

Denigrating & Discriminatory Study Staff Behaviors Harming Patient Enrollment Diversity

Tufts Csdd Impact Report

L Darling-Hammond

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Broadly Engaged Team Science in Clinical and Translational Research Debra Lerner, Marisha E. Palm, Thomas W. Concannon, 2022-02-27 Despite the large U S investment in health science and the vast and growing body of peer reviewed research findings it has produced a compelling body of evidence suggests that research too often has been slow inefficient and fallen short of desired impacts on health A key question is how research might be changed to be more innovative less wasteful and more responsive to unmet health needs One emerging response within clinical and translational science is to advance an approach that attempts to close the gap between research scientists and key stakeholders the individuals and groups responsible for or affected by health related decisions Broadly engaged team science promises to support this aim by transforming the gold standard multi disciplinary team science to include key stakeholders in activities across the research spectrum These new roles and responsibilities range from generating research questions to implementing research projects to aiding in the translation of discoveries from the laboratory to the community A transition to broadly engaged team science reflects the idea that inclusivity and a diversity of perspectives are necessary to achieving progress in addressing complex health issues while representing a new benchmark for ethical research practice. This is one of the first collections of papers describing how clinical and translational science researchers are defining and implementing new research practices and the successes and challenges involved This book represents a first and critical step towards organizing knowledge of broadly engaged team science and advancing the development of evidence based practices Written in an accessible style this book is intended to highlight the breadth of broadly engaged team science within one community motivate researchers and stakeholders to build inclusive teams bring rigor to often informal stakeholder engagement research practices and encourage people to think more broadly about the development of scientific knowledge It includes examples of multi disciplinary broadly engaged team science projects the perspectives of academic leaders about the changes needed to encourage scientists to conduct broadly engaged team science and a resource directory Pediatric Drug Development Andrew E. Mulberg, Steven A. Silber, John N. van den Anker, 2011-09-20 Pediatric Drug Development Concepts and Applications is designed as a reference and textbook and is meant to address the science of differences between the pediatric and adult subject in the development of pharmaceutical products Considered are the ethics and medical needs of proper understanding the pediatric and adult differences the business case for proper development of drugs for children as well as the technical feasibility studies and processes that are necessary for a proper pediatric drug development program The applications of these approaches will benefit all stakeholders and ultimately not only educate but also provide better and safer drugs for pediatric patients Global Clinical Trials Playbook Menghis Bairu, Richard Chin, 2012-04-20 Pharmaceuticals companies biotech companies and CROs regardless of size all face the same challenge of managing costs and operational execution associated with bringing a valuable drugs and devices to market Because of timeline pressures and cost as well as the growing interest

in neglected diseases and diseases affecting the emerging nations clinical trials are increasingly conducted in emerging markets and developing countries where infrastructure leadership skilled personnel and a governance are at a premium Working with academics regulatory professionals safety officers experts from the pharma industry and CROs the editors have put together this up to date step by step guide book to building and enhancing global clinical trial capacity in emerging markets and developing countries This book covers the design conduct and tools to build and or enhance human capacity to execute such trials appealing to individuals in health ministries pharmaceutical companies world health organizations academia industry and non governmental organizations NGOs who are managing global clinical trials Gives medical professionals the business tools needed to effectively execute clinical trials throughout the world Provides real world international examples which illustrate the practical translation of principles Includes forms templates and additional references for standardization in a number of global scenarios **Re-inventing Drug Development** Jeffrey S. Handen, 2014-10-28 The biopharmaceutical industry has entered an era of unprecedented change and challenge characterized by increasing pricing pressures rising rates of attrition in the product development lifecycle and decreasing scientific innovation. The most successful products are losing patent protection and pipelines have been unable to fill the gap Fundamentals of Drug Development Jeffrey S. Barrett, 2022-07-13 Fundamentals of DRUG DEVELOPMENT Enables This readers to understand the process of pharmaceutical research its regulatory basis and how it fits into the global healthcare environment This book discusses how to conduct pharmaceutical research and the context for how the industry fits into global healthcare Holistically the well qualified author helps readers and students of drug development appreciate the time and expense of the process Specifically the work identifies the emerging trends shaping the future of drug development along with important related topics like generic drugs data sharing and collaboration To aid in seamless reader comprehension the book includes a glossary of terms and a self assessment guiz for each chapter at the end PowerPoint slides are also available as an online ancillary for adopting professors Sample topics covered in the book include Drug development and its phases Decision making processes drug development milestones and compound progression metrics The various disciplines involved along with an assessment of the complexity and risks associated across the stages of development Differences in the nature and scope of development programs due to the therapeutic area of interest Associated costs and resources required Graduate students and professors teaching courses in drug development drug discovery pharmaceuticals medicinal chemistry and drug synthesis will be able to use this book as a complete resource for understanding all the complexities and nuances involved in the drug development process Annual Reports in Medicinal Chemistry, 2010-11-25 Annual Reports in Medicinal Chemistry provides timely and critical reviews of important topics in medicinal chemistry together with an emphasis on emerging topics in the biological sciences which are expected to provide the basis for entirely new future therapies Covers findings related to cardiovascular inflammation and pulmonary diseases

