# An Open-Label, Multi-Center Controlled Clinical Trial of Eculizumab in Adult Patients with Plasma Therapy-Sensitive Atypical Hemolytic-Uremic Syndrome (AHUS)

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Objectives: The following trial objectives for adult patients with plasma therapy-sensitive Atypical Hemolytic-Uremic Syndrome (aHUS) are to:Primary:\* Assess the effect of eculizumab on TMA-Event Free status defined as the absence of [1] decrease in...

**Ethical review** Approved WMO

**Status** Recruitment stopped

**Health condition type** Haemolyses and related conditions

Study type Interventional

# **Summary**

#### ID

NL-OMON33473

#### Source

**ToetsingOnline** 

**Brief title** 

C08-003A

#### Condition

Haemolyses and related conditions

## **Synonym**

atypical hemolytic uremic syndrome

## Research involving

Human

Sponsors and support

**Primary sponsor:** Alexion Pharmaceuticals

Source(s) of monetary or material Support: Alexion

Intervention

**Keyword:** adults, aHUS, eculizumab, therapy-sensitive

**Outcome measures** 

**Primary outcome** 

**Primary Endpoint:** 

\* The primary efficacy endpoint for this protocol is TMA-Event Free status

defined as the absence of [1] decrease in platelet count of >25% from the

Platelet Count Pre-PT Baseline Set-Point; [2] PT while the patient is receiving

eculizumab, and

[3] new dialysis for at least 12 weeks in adult patients with plasma

therapysensitive Atypical Hemolytic-Uremic Syndrome (aHUS). Dialysis events

occurring within the 14 days after the first dose of Investigational Product

will not be considered as a new Treatment Period dialysis event. In addition,

dialysis events that commence within the 14 days before the first dose of

Investigational Product and continue up to 14 days after the first dose of

Investigational Product will not be considered a new Treatment Period dialysis

event;

**Secondary outcome** 

Secondary Endpoints:

\* TMA Intervention Rate (# PT and # Dialysis Events/Patient/Day) during the

Treatment Period compared with the TMA Intervention Rate prior to the first

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dose of Investigational Product;

- \* Platelet count change from baseline (Platelet Count Pre-PT Baseline Set-Point value) after first dose of Investigational Product;
- \* Change in LDH from baseline after first dose of Investigational Product;
- \* Change in Quality of Life measures \* EuroQol 5D from baseline after first dose of Investigational Product;
- \* Change in renal function parameters as assessed by change in CKD stage from baseline after the first dose of Investigational Product;
- \* TMA Remission;
- \* Safety and tolerability of eculizumab;
- \* PK and PD parameters during induction and maintenance phases of treatment.

## **Exploratory Endpoints:**

- \* Weekly TMA-related thrombocytopenia resolution rate defined as the absence of [1] platelet count decrease > 25% from the Platelet Count Pre-PT Baseline Set Point and [2] platelet count < 150 x 109/L;
- \* Changes in parameters associated with intravascular hemolysis, such as in haptoglobin and schistocytes, from baseline after first dose of Investigational Product;

# **Study description**

## **Background summary**

Because of the severe unmet medical need in the treatment of patients with this serious and lifethreatening rare disorder, and the demonstrated activity

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of eculizumab to reduce thrombotic microangiopathy (TMA); individual physicians have chosen to utilize eculizumab in the treatment of their severely affected aHUS patients. To date, there is limited clinical experience in four therapy-resistant patients, of which one young child, and one therapy-sensitive patient. These initial results in severely ill aHUS patients have generated the hypothesis to confirm whether eculizumab treatment will improve the TMA condition of patients affected by aHUS.

See also page 28 and 29 of the protocol; 7.5.1 Eculizumab in aHUS Patients

## Study objective

Objectives: The following trial objectives for adult patients with plasma therapy-sensitive Atypical Hemolytic-Uremic Syndrome (aHUS) are to: Primary:

\* Assess the effect of eculizumab on TMA-Event Free status defined as the absence of [1] decrease in platelet count of >25% from the Platelet Count Pre-PT Baseline Set-Point; [2] PT while the patient is receiving eculizumab, and [3] new dialysis for at least 12 weeks in adult patients with plasma therapy-sensitive Atypical Hemolytic-Uremic Syndrome (aHUS).

