

Company Presentation
May 2022



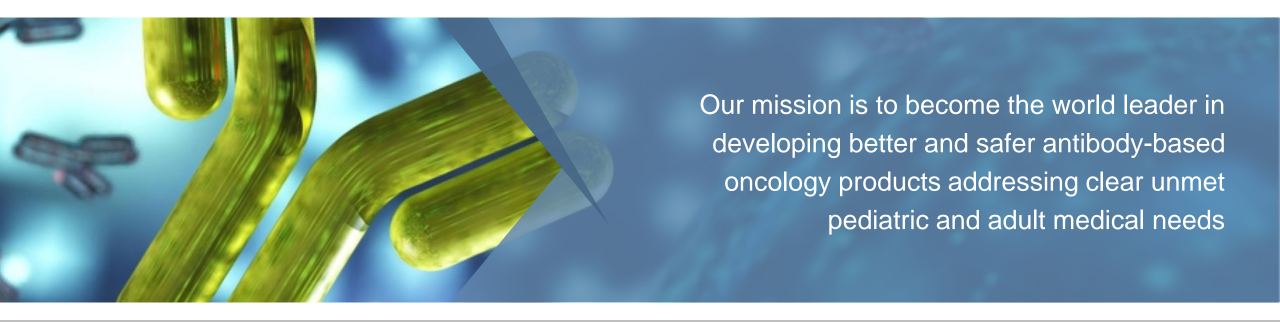
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This presentation contains forward-looking statements within the meaning of the US Private Securities Litigation Reform Act of 1995. The forward-looking statements involve substantial risks and uncertainties. All statements, other than statements of historical facts, contained in this presentation, including statements regarding our strategy, future operations, future financial position, future revenue, projected costs, prospects, plans and objectives of management and expected market growth are forward-looking statements. The words "anticipate," "believe," "continue," "could," "estimate," "expect," "intend," "may," "might," "plan," "potential," "predict," "project," "should," "target," "would" and similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. Such statements include, but are not limited to, statements about regulatory approvals, clinical trial timing and plans, the achievement of clinical and commercial milestones, future financial and operating results, business strategies, market opportunities, financing, and other statements that are not historical facts. Our product candidates and related technologies are novel approaches to cancer treatment that present significant challenges. Actual results may differ materially from those indicated by such forward-looking statements as a result of various factors, including but not limited to: risks associated with our financial condition and need for additional capital; risks associated with the Company's development work, including any delays or changes to the timing, cost and success of our product development activities and clinical trials including if we encounter difficulties enrolling patients in our clinical trials; the risks of delays in FDA and/or EU approval of our drug candidates or failure to receive approval; the risks related to commercializing any approved new pharmaceutical product including the rate and degree of market acceptance of our product candidates; development of our sales and marketing capabilities and risks associated with failure to obtain sufficient reimbursement for our products; risks related to our dependence on third parties including for conduct of clinical testing and product manufacture; our inability to enter into collaboration or alliances with partners; risks associated with protection of our intellectual property rights; and other risks and uncertainties affecting the Company including those described in the "Risk Factors" section included in our Annual Report on Form 10-K and other documents the Company files from time to time with the Securities and Exchange Commission. Any forward-looking statements contained in this presentation speak only as of the date hereof, and the Company undertakes no obligation to update any forward-looking statement, whether as a result of new information, future events or otherwise.

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MISSION



Y-mAbs Platforms to Drive Sustainable Long-term Value

Innovative Platforms

Antibodies and Vaccines

Radio-immunotherapy

SADA Platform Liquid RadiationTM 2022-2024 Milestones

Initiaton of Naxitamab label expansion into Osteosarcoma and adult indications

Resubmission and potential approval of omburtamab in 2022

Partnership potential for SADA Technology and antibody portfolio

Commercial Leverage

DANYELZA

(naxitamab-gqgk)

Anti-GD2 Antibody marketed for R/R High-Risk Neuroblastoma

omburtamab

BLA submitted

Anti-B7-H3 Antibody for CNS/LM from Neuroblastoma

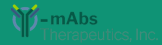
MAA submitted

Anti-B7-H3 Antibody for CNS/LM from Neuroblastoma

Capital Efficiency

\$156.7 million in cash and equivalents as of March 31, 2022

Three candidates
eligible for **Priority Review Vouchers** on
approval
(each currently
sell for ~\$100m)



