UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

Form 8-K

Current Report

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): September 13, 2016

CYTORI THERAPEUTICS, INC.

(Exact name of registrant as specified in its charter)

Delaware

(State or Other Jurisdiction of Incorporation)

001-34375 (Commission File Number) 33-0827593

(I.R.S. Employer Identification Number)

3020 Callan Road, San Diego, California 92121 (Address of principal executive offices, with zip code)

(858) 458-0900

(Registrant's telephone number, including area code)

n/a

(Former name or former address, if changed since last report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions (*see* General Instruction A.2. below):

□ Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

□ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

Dere-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

Dere-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Item 7.01 Regulation FD Disclosure

A copy of an investor slide presentation that Cytori Therapeutics, Inc. (the "Company") will use during a presentation at the 18th Annual Rodman and Renshaw Global Investment Conference on Tuesday, September 13, 2016 at 10:00AM Eastern Time (7:00 AM Pacific Time) at the Lotte New York Palace Hotel in New York, is attached to this Current Report on Form 8-K ("Current Report") as Exhibit 99.1 and is incorporated by reference herein. Additionally, the Company has posted the slide presentation on the Company's Investor Relations website at http://ir.cytori.com and maintains the current version of its corporate presentation at such website.

The information contained in this Item 7.01 and Exhibit 99.1 hereto shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as shall be expressly set forth by reference in such a filing.

Item 9.01. Financial Statements and Exhibits.

(d) Exhibits.

99.1 Investor Presentation Material

WEST\253929622.1

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the Registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

CYTORI THERAPEUTICS, INC.

/s/ Jeremy Hayden Name: Jeremy Hayden Title: General Counsel and VP of Business Development

Date: September 13, 2016

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No. 99.1

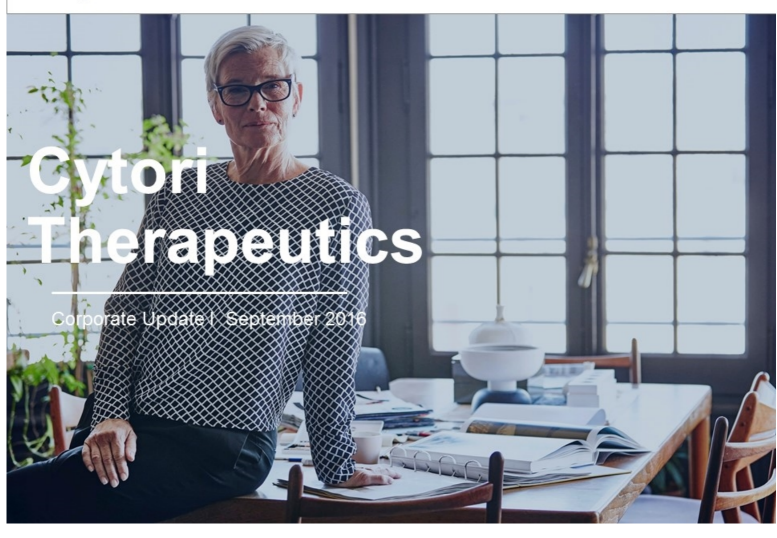
I Investor Presentation Material

Description

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Scytori

Exhibit 99.1 Enhancing lives through novel cell therapies



Forward Looking Statements and Disclaimers

This presentation contains certain 'forward-looking statements' about Cytori Therapeutics, Inc. All statements, other than statements of historical fact, that address activities, events or developments that we intend, expect, project, believe or anticipate will or may occur in the future are forward-looking statements. Such statements are based upon certain assumptions and assessments made by our management in light of their experience and their perception of historical trends, current conditions, expected future developments and other factors they believe to be appropriate.

