Long-term efficacy and safety of inotersen for hereditary transthyretin amyloidosis: NEURO-TTR open-label extension 3-year update

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Abstract

Background Hereditary transthyretin amyloidosis (hATTR/ATTRv) results from the deposition of misfolded transthyretin (TTR) throughout the body, including peripheral nerves. Inotersen, an antisense oligonucleotide inhibitor of hepatic TTR production, demonstrated a favorable efficacy and safety profile in patients with the polyneuropathy associated with hATTR in the NEURO-TTR (NCT01737398) study. We report longer-term efficacy and safety data for inotersen, with a median treatment exposure of 3 years.

Methods Patients who satisfactorily completed NEURO-TTR were enrolled in its open-label extension (OLE) study. Efficacy assessments included the modified Neuropathy Impairment Score + 7 (mNIS + 7), Norfolk Quality of Life–Diabetic Neuropathy (Norfolk QoL-DN) questionnaire total score, and the Short Form 36 (SF-36v2) Health Survey Physical Component Summary score. Safety and tolerability were also assessed. Efficacy is reported for patients living in Europe and North America (this cohort completed the study approximately 9 months before the remaining group of patients outside these regions); safety is reported for the full safety dataset, comprising patients living in Europe, North America, and Latin America/Australasia. This study is registered with ClinicalTrials.gov, identifier NCT02175004.

Results In the Europe and North America cohort of the NEURO-TTR study, 113/141 patients (80.1%) completed the study, and 109 patients participated in the OLE study. A total of 70 patients continued to receive inotersen (inotersen–inotersen) and 39 switched from placebo to inotersen (placebo–inotersen). The placebo–inotersen group demonstrated sustained improvement in neurological disease progression as measured by mNIS + 7, compared with predicted worsening based on projection of the NEURO-TTR placebo data (estimated natural history). The inotersen–inotersen group demonstrated sustained benefit, as measured by mNIS + 7, Norfolk QoL-DN, and SF-36v2, compared with estimated natural history as well as compared with the placebo–inotersen group. With a maximum exposure of 6.2 years, inotersen was not associated with any additional safety concerns or increased toxicity in the OLE study. Platelet and renal monitoring were effective in reducing the risk of severe adverse events in the OLE study.

Conclusion Inotersen treatment for > 3 years slowed progression of the polyneuropathy associated with hATTR, and no new safety signals were observed.

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Co-author Merrill D. Benson, MD, died September 11, 2021.

A complete list of NEURO-TTR Open-Label Extension investigators is provided in Online Resource 6.

Extended author information available on the last page of the article
Keywords Clinical trial · Familial amyloid polyneuropathy · Inotersen · Hereditary transthyretin amyloidosis · Peripheral neuropathies · Polyneuropathy

Introduction

Hereditary transthyretin amyloidosis (hATTR or ATTRv [variant]) is a rapidly progressive, debilitating, and ultimately fatal disease resulting from the accumulation of transthyretin (TTR) amyloid fibrils in peripheral nerves and various other organs and tissues throughout the body [1]. It is caused by mutations in the TTR gene that are thought to induce changes in the protein’s normally tetrameric structure, such that it dissociates to monomer subunits which then misfolds and have a greater propensity for aggregation and the formation of amyloid fibrils [1, 2]. In patients with hATTR, systemic deposition of amyloid fibrils causes debilitating sensorimotor peripheral neuropathy and autonomic neuropathy, cardiomyopathy with associated heart failure, as well as ocular and renal disturbances [1, 3–6]. hATTR can manifest with a predominant polyneuropathy or cardiomyopathy phenotype, or a mixture of both, depending on the TTR variant [5, 7, 8].

Early signs/symptoms of hATTR include pain and loss of temperature sensation, along with digestive problems and genitourinary dysfunction [9, 10]. As the disease course progresses, patients experience muscle weakness and balance abnormalities [10]. If left untreated, polyneuropathy associated with hATTR has a rapidly progressive course, with associated walking difficulties and loss of ambulation as well as malnutrition [11].

