Efficacy and safety of furosemide for prevention of intradialytic hypotension in haemodialysis patients: protocol for a multicentre randomised controlled trial

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ABSTRACT

Introduction Intradialytic hypotension (IDH) is a frequent and serious complication of maintaining haemodialysis (HD) patients and associated with subsequent cardiovascular events and higher mortality. Furosemide is commonly used in non-dialysis chronic kidney disease patients and can effectively manage the volume and blood pressure. However, these agents are often discontinued on initiation of dialysis. Two large observational studies have demonstrated that furosemide can lower the rate of IDH episodes. However, there is still no randomised controlled trial (RCT) to investigate the efficacy and safety of furosemide for prevention of IDH in HD patients. The purpose of this study was to assess the efficacy of furosemide in reducing IDH in HD patients with residual renal function.

Methods and analysis A two-arm, parallel, multicentre RCT will be conducted at 12 hospitals in China. An estimated sample of 560 HD patients will be recruited. Eligible patients will be randomly assigned to treatment group (patients receive oral furosemide 80 mg/day; after a 2-week treatment, if their urine volume is less than 400 mL/day, the dose of furosemide is adjusted to 160 mg/day) and blank control group via a central randomisation system using 1:1 ratio. The primary outcome is the occurrence of IDH. Outcome assessors and data analysts will be blinded and participants will be asked not to reveal their allocation to assessors. The outcome analyses will be performed both on the intention-to-treat, which includes all patients randomised, and per-protocol population, which includes eligible patients who adhere to the planned treatment and follow-ups.

Ethics and dissemination The trial protocol has been approved by the Biomedical Research Ethics Committee of West China Hospital of Sichuan University (2019.385). Results will be presented at national and international conferences and published in peer-reviewed journals.

Trial registration number ChiCTR2000039724.

INTRODUCTION

Intradialytic hypotension (IDH) is a common and serious complication of haemodialysis (HD) patients, with an incidence of up to 20%–30% of all dialysis sessions.1 The frequent occurrence of IDH increases the risk of thrombosis in vascular access,3 inadequate dialysis,4 cardiovascular disease5 and mortality.3 6 7 An absolute nadir intradialytic systolic blood pressure (SBP) less than 90–100 mm Hg was most potently associated with increase of about 30% mortality for patients with all range of predialysis blood pressure.3

A series of factors may drive IDH through multiple mechanistic pathways. When the rapid removal of a large volume of blood water and solute overwhelmed normal physical compensatory mechanisms during dialysis therapy, the circulating volume and plasma refilling reduced, and followed, blood pressure.1 Based on the multifactorial pathophysiology of IDH, a number of strategies had been developed to reduce the frequency and severity of IDH, including the modulation of ultrafiltration (UF),8 qualitative changes in dialysate composition9 and lowering of dialysate temperature,10 however, all of these had a limited effect.11 No other effective pharmacological approach is recommended to address IDH except adjusting antihypertensive drugs.

Furosemide, a common oral diuretic, is widely prescribed in chronic kidney disease (CKD) for managing volume and blood pressure. As a loop diuretic, furosemide increases...
the excretion of water, sodium and chlorine by inhibiting sodium and chloride reabsorption at the level of the thick ascending limb of the loop of Henle.\textsuperscript{12} Nonetheless, the role of diuretics in advanced CKD is not fully understood. Once patients start dialysis, it is often believed that dialysis can manage fluid overload and the use of diuretics is discontinued. Indeed for HD patients with residual renal function, diuretics can promote urine output, stabilise volume control, avoid ultrafiltration and thus reduce hypotension during dialysis.\textsuperscript{12}

