

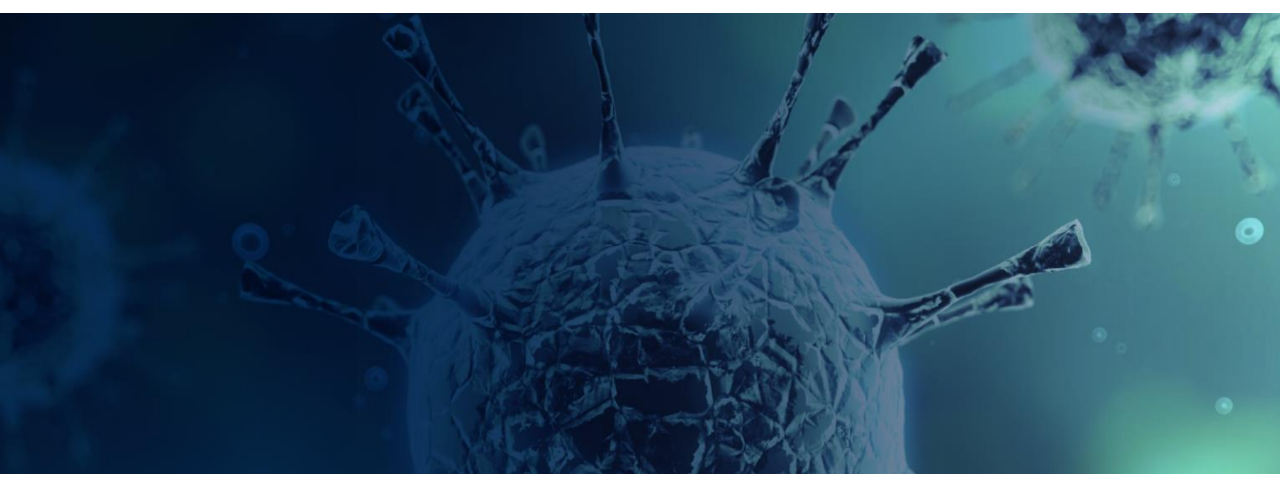
Developing Next Generation Programmed T Cell Therapies

August 2021

Disclaimer



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Lead Clinical Programs
Striving for best-in-class therapies

Focused on delivering obecel, a potentially transformational treatment for Adult Acute
Lymphoblastic Leukemia (ALL), as well as exploring activity in additional B-cell malignancies

Full data for obe-cel (FELIX) trial in adult expected in 2022

Obe-cel data in broader NHL indications expected in Q4 2021, Obe-cel data in PCNSL in Q1 2022, AUTO1/22 in pALL expected in Q4 2021

- Additional value steps in T cell lymphoma and first solid tumor indication
- Broad preclinical pipeline of next generation programs expected to transition to clinical stage in 2021/2022
- Scalable, fully enclosed manufacturing platform

Broad pipeline of clinical programs

Aut•lus

Designed to address limitations of current T cell therapies

PRODUCT	INDICATION	TARGET	PHASE 1/2	PIVOTAL*
obe-cel	Adult ALL	CD19	ALLCAR19	FELIX
obe-cel	NHL [†]	CD19	ALLCAR19	
obe-cel	PCNSL ^{††}	CD19	CAROUSEL	
AUTO1/22	Pediatric ALL	CD19 & CD22	CARPALL	
AUTO3	DLBCL	CD19 & CD22	ALEXANDER	To be partnered
AUTO4	TRBC1+ Peripheral TCL	TRBC1	LibrA T1	

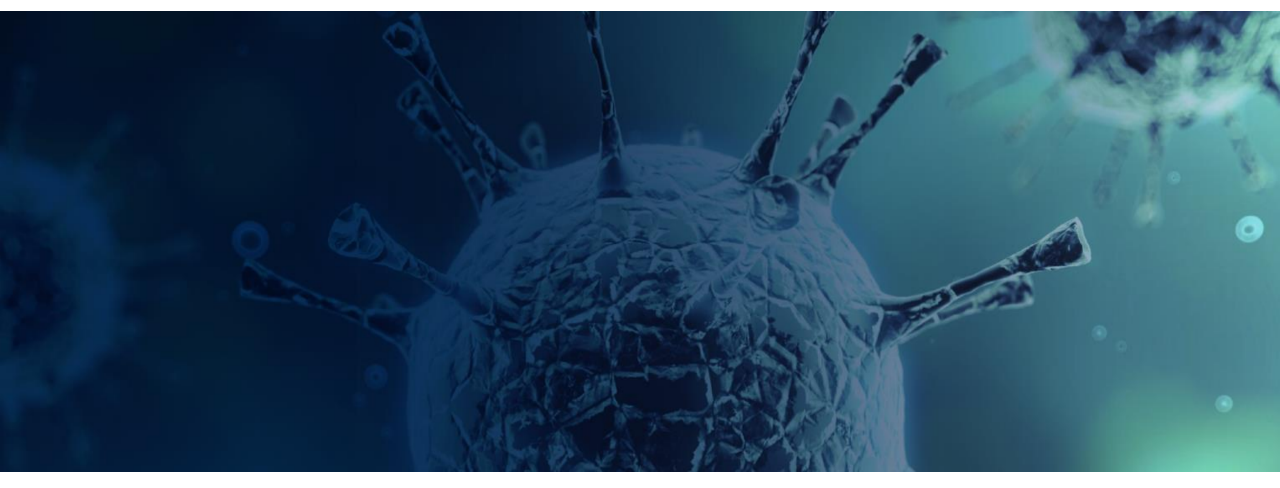
B Cell Malignancies



^{*}Subject to confirmation by regulatory authorities

[†] Non-Hodgkin lymphoma

^{††}PCNSL = Primary CNS Lymphoma



Adult Acute Lymphoblastic Leukemia
Obe-cel— Potential as a standalone therapy

No approved CAR T therapy for adult ALL patients



Successful therapy requires high level of activity and sustained persistence paired with good tolerability

ALL is a significant opportunity

Up to **8,400*** new cases of adult ALL diagnosed yearly worldwide

Estimated R/R patients in US & EU **3,000** addressable patient population in last line setting

