#### QULIPTA- atogepant tablet AbbVie Inc.

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#### HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use QULIPTA safely and effectively. See full prescribing information for QULIPTA.

 QULIPTA® (atogepant) tablets, for oral use Initial U.S. Approval: 2021
 RECENT MAJOR CHANGES

 Indications and Usage (1)
 4/2023

 Dosage and Administration (2.1, 2.2) 4/2023
 Contraindications (4)

 Varnings and Precautions (5.1)
 4/2023

 Warnings and Precautions (5.1)
 4/2023

 QULIPTA is a calcitonin gene-related peptide receptor antagonist indicated for the preventive treatment of migraine in adults. (1)

- ----- DOSAGE AND ADMINISTRATION
- QULIPTA is taken orally with or without food. (2.1)
- For episodic migraine, the recommended dosage is 10 mg, 30 mg, or 60 mg taken once daily. (2.1)
- For chronic migraine, the recommended dosage is 60 mg taken once daily. (2.1)
- Severe Renal Impairment or End-Stage Renal Disease (2.2, 8.6):
  - Episodic migraine: 10 mg once daily.
  - Chronic migraine: Avoid use.

DOSAGE FORMS AND STRENGTHS
Tablets: 10 mg, 30 mg, and 60 mg. (3)
CONTRAINDICATIONS
Patients with a history of hypersensitivity to atogepant or to any of the components of QULIPTA. (4)
WARNINGS AND PRECAUTIONS
If a hypersensitivity reaction occurs, discontinue QULIPTA and initiate appropriate therapy. Severe hypersensitivity reactions have included anaphylaxis and dyspnea. These reactions can occur days after administration. (5.1)
ADVERSE REACTIONS
The most common adverse reactions (at least 4% and greater than placebo) are nausea, constipation, and fatigue/somnolence. (6.1)

## To report SUSPECTED ADVERSE REACTIONS, contact AbbVie at 1-800-678-1605 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

----- DRUG INTERACTIONS ------

Recommended dosage modifications:

- Strong CYP3A4 Inhibitor (2.2, 7.1):
   Episodic migraine: 10 mg once daily.
   Chronic migraine: avoid use.
- Strong, Moderate, or Weak CYP3A4 Inducers (2.2, 7.2):
   C Episodic migraine: 30 mg or 60 mg once daily.
   Chronic migraine: avoid use.
- OATP Inhibitors (2.2, 7.3):
   Episodic migraine: 10 mg or 30 mg once daily.
   Chronic migraine: 30 mg once daily.

• Pregnancy: Based on animal data, may cause fetal harm. (8.1)

• Avoid use in patients with severe hepatic impairment. (8.7)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

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#### FULL PRESCRIBING INFORMATION

### **1** INDICATIONS AND USAGE

QULIPTA is indicated for the preventive treatment of migraine in adults.

#### 2 DOSAGE AND ADMINISTRATION

#### 2.1 Recommended Dosage

QULIPTA is taken orally with or without food.

#### Episodic Migraine

The recommended dosage of QULIPTA for episodic migraine is 10 mg, 30 mg, or 60 mg taken once daily.

#### Chronic Migraine

The recommended dosage of QULIPTA for chronic migraine is 60 mg taken once daily.

#### 2.2 Dosage Modifications

Dosing modifications for concomitant use of specific drugs and for patients with renal impairment are provided in Table 1.

## Table 1: Dosage Modifications for Drug Interactions and for SpecificPopulations

Dosage Modifications	Recommended Once Daily Dosage for Episodic Migraine	Usage and Recommended Once Daily Dosage for Chronic Migraine			
Concomitant Drug [see Dru	g Interactions (7	)]			
Strong CYP3A4 Inhibitors (7.1)	10 mg	Avoid use			
Strong, Moderate, or Weak CYP3A4 Inducers (7.2)	30 mg or 60 mg	Avoid use			
OATP Inhibitors (7.3)	10 mg or 30 mg	30 mg			
Renal Impairment [see Use in Specific Populations (8)]					
Severe Renal Impairment and End-Stage Renal Disease (CLcr <30 mL/min) <b>(</b> 8.6 <b>)</b>	10 mg	Avoid use			

## **3 DOSAGE FORMS AND STRENGTHS**

QULIPTA 10 mg is supplied as white to off-white, round biconvex tablets debossed with "A" and "10" on one side.

QULIPTA 30 mg is supplied as white to off-white, oval biconvex tablets debossed with "A30" on one side.

QULIPTA 60 mg is supplied as white to off-white, oval biconvex tablets debossed with "A60" on one side.

## 4 **CONTRAINDICATIONS**

QULIPTA is contraindicated in patients with a history of hypersensitivity to atogepant or any of the components of QULIPTA. Reactions have included anaphylaxis and dyspnea [see Warnings and Precautions (5.1)].

## 5 WARNINGS AND PRECAUTIONS

## 5.1 Hypersensitivity Reactions

Hypersensitivity reactions, including anaphylaxis, dyspnea, rash, pruritus, urticaria, and facial edema, have been reported with use of QULIPTA [see Adverse Reactions (6.2)]. Hypersensitivity reactions can occur days after administration. If a hypersensitivity reaction occurs, discontinue QULIPTA and institute appropriate therapy [see Contraindications (4)].

## 6 ADVERSE REACTIONS

## 6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The safety of QULIPTA was evaluated in 2657 patients with migraine who received at least one dose of QULIPTA. Of these, 1225 patients were exposed to QULIPTA for at least 6 months, and 826 patients were exposed for 12 months.

In the 12-week, placebo-controlled clinical studies (Studies 1, 2, and 3), 314 patients received at least one dose of QULIPTA 10 mg once daily, 411 patients received at least one dose of QULIPTA 30 mg once daily, 678 patients received at least one dose of QULIPTA 60 mg once daily, and 663 patients received placebo [see Clinical Studies (14)]. Approximately 88% were female, 75% were White, 13% were Black, 10% were Asian, and 10% were of Hispanic or Latino ethnicity. The mean age at study entry was 41 years (range 18 to 74 years).

The most common adverse reactions (incidence at least 4% and greater than placebo) are nausea, constipation, and fatigue/somnolence.

Table 2 summarizes the adverse reactions that occurred during Studies 1, 2, and 3.

# Table 2: Adverse Reactions Occurring with an Incidence of At Least2% for QULIPTA and Greater than Placebo in Studies 1, 2, and 3\*

Placebo (N= 663)	QULIPTA 10 mg (N=314)	QULIPTA 30 mg (N=411)	QULIPTA 60 mg (N=678)
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	/0	%	%	%
Nausea	3	5	6	9
Constipation	2	6	6	8
Fatigue/Somnolence	4	4	4	5
Decreased Appetite	<1	2	1	3
Dizziness	2	2	2	3

\* 10 mg and 30 mg incidence from Studies 1 and 2; 60 mg pooled incidence from Studies 1, 2, and 3.

The adverse reactions that most commonly led to discontinuation of QULIPTA in these studies were nausea (0.6%), constipation (0.5%), and fatigue/somnolence (0.2%).

#### Liver Enzyme Elevations

In Study 1, Study 2, and Study 3, the rate of transaminase elevations over 3 times the upper limit of normal was similar between patients treated with QULIPTA (0.9%) and those treated with placebo (1.2%). However, there were cases with transaminase elevations over 3 times the upper limit of normal that were temporally associated with QULIPTA treatment; these were asymptomatic and resolved within 8 weeks of discontinuation. There were no cases of severe liver injury or jaundice.

