

Acute Treatments for Migraine

Evidence Report

January 10, 2020

Prepared for



Please note there have been significant changes to this evidence report. Please refer to ICER's final evidence report.

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Steven Atlas served as the lead author for the report. Foluso Agboola led the systematic review and authorship of the comparative clinical effectiveness section in collaboration with Noemi Fluetsch and Eric Borrelli. Rick Chapman was responsible for oversight of the cost-effectiveness analyses and developed the budget impact model. Molly Beinfeld authored the section on coverage policies. David Rind and Steve Pearson provided methodologic guidance on the clinical and economic evaluations. Daniel Touchette and Todd Lee led the UIC modeling group and development of the cost-effectiveness model. The UIC team would like to thank Mrinmayee Joshi and Danny Quach for their contributions. The role of the UIC modeling group is limited to the development of the cost-effectiveness model, and the resulting ICER reports do not necessarily represent the views of UIC. None of the authors above disclosed any conflicts of interest.

About ICER

The Institute for Clinical and Economic Review (ICER) is an independent non-profit research organization that evaluates medical evidence and convenes public deliberative bodies to help stakeholders interpret and apply evidence to improve patient outcomes and control costs. Through all its work, ICER seeks to help create a future in which collaborative efforts to move evidence into action provide the foundation for a more effective, efficient, and just health care system. More information about ICER is available at <u>http://www.icer-review.org.</u>

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About Midwest CEPAC

The Midwest Comparative Effectiveness Public Advisory Council (Midwest CEPAC) – a core program of ICER – provides a public venue in which the evidence on the effectiveness and value of health care services can be discussed with the input of all stakeholders. Midwest CEPAC seeks to help patients, clinicians, insurers, and policymakers interpret and use evidence to improve the quality and value of health care.

The Midwest CEPAC is an independent committee of medical evidence experts from across the Midwest, with a mix of practicing clinicians, methodologists, and leaders in patient engagement and advocacy. All Council members meet strict conflict of interest guidelines and are convened to discuss the evidence summarized in ICER reports and vote on the comparative clinical effectiveness and value of medical interventions. More information about Midwest CEPAC is available at https://icer-review.org/programs/midwest-cepac/.

The findings contained within this report are current as of the date of publication. Readers should be aware that new evidence may emerge following the publication of this report that could potentially influence the results. ICER may revisit its analyses in a formal update to this report in the future.

The economic models used in ICER reports are intended to compare the clinical outcomes, expected costs, and cost effectiveness of different care pathways for broad groups of patients. Model results therefore represent average findings across patients and should not be presumed to represent the clinical or cost outcomes for any specific patient. In addition, data inputs to ICER models often come from clinical trials; patients in these trials and provider prescribing patterns may differ in real-world practice settings.

In the development of this report, ICER's researchers consulted with several clinical experts, patients, manufacturers and other stakeholders. The following clinical experts provided input that helped guide the ICER team as we shaped our scope and report. None of these individuals is responsible for the final contents of this report or should be assumed to support any part of this report, which is solely the work of the ICER team and its affiliated researchers.

For a complete list of stakeholders from whom we requested input, please visit: https://icer-review.org/material/acute-migraine-stakeholder-list/

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CHAMP (Coalition for Headache and Migraine Patients)

CHAMP accepts funding from a range of sources, including most medicine and device manufacturers that provide products in the headache, migraine and cluster space. Specifically, CHAMP receives sponsorship support from some of the companies that have products included in this review of Acute Treatments for Migraine.

Mia Minen, MD, MPH Assistant Professor, Department of Neurology NYU Langone NYU has IP rights to the RELAXaHEAD app of which Dr. Minen is the creator.

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List of Acronyms Used in this Report

95%CI95% confidence interval95%CrI95% credible intervalAAFPAmerican Academy of Family PhysiciansAANAmerican Academy of NeurologyAEAdverse eventACSAcute coronary syndromeAHRQAgency for Healthcare Research and QualityAHSAmerican Headache SocietyALTAlanine AminotransferaseASTAspartate AminotransferaseBCBSKCBlue Cross Blue Shield of Kansas CityBMIBody mass indexCADTHCanadian Authority for Drugs and Technologies in HealthCGRPCalcitonin gene-related peptideCHAMPCoalition for Headache and Migraine PatientsCHSCanadian Headache SocietyCMSCenters for Medicare and Medicaid ServicesCNSCentral nervous systemCVDCardiovascular diseaseD/CDiscontinuationHIVHuman Immunodeficiency VirusICERInstitute for Clinical and Economic ReviewICHDInternational Classification of Headache DisordersITTIntention-to-treatECGEchocardiogramEDEmergency departmentExcl.ExcludingEQ-5D-5LEuroQol 5-Dimension 5-Level ScaleevLYGEqual value of life years gainedFDAFood and Drug AdministrationGIGastrointestinalMBSMost bothersome symptommgMilligramMIMyocardial infarctionMIDASMigraine Diability AssessmentmITTM	5-HT	5-hydroxytryptamine
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N/A Not applicable	N/A	Not applicable

NCD	National Coverage Determinations
NICE	National Institute for Health and Care Excellence
NMA	Network meta-analysis
NR	Not reported
n.s.	Not significant
NSAIDs	Nonsteroidal anti-inflammatory drugs
OLE	Open label extension
OR	Odds ratio
PCE	Personal Consumption Expenditures Price Index
PCI	Percutaneous coronary intervention
PGIC	Patient Global Impression of Change
PICOTS	Populations, Interventions, Comparators, Outcomes, Timing, Settings
PRISMA	Preferred Reporting Items for Systematic Reviews and Meta-Analyses
PRN	As needed
QALY	Quality-adjusted life year
QOD	Every other day
RCT	Randomized Controlled Trial
SAE	Serious adverse event
SD	Standard deviation
TEAE	Treatment-emergent adverse event
TIA	Transient ischemic attack
SUD	Substance use disorder
ULN	Upper Limit Normal
US	United States
USPSTF	US Preventive Services Task Force

Executive Summary

Background

Migraine is a common cause of headache and is characterized by episodic, recurrent attacks that are classically pulsatile or throbbing, frequently involve one side of the head, and are associated with nausea and sensitivity to external stimuli such as light, sound, and smells. Migraine attacks vary in their frequency and intensity, but when severe can be a disabling, chronic condition that can impact all aspects of life including personal relationships and ability to work.¹ An estimated 40 million adults or 12-15% of adults in the United States (US) report migraine or severe headaches.^{2 3} Patients with migraine have higher costs of care, decreased work productivity, increased disability claims and account for \$11-50 billion in total costs.^{4-6 7,8 9}

The precise cause of migraine is not known and there is no specific test to confirm the diagnosis.¹⁰⁻¹² Migraine often starts in early adulthood, is more common in women, runs in families, and attacks can be triggered by a variety of predisposing factors such as stress and certain stimuli, activities and foods.^{2,3,13,14} Treatment broadly includes acute therapies to quickly abort episodic symptoms and ongoing therapies to reduce the frequency of attacks.¹² This review examines acute treatments for migraine attacks. Early acute treatment is especially helpful for individuals with aura (focal neurologic symptoms, frequently involving the visual system) that precede the onset of the headache. For those not responding to over-the-counter nonspecific pain medications or with moderate or severe symptoms, the use of specific migraine medications is recommended.

The most commonly used migraine specific medication class for acute treatment are "triptans" (5-hydroxytryptamine (5-HT) 1b/1d receptor agonists) available as pills, nasal sprays, and for injection under the skin.¹¹ Though effective and safe for patients with migraine, for many patients triptans are not adequately helpful or lose efficacy over time, have intolerable side effects, or have contraindications to their use (e.g., cardiovascular disease).^{15,16} The need for new therapeutic options is highlighted by the persistent use of medications, such as barbiturates and opioids that have the potential for misuse, and recognition that frequent use of acute medications can lead to medication overuse headaches.

New therapeutic classes include calcitonin gene-related peptide (CGRP) antagonists and 5hydroxytryptamine (5-HT) 1f agonists. Interest in CGRP antagonists has been driven by the observation that administration of CGRP can trigger acute headache and delayed migraine-like attacks.^{17,18} In addition, monoclonal antibodies targeting the CGRP receptor are being used for migraine prophylaxis.¹⁹ Two new oral CGRP receptor antagonists, ubrogepant (Ubrelvy™, Allergan, FDA approved on December 23, 2019) and rimegepant (under FDA review) have been studied for acute treatment of migraine attacks (class is referred to as "gepants"). Lasmiditan (Reyvow™, Lilly), a selective 5-HT 1f agonist (also referred to as a "ditan") approved on October 11, 2019 by the FDA for acute treatment of migraine, is thought to work in a similar manner to the triptans. Unlike the triptans, the gepants and lasmiditan do not have vasoconstrictive effects.^{16,20,21}

Insights Gained from Discussions with Patients and Patient Groups

Discussions with individual patients and patient advocacy groups identified important insights. We received numerous comments in which patients with migraine describe different personal stories and highlighted common themes that emphasize migraine as an episodic and chronic disease that can profoundly affect all aspects of their lives and the lives of those close to them. Though some have derived benefit from existing therapies, not all respond, headaches can recur as treatment wears off during the acute episode, response can vary from one migraine attack to another, and response can decrease over time with repeated episodic use. For others, side effects have led them to stop therapy or they have contraindications to the use of certain therapies. The net result is that for many patients with moderate or severe migraine headaches there is no single or combined therapy that offers them reliable, long-term control of their acute attacks.

A wide range of deficiencies with currently available acute treatments for migraine were noted.

- Despite a number of non-prescription and prescription medications, used alone or in combination, many patients cannot reliably prevent or abort migraine attacks.
- Available therapies do not provide symptom relief from migraine attacks with minimal side effects for many individuals.
- Triptans are effective in acute therapy for migraines but for many individuals they do not work, have intolerable side effects, or have contraindications to their use.
- For these reasons, patient turn to other medications such as opioids, barbiturates and antiemetics, but these also have limited benefit, acute side effects or risks with long-term use.

The profound impact of migraine on the lives of patients with migraine was emphasized.

- Migraine often develops in individuals during adolescence and young adulthood; formative educational years, where it can prevent them from reaching their full academic potential.
- Unpredictability of migraine attacks can result in anxiety from not knowing when the next attack will come, impacting individuals even when they do not have migraine symptoms.
- Frequent, severe attacks can have a dramatic impact on quality of life that may not be fully appreciated by the general public and even health professionals.
- As a result, migraine is a chronic condition that affects patients throughout their lives, disrupting personal relationships with friends and family.

The toll on patients with migraine includes important economic consequences.

- If the migraine attack is not aborted quickly and without medication related side effects, ability to work or work productively is profoundly affected.
- Acute treatments for migraine that work quickly and without side effects critically impact the ability to continue to work following a migraine attack.
- Frequent, severe and unpredictable migraine attacks combine to impact the ability to work, productivity when working, and risk of disability.
- The net result can be long-term un/under-employment with major socioeconomic cost that can have a long-term negative economic impact on the patient and her/his family.

Use of opioids and barbiturates for acute migraine is driven by limitations of existing therapies.

- Though recognized as having limited effectiveness, acute side effects, the potential for causing medication overuse headache and misuse, doctors end up prescribing them.
- New therapeutic classes, especially ones without side effects or limitations seen with triptans, may have a broader potential impact on the opioid crisis in the US.

Patient advocacy organizations also raised systematic issues that they felt needed to be addressed.

- Common outcome measures required by the FDA to obtain approval for new drugs may not adequately capture the impact of migraine on overall quality of life.
- Specifically, single dose studies are not designed to assess whether new therapies decrease the frequency of attacks over time or prevent medication overuse headaches.
- Successful migraine treatment may also help patients with other illnesses, such as anxiety and depression, that are impacted by frequent, unpredictable and severe attacks.

Potential Cost-Saving Measures in Migraine

Allergan suggested that opioids represent a low-value service that could be reduced.

Comparative Clinical Effectiveness

We evaluated the comparative clinical effectiveness and safety of lasmiditan, rimegepant and ubrogepant for the acute treatment of patients with migraine. Comparators of interest included: 1) no additional migraine-specific acute treatment (i.e., placebo arms of clinical trials) for patients with migraine attacks not adequately treated with non-prescription medicines and for whom triptans have not been effective, are not tolerated, or are contraindicated, and 2) triptans (eletriptan and sumatriptan) for patients who have migraine attacks that have not adequately responded to non-prescription medicines. The specific triptans were chosen because sumatriptan is one of the most

widely used triptans in clinical practice and eletriptan was shown in a recent network meta-analysis to be one of the most efficacious and well tolerated.^{15,22} We only examined oral triptan formulations because the new agents under review are all orally available.

We identified three RCTs of lasmiditan (1 Phase II and 2 Phase III),²³⁻²⁵ four RCTs of rimegepant (1 Phase II and 3 Phase III),²⁶⁻²⁹ and three RCTs of ubrogepant (1 Phase II and 2 Phase III)³⁰⁻³². All the RCTs of the interventions are placebo-controlled, except for one Phase II trial of rimegepant that also included sumatriptan as an active control arm.²⁹ We did not identify any trials comparing lasmiditan or ubrogepant to a triptan. In addition, we identified 23 RCTs of triptans (18 placebo-controlled trials of sumatriptan, three placebo-controlled trials of eletriptan and two head-to-head trials of sumatriptan and eletriptan) that met our inclusion criteria. ³³⁻⁵⁴

All the identified studies were large multicenter studies focused on the treatment of a singlemigraine attack. The trials enrolled patients who had at least a one-year history of migraine with or without aura as specified by the International Classification of Headache Disorders (ICHD) diagnostic criteria, who experienced two to eight migraine attacks (1 to 6 in triptan trials) of moderate to severe intensity per month, with age of onset before 50 years. Over 80% of the patients were female and the average age was approximately 40 years in each trial. Patients had been living with migraine for approximately 20 years, had an average of three to five migraine attacks per month, and about 20% to 25% of patients in the trials of the interventions were on preventive migraine medication. Characteristics of the treated migraine attack were generally similar across trials, with more patients having moderate than severe headache pain intensity (70% vs. 30%) at baseline. Photophobia was the most common other symptom reported (75% to 90% of patients) and was reported as the most bothersome symptom by 50% to 60% of patients. Approximately 40% to 65% of patients reported nausea, and 55% to 75% of patients reported phonophobia.

We considered all trials sufficiently similar to include in network meta-analyses.

Clinical Benefits

Pain Freedom and Pain Relief at Two Hours

The primary efficacy endpoint in all trials of lasmiditan and CGRP antagonists was freedom from pain at two hours after treatment, before the use of any rescue medication. Pain relief, defined as a decrease in headache pain from moderate or severe at baseline to mild or no pain at two hours after treatment and before taking any rescue medication was measured as a secondary outcome in the trials. Patients with moderate or severe pain who achieve pain freedom would also be counted as having pain relief. Overall, a greater proportion of patients achieved freedom from pain and pain relief at two hours post dose with the interventions compared to placebo (see Table ES1).

Table ES1. Phase III Results of Lasmiditan, Rimegepant and Ubrogepant. Pain Freedom and PainRelief at 2-Hours

Intervention (Trial)	Arms	Headache Pain Freedom at 2- Hours	Headache Pain Relief at 2-Hours	
		n/N (%)	n/N (%)	
Lasmiditan	Lasmiditan 200mg	167/518 (32.2)	330/555 (59.5)	
(SAMURAI){Goadsby, 2019, 2008}	Lasmiditan 100mg	142/503 (28.2)	334/562 (59.4)	
	Placebo	80/524 (15.3)	234/554 (42.2)	
	Lasmiditan 200mg	205/528 (38.8)	367/565 (65.0)	
Lasmiditan (SPARTAN){Kuca, 2018, 2006}	Lasmiditan 100mg	167/532 (31.4)	370/571 (64.8)	
	Placebo	115/540 (21.3)	274/576 (47.7)	
Rimegepant	Rimegepant 75mg	104/543 (19.2)	304/543 (56.0)	
(Study 301){Lipton, 2018, 1011}	Placebo	77/541 (14.2)	247/541 (45.7)	
Rimegepant	Rimegepant 75mg	105/537 (19.6)	312/537 (58.1)	
(Study 302){Lipton, 2019, 1012}	Placebo	64/535 (12.0)	229/535 (42.8)	
Rimegepant	Rimegepant 75mg	142/669 (21.2)	397/669 (59.3)	
(Study 303){Croop, 2019, 2003}	Placebo	74/682 (10.9)	295/682 (43.3)	
Ubrogepant	Ubrogepant 100mg	95/448 (21.2)	275/448 (61.4)	
(ACHIEVE I){Dodick, 2019, 1058}	Ubrogepant 50mg	81/422 (19.2)	257/422 (60.7)	
	Placebo	54/456 (11.8)	224/456 (49.1)	
Ubrogepant	Ubrogepant 50mg	101/464 (21.8)	291/464 (62.7)	
(ACHIEVE II){Lipton, 2019, 1057}	Placebo	65/456 (14.3)	220/456 (48.2)	

95% CI: 95% confidence interval, mg: milligrams, n: number of participants, N: total number of participants, vs: versus

Results of the NMA model are presented in terms of the odds ratio (OR) of freedom from pain (or pain relief) for each intervention versus placebo, sumatriptan and eletriptan (Table ES2. and Table 3). ORs above 1 indicate higher odds of pain freedom at two hours with the active intervention versus comparator while ORs below 1 indicate lower odds. Lasmiditan, rimegepant, and ubrogepant all had higher odds of achieving pain freedom at two hours versus placebo. Compared to each other, none of the interventions showed statistically significant differences, though lasmiditan showed a statistically nonsignificant, higher odds of achieving pain freedom. In contrast, all interventions showed lower odds of achieving pain freedom compared to eletriptan and sumatriptan. However, statistical significance was not reached for lasmiditan versus sumatriptan. Similar trends were observed for pain relief at two hours (Table ES3).

Lasmiditan (100/200 mg)					
1.43 (0.97, 2.06)	Rimegepant 75 mg				
1.43 (0.93, 2.14)	1 (0.69, 1.46)	Ubrogepant (50/100 mg)		_	
0.73 (0.53, 1.06)	0.51 (0.39, 0.7)	0.52 (0.37, 0.74)	Sumatriptan (50/100 mg)		_
0.54 (0.36, 0.85)	0.38 (0.27, 0.57)	0.38 (0.26, 0.59)	0.73 (0.57, 0.97)	Eletriptan 40 mg	
3.01 (2.2, 4.14)	2.11 (1.67, 2.72)	2.12 (1.58, 2.88)	4.09 (3.43, 4.82)	5.6 (4.14, 7.23)	Placebo

Table ES2. NMA results. Interventions and Comparators. Pain Freedom at 2-Hours

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Table ES3. NMA results. Interventions and Comparators. Pain Relief at 2-Hours

Lasmiditan (100/200 mg)		_			
1.16 (0.87, 1.52)	Rimegepant 75 mg				
1.15 (0.85, 1.58)	1 (0.75, 1.34)	Ubrogepant (50/100 mg)			
0.84 (0.67, 1.13)	0.73 (0.58, 0.96)	0.73 (0.55, 1)	Sumatriptan (50/100 mg)		_
0.61 (0.44, 0.88)	0.52 (0.38, 0.76)	0.52 (0.37, 0.78)	0.72 (0.58, 0.89)	Eletriptan 40 mg	
2.53 (2.04, 3.25)	2.19 (1.8, 2.76)	2.19 (1.7, 2.89)	2.99 (2.65, 3.34)	4.18 (3.32, 5.14)	Placebo

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Sustained Pain Freedom

Sustained pain freedom refers to individuals who were pain free at two hours and maintained pain freedom with no use of rescue medication or relapse within 24 (sustained pain freedom at 24 hours) or 48 hours (sustained pain freedom at 48 hours) after the initial treatment. The results of the NMA results on 24 hours sustained pain freedom are presented in Table ES4. Similar to the two-hour results, a greater proportion of patients on the interventions achieved sustained pain freedom at 24 hours versus placebo. Although all interventions showed lower odds of achieving sustained pain freedom at 24 hours compared to sumatriptan and eletriptan, these were not statistically significant. Similarly, the interventions were not statistically significantly different from each other.

Table ES4. NMA Results. All Interventions and Comparators. Sustained Pain Freedom at 24-Hours

Lasmiditan (100/200 mg)					
1.16 (0.67, 1.94)	Rimegepant (75 mg)				
1.26 (0.72, 2.11)	1.08 (0.67, 1.74)	Ubrogepant (50/100 mg)			
0.83 (0.5, 1.44)	0.71 (0.48, 1.12)	0.66 (0.41, 1.12)	Sumatriptan		
0.73 (0.34, 1.53)	0.63 (0.32, 1.22)	0.59 (0.28, 1.18)	0.89 (0.44, 1.69)	Eletriptan	
2.92 (1.89, 4.5)	2.51 (1.89, 3.46)	2.32 (1.62, 3.46)	3.53 (2.52, 4.77)	3.97 (2.24, 7.36)	Placebo

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Freedom from Most Bothersome Symptom (MBS)

Absence of the most bothersome migraine associated symptom (i.e. phonophobia, photophobia, or nausea) at two hours after treatment was measured as a co-primary endpoint in the Phase III trials of lasmiditan, rimegepant and ubrogepant. However, none of the triptan studies assessed freedom from MBS as an outcome. The NMA results showed that lasmiditan (1.69, 95% CrI: 1.33, 2.14), rimegepant (1.58, 95% CrI: 1.29, 1.94), and ubrogepant (1.64, 95% CrI: 1.28, 2.12) all had higher odds of achieving freedom from MBS at two hours post dose compared to placebo. However, compared to each other, none of the interventions showed a statistically significant difference.

Disability

Functional disability assessed at two hours was measured as a secondary outcome in all the Phase III trials of the interventions, but not consistently in the triptan studies. As such we included only the seven Phase III trials in our NMA and compared the interventions to each other and to placebo. The NMA results showed that lasmiditan (1.70, 95% CrI: 1.32, 2.20), rimegepant (1.72, 95% CrI: 1.38, 2.14), and ubrogepant (1.51, 95% CrI: 1.15, 1.96) all had higher odds of achieving no disability at two hours post dose compared to placebo. However, compared to each other, none of the interventions showed a statistically significant difference.

Harms

Harms assessed in the single-attack trials include treatment-emergent adverse events (TEAEs), serious adverse events (AEs), and any AE reported by at least 5% of a trial arm. Overall, the AEs observed in these trials were mild or moderate in intensity. The NMA results showed there were no differences in the odds of any AE and TEAE between rimegepant and ubrogepant versus placebo and triptans in the single-attack trials. However, lasmiditan had higher odds of causing TEAE

compared to placebo (5.99, 95% Crl: 3.3, 12.52, Table 3.15), rimegepant (4.00, 95% Crl: 1.38, 12.04), ubrogepant (5.10, 95% Crl: 2.31, 12.95), and sumatriptan (2.57, 95% Crl: 1.3, 6.07). Similar results were seen for any AE.

Nausea was among the most commonly reported AEs in the ubrogepant and rimegepant trials (1% to 3%). In the lasmiditan trials, central nervous system (CNS)-related AEs (e.g., dizziness [16-18%], somnolence [5-6%], paresthesia [2-7%)) were the most frequently reported AEs, with dizziness the most common. Results of the NMA on the incidence of dizziness across trials showed that lasmiditan had higher odds of causing dizziness compared to placebo (8.43, 95% Crl: 4.88, 19.35, see Table 3.16), rimegepant (7.02, 95% Crl: 2.2, 25.63), ubrogepant (4.95, 95% Crl: 1.67, 15.92), sumatriptan (4.09, 95% Crl: 2, 10.6), and eletriptan (3.97, 95% Crl: 1.44, 12.41).

In the open-label extension (OLE) study of lasmiditan, 12.8% of patients discontinued the trial due to adverse events (11.2% of patients in the 100 mg group, and 14.4% in the 200 mg group), and dizziness was reported to be the most common AE leading to discontinuation (2.7% of patients in the 100 mg group, and 4.3% of patients in 200 mg group). There was no incidence of abuse, misuse, or diversion related to the CNS effects of lasmiditan. Due to concerns about somnolence with lasmiditan, the FDA label advises that patients should not drive or operate machinery within 8 hours of taking a dose.⁵⁵ Compared to the lasmiditan OLE, rates of discontinuation were lower in the OLEs of rimegepant and ubrogepant (see Table 3.18).

Controversies and Uncertainties

We primarily used indirect quantitative methods (network meta-analyses) to compare lasmiditan, rimegepant and ubrogepant to each other because there were no head-to-head studies, and only one trial compared one of the interventions versus a triptan (rimegepant vs sumatriptan). Such indirect analyses have more uncertainty than had the therapies been compared directly.

The primary outcomes reported included efficacy and side effects of a single dose of each drug compared to placebo at two hours after initial study medication. Though patient and patient advocates highlighted the importance of outcomes after two hours, protocols for use of rescue medications and additional study medication dosing differed markedly among the trials making it difficult to assess the benefits of these drugs after two hours. While we looked at outcomes up to 48 hours, potentially important differences in efficacy among medications could be missed.

Limitations of current therapies including triptans has led to considerable interest in new therapies for acute treatment of migraine. How helpful these new drugs will be over time for these patients in terms of effectiveness and tolerance is uncertain. Though potentially an option for those with absolute or relative contraindications to triptans, such as heart disease, there is little clinical information on the safety of these new therapies for these individuals. Since most data presented results of these drugs for treatment of a single migraine attack, it is uncertain about their outcomes when used over time for repeated attacks. Important long-term outcomes such as the effect of these medications on potentially decreasing the frequency of migraine attacks, the occurrence of medication overuse headaches, and the need for other therapies such as opioids and barbiturates are currently not known. It is hoped that having more treatments for migraine can reduce use of opioids and thus the risk for opioid misuse.

Finally, migraine can have a dramatic impact on quality of life and ability to work for those with frequent, severe and unpredictable attacks. It is uncertain if these new therapies may help improve quality of life and work and productivity outcomes over time.

Summary and Comment

Lasmiditan, Rimegepant, and Ubrogepant Versus No Additional Migraine-Specific Acute Treatment (Placebo) or Triptans (Sumatriptan and Eletriptan)

Results from clinical trials and from our NMAs suggest that lasmiditan, rimegepant, and ubrogepant decrease symptoms of migraine attacks and improve function compared to placebo. Few harms were seen in the single-dose trials of lasmiditan, rimegepant and ubrogepant. However, lasmiditan showed a higher incidence of CNS related AEs (e.g., dizziness, somnolence, paresthesia) in the clinical trials.

Population 1: For adults (18 years and older) with moderate-severe migraine attacks that have not responded to non-prescription medicines and for whom triptans have not been effective, are not tolerated, or are contraindicated:

• We consider the evidence on lasmiditan, rimegepant and ubrogepant compared to placebo to be "incremental or better" (B+), demonstrating a moderate certainty of a small or substantial health benefit, with a high certainty of at least a small net health benefit.

Population 2: For adults (18 years and older) with migraine attacks that have not responded to non-prescription medicines (and are eligible to use triptans):

• We consider the evidence on lasmiditan, rimegepant and ubrogepant compared to triptans to be "comparable or inferior" (C-), demonstrating moderate certainty that the point estimate for comparative net health benefit is either comparable or inferior. Based on the results of the NMAs, rimegepant and ubrogepant appear to be less efficacious than triptans (sumatriptan and eletriptan) but have comparable short-term adverse events. For lasmiditan, the results of the NMAs suggest it is less efficacious than triptans, but the NMAs do not exclude comparable efficacy compared to sumatriptan. In terms of adverse events, the NMA results suggest a higher incidence with lasmiditan compared to triptans.

For all adults with migraine attacks:

 We consider the evidence on rimegepant and ubrogepant to be "comparable" (C), demonstrating a high certainty of a comparable net health benefit. For lasmiditan, the results of the NMAs suggest it may be slightly more efficacious than rimegepant and ubrogepant. However, the NMAs do not exclude comparable efficacy. Patients treated with lasmiditan had more adverse events and more of them discontinued treatment than patients treated with rimegepant or ubrogepant. We believe any possible greater efficacy of lasmiditan is at best balanced by these adverse events and may be outweighed by them, and thus we consider the evidence on lasmiditan compared to rimegepant and ubrogepant to be "comparable or inferior" (C-).

Long-Term Cost Effectiveness

Model Overview

The primary aim of this analysis was to estimate the cost effectiveness of lasmiditan, rimegepant, and ubrogepant among adults for the acute treatment of migraine using a decision analytic model. In the model, lasmiditan, rimegepant, and ubrogepant were compared with each other and to three comparators in separate analyses across two distinct populations. For the first comparison, we included patients who had migraine attacks that did not respond to non-prescription medicines and for whom triptans had not been effective, were not tolerated or were contraindicated. In this group, we compared lasmiditan, rimegepant, and ubrogepant to each other and to no additional migraine-specific acute treatment. For this analysis, no additional migraine-specific acute treatment was estimated by the placebo arms of the clinical trials, although we recognized that in the real-world, patients may use previously failed or untried over the counter and prescription treatments for acute migraine including analgesics. For the second comparison, we included patients who had migraine attacks that did not respond adequately to non-prescription medicines. In this analysis, we compared lasmiditan, rimegepant, and ubrogepant to each other and to two triptans: sumatriptan and eletriptan. Sumatriptan was chosen because it is one of the most widely used triptans in clinical practice; and eletriptan, a newer triptan, was shown in a recent network meta-analysis to be one of the most efficacious and well tolerated. Since these new agents under review are all available as oral preparations, we focused our comparison of triptans on the oral formulations.

We developed a *de novo* semi-Markov model with time-varying proportions of patients with response to treatment. The outcomes of interest included the incremental cost per quality-adjusted life year (QALY) gained, life-years gained, <u>equal value of life years gained (evLYG)</u>, and cost per hour of migraine pain avoided. The model was informed by a network meta-analysis of key

clinical trials and prior relevant economic models, systematic literature reviews, and input from stakeholders. The base case used a US health sector perspective with costs and outcomes discounted at 3% annually. The model cycle was 48 hours and the time horizon was two years.

Upon model entry, hypothetical patients entered one of two Markov states, either having a migraine or not having a migraine, based on the average daily rate of migraines. Among patients in the migraine health state, patients were classified as having moderate or severe migraine pain. The treatment response was evaluated at 2, 8, 24 and 48 hours. Patients could have complete resolution of migraine pain (pain freedom), improvement in migraine pain without complete resolution (pain relief) or no improvement. Patients with pain relief at each of the time points were classified as having mild migraine pain. The level of migraine pain was linked to utility values from the EQ-5D. Treatment response was linked with the probability of requiring a provider office visit, emergency department visit or hospitalization due to migraine. Rates of adverse events were linked to disutility values. Over time, patients were allowed to discontinue treatment due to side effects or insufficient effectiveness.

Key Assumptions

The model required several assumptions, which are described below.

Table ES5.	Key Model Assumptions
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Assumption	Rationale
Mortality is not associated with acute treatment for	There have been no demonstrated mortality benefits
migraine.	with treatment of migraine pain and other symptoms.
Acute treatment of migraine with lasmiditan, rimegepant, ubrogepant, and triptans does not affect migraine frequency.	Studies evaluating new migraine therapies were either short-term single episode studies or non-controlled open label studies and were not designed to demonstrate changes in migraine frequency with treatment. Longer- term, uncontrolled, open-label studies suffer from a possible placebo effect and a high likelihood that regression to the mean may affect the study's results. Should stronger evidence suggest that migraine frequency and/or characteristics are modified with acute treatments for migraine, this assumption will be reevaluated.
Patients receiving no benefit from treatment discontinued the medication in the first year of treatment only. There was no discontinuation for lack of effectiveness in the second year of the model.	Data describing treatment discontinuation due to lack of effect was obtained from a study in which follow up lasted for 12 months. ⁵⁶ It is unlikely that the majority of patients receiving no or suboptimal benefit would continue taking a medication beyond 12 months.
Patients who did not respond to acute treatments for migraine were assumed to have moderate or severe pain, in proportion to what was observed at baseline.	Sufficiently detailed data evaluating those who did not respond was not uniformly available from clinical trials. This assumption was necessary to assign utility values to those who did not respond to therapy.
If a migraine treatment resulted in migraine pain of "no pain" or "mild pain" at 2 hours, a person would be able to work.	· · ·

Model Inputs

Two-hour response to acute treatments for migraine was estimated using data directly from clinical trials included in a network meta-analysis. The proportion of patients who were pain free in clinical trials were considered to have "no pain" at the two-hour time point. The proportion of patients with "mild pain" were those who had pain relief but were not pain free. Those with no response remained in moderate or severe pain in proportion to what was observed at baseline. In clinical trials evaluating lasmiditan, rimegepant, and ubrogepant, some patients who responded at two hours subsequently lost response to treatment between 2 and 24 hours. The proportion of patients maintaining response at 24 hours was based on estimates from the network meta-analysis of

clinical trials. For the patients who lost response, we assumed the maximal proportion lost response at eight hours. After 8 hours, patients regained response where the rate of response among this group at 24 hours was equivalent to the placebo response rate. All patients responding at 2 hours were assumed to have treatment response at 48 hours.

Among patients who did not respond at two hours, the rate of response observed in this group was based on the rate of placebo response at 8 and 24 hours. Response at 48 hours was calculated by adding all two-hour responders to the placebo response for non-responders at two hours. Estimates of treatment response at 2, 8, 12, 24, and 48 hours are shown in the full report, Table 4.3.

The utilities used in the analysis were derived from published literature that estimated migrainespecific utility values using the EQ-5D and stratified by the severity of the migraine. The utility values used in the model were 0.959 for pain free, 0.835 for mild pain, 0.773 for moderate pain and 0.440 for severe pain. Hospitalized patients were assigned a disutility of -0.5 for 48 hours; those admitted to the emergency department were assigned a disutility of -0.5 for 24 hours. We did not include a disutility score for patients suffering from nausea and/or vomiting, photophobia, or phonophobia due to lack of data. Disutility of other adverse events, including drowsiness, dizziness, fatigue and paresthesia, were included in the model.

At the time of publishing this report, the prices for lasmiditan, rimegepant, and ubrogepant were not available. We therefore estimated the prices of lasmiditan, rimegepant, and ubrogepant for the model based on an opinion article that estimated that ubrogepant would have a 20% premium to branded Imitrex.⁵⁷ We applied the same premium to lasmiditan and rimegepant. All estimates generated in the model used these placeholder prices. Costs for sumatriptan and eletriptan were derived using wholesale acquisition cost (WAC). Costs for treatments for the usual care arm were estimated using the WAC prices for a prevalent mix of treatments.

Table ES6. Drug Cost per Dose

Drug	WAC	Notes
Lasmiditan	n/a (Used \$78.38)	20% premium pricing above Imitrex
Rimegepant	n/a (Used \$78.38)	20% premium pricing above Imitrex
Ubrogepant	n/a (Used \$78.38)	20% premium pricing above Imitrex
Sumatriptan, Oral tablets 50 mg 100 mg	\$1.04	
Eletriptan 40 mg	\$11.95	
Usual Care (mix)	\$4.81	

mg: milligrams, WAC: wholesale acquisition cost

Base-Case Results

The base-case results using the placeholder prices for lasmiditan, rimegepant, and ubrogepant are reported in Table ES7.

Table ES7. Base-Case Results for Lasmiditan, Rimegepant, Ubrogepant, Sumatriptan, Eletriptan,and Usual Care*

Treatment	Drug Cost (per year)**	Total Cost**	QALYs	Life Years	evLYG	Hours of Pain
Lasmiditan	\$3,970	\$13,640	1.8252	1.95	1.8252	1,743
Rimegepant	\$3,970	\$14,500	1.8222	1.95	1.8222	1,870
Ubrogepant	\$3,970	\$14,510	1.8221	1.95	1.8221	1,876
Sumatriptan	\$50	\$6,630	1.8264	1.95	1.8264	1,611
Eletriptan	\$590	\$6,790	1.8293	1.95	1.8293	1,484
Usual Care	\$0	\$10,050	1.8142	1.95	1.8142	2,100

evLYG: equal value of life years gained, LY: life year, QALY: quality-adjust life year

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

**Drug costs per year were calculated without accounting for discontinuation of the drug. Total costs take into account discontinuation and costs of alternative treatments.

The incremental cost-effectiveness results are reported in Table ES8. When evaluating the use of lasmiditan, rimegepant, and ubrogepant using the place-holder prices in Population 1, the ICERs for lasmiditan, rimegepant, and ubrogepant compared with usual care were \$327,700, \$559,500, and \$569,600 per QALY gained, respectively. When compared with each other and at the

placeholder prices used in the model and point estimates derived from the network metaanalysis, lasmiditan dominated rimegepant and ubrogepant, being more effective and less costly. However, there was significant overlap in the confidence intervals for lasmiditan and the point estimates for rimegepant and ubrogepant. Rimegepant and ubrogepant had nearly identical total costs, QALYs, and cost-effectiveness. The incremental cost effectiveness of lasmiditan, rimegepant, and ubrogepant will be dependent on the actual pricing of the three therapies. In Population 2, both sumatriptan and eletriptan produced higher QALYs at a lower total cost, and therefore dominated lasmiditan, rimegepant, and ubrogepant.

Treatment	Comparator	Cost per QALY Gained	Cost per Hour of Pain Avoided				
	Population 1						
Lasmiditan	vs. Usual Care	\$327,700	\$10.10				
Rimegepant	vs. Usual Care	\$559,500	\$19.41				
Ubrogepant	vs. Usual	\$569,600	\$19.41				
	Population 2						
Lasmiditan	vs. Sumatriptan	Dominated	Dominated				
Rimegepant	vs. Sumatriptan	Dominated	Dominated				
Ubrogepant	vs. Sumatriptan	Dominated	Dominated				
Lasmiditan	vs. Eletriptan	Dominated	Dominated				
Rimegepant	vs. Eletriptan	Dominated	Dominated				
Ubrogepant	vs. Eletriptan	Dominated	Dominated				

Table ES8. Incremental Cost-Effectiveness Ratios for the Base Case

QALY: quality-adjusted life years

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant

Sensitivity and Scenario Analyses

We conducted sensitivity analyses and scenario analyses to assess the impact of all model parameters on the estimated cost-effectiveness in population 1. The model was sensitive to several of the model inputs. However, in one-way sensitivity analyses, none of the individual model inputs that were varied resulted in an ICER below \$150,000 per QALY gained when using the assumed placeholder costs.

Probabilistic sensitivity analyses were also conducted to assess the variation across all parameters with 1,000 Monte Carlo simulations. When compared to usual care, none of the new treatments had cost-utility ratios in any of the iterations that were below \$50,000 per QALY gained or \$150,000 per QALY gained. At a cost-effectiveness threshold of \$250,000 per QALY gained, lasmiditan achieved cost-effectiveness in 9.4% and ubrogepant 0.2% of the trials.

Table ES9. Probabilistic Sensitivity Analysis Results Proportion of ICERs below specified
Thresholds for Lasmiditan, Rimegepant, Ubrogepant Compared with Usual Care (Placebo)*

Treatment Compared with Usual Care	Cost- Effective at \$50,000 per QALY	Cost- Effective at \$100,000 per QALY	Cost- Effective at \$150,000 per QALY	Cost- Effective at \$200,000 per QALY	Cost- Effective at \$250,000 per QALY
Lasmiditan	0.0%	0.0%	0.0%	0.2%	9.4%
Rimegepant	0.0%	0.0%	0.0%	0.0%	0.0%
Ubrogepant	0.0%	0.0%	0.0%	0.1%	0.2%

QALY: quality-adjusted life year

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

Scenario Analyses

The modified societal perspective included potential labor benefits for reduced migraine pain in the analysis. The ICERs for lasmiditan, rimegepant, and ubrogepant compared with usual care were \$207,800, \$422,900, and \$430,900 per QALY gained, respectively.

Threshold Analyses Results

Average annual prices that would result in willingness-to-pay thresholds of \$50,000 to \$150,000 per QALY gained for Population 1 are shown in table 4.12 below.

	Annual Price to Achieve \$50,000 per QALY	Annual Price to Achieve \$100,000 per QALY	Annual Price to Achieve \$150,000 per QALY
Lasmiditan	\$2,390	\$2,770	\$3,150
Rimegepant	\$1,960	\$2,210	\$2,460
Ubrogepant	\$1,950	\$2,200	\$2,440

Table ES10. Threshold Analysis Results for Population 1 (Patients Who Cannot Take Triptans)

QALY: quality-adjusted life year

Model Validation

Model validation followed standard practices in the field. We tested all mathematical functions in the model to ensure they were consistent with the report (and supplemental Appendix materials). Model calculations were verified, and model input parameters were varied to evaluate face validity of changes in results. We also conducted sensitivity analyses with null input values to ensure the model was producing findings consistent with expectations. Model validation was also conducted in terms of comparisons to other model findings.

Summary and Comment

In our analysis of the cost effectiveness of lasmiditan, rimegepant, and ubrogepant, we found that for patients for whom triptans are not effective, not tolerated, or are contraindicated (Population 1), if these drugs are priced with the place-holder prices used in this analysis, they will exceed commonly accepted thresholds for cost effectiveness. Also, they will be dominated by sumatriptan and eletriptan in patients who can take triptans (Population 2) in that sumatriptan and eletriptan are both more effective and less expensive than these newer agents.

When compared with usual care in patients in whom triptans are not effective, not tolerated, or are contraindicated, these new acute treatments for migraine provide utility gains. Pricing of these drugs will determine whether they are cost effective at commonly used thresholds in patients who cannot take triptan medications.

Potential Other Benefits and Contextual Considerations

Our reviews seek to provide information on potential other benefits offered by the intervention to the individual patient, caregivers, the delivery system, other patients, or the public that would not have been considered as part of the evidence on comparative clinical effectiveness. These elements are listed in the table below.

Potential Other Benefits

Table ES11. Potential Other Benefits

Other Benefits	Description
This intervention offers reduced complexity that will significantly improve patient outcomes.	Similar to most triptans, lasmiditan, rimegepant and ubrogepant are orally available medications and would not be expected to increase the complexity of care. The favorable side effects seen to date with rimegepant and ubrogepant, similar to those seen with placebo, may make these medications attractive to patients and clinicians. The restriction on driving after taking lasmiditan is a potential disadvantage of that therapy.
This intervention will reduce important health disparities across racial, ethnic, gender, socio-economic, or regional categories.	Not applicable
This intervention will significantly reduce caregiver or broader family burden.	New therapies for acute treatment of migraine may reduce caregiver and family burden if outcomes are improved for those in whom existing therapies do not effectively and safely control symptoms.
This intervention offers a novel mechanism of action or approach that will allow successful treatment of many patients for whom other available treatments have failed.	These new therapies reflect translational research in which improved understanding of the mechanisms of disease has led to new therapeutics. Lasmiditan, approved for migraine attacks, targets the 5HT1F (5-hydroxytryptamine 1F) receptor, and unlike the triptans does not induce vasoconstriction. The gepants, target CGRP, a peptide neural transmitter found in the pathways that play an important role in migraine. Ubrogepant is the first approved small molecule gepant and rimegepant is under review.
This intervention will have a significant impact on improving return to work and/or overall productivity.	The availability of new treatments for migraine is likely to allow some patients to remain at work in situations where they would otherwise have needed to miss or leave work. The restriction on driving after taking lasmiditan may negatively impact work/productivity outcomes.
Other important benefits or disadvantages that should have an important role in judgments of the value of this intervention.	Patients and advocates expressed the hope that these new therapies for patients with migraine may provide an effective and safe alternative for individuals who may turn to opioids and barbiturates because of limitations of existing therapies.

Contextual Considerations

Table ES12. Potential Contextual Considerations

Contextual Consideration	Description
This intervention is intended for the care of individuals with a condition of particularly high severity in terms of impact on length of life and/or quality of life. This intervention is intended for the care of individuals with a condition that represents a	For patients with frequent and severe migraine attacks that have not responded to other therapies or have had intolerable side effects or contraindications to their use, these new therapies may offer a new treatment option. For some individuals with migraine, it is a frequent, unpredictable and disabling condition that impacts all
particularly high lifetime burden of illness. This intervention is the first to offer any improvement for patients with this condition.	aspects of life. There are currently available over the counter and FDA approved medications for patients with migraine attacks.
Compared to "the comparator", there is significant uncertainty about the long-term risk of serious side effects of this intervention.	For patients who improve with lasmiditan, rimegepant or ubrogepant and have tolerable side effects, it is expected that prolonged use for migraine attacks will be recommended. Questions remain about the development of new side effects and the risk of medication overuse headaches with frequent use over time.
Compared to "the comparator", there is significant uncertainty about the magnitude or durability of the long-term benefits of this intervention.	For new medications that have mainly been evaluated in single dose comparative trials or non-comparative open- label studies of up to a year, their long-term benefits are uncertain relative to other therapies that have years of experience.
There are additional contextual considerations that should have an important role in judgments of the value of this intervention.	Lasmiditan, rimegepant and ubrogepant have not been shown to cause vasoconstriction, but whether they are free of cardiovascular adverse effects, particularly in those with cardiovascular disease or at high risk, remains to be proven.

Value-Based Price Benchmarks

Annual value-based price benchmarks (VBPBs) of these drugs (vs. usual care) are presented in Table ES13. The VBPB for a drug is defined as the price range that would achieve incremental costeffectiveness ratios between \$100,000 and \$150,000 per QALY gained. While the results of the NMAs suggest that lasmiditan may be slightly more efficacious than rimegepant and ubrogepant, they do not exclude comparable efficacy given the overlapping confidence intervals. Additionally, lasmiditan treatment results in more adverse events and is discontinued more frequently. Given that we felt the net benefits of the therapies were relatively similar as reflected in our comparative evidence ratings, we developed a range of value-based price benchmarks across all three drugs, using the range of threshold prices reported in Section 4.3 so as to avoid suggesting greater certainty in the individual threshold prices than is warranted.

For these drugs, price discounts of approximately 30% to 46% from the assumed list price would be required to reach the \$150,000 per QALY threshold price, range (Table ES13). Price discounts of approximately 39% to 51% from assumed list prices would be required to reach the \$100,000 per QALY threshold price range. Note that these discounts are from the assumed placeholder prices, and not from actual list prices, which are not yet known.

