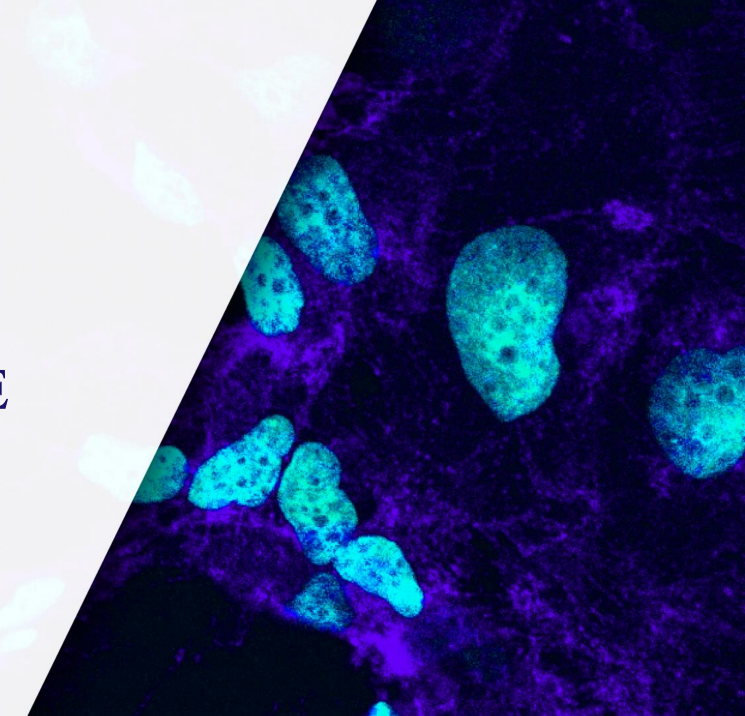


UNMASKING CANCER CELL CAMOUFLAGE

COMPANY PRESENTATION | **June 2022**



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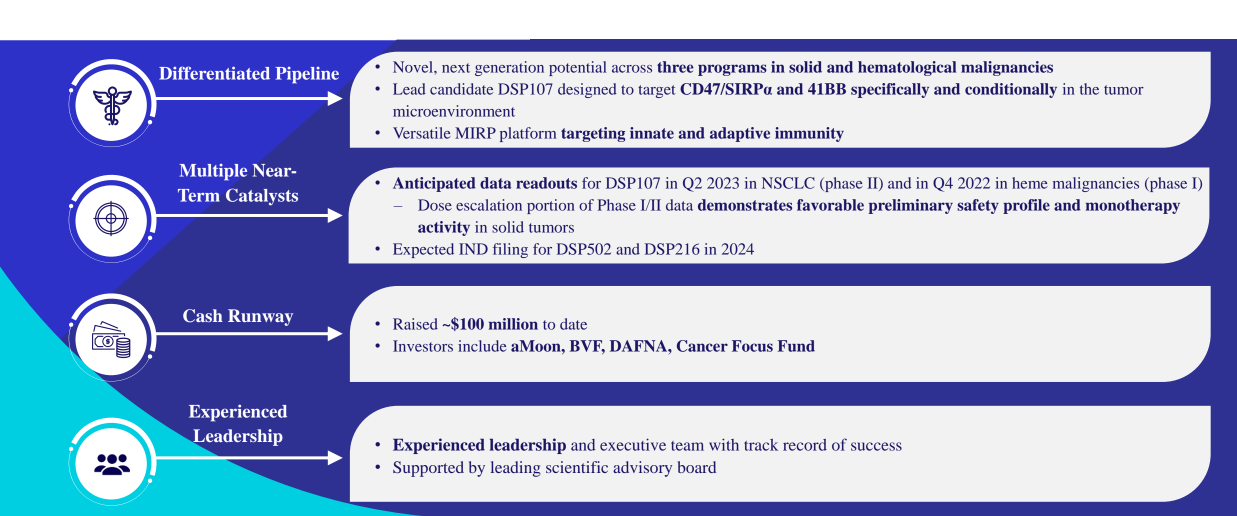
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Company Highlights



Experienced Leadership Team



Aron Knickerbocker, MBA

Board Chairman



Bristol Myers Squibb





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





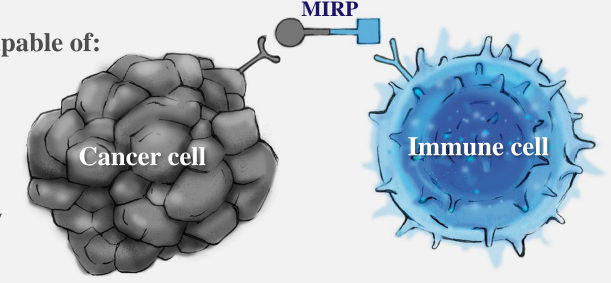
Versatile Multifunctional Immunotherapeutic Platform for Solid and Hematological Malignancies

MULTIFUNCTIONAL IMMUNE RECRUITMENT PROTEIN (MIRP)

