A Framework for Estimating the Eligible Patient Population for New Migraine Acute Therapies in the United States

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ABSTRACT

Introduction: Migraine is associated with considerable disability for patients not adequately managed with current standards of care. New acute therapies may offer relief for this population of patients; however, population size and associated potential costs of new therapies are unclear. In this study, a conceptual framework was developed to estimate anticipated use of new acute therapies.

Methods: Targeted literature review (TLR) was conducted to identify factors affecting access to migraine-specific acute therapies, and characteristics of individuals who would be eligible for new acute therapies. Findings from the TLR were combined to create a framework for estimating the size of the eligible patient population. This framework was used to calculate two estimates of the eligible patient population by applying parameters (i) identified in the TLR and (ii) from a recent budget-impact analysis (BIA).

Results: The primary factors affecting access to migraine-specific acute therapies identified in the TLR were consulting a healthcare professional for headache, receiving a migraine diagnosis, and receiving a prescription for migraine-specific treatment. Characteristics of individuals likely to use new acute therapies reflected in the TLR were contraindication to triptans, or failure to respond to/tolerate at least two oral triptans. Application of the framework suggested that 15–25% of individuals with migraine would be eligible for new acute therapies.

Conclusion: A limited number of patients currently use migraine-specific acute therapies. Among such patients, a significant proportion do not have adequate symptom control. Accordingly, a minority of individuals with migraine may be expected to use new acute therapies. The framework developed in this study is intended to facilitate estimating the eligible patient population in assessments of costs of new acute therapies. Such assessments should also consider recommendations that patients have access to multiple types of acute therapies, which may yield savings from reduced medication-overuse headache (MOH), progression to chronic migraine, and urgent-
Keywords: Migraine, acute; CGRP; 5-HT\textsubscript{1}\textalpha; Rimegepant; Ubrogepant; Lasmiditan; Triptan; Eligibility; Budget impact; Model

### Key Summary Points

**Why carry out this study?**

Migraine is a common condition affecting approximately 30 million adults and 9 million children and adolescents in the US; symptoms and disability during an attack are managed with acute treatments, including simple analgesics, nonsteroidal anti-inflammatory drugs (NSAIDs), opioid analgesics, butalbital-containing analgesic products, and over-the-counter combinations of analgesics and caffeine, as well as prescription migraine-specific agents which historically have included triptans and ergot derivatives (e.g., dihydroergotamine).

Triptans are commonly used for acute treatment of migraine attacks. However, some patients may not be adequately managed with triptans because of lack of efficacy; intolerable side effects known as “triptan sensations” (including nausea, fatigue, malaise, rapid heart rate, feelings of tingling, numbness, warmth, and chest/neck pressure or tightness); and safety concerns for those with a history of vascular disease, multiple risk factors for vascular disease (such as hypertension and diabetes), and during pregnancy.

The US Food and Drug Administration (FDA) has recently approved three new acute treatments for migraine—rimegepant (NURTEC\textsuperscript{TM} ODT) and ubrogepant (UBRELVY\textsuperscript{®}), both CGRP receptor antagonists, and lasmiditan (REYVOW\textsuperscript{®}), a 5-HT\textsubscript{1}\textalpha; receptor agonist—which the American Headache Society (AHS) has recommended for patients who have contraindications to triptans or who have failed to respond to or tolerate at least two oral triptans.

Understanding the size of the patient population likely to use new treatments may be of interest to payers and health systems; in this study, we therefore developed a conceptual framework for estimating anticipated use of new acute therapies, based on a targeted literature review (TLR) and insights from clinical experience.

**What was learned from the study?**

A minority of individuals with migraine (15%–25%) may be expected to use new acute therapies, given that only a limited proportion of patients currently use migraine-specific acute therapies. Among such patients, a significant proportion do not have adequate symptom control.

The framework developed in this study is intended to facilitate estimating the eligible patient population in assessments of costs of new acute therapies. Such assessments should also consider recommendations that patients have access to multiple types of acute therapies, which may yield savings from reduced MOH, progression to chronic migraine, and urgent-care costs.

