

Corporate Presentation

NASDAQ: CRIS



Cautionary Note Regarding Forward Looking Statements

This presentation contains certain forward-looking statements about Curis, Inc. ("we," "us," or the "Company") within the meaning of the Private Securities Litigation Reform Act of 1995, as amended. Words such as "expect(s)," "believe(s)," "may," "anticipate(s)," "focus(es)," "plans," "mission," "strategy," "potential," "estimate(s)", "intend," "project," "seek," "should," "would" and similar expressions are intended to identify forward-looking statements. Forward-looking statements are statements that are not historical facts, reflect management's expectations as of the date of this presentation, and involve important risks and uncertainties. Forward-looking statements herein include, but are not limited to, statements with respect to the timing and results of future clinical and pre-clinical milestones; the timing of future preclinical studies and clinical trials and results of these studies and trials; the clinical and therapeutic potential of our drug candidates; and management's ability to successfully achieve its goals. These forward-looking statements are based on our current expectations and may differ materially from actual results due to a variety of important factors including, without limitation, risks relating to: whether any of our drug candidates will advance further in the clinical development process and whether and when, if at all, they will receive approval from the U.S. Food and Drug Administration or equivalent foreign regulatory agencies; whether historical preclinical results will be predictive of future clinical trial results; whether historical clinical trial results will be predictive of future trial results; whether any of our drug candidate discovery and development efforts will be successful; whether any of our drug candidates will be successfully marketed if approved; our ability to achieve the benefits contemplated by our collaboration agreements; management's ability to successfully achieve its goals; the sufficiency of our cash resources; our ability to raise additional capital to fund our operations on terms acceptable to us or the use of proceeds of any offering of securities or other financing; general economic conditions; competition; and the other risk factors contained in our periodic and interim reports filed with the Securities and Exchange Commission which are available on the SEC website at www.sec.gov. You are cautioned not to place undue reliance on these forwardlooking statements that speak only as of the date hereof, and we do not undertake any obligation to revise and disseminate forward-looking statements to reflect events or circumstances after the date hereof, or to reflect the occurrence of or non-occurrence of any events, except as required by law.

This presentation is not an offer to sell or the solicitation of an offer to buy any securities, and shall not constitute an offer, solicitation, or sale in any jurisdiction in which such offer, solicitation, or sale is unlawful.

Curis Mission & Strategy

Developing the New Generation of Targeted Cancer Drugs



Company

Investment Thesis	Curis seeks to develop novel, first-in-class, cancer therapeutics that we believe have significant potential in areas of unmet patient need		
Robust Pipeline	 CA-4948: first-in-class suppressor of the TLR Pathway There are no drugs currently approved that block the entire TLR pathway CI-8993: first-in-class antagonist of the VISTA pathway There are no drugs currently approved for VISTA inhibition Fimepinostat: first-in-class suppressor of MYC There are no drugs currently approved for MYC inhibition 		
Corporate	 Experienced management team with proven capabilities Curis R&D pioneered the first-in-class inhibitor of the Hedgehog pathway (Erivedge[®]) partnered with and commercialized by Genentech/Roche for advanced basal cell carcinoma Cash, cash equivalents and investments of approximately \$20.5M as of Dec 31, 2019 		

Evolution of Curis



Progressing through Clinical Studies on the Path to Potential Registration

2018	2019	2020
Regulatory Planning	Clinical Execution	Expansion
Work with investigators and FDA to determine optimal clinical path Identify patient populations and initiate clinical studies	 Report preliminary efficacy data for CA-4948 Ph1 study in NHL Evaluate additional opportunities for CA-4948 in AML/MDS and work with investigators to design clinical studies 	 Continue dose escalation study of CA-4948 to determine MTD or RP2D and report updated efficacy data from Ph1 study Initiate new single-agent study of CA-4948 in AML/MDS patients w/ specific spliceosome mutations