As there is no mortality effect in the model, cost per LY gained is not relevant, and the cost per evLYG is essentially the same as the cost per QALY gained. We therefore do not report VBPBs for these in the table below.

Table ES13. Value-Based Price Benchmark Ranges for Lasmiditan, Rimegepant, and Ubrogepantversus Usual Care in Population 1 (Patients Who Cannot Take Triptans)

	Assumed Annual Price*	Annual Price at \$100,000 Threshold	Annual Price at \$150,000 Threshold
Drug Price	\$4,515	\$2,200-\$2,770	\$2,440-\$3,150
Discount from Assumed Price*		39% to 51%	30% to 46%

QALY: quality-adjusted life year, WAC: wholesale acquisition cost

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

Potential Budget Impact

We used the cost-effectiveness model to estimate the potential total budgetary impact of each drug (lasmiditan, rimegepant, and ubrogepant) added to usual care for prevalent individuals in the United States (US) aged 18 years and over experiencing migraines requiring acute treatment, with or without aura. Potential budget impact was defined as the total differential cost of using each new therapy rather than relevant existing therapy for the treated population, calculated as differential health care costs (including drug costs) minus any offsets in these costs from averted health care events. We used the same assumed placeholder price and threshold prices for each drug in our estimates of budget impact, rather than using the three threshold prices calculated for each drug in Section 4.3. As mentioned above, while the results of the NMAs suggest that lasmiditan may be slightly more efficacious than rimegepant and ubrogepant, they do not exclude comparable efficacy given the overlapping confidence intervals and higher adverse event and discontinuation rates with lasmiditan. We therefore used a blended range of prices in our potential budget impact analyses, using the same \$50,000, \$100,000, and \$150,000 cost-effectiveness threshold price for each drug. From the threshold prices for the three drugs, we used the lowest price for the \$50,000 per QALY threshold (\$1,950) and for the \$100,000 threshold (\$2,200), and the

highest price for the \$150,000 per QALY threshold (\$3,150). We also included a scenario analysis where the frequency of migraines is assumed to decrease over time. All costs were undiscounted and estimated over a five-year time horizon, given the potential for cost offsets to accrue over time and to allow a more realistic impact on the number of patients treated with the new therapy.

This potential budget impact analysis does not include the population cohort of patients with migraines who are eligible for treatment with triptans, as sumatriptan and eletriptan dominated these drugs in our cost-effectiveness analysis. This potential budget impact analysis includes the cohort of patients who had migraine attacks that did not respond to non-prescription medicines and for whom triptans had not been effective, were not tolerated, or were contraindicated. Using data from the literature, we estimate the size of the potential candidate population for treatment in the average 2020-2024 estimated US adult population as approximately 6.4 million patients, or approximately 1.3 million patients each year over five years.

Base-Case Results

For lasmiditan, as shown in Figure ES1, approximately 12% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at the assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000. All eligible patients could be treated at the \$100,000 and \$50,000 threshold prices, with estimated potential budget impact of approximately 62% of the threshold at the \$100,000 threshold price.

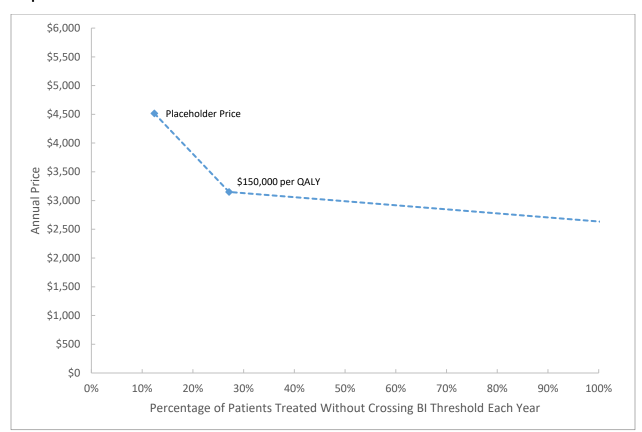
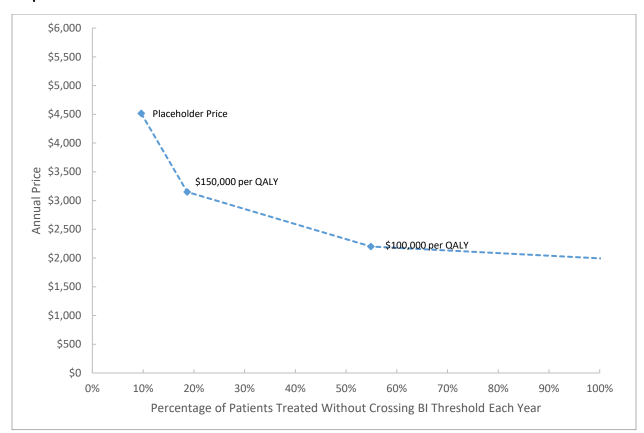
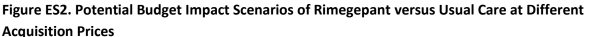


Figure ES1. Potential Budget Impact Scenarios of Lasmiditan versus Usual Care at Different Acquisition Prices

BI: budget impact, QALY: quality-adjusted life year

For rimegepant, as shown in Figure ES2, approximately 10% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at rimegepant's assumed placeholder price. Approximately 18% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price, increasing to approximately 55% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 91% of the threshold.





BI: budget impact, QALY: quality-adjusted life year

For ubrogepant, as shown in Figure ES3, approximately 10% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at ubrogepant's assumed placeholder price. Approximately 19% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price, increasing to approximately 55% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 91% of the threshold.

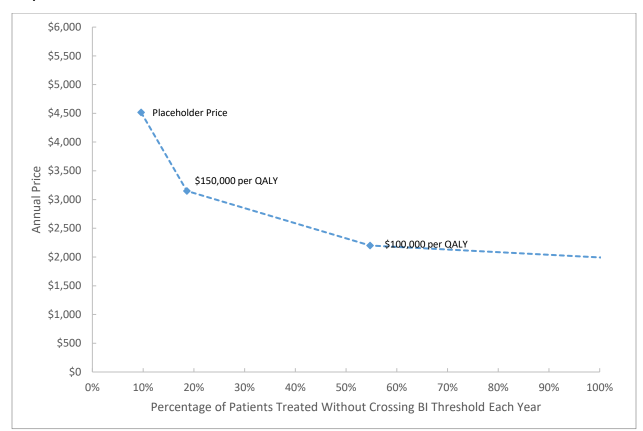


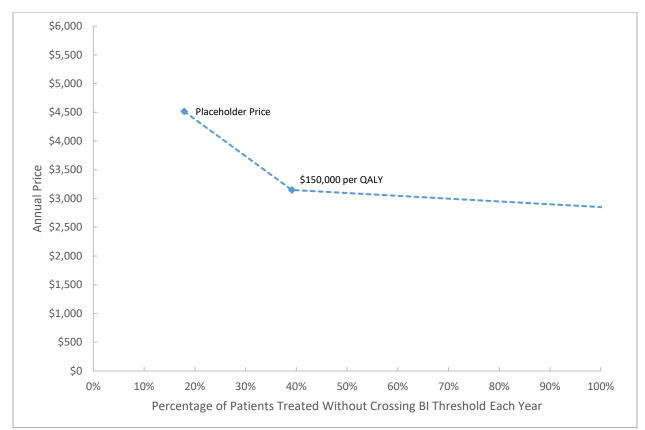
Figure ES3. Potential Budget Impact Scenarios of Ubrogepant versus Usual Care at Different Acquisition Prices

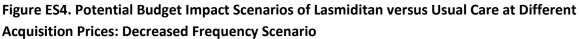
BI: budget impact, QALY: quality-adjusted life year

Scenario Results

Data from a long-term open label safety study suggests that the frequency of migraines decreased over time. While this single-arm trial was not designed to evaluate whether the same effect was observed in a control population, decreasing migraine frequency over time could have a significant impact on budget impact analyses. We therefore created a scenario analysis where we modeled the potential budget impact of these treatments if migraine frequency decreases over time.

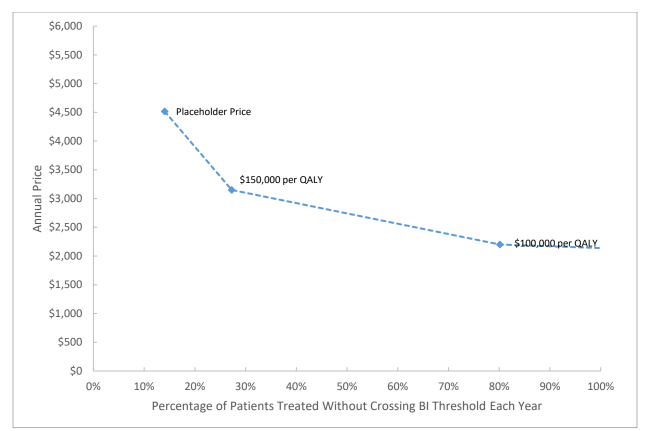
For lasmiditan in this scenario, as shown in Figure ES4, approximately 18% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at lasmiditan's assumed placeholder price. Approximately 39% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000. All eligible patients could be treated at the \$100,000 and \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 43% of the threshold at the \$100,000 threshold price

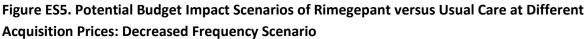




BI: budget impact, QALY: quality-adjusted life year

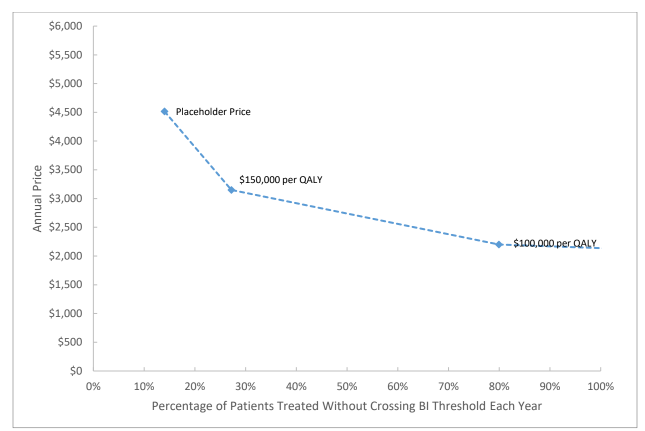
For rimegepant in this decreased frequency scenario, approximately 14% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at rimegepant's assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price, increasing to approximately 80% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 62% of the threshold.

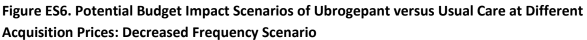




BI: budget impact, QALY: quality-adjusted life year

For ubrogepant in this scenario, approximately 14% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at ubrogepant's assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000, increasing to approximately 80% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 63% of the threshold.





BI: budget impact, QALY: quality-adjusted life year

1. Introduction

1.1 Background

Migraine is a common, typically episodic cause of disabling headache often associated with nausea and sensitivity to light and sound. Approximately 40 million adults (12-15%) in the United States (US) have reported migraine or severe headaches.^{2,3} The hallmark of migraine is recurrent attacks characterized by headache that is often but not always one-sided and described as pulsatile or throbbing. In addition to headache, other symptoms may start right before or occur with the headache including nausea with or without vomiting, and sensitivity to external stimuli such as light, sound, and smells. The frequency of attacks and the intensity of symptoms vary widely, but when frequent and severe, migraine can be a disabling, chronic condition that can impact all aspects of life including personal relationships and ability to work.¹ Patients with migraine have increased use of health care resources including visits to health care providers and emergency departments.^{4,5} Overall cost of health care for those with migraine are estimated to be \$11-50 billion dollars in the US.^{4,6} Direct health care costs as well as indirect costs associated with decreased productivity, work loss and disability claims are higher for those with migraine,⁷⁻⁹ and migraine is one of the most common causes of disability worldwide.⁵⁸

Diagnosis of migraine is based upon patient-reported symptoms, history, and physical examination findings; there is no test available that confirms the diagnosis.¹⁰ This may partly explain why many individuals with migraine may be incorrectly diagnosed.^{12 12} Clinical criteria broadly include the frequency and nature of the headache and the presence or absence of aura. Aura refers to a gradual onset of sensory or motor symptoms either before the onset of headache or as part of the headache. Though some patients do not have aura, the most common are visual symptoms such as seeing bright lines, shapes, or objects.¹² Headache features associated with a diagnosis of migraine include location on one side of the head, pulsating quality, moderate or severe pain intensity, and known triggers. Migraine is more common in women than men,¹³ and in those aged 18 to 44 years.^{2,3} A genetic predisposition to migraines is thought to account for their tendency to run in families. The precise cause of migraines is not known, but hypersensitivity of the brain to external stimuli and internal factors lead to activation of the trigeminovascular system of nerves that result in blood vessel and pain responses.¹¹ Predisposing factors associated with migraine attacks include emotional stress, menstruation, visual stimuli, changes in weather, and certain foods and activities.¹⁴

Treatment of migraine broadly focuses on two strategies: preventive therapy to reduce the frequency of attacks or acute therapy meant to quickly abort episodic symptoms, which is usually more effective the sooner it is given.¹² Acute treatments are referred to by a number of other terms including "abortive treatment," and "symptomatic treatment"; we will use the term "acute

treatment" in this document. Early acute treatment is especially helpful for individuals with aura that precedes the onset of the headache. The choice of therapy is based upon symptom frequency, severity, and the presence of nausea and vomiting. For individuals with mild symptoms, first-line over-the-counter nonspecific pain medications include aspirin, nonsteroidal anti-inflammatory drugs (NSAIDs) such as ibuprofen and naproxen, and acetaminophen. There are also combination preparations with caffeine, but caffeine withdrawal headaches can occur with frequent use. Other strategies such as lying down in a quiet and dark room are also helpful, and a nap or sleep sometimes lead to relief.

For individuals with moderate or severe symptoms or lack of response to nonspecific pain medications, the use of specific migraine medications is recommended. The most commonly used migraine specific medication class targets the 5-hydroxytryptamine (5-HT) or serotonin receptor. Seven 5-HT 1b/1d agonists or "triptans" are US Food and Drug Administration (FDA) approved for acute treatment of migraine attacks.¹¹ Triptans are available as pills, nasal sprays, and for injection under the skin, with non-oral routes of administration typically for those with severe headache accompanied by nausea and/or vomiting. Though effective and safe for many patients with migraine, triptans are labeled as contraindicated in patients with known cardiovascular disease because of their vasoconstrictive effects, but observational studies have not identified major cardiovascular risk as used in clinical practice.⁵⁹ Similarly, despite a reported possibility of serotonin syndrome in patients who combine triptans with selective serotonin and serotonin-norepinephrine reuptake inhibitors, the actual risk appears to be extremely low.^{60,61}

Ergotamine preparations also represent migraine-specific treatment, but side effects and limited efficacy have resulted in their being much less commonly used since the introduction of triptans. Non-specific pain medications, such as barbiturates and opioids, have similar limitations as well as the potential for tolerance and misuse, and have led to their being reserved for patients unresponsive to other therapies. For patients with associated nausea and vomiting, antiemetics are used but generally in addition to other medications. For most individuals with migraine, treatment focuses on episodic intervention. However, for the one-quarter to one-third of patients with severe and frequent attacks, medications to prevent migraine attacks are recommended.¹² This is important because medication overuse headache can result from frequent administration of acute medications for migraine attack, especially with nonspecific pain medications such as opioids, barbiturates, and combination agents. However, the prevalence of medication overuse headaches varies widely based upon differences in definitions and the population assessed.^{62,63}

Interventions: Calcitonin gene-related peptide (CGRP) antagonists (rimegepant, ubrogepant) and 5-hydroxytryptamine (5-HT) 1f agonist (lasmiditan)

Many individuals do not adequately respond to multiple different medications for acute treatment, demonstrating a need for new therapeutic options. For example, studies of triptans often

demonstrate response rates of 40-75%,¹⁵ and decreased response over time can also be seen in some individuals.¹⁶ One new target for therapy is calcitonin gene-related peptide (CGRP). Interest in agents that target CGRP is based upon it being expressed in trigeminal ganglia nerves involved in the vasodilatory component of neurogenic inflammation, and administration of CGRP can trigger acute headache and delayed migraine-like attacks.^{17,18} Injectable monoclonal antibodies targeting the CGRP receptor recently began being used for migraine prophylaxis, and there are two new oral CGRP receptor antagonists for acute treatment of migraine attacks: ubrogepant (Ubrelvy™, Allergan), approved on December 23, 2019 by the FDA, and rimegepant, under review by the FDA. ^{19,64} This new class of medications has been referred to as "gepants." Another new acute treatment for migraine is lasmiditan (Reyvow™, Lilly), a selective 5-HT 1f agonist (also referred to as a "ditan"), that was approved on October 11, 2019 by the FDA. Unlike triptans that cause vasoconstrictive effects on cranial and coronary blood vessels via the 5-HT 1b receptor, the gepants and lasmiditan have not been shown to cause vasoconstriction but maintain activity for acute treatment of migraine.^{16,20,21}

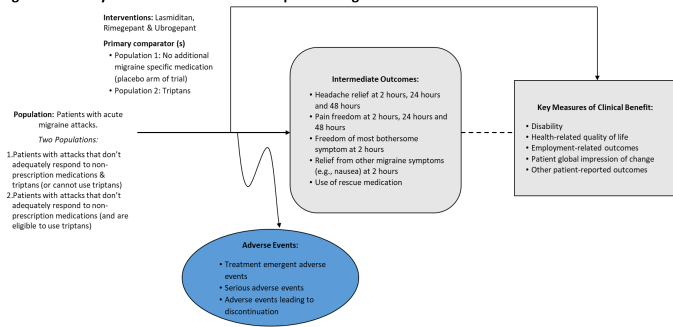
1.2 Scope of the Assessment

This review evaluates the comparative clinical effectiveness and economic impacts of lasmiditan, rimegepant, and ubrogepant for acute treatment of migraine. Evidence was collected from available randomized controlled trials, non-randomized clinical trials, comparative observational studies, as well as high-quality systematic reviews. We limited our review to those studies that captured the outcomes of interest. We included all randomized controlled trials (RCTs) and sought evidence on lasmiditan, rimegepant, and ubrogepant from non-randomized controlled trials and observational studies. We supplemented our review of published studies with data from conference proceedings, regulatory documents, information submitted by manufacturers, and other grey literature when the evidence met ICER standards (for more information, see https://icer-review.org/methodology/icers-methods/icer-value-assessment-framework-2/grey-literature-policy/). We sought head-to-head studies of lasmiditan, rimegepant, and ubrogepant and comparators to evaluate the feasibility of a network meta-analyses of selected outcomes.

Analytic Framework

The general analytic framework for assessment of acute therapies for migraine is depicted in Figure 1.1.





The diagram begins with the population of interest on the left. Actions, such as treatment, are depicted with solid arrows which link the population to outcomes. For example, a treatment may be associated with specific clinical or health outcomes. Outcomes are listed in the shaded boxes: those within the rounded boxes are intermediate outcomes (e.g., change in blood pressure), and those within the squared-off boxes are key measures of clinical benefit (e.g., health-related quality of life). The key measures of clinical benefit are linked to intermediate outcomes via a dashed line, as the relationship between these two types of outcomes may not always be validated. Curved arrows lead to the adverse events of an action (typically treatment), which are listed within the blue ellipsis.⁶⁵

Populations

The population of focus for this review was adults ages 18 years and older with a diagnosis of migraine, with or without aura as specified by the ICHD diagnostic criteria. We evaluated two populations of patients with migraine:

- Patients who have migraine attacks that have not adequately responded to nonprescription medicines and for whom triptans have not been effective, are not tolerated, or are contraindicated.
- 2. Patients who have migraine attacks that have not adequately responded to nonprescription medicines (and are eligible to use triptans).

For both populations, we also sought evidence on subgroups of interest, such as: a) patients considered to have chronic migraine (>15 headache days per month); b) patients currently receiving preventive migraine medication.

Interventions

The following new therapies were evaluated:

- Lasmiditan
- Rimegepant
- Ubrogepant

Comparators

For Population 1, we compared lasmiditan, rimegepant, and ubrogepant to each other and to no additional migraine-specific acute treatment. For the purpose of this review, no additional migraine-specific acute treatment was estimated by the placebo arms of the clinical trials, although we recognized that in the real-world patients may use failed over-the-counter analgesics including analgesics marketed as effective for acute treatment of migraine.

For Population 2, we compared lasmiditan, rimegepant, and ubrogepant to each other and to two triptans: sumatriptan and eletriptan. Sumatriptan was chosen because it is one of the most widely used triptans in clinical practice; and eletriptan, a newer triptan, was shown in a recent network meta-analysis to be one of the most efficacious and well tolerated.^{15,22} Since these new agents under review are all orally available, we focused our comparison of triptans on the oral formulations.

Outcomes

We looked for evidence on the following outcomes of interest.

Efficacy Outcomes:

- Headache relief at two hours
- Sustained headache relief (at 24 hours and 48 hours)
- Pain freedom at two hours
- Sustained pain freedom (at 24 and 48 hours)
- Freedom from most bothersome symptom (MBS) at two hours
- Relief from other migraine symptoms (e.g., photophobia, phonophobia, nausea, vomiting) at two hours
- Headache relief and pain freedom at 24 and 48 hours
- Patient global impression of change
- Use of rescue medication
- Disability
- Health-related quality of life

- Other patient-reported outcomes (e.g., depression, anxiety, and difficulties in interpersonal relationships)
- Employment-related outcomes (e.g., unemployment, work productivity loss, absenteeism)

Safety Outcomes:

- Serious adverse events
- Adverse events leading to discontinuation
- Treatment-emergent adverse events (e.g.)
 - o Dizziness
 - o Nausea
 - o Paresthesia
 - o Somnolence
- Medication overuse headache

Timing

Evidence on intervention effectiveness and safety was derived from studies of any duration, as long as they met the study design criteria set forth above and measure the outcomes of interest.

Settings

All relevant settings were considered, with a focus on outpatient settings in the United States.

1.3 Definitions

Clinical Outcome Measures

Outcomes of clinical trials of acute treatment of migraine commonly include relief of symptoms including pain, nausea/vomiting, photophobia and phonophobia. <u>Pain freedom</u> is defined as a reduction in severity of headache from mild, moderate or severe pain at baseline to none at a given follow-up time point. <u>Freedom from most bothersome symptoms (MBS)</u> refers to total absence of nausea/vomiting, photophonia or phonophobia at a given follow-up time point. <u>Pain relief</u> is defined as having mild to no pain at a given follow-up time point. The primary efficacy time point for phase 3 trials of lasmiditan, rimegepant and ubrogepant was at 2 hours after the first dose of the study drug. <u>Sustained symptom response</u> after 2-hours refers to those with an initial response that is sustained at subsequent follow-up time points without the use of repeat dosing or rescue medications. Censored outcomes after 2 hours that exclude those with repeat dosing or rescue medications are meant to maintain initial randomization to study drug or placebo but are less useful when estimating outcomes for an entire population at varying time points. As a result, uncensored outcomes after 2 hours were examined with the recognition that such outcomes may

include the benefit of rescue medications or simply the passage of time. Finally, even uncensored outcomes over time using Kaplan-Meier methods do not account for changes in symptoms after the initial outcome response.

Patients' Global Impression of Change (PGIC)

The PGIC is a seven-point scale reflecting patients' rating of overall improvement. It ranges from 1 ("very much worse") to 7 ("very much better").

Migraine Disability Assessment Test (MIDAS)

The Migraine Disability Assessment (MIDAS) is a brief, 7-item, self-administered questionnaire designed to quantify headache-related disability.⁶⁶ Respondents answer five questions about activity limitations in the past 3 months due to migraine including (1) missed work or school days, (2) missed household chores days, (3) missed non-work activity days, and days at work or school (4) plus days of household chores (5) where productivity was reduced by half or more. Two additional questions about the number of headaches and average pain level associated with headaches over the past 3 months are not used in deriving the MIDAS score, but they are for use by the respondent's clinician. The MIDAS score is the sum of the number of days reported for each of the five questions. Respondents with a MIDAS score of 0-5 are rated as having little or no disability, 6-10 as having mild disability, 11-20 as having moderate disability, and 21 or greater as having severe disability.

1.4 Insights Gained from Discussions with Patients and Patient Groups

In developing and executing this report, we received valuable input from individual patients and patient advocacy groups throughout the scoping and evidence development process. We received public comments on our draft scoping document from the following patient advocacy organizations: the Coalition for Headache And Migraine Patients (CHAMP), the Headache & Migraine Policy Forum, and the Institute for Patient Access. We also conducted scoping calls with the Alliance for Patient Access, American Headache Foundation, American Migraine Foundation, CHAMP, Golden Graine Blog, Headache & Migraine Policy Forum, Miles for Migraine, and the National Headache Foundation. Below we summarize the key insights derived from this input.

Patients with migraine describe different personal stories, but they identified common themes that emphasize migraine as an episodic and chronic disease that can profoundly affect all aspects of their lives and the lives of those close to them. Though some have derived benefit from existing therapies, not all respond and response to individual attacks can be variable. For others, side effects have led them to have to stop therapy. Patients also report recurrence of headaches as medications wear off during the acute episode or medication overuse headaches from frequent dosing for acute attacks. The net result is that for many patients with moderate or severe migraine headaches there is no single or combined therapy that offers them control of their acute attacks.

Patients and patient advocacy groups highlighted the deficiencies with currently available acute treatments for migraine. Despite a wide range of medications, both non-prescription and prescription, used alone or in combination, many patients are not able to reliably prevent or abort migraine attacks, either because therapies do not work, lose efficacy or have intolerable side effects. The result is that currently available therapies do not provide symptom relief from migraine attacks with minimal side effects for many individuals. Patients and advocacy groups noted that triptans represented a major advance in acute therapy for migraines when introduced over 20 years ago. However, many individuals cannot use triptans either because they do not work, have intolerable side effects such as flushing, numbness or chest pain, or have contraindications to their use such as existing cardiovascular disease. Because of limitations with triptans, patient often turn to other medications such as anti-emetics, barbiturates and opioids, but these also have limited benefit, acute side effects and important risks associated with long-term use.

A patient with episodic migraine describes her experience with available therapies in her public comments on the ICER draft evidence report: "I eagerly tried sumatriptan when it first hit the market in the 90s. I had a severe adverse reaction to it including severe tachycardia, shortness of breath, and my headache got much, much worse. Over the years I have tried various triptans again as new ones have hit the market or because my doctor wanted to rule them out again. I have always had the same reaction to the medications. DHE has not helped in years either. It used to work if I treated an attack when it was starting, but it no longer helps, and I often wake with a migraine attack already in progress anyway. For acute treatment, I've tried opiates and NSAIDS as well. Nothing helps and they actually seem to make things worse. For now, I do nothing to treat my attacks and it is no way to live. Some days I feel frantic for relief from the pain and other symptoms, but there is nowhere to turn. I am trapped with this. I desperately need access to new types of acute treatments."

The profound impact of migraine on the lives of patients with moderate and severe migraine was also emphasized. Migraine often develops in individuals during adolescence and young adulthood. Frequent, severe attacks can have a dramatic impact on quality of life that may not be fully appreciated by the general public and even health professionals. Stakeholders indicated that migraine attacks, especially when severe, recurrent and poorly controlled can be disabling. One patient commented: "Two years after being diagnosed with chronic, intractable migraine, I had to stop working in a career that I truly loved and for a company that was incredibly supportive of my illness. I also was in my second year of grad school at Georgetown University. The migraine thief took all of that away from me." As mentioned in this patient's story, when migraine attacks occur during formative educational years, it can prevent individuals from reaching their full academic potential. Patients also highlighted that the unpredictability of migraine attacks can result in anxiety from not knowing when the next attack will come, thus affecting individuals even when

they do not have migraine symptoms. The net effect is that migraine is an episodic and chronic condition that affects patients throughout their lives, disrupting personal relationships with friends and family, and their ability to work. The toll on patients with migraine also includes important economic consequences. For many individuals with migraine, attack severity disrupts daily life, often unpredictably. If the migraine attack is not aborted quickly and without medication related side effects, ability to work or work productively is profoundly affected. The combination of frequent, severe and unpredictable migraine attacks impacts ability to work, increases the risk of disability, and can have a long-term negative economic impact on the patient and her/his family. Patients and patient advocates recognize the critical importance of acute treatments for migraine that work quickly and without side effects on the ability to continue to work on the day of a migraine attack. Whether patients cannot work at all, work intermittently or part-time, or were less productive at work because of symptoms of migraine or side effects of therapies, the net result can be long-term un/under-employment with major socioeconomic costs.

Patients and advocates emphasized that because many patients do not find triptans effective or have side effects or contraindications to their use, doctors end up prescribing barbiturates and opioids. Though recognized as having limited effectiveness, acute side effects, the potential for causing medication overuse headaches, and a misuse potential, desperate patients frequently end up being prescribed these medications (for a small percentage of patients with difficult to treat migraine, barbiturates and opioids may be appropriate). The importance of new therapeutic classes, especially ones without side effects or limitations to use as seen with triptans, is important for managing patients with migraine attacks and may also have a broader potential impact on the opioid crisis in the US.

Finally, patient advocacy organizations also raised systematic issues that they felt needed to be addressed. They highlighted that common outcome measures required by the FDA to obtain approval for new drugs may not adequately capture the impact of migraine on things that affect the overall quality of life of migraine patients including relationships, work, and family issues. For example, outcomes of single dose efficacy studies are not designed to assess whether new therapies can decrease the frequency of migraine attacks over time or prevent medication overuse headaches. They felt this to be particularly important for patients with frequent and severe migraine attacks who have not responded to, are intolerant of, or unable to take triptans. Moreover, patients with migraine may have other illnesses, such as anxiety and depression, that are impacted by frequent, unpredictable and severe migraine symptoms. Successful treatment of migraine attacks may also help with these other conditions.

1.5. Potential Cost-Saving Measures in Migraine

ICER includes in its reports information on wasteful or lower-value services in the same clinical area that could be reduced or eliminated to create headroom in health care budgets for higher-value innovative services (for more information, see https://icer-review.org/final-vaf-2017-2019/). These services are ones that would not be directly affected by therapies for migraine (e.g., reduction in ED visits), as these services will be captured in the economic model. Rather, we are seeking services used in the current management of migraine beyond the potential offsets that arise from a new intervention. During stakeholder engagement and public comment periods, ICER encouraged all stakeholders to suggest services (including treatments and mechanisms of care) currently used for patients with migraine that could be reduced, eliminated, or made more efficient.

For this review, we received one such suggestion: Allergan and some patient groups noted that opioids for acute treatment of migraines are discouraged by guidelines and yet remain overused. Allergan suggested that opioids represent a low-value service that could be reduced.

2. Summary of Coverage Policies and Clinical Guidelines

2.1 Coverage Policies

To understand the insurance landscape for acute treatments of migraine relevant to this review, we reviewed National and Local Coverage Determinations (NCDs and LCDs) from the Centers for Medicare and Medicaid Services (CMS), and publicly available coverage policies from representative national plans (Aetna and Cigna), national and regional private payers (HealthPartners and Blue Cross Blue Shield of Kansas City) and state Medicaid plans (MO Healthnet and IL Health and Family Services). We surveyed the coverage policies for lasmiditan, rimegepant, ubrogepant, and oral triptans (with special focus on sumatriptan and eletriptan). No coverage policies, nor any NCDs or LCDs, for lasmiditan and oral CGRP antagonists rimegepant and ubrogepant were yet available at the time of this report. The FDA recently approved lasmiditan on October 11, 2019 for acute treatment of migraine. Approval is pending for rimegepant and ubrogepant.

On the national level, generic sumatriptan and eletriptan tablets are on the preferred drug list as step 1, tier 2 or high cost generic formulary without prior authorization, however quantity limits apply (between 9 and 12 tablets per month). Brand name versions are typically non-preferred and require prior authorization^{67,68} or are step 2.⁶⁹

2.2 Clinical Guidelines

American Headache Society (AHS)

The American Headache Society (AHS) 2015 guideline for acute treatment of migraine labeled several medications as Level A (established as effective for acute migraines based on available evidence): almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, sumatriptan (oral, nasal spray, patch and subcutaneous), zolmitriptan (oral and nasal spray), acetaminophen, ergots, NSAIDS, butorphanol nasal spray and acetaminophen/aspirin/caffeine and sumatriptan/naproxen combination therapies.⁷⁰ The society acknowledged that there are many acute migraine treatments with strong evidence to support their efficacy, but that clinicians should also consider potential side effects and adverse events when prescribing medications for acute migraine. Further the society indicated that opioids, such as butorphanol, codeine and tramadol, though probably effective, are not recommended for regular use.

American Academy of Family Physicians (AAFP)

The American Academy of Family Physicians (AAFP) issued a 2012 guideline on the acute treatment of migraines in the emergency setting.⁷¹ They concluded there is moderate evidence to support the use of neuroleptics, NSAIDS and injectable sumatriptan for the ability to achieve pain-free status in 1-2 hours, moderate evidence to support neuroleptics and injectable sumatriptan for the ability to provide headache relief at 1-2 hours, and moderate evidence to support the use of neuroleptics, metoclopramide, opioids and injectable sumatriptan for the ability to reduce pain intensity.

American Academy of Neurology (AAN) - Choosing Wisely

In 2013 the American Academy of Neurology (AAN) and Choosing Wisely issued a joint statement recommending that the use of opioids or butalbital for acute treatment of migraine be avoided except as a last resort because other more effective treatments are available and frequent use can worsen headache. Opioids should be reserved only for those patients who fail other treatments or cannot take migraine-specific treatments.⁷²

Canadian Headache Society (CHS)

A 2013 Canadian Headache Society (CHS) guideline gave twelve medications a strong recommendation for use in acute migraine: almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, sumatriptan, zolmitriptan, aspirin, ibuprofen, naproxen sodium, diclofenac potassium and acetaminophen. Four received a weak recommendation: dihydroergotamine, ergotamine, codeine-containing combination analgesics and tramadol-containing combination analgesics.⁷³ Ergotamine, butorphanol, codeine, butalbital and tramadol-containing medications were not recommended or were strongly recommended against. The society acknowledged that several trials of acute treatments might be required before finding the right approach for a specific patient

and that a rescue plan should be in place if acute treatment is insufficient. The society recommends triptans for the acute treatment of migraine attacks that are likely to become moderate or severe and if a patient does not respond well to one triptan or tolerates it poorly, other triptans should be tried (after 24 hours). If response to sumatriptan is inadequate, the society suggests considering adding an NSAID simultaneously with the triptan. Finally, patients with moderate to severe migraine attacks should take triptans as early in the attack as possible.

Canadian Authority for Drugs and Technologies in Health (CADTH)

In a 2012 systematic review of the safety of triptans, the Canadian Authority for Drugs and Technologies in Health (CADTH) found no consistent differences in the occurrence of adverse events (AEs) between triptans, although a dose-response relationship for oral sumatriptan was observed.⁷⁴ AEs for sumatriptan include dizziness, drowsiness, paresthesia, nausea and fatigue, but are generally mild and self-limiting. Overall incidence of withdrawal due to AEs for all doses of sumatriptan was 1.6% compared to 0.68% for placebo.

A 2007 CADTH review assessed the cost effectiveness of triptans for acute treatment of migraines. They found no evidence that one triptan was more effective than another and concluded that more research is needed to establish differences in benefits and harms between triptans.⁷⁵ The costeffectiveness studies included in the review mostly only included drug costs, making them difficult to interpret from a broader system or societal perspective.

National Institute for Health and Care Excellence (NICE)

We reviewed clinical guidelines for migraine from the National Institute for Health and Care Excellence (NICE), last updated in 2015.⁷⁶ For acute treatment of migraine, NICE recommends oral triptans in combination with NSAIDs, aspirin or paracetamol. NICE suggests starting with the lowest cost triptan, followed by other triptans if treatment is ineffective. Furthermore, NICE recommends an anti-emetic drug in addition to acute treatment, even in the absence of nausea but recommends against non-migraine specific pain medications such as ergots or opioids.

NICE currently has three reviews of injectable CGRP antagonists for preventing migraine: erenumab (publication TBD), fremanezumab (April 15 2020) and galcanezumab (publication TBD).⁷⁷⁻⁷⁹ Preliminary recommendations from NICE state that erenumab is not a recommended first-line treatment for preventing migraines.⁸⁰ If a patient does not respond to beta-blockers, antidepressants, and anti-epileptics, another oral preventive drug or Botox should be offered first. Erenumab is an option when at least three treatments have failed to prevent migraine.

3.1 Overview

To inform our review of the comparative clinical effectiveness of lasmiditan, rimegepant, and ubrogepant for acute treatment of migraine, we systematically identified and synthesized the existing evidence from available clinical studies. Full PICOTS criteria were described in Section 1.2. In brief, we compared the efficacy, safety, and effectiveness of lasmiditan, rimegepant, and ubrogepant to each other. In addition, we compared all three interventions to no additional migraine-specific acute treatment (placebo) and triptans (sumatriptan and eletriptan). Our review focused on clinical benefits, as well as potential harms. We sought evidence on all outcomes listed in Section 1.2. Methods and findings of our review of the clinical evidence are described in the sections that follow.

3.2 Methods

Data Sources and Searches

Procedures for the systematic literature review assessing the evidence on lasmiditan, rimegepant, and ubrogepant for acute treatment of migraine followed established best methods.^{81,82} The review was conducted in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines.⁸³ The PRISMA guidelines include a list of 27 checklist items, which are listed in Appendix Table A1.

We searched MEDLINE, EMBASE, Cochrane Database of Systematic Reviews, and Cochrane Central Register of Controlled Trials for relevant studies. Each search was limited to English language studies of human subjects and excluded articles indexed as guidelines, letters, editorials, narrative reviews, case reports, or news items. We included abstracts from conference proceedings identified from the systematic literature search. All search strategies were generated utilizing the Population, Intervention, and Study Design elements described in Section 1.2.

We identified a recent systematic review and network meta-analysis of triptans which followed a similar scope to the one planned for this review, with literature search end date of 2016.²² RCTs of sumatriptan and eletriptan that met our criteria from the systematic review were identified. In addition, we searched for new evidence on sumatriptan and eletriptan that has emerged since 2016 by conducting an updated systematic literature search. However, we conducted a de novo search for lasmiditan, rimegepant, and ubrogepant. The search strategies included a combination of indexing terms (MeSH terms in MEDLINE and EMTREE terms in EMBASE), as well as free-text terms, and are presented in Appendix Tables A2 – A5. The date of the most recent search is August 21, 2019.

To supplement the database searches, we performed manual checks of the reference lists of included trials and recent systematic reviews of the intervention and individual comparators and invited key stakeholders to share references germane to the scope of this project. We also supplemented our review of published studies with data from conference proceedings, regulatory documents, information submitted by manufacturers, and other grey literature when the evidence meets ICER standards (for more information, see https://icer-review.org/methodology/icers-methods/icer-value-assessment-framework-2/grey-literature-policy/).

Study Selection

After removal of duplicate citations, references went through two levels of screening at both the abstract and full-text levels. Three reviewers independently screened the titles and abstracts of all publications identified using DistillerSR (Evidence Partners, Ottawa, Canada) and disagreements were resolved through consensus.

Studies that did not meet the PICOTS criteria defined above, were excluded. No study was excluded at abstract level screening due to insufficient information. Citations accepted during abstract-level screening were reviewed as full text. Reasons for exclusion were categorized according to the PICOTS elements.

Data Extraction and Quality Assessment

Two reviewers extracted data from the full set of included studies into an excel spreadsheet. Extracted data were independently verified by another researcher. Data elements included a description of patient populations, sample size, duration of follow-up, study design features (e.g., RCT or open label), interventions (drug, dosage), outcome assessments (e.g., timing and definitions), results, and quality assessment for each study. We used criteria employed by the US Preventive Services Task Force (USPSTF) that included presence of comparable groups, non-differential loss to follow-up, use of blinding, clear definition of interventions and outcomes, and appropriate handling of missing data to assess the quality of clinical trials and classify into categories "good," "fair," or "poor."⁸⁴ For more information on data extraction and quality assessment, refer to Appendix D.

Assessment of Level of Certainty in Evidence

We used the <u>ICER Evidence Rating Matrix</u> to evaluate the level of certainty in the available evidence of a net health benefit among each of the interventions of focus (see Appendix D).⁸⁵

Assessment of Bias

As part of our quality assessment, we evaluated the evidence base for the presence of potential publication bias. Given the emerging nature of the evidence base for newer treatments, we performed an assessment of publication bias for "lasmiditan", "rimegepant", and "ubrogepant" using the <u>ClinicalTrials.gov</u> database of trials. We scanned the site to identify studies completed more than two years ago that would have met our inclusion criteria and for which no findings have been published. Any such studies may indicate whether there is bias in the published literature. For this review, we did not find evidence of any study completed more than two years ago that has not subsequently been published.

Data Synthesis and Statistical Analyses

Data on outcome results were abstracted in evidence tables (see Appendix Tables D1-D14) and synthesized quantitatively and qualitatively in the body of the review. Data from OLEs and studies were described narratively only and not included in the quantitative syntheses. Using the available trial data, we conducted network meta-analyses (NMAs) for each outcome of interest when data existed on all the interventions of interest from at least three trials that were sufficiently similar in population, interventions, outcome definition, time point, and other characteristics. Based in part on availability of data from sufficiently similar trials, we conducted NMAs on the following outcomes: pain freedom, pain relief, freedom from the most bothersome symptom, disability, adverse events, and treatment-emergent adverse events. For the NMA, we used the 2- and 24-hour timepoints as available in each of the studies that reported on these outcomes. Due to inconsistent or limited reporting of data across studies, freedom from other migraine symptoms, use of rescue medication and patient global impression of change are described only in a narrative fashion.

All NMAs were conducted in a Bayesian framework with random effects on the treatment parameters using the *gemtc* package in R.⁸⁶ The outcomes were all binary and were analysed using a binomial likelihood and logit link.⁸⁷ We conducted network meta-regression to adjust for differences in placebo group response rate in the NMAs. Goodness of fit of the analyses with and without adjustment for differences in placebo arm response were assessed, and we present the results of the adjusted NMA model where it provided a better fit of the data. Tabular results below were presented for the treatment effects (odds ratio [OR]) of each intervention versus placebo along with 95% credible intervals (95% Crl). The expected proportion of patients experiencing the outcome were also presented when anchoring to the average placebo effect observed across the trials. Additional details regarding the analysis methods, network diagrams, as well as the results of unadjusted NMAs are provided in Appendix D.

3.3 Results

Study Selection

Our literature search identified a total of 323 potentially relevant references (see Appendix A Figure A1). We included 40 references, of which 37 references were on comparative clinical trials and three were open label extension studies (OLEs). These references consisted of 31 publications and nine conference abstracts. Primary reasons for study exclusion included use of interventions or comparators outside of our scope (e.g., subcutaneous sumatriptan), wrong study population (e.g., pediatric population), and conference abstracts with duplicate data as the full-text publications. In addition, because the trials of lasmiditan, rimegepant and ubrogepant included patients with moderate to severe acute migraine, we excluded studies of triptans that evaluated only mild cases of acute migraine.

The 37 references of comparative trials correspond to 33 trials, of which 10 trials (15 references) assessed lasmiditan or the CGRP antagonists, and 23 trials (22 references) assessed one or more of the comparators of interest. We identified only one head-to-head trial of one of the interventions versus a comparator of interest (rimegepant vs sumatriptan). Below, we describe the trials and efficacy results, followed by a discussion of the tolerability and harms.

Quality of Individual Studies

We highlighted the information on the quality of all trials (published and unpublished) using criteria from the U.S. Preventive Services Task Force (USPSTF) in Appendix Table D4. The trials of lasmiditan, rimegepant and ubrogepant had comparable arms at baseline, did not have differential attrition, were patient and physician/investigator blinded, had clear definitions of intervention and outcomes, and used an intent-to-treat analysis or a modified version. As such, we rated all three lasmiditan trials, the three published rimegepant trials, and all three ubrogepant trials to be of good quality. We did not assign an overall quality rating to the unpublished rimegepant trial (Study 301) obtained from grey literature sources (i.e. conference proceedings).

The triptan trials had ratings of good (19 trials) or fair (4 trials). Reasons for lower ratings include a lack of clear reporting on the comparability of the arms at baseline or the use of per-protocol as the primary method of analysis. Detailed information on the ratings can be found in Appendix Tables D4.

Overview of Studies

Lasmiditan, Rimegepant and Ubrogepant versus No Additional Migraine-Specific Acute Treatment (Placebo-controlled studies)

We identified three RCTs of lasmiditan (1 Phase II and 2 Phase III),²³⁻²⁵ four RCTs of rimegepant (1 Phase II and 3 Phase III),²⁶⁻²⁹ and three RCTs of ubrogepant (1 Phase II and 2 Phase III)^{30,31 32} versus placebo. Currently, one of the Phase III trials of rimegepant is unpublished and data for this study was obtained from conference abstracts.

All the identified studies were large multicenter studies, conducted predominantly in the United States, and were all focused on the treatment of a single-migraine attack. The trials enrolled patients who had at least a one-year history of migraine with or without aura as specified by the International Classification of Headache Disorders (ICHD) diagnostic criteria, who experienced two to eight migraine attacks of moderate to severe intensity per month, with age of onset before 50 years. Patients who met the eligibility criteria were randomized to intervention or placebo group and were asked to treat a single migraine attack of moderate or severe intensity within a maximum of four hours of onset. Patients and investigators were blinded to treatment assignment. Patients used an electronic diary to record their baseline migraine severity, other migraine-associated symptoms (e.g., photophobia, nausea, phonophobia), and response at different time intervals after taking the study drug over a 48-hour period. The trials reported results based on modified intention to treat populations, eliminating patients who did not experience a moderate to severe migraine event during the study period, so the number of participants included in the effect estimates for the outcomes in each trial were often less than the number of patients randomized.

