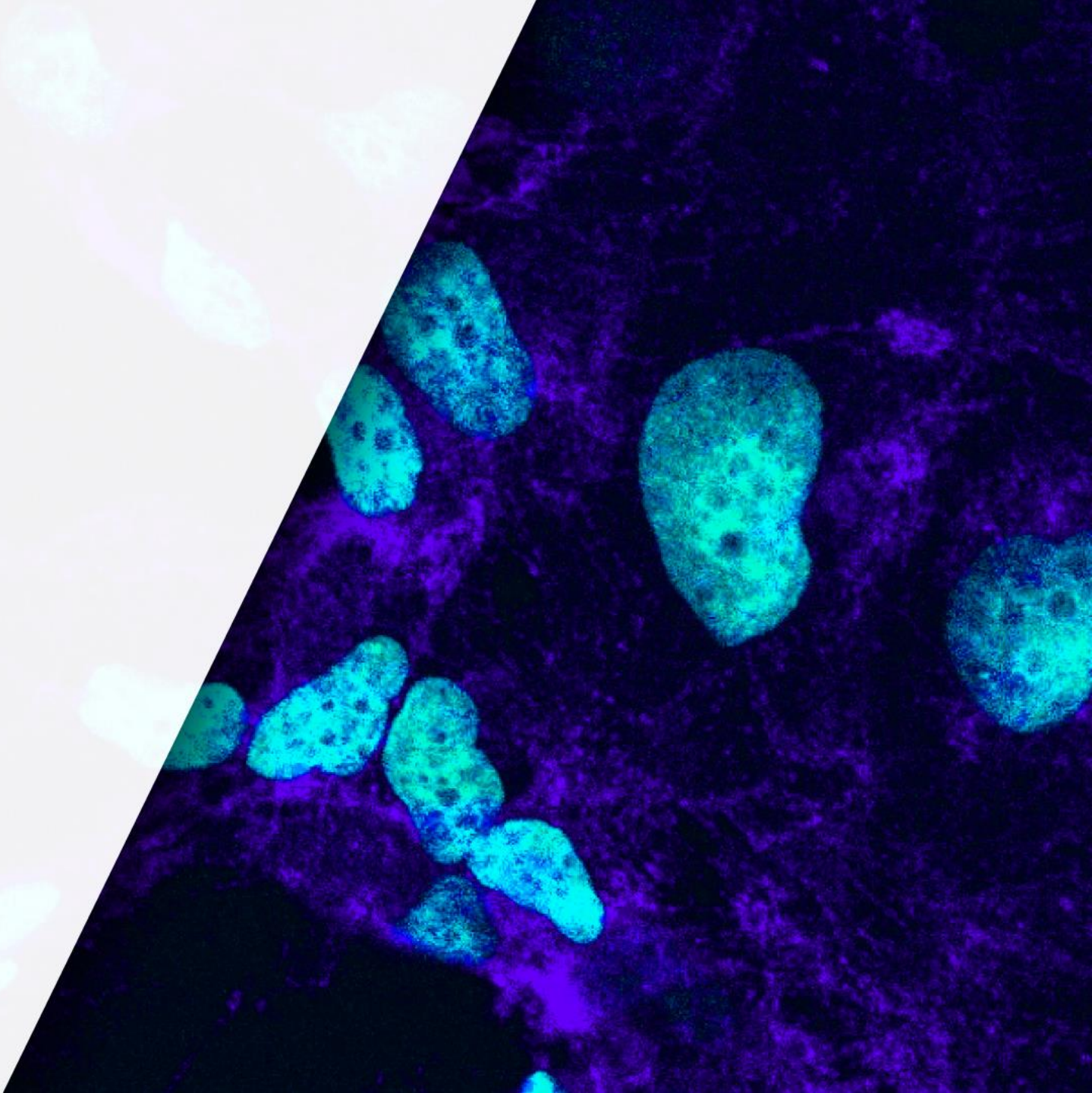


# UNMASKING CANCER CELL CAMOUFLAGE

COMPANY PRESENTATION | June 2022



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# Company Highlights



## Differentiated Pipeline

- Novel, next generation potential across **three programs in solid and hematological malignancies**
- Lead candidate DSP107 designed to target **CD47/SIRP $\alpha$  and 41BB specifically and conditionally** in the tumor microenvironment
- Versatile MIRP platform **targeting innate and adaptive immunity**



## Multiple Near-Term Catalysts

- **Anticipated data readouts** for DSP107 in Q2 2023 in NSCLC (phase II) and in Q4 2022 in heme malignancies (phase I)
  - Dose escalation portion of Phase I/II data **demonstrates favorable preliminary safety profile and monotherapy activity** in solid tumors
- Expected IND filing for DSP502 and DSP216 in 2024



## Cash Runway

- Raised **~\$100 million** to date
- Investors include **aMoon, BVF, DAFNA, Cancer Focus Fund**



## Experienced Leadership

- **Experienced leadership** and executive team with track record of success
- Supported by leading scientific advisory board

# Experienced Leadership Team



**Aron Knickerbocker, MBA**  
Board Chairman



**Yaron Pereg, PhD**  
Chief Executive Officer



**Tomer Cohen, MBA**  
Chief Financial Officer



**Adam Foley-Comer, MD**  
Chief Medical Officer



**Ayelet Chajut, PhD**  
Chief Technology Officer



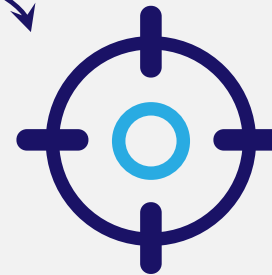


Effectively treating  
cancer requires a  
multifaceted approach



Selectively disabling  
cancer defense mechanism

WHILE



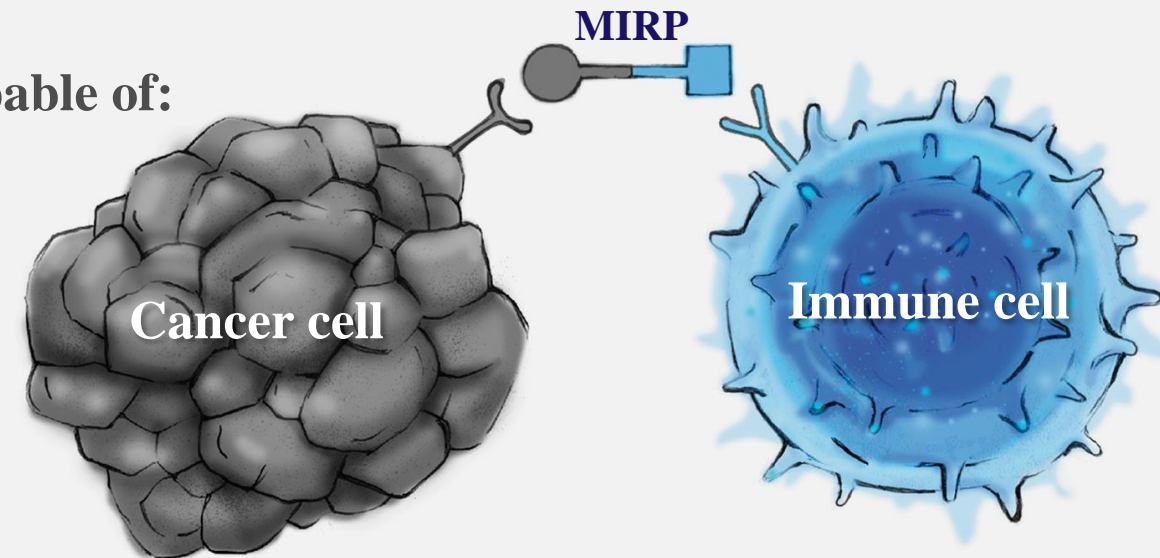
Recruiting a local  
targeted immune attack

# Versatile Multifunctional Immunotherapeutic Platform for Solid and Hematological Malignancies

## MULTIFUNCTIONAL IMMUNE RECRUITMENT PROTEIN (MIRP)

Enabled us to design dual-targeting fusion proteins capable of:

- 1 Inhibiting key evasion markers on cancer cells
- 2 Activating innate and adaptive anti-tumor immunity



# MIRP Platform - Customized, Modular and Flexible Design

## ECD Type I protein

Ig superfamily

- ✓ Receptors
- ✓ Ligands
- ✓ Tumor antigens

(e.g. PD-L1, PVR, CD47, HLA-G)

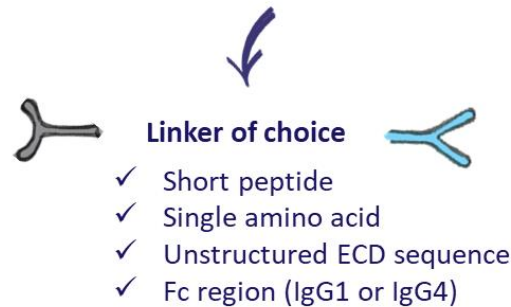


## ECD Type I or II protein

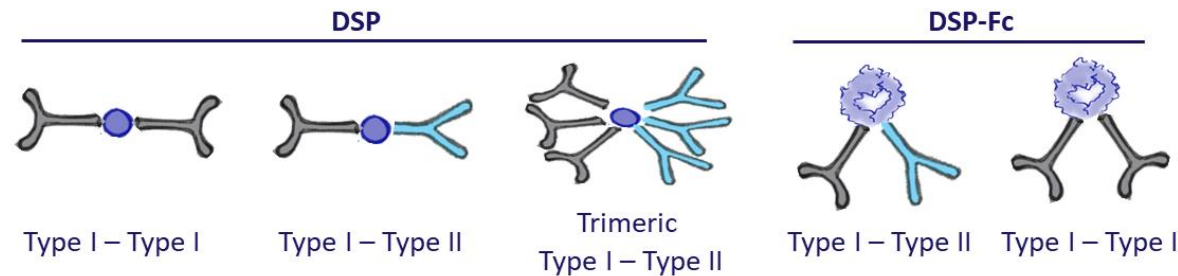
Ig superfamily or TNF superfamily

- ✓ Receptors
- ✓ Ligands
- ✓ Immune checkpoint

(e.g. 4-1BBL, TIGIT, PD1, LILRB)



## Customized composition



# Enhanced, Tumor-Localized Immune Cell Recruitment & Activation With Potential to Improve Safety and Efficacy

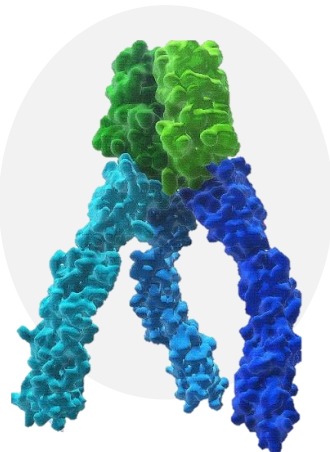
## DSP (Dual Signaling Protein)

Combined checkpoint inhibition and  
immune co-stimulation

### DSP107

**4-1BB activator** –  
*CD47-conditional  
T-cell activation*

**CD47 inhibitor** –  
*Trimeric binding  
for cancer specific  
blocking*



## DSP-Fc (Dual Signaling Protein With Fc Domain)

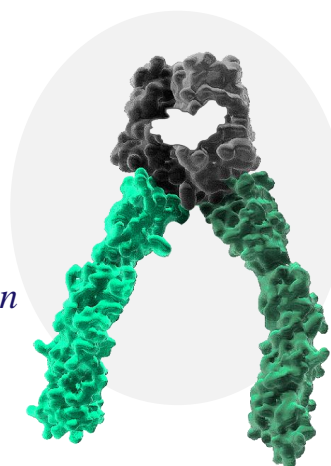
Dual checkpoint inhibition for tumoricidal response

