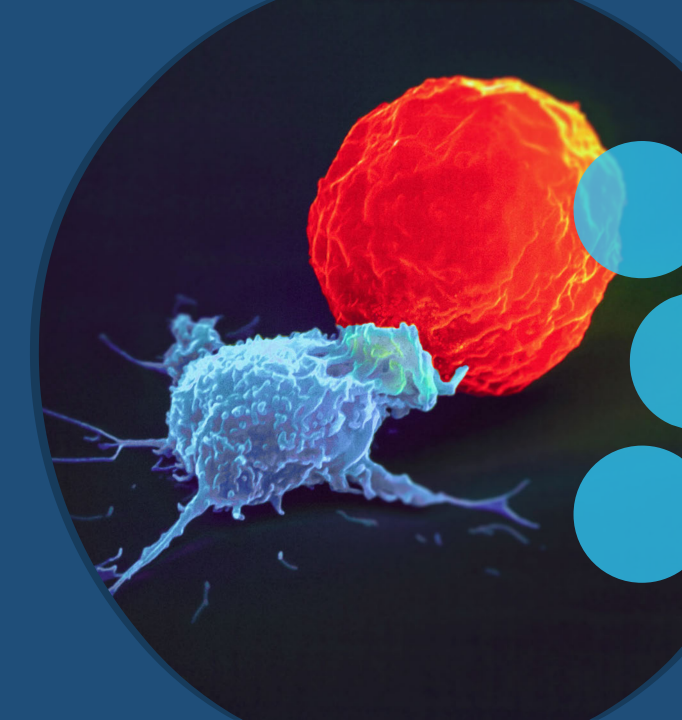


### Allogeneic Natural Killer Cells Engineered to Beat Cancer

October 2020



# Forward looking statements

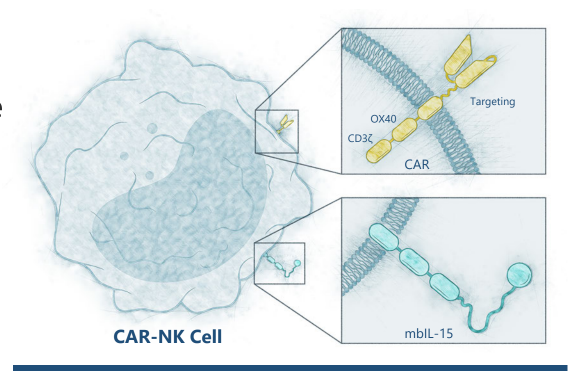
This presentation contains forward-looking statements, within the meaning of the Private Securities Litigation Reform Act of 1995, regarding future events and the future results of the company that are based on current expectations, estimates, forecasts, and projections about the industry in which the company operates and the future of our business, future plans and strategies, projections, anticipated trends and events, the economy, and other future conditions, and the beliefs and assumptions of the management of the company. Words such as "address," "anticipate," "believe," "consider," "continue," "develop," "estimate," "expect," "further," "goal," "intend," "may," "plan," "potential," "project," "seek," "should," "target," "will," variations of such words, and similar expressions are intended to identify such forward-looking statements. Such statements reflect the current views of the company and its management with respect to future events and are subject to inherent risks, uncertainties, and changes in circumstances that are difficult to predict and may be outside our control. Therefore, you should not rely on any of these forward-looking statements. Should one or more of these risks or uncertainties materialize, or should underlying assumptions prove incorrect, the company's actual results, performance, or achievements could differ materially from the results expressed in, or implied by, these forward-looking statements. Please see section entitled "Risk Factors" in our quarterly and periodic filings for a description of these risks and uncertainties. This presentation has been prepared by the company based on information it has obtained from sources it believes to be reliable. Summaries of documents contained in this presentation may not be complete. The company does not represent that the information herein is complete. The information in this presentation is current only as of the date on the cover, and the company's business or financial condition and other information in this presentation may change after that date. The company undertakes no obligation to update any forward-looking statements in order to reflect any event or circumstance occurring after the date of this presentation or currently unknown facts or conditions.