All trials provided for the use of additional, rescue treatment for patients not responding to the initial study drug or having recurrent symptoms after initial benefit, but there were differences in the rescue treatments permitted and their timing and combinations. The lasmiditan and ubrogepant trials permitted the use of an optional second dose (randomized in the lasmiditan trials and open label in the ubrogepant trials). In terms of rescue medications allowed, the ubrogepant trials permitted patients to take their usual acute care treatment (including triptans and ergots), while the lasmiditan and rimegepant trials only allowed the use of non-specific migraine medication such as NSAIDS. The use of other medications was permitted between two and 24 hours after initial dosing in the lasmiditan trials and between two and 48 hours after initial dosing in the ubrogepant trials, if needed.

Appendix Tables D1 and D2 contains the key study design and baseline characteristics of each RCT. A summary is presented in Table 3.1. Over 80% of the patients were female and the average age was approximately 40 years in each trial. Patients had been living with migraine for approximately 20 years, had an average of three to five migraine attacks per month, and about 20% to 25% of patients in the trials were on preventive migraine medication. Characteristics of the treated migraine attack were generally similar across trials, with a distribution of approximately 30% and 70% for severe and moderate headache pain intensity, respectively. Photophobia was the most common other symptom reported (75% to 90% of patients) and was reported as the most bothersome symptom by 50% to 60% of patients. Approximately 40% to 65% of patients reported nausea, and 55% to 75% of patients reported phonophobia.

All trials excluded patients who had more than 15 days of headache per month, and patients who had clinically significant, unstable or recently diagnosed cardiovascular disease (e.g., coronary artery disease, uncontrolled hypertension) were excluded. Patients who initiated or changed preventative medication within 3 months were excluded from the lasmiditan trials.

The primary efficacy endpoint in all trials was freedom from pain at two hours after treatment, before the use of any rescue medication. Pain intensity was measured on a four-point Likert scale (0=none, 1=mild, 2=moderate. 3=severe). Most trials assessed freedom from the most bothersome symptom associated with migraine (MBS) (i.e. phonophobia, photophobia or nausea) at two hours as a co-primary endpoint. MBS was measured using a binary scale (0=absent, 1=present). The main secondary efficacy endpoints assessed in the trials included: 1) those assessed at two hours: headache pain relief (defined as reduction in pain severity from moderate or severe to mild or none), photophobia, phonophobia, nausea, ability to function normally, 2) those assessed at 24 and 48 hours: sustained freedom from pain, sustained freedom from MBS, and sustained pain relief. Sustained response was in those with a response at 2 hours who did not experience subsequent recurrence or use of rescue medications.

Lasmiditan, Rimegepant and Ubrogepant versus Triptans (Sumatriptan and Eletriptan)

We identified one placebo-controlled Phase II trial of rimegepant that included sumatriptan as an active control arm .²⁹ However, the trial did not report any statistical comparison between rimegepant and sumatriptan. We did not identify any trials comparing lasmiditan or ubrogepant to a triptan. As such, our assessment of these interventions versus triptans (sumatriptan and eletriptan) is informed by indirect comparisons (i.e. network meta-analysis). In all, we included 33 trials (23 triptan RCTs and 10 RCTts of the interventions including the Phase II trial of rimegepant with an active sumatriptan arm) to inform the indirect comparison. The 23 triptan RCTs ³³⁻⁵⁴ had comparable baseline characteristics to the other trials of the interventions described above. Of the 23 triptan studies, 18 were placebo-controlled trials of sumatriptan, three were placebo-controlled trials of eletriptan and two were head-to-head trials of sumatriptan and eletriptan with placebo arms.

As with the lasmiditan and the CGRP receptor antagonist trials, the majority of the included triptan studies were large multicenter studies, conducted in a variety of countries around the world and were focused on the treatment of a single-migraine attack. However, we included one trial that evaluated multiple migraine attacks (Pfaffenrath 1998) because it presented data on the first

migraine attack separately.⁴⁵ Patients and investigators were blinded to treatment assignment, and most of the trials permitted the use of rescue medication between 2 and 24 hours after initial dosing, if needed. The studies included patients who met the ICHD diagnostic criteria and had inclusion and exclusion criteria sufficiently comparable to the trials of lasmiditan and CGRP antagonist. The majority of studies included patients with a history of one to six migraine attacks of moderate to severe intensity per month. Most trials excluded patients with cardiovascular disease (e.g., cardiac ischemia, atherosclerosis, cardiac arrhythmia or uncontrolled hypertension).

Similar to the lasmiditan and CGRP antagonist trials, the majority of patients were female, the average age was approximately 40 years in each trial, and patients had been living with migraine for approximately 20 years. Patients in the eletriptan studies had an average of three to eight migraine attacks per month. Patients in the sumatriptan studies reported a range of one to eight attacks per month. Where reported, the distribution of treated migraine ranged from approximately 30% to 70% for severe headache pain intensity. Appendix Tables D1 and D2 contain the baseline characteristics of all the included triptan studies. A summary is presented in Table 3.1.

21 triptan trials evaluated pain relief at two hours post dose. Sixteen triptan trials reported freedom from pain at two hours post dose. None of the trials assessed freedom from the most bothersome symptom as an outcome. Other secondary outcomes evaluated in the triptan studies include sustained freedom from pain at 24 hours (6 trials) and sustained pain relief at 24 hours (10 trials).

Table 3.1: Overview of the Randomized Controlled Trials

Drug	Trials	N	Characteris	tics of Attacks
			Pain Intensity	Baseline Symptoms
Lasmiditan vs. Placebo	3 trials: SAMURAI SPARTAN Farkkila 2012	4, 291	Severe: 30 – 40% About 1-4% mild attacks and the remaining were moderate pain intensity attacks.	Nausea: 40 -65% Phonophobia: 60 -65% Photophobia: 75 -80%
Rimegepant vs. Placebo	4 trials: Study 301 Study 302 Study 303 Marcus 2014*	3, 869	Severe & Moderate: 100% (distribution not reported). No mild intensity attacks.	Nausea: 60% Phonophobia: 70% Photophobia: 80 - 90%
Ubrogepant vs. Placebo	3 trials: ACHIEVE I ACHIEVE II Voss 2016	3,105	Severe: 30 – 40% The remaining were moderate pain intensity attacks. No mild intensity attacks.	Nausea: 55% Phonophobia: 75% Photophobia: 90%
	Triptan	studies includ	ed in the NMA	
Sumatriptan vs. Placebo	18 trials	8,489	In 11 trials Severe: 30 – 70% Two trials included 5% to 10% mild intensity attacks. The remaining were moderate pain intensity attacks In 7 trials, Severe & moderate: 100% (distribution not reported).	Nausea: 50 – 70% Phonophobia: 70-75% Photophobia: 80-90%
Eletriptan vs. Placebo	3 trials	1,085	Severe: 50% The remaining were moderate pain intensity attacks. No mild intensity attacks.	Nausea: 50 – 65% Phonophobia: 70% Photophobia: 75-80%
Eletriptan vs. Sumatriptan	2 trials [†]	2,479	Severe: 40-45% The remaining were moderate pain intensity attacks. No mild intensity attacks.	Nausea: 50-65% Phonophobia: 65% Photophobia: 75%

N: total number of participants, NMA: network meta-analysis, vs.: versus

*Marcus 2014 includes an active comparator arm (sumatriptan)

+Includes a placebo comparator arm

Long-Term Studies of Lasmiditan, Rimegepant and Ubrogepant

We identified three ongoing 12-month open label extension studies (OLEs) of repeated use of acute medication for migraine over the study period, one on each intervention of interest. In the lasmiditan OLE study (GLADIATOR), interested patients who had completed either of the two-single attack Phase III RCTs with lasmiditan were randomized to receive either 100 mg lasmiditan or 200 mg lasmiditan.⁵⁶ Similar to the RCTs, patients enrolled in GLADIATOR were asked to treat moderate or severe attacks and were allowed to use a second dose of the medication after two hours. The rimegepant long term OLE study (Lipton 2019) evaluated the use of once daily rimegepant taken as needed (PRN) versus scheduled dosing (every other day) plus as needed use.⁸⁸ In the ubrogepant OLE (Ailani 2019), patients who had completed the two Phase III RCTs of ubrogepant were rerandomized to receive usual care or one of two doses of ubrogepant (50 mg or 100 mg).⁸⁹ Patients were instructed to treat up to eight attacks of any severity every four weeks and could use a second dose of the medication for non-response or recurrence. The trials primarily assessed the long-term safety and tolerability of the interventions. In addition, efficacy outcomes related to potential preventive effects of these medications (e.g., reduction in migraine days per month) were also

Clinical Benefits

As described in Section 1.2 of this report, we sought evidence on the following intermediate outcomes: pain freedom, freedom from most bothersome symptom (i.e. phonophobia, photophobia, and nausea), headache relief, and use of rescue medication. We found data to on all the intermediate outcomes for the three interventions of interest. We also sought evidence on the key measures of clinical benefit including disability, health-related quality of life, employment-related outcomes, and other patient reported outcomes. We found data on disability and patient reported global impression of change but did not find any data on the other outcomes. In addition, we also describe the available evidence on reduction in migraine days per month available in the identified trials, although we did not perform a systematic review specifically to evaluate this outcome.

For the interventions that evaluated more than one dose in the clinical trials (lasmiditan and ubrogepant), we describe the results observed in all arms of the trials. However, for the purpose of the NMAs, we pooled the two highest doses into one i.e. 100 mg and 200 mg arms of the lasmiditan trials were pooled into one arm (lasmiditan 100/200 mg), and 50 mg and 100 mg arms of the ubrogepant trials were pooled into one arm (ubrogepant 50/100 mg). The lower doses (50 mg lasmiditan and 25 mg ubrogepant) were not included in the NMA because these doses were not consistently evaluated in the Phase III trials and were not included in the long-term open label extension studies.

Freedom from Pain at Two Hours

This was defined as the presence of no pain at two hours after treatment in a person who had mild, moderate or severe pain and before the use of any rescue medication. In the individual Phase III clinical trials of the interventions presented in Table 3.2, lasmiditan (50 mg, 100 mg, or 200 mg), rimegepant (75 mg) and ubrogepant (25 mg, 50 mg or 100 mg) all resulted in a greater proportion of patients being free from pain at two hours post dose compared with patients receiving placebo (Table 3.2). A similar pattern was observed in the Phase II studies of the interventions and the triptan studies.

In total, 26 trials (3 lasmiditan trials,²³⁻²⁵ 4 rimegepant trials including 1 trial that included sumatriptan as an active comparator arm,²⁶⁻²⁹ 3 ubrogepant trials,³⁰⁻³² and 16 triptan studies^{33-38,42-44,46-48,50-52}) reported on the proportion of patients with pain freedom at two hours. We considered all 26 trials sufficiently similar to include in the NMA. Appendix Table D5 provides the data for the NMA, including the sample size and the number of patients who reported pain freedom.

The NMA model that adjusted for placebo response provided a better fit and the results are presented in Table 3.3 and Table 3.5 (unadjusted NMA results are provided in Appendix D). The results are presented in terms of the odds ratio (OR) of freedom from pain for each intervention versus placebo, sumatriptan and eletriptan. ORs above 1 indicate higher odds of pain freedom at two hours with the active intervention versus comparator while ORs below 1 indicate lower odds. Lasmiditan (OR: 3.01; 95% CrI: 2.2 to 4.14), rimegepant (OR: 2.11; 95% CrI: 1.67 to 2.72), and ubrogepant (OR: 2.12; 95% CrI: 1.58 to 2.88) all had higher odds of achieving pain freedom at two hours versus placebo. Compared to each other, none of the interventions showed statistically significant differences, though lasmiditan showed a statistically non-significant, higher odds of achieving pain freedom. In contrast, all interventions showed lower odds of achieving pain freedom at two hours compared to sumatriptan (lasmiditan: 0.73, rimegepant: 0.51, ubrogepant: 0.52) and eletriptan (lasmiditan: 0.54, rimegepant: 0.38, ubrogepant: 0.38). Of note, statistical significance was not reached for lasmiditan versus sumatriptan.

Based on the estimated odds ratios, the expected proportion of patients achieving pain freedom at two hours was 28% for lasmiditan, 21% for rimegepant, 21% for ubrogepant, 35% for sumatriptan and 42% for eletriptan (Table 3.5).

Pain Relief at Two Hours

Pain relief was defined as a decrease in headache pain from moderate or severe at baseline to mild or no pain at two hours after treatment and before taking any rescue medication. Patients with moderate or severe pain who achieve pain freedom would also be counted as having pain relief. In the individual Phase III clinical trials of the interventions, lasmiditan (50 mg, 100 mg, or 200 mg), rimegepant (75 mg) and ubrogepant (25 mg, 50 mg or 100 mg) all resulted in a greater proportion of patients experiencing pain relief at two hours post dose compared with patients on placebo (Table 3.2).

We included 31 trials in the NMA (3 lasmiditan trials,²³⁻²⁵ 4 rimegepant trials including 1 trial that included sumatriptan as an active comparator arm,²⁶⁻²⁹ 3 ubrogepant trials,³⁰⁻³² and 21 triptan studies³³⁻⁵²). Appendix Table D5 provides the trial data included in the NMA, which are the sample size and the number of patients who reported pain relief.

The NMA model adjusted for placebo response provided a better fit and the results are presented in Table 3.4 and Table 3.5 (unadjusted NMA results are provided in Appendix D). The results of the NMA are presented in terms of the odds ratio (OR) of relief from pain for each intervention versus placebo, sumatriptan and eletriptan. Lasmiditan (OR: 2.53; 95% CrI: 2.04 to 3.25), rimegepant (OR: 2.19; 95% CrI: 1.8 to 2.76), and ubrogepant (OR: 2.19; 95% CrI: 1.7 to 2.89) all had higher odds of achieving pain freedom at two hours versus placebo. Compared to each other, none of the interventions showed a statistically significant difference, though lasmiditan showed a statistically non-significant, higher odds of achieving pain relief. Compared to sumatriptan, all interventions showed lower odds of achieving pain relief, however, only rimegepant was statistically significantly worse (OR: 0.73; 95% CrI: 0.58 to 0.96). Results compared to eletriptan also showed lower odds of achieving pain relief at two hours for the three interventions, and all were statistically significant (lasmiditan: 0.61, rimegepant: 0.52, ubrogepant: 0.52).

Based on the estimated odds ratios, the expected proportion of patients achieving pain relief at two hours was 58% for lasmiditan, 54% for rimegepant, 54% for ubrogepant, 62% for sumatriptan and 69% for eletriptan (Table 3.5).

Intervention		Headache Pai	n Freedom at 2-Hours	Headache Pa	Headache Pain Relief at 2-Hours	
(Trial)	Arms	n/N (%)	Odds Ratio vs. Placebo (95%Cl), p-value	n/N (%)	Odds Ratio vs. Placebo (95%Cl), p-value	
1 constitution	Lasmiditan 200mg	167/518 (32.2)	2.6 (2.0, 3.6), <0.001	330/555 (59.5)	2.5 (1.9, 3.3), <0.001	
Lasmiditan (SAMURAI) ²⁴	Lasmiditan 100mg	142/503 (28.2)	2.2 (1.6, 3.0), <0.001	334/562 (59.4)	2.4 (1.8, 3.1), <0.001	
	Placebo	80/524 (15.3)		234/554 (42.2)		
	Lasmiditan 200mg	205/528 (38.8)	2.3 (1.8, 3.1), <0.001	367/565 (65.0)	2.4 (1.8, 3.1), <0.001	
Lasmiditan	Lasmiditan 100mg	167/532 (31.4)	1.7 (1.3, 2.2), <0.001	370/571 (64.8)	2.3 (1.7, 2.9), <0.001	
(SPARTAN) ²³	Lasmiditan 50mg	159/556 (28.6)	1.5 (1.1, 1.9), 0.003	353/598 (59.0)	1.7 (1.3, 2.2), <0.001	
	Placebo	115/540 (21.3)		274/576 (47.7)		
Rimegepant	Rimegepant 75mg	104/543 (19.2)	1.4 (1.0, 2.0), 0.03	304/543 (56.0)	1.5 (1.2, 1.9), <0.001	
(Study 301) ²⁷	Placebo	77/541 (14.2)	1.4 (1.0, 2.0), 0.05	247/541 (45.7)		
Rimegepant	Rimegepant 75mg	105/537 (19.6)	1.8 (1.3, 2.5), <0.001	312/537 (58.1)	1.9 (1.5, 1.3), <0.0001	
(Study 302) ²⁶	Placebo	64/535 (12.0)	1.8 (1.3, 2.3), <0.001	229/535 (42.8)		
Rimegepant	Rimegepant 75mg	142/669 (21.2)	2.2 (1.6, 3.0), <0.0001	397/669 (59.3)	1.9 (1.5, 2.4), <0.0001	
(Study 303) ²⁸	Placebo	74/682 (10.9)	2.2 (1.0, 5.0), <0.0001	295/682 (43.3)	1.9 (1.3, 2.4), <0.0001	
Ubrogepant	Ubrogepant 100mg	95/448 (21.2)	2.0 (1.4, 3.0), 0.0003	275/448 (61.4)	1.7 (1.3, 2.2), 0.0023	
(ACHIEVE I) ³¹	Ubrogepant 50mg	81/422 (19.2)	1.8 (1.3, 2.7), 0.0023	257/422 (60.7)	1.7 (1.3, 2.2), 0.0023	
	Placebo	54/456 (11.8)		224/456 (49.1)		
Ubrogepant	Ubrogepant 50mg	101/464 (21.8)	1.6 (1.1, 2.3), 0.01	291/464 (62.7)	1.8 (1.4, 2.3), 0.01	
(ACHIEVE II) ³⁰	Ubrogepant 25mg	90/435 (20.7)	1.6 (1.1, 2.2), 0.03	263/435 (60.5)	1.7 (1.3, 2.2), 0.07	
	Placebo	65/456 (14.3)		220/456 (48.2)		

Table 3.2: Phase III Results of Lasmiditan, Rimegepant and Ubrogepant. Pain Freedom and Pain Relief at 2-Hours.

95% CI: 95% confidence interval, mg: milligrams, n: number of participants, N: total number of participants, vs.: versus

Lasmiditan (100/200 mg)					
	Rimegepant				
1.43 (0.97, 2.06)	75 mg		_		
		Ubrogepant			
1.43 (0.93, 2.14)	1 (0.69, 1.46)	(50/100 mg)		_	
			Sumatriptan		
0.73 (0.53, 1.06)	0.51 (0.39, 0.7)	0.52 (0.37, 0.74)	(50/100 mg)		-
0.54 (0.36, 0.85)	0.38 (0.27, 0.57)	0.38 (0.26, 0.59)	0.73 (0.57, 0.97)	Eletriptan 40 mg	
3.01 (2.2, 4.14)	2.11 (1.67, 2.72)	2.12 (1.58, 2.88)	4.09 (3.43, 4.82)	5.6 (4.14, 7.23)	Placebo

Table 3.3: NMA results. Interventions and Comparators. Pain Freedom at 2-Hours

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Table 3.4: NMA results. Interventions and Comparators. Pain Relief at 2-Hours

Lasmiditan (100/200 mg)					
	Rimegepant				
1.16 (0.87, 1.52)	75 mg		_		
		Ubrogepant			
1.15 (0.85, 1.58)	1 (0.75, 1.34)	(50/100 mg)			
			Sumatriptan		
0.84 (0.67, 1.13)	0.73 (0.58, 0.96)	0.73 (0.55, 1)	(50/100 mg)		
0.61 (0.44, 0.88)	0.52 (0.38, 0.76)	0.52 (0.37, 0.78)	0.72 (0.58, 0.89)	Eletriptan 40 mg	
2.53 (2.04, 3.25)	2.19 (1.8, 2.76)	2.19 (1.7, 2.89)	2.99 (2.65, 3.34)	4.18 (3.32, 5.14)	Placebo

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Table 3.5. NMA results versus Placebo. Pain Freedom and Pain Relief at 2-Hours

	Pain Freedo	om at 2-Hours	Pain Relief at 2-Hours		
	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion with Pain Freedom (95% Crl)	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion with Pain Relief (95% Crl)	
Placebo	Reference	0.11	Reference	0.35	
Lasmiditan (100/200 mg)	3.01 (2.2, 4.14)	0.28 (0.22, 0.35)	2.53 (2.04, 3.25)	0.58 (0.52, 0.63)	
Rimegepant (75 mg)	2.11 (1.67, 2.72)	0.21 (0.18, 0.26)	2.19 (1.8, 2.76)	0.54 (0.49, 0.6)	
Ubrogepant (50/100 mg)	2.12 (1.58, 2.88)	0.21 (0.17, 0.27)	2.19 (1.7, 2.89)	0.54 (0.48, 0.61)	
Sumatriptan (50/100 mg)	4.09 (3.43, 4.82)	0.35 (0.31, 0.38)	2.99 (2.65, 3.34)	0.62 (0.59, 0.64)	
Eletriptan (40 mg)	5.6 (4.14, 7.23)	0.42 (0.35, 0.48)	4.18 (3.32, 5.14)	0.69 (0.64, 0.73)	

95% CrI: 95% credible interval, mg: milligrams, vs.: versus

Sustained Pain Freedom

Sustained pain freedom was defined as the percentage of subjects who were pain free at two hours and maintained pain freedom with no use of rescue medication or relapse within 24 (sustained pain freedom at 24 hours) or 48 hours (sustained pain freedom at 48 hours) after the initial treatment. In the individual Phase III clinical trials of the interventions, lasmiditan (50 mg, 100 mg, or 200 mg), rimegepant (75 mg) and 100 mg ubrogepant all resulted in a greater proportion of patients experiencing sustained pain freedom at 24 hours and 48 hours compared with placebo (Table 3.6). The other two doses of ubrogepant (25 mg and 50 mg) were not statistically significantly different from placebo on sustained pain freedom at 24 hours (Table 3.6).

Mainly because of data availability, we conducted NMA only for the 24 hours sustained pain freedom outcome. In total, we identified 15 trials (2 lasmiditan,^{23,24} 4 rimegepant trials including 1 head-to head versus sumatriptan,²⁶⁻²⁹ 3 ubrogepant,³⁰⁻³² and 6 triptan studies^{34,36,46-48}) sufficiently similar to include in the NMA. Appendix Table D6 provides the data for the NMA, including the sample size and the number of patients who reported sustained pain freedom.

The NMA model adjusted for placebo response provided a better fit and the results are presented in Table 3.7 and Table 3.8 (unadjusted NMA results are provided in Appendix D). Consistent with the trials, the NMA results showed that lasmiditan (OR: 2.92; 95% CI: 1.89 to 4.5), rimegepant (OR: 2.51; 95% CI: 1.89 to 3.46), and ubrogepant (OR: 2.32; 95% CI: 1.62 to 3.46) all had higher odds of achieving sustained pain freedom at 24 hours versus placebo. Compared to the triptans, although all interventions showed lower odds of achieving sustained pain freedom at 24 hours compared to sumatriptan (lasmiditan: 0.83, rimegepant: 0.71, ubrogepant: 0.66) and eletriptan (lasmiditan: 0.73, rimegepant: 0.63, ubrogepant: 0.59), these were not statistically significant. Similarly, the interventions were not statistically significantly different from each other (Table 3.7).

Based on the estimated odds ratio, the expected proportion of patients achieving sustained pain freedom at 24 hours was 19% for lasmiditan, 17% for rimegepant, 16% for ubrogepant, 22% for sumatriptan and 24% for eletriptan (Table 3.8). Of note, because of recurrent symptoms after two hours, the number of patients with sustained pain freedom at 24 hours was less than those achieving pain freedom at two hours (see Table 3.8).

Sustained relief is based on a concept similar to sustained pain freedom. It was defined as the percentage of subjects who had pain relief at two hours with no use of rescue medication or relapse at follow-up after the initial treatment. We found no data on sustained pain relief for lasmiditan. In total, we included the four rimegepant trials, the three ubrogepant trials and 10 triptan trials for the NMA on sustained pain relief (see Appendix Table D6). The results of the NMA on sustained pain relief followed a similar pattern as the 24 hours sustained pain freedom (see Appendix Table D15).

Intervention		Sustained Pain Freedom at 24-Hours		Sustained Pain Freedom at 48-Hours	
(Trial)	Arms	n/N (%)	Odds Ratio vs. Placebo (95%Cl), p-value	n/N (%)	Odds Ratio vs. Placebo (95%Cl), p-value
	Lasmiditan 200mg	103/555 (18.6)	2.8 (1.9, 4.1), <0.001	91/555 (16.4)	2.4 (1.6, 3.5), <0.001
Lasmiditan (SAMURAI) ²⁴	Lasmiditan 100mg	83/562 (14.8)	2.1 (1.4, 3.1), <0.001	84/562 (14.9)	2.1 (1.5, 3.2), <0.001
	Placebo	42/554 (7.6)		42/554 (7.6)	
	Lasmiditan 200mg	128/565 (22.7)	1.9 (1.4, 2.6), <0.001	111/565 (19.6)	1.8 (1.3, 2.5), <0.001
Lasmiditan	Lasmiditan 100mg	102/571 (17.9)	1.4 (1.0, 1.9), 0.021	86/571 (15.1)	1.3 (0.9, 1.9), 0.058
(SPARTAN) ²³	Lasmiditan 50mg	103/598 (17.2)	1.3 (1.0, 1.9), 0.036	89/598 (14.9)	1.3 (0.9, 1.8), 0.065
	Placebo	77/576 (13.4)		68/576 (11.8)	
Rimegepant	Rimegepant 75mg	76/543 (14.0)	18(12)27)0002*	63/543 (11.6)	
(Study 301) ²⁷	Placebo	44/541 (8.1)		39/541 (7.2)	1.7 (1.1, 2.6), 0.013*
Rimegepant	Rimegepant 75mg	66/537 (12.3)	1.8 (1.2, 2.8), 0.004*	53/537 (9.9)	1.7 (1.1, 2.7), 0.02*
(Study 302) ²⁶	Placebo	38/535 (7.1)	1.8 (1.2, 2.8), 0.004	32/535 (6.0)	1.7 (1.1, 2.7), 0.02
Rimegepant	Rimegepant 75mg	105/669 (15.7)	3.2 (2.1, 4.7), <0.0001*	90/669 (13.5)	2.7 (1.8, 4.1), <0.0001*
(Study 303) ²⁸	Placebo	38/682 (5.6)	5.2 (2.1, 4.7), \0.0001	37/682 (5.4)	2.7 (1.0, 4.1), <0.0001
Libus sevent	Ubrogepant 100mg	68/441 (15.4)	2.0 (1.3, 3.0), 0.0037		
Ubrogepant (ACHIEVE I) ⁹⁰	Ubrogepant 50mg	53/418 (12.7)	1.6 (1.0, 2.4), n.s.	NR	
	Placebo	39/452 (8.6)			
	Ubrogepant 50mg	66/457 (14.4)	1.9 (1.2, 2.8), 0.01		
Ubrogepant (ACHIEVE II) ³⁰	Ubrogepant 25mg	55/432 (12.7)	1.6 (1.0, 1.8), n.s.	NR	
	Placebo	37/451 (8.2)			

Table 3.6: Phase III Results of Lasmiditan, Rimegepant and Ubrogepant. Sustained Pain Freedom at 24- and 48-Hours

*Odds ratio estimated

95%CI: 95% confidence interval, mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, vs.: versus

Lasmiditan (100/200 mg)					
	Rimegepant (75				
1.16 (0.67, 1.94)	mg)		_		
		Ubrogepant			
1.26 (0.72, 2.11)	1.08 (0.67, 1.74)	(50/100 mg)		_	
0.83 (0.5, 1.44)	0.71 (0.48, 1.12)	0.66 (0.41, 1.12)	Sumatriptan		
0.73 (0.34, 1.53)	0.63 (0.32, 1.22)	0.59 (0.28, 1.18)	0.89 (0.44, 1.69)	Eletriptan	
2.92 (1.89, 4.5)	2.51 (1.89, 3.46)	2.32 (1.62, 3.46)	3.53 (2.52, 4.77)	3.97 (2.24, 7.36)	Placebo

Table 3.7. NMA Results. All Interventions and Comparators. Sustained Pain Freedom at 24-Hours

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Table 3.8. NMA Results versus Placebo.Sustained Pain Freedom at 24-Hours Compared to PainFreedom Achieved at 2-Hours

	Sustained Pain Fr	eedom at 24-hours	Pain Freedom at 2-hours		
	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion with Sustained Pain Freedom (95% Crl)	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion with Pain Freedom (95% Crl)	
Placebo	Reference	0.07	Reference	0.11	
Lasmiditan 100/200 mg	2.92 (1.89, 4.5)	0.19 (0.13, 0.26)	3.01 (2.2, 4.14)	0.28 (0.22, 0.35)	
Rimegepant 75 mg	2.51 (1.89, 3.46)	0.17 (0.13, 0.22)	2.11 (1.67, 2.72)	0.21 (0.18, 0.26)	
Ubrogepant 50/100 mg	2.32 (1.62, 3.46)	0.16 (0.11, 0.22)	2.12 (1.58, 2.88)	0.21 (0.17, 0.27)	
Sumatriptan 50/100 mg	3.53 (2.52, 4.77)	0.22 (0.17, 0.27)	4.09 (3.43, 4.82)	0.35 (0.31, 0.38)	
Eletriptan 40 mg	3.97 (2.24, 7.36)	0.24 (0.15, 0.37)	5.6 (4.14, 7.23)	0.42 (0.35, 0.48)	

95% CrI: 95% credible interval, mg: milligrams, NA: not available, vs.: versus

Freedom from Most Bothersome Symptom (MBS)

Absence of the most bothersome migraine associated symptom (i.e. phonophobia, photophobia, or nausea) at two hours after treatment was measured as a co-primary endpoint in the Phase III trials of lasmiditan, rimegepant and ubrogepant. None of the Phase II studies of the interventions or the triptan studies assessed freedom from MBS as an outcome. As such we included only the seven Phase III trials in our NMA and compared the interventions to each other and to placebo.^{23,24,26-28,30,31,91}

Table 3.9 presents the results of the Phase III trials. A greater proportion of patients on lasmiditan (50 mg, 100 mg, or 200 mg), rimegepant (75 mg) or ubrogepant (25 mg, 50 mg or 100 mg) experienced freedom from MBS at two hours post dose compared with patients on placebo. The unadjusted NMA comparing the interventions to each other provided a better fit and the results are presented in Table 3.10. The results showed that lasmiditan (OR: 1.99; 95% CI: 1.03 to 3.9), rimegepant (OR: 2.57; 95% CI: 1.61 to 4.26), and ubrogepant (OR: 2.09; 95% CI: 1.19 to 3.9) all had higher odds of achieving freedom from MBS at two hours post dose compared to placebo. However, compared to each other, none of the interventions showed a statistically significant difference. Based on the estimated odds ratio, the expected proportion of patients achieving freedom from MBS at two hours was 40% for lasmiditan, 38% for rimegepant, and 39% for ubrogepant.

Intervention		Freedom From Mo	ost Bothersome Symptom at 2-Hours
(Trial)	Arms	n/N (%)	Odds Ratio vs. Placebo (95%Cl), p-value
	Lasmiditan 200mg	196/481 (40.7)	1.6 (1.3, 2.1), <0.001
Lasmiditan (SAMURAI) ²⁴	Lasmiditan 100mg	192/469 (40.9)	1.7 (1.3, 2.2), <0.001
	Placebo	144/488 (29.5)	
	Lasmiditan 200mg	235/483 (48.7)	1.9 (1.4, 2.4), <0.001
Lasmiditan	Lasmiditan 100mg	221/500 (44.2)	1.6 (1.2, 2.0), <0.001
(SPARTAN) ²³	Lasmiditan 50mg	209/512 (40.8)	1.4 (1.1, 1.8), 0.009
	Placebo	172/514 (33.5)	
Rimegepant (Study 301) ²⁷	Rimegepant 75mg	199/543 (36.6)	1.5 (1.2, 2.0), 0.002
	Placebo	150/541 (27.7)	1.3 (1.2, 2.0), 0.002
Rimegepant	Rimegepant 75mg	202/537 (37.6)	1.8 (1.4, 2.3), <0.0001
(Study 302) ²⁶	Placebo	135/535 (25.2)	1.8 (1.4, 2.3), <0.0001
Rimegepant	Rimegepant 75mg	235/669 (35.1)	1.5 (1.2, 1.9), 0.001
(Study 303) ²⁸	Placebo	183/682 (26.8)	1.5 (1.2, 1.9), 0.001
	Ubrogepant 100mg	169/448 (37.7)	1.6 (1.2, 2.2), 0.0023
Ubrogepant (ACHIEVE I) ⁹⁰	Ubrogepant 50mg	163/420 (38.6)	1.7 (1.3, 2.3), 0.0023
	Placebo	127/454 (27.8)	
	Ubrogepant 50mg	180/463 (38.9)	1.7 (1.3, 2.2), 0.01
Ubrogepant (ACHIEVE II) ³⁰	Ubrogepant 25mg	148/434 (34.1)	1.4 (1.0, 1.8), 0.07
	Placebo	125/456 (27.4)	

Table 3.9: Phase III Results of Lasmiditan, Rimegepant and Ubrogepant. MBS Freedom at 2-Hours

mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, vs: versus

Table 3.10. NMA Results. Interventions and Comparators. Freedom from MBS at 2-Hours

Lasmiditan (100/200 mg)			
1.07 (0.78, 1.46)	Rimegepant (75 mg)		
1.03 (0.73, 1.45)	0.96 (0.69, 1.33)	Ubrogepant (50/100 mg)	
1.69 (1.33, 2.14)	1.58 (1.29, 1.94)	1.64 (1.28, 2.12)	Placebo

mg: milligrams

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Freedom from Other Migraine Symptoms (phonophobia, photophobia and nausea)

Freedom from phonophobia, photophobia, and nausea were assessed as secondary outcomes in the trials of lasmiditan, rimegepant and ubrogepant. However, there was a lack of consistency in how these outcomes were analyzed across trials. In the rimegepant trials, freedom from migraine associated symptoms were evaluated correctly among patients who exhibited these symptoms at baseline, while the trials of lasmiditan and ubrogepant evaluated these outcomes among all patients, irrespective of their baseline symptoms. As such we did not quantitatively compare the drugs to each other on these outcomes.

Regardless of how the trials evaluated these outcomes, all three interventions were not different from placebo in achieving freedom from nausea at two hours in any of the Phase III trials. All interventions had higher odds of achieving freedom from phonophobia and freedom from photophobia at two hours post dose compared to placebo (Appendix Table D8).

Use of Rescue Medication

Due to differences in the design of the trials related to the use of rescue medication (e.g. open label second dose vs. randomized; NSAID vs. usual acute migraine treatment), we could not quantitatively compare the interventions to each other on this outcome (see Table 3.11). In general, patients who were randomized to the interventions were less likely to use a second dose or another medication for rescue compared to patients on placebo.

	Lasmiditan Phase III	Rimegepant Phase III Trials	Ubrogepant Phase III Trials
	Trials		
	Timing and Indi	cation for Rescue Medication	
Initial Response	Rescue medication could be used within 24 hours if pain freedom not achieved at 2 hours.	Rescue medication could be used within 48 hours if pain relief not achieved at 2 hours.	Rescue medication could be used within 48 hours if pain relief not achieved at 2 hours.
Recurrence	Patients could take a rescue medication for recurrence within 24 hours	Patients could take a rescue medication for recurrence within 48 hours	Patients could take a rescue medication for recurrence within 48 hours
	Rescue	Medication Allowed	
Second dose of study	Patients were re-	Patients were not given an	Patients were given an
Medication	randomized to an optional second dose of placebo or lasmiditan. Second dose only taken if another rescue medication has not been used.	optional second dose	optional second dose (those on placebo were given placebo and others were re- randomized to placebo or ubrogepant). Second dose only taken if another rescue medication has not been used.
Other Medications	Triptans, ergots, opioids and barbiturates were <u>not</u> <u>allowed</u> . Patients could take other over the counter medications of choice.	Triptans, ergots, opioids and barbiturates were <u>not allowed</u> within 48 hours. Patients could take aspirin, NSAIDs, acetaminophen, antiemetics, or baclofen.	Patients could take triptans, ergots, NSAIDs, acetaminophen, opioids, or other over the counter medications.

Table 3.11. Use of Rescue Medication after 2 Hours

In the Phase III trials of lasmiditan, all patients were randomly allocated to an optional second dose of the study drug. Patient with persistent or recurrent pain wanting to take additional treatment could take the optional second dose or their own rescue medication within two to 48 hours after the initial dose. The second dose was used between two and 24 hours in 32% to 39% of the lasmiditan group (200/100 mg) versus 60% of the placebo group in the SPARTAN trial; and 20% to 35% of the lasmiditan group (200/100/50 mg) versus 40% of the placebo in the SAMURAI trial.^{23,24} Of these second doses, approximately 95% were taken as rescue medication, while the remaining were taken for pain recurrence.

The rimegepant trials did not provide patients with an optional second dose of study medication but allowed the use of rescue medications. Across the four rimegepant trials, 14% to 21% of patients on rimegepant used a rescue therapy compared to 30% to 37% for patients on placebo.²⁶⁻²⁹

In the Phase III trials of ubrogepant, patients were re-randomized to an optional second dose of ubrogepant. Patient with persistent or recurrent pain wanting to take additional treatment could opt to take the optional second dose or their own rescue medication within two to 48 hours after the initial dose. In the pooled ubrogepant group, 38% of patients used an optional second dose compared with 43% in the placebo group. Rates of rescue medication use after the first dose was approximately 15% in the ubrogepant group versus 21% to 29% in the placebo group.

Disability

Functional disability was measured as a secondary outcome in all the Phase III trials of the interventions. This was assessed at two hours after initial treatment, before the use of rescue medication with a four-point functional disability scale (0=no disability [i.e. ability to function normally]; 1=mild disability [i.e. ability to perform all activities of daily living but with some difficulty]; 2=moderate disability [unable to perform certain activities of daily living]; 3=severe disability [i.e. unable to perform most to all activities of daily living or requiring bed rest]). This outcome was not consistently evaluated in the included triptan studies. As such we included only the seven Phase III trials in our NMA and compared the interventions to each other and to placebo. ^{23,24,26-28,90,91}

Table 3.12 presents the results of the Phase III trials. A greater proportion of patients on lasmiditan (50 mg, 100 mg, or 200 mg), rimegepant (75 mg) and ubrogepant (25 mg, 50 mg or 100 mg) were able to function normally at two hours post dose compared with patients on placebo. The unadjusted NMA comparing the interventions to each other provided a better fit and the results are presented in Table 3.13. The NMA showed that lasmiditan (OR:1.7; 95% CI:1.32 to 2.20), rimegepant (OR:1.72; 95% CI: 1.38 to 2.14), and ubrogepant (OR: 1.51; 95% CI: 1.15 to 1.96) all had higher odds of achieving no disability at two hours post dose compared to placebo. However, compared to each other, none of the interventions showed a statistically significant difference (Table 3.13). Based on the estimated odds ratio, the expected proportion of patients who could function normally at two hours post dose was 38% for lasmiditan, 38% for rimegepant, and 35% for ubrogepant.

Table 3.12. Phase III results of Lasmiditan, Rimegepant and Ubrogepant. Ability to FunctionNormally at 2-Hours

Intervention		Ability to Function Normally at 2-Hours		
(Trial)	Arms	n/N (%)	p-value vs. Placebo	
Lasmiditan (SAMURAI) ²⁴	Lasmiditan 200mg	180/555 (32.4)	<0.001	
	Lasmiditan 100mg	181/562 (32.2)	<0.001	
	Placebo	119/554 (21.5)	Reference	
	Lasmiditan 200mg	209/565 (37.0)	<0.001	
Lasmiditan	Lasmiditan 100mg	193/571 (33.8)	<0.001	
(SPARTAN) ²³	Lasmiditan 50mg	187/598 (31.3)	0.019	
	Placebo	143/576 (24.8)	Reference	
Rimegepant	Rimegepant 75mg	181/543 (33.3)	<0.0001	
(Study 301) ²⁷	Placebo	118/541 (21.8)	<0.0001	
Rimegepant	Rimegepant 75mg	175/537 (32.6)	NR	
(Study 302) ²⁶	Placebo	125/535 (23.4)		
Rimegepant	Rimegepant 75mg	255/669 (38.1)	NR	
(Study 303) ²⁸	Placebo	176/682 (25.8)		
Librogonant	Ubrogepant 100mg	193/423 (42.9)	<0.01	
Ubrogepant (ACHIEVE I) ⁹⁰	Ubrogepant 50mg	172/448 (40.6)	<0.01	
	Placebo	136/456 (29.8)	Reference	
	Ubrogepant 50mg	188/464 (40.5)	<0.01	
Ubrogepant (ACHIEVE II) ³⁰	Ubrogepant 25mg	185/435 (42.6)	<0.01	
	Placebo	156/456 (34.2)	Reference	

mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, vs.: versus

Table 3.13. NMA results. Interventions and Comparators. Ability to Function Normally at 2-Hours

Lasmiditan (100/200 mg)			
0.99 (0.71, 1.39)	Rimegepant (75 mg)		_
1.13 (0.78, 1.64)	1.14 (0.81, 1.62)	Ubrogepant (50/100 mg)	
1.7 (1.32, 2.2)	1.72 (1.38, 2.14)	1.51 (1.15, 1.96)	Placebo

Legend: Each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

mg: milligrams

Patient Global Impression of Change

Patient global impression of change (PGIC) was measured as a secondary outcome in the Phase III trials of lasmiditan and ubrogepant. This was assessed at two hours after initial treatment, before the use of rescue medication with a seven-point scale (1=very much worse; 2=much worse; 3=a little worse; 4=no change; 5= a little better; 6=much better; 7=very much better). The results of the trials showed that a higher proportion of ubrogepant or lasmiditan-treated patients indicated their migraine was much better/very much better at two hours post dose compared with placebo-treated patients (Table 3.14). We did not identify any PGIC data on rimegepant.

Trial	Arms	Ν	PGIC (% That Achieved "Very Much Better and Much Better")	p-value vs. Placebo
SAMURAI ²⁴	Lasmiditan 200mg	555	37.9	<0.001
	Lasmiditan 100mg	562	37.2	<0.001
	Placebo	554	21.8	Reference
SPARTAN ²³	Lasmiditan 200mg	565	42.5	<0.001
	Lasmiditan 100mg	571	41.2	<0.001
	Lasmiditan 50mg	598	36.6	<0.001
	Placebo	576	28.0	Reference
ACHIEVE I ⁹⁰	Ubrogepant 50mg	297	34.3	<0.001
	Ubrogepant 100mg	299	34.4	<0.001
	Placebo	313	22.0	Reference
ACHIEVE II ³⁰	Ubrogepant 50mg	392	33.4	<0.001
	Ubrogepant 25mg	435	34.1	<0.001
	Placebo	376	20.7	Reference

Table 3.14. Phase III Results of Lasmiditan and Ubrogepant. PGIC at 2-Hours.

mg: milligrams, N: total number of participants, PGIC: Patient Global Impression of Change, vs.: versus

Reduction in Migraine Days per Month

Stakeholders identified that decreased frequency and severity of migraine attacks was a potential benefit of lasmiditan, rimegepant and ubrogepant when used over time, something that had not been shown with the use of triptans. We did not perform a systematic review specifically to address this issue, however we examined this potential benefit and our interpretation of the evidence.

The available Phase III RCTs on the interventions of interest are short-term single dose studies, and so were not designed to provide information on changes in migraine frequency or severity over time. Evidence related to this outcome was all from long-term open label extension (OLE) studies that were uncontrolled. Specifically, we identified two OLE studies (GLADIATOR and Lipton 2019) that evaluated this outcome.^{56,88}

In GLADIATOR, two lasmiditan doses (100 mg and 200 mg) taken as needed were evaluated in 2,037 patients over one year, but only 847 patients completed the study.⁵⁶ Overall, the mean number of migraine days per month was reported to have decreased from a baseline rate of 15.5 days per month to 8.2 days per month in the 200 mg lasmiditan group (mean change -7.3 migraine days/month) and to 8.8 days per month in the 100 mg lasmiditan group (mean change -6.7 migraine days/month) at one year. In addition, the migraine disability assessment (MIDAS) score was reported to be reduced by approximately 50% in both groups by the end of the first year.

Lipton 2019 evaluated 75 mg rimegepant taken as needed (PRN group, n=1,498) or on schedule (taken every other day) plus as needed (QOD+PRN group, n=286) over one year, but patient followup over time was not reported.⁸⁸ At three months, the trial reported a mean reduction of 4 migraine days per month among patients observed to have 14 or more migraine days/month at baseline (in both rimegepant group). For patients in the QOD+PRN group, approximately half reported a \geq 50% reduction from baseline in the frequency of monthly migraine days of moderate to severe pain intensity at three months, regardless of baseline migraine days.

While the results of these studies reported a decreasing frequency of migraine attacks over time, we were concerned about study design and reporting issues that may bias these results. We felt that patients with a high frequency of attacks at baseline may experience decreases over time simply due to regression to the mean. Because these were uncontrolled studies without a placebo arm, it is not possible to differentiate regression to the mean from placebo effect or from an actual benefit. We were also concerned that patients who may have had the greatest migraine burden and were not benefitting from therapy might drop out over time, leaving patients at later follow-up points who were having fewer migraines at baseline and thus overestimating any decrease in migraine frequency or severity.

Several lines of evidence support our concerns about regression to the mean as playing a prominent role in the reported data from OLE trials. First, it is notable that therapies with very different mechanisms of action (lasmiditan and rimegepant) should both show reductions in headache frequency over time when prior acute migraine therapies have not done so in controlled trials. Moreover, it is unexpected that lasmiditan, which works through a mechanism closely related to triptans, would show this benefit when triptans are not believed to have such a benefit. To explore this issue further, we reviewed a trial comparing telcagepant (a gepant) with rizatriptan (a triptan) in more than 1000 patients.⁹² We reproduce below a figure showing similar reduction in headache frequency over time including in the triptan arm, as would be expected with regression to the mean (Figure 3.1).

