TRITON-CM: A Phase 3 Study to Evaluate the Efficacy and Safety of Nucresiran in Patients with Transthyretin Amyloidosis with Cardiomyopathy

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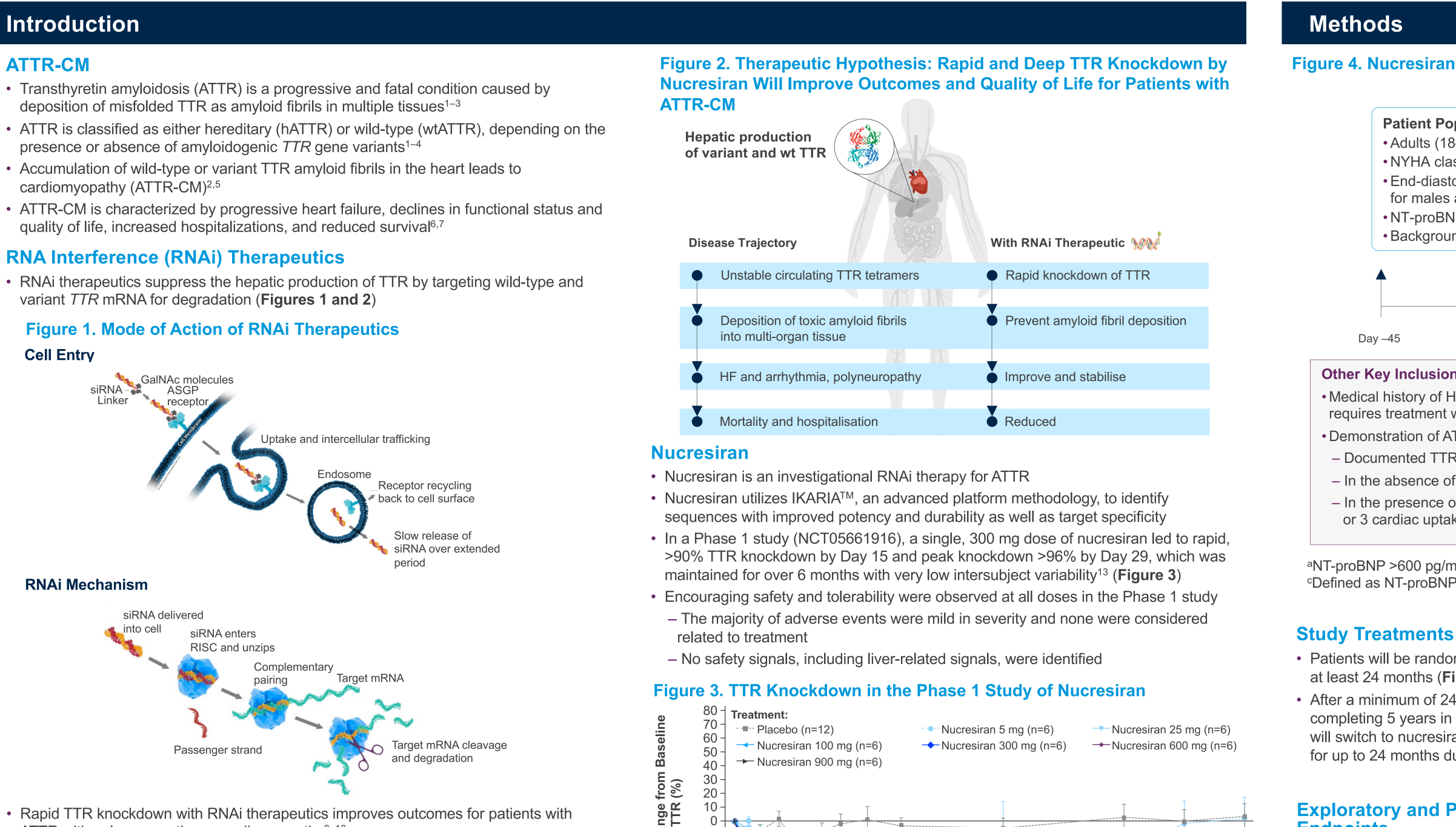
Conclusions

Introduction

ATTR-CM

- Transthyretin amyloidosis (ATTR) is a progressive and fatal condition caused by deposition of misfolded TTR as amyloid fibrils in multiple tissues^{1–3}
- ATTR is classified as either hereditary (hATTR) or wild-type (wtATTR), depending on the presence or absence of amyloidogenic *TTR* gene variants^{1–4}
- Accumulation of wild-type or variant TTR amyloid fibrils in the heart leads to cardiomyopathy (ATTR-CM)^{2,5}
- ATTR-CM is characterized by progressive heart failure, declines in functional status and quality of life, increased hospitalizations, and reduced survival^{6,7}

