



Conditionally-Active Antibody Therapeutics for the treatment of cancer

NASDAQ: CTMX

CORPORATE OVERVIEW | JANUARY 2021

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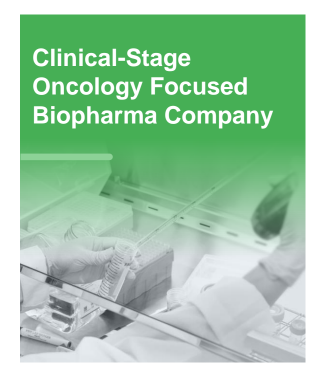
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Company Snapshot



Conditionally-Active Antibodies

- Innovative targeting strategy
- Leverages tumor microenvironment
- Opens previously undruggable target space
- Leaders in field

Foundational Partnerships

- AbbVie, Amgen, Astellas & BMS
- Retained certain US rights

Key 2021 Milestones

- CX-2009 initial Phase 2 data in breast cancer
- CX-2029 initial Phase 2 expansion cohort data
- Next IND filings

Strong Balance Sheet

- \$321M cash at end of Q3 2020
- No debt



Experienced Leadership



Sean A. McCarthy, D. Phil.

President, Chief Executive Officer and Chairman
>20 years of experience in biotech with roles in R&D, business development, financing and general management



Amy C. Peterson, M.D.
EVP, Chief Development Officer
>15 years of leadership experience in oncology drug
development



Alison L. Hannah, M.D.
SVP, Chief Medical Officer
>30 years of experience in investigational cancer therapy development



Carlos Campoy SVP, Chief Financial Officer >30 years of financial and leadership experience, mostly with publicly-held healthcare and biopharmaceutical companies



Marcia P. Belvin, Ph.D.

SVP, Head of Research
>20 years of experience in preclinical pipeline discovery and development in oncology









































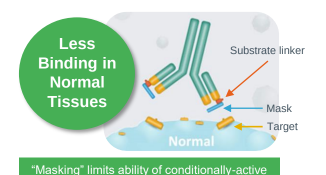


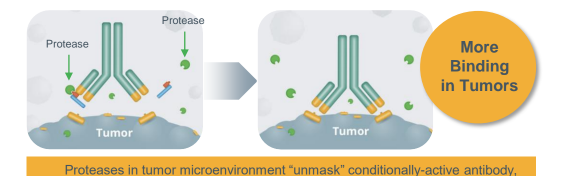
Broad Clinical and Preclinical Pipeline with Multiple Phase 2 Readouts 2021+

	PRODUCT CANDIDATE	PROBODY TARGET	INDICATION	IND-ENABLING	PHASE 1	PHASE 2	COMMERCIAL RIGHTS	
CONDITIONAL ADCs	CX-2009	CD166-DM4	Breast Cancer	Arm A: monotherapy in advanced, metastatic HR+/HER2 non-amplified BC Arm B: monotherapy in advanced, metastatic TNBC Expected Arm C: in combination with CX-072 in advanced, metastatic TNBC Q4 2021		CYTOMX		
	CX-2029	CD71-MMAE	Multiple Cohorts	Cohort 1: sqNSCLC Cohort 2: HNSCC Cohort 3: Esophageal cancer Cohort 4: DLBCL Initial Data Expected Q4 2021			CYTOMX abbvie	
	CX-2043	EpCAM- DM21	Solid Tumors	Target IND 2021			CYTOMX	
IMMUNO- ONCOLOGY	BMS-986249	CTLA-4	1L Melanoma	Randomized: + nivolumab vs.	ipilimumab + nivolumab vs. nivol	umab	ر ^{الا} Bristol Myers Squibb ٔ	
	BMS-986288	CTLA-4 a-Fucosylated	Solid Tumors	Dose escalation: +/- nivolumab			C 2	
	CX-904	EGFR + CD3 T-Cell Bispecific	ТВА	Target IND 2021			CYTOMX AMGEN	



