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Sociodemographic disparities in adult-child informal caregiving intensity in the US:
Results from the new National Study of Caregiving

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Abstract: Nearly 60 million informal caregivers provide care to aging adults. Despite its many benefits, high intensity caregiving can impact caregiver health and quality-of-life. Therefore, the objective of our study was to assess socioeconomic and demographic disparities in caregiving intensity among informal caregivers. Using a randomized, nationally representative database of 1014 informal offspring caregivers from Medicare enrollment databases, the associations between informal caregiving intensity and age, race/ethnicity, and income were examined using binary and ordinal logistic regression. Caregiving intensity varied by demographics. High ADL caregiving was highest among Black, non-Hispanic caregivers. High IADL caregiving and high number of hours spent caregiving was highest in females, and non-White caregivers. Although the overall association between caregiving intensity and income was not significant, when stratified by race/ethnicity, this association was positive for White caregivers and negative for non-White caregivers. Health care providers frequently interact with informal caregivers and should be aware of trends in caregiving and the needs and supports available to ameliorate caregiver burden. In order to protect caregivers, policies and programs designed to promote wellbeing and mitigate the potential harms of caregiving to health should consider these findings.
Introduction:

Over 34 million American adults have provided informal care to an adult over the age of 50 within the last 12 months, 47% of whom were caring for a parent (National Alliance for Caregiving, 2015). These figures are only expected to rise as the number of Americans over 65 years of age grows from 47.5 million in 2015 to 98 million by 2060 (Administration on Aging, 2016). Informal caregiving, the unpaid care and support family members and friends voluntarily provide to individuals who are unable to function independently, has numerous benefits to care recipients and society as a whole. These include savings to the national economy, prevention of hospitalization and institutionalization, and allowing older residents to remain in their own homes (Chari, Engberg, Ray, & Mehrotra, 2015). Informal caregivers are estimated to have spent over 30 billion hours a year providing care to disabled or chronically ill individuals, with an opportunity cost savings of $522 billion per year that would otherwise be spent on formal care and institutionalization (Chari, Engberg, Ray, & Mehrotra, 2015).

Despite these benefits of informal caregiving to the care recipient and to the national economy, numerous negative effects associated with caregiving have been well documented in the literature. Many studies have demonstrated the negative impacts on caregiver health-related quality of life, including physical and emotional health consequences, that can occur as a result of providing care, such as anxiety and depression (Cannuscio, et al., 2002; Ho, Chan, Woo, Chong, & Sham, 2009; Macneil et al., 2010; National Alliance for Caregiving, 2015; Pinquart & Sorensen, 2011; Schultz & Sherwood, 2008). These negative health-related quality of life outcomes are commonly referred to as caregiver stress, strain, or burden. Moreover, informal caregivers providing a high amount of care may be particularly vulnerable to the effects of caregiver strain, and may differ in substantial ways from those providing less care, such as their employment status, the type of caregiving duties they provide, and the impact caregiving has on them (Jacobs, Laporte, Van Houtven, & Coyte, 2014; National Alliance for Caregiving, 2015). The duties caregivers provide
may vary by sociodemographic factors, including race and gender, where female and non-white caregivers were more likely to provide higher amounts of informal care than their counterparts (National Alliance for Caregiving, 2015). However findings are mixed. Few studies have examined differences in caregiving intensity by socioeconomic and demographic factors using a nationally representative dataset.

There is a critical need to better understand who is providing informal care to aging parents, and how to best provide the support, assistance, and resources caregivers may need. To further the understanding of who is providing informal care, the objectives of this study were to examine the demographic profile of informal adult-child caregivers in the U.S., and to assess the sociodemographic differences in caregiving duties (caregiving “intensity”) among this population of informal caregivers.

**Methods:**

**Study population**

The data were obtained from the 2011 National Study of Caregiving (NSOC) dataset, a nationally representative sample of informal caregivers. The NSOC identified caregivers of National Health and Aging Trends Study (NHATS) participants who were receiving assistance in self-care, mobility, medical, or household activities. These caregivers were then contacted to participate in a one-time, cross-sectional assessment of caregiving that included questions on caregiving activities, duration, intensity, and demographics. This analysis focused on adult children caregivers to older adult parents (n = 1014), a subset of informal caregivers.

**Outcome variables: Caregiving intensity**

Four individual measures of caregiving duties that are most common in the literature were used to assess caregiving intensity: 1) Number of Activities of Daily Living (ADLs) performed, 2) Number of Instrumental Activities of Daily Living (IADLs) performed, 3) Hours of caregiving
provided per month, and 4) Duration (years) of caregiving. To measure these four intensity
domains, items from the NSOC questionnaire assessing aspects of caregiver duties were used. ADLs
refer to daily self-care activities that are necessary for fundamental functioning. This was measured
by the number of personal care activities caregivers helped with each month, including eating,
bathing, dressing, toileting, and helping care recipient move around. IADLs consist of other
caregiving activities not necessary for fundamental functioning, but allow an individual to live
independently. This domain included the number of instrumental activities caregivers helped their
parent with, including medication management, scheduling medical appointments, and other health
and hygiene-related tasks. The last two intensity domains were calculated based on the average
number of hours spent caregiving in the last month, and average number of years providing care.
The top quartile (25%) of each individual intensity domain were considered ‘High Intensity
Caregivers’, while the bottom 75% were considered ‘Low Intensity Caregivers’.