### Highlights: Nkarta engineered CAR-NKs

# Natural Killer cells are the cornerstone of innate immune surveillance

- » Allogeneic and off-the-shelf with attractive cost of manufacturing
- » Proprietary expansion, persistence, tumor targeting and cryopreservation technologies
- » Potential for outpatient administration
- » First IND cleared 3Q 2020; next clinical trial clearance anticipated 1Q 2021
- » IPO gross proceeds of \$290 M, July 2020



Targeting receptor, OX40 costimulatory domain, CD3ζ signaling moiety, membrane bound IL-15



# Nkarta proprietary technologies



### **Expansion**

Co-culture with proprietary K562 stimulatory cell line to achieve high cell doses



### Persistence

Expression of proprietary membrane bound IL-15 to enhance time in circulation



### **Targeting**

Engineered for expression of optimized CARs



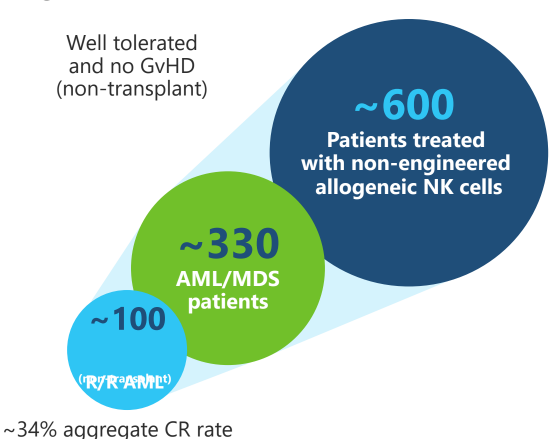
### **Cryopreservation**

Maintains NK cell viability and potency



# Extensive clinical experience validates NK approach

### Patients have been treated with nonengineered NK cells across ~30 studies



Velluchamy 2017; Nkarta systematic literature review. CR: Complete remission.

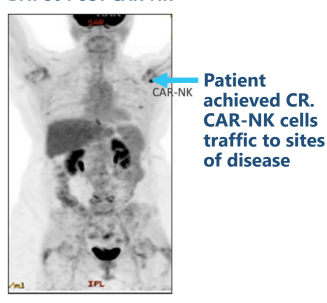
# MD Anderson study with CD19 CAR-NK cells - Published in New England Journal of Medicine, Feb 2020

- √ 7 / 11 CRs in patients with advanced B-cell malignancies
- ✓ No reported CRS, GvHD or neurotoxicity

#### **PRE-TREATMENT**



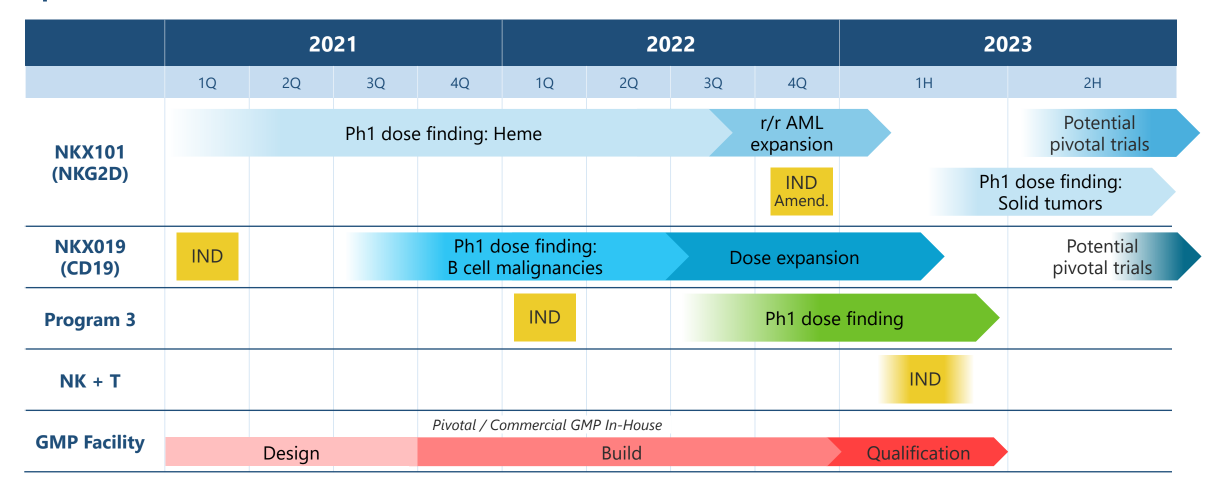
**DAY 30 POST CAR-NK** 



Katayoun Rezvani, M.D., Ph.D., et al., N Engl J Med 2020, 382:545-553. DOI: 10.1056/NEJMoa1910607. Takeda Investor Day 2019. CRS: Cytokine release syndrome. GvHD: Graft versus host disease.



