Carer-Employee Burden

Carer burden refers to the perceived stress associated with the carer role, and is often associated with poor physical and mental health of their carer-employees. This burden has potential ramifications on their capacity to maintain both roles, leading to deterioration of health and job performance. For example, carers have a higher risk for developing depression, sleep disturbance, psychosocial issues, and increased stress levels when compared with non-carers. In addition, lack of carer programs in the workplace can result in an increase of absenteeism (missing work), presenteeism (distraction), and issues with retention and productivity. We have estimated previously that the economic consequence of care provision costs the specific workplace that is the focus of this study, approximately $8674 CAD per carer-employee. A more conservative estimate found that, on average, American employers may lose at least $2400 USD per employee due to lack of carer workplace programs that result in loss of productivity and absenteeism. Given that the number of carer-employees in Canada are projected to rise with the aging population, there will be a growing need for recognition and support of carer-employees from employers and policy-makers. One strategy to be considered by employers is the implementation of carer-friendly workplace programs (CFWPs) such as: flexible working hours, telecommuting, support services, paid or unpaid leave, financial assistance, and culture change.

This study seeks to explore the effects of implementing a CFWP on carer-employee’s health in a university workplace. The following section presents a literature review discussing: carer-employee burden, the relationship to physical and mental health in Canada, current state of CFWPs, and evidence for carer interventions. This section is followed by a presentation of our research objectives, before continuing with methodology, results, and finally, an argument for greater integration of CFWPs in the discussion and conclusion.

LITERATURE REVIEW

Carer-Employee Burden

Carer burden refers to the perceived stress associated with the carer role, and is often associated with poor physical and mental health, or age-related conditions. In Canada, over 8 million individuals aged 65 years or older globally, with this number expected to burgeon to 1.5 billion in 2050. Demographic trends in Canada, such as: the growing labor force participation of women, declining average household family size, and rising retirement age have led to increased pressure on the working population to provide informal care to the elderly population, as they increase in number. This research defines care responsibilities often include: personal/medical assistance, transportation, emotional support, management of finances and medical appointments, shopping, meal preparation, and housework. These unpaid services performed by carer-employees save the Canadian government $25 billion annually in health expenditures. It can be surmised then, that carer-employees produce significant and invaluable contributions to the healthcare sector on top of paid work obligations.

Demands of care and work have increased in recent years, due to the deinstitutionalization of formal care, together with the movement towards a connected global economy. This has led to a growing burden on carer-employees. This burden has potential ramifications on their capacity to maintain both roles, leading to deterioration of health and job performance. For example, carers have a higher risk for developing depression, sleep disturbance, psychosocial issues, and increased stress levels when compared with non-carers. In addition, lack of carer programs in the workplace can result in an increase of absenteeism (missing work), presenteeism (distraction), and issues with retention and productivity. We have estimated previously that the economic consequence of care provision costs the specific workplace that is the focus of this study, approximately $8674 CAD per carer-employee. A more conservative estimate found that, on average, American employers may lose at least $2400 USD per employee due to lack of carer workplace programs that result in loss of productivity and absenteeism. Given that the number of carer-employees in Canada are projected to rise with the aging population, there will be a growing need for recognition and support of carer-employees from employers and policy-makers. One strategy to be considered by employers is the implementation of carer-friendly workplace programs (CFWPs) such as: flexible working hours, telecommuting, support services, paid or unpaid leave, financial assistance, and culture change.

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LITERATURE REVIEW

Carer-Employee Burden

Carer burden refers to the perceived stress associated with the carer role, and is often associated with poor physical and mental health
health. Burden can be both objective, denoted by physical, financial, and social demands of care provision, as well as subjective, relating to the carer’s reaction towards performing care tasks. The majority (89%) of Canadian carers report caregiving for over 1 year. Generally, carer burden results in depressed mood, greater perceived difficulty in performing care work, anxiety, sleep disturbance, and social isolation. Carer burden is a recognized clinical condition, reached when carer burden results in excessive feelings of exhaustion from strain in the carer’s physical, psychosocial, financial, and emotional capacities. Most commonly, burden and eventually burnout, arises when paid work and personal obligations are incongruent with each other, and there is a lack of services and accommodations to assist in negotiating work-life balance.

The majority of carers are often women, with approximately 54% of Canadian carers identifying as women in 2012. Greater carer burden is experienced both acutely and chronically in female carers as compared with male carers, with female carers being significantly more likely to be responsible for tasks such as personal care, medical assistance, housework, emotional support, and meal preparation. These tasks are generally time sensitive and urgent, allowing for less flexibility in the carer’s schedule. Male carers are significantly more likely to provide physical assistance in house maintenance and repairs, which generally allows for greater flexibility to schedule such commitments outside of traditional working hours. This division of tasks is reflected in the time spent on care-related tasks; women are more likely than men to spend more than 20 hours per week on caregiving related activities than men, with 17.6% of female carers doing so as compared with 11.5% of male carers. Additionally, women average approximately 5.8 years of caregiving throughout their lives, while men average approximately 3.4 years. These duties result in women being more time-constrained and for longer periods of time, which impact their health as well as ability to perform in other areas of their lives.