Patients experience substantial disease burden and a rapid decline in quality of life (QoL), with progressive neuropathy, cardiomyopathy, and psychosocial manifestations affecting every aspect of their lives [11–15]. Therefore, early diagnosis and implementation of treatment is imperative to slow progression of the disease [16]. In the global, randomized, double-blind, placebo-controlled pivotal NEURO-TTR trial (NCT01737398) [17] consisted of a ≤ 4-week screening period, a treatment period of up to 260 weeks, and a 12-week post-treatment evaluation period. The study design and inclusion/exclusion criteria for the NEURO-TTR [17] and NEURO-TTR OLE studies have been previously published [18]. Briefly, patients with stage 1 or 2 polyneuropathy associated with hATTR and a Neuropathy Impairment Score ≥ 10 and ≤ 130 who satisfactorily completed the randomized NEURO-TTR study could enter the OLE study and receive 300 mg inotersen once weekly via subcutaneous injection for up to 260 weeks (5 years). One patient who did not complete the NEURO-TTR study was nonetheless allowed to enter the OLE. Patients receiving inotersen in NEURO-TTR continued to receive inotersen in the OLE study and are hereafter referred to as the inotersen–inotersen group; patients receiving placebo in NEURO-TTR switched to inotersen in the OLE study and are hereafter referred to as the placebo–inotersen group.

The trial protocol for the OLE study was approved by the relevant institutional review boards or local ethics committees and regulatory authorities. The trial was conducted in accordance with Good Clinical Practice guidelines of the International Conference on Harmonisation and the principles of the Declaration of Helsinki. All patients provided written informed consent to participate in the study.

Outcomes

The primary objective of the OLE study was the evaluation of the safety of extended dosing with inotersen. Safety assessments included the incidence of adverse events (AEs) and treatment-emergent AEs (TEAEs) reported during the study per National Cancer Institute Common Terminology Criteria for Adverse Events, version 4.03. The change in TTR levels from baseline up to week 156 was determined during the study. Efficacy endpoints included change from baseline up to week 156 in the (a) modified Neuropathy Impairment Score + 7 composite score (mNIS + 7; range −22.3 to 346.3, with higher scores indicating poorer function), (b) Norfolk Quality of Life–Diabetic Neuropathy questionnaire scores (Norfolk QoL-DN; range −4 to 136, with higher scores indicating poorer QoL), and (c) 36-Item
Short-Form Health Survey (SF-36) questionnaire scores (range, 0 to 100; lower score indicating worse QoL).

Statistical analysis

The OLE baseline values were carried forward from the week 65 visit of the NEURO-TTR study for the SF-36 and from the week 66 visit for mNIS + 7 and QoL-DN measures. Responders were defined as patients whose change in score was less than the responder definition (RD) threshold for the mNIS + 7 and Norfolk QoL-DN measures and greater than the RD threshold for the SF-36; that is, their change in score did not indicate clinical worsening (i.e., responders included both patients with no clinically relevant change in scores plus patients who improved with treatment). Using data from the NEURO-TTR study, the RD thresholds the Norfolk QOL-DN (8.8 points) were estimated using both anchor-based and distribution-based methods and for the mNIS + 7 (12.2 points) were estimated using distribution-based methods only, because there were no appropriate anchor measures available [19]. The threshold for the SF-36 PCS (−5 points) was based on studies from the literature that utilized both anchor-based and distribution-based methods [20–22]. For each estimated RD threshold, responder analyses compared the proportion of patients on the mNIS + 7, Norfolk QOL-DN, and SF-36 PCS between inotersen—inotersen and placebo—inotersen groups at each visit up to week 156.

As of July 28, 2020, efficacy was reported for the Europe and North America cohort that received ≥1 dose of inotersen in the OLE and had ≥1 post-baseline efficacy assessment (full analysis set). This cohort completed database lock approximately 9 months before the remaining group of patients in the Latin America/Australasia cohort. For safety analyses, data are reported for the full safety dataset, which comprises patients from Europe, North America, and Latin America/Australasia.

