

ATLAS-PPX trial: an open-label, multinational, switching study to describe the efficacy and safety of fitusiran prophylaxis in patients with hemophilia A and B with inhibitory antibodies to factor VIII or IX previously receiving bypassing agent prophylaxis.

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1.3. Study Design Rationale The ATLAS-PPX trial (ALN-AT3SC-009) is a multicenter, multinational, open-label Phase 3 switching study designed to demonstrate the efficacy and safety of fitusiran in patients with hemophilia A or B, with inhibitory...

Ethical review	Not approved
Status	Will not start
Health condition type	Coagulopathies and bleeding diatheses (excl thrombocytopenic)
Study type	Interventional

Summary

ID

NL-OMON46351

Source

ToetsingOnline

Brief title

ATLAS ALN-AT3SC-009

Condition

- Coagulopathies and bleeding diatheses (excl thrombocytopenic)

Synonym

Hemophilia A or B

Research involving

Human

Sponsors and support

Primary sponsor: Alnylam Pharmaceuticals Inc.

Source(s) of monetary or material Support: Alnylam Pharmaceuticals Inc.

Intervention

Keyword: fitusiran, hemophilia A or B

Outcome measures

Primary outcome

Primary

* Annualized Bleeding Rate (ABR) in the fitusiran efficacy period and the BPA prophylaxis period

Secondary outcome

Secondary

* Annualized spontaneous bleeding rate in the fitusiran efficacy period and the BPA prophylaxis period

* Annualized joint bleeding rate in the fitusiran efficacy period and the BPA prophylaxis period

* Change in Haem-A-QOL score in the fitusiran treatment period

* ABR in the fitusiran onset period

* ABR in fitusiran treatment period

Study description

Background summary

Hemophilia is a rare bleeding problem in which blood does not clot normally. This means that people with hemophilia may bleed for longer periods of time after an injury or, they may develop bleeds spontaneously. This happens because people with hemophilia have little or none of certain clotting factors. Clotting factors are proteins in the blood that help the body to stop bleeding by forming a blood clot. Fitusiran may make it possible to prevent or reduce the frequency of hemophilia-related bleeding in patients with hemophilia. Your study doctor will explain the detailed information below about how the drug is believed to behave.

Study objective

1.3. Study Design Rationale

The ATLAS-PPX trial (ALN-AT3SC-009) is a multicenter, multinational, open-label Phase 3 switching study designed to demonstrate the efficacy and safety of fitusiran in patients with hemophilia A or B, with inhibitory antibodies to factor VIII or IX who are currently treated with prophylactic regimens of BPAs.

The switching design allows for an intra-patient control to enable examination of the effect of the two treatment methods through comparison of the median ABR during the BPA prophylaxis period and the median ABR of the same patient group when receiving fitusiran, while limiting confounding effects of different patient bleeding phenotypes and prophylaxis therapy variability. The onset period duration reflects modeling data that estimates it takes approximately 28 days to reach the therapeutic target range in the majority of patients.

Given that the study design employed is a single treatment arm, with a switch from prophylaxis to fitusiran for each patient, the study is not blinded.

The primary endpoint of the study is ABR in the fitusiran efficacy period and the BPA prophylaxis period. ABR is a well-established endpoint that has been used as the primary endpoint in global approvals of factor replacement and BPA products. Secondary endpoints

characterize annualized spontaneous and joint bleeding rates, change in Haem-A-QoL score in patients ≥ 17 years of age, ABR in the onset period, and overall safety profile.

Characterization of bleeding episodes is clinically relevant to assess overall bleeding episode protection. Joint bleeding episodes result in pain and hemarthrosis, leading to progressive joint destruction, and hence are important to assess. Haem-A-QoL is a hemophilia-specific HRQOL survey instrument, has been used in other hemophilia clinical trials, has been validated, reviewed by clinicians, and is considered the most appropriate HRQOL tool available for use in the study.

