August 3, 2022

Gaps in Access to Paid Leave May Lead to Financial Hardships, Report Finds

The Covid-19 pandemic helped shed light on the need for increased access to paid leave for U.S. workers.

Yet more than two years into the pandemic, there are still gaps in access to those benefits. That leaves many workers vulnerable to financial hardships, a new report from the Urban Institute finds.

From: CNBC | Published: July 28, 2022

Caring for a Loved One at Home? State Programs May Pay for Your Help

Nearly 40 million Americans are unpaid caregivers for older or disabled family members or friends. If you’re part of this group, a Medicaid-based Home and Community-Based Services (HCBS) program may pay you to care for a loved one at home.

From: Seniors Matter | Published: July 26, 2022

Caring for My Elderly Parents Taught Me a Hard Lesson About Burnout

Since I began heavy caretaking for my parents last November, everyone around me warned about caregiver burnout. And this month I ran straight into my caregiving wall.

From: MSNBC | Published: July 27, 2022
World Carers Conversation 2022 Session Recordings

World Carers Conversation returned May 19, 2022 as a global, virtual event with the support of EMD Serono and Embracing Carers. Innovators, researchers, policymakers and carer advocates from around the globe learned about advancements in the field, got inspired, and made connections that matter for the future of carers everywhere. These recordings cover the primary conversations and topics discussed.

To view the slide deck used in the event, click here. To view the recordings, click below.

[VIEW RECORDINGS]

Global Voices of Caregiving: A PhotoVoice Project

As part of the World Carers Conversation 2022, NAC worked with Chief Story Officer Anne Levy and caregivers from around the globe to help them tell their own story through a series of photos. These photos addressed simple questions about being a caregiver, such as what works, what doesn’t work, and what a world that embraces and supports caregivers would look like. Each one of these stories is unique to the caregiver’s own experience and tells a diverse story of what it’s like to care for others, an act which connects us all.

To view and hear these stories, please click on the red button below.

[VIEW STORIES]
The National Alliance for Caregiving is proud to present Reimagining Clinical Trial Recruitment Through a Family-Centered Lens: Caregiver Recommendations for Enhancing Clinical Trial Participation Diversity, a new report developed with sponsorship from Travere Therapeutics. Building on the priority area of clinical trials recruitment and retention identified in NAC’s 2019 report Paving the Path for Family-Centered Design: A National Report on Family Caregiver Roles in Medical Product Development, this report synthesizes outputs from a series of three roundtables. In collaboration with Health Leads, NAC mobilized its network of researchers, policymakers, innovators, patients, families and caregivers to discuss and develop strategies for improving how researchers engage caregivers from populations most impacted by structural inequities in order to better understand how to increase representation of diverse patients in clinical trials.

Included in this report:

- Summary of caregiver expert convening including their key insights on caregivers’ role in clinical trials and barriers to patient and caregiver participation;
- Summary of clinical trial roundtable including current strategies that improve caregiver engagement and support caregiver roles related to trials;
- Direct quotations from caregiver experts about their experiences with clinical trial participation;
- Outputs of a co-design session with both caregiver experts and clinical trial experts;
- Individual-level, community-level and systems-level focus areas for acting on these co-developed recommendations.

NAC’s report underlines that engaging caregivers more effectively in the clinical trials process to promote better trial enrollment diversity requires investment and collaboration from not only industry partners and sponsors, but also from federal regulatory agencies, policymakers, healthcare leaders, community care providers and patients and caregivers themselves. Recognizing the value – the contributions, the expertise, the health impact – of what caregivers do, as partners in innovation, is a key part of creating a more equitable, person and family-centered health care system.

To read the report and access our findings, click on the red button below.
The Circle of Care Guidebook Series

The National Alliance for Caregiving’s Circle of Care library is a series of guidebooks dedicated to providing resources, support and information to caregivers caring for someone in a specific disease space. Part of an ongoing effort of NAC, these first three guidebooks were developed with the assistance of both patient-advocacy and consumer-facing partners:

- Circle of Care: A Guidebook for Mental Health Caregivers
- Circle of Care: A Guidebook for Caregivers of Children and Adolescents Managing Crohn’s Disease
- The Circle of Care Guidebook for Caregivers of Children with Rare and/or Serious Diseases

NAC is proud to lend their expertise on caregiving through these guidebooks. To access the guidebooks for yourself and to share them with other caregivers in your life, please visit the link below.

VIEW THE GUIDEBOOKS

EVENTS & OPPORTUNITIES

AUGUST

CONFIDENCE: Financial Education & Training for Latino Family Caregivers

Thursdays, July 21 - August 18, 2022, 4:00 PM – 7:00 PM EDT | Online

CONFIDENCE is a free, 5-week virtual financial education program made for Latino family caregivers who help a loved one with dementia, including Alzheimer’s Disease, and/or memory problems.
As part of this course hosted by the University of Southern California's School of Gerontology, you will learn how to:

- Lower your out-of-pocket care expenses
- Find community resources
- Manage someone else's money
- Balance employment and caregiving
- ... and more!

