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FACULTY IN MEDICINAL CHEMISTRY
Kuberan (Kuby) Balagurunathan
Professor of Medicinal Chemistry

Email: kuby.balagurunathan@utah.edu
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Kuberan%20B.php

Education & Training:
- M.S., 1994, Indian Institute of Technology, Madras
- Ph.D., 2000, University of Iowa
- Post-Doctoral Training, 2001-2004, MIT

Research Interests:
HEPARAN SULFATE PROTEOGLYCANS: Biosynthesis, Structures and Functions.

In the post-genomics era, it is now accepted that complex glycoconjugates such as proteoglycans regulate numerous patho-physiological processes in all living species. They carry enormous structural information in terms of sulfation, epimerization, domain organization, chain length, number of chains and type of chains along with their core proteins. Production of proteoglycans with such high complexity occurs in template-independent fashion seamlessly, yet our understanding of their biosynthesis, structures and functions is somewhat incomplete and imperfect. We are developing a wide variety of chemical biology tools to define the biosynthetic pathways of heparan sulfate and related glycosaminoglycans (GAG) such as chondroitin sulfate and dermatan sulfate. We synthesize heparin and heparan sulfate structures with a dozen recombinant enzymes to define the structural basis for the interactions of growth factors and Heparan sulfates and the subsequent biological actions. We recently found that a library of click-xylosides produce distinct GAG chains in cellular systems and proposed a GAGOSOME model for the dynamic regulation of combinatorial GAG biosynthesis. These molecular tools are currently used in the lab to define the snap shots of biosynthetic events and signaling events that are associated with development and diseases with a final goal to advance the study of heparanomics.
Amy Barrios  
Professor of Medicinal Chemistry  

Email: amy.barrios@utah.edu  
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Barrios_A.php  

Education and Training:  
- B.S., 1995, University of Utah, Salt Lake City, UT, Chemistry  
- Ph.D., 2000, Massachusetts Institute of Technology, Cambridge, MA, Inorganic Chemistry  
- NIH Postdoctoral Fellow, 2000-2003, Department of Pharmaceutical Chemistry, University of California, San Francisco  

Research Interests:  
Chemical probes for visualizing PTP activity: Protein tyrosine phosphatases (PTPs) play critical roles in cellular signaling, regulating tyrosine phosphorylation through hydrolysis of the tyrosine phosphate in a temporally, spatially and regioselectively controlled manner. In contrast to their counterparts, the protein tyrosine kinases (PTKs), the substrate selectivity, biological regulation and specific roles of PTPs are relatively poorly understood. However, aberrant phosphorytrosine-dependent cellular signaling plays an important role in many human diseases, including cancer, diabetes and autoimmunity. PTK-targeted drugs have hit the market with considerable success as anticancer agents, but no PTP-targeted drugs have been developed to date. In this project, our aim is to develop novel PTP-targeted chemical probes that can be used to elucidate the biological roles of PTPs and can serve as lead compounds in the development of PTP-targeted therapeutics. For example, we designed the phosphocoumaryl amino acid pCAP as a fluorogenic phosphotyrosine mimic. This probe has been invaluable in allowing us to profile the substrate selectivity of PTPs, perform several high-throughput screens to identify novel PTP inhibitors, and visualize PTP activity both directly in cells and in cell lysates through polyacrylamide gel electrophoresis. Current work includes characterizing and optimizing the new inhibitors we have discovered and developing novel activity-based probes for PTPs.  

Understanding the biological action of metal-based drugs: While the majority of drug molecules are organic compounds, several very successful drugs contain metal ions. Certainly the most well-know (and well-studied) example is cisplatin, a platinum containing anticancer agent, but other examples include auranofin, a gold-containing antiarthritic agent; Pepto-Bismol®, a bismuth-containing treatment for gastrointestinal problems; and imaging agents such as magnevist (a gadolinium-based MRI contrast agent) and cardiolyte (a technetium-based radioimaging agent). In our lab, we have been studying the ability of auranofin and auranofin analogs to inhibit enzyme activity as one possible mechanism of action in the body. Au(I)-based compounds such as auranofin inhibit thiol-dependent enzymes, and we have demonstrated that, by tuning the ligands bound to the Au(I) ion, we can tune the selectivity and potency of the Au(I)-mediated inhibition. The relative potencies and selectivities of the new complexes hold up not only in vitro but also in vivo.  

Designing redox sensors: A recent area of emphasis for our lab is the development of fluorogenic chemical probes that can be used to image the production of redox active species in vivo. Our first efforts in this field are aimed at developing hydrogen peroxide sensors that can be delivered to a specific subcellular location (i.e. the cell surface, the cytosol, the mitochondria, etc.) and at developing hydrogen sulfide sensors based on fluorogenic organometallic compounds.
Grzegorz (Greg) Bulaj  
Associate Professor of Medicinal Chemistry

Email: Grzegorz.Bulaj@hsc.utah.edu  
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Bulaj_G.php

Education and Training:  
- M.Sc., 1989, Biochemistry, University of Wroclaw  
- Ph.D., 1993, University of Wroclaw, Poland

Research Interests:  
Our research is focused on creating digital therapeutics (mobile medical apps) for epilepsy, pain, depression and cancer. Digital therapeutics are mobile apps which receive the FDA regulatory status of “software as a medical device”. Mobile medical apps can be combined with pharmaceutical drugs yielding drug-device combination products. The long-term goal of our research is to develop drug-device combination therapies which target chronic diseases at both pharmacological and behavioral levels.

Examples of our projects include:  
1. Development of mobile medical apps for the treatment of epilepsy:  
2. Development of music streaming as adjunct digital therapies for depression:  
3. Development of a mobile game coupling physical exercise and empowerment for children with cancer undergoing chemotherapy:  
Tom Cheatham III
Professor of Medicinal Chemistry;
Director, Center for High Performance Computing;

Email: tec3@utah.edu
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Cheatham_T.php
http://www.chpc.utah.edu/~cheatham

Education and Training:
• B.S., 1989, Middlebury College, Middlebury, VT, Chemistry (Honors)
• B.S., 1989, Middlebury, VT, Mathematics and Computer Science
• Ph.D., 1997, University of California, San Francisco, Pharmaceutical Chemistry

Research Interests:
The people in our lab use and develop molecular dynamics, free energy simulation, and trajectory analysis methodologies in applications aimed at better understanding biomolecular structure, dynamics and interactions. A strong focus of our funded efforts centers on the reliable representation of nucleic acid systems (DNA and RNA) in solution. For example, we helped solved the NMR structure of the drug-bound Hepatitis C virus IRES structure shown on the left. Based on this (and related structures), we can now apply CADD methods and simulation to better understand and design potential new Hepatitis C therapeutics. In addition, large efforts are underway to better characterize RNA structure and force fields through simulation of a large number of commonly observed RNA structural motifs and a large variety of NMR and crystal structures. We are also involved with international collaborative efforts to understand DNA structure, for example through the ABC consortium and long simulations of DNA.

Critical to reliable representation of the structure, dynamics and interactions is not only trying to simulation the biomolecules in their native solution environment but to also both critically assess and validate the simulation results with experiment. Our group focuses on both brute-force and enhanced sampling/ensemble-based simulation using available high performance computational resources at the University of Utah (www.chpc.utah.edu) and elsewhere. Outside resources include large allocations of computer time from Blue Waters, XSEDE (www.xsede.org), on the Anton machine at PSC and from other sources. With these resources we also are able to expose and overcome limitations in the methods and force fields.
Darrell Davis  
Professor and Chair of Medicinal Chemistry

Email: darrell.davis@utah.edu  
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Davis_D.php

Education and Training:
- B.S., 1982 Chemistry, University of Puget Sound
- Ph.D., 1988, Organic Chemistry, University of Utah

Research Interests:
My laboratory is involved in the study of nucleic acid and protein structure using high-field NMR spectroscopy. We have recently developed a structure-based drug design program focused on discovering and optimizing small molecules that interact with biomedically relevant RNA targets. NMR spectroscopy is uniquely suited to solving the 3D structures of RNA domains in complex with inhibitor molecules, and NMR also is a unique tool for identifying lead compounds that only interact weakly with macromolecules. The University of Utah has an outstanding biomolecular NMR facility with 500, and 600 MHz instruments locally, and access to 800 and 900 MHz instruments at the University of Colorado.

Hepatitis C virus (HCV) infection is a major cause of liver cancer in the US and liver disease associated with HCV accounts for the majority of liver transplants. In the developing world, a high percentage of HIV patients are also co-infected with HCV, presenting a particularly challenging health problem. The 5’ un untranslated region of the HCV RNA genome contains a large structured domain that serves as an IRES (internal ribosome entry site) that enables 5’ cap independent RNA translation. The IRES of HCV is an attractive therapeutic target since it is crucial for HCV replication. The RNA has a well-defined structure, raising the possibility for developing targeted therapeutics against HCV.

Our laboratory has solved the structure of a functionally important domain of the HCV IRES RNA in complex with an inhibitor of viral replication. Current research in the laboratory involves using NMR to screen for additional inhibitors that bind this target. We are also using NMR for a structure-based drug design initiative aimed at developing next-generation inhibitors with improved potency. The structure-based design project is multi-disciplinary, with a computational chemistry component in collaboration with the Cheatham laboratory, and a synthetic chemistry initiative in collaboration with the Rainier laboratory.
Raphael Franzini
Assistant Professor of Medicinal Chemistry

Email: raphael.franzini@utah.edu
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Franzini_R.php

Education and training:
- M.Sc., 2005 Ecole Polytechnique Fédérale de Lausanne
- Ph.D., 2010 Stanford University
- Postdoctoral Fellow, 2012-2015, ETH Zürich

Research Interest:
My research group focuses on the interfaces of Chemistry, Biology and Medicine with the aim of developing novel types of therapeutic agents, imaging probes and diagnostic assays. One particular research goal is the development of DNA-encoded chemical libraries as prospective tools for drug discovery and their application to lead development for cancer-associated targets. DNA-encoded libraries are collections of compounds in which each small molecule is uniquely encoded by a covalently linked DNA sequence. Panning encoded libraries for the protein of interest enriches target-binding molecules and high-throughput sequencing of the DNA-barcodes enable the straightforward identification of the corresponding structures. Encoded library technology allows screening ultra-large compound collections in a one-pot protocol. In addition to setting up a platform of libraries and screening them for drug candidates, we aim to expand this technology beyond the identification of affinity ligands and to constantly improve methodologies for library synthesis, encoding and screening.

