

Research & Development Day

Accelerating the next generation of therapeutics to improve the standard of care for the most challenging diseases in cancer and autoimmune disease

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Today's Agenda



08:30 AM	Zymeworks R&D Vision Dr. Paul Moore	10:20 AM	Beyond Solid Tumors Dr. Paul Moore Dr. Alexey Berezhnoy
08:50 AM	Gynecological Cancer Dr. Susana Banerjee	10:50 AM	Next Generation Technologies Stuart Barnscher
09:00 AM	Lung Cancer Dr. Hatim Husain		Dr. Jamie Rich Dr. Nina Weisser Dr. Thomas Spreter Von Kreudenstein
09:10 AM	Gastrointestinal Cancer Dr. Jaffer Ajani	11:20 AM	Closing Remarks Kenneth Galbraith
09:20 AM	Solid Tumor Program Dr. Jeff Smith Dr. Paul Moore	11:30 AM	Q&A Session #2
09:50 AM	Q&A Session #1	12:00 PM	Networking Session



ZYMEWORKS R&D STRATEGY

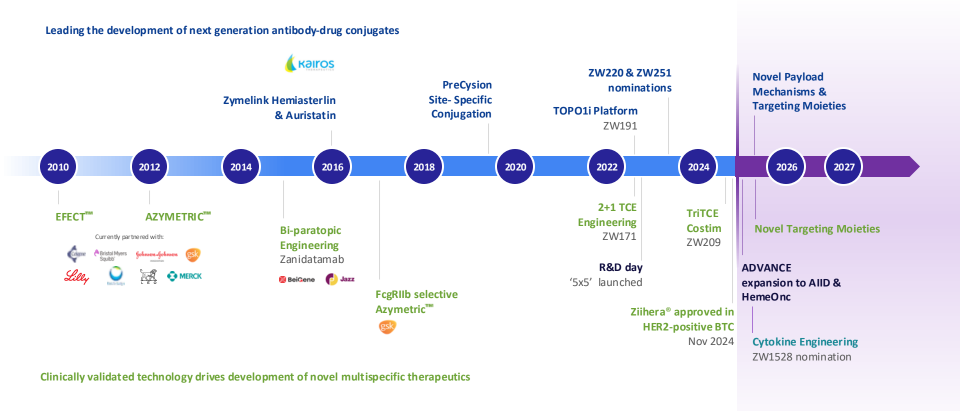
Zymeworks R&D Strategy

Paul Moore, PhD Chief Scientific Officer



10+ Years of Pioneering Multi-Functional Antibody Development





Dedicated to Advancing Targeted Therapies that Address Some of The Most Challenging Diseases, Including Aggressive Cancers and Autoimmune Disorders



Strategic
Partnerships

Collaborating with industry leaders to accelerate impact

Extended the reach of therapeutic candidates, while **validating our innovative approach** through strategic partnerships with companies including Jazz, BeiGene, J&J, and others.

Internally Developed Approved Drug

Ziihera® (zanidatamab-hrii)

(HER2 bispecific antibody)

Licensed to Jazz and BeiGene

2L BTC (IHC3+) U.S. FDA Approval

1L BTC confirmatory trial ongoing

Phase 3 GEA top-line PFS readout estimated 2Q25

Wholly-Owned Candidates

Multiple Modalities and Therapeutic Areas

2 Clinical Stage Assets in Phase 1 Trials: ZW171 & ZW191

2 INDs Planned in 2025: ZW220 & ZW251

2 INDs Planned in 2026: ZW209 & ZW1528

Azymetric™ – Adaptable to Different Formats and Applications



Engineering

Set of transferable mutations supporting pure and stable Fc heterodimer formation with exclusive chain pairing during co-expression

Libraries of constant domain Fab mutations available for kappa/kappa, kappa/lamda and lambda/lambda bispecific LC combinations

Flexibility

Can employ novel or existing antibody paratopes; human (IgG1, IgG2A, IgG4) and mouse frameworks; other CH2 and glyco-engineering approaches (eg YTE). Compatible with linker/payload conjugation

High-throughput Screening

Best-in-class activity requires screening of alternative targets, epitopes, sequences, target engagement geometries, and mechanisms of action (blocking, lytic, ADC)

Highly Manufacturable

Antibody like yields/stability; leveraged by multiple pharma/biotech with various clinical stage programs in development

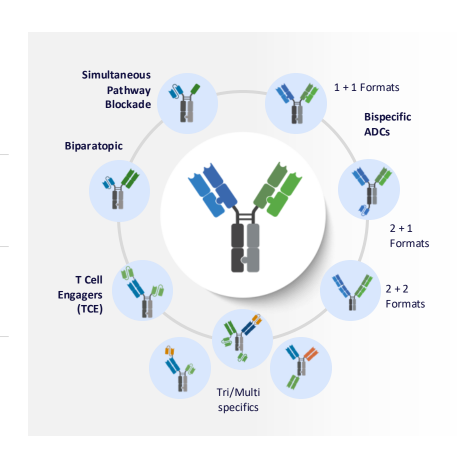








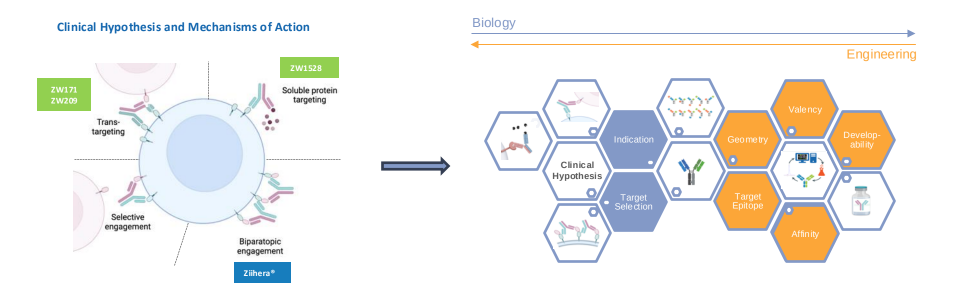




Multispecific Antibody Development Requires Optimization of Multiple Parameters Specific to Desired MOA



Understanding the interplay of antibody geometry with optimal paratope affinity, valency, and target epitope is critical to identifying best-in-class multispecific antibody therapeutics



Zymeworks' Multispecific Engineering Approach – Key Expertise in Format and Geometry Screening to Identify Differentiated Activity



ADVANCE

- Potential best-in-class activity requires screening of epitopes, affinities and target engagement geometries
- Unique flexibility of Azymetric[™]
 enables format and affinity
 optimization for potential best-inclass attributes
- Discovery of unique biology and differentiation to combination approaches

Biparatopic

Zanidatamab

- Optimization of affinity and format for highest biparatopic activity
- Unique biparatopic MOA
- · Superior activity to combination



2+1 TCE

ZW171 (2+1 MSLN TCE)

- Avidity optimization to prevent normal tissues tox
- Avidity and format optimization to not bind shed MSLN
- Synapse optimization for high activity with minimal cytokine release



Multi-Cytokine Blocker

ZW1528 (IL4Rα -IL33)

- IgG-like format, manufacturability and PK
- IL4Ra and IL33 blockade equivalent benchmarks
- Unique bispecific activity, potentially superior to combination

Trispecific T Cell Engager

ZW209 (CD28 TriTCE)

- Discovery of novel format to prevent non-specific T cell activation
- · Conditional CD28 activation
- Synapse optimization for balanced Signal 1 plus
 Signal 2