The Probody[®] Therapeutic Platform: Conditionally-Active Antibodies



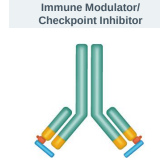


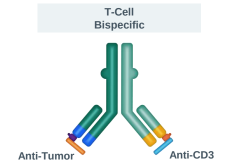
allowing more binding to tumor cells

Antibody-Drug Conjugate

Linker-payload

antibody to bind to healthy tissues







Antibody-Drug Conjugates for Cancer are a Major Opportunity

Recent Approvals and Transactions Underscore High Potential of Class





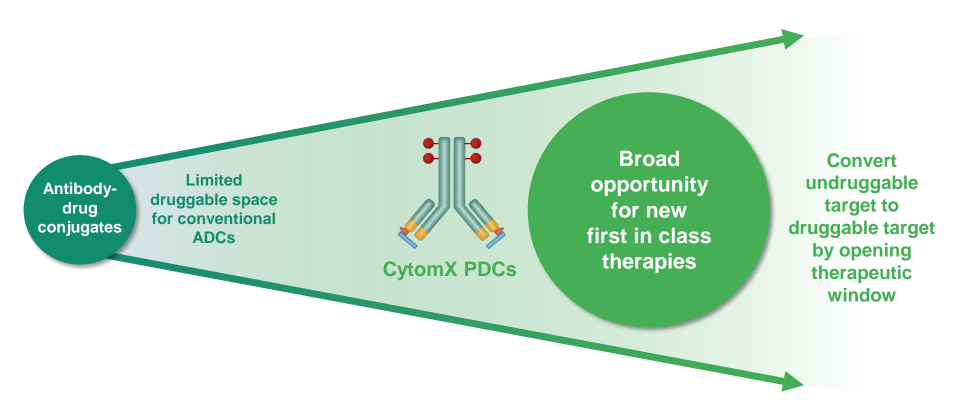








Probody Drug Conjugates Expand ADC Target Landscape







Substantial Unmet Need Remains in Breast Cancer

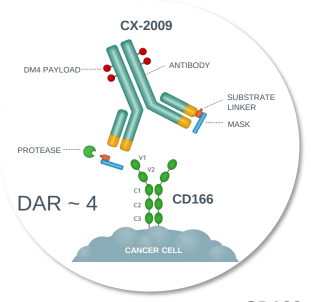


Breast cancer is the 2nd leading cause of cancer deaths in women¹

- ~80% of breast cancer is HER2 non-amplified
- Despite recent advances, new therapies are needed, especially in the metastatic setting
- CD166 is broadly and highly expressed in HER2 non-amplified breast cancer



CX-2009: A Probody Drug Conjugate Targeting CD166 (ALCAM*)



- CD166 expression in normal cells limits development of a conventional ADC (e.g., Lung, GI tissues, Liver)
- CX-2009 is a first-in-class anti-CD166 Probody conjugated to the maytansinoid cytotoxic payload DM4
- Designed to target CD166 towards tumor tissue, away from healthy tissue
- CD166 expressed on many other cancer types → future opportunity (e.g., Ovarian, Lung, HNSCC)

CD166 Expression by IHC

Breast Cancer



Lung Cancer



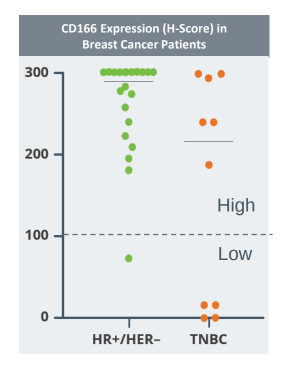
Ovarian Cancer





Phase 1 Enrolled 39 Patients with Breast Cancer at Doses 0.25-10 mg/kg

	Overall (n=39)	HR+/HER2- (n=28)	TNBC (n=11)
Median age, years (range)	53 (31-77)	54 (37-77)	45 (31-68)
White/Asian/Hawaiian/Unk/Other, n	30/1/1/5/2	21/0/1/5/1	9/1/0/0/1
ECOG PS 0/1	17/22	12/16	5/6
Median no. of prior regimens, (range)	7.5 (3-16)	8 (4-16)	7 (3-11)
Prior platinum, n (%)	15 (38.5%)	6 (21.4%)	9 (81.8%)
Prior microtubule inhibitor, n (%)	37 (94.9%)	26 (92.9%)	11 (100%)
Prior CDK4/6 inhibitor, n (%)	17 (43.6%)	17 (60.7%)	0
Prior anti-PD-I or PD-L1, n (%)	6 (15.4%)	2 (7.1%)	4 (36.4%)
CD166 High/Low/Unknown	32/5/2	26/1/1	6/4/1
Median no. of CX-2009 doses (range)	2 (1-25)	2.5 (1-25)	2 (1-14)