### DIGITAL FEATURES

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### INTRODUCTION

Migraine is a neurological condition that manifests as recurrent attacks of headache lasting approximately 4–72 h [1]. The headache may be unilateral and have a pulsating or throbbing quality, is generally worsened by physical activity, and may be associated with nausea/vomiting and/or sensitivity to light and sound.
If unsuccessfully treated, headache severity may progress throughout an attack. Migraine is classified on the basis of monthly frequency of headache days as either episodic (1–14 monthly headache days, MHDs) or chronic (at least 15 MHDs, with features of migraine headache on at least 8 days per month [2]) [1]. In the 2016 Global Burden of Disease Study, migraine was the leading cause of years lived with disability globally in patients younger than 50 years of age, and second only to lower back pain in all patients [3].

Treatments for migraine are typically distinguished as “acute” or “preventive”. Acute treatment is intended to alleviate migraine pain and associated symptoms and to restore ability to function during a migraine attack, while preventive treatment aims to reduce attack frequency, severity, duration, and disability and may improve response to acute therapies [4]. Acute treatments used to treat mild-to-moderate headache include simple analgesics (e.g., aspirin, acetaminophen), nonsteroidal anti-inflammatory drugs (NSAIDs, e.g., ibuprofen, naproxen), opioid analgesics, butalbital-containing analgesic products, and over-the-counter combinations of analgesics including caffeine (e.g., aspirin + acetaminophen + caffeine) [4, 5]. Following diagnosis of migraine, migraine-specific agents may be prescribed for acute treatment, including triptans and ergot derivatives. While effective acute treatment may alleviate symptoms and reduce disability for patients, suboptimal acute treatment may lead to an increase in headache frequency and disability [4]. In particular, overuse of acute treatments is associated with medication-overuse headache (MOH) [6], and increased headache frequency is associated with increased risk of progression from episodic to chronic migraine [7]. Treatment guidelines stress that successful management of an attack is most likely if acute therapy is taken early, and if patients understand the condition and have a treatment strategy [4, 5].

The US Food and Drug Administration (FDA) has recently approved several new oral therapies for acute treatment of migraine, including rimegepant (NURTEC™ ODT, Biohaven Pharmaceuticals) and ubrogepant (UBRELVY™, Allergan), both CGRP receptor antagonists, as well as lasmiditan (REYVOW®; Eli Lilly and Company), a 5-HT1F receptor agonist (and first in its class of the “ditans”). While triptans may provide meaningful relief, patients who experience inadequate relief may “cycle” through the multiple formulations of triptans (seven triptans are available, some with multiple routes of administration [5]), and may use more than the recommended number of doses per month and risk MOH [5]. Some patients may not tolerate side effects of triptans, known as “triptan sensations”, which may include nausea, fatigue, malaise, rapid heart rate, feelings of tingling, numbness, warmth, and pressure or tightness in the chest or neck [8]. Other patients may not be candidates for the triptans. As 5-HT1B/1D agonists, vasoconstriction (attributed to their 1B agonist effects) potentially makes them unsafe for patients with history of vascular disease (either cardiovascular, cerebrovascular, or peripheral vascular disease) or with multiple risk factors for vascular disease. Recently, the American Headache Society (AHS) indicated that new acute treatments should be considered in patients who have contraindications to triptans or who have failed to respond to or tolerate at least two oral triptans [4].

Migraine is a common condition, estimated to affect approximately 30 million adults in the US (15% of the adult population), as well as 9 million children and adolescents. Understanding the size of the patient population likely to use new treatments may therefore be of significant interest to payers and health systems [9]. In this study, we developed a conceptual framework for estimating anticipated use of new acute therapies, based on a targeted literature review (TLR) and insights from clinical experience. The framework identifies factors affecting access to migraine-specific (i.e., prescription) acute therapies and the characteristics of patients who would use new acute therapies. We then illustrated use of the framework by estimating the size of the patient population eligible for new migraine-specific acute therapies in the US, applying parameter values (i) identified in the TLR and (ii) modeled in a recent budget-impact analysis (BIA) of new acute therapies conducted by the Institute for
Clinical and Economic Review (ICER). Finally, considerations for modeling the costs of new acute treatments for migraine, beyond those for estimating the eligible patient population, are discussed on the basis of results of the TLR.

