





empowering empowering HVES

Sarcomere directed therapies

2023 CYTOKINETICS, All Rights Reserved. YTOKINETICS® and the C-shaped logo are registered trademarks of Cytokinetics in the U.S. and certain other countrie ctual patients who consented to use of their name, image, and condition.

Forward-Looking Statements

This Presentation contains forward-looking statements for purposes of the Private Securities Litigation Reform Act of 1995 (the "Act"). Cytokinetics disclaims any intent or obligation to update these forward-looking statements and claims the protection of the Act's Safe Harbor for forward-looking statements. Examples of such statements include, but are not limited to, statements express or implied related Cytokinetics' research and development and commercial readiness activities, including the initiation, conduct, design, enrollment, progress, continuation, completion, timing and results of clinical trials, projections regarding growing prevalence, low survival rates and market opportunity in heart failure, hypertrophic cardiomyopathy (HCM) or heart failure with preserved ejection fraction (HFpEF); projections regarding the size of the addressable patient population for *omecamtiv mecarbil, aficamten,* CK-136, CK-586 or any of our other drug candidates; Cytokinetics' commercial readiness for *omecamtiv mecarbil*; the likelihood and/or timing of regulatory approval for our new drug application for *omecamtiv mecarbil* or any future new drug application for any of our other drug candidates; the timing of commencement of a second phase 3 clinical trial of *aficamten* as a monotherapy in patients with obstructive HCM, the timing of commencement of a phase 3 clinical trial of *aficamten* in nonobstructive HCM, the efficacy or safety of omecamtiv mecarbil, aficamten, CK-136, CK-586 or any of our other drug candidates, our ability to fully enroll or to announce the results of any of our clinical trials by any particular date; Cytokinetics' cash expenditures or runway; the results of any of our interactions with the FDA or any other regulatory authority regarding omecamtiv mecarbil or any of our other drug candidates; the properties, potential benefits and commercial potential of aficamten, omecamtiv mecarbil, CK-136, CK-586 or any of Cytokinetics' other drug candidates. Such statements are based on management's current expectations; but actual results may differ materially due to various risks and uncertainties, including, but not limited to, potential difficulties or delays in the development, testing, regulatory approvals for trial commencement, progression or product sale or manufacturing, or production of Cytokinetics' drug candidates that could slow or prevent clinical development or product approval, including risks that current and past results of clinical trials or preclinical studies may not be indicative of future clinical trial results, patient enrollment for or conduct of clinical trials may be difficult or delayed, Cytokinetics' drug candidates may have adverse side effects or inadequate therapeutic efficacy, the FDA or foreign regulatory agencies may delay or limit Cytokinetics' ability to conduct clinical trials, and Cytokinetics may be unable to obtain or maintain patent or trade secret protection for its intellectual property; Cytokinetics may incur unanticipated research, development and other costs or be unable to obtain financing necessary to conduct development of its products; standards of care may change, rendering Cytokinetics' drug candidates obsolete; and competitive products or alternative therapies may be developed by others for the treatment of indications Cytokinetics' drug candidates and potential drug candidates may target. These forward-looking statements speak only as of the date they are made, and Cytokinetics undertakes no obligation to subsequently update any such statement, except as required by law. For further information regarding these and other risks related to Cytokinetics' business, investors should consult Cytokinetics' filings with the Securities and Exchange Commission (the "SEC").

Our Mission

To bring forward new medicines to improve the healthspan of people with devastating cardiovascular and neuromuscular diseases of impaired muscle function.



Achieve regulatory approvals for at least two drugs arising from our pipeline

Build commercial capabilities to market and sell our medicines reflective of their innovation and value

Generate sustainable and growing revenues from product sales

• Double our development pipeline to include ten therapeutic programs

• Expand our discovery platform to muscle energetics, growth and metabolism

Be the science-driven company people want to join and partner with

Our vision is to be the

leading muscle biology

biopharma company that meaningfully improves the lives of patients with diseases of impaired muscle function through access to our

pioneering medicines

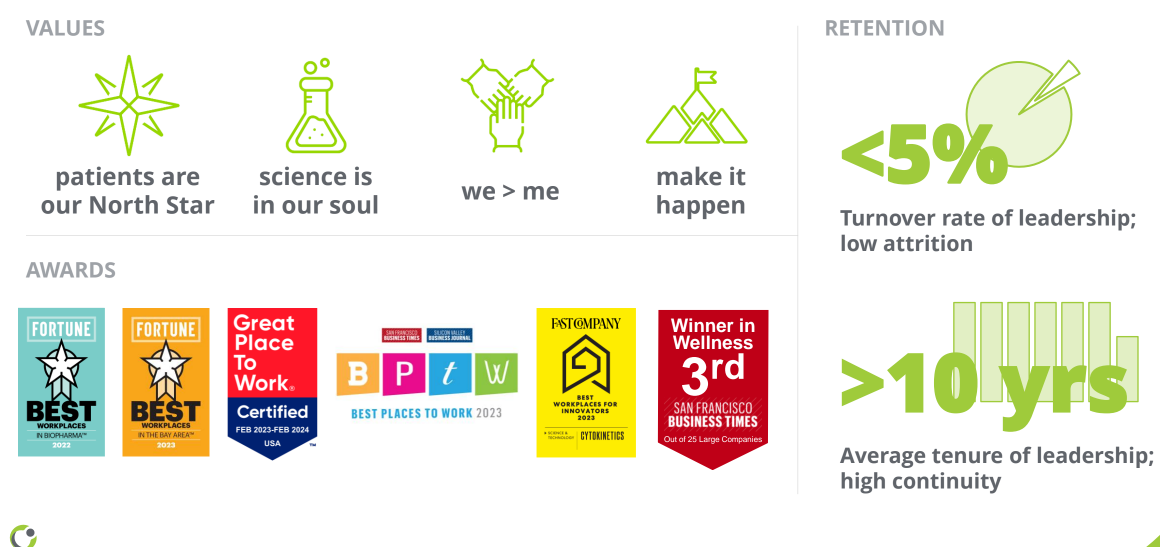
2025 Leading with Science, Delivering for Patients

VISI()I

As always, we will support disease advocacy groups elevating the patient voice and live by our values of integrity, fairness and compassion in all that we do.

Cytokinetics

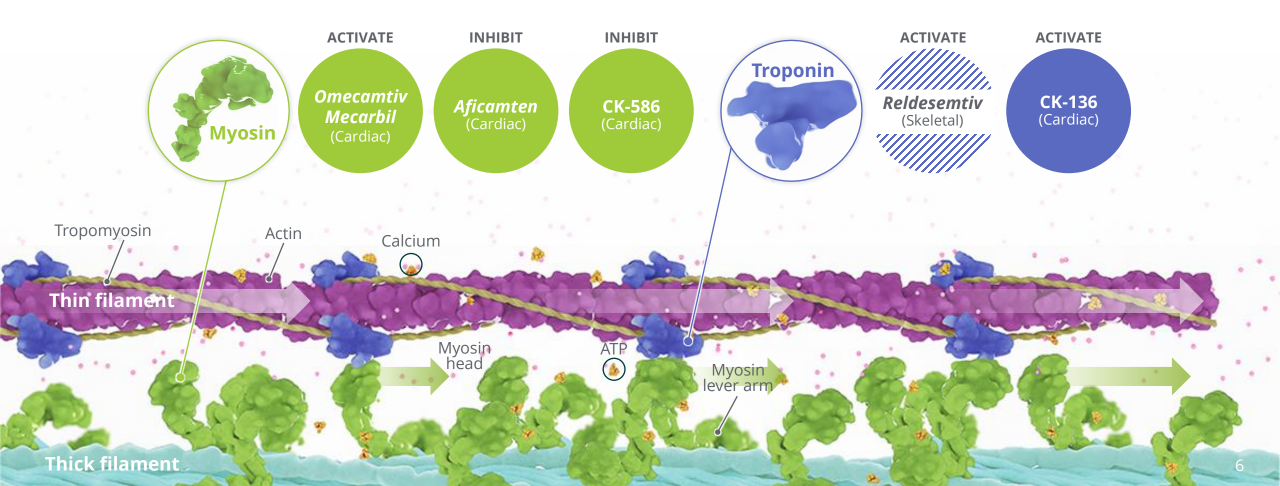
A Great Place to Work; Uncommon Continuity of Team



Cytokinetics

Sarcomere Directed Drug Development

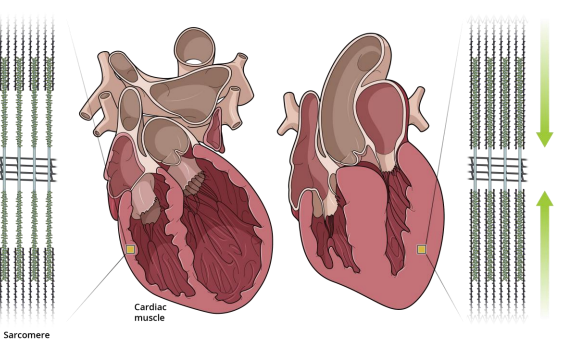
The sarcomere is a molecular structure found in skeletal and cardiac muscle that enables myocytes to contract and generate force



Contractile Dysfunction Underlies Cardiac Diseases

Decreased Cardiac Contractility

- Heart Failure with Reduced Ejection Fraction (HFrEF)
- Genetic Dilated Cardiomyopathy
- Pulmonary Hypertension with Right Ventricular Heart Failure