To register for this class, email Alex Gonzalez by clicking on the red button below.

**EMAIL TO REGISTER**

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**Engaging Caregivers: A Program for Healthcare workers**

**August 3rd, 10th, and 17th, 2022**

WellMed's Caregiver Teleconnection sessions are one hour learning sessions for caregivers of older persons. Experts from around the country share information important to caregiving. Caregivers will have the opportunity to ask questions as well as hear from other caregivers.

Our sessions are geared for the family members taking care of a loved one over the age of 60. Professionals are also welcome to participate as well.

Learn about defining roles and responsibilities with patients and caregivers, the role of diversity and culture, and where to find resources to help with difficult situations and elder abuse.

Upcoming Caregiver Teleconnection sessions will be held on August 3rd, 10th, and 17th. Click on any of the three links below to learn more or register for each event. A certificate of attendance will be provided upon request.

Registration links:

August 3rd at 11:00 am EST: [Defining Caregivers & Recognizing Their Experiences](#)

August 10th at 11:00 am EST: [Difficult Situations & Elder Abuse](#)

August 17th at 11:00 am at EST: [Diversity & Caregiving](#)

To view the Caregiver Teleconnection flyer, click the red button below.

**VIEW THE FLYER**
Why Do They Do That? Managing Common Challenging Dementia-Related Behaviors

August 4, 2022 | 11:00 AM - 12:15 PM EST | Online

Conversations with Dr. Tam Cummings - A Monthly Education Series for the Dementia Caregiver

Dr. Tam Cummings will discuss common behaviors witnessed by dementia family caregivers including:

- Cursing
- Movement changes
- Hunting and gathering
- Accusations of theft
- Taking away the car keys
- Sundowning
- Bathroom issues, including toileting and bathing
- Bedtime issues

We will also review the five senses which addresses changes leading to potentially dangerous or annoying behaviors. Stress relief tips will be shared, and handouts will include materials to self-measure caregiver stress and self-compassion.

Questions can be directed to VirtualSeminars@arden-courts.org.

To register for this event, click the red button below.

Cultivating Positivity in Caregiving: Research-Based Ways to Promote Happiness and Well-being

August 4, 2022 | 12:30 PM - 1:30 PM EST | Online

Research shows that caregiving and navigating day-to-day life with memory and movement disorders can be positively impacted by planting seeds of happiness.

Join Banner Alzheimer’s Institute & Banner Sun Health Research Institute as we share research-based ideas to cultivate happiness, well-being and success for caregivers and people living with memory and movement disorders. Research proves that happiness not only lifts spirits, it can also improve overall quality of life and well-being. Learn research-based tips to improve coping skills, emotional well-being and quality of life throughout the dementia journey.

To register for this event, click the red button below.
Peer Support Mentoring Program – Spirituality Calls

The VA Peer Support Mentoring (PSM) Program and the National Chaplain Office will be hosting a 12-month series addressing spirituality and caregiving. These calls are open to caregivers of Veterans enrolled in both the Program of General Caregiver Support Services (PGCSS) and the Program of Comprehensive Assistance for Family Caregivers (PACAF).

Calls will be held on the first Friday of each month from 12:00 p.m. to 12:30 p.m. ET (11:00 a.m. to 11:30 a.m. CT; 10:00 a.m. to 10:30 a.m. MT; 9:00 a.m. to 9:30 a.m. Pacific) and will be hosted on WebEx using the dial-in information below. There is NO registration required. Meeting Call in number: USA Toll-free Number 1-833-558-0712. Meeting number (access code): 2762 030 9378

Topics:
- August 2022 - Spirituality and Grief
- September 2022 - Spirituality and Self Reflection

To learn more, click the red button below.

Webinar: Value-based Care and the Geriatric ED

August 8, 2022 | 3:00 PM - 4:00 PM EST | Online

Join Geriatric Emergency Department Collaborative (GEDC) for this webinar on of Value-Based Care and how it is relevant to older adult care in the emergency department.

Goals of this webinar

1. To describe the concept of Value-Based Care and how it is relevant to older adult care in the emergency department
2. To review the connection between Accountable Care Organizations and Geriatric EDs
3. To discuss opportunities and strategies for Geriatric ED champions to partner with local Value-Based Care organizations
Approximately 10% of individuals dually eligible for Medicare and Medicaid under the age of 65 have an intellectual disability or related condition. In general, Americans living with disabilities receive less preventive care, have a higher incidence of chronic conditions, and visit the hospital and emergency department more often — leading to much higher health care spending than for adults in the general population. Providers and health plans are increasingly using personalized approaches in a managed long-term services and supports (LTSS) environment to support people with intellectual and developmental disabilities (I/DD) in achieving their individual goals.

A recent policy brief by the Institute on Community Integration at The University of Minnesota estimated that at the current rate of closure, all large state-run I/DD facilities could either close or limit service to fewer than 15 people per setting by 2025. The trend of continued facility closures means the number of people with I/DD living in smaller person-centered community settings will continue to grow, placing increased responsibility on more community-based providers and health plans to better serve people with I/DD. Enhanced care coordination strategies, increased caregiver support options, and improved awareness within provider networks must be part of the plan to further support the unique health care needs of people with I/DD.