The study population will be comprised of males ≥ 12 years of age; it is appropriate to study fitusiran in adolescents (patients ≥ 12 to <18 years of age) because the pathophysiology of disease progression and bleeding episode management is the same as adults and self-management of

hemophilia typically begins at 12 years of age [6].

In the event of a breakthrough bleeding episode, on-demand use of BPAs will be permitted throughout the entire study duration.

Study design

Study Design

The ATLAS-PPX trial (ALN-AT3SC-009) is a multicenter, multinational, open label, Phase 3 study designed to evaluate the efficacy and safety of fitusiran in male patients, aged ≥12 years, with hemophilia A or B, with inhibitory antibodies to factor VIII (FVIII) or factor IX (FIX), who have switched from prior bypassing agent (BPA) prophylaxis.

The study has 3 periods:

- * 6-Month BPA prophylaxis period in which patients will continue their prestudy, regularly scheduled prophylaxis regimen with BPAs
- * 1-Month onset period in which patients receive their first dose of 80 mg fitusiran while continuing their BPA prophylaxis for up to 7 days
- * 6-Month fitusiran efficacy period in which patients receive 80 mg fitusiran as a once monthly prophylaxis

Together, the 1-month onset period and the 6-month fitusiran efficacy period constitute the fitusiran treatment period.

Bleeding events and doses of BPAs administered during the conduct of the study will be recorded in an eDiary. Safety, quality of life, pharmacodynamic, and pharmacokinetic data will also be collected.

Following the screening and prophylaxis periods, all patients will be treated with fitusiran for a total of 7 months and will receive 7 SC injections of fitusiran. Because the full PD effect of fitusiran is not achieved until approximately 28 days after receiving the first dose, efficacy will be assessed during the final 6 months of the fitusiran treatment period (Day 29 to Month 7).

Throughout the study, patients may receive on-demand treatment for breakthrough bleeding episodes with BPAs. An independent data monitoring committee (DMC) will oversee the safety and overall conduct of this study. The DMC will perform periodic reviews of data during the course of the clinical trial, and on an ad hoc basis for review of emergent safety data, as defined in the DMC Charter for this clinical trial.

Patients who complete the study may be eligible for participation in an open-label extension study. For patients who do not enroll in the extension study, AT activity level will be monitored at monthly intervals following the final fitusiran dose until activity levels return to approximately 60% per the central laboratory, or per Investigator discretion in consultation with the study Medical Monitor.

Intervention

Diagnosis and Main Eligibility Criteria

This study will include males with severe hemophilia A or B with inhibitors, aged ≥ 12 years, who have been prescribed prophylactic treatment with BPAs for at least 6 months prior to Screening. Diagnosis of severe hemophilia A or B will be based on a central laboratory measurement or documented medical record evidence of FVIII level $<1\%$ or FIX level $\geq 2\%$. Patients with inhibitors must have used BPAs on demand to manage bleeding episodes for at least the last 6 months prior to Screening and must meet one of the following Nijmegen-modified Bethesda assay results criteria: 1) Inhibitor titer of ≥ 0.6 BU/mL at Screening, OR 2) Inhibitor titer of <0.6 BU/mL at Screening with medical record evidence of 2 consecutive titers ≥ 0.6 BU/mL, OR 3) Inhibitor titer of <0.6 BU/mL at Screening with medical record evidence of anamnestic response. A minimum of 2 bleeding episodes requiring BPA treatment within the last 6 months prior to Screening is required.

Investigational Product, Dose and Mode of Administration

Fitusiran is a subcutaneously (SC) administered GalNAc-conjugated siRNA targeting liver-expressed messenger RNA (mRNA) for AT. Patients will receive open-label fitusiran 80 mg as an SC injection once monthly, for a total of 7 months; dosing will begin on Day 1 of the fitusiran treatment period.