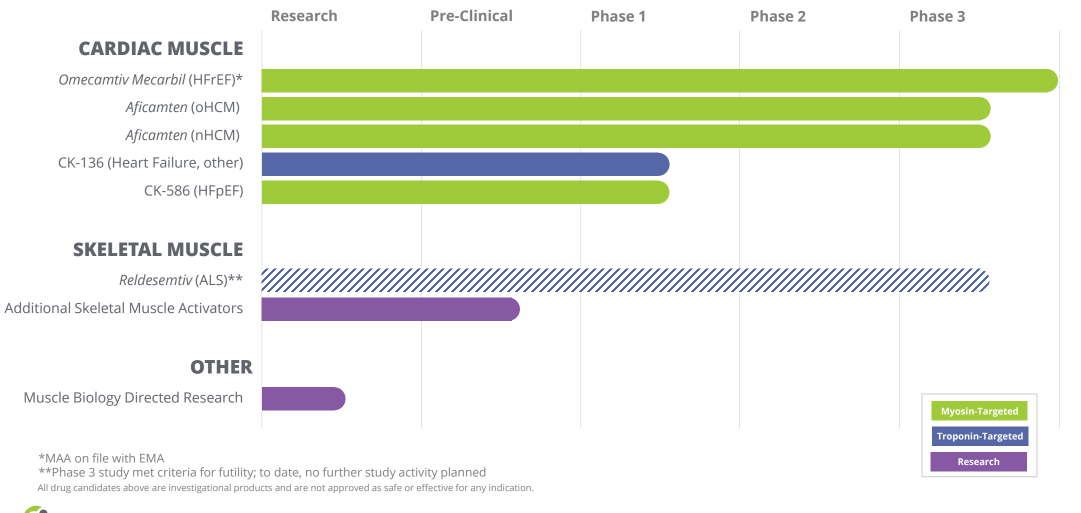


Increased / Preserved Cardiac Contractility

- Non-obstructive Hypertrophic Cardiomyopathy (nHCM)
- Obstructive Hypertrophic Cardiomyopathy (oHCM)
- Heart Failure with Preserved Ejection Fraction (certain HFpEF subsets)

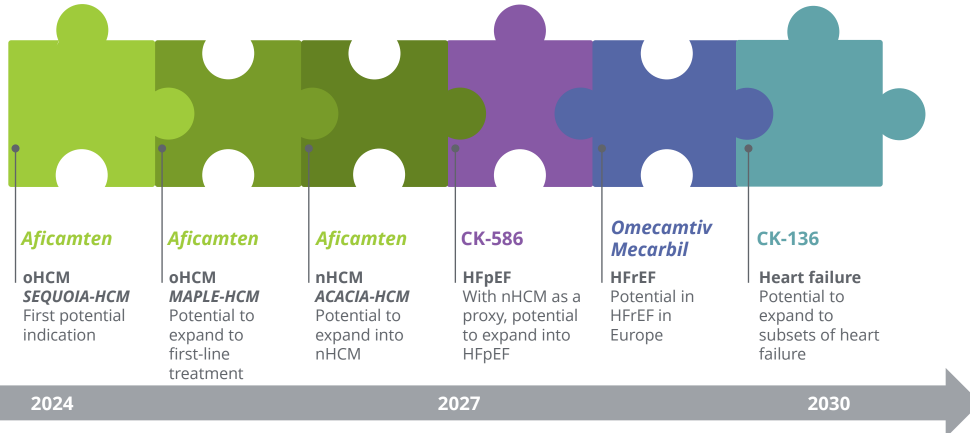


Pipeline of Novel Muscle-Directed Drug Candidates



Building a Specialty Cardiology Franchise Anchored by *Aficamten* Addressing severely ill and underserved populations in need of new therapies

Strategic expansion of clinical development program to various patient populations fuels leadership in cardiology



Aficamten, CK-586, omecamtiv mecarbil, and CK-136 are investigational drugs and have not been approved. Their safety and efficacy have not been established.



Sarcomere Directed Drug Development

Specialty Cardiovascular Portfolio

Aficamten Omecamtiv Mecarbil Emerging Pipeline – CK-136 & CK-586



Omecamtiv mecarbil , aficamten, CK-136 and CK-586 are investigational agents and have not been approved for use by the U.S. Food & Drug Administration (FDA) or any regulatory agency. The safety and effectiveness of these product has not been established.

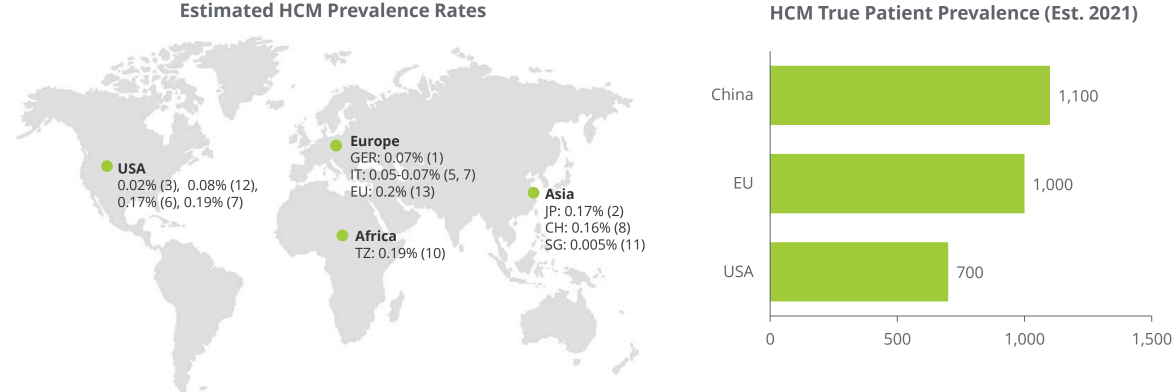
Aficamten



Aficamten is an investigational agent and has not been approved for use by the U.S. Food & Drug Administration (FDA) or any regulatory agency. The safety and effectiveness of this product has not been established.

HCM Prevalence: Significant and Growing Globally

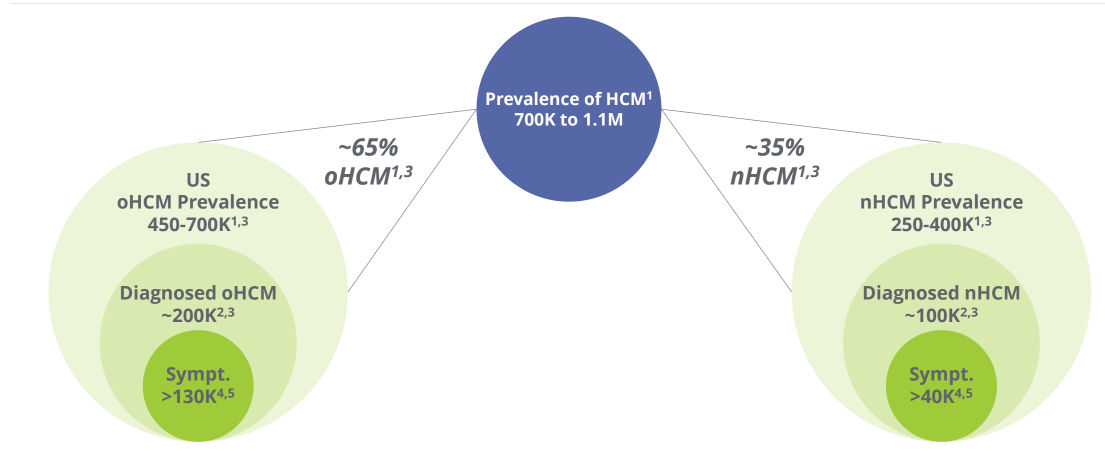
HCM prevalence estimates vary across geography and over time



Source: 1. Husser et al 2018 doi.org/10.1371/journal.pone.0196612; 2. Hada et al 10.1016/s0002-9149(87)80107-8; 3. Codd 1989 10.1161/01.cir.80.3.564; 4. Maron et al 1995 10.1161/01.cir.92.4.785; 5. Corrado et al 1998 10.1056/NEJM199808063390602; 6. Maron et all 1999 10.1001/jama.281.7.650; 7. Nistri et al 2003 10.1016/s0002-9149(03)00132-2; 8. Zou et al 2004 10.1093/aje/kwh090; 9. Maron 2004 https://doi.org/10.1016/j.amjmed.2003.10.012; 10. Maro 2006 10.1258/004947506778604904; 11. Ng et al 2011 10.1093/europace/eur051; 12. Butzner et al 2021 10.1016/j.amjcard.2021.08.024; 13. Cardim et al 2011 10.1016/j.repc.2011.09.005



Opportunity for CMIs in Diagnosed, Symptomatic HCM Patients Potential for nearly 200K patients eligible for CMIs in 2025



Projections and forecasts for illustration.

1. Cardiovascular Research Group: CVrg Market Strategies: Heart Failure, p 48, Q4 2022; Maron BJ: et al.: Prevalence of Hypertrophic Cardiomyopathy In A General Population of Young Adults, Circulation 1995;92;785-789; Semsarian C. et al: New Perspectives on the Prevalence of Hypertrophic Cardiomyopathy, J. Am, Coll. Cardiol. 2015; 65: 1249-1254;

2. DoF: SHA; Symphony PTD (Patient Transaction Data): Includes patients diagnosed since 2016 and having any HC transaction in the claims data universe in the last year June 2022-May 2023);

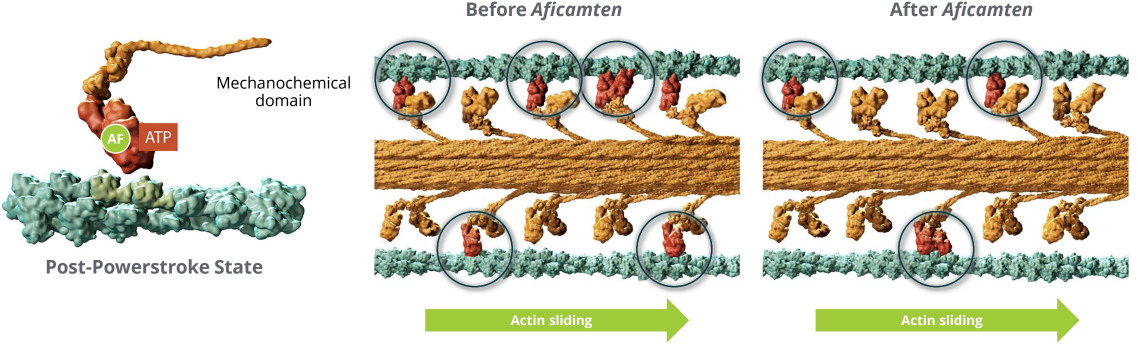
3. Lu DY et al: Clinical Outcomes in Patients With Nonobstructive, Labile, and Obstructive Hypertrophic Cardiomyopathy. J. Am. Heart Assoc.2018;7:1-11 4) DoF: SHA Symphony PTD (Patient Transaction Data) includes any patients with symptoms in the last 2 years: angina, dyspnea, fatigue, palpitations, syncope, tachycardia; and/or treatments in the past 2 years: bb, ccb, dyso, ralo, Camzyos; 5) DoF Primary market research: 443 HCPs treating HCM - % of nHCM patients not considered under control with current SOC.