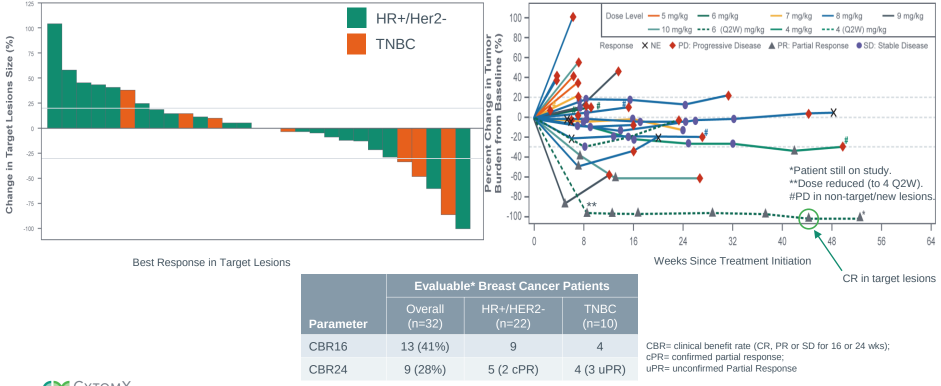
HR+/HER2- : Hormone Receptor positive and HER2 non-amplified breast cancer;

TNBC: Triple negative breast cancer



Observed Clinical Activity in Breast Cancer with CX-2009 at Doses ≥4 mg/kg Q3W

Breast cancer patients with measurable disease who received ≥ 4 mg/kg CX-2009 and had a post-baseline assessment

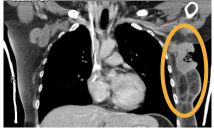


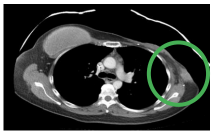
*Includes those with non-measurable but evaluable (e.g. bone-only) disease

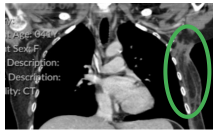
Partial Response to CX-2009 in Patient with TNBC Refractory to Pembrolizumab+Paclitaxel and to Sacituzumab Govitecan

BASELINE













- 41-year-old treated at 8 mg/kg
- Prior treatment for metastatic disease:
 - Pembrolizumab + paclitaxel (best response = PD)
 - Sacituzumab govitecan (best response = PD)
- Baseline: ulcerating skin lesions on chest wall and axillary nodal metastasis
- First scan (Week 8): 48% reduction in target lesions
- Dose interruption (week 9 -16) for keratitis (resolved), disease progressed before treatment could be re-initiated



CX-2009: Phase 1 Tolerability Supports Phase 2 Dose of 7mg/kg

		RP2D			
	< 6 mg/kg (n=38)	7 mg/kg (n=12)	8 mg/kg (n=22)	9 mg/kg (n=9)	10 mg/kg (n=8)
TRAE (Grade 3+)	16%	33%	64%	56%	50%
TEAE leading to Discontinuation	13%	8%	14%	22%	13%
DLT (n)	0	0	1	0	0
TR SAEs	0	17%	27%	22%	13%
Ocular Toxicity (any grade)*	26%	25%	59%	56%	75%
Ocular Toxicity (Grade 3+)	3%	0	14%	33%	13%

CX-2009 was generally well tolerated at doses ≤ 7 mg/kg (toxicity profile consistent with payload: ocular, neuropathic and hepatic)

Ocular toxicities appeared dose dependent in frequency and severity

Selection of 7 mg/kg Q3W as RP2D is supported by activity, tolerability and PK/PD modeling

RP2D= Recommended Phase 2 Dose

Data presented SABCS 2020 Data cut-off: August 2020 15

^{*}Ocular prophylaxis was optional; future studies will incorporate mandatory ocular prophylaxis

CX-2009 Breast Cancer Phase 2 Study Design

Monotherapy (7mg/kg Q3W) and Combination with Pacmilimab (CX-072; anti-PD-L1) In Advanced, HER2 non-Amplified Breast Cancer

Key Eligibility

Ocular prophylaxis required

HR+/HER2 non-amplified

- 0 2 prior cytotoxics for advanced disease
- Measurable disease required
- No active corneal disease

