Giving Social Support at Work May Reduce Inflammation on Employees Themselves: A Participatory Workplace Intervention Study Among Japanese Hospital Nurses

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Research note

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Abstract

Objective

Previously, we reported that the participatory workplace intervention was effective in reducing stress-related inflammatory markers, i.e., interleukin-6, among 31 Japanese female nurses. During the analysis, we recognized that our intervention might have increased prosocial behaviors like giving social support to others in some participants. Based on this assumption, we ran a secondary analysis, which examined the effect of giving social support to others on inflammatory markers, autonomic nervous activity, and perceived job stress before and after a 5-month intervention. We divided participants into two groups; those who had increased scores on giving social support after the intervention (Group 1, n = 13), and those who had decreased/unchanged in the scores (Group 2, n = 17). Friedman test was used to examine the changes in outcome measures by the group.

Results

Group 1 showed significant decreases in interferon-γ, interleukin-6, and interleukin-12/23p40 immediately after the intervention, while interleukin-12/23p40 remained stably decreased three months later; Group 2 did not show changes in these markers. No significant changes were observed regarding autonomic nervous activity and perceived job stress. This study presented a significant insight that giving social support at work may provide health benefits towards employees themselves, via decreasing inflammation in the body.

Introduction

Organizational-level participatory workplace intervention, which aims to improve work environment and employees’ health, is more likely to produce sustainable effects than interventions targeting an individual because employees take an active part in identifying problems and giving possible solutions by themselves [1, 2]. In our previous study, we reported that a 5-month lasting organizational-level participatory workplace intervention was effective in reducing stress-related inflammatory markers as represented by interferon (IFN)-γ, interleukin (IL)-6, and IL-12/23p40, and IL-15 among 31 Japanese female nurses [3]. During this course, we recognized that our intervention might have stimulated prosocial helping behaviors like giving social support to others in some participants. Based on this assumption, we decided to run a secondary analysis focusing on giving social support and physiological responses.

It is well documented that social support acts as a stress buffer, which contributes to improving mental and physical health [4–6]. With regard to physiological markers, a number of studies reported the existence of positive associations between receiving/perceived social support and inflammatory markers and autonomic nervous activities (ANA) [7–10]. These studies mainly focused on receiving social support at work rather than giving social support. There is a lack of evidence on the effects of giving social support on physiological outcomes. Although limited studies on giving social support, two intervention studies examined the effects of giving social support on physiological responses (inflammatory markers,
heart rate, blood pressure, salivary alpha-amylase, and salivary cortisol) among healthy individuals [11, 12]. These studies revealed that giving social support contributed to decreasing inflammatory markers, systolic blood pressure, and salivary alpha-amylase [11, 12]. However, the study settings were experimental, i.e., the intervention was to imagine someone whom participants wanted to support and write a supporting letter to him/her, etc. To the best of our knowledge, no organizational-level studies to date have examined giving social support and physiological outcomes in a work setting.

Therefore, the present study aimed to explore how changes in giving social support to others at work affect physiological responses among Japanese female nurses. We hypothesized that those who increased giving social support by the intervention would have a positive effect on inflammation and ANA; those who had increased scores on giving social support after the intervention exerted decreased inflammatory makers and ANA to healthier status compared to those who had no change or decreased scores.

Methods

Participants and procedure

We carried out a participatory workplace improvement intervention [13–22] from August 2017 to February 2018. Briefly, the participatory workplace improvement intervention is that employees at the workplace actively take part in identifying workplace problems, find feasible actions/solutions, and work towards improvement. We recruited nurses (n = 144) working at a hospital with 150 beds in the southern part of Japan. A total of 36 nurses agreed to participate in this study. We conducted evaluations before the intervention for baseline (T1), within a week after the end of the intervention to assess immediate effects (T2), and 3 months after the end of the intervention to assess prolonged and lasting effects (T3). We excluded participants who became pregnant during the study period (n = 1), missed evaluations (n = 3) and had incomplete responses in giving social support scores (n = 1). A male participant (n = 1) was also excluded because of possible sex differences in outcome measures. Therefore, a total of 30 female nurses were submitted to the final analysis.

Measures

Sociodemographic, lifestyle, health, and occupational conditions

We used the self-administered questionnaire to assess participants’ sociodemographic and job-related characteristics including social support at work and perceived psychosocial job stress.

