

sessions were 1 hour, and 15 minutes. Exams given post class session with a mean score of 22.8/30 for session 1 and a mean score of 23.80/30 for session 2. RESULTS/ANTICIPATED RESULTS: The Community Research Academy (CRA) has thus far held for two sessions. Of the 45 enrollees, 20 completed the entire program. Of those, ten have joined the CAB board; and three have actively participated in the pilot award review process. One of the CRA graduates will be publishing her photo novella assignment in Health Affairs. One person changed careers; and several people have taken all the classes for a second time. Many participants originally came from our faith-based connections. Now word of mouth is expanding the program. DISCUSSION/SIGNIFICANCE: Community engagement ensures that the fresh voices of diverse populations are involved in translational science. Their input ultimately leads to creating novel clinical innovations; such community-driven ingenuity which addresses the deeply felt needs of those communities.

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Meeting the Needs of Transgender People through Community Engagement

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OBJECTIVES/GOALS: Alachua TranQuility was formed in early 2016 to address needs of transgender people in Alachua County, Florida and surrounding areas. Increasing awareness of gender identity issues created a growing population in need of additional health and social services to counter the negative health outcomes traditionally experienced by transgender people. METHODS/STUDY POPULATION: The group met monthly at UF HealthStreet, a CTSA community engagement program, where social and medical referrals and opportunities to participate in research were made available to attendees. Those attending included transgender and gender non-conforming individuals as well as family, friends and other allies. Outreach was conducted through social media, physical flyers, and tabling at community events. TranQuility partnered with local LGBTQ organizations to promote activities. Leadership was provided by a steering committee that changed over time, with feedback from attendees guiding planning. Simultaneous to TranQuility's founding, a Youth Gender Clinic was established at UF Health, clearly demonstrating the need for services aimed at the transgender population. RESULTS/ANTICIPATED RESULTS: Between February 2016 and September 2019, attendance at meetings was logged per HealthStreet protocols. Age of attendees ranged from 10 to 75, and most were non-Hispanic whites. More than 300 individuals attended meetings across time, and many attended multiple meetings, with greater than 15% having attended five or more meetings. Attendance averaged around 30+ each month, with a high of 76. Educational programming was offered, such as information on hormone therapy and name/gender marker change, and a parent support group was formed to meet concurrently with the main group. Social opportunities were very popular, as a safe place for people to present as their authentic selves. The coronavirus pandemic curtailed in-person meetings at HealthStreet in 2020, but the group plans to return to this format in 2022. DISCUSSION/SIGNIFICANCE: TranQuility has become an established organization to which other groups provide referrals. Qualitatively it is clear that many people have been positively impacted. An ongoing discussion for the group, however, has been to improve outreach to transgender people of color, who typically suffer from the most discrimination and anti-trans violence.

Regulatory Science

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Enhancing Research Ethics, Equity and Protections for Uninsured Study Participants

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OBJECTIVES/GOALS: To increase diversity in clinical and translational research (CTR), to strengthen protections for uninsured and under-insured study participants from vulnerable groups by addressing their medical, ancillary care and psycho-social needs and to develop a systematic ethically sound approach to addressing such needs in the study protocol and budget. METHODS/STUDY POPULATION: We conducted ethical analyses of: (1) the regulatory and ethics scholarship concerning protections and duty of care to research participants from vulnerable groups, and 2) arguments concerning the nature and scope of ancillary care obligations of researchers, as well as 3) a review of the applicable local, federal, and international regulations and practices concerning the duty of care to CTR participants and potential participants who are uninsured, under-insured and/or undocumented members of vulnerable groups. RESULTS/ANTICIPATED RESULTS: Uninsured and underinsured study participants pose major ethical challenges for CTR as medical needs arising during a study are usually covered by the participants own insurance. Lack of health insurance increases vulnerability and creates (1) a barrier to research participation for members of socially disenfranchised groups, (2) risk of discriminatory exclusion of such participants from clinical studies, and (3) inter-institutional inconsistencies in meeting their medical needs; thus limiting diversity in CTR. To address the challenges, we propose an inclusive, systematic, ethically sound approach, which deliberately plans for and provides resources within a study protocol to address the medical and psycho-social needs of uninsured participants during and beyond the study. DISCUSSION/SIGNIFICANCE: Including diverse participants in CTR ensures data quality and social justice. PIs and IRBs should adopt an inclusive approach to the medical needs of vulnerable uninsured participants, plan for their medical and ancillary care needs in the protocol and budget, list community resources, provide follow-up support and note assistance in their files.