Further research interests include the development and optimization of ligand-based tumor targeting strategies as therapeutic and imaging modalities and binary molecular probes for clinical diagnostics.
Margo Haygood
Research Professor of Medicinal Chemistry

Email: margo.haygood@utah.edu
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Haygood_M.php

Education and Training:
- B.A., 1976, magna cum laude, Harvard University, History and Science
- Monbusho scholar, 1979-1981, Tokyo University
- Ph.D., 1984, Marine Biology, Scripps Institution of Oceanography, University of California, San Diego

Research Interest:
Marine microbiology and marine biotechnology:
Marine Invertebrate symbioses: Well known for in-depth studies of marine invertebrate symbioses, particularly the association between the bryozoan Bugula neritina and its symbiont Endobugula sertula, which produces the anticancer bryostatins to protect the bryozoan’s offspring from predation. Due in large part to our work, this is now the best-understood example of a marine chemical defense symbiosis. More recently, demonstrated that shipworm symbionts, in addition to their known nutritional role, also contribute bioactive secondary metabolites to the association, for example:

Iron acquisition by marine bacteria:
Long term collaboration with Alison Butler of UC Santa Barbara on siderophores, molecules used in iron binding and transport, in marine bacteria. Established the prevalence of a new class of amphiphilic siderophores typical in marine bacteria, for example:
Chris Ireland
Distinguished Professor of Medicinal Chemistry;
Past Dean of the College of Pharmacy;

Email: cireland@pharm.utah.edu
Website: https://pharmacy.utah.edu/medchem/faculty/current-faculty/Ireland_C.php

Education and Training:
- B.S., 1973, University of California at San Diego, CA, Chemistry
- Ph.D., 1977, Scripps Institution of Oceanography, La Jolla, CA, Oceanography

Research Interests:
Our research focuses on the isolation and structure determination of compounds from marine invertebrates, the detection of their effects on biological systems, and the characterization of their mode of action and structural determinants contributing to their biological activity. Our emphasis on organisms from tropical reefs is based on the hypothesis that competitive environments select for potent biological activities. Reefs are highly diverse ecosystems that are subject to a variety of selective mechanisms such as predation, fouling by microorganisms, and limited benthic substrates for settlement of sessile organisms. A common strategy for competing in such environments adopted by several organisms that are structurally vulnerable to these pressures is the production or accumulation of chemicals that can be used defensively or offensively.

Although the human body and its pathogens are foreign to a typical marine community, many of the biological pathways implicated in diseases such as cancer, infection, and autoimmune disorders are similar enough to those of competitors in tropical reefs that high potency of biological activity is frequently found in marine natural products. While the ecological roles of most marine natural products are unknown, the high potency and selectivity observed in marine natural products and the observation that structure-activity relationships are often highly optimized in marine natural products suggests that acquiring and maintaining high potency in these systems is a dominant selective pressure acting on their biosynthetic pathways. Add in the fact that marine invertebrates often host diverse and complex microbial communities where invasive disruption on a short time scale is a distinct possibility and that sponges and, to a lesser extent, ascidians have long evolutionary histories among the metazoans and it is no wonder that such potent compounds are to be found in these organisms.

Read more at https://pharmacy.utah.edu/medchem/faculty/current-faculty/Ireland_C.php
Eric Schmidt  
Professor of Medicinal Chemistry

Email: ews1@utah.edu  
Website: http://www.bioscience.utah.edu/faculty/schmidt/schmidt.php

Education and Training:
- B.S., 1994, University of California at San Diego  
- Ph.D., 1999, Scripps Institution of Oceanography  
- NIH Fellowship, 1999-2001, Johns Hopkins University

Research Interests:
Secondary metabolites (natural products) are central both to interactions between organisms and to human health. The goals of my lab are (i) to understand the basic biology and chemistry of secondary metabolism; (ii) to apply that understanding to synthetic biology and metabolite engineering; (iii) to understand the chemical mechanisms of biosynthetic enzymes; and (iv) to discover and develop new drug lead natural products.

Marine animals are renowned sources of natural products with pharmaceutical potential. Research shows that several compounds are not made by the animals themselves, but rather by symbiotic bacteria. My lab is working to achieve a fundamental understanding of the players involved in synthesizing secondary metabolites in marine animals. We apply this knowledge in the discovery and engineering of small molecules with therapeutic potential.

Research area 2. RiPP biosynthesis.  
The ribosomally synthesized natural products (RiPPs) are among the most ubiquitous and abundant bioactive compounds in nature. We study how they are synthesized from a biochemical and synthetic biology perspective.

Cultivated bacteria are rich sources for discovery of new bioactive agents. In a collaborative project with researchers in the Philippines and the US, we are exploring the potential of mollusk symbionts to synthesize natural products that affect neurons and that have potential to treat disease.
Jaclyn M. Winter
Assistant Professor of Medicinal Chemistry

Email: jaclyn.winter@utah.edu
Website: https://www.bioscience.utah.edu/faculty/winter/index.php

Education and Training:
- B.S., 2004, State University of New York College at Fredonia, NY, Chemistry and Molecular Genetics
- Ph. D., 2010, Scripps Institution of Oceanography, UCSD, Marine Natural Product Biosynthesis
- Postdoctoral Fellow, 2010-2011 Leibniz Institute for Natural Product Research and Infection Biology, Hans Knöll Institute, Germany
- Postdoctoral Fellow, 2011-2014, University of California at Los Angeles

Research Interests:
Secondary metabolites are specialized small molecules produced in nature and often possess a variety of biological activities that can be used toward improving our quality of life. These molecules possess exquisite chemical diversity and are often an inspiration for the development of new pharmaceutical agents. At a time when antibiotic resistant bacterial infections are reaching epidemic proportions, there is an urgent need to discover new therapeutic agents. It has been shown that biological pressures influence the structural diversity of compounds produced in nature and marine-derived microorganisms often contain specialized enzymes not found in their terrestrial counterparts. Thus, these specialized microorganisms serve as an ideal resource for drug discovery efforts and for the characterization of novel biosynthetic enzymes.

Our lab is focused on 1) elucidating the biosynthetic blueprint that nature uses for assembling biologically active compounds in bacteria and fungi, 2) manipulating and reprogramming biosynthetic systems to generate new compounds with enhanced biological activities and 3) developing individual enzymes that carry out complicated reactions into renewable and environmentally friendly biocatalysts. These enzymes can be engineered to enhance the efficacy of existing therapeutics or be used in the synthesis or semisynthesis of pharmaceutically important compounds.
FACULTY
IN
PHARMACEUTICS
&
PHARMACEUTICAL CHEMISTRY
You Han Bae
Distinguished Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: you.bae@utah.edu
Website: https://pharmacy.utah.edu/pharmaceutics/faculty/directory/faculty/YHBae.php

Education and Training:
- B.S., 1980, Seoul National University, Seoul, South Korea, Chemical Engineering
- Ph.D., 1988, University of Utah, Salt Lake City, UT, Pharmaceutics

Research Interests:
Our main research interests are as follows:
- Polymeric delivery systems for low molecular and macromolecular drugs and cells.
- Tumor pH targeting
- Oral delivery of genes, biologics and anticancer drugs
Mingnan Chen
Assistant Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: mingnan.chen@utah.edu
Website: http://pharmacy.utah.edu/pharmaceutics/faculty/directory/faculty/mchen.php

Education & Training:
- B.A., 1996, Jimei University, Xiamen, China, Aquaculture
- M.Sc., 1999, Peking University, Beijing China, Biological Sciences
- Ph.D., 2007, University of Connecticut, Storrs, CT, USA, Pharmaceutical Sciences

Research Interests:
Dr. Mingnan Chen joined the Department of Pharmaceutics and Pharmaceutical Chemistry as an Assistant Professor. Besides teaching professional and graduate students in the College of Pharmacy, Dr. Chen is interested in the innovation and study of drug delivery strategies to elevate medical treatments including cancer therapy in particular.

His research program at the University of Utah will start with two immediate projects: 1) developing drug targeting strategies toward metastastic cancer cells, and 2) developing polypeptide-base nanocarriers for cancer vaccines. Before assuming the position at the University of Utah, Dr. Chen received postdoctoral training at Duke University in engineering cancer drug carriers by using recombinant polypeptides. He also characterized pharmaceutical, therapeutical, pharmacological properties of the carriers. Earlier at the University of Connecticut, Dr. Chen spent his Ph.D. study in exploring a quality control mechanism in the MHC Class I antigen presentation of human immunity. Dr. Chen has published 9 peer-reviewed articles and one book chapter and recently received the NIH career development award (K99) in cancer nanotechnology.
Shuyun Dong
Research Assistant Professor of Pharmaceutics & Pharmaceutical Chemistry

Email: shuyun.dong@utah.edu

Education and training:
- M.D., 1993, West China University of Medical Sciences, Chengdu, China
- M.P.H., 1996, in Environmental Health, School of Public Health, Sun Yat-sen, University of Medical Sciences, Guangzhou, China
- Ph.D., 2008, in Molecular Genetics and Microbiology, University of Massachusetts Medical School, Worcester, MA

Research Interests:
My long-term goal is to conduct research that aims to improve our understanding and treatment of neurologic diseases and cancers through manipulating RNA. During my Ph.D. work, I systematically studied several key events in mRNA life cycle including: splicing, localization, translation and degradation. My original findings were published in Molecular Cell and PLOS Biology. My postdoctoral research focused on understanding the alternative splicing of RNA and developing novel therapeutics for RNA splicing related diseases. My postdoctoral works were published in journals including Nature Molecular Page, Nature Protocols, Molecular Biosystems, and The Journal of Biological Chemistry. One of my articles was selected as Paper of This Week by The Journal of Biological Chemistry.

My present research interests at the University of Utah include: Engineering and delivering therapeutic RNA-binding proteins, developing therapeutic approaches based on mechanics of RNA alternative splicing and degradation.
Darrell R. Galloway  
Research Professor of Pharmaceutics and Pharmaceutical Chemistry  

Email: darrell.r.galloway@utah.edu  

Education and training:  
- B.S. Microbiology, 1972, California State University, Los Angeles, CA  
- Ph.D., Biochemistry, 1977, University of California, Riverside, Ca  
- Postdoc, Immunochemistry, 1980 Scripps Clinic & Research Foundation, La Jolla, CA  

Research Interests:  
Dr. Galloway's research interests have always centered around studies of infectious disease, principally vaccine development and/or immunotherapy. He is a recognized expert in the field of bacterial toxins where he has published extensively on structure-function studies of Pseudomonas exotoxin A, as well as genetic regulation of exotoxin production. Current project work includes two plague vaccine studies using novel adjuvant formulations. A more recent interest involves studies to determine the basis of microbial persistence of *Borrelia burgdorferi*, the causative agent of Lyme disease, and Burkholderia species which are associated with melliodosis or glanders, depending on the infecting species of Burkholderia. Microbial persistence, which is widely recognized yet not well understood, is associated with chronic infection, as well as being a form of antimicrobial resistance and thus constitutes a major area of clinical concern for many bacterial pathogens. The development of therapeutic strategies to identify and target persistent forms of microbial pathogens is a goal of these studies.
Hamid Ghandehari
Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: hamid.ghandehari@pharm.utah.edu
Website: www.ghandeharilab.utah.edu

Education and Training:
- B.S., Pharmacy, University of Utah
- Ph.D., Pharmaceutics and Pharmaceutical Chemistry, University of Utah

