



CREATING TOMORROW, TODAY.

OTCQB: CYTR

Corporate Overview June 2021

Non-Confidential

CytRx Safe Harbor Statement

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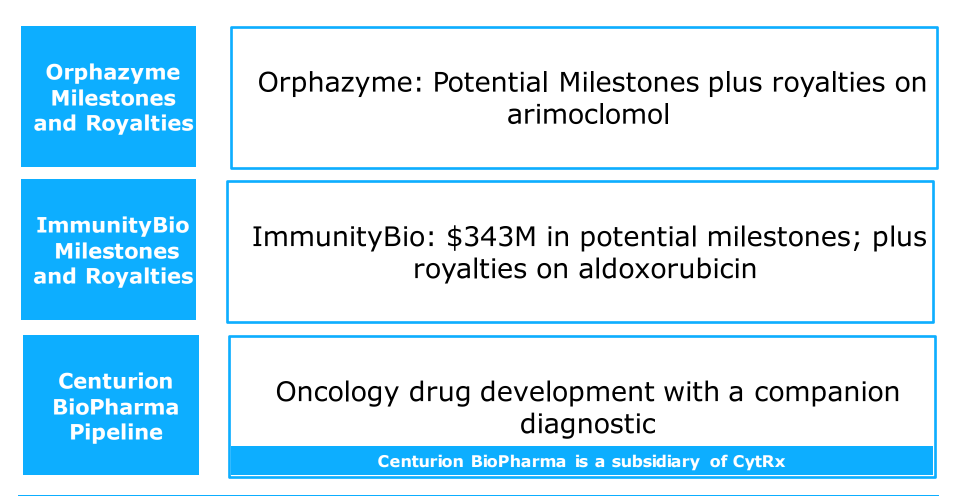


CytRx Highlights

- CytRx's milestone and royalty agreement with Orphazyme for arimoclomol could represent potential near term payments to CytRx
- Orphazyme has submitted a Marketing Authorisation Application with EMEA authorities for arimoclomol for NPC
- ImmunityBio has initiated a Phase 2 registrational-intent study for first-line and second-line locally advanced or metastatic pancreatic cancer, which includes aldoxorubicin
- Centurion BioPharma is a private oncology drug development company focused on cancer and has completed extensive pre-clinical work for its ultra high potency LADRTM drug candidates and albumin companion diagnostic (ACDx)



CytRx has potential milestone/royalty payments and a subsidiary called Centurion BioPharma





CytRx milestones and royalties from Orphazyme for Arimoclomol

Orphazyme Milestones and Royalties

Orphazyme: Potential milestones in addition to royalties on arimoclomol

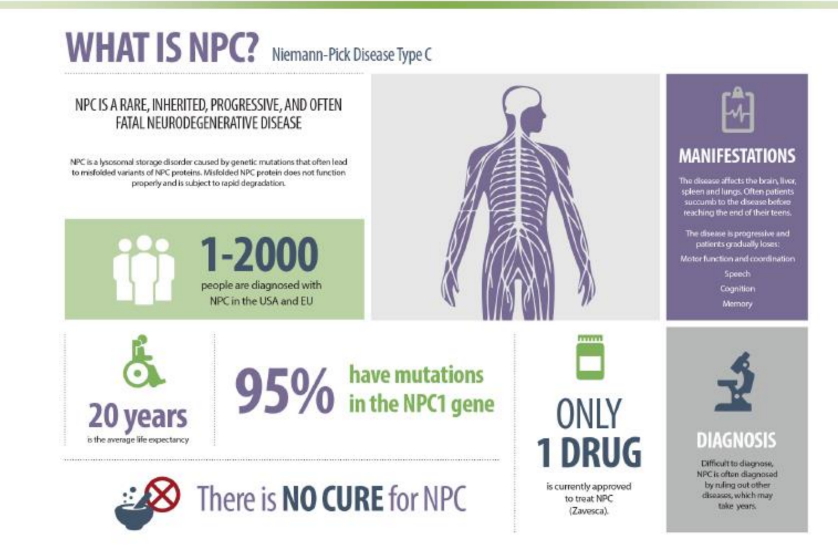
Niemann-Pick disease ("NPC")

- Orphazyme submitted an MAA with the EMA for arimoclomol for NPC, with European regulatory CHMP opinion expected in Q4/21
- Orphazyme filed an NDA with the FDA with Priority Review and received a Complete Response Letter on June 17, 2021; they will be working with the regulators to find a possible path forward.
- Orphazyme launched an Early Access Program for NPC in January 2020 to further accelerate access to treatment with arimoclomol for people living with NPC.
- Total worldwide patients approximately 3,000.
- Expected price range is \$300,000 \$600,000; EU market potential \$300 Million.
- Go to market in EU/RoW H1 2022.