Increased Complexity

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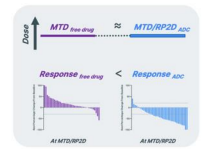
Historical Observations Guide Our Approach to ADC Design

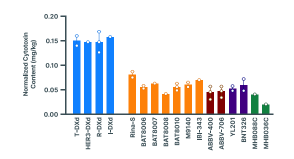


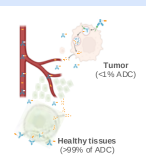
Conjugation does not improve payload MTD¹

More potent payloads limit protein dose²

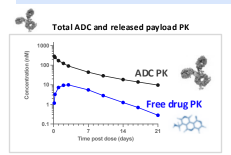
Most ADC catabolism occurs in normal tissue³

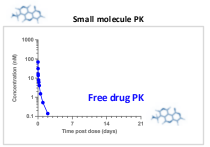




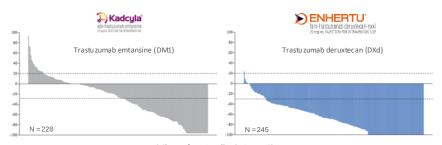


4 Bystander activity is a key payload feature²





The right payload mechanism matters⁴



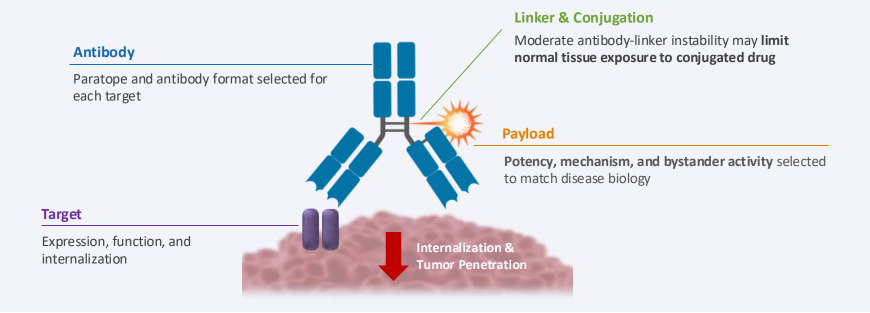
% Change from Baseline in Tumor Size

^{1.} R. Colombo and J.R. Rich, Cancer Cell 2022, 40(11):1255-1263; 2. R. Colombo et al, Cancer Discov 2024, 14(11):2089-2108; 3. A.T. Lucas et al, Antibodies 2019, 8(1):3; 4. J. Cortés et al, N Engl J Med 2022, 386:1143-1154.

Building Clinically and Commercially Relevant ADCs Requires Careful Selection of Target, Antibody, Linker/Conjugation, and Payload



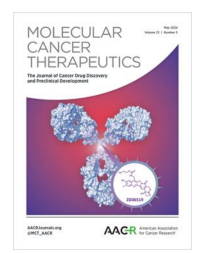
Historical Observations Guide Our Approach to ADC Design

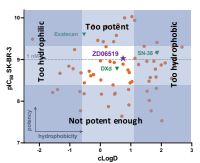


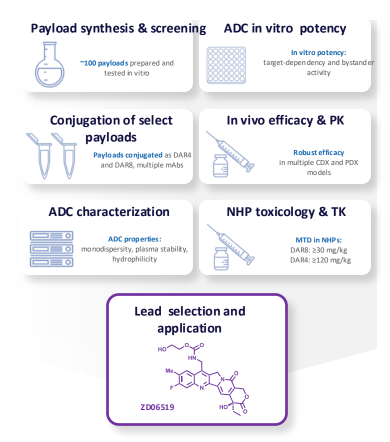
1. Colombo R, Rich JR. Cancer Cell 2022 (40), 1255-1263; 2. Colombo R, Barnscher SD, Rich, JR. Cancer Res 2023, 83 (7). Abstract #1538 presented at AACR 2023.

Zymeworks Topoisomerase ADC Platform Exemplifies Our Philosophy and Enables Our Pipeline

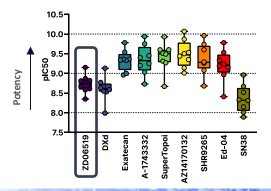








- Moderate potency to enable higher ADC dose
- Bystander active
- ZW191 first in human trial (NCT0655574)
- ZW220 and ZW251 expected to enter clinic in 2025



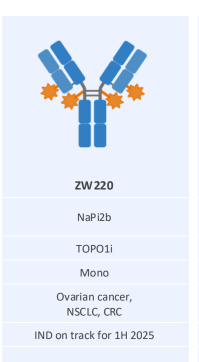
M.E. Petersen, M.G. Brantetal, Mol Cancer Ther 2024, 23(5):606-618

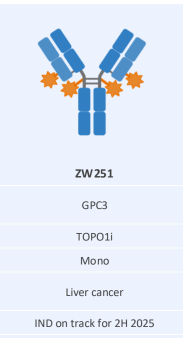
Zymeworks' TOPO1i ADC Assets Have Potential to Address Multiple Diverse **Patient Populations**

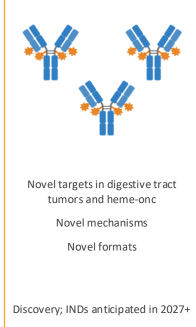


ADVANCE

ZW 191 FRα TOPO1i Mono Ovarian cancer, endometrial cancer, and NSCLC Phase 1







Making a Meaningful Difference

Target

Antibody

Stage

Payload mechanism

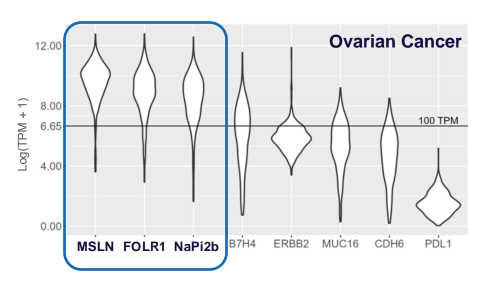
Potential Indications

Target Selection Driven by Expression Profile, Biology and Clinical Precedence



Selection of an ADC or TCE strategy is driven by target expression, biology including internalization rate, clinical precedence and differentiation to prior therapeutic programs

mRNA Expression Profile of Select Cancer Target in Ovarian Cancer (N=421)



A balanced portfolio of ADCs targeting clinically validated FRα and NaPi2b, along with a T cell engager targeting MSLN, ensures comprehensive coverage and risk mitigation for ovarian cancer and NSCLC, providing a diversified therapeutic focus on ovarian and lung cancers.

