# Vutrisiran Reduces Days Lost to Death and/or Hospitalization Versus Placebo in Patients with Transthyretin Amyloidosis with Cardiomyopathy in the HELIOS-B Trial

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# **Conclusions**

- In this post hoc analysis from HELIOS-B, vutrisiran reduced the number of days lost to death and/or hospitalization (all-cause or CV-related) versus placebo.
- Patients receiving vutrisiran were more likely (64% increased odds) to be completely free from days lost to death or hospitalization compared with patients receiving placebo.
- When days lost to impaired well-being (according to KCCQ and NYHA class) were additionally accounted for, the beneficial effects of vutrisiran were even greater.



In patients with ATTR-CM, vutrisiran decreased the number of days lost to death and/or hospitalization by more than 1 month versus placebo over a 3-year period.

## Introduction

#### **Transthyretin Amyloidosis with Cardiomyopathy**

- ATTR-CM is a progressive, debilitating, and ultimately fatal disease caused by the accumulation of misfolded TTR protein (variant or wild-type) as amyloid fibrils in the heart. 1–3

  HELIOS-B Study
- Vutrisiran, an RNAi therapeutic that reduces the production of variant and wild-type TTR, was evaluated in patients with ATTR-CM in the Phase 3 HELIOS-B study (NCT04153149).
- In HELIOS-B, vutrisiran significantly reduced the risk of the composite primary endpoint of all-cause mortality and recurrent CV events versus placebo and met all secondary endpoints in the overall population and the monotherapy population (those not receiving tafamidis, another approved therapy for ATTR-CM, at baseline).4

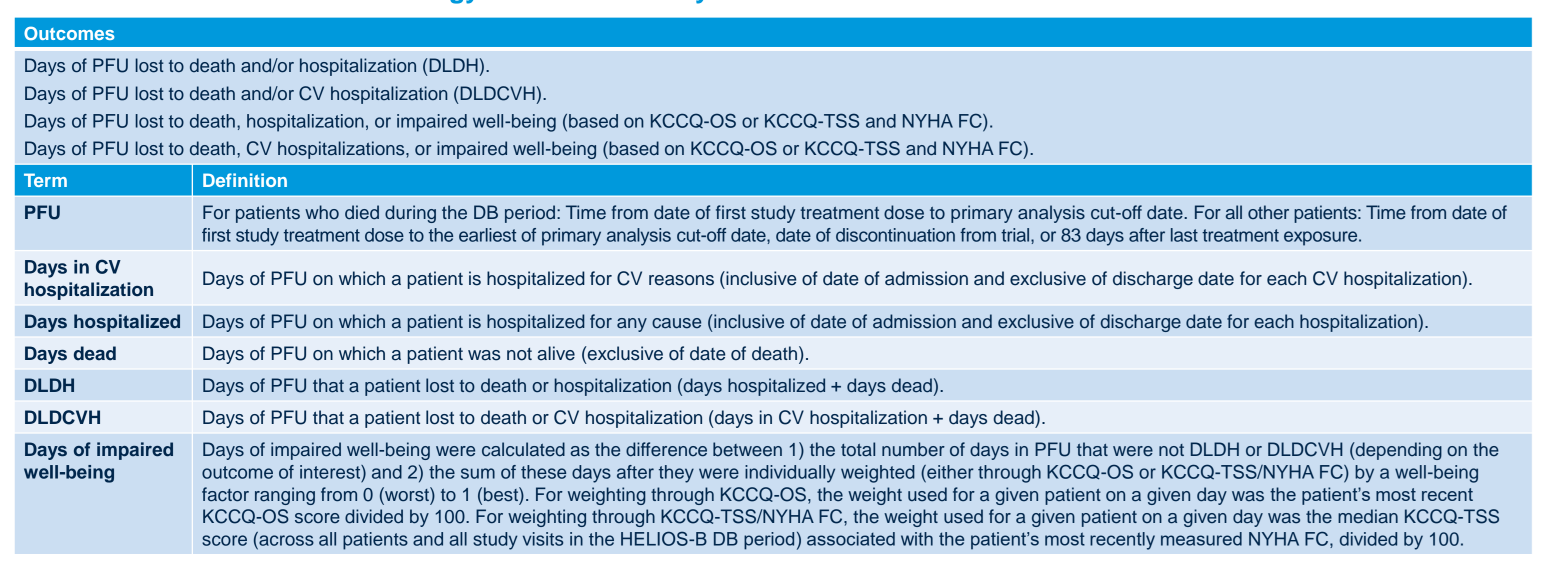
#### **Objective**

• To further characterize the benefits of vutrisiran on all-cause mortality and recurrent CV events, in a way that reflects the practical impacts of how early or late deaths occurred during follow-up and/or how long CV events lasted, this post hoc analysis from HELIOS-B assessed the effect of vutrisiran versus placebo on days lost to death and/or hospitalization in the overall and monotherapy populations.