# Pipeline



NKX101 heme IND cleared 3Q20; in-house clinical GMP facility completed mid 2020



# Leadership

MANAGEMENT TEAM		BOARD	
Paul Hastings President & CEO	OncoMed QLT Inc. AXYS CHIRON	<b>Ali Behbahani, MD</b> Chairman	NEA
Ralph Brandenberger, PhD	Baxalta Geron Xcelera	<b>Tiba Aynechi, PhD</b> Director	novo holdings
VP, Technical Operations  Nadir Mahmood, PhD	NEURONA THERAPEUTICS	Fouad Azzam, PhD Director	LSP
Chief Financial and Business Officer	SECOND GENOME THE MICROBIOME COMPANY  Biopharmaceuticals  Goldman Sactis	Mike Dybbs, PhD Director	SAMSARA BIOGAPITAL
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		Zach Scheiner, PhD Director	RACAPITAL
		Laura Shawver, PhD Director	SILVERBACK" THERAPEUTICS





# Platform

# Nkarta CAR-NKs: engineered to enhance activity

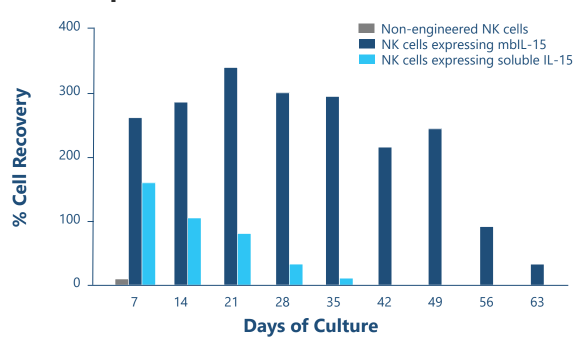
	APPROVED CAR-T THERAPIES	ALLO CAR-T THERAPIES	NK CELLS	CAR-NK CELLS
OPPORTUNITY FOR IMPROVED SOLID TUMOR ACTIVITY				$\checkmark$
PERSISTENCE	$\checkmark\checkmark$	$\checkmark$		$\checkmark$
LOW GVHD RISK	$\checkmark$	TBD	$\checkmark$	$\checkmark$
LOW RISK OF CRS OR NEUROTOXICITY			$\checkmark$	$\checkmark$
ALLOGENEIC, OFF-THE-SHELF MANUFACTURING		$\checkmark$		$\checkmark$
COST OF MANUFACTURING	+++	++	++	+

Nkarta's platform is designed to generate CAR-NKs engineered to address the limitations of current CAR-T therapies, including safety concerns, tumor targeting, manufacturing time and COGS

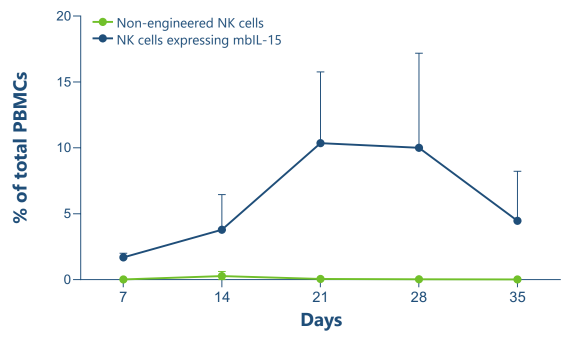


### Superior persistence from membrane bound IL-15

### *In vitro* persistence



### In vivo persistence and expansion in NSG mice



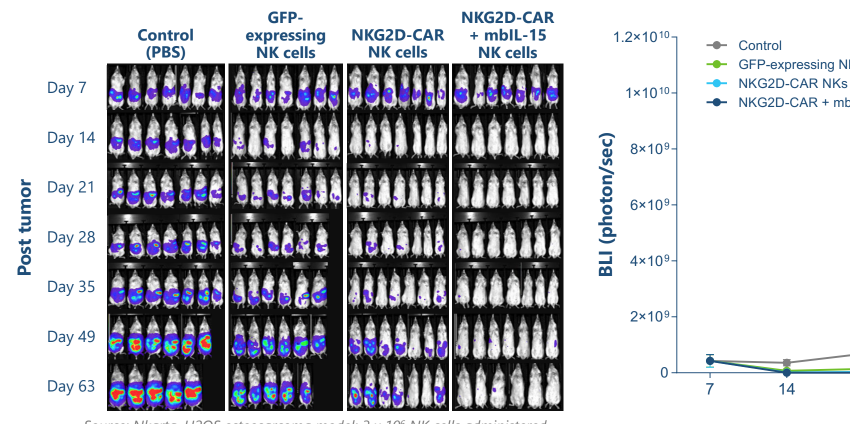
Source: Nkarta. N = 5 per arm.