Examines issues in oncology from mTor inhibitors to drug targets Incorporates up to date information on drug design and discovery including delivery to market **Phenotypic Drug Discovery** Beverley Isherwood, Angelique Augustin, 2020-12-09 Phenotypic drug discovery has been highlighted in the past decade as an important strategy in the discovery of new medical entities How many marketed drugs are derived from phenotypic screens From the most recent examples what were the factors enabling target identification and validation This book answers these questions by elaborating on fundamental capabilities required for phenotypic drug discovery and using case studies to illustrate approaches and key success factors Written and edited by experienced practitioners from both industry and academia this publication will equip researchers with a thought provoking guide to the application and future development of contemporary phenotypic drug discovery for clinical Principles and Practice of Clinical Research John I. Gallin, Frederick P Ognibene, 2012-07-10 The third success edition of this innovative work again provides a unique perspective on the clinical discovery process by providing input from experts within the NIH on the principles and practice of clinical research Molecular medicine genomics and proteomics have opened vast opportunities for translation of basic science observations to the bedside through clinical research As an introductory reference it gives clinical investigators in all fields an awareness of the tools required to ensure research protocols are well designed and comply with the rigorous regulatory requirements necessary to maximize the safety of research subjects Complete with sections on the history of clinical research and ethics copious figures and charts and sample documents it serves as an excellent companion text for any course on clinical research and as a must have reference for seasoned researchers Incorporates new chapters on Managing Conflicts of Interest in Human Subjects Research Clinical Research from the Patient's Perspective The Clinical Researcher and the Media Data Management in Clinical Research Evaluation of a Protocol Budget Clinical Research from the Industry Perspective and Genetics in Clinical Research Addresses the vast opportunities for translation of basic science observations to the bedside through clinical research Delves into data management and addresses how to collect data and use it for discovery Contains valuable up to date information on how to obtain funding from the federal government Portfolio, Program, and Project Management in the Pharmaceutical and Biotechnology Industries Pete Harpum, 2011-09-20 This book describes the way that pharmaceutical projects and programs are currently managed and offers views from many highly experienced practitioners from within the industry on future directions for drug program management The book integrates portfolio program and project management processes as fundamental for effective and efficient drug product development Contributing expert authors provide their view of how the projectization approach can be taken forward by the drug industry over the coming years Drug and Biological Development Ronald P. Evens, 2007-08-14 This book offers a complete discussion of product development in the pharmaceutical and biotechnology industries from discovery to product launch through life cycle management The book is organized for optimal usefulness in the education and training of health care professionals MD PharmD PhD at universities