Aficamten: Proposed Mechanism of Action

Aficamten stabilizes myosin in the released post-powerstroke state unable to hydrolyze ATP

"Fewer hands pulling on the rope"



Aficamten is an investigational agent and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Aficamten: Aspirational Target Profile Potential next-in-class cardiac myosin inhibitor



Aspirational information. Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



 Not for Promotional Use, For Investors Only

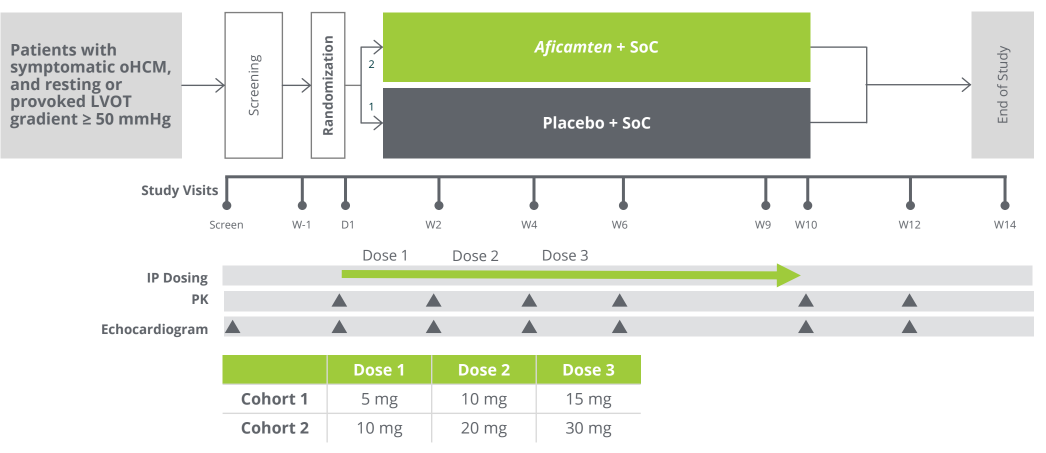
 OVERVIEW
 AFICAMTEN
 OMECAMTIV MECARBIL
 EMERGING PIPELINE
 CORPORATE PROFILE

REDWOOD-HCM: Cohorts 1 & 2



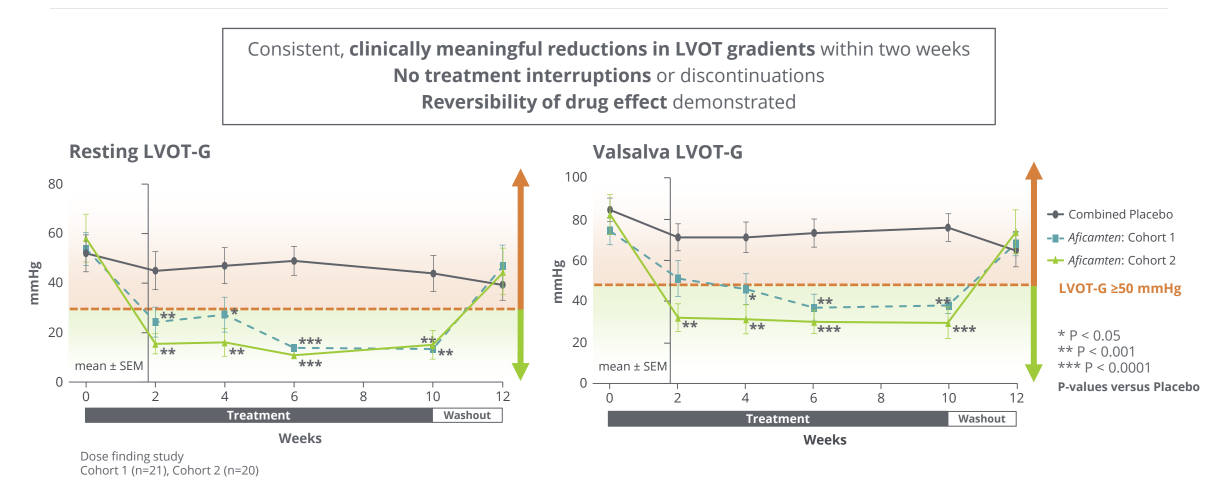
Patients with symptomatic oHCM on background therapy excluding disopyramide

Two sequential dose-finding cohorts





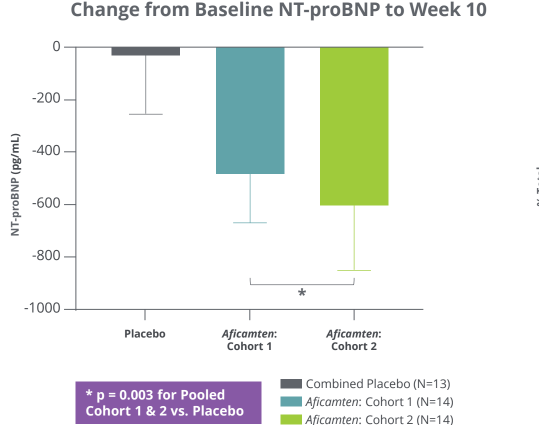
REDWOOD-HCM: Robust Reduction of LVOT Gradients REDWOOD



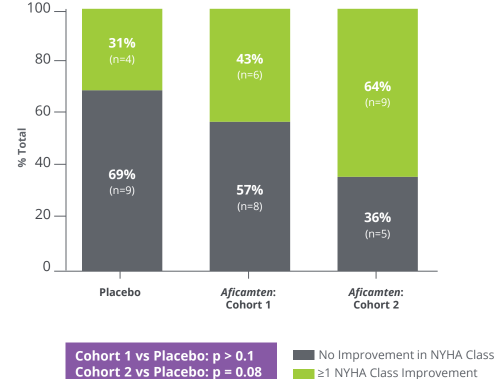
Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established. Maron M, et. al. Phase 2 Study of *Aficamten* in Patients With Obstructive Hypertrophic Cardiomyopathy. JACC. January 2023.

Change from Baseline in NT-proBNP & NYHA Class Cohorts 1 & 2





Improvement in Heart Failure Symptoms (NYHA Class)



Week 10 Responder Definition: Improvement in NYHA Class ≥1

Aficamten is an investigational agent and is not approved by any regulatory agency. Its safety and efficacy have not been established. Maron M, Abraham T, Masri A, et al. "REDWOOD-HCM: A Randomized, Double-blind, Placebo-controlled, Dose-finding Trial of the Cardiac Myosin Inhibitor, *Aficamten*, In Obstructive Hypertrophic Cardiomyopathy". HFSA 2021.



OVERVIEW AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

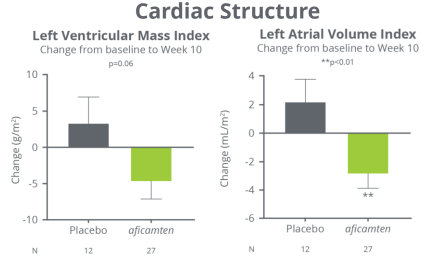
Improved Cardiac Structure and Diastolic Function



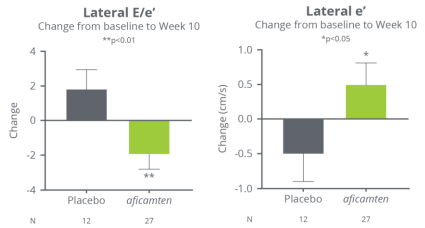
Cohorts 1 & 2: Early signs of improvement in cardiac structure and myocardial relaxation

Treatment with *aficamten* for 10 weeks resulted in:

- Significant reduction in left atrial volume index
- Trend towards a **reduction in LV mass index**
- Improved diastolic function
 - reduction in lateral E/e' (p<0.01)
 - increase in lateral e' (p<0.05))



Diastolic Function

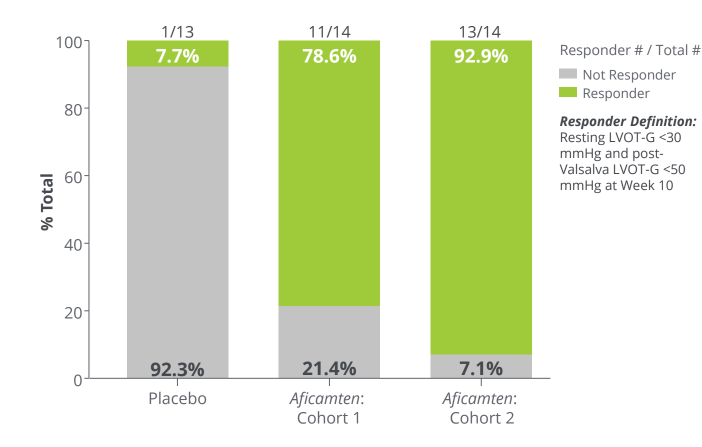


Aficamten is an investigational agent and is not approved by any regulatory agency. Its safety and efficacy have not been established. Abraham T. et al. "Early Cardiac Structural and Functional Reverse Remodeling in Obstructive Hypertrophic Cardiomyopathy after 10 Weeks of *Aficamten* Therapy: Analyses from REDWOOD-HCM". ASE 2022.