This webinar will discuss strategies for providing coordinated care for individuals with I/DD, including person-centered medical assessment considerations for individuals with I/DD, risk stratification strategies to drive interventions, and family or caregiver support, education, and outreach. Speakers will describe the current gaps in care for individuals with I/DD and share strategies for delivering person-centered care for individuals with I/DD through use of interdisciplinary teams (IDTs) and innovative care coordination models. The event will conclude with an opportunity for participants to ask the featured speakers questions during a live Q&A session.

By the end of this webinar, participants should be able to:
1. Describe the current gaps in care for individuals with I/DD;
2. Provide strategies for delivering person-centered care for individuals with I/DD through use of innovative care coordination models; and
3. Understand the impact of these strategies on outcomes of care.
Caregiving Basics: Medication Administration

August 9, 2022 | 9:00 PM - 10:00 PM EST | Online

This training will go over everything you need to know about Medication Administration, basic procedures, and protocols of medication administration to comply with the title code regulations 22 requirements to provide training to direct care staff who administer and manage medication for residents.

As DSP and Direct Care Staff, we play a crucial role when it comes to administering medication to patients. This training will provide you with an overview of how to successfully administer it. Along with some challenges you may face and how to best address them. The Training will be conducted by our Pharmacist Partner; by the end of this training, you will be able to:

1. Understand your roles and responsibilities when administering medications.
2. Understand some challenges you may face and how to best address them.
3. Understand day-to-day situations that may cause errors when doing so.

To register for this event, click the red button below.

Healthy Aging: Creating Age-Friendly Health Systems

August 10, 2022 | 1:00 PM - 2:00 PM EST | Online

The nation’s adult population over age 65 is projected to reach 83.7 million by the year
2050, an increase from 21% of the population in 2012 to more than 39% in 2050. Age-Friendly health systems is an initiative of The John A. Hartford Foundation and the Institute for Healthcare Improvement in partnership with the American Hospital Association and the Catholic Health Association of the United States. The initiative is designed to meet the needs of older adults, looking beyond acute events, engaging the whole community, and achieving better health for older adults. By focusing on four key areas—what matters, medications, mobility and mentation—we aim to improve patient care, safety and outcomes; and reduce length of stay and readmissions. Join this webinar to learn about the importance and impact of the 4Ms Framework, hear bright spots in the field integrating value-based, Age-Friendly care and understand the benefits of joining the Action Community.

To learn more about this webinar, call the American Hospital Association at (312)422-2609.

To register for this event, click the red button below.

REGISTER

Take a Breath! Tips from a Caregiver Coach

August 11, 2022 | 12:00 PM - 1:00 PM EST | Online

Caregiving is complicated. Knowing where to turn and how to care for yourself can be overwhelming. Caregiver coaches get it—they have real-life experience, tips, and tools to help you! Join the Ontario Caregiver Organization’s webinar with a Caregiver Coach to:

• Discover helpful OCO tools and resources
• Try some practical 5-minute self-help activities you can do TODAY
• Learn how to create a self-care plan with helpful tips

Our Speaker
Suzanne McKenna has been a caregiver for 15 years for her son with a severe traumatic brain injury due to an accident. With a certification in Motivational Interviewing/Counselling and ten years of work experience as a System Navigator, she has many skills in compassionate support. As a caregiver coach and mentor, Suzanne guides the caregiver in building confidence through their ups and downs, helps find resources and encourages them to find time for themselves without guilt.

Information
• Please note that this presentation is available in English only.
• For any difficulty with registration or any other questions, please contact us at info@ontariocaregiver.ca or call 1-888-877-1626 X 1011
Balancing Caregiving and Work

August 11, 2022 | 6:30 PM - 7:30 PM EST | Online

Join the Ontario Caregiver Association for a webinar about striking the right balance between caregiving and professional commitments. Learn more about caregiving experiences, explore the best strategies for balancing work and caregiving, and gain access to resources focused on their needs.

The Caregiver Circle

August 11, 2022 | 7:00 PM - 9:00 PM EST | Online

The Caregiver Connection brings family caregivers, expert guests, and those who may become caregivers together every month to talk about challenges, learn about specific areas of knowledge and to share. Our aim is to help you be more proactive, find the gifts and hear from knowledgeable people in the caregiving space.
10 Warning Signs of Alzheimer's

August 16, 2022 | 4:00 PM - 5:30 PM EST | Online

Alzheimer's and other dementias cause changes in memory, thinking and behavior that interfere with daily life. Join Kadlec Community Health to learn about 10 common warning signs and what to watch for in yourself and others.

To register for this event, click the red button below.

Dementia 101: The Basics

August 17, 2022 | 6:00 PM - 7:00 PM EST | Online

If you’ve heard of dementia (or many of the other "memory loss" terms), but aren't sure what it is or you want to learn more, this webinar can help! Spend an hour learning the basics of all things "dementia". You're invited to bring your questions for an open Q+A!

