Time to End of Progression of Hereditary Angioedema Attacks Treated with Sebetralstat

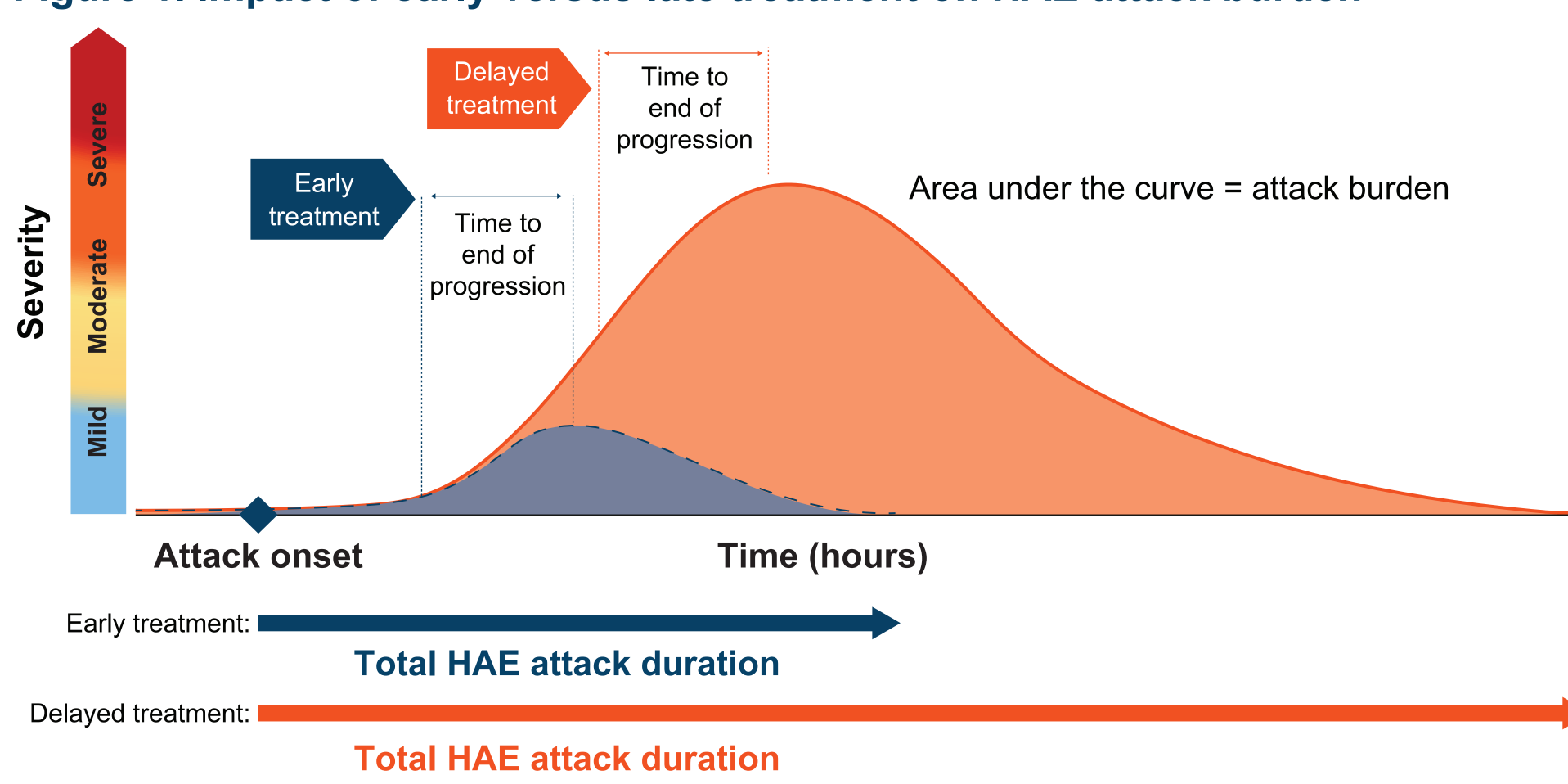
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Background

