Long-Term Efficacy and Safety of Oral Deucrictibant, 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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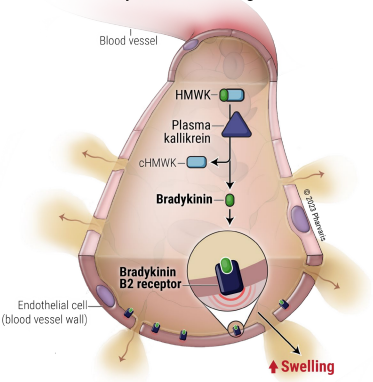
H.F.: BioCryst, CSL Behring, Intellia, KalVista, ONO Pharma, Pharming, Pharvaris, Takeda; E.A-P.: Astria, BioCryst, BioMarin, CSL Behring, Intellia, KalVista, Pharming, Pharvaris, Takeda; L.B.: BioCryst, Blueprint, CSL Behring, Novartis, Takeda; H.C.: AstraZeneca (Alexion), CSL Behring, KalVista, Merck, Novartis, Pharming, Pharvaris, Roche, Sanofi, Sobi, Takeda; D.G.: Pharming, Takeda; R.H.: BioCryst, CSL Behring, KalVista, Pharming, Pharvaris, Takeda; R.L.: BioCryst, CSL Behring, Ionis, KalVista, Novartis, Pharming, Pharvaris, Takeda, Teva; G.S.: Pharvaris, Takeda; M.Sta.: Pharming, Pharvaris, Sobi; M.Sto.: BioCryst, CSL Behring, KalVista, Pharming, Takeda; A.V.: AstraZeneca, Berlin-Chemie/Menarini Group, CSL Behring, KalVista, Novartis, Pharming, Pharvaris, Sobi, Takeda; J.S., L.Z., M.Y., G.G.: employees of Pharvaris, hold stocks in Pharvaris; P.L.: employee of Pharvaris, holds stocks/stock options in Pharvaris; M.M.: Adverum, Attune, BioCryst, CSL Behring, KalVista, Pharming, Pharvaris, Takeda.

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



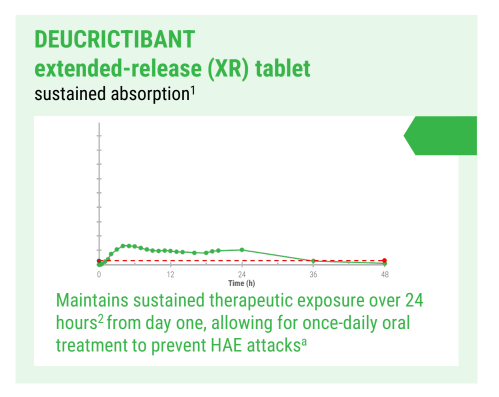


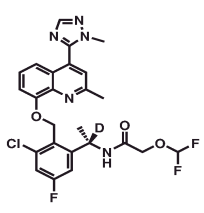
- International guidelines recommend that HAE attacks are treated as early as possible.<sup>1-3</sup>
- Burden associated with parenteral administration of currently approved on-demand medications<sup>4-8</sup> leads to treatment of a number of HAE attacks being delayed or forgone.<sup>9-13</sup>
- An unmet need exists for on-demand oral therapies that are effective and well-tolerated and may reduce the treatment burden enabling prompt administration.<sup>13</sup>

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.