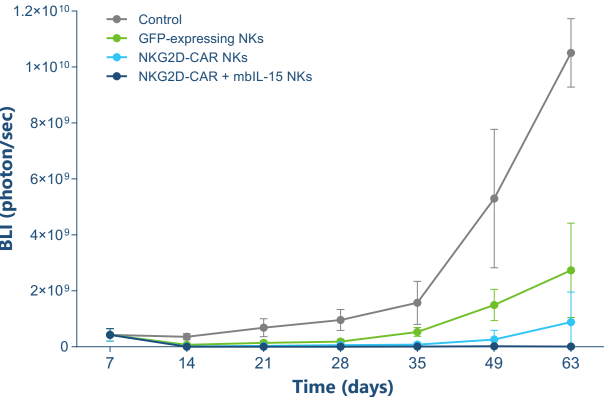
Source: Imamura, Blood 2014

NK cells engineered to express membrane-bound IL-15 (mbIL-15) demonstrate superior persistence as compared to unmodified NK cells



# Persistence and targeting to maximize activity



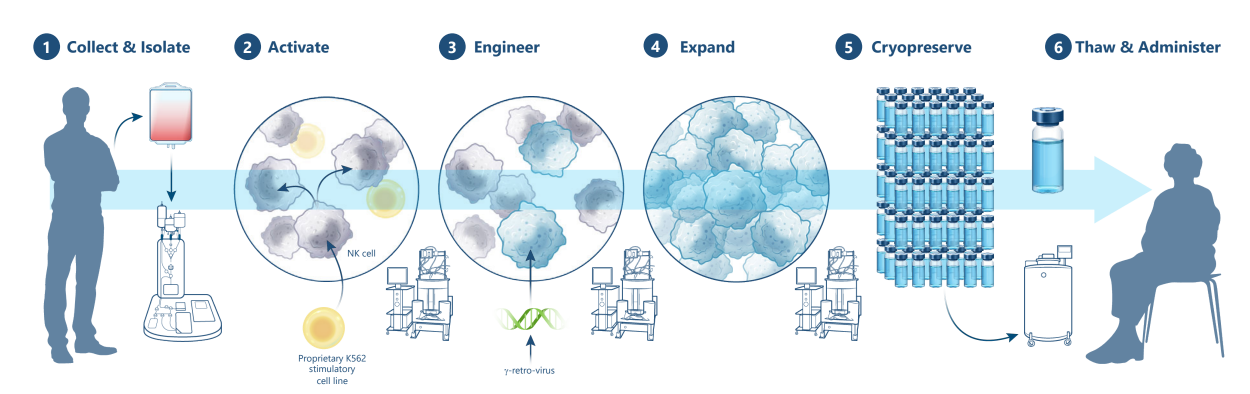


Source: Nkarta. U2OS osteosarcoma model;  $3 \times 10^6$  NK cells administered on D7. Graphical data at right are average BLI of mice above.

NK cells demonstrate enhanced tumor killing when engineered for targeting and mblL-15 expression



# Allogeneic, commercially-enabling manufacturing



NK cells collected from healthy donors by leukapheresis.

NK cell expansion using proprietary stimulatory cells.

Expanded NK cells are transduced to express mblL-15 and CAR.

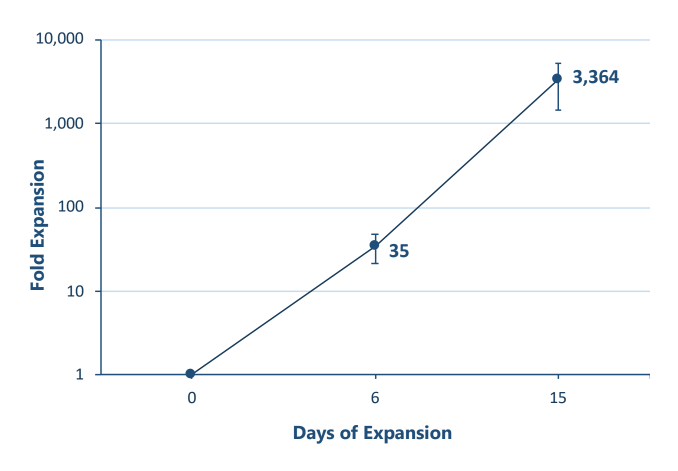
Continued expansion driven by mblL-15.

