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# Corporate Presentation

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# RIGHT TARGETS. RIGHT CELLS. RIGHT PLACE.

Our goal is to orchestrate the immune system to target oncogenic driver mutations and deliver transformative therapies for patients with solid tumors



# Experienced Management Team Supported by Blue-Chip Investor Syndicate

Executive Leadership

Board of Directors

Investors





# **Co-Founders**

# Scientific Advisors



Phil Greenberg, MD Scientific Co-Founder

Fred Hutch Cancer Center



Aude Chapuis, MD Scientific Co-Founder

W 🚀 Fred Hutch



Tom Schmitt, PhD Scientific Co-Founder

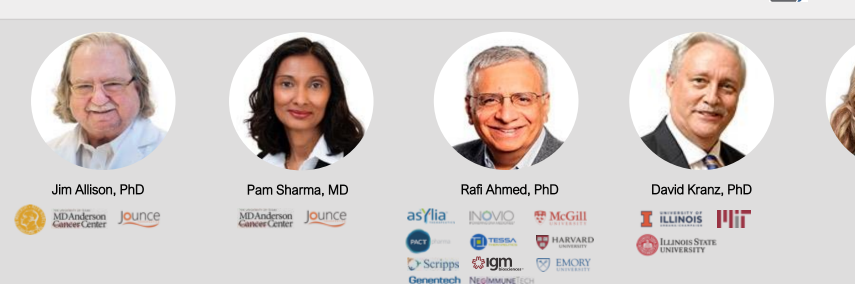
W Kred Hutch



Chris Klebanoff, MD Scientific Co-Founder













Cancer cells are dependent on oncogenic drivers KRAS mutations are present in 30% of all solid tumors Targeting KRAS has been clinically de-risked by approved G12C therapies Affini-T TCRs have high specificity for KRAS and other oncogenic drivers



Oncogenic driver mutations initiate and maintain cancer growth, and are present in each tumor cell

Minimizes tumor heterogeneity and escape mechanisms

KRAS represents the most frequently mutated oncogene in difficult-to-treat solid tumors

Provides impact for a high unmet medical need

Recent drug approvals demonstrate single agent activity but need improved duration of response

Robust Pharma interest for drugs targeting KRAS



Affini-T leverages TCRs to attack only cancer cells, utilizing synthetic biology to enhance persistence

Therapeutic modality with clinical PoC

Tran et al. NEJM (2016), Leidner et al. NEJM (2022), From Joglekar AV and Li G, Nat Methods (2021), From Junttila MR and Sauvage FJ, Nature (2013), From Prior et al. Cancer Res (2020)



### **RIGHT TARGETS**

Targeting oncogenic drivers optimizes solid tumor target specificity and leverages cancer dependency

## **RIGHT CELLS**

Building a path to persistence using T cells enriched for stemness and coordinating a CD4/CD8 T cell response

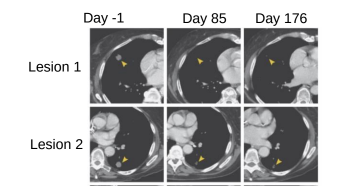
## **RIGHT PLACE**

Synthetic biology empowers T cells to convert signals in the hostile TME into T cell survival and proliferation drives

The NEW ENGLAND JOURNAL of MEDICINE

#### Neoantigen T-Cell Receptor Gene Therapy in Pancreatic Cancer

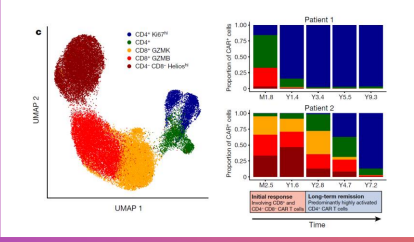
– Leidner and Tran et al.



#### nature

# Decade-long leukaemia remissions with persistence of CD4 $^{\scriptscriptstyle +}$ CAR T cells

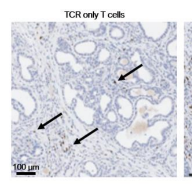
- Melenhorst and June et al.



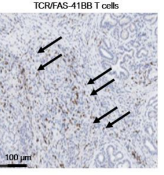


# A Fas-4-1BB fusion protein converts a death to a pro-survival signal and enhances T cell therapy

- Oda and Greenberg et al.



No / limited T cell invasion in tumor bed (blue)



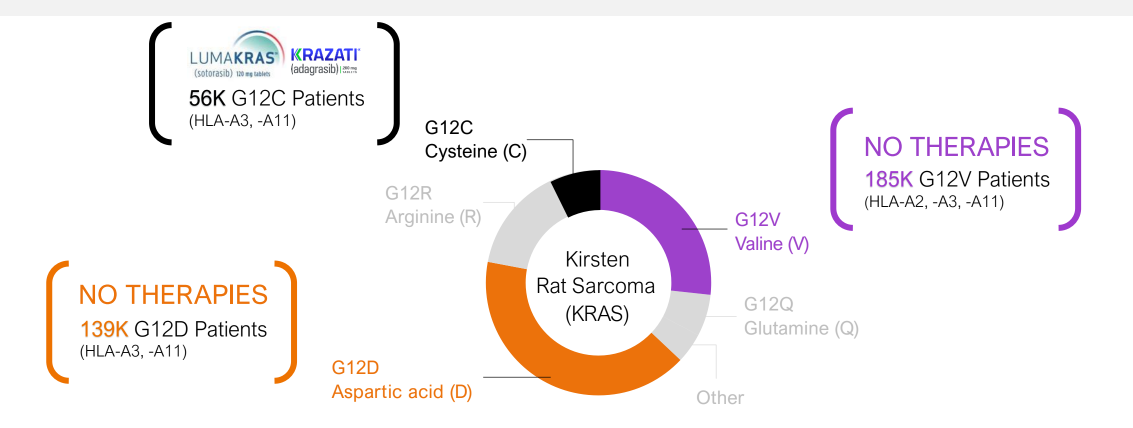
Tumor bed (blue) are invaded by T cells (brown