## Secondary:

- \* Evaluate additional efficacy endpoints such as the effect of eculizumab on:
- \* TMA Intervention Rate (# PT and # Dialysis Events/Patient/Day) during the Treatment Period compared with the TMA Intervention Rate prior to the first dose of Investigational Product.
- \* Reduction of thrombotic microangiopathy (TMA) as indicated by thrombocytopenia as measured by platelet count change from baseline through the treatment period.
- \* Key Hemolytic measures.
- \* Quality of Life measures.
- \* Renal function measures.
- \* TMA Remission.

## Study design

After screening patient will start with the observation phase and come to the hospital for a weekly visit for 8 weeks.

Patients will be treated with eculizumab for 26 weeks and will have post-treatment assessments performed at 1 week, 2 weeks, 4 weeks and 8 weeks after the last dose of eculizumab. Total trial duration for each patient is approximately 35 weeks (3-day Screening Period, 26 Week Treatment Period, 8-Week Post-Treatment Follow-up Period). The estimated duration of the study, including an estimated 6 month enrollment period is approximately 14 months. Patients may be eligible to enroll in an open-label extension study after

completing study evaluations to Week 26.

Patients who prematurely discontinue investigational product during the study or who do not enter the extension study will require follow-up contacts for 8 weeks after the last dose of eculizumab.

#### Intervention

Eculizumab 900 mg or 1200 mg will be administered intravenously according to the following regimens:

- \* Induction Period: patients will receive eculizumab 900mg via IV infusion over approximately 35 minutes once a week (every  $7 \pm 2$  days) for 4 weeks followed by 1200mg eculizumab for the 5th dose ( $7 \pm 2$  days) later.
- \* Maintenance Period: patients will receive eculizumab 1200mg via IV infusion over approximately 35 minutes every two weeks (every 14 ± 2 days).
- \* If the physician administers plasmapheresis, plasma exchange or FFP, 600 mg eculizumab must be administered (i) within 60 minutes after each 1 volume plasmapheresis or plasma exchange and (ii) within 60 minutes prior to each 3 units

of FFP infusion, respectively. Please refer to Section 11.2.1.1 Plasma Therapy for details.

## Study burden and risks

For most patients, placements of an IV catheter and needle punctures for blood draws are usually well tolerated. However, they rarely may cause pain, bleeding, bruising, swelling, clotting, leakage of drug, and possibly infection at the needle or catheter site.

Patients will receive a vaccine against Neisseria meningitidis that can cause adverse reactions. Different types of meningococcal vaccines are available. The study doctor may choose the most appropriate according to the condition. The electrocardiogram is a painless procedure that traces the activity of the heart.

Plasma therapy procedures may be performed by your study doctor, at his/her discretion. The plasma for the transfusion may come from different blood supply sources depending on the medical institution. Although eculizumab is being tested for the condition of the patient, there is no guarantee that they will not be exposed to increased risks of plasma transfusion reactions after being treated with eculizumab.

## **Contacts**

#### **Public**

Alexion Pharmaceuticals

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#### **Scientific**

**Alexion Pharmaceuticals** 

54/56 Avenue Hoche 75008 Paris FR

## **Trial sites**

## **Listed location countries**

Netherlands

# **Eligibility criteria**

## Age

Adults (18-64 years) Elderly (65 years and older)

## Inclusion criteria

- 1. Male or female patients \*18 years of age who have been diagnosed with Atypical Hemolytic-Uremic Syndrome (aHUS).
- 2. Patients must be receiving PT for aHUS and must be observed to (i) receive \*1 PT treatment every two weeks and no more than 3 PT treatments/week (at an unchanged frequency) for at least 8 weeks immediately prior to first dose of Investigational Product and (ii) receive the same volume of PP or PE and units of FFP for at least 8 weeks immediately prior to first dose of Investigational Product.
- 3. Platelet Count Pre-PT Baseline Set-Point (collected immediately prior to the Qualifying PT Episode) is within 75% of the average of the pre-PT platelet counts collected at Screening and during the Observation Period.
- 4. Known complement regulatory protein genetic abnormality, i.e., a mutation in Complement Protein 3, factor H or associated factor, factor I, or membrane cofactor protein 1 (MCP-1) or known Factor B gain-of-function mutation, or known anti-CFH antibody (\*aHUS lesions\*).
- \* Patients diagnosed with aHUS with any of these aHUS lesions are eligible and will be assigned to one of the following parallel categories during the treatment period of the trial:
- \* (Category 1) Complement Protein 3 or factor H or factor I functional

deficiency or abnormal factor interaction (C3/CFH/CFI FFP Group);