Response Rates on Treatment with *Aficamten*





- Consistent, clinically meaningful reductions in LVOT gradients within two weeks
- No treatment interruptions or discontinuations
- No treatment-related SAEs
- Reversibility of drug effect
 demonstrated
- Statistically significant reductions in NT-proBNP
- Improvement in NYHA class

Aficamten is an investigational agent and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Maron M, Abraham T, Masri A, et al. "REDWOOD-HCM: A Randomized, Double-blind, Placebo-controlled, Dose-finding Trial of the Cardiac Myosin Inhibitor, *Aficamten*, In Obstructive Hypertrophic Cardiomyopathy". HFSA 2021.

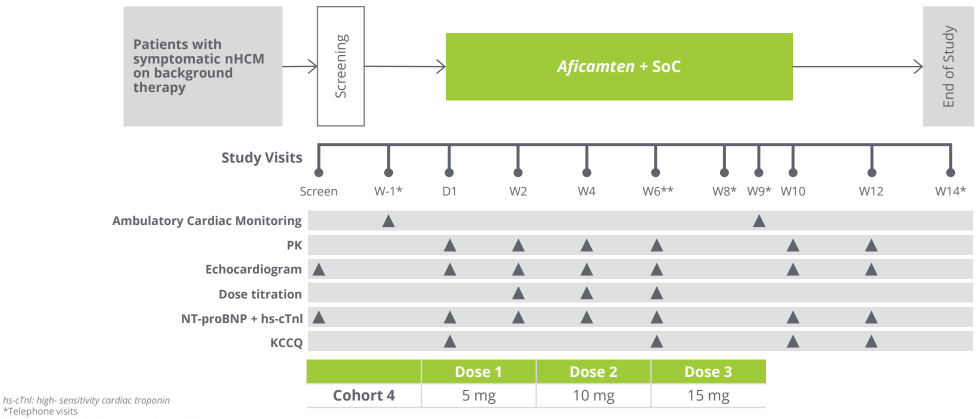


REDWOOD-HCM: Cohort 4



Patients with symptomatic nHCM on background therapy

Results presented at ESC Heart Failure 2023



*Telephone visits **Patient can only be down-titrated at Week 6

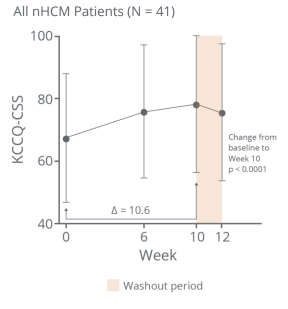


Significant Improvements in KCCQ & NYHA Class Cohort 4

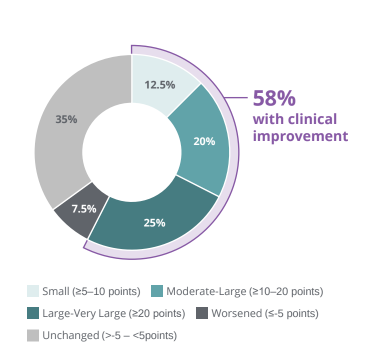


85% of patients achieved 15 mg dose; no discontinuations due to adverse events

Kansas City Cardiomyopathy Questionnaire Mean improvement in KCCQ of 10.6 points

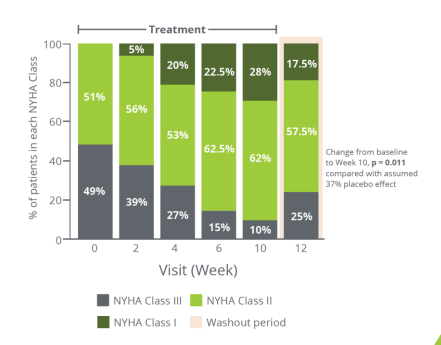


Data presented as mean and standard deviation



Categorical Changes at Week 10 in KCCQ-CSS

NYHA Functional Class 56% of patients improved by ≥1 NYHA class



Aficamten is an investigational agent and is not approved by any regulatory agency. Its safety and efficacy have not been established. Masri A. et al. "*Aficamten* in Patients with Symptomatic Non-Obstructive Hypertrophic Cardiomyopathy (REDWOOD-HCM Cohort 4)". ESC HF 2023.



Change in Baseline in Biomarkers & Angina Frequency Cohort 4

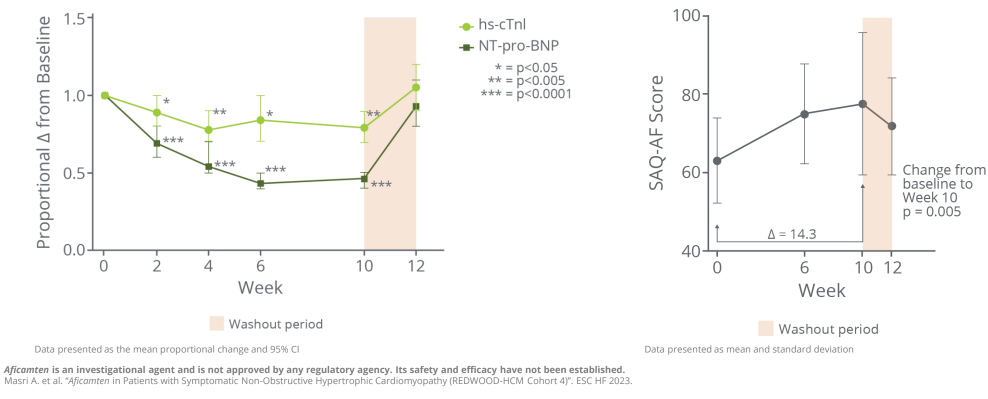
Proportional Change from Baseline in Cardiac Biomarkers

Mean reduction in high-sensitivity cardiac troponin of 21% Mean reduction in NT-proBNP of 55%

Seattle Angina Questionnaire Angina Frequency (SAQ-AF)

Reduction in frequency of angina from daily or weekly, to weekly or monthly

REDWOOD



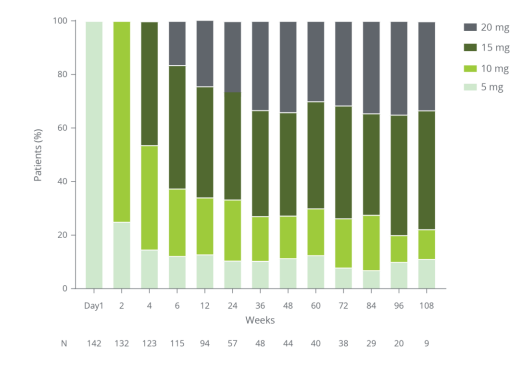
FOREST-HCM: Baseline Characteristics



Baseline characteristics indicate substantial disease burden; ~2/3 patients achieving 15 or 20 mg

FOREST-HCM
оНСМ
N=143*
60.4 (13.2)
65 (45.5)
29.2 (4.5)
82 (58)
60 (42)
40 (28.0)
90 (62.9)
14 (9.8)
27 (18.9)
69 (5)
56.8 (33.2)
93.1 (37.9)

Dose of Aficamten



Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Cytokinetics

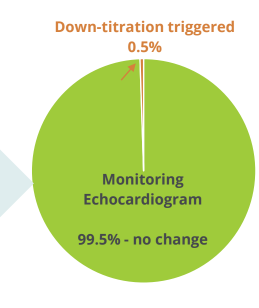
No Dose Interruptions During Titration; Few Dose Reductions During Maintenance Forest

Dose Titration Phase

- No treatment-related LVEF <50% during the titration period
- Of the 94 patients having completed the titration period,
 ~2/3 are receiving 15 and 20 mg qd
- Approximately 30% of patients have reduced doses or discontinued background therapy at the discretion of the treating physician and/or request from the patient

Maintenance Phase

- 579 monitoring echocardiograms completed* in oHCM patients
- None with LVEF <40% requiring treatment interruption
- 3 patients (0.5%) with LVEF <50%
 - Two asymptomatic patients (LVEF of 47% and 49%) resulting in perprotocol dose reduction
 - One patient with atrial fibrillation (unrelated) and LVEF of 47%
 - All 3 patients are currently receiving *aficamten* with apparent relief from obstruction, symptoms & improved biomarkers



Target dose defined as achieved if Valsalva LVOT-G \leq 30 mmHg or no dose change for 2 consecutive visits

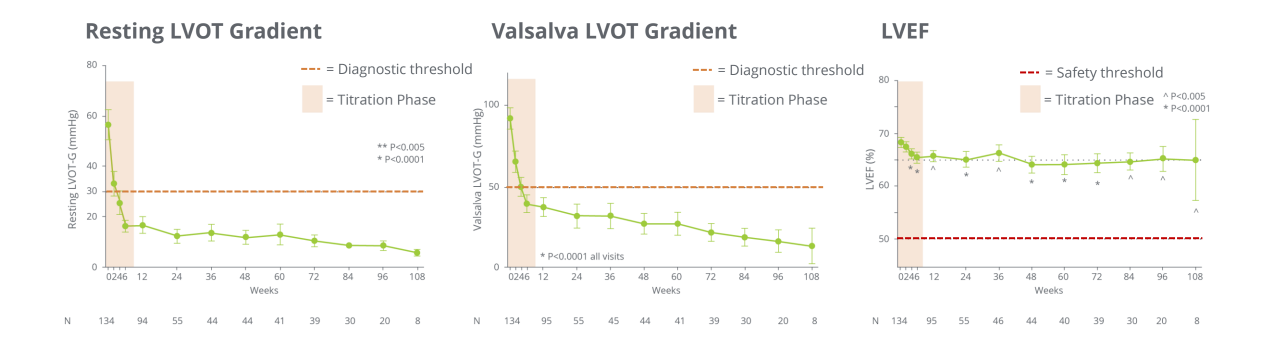
Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established. * *As of Sept 15, 2023.*



Durable Effects of Aficamten on LVOT-G & LVEF



Resting & provoked gradients remain below diagnostic threshold for >2 years, LVEF remains flat after titration