Carer burden can include financial commitments such as out of pockets costs of medication and transportation, which may cost on average $3300 CAD annually. Given this, the financial burden of caring influences lower-income carer-employees significantly more than well-resourced carer-employees. Low-income carer-employees were found to have a higher probability of suffering from mental health issues, when compared with higher-income carer-employees. A high carer burden can manifest in other dimensions of carer-employee’s lives, such as quality of sleep and mental health, compromising their wellbeing and productivity at work.

Effects on Work and Health

Carer-employees may accrue a multitude of health and employment impacts due to the burden of managing their work-life balance. This is exacerbated by the often-chronic nature of caring. There is an observed positive correlation between reported carer burden and length of time spent as a carer, in which burden is greater the longer one has been in a carer role. Other adverse effects incurred due to carer-employee responsibilities include: poor morale, reduction in quality of interpersonal relationships, reduced social activities, and loss of one’s sense of independence. This results in poorer overall quality of life in carer-employees, leaving many carer-employees feeling isolated, frustrated, fatigued, and stressed. Approximately 38% of Canadians report overall health declining due to caregiving duties.

For many carers, carer burden and time constraints lead to diminished capacity to engage in paid work. Forty three percent (43%) of Canadian carers in 2012 reported missing partial or full days of work due to caring responsibilities, while 15% reduced their work hours to accommodate their work-life balance. Of those 43%, a further 14% lost employer provided health benefits, pension, and insurance as a direct result of reduced engagement in the workforce. From 2003 to 2008, net lost wages because of caregiving duties amounted to $336.8 million CAD annually, representing a direct loss of financial capital for carer-employees. We estimated that the annual economic cost absorbed by carer-employees at the research worksite of concern herein was approximately $32.922 per carer, due to loss of income, out of pocket caregiving expenses, and opportunity cost of caring labor.

Perceived burden is the result of a variety of dynamic factors, including: intensity of care, sex, care recipient condition, and carers’ financial status—all of which uniquely impact work-life balance. Understanding how each of these factors interact dynamically to form the carer-employee experience may shed light on how carer burden and health outcomes can be improved through CFWPs to produce healthier and productive work environments.

Workplace Accommodations

CFWPs are designed modifications to organizational structure, policies, or programs for the purpose of supporting employee work-life balance and reducing conflicts for carer-employees. Interest in implementation of CFWPs has been growing in recent years, with the finance, technology, and healthcare sectors leading in terms of present CFWP prevalence. Canadian businesses have been losing an approximate $1.3 billion worth of productive hours annually due to caregiving demands on employee time. Recognizing this, many workplaces are motivated to adopt various CFWPs to avert these costs as well as additional benefits such as increased employee satisfaction, and positive company reputation. The most common forms of CFWPs are currently flexible working arrangements, support services (eg, counselling), and paid/unpaid leaves. Within the Canadian context, little is known about the efficiency and uptake of these CFWPs in any form.

Carer Interventions

Interventions targeted towards carers generally fall into the following types: education, behavioral change, case study, support services, and psychosocial (eg, counseling, therapy). Carer interventions are developed with the aim of being therapeutic while building supportive capacity for carers. Common intervention protocols have intentionally sought to build: coping skills, knowledge of available resources, knowledge of how to best manage care recipient conditions, psychosocial health, carer burden, and planning skills.

Generally, carer interventions tend to produce modest but positive effects. This is largely theorized to be the result of the multidimensional and complicated nature of carer burden. Multicomponent interventions tend to show more pronounced intervention effects, likely due to a broader range of resources offered to carers. A 2002 meta-analysis of 78 carer interventions found two broad classes of interventions: (1) respite-type interventions aimed at reducing objective burden, and; (2) psychoeducational interventions targeting wellbeing. In addition, intervention effects related to building carer capacity and knowledge were found to have greatest short-term changes.

While workplace carer interventions have not been extensively explored, there is evidence that workplaces may be useful interventions sites for health factors by reducing work-family conflict. Modification to employee work environment, such as increased opportunity for job control and decision making at work or increased supports to decrease workload, has been associated with improved mental health and physical health in employees. This finding was reflected in the results of a meta-analysis of workplace interventions for healthcare workers, where work schedules conducive to work-life balance were associated with significant reductions in stress. Given the increasing elderly population and demands of eldercare, carer burden greatly contributes to
work-family conflict. Workplace interventions for carer-employees may offer a preventative solution to employee stress and burnout.