- \* (Category 2) Factor B Gain of Function;
- \* (Category 3) Anti-CFH Antibody (Anti-CFH Group);
- \* (Category 4) MCP-1 deficiency (MCP-1 Group);
- 5. Patients diagnosed with HUS of the atypical type without documented complement regulatory protein genetic abnormality or known anti-CFH antibody are eligible if other etiologies of HUS have been ruled out as confirmed in the Exclusion Criteria (i.e., including Shiga-toxin negative, non-infectious, non-drug-exposure-related [e.g., cyclosporine]), no known HIV positivity, and anti-phospholipid antibody negative). Thrombotic thrombocytopenic purpura also must be ruled out (i.e., ADAMTS-13 activity must be > 5%; see Exclusion Criteria). Patients meeting these conditions will be assigned to Category 5. In addition, these patients will undergo genetic testing to determine if a mutation can be identified. If a mutation is identified, the patient will be reassigned to the appropriate category.
- 6. Lactate dehydrogenase (LDH) level \* ULN.
- 7. Creatinine level \* ULN for age.
- 8. Sexually active women of childbearing potential must be practicing an effective, reliable and medically acceptable contraceptive regimen during the entire duration of the study, including the follow-up period.
- 9. Able to give written informed consent.
- 10. Able and willing to comply with study procedures.

## **Exclusion criteria**

- 1. ADAMTS-13 inhibitor or deficiency (i.e., ADAMTS-13 activity <5%) as measured at the screening visit.
- 2. Malignancy.
- 3. Typical HUS (Shiga toxin +).
- 4. Known HIV infection.
- 5. Identified drug exposure-related HUS.
- 6. Infection-related HUS.
- 7. Patients with a confirmed diagnosis of sepsis defined as positive blood cultures within 7 days of the screening visit and not treated with antibiotics to which the organism is sensitive.
- 8. Presence or suspicion of active and untreated systemic bacterial infection that, in the opinion of the Investigator confounds an accurate diagnosis of aHUS or impedes the ability to manage the aHUS disease.
- 9. Pregnancy or lactation.
- 10. Unresolved meningococcal disease.
- 11. Known Systemic Lupus Erythematosus (SLE) or antiphospholipid antibody positivity or syndrome.
- 12. Any medical or psychological condition that, in the opinion of the investigator, could increase the patient\*s risk by participating in the study or confound the outcome of the study.
- 13. Patients receiving IVIg or Rituximab therapy.
- 14. Patients receiving other immunosuppressive therapies such as steroids, mTOR

inhibitors or FK506 inhibitors are excluded unless: [1] part of a post-transplant antirejection regime, [2] patient has confirmed anti-CFH antibody requiring

immunosuppressive therapy and [3] dose of such medications have been unchanged for at least 4 weeks prior to the screening period.

- 15. Patients receiving Erythrocyte Stimulating Agents (ESAs) unless already on a stable dose for at least 4 weeks prior to the screening period.
- 16. Participation in any other investigational drug trial or exposure to other investigational agent, device, or procedures beginning 4 weeks prior to screening and throughout the entire trial.

# Study design

## **Design**

Study phase: 2

Study type: Interventional

Masking: Open (masking not used)

Control: Uncontrolled

Primary purpose: Treatment

### Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 01-10-2009

Enrollment: 1

Type: Actual

## Medical products/devices used

Product type: Medicine

Brand name: eculizumab

Generic name: Soliris

Registration: Yes - NL outside intended use

# **Ethics review**

## Approved WMO

Date: 19-03-2009

Application type: First submission

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 12-08-2009

Application type: First submission

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 30-11-2009

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

Approved WMO

Date: 22-10-2013

Application type: Amendment

Review commission: CMO regio Arnhem-Nijmegen (Nijmegen)

# **Study registrations**

## Followed up by the following (possibly more current) registration

No registrations found.

## Other (possibly less up-to-date) registrations in this register

No registrations found.

# In other registers

Register ID

EudraCT EUCTR2008-006954-17-NL

ClinicalTrials.gov NCT00838513 CCMO NL27321.091.09