To register for this event, click the red button below.
Spotlight Series: Let’s Talk About Driving

August 22, 2022 | 2:30 PM - 4:00 PM EST | Online

This educational event hosted by the Alzheimer Society of Durham Region (Canada) will discuss driving from the perspective of an older adults, people living with Mild Cognitive Impairment or people living with dementia and their families. Guest speakers from Alzheimer Society Southwest Partners and the Thames Valley Health Team will provide information on warning signs, retirement from driving, and the role of healthcare providers and the Canadian Ministry of Transportation. All populations are welcome to attend this presentation.

**Healthcare providers are welcome to attend, but this event will not provide training on how to assess fitness to drive.**

This session will be hosted on Zoom (with a phone in option)

Zoom session: Once you have registered for the session, you will receive a link with instructions on how to connect.

For Technological Support please contact the Community Education Team for education@alzheimerdurham.com

To register for this event, click the red button below.

REGISTER

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Emergency Preparedness Planning For Persons With Disabilities Webinar

August 24, 2022 | 2:30 PM - 4:00 PM EST | Online

During emergencies and natural disasters, health plans play a critical role in addressing and protecting their members’ health, safety, and independence. For persons with disabilities and their caregivers, it is crucial for health plans to consider individual circumstances and needs in order to effectively prepare for emergencies and disasters. This webinar will provide information on creating and strengthening emergency practices embedded in individualized care plans and health plan processes. Enhanced emergency practices will assist members and their caregivers in navigating emergencies and natural disasters.

By the end of this webinar, participants should be able to:

1. Understand populations and identify persons with access and functional needs (AFN) that may include individuals with disabilities, limited English proficiency, limited access to transportation and financial resources, older adults and others deemed at-risk.
2. Identify the role of care management and care coordination in including emergency preparedness planning in member care plans, including identifying members at high risk during emergencies.
3. Identify steps health plans can take to prepare members and their caregivers for emergencies and natural disasters.
4. Understand the benefits of emergency preparedness for dually eligible individuals.

Intended Audience:
This webinar is intended for providers, health plans, and health care organizations interested in creating or bolstering their emergency preparedness processes for individuals with disabilities.

To register for this event, click the red button below.
The Grief Journey of the Dementia Caregiver

August 31, 2022 | 1:00 PM - 2:00 PM EST | Online

This program offers clinicians a breakdown of the types and stages of grief that affect caregivers of persons living with dementia. We will discuss how to identify the area of stress that is most burdensome to each particular caregiver and ways to lessen that burden in order to help that caregiver continue to provide optimal care.

To register for this event, click the red button below.

Integrating Technology into Caregiving

August 31, 2022 | 7:00 PM - 9:00 PM EST | Online

Join us for a Zoom webinar where we learn how to integrate technology into caregiving for a person living with dementia.

Have you ever wondered if there is technology available that can assist you with caregiving and enhancing safety in the home?

Join us for a presentation with guest speakers Jennifer Watt, Occupational Therapist with the Behavioural Response Team in Geriatric Mental Health at LHSC and technology expert Ron Beleno, Consultant, Coach and Mentor from RB33. As a caregiver to his father who lived with Alzheimer's for 10+ years, Ron utilized technology and creative strategies to support his family's life to live well and as best as possible.
Together, Jenny and Ron will showcase a broad range of tools available to help you with caregiving and address everyday challenges that you might be facing, or potentially could face in the future. They will share information on what these tools can be used for taking into consideration the person living with dementia’s right to privacy, their cost, where to get them, and how to set them up.

Please come ready with your questions about caregiving and technology.

To register for this event, click the red button below.

REGISTER

SEPTEMBER

Webinar: Navigating a Dementia Diagnosis and Care Needs

September 7, 2022 | 1:00 PM - 2:00 PM EST | Online

Knowing the warning signs of Alzheimer’s disease is important for early detection and diagnosis. Family members often are the first to notice the signs, but may struggle on how to approach their loved one about scheduling an appointment with their doctor. Once they do broach the subject, they may be met with resistance or denial. If a diagnosis is received, the individual and their family may be at a loss for what to do next and what to expect for their loved one’s care needs. As professionals, it is important to help older adults and their families navigate life after a dementia diagnosis by providing them with useful information and support. Join this webinar during World Alzheimer’s Month to learn more about what to do next after a dementia diagnosis, what changes to expect and how to best care for someone living with dementia.