HIGH UNMET MEDICAL NEED

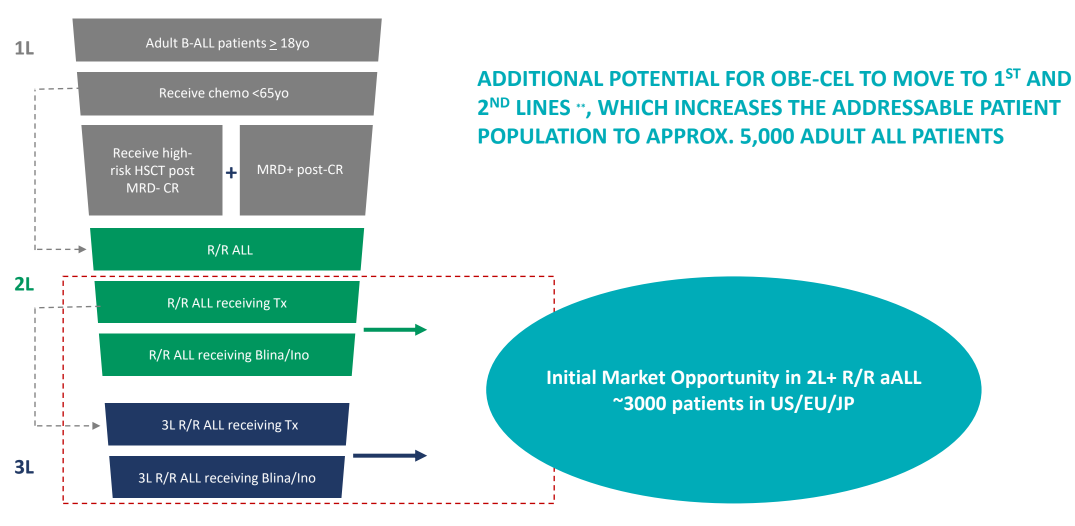
- Combination chemotherapy enables 90% of adult ALL patients to experience CR, but only 30% to 40% will achieve long-term remission
- Median overall survival is < 1 year in r/r ALL
- Only redirected T cell therapy for adult patients is blinatumomab
- CAR T therapies are highly active, but require subsequent allograft to achieve durability
- Patients are generally more fragile with co-morbidities, yet CAR T toxicities in this setting have been notable with high incidences of severe CRS and cases of fatal neurotoxicity
- Opportunity to progress to earlier lines of treatment and expand the addressable patient population

OBE-CEL GRANTED ORPHAN DRUG DESIGNATION BY FDA FOR ALL, PRIME DESIGNATION IN R/R B-ALL BY EMA AND ILAP DESIGNATION BY MHRA IN ADULT R/R B-ALL

^{*}SEER and EUCAN estimates (respectively) for US and EU epi

Adult ALL is a promising commercial opportunity with limited competition





^{*}Company estimate, based on US, EU5 and Japan

^{**}Subject to successful clinical progress

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Key features of a successful CAR T Cell Therapy for adult ALL



Obe-cel is uniquely placed to address current limitations of therapy

Challenge	Product Property	CAR T Feature	Benefit
Fast proliferating disease	Very high level of anti- leukemic activity	Rapid CAR T mediated kill and high level of CAR T expansion	High response rates
Almost stem cell like nature of leukemic cells	Sustain long term pressure on leukemia	Long CAR T persistence	Durable responses
Poor patient condition	Good tolerability	Minimize high grade CRS and NT	Manageable AE profile

Obe-cel has potential for transformational outcomes in Adult ALL



Data cut-off date May 17, 2021

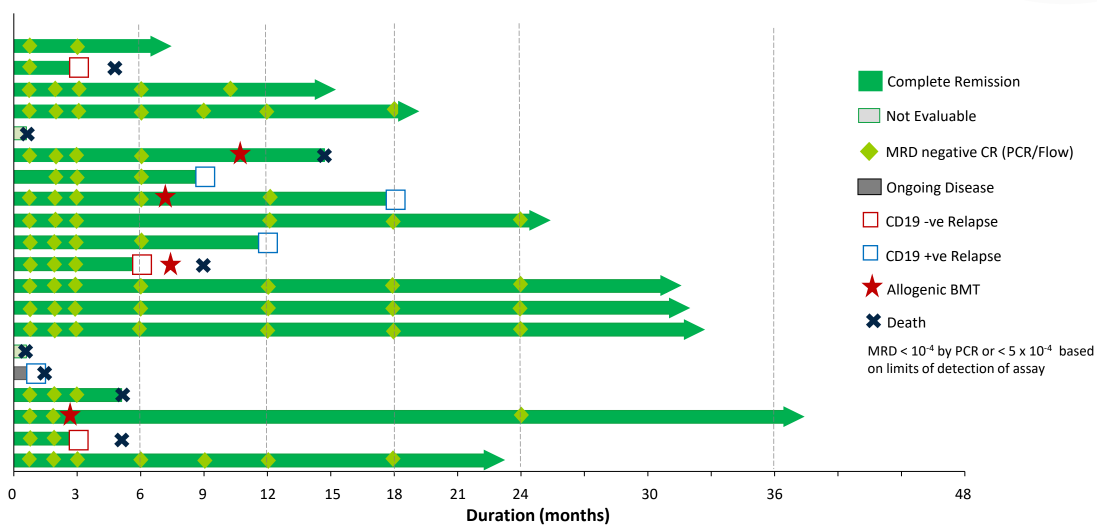
- High level of sustained CR achieved without subsequent stem cell transplant
- Durability of remissions highly encouraging
 - Across all treated patients, event free survival (EFS) at twelve and twenty-four months of 50%
- Obe-cel well tolerated, despite heavily pre-treated patients with high disease burden
 - No patients experienced ≥ Grade 3 cytokine release syndrome (CRS)
 - 20% of patients experienced any grade ICANS*, swiftly resolved with steroids
- Phase 1b/2 potential pivotal study underway, expect full data in 2022
- Adult ALL represents a sizeable market opportunity addressable with focused commercial footprint

¹⁰

Obe-cel shows sustained stand-alone activity in adult ALL patients



ALLCAR-19 Phase 1 EHA data cut - May 17 2021

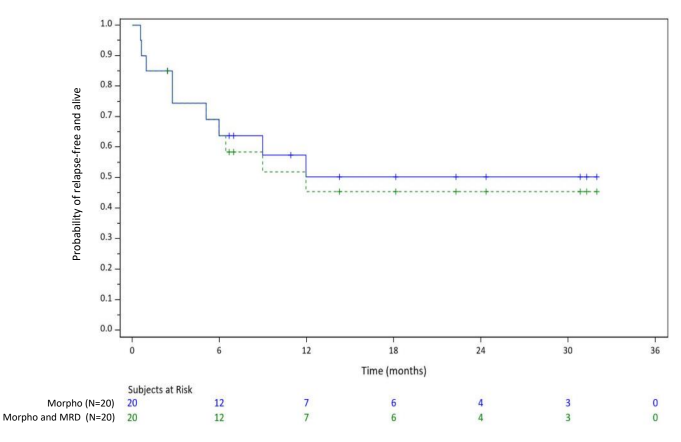


Obe-cel morphological event-free survival of 50.2% at 24 months



MRD and morphological EFS curves are superimposable with a plateau seen from 12 months