The format is a set of figures tables and lists along with detailed narrative descriptions including real life examples illustrations controversies in industry and references The editors and authors of the book are industry and research experts Curious 2018 Ulrich A.K. Betz, 2019-08-13 This book expands the debate on the future of in a variety of disciplines science and technology at the Curious 2018 Future Insight Conference held on the occasion of Merck s 350 year anniversary In the respective chapters some of the world's top scientists managers and entrepreneurs explore breakthrough technologies and how they can be applied to make a better world for humanity Divided into three parts the book begins with an introduction to the vision of the conference and to the importance of curiosity for innovation while also exploring the latest scientific developments that are shaping the future of healthcare medicine the life and material sciences digitalization and new ways of working together In the second part particular attention is paid to new therapies and diagnostics here readers will learn how synthetic biology and chemistry are being used to solve problems that are essential to the future of humanity The role of in silico research is also discussed In the final part of the book readers will find some thoughts on ethical principles guiding our application of science and technology to create a bright future for humanity Given its interdisciplinary appeal the book will inspire curiosity in a wide readership from scholars and researchers to professionals with an interest in exploring the future of science and technology solving the problems of today and paving the way for a better tomorrow Chapters 1 2 3 and 17 are available open access under a Creative Commons Attribution NonCommercial 4 0 International License via link springer com Translational Surgery Adam E.M. Eltorai, 2023-03-22 Translational Surgery covers the principles of evidence based medicine and applies these principles to the design of translational investigations. The reader will come to fully understand important concepts including case control studies prospective cohort studies randomized trials and reliability studies Investigators will benefit from greater confidence in their ability to initiate and execute their own investigations avoid common pitfalls in surgical research and know what is needed for collaboration Further this title is an indispensable tool in grant writing and funding efforts The practical straightforward approach helps the translational research navigate challenging considerations in study design and implementation The book provides valuable discussions of the critical appraisal of published studies in surgery allowing the reader to learn how to evaluate the quality of such studies Thus they will improve at measuring outcomes making effective use of all types of evidence in patient care In short this practical guidebook will be of interest to every surgeon or surgical researcher who has ever had a good clinical idea but not the knowledge of how to test it Focuses on translational research in Surgery covering the principles of evidence based medicine and applying those principles to the design of translational investigations Provides a practical straightforward approach to help surgeons and researchers navigate challenging aspects of study design and implementation Details valuable discussions on the critical appraisal of published studies in Surgery allowing the reader to effectively use all types of evidence for patient care BNA's Health Care Policy Report ,2002 Miracle Cure Sally Pipes, Fraser Institute

(Vancouver, B.C.), Pacific Research Institute for Public Policy, 2004 America has wealth innovation and access to the best of everything So why is our health care system so broken Why does it cost more than ever and deliver less How do we solve the problems of the uninsured and seniors who lack drug coverage And equally important why is the Canadian system widely touted as a sparkling example of compassion and universal access actually a disastrous model to be avoided of Drug Safety Institute of Medicine, Board on Population Health and Public Health Practice, Committee on the Assessment of the US Drug Safety System, 2007-02-27 In the wake of publicity and congressional attention to drug safety issues the Food and Drug Administration FDA requested the Institute of Medicine assess the drug safety system The committee reported that a lack of clear regulatory authority chronic underfunding organizational problems and a scarcity of post approval data about drugs risks and benefits have hampered the FDAs ability to evaluate and address the safety of prescription drugs after they have reached the market Noting that resources and therefore efforts to monitor medications risk benefit profiles taper off after approval The Future of Drug Safety offers a broad set of recommendations to ensure that consideration of safety extends from before product approval through the entire time the product is marketed and used Green and Sustainable Pharmacy Klaus Kümmerer, Maximilian Hempel, 2010-06-16 Within recent years pharmaceuticals have come into focus as contaminants of the environment see for example K mmerer K editor Pharmaceuticals in the Environment At the same time the issue of sustainable chemistry gained momentum Bringing both together would result in sustainable pharmacy Sustainable pharmacy is a totally new issue and approach It addresses environmental economical and social aspects of pharmacy In the present stage the focus will be on environmental issues along the whole lifecycle of a pharmaceutical entity That is dealing with resources and energy input but also with waste issues for example during the synthesis and production of an active pharmaceutical ingredient Furthermore it would also look on the compounds themselves and will aim to improve the degradability of the compounds after their use in the environment to reduce the environmental risk caused by pharmaceuticals in the environment Another issue is the people using pharmaceuticals such as pharmacists medical doctors and patients How can they contribute to more efficient use of pharmaceuticals with less environmental burden and less risk for drinking water The book Sustainable Pharmacy will address all these issues and will be the first one dealing with this important topic Global Clinical Trials for Alzheimer's Disease Debbie N. Cote, 2013-08-28 About 2% of the United States population gets involved with clinical research trials each year among those people who suffer from severe chronic illnesses including Alzheimer's disease only 6% participate As a result an increasing number of clinical trials are delayed because too few people ever know they had an opportunity to get involved Recruitment strategies consist of effective campaigns that reach the intended population of potential patients A strategy using several tactics will most likely achieve the best outcome The art of patient recruitment is to cast a wide net combining a patient centric enrollment focus involving the use of social media and conventional media coupled with customer friendly protocols Investigative sites providing a high