- Global treatment guidelines recommend that people living with hereditary angioedema (HAE) treat attacks with on-demand medication as early as possible after recognizing the onset^{1,2}
- Early treatment is indicated to arrest progression of swelling, thereby reducing severity and morbidity, limit potential spread of the HAE attack from the initial site, shorten total attack duration, and reduce disruption in daily activities^{1,2} (Figure 1)
- Currently, all approved on-demand therapies require injections and are associated with delays, which may lead to progression and therefore increased symptom burden³⁻⁹
- Sebetralstat is an oral plasma kallikrein inhibitor being evaluated for on-demand treatment of HAE attacks¹⁰
- In pharmacologic studies, sebetralstat achieved rapid plasma exposure and near-complete inhibition of plasma kallikrein within 15 minutes in patients with HAE⁹

Figure 1. Impact of early versus late treatment on HAE attack burden



Objective

 This post hoc analysis examined the time to end of HAE attack progression following treatment with sebetralstat in the phase 3 KONFIDENT trial and KONFIDENT-S open-label extension (OLE) study

Methods

Study Design

HAE, hereditary angioedema.

- KONFIDENT (NCT05259917) was a phase 3, double-blind, randomized, placebo-controlled, 3-way crossover trial¹⁰
- Adults and adolescents (≥12 years old) with HAE due to C1-inhibitor deficiency (HAE-C1INH) and ≥2 documented HAE attacks within 3 months were randomly assigned to 1 of 6 treatment sequences in which 3 eligible HAE attacks were treated with 1 to 2 doses of sebetralstat 300 mg, sebetralstat 600 mg, or placebo
- KONFIDENT-S is an ongoing multicenter OLE study (NCT05505916)
- Adults and adolescents (≥12 years old) with HAE-C1INH and ≥2 documented HAE attacks within 3 months or who completed the phase 3 KONFIDENT trial self-administered sebetralstat 600 mg (2 × 300-mg tablets) for each HAE attack as early as possible after onset
- In both KONFIDENT and KONFIDENT-S, Patient Global Impression of Severity (PGI-S) ratings from "None" to "Very Severe" were recorded at the time of treatment, every 0.5 hours during the first 4 hours after taking the trial agent, and every hour from 5 to 12 hours

Effectiveness Endpoint

- End of progression was analyzed post hoc as the time at which the worst HAE attack severity was recorded within 4 hours on the PGI-S scale for attacks treated with sebetralstat
- HAE attacks treated with conventional therapy were censored at the time of use
- HAE attacks with no post baseline assessments were excluded (except for attacks treated with conventional therapy)

Participants and HAE Attacks

Overall, data from 110 participants from the KONFIDENT trial and 134 participants from the KONFIDENT-S study (**Table 1**) who treated 172 and 1591 HAE attacks with sebetralstat (Table 2), respectively, were analyzed post hoc

Table 1. Demographics and disease characteristics

	KONFIDENT ¹⁰ N=110	KONFIDENT-S N=134
Age, mean (range), years	39.5 (25.0 to 49.0)	35.0 (21.0 to 49.0)
Sex, female, n (%)	66 (60.0)	87 (64.9)
Race, n (%)		
White	92 (83.6)	99 (73.9)
Asian	10 (9.1)	17 (12.7)
Black or African American	1 (0.9)	0
Other or not reported ^a	7 (6.4)	18 (13.4)
BMI, mean (range), kg/m ²	26.2 (22.8 to 31.7)	25.4 (22.3 to 30.6)
HAE-C1INH-Type1, n (%)	101 (91.8)	125 (93.3)
Treatment regimen, LTP + on-demand, n (%)	24 (21.8)	35 (26.1)

BMI, body-mass index; HAE-C1INH, hereditary angioedema due to C1 inhibitor deficiency; LTP, long-term prophylaxis; N/n, number of participants ^aIncludes 1 participant in KONFIDENT-S who identified as "Multiple." Data cutoff date was January 31, 2024 for KONFIDENT and September 14, 2024 for KONFIDENT-S.

Table 2. Sebetralstat-treated HAE attack characteristics

	KONFIDENT Sebetralstat 300 mg n=84 ^a	KONFIDENT Sebetralstat 600 mg n=88 ^a	KONFIDENT-S Sebetralstat 600 mg n=1591a
HAE attack location, n (%) ^b			
Subcutaneous	48 (57.1)	49 (55.7)	930 (58.5)
Mucosal	35 (41.7)	39 (44.3)	650 (40.9)
Abdominal	32 (38.1)	35 (39.8)	603 (37.9)
Laryngeal	1 (1.2)	1 (1.1)	30 (1.9)
Baseline PGI-S rating, n (%) ^{c,d}			
Mild	36 (42.9)	40 (45.5)	594 (37.3)
Moderate	33 (39.3)	30 (34.1)	668 (42.0)
Severe	12 (14.3)	16 (18.2)	262 (16.5)
Very severe	2 (2.4)	2 (2.3)	58 (3.6)
Time to treatment, median (IQR), min	36.0 (7.0-131.0)	38.5 (4.5-141.0)	14.5 (1.0-76.0)

