

Long-Term Efficacy and Safety of Oral Deucricitibant, a Bradykinin B2 Receptor Antagonist, in Treatment of Hereditary Angioedema Attacks: Results of the RAPIDe-2 Extension Study

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Conflicts of interest disclosure

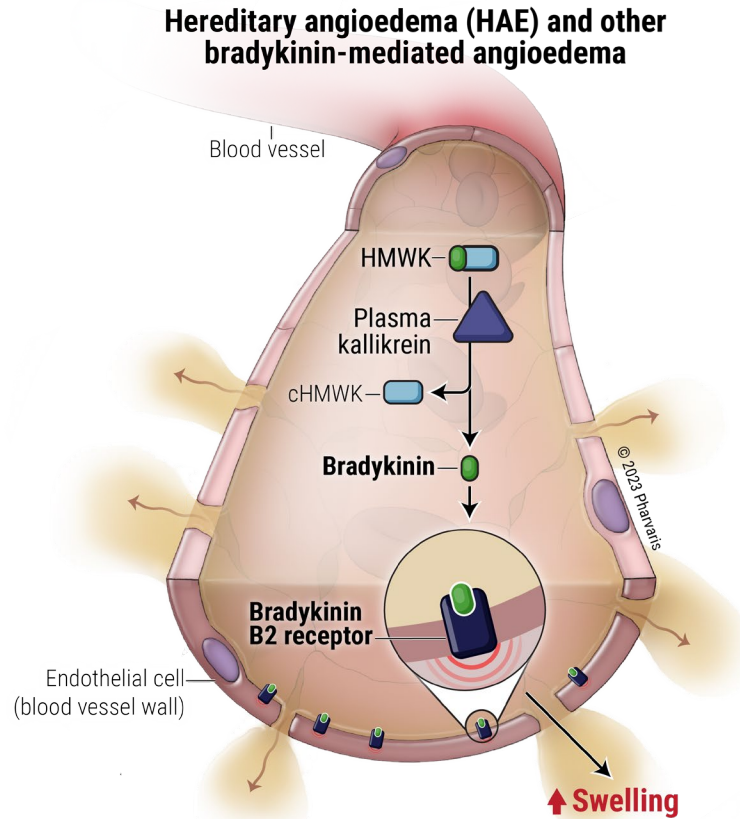
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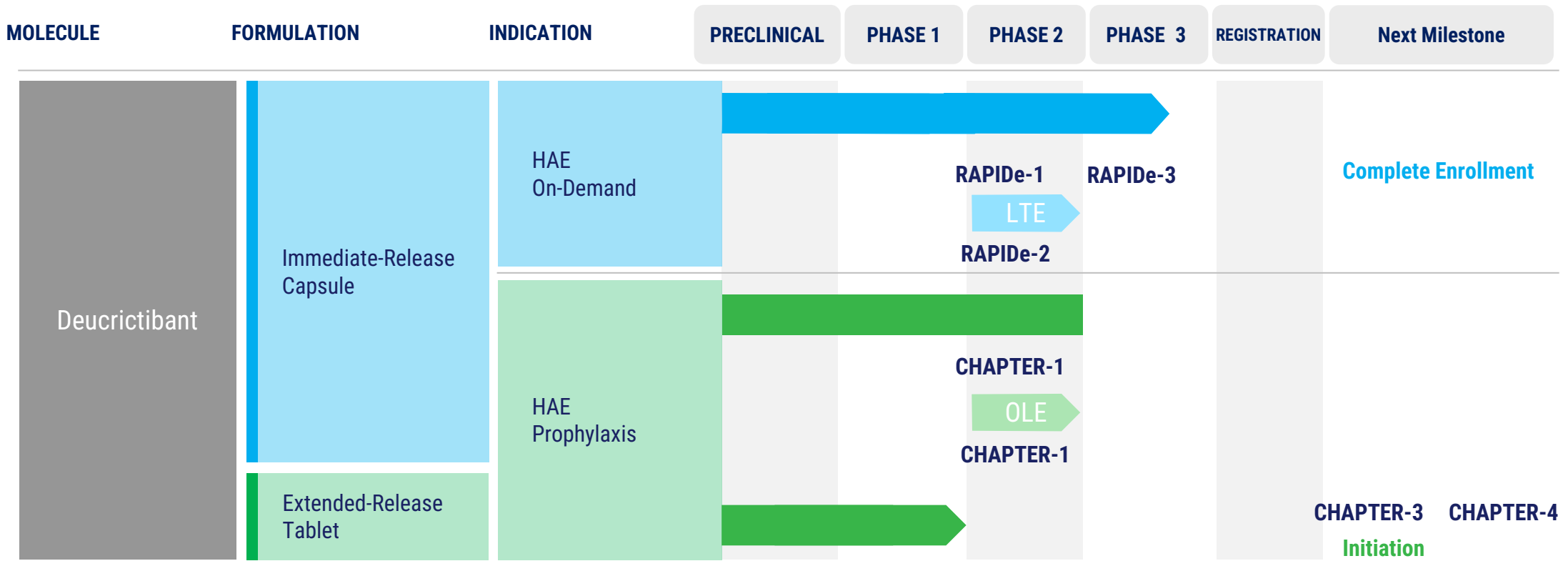
Hereditary angioedema (HAE) is a bradykinin-mediated condition with unmet medical needs



