Self-care interventions in stroke survivor–caregiver dyads: a protocol for systematic review and meta-analysis

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ABSTRACT

Introduction Stroke is known as one of the leading causes of mortality and disability worldwide. Self-care plays a significant role in improving the quality of life, self-efficacy and many other outcomes of stroke survivors. However, it is a dyadic phenomenon where patient self-care and the caregiver contribution to self-care are interrelated in terms of predictors and outcomes. Currently, there is still no systematic assessment conducted to examine the overall effectiveness of self-care interventions carried out in stroke survivor–caregiver dyads and explore the effect on stroke survivor and/or caregiver outcomes.

Methods We plan to conduct a systematic review and meta-analysis of the evidence regarding the self-care interventions carried out in stroke survivor–caregiver dyads. We will undertake a systematic search of multiple databases including PubMed, Web of Science, CINAHL, PsycINFO, EMBASE, Cochrane Central Register of Controlled Trials and four Chinese databases (CNKI, CBM, WANFANG and VIP) from inception to July 2021 for the purpose of collecting the relevant articles. The eligible studies are defined as those original researches, written in English or Chinese, on self-care interventions in stroke survivor–caregiver dyad samples. Two independent researchers will be deployed to identify the eligible trials according to the selection criteria and extract the relevant data. The Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols checklist has been used for this protocol. We will use the Cochrane Risk for Bias tool to assess the risk of bias for randomised controlled trials.

Ethics and dissemination In our review, any identifiable patient data will be excluded, which removes the need for ethical approval and participant consent. The final results of our study will be published in an open-access peer-reviewed journal, and abstract will be presented at suitable national/international conferences.

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INTRODUCTION

Stroke is known as one of the leading causes of mortality and disability that incurs significant societal cost and burden.1 According to the Global Burden of Disease Study, there were about 80 million stroke survivors worldwide in 2016, apart from 13.7 million new stroke survivors each year.2 According to a recent systematic review (SR), the cumulative recurrence rate reached 7.7% and 9.5% 3 months and 6 months after stroke, respectively, with 13.2% and 34.8% of all stroke inpatients getting hospitalised due to the recurrence of stroke.3 The poor functional, cognitive and psychological outcomes manifested after stroke, such as haemiplegia, aphasia and depression, have negative effects on a large proportion of long-term stroke survivors.4 In some studies, it has been demonstrated that the good self-care after stroke can contribute to the recovery of various physiological functions for patients, thus improving the quality of their life. In the meantime, stroke recurrence is prevented and the readmission rate is reduced.5 6 In 8760 hours throughout a year, patients spend merely 0.001% of their time with medical staff, which means that all other activities related to health maintenance, monitoring and management are carried out either by individuals or by patients and their families as the self-care activities performed outside the clinical or hospital environment.5 The study of Haley showed that stroke survivor and caregiver reports of engagement were...
closely correlated with each other (r=0.89). In practice, self-care is often cocare for stroke survivors. That is to say, patients often practice stroke self-care together with their informal caregivers, especially those family members who exert significant influence on the self-care process. Additionally, there have been more and more theoretical and practical studies indicating that stroke can be viewed as a dyad event, which can affect the physical, psychological and social functions of patients and their caregivers.

Self-care is defined as a naturalistic decision-making process aimed to address both the prevention and management of chronic illness. 

According to the Middle-Range Theory of Self-Care of Chronic Illness, self-care is interpreted as a process of maintaining health. 

Besides, it can be regarded as an overarching construct based on the three critical concepts including self-care maintenance, self-care monitoring and self-care management. As for self-care maintenance, it refers to those behaviours performed to improve well-being, maintain health or ensure physical and emotional stability. Self-care monitoring is defined as a process of routine, vigilant body monitoring, surveillance or ‘body listening’. Self-care management requires an evaluation of the changes in various physical and emotional signs and symptoms to determine whether it is necessary to take action in response. The performance of sufficient self-care is supposed to encompass all three behaviours because these three concepts are closely related to each other. 

Self-care represents a fundamental element of the treatment received by those patients with chronic conditions and is a major focus of many interventions. Self-care interventions can play a role in increasing choice, improving accessibility and affordability as well as presenting opportunities for individuals to make informed decisions on their health and healthcare. 