RESEARCH OBJECTIVES

This CFWP intervention pilot program is divided into three phases, with the overall goal of building the business case regarding the implementation of workplace programs for carer-employees. Approved by the university’s Research Ethics Board (ISRCTN 16187974), this study pertains specifically to the first phase (Phase A), in which a web-based decision tool was developed as an intervention for improving carer-employee health outcomes. This specific paper examines quantitative health outcomes at the conclusion of Phase A.

We aim to evaluate the effectiveness of an educational CFWP when implemented in the work environment over time. More specifically, this research asks the following question: does the implementation of CFWPs improve the health condition of carer-employees over time? The results will offer actionable evidence to policy decision makers on the benefits of CFWPs, while providing guidelines regarding the organizational changes necessary for the inclusion of carer-employees in healthy productive workplaces.

In keeping with the literature review, our CFWP intervention explores educational effects of a CFWP intervention on carer-employee’s health and wellbeing. The multicomponent intervention was designed to include both an educational and behavior change component, to maximize potential intervention effects. Additionally, our participants include carers of care recipients of all disease types, in order to yield more inclusive narratives.

METHODS

As part of a larger research program, the full methodology has previously been published.1 Phase A employed the use of a post-intervention design to empirically quantify changes in carer-employee’s health following the CFWP intervention. Interview survey measures used to capture carer-employee health include: self-reported health; psychosocial health, and depression. The intervention took place between the pre-test (Time 1 or T1) and post-test (Time 2 or T2), while the terminal point (Time 3 or T3) survey was implemented approximately 6 months after T2.

Recruitment and Demographics

This study was conducted at a large university located in Southern Ontario, Canada. Individual participants employed at the university were invited to participate in the study through an extensive multi-pronged recruitment strategy which encompassed: postcard and poster advertisements, electronic ads on TV displays screens, advertisements via e-mail lists, word of mouth, and, cold calling across various departments.

Eligibility criteria of participants included: working full-time in some capacity at the institution concerned while also actively caring for an adult dependent. The study procedure and purpose were verbally explained to each potential participant over email or phone. Interested participants were provided with a written consent form to review and sign. Before each data collection point, oral consent from participants were confirmed when verbally reminded of the study’s goals, design, and their rights as participants to withdraw at any time or choose not to answer any questions. A total of 21 participants made up the final convenience sample.

Each participant completed the same CFWP interview survey, where researchers would verbally ask each question (either in person or verbally over the phone) and record participant responses on an electronic copy of the survey. Survey data were collected at three different time periods (T1 to T3) across a 12-month period. The first T1 surveys were collected in the spring of 2016 and the last T3 surveys were concluded January 2018.

Table 1 depicts fluctuations in the sociodemographic profile of each participant pool across time. Overall, our sample population appears to be fairly homogenous. The participant population were female-skewed, mostly over the age of 45 years old, Caucasian, highly educated, and likely to have a household income of over $70k CAD per annum.

Marital status, income, sex, education, years are current job, and number of care-recipient tend to be fairly stable over time, even with participant drop-out. Overall, the sociodemographic profile of withdrawing participants was fairly similar to the remaining pool. While we cannot argue for random distribution of participant drop-out, we observe that there does not appear to be an overt sociodemographic pattern of participant withdrawals.

Participant Drop-Out

Forty-three (43) participants were originally recruited at T1. From T1 to T2, 13 of the 43 participants withdrew (N = 30). By the end of this study (T3), only half of the sample size (n = 21) was left due to the disqualification of participants based on the inclusivity criteria. Disqualification was due to: (1) the care-recipient passing away (two participants); (2) the participant no longer providing informal care (two participants); (3) the participant no longer working full-time (three participants). Ten participants withdrew from the research program for reasons not disclosed, while an additional two participants withdrew citing time constraints due to work and family responsibilities. Three participants were non-responsive or had invalid responses. Data from these participants were removed from the study, as per ethics guidelines.

Intervention

The educational intervention itself consisted of an in-person meeting with each individual participant, which occurred between T1 and T2 with the first author. Demographic and employment data were collected from the participants and entered into a web-based decision tool, which then produced a customized list of tailored caregiving and work-related resources and programs at the community, workplace, provincial and federal level. These options were explored in detail between the researcher and participant. Following this, participants were provided a list of behavior change goals from a checklist and encouraged to focus on two items to attempt to accomplish by the next meeting. Participant progress on the checklist activity was then followed up with at T2 and T3.