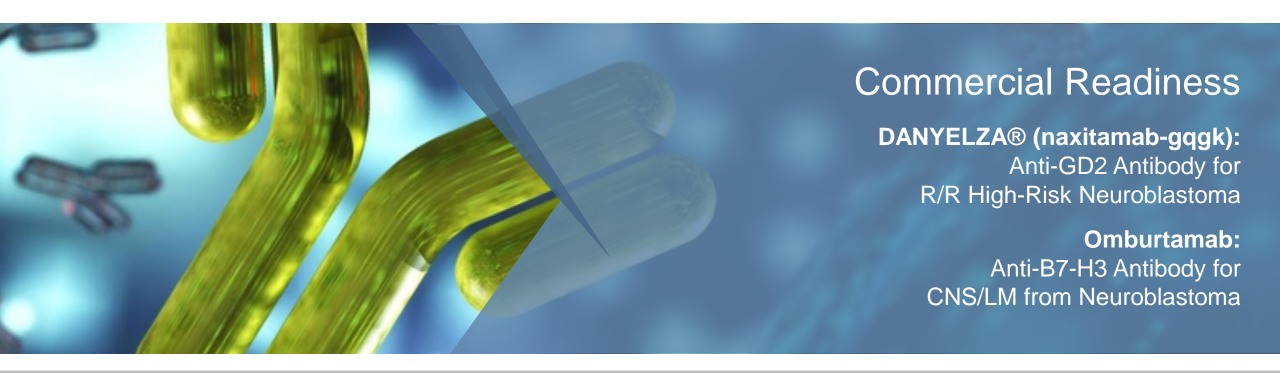
Strong Pipeline

Programs	Preclinical	Phase 1	Phase 2/Pivotal Study	Approved	Next Anticipated Milestones
Lood Condidates	DANYELZA (naxitamab-go	igk)			FDA approved
Lead Candidates	¹³¹ I-omburtamab		RPDD¹ ❖		BLA resubmitted Q1 2022, MAA submitted Q2 2021
Vaccine	GD2-GD3 Vaccine		RPDD ❖		Multicenter Phase 2 study being planned
Radiotherapy	¹⁷⁷ Lu-omburtamab-DTPA		RPDD 🕢		Medulloblastoma
SADA Technology	GD2-SADA				GD2 Positive solid tumors – IND filed December 2021. Additional INDs being prepared

¹ Indicates eligibility for a Priority Review Voucher (PRV) on approval







Commercial Opportunities – DANYELZA and omburtamab

Compound	Indication	Total Incidence per Year (US)	Addressable Patient Population per Year (US)
GD2	Neuroblastoma – 2 nd Line	300	300
DANYELZA (naxitamab)	Neuroblastoma – Front Line	800	450
	Osteosarcoma – 2 nd Line	450	200
	Neuroblastoma Metastatic to the Central Nervous System (CNS/LM from NB)	80	80
B7-H3 omburtamab	Diffuse Intrinsic Pontine Glioma (DIPG)	300	300
	Desmoplastic Small Round Cell Tumors (DSRCT)	100	100



DANYELZA: Only FDA-Approved Medicine for R/R NB Patients



FDA approval for patients with R/R Neuroblastoma (NB)

- Rapid infusion, less pain, fewer hospitalization days, high degree of treatment compliance
- Outpatient treatment



Neuroblastoma

- NB forms in certain types of nerve tissue. It most frequently starts from one of the adrenal glands but can also develop in the neck, chest, abdomen or spine.
- NB is the most common cancer in babies and the third-most common cancer in children



U.S. commercial launch performance

- Q1 net sales of \$10.5 million
- 34 active sites across the U.S.



Solid drivers of market uptake

- Pre-launch efforts drove market development and market access
- Significant engagement with key stakeholders





DANYLEZA: Primary and Secondary Refractory Patients

Study 12-230 (SIOP October 2019 - Investigator evaluated responses)

23 evaluable patients with primary refractory high-risk NB: 78% ORR

50% two-year progression free survival (PFS) was observed

Study population of 35 secondary refractory patients with relapsed NB resistant to salvage therapy: 37% ORR

36% two-year PFS was observed

Study 201 (SIOP October 2020 - Independent review assessment)

25 patients enrolled, of which 22 patients were evaluable: 68% ORR and 59% CR

Bone marrow (BM) clearance in subjects with positive BM at trial start: CR in BM was observed in 7 of 9 subjects.