The forward-looking statements included in this presentation, involve known and unknown risks that relate to future events or our future financial performance and the actual results could differ materially from those discussed in this presentation. Some of those forward-looking statements include statements regarding: our financial condition and prospects; our commercialized and pipeline products and technologies; the timing and conduct of our clinical trials and other parties' clinical trials involving Cytori Cell Therapy, including associated financial, clinical and regulatory burdens and projected timing for trial approval, enrollment and completion; the various medical indications and markets that may be addressed by Cytori Cell Therapy; the potential effectiveness of Cytori Cell Therapy, including clinical outcomes; conduct of our European managed access program; anticipated uses of clinical trial data; regulatory, reimbursement and commercial strategies and pathways; potential costs and other adverse effects of diseases targeted for treatment by our products, including the Celution system, and; anticipated future funding and contract revenues. Some risks and uncertainties related to such forward looking statements include risks and uncertainties regarding: the funding, conduct and completion of our clinical trials and other parties' clinical trials involving Cytori Cell Therapy; our ability to successfully execute our managed access program; uncertain clinical outcomes; regulatory uncertainties (including potentially adverse decisions regarding our existing and expected regulatory registrations, approvals and authorizations), unfavorable reimbursement outcomes; inability to access sufficient capital on acceptable terms (including inability to fund, or find third party sources to fund, our proposed clinical trials or continued development of our technologies), failure to maintain our substantially reduced cash burn; failure to achieve projected product revenue and contract revenue growth; our and our partners' failure to launch products and grow revenues in markets where we currently forecast sales; our abilities to service, pay and/or refinance our corporate debt; availability of future government funding and changes in government procurement priorities; the U.S. federal government's ability to reduce, modify or terminate the BARDA contract if it determines it is in its best interests to do so; increasing or unanticipated competitive pressures; potential performance issues with our products and technologies; lack of customer acceptance of our technologies; inability to find commercial partners for our therapies; and other risks and uncertainties described under the "Risk Factors" section in our Securities and Exchange Commission Filings on Form 10-K and Form 10-Q. These risks and uncertainties may cause our actual results to differ materially from those discussed in this presentation. We advise reading our most recent annual report on Form 10-K and quarterly reports on Form 10-Q filed with the U.S. Securities and Exchange Commission for a more detailed description of these risks.

The forward-looking statements contained in this presentation represent our estimates and assumptions only as of the date of this presentation and we undertake no duty or obligation to update or revise publicly any forward-looking statements contained in this presentation as a result of new information, future events or changes in our expectations. **Caution: Within the U.S., the Celution System is an investigational device limited by U.S. law to investigational use.** The following trademarks are owned by Cytori Therapeutics: Celase, Celution, Celution (with design), Cytori Therapeutics, Cytori (with design) and Cytori Cell Therapy. All third party trademarks are the property of their respective owners.

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Cytori Overview

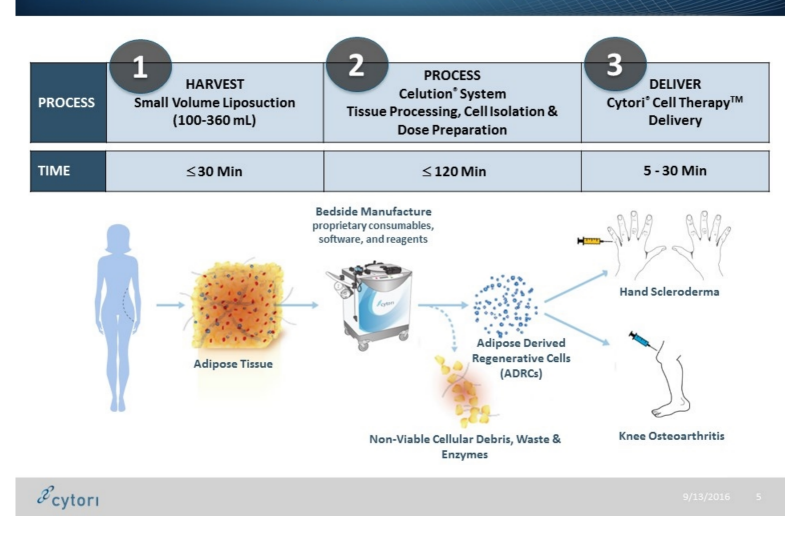
- Late stage cell therapy company, forecasted breakeven 2018
- Three phase III trials complete or enrolling
- Fully enrolled US phase III scleroderma trial, data 2017
- Recent US Phase II trial readout in knee osteoarthritis
- Up to \$106m US government (BARDA) contract, anticipate clinical trial milestone 2017
- Product revenue growth & contracting revenue- narrowing burn



