# A Phase 4 Randomized Double-blind Placebo-Controlled Study of Rimegepant for Acute Treatment of Migraine in Adults Unsuitable for Triptan Use

Messoud Ashina<sup>1</sup>, Peter McAllister<sup>2</sup>, Luz M Ramirez<sup>3</sup>, Catherine Nalpas<sup>4</sup>, Alexandra Thiry<sup>5</sup>, Lucy Abraham<sup>6</sup>, Robert Fountaine<sup>5</sup>, Terence Fullerton<sup>5</sup>

<sup>1</sup>Danish Headache Center, Rigshospitalet, Copenhagen, Denmark; <sup>2</sup>New England Institute for Neurology and Headache, Stamford, CT, USA; <sup>3</sup>Pfizer Inc, Princeton, NJ, USA; <sup>4</sup>Pfizer Inc, Paris, France; <sup>5</sup>Pfizer Inc, Groton, CT, USA; <sup>6</sup>Pfizer R&D UK Ltd, Tadworth, UK

### **BACKGROUND**

- There is an unmet treatment need for individuals with migraine who are unsuitable for triptans due to insufficient response, intolerance, or contraindication.1-3
- Post-hoc subgroup analyses from previous phase 3 trials suggest that rimegepant, an oral calcitonin gene-related peptide (CGRP) receptor antagonist, may be effective for acute treatment of migraine in individuals who previously discontinued triptans.4
- Prospectively designed trials in individuals unsuitable for triptans have not previously been conducted with gepants.

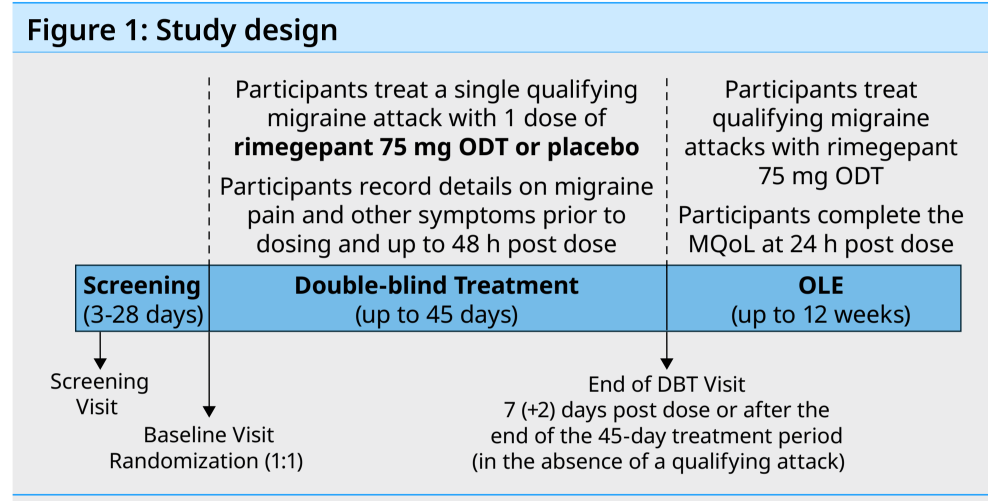
# **OBJECTIVE**

• To investigate the efficacy and tolerability of rimegepant for acute treatment of migraine in individuals unsuitable for triptans due to a documented history of prior inadequate response and/or intolerance to multiple agents, or due to the presence of a contraindication.

# **METHODS**

#### STUDY DESIGN

 This was a phase 4, multinational, randomized, double-blind, placebo-controlled study (NCT05509400; **Figure 1**).



DBT=double-blind treatment; MQoL=Migraine Quality of Life Questionnaire; ODT=orally disintegrating tablet; OLE=open-label extension

### **PARTICIPANTS**

- Eligible participants were aged ≥18 years with ≥1-year history of migraine attacks (with or without aura), migraine onset prior to age 50 years, migraine attacks lasting an average of 4–72 h if untreated, and an average of 4–14 migraine days per month in the 3 months prior to screening.
- Participants were unsuitable for triptan therapy due to documented (A) history of prior intolerance or lack of efficacy to ≥2 triptans or (B) the presence of a contraindication.
- Documentation was within the medical/pharmacy record complemented by participant interview if needed – or via principal investigator interview of the treating physician.
- Participants on stable (≥3 months) preventive migraine treatment (excluding CGRP antagonists) were eligible.

