

Delivering More For Patients Positively Impacting Healthcare



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Nasdaq: PSTV

Biotech Showcase January 14, 2020

Forward-Looking Statements

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. These statements are based on our management's current beliefs, expectations and assumptions about future events, conditions and results and on information currently available to us.

All statements, other than statements of historical fact, included herein regarding our strategy, future operations, financial position, future revenues, projected costs, plans, prospects and objectives are forward-looking statements. Words such as "expect," "anticipate," "intend," "plan," "believe," "seek," "estimate," "think," "may," "could," "will," "would," "should," "continue," "potential," "likely," "opportunity" and similar expressions or variations of such words are intended to identify forward-looking statements, but are not the exclusive means of identifying forward-looking statements. Additionally, forward-looking statements include, without limitation, statements about our anticipated expenditures, including those related to pre-clinical and clinical trials and research studies and development, sales and marketing, and general and administrative expenses, the potential size of the market for our products, future development and/or expansion of our products in our markets, our ability to generate product or development revenues and the sources of such revenues, our ability to effectively manage our gross profit margins, our ability to obtain and maintain regulatory approvals, expectations as to our future performance, liquidity and capital resources, including our potential need for additional financing and the availability thereof, our ability to continue as a going concern, our ability to remain listed on the Nasdag Capital Market, our ability to repay or refinance some or all of our outstanding indebtedness and our ability to raise capital in the future, and the potential enhancement of our cash position through development, marketing, and licensing arrangements. Such statements are based on currently available operating, financial and competitive information and are subject to various risks, uncertainties and assumptions that could cause actual results to differ materially from those anticipated or implied in our forward-looking statements due to a number of factors including, but not limited to, the early stage of our product candidates, the results of our research and development activities, including uncertainties relating to the clinical trials of our product candidates, our need and ability to raise additional cash, the outcome of our partnering/licensing efforts, risks associated with laws or regulatory requirements applicable to us, market conditions, product performance, potential litigation, competition within the pharmaceutical industry, and other factors set forth in our most recent Annual Report on Form 10-K and our Quarterly Reports on Form 10-Q, as well as any amendments thereto, filed with the Securities and Exchange Commission. Given these risks, uncertainties and other factors, many of which are beyond our control, you should not place undue reliance on these forward-looking statements.

Except as required by law, we assume no obligation to update these forward-looking statements publicly, or to revise any forward-looking statements to reflect events or developments occurring after the date of this presentation, even if new information becomes available in the future.



Our Mission

Our Vision

We believe in the importance of discovering, developing, and delivering complex and innovative treatments for patients battling cancer and rare diseases.

We envision medicines that significantly enhance clinical and economic outcomes for patients and providers and forever change the future of human health worldwide.



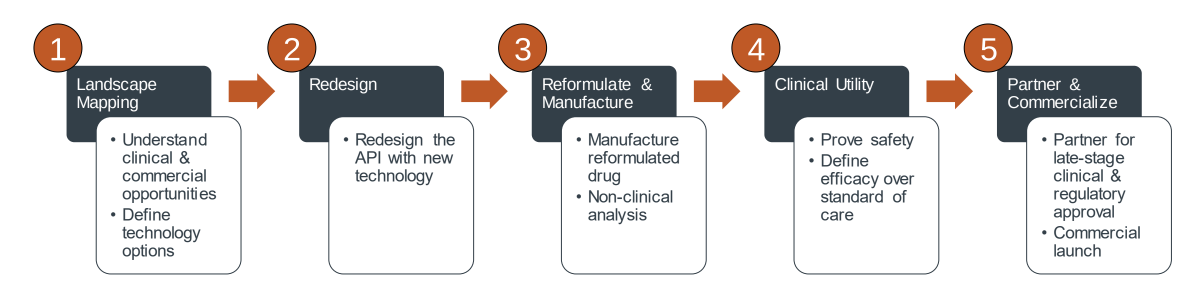
2019 Transition Achieved, 2020 Poised for Expansion

- + Strategic Transition to Clinical-Stage Oncology / Rare Disease Portfolio
 - + Divested legacy assets
 - + Pivot emphasis to first 2 portfolio drugs
 - + Actively evaluating potential portfolio additions
- Achieved Long-Term Capitalization
- + Right-Sized the Company
 - + Optimized cost structure
 - + Virtualized operations
 - + Strategic move of HQ to Texas
 - + Built for scale at sustainable margin