High-Risk Neuroblastoma Frontline Treatment Regimes

Dinutuximab 10-20 hours infusion (x4 per week)

Induction

- Chemotherapy
- Stem Cell Collection
- Surgical Resection of Primary tumor

Consolidation

- High Dose Chemotherapy with ASCT
- Radiation Therapy

Maintenance

- Immunotherapy
- Cis-Retinoic Acid
- IL-2

Naxitamab app. 30-60 min infusion (x3 per week)

Induction

- Chemotherapy
- Stem Cell Collection
- Surgical Resection of Primary tumor

Consolidation

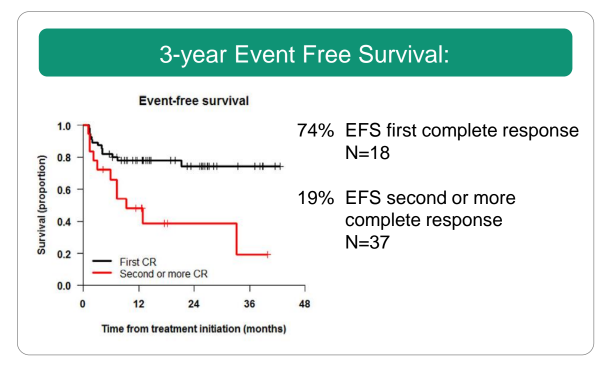
- High Dose Chemotherapy without ASCT
- Radiation Therapy

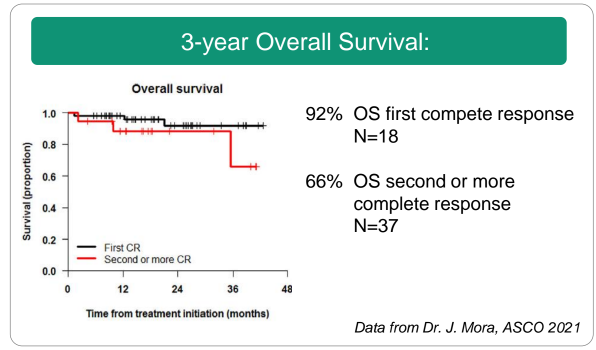
Maintenance

- Immunotherapy
- Cis-Retinoic Acid



Naxitamab: Frontline NB Data without Standard ASCT



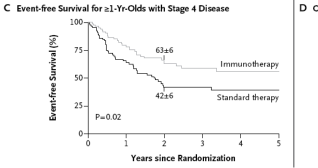


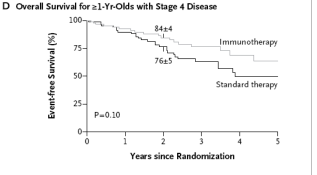


ORIGINAL ARTICLE

Anti-GD2 Antibody with GM-CSF, Interleukin-2, and Isotretinoin for Neuroblastoma

Yu, AL, et. al, New England Journal of Medicine, 2010





Naxitamab – Potential Label Expansion - Anticipated Study Initiation

Cancer indications	Treatable patient population (US)	GD2 expression	2020 2021 2022 2023 2024
Neuroblastoma FDA approval for R/R HR-NB in Nov 2020	800	~ 99-100%	R/R - High Risk Neuroblastoma Front-line HR Neuroblastoma Chemo combo in R/R HR NB
Relapsed/Recurrent Osteosarcoma	200	~88%	Relapsed Osteosarcoma Study 15-096 Pivotal RCT
Soft-Tissue Sarcomas	2,900 (1st-line population)	>90%	Phase 1/2
Triple Negative Breast Cancer	8,900 (2 nd line & 3 rd line <i>plus</i>)	>50%	Phase 1/2
Melanoma - Newly unresectable & met.	11,400 (2 nd line & 3 rd line <i>plus</i>)	>50%	Phase 1/2



Naxitamab: Key Takeaways

Addresses Significant Unmet Needs in R/R High-Risk NB • Potential to Expand to Broader Populations