### Methods

- Patients with ATTR-CM were randomized 1:1 in the Phase 3 HELIOS-B study to receive sc vutrisiran 25 mg or placebo every 3 months for up to 36 months in the DB period.<sup>4</sup>
- Outcomes reflecting days lost to death, hospitalization, and/or impaired well-being were assessed based on mortality and hospitalizations continuously ascertained over the DB period (Table 1).

#### **Table 1. Outcomes and Terminology Used in the Analysis**



#### **Statistical Analyses**

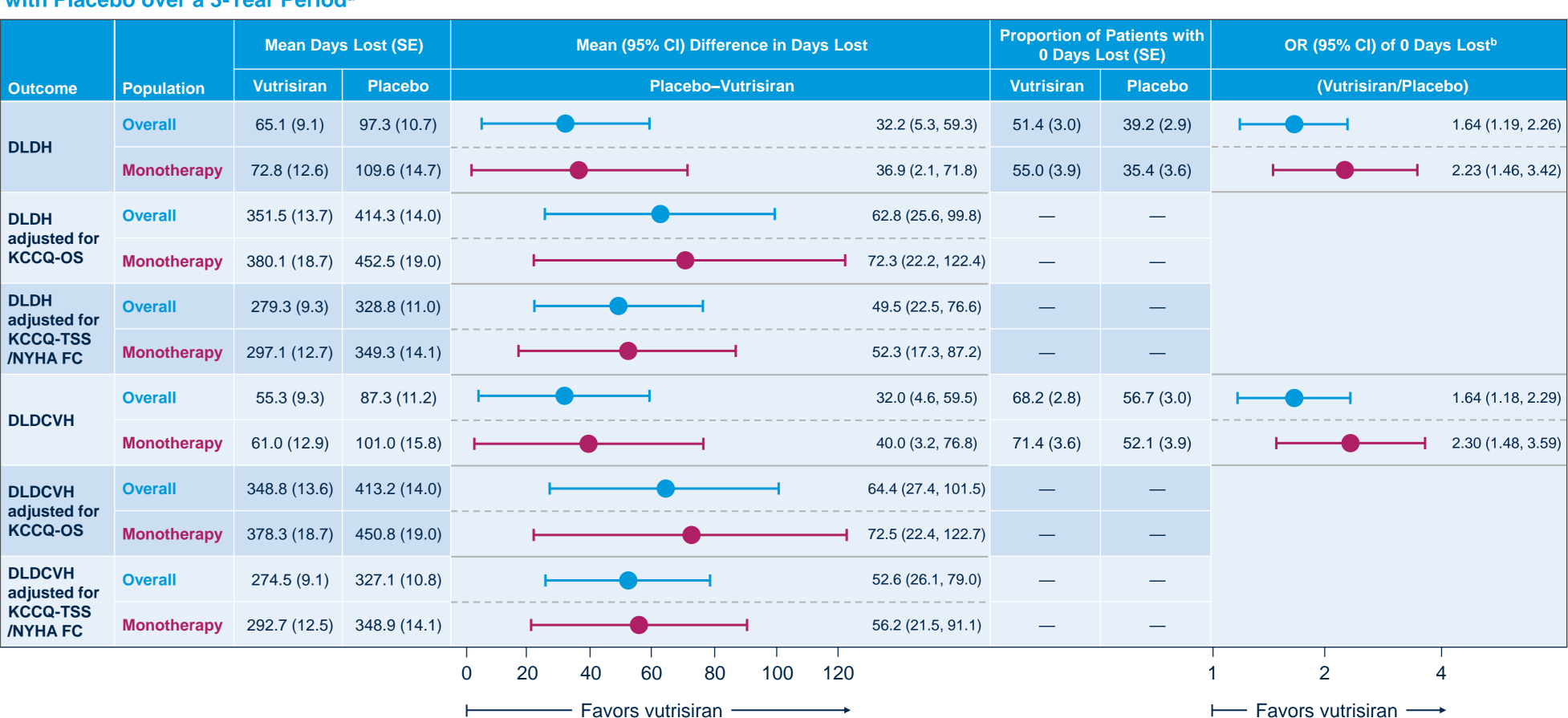
- Outcomes were analyzed as a proportion out of the PFU.
   Outcomes with ≥10 patients having a value of 0 days lost were fitted to a zero-inflated beta model.
  - Otherwise, the outcome was fitted to a beta model with patients having 0 days lost adjusted to have a value of 1 / PFU.
- Each of the location and shape parameters (and mixture parameter when fitted to the zero-inflated beta model) included treatment, log-transformed PFU, log-transformed NT-proBNP, age group, baseline tafamidis use, and treatment by baseline tafamidis use interaction as covariates.
  - In analyzing the vutrisiran monotherapy population, the models excluded the baseline tafamidis terms.
    Results are derived from a model-based estimation for a patient with 3 years of PFU.