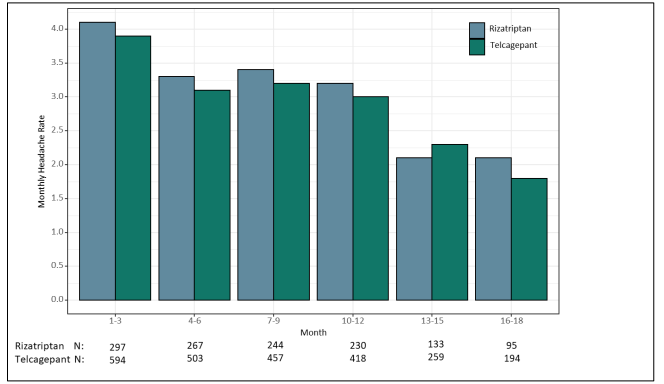
Loss of follow-up over time in the GLADIATOR trial was large (51.7%) and suggests that using the larger denominator at baseline but a smaller one at follow-up may affect the reported results. The most common reason for discontinuation was "patient request' (21.8%), which likely referred to those patients who discontinued the medication for lack of effect. In the rimegepant OLE trial,

information was obtained from a conference abstract, so information on dropout is unclear. Based on the data in the poster, only 17.6% of patients that were evaluated at 12 weeks were included in the reported analysis.

Finally, regarding placebo effect, we note that the response rate in the placebo arms of the single dose RCTs ranged from 25% to 51%, and that in ICER's prior report on migraine prevention, the placebo response rate for prophylactic therapy ranged from 10% to 62%. In addition, the mean decrease in migraine-specific days and migraine-specific medication consumption per month was considerably smaller in RCTs of CGRP monoclonal antibodies for prevention of migraine attacks.⁶⁴

Given these concerns, we do not feel that current evidence supports a conclusion that treatment with lasmiditan, rimegepant, or ubrogepant decreases migraine frequency over time. A placebocontrolled trial would likely be needed to explore this issue, and in the absence of such a trial, we do not think patients or clinicians should select one of these medications based upon such a treatment-specific benefit.

Figure 3.1. Mean monthly headache rate. Adapted from "Long-Term Tolerability of Telcagepant for Acute Treatment of Migraine in a Randomized Trial," by Connor KM, Aurora SK, Loeys T, et al. Headache. 2011 Jan;51(1):73-84



N: total number of participants

Harms

Randomized Controlled Trials

The majority of adverse events observed in the single-attack trials were mild or moderate in intensity. Adverse events (AEs) with incidence ≥5% in any of the treatment arm are presented in Appendix Table D10. In the lasmiditan trials, central nervous system (CNS)-related AEs (e.g., dizziness, somnolence, paresthesia) were the most frequently reported AE, with dizziness the most common. Nausea was among the most commonly reported AE in the ubrogepant and rimegepant trials (1% to 3%). In general, there was a low incidence of serious adverse events in these trials. There was a low or no incidence of cardiovascular related AEs in the trials.

Table 3.15 presents the data on AEs, treatment-emergent AEs (TEAEs), and most frequent AEs from the Phase III trials of the interventions. In the Phase III trials, TEAEs among patients on placebo ranged from 1% to 3%, while they ranged from 6% to 12% in patients on CGRP antagonists and 32% to 38% among those on lasmiditan. In total, 24 trials (including the Phase II trials and the triptan studies) reported on the number of patients who experienced any type of adverse event (any AE)^{23,24,26-28,32,33,35-37,39,40,42,44,45,47,48,50-53,90,91} and 16 trials (including the Phase II trials and the triptan studies) reported on the number of patients who experienced any treatment emergent adverse event (any TEAE).^{23,25,27,28,32,33,640,46,50,51,53,54,90,91} We considered all the trials sufficiently similar to include in the NMA. Appendix Table D7 provides the data for the NMA, including the sample size and the number of patients who reported pain freedom.

The unadjusted NMAs on any AE and TEAE provided a better fit and the results are presented in Table 3.15 and Appendix Table D16-D18. The NMA results are expressed as Ors, where values greater than one indicate a higher odd of any AE or TEAE for the active therapy versus placebo. Lasmiditan had higher odds of any AE compared to placebo (3.91, 95% Crl: 2.45, 6.25, Table 3.15), rimegepant (3.13, 95% Crl: 1.69, 5.82), ubrogepant (3.51, 95% Crl: 1.86, 6.61), sumatriptan (2.16, 95% Crl: 1.27, 3.56), and eletriptan (3.66, 95% Crl: 2.03, 6.51) (Appendix Table D16). Compared to placebo, both rimegepant and ubrogepant had point estimates with higher odds of any AE, but these were not statistically significant. There was also no statistically significant difference between rimegepant and ubrogepant, and these agents versus the triptans. Based on the estimated odds ratio, the expected proportion of patients achieving any AE was 50% for lasmiditan, 24% for rimegepant, 22% for ubrogepant, 31% for sumatriptan and 21% for eletriptan (Table 3.15).

In terms of TEAEs, lasmiditan had higher odds of TEAE compared to placebo (5.99, 95% Crl: 3.3, 12.52, Table 3.15), rimegepant (4.00, 95% Crl: 1.38, 12.04), ubrogepant (5.10, 95% Crl: 2.31, 12.95), and sumatriptan (2.57, 95% Crl: 1.3, 6.07). The point estimate compared to eletriptan was 3.27, however it was not statistically significant (95% Crl: 1, 11.83). Both rimegepant and ubrogepant were not statistically significantly different from placebo, sumatriptan, and eletriptan (Appendix Table D17). However, both rimegepant and ubrogepant had point estimates with lower odds of

TEAEs compared to sumatriptan and eletriptan. Based on the estimated odds ratio, the expected proportion of patients achieving any AE was 42% for lasmiditan, 15% for rimegepant, 12% for ubrogepant, 22% for sumatriptan and 18% for eletriptan (Table 3.15).

We also quantitatively compared the incidence of dizziness, the most frequent AE that was consistently reported in the trials. Lasmiditan had higher odds of causing dizziness compared to placebo (8.43, 95% Crl: 4.88, 19.35, Table 3.17), rimegepant (7.02, 95% Crl: 2.2, 25.63), ubrogepant (4.95, 95% Crl: 1.67, 15.92), sumatriptan (4.09, 95% Crl: 2, 10.6), and eletriptan (3.97, 95% Crl: 1.44, 12.41) (Appendix Table D18). Based on the estimated odds ratios, the expected proportion of patients experiencing dizziness was 14% for lasmiditan, 2% for rimegepant, 3% for ubrogepant, 4% for sumatriptan and 4% for eletriptan (Table 3.17).

Intervention (Trial)	Arms	N	SAEs, n (%)	Any Aes, n (%)	TEAEs, n (%)	Dizziness, n (%)	Somnolence, n (%)	Paresthesia, n (%)	Nausea, n (%)
	Lasmiditan 200mg	609	2 (0.3)	260 (42.7)	237 (38.9)	99 (16.3)	33 (5.4)	48 (7.9)	32 (5.3)
Lasmiditan (SAMURAI) ²⁴	Lasmiditan 100mg	630	0 (0)	229 (36.3)	205 (32.5)	79 (12.5)	36 (5.7)	36 (5.7)	19 (3.0)
	Placebo	617	1 (0.2)	101 (16.4)	78 (12.6)	21 (3.4)	14 (2.3)	13 (2.1)	12 (1.9)
	Lasmiditan 200mg	649	1 (0.2)	253 (39.0)	NR	117 (18.0)	42 (6.5)	43 (6.6)	17 (2.6)
Lasmiditan	Lasmiditan 100mg	635	1 (0.2)	230 (36.2)	NR	115 (18.1)	29 (4.6)	37 (5.8)	21 (3.3)
(SPARTAN) ²³	Lasmiditan 50mg	654	0 (0)	167 (25.5)	NR	56 (8.6)	35 (5.4)	16 (2.4)	18 (2.8)
	Placebo	645	0 (0)	75 (11.6)	NR	16 (2.5)	13 (2.0)	6 (0.9)	8 (1.2)
Rimegepant	Rimegepant 75mg	546	2 (0.4)	69 (12.6)	3 (0.5)	4 (0.7)	NR	NR	5 (0.9)
(Study 301) ²⁷	Placebo	549	1 (0.2)	59 (10.7)	1 (0.2)	2 (0.4)	NR	NR	6 (1.1)
Rimegepant	Rimegepant 75mg	537	1 (0.2)	93 (17.3)	NR	NR	NR	NR	10 (1.8)
(Study 302) ²⁶	Placebo	535	2 (0.4)	77 (14.4)	NR	NR	NR	NR	6 (1.1)
Rimegepant	Rimegepant 75mg	682	0 (0)	90 (13.5)	47 (6.9)	6 (0.9)	NR	NR	11 (1.6)
(Study 303) ²⁸	Placebo	693	0 (0)	73 (10.5)	36 (5.2)	7 (1.0)	NR	NR	3 (0.4)
Ubrogepant	Ubrogepant 100mg	485	2 (0.4)	79 (16.3)	58 (12.0)	7 (1.4)	12 (2.5)	NR	20 (4.1)
(ACHIEVE I) ⁹⁰	Ubrogepant 50mg	466	3 (0.6)	44 (9.4)	27 (5.8)	4 (0.9)	3 (0.6)	NR	8 (1.7)
	Placebo	485	0 (0)	62 (12.8)	41 (8.5)	3 (0.6)	4 (0.8)	NR	8 (1.6)
Ubrogepant	Ubrogepant 50mg	488	0 (0)	63 (12.9)	42 (8.6)	7 (1.4)	4 (0.8)	NR	10 (2.0)
(ACHIEVE II) ³⁰	Ubrogepant 25mg	478	0 (0)	44 (9.2)	30 (6.3)	10 (2.1)	4 (0.8)	NR	12 (2.5)
	Placebo	499	0 (0)	51 (10.2)	30 (6.0)	8 (1.6)	2 (0.4)	NR	10 (2.0)

Table 3.16. Adverse Events. Phase III Single-Attack Trials of Lasmiditan, Rimegepant, and Ubrogepant

AEs: adverse events, mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, SAEs: serious adverse events, TEAEs:

treatment-emergent adverse events

Table 3.17. NMA results.	Any Adverse Event and Treatment Emergent Adverse Event (Single-
Attack RCTs)	

	Any Advers	se Event (AE)	Treatment Emergent Adverse Event (TEAE)		
	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion with Any AE (95% Crl)	Odds Ratio vs. Placebo (95% CrI)	Expected Proportion with TEAEs (95% Crl)	
Placebo	Reference	0.20	Reference	0.13	
Lasmiditan	3.91 (2.45, 6.25)	0.5 (0.38, 0.61)	5.99 (3.3, 12.52)	0.42 (0.29, 0.6)	
Rimegepant	1.25 (0.83, 1.87)	0.24 (0.17, 0.32)	1.5 (0.67, 3.71)	0.15 (0.08, 0.31)	
Ubrogepant	1.11 (0.73, 1.71)	0.22 (0.16, 0.3)	1.17 (0.68, 2.03)	0.12 (0.08, 0.2)	
Sumatriptan	1.82 (1.48, 2.27)	0.31 (0.27, 0.36)	2.33 (1.58, 3.29)	0.22 (0.16, 0.29)	
Eletriptan	1.07 (0.76, 1.52)	0.21 (0.16, 0.28)	1.83 (0.65, 5.24)	0.18 (0.07, 0.39)	

95% CrI: 95% credible interval, vs.: versus

Table 3.18. NMA Results. Dizziness (Single-Attack RCTs)

	Odds Ratio vs. Placebo (95% Crl)	Expected Proportion With Dizziness (95% Crl)
Placebo	NA	0.02
Lasmiditan	8.43 (4.88, 19.35)	0.14 (0.09, 0.27)
Rimegepant	1.22 (0.44, 3.48)	0.02 (0.01, 0.06)
Ubrogepant	1.73 (0.73 <i>,</i> 4.52)	0.03 (0.01, 0.08)
Sumatriptan	2.07 (1.3, 3.34)	0.04 (0.02, 0.06)
Eletriptan	2.14 (0.96, 5.11)	0.04 (0.02, 0.09)

95% CrI: 95% credible interval, NA: not available, vs.: versus

Long-Term Studies

We present data on any AE and discontinuation due to AEs from the interim analysis of the OLEs of the interventions in Table 3.18. The majority of AEs observed in these trials were mild or moderate in intensity. Similar to the RCTs, most of the AEs observed in the OLE of lasmiditan after 12 months of follow up were CNS-related, with the most frequently reported event being dizziness (21.3% of patients in the 100 mg group, and 15.8% in the 200 mg group). Somnolence occurred in 8-9% of patients and paresthesia occurred in 5-8% of patients.

In total, 12.8% of patients discontinued the trial due to adverse events (11.2% of patients in the 100 mg group, and 14.4% in the 200 mg group), and dizziness was reported to be the most common AE leading to discontinuation (2.7% of patients in the 100 mg group, and 4.3% of patients in 200 mg group). There was no incidence of abuse, misuse, or diversion related to the CNS effects of lasmiditan. Of note, one patient on lasmiditan experienced a road traffic accident during the OLE, although dosing was reported to have occurred two days before the accident, and the patient was also on concomitant medications that have CNS-related effect (lithium and quetiapine). Due to concerns about somnolence with lasmiditan, the FDA label advises that patients should not drive or

operate machinery within 8 hours of taking a dose.⁵⁵ Compared to the lasmiditan OLE, rates of discontinuation were lower in the OLEs of rimegepant and ubrogepant (Table 3.18).

Table 3.19. Adverse Events and Discontinuation due to Adverse Events. Results of 12-monthsOLEs

Intervention (Trial)	Arms	N	Discontinuation due to AE, n (%)	SAEs, n (%)	Any AE, n (%)	Dizziness, n (%)
	Lasmiditan 200mg	1015	146 (14.4)	32 (3.2)	731 (72.0)	217 (21.3)
Lasmiditan (GLADIATOR) ⁵⁶	Lasmiditan 100mg	963	108 (11.2)	28 (2.9)	636 (66.0)	153 (15.8)
Rimegepant (Study 201) ⁸⁸	Rimegepant 75mg	1784	48 (2.7)	45 (2.5)	1062 (59.5)	39 (2.2)
Ubrogepant	Ubrogepant 100mg	409	11 (2.7)	12 (2.9)	297 (72.6)	12 (2.9)
(NCT02873221) ^{89,93,94}	Ubrogepant 50mg	417	9 (2.2)	9 (2.2)	268 (66.3)	5 (1.2)

AE: adverse event, mg: milligrams, n: number of participants, N: total number of participants, SAEs: serious adverse events

Subgroup Analyses

Prior Use of Triptans:

We identified two subgroup analyses that evaluated outcomes among patients in the lasmiditan and ubrogepant trials based upon their prior use of triptans (Knivel 2018 and Blumenfeld 2019).

Knivel 2018 was a pooled analysis of the Phase III trials of lasmiditan (SAMURAI and SPARTAN). At baseline, patients had rated themselves as good, poor, or nonresponders based on three months historical triptan use. The analysis included only patients that were randomized to receive either lasmiditan 100 mg or 200 mg, or placebo in the RCTs. The results showed no significant difference in the benefit of lasmiditan 200 mg versus placebo (on headache pain freedom, MBS freedom, and headache pain relief) in the different triptan responder subgroups.⁹⁵

Blumenfeld 2019 was a pooled analysis of the Phase III trials of ubrogepant (ACHIEVE I and II). At baseline, patients were categorized as triptan-responder, triptan-insufficient responder (includes lack of efficacy, tolerability or contraindications), or triptan-naïve, based on historical experience. Although, higher response rates were observed for ubrogepant 50mg versus placebo in the triptan-responder (2-hour pain freedom OR 2.03; 95%CI: 1.32, 3.11) and triptan-insufficient responder subgroups (2-hour pain freedom OR 2.16; 95%CI: 1.19, 3.95) compared to triptan-naïve subgroup (2-hour pain freedom OR 1.37; 95%CI: 0.94, 2.01), the benefit of ubrogepant 50 mg versus placebo was not significantly different (on 2-hours pain freedom [p=0.29), 2-hours freedom from MBS

[p=0.70]) among the three triptan subgroups, indicating comparable treatment effect regardless of historical triptan experience.⁹⁶

Patients Receiving Migraine Preventive Medications

Monoclonal CGRP antagonists for prevention were not permitted in the lasmiditan trials, use was not permitted within 3 months of enrollment in the ubrogepant trials, and their use is not specifically mentioned in the rimegepant trials. We identified two subgroup analyses that evaluated patients on migraine preventive medications in the trials of lasmiditan and rimegepant (Loo 2019 and Dodick 2019).

Loo 2019 was a pooled analysis of the Phase III trials of lasmiditan (SAMURAI and SPARTAN). The two RCTs allowed patients to continue migraine preventives as long as doses were stable for three months prior to screening and were unchanged during the study. Approximately 18% of patients were on migraine preventive treatments (n=698). The results of the analysis showed that 200 mg lasmiditan was more effective than placebo in achieving pain freedom at two hours for the subgroup using (OR 3.3; 95%CI: 1.9 to 5.7) and not using (OR 2.3: 95%CI: 1.9 -2.9) migraine preventive medications. There was no significant difference in the benefit of all lasmiditan doses versus placebo between patients using or not using migraine preventives (all interaction p-values >=0.1). Rates of adverse events were also similar for patients using and not using preventive medications.⁹⁷

Dodick 2019 was a pooled analysis of the Phase III trials of rimegepant (Study 301, 302, and 303). In total, approximately 16% of the total patients were using preventive medication (rimegepant n=272, placebo n=275). The results showed rimegepant was more effective than placebo in achieving pain freedom at two hours in the subgroup using (20.6% vs. 10.2; p=0.007) and not using (20% vs. 12.6%; p<0.0001) migraine preventive medications, with no significant difference between the two subgroups. Similar trend was observed for the co-primary outcome (freedom from MBS).⁹⁸

Controversies and Uncertainties

Feedback received during this project recommended only comparing the new drugs to placebo, and to each other, for patients in whom triptans have not been effective, are not tolerated or are contraindicated. However, given the availability of triptans for acute treatment of migraine, we also sought to compare these interventions to triptans for patients who do not adequately respond to non-prescription medications and are eligible to use triptans.

We identified 10 RCTs (3 for lasmiditan, 4 for rimegepant and 3 for ubrogepant) comparing the interventions to placebo, but we found only one head-to-head trial of one of the interventions versus a triptan (rimegepant vs sumatriptan). There was no study directly comparing the interventions to each other. Since head-to-head data were generally lacking for the comparisons

between agents, indirect quantitative methods (network meta-analyses) were used. These indirect techniques necessarily have more uncertainty than had the therapies been compared directly.

Patient and patient advocates were concerned that the primary outcomes in the RCTs did not fully reflect the potential benefits of these new therapies. We reported on primary efficacy and side effects of treatment at two hours after initial study medication. After two hours, there were important differences among trial protocols for use of rescue medications and additional study medication dosing (both blinded and open label). Though censoring patients who use additional treatments after 2 hours maintains the placebo-controlled nature of the study, reported outcomes over time reflect only a fraction of the entire study population. In addition, most of the RCTs did not present data on what happened to patients who had pain freedom or relief at two hours but then had a subsequent recurrence of pain, or on the time to pain freedom or relief in patients who did not achieve that outcome at two hours. Though we used best available data for outcomes after 2 hours, potentially important differences in efficacy between medications could be missed.

The RCTs present data on efficacy of treatment for a single migraine attack. There is uncertainty about efficacy over time when these medications are used for repeated attacks over the course of a year or longer. Since migraine can impact quality of life for those with frequent, severe and unpredictable attacks, it is uncertain if these new therapies may favorably impact quality of life measures and work and productivity outcomes over time. Data were also limited for subgroups of interest, including patients not responding to triptans, patients intolerant of triptans, and patients taking CGRP monoclonal antagonists for prevention.

Interest in new therapies for acute treatment of migraine are driven in part by data showing low rates of use of triptans among migraine patients, reflecting lack of effectiveness or intolerance. The medications studied had different rates and types of side effects. It is uncertain how differing rates of side effects will affect patient use and satisfaction over time. Single administration RCTs do not provide useful information for understanding this.

Although triptans are considered to have safety concerns related to vasoconstrictive effects and, when used with certain other medications such as SSRIs, carry a risk of serotonin syndrome, decades of use have suggested that these complications may be extremely infrequent in clinical practice. In contrast, the newer agents are touted as potentially safer, but we have much less clinical information to demonstrate long-term safety at this time.

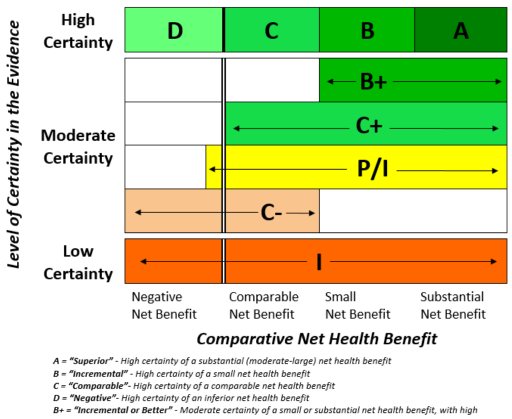
The effect of the newer therapies on migraine frequency over time is uncertain. We heard from multiple stakeholders that decreasing migraine frequency may be an important benefit of these therapies. However, as discussed above, we do not consider it proven that the observed decrease in migraine frequency is due to the treatments. Additionally, it is unknown whether medication overuse headache can occur with these treatments and, if so, whether this occurs more or less frequently than with triptans.

Though migraine is associated with other comorbid conditions and death, it is not known if more effective medications to treat acute migraine episodes may decrease these -longer-term risks.

Because of limitations of existing therapies, there are many individuals in whom no effective, reliable treatment is available. It is hoped that having more treatments for migraine can reduce use of opioids and thus the risk for opioid misuse. Data on this are not yet available.

3.4 Summary and Comment

Figure 3.2. ICER Evidence Rating Matrix



Comparative Clinical Effectiveness

B+ = "incremental or Better" - Moderate certainty of a small or substantial net nealth benefit, with high certainty of at least a small net health benefit
C+ = "Comparable or Better" - Moderate certainty of a comparable, small, or substantial net health benefit, with high certainty of at least a comparable net health benefit

P/I = "Promising but Inconclusive" - Moderate certainty of a comparable, small, or substantial net health benefit, and a small (but nonzero) likelihood of a negative net health benefit

C- = "Comparable or Inferior" - Moderate certainty that the point estimate for comparative net health benefit is either comparable or inferior

I = "Insufficient" - Any situation in which the level of certainty in the evidence is low

Lasmiditan, Rimegepant, and Ubrogepant Versus No Additional Migraine-Specific Acute Treatment (Placebo) or Triptans (Sumatriptan and Eletriptan)

Results from clinical trials and from our NMAs suggest that lasmiditan, rimegepant and ubrogepant decrease symptoms of migraine attacks and improve function compared to placebo. Few harms were seen in the single-dose trials of lasmiditan, rimegepant and ubrogepant. However, lasmiditan showed a higher incidence of CNS related AEs (e.g., dizziness, somnolence, paresthesia) in the clinical trials. Below, we provide summary of the evidence for each drug.

Lasmiditan

- Efficacy (RCTs): Results from single-dose clinical trials suggest a greater proportion of patients achieved freedom from pain (OR 1.5-2.6), relief from pain (OR 1.7-2.5), freedom from MBS (OR 1.4-1.9), and ability to function normally (OR 1.7) at two hours post dose, as well as sustained freedom from pain at 24-and 48-hours (OR 1.3 -2.8) with lasmiditan compared with placebo.
- Efficacy (NMA): Results suggest a higher proportion of patients on lasmiditan achieved pain freedom (OR 1.43) and pain relief (OR 1.15-1.16) at two hours compared to rimegepant and ubrogepant, however, these were not statistically significant. Compared to triptans, a lesser proportion of patients on lasmiditan achieved freedom from pain (OR 0.54) and relief from pain (OR 0.61) at two hours post dose versus eletriptan; the results versus sumatriptan followed the same trend but were not statistically significant.
- Safety: Lasmiditan showed a higher incidence of TEAE compared to placebo in single-dose trials, although the majority were mild or moderate in intensity. Specifically, there was a higher incidence of CNS related AEs, with dizziness the most common. NMA results suggest a higher incidence of TEAE compared to rimegepant, ubrogepant and triptans. In the ongoing 12-month extension study, 12.8% of patients discontinued the trial due to adverse events.

Rimegepant

- Efficacy (RCTs): Results from single-dose clinical trials suggest a greater proportion of patients achieved freedom from pain (OR 1.4-2.2), relief from pain (OR 1.5-1.9), freedom from MBS (OR 1.5-1.8), and ability to function normally (OR 1.7) at two hours post dose, as well as sustained freedom from pain at 24- and 48-hours (OR 1.7-3.2) with rimegepant compared with placebo.
- Efficacy (NMA): Results suggest no significant differences between rimegepant compared to ubrogepant (OR 1.00) and lasmiditan (see above) on pain freedom and pain relief at two hours. However, compared to triptans, lesser proportion of patients achieved freedom from pain (OR 0.38-0.51) and relief from pain (OR 0.52-0.73) at two hours post dose with rimegepant compared with triptans.
- Safety: Rimegepant was generally well tolerated in the single-dose trials, showing a similar rate of TEAE compared to placebo. NMA results also suggest comparable incidence of TEAE relative to ubrogepant and triptans, and a lower incidence compared to lasmiditan. In the ongoing 12-month extension study, 2.7% of patients discontinued the trial due to adverse events.

Ubrogepant

- Efficacy (RCTs): Results from single-dose clinical trials suggest a greater proportion of patients achieved freedom from pain (OR 1.5-2.0), relief from pain (OR 1.7-1.8), freedom from MBS (OR 1.4-1.7), and ability to function normally (OR 1.5) at two hours post dose, as well as sustained freedom from pain at 24-hours (OR 1.6 2.0) with ubrogepant compared with placebo.
- Efficacy (NMA): Results suggest no significant differences between ubrogepant compared to rimegepant (OR 1.00) and lasmiditan (see above) on pain freedom and pain relief at two hours. However, compared to triptans, lesser proportion of patients achieved freedom from pain (OR 0.38-0.52) and relief from pain (OR 0.52-0.73) at two hours post dose, with ubrogepant compared with triptans.
- Safety: Ubrogepant was generally well tolerated in the single-dose trials, showing a similar rate of TEAE compared to placebo. NMA results also suggest comparable incidence of TEAE relative to rimegepant and triptans, and a lower incidence compared to lasmiditan. In the ongoing 12-month extension study, 2.2% of patients discontinued the trial due to adverse events.

Hence, we rated the evidence as follows:

Population 1: For adults (18 years and older) with moderate-severe migraine attacks that have not responded to non-prescription medicines and for whom triptans have not been effective, are not tolerated, or are contraindicated:

• We consider the evidence on lasmiditan, rimegepant and ubrogepant compared to placebo to be "incremental or better" (B+), demonstrating a moderate certainty of a small or substantial health benefit, with a high certainty of at least a small net health benefit.

Population 2: For adults (18 years and older) with migraine attacks that have not responded to non-prescription medicines (and are eligible to use triptans):

Based on the results of the NMAs, rimegepant and ubrogepant appear to be less efficacious than triptans (sumatriptan and eletriptan) but have comparable short-term adverse events. Thus, we consider the evidence on rimegepant and ubrogepant compared to triptans to be "comparable or inferior" (C-), demonstrating moderate certainty that the comparative net health benefit is either comparable or inferior. For lasmiditan, the results of the NMAs suggest it is less efficacious than triptans. However, compared to sumatriptan, the NMAs do not exclude comparable efficacy. In terms of adverse events, the NMA results suggest a higher incidence with lasmiditan compared to triptans. Thus, we consider the evidence on lasmiditan compared to triptans to be "comparable or inferior" (C-).

For all adults with migraine attacks:

 We consider the evidence on rimegepant and ubrogepant to be "comparable" (C), demonstrating a high certainty of a comparable net health benefit. For lasmiditan, the results of the NMAs suggest it may be slightly more efficacious than rimegepant and ubrogepant. However, the NMAs do not exclude comparable efficacy. Patients treated with lasmiditan had more adverse events and more of them discontinued treatment than patients treated with rimegepant or ubrogepant. We believe any possible greater efficacy of lasmiditan is at best balanced by these adverse events and may be outweighed by them, and thus we consider the evidence on lasmiditan compared to rimegepant and ubrogepant to be "comparable or inferior" (C-).

Table 3.21. ICER Ratings on the Comparative Net Health Benefit of Interventions versusComparators

Population	Population 1	Population 2
Interventions	Versus No Treatment	Versus Triptans (sumatriptan and eletriptan)
Lasmiditan	B+	C-
Rimegepant	B+	C-
Ubrogepant	В+	C-

Population 1: Patients with migraine-attacks that have not responded to non-prescription medicines and for whom triptans have not been effective, are not tolerated, or are contraindicated
 Population 2: Patients with migraine-attacks that have not responded to non-prescription medicines (and are eligible to use triptans)

Table 3.22. ICER Ratings on the Comparative Net Health Benefit of Interventions versus EachOther

Population		For All Patients					
Interventions	Versus Lasmiditan	Versus Rimegepant	Versus Ubrogepant				
Lasmiditan		C-	C-				
Rimegepant	C+		С				
Ubrogepant	C+	С					

Note: The table should be read row-to-column. For example, there is moderate certainty that the point estimate for comparative net health benefit of lasmiditan is either comparable or inferior to rimegepant (C-). Conversely, there is moderate certainty of comparable, small or substantial health benefit, with at least a high certainty of at least a comparable health benefit of Rimegepant compared to lasmiditan (C+).

Please note there have been significant changes to this evidence report. Please refer to ICER's final evidence report.

4. Long-Term Cost Effectiveness

4.1 Overview

The primary aim of this economic evaluation was to estimate the cost effectiveness of lasmiditan, rimegepant, and ubrogepant for the acute treatment of migraine using a de novo decision analytic model. The outcomes of interest included the incremental cost per quality-adjusted life year (QALY) gained, life-years gained, equal value of life years gained (evLYG), and cost per hour of migraine pain avoided. An analysis of the incremental cost per evLYG is included in this report to complement the cost per QALY calculations and provide policymakers with a broader view of cost effectiveness. A description of the methodology used to derive the evLYG can be found in Appendix E. Lasmiditan, rimegepant, and ubrogepant were compared with each other and to three comparators in separate analyses representing two distinct populations. For the first comparison, we evaluated lasmiditan, rimegepant, and ubrogepant to each other and to no additional migrainespecific acute treatment. For the purpose of this review, no additional migraine-specific acute treatment was estimated by the placebo arms of the clinical trials, although we recognized that in the real-world, patients may use previously failed or untried over-the-counter and prescription treatments for acute migraine including analgesics. For the second comparison, we evaluated lasmiditan, rimegepant, and ubrogepant to each other and to two triptans: sumatriptan and eletriptan. Sumatriptan was chosen because it is one of the most widely used triptans in clinical practice; and eletriptan, a newer triptan, was shown in a recent network meta-analysis to be one of the most efficacious and well tolerated. Since these new agents under review are all orally available, we focused our comparison of triptans on the oral formulations. All costs and outcomes were discounted at a rate of 3%. For this aim, the base-case analysis was conducted using a health care sector perspective (i.e., focus on direct medical care costs only) and a two-year time horizon. Longer time horizons and productivity gains with treatment were considered in scenario analyses. The model was developed in Microsoft Excel 2016 (Redmond, WA).

4.2 Methods

Model Structure

For the cost-effectiveness analysis, we developed a *de novo* semi-Markov model with time-varying proportions of patients with response to treatment. The model was informed by a network metaanalysis of key clinical trials and prior relevant economic models, systematic literature reviews, and input from diverse stakeholders (patients, advocacy groups, clinicians, payers, researchers, and manufacturers of these agents). The base case used a US health sector perspective. Costs and outcomes were discounted at 3% annually. The model cycle was 48 hours based on the typical duration of clinical trials evaluating acute migraine treatments. The model evaluated two hypothetical cohorts of patients requiring acute treatment for migraine, all being treated with lasmiditan, rimegepant, ubrogepant, or usual care in the first population and all being treated with lasmiditan, rimegepant, ubrogepant, sumatriptan, or eletriptan in the second population.

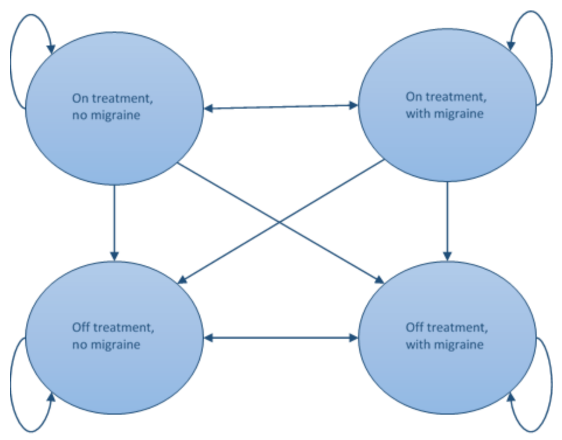
As shown in the model schematic (Figure 4.1), simulated patients entered the model through one of two Markov states, "On treatment, no migraine" or "On treatment, with migraine," according to the average daily probability of having a migraine in the target population (i.e., 4.8 migraines per month, corresponding to a probability of 0.316 migraines in each 48-hour period).

Those patients entering the "On treatment, with migraine" Markov state received the assigned acute initial treatment for migraine (i.e., lasmiditan, rimegepant, ubrogepant, sumatriptan, eletriptan, or usual care). Initial treatment resulted in some proportion of patients achieving complete resolution of migraine pain (pain freedom), an improvement in migraine pain without complete resolution (pain relief), or no improvement in migraine pain at each of four time points: 2, 8, 24, and 48 hours.

Over time, patients were allowed to discontinue treatment due to side effects or insufficient effectiveness. For patients who discontinued treatment due to side effects, 12-month treatment-specific discontinuation rates were used. For patients who discontinued treatment due to insufficient effectiveness, the proportion of patients remaining in the "On treatment, with migraine" Markov state who received benefit from therapy increased, to maintain the *total* proportion of patients remaining in the "On treatment over time. Since the absolute effectiveness gains of patients remaining in the "On treatment, with migraine" Markov state is not known, this estimate was subjected to a modifier, that was set at 50% benefit for the base case.

Patients who discontinued treatment transitioned to the "Off treatment, no migraine" or "Off treatment, with migraine" Markov states according to the observed probability of discontinuation derived from Brandes et al.⁵⁶ The model was designed with the assumption that patients who discontinued treatment would not return to either of the "On treatment, no migraine" or "On treatment, with migraine" Markov states. Patients transitioned between the "Off treatment, no migraine" and "Off treatment, with migraine" states according to the average probability of having a migraine every 48 hours, similar to those on the initial treatment.





Target Population

The population of focus for the economic evaluation was the prevalent cohort of individuals in the United States (US) aged 18 years and over experiencing migraines requiring acute treatment, with or without aura as specified by the International Classification of Headache Disorders (ICHD) 3 diagnostic criteria.⁹⁹ Two separate cohorts of patients were evaluated using different comparators. The first cohort was comprised of patients who had migraine attacks that did not respond to non-prescription medicines and for whom triptans had not been effective, were not tolerated, or were contraindicated. The second cohort was comprised of patients who had migraine attacks that did not respond adequately to non-prescription medicines, such as non-steroidal anti-inflammatory agents. In this cohort, comparisons were made among lasmiditan, rimegepant, and ubrogepant, and two commonly used oral triptans with different effectiveness and cost, sumatriptan and eletriptan, representing a range of triptan medications. The baseline patient characteristics are presented in Table 4.1.

Table 4.1. Base-Case Model Cohort Characteristics

Baseline Characteristics	Value	Source
Mean Age, years (SD)	40.8	Croop 2019 ²⁸
Female, %	86.0	Lipton 2019 ²⁶
Migraine Days per Month at Baseline	4.8	Doty 2019 ¹⁰⁰

Treatment Strategies

Interventions included in the models were lasmiditan 100-200 mg, rimegepant 75 mg, and ubrogepant 50-100 mg. The comparators depended on the population being evaluated. In Population 1 (i.e., patients in whom prior treatment with non-prescription medicines failed and for whom triptans were not effective, were not tolerated, or were contraindicated), the interventions were compared with each other and with usual care, represented by the placebo arm from clinical trials. In Population 2, the interventions were compared with each other and with sumatriptan 50-100 mg and eletriptan 40 mg.

Key Model Characteristics and Assumptions

The model required several assumptions. Key model assumptions and rationale for the assumptions are presented in table 4.2.

Assumption	Rationale
Mortality is not associated with acute treatment for	There have been no demonstrated mortality benefits
migraine.	with treatment of migraine pain and other symptoms.
Acute treatment of migraine with lasmiditan,	Studies evaluating new migraine therapies were either
rimegepant, ubrogepant, and triptans does not affect	short-term single episode studies or non-controlled
migraine frequency.	open label studies and were not designed to
	demonstrate changes in migraine frequency with
	treatment. Longer-term, uncontrolled, open-label
	studies suffer from a possible placebo effect and a
	high likelihood that regression to the mean may affect
	the study's results. Should stronger evidence suggest
	that migraine frequency and/or characteristics are
	modified with acute treatments for migraine, this
	assumption will be reevaluated.
A two-year time horizon is sufficient to estimate the	Compared with many other chronic conditions
cost effectiveness of acute treatments for migraine.	modeled using Markov models, migraine onset is
	rapid, and resolution occurs quickly. Since costs are
	incurred with each treatment and benefits are
	observed immediately, we believe that a two-year
	time horizon will be sufficient to estimate a stable
	incremental cost-effectiveness ratio for the acute

Table 4.2. Key Model Assumptions

Assumption	Rationale
	treatment of migraine. We will test this assumption by extending the time horizon to 5 years and determining whether the cost effectiveness of therapies appreciably change.
Patients who have discontinued treatment received some other medication with a response similar to those in the placebo arm from clinical trials.	This analysis was intended to evaluate the cost effectiveness of new acute treatments for migraine. Since there are a variety of medications available for acute migraine, with varying effectiveness and cost, that could be used in the event that patients discontinued one of the new acute treatments, there was no single alternative available for the model. The discontinuation rates of the new treatments appear to be relatively similar from single arm continuation safety studies, so the impact of this assumption is expected to be minimal. In addition, the cost and effectiveness of the acute treatment used for those who discontinue lasmiditan, rimegepant, and ubrogepant will be subjected to a two-way sensitivity analysis to determine the potential impact of this assumption on the cost-effectiveness results.
Patients receiving no benefit from treatment discontinued the medication in the first year of treatment only. There was no discontinuation for lack of effectiveness in the second year of the model.	Data describing treatment discontinuation due to lack of effect was obtained from a study in which follow up lasted for 12 months. ⁵⁶ It is unlikely that the majority of patients receiving no or suboptimal benefit would continue taking a medication beyond 12 months.
Patients who did not respond to acute treatments for migraine were assumed to have moderate or severe pain, in proportion to what was observed at baseline.	Sufficiently detailed data evaluating those who did not respond was not uniformly available from clinical trials. This assumption was necessary to assign utility values to those who did not respond to therapy.
Adverse drug events last for 8 hours.	Symptoms of drowsiness, dizziness, fatigue, and paresthesia were more frequent than placebo with certain acute treatments of migraine. The mean time that patients suffered from these treatment-emergent adverse events was not described in studies. In order to determine QALYs lost due to treatment-emergent adverse events, a duration of the event had to be assumed.

Assumption	Rationale
Discontinuations due to "patient request" in the	Discontinuation probability and reasons for
GLADIATOR study represent discontinuations due to	discontinuation are not reported for acute treatments
lack of treatment effect. ⁵⁶ Given the similarity in	for acute migraine. This study described
treatment response among lasmiditan, rimegepant,	discontinuation reasons but did not include a category
and ubrogepant, we assumed that treatment	stating whether discontinuation was for lack of
discontinuation due to lack of effectiveness would be	effectiveness. Given the other categories for
similar.	discontinuation, this category of "patient request" was
	likely to represent patients who did not derive benefit
	from treatment. Assuming patients would continue
	treatment, even when it wasn't effective, would bias
	the analysis against lasmiditan, rimegepant, and
	ubrogepant, when compared to usual care.
If a migraine treatment resulted in migraine pain of	The impact of migraine on productivity is important to
"no pain" or "mild pain" at 2 hours, a person would	patients. However, clinical trials did not evaluate work
be able to work.	productivity. Studies that have evaluated work
	productivity have assessed the impact of migraine on
	productivity (primarily absenteeism) but have not
	assessed the impact of treatment and time to pain
	and/or symptom relief on productivity. This
	assumption was necessary to apply results of
	productivity studies in migraine patients to this model
	for the scenario analysis evaluating a modified societal
	perspective.

Model Inputs

Clinical Inputs

Short-term clinical inputs for the effectiveness of acute treatments for migraine and the comparators were derived from a network meta-analysis of clinical trials evaluating lasmiditan, rimegepant, ubrogepant, sumatriptan, and eletriptan compared with placebo and with each other, where such studies existed.

Clinical Probabilities/Response to Treatment

The decision model was evaluated over a two-year time horizon with 48-hour cycles. The probability of having a migraine in each cycle was estimated using the number of migraine days per month from patients enrolled in clinical trials. Within each cycle, the proportions of patients with severe, moderate, mild, or no pain were evaluated at baseline, 2, 8, 24, and 48 hours using data from clinical trials. Patients **without** migraine had no pain for the entire 48-hour cycle. Patients **with** migraine started in severe or moderate pain, derived from the average proportions of patients with moderate or severe pain at baseline from clinical trials.

Two-hour response to acute treatments for migraine was estimated using data directly from clinical trials included in a network meta-analysis described earlier in this report. The proportion of patients who were pain free in clinical trials were considered to have "no pain" at the 2-hour time point. Since the proportion of patients who had pain relief in clinical trials included those who were pain free, the proportion who were pain free was subtracted from those with pain relief to estimate the proportion of patients with "mild pain" at 2 hours and for all subsequent time points. Those who did not have a response in clinical trials were assumed to have moderate or severe pain, in proportion to what was observed at baseline.

In clinical trials evaluating lasmiditan, rimegepant, and ubrogepant, some patients who responded at two hours subsequently lost response to treatment between 2 and 24 hours. The proportion of patients who did not lose response at 24 hours were considered to have maintained response over that time. For the proportion of patients who did lose response as estimated in the network meta-analysis, we assumed the maximal proportion lost response at eight hours with a linear loss from two to eight hours. After eight hours, we assumed that patients regained response such that at 24 hours the patients who had lost response had the same response rate as in the placebo response from Dodick.¹⁰¹ This return of response was assumed to be linear from eight to 24 hours. All patients responding at 2 hours were also assumed to have response at 48 hours.

Patients who did not respond at two hours were similarly assumed to achieve response at eight and 24 hours as per the placebo response from Dodick,¹⁰¹ with linear achievement of response between two and eight hours, and then a separate linear response between eight and 24 hours. Response at 48 hours was similarly calculated by adding all two-hour responders to the placebo response for non-responders at two hours. The proportion of patients with moderate or severe migraine pain was calculated by multiplying the proportion of non-responders (i.e., 1 - responders) at 2, 8, 24, and 48 hours by the proportion of patients with "moderate pain" and/or "severe pain" at baseline.

Level of Migraine						
Pain at	Lasmiditan	Rimegepant	Ubrogepant	Sumatriptan	Eletriptan	Usual Care
Timepoints, %						
Baseline (0h), %						
None	0.0	0.0	0.0	0.0	0.0	0.0
Mild	0.0	0.0	0.0	0.0	0.0	0.0
Moderate	66.6	66.6	66.6	66.6	66.6	66.6
Severe	33.4	33.4	33.4	33.4	33.4	33.4
2h, %						
None	28.0	21.0	21.0	35.0	42.0	11.0
Mild	30.0	33.0	33.0	27.0	27.0	24.0
Moderate	27.9	30.6	30.6	25.3	20.6	43.3
Severe	14.0	15.4	15.4	12.7	10.3	21.7
8h, %						
None	59.5	58.5	58.0	61.0	62.0	53.5
Mild	29.9	31.2	31.4	29.1	29.9	32.8
Moderate	7.1	6.8	7.1	6.6	5.4	9.1
Severe	3.5	3.4	3.5	3.3	2.7	4.6
24h, %						
None	74.3	71.9	71.9	76.8	79.3	68.3
Mild	19.0	20.9	20.9	17.2	15.8	21.5
Moderate	4.4	4.8	4.8	4.0	3.2	6.8
Severe	2.2	2.4	2.4	2.0	1.6	3.4
48h						
None	81.8	80.0	80.0	83.6	85.3	77.4
Mild	12.4	13.6	13.7	11.2	10.4	13.6
Moderate	3.8	4.2	4.2	3.5	2.8	5.9
Severe	1.9	2.1	2.1	1.7	1.4	3.0

Table 4.3. Treatment Response Used in Model

The probability of having migraine-related provider office visits or of being admitted to the emergency department or hospital were determined for patients with persistent pain, derived from Silberstein et al.¹⁰² To estimate the probability of having a migraine-related provider office, emergency, or hospital visit during a migraine, these rates were divided by the baseline number of migraines with severe headache pain per year. In the model, provider office, emergency department, and hospital visits were assumed to occur only in patients who had migraine pain lasting 12 hours. A ratio of having moderate or severe pain at 12 hours with a specific treatment compared with placebo was used to adjust the likelihood of requiring a provider office, emergency department, or hospital visit due to migraine. Therefore, more effective therapies reducing headache pain at 12 hours resulted in fewer health care visits than did less effective therapies.

Model Input	12-Month Value	Per Migraine Probability	Source
Mean Number of Migraine-Related	2.2	3.8%	Silberstein
Health Care Provider Visits			2018 ¹⁰²
Mean Number of Migraine-Related	1.2	2.1%	
Emergency Department Visits			
Mean Number of Migraine-Related	0.4	0.7%	
Hospitalizations			

Table 4.4. Non-Treatment Dependent Values Used to Calculate Model Event Probabilities

Discontinuation

Treatment discontinuation probabilities due to lack of response were derived from the GLADIATOR long-term safety study of lasmiditan.⁵⁶ We assumed that "patient request" referred to those patients who discontinued the medication for lack of effect. Discontinuation was primarily due to "patient request" (21.8%) and adverse events (12.8%). Long-term data on treatment discontinuation due to lack of effectiveness were not available for other treatments. Since lasmiditan, rimegepant, and ubrogepant all show similar effectiveness, we assumed that discontinuation for lack of effectiveness would also be similar among all treatments. We also assumed that discontinuation of triptans due to lack of effectiveness was the same as that of the newer acute treatments for migraine. Discontinuation due to lack of effectiveness was set to 0% after one year.