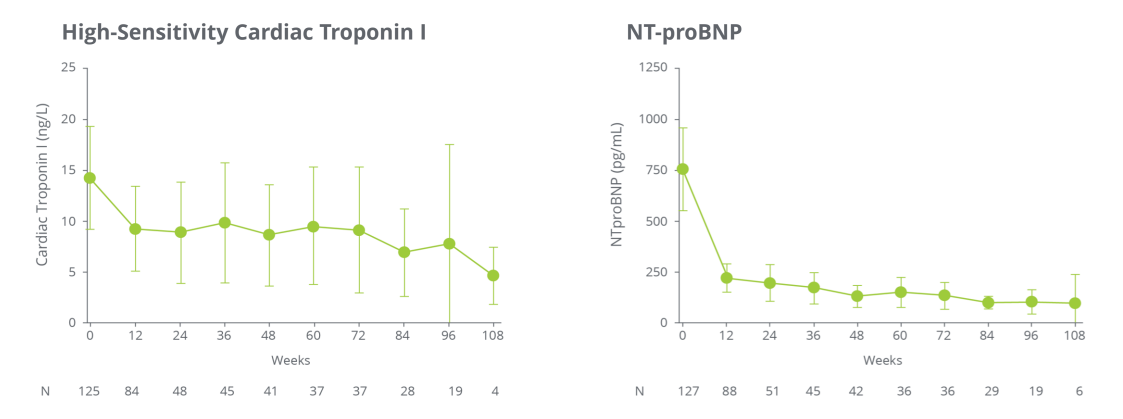


Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Durable Effects of *Aficamten* on Biomarkers



Sustained relative reductions in high-sensitivity Troponin I (~30%) & NT-proBNP (~70%) observed



Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



OVERVIEW AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

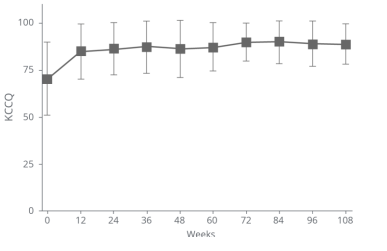
Not for Promotional Use, For Investors Only overview AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

Durable Effects of Aficamten on Clinical Endpoints

KCCQ-CSS

Cytokinetics

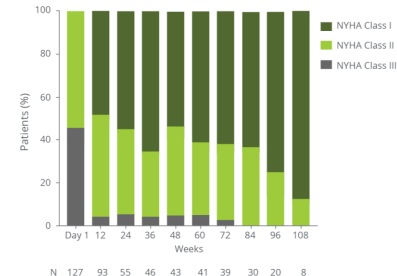
71% of patients had \geq 5-point KCCQ-CSS increase 30% of patients had \geq 10-point KCCQ-CSS increase



NYHA Class

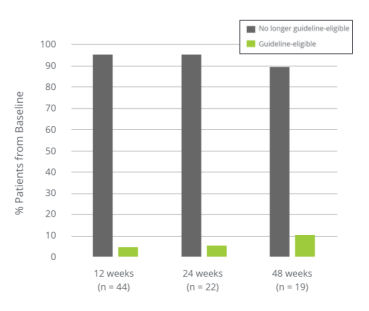
~50% of patients were asymptomatic at 1 year

>80% of patients improved ≥1 NYHA Class at every visit after initiation of *aficamten*



Guideline-Eligible for SRT

90% of SRT-eligible patients at baseline are no longer SRT-eligible



28

Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Safety Data: Phase 2 & OLE



• <u>oHCM</u> → <u>Cohorts 1, 2, & 3:</u> After 10-weeks of treatment

- 2 SAEs reported in 41 *aficamten*-treated → none were related to *aficamten* treatment
- No treatment interruptions or discontinuations
- Transient and asymptomatic decrease in LVEF < 50% occurred in 2 of 41 *aficamten*-treated patients

<u>nHCM</u> → <u>Cohort 4</u>: After 10-weeks of treatment

Cvtokinetics

- Well tolerated 85% achieved maximal dose (15 mg)
- Transient and asymptomatic decrease in LVEF < 50% occurred in 3 of 41 *aficamten*-treated patients
- One death unrelated to *aficamten* treatment sudden cardiac death (SCD) in patient with history of aborted SCD x 2 prior to participation. Two days before event, LVEF was normal, NT-proBNP was lower and plasma concentration of *aficamten* was within the expected range



- Almost all eligible patients choose to participate in the OLE
- Echocardiography-guided dose titration of *aficamten* is **managed entirely by the treating physicians**
- 2/3 of patients achieve **higher doses;** no low LVEF events requiring treatment interruption
- 94 patients have completed the titration period none have experienced LVEF <50%
- 99.5% of monitoring echocardiograms have not led to a dose reduction
- Clinical, hemodynamic & biochemical markers of efficacy continue to indicate sustained efficacy following exposures for > 2-years
- Of the patients that are guideline-eligible for septal reduction therapies at baseline, ~90% are no longer eligible after dose titration
- *Aficamten* has been **generally well-tolerated**, with 60% of patients experiencing at least one treatment emergent adverse event (TEAE) but there were no treatment-related serious adverse events (SAEs) as assessed by investigators, and no patient deaths

SEQUOIA-HCM: Phase 3 Trial



Completed enrollment; expect topline results in late December

Primary endpoint: Change in pVO₂ by CPET from baseline to Week 24

Secondary objectives include measuring change in KCCQ & improvement in NYHA class at week 12 and 24

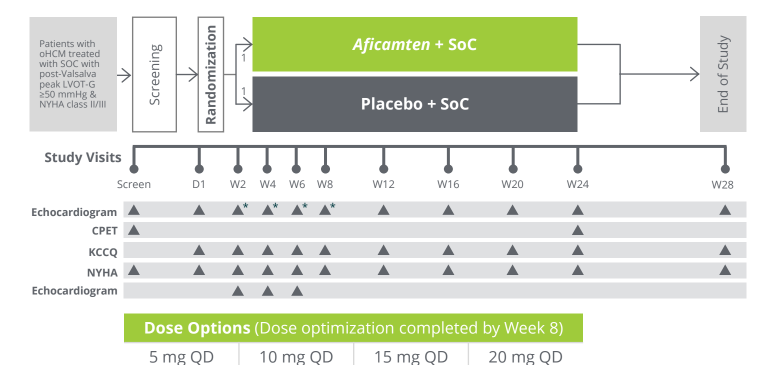
Enrolled 282 patients treated with standard of care with:

- resting LVOT-G ≥30 mmHg,
- post-Valsalva LVOT-G ≥50 mmHg,
- NYHA Class II or III,
- exercise performance <80% predicted

Individualized dose up-titration based on echocardiography: LVEF ≥55%, post-Valsalva LVOT-G ≥30 mmHg

SOC: standard of care * Focused echocardiogram

Cvtokinetics



SEQUOIA-HCM: Baseline Characteristics



Baseline characteristics reflect highly symptomatic patient population with reduced exercise capacity

- Significant symptom burden despite background therapy
- 61% of patients on beta-blockers
- Baseline pVO2 reflects patient population with reduced exercise capacity

a Unless otherwise indicated. b >100% total due to overlap in ethnicity and race. c NYHA FC III and any LVOTO ≥50 mmHg d Combines hypertension and essential hypertension. e Combines T2DM, T1DM, and DM CCB, calcium channel blocker; DM, diabetes mellitus, including types 1 and 2; IQR, interquartile range

Baseline Characteristics (N=282)	n (%) or Mean (SD) ^a	Baseline Characteristics (N=282)	n (%) or Mean (SD)ª
Demographics		HCM Medical Therapies	
Age, years	59.1 (12.9)	Beta-blocker	172 (61.0)
Female	114 (40.4)	Non-dihydropyridine calcium	75 (26.6)
Race/ethnicity ^b		channel blocker	
White	222 (78.7)	Disopyramide	36 (12.8)
Black	3 (1.1)	HCM Symptoms	
Asian	53 (18.8)	KCCQ-CSS	74.7 (18.0)
Hispanic	9 (3.2)	NYHA class II/III/IV	214 (75.9)
Other	4 (1.4)		67 (23.8)
Region			1 (0.4)
United States	94 (33.3)	SRT guideline eligible	68 (24.1)
China	46 (16.3)	Comorbidities	
Europe and Israel	142 (50.4)	Hypertension ^d	136 (48.2)
Vital Signs		Diabetes ^e	24 (8.5)
Weight, kg	81.6 (15.7)	Permanent atrial fibrillation	1 (0.4)
Body mass index, kg/m ²	28.1 (3.7)	Paroxysmal atrial fibrillation	40 (14.2)
Systolic blood pressure, mmHg	125.3 (16.1)	CPET Metrics	
Diastolic blood pressure, mmHg	74.4 (10.6)	Treadmill	1EE (EE O)
Heart rate, bpm	65.6 (11.2)		155 (55.0)
HCM History		Peak VO ₂ , mL/kg/min	18.5 (4.5)
History of known HCM-causing	48 (17.0)	Peak VO ₂ , % of predicted	56.9 (11.8)
gene mutation		maximum ^f	
Positive family history of HCM	71 (25.2)	Total workload, watts	122.4 (41.2)
Time since initial HCM diagnosis,	5.9 (1.7 – 8.5)	Biomarker	
median (IQR), years		hs-cTnl median (IQR), ng/L	21.1 (7.7 – 27.3)

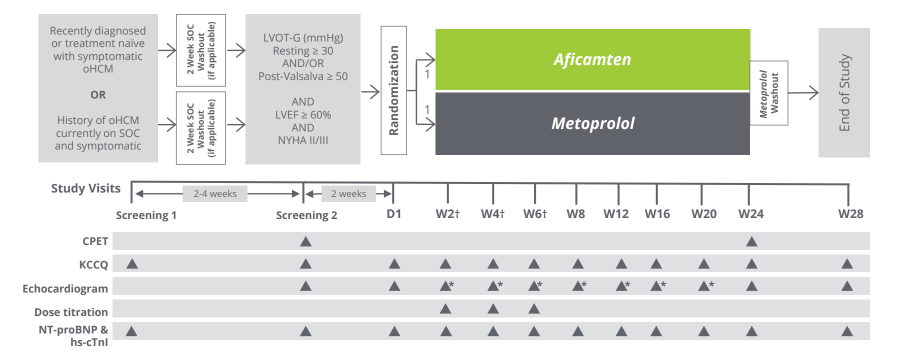


MAPLE-HCM: Phase 3 Monotherapy Trial Open to enrollment



Active-comparator trial of *aficamten* as monotherapy vs. *metoprolol* in patients with oHCM

- Trial to enroll approximately 170 patients
- Primary endpoint: change in peak VO2, assessed by CPET from baseline to Week 24
- Secondary endpoints: change in NYHA class, KCCQ, NT-proBNP, and measures of structural remodeling



SOC: standard of care * Focused echocardiogram

Not for Promotional Use, For Investors Only OVERVIEW AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

d 4-week follow up after last dose

^c Site-read focused echocardiogram for titration visit (sole criterion). *Aficamten* dose range 5-20 mg.