### DSP502

**PVR inhibitor** –  
*Dual PD1/TIGIT  
inhibition with DNAMI  
potentiation potential*

**PD-L1 inhibitor** –  
*T and NK cell activation*

**Active IgG1 Fc** –  
*Half-life extension  
potential, ADCC activity*

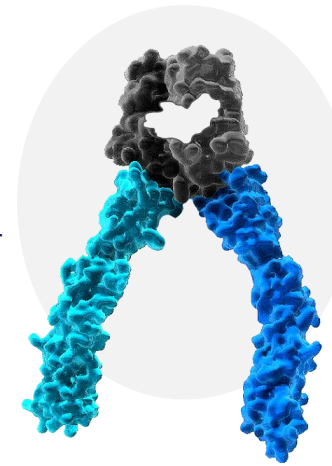


### DSP216

**CD47 inhibitor** –  
*Avidity driven for  
cancer specific  
blocking*

**HLA-G inhibitor** –  
*Inhibition of  
LILRB1, LILRB2*

**Inactive Fc** –  
*Half-life extension  
potential*





# Wholly Owned, Focused and Differentiated Pipeline

Program	Targets	Indications	Discovery	Preclinical	Phase 1	Phase 2	Phase 3	Anticipated Milestones
DSP107	CD47 4-1BB	Advanced Solid Tumors, NSCLC	DSP107 ± atezolizumab*					Phase II interim data Q2/23
		AML / MDS	DSP107 ± azacitidine + venetoclax					Phase Ib interim data Q4/22
DSP502	PVR PD-L1	Oncology						IND submission H1 2024
DSP216	HLA-G CD47	Oncology						IND submission H2 2024



\*Clinical trial collaboration and supply agreement with Roche for the PD-L1 inhibitor atezolizumab (TECENTRIQ®)

# Key Anticipated Milestones

Program MOA	Indication	2022		2023		2024	
		1H	2H	1H	2H	1H	2H
DSP107 CD47 inhibitor 4-1BB activator	Solid Tumors, NSCLC	Ph I Topline Results		Ph I/II NSCLC Interim Results	Ph I/II NSCLC Topline Results		
	AML/MDS	Ph Ib AML Interim Results		Ph Ib AML Topline Results			
DSP502 PVR inhibitor PD-L1 inhibitor Active IgG1 Fc	Oncology	IND-enabling activities				File IND	
DSP216 CD47 inhibitor HLA-G inhibitor Inactive Fc	Oncology	IND-enabling activities					File IND

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# DSP107

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MIRP Type

**DSP**

Targets

**CD47, 4-1BB**

Primary Cell Target

**mφ macrophages, T cells**

Mechanism of Action

**Unleash mφ via ‘Don’t Eat Me’ blockade, Activate 4-1BB+ T cells**

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# DSP107 – Differentiated CD47 Targeting Compound

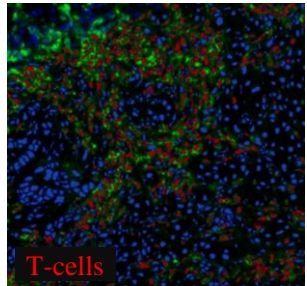
First-generation CD47 blocking agents are mainly active in hematological malignancies and limited by anemia

**Dual MOA**  
designed to activate innate and  
adaptive immunity

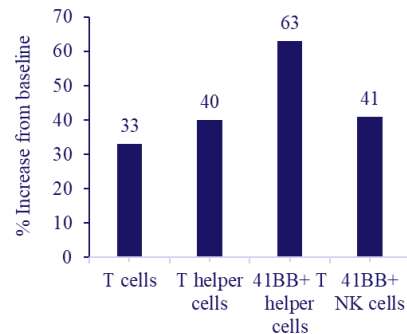
**Favorable safety**  
without hematological  
toxicities observed

**Strongly positioned**  
for treatment of solid and  
hematological malignancies

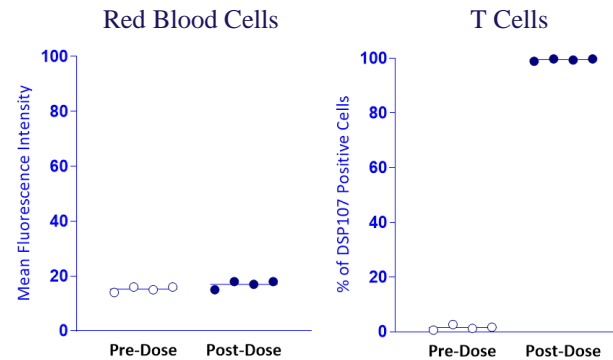
Phase I data demonstrates unique and differentiated features



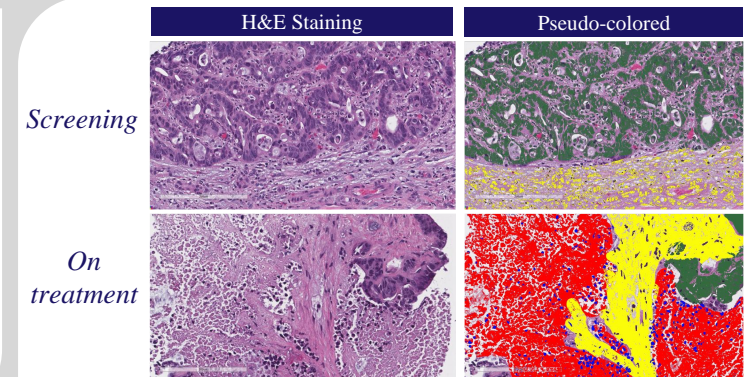
Post DSP107  
treatment biopsy data



Triggering T and NK-cell infiltration  
into tumor microenvironment



No binding to red blood cells observed



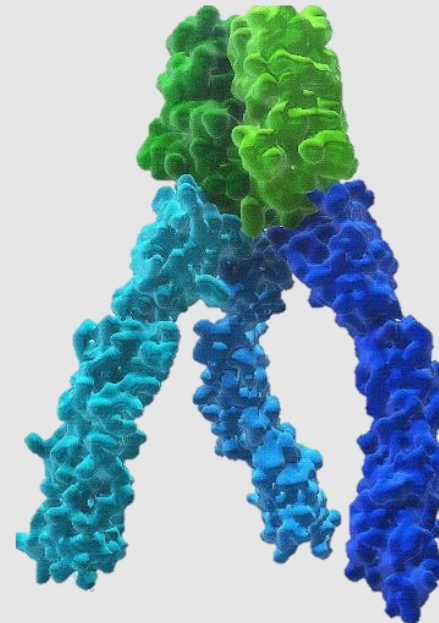
Increasing tumor necrosis (red) and  
immune cell infiltration (blue)

# Trimeric Structure Design For Tumor Selectivity and Improved Safety





## Trimeric ligand ends designed to enable:

- 1 Cancer selective binding to overexpressed CD47 on tumor cells, minimizing RBC binding and associated toxicities
- 2 Conditional 4-1BB mediated T cell activation dependent on trimeric binding to CD47 on cancer cells

Trimeric 4-1BBL



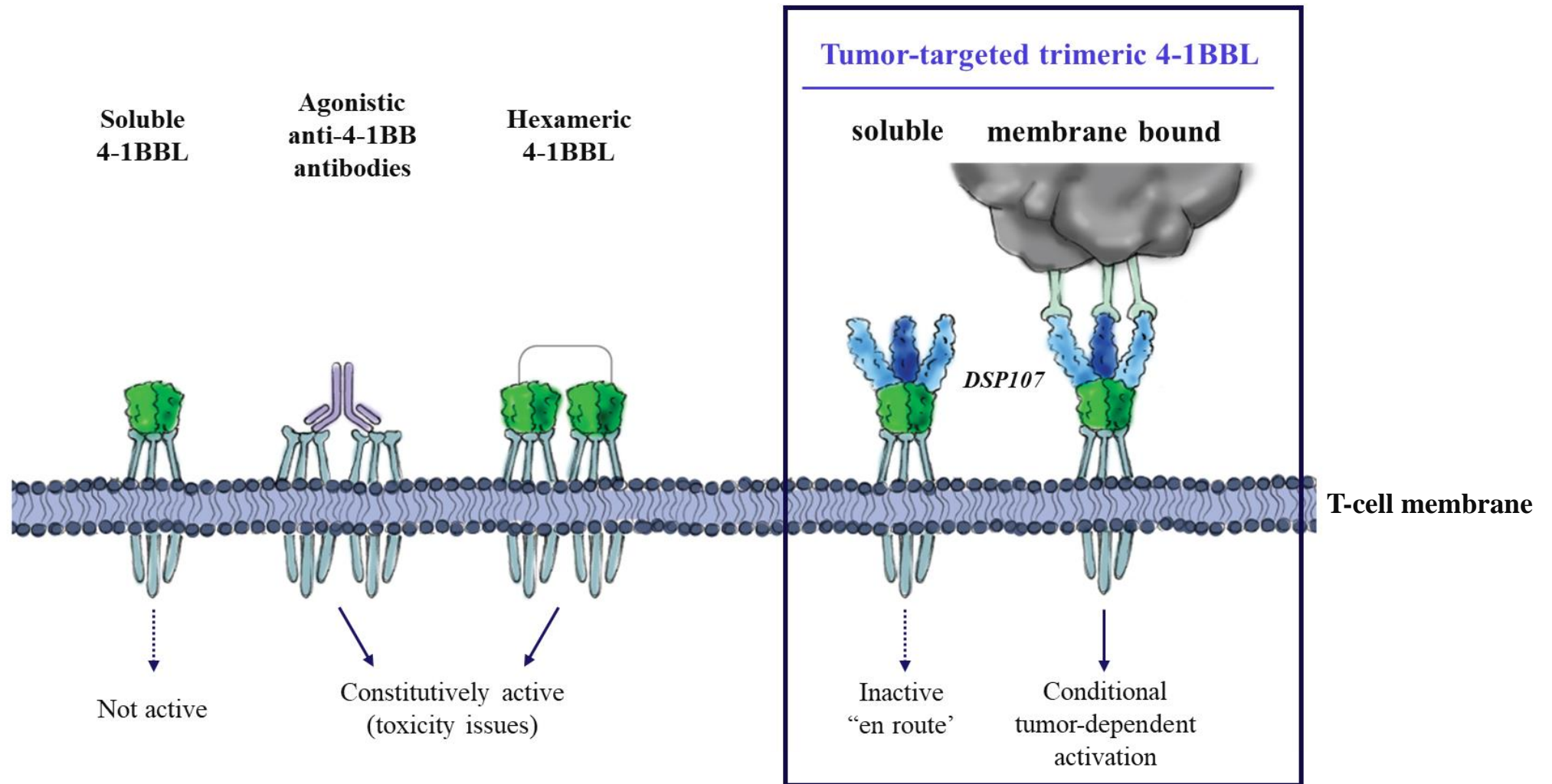
3 SIRP $\alpha$  for  
CD47 Checkpoint Targeting

-  Cytolytic T cell activation
-  T cell Proliferation
-  Checkpoint inhibition
-  Tumor microenvironment modulation