To calculate the composite intensity measure, each of the four individual measures of
caregiving intensity scored one point if considered 'high intensity', and zero points if considered
“low intensity”. This composite score ranged from zero (provided no high intensity care in any of
the four individual caregiving measures) to four (provided high intensity care in all four individual
caregiver domains).

Exposure variables: Caregiver demographics

Four demographic characteristics of caregiver respondents identified in previous studies of
caregiving intensity were assessed, to include caregiver age, gender, race/ethnicity, and annual
household income (Navaie-Waliser, Spriggs, & Feldman, 2002; Navaie-Waliser, et al, 2002;
Demographic age was categorized into 10-year age groups (< 45, 45-54, 55-64, and 65+).
Race/ethnicity was based on three calculated domains (non-Hispanic White, non-Hispanic Black,
and ‘Other’ (Hispanic, American Indian, Native Hawaiian, Pacific Island, other non-Hispanic)).
Income was grouped into four $25,000 intervals ($\leq 24,999$, $25,000$-$49,999$, $50,000$-$74,999$, and $75,000$ or more).

**Additional demographic confounders and covariates**

Other key confounders and covariates commonly used in studies of disparities in caregiver intensity and are shown to affect caregiver intensity, including caregiver marital status (Brody, Litvin, Hoffman, & Kleban, 1995; Dentinger & Clarkberg, 2002), presence of a child (under 18) living in the home (Cohen, Cook, Sando, Brown, & Longo, 2017; Grundy & Henretta, 2006), and caregiver co-resident status (care recipient and caregiver reside in same home) (Tennstedt, Crawford, & McKinlay, 1993) were also assessed.

**Data analysis**

Univariate and bivariate analyses were used for all primary outcome and exposure variables to assess individual measures of high intensity caregiving. For the composite measure of high intensity caregiving, ordinal logistic regression models were used to calculate adjusted odds ratios and 95% confidence intervals (CI), adjusting for covariates. Pairwise deletion was used to handle missing values for each model. SAS 9.3 (Cary, NC) was used for all analyses.

**Results:**

**Demographics of NSOC adult-child caregivers**

The demographic breakdown for this sample of adult children caregivers is found in Table 1. The average age of caregivers was 54.6 years old. Sixty-nine percent of respondents were female, while 31% were male. Respondents reported an average annual income of $56,582. Sixty percent of respondents identified as non-Hispanic White, 31% as non-Hispanic Black, and 9% as another racial/ethnic group. Adult children caregivers spent an average of 85 hours a month providing care, and had been caring for an elderly parent for 5.6 years.

**Individual measures of Caregiving Intensity**
High ADL caregiving was most prevalent in caregivers aged 45-54 (28.1%) and non-Hispanic Black caregivers (33.9%). High IADL caregiving was significantly higher in females (30.3%) than in males (20.6%). Compared to White caregivers, high IADL caregiving was significantly higher among non-Hispanic Black caregivers (33.5%) and caregivers of ‘other’ racial/ethnic groups (30.9%). A high number of hours spent caregiving was highest in females (27.9%), non-Hispanic Black caregivers (36.1%) and ‘other’ race/ethnicities (30.8%), and individuals earning less than $25,000 per year (37.6%). High years of caregiving was highest in non-Hispanic Black caregivers (33.8%) and caregivers of ‘other’ race/ethnicities (35.7%). All results are displayed in Table 2.

Composite measure of caregiving intensity

Differences in high intensity caregiving varied by gender, race, and other sociodemographic factors (Figure 1). Female caregivers had higher odds of providing high intensity care than their male caregiver counterparts (OR = 1.43, CI [1.03, 1.99]). The odds of providing high intensity caregiving was greater for non-White caregivers (caregivers of Black and ‘other’ race/ethnicities) than White caregivers (OR = 1.86, CI [1.30, 2.64]). Co-resident caregivers were more likely to have provided high intensity caregiving than caregivers not residing with their care recipient (OR = 1.70, CI [1.19, 2.42]). Additionally, a negative association between annual income and high intensity caregiving was observed: as caregivers’ annual income decreased, the odds of providing high intensity caregiving significantly increased (p < 0.001).