TME = Tumor Microenvironment



# Targeting KRAS: Large Unaddressed Patient Populations with G12 Mutations

Addressable KRAS G12 mutations across multiple solid tumor populations: ~ 380K annually



Affini-T estimates based on 2020 American Cancer Society (USA) and WHO IARC (EU, UK, CN, and JP) incidence (newly diagnosed patients/yr), frequencies of mutations from Hofmann et al. Cancer Discovery (2022), and HLA frequencies from National Marrow Donor Program (USA, census adjusted) and Immune Epitope Database (Europe, CN, JP). Indications include lung, CRC, PDAC, and endometrial. Total number of lung cancer patients was adjusted by 40% for lung adenocarcinoma. "Market Report: Lumakras Drug Clinical Insight & Sales Forecast 2026. ID 5368077.



# Extensive Portfolio of Proprietary Technologies

	TAILOR <sup>TM</sup> TCR Discovery	TUNE <sup>™</sup> Synthetic Biology	THRIVE <sup>TM</sup> Engineering and Manufacturing
<b>Tools</b> Build a state-of-the- art platform	<ul> <li>Predictive algorithms and machine learning</li> <li>High throughput screening</li> </ul>	<ul> <li>Switch receptors</li> <li>Co-receptors</li> <li>Others (e.g., armoring, persistence)</li> </ul>	<ul> <li>Single LVV transduction of CD4/CD8 T Cells</li> <li>Gene editing with Type II and Type V systems</li> <li>Cryopreserved cell product</li> </ul>
Rationale Equip cells to sustain anti- tumor response	<ul> <li>Leverage tumor dependency</li> <li>Expand patient access globally</li> </ul>	<ul> <li>Improve T cell persistence in TME to enhance therapeutic durability</li> <li>Reduce/prevent T cell exhaustion</li> </ul>	<ul> <li>Gene knock-out and non-viral knock-in</li> <li>High yield TCR+ T cells per run of naïve and memory T cells</li> </ul>
Approach Streamline processes and materials	Avidity	Transgenic TCR Co-Receptor Switch Receptor	CD8 T Cell CD8 T Cell CD8 T Cell CD4 T Cell CD4 T Cell

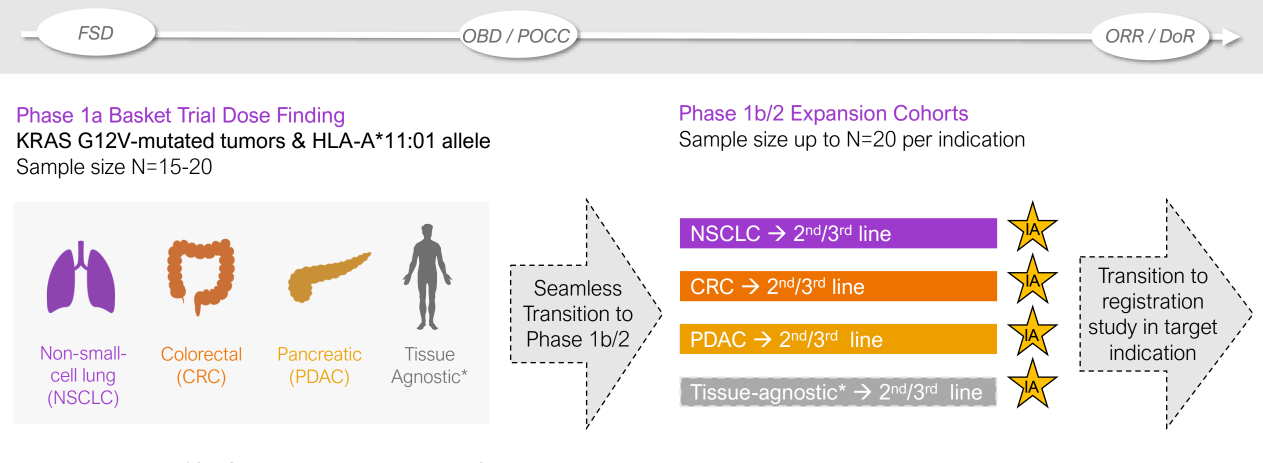


# First-In-Class Potential for Multiple Products Targeting Oncogenic Drivers in Solid Tumors

Program	Target	Sup Rio	Discovery	Preclinical	Clinical			
Program	Target	Syn Bio	Discovery	Flechinical	Clinical			
Oncogenic Driver Programs								
AFNT-111		CD8 Co-Receptor	HLA-A11		*IND Submission 2H '23			
AFNT-211	KRAS G12V	CD8 CoR + FAS-41BB	HLA-A11		*IND Submission 2H '23			
		TUNE™	HLA-A2	1 1 1				
		TUNE™	HLA-A3					
AFNT-212	KRAS G12D	CD8 CoR + TUNE™	HLA-A11		*IND Submission 2024			
		TUNE™	HLA-A3					
	KRAS G12C	TUNE™	Multiple					
	P53 R175H	TUNE™	HLA-A2					
	PIK3CA	TUNE™	Multiple					
	Undisclosed	TUNE™	Multiple	-       				
Viral Antigen Driver Programs								
	Undisclosed	TUNE™	Multiple					
*Planned submission dates								
CD8 CoR = CD8 Co-Receptor								



# AFNT-211: Early-Phase Clinical Development Plan



• Approximately 10 US clinical trial sites planned for Phase 1a dose finding

- Expand clinical trial sites to 35-40 in US, EU5, and CAN
- Accelerated approval based on ORR & DoR data
- Total sample size N=~80 per indication