Despite a series of SRs suggesting that the self-care interventions/programmes, which target only patients improved the quality of life (QoL), self-efficacy, coping ability as well as activities of daily living (ADL) and social participation for patients with stroke, which makes them different from inactive or active (usual care) control interventions, there remain some limitations on these self-care interventions, for example, the self-management compliance of stroke survivors was poor, some self-care skills could not be developed independently by them, and the contributions of stroke caregivers could not be ignored in the self-care of stroke.

Typically, two individuals (such as husband and wife) maintaining a sociologically important relationship are defined as a dyad. According to the theory of dyadic illness management, the patients suffering chronic diseases and their caregivers take joint participation in disease management, which involves decision-making, emotional communication support, the management of changes in physical function and general health behaviours. The ultimate goal of doing this is to promote the dual health of patients and their caregivers. In previous studies, it has been demonstrated that the health outcomes of stroke survivors and caregivers affected each other and was greatly impacted by the physical function changes of the survivor. Therefore, the dyadic approaches to stroke care that recognise the interdependence of dyads are required.

The self-care interventions carried out in stroke survivor–caregiver dyads are delivered to both a patient and his/her informal caregiver with the expectations that both dyad members will be actively engaged in the stroke self-care of patients. Given the significance of dyadic perspectives in stroke self-care, it is essential to summarise the self-care interventions conducted in stroke survivor–caregiver dyads through meta-analysis (MA), thus identifying the most effective intervention in improving health-related outcomes for both stroke survivors and caregivers.

OBJECTIVES

The purpose of our review is to integrate the scientific evidence on dyadic self-care interventions for enhancing the health-related outcomes for stroke survivors and caregivers, which entails an understanding of the influencing factors in the success of interventions with health-related outcomes for stroke survivors and caregivers. Therefore, attempt will be made to answer the following research questions. First, what is the overall effectiveness of self-care interventions carried out in stroke survivor–caregiver dyads with health-related outcomes for stroke survivors, covering the QoL, self-efficacy, coping ability, emotional functioning, stroke self-care behaviours, social participation and ADL, and for caregivers, covering the QoL, self-efficacy, burden, coping ability and emotional functioning. Second, how do the effects of dyadic self-care interventions on health-related outcomes vary according to the characteristics of study design and the risk of bias? Finally, how do the effects of dyadic self-care interventions on health-related outcomes vary depending on the characteristics of intervention, such as the type of patient populations, intervention content, the combination of intervention components, intervention dose and the composition of interventionists?

METHODOLOGY

Our study will use a SR and MA, which will consider both randomised controlled trials and non-randomised trials (prospective and retrospective observational studies). The Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (PRISMA-P) statement has been used in the preparation of our protocol. And we will
follow the PRISMA guidelines and standard methods for conducting an SR and MA, while the random effect MA will be conducted to thoroughly evaluate the research results. In addition, exploratory moderator analysis will be carried out to account for heterogeneity.

**Selection criteria**

Our study aims to identify the studies published from inception to July 2021 of stroke survivor–caregiver dyads. According to the latest science papers on stroke self-care in survivor–caregiver dyads, the intervention included in our review is supposed to involve at least one of the following elements: (a) self-monitoring coping with the condition, (b) performing regular medication, exercising and other self-sustaining behaviours, (c) goal setting, (d) decision-making, (e) problem-solving, (f) an alternative method designed to facilitate behaviour change and improvements in physical and psychological functioning. The inclusion criteria applied to all papers are detailed as follows. The participants are aged 18 or over; dyad consists of a patient with stroke and at least one informal caregiver; both dyad members should be targeted for the intervention, present at the intervention outcomes (primary or secondary) from both OR either one of the participants measured; randomised controlled trials are conducted to include cross-over studies and cluster studies. What to be excluded from consideration includes meta-analyses and SRs, duplicative papers, case reports, opinion pieces, editorials and letters written to the editor. Besides, quasi-experiment trials or other intervention trials and the interventions that provided education or exercise only to participants will be excluded. In addition, we will exclude the studies of patients with cognitive impairment or institutionalised individuals because their stroke self-care is commonly managed by medical professionals. Additionally, we will include the studies under review, which may provide additional components about self-care interventions. Our study will include only those papers published in English or Chinese and conducted among human subjects only. However, we will also include non-English/Chinese studies if an English-language version of the abstract containing sufficient data is available for the calculation of effect size (ES).