Survey Instruments

The selection of instruments and variables captured in the survey was designed by the Principle Investigator in consultation with economists and work-family researchers. The survey captured a diverse set of variables; however, only health related variables are reported in this paper. The three manifest scales used for analysis in this paper include: self-reported health (SF-12), carer reaction (CRA), and depression (CES-D). All instruments were selected for their validity, moderate to excellent internal consistency and reliability, in addition to their prevalence in clinical health and sociological literature, particularly in American and European contexts.42–46

The SF-12 manifest scale contained 12 items that evaluated participants’ perception of their own physical and mental wellbeing. Examples of topics probed include: level of physical activity, self-assessments of pain/discomfort, mood, emotional problems, and execution of routine tasks. The CRA scale contains 24 items and probed dimensions of psychosocial health pertaining to financial burden, caring activity, familial support, physical/emotional strain, and attitude towards caring. The CES-D is a 10-item scale investigating frequency and intensity of depressive symptoms. Most of the answer options for each scale were formatted using a Likert scale, such as: from “Excellent” (a score of 1) to “Poor” (a score of 5),
and “All of the time” (a score of 5) to “None of the time” (a score of 1). All three scales explore multidimensional aspects of health.

Analysis of all data was conducted on R version 3.5.3 (R Foundation for Statistical Computing, Vienna, Austria). All items were converted to numerical values and coded in the same direction, such that lower scale sums are associated with positive outcomes and higher scale sums indicated more adverse outcomes. All scales were individually tested for reliability within the context of our study by creating a time-series correlation matrix and using the test-retest via intra-class correlation (ICC) coefficient method using data collected from all time points.

A health condition scale was generated by summing up all three scales at each time point, due to the multidimensional nature of the scales investigating overlapping phenomena and high reliability of the scales. This health condition scale was used as the dependent variable in the linear mixed modeling (LMM). The complete longitudinal dataset contained less than 5% missingness where instances of missing data were randomly distributed single items within the aforementioned scales. Missing data were therefore imputed using multiple imputation via predictive mean matching with five iterations, if the total number of missing items were less than 30% of the total items in the scale; otherwise, the scale would be excluded from analysis for that specific participant.47 The datasets from all three time periods were cleaned up and aggregated into long format (longitudinal) to conduct linear mixed modelling (LMM).

**Linear Mixed Modeling**

LMM was selected as the most appropriate method for analysis due to its ability to: (1) handle longitudinal data; and, (2) analyze the changes in health condition as a random effect measurement for the intervention. LMM is an extension of the simple linear regression model for longitudinal or group structured data.48 While linear regressions measure single fixed effects of a dependent variable, LMM measures the random effects, or differences, between grouped data in addition to these fixed effects. Fixed effects represent a scaled relationship between the predictor and explanatory variables, in this case, the relationship between participants’ health scores and time (T1 to T3).49 Random effects are represented by a categorical variable with levels assumed to be a random selection from a hypothetical larger population.49 They can act as a grouping factor to be controlled. In the case of this analysis, participant ID numbers were used as random effects to account for variation in health score between participants.49 In layman terms, this analysis is identifying associations between the health condition of participants and time, specifically examining the impact of the intervention on participant health, while controlling for inter-participant variation. Each time period (T1

| TABLE 1. Descriptive Statistics Containing General Demographics of Sample |
|-----------------------------|------------------|-----------------|-----------------|
| Variable                   | Value            | Time 1 (n = 43) | Time 2 (n = 30) | Time 3 (n = 21) |
| Age                        | 18–45            | 30.2%           | 20.0%           | 38.1%           |
|                            | 46+              | 69.8%           | 80.0%           | 61.9%           |
| Gender                     | Male             | 11.6%           | 10%             | 14.3%           |
|                            | Female           | 88.4%           | 90%             | 85.7%           |
| Marital                    | Married /Common-in law | 58.1%      | 56.6%           | 57.1%           |
|                            | Widowed, divorced, separated | 16.3%      | 13.4%           | 19.1%           |
|                            | Single           | 23.3%           | 26.7%           | 23.8%           |
|                            | Other            | 2.3%            | 3.3%            | 0.0%            |
| Race                       | Euro/Caucasian   | 100%            | 100%            | 100%            |
| Highest education          | College GCEP or less | 30.2%      | 23.3%           | 19.0%           |
|                            | Bachelors        | 32.6%           | 36.7%           | 28.6%           |
|                            | Graduate         | 37.2%           | 40.0%           | 52.4%           |
| Household income           | $15k–29.9k       | 2.3%            | 0.0%            | 0.0%            |
|                            | $30k–49.9k       | 7.0%            | 0.0%            | 4.7%            |
|                            | $50k–69.9k       | 9.3%            | 23.3%           | 14.3%           |
|                            | $70k–99.9k       | 27.9%           | 13.3%           | 14.3%           |
|                            | $100k+           | 46.5%           | 43.3%           | 61.9%           |
|                            | Prefer not to answer | 7.0%      | 20.0%           | 4.7%            |
| Place of residence         | Hamilton Metro.  | 67.4%           | 70.2%           | 67.0%           |
|                            | GTA              | 21.0%           | 19.9%           | 19.0%           |
|                            | Other            | 11.6%           | 9.9%            | 14.0%           |
| Yrs. at current job        | Less than 5 yrs  | 37.2%           | 36.7%           | 47.6%           |
|                            | 5–10 yrs         | 27.9%           | 27.0%           | 19.0%           |
|                            | 11–15 yrs        | 16.2%           | 16.7%           | 14.3%           |
|                            | 16–20 yrs        | 6.9%            | 3.3%            | 0.0%            |
|                            | 21+ yrs          | 9.3%            | 10.0%           | 14.3%           |
| Current health             | Poor             | 2.3%            | 0.0%            | 0.0%            |
|                            | Fair             | 9.3%            | 3.3%            | 14.3%           |
|                            | Good             | 34.8%           | 23.3%           | 28.6%           |
|                            | Very good        | 34.8%           | 60.0%           | 33.3%           |
|                            | Excellent        | 18.6%           | 13.3%           | 23.8%           |
| Number of care-recipients  | 1                | 62.7%           | 63.3%           | 76.2%           |
|                            | 2                | 23.3%           | 30.0%           | 14.3%           |
|                            | 3                | 14.0%           | 6.7%            | 4.8%            |
| Did the caregiver postpone their career or education? | No | 46.5% | 80.0% | 76.2% |
|                            | Yes              | 20.9%           | 16.7%           | 23.8%           |
|                            | N/A              | 32.6%           | 3.3%            | 0.0%            |