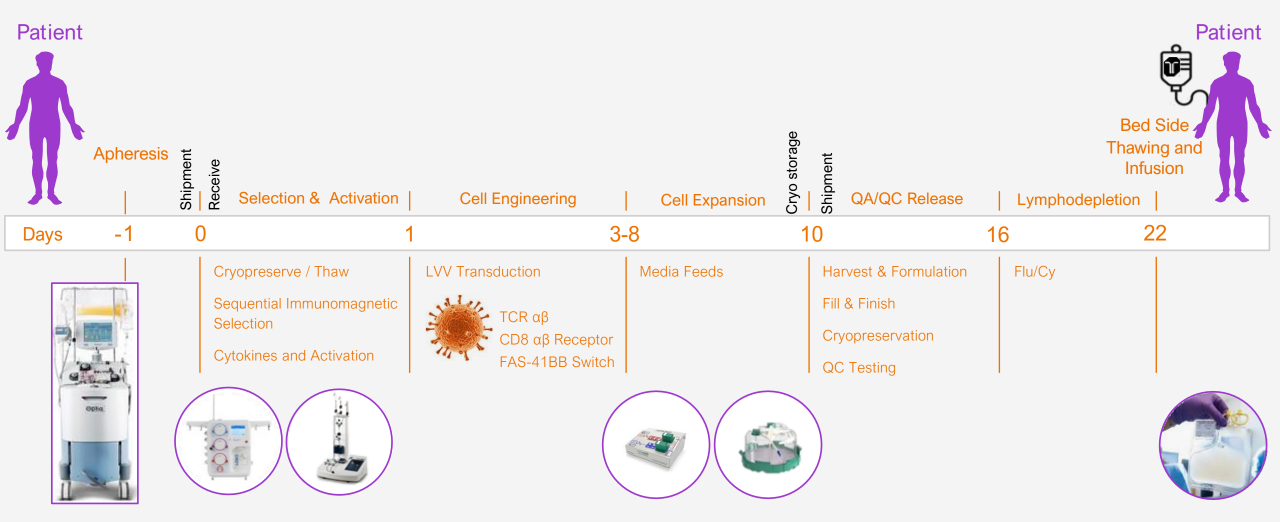
#### \*Excluding primary brain tumors

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ORR – Overall Response Rate DoR – Duration of Response IA – Interim Analysis