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

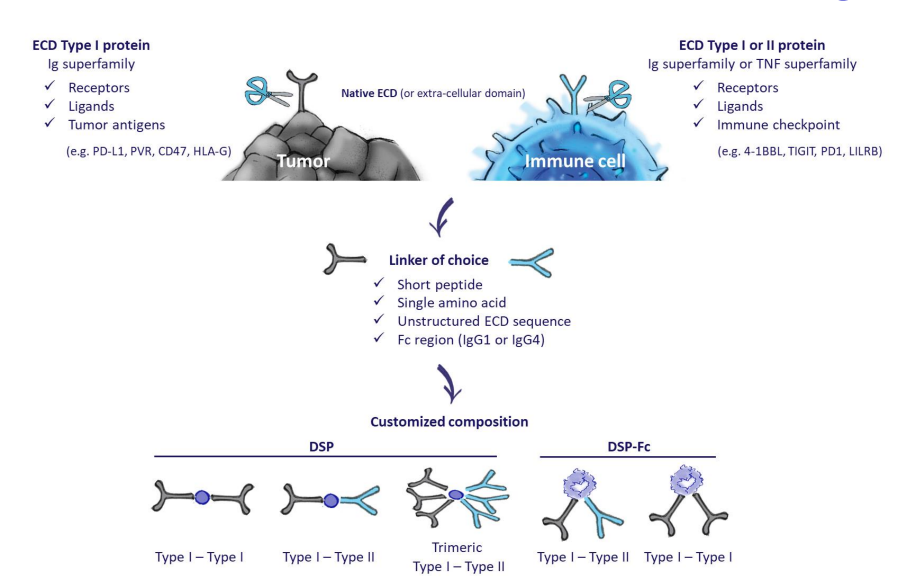
1 Inhibiting key evasion markers on cancer cells

2 Activating innate and adaptive anti-tumor immunity





MIRP Platform - Customized, Modular and Flexible Design





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

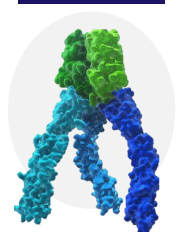
DSP (Dual Signaling Protein)

Combined checkpoint inhibition and immune co-stimulation

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

CD47 inhibitor – Trimeric binding for cancer specific blocking





DSP-Fc (Dual Signaling Protein With Fc Domain)

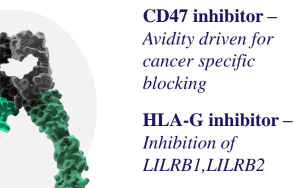
Dual checkpoint inhibition for tumoricidal response

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

PD-L1 inhibitor – T and NK cell activation

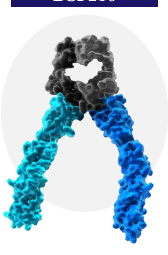
Active IgG1 Fc -Half-life extension potential, ADCC activity





Inactive Fc – Half-life extension potential

DSP216





Wholly Owned, Focused and Differentiated Pipeline

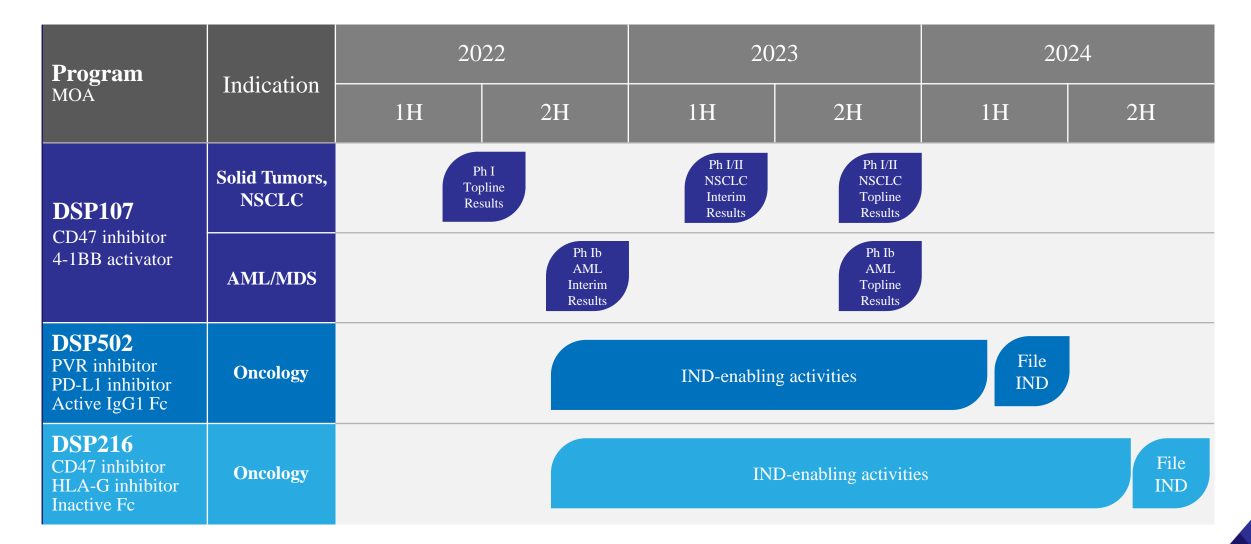
Program	Targets	Indications	Discovery	Preclinical	Phase 1	Phase 2	Phase 3	Anticipated Milestones
DSP107	CD47 4-1BB	Advanced Solid Tumors, NSCLC	DSP107 ± ate	zolizumab*				Phase II interim data Q2/23
		AML/MDS	DSP107 ± aza	citidine + veneto	clax 🖒			Phase Ib interim data Q4/22
DSP502	PVR PD-L1	Oncology		(\$				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