Results

Study population

A total of 139 (80%) patients completed the NEURO-TTR study, including 113 patients in the Europe and North America cohort [17]. Of the 139 patients who completed the NEURO-TTR study, 135 (97%) participated in the OLE study, of whom 109 patients comprised the Europe and North America cohort. A total of 70 patients in the Europe and North America cohort continued to receive inotersen (inotersen—inotersen) and 39 switched from placebo to inotersen (placebo—inotersen) (Online Resource 1, Fig. 1). The primary reasons for treatment discontinuation are specified in Online Resource 1, Fig. 1. Of the 108 patients who discontinued the OLE study, 43 (39.8%) switched to a commercial drug (Tegsedi®): 25 patients from the inotersen—inotersen treatment group and 18 patients from the placebo—inotersen group. In the OLE study, baseline demographics and clinical characteristics were generally balanced between the inotersen—inotersen and placebo—inotersen groups (Table 1). A higher proportion of patients in the inotersen—inotersen group had a polyneuropathy disability score of I/II at baseline compared with the placebo—inotersen group (65.7% vs. 53.8%).

With extended follow-up, median treatment exposure for the safety population in the OLE study was 987 days (range 1–1814 days), and the longest combined exposure of inotersen during the NEURO-TTR plus OLE study, measured as the time from the first dose to the last dose, was 2270 days (6.2 years). Median treatment exposure for
the inotersen—inotersen and placebo—inotersen groups was 1063 days (1–1814 days) and 938 days (106–1728 days), respectively. Dose interruption and reasons for dose interruption are provided in Online Resource 2.

**Pharmacodynamics**

In the OLE study, reduction in TTR level was sustained in the inotersen—inotersen group to week 156, reaching a median nadir of 73% relative to NEURO-TTR baseline (Fig. 1). In the placebo—inotersen group, there was a notable reduction in TTR level by OLE week 7, when the first measurement was taken (Fig. 1). The reduction continued through week 156 in the OLE study, reaching a median nadir of 66% relative to OLE baseline.

**Efficacy**

**Neuropathy impairment (mNIS + 7)**

The inotersen—inotersen group demonstrated sustained benefit, as measured by mNIS + 7, where higher scores are indicative of poorer neurological function, compared with the placebo—inotersen group (Fig. 2A). The mean change from NEURO-TTR study baseline to OLE study baseline, as well as weeks 52, 104, and 156 for the mNIS + 7, was 4.0, 9.5, 17.8, and 19.2, respectively, in the inotersen—inotersen group. The placebo—inotersen group demonstrated sustained improvement in neurological disease progression, compared with predicted worsening based on a projection of the placebo data (estimated natural history). The corresponding mean change from NEURO-TTR...
Fig. 2  Mean change from NEURO-TTR baseline to OLE week 156 in efficacy measures. Mean (± SE) change from NEURO-TTR baseline in A the Modified Neuropathy Impairment Score + 7 (mNIS + 7); B the Norfolk Quality of Life–Diabetic Neuropathy Questionnaire Total Score (QoL-DN); C the 36-Item Short-Form Health Survey, version 2 (SF-36), Physical Component Summary score (PCS). The vertical dashed line represents OLE baseline (OLE week 0). Sample sizes for each time point and treatment group are indicated under the figure. The OLE baseline values were carried forward from the week 65 visit of the NEURO-TTR study for the SF-36 and from the week 66 visit for mNIS + 7 and QoL-DN measures. The OLE baseline on the graph is noted as occurring 4 weeks after the end of the NEURO-TTR study because the maximum screening period in the OLE study was generally 4 weeks. OLE, open-label extension, SE, standard error.
### Percent of Responders (∆mNIS+7 < 12.2)

| Time (weeks) | Placebo-inotersen, n | Inotersen-inotersen, n |
|--------------|----------------------|------------------------|
| Baseline     | 38                   | 67                     |
| 1            | 39                   | 67                     |
| 2            | 39                   | 66                     |
| 3            | 36                   | 65                     |
| 4            | 26                   | 60                     |
| 5            | 22                   | 56                     |
| 6            | 22                   | 45                     |
| 7            | 20                   | 39                     |
| 8            | 16                   | 30                     |

