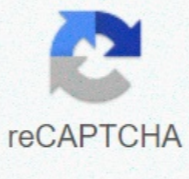




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Introduction to the principles and practices of the clinical research course (IPPCR) by the National Institutes of Health Clinical CentersteTettemberber 12, 2016 - 14 April 2017 Online introduction of clinical research principles and practice (IPPCR) is a course to train participants on how to effectively conduct clinical research. This course will be of interest to doctors, scientists, medical students, nurses, public health professionals and all other health professionals who plan a career in clinical research. To a reminder for all CTSI trainees, this course is one of the many possible electives to CTSI Training Program in Translational Science Track 2 Student can select. It is also one of the requirements that a student of UCLA routes needs to complete at some point by the end of their 4th year. Course information: the course is conducted entirely online See the program of the course IPPCR, Clická here is not the fee to register for the Coursthere is no requirement for attendance for the coursethere is not the academic credit or continuing medical training (CME), the credit offered for the Certificate of Electronic Completion Coursean will be awarded to the successful completion of the course, which is based on the reception of the degree pass of 75% or higher on an open book67. The textbook can be purchased from several online book retailers to register for the course, it is necessary to be an NIH employee or an affiliate participant with one of the approved remote sites of the course is not affiliated with an approved remote site, you can apply To start your remote site as long as your site meets all the requirements of the remote site for more information on the course and registration, visit the course website . The deadline for registration is due to be mid-December. For questions, please email the course coordinator ATIIIPPCR2@mail.nih.gov Skip to main content published by Angela Zito Friday 20 August 2021 in Ads. Provide an overview of the basic biostatistic and epidemiological methods involved in conducting clinical research. Describe the principles involved in ethical, legal and regulatory issues in the search for clinical human subjects, including the role of institutional review cards (IRBS). Describe the principles and issues involved in monitoring patient-oriented research. Describe the required infrastructure in the execution of clinical research and the stages involved in the development and funding of research studies . This course will be of interest to doctors, scientists, medical students and dental students, nurses, public health professionals and others who lead or plan a career in clinical research. Important dates Registration 1 September 2021 á € *1 July 2022 (4pm US EST) Course 1 September 2021 á € *1 August 2022 Final exam 1 September 2021 á € *28 July 2022 (12pm UST) Opened on 1 September here: Tags: Career Exploration Opportunities, Career Options: Clinical Research, Skill Development The introduction to the principles and practice of clinical research (IPPCR) Course Trains Recorders on how to perform clinical research effectively and safely. The course focuses on the spectrum of clinical research and research process by highlighting biostatistic methods andstudy design, protocol preparation, patient monitoring, quality assurance, ethical and legal issues, and much more. The course's objectives offer an overview of the basic biostatistic and epidemiological methods involved in conducting clinical research. describe the principlesNelle questioni etiche, legali e regolamentari nella ricerca sui soggetti umni cliniche, compreso il ruolo delle schede di revisione istituzionale (IRBS). Descriptionverre i principi e le questioni coinvolte nel monitorare la ricerca orientata al paziente. Descriptionverre l'infrastruttura richiesta nell'esecuzione della ricerca clinica e nelle fasi coinvolte nello sviluppo e nei finanziamenti di studi di ricerca. Audience planned Question corso sarà di interest per i medici, gli scienziati, gli studenti medici and dentistici, gli infermieri, gli infermieri, i professionisti della sanità pubblica and altri che conducono o pianificano una carriera nella ricerca clinica. Direttori del corso "Corse Co-Director Dr. John Gallin serves come istituto nazionale del direttore Associao della salute per la ricerca clinica e il Chief Scientific Officer del Centro clinico NIH. Ha pubblicato più di 365 articoli in rivistehe e ha modito due libri di testo á€ "" infiammazione, principi di base e correlati clinici " (Lippincott, Williams and Wilkins, 1999, ora in 3a edizione) e" Principi e practitioner della ricerca clinica " (Stampa accademica, ora in 4zione edizione). Il Dr. Gallin is a member of the American società per l'indagine clinica, l'Associazione dei medici americani, dell'Accademia Nazionale della Medicina, ed è un Maestro del Collegio Americano dei Medici, Anne Zajicek, MD, Pharm.D., FAAP á€ "Course - Co-registration Dr. Zajicek è un pediatra certo del consiglio di amministrazione e pharmacologo clinico pediatrico che attualy funge da vicedirettore dell'Ufficio della ricerca clinica presso i National Institutes of Health. Laura Lee Johnson, Ph.D. - "Course Co-Director Laura Lee Johnson, Ph.D. E il collegamento per lo sviluppo di farmaci focalizzato del paziente e il direttore della divisione per l'ufficio di Biostatistica presso il Centro U.S. Food and Drug Administration (FDA) per la valutazione della drug e la ricerca (Cdr). Si è specializzata nella progettazione, dell'attuazione e dell'analisi degli studi di ricerca di tutte le dimensioni e dello strumento di misurazione e dello sviluppo endpoint. Prima di lavorare alla FDA ha trascorso oltre un decennio agli Istituti nazionali di salute degli Stati Uniti che lavorano e superavano i programmi di ricerca e ricerca sulla ricerca clinica. Lisa Cordes, Pharm.D., BCACP, BCOP - Corse Co-Direttore Dr. Lisa M. Cordes è uno specialista ed educatore di Oncology Clinical Pharmacy per gli Istituti di salute nazionali. Nella his attuale posizione, fornisce supporto clinico e protocollo alla filiale dei neonati genitainari e al programma di farmacologia clinica del National Cancer Institute, ed è il co-direttore dei principi del corso di farmacologia clinica. > ?The video archives and conference performances provided by national health institutes (NIH) Clinical Center Office of Clinical Research Training and Medical Education. Access to past course materials is available for free. Go to: The large thematic areas (modules) include: Ethical design and statistics, jurisprudence and regulatory considerations Preparation and monitoring Clinical studies Lessons include: design of epidemiological studies Participant Studio secondary data / analysis Meta ethical principles in clinical research protocol Development and testing Data and security questionnaires Monitoring Community research committees 1. 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