Studies 12-230 and 201 formed primary basis of approval in November 2020





Granted ODD and BTD. Frontline studies ongoing



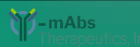


US commercialization in highrisk NB initiated. Chinese partnership with SciClone

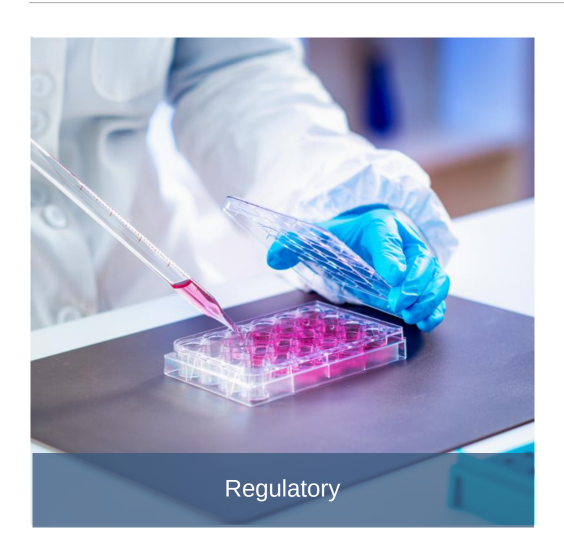




Multiple potential advantages over other GD2 targeting antibody-based therapies: Modest toxicity, shorter infusion time, ability to be administered in outpatient setting



Omburtamab: Regulatory Path to BLA Approval



U.S.

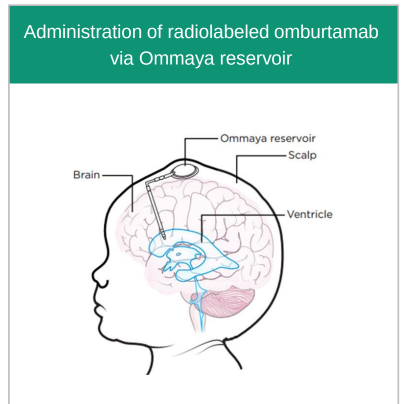
Studies 03-133 and 101 formed basis of BLA resubmission in Q1 2022

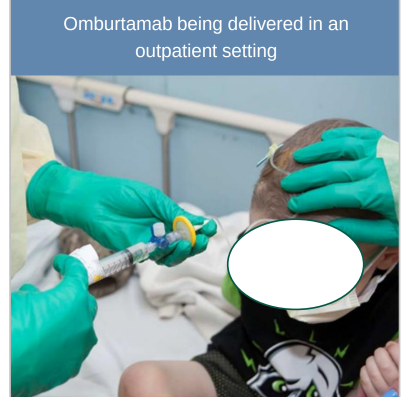
Europe:

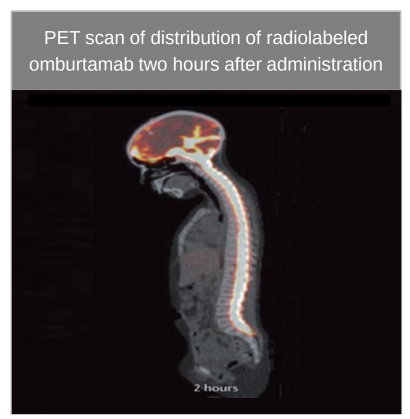
Marketing Authorization Application submitted Q2 2021

Omburtamab: Delivered in an Outpatient Setting – 2 Doses per Patient

CNS/LM from NB patients



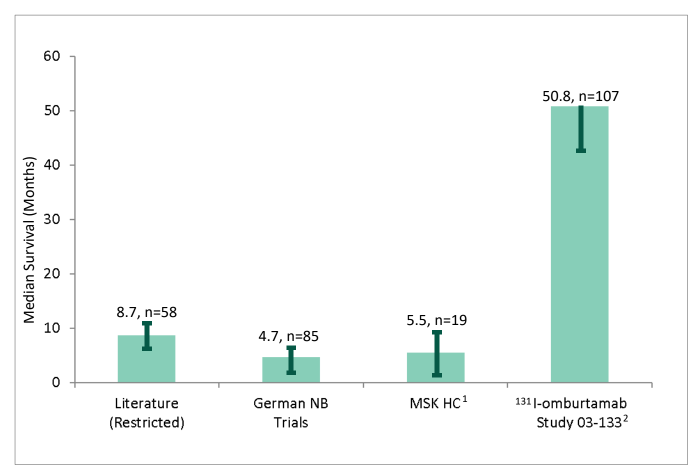




After induction treatment including all or some of the three treatments (chemotherapy, surgery, and radiation) patients will receive radiolabeled omburtamab