#### Decreases in Body Weight

In Study 1, Study 2, and Study 3, the proportion of patients with a weight decrease of at least 7% at any point was 2.5% for placebo, 3.8% for QULIPTA 10 mg, 3.2% for QULIPTA 30 mg, and 5.3% for QULIPTA 60 mg.

## 6.2 Postmarketing Experience

The following adverse reactions have been identified during post approval use of QULIPTA. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to estimate their frequency or establish a causal relationship to drug exposure.

*Immune System Disorders*: Hypersensitivity (e.g., anaphylaxis, dyspnea, rash, pruritus, urticaria, facial edema) [see Contraindications (4) and Warnings and Precautions (5.1)]

## 7 DRUG INTERACTIONS

## 7.1 CYP3A4 Inhibitors

Coadministration of QULIPTA with itraconazole, a strong CYP3A4 inhibitor, resulted in a significant increase in exposure of atogepant in healthy subjects [see Clinical Pharmacology (12.3)]. For episodic migraine, the recommended dosage of QULIPTA with concomitant use of strong CYP3A4 inhibitors is 10 mg once daily. For chronic migraine, avoid concomitant use of strong CYP3A4 inhibitors with QULIPTA [see Dosage and Administration (2.2)]. No dosage adjustment of QULIPTA is needed with concomitant use of moderate or weak CYP3A4 inhibitors.

## 7.2 CYP3A4 Inducers

Coadministration of QULIPTA with steady state rifampin, a strong CYP3A4 inducer, resulted in a significant decrease in exposure of atogepant in healthy subjects [see Clinical Pharmacology (12.3)]. Concomitant administration of QULIPTA with moderate inducers of CYP3A4 can also result in decreased exposure of atogepant. Coadministration of QULIPTA with steady-state topiramate, a weak CYP3A4 inducer, resulted in decreased exposure of atogepant in healthy subjects [see Clinical Pharmacology (12.3)].

For episodic migraine, the recommended dosage of QULIPTA with concomitant use of strong, moderate, or weak CYP3A4 inducers is 30 mg or 60 mg once daily [see Dosage and Administration (2.2)].

For chronic migraine, avoid concomitant use of strong, moderate, or weak CYP3A4 inducers with QULIPTA [see Dosage and Administration (2.2)].

## 7.3 OATP Inhibitors

Coadministration of QULIPTA with single dose rifampin, an OATP inhibitor, resulted in a significant increase in exposure of atogepant in healthy subjects [see Clinical Pharmacology (12.3)]. For episodic migraine, the recommended dosage of QULIPTA with concomitant use of OATP inhibitors is 10 mg or 30 mg once daily. For chronic migraine, the recommended dosage of QULIPTA with concomitant use of OATP inhibitors is 30 mg once daily. For chronic migraine, the recommended dosage of QULIPTA with concomitant use of OATP inhibitors is 30 mg once daily [see Dosage and Administration (2.2)].

## 8 USE IN SPECIFIC POPULATIONS

### 8.1 Pregnancy

#### <u>Risk Summary</u>

There are no adequate data on the developmental risk associated with the use of QULIPTA in pregnant women. In animal studies, oral administration of atogepant during the period of organogenesis (rats and rabbits) or throughout pregnancy and lactation (rats) resulted in adverse developmental effects (decreased fetal and offspring body weight in rats; increased incidence of fetal structural variations in rabbits) at exposures greater than those used clinically [see Data].

In the U.S. general population, the estimated background risk of major birth defects and miscarriages in clinically recognized pregnancies is 2-4% and 15-20%, respectively. The estimated rate of major birth defects (2.2%-2.9%) and miscarriage (17%) among deliveries to women with migraine are similar to rates reported in women without migraine.

### **Clinical Considerations**

Disease-Associated Maternal and/or Embryo/Fetal Risk

Published data have suggested that women with migraine may be at increased risk of preeclampsia and gestational hypertension during pregnancy.

<u>Data</u>

### Animal Data

Oral administration of atogepant (0, 5, 15, 125, or 750 mg/kg/day) to pregnant rats

during the period of organogenesis resulted in decreases in fetal body weight and in skeletal ossification at the two highest doses tested (125 and 750 mg/kg), which were not associated with maternal toxicity. At the no-effect dose (15 mg/kg/day) for adverse effects on embryofetal development, plasma exposure (AUC) was approximately 4 times that in humans at the maximum recommended human dose (MRHD) of 60 mg/day.

Oral administration of atogepant (0, 30, 90, or 130 mg/kg/day) to pregnant rabbits during the period of organogenesis resulted in an increase in fetal visceral and skeletal variations at the highest dose tested (130 mg/kg/day), which was associated with minimal maternal toxicity. At the no-effect dose (90 mg/kg/day) for adverse effects on embryofetal development, plasma exposure (AUC) was approximately 3 times that in humans at the MRHD.

Oral administration of atogepant (0, 15, 45, or 125 mg/kg/day) to rats throughout gestation and lactation resulted in decreased pup body weight at the highest dose tested (125 mg/kg/day), which persisted into adulthood. At the no-effect dose (45 mg/kg/day) for adverse effects on pre- and postnatal development, plasma exposure (AUC) was approximately 5 times that in humans at the MRHD.

## 8.2 Lactation

There are no data on the presence of atogepant in human milk, the effects of atogepant on the breastfed infant, or the effects of atogepant on milk production. In lactating rats, oral dosing with atogepant resulted in levels of atogepant in milk approximately 2-fold higher than that in maternal plasma. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for QULIPTA and any potential adverse effects on the breastfed infant from QULIPTA or from the underlying maternal condition.

## 8.4 Pediatric Use

Safety and effectiveness in pediatric patients have not been established.

## 8.5 Geriatric Use

Population pharmacokinetic modeling suggests no clinically significant pharmacokinetic differences between elderly and younger subjects. Clinical studies of QULIPTA did not include sufficient numbers of patients aged 65 years and over to determine whether they respond differently from younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.

## 8.6 Renal Impairment

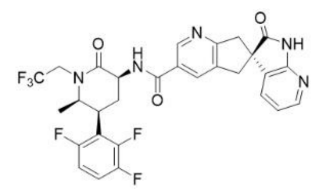
The renal route of elimination plays a minor role in the clearance of atogepant [see Clinical Pharmacology (12.3)]. For episodic migraine, in patients with severe renal impairment (CLcr 15-29 mL/min) and in patients with end-stage renal disease (ESRD) (CLcr <15 mL/min), the recommended dosage of QULIPTA is 10 mg once daily; in patients with ESRD undergoing intermittent dialysis, QULIPTA should preferably be taken after dialysis [see Dosage and Administration (2.2)]. For chronic migraine, avoid use of QULIPTA in patients with severe renal impairment and in patients with ESRD. No dose adjustment is recommended for patients with mild or moderate renal impairment.

## 8.7 Hepatic Impairment

No dose adjustment of QULIPTA is recommended for patients with mild or moderate hepatic impairment. Avoid use of QULIPTA in patients with severe hepatic impairment [see Adverse Reactions (6.1) and Clinical Pharmacology (12.3)].