**Data sources and search strategy**

We will conduct a systematic search of electronic database, including PubMed, Web of Science, CINAHL, PsycINFO, EMBASE, Cochrane Central Register of Controlled Trials and four Chinese databases (National Knowledge Infrastructure (CNKI), Chinese Biomedical Literature database (CBM), Wanfang Digital Periodicals (WANFANG) and Chinese Science and Technology Periodicals (VIP) database) from inception to July 2021, for the collection of relevant articles. To apply a variety of different search strategies is conducive to eliminating the risk of bias in MA results. To avoid the prospect of missing the eligible literature, we will conduct ancestry searches using the reference lists of eligible studies and conduct a review of the articles published over the past decade. Besides, we will search those unpublished studies and contact the experts in the field of stroke and self-care to identify any additional trials. The search strategy adopted for PubMed has been reviewed by the experts in different fields of stroke care, as listed in table 1. See online supplemental file 1 for full search strategies for all databases.

**Methodological quality assessment**

We will use the Cochrane Collaboration’s Risk for Bias tool to evaluate the risk of bias and the quality of the studies in the form of randomised controlled trials, as shown in table 2. Also, this tool allows each domain of potential bias to be classified either as ‘low risk’, ‘high risk’ or ‘unclear risk’. Apart from that, consideration will be given to methodological quality as an empirical question, the coding data about aspects of study quality and the potential risks of bias. Then, the data will be subjected to moderator analyses for establishing whether the potential risks of bias are significantly associated with the ES of the study. Two researchers will carry out this process independently, and any disagreement will be resolved through discussion and consultation with the third researcher, if necessary. The selection process will be presented as a PRISMA flow diagram (figure 1).

**Data collection and extraction**

We will use an EndNote V.X9 database to manage all citations from searches, so as to facilitate the screening of titles and abstracts. The two well-trained researchers (WW and BL) will screen a random sample of 50 studies in line with the study selection criteria to assess the consistency of literature selection. Once the interrater agreement measured by Kappa falls below 0.8, secondary training will be provided. The two researchers will screen all the initial studies independently if the agreement has been reached. Otherwise, the third researcher (YM) will intervene to resolve the disagreement between the first two reviewers. In the screening performed by the two well-trained researchers, titles and abstracts will be examined to evaluate whether the study involves an intervention purposed to improve an aspect of stroke self-care, with any possibly eligible citations marked for the retrieval of the entire paper. Any discrepancies will be discussed whenever necessary until the final studies are all identified.

These two researchers (WW and BL) will carry out data extraction independently. First, the data will be extracted from a random sample of 10 studies to verify the consistency of data extraction between WW and BL. If the level of agreement falls below 0.8, the causes will be explored and adjustment will be made. Data extraction will cover the following information: (a) study characteristics, including the first author’s name, publication year, country, title, studies design and settings, (b) participants, including the type of dyads, sample sizes, inclusion and exclusion criteria, sex ratio and average age, (c) methodological quality of trials, including the details on random process, blinding, dropout, reporting and
We will calculate descriptive statistics using SPSS V.21.0, so as to find out the basic characteristics of the included studies, study samples, interventions and others. When the results obtained from the same study are reported through different papers, they will be grouped as companion papers and the results will be reported as a single study. The risk ratio with 95% CI will be used to express the estimate of the effect for dichotomous outcome. When the same outcome is measured in different ways, the standardised mean difference with 95% CI will be used to indicate the magnitude of the intervention effect. The continuous and dichotomous data will be pooled using the inverse variance method and the Mantel-Haenszel method, respectively. The heterogeneity among the studies will be determined by means of $\chi^2$ test. When $p\geq0.1$ or $I^2\leq50\%$, the fixed effect model will be used to conduct MA. When $p<0.1$ or $I^2>50\%$, it is considered that there is statistical heterogeneity among the studies, and the causes of heterogeneity will be analysed. In case of clinical heterogeneity, subgroup analysis or sensitivity analysis will be conducted depending on its source. In case of no significant clinical heterogeneity, MA will be conducted using random effect model. We will conduct the MA with the assistance of Review Manager V.5.3 software. A narrative summary of the results obtained from individual studies will be made if an MA is not viable.

**Validity, reliability and rigour**

We will conduct our study according to the best practices in SR and MA methods. The results of SR and MA will be reported according to the PRISMA guidelines.