- International guidelines recommend that HAE attacks are **treated as early as possible**.¹⁻³
- Burden associated with **parenteral administration** of currently approved on-demand medications⁴⁻⁸ leads to treatment of a number of HAE attacks being **delayed or forgone**.⁹⁻¹³
- An unmet need exists for **on-demand oral** therapies that are effective and well-tolerated and may reduce the treatment burden enabling prompt administration.¹³

cHMWK, cleaved HMWK; HAE, hereditary angioedema; HMWK, high-molecular-weight kininogen. **1.** Betschel S, et al. *Allergy Asthma Clin Immunol*. 2019;15:72. **2.** Busse PJ, et al. *J Allergy Clin Immunol Pract*. 2021;9:132-50. **3.** Maurer M, et al. *Allergy*. 2022;77:1961-90. **4.** Berinert®. Package insert. Accessed September 16, 2024. <https://labeling.cslbehring.com/pi/us/berinert/en/berinert-prescribing-information.pdf>; **5.** Cinryze®. Summary of product characteristics. Accessed September 16. https://www.ema.europa.eu/en/documents/product-information/cinryze-epar-product-information_en.pdf; **6.** Firazyr®. Package insert. Accessed September 16, 2024. https://www.shirecontent.com/PI/PDFs/Firazyr_USA_ENG.pdf; **7.** Kalbitor®. Package insert. Accessed September 16, 2024. https://www.shirecontent.com/PI/PDFs/Kalbitor_USA_ENG.pdf; **8.** Ruconest®. Package insert. Accessed September 19, 2024. https://www.ruconest.com/wp-content/uploads/Ruconest_PL_Apr2020.pdf; **9.** Burnette A, et al. Presented at: AAAAI; February 24–27, 2023; San Antonio, TX, USA. **10.** Tuong LA, et al. *Allergy Asthma Proc* 2014;35:250-4. **11.** Center for Biologics Evaluation and Research. The voice of the patient—Hereditary angioedema. US Food and Drug Administration. Accessed September 19, 2024. <https://www.fda.gov/media/113509/download>; **12.** Radojicic C, et al. Presented at: AAAAI; February 24–27, 2023; San Antonio, TX, USA. **13.** Mendevil J, et al. Presented at: ACAAI; November 9–13, 2023; Anaheim, CA, USA.