Cytori Cell Therapy: Clinical Pipeline

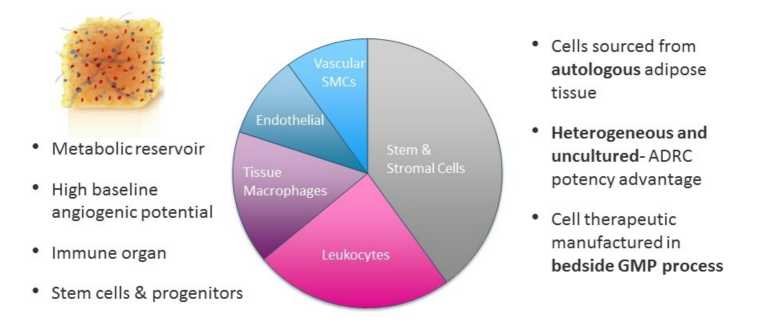
Therapeutic	Pre-Clinical	Phase I/II	Phase III		Market (Estimate)	
Scleroder	ma Associated Hand Dysfo	unction				
ECCS-50			Enrollment Complete		>\$1B	
ECCS-50			Enrolling ¹		>\$500M	
Knee Oste	oarthritis					
ECCO-50	Τορ	p-line preliminary results			>\$3B	
Urinary In	continence					
ECCI-50			Enrolling ²		>\$75M	
Cutaneou	s Radiation & Thermal Inj	ury				
DCCT-10	Preclinical ³				>\$50M	
¹ Cytori-supported, Investigator-ini ² Japan Govt Sponsorship ³ Funded by BARDA (US Govt.)						
Pcytori						
DCCT-10 ¹ Cytori-supported, Investigator-ini ¹ Japan Govt Sponsorship	Preclinical ³	ury				6

Cytori Cell Therapy: Same Day Procedure



Cytori Cell Therapy: Why Adipose?

Adipose-derived regenerative cells- Clinical grade, heterogeneous cell population highly-enriched for adipose-derived stem, stromal, vascular, and immunoregulatory cell types

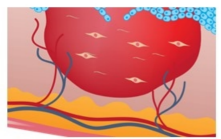


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Cytori Cell Therapy: Mechanism of Action

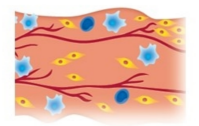
Cytori Cell Therapy is being developed with the goal of beneficially modulating multiple key pathologic processes which are anticipated to reduce pain and disability and improve quality of life

Angiogenesis/Vasculopathy



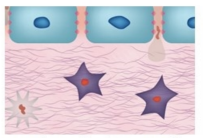
- Promotes angiogenesis
- Normalization of vessel architecture
- Improved vasomotor function¹⁻⁵

Inflammation



- Modulates expression of proand anti-inflammatory factors
- Modulates the function of proand anti-inflammatory cells^{3, 6-9,}

Fibrosis/Wound Remodeling



- Reduces development of fibrosis
- Remodels existing fibrosis^{2,10,11}

1. Foubert et al (2015); 2. Koh et al (2011); 3. Premaratne (2011); 4. Morris et al (2015); 5. Eguchi et al (2015); 6. Feng et al (2010); 7. Hao et al (2014); 8. Dong et al (2013); 9. Data on file (Cytori); 10. Serratrice et al (2014); 11. Data on file (Cytori)



Lead Indication: Scleroderma



Scleroderma

Scleroderma or Systemic Sclerosis

- Rare autoimmune condition
- Affects Women: Men, 4:1
- US Prevalence: 50,000 patients
- >90% of patients have hand disability
 - Fibrosis, pain, and edema result in diminished mobility and hand function even with standard medical care
 - Severe vasomotor symptoms

Pathophysiology





Raynaud's Phenomenon

Ulceration and Edema



Cytori Cell Therapy

Preclinical and in vitro studies report modulation of perivascular inflammation, improved endothelial function, and reduction of extracellular matrix (fibrosis)

Images reproduced with permission of the nonprofit International Scleroderma Network at sclero.org Image on left by D Niklas, https://commons.wikimedia.org/wiki/File:Raynaud-Syndrom.JPG used under CC license Image on right reproduced with permission of the nonprofit International Scleroderma Network at sclero.org

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Scleroderma: Market Overview