• RNAi therapeutics suppress the hepatic production of TTR by targeting wild-type and variant *TTR* mRNA for degradation (**Figures 1 and 2**)



- Rapid TTR knockdown with RNAi therapeutics improves outcomes for patients with ATTR with polyneuropathy or cardiomyopathy^{8–10}
- Most recently, the HELIOS-B study showed that vutrisiran knockdown of TTR improved outcomes for patients with ATTR-CM across multiple domains, including reducing CV events and all-cause mortality and improving functional capacity and quality of life¹⁰
- The positive effects observed in HELIOS-B add to the growing evidence that RNAi-based therapeutics that knockdown TTR can improve clinical outcomes in ATTR-CM
- Greater TTR knockdown offers the potential for further improving outcomes, as has been observed in other forms of amyloidosis^{11,12}

-10 -20

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-90

-100

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• TRITON-CM is a global, randomized, double-blind, event-driven Phase 3 CV outcomes study that will investigate the efficacy and safety of nucresiran in patients with transthyretin amyloidosis with cardiomyopathy (ATTR-CM) • Nucresiran is an investigational RNAi therapeutic with advanced target specificity that is capable of rapid, deep, and durable knockdown of transthyretin (TTR) with low interpatient variability • TRITON-CM will test the hypothesis that nucresiran can reduce all-cause mortality and recurrent CV events in patients with ATTR-CM

Exploratory and Pharmacokinetic/Pharmacodynamic (PK/PD) Endpoints

270

180

Visit (Days)

12. Lachmann et al. N Engl J Med 2007;356:2361–7; 13. Murad A, et al. American Heart Association, 16–18 November 2024. 13. Gillmore et al. Eur Heart J 2018;39:2799–806.

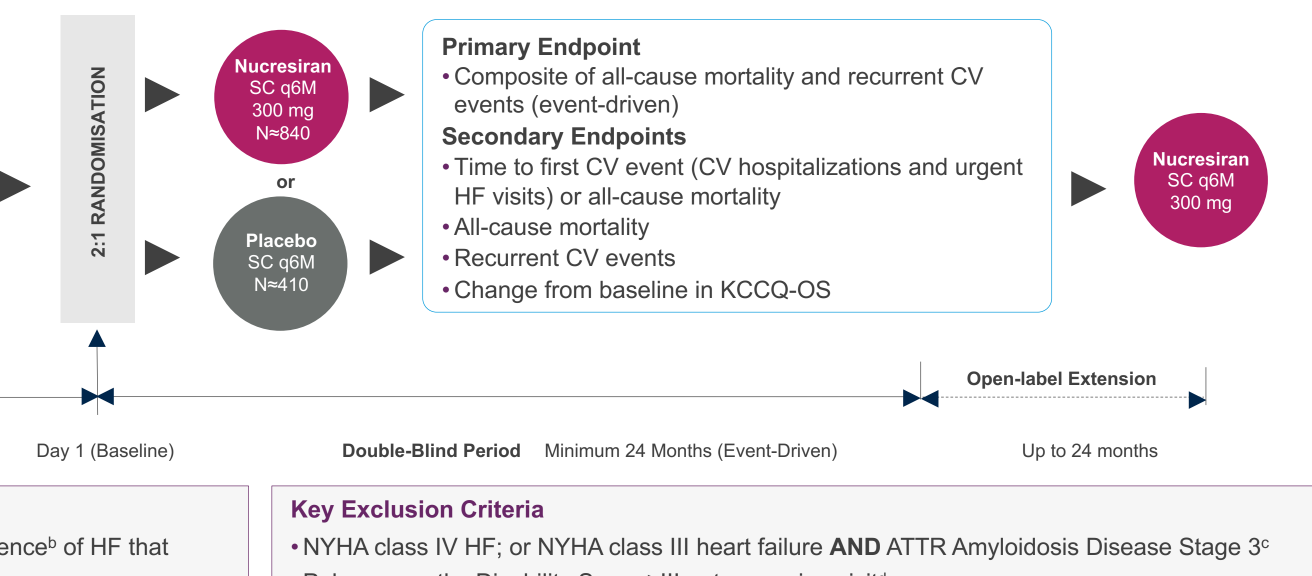
315

360

Figure 4. Nucresiran TRITON-CM Phase 3 Study Design

Patient Population: N≈1250

- Adults (18–85 years) with wild-type or variant ATTR-CM •NYHA class I–III
- End-diastolic interventricular septal wall thickness >12 mm
- for males and >11 mm for females by echocardiography
- NT-proBNP >300 and <8500 ng/L at screening^a
- Background TTR stabilizer permitted



