

Clinical Trial Protocol

Iranian Registry of Clinical Trials

11 Sep 2026

Evaluating the effect of the comprehensive drug protocol for spasticity in cerebral palsy patients in terms of gait analysis, clinical examination, radiographic changes, complications and quality of life.

Protocol summary

Study aim

Determining the effectiveness of a comprehensive drug treatment protocol for spasticity in patients with cerebral palsy. Analysis of gait analysis, clinical examination, radiographic examination, drug side effects, quality of life.

Design

single arm group randomized trial, phase 2, group design of 31 patients

Settings and conduct

This study will be conducted in the orthopedic clinics of Tehran University of Medical Sciences on spastic cerebral palsy patients who have been included in the study according to the inclusion and exclusion criteria, based on a drug therapy protocol.

Participants/Inclusion and exclusion criteria

Patients with spastic cerebral palsy with or without a history of previous drug treatment and without a history of surgery within the past year.

Intervention groups

In this study, the effectiveness of drug therapy in cerebral palsy patients who were included in the study according to the inclusion and exclusion criteria will be examined according to the following protocol. Choice of treatment type: -Generalized spasticity treatment 1-First line: Baclofen (maximum dose 2.5mg/kg/day), Tizanidine (4mg/kg/day) 2-Second line: Dantrolene (12mg/kg/day), Diazepam (0.01-0.3mg/kg/day) 3-Third line (intraspinal baclofen (50 micrograms on the first day, 75 micrograms on the second day, 100 micrograms on the third day) - Treatment with local aim (both in generalized and local spasticity) 1-Botulinum toxin (upper, lower limbs): 30U/kg After giving the above drugs to each of the patients with the specified dosage in the first three visits, 3 and 6 months later, the patients are examined with examination and radiology and the results of the study are recorded

Main outcome variables

spasticity :Range of motion:drug side effect

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20241215064059N1**

Registration date: **2025-01-04, 1403/10/15**

Registration timing: **prospective**

Last update: **2025-01-04, 1403/10/15**

Update count: **0**

Registration date

2025-01-04, 1403/10/15

Registrant information

Name

Mohammad Amin Choobdar

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 11 4424 4601

Email address

dr.amin.choobdar@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2025-01-17, 1403/10/28

Expected recruitment end date

2025-02-16, 1403/11/28

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date
empty

Scientific title
Evaluating the effect of the comprehensive drug protocol for spasticity in cerebral palsy patients in terms of gait analysis, clinical examination, radiographic changes, complications and quality of life.

Public title
Investigating the effect of drug treatment protocol in cerebral

Purpose
Treatment

Inclusion/Exclusion criteria
Inclusion criteria:
Patients with spastic cerebral palsy with or without a history of previous drug treatment without a history of surgery within the past year
Exclusion criteria:
Patients who have not returned for a follow-up visit during this period Patients who have had surgery within the past year

Age
No age limit

Gender
Both

Phase
2

Groups that have been masked
No information

Sample size
Target sample size: 31

Randomization (investigator's opinion)
N/A

Randomization description

Blinding (investigator's opinion)
Not blinded

Blinding description

Placebo
Not used

Assignment
Single

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee
Name of ethics committee
Ethics committee of tehran University of Medical Sciences
Street address
south Eskandari,Shokoufe ave
City
Tehran

Province
Tehran
Postal code
1319764853
Approval date
2024-06-18, 1403/03/29
Ethics committee reference number
IR.TUMS.IKHC.REC.1403.222

Health conditions studied

1

Description of health condition studied
spastic cerebral palsy
ICD-10 code
G80
ICD-10 code description
Cerebral palsy

Primary outcomes

1

Description
spasticity
Timepoint
The beginning of the study, 3 and 6 months later
Method of measurement
Ashworth scale

2

Description
range of motion
Timepoint
The beginning of the study, 3 and 6 months later
Method of measurement
Goniometer