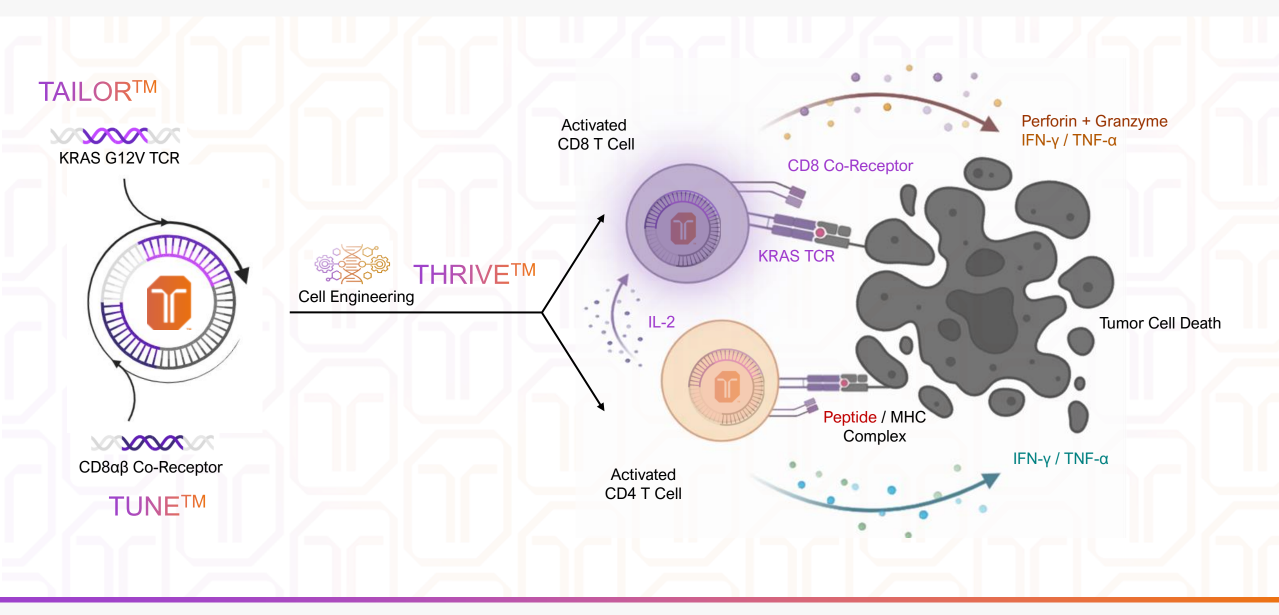


### 10-Day Autologous Manufacturing Process



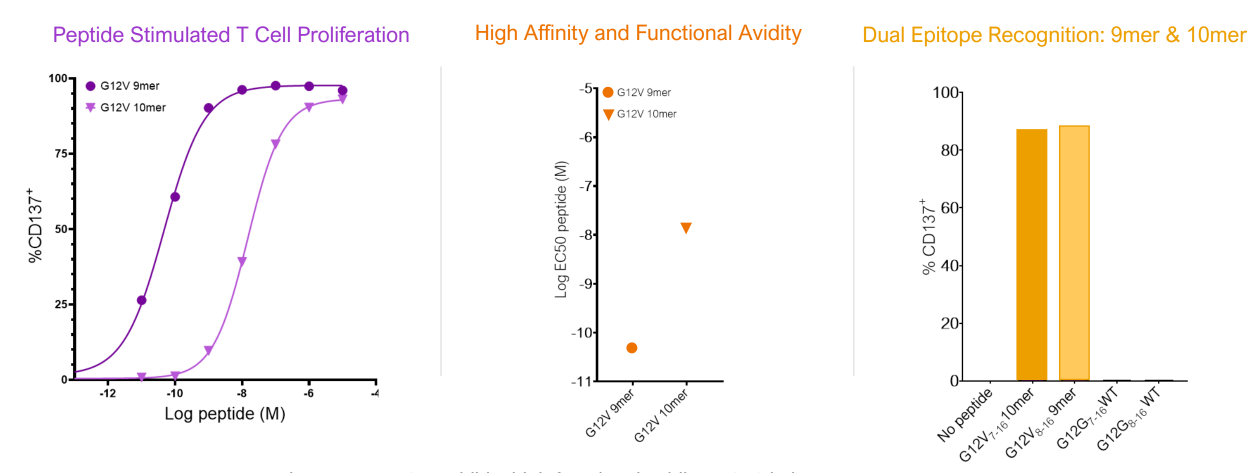


# AFNT-111: KRAS A11 G12V TCR Engineered T Cells + CD8 Co-Receptor





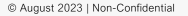
# Robust Preclinical Avidity of KRAS A11 G12V TCR Candidate



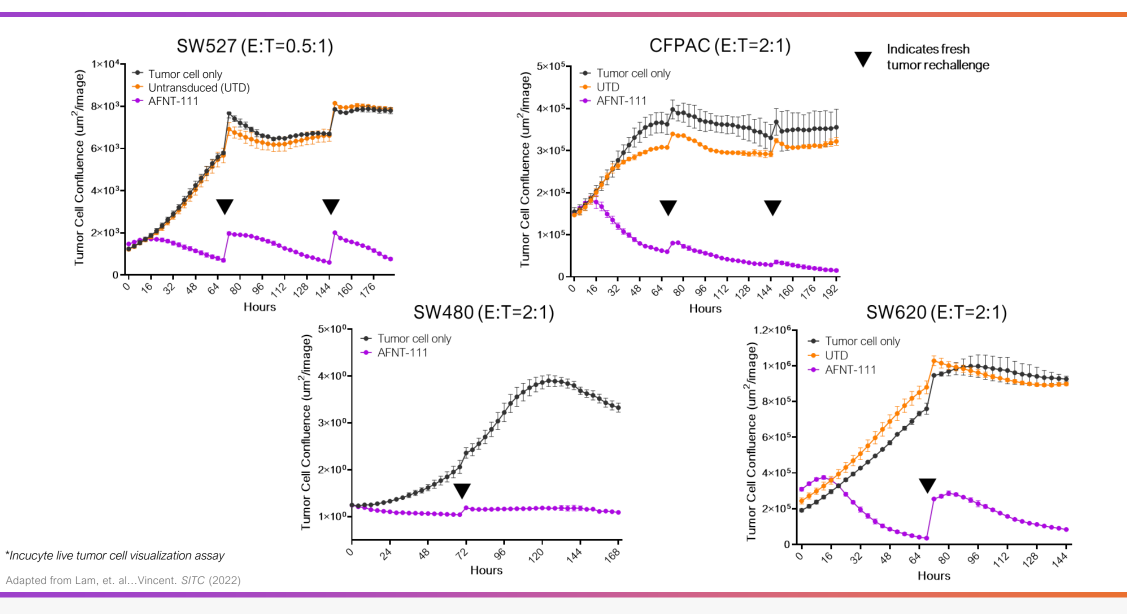
- Lead AFNT-111 TCR exhibits high functional avidity: EC50 in low pM
- Recognition of both 9mer and 10mer KRAS G12V epitopes with no reactivity to WT KRAS

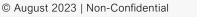
\*Peptide dose response CD137 T cell activation assay

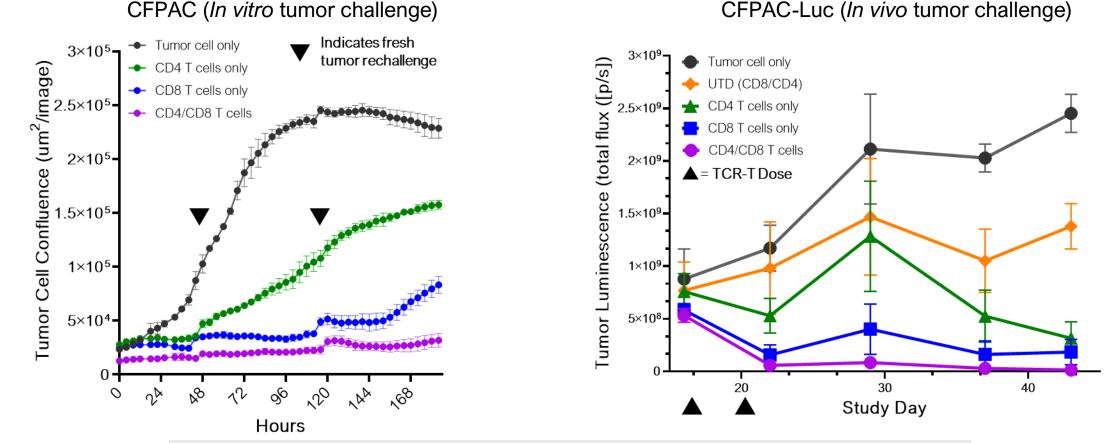
Adapted from Lam, et. al...Vincent. SITC (2022)



Sustained Cancer Cell Killing with AFNT-111 TCR for In Vitro Repeat Tumor Challenge Assay







CFPAC (In vitro tumor challenge)

CD8<sup>+</sup> T cell killing is enhanced by the presence of CD4<sup>+</sup> T cells both *in vitro* and *in vivo* 

