Harnessing DNA replication stress to induce cell death has been a successful approach in cancer therapy. One attractive but underexplored strategy to selectively enhance replication stress in cancer cells is to inhibit the licensing of replication origins, i.e., the loading of replicative helicases onto DNA. We hypothesize that origin licensing inhibitors would serve as powerful cancer drugs, either in stand-alone or combination therapy, as they sensitize cancer cells to replication stress through a mechanism distinct from established chemotherapeutics, eventually causing cell death. Here, we propose to use high-throughput biochemical screening to identify small-molecule origin licensing inhibitors. Positive hits will be tested in biochemical and cellular assays to establish dose-responses and identify compounds that are most effective at inhibiting DNA replication initiation. These studies will be combined with X-ray crystallography and cryo-electron microscopy to understand the structural mechanisms by which identified compounds selectively inhibit replicative helicase loading. The outcomes of these efforts will pave the way for the development of the first origin licensing inhibitors as anticancer therapeutics and, in the long term, have the potential to yield a novel class of cancer drugs.
This proposal is focused on pancreatic ductal adenocarcinoma (PDAC) – a highly lethal cancer with poor treatment options. We study alternatively spliced Tissue Factor (asTF), a protein whose expression is high in tumor tissue and blood of patients with PDAC. asTF activates integrins, which promotes proliferation, migration, and metastasis of cancer cells. We developed 1) a humanized, inhibitory monoclonal anti-asTF antibody termed hRabMab1 with a favorable pharmacokinetic profile and tumor-suppressing properties in vivo; and 2) asTF-specific ELISA. The Specific Aims we propose are: 1. Test the hypothesis that hRabMab1, a first-in-class biologic, may help treat PDAC. Our preliminary data shows that hRabMab1 suppresses the growth of pre-formed, orthotopically grown PDAC tumors when administered intravenously as a single agent. We will assess hRabMab1’s ability to suppress the growth of several PDAC cell lines and patient-derived xenografts in combination with standard-of-care regimens; and hRabMab1’s ability to impede metastases. 2. Test the hypothesis that measuring circulating asTF may help evaluate response to therapy. We found that asTF levels in plasma drop post-treatment; this extends our earlier findings that plasma asTF levels are high in PDAC patients compared to healthy subjects, and positively correlate with non-resectability. In this aim, we will determine if asTF levels rise with recurrence post-surgery. In partnership with investigators conducting a Phase II trial examining an adaptive approach to neoadjuvant therapy, we will determine if plasma asTF rises in those subjects who recur, and whether said rise precedes the emergence of conventional signs of recurrence. Our studies will help us validate asTF as a new target and a biomarker in PDAC, bringing this technology closer to the clinic.
Early cancer detection is a critical factor in patient survival. However, sensitive and specific diagnostic tools are limited. Many cancers exhibit genome instability, a hallmark of cancer. Among the causes of genome instability are mutations in DNA repair genes, including the hereditary breast cancer susceptibility genes, BRCA1 and BRCA2. Mutations in BRCA1/2 lead to elevated levels of R-loops, nucleic acid structures that form when nascent RNA hybridizes to the template DNA. Although R-loops have regulatory roles in the cell, their deregulation causes DNA damage and genome instability. We have identified RNA-DNA hybrids as a product of R-loop processing, which accumulate in the cytoplasm of cells lacking BRCA1/2. We hypothesize that these hybrids could be a circulating biomarker suitable for liquid biopsy and thus, early cancer detection. Our approach has three aims with the overarching goal of evaluating hybrids in the serum of BRCA1/2-deficient high-risk individuals. We plan to detect and sequence extracellular hybrids from BRCA1/2-deficient cell lines (aim 1) and serum samples (aim 2), and test a quantitative qPCR-based assay for serum hybrid analysis (aim 3). A highly sensitive, non-invasive, and inexpensive method for early cancer detection in individuals of elevated genetic risk would transform current management and surveillance schemes.
Alzheimer’s disease (AD) is a progressive neurodegenerative disease that affects over 5 million people in the United States and is characterized by amyloid plaques, tau tangles, loss of synapses and ultimately, neuronal death. FDA-approved drugs for AD treat the symptoms of the disease but do not improve the underlying cell damage that leads to disease progression, highlighting the need for novel neuroprotective therapeutic options. Dysfunction in the autophagic and lysosomal pathways, which are critical for protein and organelle homeostasis in neurons, is associated with different stages of the disease, and thus modulation of autophagy has emerged as a new strategy for developing AD-targeted drugs. To this end, we have identified novel, small-molecule autophagy activators in a high-throughput screen that rescued key pathological features in human neuronal culture models of the disease. We hypothesize that these novel activators will restore optimal autophagy, clear protein aggregates, prevent disease progression, and ameliorate AD symptoms in the 5xFAD mouse model. We will test this hypothesis through determination of the drug metabolism and pharmacokinetic parameters of these two autophagy activators to identify a lead compound and evaluation of the in vivo efficacy of this compound in 5xFAD mice.
“Aerosolized HIV Inhibitors for the Treatment of Laryngopharyngeal Reflux (LPR)”

More than 20% of the US population suffer from LPR with no effective medical therapy. We have identified two, well tolerated, drugs which hold promise for the treatment of LPR. While the FDA has approved a proof-of-concept clinical trial using the high dose oral formulation (Transformational Application), the long-term objective is to develop a metered inhaler for local delivery. Local administration of a low dose to the laryngopharynx can achieve the same therapeutically effective drug level as the high dose, oral formulation. It follows that the use of a lower dose will be associated with fewer and/or less severe side effects. We therefore propose to:

Specific Aim 1: Develop a water-based formulation of fosamprenavir and darunavir that is amenable for inhalation administration in humans, and

Specific Aim 2: Compare the dose response of oral and inhalation administration of fosamprenavir and darunavir for pepsin-mediated laryngeal epithelial damage in our established LPR in vivo mouse model.

Aerosol formulation offers the potential for generating Intellectual Property and thereby the means to connect with an industrial partner to carry out the detailed drug product development needed for an effective treatment for the millions of people in the US that suffer from LPR.
Allergies are a result of allergen proteins cross-linking allergen-specific IgE (sIgE) on the surface of mast cells and basophils. The diversity and complexity of allergen epitopes, and high-affinity of the sIgE–allergen interaction have impeded the development of allergen-specific inhibitors of allergic responses. This study presents the design of food allergen-specific sIgE inhibitors termed covalent heterobivalent inhibitors (cHBIs) that selectively form covalent bonds to only sIgEs, thereby permanently inhibiting them. We have developed peanut-specific inhibitors that have demonstrated efficacy and specificity in blocking mast cell or basophil degranulation and anaphylaxis using in vitro assays, ex vivo samples from peanut-allergic patients, and humanized mouse models. The next critical step in moving these inhibitors to testing in humans is more formal toxicology analysis and establishing efficacy of the inhibitors for cutaneous anaphylaxis as a prelude to testing inhibition of skin prick testing. Thus, the specific aims of this application are to 1. Validate the inhibitory potential of cHBIs for cutaneous reactivity in humanized mouse models; and 2. Assess the cHBI toxicity and pharmacokinetics in non-human primate models. Together, these Aims will provide the critical data required to move forward with IND approval and preparation for phase I clinical trials.
The overall aim of this application is to discover novel therapeutics for Plasmodium falciparum malaria. P. falciparum is a leading cause of morbidity and mortality in developing countries, infecting hundreds of millions of individuals and killing over 300,000 children each year. The spread of parasites resistant to the artemisinin family of compounds threatens recent progress achieved by antimalarial campaigns and underscores the urgent need to identify new anti-malarial drugs.

In previous work, we discovered PfGARP, a previously unrecognized vaccine candidate found only in P. falciparum. Antibodies to the highly invariant carboxyl-terminal of PfGARP (PfGARP-A, aa 411–673) inhibit parasite growth in vitro by 99% compared to controls (P < 0.001) by killing trophozoite stage parasites. In confocal and transmission electron microscopy studies, PfGARP localized to the exofacial surface of the RBC membrane in trophozoite and early schizont infected RBCs, but not to other parasite stages or uninfected RBCs. Importantly, the growth inhibition assays are performed in the absence of any immune effector molecules (complement) or cells—thus the remarkable anti-parasite effect of anti-PfGARP results from antibody binding alone. This is further supported by the killing effect of recombinant mAb (KD 2.9 nM, 95% CI = 1.3 – 5.9 nM) and its rec monovalent Fab that target aa 443-459 (VKNVIEDEDKDGEVEIIN) of PfGARP.

The Scientific Premise of this application is that PfGARP is a high-value druggable target based on: 1) its surface expression on infected RBCs, 2) the absence of any significant amino acid homology with human host proteins, and 3) the ability of antibody binding to PfGARP to kill essentially all parasites within 12-24 hours.

In the current proposal, we will: 1) conduct a targeted, high-throughput drug screen to discover drugs which mimic the lethal activity of antibodies recognizing PfGARP, 2) optimize and down select these candidates, and 3) validate these new drug candidates in a humanized mouse model of P. falciparum.
Multiple myeloma (MM) is a malignancy of plasma cells that is the second most common hematologic malignancy (20% of all cases), and remains incurable for almost all patients. The primary cause of treatment failure is upregulation of pro-survival resistance mechanisms, and identifying these mechanisms remains key for new therapeutic development. We have recently reported that the serine-threonine kinase PIM2 has a major pro-survival role in MM, and that inhibition of PIM2 with a first-in-class PIM2-selective non-ATP-competitive kinase inhibitor JP11646 (JP) caused significant MM cell death in vitro and in vivo in preclinical MM models. Unexpectedly, JP was much more effective than the ATP-competitive PIM kinase inhibitors due to JP’s unique ability to downregulate of PIM2 gene and protein expression. This has led to our findings that PIM2 has critical but previously unrecognized kinase-independent (KI) functions, which would be a change in paradigm. One such PIM2 KI function that is inhibited by JP is induction of the expression (mRNA and protein) of the c-Myc oncogene, which is dysregulated in ~50% of all human cancers, including MM. Our additional findings suggest a completely novel mechanism where JP disrupts PIM2 interaction with a partner protein (possibly c-Myc itself) that results in downregulation of c-Myc expression and loss of c-Myc driven PIM2 gene expression, collapsing a self-reinforcing loop that sustains both PIM2 and c-Myc expression. Targeting c-Myc expression through PIM2 inhibition represents an entirely unexplored therapeutic approach. The overall goal of this proposal is to develop JP11646 for the treatment of refractory/relapsed MM, and potentially for other cancers. The Specific Aims of the proposal are:

Aim 1. Define the kinase-independent mechanisms by which PIM2 supports MM survival, and how JP11646 inhibits these.

Aim 2. Conduct preclinical and IND enabling studies for a phase I clinical trial of JP11646 in refractory/relapsed multiple myeloma.
Recent evidence has associated overactivation of canonical Wnt signaling through β-catenin (cWnt) with multiple pathological processes in osteoarthritis (OA). The cWnt agonist, R-spondin 2 (Rspo2), is a soluble ligand that our preliminary data demonstrates is sufficient to induce joint degeneration in vivo, making it a promising therapeutic target. The antidepressant drug Mianserin was recently identified as an Rspo2 inhibitor by blocking Lgr5 binding. Our overall objective is to elucidate the role of Rspo2 in OA pathogenesis and test the efficacy of sustained intra-articular Rspo2 inhibition using Mianserin-loaded polymeric microspheres.

We will:
Aim 1: Demonstrate the joint-protective effect of Rspo2 ablation. We hypothesize that joint injury-induced Rspo2 potentiates OA by activating cWnt signaling in multiple intra-articular cell types, promoting pathology. Using a global, inducible Rosa26-CreERT2; Rspo2flox mouse, we will globally ablate Rspo2 at the time of injury (Rspo2cKO) and comprehensively evaluate OA severity.

Aim 2: Test the efficacy of sustained intra-articular Rspo2 inhibitor therapy as a novel OA treatment. We will formulate PLGA microspheres to deliver the Rspo2 inhibitor Mianserin to the joint, assessing dose-dependent pharmacokinetics and pharmacodynamics of Mianserin release in vivo. Then, we will test the OA disease-modifying effects of Mianserin-loaded PLGA-MS in mice, benchmarking against repeated direct Mianserin injections.
CAR-T immunotherapies have produced remarkable clinical responses, but several challenges remain including disease relapse due to antigen escape/decreased CAR-T persistence, fatal side effects (cytokine release syndrome, neurotoxicity), the cost and time to produce CAR-T cells. We have developed novel B cell activating factor (BAFF) ligand-based BAFF CAR with multi-receptor specificity to target BAFF receptors (BCMA, TACI, BAFF-R) expressed by B cell cancers. We deliver BAFF-CAR to human NK cells using a non-viral TcBuster transposon platform, enabling safer, simpler, and more cost-effective CAR-NK production. Our preliminary data demonstrates in vitro and in vivo cytotoxicity of BAFF CAR-T cells against human malignant B cell lines. The overall objective of this proposal is to develop, optimize and evaluate BAFF-CAR NK cell therapy for the treatment of B cell malignancies. Aim 1: Achieve stable, non-viral integration of various BAFF CAR constructs in human NK cells. Aim 2: In vitro functional validation and characterization of BAFF CAR-NK. Aim 3: In vivo evaluation of dosing and efficacy of BAFF-CAR-NK cells using mantle cell lymphoma (MCL) xenograft NSG mouse model. Persistence of engineered NK cells will be evaluated in parallel with efficacy, as well as examination of preliminary safety endpoints and disease relapse.
DNA polymerase (Polq) is a unique polymerase-helicase DNA repair protein that has recently been validated as a synthetic lethal drug target in cancer cells harboring mutations in BRCA1 or BRCA2 (BRCA) tumor suppressor proteins. Polq promotes DNA repair via the microhomology-mediated end-joining (MMEJ) pathway (also referred to as alternative end-joining). Polq is essential for BRCA-deficient cells, but is dispensable for normal cells and Polq null mice show no phenotypes. Thus, Polq is synthetic lethal with BRCA1/2. Consistent with this, recently published Polq inhibitors (Polqi) selectively kill BRCA-deficient cells, while showing no effects in BRCA-proficient cells. Yet, whether these early stage Polqi will lead to clinically effective drugs is unknown.

Towards the goal of developing a clinically effective Polqi, our laboratory and collaborators have taken a different approach for targeting Polq: to develop a Proteolysis Targeted Chimera (PROTAC) Polq degrader to completely abolish all Polq activities in cancer cells. Because Polq null mice are healthy with no phenotypes, simultaneous inactivation of Polq polymerase (Polq-pol) and helicase (Polq-hel) enzymatic domains is expected to be a safe and a highly effective therapeutic method. Based on this rationale, we plan to develop a Polq PROTAC targeting the Polq-pol domain as a unique therapeutic for treating BRCA-mutant cancers.
Our drug discovery effort seeks to develop brain-penetrable and orally available small molecule pharmacological chaperones to restore defective acid β-glucosidase (GCase) as a new disease-modifying treatment for neuronopathic Gaucher disease and Parkinson’s disease. We have identified novel small molecules that are non-inhibitory chaperones capable of improving multiple important endpoints: a) GCase activity, b) GCase substrate reduction and c) penetration through the blood brain barrier. We are presently at the critical optimization stage and will select lead compounds to advance as drug candidates for preclinical studies. In this proposal, we will optimize our lead compounds to improve activity and bioavailability. Newly designed compounds will be screened to increase GCase activity using cell-based multi-well screening platform. The chaperone property of selected compounds will be evaluated in fibroblasts and neurons carrying common GBA1 mutations found in Gaucher disease and Parkinson’s disease patients. Selected compounds will be assessed for CNS drug properties, biodistribution, and preclinical evaluation in our diseased mouse models. Upon completion of the proposed milestones, we will have identified promising drug candidates of non-inhibitory chaperones that will be implemented for IND-enabling studies and advance a drug candidate into clinical trials to test the compound’s benefit for patients with Gaucher and Parkinson’s diseases.
Sherry Thornton, Ph.D. - 2021 Awardee

Rheumatology
Children's Hospital Medical Center dba Cincinnati Children’s Hospital Medical

“Biomarkers for the Advancement of Targeted Therapies for Children with Juvenile Arthritis”

Juvenile idiopathic arthritis (JIA) is the most common chronic inflammatory rheumatic disease of childhood, affecting 1:1,000 children worldwide. This prevalence is about the same as juvenile diabetes. Currently, physicians lack laboratory tests to support them in the initial choice and sequence of medications prescribed to a child with JIA. This deficiency increases not only health care cost but also adversely impacts the prognosis of children with JIA. What is needed is a JIA-Rx Biomarker Panel, i.e., a comprehensive, yet concise group of biomarkers that helps anticipate the response of a child with JIA to a given immunosuppressive treatment. Over the next 1 year, we will (1) delineate serum biomarkers that individually, or in combination, forecast response of JIA therapy; (2) develop a mathematical algorithm to provide a numeric estimate for the probability of experiencing response to therapy, and (3) test the performance of these candidate biomarkers and algorithm in children with JIA who are newly started on TNF-blocking medications. We will use serum samples, gene expression RNAseq data, and clinical information from 187 children with JIA who participated in a 44-week, double-blinded international clinical trial (NCT02592434) of tofacitinib. Data and samples have been donated by Pfizer to CCHMC Rheumatology. We will enroll 60 children with JIA who are newly prescribed TNF inhibiting medication and study them prospectively. We expect to delineate an actionable JIA-Rx Biomarker Panel and interpretation rules to immediately support treatment decisions for children with JIA.
Triple-negative breast cancer (TNBC) is the most aggressive breast cancer. Treatment options for this subtype are limited, as it lacks expression of ER, PR, and HER2. Immune checkpoint inhibitors (ICI) have shown promising effects, but response rates are low, underscoring the need for combination therapy. However, there are many critical roadblocks for developing combination regiments, including identifying which combinations will be effective, and identifying biomarkers to select patients that are most likely to benefit from treatment.