DSP107

MIRP Type	DSP
Targets	CD47, 4-1BB
Primary Cell Target	mø macrophages, T cells
Mechanism of Action	Unleash mφ via 'Don't Eat Me' blockade, Activate 4-1BB+ T cells

DSP107 – Differentiated CD47 Targeting Compound

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

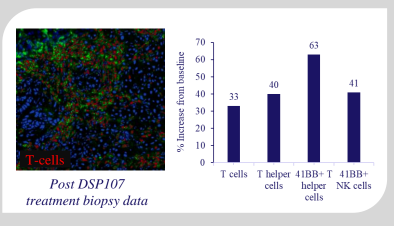
Dual MOA

designed to activate innate and adaptive immunity

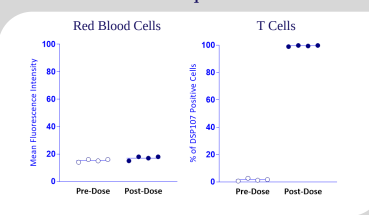
Favorable safety without hematological toxicities observed

Strongly positioned for treatment of solid and hematological malignancies

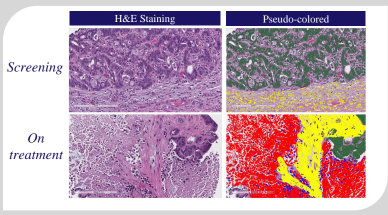
Phase I data demonstrates unique and differentiated features



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:

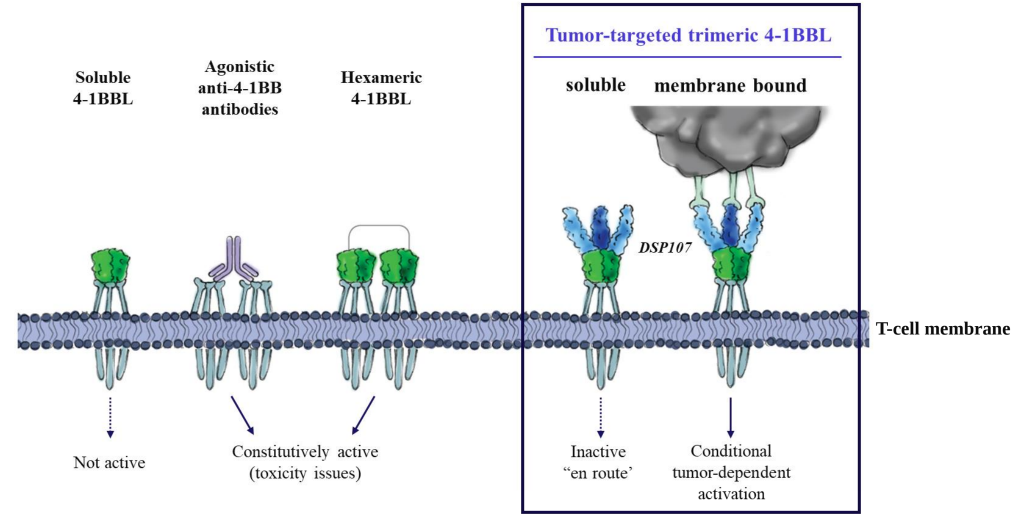
- Cancer selective binding to overexpressed CD47 on tumor cells, minimizing RBC binding and associated toxicities
- Conditional 4-1BB mediated T cell activation dependent on trimeric binding to CD47 on cancer cells

Trimeric 4-1BBL **Cytolytic T cell activation T cell Proliferation Checkpoint inhibition Tumor microenvironment** modulation 3 SIRPα for **CD47** Checkpoint Targeting

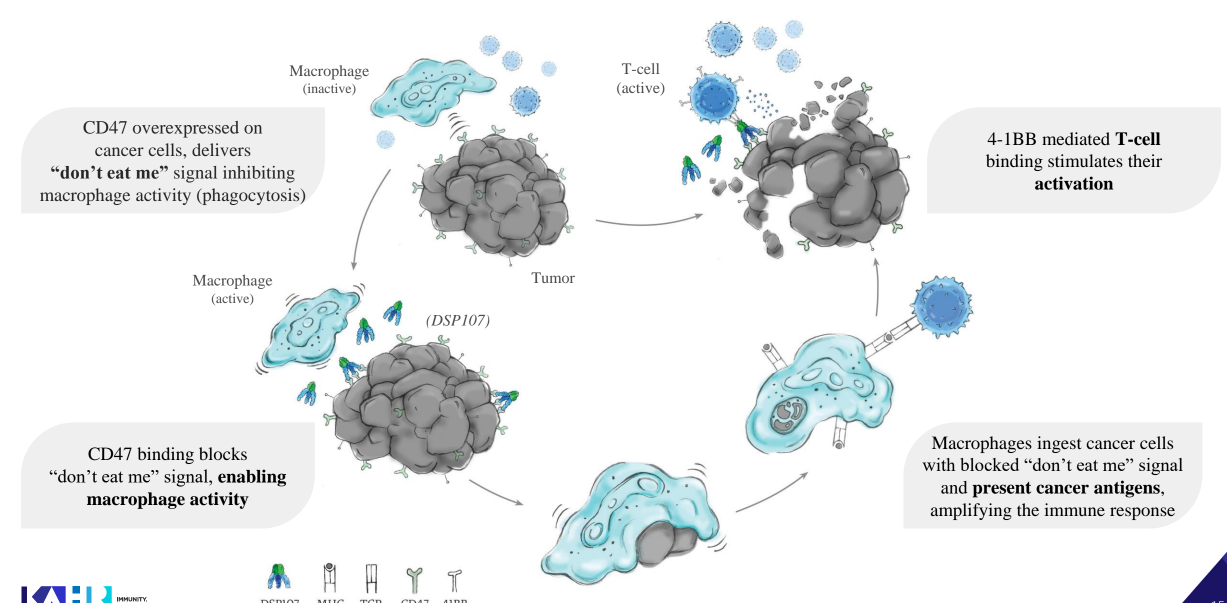


DSP107 Structure Designed to Safely Unlock 4-1BB Potential

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

DSP107 combination with therapeutic Antibodies IgG1 mAbs

(cetuximab, trastuzumab...)