### Percent of Responders (∆Norfolk QOL-DN < 8.8)

| Time (weeks) | Placebo-inotersen, n | Inotersen-inotersen, n |
|--------------|----------------------|------------------------|
| Baseline     | 39                   | 66                     |
| 1            | 39                   | 65                     |
| 2            | 39                   | 65                     |
| 3            | 35                   | 62                     |
| 4            | 28                   | 58                     |
| 5            | 23                   | 50                     |
| 6            | 25                   | 44                     |
| 7            | 21                   | 35                     |

### Percent of Responders (∆PCS > 5)

| Time (weeks) | Placebo-inotersen, n | Inotersen-inotersen, n |
|--------------|----------------------|------------------------|
| Baseline     | 39                   | 66                     |
| 1            | 39                   | 64                     |
| 2            | 38                   | 63                     |
| 3            | 34                   | 62                     |
| 4            | 27                   | 58                     |
| 5            | 24                   | 49                     |
| 6            | 23                   | 43                     |
| 7            | 20                   | 35                     |
Responder analyses for the NEURO-TTR Europe and North America cohort. Data are presented for A the Modified Neuropathy Impairment Score +7 Neurophysiological Tests Composite Score (mNIS +7), B the Norfolk Quality of Life–Diabetic Neuropathy Questionnaire Total Score (QoL-DN), and C the 36-Item Short-Form Health Survey, version 2 (SF-36), Physical Component Summary score (PCS). A patient was classified as a responder at various time points through week 156 if they showed no clinically meaningful progression based on the applicable threshold and had a recorded measurement at baseline and at the relevant time point. The OLE baseline on the graph is noted as occurring 4 weeks after the end of the NEURO-TTR study because the maximum screening period in the OLE study was generally 4 weeks. A For the inotersen–inotersen group, responders at all visits are based on change <12.2 points from NEURO-TTR baseline. For the placebo–inotersen group, responders at OLE-TTR week 35 and week 66 visits and OLE baseline visits are based on change <12.2 points from NEURO-TTR baseline, while responders at all post-baseline OLE visits are based on change <12.2 points from OLE baseline. B For the inotersen–inotersen group, responders at all visits are based on change <8.8 points from NEURO-TTR baseline. For the placebo–inotersen group, responders at NEURO-TTR week 35 and week 66 visits and OLE baseline visits are based on change <8.8 points from NEURO-TTR baseline, while responders at all post-baseline OLE visits are based on change <8.8 points from OLE baseline. C For the inotersen–inotersen group, responders at all visits are based on change >–5 points from NEURO-TTR baseline. For the placebo–inotersen group, responders at NEURO-TTR week 35 and week 66 visits and OLE baseline visits are based on change >–5 points from NEURO-TTR baseline, while responders at all post-baseline OLE visits are based on change >–5 points from OLE baseline. 

baseline for the placebo–inotersen group was 25.3, 30.5, 33.4, and 40.6. Compared with projected natural history data, the mean change from NEURO-TTR baseline for mNIS +7 was 62.2 points lower in the inotersen–inotersen group and 40.7 points lower in the placebo–inotersen group at week 156.

Using a responder definition of a change of < 8.8 points in the Norfolk QoL-DN relative to NEURO-TTR baseline, improvement occurred in 61%, 60%, and 49% of patients in the inotersen–inotersen group after 52, 104, and 156 weeks of cumulative treatment during the OLE study (Fig. 3B). At the same time points, improvement in the Norfolk QoL-DN occurred in 75%, 68%, and 65% of patients, respectively (Fig. 3B) in the placebo–inotersen group, relative to OLE study baseline. Improvement of Norfolk QoL-DN was also observed for both inotersen–inotersen and placebo–inotersen groups as indicated by a < 0.0-point change (Online Resource 3).

The improvement in Norfolk QoL-DN compared with natural history was maintained for the inotersen–inotersen and placebo–inotersen groups for stage 1 disease (but not stage 2 disease), those both with and without a V30M mutation, and those receiving prior treatment (with or without stabilizers) (Online Resource 4). Across these three baseline characteristics, there was no clear separation over time between the inotersen–inotersen and placebo–inotersen groups.