Research Interests:
The main focus of research in the Ghandehari laboratory is the development of novel methods for controlled delivery of bioactive agents. Four areas are being explored:
- Genetically engineered polymers for gene delivery
- Water-soluble polymers for targeted delivery
- Poly (amidoamine) dendrimers for oral delivery
- Inorganic nanoconstructs for controlled chemical delivery
David W. Grainger
Distinguished Professor of Pharmaceutics and Pharmaceutical Chemistry;
past Inaugural George S. & Dolores Doré Eccles Presidential Endowed Chair
of Pharmaceutics and Pharmaceutical Chemistry;
Professor and Chair of Biomedical Engineering

Email: david.grainger@utah.edu
Website: http://pharmacy.utah.edu/pharmaceutics/faculty/directory/faculty/
dgrainger.php

Education and Training:
- B.A., 1983, Dartmouth College, Hanover, NH, Engineering and Chemistry
- Ph.D., 1987, University of Utah, Salt Lake City, UT, Pharmaceutical Chemistry
- Alexander von Humboldt Postdoctoral Fellowship, 1988-89, Mainz, Germany

Research Interests:
Bioanalytical sensing and microarrays, cell-surface interactions, infection and bacterial-surface interactions, drug delivery, antibody drugs, novel vaccines

Our main research interests are as follows:
- Polymeric biomaterials interactions with cells
- Medical device infection and bacterial colonization
- Nucleic acid and protein microarray performance
- Macrophage interactions with biomaterials
- Novel live vaccine and protein delivery
- Bioanalytical sensing and device miniaturization
James N. Herron
Associate Professor of Pharmaceutics and Pharmaceutical Chemistry;
Executive Associate Dean for Professional Education, College of Pharmacy;

Email: james.herron@utah.edu
Website: https://pharmacy.utah.edu/pharmaceutics/faculty/directory/faculty/JHerron.php

Education & Training:
- B.S., Biology, University of Illinois Urban-Champaign, 1976
- M.S., Microbiology, University of Illinois, Urban-Champaign, 1979
- Ph.D., Microbiology, University of Illinois Urban-Champaign, 1981

Research Interests:
Dr. Herron is interested in translational research – specifically, translating innovative near-patient diagnostics technologies from the laboratory into the clinic. Presently, his research is focused in three areas: 1) optical biosensor applications in biodefense, diagnostics, companion diagnostics, environmental and food testing, and personalized medicine; 2) high-throughput homogeneous fluorescence assays for use in biodefense, diagnostics, and quality control and assurance; and 3) dynamic light scattering as a means of detecting intravenous (IV) drug incompatibilities, when co-administered through the same IV line. He has published 113 full-length research articles and 107 abstracts. He also edited a book entitled “Physical Methods to Characterize Pharmaceutical Proteins.”

Dr. Herron has 32 US and 41 foreign patents, and has co-founded three different point-of-care diagnostics companies to commercialize biosensor technology developed in his laboratory. He has also consulted for 3M Corporation, Abbott Laboratories, Agri-Analysis, ARUP Laboratories, Echelon Biosciences, Johnson & Johnson, Kansas Technology Enterprise Corporation, and mBiO Diagnostics. He also chaired the translational development subcommittee of the Rocky Mountain Regional Center of Excellence in Biodefense and Emerging Infectious Disease (RMRCE), a NIH-sponsored center that supports basic and translational research in biodefense and infectious disease, with focus on developing countermeasures for both.
Kyungsook Kim
Research Assistant Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: kyungsook.kim@utah.edu

Education & Training:
- B.S., 2007, Chemical engineering, Pukyong National University, Korea
- Ph.D., 2013, Bioengineering, Tokyo Women’s Medical University, Japan
- Postdoctoral fellow, 2015, Biomedical Engineering, Tufts University, USA

Research Interests:
Dr. Kim is a biomedical engineer in the College of Pharmacy's Department of Pharmaceutical Chemistry who serves as a co-investigator for CSTEC@Utah's University Technology Acceleration Grant (UTAG). Dr. Kim's current research is developing allogeneic cell sheets, which have shown high transplantation efficacy to targeted tissues and release anti-inflammatory and anti-fibrotic cytokines continuously to damaged organs. This cell sheet tissue engineering technology has a wide range of applications, including the treatment of renal fibrosis, partial thickness cartilage defects, and salivary gland disorders. Dr. Kim is collaborating with the School of Medicine and the School of Dentistry at The University of Utah to develop and translate allogeneic cell sheet technology for a variety of clinical applications.
Sung Wan Kim  
Distinguished Professor of Pharmaceutics & Pharmaceutical Chemistry;  
Distinguished Professor of Bioengineering;  

Email: SW.Kim@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmaceutics/faculty/directory/faculty/SWKim.php  

Education and Training:  
- B.S., 1963, in Chemistry from Seoul National University, Korea  
- M.S., 1965, in Physical Chemistry from Seoul National University, Korea  
- Ph.D., 1969 in Chemistry from University of Utah  

Research Interests:  
Dr. Kim’s current research includes the design of polymeric carriers for therapeutic gene delivery.  

Professor Sung Wan Kim was Director of the Center for Controlled Chemical Delivery at the University of Utah from 1985-2006. He is a pioneer in drug delivery research and has engaged in his research since 1974 in the areas of hydrogels, biodegradable drug conjugates, self-regulating drug delivery and stimuli sensitive polymers. He also worked extensively in medical polymers, especially blood compatible polymers. Dr. Kim’s present research includes design of novel polymers for the delivery of protein drugs, cells and genes.  

Dr. Kim has been elected to three U.S. national academies: National Academy of Medicine (1999), National Academy of Engineering (2003) and National Academy of Inventors (2014). Dr. Kim has received other numerous awards; among them are the Terumo Global Science Prize (2014), Research Achievement Award-Pharmaceutical Sciences World Congress (2004), Rosenblatt Prize (2003), Ho-Am Prize (2003), AACP Volwiler Award (2002), American Association of Pharmaceutical Scientists (AAPS) Dale Wurster Award (1998), Controlled Release Society (CRS) Founders Award (1995), and the Clemson Basic Biomaterials Award (1987). These awards are the highest scientific awards from their respective societies. He is the Founder and served as Co-Chairman of the International Symposium on Recent Advances in Drug Delivery, Salt Lake City, 1983-2005.  

In 2006, Dr. Kim received an honorary doctorate degree from the University of Twente. From 2004 to present, Dr. Kim is a Hanyang Distinguished Professor at Hanyang University. Dr. Kim to date has published over 500 papers and owns 35 U.S. Patents. He has trained over 150 scientists.
Jindřich Henry Kopeček
Distinguished Professor of Pharmaceutics and Pharmaceutical Chemistry;
Distinguished Professor of Bioengineering;

Email: jindrich.kopececk@utah.edu
Website: http://www.pharmacy.utah.edu/pharmaceutics/groups/kopecek/

Education & Training:
- M.S., 1961, Institute of Chemical Technology, Prague, Macromolecular Chemistry
- Ph.D., 1965, Institute of Macromolecular Chemistry, Prague, Macromolecular Chemistry
- Postdoctoral Fellow, 1967-1968, National Research Council of Canada, Ottawa
- D.Sc., 1990, Czechoslovak Academy of Sciences, Chemistry

Research Interests:
Research in the Kopeček Biomedical Polymers Laboratory focuses on two main areas:

Macromolecular therapeutics. Recent research focuses on the design of backbone degradable, long-circulating polymer carriers that contain enzymatically degradable oligopeptide sequences in the main chain. These \(N\)-(2-hydroxypropyl)methacrylamide (HPMA) copolymer-drug (epirubicin, gemcitabine, paclitaxel) conjugates have demonstrated enhanced efficacy in the treatment of animal tumor models when compared to non-degradable HPMA copolymer conjugates or free drugs. Combination therapy targeting both tumor-initiating and differentiated cancer cell populations is also studied.

Drug-free macromolecular therapeutics. A new therapeutic approach for the treatment of B cell malignancies has been developed. The effectiveness of the new system is based on biorecognition events without the participation of low molecular weight drugs. Apoptosis of cells can be initiated by the biorecognition of complementary peptide/oligonucleotide motifs at the cell surface resulting in the crosslinking of slowly internalizing receptors. Drug-free macromolecular therapeutics (DFMT) are composed from two nanoconjugates: 1) bispecific engager, Fab'-'MOTIF1 (anti-CD20 Fab' fragment conjugated with coiled-coil forming peptide or morpholino oligonucleotide), and 2) a crosslinking (effector) component \(P\)-(MOTIF2)\(_x\) \(N\)-(2-hydroxypropyl)methacrylamide (HPMA) copolymer or human serum albumin grafted with multiple copies of complementary oligopeptide or morpholino oligonucleotide). The efficacy of the system was validated in vitro, in vivo and on cells isolated from patients with various subtypes of B cell malignancies.
Carol Lim  
Associate Professor and Interim Chair of Pharmaceutics and Pharmaceutical Chemistry  

Email: carol.lim@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmaceutics/groups/lim-lab/  

Education and Training:  
- B.S., 1987, Purdue University, W. Lafayette, IN, Pharmacy  
- Ph.D., 1996, University of California, San Francisco, CA, Pharmaceutical Chemistry  
- Post-Doc, 1999, National Institutes of Health, Bethesda, MD, NCI, PRAT Fellow  

Research Interests:  
Globally, our research is focused in 2 disease areas: breast cancer and leukemia. Our research focuses on:  
- Trafficking of human progesterone receptor (PR), including mechanisms of export of A and B isoforms of PR  
- Import kinetics of PR into the nucleus and the correlation to agonist dose and gene activation, which involves cellular kinetics and time-lapse microscopy  
- Use of signal sequences (including hormone inducible signals) for regulated, controlled delivery of gene therapy products (therapeutic targets include PR-positive breast cancers and chronic myelogenous leukemia).
John W. Mauger  
Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: john.mauger@hsc.utah.edu  
Website: https://pharmacy.utah.edu/pharmaceutics/faculty/directory/

Education and Training:
- B.S. Pharmacy, 1965, Union University Albany College of Pharmacy
- M.S., 1967, University of Rhode Island
- Ph.D., 1971, University of Rhode Island

Research Interests:
Dr. Mauger’s research is focused on developing standards for pharmaceutical dosage forms that can be used to assure safe and effective medicines. In particular, the application of sound and innovative measurement science to test the dissolution properties of solid dosage forms remains as one of the most useful tools for quality control purposes to assure that the active ingredient is released for therapeutic effect. These standards and tests are recognized for their importance to public health by the United States Pharmacopeia and the U.S. Food and Drug Administration.
Shawn C. Owen  
Assistant Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: shawn.owen@hsc.utah.edu  
Website: http://www.owenlab.com

Education and Training:  
- B.S., 2005, in Chemistry; B.A. in Chinese, 2005, University of Utah  
- Ph.D., 2009, in Pharmaceutics and Pharmaceutical Chemistry, University of Utah  
- Postdoctoral Fellow, 2009-2013, University of Toronto