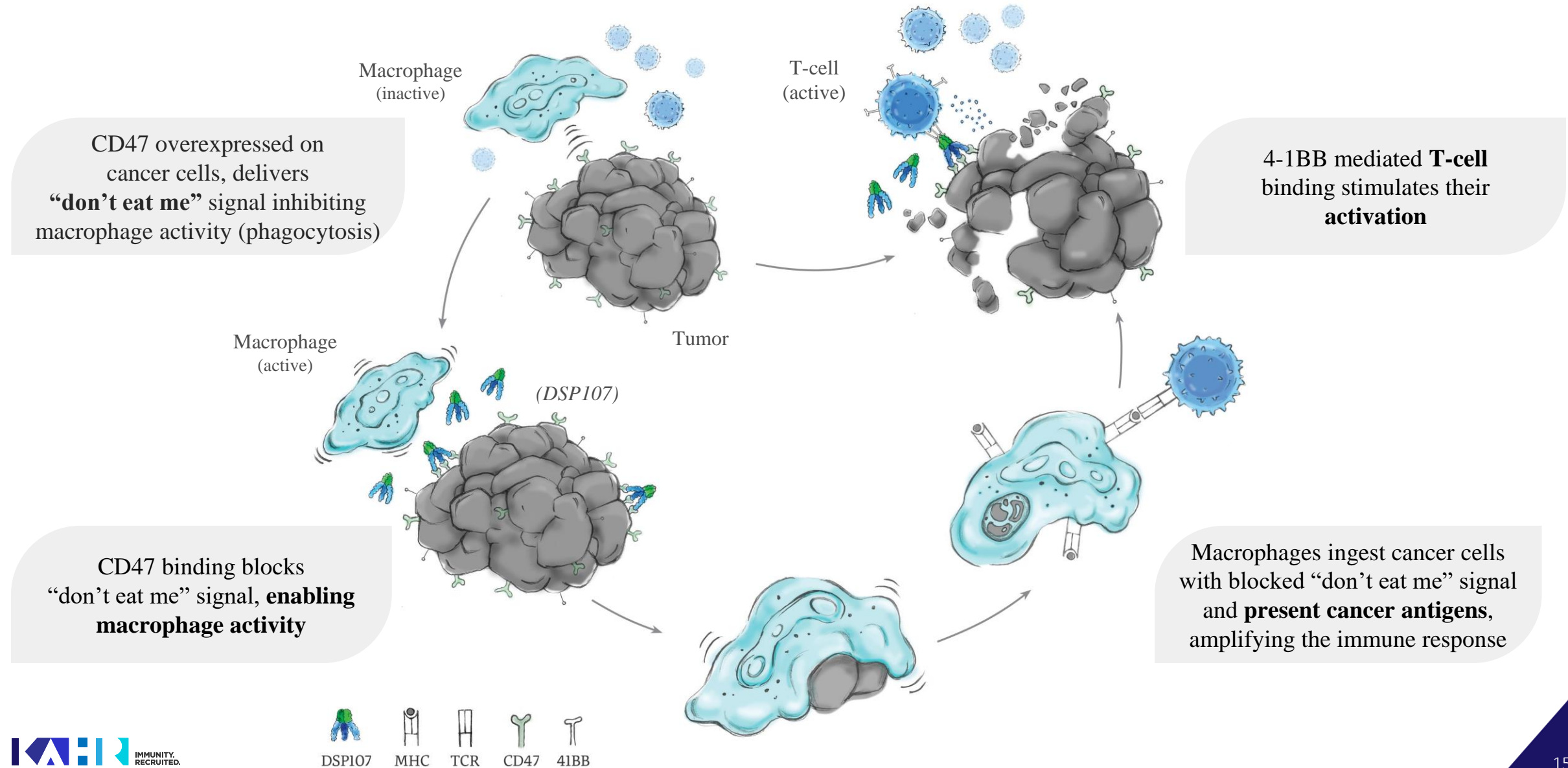


# DSP107 Structure Designed to Safely Unlock 4-1BB Potential

Phase I/II Data Demonstrated Absence of 4-1BB Related Hepatotoxicity



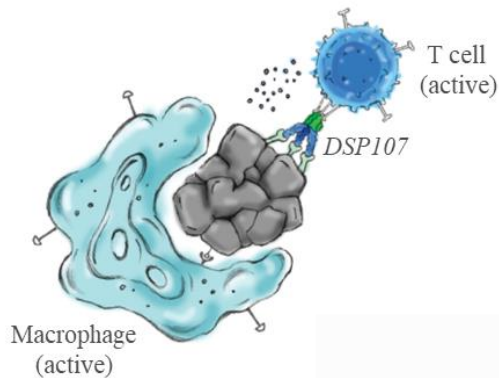
# Designed for Synergistic Innate & Adaptive Immune Activation



# DSP107 Potential as a Monotherapy and in Combination Therapies

## DSP107 monotherapy

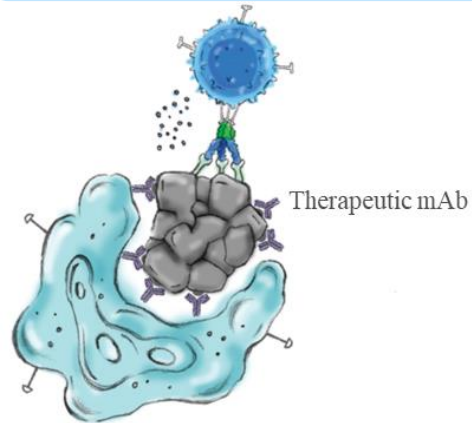
Triggers macrophage mediated phagocytosis and T cell cytotoxicity



## DSP107 combination with therapeutic Antibodies IgG1 mAbs

(cetuximab, trastuzumab...)

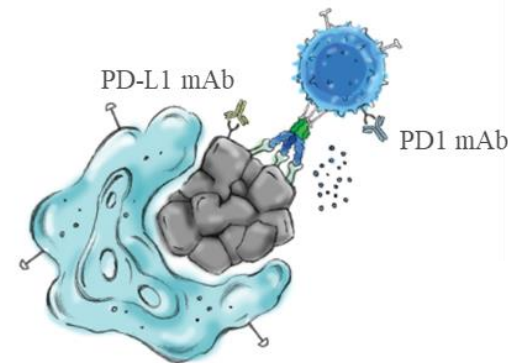
Enhances antibody-dependent cellular phagocytosis (ADCP)



## DSP107 combination with PD1/PD-L1 Checkpoint Inhibitors

(atezolizumab, pembrolizumab...)

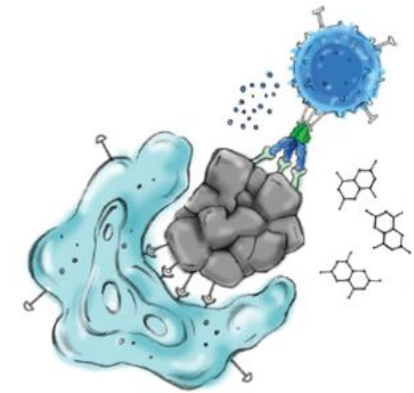
Enhances T cell activation



## DSP107 combination with pro-apoptotic agents

(chemotherapy, hypomethylating agents and BCL2 inhibitors)

Increases “eat me” signals



# Preclinical Studies Support Differentiated Potential Dual MoA

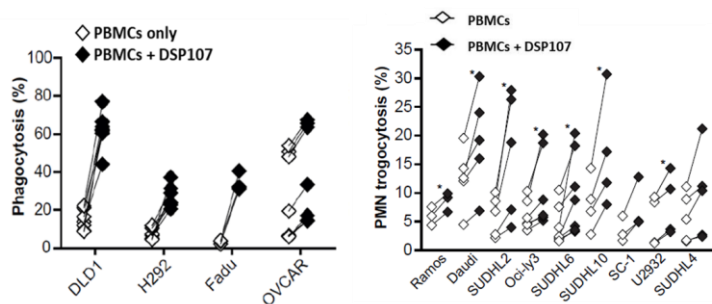
**Dual MOA**  
designed to activate innate and  
adaptive immunity

**Favorable safety**  
without hematological or hepato-  
toxicities in NHP observed

**Monotherapy potential**  
for treatment of solid and  
hematological malignancies

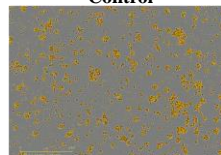
## Comprehensive preclinical package demonstrated differentiated features

Carcinoma and Lymphoma cell lines

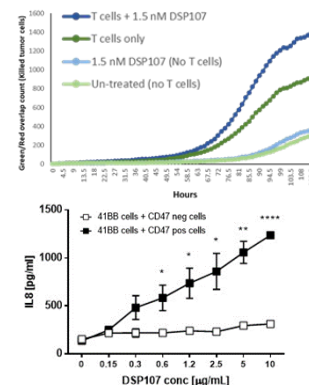
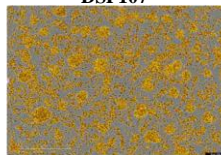


Triggering cancer cell death by  
phagocytosis as a single agent

Control

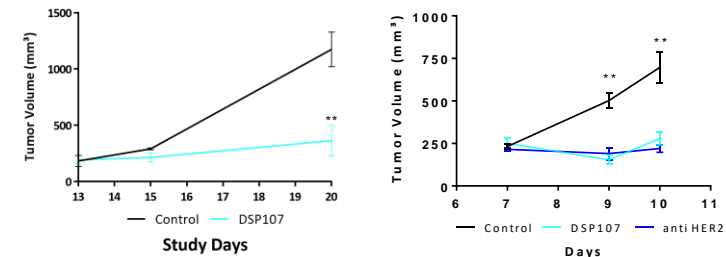


DSP107



Inducing 4-1BB activation, T-cell proliferation  
and killing potential against cancer cells

Lymphoma and Ovarian Carcinoma models

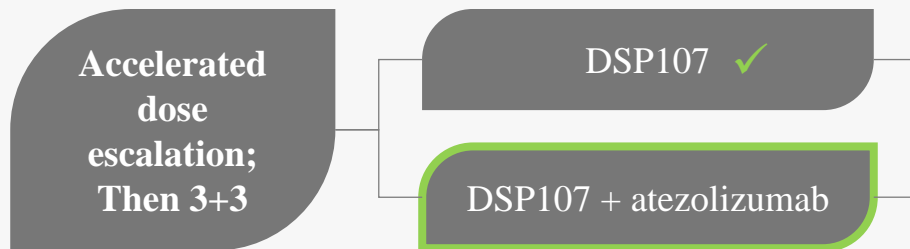


Demonstrating single agent anti tumor  
activity in mice models

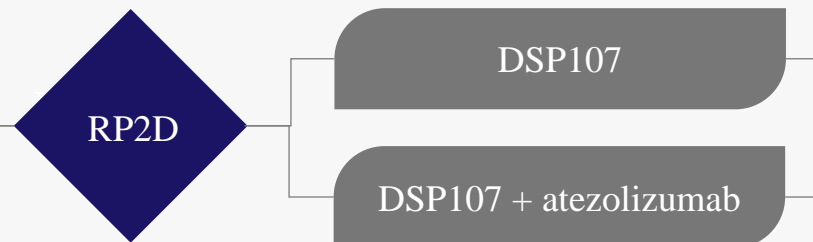
# DSP107 Clinical Program

## Advanced Solid Tumors

### Phase 1 – Dose escalation (n ~ 30)



### Phase 2 – Expansion Cohort (n = 50)



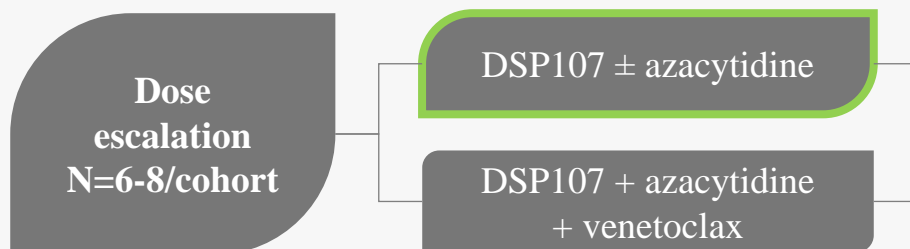
### Potential regulatory path



**Enrolling sites:** Pittsburgh, Colorado, Kansas, Thomas Jefferson; Sites under evaluation: San-Diego, Augusta, Chapel Hill, University of Texas