Niemann-Pick Disease Type C (NPC)

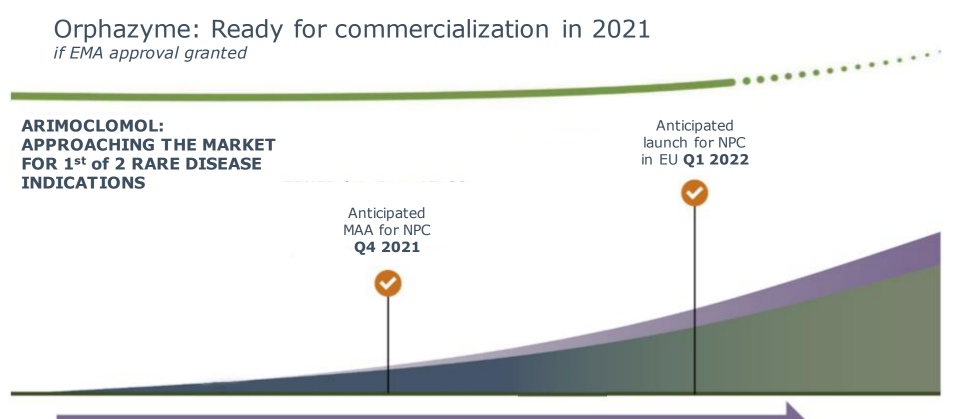




Source: www.orphazyme.com

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Orphazyme ready for commercialization in 2021 for arimoclomol



Building a highly specialized commercial footprint in US and EU



CytRx potential milestones and royalties from ImmunityBio for aldoxorubicin

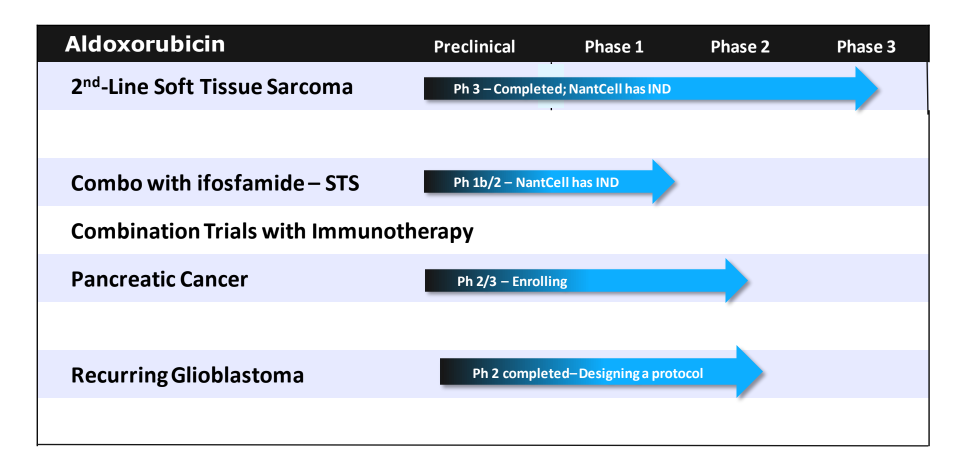
ImmunityBio Milestones and Royalties

ImmunityBio: up to \$343M in milestones In addition to royalties on aldoxorubicin

- ImmunityBio has merged with NantKwest and trades under IBRX
- ImmunityBio has highlighted aldoxorubicin as one of three separate modalities of its platform
- ImmunityBio announced initiation of a phase 2 registrational-intent study using aldoxorubicin in combination with immunotherapy in metastatic pancreatic cancer
- ImmunityBio, to date, plans to use aldoxorubicin in studies in glioblastoma, in addition to metastatic pancreatic cancer.
- CytRx is entitled to increasing double-digit royalties on aldoxorubicin for soft tissue sarcomas and increasing single-digit royalties for all other indications
- ImmunityBio is reviewing options in Soft Tissue Sarcoma