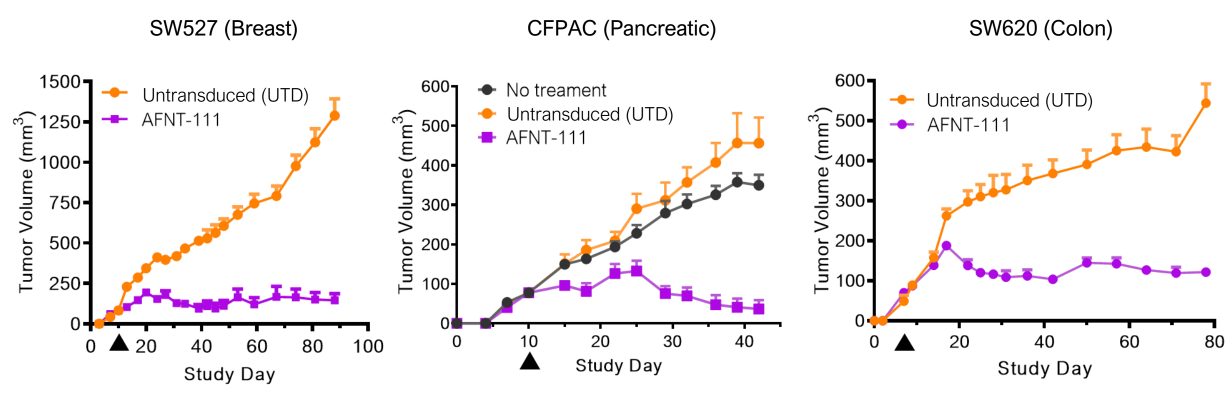
\*Incucyte live tumor cell visualization assay; Total T cells kept constant in CD4 only, CD8 only and CD4/CD8 groups

NSG mice randomized after IP tumor implantation (5 mouse/group) Dose: two administrations of 7x10<sup>6</sup> (D8) and 8x10<sup>6</sup> (D21) AFNT-111 TCR-T cells

Adapted from Lam, et. al...Vincent. SITC (2022)



# AFNT-111: T Cells Targeting KRAS G12V Reduce Tumor Volume Across Several In Vivo Tumor Models



#### $\blacktriangle$ = TCR-T Dose

- NSG mice randomized after SC tumor implantation (5 mouse/group)
- Dose: single IV administration of 3x10<sup>6</sup> AFNT-111 TCR-T cells on D7

- NSG mice randomized after SC tumor implantation (5 mouse/group)
- Dose: single IV administration of 1x10<sup>7</sup> AFNT-111 TCR-T cells on D10

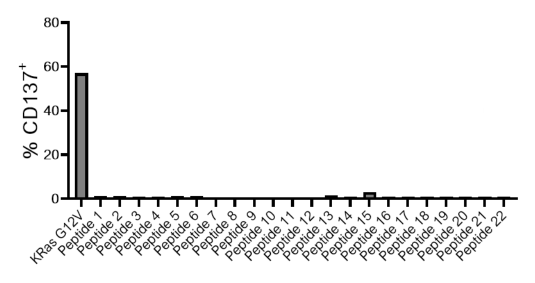
- NSG mice randomized after SC tumor implantation (5 mouse/group)
- Dose: single IV administration of 3x10<sup>6</sup>
   AFNT-111 TCR-T cells on D9

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Adapted from Lam, et. al...Vincent. SITC (2022)

## **Cross-Reactivity**

XScan cross-reactivity assay used to define the peptide recognition motif of AFNT-111 yielded a tolerable preclinical safety profile

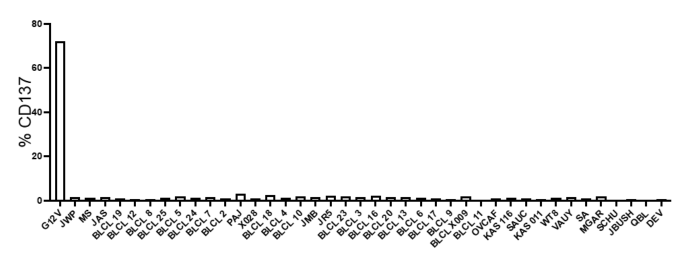


• No cross-reactive self-peptides of concern identified out of 22 identified via XScan and *in silico* analysis

Adapted from Lam, et. al...Vincent. SITC (2022)

### Allo-Reactivity

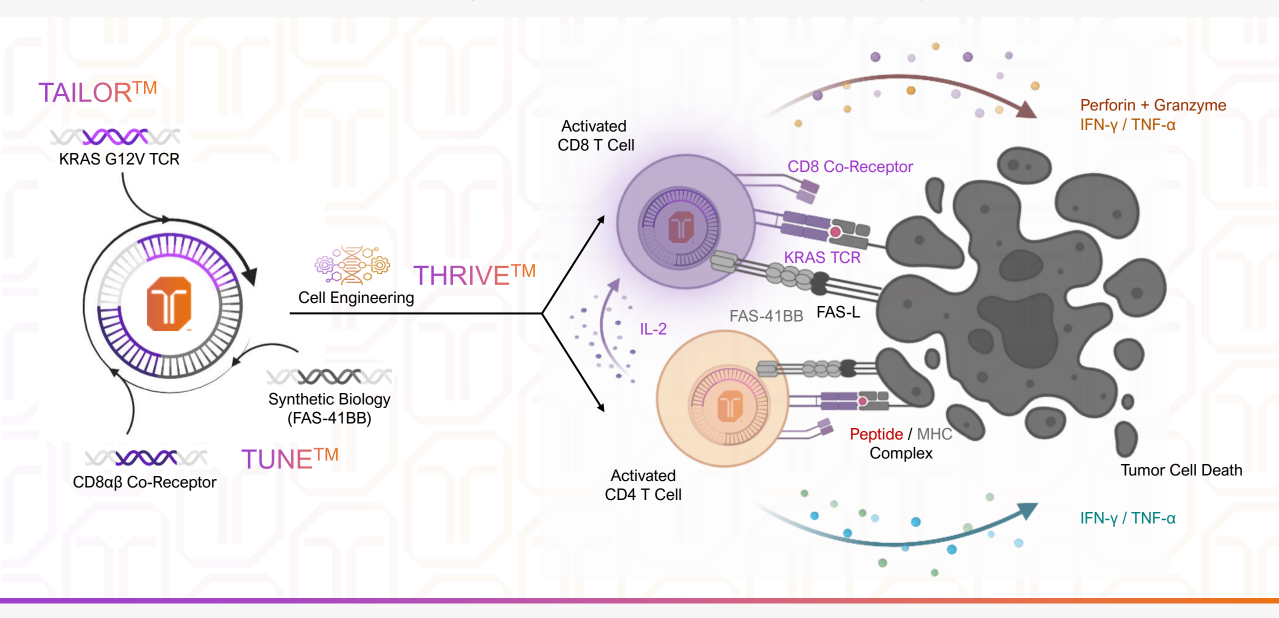
Allo-reactivity screen conducted with 40 B-Lymphoblastoid Cell Lines (B-LCL) had no responses detected