Health-related QoL (SF-36 PCS)

The SF-36 is a non-disease-specific health questionnaire that assesses the patient’s perceived functional health, and higher scores are indicative of better function. As seen in the inotersen–inotersen group, early initiation of inotersen led to sustained benefit in health-related QoL compared with the placebo–inotersen group (Fig. 2C). For the inotersen–inotersen group, the mean change from NEURO-TTR baseline to OLE baseline and 52, 104, and 156 weeks was 11.6, 9.0, 12.3, and 17.3, respectively. Relative to the placebo-slope extrapolation, the mean change from NEURO-TTR baseline to OLE week 156 was 23.5 points lower in the inotersen–inotersen group and 15.9 points lower in the placebo–inotersen group.

Using a responder definition of a change of < 8.8 points in the Norfolk QoL-DN relative to NEURO-TTR baseline, improvement occurred in 61%, 60%, and 49% of patients in the inotersen–inotersen group after 52, 104, and 156 weeks of cumulative treatment during the OLE study (Fig. 3B). At the same time points, improvement in the Norfolk QoL-DN occurred in 75%, 68%, and 65% of patients, respectively (Fig. 3B) in the placebo–inotersen group, relative to OLE study baseline. Improvement of Norfolk QoL-DN was also observed for both inotersen–inotersen and placebo–inotersen groups as indicated by a < 0.0-point change (Online Resource 3).

The improvement in Norfolk QoL-DN compared with natural history was maintained for the inotersen–inotersen and placebo–inotersen groups for stage 1 disease (but not stage 2 disease), those both with and without a V30M mutation, and those receiving prior treatment (with or without stabilizers) (Online Resource 4). Across these three baseline characteristics, there was no clear separation over time between the inotersen–inotersen and placebo–inotersen groups.

Neuropathy-related QoL (Norfolk QoL-DN)

The inotersen–inotersen group demonstrated sustained benefit compared with the placebo–inotersen group, as measured by the Norfolk QoL-DN questionnaire in which higher scores are also indicative of poorer function (Fig. 2B). For the inotersen–inotersen group, the mean changes in the Norfolk QoL-DN from NEURO-TTR baseline to OLE baseline, 52, 104, and 156 weeks were 1.9, 5.1, 4.7, and 9.8, respectively. The placebo–inotersen group demonstrated stabilization in neuropathy-related QoL. The mean changes in Norfolk QoL-DN from NEURO-TTR baseline to OLE baseline and 52, 104, and 156 weeks were 11.6, 9.0, 12.3, and 17.3, respectively. Relative to the placebo-slope extrapolation, the mean change from NEURO-TTR baseline to OLE week 156 was 23.5 points lower in the inotersen–inotersen group and 15.9 points lower in the placebo–inotersen group.

Using a responder definition of a change of < 8.8 points in the Norfolk QoL-DN relative to NEURO-TTR baseline, improvement occurred in 61%, 60%, and 49% of patients in the inotersen–inotersen group after 52, 104, and 156 weeks of cumulative treatment during the OLE study (Fig. 3B). At the same time points, improvement in the Norfolk QoL-DN occurred in 75%, 68%, and 65% of patients, respectively (Fig. 3B) in the placebo–inotersen group, relative to OLE study baseline. Improvement of Norfolk QoL-DN was also observed for both inotersen–inotersen and placebo–inotersen groups as indicated by a < 0.0-point change (Online Resource 3).

The improvement in Norfolk QoL-DN compared with natural history was maintained for the inotersen–inotersen and placebo–inotersen groups for stage 1 disease (but not stage 2 disease), those both with and without a V30M mutation, and those receiving prior treatment (with or without stabilizers) (Online Resource 4). Across these three baseline characteristics, there was no clear separation over time between the inotersen–inotersen and placebo–inotersen groups.
PCS scores were 8.6 points higher in the inotersen–inotersen group and 4.9 points higher in the placebo–inotersen group.