METHODS

The TLR was conducted using the MEDLINE database to identify literature published from 2000 to 2021 on real-world use of migraine-specific (i.e., prescription) acute therapies. Animal studies were excluded, and articles were limited to those published in English and with an abstract. Results of the search were then screened on the basis of title to restrict to real-world studies of acute-therapy use and acute-therapy recommendations/guidelines. Types of results excluded in the screening step included descriptive publications (e.g., reporting on symptoms and complications, pathophysiology, patient-reported outcome tools, and biological mechanisms of side effects of treatments), prospective studies of efficacy of migraine therapies (including acute and preventive), cost-effectiveness analyses, and other studies of non-acute therapies (including non-migraine specific and preventive). Eligibility of remaining publications was confirmed on the basis of review of abstracts; publications were excluded if their abstract suggested they were review articles of efficacy and safety results or studies of non-US populations. A publication’s full text was reviewed to confirm eligibility when deemed necessary by the reviewers (TO and ZH). Following confirmation of eligibility and review of the publications, their findings were summarized by the reviewers and discussed among all authors of this study. Two group discussions were held to review findings of the TLR.

The conceptual framework was developed by relating findings of the TLR and group discussions to the American Headache Society (AHS) recommendation that new acute treatments should be considered in patients who have contraindications to triptans or who have failed to respond to or tolerate at least two oral triptans [4]. Specifically, to estimate the patient population eligible for new acute therapies, factors affecting access to migraine-specific acute therapies were first specified. Then, among individuals who would access migraine-specific acute therapies, characteristics of those who would be eligible for new acute therapies were further specified. For illustration of use of the framework, two estimates of the eligible patient population were calculated by applying parameters (i) identified in the TLR and (ii) from the BIA released on February 25, 2020 in ICER’s Final Evidence Report on treatments for “acute migraine” [10].

This analysis was based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

RESULTS

The TLR included 236 results meeting the search terms and restrictions, 56 of which were retained after screening and eligibility reviews, as illustrated in the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) flow diagram in Fig. 1.

Findings of publications retrieved in the TLR revealed several prerequisites for patient access to migraine-specific acute therapies. Real-world studies suggest that in the US, individuals meeting diagnostic criteria for migraine may not consult a healthcare professional routinely (only 45.5% sought care for their headache in the last year [11] and 79.8% in their lifetime [12]). As described by Lipton et al. [11], the consulting rate is significantly higher in those with insurance coverage vs. those without. Among individuals who do consult a healthcare professional for acute headache, evidence indicates that the majority meet diagnostic criteria for migraine (84% in outpatient clinics [13] and 95% in the emergency department (ED) [14]); nonetheless, a diagnosis of migraine may not be reached. In US studies, the diagnosis rate of migraine among patients meeting diagnostic criteria varies considerably, from 45.7% in the Migraine in America Symptoms and Treatment (MAST) study [12] to 86.7% in the American Migraine Prevalence and Prevention (AMPP)
study [11]. Reaching a diagnosis is particularly challenging in the ED setting [14] as conditions such as anxiety and depression are commonly comorbid with migraine and may confound the cause of a patient’s symptoms [15]. Among consulting patients, women appear to be more likely to receive a migraine diagnosis than men [11, 12].