Research Interests:  
- Antibody-based Therapeutics and Diagnostics  
- Hydrogels for Tissue Engineering  
- Cell-mediated Treatment  
- Biotherapeutics Stability
Jiyuan (Jane) Yang  
Research Professor of Pharmaceutics and Pharmaceutical Chemistry

Email: jiyuan.yang@utah.edu  
Website: https://pharmacy.utah.edu/pharmaceutics/faculty/directory/research_faculty.php

Education and Training:
- B.S., 1986, Tsinghua University, Beijing, China, Chemical Engineering  
- M.S., 1989, Dalian Institute of Technology, China, Polymer Materials  
- Ph.D., 2001, Peking University, China, Biomedical Polymer Science  
- Postdoctoral Fellow, 2002-2003, Institute Curie, Paris, France, Liquid Crystal Elastomers (Artificial Muscle)

Research Interests:
Having been working on biomaterials and drug delivery nearly 30 years, I have expertise in the design, synthesis and characterization of polymers and bioconjugates. From injectable biodegradable vaccine-delivery microspheres to new constructs of antibody-drug conjugates, I have accumulated a broad background with specific expertise in the synthesis and evaluation of various nanomedicines, in particular N-(2-hydroxypropyl)methacrylamide (HPMA)-based bioconjugates. For example, a patented technology relevant to more effective 2nd generation HPMA-based polymer-drug conjugates has been developed in our laboratory. This new synthetic strategy endorses inert HPMA polymer with biodegradability, which permits to prepare high molecular weight HPMA-drug conjugates with simple linear architecture while maintaining good biocompatibility. Various preclinical studies demonstrate distinct advantages over current clinical chemotherapeutic agents on tumor inhibition, indicating great potential to enhance cancer therapy.

I also work on new combination treatment strategies. In addition, our recent work on new construct of antibody-drug conjugates has won ‘2017 Best Research Paper in European Journal of Pharmaceutical Sciences’, in which I served as corresponding author. This design integrates the high specificity of antibody-drug conjugates with advantages of macromolecular therapeutics.
FACULTY
IN
PHARMACOLOGY
&
TOXICOLOGY
Louis Barrows
Professor of Pharmacology and Toxicology

Email: lbarrows@deans.pharm.utah.edu
Website: https://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Barrows-L.php

Education and Training:
- B.S., 1975, California Polytechnic State University
- Ph.D., 1980, University of California, Irvine, Pharmacology.

Research Interests:
My laboratory is dedicated to the discovery of new anti-cancer and anti-infective agents. Much of what we do can be considered natural products drug discovery. We identify new drug leads based on their novel chemical structure or mechanism of action. Extracts of macro- and microorganisms from coral reefs and tropical rain forests provide the new molecules we isolate and evaluate. Determination of the molecular actions of new molecules and determination of the precise cellular consequences of their activity is often the basis of student doctoral projects. We take bioactive organisms and molecules all the way from the source to the sequencing gel, and then into animal models of human disease.

Our recent major project has focused on linking scientific discovery with conservation and social progress in Papua New Guinea (the “PNG ICBG”). We have supported the development of a research lab in the University of Papua New Guinea (UPNG), School of Medicine and Health Sciences, and we have funded the botanical survey work of the National Herbarium in the School of Natural and Physical Sciences. This collaboration has permitted UPNG students to study here in Utah, and also for Utah students to spend time in PNG offering workshops and hands-on training. We have assisted in many UPNG Honor’s and Master’s students projects aimed at validating traditional phytomedicines, and identifying their active components. Our work has also funded numerous community education and conservation outreach efforts, which we have linked to the botanical survey expeditions. Our work in PNG has broadened our research interests to include the discovery of new molecules and mechanisms of anti-HIV and anti-TB activity.
Donald Blumenthal
Associate Professor of Pharmacology and Toxicology;
Associate Dean for Interprofessional Education and Assessment;
Adjunct Associate Professor of Biochemistry;
Adjunct Associate Professor of Biomedical Informatics;
Associate Editor of Goodman & Gilman's The Pharmacological Basis of Therapeutics,
On-line edition

Email: Don.Blumenthal@pharm.utah.edu
Website: http://pharmacy.utah.edu/phar motox/faculty/current-faculty/Blumenthal-D.php

Education and Training:
- B.A., 1975, University of California, San Diego
- Ph.D., 1980, University of California, San Diego, Physiology and Pharmacology

Research Interests:
Much of my laboratory's current basic research efforts are directed towards biophysical studies of protein kinases and other signal transduction molecules using fluorescence, circular dichroism (CD), small-angle x-ray (SAXS) and neutron scattering (SANS), and molecular dynamics (MD). We are using these methods to better understand the large-scale dynamic properties of these important drug targets and their role in normal cell function and disease.

We have also recently begun to study human acetylcholinesterase, the enzyme that hydrolyzes the neurotransmitter acetylcholine. This enzyme is the target of nerve gas agents and several drugs that have therapeutic application in treating glaucoma and Alzheimer's disease. We are interested in developing better antidotes to inhibitors of this enzyme by studying the conformational dynamics of the protein using techniques such as SAXS, MD simulations, and neutron and X-ray diffraction.
Marco Bortolato  
Associate Professor of Pharmacology and Toxicology  

Email: marco.bortolato@utah.edu  
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Bortolato-M.php  

Education and Training:  
- M.D., 2000, University of Cagliari, Italy  
- Ph.D., 2004, Pharmacology, University of Cagliari, Italy  
- Specialization in Neurology 2009, University of Cagliari, Italy  

Research Interests:  
My research is primarily focused on the characterization of the biological bases of neurodevelopmental disorders through the employment of behavioral tests in animal models. In particular, my key scientific interest is the identification of the mechanisms of interaction between lipid mediators (including neuroactive steroids and endocannabinoids) and key brain neurotransmitters, such as dopamine and serotonin. To this end, we employ a broad array of behavioral, biochemical and molecular biological techniques, as well as stereotactic surgery and HPLC.

The long-term goal of this research is the development of novel therapeutic agents for the treatment of pathological aggression, autism-spectrum disorder, Tourette syndrome, schizophrenia and impulse-control disorders (such as pathological gambling).

My laboratory is currently involved in two major translational research projects. The first target of our investigations is the characterization of the molecular substrates of gene x environment interactions in impulsive aggression. The second aim of our research is the development of novel steroid- and cannabinoid-based tools for the therapy of impulse-control disorders, Tourette syndrome, autism and schizophrenia. Both projects involve multiple collaborations with several basic and clinical scientists in US, Canada, Italy, France and Germany.
**William Crowley**  
**Professor of Pharmacology and Toxicology**

**Email:** William.Crowley@deans.pharm.utah.edu  
**Website:** [http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Crowley-W.php](http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Crowley-W.php)

**Education and Training:**
- B.S., 1970, University of Connecticut
- M.S., 1972, Villanova University
- Ph.D., 1976, Rutgers University

**Research Interests:**
My laboratory investigates the neuroendocrine and neurochemical factors that regulate the secretion of anterior and posterior pituitary hormones. In particular, we have focused on identifying and characterizing the actions of brain neurotransmitters and peptides that participate in the regulation of the secretion of luteinizing hormone, the anterior pituitary hormone responsible for ovulation, and prolactin, the anterior pituitary hormone that controls milk secretion in lactation. Multidisciplinary approaches are used in these investigations, including in vivo microdialysis to study release of brain neurotransmitters and peptides in discrete brain areas, biochemical measurements of neurotransmitters and peptides, measurements of expression of specific messenger RNAs involved in neurotransmitter and peptide transmission, neurotransmitter and peptide receptor binding, and various approaches towards studying signal transduction mechanisms. Studies are done in whole animal and in isolated brain tissues and cell culture. Current studies are focused on the actions of neuropeptide tyrosine (neuropeptide Y), which plays a central role in controlling the release of these hormones and in regulating food intake during lactation.

A second line of investigations employs all of these methods to identify and study actions of the neurotransmitter and peptide systems that control the secretion of the neurohypophyseal hormones, oxytocin, which is important for milk release during lactation, and vasopressin, which participates in control of body fluid homeostasis. This work is now focusing on the actions of ovarian hormones on brain transmitters, particularly norepinephrine, glutamate and γ-aminobutyric acid, and peptides in late pregnancy that increase the activity of the oxytocin neurosecretory system in preparation for birth and lactation.
Kristen Keefe
Professor of Pharmacology and Toxicology

Email: K.Keefe@utah.edu
Website: http://pharmacy.utah.edu/pharmtoc/faculty/current-faculty/Keefe-K.php

Education and Training:
- B.S., 1984, Case Western Reserve University
- M.S., 1989, University of Pittsburgh
- Ph.D., 1992, University of Pittsburgh
- Post-doctoral fellow, 1992-1995, NIMH

Research Interests:
My laboratory is interested in the structure and function of the basal ganglia, a group of subcortical nuclei in the brain involved in the control of movement and cognition. The importance of the basal ganglia for normal behavior is highlighted by the profound deficits observed in patients with Parkinson's disease, Huntington's disease, schizophrenia, and drug addiction -- diseases that are associated with dysfunction in the basal ganglia. Our work determines the influence of both endogenous and exogenous chemicals on the function of neurons in the basal ganglia in an attempt to better understand 1) the role that glutamate (via NMDA receptors) and monoamines (dopamine and serotonin) play in regulating the activity of basal ganglia nuclei, 2) the mechanisms by which drugs of abuse that affect the basal ganglia exert both their acute and long-lasting adverse effects, and 3) the mechanisms by which the function of the basal ganglia can be beneficially altered by drugs to better treat sequelae associated with dysfunction in these nuclei. We use numerous techniques to examine the effects of both endogenous and exogenous drugs on basal ganglia function. These techniques include: 1) In vivo microdialysis in the brain of awake animals to examine changes in the release of amino acid, monoamine, and neuropeptide neurotransmitters. 2) In situ hybridization histochemistry to measure changes in the levels of messenger RNAs in brain neurons to examine short- and long-term changes in gene expression in neurons of the basal ganglia. 3) Immunohistochemistry to examine changes in protein expression in defined basal ganglia neurons and nuclei. 4) Behavioral analyses of learning and memory processes mediated by corticostriatal circuits. Coupling these techniques, we can begin to understand how neurotransmitters and drugs acutely affect the function of basal ganglia neurons and the neuroadaptive changes that occur in response to neural injury in the basal ganglia and exposure to therapeutic and abused drugs.
Mei Yee Koh  
Assistant Professor of Pharmacology and Toxicology

Email: mei.koh@utah.edu  
Website: https://pharmacy.utah.edu/pharmtox/faculty/current-faculty/koh-mei.php  

Education and Training:
- PhD 2004, University of Manchester, Molecular Pharmacology  
- BSc (Honors) 2000, University of Manchester, Biochemistry with Biotechnology

Research Interests:
All solid tumors and their metastases suffer from regions of oxygen deprivation, also known as hypoxia. This occurs as a result of the diffusion limitation of oxygen, and to the highly proliferative nature of cancer cells. Hypoxic tumors are intrinsically resistant to radiation and chemotherapy, and patients with these tumors are likely to have a poorer prognosis. My lab aims to determine how tumor hypoxia, and the hypoxia-inducible factors, \textit{HIF-1} and \textit{HIF-2}, drive outcomes that promote cancer progression and resistance to therapy. These outcomes include ‘reprogramming’ of cancer cells to favor increased growth (such as through increased iron uptake), reversion to a multipotent stem-like (and more aggressive) phenotype, and increased metastasis. The hypoxic tumor microenvironment also alters the patient’s immune cells that are in proximity to the tumor, suppressing their ability to eliminate cancer cells. The overall goal of my lab is to identify new therapeutic strategies for cancer by targeting components of the tumor and the tumor microenvironment that drive cancer progression.