This project is based on our groundbreaking discovery of the role of CARM1-mediated BAF155 methylation in modulating the efficacy of immunotherapy. We found that pharmacological inhibition of CARM1 boosted host immune responses by enhancing cytotoxic T cell activity and tumor infiltration, leading to tumor regression. Moreover, a combination of CARM1 inhibitor with ICI synergistically inhibited tumor growth. CARM1 is overexpressed in TNBC and is a druggable enzyme with several pharmaceutical inhibitors. Our studies suggest that inhibiting CARM1 methylation of BAF155 primes tumors to become immunologically ‘hot’, and sensitizes BC to ICIs.

Furthermore, methylated BAF155 (me-BAF155) is readily detected in circulating tumor cells of metastatic BC patients, making me-BAF155 a potential biomarker for selecting patients for the combined CARM1i and immunotherapy.
The peripheral nervous system is an appealing site for devising new analgesic treatments since primary sensory neurons (PSN) play a central role in the development and maintenance of chronic pain, and can be safely medicated in a highly targeted fashion. The Nav1.7 sodium channel plays fundamental roles in mediating pain, so development of PSN-targeted NaV1.7 inhibitors could provide highly effective analgesia. However, success in creating small molecule NaV1.7 inhibitors remains elusive. For a new approach, we propose a strategy in which small peptides derived from Nav1.7 ion channels are developed as interfering peptide aptamers (iPAs) for highly effective and selective blockade of pain signaling. These will be delivered selectively to anatomically targeted sensory neurons by recombinant adeno-associated viral (AAV) vectors to achieve analgesia without side effects or addiction potential. Candidate peptides will be identified by combined computational and experimental strategies and tested for selective and effective NaV1.7 channel blockade. Analgesic efficacy will be tested in established animal pain models representing neuropathic and arthritis pain, following AAV-NaV1.7iPA injection into peripheral segmental nerves. Preliminary observations with a prototypic NaV1.7iPA candidate predict highly safe and effective analgesia. Achievement of these goals will provide candidate agents suitable for development in the subsequent transformational phase.
Glioblastoma is an incurable primary brain tumor that is characterized by regions of hypoxia and marked resistance to radiation. A major roadblock in the treatment of glioblastoma is the inability to kill the cells that give rise to disease recurrence, the glioma stem-like cells (GSCs). GSCs are highly resistant to standard cytotoxic treatments, have a high capacity for self-renewal, and are frequently located in hypoxic areas, which renders them even more difficult to kill with radiation and chemotherapy. Here, we propose to develop a new therapeutic strategy to kill these GSCs, with the long-term goal of extending glioblastoma patient survival. We have found that the long non-coding RNA (lncRNA) Lucat1 is an important regulator of GSC response to hypoxia. Lucat1 is frequently overexpressed in glioblastoma and is associated with poor prognosis. Our data support that Lucat1 is induced by hypoxia and forms a positive regulatory loop to promote HIF1alpha-mediated signaling. Lucat1 helps to maintain GSCs in hypoxia and promote tumor growth. By targeting Lucat1, GSCs fail to adapt to hypoxia, resulting in cell death. Silencing Lucat1 extends animal survival in mouse models of glioblastoma. Importantly, our data suggest that targeting Lucat1 has fewer side effects than direct inhibition of hypoxia regulator HIF1alpha. In this study, we propose to develop and optimize novel anti-sense oligonucleotide (ASO) therapeutics that inhibit Lucat1-HIF1alpha signaling and test the efficacy of these ASOs in mouse models of glioblastoma. Our findings will provide a new therapeutic approach for targeting GSCs in hypoxia to improve glioblastoma control.
Lupus, incurable autoimmune disease affects round 1.5 million persons living in the United States. Among the main risk factors for poor disease outcome is the presence of lupus nephritis (LN). The applicants have discovered and initially validated a panel of LN-biomarkers [NGAL, MCP-1, Kim-1, ceruloplasmin, adiponectin, hemopexin]. Considering the urine levels of these LN-biomarker we delineated an algorithm, the Renal Activity Lupus Index (RAIL), where higher scores reflect more kidney inflammation as seen on kidney biopsy. Changes in RAIL scores allow us to predict LN flares and LN response to therapy at least 3 months earlier than when using current clinical tools. Further, our pilot studies suggest that the RAIL score can be used to refine the dosing of current LN therapies; we observed that high-dose intravenous steroids lead to a dramatic reduction of the RAIL score that does not occur when only daily oral steroids are used for LN therapy.

Critical Scientific Roadblock to be addressed: To enable the application of these highly promising LN-biomarkers and the RAIL algorithm in clinical care and for research, following work needs to be conducted: (1) development of a high-quality multiplex assay (RAIL-MPLA) to rapidly measure all of the LN-biomarkers concurrently; (2) additional validation of the RAIL in an independent cohort of adults and children.

Approach: Working together with Ethos Research & Development and pharmaceutical companies, we will: (1) develop a RAIL-MPLA; (2) define changes of RAIL scores that reflect clinically relevant improvement of LN and reference values for absent, controlled and active LN across age-ranges; and (3) test the hypothesis that high-dose intravenous steroids result in a significantly higher reduction of the RAIL score compared to standard doses of oral steroids alone.

Expected Results/Deliverables:
- Develop a novel clinically actionable multiplex assay that accurately quantifies the biomarkers used in the RAIL within 4 hours of sample receipt.
- Increase the usability of RAIL by refining the interpretation of RAIL scores
- Immediately advance the treatment of LN by providing the scientific underpinning for the preferred use of intravenous rather than oral steroids alone.
Women with early-stage estrogen receptor-positive (ER+) breast cancer face a persistent risk of distant recurrence and breast cancer death up to 20 years post diagnosis. We have recently reported the 20 year follow-up of nearly 2000 early-stage breast cancer patients with accompanying tumor molecular profiling. This work identified 11 integrative clusters (ICs) of breast cancer, defined based on integration of genomic copy number alterations and transcriptional profiles, with markedly variable risk of recurrence over time. Four of these ICs had exceedingly high risk of recurrence – over 40 percent 20 years post diagnosis – and together comprised nearly one-quarter of ER+/HER2- patients. Importantly, each of these ICs harbored a characteristic and putatively targetable gene amplifications. We hypothesize that therapy targeted at the driver genes present in each of the high-risk ER+/HER2- IC groups will reduce tumor cell proliferation, providing a path forward to design targeted therapeutic strategies aimed at preventing recurrence and reducing mortality associated with lethal metastatic breast cancer. Having identified and validated biomarkers of breast cancer relapse, several of which themselves represent potential new therapeutic targets, we propose to functionally characterize the drivers of the high-risk Integrative Clusters (ICs), to uncover their oncogenic dependencies and to elucidate mechanisms of resistance to endocrine and targeted therapies. Accordingly, we will leverage advances in CRISPR technology to perform pooled multiplexed CRISPR/dCas9-interference (CRISPRi) screens in representative 2D cell lines and 3D spheroid cultures to delineate 3D-specific vulnerabilities (Aim 1). In tandem we propose reciprocal experiments in which we oncogene engineer non-malignant mammary cells via CRISPR/dCas9-activation (CRISPRa) pooled screens to define the genes that promote malignant growth (Aim 2). Further, we will exploit a powerful cellular barcoding techniques to trace clonal dynamics during treatment and delineate the functional determinants of response to targeted and endocrine therapies in representative patient-derived organoid models (Aim 3). These complementary approaches will systematically define novel therapeutic targets and mediators of drug sensitivity in aggressive ER+ breast cancer, thus informing subsequent clinical trials.
Richard Harvey, M.D. - 2020 Awardee

Physical Medicine and Rehabilitation and Physical therapy and Human Movement Sciences
Rehabilitation Institute of Chicago


The purpose of this project is to develop a usable bedside hand therapy device that takes advantage of early neuroplasticity following stroke and cervical spinal cord injury (SCI) to retrain functionally relevant movement in individual fingers of the neurologically impaired hand. It is well established in neuroscience that neural recovery and neuroplasticity at the level of cortex in animals and humans is dependent on active motor practice. In this Catalyst project we will determine and assure that the second-generation device we have develop, which has not yet been used in humans, is usable in the clinical setting and that patients with stroke and cervical SCI find it a meaningful tool for relearning hand movement. We will also assess whether use of the device daily for 2 weeks results in improved functional hand use, improve finger strength and improved ability to control individual finger movements. Although these abilities are foundational to functional manipulation of objects with the hand, there is presently no specific therapeutic interventions, nor time available in conventional early (acute) rehabilitation to address these goals. Thus, early intensive hand recovery is an obvious but unmet need in neurorehabilitation. Our goals for this one-year project are determine if the device:
1. can facilitate improve hand function.
2. can improve finger strength and individuation.
3. is usable in a clinical setting and serves as a meaningful therapy tool to patients with stroke and cervical SCI.

If successful, this project will lead to a refined therapy protocol in preparation for a larger clinical trial to establish efficacy of this training approach for both stroke and cervical spinal cord injury. We chose to study both patient populations as we believe this tool can have benefit for both groups and to study both provides generalizability to a wider group of patients needing rehabilitation. If we achieve our long-term goal, this device will evaluated in a large multi-center pivotal trial in preparation for transition to the clinical marketplace as new rehabilitation technology that has proven scientific evidence supporting its efficacy in early hand rehabilitation.
Treatment resistance causes nearly all cancer deaths. Novel treatments that bypass this resistance and inhibit the biology that drives cancer progression are needed to improve patient survival. Using prostate cancer (CaP) as model, we isolated citron kinase (CIT) as a target for such a therapy. CIT is a previously unrecognized novel druggable target that is functionally distinct from current targets yet is a critical determinant of cancer cell proliferation and induces cancer growth and treatment resistance. CIT silencing inhibited growth of cell lines and xenografts representing different stages of CaP progression and diverse forms of treatment resistance but not of benign epithelial prostate cells. CIT’s stimulation of cancer progression relied entirely on its kinase activity, isolating CIT’s kinase activity as a completely new target. A specific CIT inhibitor has not yet been developed. By analyzing the target spectrum of kinase inhibitors and CIT kinase assays, we identified 3 multikinase inhibitors that inhibit CIT, decrease CaP cell proliferation and inhibit CIT substrate phosphorylation. Because these drugs are not selective for CIT and thus not likely to succeed as cancer therapies, we developed a lead compound that inhibits CIT at low doses, has improved CIT selectivity and promising DMPK. Our objective is to develop our lead compound into first-in-class selective CIT inhibitor as a novel cancer treatment. We hypothesize that novel CIT inhibitors will overcome acquired resistance to conventional cancer therapies, which will be reflected in the phosphorylation status of CIT substrates. We will test 2 Specific Aims:
1. To determine the therapeutic efficacy of novel CIT inhibitors during cancer progression using multiple chemistry and crystallography approaches and kinome screens to improve CIT specificity and DMPK and verifying growth inhibition of clinical relevant treatment-resistant cancer models.
2. To determine the substrates by which CIT kinase action conveys aggressive cancer behavior using integrated state-of-the-art biotin-based proximity ligation assays, kinase substrate arrays, and mass spectrometry approaches.
We expect these studies to have a significant positive impact because they will provide CIT inhibition as an entirely new and functionally diverse cancer treatment and the phosphorylation status of CIT substrates as a treatment-specific biomarker of response.
Declining immune function is well described in the elderly, and leads to increased risk and severity of infection, poorer control of cancer, and impaired responses to vaccination, all major public health concerns. Further, the elderly population is at high risk of mortality if infected with new viral pathogens, such as SARS-CoV-2. So, even if vaccines are quickly developed against emerging pathogens, they are likely to have reduced efficacy in the elderly. Thus, there is an urgent need to identify strategies to increase vaccine responses in elderly humans.

Our exciting and newly published data show that levels of IL-10, an immune suppressive cytokine, are increased dramatically with age. Further, we found that the majority of this IL-10 is produced by a population of cells called T follicular helper (Tfh) cells, which are normally critical for productive antibody responses. Notably, these aged, IL-10-producing Tfh cells are found in both mice and humans. Strikingly, neutralization of IL-10R signaling in aged mice substantially increases antibody responses, nearly to the levels observed in young mice. In monkeys, we defined the pharmacokinetics of an IL-10 neutralizing antibody as well as a soluble receptor. Given these compelling data, we hypothesize that Tfh10 cells play an important role in impaired age-related vaccine responsiveness that is conserved between mice, monkeys, and humans. Here, we propose to determine if short-term IL-10 blockade will restore Flu vaccine responsiveness in aged macaques. We propose an iterative approach, assessing whether concomitant blockade of IL-10 alongside flu vaccination will: (i) increase anti-influenza neutralizing antibody responses in response vaccination without increasing systemic inflammation (ii) promote resistance to influenza challenge. These studies will likely have important translational implications, as a short-term blockade of IL-10 signaling could be envisioned to improve vaccine responsiveness in elderly humans. Our long-term goal is to develop therapeutic strategies to enhance protective immune responses in the elderly.
“Development of Spherical Nucleic Acid-Based Anti-Sense Oligos for the Therapeutic Treatment of Pediatric Epilepsy Disorders”

Heterozygous mutations in KCNQ2, which encodes a pore-forming K+ channel subunit responsible for neuronal M-current, cause neonatal epileptic encephalopathy (EE), a complex disorder presenting with severe early-onset seizures and impaired neurodevelopment. The condition is exceptionally difficult to treat, partially because the effects of KCNQ2 mutations on the development and function of human neurons are unknown. Using induced pluripotent stem cells (iPSCs) and gene editing we have established KCNQ2-EE disease model systems, and measured the functional properties of patient-derived neurons using electrophysiological and optical approaches at single-cell resolution. We find that patient-derived excitatory neurons develop intrinsic and network hyperexcitability that mimics epileptic electroencephalogram (EEG) activity. Using heterologous expression systems, we have also found that some disease-causing variants exhibit a dominant-negative effect, reducing the channel activity by more than 50%. We hypothesize that deleting the mutant transcript will restore channel activity and alleviate the associated neuronal firing defects. Here, we propose to collaborate with Exicure Inc, to design, screen and test allele-specific antisense oligonucleotides (ASOs), which will target and degrade the mutant KCNQ2 allele. Exicure, is a clinical-stage biotechnology company developing spherical nucleic acid (SNA) based ASOs for genetic disorders. We will assess the specificity of ASOs using established digital droplet PCR assays and examine the efficacy of ASOs by measuring their ability to restore the firing activity of patient-derived neurons. If successful, our work will provide a rational therapeutic approach for this devastating disease.
Intellectual disabilities are common and carry high lifetime costs to families. We have identified a new neurodevelopmental disorder caused by loss-of-function mutations in the mitochondrial enzyme glutamate pyruvate transaminase 2 (GPT2). GPT2 disease involves postnatal microcephaly, cognitive disability, epilepsy, and progressive spastic paraplegia. GPT2 localizes to mitochondria and catalyzes the reversible addition of an amino group from glutamate to pyruvate, yielding alanine and alpha-ketoglutarate, a metabolite in the tricarboxylic acid (TCA) cycle. Metabolic diseases in children, such as GPT2 disease, may be amenable to treatments, via dietary restrictions or supplements, that can prevent intellectual disabilities when treated early. With the prospect of newborn screening supported by genome-wide sequencing on the horizon, there will be new opportunities to intervene in childhood brain disease. Therefore, we rapidly need to determine which new neurometabolic diseases may be amenable to interventions in early childhood. Our preliminary data provide support for disease mechanisms wherein GPT2 plays a critical role in alanine synthesis, as well as in neuronal anaplerosis. Anaplerosis (filling-up) is the metabolic process whereby TCA cycle intermediates are replenished. Anaplerosis is important during high biosynthetic demand, such as during brain development, when TCA cycle intermediates are consumed for synthesis of macromolecules for cell growth. We have established potential treatment strategies for GPT2 disease that may be implemented in the near term. To guide these interventions, we propose to complete needed pre-clinical studies. Our overriding hypothesis is that GPT2 is required for metabolic mechanisms central to neuronal and axonal growth during brain development, particularly in long projection neurons of the motor system. In Aim 1, we will test mechanism-based treatments in our mouse model of GPT2 disease. Our Gpt2-null mouse recapitulates key aspects of disease, such as motor abnormalities, akin to spastic paraplegia seen in patients. In Aim 2, to translate our basic findings to the clinic, we will establish the infrastructure and collaborations, including with industry, to develop a protocol for biomarker studies and a clinical trial in patients. The impact of this research is that we will establish the basis for therapeutics in a childhood disease for which there is no current treatment.
Lonnie Shea, Ph.D. - 2020 Awardee

Biomedical Engineering
University of Michigan

“A Cell Capture Implant to Predict Acute Cardiac Allograft Rejection”

Over 36,000 solid organ transplants are conducted annually in the US, costing $30 billion. Immunosuppressive drugs protect these donor grafts from acute rejection but increase the risk of opportunistic infections and cancer, especially in pediatric transplant recipients, who require immune suppression for decades. As there is no method for determining which grafts will be rejected, immunosuppression is aggressively applied in a one-size-fits-all approach.