Randomization (1:1) N = 420

Screening

 \rightarrow

i

ACACIA-HCM: Pivotal Phase 3 Trial in nHCM Open to enrollment

Patients with:

Symptomatic

(NYHA Class

II/III) nHCM,

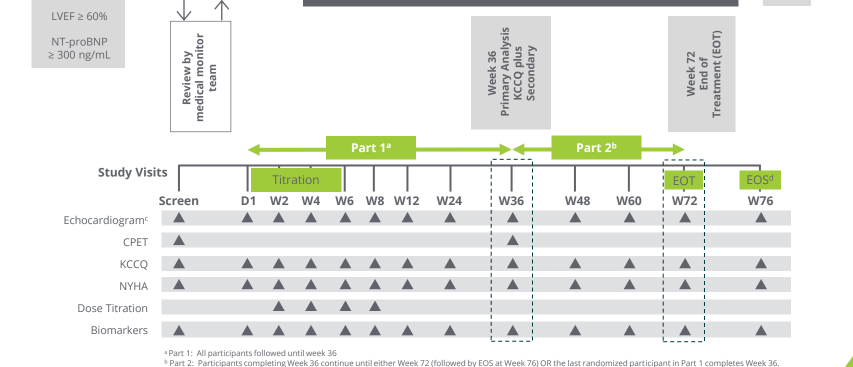
KCCQ-CSS

 \geq 30 and \leq 85

- Trial to enroll approximately 420 symptomatic nHCM patients
- Primary endpoint: change in KCCQ Clinical Summary Score from baseline to Week 36
- **5-20 mg doses**; 6-week titration period
- Secondary endpoints:
 - Change in pVO2, Ve/VCO2,
 - Left atrial volume index (LAVI)
 - NT-proBNP

Cytokinetics

- Proportion of patients with ≥1 class improvement in NYHA from baseline to Week 36
- Time to first cardiovascular event



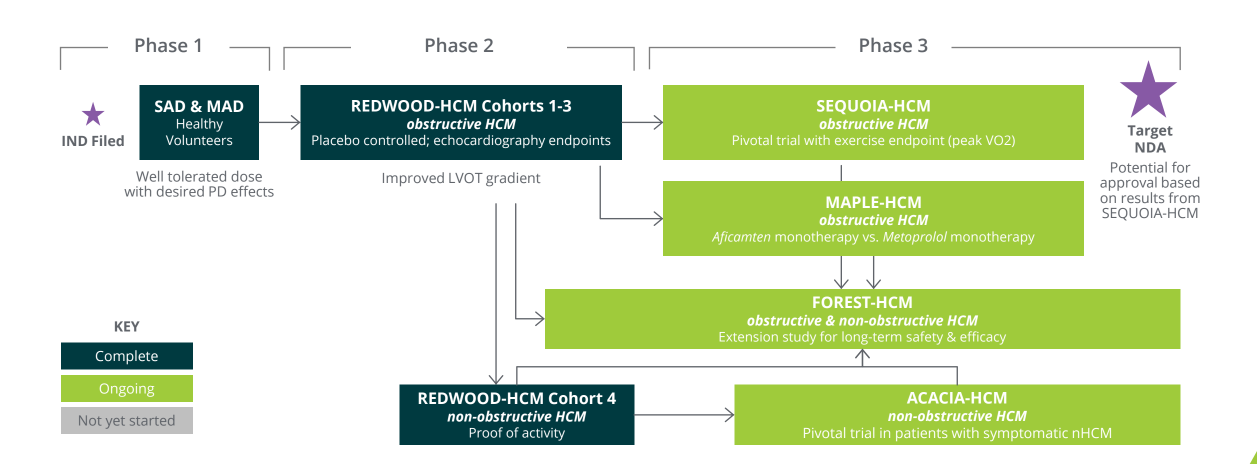
Aficamten

Placebo



End of Study (EOS)

Aficamten: Clinical Development Plan for HCM



Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

Cytokinetics

Aficamten: Planned Commercial Approach Driven by a relentless focus on our North Star: the HCM patient

Learn	Design	Build
Leverage deep	Engage with all	Tap into deep functional
understanding of	stakeholders to design	experience to build
patients, HCPs, payers,	an optimal customer	operational excellence
and community	experience	across launch functions

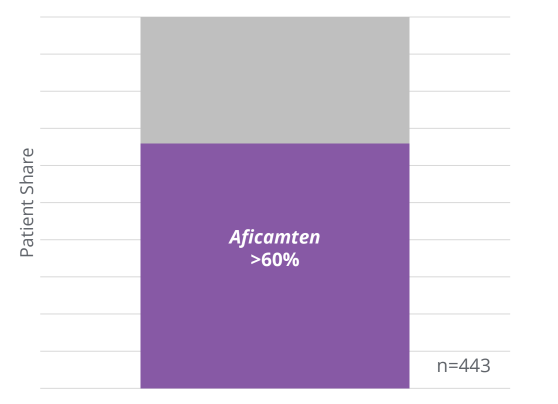
Our Focus to Date

Our 2024 Focus

Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



oHCM CMI Preference Shares in Eligible Patient Population*

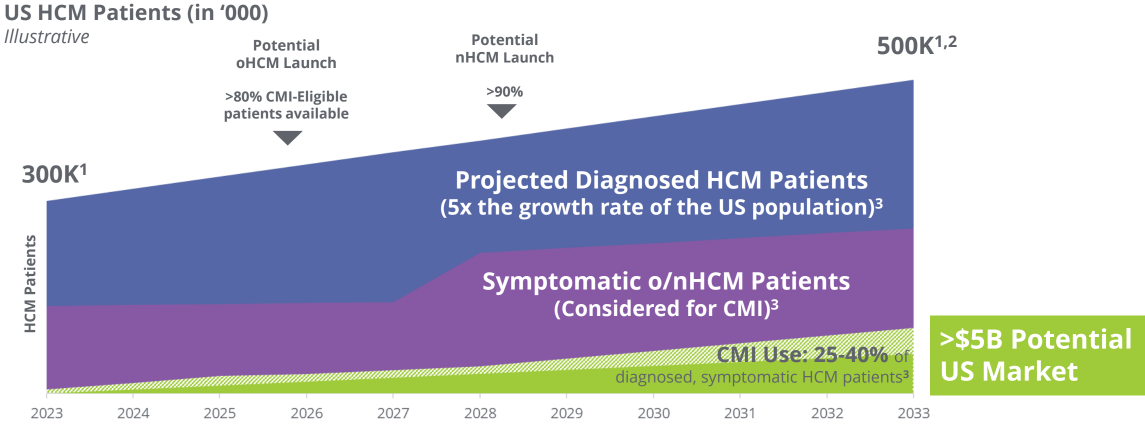


- Potential target product profile for *aficamten* interest creates **share opportunity** in newly treated CMI patients
- Aficamten is also expected to expand the total CMI market
- Key attributes that may drive preference include the potential for:
 - LVOT gradient reduction
 - Change in NYHA Functional Class
 - Pharmacodynamics/LVEF maintenance
 - Change in KCCQ
 - Absence of DDI

Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established. *Source: *Aficamten* Impact of Product Attributes on Product Preference Share n=443 cardiologists, Quantitative research including conjoint - Cogent



If *Aficamten* is Approved, Expect Majority of CMI-Eligible Patients Available at Launch **Diagnosis of HCM anticipated to grow 5x the rate of the general population**

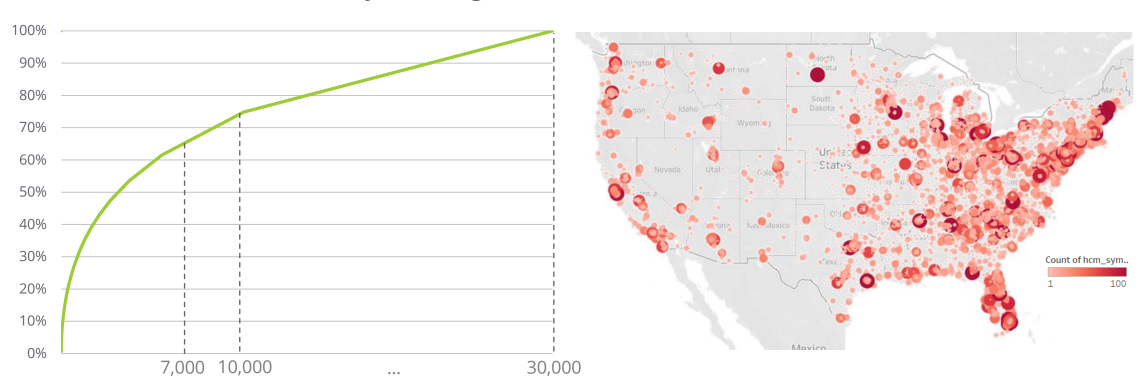


Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established. Projections and forecasts for illustration

Source: 1) DoF: SHA; Symphony PTD (Patient Transaction Data): Includes patients diagnosed since 2016 and having any HC transaction in the claims data universe in the last year June 2022-May 2023); 2) Butzner et al 2021 estimated a 8% growth rate in diagnosed HCM patients between 2013-2019 <u>https://www.ajconline.org/article/S0002-9149(21)00783-9/fulltext</u>; CYTK is forecasting an average growth rate of 5% over the coming decade; 3) Internal forecasts



