Family caregivers are the most important source of support for people with chronic or other health conditions, disabilities, or functional limitations. Millions of family caregivers help their loved ones live at home and in their communities, providing the bulk of this assistance. About 40 million family caregivers provide about $470 billion annually in unpaid care to their loved ones.\(^1\) According to Caregiving in the US 2015, about 3.7 million family caregivers provide care to a child under age 18 because of a medical, behavioral, or other condition or disability and 6.5 million family caregivers assist both adults and children.\(^2\) Estimates show 3.5 million individuals with intellectual or developmental disabilities (I/DD) live with family caregivers, of whom over 850,000 are age 60+.\(^3\)

**What do family caregivers do?**

Family caregivers help with activities such as eating, bathing, dressing, transportation, and managing finances; perform medical/nursing tasks such as wound care and managing multiple, complex medications; arrange and coordinate care among multiple providers and settings; and pay for services to help their loved ones, such as home modifications, transportation, or a home care aide. The assistance family caregivers provide saves taxpayer dollars by helping to delay or prevent more costly nursing home care and unnecessary hospital stays. Family caregivers spend on average 24 hours a week caring for their loved one; almost one-third of family caregivers provide an average of 62 hours of care a week.\(^4\)

**What challenges do family caregivers face?**

Family caregivers take on physical, emotional, and financial challenges. They commonly experience emotional strain and mental health problems, especially depression, and have poorer physical health than noncaregivers.\(^5\) Family caregivers generally do not receive training and other assistance to help them provide care.\(^6\) Too often family caregivers aren’t even recognized and included, as appropriate, by health care and social service providers helping their loved ones. Navigating, locating, and coordinating fragmented services is too often bewildering, complex, and very time consuming.

Most family caregivers are employed and juggle work and caregiving responsibilities. They often make workplace accommodations because of caregiving, up to and including leaving their jobs.\(^7\) Employers have an interest in supporting family caregiver employees so they do not lose talented workers. Family caregivers (age 50 and older) who leave the workforce to care for a parent lose, on average, nearly $304,000 in wages and benefits over their lifetime. These estimates range from $283,716 for men to $324,044 for women.\(^8\) In addition, family caregivers may pay out-of-pocket for services whose costs add up over time.

Family caregivers will only face greater strains in the future as the “caregiver support ratio” – the number of potential family caregivers aged 45-64 for each person aged 80+ - shrinks. In 2010, the ratio was more than seven potential caregivers for every person in the high-risk years of 80-plus. By 2030, this ratio is projected to decline sharply to 4 to 1 and to less than 3 to 1 in 2050.\(^9\)
Why is a strategy to support family caregivers important?
If family caregivers were no longer available, the economic cost to the U.S. health care and long-term services and supports (LTSS) systems would increase astronomically. Our country relies on the contributions family caregivers make and should recognize and support them. Supporting family caregivers helps the caregivers themselves, the millions of individuals who rely on them, and also the economy and the workplaces who benefit from the contributions of family caregivers. Congress has passed and the President has signed into law the Recognize, Assist, Include, Support, and Engage (RAISE) Family Caregivers Act (P.L. 115-119). This legislation was sponsored by Senators Susan Collins (R-ME) and Tammy Baldwin (D-WI) and Representatives Gregg Harper (R-MS) and Kathy Castor (D-FL). The law will implement the bipartisan recommendation of the federal Commission on Long-Term Care that Congress require the development of a strategy to support family caregivers, similar in scope to the national strategy developed to address Alzheimer’s disease. The law is also consistent with a 2016 recommendation by the National Academies of Sciences, Engineering, and Medicine calling for a family caregiver strategy. Over 80 national organizations supported the legislation.

What does the RAISE Family Caregivers Act do?
It requires the development, maintenance, and updating of a strategy to recognize and support family caregivers by the HHS Secretary. A family caregiver is an adult family member or other individual who has a significant relationship with, and who provides a broad range of assistance to, an individual with a chronic or other health condition, disability, or functional limitation. The law brings together representatives of the private and public sectors, such as family caregivers, older adults with LTSS needs and persons with disabilities, veterans, health care and LTSS providers, employers, state and local officials, and others on an advisory council to advise and make recommendations regarding the strategy. The advisory council meetings will be open to the public and there will be opportunities for public input. The strategy will identify recommended actions that communities, providers, government, and others are taking or may take to recognize and support family caregivers, including with respect to:

- Promoting greater adoption of person-and family-centered care in all health and LTSS settings, with the person and the family caregiver (as appropriate) at the center of care teams;
- Assessment and service planning (including care transitions and coordination) involving care recipients and family caregivers;
- Information, education, training supports, referral, and care coordination;
- Respite options;
- Financial security and workplace issues.

There are 18 months for the development of the initial strategy under the law, followed by updates of the strategy biennially, and a sunset of the law three years after enactment. The law will improve the collection and sharing of information, including related to evidence-based or promising practices and innovative models regarding family caregiving, as well as better coordinate, assess, maximize the effectiveness, and avoid unnecessary duplication of existing federal government programs and activities to recognize and support family caregivers. The strategy and work around it could help support and inform state and local efforts to support family caregivers.

January 2018

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