

Clinical Trial Protocol

Iranian Registry of Clinical Trials

09 Jun 2026

In healthy human volunteers; study of Insulin Glargine (Basagine™) after single subcutaneous injection of 0.2 unit/Kg dose; compared with nothing and primarily to establish the PK parameters Cmax, Tmax and AUC in Pakistani population.

Protocol summary

Summary

Study Title: Pharmacokinetic evaluation of Insulin-Glargine (rDNA origin) 100 u/ml (Basagine™) in local healthy human volunteers after administration of a 0.2 units/Kg subcutaneous dose. Study Design: Open label, single period, single treatment, and single dose study in fourteen healthy volunteers. Goal/Objective: To determine AUC_{0-t}, C_{max} and, T_{max} of the drug in Pakistani population Inclusion Criteria: Healthy male/female, Age: 18 to 50 years, BMI: 18-26 Kg/m², Fasting plasma glucose: <6.0 mmol/l, non-smoker, non-alcoholic and no drug abuse. Exclusion criteria: Volunteers with any neuropathy, diabetes mellitus, Hypertension and any Cardio Vascular disease, positive test results for HBsAg, anti HCV and HIV antibody. Findings of hematology, biochemistry and ECG are out of range revealing clinical significance. Treatments: Single subcutaneous dose of Insulin-Glargine (rDNA origin; Basagine™) 0.2 units/Kg. Afterwards 5ml blood would be collected at 30 min, 15 min and 0 min before and 1, 2, 3, 4, 6, 8, 12, 16, 20, 22, 23, 24, 25 hours after drug administration. 120-240 ml 20% glucose will be given orally every 30 minutes to keep the blood glucose levels above 100mg/dl. Safety: Through physical examination, vital sign, adverse events, glucose monitoring and lab test monitoring on screening and at follow up after 72 hours. Throughout the study blood glucose levels would be monitored to maintain the blood glucose level between 3.3 to 7.7 mmol/liter. Measurements/ Analysis: Insulin and C-peptide through Chemiluminescence. Pharmacokinetic parameters would be determined by model independent method using PPstat soft ware. Ethical consideration: Approved from IRB and full compliance to ICH-GCP.

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT201203227978N2**

Registration date: **2012-04-01, 1391/01/13**

Registration timing: **prospective**

Last update:

Update count: **0**

Registration date

2012-04-01, 1391/01/13

Registrant information

Name

Tasneem Ahmad

Name of organization / entity

Pharma Professional Services, Karachi, Pakistan

Country

Pakistan

Phone

(92-21) 34972358, 34820573

Email address

tasneem.ahmed@iccs.edu

Recruitment status

Recruitment complete

Funding source

Getz Pharma (Pvt) Ltd; Karachi, Pakistan

Expected recruitment start date

2012-06-22, 1391/04/02

Expected recruitment end date

2012-06-22, 1391/04/02

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

In healthy human volunteers; study of Insulin Glargine (Basagine™) after single subcutaneous injection of 0.2 unit/Kg dose; compared with nothing and primarily to establish the PK parameters Cmax, Tmax and AUC in Pakistani population.

Public title

A STUDY OF BASAGINE™ IN LOCAL HEALTHY HUMAN SUBJECTS TO EVALUATE ITS PHARMACOKINETIC BEHAVIOUR

Purpose

Other

Inclusion/Exclusion criteria

INCLUSION: Healthy male/female, Age:18 to 50 years, BMI: 18-26 Kg/m2, Fasting plasma glucose: <6.0 mmol/l (<108 mg/dl) or normal range and Non-smoker, non-alcoholic
EXCLUSION: • Volunteers with any neuropathy, diabetes mellitus. • Any active allergic disease or a history of significant allergic disease. • Presence of renal, hepatic or gastrointestinal disease known to interfere with the drug absorption, distribution, metabolism or elimination with in last year. • Subject demonstrates protocol non-compliance (e.g. uncooperative attitude, & inability to finish study). • Participation in another study within last 2 months of 1st drug administration. • If donated blood within last 2 months preceding the study. • Age below 18 years and above 50yr. • Smoking within last 3 months prior to the drug administration and 6 hours after drug administration. • Ingestion of OTC drug (except Paracetamol) within last 14 days of 1st drug administration. • Participants with insufficient organ and/or bone marrow dysfunction. • Ingestion of investigational drug within 1 year prior to 1st drug administration. • Participants with an uncontrolled medical condition (i.e., hypertension, cardiac arrhythmias, CHF) that places the patient at risk by participating in the study. • Results of CBC, LFT, serum creatinine, fasting plasma glucose indicating significantly out of range and abnormal ECG. • Participants with any physical/mental disability • Subjects with known HIV, hepatitis B or hepatitis C infection, or autoimmune diseases. • History of major organ transplantation, new onset diabetes, unstable thyroid function. • Alcohol or drug abuse within the past year. • Known hypersensitivity to investigational drug • Positive drug of abuse test and alcohol test. • Intake of gutka, pan and any other thing containing nicotine 48 hours before and during study.