Cardiologists Located in Concentrated Geographic Clusters Across the US **75% of the HCM patient volume is treated by 10,000 cardiologists**



Geographic Distribution of HCM Patients

Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

HCM Patient Concentration by Cardiologist

Note: includes only patients who are treated by a cardiologist - not all patients see a cardiologist; sample of 67K HCM patients Source: Symphony PTD (Patient Transaction Data); mapping of HCPs to HCOs using Definitive Healthcare Data 2023 and 7/2023 mapping; Patient volume by dominant Cardiologist Location 7/2023



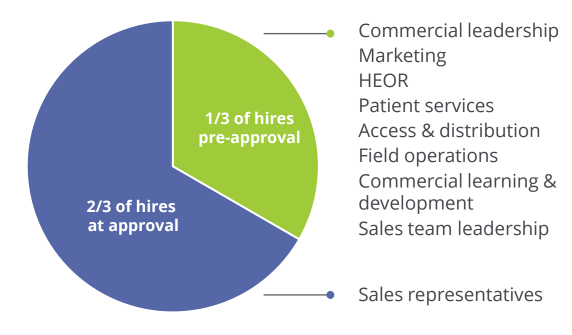
Not for Promotional Use, For Investors Only

OVERVIEW AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

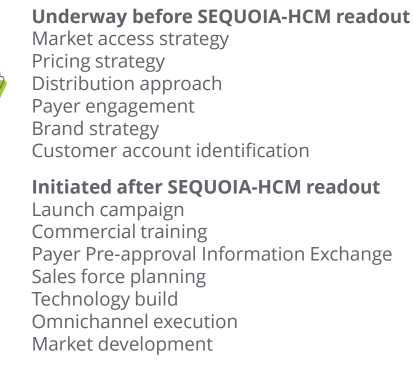
Gated Build of Commercial Infrastructure

Majority of spending to occur closer to approval in 2025

2/3 of hiring to occur at-approval



Activities initiated upon key de-risking events





 $\sqrt{2}$

Initiated upon FDA approval Media purchases Patient support programs

Aficamten is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



Omecamtiv Mecarbil



Omecamtiv mecarbil is an investigational agent and has not been approved for use by the U.S. Food & Drug Administration (FDA) or any regulatory agency. The safety and effectiveness of this product has not been established.

Omecamtiv Mecarbil: Current Status

No current plans to conduct additional clinical trial of *omecamtiv mecarbil*

Received CRL from FDA Feb 28, 2023

GALACTIC-HF not sufficiently persuasive to establish substantial evidence of effectiveness for reducing the risk of heart failure events and cardiovascular death in adults with chronic HFrEF



• Submitted Formal Dispute Resolution Request to the FDA

• Continue to pursue **approval** of *omecamtiv mecarbil* in Europe

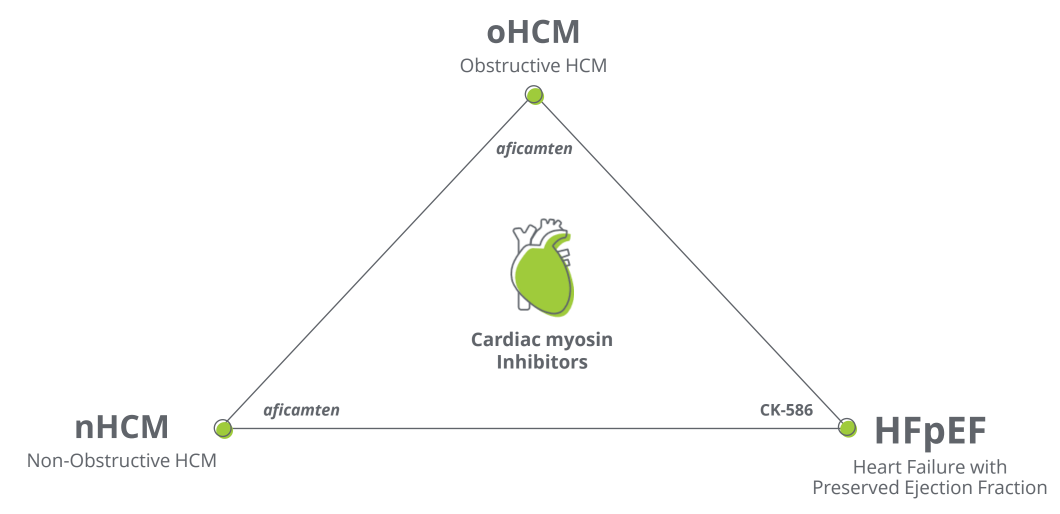


Emerging Cardiovascular Pipeline CK-136 & CK-586



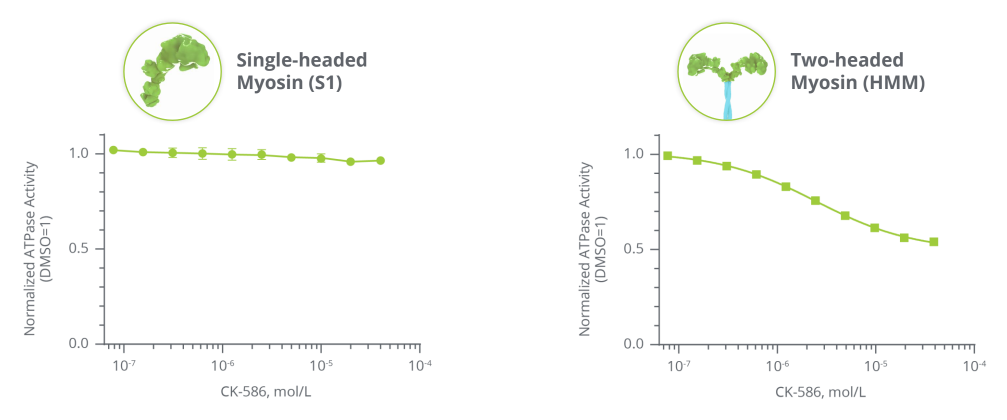
CK-136 and CK-586 are investigational agents and have not been approved for use by the U.S. Food & Drug Administration (FDA) or any regulatory agency. The safety and effectiveness of these products have not been established.

Novel Approach May Address Multiple Unmet Patient Needs



CK-586: Distinct Mechanism of Action from Aficamten

CK-586 inhibits actin-activated ATPase of HMM only; aficamten inhibits both S1 and HMM



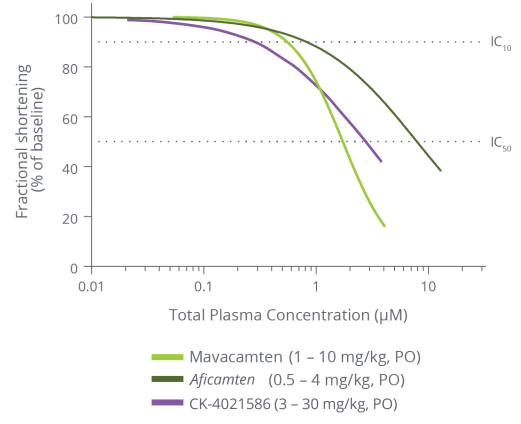
Based on preclinical testing

CK-586 is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



CK-586: Shallow In Vivo Concentration-Response

CK-586 is predicted to have a shorter half-life in humans than *aficamten*



Pharmacodynamic window Fractional shortening IC ₅₀ /IC ₁₀ ratio		
mavacamten	2.8x	
aficamten	9.9x	
CK-586	9.3x	

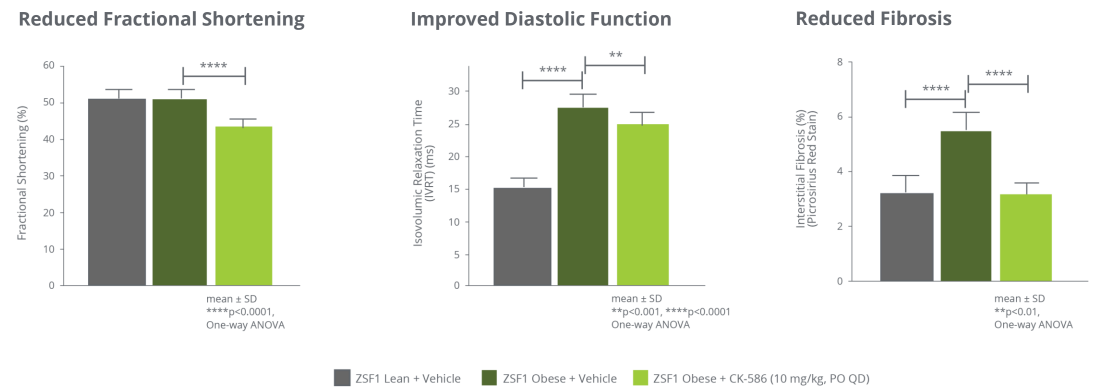
 IC_{10} : plasma concentration at 10% relative reduction in fractional shortening IC_{50} : plasma concentration at 50% relative reduction in fractional shortening

Compound half-life in humans	Actual	Predicted
aficamten	~3 days	2.8 days
CK-586	TBD	15 hours

CK-586 is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.