## Hematological Malignancies

### Phase I Dose escalation (n ~ 36)



**Phase 2 – Expansion**  
Details to be announced following EOPI meeting with the FDA

**Lead site:** MD Anderson Cancer Center

*RP2D: Recommended Phase 2 dose; NSCLC: Non-small cell lung cancer; MDS: Myelodysplastic syndromes; AML: Acute myeloid leukemia*

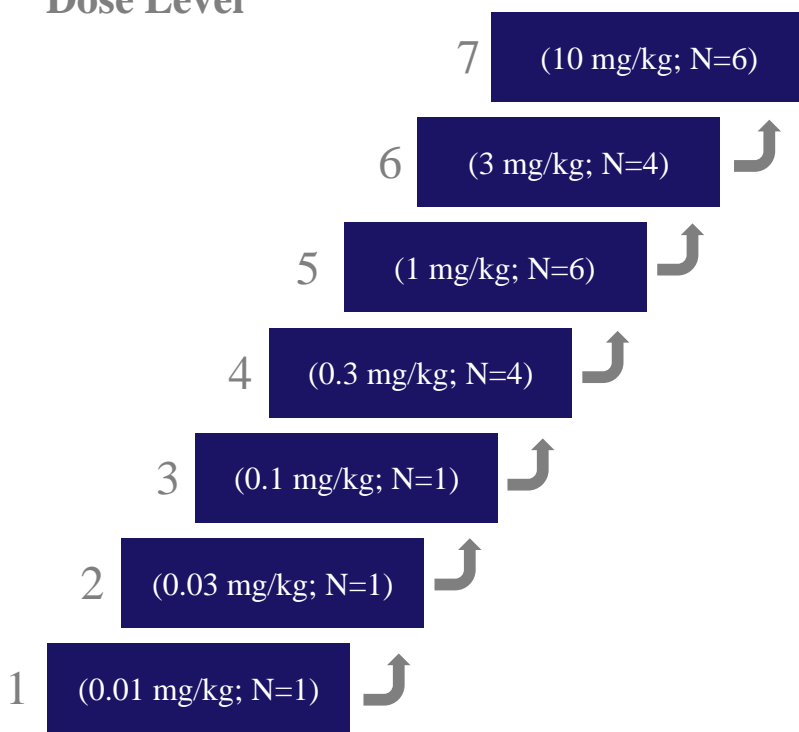


# Trial Design and Key Inclusion Criteria

## Part 1 – Monotherapy and Combination Dose Escalation in Advanced Solid Tumor Patients

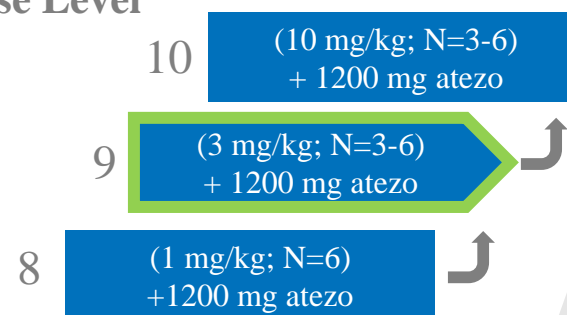
### DSP107 Monotherapy

#### Dose Level



### DSP107 + atezolizumab Combination

#### Dose Level



### Trial Design:

- Patients with advanced solid tumors (N=30) not suitable for curative therapy and without approved treatment options
- IV administration once weekly
- Accelerated dose escalation in single patient cohorts followed by standard 3+3 design

### Key Inclusion Criteria:

- Histologically confirmed advanced solid tumor with no approved therapeutic options
- Age 18 years or older
- ECOG performance status 0 or 1
- Measurable disease per RECIST v 1.1

# Patients With Advanced Solid Tumors

Nearly Half Failed Prior Immunotherapy and/or Cold Tumors

Characteristics	
<b>Total number of patients</b>	N = 23 (cohorts 1 – 7)
<b>Sex</b>	10 (43%) ♀; 13 (57%) ♂
<b>Age</b>	Median 63 (Range 29-78)
<b>Tumor types</b>	
Colorectal	7 (30%)
Pancreas	5 (22%)
Head and Neck	3 (13%)
NSCLC	1 (4%)
Ovarian	2 (9%)
Rare tumor types	5 (22%)
<b>Previous lines of therapy</b>	Median 3 (Range 2-8)
<b>PD1/PD-L1 experienced</b>	11 (48%)

# Well Tolerated Without DLTs, Hematological or Hepato-Toxicities

## Summary

- DSP107 doses up to and including 10 mg/kg well tolerated
- No DLTs and no treatment-related SAEs
- No hematological toxicities
- No hepato-toxicities
- Very few AEs considered related to DSP107 and almost all mild or moderate in severity
- Most related AEs Grade 1-2 in severity. Only 2 related Grade 3 AEs – transient hypertension and fatigue (at EOT visit)

Now treating patients with DSP107 and Atezolizumab combination

## Treatment-Related AEs in $\geq 2$ Patients

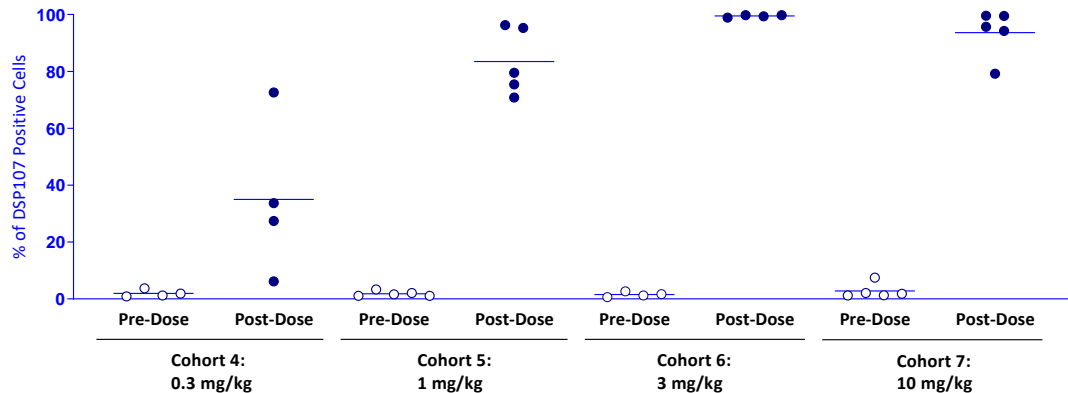
Total No of Patients	N = 23 (cohorts 1 - 7)
Treatment-related AEs (any grade)	n (%)
Any	16 (70)
IRR*	7 (30)
Diarrhea	4 (17)
Fatigue	4 (17)
Nausea	3 (13)
Constipation	2 (9)

\*IRRs Grade 1-2 in severity. Easily abrogated (except 10-004 and 11-012 who withdrew consent) in subsequent infusions by reduced rate of infusion and concomitant IV fluids.

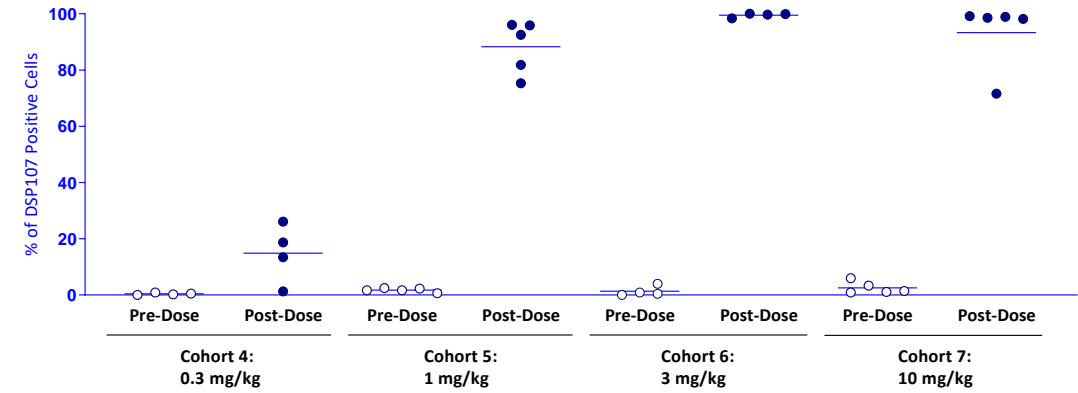
# Receptor Occupancy Data Showed Immune Cell Target Engagement With No RBC Binding

Dose dependent target engagement achieved with 100% receptor occupancy on circulating immune cells observed from 3 mg/kg

## Binding to T cells



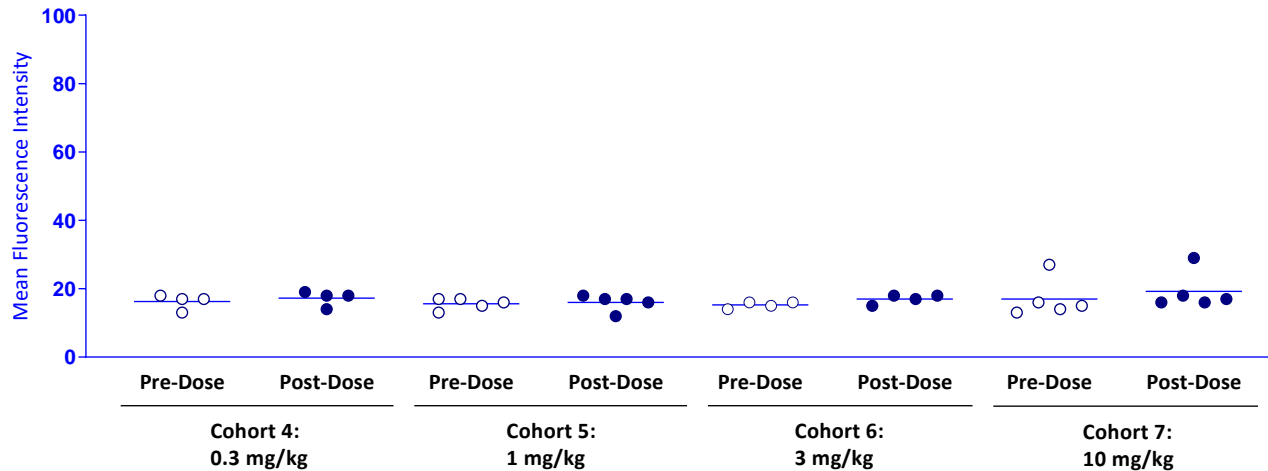
## Binding to NK cells



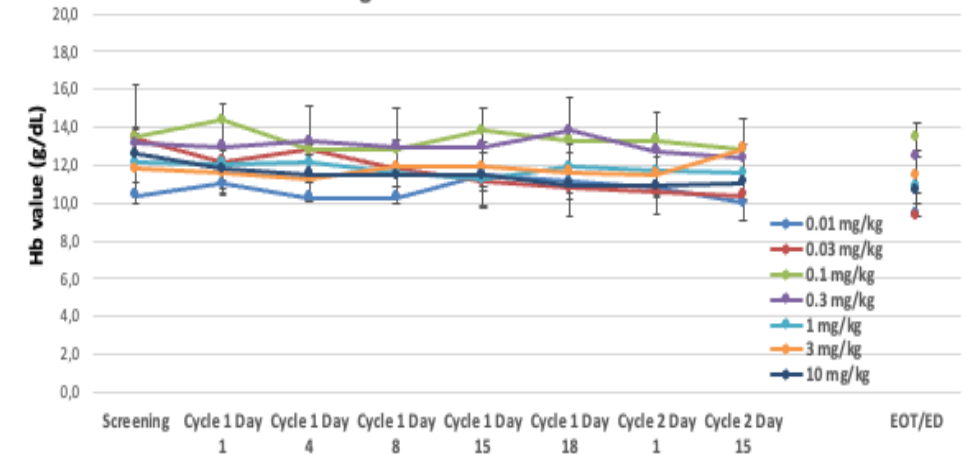
# DSP107 Does Not Bind Red Blood Cells

Resulting in Favorable Safety Profile With No Anemia or Antigen Sink Issues

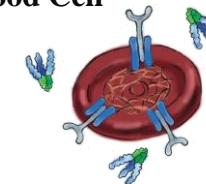
## Binding to Red Blood Cells



## Mean Hemoglobin Values on DSP107 Treatment



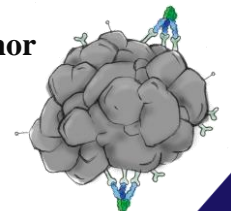
## Red Blood Cell



On RBCs CD47 protein complex is anchored to cytoskeleton resulting in its immobilization and low affinity of DSP107 to the monomeric CD47