The PLUS Development Model







Putting the 'Plus' in PLUS

- + Clinically or commercially proven drugs + novel delivery technology = improved efficacy for patients
- + PLUS Current Pipeline:

THERAPEUTICS

- + Serve unmet or underserved medical needs
- + Deliver improved efficacy through technology innovation
- + Addressable market opportunity of \$250M+
- + DocePLUS[™] -- proprietary innovative liposomal docetaxel formulation
 - + Potential benefits over comparative drugs
 - + Published U.S. Phase 1 clinical trial results
 - + Received FDA orphan drug designation for small cell lung cancer
 - + Confirmed with FDA that 505(b)(2) NDA pathway appears to be an acceptable approach
 - + Preparing for Phase 2 clinical trial for small cell lung cancer
- + DoxoPLUS[™] -- complex generic liposomal doxorubicin
 - + Completed clinical trial versus CAELYX® for EU bioequivalence
 - + cGMP validated facility for R&D, manufacturing, analytical testing
 - + Seeking divestiture to focus on innovative product candidates
- + Ongoing evaluation of new product candidates

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Capabilities and Experience

- + Decades of drug, nanotechnology, and combination product development experience
- + Management team with successful track record in completing clinical trials and launching novel products in the U.S. and internationally
- + A fully GMP-compliant, validated, state-of-the-art, U.S. development and manufacturing facility supported by an ICH-compliant Quality Management System
- + A 10,000 square foot facility with commercial scale negative pressure manufacturing suites with ISO Class 7 and 8 cleanrooms, onsite WFI and full handling capability for hazardous cytotoxics

+ A dedicated analytical chemistry development and validation lab

Pipeline



DocePLUS™

Proprietary Innovative IV Albumin-Stabilized PEGylated Liposomal Docetaxel*



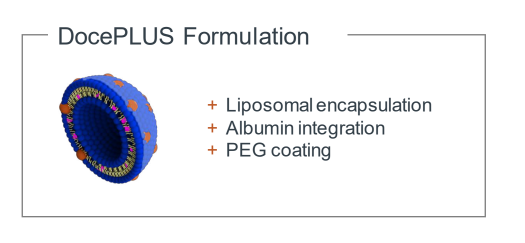
DoxoPLUS™

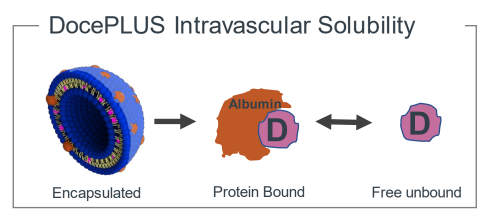
Complex Generic IV PEGylated Liposomal Doxorubicin Hydrochloride**

PROGRAM	CANCER TARGET	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	REGULATORY REVIEW	COMMERCIALIZATION RIGHTS
DocePLUS™	Small Cell Lung						PLUS - Global
PROGRAM	CANCER TARGET	PRECLINICAL	В	IOEQUIVALEN)E	REGULATORY REVIEW	COMMERCIALIZATION RIGHTS



DocePLUS[™] Design Rationale, PK and PD





- + DocePLUS Liposomal Encapsulation
 - + Free unbound docetaxel distributed to tissues
 - + Encapsulated AUC 2-3X greater than unencapsulated¹
 - + Intravascular depo & reduced clearance, lower V_{ss}²
- + DocePLUS Albumin Integration
 - + Liposomal stabilization
 - + Docetaxel binds to albumin in plasma²
- + DocePLUS Polyethylene Glycol Coating (PEG)
 - + Reduced macrophage uptake
- + Compared to TAXOTERE®/Tween 80
 - + Surfactant increases unbound docetaxel, ~15%²
 - + DocePLUS free docetaxel C_{max} 3.3x less TAXOTERE³
 - + AAG (α1-acid glycoprotein) acute phase reactant also responsible for plasma protein binding & variable in cancer patients²

TAXOTERE is a registered trademark of Sanofi

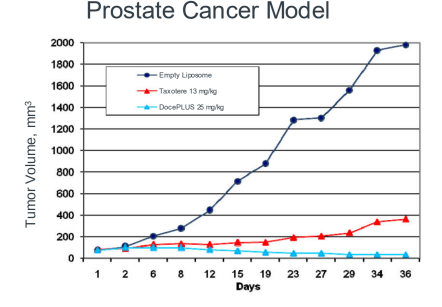
l Mahalingham et al 2014

- 2 Loos et al., Clin Pharmacol Ther 2003; 74: 364-371
- 3 NIH National Characterization Laboratory, NCL200911A