TNBC

- CD166 High
- ≥ 1 and ≤ 3 priors for advanced disease
- · Measurable disease required
- Treated/stable brain metastases allowed
- No active corneal disease
- Arm C exclusion criteria:
 - PD-L1 negative/unknown
 - I/O refractory
 - History of or active autoimmune condition

Breast Cancer SubType

Arm A

HR+/HER2 non-amp (n~40*) CX-2009

Arm B

TNBC (n~40*) CX-2009

Arm C

TNBC (n~40*) CX-2009 + CX-072**

Endpoints

Primary: Overall Response Rate (ORR) by central review

Secondary: ORR (Inv), PFS, DCR, CBR24, DoR, OS, Safety, PK, ADA

Exploratory: Biomarker correlation with outcome

Readout: Initial data expected Q4 2021

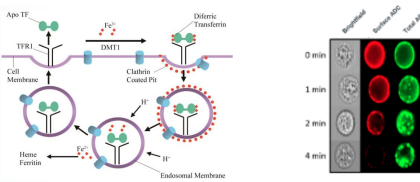




CD71 (Transferrin Receptor)



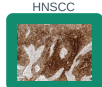
- Highly expressed tumor antigen
- "Professional internalizer" ideally suited to delivery of cytotoxic payloads to cancer cells
- Undruggable target with conventional antibody approaches due to normal tissue biology
- Probody strategy open therapeutic window by limiting normal tissue binding
- Potentially paradigm shifting anti-cancer agent with first in class potential

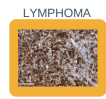


Elliott and Head. J Cancer Ther. 2012;3:278-311.







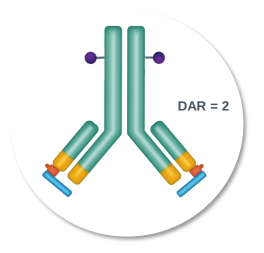


CD71 Expression by IHC

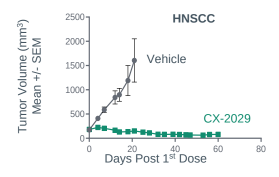


CX-2029: Potentially Paradigm Shifting Anti-Cancer Agent





- Unmasked ADC is lethal in preclinical models at subtherapeutic doses
- Therapeutic range for CX-2029 conditional ADC predicted in patients 2-4mg/kg
- Hematologic toxicity dose limiting in preclinical studies





Phase 1 Dose Escalation Study Evaluated CX-2029 Q3W in 45 Patients with Solid Tumors





Key Eligibility Criteria

- · Metastatic or locally advanced unresectable solid tumor
- · Archival tissue or biopsy available for tissue analyses
- · Stable brain metastases permitted

Exclusions:

- Transfusion-dependent anemia or iron metabolism disorders
- · Grade 2 or higher neuropathy

K D :	AII O 1 4
Key Patient Demographics	All Cohorts (n=45)
Age, median (min, max)	60 (31, 75)
Baseline ECOG 0 / 1, %	29 / 71
CD71 IHC staining, n (%) High expression [2+/3+] Low expression [0/1+] Unknown	15 (33) 16 (36) 14 (31)
Tumor types, n (%) NSCLC Squamous NSCLC HNSCC Colorectal cancer Other*	9 (20) 4 (9) 8 (18) 7 (16) 21 (46)
Median priors (min, max)	3 (1, 16)

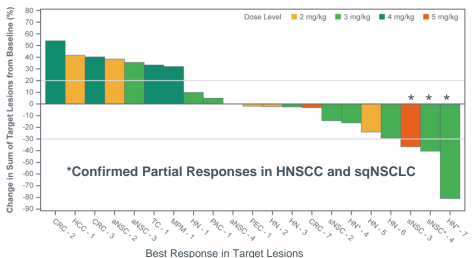
*Other tumor types include sarcoma (4), Prostate (3), parotid gland (3); ovarian (2); melanoma (n=1); endometrial (1); hepatocellular (1); mesothelioma (1); ocular melanoma (1); pancreatic (1); perivascular epithelioid (1); thymoma (1); thyroid (1).