Among patients diagnosed with migraine and who can therefore be prescribed migraine-specific acute therapies, triptans are commonly recommended as first-line options [4, 5]. However, not all patients are successfully managed with triptans for several reasons, including lack of efficacy and intolerable side effects. In the US, seven types of triptans are available [5], several of which have multiple routes of administration (oral, oral disintegrating tablet, nasal spray, injection) and may be available as generic or branded products [5]. Evidence suggests that determining the triptan formulation with optimal efficacy for a certain patient may therefore require trial of multiple triptans (“cycling”) [16, 17]. However, ability to tolerate the trial of multiple triptans may be limited given their potential known side effects (“triptan sensations”). Further, a significant proportion of patients who might benefit from migraine-specific acute therapy may not be candidates for use of triptans. In patients with vascular disease or with multiple risk factors for vascular disease (e.g., advanced age, hypertension, smoking, obesity, diabetes, family history of coronary artery disease), vasoconstriction attributed to triptans’ 1B agonist effects poses safety concerns [6, 18–20]. In addition, during pregnancy, the benefit of triptans should be weighed against their risk (i.e., they are deemed FDA Category C—see note in supplementary information) [6, 21].

The aforementioned potential limitations of triptans likely contribute to their relatively low
use observed in real-world studies, estimated at 20–30% of individuals meeting diagnostic criteria for migraine (including ca. 20% in the AMPP study [22], 23% in the Chronic Migraine Epidemiology and Outcomes (CaMEO) study [23], 26% in the MAST study [24], and 29% in the Medical Expenditure Panel Survey (MEPS) [25]). Beyond current users of triptans, a significant proportion of patients may have discontinued use; in the CaMEO study, compared to the 23% currently using migraine-specific acute therapy, 13% were former users, among whom only one in five reported being able to work or function normally with a headache [26]. Consistent with the CaMEO study, in a recent study of eligibility for acute treatments using a web-based survey of a representative US sample [27], 18.2% of individuals meeting diagnostic criteria had contraindications to triptans, and 8.6% reported prior use of triptans but did not use them currently [27]. Among patients currently using triptans, approximately half experienced “poor” or “very poor” efficacy according to the migraine Treatment Optimization Questionnaire (mTOQ) (mTOQ < 6) [27].

Figure 2 presents the conceptual framework developed, the steps of which include (1) consulting a healthcare professional for headache, (2) receiving a migraine diagnosis, (3) receiving a prescription for acute migraine-specific treatment, and (4) being inadequately managed on triptans because of lack of efficacy, intolerance, or safety concerns.

The conceptual framework was applied using parameter values identified in the TLR, which yielded an estimate that 21.6% of prevalent migraine patients might be eligible for new acute therapies. Specifically, the AMPP study reports that in respondents meeting diagnostic criteria for migraine, 45.5% consulted a healthcare professional, of whom 86.7% received a medical diagnosis of migraine, and 66.7% of those diagnosed currently used migraine-specific acute therapies [11]. These parameter values were applied in steps 1–3 of the conceptual framework, as reflected in Fig. 2. In step 4, the percentage of migraine-specific acute therapy users inadequately managed with triptans was modeled on the basis of the OVERCOME (ObserVational survey of the Epidemiology, tReatment, and Care Of MigrainE) study [27], due to its recency and distinction of causes of inadequate management with triptans (i.e., lack of efficacy, intolerance, or contraindication). As described in notes to Fig. 2, OVERCOME found that among the prevalent migraine population, 18.2% had contraindications to triptans, and among those with history of use of at least one triptan for acute treatment, patients reported inadequate efficacy or discontinuation of triptans twice as often as satisfactory efficacy. In particular, on the basis of the migraine Treatment Optimization Questionnaire (mTOQ), triptan users who experienced “poor” or “very poor” efficacy (mTOQ < 6) or discontinued were approximately double in number vs. those who experienced “moderate” to “maximum” efficacy (mTOQ ≥ 6).

The conceptual framework was also applied using parameter values reported in the ICER BIA, which yielded an estimate that 17.1% of prevalent migraine patients might be eligible for new acute therapies. In the ICER BIA, two parameters were used to estimate the proportion of prevalent patients eligible for new acute treatments. First, a percentage of prevalent patients using prescription acute treatments was applied. On the basis of the AMPP, ICER modeled that 48.9% of prevalent patients use prescription migraine medications. Second, a percentage of prescription-medication users for whom triptans would not be effective or tolerated was applied. ICER modeled this at 35%, based on an estimate that triptans are effective in 60–70% of migraine patients who have used them [28].