Treatment-specific discontinuation rates due to adverse drug events were obtained from longer term observational studies.^{56,88,89} We assumed that adverse events were not related to patient response. Therefore, patients discontinuing treatment due to an adverse event were proportionally removed from all response categories (i.e. pain free, pain relief, and non-responders). Discontinuation due to adverse drug events was set to 0% after two years in the sensitivity analysis evaluating longer time horizons.

<u>Mortality</u>

Therapies for migraine have not demonstrated differences in mortality, nor has a mechanism for differential survival with the current treatments been proposed. In addition, the model used a short time horizon of two years to generate the incremental cost-effectiveness estimates for the new therapies. Given the relatively young age of the population being evaluated and associated low mortality rate, mortality was not included in the model.

<u>Adverse Events</u>

All adverse events occurring in at least 5% of patients, and their disutilities, were included in the analysis. In addition, fatigue was included even when it did not reach an incidence of 3%, as it had a larger impact on patient utility. Adverse events were assumed to last for 8 hours. Discontinuation due to adverse events was also included in the analysis.

Adverse Event	Drug	Frequency, %	Disutility	References
Drowsiness	Lasmiditan	5.5	-0.028	Krege 2019 ¹⁰³ Matza 2019 ¹⁰⁴
Dizziness	Lasmiditan	14.7	-0.021	Krege 2019 ¹⁰³ Matza, 2019 ¹⁰⁴
	Lasmiditan	3.8	-0.069	Krege 2019 ¹⁰³ Matza 2019 ¹⁰⁴
Fatigue	Sumatriptan	3.0	-0.069	Imitrex FDA label ¹⁰⁵ Matza, 2019 ¹⁰⁴
	Eletriptan	10.0	-0.069	Relpax FDA label ¹⁰⁶ Matza 2019 ¹⁰⁴
	Lasmiditan	5.7	-0.013	Krege 2019 ¹⁰³ Matza 2019 ¹⁰⁴
Paresthesia	Sumatriptan	5.0	-0.013	Imitrex FDA label ¹⁰⁵ Matza, 2019 ¹⁰⁴
	Eletriptan	4.0	-0.013	Relpax FDA label ¹⁰⁶ Matza 2019 ¹⁰⁴

Health State Utilities

Table 4.6 shows health state utility values used in the model. Utilities were derived from published literature that estimated migraine-specific utility values using the EQ-5D and stratified by the severity of the migraine. For patients without migraine, a utility associated with "no pain" derived from Xu et al. was used.¹⁰⁷ For patients with migraine, we first estimated the proportion of patients with no, mild, moderate, or severe pain at 0 (baseline), 2, 8, 24, and 48 hours. The trapezoidal method for estimating area under the curve was then used to derive the proportion of patients with no, mild, moderate, or severe pain between 0-2 hours, 2-8 hours, 8-24 hours, and 24-48 hours. Utility estimates from Xu et al., shown in Table 4.5, were applied to these proportions for the appropriate amount of time (e.g., 16 hours for the 8-24-hour time period).¹⁰⁷

Disutilities of -0.5 were assumed for those patients who were hospitalized or required an ED visit. Hospitalizations were assumed to last for 2 days, ED visits for 1 day. We did not include a disutility score for patients suffering from nausea and/or vomiting, photophobia, or phonophobia due to lack of data.

Disutility of adverse events were estimated from the rate of the events, the associated disutility for the event, and an assumed duration of eight hours. The disutility values are noted in the table included in the above section on adverse events.

	Migra	ine-Specific Utility	Value	
Migraine Symptom	Mean Value	95% CI	Method	Source
Severe Pain	0.440	(0.374, 0.502)	EQ-5D	Xu 2011 ¹⁰⁷
Moderate Pain	0.773	(0.755, 0.789)	EQ-5D	Xu 2011 ¹⁰⁷
Mild Pain	0.835	(0.790, 0.883)	EQ-5D	Xu 2011 ¹⁰⁷
Pain free	0.959	(0.896, 0.967)	EQ-5D	Xu 2011 ¹⁰⁷
Nausea/vomiting	Estimate not found in literature search	Estimate not found in literature search	Estimate not found in literature search	
Photophobia	Estimate not found in literature search	Estimate not found in literature search	Estimate not found in literature search	
Phonophobia	Estimate not found in literature search	Estimate not found in literature search	Estimate not found in literature search	
Hospitalization	-0.5 (for 2 days)			Assumed
Emergency Department Visit	-0.5 (for 1 day)			Assumed
Adverse Events	-0.013—0.069		Time Trade Off	Matza 2019 ¹⁰⁴

Table 4.6. Utility Values for Health States

Economic Inputs

Drug Utilization

Drug utilization for acute treatments for migraine evaluated in this model, used to determine costs, are shown in Table 4.7. When available, the approved indication dosage will be used to model drug costs.

Generic Name	Lasmiditan	Rimegepant	Ubrogepant	Sumatriptan	Eletriptan	Sources
Brand name	Reyvow	Investigational	Ubrelvy	lmitrex, others	Relpax	
Manufacturer	Eli Lilly	Biohaven	Allergan			
Route of Administration	Oral	Oral	Oral	Oral	Oral	
Dosing	50 mg, 100 mg, or 200 mg orally; No more than one dose in 24 hours.	Approved dosing information not available	50-100 mg orally; may repeat after 2 hours; maximum dose is 200 mg/24 hours	50-100 mg orally; may repeat after 2 hours; maximum dose: 200 mg/24 hours	40 mg; may repeat after 2 hours; maximum dose: 80 mg/24 hours	Reyvow prescribing information Ubrelvy prescribing information Micromedex online

Table 4.7. Treatment Regimen Recommended Dosage

Drug Costs

At the time of publishing this report, the prices for lasmiditan, rimegepant, and ubrogepant were not available. We therefore estimated the prices of lasmiditan, rimegepant, and ubrogepant for the model based on an opinion article that estimated that ubrogepant would have a 20% premium to branded Imitrex.⁵⁷ We applied the same premium to lasmiditan and rimegepant. All estimates generated in the model used these placeholder prices. Costs for sumatriptan and eletriptan were derived using wholesale acquisition cost (WAC) from Redbook and shown in Table 4.8.¹⁰⁸ Aligning with the <u>ICER Reference Case (http://icer-review.org/wp-</u>

<u>content/uploads/2018/07/ICER Reference Case July-2018.pdf</u>), we have used the WAC to price these treatments, as they are currently available as generic medications. Costs for treatments for the usual care arm were estimated using a prevalent mix of treatments, estimated from Ford et al , and applying WAC prices from Redbook.^{108,109} Since triptans were not indicated for Population 1 and were the comparators for Population 2, we removed triptans from the prevalent mix reported and adjusted the remaining treatments accordingly. After the removal of triptans, the resulting mix of treatments and proportion of patients in which they were used were as follows: butalbital/caffeine/acetaminophen (11.3%), ibuprofen (38.2%), naproxen (32.1%), opioids (28.3%).

Table	4.8.	Drug	Cost	per	Dose
-------	------	------	------	-----	------

Drug	WAC	Notes	Source
Lasmiditan	n/a	20% premium pricing above	Kish 2018 ⁵⁷
	(Used \$78.38)	Imitrex	Micromedex ¹⁰⁸
Dimeronent	n/a	20% premium pricing above	Kish 2018 ⁵⁷
Rimegepant	(Used \$78.38)	Imitrex	Micromedex ¹⁰⁸
Librogonant	n/a	20% premium pricing above	Kish 2018 ⁵⁷
Ubrogepant	(Used \$78.38)	Imitrex	Micromedex ¹⁰⁸
Sumatriptan,			
Oral tablets	\$1.04		Redbook Online from Micromedex ¹⁰⁸
50 mg	\$1.04		Reabook Online from Micromedex
100 mg			
Eletriptan			Redbook Online from Micromedex ¹⁰⁸
40 mg	\$11.95		Reabook Online from Micromedex-
Usual Care	¢4 01		Ford 2017 ¹⁰⁹
(mix)	\$4.81		Micromedex ¹⁰⁸

Non-Drug Health Care Costs

In the model, the non-drug health care costs for the acute treatment of migraine included only those costs demonstrated to be associated with treatment. Costs associated with provider office visits, emergency department visits, and hospitalizations were included, as a rapid decrease in pain and other migraine symptoms were likely to be impacted by improved migraine pain. To estimate the cost of hospitalization, the most recently available year (2016) mean cost of hospitalizations for ICD-10 codes G43.xxx were obtained from the online Health Care Utilization Project (HCUP.net).¹¹⁰ The cost of emergency department visits was estimated by obtaining the total ED facility and doctor's fees from the Medicare Expenditure Panel Survey online tool (MEPS.AHRQ.gov).¹¹¹ The 2019 Center for Medicare & Medicaid Services physician fee schedule was used to estimate the cost of a provider office visit. We assumed a level 2 physician office visit (HCPCS code 99212) for a migraine-related visit. All costs were inflated to 2019 US dollars using the Health Care component of the Bureau of Economic Analysis Personal Consumption Expenditures Price Index (PCE) as per ICER's Reference Case. These costs are shown in Table 4.4.

We included the potential impact of therapies for migraine on productivity losses in a scenario analysis. We used estimates for productivity losses resulting from migraine derived from Mesalli et al. 2016, which captures presenteeism productivity loss, days missed, and losses in housework conducted for full-time employees, part-time employees, and those with other employment status.⁷ The total productivity loss costs for acute migraines were estimated to be \$245 per month. We used an assumption that if a migraine responded to treatment quickly (i.e. within 2 hours), people would be able to begin, continue, or return to work. Productivity gains due to effective treatment were estimated by applying a calculated benefit per migraine at 2 hours to all patients with no pain or mild pain.

Sensitivity Analyses

One-way sensitivity analyses were conducted on all model inputs to identify the impact of parameter uncertainty and key drivers of model outcomes. Probabilistic sensitivity analyses were performed by jointly varying sensitive model parameters over 5,000 simulations and calculating 95% credible range estimates for each model outcome based on the results.

Scenario Analyses

We employed four scenario analyses. In the first scenario analysis, we evaluated the impact of productivity gains added to the base-case analysis, using a modified societal perspective. In this analysis, patients with mild or no pain at 2 hours were assumed able to continue working.

The second scenario analysis evaluated the impact of increased effectiveness for rimegepant and ubrogepant after 2 hours. There is limited evidence that an increasing number of patients taking rimegepant and ubrogepant may benefit from active treatment compared with placebo after 2 hours.^{26,30} These data were not included in the base case, since the evidence came from exploratory analyses that censored patients who received rescue medications. There were differential rates of censoring between active treatments and placebo in both study reports evaluating rimegepant and ubrogepant, which could lead to a bias. We applied a rate ratio between active treatment and placebo to those patients who did not receive benefit at two hours to replicate the possible benefit observed in these exploratory analyses.

The third scenario analysis evaluated the impact of decreased migraine frequency in the population over time. Evidence from long-term safety trials suggests that migraine frequency may have decreased over time.⁵⁶ One non-controlled, observational study showed that in patients who were observed for one year, migraine frequency decreased from a mean of approximately 6 migraines per quarter to 3.7 migraines per quarter. Since a reduction in migraine frequency would have an impact on medication costs, we conducted a scenario analysis to evaluate the impact of decreasing migraine frequency on cost effectiveness and to generate inputs for the budget impact analysis.

The fourth scenario analysis extended the time horizon to five years to assess whether a longer timeline impacted the cost effectiveness of treatments.

Threshold Analyses

Threshold analyses were conducted for population 1 to determine the price required to result in willingness-to-pay thresholds of \$50,000, \$100,000, and \$150,000 per QALY gained.

Model Validation

We have and will use several approaches to validate the model. First, we provided preliminary methods and results to manufacturers, patient groups, and clinical experts. Based on feedback from these groups, we refined model structure and data inputs used in the model. We performed model verification for model calculations using internal reviewers and varied model input parameters to evaluate face validity of changes in results. Finally, we provided the manufacturers of rimegepant, ubrogepant and lasmiditan an opportunity to review and comment on the most recent draft of the model base case during the comment period for the draft report.

4.3 Results

Base-Case Results

Since the prices for lasmiditan, rimegepant, and ubrogepant were not available at the time of publishing this draft document, we used prices in the model that were 20% above those for branded Imitrex for all reported results and sensitivity analyses. The total discounted lifetime costs, QALYs, LY, evLYG, and mean hours of migraine pain per attack are shown for lasmiditan, rimegepant, ubrogepant, sumatriptan, eletriptan, and usual care in Table 4.9.

Table 4.9. Base-Case Results for Lasmiditan, Rimegepant, Ubrogepant, Sumatriptan, Eletriptan,and Usual Care*

Treatment	Drug Cost (per year)**	Total Cost**	QALYs	Life Years	evLYG	Hours of Pain
Lasmiditan	\$3,970	\$13,640	1.8252	1.95	1.8252	1,743
Rimegepant	\$3,970	\$14,500	1.8222	1.95	1.8222	1,870
Ubrogepant	\$3,970	\$14,510	1.8221	1.95	1.8221	1,876
Sumatriptan	\$50	\$6,630	1.8264	1.95	1.8264	1,611
Eletriptan	\$590	\$6,790	1.8293	1.95	1.8293	1,484
Usual Care	\$0	\$10,050	1.8142	1.95	1.8142	2,100

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

**Drug costs per year were calculated without accounting for discontinuation of the drug. Total costs take into account discontinuation and costs of alternative treatments.

QALY: quality-adjust life year; LY: life year; evLYG: equal value of life years gained

Cost per QALY gained for the primary comparisons are shown in Table 4.10. When evaluating the use of lasmiditan, rimegepant, and ubrogepant using the place-holder prices in Population 1, the ICERs for lasmiditan, rimegepant, and ubrogepant compared with usual care were \$327,700,

\$559,500, and \$569,600 per QALY gained, respectively. When compared with each other and at the placeholder prices used in the model and point estimates derived from the network meta-analyses, lasmiditan dominated rimegepant and ubrogepant, being more effective and less costly. However, there was significant overlap in the confidence intervals for lasmiditan and the point estimates for rimegepant and ubrogepant. Rimegepant and ubrogepant had nearly identical total costs, QALYs, and cost effectiveness. The incremental cost effectiveness of lasmiditan, rimegepant, and ubrogepant, will be dependent on the actual pricing of the three therapies. In Population 2, both sumatriptan and eletriptan produced higher QALYs at a lower total cost, and therefore dominated lasmiditan, rimegepant, and ubrogepant. As there is no mortality effect in the model, cost per LY gained is not relevant, and the cost per evLYG is essentially the same as the cost per QALY gained.

Treatment	Comparator	Cost per QALY Gained	Cost per Hour of Pain Avoided
	Popula	ation 1	
Lasmiditan	Usual Care	\$327,700	\$10.10
Rimegepant	Usual Care	\$559,500	\$19.41
Ubrogepant	Usual Care	\$569,600	\$19.41
	Popula	ation 2	
Lasmiditan	Sumatriptan	Dominated	Dominated
Rimegepant	Sumatriptan	Dominated	Dominated
Ubrogepant	Sumatriptan	Dominated	Dominated
Lasmiditan	Eletriptan	Dominated	Dominated
Rimegepant	Eletriptan	Dominated	Dominated
Ubrogepant	Eletriptan	Dominated	Dominated

Table 4.10. Incremental Cost-Effectiveness Ratios for the Base Case *

QALY: quality-adjusted life years

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant

Differences from Draft Evidence Report

Several changes and corrections were responsible for revisions between the results presented in the Draft Evidence Report and this report:

• Based on stakeholder input, we used the same placeholder price for lasmiditan, rimegepant, and ubrogepant rather than assuming a slightly lower price for lasmiditan. The actual prices are still not known.

- Based on input from multiple stakeholders, the NMA was changed to adjust for placebo response rates. This improved the effects of all three agents on "pain freedom" and "pain response" at 2 hours, however this had the greatest beneficial change for lasmiditan.
- Based on stakeholder review, an error was corrected that had led to underestimating QALYs gained from avoiding emergency department and hospital visits.
- We corrected an error that resulted in overestimating the number of hours of pain experienced by patients.

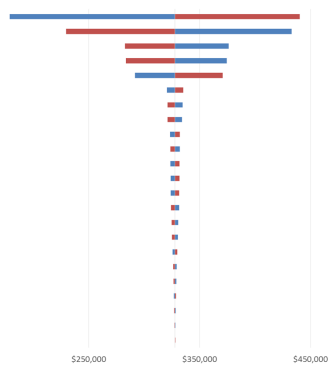
Sensitivity Analysis Results

The model was sensitive to many of the model inputs. However, in one-way sensitivity analysis, none of the individual model inputs being varied resulted in an ICER of below \$150,000 per QALY gained when using the assumed placeholder costs.

Figures 4.2. Tornado Diagrams for One-Way Sensitivity Analyses of Lasmiditan, Rimegepant, and Ubrogepant Compared with Usual Care (Placebo)

Figure 4.2a. Model Probabilities, Lasmiditan versus Usual Care (Placebo)

	Base-Case	Min	Max
Migraine frequency (new migraines per month)	4.8	3.0	8.0
Probability of hospitalizations	0.7%	0.2%	1.2%
Probability of 24h pain relief, LAS	93.4%	92.9%	94.0%
Probability of ED visits	2.1%	1.3%	2.9%
Probability of 24h pain relief, UC	89.8%	89.1%	90.5%
Probability of 24h pain free, UC	68.3%	67.2%	69.3%
Probability of 24h pain free, LAS	74.3%	73.4%	75.3%
Probability of improved effectiveness in pts who don't discontinue	50.0%	0.0%	100.0%
Probability of 8h pain free, UC	53.5%	52.4%	54.6%
Probability of 8h pain free LAS	59.5%	58.4%	60.6%
Probability of 8h pain relief, UC	86.3%	85.5%	87.1%
Probability of 48h pain free, UC	77.5%	76.6%	78.4%
Probability of 48h pain relief, UC	91.1%	90.4%	91.7%
Probability of 48h pain free, LAS	81.8%	80.9%	82.7%
Probability of 48h pain relief, LAS	94.2%	93.7%	94.8%
Probability of 8h pain relief, LAS	89.4%	88.7%	90.1%
Probability of 2h pain relief, UC	35.0%	33.9%	36.1%
Probability of 2h pain free, LAS	28.0%	27.0%	29.0%
Probability of provider visits due to nausea/vomiting	3.8%	2.7%	4.9%
Probability of 2h pain free, UC	11.0%	10.3%	11.7%
Probability of 2h pain relief, LAS	58.0%	56.9%	59.1%
Probability of discontinuing due to lack of effect, LAS	21.8%	20.9%	22.7%
Discontinuation rate due to adverse events, LAS	12.8%	12.0%	13.6%

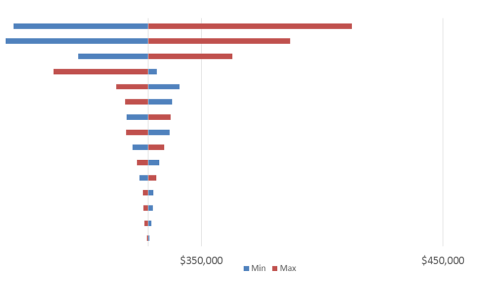


Min Max

LAS: Lasmiditan; UC: Usual Care; ED: emergency department

Figure 4.2b. Model Costs and Utilities, Lasmiditan versus Usual Care (Placebo)	
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	Base-Case	Min	Max
Disutility, ED visit	0.00	-0.01	0.00
Price, LAS	\$78	\$71	\$86
Disutility, hospitalization	0.00	-0.01	0.00
Price, UC	\$5	\$4	\$10
Hospitalization cost per visit	\$6,840	\$6,156	\$7,524
Utility, no migraine	0.96	0.95	0.97
Utility, moderate migraine	0.77	0.75	0.80
ED cost per visit	\$1,555	\$1,400	\$1,711
Utility, severe migraine	0.44	0.41	0.47
Disutility, dizziness	-0.02	-0.03	-0.01
Utility, mild migraine	0.84	0.81	0.86
Disutility, fatigue	-0.07	-0.09	-0.04
Disutility, drowsiness	-0.03	-0.04	-0.01
Disutility, parasthesia	-0.01	-0.02	0.00
Provider office visit cost per visit	\$46	\$41	\$50



LAS: lasmiditan; UC: Usual Care; ED: emergency department

Figure 4.2c. Model Probabilities, Rimegepant versus Usual Care (Placebo)

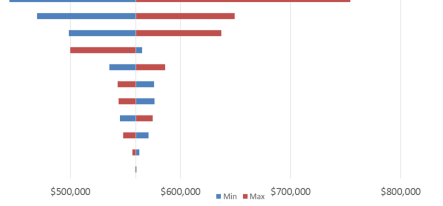
	Base Case	Min	Max
Migraine frequency (new migraines per month)	4.8	3.0	8.0
Probability of hospitalizations	0.7%	0.2%	1.2%
Probability of 24h pain relief, RIM	92.8%	92.2%	93.4%
Probability of 24h pain relief, UC	89.8%	89.1%	90.5%
Probability of ED visits	2.1%	1.3%	2.9%
Probability of improved effectiveness in pts who don't discontinue	50.0%	0.0%	100.0%
Probability of 24h pain free, RIM	71.9%	70.8%	72.9%
Probability of 24h pain free, UC	68.3%	67.2%	69.3%
Probability of 8h pain free RIM	58.5%	57.3%	59.7%
Probability of 8h pain free, UC	53.5%	52.4%	54.6%
Probability of 8h pain relief, UC	86.3%	85.5%	87.1%
Probability of 48h pain free, RIM	80.0%	79.1%	81.0%
Probability of 48h pain free, UC	77.5%	76.6%	78.4%
Probability of 8h pain relief, RIM	89.7%	89.0%	90.5%
Probability of 48h pain relief, UC	91.1%	90.4%	91.7%
Probability of 48h pain relief, RIM	93.7%	93.1%	94.3%
Probability of 2h pain relief, RIM	54.0%	52.8%	55.2%
Probability of 2h pain relief, UC	35.0%	33.9%	36.1%
Probability of 2h pain free, RIM	21.0%	20.0%	22.0%
Probability of 2h pain free, UC	11.0%	10.3%	11.7%
Probability of provider visits due to nausea/vomiting	3.8%	2.7%	4.9%
Probability of discontinuing due to lack of effect, RIM	21.8%	20.9%	22.7%
Discontinuation rate due to adverse events, RIM	2.7%	2.3%	3.1%
			\$35

Min Max

RIM: rimegepant; UC: usual care; ED: emergency department

	Base-Case	Min	Max
Disutility, ED visit	0.00	-0.01	0.00
Price, RIM	\$78	\$71	\$86
Disutility, hospitalization	0.00	-0.01	0.00
Price, UC	\$5	\$4	\$10
Utility, moderate migraine	0.77	0.75	0.80
Hospitalization cost per visit	\$6,840	\$6,156	\$7,524
Utility, no migraine	0.96	0.95	0.97
Utility, severe migraine	0.44	0.41	0.47
ED cost per visit	\$1,555	\$1,400	\$1,711
Utility, mild migraine	0.84	0.81	0.86
Provider office visit cost per visit	\$46	\$41	\$50

Figure 4.2d. Model Costs and Utilities, Rimegepant versus Usual Care (Placebo)



RIM: rimegepant; UC: usual care; ED: emergency department

Figure 4.2e. Model Probabilities, Ubrogepant versus Usual Care (Placebo)

	Base-Case	Min	Max
Migraine frequency (new migraines per month)	4.8	3.0	8.0
Probability of 24h pain relief, UBR	92.8%	91.9%	93.7%
Probability of hospitalizations	0.7%	0.2%	1.2%
Probability of 24h pain relief, UC	89.8%	89.1%	90.5%
Probability of ED visits	2.1%	1.3%	2.9%
Probability of improved effectiveness in pts who don't discontinue	50.0%	0.0%	100.0%
Probability of 24h pain free, UBR	71.9%	70.3%	73.4%
Probability of 24h pain free, UC	68.3%	67.2%	69.3%
Probability of 8h pain free UBR	58.0%	56.3%	59.7%
Probability of 48h pain relief, UBR	80.0%	78.6%	81.4%
Probability of 8h pain relief, UBR	89.4%	88.3%	90.5%
Probability of 48h pain relief, UBR	93.7%	92.8%	94.5%
Probability of 8h pain free, UC	53.5%	52.4%	54.6%
Probability of 8h pain relief, UC	86.3%	85.5%	87.1%
Probability of 48h pain free, UC	77.5%	76.6%	78.4%
Probability of 48h pain relief, UC	91.1%	90.4%	91.7%
Probability of 2h pain relief, UBR	54.0%	52.3%	55.7%
Probability of 2h pain free, UBR	21.0%	19.6%	22.4%
Probability of 2h pain relief, UC	35.0%	33.9%	36.1%
Probability of 2h pain free, UC	11.0%	10.3%	11.7%
Probability of provider visits due to nausea/vomiting	3.8%	2.7%	4.9%
Probability of discontinuing due to lack of effect, UBR	21.8%	20.9%	22.7%
Discontinuation rate due to adverse events, UBR	2.5%	2.0%	3.0%
			\$350

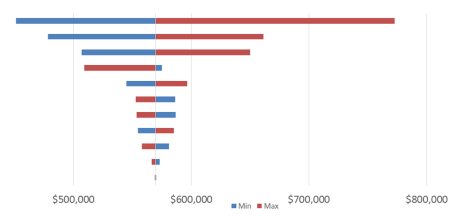
Min Max

UBR: ubrogepant; UC: usual care; ED: emergency department

\$750,000

Figure 4.2f. Model Costs and Utilities.	Ubrogepant versus Usual Care (Placebo)
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	Base Case	Min	Max
Disutility, ED visit	0.00	-0.01	0.00
Price, UBR	\$78	\$71	\$86
Disutility, hospitalization	0.00	-0.01	0.00
Price, UC	\$5	\$4	\$10
Utility, moderate migraine	0.77	0.75	0.80
Hospitalization cost per visit	\$6,840	\$6,156	\$7,524
Utility, no migraine	0.96	0.95	0.97
Utility, severe migraine	0.44	0.41	0.47
ED cost per visit	\$1,555	\$1,400	\$1,711
Utility, mild migraine	0.84	0.81	0.86
Provider office visit cost per visit	\$46	\$41	\$50



UBR: ubrogepant; UC: usual care; ED: emergency department

Using the placeholder prices, none of the treatments achieved cost effectiveness between thresholds of \$50,000 per QALY gained and \$150,000 per QALY gained in any of the probabilistic sensitivity analysis runs. At a cost-effectiveness threshold of \$250,000 per QALY gained, lasmiditan achieved cost effectiveness in 9.4% and ubrogepant 0.2% of the trials. One-way sensitivity analyses were not conducted when comparing the lasmiditan, rimegepant, and ubrogepant to each other, as none of the treatments were statistically different from each other in terms of effectiveness.

 Table 4.11. Probabilistic Sensitivity Analysis Results: Lasmiditan, Rimegepant, Ubrogepant

 Compared with Usual Care (Placebo)*

Treatment Compared with Usual Care	Cost-Effective at \$50,000 per QALY	Cost-Effective at \$100,000 per QALY	Cost-Effective at \$150,000 per QALY	Cost-Effective at \$200,000 per QALY	Cost-Effective at \$250,000 per QALY
Lasmiditan	0.0%	0.0%	0.0%	0.2%	9.4%
Rimegepant	0.0%	0.0%	0.0%	0.0%	0.0%
Ubrogepant	0.0%	0.0%	0.0%	0.1%	0.2%

QALY: quality-adjusted life year

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

Scenario Analyses Results

Modified Societal Perspective

The modified societal perspective included potential labor benefits for reduced migraine pain in the analysis. The ICERs for lasmiditan, rimegepant, and ubrogepant compared with usual care were \$207,800, \$422,900, and \$430,900 per QALY gained, respectively.

Other Scenarios

Exploratory analyses suggest that rimegepant and ubrogepant have a delayed effect beyond two hours. The scenario analysis evaluating this delayed effect resulted in a cost-effectiveness estimate of \$138,000 per QALY gained. Should these exploratory analyses be confirmed with an effect size compared to placebo similar to what was observed, and assuming that such an effect is unique to gepants, then rimegepant and ubrogepant may be cost-effective compared with placebo and would dominate lasmiditan.

Data from a long-term open-label study suggested that the frequency of migraines decreased over time.⁵⁶ In the scenario analysis evaluating the effect of a decreasing migraine frequency over time, total costs were lower, QALYs were higher, and hours of pain were lower for all treatments, including usual care. The cost-effectiveness ratios were similar to the base case.

Extending the timeline to 5 years had almost no effect on the cost-effectiveness estimates for the treatments. The ICERs at 5 years for lasmiditan, rimegepant, and ubrogepant compared with usual care were \$326,300, \$552,100, and \$562,000 per QALY gained, respectively.

The full results of all scenario analyses are presented in the Appendix as tables.

Threshold Analyses Results

Average annual prices that would result in willingness-to-pay thresholds of \$50,000 to \$150,000 per QALY gained for Population 1 are shown in table 4.13 below.

	Annual Price to Achieve \$50,000 per QALY	Annual Price to Achieve \$100,000 per QALY	Annual Price to Achieve \$150,000 per QALY
Lasmiditan	\$2,390	\$2,770	\$3,150
Rimegepant	\$1,960	\$2,210	\$2,460
Ubrogepant	\$1,950	\$2,200	\$2,440

Table 4.12. Threshold Analysis Results for Population 1 (Patients Who Cannot Take Triptans)

QALY: quality-adjusted life year

Model Validation

Model validation followed standard practices in the field. We tested all mathematical functions in the model to ensure they were consistent with the report (and supplemental Appendix materials). Model calculations were verified, and model input parameters were varied to evaluate face validity of changes in results. We also conducted sensitivity analyses with null input values to ensure the model was producing findings consistent with expectations. Model validation was also conducted in terms of comparisons to other model findings. We searched the literature to identify models that were similar to our analysis, with comparable populations, settings, perspective, and treatments.

Prior Economic Models

Our systematic review identified 28 potential pharmacoeconomic analyses of migraine therapies. We reviewed all 28 identified studies and found very few economic models for chronic treatment that involved a Markov model or long-term analysis. Also, extremely few included utilities or QALYs as an outcome. When developing the current model, we combined aspects of models for chronic migraine with other aspects from decision trees of acute migraine.¹¹²⁻¹²³

Some of the prior cost-effectiveness analyses that were most useful in developing our model examined preventive treatments for episodic and chronic migraine, including topiramate¹¹³ and

more recently erenumab.^{122,123} We identified three economic analyses of triptans for acute treatment of migraines. Perfetto et al. used a composite outcome measure to compare six triptans on cost per successfully treated patient, with successfully treated defined as requiring only one dose per attack during a 24-hour period.¹¹⁴ They estimated that eletriptan 40 mg would have the lowest cost per successfully treated patient compared to other triptans. Mullins et al. conducted a similar analysis from a Medicaid perspective, and again found that eletriptan had the lowest cost per successfully treated patient.¹¹⁸ Ramsberg and Henriksson analyzed the cost effectiveness of triptan treatment for a single attack from a Swedish societal perspective. They compared the cost per sustained pain-free response without adverse event and found that rizatriptan 10 mg and eletriptan 40 mg had the highest probability of cost effectiveness. However, none of these studies extended beyond the 24-hour time horizon nor estimated cost per LY or QALY ratios, and so could not be directly compared with the current analysis.

4.4 Summary and Comment

In our analysis of the cost effectiveness of lasmiditan, rimegepant, and ubrogepant, we found that for patients for whom triptans are not effective, not tolerated, or are contraindicated (Population 1), if these drugs are priced with the place-holder prices used in this analysis, they will exceed commonly accepted thresholds for cost effectiveness. Also, they will be dominated by sumatriptan and eletriptan in patients who can take triptans (Population 2) in that sumatriptan and eletriptan are both more effective and less expensive than these newer agents. However, prices for these therapies have not been released by the manufacturers and if the prices are set below those for the triptans, this conclusion would change.

Limitations

This analysis has several limitations and assumptions that must be considered when evaluating the results. We acknowledge that there is considerable heterogeneity among and even within individuals with migraine in terms of the frequency, severity, and unpredictability of attacks over time. Levels of pain severity (i.e., no, mild, moderate, or severe pain) were not reported in clinical trials. Instead, clinical trials used "freedom from pain" and "pain relief" at 2 hours as their primary outcomes. In addition, response to treatment was not reported for patients who did not have freedom from pain or pain relief at two hours. We therefore had to reconstruct pain levels to be able to apply utilities to the data. In doing so, we took a conservative approach to mapping "pain relief" to levels of pain, with patients potentially deriving more benefit from treatment than was likely observed in clinical trials. The result is that the model relies heavily on the outcomes of "pain freedom" or "pain relief" at two hours. As mentioned in the Controversies and Uncertainties section, because of study design characteristics requiring our reliance on 2-hour and 24-hour outcomes for the network meta-analysis and model, important differences among lasmiditan, rimegepant, and ubrogepant may not have been reflected well in the model. The model estimated

the effects of treatments in patients who were not "pain free" or did not have "pain relief" between 2 and 24 hours. Therefore, the benefit of these new medications may have been under- or overestimated if their relative benefit compared to placebo changed in the time period between 2 and 8 hours, affecting the estimated cost-effectiveness ratio. The scenario analysis evaluating an improved effect with ubrogepant (and similar effects observed with rimegepant) at 8 hours resulted in a dramatically different cost-effectiveness threshold. More research is needed to determine whether these delayed benefits beyond 2 hours are real and to provide a better estimate of the effect size.

Other limitations include that the probability of discontinuing a medication due to ineffective treatment was unknown for rimegepant, ubrogepant, sumatriptan, and eletriptan. As a result, we had to use an estimate derived from lasmiditan. Also, the probabilities for discontinuation due to adverse events were not available for sumatriptan alone or eletriptan. A rate from a trial evaluating sumatriptan plus naproxen was used for both treatments.

Importantly, prices for these therapies have not been released by the manufacturers, precluding final determination of their cost effectiveness based on actual pricing.

Conclusions

In our analysis of the cost effectiveness of lasmiditan, rimegepant, and ubrogepant, we found that for patients for whom triptans are not effective, not tolerated, or are contraindicated (Population 1), if these drugs are priced with the place-holder prices used in this analysis, they will exceed commonly accepted thresholds for cost effectiveness. Also, they will be dominated by sumatriptan and eletriptan in patients who can take triptans (Population 2), in that sumatriptan and eletriptan are both more effective and less expensive than these newer agents.

When compared with usual care in patients in whom triptans are not effective, not tolerated, or are contraindicated, these new acute treatments for migraine provide utility gains. Pricing of these drugs will determine whether they are cost effective at commonly used thresholds in patients who cannot take triptan medications. Also, due to the designs of the clinical trials, there remains considerable uncertainty surrounding estimating the impact of these acute treatments for migraine on patient quality of life, further complicating the estimation of their cost effectiveness.

5. Potential Other Benefits and Contextual Considerations

Our reviews seek to provide information on potential other benefits offered by the intervention to the individual patient, caregivers, the delivery system, other patients, or the public that would not have been considered as part of the evidence on comparative clinical effectiveness. We also recognize that there may be broader contextual issues related to the severity of the condition, whether other treatments are available, and ethical, legal, or other societal priorities that influence the relative value of illnesses and interventions. These general elements are listed in the table below, and the subsequent text provides detail about the elements that are applicable to the comparison of lasmiditan, rimegepant and ubrogepant to placebo and triptans (eletriptan and sumatriptan). We sought input from stakeholders, including individual patients, patient advocacy organizations, clinicians, and manufacturers, to inform the contents of this section.

Each ICER review culminates in a public meeting of an independent voting Council of clinicians, patients, and health services researchers. As part of their deliberations, Council members will judge whether a treatment may substantially impact the considerations listed in Table 5.1. The presence of substantial other benefits or contextual considerations may shift a council member's vote on an intervention's long-term value for money to a different category than would be indicated by the clinical evidence and cost-effectiveness analyses alone. For example, a council member may initially consider a therapy with an incremental cost-effectiveness ratio of \$150,000 per QALY to represent low long-term value for money. However, the Council member may vote for a higher value category if they consider the treatment to bring substantial other benefits or contextual considerations. Conversely, disadvantages associated with a treatment may lead a Council member to vote for a lower value category. A Council member may also determine that there are no other benefits or contextual considerations substantial enough to shift their vote. All factors that are considered in the voting process are outlined in ICER's <u>value assessment framework</u>. The content of these deliberations is described in the last chapter of ICER's Final Evidence Report, which is released after the public meeting.

This section, as well as the Council's deliberation, provides stakeholders with information to inform their decisions on a range of issues, including shared decision-making between patients and clinicians, coverage policy development, and pricing negotiations.

Table 5.1. Potential Other Benefits or Contextual Considerations (Not Specific to Any Disease or Therapy)

Potential Other Benefits
This intervention offers reduced complexity that will significantly improve patient outcomes.
This intervention will reduce important health disparities across racial, ethnic, gender, socio-economic, or
regional categories.
This intervention will significantly reduce caregiver or broader family burden.
This intervention offers a novel mechanism of action or approach that will allow successful treatment of many patients for whom other available treatments have failed.
This intervention will have a significant impact on improving return to work and/or overall productivity.
Other important benefits or disadvantages that should have an important role in judgments of the value of this
intervention.
Potential Other Contextual Considerations
This intervention is intended for the care of individuals with a condition of particularly high severity in terms of impact on length of life and/or quality of life.
This intervention is intended for the care of individuals with a condition that represents a particularly high lifetime burden of illness.
This intervention is the first to offer any improvement for patients with this condition.
There is significant uncertainty about the long-term risk of serious side effects of this intervention.
There is significant uncertainty about the magnitude or durability of the long-term benefits of this intervention.
There are additional contextual considerations that should have an important role in judgments of the value of this intervention.

5.1 Potential Other Benefits

For patients with migraine attacks, lasmiditan, rimegepant and ubrogepant represent the first new drugs for acute treatment with novel mechanisms of action to be submitted for FDA approval in over 20 years. Lasmiditan was approved on October 11, 2019 by the FDA for acute treatment of migraine and rimegepant and ubrogepant remain under review. These new therapies reflect translational research in which improved understanding of the mechanisms of disease has led to new therapeutics. Lasmiditan, the first ditan approved for use in the US, targets the 5HT1F (5-hydroxytryptamine 1F) receptor, and unlike the triptans does not induce vasoconstriction. The gepants, target CGRP, a peptide neural transmitter found in the pathways that play an important role in migraine. Monoclonal drugs that block CGRP have already been approved by the FDA for migraine prevention. Rimegepant and ubrogepant are the first small molecule gepants under review for relieving migraine attacks.

Similar to most triptans, lasmiditan, rimegepant and ubrogepant are orally available medications and would not be expected to increase the complexity of care. The favorable side effects seen to date with rimegepant and ubrogepant, similar to those seen with placebo, may make these medications attractive to patients and clinicians. The restriction on driving after taking lasmiditan is a potential other disadvantage of that therapy. Patients and advocates expressed the hope that these new therapies for patients with migraine may provide an effective and safe alternative for individuals who may turn to opioids and barbiturates because existing therapies are not effective, have intolerable side effects, or are not recommended because of the risk of misuse.

5.2 Contextual Considerations

For new medications that have mainly been evaluated in single dose comparative trials or noncomparative open-label studies of up to a year, there is uncertainty about their effects in actual clinical practice over time. Available data suggests that patients can use lasmiditan, rimegepant and ubrogepant for up to a year. However, the long-term benefits and harms of lasmiditan, rimegepant and ubrogepant are uncertain relative to other therapies that have years of experience.

For patients who improve with lasmiditan, rimegepant or ubrogepant and have tolerable side effects, it is expected that prolonged use for migraine attacks will be recommended. Questions remain about the duration of effectiveness, development of new side effects, and the risk of medication overuse headaches with frequent use. Lasmiditan, rimegepant and ubrogepant have not been shown to cause vasoconstriction, but whether they are free of cardiovascular adverse effects, particularly in those with cardiovascular disease or at high risk, remains to be proven.

The availability of new treatments for migraine is likely to allow some patients to remain at work in situations where they would otherwise have needed to miss or leave work.

6. Value-Based Price Benchmarks

Annual value-based price benchmarks (VBPBs) of these drugs (vs. usual care) are presented in Table 6.1. The VBPB for a drug is defined as the price range that would achieve incremental costeffectiveness ratios between \$100,000 and \$150,000 per QALY gained. While the results of the NMAs suggest that lasmiditan may be slightly more efficacious than rimegepant and ubrogepant, they do not exclude comparable efficacy given the overlapping confidence intervals. Additionally, lasmitidan treatment results in more adverse events and is discontinued more frequently. Given that we we felt the net benefits of the therapies were relatively similar as reflected in our comparative evidence ratings, we developed a range of value-based price benchmarks across all three drugs, using the range of threshold prices reported in Section 4.3 so as to avoid suggesting greater certainty in the individual threshold prices than is warranted.

For these drugs, price discounts of approximately 30% to 46% from the assumed list price would be required to reach the \$150,000 per QALY threshold price range (Table 6.1). Price discounts of approximately 39% to 51% from assumed list prices would be required to reach the \$100,000 per QALY threshold price range. Note that these discounts are from the assumed placeholder prices, and not from actual list prices, which are not yet known.

As there is no mortality effect in the model, cost per LY gained is not relevant, and the cost per evLYG is essentially the same as the cost per QALY gained. We therefore do not report VBPBs for these in the table below.

Table 6.1. Value-Based Price Benchmark Ranges for Lasmiditan, Rimegepant, and Ubrogepant versus Usual Care in Population 1 (Patients Who Cannot Take Triptans)

	Assumed Annual Price*	Annual Price at \$100,000 Threshold	Annual Price at \$150,000 Threshold
Drug Price	\$4,515	\$2,200-\$2,770	\$2,440-\$3,150
Discount from Assumed Price*		39% to 51%	30% to 46%

QALY: quality-adjusted life year, WAC: wholesale acquisition cost

*Using assumed placeholder prices for lasmiditan, rimegepant, and ubrogepant.

7. Potential Budget Impact

7.1 Overview

We used the cost-effectiveness model to estimate the potential total budgetary impact of each drug (lasmiditan, rimegepant, and ubrogepant) for prevalent individuals in the United States (US) aged 18 years and over experiencing migraines requiring acute treatment, with or without aura. We also included a scenario analysis where the frequency of migraines is assumed to decrease over time.

We used the same assumed placeholder price and threshold prices for each drug in our estimates of budget impact, rather than using the three threshold prices calculated for each drug in Section 4.3. As mentioned above, while the results of the NMAs suggest that lasmiditan may be slightly more efficacious than rimegepant and ubrogepant, they do not exclude comparable efficacy given the overlapping confidence intervals and higher adverse event and discontinuation rates with lasmiditan. We therefore used a blended range of prices in our potential budget impact analyses, using the same \$50,000, \$100,000, and \$150,000 cost-effectiveness threshold price for each drug. From the threshold prices for the three drugs, we used the lowest price for the \$50,000 per QALY threshold (\$1,950) and for the \$100,000 threshold (\$2,200), and the highest price for the \$150,000 per QALY threshold (\$3,150).

7.2 Methods

We used results from the same model employed for the cost-effectiveness analyses to estimate total potential budget impact. Potential budget impact was defined as the total differential cost of using each new therapy rather than relevant existing therapy for the treated population, calculated as differential health care costs (including drug costs) minus any offsets in these costs from averted health care events. All costs were undiscounted and estimated over a five-year time horizon, given the potential for cost offsets to accrue over time and to allow a more realistic impact on the number of patients treated with the new therapy.

This potential budget impact analysis does not include the population cohort of patients with migraines who are eligible for treatment with triptans, as sumatriptan and eletriptan dominated these drugs in our cost-effectiveness analysis. This potential budget impact analysis includes the cohort of patients who had migraine attacks that did not respond to non-prescription medicines and for whom triptans had not been effective, were not tolerated, or were contraindicated. To estimate the size of the potential candidate population for treatment, we first used an estimate derived from the 2012 National Health Interview Survey of 14.2% for the prevalence of US adults 18 or older reporting having migraine or severe headache.³ The American Migraine Prevalence and Prevention Study found in a survey of migraine patients that 48.9% reported using prescription

medicines (only or sometimes) for acute treatment.¹²⁴ Based on an estimate that triptans work in approximately 60% to 70% of migraine patients,¹²⁵ we assumed that 35% of migraine patients attempting prescription treatments would fall into this non-triptan cohort. We applied these estimated proportions to the average 2020-2024 estimated US adult population to arrive at an eligible population size of approximately 6.4 million patients, or approximately 1.3 million patients each year over five years. We assumed in our analysis of potential budget impact in this population that each drug would be added to usual care, rather than displacing other migraine-specific treatments.

ICER's methods for estimating potential budget impact are described in detail elsewhere¹²⁶ and have been recently <u>updated</u>. The intent of our revised approach to budgetary impact is to document the percentage of patients who could be treated at selected prices without crossing a budget impact threshold that is aligned with overall growth in the U.S. economy. For 2019-2020, the five-year annualized potential budget impact threshold that should trigger policy actions to manage access and affordability is calculated to be approximately \$819 million per year for new drugs.

7.3 Base-Case Results

Table 7.1 illustrates the five-year annualized per-patient potential budget impact of lasmiditan compared to usual care in this population. These results are based on the assumed placeholder price (\$4,515 per year), and the annual threshold prices listed above for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Table 7.1. Annualized Per-Patient Potential Budget Impact Over a Five-year Time Horizon for
Lasmiditan versus Usual Care

	Average Annual Per Patient Budget Impact			
	At Placeholder Price*	At \$150,000/QALY Price	At \$100,000/QALY Price	At \$50,000/QALY Price
Lasmiditan	\$6,950	\$5,980	\$5,300	\$5,120
Usual Care	\$5,160			
Net Impact	\$1,790	\$820	\$140	-\$40

*Assumed placeholder price.