IQR, interquartile range; n, number of attacks; PGI-S, Patient Global Impression of Severity. ^aNumber of HAE attacks with a baseline PGI-S score KONFIDENT: sebetralstat 300 mg n=83; sebetralstat 600 mg n=88. KONFIDENT-S: sebetralstat 600 mg ^bBaseline attack location was missing for 1 attack (1.2%) treated with sebetralstat 300 mg in KONFIDENT and 11 attacks (0.7%) treated with

°PGI-S rating of "None" included in "Mild": KONFIDENT sebetralstat 300 mg, 0; KONFIDENT sebetralstat 600 mg, 0; KONFIDENT-S sebetralstat 600 mg, dPGI-S missing: KONFIDENT sebetralstat 300 mg, 1 (1.2%); KONFIDENT sebetralstat 600 mg, 0; KONFIDENT-S sebetralstat 600 mg, 9 (0.6%). Data cutoff date was January 31, 2024 for KONFIDENT and September 14, 2024 for KONFIDENT-S.

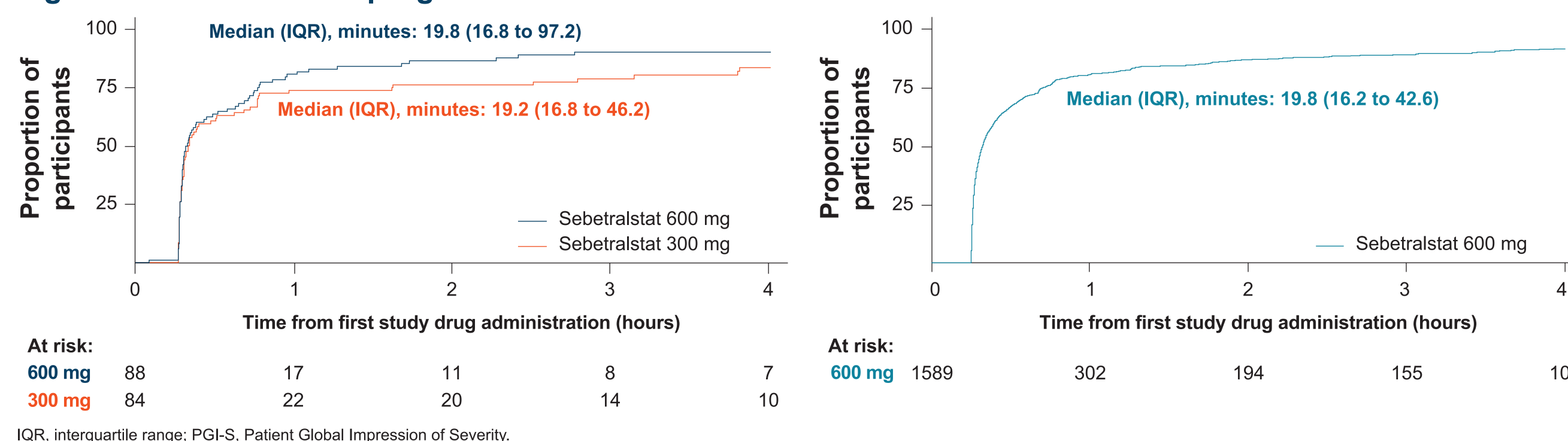
Effectiveness

sebetralstat 600 mg in KONFIDENT-S.

- The median time to end of progression was 19.2 minutes (IQR, 16.8 to 46.2) with sebetralstat 300 mg and 19.8 minutes (IQR, 16.8 to 97.2) with sebetralstat 600 mg in the KONFIDENT trial; in the KONFIDENT-S study, the median time was 19.8 minutes (IQR, 16.2 to 42.6) with sebetralstat 600 mg (Figure 2)
- In total, 69 (82.1%) HAE attacks treated with sebetralstat 300 mg, 79 (89.8%) treated with sebetralstat 600 mg in KONFIDENT, and 1437 (90.3%) treated with sebetralstat 600 mg in KONFIDENT-S reached end of progression within 4 hours

