

# The effect of mindfulness meditation on sleep quality and negative emotions in stroke patients with heart disease

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<b>Registration date</b> 24/06/2024	<b>Overall study status</b> Completed	<input type="checkbox"/> Protocol
<b>Last Edited</b> 24/06/2024	<b>Condition category</b> Mental and Behavioural Disorders	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
		<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Sleep is one of the most fundamental and essential physiological needs for humans. It plays a crucial role in the recovery of the body, the integration and consolidation of memory, and is an indispensable component of health. Neuroscientific research indicates that sleep is a state required for maintaining highly physiological functions, adaptive behaviors, and biological defense mechanisms. Sleep significantly affects both physical and mental health, as well as the onset and progression of various diseases. Mindfulness meditation, as an emerging non-pharmacological therapy, has garnered considerable research attention in the field of insomnia in recent years and is considered an effective adjunctive method for improving sleep. This study aims to observe the effects of combining oral eszopiclone with mindfulness meditation on insomnia in stroke patients with concurrent coronary heart disease through a prospective randomized controlled clinical trial, in hopes of providing a more advantageous reference for rehabilitation treatment of such patients in clinical settings.

### Who can participate?

Stroke patients with concurrent coronary heart disease whose sleep quality and negative emotions are currently affected

### What does the study involve?

Participants will be randomly allocated to two groups that will receive routine treatment, including medication therapy and conventional rehabilitation training. Medication Therapy includes antiplatelet medication, lipid-lowering and plaque-stabilization medication, blood glucose control, blood pressure management, and neurotrophic medication. Conventional Rehabilitation Training: this is tailored according to the functional impairments of the patients and includes limb motor function training in stroke patients, activities of daily living (ADL) training, and related physical therapies.

For the control group, in addition to the standard treatment, eszopiclone tablets (Disha Pharmaceutical Group Co., Ltd., approval number H20213830, 3 mg/tablet) will be administered orally before bedtime, at a dosage of 3mg once daily.

In the intervention group, mindfulness meditation training is added to the treatment regimen

based on the control group's therapy. The intervention is divided into three stages, with the mindfulness meditation protocol developed according to relevant literature.

What are the possible benefits and risks of participating?

Mindfulness meditation can effectively improve sleep quality and reduce negative emotions in stroke patients with concurrent coronary heart disease, as well as promote the recovery of limb functions. There is little evidence about the potential harmful effects of mindfulness meditation. But caution is needed.

Where is the study run from?

The Second Hospital of Hebei Medical University

When is the study starting and how long is it expected to run for?

May 2021 to April 2024

Who is funding the study?

Health Commission of Hebei Province 2022 Hebei Province Medical Science Research Project

Who is the main contact?

Guangxiao Ni, kll51x6g4@163.com

## Contact information

### Type(s)

Public, Scientific, Principal investigator

### Contact name

Dr Guangxiao Ni

### ORCID ID

<https://orcid.org/0009-0007-1984-3875>

### Contact details

The first Department of Rehabilitation, the Second Hospital of Hebei Medical University

No. 215 Heping West Road, Xinhua District

Shijiazhuang City

China

050000

+86 15230161010

kll51x6g4@163.com

## Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### Protocol serial number

2022 Hebei Province Medical Science Research Project number: 20221008

## Study information

**Scientific Title**

Efficacy of mindfulness meditation on sleep quality and negative emotions in stroke patients with concurrent coronary heart disease

**Acronym**

Effect of Mindfulness Meditation on Stroke Patient

**Study objectives**

To investigate the efficacy of mindfulness meditation on sleep quality and negative emotions in stroke patients with concurrent coronary heart disease.

**Ethics approval required**

Ethics approval required

**Ethics approval(s)**

approved 20/05/2021, Research Ethics Committee of The Second Hospital of Hebei Medical University (No. 215 Heping West Road, Xinhua District, Shijiazhuang City, 050051, China; +86 66002811; kll51x6g4@163.com), ref: 2021-R296

**Study design**

Interventional randomized controlled trial

**Primary study design**

Interventional

**Study type(s)**

Treatment

**Health condition(s) or problem(s) studied**

Sleep quality and negative emotions in stroke patients with concurrent coronary heart disease

**Interventions**

Eligible stroke patients with concurrent coronary heart disease at the Second Hospital of Hebei Medical University will be selected. The patients will be randomly divided into an intervention group and a control group. The control group will receive routine treatment plus eszopiclone tablets, while the intervention group will receive mindfulness meditation in addition to the treatment given to the control group.

Routine treatment included medication therapy and conventional rehabilitation training. Medication Therapy included: This included antiplatelet medication, lipid-lowering and plaque-stabilization medication, blood glucose control, blood pressure management, and neurotrophic medication. Conventional Rehabilitation Training: This was tailored according to the functional impairments of the patients and included limb motor function training in stroke patients, activities of daily living (ADL) training, and related physical therapies.

For the control group, in addition to the standard treatment, eszopiclone tablets (Disha Pharmaceutical Group Co., Ltd., approval number H20213830, 3 mg/tablet) were administered orally before bedtime, at a dosage of 3mg once daily.