MSLN, FOLR1 and NaPi2b are each expressed at higher level than other targets pursued in ovarian cancer or NSCLC

TCGA bulk RNA-sequencing data were obtained from TCGA-OV, workflow ST AR – Counts from https://portal.gdc.cancer.gov/repository. The median TPM (Transcript per Million) for each gene in each patient was plotted on a violin plot using ggplot 21. This dataset contains 42.1 samples (patients) from Ovarian Serous Cysta denocarinoma (OV) and 5.21 sample from lung ade no carcinoma. The width of the shape/violins indicates the density of samples

Development Pipeline



Program	Technology	Target	Indication	Discovery	Preclinical	Phase 1	Phase 2
Solid Tumor Oncology: Antibody-D	Solid Tumor Oncology: Antibody-Drug Conjugates (ADC)						
ZW191 Topo1iADC DAR 8 Fc WT	ZD06519 Payload	Frα	Gynecological Thoracic	NCT06555744			
ZW220 Topo1iADC DAR 4 Fc Mut	ZD06519 Payload	NaPi2b	Gynecological Thoracic			IND 1H 2025	
ZW251 Topo1iADC DAR 4 Fc WT	ZD06519 Payload	GPC3	Digestive System (HCC, PDAC)			IND 2H 2025	
Solid Tumor Oncology: T Cell Enga	Solid Tumor Oncology: T Cell Engagers (TCE)						
ZW171 Trivalent TCE 2+1 Format	Azymetric™ Novel anti-CD3	MSLN x CD3	Gynecological Thoracic	NCT06523803			
ZW209 Trispecific TCE Tri-TCE Costim	Azymetric [™] Novel anti-CD3 Conditional CD28	DLL3xCD3xCD28	Thoracic		IND	1H 2026	
Autoimmune and Inflammatory Disease							
ZW1528 Dual Cytokine Blocker	Azymetric™ Hetero-Fab YTE	IL4RaxIL33			IND	2H 2026	

Fc WT: Fast continuous wavelet transform; Mut: Methylmalonyl-CoA mutase; HCC: hepatoœllular carcinoma; PDAC: pancreatic ductal adenocarcinoma



SOLID TUMOR PROGRAM

ZW171 and ZW191

Jeff Smith, MD, FRCP Executive Vice President and Chief Medical Officer



Zymeworks' Pipeline has an Opportunity to Address Unmet Needs Across Indications



	Ovarian Cancer	Endometrial Cancer	NSCLC	SCLC	Pancreatic Cancer	Hepatocellular Carcinoma
ZW171 MSLN x CD3 TCE	<u>~</u>	~	~		✓	
ZW191 FRα TOPO1i ADC	~	~	~			
ZW220 NaPi2b TOPI1i ADC	<u>~</u>	~	~			
ZW251 GPC3 TOPO1i ADC						<u>~</u>
ZW209 DLL3xCD3xCD28 TCE				<u>~</u>		

Zymeworks' Therapies Actively Enrolling into Phase 1 Clinical Studies



ZW171



MSLN x CD3 bispecific T cell engager

- 2+1 format
- Designed for enhanced therapeutic window
- Tumor selective binding and cytotoxicity
- Maintains potency in presence of soluble MSLN
- Reduced T cell binding to mitigate risk of CRS

ZW191



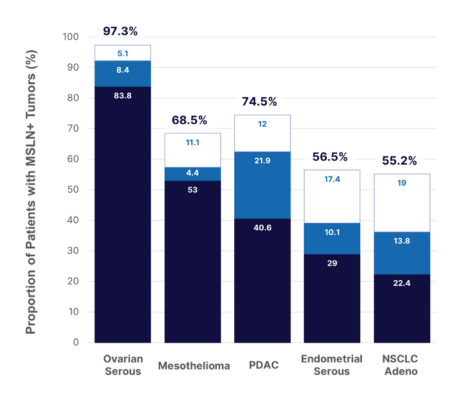
FRα-targeting ADC

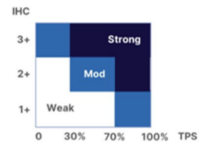
- Antibody selected to improve upon current in-clinic models
- Proprietary TOPO1i payload selected for gynecological cancer tissue response
- DAR 8 selected for protein dose vs protein expression

ADC: artibody-drug conjugate; CD3: cluster of differentiation 3 protein complex and T cell co-receptor; CRS: cytokine release syndrome; DAR: drug-to-artibody ratio; FRa: folate receptor alpha; MSLN: mesighelin; TOPOfic hord-somerase-1 inhibitor.

ZW171: Mesothelin Expression Is Frequent in Ovarian Cancer, Endometrial Cancer, NSCLC, PDAC and Other Malignancies





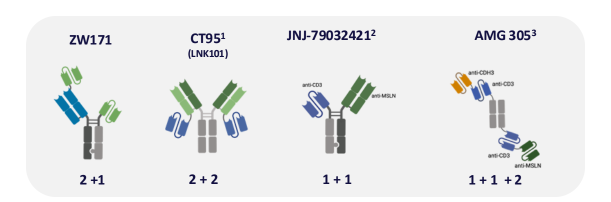


Adapted from Weidemann S, et al Biomedicines. 2021;9(4):397. CRC: cdorectal cancer; FRa: foldate receptor alpha; GEA: gastroes ophageal adenocarcinoma; IHC: immunohistochemistry; Mod: moderate; MSLN: mesothelin; NSCLC: non-small cell lung cancer; PDAC: pancreatic ductal adenocarcinoma

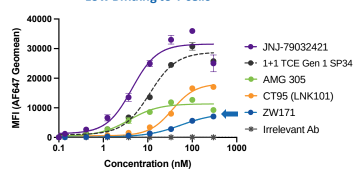
ZW171 Exhibits a Wider Therapeutic Window Compared to Next Gen MSLN TCEs



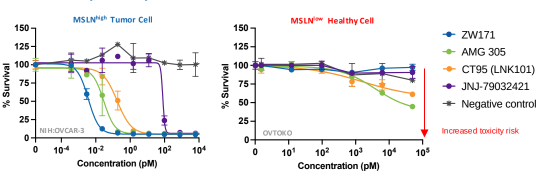
- Enhanced tumor selective cytotoxicity
- No targeting of normal tissues
- Low affinity CD3 binding to mitigate peripheral T cell binding and cytokine release
- Maintains potency in the presence of soluble MSLN



Low Binding to T cells



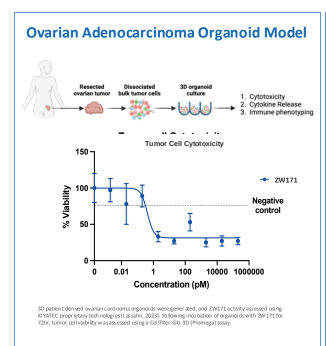
Potent Cytotoxicity in MSLN⁺ Tumor Cells but not Normal Cells

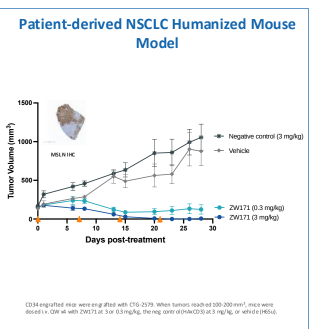


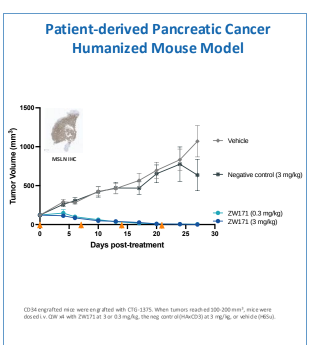
Tumor cell lines were occultured with human PBMCsatan ET ratio of 5:1. Test articles were thrated and added to week in duplicate. After 72hr, tumor cell survival wasassessed by high-content imaging, longative control. HACCO3.