Participants in this webinar will be able to:

- Review the 10 warning signs of Alzheimer’s disease and the importance of diagnosis
- Discuss what to expect after a diagnosis and what changes are likely to occur
- Learn communication strategies and care tips to best support the individual
- Discover resources for professionals and families for diagnosis, care and support
Hopeful Alzheimer's & Caregivers' Symposium

September 10, 2022 | 11:00 AM - 3:00 PM EST | Online

The Hopeful Alzheimer's & Caregivers' Symposium provides education and information for caregivers of loved ones suffering from Alzheimer's disease. The 2022 program will include a video of persons suffering from various types of dementia. Additionally, some of these individuals will join us and answer questions. Dr. Monica Parker, Director of the Goizueta Alzheimer's Disease Research Center at Emory University will discuss the illness and the latest research findings. Mrs. Shondale Coleman of Amedicis Hospice will discuss how to access hospice services, once a person has an Alzheimer's diagnosis. Attorney Joseph Gilsoul will discuss how to apply for Social Security Disability benefits, once an Alzheimer's diagnosis is received.

To register for this event, click the red button below.

REGISTER >

Caregiver Educational Series - "Incontinence Products"

September 13, 2022 | 12:45 PM - 2:30 15 EST | Online

This webinar hosted by the Area Office on Aging of Northwestern Ohio will discuss incontinence products in the context of caregiving. It will be presented by Janet Firestone, National Account Executive Medicaid, Principle Business Enterprises/Tranquility Products.

To RSVP and obtain access instructions, call Jennifer Forshey at (419) 725-6983, or email her by clicking on the red button below

To register for this event, click the red button below.

RSVP >
Dementia Information Series

September 14, 2022 through November 2, 2022 | 7:00 PM - 9:00 PM EST | Online

The Dementia Information Series is a comprehensive 8-week series on **Wednesday evenings** designed for families who are caring for a loved one with dementia or Alzheimer's disease. You can understand the disease, learn how it affects your relative, how to access support in the home and community, how to cope, and how to communicate.

Register as early as possible to reserve a spot for this popular education event.

To register for this event, click the red button below.

REGISTER >

Request for Applications for NIA's Career Development Award Program

**Deadline to apply: September 15, 2022**

The National Institute for Aging's IMPACT Collaboratory seeks to build the nation's capacity to conduct pragmatic clinical trials of interventions embedded within health care systems for people living with dementia and their care partners.

IMPACT Collaboratory has issued a request for applications for its Career Development Award Program. The award is intended for scientists pursuing careers in embedded pragmatic clinical trials for people living with Alzheimer's disease and Alzheimer's disease-related dementias (AD/ADRD) and their care partners.

To learn more, click [here](#) or on the red button below.

LEARN MORE >
Virtual Duke Caregiver Community Event

September 20, 2022

Duke University is hosting a free, virtual event that will offer you the chance to learn about the latest disease research, understand advancements in caregiving, and discover ways to find support as you care for loved ones. After the event is held on September 20, 2022, its presentations and information sessions can be accessed online by those who register. Materials can be viewed at any time on your computer or mobile device through September 2023.

Conference materials will be available virtually in English and Spanish. Presentations will go live on Sept. 20 and cover these topics and more:

- African-American, Hispanic, Latino, and LGBTQ+ experiences in caregiving
- Challenges of long-distance caregivers
- Physical, occupational, and speech therapy options
- Adult guardianship benefits and misconceptions
- Finding help for mental health care
- Advancements in technology to understand health needs

To learn more about this event, click here. To register, click the red button below.

REGISTRER >

OCTOBER

Alzheimer's Program: When Living at Home is No Longer an Option

October 11, 2022 | 4:00 PM - 5:30 PM EST | Online
Changes in thinking may reduce your ability to make appropriate decisions about self-care and your day-to-day needs as the disease progresses. You may be at increased risk for harm, falls, wandering and/or malnutrition. You also may have difficulty managing personal hygiene or household tasks, which can lead to unsafe living conditions. Plan ahead for how you will address your basic needs, including housing, meals and physical care.

If friends or family have expressed concern about your ability to perform certain roles or tasks, listen to their observations. Alzheimer's will eventually limit your own insights as to what you can safely do.

To register, click the red button below.
Social Engagement Study for Care Partners

The University of Illinois is seeking participants for a study about the importance of social engagement in the lives of those caring for loved ones with dementia. In order to be eligible, individuals must be over the age of 60, fluent in English, have access to a computer and internet connection, and be currently caring for a relative with dementia. In addition, participants must be able to attend two 45-minutes sessions each week for four weeks.

To learn more about this study, click the red button below.

LEARN MORE

Lung Cancer Registry

The Lung Cancer Registry at the GO2 Foundation for Lung Cancer has initiated a new worldwide survey focused on the experiences of those caring for patients with lung cancer. The information collected will directly guide caregiver support programs and resource development.

To complete this survey, join the Lung Cancer Registry. Once you log in, you will be taken to the ‘Home’ page of the Lung Cancer Registry where you will find the Caregiver Survey. Every response is the start of a valued conversation.

Anyone with questions or concerns, please feel free to contact a registry coordinator at registry@go2foundation.org.

To participate in the study, click the red button below.

PARTICIPATE IN THE STUDY
Brain Donation: A Gift for Future Generations

Brain donation helps researchers study brain disorders, such as Alzheimer’s disease and related dementias, that affect millions of people. Learn about why people donate their brains, the process of brain donation, and how you can enroll to make this generous gift.

