Perceived support from a caregiver's social ties predicts subsequent care-recipient health

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ABSTRACT

Most social support research has examined support from an individual patient perspective and does not model the broader social context of support felt by caregivers. Understanding how social support networks may complement healthcare services is critical, considering the aging population, as social support networks may be a valuable resource to offset some of the demands placed on the healthcare system. We sought to identify how caregivers’ perceived organizational and interpersonal support from their social support network influences care-recipient health.

We created a dyadic dataset of care-recipient and caregivers from the first two rounds of the National Health and Aging Trends survey (2011, 2012) and the first round of the associated National Study of Caregivers survey (2011). Using structural equation modeling, we explored how caregivers’ perceived social support is associated with caregiver confidence to provide care, and is associated with care-recipient health outcomes at two time points. All data were analyzed in 2016.

Social engagement with members from caregivers’ social support networks was positively associated with caregiver confidence, and social engagement and confidence were positively associated with care-recipient health at time 1. Social engagement positively predicted patient health at time 2 controlling for time 1. Conversely, use of organizational support negatively predicted care-recipient health at time 2.

Care-recipients experience better health outcomes when caregivers are able to be more engaged with members of their social support network.
health outcomes associated with care provision. However, understanding scenarios in which caregiver social support may also be protective could be beneficial as social support is associated with caregiver and care-recipient outcomes (Pakenham et al., 2007; Thorpe et al., 2015; Weaver et al., 2011; Gere et al., 2014). Recent studies have looked at various factors in the context of care-recipient/caregiver dyads (Thorpe et al., 2015; Weaver et al., 2011; Gere et al., 2014). These studies have shown that behavioral concordance, such as smoking behavior (Weaver et al., 2011), are associated with care-recipient and caregiver psychological and emotional health. These studies have provided compelling evidence for the need to understand disease outcomes in a broader context.

In the following analysis, we sought to identify how various organizational (e.g. social workers) and interpersonal (e.g. family) sources within a caregivers’ social support network influence care-recipient health. We expect that the extent to which caregivers perceive themselves to be supported as they provide should affect their confidence in their ability to provide care and have an effect on the health of care-recipient as an indirect consequence that underscores the importance of investing in social support systems not only for care-recipient, but also for those who work with them. To test this argument, we constructed a unique dataset using existing data to look at the predictive power of caregiver’s perceived social support on the health outcomes of their care-recipients.

1. Design and methods

We obtained care-recipient data from the first (2011) and second (2012) waves of the National Health and Aging Trends Study (NHATS) (Kasper and Freedman, 2014). The NHATS is nationally-representative of Medicare beneficiaries age 65 and older and includes a variety of constructs relevant to disease type, management, and consequences collected through in-person interviews conducted by trained research personnel. Although NHATS allows for proxy respondents when participants are unable to respond on their own, for consistency across the sample we focus here on those who provided their own data as our outcome is self-reported health status. Further, of self-reporting participants, those reporting Alzheimer’s or Dementia (< 5%) were excluded as the experiential and care-provision burdens associated with such cognitive impairment are fundamentally different from other morbidities measured in the NHATS (e.g. diabetes, arthritis, high blood pressure). Caregiver data was obtained from the associated National Study of Caregiving (NSOC), which was collected in 2011 (Kasper et al., 2014). NSOC data were collected via telephone-interviews with trained research personnel. Caregivers were eligible if 1) identified as a caregiver by an NHATS participant, and 2) indicated that they provided unpaid care to the nominating participant Measures included constructs such as the consequences of caregiving, social support, and caregiver health.

In order to access sensitive NHATS data (e.g. participant age) and NSOC data, we obtained approval to access sensitive data from the NHATS Data Confidentiality Committee. We downloaded and merged public and sensitive datasets for wave 1 (N = 8245), wave 2 (N = 7075) NHATS participants, and respective caregiver data from NSOC (N = 2007). The current study focuses on a subset of patients who responded to both waves 1 and 2), lived in traditional community residences, and identified at least one informal caregiver. Because patients could have up to five caregivers, caregiver data was averaged to avoid over-representing care-recipient with multiple caregivers. We also created an additional variable to capture the number of caregivers for any one particular patient. Our analysis consists of N = 780 care-recipient and their associated caregivers (N = 1412). It is important to note that while the NHATS and NSOCs datasets are weighted, we do not use sample weights as the dyads in this analysis are distinguishable (i.e. non-independent). Considering the purpose of data weights is to define the total population, it is unclear how to define the population of care-recipients and caregivers, especially when a care-recipient has multiple caregivers, and covariation is a function of common influences (e.g. geography) and within-dyad relational aspects that affect thoughts, feelings, and behaviors. Thus, the use of such weights would be mis-leading and not contribute to the overall goal of understanding the relationships between these variables in order to better understand how to bolster caregiving efforts.