Using a responder definition of a change of −5 points relative to the NEURO-TTR baseline, the percentage of responders based on the SF-36 PCS was 76%, 76%, and 80% of patients in the inotersen–inotersen group after 52, 104, and 156 weeks of cumulative treatment during the OLE study (Fig. 3C). At similar time points for patients in the placebo–inotersen group, the percentage of responders based on the SF-36 PCS was 81%, 74%, and 75% (Fig. 3C) relative to OLE study baseline. Improvement in SF-36 PCS was also observed for both the inotersen–inotersen and placebo–inotersen groups as indicated by a >0.0-point change (Online Resource 3).

The improvement in SF-36 PCS compared with natural history was maintained for the inotersen–inotersen and placebo–inotersen groups irrespective of genetics (V30M mutation vs. no V30M mutation), disease stage (stage 1 vs. stage 2), and prior treatment (with or without stabilizers) (Online Resource 4). Across these three baseline characteristics, there was no clear separation over time between the inotersen–inotersen and placebo–inotersen groups.

Safety and tolerability

Safety of inotersen in the OLE study is summarized in Table 2. The most common (≥10%) TEAEs across all patients in the OLE were thrombocytopenia (28.1%), diarrhea (25.9%), nausea (25.2%), urinary tract infection (24.2%), fall (22.2%), fatigue (20.0%), vomiting (20.0%), chills (17.8%), peripheral edema (17.0%), injection site pain (13.3%), injection site erythema (11.9%), constipation (11.9%), syncope (11.1%), dyspnea (11.1%), headache (11.1%), pyrexia (10.4%), and anemia (10.4%).

| Event, n (%) | Total (n=135) |
|-------------|--------------|
| Treatment exposure during OLE study, median (range), days | 987 (1–1814) |
| Any TEAEs | 132 (97.8) |
| TEAEs related to study treatment | 99 (73.3) |
| TEAEs leading to permanent discontinuation of inotersen | 30 (22.2) |
| TEAEs leading to dose reduction | 13 (9.6) |
| TEAEs leading to dose interruption/delay | 66 (48.9) |
| Serious TEAEs | 64 (47.4) |
| Serious TEAEs related to study treatment | 6 (4.4) |
| Fatal TEAEs | 16 (11.9) |
| Fatal TEAEs related to study treatment | 0 |

OLE open-label extension, TEAE treatment-emergent adverse event

*Represents all patients from NEURO-TTR OLE

*Includes time on drug only; excludes dose holidays

TEAEs leading to dose interruption, dose reduction, or study drug delay occurred in 79 (58.5%) patients. Overall, 30 (22.2%) patients discontinued because of a TEAE: 23 (27.1%) patients in the inotersen–inotersen group (thrombocytopenia [n=4], glomerulonephritis [n=1]) and seven (14.0%) patients in the placebo–inotersen group.

A total of 64 (47.4%) patients experienced a serious TEAE: 46 (54.1%) patients in the inotersen–inotersen group and 18 (36.0%) patients in the placebo–inotersen group. Few were considered related to inotersen (n=6; 4.4%); these included thrombocytopenia (n=2 in the inotersen–inotersen group), nausea (n=1 in the placebo–inotersen group), chills (n=1 in the inotersen–inotersen group), liver disorder (n=1 in the inotersen–inotersen group), and hypertension (n=1 in the inotersen–inotersen group). Study drug was interrupted for both events of thrombocytopenia (platelet count nadir of 32 × 10^9/L [day 350] and 43 × 10^9/L [day 519], respectively); neither patient permanently discontinued study drug nor withdrew from the study due to these events of thrombocytopenia. The declines in platelet count were reversed on interruption of dosing and the events were not associated with bleeding. For the event of nausea, the drug dose was interrupted, and the event recovered/resolved. For the event of hypertension, the drug was withdrawn and the event recovered/resolved. The event of chills was recovered/resolved without any changes to drug dose. For the event of liver disorder, elevations were predominantly seen in aspartate aminotransferase (AST) (less than 3× upper limit of normal [ULN]) and alkaline phosphatase (less than 2× ULN), with no increase in total bilirubin levels. Partial resolution of the enzyme elevations was observed during continued inotersen treatment and complete resolution occurred after discontinuation of treatment. Liver biopsy showed evidence of congestion thought to be secondary to transient hepatic venous outflow obstruction rather than hepatotoxicity.
Overall, 16 (11.9%) patients died (Table 2) in the OLE study, and none of these deaths were considered related to treatment.