## 11 DESCRIPTION

The active ingredient of QULIPTA is atogepant, a calcitonin gene-related peptide (CGRP) receptor antagonist. The chemical name of atogepant is (3'S)-*N*-[(3*S*,5*S*,6*R*)-6-methyl-2-oxo-1-(2,2,2-trifluoroethyl)-5-(2,3,6-trifluorophenyl)piperidin-3-yl]-2'-oxo-1',2',5,7-tetrahydrospiro[cyclopenta[*b*]pyridine-6,3'-pyrrolo[2,3-*b*]pyridine]-3-carboxamide, and it has the following structural formula:



The molecular formula is  $C_{29}H_{23}F_6N_5O_3$  and molecular weight is 603.5. Atogepant is a white to off-white powder. It is freely soluble in ethanol, soluble in methanol, sparingly soluble in acetone, slightly soluble in acetonitrile, and practically insoluble in water.

QULIPTA is available as tablets for oral administration containing 10 mg, 30 mg, or 60 mg atogepant. The inactive ingredients include colloidal silicon dioxide, croscarmellose sodium, mannitol, microcrystalline cellulose, polyvinylpyrrolidone vinyl acetate copolymer, sodium chloride, sodium stearyl fumarate, and vitamin E polyethylene glycol succinate.

## 12 CLINICAL PHARMACOLOGY

## 12.1 Mechanism of Action

Atogepant is a calcitonin gene-related peptide (CGRP) receptor antagonist.

## 12.2 Pharmacodynamics

Cardiac Electrophysiology

At a dose 5 times the maximum recommended daily dose, QULIPTA does not prolong the QT interval to any clinically relevant extent.

## 12.3 Pharmacokinetics

**Absorption** 

Following oral administration of QULIPTA, atogepant is absorbed with peak plasma concentrations at approximately 1 to 2 hours. Atogepant displays dose-proportional pharmacokinetics up to 170 mg per day (approximately 3 times the highest recommended dosage), with no accumulation.

### Effect of Food

When QULIPTA was administered with a high-fat meal, the food effect was not significant (AUC and  $C_{max}$  were reduced by approximately 18% and 22%, respectively, with no effect on median time to maximum atogepant plasma concentration). QULIPTA was administered without regard to food in clinical efficacy studies.

## **Distribution**

Plasma protein binding of atogepant was not concentration-dependent in the range of 0.1 to 10  $\mu$ M; the unbound fraction of atogepant was approximately 4.7% in human plasma. The mean apparent volume of distribution of atogepant (Vz/F) after oral administration is approximately 292 L.

## <u>Elimination</u>

## Metabolism

Atogepant is eliminated mainly through metabolism, primarily by CYP3A4. The parent compound (atogepant), and a glucuronide conjugate metabolite (M23) were the most prevalent circulating components in human plasma.

## Excretion

The elimination half-life of atogepant is approximately 11 hours. The mean apparent oral clearance (CL/F) of atogepant is approximately 19 L/hr. Following single oral dose of 50 mg <sup>14</sup>C-atogepant to healthy male subjects, 42% and 5% of the dose was recovered as unchanged atogepant in feces and urine, respectively.

## Specific Populations

## Patients with Renal Impairment

The renal route of elimination plays a minor role in the clearance of atogepant. Based on a population pharmacokinetic analysis, there is no significant difference in the pharmacokinetics of atogepant in patients with mild or moderate renal impairment (CLcr 30-89 mL/min) relative to those with normal renal function (CLcr >90 mL/min). Patients with severe renal impairment or end-stage renal disease (ESRD; CLcr <30 mL/min) have not been studied [see Dosage and Administration (2.2) and Use in Specific Populations (8.6)].

## Patients with Hepatic Impairment

In patients with pre-existing mild (Child-Pugh Class A), moderate (Child-Pugh Class B), or severe (Child-Pugh Class C) hepatic impairment, the total atogepant exposure was increased by 24%, 15%, and 38%, respectively. Due to a potential for liver injury in patients with severe hepatic impairment, avoid use of QULIPTA in patients with severe hepatic impairment *[see Use in Specific Populations (8.7)]*.

## Other Specific Populations

Based on a population pharmacokinetic analysis, age, sex, race, and body weight did not

have a significant effect on the pharmacokinetics (C<sub>max</sub> and AUC) of atogepant. Therefore, no dose adjustments are warranted based on these factors.

#### **Drug Interactions**

In Vitro Studies

### Enzymes

In vitro, atogepant is not an inhibitor for CYPs 3A4, 1A2, 2B6, 2C8, 2C9, 2C19, or 2D6 at clinically relevant concentrations. Atogepant does not inhibit MAO-A or UGT1A1 at clinically relevant concentrations. Atogepant is not anticipated to be a clinically significant perpetrator of drug-drug interactions through CYP450s, MAO-A, or UGT1A1 inhibition.

Atogepant is not an inducer of CYP1A2, CYP2B6, or CYP3A4 at clinically relevant concentrations.

### Transporters

Atogepant is a substrate of P-gp, BCRP, OATP1B1, OATP1B3, and OAT1. Dose adjustment for concomitant use of QULIPTA with inhibitors of OATP is recommended based on a clinical interaction study with a OATP inhibitor [see Dosage and Administration (2.2)].

Coadministration of atogepant with BCRP and/or P-gp inhibitors is not expected to increase the exposure of atogepant. Atogepant is not a substrate of OAT3, OCT2, or MATE1.

Atogepant is not an inhibitor of P-gp, BCRP, OAT1, OAT3, NTCP, BSEP, MRP3, or MRP4 at clinically relevant concentrations. Atogepant is a weak inhibitor of OATP1B1, OATP1B3, OCT1, and MATE1. No clinical drug interactions are expected for atogepant as a perpetrator with these transporters.

### In Vivo Studies

CYP3A4 Inhibitors

Co-administration of QULIPTA with itraconazole, a strong CYP3A4 inhibitor, resulted in a clinically significant increase ( $C_{max}$  by 2.15-fold and AUC by 5.5-fold) in the exposure of atogepant in healthy subjects [see Drug Interactions (7.1)].

Physiologically based pharmacokinetic (PBPK) modeling suggested co-administration of QULIPTA with moderate or weak CYP3A4 inhibitors increase atogepant AUC by 1.7- and 1.1-fold, respectively. The changes in atogepant exposure when coadministered with weak or moderate CYP3A4 inhibitors are not expected to be clinically significant.

### CYP3A4 Inducers

Co-administration of QULIPTA with rifampin, a strong CYP3A4 inducer, decreased atogepant AUC by 60% and  $C_{max}$  by 30% in healthy subjects [see Drug Interactions (7.2)]. No dedicated drug interaction studies were conducted to assess concomitant use with moderate CYP3A4 inducers. Moderate inducers of CYP3A4 can decrease atogepant exposure [see Drug Interactions (7.2)]. Co-administration of QULIPTA with topiramate, a weak inducer of CYP3A4, decreased atogepant mean steady-state AUC  $_{0-\tau}$  by 25% and mean steady-state C<sub>max</sub> by 24% in healthy subjects [see Drug Interactions (7.2)].

BCRP/OATP/P-gp Inhibitors

Co-administration of QULIPTA with single dose rifampin, an OATP inhibitor, increased atogepant AUC by 2.85-fold and C<sub>max</sub> by 2.23-fold in healthy subjects [see Drug Interactions (7.3)].

Co-administration of QULIPTA with quinidine, a P-gp inhibitor, increased atogepant AUC by 26% and  $C_{max}$  by 4% in healthy subjects. The changes in atogepant exposure when co-administered with P-gp inhibitors are not expected to be clinically significant.