4 Bruno et al., Am J Health-Syst Pharm Vol 54 Dec 15 1997 Supp 2

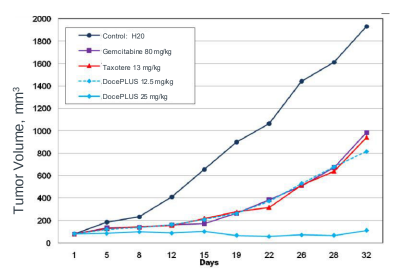
DocePLUS[™] Preclinical Findings

- Preclinical pharmacokinetic data support elevated AUC and C_{max} of docetaxel for nanoparticle formulation
- + Biodistribution show preferential uptake to lung & liver
- + Immunodeficient xenograft tumor models in mice show
 DocePLUS[™] efficacy against a range of human tumor types
 - + Lung, prostate, pancreatic, mesothelioma



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Pancreatic Cancer Model



DocePLUS[™] Phase 1 Study Key Findings

+ First-in-human, open-label, dose escalation (15-110 mg/m²) study of DocePLUS[™] in 29 heavily pretreated patients (~33% had prior TAXOTERE® exposure) with advanced solid tumors at 2 U.S. sites

+ Safety

- + DocePLUS[™] achieved a 20% increase in MTD vs. standard docetaxel
- + 10 patients treated at MTD
 - + Treatment-emergent adverse events included neutropenia, anemia, fatigue, and nausea
- + ~2-3 fold increase in AUC compared with free (nonencapsulated)docetaxel
- + Efficacy
 - + 22 of 29 patients (76%) experienced stable disease
 - + 1 patient with NSCLC and previous exposure to docetaxel had confirmed partial response to DocePLUS™
 - + 2 prostate cancer patients had PSA reduction >95%: 1 patient was progression free for 54 weeks

DIUS THERAPEUTICS

Cancer Chemother Pharmacol (2014) 74:1241-1250 DOI 10.1007/s00280-014-2602-x

ORIGINAL ARTICLE

Phase I study of intravenously administered ATI-1123, a liposomal docetaxel formulation in patients with advanced solid tumors

Devalingam Mahalingam · John J. Nemunaitis · Laeeq Malik · John Sarantopoulos · Steven Weitman Kamalesh Sankhala · Jessica Hart · Ahmed Kousba · Nicole S. Gallegos · Gavin Anderson · John Charles Jon M. Rogers · Neil N. Senzer · Alain C. Mita

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Abstract

etaxel and may be administered without the premedications and hypersensitivity reactions. This Phase I study examines the safety, tolerability, pharmacokinetics (PKs), and antitumor activity of ATI-1123. Methods Patients with advanced solid malignancies Conclusions ATI-1123 demonstrated an acceptable tol-1-h every 3 weeks. The dosing commenced using an accelerated titration design and was followed by a modified

3 + 3 Fibonacci schema to determine maximally tolerated dose (MTD). Plasma was analyzed for encapsulated/nonencapsulated docetaxel; PK analyses were performed using model independent method. Response was assessed using RECIST criteria.

Results In total, 29 patients received doses ranging from 15 to 110 mg/m2. At 110 mg/m2, two of six patients expe rienced dose-limiting toxicities including grade 3 stomatitis and febrile neutropenia. The 90 mg/m2 cohort was

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expanded to ten patients and identified as the MTD. The Purpose ATI-1123 is a liposomal formulation of doc- most common adverse events were fatigue, nausea, neutropenia, anemia, anorexia, and diarrhea. ATI-1123 exhibited linear and dose proportional PKs. One patient with lung cancer had confirmed partial response, and stable disease was observed in 75 % patients.

received escalating doses of ATI-1123 intravenously over erability and favorable PK profile in patients with solid tumors. Our results provide support for Phase II trials to determine the antitumor activity of this drug.