Current Standard of Care

- No therapies approved for treatment of hand dysfunction in scleroderma patients ٠
- Existing 1st and 2nd line treatments for treatment of Raynaud's Phenomenon or other aspects of scleroderma are often inadequate and/or poorly tolerated
- Existing 3rd line treatments are costly (\$30-\$100k) and often very poorly tolerated •

Diagnosis • Average age: 30's-50's	<u>1st/2nd Line Therapies</u> Inadequately effective and/or poorly tolerated in ~50% of patients ^{1,2}	<u>3rd Line Therapies</u> Expensive, often poorly-tolerated doses titrated to tolerance rather than to symptom relief	
	 Calcium channel blockers (eg: nifedipine) PDE5 inhibitors (eg: sildenafil) Topical nitrates Side effects: headache, dizziness, flushing, tachycardia, and edema 	 Endothelin-1 receptor antagonist (eg: Bosentan) Intravenous (IV) prostaglandin (PG) analog (eg: Iloprost) Pain due to severe ischemia may require the use of analgesics Immunosuppressive agents (eg: methotrexate, cyclophosphamide, azathioprine, mycophenolate) Surgical sympathectomy 	

Scleroderma: Treatment Approach

- Ambulatory
- Procedure room
- Local or mild conscious sedation
- Single administration ECCS-50
- 0.5cc injection to each side of each finger





Pilot/Phase I SCLERADEC I Trial

	SCLERADEC I	
Study size	12	
Randomization	Open label	
Administration	Single administration (~4m cells/finger)	
Sites	Sites Single site - Marseille, France	
Endpoints	 Cochin Hand Function Scale Raynaud's Condition Score Scleroderma Health Assessment Questionnaire Pain Modified Rodnan Skin Score Capillaroscopy Adverse events Other 	
Follow-Up	24 months	
Status	Complete	



- Six and 12 month data published^{1,2}
- 24 month data presented at Systemic Sclerosis World Congress in Lisbon, Portugal, February 19, 2016

1. Granel et al (2014); Ann Rheum Dis Aug 11: doi: 10.1136/annrheumdis-2014-205681 2. Guillaume-Jugnot et al (2015) Rheumatol. 10.1093/rheumatology/kev323

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SCLERADEC I Improvement Through 24 months

ECCS-50 Treatment led to improvement in hand function, Raynaud's phenomenon, and pain



Key Observation:

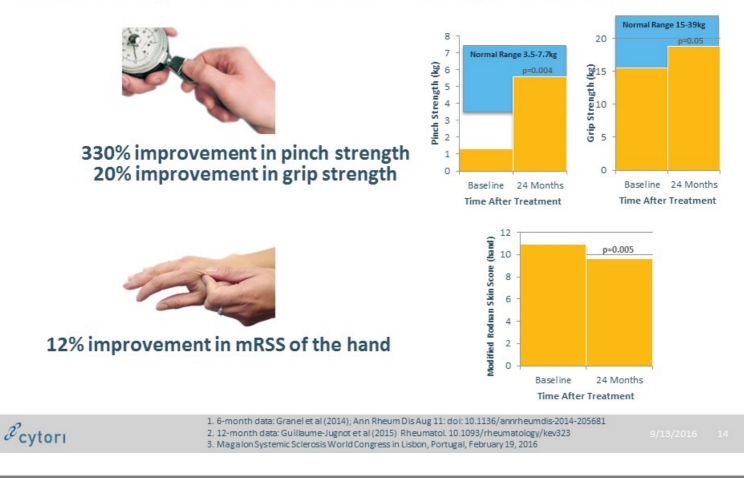
- Concordant reduction (~50%) in four key symptomatic patient reported outcomes
- Efficacy sustained to two years following a single treatment



1. 6-month data: Granel et al (2014); Ann Rheum Dis Aug 11: doi: 10.1136/annrheumdis-2014-205681 2. 12-month data: Guillaume-Jugnot et al (2015) Rheumatol. 10.1093/rheumatology/kev323 3. Magalon Systemic Sclerosis World Congress in Lisbon, Portugal, February 19, 2016