Screening Period

Other Key Inclusion Criteria

- Medical history of HF with ≥1 prior hospitalization for HF **OR** clinical evidence^b of HF that requires treatment with a diuretic
- Demonstration of ATTR cardiomyopathy by one of the following:
- Documented TTR amyloid deposits in cardiac tissue regardless of MGUS status
- In the absence of MGUS, ^{99m}Tc scintigraphy with Grade 2 or 3 cardiac uptake
- In the presence of MGUS, documentation of TTR protein in noncardiac tissue **AND** Grade 2 or 3 cardiac uptake on ^{99m}Tc scintigraphy

^aNT-proBNP >600 pg/mL and <8500 pg/mL for patients with atrial fibrillation. ^bManifested by signs and symptoms of volume overload or elevated intracardiac pressures. ^cDefined as NT-proBNP >3000 ng/L and eGFR <45 mL/min.¹³ ^dRequires cane or stick to walk or is wheelchair-bound due to polyneuropathy.

• Patients will be randomized 2:1 to receive nucresiran 300 mg SC or placebo SC q6M for at least 24 months (**Figure 4**)

• After a minimum of 24 months and the double-blind period has ended OR after completing 5 years in the double-blind period, patients initially randomized to placebo will switch to nucresiran and all patients will receive open-label treatment with nucresiran for up to 24 months during the extension period

• Exploratory endpoints include the change from baseline in: cardiac biomarkers (NT-proBNP, troponin I), echocardiographic parameters, NAC stage, NYHA class, EuroQoL-5 Dimensions, neurofilament light chain, and oral diuretic intensification/initiation

• PK/PD endpoints include change from baseline in: serum TTR, plasma PK exposure, and the frequency and titers of anti-drug antibodies

Statistical Analysis

- The analysis timing is event-driven, requiring a prespecified number of primary composite endpoint events
- The composite endpoint of all-cause mortality and recurrent CV events will be analyzed by a modified Anderson–Gill model
- using a log-rank test; hazard ratio will be estimated using a Cox proportional hazards model rate ratio will be generated
- Time to first CV event or all-cause mortality, and time to death from any cause will be analyzed • Recurrent CV events will be analyzed using a negative binomial regression model; relative
- Change from baseline in KCCQ-OS will be analyzed using a mixed-effects model of repeated measures approach

Study Status and Timeline

- The study design, including inclusion and exclusion criteria, have been finalized • Enrolment of adult patients with ATTR-CM is expected to begin in 2025

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Abbreviations: ASGP, asialoglycoprotein; ATTR, transthyretin amyloidosis; ATTR, transthyretin anyloidosis; ATTR, hereditary ATTR; HCP, healthcare professional; HF, heart failure; KCCQ-OS, Kansas City Cardiovascular; eGFR, estimated glomerular filtration rate; GalNAc, N-Acetylgalactosamine; hATTR, hereditary ATTR; HCP, healthcare professional; HF, heart failure; KCCQ-OS, Kansas City Cardiovascular; eGFR, estimated glomerular filtration rate; GalNAc, N-Acetylgalactosamine; hATTR, hereditary ATTR; HCP, healthcare professional; HF, heart failure; KCCQ-OS, Kansas City Cardiovascular; eGFR, estimated glomerular filtration rate; GalNAc, NAC, National Amyloidosis Centre; NT-proBNP, N-terminal prohormone of B-type natrivetic peptide; NYHA, New York Heart Association; PD, pharmacodynamics; PK, pharmacokinetics; q6M, every 6 months; RISC, RNA-induced silencing complex; RNAi, small interference; SC, subcutaneous; SEM, standard error of the mean; siRNA, small interfering RNA; Tc, technetium; TTR, transthyretin; wt, wild-type; wtATTR, wild type-ATTR.



- Polyneuropathy Disability Score ≥IIIa at screening visit^d
- Estimated glomerular filtration rate (eGFR) <30 mL/min/1.73 m²
- Non-TTR cardiomyopathy, hypertensive cardiomyopathy, cardiomyopathy due to valvular heart disease, or cardiomyopathy due to ischemic heart disease
- Received prior TTR-lowering therapy **OR** plan for or anticipate beginning treatment during screening or the first 24 months following randomization