Multiple items comprised our key predictor variables. Four computed variables represented four domains within the broader context of social support, and a single item represented the mediating variable of caregiver confidence.

Aide finding support was computed from six-items which asked caregivers to identify who provided aide in caregivers’ search for support. These sources included government/community agency, medical care provider or social worker, church/synagogue, employer, self-guided or from a friend, and other sources of support. Caregivers responded yes or no to each of the six sources. Responses to the six sources were summed to capture the total number of sources who aided caregivers’ search to find support.

Use of organized supportive services was computed by summing the number of organizational supports (respite care, training, and financial help; three items) caregivers indicated having used in the past year.

Social engagement was computed from five items which asked caregivers to identify whether or not in the past month they attended religious services, went to a support group, visited with friends or family not living in the same home, participated in club meetings or group activities, or went out for enjoyment.

Instrumental support was computed from two items which asked if caregivers received help from friends or family with their own daily activities, such as running errands, and if friends or family helped the caregiver provide care to the care-recipient.

Caregiver confidence was measured with one item, “helping [care-recipient] has made you more confident about your abilities,” measured on a scale of 1 = very much to 3 = not so much.

Care-recipient self-reported health at wave one and wave two were the primary outcome variables of interest. Self-reported health was measured with the item, “would you say in general your health is…” with 1 = very good and 5 = poor.

We used structural equation modeling (SEM) to assess how caregivers’ perceived organizational and interpersonal supports predict care-recipient health status as reported in the first and second waves of the NHATS survey. We controlled for the severity of care-recipient illness by accounting for the number of chronic illnesses reported by the care-recipient by creating a unique variable to represent the sum of chronic conditions reported. Additional patient demographic characteristics (i.e. patient age, gender, and educational attainment) found to be significantly correlated with the primary outcomes of interest (care-recipient self-reported health in 2011 and 2012) were controlled, as well as caregiver characteristics (i.e. caregiver health status, educational attainment, and marital status) associated with caregiver confidence in ability to provide care (hypothesized to mediate the relationship between caregiver network support and care-recipient health). To select predictors to include in the SEM, we conducted a series of preliminary correlational and multiple regression analyses to exclude non-significant predictors following the method outlined by Bugge et al. (1999). Caregiver age and duration of caregiving, for example, were not associated with caregiver confidence and were thus excluded from the model to enable clearer interpretation of the relationship between various aspects of social support and caregiver confidence and care-recipient self-reported health. Intensity of caregiving was removed as it is highly correlated with numerous care-recipient factors such as health status and number of chronic diseases, thus introducing multicollinearity that would confound the SEM results. We used SEM with maximum likelihood estimation in STATA version 13. To assess model fit, multiple indicators were consulted: chi-square,
Table 1 Demographic characteristics of care-recipient (NHATS 2011, 2012) and caregivers (NSOC 2011).