Morphological and Molecular EFS among all infused patients in ALLCAR19



		All infused patients	Closed Process
	N	20	14
	ORR	85%	93%
	MRD Neg CR	85%	93%
DOR			
	Median	Not reached	Not reached
	12 months	64%	64%
Morph. EFS			
	Median	Not reached	Not reached
	12 months	50.2%	60%
	24 months	50.2%	60%
Molecular EFS			
	Median	12 months	Not reached
	12 months	45%	54%
	24 months	45%	54%

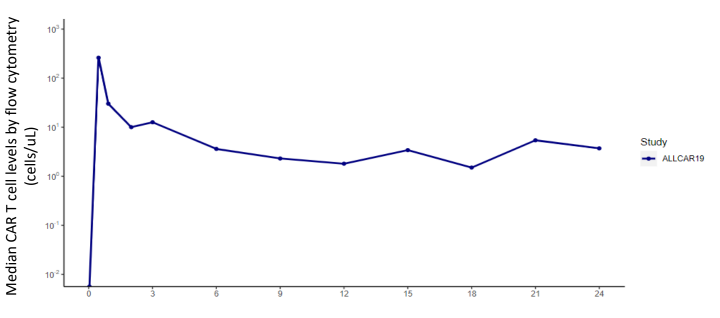
Event for morphological EFS = death or morphological relapse Event for molecular EFS = death, morphological relapse, or molecular relapse (i.e. MRD > 0.01%) Data Cut-off 17-May-2021

Obe-cel expansion characteristics support its differentiated profile



Data so far points to a transformational product with ability to maintain pressure on tumor

Median CAR T cell levels in peripheral blood



Time from	CART	T infusion	(months)

	ALLCAR-19 Phase 1
N	20
CRS Any Grade	55%
CRS Grade ≥ 3	0
NE / ICANS Any Grade	20%
NE / ICANS Grade ≥ 3	15%
Treatment for CRS and/or ICANS	
Tocilizumab	35%
Steroids	20%
Vasopressor	0



	ZUMA-3¹ Phase 2
	Tecartus
N	55
ORR (CR/CRi)	71%
EFS	~45% (12 m), ~25% (18 m)
CRS ≥ Grade 3†	24%
Neurotox ≥ Grade 3†	25%
Other notable observations	40% vasopressor use

Standard of Care		
Blinatumumab ²	Inotuzumab ³	
271	109	
44%	80.7%	
31% (6 m)	mPFS 5m	
3%	0%	
13%	0%	
NA	14% Hepatic VoD	

- Approximately 50% of blinatumomab and inotuzumab patients received subsequent HSCT
- Veno-Occlusive Disease (VoD) during treatment and following subsequent HSCT, with the latter causing a higher post-HSCT non-relapse mortality rate, has limited inotuzumab uptake

Duration of follow-up is calculated from CAR T infusion to data cutoff. EFS for ZUMA-3 were estimated based on the KM curve#

- 1. Shah et al. Lancet 2021
- 2. Kantarjian et al., 2017/ USPI (product label)
- 3. Kantarjian et al., 2016/ USPI (product label)

Obe-cel potentially differentiated on efficacy, durability and safety



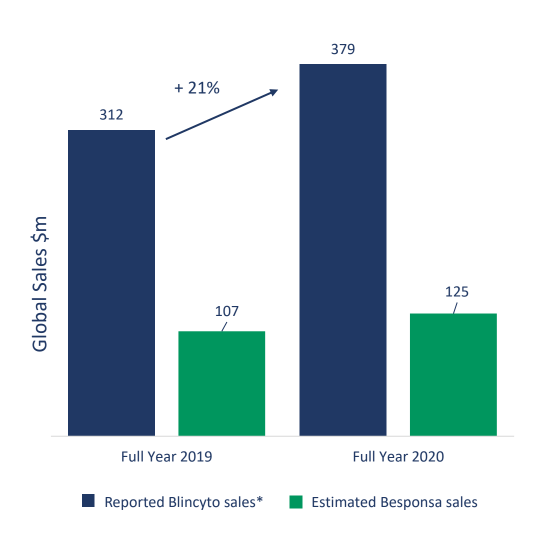
Unique CAR T design drives differentiated product profile

○ Obe-cel has a high level of MRD-negative CR (85%)
 Morphological EFS for obe-cel at 24 months was 50.2%
 Long term CAR T persistence drives durability of effect
 Obe-cel has a favorable safety profile

Obe-cel could launch into an expanding market

Aut•lus

Benefitting from a potentially superior clinical profile



- Blincyto sales price estimated to be \$178k[±] (based on 2 cycles)
 resulting in approx. 2,100 commercial patients (of which approx. 85% are >18 years **)
- Growth attributed by Amgen* to broader uptake and expansion in community settings, continued strong growth at 29% y-o-y for Q4
- O Kymriah is priced at \$475k in pediatric ALL. Breyanzi (lisocabtagene maraleucel) is priced at \$410k in DLBCL^{±±}.
- Breyanzi and other CAR T cell therapies are expanding delivery center footprint
- Tecartus (brexucabtagene autoleucel) is expected to establish CAR T use in adult ALL
- Obe-cel expected to have a superior clinical profile
 - Has potential to be the only curative therapy with tolerability profile to take advantage of expanding delivery footprint

^{**} Komodo Health 2015 – 2020

[±] https://www.medscape.com/viewarticle/836879

^{± ±} Bristol Myers finally wins FDA approval for cancer cell therapy | BioPharma Dive

Preliminary Ph1 data supports development as a standalone therapy



Obe-cel is the first Autolus program to move into a pivotal program

Pivotal program,
FELIX, in adult ALL
enrolling with full
data targeted in 2022

CTA approved by the MHRA in January 2020 and US IND accepted by the FDA in April 2020

