

**CENTER FOR DRUG EVALUATION AND
RESEARCH**

APPLICATION NUMBER:

219083Orig1s000

LABELING

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

MYQORZO™ (aficamten) tablets, for oral use
Initial U.S. Approval: 2025

WARNING: RISK OF HEART FAILURE
See full prescribing information for complete boxed warning.

- MYQORZO can cause heart failure due to systolic dysfunction. (5.1)
- Echocardiogram assessments of left ventricular ejection fraction (LVEF) are required before and during MYQORZO use. (2.1)
- Initiation in patients with left ventricular ejection fraction (LVEF) <55% is not recommended. (2.1)
- Decrease dose if LVEF <50% and ≥40%. Interrupt dosing if LVEF <40% or if worsening clinical status. (2.2)
- MYQORZO is available only through a restricted program called the MYQORZO REMS Program. (5.2)

INDICATIONS AND USAGE
MYQORZO is a cardiac myosin inhibitor indicated for the treatment of adults with symptomatic obstructive hypertrophic cardiomyopathy (oHCM) to improve functional capacity and symptoms. (1)

DOSAGE AND ADMINISTRATION

- Recommended starting dose is 5 mg orally once daily. (2.1)
- Dosage is individualized based on echocardiographic assessments and clinical status. Refer to Full Prescribing Information for instructions on dosage modification. (2.1, 2.2)

DOSAGE FORMS AND STRENGTHS
Film-coated tablets: 5 mg, 10 mg, 15 mg, 20 mg. (3)

CONTRAINDICATIONS

- Rifampin (4)

WARNINGS AND PRECAUTIONS
Risk of Heart Failure: Consider decreasing dose or dose interruption in patients with serious intercurrent illness. (2.2, 5.1)
Drug Interactions Leading to Increased Risk of Heart Failure or Loss of Effectiveness: Advise patients of the potential for drug interactions. (2.3, 5.3, 17)

ADVERSE REACTIONS
The most common adverse reaction was hypertension (8%). (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Cytokinetics at 1-833-633-2986 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG INTERACTIONS
Drugs that inhibit multiple pathways of MYQORZO elimination, strong CYP2C9 inhibitors, or moderate-to-strong CYP3A inducers may increase risk of heart failure. MYQORZO dose reduction and additional monitoring may be required when initiating or discontinuing these drugs. (2.3, 7.1)

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved Medication Guide.

Revised: 12/2025

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FULL PRESCRIBING INFORMATION

WARNING: RISK OF HEART FAILURE

MYQORZO reduces left ventricular ejection fraction (LVEF) and can cause heart failure due to systolic dysfunction [see *Warnings and Precautions* (5.1)].

Echocardiogram assessments are required prior to and during treatment with MYQORZO to monitor for systolic dysfunction. Initiation of MYQORZO in patients with LVEF <55% is not recommended. Decrease the dose of MYQORZO if LVEF is <50% and ≥40% [see *Dosage and Administration* (2.2) and *Warnings and Precautions* (5.1)]. Interrupt the dose of MYQORZO if LVEF <40% or if the patient experiences heart failure symptoms or worsening clinical status due to systolic dysfunction [see *Dosage and Administration* (2.2)].

Because of the risk of heart failure due to systolic dysfunction, MYQORZO is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called the MYQORZO REMS Program [see *Warnings and Precautions* (5.2)].

1 INDICATIONS AND USAGE

MYQORZO is indicated for the treatment of adults with symptomatic obstructive hypertrophic cardiomyopathy (oHCM) to improve functional capacity and symptoms.

2 DOSAGE AND ADMINISTRATION

2.1 Evaluation Before and During Use of MYQORZO

Initiation or up-titration of MYQORZO in patients with LVEF <55% is not recommended.

Patients may develop heart failure while taking MYQORZO. Regular LVEF and Valsalva left ventricular outflow tract gradient (LVOT-G) assessment is needed for titration to achieve an appropriate target Valsalva LVOT-G, while maintaining LVEF ≥50% and avoiding heart failure symptoms.

2.2 Recommended Dosage and Administration

The recommended starting dose of MYQORZO is 5 mg orally once daily. Increase the dose every 2 to 8 weeks by 5 mg until a maintenance dose or the maximum recommended dose of 20 mg once daily is achieved. The maintenance dose of MYQORZO is individualized based on the patient's LVEF and LVOT-G. Recommendations for dosing based on LVEF and LVOT-G criteria are provided in [Table 1](#).

Table 1: Dose Adjustment of MYQORZO

LVEF	Valsalva LVOT-G	Dose Adjustment
≥55%	≥30 mmHg	Increase dose by 5 mg (up to the maximum dose of 20 mg once daily)
≥55%	<30 mmHg	Maintain Dose
<55% and ≥50%	Any	Maintain Dose
<50% and ≥40%	Any	Decrease dose by 5 mg ¹ If already on 5 mg, interrupt treatment for at least 7 days
<40%	Any	Interrupt treatment for at least 7 days

1. Decrease dose as follows: 20 mg to 15 mg; 15 mg to 10 mg; 10 mg to 5 mg

Perform an echocardiographic assessment 2 to 8 weeks after initiation of treatment or any dose adjustment (e.g., due to LVEF and LVOT-G criteria or drug interaction). After a treatment interruption due to low LVEF, resume treatment, no earlier than 7 days, when LVEF ≥55% and re-initiate dose titration at the starting dose of 5 mg (see [Table 1](#)).

After the maintenance dose has been established, assess LVEF and Valsalva LVOT-G every 6 months, or every 3 months in patients with LVEF <55% to ≥50%. Consider monitoring LVEF and adjust the dose per [Table 1](#) as needed, in patients with an intercurrent illness (e.g., severe infection or COVID-19), new arrhythmia (e.g., new or uncontrolled atrial fibrillation or other uncontrolled tachyarrhythmia) or any other conditions that may impair systolic function. Do not increase the dose until the intercurrent illness or new arrhythmia has resolved or stabilized.

MYQORZO should be taken once daily with or without meals at about the same time every day. Swallow tablets whole.

2.3 Dosage Modifications for Drug Interactions

Initiate MYQORZO at the recommended starting dose of 5 mg once daily in patients who are on stable therapy with fluconazole, voriconazole, fluvoxamine, strong CYP2C9 inhibitors, or in patients discontinuing a moderate to strong CYP3A inducer.

Concomitant Administration with Fluconazole or Voriconazole

In patients who initiate fluconazole (if used for more than 3 days) or voriconazole, reduce the dose of MYQORZO to 5 mg if they are currently receiving 15 mg or 20 mg. Avoid concomitant use if patients are currently receiving MYQORZO 5 mg or 10 mg. Assess LVEF and LVOT-G 2 to 8 weeks after initiation of fluconazole or voriconazole and titrate the dose of MYQORZO according to [Table 1](#) [see *Dosage and Administration* (2.2), *Warnings and Precautions* (5.3) and *Drug Interactions* (7.1)].

Concomitant Administration with Fluvoxamine or Strong CYP2C9 Inhibitors

In patients who initiate fluvoxamine or a strong CYP2C9 inhibitor, reduce the dose of MYQORZO (20 mg to 10 mg; 15 mg to 5 mg; 10 mg to 5 mg). For patients currently receiving MYQORZO 5 mg, maintain the 5 mg dose. Assess LVEF and LVOT-G 2 to 8 weeks after inhibitor initiation and titrate

the dose of MYQORZO according to [Table 1](#) [see *Dosage and Administration (2.2)*, *Warnings and Precautions (5.3)* and *Drug Interactions (7.1)*].

Concomitant Administration with Moderate to Strong CYP3A Inducers

In patients who are on stable therapy with moderate to strong CYP3A inducers (e.g., carbamazepine), when discontinuing these medications, reduce the dose of MYQORZO (20 mg to 10 mg; 15 mg to 5 mg; 10 mg to 5 mg). For patients currently receiving MYQORZO 5 mg, maintain the 5 mg dose. Assess LVEF and LVOT-G 2 to 8 weeks after inducer discontinuation and titrate the dose of MYQORZO according to [Table 1](#) [see *Dosage and Administration (2.2)*, *Warnings and Precautions (5.3)* and *Drug Interactions (7.1)*].