We have developed a novel implantable scaffold device to monitor recipient immune response to the transplant that would enable the use of personalized immunosuppression while maintaining graft acceptance. We employed these scaffold devices in a murine skin allograft model to develop a highly sensitive and specific scoring system from a 19 gene biomarker signature to distinguish between recipients with healthy grafts and recipients with rejecting grafts.

In heart transplantation, organ scarcity requires that clinicians remain vigilant in preventing acute cardiac allograft rejection (ACAR) through frequent graft biopsy and aggressive immunosuppression, at the expense of toxicities due to over-suppression. A novel surveillance method is urgently needed to calculate the early risk of transplant rejection to allow personalized immunosuppression regimes. Here, we present a novel subcutaneous scaffold implant that collects biomarkers of graft health that will predict ACAR onset to preserve graft function while reducing the need for frequent graft biopsy and overly aggressive immunosuppression. In this proposed work, we will employ these minimally-invasive cell capture scaffolds in four cohorts of heterotopic heart transplants in mice in which we will identify a signature gene panel of graft rejection, validate this scaffold biomarker signature, assess the predictive power of the signature, and employ the biomarker signature to monitor the immunosuppression response. We hypothesize that the dynamic changes captured in the scaffold device as the allograft is recognized as non-self by the recipient immune system will enable the development of a novel predictive biomarker panel for ACAR. This minimally-invasive immune cell-capturing scaffold could prove transformative in directing and personalizing immunosuppression in solid organ transplant recipients.
David Spiegel, M.D., Ph.D. - 2020 Awardee

Chemistry
Yale School of Medicine

“Development of Bifunctional Molecules that Cross the Blood-Brain Barrier and Degrade Pathogenic Neurological Proteins”

Unique challenges are presented when treating neurological disease due to the physical obstacle formed by the Blood-Brain Barrier (BBB). Many neurodegenerative diseases result from the accumulation of pathogenic proteins and could potentially be avoided with the early detection and removal of these species. The Spiegel lab is developing a novel class of bioactive molecules that can bind proteins non-covalently and chaperone them across the BBB. These small molecules will be used for the removal of pathogenic proteins from the brain as a novel therapeutic approach to the treatment of neurological diseases. The bifunctional small molecules consist of a protein-targeting motif and a motif for the transport of the target protein across the blood-brain barrier, and subsequent degradation.
Devastating and persistent functional deficits occur after Spinal Cord Injury (SCI), despite survival of nearly all neurons. Partial recovery would be adequate for patients to gain a level of independence in wheelchair transfers, bowel and bladder management, and locomotion. Today, there is no approved medical therapy for the over 300,000 individuals in the USA with SCI, rendering therapy a totally unmet medical need. Because the primary cause of disability is disconnection of networks by axon transection, axon regrowth has the potential to provide recovery by restoring connectivity, without requiring “new” cells. Unfortunately, without therapeutic intervention, the adult brain and spinal cord produce extremely limited reparative axon growth after damage.

We surveyed the mouse genome in an unbiased approach for loci with undiscovered functions in axon regeneration. This screen identified 400 genes whose suppression yielded greater axon regeneration in vitro. In Preliminary Studies, we leveraged the in vitro results to identify pathways that might be targeted to improve CNS neural repair in vivo using a simple optic nerve crush model. In control eyes, very few axons regenerate, but knockdown of 40 of the in vitro hit genes produced significantly increased regeneration. These genes were validated gene editing using CRISPR/Cas9 in vitro and in vivo.

In the current proposal, we will test two of these novel regeneration genes as attractive therapeutic candidates for their ability to promote axon growth and behavioral recovery after spinal cord injury in mice. We hypothesize that their demonstrated axon regeneration activity will translate into improved neural repair. Validation of this hypothesis will support development of their potential in translational research.
The goal of this project is to develop an effective therapy for Idiopathic pulmonary fibrosis (IPF), a devastating lung disease that progressively scars the lung and has a prognosis worse than many aggressive cancers. A defining feature of IPF is the differentiation of fibroblasts into myofibroblasts, followed by the production of massive amounts of extracellular matrix (ECM) and the formation of collagen-rich scars, which restrict movement of the lung and impede respiratory gas exchange. While two anti-fibrotic drugs are approved to treat IPF, neither therapy is highly effective. Our long-term objective, and the focus of this high-risk, high-yield application is to develop a new, effective therapy for IPF. Previous work has shown that differentiation of embryonic and hematopoietic stem cells requires the transient decondensation of chromatin at sites of repressed genes and that failure to do so prevents the binding of critical differentiation transcription factors (TF). The hallmark of condensed arrays of nucleosomes in chromatin is H3K27me3, and transient global removal of this mark is required for activating repressed genes during differentiation of stem and progenitor cells. The same mechanism was also found in more differentiated naive T cells during their differentiation into Th1 and Th2 helper cells, leading us to hypothesize that removal of H3K27me3 chromatin marks might also dictate the differentiation of fibroblasts into myofibroblasts. Because transient de-condensation of the H3K27me3-marked chromatin is achieved by the activities of the H3K27me3 de-methylases (KDM) UTX and JMJD3, we hypothesized that inhibiting these enzymes via a small molecule ‘epigenetic’ inhibitor GSKJ4 would arrest pulmonary fibrosis. In preliminary studies presented here, we show that GSKJ4 effectively blocks the formation of myofibroblasts in vitro and limits the production of extracellular matrix genes in experimentally induced IPF mouse models. We now wish to extend these observations in hopes of moving this concept to the clinic. To achieve this goal, we propose 3 rationally designed Specific Aims that are mechanistically link but fully independent. Studies in this application aim to challenge current approaches to the treatment of IPF and provide support for targeting nascent chromatin in the treatment of a wide assortment of fibrotic diseases.
Chronic Obstructive Pulmonary Disease (COPD) affects 13.5 million (M) people in the US with a regional age-standardized death rate that will be the 4th leading cause of death by 2020. This obstructive lung disease is characterized by acute episodic decompensations, “exacerbations” associated with increased symptoms that result in substantial morbidity and costs through unscheduled hospital visits. Importantly, exacerbations are associated with more rapid declines in pulmonary function for which no effective therapies exist. These patients suffer substantially reduced quality of life and over half will die of acute-on-chronic respiratory failure.

COPD is initially triggered by oxidative airway injury that results in the activation of a pathogenic mesenchymal myofibroblast population in the small bronchioles of the airway. Myofibroblasts produce fibronectin and collagen that reduce the normal elastic properties of the lung and impair gas exchange.

The broad goal of this Catalyst Award is to advance first in-class therapeutics that target myofibroblast transdifferentiation. This advance will come in two stages, each a focus in this application. First, there are no high resolution, non-invasive methods for detection of airway remodeling or determination of therapeutic response. We will validate an integrated proteomic and imaging diagnostic for airway remodeling based on our unbiased pharmacoproteomics study of inhibiting BRD4 in airway remodeling. Quantitative selective reaction monitoring assays of a panel of airway remodeling proteins will be paired with minimally invasive optical imaging of mucosal collagen deposition using optical coherence tomography (OCT). Second, we will advance a lead highly selective BRD4 inhibitor to an aerosol formulation encapsulated in proprietary Dendron Micelles. Understanding that foci of myofibroblasts are initially formed in the small bronchioles, we will develop aerosolized therapeutics that target fibronectin-rich foci. Nanoparticle formulation have additional advantages that they will enhance duration of effect, and reduce potential systemic toxicity. We will demonstrate enrichment using advance mass spectrometry based imaging, and demonstrate efficacy in an established model of COPD. By advancing, in parallel, minimally invasive biomarkers and targeted therapeutics, we will be uniquely poised to translate these into first-in human studies.
The gene encoding RUNX1 (AML1) is disrupted by the t(8;21) that is associated with 4-12% of adult acute myeloid leukemia (AML) patients and ~13% of pediatric AML patients. The t(8;21) results in a fusion protein containing the N-terminus of RUNX1, including the Runt domain, fused to almost all of ETO. The RUNX1-ETO (also called AML1-ETO) fusion protein acts as a dominant repressor of RUNX1 function, dysregulating the expression of multiple genes required for normal hematopoiesis and, in cooperation with secondary mutations, leads to the development of leukemia. The RUNX1-ETO fusion protein has clearly been established as the primary driver of t(8;21) AML. About 60-70% of t(8;21) patients are alive at 5 years, however disease recurrence is the major treatment failure with 30-40% of these patients relapsing after standard intensive chemotherapy, highlighting the need for new approaches to treatment. The standard chemotherapy used to treat these patients has serious long-term side effects, which is particularly problematic for pediatric patients who will deal with these effects throughout their lives as well as for older patients who can’t tolerate standard chemotherapy as well. In order to overcome this, it is essential to develop drugs which directly and selectively target the RUNX1-ETO fusion protein driver to treat the disease.

We propose to develop small molecule inhibitors of RUNX1-ETO that block its ability to bind to DNA and which selectively inhibit RUNX1-ETO while having minimal if any effect on wildtype RUNX function. Such a high level of selectivity of action has rarely been achieved, but we have previously done so in targeting the CBFβ-SMMHC fusion protein driver in inv(16) AML. We are proposing to develop a hetero-bivalent compound which targets both the Runt domain of RUNX1 (DNA binding domain) and the TAF domain of ETO (the nearest ETO domain to the Runt domain in the fusion protein). We have screened and optimized a compound which binds to the TAF domain displacing HEB with a low µM IC50. We have identified compounds which covalently react with Cys residues on the Runt domain. For this grant, we will focus on covalently linking the TAF domain inhibitor to optimized Cys reactive compounds. Linker length and chemical structure will be varied to identify optimal linkers. Compounds will be evaluated in t(8;21) cell lines for their ability to inhibit growth, alter the expression of RUNX1-ETO target genes (qPCR), and ability to inhibit binding of RUNX1-ETO to its genomic sites (ChIP).
“Developing Therapeutic Agents to Rescue Neurotoxicity of FUS Aberrant Phase Transition”

It is now universally appreciated that accumulation of misfolded proteins, which can acquire alternative proteotoxic states, causes a series of deleterious molecular events resulting in numerous lethal neurodegenerative diseases. Among these, amyotrophic lateral sclerosis (ALS) is the most common motor neuron disease. There are no effective therapies for ALS and very few treatment options. Therefore, new therapeutic target and strategies are greatly needed. ALS-linked mutations have been discovered in several prion-like domain (PrLD) containing nuclear RNA-binding proteins (RBPs) including FUS, which is a stress granule (SG) component. Cytoplasmic mislocalization and inclusion formation, which are common pathological features of FUS proteinopathies, have been connected to persistent SGs. Upon stress, ALS disease proteins are recruited to SGs, which are reversible cytoplasmic membraneless organelles that form through Liquid-liquid phase separation (LLPS) and behave like liquid-droplets. Because SGs condensate ALS disease proteins such as FUS that are intrinsically aggregation-prone, if SGs are not cleared in time, FUS can go through aberrant phase transition to form solid fibrils that can induce toxicity and neurodegeneration. The long-term goal of this project includes understanding the molecular mechanisms underlying the aberrant phase transitions of SGs and leveraging our understanding of aberrant phase transition to develop therapeutic agents to mitigate the neurotoxicity of these pathological events. This proposal focuses on developing two types of therapeutic agents to rescue FUS neurotoxicity caused by aberrant phase transition. We have discovered a novel function of nuclear import receptor-Kapbeta2 in reversing FUS aberrant phase transition and aggregation. However, ALS-causing mutations in FUS PY-NLS such as P525L reduces Kapbeta2’s activity as protein disaggregase. Therefore, the first goal of the proposal is to discover small molecules that can enhance Kapbeta2’s activity to reverse FUS P525L aberrant phase transition. Kapbeta2 reverses FUS aberrant phase transition by binding to the nuclear localization signal PY-NLS in the C-terminus of FUS. Thus, other FUS-binding biomolecules might also prevent and reverse FUS LLPS and aggregation. Indeed, our preliminary data show FUS-binding RNA can prevent and reverse FUS aggregation. Therefore, the second aim of the proposal focuses on developing RNA oligonucleotides to reverse FUS aberrant phase transition and defining their therapeutic potential in mitigating FUS neurotoxicity. The in vitro activities of the agents developed in this proposal will be characterized using pure protein biochemical and biophysical assays. Top ranking agents will then be validated in iPSC-derived motor neurons for their ability to mitigate FUS aberrant phase transition and the resulted neurotoxicity.
Venous thromboembolism (VTE), comprised of deep venous thrombosis and secondary pulmonary embolism, affects 900,000 people and is the third leading cause of cardiovascular death. Current therapies are limited to anticoagulation, which carries significant bleeding risks and don’t address the inflammatory processes that initiate and propagate VTE. Therapies that directly target inflammatory processes would address the underlying catalyst of disease and have the potential to fundamentally transform the treatment of VTE.

A distinguishing feature of VTE is the recruitment of leukocytes to the thrombus. We discovered that this is driven by marked neutrophil activation, the expulsion of DNA as neutrophil extracellular traps (NETs), and inflammasome activation to release a potent inflammatory cytokine, interleukin 1-beta (IL-1beta). This phenomenon further polarizes the vascular environment in a self-amplifying loop culminating in VTE. We showed that hyperactive neutrophils are more prone to form NET ‘scaffolds’ for thrombus expansion, and that systemic delivery of anti-IL-1beta antibodies (Ab) can markedly reduce neutrophil activation during thrombogenesis. There is currently no therapeutic approach to specifically target neutrophils, and systemic IL-1beta inhibition is limited by suppression of other innate immune functions essential in host-defense. This represents a critical roadblock in the translation of our findings into the clinic for patients with VTE.

Our goal in this Falk Catalyst proposal is to overcome this current roadblock by developing a synthetic bio-functional molecule to precisely target neutrophils for inhibition during thrombogenesis. This proposal builds on an existing collaboration between experts in venous thrombo-inflammation and vascular-targeting drug carrier design to develop a neutrophil-tropic particle that directs to the inflamed neutrophil and arrests thrombosis expansion. The Aims outlined here will optimize our novel nanoparticles to exclusively target to neutrophils (Aim 1, Particle optimization); and determine whether this approach effectively reduces in vivo and in vitro neutrophil activation and venous thrombogenesis (Aim 2, Efficacy). Our proposal is driven by our innovative discoveries, expert cross-disciplinary collaboration, and strong preliminary data, with the long-term objective of developing a neutrophil-inhibiting ‘precision’ therapy for patients who suffer from venous thromboembolism. The results of these studies will provide a solid rationale for us to translate our discoveries during the Transformational phase of this Award, and form the basis of an innovative and more complete strategy to treat one of the leading cardiovascular diseases.