- Ph1b run-in component, prior to single arm Ph2 pivotal study
- 100 relapsed/refractory adult ALL patients
- Primary endpoint: Overall Complete Response Rate (CR/CRi)
- Secondary endpoints: include MRDnegative CR EFS and DoR

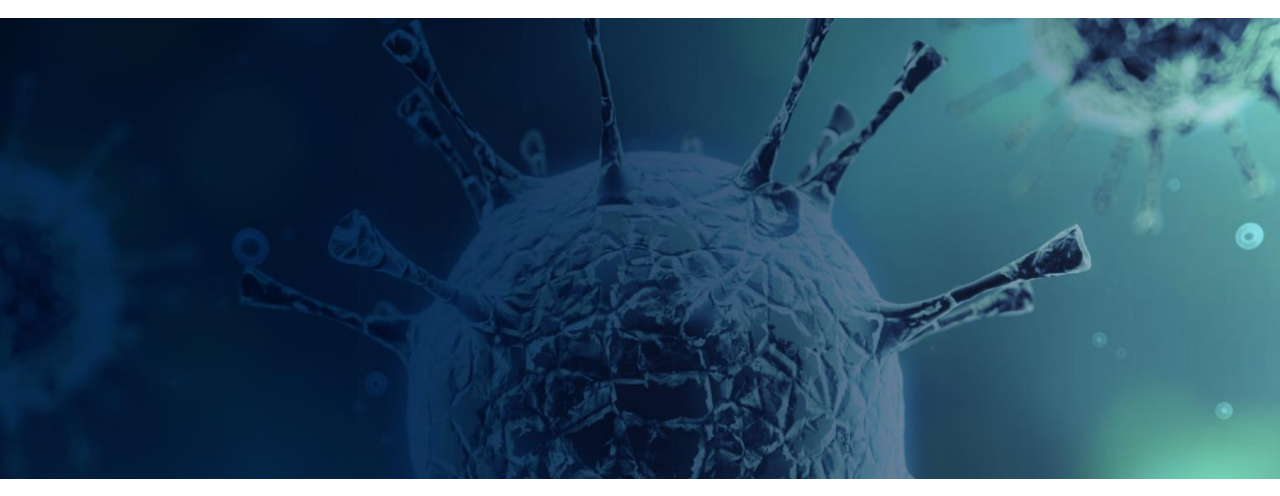
Unique profile of obe-cel offers potential across broader indications



Evaluation of obe-cel activity in additional B-Cell malignancies to capitalize on potential market opportunity

PRODUCT	INDICATION	TARGET	PHASE 1	PHASE 1B/2
Obe-cel	Adult ALL	CD19	ALLCAR-19 *	FELIX
Obe-cel	B-NHL & CLL	CD19	ALLCAR-19 Ext *	
Obe-cel	Primary CNS Lymphoma	CD19	CAROUSEL *	
AUTO1/22	Pediatric ALL	CD19 & CD22	CARPALL *	

OPPORTUNITY TO PURSUE IN EARLIER LINES OF THERAPY AND INDICATIONS OF ADULT ALL



Pediatric ALL

AUTO1/22 CD19 & CD22 targeting CAR-T for r/r pediatric ALL



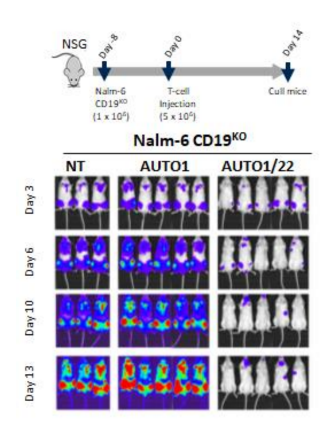
Adding a high sensitivity CD22 CAR to AUTO1 to minimize relapses due to CD19 antigen loss

Primary driver for relapse in pALL is CD19 antigen loss

	pALL		
	*AUTO1	#Kymriah	
n	14	75	
CR Rate	86%	81%	
EFS 12m	54%	50%	
EL2 TZIII	(95% CI, 24 to 76)	(95% CI, 35 to 64)	
No. of CD19 negative relapses	5/6	15/22	
CRS ≥ G3	0%	47%	
NTX ≥ G3	7%	13%	

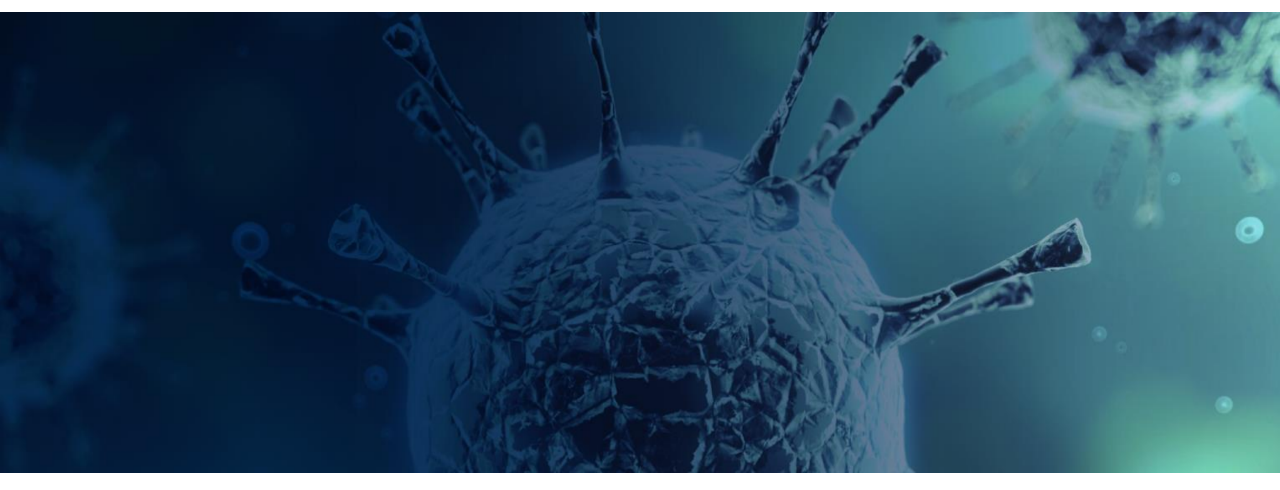
^{*}Ghorashian S, Pule MA, Amrolia P et al. Nature Medicine 201