ZW171 Mediates Strong Anti-Tumor Activity in Patient-derived *Ex Vivo* and *In Vivo* Models







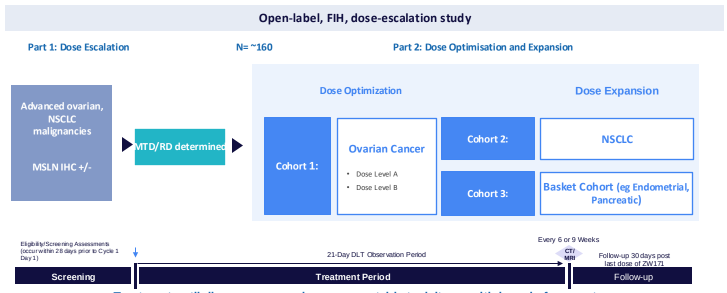


Lassahn et al., Abstract 2275: Predinical testing of therapeutic biologics using patient-derived 3D spheroids. Cancer Res 1 April 2023; 83 (7_Supplement): 2275 IHC: immunohistochemistry. MSLN: mesothelin: NSCLC: non-small cell lung cancer

ZW171 Global Phase 1 Study in MSLN-Expressing Solid Tumors



(NCT06523803)

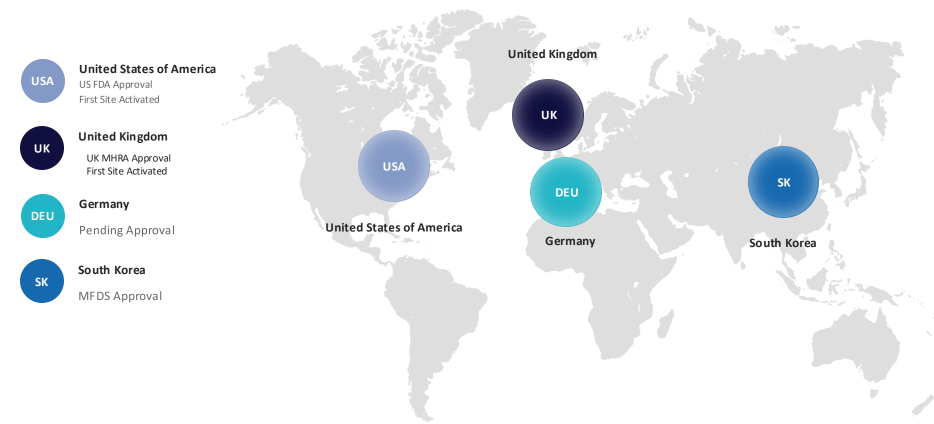


Treatment until disease progression, unacceptable toxicity, or withdrawal of consent

ADA: anti-drug artibodies; of RR: confirmed objective response rate; DL: dose level; DOR: duration of response; ECOG: Eastern Cooperative Oncology Group; FIH: first in human; GEA: gastrointestinal adenocarcinomas; HC: immunohistochemistry; MTD: maximum blerated dose; MSLN: mesothelin; mTDI: modified boxidity probability interval; NSCLC: non-small cell lung cancer; OBD: optimal biological dose; OS: overall survival; PFS: progression-free survival; PFC: pharmacokinetics; pts: patients; PS: preferred status; QSW: every 3 weeks; RD: recommended dose; SOC: standard of care; TBC: to be confirmed

ZW171 Clinical Development Progress – Dose Escalation Territories



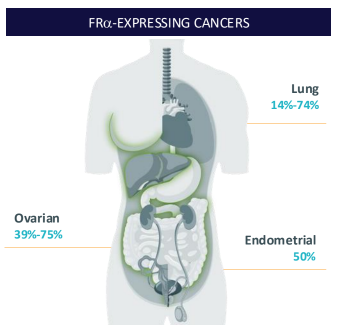


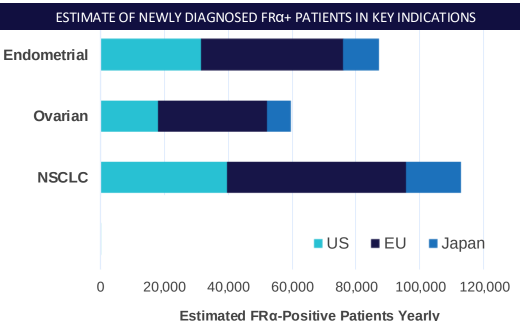
US FDA:U.S. Food and Drug Administration; MFDS: Ministry of Food and Drug Safety.

FRα-expressing Cancers Represent a Significant Commercial Opportunity¹⁻⁷



- Potential first and best-in-class in FRα-high endometrial, NSCLC, TNBC, and FRα-mid/low solid tumors
- Potential best-in-class opportunity in FRα-high ovarian cancer





Estimated FRα-Positive Patients Yearly

FRC: folte receptor alpha; NSCLC: non-small cell lung career; TNBC: triple regative breast cancer

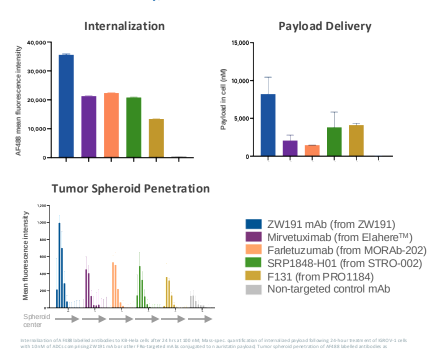
1. Send S, et al. Int J Clin Exp Pathol. 2015;8(5):5633-5641; 2. Omote S, et al. Med Mol Morphol. 2018;51(4):237-243; 3. Oza AM. SGO. 2021; 4. O'Shannessy DJ, et al. Oncotarget. 2012;3(4):414-425; 5. Nunez MJ, et al.

1 Thorac Oncol. 2012;7(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Mod Pathol. 2011;24(9):1221-1228; 7. Scaranti MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6. D'Angelica MJ, et al. Nat Rev Clin Oncol. 2012;17(5):833-840; 6.

ZW191: Key Design Considerations



ZW191's Novel mAb Drives Superior Internalization, Payload Delivery, and Tissue Penetration

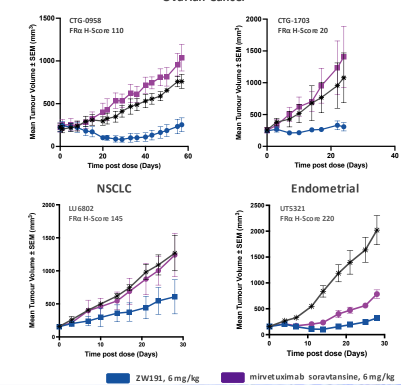


Wong Jetal, Abstract #3127 presented at American Association for Cancer Research annual meeting 2024 Lawn S. et al. Abstract #1862 presented at American Association for Cancer Research annual meeting 2024

quantified by high content imaging of spheroid layers at 24 hours post-treatment at 50 nM.