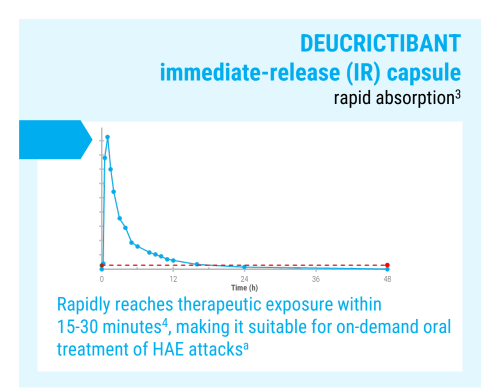
3. Maurer M. et al. *Allergy*. 2022;77:1961-90. 4. Berinert®. Package insert. Accessed September 16, 2024. https://labeling.cslbehring.com/pi/us/berinert/en/berinert-prescribing-information.pdf; 5. Cinryze®. Summary of product characteristics. Accessed September 16. https://www.ema.europa.eu/en/documents/product-information/cinryze-epar-product-information\_en.pdf; 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.ruconest.com/wp-content/uploads/Ruconest\_PI\_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.

# Two investigational oral therapies with the same active ingredient for the prophylactic and on-demand treatment of HAE attacks





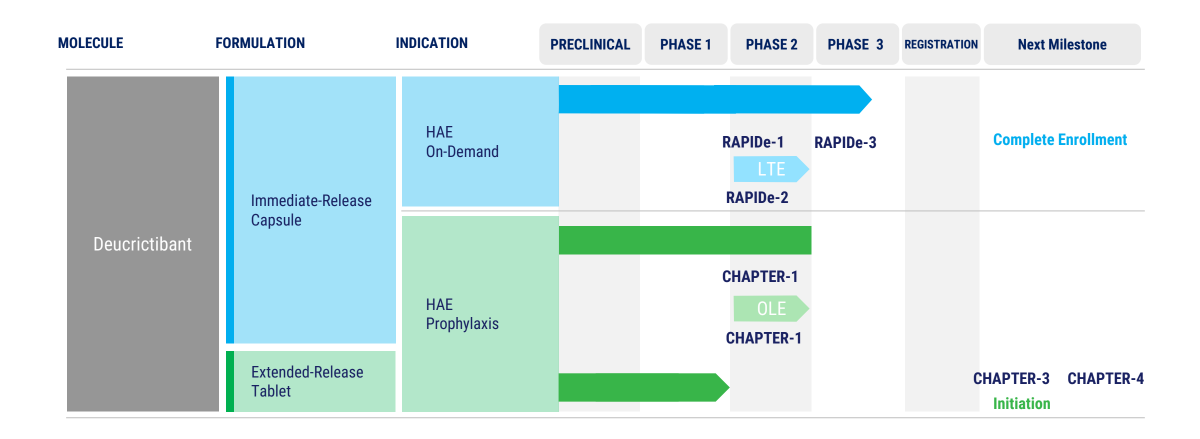
deucrictibant



### Two oral products with the same active ingredient for the prevention and treatment of HAE attacks

HAE, hereditary angioedema. <sup>a</sup>Aspirational; to be confirmed with clinical data from Phase 3 studies. **1.** Company data: single-dose cross-over PK study in healthy volunteers (n=14) under fasting conditions. **2.** Lesage A et al. Presented at IDDST; May 22-24, 2024. **3.** Crabbe et al. Presented at AAAAI: Feb 26-Mar 1, 2021. **4.** Maurer M. et al. Presented at AAAAI: Feb 24-27, 2023: San Antonio, TX, USA.

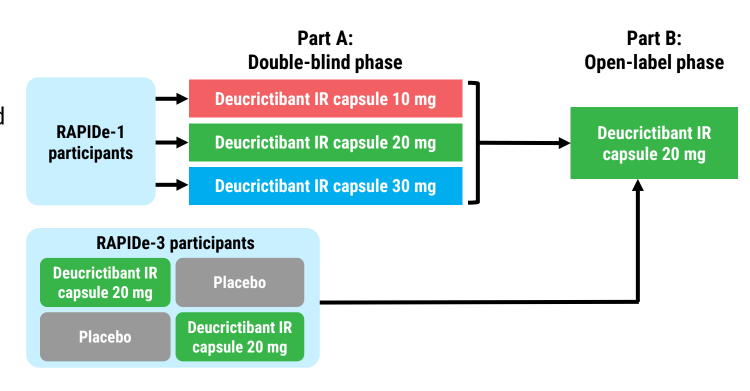
#### Deucrictibant development program in HAE



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/NCT05396105. **3**. RAPIDe-3. ClinicalTrials.gov identifier: NCT06343779. Accessed September 23, 2024. https://www.clinicaltrials.gov/study/NCT05047185. Accessed September 23, 2024. https://www.clinicaltrials.gov/study/NCT05047185.