Harvesting and cryopreservation of final cell product.

NK product candidate is thawed for off-the-shelf administration to patients.



# Proprietary expansion to enable large scale manufacturing

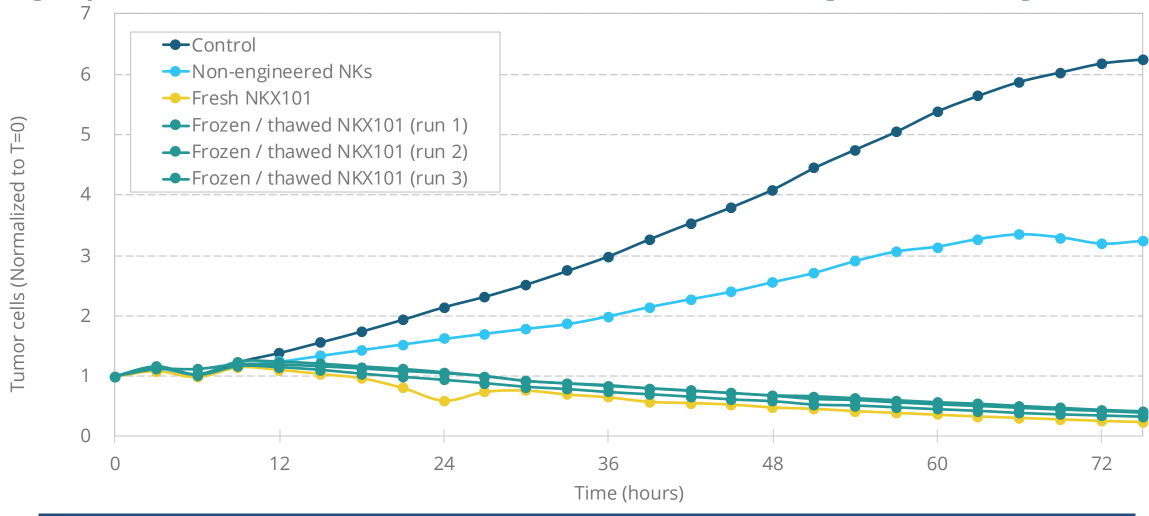


- Extensive optimization enables truly off-the-shelf products
- » In-house cGMP manufacturing suite construction recently completed
- Projected cost of commercial manufacturing at peak:~\$2,000 / dose (500 doses / batch)

Data above are from the process development of NKX019 for cGMP manufacturing and are an average of 5 expansions from 4 different donors.



# Cryopreserved NKX101 retains in vitro cytotoxicity



Cryopreserved NKX101 retains cytotoxicity similar to fresh NKX101 in a long-term assay

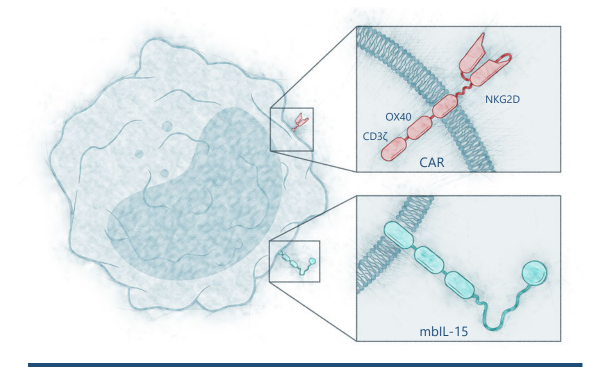




Pipeline

## NKX101: CAR-NK targeting NKG2D ligands

- » NKG2D receptor is primary driver of NK cell activation and tumor killing
- » > 10x increase in NKG2D expression vs. non-engineered NK cells
- » OX40 selected based on superiority vs. other costimulatory domains
- » Targets of NKG2D are selectively over-expressed in cancer cells



NKX101: NKG2D activating receptor, OX40 costimulatory domain, CD3ζ signaling moiety, membrane bound IL-15



# NKG2D: ligands enriched in tumors, demonstrated responses

# NKG2D ligand expression is documented in multiple tumor types

#### **TUMOR TYPE** REFERENCE AML, ALL, CML, CLL Hilpert, J Immunol 2012 **MULTIPLE MYELOMA** Carbone. Blood 2005 HCC Kamimura, J Hep 2012 **BREAST** de Kruif, BMC Can 2012 **OVARIAN** McGilvray, Int J Can 2010 Okita, Can Imm Immunother 2016 LUNG **COLON** McGilvray, CCR 2009 **MELANOMA** Vetter, J Inv Derm 2002 **OSTEOSARCOMA** Lu, Neoplasma 2008 **GLIOMA** Weiss, CCR 2018