Anti-tumor Activity Across Multiple Tumor Types And Range of FRα Expression (PDX models)

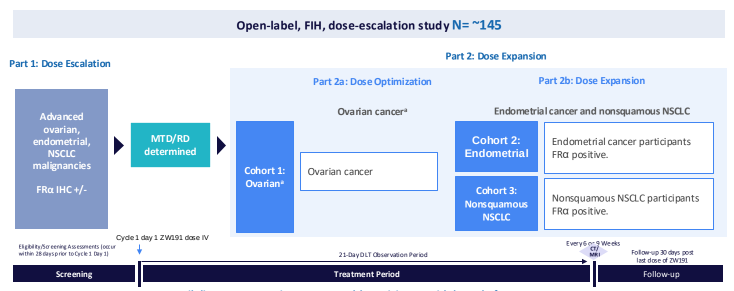




ZW191: Global Phase 1 Study in FRα-Expressing Solid Tumors



(NCT06555744)



Treatment until disease progression, unacceptable toxicity, or withdrawal of consent

*Ovarian cancer includes primary peritoneal and fallopian tube cancers. *Part 2 will be initiated at dose levels (RDEs) based on the SMC's comprehensive analysis of safety, tolerability, clinical PK, PD, and preliminary antitumor activity data from Part 1. The Part 2 selected doses will be decided at SMC meetings and could be the MTD or RDEs based on comprehensive analysis of safety, tolerability, clinical PK, PD, and antitumor activity data from each part 1. The RDE dose levels may vary across the tumor types in Cohorts 1, 2, and 3. *Timed from cycle 1 day 1. Q6W (every 6 weeks) for the first 4 assessments and then Q9W (every 9 weeks) thereafter. Clinical Trials, gov ID: NCT06555744.

ADA: anti-drug anti-drug anti-body; ADC: antibody-drug conjugate; AE: adverse event; AESI: adverse event of special interest; CNS: central nervous system; CT/MRI: computed tomography/magnetic resonance imaging; ECOG PS: Eastern Cooperative Oncology Group performance status; FRa: folate receptor alpha; IHC: immunohistochemistry; IV: intravenous; MTD: maximum tolerated dose; NSCLC: non-small cell lung cancer; ORR: objective response rate; PD: pharmacodynamics; PK: pharmacokinetics; Q3W: every 3 weeks; RDE: recommended dose for expansion; RECIST: Response Evaluation Criteria in Solid Tumors; SAE: serious adverse event; SMC: safety monitoring committee; SOC: standard of care; TBD: to be determined: TOPO1i: toposomerase-1 inhibitor.

ZW191 Clinical Development Progress

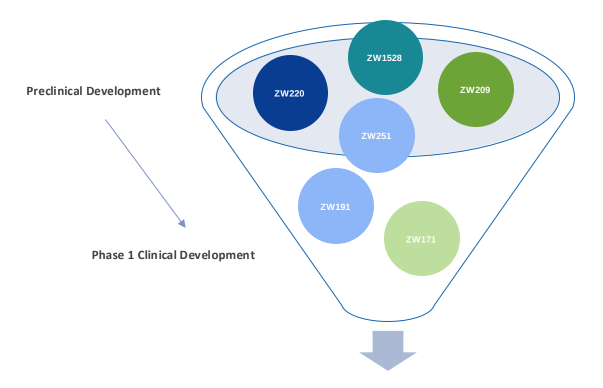




USFDA: U.S. Food and Drug Administration; PDMA: Prescription Drug Marketing Act; TGA: Therapeutic Goods Administration; MFDS: Ministry of Food and Drug Safety; HAS: Health Sciences Authority

Multiple Candidates in Development Offer Strategic Pivot Points





Decision Pathway Factors

- Optimal dose
- Tolerability and safety
- Early signs of anti-tumor activity
- Proof of targeted therapy: strong relationship between expression levels and anti-tumor activity
- Competitive landscape

Pipeline Resource Allocation

Partnership Optionality

Combination Approaches

Accelerated Development into Phase 2/3



SOLID TUMOR PROGRAM

ZW220 and ZW251

Paul Moore, PhD Chief Scientific Officer



Zymeworks' Therapies on Track for 2025 IND Submission



ZW220



NaPi2b-targeting ADC

- TOPO1i payload selected for gynecological and lung cancer tissue response
- DAR4 selected to balance desired antitumor activity with potential for on-target toxicities
- FcγR silenced to potentially minimize toxicities driven by cellular uptake via FcγR

ZW251

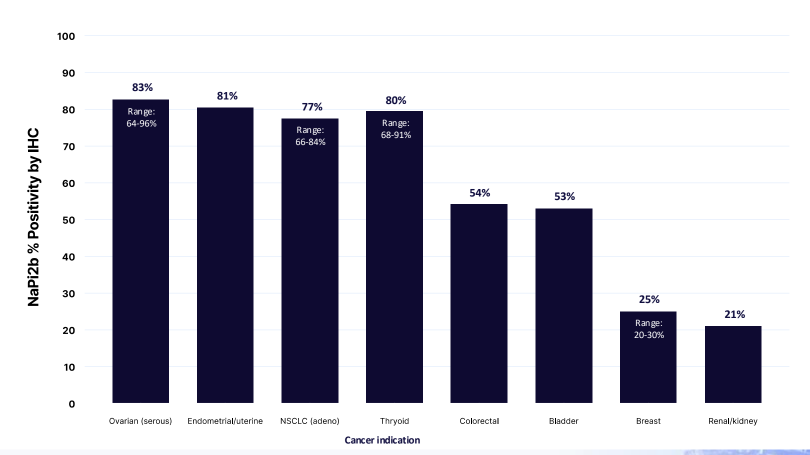


GPC3-targeting ADC

- Designed to target GPC3 that is overexpressed in majority of Hepatocellular carcinoma patients
- TOPO1i payload with bystander activity selected for gastrointestinal cancer tissue response
- DAR4 selected for protein dose vs protein expression

NaPi2b is Overexpressed in Multiple Cancers with High Unmet Medical Need

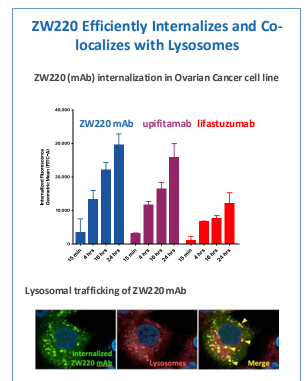


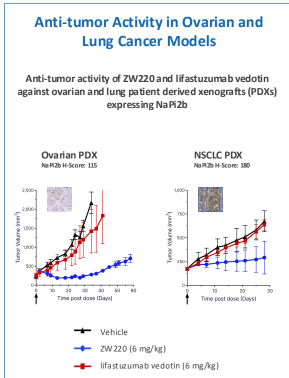


- 1) Baner jee et al. 2023. ESM O #145
- 2) Ri chard son et al. 2022. SGO #76
- 3) Levan et al. 2017. BMC Cancer
- 4) Lin et al. 2015. Clin Cancer Res
- 5) Lopes dos Santo set al. 2013. PLoS One Endo metrial/u terine
- 1) Horsley et al. 2024. Cancer Res #5085
- 1) Horsley et al. 2024. Cancer Res #5085
- 2) Heyn emann et al. 2022. Clin Lung Cancer
- 3) Yu et al. 2018. IASLC #12636 4) Zhang et al. 2017. Tu mo r Bi ology
- 5) Lin et al. 2015. Clin Cancer Res
- 1) Hakim et al. 2021. An al Cell Pathol
- 2) Lin et al. 2015. Clin Cancer Res Colorectal
- 1) Liu et al. 2018. Biomed Pharmacother
- 1) Ye et al. 2017. Cell Death Dis
- 1) Lopes dos Santo set al. 2013. PLoS One 2) Kiyamoya et al. 2011. Exp On col
- Ren al /kidney
- 1) Lopes dos Santo set al. 2013. PLoS One

ZW220: Potential Utility in Multiple Cancers; on Track for Clinical Studies in 1H 2025^{1,2,3}



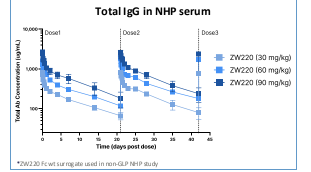






Non-GLP toxicology study in non-human primates dosed 3 times every 3 weeks

Dose	MTD	T _{1/2} (day)		
30 mg/kg		10.3		
60 mg/kg	≥ 90 mg/kg	9.8		
90 mg/kg		8.0		



mAb: monoclonal antibody; PDX: patient derived xenograft; MTD: maximum tolerated dose; T1/2: half-life; GLP: good labor atory practice

1. Hemandez Rojas A et al., Abstract #1533 presented at AACR 2023; 2. Hemandez Rojas A et al. Presentation at World ADC 2023; 3. Hemandez Rojas A et al. Eur. J. Cancer (2024), 211, 114555.