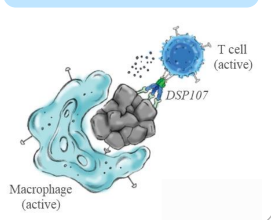
DSP107 combination with PD1/PD-L1 Checkpoint Inhibitors

(atezolizumab, pembrolizumab...)

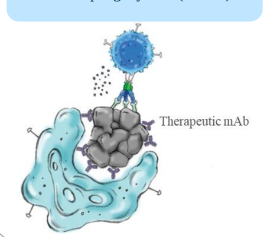
DSP107 combination with pro-apoptotic agents

(chemotherapy, hypomethylating agents and BCL2 inhibitors)

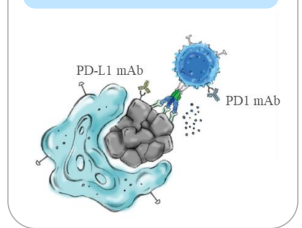
Triggers macrophage mediated phagocytosis and T cell cytotoxicity



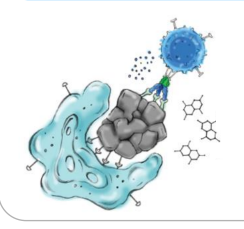
Enhances antibody-dependent cellular phagocytosis (ADCP)



Enhances T cell activation



Increases "eat me" signals







Preclinical Studies Support Differentiated Potential Dual MoA

Dual MOA

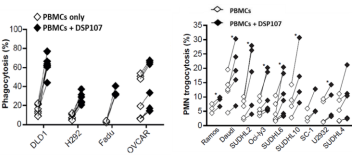
designed to activate innate and adaptive immunity

Favorable safety without hematological or hepatotoxicities in NHP observed

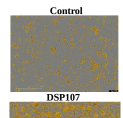
Monotherapy potential for treatment of solid and hematological malignancies

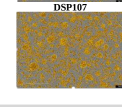
Comprehensive preclinical package demonstrated differentiated features

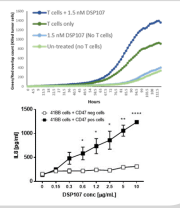




Triggering cancer cell death by phagocytosis as a single agent

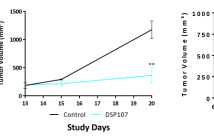


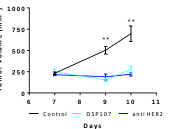




Inducing 4-1BB activation, T-cell proliferation and killing potential against cancer cells

Lymphoma and Ovarian Carcinoma models

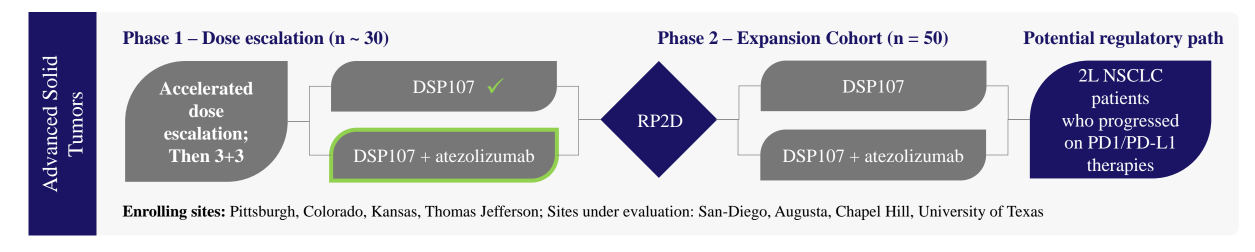




Demonstrating single agent anti tumor activity in mice models



DSP107 Clinical Program

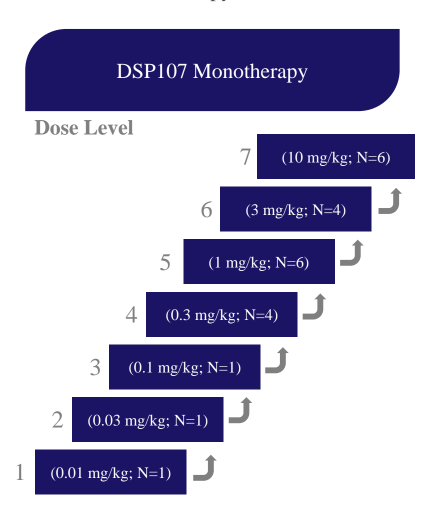


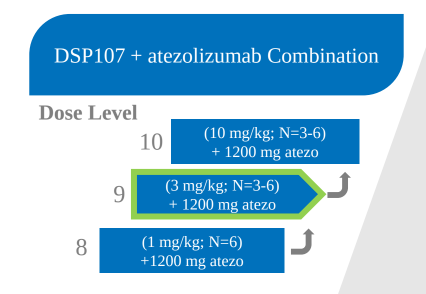
Phase I Dose escalation (n ~ 36) Hematological Malignancies **Phase 2 – Expansion** $\overline{\text{DSP107}} \pm \text{azacytidine}$ High risk R/R Dose Details to be announced MDS/CMML and AML RP2D escalation following EOPI meeting who failed up to 2 prior DSP107 + azacytidine N=6-8/cohort with the FDA therapeutic regimes + venetoclax Lead site: MD Anderson Cancer Center