In the observation group, mindfulness meditation training was added to the treatment regimen based on the control group's therapy. The intervention was divided into three stages, with the mindfulness meditation protocol developed according to relevant literature.

1. Introduction Stage (Week 1)
2. Maintenance Stage (Week 2)
3. Consolidation Stage (Weeks 3 to 6)

Participating patients were required to practice mindfulness meditation for 45 minutes each night before bed, record the time and their feelings after completing the practice, and continue this training for 6 weeks, with focused feedback and Q&A sessions in a WeChat group every Sunday. After 6 weeks of treatment, participants from both groups were to gradually reduce and then cease using eszopiclone within a maximum of 2 weeks.

Data will be collected before the intervention, and/or after 6 and 12 weeks of the intervention, including general information, Pittsburgh Sleep Quality Index (PSQI), Self-Rating Anxiety Scale (SAS) scores, Self-Rating Depression Scale (SDS) scores, and Fugl-Meyer Assessment (FMA) scores. Demographic data comprises gender, age, BMI, disease duration, education level, stroke location, stroke region, marital status, place of residence, limb disability, hypertension, diabetes, and per capita monthly income.

The PSQI will be used to assess the patient's sleep quality. The total score ranges from 0 to 21, with scores >7 indicating sleep disturbances. The higher the score, the poorer the sleep quality. The SDS and SAS were used to evaluate the visitor's depression and anxiety levels. An SDS score > 72 indicates severe depression; an SDS score of 63-72 suggests moderate depression; an SDS score of 53-62 indicates mild depression; an SDS score <53 signifies the absence of depression. A SAS score >69 denotes severe anxiety that requires immediate referral to a professional psychology institution; a SAS score of 60-69 indicates moderate anxiety, with the testee often feeling anxious recently, yet generally able to self-regulate; a SAS score of 50-59 suggests mild anxiety, with occasional anxious experiences recently, but with minor symptoms that usually quickly resolve after timely adjustment; a SAS score <50 signifies the absence of anxiety, considered a normal state. Motor function will be assessed using the FMA, including the Upper Limb Motor Function Assessment (comprising 33 items, with a total score of 66) and the Lower Limb Motor Function Assessment (comprising 17 items, with a total score of 34). A higher score indicates better motor function of the affected limb.

### **Intervention Type**

Mixed

### **Primary outcome(s)**

The following primary outcome measures will be assessed before the intervention, and/or after 6 and 12 weeks of the intervention:

1. Sleep quality measured using the Pittsburgh Sleep Quality Index (PSQI)
2. Anxiety measured using the Self-Rating Anxiety Scale (SAS)
3. Depression measured using Self-Rating Depression Scale (SDS)
4. Motor function measured using the Fugl-Meyer Assessment (FMA)

### **Key secondary outcome(s)**

There are no secondary outcome measures

### **Completion date**

30/04/2024

# Eligibility

## Key inclusion criteria

1. First onset of ischemic stroke confirmed via imaging, with a prior clear diagnosis of concurrent coronary heart disease
2. Presence of insomnia that appeared either after the stroke or within one month before the stroke onset, with a Pittsburgh Sleep Quality Index (PSQI) score of  $\geq 7$
3. Self-Rating Anxiety Scale (SAS) score of  $\geq 50$  and Self-Rating Depression Scale (SDS) score of  $\geq 53$
4. Clear consciousness, normal cognitive function, and ability to cooperate with treatment
5. Voluntary signing of informed consent by the patient and their family for participation in this trial

## Participant type(s)

Patient

## Healthy volunteers allowed

No

## Age group

Mixed

## Sex

All

## Total final enrolment

72

## Key exclusion criteria

1. Aphasia, abnormal mental behavior post-AIS, or severe consciousness disorders
2. Existence of unstable angina or myocardial infarction within the past two weeks
3. Significant organ dysfunction
4. Patients with malignant tumors
5. Existence of sleep disorders for more than one month before the stroke onset, or sleep disorders caused by medications or other systemic diseases
6. History of allergy to eszopiclone or similar medications

## Date of first enrolment

01/05/2023

## Date of final enrolment

20/04/2024

# Locations

## Countries of recruitment

China

**Study participating centre**  
**The Second Hospital of Hebei Medical University**  
No. 215 Heping West Road, Xinhua District  
Shijiazhuang City  
China  
050000

## Sponsor information

**Organisation**  
Second Hospital of Hebei Medical University

**ROR**  
<https://ror.org/015ycqv20>

## Funder(s)

**Funder type**  
Government

**Funder Name**  
Health Commission of Hebei Province

**Alternative Name(s)**  
Hebei Provincial Health Commission,

**Funding Body Type**  
Government organisation

**Funding Body Subtype**  
Local government

**Location**  
China

## Results and Publications

**Individual participant data (IPD) sharing plan**  
The datasets generated and/or analysed during the current study will be available upon request from Guangxiao Ni (kll51x6g4@163.com)

**IPD sharing plan summary**

Available on request