Note: due to rounding, some of the percentages may not add up to 100%.
to T3) in the regression was indicated by a dummy variable, while an additional dummy variable was created using time data to denote whether the intervention had occurred (T2 and T3) or not (T1).

Three random intercept LMM were created to investigate if there is an association between health scores and time as well as which time periods indicate greater changes to the participant’s health. The first model compares changes between participants health amongst the entire intervention, where T2 and T3 are aggregated as a dummy variable. The second and third model expands upon the first by examining the specific effects of the T2 or T3 dummy variables respectively, as a separate independent variable within the entire intervention.

RESULTS

Reliability Analysis

The correlation matrix reported the manifest scales across all time periods to be: reliable; somewhat consistent; showing a significant increase from T1 to T2, and; having relatively strong linkages (Table 2). The strength of all correlations are classified as strong (0.5+) with a range of 0.58 to 0.92, indicating strong correlations between all the scales. 50

The ICC test-retest analysis shown in Table 3 conveys a range of good (depression and SF-12 scales) to excellent (psychosocial and health score) inter-rater agreement measures. 44 This indicates that the manifest scales in all time periods are reliable and consistent for further analysis.

Descriptive Statistics

Figure 1 displays the health score of each participant over time, and Table 4 shows mean values and standard deviation of the health score. The majority of the participants’ health improved greatly between pre-intervention (T1) to the first wave of the intervention (T2), as noted by the downward trend in health condition score (from 100 at T1 to 85.8 at T2). Between T2 and T3, the plot (Fig. 1) illustrates mixed results with some participants getting a bit worse while others not improving much relative to the changes from T1 to T2. Overall, the health condition score increased from 85.5 at T2 to 89.6 at T3 (Table 4). The average health condition scores from T1 to T3 conveys a downward trend, from 100 (T1) to 89.6 (T3).

Power Analysis

Given the small sample size, a power analysis was necessary to validate the results of the analysis. Figure 2 depicts a plot of minimum sample size required for a specific effect size. In the context of our study, we set our type I error at 0.05 (paired), and the statistical power (1 – β) was set at 0.8 to test the validity of intervention when the effect size goes from about 1.0 to 0.1. Based on Cohen guidelines, 51 a moderate to large effect size (d ≥ 0.5) requires a sample size of approximately 20 or less. Based on this result, we can verify the validity of our results with a small sample size.

Principal Component Analysis

A socioeconomic index was generated using principal component analysis (PCA) to determine the extent of social and demographic variation in our final T3 sample.32 This was done as the small sample size in our study precluded the addition of sociodemographic and economic variables in our LMM models, and there was a need to control these variables. PCA was conducted to reduce the dimensionality of socioeconomic variables, such as age, sex, education level, income and marital status, through creating an index that examines total variance. The first principal component (PC1) indicates the orthogonal linear combination of the variables that accounts for the maximum variance in the data. 53 Subsequent components capture unrelated and lesser variation; thus, PC1 provides the most information. 54 Socioeconomic and demographic data for each participant was converted into a binary format, and tetrachoric correlation was used for the PCA index, as seen in Table 5 below.

Table 6 depicts the variable loadings resulting from the PCA, as well as the proportion of variance of each component. Principal component loadings represent eigenvalues that indicate weights of each variable on the principal component. 55 PC1 had a proportion variance of 38%, in contrast to PC2 and PC3, which had 32% and 23% respectively. Only the first three components (PC1 to PC3) were retained, as the variance in the first three components account for 93% of the total variation when the proportion of variance score was summed.

PC1 explains the largest variance of the data and has large positive loadings on income and marital status together with large negative loadings on sex; this indicates that income and marital status are inversely proportional to sex. PC2 had large positive loadings on age and sex, while having large negative loading on education. PC3 contains large negatively loadings on sex, marital status, and education only.