level of customer satisfaction may result in accelerating enrolment improving retention and thereby reducing the overall The Politics of Precaution David Vogel, 2012-04-29 The Politics of Precaution examines the politics of study cost consumer and environmental risk regulation in the United States and Europe over the last five decades explaining why America and Europe have often regulated a wide range of similar risks differently It finds that between 1960 and 1990 American health safety and environmental regulations were more stringent risk averse comprehensive and innovative than those adopted in Europe But since around 1990 the book shows global regulatory leadership has shifted to Europe What explains this striking reversal David Vogel takes an in depth comparative look at European and American policies toward a range of consumer and environmental risks including vehicle air pollution ozone depletion climate change beef and milk hormones genetically modified agriculture antibiotics in animal feed pesticides cosmetic safety and hazardous substances in electronic products He traces how concerns over such risks and pressure on political leaders to do something about them have risen among the European public but declined among Americans Vogel explores how policymakers in Europe have grown supportive of more stringent regulations while those in the United States have become sharply polarized along partisan lines And as European policymakers have grown more willing to regulate risks on precautionary grounds increasingly skeptical American policymakers have called for higher levels of scientific certainty before imposing additional regulatory controls on business Proteomics in Drug Research Michael Hamacher, Katrin Marcus, Kai Stühler, Andre van Hall, Bettina Warscheid, Helmut E. Meyer, 2006-03-31 From skillful handling of the wide range of technologies to successful applications in drug discovery this handbook has all the information professional proteomics users need Edited by experts working at one of the hot spots in European proteomic research the numerous contributions by experts from the pharmaceutical industry and public proteomics consortia to provide the necessary perspective on current trends and developments in this exciting field Following an introductory chapter the book moves on to proteomic technologies such as protein biochips protein protein interactions and proteome analysis in situ The section on applications includes bioinformatics Alzheimer's disease neuroproteomics plasma and T cell proteomics differential phosphoproteome analysis and biomarkers as well as pharmacogenomics Invaluable reading for medicinal and pharmaceutical chemists gene technologists molecular biologists and those working in the pharmaceutical industry The Agile Approach to Adaptive Research Michael J. Rosenberg, 2010-02-18 Apply adaptive research to improve results in drug development The pharmaceutical industry today faces a deepening crisis inefficiency in its core business the development of new drugs The Agile Approach to Adaptive Research offers a solution It outlines how adaptive research using already available tools and techniques can enable the industry to streamline clinical trials and reach decision points faster and more efficiently With a wealth of real world cases and examples author Michael Rosenberg gives readers a practical overview of drug development the problems inherent in current practices and the advantages of adaptive research technology and methods He explains the concepts principles

and specific techniques of adaptive research and demonstrates why it is an essential evolutionary step toward improving drug research and development Chapters explore such subjects as The adaptive concept Design and operational adaptations Sample size reestimation Agile clinical development Safety and dose finding Statistics in adaptive research including frequentist and Bayesian approaches Data management technologies The future of clinical development By combining centuries old intellectual foundations recent technological advances and modern management techniques adaptive research preserves the integrity and validity of clinical research but dramatically improves efficiency

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