# Reporting of Administration Site Reactions with Parenteral Drugs for the On-Demand Treatment of Hereditary Angioedema Attacks – Analysis of the FAERS Database 2009 to 2022 Raffi Tachdjian<sup>1</sup>, Sinisa Savic<sup>2</sup>, Moshe Fridman<sup>3</sup>, Joao Frade<sup>4</sup>, Paul K. Audhya<sup>4</sup>

## Background

- Hereditary angioedema (HAE) is a rare genetic disease associated with recurrent and unpredictable episodic attacks of tissue swelling, which may be life-threatening if involving the airway
- Treatment guidelines recommend that patients should have access to on-demand therapy to treat attacks as early as possible to reduce the severity and duration of the attack<sup>1-3</sup>
- Currently available on-demand treatments for HAE attacks are administered subcutaneously or intravenously<sup>3</sup>
- The most common adverse events reported in clinical trials and post-marketing reports for most of the on-demand drugs are associated with administration site reactions, which range from 3% to 97% of patients<sup>4-6</sup>
- Few studies have examined the real-world burden of administration site reactions with parenteral on-demand treatments
- The current real-world study was driven by two objectives: first, to describe reported rates of all administration site adverse drug reactions (ADRs) for approved on-demand HAE therapies using the FDA's Adverse Event Reporting System (FAERS); and second, to detect administration site ADRs from clinical trials with higher-than-expected reporting rates compared to other parental drugs in the FDA registry

## Methods

### Data Source

- We searched FAERS data from 10/1/2009 to 3/31/2022 for human C1 inhibitor (pdC1-INH). ecallantide, icatibant, and recombinant C1 inhibitor (rhC1-INH)
- The FAERS database contains information on adverse event and medication error reports submitted to FDA by healthcare professionals (such as physicians, pharmacists, nurses, and others) and consumers (such as patients, family members, lawyers, and others)
- Cases were only included if the HAE drug was listed as the 'primary suspect' potentially leading to an administration site ADR

### Variable Creation

- For the first objective, to describe the reported rates of all administration site ADRs, the ADR preferred terms from the Medical Dictionary for Regulatory Activities Preferred Terms (MedDRA) were grouped into an ADR domain based on semantic and/or clinical similarity
- This process resulted in 18 overarching ADR domains (**Table 1**). For each drug and ADR domain, the number of reports was calculated per year from the time of the US approval through 3/31/2022
- For the second objective, only the preferred terms associated with injection site ADRs identified from clinical trials and denoted on approved HAE drug US package inserts were included (Table 2)
- An FDA report with at least one of the preferred terms in the composite was flagged as an ADR in this analysis
- For each drug-event pair, the reporting odds ratio (ROR) and the empirical Bayesian geometric mean (EBGM) were calculated to detect drug-ADR pairs with higher-than-expected reporting rates compared to all other drugs with the same route of administration in the FDA registry
- ROR estimates with a two-sided lower 95% confidence bound >1.0 were considered significant
- One-sided 95% lower confidence bound of the EBGM was generated, with values >1 considered significant. A strong signal of a disproportionately high event rate for a drug-event pair was declared when both the ROR and EBGM were significant
- When both RORs and EBGM values were ≥1, but not both significant, it was reported as an indication of a trend toward higher-than-expected reporting rates<sup>7</sup>

### Acknowledaments

The authors wish to thank Jason Allaire, PhD of Generativity Health Outcomes Research for his assistance with this poster. Funding for Dr. Allaire was provided by KalVista Pharmaceuticals.