PBPK modeling suggests that co-administration of QULIPTA with BCRP inhibitors increases atogepant exposure by 1.2-fold. This increase is not expected to be clinically significant.

#### Other Drug Interaction Evaluations

Co-administration of QULIPTA with oral contraceptive components ethinyl estradiol and levonorgestrel, famotidine, esomeprazole, acetaminophen, naproxen, sumatriptan, or ubrogepant did not result in significant pharmacokinetic interactions for either atogepant or co-administered drugs. Co-administration of QULIPTA with topiramate did not result in clinically significant changes in the pharmacokinetics of topiramate.

## 13 NONCLINICAL TOXICOLOGY

## 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

#### **Carcinogenicity**

Atogepant was administered orally to mice (0, 5, 20, or 75 mg/kg/day in males; 0, 5, 30, 160 mg/kg/day in females) and rats (0, 10, 20, or 100 mg/kg in males; 0, 25, 65, or 200 mg/kg in females) for up to 2 years. There was no evidence of drug-related tumors in either species. Plasma exposures at the highest doses tested in mice and rats were approximately 8 and 20-35 times, respectively, that in humans at the maximum recommended human dose (MRHD) of 60 mg/day.

#### <u>Mutagenicity</u>

Atogepant was negative in in vitro (Ames, chromosomal aberration test in Chinese Hamster Ovary cells) and in vivo (rat bone marrow micronucleus) assays.

### Impairment of Fertility

Oral administration of atogepant (0, 5, 20, or 125 mg/kg/day) to male and female rats prior to and during mating and continuing in females to Gestation Day 7 resulted in no adverse effects on fertility or reproductive performance. Plasma exposures (AUC) at the highest dose tested are approximately 15 times that in humans at the MRHD.

## 14 CLINICAL STUDIES

## 14.1 Episodic Migraine

The efficacy of QULIPTA for the preventive treatment of episodic migraine in adults was demonstrated in two randomized, multicenter, double-blind, placebo-controlled studies (Study 1 and Study 2). The studies enrolled patients with at least a 1-year history of migraine with or without aura, according to the International Classification of

Headache Disorders (ICHD-3) diagnostic criteria.

In Study 1 (NCT03777059), 910 patients were randomized 1:1:1:1 to receive QULIPTA 10 mg (N = 222), QULIPTA 30 mg (N = 230), QULIPTA 60 mg (N = 235), or placebo (N = 223), once daily for 12 weeks. In Study 2 (NCT02848326), 652 patients were randomized 1:2:2:2 to receive QULIPTA 10 mg (N = 94), QULIPTA 30 mg (N = 185), QULIPTA 60 mg (N = 187), or placebo (N = 186), once daily for 12 weeks. In both studies, patients were allowed to use acute headache treatments (i.e., triptans, ergotamine derivatives, NSAIDs, acetaminophen, and opioids) as needed. The use of a concomitant medication that acts on the CGRP pathway was not permitted for either acute or preventive treatment of migraine. The studies excluded patients with myocardial infarction, stroke, or transient ischemic attacks within six months prior to screening.

### Study 1

The primary efficacy endpoint was the change from baseline in mean monthly migraine days (MMD) across the 12-week treatment period. Secondary endpoints included the change from baseline in mean monthly headache days, the change from baseline in mean monthly acute medication use days, the proportion of patients achieving at least a 50% reduction from baseline in mean MMD (3-month average), the change from baseline in mean monthly Activity Impairment in Migraine-Diary (AIM-D) Performance of Daily Activities (PDA) domain scores, the change from baseline in mean monthly AIM-D Physical Impairment (PI) domain scores, across the 12-week treatment period, and the change from baseline at Week 12 for Migraine Specific Quality of Life Questionnaire version 2.1 (MSQ v2.1) Role Function-Restrictive (RFR) domain scores.

The AIM-D evaluates difficulty with performance of daily activities (PDA domain) and physical impairment (PI domain) due to migraine, with scores ranging from 0 to 100. Higher scores indicate greater impact of migraine, and reductions from baseline indicate improvement. The MSQ v2.1 Role Function-Restrictive (RFR) domain score assesses how often migraine impacts function related to daily social and work-related activities over the past 4 weeks, with scores ranging from 0 to 100. Higher scores indicate lesser impact of migraine on daily activities, and increases from baseline indicate improvement.

Patients had a mean age of 42 years (range 18 to 73 years), 89% were female, 83% were White, 14% were Black, and 9% were of Hispanic or Latino ethnicity. The mean migraine frequency at baseline was approximately 8 migraine days per month and was similar across treatment groups. A total of 805 (88%) patients completed the 12-week double-blind study period. Key efficacy results of Study 1 are summarized in Table 3.

Monthly Migraine	QULIPTA 10 mg N=214 Davs (MMD) acro	QULIPTA 30 mg N=223 oss 12 weeks	QULIPTA 60 mg N=222	Placebo N=214
Baseline	7.5	7.9	7.8	7.5
Mean change from baseline	-3.7	-3.9	-4.2	-2.5
Difference from placebo	-1.2	-1.4	-1.7	

## Table 3: Efficacy Endpoints in Study 1

1 <0.001
<u></u>
9.0 8.4
-4.2 -2.5
-1.7
1 <0.001
2 weeks
6.9 6.5
-3.9 -2.4
-1.5
1 <0.001
ł
61 29
32
1 <0.001
I
46.8 46.8
31.3 20.5
10.8
1 <0.001
I
15.9 15.2
-9.4 -6.1
-3.3
1 <0.001
11.6 11.2
-6.5 -4.0
-2.5
<0.001

\*\*\* Activity Impairment in Migraine-Diary Physical Impairment domain score <sup>†</sup>Not statistically significant (NS) Figure 1 shows the mean change from baseline in MMD in Study 1. Patients treated with QULIPTA had greater mean decreases from baseline in MMD across the 12-week treatment period compared to patients who received placebo.

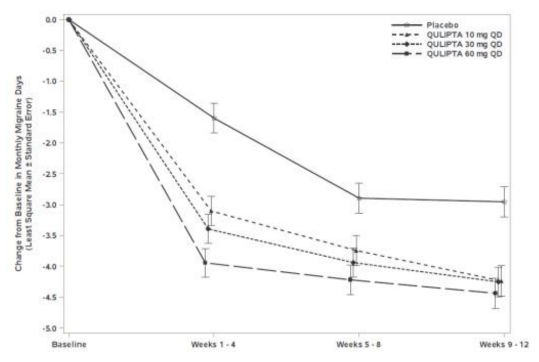
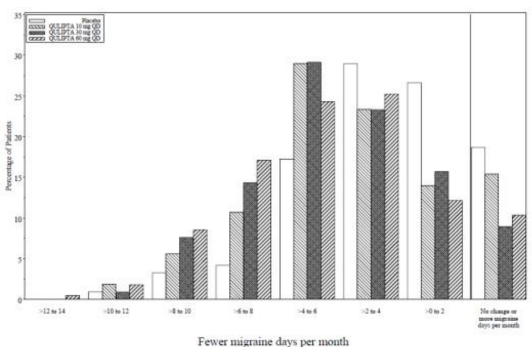


Figure 1: Change from Baseline in Monthly Migraine Days in Study 1

Figure 2 shows the distribution of change from baseline in mean MMD across the 12week treatment period, in 2-day increments, by treatment group. A treatment benefit over placebo for all doses of QULIPTA is seen across a range of mean changes from baseline in MMD.