Tumor site-specific research programs are described at https://pharmacy.utah.edu/pharmtox/faculty/current-faculty/koh-mei.php
Cameron S. Metcalf  
Research Assistant Professor of Pharmacology and Toxicology;  
Associate Director, Anticonvulsant Drug Development Program;  

Email: cameron.s.metcalf@utah.edu  
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/cameron-metcalf.php  

Education and Training:  
- B.S., 2001, University of Utah, Salt Lake City, UT, Biology and Human Development/Family Studies  
- Ph.D., 2008, University of Utah, Salt Lake City, UT, Pharmacology  

Research Interests:  
Dr. Metcalf is a Research Assistant Professor in Pharmacology and Toxicology. He is also a Co-Investigator and the Associate Director of the Anticonvulsant Drug Development Program. Dr. Metcalf’s primary research interests include the evaluation and advancement of novel therapies for the treatment of epilepsy and pain. This work also involves ongoing efforts to identify and validate new animal models that can be used to screen and differentiate lead compounds. Approximately one-third of patients with epilepsy continue to have seizures despite treatment with one or more drugs. In addition, there are several special populations of epilepsy patients, including pediatric epilepsies and genetic syndromes, for which there are little or no effective therapies. Therefore, the development of new compounds for various forms of epilepsy is of critical importance for the ongoing treatment of epilepsy.  
For several years, Dr. Metcalf has also been interested in the development of neuropeptide compounds for the treatment of epilepsy and pain. Both epilepsy and pain are conditions where currently available treatments are often inadequate for many patients, and novel targets for therapies such as neuropeptides may offer new therapeutic opportunities. Dr. Metcalf’s work in this area has included both pre-clinical screening of novel compounds and the late-stage testing of lead candidate compounds. Dr. Metcalf’s ongoing work in this area includes testing of analogs of the neuropeptide galanin, alone and in combination with other drugs, in models of epilepsy and pain, in order to identify a lead compound for progression into clinical testing. Finally, Dr. Metcalf is also interested in the mechanisms of and potential clinical interventions for sudden unexpected death in epilepsy (SUDEP). While it is known that individuals with epilepsy, and particularly those with poorly controlled seizures, are at an increased risk for SUDEP, the mechanisms for this condition are not well understood. Therefore, understanding of risk factors, biomarkers, and potential clinical interventions to prevent SUDEP are an area of unmet need in epilepsy.
David Moody
Research Professor of Pharmacology and Toxicology

Email: david.moody@utah.edu
Website: http://pharmacy.utah.edu/pharmtx/faculty/current-faculty/Moody-D.php

Education and Training:
- B.A., 1972, University of Kansas, Chemistry
- Ph.D., 1977, University of Kansas, Experimental Pathology
- Postdoctoral Fellow, 1977-1980, University of California

Research Interests:
Our laboratory has used the power of mass spectrometry (MS) coupled to liquid or gas chromatography to measure amounts of drugs, drug metabolites and other xenobiotics in biological samples. We often use this technology to provide analytical services to other investigators, either commercially or collaboratively. When funding for research is obtained, we have used MS techniques to study in vitro drug metabolism. This includes use of animal models, human liver microsomes, cDNA-expressed drug metabolizing enzymes and human clinical studies. We have examined the role of cytochrome P450s (CYP) 2D1 in rat metabolism of amphetamine, the involvement of different CYPs in the metabolism of l-acetylmethadol (LAAM) and methadone, and the involvement of both CYPs and glucuronosyltransferases (UGT) in the metabolism of buprenorphine. Currently we are studying in vitro inhibition of the metabolism of oxycodone, methadone and buprenorphine.
Philip Moos
Associate Professor of Pharmacology and Toxicology

Email: philip.moos@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmpo/faculty/current-faculty/Moos-P.php

Education and Training:
- B.S., 1989, University of Colorado, Boulder, CO. Aerospace Engineering Sciences,
- M.S., 1990, University of Colorado, Boulder, CO. Aerospace Engineering Sciences,
- Ph.D., 1994, Kansas State University, Manhattan, KS. Biology,

Research Interests:
The Moos laboratory is broadly interested in understanding cancer susceptibility and the development of resistance. Research efforts are focused on cancer genomics, mechanisms of disease, modifiers of cancer risk, and drug susceptibility.

Current work is primarily in three arenas: 1) tumor heterogeneity and the role genotype and phenotype in the response to therapeutics during cancer progression, 2) combining genomics and drug screens to identify novel therapeutic agents, and 3) genomic evaluation of nanoparticles for inflammatory potential or other toxic adverse effects.

The first and second area of research are collaborations with Dr. Bild and focus on tumor heterogeneity. Most tumors are thought to have monoclonal origin but by the time they are diagnosed, they are heterogeneous in terms of the subclonal structure driving their growth. We are involved in a collaboration where we are using various genomic strategies, from whole genome sequencing to single cell-RNA sequencing to elucidate the subclonal structure and pathways that dominate the subclones. We are also testing unique drugs and drug combinations to identify new potential strategies for therapeutic intervention.

The third and newer, area of research utilizes transcript profiling and apoptotic studies to evaluate particulate matter toxicity and inflammatory potential. We have found that certain metal oxide manufactured nanoparticles have significant toxicity and induce a cellular response to the stress of the nanoparticles. Nanomaterials provide many benefits and are being utilized more broadly so it is important to understand the potential toxicities, and mitigation strategies prior to use. Therefore, we collaborate with Dr. Ghandehari to understand the basis of the toxicology of materials with the potential for biomedical application. We also collaborate with Dr. Reilly to evaluate the genomic consequences of airborne pollutants that are risk factors for diseases like asthma.
Randall T. Peterson  
L. S. Skaggs Presidential Endowed Chair and Dean, College of Pharmacy;  
Professor of Pharmacology and Toxicology;

Email: randall.peterson@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/randall-peterson.php  

Education and Training:  
- B.S., in Molecular Biology, Brigham Young University  
- Ph.D., in Biochemistry, Harvard University  
- Postdoctoral Fellow, Massachusetts General Hospital and Harvard Medical School  

Research Interests:  

Developmental Biology: Small molecules are powerful tools for studying developmental biology because they provide timing and dosage control over developmental pathways that is difficult to achieve with genetic mutations. Unfortunately, only a handful of developmental pathways can currently be targeted with small molecules. We are discovering novel chemical modifiers of developmental pathways by exposing zebrafish embryos to libraries of structurally diverse small molecules and identifying those that induce specific developmental defects. Using screens of this type, we have discovered dozens of compounds that cause specific defects in hematopoiesis, cardiac physiology, embryonic patterning, pigmentation, and morphogenesis of the heart, brain, ear, and eye and germ cell lineage.

Disease Physiology: One focus of our group is modeling human diseases in zebrafish. We then use the models to screen large chemical libraries for small molecule modulators of the disease-related phenotypes. The compounds we discover help us elucidate disease mechanisms and serve as starting points for developing new drug candidates.

Disease physiology is often complex and involves interactions between multiple organs and tissue types. Consequently, many diseases cannot be studied effectively using in vitro assays. The zebrafish is an excellent vertebrate model system to study many complex, non-cell autonomous diseases because the diseases can be studied in a native, whole-organism setting. In addition, compounds discovered in zebrafish screens have the advantage of having been selected for their ability to be active, efficacious, and well tolerated in animals.

Neuroscience & Behavior: Behaviors are accessible readouts of the molecular pathways that control neuronal signaling. Our group develops tools and techniques for comprehensive and high-throughput behavioral phenotyping in the zebrafish. These tools have some potential to improve our understanding of the neuronal signaling and may accelerate the pace of neuroactive drug discovery.

Zebrafish Reverse Genetics: Zebrafish have proven to be a powerful genetic tool over the years, primarily through forward genetic screens where fish are mutagenized (typically with chemical agent) and screened for obvious defects. We are now on the verge of the next exciting step in zebrafish genetics: reverse genetics! Using targeted DNA disruption, we are now making designer mutations in specific genes of interest. Here are some of the recent papers describing three different processes.
Christopher Reilly  
Professor of Pharmacology and Toxicology

Email: chris.reilly@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Reilly-C.php

Education and Training:
- B.S., 1994, University of Utah. Biology, minor in Chemistry  
- Ph.D., 1999, Utah State University, Toxicology

Research Interests:
My laboratory has three areas of research: TRP ion channels in lung physiology and disease, mechanistic studies of drug metabolizing enzymes, and biological applications of mass spectrometry.

1. TRP ion channels are a family of proteins that exhibit unique functional properties and associated physiological functions. We are interested in how TRP channels regulate physiological and adverse events in the lung, with particular emphasis on the ability of these channels to elicit deleterious pulmonary inflammation and lung cell death when activated by endogenous and/or exogenous agonists; pulmonary inflammation and acute lung damage are two critical components of lung diseases such as asthma, chronic obstructive pulmonary disease (COPD) and emphysema, fibrosis, and acute lung injury/ARDS. Our research is to establish roles of different members of the TRP ion channel family in the development and progression of lung injury and diseases and we have an active research program investigating receptor targets of pneumotoxicants and potential therapeutic inhibitory chemicals, the identification of specific cellular pathways that modulate deleterious and/or beneficial responses of lung cells to TRP channel agonists, and evaluation of TRP channel-dependent pathways in adverse outcomes in experimentally-induced disease states. Currently we are funded by the National Institute of Environmental Health Sciences (NIEHS) to determine how different components of polluted air adversely affect the human respiratory system.

2. Xenobiotic metabolism is a collective process by which chemicals that enter our body are modified. We are interested in how human cytochrome P450 enzymes transform chemicals to pharmacologically and/or toxicologically inactive vs. toxic reactive intermediates that damage cellular macromolecules, cells, and organ systems. We have two primary areas of focus: 1) enzymatic and chemical mechanisms of oxygenation vs. dehydrogenation of substrates and analysis of modified biological macromolecules (i.e., DNA and protein) by electrophiles, particularly with respect to such processes in lung tissue; and 2 the elucidation of how variations in drug metabolism mechanisms and efficiency dictate the pharmacological and/or toxicological properties of inhaled and/or systemically-delivered therapeutic drugs. Currently research in this area is supported by grants from the National Institute of General Medical Sciences (NIGMS) and Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD).