# **TREATMENT**

- Participants treated a single qualifying migraine attack with rimegepant 75 mg orally disintegrating tablet (ODT) or placebo.
- A qualifying migraine attack was defined as an attack of moderate or severe pain intensity first treated with study intervention, not with non-study medication (eg, NSAID).
- Participants rated migraine pain and other symptoms prior to dosing and up to 48 h post dose.

# **ENDPOINTS**

- The primary endpoint was the percentage of participants with migraine pain relief (no or mild migraine pain) at 2 h post dose.
- Key secondary endpoints included the percentage of participants with migraine pain freedom (no migraine pain) at 2 h post dose, rescue medication use within 24 h post dose, return to normal function at 2 h post dose, sustained return to normal function from 2–24 h and from 2–48 h post dose, sustained migraine pain relief from 2–24 h and from 2–48 h post dose, sustained pain freedom from 2–24 h and from 2–48 h post dose, and most bothersome symptom (MBS) freedom (absence of nausea, photophobia, or phonophobia) at 2 h post dose.
- On-treatment safety included the number and percentage of participants with adverse events (AEs), serious AEs, grade 3 or 4 clinical laboratory abnormalities, and liver function test elevations.

# **ANALYSIS**

- Efficacy was assessed in all participants who were randomized once, had a qualifying migraine attack at time of dosing, took double-blind study intervention, and had post-dose efficacy data.
- Treatment groups were compared using Mantel-Haenszel risk estimation. Type I error was controlled using hierarchical testing whereby the primary endpoint was evaluated at a 2-sided alpha level of 0.05.
- If the primary endpoint was significant, key secondary endpoints were each tested at a 2-sided alpha level of 0.05 in the pre-specified order.
- Safety was summarized descriptively in all participants who took double-blind study intervention.

# **RESULTS**

#### **PARTICIPANTS**

- 585 participants took double-blind study intervention (rimegepant, n=295; placebo, n=290) and 570 were analyzed for efficacy (rimegepant, n=286; placebo, n=284).
- Demographic and clinical characteristics were similar between treatment groups (**Table 1**).
- 93.5% of participants analyzed for efficacy had a documented failure to ≥2 triptans due to lack of efficacy and/or prior intolerance and 9.1% had a contraindication (**Table 1**).

#### Table 1: Summary of demographics and baseline clinical characteristicsa

	Rimegepant 75 mg	Placebo
Demographic/Characteristic	n=295	n=290
Age, mean (SD), years	43.0 (11.8)	42.7 (11.5)
Sex, n (%)		
Female	260 (88.1)	261 (90.0)
Male	35 (11.9)	29 (10.0)
Race, n (%) <sup>b</sup>		
White	52 (91.2)	45 (83.3)
Black or African American	5 (8.8)	8 (14.8)
Multiple	0	1 (1.9)
Body mass index, mean (SD), kg/m <sup>2</sup>	25.3 (4.3)	25.5 (4.5)
Age at migraine onset, mean (SD), years <sup>c</sup>	19.7 (9.0)	19.7 (9.5)
Number of moderate to severe migraine days per month in the previous 3 months, mean (SD) <sup>c</sup>	6.7 (2.5)	6.6 (2.6)
Average duration of untreated attacks, mean (SD), h <sup>c</sup>	41.3 (21.6)	43.0 (20.2)
Primary migraine type, n (%) <sup>c</sup>		
Without aura	227 (76.9)	222 (76.6)
With aura	68 (23.1)	68 (23.4)
Reason for triptan unsuitability, n (%)	Overall (N=570)d	
Documented failure to ≥2 triptans	533 (93.5)	
With ≥1 reason due to lack of efficacy	484 (84.9)	
With ≥1 reason due to prior intolerance	174 (30.5)	
With ≥1 reason due to lack of efficacy and	125 (21.9)	
≥1 reason due to prior intolerance		
Contraindication to triptans	52 (9.1)	

<sup>a</sup> Summarized in all participants who took double-blind study intervention unless noted. <sup>b</sup> Race was only assessed among US participants (rimegepant, n=57; placebo, n=54).