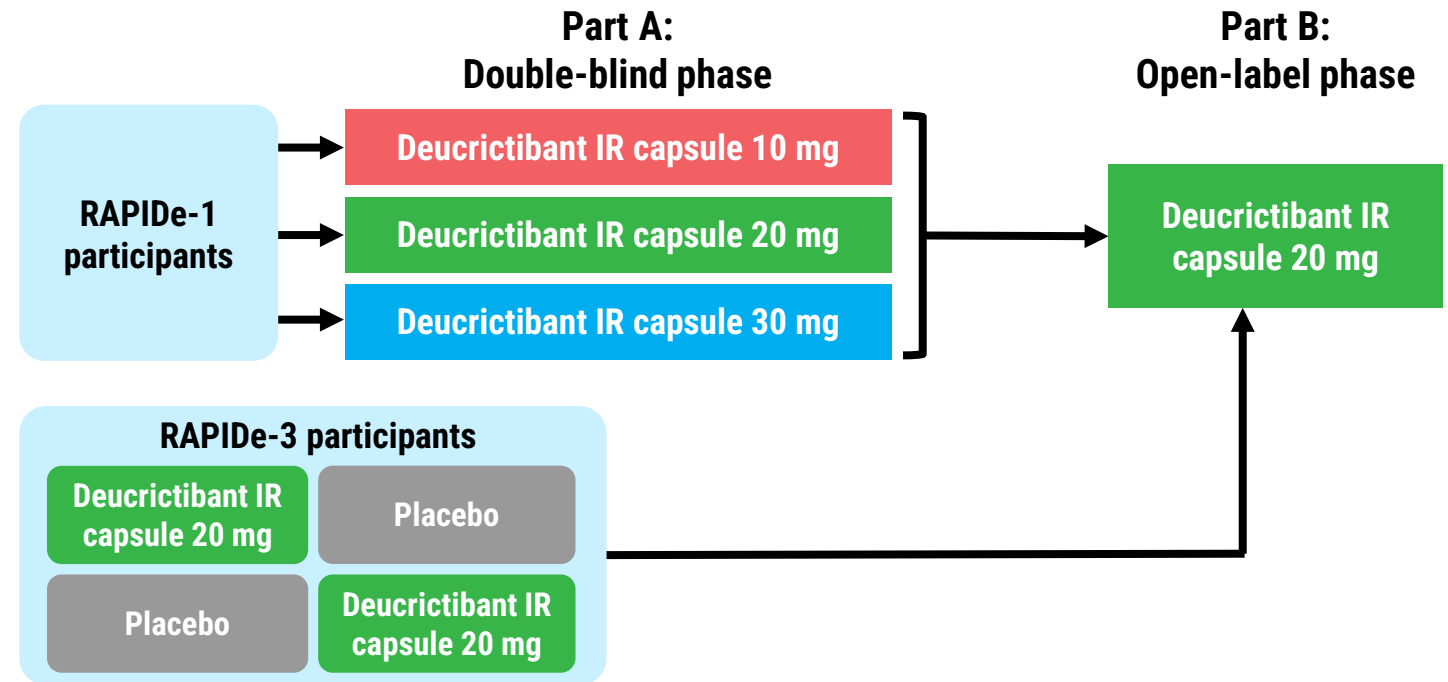
Deucricitibant development program in HAE



HAE, hereditary angioedema; LTE, long-term extension; OLE, open-label extension. **1.** RAPIDe-1. ClinicalTrials.gov identifier: NCT04618211. Accessed September 23, 2024. <https://www.clinicaltrials.gov/study/NCT04618211>. **2.** RAPIDe-2. ClinicalTrials.gov identifier: NCT05396105. Accessed September 23, 2024. <https://www.clinicaltrials.gov/study/NCT05396105>. **3.** RAPIDe-3. ClinicalTrials.gov identifier: NCT06343779. Accessed September 23, 2024. <https://www.clinicaltrials.gov/study/NCT06343779>. **4.** CHAPTER-1. ClinicalTrials.gov identifier: NCT05047185. Accessed September 23, 2024. <https://www.clinicaltrials.gov/study/NCT05047185>.