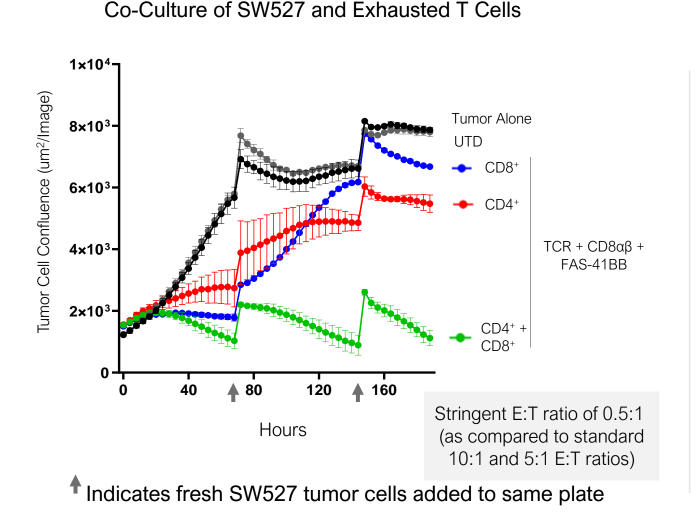
- No allo-reactive responses detected
- B-LCL library covers >95% of the most common HLA alleles



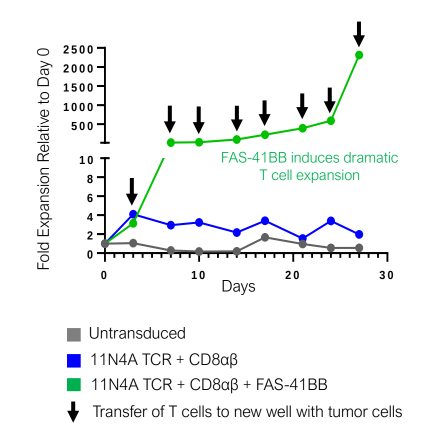
AFNT-211: KRAS A11 G12V TCR Engineered T Cells + FAS-41BB Durability Switch Receptor



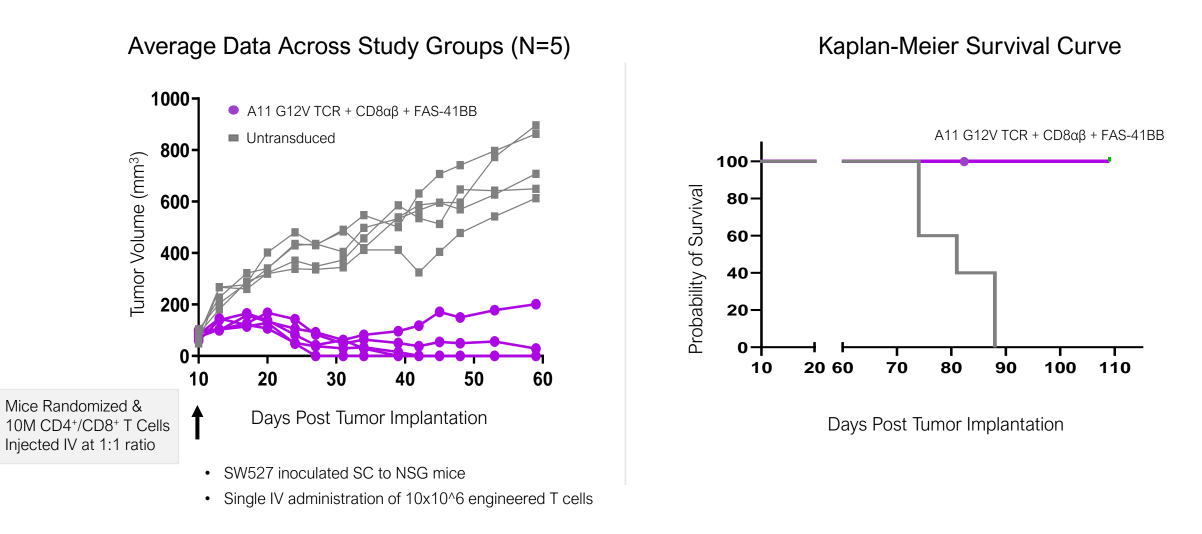




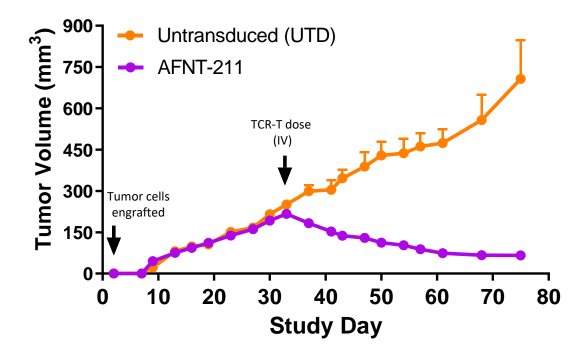
#### FAS-41BB Increases Cell Proliferation in SW527 Model