Study 2

The primary efficacy endpoint was the change from baseline in mean monthly migraine days across the 12-week treatment period.

Patients had a mean age of 40 years (range: 18 to 74 years), 87% were female, 76% were White, 20% were Black, and 15% were of Hispanic or Latino ethnicity. The mean migraine frequency at baseline was approximately 8 migraine days per month. A total of 541 (83%) patients completed the 12-week double-blind study period.

In Study 2, there was a significantly greater reduction in mean monthly migraine days across the 12-week treatment period in all three QULIPTA treatment groups, compared with placebo, as summarized in Table 4.

	QULIPTA 10 mg N=92	QULIPTA 30 mg N=182	QULIPTA 60 mg N=177	Placebo N=178
Monthly Migraine Days (MM	1D) across 1	.2 weeks		
Baseline	7.6	7.6	7.7	7.8
Mean change from baseline	-4.0	-3.8	-3.6	-2.8
Difference from placebo	-1.1	-0.9	-0.7	
<i>p</i> -value	0.024	0.039	0.039	
Monthly Headache Days ac	ross 12 wee	eks		
Baseline	8.9	8.7	8.9	9.1
Mean change from baseline	-4.3	-4.2	-3.9	-2.9
Difference from placebo	-1.4	-1.2	-0.9	
<i>p</i> -value	0.024	0.039	0.039	

### Table 4: Efficacy Endpoints in Study 2

Figure 3 shows the mean change from baseline in MMD in Study 2. Patients treated with QULIPTA had greater mean decreases from baseline in MMD across the 12-week treatment period compared to patients who received placebo.

#### Figure 3: Change from Baseline in Monthly Migraine Days in Study 2

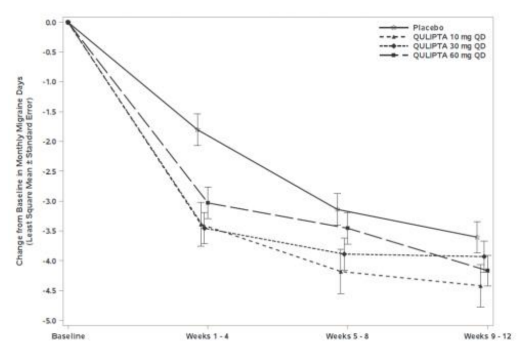
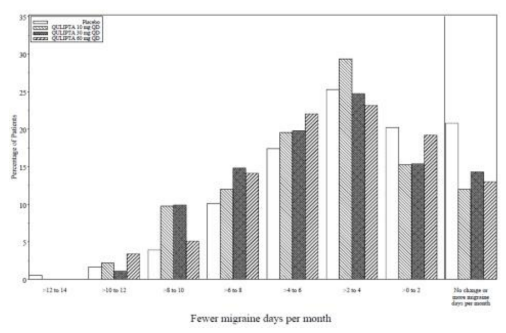


Figure 4 shows the distribution of change from baseline in mean MMD across the 12week treatment period, in 2-day increments, by treatment group. A treatment benefit over placebo for all doses of QULIPTA is seen across a range of mean changes from baseline in MMD.

Figure 4: Distribution of Change from Baseline in Mean Monthly Migraine Days by Treatment Group in Study 2



### 14.2 Chronic Migraine

### Study 3

The efficacy of QULIPTA for the preventive treatment of chronic migraine in adults was demonstrated in a randomized, multicenter, double-blind, placebo-controlled study (Study 3). The study enrolled patients with at least a 1-year history of chronic migraine, according to the ICHD-3 diagnostic criteria.

Study 3 (NCT03855137) included randomization of patients to QULIPTA 60 mg once daily (N = 262) or placebo (N = 259) for 12 weeks. A subset of patients (11%) was allowed to use one concomitant migraine preventive medication. Patients were allowed to use acute headache treatments (i.e., triptans, ergotamine derivatives, NSAIDs, acetaminophen, and opioids) as needed. Patients with medication overuse headache also were enrolled. The use of a concomitant medication that acts on the CGRP pathway was not permitted for either acute or preventive treatment of migraine. The study excluded patients with myocardial infarction, stroke, or transient ischemic attacks within six months prior to screening.

The primary efficacy endpoint was the change from baseline in mean MMD across the 12-week treatment period. Secondary endpoints included the change from baseline in mean monthly headache days, the change from baseline in mean monthly acute medication use days, the proportion of patients achieving at least a 50% reduction from baseline in mean MMD (3-month average), the change from baseline in mean monthly AIM-D PDA domain scores, the change from baseline in mean monthly AIM-D PI domain scores, across the 12-week treatment period, and the change from baseline at Week 12 for MSQ v2.1 RFR domain scores.

Patients had a mean age of 42 years (range 18 to 74 years), 87% were female, 60% were White, 3% were Black, 36% were Asian, and 4% were of Hispanic or Latino ethnicity. The mean migraine frequency at baseline was approximately 19 migraine days per month and was similar across treatment groups. A total of 463 (89%) of these patients completed the 12-week double-blind study period.

Key efficacy results of Study 3 are summarized in Table 5.

	QULIPTA 60 mg QD N=256	Placebo N=246
Monthly Migraine Days (MMD)	across 12 weeks	
Baseline	19.2	18.9
Mean change from baseline	-6.9	-5.1
Difference from placebo	-1.8	
<i>p</i> -value	< 0.001	
Monthly Headache Days acros	s 12 weeks	
Baseline	21.5	21.4
Mean change from baseline	-7.0	-5.1
Difference from placebo	-1.9	
<i>p</i> -value	< 0.001	
Monthly Acute Medication Use	e Days across 12 we	eks
Baseline	15.5	15.4
Mean change from baseline	-6.2	-4.1
Difference from placebo	-2.1	
<i>p</i> -value	< 0.001	
≥ 50% MMD Responders acro	ss 12 weeks	
% Responders	41	26

### Table 5: Efficacy Endpoints in Study 3

Difference from placebo (%)	15	
<i>p</i> -value	< 0.001	
MSQ v2.1 RFR Domain* at week	12	
Baseline	43.4	43.9
Mean change from baseline	23.3	17.2
Difference from placebo	6.2	
p-value	< 0.001	
AIM-D PDA Domain** across 12	weeks	
Baseline	31.2	29.5
Mean change from baseline	-12.8	-9.4
Difference from placebo	-3.4	
p-value	< 0.001	
AIM-D PI Domain*** across 12 w	veeks	
Baseline	27.1	25.2
Mean change from baseline	-10.6	-7.9
Difference from placebo	-2.7	
p-value	0.003	

\* Migraine Specific Quality of Life Questionnaire version 2.1 Role Function-Restrictive domain score

\*\* Activity Impairment in Migraine-Diary Performance of Daily Activities domain score

\*\*\* Activity Impairment in Migraine-Diary Physical Impairment domain score

Figure 5 shows the mean change from baseline in MMD in Study 3. Patients treated with QULIPTA had greater mean decreases from baseline in MMD across the 12-week treatment period compared to patients who received placebo.