RAPIDe-2 objectives and study design

- RAPIDe-2 (NCT05396105)¹ is an ongoing, two-part, Phase 2/3 extension study evaluating long-term safety and efficacy of orally-administered deucricitbant IR capsule for the treatment of HAE attacks.
 - Part A enrolls adult (≥ 18 years) participants who completed RAPIDe-1.
 - In Part A, participants continue self-administering the same double-blinded dose of deucricitbant IR capsule (10 mg, 20 mg, or 30 mg) received in RAPIDe-1 to treat qualifying non-laryngeal attacks (≥ 1 symptom with Visual Analogue Scale score ≥ 30), and laryngeal attacks presenting without breathing difficulties.
 - This presentation shows data from the RAPIDe-2 Part A combined-dose group at the date of cutoff.



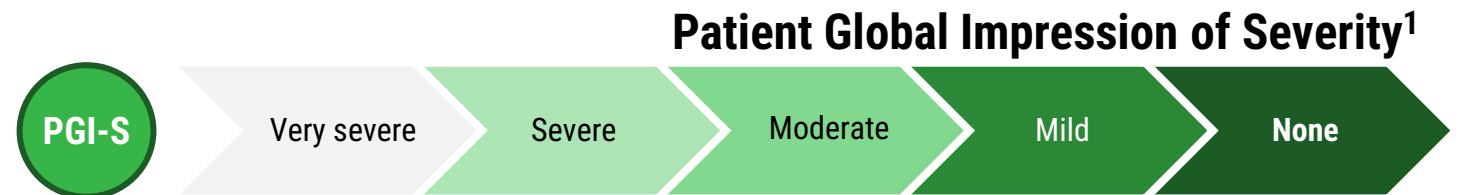
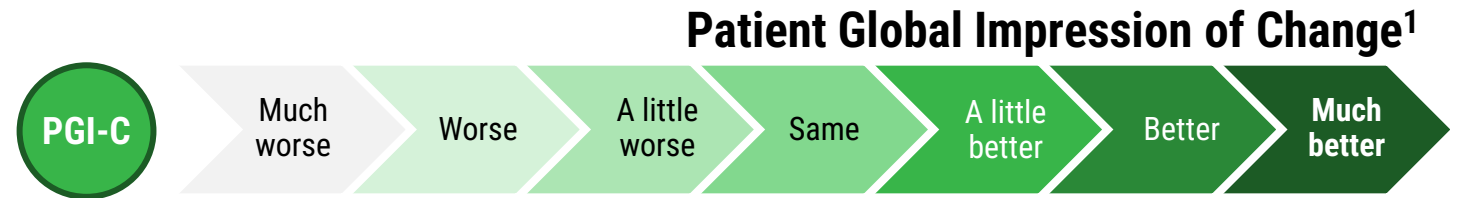
Study endpoints

- **Primary endpoint:** Safety, including TEAEs, clinical laboratory tests, vital signs, and ECG findings.
- **Efficacy:** Assessed using PRO tools.
- **Key efficacy endpoints:**

- **Time to onset of symptom relief:**
PGI-C rating of at least “a little better” for 2 consecutive timepoints by 12 hours post-treatment.

- **Time to reduction in attack severity:**
≥1 level reduction in PGI-S from pre-treatment for 2 consecutive timepoints by 12 hours post-treatment.

- **Proportion of attacks achieving complete attack resolution:**
PGI-S rating of “none” at 24 hours post-treatment.



Baseline characteristics

- 265 attacks from 17 participants included in the mITT efficacy analysis set (data cutoff: 01 March 2024).^a
- 337 attacks from 19 participants included in the safety analysis set (data cutoff: 10 June 2024).^b
 - 7 of 337 attacks were laryngeal.
- Baseline characteristics consistent with the RAPIDe-1 Phase 2 trial.