Factor scores were generated by multiplying principal component loadings with the proportion of variance for each variable and summed through PC1 to PC3 (Table 6). The absolute value of each factor score represents the weighting of each respective variable (either as positive or negative) of the carer-employee’s socio-economic status. Age (46+ years), marital status (married), and income ($70k+) are positive, indicating a higher socio-economic status of the carer-employee's socio-economic status.

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### Table 2. Correlation Matrix With Respective Correlation Coefficients of Health Indices Across Three Time Periods

| Correlation               | T1        | T2        | T3        |
|---------------------------|-----------|-----------|-----------|
| Psychosocial to depression| 0.62      | 0.87      | 0.71      |
| SF-12 to depression       | 0.62      | 0.92      | 0.67      |
| SF-12 to psychosocial     | 0.58      | 0.78      | 0.65      |

SF-12, self-reported health.

### Table 3. Test-Retest One-Way Random Effects Model Using Single Measures

| Variable     | ICC   | Lower | Upper  | F Test with True Value 0 |
|--------------|-------|-------|--------|-------------------------|
| Depression   | 0.633 | 0.402 | 0.813  | 6.18                    |
| SF-12        | 0.713 | 0.510 | 0.800  | 8.46                    |
| Psychosocial | 0.801 | 0.642 | 0.910  | 13.1                    |
| Health score | 0.834 | 0.696 | 0.922  | 16.1                    |

ICC, intra-class correlation; SF-12, self-reported health.
employee. These findings match the demographic profile found in the caregiving literature.\textsuperscript{16,17} Interestingly, education (Bachelors or higher) and sex (female) reveal negative scores, thus, lowering the socioeconomic status of the carer-employee. A plausible explanation for sex may be that women are more likely to be caregivers when compared with men (as noted earlier), which would hinder their career potential. A higher education may entail more job responsibilities and, together with the responsibilities of caregiving, may leave the carer-employee stressed. Overall, socioeconomic variation within our sample population is not strong, suggesting that these demographic variables are likely not underlying confounding factors within the analysis.

**Linear Mixed Models**

All three models reported similar goodness of fit and statistically significant intercepts on both random effects and fixed effects (Tables 7–9). The first model (Table 7) looked at intervention effects over the whole study (T1 to T3), with T2 and T3 variable aggregated. The model showed the intercepts on both random and fixed effects to be statistically significant with relatively low standard errors. The coefficient of overall time of intervention was approximately –12 and statistically significant. This indicates that from T1 to the combined time of intervention (T2 and T3), the participants reduced their health condition score by –12.

The second model (Table 8) used the same aggregation of T2 and T3 to examine intervention effects over the entire study, but used T2 as a factor in the regression equation. Model 2 had a coefficient of –10 for the overall intervention, when factoring T2 as a dummy variable, while the T2 variable had a coefficient of –4, though this was not statistically significant.

For the third model (Table 9), the overall time of the intervention had a coefficient of –14, with statistical significance. The dummy variable of T3 displays an approximate increase of 4, despite not being statistically significant. This finding similarly reflects the plot in Fig. 1, which illustrates that some participants increased their health condition score after T2. In general, when implementing the overall intervention, the health condition score of the participants is reduced, which improves their general wellbeing. Overall, the first model seems to be the most appropriate model for the data, as it is simpler and has slightly better Akaike information criterion (AIC) and Bayesian information criterion (BIC) scores.

### TABLE 4. Aggregated Health Scale Means at Each Time Point (T1–T3)

| Time Period | Mean Aggregated Health Condition Score (N = 21) | Standard Deviation |
|-------------|-----------------------------------------------|--------------------|
| T1          | 100                                           | 25.5               |
| T2          | 85.8                                          | 27.2               |
| T3          | 89.6                                          | 30.1               |

FIGURE 1. Individual participant aggregated health condition scores over time (T1 to T3), where each plot represents a participant. Participant numbers are labeled above each plot.
FIGURE 2. Minimum sample size required for effective changes, where greater sample sizes are needed to verify smaller effect sizes.

TABLE 5. Assignment of Participant Categorical Data into Binary Values

| Variable          | Reference          |
|-------------------|--------------------|
| Age               | 1 = 46+ yrs        |
| Marital           | 1 = Married/Common-law |
| Education         | 1 = Bachelor’s degree or higher |
| Income (annual)   | 1 = $70k+         |
| Gender            | 1 = Female         |

TABLE 6. Variable Loadings of First Three Principal Components

| Variable     | PC1    | PC2    | PC3    | Factor Score | Mean Index |
|--------------|--------|--------|--------|--------------|------------|
| Age          | 0.337  | 0.603  | N/A    | 0.2580       | 0.2434     |
| Gender       | −0.422 | 0.462  | −0.536 | −0.1320      | −0.1261    |
| Marital      | 0.489  | 0.210  | −0.601 | 0.1151       | 0.0722     |
| Education    | 0.219  | 0.511  | 0.576  | 0.3748       | 0.2965     |
| Income       | 0.649  | −0.343 | −0.123 | 0.1067       | 0.0932     |
| Proportion of variance | 0.38 | 0.32 | 0.23 | N/A | N/A |