> Keywords ATI-1123 · Docetaxel · Safety · Tolerability · Pharmacokinetics

Introduction

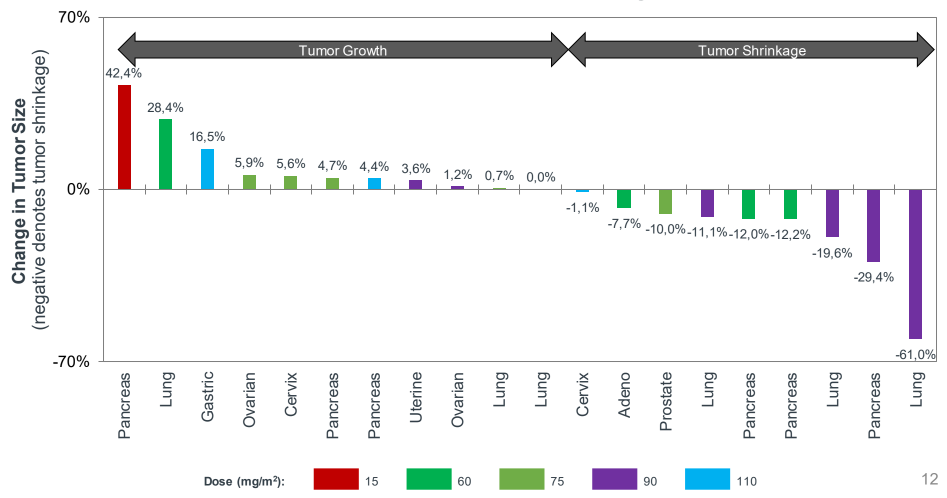
Docetaxel is an antimitotic agent which hinds to the beta subunit of tubulin and causes stabilization of tubulin polymerization. This stabilization results in cell cycle arrest at the G₂/M phase, thus inhibiting mitosis [1]. It is a poorly water soluble semisynthetic taxane analogue, commonly used in the treatment of a variety of solid tumors including head and neck, non-small cell lung, prostate, breast, and gastric cancer 12-41. The current recommended regimen for docetaxel is 60-100 mg/m2 administered over 1-h every 3 weeks, depending upon the indication [5].

Taxotere[®] is the standard formulation of docetaxel with well-established safety and efficacy when administered in 3-week cycles [6]. Because of its poor water solubility, it is formulated with solvents that can potentially contribute to treatment-related adverse events such as hypersensitivity reaction. Current docetaxel and other taxane formulations often complicate drug delivery and can alter both the PK and toxicity profiles. These problematic issues have spurred

DocePLUS[™] Clinical Antitumor Activity

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Patients in Order of % Change



DocePLUS[™] Development Plan

- + Underserved Market Opportunity
 - + Small Cell Lung Cancer; >\$250M annual revenue potential
- + Therapeutic redesign: known API PLUS new technology
 - + Docetaxel PLUS protein-stabilized PEGylated liposome encapsulation
- + Regulatory path
 - + Orphan, 505(b)(2)
- + Therapeutic reformulation: CMC, nonclinical
 - + Improved drug performance
- + Clinical

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- + Phase 1 complete
- + Phase 2 ready
- + Commercial: partner or direct sales

Why Small Cell Lung Cancer?

+ Lung Cancer¹

- + 2nd most common cancer
- + 14% of all new cancer cases
- + Leading cause of cancer-related death in U.S.
- + Diagnosed at median age of 70
- + 10-15% classified as Small Cell Lung Cancer¹
 - + Smoking is #1 risk factor²
 - + Low 5-year survival of 8-31%³
 - + Most patients relapse following first-line therapy⁴
 - + HYCAMTIN® (topotecan), first FDA-approved in 1998, is indicated for platinum-sensitive patients who relapse
 >60 days after start of first-line therapy⁵



1 https://www.cancer.org/cancer/small-cell-lung-cancer/about/key-statistics.html
2 https://www.cancer.org/cancer/small-cell-lung-cancer/causes-risks-prevention/risk-f
3 https://www.cancer.net/cancer-types/lung-cancer-small-cell/statistics
4 https://ascopubs.org/doi/full/10.1200/J0P.18.00204
5 https://www.accessdata.fda.gov/drugsatfda_docs/label/2018/020671s0231b1.pdf

