Comparison of surgical strategies in patients with spontaneous intracerebral haemorrhage: a protocol for a network meta-analysis

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ABSTRACT

Introduction Spontaneous intracerebral haemorrhage (sICH) is a severe stroke subtype. The effective therapies for patients with sICH are still unclear, and the role of surgical treatment in sICH management is still controversial. Although some large trials did not show that surgery could benefit patients with sICH, some other studies suggested that some specific surgical strategies can have potential benefits to these patients. For a better understanding of the surgical treatment in patients with sICH, it is necessary to conduct a network meta-analysis to compare the effects of medical treatment and different surgical methods comprehensively.

Methods and analysis This protocol has been reported following the Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols. Related studies until August 2018 will be searched in the following databases: PubMed, Embase, Scopus, Web of Science, Cochrane Library, China National Knowledge Infrastructure (CNKI), VIP and Wanfang. Randomised controlled trials and non-randomised prospective studies comparing at least two different interventions in patients with sICH will be included. Quality assessment will be conducted using Cochrane Collaboration’s tool or Newcastle-Ottawa Scale based on their study designs. The primary outcome will be functional outcome and the secondary outcome will be mortality. Pairwise and network meta-analysis will be conducted using STATA V.14 (StataCorp, College Station, Texas, USA). Mean ranks and the surface under the cumulative ranking curve will be used to evaluate every intervention. Statistical inconsistence assessment, subgroup analysis, sensitivity analysis and publication bias assessment will be performed.

Ethics and dissemination Ethics approval is not necessary because this study will be based on publications. The results of this study will be published in a peer-reviewed journal.

PROSPERO registration number CRD42018112239

INTRODUCTION

Spontaneous intracerebral haemorrhage (sICH) is the second most common and the deadliest stroke subtype. However, the effective therapies for patients with sICH are still unclear, and the role of surgical treatment in sICH management is still controversial. Theoretically, surgical removal of haematoma can reduce mass effect and secondary brain injury after sICH. However, the International Surgical Trial in Intracerebral Haemorrhage (STICH), a large randomised controlled trial (RCT) including 1033 patients, did not find that early surgery could benefit patients with supratentorial sICH. In another study including 601 patients, named STICH 2, early surgery also could not improve outcome in patients with lobar sICH. In these two studies, the surgical methods were not restricted and most patients received craniotomy, although endoscopy and stereotaxy were used in some cases. However, different surgical methods have various effects on patients with sICH. Some other studies have been done to explore if specific surgical strategies can improve outcome in patients with sICH. Teernstra et al performed a multicentre RCT including 71 patients and found that stereotactic aspiration plus urokinase could be safely used in patients with sICH. Another
important RCT, Minimally Invasive Surgery and Alteplase for ICH Evacuation, which was conducted by Hanley et al., also showed that stereotactic aspiration plus alteplase was safe for haematoma removal in patients with sICH. Vespa et al enrolled 20 patients with sICH in their RCT, named Intraoperative Stereotactic Computed Tomography-Guided Endoscopic Surgery, and showed that early endoscopic surgery was safe in patients with sICH. The results of these studies have suggested that some specific surgical strategies may have potential benefits to patients with sICH.

Although some meta-analyses about surgical treatment in patients with sICH have been published previously, obvious limitations exist in these studies. For example, both stereotactic aspiration and endoscopic surgery were considered as minimally invasive surgery and compared with medical treatment or craniotomy in Scaggiante et al’s study. In Xia et al’s study, minimally invasive surgery including stereotactic aspiration and endoscopic surgery was only compared with craniotomy. In another study by Ye et al, only endoscopic surgery and craniotomy were analysed. According to its methodology, a network meta-analysis can assess the relative effectiveness of different therapies together and estimate the rank of these therapies. For a better understanding of the surgical treatment in patients with sICH, it is necessary to conduct a network meta-analysis to compare the effects of medical treatment and different surgical methods comprehensively.

OBJECTIVE
This study aimed to compare the efficacy and safety of medical treatment, craniotomy, stereotactic aspiration, endoscopic surgery and decompressive craniectomy in patients with sICH using Bayesian network meta-analysis.

METHODS AND ANALYSIS
Design
This study will be conducted using the Bayesian network meta-analysis. The protocol has been reported following the Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (see online supplement 1).

Registration information
This protocol has been registered on the International Prospective Register of Systematic Review (PROSPERO). The PROSPERO registration number is CRD42018112239.

Patient and public involvement
This protocol was designed for a network meta-analysis based on the existing literature. Therefore, the patients or the public were not involved.

Information source and search strategy
We will conduct the literature search for the related RCTs and non-randomised prospective studies until August 2018 in the following databases: PubMed, Embase, Scopus, Web of Science, Cochrane Library, CNKI, VIP and Wanfang. No restrictions on language will be set. The detailed search strategy is shown in the online supplement 2.

Eligibility criteria
Type of patients
This study will include adult patients diagnosed by CT or MRI. Studies about secondary intracerebral haemorrhage, primary intraventricular haemorrhage, subarachnoid haemorrhage or ischaemic stroke will not be included.