TABLE 7. Random Intercept Model 1: Intervention (T2 and T3 Aggregated)

**Formula:** Health Condition ~ Intervention + (1|ID)

**Random Effects**

| Groups         | Name   | Lower Variance | Variance | Upper Variance | Std. Error | Pr (>|t|) | Sig. |
|----------------|--------|----------------|----------|----------------|------------|----------|------|
| ID             | Intercept | 468.6         | 647.4    | 952.8          | 3.21       | 0.002    | **   |
| Residual       | Intercept | 61.1          | 84.6     | 124.2          | 1.16       | 0.252    |      |

**Fixed Effects**

| Variable   | Estimate | Std. Error | Df | t-Value | Pr (>|t|) | Sig. |
|------------|----------|------------|----|---------|----------|------|
| Intercept  | 99.7     | 5.90       | 24.6 | 16.9   | 0.000    | ***  |
| Intervention | −12.0   | 2.46       | 42.0 | −4.90  | 0.000    | ***  |

AIC = 533.0; BIC = 541.5.
*Denotes 0.01–0.05 P-value.
**Denotes 0.001–0.01 P-value.
***Denotes P-value < 0.001.
DISCUSSION

This study examines the effectiveness of CFWPs within the workplace environment by assessing changes in carer-employees’ health. To our knowledge, this is the first study of its kind, where the relationship between a workplace-based CFWP intervention for carer-employees, and the mitigation of negative effects with respect to physical and emotional health, psychosocial health, and mental aptitude has been studied. Data were cleaned and validated for suitability prior to all analysis. Due to the small sample size, several preliminary analyses, such as: reliability analysis, correlation matrices, power analysis, and PCA, had to be conducted to confirm appropriateness of our data for final analysis.

Our findings suggest that the original objective of this paper was met; the workplace intervention was successful in reducing self-reported health, psychosocial health, and depression outcome variables, resulting in overall improvements of carer-employees’ health from T1 to T3. Compared with the baseline measurements of T1, all of the participants (N = 21) who received the standardized intervention generally reported significantly fewer or reduced symptoms of depression, psychosocial burden, and poor self-reported health during the post intervention assessment (T2) (Fig. 1). These effects were measured to be statistically significant (P < 0.05) within and between participants (Tables 7–9). While we observed an overall decline in adverse health variables throughout the whole project (T1 to T3), transformations were non-statistically significant (P > 0.05) between T2 to T3 (Tables 8 and 9), suggesting that intervention effects were maintained over time. The improvement in health outcomes are evident across all three LMM models.

Model 2 and 3 (Tables 8 and 9) explored the effects of T2 and T3 as a dummy variable in the regression equation. However, the exact effects of T2 and T3 specifically on the intervention were inconclusive due to the non-significance of T2 and/or T3 as a regression variable, despite the models indicating significant overall intervention effects. With that, we highlight Model 1 is the best representation of the data, as it is the simplest model that conveys the effectiveness of the intervention over time.

While existing evidence for carer interventions are limited, our findings generally align with evidence from the literature, in which carer health can be improved by interventions. A meta-analysis of 127 interventions found that psychoeducational interventions are able to produce significant effects on carer burden, depression, subjective wellbeing, carer knowledge, and care recipient symptoms, although these effects are most pronounced in interventions with active carer participation rather than information only.38 Our study differs in that we were able to produce significant effects on carer health with a largely informational intervention. This may reflect the setting of our intervention, in which information and resources were provided in person and discussed with the...
Carer-employees. This dialogue between the research and participant may allow greater information processing, understanding, feedback, and encourage greater positive behavior change than solely an information exchange.

Beyond short-term intervention effects, few studies have examined the long-term maintenance of intervention effects, with existing literature having mixed results. In previous literature, the magnitude of observed educational intervention effects on carer burden tend to decrease over time.56,57 This may be due to the fact that carer burden, stress, and mental health are complex multidimensional phenomena, that may not be easily treated with educational interventions.37,39 Indeed, interventions with more psychosocial components appear to have a higher maintenance rate of effects; Vázquez et al.58 were able to produce a significant reduction in major depressive episodes between the intervention groups and control groups 12 months after a cognitive-behavioral intervention for carers. In addition, the New York University Caregiver Intervention (NYUCI), a popular psychosocial intervention for dementia carers, has extensive evidence from several clinical randomized controlled trials demonstrating long-term (12 months post-intervention) reduction of depression symptoms in carers.59–61

We believe our study is unique in carer literature in that we are evaluating a workplace based intervention for carer-employees with carer-specific resources from multiple avenues, including workplace resources, as well as governmental and community resources. The workplace setting and composition of our sample, composed entirely of carer-employees, differs from other caregiving intervention research, as the context of the workplace and employment introduces additional nuances, such as work culture or supervisory relationships, that may complicate compliance with the intervention.

