

Clinical Trial Protocol

Iranian Registry of Clinical Trials

18 Sep 2026

In- Vivo Bioequivalence study of Empagliflozin-Metformin tablet (EMPATUS Plus® 25/1000 mg ,Pars Darou Pharma, Iran) with brand drug (Synjardy® 25/1000 mg, Boehringer Ingelheim, Germany) in Iranian healthy

Protocol summary

Study aim

In- Vivo Bioequivalence study of Empagliflozin-Metformin tablet (EMPATUS Plus® 25/1000 mg ,Pars Darou Pharma, Iran) with brand drug (Synjardy® 25/1000 mg, Boehringer Ingelheim, Germany) in Iranian healthy

Design

In-vivo bioequivalence study of EMPATUS Plus tablet Pars Darou Pharma. in compared with reference drug (Synjardy, Boehringer Ingelheim, German). The study design is single blind, Cross-over, two period & two groups (Intervention and control) with one week wash-out time. Randomization of the drug types in each period will perform by paper lottery randomization.

Settings and conduct

Study background is biopharmacy and pharmacokinetics. Place of study is Simin Baspar Tayf-Gostar Inc, Tabriz. Study blinding will done by taking the test and reference drugs out of their original packaging and put them in the similar packaging. The volunteers will not be informed about the type of received drug. In cross-over design, the test drug will taken by group 1 in first period and by group 2 in second period. The reference drug also will taken by group 2 in the first period and by group 1 in the second period.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Age between 18-55 years, body mass index (BMI) in the range of 18-28. Exclusion criteria: History of heart, kidney and liver disease, Pregnancy, Drug addiction, Smoking

Intervention groups

Single dose EMPATUS Plus tablet Pars Darou Pharma. Control group: brand drugs (Synjardy, Boehringer Ingelheim, German)

Main outcome variables

Plasma drug concentration

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20200105046010N60**

Registration date: **2022-05-05, 1401/02/15**

Registration timing: **registered_while_recruiting**

Last update: **2022-05-05, 1401/02/15**

Update count: **0**

Registration date

2022-05-05, 1401/02/15

Registrant information

Name

Javad Shokri

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 41 3661 4125

Email address

shokri.j@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2022-03-09, 1400/12/18

Expected recruitment end date

2022-09-09, 1401/06/18

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty	Street address Third floor; Central building; Tabriz University of Medical Sciences; Dneshgah St.
Scientific title In- Vivo Bioequivalence study of Empagliflozin-Metformin tablet (EMPATUS Plus® 25/1000 mg ,Pars Darou Pharma, Iran) with brand drug (Synjardy® 25/1000 mg, Boehringer Ingelheim, Germany) in Iranian healthy	City Tabriz
Public title In-vivo Bioequivalence Test of Empagliflozin-Metformin tablets with brand drug (Synjardy® 25/1000 mg, Boehringer Ingelheim, Germany)	Province East Azarbaijan
Purpose Treatment	Postal code 5166614766
Inclusion/Exclusion criteria Inclusion criteria: General health Body mass index between18-28 Informed consent Being at the age of 18-55 years old Exclusion criteria: Smoking A history of cardiovascular disease A history of liver & kidney disease Pregnancy Alcohol & Drug addiction Hypersensitivity to the drug	Approval date 2021-11-29, 1400/09/08
Age From 18 years old to 55 years old	Ethics committee reference number IR.TBZMED.REC.1400.825
Gender Both	Health conditions studied 1 Description of health condition studied Bio equivalence test ICD-10 code ICD-10 code description
Phase Bioequivalence	Primary outcomes 1 Description Plasma drug concentration Timepoint Sampling times in this study will be 0, 0:30, 1, 1:30, 2, 2:30, 3, 4, 6, 8, 10, 12, 24, 48 hours after prescribing the tablet. Method of measurement High performance liquid chromatography with tandem mass spectroscopy detector
Groups that have been masked <i>No information</i>	Secondary outcomes empty
Sample size Target sample size: 24	Intervention groups 1 Description Intervention group: One test tablet (EMPATUS Plus® 25/1000 mg ,Pars Darou Pharma, Iran) will be received. Blood samples will be taken for 72 hours at the mentioned times after drug administration and the concentration of the drug in Plasma samples will be measured by liquid chromatography with mass spectroscopy detector
Randomization (investigator's opinion) Randomized	Category Treatment - Other
Randomization description For this purpose, A 24- persons group will be selected and divided to two 12-persons groups randomly. The names of all volunteers will be written on paper pieces and wrapped in aluminum foils. The first 12 papers will randomly be withdrawn from bottle will be selected as group A and others will be categorized in group B.	2 Description Control group: One Reference tablet (Synjardy® 25/1000 mg, Boehringer Ingelheim, German) will be received. Blood samples will be taken for 72 hours at the
Blinding (investigator's opinion) Not blinded	
Blinding description	
Placebo Not used	
Assignment Crossover	
Other design features	
Secondary Ids empty	
Ethics committees 1 Ethics committee Name of ethics committee Ethics Committee of Tabriz University of Medical Sciences	

mentioned times after drug administration and the concentration of drug in plasma samples will be measured by liquid chromatography with mass spectroscopy detector.

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Simin Baspar Teyf Gostar Company

Full name of responsible person

Javad Shokri

Street address

No.48, Ferdos square

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Tabriz

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East Azarbaijan

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5167874434

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+98 41 3384 2724

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Shokri.j@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Pars Darou Pharmaceutical Company

Full name of responsible person

Amir Garmabadri

Street address

Tehran, the first square of Tehranpars, in front of 144 East Street, No.13

City

Tehran

Province

Tehran

Postal code

1654713691

Phone

+98 21 7770 4061

Email

info@parsdarou.ir

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Pars Darou Pharmaceutical Company

Proportion provided by this source

100

Public or private sector

Private

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

Persons

Person responsible for general inquiries

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Javad Shokri

Position

Professor

Latest degree

Ph.D.

Other areas of specialty/work

Medical Pharmacy

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Person responsible for scientific inquiries

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Full name of responsible person

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Position

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Latest degree

Ph.D.

Other areas of specialty/work

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Person responsible for updating data

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Dariush.omidfar

Position

Lab. Manager

Latest degree

Master

Other areas of specialty/work

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

No - There is not a plan to make this available

Justification/reason for indecision/not sharing IPD

All information and data of the study will remain secured

based on the agreement established between researcher and drug producer.

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

No - There is not 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

Only protocol and methods of study are sharable

When the data will become available and for how long

The access to data will be possible after finishing of project (almost 6 months after receiving of IRCT Code).

To whom data/document is available

Pharmaceutical and medical sciences researchers

Under which criteria data/document could be used

Projects information's for any publications is not allowed.

From where data/document is obtainable

By email to the project manager (shokri.j@gmail.com)

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

These information are confidential and be under disposal of the project's contractor. Upon request, the information will be accessed to the applicant by the Executor's email after receiving contractor's consent.

Comments