SCLERADEC I- Other Endpoints

Sustained improvement in hand strength & skin stiffness

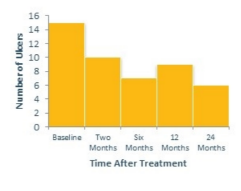


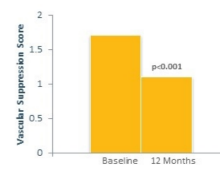
SCLERADEC I- Other Endpoints

Reduction in digital ulcers, improved microvascular architecture



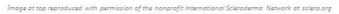
40% improvement in number of ulcers





30-35% improvement in vascular suppression score

VSS data at 24 months not available





6-month data: Granel et al (2014); Ann Rheum Dis Aug 11: doi: 10.1136/annrheumdis-2014-205681
 2. 12-month data: Guillaume-Jugnot et al (2015) Rheumatol. 10.1093/rheumatology/kev323
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Clinical/Regulatory Strategy

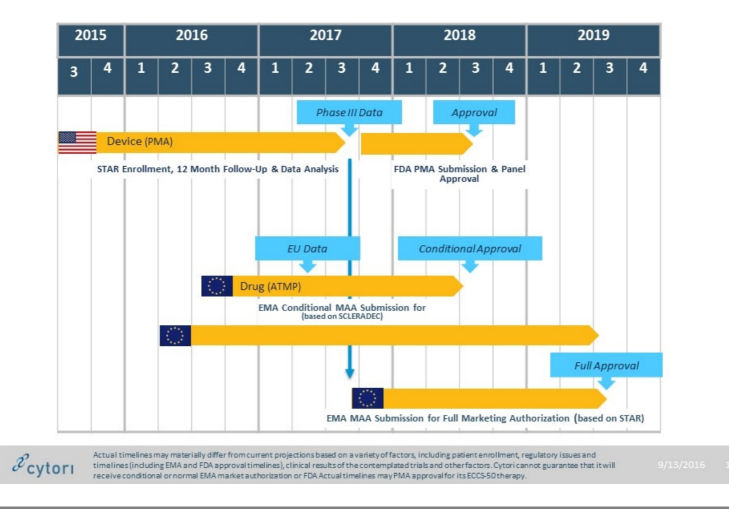
- EU SCLERADEC I trial data used to support US FDA STAR trial approval, potential EU Conditional Marketing Authorization
- US FDA STAR trial for US PMA approval
- US STAR trial ± SCLERADEC II to obtain Full Marketing Authorization

	STAR (Phase III)	SCLERADEC II (Phase III)*	
Study size	88	40	
Randomization	1:1, active: placebo	1:1 (dose from Pilot, placebo)	
Crossover	Placebo, crossover at 48 weeks Placebo, crossover at 24 weeks (
Sites	Up to 20 in USA	6 France	
Primary Endpoint	Cochin Hand Function Score (CHFS) at 6 months	Cochin Hand Function Score at 3 months	
Secondary Endpoints	CHFS, Raynaud's Condition Score, Scleroderma Health Assessment Questionnaire, Pain, Modified Rodnan Skin Score, Hand Mobility in Scleroderma Test, Adverse events CHFS, Raynaud's Condition S Scleroderma Health Assessm Questionnaire, Pain, Modifie Score, Capillaroscopy, Adver		
Follow-Up	48 weeks 24 weeks		
Status	Enrolled, Data in mid-2017	Enrolling	



* Investigator-initiated tria

Scleroderma - Projected Development Timeline





1	Provide ethical and compliant access to Cytori Cell Therapy [™] , ECCS-50, for hand scleroderma patients prior to EMA marketing authorization
2	Increase awareness of and facilitate a positive experience with Cytori Cell Therapy [™] among healthcare providers in advance of commercial launch
3	Track and collect key program data and documentation providing valuable insight regarding the demand for and use of Cytori Cell Therapy™
4	Implement a chargeable program in EMEA countries where regulations allow
5	Launch the program in Q1 and begin treating patients in Y1 and close the program once reimbursement is attained in each EMEA country
	IDIS
	— Managed Access



<u>Pipeline Indications</u> Knee Osteoarthritis Urinary Incontinence Radiation/Thermal Burn