# Clinical responses observed in R/R AML with non-engineered allo-NKs validate NKG2D

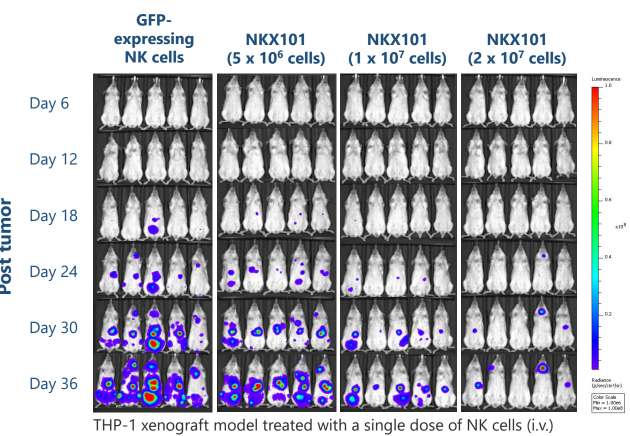
STUDY	RESPONSES*
Bachanova, Crit Rev Oncog 2014, A+B cohort	9 / 42 (21%)
Bachanova, Crit Rev Oncog 2014, C cohort	8 / 15 (53%)
Curti, Blood 2011	1 / 5 (20%)
Kottaridis, PLOS One 2015	1 / 1 (100%)
Miller, Blood 2005	5 / 19 (26%)
Romee, Sci Transl Med 2016	5 / 9 (56%)
Rubnitz, Pediatr Blood Cancer 2015	6 / 12 (50%)
OVERALL	35 / 103 (34%)
*AMI responses in nationts with morphologic disease at ho	iseline as renorted

<sup>\*</sup>AML responses in patients with morphologic disease at baseline as reported in individual trials, patients with CR at study entry excluded from summary. The 35 responses include 20 CR, 12 CRi, 2 CRp and 1 MLFS.



### NKX101: Acute myeloid leukemia (AML)

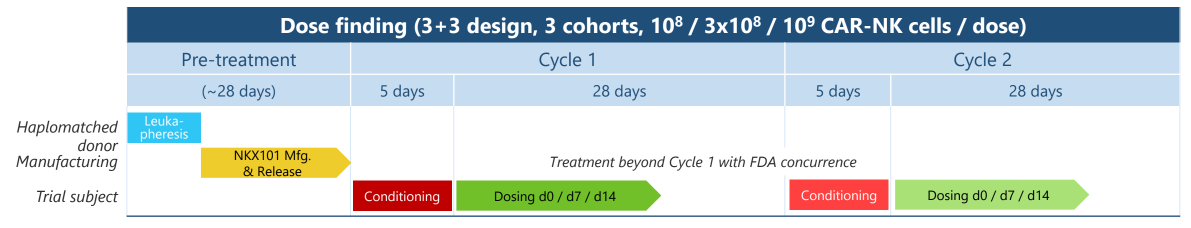
- » AML US incidence: ~21K / yr
  - 5-year survival rate ~28%
- NKG2D targets are over-expressed in AML blasts
- » Clinical activity with non-engineered NKs
- IND cleared July 2020
- Phase 1 in r/r AML and higher-risk MDS: FPI expected 4Q20



2 days after tumor injection



# NKX101: Heme dose finding and expansion



	Dose expansion (6 subjects with unrelated donor)							
		Cycle 1		Cycle 2				
		5 days	28 days	5 days	28 days			
		Treatment beyond Cycle 1 with FDA concurrence						
Trial subject		Conditioning	Dosing d0 / d7 / d14	Conditioning	Dosing d0 / d7 / d14			

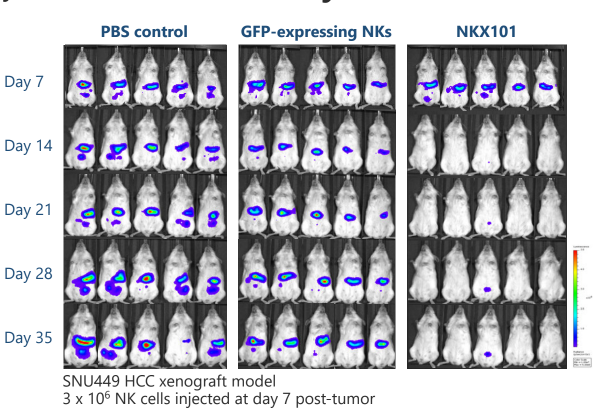
In addition to haplomatched subjects, the dose expansion cohort is designed to evaluate subjects treated with off-the-shelf NKX101 – our expectation for pivotal trials and commercial use