2.4 Missed Dose

If a dose is missed, it should be taken as soon as possible on the same day. The next scheduled dose should be taken at the usual time the following day. Two doses of MYQORZO should not be taken on the same day.

3 DOSAGE FORMS AND STRENGTHS

MYQORZO is available as tablets in the following strengths:

Tablets: 5 mg - purple, round tablet debossed with “5” on one side and “CK” on the other side.

Tablets: 10 mg - purple, triangular tablet debossed with “10” on one side and “CK” on the other side.

Tablets: 15 mg - purple, pentagonal tablet debossed with “15” on one side and “CK” on the other side.

Tablets: 20 mg - purple, oval tablet debossed with “20” on one side and “CK” on the other side.

4 CONTRAINDICATIONS

MYQORZO is contraindicated with concomitant use of rifampin [see *Warnings and Precautions (5.3)* and *Drug Interactions (7.1)*].

5 WARNINGS AND PRECAUTIONS

5.1 Heart Failure

MYQORZO reduces cardiac contractility, which can reduce LVEF and cause heart failure.

Patients who experience a serious intercurrent illness (e.g., serious infection) or arrhythmia (e.g., new or uncontrolled atrial fibrillation) may be at greater risk of developing systolic dysfunction and heart failure [see *Clinical Trial Experience (6.1)*]. Asymptomatic LVEF reduction, intercurrent illnesses, and arrhythmias require additional monitoring considerations [see *Dosage and Administration (2.2)*].

Assess the patient’s clinical status and LVEF prior to and regularly during treatment and adjust the MYQORZO dose accordingly [see *Dosage and Administration (2.3)*]. New or worsening arrhythmia, dyspnea, chest pain, fatigue, leg edema, or elevations in N-terminal pro-B-type natriuretic peptide (NT-proBNP) may be signs and symptoms of heart failure and should prompt an evaluation of cardiac function.

Initiation of MYQORZO in patients with LVEF <55% is not recommended.

5.2 MYQORZO REMS Program

MYQORZO is available only through a restricted program called the MYQORZO REMS Program, because of the risk of heart failure due to systolic dysfunction [see *Warnings and Precautions (5.1)*].

Notable requirements of the MYQORZO REMS Program include the following:

- Prescribers must be certified by enrolling in the MYQORZO REMS Program.
- Patients must enroll in the MYQORZO REMS Program and comply with ongoing monitoring requirements [see *Dosage and Administration* (2.1)].
- Pharmacies must be certified by enrolling in the MYQORZO REMS Program and must only dispense to patients who are authorized to receive MYQORZO.
- Wholesalers and distributors must only distribute to certified pharmacies.

Further information is available at www.MYQORZOREMS.com or by telephone at 1-844-285-7367.

5.3 Cytochrome P450 Interactions Leading to Heart Failure or Loss of Effectiveness

MYQORZO is metabolized primarily by CYP2C9, and to a lesser extent by CYP3A, CYP2D6, and CYP2C19 enzymes. Initiation of medications that inhibit multiple P450 pathways of MYQORZO elimination (e.g., fluconazole, voriconazole, fluvoxamine) or strong CYP2C9 inhibitors, and discontinuation of moderate-to-strong CYP3A inducers may lead to increased blood concentrations of aficamten and increase the risk of heart failure due to systolic dysfunction [see *Contraindications* (4), *Warnings and Precautions* (5.1), and *Drug Interactions* (7.1)]. Conversely, initiation of medications that induce P450 pathways of MYQORZO (e.g., rifampin, moderate-to-strong CYP3A inducers) may lead to decreased blood concentrations of aficamten and potential loss of effectiveness [see *Contraindications* (4) and *Drug Interactions* (7.1)]. Assess LVEF 2 to 8 weeks after initiation of such inhibitors or after discontinuation of such inducers and adjust the dose of MYQORZO accordingly [see *Dosage and Administration* (2.3)].

Advise patients of the potential for drug interactions. Advise patients to inform their healthcare provider of all concomitant medications prior to and during MYQORZO treatment [see *Drug Interactions* (7.1) and *Patient Counseling Information* (17)].

6 ADVERSE REACTIONS

The following adverse reaction is discussed in other sections of labeling:

- Heart Failure [see *Warnings and Precautions* (5.1)]

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 MYQORZO was evaluated in SEQUOIA-HCM, a phase 3, randomized, double-blind, placebo-controlled study [see *Clinical Studies* (14)]. Of the 282 adults with oHCM, 142 patients received daily doses of MYQORZO (initiated at 5 mg and titrated up to a maximum dose of 20 mg) and 140 patients received placebo. The median treatment duration for patients receiving MYQORZO was ~24 weeks (range 4 to 29 weeks).

Hypertension (8% vs. 2%) was the only adverse reaction occurring in >5% of patients and more commonly on MYQORZO than on placebo.

Eligible oHCM patients were able to participate in an ongoing, open-label, single-arm, long-term safety study (FOREST-HCM). Based on available data, the safety profile of MYQORZO in FOREST-HCM was similar to that observed in SEQUOIA-HCM.

Effects on Systolic Function

In SEQUOIA-HCM, the mean (SD) resting LVEF at baseline was 75% (6) in both treatment groups. Consistent with the mechanism of action of MYQORZO, LS mean (SE) change from baseline in LVEF

was -7% (0.6) in the MYQORZO group and -2% (0.6) in the placebo group at the end of the 24-week treatment period. Four weeks after the end of treatment, mean LVEF was similar between the MYQORZO and placebo groups. During the 24-week treatment period, 5 (4%) patients in the MYQORZO group and 1 (1%) patient in the placebo group experienced a reversible reduction in LVEF to <50% (median LVEF: 47%; range 34% - 49% for these 5 patients in the MYQORZO group). Reductions in LVEF to <50% did not require treatment interruption and were not associated with clinical heart failure [see *Warnings and Precautions* (5.1)].

Effects on Blood Pressure

In SEQUOIA-HCM, the mean (SD) systolic/diastolic blood pressure (SBP/DBP) at baseline was 125(16)/75(11) mmHg for patients in the MYQORZO group and 126(16)/74(11) mmHg in the placebo group. There was a greater mean change from baseline in SBP/DBP (SD) in the MYQORZO group compared to the placebo group at the end of the 24-week treatment period [2(13)/3(8) mmHg and -3(14)/-1(9) mmHg, respectively]. Four weeks after the end of treatment, the mean SBP/DBP was similar between the MYQORZO and placebo groups. In SEQUOIA-HCM, SBP ≥160 mmHg was observed in approximately 16% of patients in the MYQORZO group and 8% of patients in the placebo group. MYQORZO-associated increases in blood pressure are consistent with relief of LVOT obstruction and improved cardiac output.

7 DRUG INTERACTIONS

7.1. Potential for Other Drugs to Affect Plasma Concentrations of MYQORZO

Aficamten is primarily metabolized by CYP2C9 and, to a lesser extent by CYP3A, CYP2D6, and CYP2C19. Concomitant administration of drugs that inhibit multiple P450 pathways of aficamten elimination, strong inhibitors of CYP2C9, and moderate-to-strong inducers of CYP3A, may affect the exposure of aficamten [see *Warnings and Precautions* (5.2) and *Clinical Pharmacology* (12.3)] (see Table 2).