The effect of furosemide in dialysis has been investigated in several observational studies. Recently, one large observational study enrolling 12971 HD patients showed that loop diuretics can reduce dialysis hypotension significantly compared with control group (RR 0.95, 95% CI, 0.92 to 0.99).\textsuperscript{13} These results were consistent with the previous Dialysis Outcome and Practice Pattern Study (DOOPS), which reported that diuretics reduce 45% hypotensive episodes and furosemide was the mostly widely used loop diuretic.\textsuperscript{14} Accumulating evidence showed that the use of loop diuretics can increase urine volume and sodium excretion, decrease the rate of weight gain in dialysis and reduce the risk of hospitalisation.\textsuperscript{15−18} while some other studies have shown that furosemide failed to delay kidney function decline or reduce mortality in dialysis patients.\textsuperscript{16,17} Unfortunately, prospective randomised studies evaluating furosemide therapy for IDH are lacking.

In this study, we aim to conduct a prospective multi-centre randomised controlled trial (RCT) to primarily examine if furosemide can reduce IDH in HD patients with residual renal function.

**METHODS AND ANALYSIS**

We followed the Standard Protocol Items: Recommendations for Interventional Trials statement for the reporting of our trial protocol.\textsuperscript{18}

**Study design and setting**

A two-arm, parallel, blank-control, multicentre RCT evaluating the efficacy and safety of furosemide for prevention of IDH in patients will be conducted at West China Hospital of Sichuan University and other eleven dialysis centres in China. All the study centres can perform in-centre HD three times a week. We plan to recruit participants at each HD unit from June 2021 by the nephrologist and expect to end in March 2022. The eligible and consented patients will be randomly assigned to receive 6 months of furosemide treatment (n=280) or blank control group (n=280) at a 1:1 ratio. The intervention between groups, including dialysis prescription and artificial kidney machines, will not be altered during the study period. All participants will be followed at least 1 year. The frequency of IDH episodes and other outcomes will be assessed at each HD session. This study has been approved by the biomedical research ethics committee, West China Hospital of Sichuan University (2019.385). The study will be conducted in line with the declaration of Helsinki. A flow chart of the trial procedure is summarised in figure 1.

**Participants**

All participants aged 18 years or older and diagnosed with end-stage renal disease according to the Kidney Disease: Improving Global Outcomes guideline\textsuperscript{19} at each HD unit will be assessed for eligibility. The inclusion criteria are as follows: participants who have been treated with HD three times a week for more than 3 months, with residual renal function of more than 200 mL urine output/day (based on a 24-hour urine collection, measured by a 200 mL measuring cup)\textsuperscript{14} and consent to participate in the study. Participants with SBP ≤100 mm Hg, a history of liver cirrhosis or concurrent liver impairment, experienced congestive heart failure (ejection fraction ≤50%), acute myocardial infarction or stroke in the preceding 6 months, used any loop diuretics in the preceding 1 month, planned to receive a kidney transplant during the study period, or any other condition that the research staff judged as hard to complete or comply with or otherwise unsuitable for the study will be excluded.

The screening will be conducted by the research nephrologist through necessary physical examination and clinical tests. Eligible patients will be informed clearly of the aim and procedure of the research both verbally and via an information sheet. A consent form is needed to be completed and signed by each individual who agrees to participate before enrolment. Information of all the patients will keep confidential and they will have the right to withdraw without prejudice at any time.

**Randomisation**

Eligible and consented patients will be randomly assigned to receive 6 months of furosemide treatment (n=280) or blank control group (n=280) at a 1:1 ratio via a central computer-generated randomisation sequence. We will use a block randomisation in block sizes of 4 or 6 to keep balance between groups. The randomised sequence will be stored by the study coordinator and keep secret from the research nephrologist who recruits and evaluates the patients. Only after the informed consent is obtained, the allocation information will be provided to the patient’s HD doctor, who will implement the intervention according to the study protocol. The study coordinator must ensure that the informed consent form has been obtained from each participant before randomisation to avoid selection bias.