Trial Design and Key Inclusion Criteria

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





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

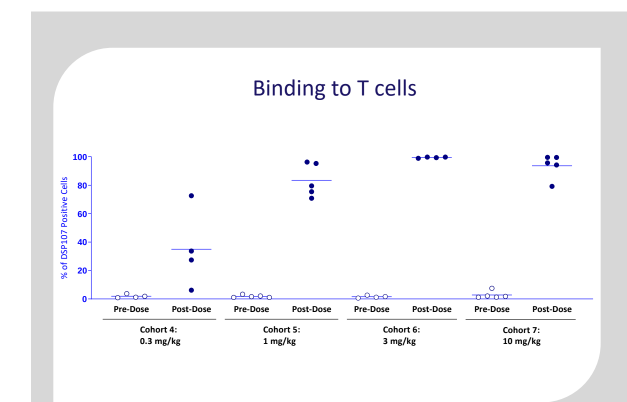
Treatment-Related AEs in ≥ 2 Patients		
Total No of Patients	N = 23 (cohorts 1 - 7)	
Treatment-related AEs (any grade)	n (%)	
Any	16 (70)	
IRR*	7 (30)	
Diarrhea	4 (17)	
Fatigue	4 (17)	
Nausea	3 (13)	
Constipation	2 (9)	

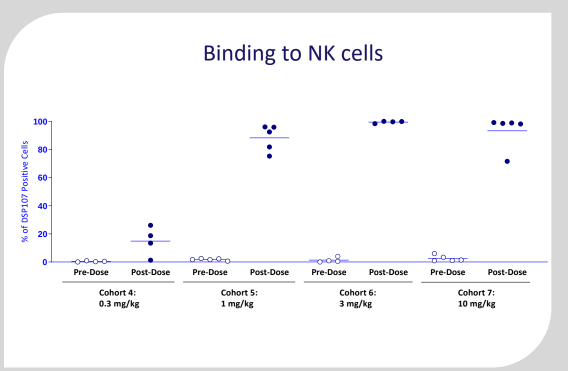
^{*}IRRs Grade 1-2 in severity. Easily abrogated (except 10-004 and 11-012 who withdrew consent) in subsequent infusions by reduced rate of infusion and concomitant IV fluids.



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

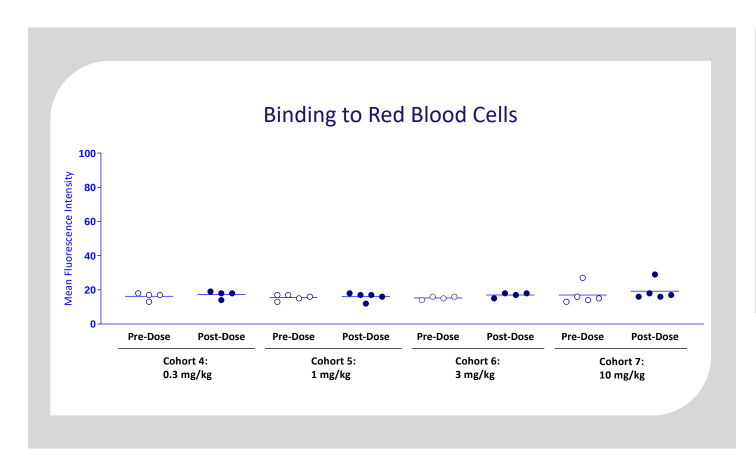


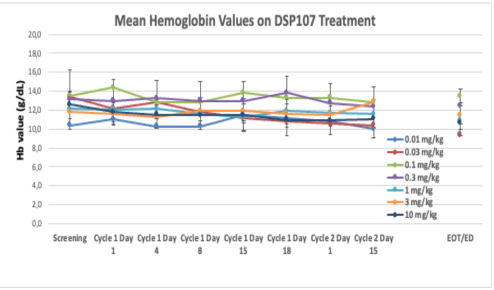


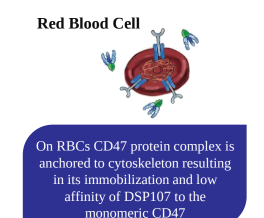


DSP107 Does Not Bind Red Blood Cells

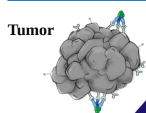
Resulting in Favorable Safety Profile With No Anemia or Antigen Sink Issues