Our brains are amazing, intricate networks that help us think, love, and breathe. But sometimes things go awry and cause brain disorders, such as Alzheimer’s disease and related dementias. By studying the brains of people who have died — both those who had a brain disorder and those who were healthy during life — researchers learn more about how types of dementia affect the brain and how we might better treat and prevent them. Brain donation provides an opportunity to help researchers better understand these disorders, which can lead to improved treatments for future generations.

While many people think that signing up to be an organ donor includes donating their brain, the purpose and the process of brain donation are different. Rather than helping to keep others alive, such as with kidney donation, brain donation helps advance scientific research. One donated brain can make a huge impact, potentially providing information for hundreds of studies. But many brains are needed from diverse populations and ages to help researchers investigate the causes of disease and to develop more effective therapies that can then be applied broadly.

Researchers use donated brain tissue to study brain diseases that affect millions of people. These diseases include Alzheimer’s disease, Lewy body dementia, frontotemporal disorders, mixed dementia, Parkinson’s, and Huntington’s disease, as well as brain injuries such as trauma and stroke.

To learn more about brain donation, click the red button below.

University of Georgia Survey on Access to Health Services
(Georgia residents only)

Elisa Childs, a doctoral student at the University of Georgia’s School of Social Work, is conducting a study examining older Georgians’ access to health care. Georgia has one of the fastest-growing populations of adults 65 and older yet has some of the worst access to health care in the country.

If you are a Georgian who is at least 65 in age, your participation in this 15-minute survey can help bring attention to this critical issue. Survey results will be used for Ms. Childs’s doctoral dissertation and will potentially serve as the foundation for federal and state grant applications aimed to increase access, minimize disparities, and improve health outcomes.
Study: Caregiver Perspectives on Alzheimer’s Disease and Related Dementias Clinical Trials

Our team at the University of Illinois Urbana-Champaign is conducting a study to learn more about engaging family caregivers of racial/ethnic minority participants in clinical trials for Alzheimer’s Disease and related dementias (ADRD).

We’d like to learn more about your experiences with helping your relative navigate participation in clinical trials for ADRD. We hope to learn about ways that we can better support family caregivers so they can better support their relative through these trials.

To participate in this study, the participant must be:

- Age 18 or older
- Speak and understand English
- A caregiver for someone with ADRD who has either a) received information about a clinical trial, b) been invited to participate in a clinical trial, or c) has participated in a clinical trial
- Identify as part of a racial or ethnic minority community

Participation includes a 25-30 minute virtual interview. Participant will receive a $25 gift card upon completing the interview.

Please contact Dr. Minakshi Raj if you are interested in participating by clicking the link below.

Participants Needed for Study on Caregiving for Family Member with Memory Problems

Are you helping a family member with memory problems? The Virginia Tech Center for Gerontology is seeking grandchildren, siblings, nieces/nephews, and step family members (any relationship) living in Virginia to participate in a research study about caring for a relative with dementia living in the community (not in an assisted living or nursing home). This study involves one initial phone interview (approx. 70 minutes) and a brief phone interview for 8 days in a row (15-20 minutes each evening).

You will be asked questions about your experiences providing care, family involvement in care, and other caregiving-related topics. Participation is voluntary and may be terminated at any time.

Please contact Dr. Minakshi Raj if you are interested in participating by clicking the link below.
and use of paid services, and challenges you face caring for your relative.

Compensation is $110 for full participation (VT IRB# 20-742). Interested? Visit the study website careex.isce.vt.edu, call: 540-231-9250 or email us at careex@vt.edu.

To learn more, click the red button below.

IN CASE YOU MISSED IT...

AFPHS Training – Using Data to Address Social Determinants of Health

Data systems that include information on the health of older adults are crucial to developing programs and services that meet their health and social needs. Organizations and agencies can use this information to target resources, identify community partners, support grant applications, and develop strategies to address health disparities and social needs in their communities.

The July 2022 AFPHS training focused on the importance of data in addressing social determinants of health among older adults. This session highlighted how some states have developed and used data on older adults to advance healthy aging in their communities.

RESEARCH & RESOURCES

Alzheimers.gov

Finding Dementia Care and Local Services

A person with dementia will need more care as symptoms worsen over time. Problems
Policies and Resources Identified by Youth as Being Important to Prepare for Caregiving Responsibilities

The Journal of the American Board of Family Medicine's July edition features a new article examining ways to support young caregivers.

**Background:** Youth are increasingly upholding significant caregiving responsibilities. These caregiving responsibilities can have emotional, educational, and professional impacts on youth and young adults. And yet, policies and resources focus on adult caregivers and are limited in supporting young caregivers. The purpose of this study was to describe the different types of support that youth identify as being important to prepare to take care of an adult relative.

**Methods:** We conducted an open-ended, text-message based poll of youth ages 14 to 24 in August 2020. We conducted a content analysis to categorize and describe the different types of support respondents identified in their responses. We compared types of support identified by age-group, gender identity, and prior caregiving experience.