#### .1–3

Results

- The analysis included data from 654 patients (vutrisiran, n=326; placebo, n=328), with 395 patients in the monotherapy subgroup (vutrisiran, n=196; placebo, n=199).
- Baseline demographic and clinical characteristics were generally well balanced between the vutrisiran and placebo groups, except that NT-proBNP and troponin I levels were higher (indicating greater disease severity) in the vutrisiran group than in the placebo group within the monotherapy population.

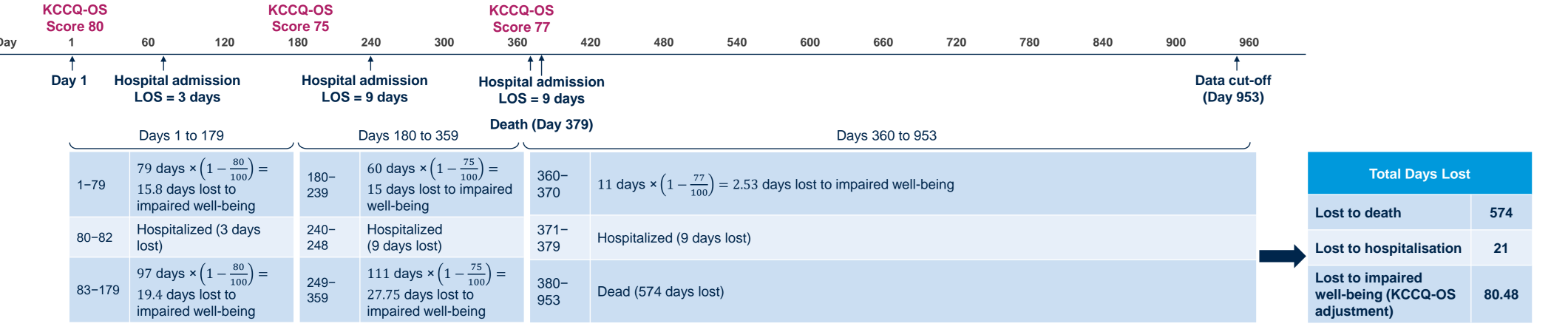
# Figure 2. Vutrisiran Reduced the Mean Days Lost to Death and/or Hospitalization and Mean Days Lost to Death and/or CV Hospitalization Compared with Placebo over a 3-Year Period<sup>a</sup>



<sup>a</sup>Data derived from a model-based estimation for a patient with 3 years of PFU. <sup>b</sup>When adjusting for impaired well-being, no patients had 0 days lost. Therefore, there is no OR listed for outcomes that involve adjustment for impaired well-being.

- In the overall population, vutrisiran treatment resulted in a mean of 32.2 (95% CI 5.3, 59.3) fewer DLDH and 32.0 (95% CI 4.6, 59.5) fewer DLDCVH over 3 years versus placebo (Figure 2).
  - Patients treated with vutrisiran had 64% greater odds of being completely free from DLDH or DLDCVH during HELIOS-B follow-up when compared with patients receiving placebo.
- In analyses that assessed days of impaired well-being (for days alive and not hospitalized) as a contributor to days lost, a larger effect of vutrisiran in preventing days lost was observed.
  - In the overall population, the treatment effect of vutrisiran translated to a mean of 62.8 (95% CI 25.6, 99.8) fewer DLDH versus placebo over a 3-year period when weighted by KCCQ-OS, and a mean of 49.5 (95% CI 22.5, 76.6) fewer DLDH versus placebo when weighted by KCCQ-TSS and NYHA FC collectively (**Figure 2**).
- Similarly, in the overall population, the treatment effect of vutrisiran translated to a mean of 64.4 (95% CI 27.4, 101.5) fewer DLDCVH versus placebo over a 3-year period when weighted by KCCQ-OS, and a mean of 52.6 (95% CI 26.1, 79.0) fewer DLDCVH versus placebo when weighted by KCCQ-TSS and NYHA FC collectively (Figure 2).
- Results observed in the monotherapy population were generally similar to those in the overall population (Figure 2).

# Figure 1: Example Calculation of Days Lost to Death, Hospitalization, and Days of Impaired Well-Being from a Hypothetical Patient



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