

SRI 2016
UC Santa Cruz Faculty

This is a list of faculty members and a brief summary of their research interests. For more information on the faculty listed below, please check the following websites.

<http://www.chemistry.ucsc.edu/faculty/index.html>

<http://www.mcd.ucsc.edu/faculty.html>

<http://www.metx.ucsc.edu/faculty/index.html>

<http://www.bme.ucsc.edu/people/faculty>

MARK AKESON, PROFESSOR OF BIOMOLECULAR ENGINEERING

Nanopores in lipid-bilayer membranes have the potential to be used to characterize DNA and RNA. Mark Akeson's primary research area concerns the manner in which these linear macromolecules traverse nanoscopic channels. He hopes to apply these studies of DNA structure and dynamics to HIV treatment and personal genomics.

ALEX AYZNER, ASSISTANT PROFESSOR OF CHEMISTRY

We are broadly interested in understanding the influence of molecular and macromolecular structure of small-molecule and polymeric organic semiconductors on the efficiency and dynamics of light harvesting. Current work in the group aims to address the three fundamental steps involved in photoelectric and photochemical conversion: energy transfer, interfacial electron transfer, and long-range charge transport.

NEEDHI BHALA, ASSISTANT PROFESSOR OF MCD BIOLOGY

We are interested in the mechanisms that ensure that chromosomes segregate correctly during cell division, particularly in meiosis. Defects in meiosis can generate gametes with the incorrect number of chromosomes and typically produce inviable embryos. In some cases, the presence of an extra copy of a chromosome results in serious developmental disorders, such as Down and Klinefelters syndrome. We combine genetic and biochemical approaches with high-resolution microscopy and cytological techniques to gain a more informed view of how molecular events during meiosis govern and are governed by higher-order chromosome behavior.

REBECCA BRASLAU, PROFESSOR OF CHEMISTRY

Our research group is focused on reactions involving free radical intermediates. Much of the work involves the synthesis and use of designed nitroxides for the preparation of specialized polymers for applications in nanotechnology, and as sensors via fluorescence quenching. We are also engaged in the development of new synthetic methodologies via free radical intermediates.

MANEL CAMPS, ASSISTANT PROFESSOR OF MICROBIOLOGY & ENVIRONMENTAL TOXICOLOGY

Spontaneous DNA Methylation results from methyl donors reacting with DNA. DNA methylation is a potent carcinogen. Paradoxically, in addition to being carcinogenic, methylating agents are also mainstays for cancer treatment. Among his research interests, Dr. Camps examines the molecular mechanisms of methylating agent toxicity to design safer and more effective strategies for cancer chemotherapy.

SHAOWEI CHEN, PROFESSOR OF CHEMISTRY

This research is centered around electron transfer chemistry at the nanoscale; more specifically, we are interested in the electron transfer properties of nanometer-sized particle molecules and their organized assemblies. Our strategy is to employ a series of chemical as well as physical manipulations to shed light onto the molecular origin of these unprecedented electrochemical phenomena. Currently there are three major research projects in my laboratory: 1) rectification of nanoparticle quantized charge transfer, 2) solid-state electronic conductivity of nanoparticle ensembles, 3) magnetochemistry of nanoparticle quantized charging.

PHILLIP CREWS, DISTINGUISHED PROFESSOR OF CHEMISTRY

Application of nuclear magnetic resonance to problems of organic structure, marine organic chemistry – isolation and structure determination of biologically important substances, especially for marine sponges and their associated microorganisms.

OLOF EINERSDOTTIR, PROFESSOR OF CHEMISTRY

Cytochrome oxidase is an oxidizing enzyme found in mitochondria and important in cell respiration as an agent of electron transfer from certain cytochrome molecules to oxygen molecules. It is the last enzyme in the respiratory electron transport chain. The primary aim of my research is to understand the mechanism of coupled electron transfer and proton translocation in cytochrome oxidases.

RUSS FLEGAL, PROFESSOR OF ENVIRONMENTAL TOXICOLOGY

My research is on the natural biogeochemical cycles of trace elements in the environment and the perturbation of those cycles by anthropogenic processes. The primary focus is on the biogeochemical cycle of lead in preindustrial and contemporary environments. This includes studies of lead cycles in the world's oceans, Mediterranean Sea, Great Lakes, European Alps, and Antarctic. I am also involved with research on the biogeochemical cycling of other trace elements in aquatic systems. This is focused on the cycling of heavy metals and rare earth elements in the San Francisco Bay estuarine system and the northeast Pacific coastal zone. These studies involve analyses of elemental concentrations and isotopic compositions in aerosols, water, sediments, and organisms.