#### Figure 5: Change from Baseline in Monthly Migraine Days in Study 3

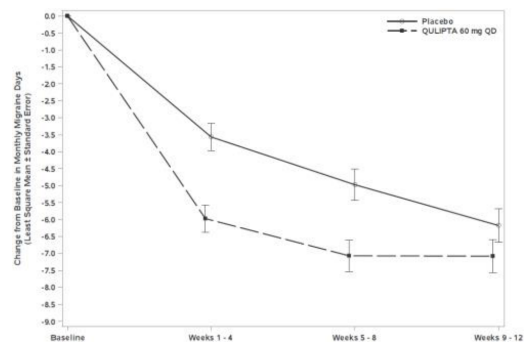
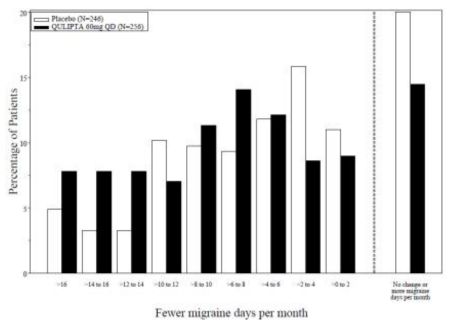


Figure 6 shows the distribution of change from baseline in mean MMD across the 12-

week treatment period, in 2-day increments, by treatment group. A treatment benefit of QULIPTA over placebo is seen across a range of mean changes from baseline in MMD.





## 16 HOW SUPPLIED/STORAGE AND HANDLING

### 16.1 How Supplied

QULIPTA 10 mg is supplied as white to off-white, round biconvex tablets debossed with "A" and "10" on one side in the following packaging presentations:

• Bottle of 30, NDC: 0074-7095-30

QULIPTA 30 mg is supplied as white to off-white, oval biconvex tablets debossed with "A30" on one side in the following packaging presentations:

• Bottle of 30, NDC: 0074-7096-30

QULIPTA 60 mg is supplied as white to off-white, oval biconvex tablets debossed with "A60" on one side in the following packaging presentations:

• Bottle of 30, NDC: 0074-7094-30

### 16.2 Storage and Handling

Store between 20°C and 25°C (68°F and 77°F): excursions permitted between 15°C and 30°C (59°F and 86°F) [see USP Controlled Room Temperature].

## 17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Patient Information).

Hypersensitivity Reactions

Inform patients about the signs and symptoms of hypersensitivity reactions and that

these reactions can occur with QULIPTA. Advise patients to discontinue QULIPTA and seek immediate medical attention if they experience any symptoms of a hypersensitivity reaction *[see Warnings and Precautions (5.1)]*.

#### **Drug Interactions**

Inform patients that QULIPTA may interact with certain other drugs, and that dosage modifications of QULIPTA may be recommended when used with some other drugs. Advise patients to report to their healthcare provider the use of any other prescription medications, over-the-counter medications, herbal products, or grapefruit juice [see Dosage and Administration (2.2) and Drug Interactions (7.1, 7.2, 7.3)].

Manufactured for: AbbVie Inc. North Chicago, IL 60064

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#### PATIENT INFORMATION QULIPTA<sup>®</sup> (kew-LIP-tah)

(atogepant) tablets, for oral use

#### What is QULIPTA?

• QULIPTA is a prescription medicine used for the preventive treatment of migraine in adults.

It is not known if QULIPTA is safe and effective in children.

#### Do not take QULIPTA if you:

• have had an allergic reaction to atogepant or any ingredients in QULIPTA. See the end of this Patient Information leaflet for a complete list of ingredients in QULIPTA.

## Before you take QULIPTA tell your healthcare provider about all of your medical conditions, including if you:

- have kidney problems or are on dialysis.
- have liver problems.
- are pregnant or plan to become pregnant. It is not known if QULIPTA will harm your unborn baby.
- are breastfeeding or plan to breastfeed. It is not known if QULIPTA passes into your breast milk. Talk to your healthcare provider about the best way to feed your baby while taking QULIPTA.

**Tell your healthcare provider about all the medicines you take,** including prescription and over-the-counter medicines, vitamins, and herbal supplements. QULIPTA may affect the way other medicines work, and other medicines may affect how QULIPTA works. Your healthcare provider may need to change the dose of QULIPTA when taken with certain other medicines.

Keep a list of medicines you take to show to your healthcare provider or pharmacist when you get a new medicine.

## How should I take QULIPTA?

- Take QULIPTA by mouth with or without food.
- Take QULIPTA exactly as your healthcare provider tells you to take it.

#### What are the possible side effects of QULIPTA? QULIPTA can cause serious side effects, including:

Allergic (hypersensitivity) reactions, including anaphylaxis: Serious allergic reactions can happen when you take QULIPTA or days after. Stop taking QULIPTA and get emergency medical help right away if you get any of the following symptoms, which may be part of a serious allergic reaction:

- swelling of the face, lips, or tongue itching
- trouble breathing
   • hives
- rash

**The most common side effects of QULIPTA include:** nausea, constipation, and fatigue/sleepiness.

These are not all of the possible side effects of QULIPTA. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

#### How should I store QULIPTA?

• Store QULIPTA at room temperature between 68°F to 77°F (20°C to 25°C).

## Keep QULIPTA and all medicines out of the reach of children.

### General information about the safe and effective use of QULIPTA.

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not use QULIPTA for a condition for which it was not prescribed. Do not give QULIPTA to other people, even if they have the same symptoms you have. It may harm them. You can ask your pharmacist or healthcare provider for information about QULIPTA that is written for health professionals.

#### What are the ingredients in QULIPTA?

Active ingredient: atogepant

**Inactive ingredients:** colloidal silicon dioxide, croscarmellose sodium, mannitol, microcrystalline cellulose, polyvinylpyrrolidone vinyl acetate copolymer, sodium chloride, sodium stearyl fumarate, and vitamin E polyethylene glycol succinate.

Manufactured for: AbbVie Inc. North Chicago, IL 60064 © 2023 AbbVie. All rights reserved. QULIPTA and its design are trademarks of Allergan Pharmaceuticals International Limited, an AbbVie company.

This Patient Information has been approved by the U.S. Food and DrugAdministrationRevised: 6/2023

20078877

## PRINCIPAL DISPLAY PANEL

NDC 0074-**7095**-30 Rx Only **QULIPTA**<sup>®</sup> (atogepant) tablets **10 mg** 