Knee Osteoarthritis

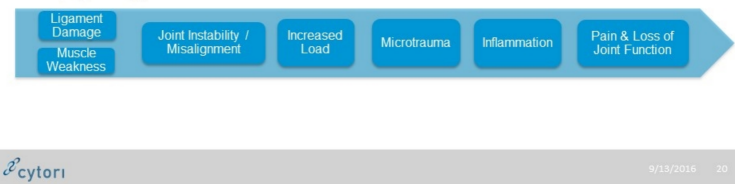
Osteoarthritis

- Progressive loss of joint function
- Imbalance between anabolic (cartilage-forming) and catabolic (cartilage-destroying) processes driven by synovial inflammation
- Distinct from RA

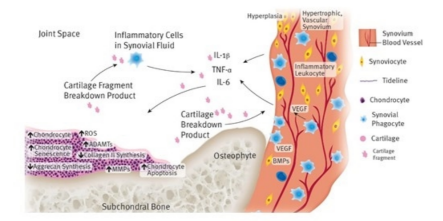
Pathophysiology

Epidemiology

- OA is the most common form of arthritis
- 13.9% of adults <u>></u>25 years
- 33.6% (12.4 million) ≥65 years
- Estimated 26.9 million US adults (2005)



Scientific Rationale: Cytori Cell Therapy in OA



- Pathophysiology of OA (persistent synovial inflammation leading to cartilage destruction) overlaps with other clinical indications in which Cytori Cell Therapy shown to have impact
- Combination of veterinary, preclinical, *in vitro*, and pilot clinical data indicate significant potential for symptomatic improvement and potential disease modification

Opportunity: Biologic/Cell Therapy to better address gap between analgesics and surgical management

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ACT-OA Trial

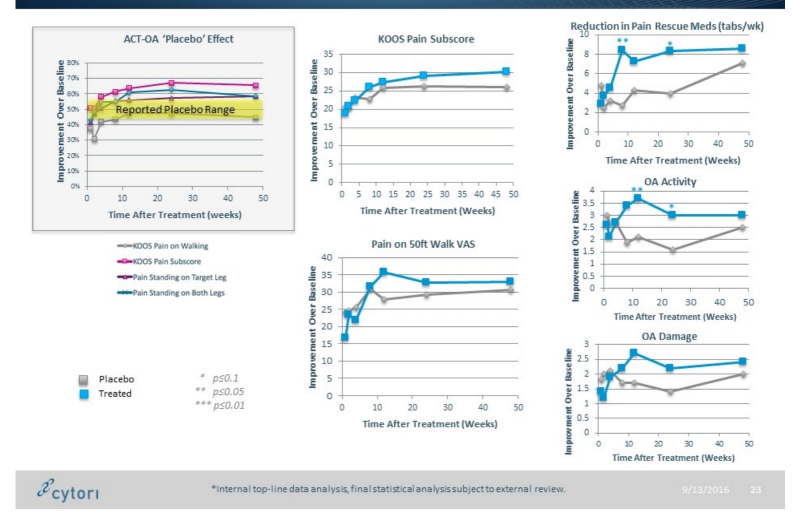
	Phase II (ACT-OA)	
Study Size	94 enrolled	
Randomization	n 1:1:1 (low dose, high dose, placebo)	
Sites	12 US	
Primary Endpoint	KOOS - pain on walking @ 12 weeks, not powered	
Secondary Endpoints	KOOS, pain/function questionnaires, disease activity pain meds, SF-36, MRI@ 48 weeks	
Follow-Up	48 weeks	
Status	Enrolled, completed 48 week topline assessment	
Next Steps Partnership discussion ongoing for Phase III/commercial		

48 Week Preliminary Data- Top-line Results

- No SAEs related to cell therapy or procedure
- Consistent 12, 24, 48 week trends favoring cell therapy effect in patient reported outcomes
- Pain PROs, pain on walking question @12 weeks- trends not reaching statistical significance
- Substantial effect from baseline to 12, 24, 48 week, active vs. placebo smaller on relative basis
- · Consistent effect in MOAKS/MRI imaging in several parameters at 48 weeks

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ACT-OA Trial Preliminary Top-line 48 Week Data, Patient Reported Outcomes



ACT-OA Trial Preliminary Top-line 48 Week Data, MRI or MOAKS



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Stress Urinary Incontinence Program