Table 2: Established and Potentially Significant Pharmacokinetic Drug-Drug Interactions with MYQORZO

Inhibitors of Multiple CYPs		
Fluvoxamine	Clinical Impact	Fluvoxamine is a strong CYP2C19 inhibitor, weak to moderate CYP3A inhibitor, weak CYP2C9 inhibitor, and weak CYP2D6 inhibitor. Coadministration of fluvoxamine is predicted to increase aficamten exposure, which may increase the risk of developing heart failure due to systolic dysfunction.
	Prevention or Management	Initiate MYQORZO at 5 mg in patients on stable therapy with fluvoxamine. In patients who are on MYQORZO treatment and intend to initiate fluvoxamine: Reduce dose of MYQORZO (i.e., 20 mg to 10 mg, 15 mg to 5 mg, or 10 mg to 5 mg) or continue 5 mg [see <i>Dosage and Administration</i> (2.3)].

Fluconazole or Voriconazole	Clinical Impact	Fluconazole is a moderate CYP2C9 inhibitor, moderate CYP3A inhibitor, and strong CYP2C19 inhibitor. Voriconazole is a strong CYP3A inhibitor, moderate CYP2C19 inhibitor, and weak CYP2C9 inhibitor. Coadministration with fluconazole is observed, and voriconazole is predicted, to increase aficamten exposure, which may increase the risk of developing heart failure due to systolic dysfunction [see <i>Clinical Pharmacology</i> (12.3)].
	Prevention or Management	Initiate MYQORZO at 5 mg in patients on stable therapy with fluconazole or voriconazole. In patients currently receiving MYQORZO who intend to initiate fluconazole (if used more than 3 days) or voriconazole: Reduce dose of MYQORZO to 5 mg if they are currently receiving MYQORZO 15 mg or 20 mg. Avoid concomitant use if currently receiving MYQORZO 5 mg or 10 mg [see <i>Dosage and Administration</i> (2.3)].
Strong CYP2C9 inhibitors		
Clinical Impact		Coadministration with a strong CYP2C9 inhibitor is predicted to increase the exposure of aficamten, which may increase the risk of developing heart failure due to systolic dysfunction [see <i>Clinical Pharmacology</i> (12.3)].
Prevention or Management		Initiate MYQORZO at 5 mg in patients who are on stable therapy with a strong CYP2C9 inhibitor. In patients who are on MYQORZO treatment and intend to initiate a strong CYP2C9 inhibitor: Reduce dose of MYQORZO (i.e., 20 mg to 10 mg, 15 mg to 5 mg, or 10 mg to 5 mg) or continue 5 mg [see <i>Dosage and administration</i> (2.3)].
CYP Inducers		
Rifampin	Clinical Impact	Rifampin is a strong CYP3A inducer, strong CYP2C19 inducer, and moderate CYP2C9 inducer. Coadministration of rifampin decreases aficamten exposure, which may reduce the effectiveness of MYQORZO [see <i>Contraindications</i> (4) and <i>Clinical Pharmacology</i> (12.3)].
	Prevention of Management	Concomitant use with rifampin is contraindicated.
Other moderate-to-strong CYP3A inducers	Clinical Impact	Concomitant use with a moderate to strong CYP3A inducer decreases aficamten exposure, which may reduce MYQORZO effectiveness [see <i>Clinical Pharmacology</i> (12.3)]. The risk of heart failure due to systolic dysfunction may increase with discontinuation of these inducers.

	Prevention or Management	In patients who intend to discontinue a moderate to strong CYP3A Inducer: Reduce dose of MYQORZO (20 mg to 10 mg; 15 mg to 5 mg; 10 mg to 5 mg) or continue 5 mg [see <i>Dosage and Administration</i> (2.3)].
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8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

There are no available data on the use of MYQORZO during pregnancy to evaluate for a drug-associated risk of major birth defects, miscarriage, or other adverse maternal or fetal outcomes. The underlying maternal condition during pregnancy poses a risk to the mother and fetus (see *Clinical Considerations*). In embryo-fetal and pre- and postnatal development studies, when pregnant rats were administered aficamten during the period of organogenesis, aficamten increased the incidence of structural malformations at exposures ≥ 4 -times the maximum recommended human dose (MRHD) of 20 mg based on free area under the concentration curve (AUC). No adverse effects on development were seen at 3-times the MRHD exposure (see *Data*).

The background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defects, loss, and other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively.

There is a pregnancy safety study for MYQORZO. If MYQORZO is administered during pregnancy, or if a patient becomes pregnant while receiving MYQORZO or within 3 weeks after the last dose of MYQORZO, healthcare providers should report MYQORZO exposure by contacting Cytokinetics, Inc. at 1-833-633-2986.

Clinical Considerations

Disease-Associated Maternal and Embryo-Fetal Risk

Obstructive HCM in pregnancy has been associated with increased risk for preterm birth.

Data

Animal Data

Aficamten given to pregnant rats (2, 6 and 9 mg/kg/day) during the period of organogenesis was associated with increased external fetal malformations (kinked tail) at aficamten exposures 5-times the clinical exposures at the MRHD based on free AUC, with uncertain human relevance. Increased post-implantation loss (early and late resorptions) and decreased mean fetal body weight occurred in the presence of significant maternal toxicity at exposures 5-times the clinical exposures at the MRHD. No adverse effects on embryofetal development were observed at 6 mg/kg/day, representing 3-times the clinical exposure at the MRHD.

Pregnant rabbits given aficamten (5, 10 and 20/15 mg/kg/day) during the period of embryofetal organogenesis showed a numerical increase in post-implantation loss (late resorptions) and fetuses with cardiac malformations at 10 mg/kg/day. However, no increase in malformations was observed at the high dose of 20/15 mg/kg/day, a dose that caused significant maternal toxicity (leading to dose reduction from 20 to 15 mg/kg/day during the study) at clinically relevant exposures.

Pregnant rats given aficamten (0.5, 1.5, 6 mg/kg/day) during embryofetal organogenesis through offspring weaning in a pre- and postnatal development study had reduced offspring viability at the 6 mg/kg/day in the presence of maternal toxicity, at 4-times the clinical exposure at the MRHD based

on free AUC. An increased number of offspring with bent tails was also observed at the 6 mg/kg/day. No adverse maternal toxicity or developmental effects occurred at 1.5 mg/kg/day, which corresponds to clinical exposure at the MRHD. Aficamten did not affect neurobehavioral parameters, sexual maturation, or reproductive function of the offspring at any dose.

8.2 Lactation

Risk Summary

There are no data on the presence of aficamten in human milk, the effects on the breastfed infant, or the effects on milk production. Aficamten was detected in the plasma of rat pups when dams were dosed with the drug orally during the lactation period. When a drug is present in animal milk, it is likely that the drug will be present in human milk. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for MYQORZO and any potential adverse effects on the breastfed infant from MYQORZO or from the underlying maternal condition.

8.4 Pediatric Use

The safety and effectiveness of MYQORZO in pediatric patients have not been established.

8.5 Geriatric Use

There were 57 (40%) MYQORZO-treated patients, aged 65 years and older, in SEQUOIA-HCM [see *Clinical Studies* (14)]. Of the total number of MYQORZO-treated patients in SEQUOIA-HCM, 45 (32%) patients were 65 to 74 years of age, and 12 (8%) patients were 75 years of age and older.

No overall differences in safety or effectiveness of MYQORZO have been observed between patients 65 years of age and older and younger adult patients.

10 OVERDOSAGE

Clinical Experience and Effects

There have been no reports of overdose with MYQORZO. Cardiovascular effects of overdose may include reduced LVEF, heart failure, and hypotension. Signs and symptoms of heart failure such as dyspnea, fatigue, leg edema, or elevations in NT-proBNP should prompt an evaluation of cardiac function [see *Warnings and Precautions* (5.1)].

Management

Discontinue MYQORZO treatment. Provide medically supportive measures to maintain hemodynamic stability and monitor left ventricular function.