Omburtamab: Clinical Overview

Studies 03-133 and 101: 131 I-omburtamab Improves Survival in CNS/LM from NB Patients

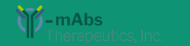


Number of patients in the full analysis set	24		
Objective Radiographic Response (CR and PR), N (%)	4 (40.0)		
[95% CI*]	[12.2;73.8]		
Best Overall Radiographic Response			
Complete response	2 (20.0)		
Partial response	2 (20.0)		
Stable disease	5 (50.0)		
Progressive disease	1 (10.0)		
Total	10 (100.0)		
No evidence of disease / Not evaluable (N)	14		
N: Number of subjects, %: Percentage of subjects Best overall radiographic response is assessed at Week 26 by independent Review of images.			

Results confirm the direct anti-tumor effect of ¹³¹I-omburtamab

Disease Control at Week 26 in 9 out 10 pts (90%)

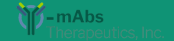
² ¹³¹I-omburtamab = Patients with CNS/LM treated under Study 03-133



¹ MSK HC = neuroblastoma patients with CNS/LM treated at MSK prior to 2003

Omburtamab: Label Expansion Through Broad Clinical Platform

Omburtamab (B7-H3)	Phase 1	Phase 2/Pivotal Study	Highlights
Accelerated Pathway	Phase 2: CNS/LM from NB (Pedia	atric) – Study 101	Multi-center PK study; resubmission of BLA in Q1 2021 – MAA submitted in Q2 2021
	Phase 1: CNS/LM – Study 03-133		MSK single-center efficacy data
Label Expansion	Phase 1: DIPG – Study 11-011		Study update presented at ASCO 2021
	Phase 2: DIPG multi-center		Multi-center study, IND filed Q4 2021
	Phase 2: DSRCT – Study 19-182		Study update from Phase 1 presented at CTOS Nov 2019



Omburtamab: Key Takeaways

Addresses Significant Unmet Needs and has the Potential to Expand its Application to Broader Populations





No approved products for patients with R/R NB who have CNS/LM from NB

Goal of treatment is generally palliative





Granted ODD, BTD, and RPDD; May qualify for a sBLA for DIPG and DSRCT assuming positive pivotal data





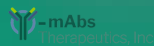
Demonstrated median OS of approximately 51 months

Historical median OS of ~6-9 months and no expected five-year survival

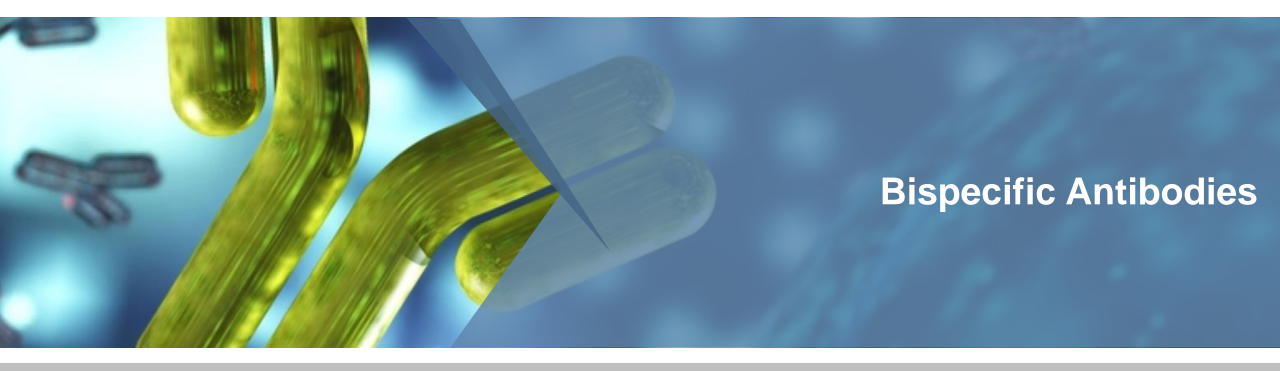




Studies 03-133 and 101 formed primary basis for BLA resubmission for CNS/LM from NB in Q1 2021. MAA submitted in Q2 2021. Large potential market opportunity for the treatment of LM from tumors expressing B7-H3