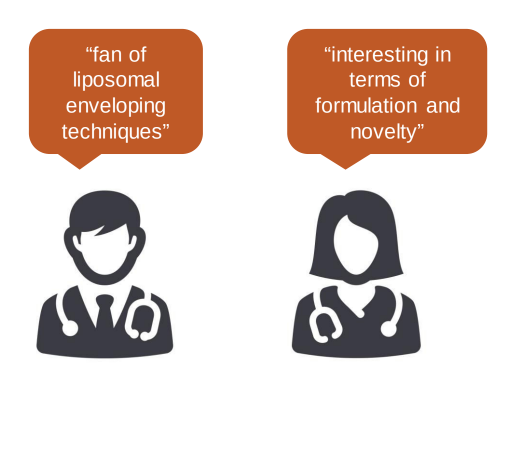




Unmet Needs for Second-Line SCLC Treatment

- + DocePLUS: rationale for use in SCLC
 - + Biodistribution favorable for pulmonary delivery
 - Phase I potential efficacy signal in lung cancer (NSCLC)
- + Single-Agent Chemotherapy with HYCAMTIN®
 - + Demonstrated activity in SCLC
 - In randomized clinical trial, key endpoints (ORR, DoR, TTP, OS) were not statistically improved over CAV treatment
 - + Adverse reactions included neutropenia, anemia, and thrombocytopenia
 - Difficult dosing schedule: 1.5 mg/m² IV infusion over 30 minutes daily on days 1 to 5 of each 21-day cycle

DocePLUS™ Market Research



In considering 10 solid tumors and 3 target product profiles, oncologists viewed SCLC as an attractive Phase 2 study indication

- + Low cure rate, rapid progression, low survival
- + Lack of effective treatment options
- + For second-line, view current standard of care as toxic, inconvenient, and minimally efficacious
- + Expect shorter trial and easier patient recruitment (relative)

Oncologists also cited pancreatic cancer as an attractive target and described a high willingness to use DocePLUS[™] if FDAapproved for this indication

- + Limited treatment options
- + Difficulties in achieving good results with new treatments



DocePLUS[™] Proposed Market Differentiation in SCLC

Compared to $\ensuremath{\mathsf{HYCAMTIN}}\xspace{\mathbb{R}}$

- Convenient dosing schedule: IV infusion over 60 minutes on only day 1 of each 21-day cycle
- + Potential improved biodistribution to lung
- + Potential for improved safety & efficacy

Compared to TAXOTERE®

- Eliminates need for Tween 80 surfactant and associated with hypersensitivity reactions by encapsulating docetaxel in lipid bilayer of a liposome and albumin
- + Potential improved pharmacokinetics & pharmacodynamics^{1,2,3}
- + Potential for improved safety & efficacy



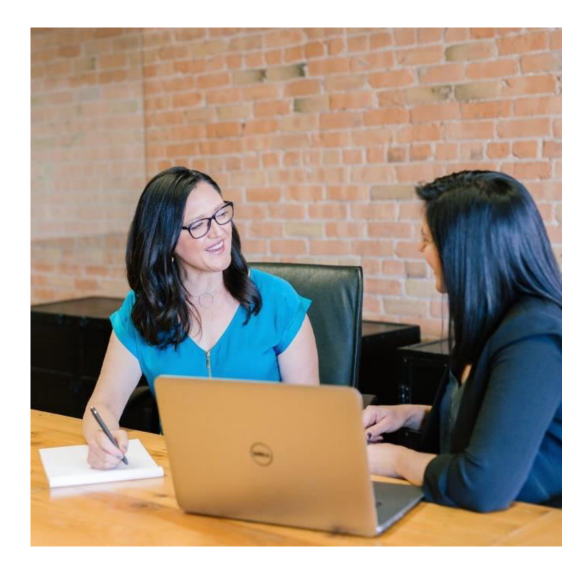
¹ Mahalingham et al 2014

^{17 2} Loos et al., Clin Pharmacol Ther 2003; 74: 364-371 HYCAMTIN is a registered trademark of Nova 3 NIH National Characterization Laboratory, NCL200911A TAXOTERE is a registered trademark of Sand



Corporate Information

- + Headquarters: Austin, Texas
- + Manufacturing: San Antonio, Texas
- + Employees: 10 full-time & 3 part-time
- + Nasdaq Symbol: PSTV
- + Website: plustherapeutics.com



2020 Key Milestones

+ DocePLUS[™] Phase 2 Clinical Trial in Small Cell Lung Cancer

- + Q1 2020: Complete 3rd party testing & manufacturing contracts
- + H1 2020: Complete FDA clinical trial development plan
- + Mid 2020: Clinical manufacturing lots
- + Q3 2020: 1st patient enrolled
- + Mid 2021: Patient enrollment complete
- Pipeline opportunities
 - + Expand DocePLUS[™] indications
 - + Acquire and/or in-license new drugs for oncology and rare disease indications
- + Partner/Divest DoxoPLUS[™] complex generic asset



Capitalization Summary

Select Data †				
Cash	\$21.4M*			
Common Shares Outstanding	2,841,588			
Options Outstanding	253,818			
Warrants Outstanding	7,238,400			
Senior Term Loan	\$9.3M			





Thank You

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Marc Hedrick, MD President & CEO

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