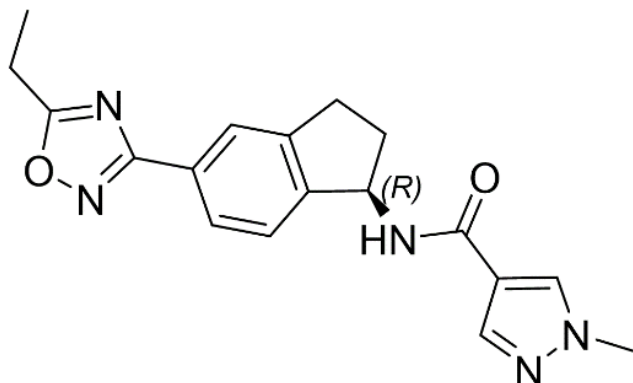
Consider contacting the Poison Help line (1-800-222-1222) or a medical toxicologist for additional overdose management recommendations.

11 DESCRIPTION

MYQORZO tablets for oral use contain aficamten, a cardiac myosin inhibitor.

The chemical name of aficamten is (R)-N-(5-(5-ethyl-1,2,4-oxadiazol-3-yl)-2,3-dihydro-1H-inden-1-yl)-1-methyl-1H-pyrazole-4-carboxamide. The molecular formula is C₁₈H₁₉N₅O₂ and the molecular weight is 337.38 g/mol.

The structural formula of aficamten is:



Aficamten is a white to off-white to yellow to grey solid that is practically insoluble in water and aqueous buffers (pH 2 – 9), sparingly soluble in acetone and ethanol, and soluble in N-methylpyrrolidone (NMP).

MYQORZO is supplied as tablets containing 5 mg, 10 mg, 15 mg, or 20 mg of aficamten per tablet as the active ingredient and the following inactive ingredients: croscarmellose sodium, hydroxypropyl cellulose, magnesium stearate (non-bovine), mannitol, microcrystalline cellulose, and sodium lauryl sulfate. The tablet coating contains carmine, FD&C Blue No. 2, glyceryl mono and dicaprylate, polyvinyl alcohol, polyvinyl alcohol graft polyethylene copolymer, talc and titanium dioxide.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Aficamten is an allosteric and reversible inhibitor of cardiac myosin motor activity. Aficamten reduces the force generated by myosin at the cardiac sarcomere, which contributes to the pathophysiology of HCM. In patients with HCM, myosin inhibition with aficamten reduces cardiac contractility and left ventricular outflow tract (LVOT) obstruction.

12.2 Pharmacodynamics

Left Ventricular Ejection Fraction and Left Ventricular Outflow Tract Obstruction

In SEQUOIA-HCM, reductions in LVOT-G were observed at 2 weeks after initiating treatment, and LVOT-G continued to decrease through Week 8. The reduction in LVOT-G was sustained through the remainder of the 24-week trial. At Week 24, the mean (SD) change from baseline in resting and Valsalva LVOT-G was -35 (29) mmHg and -48 (37) mmHg, respectively, for the MYQORZO group and 4 (35) mmHg and 2 (35) mmHg, respectively, for the placebo group. The reduction in LVOT-G was accompanied by a greater decrease in LVEF in the MYQORZO group compared to the placebo group with mean (SD) change from baseline in LVEF of -7 (7)% and -2 (6)%, respectively. Four weeks after discontinuation of treatment, LVEF and Valsalva LVOT-G returned to baseline.

Cardiac Structure

In SEQUOIA-HCM, maximal left ventricular wall thickness, left ventricular mass index (LVMI) and left atrial volume index (LAVI) were measured at baseline and Week 24. At Week 24, the mean (SD) maximal left ventricular wall thickness decreased in the MYQORZO group by -0.2 (0.3) cm and in the placebo group by -0.1 (0.3) cm. LVMI decreased in the MYQORZO group by -5.3 (24) g/m² and increased in the placebo group by 5.8 (25) g/m². At Week 24, LAVI decreased in the MYQORZO group by -2.6 (8) mL/m² and increased in the placebo group by 1.4 (8) mL/m². The clinical significance of these findings is unknown.

Cardiac Biomarkers

In SEQUOIA-HCM, reductions in NT-proBNP were observed in the MYQORZO group starting at Week 2 and were sustained through Week 24. At Week 24, NT-proBNP was 80% lower with MYQORZO compared to placebo (proportion of geometric mean ratio between MYQORZO and placebo, 0.20 [95% CI: 0.17, 0.23]). At both Week 12 and 24, troponin I was 40% lower with MYQORZO compared to placebo (proportion of geometric mean ratio between the two groups, 0.57 [95% CI: 0.51, 0.64]).

The clinical significance of the NT-proBNP and troponin findings is unknown.

Cardiac Electrophysiology

The results of a thorough QT study demonstrated a lack of QTc prolongation across the therapeutic concentration range of MYQORZO. At a single dose of 50 mg (similar exposure to 20 mg daily dosing to steady-state), the upper limit of the predicted placebo-corrected change from baseline in QT interval corrected by Fridericia ($\Delta\Delta\text{QTcF}$) 90% confidence interval for MYQORZO was <10 msec.

12.3 Pharmacokinetics

Aficamten exposure increases dose proportionally following single doses of 1 mg to 75 mg and multiple once daily doses of 5 mg to 20 mg. Aficamten pharmacokinetics was comparable between healthy subjects and patients with oHCM. Geometric mean accumulation ratios for aficamten were similar across dose levels and ranged from 4.6 to 4.8. Steady state is predicted to be reached following approximately 17 days of once daily dosing of MYQORZO in patients with oHCM.

Absorption

Aficamten is rapidly absorbed with a median time to maximum concentration (T_{max}) of 1.5 to 2.0 hours. The bioavailability of aficamten after oral administration is unknown.

Effect of Food

No clinically significant differences in aficamten AUC and C_{max} were observed following its administration with a high-fat, high-calorie meal.

Distribution

Aficamten is approximately 90% bound to plasma proteins and demonstrates a volume of distribution of 313 L. The blood to plasma ratio of aficamten was 0.94.

Elimination

Aficamten has a median terminal half-life ($t_{1/2}$) of approximately 80 hours in patients with oHCM. At steady-state, the peak-to-trough plasma concentration ratio with once daily dosing is approximately 1.2. The total clearance is 2.6 L/hr and the renal clearance is <0.1% of total clearance.

Metabolism

Aficamten is extensively metabolized in humans, primarily through CYP2C9 with contributions by CYP3A, CYP2D6, and CYP2C19. Aficamten is primarily metabolized to two pharmacologically inactive metabolites, CK-3834282 and CK-3834283, that circulate at approximately 56% and 103% of parent in plasma, respectively.

Excretion

Following a single 20 mg radiolabeled dose of aficamten, 32% (0.06% unchanged aficamten) was excreted in urine and 58% (5.1% unchanged aficamten) was excreted in feces.

Specific Populations

No clinically meaningful differences in the pharmacokinetics of aficamten were observed based on age (18 – 83 years), race, mild to moderate hepatic impairment (Child-Pugh Classes A and B), or mild to moderate renal impairment (estimated glomerular filtration rate (eGFR) ≥ 30 mL/min).

normalized by body surface area and based on the modification of diet in renal disease (MDRD) equation). The effects of severe renal (eGFR <30 mL/min based on MDRD equation) or hepatic (Child-Pugh Class C) impairment are unknown.

Small differences in aficamten exposure with sex and body weight were observed. Female patients demonstrated a 31% higher mean exposure (AUC) compared to male patients. The highest body weight quartile of patients (median: 105 kg) demonstrated a 34% lower aficamten mean exposure (AUC) than the lowest body weight quartile of patients (median: 64 kg). These differences are not clinically significant.

Drug Interaction Studies

Clinical Studies and Model-Informed Approaches

Inhibitors and Inducers of CYP enzymes: The predicted and observed effect of coadministration of select CYP inhibitors and inducers on the plasma exposure of aficamten is provided in [Table 3](#).