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





DSP107 Treatment Resulted in Tumor Necrosis

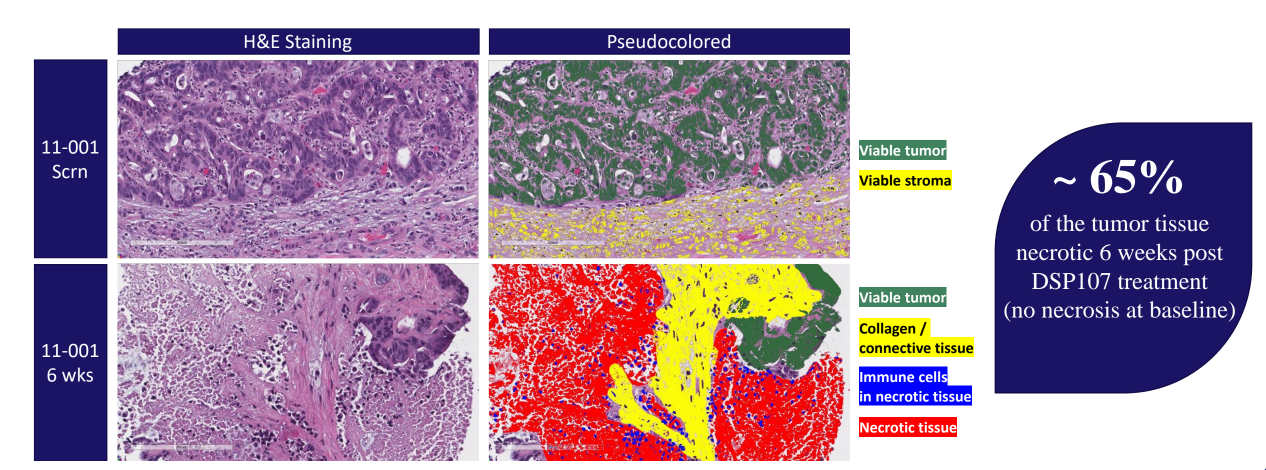
Key paired biopsies data

- 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
11-001			6 weeks	65
11-002	0.3	Colorectal	Screening	2
			6 weeks	35
10.002	1	Pancreatic	Screening	10
10-003			6 weeks	50
13-005	1	Pancreatic	Screening	4
	1		6 weeks	3



Case study: Increased Tumor Necrosis Associated With Immune Cell Infiltration

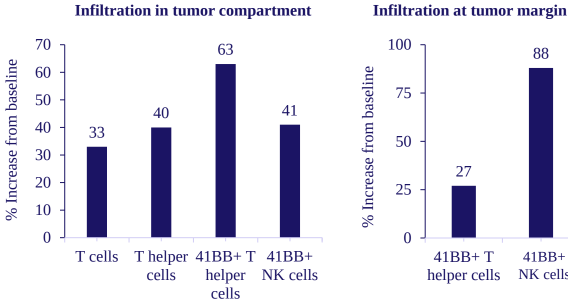


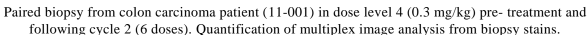
Paired biopsy from colon carcinoma patient (11-001) in dose level 4 (0.3 mg/kg) pre- treatment and following cycle 2 (6 doses)

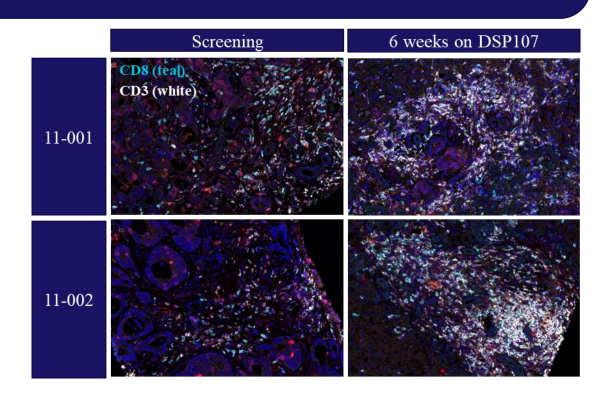


Case Study: Paired Biopsies Demonstrated Adaptive Immune **Engagement in Ongoing Clinical Trial**

6-wks post DSP107 treatment







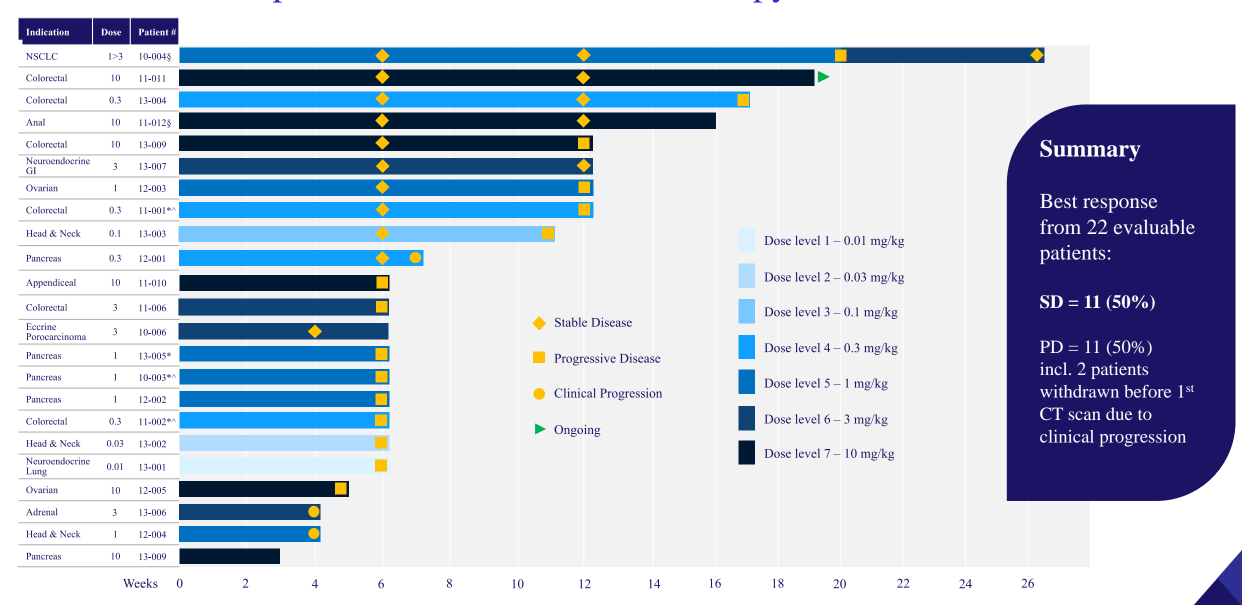
Significant infiltration of T cells and NK cells in both the tumor compartment and at the tumor margin following DSP107 treatment