**Outcome measures**

We will conduct a series of meta-analyses on multiple health-related outcomes for both stroke survivors and caregivers. As for the primary outcomes of stroke survivors and caregivers, QoL will be measured using generic or condition-specific scales, such as the EQ5D and the Stroke-Specific Quality of Life Scale. Concerning the secondary outcomes of stroke survivors, self-efficacy, coping ability, emotional functioning, self-care behaviour, social participation and ADL will be included. The secondary outcomes of stroke caregivers include self-efficacy, burden, coping ability and emotional functioning.

**Statistical analysis**

We will calculate descriptive statistics using SPSS V.21.0, so as to find out the basic characteristics of the included studies, study samples, interventions and others. When the results obtained from the same study are reported through different papers, they will be grouped as companion papers and the results will be reported as a single study. The risk ratio with 95% CI will be used to express the estimate of the effect for dichotomous outcome. When the same outcome is measured in different ways, the standardised mean difference with 95% CI will be used to indicate the magnitude of the intervention effect. The continuous and dichotomous data will be pooled using the inverse variance method and the Mantel-Haenszel method, respectively. The heterogeneity among the studies will be determined by means of $\chi^2$ test. When $p\geq0.1$ or $I^2\leq50\%$, the fixed effect model will be used to conduct MA. When $p<0.1$ or $I^2>50\%$, it is considered that there is statistical heterogeneity among the studies, and the causes of heterogeneity will be analysed. In case of clinical heterogeneity, subgroup analysis or sensitivity analysis will be conducted depending on its source. In case of no significant clinical heterogeneity, MA will be conducted using random effect model. We will conduct the MA with the assistance of Review Manager V.5.3 software. A narrative summary of the results obtained from individual studies will be made if an MA is not viable.

**Validity, reliability and rigour**

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DISCUSSIONS

In our study, we will compare the efficacy of self-care intervention in stroke survivor–caregiver dyads, thus providing data on the overall effects and identifying the intervention components associated with the most significant improvement of outcomes for patients and/or caregivers. The results obtained from our study may indicate the direction of further studies on the development of self-care interventions for specific patient groups.

In our study, we will conduct analysis to determine the effect of dyadic self-care interventions on stroke survivor (including QoL, self-efficacy, coping ability, emotional functioning, stroke self-care behaviours, social participation and ADL) and/or caregiver outcomes (including QoL, self-efficacy, burden, coping ability and emotional functioning). This is expected to be the most extensive MA of self-care interventions in stroke survivor–caregiver dyads to date. Through our study, the entry point and components of most effective intervention for the
development of dyadic stroke self-care programme can be identified, which will provide a basis for developing the best method of developing objective measures and interventions with stroke self-care, so as to improve health outcomes for stroke survivors and caregivers. Furthermore, these significant findings will be vital to assisting policymakers and researchers in synthesising a large and complex literature.

Limitations
Since this is an understudied research area, it is likely that there will be very few studies identified in our review. Also, heterogeneity will arise in MA due to the variation in clinical and methodological characteristics. If it is necessary to amend our protocol considering the potential limitations as mentioned above in this study, the date of each amendment will be presented with a description of the change and corresponding rationale.

PATIENT AND PUBLIC INVOLVEMENT
There was no time allocated to patient and public involvement, particularly in the context of the current COVID-19 pandemic, so we were unable to involve patients. However, our SR protocol follows a standardised approach as per PRISMA-P guidelines.

ETHICS APPROVAL STATEMENT AND DISSEMINATION
We will exclude any identifiable patient data. Therefore, ethical approval and participant consent are not required. To conclude the study, we would like to publish our study in a peer-reviewed journal for the findings to be widely disseminated, and our abstract will be presented at suitable national/international conferences.

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Contributors
All authors have given approval to the final version for publication.

The following shows the contributor statement in our study: (1) Study concept and design: WW and BL. (2) Acquisition of data: WW, BL and YM. (3) Drafting of the manuscript: WW, BL, YM and BZ. (4) Critical revision of the manuscript for important intellectual content: all authors. (5) Supervision: ZZ.

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Competing interests
None declared.

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Not applicable.

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Supplemental material
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REFERENCES


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32 Hróbjartsson A, Bottrum I, Turner L. Assessing risk of bias in randomised clinical trials included in Cochrane reviews: the why is easy, the how is a challenge. Cochrane Database Syst Rev 2013;4:CD000058.