High affinity/avidity of DSP107 to CD47 clusters on cancer cells

## Tumor





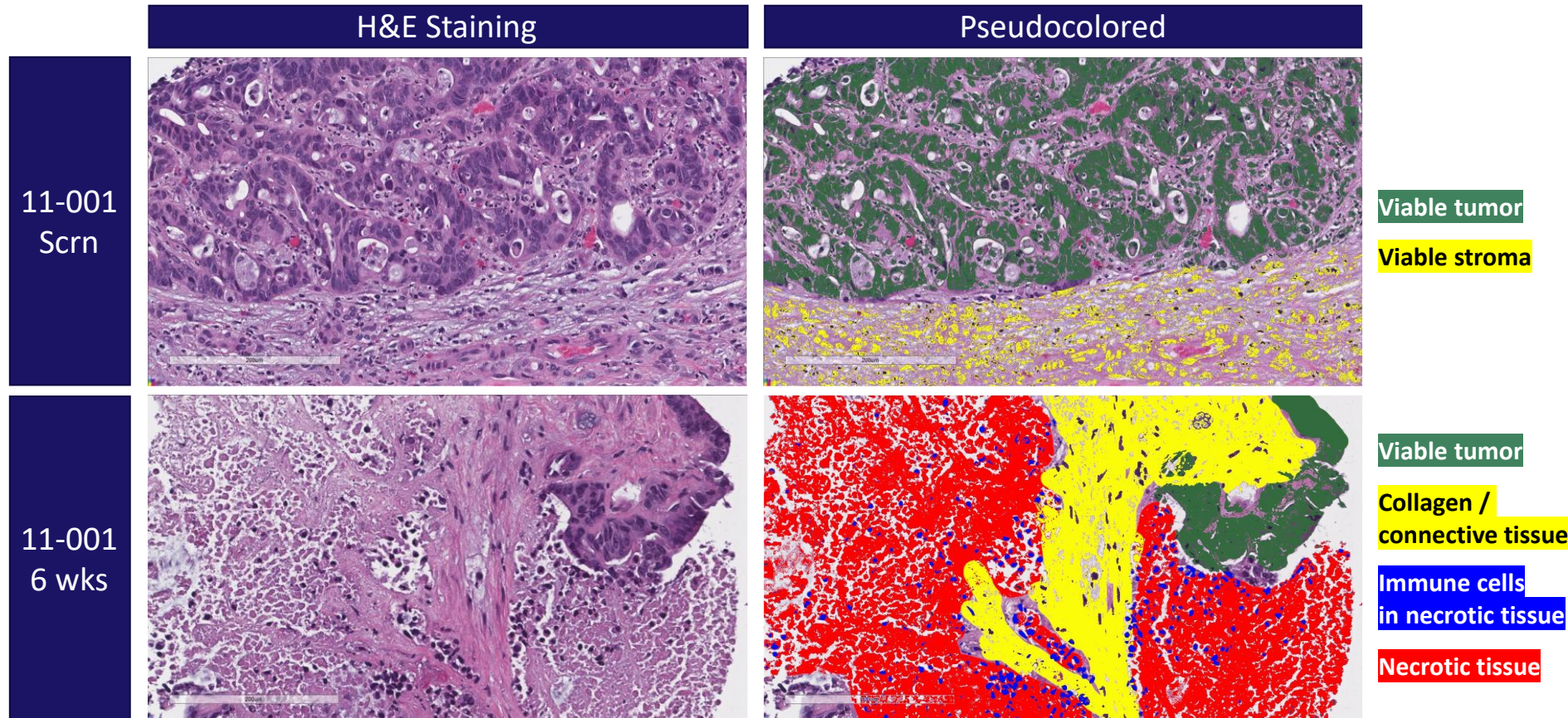
# DSP107 Treatment Resulted in Tumor Necrosis

## Key paired biopsies data

- All biopsies collected from hepatic metastases pre-treatment and following cycle 2 (6 doses)
- H&E stained slides assessed by independent, blinded pathologist
- In 3 out of 4 paired biopsies significant increase in necrotic (dead) tumor tissue was observed
- Necrosis associated with immune cell infiltration; no evidence of vascular necrosis

Patient Number	Dose (mg/kg)	Tumor type	Timepoint	% Necrosis
11-001	0.3	Colorectal	Screening	0
			6 weeks	65
11-002	0.3	Colorectal	Screening	2
			6 weeks	35
10-003	1	Pancreatic	Screening	10
			6 weeks	50
13-005	1	Pancreatic	Screening	4
			6 weeks	3

# Case study: Increased Tumor Necrosis Associated With Immune Cell Infiltration



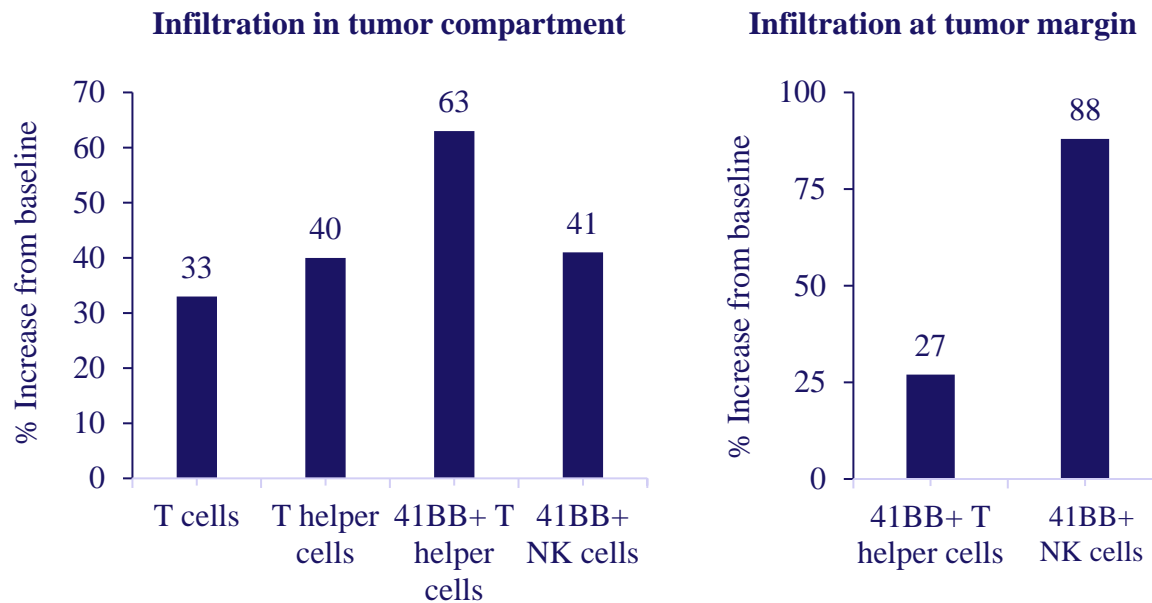
**~ 65%**  
of the tumor tissue  
necrotic 6 weeks post  
DSP107 treatment  
(no necrosis at baseline)

Paired biopsy from colon carcinoma patient (11-001) in dose level 4 (0.3 mg/kg) pre- treatment and following cycle 2 (6 doses)