Adapted from He, et. al...Vincent. SITC (2022)



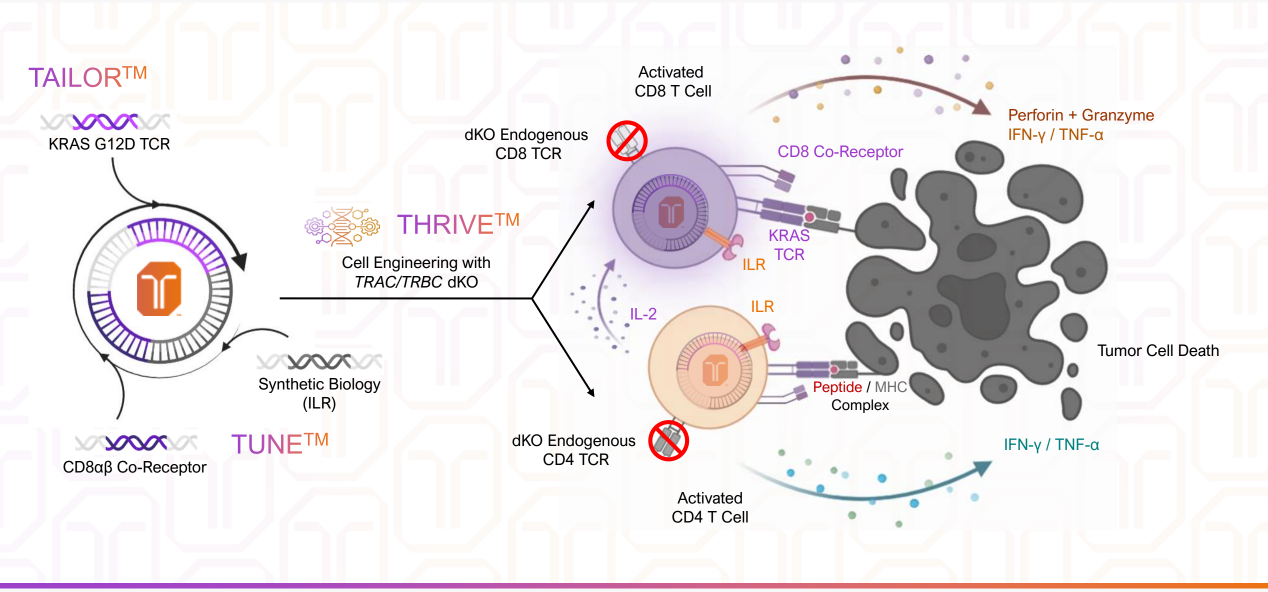
Adapted from He, et. al...Vincent. SITC (2022)



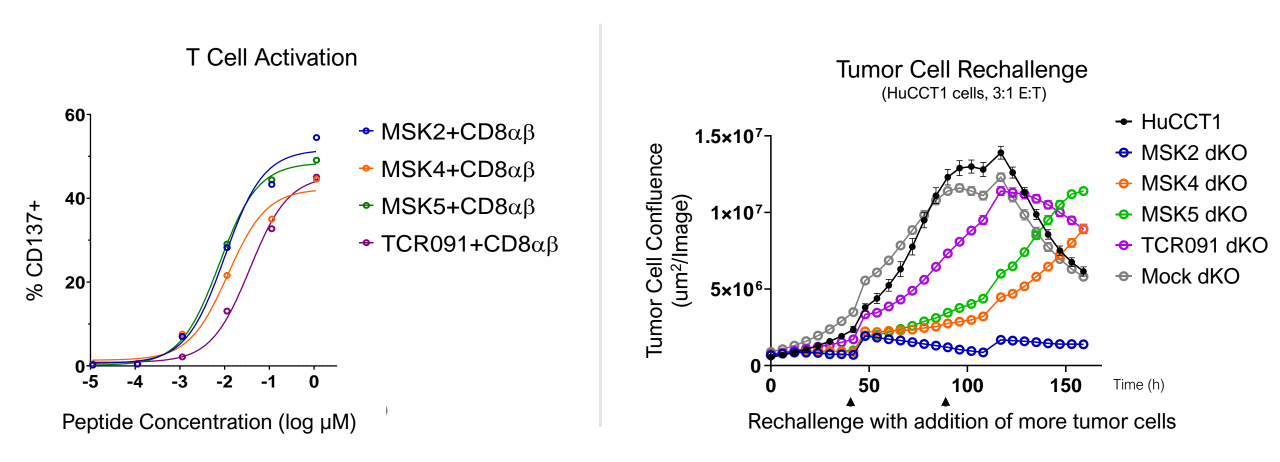
- Large tumors ~300mm^3
- 5x10^6 AFNT-211 cells administered IV on D33 post-tumor SW527 engraftment

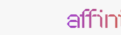


AFNT-212: KRAS A11 G12D TCR Engineered T Cells + Durability Switch Receptor + Gene Editing