ZW251: Potential Utility in Hepatocellular Carcinoma^{1,2,3}; on Track for Clinical Studies in 2025



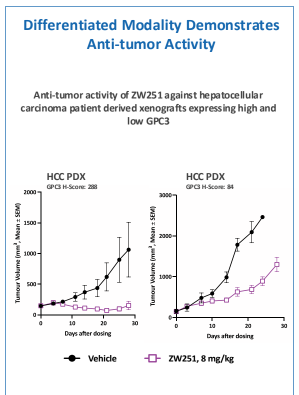
Robust ADC Internalization and Cytotoxicity ZW251 internalized in HCC cell line Internalization visualized after 24-hour treatment Tumor spheroid cytotoxicity in HCC cell line

0.0001 0.001 0.01

0.1

Concentration (nM)

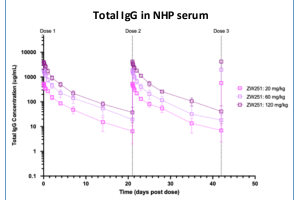
Cytotoxicity assessed by cell line spheroids (treatment over 4 days)



Impressive Tolerability and Doseproportional PK in NHP

Non-GLP toxicology study in non-human primates dosed 3 times every 3 weeks

Dose	MTD	T _{1/2} (day)		
20 mg/kg		4.6		
60 mg/kg	≥ 120 mg/kg	4.8		
120 mg/kg		5.4		



HCC: Hepatocellular carcinoma; PDX: patient derived xe no graft; MTD: maximum tolerated dose; T1/2: half-life; GLP: good laboratory practice

^{1.} Madera L et al., Abstract#2658 presented at AACR 2023; 2. Madera L et al., presentation at World ADC 2023; 3. Madera L et al., Abstract#177 presented at EORT C-NCI-AACR 2024



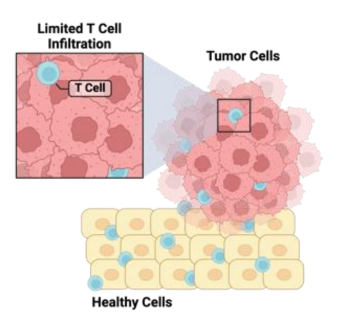
SOLID TUMOR PROGRAM

Product Candidate Nomination: ZW209
DLL3 x CD3 x CD28 TriTCE

Overcoming Lack of Efficacy and Durability of T cell Mediated Responses in Solid Tumors Through Incorporation of CD28 Co-stimulation



Low T cell infiltration and T cell anergy remain challenges in the treatment of solid tumors



Expected Benefit

Opportunity

 Enhances proliferation and survival of CD8 and CD4 T cells¹

CD28

- Ability to expand and maintain Tpex and prevent Tex²
- CD28 signaling critical for Teff expansion and epitope spreading^{3,4}

- No signaling independent of TCR
- Opportunity to engineer safe and effective conditional co-stimulation

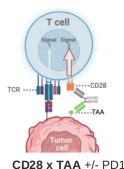
Arvedson T et al Ann Rev Cancer Biol 2022

¹ Lot ze et al., Nature Reviews Immunol 2024; ² Humblin et al., 2023, Sci. Immunol.; ³ Prokhnevska et al., Immunit y 2023; ⁴ Friedrich et al., Camcer Cell 2023

CD28 Co-stimulatory T Cell Engager Approaches

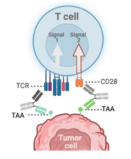


Bispecific CD28 T cell Engagers



Limitations:

Initial clinical activity for CD28-TAA +PD1, but potential toxicity due to autoreactive T cells¹

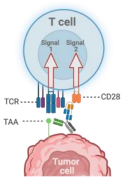


CD28 x TAA + CD3 x TAA

Limitations:

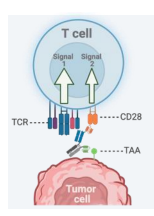
- Optimized for single agent activity and strong CD28 agonism, potential for similar toxicity to CD28-TAA and difficult to optimize by dose adjustment
- Exposure of two molecules at required dose levels potentially suboptimal

Trispecific CD28 T cell Engagers



First Generation:

- High affinity CD3 and CD28 superagonist paratopes^{2,3}
- T cell binding, activation and TMDD observed in periphery^{2,3}
- Target-independent activity and T cell activation



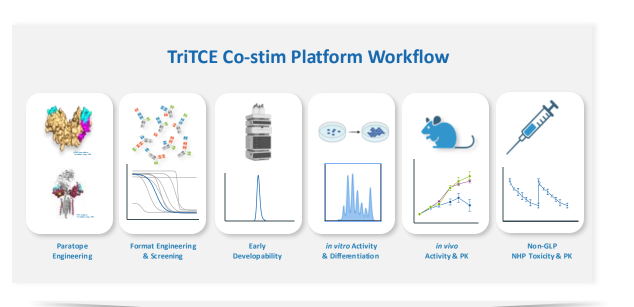
Zymeworks' Next Generation Solution:

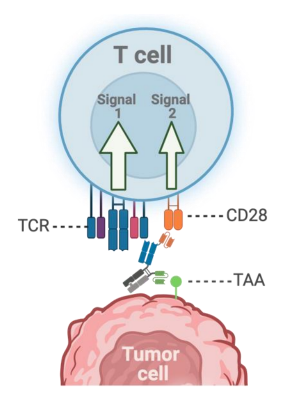
- Balanced low affinity CD3 and CD28 engagement
- Conditional CD28 binding that only binds in cis with CD3 engagement
- Strict target-dependent activity and T cell activation
- Identified via Azymetric[™] screening of various antibody geometries and CD3 and CD28 paratope affinities

Estein et al., Journal Clinical Oncology (2023); 2 Seung et al., Nature (2022); 3 Promsote et al., Nature Communications (2023)

TriTCE Co-stim: A Next Generation Trispecific T Cell Engager Platform





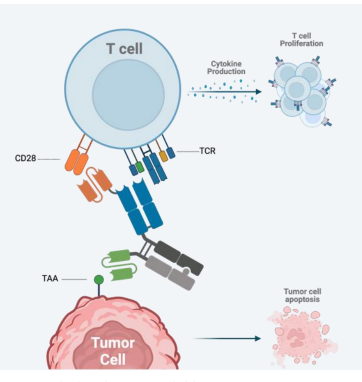


TriTCE Co-stim Lead Format Selection

TriTCE Co-stim Designed to Optimize T cell Binding, Activation and Anti-Tumor Activity



