The Caregiver Role for a Family Member with Dementia
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Background
- In the U.S., dementia affects an estimated 5.7 million Americans, and this number is expected to rise to nearly 14 million by 2050 (Alzheimer’s Association, 2018).
- Approximately 70-80% of people with dementia reside with a caregiver (Roberts & Struckmeyer, 2018).
- More than 16.1 million Americans currently care for individuals with dementia without pay, most of this care being provided by their loved ones (Alzheimer’s Association, 2018).
- Caring for a loved one with dementia is associated with emotional, physical, and financial repercussions (Warchol-Biedermann et al., 2014).

Purpose
- The objectives of this mixed method study was to assess the physical, emotional, and psychological effects of being a caregiver for someone with dementia and the impact of using an adult day service.

Theoretical Framework
- Caregiver resilience is the successful use of coping strategies that emphasize the perspective of resilience rather than burden (Roberts & Struckmeyer, 2018).
- Resilience enhances the ability to maintain normal or enhanced functioning during stressful caregiving experiences.
- Fitting with the conceptualization of resilience, caregivers experience subtle but long-term role changes, both physical and emotional.
- Caregiver respite is one factor that may sustain resilience over the continuum of care for family members caring for an individual with dementia.

Method

Approval: University IRB
Setting: Private, faith-based, adult day service center
Design: Cross-sectional, descriptive
Sample: Caregivers of those with dementia
N = 10
Qualitative: Two Focus groups: Structured, open-ended questions
Quantitative: The Kingston Caregiver Stress Scale (KCSS) (Hopkins & Klik, 2016)
Analysis: Interviews audio recorded, transcribed verbatim & descriptive statistics

Results

Caregiver Characteristics

| Age: \( M = 65.0 \text{ SD} = 10.58 \) |
| Gender: 60% male, 40% female |
| Race: 90% White, 10% Hispanic |
| Employment: 50% employed outside the home |
| Years of Providing Care: 6.1, SD = 6.0 |
| Years of Using Day Service: 1.8, SD = 0.98 |

Caregiver’s Relation to Client

Reasons for Using Services

Kingston Caregiver Stress Scale
- Caregiver stress: 3 Domains
- Care giving, family, & financial issues
- Scores can range 10-50; higher scores = more stress
- KCSS results: Ranged = 15 – 38; \( M = 30.5, SD = 9.07 \)

Themes

Most Common Emotional Responses: The participants described a variety of emotional responses experienced while providing care for their loved one. These emotions included feelings of grief, guilt, denial, frustration, and despair.

“Because I think the hardest thing I had to deal with was it’s not going to get any better. There is no cure. I have to accept that….it's only going to get worse.”
“Id be crying every day. It's mourning. There's grief. It's a grief, and it's continuous.”
“I think sometimes you almost feel guilty when you wish that it was over.”

Benefits of Adult Day Service: Benefits of Adult Day Services, the participants were grateful for the security and peace of mind provided because it benefitted their lives and helped promote resilience.

“I was able to go out and at least run, go to the YMCA….. Because as you know, there’s a lot of stress. And the physical activity was what saved me.”
“So, they’re a godsend, because that’s a break for me. I try to work during those hours.”
“And, it’s the best thing I every did…” “It saved me.”

Overwhelming Impact on Daily Life: Participants expressed concerns with role overload that changed the way they lived their daily lives. Not only did the caregiver role impact them, but the role impacted family members as well.

“Another kids, sometimes I feel bad that it’s compromising kind of their — I don’t know social lives.”
“You have to take over everything.”
“There’s very little time in my life.”
“Grocery shopping, making food, sleeping in a chair next to his bed. I had to know, learn everything…”

Caregiver Social Isolation: Participants expressed concerns of feeling disengaged from their lives, friends, and social connections. Some even expressed feelings of being a burden to others.

“You have to cut back on things you love. You can’t do the hobbies you used to do. You can’t do the, you know, meet the friend for lunch on any given day. You can’t do those simple things that used to bring you a little bit of joy.”
“You don’t want to ruin somebody else’s time, because usually that topic comes up.”

Conclusions/Implications for Practice

- Respite services, such as adult day centers, provide caregivers opportunities for self-care, the ability to perform daily tasks, and result in decreased emotional burden.
- Qualitative and quantitative data reflected stressful aspects of the caregiver role.
- Using respite services promotes resiliency in the caregivers and supports a healthier mental, emotional, social, and physical well-being.
- Education is needed to inform the public of the resources available when caring for someone with dementia.

Future Research

- Investigate the barriers caregivers face when learning about an adult day center.
- Further assessment of caregivers’ experiences as their loved one’s disease continues to progress.
- Explore the impact of adult day services on the person with dementia.

Acknowledgements: Funding from The Kreft Endowment