CD33xCD3

Non-clinical package

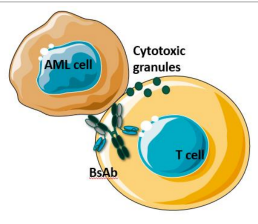
- Full non-clinical package submitted to support Y-mAbs IND.
- No FDA hold issues identified.

Clinical

- Clinical Protocol Submitted for Phase 1.
- No FDA hold issues identified.
- Collaboration with COG who has identified approx. 15 sites

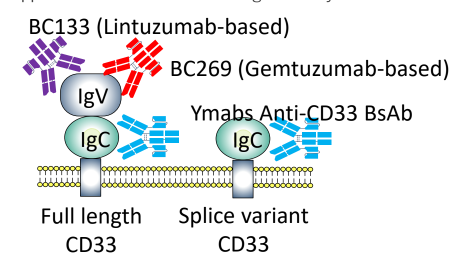
IND

Expect IND to open Q2 2022



Y-mAbs Biclone CD33xCD3 binds to IgC region allowing binding to both full length and splice variant (rs12459419; reduced / no expression of IgV domain)

Approved CD33 mAb binds to region V only

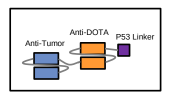


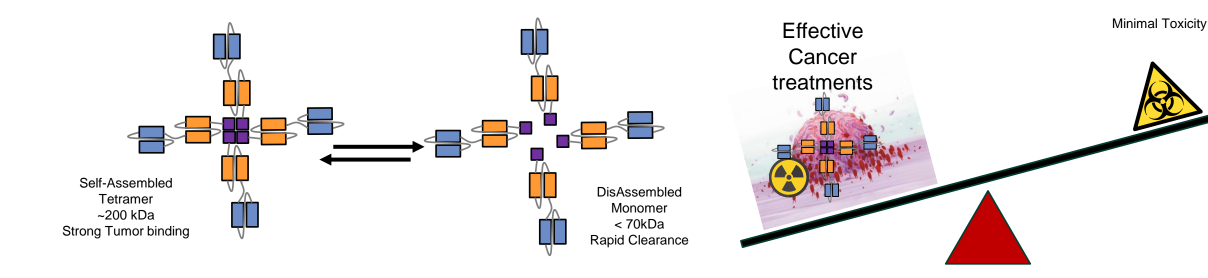




SADA Technology - High affinity for tumor targets and rapid clearance from blood stream

SADA domains uniquely selected to allow proteins to change size based on concentration





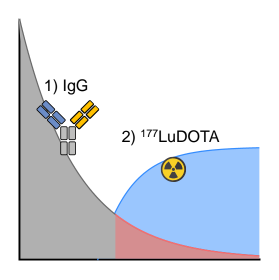
Adapted from Santich et al. Clin Canc Res 2020



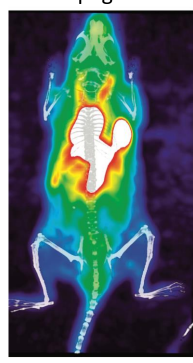
SADA has high tumor uptake with minimal exposure to all other tissues

Conventional antibody's persistence in blood stream leads to substantial unwanted exposure

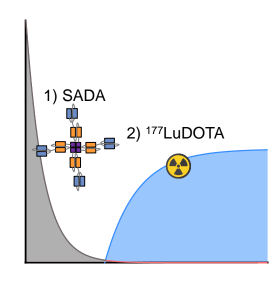
Pretargeted Radioimmunotherapy (2-step, TI = ~20:1)



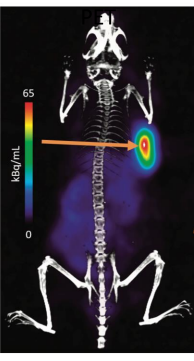
2-step IgG PET



Pretargeted SADA Technology (2-step, TI > 50:1)