**Interventions**

Following enrolment, participants will undergo a 1-week baseline period of comprehensive clinical assessment, including dry weight reduction, medication review and standardisation of their dialysis prescription. After randomisation, patients in both groups will receive HD three times a week over a 6-month treatment period. The dialysis prescription, dialysate and artificial kidney machines will
Assessed for eligibility

Excluded
- Not meeting inclusion criteria
- Refuse to participate
- Other reasons

Randomisation (n=560)

Furosemide treatment group (n=280):
patients receive oral furosemide 80mg/day, after
two-week treatment, if their urine volume less
than 400mL/day, the dose of furosemide adjust
to 160 mg/day

Blank control group (n=280):
no furosemide treatment

Follow up one year
Primary outcome: intradialytic hypotension
Secondary outcomes: hospitalization, all-cause
mortality, cardiac mortality, cardiovascular events,
intradialytic weight gain, adverse events.

Figure 1  The summarised design of the trial.

not be altered during the treatment period. UF volumes
will be adapted to reach <5% of dry weight during each
HD session. Dry weight will be determined clinically by the
patient’s attending nephrologist. The HD nurse assigned
to each individual participant will record haemodynamic
parameters, treatment parameters and IDH-related inter-
ventions on standard clinical HD run-sheets.

Treatment group and blank group

Eligible patients will be randomly assigned to treatment
group or blank group. The patients in the treatment
group will receive oral furosemide 80 mg/day. After
a 2-week treatment, if their urine volume was less than
400mL/day, the dose of furosemide will be adjusted to
160 mg/day. Furosemide treatment group will last for 6
months. Once the patient’s daily urine output is less than
200mL, the use of furosemide will be discontinued. The
patients in the blank group will receive no intervention,
just as usual. Both groups will be followed up for 1 year.

Primary outcome and definition of ‘IDH’

The primary outcome measures the rate of symptomatic
IDH, and any IDH-related nursing interventions to treat
hypotension episodes during each dialysis session. The
definition of IDH is SBP <90 mm Hg (among patients
with predialysis SBP <160 mm Hg) or SBP <100 mm Hg
(among patients with predialysis BP ≥160 mm Hg). Blood
pressure will be measured before dialysis, every
30 min during HD and after dialysis in each HD session
of the study. IDH-related interventions will be defined as
the use of the Trendelenburg position, manual reduction
of UF rate, infusion of isotonic saline or hypertonic fluid,
lowering of dialysate temperature or dialysis cessation.
The number of symptomatic IDH episodes along with the
duration of each dialysis treatment will be captured. The
rate of IDH for each session will be calculated by dividing
the number of episodes by the duration of the session
in hours. Hypotensive events and symptoms (headache,
cramps, nausea and vomiting) will be recorded and anal-
ysed as both the number of episodes and time of occur-
rence since the beginning of the HD session.

Secondary outcomes

The secondary outcomes are hospitalisation, all-cause
mortality, cardiac mortality, cardiovascular events, inter-
dialytic weight gain, dialysis symptoms and any adverse
events. These data will be collected for all periods of
the study. The outcome measurement time points are
provided in detail in table 1.
Patient safety
Any adverse events (described as unfavourable or unintended signs, symptoms or diseases occurring after treatment) related to furosemide therapy will be observed and reported by patients and practitioners during each dialysis session. In addition, all vital signs and adverse events will be measured and recorded at each dialysis session.

Blinding
Study investigators, nephrologists and participants will be aware of the treatment allocation. HD nurses, outcome assessors and data analysts will be blinded and participants will be asked not to reveal their allocation to assessors. In addition, we will blind interpretation of the study results to minimise misleading data interpretation.

Sample size determination
Our primary study hypothesis is that furosemide treatment would achieve more reduction in the rate of IDH than the blank control group in HD patients. According to the previous study, the rate of IDH was 0.25, compared with the control group, or that furosemide could reduce the rate of IDH was 0.55. Sample size calculations were conducted using G-Power V3.1, with an $\alpha$ value of 0.05 and power of 80%. The relative risk of IDH for treatment group relative to controls is 0.6, and a sample size of 250 patients per arm is required. The dropout rate of furosemide treatment during the study was estimated to be 10%, so a minimum sample size of 560 patients will be needed in each group.