Use of the conceptual framework developed in this study thus illustrated that 15–25% of prevalent migraine patients might be eligible for new acute treatments, as presented in Fig. 2. While estimates based on the two sets of population mean parameter values considered were similar, results of the TLR suggest that they may be sensitive to characteristics of the patient population considered (e.g., see Table 1 of Lipton et al. [11]) and nature of the setting of care (e.g., lower diagnosis rates in the ED [14]).
DISCUSSION

While the conceptual framework developed in this study provides guidance for estimating the number of patients who may try new acute therapies, assessment of total costs also requires estimating the frequency of use of new therapies and the impact on frequency of use of existing therapies. BIAs, including the ICER BIA considered in this report, often assume that eligible patients would use a new therapy rather than an alternative—e.g., that patients would use rimegepant, ubrogepant, or lasmiditan instead of a triptan. However, in acute treatment for migraine, such a substitution may not occur in practice. Some management strategies suggest using a stratified approach [5, 29–33], involving use of different acute therapies.
depending on the severity of a migraine attack, recognizing that the attacks experienced by a single patient may vary considerably in clinical features and severity [4, 34, 35]. Other strategies involve “step care within attacks” [5], whereby a patient would use a simple analgesic initially, but switch to more potent medications later in the same attack, as necessary. Considering that different acute therapies may be needed for effective management in different attacks or within an attack, patients may require access to an array of acute therapies in order to follow a successful management strategy [30, 35–37].

If access to new acute therapies allows patients to follow their recommended management strategy more effectively, assessments of total costs should account for the potential cost offsets associated with improved effectiveness of management [33]. Ineffective acute treatment may lead to MOH [4, 5] or progression to chronic migraine [7, 38], which are associated with higher healthcare resource use and costs [39]. In particular, suboptimal acute management of migraine increases the likelihood of an ED visit [40], and annual frequency of ED visits is associated with higher annual healthcare costs ($3125 in incremental annual costs associated with at least two annual ED visits [41]). In migraine patients, opioids are most commonly prescribed in the ED [42], for patients who have not been able to achieve effective acute management of migraine, and are associated with significantly higher annual healthcare costs (incremental costs of $1985 for 1–2 annual prescriptions, $4107 for 3–6, and $6332 for at least 7 [41]).

Finally, in assessing the costs of acute treatments for migraine, healthcare decision-makers may benefit from considering patient and physician education initiatives around effective acute treatment strategies. Many patients with migraine have never used migraine-specific acute therapies [23, 43] and among those who have, a significant proportion may delay use of acute treatments to a point where they are of lesser effectiveness [44, 45]. Guidelines for acute treatment therefore stress the importance of ensuring that patients understand the condition and their treatment strategy, and that acute therapy be taken early in attacks [4, 5]. Education initiatives may therefore help to minimize ineffective use of acute therapies and associated healthcare resource use (e.g., ED visits) [46, 47].

CONCLUSIONS

In this study, we conducted a TLR which was used to inform development of a conceptual framework for estimating the eligible patient population for new acute therapies for migraine in the US. Using the framework, we illustrated that applying population mean parameter values (i) identified in the TLR and (ii) used in ICER’s BIA of new acute therapies suggested that 15–25% of prevalent migraine patients might be eligible for new acute therapies. Beyond estimation of the eligible patient population, other important factors to consider in assessment of total costs of new acute therapies include that effective management strategies call for access to multiple acute therapies, and that ineffective acute treatment is associated both with considerable burden for patients (including risk of MOH and progression to chronic migraine) as well as significantly higher healthcare resource-use costs. Access to effective acute therapies, for use when necessary, coupled with patient and physician education on recommended management strategies, may significantly impact patient outcomes and costs.

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**Compliance with Ethics Guidelines.** This article is based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

**Data Availability.** Data sharing is not applicable to this article as no datasets were generated or analyzed during the current study.

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