February 19, 2020

Dr. Marcella Nunez-Smith  
Chair, Biden-Harris COVID-19 Health Equity Task Force  
Office of Minority Health  
Department of Health and Human Services  
Tower Oaks Building, Suite 100  
1101 Wootton Parkway  
Rockville, MD 20852  

Re: Summary Recommendations to Combat COVID-19 Disparities with a Comprehensive Research Agenda on Disability, Independent Living, and Rehabilitation

Dear Dr. Nunez-Smith:

On behalf of the Disability and Rehabilitation Research Coalition (DRRC), we are pleased to share with you and your colleagues on the COVID-19 Health Equity Task Force our recommendations for improving the lives of Americans with disabilities by enhancing the federal commitment to disability, independent living, and rehabilitation research. We understand that addressing the health and social disparities faced by people with disabilities (in many cases preexisting, but significantly exacerbated by the COVID-19 pandemic) is a key facet of the mission of the Task Force, and we look forward to working with you and your team on this critical issue. In particular, this letter addresses:

- Critical COVID-19 Research on Disability, Independent Living, and Rehabilitation  
- Collection of Demographic Data on COVID-19 Including Disability Status Data  
- Support for COVID-Specific Disability and Rehabilitation Research at the Federal Level

The DRRC is a coalition of more than 20 national research, clinical, and consumer non-profit organizations committed to improving the science of rehabilitation, disability, and independent living¹. The DRRC seeks to maximize the return on the federal research investment in these areas with the goal of improving the ability of Americans with disabilities to live and function as independently as possible following an injury, illness, disability, or chronic condition. The coalition plays a leadership role in coordinating the activities of stakeholders to increase and leverage federal resources devoted to research and development in these domains.

¹ Please see Attachment A for a complete list of DRRC members.
The DRRC believes that disability and rehabilitation research plays a critical role in enabling and empowering individuals with disabilities to live the American dream, consistent with the notion that disability, like race and gender, is a natural part of the human experience and the goals of federal disability policy articulated in the Americans with Disabilities Act (ADA) – equality of opportunity, full participation, independent living, and economic self-sufficiency. This research is also crucial for identifying and eliminating the inequities faced by people with disabilities in many areas, particularly with regards to the impact of COVID-19.

We appreciate the focus that the Biden-Harris Administration has placed thus far on disability issues and look forward to working with the Task Force to advance federal public policy impacting people with disabilities. Significant disability and rehabilitation research, including translational research, is necessary to develop new and more effective approaches to test and evaluate the costs and benefits of current services, supports, treatments, and devices and to use these findings to affect future programs and maximize benefits to costs. Therefore, the DRRC recommends that the Task Force work with others in the Administration to support the development and implementation of a comprehensive disability and rehabilitation research agenda, including the priorities enumerated below. We also provide as attachments to this letter more detailed documents providing further information on these recommendations.

I. Recognize and Conduct Critical COVID-19 Research on Disability, Independent Living, and Rehabilitation Research

As the Biden-Harris Administration implements its effort to combat the COVID-19 pandemic and prepare for future global health crises, we urge the Task Force to help ensure that the federal research agenda encompass the pandemic’s impact on people with disabilities and chronic conditions. Including this population in COVID-19 research will boost the pandemic response, more efficiently direct federal resources, and inform risk communication development. In addition to the specific interactions of this virus with pre-existing disabilities and the disparities in complications and outcomes faced by disabilities, it is also crucial to understand the ripple effects of the pandemic on disability populations, including increased barriers to accessing health care and social services, loss of employment, social isolation, and other disparate impacts on people with disabilities.

We have included a recent letter (Attachment B-1) from the DRRC to NIH Director Francis Collins outlining the importance of a dedicated disability and rehabilitation-focused federal research effort into COVID-19 and proposing an agenda of key themes and topics for investigation.

II. Ensure Collection of Key Demographic Data on COVID-19 to Identify and Address Disparities

Among all efforts to collect data regarding COVID-19 testing, diagnoses, treatment, and outcomes, we urge the Task Force to help ensure that data is disaggregated by disability status, as well as other key demographic factors. Without collecting data disaggregated by disability status associated with COVID-19, people with disabilities will remain at greater risk of disease and death. This is an essential step to supporting the federal pandemic response and addressing the disparate impact of COVID-19 on people with disabilities and other minority populations.
We appreciate the steps the Biden Administration has taken thus far to improve data collection, including establishing the Interagency Working Group on Equitable Data, and hope these efforts will continue to be expanded to ensure that COVID-19 data collection address populations facing health inequities.

*We have included a separate letter (Attachment C-1) from the DRRC regarding the collection of disability status data in all COVID-19 related research and how this data can be used to help address the disparities exacerbated by the pandemic.*

### III. Increase COVID-Specific Funding for Disability and Rehabilitation Research

DRRC believes that disability and rehabilitation research programs have been chronically underfunded for many years, especially considering the magnitude of the aging population and the current and future veterans’ and civilians’ need for rehabilitation, independent living, and disability-related services, supports, treatments, and devices. This research has become even more critical to address the additional needs associated with understanding the impact of the COVID-19 pandemic on people with disabilities. DRRC urges the Task Force to support in its recommendations to the President significant increases in funding for the various federal agencies supporting and conducting this research, especially the National Institute for Disability, Independent Living, and Rehabilitation Research (NIDILRR), housed within the Administration for Community Living.

Additionally, we note that the important work of coordinating government-wide research to address the needs of people with disabilities during the COVID-19 pandemic should closely involve the Interagency Committee on Disability Research (ICDR). We also note that the ICDR in recent years has not received sufficient commitments of resources and high-level staff to appropriately carry out its mission. We believe that ICDR has a critical role to play in advancing disability research that addresses the inequities caused and deepened by COVID-19 and we encourage the Task Force to recommend that the Biden Administration reinvigorate ICDR and provide the necessary infrastructure support to improve its functioning.