3. Mass spectrometry is a versatile and powerful analytical technology that has revolutionized mechanistic pharmacology and toxicology studies. We use mass spectrometry to quantify a variety of substances in diverse matrices, to establish structures and metabolic origins of drug metabolites in P450 reactions, to discover unknown bioactive substances that have measurable biological activity using metabolomics-based methodology, and to identify macromolecular targets of toxic electrophiles generated by P450s. Mass spectrometric analysis is a routine approach used by our laboratory and it is supported by a variety of research grants, subcontracts, and collaborations.
Joseph Rower
Research Assistant Professor of Pharmacology and Toxicology

Email: joseph.rower@hsc.utah.edu
Website: https://pharmacy.utah.edu/pharmtox/faculty/current-faculty/rower-joseph.php

Education and Training:
Postdoctoral Fellowship
- University of Utah Pediatric Clinical Pharmacology
- University of Colorado Anschutz Medical Campus, Antiviral Pharmacology
- University of Colorado Anschutz Medical Campus, Systems Biology

Doctoral Training
- University of Colorado Anschutz Medical Campus, Pharmaceutical Sciences

Undergraduate
- California Lutheran University Chemistry, Mathematics

Research Interests:
Dr. Joseph Rower is a Research Assistant Professor in the Department of Pharmacology and Toxicology. Dr. Rower completed his graduate work at the University of Colorado, Anschutz Medical Campus where his research focused on the unique pharmacology of nucleoside reverse transcriptase inhibitors used to treat and prevent HIV infection.

Dr. Rower’s current research focuses on improving the clinical care of children. Specifically, his research focuses on validating bioanalytical methods for determining drug concentrations from small blood volumes. This research has the potential to revolutionize pediatric clinical pharmacology studies, especially in neonates and other populations where blood volume is limited. These advances will enable future studies which can better define the impact of age on the pharmacokinetic-pharmacodynamic relationships. Dr. Rower is especially interested in defining these relationships for drugs used to treat infectious diseases and in transplantation.
Misty D. Smith  
Research Assistant Professor of Pharmacology and Toxicology

Email: misty.smith@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Smith-M.php

Education and Training:
- B.S., 1996, Muskingum University, OH, Biology
- Ph.D., 2001, University of South Carolina, Biomedical Sciences (Pharmacology and Physiology)

Research Interests:
Dr. Smith is a Research Assistant Professor in the Department of Pharmacology and Toxicology in the College of Pharmacy whom serves as a co-investigator in the NIH-sponsored Anticonvulsant Drug Development (ADD) Program. Her primary research interests include the evaluation and advancement of novel therapies for the treatment of epilepsy and pain. A focus of her research is the investigation of the therapeutic potential of novel compounds, such as cannabidiol, for the treatment of chronic neurological disorders and their comorbidities, as well identifying potential drug-drug interactions of cannabidiol with other concomitant therapeutics. With nearly two decades experience as a behavioral pharmacologist, Dr. Smith’s research contributes to ongoing efforts to identify and validate new animal models which can be used to screen and differentiate lead compounds. Additional related research interests of Dr. Smith include (1) the pharmacotherapeutic potential of novel neuropeptides in the treatment of chronic pain, substance abuse and other neuropsychiatric conditions and (2) sex differences in disease pathophysiology, behavior and drug responses. In addition to her contributions to the Department of Pharmacology and Toxicology in the College of Pharmacy, Dr. Smith holds a joint appointment as a Research Assistant Professor in the Oral Biology, Medicine, and Pathology Section of the School of Dentistry at the University of Utah.
Shawshank Tandon
Research Assistant Professor of Pharmacology and Toxicology

Email: shashank.tandon@utah.edu
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/shashank-tandon.php

Education & Training:
- B.Sc. (Hons.) (Human Biology), All India Institute of Medical Sciences, India, 2001
- M.Sc. (Physiology), All India Institute of Medical Sciences, India, 2003
- PhD (Neuroscience), National Brain Research Centre, India, 2010

Research Interests:
My research focuses on understanding the neural correlates of reward and aversion and how they play a role in modulating appetitive and consummatory behavior for natural rewards and drugs of abuse. To this end, I am employing electrophysiological and behavioral techniques in rats to determine the brain mechanisms mediating appetitive behavior which drives intake of highly palatable high fat/sugar food as well as drugs of abuse. Specifically, I have been interested in determining the role of opioid signaling in the nucleus accumbens (NAcc), an important node in the reward pathway, in normal feeding and binge eating behavior. My current research focuses on the brain circuitry regulating ethanol intake, with a goal of determining the role of the lateral habenula (LHb), a crucial area in which aversive stimuli are encoded, in regulating voluntary ethanol consumption during early and late stage of alcohol abuse. My long-term research goal is to determine the neural mechanism through which different factors (e.g. stress, negative affective state) can increase vulnerability to develop drug addiction as well as increase risk for chronic relapse in post-dependent state.
Peter West
Research Assistant Professor Pharmacology and Toxicology

Email: peter.west@utah.edu
Website: http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/West-P.php

Education and Training:
- B.S., 1997, Lehigh University, Biochemistry
- Ph.D., 2003, University of Utah, Neuroscience

Research Interests:
Dr. West is a staff scientist in the NIH sponsored Anticonvulsant Drug Development Program where he directs studies determining the electrophysiological mechanism of action of proprietary investigational compounds. Additionally, his research is focused on the pharmacological treatment of diseases that affect cognition such as Epilepsy, Down syndrome, and Alzheimer's disease. In order to identify novel molecular targets and test potential treatments, an understanding of the pathophysiological basis of cognitive deficit in these diseases must first be obtained and preclinical model systems must be developed. Presently, studies intended to characterize synaptic plasticity deficits and cognitive dysfunction in animal models of Epilepsy are underway. Furthermore, these projects seek to discover and characterize novel treatments to correct these deficits or to treat seizures and prevent epileptogenesis without affecting synaptic plasticity associated with learning and memory. To achieve these goals, the lab currently uses electrophysiological, pharmacological, behavioral, and immunohistochemical techniques. Of particular note, Dr. West's laboratory uses specialized equipment which allows the experimenter to perform simultaneous recordings from multiple brain slices, thus allowing for the high-throughput screening of compounds for their effects on long-term synaptic plasticity in a manner that accounts for the day-to-day variability often observed in physiological experiments that use in-vitro brain slices.
Karen Wilcox  
Professor of Pharmacology and Toxicology

Email:  kwilcox@deans.pharm.utah.edu  
Website:  http://pharmacy.utah.edu/pharmtox/faculty/current-faculty/Wilcox-K.php

Education and Training:
- B.A., 1981, Allegheny College
- Ph.D., 1993, University of Pennsylvania

Research Interests:
The Wilcox laboratory is interested in understanding basic mechanisms underlying epileptogenesis, seizure generation, and therapy-resistance to anticonvulsant drugs. To achieve these goals, we use electrophysiological, calcium imaging, pharmacological, behavioral, genetic, immunoblot, and immunohistochemical techniques in a variety of in vitro preparations and animal models of epilepsy. Our working hypothesis is that insight into disease-induced changes in neuronal and glial function will provide new avenues for therapeutic interventions in patients at risk for developing epilepsy or those patients who are refractory to current treatment options. To that end, I am also a Co-Investigator in the Anticonvulsant Drug Development (ADD) Program (Principal Investigator, H. Steve White, Ph.D.) and direct studies determining the neuroprotection potential of proprietary investigational compounds through a contract with NINDS at the National Institutes of Health. Finally, I am actively seeking rotation students as well as dissertation student.
FACULTY IN PHARMACOTHERAPY
Jennifer Babin
Assistant Professor (Clinical) of Pharmacotherapy

Email: jennifer.babin@hsc.utah.edu
Website: https://pharmacy.utah.edu/pharmacotherapy/faculty/jbabin.php

Education & Training:
- B.S., 2009, University of Alabama, Tuscaloosa, AL, Biology
- Pharm.D., 2013, Auburn University, Auburn, AL
- PGY1 Pharmacy Practice Residency, 2014, Greenville Health System, Greenville, SC
- PGY2 Internal Medicine Residency, 2015, University of Utah Health Care, Salt Lake City, UT

Research Interests:
Dr. Babin is a clinical pharmacist practicing at the University of Utah Hospital in internal medicine. Her research interests include topics relating to the care of internal medicine patients, patient education, and interprofessional education.
Joseph Biskupiak  
Research Professor of Pharmacotherapy;  
Director, Pharmacotherapy Outcomes Research Center;  

Email: joseph.biskupiak@pharm.utah.edu  
Website: https://pharmacy.utah.edu/pharmacotherapy/faculty/Biskupiak.php  

Education & Training:  
- B.S., 1979, University of Connecticut, Chemistry  
- M.B.A., 1995, Seattle University  
- Ph.D., 1985, University of Utah, Medicinal Chemistry  

Research Interests:  
Outcomes research, health services research, US health care delivery system.
Diana Brixner
Professor of Pharmacotherapy;
Executive Director, Outcomes Research Center;
Director of Outcomes, Program in Personalized Health Care;
President, Academy of Managed Care Pharmacy;
Director of the Graduate Program for the Department of Pharmacotherapy;

Email: diana.brixner@utah.edu
Website: https://pharmacy.utah.edu/pharmacotherapy/faculty/dbrixner.php

Education & Training:
• B.S., 1982, University of Rhode Island, Pharmacy
• Ph.D., 1987, University of Utah, Medicinal Chemistry

Research Interests:
Currently I am a Professor in the Department of Pharmacotherapy, Adjunct Professor in the Department of Population Health Sciences and Pediatrics in the Divisions of Health Systems Innovation Research and Clinical Pharmacology and Executive Director of the Pharmacotherapy Outcomes Research Center at the University of Utah. My research focus is on the design, conduct, and communication of pharmacoeconomic and outcomes research studies to demonstrate the value of pharmaceutical and related therapies from the perspective of the private and public payer. I am also a Research Associate at the Institute of Public Health, Medical Decision Making and Health Technology Assessment in the Department of Public Health and Health Technology Assessment at UMIT - University for Health Sciences, Medical Informatics and Technology in Hall i.T., Austria. This appointment supports my international collaborations in oncology research, personalized medicine and value assessment. During my career, I have published over 100 articles in peer-reviewed journals, authored three book chapters, have one issued patent, and have been an invited speaker at a variety of national and international professional meetings. I am a founding member, served on the Executive Board, and currently serve on the Health Policy and Science Council and am a past president of the International Society of Pharmacoeconomics and Outcomes Research (ISPOR). In addition, I am a long-standing member and immediate past president of the Academy of Managed Care Pharmacy (AMCP), and am also a Fellow of the Academy.
Nicholas Cox
Assistant Professor (Clinical) of Pharmacotherapy;
Clinical Pharmacist, Westridge Health Center, University of Utah Health;

Email: Nicholas.Cox@pharm.utah.edu
Website: https://faculty.utah.edu/u0511019-NICHOLAS_WILLIAM_COX/hm/index.html