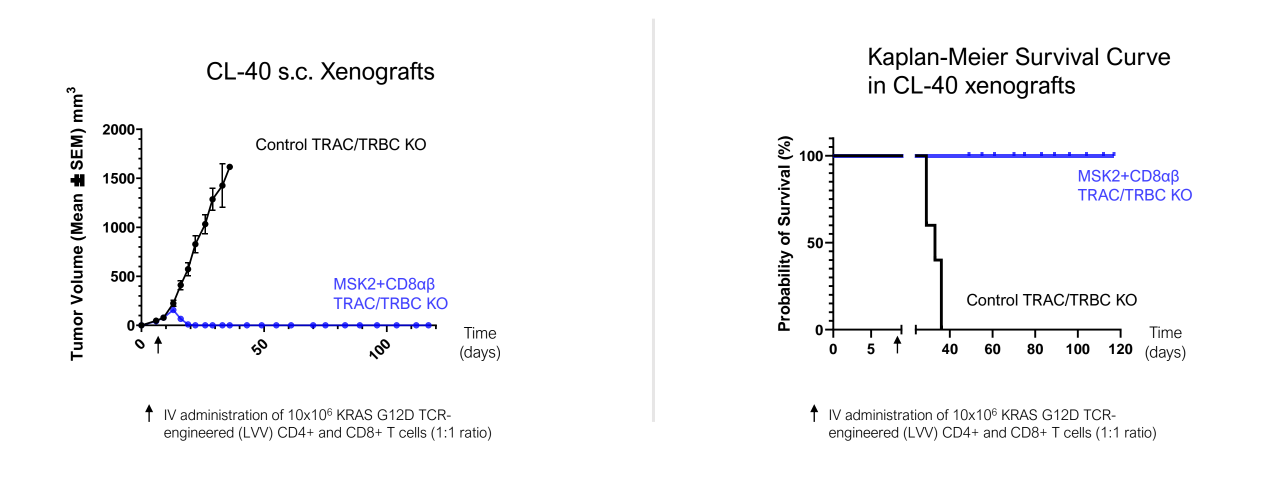






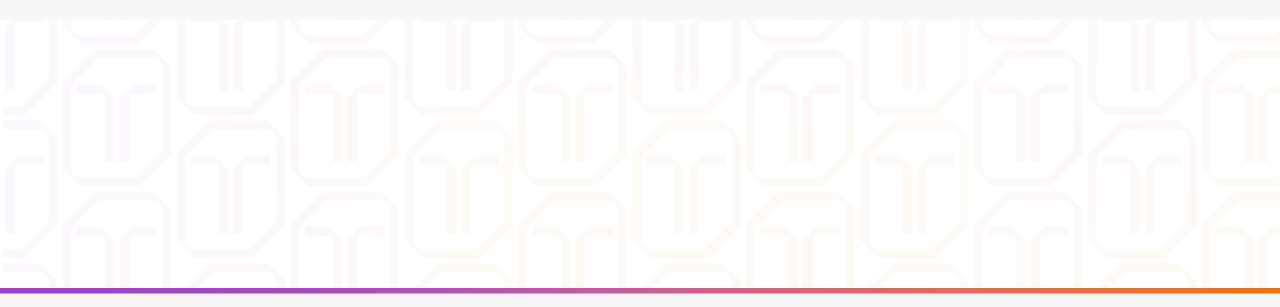


KRAS G12D TCR-T Cells with Endogenous *TRAC/TRBC* dKO Show Robust Preclinical Activity in CL-40 Established Tumors *In Vivo* 



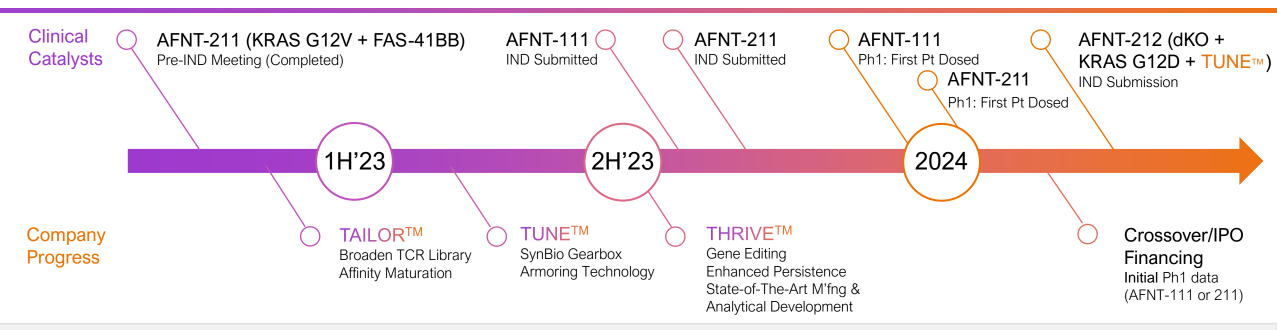
Adapted from Gupta, et. al...Vincent. AACR (2023)

# The Affini-T Opportunity





# Affini-T: Current Status and Key Clinical Catalysts



Affini-T is the premier Precision Immunotherapy company targeting oncogenic driver mutations to develop curative therapies for patients with solid tumors

#### Funding, Team and Culture

- \$194M raised to date
- Skilled team of >90
- Headquartered in Watertown, MA

#### Strategic Partnerships

- <u>The Fred Hutch</u> and <u>Memorial Sloan Kettering</u> TCR discovery and synthetic biology
- <u>ElevateBio</u> phase-appropriate M'fng capabilities with path to commercialization
- <u>Metagenomi</u> world class gene editing capabilities to power next-gen products
- <u>Adimab</u> affinity maturation and T cell engager constructs

#### Programs Tracked against Key Clinical Catalysts

- AFNT-211 and AFNT-111: lead assets targeting KRAS G12V completed positive pre-INDs and tracking for 2H23 INDs
- AFNT-212: first gene edited asset targeting KRAS G12D transitioning to IND enabling studies
- Robust discovery pipeline resourced to target additional oncogenic drivers including P53 and PIK3CA
- Establishing non-viral knock-in gene transfer product platform