*We have included a letter from the DRRC (Attachment D) and a 2020 white paper from the National Association of Rehabilitation Research and Training Centers (NARRTC) [Attachment E-1] detailing the specific need for additional funding at NIDILRR to conduct applied research relating to the COVID-19 pandemic.*

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We greatly appreciate your consideration of our recommendations to enhance the federal research agenda regarding disability, independent living, and rehabilitation. The DRRC stands ready to serve as a resource for the Task Force to answer any questions that may arise and provide technical expertise on behalf of the field. If we can be of any assistance, please do not hesitate to contact the DRRC coordinators, Peter Thomas, Joe Nahra, and Bobby Silverstein, at 202-466-6550 or by email at Peter.Thomas@PowersLaw.com, Joseph.Nahra@PowersLaw.com, and Bobby.Silverstein@PowersLaw.com.
Sincerely,

The Disability and Rehabilitation Research Coalition (DRRC)

Bobby Silverstein, DRRC Coordinator
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Joe Nahra, DRRC Coordinator
Joseph.Nahra@PowersLaw.com

CC: Andrew Imparato, JD, Member, COVID-19 Health Equity Task Force
Alison Barkoff, Acting Administrator and Assistant Secretary for Aging, Administration for Community Living
The DRRC is a coalition of 26 national non-profit organizations committed to improving the science of disability, independent living, and rehabilitation. The DRRC seeks to maximize the return on the federal investment in disability, independent living, and rehabilitation research and development with the goal of improving the ability of Americans with disabilities to live and function as independently as possible. The coalition plays a leadership role in increasing and leveraging federal resources devoted to disability, independent living, and rehabilitation research.

**DRRC Members**

- Academy of Spinal Cord Injury Professionals
- American Academy of Orthotists & Prosthetists
- **American Academy of Physical Medicine & Rehabilitation***
- American Association on Health and Disability
- **American Congress of Rehabilitation Medicine***
- American Medical Rehabilitation Providers Association
- American Music Therapy Association
- **American Occupational Therapy Association***
- **American Physical Therapy Association***
- American Speech-Language-Hearing Association
- American Therapeutic Recreation Association
- Amputee Coalition of America
- **Association of Academic Physiatrists***
- Association of Rehabilitation Nurses
- Association of University Centers on Disabilities
- **Brain Injury Association of America***
- Child Neurology Society
- Christopher and Dana Reeve Foundation
- National Association for the Advancement of Orthotics & Prosthetics
- **National Association of Rehabilitation Research Training Centers***
- National Association of State Head Injury Administrators
- National Multiple Sclerosis Society
- National Neurotrauma Society
- Paralyzed Veterans of America
- Rehabilitation Engineering and Assistive Technology Society of North America
- United Spinal Association

***DRRC Steering Committee Member***

For more information:

Contact the DRRC coordinators at 202-466-6550 or via email at:

Peter Thomas: Peter.Thomas@powerslaw.com
Bobby Silverstein: Bobby.Silverstein@powerslaw.com
Joe Nahra: Joseph.Nahra@powerslaw.com
September 8, 2020

Francis S. Collins, M.D., Ph.D.
Director
National Institutes of Health
1 Center Drive
Bethesda, MD 20814

Re: Support for Federal Investment into COVID-19 Research on Disability, Independent Living, and Rehabilitation

Dear Director Collins:

On behalf of the Disability and Rehabilitation Research Coalition (DRRC), we write to express our strong support for the critical research being conducted on the COVID-19 pandemic and urge that this research encompass the pandemic’s impact on people with disabilities and chronic conditions. Including this population in COVID-19 research will boost the pandemic response, more efficiently direct federal resources, and inform risk communication development. In addition, it is crucial that federal research develop a broader understanding of the specific interactions of this virus with pre-existing disabilities and the disparities in complications and outcomes faced by people with disabilities, who already tend to have poorer health and health outcomes due to entrenched disparities in the health care system.

It is already clear that people with disabilities have greater vulnerability to the COVID-19 virus, including a higher risk of exposure and infection, especially for those living in congregate settings. Additionally, people with disabilities face barriers to accessing care that have only been exacerbated during the pandemic, making disability-focused research a critical priority for the federal government.

We have included below an appendix outlining research topics, developed by members of the DRRC conducting clinical research across the country, for inclusion in the federal research agenda on COVID-19. These topics cover a wide spectrum and are organized into four thematic categories:

- **Ensuring Inclusion of People with Disabilities in Existing COVID-19 Research** (e.g., collecting and reporting disability status information in all ongoing collections; tracking sequelae of COVID-19 in the context of disability; and studying acute, subacute, and chronic effects of virus on recovering patients)
- **Disability-Specific COVID-19 Research** (e.g., sequelae of COVID-19 that may result in disability; disparities in COVID-19 outcomes and potential heightened risks of infection)
among people with disabilities; and development of a national database or registry to track outcomes for COVID-positive patients)

- **Rehabilitation Interventions for COVID-19 Treatment and Recovery** (e.g., examination and comparative effectiveness analysis of alternative treatments for COVID-19 patients with complex needs, including ventilation and respiratory support; opportunities to expand and increase access to tele-rehabilitation for COVID-19 treatment; and potential treatments to regain function during the course of recovery)

- **Immediate Impacts of COVID-19 Pandemic on People with Disabilities** (e.g., disparities and patterns in furlough, loss of employment, and return to employment for people with disabilities during the pandemic; impacts of social isolation and quarantine policies on older adults and people with disabilities; and barriers in access to services caused by the pandemic, including prevention, wellness, and fitness programs)

As the Department of Health and Human Services (HHS) and agencies within the Department, including the National Institutes of Health (NIH), the Centers for Disease Control and Prevention (CDC), and the National Institute on Disability, Independent Living, and Rehabilitation Research (NIDILRR) draft, revise, and update their respective strategic plans and budget requests in response to the pandemic, it is crucial that there be a recognized and explicit inclusion of research specifically relating to people with disabilities.

We urge the leadership of these agencies to consult with career disability researchers, medical experts, advocacy organizations, and other stakeholders with the necessary expertise and experience within the disability community in order to develop research plans that will appropriately address the needs of these populations. As trusted thought leaders and health care service providers, these individuals and organizations should be used as a resource to develop research toward creating holistic solutions to remedy the inequitable impact of this and future public health crises.

We appreciate your consideration of our recommendations to integrate rehabilitation and disability considerations into the federal research structure for COVID-19. The DRRC stands ready to serve as a resource to answer any questions that may arise and provide technical expertise on behalf of the field. If we can be of any assistance, please do not hesitate to contact the DRRC coordinators, Peter Thomas, Joe Nahra, and Bobby Silverstein, at 202-466-6550 or by email at Peter.Thomas@PowersLaw.com, Joseph.Nahra@PowersLaw.com, and Bobby.Silverstein@PowersLaw.com.