Type of studies
This study will include RCTs or non-randomised prospective studies. Retrospective studies, case reports, case series or reviews will not be included in this study.

Type of interventions
This study will include studies comparing at least two different interventions among the following interventions: medical treatment, craniotomy, stereotactic aspiration, endoscopic surgery and decompressive craniectomy. Studies will be excluded if there is no restriction for surgical methods.

Type of outcomes
The primary outcome will be functional outcome at the end of follow-up. Functional outcome will be dichotomised to good and poor according to the scale and threshold in each study. The secondary outcome will be mortality at the end of follow-up. The time point for outcomes will be the longest follow-up time in each study.

Study selection
After removing duplicate, titles and abstracts of all records will be screened by two authors (ZY and JZ) independently. Any record that does not meet the eligibility criteria will be removed. Full-text papers of the remaining studies will be obtained and screened by two authors independently. Only studies meeting the eligibility criteria will be finally included. If studies have duplicate data, only the study with larger sample size and longer follow-up time will be included. Any disagreement between two authors will be solved by another author (RG).

Data extraction
Based on a pre-established extraction form, two authors (ZY and JZ) will independently extract data from all included studies. The following information will be extracted: first author, year of publication, area, study duration, sample size, age, percentage of female, time from onset to surgery, inclusion/exclusion criteria, detailed intervention in each group, number of patients in each group, follow-up time and outcomes in each group. If some data cannot be obtained from the papers directly, we will try to contact the authors to obtain those data. Any disagreement between two authors will be solved by consensus and all data will be checked by another author (RG).
Risk of bias assessment
The quality of all RCTs will be assessed using Cochrane Collaboration’s tool. The quality of all non-randomised prospective studies will be assessed using Newcastle-Ottawa Scale. Two authors (ZY and JZ) will conduct quality assessment independently and any disagreement will be solved by discussion with another author (RG).

Data synthesis
When quantitative analysis cannot be conducted, we will narratively describe the results. If quantitative analysis is feasible, all of the following statistical analyses will be conducted using STATA V.14 (StataCorp, College Station, Texas, USA).

Direct comparisons of interventions
Conventional pairwise meta-analyses between different interventions will be first conducted if at least two studies provide relevant data. DerSimonian–Laird method and random effects model will be used.14 I² statistic will be used to evaluate heterogeneity among included studies.15

Indirect and mixed comparisons of interventions
Network meta-analysis will be performed with a random effects model reported in the previous study.16 Interactions among all included studies will be shown in the network geometry, and the contribution plot for the network will show the contributions of direct comparison.17 Mean ranks and the surface under the cumulative ranking curve will be used to evaluate every intervention for both functional outcome and mortality in patients with sICH.18

Statistical inconsistency assessment
Both global and local methods will be used to assess the inconsistency between direct and indirect comparison. For global method, the design-by-treatment model will be used.19 The loop-specific method will be adopted to assess the inconsistency locally.20

Subgroup analysis and sensitivity analysis
If possible, subgroup analyses will be performed based on age, gender, race, time from onset to surgery, Glasgow Coma Scale score, baseline haematoma volume and haematoma location. Sensitivity analysis will be performed to check the stability of the results by excluding each study.

Publication bias
Potential publication bias in the network meta-analysis will be assessed using a network funnel plot.

Quality of evidence
The evidence quality will be assessed following the Grading of Recommendations, Assessment, Development and Evaluation approach for rating the quality of treatment effect estimations from the network meta-analysis.21

DISCUSSION
This will be the first network meta-analysis that comprehensively compares different surgical strategies in patients with sICH. We will include non-randomised prospective studies to strengthen the statistical power of this network meta-analysis because the number of related randomised controlled studies is still limited. We hope the findings of this network meta-analysis can provide more information about the efficacy and safety of different surgical strategies in patients with sICH, which can help both clinical practice and study design in the future. However, limitations will still exist in this network meta-analysis. First, retrospective studies will be excluded considering their potentially low quality. However, exclusion of those studies will also increase potential publication bias. Moreover, some included studies may have inferior quality, which decreases the significance of the results in this network meta-analysis. Furthermore, potentially high heterogeneity among different studies may influence the final results of this network meta-analysis.

Ethics and dissemination
Ethical issues
Neither ethics approval nor patient consent is necessary because this network meta-analysis will be based on publications.

Publication plan
This protocol has been successfully registered on PROSPERO. The final results of this study will be published in a peer-reviewed journal.

Contributors
ZY, JZ and HL designed this protocol. ZY and JZ tested the feasibility of this protocol. The evidence quality will be assessed using Cochrane Collaboration’s tool. The quality of all non-randomised prospective studies will be assessed using Newcastle-Ottawa Scale. Two authors (ZY and JZ) will conduct quality assessment independently and any disagreement will be solved by discussion with another author (RG).

Project of this study will be assessed using Newcastle-Ottawa Scale. Two authors (ZY and JZ) will conduct quality assessment independently and any disagreement will be solved by discussion with another author (RG).

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REFERENCES