	Deucricitbant IR capsule (All doses)
Number of attacks treated^c	337
Number of participants^c	19
Age in years, mean (SD)	42.7 (17.6)
Sex: Male/female, n (%)	7 (36.8) / 12 (63.2)
Race: White/other	18 / 1
BMI, mean (SD)	27.0 (3.8)
Years since HAE diagnosis, mean (SD)	21.7 (15.2)
HAE type, n (%)	
HAE-1	17 (89.5)
HAE-2	2 (10.5)

BMI, body mass index; HAE, hereditary angioedema; IR, immediate-release; mITT, modified intention-to-treat; PGI-C, Patient Global Impression of Change; SD, standard deviation. ^aAll participants who had ≥1 attack treated with deucricitbant and non-missing PGI-C results from ≥1 post-treatment timepoint. ^bAll participants who received any dose of deucricitbant in the study. ^cNumber by the cutoff date of 10 June 2024.

Deucricitibant was well-tolerated across all doses

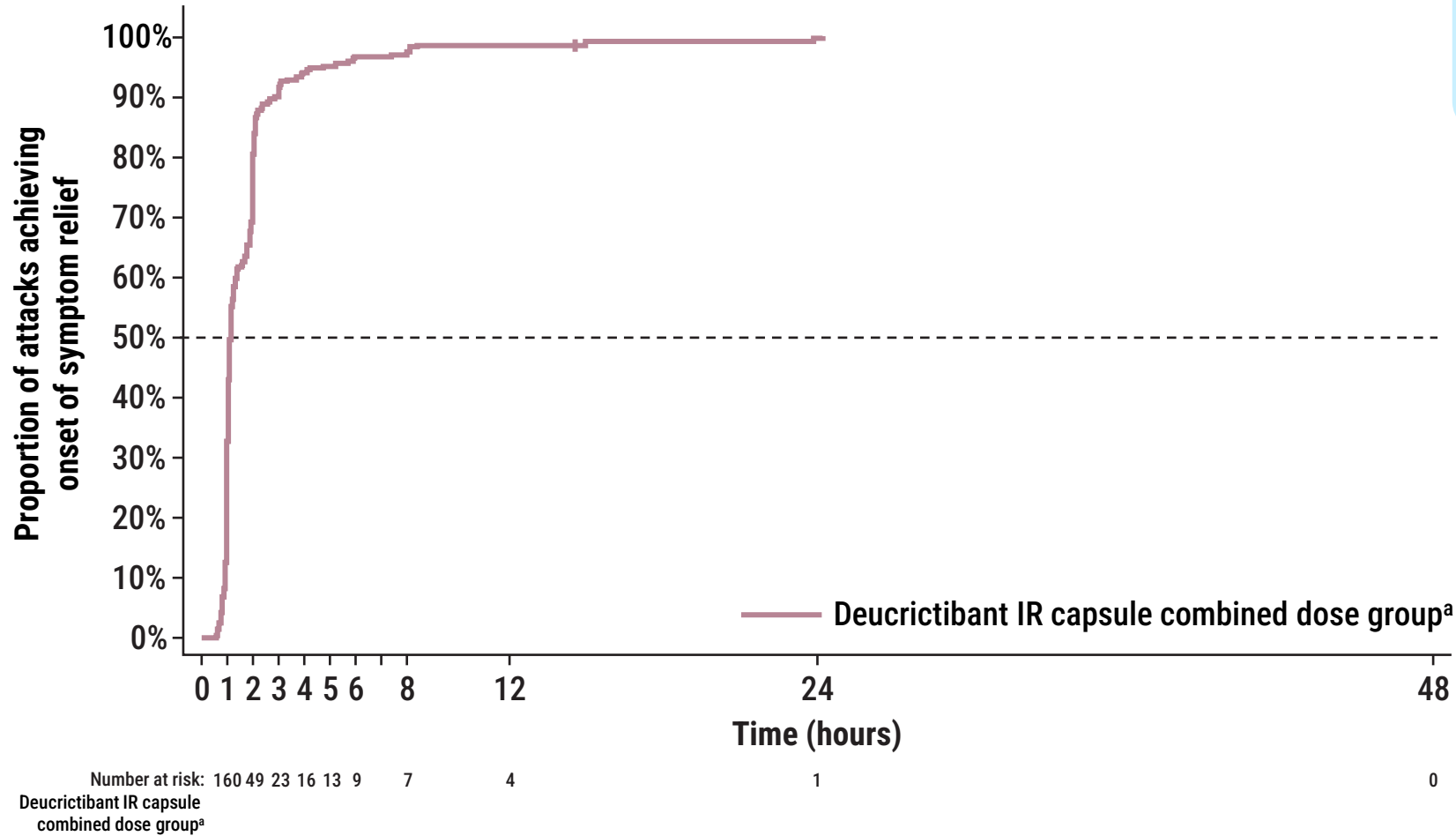
- No treatment-related TEAEs.
- No treatment-related serious or severe TEAEs, no treatment-related TEAEs in laboratory parameters, vital signs, or ECG findings.
- No TEAEs leading to treatment discontinuation, study withdrawal, or death.