Sincerely,

The Undersigned Members of the Disability and Rehabilitation Research Coalition

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National Multiple Sclerosis Society
Paralyzed Veterans of America
Rehabilitation Engineering and Assistive Technology Society of North America
United Spinal Association

CC:
Dr. Alison Cernich, Deputy Director, National Institute of Child Health and Human Development
Dr. Theresa Cruz, Director, National Center for Medical Rehabilitation Research
Lisa Kaeser, Chief, Office of Legislation and Public Policy, National Institute of Child Health and Human Development
COVID-19 Research Topics on Disability, Rehabilitation, and Independent Living

1. Ensuring Inclusion of People with Disabilities in Existing COVID-19 Research

As the federal government has turned its focus to pandemic response, research agencies have embarked on significant and widespread efforts to develop and advance fundamental knowledge of the novel coronavirus, including the viral biology, processes of infection and transmission, and the impact of the disease on COVID-positive patients. These efforts are critical to informing public health actions to combat the virus. However, it is essential that any COVID-19 research include people with disabilities, especially regarding broad data collection and related analyses. Without collecting data associated with COVID-19 disaggregated by disability status and other demographic factors, people with disabilities will remain at greater risk of disease and death. Inclusive research on COVID-19 will boost the pandemic response, more efficiently direct critical resources, and inform risk communication development. Existing research projects should ensure that individuals with disabilities are included in the studied populations and that data collection appropriately identifies and tracks these individuals, in order to properly reflect the wider population that continues to be impacted by COVID-19. Potential research topics under this theme may include:

- Sequelae of COVID-19 in the context of disability, including impact of virus on mobility, cognition, and functional performance.
- Acute, subacute, and chronic effects of virus on recovering patients, including cognitive changes, fatigue, endurance, activity tolerance, quality of life, and psychosocial/mental health functioning.
- Inclusion of people with disabilities and collection of disability status information in all ongoing research on COVID-19.
- Disability status changes by incidence and disparity, including cross-references with demographic and geographic factors.

2. Disability-Specific COVID-19 Research

As researchers and clinicians are beginning to better understand the impact of the COVID-19 virus, it is clear that infection can result in serious, extensive, and potentially long-lasting damage beyond the commonly understood respiratory symptoms. It seems likely that there will be a population of previously nondisabled individuals who will develop short-term or even permanent disabilities as a result of COVID-19. It is also clear that people with pre-existing chronic health conditions and disabilities will be at higher risk for serious complications from the disease. It is essential that the federal government work to understand the specific interactions of this virus with pre-existing disabilities and the disparities in complications and outcomes faced by people with disabilities, who already tend to have poorer health and health outcomes due to entrenched disparities in the health care system. Additionally, research must be conducted to better understand the long-term impacts of the disease and the potentially permanent complications related to COVID-19 infection. Potential research topics under this theme may include:

- Sequelae of COVID-19 that result in disability, among people with and without pre-existing disabling conditions.
Disparities in COVID-19 outcomes and potential heightened risks of viral infection for people with pre-existing disabilities.

COVID-related brain and other body organ damage.
  - Extent/permanence of observed neurologic symptoms.
  - Impact of prolonged ventilator treatment.
  - Prevalence/association of additional comorbidities due to COVID (e.g., stroke, deep vein thromboses, myocardial infarctions, etc.)
  - Impact of prolonged isolation during treatment
  - Long-term impact(s) on physical and cognitive function/outcomes.

Development of a national database or registry to track outcomes (including mortality, function, community integration, daily living, and employment) for people diagnosed with COVID-19, similar to the TBI, SCI, and Burn Model Systems.

Enhancement of existing Model Systems databases with inclusion of information on the impact of COVID-19 among patients in the databases.

Systematic evaluation of consumer experiences regarding COVID-19 testing, treatment, and rehabilitation/habilitation among persons with disabilities to inform guidance for improved access and accessibility to health services.

3. Rehabilitation Interventions for COVID-19 Treatment and Recovery

The breadth of the potential complications associated with COVID infections necessitate a broad treatment portfolio to aid in recovery from the virus. In order to assist with treatment of some specific conditions of the virus (e.g., respiratory issues resulting from ventilator support, cognitive “fog,” and general motor function), many patients will need some form of short- or long-term rehabilitation in order to regain lost function. As rehabilitation interventions are used in a variety of settings to enhance COVID-19 recovery, it is important that research into the effectiveness of such treatments be conducted, including best practices, outcomes, and COVID-specific training for rehabilitation providers. Additionally, patients who were in need of rehabilitation prior to contracting COVID-19 will likely still need these services, depending on the severity of their case – the practice of rehabilitation will necessarily shift for these patients and research should examine the impacts of these changes. Potential research topics under this theme may include:

- Potential alternative treatments for COVID-19 patients in need of ventilation, e.g., forced inspiratory training.
- Development of standard outcome and quality measures for individuals with COVID-19 receiving rehabilitation services.
- Evaluation of respiratory support needs and outcomes among persons with disabilities who receive complementary interventions (i.e., music-based and other interventions with therapy support vs. without)
- Best practices for the provision of rehabilitation services such as physical and occupational therapy during acute and post-acute disease management to restore or maintain mobility and function to achieve the highest degree of independence, and methods to measure the results and effectiveness of such treatments.
- Impact of rehabilitation therapy (including physical, occupational, and speech-language pathology) on improving function post-recovery for COVID-19 patients and associated training for family and caregivers.
• Feasibility of validated instruments for assessing persons with disabilities experiencing prolonged disorders of consciousness (PODC) following COVID-19 ventilator support.

• Opportunities to expand and increase equitable access to and permanent coverage of tele-rehabilitation for COVID-19 treatment and recovery.

• Roles and efficacy of rehabilitation therapy provided via telehealth (including physical, occupational, and speech language pathology) for improving functional capacity and maximizing independent living skills during and following COVID-19 infection.

• Development of best practices for training and implementation of COVID-19 rehabilitation programs.

• Potential treatment(s) and course of recovery of physical, cognitive, and psychosocial functioning post-COVID.

• Use of crisis standards of care to understand the scope of potential biases experienced by people with disabilities when receiving care during a health crisis and inform ethical guidance for service providers and caregivers.

• Recognition by insurers and third-party administrators of the appropriateness of and payment for post-COVID rehabilitation services.

• Importance of reduced cost-sharing and administrative burden in eliminating barriers to care.