Education & Training:
- PharmD, 2015, University of Utah, Salt Lake City, UT
- PGY-1 Pharmacy Practice Residency, University of Utah Health, Salt Lake City, UT
- PGY-2 Ambulatory Care Pharmacy Residency, University of Utah Health, Salt Lake City, UT

Research Interests:
Dr. Cox is a clinical pharmacist whose current research interests and projects involve quality improvement initiatives, chronic pain, primary care, scholarship of teaching and learning, and others.
Barbara Crouch
Professor (Clinical) of Pharmacotherapy;
Executive Director, Utah Poison Control Center;

Email: barbara.crouch@hsc.utah.edu
Website: www.utahpoisoncontrol.org
http://pharmacy.utah.edu/pharmacotherapy/faculty/crouch.php

Education & Training:
- B.S., 1982, Philadelphia College of Pharmacy and Science, Pharmacy
- Pharm.D., 1984, Jointly administered by the University of Texas at Austin and the University of Texas Health Science Center at San Antonio, Pharmacy
- M.S. MSPH, 1994, University of Utah, Public Health
- Clinical Toxicology Fellowship, Maryland Poison Center, University of Maryland School of Pharmacy

Research Interests:
My primary research involves epidemiology of poison exposures electronic health information exchange between poison centers and emergency departments and factors that affect poison center utilization. Current projects include demonstrating feasibility of electronically exchanging information between poison centers and health information exchange, analyzing a national poison database to identify trends in prazosin exposures reported to US poison centers and a study of caregivers knowledge, attitudes and behaviors about the poison center. Past research projects include the impact of communication to adherence to poison center recommendations, characterizing trends in teen OTC drug abuse in Utah and an analysis of poison exposures in older adults.
Karen M. Gunning
Professor (Clinical) of Pharmacotherapy;
Adjunct Professor of Family & Preventive Medicine;
Associate Dean of Community Engagement;
Clinical Pharmacist – Sugarhouse Health Center Family Medicine;

Email: karen.gunning@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/gunning.php

Education & Training:
- B.S., 1995, Oregon State University, Pharmacy
- Pharm.D., 1997, University of Utah
- Specialized Residency in Family Medicine, 1998, University of Washington

Clinical Specialty: Family Medicine

Research Interests:
- Population health and continuous quality improvement for chronic conditions in primary care
- Medication safety in the ambulatory care setting - particularly in pain management, and diabetes.
- Scholarship of teaching and learning as it relates to family medicine residents and pharmacy students and residents.
- Women’s and Men’s health, Care of Transgender Patients.
Holly Gurgle
Associate Professor (Clinical) of Pharmacotherapy;
Vice Chair of Teaching of Pharmacotherapy;
Clinical Pharmacist, ARUP Family Health Clinic;

Email: holly.gurgle@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/hgurgle.php

Education & Training:
- B.S., in Biochemistry, 2002-2006, The University of Virginia, Charlottesville, Virginia
- PharmD, 2006-2010, Virginia Commonwealth University, Richmond, Virginia
- PGY1 Pharmacy Practice Residency, 2010-2011, University of Washington Medicine, Seattle, Washington
- PGY2 Ambulatory/Primary Care Pharmacy Residency, 2011-2012, University of Washington Medicine, Harborview Medical Center, Seattle, Washington

Clinical Specialty: Family Medicine, Primary Care

Research Interests:
Research, teaching, and practice interests include: cardiometabolic disorders (diabetes, lipids, obesity), preventative medicine, immunizations, and innovative team-based primary care practice models.
Lauren Heath  
Assistant Professor (Clinical) of Pharmacotherapy

Email: lauren.heath@utah.edu

Education & Training:
- B.S. in biochemistry, 2010, University of Michigan – Ann Arbor, MI
- Pharm.D., 2014, University of Michigan College of Pharmacy, MI
- PGY1 Pharmacy Practice Residency, 2015, University of Southern California, CA
- M.S. in Clinical Science, 2017, University of Colorado Denver, CO
- Outcomes Research Fellowship in Ambulatory Care, 2018, Kaiser Permanente Colorado and University of Colorado School of Pharmacy, CO

Research Interests:
Research interests include health services research, implementation science, and evidence-based health care. Other interests include evaluating the impact of comorbid conditions on clinical and health care outcomes and the public and other healthcare professional's perceptions of clinical pharmacists.
Nakyung Jeon
Assistant Professor of Pharmacotherapy

Email: nakyung.jeon@utah.edu
Website: https://faculty.utah.edu/u6020442-NAKYUNG_JEON/hm/index.html

Education & Training:
- B.S., 2009, Ewha Womens University, Seoul, South Korea, Pharmacy
- M.S., 2011, Seoul National University, Seoul, South Korea, Public Health
- Ph.D., 2017, University of Florida, Pharmacy (Pharmacoepidemiology)

Research Interests:
Nakyung’s research interests focus on drug safety evaluations, and the prevention of inappropriate medication use. Clinical areas of drug safety studies include in particular pediatrics, psychiatry, and acute kidney injury. She is currently developing a research program to identify the role of psychotropic medications in unplanned pregnancy and pregnancy outcomes in teenage girls, using epidemiologic study designs and advanced analytical methods. Her interest in quality of care issues focuses on the development of clinical decision support systems (to prevent adverse drug events) and the development of quality metrics (to evaluate quality of patient care across hospitals).

Nakyung’s professional background is in pharmacy and public health. Prior to pursuing Ph.D. with concentration in pharmacoepidemiology, she practiced as a staff pharmacist in tertiary care hospitals and received master’s degree in public health with a focus on psychosocial and behavioral research. During her doctoral training, she led several projects under a large grant funded by American Society of Hospital System Pharmacists Foundation (Primary Investigator: Dr. Almut G. Winterstein) which developed 16 individual predictive models to identify patients at higher risk for different types of preventable adverse drug events (pADEs) in inpatient setting using electronic health records (EHR). Publications directly related to this project can be found here: https://www.ncbi.nlm.nih.gov/sites/myncbi/1NkPvAvH3D_Uil/bibliography/57196714/public/?sort=date&direction=ascending. Also, she worked for a Centers for Medicare & Medicaid Services (CMS) funded project, titled Inpatient Psychiatric Facility Quality Reporting Program, which develops, maintains, re-evaluates and supports implementation of quality outcome and process measures for psychiatric inpatients.
Kibum Kim  
Research Assistant Professor of Pharmacotherapy

Email: kibum.kim@pharm.utah.edu  
Website: https://faculty.utah.edu/u6008102-KIBUM_KIM/hm/index.html

Education & Training:
- 2016-2018, Post-Doctoral Fellow, Pharmacotherapy Outcomes Research Center, University of Utah, Salt Lake City, Utah
- 2011-2016, PhD in Pharmacy specialized in pharmacoeconomics and outcomes research, University of Illinois at Chicago, Chicago, Illinois
- 2006-2008, Master of Science in Pharmacy specialized in Pharmacokinetics and Pharmacodynamics, Catholic University of Daegu, Gyeongbuk Province, South Korea
- 2002-2006, Bachelor in Pharmacy, Catholic University of Daegu, Gyeongbuk Province, South Korea

Research Interests:
Kibum Kim is a Research Assistant Professor in the Department of Pharmacotherapy and at the Pharmacotherapy Outcomes Research Center, University of Utah. He graduated from University of Illinois at Chicago with his PhD in Pharmacy, specialized in pharmacoeconomics and outcomes research. Prior to working at University of Utah, he interned at Abbvie, Inc., Takeda Pharmaceuticals, and eMax Health Company where he assisted multiple market access projects. He also had years’ practical experience from community pharmacies. Dr. Kim’s research interest includes contemporary trends in medication utilization, outcome analysis using sophisticated statistical approach and model-based cost-effectiveness analysis. During his career, Dr. Kim has presented and published his researches in clinical and economic outcomes of treatment options for diabetes, cardiovascular disorders, and oncology. His recent studies focus on the clinical utility of molecular diagnostics, therapeutic drug monitoring, and precision medicine.

Main research interests are as follows:
- Utility of precision medicine and molecular diagnostic testing
- Utilization and outcomes of antiplatelet therapy and anti-coagulation management
- Glycemic control associated with anti-diabetic agent use and concurrent medication to prevent patient from long-term complications
- Resource utilization and healthcare cost across multiple chronic conditions
- Research methods in health outcomes research using real-world databases
Joanne LaFleur
Associate Professor of Pharmacotherapy

Email: joanne.lafleur@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/Lafleur.php

Education & Training:
- B.S., 2001, University of Utah, Salt Lake City, UT, Pharmacy Practice
- Pharm.D., 2003, University of Utah, Salt Lake City, UT, Pharmacy Practice
- M.S.P.H., 2005, University of Utah, Salt Lake City, UT, Public Health

Research Interests:
- Pharmacoeconomics Outcomes research
- Patient adherence and persistence
- Medicaid and public health policy
- Chronic pain syndromes
- Primary and secondary prevention of cardiovascular disease
- Risk factors for osteoporotic fracture
Joanita Lake
Research Assistant Professor of Pharmacotherapy

Email: joanita.lake@pharm.utah.edu
Website: https://pharmacy.utah.edu/pharmacotherapy/about-us/staff/lake.php

Education & Training:
- B.Pharm, 2000, University of Port Elizabeth, South Africa, Pharmacy
- MSc (Oxon), 2008, University of Oxford, U.K., Evidence-based Health Care (EBHC)

Research Interests:
Her research interests lie primarily in the areas of evidence-based health care, drug safety, drug utilization review, prescribing practices, pharmacoepidemiology, pharmacoeconomics, and health policy. Her recent research projects include systematic reviews on opioid misuse and abuse. She is a clinical pharmacist at the Drug Regimen Review Center (DRRC).
Alisyn May
Assistant Professor (Clinical) of Pharmacotherapy

Email: alisyn.may@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/ahansen.php

Education & Training:
- B.S., 2011, in Chemistry/Health Sciences, Wayne State College, Wayne, NE
- PharmD, 2013, University of Nebraska Medical Center, Omaha, NE
- PGY1 Pharmacy Practice Residency, 2013-2014, University of Utah Hospitals & Clinics, Salt Lake City, UT
- PGY2 Ambulatory Care & Family Medicine Residency, 2014-2015, University of Utah Hospitals & Clinics, Salt Lake City, UT

Clinical/Research Interests:
- Research in the area of ambulatory care, family medicine, and chronic disease state management.
- Interested in studying the role of pharmacist in the primary care setting, novel ways to engage patients in their healthcare, and resiliency & burnout in clinical and faculty pharmacists.
- Research interest in transitions of care, resiliency/burnout, shared decision making, diabetes, chronic disease state management
Krystal Moorman  
Assistant Professor (Clinical) of Pharmacotherapy

Email: krystal.moorman@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/kmoorman.php

Education & Training:  
- PharmD, 2000-2004, Medical University of South Carolina  
- PGY-1 residency, 2004-2005, Medical University of South Carolina  
- PGY-2 residency in Drug Information Practice, 2005-2006, Medical University of South Carolina

Clinical Specialty: Community Practice

Research Interests:  
Dr. Moorman has research interests in medication safety, medication therapy management, community pharmacy practice, and drug information.
Mark Munger
Professor of Pharmacotherapy;
Adjunct Professor, Internal Medicine (Cardiology);
Associate Dean, College Affairs, CoP;

Email: mmunger@hsc.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/mmunger.php

Education and Training:
- B.S.(Pharmacy), 1980, Oregon State University
- Pharm.D., 1984, University of Illinois at Chicago
- Clinical Pharmacology Research Fellowship, 1988, Case Western Reserve University School of Medicine

Research Interests:
The Munger Research Group is focusing on repurposing drugs to address major public health concerns, unique value propositions and differentiation, with the potential for commercial product value. Currently we have two drugs in research development.