| Variable | Care-recipient \((N = 780)\) | Caregiver \((N = 1412)\) |
|----------|-------------------------------|--------------------------|
| Age at first interview mean (SD)/range | 80 (8.21) 65–101 | 56 (13.08) 18–92 |
| Chronic disease | | |
| Heart attack | 159 20 | 98 7 |
| Heart disease | 199 26 | 102 7 |
| High blood pressure | 582 75 | 639 45 |
| Arthritis | 560 72 | 545 39 |
| Osteoporosis | 221 28 | 168 12 |
| Diabetes | 276 35 | 233 17 |
| Lung disease | 166 21 | 192 14 |
| Cancer | 223 29 | 147 10 |
| Gender | | |
| Female | 540 69 | 916 65 |
| Education | | |
| Less than high school | 278 36 | 125 9 |
| High school diploma/GED | 268 34 | 340 24 |
| Some college/associate's degree | 130 17 | 291 21 |
| Bachelor's degree or higher | 103 13 | 302 21 |
| Health status (wave 1) | | |
| Excellent | 122 16 | 266 19 |
| Very good | 238 30 | 430 30 |
| Good | 256 33 | 397 28 |
| Fair | 131 17 | 240 17 |
| Poor | 33 4 | 59 4 |
| Don't know/refused | 0 0 | 20 1 |
| Health status (wave 2) | | |
| Excellent | 112 14 | | |
| Very good | 252 32 | | |
| Good | 255 33 | | |
| Fair | 126 16 | | |
| Poor | 35 4 | | |
| Care-recipient/caregiver relationship | | |
| Daughter | 472 33 | | |
| Spouse/partner | 335 24 | | |
| Son | 211 15 | | |
| Grandchild | 92 7 | | |
| Child-in-law | 77 5 | | |
| Friend | 70 5 | | |
| Sibling | 46 3 | | |
| Niece/nephew | 36 3 | | |
| Other relative | 44 3 | | |
| Other nonrelative | 29 2 | | |

* Care-recipient and caregivers were allowed to select more than one chronic health condition.
* One care-recipient and 354 caregivers did not provide educational attainment data.

3. Discussion

We conclude that care-recipients experience better health when caregivers perceive more support from and engage more often with their own interpersonal connections. Our hypothesis was partially supported as the effect of social engagement and instrumental support on care-recipient self-reported health status was mediated through caregiver confidence in their ability to provide care. However, it is important to note that the paths associated with the use of organizational support and instrumental support were negative predictors. This is likely explained by the fact that care-recipient who are in worse condition are likely to be eligible for additional organizational resources and intensive medical care. Instrumental support was negatively associated with caregiver confidence, but the net effect on care-recipient health was positive. Thus, this finding is encouraging as it may be that caregiver confidence decreased decreased due to the severity of the care-recipient’s condition. However, social engagement was positively associated instrumental support, again highlighting that care-recipient of caregivers with stronger social networks experience better health outcomes.

A unique finding that we did not predict was that social engagement was not only mediated by caregiver confidence in their ability to provide care, but also had a direct positive association with care-recipient health at wave one, and wave two controlling for wave one. These findings highlight the importance of investing in social programs that enable caregivers to maintain interpersonal relationships and the ability to partake in activities they enjoy. Determining how to effectively encourage and support caregiver social engagement is an important next step for intervention research. We know from previous research that care-recipient-perceived social support is associated with lower care-recipient morbidity and mortality (Uchino et al., 1996), and that caregiver factors, such as functional limitations, impact care-recipient outcomes (Thorpe et al., 2015). This study builds upon existing research by demonstrating that acknowledging the role of the caregiver social support network provides a broader narrative as to how support environments influence care-recipient health and suggests the potential importance of investing in social support structures for caregivers as well as for their care-recipients.

Given the nature of secondary data analysis, important limitations must be considered. Inherent to any secondary analysis, the data were not collected for the exact purpose of the current study. Validated measures often used in the literature to measure perceived social support were not included in the NHATS or NSOC surveys, therefore, we constructed composite variables to understand the relationships between constructs. Caregiver confidence and care-recipient self-reported health were represented by single items, as this is what was available in the dataset, representing a limitation to this analysis. Thus, additional research should employ validated measures to better understand the complexities of this larger social context. However, the strengths of this analysis outweigh the limitations as these datasets offered a unique opportunity to begin exploring the effects of caregivers’ social networks on care-recipient outcomes over time, which is currently an aspect of the caregiver-patient relationship that is absent in the literature.

By 2029, when the last of the baby boomer generation has retired, over 20% of the U.S. population will be over the age of 65 (Colby and Ortman, 2014). As the age of the U.S. population shifts, informal caregiving will likely become an imperative compliment to organizational healthcare services. Thus, understanding how to best support informal caregivers is critical to the health and wellbeing of caregivers and care-
recipients. Future research designed to answer questions specific to this context is greatly needed.

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Fig. 1. Impact of Caregivers' (N = 1412) Perceived Social Support on Longitudinal Care-recipient (N = 780) Health.
Note. Data for care-recipients is from the first (2011) and second (2012) waves of NHATS and data for caregivers was collected from the first (2011) wave of NSOC as the second NSOC dataset was not available at the time of analysis (2016). *p < 0.05; **p < 0.01; ***p < 0.001.