4. Immediate Impacts of COVID-19 Pandemic on People with Disabilities

The pandemic has not only impacted the country’s health care system but has upended nearly every facet of the nation’s infrastructure. The pandemic’s quarantining and social distancing requirements, along with the associated economic recession, have already had dire and disproportionate consequences for people with disabilities and other vulnerable populations. People with disabilities already face significant systemic inequities in income, education, employment, housing, transportation, and community participation. These conditions have been exacerbated in many cases by the indirect effects of COVID-19, necessitating an informed, coordinated, and sustained federal response. Applied research into the impact of the pandemic, including on employment, education, and access to services for people with disabilities, is critical to supporting this population during the pandemic and the prolonged aftereffects. Of course, the non-COVID-related health concerns of people with disabilities have not diminished, and it is similarly important for federal research to explore how health care and rehabilitation for these individuals has been impacted by the system’s pivot to a COVID-19 focus. Potential research topics under this theme may include:

Employment

• Comparative likelihood of furlough/loss of employment for workers with disabilities.

• Disparities in return-to-work patterns and employment for people with disabilities.

• Use of SSI/SSDI vs. unemployment insurance for workers with disabilities furloughed/terminated due to the pandemic.

• Impact of the pandemic on the transition from school to work for youth with disabilities.

• Impact of the pandemic on access to and effectiveness of supportive services toward the school-to-work transition and career development in young adulthood for people with disabilities.

• Disparities in return-to-work patterns, employment, access to and effectiveness of supportive services for people with disabilities during and post-COVID.
Quarantine/Social Distancing

- Impact of and disparities in social isolation and occupational deprivation on older adults and people with disabilities.
  - Equitable and accessible interventions to reduce social isolation during times of quarantine.
- Metric development for measuring community participation and re-engagement following COVID-19.
- Impact of quarantining/social distancing on individuals with serious mental illness.
  - Shutdown of public transportation systems.
  - Heightened structural risk for individuals in group, residential, or inpatient settings.
  - Loss of income for individuals on SSI/SSDI.
- Psychosocial impact of social distancing on people with disabilities, especially youth with disabilities, and related impact on other areas of function.
- Effectiveness of community-based interventions and innovations in creative arts therapies to address social isolation for people with disabilities, especially those living in congregate care settings.
- Effectiveness of different media and tele-rehabilitation services for teaching students with disabilities during periods of isolation/distancing.
- Impact of social distancing and COVID-19 response on treatment/rehabilitation and associated outcomes for people with pre-existing disabilities, chronic conditions, and injuries.
- Opportunities for existing providers of services for people with disabilities to implement telehealth and mobile health services.
- Opportunities to support people with disabilities, students, and caregivers directly through the use of telehealth and mobile health technology and services.
- Efficacy of in-person, telehealth, and virtual therapy interventions (including physical, occupational, and recreational therapy) focused on mindfulness, physical activity, exercise, and stress management during quarantine, including for people with physical, cognitive, and sensory impairments.
- Effectiveness of the use of prescribed video games, mindfulness/medication apps (like Headspace), and at-home biofeedback tools to reduce depression, anxiety, and mood issues in adults and behavior and academic difficulties of students with IEPs/behavioral intervention plans during the pandemic.

Access to Essential Services

- Impact on availability and quality of care provided by health care providers including peer specialists/peer providers due to social distancing requirements and limits on in-person visits.
- Impact of COVID-19 on access to non-COVID health care services for people with disabilities.
- Impact of COVID-19 recession on budgets for state and local services provided to people with disabilities.
- Impact of stay-at-home orders and safety of public transportation to accessing COVID-19 testing facilities for people with disabilities.
- Availability of home- and community-based services and supports for adults with disabilities due to the pandemic, especially a ready-trained and reliable workforce of personal care assistants and an affordable supply of essential personal protective equipment (PPE) and products.
• Availability of assistive devices and accessible emergency response technologies. Maintaining an adequate supply chain of assistive devices that can be delivered and/or repaired in a timely manner.
• Opportunity for community health workers to work as social “first responders” for people with disabilities and concomitant health disparities during COVID-related workforce shortages and how to receive payment for these services.
• Opportunities to develop/improve telehealth interventions to address needs of people with disabilities.
• Effectiveness of expanded telehealth services and related treatment guidelines for people with disabilities.
• Access to critical supplies for people with limited mobility, low vision, blindness, and other sensory and/or cognitive disabilities during pandemics.
• Potential protective factors against COVID-19 and related complications of social safety net programs (including SNAP, SSI, and housing supports) for people with disabilities.
• Impact of increased food insecurity on people with disabilities due to quarantines, limited availability of food delivery services, limited food products, especially for those already living in food deserts and/or increased demand on federal, state, and local nutrition assistance programs and reductions in income/resources.
• Impact of increased housing insecurity on people with disabilities.
• Interventions to boost the impact of housing/food assistance programs for people with disabilities during and post-COVID.
• Investigation of disparities faced by people with long-term disabilities pre- and post-COVID.
  o Community living, community participation, work/economic equity.
• Disparities in access to rehabilitation services based on social determinants of health and impact of lack of access to treatment pre- and post-COVID, including economic stability, education, health and health care, neighborhood and built environment, and social and community context.
February 19, 2020

Dr. Marcella Nunez-Smith
Chair, Biden-Harris COVID-19 Health Equity Task Force
Office of Minority Health
Department of Health and Human Services
Tower Oaks Building, Suite 100
1101 Wootton Parkway
Rockville, MD 20852

Re: Collection of Disability Status Data in Mandated Reporting Regarding COVID-19 Testing, Diagnoses, Treatment, and Outcomes

Dear Dr. Nunez-Smith:

On behalf of the Disability and Rehabilitation Research Coalition (DRRC) and the undersigned disability and rehabilitation stakeholder organizations, we urge you and the Health Equity Task Force to recommend that the Centers for Disease Control and Prevention (CDC) and other program operating components (POCs) within the Department of Health and Human Services (HHS) collect and report data on demographic characteristics, de-identified and disaggregated by disability status, to better inform the COVID-19 pandemic response. This would also help achieve the goals of President Biden’s Executive Order 13985, Advancing Racial Equity and Support for Underserved Communities Through the Federal Government. As you know, that order establishes an Interagency Working Group on Equitable Data, and we believe that ensuring and standardizing the collection of disability status data regarding the COVID-19 pandemic would be an important first step towards fulfilling the mission of the Data Working Group. The data should, among other things, address:

- Testing and diagnoses;
- Treatment (including admissions and denials to hospitals and intensive care units, denials of needed accommodations, virus interactions with chronic disease and disability, and recovery patterns); and
- Outcomes (including morbidities, mortalities, rehabilitation, and recovery).