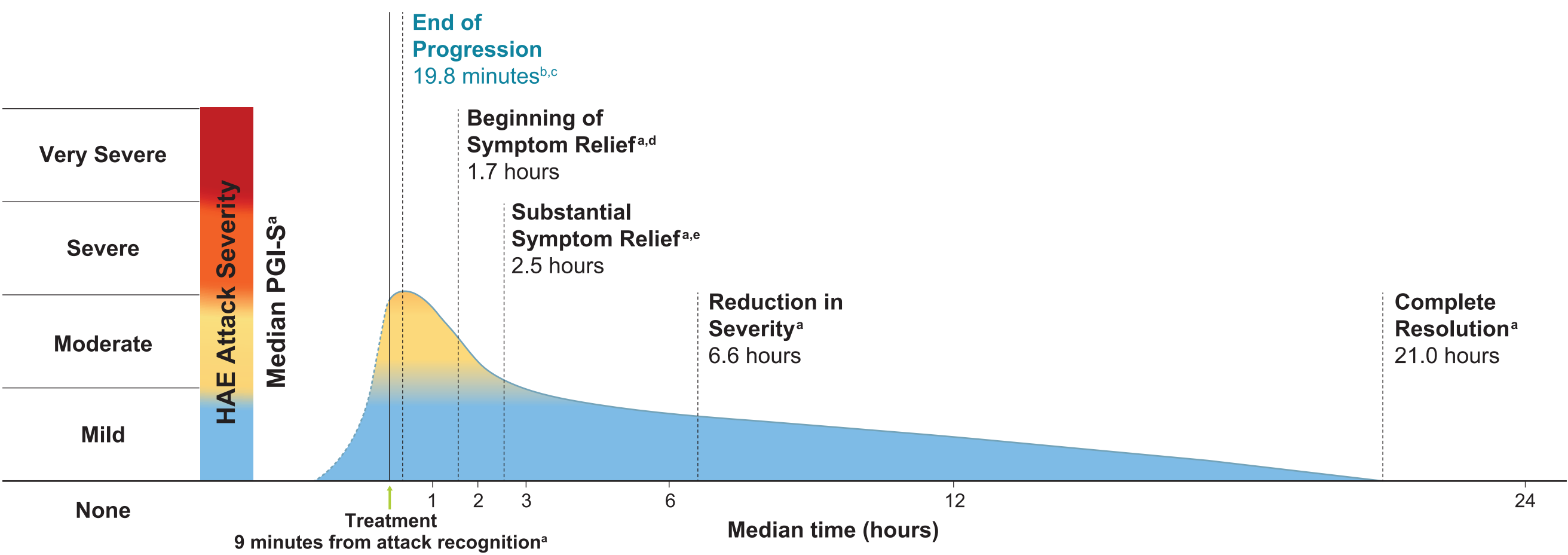
Results

Figure 2. Time to end of progression



Time to end of progression was defined as the time at which the worst HAE attack severity was recorded using the 5-point PGI-S scale within 4 hours. Data cutoff date was January 31, 2024 for KONFIDENT and September 14, 2024 for KONFIDENT-S.

Figure 3. Time course of sebetralstat-treated HAE attacks (illustrative)



HAE, hereditary angioedema; OLE, open-label extension; PGI-C, Patient Global Impression of Change; PGI-S, Patient Global Impression of Severity. ^aBased on data cutoff date of January 31, 2024 (640 attacks treated with sebetralstat 600 mg). ¹²

Graph generated based on median time to endpoints as observed in the KONFIDENT-S OLE study. Area under the curve represents the treatment burden.

^bBased on data cutoff date of September 14, 2024 (1706 attacks). ^cTime to end of progression was defined as the time at which the worst HAE attack severity was recorded on the PGI-S scale within 4 hours. ^dDefined as a PGI-C rating of at least "A Little Better" for at least 2 consecutive time points (with missing data entries in between) within 12 hours. ^eDefined as time to substantial reduction of symptom burden (PGI-S of "Mild" for 2 consecutive time points) within 24 hours.

Conclusions

- Results of this post hoc analysis suggested that sebetralstat ended the progression of the HAE attack early, with a median of 19.8 minutes in KONFIDENT-S
- These results are consistent with those observed with sebetralstat 300 mg and sebetralstat 600 mg in the phase 3 KONFIDENT trial
- The time to end of HAE attack progression closely followed the expected time to near-complete inhibition of plasma kallikrein (15 minutes) based on prior pharmacodynamic studies¹¹
- Oral sebetralstat allowed for rapid treatment and halted HAE attack progression early in patients with HAE

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