Discussion:

This study sought to better understand who is providing high intensity informal care to an older parent. Our findings indicate that there are notable differences in caregiving intensity that vary by caregiver socioeconomic and demographic factors. High intensity caregiving was most prevalent among females, non-White caregivers, those living with their care recipient, and low
income caregivers. These results are in line with previous studies that report greater caregiving responsibilities among females, racial/ethnic minorities, shared co-residence, and low income caregivers (Kim, Chang, Rose & Kim, 2012; Pinquart & Sörensen, 2005; National Alliance for Caregiving, 2015). However, the observed associations were dependent on the type of care being provided, such that certain caregivers were more likely to provide high intensity care in some domains of caregiving than others. Surprisingly, no differences were observed between males and females in high ADL caregiving. Historically, this has not been the case, and could be a result of the changing face of informal caregivers, or in how the caregivers were selected for inclusion in this sample.

While our results largely confirm previous findings, our analysis adds to the current body of caregiver research in several ways. First, the analysis uses a nationally representative sample of caregivers previously identified by their care recipient. This is an important distinction from other representative samples where respondents self-identify as caregivers. Second, our focus on adult child caregivers sheds light on who is providing high intensity care to an aging parent. Investigating adult child caregivers – an important and large subset of caregivers – is important, as they differ from other types of caregivers (i.e. spousal) in significant ways. Examining this group separately is recommended, as significant differences in caregiver characteristics, needs, and burden have been noted (Chappell, Dujela & Smith, 2014; Pinquart & Sörenson, 2011). Third, this analysis compares caregivers based on the level of care they provided (high vs. low intensity).

Caregiving intensity, whether measured by the type or amount of assistance provided is associated with various health effects and quality of life outcomes (Shultz & Sherwood, 2008). Numerous studies have shown any type of informal caregiving can result in negative physical and emotional health consequences for the caregiver, often referred to as caregiving-related stress or burden (Cannuscio, et al., 2002; Ho, Chan, Woo, Chong, & Sham, 2009; Macneil et al., 2010; National Alliance for Caregiving, 2015; Pinquart & Sörensen, 2011; Schultz & Sherwood, 2008). With noted
differences in the intensity of care being provided, we anticipate implications for caregiver health and quality of life that also vary by sociodemographics. Research suggests that differences exist in caregiver quality of life among male and female caregivers, caregivers of different racial and ethnic groups, ages (Neugaard, Andresen, McKune & Jamoom, 2008; Anderson et al., 2013; Covinsky, 2003), and income levels (Williams, Forbes, Mitchell, Essar & Corbett, 2003).

Health care professionals should be aware of trends in high intensity caregiving among informal caregivers providing high intensity care to an older parent. Gerontological nurses may have frequent contact with aging adults and their informal caregivers who are susceptible to or exhibiting signs of burnout and quality of life concerns. Their role in facilitative informal caregiving has expanded from being primary caregivers to teaching and assisting family members to provide care (Schulz & Sherwood, 2008). Nurses that provide informal caregivers a temporary break from their caregiving responsibilities can significantly improve caregiver health and quality of life (Lopez-Hartmann, Wens, Verhoeven, & Remmen, 2012) and reduce subsequent caregiver burden (Horton-Deutsch, Farran, Choi, & Fogg, 2002).

This group of health providers are especially poised to recognize symptoms of distress and burnout, and offer appropriate resources for caregivers in need of additional supports. Such support is facilitated through communication between nurses and informal caregivers, building relationships with informal caregivers, and creating a culture of trust with the family of the care recipient (Weman & Fagerberg, 2006). However, the availability of healthcare providers trained to provide such support may depend upon the services and resources that are available at the local level and may not be equitable for all informal caregiver populations throughout the US.

Limitations:

When interpreting these findings, there are a few important limitations to note. First, due to the cross-sectional nature of this study, we are unable to determine causal relationships between caregiver demographics and high intensity caregiving. A second wave of NSOC data will be available
within the year and future analyses may be able to determine causality. Second, we did not look at
caregiver employment, which may impact caregivers’ ability to provide different types and levels of
care. Third, caregiving responsibilities was dichotomized into high and low caregiving intensity,
rather than assessed along a gradient. As such, a composite analysis was conducted to assess
overall caregiving intensity as a continuous measure composed of multiple types of caregiving (e.g.
ADLs, IADLs, and hours per month). Next, all measures were self-reported, which may bias the
results toward more socially acceptable responses to the measures examined in the study. Sample
weights were not used in the analysis, as the importance of including weights in regression models
such as these in which descriptive population parameters are not being estimated are a subject of
debate in the survey analysis literature. Lastly, we assumed that missing data were missing at
random, and therefore did not impute missing values, and instead handled the issue of missing
values through the use of pairwise deletion.

Conclusions:

Our findings show both females and non-White caregivers are more likely to provide high
intensity care, though the intersection between these two constructs and the influence on outcomes
is not well known. What remains to be seen is the impact sociodemographic interactions have on
caregiver intensity and caregiver health outcomes. In addition, as the number of aging adults
increase and the face of caregiving evolves, we anticipate the sociodemographics of caregivers
providing high intensity care will also change over time, as well as their needs and necessary
supports. Consequently, policies and programs designed to promote caregiver health and quality of
life should consider these important sociodemographic disparities to protect and support this vital
component of the US health care system.