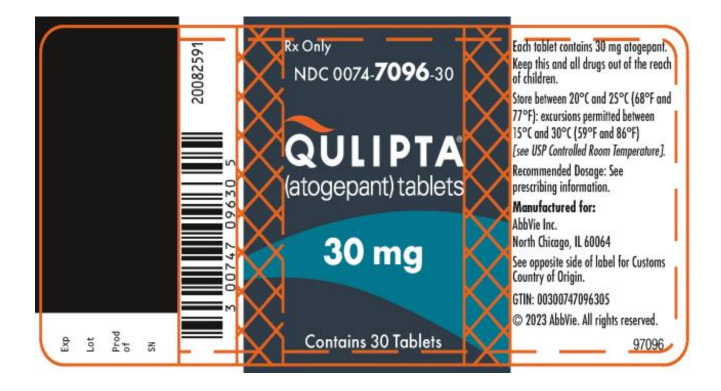
Contains 30 Tablets



### PRINCIPAL DISPLAY PANEL

NDC 0074-**7096**-30 Rx Only **QULIPTA**<sup>®</sup> (atogepant) tablets

**30 mg** Contains 30 Tablets



## PRINCIPAL DISPLAY PANEL

NDC 0074-**7094**-30 Rx Only **QULIPTA®** (atogepant) tablets

### 60 mg

Contains 30 Tablets



Due due toto							
Product Inform	nation						
Product Type		HUMAN PRESCR	IPTION DRUG	Item C	ode (Source)	NDC	:0074-7095
Route of Adminis	tration	ORAL ORAL					
Active Ingredie	nt/Active	Moiety					
gg	-	dient Name			Basis of Stre	nath	Strength
ATOGEPANT (UNII: 7	•		II:7CRV8RR151)		ATOGEPANT		10 mg
	, (						
Inactive Ingred	lients						
		Ingredier	nt Name				Strength
SILICON DIOXIDE (U	INII: ETJ7Z6XE	8U4)					
CROSCARMELLOSE	SODIUM (UN	III: M28OL1HH48)					
MANNITOL (UNII: 30)	WL53L36A)						
MICROCRYSTALLINE		E (UNII: OP1R32D	61U)				
COPOVIDONE K25-3	<b>31</b> (UNII: D9C	330MD8B)					
SODIUM CHLORIDE	(UNII: 451W47	7IQ8X)					
SODIUM STEARYL F	UMARATE (U	NII: 7CV7WJK4UI)					
VITAMIN E POLYETH	IYLENE GLY	COL SUCCINATI	(UNII: 003590U1	F2)			
Product Charac	torictics						
		to	<b>6</b>				
Color	whi		Score			o score	
Shape 	ROU	JND	Size				
Flavor	Imprint Code A;10						
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## QULIPTA

atogepant tablet

**Product Information** 

Product Type		HUMAN PRES	CRIPTION DRUG	ltem	Code (Source)	NDC	2:0074-7096
Route of Admin	istration	ORAL					
Active Ingred	lient/Activ	/e Moietv					
		redient Name	2		Basis of St	renath	Strengt
ATOGEPANT (UNII					ATOGEPANT		30 mg
Inactive Ingre	edients						
		Ingred	ient Name				Strength
SILICON DIOXIDE	(UNII: ETJ7Z	6XBU4)					
CROSCARMELLOS	SE SODIUM	(UNII: M28OL1HH	48)				
MANNITOL (UNII: 3	30WL53L36A)						
MICROCRYSTALL			32D61U)				
COPOVIDONE K2							
SODIUM CHLORIE							
SODIUM STEARYI							
VITAMIN E POLYE	THYLENE G	LYCOL SUCCIN	<b>ATE</b> (UNII: 003590U)	1F2)			
Product Char	acteristic		-				
Color		white	Score			no score	
Shape		OVAL	Size			7mm	
Flavor			Imprint Code		ŀ	430	
Contains							
Packaging							
# Item Code	i	Package Des	cription	Mark	eting Start Date		eting End Date
<b>1</b> NDC:0074-7096-30	- 30 in 1 BO Product	TTLE; Type 0: No	t a Combination	09/30/20	21		
<b>2</b> NDC:0074-7096-04	- 4 in 1 BOT Product	TLE; Type 0: Not	a Combination	10/06/20	21		
Marketing	Informa	ation					
Marketing Category	Appli	cation Numbe Citat	er or Monograph ion	Ma	rketing Start Date	Mar	keting End Date
NDA	NDA2152	206		09/30	/2021		
QULIPTA							
atogepant table	t						
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Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0074-7094