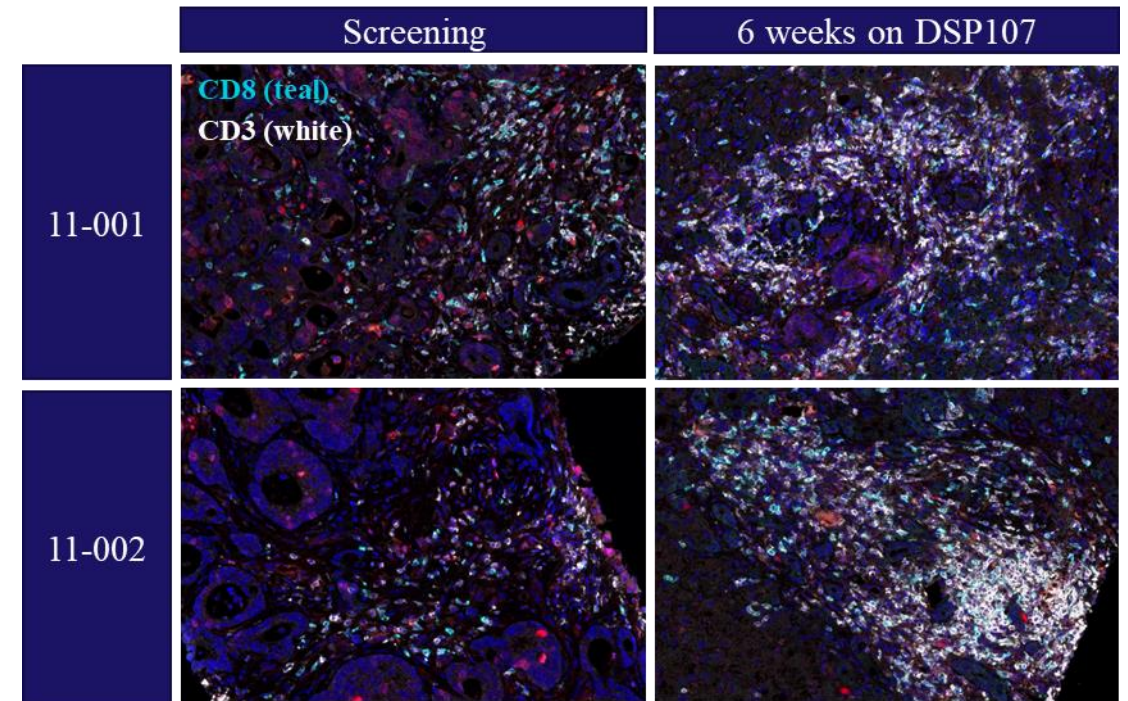


# Case Study: Paired Biopsies Demonstrated Adaptive Immune Engagement in Ongoing Clinical Trial

6-wks post DSP107 treatment

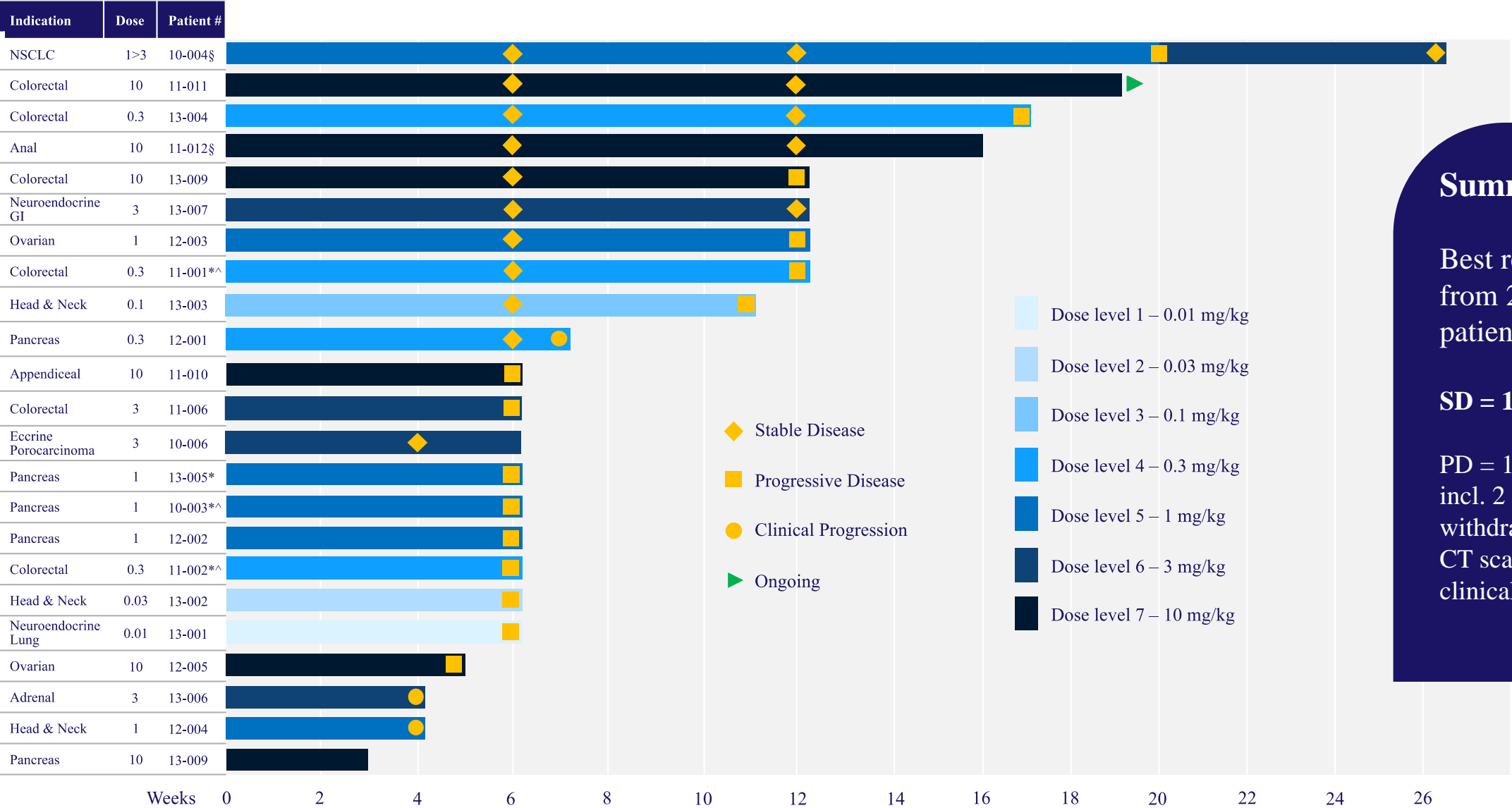


Paired biopsy from colon carcinoma patient (11-001) in dose level 4 (0.3 mg/kg) pre- treatment and following cycle 2 (6 doses). Quantification of multiplex image analysis from biopsy stains.



*Significant infiltration of T cells and NK cells in both the tumor compartment and at the tumor margin following DSP107 treatment*

# Best Overall Response After DSP107 Monotherapy



## Summary

Best response from 22 evaluable patients:

SD = 11 (50%)

PD = 11 (50%)  
incl. 2 patients withdrawn before 1<sup>st</sup> CT scan due to clinical progression

# DSP107 Phase I/II Data: Favorable Preliminary Safety and Activity in Advanced Solid Tumors

## Clinical Overview

- DSP107 as monotherapy and in combination with Atezolizumab is being evaluated in a dose escalation trial
- 23 patients with diverse solid tumors have been treated with DSP107 monotherapy with 22 patients evaluable for efficacy analysis
- Now treating patients with DSP107 and Atezolizumab
- Phase Ib in r/r AML and MDS is ongoing assessing the safety of DSP107 monotherapy, DSP107 with Azacitidine and DSP107 with Azacitidine plus Venetoclax

## Key Findings

- Low-grade AEs with no DLTs, no hematological toxicities and no hepato-toxicities
- Receptor occupancy data suggesting lack of RBC binding and immune cell engagement
- Increased immune cell infiltration into the tumor and tumor necrosis
- 50% DCR in difficult to treat phase I patients

Next steps: Further evaluate safety and efficacy of DSP107 monotherapy and combination with SOC therapies

# DSP107 Highlights



## MOA

- CD47 inhibition (Cancer specific)
- 4-1BB activation (CD47-conditional)



## Potential Efficacy - Preclinical

- Activates T cells, increases IFN $\gamma$  secretion and anti-tumor killing
- Increased macrophage phagocytosis of tumor cells
- Augments mAbs' ADCC phagocytosis of cancer cells
- Potential efficacy as monotherapy and synergistic activity in combination



## Differentiation

- Activates both adaptive and innate immunity
- No RBC binding observed
- Potentially favorable safety profile suitable for combination therapy



## Potential Safety - Preclinical

- No binding to human RBCs observed
- No CD47 related hematological toxicities observed
- No 4-1BB related hepato-toxicities observed



## Clinical Opportunities

- Solid tumors
  - DSP107  $\pm$  atezolizumab (intended to enhance T cell activation)
- Hematological Malignancies
  - DSP107  $\pm$  azacytidine + venetoclax (intended to enhance eat me signal)



## Preliminary Clinical Data

Well tolerated

- No DLTs
- No hematological toxicities
- No hepato-toxicities

Preliminary efficacy signals as a monotherapy in solid tumors

- Paired biopsies demonstrate tumor necrosis
- Stable disease achieved in eleven of twenty-two patients (50%)



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# DSP502

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MIRP Type

**DSP-Fc**

Targets

**PVR, PD-L1, FcR**

Primary Cell Target

**NK cells, T effector cells**

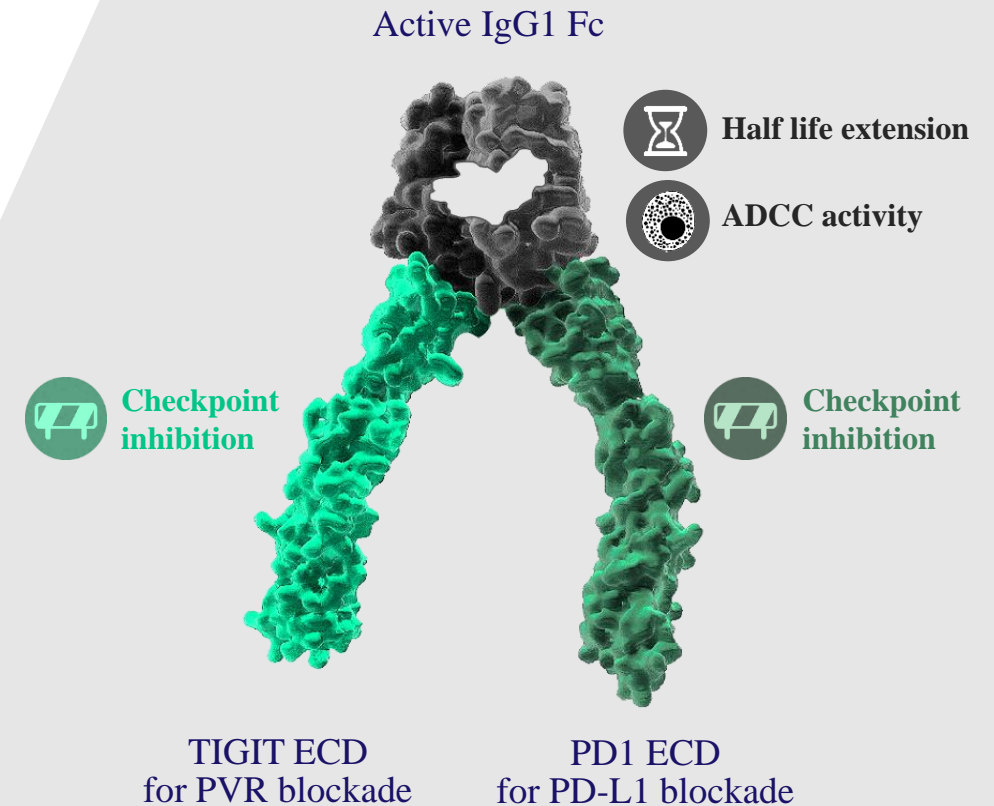
Mechanism of Action

**Dual checkpoint inhibition designed to unleash NK and T-cells, ADCC**

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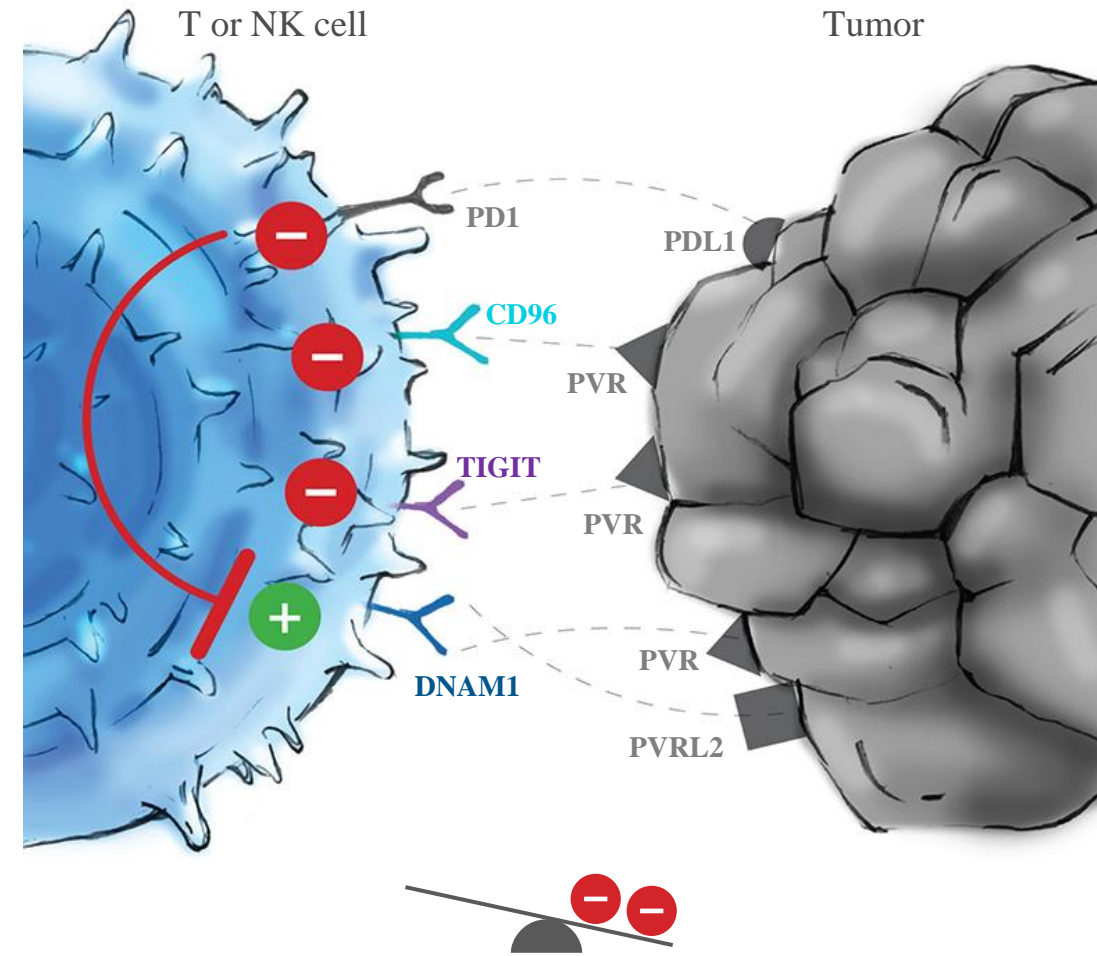
# Dual Checkpoint Binding Aimed at Enhancing Selectivity and Synergistic Immunity