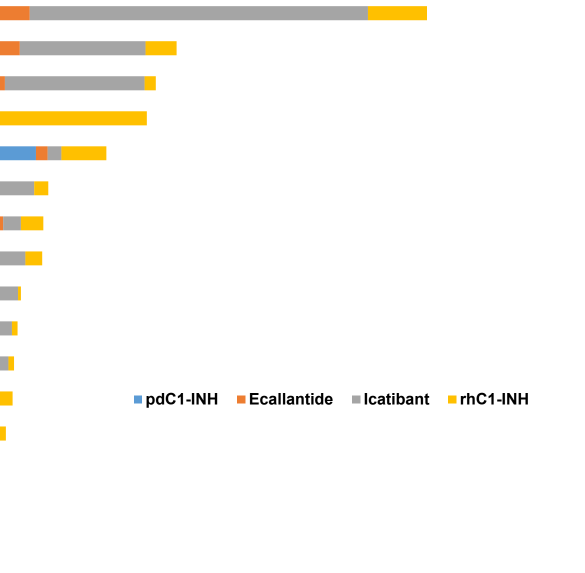
Disclosures Raffi Tachdjian discloses the following relationships with KalVista Pharmaceuticals (honoraria for advisory work), Takeda (honoraria for research, speaking, and advisory work), CSL Behring (honoraria for research, speaking, and advisory work), Pharming (honoraria for speaking and advisory work), Biocryst (honoraria for research, speaking and advisory work), Pharvaris (research grant). Sinisa Savic discloses the following relationships with KalVista Pharmaceuticals (honoraria for expenses and advisory work), CSL Behring (honoraria for expenses, advisory work, and research), Novartis (honoraria for expenses, advisory work, and research), and Biocryst (honoraria for advisory work). Moshe Fridman reports receiving consulting fee from KalVista Pharmaceuticals. Joao Frade and Paul K. Audhya are employees and own stock in KalVista Pharmaceuticals.

<sup>1</sup>UCLA School of Medicine, Los Angeles, CA, USA; <sup>2</sup>School of Medicine, University of Leeds, England; <sup>3</sup>AMF Consulting, Los Angeles, CA, USA; <sup>4</sup>KalVista Pharmaceuticals Cambridge, MA, USA

dministration Site ADR Domain	Administration Site ADR			
ncorrect Route of Product	Incorrect route of product administration			
Administration		Figure 1. Specific administration site		
Poor Venous Access	Poor venous access	demand HAE therapy Site Pain		
Site Pain	Infusion site pain	Site Swelling		
Site Bruising	Injection site pain	Site Swelling		
	Administration site pain	Site Erythema		
	Application site pain	Access Site Complication/Malfunction		
	Instillation site pain Vessel puncture site pain	Access Site Complication/Malfunction		
	Injection site bruising	Incorrect Route of Product Administration		
	Administration site bruise	Site Related Reaction		
	Infusion site bruising			
	Catheter site bruise	Poor Venous Access		
Site Emithema	Vessel puncture site bruise	Sito Bruising		
Site Erythema	Infusion site erythema Injection site erythema	Site Bruising		
	Catheter site erythema	Site Pruritus		
	Application site erythema	Site Urticere		
Site Swelling	Injection site swelling	Site Urticara		
	Infusion site swelling	Site Rash		
	Injection site edema	Oite Infection		
	Local swelling	Site Infection		
	Application site swelling Vascular access site swelling	Site Hemorrhage		
	Catheter site swelling			
Site Extravasation	Infusion site extravasation	Site Extravasation		
	Injection site extravasation	Site Mass		
	Catheter site extravasation			
Site Rash	Infusion site rash	Site Warmth		
	Catheter site rash	Site Vesicles		
	Injection site rash Application site rash			
Site Related Reaction	Infusion related reaction	Site Nodule		
	Injection related reaction			
	Injection site reaction	U		
	Infusion site reaction			
Site Hemorrhage	Infusion site hemorrhage	<ul> <li>The five most frequently reported ad</li> </ul>		
	Incision site hemorrhage	pain, site swelling, site erythema, ac		
	Injection site hemorrhage	· · · ·		
	Medical device site hemorrhage Application site hemorrhage	route of product administration (Figu		
	Catheter site hemorrhage	<ul> <li>Access site complications/malful</li> </ul>		
	Vascular access site hemorrhage			
Site Mass	Infusion site mass	reports per year		
	Injection site mass	<ul> <li>Icatibant had the highest reported ra</li> </ul>		
Site Nodule	Infusion site nodule	<b>e</b> .		
Site Infection Site Vesicles	Injection site nodule Injection site infection	(17.9 per year), site swelling (6.7 per		
	Vascular access site infection	had the highest rate of incorrect rout		
	Catheter site infection	Ũ		
	Infusion site infection	Figure 2. Total administration site A		
	Medical device site infection	demand HAE therapy		
	Injection site vesicles	demand the therapy		
	Application site vesicles	50		
Site Warmth	Injection site warmth	45		
Site Pruritus	Application site warmth Injection site pruritus	45		
Site Urticaria	Application site prunitus	40		
	Infusion site pruritus	95		
	Injection site urticaria	35		
	Infusion site urticaria	<u>ه</u> 30		
Access Site Complication/Malfunction	Vascular access complication			
	Vascular access site complication	<u></u> ້ອ 25		
	Vascular access malfunction	ස් 20		
le 2. ADR composites and	d associated preferred terms			
-	_	15 12.6		
Composite	Preferred Terms	10 9.2		
Administration Site Reactions	Injection site reactions			
	Injection site pain	5		
	Injection site redness	o		
	Injection site pruritus	pdC1-INH Ecallantid		
	Injection site irritation			
	Injection site urticaria Injection site bruising	<ul> <li>Figure 2 provides the total number of</li> </ul>		
	Injection site bruising	<b>.</b> .		
	Injection site hypoesthesia	HAE therapies		
	Injection site edema	a lootibout bod the meat administration		
	Injection site swelling	<ul> <li>Icatibant had the most administration</li> </ul>		
	Injection site warmth	had 24.3 ADRs reported per year		
	Injection site burning	amount of ADRs reported per yea		
	Injection site numbness			
	Injection site pressure sensation			