Monitoring
We will develop an independent data monitoring committee (DMC), responsible for the monitoring the quality and regulatory compliance of the trial, as well as ensuring the safety of participating patients. The DMC will consist of six members with expertise in nephrology,

| Table 1 | Timing of visits and data collection |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|               | Screening 1 week | Baseline period 0 week | Treatment period 1 week | Follow-up period 1 year |
| Patient        |                 |                 |                 |                 |
| Eligibility    | X               |                 |                 |                 |
| Informed consent | X              |                 |                 |                 |
| Demographics and medical history | X |                 |                 |                 |
| Physical examination | X |                 |                 |                 |
| Type of vascular access | X |                 |                 |                 |
| Charlson comorbidity index score | X |                 |                 |                 |
| Randomisation  |                 | X               |                 |                 |
| Intervention   |                 |                 |                 |                 |
| Treatment group |                 |                 |                 |                 |
| Blank group    |                 |                 |                 |                 |
| Comparison     |                 |                 |                 |                 |
| outcomes       |                 |                 |                 |                 |
| Hypotension episodes | X | X               |                 |                 |
| Blood pressure | X               | X               |                 |                 |
| IDH-related nursing interventions | X | X               |                 |                 |
| Dialysis symptoms | X | X               |                 |                 |
| Interdialytic weight gain | X | X               |                 |                 |
| Hospitalisation |                 | X               | X               |                 |
| All-cause mortality | X | X               |                 |                 |
| Cardiac mortality | X | X               |                 |                 |
| Cardiovascular events | X | X               |                 |                 |
| Episode of frank (or suspected) sepsis | X | X               |                 |                 |
| Participant safety |             |                 |                 |                 |
| Adverse effects | X               | X               | X               | X               |

IDH, intradialytic hypotension.
dialysis management, dialysis care, trial methodology and biostatistics. The DMC meeting will be held once in 3 months; at the request of DMC, the meeting may take place every 2 months. We will develop a procedural document for the DMC meeting, and strictly follow the document.

**Patient and public involvement**
No patient involved.

**Statistical analysis**
Group comparisons will be undertaken using \( \chi^2 \) tests for categorical characteristics, and either analysis of t-test or Wilcoxon rank sum tests for continuous variables. The outcome analyses will be performed both on the intention-to-treat, which includes all patients randomised, and per-protocol population, which includes eligible patients who adhere to the planned treatment and follow-up.

**Ethics and dissemination**
The trial protocol has been approved by the Human Research Ethics Committee of West China Hospital of Sichuan University and other 11 dialysis centres in China. Participants were informed of the aims and nature of the research both verbally and via an information sheet. They were requested to complete an informed written consent before enrolment. Confidentiality of the information provided by the participants and their right to withdraw without prejudice are enforced throughout the study duration.

All individual data are securely stored, password protected and accessible by the research team only. The results of the trial will be presented at national and international scientific conferences and be submitted for publication in peer-reviewed journals.

**DISCUSSION**
This trial is expected to provide convincing evidence about the preventive effect of furosemide therapy for IDH. Although number of strategies have been developed to reduce the frequency and severity of IDH, all of those are difficult to conduct and hard to promote in clinical practices. To our knowledge, no effective pharmacological approach is recommended to address IDH except adjusting antihypertensive drugs.