All annualized costs include drug and non-drug health care costs.

QALY: quality-adjusted life year

For lasmiditan, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$1,790 versus usual care. Its average annualized potential budget impact versus usual care at the threshold prices for \$50,000 to \$150,000 per QALY ranged from cost-saving to approximately \$820 per patient.

In this population, as shown in Figure 7.1, approximately 12% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at lasmiditan's assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000. All eligible patients could be treated at the \$100,000 and \$50,000 threshold prices, with estimated potential budget impact of approximately 62% of the threshold at the \$100,000 threshold price.

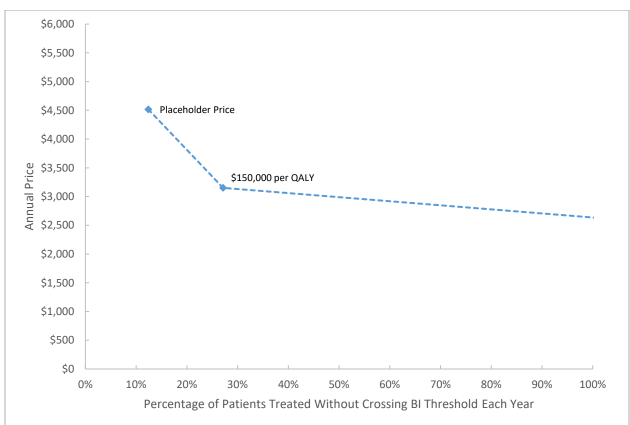


Figure 7.1. Potential Budget Impact Scenarios of Lasmiditan Usual Care at Different Acquisition Prices

*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

Table 7.2 illustrates the five-year annualized per-patient potential budget impact of rimegepant compared to usual care in the same population. These results are based on the assumed placeholder price (\$4,515 per year), and the annual threshold prices listed above for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Table 7.2. Annualized Per-Patient Potential Budget Impact Over a Five-year Time Horizon forRimegepant versus Usual Care

	Average Annual Per Patient Budget Impact				
	At Placeholder Price*	At Price to Reach \$150,000/QALY	At Price to Reach \$100,000/QALY	At Price to Reach \$50,000/QALY	
Rimegepant	\$7,420	\$6,330	\$5,560	\$5,360	
Usual Care		\$5,160			
Net Impact	\$2,260	\$1,170	\$400	\$200	

All annualized costs include drug and non-drug health care costs.

QALY: quality-adjusted life year

*Assumed placeholder price.

For rimegepant, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$2,260 versus usual care. Its average annualized potential budget impact versus usual care at the threshold prices for \$50,000 to \$150,000 per QALY ranged from approximately \$200 per patient to approximately \$1,170 per patient.

As shown in Figure 7.2, approximately 10% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at rimegepant's assumed placeholder price. Approximately 18% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price, increasing to approximately 55% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 91% of the threshold.

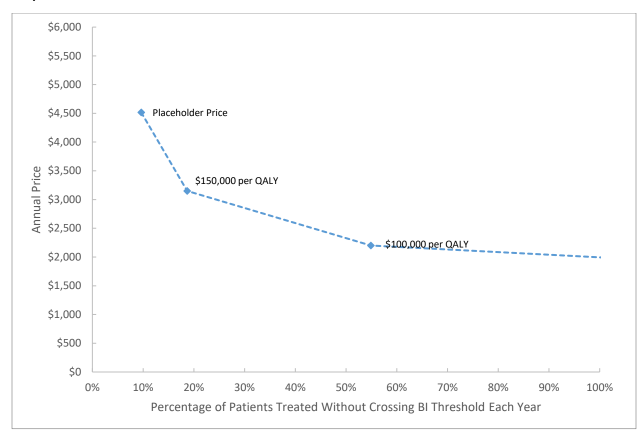


Figure 7.2. Potential Budget Impact Scenarios of Rimegepant vs. Usual Care at Different Acquisition Prices

*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

Table 7.3 illustrates the five-year annualized per-patient potential budget impact of ubrogepant compared to usual care in this population. These results are based on the assumed placeholder price (\$4,515 per year), and the annual threshold prices listed above for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Table 7.3. Annualized Per-Patient Potential Budget Impact Over a Five-year Time Horizon forUbrogepant versus Usual Care

	Average Annual Per Patient Budget Impact					
	At Placeholder Price*					
Ubrogepant	\$7,430	\$6,330	\$5,560	\$5,360		
Usual Care	\$5,160					
Net Impact	\$2,270	\$1,170	\$400	\$200		

All annualized costs include drug and non-drug health care costs.

QALY: quality-adjusted life year

*Assumed placeholder price.

For ubrogepant, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$2,270 versus usual care. Its average annualized potential budget impact versus usual care at the threshold prices for \$50,000 to \$150,000 per QALY ranged from approximately \$200 per patient to approximately \$1,170 per patient.

As shown in Figure 7.3, approximately 10% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at ubrogepant's assumed placeholder price. Approximately 19% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price, increasing to approximately 55% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 91% of the threshold.

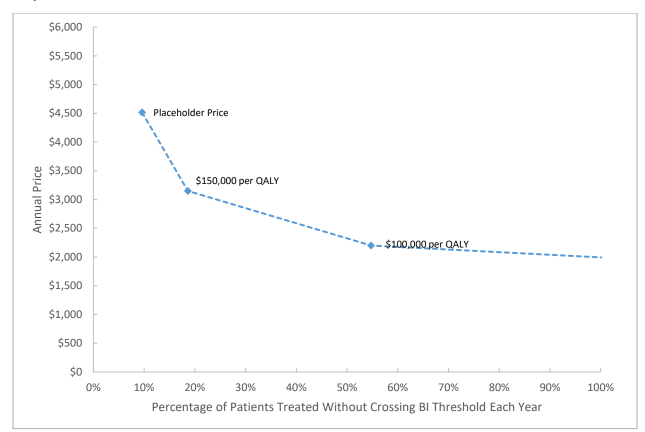


Figure 7.3. Potential Budget Impact Scenarios of Ubrogepant versus Usual Care at Different Acquisition Prices

*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

While we used the same prices for all three drugs, the potential budget impact results for each drug are different from each other, especially those for lasmiditan compared to rimegepant and ubrogepant. This is because of differences in efficacy and discontinuation rates across the drugs.

7.4 Scenario Results

Data from a long-term open label safety study suggests that the frequency of migraines decreased over time.⁵⁶ While this single-arm trial was not designed to evaluate whether the same effect was observed in a control population, decreasing migraine frequency over time could have a significant impact on budget impact analyses. We therefore created a scenario analysis where we modeled the potential budget impact of these treatments if migraine frequency decreases over time.

Table 7.4 illustrates the five-year annualized per-patient potential budget impact of lasmiditan compared to usual care under this scenario. These results are based on the assumed placeholder

price (\$4,515 per year), and the same annual prices for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Average Annual Per Patient Budget Impact				:
	At Placeholder Price*	At Price to Reach \$150,000/QALY	At Price to Reach \$100,000/QALY	At Price to Reach \$50,000/QALY
Lasmiditan	\$4,880	\$4,180	\$3,690	\$3,570
Usual Care	\$3,590			
Net Impact	\$1,290	\$590	\$100	-\$30

Table 7.4. Annualized Per-Patient Potential Budget Impact Over a Five-year Time Horizon for Lasmiditan versus Usual Care: Decreased Frequency Scenario

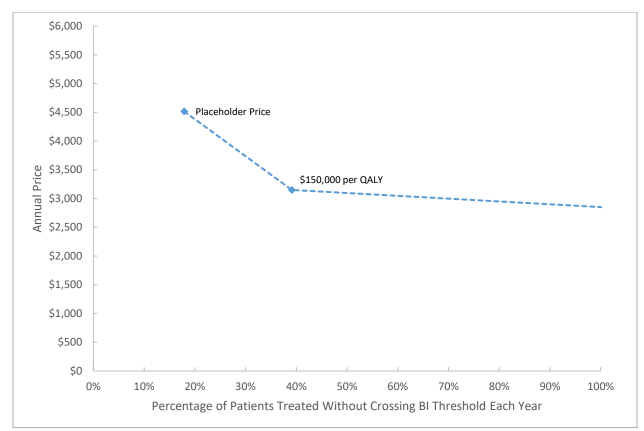
*Assumed placeholder price.

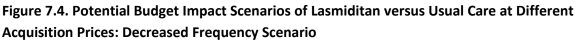
All annualized costs include drug and non-drug health care costs.

QALY: quality-adjusted life year

For lasmiditan, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$1,290 versus usual care. Its average annualized potential budget impact versus usual care at the threshold prices for \$50,000 to \$150,000 per QALY ranged from cost-saving to approximately \$590 per patient.

In this scenario, as shown in Figure 7.4, approximately 18% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at lasmiditan's assumed placeholder price. Approximately 39% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000. All eligible patients could be treated at the \$100,000 and \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 43% of the threshold at the \$100,000 threshold price.





*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

Table 7.5 illustrates the five-year annualized per-patient potential budget impact of rimegepant compared to usual care in the decreased frequency scenario. These results are based on the assumed placeholder price (\$4,515 per year), and the same annual prices for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Table 7.5. Annualized Per-Patient Potential Budget Impact Over a Five-year Time Horizon forRimegepant versus Usual Care: Decreased Frequency Scenario

Average Annual Per Patient Budget Impact				
	At Placeholder Price*	At Price to Reach \$150,000/QALY	At Price to Reach \$100,000/QALY	At Price to Reach \$50,000/QALY
Rimegepant	\$5,200	\$4,420	\$3,880	\$3,730
Usual Care		\$3,590		
Net Impact	\$1,600	\$830	\$280	\$140

All annualized costs include drug and non-drug health care costs.

QALY: quality-adjusted life year

*Assumed placeholder price.

For rimegepant in this decreased frequency scenario, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$1,600 versus usual care. Its average annualized potential budget impact versus usual care at the threshold prices to reach cost-effectiveness thresholds of \$50,000 to \$150,000 per QALY ranged from approximately \$140 per patient to approximately \$830 per patient.

As shown in Figure 7.5, approximately 14% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at rimegepant's assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at the \$150,000 threshold price , increasing to approximately 80% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 62% of the threshold.

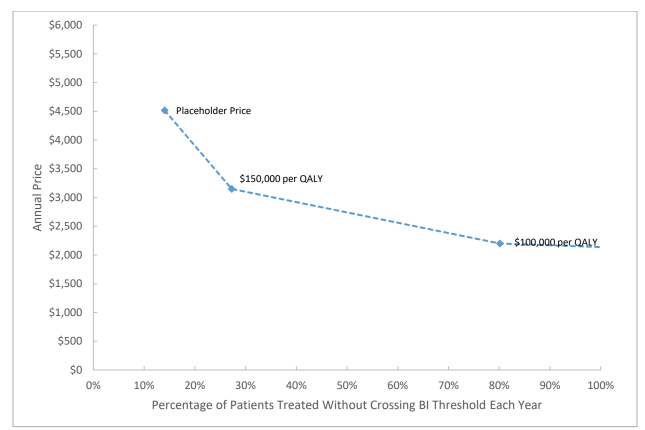


Figure 7.5. Potential Budget Impact Scenarios of Rimegepant versus Usual Care at Different Acquisition Prices: Decreased Frequency Scenario

*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

Table 7.6 illustrates the five-year annualized per-patient potential budget impact of ubrogepant compared to usual care under the decreased frequency scenario. These results are based on the assumed placeholder price (\$4,515 per year), and the same annual prices for thresholds of \$150,000, \$100,000, and \$50,000 per QALY versus usual care (\$3,150, \$2,200, and \$1,950, respectively).

Table 7.6. Annualized Per-Patient Potential Budget Impact over a Five-year Time Horizon forUbrogepant versus Usual Care: Decreased Frequency Scenario

	Average Annual Per Patient Budget Impact			
	At Placeholder Price*	At Price to Reach \$150,000/QALY	At Price to Reach \$100,000/QALY	At Price to Reach \$50,000/QALY
Ubrogepant	\$5,200	\$4,420	\$3,880	\$3,730
Usual Care	\$3,590			
Net Impact	\$1,610	\$830	\$280	\$140

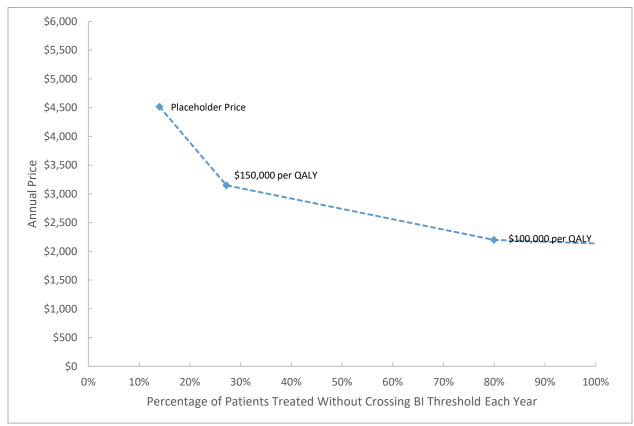
QALY: quality-adjusted life year

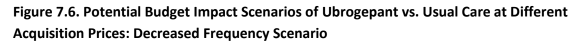
*Assumed placeholder price

All annualized costs include drug and non-drug health care costs.

For ubrogepant in this scenario, the average annualized potential budgetary impact when using its assumed placeholder price was an additional per-patient cost of approximately \$1,610 versus usual care. Its average annualized potential budget impact versus usual care at threshold prices to reach cost-effectiveness thresholds of \$50,000 to \$150,000 per QALY ranged from approximately \$140 per patient to approximately \$830 per patient.

As shown in Figure 7.6, approximately 14% of eligible patients could be treated in a given year without crossing the ICER budget impact threshold of \$819 million at ubrogepant's assumed placeholder price. Approximately 27% of eligible patients could be treated without crossing the budget impact threshold at its price to reach the cost-effectiveness threshold of \$150,000, increasing to approximately 80% at the price to reach \$100,000 per QALY. All eligible patients could be treated at the \$50,000 per QALY threshold price, with estimated potential budget impact of approximately 63% of the threshold.





*Prices used are derived from a single price range for all three drugs rather than the individual drug threshold prices.

BI: budget impact, QALY: quality-adjusted life year

This is the third ICER review of interventions for migraine.

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APPENDICES

Appendix A. Search Strategies and Results

Table A1. PRISMA 2009 Checklist

	#	Checklist item
		TITLE
Title	1	Identify the report as a systematic review, meta-analysis, or both.
		ABSTRACT
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.
		INTRODUCTION
Rationale	3	Describe the rationale for the review in the context of what is already known.
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).
		METHODS
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.

	#	Checklist item
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I ²) for each meta-analysis.
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.
		RESULTS
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).
		DISCUSSION
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.
		FUNDING
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.

From: Moher D, Liberati A, Tetzlaff J, Altman DG. The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoS Med 6(6): e1000097. doi:10.1371/journal.pmed1000097

Search Strategies for Acute Treatments for Migraine

Table A2. Search Strategy of Medline 1996 to Present with Daily Update and Cochrane CentralRegister of Controlled Trials (via Ovid) - Lasmiditan/Rimegepant/Ubrogepant

#	Search Terms
1	exp migraine disorders/
2	exp migraine with aura/
3	exp migraine without aura/
4	((acute AND migraine*) OR migraine* OR migraine syndrome OR migraine disorder).ti,ab.
5	OR/1-4
6	(lasmiditan OR COL-144 OR LY573144 OR rimegepant OR BHV-3000 OR BMS-927711 OR ubrogepant
	OR MK-1602).ti,ab.
7	5 AND 6
8	(animals not (humans and animals)).sh.
9	7 NOT 8
10	(addresses or autobiography or bibliography or biography or clinical trial, phase I or comment or
	congresses or consensus development conference or duplicate publication or editorial or guideline or
	in vitro or interview or lecture or legal cases or legislation or letter or news or newspaper article or
	patient education handout or periodical index or personal narratives or portraits or practice
	guideline or review or video audio media).pt.
11	9 NOT 10
12	Limit 11 to English language
13	Remove duplicates from 12

Table A3. Search Strategy of Medline 1996 to Present with Daily Update and Cochrane CentralRegister of Controlled Trials (via Ovid) – Sumatriptan & Eletriptan (updated)

#	Search Terms
1	exp migraine disorders/
2	exp migraine with aura/
3	exp migraine without aura/
4	((acute AND migraine*) OR migraine* OR migraine syndrome OR migraine disorder).ti,ab.
5	OR/1-4
6	(sumatriptan OR eletriptan).ti,ab.
7	5 AND 6
8	(animals not (humans and animals)).sh.
9	7 NOT 8
10	(addresses or autobiography or bibliography or biography or clinical trial, phase I or comment or
	congresses or consensus development conference or duplicate publication or editorial or guideline or
	in vitro or interview or lecture or legal cases or legislation or letter or news or newspaper article or
	patient education handout or periodical index or personal narratives or portraits or practice
	guideline or review or video audio media).pt.
11	9 NOT 10
12	Limit 11 to English language
13	limit 12 to yr="2016- Current"
14	Remove duplicates from 13

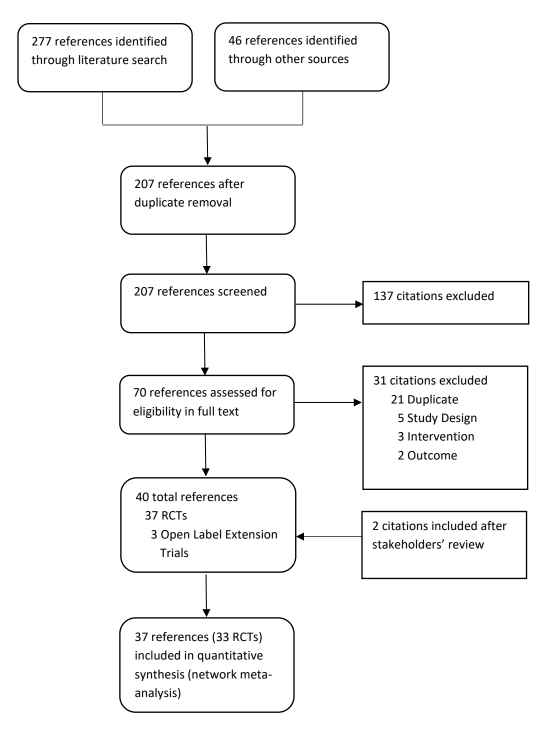
Table A4. Search Strategy of EMBASE Search - Lasmiditan/Rimegepant/Ubrogepant

#	Search Terms
#1	acute AND ('migraine'/exp OR migraine)
#2	'lasmiditan'/exp OR 'lasmiditan' OR 'COL-144' OR 'LY573144'
#3	'rimegepant'/exp OR 'rimegepant' OR 'BHV-3000' OR 'BMS-927711'
#4	'ubrogepant'/exp OR 'ubrogepant' OR 'MK-1602'
#5	#2 OR #3 OR #4
#6	#1 AND #5
#7	'animal'/exp or 'nonhuman'/exp or 'animal experiment'/exp NOT 'human'/exp
#8	#6 NOT #7
#9	#8 AND [english]/lim
#10	#9 AND [medline]/lim
#11	#9 NOT #10
#12	#11 NOT ('case report'/de OR 'human tissue'/de OR 'nonhuman'/de OR 'practice guideline'/de OR
	'questionnaire'/de OR 'chapter'/it OR 'conference review'/it OR 'editorial'/it OR 'letter'/it OR 'note'/it
	OR 'review'/it OR 'short survey'/it)

	As search stategy of Emphase search - sumariptan a Electriptan (aparted)
#	Search Terms
#1	acute AND ('migraine'/exp OR migraine)
#2	'Sumatriptan'/exp OR 'Sumatriptan'
#3	'eletriptan'/exp OR 'eletriptan'
#4	#2 OR #3
#5	#1 AND #4
#6	'animal'/exp or 'nonhuman'/exp or 'animal experiment'/exp NOT 'human'/exp
#7	#5 NOT #6
#8	#7 AND [english]/lim
#9	#8 AND [medline]/lim
#10	#8 NOT #9
#11	#10 AND [01-01-2016]/sd
#12	#11 NOT ('case report'/de OR 'human tissue'/de OR 'nonhuman'/de OR 'practice guideline'/de OR
	'questionnaire'/de OR 'chapter'/it OR 'conference review'/it OR 'editorial'/it OR 'letter'/it OR 'note'/it
	OR 'review'/it OR 'short survey'/it)
#13	#12 AND 'randomized controlled trial'/exp OR 'controlled clinical trial'/exp OR random*:ti,ab OR
	placebo:ti,ab OR 'drug therapy':lnk OR trial:ti,ab OR groups:ti,ab

Table A5. Search Strategy of EMBASE Search – Sumatriptan & Eletriptan (updated)

Figure A1. PRISMA flow Chart Showing Results of Literature Search for Acute Treatments for Migraine



RCT: randomized control trial

<u>Appendix B. Previous Systematic Reviews and</u> <u>Technology Assessments</u>

Xu F, Sun W. Network Meta-Analysis of Calcitonin Gene-Related Peptide Receptor Antagonists for the Acute Treatment of Migraine. *Frontiers in pharmacology.* 2019;10:795.

The investigators performed a network meta-analysis (NMA) to indirectly compare and rank six different calcitonin gene-related peptide (CGRP) receptor antagonists (telcagepant, olcegepant, BI 44370, rimegepant, MK3207, and ubrogepant) for the acute treatment of migraine. Ten randomized controlled trials (RCTs) in adult patients with migraine were included in the quantitative analysis. Efficacy was evaluated based on pain-freedom at 2-hours, and safety was assessed based on the occurrence of adverse events (AEs) and drug-related AEs. Olcegepant, ubrogepant, and BI 44370 were statistically significantly better than placebo in achieving pain freedom at 2-hours. In addition, olcegepant was found to show greater efficacy than the other CGRP receptor antagonists and to be marginally more efficacious than triptans, however, statistical significance was not reached. Telcagepant, olcegepant, MK3207, rimegepant, and ubrogepant were found to have a safety profile comparable to placebo, while BI 44370 was associated with an increased risk for AEs. Of note, research regarding olcegepant, telcagepant, BI 44370, and MK3207 has been discontinued, primarily due to concerns of hepatoxicity.

Thorlund K, Toor K, Wu P, et al. Comparative tolerability of treatments for acute migraine: A network meta-analysis. *Cephalalgia: an international journal of headache.* 2017;37(10):965-978.

This systematic literature review and NMA was conducted to evaluate the comparative tolerability of acute treatments for migraine with regards to AEs, treatment-related AEs (TRAEs), and serious AEs (SAEs). The NMA included 141 RCTs, comparing acute oral treatments for migraine in adults, including triptans (almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, sumatriptan, and zolmitriptan), NSAIDs (diclofenac, ibuprofen, naproxen, and selective COX-2 inhibitors), acetaminophen, as well as ergotamines. Triptans were generally associated with the highest odds ratios (ORs) for the occurrence of any AEs and TRAEs (i.e. fatigue, dizziness, chest discomfort, somnolence and nausea). Specifically, sumatriptan, eletriptan, rizatriptan, zolmitriptan, and the combination treatment of sumatriptan and naproxen had statistically significant higher odds of TRAE occurring compared with placebo. Among the non-triptans and non-triptans were not associated with increased odds of SAEs compared to placebo. The authors concluded however that differences in safety profiles were not large enough to necessitate prioritizing one treatment over another.

Xu H, Han W, Wang J, Li M. Network meta-analysis of migraine disorder treatment by NSAIDs and triptans. *J Headache Pain.* 2016;17(1):113.

The investigators performed an NMA to compare the relative efficacy and tolerability of NSAIDs and triptans in the acute treatment for migraine in adults. Eighty-eight RCTs pertaining to sumatriptan, zolmitriptan, almotriptan, rizatriptan, naratriptan, eletriptan, ibuprofen, sumatriptan-naproxen, diclofenac-potassium, and aspirin were included in the analysis. Efficacy was evaluated based on pain-freedom, pain-relief, absence of nausea, rate of recurrence, and the use of rescue medication. Safety was evaluated based on the occurrence of AEs. With regards to pain-freedom and pain-relief at 2-hours, all treatments included in the NMA were found to be statistically more effective than placebo. Eletriptan exhibited superior efficacy over sumatriptan, zolmitriptan, almotriptan, ibuprofen, and aspirin with regards to 2-hour pain-freedom, while rizatriptan was superior to sumatriptan, zolmitran, almotriptan, ibuprofen, and aspirin. The difference between eletriptan and rizatriptan was not found to be statistically significant. With regards to absence of nausea at 2hours, rizatriptan was found to have better efficacy outcomes compared to sumatriptan, while no other meaningful differences were found between the other treatments including placebo. The AE incidence of sumatriptan was higher compared to diclofenac-potassium, ibuprofen, and almotriptan. Similarly, the safety profile for naratriptan was found to be inferior to that of ibuprofen and diclofenac-potassium. Results overall suggested that eletriptan exhibited the best efficacy results while also having an acceptable safety profile. Sumatriptan-naproxen and diclofenac-potassium also showed favorable efficacy as well as tolerability, while ibuprofen appeared the best tolerated treatment option. The authors concluded that eletriptan may be the most suitable treatment option for the acute treatment of migraines when taking both efficacy and safety outcomes into account. Additionally, ibuprofen was also considered to be an appropriate treatment option due to its excellent safety profile.

Cameron C, Kelly S, Hsieh SC, et al. Triptans in the Acute Treatment of Migraine: A Systematic Review and Network Meta-Analysis. *Headache.* 2015;55 Suppl 4:221-235.

This systematic review and NMA sought to compare triptans (almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, sumatriptan, and zolmitriptan) to each other, versus placebo, and versus other acute migraine treatments such as NSAIDs, ASA, acetaminophen, ergotamines, opioids, or antiemetics. A total of 133 single-attack RCTs evaluating acute treatments for migraines in adults were included in the quantitative analysis. Efficacy was evaluated based on pain-freedom and headache relief at 2-hours, sustained pain-freedom and headache relief at 24-hours, as well as the use of rescue medication. Results found that rizatriptan (oral), eletriptan (oral), and sumatriptan (subcutaneous injection) have the largest effect on 2-hour pain-freedom among all monotherapies. With respect to 2-hour pain-relief, sumatriptan (subcutaneous injection), rizatriptan (oral), and zolmitriptan (oral) showed the largest effect compared to the other monotherapies. Eletriptan (oral) and rizatriptan (oral) exhibited the largest effect on sustained freedom of pain, while zolmitriptan (oral) and eletriptan (oral) were found to be most efficacious with respect to sustained

pain relief. Participants treated with eletriptan (oral) and zolmitriptan (oral) required the least amount of rescue medications, while those treated with NSAIDs, sumatriptan (oral), and ASA required the most doses. The authors concluded that the majority of triptans, with the exception of frovatriptan and naratriptan, are comparable in terms of efficacy. However, it was suggested that eletriptan and rizatriptan may be slightly superior in providing pain relief.

Appendix C. Ongoing Studies

Title/ Trial Sponsor	Study Design	Comparators	Patient Population	Key Outcomes	Estimated Completion Date
			Lasmiditan		
Randomized Controlled Trial of Lasmiditan Over Four Migraine Attacks NCT03670810 Sponsor: Eli Lilly	Phase 3, Randomized, double-blind, parallel assignment <u>Estimated N</u> : 1600 <u>Time Frame</u> : 16 weeks	 Lasmiditan high dose Lasmiditan low dose Placebo 	Inclusions:≥18 years; Migraine with or without aura; History ofdisabling migraine for at least 1 year; Migraine onsetbefore the age of 50 years; 3 to 8 migraineattacks/month (<15 headache days/month) duringthe past 3 months; MIDAS score ≥11Exclusion:Known hypersensitivity to lasmiditan; History ofhemorrhagic stroke, epilepsy, or any other conditionplacing the participant at increased risk of seizures;History of recurrent dizziness and/or vertigo; Historyof diabetes mellitus with complications; History oforthostatic hypotension with syncope; Significantrenal or hepatic impairment; Participants who aredeemed to be at significant risk for suicide; History ofchronic migraine or other forms of primary orsecondary chronic headache disorder within past 12months; Use of more than 3 doses/month of eitheropioids or barbiturates; Initiation of or a change inconcomitant medication to reduce the frequency ofmigraine episodes within 3 months prior to screening;SUD within 1 year prior to screening; Currentlyenrolled in any other clinical study involving aninvestigational product	Primary Outcomes: Pain freedom at 2-hours postdose during the first attack; Pain freedom at 2- hours postdose in at least 2 out of 3 attacks <u>Secondary Outcomes</u> : 2-hour pain freedom; Freedom of MBS; 24-hour sustained pain freedom; Use of rescue medication; Freedom of associated symptoms at 2-hours; Migraine recurrence at 24- hours; Pain freedom, pain relief, freedom from MBS, and no disability at 2- hours; Change in MIDAS score; No disability at 2- hours; PGI-C at 2-hours; MQoLQ score at 24-hours; Patient satisfaction; Change in EQ-5D-5L at 24- hours	March 2020
RandoMized, Double-blind, Placebo-controlled Trial Of Lasmiditan in a Single	Phase 2, Randomized, double-blind, parallel assignment	 Lasmiditan high dose Lasmiditan mid dose 	Inclusions: ≥18 years; Migraine with or without aura; History of disabling migraine for at least 1 year; MIDAS score ≥11; Migraine onset before the age of 50 years;	Primary Ouctomes: Pain freedom at 2-hours (high dose) Secondary Outcomes:	March 2020

Title/ Trial Sponsor	Study Design	Comparators	Patient Population	Key Outcomes	Estimated Completion Date
Migraine Attack in Japanese Patients Suffering From Migraine With or Without Aura - the MONONOFU Study NCT03962738 Sponsor: Eli Lilly	Estimated N: 36 <u>Time Frame</u> : up to 50 days	 Lasmiditan low dose Placebo 	History of 3 to 8 migraine attacks/month and <15 headache days/month during the past 3 months <u>Exclusions</u> : Known hypersensitivity to lasmiditan; History of hemorrhagic stroke, epilepsy, or any other condition placing the patient at increased risk of seizures; History of recurrent dizziness and/or vertigo; History of diabetes mellitus with complications; History of orthostatic hypotension with syncope	Pain freedom in each dose group at 2-hours; Pain relief at 2-hours; Freedom of MBS at 2-hours; 24- and 48-hour sustained pain freedom; Freedom of phonophobia, photophobia, nausea, and vomiting; No disability at 2- hours; Change in EQ-5D-5L at 24-hours; PGI-C at 2- hours; MQoLQ score at 24- hours	
			Rimegepant		
An Open-label, Intermediate-size, Expanded Access Study of BHV-3000 in the Acute Treatment of Migraine NCT03934086 Sponsor: Biohaven Pharmaceuticals, Inc.	Expanded Access		Inclusions:Patients who participated in a previous BHV-3000/Rimegepant Clinical TrialExclusions:History of basilar migraine or hemiplegic migraine;History with current evidence of uncontrolled,unstable or recently diagnosed cardiovasculardisease; HIV; Uncontrolled hypertension or diabetes;Current diagnosis of major depression, other painsyndromes, psychiatric conditions, dementia, orsignificant neurological disorders (other thanmigraine) that might interfere with studyassessments; History of gastric, or small intestinalsurgery, or disease that causes malabsorption	The purpose of this protocol is to allow subjects who completed any BHV3000 (rimegepant) clinical study to continue to have access to rimegepant while collecting ongoing safety data	

Title/ Trial Sponsor	Study Design	Comparators	Patient Population	Key Outcomes	Estimated Completion Date

ALT: Alanine Aminotransferase, AST: Aspartate Aminotransferase, BMI: Body mass index, EQ-5D-5L: EuroQol 5-Dimension 5-Level Scale, HbA1c: Hemoglobulin A1c, HIV: Human Immunodeficiency Virus, MBS: most bothersome symptom, MIDAS: Migraine Disability Assessment Test, MQoLQ: Migraine Quality of Life Questionnaire, N: total number, PGI-C: Patient Global Impression of Change, ULN: Upper Limit

Source: www.ClinicalTrials.gov (NOTE: studies listed on site include both clinical trials and observational studies)

Appendix D. Comparative Clinical Effectiveness Supplemental Information

We performed screening at both the abstract and full-text level. A single investigator screened all abstracts identified through electronic searches according to the inclusion and exclusion criteria described earlier. We did not exclude any study at abstract-level screening due to insufficient information. For example, an abstract that did not report an outcome of interest would be accepted for further review in full text. We retrieved the citations that were accepted during abstract-level screening for full text appraisal. One investigator reviewed full papers and provided justification for exclusion of each excluded study.

We used criteria published by the US Preventive Services Task Force (USPSTF) to assess the quality of RCTs and comparative cohort studies, using the categories "good," "fair," or "poor" (see Appendix Table F2).¹²⁷ Guidance for quality ratings using these criteria is presented below, as is a description of any modifications we made to these ratings specific to the purposes of this review.

Good: Meets all criteria: Comparable groups are assembled initially and maintained throughout the study; reliable and valid measurement instruments are used and applied equally to the groups; interventions are spelled out clearly; all important outcomes are considered; and appropriate attention is paid to confounders in analysis. In addition, intention to treat analysis is used for RCTs.

Fair: Studies were graded "fair" if any or all of the following problems occur, without the fatal flaws noted in the "poor" category below: Generally comparable groups are assembled initially but some question remains whether some (although not major) differences occurred with follow-up; measurement instruments are acceptable (although not the best) and generally applied equally; some but not all important outcomes are considered; and some but not all potential confounders are addressed. Intention to treat analysis is done for RCTs.

Poor: Studies were graded "poor" if any of the following fatal flaws exists: Groups assembled initially are not close to being comparable or maintained throughout the study; unreliable or invalid measurement instruments are used or not applied equally among groups (including not masking outcome assessment); and key confounders are given little or no attention. For RCTs, intention to treat analysis is lacking.

Note that case series are not considered under this rating system—because of the lack of comparator, these are generally considered to be of poor quality.

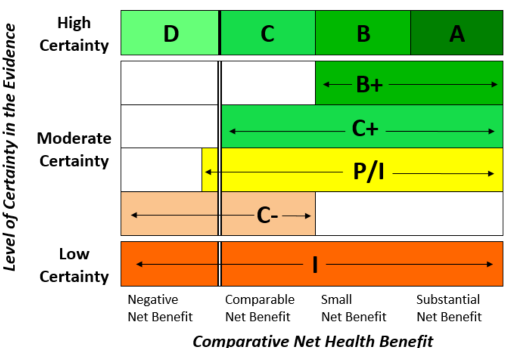
ICER Evidence Rating

We used the ICER Evidence Rating Matrix (see Figure D1) to evaluate the evidence for a variety of outcomes. The evidence rating reflects a joint judgment of two critical components:

The magnitude of the difference between a therapeutic agent and its comparator in "net health benefit" – the balance between clinical benefits and risks and/or adverse effects AND

The level of certainty in the best point estimate of net health benefit.⁸⁵

Figure D1. ICER Evidence Rating Matrix



Comparative Clinical Effectiveness

A = "Superior" - High certainty of a substantial (moderate-large) net health benefit

B = "Incremental" - High certainty of a small net health benefit

C = "Comparable"- High certainty of a comparable net health benefit

D = "Negative"- High certainty of an inferior net health benefit

B+ = "Incremental or Better" - Moderate certainty of a small or substantial net health benefit, with high certainty of at least a small net health benefit

C+ = "Comparable or Better" - Moderate certainty of a comparable, small, or substantial net health benefit, with high certainty of at least a comparable net health benefit

P/I = "**Promising but Inconclusive**" - Moderate certainty of a comparable, small, or substantial net health benefit, and a small (but nonzero) likelihood of a negative net health benefit

C- = "Comparable or Inferior" - Moderate certainty that the point estimate for comparative net health benefit is either comparable or inferior

I = "Insufficient" - Any situation in which the level of certainty in the evidence is low

Trial	Arm	Ν	Age, Mean Years (SD)	Female, n (%)	History of Migraine, Mean Years (SD)	Migraine Attacks/ Month in Past 3 Months, Mean (SD)
			Lasmiditan			
	Lasmiditan 200mg	609	41.4 (12.0)	515 (84.6)	18.9 (13.1)	5.3 (2.3)
SAMURAI ²³	Lasmiditan 100mg	630	42.2 (11.7)	512 (81.3)	19.7 (13.0)	5.1 (1.8)
	Placebo	617	42.4 (12.3)	525 (85.1)	19.3 (12.7)	5.1 (1.8)
	Lasmiditan 200mg	649	41.8 (12.4)	536 (82.6)	17.6 (12.6)	5.3 (1.9)
SPARTAN ²⁴	Lasmiditan 100mg	635	43.4 (12.6)	539 (84.9)	19.2 (13.6)	5.3 (1.9)
SFANTAN	Lasmiditan 50mg	654	42.8 (13.2)	554 (84.7)	18.6 (12.9)	5.2 (2.0)
	Placebo	645	42.6 (12.9)	545 (84.5)	17.9 (12.8)	5.5 (2.4)
	Lasmiditan 200mg	71	39.5 (10.3)	65 (91.5)		3.3 (1.9)
For the 2012 ²⁵	Lasmiditan 100mg	82	42.0 (10.6)	68 (82.9)	ND	3.3 (1.7)
Farkkila 2012 ²⁵	Lasmiditan 50mg	82	40.4 (12.5)	69 (84.1)	NR	3.3 (1.6)
	Placebo	86	40.5 (10.3)	75 (87.2)		3.1 (1.7)
			Rimegepant			
Charles 204 ²⁷	Rimegepant 75mg	543	41.9 (12.3)	464 (85.5)	ND	4.8 (1.7)
Study 301 ²⁷	Placebo	541	41.3 (12.1)	463 (85.6)	NR	4.7 (1.8)
Study 302 ²⁶	Rimegepant 75mg	537	40.2 (11.9)	479 (89.2)	NR	4.5 (1.9)
	Placebo	535	40.9 (12.1)	472 (88.2)	INIT	4.6 (1.8)
Study 303 ²⁸	Rimegepant 75mg	669	40.3 (12.1)	568 (84.9)	NR	4.6 (1.8)
	Placebo	682	40.0 (11.9)	579 (84.9)	INIT	4.5 (1.8)
	Rimegepant 75mg	91	38.5 (11.9)	81 (89.0)		3.9 (1.7)*
Marcus 2014 ²⁹	Sumatriptan 100mg	109	40.6 (10.5)	91 (83.5)	NR	4.1 (1.8)*
	Placebo	229	37.9 (11.4)	196 (85.6)		4.0 (1.8)*
			Ubrogepant			
	Ubrogepant 100mg	485	40.6 (12.0)	418 (86.2)	18.9 (12.3)	4.6 (1.8)
ACHIEVE I ³¹	Ubrogepant 50mg	466	40.1 (11.7)	418 (89.7)	17.9 (11.9)	4.6 (1.9)
	Placebo	485	40.9 (11.7)	430 (88.7)	19.1 (12.3)	4.4 (1.7)
	Ubrogepant 50mg	488	41.2 (12.5)	444 (91.0)	18.1 (12.3)	4.4 (1.8)
ACHIEVE II ³⁰	Ubrogepant 25mg	478	41.6 (12.4)	431 (90.2)	18.9 (12.2)	4.8 (1.8)

Table D1. Key Baseline Characteristics of Patients in the Trials of Lasmiditan, Rimegepant, Ubrogepant and Triptans

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	Placebo	499	41.7 (12.1)	442 (88.6)	19.2 (12.6)	4.6 (1.8)
	Ubrogepant 100mg	102	41.9 (11.0)	90 (88.2)		
V 201 C ³²	Ubrogepant 50mg	106	40.7 (12.3)	92 (86.8)	ND	
Voss 2016 ³²	Ubrogepant 25mg	104	41.4 (11.5)	91 (87.5)	NR	NR
	Placebo	113	40.5 (11.7)	99 (87.6)		
			Triptans			
Diener 2002 ³⁴	Eletriptan 40mg	210	40 (11.0)	181 (86.2)	Range: 10.9 - 23.3	Range: 6.7 - 8.0
	Placebo	106	42 (11.0)	91 (85.8)	Kange. 10.9 - 25.5	Kalige. 0.7 - 0.0
Steiner 2003 ⁴⁸	Eletriptan 40mg	392	40.3 (10.4)	345 (88.0)	16.6 (12.1)	2.5 (1.3)
Stemer 2005	Placebo	144	39.9 (10.6)	124 (86.0)	16.2 (12.1)	2.6 (1.3)
Garcia-Ramos 2003 ³⁶	Eletriptan 40mg	192	36.3 (11.1)	152 (79)	10.3 (9.7)	2.8 (NR)
	Placebo	92	36.4 (11.1)	75 (82)	11.9 (10.4)	2.8 (NR)
The EMSASI Study Group 2004 ⁵¹	Sumatriptan 50mg	226	38.2 (12.5)	182 (80.5)	With aura: 19.4 (14.0) Without aura: 16.0 (12.7)	NR
The EIVISASI Study Group 2004-2	Placebo	222	38.3 (12.2)	180 (81.1)	With aura: 18.9 (13.0) Without aura: 15.1 (11.6)	NK
Diener 2004 ³³	Sumatriptan 50mg	135	43.7 (12.1)	111 (82.2)	ND	NR
Diener 2004-4	Placebo	152	41.9 (11.7)	127 (83.6)	NR	INK
C 1 2222 ³⁷	Sumatriptan 100mg	504	38.0 (10.6)	424 (84.0)	ND	2.8 (1.4)
Geraud 2000 ³⁷	Placebo	56	37.9 (9.7)	49 (86.0)	NR	2.7 (1.3)
	Sumatriptan 100mg	462	41.5 (11.2)	389 (84.2)		
Sheftell 2005 ⁴⁶ -Study 1	Sumatriptan 50mg	448	41.6 (10.8)	380 (84.9)	NR	NR
	Placebo	456	41.2 (10.8)	401 (87.9)		
	Sumatriptan 100mg	440	40.2 (10.8)	361 (82.0)		
Sheftell 2005 ⁴⁶ - Study 2	Sumatriptan 50mg	454	39.9 (10.8)	387 (85.2)	NR	NR
	Placebo	436	39.2 (10.5)	378 (86.7)		
Havanka 2000 ³⁹	Sumatriptan 100mg	98	NR	89 (89.0)	NR	NR
	Placebo	91		81 (89.0)	IVIN	
Smith 2005 ⁴⁷	Sumatriptan 50mg	229	41.2 (11.3)	208 (90.8)	21.5 (NR)	NR
- Smith 2005	Placebo	242	41.2 (10.2)	214 (88.4)	20.0 (NR)	
Tfelt-Hansen 1995 ⁴⁹	Sumatriptan 100mg	139	39 (Range: 18 - 58)	108 (77.7)	18 (Range: 1 - 50)	3.3 (Range: 2 - 6)
	Placebo	137	39 (Range: 18 - 63)	106 (77.4)	19 (Range: 1 - 51)	3.4 (Range: 2 - 8)

	Sumatriptan 100mg	46	40 (10.0)	39 (84.8)		
Myllyla 1998 ⁴³	Placebo	48	39 (9.5)	45 (93.8)	NR	NR
	Sumatriptan 100mg	388	39.2 (10.1)	309 (79.6)		
Tfelt-Hansen 1998 ⁵⁰	Placebo	160	38.3 (10.3)	132 (82.5)	NR	NR
	Sumatriptan 100mg	194	42.0 (10.5)	162 (83.5)		
Dowson 2002 ³⁵	Placebo	99	40.2 (10.1)	88 (88.9)	NR	NR
	Sumatriptan 50mg	144	41.1 (9.9)	130 (90.3)		
Kudrow 2005 ⁴⁰	Placebo	141	39.0 (9.8)	124 (87.9)	NR	NR
	Sumatriptan 50mg					
Lines 2001 ⁴¹	Placebo	No baseli	ine characteristics repo	orted		
Nappi 1994 ⁴⁴	Sumatriptan 100mg	158	38 (9)	120 (76)	Median: 17.5	NR
Mappi 1994	Placebo	86	38 (11)	68 (79)	Median: 18.0	
	Sumatriptan 100mg	298	40.0	247 (82.9)	17.2 (NR)	
Pfaffenrath 1998 ⁴⁵	Sumatriptan 50mg	303	40.4	266 (87.8)	17.2 (NR)	NR
	Placebo	99	40.4 (10.7)	80 (80.8)	18.0 (NR)	
	Sumatriptan 100mg	148	42 (10)	128 (86.5)	Median: 20.0	
Oral Sumatriptan International Multiple-Dose Study Group 1991 ⁵²	Placebo	84	40 (10)	70 (83.3)	Median: 18.0	NR
	Eletriptan 40mg	822	41.1 (10.8)	716 (87.0)	13.4 (11.3)	2.7 (1.3)
Mathew 2003 ⁴²	Sumatriptan 100mg	831	41.8 (10.4)	715 (86.0)	14.0 (11.2)	2.7 (1.3)
	Placebo	419	41.6 (10.6)	365 (87.0)	13.6 (11.5)	2.8 (1.4)
	Eletriptan 40mg	136	41 (11)	115 (84.6)		
Goadsby 2000 ³⁸	Eletriptan 20mg	144	40 (11)	118 (81.9)	NR	NR
	Sumatriptan 100mg	129	40 (10)	108 (83.7)	INIT	INT
	Placebo	142	41 (10)	113 79.6)		
Kolodny 2004 ⁵³	Sumatriptan 50mg	285			ross group. Average age in stud	ly is 40 years, and patients
	Placebo	288	were predominantly	female (86%)		
Pini 1995 ⁵⁴	Sumatriptan 100mg	151	37	186 (78)		4 per month (45%); 1-3 per month (48%); Daily (2.6%)
	Placebo	87				4 per month (42%); 1-3 per month (47%); Daily (9%)

Page 120 Return to Table of Contents mg: milligram, n: number of participants, N: total number of particiants, NR: not reported, SD: standard deviation *in the past 12 months