Table 3: In Vivo Observed and Predicted Effects of CYP Inhibitors and Inducers on Aficamten Plasma Exposure

Concomitant Medication	Dosage	Effect on Aficamten Mean AUC
Inhibitors of Individual CYP Enzymes		
Sulfaphenazole ¹ - <i>Strong CYP2C9 Inhibitor</i>	2000 mg once daily	AUC: ↑ 113%
Itraconazole ² - <i>Strong CYP3A Inhibitor</i>	200 mg once daily	AUC: ↑ 26%
Paroxetine ² – <i>Strong CYP2D6 Inhibitor</i>	40 mg once daily	AUC: ↑ 27%
Inhibitors of Multiple CYP Enzymes		
Fluconazole ² - <i>Strong CYP2C19, Moderate CYP3A, and Moderate CYP2C9 Inhibitor</i>	400 mg once daily	AUC: ↑ 278%
	150 mg x 2 doses; separated by 48 hours	AUC: No change
Fluvoxamine ¹ - <i>Strong CYP2C19, Weak-to-Moderate CYP3A, Weak CYP2C9, and Weak CYP2D6 Inhibitor</i>	150 mg twice daily	AUC: ↑ 116%
Voriconazole ¹ - <i>Strong CYP3A, Moderate CYP2C19, and Weak CYP2C9 inhibitor</i>	200 mg twice daily	AUC: ↑ 94%
Fluoxetine ² - <i>Strong CYP2D6 and Strong CYP2C19 Inhibitor</i>	40 mg once daily	AUC: ↑ 32%
CYP Enzyme Inducers		
Rifampin ¹ - <i>Strong CYP3A, Strong CYP2C19, and Moderate CYP2C9 Inducer</i>	600 mg once daily	AUC: ↓ 79%
Carbamazepine ² – <i>Moderate-to-strong CYP3A and Weak CYP2C9 Inducer</i>	300 mg twice daily	AUC: ↓ 51%

1. Predicted via physiologically-based pharmacokinetic modeling.

2. Observed.

P-gp Transporter Substrates: Concomitant administration of a single 20 mg dose of aficamten increased dabigatran (administered as dabigatran etexilate) mean AUC_{inf} by 26%.

In Vitro Drug Interaction Studies

CYP enzymes: Aficamten is not expected to inhibit CYP1A2, CYP2B6, CYP2C8, CYP2C19, CYP2D6, or CYP3A4 at clinically relevant concentrations. Aficamten is not expected to induce CYP1A2, CYP2B6 or CYP3A4 at clinically relevant concentrations.

Transporter systems: Aficamten is not a sensitive substrate of drug transporters. Aficamten is not expected to inhibit BCRP, OCT2, OCT3, OAT1, OAT3, OATP1B1, OATP1B3, MATE1 or MATE2-K drug transporters at clinically relevant concentrations.

12.5 Pharmacogenomics

Aficamten is primarily metabolized by the polymorphic enzyme, CYP2C9, with lesser contributions by other polymorphic enzymes (e.g., CYP2D6 and CYP2C19) [see *Clinical Pharmacology* (12.3)]. The impact of CYP2C9 genetic variants on the pharmacokinetics of aficamten has not been directly evaluated, though, a similar effect on exposure is expected as seen with strong CYP2C9 inhibitors [see *Drug Interactions* (7.1)].

No additional dosage modifications are required for patients who are CYP2C9 poor metabolizers, as MYQORZO dosage titration and modifications account for CYP2C9 metabolizer status [see *Dosage and Administration* (2.2) and (2.3)].

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis

The carcinogenic potential of aficamten was assessed in a 26-week study in transgenic rasH2 mice and a 2-year study in Sprague Dawley rats. There was no evidence of carcinogenicity with oral administration of aficamten at dose levels up to 2.0 mg/kg/day in the mouse or 3 mg/kg/day in the rat (corresponding to 4-fold the MRHD based on free AUC).

Mutagenesis

Aficamten was not genotoxic in the in vitro bacterial (Ames) mutagenicity assay or the combined in vivo micronucleus assay and comet assay in rat.

Impairment of Fertility

In a fertility and early embryonic development toxicology study in the rat, aficamten was administered at doses up to 6 mg/kg/day in females and 3 mg/kg/day in males, with no drug-related effects on mating, fertility, estrous cycling, male reproductive assessments, or early embryonic survival.

14 CLINICAL STUDIES

The efficacy of MYQORZO was evaluated in SEQUOIA-HCM (NCT05186818), a phase 3, multicenter, randomized, double-blind, placebo-controlled study in 282 adults (142 aficamten, 140 placebo) with symptomatic New York Heart Association (NYHA) class II and III oHCM, LVEF $\geq 60\%$, and resting and post-Valsalva peak LVOT-G ≥ 30 and ≥ 50 mmHg at screening, respectively. Patients who completed SEQUOIA-HCM were eligible to participate in an ongoing, open-label, single-arm extension study (FOREST-HCM). Patients with a known infiltrative or storage disorder causing cardiac hypertrophy such as Noonan syndrome, Fabry disease or amyloidosis were excluded.

Patients were randomized in a 1:1 ratio to receive either MYQORZO or placebo once daily for 24 weeks. Randomization was stratified by use of beta-blockers (yes or no) and cardiopulmonary exercise testing (CPET) exercise modality (treadmill or bicycle)

At baseline, 76% of the randomized patients were NYHA class II and 24% were NYHA class III. The mean peak oxygen uptake (pVO₂) by CPET at baseline was 18.5 mL/kg/min with 55% using treadmill and 45% using bicycle, respectively. At baseline, the median LVEF was 76%, the mean resting LVOT-G was 55 mmHg, the mean Valsalva LVOTG was 83 mmHg, and mean Kansas City Cardiomyopathy Questionnaire – Clinical Summary Score (KCCQ-CSS) was 74. At baseline, 61% of patients were on beta-blockers, 29% were on non-dihydropyridine calcium channel blockers, 13% were on disopyramide, and 15% were not taking any background medication for oHCM.

Groups were balanced with respect to age (mean 59 years; range 18 to 84 years), sex (59% male), race (79% White, 19% Asian and 1% Black or African American), body mass index (mean 28 kg/m²), heart rate (mean 66 bpm) and blood pressure (mean 124/74 mmHg).

Patients were initiated on MYQORZO at a dose of 5 mg once daily. Doses were individually titrated (or sham-titrated if on placebo) at Week 2, 4 and 6 if Valsalva LVOT-G was ≥30 mmHg and LVEF was ≥55% in 5 mg dose increments up to a maximum dose of 20 mg once daily. At Week 24, in the MYQORZO group, 46% of patients were on the 20 mg dose, 35% were on the 15 mg dose, 15% were on the 10 mg dose and 4% were on the 5 mg dose.

Primary Endpoint - Peak Oxygen Uptake (pVO₂)

In SEQUOIA-HCM, the primary endpoint of change from baseline in pVO₂ to Week 24 was greater with MYQORZO compared to placebo, as shown in [Table 4](#).