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Ethical Considerations of Decentralized Clinical Trials

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OBJECTIVES/GOALS: Our goal is three-fold: (1) to enhance protections for research participants in decentralized clinical trials (DCT) by (2) identifying the ethical and regulatory challenges posed by DCT and (3) considering possible solutions to the ethical and regulatory challenges of DCT. METHODS/STUDY POPULATION: A literature review was conducted to identify the ethical and regulatory challenges of DCT. The review showed that, with few exceptions, the publications on DCT have been written by IT experts, researchers or representatives of the pharma industry. There are hardly any independent or multidisciplinary analyses of DCT, e.g., by ethicists, medical sociologists or patients. This suggests that, currently, the

potential advantages of digitized clinical research are still to be tested empirically and confirmed independently. Ethical analyses of DCT are particularly scant, which points to a gap in the current understanding of the ethical and regulatory implications of DCT and the issues they raise. Our research used the method of ethical analysis to examine the ethical challenges and to consider possible solutions. RESULTS/ANTICIPATED RESULTS: We identify the following ethical challenges of DCT: (1) challenges related to modifications of research protocols to allow for decentralized activities; (2) lack of expertise for ethical review and approval of digital tools; (3) different regulatory standards of privacy for clinical trials and for commercial digital tools, (4) risks to participants privacy and confidentiality involving both data and physical privacy; (5) the impact of the digital divide on DCT, (6) technology-related subject selection bias and marginalization of vulnerable groups, (7) imposing new burdens on research subjects and caregivers due to technology needs and smaller research teams, (8) restricted access to the study team, and (9) impact of digital technology on informed consent and participant understanding. DISCUSSION/SIGNIFICANCE: DCT became the primary form of clinical research during the COVID-19 pandemic. DCT rely on a wide range of internet-based tools for recruitment, informed consent, data collection, health monitoring, and communication with participants. The tools pose ethical and regulatory challenges, which should be addressed to ensure participant well-being.

Research Management, Operations, and Administration

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Health Equity Starts with Us: Recommendations from the Indiana Clinical and Translational Sciences Institute Racial Justice and Health Equity Task Force

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OBJECTIVES/GOALS: The Indiana CTSI Strategy Committee charged the Racial Justice and Health Equity Taskforce to identify priorities with short-term and long-term goals consistent with the I-CTSI mission. In addition, I-CTSI leadership asked for a general description of current state and the resources necessary to achieve the proposed goals. METHODS/STUDY POPULATION: The Taskforce applied an inclusive excellence model to the way we look at the I-CTSI structure, policies, and programs while performing an environmental scan within and across I-CTSI partner institutions. In order to reach equitable solutions and consensus, listening tours were held with partner stakeholders guided by the SOAR framework for strategic planning. This approach allowed us to assess current resources, needs, and gaps across the system, along with a baseline of measures currently monitored. Taskforce members openly discussed strengths and opportunities for enhancement of current programs and services. In addition, these conversations offered an opportunity to disrupt existing practices and through collective agency we identified priority areas that promote equity, diversity and inclusion. RESULTS/ANTICIPATED RESULTS: The Taskforce identified recurring themes in conversations with all partners, which led to the formation of three working groups

that examined recruitment broadly: workforce, staffing, and research participation; professional development across all stakeholders from community members to I-CTSI staff; and data-centered metrics informing current state, decision-making, and accountability. Recommendations included these priorities, content, and implementation strategies. The Taskforce delivered a report to the I-CTSI leadership fostering the promotion of diversity, equity and inclusion along with a systematic collection of gender, race, and ethnicity data for individuals utilizing I-CTSI services and resources requiring additional metrics and tracking. DISCUSSION/SIGNIFICANCE: The pandemic shed light on the manner in which marginalized groups are rendered particularly vulnerable to death and disease by systemic and structural racism. The I-CTSI recognized that we cannot advance population health without attending to root causes of inequity and that includes our internal structure. We offer a potential model for other CTSAs.

Team Science

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A CTS Team Approach to Incorporating Black Womens Conceptualization of Trust in the development of a Mindfulness Practice tool tailored for the Reproductive Provider Space

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OBJECTIVES/GOALS: The goals of the proposed project are to 1) identify the critical components of trust required by Black Women to build a visible expression of trust within their reproductive health space, and 2) incorporate the elements of trust identified into a mindfulness-based communication tool, for reproductive health care providers. METHODS/STUDY POPULATION: We will be applying a mixed-methods approach inclusive of questionnaires and focus groups. Aim 1: Our study population consists of persons who identify as 18+ Black Women in Alachua County, Florida. Aim 2: Our Population within the reproductive health space (e.g., ob-gyn, midwives, nurse practitioners, etc...) at the University of Florida Shands Hospital. RESULTS/ANTICIPATED RESULTS: We anticipate that our results will fully engage with our Black Women in dialogues about key components of trust they want to experience and have within their reproductive space. This will inform the development of the mindfulness-based tool which will be incorporated with continuous insights with our Black Women. The proposed research project will contribute to the call for health equity. We aim to address this barrier by recognizing the agency of Black women to conceptualize trust on their terms, which we then would incorporate into applied trainings of reproductive health providers to establish trust. Health equity will take a considerable amount of social change at many levels and this study aims to better understand the process. DISCUSSION/SIGNIFICANCE: Structural racism and health disparities are linked with the disproportionate loss of Black women and associated negative birth outcomes. Our work will provide needed insights regarding mistrust in patient-provider relationships. Inform the development of a feasible tool challenging implicit bias in health provider spaces.