41BB+

NK cells



Best Overall Response After DSP107 Monotherapy





^{*}Paired biopsy data available; ^Necrosis induction evident in "on treatment" biopsy

[§] Patient withdrawn due to Grade 2 IRRs not controlled with standard supportive measures

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 IFNy secretion and anti-tumor killing
- Increased macrophage phagocytosis of tumor cells
- Augments mAbs' ADCP 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 ± atezolizumab (intended to enhance T cell activation)
- Hematological Malignancies
 - DSP107 ± 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%)



DSP502

MIRP Type **DSP-Fc**

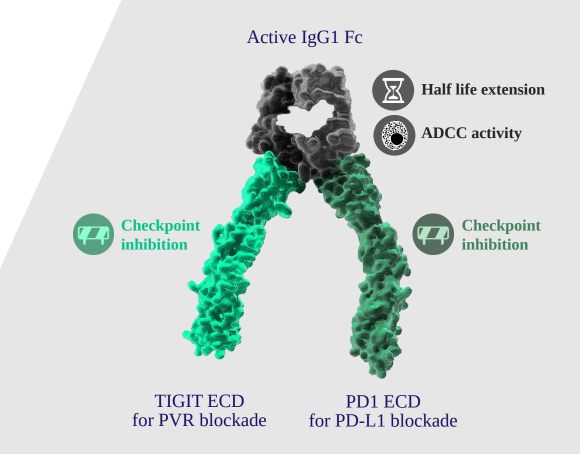
Targets PVR, PD-L1, FcR

Primary Cell Target NK cells, T effector cells

Mechanism of Action
Dual checkpoint inhibition designed to unleash NK and T-cells, ADCC

Dual Checkpoint Binding Aimed at Enhancing Selectivity and Synergistic Immunity

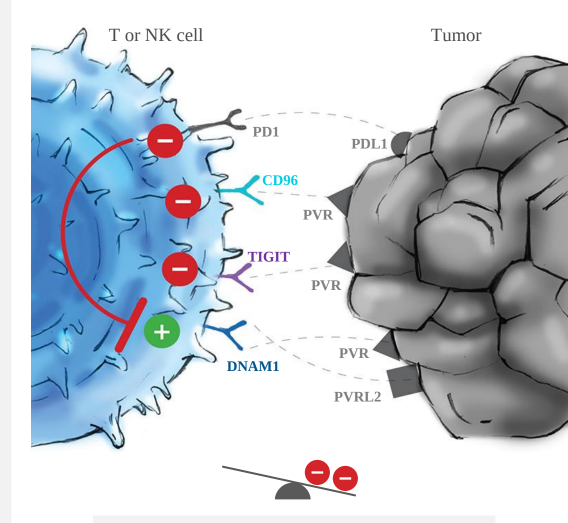
- Dual binding to overexpressed checkpoints may enable high tumor specificity
- Potential anti-tumor immunity via simultaneous checkpoint inhibition of PVR and PD-L1
- Active Fc may extend half-life and enhance ADCC activity





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















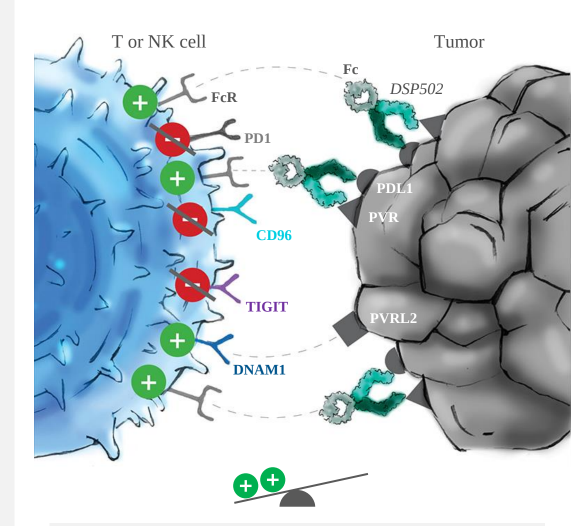
Synergistic Dual Checkpoint Inhibition for Robust Anti-tumor Immunity

- 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)
— TIGIT		✓
C D96		_
+ DNAM1		_
PD-1		_