### Results

### ADRs per year by FDA-approved parenteral on



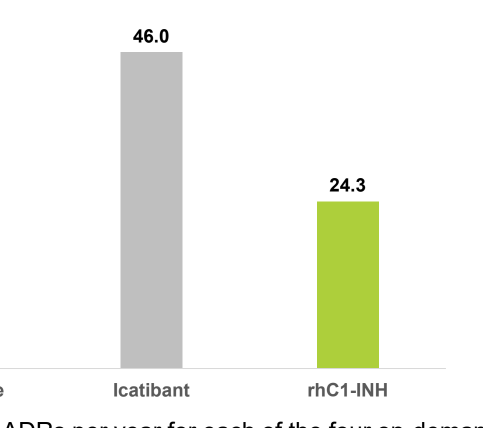


dministration site ADR domains included injection site ccess site complications/malfunctions, and incorrect ure 1)

unctions were only reported for rhC1-INH with 9.5

ate of administration site ADRs per year for site pain r year), and site erythema (7.4 per year). PdC1-INH Ite of product administration at 3.7 per year

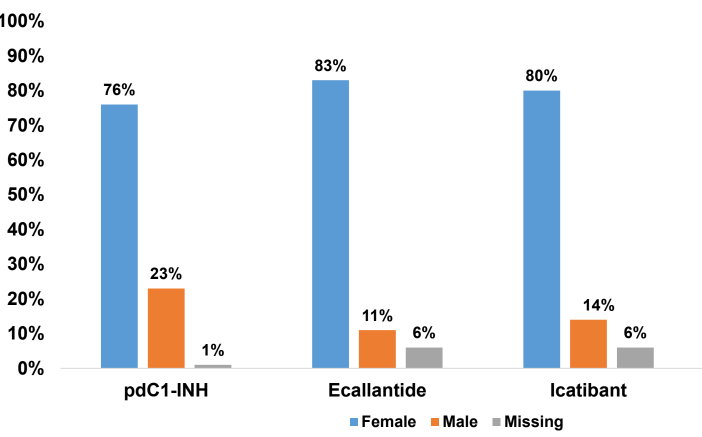
### ADRs per year by FDA-approved parenteral on-



f ADRs per year for each of the four on-demand

#### ation site ADRs reported per year with 46, rhC1-INH r, while pdC1-INH and ecallantide had a similar ear (12.6 vs 9.2, respectively)