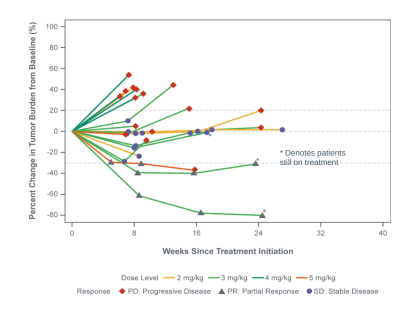
Observed Clinical Activity with CX-2029 at Doses ≥2 mg/kg Q 3 Weeks



Patients with measurable disease who received ≥ 2 mg/kg CX-2029 and had a post-baseline assessment



CRC=Colorectal Cancer, HCC=Hepatocellular carcinoma, aNSC=Non-small cell lung adenocarcinoma, TC=Thyroid carcinoma MPM=Malignant pleural mesothelioma, HN=Head and neck squamous cell carcinoma, PAC=Pancreatic cancer, PEC=Perivascular epithelioid cell tumor, sNSC=Non-small cell lung squamous carcinoma



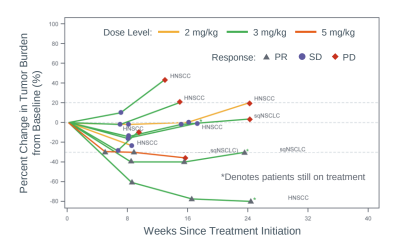


Observed Clinical Activity in sqNSCLC and HNSCC with CX-2029 at Doses ≥2 mg/kg Q 3 Weeks



sqNSCLC or HNSCC patients with measurable disease, received ≥ 2 mg/kg CX-2029, and had a post-baseline assessment





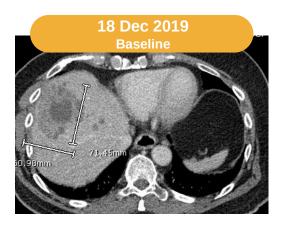
1 patient with sqNSCLC was dosed at 1mg/kg; 1 patient with HNSCC came off study without a post-baseline assessment

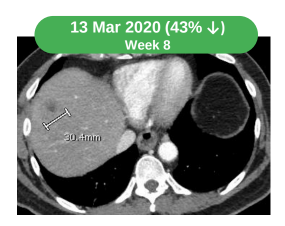


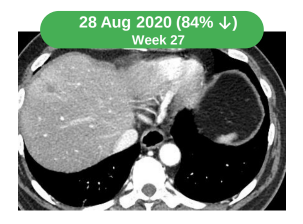
CX-2029 Case Study: Patient with HNSCC



- Nasopharyngeal carcinoma (Diagnosed in February 2018)
- Prior therapies: docetaxel/5FU/cisplatin with radiation; high-dose cisplatin; investigational agent (sEphB4-HSA)
 + pembrolizumab (best response was PD)
- CX-2029 treatment initiated (January 2020)
- Partial response at Week 8 confirmed 8 weeks later. Dose reduced to 2 mg/kg; additional shrinkage of liver target lesion seen.









CX-2029 Phase 1 Tolerability Supports Phase 2 Dose of 3mg/kg



Generally Well Tolerated to 3 mg/kg with Manageable Adverse Events

	RP2D				
Treatment-Related Grade 3+ AEs (≥2 patients)	1.0 mg/kg (n=3)	2.0 mg/kg (n=8)	3.0 mg/kg (n=12)	4.0 mg/kg (n=6)	5.0 mg/kg (n=4)
Anemia	33%	63%	58%	83%	100%
Neutropenia	0	0	33%	50%	75%
Leukopenia	0	0	8%	33%	50%
Infusion-related reaction	0	13%	0	17%	0

- > 90% masking maintained in circulation
- Most frequent Grade 3+ AE was anemia
 - Managed with red blood cell transfusions, growth factor support and/or dose delays/reductions
 - Likely multi-factorial including CD71 biology and MMAE payload
- 3 mg/kg Q3W selected as Phase 2 dose



Phase 2 Expansion Underway to Evaluate CX-2029 in Four Cohorts



Monotherapy at 3mg/kg Q3W

Eligibility

sqNSCLC, HNSCC and esophageal

- Prior therapy must include prior platinum and a checkpoint inhibitor (alone or in combination; if approved by the local Health Authority).
- For esophageal: squamous, adenocarcinoma or GE junction; prior HER2-targeted therapy if tumor is HER2+
- Documented progression after at least one prior regimen for advanced disease

