Reducing Burden on Hispanic Caregivers of Relatives with Dementia – The Northern Manhattan Hispanic Caregiver Intervention Effectiveness Study (NHiCE)

What was the research about?
Dementia is a loss in mental ability that interferes with daily life. People with dementia often lose the ability to care for themselves. About one in eight people over 65 have dementia. Unpaid caregivers, such as family members, often take care of people with dementia. Caring for a person with dementia can be hard and, at times, overwhelming.

Dementia is more common among Hispanic people than other groups. Hispanic caregivers also have higher rates of burden, or personal strain, and depression than other caregivers.

The study compared two programs designed to help caregivers:

- The New York University Caregiver Intervention, or NYUCI
- The Resources for Enhancing Alzheimer’s Caregiver Health: Offering Useful Treatments, or REACH OUT

The research team compared how well the two programs helped Hispanic caregivers by

- Reducing the burden that caregivers feel
- Reducing signs of depression among caregivers

What were the results?
After six months, the two programs worked about the same to help caregivers. The caregivers in both programs had less burden and stress than they did at the start of the study. Caregivers’ signs of depression and physical function didn’t change during the study.

In both programs, the reductions in burden and stress were larger for caregivers who were caring for their spouse than for those who were caring for their parent.

Who was in the study?
The study included 221 Hispanic caregivers who lived in New York City and took care of a relative with dementia. About 63 percent of caregivers were caring for their parent, and 32 percent were caring for their spouse. The average age of caregivers was 58, and 83 percent were women.
What did the research team do?
The research team assigned caregivers to one of the two programs by chance. Both programs offered six sessions over the course of six months. Each session ranged from 60 to 90 minutes. Social workers, who spoke both English and Spanish, led both programs.

Caregivers in the NYUCI group received six counseling sessions in their home or at another convenient place. Four of the sessions were family sessions. The caregiver could choose what family members or close friends to include. The caregiver, along with the family when present, decided on the focus for each session.

Caregivers in the REACH OUT group received six in-home sessions. The sessions focused on skill building and problem solving. The caregivers created an action plan to address their problems. They also received information about dementia, self-care, safety, and other caregiver issues.

Caregivers took surveys before starting the programs and again six months after the programs ended. The surveys asked about burden, depression, stress, and physical function. The research team looked for changes in the caregivers’ answers to the survey questions.

What were the limits of the study?
The study only included family caregivers in New York City. The results may be different in other places. Also, the study may have been too short to see changes in signs of depression or physical function.

Future research could test the programs in different places or for a longer time.

How can people use the results?
Both REACH OUT and NYUCI reduced feelings of burden and stress in Hispanic caregivers of relatives with dementia. Organizations that serve Hispanic caregivers could consider offering such programs.

To learn more about this project, visit pcori.org/Luchsinger068.