Pipeline

Of Oncology Drug Candidates

		PRE-CLINICAL		CLIN	IICAL		MARKETED
	Indication	Proof of Principle	Safety	Dose Optimization	Clinical Activity	Pivotal	Commercial
Heme Malignancies							
CA-4948 * IRAK4	MYD88/TLR-altered NHL						
CA-4948 * IRAK4	IL-1R/TLR-altered AML, MDS						
Fimepinostat HDAC/PI3K	MYC-altered Cancers						
Immune Checkpoint	Inhibitors						
CI-8993 ** VISTA	VISTA-expressing Cancers						
CA-327 * PDL1/TIM3	PDL1/TIM3-expressing Cancers						
CA-170 * PDL1/VISTA	PDL1/VISTA-expressing Cancers						
Anaround Dava							
Erivedge*** Hedgehog	Basal Cell Carcinoma						

** Option to licensed from Murigene
 *** Option to license IP from ImmuNext
 Generatech **** IP licensed to Generatech (Curis receives royalty income)



Targeted Programs in Heme MalignanciesCA-4948: In development for treatment of TLR-altered cancers



CA-4948 Overview



In Development for Patients with MYD88/TLR-Altered Disease

	Profile			
Value Proposition	 First-in-class IRAK4 inhibitor in cancer Specific malignancies have overactivity of the myddosome/TLR pathway (which is dependent upon IRAK4) Composition-of-matter IP extends into 2035 	Designed to be best-in-class IRAK4 inhibitor ¹	Kinase IRAK4 IRAK1 IRAK2 IRAK3	Affinity K _d (nM) 23 12,000 >20,000 8,500
Population	Lymphoma: Patients w/ over-activated myddosome/TLR pathway; Patients treated w/ ibrutinib (IRAK4i has strong synergy with BTKi) Leukemia: Patients whose tumors have splicing mutations that overexpress IRAK4	Potent suppressor	Control 10 1.0	5 μM P-IKKα/β
Product Candidate Description	 Potent and orally bioavailable inhibitor of IRAK4 for treatment of MYD88-altered tumors and augmentation of BTK inhibition 	or signal transduction-	Phospho-protein lev after treatment w	P-ERK els in AML cells <i>r</i> ith CA-4948

CA-4948 Mechanism of Action

CURIS

In Development for Patients with MYD88/TLR-Altered Disease

Inhibiting either of these two pathways should provide benefit to patients with B cell lymphoma, CA-4948 targets oncogenic activity in the TLR pathway by blocking IRAK4



CA-4948 in Lymphoma



Preliminary Phase 1 Data Demonstrate Tolerability, PK, PD, and Anti-Cancer Activity



Patient Population

• Patients with R/R Lymphoma (incl DLBCL, WM, and patients with MYD88-altered disease)

Treatment

• Oral, once-daily (QD) or twice-daily (BID), dosing in continuous 21-day cycles

Objective

- Safety/tolerability during dose escalation
- Efficacy during expansion

Preliminary Phase 1 Data Readout

- Generally well tolerated
- Favorable PK profile, PD, and anti-cancer activity
- 5 of 6 patients dosed at 200mg or higher have seen reduction
- Report updated safety and efficacy data in 2020

CA-4948 in Lymphoma



Single Agent Dose Response Observed as Phase 1 Study Advances to Therapeutic Dose Levels



CA-4948 in R/R Lymphoma



Single Agent Dose Response Observed as Phase 1 Study Advances to Therapeutic Dose Levels



Dose Response Observed Increased tumor reduction observed as patient increased dose

CA-4948 in R/R Lymphoma



Combination with BTKi leads to Potent Anti-Cancer Activity in MYD88-altered DLBCL Models

Preclinical Anti-Cancer Activity

Potent as Single Agent

- Anti-cancer activity demonstrated in MYD88-altered DLBCL
- Interim readout in dose escalation study shows clear dose response
- We intend to continue dose escalation to further enhance efficacy until MTD / R2PD

Strong Synergy in Combination

- Anti-cancer activity demonstrated to be highly synergistic with BTK inhibition
- Evaluating potential clinical development strategies for CA-4948 with BTKi to assess anti-cancer activity

Preclinical Anti-Cancer Activity in MYD88-altered DLBCL¹ (OCI-Ly10)



