Down Syndrome Research: The Intersection of Basic Science and Clinical Cohort Development November 9t10, 2020 NIH-Sponsored Virtual Meeting

EXECUTIVE SUMMARY

On November 910, 2020, the National Institutes of Health (NIH) in Bethesda, MD, sponsored a virtual workshop of the INCLUDE Project Nvestigation of Co-occurring conditions across the span to Understand Down syndrom (Down Syndrome Research: The Intertiment of Basic Science and Clinical Cohort Development._Representatives from NIH, basic and clinical research set for the span of the set of the s

Keynote Presentations : Perspectives from Research Study Participants

Day 1 of the meeting included keynote presentations **by y**oungpeople with DS and the if a milies who have participated inclinical studies of DS. The esenters offered their personal views on the importance of engaging with participants throughout the course of the clinical trial, making the experience personal and relevant, and sharing the outcomes of the study addition they said their research experience gave them a familiarity with hospitals and health providers, which made going to see the doctor a more positive experience. They asked that investigators to schedule invasive research procedures such as blood **docros** incide with the participants informed about research updates using social media and understandable language and educate and engage potential candidates about clinical trials. They suggested that more information about the transition from adolescence to young adulthood is needed.

Day 1 Reports from Working Groups and Breakout Sessions

Presentations on conorbidities associated with DS follow**the** keynote presentations, with discussions on neurodevelopment, behavior, cardiovascular dise**ase** pulmonary hypertension, and respiratory and airway conditions. Additional presentations were given on canaetoimmunity and infectionsendocrine, metabolic, and skeletabonditions and aging and Alzheimerdisease(AD) The meeting then divided intereakout Groups 1 (Development and Behavior), 2 (Heart and Lung), 3 (Cancer and mmunity), and 4 (Aging and Metabolic Conditions). The breakout groups identifies to ecommon themes, including e need for longitudinal cohort studies with welvalidated endpoints, better animal and cellular models for preclinical data, more cohort diversity, integration of adult and pediatric cohoits a single cohort across the lifespan, collection of samples of convenience from routine surgical procedures, and better harmonization and linkage of databases. On the basic science side breakout groups discussed the need to bring together infation on phenotypes of various mouse models, provide more funding opportunities for model development, and develop induced stem cells to generate lines from propenewith DS It was announced during Dayof the meeting thatwhole genome sequencing data 20,600 people with DS would soon be available to be shared with the community.

Day 2 Session on Basic ScienceModel Systems and Tools to Advance Down Syndrome Basic and Preclinical Science

This session focused the current state of DSmouse models An overarching suewasthe importance of knowing the background of the mouse model and in research studies cause many factors can affect the mouse phenotypesuch as the strain of the mouse and/ow the model was derived Dne promising rodel is the TcMAC2 mouse, which has an tificial chromosome containing the long arm of human chromosome 21, retains 93% of the human chromosome protein coding genes, and is not mode was also begun to put the human chromosome 21 in rats, which the date the human centromere better than miced, unlike mice, are rarely mosaic Research using human induced pluripotent stem cell (SPSCs) is moving forwald vestigators an now use patient derived iPSC to study condition common people with DS such as congenital heart defects, intellectual disability, and D. More researchers are now using threferensional cell cultures that allow cells to self-organize into organoids. This method supports greater nursible cell types and cell interactions than or dimensional cell cultures that allow cells to self-organize into organoids. This method supports greater nursible cell types and cell lines containing the presentilin mutation from patients with familial AD to use in threferensional cultures. This presentation also described studies of theole of extracellular vesicles in AD pathogenesis

Day 2 Session onCohort Development: INCLUDE Data Coordinating Center and Existing and Future Cohorts

NIH has funded multiple projects in support of development of DS cohort studies. One such effortsinvolv creation of a data coordinating centerD(CC) and a data portal to standardize, harmonize, and aggregate DS data into a virtual biorepository with a goal of providing data access and analysis tools for transformative DS research. The findings of a surveof 57 existing cohorts and databases related to DS research as a starting point for the DCC another presentation described variety of options for linking data, including Global

Unique Identifiers (GUIDs), PCORnet, Datavant, and a refedralmodel that is being used in the DS