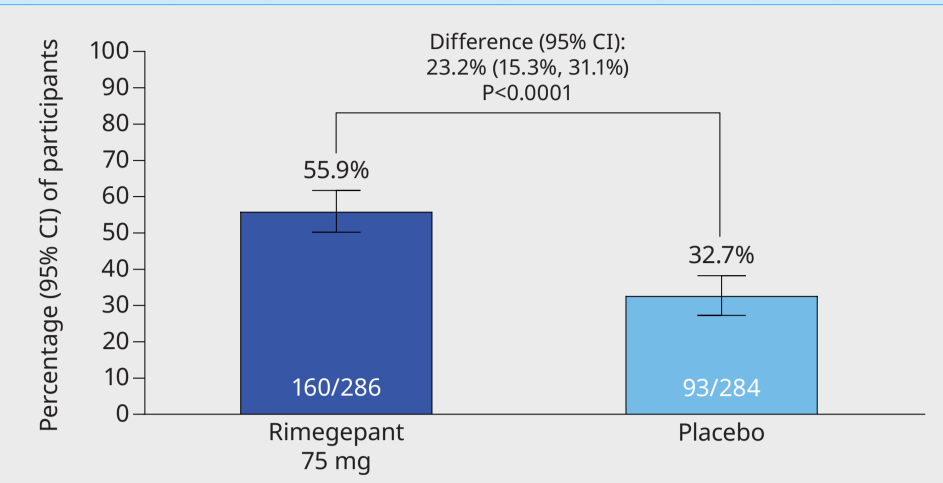
<sup>c</sup> Based on self-reported migraine history.

d Summarized in all participants who were randomized once, had a qualifying migraine attack at time of dosing, took double-blind study intervention, and had post-dose efficacy data (rimegepant, n=286; placebo, n=284).

# **EFFICACY**

- Rimegepant was superior to placebo for the primary endpoint of migraine pain relief at 2 h post dose (Figure 2).
- The percentage of participants with pain relief at 2 h was 55.9% for rimegepant and 32.7% for placebo; difference: 23.2% (95% CI, 15.3, 31.1); P<0.0001.

#### Figure 2: Migraine pain relief at 2 h pose dose (primary endpoint)



Analyzed in all participants who were randomized only once, had a qualifying migraine attack at the time of dosing, took double-blind study intervention, and had post dose efficacy data. Groups were compared using Mantel-Haenszel risk estimation.

Rimegepant was also superior to placebo for all 10 alpha-protected key secondary endpoints (Figure 3).

- This included assessments of acute effects at 2 h post dose (migraine pain freedom, return to normal function, MBS freedom), assessments of sustained effects from 2–24 and 2–48 h post dose (migraine pain relief, migraine pain freedom, return to normal function), and assessment of rescue medication use within 24 h post dose.

#### **SAFETY**

- AE frequency was similar in the rimegepant (12.5%) and placebo (12.1%) groups (**Table 2**).
- Only nasopharyngitis occurred in ≥1% of participants in the rimegepant group (rimegepant, 1.7%; placebo, 1.0%).
- No severe AEs, serious AEs, grade 3 or 4 laboratory test abnormalities, alanine aminotransferase or aspartate aminotransferase levels >3x upper limit of normal (ULN), or total bilirubin levels >1.5x ULN were reported among rimegepant-treated participants.

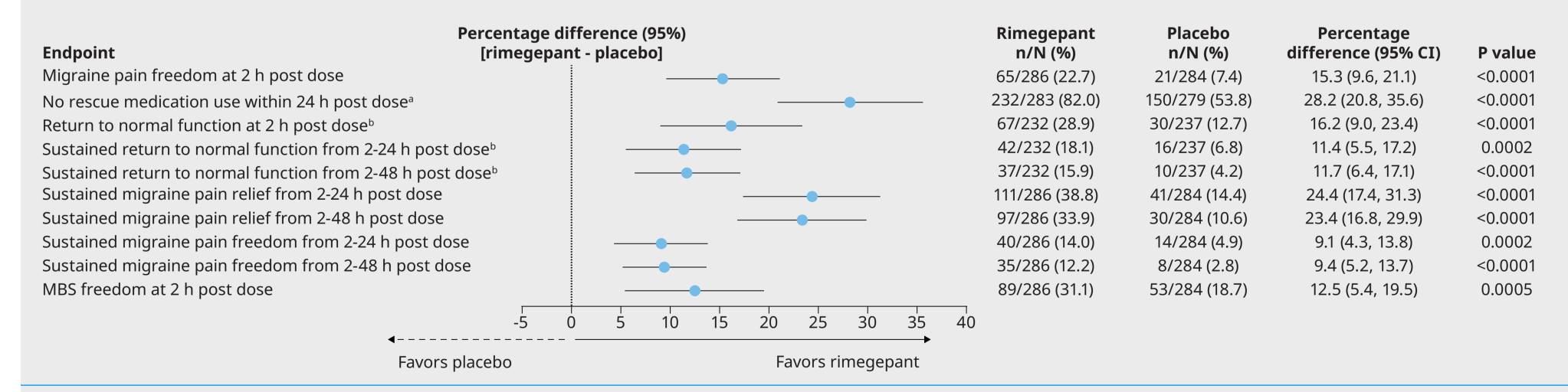
# Table 2: Summary of on-treatment adverse events<sup>a</sup>