- 1 Dual binding to overexpressed checkpoints may enable high tumor specificity
- 2 Potential anti-tumor immunity via simultaneous checkpoint inhibition of PVR and PD-L1
- 3 Active Fc may extend half-life and enhance ADCC activity



# Novel, Potential Next-Generation Approach to Checkpoint Inhibition

- PVR is a ligand for inhibitory (TIGIT, CD96) and stimulatory (DNAM1) immune checkpoint receptors
- Tumor cells overexpress PVR to evade immune surveillance
- Overexpression of PVR is associated with resistance to checkpoint therapy in NSCLC and melanoma patients
- Inhibition of TIGIT/PVR is efficacious in combination with PD1 blockade
- PD1 inactivates DNAM1 stimulatory activity



Immunosuppression is the net effect of PVR and PD1 overexpression in the TME

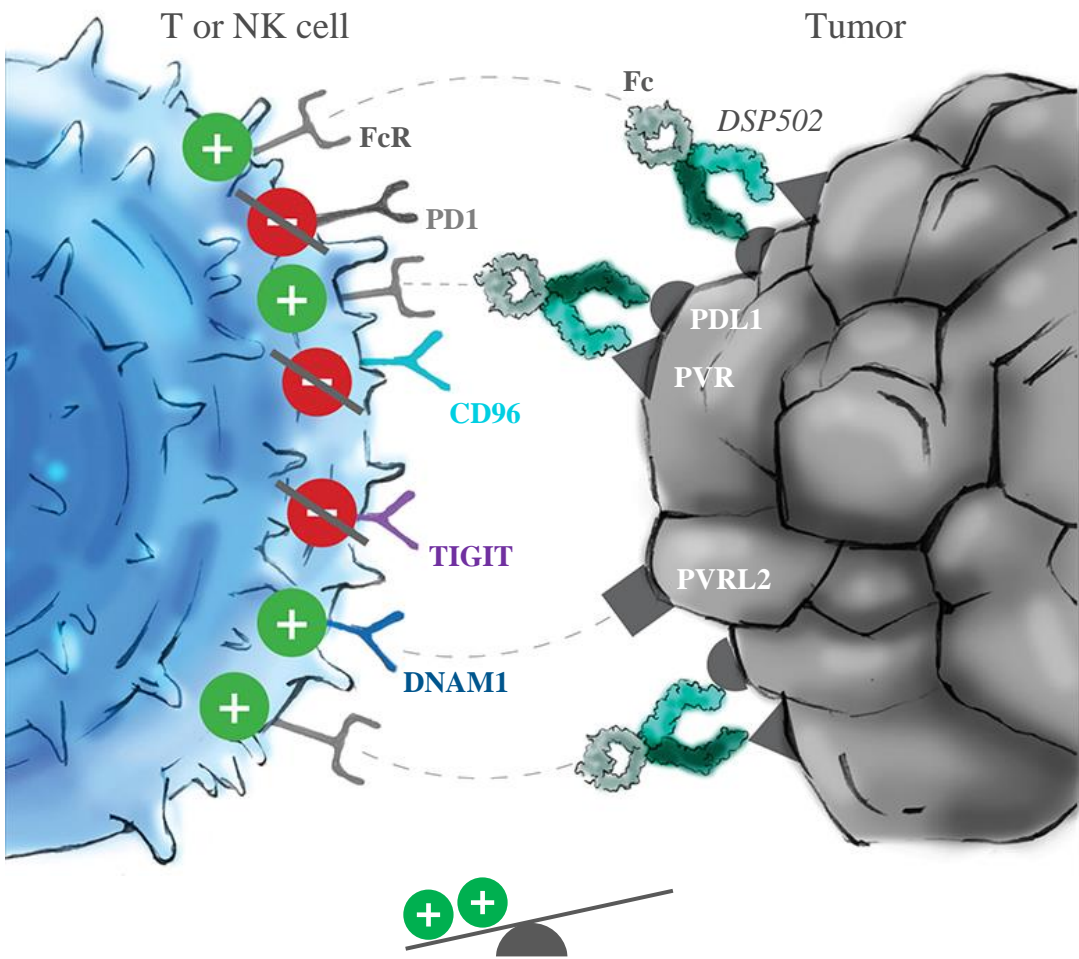


# Synergistic Dual Checkpoint Inhibition for Robust Anti-tumor Immunity

- 1 Simultaneous TIGIT, CD96 and PD-L1 inhibition with DNAM1 costimulation for enhanced anti-tumor immunity
- 2 Designed to activate both, T cells and NK cells

Effect	Dual PVR and PD-L1 Targeting (KAHR's approach)	TIGIT Ab* (Competitors)
<div>-</div> TIGIT	<div>✓</div>	<div>✓</div>
<div>-</div> CD96	<div>✓</div>	<div>-</div>
<div>+</div> DNAM1	<div>✓</div>	<div>-</div>
<div>-</div> PD-1	<div>✓</div>	<div>-</div>

\*Company has not undertaken comparative trials of DSP502 against the identified competitors



DSP502 blocks PVR and PD1 signaling for enhanced immune activation and anti-tumor immunity



# DSP502 – Differentiated TIGIT/PD1 Targeting Compound

Potential next-generation capabilities

## Dual MOA

designed to activate innate and adaptive immunity

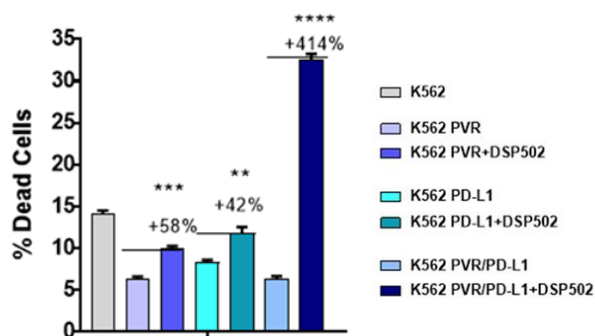
## High Tumor Specificity

Concomitant binding to PVR and PD-L1 required for its activity

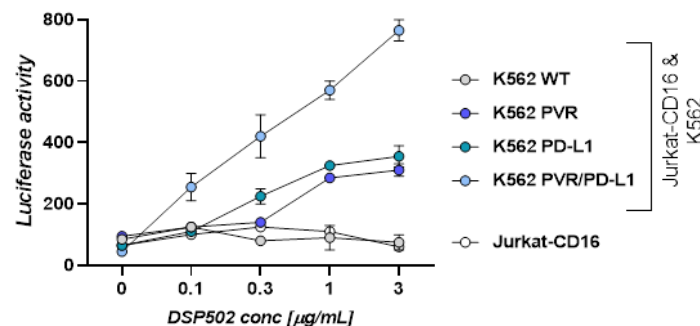
## Designed to Have Unique Features

Multiple functionalities that act simultaneously for synergistic effect

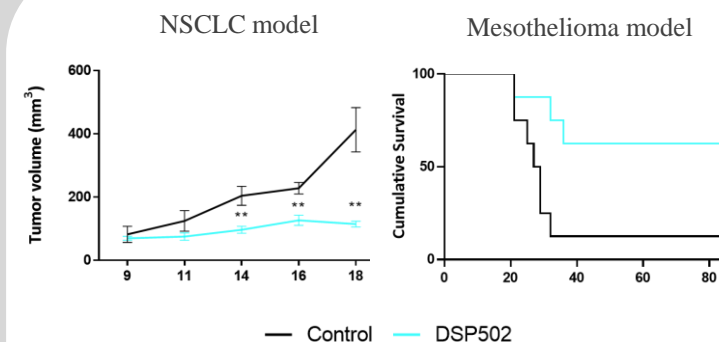
## Differentiated mechanism of action



Enhanced NK cells cancer killing potential



Augmented NK cells ADCC activity



Demonstrated potent single agent anti tumor activity

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# DSP216

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MIRP Type

**DSP-Fc**

Targets

**CD47, HLA-G**

Primary Cell Target

**mφ macrophages, T effector cells, NK cells, Myeloid cells**

Mechanism of Action

**Dual checkpoint inhibition unleash macrophage, NK and Teff**

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# Dual Checkpoint Binding Designed For Enhanced Selectivity and Broad Immunity

1

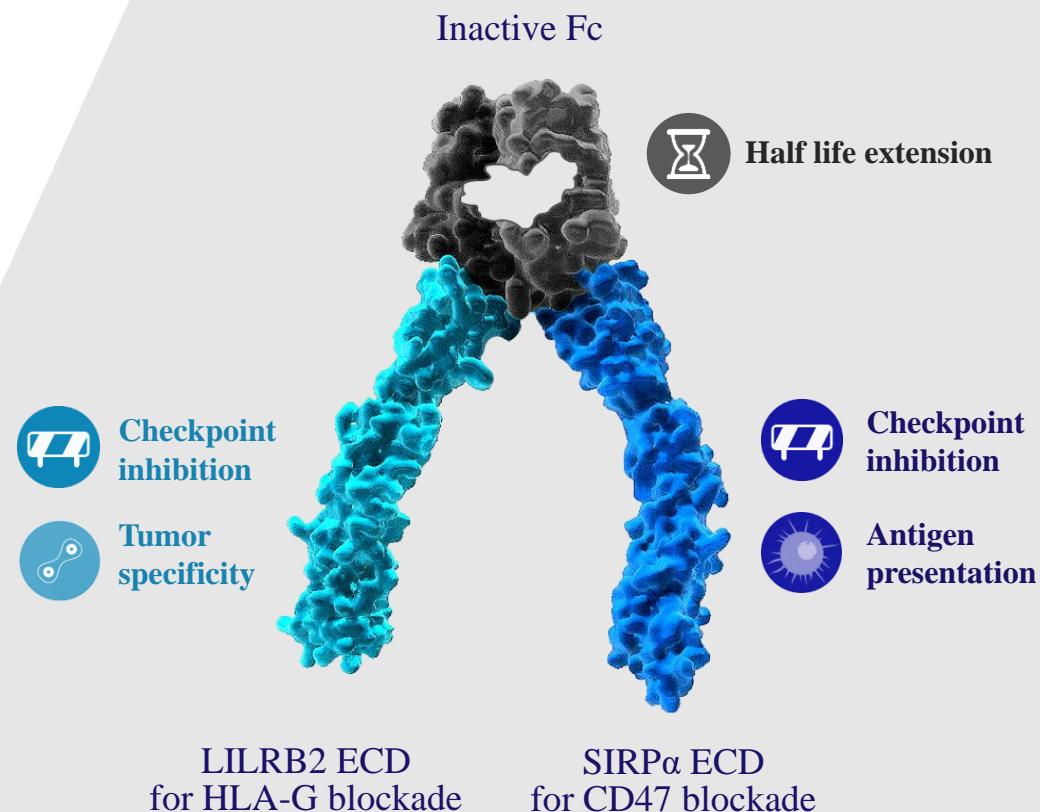
Dual binding to overexpressed cancer checkpoints may enable high tumor specificity