CK-586 is Efficacious in ZSF1 Obese Rat Model of HFpEF Model is representative of hypertensive, diabetic, metabolic aspects of HFpEF

10 weeks of treatment improved diastolic function and reduced cardiac fibrosis



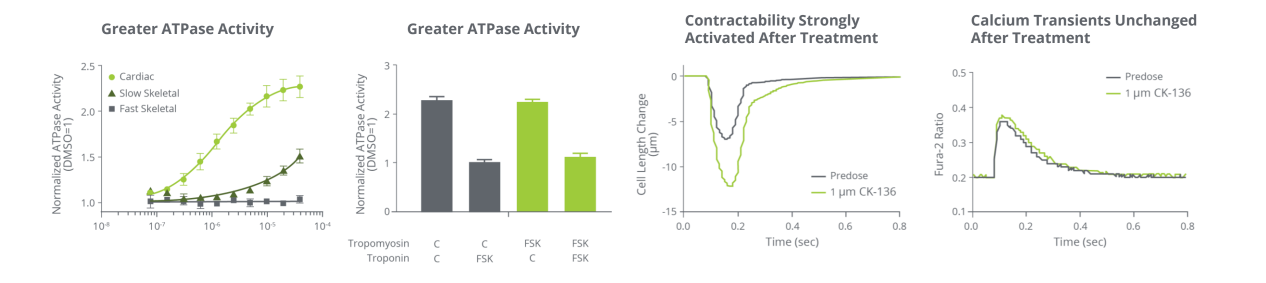
CK-586 is an investigational drug and is not approved by any regulatory agency. Its safety and efficacy have not been established.



CK-136: Mechanism of Action

Key biochemical and cellular features

The first selective cardiac troponin activator



¹PD Window = Maximum Tolerated Concentration (MTC) / 10% increase in Fractional Shortening (FS)

CK-136 is an investigational agent and has not been approved for use by any regulatory agency. Its safety and efficacy have not been established.

Cytokinetics

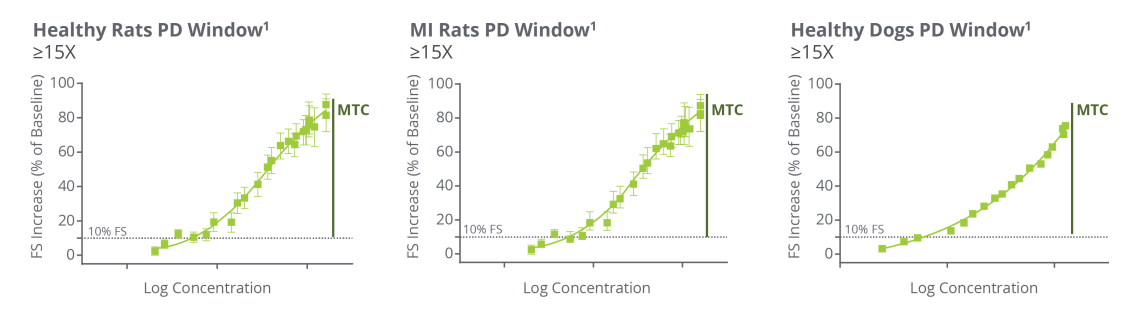
Not for Promotional Use, For Investors Only overview AFICAMTEN OMECAMTIV MECARBIL EMERGING PIPELINE CORPORATE PROFILE

CK-136: Exposure Response Relationship

Exposure-response of troponin activator is shallower than myosin activator

Analyzing single ascending dose data from Phase 1 study

Animal Models of Cardiac Function



¹PD Window = Maximum Tolerated Concentration (MTC) / 10% increase in Fractional Shortening (FS)

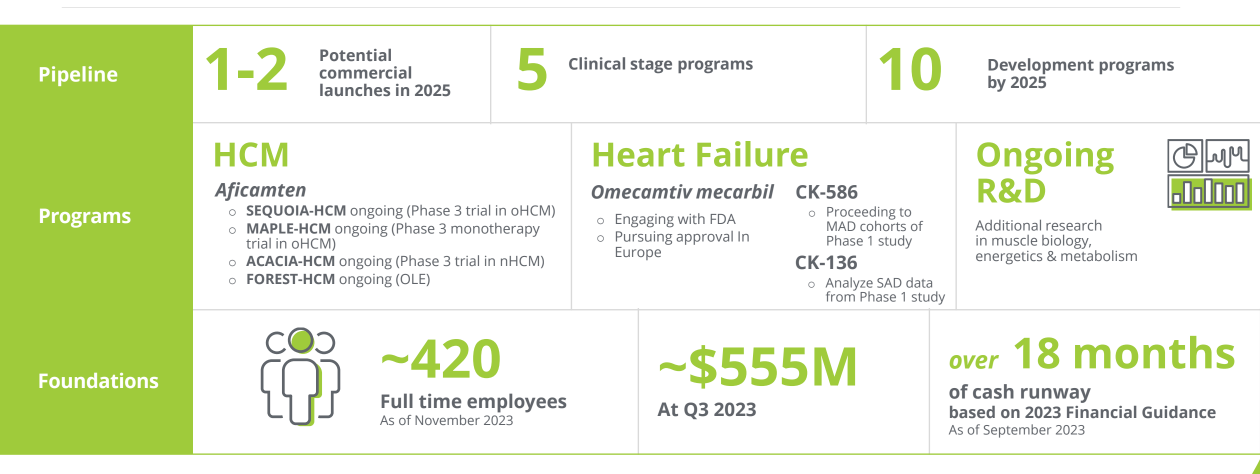
CK-136 is an investigational agent and has not been approved for use by any regulatory agency. Its safety and efficacy have not been established.

Sarcomere Directed Therapies

Corporate Profile



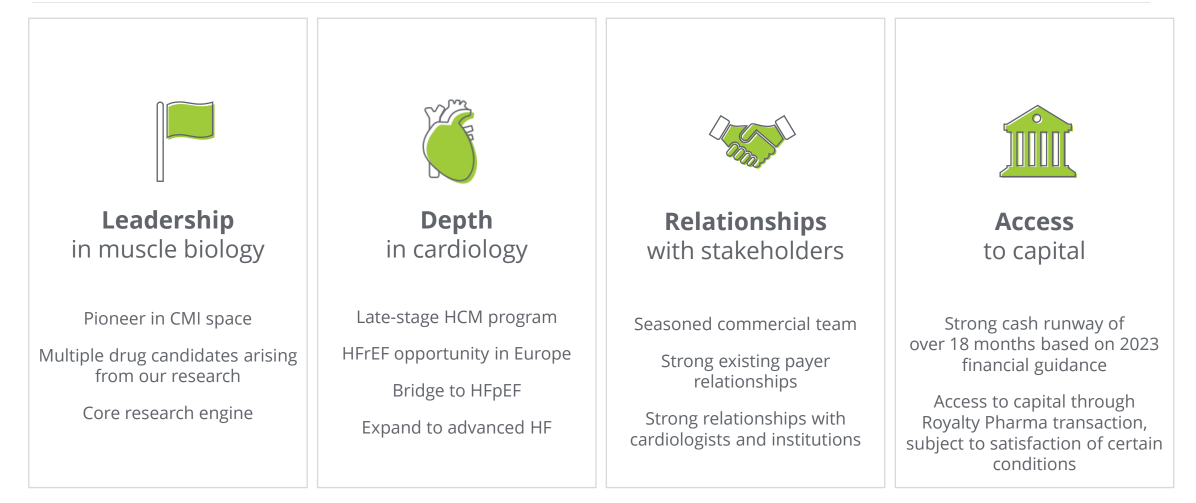
Robust Pipeline, Solid Financial Position



Timelines and milestones reflect Cytokinetics' current expectations and beliefs



Cytokinetics: Uniquely Positioned for Success



CMI: cardiac myosin inhibitor



Balance Sheet & Financial Guidance

Over 18 months of cash runway based on 2023 guidance

2023 Condensed Balance Sheet

As of 9/30/2023	in millions
	Total
Cash and investments	\$554.7
Accounts receivable	\$2.5
PPE	\$75.6
Leased assets	\$79.9
Other assets	\$27.9
Total Assets	\$740.6
Convertible Debt	\$545.0
Liability related to sale of future royalties	\$370.0
Lease liability	\$122.2
Other liabilities	\$142.2
Total Liabilities	\$1,179.4
Working capital	\$662.9
Accumulated deficit	(\$1,975.3)
Stockholders' deficit	(\$438.8)
Wtd Avg Basic Shares Outstanding (million)	96.1

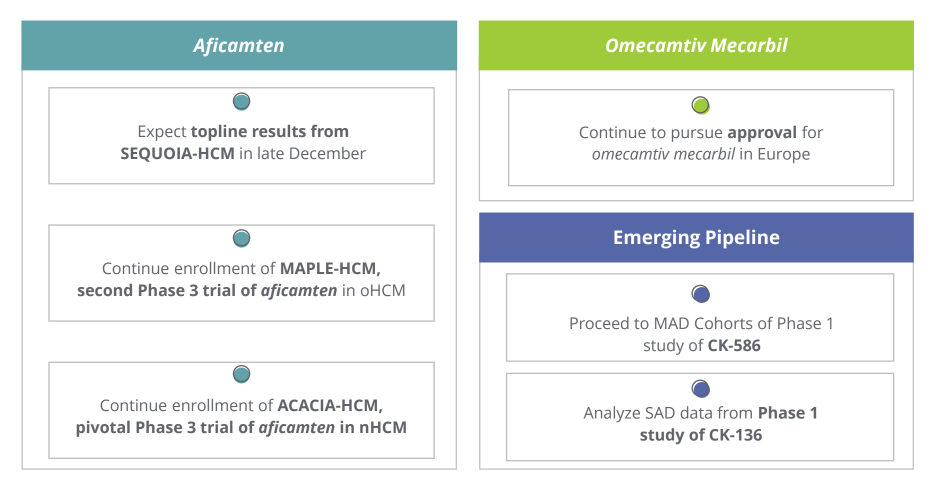
1. Cytokinetics internal planning data. Outside services spend for clinical trials, CMC and toxicology studies

2023 Financial Guidance

Net	~ \$310-320
Cash Operating Expenses	\$390-410
Cash Revenue	\$5
	Total
	In millions



Expected 2023 Milestones



Aficamten, omecamtiv mecarbil, CK-586 and CK-136 are investigational drugs and have not been approved. Their safety and efficacy have not been established.







Thank You

Sarcomere directed therapies



023 CYTOKINETICS, All Rights Reserved. "OKINETICS® and the C-shaped logo are registered trademarks of Cytokinetics in the U.S. and certain other countri ual patients who consented to use of their name, image, and condition.