*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

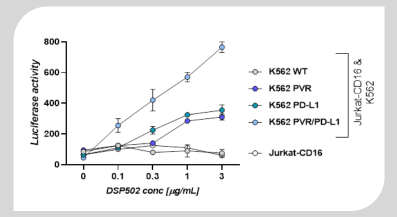
High Tumor Specificity
Concomitant binding to PVR and
PD-L1 required for its activity

Designed to Have Unique FeaturesMultiple functionalities that act
simultaneously for synergistic effect

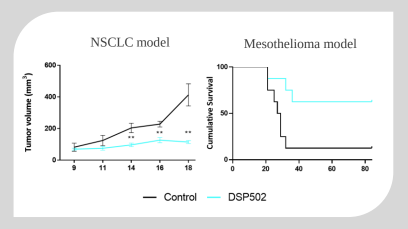
35 30-4414% 30-K562 PVR K562 PVR+DSP502 K562 PVR+DSP502 K562 PVR/PD-L1 K562 PVR/PD-L1 K562 PVR/PD-L1 K562 PVR/PD-L1 K562 PVR/PD-L1

Enhanced NK cells cancer killing potential

Differentiated mechanism of action



Augmented NK cells ADCC activity



Demonstrated potent single agent anti tumor activity

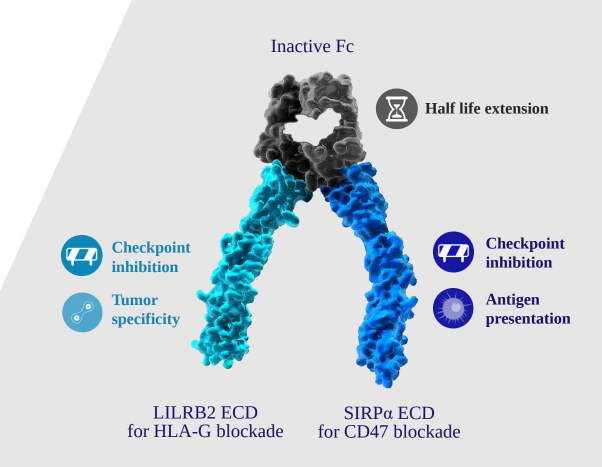


DSP216

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

Dual Checkpoint Binding Designed For Enhanced Selectivity and Broad Immunity

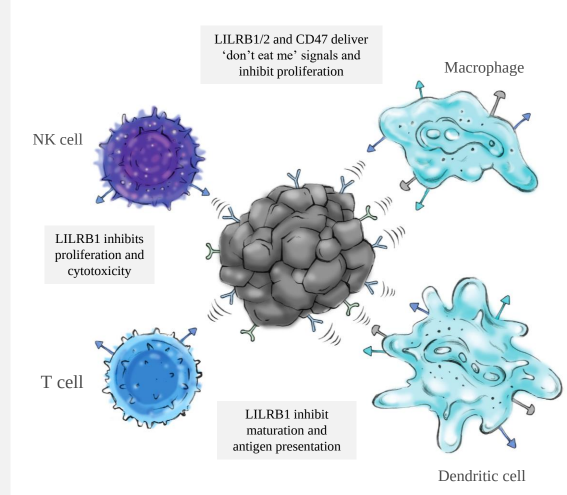
- Dual binding to overexpressed cancer checkpoints may enable high tumor specificity
- 2 HLA-G and CD47 blockade designed to activate innate and adaptive immunity
- Inactive Fc may extend half-life





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





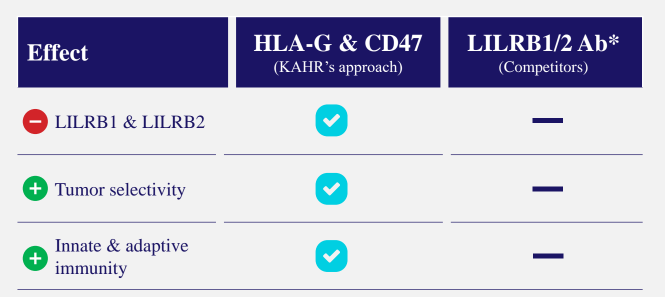


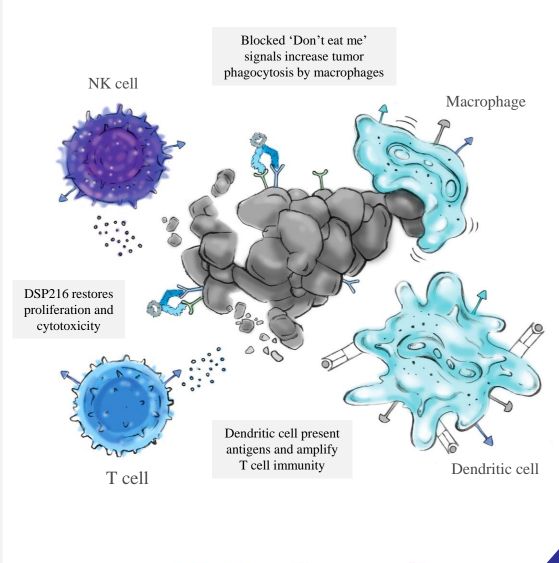




Dual Targeting Designed For Robust Tumor Selective Innate & Adaptive Immune Activation

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



















CORPORATE HIGHLIGHTS

Multifunctional Cancer Immunotherapy Candidates Targeting Innate and Adaptive Immune Cells



NOVEL MIRPsTM

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





IP

13 families4 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

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