### NKX101: Solid tumors

- » Liver & bile cancer US incidence: ~42K / yr
  - 5-year survival rate ~18%
- » NKG2D targets over-expressed on HCC and CRC cells
- » NK cells are important in liver immunity and tumor surveillance
- » Activity of non-engineered NK cells in HCC/ICC: 3/16 PRs
- » Planned Phase 1: Locoregional delivery using SOC technique in 1° liver cancer or liver metastases

### **NKX101 activity in NSG mice**



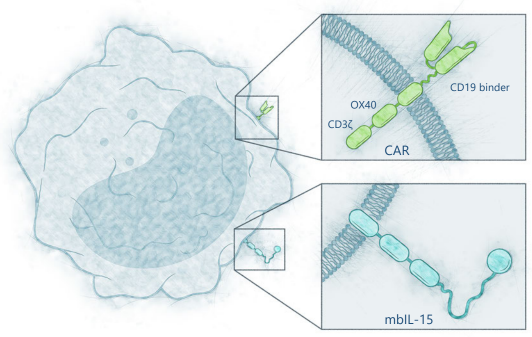
HCC: Hepatocellular carcinoma. CRC: Colorectal cancer. Sources: SEER database; Sun Act Pharm Sin 2015; Kamimura, J Hepatology, 2012; Kamiya et. al, Cancer Immunol Res 2016; Qin 2017

Post tumor



### NKX019: CD19 targeted CAR-NK; IND 1Q21

- » Large opportunity after CAR-T approvals:
  - Gr3+ CRS: 13–49%; Gr3+ neurotoxicity: 18–31%
  - Limited number of specialized sites can treat
  - 9–34% of patients in pivotal trials didn't receive cells (primarily due to mfg. challenges)
- » Rezvani (MDACC / Takeda) CAR19-NK:
  - 7 / 11 CRs in patients with B-cell malignancies (median 4 prior rounds Tx)
  - No reported CRS, GvHD or neurotoxicity
- » Phase 1 in B-cell malignancies
  - Off-the-shelf NKX019
  - 3x10<sup>8</sup> starting dose, 2 dose finding cohorts
  - Several dose expansion cohorts thereafter

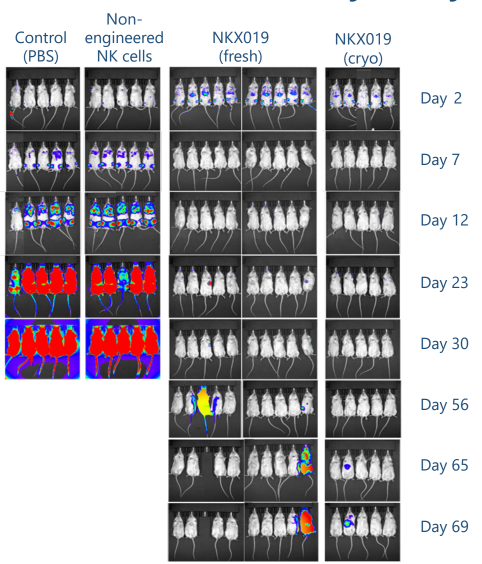


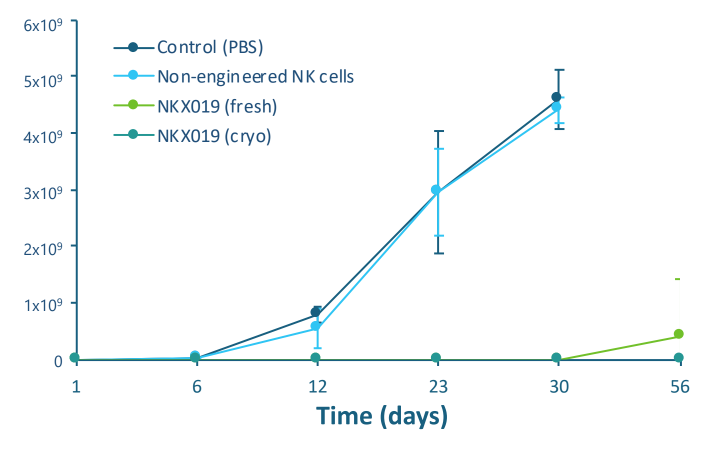
NKX019: Proprietary CD19 binder, OX40 costimulatory domain, CD3ζ signaling moiety, membrane bound IL-15

Sources: Kymriah and Yescarta package inserts; Rezvani NEJM 2020. Per NEJM publication, CR/SD patient achieved a CR for Richter's transformation and SD for underlying CLL.