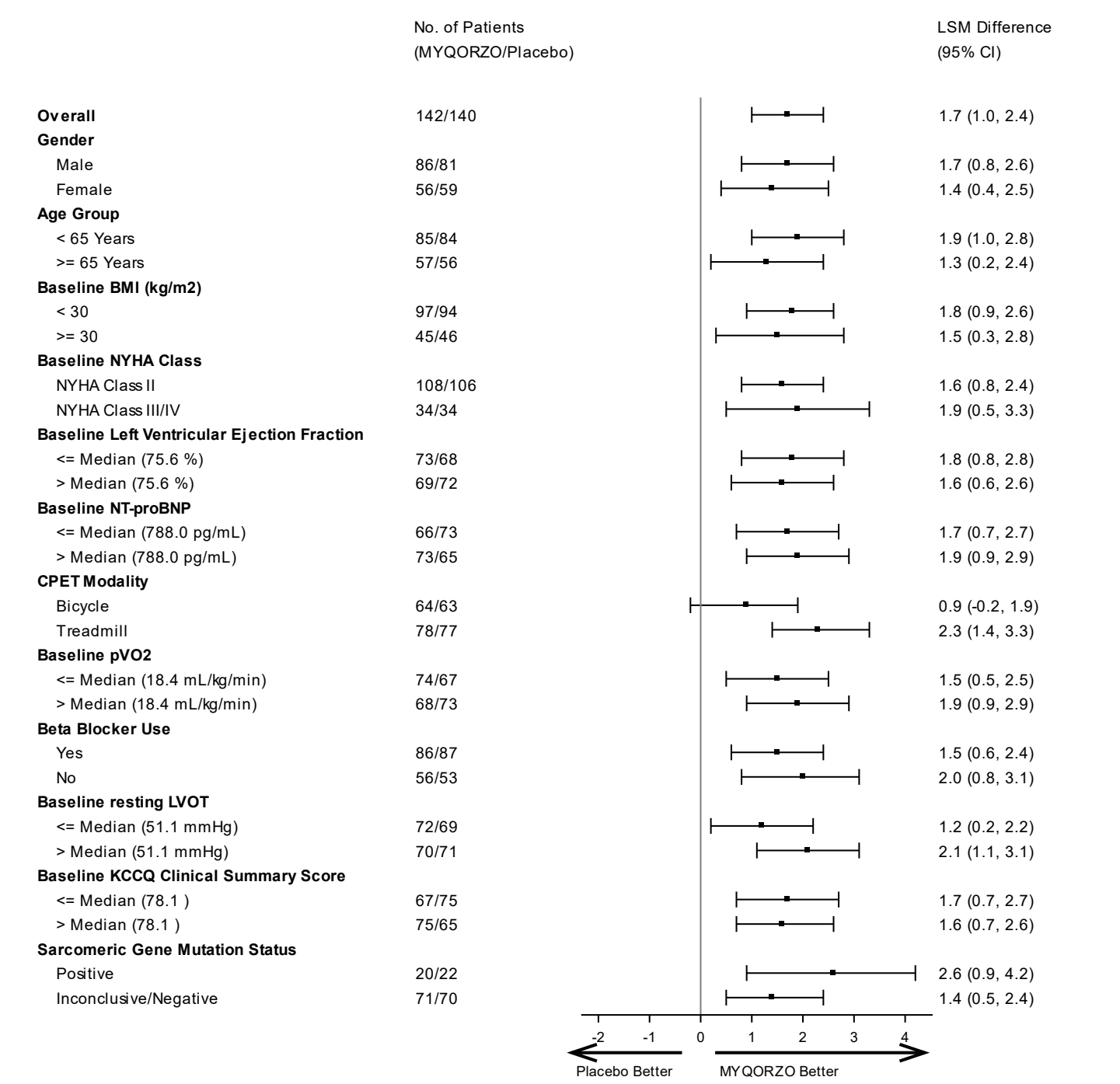
Table 4: Change from Baseline in pVO₂ by CPET at Week 24 in SEQUOIA-HCM

pVO ₂	MYQORZO (N = 142)	Placebo (N = 140)
Baseline (mL/min/kg), mean (SD)	18.4 (4.4)	18.6 (4.5)
Change from baseline to Week 24		
LS mean (SE) ¹	1.7 (0.3)	0.0 (0.3)
LS mean difference vs. placebo (95% CI) ¹	1.7 (1.0, 2.4)	
p-value	<0.0001	

1. LS mean (SE), and LS mean difference (95% CI) were estimated from an ANCOVA model. A placebo-based multiple imputation approach was used to impute the missing data.

A range of demographic characteristics, baseline disease characteristics, and baseline concomitant medications (e.g., use of beta-blockers) were examined for their influence on outcomes. Results of the primary endpoint analysis consistently favored MYQORZO across all subgroups analyzed ([Figure 1](#)).

Figure 1: Forest Plot for Change in Baseline pVO₂ by CPET at Week 24 by Subgroups in SEQUOIA-HCM



BMI: Body Mass Index; NYHA: New York Heart Association; NT-proBNP: N-terminal pro-B-type natriuretic peptic; CPET: Cardiac Pulmonary Exercise Test; pVO₂: Peak Oxygen Uptake; LVOT: left ventricular outflow tract; KCCQ CSS: Kansas City Cardiomyopathy Questionnaire – Clinical Summary Score.

Note: The figure above presents effects in various subgroups, all of which are baseline characteristics. The 95% confidence limits that are shown do not take into account the number of comparisons made and may not reflect the effect of a particular factor after adjustment for all other factors. Apparent homogeneity or heterogeneity among groups should not be over-interprete

Secondary Endpoints

The treatment effects of MYQORZO on health status, functional capacity, and LVOT obstruction were assessed by change in the KCCQ-CSS, proportion of patients with ≥ 1 class improvement in NYHA functional class, change from baseline in Valsalva LVOT-G, proportion of patients with Valsalva LVOT-G ≤ 30 mmHg, duration of eligibility for septal reduction therapy (SRT), and change in total workload during CPET. At Week 24, patients receiving MYQORZO had greater improvement compared to the placebo group across all secondary points (Table 5, Figure 2 and Figure 3).

Table 5. Results of Secondary Efficacy Endpoints in SEQUOIA-HCM

Endpoints	MYQORZO N=142	Placebo N=140	LS Mean Difference (95% CI)	p-Value
Mean Change from Baseline in KCCQ-CSS^{1,3}				
Week 12	11.1 (0.9)	4.0 (0.9)	7.0 (4.6, 9.5)	<0.0001
Week 24	11.6 (1.0)	4.3 (1.0)	7.3 (4.6, 10.1)	<0.0001
Mean Change from Baseline in Valsalva LVOT-G (mmHg)¹				
Week 12	-45.0 (2.4)	2.7 (2.4)	-47.7 (-54.5, -41.0)	<0.0001
Week 24	-46.7 (2.5)	2.2 (2.5)	-48.8 (-55.7, -42.0)	<0.0001
Mean Duration of SRT Eligibility¹				
Days spent SRT-eligible during 24 Weeks of Treatment	35 (8)	113 (8)	-78 (-100, -56)	<0.0001
Mean Change from Baseline in Total Workload (Watts) during CPET¹				
Week 24	13.1 (2.1)	0.9 (2.2)	12.2 (6.3, 18.1)	<0.0001
Endpoints	MYQORZO N=142	Placebo N=140	Odds Ratio (95% CI)	p-Value
Proportion of Patients with ≥ 1 NYHA Class Improvement²				
Week 12	69 (48.6)	25 (17.9)	OR: 4.6 (2.6, 8.4)	<0.0001
Week 24	83 (58.5)	34 (24.3)	OR: 4.4 (2.6, 7.6)	<0.0001
Proportion of Patients with Valsalva LVOT-G ≤ 30 (mmHg)²				
Week 12	73 (51.4)	8 (5.7)	OR: 16.9 (7.7, 37.2)	<0.0001
Week 24	70 (49.3)	5 (3.6)	OR: 22.9 (8.8, 59.6)	<0.0001

CPET: Cardiac Pulmonary Exercise Test; KCCQ CSS: Kansas City Cardiomyopathy Questionnaire – Clinical Summary Score; NYHA: New York Heart Association; LVOT-G: left ventricular outflow tract gradient; SRT: septal reduction therapy.

1. LS means (SE) and LS mean difference (95% CI) was estimated from an ANCOVA model (mixed model with repeated measures was used for longitudinal data) for continuous endpoints. Placebo-based multiple imputation approach was used to impute the missing data.
2. The number (percentage) of responders and common Odds Ratio (OR) obtained using Cochran-Mantel-Haenszel method (95% CI) are presented for binary endpoints.
3. The KCCQ-CSS score ranges from 0 to 100, with higher scores indicating better health status.

