Vutrisiran Clinical Development Program

Vutrisiran is an RNA interference (RNAi) therapeutic administered via subcutaneous injection once every three months (quarterly) for the treatment of the polyneuropathy of hereditary transthyretin-mediated (hATTR) amyloidosis in adults. Where it is approved, it is marketed as AMVUTTRA™ (vutrisiran).

Vutrisiran is also being evaluated as an investigational therapy in the HELIOS-B\(^1\) Phase 3 study for the treatment of adult patients with transthyretin-mediated (ATTR) amyloidosis with cardiomyopathy, including both hATTR and wild-type ATTR (wtATTR) amyloidosis.

**HELIOS-A**

HELIOS-A is a Phase 3 global, randomized, open-label study to evaluate the efficacy and safety of vutrisiran in adult patients with hATTR amyloidosis with polyneuropathy.\(^2\)

**Study Status**
- Enrollment is complete with 164 patients.

**Study Design**
- Patients were randomized 3:1 to receive either 25 mg of vutrisiran via subcutaneous injection once every three months or 0.3 mg/kg of patisiran via IV infusion once every three weeks (as a reference group), for 18 months. Following the 18-month treatment period, patients were eligible to receive either 25 mg vutrisiran once every three months or 50 mg vutrisiran once every six months for an additional 18 months as part of the randomized treatment extension period.\(^3\)
- For the primary and most secondary and exploratory efficacy endpoints, the vutrisiran arm is compared to the placebo arm of the APOLLO Phase 3 study.\(^3,4\)

**Primary Endpoint**
The primary endpoint of HELIOS-A was the change from baseline in the modified Neuropathy Impairment Score +7 (mNIS+7) at 9 months.

**Secondary Endpoints**

<table>
<thead>
<tr>
<th>Secondary Endpoint</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>Change from baseline in Norfolk Quality of Life-Diabetic Neuropathy (QOL-DN) Score at 9 and 18 months</td>
<td>The Norfolk QoL-DN questionnaire is a standardized 35-item patient-reported outcomes measure that is sensitive to the different features of diabetic neuropathy – small fiber, large fiber, and autonomic nerve function, symptoms, and activities of daily living – which may impact quality of life. It is validated for hATTR amyloidosis with polyneuropathy. The minimum and maximum values are -4 and 136, respectively. A higher score indicates a worse outcome.(^5,6)</td>
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<tr>
<td>Change from baseline in timed 10-meter walk test (10-MWT) at 9 and 18 months</td>
<td>A test of ambulatory function that measures a patient’s speed in walking 10 meters.(^8)</td>
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<tr>
<td>Change from baseline in modified Neuropathy Impairment Score +7 (mNIS+7) at 18 months</td>
<td>The mNIS+7 is a composite score that quantifies motor, sensory, and autonomic neurologic impairment due to injury of large and small nerves. The minimum and maximum values are 0 and 304, respectively. A higher score indicates a worse outcome.(^9)</td>
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<tr>
<td>Change from baseline in modified Body Mass Index (mBMI) at 18 months</td>
<td>A measure of nutritional status calculated as the product of body mass index and serum albumin.(^4,10) Lower mBMI indicates worse nutritional status.</td>
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<tr>
<td>Change from baseline in Rasch-built Overall Disability Scale (R-ODS) at 18 months</td>
<td>R-ODS is comprised of a 24-item linearly weighted scale that specifically captures activity and social participation limitations. The minimum and maximum values are 0 and 48, respectively. A higher score indicates a better outcome.(^4,11)</td>
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<tr>
<td>Percentage reduction in serum transthyretin (TTR) levels through 18 months</td>
<td>Unlike other endpoints, for this measure the vutrisiran arm will be compared to the within-study patisiran arm.(^7)</td>
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HELIOS-B

HELIOS-B is a global, Phase 3, randomized, double-blind, placebo-controlled study to evaluate the efficacy and safety of vutrisiran in adult patients with ATTR amyloidosis with cardiomyopathy (including both hATTR and wtATTR amyloidosis).1

Study Status
• Enrollment is complete with 655 patients.

Study Design
• Patients will be randomized on a 1:1 basis to receive either 25 mg of vutrisiran or placebo administered as a subcutaneous injection once every three months for up to 36 months.

Primary Endpoint
The primary endpoint will evaluate the efficacy of vutrisiran versus placebo on the composite endpoint of all-cause mortality and recurrent cardiovascular (CV) events (CV hospitalizations and urgent heart failure (HF) visits) at 30-36 months.

Secondary Endpoints

| Change from baseline in 6-minute walk test (6-MWT) at 30 months | An assessment of functional exercise capacity, measuring how far a patient can walk in six minutes along a prescribed course.1,12 |
| Change from baseline in Kansas City Cardiomyopathy Questionnaire Overall Summary (KCCQ-OS) at 30 months | The KCCQ is a 23-item self-administered questionnaire quantifying 6 domains (symptoms, physical function, quality of life, social limitation, self-efficacy, and symptom stability) and 2 summary scores (clinical and overall summary [OS]). Scores are transformed to a range of 0-100, in which higher scores reflect better health status.1 |
| Change from baseline in mean left ventricular (LV) wall thickness and global longitudinal strain at 30 months | Echocardiographic assessments of cardiac structure and function.1 |
| Composite endpoint of all-cause mortality and recurrent all-cause hospitalizations and urgent HF visits at 30-36 months | All-cause mortality, recurrent all-cause hospitalizations and urgent HF visits.1 |
| All-cause mortality at 30-36 months | Deaths from any cause.1 |
| Rate of recurrent CV events at 30-36 months | Recurrent CV events (CV hospitalizations and urgent HF visits).1 |
| Change from baseline in N-terminal prohormone B-type natriuretic peptide (NT-proBNP) at 30 months | A biomarker for the severity of heart failure.1,13 |

For more information on HELIOS-A (NCT03759379) and HELIOS-B (NCT04153149) please visit www.clinicaltrials.gov or contact media@alnylam.com.

Current information as of June 2022.