#### RAPIDe-2 objectives and study design

- RAPIDe-2 (NCT05396105)<sup>1</sup> is an ongoing, two-part, Phase 2/3 extension study evaluating long-term safety and efficacy of orally-administered deucrictibant 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 deucrictibant 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).<sup>a</sup>
- 337 attacks from 19 participants included in the safety analysis set (data cutoff: 10 June 2024).
  - 7 of 337 attacks were laryngeal.
- Baseline characteristics consistent with the RAPIDe-1 Phase 2 trial.

	Deucrictibant IR capsule (All doses)
Number of attacks treated <sup>c</sup>	337
Number of participants <sup>c</sup>	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. <sup>a</sup>All participants who had ≥1 attack treated with deucrictibant and non-missing PGI-C results from ≥1 post-treatment timepoint. <sup>b</sup>All participants who received any dose of deucrictibant in the study. <sup>c</sup>Number by the cutoff date of 10 June 2024.

#### Deucrictibant was well-tolerated across all doses

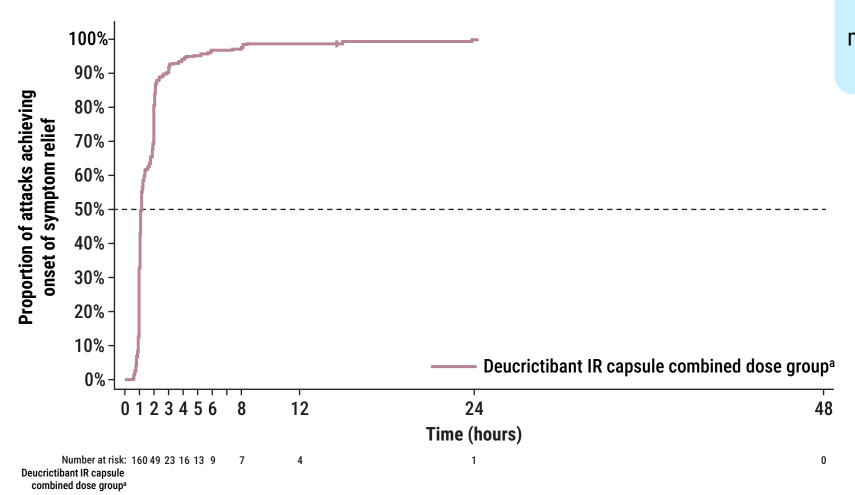
- No treatment-related TEAEs.
- No treatment-related serious or severe TEAEs, no treatment-related TEAEs in laboratory parameters, vital signs, or ECG findings.
- No TEAEs leading to treatment discontinuation, study withdrawal, or death.

#### TEAEs within 5 days after administration of study drug

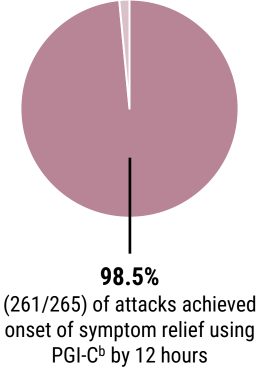
Adverse events	Deucrictibant IR capsule (All doses)
Number of attacks treated <sup>a</sup>	337
Number of participants <sup>a</sup>	19
Attacks with any TEAE, n (%)	13 (3.9)
Treatment-related TEAEs, n	0
Serious TEAEs, n	1 <sup>b</sup>
Treatment-related serious TEAEs, n	0
TEAEs leading to study drug discontinuation 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). <sup>a</sup>Number in the safety analysis set (data cutoff: 10 June 2024). <sup>b</sup>Tooth caries unrelated to treatment.