**Results:** Most respondents (42.2%) identified education (eg, skills training) as being an important resource. Other types of support reported included financial support (eg, assistive programs), workplace policies (eg, paid leave), mental health support, and professional support.

**Discussion:** Policy makers should extend existing policies (eg, Family and Medical Leave Act) to include and consider the circumstances of youth and young adults. Policies enabling young caregivers to actively participate in their adult relative's health care visits could be critical to preparing youth for the skills required and the physical and emotional demands associated with caregiving. Coordinated efforts between health and education systems could support youth in learning information about caregiving, medical decision making, and medical tasks.

To read the article, click the red button below.
The Impact of COVID-19 on Carers: An International Perspective

The International Alliance of Carer Organizations has produced a new report that explores international perspectives on caregiving during the Covid-19 pandemic. Building on IACO members' reports, statements and surveys, this study provides an international view of caregivers' needs and the measures individual countries have taken to support carers during the COVID-19 pandemic and beyond.

To view the report, click the red button below.

Alzheimer’s Disease Should Stop Driving

Good drivers are alert, think clearly, and make good decisions. When people with Alzheimer's disease are not able to do these things, they should stop driving. But some people may not want to stop driving or even think there is a problem.

As the caregiver, you must talk with the person about the need to stop driving. Do this in a caring way. Understand how unhappy the person may be to admit that he or she has reached this new stage.
What is Parkinson’s Disease?

Parkinson’s disease is a brain disorder that causes unintended or uncontrollable movements, such as shaking, stiffness, and difficulty with balance and coordination. Symptoms usually begin gradually and worsen over time. Common symptoms include:

- Tremor in hands, arms, legs, jaw, or head
- Muscle stiffness, where muscle remains contracted for a long time
- Slowness of movement
- Impaired balance and coordination, sometimes leading to falls

While the exact cause of the disease remains unknown, many researchers believe that Parkinson’s results from a combination of genetic and environmental factors. There is no cure for Parkinson’s, but there are medicines, surgical treatments, and other therapies that can relieve some of the symptoms associated with the disease.

To learn more about Parkinson’s disease, click the red button below.

How To Evaluate Health Information on the Internet

The internet makes finding health information easy and fast. But, it can also lead you to a lot of false and misleading information. The National Institutes of Health offers tips on how to decide whether the health information you find on the internet is reliable.

To learn more about assessing health information online, click the red button below.
Know the Signs of Elder Abuse and How to Get Help

The mistreatment of older adults can be by family members, strangers, health care providers, caregivers, or friends. Abuse can happen to any older adult, but often affects those who depend on others for help with activities of everyday life. Learn how to recognize some of the signs of elder abuse so you can step in and help. For example, you may notice that the older adult:

- Seems depressed, confused, or withdrawn
- Appears dirty, underfed, or dehydrated
- Has unexplained bruises, burns, cuts, or scars
- Has unpaid bills or recent changes in banking or spending patterns

To learn more about the signs of elder abuse, click the red button below.
A Workbook for Your Workplace Wellness

Our work life now includes our life’s work – caring for a family member with a chronic illness or disease or injury. How do we make room for both experiences in the workplace? How do we take time during our work day to manage our stress and worries? How do we console our colleagues who grieve?

The Caregiving Years Training Academy has come up with a free new tool to caregivers for managing caregiving, grieving and working. A Workbook for Your Workplace Wellness shares tools to help you manage your experiences as an employee, a co-worker and a family caregiver.

The workbook includes a Worksheets section which features tools to use daily and weekly to manage your stress, grief and wellness.

To learn more, click here. To view or download the workbook, click the red button below.

VIEW THE WORKBOOK

VA and Rosalynn Carter Institute for Caregivers Launch Partnership

The VA is committed to caring for Veterans, caregivers, families and survivors. Strong partnerships within and outside of the VA are critical to ensuring caregivers receive the support they need. That’s why the VA is partnering with the Rosalynn Carter Institute for Caregivers (RCI) to further promote the health and well-being of those who care for our nation’s Veterans.
The VA and Rosalynn Carter Institute for Caregivers launched the new partnership to support caregivers of Veterans, caregivers who are Veterans, and caregivers who work for the VA. Both organizations will work collaboratively to increase awareness of the needs of caregivers of Veterans, increase caregiver access to programs, services and resources, and improve caregiver outcomes.

For more information about this partnership and the VA’s Caregiver Support Program, click the red button below.

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Get the Support You Need

The Operation Family Caregiver (OFC) program at the Rosalynn Carter Institute for Caregivers helps create stronger, healthier, more resilient families by supporting the unsung heroes behind our heroes: the caregivers. These friends and family who take care of America’s wounded warriors do so, in many cases, at the expense of their own health and well-being. The challenges that our veterans face can affect the entire family, but services and programs rarely focus on the caregiver.

OFC provides eight free and confidential one-on-one coaching to the families, friends and supporters of those who have served our nation and returned home with either visible or invisible injuries. OFC is tailored to each caregiver’s unique needs and delivered virtually on their schedule.