CAMILLA FORSBERG, ASSISTANT PROFESSOR OF BIOMOLECULAR ENGINEERING

The Forsberg lab focuses on stem cell fate decisions of the blood system, seeking to understand their molecular determinants in order to prevent and treat both genetic and acquired disorders of the hematopoietic system. We tackle questions regarding these decisions from multiple angles - by *in vivo* and *in vitro* experimental approaches, by focusing on specific molecules as well as analyzing global changes.

GRANT HARTZOG, PROFESSOR OF MCD BIOLOGY

1) We are interested in understanding the role chromatin plays in gene expression and the mechanisms by which chromatin structure is manipulated to regulate transcription. The spatial arrangement of nucleosomes on a gene influences its transcription by modulating the ability of regulatory proteins to access specific DNA sequences and by modulating the rate at which RNA polymerase travels down genes. We study this problem in the yeast, *Saccharomyces cerevisiae*, using biochemistry and genetics. 2) We are also interested in isolating new bacteriophage (bacterial viruses) which may lead to identification of novel genes and proteins. In addition, new bacteriophage may provide tools enabling study of their bacterial hosts.

THEODORE HOLMAN, PROFESSOR OF CHEMISTRY AND BIOCHEMISTRY

Bioinorganic Chemistry. Many critical processes require metal ions, such as respiration, photosynthesis and signal transduction. Our research group utilizes enzymology, site directed mutagenesis, molecular biology, inorganic chemistry, and spectroscopy to investigate the biological function of metalloproteins from a rigorous chemical perspective. Specifically, we study the biological function of lipoygenase, an enzyme in humans that has been implicated in a variety of diseases (asthma, atherosclerosis, and cancer), in order to understand its mechanism better and discover novel inhibitors.

MELISSA JURICA, PROFESSOR OF MCD BIOLOGY

My lab investigates the cellular machinery responsible for editing the information contained in the RNA transcripts of nearly all of human genes. This machinery, called the spliceosome, splices out intron sequences that interrupt gene transcripts and joins exon sequences to make messenger RNAs that correctly encode for proteins. The goal of our research is to understand how the spliceosome is assembled and how it catalyzes the splicing reaction.

ROHINTON KAMAKAKA, PROFESSOR OF MCD BIOLOGY

Research in our laboratory is devoted to understanding the mechanism by which the genome is partitioned into structural and functional units. We employ molecular and genetic analysis coupled with biochemical experiments to explore the issues of genome organization. We are presently working on 1) the architecture of the silenced chromatin domains and 2) the mechanism by which chromatin domains are delimited.

DOUG KELLOGG, PROFESSOR OF MCD BIOLOGY

Cells show extraordinary diversity in size and shape. Generation of diverse sizes and shapes requires regulation of the amount and location of growth, as well as coordination of cell growth with cell division. The mechanisms by which cells regulate cell growth and size are poorly understood and represent one of the most important and fundamental unsolved problems in cell biology. The goal of our work is to elucidate these mechanisms.

DAVE KLIGER, DISTINGUISHED PROFESSOR OF CHEMISTRY

My research group spans the fields of physical chemistry and biophysics. We have been involved in developing a variety of spectroscopic techniques and applying them to a wide range of photochemical, photophysical, and photobiological problems. These techniques provide a powerful set of tools for studying molecular dynamic processes such as: 1. the mechanism of activation of visual pigments, 2. the mechanisms of function of the plant regulatory protein phytochrome and a variety of heme proteins, and 3. the early events in the folding of a variety of proteins and DNA.

JEREMY LEE, LECTURER OF MCD BIOLOGY

I collaborate with colleagues at Drexel University in a research program to identify regulators of the amyloid precursor protein (APP), a central protein in Alzheimer's disease pathology. APP, which is expressed in neurons and other cell types, normally undergoes proteolytic cleavage; among the products of this cleavage is the amyloid beta (A β) peptide. In individuals with Alzheimer's disease, the levels of this peptide are increased above normal levels and the excess forms amyloid plaques. Our goal is to identify factors that regulate cleavage of APP and generation of the A β peptide.

YAT LI, ASSISTANT PROFESSOR OF CHEMISTRY

The primary aim of my research is to address fundamental science in low-dimensional materials through bottom-up paradigm, specifically consists of three interrelated areas including (i) design and rational synthesis of functional nanomaterials, (ii) investigation of their fundamental properties and (iii) use these novel nanomaterials to address the scientific issues in photovoltaic, nanoscale photonics and electronics.

SCOTT LOKEY, PROFESSOR OF CHEMISTRY

The primary goal is to develop chemical tools with which to study complex biological processes. We have three specific areas of interest: 1. the synthesis of combinatorial libraries inspired by cyclic peptide natural products; 2. the study of membrane permeability using cyclic peptide model systems; 3. the development of cell-based assays to identify novel compounds with potent biological activity.