AUTO1/22 Dual Targeting CAR T to CD19 and CD22



[#]Maude et al., NEJM 2018





Indolent NHL

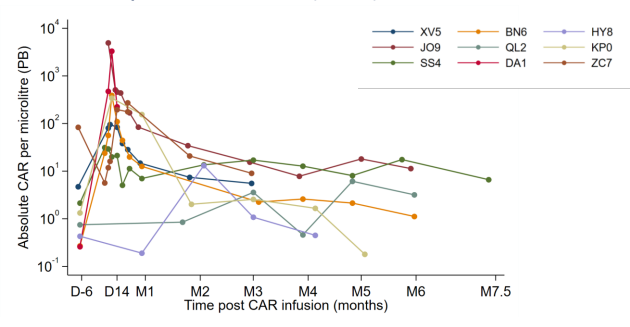
obe-cel



Obe-cel shows excellent T cell expansion and engraftment with a manageable safety profile

Baseline Characteristics	N=9
Median age, years (range)	56 (39 - 68)
Gender	7M/ 2F
DiseaseFollicular LymphomaMantle Cell Lymphoma	7(78%) 2 (22%)
Lines of treatment	3 (2-5) 4 (44%) 1 (11%)
Stage of disease at screening • Stage I/II • Stage III/IV	0/9 9/9
Bridging therapy Chemotherapy alone Radiotherapy + steroids Chemo + Radiotherapy Nil	7 2 0 0

CAR T Cell Expansion in Cohort D (B-NHL)

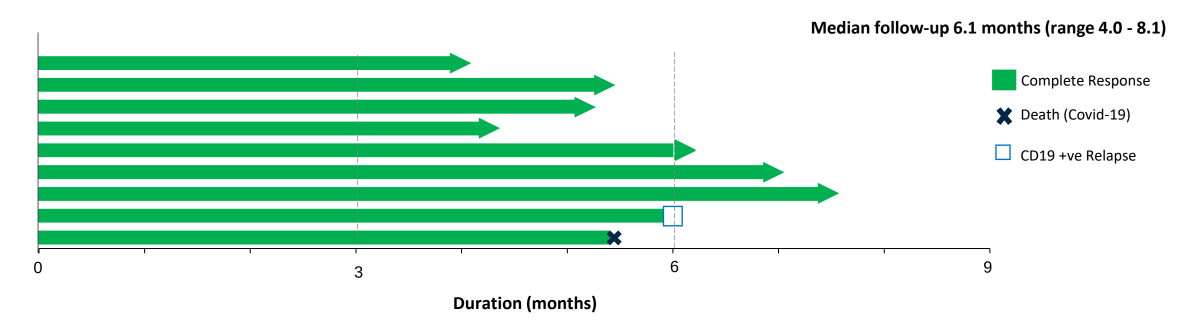


Immunotoxicity in Cohort D (B-NHL)

CRS (ASTCT criteria*)	Neurotoxicity (ICANS)
• CRS (any) in 5/9	• ICANS in 0/9
• Grade 2 in 1/9	• Grade 2 in 0/9
• ≥ Grade 3 CRS in 0/9	• Grade 3 in 0/9
Tocilizumab used in 2 patients	



All patients treated achieved a metabolic Complete Response (CR)

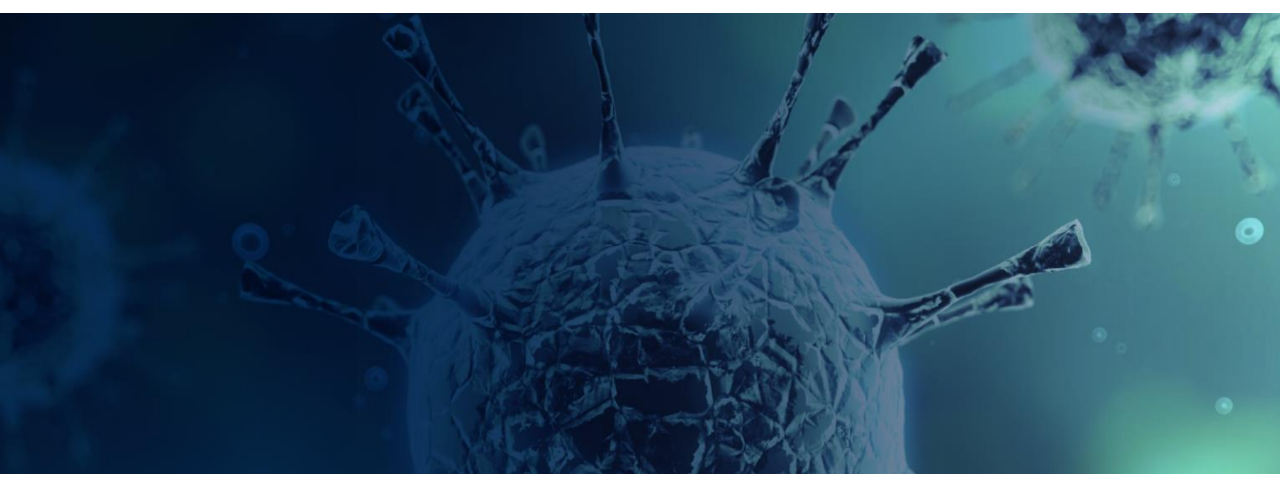


- 9/9 patients in the indolent B-NHL cohort achieved metabolic CR by month 3
- \circ 8/9 disease-free at last follow-up (median F/U = 6.1 months; range 4.0 8.1m)
- 1/9 patients died on study from COVID-19 whilst in remission at month 6 of follow-up
- 1/9 relapsed with small volume subcutaneous CD19+ disease, salvaged with radiotherapy
- o 0/9 patients experienced ICANS of any grade or ≥ grade 3 CRS

Summary and next steps for obe-cel outside of adult ALL



Initial obe-cel efficacy data in the iNHL cohort is encouraging
 Obe-cel continues to show a consistent and favorable safety profile across all indications evaluated
 Additional cohorts being explored, and further data planned for Q4 2021
 CAROUSEL study in PCNSL open with data expected in Q1 2022
 AUTO1/22 (CARPALL study) in pediatric ALL open with data expected in Q4 2021