### NKX019: Activity in lymphoma model





Nalm-6 lymphoma model. 10<sup>7</sup> cells administered one day post tumor. Graphical data above are an average of mouse luminescence at left. "Cryo" denotes cryopreserved then thawed NKX019.

NKX019 production under optimized conditions allows cryopreservation with retention of *in vivo* activity





# Corporate

### Intellectual property

### **PLATFORM**

### **NK cell expansion**

- » Multiple issued patents and pending applications
- » Compositions and methods of expansion/treatment
- » Expiry ~2024 to ~2038

### **NK cell persistence**

- » Allowed US application and multiple pending OUS applications
- » Expiry ~2035

### **Pipeline**

- » Provisional applications
- » Compositions & treatment methods
- » Expiry ~2039 to ~2040

### **NKX101**

### NKG2D target

- » Issued US patents and multiple pending US/OUS/PCT applications
- » Claims to various NKG2D targeting constructs & treatment methods
- » Expiry ~2034 to ~2039

### **Local NKX101 delivery**

- » Provisional applications
- » Local delivery to tumors
- » Expiry ~2039

### **Combo Therapy**

- » Provisional applications
- » NKG2D construct + adjunct therapy
- » Expiry ~2039

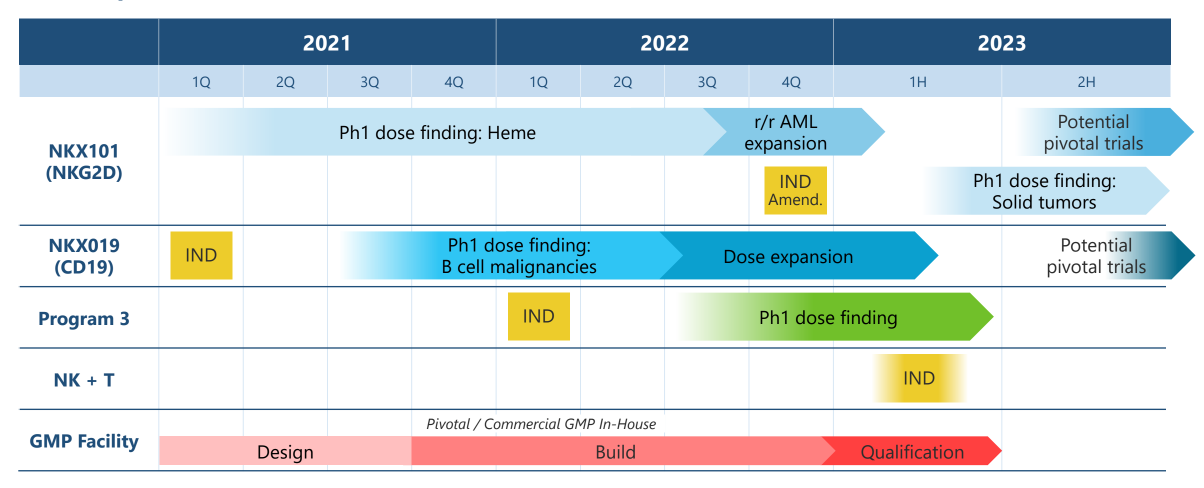
### **NKX019**

#### **CD19**

- » Provisional applications
- » Cells expressing tumor-targeting receptor & cytotoxic effector
- » Expiry ~2040



# **Anticipated Milestones**



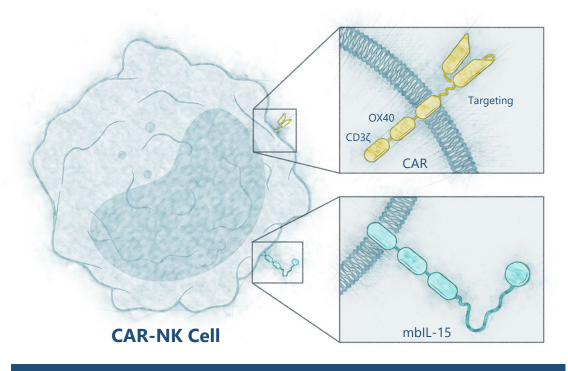
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