GLENN MILLHAUSER, PROFESSOR OF CHEMISTRY AND BIOCHEMISTRY

Research in our laboratory ranges from physical chemistry to biochemistry. Three examples of our research are as follows: 1. Understanding how the primary sequence of peptides controls secondary structure in solution. 2. Use of ESR to monitor kinetic processes involved in the production of amyloid - a protein-rich deposit that appears to be the causative agent in Alzheimer's Disease and, perhaps, in the prion diseases such as Mad Cow Disease. 3. Use of NMR to solve the solution structures of small proteins; in particular, human agouti related protein (AGRP), a signaling molecule that plays a major role in the control of hunger and fat deposition.

SCOTT OLIVER, PROFESSOR OF CHEMISTRY

The Oliver research group deals with several ongoing materials chemistry projects. ACCESS students would join our efforts on the synthesis of new inorganic materials. We are working towards the discovery of new mineral-like structures with a positive charge, for environmental application as adsorbents of anionic pollutants.

KAREN OTTEMAN, PROFESSOR OF MICROBIOLOGY

The Ottemann laboratory investigates how bacteria translate chemical and physical cues in their host environment into a response that allows them to colonize a mammalian host. Our lab members are particularly interested in how pathogens use swimming during infection, using the bacterium *Helicobacter pylori* as a model for this ability. Their chemotaxis system allows us to examine the types of cues sensed by mammalian pathogens.

CARRIE PARTCH, ASSISTANT PROFESSOR OF CHEMISTRY AND BIOCHEMISTRY

Our goal is to identify the fundamental basis of crosstalk between the circadian clock and DNA damage checkpoint response to understand how disruption of the clock regulates oncogenesis. We will achieve this by integrating diverse experimental approaches from cell biology to solution NMR spectroscopy to gain an atomic-level understanding of the clock, as well as providing new targets and/or a temporal basis for improved therapeutic intervention in cancer.

NADER POURMAND, ASSISTANT PROFESSOR OF BIOMOLECULAR ENGINEERING

The Pourmand lab develops genome sequencing technology and uses high-throughput sequencing and bioinformatics to understand genetic variation and its function in health, disease, and biological systems (UCSC Genome Sequencing Center). They also develop nanoscale sensors that can be used for biological research at the cellular level (Biosensors and Bioelectrical Technology group).

SETH RUBIN, ASSOCIATE PROFESSOR OF CHEMISTRY AND BIOCHEMISTRY

Our research interests are in understanding the biochemical mechanisms that control the eukaryotic cell cycle. Our lab seeks to elucidate the biochemical determinants of protein interaction affinity and specificity and how these factors are affected by regulatory modifications to protein composition and structure. We apply a variety of structural and biochemical techniques to attain atomic resolution structures of protein complexes and to learn how structural changes and chemical modifications affect biological function.

CHAD SALTIKOV, ASSOCIATE PROFESSOR OF MICROBIOLOGY AND ENVIRONMENTAL TOXICOLOGY

My lab focuses on understanding how microbial anaerobic respiratory processes influence the fate and transport of pollutants in the environment with a specific interest in the metalloid arsenic. Microbial metabolic activity is increasingly thought to play a significant role in controlling the fate and transport of toxic metals in aquatic systems. In the case of arsenic, two primary redox reactions have been described: (i) the oxidation of arsenite (As(III)) to arsenate (As(V)) and (ii) the reduction of As(V) to As(III). Current research projects are directed at: (i) constructing a whole-cell model for how bacteria respire As(V), (ii) investigating the diversity of the *arrA* functional gene, and (iii) applying this molecular biological information to understanding how microbial redox processes affect the As biogeochemical cycle.

DON SMITH, PROFESSOR OF ENVIRONMENTAL TOXICOLOGY

Our research seeks to understand the mechanistic basis and functional outcomes of metal toxicity. Our efforts have centered around the effects of lead in model systems and humans, with emphasis on the study of therapeutic treatments for lead poisoning. We have also become quite interested in the neurotoxicology of manganese and other redox active metals, and how they contribute to neurologic disease. We use an array of analytical, biochemical, and molecular techniques to investigate basic mechanisms of action at the biochemical/molecular level, and the functional outcomes at the organ and whole organism level.

MICHAEL STONE, ASSISTANT PROFESSOR OF CHEMISTRY AND BIOCHEMISTRY

Using single molecule techniques, the Stone laboratory studies the structure and function of telomerase, an enzyme that synthesizes telomere DNA. Telomeres are specialized chromatin structures that prevent deleterious chromosome fusion events by differentiating normal chromosome ends from sites of DNA damage. Since telomerase activation is restricted to rapidly dividing cell types (such as stem cells and the majority of human tumors), it is of direct medical importance to understand fundamental mechanisms governing its assembly and function.