The University of Michigan has a unique history of organizing around research questions across disciplines, providing a robust foundation for the Catalyst and Transformational phases of this program.
Major depressive disorder (MDD) is among the leading causes of illness and disability worldwide. It is a severe and life-threatening disease. Childhood trauma and insulin resistance are known risk factors for MDD. The pathophysiology of MDD remains poorly understood, and there is a serious dearth of new druggable biological targets to guide the development of improved therapeutics. This proposal outlines a research plan to identify a fundamentally new drug therapy for MDD.

In rodent models of depression, the biomolecule acetyl-carnitine promotes a rapid antidepressant response. Endogenous acetyl-carnitine levels are also lower in the plasma of patients with MDD compared with age- and sex-matched healthy controls. Several older studies involving human subjects have also reported neurophysiological and neuropharmacological effects of oral acetyl-carnitine, although these studies were either uncontrolled or under-powered. While not a typical drug-like substance, acetyl-carnitine has properties that make it a promising lead for our purposes. It is classified as a Generally Regarded as Safe (GRAS) substance by the FDA, and is orally bioavailable, while also being transported across the blood-brain barrier.

Our goals for this Catalyst Research Award are to assemble a multidisciplinary team of researchers with relevant backgrounds in chemistry, neuropharmacology, and psychiatry to launch a comprehensive drug discovery effort aimed at optimizing the activity of acetyl-carnitine in MDD. Our technical aims are to: (1) Implement robust in vitro and in vivo assays based on our model for the mode of action of acetyl-carnitine in MDD; (2) Design and synthesize acetyl-carnitine analogs, and test them in the above assays; and (3) Compare the activity of the most promising analog versus acetyl-carnitine in a rodent model of depression. If successful, these team-building and technical achievements will pave the way for a more intensive pursuit of a safe and effective drug candidate for MDD therapy under the Transformational Research Award Program.
Infectious diseases caused by bacteria, viruses, fungi, and single-cell parasites kill millions of people worldwide, each year. Although these microorganisms are very different in many ways, the first step of infection with each of them is microbial adherence to human or host tissues. Regardless of the microorganism, adherence essentially always involves binding between a protein on the surface of the microbe and a carbohydrate ligand on the host. Without this binding, microbes cannot infect host tissues and cause disease. It is clear that blocking microbial adherence mechanisms shared among microbes would be a powerful and nearly universal way to treat or prevent infectious diseases. However, no conserved aspects of microbial adherence have ever been identified and there are no proposed strategies to target this type of dangerous microbial behavior.

We have identified a conserved amino acid motif (CAMo-1) with an associated molecular feature in a large number of adherence proteins from bacteria, viruses, fungi, and protozoa. In our preliminary work, we identified this novel drug target and validated it experimentally by showing that a host protein binding to this structure blocks microbial adherence to human cells. The goal of this study is now to identify and develop small molecules that bind to this structural feature and block microbial adherence to host cells for treatment and prevention of these infectious diseases. We hypothesize that we can identify specific small molecules that bind to microbial adherence proteins at CAMo-1 to block the attachment of pathogenic microbes to human tissues. We will test this hypothesis using three different screening approaches as outlined in our specific aims: Aim #1) Use purified recombinant proteins to identify small molecules that bind to microbial adherence proteins at CAMo-1. Aim #2) Use bead-based microbial mimics to identify small molecules that prevent microbial adherence protein binding to host cells. Aim #3) Use live microbes to identify small molecules that prevent microbe binding to host cells. In each of these aims we will be performing primary and secondary screens using diversity and fragment small molecule libraries. Successful completion of this study would provide small molecule leads for development of a completely new type of antimicrobial drug designed to target dangerous microbial behaviors, regardless of the microorganism. This would be a paradigm shift in how these diseases are conceptualized and treated and a major advance towards our long-term objective to provide cures for infectious diseases in which no cures currently exist.
Diabetic patients often develop neuropathies and have a greater risk for vascular or Alzheimer’s dementia compared to those without this disease. The epidemic of these diseases indicates a need to improve the treatment of diabetes and reduce neurodegeneration. Patients with diabetes must use increasingly higher doses of therapeutics over time, which increases the risk for side effects. These side-effects are partly a consequence of the fact that insulin is a growth factor that recruits glucose for anabolic processes. Consequently, diabetic patients treated with insulin are prone to weight gain and dyslipidemia that increases the risk of cardiovascular mortality. In this work, a series of short peptides are being designed and synthesized, which undergo self-assembly into nanofibers upon exposure to physiological conditions. The positive charge on the nanofibers electrostatically binds negatively charged molecules, including insulin, and can concomitantly bind to critical receptors, i.e. LepR, to stimulate glucose uptake as a cytokine. One peptide, AAC2, alone and bound to insulin, rescued mice from T1D and T2D in preliminary studies. These preliminary data provide the basis for a central hypothesis/question to be tested in this work: Can these peptides improve glucose tolerance and reduce neurodegeneration related to diabetes, based on their interaction with LepR? The overall goal is to develop and optimize this class of therapeutics with dual properties: (1) as a scaffold to bind and stabilize hormones, such as insulin, and (2) as a concomitant activator of LepR to improve glucose uptake, energy balance, and to reduce associated neurological complications. The specific aims of this work are: (1) to optimize the structure of AAC, (2) to modulate the length and stability of the AAC nanofibers, (3) to examine the glycemic and antioxidant effects of AAC variants in vitro, and a future aim will be to (4) to determine the pharmacokinetic and pharmacodynamic profile of the candidate AAC molecule in Akita mouse models of T1D and high-fat diet induced model of T2D.
Kawasaki Disease (KD) is the leading cause of acquired heart disease in children in developed nations. KD can result in coronary artery aneurysms that can lead to lifelong heart disease, myocardial infarction, and death. The clinical and epidemiologic features support an infectious etiology in genetically susceptible children, but the cause has eluded more than 50 years of study. Delayed and missed diagnoses increase the risk of coronary artery aneurysms. The development of urgently needed diagnostic tests and improved therapies are dependent upon identifying the etiology. In this study, we propose to determine if a new human hepacivirus is an etiologic agent of KD. Recent studies have shown that 1-2 weeks after infection with any specific pathogen, 75% of peripheral blood plasmablasts target that specific infection. In preliminary studies, we analyzed the peripheral blood plasmablast response at 1-3 weeks after KD fever onset using single cell RT-PCR and made 61 monoclonal antibodies (Mab) from these plasmablasts. We used these Mab to determine their target antigens. We found that 33/61 Mab, derived from 9/11 KD patients, identify intracytoplasmic virus-like inclusion bodies (ICI) in ciliated bronchial epithelium of KD children but not infant controls. Using a viral peptide discovery array and/or ELISA, we found that 6 of the 33 (18%) Mab, derived from 3 KD patients with coronary artery aneurysms, recognize multiple similar peptides of hepacivirus non-structural protein 4A (NS4A). An optimized NS4A peptide completely blocks binding of these Mab to KD ICI, indicating the presence of a hepacivirus-like protein in the ICI. We hypothesize that at least a subset of KD cases are due to a previously unidentified hepacivirus. To test this hypothesis, we will identify the KD-associated hepacivirus using a specifically designed pathway for bioinformatics analysis of our KD tissue RNAseq dataset containing >4 billion reads. In addition, we will obtain additional viral proteome data by screening viral peptide arrays/phage display libraries with KD Mab that bind to KD ICI but do not recognize NS4A. We will test KD patients and childhood controls for serologic response to and presence of KD-associated hepacivirus. These studies will lead to improved diagnosis and treatment of KD, enable prevention, and reduce healthcare costs from the long-term consequences of coronary artery aneurysms arising in young childhood.
Melanoma, the deadliest skin cancer, was diagnosed in approximately 90,000 Americans and led to over 9,000 deaths in 2019. While BRAF inhibitors and checkpoint inhibitors have revolutionized treatment of metastatic melanoma, these therapies are limited by rapid development of resistance and low response rates, respectively. New therapies, and entirely novel drug targets, are therefore critically needed, ideally targeting not only melanomas but also congenital giant nevi that are precursors to melanomas and are currently removed by repeated surgery. An innovative approach originally developed by Prof. Craig Crews at Yale University, leverages molecules called proteolysis targeting chimeras, or PROTACs, to degrade – rather than inhibit – previously “undruggable” classes of proteins, including transcription factors. PROTACs also have the potential to enhance immunotherapy because the proteolytic peptides arising from target protein degradation are immunogenic. In this work, we propose development of PROTACs targeting a melanocyte-specific transcription factor, SOX10, that is required for melanoma cell, normal melanocyte and nevus cell proliferation. PROTACs consist of a ligand to the protein of interest, a flexible linker, and an E3 ubiquitin ligase ligand. We will therefore first develop a ligand to SOX10 using two high-throughput screening approaches for which preliminary feasibility has been established (Specific Aim 1). While ligand development is in progress, we will utilize a previously reported biotechnology-based strategy to demonstrate that PROTAC-mediated degradation of an engineered SOX10 fusion protein kills melanoma cells (Aim 2). Finally, we will create a small PROTAC library based on our novel SOX10 ligands, from which optimal members will be selected based on their ability to induce degradation of the transcription factor, inhibit melanoma cell, normal melanocytes and nevus cell proliferation, and eliminate melanocytes from human skin tissue in organotypic culture (Aim 3). In the future, we envision that the best PROTACs identified in this study will be further optimized and elaborated into drug candidates for testing in animal models prior to translation to the clinic. More broadly, this work will advance current cutting-edge efforts to drug transcription factors that have been refractory to traditional inhibitor development.
ALS patients desperately need new approaches and directions to combat this disorder. While motor neuron loss in the spinal cord has been a central area of research focus for this disease, there is extensive pathology in ALS that spans from the neuromuscular synapses to the spinal cord, to the lateral corticospinal tract, and to the cerebral cortex. It is still not known how and where this disease begins nor the molecular mechanisms of how the disease progresses once it starts.

This application represents an innovative and uniquely humanized, collaborative approach for human ALS. It will combine clinical and basic research on rapidly-acquired human postmortem tissues to identify and validate new targets for drug development using a novel injury based model of disease progression. While gene mutations in animal models that rely on rarer forms of the disease with distinct genetic abnormalities have been used extensively, these animal models have not yielded effective therapeutics. The unique aspect here is that we instead focus on targeting disease progression once the disease has already begun, as it does when our patients first present with their disease. Our central hypothesis to be tested here is that the degree of pathological change will correlate with a distinct group of genes, proteins, and biological pathways that will lead us to novel drug targets of disease progression. Currently, we have leads to suggest that inflammation is key to this as well as a custom designed, patented fusion protein that targets this. Targeting disease progression offers a ‘final common pathway’ approach to treat ALL ALS patients, not just those with rare genetic abnormalities.

The catalyst phase will bring together a diverse team for genomic analysis and mining of our novel tissue repository. Our group has pioneered human tissue functional genomics, cellular predictions, biomarker identification, and drug target identification from human tissues. In parallel, it will build a new animal model of ALS disease progression by bringing together a multidisciplinary team to test targets generated from the human tissue data. These parallel teams will come together for the transformational phase where our top drug targets will be translated into small molecular and/or biologic therapeutics to test on the new animal model of ALS disease progression. In this later phase, our goal will be to select our top three drugs for preclinical testing, toxicology, and filing of IND applications.
Disorders affecting urea metabolism are among the most common inborn errors of metabolism in the liver. Defects in metabolism of waste nitrogen from the breakdown of protein and other nitrogen-containing molecules lead to elevated blood ammonium level that causes neurotoxicity and can be fatal. Currently, low protein diets and liver transplantation are the only available therapies for disorders of urea metabolism. Procurement of cadaveric livers or use of hepatocytes/stem cell-based replacement therapies, has remained a significant challenge, driving intense research towards developing alternative liver regeneration strategies for human application. Our own current focus leverages a “self-condensation” culturing methodology wherein human induced pluripotent stem cells (hiPSC) are developmentally specified and together with vascular progenitors, self-organized into 3-D vascularized miniature livers (“organoids”). Our success with this approach is highlighted by rescue of a mouse model of liver disease using these cultured hiPSC-derived human liver organoid (hiPSC-HLO) transplants.

More recently we showed that hiPSC-HLO can be produced at a scale and purity for detailed testing in animal models. Building on this recent success, we now propose to conduct a preclinical study for the treatment of newborns who have congenital rare urea cycle disorders, more specifically, ornithine transcarbamylase deficiency (OTCD). The initial goal is to provide bridge therapies during the waiting period for liver transplants when surgery is not yet feasible and a liver not available. Our interim goals are (1) To determine if hiPSC-HLO transplants provide pivotal efficacy in alleviating urea cycle disease in our rodent models, and (2) To assure safety without tumorigenic complications by defining critical quality attributes (CQAs) in the hiPSC-HLO manufacturing process. The proposed analyses utilizing urea cycle disorder mice (generated under a severely immunocompromised background) will make it possible to determine the efficacy for correcting hyperammonemia, survival and behavioral deficits. Concurrent monitoring of tumor formation will ensure safety of our proposed organoid based approach.

In addition to providing preclinical assessment of utility of hiPSC-HLO in the OTCD model, our proposal has broader implications for efficacy in multiple hepatic dysfunctions related to protein metabolism, bile acid synthesis and export, and coagulation factor synthesis. This will expand the utility of our methodology for many other clinical indications such as liver cirrhosis, which is a significant cause of global health burden with more than one million deaths per annum.
Ventilatory support, including oxygen supplementation, has saved countless premature infants, yet these therapies have also led to bronchopulmonary dysplasia (BPD) in premature babies, which can persist into adolescence and adulthood. The pathology of this disease is characterized by alveolar and vascular simplification in the lungs. Although current therapies, including antenatal steroids and surfactant, have greatly improved survival of premature infants, the prevalence of BPD and its consequences on the lung have not been reduced. My long-term objective is to develop effective drugs to prevent or treat BPD.

Carnitine palmitoyltransferase 1 (Cpt1), a rate-limiting enzyme of the carnitine shuttle system for beta-oxidation during fatty acid oxidation (FAO), may be key for developing these drugs. Cpt1 has three isoforms: Cpt1a, Cpt1b and Cpt1c, of which Cpt1a has acyltransferase activity and a high affinity to carnitine to generate acylcarnitine for mitochondrial transport. My previous studies showed for the first time that hyperoxic exposure reduces levels of Cpt1a and FAO in neonatal lung endothelial cells (ECs), leading to apoptosis. Pharmacological inhibition or genetic deletion of Cpt1a aggravates hyperoxia-induced alveolar and vascular simplification, characteristics of BPD, in neonatal mouse model of BPD. This proof-of-concept study suggests that Cpt1a reduction causes hyperoxic lung injury in neonates. Nevertheless, whether enhancing Cpt1a attenuates neonatal hyperoxic lung injury remains unclear.

We hypothesize that enhancing Cpt1a level and activity in neonatal mice ameliorates hyperoxia-induced persisting lung injury into adulthood. To test this hypothesis, we propose two Specific Aims using pharmacological activators (i.e., C89b and L-carnitine) and genetic approaches (i.e., Cpt1a gene knockout and overexpression) to determine their impact on lung injury and repair.

Aim 1: Determine whether overexpression and activation of Cpt1a reduces hyperoxia-induced lung EC dysfunction in vitro. We will genetically overexpress and pharmacologically activate Cpt1 to determine whether this attenuates hyperoxia-induced lung EC dysfunction, including apoptosis, reduced proliferation, migration, and angiogenesis.

Aim 2: Validate whether targeting Cpt1a protects against hyperoxia-induced persisting lung injury in mice. We will expose neonatal mice (<12 h old) to different concentrations of hyperoxia for 3 days, and allow them to recover in air until adulthood. These mice will be treated with L-carnitine or C89b, or Cpt1a plasmids will be transfected into lungs to determine whether this ameliorates hyperoxia-induced lung function decline as well as alveolar and vascular simplification.

These proposed studies will set the groundwork for Cpt1a as a novel therapeutic target for preventing BPD, potentially impacting tens of thousands of lives.
Towards Predicting and Preventing the Development of Severe Dengue

Dengue virus (DENV) is a global health threat for which no effective vaccines or approved antivirals exist. 5-20% of symptomatic patients progress to severe dengue (SD), manifested by complications and sometimes death. Early administration of supportive care reduces mortality, however, there are no accurate means to predict which patients will progress to SD. Our goal is to better understand the virus-host interplay involved in SD pathogenesis in order to develop prognostic tools for early identification of patients at risk for progression to SD and host-targeted antivirals.