Conditional CD28 Co-stimulation and Obligate cis T cell Binding



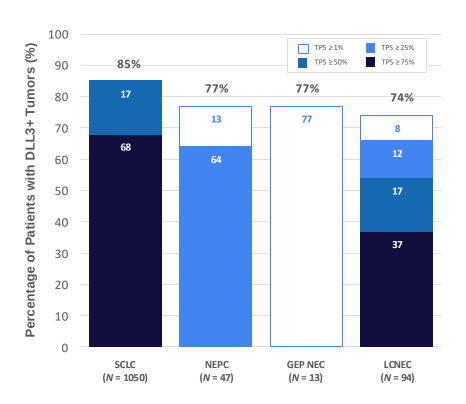
	Design Feature	Expected Benefit
1	Balanced activation of CD3 and CD28	Potential to provide more durable responses and activate T cell responses in 'cold' tumors with lower T cell infiltration
2	Low affinity CD3 and CD28 binding	Prevents overactivation of T cells and reduces risk of CRS and irAEs
3	Obligate <i>cis</i> T cell binding	No T cell-to-T cell bridging or T cell fratricide
4	Conditional CD28 engagement	Requires co-engagement of CD3
5	Enhanced target- dependent activity	Low T cell binding and no T cell activation in periphery or absence of tumor target

CRS: Cytokine release syndrome; ir AEs: immune-related adverse events

DLL3 is an Ideal Target to Evaluate TriTCE Co-stim Platform, with Opportunities in Multiple Cancers



- Responsiveness of DLL3-expressing tumors to TCE modality validated with Imdelltra[™] and other DLL3 bispecific TCEs; however, opportunity for improved responses remains
- DLL3 is expressed on the surface of SCLC and other neuroendocrine tumors but rarely on the surface of normal cells
- Clean expression profile and absence of on-target, off-tumor side effects observed for DLL3 x CD3 bispecifics provides ideal TriTCE Co-Stim target profile

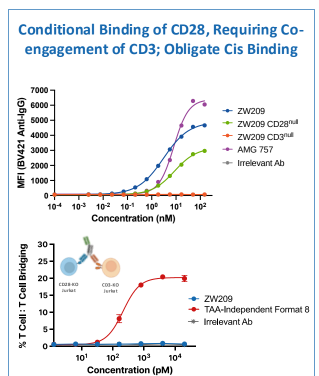


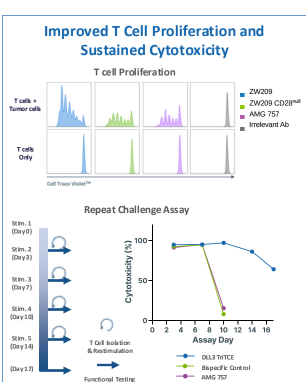
Adapted from: Rojo Fetal Lung Cancer 2020. International real-world study of DLI3 expression in patients with small cell lungcancer. Puca Letal Delta-like protein 3 expression and therapeutic targeting in neuroendocrine prostate cancer. Sci Transl Med. 2019. 11: eaav0891. Liverani C et al Endocrine Pathol 2021. Diagnostic and Predictive Role of DLI3 expression in Gastroenter opancreatic Neuroendocrine Neoplasms. 32:309-27. Hermans BCM et al. DLI3 expression in large cell neuroendocrine carcinoma (LCNEC) and association with molecular subtypes and neuroendocrine profile. Lung Cancer 2019. 138:102-8. TC. T. Cell enacer.

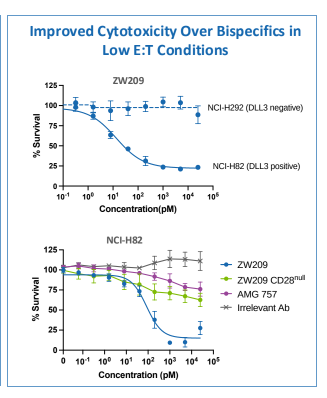
ZW209: Mediates Enhanced and Sustained Cytotoxicity



ZW209 demonstrates conditional CD28 binding and target-dependent anti-tumor activity

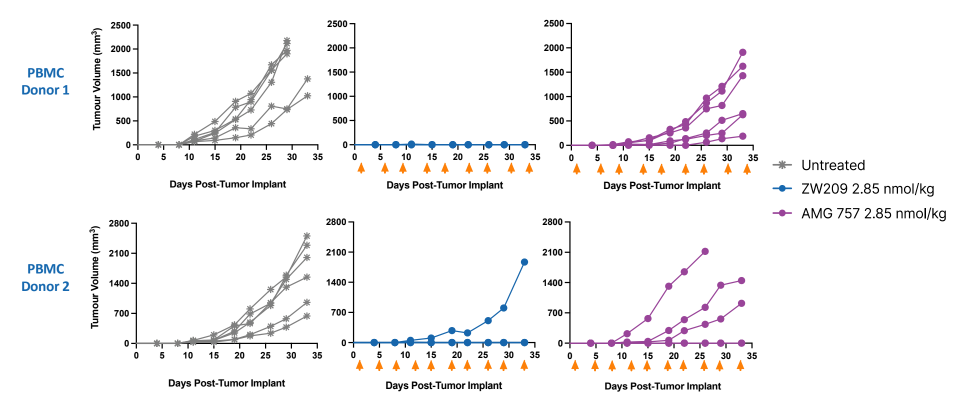






ZW209 Mediates Enhanced Anti-Tumor Activity *In Vivo* Compared to Benchmark Bispecific TCE in Humanized SCLC Models

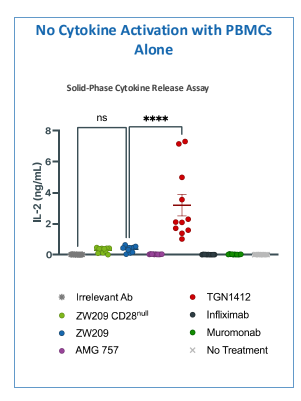


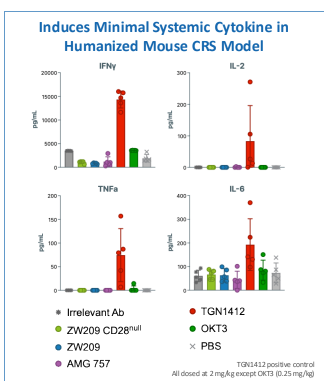


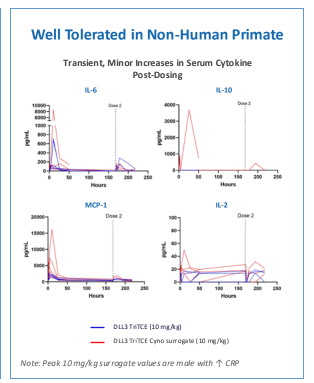
AMG 757 produced by Zymeworks for internal preclinical studies.