TEAEs within 5 days after administration of study drug

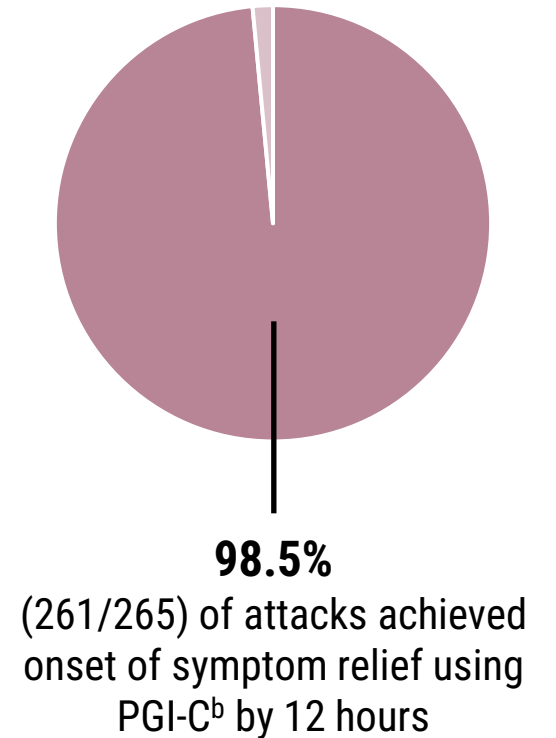
Adverse events	Deucricitibant IR capsule (All doses)
Number of attacks treated ^a	337
Number of participants ^a	19
Attacks with any TEAE, n (%)	13 (3.9)
Treatment-related TEAEs, n	0
Serious TEAEs, n	1 ^b
Treatment-related serious TEAEs, n	0
TEAEs leading to study drug discontinuation, study withdrawal, or death, n	0

ECG, electrocardiogram; IR, immediate-release; TEAE, treatment-emergent adverse event (defined as adverse event occurring during time window from first study drug administration). ^aNumber in the safety analysis set (data cutoff: 10 June 2024). ^bTooth caries unrelated to treatment.

1.1 hours median time to onset of symptom relief using PGI-C



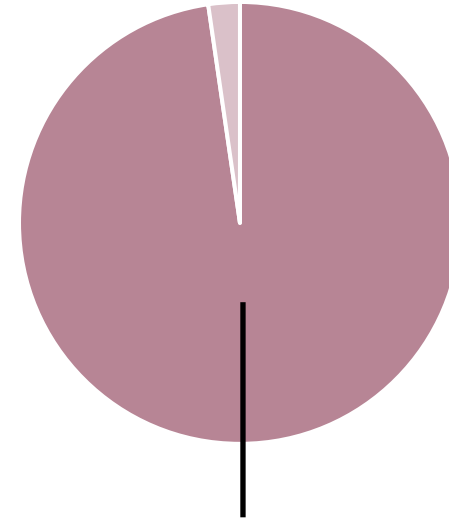
1.1 hours (95% CI, 1.0, 1.2)
median time to onset of symptom relief
using PGI-C^b



CI, confidence interval; IR, immediate-release; PGI-C, Patient Global Impression of Change. ^aIncludes 10 mg, 20 mg, and 30 mg dose groups. ^bPGI-C rating of at least "a little better" for 2 consecutive timepoints by 12 hours post-treatment.