Figure 2: Change from Baseline in KCCQ-CSS through Week 24

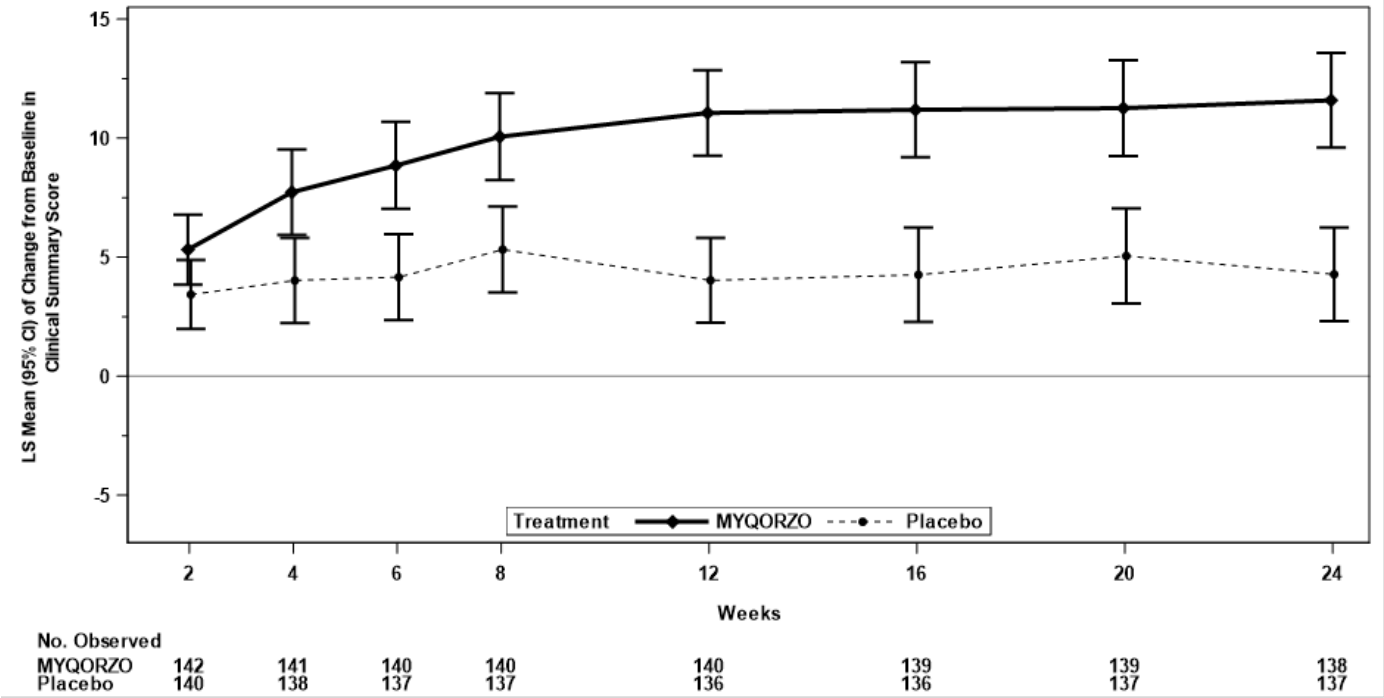
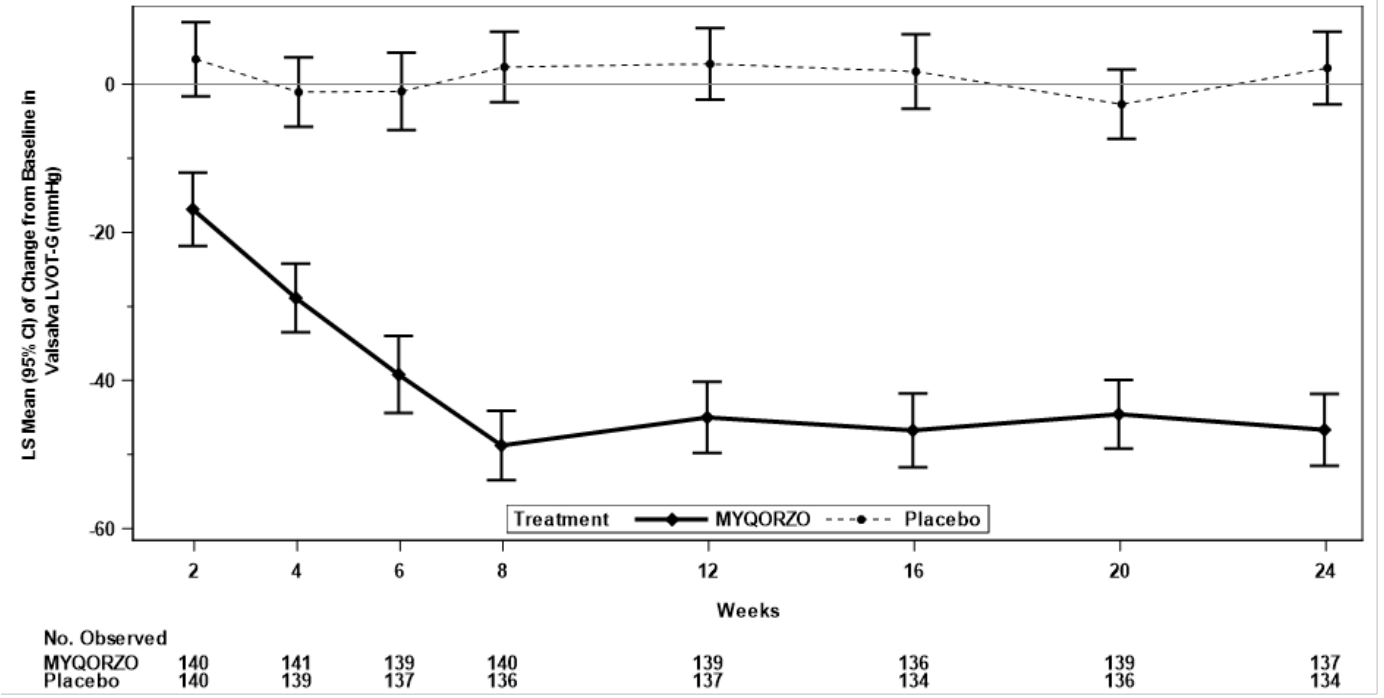


Figure 3: Change from Baseline in Valsalva LVOT-G through Week 24



16 HOW SUPPLIED/STORAGE AND HANDLING

MYQORZO is supplied as purple, film-coated tablets containing 5 mg, 10 mg, 15 mg, or 20 mg aficamten. The tablets are debossed on one side with “CK” and the other side with “5”, “10”, “15” and “20” for the 5, 10, 15, and 20 mg strength, respectively.

Tablets are supplied in bottles with child-resistant closure as follows:

5 mg round tablets	Bottles of 30 tablets	NDC: 82112-105-30
10 mg triangular tablets	Bottles of 30 tablets	NDC: 82112-110-30
15 mg pentagonal tablets	Bottles of 30 tablets	NDC: 82112-115-30
20 mg oval tablets	Bottles of 30 tablets	NDC: 82112-120-30

Storage

Store at 20°C to 25°C (68°F to 77°F); excursions permitted between 15°C to 30°C (59°F to 86°F) [see *USP controlled Room Temperature*].

17 PATIENT COUNSELING INFORMATION

Advise the patient and/or caregiver to read the FDA-approved patient labeling (Medication Guide).

Heart Failure

Inform patients that cardiac function monitoring using echocardiography must be performed before and while on treatment to monitor for signs and symptoms of heart failure [see *Dosage and Administration* (2.1), *Warnings and Precautions* (5.1)]. Advise patients to report any signs or symptoms of heart failure immediately to their healthcare provider.

MYQORZO REMS Program

MYQORZO is available only through a restricted program called the MYQORZO REMS Program [see *Warnings and Precautions* (5.2)]. Inform the patient of the following notable requirements:

Patients must enroll in the program and comply with ongoing monitoring requirements [see *Warnings and Precautions* (5.1) and (5.2)]

MYQORZO is only prescribed by certified healthcare providers and only dispensed from certified pharmacies participating in the program. Provide patients with the telephone number and website for information on how to obtain the product [see *Warnings and Precautions* (5.2)].

Pregnancy

Advise patients that there is a pregnancy safety study for MYQORZO. Advise patients to report pregnancy to their prescriber if they become pregnant while receiving MYQORZO or within 3 weeks after their last dose of MYQORZO [see *Use in Specific Populations* (8.1)].

