

# A comparative, open-label, randomised, cross-over phase I trial in healthy volunteers to investigate the relative efficacy, safety and tolerability of OctaplasLG™ versus Octaplas® SD

<b>Submission date</b>	<b>Recruitment status</b>	<input type="checkbox"/> Prospectively registered
03/09/2009	No longer recruiting	<input type="checkbox"/> Protocol
<b>Registration date</b>	<b>Overall study status</b>	<input type="checkbox"/> Statistical analysis plan
04/09/2009	Completed	<input type="checkbox"/> Results
<b>Last Edited</b>	<b>Condition category</b>	<input type="checkbox"/> Individual participant data
28/10/2009	Injury, Occupational Diseases, Poisoning	<input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

Not provided at time of registration

## Contact information

### Type(s)

Scientific

### Contact name

Dr Claudia Walasek

### Contact details

Oberlaaerstrasse 235

Vienna

Austria

1100

+43 (0)1 61032 1791

claudia.walasek@octapharma.com

## Additional identifiers

### Clinical Trials Information System (CTIS)

2009-012856-26

### Protocol serial number

LAS-203

# Study information

## Scientific Title

### Study objectives

Comparison of efficacy, safety and tolerability of OctaplasLG™ versus Octaplas® SD plasma in healthy volunteers.

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Local medical ethics committee (ethikkommission der med Uni Wien und des Allg Krankenhauses der Stadt Wien AKH) approved on the 15th July 2009 (ref: 460/2009)

### Study design

Open-label block randomised cross-over phase I study

### Primary study design

Interventional

### Study type(s)

Treatment

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

Safety/efficacy/tolerability of plasma products

### Interventions

The treatment day will start with plasmapheresis (600 ml) then transfusion of either OctaplasLG™ or Octaplas® SD will be randomly assigned. Safety, efficacy and tolerability will be assessed by clinical and laboratory parameters (haematology, coagulation factors, haemostatic parameters, chemistry). All these parameters will be collected before and immediately after plasmapheresis (PP), then 15 minutes, 2 hours, 24 hours and 7 days after end of IMP administration. Treatment sequence is either OctaplasLG™ or Octaplas® SD or vice versa after a minimal wash out period of 1 month. The overall duration per subject will be 1.5 months and a treatment performed on 2 days.

### Intervention Type

Drug

### Phase

Phase I

### Drug/device/biological/vaccine name(s)

OctaplasLG™, Octaplas® SD

### Primary outcome(s)

## 1. Coagulation factors

2. Activated partial thromboplastin time (aPTT), prothrombin time (PT), protein C

All primary and secondary endpoints will be measured before and immediately after PP and at 15 minutes, 2 hours and 24 hours post-transfusion of IMP. Haematology and clinical chemistry will be measured 7 days after end of IMP administration.

## Key secondary outcome(s)

1. Haematology: red blood cell (RBC) count, white blood cell (WBC) count, platelets, haematocrit (Hct), haemoglobin (Hb), and plasmin inhibitor, Protein S
2. Clinical Chemistry: electrolytes, creatinine, alanine aminotransferase (ALAT), gamma-glutamyl transferase (GGT), total protein (TP)
3. Overall tolerability, vital parameters

All primary and secondary endpoints will be measured before and immediately after PP and at 15 minutes, 2 hours and 24 hours post-transfusion of IMP. Haematology and clinical chemistry will be measured 7 days after end of IMP administration.

## Completion date

01/10/2010

# Eligibility

## Key inclusion criteria

1. Subject must be capable of understanding and complying with all aspects of the protocol
2. Signed informed consent
3. Subject must be capable of understanding the plasmapheresis information sheet and sign it
4. Healthy male or female volunteers, aged 18 years or above
5. Women must have a negative pregnancy test (human chorionic gonadotrophin [HCG]-based assay)
6. Women must have sufficient methods of contraception (e.g. intrauterine device, oral contraception, etc.)
7. Subjects must have no clinically relevant abnormalities in medical history and general physical examination
8. Standard health insurance

## Participant type(s)

Patient

## Healthy volunteers allowed

No

## Age group

Adult

## Lower age limit

18 years

## Sex

All

## **Key exclusion criteria**

1. Pregnancy or lactation
2. Tattoos within the last 3 months
3. Subject was treated therapeutically with FFP, blood or plasma derived products in the previous 6 months
4. Subjects have a hypersensitivity to blood products or plasma protein
5. History of angioedema
6. History of coagulation or bleeding disorder or any other known abnormality affecting coagulation, fibrinolysis or platelet function
7. Any clinically significant abnormal laboratory values
8. IgA deficiency
9. Seropositivity for HBs-Ag, HCV, HIV-1/2 antibodies
10. Symptoms of a clinically relevant illness within 3 weeks before the first trial day
11. Subjects with a history of, or suspected, drug or alcohol abuse
12. Subjects currently participating in another clinical study
13. Any IMP administration within the last 4 weeks

## **Date of first enrolment**

01/07/2009

## **Date of final enrolment**

01/10/2010

## **Locations**

### **Countries of recruitment**

Austria

### **Study participating centre**

Oberlaaerstrasse 235

Vienna

Austria

1100

## **Sponsor information**

### **Organisation**

Octapharma AG (Switzerland)

### **ROR**

<https://ror.org/002k5fe57>

## **Funder(s)**

**Funder type**

Industry

**Funder Name**

Octapharma AG (Switzerland)

## Results and Publications

### Individual participant data (IPD) sharing plan

#### IPD sharing plan summary

Not provided at time of registration

#### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>	Participant information sheet	11/11/2025	11/11/2025	No	Yes