CytRx partnered Pipeline with ImmunityBio - aldoxorubicin





Update from NantKwest/ImmunityBio at JP Morgan Conference in January 2021

Metastatic Pancreatic Cancer QUILT-88: early indications of increased survival rate with no other approved treatment options

- In initial QUILT trials, median overall survival rate more than doubled compared to historical controls
- A single-arm Phase 2 trial was initiated in October 2020, for which the primary endpoint is overall survival and 83% of patients enrolled with second-line or greater pancreatic cancer remain alive to date
- Former Senate Majority Leader Harry Reid's stage IV pancreatic cancer is now in "complete remission" after receiving this experimental combination immunotherapy that included aldoxorubicin
- Initiation of a <u>Registrational-Intent</u> Phase 2 randomized, three-cohort, open-label study for first and second-line treatment of locally advanced or metastatic pancreatic cancer
- Randomized trials in first and second-line pancreatic cancer are actively recruiting at three sites with more than 50 patients enrolled or being evaluated in QUILT-88 to date



CytRx subsidiary Centurion BioPharma has an oncology preclinical pipeline and diagnostic

Centurion BioPharma Pipeline

Oncology drug development with a companion diagnostic

LADR[™] (linker activated drug release) <u>albumin</u> binding drug conjugates

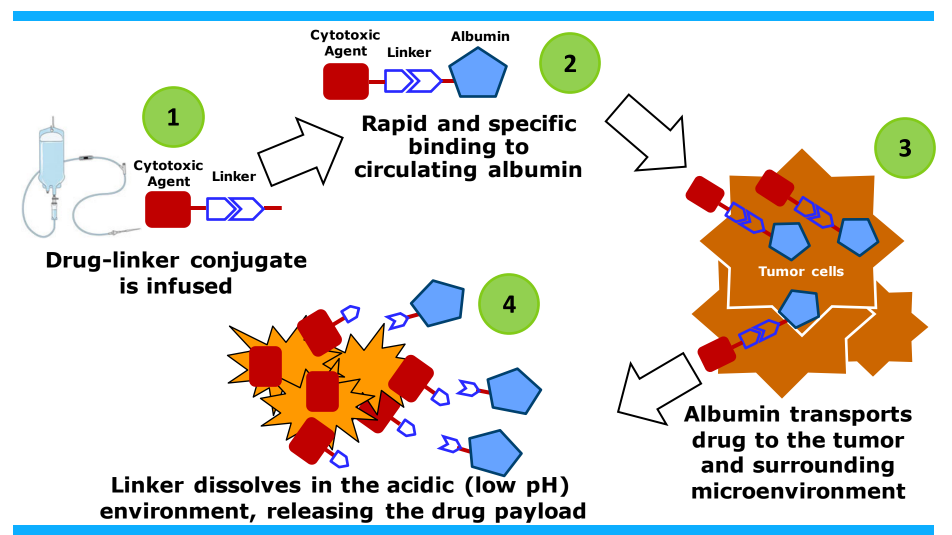
LADR-7 (auristatin) LADR-8 (auristatin) LADR-9 (maytansinoid) LADR-10 (maytansinoid)

Albumin companion diagnostic (ACDx)

identifies tumors eligible for treatment with LADR[™]



LADRTM Mechanism of Action





Recent and Upcoming Catalysts

2020-2021

- ✓ 1H 2020: Orphazyme filed for FDA approval for arimoclomol in Niemann-Pick Type C disease with a target action date of 06/17/21
- ✓ 2H 2020: Orphazyme has submitted for EMEA (Europe) approval for arimoclomol in Niemann-Pick Type C ("NPC") disease
- ✓Q4 2021: Orphazyme expecting European regulatory CHMP opinion for arimoclomol in NPC
- 2021: Upon approval, CytRx is to receive a \$6 million milestone payment if Europe and Japan are approved (\$4 million for Europe and \$2 million for Japan)



Financial Summary

 Cash Position (03/31/21) 	\$9.3M
 No Debt 	
 Shares Outstanding 	36.5M
 Options Weighted-average strike price: \$7.43 	3.2M
 Fully-Diluted Share Count (3/31/2021) 	39.7M



Summary

- Orphazyme could deliver milestones and royalties
- ImmunityBio could deliver milestones and royalties
- Cash burn rate is ~\$430k per month
- Potential to shelter future income with non-restrictive net-operating carry-forward losses ("NOL's") of \$258 million