DLBCL

 Progression after at least 2 prior regimens (one of which must be anti-CD20 based therapy); not a candidate for stem cell transplant

Cancer Type

sqNSCLC

n~25*

HNSCC

n~25*

Esophageal/GEJ

n~25*

DLBCL

n~25*

*Evaluable

Endpoints

Primary: Overall Response Rate (ORR) by local investigator

Secondary: PFS, DCR, CBR24, DoR, OS, Safety, PK, ADA, TTR

Exploratory: Biomarker correlation with outcome

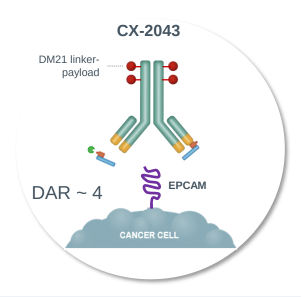
Readout: Initial data expected Q4 2021







CX-2043: Conditional ADC Targeting EpCAM/TROP-1

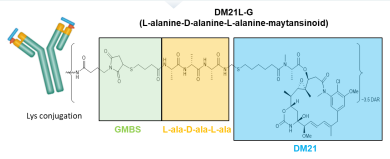


- CX-2043 generated in collaboration with Immunogen
- CytomX retains WW development and commercial rights

Target Background

- Epithelial cell marker; Highly expressed on solid tumors
- EpCAM-targeted therapies can be active when delivered locally
- On-target / off-tumor toxicities limit systemic delivery

CX-2043: EpCAM-targeting PDC



- Next-generation linker-payload system with enhanced stability and improved bystander activity
- Probody platform alleviates on-target / off-tumor toxicity (pancreatitis, GI tox)



Presented at EORTC-NCI-AACR 2020



Strong Alliances Advancing Multiple Programs and Probody Formats





CHECKPOINT INHIBITORS

LEAD PROGRAMS: Expanding Therapeutic Window for CTLA-4

BMS-986249 ipilimumab Probody in melanoma Phase 2

BMS-986288 non-fucosylated ipilimumab Probody in Phase 1





PROBODY DRUG CONJUGATES

LEAD PROGRAM: CD71 (CX-2029)

Global co-development alliance

CytomX retained US rights (35%) and >20% royalties ex-US





T-CELL BISPECIFICS

LEAD PROGRAM: CX-904

EGFR-CD3 conditional T-Cell bispecific

IND enabling studies for potential 2021 IND





T-CELL BISPECIFICS

Conditional T-Cell Bispecifics

Alliance formed March 2020

\$80 million upfront



Strong Balance Sheet to Support Pipeline and Operations



\$321M in cash as of Sept. 30th 2020



\$130M of non-dilutive capital in 2020

- > \$80M upfront from Astellas
- > \$40M CX-2029 milestone from AbbVie
- > \$10M anti-CTLA-4 milestone from BMS



No debt



46.2M shares outstanding



Leadership in Conditionally-Active Antibodies with Validated Platform

Summary

- Versatile, multi-modality platform
- Five clinical stage assets
- 2 conditional ADCs in Phase 2
 - CX-2009, CX-2029
- 2 conditional checkpoint inhibitors in Phase 2
 - CX-072 (+ CX-2009)
 - BMS-986249
- Emerging T-cell bispecifics
- Robust platform and preclinical pipeline
- Strong alliances

2021 **Priorities**

- Patient enrollment into CX-2009 Ph 2 study
 - HR+/HER2 non-amplified breast cancer
 - TNBC +/- CX-072
 - Initial data expected Q4 2021
- Patient enrollment into CX-2029 Ph 2 expansions
 - sqNSCLC, HNSCC, esophageal, DLBCL
 - Initial data expected Q4 2021
- IND submission
 - CX-2043
 - CX-904
- Continued progress within partnerships



CytomX Therapeutics Inc.

Our VISION

Our **PLATFORM**

Our **PRODUCTS**

Our **TOMORROW**



Create

a new approach to the treatment of cancer by improved tumor targeting



Lead

in conditional activation of antibody-drug conjugates and other modalities



Advance

a broad clinical pipeline of anti-cancer therapies in areas of significant unmet need



Build

a long-term, commercial stage, multi-product enterprise







Conditionally-Active Antibody Therapeutics for the treatment of cancer

NASDAQ: CTMX

CORPORATE OVERVIEW | JANUARY 2021

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