Despite this, our results indicated a tangible effect of the workplace intervention on carer-employee health outcomes, specific to self-report health, psychosocial health, and depression. Further research, such as modifying the intervention design to include counselling components and extending the time period, may be required to expand our findings. One potential adaptation particularly amendable to workplaces is the introduction of multi-level interventions, which calls for multiple targeted interventions at the individual and organizational levels.62

**IMPORTANT IMPLICATIONS**

Our findings affirm that our designed intervention may serve as an effective workplace accommodation tool for uptake by organizations to avert not only the clinical burden of caring but also protect employee health. While this study was not a clinical study, our findings may be useful for policy-makers, HR professionals, occupational health specialists, managers and employers, as our outcomes demonstrates effectiveness of an educational intervention at improving employee health, particularly mental health. The impact of such an intervention in the workplace is twofold: (1) employees’ health may be improved longitudinally, resulting in less sick days and reduced absenteeism and; (2) enhanced employer reputation, resulting in increased employee satisfaction, productivity, and retention as a result of reduced conflict between work and non-work.5,64 Our study is supportive of the estimations made in a prior paper estimating the economic effects of our intervention, working the cost of different effectiveness models and our intervention may have saved the workplace between $48,010 to $676,657 CAD.10 While we did not evaluate work-related variables in this paper, our findings specific to intervention outcomes are encouraging, as increases in health outcomes are associated with work performance.65

The caregiving population is expected to increase in the coming years and is poised to become a dominant issue for employers. When supports for carers are inadequate, carer burden can progress into carer burnout, a serious mental health condition that can lead to deterioration in many facets of carer’s lives, including paid work performance and health. Carer educational interventions in the workplace offer several unique advantages: (1) ability to reach large audiences at once, (2) utilization of existing infrastructure (HR staff, existing work policies) to augment interventions, and (3) benefit to the bottom line.

In the past, issues regarding child care prompted employers to implement flexible work arrangements, and maternity/paternity leaves in order the keep the workforce healthy and aid their employees in maintenance of their multiple obligations.66 Similarly, eldercare issues should now receive the same attention and support from workplaces as our global population ages. We conclude that this intervention, an education resource for carer-employees, is a feasible and effective strategy for uptake by workplaces.

It is important that we underscore here that, although our intervention demonstrated substantive influence in reducing negative health outcomes, the nature of our recruitment strategies likely yielded high-interest carer-employees who were seeking and/or open to CFWP accommodations. While we have observed effectiveness of our designed intervention in a convenience sample of carer-employee participants in researcher-controlled conditions, we cannot make the same argument for the broader working environment. As the data collection points and intervention session were conducted one-on-one in-person, between the participants and researcher, it is possible that participants may have modified their behavior during these meetings. Additional studies could assess: reproducibility of results, barriers to implementation, and uptake of a standardized intervention in the work setting.

We recognize that this intervention study has a number of limitations, the foremost being the small sample size. Of the initial 43 participants at T1, only 21 participants remained by T3, creating only 21 sets of complete longitudinal data. Despite the power analysis and PCA, we are unable to determine if there were systemic or selection biases present in the population that left the study. Given the nature of the carer-employee role, our study targeted a population known for being highly stressed and over-burdened, leading to difficulties in recruitment and retention. It is possible that the participants engaged in our research are not representative of the whole carer population, as many highly burdened carers would likely not be able to join our study. Furthermore, those who withdrew from our study likely experienced high stress and work-life conflict, resulting in their departure. This may render fluctuation effects more pronounced in our analyses, and potentially skew our produced results.67 A further limitation includes the lack of a control group in our study to compare intervention results, which may lead to misidentification of random noise in our data.65 Lastly, our intervention study was conducted solely on university employees; further studies may need to be done to determine effectiveness of intervention protocols in different sectors and workplaces.

**CONCLUSIONS**

Carer-employees are a large and important segment of Canadian society, and given Canada’s aging demographic, provide an innumerable service to their families as well as society. This underscores the need for recognition and support for CEs from a wider audience, including not only employers, but also public health and occupational health specialists. In our examination of the effects of the CFWP intervention, our current understanding is that the intervention was generally successful in decreasing negative health outcomes longitudinally in our population of university-employed carer-employees. This intervention can be considered effective in improving carer-employees overall health outcomes, such as self-reported health, psychosocial health, and depression, over time.
Regardless, the limitations present in this study calls for the need to carry out similar types of studies with larger sample sizes and in different workplaces. Subsequent studies should focus on applicability of our results in other employment sectors, while identifying barriers to uptake within working environments.

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