97.7% of attacks achieved reduction in attack severity by 12 hours using PGI-S

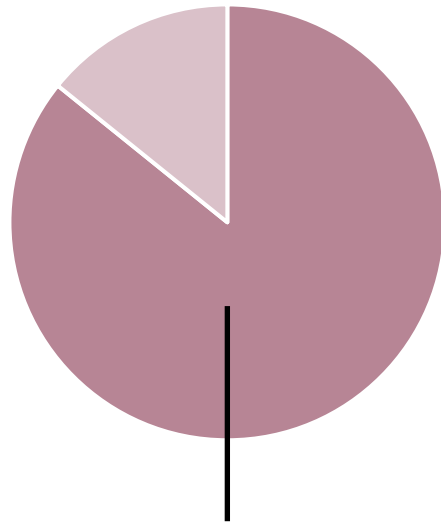
2.6 hours (95% CI, 2.0, 2.9)
median time to reduction in attack severity
using PGI-S^a



97.7%
(255/261^b) of attacks achieved reduction in
attack severity by 12 hours using PGI-S^a

85.8% of attacks achieved complete attack resolution within 24 hours with a median time of 11.5 hours

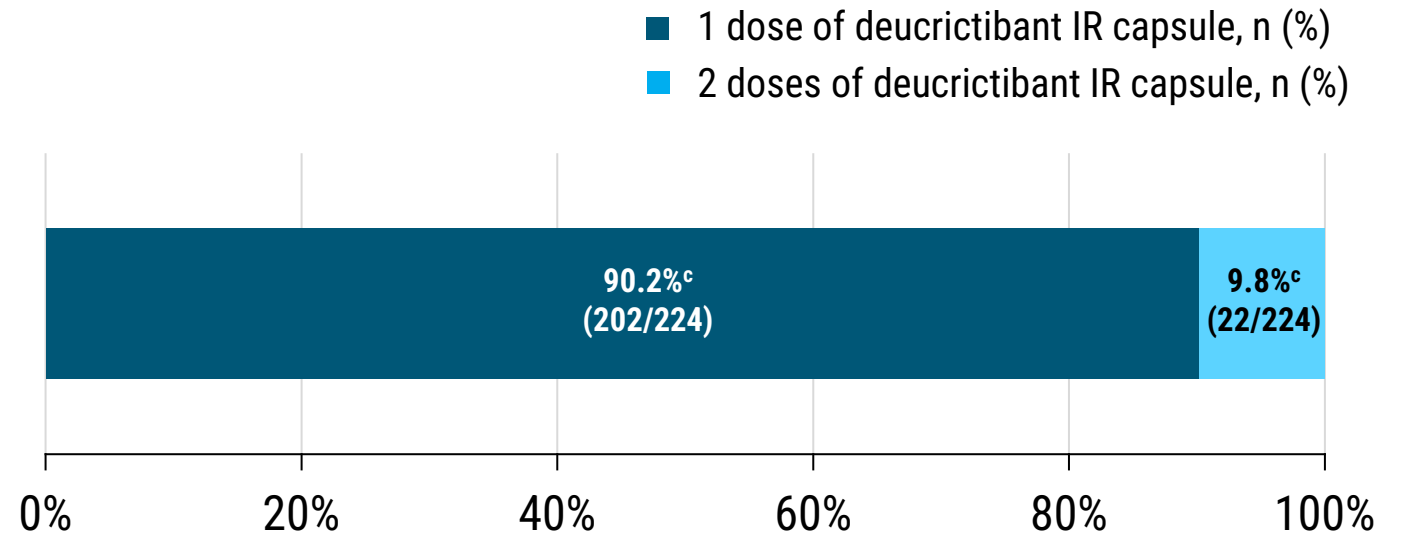
11.5 hours (95% CI, 11.0, 13.0)
median time to complete attack resolution
using PGI-S^a



85.8%
(224/261^b) of attacks achieved
complete attack resolution within 24 hours
using PGI-S^a

90.2% (202/224) of attacks achieved this milestone
with a single dose of deucricitbant IR capsule.