Trial	Arm		Headache P	ain Intensity,	n (%)		Baseline Symptoms, n (%)					MBS, n (%)			
		N	Severe	Moderate	Mild	N	Phono- phobia	Photo- phobia	Nausea	Vomiting	N	Phono- phobia	Photo- phobia	Nausea	
						Lasmic	ditan								
SAMURAI ²³	Lasmiditan 200mg	518	148 (28.6)	355 (68.5)	15 (2.9)	518	322 (62.2)	391 (75.5)	232 (44.8)	NR	481	96 (20.0)	267 (55.5)	118 (24.5)	
	Lasmiditan 100mg	503	132 (26.2)	366 (72.8)	5 (1.0)	503	303 (60.2)	386 (76.7)	210 (41.7)		469	117 (24.9)	237 (50.5)	115 (24.5)	
	Placebo	524	145 (27.7)	370 (70.6)	9 (1.7)	524	327 (62.4)	416 (79.4)	221 (42.2)		488	104 (21.3)	269 (55.1)	115 (23.6)	
SPARTAN ²⁴	Lasmiditan 200mg	528	147 (27.8)	374 (70.8)	7 (1.3)	528	326 (61.7)	397 (75.2)	219 (41.5)	NR	483	110 (20.8)	269 (50.9)	104 (19.7)	
	Lasmiditan 100mg	532	159 (29.9)	364 (68.4)	9 (1.7)	532	345 (64.8)	406 (76.3)	235 (44.2)		500	110 (20.7)	276 (51.9)	114 (21.4)	
	Lasmiditan 50mg	556	152 (27.3)	392 (70.5)	12 (2.2)	556	330 (59.4)	427 (76.8)	245 (44.1)		512	108 (19.4)	277 (49.8)	127 (22.8)	
	Placebo	540	165 (30.6)	369 (68.3)	5 (0.9)	540	353 (65.4)	419 (77.6)	249 (46.1)		514	119 (22.0)	268 (49.6)	127 (23.5)	
Farkkila 2012 ²⁵	Lasmiditan 200mg	71	34 (48.0)†	36 (51.0)†	0 (0)	71	48 (66.4)*	57 (79.8)*	48 (66.6)*	1 (0.1)*	NR				
	Lasmiditan 100mg	82	33 (40.0)	49 (60.0)	0 (0)	82	52 (63.2)*	61 (73.9)*	43 (51.4)*	3 (2.6)*					
	Lasmiditan 50mg	82	32 (39.0)†	49 (60.0)†	0 (0)	82	56 (68.2)*	59 (72.0)*	48 (58.1)*	3 (2.8)*					
	Placebo	86	34 (40.0)†	51 (59.0)†	0 (0)	86	56 (64.2)*	66 (76.3)#	52 (60.4)*	8 (8.7)*					

Table D2. Baseline Characteristics of Treated Migraine Attacks in the Trials of Lasmiditan, Rimegepant, Ubrogepant, and Triptans

Rimegepant														
Study 301 ²⁷	Rimegepant 75mg	NR				NR					543	89	302	152
												(16.4)‡	(55.6)‡	(28.0)‡
	Placebo										541	101	302	138
					1							(18.7)‡	(55.8)‡	(25.5)‡
Study 302 ²⁶	Rimegepant 75mg	537	537 (100)#		0 (0)	537	362	489	355	NR	537	72	277	169
			505 (400) #		0 (0)		(67.4)	(91.1)	(66.1)			(13.4)	(51.6)	(31.5)
	Placebo	535	535 (100)#		0 (0)	535	374	477	336		535	92	279	148
Study 303 ²⁸	Dimogenent 75mg	669	660 (100)#		0 (0)	NR	(69.9)	(89.2)	(62.8)		669	(17.2) 108	(52.1) 359	(27.7) 189
Study 303	Rimegepant 75mg		669 (100)#			INK								
R.d	Placebo	682	682 (100)#		0 (0)						682	101	374	195
Marcus 2014 ²⁹	Rimegepant 75mg	91	91 (100)#		0 (0)	NR					NR			
2014	Sumatriptan 100mg	109	109 (100)#		0 (0)									
	Placebo	229	229 (100)#		0 (0)									
	Flacebo	229	229 (100)#			 Ubroge	anant							
ACHIEVE I ³¹	Ubrogepant	448	160 (35.7)	288 (64.3)	0 (0)	448	360	391	274	18 (4.0)	448	116	246	86
	100mg	0	100 (33.7)	200 (04.3)	0 (0)	0	(80.4)	(87.3)	(61.2)	10 (4.0)	0	(25.9)		(19.2)
	Ubrogepant 50mg	423	163 (38.5)	260 (61.5)	0 (0)	423	315	390	237	27 (6.4)	423	82	248	90
			()	()	- (-)		(74.5)	(92.2)	(56.0)			(19.4)		(21.3)
	Placebo	456	169 (37.1)	287 (62.9)	0 (0)	456	362	416	292	26 (5.7)	456	98	254	102
							(79.4)	(91.2)	(64.0)			(21.5)	(55.7)	(22.4)
ACHIEVE II ³⁰	Ubrogepant 50mg	488	175 (37.7)	289 (62.3)	0 (0)	488	374	420	297	21 (4.5)	488	115	265	83
							(80.6)	(90.5)	(64.0)			(24.8)	(57.1)	(17.9)
	Ubrogepant 25mg	478	178 (40.9)	257 (59.1)	0 (0)	478	353	399	284	19 (4.4)	478	102	257	75
							(81.1)	(91.7)	(65.3)			(23.4)	(59.1)	(17.2)
	Placebo	499	198 (43.4)	258 (56.6)	0 (0)	499	370	404	279	22 (4.8)	499	136	245	75
Non 2016 ³²	Libus sevent	102			0 (0)	100	(81.1)	(88.6)	(61.2)	4 (2.0)	ND	(29.8)	(53.7)	(16.4)
Voss 2016 ³²	Ubrogepant 100mg	102	27 (26.5)	75 (73.5)	0 (0)	102	79 (77.5)	85 (83.3)	58 (56.9)	4 (3.9)	NR			
	Ubrogepant 50mg	106	31 (29.2)	75 (70.8)	0 (0)	D)10678 (72.6)88 (83.0)57 (53.8)5 (4.7)								
	Ubrogepant 25mg	104	38 (36.5)	65 (62.5)	0 (0)	104	82 (78.8)	94 (90.4)	53 (51.0)	2 (1.9)				

	Placebo	113	41 (36.3)	72 (65.7)	0 (0)	113	87 (77.0)	100 (88.5)	65 (57.5)	2 (1.8)	
						Tript	ans				
Diener 2002 ³⁴	Eletriptan 40mg	210	97 (46.2)	113 (53.8)	0 (0)	210	155 (73.8)	153 (72.9)	143 (68.1)	21 (10.0)	NR
2002	Placebo	106	51 (48.1)	55 (51.9)	0 (0)	106	75 (70.8)	80 (75.5)	72 (67.9)	12 (11.3)	
Steiner	Eletriptan 40mg	392	185 (47.0)	207 (53.0)	NR	392	290 (74.0)	306 (78.0)	255 (65.0)	NR	NR
2003 ⁴⁸	Placebo	144	67 (46.0)	77 (54.0)		144	103 (71.0)	114 (79.0)	87 (60.0)		
Garcia-	Eletriptan 40mg	192	102 (53)	90 (47)#	NR	192	NR		102 (52)	NR	NR
Ramos 2003 ³⁶	Placebo	92	42 (46)	50 (54)#		92			47 (51)		
The EMSASI Study	Sumatriptan 50mg	226	113 (50.0)	113 (50.0)	NR	224	129 (57.6)	148 (66.1)	NR	39 (17.4)	NR
Group 2004 ⁵¹	Placebo	222	107 (48.2)	115 (51.2)		222	128 (57.7)	138 (62.2)		33 (14.9)	
Diener 2004 ³³	Sumatriptan 50mg	135	135 (100)#		0 (0)	NR					NR
2004	Placebo	152	152 (100)#		0 (0)						
Geraud 2000 ³⁷	Sumatriptan 100mg	503	192 (38.0)	310 (62.0)	1 (0.2)	503	356 (70.7)	346 (68.8)	273 (54.3)	NR	NR
2000	Placebo	55	18 (33.0)	37 (67.0)	0 (0)	55	43 (78.2)	42 (76.4)	25 (45.5)		
Sheftell	Sumatriptan 100mg	488	488 (100)#		0 (0)	NR					NR
2005 ⁴⁶ - Study 1	Sumatriptan 50mg	494	494 (100)#		0 (0)						
	Placebo	495	495 (100)#		0 (0)						
Sheftell 2005 ⁴⁶ -	Sumatriptan 100mg	485	485 (100)#		0 (0)	NR					NR
Study 2	Sumatriptan 50mg	496	496 (100)#		0 (0)						

	Placebo	494	494 (100)#		0 (0)						
Havanka	Sumatriptan	98	68 (69.0)	31 (31.0)	0 (0)	98	NR		77 (78.0)	NR	NR
2000 ³⁹	100mg										
2000	Placebo	91	69 (75.0)	23 (75.0)	0 (0)	91			72 (79.0)		
Smith	Sumatriptan	229	229 (100)#		0 (0)	NR					NR
2005 ⁴⁷	50mg				a (a)						
	Placebo	242	242 (100)#		0 (0)						
Tfelt-	Sumatriptan	122	40 (32.8)	82 (67.2)	0 (0)	122	NR		84 (68.9)	10 (8.2)	NR
Hansen 1995 ⁴⁹	100mg	120	42 (22 2)	OA(CC T)	0 (0)	120			01 (C1 2)	11 (0 7)	
1995**	Placebo	126	42 (33.3)	84 (66.7)	0 (0)	126	20/45	20/45	81 (64.3)	11 (8.7)	ND
Myllyla	Sumatriptan 100mg	46	46 (100)#		0 (0)	46	30/45 (66.7)	38/45 (84.4)	20 (43.5)	2/45 (4.4)	NR
1998 ⁴³	Placebo	48	48 (100)#		0 (0)	48	33 (68.8)	42 (87.5)	20 (41.7)	4 (8.3)	
Tfelt-	Sumatriptan	388	196 (50.5)	191 (49.2)	0 (0)	NR	33 (00.0)	42 (07.3)	20 (41.7)	4 (0.5)	NR
Hansen	100mg	500	190 (90.9)	191 (49.2)	0 (0)						
1998 ⁵⁰	Placebo	160	84 (52.5)	75 (46.9)	0 (0)						
	Sumatriptan	194	82 (42.3)	111 (57.2)	0 (0)	NR					NR
Dowson 2002 ³⁵	100mg										
200200	Placebo	99	32 (32.3)	67 (67.7)	0 (0)						
	Sumatriptan	144	47 (32.9)	96 (67.1)	0 (0)	144	104	125	95 (66.4)	3 (2.1)	NR
Kudrow	50mg						(72.7)	(87.4)			
2005 ⁴⁰	Placebo	141	56 (39.7)	85 (60.3)	0 (0)	141	106	134	97 (68.8)	7 (5.0)	
							(75.2)	(95.0)			
	Sumatriptan	No ba	aseline charac	teristics repor	ted						
Lines 2001 ⁴¹	50mg Placebo										
		158	77 (48.7)	71 (44.9)	10 (6.4)	NR					NR
Nappi	Sumatriptan 100mg	128	// (48./)	71 (44.9)	10 (0.4)	INK					NK
1994 ⁴⁴	Placebo	86	40 (46.5)	41 (47.7)	5 (5.8)						
	Sumatriptan	298	40 (40.3) 277 (93.0)	41 (47.7)	NR	NR					NR
Pfaffenrath	100mg	290	277 (95.0)		INIT	INIT					INT
1998 ⁴⁵	Sumatriptan	303	285 (94.1)		NR						
	50mg										

	Placebo	99	91 (91.9)		NR						
Oral Sumatriptan Internation	Sumatriptan 100mg	148	52 (35.1)	79 (53.4)	17 (11.5)	NR					NR
al Multiple- Dose Study Group 1991 ⁵²	Placebo	84	27 (32.1)	51 (60.8)	6 (7.1)						
	Eletriptan 40mg	822	321 (39.0)	501 (61.0)	0 (0)	822	526 (64.0)	592 (72.0)	510 (62.0)	NR	NR
Mathew 2003 ⁴²	Sumatriptan 100mg	831	341 (41.0)	490 (59.0)	0 (0)	831	557 (67.0)	624 (75.0)	516 (62.0)		
	Placebo	419	172 (41.0)	247 (59.0)	0 (0)	419	269 (64.0)	315 (75.0)	269 (64.0)		
	Eletriptan 40mg	136	63 (46.3)	68 (50.0)	NR	136	NR		83 (61.0)	11 (8.1)	NR
Goadsby	Eletriptan 20mg	144	62 (43.1)	82 (56.9)		144			91 (63.2)	8 (5.6)	
2000 ³⁸	Sumatriptan 100mg	129	56 (43.4)	71 (55.0)		129			82 (63.6)	14 (10.9)	
	Placebo	142	66 (46.5)	74 (52.1)		142			90 (63.4)	12(8.5)	

MBS: most bothersome symptom, mg: milligram, n: number of participants, N: total number of participants, NR: not reported.

* Data are digitized and should be interpreted with caution,

⁺ due to missing data, percentages do not add up to 100%,

‡ historical, # assumption made based on study protocol

Trial (NCT) & Author	Design and duration of follow up	Interventions & dosing procedure	Inclusion Criteria	Exclusion Criteria		
Aution		La	smiditan			
SAMURAI (NCT02439320) Kuca 2018 ²³	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit 7 days after treated migraine attack	Lasmiditan (100 or 200mg) vs placebo - study medication to be taken within 4-hours of migraine onset (moderate to severe pain); second dose for rescue allowed 2- 24 hours after first dose	Adults ≥18 years; ≥1-year history of disabling migraines with or without aura; onset before age 50; 3-8 migraine attacks/month (<15 headache days/month)	History of chronic migraine or other forms of primary or secondary headache disorder in past 12 months; ≥15 headache days/month within past 12 months; initiation of or change in migraine preventative medication within 3 months; known coronary artery disease; clinically significant arrythmia; uncontrolled hypertension; condition increasing risk of seizures		
SPARTAN (NCT02605174) Goadsby 2019 ²⁴	Prospective, randomized, double- blind, placebo controlled, multicentre phase III, single attack study; follow-up visit 7 days after treated migraine attack	Lasmiditan (50, 100, or 200mg) vs placebo - study medication to be taken within 4-hours of migraine onset (moderate to severe pain); second dose for rescue or recurrence allowed 2-24 hours after first dose	Adults ≥18 years; ≥1-year history of disabling migraines with or without aura; MIDAS score ≥11; onset before age 50; 3-8 migraine attacks/month (<15 headache days/month)	History of chronic migraine; other forms of primary or secondary headache disorder; ≥15 headache days/month within past 12 months; condition increasing risk of seizures; recurrent dizziness or vertigo; diabetes mellitus with complications; orthostatic hypotension with syncope; renal or hepatic impairment; current SUD within past 3 years; imminent risk of suicide or suicide attempt within past 6 months		

Table D3. Study Designs of the Trials on Lasmiditan, Rimegepant, Ubrogepant

Trial (NCT) & Author	Design and duration of follow up	Interventions & dosing procedure	Inclusion Criteria	Exclusion Criteria
Farkkila 2012 ²⁵	Randomized, double- blind, parallel-group, multicentre, single attack, dose-ranging study (Phase II); follow- up visit within 14 days of treated migraine attack	Lasmiditan (50, 100, 200, or 400mg) vs placebo - study medication to be taken within 4- hours of migraine onset (moderate to severe pain); second dose for rescue allowed (excl. triptans or ergotamines) 2-hours after first dose	Adults ≥18 years; ≥1-year history of acute migraines with or without aura; onset before age 50; 1-8 migraine attacks/month	Use of migraine prophylaxis (unless discontinued at least 15 days prior to screening), vasoactive drugs, serotonin reuptake inhibitors, or known cyto chrome P450 inhibitors
		Rin	negepant	
Study 301 (NCT03235479) - not yet published Lipton 2018 ²⁷	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit within 7 days of treated migraine attack	Rimegepant (75mg) vs placebo; rescue medication was allowed within 24-hours	Adults ≥18 years of age; ≥1-year history of migraine; 2-8 migraine attacks/month (moderate to severe intensity); <15 headache days/month within the past 3 months; patients receiving preventative migraine medications had to be receiving stable dose for at least 3 months before trial entry	HIV; uncontrolled, unstable or recently diagnosed CVD; patients with MI, ACS, PCI, cardiac surgery, stroke, or TIA within 6 months of screening; uncontrolled hypertension or diabetes; current diagnosis of major depression, other pain syndromes, psychiatric conditions, dementia, or significant neurologic conditions; history of GI surgery or disease that causes malabsorption; SUD within past 12 months

Trial (NCT) & Author	Design and duration of follow up	Interventions & dosing procedure	Inclusion Criteria	Exclusion Criteria
Study 302 (NCT03237845) Lipton 2019 ²⁶	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit within 7 days of treated migraine attack	Rimegepant (75mg) vs placebo - study medication to be taken when migraine of moderate to severe intensity occurred; use of second dose as rescue medication was allowed within 24-hours	Adults ≥18 years of age; ≥1-year history of migraine with or without aura; onset before age 50; 2-8 migraine attacks/month (moderate to severe intensity); <15 days/month with headache within the past 3 months; Patients receiving preventative migraine medications had to be receiving stable dose for at least 3 months before trial entry	History of any clinically significant or unstable medical condition, including alcohol or drug abuse and substance-use disorder; Use of any biologic investigational agents within 90 days of baseline visit; received nonbiologic investigational agents within 30 days before baseline visit
Study 303 (NCT03461757) Croop 2019 ²⁸	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit within 7 days of treated migraine attack	Rimegepant (75mg) vs placebo - study medication to be taken when migraine attack of moderate to severe intensity occurred; rescue medications (eg, aspirin, ibuprofen, acetaminophen [up to 1000 mg/day], naproxen[or any other NSAIDs], antiemetics, or baclofen) after 2-hours postdose	Adults ≥18 years of age; ≥1-year history of migraine with or without aura; onset before age 50; 2-8 migraine attacks/month (moderate to severe intensity); <15 days per month with headache within the past 3 months	SUD within past 12 months; history of drug or other allergy that made them unsuitable for participation; ECG or laboratory test findings that raised safety or tolerability concerns
Marcus 2014 ²⁹	Randomized, double- blind, multicentre, placebo-controlled, phase II, single attack study; follow-up visit within 7 days of treated migraine attack	Rimegepant (10, 25, 75, 150, 300, or 600mg) vs sumatriptan (100mg) and placebo - study medication to be taken at onset of moderate to severe migraine; use of rescue medication (aspirin, ibuprofen, acetaminophen, NSAIDs, anti-emetics, or baclofen) allowed 2-hours post-dose	Adults aged 18-65 years; ≥1-year history of migraine with or without aura; onset before age 50; duration of migraine attack 4-72 hours if untreated; 2-7 attacks/month (moderate to severe intensity) in 3 months prior to study; < 15 headache days/month in previous 3 months	<u>General</u> : History of stroke/transient ischemic attacks, ischemic heart disease, coronary artery vasospasm, other significant underlying CVD, uncontrolled hypertension or diabetes, HIV; current diagnosis of major depression, other pain syndromes, psychiatric conditions, dementia, or significant neurological disorders, other than migraine; SUD within the past 12 months. <u>For sumatriptan</u> : history of basilar-type or hemiplegic migraine; nonresponse to triptans

Trial (NCT) & Author	Design and duration of follow up	Interventions & dosing procedure	Inclusion Criteria	Exclusion Criteria
		Ub	rogepant	
ACHIEVE I (NCT02828020) Dodick 2018 ³¹	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit within 7 days of treated migraine attack	Ubrogepant (50 or 100mg) vs placebo, second dose or rescue medication allowed in patients with inadequate response or headache recurrence	Adults 18-75 years old; ≥1-year history of migraines with or without aura; onset before age 50; duration of migraine attack 4-72h and separated by ≥48h; 2-8 migraine attacks/month with moderate to severe headache pain in previous 3 months	Taken medication for acute treatment of headache on ≥10 days/month in previous 3 months; history of aura with diplopia or impairment of level of consciousness, hemiplegic or retinal migraine; current diagnosis of new persistent daily headache, trigeminal autonomic cephalgia, or painful cranial neuropathy; required hospital treatment of a migraine attack ≥3 times in previous 6 months; chronic non- headache pain condition requiring daily pain medication; history of malignancy in the prior 5 years; history of any prior GI conditions that may affect the absorption or metabolism; history of hepatitis within previous 6 months
ACHIEVE II (NCT02867709) Lipton 2019 ³⁰	Randomized, double- blind, placebo- controlled, multicentre, phase III, single attack study; follow-up visit within 7 days of treated migraine attack	Ubrogepant (25, 50, or 100mg) vs placebo, second dose or rescue medication allowed in patients with inadequate response or headache recurrence	Adults 18-75 years old; ≥1-year history of migraines with or without aura; onset before age 50; duration of migraine attack 4-72 hours and separated by ≥48 hours; 2-8 migraine attacks/month (moderate to severe intensity) in previous 3 months	Taken medication for acute treatment of headache on ≥10 days/month in the previous 3 months; history of migraine aura with diplopia or impairment of level of consciousness, hemiplegic or retinal migraine; current diagnosis of new persistent daily headache, trigeminal autonomic cephalgia, or painful cranial neuropathy; required hospital treatment of a migraine attack ≥3 times in previous 6 months; chronic non- headache pain condition requiring daily pain medication; history of malignancy in the prior 5 years; history of any prior GI conditions that may affect the absorption or metabolism; history of hepatitis within previous 6 months

Trial (NCT) & Author	Design and duration of follow up	Interventions & dosing procedure	Inclusion Criteria	Exclusion Criteria
Voss 2016 ³²	Randomized, double- blind, placebo- controlled, Phase lib, single attack study; follow-up visit five days post-treatment	Ubrogepant (1, 10, 25, 50, or 100mg) vs placebo - study drug to be taken to treat a migraine of moderate to severe intensity; non-study medication allowed as rescue or recurrence treatment	Adults ≥18 years; ≥1-year history of acute migraines with or without aura; onset before age 50; 1-8 migraine attacks/month	Difficulty distinguishing migraine attacks from tension type headaches; uncontrolled hypertension; basilar-type or hemiplegic migraine headache; >15 headache days/month or had taken medication for acute headache on >10 days/month in the three months prior to screening; acute attack within past 2 months that required inpatient or ER treatment; use of an opioid or barbiturate for migraine in the past 2 months; recent change in dose of migraine- prophylactic medication

ACS: acute coronary syndrome, CVD: cardiovascular disease, ECG: echocardiogram, excl: excluding, GI: gastrointestinal, HIV: human immunodeficiency virus, MI: myocardial infarction, PCI: percutaneous coronary intervention, SUD: substance use disorder, TIA: transient ischemic attack

Trial	Comparable Groups	Non- differential Follow-up	Patient/ Investigator Blinding (Double-Blind)	Clear Definition of Intervention	Clear Definition of Outcomes	Selective outcome reporting	Measurements Valid	Intention- to-Treat Analysis	Approach to Missing Data	UPSTF Rating
				Lasmidi	itan					
SAMURAI ²⁴	Yes	No	Yes	Yes	Yes	No	Yes	mITT†	N/A	good
SPARTAN ²³	Yes	No	Yes	Yes	Yes	No	Yes	mITT†	N/A	good
Farkkila 2012 ²⁵	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
	_			Rimege	pant					
Study 301 ²⁷	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	*
Study 302 ²⁶	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
Study 303 ²⁸	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
Marcus 2014 ²⁹	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
				Ubroger	pant					
ACHIEVE I ³¹	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
ACHIEVE II ³⁰	Yes	No	Yes	Yes	Yes	No	Yes	mITT	N/A	good
Voss 2016 ³²	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
				Tripta	ns					
Diener 2002 ³⁴	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Steiner 2003 ⁴⁸	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Garcia-Ramos 2003 ³⁶	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
The EMSASI Study Group 2004 ⁵¹	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Diener 2004 ³³	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Geraud 2000 ³⁷	Yes	No	Yes	Yes	Yes	No	Yes	All-treated	N/A	good
Sheftell 2005 ⁴⁶ – Study 1	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Sheftell 2005 ⁴⁶ – Study 2	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good

Table D4. Quality Ratings for Trials of Lasmiditan, Rimegepant, Ubrogepant and Triptans

Trial	Comparable Groups	Non- differential Follow-up	Patient/ Investigator Blinding (Double-Blind)	Clear Definition of Intervention	Clear Definition of Outcomes	Selective outcome reporting	Measurements Valid	Intention- to-Treat Analysis	Approach to Missing Data	UPSTF Rating
Havanka 2000 ³⁹	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Smith 2005 ⁴⁷	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Tfelt-Hansen 1995 ⁴⁹	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Myllyla 1998 ⁴³	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Tfelt-Hansen 1998 ⁵⁰	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Dowson 2002 ³⁵	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Kudrow 2005 ⁴⁰	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Lines 2001 ⁴¹	‡	No	Yes	Yes	Yes	No	Yes	mITT	N/A	fair
Nappi 1994 ⁴⁴	Yes	No	Yes	Yes	Yes	No	Yes	Per- protocol	N/A	fair
Pfaffenrath 1998 ⁴⁵	Yes	No	Yes	Yes	Yes	No	Yes	Per- protocol	N/A	fair
Oral Sumatriptan International Multiple- Dose Study Group 1991 ⁵²	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Mathew 2003 ⁴²	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Goadsby 2000 ³⁸	Yes	No	Yes	Yes	Yes	No	Yes	ITT	N/A	good
Kolodny 2004 ⁵³	‡	No	Yes	Yes	Yes	No	Yes	mITT	N/A	fair
Pini 1995 ⁵⁴	Yes	No	Yes	Yes	Yes	No	Yes	NR	N/A	good

ITT: intention-to-treat, mITT: modified intention-to-treat, N/A: not applicable, USPSTF: US Preventive Services Task Force

*Data was only available in grey literature. Due to this, we did not assign an overall quality rating for the trials and were not able to assess selective outcome reporting. We will assign an overall quality rating and update quality categories where necessary upon publication of peer-reviewed results.

⁺ Primary outcomes were analyzed with a modified intention-to-treat and secondary outcomes with intention-to-treat.

‡Baseline characteristics were stated to be similar between both intervention arms, however specific values were not reported

Data included in the NMA

Table D5. Efficacy Outcomes at 2-hours

Trial	Arms	Headac	he Pain F	reedom	Head	ache Pain	Relief	Free of MBS			Ability to Function Normally		
ITIAI	Arms	n	Ν	%	n	Ν	%	n	N	%	n	N	%
				ĺ	Lasmiditan								
	Lasmiditan 200mg	167	518	32.2	330	555	59.5	196	481	40.7	180	555	32.4
SAMURAI ²⁴	Lasmiditan 100mg	142	503	28.2	334	562	59.4	192	469	40.9	181	562	32.2
	Placebo	80	524	15.3	234	554	42.2	144	488	29.5	119	554	21.5
	Lasmiditan 200mg	205	528	38.8	367	565	65.0	235	483	48.7	209	565	37.0
SPARTAN ²³	Lasmiditan 100mg	167	532	31.4	370	571	64.8	221	500	44.2	193	571	33.8
	Placebo	115	540	21.3	274	576	47.6	172	514	33.5	143	576	24.8
	Lasmiditan 200mg	13	69	18.8	35	69	50.7						
Farkkila 2012 ²⁵	Lasmiditan 100mg	11	81	13.6	52	81	64.2	NR			NR		
	Placebo	6	81	7.4	21	81	25.9						
				F	limegepant	t							
Study 301 ²⁷	Rimegepant 75mg	104	543	19.2	304	543	56.0	199	543	36.6	181	543	33.3
Study SUI	Placebo	77	541	14.2	247	541	45.7	150	541	27.7	118	541	21.8
Study 302 ²⁶	Rimegepant 75mg	105	537	19.6	312	537	58.1	202	537	37.6	175	537	32.6
Study 302	Placebo	64	535	12.0	229	535	42.8	135	535	25.2	125	535	23.4
Study 303 ²⁸	Rimegepant 75mg	142	669	21.2	397	669	59.3	235	669	35.1	225	669	38.1
5100 505	Placebo	74	682	10.9	295	682	43.3	183	682	26.8	176	682	25.8
	Rimegepant 75mg	27†	86†	31.4†	62	86	72.1						
Marcus 2014 ²⁹	Sumatriptan 100mg	35†	100†	35.0†	72	100	72.0	NR			NR		
	Placebo	31†	203†	15.3†	104	203	51.2						
				ι	Ibrogepant	t							
	Ubrogepant 100mg	95	448	21.2	275	448	61.4	169	448	37.7	193	448	42.9
ACHIEVE I ³¹	Ubrogepant 50mg	81	422	19.2	256	422	60.7	162	420	38.6	171	423	40.6
	Placebo	54	456	11.8	224	456	49.1	126	454	27.8	136	456	29.8
ACHIEVE II ³⁰	Ubrogepant 50mg	101	464	21.8	291	464	62.7	180	463	38.9	188	464	40.5

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Trial	Arms	Headac	he Pain F	reedom	Head	ache Pain	Relief	l	Free of N	IBS	Ability to Function Normally		
I I Idi	ATTIS	n	N	%	n	Ν	%	n	N	%	n	N	%
	Placebo	65	456	14.3	220	456	48.2	125	456	27.4	156	456	34.2
	Ubrogepant 100mg	26	102	25.5	60	102	58.8						
Voss 2016 ³²	Ubrogepant 50mg	22	106	20.8	60	106	56.6	NR			NR		
	Placebo	10	113	8.8	50	113	44.2						
					Triptans								
Diener 2002 ³⁴	Eletriptan 40mg	58	206	28.2	111	206	53.9	NR			NR		
Diener 2002	Placebo	5	102	4.9	21	102	20.6	INK			INK		
Steiner 2003 ⁴⁸	Eletriptan 40mg	115	359	32.0	229	359	63.8	ND			NR		
Steiner 2003	Placebo	8	135	5.9	30	135	22.2	NR			NK		
Garcia-Ramos 2003 ³⁶	Eletriptan 40mg	67	192	35.0	108	192	56.0	NR			NR		
Garcia-Ramos 2003	Placebo	17	91	19.0	28†	91†	31.0†	INK			INK		
The EMSASI Study Group	Sumatriptan 50mg	83	224	37.1	125	224	55.8	ND			NR		
2004 ⁵¹	Placebo	28	222	12.6	68	222	30.6	NR					
Diener 2004 ³³	Sumatriptan 50mg	33	135	24.4	66	135	48.8	NR			NR		
Dieliei 2004	Placebo	22	152	14.5	50	152	32.9	ININ			INK		
Geraud 2000 ³⁷	Sumatriptan 100mg	150	499	30.1	304	498	61.0	NR			NR		
Geradu 2000	Placebo	7	55	12.7	24	55	43.6	INK			INK		
	Sumatriptan 100mg	219	462	47.4	331	462	71.6						
Sheftell 2005 ⁴⁶ - Study 1	Sumatriptan 50mg	180	448	40.2	310	448	69.2	NR			NR		
	Placebo	84	456	18.4	208	456	45.6						
	Sumatriptan 100mg	207	440	47.0	318	440	72.3						
Sheftell 2005 ⁴⁶ - Study 2	Sumatriptan 50mg	178	454	39.2	293	454	64.5	NR			NR		
	Placebo	53	436	12.2	167	436	38.3						
Havanka 2000 ³⁹	Sumatriptan 100mg	NR	NR	NR	59	98	60.0	NR			NR		
	Placebo	NR	NR	NR	29	91	31.0						
Smith 2005 ⁴⁷	Sumatriptan 50mg	45	226	20.0	111	226	49.0	NR			NR		
-5mm 2005	Placebo	14	241	6.0	65	241	27.0						
Tfelt-Hansen 1995 ⁴⁹	Sumatriptan 100mg	NR	NR	NR	63	119	52.9	NR			NR		
	Placebo	NR	NR	NR	30	124	24.2				NIT.		

Trial	Arms	Headac	ne Pain F	reedom	Heada	ache Pain	Relief	F	ree of MB	S	Ability to Function Normally		
ITIdi	Arms	n	N	%	n	N	%	n	N	%	n	N	%
Myllyla 1998 ⁴³	Sumatriptan 100mg	21	42	50.0	33	42	78.6	NR			NR		
	Placebo	3	41	7.3	12	41	29.3	ININ			INIT		
Tfelt-Hansen 1998 ⁵⁰	Sumatriptan 100mg	127	387	32.8	239	387	61.8	NR					
	Placebo	15	159	9.4	64	159	40.3	INIX					
Dowson 2002 ³⁵	Sumatriptan 100mg	65	193	33.7	123	193	63.7	NR			NR		
	Placebo	15	99	15.2	42	99	42.4						
Kudrow 2005 ⁴⁰	Sumatriptan 50mg	NR	NR	NR	60	144	42.0	NR			NR		
	Placebo	NR	NR	NR	42	141	30.0						
Lines 2001 ⁴¹	Sumatriptan 50mg	NR	NR	NR	239	356	67.0	NR		NR NR			
	Placebo	NR	NR	NR	18	80	23.0						
Nappi 1994 ⁴⁴	Sumatriptan 100mg	34	142	24.0	73	142	51.4	NR			NR		
	Placebo	10	81	12.0	25	81	30.9	INK					
	Sumatriptan 100mg	NR	NR	NR	177†	298†	59.5†						
Pfaffenrath 1998 ⁴⁵	Sumatriptan 50mg	NR	NR	NR	180†	303†	59.5†	NR					
	Placebo	NR	NR	NR	28†	99†	28.1†						
Oral Sumatriptan	Sumatriptan 100mg	38	148	26.0	74	148	50.0						
International Multiple- Dose Study Group 1991 ⁵²	Placebo	4	84	5.0	16	84	19.0	NR			NR		
	Eletriptan 40mg	281	779	36.0	522	779	67.0						
Mathew 2003 ⁴²	Sumatriptan 100mg	216	799	27.0	472	799	59.0			NR			
	Placebo	21	404	5.0	105	404	26.0						
	Eletriptan 40mg	34	117	29.0	76	117	65.0						
Goadsby 2000 ³⁸	Sumatriptan 100mg	26	115	23.0	63	115	55.0			NR			
	Placebo	8	126	6.0	30	126	24.0						

MBS: most bothersome symptom, mg: milligram, n: number of participants, N: total number of participants, NR: not reported

⁺ Data are digitized and should be interpreted with caution

Table D6. Sustained Efficacy Outcomes

Trial	Arms	Sustained	Pain Freedom	, 24-hours	Sustaine	d Pain Fre hours	edom, 48-	Sustained Pain Relief, 24- hours		
		n	N	%	n	N	%	n	N	%
			asmiditan							
	Lasmiditan 200mg	103	555	18.6	111	565	19.6	NR		
SAMURAI ²³	Lasmiditan 100mg	83	562	14.8	86	571	15.1			
	Placebo	42	554	7.6	89	598	14.9			
	Lasmiditan 200mg	128	565	22.7	68	576	11.8	NR		
SPARTAN ²⁴	Lasmiditan 100mg	102	571	17.9	91	555	16.4			
	Placebo	77	576	13.4	42	554	7.6			
	Lasmiditan 200mg	NR			NR			NR		
Farkkila 2012 ²⁵	Lasmiditan 100mg									
	Placebo									
		R	imegepant							
Study 301 ²⁷	Rimegepant 75mg	76	543	14.0	90	669	13.5	211	543	38.9
Study SOI	Placebo	44	541	8.1	37	682	5.4	151	541	27.9
Study 302 ²⁶	Rimegepant 75mg	66	537	12.3	53	537	9.9	229	537	42.6
Study 502	Placebo	38	535	7.1	32	535	6.0	142	535	26.5
Study 303 ²⁸	Rimegepant 75mg	105	669	15.7	63	543	11.6	320	669	47.8
Study 505	Placebo	38	682	5.6	39	541	7.2	189	682	27.7
	Rimegepant 75mg	24	86	27.9	24	86	27.9	60	86	69.8
Marcus 2014 ²⁹	Sumatriptan 100mg	26	100	26.0	26	100	26.0	63	100	63.0
	Placebo	15	203	7.4	15	203	7.4	86	203	42.4
		U	brogepant							
	Ubrogepant 100mg	68	441	15.4	NR			165	434	38.0
ACHIEVE I ³¹	Ubrogepant 50mg	53	418	12.7				150	413	36.3
	Placebo	39	452	8.6				93	447	20.8
ACHIEVE II ³⁰	Ubrogepant 50mg	66	457	14.4	NR			165	449	36.7
	Placebo	37	451	8.2				93	443	21.0
Voss 2016 ³²	Ubrogepant 100mg	22	102	21.6	21	102	20.6	47	102	46.1

Trial	Arms	Sustaine	ustained Pain Freedom, 24-hours			Sustained Pain Freedom, 48- hours			Sustained Pain Relief, hours		
		n	N	%	n	N	%	n	N	%	
	Ubrogepant 50mg	16	106	15.1	15	106	14.2	48	106	45.3	
	Placebo	7	113	6.2	7	113	6.2	32	113	28.3	
			Triptans								
Diener 2002 ³⁴	Eletriptan 40mg	42*	209*	20.0*	NR			84*	209*	40.1*	
	Placebo	2*	104*	1.7*				7*	104*	7.0*	
Steiner 2003 ⁴⁸	Eletriptan 40mg	75	349	21.5	NR			151	345	43.8	
Stemer 2005	Placebo	6	134	4.5				14	131	10.7	
Garcia-Ramos 2003 ³⁶	Eletriptan 40mg	37	168	22.0	NR			64	168	38.0	
Garcia-Ramos 2003	Placebo	10	85	12.0				16	85	19.0	
	Sumatriptan 50mg	NR			NR			NR			
The EMSASI Study Group 2004 ⁵¹	Placebo										
	Sumatriptan 50mg	NR			NR			97	135	71.4	
Diener 2004 ³³	Placebo							101	152	66.4	
	Sumatriptan 100mg	NR			NR			195	498	39.2	
Geraud 2000 ³⁷	Placebo							14	55	25.5	
	Sumatriptan 100mg	107	426	25.1	NR			163	420	38.8	
Sheftell 2005 ⁴⁶ - Study 1	Sumatriptan 50mg	85	419	20.3				154	405	38.0	
	Placebo	46	449	10.2				92	446	20.6	
	Sumatriptan 100mg	108	424	25.5	NR			181	421	43.0	
Sheftell 2005 ⁴⁶ - Study 2	Sumatriptan 50mg	96	442	21.7				173	437	39.6	
	Placebo	21	430	4.9				69	429	16.1	
	Sumatriptan 100mg	NR			NR			44	98	44.0	
Havanka 2000 ³⁹	Placebo							20	91	22.0	
Smith 2005 ⁴⁷	Sumatriptan 50mg	25	226	11.0	NR			66	226	29.0	
<u></u>	Placebo	12	241	5.0				41	241	17.0	
	Sumatriptan 100mg	NR			NR			NR			
Tfelt-Hansen 1995 ⁴⁹	Placebo										

Trial	Arms	Sustained	Pain Freedom,	24-hours	Sustaine	d Pain Fre hours	edom, 48-	hours			
		n	N	%	n	Ν	%	n	N	%	
Myllyla 1998 ⁴³	Sumatriptan 100mg	NR			NR			NR			
	Placebo										
Tfelt-Hansen 1998 ⁵⁰	Sumatriptan 100mg	NR			NR			NR			
	Placebo										
Dowson 2002 ³⁵	Sumatriptan 100mg	NR			NR			NR			
Dowson 2002	Placebo										
Kudrow 2005 ⁴⁰	Sumatriptan 50mg	NR			NR			NR			
	Placebo										
Lines 2001 ⁴¹	Sumatriptan 50mg	NR			NR			NR			
Lines 2001	Placebo										
Nappi 1994 ⁴⁴	Sumatriptan 100mg	NR			NR			NR			
	Placebo										
	Sumatriptan 100mg	NR			NR			NR			
Pfaffenrath 1998 ⁴⁵	Sumatriptan 50mg				NR			NR			
	Placebo										
	Sumatriptan 100mg	NR									
Oral Sumatriptan International Multiple-					NR			NR			
Dose Study Group 1991 ⁵²	Placebo										
	Eletriptan 40mg	NR			NR			342	795	43	
Mathew 2003 ⁴²	Sumatriptan 100mg							276	812	34	
	Placebo							58	414	14	
	Eletriptan 40mg	NR			NR			NR			
Goadsby 2000 ³⁸	Sumatriptan 100mg										
	Placebo										

mg: milligram, n: number of participants, N: total number of participants, NR: not reported.

*Data are digitized and should be interpreted with caution

Table D7. Adverse Events

Trial	A 1100 a		Any A	E		TEAEs		Nausea			Dizziness			Somnolence		
ITIdi	Arms	n	Ν	%	n	Ν	%	n	N	%	n	Ν	%	n	Ν	%
				La	ismidita	an	-									
	Lasmiditan 200mg	260	609	42.7	237	609	38.9	32	609	5.3	99	609	16.3	33	609	5.4
SAMURAI ²³	Lasmiditan 100mg	229	630	36.3	205	630	32.5	19	630	3.0	79	630	12.5	36	630	5.7
	Placebo	101	617	16.4	78	617	12.6	12	617	1.9	21	617	3.4	14	617	2.3
	Lasmiditan 200mg	253	649	39.0				17	649	2.6	117	649	18.0	42	649	6.5
SPARTAN ²⁴	Lasmiditan 100mg	230	635	36.2	NR			21	635	3.3	115	635	18.1	29	635	4.6
	Lasmiditan 50mg	167	654	25.5				18	654	2.8	56	654	8.6	35	654	5.4
	Placebo	75	645	11.6				8	645	1.2	16	645	2.5	13	645	2
Farkkila 2012 ²⁵	Lasmiditan 200mg				61	71	85.9	2	71	2.8	27	71	38.0	8	71	11.3
	Lasmiditan 100mg	NR			59	82	72.0	8	82	9.8	21	82	25.6	10	82	12.2
	Lasmiditan 50mg				53	82	64.6	4	82	4.9	19	82	23.2	8	82	9.8
	Placebo				19	86	22.1	0	86	0	0	86	0	2	86	2.3
				Rii	megepa	int										
Study 301 ²⁷	Rimegepant 75mg	69	546	12.6	3	546	0.5	5	546	0.9	4	546	0.7	NR		
	Placebo	59	549	10.7	1	549	0.2	6	549	1.1	2	549	0.4	INIX		
Study 302 ²⁶	Rimegepant 75mg	93	537	17.3	NR			10	537	1.8	NR			NR		
51449 502	Placebo	77	535	14.4				6	535	1.1						
Study 303 ²⁸	Rimegepant 75mg	90	682	13.5	47	682	6.9	11	682	1.6	6	682	0.9	NR		
	Placebo	73	693	10.7	36	693	5.2	3	693	0.4	7	693	1.0			
	Rimegepant 75mg							3	86	3.5	1	86	1.2			
Marcus 2014 ²⁹	Sumatriptan 100mg	NR			NR			2	100	2.0	1	100	1.0	NR		
							5	209	2.4	2	209	1.0				
				Uk	orogepa	int										
ACHIEVE I ³¹	Ubrogepant 100mg	79	485	16.3	58	485	12.0	16	485	3.3	7	485	1.4	11	485	2.3

Trial	Arms		Any A	E		TEAEs			Nause	а	Dizziness			Somnolence			
I lai	AIIIIS	n	Ν	%	n	Ν	%	n	Ν	%	n	Ν	%	n	Ν	%	
	Ubrogepant 50mg	44	466	9.4	27	466	5.8	7	466	1.5	4	466	0.9	3	466	0.6	
	Placebo	62	485	12.8	41	485	8.5	8	485	1.6	3	485	0.6	4	485	0.8	
	Ubrogepant 50mg	63	488	12.9	42	488	8.6	10	488	2.0	7	488	1.4	4	488	0.8	
ACHIEVE II ³⁰	Ubrogepant 25mg	44	478	9.2	30	478	6.3	12	478	2.5	10	478	2.1	4	478	0.8	
	Placebo	51	499	10.2	30	499	6.0	10	499	2.0	8	499	1.6	2	499	0.4	
	Ubrogepant 100mg	30	102	29.4	25	102	24.5	7	102	6.9	6	102	5.9	4	102	3.9	
Voss 2016 ³²	Ubrogepant 50mg	23	107	21.5	18	107	16.8	8	107	7.5	2	107	1.9	3	107	2.8	
V055 2016	Ubrogepant 25mg	21	103	20.4	14	103	13.6	6	103	5.8	3	103	2.9	5	103	4.9	
	Placebo	28	113	24.8	23	113	20.4	4	113	3.5	1	113	0.9	6	113	5.3	
					Triptan	s											
Diener 2002 ³⁴	Eletriptan 40mg	NR			NR			10	210	4.8	10	210	4.8	5	210	2.4	
	Placebo				INIX			7	106	6.6	2	106	3.8	2	106	1.9	
Steiner 2003 ⁴⁸	Eletriptan 40mg	117	392	30	NR	6 3 NR						392	1.5	9	392	2.3	
Steller 2005	Placebo	57	144	40	INIX						2	144	1.4	0	0	0	
Garcia-Ramos 2003 ³⁶	Eletriptan 40mg	60	192	31	50	192	26	17	192	8.9	12	192	6.3	10	192	5.2	
	Placebo	32	92	35	15	92	16	13	92	14.1	3	92	3.3	2	92	2.2	
The EMSASI Study Group 2004 ⁵¹	Sumatriptan 50mg	44	224	19.8	15	224	6.6	NR			NR			NR	ND		
	Placebo	32	222	14.4	10	222	4.5										
Diener 2004 ³³	Sumatriptan 50mg	19	135	14.1	9	135	6.7	NR			NR			NR			
	Placebo	16	153	10.5	6	153	3.9										
Geraud 2000 ³⁷	Sumatriptan 100mg	279	492	56.7	NR			35	492	7.1	46	492	9.3	29	492	5.9	
	Placebo	13	56	23.2				1	56	1.8	1	56	1.8	2	56	3.6	
	Sumatriptan 100mg				57	488	11.7	13	488	2.7							
Sheftell 2005 ⁴⁶ - Study 1	Sumatriptan 50mg	NR			40	494	8.1	11	494	2.2	NR			NR			
	Placebo				17	495	3.4	5	495	1							
Sheftell 2005 ⁴⁶ - Study 2	Sumatriptan 100mg	NR			94	485	19.4	16	485	3.3	NR			NR			
	Sumatriptan 50mg				58	496	11.7	10	496	2							

Trial	Arms		Any A	E		TEAEs			Nause	a	Dizziness			Somnolence				
i fidi	Arms	n	Ν	%	n	Ν	%	n	Ν	%	n	Ν	%	n	Ν	%		
	Placebo				25	494	5.1	5	494	1								
Havanka 2000 ³⁹	Sumatriptan 100mg	25	98	26	NR			1	98	1.0	NR			NR				
	Placebo	21	91	23	INIT			1	91	1.1	INIT			INIT				
Smith 2005 ⁴⁷	Sumatriptan 50mg	55	229	24	NR			3	229	1.3	11	229	4.8	6	229	2.6		
51111 2005	Placebo	36	242	15	INIX			4	242	1.7	8	242	3.3	0	0	0		
Tfelt-Hansen 1995 ⁴⁹	Sumatriptan 100mg	35	125	28	NR			14	125	11.2	3	125	2.4	6	125	4.8		
Treit-Hallsell 1995	Placebo	16	126	13	INIX			11	126	8.7	1	126	0.8	0	0	0		
Myllyla 1998 ⁴³	Sumatriptan 100mg	17	46	38	NR			8	46	17.4	NR			NR	NR			
	Placebo	9	48	19				2	48	4.2								
Tfelt-Hansen 1998⁵⁰	Sumatriptan 100mg	202	388	52.1	160	388	41.2	35	388	9	35	388	9	28	388	7.2		
	Placebo	51	160	31.9	32	160	20	4	160	2.5	6	160	3.8	9	160	5.6		
Dowson 2002 ³⁵	Sumatriptan 100mg	43	194	22.2	NR			NR			4	194	2.1	4	194	2.1		
	Placebo	6	99	6.1	i Ni V							99	2	0	0	0		
Kudrow 2005 ⁴⁰	Sumatriptan 50mg	45	141	31.9	30	141	21.3	6	141	4.3	3	141	2.1	3	141	2.1		
	Placebo	41	140	29.3	24	140	17.1	2	140	1.4	4	140	2.9	3	140	2.1		
Lines 2001 ⁴¹	Sumatriptan 50mg	NR			NR			NR			NR			NR				
	Placebo					INK		INT			i i i i							
Nappi 1994 ⁴⁴	Sumatriptan 100mg	47	162	29	NR			12	162	7.4	NR			NR				
	Placebo	14	88	15.9				6	88	6.8	i i i i			, , , , , , , , , , , , , , , , , , ,				
	Sumatriptan 100mg	111	298	37.2				13	298	4.4	14	298	4.7					
Pfaffenrath 1998 ⁴⁵	Sumatriptan 50mg	82	303	27.1	NR			18	303	5.9	4	303	1.3	NR				
	Placebo	20	99	20.2				2	99	2	2	99	2					
	Sumatriptan 100mg	57	149	38				12	149	8	7 149 5		5					
Oral Sumatriptan International					NR									NR				
Multiple-Dose Study Group 1991 ⁵²	Placebo	19	84	23				5	84	6	2 84 2		2					
											2 04 2							
Mathew 2003 ⁴²	Eletriptan 40mg	259	835	31	NR			99	835	11.9	NR			NR				
	Sumatriptan 100mg	314	849	37				125	849	14.7				INK				

Trial	Arms		Any A	E	TEAEs			Nausea			Dizziness			Somnolence			
ITIdi	ATTIS	n	Ν	%	n	Ν	%	n	Ν	%	n	Ν	%	n	N	%	
	Placebo	146	429	34				54	429	12.6							
Goadsby 2000 ³⁸	Eletriptan 40mg	47	136	34.6				2	136	1.5	5	136	3.7				
	Eletriptan 20mg	49	144	34	NR			4	144	2.8	3	144	2.1	NR			
	Sumatriptan 100mg	52	129	40.3				4	129	3.1	5	129	3.9	INIT			
	Placebo	24	142	16.9				1	142	0.7	1	142	0.7				
Kolodny 2004	Sumatriptan 50mg	142	287	49.5	110	287	38.3	19	287	6.6	30	287	10.5	18	287	6.3	
	Placebo	102	288	35.4	61	288	21.2	12	288	4.2	13	288	4.5	13	288	4.5	
Pini 1995	Sumatriptan 100mg	NR			18	151	12	NR			NR			NR			
	Placebo	INK			6	87	7	INIX			INK			INK			

AE: adverse event, mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, TEAE: treatment-emergent adverse event.