Without collecting data disaggregated by disability status associated with COVID-19, people with disabilities will remain at greater risk of disease and death. Health data collected regarding disability status will boost the pandemic response, more efficiently direct critical resources, and inform risk communication development.1 In the appendix below, we outline some of the ways

1 As you know, the recently passed Paycheck Protection Program and Health Care Enhancement Act (P.L. 116-139) includes provisions requiring the regular reporting of demographic data on individuals tested for COVID-19, as well as diagnoses, hospitalizations, and deaths. The legislative text specifically references “race, ethnicity, age, sex,
in which people with disabilities are experiencing additional, disparate impacts of the COVID-19 crisis beyond the infection itself. These only serve to reiterate the importance of widespread and comprehensive data collection to understand these and other burdens faced by the disability community.

We recognize that recently enacted legislation (P.L. 116-139) mandates the collection of race and ethnicity data regarding COVID-19 testing and morbidity. This data will be critical to document the disproportionate impact of the pandemic on people of color and to begin to develop measures to mitigate immediate and long-term issues of health disparities and access to treatment. However, we will not be able to understand the intersectionality of race, age, disability, geography, and chronic illness without collecting disability data. Current case reporting forms used by the CDC to track COVID-19 cases do not include disability identifiers, and as such, we do not know the full impact of the virus on people with disabilities. It is essential that the public health response include an understanding of the impact that disability status has on increasing risk and vulnerability for minority populations, older adults, those in high-risk geographic areas, and other populations.

Our request to disaggregate data by disability status reflects the underlying realities faced by people with disabilities. Many people with disabilities already face significant inequities in income, education, employment, housing, transportation, and community participation. Disability status is also correlated with poorer health and health care outcomes.

Populations with physical, cognitive, psychiatric, or communication disabilities have decreased access to preventative care, including wellness, fitness, smoking cessation and weight loss interventions. Deaf populations, for example, are 7 times more likely to have low health literacy compared to hearing populations; people with disabilities are 30% more likely to be obese, are over 2 ½ times more likely to develop diabetes and are around 3 times more likely to develop cardiovascular disease. People with disabilities face negative attitudes and biases; inaccessible environments and equipment; inaccessible written, verbal, and electronic health care communication; and failure to provide accommodations and other modifications to policies, practices, and procedures.

These disparities are likely to lead to decreased access to or denial of COVID-19 treatment. Even before the current public health emergency, cause of death for people with disabilities, including people with intellectual and developmental disabilities, was too often inaccurately attributed. Independent data collection efforts, including the recently released COVID-19 & Disability survey conducted by the American Association on Health and Disability, have shown that people with disabilities are already facing significant disruptions in their ability to access regular health care.

guidance on data collection standards under this section in 2011.
care treatment during the outbreak.\textsuperscript{2} This underscores the importance of a coordinated federal effort to collect and report critical data on COVID-19 and disability.

We also know that data alone are not enough. The federal government has a responsibility to direct support, information, and resources to people with disabilities, who are disproportionately impacted by this pandemic. Disability researchers, medical experts, and advocacy organizations have the necessary expertise to address these issues and are embedded in the disability community. These individuals and organizations are well-positioned, as trusted thought leaders and health care service providers, to collect data and conduct research toward creating holistic solutions to remedy the inequitable impact of this and future public health crises.

They are also well-situated to investigate how various demographic characteristics intersect to influence testing and treatment response. They can be deployed in partnership with public health departments to develop rapid response interventions to target the unmet needs of people with disabilities, such as the need for personal protective equipment (PPE), protections for their direct service workers, adequate medications and medical supplies, access to disability component mobile health units, COVID-19 testing in the community, and telehealth services, particularly for those who lack internet access.

The combination of limited resources and high needs creates a relatively thin “margin of health” and much greater vulnerability to communicable disease for people with disabilities. Additionally, people with disabilities face barriers to accessing care that are only exacerbated during the pandemic, including those outlined above. Consequently, people with disabilities appear to be at significantly higher risk of COVID-19 exposure, infection, serious and life-threatening symptoms, hospitalization, and death.

This grim reality is demonstrated most starkly in the shockingly high rates of COVID-19 deaths reported in nursing homes, psychiatric facilities, and other residential facilities throughout the U.S. It may also account for at least some of the racial and ethnic disparities observed thus far during the pandemic, because disability rates are much higher within African American, American Indian, and Native Alaskan populations. People with disabilities share many of the same risk factors of these populations. Real-time tracking of COVID-19 morbidity and mortality among people with disabilities and chronic conditions will be essential for developing targeted interventions in testing, treatment, economic support, and, ultimately, vaccination.

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Thank you for considering our request to recommend the government-wide collection and reporting of data on COVID-19 testing and diagnoses, treatment, and outcomes (including recovery) disaggregated by disability status. We support the critical mission of the newly formed Task Force during this unprecedented public health crisis and stand ready to provide political and

technical support for your efforts in disability measurement, needs assessment, program
development, and evaluation. If the DRRC can be of any assistance, please do not hesitate to
contact the DRRC coordinators, Bobby Silverstein, Peter Thomas, and Joe Nahra, at 202-466-6550 or by email at Peter.Thomas@PowersLaw.com, Bobby.Silverstein@PowersLaw.com, and

Sincerely,

The Undersigned Members of the Disability and Rehabilitation Research Coalition

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National Association of State Head Injury Administrators  
National Multiple Sclerosis Society  
Paralyzed Veterans of America  
United Spinal Association

Supporting Organizations

American Foundation for the Blind  
American Psychological Association  
Autistic Self Advocacy Network  
Center for Public Representation  
CommunicationFIRST  
Disability Rights Education and Defense Fund  
Easterseals  
Justice in Aging  
Lakeshore Foundation  
RespectAbility  
National Association of Councils on Developmental Disabilities  
National Center for Parent Leadership, Advocacy, and Community Empowerment
Appendix: Need for Disability Status Data Collection - Burdens Faced by People with Disabilities

We believe that people with disabilities are experiencing disproportionate and unique burdens from COVID-19 that are currently under-recognized and, therefore, under-addressed. Without systematic data collection regarding disability status, we cannot hope to mitigate or address these burdens in any meaningful way. For example:

1. People in congregate living facilities are experiencing exorbitant rates of COVID-19 infections, and in many states about half of the reported COVID-19 deaths occurred in these facilities.\(^3\) Overwhelmingly, we believe these cases represent people with disabilities who require personal care, supervision, or mobility assistance. People who require intimate personal care from direct service workers are unable to follow social distancing guidelines. Improved data collection would allow us to explore questions such as:
   a. How many individuals reside in these facilities as their “home” and how many have been placed there temporarily as a result of acute illness and a need for subacute rehabilitation or wound care?
   b. Which individuals would be able to live in the community if they had adequate accessible housing or personal assistant services?
   c. How many of these infected individuals have multiple chronic illnesses, and what are they?