We established a unique cohort in Colombia—dengue patients who present prior to progressing to SD. Moreover, we developed a novel virus-inclusive, single-cell transcriptomic (viscRNA-seq) platform, which transforms our ability to monitor host gene expression dynamics with viral abundance in thousands of individual cells. Additionally, we used a novel multi-cohort analysis of the publicly available gene expression data sets to identify and validate a 20-gene set predictive of SD. This discovery is groundbreaking since this gene set is generalizable across a variety of countries and ages. Lastly, we demonstrated a proof-of-concept for the utility of host-targeted approaches to combat DENV. We hypothesize that combining viscRNA-seq profiling in our dengue cohort samples and the multi-cohort analysis will identify biomarkers of SD whose mechanistic validation can also reveal host targets for antiviral therapy.

In Aim 1, we will conduct viscRNA-seq analysis in longitudinal PBMC samples from the Colombia cohort to map an atlas of DENV cellular targets and identify candidate biomarkers predictive of SD. To functionally validate these biomarkers and identify novel druggable host functions for antiviral therapy, we will decipher the roles of prioritized top hits identified in our viscRNA-seq studies in SD pathogenesis and the DENV life cycle.

Aim 2 will determine the feasibility and biological rationale for predicting SD by the promising prognostic 20-gene set. We will validate this gene set in silico and prospectively, monitor its dynamics during the disease course, and define its specificity. Lastly, we will decipher the roles of some of the 20-gene products in SD pathogenesis and the DENV life cycle.

This bold, interdisciplinary proposal will further transformative single-cell and bioinformatics technologies, provide insights into SD pathogenesis with an unprecedented resolution, and yield generalizable gene sets predictive of SD and host functions as candidate antiviral targets. By advancing the development of dengue prognostic assays and antiviral strategies (years 2-3), this work has the potential to improve human health.
“Development of a Novel Treatment for Organophosphorus Chemical Nerve Agent Poisoning”

Many organophosphorus (OP) compounds are chemical warfare agents or pesticides and are significant threats to human society as over 200,000 people die every year from OP exposure. These chemical agents inhibit the enzyme acetylcholinesterase (AChE) in the human body, and upon exposure to OP pesticides, AChE undergoes an aging process, rendering current oxime treatment to be ineffective. Our team has recently demonstrated the first small drug-like molecules that can reverse the effects of aging for AChE in vitro, thereby fully reversing the effects of OP exposure. The objective for this proposal is to expand on these alkylating compounds to improve selectivity, binding and reactivity with the aged forms of AChE in order to form stable adducts which can then be reactivated, and while doing so, enhance the drug-like properties for improvements in dosages and time for effective treatment. Thus, this proposal will improve the efficacy of these compounds in order to develop new therapeutics against the toxicological effect for the global use of OP pesticides.
T- and NK-cell lymphomas comprise 10% of non-Hodgkin’s lymphomas in Western countries. Specifically, advanced cutaneous T-cell lymphoma (CTCL) and peripheral T-cell lymphoma (PTCL) have a dismal survival. To date their treatment has been largely derivative or empiric, with few biomarkers to guide therapeutic selection. Three fundamental barriers have prevented successful translation in TCL: 1) the low incidence of each TCL subtype, which limits sample availability and clinical trial enrollment; 2) heterogeneity across subtypes, which further complicates biologic interrogation; and 3) a lack of faithful model systems for in vitro and in vivo studies.

To directly address the need for patient-specific biomarkers that guide therapeutic selection, we have developed a rapid parallel in vivo assay that consists of implantable microdevices (MDs) placed inside the native tumor microenvironment. Each MD contains up to 20 reservoirs, which are individually loaded with a unique agent or drug combination. It is implanted directly into tumor and remains in situ for ~24-72h. Following implantation, drugs from each reservoir are released into distinct regions of tumor tissue. A coring needle is used to retrieve the MD with surrounding tissue. Native tumor tissue surrounding MDs can be analyzed by multi-parameter immunohistochemistry or immunofluorescence, RNAseq, or other approaches to determine individual drug effects. This completely new paradigm for discerning targeted tumor-specific drug effects allows immediate readouts of drug activity directly from the affected tissue. The key advantage of this approach is the high throughput of targeted perturbations induced by released drugs within the native tumor architecture, which, through precise spatial overlay, provides a combined measurement of drug release and phenotypic readout of cell/tissue response, across a large set of drugs relevant to the treatment of TCL.

Application of this microdevice, to determine the optimal treatment for each patient, could be transformative for the care of patients with TCLs. We will test the MD in situ within tumors, a process that has never been undertaken in lymphoma models. The results of this study will serve as the basis for registering the MD with the FDA to allow investigation in patients with TCL and as preclinical data to define mechanisms of response and resistance for multiple novel agents. We propose the following Specific Aims.

Aim 1. Define the activity of FDA-approved and investigational drugs across TCL models in vivo.
Aim 2. Validate methods for multiparametric assessment of TCL biopsies with embedded MDs.
Aim 3. Finalize planning for clinical trial sample analysis.
Schizophrenia, autism spectrum disorder, bipolar disease, and epilepsy are genetically related and highly debilitating neurodevelopmental disorders that affect >3% of the world population. Current therapies for these diseases do not ameliorate all symptoms and have not improved in efficacy in decades. Our goal is to develop better drugs to treat these disorders.

TRIO is among the genes most frequently impacted by heterozygous rare damaging genetic variants in individuals with schizophrenia, autism, and related disorders. TRIO contains two guanine nucleotide exchange factor (GEF1 and GEF2) domains which catalyze GTP exchange on the Rac1 and RhoA GTPases, respectively, to regulate synapset development and function. Most of the disorder-associated mutations in TRIO are heterozygous and disrupt the GEF1 and/or GEF2 activities. Deletion of one TRIO allele in mice compromises normal brain development and function. Together, these data strongly implicate reduced TRIO function as a causative factor in neurodevelopmental disorders.

We propose to test the hypothesis that boosting TRIO’s enzymatic activities might lead to therapeutic benefits in individuals suffering from SCZ, ASD, and related disorders. Specifically, we propose to identify small molecules that can modulate TRIO’s GEF1 and GEF2 activities and to test their ability to rescue phenotypes arising from TRIO haploinsufficiency.

Aim 1. To screen for small molecule modulators of TRIO GEF1 and GEF2 activities. We will use a fluorescence-based, 384-well plate assay that measures the TRIO GEF1-mediated exchange of GTP for GDP on Rac1 to implement a pilot high throughput screen of an approximately 5,000 compound library enriched with CNS-drug like properties. We will perform a similar pilot screen on TRIO GEF2 activity on RhoA. These screens will identify both positive and negative regulators, and, because both GEF domains will be screened against the same library, hit compounds with initial selectivity will be identified and prioritized. We will validate lead compounds in cells using genetically-encoded Rac1 or RhoA FRET biosensors.

Aim 2. To test the ability of TRIO GEF modulators to rescue defects in TRIO+/- neurons. Cortical neurons and slices from TRIO+/- mice exhibit defects in dendrite structure and synaptic structure, transmission, and plasticity. We will apply our battery of quantitative cellular, anatomical, and electrophysiological assays to probe whether validated activators from the initial screen can remediate structural and functional defects in TRIO+/- neurons. Active hits from Aim 2 will ultimately be advanced into a drug discovery cascade for advancement toward a clinical candidate in future studies.
Atherosclerosis is the process underlying heart attack and stroke. Despite recent advances, atherosclerotic cardiovascular disease (CVD) remains the leading cause of death in the United States. Most current therapies are directed against cardiovascular risk factors (such as hypertension and elevated cholesterol levels). However, much of the population’s risk of developing disease occurs independently of traditional risk factors. Therapies that directly target the plaque would instead address the root cause of disease and have the potential to fundamentally transform how CVD is treated.

A characteristic feature of the atherosclerotic plaque is the pathological accumulation of diseased and dying cells in the necrotic core. We discovered that this phenomenon is driven by the marked upregulation of a key ‘don’t eat me’ molecule known as CD47. This renders vascular cells ‘inedible’ and resistant to ‘efferocytosis’ (programmed cell removal). We showed that systemic delivery of anti-CD47 antibodies (Ab) could reactivate efferocytosis within the lesion, thus dramatically reducing plaque development and vascular inflammation. However, systemic antibody-based therapy also caused off-target clearance of red blood cells. This induces an anemia which represents a critical roadblock in the translation of our findings into the clinic for CVD patients.

The broad, long-term objective of our Falk Catalyst proposal is to overcome this current roadblock by developing a ‘precision’ nanotherapy that delivers anti-CD47 therapy specifically to the plaque. We are collaborating with experts in nanomedicine, bioengineering, and immune cell biology to develop a vascular-tropic nanoparticle that homes to the inflamed macrophage and reactivates phagocytosis in the plaque, thereby preventing atherosclerosis without off-target toxicity. The aims outlined in this Catalyst Award proposal will determine whether our pro-efferocytic nanoparticles specifically accumulate in the macrophage-rich plaque and prevent atherosclerosis (Aim 1, Efficacy), and validate that this approach effectively avoids the off-target clearance of healthy tissue seen with systemic anti-CD47 therapy (Aim 2, Safety). Our proposal stems from our major innovative discovery and promising preliminary data, with the long-term goal of developing an efferocytosis-activating ‘precision’ therapy for patients who suffer from atherosclerotic CVD. The results of these studies will allow us to move towards translating our discovery into humans during the Transformational Award phase, and form the basis of a completely new strategy to prevent or even cure the leading cause of death in the United States.
Despite current medical treatment, a large number of patients with asthma develop chronic refractory asthma with long-term disability. Innovative approaches for treatment of asthma are needed to reduce the economic and social burden. People become asthmatic through the process known as ‘sensitization’ against specific allergens such as house dust mites. Dendritic cells (DCs) play a key role in this sensitization process. DCs are able to recognize allergens, and present the immune determinants of an allergen to T and B cells to generate immunoglobulin E (IgE). Thus, the allergen-reactive IgE is a hallmark for systemic sensitization of the allergen. Upon re-exposure to the already sensitized allergens, DCs capture the re-entered allergen and present the allergen to memory T and B cells, which induce a robust allergic inflammation resulting in the clinical symptoms of asthma. This indicates that DCs play an important role not only in the initial sensitization, but also exacerbation of asthma in already sensitized individuals. Intervention of DC function in this process could create a novel therapeutic opportunity for treating asthma at very early steps of allergic inflammation by blocking allergen sensitization. We found that the allergen-exposed airway epithelial cells secrete Colony Stimulating Factor-1 (CSF1) into airways where they bind to its receptor (CSF1R) on recruited dendritic cells in airways. The binding between CSF1 (ligand) and CSF1R (receptor) enhances the expression of CCR7 on DCs. CCR7 is the key molecule for DC migration and subsequent antigen presentation. We also confirmed that the small molecule inhibitor of CSF1R successfully inhibited CCR7 expression on DCs and subsequent specific IgE production. Based on this novel immunologic mechanism of allergen sensitization, we hypothesized that the inhibition of the CSF1-CSF1R pathway has a therapeutic benefit for asthma treatment. To examine the effectiveness of CSF1R inhibitor in the treatment of asthma, we have synthesized the encapsulated nanoparticle carrying CSF1R inhibitors. This particle can be delivered to the lung via inhalation or intranasal delivery. Before moving forward with a clinical trial, we will examine the effectiveness of the nanoparticle of CSF1R inhibitor and look for possible adverse reactions in an animal model of asthma. This proposal is based on our strong scientific premise that the CSF1-CSF1R pathway plays a critical role in allergen sensitization and development of subsequent allergic inflammation. The proposed experiment will set the groundwork for the potential use of CSF1R inhibitor in clinical practice.
Sepsis is a syndrome characterized by a maladaptive immune response to infection that leads to hypotension, organ dysfunction, and death. It is a common disease with an exceedingly high fatality rate and was deemed a global health priority by the World Health Organization in 2017. Despite decades of clinical and basic science research, there are no effective treatment options for sepsis other than early administration of antibiotics and supportive care. A growing body of research has illuminated the critical role of the inflammation-induced programmed cell death pathway necroptosis in the pathogenesis of sepsis. Necroptotic cell corpses are inflammatory, particularly towards myeloid cells, which can drive a vicious cycle of increased myeloid cell activation, cytokine production, and further necroptosis that sustains sepsis pathophysiology. However, the mechanism by which necroptotic corpses promote and sustain inflammation remains unresolved. To address this question, we recently identified a family of cell surface receptors that specifically bind to the surface of necroptotic cells, but not viable or apoptotic cells. To interrogate the function of these receptors and their potential contribution to the pathophysiology of sepsis, we generated recombinant decoy proteins as pharmacologic antagonists of the receptors. In vitro, these antagonists attenuated the effect of necroptotic cells to activate inflammatory signaling in macrophages. Furthermore, these antagonists dramatically improved survival in an in vivo experimental endotoxemia sepsis model. This effect was evident even when the drugs were administered well after the acute inflammatory phase; by contrast, most other experimental agents are ineffective at this time point and work only when given before or immediately after the initiation of sepsis. Thus, we have uncovered a new therapeutic strategy to break the "vicious cycle" of inflammation in sepsis by "shrouding" necroptotic corpses from immune recognition with novel therapeutic candidate drugs. The goal of our proposal is to understand the biology of these molecular sensors of necroptotic corpses, to identify their ligand(s), to understand how our agents protect animals from sepsis, and to conduct key proof-of-concept studies of the efficacy of the proteins in preparation for an Investigational New Drug (IND) application to enable clinical trials. Our efforts will therefore shed new light on the biological mechanisms of sepsis and will lay the foundation for transformative therapeutic approaches for this deadly disease.
Approximately 900,000 people in the United States are affected by ulcerative colitis (UC) and patient numbers increase yearly. While its etiology is unknown, symptoms of UC typically include chronic inflammation of the large intestines and are accompanied by numerous complications. Medical costs range from $4-14 billion annually within the United States and treatments range from oral medications to injectable agents that circulate widely throughout the body. As these treatments eventually fail, surgery is performed to remove parts of the diseased tissue. One-third of those with UC will require surgery during their lifetime, drastically affecting quality of life. The current regimen of injectable drugs to treat UC has multiple side-effects, including increased risk of certain cancers. Hence, there is still an important unmet clinical need for safe and effective treatments. Our proposed innovation is to engineer potent anti-inflammatory agents in the form of supramolecular anti-inflammatory peptide amphiphiles (AIF-PAs). The AIF-PAs will target inflamed areas in the lower gastrointestinal tract so that a high proportion of the treatment goes directly to the affected area. Hence, we propose that the non-invasive delivery of AIF-PAs will attenuate inflammation in a pre-clinical mouse model of UC that closely mimics the human condition while simultaneously promoting wound healing. In order to validate our hypothesis, we will first synthesize, characterize, and optimize supramolecular AIF-PAs that can be targeted to the site of lower gastrointestinal tract inflammation and deliver effective anti-inflammatory agents (Specific Aim 1, months 1-3). We will then evaluate the efficacy of AIF-PAs to inhibit lower gastrointestinal tract inflammation using a targeted delivery platform in an established mouse model of UC (Specific Aim 2, months 4-12). Initial data gleaned from these studies will set the foundation to initiate clinical studies that will meet our long-term objectives. The ultimate goal of this study is to create a product that can be successfully utilized in the clinical arena to treat those afflicted with varying degrees of UC. In order to meet this criteria, our AIF-PAs, in-part, need to be: 1) able to reduce lower gastrointestinal inflammation with minimal dosing, 2) synthetically reproducible on a large scale at clinical grade levels for patient delivery, 3) non-immunogenic and non-toxic, 4) biodegradable, and 5) biocompatible in humans. These issues need to be addressed in subsequent years of funding.
Antibiotic resistance is a growing health threat worldwide. Although microbes evolve resistance mechanisms inherently, resistance can be exacerbated by prophylactic and systemic antibiotic administration and lengthy exposures to antibiotics at sub-inhibitory concentrations. Localized drug delivery has the potential to provide rapid antibacterial therapy at the site of an infection, while preventing offsite toxicity and reducing susceptibility to resistance. However, most currently existing local antibiotic delivery devices are plagued by low drug loadings and release below inhibitory concentrations. We propose to develop a library of bacteria-responsive polymeric building blocks that can be used to formulate biomaterials for triggered antibiotic delivery at the site of an infection. Responsive biomaterials undergo a change in properties under specific stimuli, which can be coupled with the release of encapsulated therapeutics. Existing infection responsive materials rely on non-specific triggers, such as pH, to trigger drug release. In the proposed work, we will utilize beta-lactamases, which are bacteria produced enzymes, to serve as the bacteria-specific drug release trigger. Beta-lactamases hydrolyze the beta-lactam ring in beta-lactam antibiotics (among the most commonly prescribed drugs in the world) and are the most prevalent cause of resistance to these antibiotics. They are produced by many bacteria including the ESKAPE pathogens (Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa, and Enterobacter species) that cause nosocomial infections and commonly develop resistance. In the first aim of this project, we will synthesize polymer-beta-lactam conjugates using previously developed beta-lactam molecules. Synthetic and naturally-derived biocompatible polymers including poly(ethylene glycol), alginate, hyaluronic acid, and gellan will be explored. This polymer-beta-lactam library will be thoroughly characterized for beta-lactamase triggered degradation. In the second aim, we will develop a range of bacteria-responsive biomaterials using these building blocks. The materials will vary in size from nanoparticles for injectable therapies to micro-scale self-assembled films for device coatings to macro-scale hydrogels for antibacterial wound dressings. Antibiotics effective against ESKAPE pathogens will be incorporated into these materials. Biomaterial physical properties and beta-lactamase responsiveness will be assessed. In the final aim, we will investigate the in vitro antibacterial efficacy of these responsive materials against the ESKAPE pathogens. Results from this study will provide a strong basis for application to the Falk Transformational Awards program, in which we will optimize these materials for use in preclinical infection models, examine in vivo efficacy and biocompatibility, and evaluate antibiotic resistance development in the presence of these materials.
Premature termination codons (PTCs) are genetic mutations that result in truncated and inactive proteins. The World Health Organization estimates that over 10,000 human diseases are caused by mutations within a single gene, and 10-25% of these are believed to be caused by PTCs. While individually rare, collectively genetic diseases caused by PTCs affect many patients and their families and represent a significant burden to the health care system. Many of these diseases result in severe developmental or neurological defects or are lethal in infancy and early childhood, and almost none have effective treatments or cures. Several treatment and curative modalities for PTCs have been proposed. Suppressor tRNAs are engineered tRNAs that recognize the termination codon but add the appropriate amino acid instead of terminating the chain, enabling read-through of the PTC and the producing the functional protein. CRISPR-Cas9 guided single base editors can change individual base pairs within DNA, correcting the underlying mutations and potentially curing these diseases. Both approaches have been effective in vitro but are limited by the lack of an in vivo-compatible platform to deliver these therapies to tissues within patients. The objective of this study is to determine whether our novel, rationally engineered nanoscale PrOtein Delivery (nanoPOD) platform can deliver suppressor tRNAs (Aim 1) or CRISPR-Cas9 guided single base editors (Aim 2) in vivo and enable read-through or correction of PTCs. We will employ a powerful mouse model that contains a DNA sequence – including a PTC – from patients suffering from glycogen storage disease type 1a (GSD1a) upstream of red-shifted luciferase (akaLuc) and fluorescent reporter (Venus) proteins. If read-through or correction of the patient-derived PTC occurs, these enzymes are produced and are detectable using whole-animal bioluminescence or cryofluorescence imaging, respectively, allowing precise determination of on- and off-target delivery of our therapeutics. We will also characterize the kinetics and duration of treatment as well as monitor toxicity and immunogenicity of the nanoPOD platform. Finally, in Aim 3, we will develop a transgenic mouse model of GSD1a that incorporates the patient-derived PTC that will be used to evaluate the efficacy of our interventions in vivo to prepare for a potential Falk Transformational Award. These animals will be shared with the research community to stimulate research into GSD1a treatments and cures. The successful completion of this will lay the foundation for transformative treatments and cures for GSD1a and other devastating genetic diseases caused by PTCs.
Approximately 70% of breast cancers express estrogen receptor alpha (ERα) and are, therefore, treated with endocrine therapies. Although many patients benefit from Tamoxifen or aromatase inhibitors (AIs) in adjuvant and metastatic settings, approximately 50% of responsive tumors eventually relapse due to the development of resistance. One emerging mechanism of resistance is the clonal evolution of mutations in a “hotspot” within the ligand-binding domain (LBD) of ESR1, the gene encoding ERα. ESR1 mutations occur in 10-20% of patients with metastatic ERα-positive disease who received endocrine therapies. The mutations led to ligand-independent ERα activity that promotes tumor growth and metastasis, and reduced efficacy of ERα antagonists resulting in endocrine resistance. The mutant ERα proteins are also more resistant to selective estrogen receptor degraders (SERDs) such as faslodex. AZD9496 and GDC-0810, two newly developed, orally bioavailable selective estrogen receptor degraders (SERDs) are in clinical trials. However, both displayed mild estrogen activity in endometrial cells and increased uteri weight in rat models, raising the concern that the mixed agonist/antagonist activity may increase risk of endometrial cancer. Thus, our goal is to identify a new class of ERα blocker to treat metastatic, ERα-expressing breast cancer.

