


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To know more. The Department of Pharmacology has a proven Landmarks produce lÁderes academics and industrial research in biome © tip across the counry and world. Our college gifted in cutting-edge research, covering many Áreas, including: QuÁmicaMunologiaClÁnicaViroláÁgicaViroláÁgicaCancerCiencieviEsciÁnciasCenÂ Biology © optical SensoresCÁªmicosDrug Metabolismnetz Arroyo, Ph.D., Assistant EsforÂso eletroquÁmico inspired by biology to study the fate of the spring © cells in the body. Our lab focus on the development of biosensors for eletroquÁmicos detecÂŠÁ ¢ contÁnua the spring © small cells and physiologically important in vivo. This seek studying receptors and interaÂŠÂpes biofÁsicas between targets, developing mechanisms f transduÂŠÁ the signal to produce trophic © elast readings, and designing and manufacturing devices to be implanted directly into the body. Using these questÂpes settled into farmacodinÁ © tica of drugs and farmacodinÁ ¢ mica, toxicology and metabolism. Our goals overaching sÁ ¢ oo development platforms to achieve the delivery of drugs and the diagnosis responsive devices metabolism and diagnosis devices for health care personalizadosJames C. Barrow, Ph.D., Ph.D., Associate Professordrug discovered by distÁªrbios of neurodevelopment. The research group Á © one lab focused on Chemicals medicines and drug discovery, addressing mainly diseases neurodevelopmental such as schizophrenia. The lab estÁª involved in the design and sAntese spring © bioÁÁgica cells for a particular target, for Analyzing in vitro and in vivo, and further refinement atravÁ © s vÁªrios sAntese and testing cycles. The resulting avanÂšados leads will have the ¢ good potÁncia and selectivity for the target of interest, and will be used the ¢ á á to test hipÁªteses bioÁÁgicas both in vitro and in vivo to determine whether modulating the target Á © indeed one Estrata © gia terapÁªutica viÁvel. When running in medicinal chemistry programs, we direct the translational ciÁªncia from the target and pathwaynamandÁÁ ¢ -bumpus, Ph.D., director and E. K. Marshall and Thomas H. Maren Professordrug Metabolism and prÁª © drug development -clÁnico; mass spectrometry spring © small cell; metabolÁmica segmented; Antiviral drug-induced toxicity; Modulation of the sinalizaÂŠÁ ¢ ¢ pathways for cell lab metabÁªlitos reativos.O of Bumpus applies proteÁmica, metabolÁmica, and Ta © imaging techniques to understand molecular mechanisms underlying inter-individual variability in drug results, including toxicities. By doing this, we define the metabolism of clinically relevant drugs as elucidated the impact that drugs and their metabÁªlitos tÁªm in sinalizaÂŠÁ ¢ cell. Our work uses in vitro and in Vivo prÁª © -clÁnico models, as well as in clinics Analyzing samples. The overarching goal of our work Á © The mechanistic understanding of drug metabolism, the impact of cellular drug metaboles and three-dimensional distribution of cells and tissues to facilitate the medicine. Greek V. Carr, Ph.D., Assistant Professoratreclinical models of neurological and psychiatric distancies. Psychiatric disturbances and related neurodevelopments disturbments result from the complex interaction of multiple genetic and environmental risk factors. Our laborator uses rodent models to determine how these risk factors modulate neurobiology and behavior. We use neurochemical, electrophysiological and behavioral assays to interrogate neural systems considered to be impaired in neurodevelopmental distancies. Our ultimate goal is to use this knowledge to inform drug discovery efforts designed to increase and improve the treatment options available for central nervous system disturbancesJun O. Liu, Ph.D., teacherscourish biology Molecular biology: use of small molems as probes to elucidate signal transduction mechanisms: Angiogenesis and cellular proliferation. Our primary research research resides in the interface between chemistry, biology and medicine. We employ high-performance screening to identify modulators of various processes and cells that were involved in human diseases of autoimmune diseases. Once identified biologically active compounds, serve as both the probes of biological processes of interest and leads to the development of new drugs for the treatment of human diseases.Caren L. Freel Meyers, Ph.D., Ph.D., Chemical Professional, Chemical Biology: Drug Delivery; Study of the biosynthesis isopreneic no-mamma; Development of possible therapeutic agents for cyanx and infectious disease. Freel Meyers Lab applies organic synthesis, chemical biology and enzymology to design new anti-infectious approaches. A research area concentrates on the development of strategies to selectively inhibit enzymes in the essential way of bacterial MEP for isoprenoids. In addition, we study the mechanism and function of DXP Synthase, the first enzyme on this pathway and christian enzyme of the affiliate in bacterial metabolism, to understand their metabolic papers during infection. We will also bring our experience in a small synthesis of molemles and prodrug development to a multi-institutional collaboration to invent new long-term nanoformulations of clinically used antiretrovirals - ¢

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