### Figure 3. Sex distribution by FDA-approved parenteral on-demand HAE therapy



Female patients reported most of the reported administration site ADRs (**Figure 3**), with similar proportions of females across therapies (76% to 83%)

- PdC1-INH had the highest proportion of reports by males (23%), while ecallantide (11%) and icatibant (14%) had a similar proportion of reports by male patients
- No reports were by males who were using rhC1-INH
- The average age of HAE patients who reported administration site ADRs was similar across therapies (mean [SD], pdC1-INH, 42.9 [18.7]; icatibant, 42.3 [15.3]; rhC1-INH, 43.5 [9.4]), with the mean age of patients using ecallantide being nominally lower (37.5; [17.4])
- The age distribution was much narrower for rhC1-INH (SD=9.4) compared to the other three drugs, with standard deviations ranging from 15.3-18.7

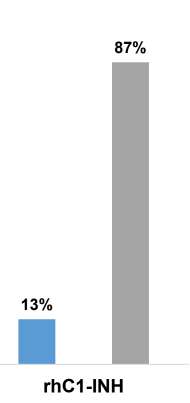
### Table 3. Reporting odds ratio and empirical Bayesian geometric mean values for HAE on-demand parenteral drugs

HAE Drug (FDA approval)	Adverse Drug Reaction	N of 'Primary suspect' cases	ROR (95% CI)	EBGM (95% CI Lower Bound)
pdC1-INH (Oct 12, 2009)	Administration site reactions composite (vs other IV) <sup>a</sup>	22	3.59 (2.36-5.46)	1.97 (1.39)
Ecallantide (Dec 1, 2009)	Administration site reactions composite (vs other SC)	41	0.39 (0.28-0.53)	0.38 (0.29)
lcatibant (Aug 25, 2011)	Administration site reactions composite (vs other SC) <sup>b</sup>	252	1.15 (1.01-1.30)	1.00 (0.90)
rhC1-INH (July 16, 2014)	Administration site reactions composite (vs other IV) <sup>b</sup>	19	2.85 (1.82-4.48)	1.32 (0.90)

CI confidence interval, ROR reporting odds ratio, EBGM empirical Bayesian geometric mean, IV intravenous, SC subcutaneous <sup>a</sup> Both ROR and EBGM lower-bound CI values were >1; ROR or EBGM lower-bound CI values >1

- The results of the predictive analysis examining the reporting rate of administration site reactions for each HAE drug are provided in **Table 3**
- PdC1-INH showed a statistically significant elevated reporting rate of injection site reactions (ROR=3.59 [2.36-5.46]; EBGM=1.97 [1.39])
- A trend toward increased reporting rate of injection site reactions was found for icatibant (ROR=1.15 [1.01-1.30]; EBGM=1.00 [0.90])
- Similarly, rhC1-INH showed a trend toward increased reporting rate of injection site reactions (ROR=2.85 [1.82-4.48]; EBGM=1.32 [0.90])