Diffuse Large B Cell Lymphoma AUTO3 — tailored for DLBCL

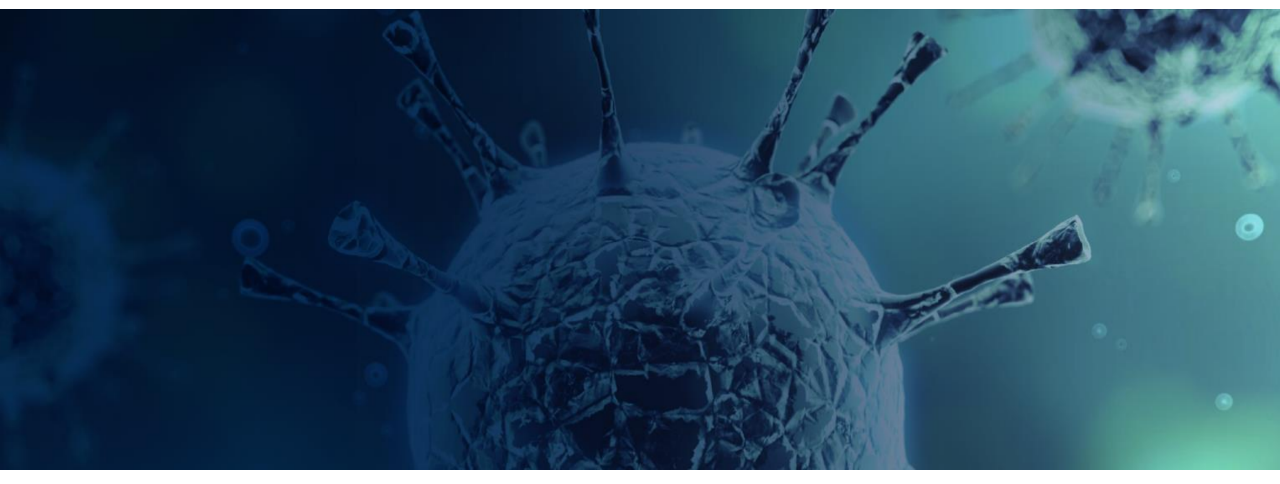
AUTO3 continues to show differentiated product profile in DLBCL

Aut lus

Data presented at ASH 2020, with data cut-off date of October 30, 2020

- Key Phase 1 observations:
 - High level of complete remissions (CR) of 51% overall
 - At the highest dose level of 450M cells the CR rate was 73%
 - Very low levels of high-grade CRS and neurotoxicity
 - AUTO3 administration together with the pembrolizumab dosing regimens (D-1 and D14/D35/D56) were well tolerated
 - Among the five patients who achieved a CR having received 3 doses of pembrolizumab, none had progressed as of the data cut-off date
 - Demonstrated feasibility to administer AUTO3 in outpatient setting

- Potential path forward for development of AUTO3
 - Phase 2 designs under evaluation:
 - 3L r/r DLBCL setting
 - 2L/3L transplant ineligible DLBCL setting
 - Planned Phase 2 dosing regimen
 - Dose range of 150M to 450M cells, as patients benefitted from therapy at 150M, 300M and 450M cell dose levels
 - 3 doses of pembrolizumab with a schedule of D-1, D28, D56
 - Implement manufacturing process enhancements (incl. stable cell line for vector manufacturing)



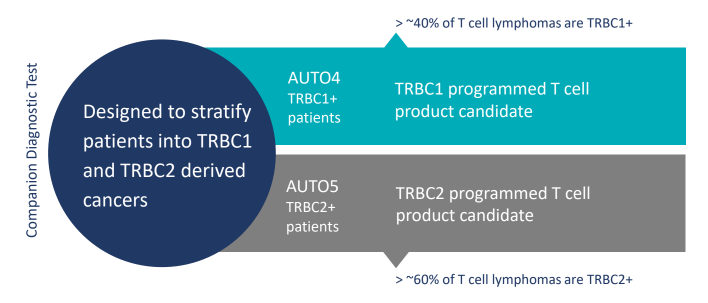
T Cell Lymphoma AUTO4 and AUTO 5 — tailored for T Cell Lymphoma

T Cell Lymphoma



No standard of care after first relapse and no T cell therapy approved

AUTOLUS USES THREE KEY ELEMENTS TO ADDRESS T CELL LYMPHOMAS—AUTO4, AUTO5 AND A COMPANION DIAGNOSTIC TEST



- T cell lymphoma is an aggressive disease with a very poor prognosis for patients
- Median 5 yrs OS: 32%
- Standard of care is variable and often based on high-dose chemotherapy and stem cell transplants
- A large portion of T cell lymphoma patients are refractory to or relapse following treatment with standard therapies
- T cell lymphomas have not, so far, benefited from advances in immunotherapeutic approaches
- AUTO4 Phase 1 interim data expected in H1 2022
- AUTO5 to enter Phase 1 study in H1 2022

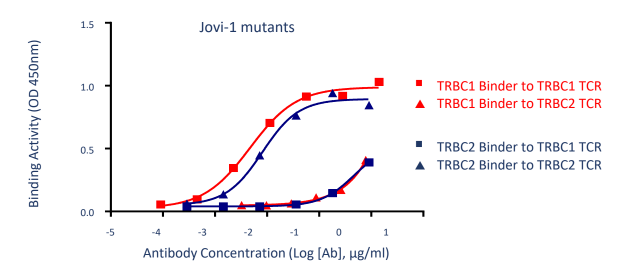
Unique targeting of TRBC1 & TRBC2 opens new therapeutic approach AUTO4/5 in Peripheral T Cell Lymphoma