Giving social support

In the questionnaire, we included questions of ‘giving’ social support to others at work, which we modified from ‘receiving’ social support in the Brief Job Stress Questionnaire [23], “How much help do you provide
to the following people?”, “How much are you relied on by the following people?”, “How well do you listen to the following people when they ask for advice on personal matters?”. Participants answered each question by “superiors”, “co-workers”, and “subordinates” with a four-point scale (1 = extremely to 4 = not at all). Cronbach's alphas for these items were > 0.630 at all-time points.

**Inflammatory markers**

We used serum interferon (IFN)-γ, interleukin (IL)-6, tumor necrosis factor (TNF)-α, IL-12/23p40, IL-15, IL-27, and high-sensitivity C-reactive protein (hs-CRP) as inflammatory markers. On the evaluation days, participants (nurses) brought their blood samples which collected between 2 pm and 5 pm in gamma-rays sterilized polyethylene-terephthalate tubes containing serum separating gel and coagulation accelerant (silica particles). We stored the samples in a cooler box (0–5°C) and transported them to our laboratory twice a day by 4:30 pm and 7:30 pm. In the laboratory, we centrifugalized them with 2,400 rpm for 10 minutes to extract 500µL of the serum and deep-freezed (-20°C) until the analysis. The level of inflammatory markers was assessed with the Enzyme Immunoassay or Chemiluminescent Enzyme Immunoassay with MESOTM QuickPlex SQ 120 (Meso Scale Diagnostic, LCC, Rockville, USA) by the analyzing company, Life Science Institute Medience Corporation, Japan. The minimum detectable level for IFN-γ, IL-6, TNF-α, IL-12/23p40, IL-15, IL-27, and hs-CRP was 0.2 pg/ml, 0.06 pg/ml, 0.04 pg/ml, 15.0 pg/ml, 2.0 pg/ml, 8 pg/ml, and 0.004 mg/dl, respectively. We calculated the values lower than them into the minimum detectable level/√2, as it was described elsewhere [24].

**Autonomic nervous activity (ANA)- Heart rate variability (HRV)**

We utilized an electrocardiograph device, Silmee Bar Type Lite (Silmee; Tokyo Denki Kagaku, Tokyo, Japan) to measure heart rate variability (HRV). Silmee automatically calculates HRV by the power spectral analysis and measures 3 sympathetic nervous activity (SNA); low-frequency HRV/total frequency HRV (standing position), mean R-R interval/R-R interval per minute (standing position), and mean R-R interval (supine-stand position), and 3 parasympathetic nervous activity (PNA) parameters; mean R-R interval (supine position), high-frequency HRV/total frequency HRV (supine position), and the standard deviation of R-R intervals (SDRR) (supine position). Silmee also calculates SNA/PNA. We measured participants’ autonomic nervous activities in two rooms at the hospital between 2 pm and 5 pm to adjust in-day fluctuation.

**Statistical Analyses**

Based on the total giving social support score at each time-point, we divided participants into two groups; those who had increased scores on giving social support to others after the intervention (Group 1, n = 13), and those who had decreased/unchanged in the scores (Group 2, n = 17). After the confirmation of non-Gaussian distribution with the Shapiro-Wilk test, we applied the Friedman test to examine changes in inflammatory markers, autonomic nervous activity, and perceived job stress by the group. We analyzed data using IBM SPSS Statistics for Windows, version 25.0 (IBM Corp., Chicago, IL, USA), and the level of significance was set at $p < 0.05$. 
Results

Table 1 shows the baseline characteristics of female nurse participants. The median age of participants was 37.0 years old for Group 1 and 38.0 years old for Group 2. More than 60% of participants were not married in both groups. Except for department G (nursing department), 4 to 6 nurses in each unit participated. More than half of the participants worked for the day shift in both groups. Over 60% of participants had 6 or more hours of sleep on workdays in both groups. Most of them in both groups were under regular menstrual cycle (> 76.5%).

Table 2 presents the changes of physiological markers overtime in Group 1. IFN-γ ($p = 0.005$), IL-6 ($p = 0.018$), and IL-12/23p40 ($p = 0.018$) were decreased at T2 compared to T1. IL-12/23p40 was decreased at T3 compared to T1 ($p = 0.013$). The overall changes of TNF-α were also significant ($p = 0.021$), but it was insignificant with Bonferroni-adjusted pairwise tests. No significant decreases were found in ANA.