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



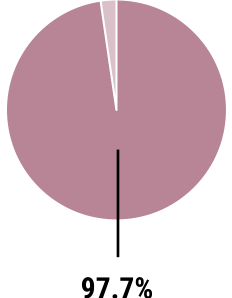
**1.1 hours** (95% CI, 1.0, 1.2) median time to onset of symptom relief using PGI-C<sup>b</sup>



Cl, confidence interval; IR, immediate-release; PGI-C, Patient Global Impression of Change. alncludes 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<sup>a</sup>

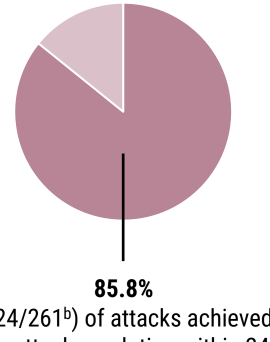


(255/261b) of attacks achieved reduction in attack severity by 12 hours using PGI-Sa

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

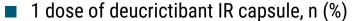
**11.5 hours** (95% CI, 11.0, 13.0) median time to complete attack resolution using PGI-Sa

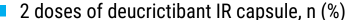
90.2% (202/224) of attacks achieved this milestone with a single dose of deucrictibant IR capsule.

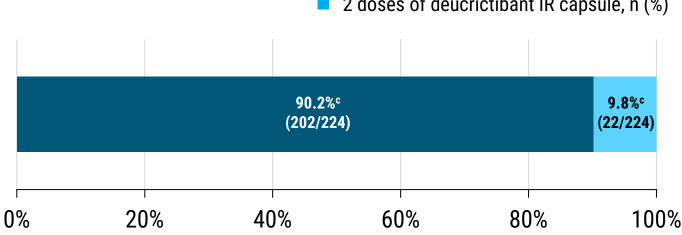


(224/261b) of attacks achieved complete attack resolution within 24 hours using PGI-Sa

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



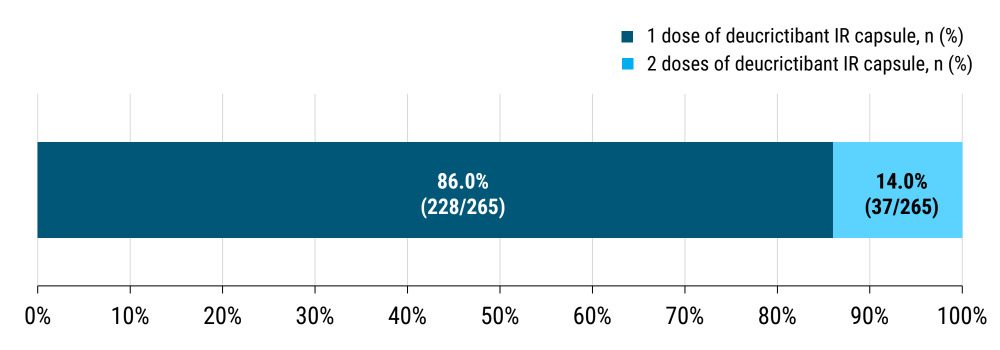




CI, confidence interval; IR, immediate-release; PGI-S, Patient Global Impression of Severity. aPGI-S rating of "none" at 24 hours post-treatment. b261 attacks have non-missing pre-treatment PGI-S. Percentage of 224 attacks achieving complete attack resolution using PGI-S within 24 hours.

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

#### Attacks treated with 1 or 2 doses of deucrictibant



#### **Conclusions**

- In the current analysis of the ongoing RAPIDe-2 Phase 2/3 extension study, deucrictibant 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 deucrictibant 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 deucrictibant IR capsule for repeat treatment of HAE attacks.

The Authors and the Sponsor would like to thank all the people with HAE as well as all study site staff who participated in the RAPIDe-2 trial.