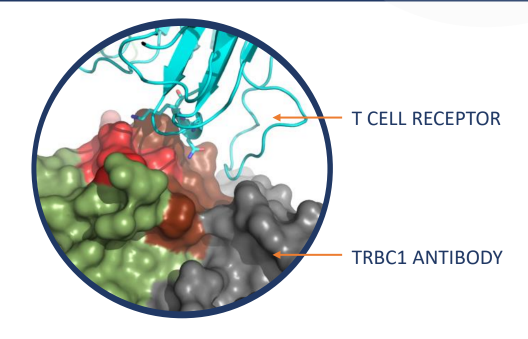


DIFFERENCES BETWEEN TRBC1 AND TRBC2 ARE SMALL

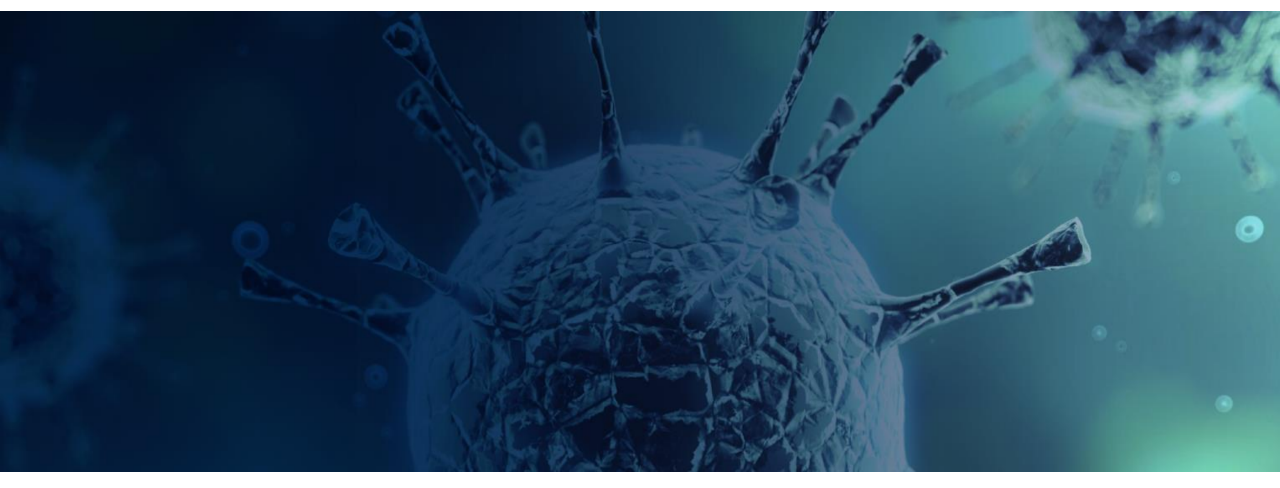
		NK-KN 4/5	F-Y 36
TRBC1	1		HTQKATLVCLATGF PDHVELSWWVNGK
TRBC2	1	EDL <mark>KN</mark> VFPPEVAVFEPSEAEIS	HTQKATLVCLATGF <mark>Y</mark> PDHVELSWWVNGK
TRBC1	51	EVHSGVSTDPQPLKEQPALNDS	RYCLSSRLRVSATFWQNPRNHFRCQVQF
TRBC2	51	EVHSGVSTDPQPLKEQPALNDS	RYCLSSRLRVSATFWQNPRNHFRCQVQF
TRBC1	101	YGLSENDEWTQDRAKPVTQIVS	AEAWGRADCGFTS <mark>V</mark> SYQQGVLSAT AEAWGRADCGFTSESYQQGVLSAT
TRBC2	101	YGLSENDEWTQDRAKPVTQIVS	AEAWGRADCGFTSESYQQGVLSAT
			V-E 135

ANTIBODY BINDING DATA





- AUTO4 clinical study, LibrA T1, in progress
- AUTO5 in late preclinical development
- Preclinical study package demonstrating selective binding and anti-tumor activity of TRBC1 and TRBC2
 CARs in vitro and in vivo

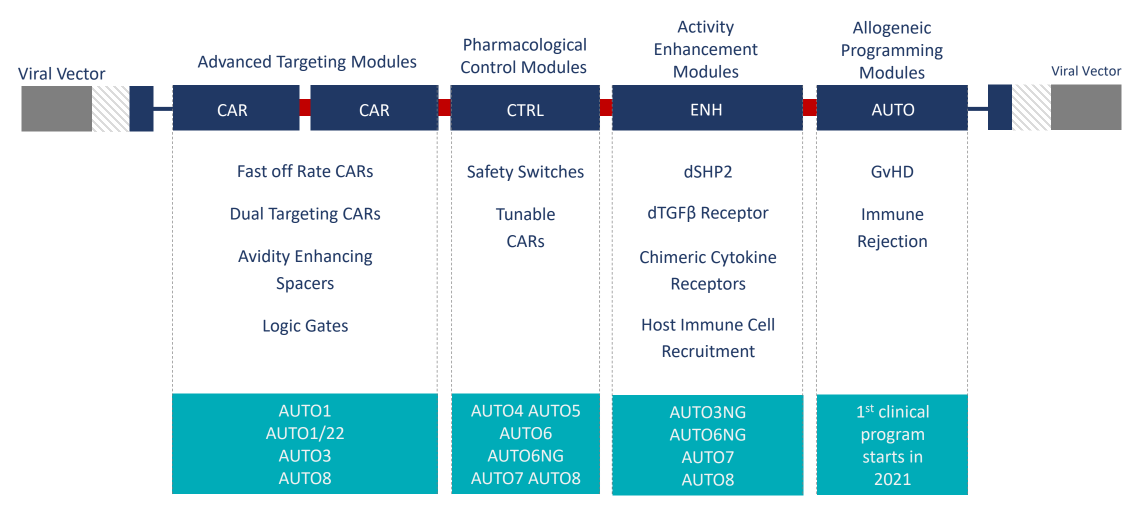


Pipeline

A broad portfolio of next generation modular T cell therapies

A broad toolkit which is core to our strategy of modular innovation Advanced T cell programming





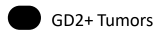
Broad pipeline of next generation programs

Aut•lus

Designed to address limitations of current T cell therapies

PRODUCT	INDICATION	TARGET	PRECLINICAL	PHASE 1*
AUTO1/22 **	Pediatric ALL	CD19 & CD22		Started Q4 2020
AUTO5	TRBC2+ Peripheral TCL	TRBC2		H1 2022
AUTO6NG**	Neuroblastoma; Other tumor types	GD2		H1 2022
AUTO7	Prostate Cancer	PSMA		H1 2022
AUTO8 **	Multiple Myeloma	BCMA & CAR X		H2 2021





Prostate Cancer



B Cell Malignancies

^{*}Planned Trial Initiations

^{**} Collaboration with UCL
NG = Next Generation, SCLC = Small Cell Lung Cancer

Modular approach designed to enhance AUTO6NG for solid tumor environment



Next generation programs powered by our proprietary technology toolbox



To provide anti-tumor activity and potential to help address neurotoxicity and pain syndrome

AUTO6

SAFETY SWITCH
To eliminate the therapy in the event of unexpected toxicities

dSHP2

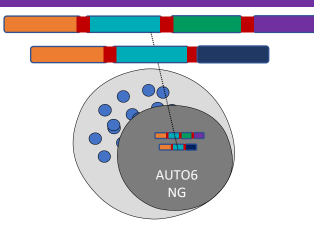
To overcome multiple checkpoint pathways