'ADRESU' Trial Objectives

- Approved, reimbursed therapy for SUI in men following prostate intervention
 - Unmet need for patients
- Support proof of concept in female incontinence

Progress/Data

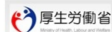
- Pilot clinical trial data published ^{1,2}
 - Increase maximum urethral closing pressure
 - Reduction 24-hour pad weight
 - Increased blood flow

Ongoing 45 pt. Multicenter Pivotal Trial

- Anticipate >50% enrollment by YE 2016
- Anticipate enrollment completion 2017

1. Gotoh et al. (2014) Int J Urology 21 (3) 294-300 2. Yamamoto et al. (2012) Int J Urology 19 (7) 652-9





GRADUATE SCHOOL OF MEDICINE NAGOYA SCHOOL OF HEALTH SCIENCES

Support

- Investigator initiated with Cytori support
- Substantial funding via Japanese Ministry of Health, Labour and Welfare

Development Plan

- Ongoing pivotal trial sufficient for approval/reimbursement
- Assuming positive data, seek approval and reimbursement based on 12 month assessment
- Potential partnering opportunity

Radiation/Thermal Burn Program

Objectives

- Development medical countermeasure for mass casualty event- thermal burn ± radiation exposure
- Proof of concept clinical data for use of Cytori Cell Therapy in wound healing

Progress/ Preclinical Data

- Improvement in multiple tissue repair parameters following administration of Cytori Cell Therapy^{1,2}
- Effective via multiple routes of administration^{1,2}
- Efficacy sustained following substantial exposure to radiation dose³

1. Foubert et al. (2015) Burns doi:10.1016/j.burns.2015.05.004 2. Foubert et al. (2015) Adv Wound Care doi:10.1089/wound.2015.0672 3. Foubert et al (manuscript in preparation)



Support





- Funded by contract of up to \$106MM from Biomedical Advanced Research and Development Authority (BARDA)
- \$18.7MM of funding allocated through September 2016

Development Plan

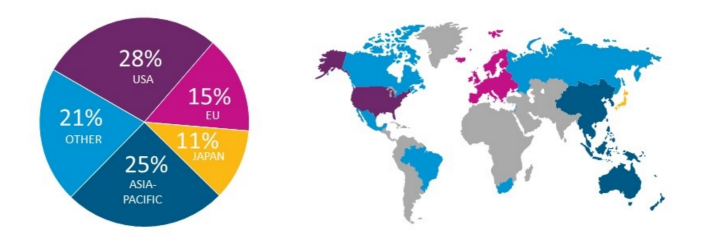
- Submit IDE application in 2016 for a pilot clinical trial
- Enrollment begins 2017
- Additional funding anticipated, pending receipt of IDE approval for clinical trial

Corporate Information



Cytori Cell Therapy: Global Patent Estate

89 patents issued worldwide; over 76 applications pending



Goal: Protect Cytori's proprietary methods and devices for manufacturing Cytori Cell Therapy, as well as methods of using Cytori Cell Therapy in the treatment of scleroderma, and several other indications, including osteoarthritis and SUI.

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Capitalization Summary

Select Data – as of 6/30/16		
Cash	~ \$20.0MM	
Senior term loan	~ \$17.7MM	
Common Shares outstanding	~ 20.5MM	
Outstanding options, RSAs and warrants	~ 4.4MM	
Fully diluted share count	~ 24.9MM	
Market capitalization	~ \$42MM*	

* Based on share price of \$2.05 at closing on September 2, 2016.



Update- Corporate Objectives & Milestones

2016 Milestones			
1 st Half	 ✓ EU MAP program launch ✓ 24 WK ACT-OA interim data evaluation ✓ SCLERADEC-I two year follow-up data ✓ Full STAR phase III trial enrollment 		
2nd Half	 ✓ 48 WK ACT-OA data evaluation ✓ Japan & MAP progress reported FDA BARDA and Orphan approvals SCLERADEC-I three year follow-up data SCLERADEC-II enrollment slow 		
2017 Milestone	es		
	 STAR Phase III one year follow-up data SCLERADEC-II 24 WK follow-up data Submit for US FDA PMA approval scleroderma Submit for EMA marketing authorization scleroderma US Phase I BARDA-funded trial enrollment Full ADRESU enrollment 		

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Thank You