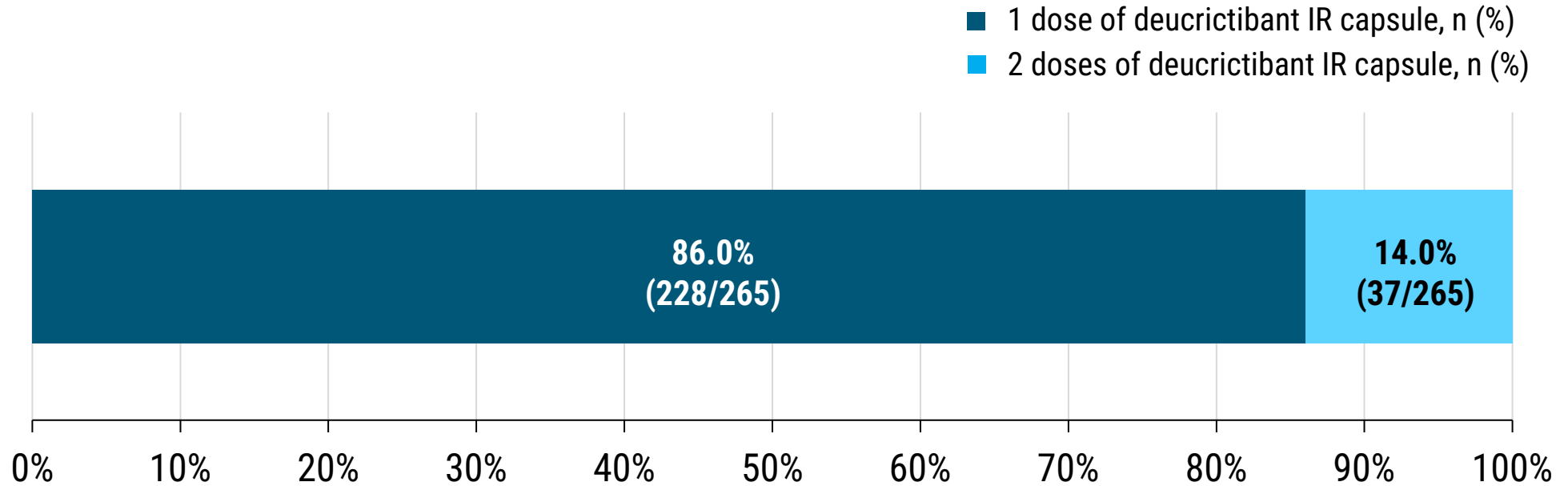
Attacks treated with 1 or 2 doses of deucricitbant prior to achieving complete attack resolution



CI, confidence interval; IR, immediate-release; PGI-S, Patient Global Impression of Severity. ^aPGI-S rating of "none" at 24 hours post-treatment. ^b261 attacks have non-missing pre-treatment PGI-S. ^cPercentage of 224 attacks achieving complete attack resolution using PGI-S within 24 hours.

Overall, 86.0% of attacks were treated with a single dose of deucricitibant

Attacks treated with 1 or 2 doses of deucricitibant



Conclusions

- In the current analysis of the ongoing RAPIDe-2 Phase 2/3 extension study, deucricitibant IR capsule was well-tolerated for all studied doses with no safety signals observed.
- Efficacy analysis showed:
 - 1.1 hours median time to onset of symptom relief using PGI-C – 98.5% of attacks by 12 hours.
 - 2.6 hours median time to reduction in attack severity using PGI-S – 97.7% of attacks by 12 hours.
 - 11.5 hours median time to complete attack resolution using PGI-S – 85.8% of attacks within 24 hours.
 - 86.0% of attacks were treated with a single dose of deucricitibant IR capsule.
- Results from the ongoing RAPIDe-2 extension are consistent with the Phase 2 RAPIDe-1 study and provide evidence on the long-term safety and efficacy of deucricitibant IR capsule for repeat treatment of HAE attacks.

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