1) Data from Curis preclinical study; Booher, et al. 4th Waldenstrom Roadmap Symposium

CA-4948 in AML/MDS

Additional Regulatory Path Identified in AML/MDS

Specific Genetic Mutations Lead to Expression of the Long Isoform of IRAK4 (IRAK4-L)

Spliceosome mutations (incl. SF3B1 and U2AF1) drive expression of IRAK4-L in >50% of AML/MDS



RIS

CA-4948 in AML/MDS



Overexpression of IRAK4-L is a Strong Predictor of Sensitivity to CA-4948







Monoclonal Antibody Immune Checkpoint Inhibitor

CI-8993: In development for treatment of VISTA-expressing cancers



CI-8993 Overview



In Development for Patients with VISTA Expressing Cancers

Profile				
Value Proposition	 First-in-class monoclonal antibody antagonist of VISTA No anti-VISTA drugs currently in the clinic Composition-of-matter IP extends into 2034 			
Population	 Patients with VISTA-expressing cancers, including Mesothelioma, NSCLC, and TNBC All patients who will receive PD1/PDL1 or CTLA4 antibody therapy Patients who have developed resistance to PD1/PDL1 or CTLA4 antibody therapy 			
Product Description	 Monoclonal antibody developed by ImmuNext/Janssen in partnership with Randy Noelle's lab at Dartmouth (the co-discoverer of VISTA) 			

CI-8993

Illustrative Example of VISTA's Role in Immune Suppression





Targeted Programs in Heme Malignancies

Fimepinostat: In development for treatment of MYC-altered cancers

Fimepinostat Overview

In Development for Patients with MYC-Altered Cancers

Profile			
Value Proposition	 First-in-class drug candidate with demonstrated anti-cancer activity as a single agent in MYC-altered patients in Ph1 and Ph2 trials Composition-of-matter IP extends into 2032 		
Population	 Patients with MYC-altered cancer (>50% of all cancers are effected by MYC)³ Collaborating with DarwinHealth on characterization of biomarkers and tumor type alignments to identify potential therapeutic opportunities 		
Product Candidate Description	 Potent and orally bioavailable dual inhibitor of HDAC and PI3K enzymes¹ Favorable safety profile in over 200 patients 		



Protein levels in DLBCL cells after treatment with Fimepinostat (Curis Preclinical Study)

Tubulin

Qian et.al. Clin Cancer Res. 2012. 18: 4104
 Sun et.al. Mol Cancer Ther. 2017. 6: 285
 Chen et al. Nature. 2018 Feb 23. 3:5

Fimepinostat Clinical Data



Provides Strong Rationale For Development in MYC-Altered Lymphoma



Monotherapy Anti-Cancer Activity

Deep responses

- 8 complete responses (CR); 6 partial responses (PR)
- 2 patients able to proceed to transplant

Durable responses

• Median duration = 13.6 months

Fast Track designation received

• Following FDA review of clinical data

Evaluating potential therapeutic opportunities for fimepinostat in research collaboration with DarwinHealth



Investment Thesis	Curis seeks to develop novel, first-in-class, cancer therapeutics that we believe have significant potential in areas of unmet patient need		
Robust Pipeline	CA-4948: CI-8993: Fimepinostat:	first-in-class suppressor of the TLR Pathway There are no drugs currently approved that block the entire TLR pathway first-in-class antagonist of the VISTA pathway There are no drugs currently approved for VISTA inhibition first-in-class suppressor of MYC There are no drugs currently approved for MYC inhibition	
Potential Upcoming Catalysts	 Updated efficacy data for CA-4948 Phase 1 study Initiation of CA-4948 Phase 1 study in AML/MDS patients with spliceosome mutations Initiation of CI-8993 dose escalation Phase 1 study 		



Leadership Team







Mark Noel Head, Intellectual Property



Reinhard von Roemling Head, Clinical Development



Raul Soikes Head, Portfolio Management



Nancy Soohoo General Counsel



William Steinkrauss Chief Financial Officer

CURIS



End of Corporate Presentation

NASDAQ: CRIS