Table 3 shows the changes of physiological markers in Group 2 over time. Neither blood inflammatory markers nor ANA showed insignificant decreases.

Tables S1 and S2 (Additional files 1 and 2) present the changes of perceived job stress over time by the groups. There were no significant decreases with Bonferroni-adjusted pairwise tests in both groups.

Discussion

This study examined the effect of giving social support to others on inflammatory markers, autonomic nervous activity, and perceived job stress before and after the participatory workplace intervention among Japanese female nurses. As we hypothesized, the group with increased levels of giving social support (Group 1) showed significant post-intervention decreases in inflammatory markers (IFN-γ, IL-6, and IL-12/23p40), while another group with decreased/unchanged levels of giving social support (Group 2) did not show such changes in these markers. ANA and perceived job stress did not show significant changes in both groups. This is one of the first studies to examine changes in giving social support to others at work after an organizational-level intervention using multiple physiological markers.

We observed decreases in inflammatory markers only in Group 1. Our finding is comparable with a past study regarding giving social support and inflammatory markers. A study by Moieni et al. reported that increases in giving social support levels are related to decreases in inflammatory markers [12]; a 6-week gratitude intervention resulted in decreases in the percentage of monocytes producing IL-6, TNF-α, and coproducing IL-6 and TNF-α via increases in support-giving among healthy middle-aged women. On the other hand, the control group did not lead to such changes. Our results also imply that giving social support to others at work may contribute to improving health by decreasing inflammatory markers on employees themselves.

Although several inflammatory markers had decreased in Group 1 after the intervention, another physiological measure (ANA) remained unchanged in the same group. The plausible explanation is that
positive outcomes may emerge at different timing in each measure [3]. Past participatory workplace intervention studies did not also obtain positive effects simultaneously in all stress-related measures they used in the intervention group [20-22, 25], despite longer intervention period compared to our study. The various types of measures and timing of evaluation may have led to disaggregated results on inflammatory markers and ANA in Group 1.

In conclusion, this study presented a significant insight that increases in giving social support to others at work may have positive health effects on employees themselves via reducing inflammation in ones’ body. Further studies with a better study design, i.e., randomized control design, are required to confirm the effects. Moreover, work remains to investigate the amount and timing of giving social support at work, and towards who (supervisor, family, etc.) provides social support is important in exerting beneficial effects on employees’ health.

Limitations

- Small sample size
- No control group
- Other confounding factors might be influenced the results
- Data analysed by two separate groups due to the non-Gaussian distribution of the data.

Abbreviations

Group 1: the group of nurses with increased scores in giving support after the intervention; Group 2: the group of nurses with decreased/unchanged scores in giving support after the intervention; ANA: Autonomic nervous activity; HRV: Heart rate variability; IFN: Interferon; IL: Interleukin; TNF: Tumor necrosis factor; hs-CRP: High sensitivity C-reactive protein; T1: baseline evaluation; T2: within a week after the intervention; T3: three months after the end of the intervention.

Declarations

Acknowledgments

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Authors’ contributions

AN designed the study and TT, AN, and AA contributed coordination. TT, AN, YO, and AA facilitated recruitment, and TT, AN, and YO facilitated the intervention. TT, NY, HK, and NS contributed data collection and TT performed serum extraction. AN supervised TT on the data analysis and paper conduct. All authors read and approved the final manuscript.
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**Availability of data and materials**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**

The study was reviewed and approved by the ethical committee of the International University of Health and Welfare (18-ml-002). This study was registered on the University Hospital Medical Information Network Clinical Trials Registry (UMIN000039836). We informed potential participants about the study aim, procedure, and confidentiality policy for individual information. Written informed consent was obtained from those who agreed to participate. At each time of the evaluation, they received a 1,000-yen gift card as a reward.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**References**


Tables

Due to technical limitations, table 1, 2 and 3 is only available as a download in the Supplemental Files section.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- TableS1group1BMCPercievedStress.pdf
- TableS2Group2BMCPercievedStress.pdf
- Table1BMCDemographics2.pdf
- Tables2group1BMCTONDOKORO.pdf
- Tables3group2BMCTONDOKORO.pdf