

NN7088-4029 A multinational, prospective, open labelled, non-controlled, non-interventional post-authorisation study of turoctocog alfa pegol (N8-GP) during long-term routine prophylaxis and treatment of bleeding episodes in patients with haemophilia A (pathfinder 9)

**First published:** 21/09/2020

**Last updated:** 08/01/2025

Study

Ongoing

## Administrative details

**EU PAS number**

EUPAS36536

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**Study ID**

39327

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**DARWIN EU® study**

No

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## Study countries

-  Austria
  -  Croatia
  -  Czechia
  -  Estonia
  -  Germany
  -  Greece
  -  Hungary
  -  Italy
  -  Lithuania
  -  Portugal
  -  Slovakia
  -  Slovenia
  -  Spain
  -  Switzerland
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## Study description

This study will collect information on side effects and how well Esperoct® (turoctocog alfa pegol (N8-GP)) works during long-term treatment (prophylaxis) in males with haemophilia A. Participants in this study will get the same treatment as they would normally get, if they were not participating in the study. All visits at the clinic are done in the same way as participants are used to, when visiting their doctor. During visits at the clinic participants might be asked for some relevant tests if considered useful by the study doctor. During the visits the study doctor might ask if participants had any side effects since the last study visit. Participants will be asked to note down in their own diary the number of bleeds and how these were treated, as well as their regular prophylaxis. Participation in the study will last for about 5-7 years, depending on when participants join the study. Participants are free to leave the study at any time and for any reason. This will not affect their current and future medical

care.

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## Study status

Ongoing

## Research institutions and networks

### Institutions

#### Novo Nordisk

**First published:** 01/02/2024

**Last updated:** 01/02/2024

Institution

## Contact details

### Study institution contact

Clinical Transparency and Medical Writing Office (1452)

Novo Nordisk A/S [pactadmin@novonordisk.com](mailto:pactadmin@novonordisk.com)

Study contact

[pactadmin@novonordisk.com](mailto:pactadmin@novonordisk.com)

### Primary lead investigator

Clinical Transparency and Medical Writing Office (1452)

Novo Nordisk A/S

Primary lead investigator

# Study timelines

## **Date when funding contract was signed**

Actual: 17/04/2020

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## **Study start date**

Planned: 01/10/2020

Actual: 23/10/2020

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## **Date of final study report**

Planned: 02/06/2028

# Sources of funding

- Pharmaceutical company and other private sector

# More details on funding

Novo Nordisk A/S

# Study protocol

[4029-protocol-eu-pas-reg-redacted.pdf](#) (1.07 MB)

[\\_ Protocol 4029 protocol eu-pas-reg redacted \(1\).pdf](#) (715.55 KB)

# Regulatory

## **Was the study required by a regulatory body?**

Yes

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## Is the study required by a Risk Management Plan (RMP)?

EU RMP category 1 (imposed as condition of marketing authorisation)

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### Regulatory procedure number

EMA/H/C/004883/0000

## Other study registration identification numbers and links

UTN: U1111-1235-6007

## Methodological aspects

### Study type

### Study type list

#### Study topic:

Human medicinal product

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#### Study type:

Non-interventional study

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#### Scope of the study:

Safety study (incl. comparative)

#### Main study objective:

The primary objective of the study is to investigate the safety of N8-GP including the PEG moiety during prophylaxis and long-term use in patients with

haemophilia A as prescribed by the physician.

## Study Design

### **Non-interventional study design**

Cohort

## Study drug and medical condition

### **Study drug International non-proprietary name (INN) or common name**

TUROCTOCOG ALFA PEGOL

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### **Medical condition to be studied**

Factor VIII deficiency

## Population studied

### **Age groups**

- Term newborn infants (0 - 27 days)
  - Infants and toddlers (28 days - 23 months)
  - Children (2 to < 12 years)
  - Adolescents (12 to < 18 years)
  - Adults (18 to < 46 years)
  - Adults (46 to < 65 years)
  - Adults (65 to < 75 years)
  - Adults (75 to < 85 years)
  - Adults (85 years and over)
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## Estimated number of subjects

60

## Study design details

### Outcomes

Number of Adverse Events (AEs) reported during the study period from inclusion of the patient until end of study, Number of Serious Adverse Events (SAEs) reported during the study period from inclusion of the patient until end of study

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### Data analysis plan

No formal testing of statistical hypothesis will be performed. All data will be presented using descriptive statistics. Categorical data will be summarized by frequency tables while continuous data will be summarized by mean, standard deviation, median, minimum and maximum value. Subgroup analysis will be presented: - By age groups (< 6 years, 6 to < 12 years, 12 to < 18 years, 18 to < 65 years, ≥ 65 years) - By severity of disease (moderate and severe) Patients who previously developed inhibitors before entering this study might be presented separately if deemed necessary.

## Documents

### Study report

[4029 progress report eu-pas-reg 01 redacted.pdf](#) (263.77 KB)

[4029 progress report eu-pas-reg 02 redacted.pdf](#) (476.39 KB)

### Study, other information

[4029 progress report eu-pas-reg 02 redacted.pdf](#) (476.39 KB)

## Data management

## ENCePP Seal

The use of the ENCePP Seal has been discontinued since February 2025. The ENCePP Seal fields are retained in the display mode for transparency but are no longer maintained.

## Data sources

### Data sources (types)

Other

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### Data sources (types), other

Prospective patient-based data collection

## Use of a Common Data Model (CDM)

### CDM mapping

No

## Data quality specifications

### Check conformance

Unknown

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### Check completeness

Unknown

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### Check stability

Unknown

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## **Check logical consistency**

Unknown

## Data characterisation

### **Data characterisation conducted**

No