Episodes of IDH can cause serious adverse events (renal ischaemia, cardiovascular disease and mortality), while the use of diuretics in HD patients has the potential to improve interdialytic fluid status. Diuretics are commonly prescribed for hypertension and volume management in CKD patients before dialysis. Currently, many clinicians discontinue these medications when individuals start dialysis, and diuretics are used infrequently in dialysis patients with residual renal function, studies in China report diuretic use between 6% and 29%.\(^{21,22}\) and the dosing is also variable, with daily furosemide dose of 40–80 mg reported in China.\(^{21,22}\) Continuation of loop diuretics after HD initiation was associated with lower rates of hospitalisation and IDH as well as lower interdialytic weight gain during the first year of dialysis.\(^{13}\)

Based on published observational studies, the use of loop diuretics can increase urine volume and sodium excretion, decrease the rate of weight gain in dialysis and reduce the risk of hospitalisation.\(^{13-15}\) Among them, furosemide proved to be effective for prevention of IDH in HD patients.\(^{13-15}\) However, to our knowledge, prospective randomised studies evaluating furosemide therapy for IDH are lacking. Considering the heavy burden of IDH, there is a call for validation of the efficacy of this cheap and convenient strategy. Through our carefully designed RCT, we will offer trustworthy evidence for the effects of furosemide for IDH.

The major concern about the development of ototoxicity and other side effect of loop diuretics in dialysis patients hindered its use in HD.\(^{12}\) Another possible reasons for discontinuation of the use of loop diuretics include the assumption that dialysis treatment alone is sufficient for management of fluid overload and under-estimation of its benefit for HD patients. A recent systematic review has found that loop diuretics may benefit HD patients by reducing the incidence rate of IDH, all-cause mortality and cardiovascular mortality for HD patients.\(^{23}\) However, evidence about its safety is still limited. The present prospective study will confirm the efficacy and gain evidence about the safety of furosemide in HD and further guide the clinical practices.

There are several limitations of this study. First, we did not administer a placebo control and dose contrast. Thus, the placebo effect may not be well parcelled out. However, it is difficult to get the placebo medicine, which is similar to furosemide in our study. Furosemide has some side effects, and blood pressure of patients vary, so it is not appropriate to administer dose contrast considering the limited sample sizes. For the safety of patients, the study gives a safe dose. Other assessments, including placebo and dose control with IDH, will be considered in the future study. Second, the definition of IDH is objective. In contrast, HD-related symptom frequency was much higher in a survey,\(^{21}\) but patients experience symptoms at varying thresholds of BP change and nadir, so it is difficult to recognise the symptom of IDH. However, hypotensive symptoms (headache, cramps, nausea and vomiting) will be recorded and analysed in our study. Our methods for recruitment, randomisation, allocation, outcome assessment and data collection methods have been carefully designed to minimise bias.

In summary, based on evidence from observational studies, furosemide seems to be a promising treatment for prevention of IDH in HD patients. Despite all the potential advantages of the use of diuretics in maintaining HD patients with residual renal function, furosemide is used infrequently in China and most countries. Our study will gain evidence from RCT and provide an effective way for reducing IDH in maintaining HD patients. Since we adopted a relatively low and safe dose of furosemide, more
prospective studies investigating the ideal dose, especially dose based on the residual kidney function of individual are needed. Further studies are expected to offer trustworthy evidence for the effects of furosemide and stimulate furosemide to be better used in HD patients despite numerous potential benefits and few side effects.

In summary, we will conduct a definitive multicentre RCT to assess the efficacy and safety of furosemide for prevention of IDH in HD patients. We will use rigorous methods to minimise bias and set up several working committees to ensure quality conduct of the trial. Our study will gain evidence from RCT and provide an effective way for reducing IDH in maintaining HD patients. Further studies are expected to offer trustworthy evidence on the effects of furosemide and stimulate furosemide to be better used in HD patients despite numerous potential benefits and few side effects.

Contributors WC, FW, YZ and LZ conceived and designed the study. WC and FW drafted the manuscript. LZ, YZ, ZC and MD critically revised the manuscript. All authors have read and approved the manuscript.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were involved in the design, or conduct, or reporting, or dissemination plans of this research. Refer to the Methods section for further details.

Patient consent for publication Parental/guardian consent obtained.

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