This study builds on our recent discovery of a natural plant product, Diptoindonesin G (Dip G) that significantly decreases ERα protein levels and, importantly, is insensitive to ESR1 mutations. We showed that Dip G acts via a mechanism distinct from all known endocrine-therapy agents. Instead of binding to ERα, the direct target of Dip G is CHIP/STUB1, an E3 ubiquitin ligase that controls ERα stability. Consequently, Dip G can degrade mutant ERα more effectively than faslodex within the therapeutic window. We hypothesize that Dip G and its analogues will be effective for treating mutant ERα expressing, endocrine-resistant tumors. This application will focus on the synthesis of novel Dip G analogues to determine the basic pharmacophore of Dip G and improve the potency and pharmacological properties of Dip G (Aim 1), which will be performed by co-I Dr. Weiping Tang (School of Pharmacy, UW-Madison). The mechanism of action of Dip G will be studied (Aim 2) and the anti-cancer effects of Dip G and its analogues will be evaluated in endocrine-resistant models and compared with other SERDs (Aim 3). By targeting mutant ERα for degradation, these novel therapeutic agents will help reversing endocrine-resistance and provide a cure to metastatic patients.
“Role of DFNA5 in the Anti-Tumor Immune Response”

Tumor immunogenicity, defined as the ability of the tumor itself to trigger an anti-tumor adaptive immune response, is one of the most important determinants of successful anti-cancer therapy. The immunogenicity of a tumor depends on its antigenicity, conferred by neo-antigens generated in the course of oncogenesis, and adjuvanticity, the latter provided by damage-associated molecular patterns (DAMPs) released from stressed or dying tumor cells by a process called immunogenic cell death (ICD). Corroborating the importance of ICD in the treatment of cancer, traditional chemo- or radiotherapies that induce ICD in tumors yield a far superior and durable response to those therapies that don’t. In addition, combinatorial strategies that boost the release of DAMPs from dying tumor cells can efficiently convert non-immunogenic cell death into a bona fide ICD. Nevertheless, despite these important insights into the key role of ICD in tumor immunogenicity, there is still much to learn about the signaling and molecular pathways that induce or modulate ICD. Hence, a better understanding of the mechanistic aspects of ICD and the signaling pathways that regulate it can help us better predict the responsiveness of certain tumors to different therapies as well as reveal candidate molecular targets to enhance ICD of tumor cells and aid in cancer treatment. We have recently discovered a signaling pathway that switches apoptotic cell death into a necrotic form of cell death, allowing efficient release of DAMPs from apoptotic cells. We discovered that activation of caspase-3 during apoptosis in transformed and non-transformed cells leads to caspase-3-mediated cleavage of the gasdermin protein DFNA5 after Asp270 generating a necrotic pore-forming DFNA5-N fragment that targets the plasma membrane to induce its permeabilization. Creation of these plasma membrane pores by the necrotic DFNA5-N fragment allows the release of intracellular DAMPs such as HMGB1, DNA, and ATP. The ability of these events to switch apoptosis into a potentially immunogenic form of cell death demonstrates a novel mechanism that could explain how immunogenicity of tumor cells arise. Therefore, the experimental plan outlined in this Catalyst Award proposal will test the novel concept that the caspase-3/DFNA5 pathway constitutes one of the major checkpoints determining the immunogenicity of cancer cells. Elucidation of the role of this pathway in the regulation of tumor immunogenicity will provide a solid rationale for a Falk Transformational Award that will allow us to exploit DFNA5 as a biomarker and a target to enhance anti-cancer therapies.
Current therapies reduce the frequency and magnitude of immune attacks in multiple sclerosis (MS) but do not stimulate remyelination. Lack of remyelinating therapies is one of the major roadblocks to treat MS patients. Although the CNS contains abundant oligodendrocyte progenitors (OPCs) capable to differentiate in myelinating oligodendrocytes, most OPCs fail to remyelinate during MS. At least two factors contribute to this failure: inadequate stimuli to promote differentiation and abnormal expression of myelinating-inhibitory signals in demyelinated axons. Although numerous mechanisms regulating OPC differentiation are known, their translation into remyelinating therapies has not been achieved yet.

Our project to the Falk Foundation will address the current roadblock in remyelination therapies by taking advantage of our successful experience with promyelinating microRNAs and our experience with extracellular microvesicles (EVs). Various micro-RNAs (i.e. miR-219) have potent proremyelinating properties, constituting excellent candidates for therapy. The main difficulty is their delivery to OPCs in the CNS during disease, with minimal off-target. EVs are ideal vehicles for CNS therapy because they can be loaded with different therapeutic biologicals, elicit limited or none toxicity and immunogenicity, and are taken up by most cells.

To test our hypothesis that receptor-mediated endocytosis of EVs loaded with proremyelinating miRs can enhance remyelination in MS, our catalyst project will design and optimize a cell-free delivery system to selectively target OPCs to improve the delivery of miRs. We will produce bone-marrow derived mesenchymal stem cells engineered to produce EVs which express surface recognition proteins with high affinity binding to PDGFrα receptors in OPCs. We have chosen PDGFrα receptors as our target receptor for endocytosis of EVs because PDGFrα is highly expressed in OPCs, the target cells for remyelination and because we have identified a set of small peptides that bind to this receptor in OPCs. We show that one of these peptides (OL-2) have the highest selectivity for OPC binding, with minimal binding to other glial cells such as astrocytes and microglia. Our catalytic proposal stems from strong preliminary data, with a defined target for surface recognition, and consolidated methods to deliver a measurable product: EVs targeting OPCs.

Upon generation of effective OPC-targeting EVs and optimization of conditions for proremyelinating miR loading, the subsequent transformative phase will determine the therapeutic effects of single vs combined miRs to promote remyelination in EAE via intrathecal injections of OPC-targeting EVs. Achievement of the transformative award will produce measurable deliverables ready for consideration for IND and clinical trials.
The overarching goal of this project is to develop a small molecule inhibitor that can transiently increase recombinant adeno-associated virus (rAAV)-mediated homologous recombination (HR) levels such that a single-dose administration in neonates or adults can treat individuals suffering from life-threatening metabolic disorders.

Even though classical rAAV vectors show early promise in the clinic, there are still limitations: (1) Inability to get life-long persistence in neonates/infants/children from any growing tissue such as the liver. (2) rAAV-vector administered into neonatal mice results in high rates (>50%) of hepatocellular carcinoma because rare integration events activated a proto-oncogenic because of the strong promoter used to drive the transgene. Additionally, nuclease-mediated rAAV approaches are being considered for in vivo genome editing. This presents several problems: (1) Multiple rAVVs are required to transduce the same cell. (2) No means to limit expression or integration of the vector containing the promoter driven nuclease. (3) Off-target cutting, immunogenicity and/or insertional mutagenic risks remain.

We developed a new technology for HR that overcome the problems cited. Our nuclease-free, promoterless AAV-mediated HR technology, named GeneRideR, uses a vector containing a ribosome skipping sequence and therapeutic protein coding sequence flanked by homology arms to an endogenous gene. After HR, transcription from the endogenous gene locus produces a chimeric mRNA producing both the endogenous and therapeutic protein. This technique has been used to treat three mouse models of human genetic liver diseases. However for many diseases, the threshold for correction will require higher rates of HR. We have recently identified 20 genes that when knocked out enhance rAAV-mediated HR. Independent validation for one of these genes confirmed that in its absence, AAV-HR is 10-fold higher. We propose to: (1) continue to validate the other genes identified in cells and mice, and (2) plan a small molecule screening strategy to identify small molecules that can be used to temporarily increase AAV-mediated HR. With as little as a 5-fold increase in HR, this technology becomes widely applicable to hundreds of diseases for which there is no current treatment.

Our ultimate goal is to make the GeneRideR technology a universal plug-play therapeutic that can be used to treat a myriad of genetic diseases by simply changing the therapeutic protein coding sequence in the vector. The major advantages of this tactic compared to current gene therapy/editing approaches is that it has the potential for single administration lifelong cure in neonates, or adults, and mitigates current concerns for vector-induced cancers.
“SIRT3 as a Therapeutic Target for Therapy Eesistant B-cell Lymphomas”

Therapy resistance remains one of the most important challenges in treating patients with aggressive B-cell lymphomas. Yet little is known about the specific mechanisms that cause chemo-immunotherapy resistance in patients. Through a rationally designed screening strategy our research identified SIRT3 as a critical driver of these resistant forms of DLBCL. SIRT3 mRNA and protein expression is highly significantly and reproducibly linked to inferior outcome in DLBCL, independent of all other known risk factors including IPI, cell of origin, double hit, etc. We showed that whereas SIRT3 does not play a role in normal B-cell development, loss of SIRT3 in mice is required for B-cells to undergo malignant transformation. Our preliminary data show that SIRT3 loss of function causes proliferation arrest and cell death in almost all DLBCL cell lines. SIRT3 is also required for human DLBCL cells to engraft and form tumors in mice. Investigation into mechanism of action show that a crucial function of SIRT3 in DLBCL cells is to maintain a constant flow of Acetyl CoA to the cytoplasm, to maintain various anabolic synthetic pathways. Another key function appears to be control of oxidative stress response. Loss of SIRT3 in DLBCL cells causes them to literally “die of starvation”, manifested by massively increased autophagy, depletion of metabolic precursors, and other features. Using a rational approach we designed small molecules that selectively inhibit mitochondrial SIRT3. These molecules exert identical effects to the genetic loss of function of SIRT3. Collectively, these data lead us to hypothesize that i) SIRT3 is a critical mediator of chemotherapy resistant DLBCL, ii) that SIRT3 mediates these effects by maintaining unique aspects of DLBCL metabolism and enabling these cells to tolerate extreme metabolic and oxidative stress characteristic of these highly proliferative tumors, iii) by the same token these features enable DLBCLs to tolerate exposure to chemo-immunotherapy, iv) SIRT3 targeted therapy will potently suppress the most aggressive and resistant forms of DLBCL and v) SIRT3 expression and metabolic profiles will serve as a useful biomarker to guide SIRT3 inhibitor deployment to the clinic. The aims of our TRP will address these various points and lead to the clinical translation of an entirely novel therapeutic concept specifically geared to those patients that most urgently need improved therapies.
Our long-term goal is to deliver Novel Bacterial Type II Topoisomerase Inhibitors (NBTIs) into clinical trials for multidrug-resistant pathogens. This application proposes the design and synthesis of NBTIs targeting MRSA with a diminished resistance potential. Our central hypothesis is that potent dual inhibition of gyrase and TopoIV will reduce the spontaneous resistance rate and that a novel linker moiety will afford improved cardiovascular safety. Our synthetic route delivers new analogs in six steps, with structural diversification in the final step. Our rationale is that the synthetic efficiency and superior physicochemical properties will enable optimal dual target inhibition while preserving cardiovascular safety. These efforts will be guided by innovative x-ray crystallographic and computational homology models.

1) Design and synthesize potent dual inhibitors with reduced cardiovascular safety liability
Our working hypothesis is that optimization of the enzyme-binding moiety will improve TopoIV inhibition. Twelve analogs will be synthesized initially. Subsequent cycles of design and synthesis, guided by results from the Aims below, seek ten-fold improvements in the TopoIV/gyrase ratio of our lead compound and in the spontaneous resistance frequency. We hypothesize that cardiovascular safety issues are primarily driven through hERG inhibition and that the lower basicity and lipophilicity of the linker will minimize hERG inhibition.