	Rimegepant	
	75 mg	Placebo
AE, n (%)	n=295	n=290
Any AE	37 (12.5)	35 (12.1)
AE related to study drug	10 (3.4)	10 (3.4)
Mild AE <sup>b</sup>	31 (10.5)	19 (6.6)
Moderate AE <sup>b</sup>	6 (2.0)	15 (5.2)
Severe AE <sup>b</sup>	0	1 (0.3)
Serious AE	0	0
Hypertension AE	1 (0.3)	0
Raynaud's phenomenon AE	1 (0.3)	0

<sup>a</sup> Summarized in all participants who took double-blind study intervention.

<sup>b</sup> Based on preferred term worst intensity. AE=adverse event

# Figure 3: Summary of key secondary endpoints



Analyzed in all participants who were randomized only once, had a qualifying migraine attack at the time of dosing, took double-blind study intervention, and had post dose efficacy data. Testing utilized a hierarchical gate-keeping approach to control type 1 error. First, the significance of the primary endpoint was evaluated at a 2-sided alpha level of 0.05 level. Because the primary endpoint was significant, key secondary endpoints were each tested hierarchically at a 2-sided alpha-level of 0.05 in the prespecified order. Groups were compared using Mantel-Haenszel risk estimation. <sup>a</sup> Direction reversed so that positive percentage favors rimegepant. Participants with a first rescue medication use date being less than or equal to the study intervention dosing date +1 day, and missing the time of first rescue

medication use, were excluded

<sup>b</sup> Among participants with any level of functional disability (mild impairment, severe impairment, or requires bedrest) at time of dosing. MBS=most bothersome symptom; n=number of participants meeting the endpoint; N=number of evaluable participants

# CONCLUSIONS

- A single dose of rimegepant 75 mg ODT demonstrated superiority over placebo for the primary endpoint and all 10 key alpha-protected secondary endpoints, with a favorable tolerability profile that was similar to placebo.
- This is the first prospective controlled study to demonstrate efficacy of a gepant for the acute treatment of migraine in participants with a documented history of being unsuitable for triptans.
- Rimegepant may be a suitable option that addresses an unmet treatment need in this patient population.
- Findings from the 12-week open-label extension phase of this trial (currently ongoing) will allow for evaluation of the effectiveness of rimegepant and provide additional safety data in this population.

# REFERENCES

1. Lipton RB, et al. Cephalalgia 2020;40 (5):437-47. 2. Dodick DW, et al. J Prim Care Community Health 2020;11:2150132720963680. 3. Lipton RB, et al. Headache 2025;65:164-79. **4.** Lipton RB, et al. Cephalalgia 2023;43:1-11.

# **DISCLOSURES**

MA: Advisory board/consultant/speaker: AbbVie, Astra Zeneca, Eli Lilly, GlaxoSmithKline, Lundbeck, Pfizer, Teva; institutional research grant: Lundbeck, Lundbeck Foundation, Novartis, Novo Nordisk Foundation; associate editor: Journal of Headache and Pain, Brain. PM: Advisory board/speaker/consultant: AbbVie, ANI, BrightMind.AI, Dompe, Lilly, Lundbeck, Pfizer. LMR, CN, LA, RF, TF: Employees of and hold stock/options in Pfizer. AT: Former employee of Biohaven Pharmaceuticals; owns stock in Biohaven Ltd; employee and owns stock/options in Pfizer.



This study was sponsored by Pfizer. Medical writing support was provided by Matt Soulsby, PhD, CMPP, of Engage Scientific Solutions and was funded by Pfizer.



**Electronic Poster:** Please scan this QR code with your smartphone app to view this poster. If you do not have a smartphone, access the poster via the https://scientificpubs.congressposter.com/p/