Cytokine Signal

IL7 CCR chimeric protein designed to improve CAR T cell persistence

dnTGFβRII Receptor

To overcome inhibitory effect of TGFβ in microenvironment



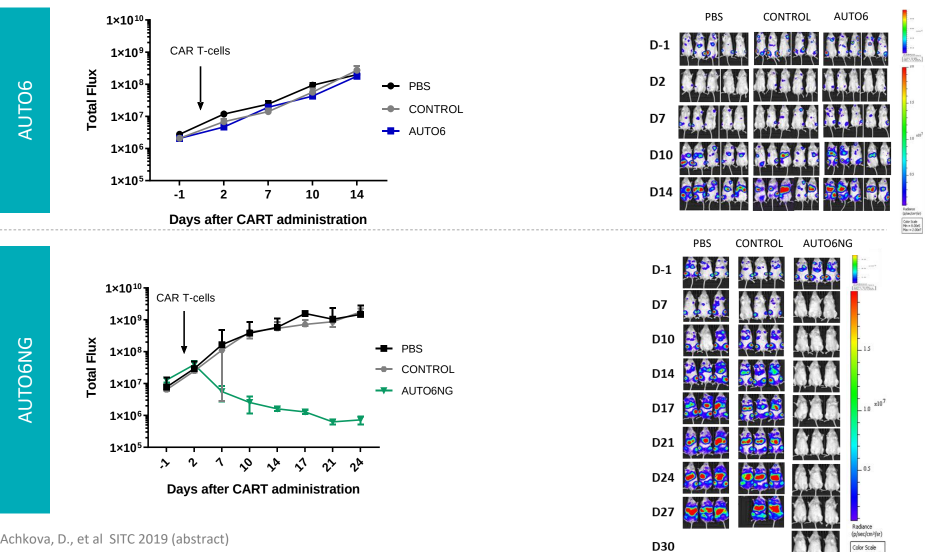
AUTO6NG:

- Utilizes GD2 CAR from AUTO6, and is further enhanced to address persistence, control and tumor defenses
- Targeting neuroblastoma, osteosarcoma, melanoma and small cell lung cancer amongst others

AUTO6NG exhibits potent anti-tumor activity in preclinical model



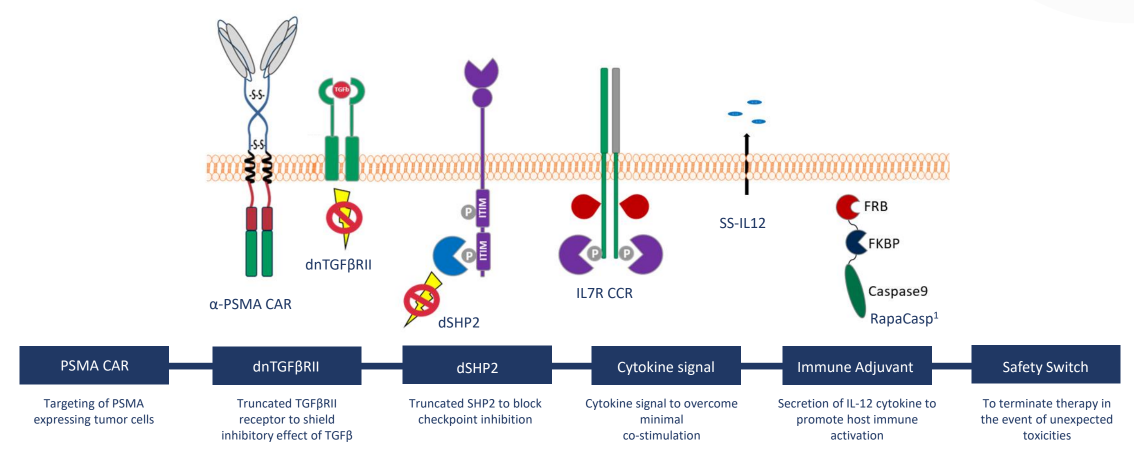
Extends survival in challenging in vivo model



AUTO7 is designed to tackle the complex solid tumor environment



Anti-PSMA humanized CAR T cell for improved persistence and resistance in Prostate Cancer



MODULES DELIVERED USING GAMMA-RETROVIRAL VECTOR



Next Steps

Multiple clinical milestones anticipated through 2021/2022



PRODUCT	INDICATION	TARGET	PHASE	NEXT MILESTONE
Obe-cel	Adult ALL	CD19	Pivotal*	FELIX data in 2022
AUTO1 /22	Pediatric ALL	CD19/CD22	Phase 1	Study enrolling, data in Q4 2021
obe-cel	B-NHL	CD19	Phase 1	Study enrolling, data in Q4 2021
obe-cel	PCNSL	CD19	Phase 1	Study enrolling, data in Q1 2022
AUTO3	DLBCL	CD19/CD22	Phase 1	Intend to partner
AUTO4	TRBC1+ Peripheral TCL	TRBC1+ Peripheral TCL	Phase 1	Phase 1 interim data H1 2022
AUTO5	TRBC2+ Peripheral TCL	TRBC2+ Peripheral TCL	Preclinical	Start Phase 1 H1 2022
AUTO6 NG	Neuroblastoma; Osteosarcoma; SCLC	GD2	Preclinical	Start Phase 1 H1 2022
AUTO7	Prostate	PSMA	Preclinical	Start Phase 1 H1 2022
AUTO8	Multiple Myeloma	BCMA/CAR-X	Preclinical	Start Phase 1 study H2 2021
ALLO Program	Undisclosed	Undisclosed	Preclinical	Start Phase 1 2021

^{*}Subject to confirmation by regulatory authorities.











Autolus poised for potential value inflection



- Obe-cel and AUTO1/22
 - Autolus' first pivotal trial (FELIX) in adult ALL. Enrollment continues and company reiterates guidance to expect data in 2022
 - Pediatric ALL—AUTO1/22 Phase 1 study started in Dec 2020. Update expected Q4 2021.
 - ALLCAR study extension in other relapsed/refractory B-NHL and CLL ongoing. Update expected Q4 2021
 - Opportunity to develop AUTO1 in Primary CNS Lymphoma, CAROUSEL study update expected Q1 2022
- O AUTO4
 - AUTO4 continues in dose escalation in a Phase 1 trial, interim data expected in H1 2022
- Autolus' solid tumor program, AUTO6NG, to enter clinic in H1 2022
- Cash balance at Jun 30, 2021, was approx. \$216 million, anticipate cash runway into H1 2023



Thank you