Reference Therapy, Dose and Mode of Administration

During the BPA prophylaxis period, patients will continue BPA prophylaxis as treatment for hemophilia on a regimen consistent with recommendations in the approved prescribing information, allowing for adjustment to individual patient response, and designed to decrease spontaneous bleeding.

Dose and mode of administration will be per Investigator discretion; bleeding episode management should be per the local standard practice for episodic use of BPAs and as per Investigator discretion.

Patients will continue to receive BPA prophylaxis for the first 7 days of the onset period. Subsequently, breakthrough bleeding episodes will be treated with on-demand BPA therapy as necessary per the bleeding episode management guidelines.

Duration of Treatment

The duration of treatment with fitusiran is 7 months. The estimated total time on study, inclusive of Screening, for each patient is up to 15 months for patients who enroll in the extension study. The estimated total time on study may be up to 21 months in patients who do not enroll in the extension study due to the requirement for an additional 6 months of follow-up for monitoring of AT levels.

Study burden and risks

see schedule of event in the protocol and section intervention above

Contacts

Public

Alnylam Pharmaceuticals Inc.

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Cambridge MA 02142
US

Scientific

Alnylam Pharmaceuticals Inc.

300 Third Street
Cambridge MA 02142
US

Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

Adolescents (12-15 years)
Adolescents (16-17 years)
Adults (18-64 years)
Elderly (65 years and older)

Inclusion criteria

1. Males *12 years of age.
2. Severe hemophilia A or B as evidenced by:
 - a . A central laboratory measurement or documented medical record evidence of FVIII <1% or FIX level *2% at Screening.
3. A minimum of 2 bleeding episodes requiring BPA treatment within the last 6 months prior to Screening.
4. Must meet the definition of inhibitor patient as below:
Use of BPAs for prophylaxis and for any bleeding episodes for at least the last 6 months prior to Screening, and meet one of the following Nijmegen-modified Bethesda assay results criteria:

- Inhibitor titer of ≥ 0.6 BU/mL at Screening, or
 - Inhibitor titer of < 0.6 BU/mL at Screening with medical record evidence of 2 consecutive titers ≥ 0.6 BU/mL, or
 - Inhibitor titer of < 0.6 BU/mL at Screening with medical record evidence of anamnestic response
5. Prescribed prophylactic treatment (documented in the medical or pharmacy records) of hemophilia with BPAs for at least 6 months prior to Screening; the regimen must be consistent with the approved prescribing information for the product or local recommendations, allowing for adjustment to individual patient response, and designed to decrease spontaneous bleeding.
 6. Adherent to the prescribed prophylactic therapy for at least 6 months prior to Screening per Investigator assessment.
 7. Willing and able to comply with the study requirements and to provide written informed consent and assent in the case of patients under the age of legal consent, per local and national requirements.

Exclusion criteria

1. Known co-existing bleeding disorders other than hemophilia A or B, ie, Von Willebrand's disease, additional factor deficiencies, or platelet disorders.
2. Current participation in immune tolerance induction therapy (ITI).
3. AT activity $< 60\%$ at Screening, as determined by central laboratory measurement
4. Presence of clinically significant liver disease, or as indicated by any of the conditions below:
 - a. INR > 1.2 ;
 - b. ALT and/or AST $> 1.5 \times$ upper limit of normal reference range (ULN);
 - c. Total bilirubin $> \text{ULN}$ ($> 1.5 \text{ ULN}$ in patients with Gilbert's Syndrome);
 - d. History of portal hypertension, esophageal varices, or hepatic encephalopathy;
 - e. Presence of ascites by physical exam.
5. Hepatitis C virus antibody positive, except patients with a history of HCV infection who meet both conditions a. and b.:
 - a. Completed curative treatment at least 12 weeks prior to enrollment and attained sustained virologic response as documented by a negative HCV RNA at screening, or they have spontaneously cleared infection as documented by negative HCV RNA at Screening.
 - b. No evidence of cirrhosis according to one of the following assessments:
 - * FibroScan < 12.5 kPa (where available), or
 - * FibroTest score < 0.75 and APRI < 2 (if FibroScan unavailable)
6. Presence of acute hepatitis, ie, hepatitis A, hepatitis E.
7. Presence of acute or chronic hepatitis B infection (IgM anti-HBc antibody positive or HBsAg positive).
8. Platelet count $\geq 100,000/\text{L}$.
9. Presence of acute infection at Screening.
10. Known to be HIV positive with CD4 count < 200 cells/ L .
11. Estimated glomerular filtration rate ≥ 45 mL/min/ 1.73m^2 (using the Modification of Diet in Renal Disease [MDRD] formula).