2

HLA-G and CD47 blockade designed to activate innate and adaptive immunity

3

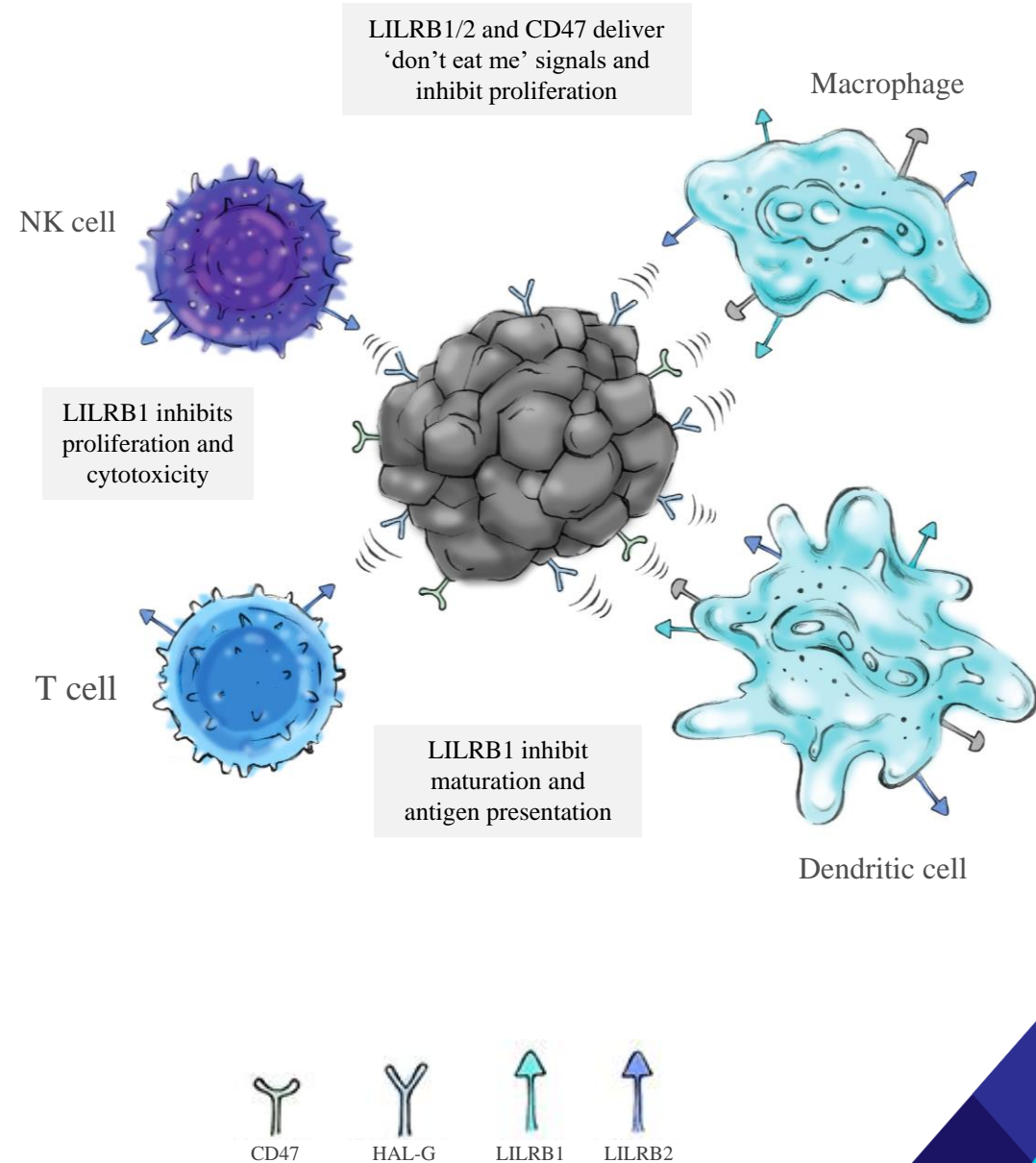
Inactive Fc may extend half-life





# HLA-G Blockade Designed To Prevent Immunotolerance Across Immune Cells

- Placenta expressed HLA-G triggers immunotolerance to prevent the mother's immune system from attacking the fetus
- Tumor cells overexpress HLA-G to evade immune surveillance
- HLA-G is a broad-range immune checkpoint that is the main ligand for the LILRB/ILT receptor family
- HLA-G inhibits multiple immune cell subsets and recruits suppressive immune cells to the tumor microenvironment

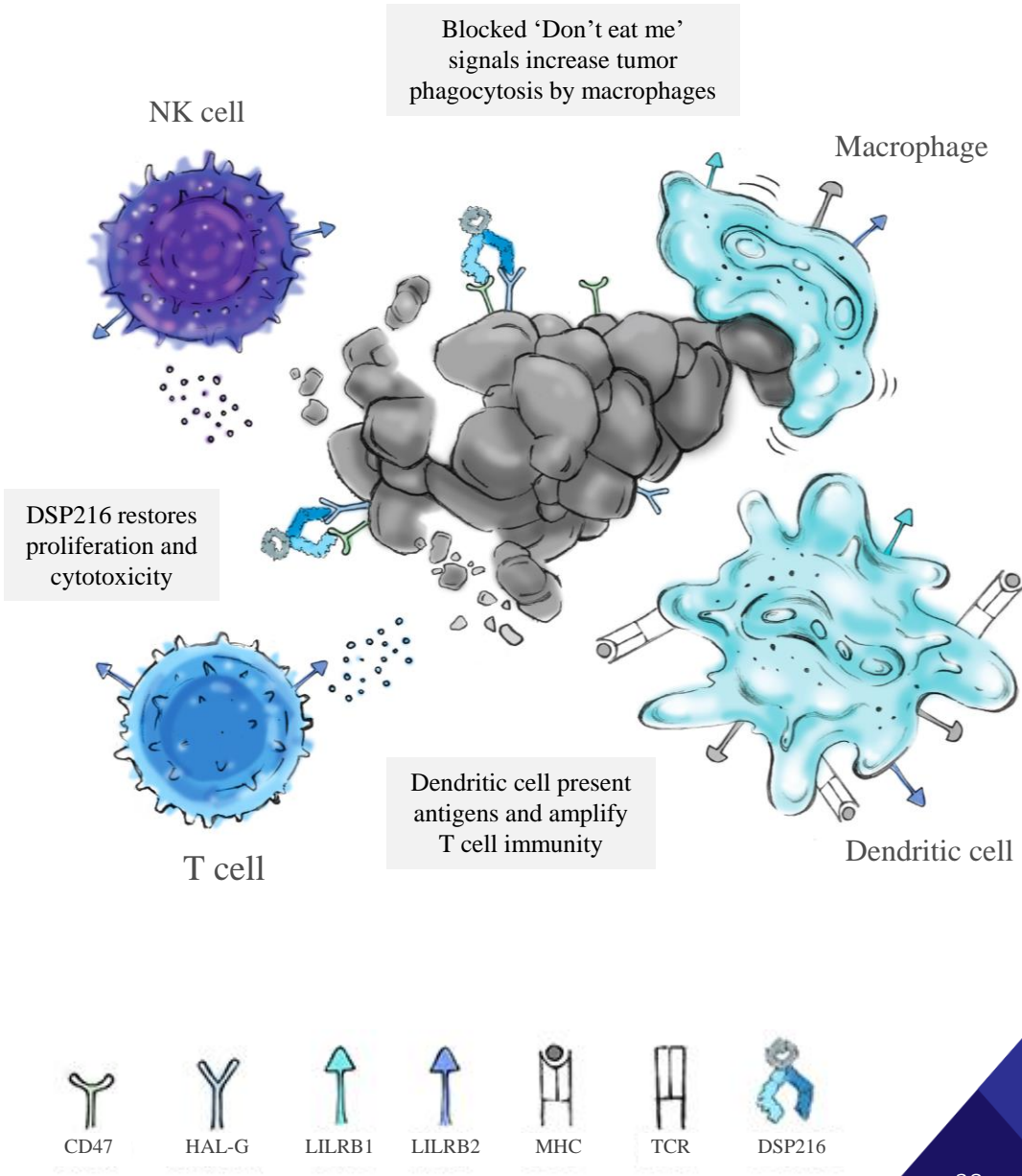


# Dual Targeting Designed For Robust Tumor Selective Innate & Adaptive Immune Activation

- 1 HLA-G blockade aims to prevent cancer immunotolerance by **multiple immune cell subsets**
- 2 CD47 blockade removes ‘don’t eat me’ signal, triggering macrophage phagocytosis of tumor cells

Effect	HLA-G & CD47 (KAHR’s approach)	LILRB1/2 Ab* (Competitors)
<div>–</div> LILRB1 & LILRB2	<div>✓</div>	<div>—</div>
<div>+</div> Tumor selectivity	<div>✓</div>	<div>—</div>
<div>+</div> Innate & adaptive immunity	<div>✓</div>	<div>—</div>

\*Company have not undertaken comparative trials of DSP502 against the identified competitors



# CORPORATE HIGHLIGHTS

# Multifunctional Cancer Immunotherapy Candidates Targeting Innate and Adaptive Immune Cells



## NOVEL MIRPs™

Multifunctional Immuno-Recruitment Proteins – versatile platform targeting both innate & adaptive immunity across cancers



## DIFFERENTIATED PIPELINE

- Novel, next-generation potential across three programs
- Lead candidate DSP107 –  
CD47 inhibition (Cancer specific)  
41BB activation (CD47-conditional)



## ANTICIPATED MILESTONES

- DSP107** | Interim Ph II solid tumor mono and combo data Q2 2023 and interim Ph I hematological malignancy data Q4 2022
- DSP502 & DSP216** | IND 2024



## MARKET

Immuno-therapeutics  
\$50.2B by 2026



## IP

13 families  
4 granted (US and other territories),  
73 pending (NP worldwide and PCT stage)



## Experienced Leadership

Management team, BOD and SAB comprised of leading experts including technology inventor, Prof. Mark Tykocinski, Dean of the School of Medicine and Provost, Thomas Jefferson University.

# Scientific Advisors and Board of Directors

## Scientific and Clinical Advisory Board

**Mark L. Tykocinski,  
MD**

KAHR technology inventor;  
BOD Observer; Provost  
Jefferson Thomas University



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Memorial Sloan Kettering  
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Medical Oncology,  
Weill Cornell



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Bioscience, co-founder of  
RayzeBio; 25+ years as a  
leader in biotech



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Chairman and owner  
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25+ yrs in biotech and  
life sciences



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Partner at ALIVE  
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investing in biotech