Age

From **18 years** old to **50 years** old

Gender

Male

Phase

4

Groups that have been masked

No information

Sample size

Target sample size: **14**

Randomization (investigator's opinion)

N/A

Randomization description**Blinding (investigator's opinion)**

Not blinded

Blinding description**Placebo**

Not used

Assignment

Single

Other design features

Open label, single period, single treatment, and single dose study in fourteen healthy volunteers.

Secondary Ids**1****Registry name**

The Universal Trial Number (UTN)

Secondary trial Id

U1111-1129-4410

Registration date

2012-03-24, 1391/01/05

Ethics committees**1****Ethics committee****Name of ethics committee**

Instituional Review Board, Civil Hospital, Dow University of Health Sciences, Karachi

Street address

Instituional Review Board, Civil Hospital, Dow University of Health Sciences, Karachi

City

Karachi

Postal code

74200

Approval date

2012-03-10, 1390/12/20

Ethics committee reference number

IRB-311/DUHS-12

Health conditions studied**1****Description of health condition studied**

Diabetes mellitus

ICD-10 code

E10-E14

ICD-10 code description

Diabetes mellitus Without complications

Primary outcomes**1****Description**

Serum levels of Insulin and C-peptide

Timepoint

30 min, 15 min before and 0, 1, 2, 3, 4, 6, 8, 12, 16, 20, 22, 23, 24, 25 hours after

Method of measurement

Serum concentrations of Insulin and C-peptide would be measured through Chemiluminescence.

Secondary outcomes**1****Description**

Pharmacokinetic parameters: C_{max}, T_{max} and AUC_{0-t}

Timepoint

Pharmacokinetic (PK) parameter will be determined: Area under the curve to the last measurable concentration (AUC_{0-t}) will be calculated by the linear trapezoidal rule. C_{max} and t_{max} would also be recorded from the drug concentration vs time profile.

Method of measurement

Pharmacokinetic (PK) parameter will be determined on the basis of measurement of exogenous and endogenous concentration of insulin; performed by means of model independent method using PPstat computer programs.

Intervention groups**1****Description**

Administration of Insulin-Glargine (rDNA origin; 100 u/ml; Basagline™) 0.2 units/Kg subcutaneous dose

Category

Treatment - Drugs

Recruitment centers**1****Recruitment center****Name of recruitment center**

Dow University of Health Sciences (DUHS), Karachi

Full name of responsible person

Dr. Khaqan Ahmad

Street address

Dow University of Health Sciences (DUHS) Ojha Campus

City

Karachi

Sponsors / Funding sources**1****Sponsor****Name of organization / entity**

Getz Pharma (Pvt) Limited

Full name of responsible person

Dr. Khawar Mehdi

Street address

Director Medical Affairs, 29-30, Secotr-27, Korangi Industrial Area

City

Karachi

Grant name**Grant code / Reference number****Is the source of funding the same sponsor organization/entity?**

Yes

Title of funding source

Getz Pharma (Pvt) Limited

Proportion provided by this source

100

Public or private sector

empty

Domestic or foreign origin

empty

Category of foreign source of funding

empty

Country of origin**Type of organization providing the funding**

empty

Person responsible for general inquiries**Contact****Name of organization / entity**

Pharma Professional Services, Karachi, Pakistan

Full name of responsible person

Ms. Ghousia Saba

Position

Chief Scientific Officer

Other areas of specialty/work**Street address**

8 Gulshan View, Gulshan-e-Iqbal, Karachi,

City

Karachi

Province

Sind

Postal code

75300

Phone

00922134972358

Fax**Email**

ghousia@phaps.com

Web page address

www.phaps.com

Person responsible for scientific inquiries**Contact****Name of organization / entity**

Pharma Professional Services, Karachi, Pakistan

Full name of responsible person

Prof. Dr. Tasneem Ahmad

Position

CEO/ B.Pharm, M.Pharm, Ph.D.

Other areas of specialty/work**Street address**

8 Gulshan View, Gulshan-e-Iqbal, Karachi,

City

Karachi

Province

Sind

Postal code

75300

Phone

00922134972358

Fax**Email**

tasneem_ahmad2001@yahoo.com

Web page address

www.phaps.com

Province

Sind

Postal code

75300

Phone

00922134972358

Fax**Email**

tasneem_ahmad2001@yahoo.com

Web page address

www.phaps.com

Person responsible for updating data**Contact****Name of organization / entity**

Pharma Professional Services, Karachi, Pakistan

Full name of responsible person

Prof. Dr. Tasneem Ahmad

Position

CEO/ B.Pharm, M.Pharm, Ph.D.

Other areas of specialty/work**Street address**

8 Gulshan View, Gulshan-e-Iqbal, Karachi,

City

Karachi

Sharing plan**Deidentified Individual Participant Data Set (IPD)***empty***Study Protocol***empty***Statistical Analysis Plan***empty***Informed Consent Form***empty***Clinical Study Report***empty***Analytic Code***empty***Data Dictionary***empty*