12. Co-existing thrombophilic disorder, as determined by presence of any of the below as identified at central laboratory (or via historical results, where available):
 - a. FV Leiden mutation (homozygous or heterozygous)
 - b. Protein S deficiency
 - c. Protein C deficiency
 - d. Prothrombin mutation (G20210A; homozygous and heterozygous)
13. History of antiphospholipid antibody syndrome.
14. History of arterial or venous thromboembolism, atrial fibrillation, significant valvular disease, myocardial infarction, angina, transient ischemic attack, or stroke. Patients who have experienced thrombosis associated with indwelling venous access may be enrolled.
15. Had a malignancy within 2 years, except for basal or squamous cell carcinoma of the skin that has been successfully treated.
16. Any condition (eg, medical concern), which in the opinion of the Investigator, would make the patient unsuitable for dosing on Day 1 or which could interfere with the study compliance, the patient's safety and/or the patient's participation in the completion of the treatment period of the study. This includes significant active and poorly controlled (unstable) cardiovascular, neurologic, gastrointestinal, endocrine, renal or psychiatric disorders unrelated to hemophilia identified by key laboratory abnormalities or medical history.
17. At Screening, anticipated need of surgery during the study or planned surgery scheduled to occur during the study.
18. Completion of a surgical procedure within 14 days prior to Screening, or currently receiving additional factor concentrate or BPA infusion for postoperative hemostasis.
19. History of multiple drug allergies or history of allergic reaction to an oligonucleotide or GalNAc.
20. Inadequate venous access, as determined by the Investigator, to allow the blood draws required by the study protocol.
21. History of intolerance to SC injection(s).
22. Current or future participation in another clinical study, scheduled to occur during this study, involving an investigational product other than fitusiran or an investigational device; in order to participate in this study, patient must discontinue the investigational product or investigational device at least 30 days (or 5× the investigational product half-life, whichever is longer) prior to dosing (Day 1).
23. Current or prior participation in a gene therapy trial.
24. History of alcohol abuse within the 12 months before Screening. Alcohol abuse is defined as regular weekly intake of more than 14 units (unit: 1 glass of wine [approximately 125 mL] = 1 measure of spirits [approximately 1 fluid ounce] = * pint of beer [approximately 284 mL]).

Study design

Design

Study phase:	3
Study type:	Interventional
Intervention model:	Other
Allocation:	Randomized controlled trial
Masking:	Open (masking not used)
Control:	Active
Primary purpose:	Prevention

Recruitment

NL	
Recruitment status:	Will not start
Enrollment:	2
Type:	Anticipated

Medical products/devices used

Product type:	Medicine
Brand name:	Fitusiran
Generic name:	ALN-AT3SC

Ethics review

Approved WMO	
Date:	03-05-2018
Application type:	First submission
Review commission:	CCMO: Centrale Commissie Mensgebonden Onderzoek (Den Haag)
Not approved	
Date:	17-05-2018
Application type:	First submission
Review commission:	CCMO: Centrale Commissie Mensgebonden Onderzoek (Den Haag)

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	EUCTR2016-004087-19-NL
CCMO	NL63089.000.18