2. We are learning from post-acute rehabilitation providers that people who were not disabled before COVID-19 infection are experiencing significant functional problems due to the virus.\(^4\) We need to be able to track this population of “newly disabled” patients to document their “long tail” of rehabilitation and recovery, and to consider the potential for permanent disability even when the initial infection has passed.

3. Direct service workers who care for people with disabilities in nursing homes and the community have been an “invisible” workforce during the COVID-19 emergency.\(^5\) Getting access to proper personal protective equipment (PPE) and infection control training has been slow to non-existent, especially for those who are independently contracted by Medicaid waiver programs to care for a single person in their home setting. This critical but low-paid workforce may be the single most important factor that keeps people with disabilities out of...

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emergency rooms, hospitals, and nursing homes. Without proper disability-focused data, interventions to improve the availability and safety of these workers are less likely.

4. For people with intellectual/developmental disabilities (I/DD), COVID-19 appears to have a particularly outsized impact. We need systematic collection of data to understand why, including the barriers to testing, risk, and adherence to infection control measures for these populations. We know, for example, that people with I/DD may have difficulty understanding the directions for tests, treatment, and other services, and they may need the support of a familiar person, take extra time, or require other accommodations. It may be difficult to quarantine a person who is COVID-positive in a group home, yet many alternative facilities being constructed for the care of COVID-19 patients who do not require hospitalization have not taken into consideration the needs of people with disabilities, leading to many being refused admission. If hospitalized, people with I/DD may need a support person to be allowed to shelter in place with them to avoid preventable problems due to poor communication, inaccurate information, or behavioral difficulties that can flare up in unfamiliar and frightening circumstances.

5. We know that people with disabilities are 30% more likely to be obese, are over 2.5 times more likely to develop diabetes, and are approximately 3 times more likely to develop cardiovascular disease. These chronic illnesses also impact COVID-19 morbidity and mortality. Better data will allow researchers to examine whether people with disabilities who have these chronic illnesses are at the same or higher risk as people without disabilities.

These examples represent only a small sampling of the burdens that people with disabilities face during the pandemic, and data collection and reporting will be the first step towards designing a comprehensive response to protect this population.

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February 19, 2020

Dr. Marcella Nunez-Smith  
Chair, Biden-Harris COVID-19 Health Equity Task Force  
Office of Minority Health  
Department of Health and Human Services  
Tower Oaks Building, Suite 100  
1101 Wootton Parkway  
Rockville, MD 20852

Re: COVID-19 Related Research: Increase Funding for National Institute on Disability, Independent Living, and Rehabilitation Research; Budget Justification Language Request on Research Priorities for Federal Research Agencies

Dear Dr. Nunez-Smith:

On behalf of the Disability and Rehabilitation Research Coalition (DRRC), we write to support a recommendation to increase COVID-19-related funding in Fiscal Year (FY) 2022 for the National Institute on Disability, Independent Living, and Rehabilitation Research (NIDILRR). We hope you will consider recommending that the President include such funding in his FY 2022 budget to support the aims of the COVID-19 Health Equity Task Force. As federal research agencies continue to grapple with the impact of the COVID-19 pandemic, the DRRC has urged the federal government to prioritize research and data collection on COVID-19 relating to people with disabilities, who are at extremely high risk for COVID infection, serious complications, and death. Increasing NIDILRR funding specifically to fund COVID-19 disability research will be crucial to developing and strengthening the federal pandemic response for people with disabilities to address preexisting, exacerbated, and newly created health and social inequities.

COVID-19 and Disability-Related Research Funding at the National Institute on Disability, Independent Living, and Rehabilitation Research

NIDILRR’s mission is to generate and promote new knowledge that improves the abilities of people with disabilities to perform activities of their choice in the community. Not only are people with disabilities an at-risk minority group facing significant health disparities even before the pandemic, case reports are indicating that the long-term clinical complications of this virus are leading previously nondisabled individuals to develop potentially lifelong disabilities as a result of their infection.

In the final FY 2021 omnibus spending bill, Congress appropriated a $1 million increase for NIDILRR’s activities. NARRTC, the professional association of NIDILRR grantees, has circulated a call to action requesting an additional $5-10 million funding increase for NIDILRR to address the urgent research and policy questions around the impact of COVID-19 on the
disability population, for a total NIDILRR budget of $118-$123 million. On behalf of the members of the DRRC, we urge you to support this funding increase in the President’s budget to ensure that NIDILRR can appropriately address the dire research needs stemming from this pandemic.

**Budget Justification Language on Disability Research Priorities at the National Institutes of Health, Centers for Disease Control and Prevention, and Other Federal Research Agencies**

While additional NIDILRR funding is crucial to developing applied research for long-term outcomes in the disability population, NIDILRR is not the only federal agency conducting critical disability research. Therefore, we are also requesting specific language to be added to the President’s budget emphasizing the importance of disability research across all federal agencies conducting COVID-19 research, expected to be a major priority for federal funding over the coming fiscal year. We respectfully propose the following language:

“The Administration directs the National Institutes of Health, the Centers for Disease Control and Prevention, and other federal agencies conducting research on COVID-19 to ensure that their research plans address and prioritize disability, independent living, and rehabilitation research, especially regarding COVID-19 testing and diagnoses; treatment, including admissions and denials to hospitals and intensive care units, denials of needed accommodations, virus interactions with chronic disease and disability, and recovery patterns; and outcomes, including morbidities, mortalities, rehabilitation, and recovery for individuals with disabilities.”

Thank you for your consideration of our requests. We look forward to continuing to work with you to advance public policy impacting people with disabilities. If you have any questions, please contact the DRRC coordinators at the email addresses listed below.

Sincerely,

Bobby Silverstein, DRRC Coordinator
Bobby.Silverstein@PowersLaw.com

Peter Thomas, DRRC Coordinator
Peter.Thomas@PowersLaw.com

Joe Nahra, DRRC Coordinator
Joseph.Nahra@PowersLaw.com
The COVID-19 Pandemic and Americans with Disabilities: A Call to Action for the National Institute on Disability, Independent Living and Rehabilitation Research

People with disabilities are at much higher risk of COVID-19 exposure, infection, hospitalization, and death than those without disabilities. This situation is most evident in the shockingly high rates of COVID-19 mortality among residents of nursing homes and other residential facilities, nearly all of whom are disabled. Most adults with disabilities live in the community, but they share many of the same risk factors as those who live in institutions (e.g. multiple comorbid health conditions, older age, frequent interactions with medical professionals and service providers). It is therefore likely that the recent surge in excess COVID-19 mortality is also concentrated among people with disabilities.