2-step SADA

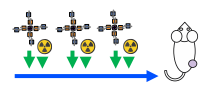


Adapted from Santich et al. Clin Canc Res 2021

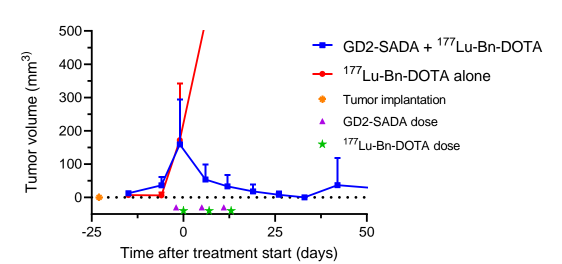


GD2-SADA in established mouse tumor models

PDX models; NB and SCLC

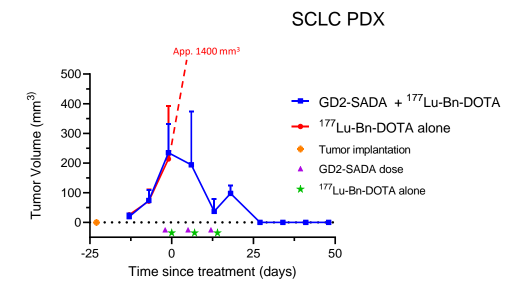


Neuroblastoma PDX



Model: BRG mice with s.c. PDX

Doses: 1x weekly for 3 weeks with 48hr interval,



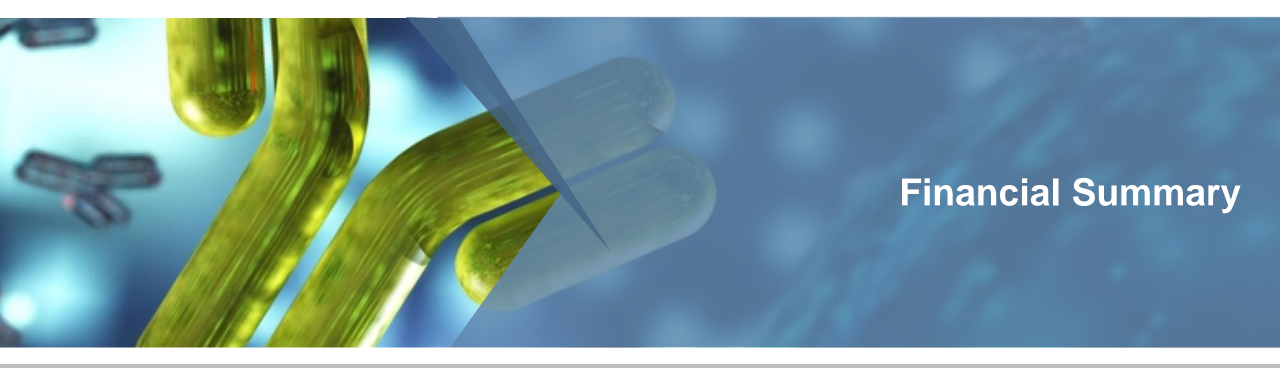
Model: BRG mice with s.c. PDX

Doses: 1x weekly for 3 weeks with 48hr interval

Adapted from Santich et al. Clin Canc Res 2020 and unpublished data







Strong Financial Position





Follow on: November 2019

\$144 Million

Follow on: February 2021

\$115 Million

Non-dilutive cash: January 2021

\$62 Million from PRV sale



1 PRV

PRV sold for \$105 Million Y-mAbs retained 60% of net proceeds \$62 million

3 RPDDs

Received for leading compounds

\$156.7 Million

of cash and cash equivalents as of March 31, 2022



Investment Highlights

	Pediatric	Adult
DANYELZA (naxitamab-gqgk) and GD2-GD3 Vaccine	High-Risk NeuroblastomaOsteosarcoma	Osteosarcoma
Omburtamab	 CNS/LM from NB (¹³¹I) DIPG and DSRCT (¹³¹I) 	
SADA Platform	Neuroblastoma	 Small Cell Lung Cancer Colon Cancer, Prostate Cancer, Breast Cancer