For more information about Operation Family Caregiver, click [here](#). To enroll, click the red button below.

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Ignite Change as a Caregiver Advocate

This fall Rosalynn Carter Institute (RCI) for Caregivers is launching its first-ever grassroots advocacy network to fundamentally change the way caregivers are seen and supported. Will you join the movement?

Over the past two years, the essential role of our country’s 53 million caregivers has been elevated to the national stage, emphasizing the fragmented, inaccessible, and oftentimes nonexistent systems in place to support this indispensable population. In the words of our founder, former First Lady Rosalynn Carter, we must “seize this moment” to affect meaningful, systemic change.

To learn more or become an advocate, contact the RCI by clicking the red button below.
Caring for a Person with Alzheimer's Disease: Your Easy-to-Use Guide

Get Alzheimer's caregiving information and advice in this comprehensive, easy-to-read guide produced by the National Institute on Aging. Learn caregiving tips, safety information, common medical problems, and how to care for yourself.

To view or download the guide, click the red button below.

VIEW THE GUIDE

MAKE YOURSELF A PRIORITY, TOO
[TIPS FOR CAREGIVERS]

Make Yourself a Priority, Too: Tips for Caregivers

Caregiving can be rewarding, but difficult. Learn how you can put yourself back on the priority list.

Share this infographic and help spread the word about caring for yourself while caring for others. Click on the social media icons above, or copy and paste the URL and post it to your account (Twitter, Facebook, etc.).

To download the infographic, click on the red button below.

DOWNLOAD THE INFOGRAPHIC

How to Share Caregiving Responsibilities with Family Members

Caring for an older family member often requires teamwork. While one sibling might be local and take on most of the everyday caregiving responsibilities, a long-distance caregiver can also have an important role.

As a long-distance caregiver, you can provide important respite to the primary caregiver and support to the aging family member.
Making the Invisible Visible: State Strategies for Identifying and Reaching Family Caregivers

Family caregivers — often relatives, friends, and neighbors — provide vital assistance to support the health and functional needs of older adults, but often face physical, emotional, and financial challenges related to their caregiving responsibilities.

Many states are seeking to better support this important workforce, particularly family caregivers in communities of color, but need an effective way to accurately identify the number, diversity, and unique needs of family caregivers in their state.

In identifying family caregivers, states should recognize cultural considerations, including distinct racial and ethnic perspectives associated with caregiving, that may have implications for identification.

This brief by the Center for Health Care Strategies presents key strategies to help states better identify and reach family caregivers.

To view this brief, click on the red button below.

AARP/National Geographic "Second Half of Life Study"

Older Americans are not only the happiest adult Americans as a whole, but also consider themselves healthier and more financially secure than those in their 40s and 50s, reveals a new study published in AARP Bulletin. The "Second Half of Life Study," conducted with National Geographic, surveyed thousands of adults age 18 to 90 to explore how Americans perceive their current life, their expectations for the future and aging in general as we emerge from the COVID pandemic. About two-thirds of adults over 80 said that they were living their "best possible life" or close to it, compared with just 1 in 5 younger adults.

The study disproves many common misconceptions about aging and suggests that greater focus should be put on adults in their 40s and 50s, who reported higher levels of stress and worry and lower levels of life satisfaction and health than older Americans.

To view the study, click on the red button below.
New Additions to the engAGED Innovations Hub

The engAGED Social Engagement Innovations Hub houses best and emerging social engagement programs. Searchable by a variety of filters—including social engagement intervention type, geographic area served or partners involved—the Innovations Hub gives you access to replicable examples to inspire and inform your organization’s social engagement efforts. Each program summary within the Innovations Hub contains all you need to know about that particular program, such as partners involved, outcomes demonstrated, lessons learned, resources needed and contact information to learn more.

To access the Hub, click the red button below.

ACCESS THE HUB

Improving Representation in Clinical Trials and Research: Building Research Equity for Women and Underrepresented Groups

The United States has long made substantial investments in clinical research with the goal
of improving the health and well-being of our nation. There is no doubt that these investments have contributed significantly to treating and preventing disease and extending human life. Nevertheless, clinical research faces a critical shortcoming. Currently, large swaths of the U.S. population, and those that often face the greatest health challenges, are less able to benefit from these discoveries because they are not adequately represented in clinical research studies. While progress has been made with representation of white women in clinical trials and clinical research, there has been little progress in the last three decades to increase participation of racial and ethnic minority population groups. This underrepresentation is compounding health disparities, with serious consequences for underrepresented groups and for the nation.

At the request of Congress, *Improving Representation in Clinical Trials and Research: Building Research Equity for Women and Underrepresented Groups*, identifies policies, procedures, programs, or projects aimed at increasing the inclusion of these groups in clinical research and the specific strategies used by those conducting clinical trials and clinical and translational research to improve diversity and inclusion. This report models the potential economic benefits of full inclusion of men, women, and racial and ethnic groups in clinical research and highlights new programs and interventions in medical centers and other clinical settings designed to increase participation.

To learn more or to access the report, click the red button below.