2) Quantify anti-MRSA potency, propensity for resistance, and safety
Previous research supports our working hypothesis that dual inhibition will reduce spontaneous resistance and improve activity against MRSA with mutated DNA gyrase. Gyrase and TopoIV inhibition will be measured for each compound alongside Staphylococcus aureus minimum inhibitory concentrations (MICs). Spontaneous mutation frequencies will be determined for our lead compound and for an optimized analog with superior dual target inhibition. MICs against a lab-generated mutant S. aureus with resistance to our lead will be used to assess whether superior dual inhibition translates to improved whole cell mutant activity. hERG inhibition and mammalian cytotoxicity (K562 cells) will be determined for prioritized compounds.

3) Optimize TopoIV inhibition guided by computational and structural methods
Our working hypothesis is that molecular-level understanding of target binding will enable optimized target inhibition. We will determine x-ray crystal structures of NBTIs in ternary complex with DNA and gyrase as proof of feasibility, and we will express TopoIV enzyme in preparation for future crystallography efforts. We will build a homology model of the NBTI, TopoIV, and DNA ternary complex as a complementary approach and use this model to inform the rational design of analogs with improved TopoIV inhibition.
Preclinical studies of primary cancer cells are done after cells are removed from patients or animals at ambient atmospheric oxygen (~21%) yet, oxygen concentrations in organs are in the ~3-10% range, with most tumors in an hypoxic environment in vivo. While effects of oxygen tension on tumor cell characteristics in vitro have been studied, it is only after the cells were first collected in ambient air. Dr. Broxmeyer’s lab recently showed that hematopoietic stem cells exposed to ambient air within minutes undergo irreversible differentiation through a phenomenon termed extra physiologic oxygen shock/stress (EPHOSS). Therefore, results of many stem cell-related studies likely need to be interpreted with caution, and re-evaluated because the cells were first collected and processed/propagated under ambient air. With growing interest in cancer stem cells (CSC), it is crucial to determine whether current methods of collecting/studying tumor cells in ambient air influence numbers, gene expression profiles, and drug sensitivity of CSCs in tumors due to EPHOSS during tissue collection. We will address this important question using breast cancer animal models and patient-derived samples, particularly focusing on metastatic cancer cells. Studies using properly collected and processed metastatic cells are needed. Recent studies showed that metastases depend on signaling networks distinct from those of primary tumors, due to independent evolution. Drugs effective against primary tumors may be ineffective against metastases. We observed that mammary tumors from MMTV-PyMT mice collected and propagated under 3% oxygen manifest more CD61+ tumor cells compared to tumors of the same mice propagated under ambient air. CD61+ mammary tumor cells have 50-fold higher CSC activity due to enhanced CD61-KRAS-RalB-NF-κB signaling, and are resistant to receptor tyrosine kinase inhibitors (RTKis). Aim 1 will functionally characterize tumor cells collected and propagated at 3% oxygen vs. ambient air for CSC activity, gene expression profiles, and sensitivity to RTKis and chemotherapy. Aim 2 will determine whether metastatic tumor cells from patient pleural effusions and ascites fluid with and without EPHOSS show distinct CSC properties and drug sensitivity using our recently modified primary cell culturing system. Our results could provide paradigm-shifting information in the CSC field for drug-screening efforts, and provide a more rational basis to change tissue collection procedures for a truer understanding of in vivo functional characteristics of CSC/metastatic tumor cells.
Mitochondrial dysfunction is an early prominent feature in patients with neurodegenerative diseases such as Alzheimer’s (AD), Parkinson’s (PD) and Huntington’s disease (HD). Significantly, we recently reported in vitro and in vivo proof of concept that suppression of mitochondrial impairment is a useful strategy for treatment in models of these diseases. We demonstrated that improving either impaired mitochondrial dynamics or aberrant mitophagy with rationally designed peptides was protective in both neurons derived from patient induced pluripotent stem cells (iPSCs) and mouse models of these diseases. Because peptides often face challenges during drug development, we sought small molecules that increase mitochondrial function as a new therapeutic approach. The objective of this study is to optimize an existing lead molecule to enable deeper in vivo evaluation of the hypothesis that enhancing mitochondrial efficacy is a novel therapeutic strategy for neurodegenerative disease. We have recently developed and validated a series of cell-based assays in 384-well format to identify small molecules that reduce mitochondrial depolarization and bioenergetic failure as well as cell death. We focus on HD that is a fatal and inherited neurodegenerative disease with no treatment available, as a disease model. High-throughput screening identified a number of molecules, including the GSK3alpha/beta inhibitor CHIR99021, as enhancing mitochondrial function and cell viability in an HD culture model. Preliminary in vivo studies found that CHIR99021 reduces neuronal loss, behavioral deficits and animal lethality in an HD R6/2 transgenic mouse line. Notably, past reports cast doubt on GSK3 as a drug target in HD, and knockdown of GSK3alpha and GSK3beta is insufficient to enhance mitochondrial function or block cell death. Among 12 GSK3 inhibitors evaluated in dose in our in vitro assays, only one other scaffold (AZD1080) is effective. These results suggest that targets beyond GSK3 likely contribute to the phenotypes seen in vivo and have led us to formulate the hypothesis that CHIR99021 that stimulates mitochondrial activity without targeting GSK3 could have therapeutic benefit for HD. In Aim 1, we will identify molecular targets beyond GSK3 that mediate the effects of AZD1080 and CHIR99021 in HD models. In Aim 2, we will assess efficacy of CHIR99021 in both HD patient neurons and a chronic HD mouse model. The successful completion of our studies will validate our approach of enhancing mitochondrial efficacy and provide a chemical lead for further drug development toward novel therapeutics for HD and a wider range of neurological disorders marked by dysfunctional mitochondria.
The healthy blood-brain barrier (BBB) physically separates the central nervous system from the bloodstream and prevents effective delivery of many therapies. However, under certain pathological conditions such as brain cancer, the BBB is disrupted such that brain cells and surrounding extracellular matrix (ECM) are exposed to the systemic circulation. For the incurable brain cancer, glioblastoma (GBM), core tumor regions often enhance on magnetic resonance imaging (MRI) due to contrast permeability across disrupted BBB, but the invasive tumor margin remains behind an intact BBB. Such BBB heterogeneity has made GBM very difficult to treat, especially to eradicate therapeutically resistant GBM cells in the invasive margin likely responsible for recurrence. To address this issue, we propose an immunotherapy that first targets BBB disruption and accumulates at tumor sites of exposed brain ECM. Then, an immunogenic peptide capable of spreading to the invasive margins is released to generate a cytotoxic T cell response throughout the tumor volume.

To generate this therapeutic platform, we use a family of antibody-like targeting molecules called Variable Lymphocyte Receptors (VLRs). VLRs are lamprey antigen receptors that are particularly adept at binding glycosylated structures and therefore well-suited for targeting glycosylation-rich brain ECM. Using state-of-the-art VLR screening paradigms, we have identified multiple VLRs that preferentially bind brain ECM. After intravenous injection into tumor-bearing mice, the VLRs home to BBB permeability sites, indicating a capacity to specifically deliver GBM therapy. In this proposal, the VLRs will be fused with immunogenic peptides (VLR-IPs) that can be activated proteolytically by the GBM microenvironment, enabling IP spread throughout the tumor volume to mediate a cytotoxic T cell response. To validate the VLR-IP platform, we will first test the capability for the VLR-IPs to activate T cells in vitro, using both human and murine lymphocyte cultures. Next, the VLR-IPs will be administered to mice bearing an orthotopic, syngeneic murine GBM and therapeutic efficacy determined. The proposed work will therefore evaluate the potential of the VLR-IP platform as an innovative GBM treatment and motivate further detailed translational studies. If successful, ECM targeting of therapeutic payloads would also become a viable approach in many other neurological diseases that exhibit BBB disruption including stroke, traumatic brain injury and multiple sclerosis.
Neurodegenerative diseases produce diverse symptoms, but share some molecular mechanisms. Pathology related to the misfolding, phosphorylation and accumulation of microtubule-associated protein Tau, has been observed in several diseases, including Alzheimer’s, Fronto-Temporal Dementia and glaucoma. Our previous work has mapped a signaling pathway in Alzheimer’s from Amyloid-ß oligomers to synapse loss through Fyn kinase, and Fyn inhibitors are being tested in a Phase 2 trial.

It is well known that Fyn associates with Tau, and our data show that Fyn inhibition reduces Tauopathy in Alzheimer’s mice. Therefore, we propose that Fyn kinase inhibition may provide effective treatment for other Tauopathies. Here, we will investigate Tau-dependent Fronto-Temporal Lobar Degeneration (FTLD-Tau) and glaucoma.

We will treat mice modeling FTLD-Tau with the Fyn inhibitor, AZD0530 (Saracatinib), at doses effective in Alzheimer’s models. We will utilize two models, a transgenic strain over-expressing human mutant P301S Tau, and intracerebral injection of pathological Tau extracted from human autopsy brain. We will assess the ability of Fyn inhibition to reduce Tau pathology, synapse loss, cell degeneration and memory impairment, each of which is well documented in this strain. Critically, our Preliminary Data show that the drug eliminates memory dysfunction in this transgenic model. In addition, we seek to validate a role for Fyn by treating mice injected with human pathological Tau, and monitoring the extent of induced mouse Tau misfolding and the spreading of pathology. Together these data will determine whether Fyn inhibition generally, and AZD0530 particularly, is a therapeutic candidate for FTLD-Tau.

Vision loss in glaucoma is driven by increased intraocular pressure (IOP), and reducing IOP is the mainstay of therapy, though benefit is partial. There is a recognized unmet need for neuroprotective therapy to prevent progressive vision loss. It has been shown that glaucomatous retina exhibits Tauopathy, and one of our team members showed that silencing Tau expression in the eye eliminates ganglion cell loss. We propose that Fyn inhibition will reduce Tauopathy in the glaucomatous eye, thereby preserving function and cell number separately from IOP lowering by a neuroprotective mechanism. Importantly, our Preliminary Data reveal that retinal pattern ERG signals, which are reduced with elevated IOP, are maintained by Fyn inhibition in a glaucoma model. We propose to assess the ability of AZD0530 treatment to preserve retinal ganglion cell numbers and function in two glaucoma models. The outcomes will provide an assessment of whether Fyn inhibition might provide the first neuroprotective therapy for glaucoma.
Despite major advancements in progression-free and overall survival through the introduction of novel agents, relapse remains a major problem in high-risk multiple myeloma (HRMM). Our group is focused on the development of novel immunotherapeutic strategies for HRMM. In this innovative study, we will combine three highly-active agents: autologous expanded natural killer cells (auto-ENKs) that avidly kill myeloma targets, the anti-SLAMF7 antibody (ab) elotuzumab (Elo) which stimulates and redirects ENKs to myeloma, and the interleukin-15 superagonist ALT-803, a cytokine uniquely able to potently stimulate the expansion, activity, and persistence of natural killer cells. This regimen will be administered after auto-stem cell transplant (ASCT) to patients with previously-treated gene expression profile-defined HRMM in a FDA- and IRB-approved clinical trial. We safely treated HRMM patients in frank relapse with ENKs in a previous clinical trial, but found that the activation state of the infused ENKs was rapidly lost, likely due to suboptimal support provided by low dose IL2 and the suppressive effects of the bone marrow microenvironment (BM-ME). We have performed preliminary studies examining ways to enhance this approach. We found that Elo substantially enhances the activity of auto-ENKs, and ALT-803 extends the activation state and promotes the proliferative capacity of ENKs in vitro. We hypothesize that our combinatorial approach will maximize the efficacy of ENKs by 1) targeting the auto-ENK to myeloma using the antibody elotuzumab; 2) delivering therapy after ASCT, when the disease burden has been significantly reduced rendering the BM-ME more conducive to immune effectors; and 3) supporting transferred ENKs with ALT-803 rather than IL2. The principal endpoint will be response. The activation and persistence, anti-MM cytolytic ability, and homing of ENK will be assessed post-infusion by studying the peripheral blood and BM compartments. Effects of ALT-803 on the post-ASCT immune reconstitution of NK and T cells will be studied. We will also explore the potential adverse impact of the tumor ME on the ENKs to identify therapeutically actionable targets and evaluate the presence of immunosuppressive T regulatory cells. We hope that this combinatorial approach will transform NK immunotherapy and can be extended to other malignancies.
MicroRNAs (miRs) comprise a large family of small, ~21–23 nucleotide noncoding RNAs that have emerged as key post-transcriptional regulators of gene expression and act by silencing the translation of target mRNAs. To date, there are ~1,000 predicted human miRs believed to control the activity of >60% of all protein-coding genes. Not surprisingly, these small RNAs have been shown to play crucial roles in nearly all aspects of human biology from development to disease. In cancer, global down-regulation of tumor suppressor miRs (TS-miRs) is commonly observed and has been demonstrated to be a causative feature in tumorigenesis. Loss can stem from genetic mutation or deletion, promoter methylation or dysregulation of miR biogenesis. Of these mechanisms, alteration of global miR biogenesis is receiving increased attention due to recent findings demonstrating the critical role that miR-binding proteins (miR-BPs) play in the inhibition of this process, ultimately stimulating TS-miR degradation and cancer development. Current strategies for restoring TS-miR function rely on replacement therapy, whereby chemically-modified and formulated mature miR mimics are delivered to cells and tissues. Despite the promise of these agents in xenograft mouse models, similar to RNAi-based therapeutics, many liabilities exist. These include the requirement for chemical modifications to enhance in vivo pharmacokinetic/pharmacodynamic properties, hybridization-based off-target effects, toxicity, drug delivery problems and high cost. As such, no miR-targeted drugs have yet been approved for use. To overcome these challenges and provide a druggable and transformative solution for restoring TS-miR levels and conquering cancer, we propose to target miR−miR-BP interactions. Our overall goal is to discover and develop small molecule-based inhibitors of these interactions to prohibit the degradation of specific TS-miRs, thus providing the basis for next-generation miR-targeted therapeutics for cancer treatment. These proposed investigations are based on the hypothesis that regulatory miR-BPs specific for select miRs have been evolved as a quality control mechanism for controlling mature TS-miR levels, and this network is hijacked in cancer. If successful, these studies will catalyze the development of new therapeutics for the treatment of cancer, in addition to the identification of new miR−miR-BP targets for future drug discovery efforts. Our Specific Aims are as follows: (1) To develop a high-throughput screening (HTS) assay of the let-7−Lin28 miR−miR-BP interaction for small molecule inhibitor discovery, and (2) To develop a strategy for the discovery of miR-BPs using chemically-modified pre-miR probes.
Jeffrey Glenn, M.D., Ph.D. - 2016 Awardee

Medicine
Stanford University

“Targeting RNA Secondary Structure--Towards a New Class of High Barrier to Resistance Antiviral Agents.”

Our broad, long-term objectives are to develop a new class of antiviral therapeutics that target essential RNA secondary structures present in viral genomes, and that can be developed for all viruses for which no effective therapies exist.

As proof-of-concept, we will seek to target the influenza virus RNA secondary structure that we have recently discovered by SHAPE (selective 2'-hydroxyl acylation analyzed by primer extension) and demonstrated to be a universal influenza virus packaging signal--absolutely conserved in all viral isolates from 1918 epidemic flu to H1N1 and H5N1 bird flu--that mediates both packaging of the influenza virus genome as well as in vivo disease. In particular, we will test our hypothesis that potent pharmacological inhibitors of the virus’ packaging and life cycle can be identified using SHAPE, which offers a dramatic means of overcoming the critical scientific roadblock of conveniently determining viral RNA secondary structure targets, and identifying molecules capable of disrupting those targets. Our specific aims are to: 1) Perform the first SHAPE-based small molecule screen, and identify small molecules capable of disrupting the RNA secondary structure of the influenza virus packaging signal; 2) Identify a candidate lead molecule by determining the anti-influenza virus activity of the most potent hits from the SHAPE screen; 3) Optimize the lead molecule by determining its structure-activity relationships (SAR) using medicinal chemistry to synthesize analogs and assessing their SHAPE and antiviral activity; 4) Nominate a pan-genotypic anti-influenza virus development candidate with the best therapeutic index.

Successful accomplishment of our specific aims will not only result in a novel universal pan-genotypic inhibitor of influenza virus with a high barrier to the development of resistance that can be developed for clinical use; the exact same approach can then also be undertaken to develop an exciting new class of drugs targeting virtually any virus of interest--thereby opening up new avenues for treating and curing a wide range of diseases.
Triple-negative breast cancer (TNBC), representing ~15% of breast cancer diagnosed across the globe, is considered as incurable malignancy. Due to the lack of surface markers for current targeted therapy, there are currently no molecularly targeted therapies approved for TNBC.