Drug Interactions

Advise patients to inform their healthcare provider of all concomitant medications they take, prior to and during MYQORZO treatment.

Instructions for Taking MYQORZO

MYQORZO tablets should be swallowed whole. Advise patients that if they miss a dose of MYQORZO, it should be taken as soon as possible on the same day. The next scheduled dose should be taken at the usual time the following day. The patient should not take two doses on the same day.

Distributed By:

Cytokinetics Incorpo

South San Francisco, CA 94080

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Patent Information: <https://cytokinetics.com/patents>

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MEDICATION GUIDE
MYQORZO (my-kor-zo)
(aficamten)
tablets, for oral use

What is the most important information I should know about MYQORZO?

MYQORZO can cause serious side effects, including:

- **Heart failure**, a condition where the heart cannot pump with enough force. Heart failure is a serious condition that can lead to death. You must have echocardiograms before and during treatment with MYQORZO and monitor for signs and symptoms of heart failure. People who develop a serious illness such as a serious infection or who develop a new or worsening irregular heartbeat have a greater risk of heart failure during treatment with MYQORZO.
Tell your healthcare provider or get medical help right away if you develop new or worsening:
 - shortness of breath
 - swelling in your legs
 - a racing sensation in your heart (palpitations)
 - fatigue
 - chest pain
 - rapid weight gain
- **The risk of heart failure is also increased when MYQORZO is taken with certain other medicines.** Tell your healthcare provider about the medicines you take, both prescribed and obtained over-the-counter, before and during your treatment with MYQORZO.
- **Because of the risk of heart failure, MYQORZO is only available through a restricted distribution program called the MYQORZO Risk Evaluation and Mitigation Strategy (REMS) Program.**
 - Your healthcare provider must be enrolled in the MYQORZO REMS Program for you to be prescribed MYQORZO.
 - Before you start treatment with MYQORZO, you must enroll in the MYQORZO REMS Program. Talk to your healthcare provider about how to enroll in the program. You will be given information about the program when you enroll.
 - Before you take MYQORZO, your healthcare provider and pharmacist will make sure you understand how to take MYQORZO safely, which will include returning for echocardiograms when advised by your healthcare provider. MYQORZO can only be dispensed by a certified pharmacy that participates in the MYQORZO REMS Program. Your healthcare provider can give you information on how to find a certified pharmacy.
 - If you have any questions about the MYQORZO REMS Program, ask your healthcare provider, go to www.MYQORZOREMS.com, or call 1-844-285-7367.
- See **“What are the possible side effects of MYQORZO?”** for information about side effects.

What is MYQORZO?

MYQORZO is a prescription medicine used to treat adults with symptomatic obstructive cardiomyopathy (oHCM) to improve functional capacity and symptoms.

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

Who should not take MYQORZO?

Do not take MYQORZO if you take a medicine called rifampin.

Before taking MYQORZO, tell your healthcare provider about all of your medical conditions, including if you:

- are pregnant or plan to become pregnant. It is not known if MYQORZO can cause harm to your unborn baby. Tell your healthcare provider if you become pregnant during treatment or within 3 weeks after the last dose of MYQORZO. There is a pregnancy safety study for MYQORZO. Your healthcare provider should report your pregnancy exposure to Cytokinetics, Inc.
- are breastfeeding or plan to breastfeed. It is not known if MYQORZO passes into your breast milk. Talk to your healthcare provider about the best way to feed your baby during treatment with MYQORZO.

Before and during MYQORZO treatment, tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Taking MYQORZO with certain medicines may lead to increased levels of MYQORZO in your blood and increase your risk of heart failure. **Do not stop or change the dose of a medicine or start a new medicine without telling your healthcare provider.**

Especially tell your healthcare provider if you take the medicines:

- fluconazole (if used for more than 3 days)
- voriconazole
- fluvoxamine

How should I take MYQORZO?

- Take MYQORZO exactly as your healthcare provider tells you to take it.
- Do not change your dose of MYQORZO without talking to your healthcare provider first.
- Take MYQORZO 1 time a day, at about the same time each day,
- Take MYQORZO with or without meals.
- Swallow the MYQORZO tablet whole.
- If you miss a dose of MYQORZO, take it as soon as possible and take your next dose at your regularly scheduled time the next day. Do not take 2 doses on the same day to make up for a missed dose.
- Your healthcare provider may change your dose or temporarily stop your treatment with MYQORZO if you have certain side effects.
- If you take too much MYQORZO, call your healthcare provider or Poison Control Help Line at 1-800-222-1222 or go to the nearest hospital emergency room right away.

What are the possible side effects of MYQORZO?**MYQORZO can cause serious side effects including:**

- **Heart failure.** See “What is the most important information I should know about MYQORZO?”

The most common side effect of MYQORZO is high blood pressure (hypertension).

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

You may also report side effects to Cytokinetics at 1-833-633-2986.

How should I store MYQORZO?

- Store MYQORZO at room temperature between 68°F to 77°F (20°C to 25°C).
- MYQORZO comes in a bottle with a child-resistant closure.

Keep MYQORZO and all medicines out of the reach of children.

General information about the safe and effective use of MYQORZO.

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

What are the ingredients in MYQORZO?

Active ingredients: aficamten

Inactive ingredients: croscarmellose sodium, hydroxypropyl cellulose, magnesium stearate (non-bovine), mannitol, microcrystalline cellulose, and sodium lauryl sulfate.

The tablet coating contains: carmine, FD&C Blue No. 2, glyceryl mono and dicaprylate, polyvinyl alcohol, polyvinyl alcohol graft polyethylene copolymer, talc and titanium dioxide.

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Patent information: <https://cytokinetix.com/patents>

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This Medication Guide has been approved by the U.S. Food and Drug Administration.

Issued: 12/2025



LOT 12345678
EXP YYYY-MM
SN 12345678901
GTIN 12345678901234

2000018329

NDC 82112-105-30
MYQORZO™
(aficamten) tablets

5 mg

Dispense with attached
Medication Guide

Rx Only **30 tablets**

Each tablet contains aficamten 5 mg
Recommended dosage:
See prescribing information.
Store at 20°C to 25°C (68°F to 77°F); excursions
permitted between 15°C to 30°C (59°F to 86°F)
[see USP controlled Room Temperature].
Keep out of reach of children.

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8211210530



LOT 12345678
EXP YYYY-MM
SN 12345678901
GTIN 12345678901234

2000018330

NDC 82112-110-30
MYQORZO™
(aficamten) tablets

10 mg

Dispense with attached
Medication Guide

Rx Only **30 tablets**

Each tablet contains aficamten 10 mg
Recommended dosage:
See prescribing information.
Store at 20°C to 25°C (68°F to 77°F); excursions
permitted between 15°C to 30°C (59°F to 86°F)
[see USP controlled Room Temperature].
Keep out of reach of children.

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8211211030



LOT 12345678
EXP YYYY-MM
SN 12345678901
GTIN 12345678901234

2000018331

NDC 82112-115-30
MYQORZO™
(aficamten) tablets

15 mg

Dispense with attached
Medication Guide

Rx Only **30 tablets**

Each tablet contains aficamten 15 mg
Recommended dosage:
See prescribing information.
Store at 20°C to 25°C (68°F to 77°F); excursions
permitted between 15°C to 30°C (59°F to 86°F)
[see USP controlled Room Temperature].
Keep out of reach of children.

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8211211530



LOT 12345678
EXP YYYY-MM
SN 12345678901
GTIN 12345678901234

2000018332

NDC 82112-120-30
MYQORZO™
(aficamten) tablets

20 mg

Dispense with attached
Medication Guide

Rx Only **30 tablets**

Each tablet contains aficamten 20 mg
Recommended dosage:
See prescribing information.
Store at 20°C to 25°C (68°F to 77°F); excursions
permitted between 15°C to 30°C (59°F to 86°F)
[see USP controlled Room Temperature].
Keep out of reach of children.

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