People with disabilities are an at-risk minority group that already faces significant inequities in income, education, employment, housing, transportation, and community participation. Disability also intersects with other sources of disadvantage and discrimination – with higher rates among racial, ethnic, gender and sexual minorities. By definition, people with disabilities also have chronic health conditions or injuries, and therefore require much higher levels of healthcare and support services than those without disabilities. This combination of limited resources and high needs creates a relatively thin “margin of health” and a much greater vulnerability to communicable disease.

A growing body of case reports also indicate that an unknown but potentially large number of survivors are experiencing long-term clinical complications of COVID-19, including impaired function of the lungs, heart, brain, kidneys and circulatory system. People with pre-existing chronic health conditions and disabilities will be at higher risk for these complications, but a portion of previously nondisabled individuals will also develop disabilities as a result of COVID-19, increasing demand for disability benefits and support services.

The pandemic and associated economic and political disruptions raise urgent research and policy questions that will require a coordinated and sustained federal response, including:

- How to protect people with disabilities from COVID-19 infection, hospitalization and death.
- How to minimize the adverse health effects of pandemic policies, including quarantine, social distancing and restricted access to medical, rehabilitative and social services.
- How to strengthen social support networks, including relationships with family, friends and peers with disabilities, and maintain critical disability services such as paratransit, rehabilitation counseling, personal assistance, and independent living services.
- How to cope with significant economic disruptions, including unemployment, wage loss, loss of health insurance, food insecurity, eviction, and catastrophic healthcare costs.
- How to synthesize new and emerging information about COVID-19 and disability, including vetting, curating, organizing and updating population-specific resources.

1 The World Health Organization defines excess mortality as a surge in population death rates above pre-crisis levels. In this case, excess mortality can be attributed to people who died at home with COVID-19 but weren’t diagnosed, and people who didn’t obtain critical medical care because of hospital crowding and/or restricted access to clinics and healthcare providers.

2 Although current public health data is limited, it seems likely that people with disabilities will also be at greater risk of “deaths of despair” from drug or alcohol abuse and suicide, due to increased unemployment, social isolation, depression and anxiety.
The National Institute on Disability, Independent Living and Rehabilitation Research (NIDILRR) is the federal agency best positioned to lead this national effort. NIDILRR is the research arm of the Administration for Community Living (ACL) at the US Department of Health and Human Services. The NIDILRR director holds a government-wide leadership role in this area, as the position is traditionally designated by the Secretary of HHS to lead the Interagency Committee on Disability Research (ICDR). ICDR promotes “coordination and cooperation among federal departments and agencies conducting disability, independent living, and rehabilitation research.” To fully assume this expanded federal role, NIDILRR will need dedicated Congressional support. Unfortunately, NIDILRR’s adjusted annual budget peaked in 2001, and did not return to the pre-sequestration funding level of $112 million until Fiscal Year (FY) 2020.

The Bipartisan Disabilities Caucus in the US House of Representatives has called for a $3m increase in NIDILRR funding for FY2021, but this allocation was recommended before COVID-19 began to sweep through the disability community like a wildfire. NARRTC, the professional association of NIDILRR grantees, now calls for an additional $5-10 million funding increase in FY2021 for COVID-19 related research and program administration, for a total NIDILRR budget of $120-$125 million.

To build the case for how this funding could be used in the disability research community, NARRTC president Jae Kennedy and executive committee member Kathleen Murphy issued a call to members, asking for examples of realizable, ready-to-implement project supplements and ideas for new research projects. In less than two weeks, we received dozens of responses, which we organize by research domain in Appendix 1. In summary:

- Many NIDILRR grantees have already modified their work to encompass COVID-19 research questions, but these efforts could be sustained and focused, without interrupting pre-pandemic programs, by providing relatively modest supplemental funding.
- More urgent research on the public health and economic consequences of the pandemic should be funded under new grant NIDILRR competitions, with expedited review and award procedures.
- The time to study the impact of COVID-19 among people with disabilities is now.

Though no researcher would have chosen to be in these circumstances, we are in the midst of an enormous natural experiment, where environmental changes have confounded the effect of many current experimental and quasi-experimental research interventions. With supplemental funding and appropriate design changes, some of these projects could pivot to timely and relevant studies of pandemic outcomes.

For example, a group activities intervention to reduce social isolation among young adults with autism spectrum disorder would be disrupted by stay-at-home orders, but a pre-post comparison could now be used to assess the impact of suddenly restricted peer networks on cognitive and emotional function. Likewise, special populations of adults with disabilities whose conditions are not studied by other federal research institutes (e.g. people with spinal cord injury, traumatic brain injury, or serious burns) are experiencing dramatic disruptions in their medical, rehabilitative and social supports, and NIDILRR researchers can help document the impact of these disruptions and then develop and test interventions to help minimize them.
Supplemental research funding for current NIDILRR projects, like the already established research supplements to existing National Institutes of Health (NIH), National Science Foundation (NSF), or Agency for Healthcare Research and Quality (AHRQ) grants, could provide invaluable insights on how to support people with serious mental or physical health problems during a public health crisis, informing not only the COVID-19 pandemic but future national emergencies.

New funding will also be required to evaluate the impact of the pandemic on people of all ages with disabilities, including children with individualized education plans (IEPs) whose schools are now closed, parents providing in-home special education, high school graduates with disabilities considering current college or work options, college students with disabilities wondering if they can and should continue their undergraduate or graduate studies, working-age adults with disabilities deciding whether to seek employment or enroll in federal disability programs, and retirees with disabilities evaluating housing options that protect their economic and physical well-being. The size and attributes of the population of newly disabled by COVID-19 must also be determined, particularly if we hope to fully realize the gains of early medical, rehabilitative and social interventions.

In the context of trillions of dollars in new federal spending, investing several million in new COVID-19 related disability research is a smart investment that will impact a wide swath of constituents. We urge the House and Senate to support this research component of the federal budget.