3

Description
drug side effect
Timepoint
The beginning of the study, 3 and 6 months later
Method of measurement
Medication side effects questionnaire

Secondary outcomes

empty

Intervention groups

1

Description
In this study, the effectiveness of drug therapy in cerebral palsy patients who were included in the study according to the inclusion and exclusion criteria will be examined according to the following protocol. Selection

of treatment type: - Treatment of generalized spasticity
 1- First line: Baclofen (maximum dose 2.5 mg/kg/day), Tizanidine (4 mg/kg/day) 2- Second line: Dantrolene (12 mg/kg/day), Diazepam (0.01-0.3 mg/kg/day) 3- Third line (intraspinal baclofen (50 micrograms on the first day, 75 micrograms on the second day, 100 micrograms on the third day) - Treatment with local aim (both in generalized and local spasticity) 1- Botulinum toxin {Dysport, Masport, Dyston} (upper, lower limbs): 30 U/kg
 Treatment based on specific conditions - Pain (botulinum toxin, Tizanidine, oral baclofen, Diazepam, Dantrolene, intraspinal baclofen) - Disorder Sleep (tizanidine, oral baclofen, diazepam) - mixed type with dystonia (botulinum toxin, oral baclofen, diazepam) - post-operative pain due to spasm (diazepam, botulinum toxin, oral baclofen) - seizures (tizanidine, diazepam * Caution regarding the use of baclofen, dantrolene and intrathecal baclofen; After giving the above drugs to each of the patients with the specified dosage in the first three visits, 3 and 6 months later, the patients are examined with examination and radiology and the results of the study are recorded. If the drug results improve, it is continued, and if the results are not favorable, the second line is started.

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Theran Imam khomeini hospital complex

Full name of responsible person

Mohamad Amin Choobdar Omrani

Street address

Keshavarz blv

City

Tehran

Province

Tehran

Postal code

۱۴۱۹۷۳۳۱۴۱

Phone

+98 21 6119 2613

Email

Imamhospital@tums.ac.ir

2

Recruitment center

Name of recruitment center

Shariati hospital

Full name of responsible person

Mohamad Amin Choobdar Omrani

Street address

north kargar ave

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Province

Tehran

Postal code

1411713135

Phone

+98 21 8490 1000

Email

shariatihosp@tums.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tehran University of Medical Sciences

Full name of responsible person

Sina Rezaei

Street address

poorsina ave

City

Tehran

Province

Tehran

Postal code

1461884513

Phone

+98 21 8163 3619

Fax

Email

aminch2009@gmail.com

Web page address

Grant name

Grant code / Reference number

Is the source of funding the same sponsor organization/entity?

Yes

Title of funding source

Tehran University of Medical Sciences

Proportion provided by this source

100

Public or private sector

Public

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

Academic

Person responsible for general inquiries

Contact

Name of organization / entity

Tehran University of Medical Sciences

Full name of responsible person

Mohammad Amin Choobdar

Position

Resident

Latest degree

Medical doctor

Other areas of specialty/work

Orthopedics

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Sharing plan**Deidentified Individual Participant Data Set (IPD)**

Undecided - It is not yet known if there will be a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Yes - There is a plan to make this available

Data Dictionary

Yes - There is a plan to make this available

Title and more details about the data/document

All data can be shared

When the data will become available and for how long

The access period begins immediately after the results are published.

To whom data/document is available

All researchers in the field of medical and pharmaceutical sciences

Under which criteria data/document could be used

Researchers in the fields of medical and pharmaceutical sciences have permission to access the data. There are no restrictions on data analysis.

From where data/document is obtainable

mohammad amin choobdar 09200567515

dr.amin.choobdar@gmail.com

What processes are involved for a request to access data/document

پس از تماس و اعلام درخواست ابتدا مدارک مربوط به فرد درخواست دهنده جهت تایید ارسال شده و طی یک هفته داده ها از طریق ایمیل تحویل خواهد شد

Comments