Ingredient Name       Basis of Strength       Strength         ATOGEPANT (UNII: 7CRV8RR151) (ATOGEPANT - UNII:7CRV8RR151)       ATOGEPANT       60 mg         Inactive Ingredients       Ingredient Name       Strength         Inactive Ingredients       Ingredient Name       Strength         Inactive Ingredients       Ingredient Name       Strength         Sulcon DIOXIDE (UNII: ETJ7Z 6XBU4)       Strength       Strength         CROSCARMELLOSE SODIUM (UNII: M280L1HH48)       MANITOL (UNII: 300453136A)       Ingredient Name       Strength         MANITOL (UNII: 300453136A)       Ingredient Name       Ingredient Name       Ingredient Name         SoDIUM CHLORIDE (UNII: 4514947108X)       Ingredient Name       Ingredient Name       Ingredient Name         SoDIUM CHLORIDE (UNII: 4514947108X)       Ingredient Name       Ingredient Name       Ingredient Name         SoDIUM CHLORIDE (UNII: 4514947108X)       Ingredient Name       Ingredient Name       Ingredient Name         SoDIUM CHLORIDE (UNII: 451497108X)       Ingredient Code       Ingredient Name       Ingredient Name         Product Characteristics       Ingrint Code       Noscore       Ingredient Name       Ingredient Name         Product Characteristics       Ingrint Code       Nafeting Start       Marketing En Date         1       NDC:007	Ro								
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Ingredient Name       Basis of Strength       Strength         ATOGEPANT (UNII: 7CRVBRR151) (ATOGEPANT - UNII:7CRVBRR151)       ATOGEPANT       60 mg         Inactive Ingredients       Ingredient Name       Strength         Inactive Ingredients       Ingredient Name       Strength         SILCON DIOXIDE (UNII: ETJ7Z 6XBU4)       Strength       Strength         CROSCARMELLOSE SODIUM (UNII: M280L1HH48)       MANITOL (UNII: 300453136A)       Strength         MICROCRYSTALLINE CELLULOSE (UNII: 01R32D61U)       Strength       Strength         COPONDONE K23-31 (UNII: 902300MB8)       SoDIUM CHLORIDE (UNII: 451W47108X)       Strength         SoDIUM CHLORIDE (UNII: 451W47108X)       Strength       Strength         SoDIUM CHLORIDE (UNII: 451W47108X)       Strength       Strength         SoDIUM CHLORIDE (UNII: 451W47108X)       Strength       Strength         Strength       Markati (UNII: 902300MB8)       Strength         SoDIUM CHLORIDE (UNII: 451W47108X)       Strength       Strength         Strength       Markating Ength       Strength       Strength         Strength       Marketing Start       Marketing Ength       Strength         Strength       Marketing Category       Marketing Ength       Strength         Strength       Strength       Strength									
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Ingredient Name       Strengt         Silicon Dioxide (UNII: ETJ7Z6XBU4)         CROSCARMELLOSE SODIUM (UNII: M280L1HH48)         MANNITOL (UNII: 30WL53L36A)         MICROCRYSTALLINE CELLULOSE (UNII: OPIR32D61U)         COPOVIDONE K25-31 (UNII: OPIR32D61U)         COPOVIDONE K25-31 (UNII: OPIR32D61U)         COPOVIDONE K25-31 (UNII: OPIR32D61U)         Sodium STEARYL FUMARATE (UNII: 7CV7WJK4UI)         VILCOL SUCCINATE (UNII: 003590U1F2)         Product Characteristics         Color       white       Score       no score         Sodium Steary Fumarate (UNII: 7CV7WJK4UI)       VILCOL SUCCINATE (UNII: 003590U1F2)         Product Characteristics         Color       white       Score       no score         Shape       OVAL       Size       9mm         Flavor       Imprint Code       A60         Contains       09/30/2021         Imprint Code       Marketing En Date         1       NDC:074-7094       Ain 1 BOTTLE; Type 0: Not a Combination			Ingr	edient Name	е		Basis of Str	ength	Strengt
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Ingredient Name       Strengt         Silicon Dioxide (UNII: ETJ726XBU4)       CROSCARMELLOSE SODIUM (UNII: M280L1HH48)         MANNITOL (UNII: 30WL53L36A)       Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Solum ChloRide CUNII: 451W47IQ8X)         Solum ChloRide (UNII: 451W47IQ8X)       Good Colspan="2">Good Colspan="2">Good Colspan="2">Good Colspan="2">Marketing Start (UNII: 7CV7WK4UI)         Yorduct Characteristics         Color       white       Score       no score         Shape       OVAL       Size       9mm         Flavor       Marketing Start       Marketing Start       Marketing Start       Marketing Start       Marketing Start       Marketing En         Application Number or Monograph       Marketing Start       Marketing Start       Marketing Start									
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SILICON DIOXIDE (UNII: ETJ7Z6XBU4) CROSCARMELLOSE SODIUM (UNII: M280L1HH48) MANNITOL (UNII: 30WL53L36A) MICROCRYSTALLINE CELLULOSE (UNII: 0P1R3ZD61U) COPOVIDONE K25-31 (UNII: 451W47IQ8X) SODIUM CHLORIDE (UNII: 1451W47IQ8X) SODIUM STEARYL FUMARATE (UNII: 7CV7WK4UI) VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 003S90U1F2) FOUCT Characteristics Color white Score no score Shape 00VAL Size 9mm Flavor 0VAL Size 9mm Flavor A60 Contains Marketing Start Marketing En Date Marketing Information Froduct Application Number or Monograph Marketing Start Date Marketing En Tote Characterist (Start Number or Monograph Marketing Start Date Marketing En Date Marketing Category Application Number or Monograph Marketing Start Date Marketing En Date Marketing Category Application Number or Monograph Marketing Start Date Marketing En Date Date Marketing En Date Marketing En D	In	nactive Ingred	dients						
CROSCARMELLOSE SODIUM (UNII: M280L1HH48)         Image: Sodium (UNII: 30WL53L36A)         Image: Sodium (UNII: 30WL57L3A)         Image: Sodium (UNII: 30WL57L4A)				Ingred	ient Name				Strength
MANNITOL (UNII: 30WL53L36A)       Image: Cellulose (UNII: 0P1R32D61U)       Image: Cellulose (UNII: 0P1R32D61U)         COPOVIDONE K25-31 (UNII: D9C330MD8B)       SODIUM CHLORIDE (UNII: 451W47IQ8X)       Image: Cellulose (UNII: 451W47IQ8X)         SODIUM CHLORIDE (UNII: 451W47IQ8X)       SODIUM STEARYL FUMARATE (UNII: 7CV7WK4UI)       Image: Cellulose (UNII: 603590U1F2)         VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 003590U1F2)       Image: Cellulose (UNII: 603590U1F2)       Image: Cellulose (UNII: 603590U1F2)         Product Characteristics         Color       white       Score       no score         Size       9mm         Imprint Code       Marketing Start       Marketing En         Contains         Package Description       Marketing Start       Marketing En         Date         Intornation         1 NDC:0074-7094-       30 in 1 BOTTLE; Type 0: Not a Combination       09/30/2021       Imprint Code       Imprint Co	SI	LICON DIOXIDE (I	JNII: ETJ7Z6	XBU4)					
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U) COPOVIDONE K25-31 (UNII: D9C330MD8B) SODIUM CHLORIDE (UNII: 451W47IQ8X) SODIUM STEARYL FUMARATE (UNII: 7CV7WjK4UI) VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 003590U1F2) Product Characteristics Color white Size no score Shape 0VAL Size 9mm Flavor A60 Contains 100 Package Description Marketing Start Date Marketing En Date 1 1 NDC:0074-7094 30 in 1 BOTTLE; Type 0: Not a Combination 9/30/2021 2 NDC:0074-7094 4 in 1 BOTTLE; Type 0: Not a Combination 10/06/2021 2 NDC:0074-7094 4 in 1 BOTTLE; Type 0: Not a Combination 10/06/2021 Marketing Information Marketing Information Marketing Start Date Marketing Start Date Marketing En Date D1/06/2021	CF	ROSCARMELLOSE	SODIUM (U	INII: M28OL1HH	48)				
SOJUM CHLORIDE (UNII: 451W47IQ8X) SOJUM STEARYL FUMARATE (UNII: 7CV7W/K4UI) VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 003S90UIF2) Product Characteristics Shape V 0VAL Size 0 9mm 0 0 soor Shape V 0VAL Size 0 9mm 0 0 soor Flavor 0 0VAL 0 0 0 0 0 0 0 soor 0 0 soor Flavor 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	M	ANNITOL (UNII: 30	WL53L36A)						
Monodia Size       no score         Shape       OVAL       Size       9mm         Flavor       Imprint Code       9mm         Contains       Imprint Code       9mm         Packaging       Marketing Start       Marketing En Date         1       NDC:0074-7094- 30       30 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       09/30/2021         2       NDC:0074-7094- 04       30 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       10/06/2021         Marketing Information       Marketing Start Date       Marketing Start Date       Marketing En Date	М	ICROCRYSTALLIN	E CELLULO	SE (UNII: OP1R3	32D61U)				
SODIUM STEARYL FUMARATE (UNII: 7CV7WjK4U)       Image: Sodium Stearyl Fumarate (UNII: 7CV7WjK4U)         VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 003S90U1F2)       Image: Sodium Stearyl Fumarate (UNII: 003S90U1F2)         Product Characteristics       white       Score       no score         Shape       OVAL       Size       9mm         Flavor       Imprint Code       A60         Contains       Imprint Code       Marketing Start Date         #       Item Code       Package Description       Marketing Start Date       Marketing En Date         1       NDC:0074-7094- 30 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       Imprint Code       Imprint Code         2       NDC:0074-7094- 4 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       Imprint Code       Imprint Code         Start Reting Information       Product       Imprint Combination Citation       Marketing Start Date       Marketing En Date									
VITAMIN E POLYETHYLENE GLYCOL SUCCINATE (UNII: 0.03S90U1F2)       Note       Note <th< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></th<>									
Moduct Characteristics         No score         Shape       OVAL       Size       9mm         Flavor       Imprint Code       9mm         Contains         Packaging         #       Item Code       Package Description       Marketing Start Date       Marketing En Date         1       NDC:0074-7094- 30       30 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       09/30/2021         2       NDC:0074-7094- 04       4 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       10/06/2021         Marketing Information         Marketing Start Category       Application Number or Monograph Citation       Marketing Start Date       Marketing En Date									
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Size       9mm         Fiavor       Size       9mm         Fiavor       Imprint Code       Ado         Contains       State       Marketing Statt       Marketing En Date         Mode       Package Description       Marketing Statt       Marketing En Date         Mode       Package Description       09/30/2021       Marketing En Date         Mode       Product       State       State       Marketing En Date         Marketing Category       Application Number or Monograph Citation       Marketing Start Date       Marketing En Date	P	roduct Chara	cteristics	5					
Fiavor       Imprint Code       A60         Fiavor       Imprint Code       A60         Operations         Package Description       Marketing Start Date       Marketing En Date         1       NDC:0074-7094- 30       30 in 1 BOTTLE; Type 0: Not a Combination Product       09/30/2021       Marketing En Date         2       NDC:0074-7094- 04       4 in 1 BOTTLE; Type 0: Not a Combination Product       10/06/2021       Imprint Code         Marketing Category         Marketing Start Date	Co	olor	v	white	Score		no score		
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