**Thrombocytopenia and glomerulonephritis**

Platelet and renal monitoring were effective in the OLE study. In the safety population, 38% (32/85) of the inotersen–inotersen group and 52% (26/50) of the placebo–inotersen group had a dose interruption due to platelet monitoring rules. In the OLE study, 50% (25/50) of patients in the placebo–inotersen group and 48% (41/85) of patients in the inotersen–inotersen group experienced confirmed platelet count decreases of < 100 × 10⁹/L. Nonetheless, grade 4 platelet count decrease was reported in none of the patients in either group (Online Resource 5, Fig. 5). In addition, no cases of acute glomerulonephritis were reported in the two groups.

**Discussion**

Currently, patients with the polyneuropathy associated with hATTR have an estimated life expectancy of 5–10 years from symptom onset [23]. Peripheral nerve function in particular has been shown to deteriorate more rapidly in patients with the polyneuropathy associated with hATTR relative to other peripheral neuropathies [23]. Therefore, effective treatment options are crucial to slow down disease progression and improve patients’ QoL [12].

The results of this OLE study confirm that long-term exposure to inotersen resulted in continued efficacy after 3 years, with no additional safety concerns or signs of increased toxicity. Mean mNIS + 7 and Norfolk QoL-DN scores were relatively similar at NEURO-TTR and OLE study baseline for the inotersen–inotersen group; however, scores for the placebo–inotersen group indicated more severe disease at OLE study baseline because of disease progression while on placebo during the NEURO-TTR study.

The progression of polyneuropathy was substantially delayed in patients treated with inotersen versus predicted natural history over a 3-year follow-up. Importantly, patients who initiated treatment with inotersen earlier derived a greater benefit with respect to neurological impairment and QoL when compared with patients who received delayed treatment with inotersen (i.e., those who switched from placebo to inotersen approximately 15 months later). A previous analysis of specific aspects of QoL from the OLE showed that treatment with inotersen stabilized most aspects of QoL for up to 3 years including activities of daily living and physical and social functioning. Moreover, the gap in QoL between those who had previously received inotersen or placebo in NEURO-TTR did not close by week 104 of the OLE, indicating the importance of early treatment for maintaining HRQoL in patients with hATTR [24]. Together, these results confirm that early diagnosis and treatment are essential for maximal benefit. Delayed treatment provides benefit by slowing disease progression and improving QoL compared with natural history.

The emerging availability of effective therapeutic options for the polyneuropathy associated with hATTR has enhanced awareness and recognition of this debilitating disease and promoted expeditious implementation of treatment, even in patients with mild disease.

Enhanced monitoring has reduced the risk of severe thrombocytopenia and acute glomerulonephritis. With a maximum exposure of up to 6.2 years, inotersen was not associated with any additional safety concerns or increased toxicity in the OLE study. Platelet and renal monitoring have been effective in the OLE study: among all patients, there were no cases of grade 4 platelet count decrease or acute glomerulonephritis.

Limitations of the current study include its open-label design with no placebo control, which could influence patient responses to QoL assessments and, to a much lesser extent, mNIS + 7 assessments. Next, inferential statistics were not completed, and the results are thus qualitative. In addition, efficacy analyses included a subgroup of the overall OLE population (albeit a fairly large subgroup, consisting of 81% of study participants). Thus, subgroup efficacy analyses based on certain baseline characteristics are difficult to interpret. In addition, all OLE studies are associated with “survivor bias,” as only data on patients continuously enrolled can be analyzed; OLE analyses generally exclude patients who drop out and may have suffered different outcomes than the continuing cohort, depending on the reason for dropout. Nonetheless, the results of this study are important in determining the long-term efficacy and safety of inotersen, even under imperfect conditions.