- References
- . Betschel S, Badiou J, Binkley K, et al. The International/Canadian Hereditary Angioedema Guideline. Allergy, Asthma & Clinical Immunology. 2019/11/25 2019;15(1):72. doi:10.1186/s13223-019-0376-8 Busse PJ, Christiansen SC, Riedl MA, et al. US HAEA Medical Advisory Board 2020 Guidelines for the Management of Hereditary Angioedema. The journal of allergy and clinical immunology In practice. Jan 2021;9(1):132-150.e3. doi:10.1016/j.jaip.2020.08.046
- 3. Maurer M, Magerl M, Betschel S, et al. The international WAO/EAACI guideline for the management of hereditary angioedema-The 2021 revision and update. Allergy. Jul 2022;77(7):1961-1990. doi:10.1111/all.15214 4. Kalbitor (ecallantide) {package insert]. Cambridge, MA: Dyax Corporation; 2009.
- Ruconest (c1 esteras e inhibitor recombinant injection) [package insert]. Warren, NJ: Pharming Healthcare Inc.; 2014.
- 6. Berinert (human c1-esterase inhibitor) [package insert]. Marburg, Germany; CSL Behring GmbH; 2009.
- 7. Bate A, Evans SJ. Quantitative signal detection using spontaneous ADR reporting. *Pharmacoepidemiol Drug Saf*. Jun 2009;18(6):427-36. doi:10.1002/pds.1742
- 8. Cicardi M, Banerji A, Bracho F, et al. Icatibant, a new bradykinin-receptor antagonist, in hereditary angioedema. *The New England journal of medicine*. Aug 5 2010;363(6):532-41. doi:10.1056/NEJMoa0906393
- 9. Lumry WR, Li HH, Levy RJ, et al. Randomized placebo-controlled trial of the bradykinin B<sub>2</sub> receptor antagonist icatibant for the treatment of acute attacks of hereditary angioedema: the FAST-3 trial. Annals of allergy, asthma & immunology: official publication of the American College of Allergy, Asthma, & Immunology. Dec 2011;107(6):529-37. doi:10.1016/j.anai.2011.08.015
- 10. Sakaeda T, Tamon A, Kadoyama K, Okuno Y. Data mining of the public version of the FDA Adverse Event Reporting System. International journal of medical sciences. 2013;10(7):796-803. doi:10.7150/ijms.6048 11. Hazell L and Shakir SA. Under-reporting of adverse drug reactions : a systematic review. Drug safety 2006; 29: 385-396. 2006/05/13. DOI: 10.2165/00002018-200629050-00003. 12. Hereditary Angioedema <u>www.fda.gov/files/about%20fda/published/The-Voice-of-the-Patient---Hereditary-Angioedema.pdf</u>



## Discussion

- The results of this real-world study suggest that all four FDA-approved ondemand therapies for HAE attacks are associated with administration site reactions
- The reported yearly rates in the FAERS database of administration site reactions were greatest for icatibant, especially for injection site pain
- This finding is not surprising given that the most common adverse effects in the pivotal FAST-1 and FAST-2 trials were injection-site reactions, which were experienced by most patients receiving icatibant in both trials (26 [96%] and 35 [97%], respectively)<sup>8,9</sup>
- Ecallantide had one of the lowest number of reported rates per year of administration site ADR in the current study, which mirrors the integrated analysis of the pivotal EDEMA3 and EDEMA4 clinical trials that found only 3% of patients reporting injection-site reactions<sup>4</sup>
- The results from the ROR and EBGM analyses indicated that administration site ADRs listed on the label of pdC1-INH were high when compared with other drugs utilizing the same administration route (Table 3)

## Limitations

- It should be noted that due to the nature of the FAERS registry, there are several limitations:
- Administration site ADRs are not exposure-adjusted and are based on spontaneous reporting; thus, they cannot be used to estimate incidence
- Reporting rates may differ among the included drugs
- Reporting rates may vary over time, with the highest rates typically in the first two years of commercial availability<sup>10</sup>
- Adverse events are significantly underreported in spontaneous reporting systems such as FAERS<sup>12</sup>

## Conclusions

- FAERS real-world data suggest that patients experience a treatment related burden due to administration site ADRs from the use of currently approved parenteral on-demand therapies for HAE attacks
- The current results are likely underestimating the real-world burden due to site administration ADRs
- These findings support the conclusions from the FDA Patient Voice Summit where patients stressed the difficulties surrounding parenteral administration and their desire for less invasive routes of administration<sup>11</sup>
- Evidence from patient-reported experiences, safety reports from clinical trials and post-marketing studies, and the real-world results presented here suggest that alternatives to on-demand therapies administered parenterally would reduce administration related burden to patients with HAE