Sincerely,

Jae Kennedy, NARRTC President, Washington State University, WA

NARRTC Executive Committee Members:

- Dana Barton, Rocky Mountain ADA Center, CO
- Jessica Chaiken, HeiTech Services Inc., MD
- Marsha Ellison, University of Massachusetts, MA
- Marianne Farkas, Boston University, MA
- Robert Gould, University of Illinois Chicago, IL
- Mark Harniss, University of Washington, WA
- Rose Mason, Purdue University, IN
- Kathleen Murphy, American Institutes for Research (AIR), TX
- Mark Odum, HeiTech Services Inc., MD
- Sarah Parker-Harris, University of Illinois Chicago, IL
- E. Sally Rogers, Boston University, MA
- Joann Starks, American Institutes for Research (AIR), TX
- Wendy Strobel Gower, Institute on Employment and Disability, Cornell University, NY
Appendix: COVID-19 Research Project Ideas from NARRTC Members

I. Examples of current and proposed COVID-19 research under NIDILRR grants

A. Current (unfunded) research initiated by current NIDILRR grantees

- National needs assessment on how COVID-19 has impacted services in Centers of Independent Living (CILs) (Lex Frieden, Collaborative on Health Reform and Independent Living, TX)
- Survey with CIL consumers in 4 states, asking how COVID-19 has affected their access to employment, education, health care, food, personal assistance services (PAS), food, and ability to live independently (Research and Training Center on Promoting Interventions for Community Living, Jean Hall, KS)
- Interviews with employees with disabilities about their changing workplace accommodation needs during the pandemic (Rob Gould, Great Lakes ADA Center, IL)
- Rapid COVID-related survey of workers with disabilities using Mturk (Amazon’s crowdsourcing platform), including questions about trust in information sources, current health status, presence of high-risk conditions that the Centers for Disease Control (CDC) has identified for COVID-19 complications, safety practices, change in access to medical and community resources, social activity and feelings of isolation (Catherine Ipsen, Research and Training Center on Disability in Rural Communities, MT)
- COVID-19 items and research questions added to a randomized trial on the efficacy of a telehealth pain self-management intervention, a national survey on employer, client, job, and environment-related barriers and facilitators for promoting job retention for people with physical disabilities, and a study on intervening early to support employment for people with Parkinson’s disease (Allen Heinemann, RRTC on Employment and Disability, IL).

B. Supplemental projects that could be conducted by current grantees

- Project to develop and implement interventions to help people who are blind or visually impaired to access the physical distancing signage that is increasingly common in buildings and public spaces (Mike May, Access Explorer, KY)
- Survey of youth and young adults with disabilities with serious mental health conditions on the impact of the pandemic on school, work and career development and related intervention development; and (Maryann Davis, Learning and Working During the Transition to Adulthood Rehabilitation Research and Training Center MA)
- COVID-19 seed grants and mentorships for faculty and postdocs at minority-serving institutions to conduct disparities research (Corey Moore, Langston University Rehabilitation Research and Training Center, OK)
- Virtual workshops and webcasts on inclusive remote learning and work, to proactively invent strategies that ensure full engagement and performance during and after the pandemic (Valerie Fletcher, New England ADA Center, MA)
• Survey of campus and vocational rehabilitation agency staff to assess support services for college students with psychiatric disabilities following campus closures and conversion to online classes (Michelle Mullen-Gonzalez, *Helping Youth on the Path to Employment*, MA)

• Evaluation of mobile health (mHealth) interventions to support the health, function, and independent living of people with disabilities and their families (Bambang Parmanto, *Rehabilitation Engineering Research Center: from Cloud to Smartphone*, PA)

• International virtual summit on emerging initiatives and innovations in increasing employment outcomes for people with disabilities in the COVID-19 crisis and its aftermath (Marianne Farkas, *Center for Psychiatric Rehabilitation*, MA)

II. New project ideas (by NIDILRR research domain)

A. Community living and participation

• Multi-measure study of social, economic and emotional well-being of families that have members with disabilities during and after the pandemic (Chung-Yi Chiu, IL; Malachy Bishop, WI)

• Mixed method study of PAS recipients, evaluating the impact of the pandemic on recruitment, retention, payment, and health practices of attendants and consumers (Jae Kennedy, WA)

• Food insecurity project for people with disabilities during and after the pandemic assessing their nutrition, body mass index (BMI), overall health, and participation in federal, state and local nutrition assistance programs (Debra Brucker, NH)

• Study of work/life balance among people with disabilities who use PAS – remote working and parenting challenges in a public health emergency (LaWanda Cook, NY)

B. Health and function

• Creation of a database of people who either: 1) acquire disabilities because of the sequelae of COVID-19, or 2) already have a disability but become infected with COVID-19; tracking clinical and psychosocial impacts of infection and treatment (Jeffrey Schneider, MA)

• Study of mental and economic stability of adults with serious mental illness as public transportation, supported employment, and disability support services are disrupted by the pandemic (Nev Jones, FL)

• Evaluate the accessibility, use and financing of telehealth and online disability support services for adults with disabilities during and after the COVID-19 pandemic (Gilbert Gimm, VA)

• Project on COVID-19 related disruptions to access to primary care, specialty care, durable medical equipment, and rehabilitative and habilitative services and associated outcomes (e.g. self-assessed health, out-of-pocket healthcare expenditures, and health related quality of life) among adults with disabilities (Noelle Kurth, KS)
C. Employment

- Research on the COVID recession and employment opportunities for workers and job-seekers with disabilities; comparing work and disability program participation rates in the current crisis and the Great Recession of 2008 (Andrew Houtenville, NH)
- Study of how the pandemic has complicated school to work transitions for youth with disabilities, with particular focus on special education, participation in unemployment and Supplemental Security Income (SSI) programs, full and part-time work (Maryann Davis, MA)
- Research on return-to-work decisions with limited job opportunities and heightened health risks (LaWanda Cook, NY)

D. Cross-cutting research activities

- Project on the comparatively low access rural residents with disabilities have to online communities, education, and employment, and how this limited internet access, speed, and penetration accentuate COVID-19 disparities (Catherine Ipsen, MT).
- Research on utilization of telehealth and mHealth services for consumers, families and providers during a pandemic (Brad Dicianno, PA).

E. Activities to promote quality and use of disability research

- Knowledge translation activities to support NIDILRR in its efforts to promote use and adoption of the new knowledge and products to respond to the impact of COVID-19 (i.e. synthesizing existing products that address similar topics, identifying and compiling an inventory of accessible products that meet the needs of specific disability communities, providing a public clearinghouse and hosting dialogues to allow for continued discussion and research in the aftermath of the virus) (Kathleen Murphy, TX).
- Classification and tracking of COVID-19 related technical assistance requests at the 10 regional Americans with Disabilities Act (ADA) centers throughout the US, and development of appropriate and consistent resources for employers and workers with disabilities (Mark Harniss, WA).