To eliminate the mortality associated with TNBC, there is an urgent need to discover novel surface markers for TNBC and develop corresponding targeted therapy for TNBC. We hypothesize that tissue factor (TF) is a novel surface marker and oncotarget in TNBC and that TF-targeted immunotherapy is effective for the treatment of TNBC. The rationale behind the hypothesis is that (i) our preliminary studies demonstrate that TF is over-expressed on TNBC cancer cells as well as the tumor neovasculature in over 85% of TNBC patients (n=14) when using standard paraffin-embedded tumor tissues or in nearly 60% of TNBC patients (n=157) when employing tissue microarray slides. Importantly, TF is not expressed on normal breast gland and quiescent endothelial cells of normal vasculature. Thus, TF is a novel surface marker in TNBC and targeting TF may constitute a novel targeted therapy for a majority of TNBC patients. (ii) Dr. Hu’s prior work demonstrated that TF-targeted therapies are effective and safe in preclinical animal studies and clinical studies. He has co-invented and patented a TF-targeting Immuno-Conjugate agent named ICON that consists of factor VII (fVII, the natural ligand to TF) fused to the Fc region of human IgG1. As a neovascular-targeting agent, ICON has shown therapeutic efficacy and safety in preclinical animal models against solid cancers, macular degeneration (AMD) and endometriosis and in a Phase I trial for AMD patients. (iii) Dr. Hu has improved ICON by reducing 50% of molecular weight and completely eliminating its pro-coagulant effects yet with intact binding to TNBC, as a second-generation TF-targeting immunotherapy agent, called L-ICON. The major goal and specific aims of this proposal are to evaluate the novel L-ICON as single agent therapy and combination therapy with current immune modulatory agents (such as immune checkpoint inhibitors) and natural killer (NK) cells for TNBC. To ultimately translate the findings into clinic for TNBC patients, Dr. Hu is collaborating with breast cancer expert and clinical trialist, GMP manufacturing NK cell expert, pathologist and biostatistician on this proposal. If successful, the significance and impact of this project is that it will provide novel targeted approaches for the treatment of TNBC and hopefully to eliminate the mortality associated with this disease.
Organ transplantation currently relies on nonspecific immunosuppressive agents (IS) given lifelong to prevent graft rejection. These drugs have significant and cumulative toxicities, including nephrotoxicity and increased rate of infections, malignancy, diabetes, and hypertension. The elusive goal since organs were first transplanted in humans is to establish donor-specific immunologic tolerance, a state where a donated organ is accepted as "self," eliminating the need for IS. Regulatory CD4+CD25+FOXP3+ T cells (Tregs) have been shown to induce tolerance in transplantation models and to be elevated in tolerant human recipients. However, there have been limited efforts to harness their therapeutic potential in the clinic.

Our long-term goal is to develop tolerogenic protocols that will lead to controlled IS minimization or withdrawal after organ transplantation. As a step in that direction, the overarching goal of this proposal is to develop a recipient derived, DONOR-specific expanded regulatory T cell "product" for use in kidney transplant recipients, to be pursued under two specific aims.

Specific Aim 1: To further define and optimize the conditions for laboratory-based expansion of allospecific Tregs

(A) Our approach will be to verify if greater expansion without the loss of efficacy can be obtained by a non-specific restimulation with anti-CD3/CD28-beads on day 14, following the initial donor-specific stimulation with "donor" irradiated B cells on Days 0 and 7. (B) After this optimization, we will make two more small-scale Tregs expansions in the laboratory and then test these for their in vivo safety and efficacy (below).

Specific Aim 2: To optimize the conditions for large scale (i.e., clinically applicable) expansion of allospecific Tregs. The technology optimized in the laboratory will be transferred to the GMP Facility where two large-scale expansions will be carried out.

The Tregs expanded in both the laboratory (n=2) and the GMP Facility (n=2) will be tested for their (a) in vivo safety, testing whether they would lose their suppressive properties following infusion and instead cause rejection of human skin grafts on NSG mice, and (b) in vivo efficacy, determining if they protect human skin grafts on humanized NSG mice from rejection. Completion of these studies will enable us to file for an IND from the FDA to perform a future phase I clinical safety trial during the Transformational Award Phase, and then to a future phase II efficacy trial. The significance of this proposal lies in its broad applicability for induction of tolerance in organ and tissue transplantation.
Here at the Rehabilitation Institute of Chicago, we treat over 55,000 patients per year with a variety of diagnoses ranging from traumatic brain injury, spinal cord injury and stroke to Parkinson's disease and cerebral palsy. Virtually all of these patients suffer from movement disorders and weakness. Currently, the gold standard for diagnosis of neuromuscular problems involves painful and sometimes impossible muscle biopsies or very qualitative clinical exams. We believe the device described in this proposal to measure muscle properties will revolutionize treatment of these problems. Using this device, patient muscle function can be diagnosed and objectively characterized in a matter of seconds. This would have dramatic impact on the course of patient care and clinical decisions since muscle structure and function are so central to patient well being. While a few tools are available to collect in vivo muscle data, none can measure sarcomere properties noninvasively or with this degree of resolution. Necessary attributes are: simultaneous sampling across millimeters of muscle tissue, real-time sampling, compatibility with gross movement, inducing no or minimal tissue damage, and resolving nanometer-scale sarcomere length changes.

To solve this challenge, we are developing a new method termed resonant reflection spectroscopy (RRS). RRS is an optical method that relates reflected spectra to the muscle sarcomere structure. Unlike current methods, RRS operates through a minimally invasive optical probe, collects large sample sizes of sarcomere lengths, and is compatible with muscle movement and activity. In the catalyst stage, our goal is to develop RRS so that it can be routinely used in clinical settings. Specific aim 1 will develop a new optical source so RRS can be applied across general muscle architectures and geometries. Specific aim 2 will result in a robust optical probe designed for use in the clinical setting. Specific aim 3 will result in a packaged RRS system that can be used in a future transformational award across clinical settings. Our ultimate goal is to revolutionize the way that our patients with neuromuscular disorders are diagnosed and treated.
This proposal will address major limitations in the clinical effectiveness of emerging cancer immunotherapy regimens. Adoptive cell therapies and administration of checkpoint blockade antibodies, which depend on CD8 T cells of the patient’s immune system to specifically kill tumor cells, are predicted to revolutionize how cancer is treated. This proposal will exploit tissue resident memory CD8+ T cells (TRM) to overcome two major limitations of cancer immunotherapy efficacy: 1) tumors represent a strongly immuno-suppressive environment, which antagonizes immune-based therapies, and 2) therapies need to localize to tumors but current approaches are systemic and not anatomically targeted. TRM are a recently defined lineage of T cells that comprise the most abundant immunological surveyors of non-lymphoid tissues in mice and humans. TRM are established throughout the body as a consequence of natural infections and vaccines. TRM reactivation, achieved via topical application of cognate peptide, rapidly licenses these cells to direct the activation of all arms of the immune system, triggering a highly immune-stimulatory environment. This response, referred to as the ‘sensing and alarm’ function, occurs locally and recruits adaptive immune cells and serum antibody to the site of TRM reactivation. Reactivated TRM also potently activate local innate responses, dendritic cell priming, and natural killer cell activation. We hypothesize that TRM reactivation can reverse the immunosuppressive tumor microenvironment and can be exploited to enhance existing adoptive cell transfer and checkpoint blockade therapies. This proposal will answer two major questions. 1) Can TRM reactivation be exploited to promote an immune-stimulatory environment within solid tumors and mediate tumor regression? 2) Does TRM reactivation recruit engineered anti-cancer chimeric antigen receptor (CAR) T cells and checkpoint blockade antibodies to solid tumors? These questions will be addressed in autochthonous inducible tumors of the female reproductive tract (uterine and/or ovarian cancer) and skin (melanoma) in mice. Successful outcome will provide proof-of-principle demonstration of a human translatable strategy for improving upon cancer immunotherapies by modifying tumor microenvironments and focusing therapies to tumor sites. Future pursuits would interrogate TRM biology, specificity, and function within human tumors and will define regimens for therapeutic application.
Perhaps the most common hallmark of cancer is activation of the MYC oncogene. Nearly all tumor cells are strictly dependent on high MYC levels in order to drive malignant phenotypes like rapid proliferation, elevated invasion and migration, reprogrammed metabolism, and survival in the presence of cellular stresses like hypoxia and genetic damage. Despite this central role in the biology of cancer, strategies for directly targeting MYC have not been successful. We report here evidence suggesting that MYC may regulate an additional hallmark of cancer, i.e. immune evasion. At the molecular level, MYC drives expression of the TNF receptor family member CD30/TNFRSF8. While CD30 expression is known to be important in rare forms of lymphoma (e.g. Cutaneous T Cell Lymphoma/CTCL and Anaplastic Large Cell Lymphoma/ALCL), we demonstrate that more common lymphoid malignancies, and also common epithelial tumors, express CD30 as a surface antigen. Remarkably, CD30 levels are controlled by MYC in all tumor types we examined. Confirming the functional relevance of these observations, we demonstrate that blocking CD30 expression in either lymphoid or epithelial tumor cells causes rapid cell death. Thus, CD30 may serve as a biomarker and potential therapeutic target in cancers with elevated MYC. Furthermore, based on published findings on CD30 in Hodgkin's Lymphoma, it likely serves as an immune checkpoint molecule to provide immune evasion properties to MYC-driven tumors. The experimental plan outlined in this Catalyst Award proposal will formally test these highly impactful predictions. Successful completion of this proposed research will provide a solid rationale for a Falk Transformational Award that explores methods for exploiting these biomarker, targeted therapy and immune checkpoint blockade strategies.
B cells critically depend on continuous survival and proliferation signals from a functional B cell receptor (BCR). In >80% of B cell malignancies, the tumor clone is driven by an oncogenic BCR-mimic. Oncogenic BCR-mimics include BCR-ABL1, EBV-encoded oncoproteins, RAS-pathway and NF-kB-pathway activating lesions. While CD25 mediates IL2 signaling on T cells, we recently revealed that CD25 is a critical feedback regulator of B cell receptor (BCR) and oncogenic BCR-mimics in human B cell malignancies. Indeed, we found that CD25 is a reliable biomarker of tumor clones driven by oncogenic BCR-mimics (e.g. BCR-ABL1, LMP2A, BRAF, NF-kB) and genetic experiments demonstrated that CD25 is critical for the initiation of B cell leukemia and lymphoma in transplant recipients. Surface expression is rapidly induced by activity of PKC-delta and NF-kB downstream of the BCR. CD25 then recruits an inhibitory complex to the surface to reduce kinase signaling downstream of the BCR or its oncogenic mimics. Analysis of three clinical cohorts revealed that high expression levels of CD25 are associated with poor clinical outcome in various B cell malignancies. While CD25 expression is associated with drug-resistance, inhibition of CD25 sensitizes multiple B cell malignancies to conventional drug-treatment.

Based on these and other findings, we propose three Aims to (1) elucidate mechanisms of CD25 regulation, (2) explore usefulness of pharmacological suppression of CD25 and (3) CD25 as target for immunotherapy in B cell malignancies.

Aim 1: Mechanisms of CD25 regulation in B cell malignancies: Which mechanisms control transcriptional regulation (e.g. Rel-A/NF-kB) and surface expression (phosphorylation of cytoplasmic tail by PKC-delta) of CD25 and how are these events related to oncogenic mimicry of BCR signaling?

Aim 2: Pharmacological suppression of CD25 in human B cell malignancies. For suppression of CD25 at the transcriptional level, we will test the Rel-A small molecule inhibitor IT-901 (genetic validation based on Relafl/fl, Nfkb1fl/fl leukemia and lymphoma cells). To inhibit translocation to the cell membrane, we will inhibit PKC-delta function by Enzastaurin and CID2858522 (genetic validation based on leukemia and lymphoma cells from Prdkcdfl/fl and analog-sensitive Prkcd mice).

Aim 3: CD25 as a target for immunotherapy of B cell malignancies. We will test the efficacy and specificity of antibody-based targeting of CD25+ B cell malignancies in humanized NSG-KitW41 mice. In two experimental series, antibody-drug conjugate (ADC) against CD25 (ADCT-301) will be studied.
Re-wiring the human brain represents a critical frontier in clinical neuroscience. Behaviors and mental function are emergent properties of large-scale neuronal networks where the connectivity strengths between nodes define the network. Correspondingly, many neurological and psychiatric symptoms arise from network-level derangements where the structural and/or functional connectivities between separate brain areas have been altered by the disease process (e.g., sub-cortical stroke, traumatic brain injury) and/or secondary maladaptive network changes. While the adult brain has little capacity to re-grow damaged long-range axons, it might be possible to restore functional connectivity by training the brain to use an alternate route connecting the areas, with the ultimate goal of improving motor/behavioral/cognitive function. However, currently there are no clinical treatments available for restoration of functional connectivity between specific brain areas.

Here, we propose to develop a non-invasive brain stimulation/recording technique to functionally re-wire the human brain. Inspired by our previous animal studies and existing literature in healthy humans using paired associative stimulation (PAS), the instrument system utilizes two transcranial magnetic stimulator (TMS) channels to activate two cortical regions in precisely orchestrated temporal patterns to selectively up- or down-regulate functional connectivity between them. To this aim, we will develop new stimulation sequences and monitor the ensuing interregional connectivity changes with source-resolved electroencephalography (EEG). To improve the navigation and targeting accuracy of brain stimulation, we will use the newest-generation small but powerful cooled TMS coils, combined with our published TMS E-field computations with individual magnetic resonance images (MRIs) and a TMS navigator device. The TMS-EEG-MRI data will be correlated with connectivity estimates from high-angular resolution diffusion imaging (HARDI) diffusion MRI tractography and resting-state functional MRI (rs-fMRI), along with behavioral measures.

In summary, this proposal develops non-invasive techniques to transform our capability to manipulate brain interregional functional connectivity in humans, therefore laying the foundation for a new class of network-level therapies in disorders that involve white matter lesions or aberrant functional connectivity due to other reasons. We build the techniques within studies in healthy human subjects and stroke/TBI patients, but the findings will be applicable to a broad range of neurological and psychiatric disorders. If the Catalyst phase of the award is successful, the techniques may be applied in a controlled clinical trial during the Transformational phase. The TMS, EEG, and MRI devices are noninvasive, safe, and already FDA-approved, and the experiments can be conducted in their allotted time frames.
Systemic lupus erythematosus (SLE) is a devastating autoimmune disease in which autoantibodies drive potentially fatal tissue damage in hundreds of thousands of individuals in the U.S. Tragically, the disease frequently affects ethnic minorities, children, and women of childbearing age. There is no cure for lupus and the current standard of care for lupus relies heavily upon the symptom-suppressing use of immunosuppressive drugs that are associated with harmful side effects. To achieve remission in lupus, new treatments must be developed to restrain the pathogenic production of autoantibodies by B cells.

The laboratory of Dr. Stephen Waggoner at Cincinnati Children's has discovered a novel and critical role for natural killer (NK) cells in suppression of antibody production by B cells. His lab is currently funded to pursue means of inhibiting this regulatory function of NK cells to improve immune responses during vaccination and chronic infection. In this proposal, an innovative alternative strategy aimed at harnessing NK cells to eliminate follicular helper T cells (TFH) and suppress B-cell production of autoantibodies is proposed as a novel means to improve treatment and potentially cure lupus. There is strong precedence for the clinical use of NK cells in therapy of cancer. Investigators at our institution are developing chimeric antigen receptors (CARS) that can be expressed in NK cells to increase their ability to eliminate patient tumors. The premise of the current proposal, supported by Dr. Waggoner's preliminary studies, is that CAR-expressing NK cells can also be used to kill normal, non-cancerous cells that are central to the pathogenesis of autoimmune diseases like lupus.

This Catalyst Award will enable development of a novel CAR construct and transduction of primary human NK cells isolated from the blood of lupus patients. The subsequent experiments supported by this award will test the hypothesis that this CAR construct endows patient NK cells with the ability to specifically recognize and kill TFH cells. Furthermore, the ability of these CAR-expressing NK cells to suppress antibody production by B cells from lupus patients in vitro and in vivo will be evaluated.

This proposal represents an important paradigm shift in NK-cell biology and highlights a transformative use of NK cells in clinical therapy of lupus. Our broad, long-term objective is to introduce CAR-expressing NK cells into patients in order to reduce autoantibody secretion and promote remission of lupus and other TFH-mediated autoimmune diseases (e.g. rheumatoid arthritis) for which we have no cure.