In the NEURO-TTR OLE study, extended treatment with inotersen for over 3 years slowed progression of the polyneuropathy associated with hATTR and maintained QoL in patients, with greater benefit observed in patients who initiated inotersen earlier versus patients with delayed treatment initiation. These long-term results further highlight the importance of early initiation of treatment. Grade 4 thrombocytopenia and acute glomerulonephritis were not reported with enhanced monitoring during the OLE study.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s00415-022-11276-8.

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Author contributions All authors were involved in data interpretation, critically revising the manuscript, and have approved the final draft of the manuscript before submission. KO, PA, and AN had full access to de-identified patient-level data, while all authors had access to aggregate study data; all authors vouch for and are accountable for the accuracy and integrity of the publication. Data review committee investigators contributed to conception and design of the study. Study investigators TB, TC, AKW, MJP, PJD, JLB, BD, PG, CW, IC, VP-B, GM, LO, JMCP, JG, AVK, AM, LG, and MG were involved in the acquisition, analysis, and interpretation of the data.

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Data availability Data that underlie the results reported in this article and the respective individual participant data will not be shared.

Declarations

Conflicts of interest TB reports being an advisory board member for Akcea Therapeutics, Pfizer, and Alnylam Pharmaceuticals; study investigator for Ionis Pharmaceuticals, Inc., and Alnylam Pharmaceuticals; and speaker for Alnylam Pharmaceuticals; and receiving honoraria for speaking for Akcea Therapeutics. TC reports receiving financial support to attend scientific meetings from Pfizer, Alnylam, Ionis, and Biogen. AKW reports being study investigator, consultant, and speaker for Ionis Pharmaceuticals, Inc.; consultant for Pfizer; and study investigator for Alnylam. MJP reports receiving honoraria from Pfizer and Alnylam Pharmaceuticals, Inc. PJD reports receiving training and consulting honoraria from Ionis Pharmaceuticals, Inc., and Alnylam Pharmaceuticals. JLB reports receiving honoraria from Ionis Pharmaceuticals, Inc., and Alnylam Pharmaceuticals; and being a study investigator for Ionis Pharmaceuticals, Inc., Alnylam Pharmaceuticals, Eidos BridgeBio, and Pfizer. BD reports being a consultant/advisory board member for Eidos and Alnylam Pharmaceuticals. PG declares no competing interests. CW reports being an advisory board member for Alnylam and Akcea Therapeutics. IC reports being a consultant for Alnylam and Pfizer; receiving honoraria for Alnylam, Pfizer, and Sanofi; and being on speaker bureaus for Alnylam, Pfizer, and Sanofi. VP-B reports receiving consulting honoraria for Alnylam Pharmaceuticals, Ionis, and Pfizer and travel honoraria from Ionis. GM declares no competing interests. LO reports receiving speaker honoraria from Alnylam Pharmaceuticals, Alnylam Pharmaceuticals, SOBI, and Pfizer. JMCP declares no competing interests. JG declares no competing interests. AVK reports receiving honoraria and fees for lectures and speakers bureaus from Alnylam Pharmaceuticals, Akcea Therapeutics, and Pfizer Inc. AM reports receiving speaker fees and consulting honoraria from Alnylam Pharmaceuticals, Akcea Therapeutics, and Pfizer Inc. LG reports receiving speaker fees and consulting honoraria from Pfizer. AN is a former employee of Akcea Therapeutics, a subsidiary of Ionis Pharmaceuticals, Inc. KO is a former employee of Ionis Pharmaceuticals, Inc. PA is a former employee of Ionis Pharmaceuticals, Inc./Akcea Therapeutics. MDB reports being study investigator for Ionis Pharmaceuticals, Inc. PG reports receiving consulting honoraria from Ionis, AbbVie, Alnylam, Amgen, Annexon, Appellis, Celgene, Janssen, Medscape, Physicians Education Resource, Prothena, Research to Practice, Teva, and Spectrum, and grants from Spectrum.

Ethical approval The trial protocol for the OLE study was approved by the relevant institutional review boards or local ethics committees and regulatory authorities. The trial was conducted in accordance with Good Clinical Practice guidelines of the International Conference on Harmonisation and the principles of the Declaration of Helsinki. All patients provided written informed consent to participate in the study.

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