Additional Efficacy Outcomes from the Trials of Lasmiditan, Rimegepant, and Ubrogepant (Not Included in the NMA)

			Phonop	hobia-Free	Photop	hobia-Free	Naus	ea-Free	Vomiting-Free		
Trial	Arms	N	n (%) OR (95%Cl), p-value		n (%)	OR (95%CI), p-value	n (%)	OR (95%CI), p-value	n (%)	OR (95%CI), p-value	
				La	smiditan						
	Lasmiditan 200mg	555	419 (75.5)	1.5 (1.1, 1.9), 0.005	379 (68.3)	2.0 (1.5, 2.6), <0.001	449 (80.9)	1.2 (0.9, 1.7), 0.153	546 (98.4)	0.9 (0.3, 2.3), 0.773	
SAMURAI ²⁴	Lasmiditan 100mg	562	426 (75.8)	1.6 (1.2, 2.0), 0.002	388 (69.0)	2.1 (1.7, 2.8), <0.001	448 (79.7)	1.2 (0.9, 1.6), 0.276	549 (97.7)	0.6 (0.3, 1.5), 0.286	
	Placebo	554	374 (67.5)		294 (53.1)		427 (77.1)		546 (98.6)		
	Lasmiditan 200mg	565	431 (76.3)	1.8 (1.4, 2.4), <0.001	391 (69.2)	2.0 (1.5, 2.6), <0.001	460 (81.4)	1.0 (0.8, 1.4), 0.834	557 (98.6)	0.6 (0.2, 1.8), 0.373	
SPARTAN ²³	Lasmiditan 100mg	571	428 (75.0)	1.7 (1.3, 2.2), <0.001	380 (66.5)	1.8 (1.4, 2.3), <0.001	468 (82.0)	1.1 (0.8, 1.5), 0.629	567 (99.3)	1.2 (0.3, 4.6), 0.749	
	Lasmiditan 50mg	598	428 (71.6)	1.4 (1.1, 1.9), 0.004	368 (61.5)	1.4 (1.1, 1.8), 0.005	473 (79.1)	0.9 (0.7, 1.2), 0.443	588 (98.3)	0.5 (0.2, 1.5), 0.229	
	Placebo	576	368 (63.9)		309 (53.6)		465 (80.7)		571 (99.1)		
	Lasmiditan 200mg	N not reported	60.5*	NR, n.s.	48.5*	NR, 0.031	64.4*	NR, n.s.	92.5*	NR, n.s.	
Farkkila 2012 ²⁵	Lasmiditan 100mg	N not reported	76.9*	NR, 0.0013	69.3*	NR, <0.0001	75.6*	NR, 0.034	99.9*	NR, 0.0027	
	Lasmiditan 50mg	N not reported	58.1*	NR, n.s.	53.4*	NR, 0.018	68.5*	NR, n.s.	94.6*	NR, n.s.	
	Placebo	N not reported	52.1*		34.9*		59.4*		88.9*		
				Rir	negepant						
Study 301 ²⁷	Rimegepant 75mg	See results column	133/345 (38.6)	1.4 (1.0, 1.9), 0.03†	164/470 (34.9)	1.6 (1.2, 2.1), <0.001†	149/318 (46.9)	1.2 (0.9, 1.7), n.s.†	NR		

Table D8. Efficacy Outcomes at 2 Hours: Associated Migraine Symptoms

			Phonop	hobia-Free	Photop	hobia-Free	Naus	ea-Free	Vomiting-Free		
Trial	Arms	N	n (%)	n (%) OR (95%Cl), p-value		OR (95%CI), p-value	n (%)	OR (95%CI), p-value	n (%)	OR (95%CI), p-value	
	Placebo	See results column	113/366 (30.9)		120/483 (24.8)		134/322 (41.6)				
Study 302 ²⁶	Rimegepant 75mg	See results column	133/362 (36.7)	1.6 (1.2, 2.2),	183/489 (37.4)	2.1 (1.6, 2.8),	171/355 (48.1)	1.2 (0.9, 1.7),	NR		
Study 302	Placebo	See results column	100/374 (26.8)	0.004†	106/477 (22.3)	<0.0001†	145/336 (43.3)	n.s.†			
Study 303 ²⁸	Rimegepant 75mg	See results column	188/451 (41.7)	1.7 (1.3, 2.2),	198/593 (33.4)	1.5 (1.2, 2.0),	203/397 (51.0)	1.3 (1.0, 1.7),	ND		
Study 505	Placebo	See results column	135/447 (30.2)	<0.001†	150/611 (24.5)	<0.001†	194/430 (45.2)	n.s.†	NR		
	Rimegepant 75mg	86	45 (52.3)	2.8 (1.7, 4.8), <0.0001 ⁺	36 (41.9)	2.3 (1.3, 3.9), 0.003†	58 (67.4)	2.0 (1.2, 3.4), 0.01†			
Marcus 2014 ²⁹	Sumatriptan 100mg	100	49 (49.0)	2.5 (1.5, 4.1), <0.001†	47 (47.0)	2.8 (1.7, 4.7), <0.0001†	60 (60.0)	1.4 (0.9, 2.4), n.s.†	NR		
	Placebo	204	57 (28.1)		49 (24.1)		104 (51.2)				
Ubrogepant							1				
	Ubrogepant 100mg	448	244 (49.0)	1.5 (1.1, 2.0), n.s.	205 (45.8)	1.8 (1.4, 2.4), 0.004	310 (69.2)	1.4 (1.0, 1.8), n.s.	NR		
ACHIEVE I ³¹	Ubrogepant 50mg	423	245 (57.9)	1.6 (1.2, 2.1), n.s.	172 (40.7)	1.6 (1.2, 2.2), n.s.	297 (70.2)	1.3 (1.0, 1.8), n.s.			
	Placebo	456	215 (47.1)		143 (31.4)		284 (62.3)				
	Ubrogepant 50mg	464	251 (54.1)	1.4 (1.1, 1.8), 0.044	203 (43.8)	1.5 (1.1, 2.0), 0.0167	331 (71.3)	1.1 (0.8, 1.5), n.s.	NR		
ACHIEVE II ³⁰	Ubrogepant 25mg	435	234 (53.6)	1.4 (1.0, 1.8), n.s.	171 (39.3)	1.3 (1.0, 1.7), n.s.	307 (70.6)	1.1 (0.8, 1.5), n.s.			
	Placebo	456	212 (46.3)		162 (35.5)		319 (70.0)				

Trial			Phonop	hobia-Free	Photop	hobia-Free	Naus	ea-Free	Vomi	ting-Free
	Arms	N	n (%)	OR (95%CI), p-value	n (%)	OR (95%CI), p-value	n (%)	OR (95%CI), p-value	Vomiting-Free n (%) OR (95%CI), p-value NR	
	Ubrogepant 100mg	102	62 (60.8)	2.1 (1.2, 3.7), 0.006†	56/102 (54.9)	2.8 (1.6, 4.9), <0.001†	72/102 (70.6)	1.4 (0.8, 2.6), n.s.†	NR	
Voss 2016 ³²	Ubrogepant 50mg	105	59 (56.2)	1.8 (1.0, 3.0), 0.04†	50 (47.6)	2.1 (1.2, 3.6), 0.0†	72 (68.6)	1.3 (0.7, 2.3), n.s.†		
	Ubrogepant 25mg	103	57 (55.3)	1.7 (1.0, 2.9), n.s.†	41 (39.8)	1.5 (0.9, 2.7), n.s.†	76 (73.8)	1.7 (0.9, 3.0), n.s.†		
	Placebo	112	47 (42.0)		34 (30.4)		70 (62.5)			

95%CI: 95% Confidence Interval, mg: milligram, n: number of participants, N: total number of participants, NR: not reported, n.s.: not significant, OR: odds ratio

* data are digitized and should be interpreted with caution

+ calculated

Table D9. Patient-Reported Outcomes at 2 Hours

	Global Impression of Change at 2 Hours, n (%)											
Trial	Arms	N	Very Much Better	Much Better	A Little Better	No Change	A Little Worse	Much Worse	Very Much Worse	p-Value vs. Placebo		
				Lasn	niditan							
	Lasmiditan 200mg	555	57 (10.3)	153 (27.6)	143 (25.8)	60 (10.8)	31 (5.6)	13 (2.3)	5 (0.9)	<0.001		
SAMURAI ²⁴	Lasmiditan 100mg	562	54 (9.6)	155 (27.6)	153 (27.2)	83 (14.8)	16 (2.8)	8 (1.4)	8 (1.4)	<0.001		
	Placebo	554	34 (6.1)	87 (15.7)	159 (28.7)	146 (26.4)	28 (5.1)	14 (2.5)	3 (0.5)			
	Lasmiditan 200mg	565	82 (14.5)	158 (28.0)	155 (27.4)	70 (12.4)	20 (3.5)	13 (2.3)	5 (0.9)	<0.001		
SPARTAN ²³	Lasmiditan 100mg	571	74 (13.0)	161 (28.2)	163 (28.5)	75 (13.1)	27 (4.7)	10 (1.8)	3 (0.5)	<0.001		
SPARTAN	Lasmiditan 50mg	598	66 (11.0)	153 (25.6)	175 (29.3)	98 (16.4)	29 (4.8)	11 (1.8)	4 (0.7)	<0.001		
	Placebo	576	46 (8.0)	115 (20.0)	162 (28.1)	152 (26.4)	25 (4.3)	15 (2.6)	1 (0.2)			
	Lasmiditan 200mg	69	19 (28.0)							n.s.		
Farkkila 2012 ²⁵	Lasmiditan 100mg	81	29 (36.0)		NR				0.0041			
	Lasmiditan 50mg	79	18 (23.0)							n.s.		
	Placebo	81	13 (16.0)	13 (16.0)								
				Rime	gepant							
Study 301 ²⁷	Rimegepant 75mg	543	NR									
5100 501	Placebo	541										
Study 302 ²⁶	Rimegepant 75mg	537	NR									
5100 302	Placebo	535										
Study 303 ²⁸	Rimegepant 75mg	669	NR									
Study 303	Placebo	682										
	Rimegepant 75mg	91										
Marcus 2014 ²⁹	Sumatriptan 100mg	109	NR									
	Placebo	229										
				Ubro	gepant							
ACHIEVE I ³¹	Ubrogepant 100mg	297	102 (34.3)							<0.001		
	Ubrogepant 50mg	299	103 (34.4)		NR					<0.001		
	Placebo	313	69 (22.0)									

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			Global Impression of Change at 2 Hours, n (%)									
Trial	Arms	N	Very Much Better	Much Better	A Little Better	No Change	A Little Worse	Much Worse	Very Much Worse	p-Value vs. Placebo		
	Ubrogepant 50mg	392	131 (33.4)							<0.001		
ACHIEVE II ³⁰	Ubrogepant 25mg	435	148 (34.1)		NR					<0.001		
	Placebo	376	78 (20.7)									
	Ubrogepant 100mg	102										
Voss 2016 ³²	Ubrogepant 50mg	106	ND									
V055 2016-	Ubrogepant 25mg	104	NR									
f	Placebo	113										

mg: milligram, n: number of participants, N: total number of participants, NR: not reported, n.s.: not significant.

Table D10. Adverse Events

Trial	Arms	N	AE Leading to D/C, n (%)	SAEs, n (%)	Death, n (%)	Any AEs, n (%)	TEAEs, n (%)	Dizziness, n (%)	Somnolence, n (%)	Paresthesia, n (%)	Serum AST or ALT Above ULN, n (%)
					La	smiditan					
	Lasmiditan 200mg	609	0 (0)	2 (0.3)	0 (0)	260 (42.7)	237 (38.9)	99 (16.3)	33 (5.4)	48 (7.9)	
SAMURAI ²⁴	Lasmiditan 100mg	630	0 (0)	0 (0)	0 (0)	229 (36.3)	205 (32.5)	79 (12.5)	36 (5.7)	36 (5.7)	NR
	Placebo	617	0 (0)	1 (0.2)	0 (0)	101 (16.4)	78 (12.6)	21 (3.4)	14 (2.3)	13 (2.1)	
	Lasmiditan 200mg	649	1 (0.2)	1 (0.2)	0 (0)	253 (39.0)		117 (18.0)	42 (6.5)	43 (6.6)	
SPARTAN ²³	Lasmiditan 100mg	635	0 (0)	1 (0.2)	0 (0)	230 (36.2)	676 (93.2)	115 (18.1)	29 (4.6)	37 (5.8)	NR
	Lasmiditan 50mg	654	0 (0)	0 (0)	0 (0)	167 (25.5)		56 (8.6)	35 (5.4)	16 (2.4)	
	Placebo	645	0 (0)	0 (0)	0 (0)	75 (11.6)		16 (2.5)	13 (2.0)	6 (0.9)	
	Lasmiditan 200mg	71	NR	28 (39.0)	0 (0)	NR	61 (86.0)	27 (38.0)	8 (11.3)	12 (17.0)	NR
Farkkila 2012 ²⁵	Lasmiditan 100mg	82	NR	23 (28.0)	0 (0)	NR	59 (72.0)	21 (26.0)	10 (12.2)	9 (11.0)	
	Lasmiditan 50mg	82	NR	16 (20.0)	0 (0)	NR	53 (65.0)	19 (23.0)	8 (9.8)	2 (2.0)	
	Placebo	86	NR	5 (6.0)	0 (0)	NR	19 (22.0)	0 (0)	2 (2.3)	2 (2.3)	
					Rin	negepant					
Study 301 ²⁷	Rimegepant 75mg	546	0 (0)	2 (0.4)	NR	69 (12.6)	3 (0.5)	4 (0.7)	NR	NR	11 (2.0)
	Placebo	549	0 (0)	1 (0.2)	NR	59 (10.7)	1 (0.2)	2 (0.4)			20 (3.6)
Study 302 ²⁶	Rimegepant 75mg	537	0 (0)	1 (0.2)	0 (0)	93 (17.3)	NR	NR	NR	NR	13 (2.4)
	Placebo	535	0 (0)	2 (0.4)	0 (0)	77 (14.4)					12 (2.2)
Study 303 ²⁸	Rimegepant 75mg	682	0 (0)	0 (0)	0 (0)	90 (13.5)	47 (6.9)	6 (0.9)	NR	NR	1 (0.1)

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Trial	Arms	N	AE Leading to D/C, n (%)	SAEs, n (%)	Death, n (%)	Any AEs, n (%)	TEAEs, n (%)	Dizziness, n (%)	Somnolence, n (%)	Paresthesia, n (%)	Serum AST or ALT Above ULN, n (%)
	Placebo	693	0 (0)	0 (0)	0 (0)	73 (10.5)	36 (5.2)	7 (1.0)			1 (0.1)
	Rimegepant 75mg	86	0 (0)	0 (0)	0 (0)			1 (1.2)		0 (0)	NR
Marcus 2014 ²⁹	Sumatriptan 100mg	100	0 (0)	0 (0)	0 (0)	NR	NR	1 (1.0)	NR	2 (2.0)	
	Placebo	209	0 (0)	0 (0)	0 (0)			2 (1.0)		2 (1.0)	
	Ubrogepant										
	Ubrogepant 100mg	485	1 (0.2)	2 (0.4)	0 (0)	79 (16.3)	58 (12.0)	7 (1.4)	12 (2.5)	NR	62 (12.9)
ACHIEVE I ³¹	Ubrogepant 50mg	466	1 (0.2)	3 (0.6)	0 (0)	44 (9.4)	27 (5.8)	4 (0.9)	3 (0.6)		NR
	Placebo	485	3 (0.6)	0 (0)	0 (0)	62 (12.8)	41 (8.5)	3 (0.6)	4 (0.8)		NR
	Ubrogepant 50mg	488	2 (0.4)	0 (0)	0 (0)	63 (12.9)	42 (8.6)	7 (1.4)	4 (0.8)		NR
ACHIEVE II ³⁰	Ubrogepant 25mg	478	1 (0.2)	0 (0)	0 (0)	44 (9.2)	30 (6.3)	10 (2.1)	4 (0.8)	NR	53 (11.2)
	Placebo	499	1 (0.2)	0 (0)	0 (0)	51 (10.2)	30 (6.0)	8 (1.6)	2 (0.4)		NR
	Ubrogepant 100mg	102	0 (0)	0 (0)	0 (0)	30 (29.4)	25 (24.5)	6 (5.9)	4 (3.9)		0 (0)
Voss 2016 ³²	Ubrogepant 50mg	107	0 (0)	2 (1.9)	0 (0)	23 (21.5)	18 (16.8)	2 (1.9)	3 (2.8)	NR	1 (0.9)
1	Ubrogepant 25mg	103	0 (0)	0 (0)	0 (0)	21 (20.4)	14 (13.6)	3 (2.9)	5 (4.9)		0 (0)
	Placebo	113	0 (0)	0 (0)	0 (0)	28 (24.8)	23 (20.4)	1 (0.9)	6 (5.3)		0 (0)

AE: adverse event, ALT: alanine aminotransferase, AST: aspartate aminotransferase, D/C: discontinuation, mg: milligram, n: number of participants, N: total number of participants, NR: not reported, TEAE: treatment-emergent adverse event, ULN: upper limit of normal.

Trial	Arms	N , (Treated		he Pain Inten Iigraine Attac			Baseli	ne Sympto	oms of Trea	ted Attacks,	n (%)	MBS of	f Treated A n (%)	ttacks,
11101		Attacks)	Severe	Moderate	Mild	None	Phono- phobia	Photo- phobia	Nausea	Vomiting	None	Phono- phobia	Photo- phobia	Nause a
Lasmiditan														
GLADIATOR ⁵⁶	Lasmiditan 200mg	1015 (8513)	2848 (33.4)	5546 (65.1)	115 (1.4)	6 (0.1)	4988 (58.6)	6322 (74.3)	3188 (37.4)	302 (3.5)	962 (11.3)	1726 (22.9)	4141/ 7550 (54.9)	1683/ 7550 (22.3)
GLADIATOK	Lasmiditan 100mg	963 (8782)	2872 (32.7)	5762 (65.6)	141 (1.6)	7 (0.7)	5609 (63.9)	6741 (76.8)	3527 (40.2)	275 (3.1)	792 (9.0)	1970/ 7987 (24.7)	4307/ 7987 (53.9)	1710/ 7987 (21.4)
						Rimege	pant							
Study 201 ⁸⁸	NR													
	_					Ubroge	pant							
NCT 02873221 ^{89,93,94}	NR													

Table D11. Open-Label Extension Studies for Lasmiditan, Rimegepant, and Ubrogepant – Baseline Characteristics

mg: milligram, n: number of participants, N: total number of participants, NR: not reported.

· • • • • • • • • • • • • • • • • • • •	crextension studies for Easimatian, Rintegepant, and Obrogepa					Enlacy Outcomes		
		Headache Pain at 2 Hoi		Free of MBS a	at 2 Hours	Number of Attacks	Reduction in Mean	
Trial	Arms	n/N (%)	p-value	n/N (%)	p-value	Treated with Second Dose, n/N (%)	Migraine Days per Month, Mean	
			Ubrogepa	nt				
	Lasmiditan 200mg	2668/8232 (32.4)	<0.001	2963/7298 (40.6)	<0.001	2776/8513 (32.6)	NR	
GLADIATOR ⁵⁶	Lasmiditan 100mg	2296/8532 (26.9)		2909/7758 (37.5)		3627/8782 (41.3)	NR	
			Rimegepa	nt				
Study 201* ⁸⁸	Rimegepant 75mg PRN (2-8) Rimegepant 75mg PRN (9- 14)	NR		NR		NR	NR	
51009 201	Rimegepant 75mg QOD + PRN	NR		NR		NR	-6.0 (at 52 weeks)†	
	Rimegepant 75mg Total	NR		NR		NR		
			Ubrogepa	nt				
NCT 02873221 ^{89,93,94}	Ubrogepant 100mg Ubrogepant	105/420 (25.0) 96/417 (23.0)	NR NR	NR		NR	NR	
	50mg Usual care							

Table D12. Open-Label Extension Studies for Lasmiditan, Rimegepant, and Ubrogepant – Efficacy Outcomes

mg: milligram, n: number of participants, N: total number of participants, NR: not reported, PRN: as needed, QOD: every other day.

*based on interim analysis at three months,

+ in patients with ≥14 headache days/month.

			_	-						
Trial	Arms	N	Any AE, n (%)	TEAE, n (%)	SAEs, n (%)	Treatment-Emergent	AE Leading to	Death, n (%)		
						SAEs, n (%)	D/C, n (%)			
	-			Lasmidit	an					
GLADIATOR	Lasmiditan 200mg	1015	731 (72.0)	528 (52.0)	32 (3.2)	3 (0.3)	146 (14.4)	0 (0)		
GLADIATON	Lasmiditan 100mg	963	636 (66.0)	434 (45.1)	28 (2.9)	6 (0.6)	108 (11.2)	0 (0)		
Rimegepant										
	Rimegepant 75mg PRN (2-8)	1017	659 (64.8)	NR	NR	NR	24 (2.4)	0 (0)		
Study 201 ^{*88}	Rimegepant 75mg PRN (9-14)	481	294 (61.1)				15 (3.1)	0 (0)		
Study 201	Rimegepant 75mg QOD + PRN	109	109 (38.1)				9 (3.1)	0 (0)		
	Rimegepant 75mg Total	1784	1062 (59.5)		45 (2.5)	9 (0.5)	48 (2.7)	0 (0)		
				Ubrogep	ant					
NCT 02873221 ^{89,93,94}	Ubrogepant 100mg	409	297 (72.6)	43 (10.5)	12 (2.9)	NR	11 (2.7)	0 (0)		
NCT 02873221-0000	Ubrogepant 50mg	417	268 (66.3)	42 (10.4)	9 (2.2)		9 (2.2)	0 (0)		
	usual care	417	271 (65.0)	65 (15.6)	17 (4.1)		NR	0 (0)		

Table D13. Open Label Extension Studies for Lasmiditan, Rimegepant, and Ubrogepant – Adverse Events I

AE: adverse event, D/C: discontinuation, n: number of participants, N: total number of participants, PRN: as needed, QOD: every other day, SAE: serious adverse events, TEAE: treatment-emergent adverse event.

*based on interim analysis at three months

Trial	Arms	N	Dizziness, n (%)	Somnolence, n (%)	Paresthesia, n (%)	Fatigue, n (%)	Nausea, n (%)	Upper Respiratory Tract Infection, n (%)		
				Lasmidit	tan					
GLADIATOR ⁵⁶	Lasmiditan 200mg	1015	217 (21.3)	95 (9.3)	85 (8.3)	63 (6.2)	53 (5.2)	NR		
GLADIATOR	Lasmiditan 100mg	963	153 (15.8)	76 (7.8)	51 (5.3)	46 (4.7)	41 (4.2)	INU		
	Rimegepant									
	Rimegepant 75mg PRN (2-8)	1017	25 (2.5)				33 (3.2)	108 (10.6)		
Study 201* ⁸⁸	Rimegepant 75mg PRN (9-14)	481	11 (2.3)	NR	NR	NR	15 (3.1)	31 (6.4)		
5000 201	Rimegepant 75mg QOD + PRN	109	3 (1.0)	NIX .	NIX.		3 (1.0)	12 (4.2)		
	Rimegepant 75mg Total	1784	39 (2.2)				51 (2.9)	151 (8.5)		
	Ubrogepant									
	Ubrogepant 100mg	409	12 (2.9)	NR	NR	NR	19 (4.6)	44 (10.8)		
NCT 02873221 ^{89,93,94}	Ubrogepant 50mg	417	5 (1.2)	NR	NR	NR	19 (4.7)	47 (11.6)		
	usual care	417	4 (1.0)	NR	NR	NR		48 (11.5)		

Table D14. Open Label Extension Studies for Lasmiditan, Rimegepant, and Ubrogepant – Adverse Events II

mg: milligrams, n: number of participants, N: total number of participants, NR: not reported, PRN: as needed, QOD: every other day.

* based on interim analysis at three months

Supplemental NMA Methods

As described in the report, we conducted random effect network meta-analyses (NMAs) where feasible. An NMA extends pairwise meta-analyses by simultaneously combining both the direct estimates (i.e., estimates obtained from head-to-head comparisons) and indirect estimates (i.e., estimates obtained from comparator[s]).^{128,129}

NMAs were conducted using a Bayesian framework. For continuous outcomes, the NMA model corresponds to a generalized linear model with identity link.⁸⁷ For binary outcomes (e.g., proportion of patients pain-free at 2 hours), the NMA model corresponds to a generalized linear model with a logit link.⁸⁷ For all analyses, we included random effects on the treatment parameters, and the amount of between-study variance (i.e., heterogeneity) was assumed constant across all treatment comparisons. We used noninformative prior distributions for all model parameters. We initially discarded the first 50,000 iterations as "burn-in" and base inferences on an additional 50,000 iterations using three chains. Convergence of chains was assessed visually using trace plots.

Furthermore, for any network where there were "loops" in evidence, we empirically compared the direct and indirect estimates to assess if the NMA consistency assumption is violated using a node-splitting approach.¹³⁰ As there was no evidence of inconsistency, we present the full NMA results in the report. All analyses were conducted in R using the gemtc package.

Supplemental NMA Results

We provide three network diagrams that represents the NMAs in the report (Figure D2, D3 and D4). To interpret the network figures, note that the lines indicate the presence of a trial directly assessing the connecting interventions, with the thickness of the line corresponding to the number of trials. The location of treatments and the distances between them does not have any meaning. The gepants are depicted in blue, lasmiditan in green, triptans in orange, and placebo in black.

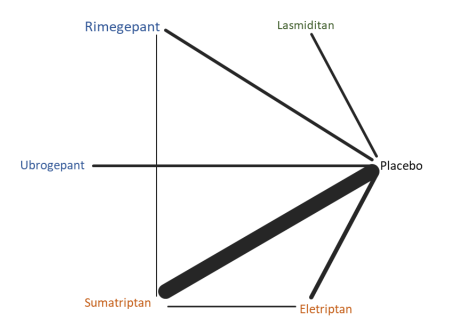


Figure D2. Network of Studies Included in the NMA of 2-hours Pain Relief (see Legend)

Legend: Figure D2 is a network of studies included in the NMA of 2-hours Pain Relief, with the thickness of the connecting lines related to the number of trials available for each pair of treatments. The NMAs of 2-hours Pain Freedom, 24 hours Sustained Pain Freedom, Any AE, TEAE, and dizziness all have a similar network diagram (not shown), with less studies contributing to the sumatriptan versus placebo connection.



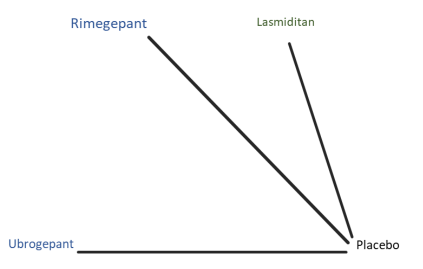
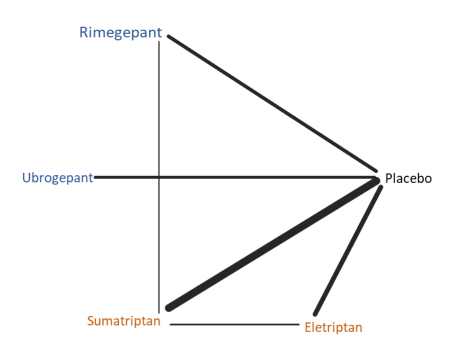


Figure D4. Network of Studies Included in the NMAs of Sustained Pain Freedom at 24 hours



Additional league tables that were not provided in the report are presented below. As stated in the report, each box represents the estimated odds ratio and 95% credible interval for the combined direct and indirect comparisons between two drugs. Estimates in bold signify that the 95% credible interval does not contain 1.

Lasmiditan		_		
1.14 (0.69, 1.84)	Rimegepant		_	
1.12 (0.65, 1.9)	0.99 (0.6, 1.61)	Ubrogepant		
0.56 (0.37, 0.88)	0.5 (0.35, 0.73)	0.5 (0.33, 0.8)	Sumatriptan	
0.37 (0.23, 0.63)	0.33 (0.21, 0.53)	0.33 (0.21, 0.57)	0.66 (0.47, 0.93)	Eletriptan
2.21 (1.53, 3.25)	1.95 (1.45, 2.69)	1.97 (1.37, 2.95)	3.91 (3.19, 4.76)	5.89 (4.23, 8.14)

Lasmiditan		_			
1.19 (0.86, 1.70)	Rimegepant		_		
1.28 (0.91, 1.91)	1.08 (0.77, 1.54)	Ubrogepant		_	
0.72 (0.55, 1.00)	0.6 (0.48, 0.79)	0.56 (0.42, 0.75)	Sumatriptan		
0.46 (0.33, 0.67)	0.39 (0.29, 0.54)	0.36 (0.25, 0.51)	0.64 (0.5, 0.81)	Eletriptan	
2.2 (1.74, 2.94)	1.84 (1.5, 2.33)	1.71 (1.31, 2.22)	3.05 (2.68, 3.45)	4.75 (3.78, 5.99)	Placebo

Table D16. All Interventions and Comparators. Pain Relief at 2 Hours (unadjusted NMA)

Table D17. All Interventions and Comparators. Sustained Pain Freedom at 24 Hours (unadjusted NMA)

Lasmiditan		_			
0.78 (0.33, 1.75)	Rimegepant		_		
0.96 (0.38, 2.27)	1.23 (0.56, 2.62)	Ubrogepant			
0.59 (0.26, 1.36)	0.76 (0.4, 1.49)	0.62 (0.29, 1.38)	Sumatriptan		
0.41 (0.15, 1.06)	0.53 (0.22, 1.24)	0.43 (0.17, 1.08)	0.7 (0.28, 1.63)	Eletriptan	
1.99 (1.03, 3.9)	2.57 (1.61, 4.26)	2.09 (1.19, 3.9)	3.39 (2.05, 5.59)	4.86 (2.43, 10.48)	Placebo

Table D18. All Interventions and Comparators. Sustained Pain Relief at 24 Hours

a) Baseline-risk Adjusted NMA

Rimegepant		_		
1.08 (0.77, 1.56)	Ubrogepant			
0.94 (0.72, 1.28)	0.87 (0.64, 1.21)	Sumatriptan		_
0.62 (0.43, 0.92)	0.57 (0.39, 0.85)	0.66 (0.48, 0.88)	Eletriptan	
2.39 (1.93, 3.05)	2.2 (1.69, 2.88)	2.53 (2.1, 3.02)	3.84 (2.9, 5.15)	Placebo

b) Unadjusted NMA

Rimegepant				
0.99 (0.63, 1.61)	Ubrogepant		_	
0.87 (0.62, 1.31)	0.88 (0.59, 1.39)	Sumatriptan		
0.47 (0.3, 0.74)	0.48 (0.28, 0.77)	0.54 (0.35, 0.77)	Eletriptan	
2.18 (1.64, 2.99)	2.21 (1.53, 3.17)	2.49 (1.93, 3.1)	4.59 (3.31, 6.66)	Placebo

Table D19. NMA results. All Interventions and Comparators. Any Adverse Event

Lasmiditan		_			
3.14 (1.64, 6)	Rimegepant		_		
3.51 (1.81, 6.85)	1.12 (0.61, 2.07)	Ubrogepant		_	
2.15 (1.23, 3.65)	0.68 (0.42, 1.1)	0.61 (0.37, 1)	Sumatriptan		_
3.64 (1.97, 6.69)	1.16 (0.66, 2.01)	1.04 (0.58, 1.83)	1.7 (1.16, 2.52)	Eletriptan	
3.91 (2.39, 6.41)	1.25 (0.82, 1.9)	1.11 (0.71, 1.74)	1.82 (1.46, 2.33)	1.07 (0.75, 1.55)	Placebo

NMA: network meta-analysis

Table D20. NMA results. All Interventions and Comparators. Treatment Emergent Adverse Events

Lasmiditan					
4.05 (1.17, 14.08)	Rimegepant		_		
5.27 (2.06, 15.44)	1.3 (0.43, 4.46)	Ubrogepant		_	
2.62 (1.15, 7.18)	0.65 (0.24, 2.11)	0.5 (0.23, 1.13)	Sumatriptan		_
3.36 (0.86, 14.55)	0.83 (0.19, 4.04)	0.64 (0.17, 2.41)	1.28 (0.35, 4.41)	Eletriptan	
6.17 (3.04, 14.45)	1.53 (0.61, 4.32)	1.17 (0.62, 2.22)	2.36 (1.43, 3.72)	1.84 (0.57, 6.04)	Placebo

NMA: network meta-analysis

Table D21. NMA results. All Interventions and Comparators. Dizziness

Lasmiditan		_			
7.19 (2.11, 28.58)	Rimegepant		_		
5.01 (1.59, 17.7)	0.7 (0.16, 2.85)	Ubrogepant		_	
4.41 (1.96, 12.7)	0.62 (0.19, 2.04)	0.89 (0.31, 2.79)	Sumatriptan		_
4.11 (1.39, 14.07)	0.57 (0.14, 2.25)	0.82 (0.23, 3.07)	0.93 (0.34, 2.42)	Eletriptan	
8.68 (4.79, 21.71)	1.23 (0.43, 3.68)	1.75 (0.72, 4.85)	1.98 (1.16, 3.47)	2.15 (0.92, 5.35)	Placebo

NMA: network meta-analysis

Appendix E. Comparative Value Supplemental Information

Table E1. Impact Inventory

	Included in T from Per		Notes on Sources (if	
Sector	Type of Impact (Add Additional Domains, as Relevant)	Health Care Sector	Societal	Quantified), Likely Magnitude & Impact (if not)
Health	Longevity effects	Х	Х	
outcomes	Health-related quality of life effects	Х	Х	
outcomes	Adverse events	Х	Х	
	Paid by third-party payers	Х	Х	
Medical costs	Paid by patients out-of-pocket			
weultal costs	Future related medical costs	Х	Х	
	Future unrelated medical costs			
	Informal Health Care S	ector		
Health-related	Patient time costs	N/A		
costs	Unpaid caregiver-time costs	N/A		
COSIS	Transportation costs	N/A		
	Non-Health Care Sec	tors		
	Labor market earnings lost	N/A	Х	
Productivity	Cost of unpaid lost productivity due to illness	N/A	х	
	Cost of uncompensated household production	N/A		
Consumption	Future consumption unrelated to health	N/A		
Social services	Cost of social services as part of intervention	N/A		
Legal/Criminal	Number of crimes related to intervention	N/A		
justice	Cost of crimes related to intervention	N/A		
Education	Impact of intervention on educational achievement of population	N/A		
Housing	Cost of home improvements, remediation	N/A		
Environment	Production of toxic waste pollution by intervention	N/A		
Other	Other impacts (if relevant)	N/A		

N/A: not applicable

Adapted from Sanders et al.¹³¹

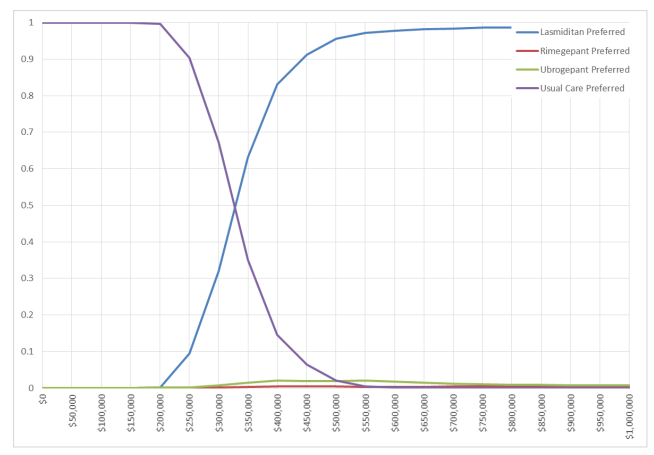


Figure E1. Probabilistic Sensitivity Analysis Results: Acceptability Curve Comparing Lasmiditan, Rimegepant, Ubrogepant, and Usual Care

QALY: quality-adjusted life year

Scenario Analysis 1: Modified Societal Perspective

Labor costs were included in scenario analysis 1. A productivity gain of \$51 per migraine was granted to patients who had "no pain" or "mild pain" at 2 hours. No productivity gain was granted to patients who had "moderate pain" or "severe pain" at 2 hours, even if they achieved a lower pain state beyond 2 hours.

Treatment	Total Cost**	QALYs	Hours of Pain	Cost per QALY Gained (Compared with Usual Care)
Lasmiditan	\$10,320	1.8252	5,524	\$207,800
Rimegepant	\$11,410	1.8222	5,925	\$422,900
Ubrogepant	\$11,420	1.8221	5,944	\$430,900
Usual Care	\$8,040	1.8142	6,652	Comparator

**Drug costs per year were calculated without accounting for discontinuation of the drug. Total costs take into account discontinuation and costs of alternative treatments.

Scenario Analysis 2: Increasing Effectiveness of Gepants Beyond 2-Hours

Exploratory analyses of the impact of rimegepant and ubrogepant on migraine pain after the 2-hour time point suggest that the effect size between active drug and placebo may increase over time.²⁶ ¹³² However, the results are likely confounded by differential dropout between treatment groups in the two arms of each study, making the effect size estimates unreliable. Using the usual care as the baseline response, we modified the model inputs by applying a rate ratio from Lipton et al.¹³² and confidential data provided to us by Allergan at the 24- and 48-hour time points to derive new effectiveness estimates for ubrogepant. Although rimegepant displayed similar results to ubrogepant in published exploratory analyses, we did not have 24- and 48-hour estimates for rimegepant. We therefore ran the scenario analysis using data supplied for ubrogepant only. The table below shows the results of adding these new effect estimates into the model.

Table E3. Costs, QALYs, and Cost-Effectiveness of Gepants Using Data from Exploratory AnalysesSuggesting Continued Growth in Effect Compared with Placebo After 2-Hour Time Point

Treatment	Total Cost**	QALYs	Hours of Pain	Cost per QALY Gained (Compared with Usual Care)
Rimegepant and/or Ubrogepant	\$12,150	1.8295	1,576	\$138,000
Usual Care	\$10,050	1.8142	2,100	Comparator

**Drug costs per year were calculated without accounting for discontinuation of the drug. Total costs take into account discontinuation and costs of alternative treatments.

Scenario Analysis 3: Decreasing Frequency of Migraines Over Time

Data from a long-term open label safety study suggested that the frequency of migraines decreased over time. However, this single arm trial was not designed to evaluate whether the same effect was observed in a control population. The potential for regression to the mean and a high rate of patient drop-out could reasonably be the source of these observed changes in migraine frequency over time. However, decreasing migraine frequency could have a significant impact on budget impact analyses. Therefore, we conducted a scenario analysis evaluating the impact of decreasing migraine frequency over time, resulting in lower total costs, higher QALYs, and fewer hours of pain for all therapies, including usual care. The cost-effectiveness ratios were similar to the base-case. The full results are shown below.

Table E4. Costs, QALYs, and Cost-Effectiveness of Treatments Including Decreasing MigraineFrequency Over Time

Treatment	Total Cost**	QALYs	Hours of Pain	Cost per QALY Gained (Compared with Usual Care)
Lasmiditan	\$9,705	1.8379	1,231	\$328,000
Rimegepant	\$10,294	1.8356	1,321	\$562,400
Ubrogepant	\$10,300	1.8355	1,325	\$572,700
Usual Care	\$7,092	1.8299	1,482	Comparator

**Drug costs per year were calculated without accounting for discontinuation of the drug. Total costs take into account discontinuation and costs of alternative treatments.

Table E5. Cost per QALY Gained and Cost per Additional Hour of Pain Avoided for Lasmiditan,Rimegepant, and Ubrogepant versus Sumatriptan and Eletriptan, with a 5-Year Time Horizon

Intervention	Total Cost	QALYs	Hours of Pain	ICER Compared with Usual Care (cost per additional QALY)	ICER Compared with Usual Care (cost per additional hour of pain avoided)
Lasmiditan	\$31,651	4.3607	4,162	\$326,285	\$8.90
Rimegepant	\$34,115	4.3556	4,466	\$552,107	\$18.24
Ubrogepant	\$34,147	4.3553	4,481	\$562,023	\$18.24
Sumatriptan	\$16,490	4.3643	3,843	Dominates	Dominates
Eletriptan	\$16,834	4.3708	3,539	Dominates	Dominates
Usual Care	\$24,019	4.3373	5,019	Comparator	Comparator

QALY: quality-adjusted life year

Description of the evLYG Calculations

The cost per <u>evLYG</u> considers any extension of life at the same "weight" no matter what treatment is being evaluated. Below are the stepwise calculations used to derive the evLYG.

- 1. First, we attribute a utility of 0.851, the age- and gender-adjusted utility of the general population in the US that are considered healthy.¹³³
- For each cycle (Cycle I) in the model where using the intervention results in additional years of life gained, we multiply this general population utility with the additional life years gained (ΔLYG).
- 3. We sum the product of the life years and average utility (cumulative LYs/cumulative QALYs) for Cycle I in the comparator arm with the value derived in Step 2 to derive the equal value of life years (evLY) for that cycle.
- 4. If no life years were gained using the intervention versus the comparator, we use the conventional utility estimate for that Cycle I.
- 5. The total evLY is then calculated as the cumulative sum of QALYs gained using the above calculations for each arm.
- 6. We use the same calculations in the comparator arm to derive its evLY.

Finally, the evLYG is the incremental difference in evLY between the intervention and the comparator arms.