1. NSAID-induced cardiovascular and renal adverse effects are associated with the mortality of approximately 8 million lives yearly. Our research has focused on reducing these effects by biologically affecting the underlying drug-induced physiology. Through combining a prostaglandin E2 analog with the NSAID diclofenac we have been able to beneficially reset cardiovascular and renal hemodynamics; and to show that cardiovascular and renal induced events are statistically and clinically improved. This agent is currently in the pre-IND stage of FDA approval.

2. We have also developing a drug for the treatment of atrial fibrillation. In collaboration with the The Ohio State University and colleagues in Italy we have developed an agent that physiologically affects Na⁺ channels which co-localize with ryanodine receptors Ca²⁺ release channels and the Na⁺/Ca²⁺ exchanger. These findings have translated into a clinical setting of a community-based historical cohort which revealed that patients treated with the agent evidenced lower incidence of AF.
Nancy Nickman
Professor of Pharmacotherapy

Email: nancy.nickman@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/nnickman.php

Education & Training:
- RPh License, 1982, Minnesota; 2011 Utah
- B.S., 1982, University of Montana, Pharmacy
- ASHP Accredited Administrative Residency 1982-1984, United & Children’s Hospital
- M.S., 1984, University of Minnesota, Hospital Pharmacy
- Ph.D., 1987, University of Minnesota, Social & Administrative Pharmacy

Research Interests:
Dr. Nickman’s teaching and research expertise includes patient-centered outcomes research to improve provision and quality of patient care based on application of industrial engineering and economic analyses. Publications and presentations include work sampling evaluations of institutional pharmacy services for re-design of structural and functional activities, pre-post analyses of the impact of technology on the practice of health professionals, simulated time-and-motion evaluations coupled with microcost analyses of issues related to medication preparation and administration, and analysis of devices intended for medication self-administration. Since 2012, Nancy has served as a Clinical Coordinator for Analytics and Outcomes for Pharmacy Services, University of Utah Health Care. For 2016-17, she is serving as co-Chair of the ASHP Section on Pharmacy Informatics and Technology (SOPIT) Pharmacy Operations Automation (POA) Section Advisory Group (SAG). From 2013-2016, she co-chaired the “Automation of the Pharmacy Enterprise” subgroup of the POA SAG and has been a POA member since 2010.
Heather Nyman  
Assistant Professor (Clinical) of Pharmacotherapy

Email: heather.nyman@hsc.utah.edu  
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/nyman.php  
Education & Training:  
- B.S., 2000, Brigham Young University, Provo, UT, Chemistry.  
- Pharm.D., 2004, University of North Carolina at Chapel Hill, Chapel Hill, NC  
- PGY1 Pharmacy Practice Residency, 2005, University of Utah Hospitals and Clinics, Salt Lake City, UT  
Research Interests:  
Dr. Nyman is a clinical pharmacist practicing at the University of Utah Hospital in internal medicine. Her research interests are in the area of nephrology, including management of anemia of chronic kidney disease and measures of kidney function for drug dosing. Current projects include an analysis of data from the VA system to compare drug dosing when using the CKD-EPI, MDRD, and Cockcroft-Gault equations.
Patricia Orlando
Associate Professor (Clinical) of Pharmacotherapy

Email: patricia.orlando@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/orlando.php

Education & Training:
- B.S. Pharmacy, 1982, University of Montana
- Pharm.D., 1985, University of Utah
- Pharmacy Practice Residency, 1985, University of Utah Hospital,
- Residency, 1986, SLC Veterans Affairs Medical Center, Geriatric Pharmacy
- Fellowship, 1987, UCLA, Infectious Diseases Pharmacotherapy

Clinical Specialty:
Geriatric Infectious Disease Pharmacotherapy
Dr. Orlando is a clinical pharmacy specialist for Infectious Diseases at the George E. Wahlen Veterans Affairs Healthcare System in Salt Lake City, Utah

Research Interests:
Role of pharmacy interventions in outpatient parenteral antimicrobial therapy (OPAT) and transitional care for the geriatric patient with infection requiring extended therapies; Antibiotic pharmacy monitoring systems for OPAT; Antibiotic education models for geriatric patients with infections; Pharmacotherapy role of pharmacy students in the management of geriatric infections.
Hanna Raber  
Assistant Professor (Clinical) of Pharmacotherapy

Email: hanna.raber@pharm.utah.edu  
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/hraber.php

Education and Training:
- Pharm.D., 2014, Drake University, Des Moines, IA
- PGY-1 and PGY-2 Ambulatory Care Residency, 2016, Saint Joseph Regional Medical Center, Mishawaka, IN

Research Interests:
Dr. Raber is a clinical faculty member who provides patient care at the University of Utah Centerville and Madsen Family Medicine Residency Clinics. Her research interests include health communication, global health, and preventative medicine.
James Ruble
Associate Professor (Clinical) of Pharmacotherapy

Email: jim.ruble@hsc.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/ruble.php

Education & Training:
- B.S., 1989, University of Utah College of Science, Biology
- B.S., 1992, University of Utah College of Pharmacy, Pharmacy
- Pharm.D., 1994, University of Utah College of Pharmacy
- J.D., 2002, University of Utah College of Law

Clinical Practice and Research Interest:
- Pharmaceutical compounding, including USP Chapters <795>, <797>, <800>
- Ethical dilemmas in health care
- Moral reasoning, moral distress, and moral fatigue in health care professionals
- Communication and Conflict Management in Health Care
- Legal requirements in health care, professional standards of care and risk management
- Intellectual Property and equipoise in healthcare and pharmaceutical research
Natalia Ruiz-Negrón  
Research Assistant Professor of Pharmacotherapy

Email: Natalia.Ruiz@pharm.utah.edu  
Website: https://faculty.utah.edu/u0710482-NATALIA_RUIZ-NEGRON/hm/index.html  

Education & Training:  
- B.S. Chemistry (biology emphasis), 2012, University of Utah, Salt Lake City, UT  
- Pharm.D., 2016, University of Utah, Salt Lake City, UT  
- Health Economics & Outcomes Research Fellowship, 2018, Pharmacotherapy Outcomes Research Center, University of Utah, Salt Lake City, UT

Research Interests:  
- Pharmacoepidemiology  
- Health disparities research  
- Pharmacoeconomics  
- Health outcomes modeling  
- Chronic diseases, including diabetes, hypertension, and hyperlipidemia
Linda Tyler
Professor (Clinical) of Pharmacotherapy;
Associate Dean, Pharmacy Practice;
Chief Pharmacy Officer, University of Utah Health Care;

Email: Linda.Tyler@hsc.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/tyler.php

Education & Training:
- B.S., 1978, University of Utah, Pharmacy
- Pharm.D., 1981, University of Utah
- Residency, 1978-1979, University of Nebraska Medical Center, Hospital Pharmacy

Research Interests:
Dr. Tyler's interests include health system pharmacy administration, drug information, medication safety, medication use and policy development, adverse drug reactions, drug shortages, literature evaluation, evidence based medicine.

Dr. Tyler serves as residency program director for the Health System Pharmacy Administration Residency (2 year program with MS degree) and co-director of the MS in Health System Pharmacy Administration.
Kyle Turner
Assistant Professor (Clinical) of Pharmacotherapy

Email: Kyle.Turner@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/kturner.php

Education & Training:
- PharmD, 2010-2014, University of Utah College of Pharmacy
- Pharmaceutical Care Leadership Residency, 2014-2016, University of Minnesota College of Pharmacy

Research Interests:
Dr. Turner is a clinical pharmacist practicing in primary care at the University of Utah Westridge Clinic. His research interests include development and impact of primary care pharmacy services, continuous quality improvement, health professional leadership development. He currently helps lead the University of Utah Interprofessional Education program’s Student Hotspotting program which engages student team with patient who have high health care utilization in an effort to reduce cost and improve outcomes.
Daniel M. Witt
Chair and Professor (Clinical) of Pharmacotherapy;
Assistant Dean for Clinical Affairs;

Email: dan.witt@pharm.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/witt.php

Education & Training:
- A.S., 1984, Snow College, Ephraim, Utah
- B.S., 1990, University of Utah, Salt Lake City, Utah, Pharmacy
- Pharm.D., 1992, University of Washington, Seattle, Washington

Research Interests:
I have been actively involved in conducting mentored research projects for more than 20 years, with the principal areas of focus being 1) documentation of the clinical and economic impact of clinical pharmacy services; 2) providing optimized management of anticoagulation therapy; and 3) providing practical evidence-based solutions to anticoagulation therapy stakeholders, including issues pertaining to anticoagulation therapy nonadherence. I joined the faculty at the University of Utah College of Pharmacy in large part to expand anticoagulation therapy research opportunities, foster additional collaborative research relationships and teams, and expand my opportunities to mentor investigators in training. An example of recent success in this regard is the Transitions of Care for Venous Thromboembolism project. The dataset created for this project is currently supporting seven research projects for PharmD students. In addition, I am a founding member of the Clinical Pharmacy Research Group (CARGO) which includes researchers from Kaiser Permanente Colorado, Intermountain Health Care, the Veterans Administration, McMaster University, the University of Michigan, and the University of Utah. CARGO has outlined an aggressive agenda of research projects, each intended to include researchers in training, who will receive mentoring from members of the group.
David Young
Professor (Clinical) of Pharmacotherapy

Email: david.young@hsc.utah.edu
Website: http://pharmacy.utah.edu/pharmacotherapy/faculty/young.php

Education & Training:
• Pharm.D., 1995, Idaho State University, Pocatello, ID, Pharmacy
• Primary Care Specialty Residency, 1996, Idaho State University, Boise, ID

Research Interests:
Dr. Young is the clinical pharmacist at the University of Utah Adult Cystic Fibrosis and Asthma Centers. Dr. Young a member of the North American Cystic Fibrosis Foundation Annual Meeting Planning and Clinical Research Committees. Dr. Young is the Co-director of the North American Cystic Fibrosis Foundation Pharmacist Mentorship Program. Dr. Young is the coordinator for PGY-2 Internal Medicine Pharmacy Residency at the University of Utah Health. Dr. Young is actively engaged in research regarding improving care for patients with cystic fibrosis and asthma.