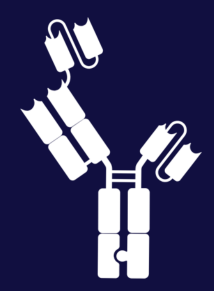
ZW209 has a Favorable Safety Profile in *In Vitro* and Animal Studies











ZW209 DLL3 TriTCE Co-stim

Therapeutic Program for the Treatment of DLL3-Expressing Solid Tumors





Design

- Trispecific TCE with potentially optimized TAA, CD3, CD28 binding affinity and geometry using Azymetric[™] and EFECT[™] platforms
- Obligate cis-T cell binding with conditional CD28 engagement



Mechanism

- Targets DLL3-expressing tumor cells and CD3 and CD28 on T cells
- DLL3-dependent T cell mediated cytotoxicity prevents activation of effector T cells in the absence of TAA



Profile & Opportunity

- Differentiated long term cytotoxicity at low effector to T cell ratios, increased T cell proliferation, survival, and anti-tumor activity with reduced cytokine release
- First in class TriTCE Co-stim opportunity on validated target
- Potential to increase durability of responses in DLL3 expressing cancers
- On track for IND submission 1H 2026





ADVANCE PROGRAM

Solid Tumors and Beyond

Paul Moore, PhD Chief Scientific Officer



AD-VAN-CE Portfolio: Progressing "First In Class" Therapeutics



- 1. Focus on novel "first in class" multi-functional therapeutics: novelty of modality, mechanism of action, and/or targeting strategy. Disruptive therapeutics with high potential benefit to patients.
- 2. Build on competitive edge in ADCs and protein engineering: cross complementary MoA and pathway axes across Zyme portfolio.
- **3. Continue to focus on select therapeutic opportunities in solid tumors:** expand portfolio coverage with GI tract and thoracic cancers.
- **4. Expand technology application to Heme-Onc, Autoimmune and Inflammatory Disease:** targeted areas conducive to multifunctional therapeutic intervention; overlap with company expertise.

Antibody-Drug Conjugates

- Novel Payload(s) beyond TOPO1i
- Bispecific/Biparatopic(s)
- Novel Targets and Target Pairs
- Payload modalities beyond cytotoxics

Cell Engagers

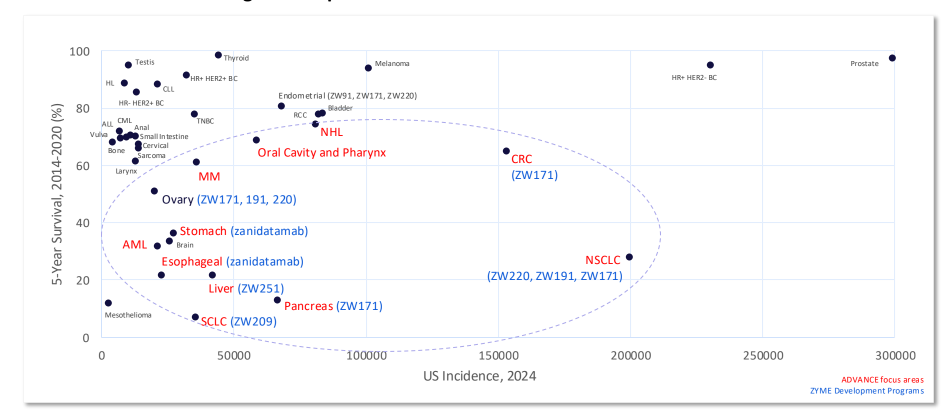
- Muti-specific T Cell Engagers
- Multi-antigen targeting
- Conditional activation
- Novel targets (e.g. proteomics)
- Intracellular antigens

Cytokine Engineering

- Tumor specific cytokine activation
- Combination Checkpoint Inhibition/cytokine activation
- Chemokine incorporation
- Multi-cytokine blockade (Autoimmune)

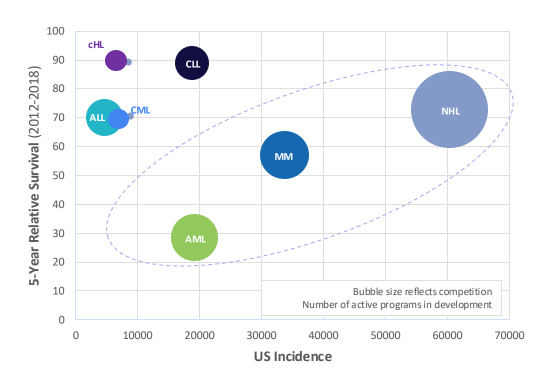
Maintain Focus on Cancers with Highest Unmet Medical Need: Increase GI Tract and Thoracic Cancer Coverage and Expand to Heme-Onc Cancers





Opportunities in NHL, MM, AML Compatible with ZYME Technology





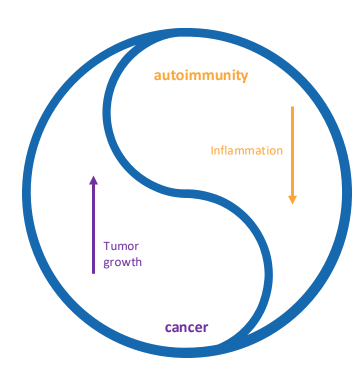
- Despite good outcomes for some, many patients with common blood cancers are not cured by frontline therapies. Patients with refractory or multi-drug resistant disease need more effective therapies.
- Outcomes for patients with AML are especially poor.
- ADCs and TCEs have proven to be important therapeutic classes to treat ALL, cHL, MM and NHLs. ZYME's differentiated technologies and next-generation multi-functional therapeutics may improve upon earlier approaches to provide better outcomes for patients.
- ZYME designed ADCs and TCEs provide opportunity for combination strategies with SOC, and potentially with each other (e.g., induction v maintenance).

Competition = program count from Cortellis filtered by indication of interest, development stage, and active developments ince 1 Jan 2018; US incidence is approximate — GLO BOCAN does not split leukemias or lymphomas by cell type; NHLs are diverse and comprised of B-, T- and NK cell neoplasms; CLL/SLL are considered NHLs, but are split out due to distinct treatment; chart reflects 5-year relative survival in aggregate but aggressive lymphomas e.g. DLBCL and indolent lymphomas e.g. FL, have dramatically different survival outcomes i.e. 5-Year Relative Survival for DLBCL is ~65%, 5-Year Relative Survival for FL is ~90%; Vç

Cancer Immunity and Autoimmunity are Two Sides of the Same Coin



- The resounding recent success of cancer immunotherapy has spurred rapid development in precision medicine
- Many cancer immunotherapy drugs aim to unleash endogenous cancer responses
- The regulatory mechanisms that hold back cancer immunity are often mechanisms that evolved to limit autoimmunity
- Therefore, by applying knowledge of these pathways, Zymeworks could address AIID
- To expand the breadth of Zymeworks' pipeline, we can take advantage of significant internal expertise and existing molecules to rapidly develop programs in AIID



Harnessing Zymeworks' Strengths to Address AIID





Autoimmune Disease:

- Chronic immune responses
- lymphocytes
- Self antigens
- Amenable to cell depletion approaches
- \rightarrow SLE
- \rightarrow RA
- \rightarrow T1D

B cell depletion and immune cell reprogramming compatible with Zymeworks' next-generation
T cell engagers.



Inflammatory Disease:

- Chronic dysregulation of inflammation
- Innate immune cells
- Environmental factors
- Amenable to cytokine blockade
- → Asthma
- \rightarrow COPD
- → Atopic dermatitis
- \rightarrow IBD

Multispecific antibody molecules to simultaneously block multiple pathways to enhance therapeutic response.

ADVANCE Program Presenters





Alex Berezhnoy, PhD Director *Immunology*



Jamie Rich, PhD Senior Director Technology ADC Therapeutic Development



Stuart Barnscher Senior Director Preclinical Programs ADC Therapeutic Development



Senior Director Multispecific Antibody Research



Nina Weisser, PhD Thomas Spreter Von Kreudensteir PhD Senior Director Protein Engineering



ADVANCE PROGRAM

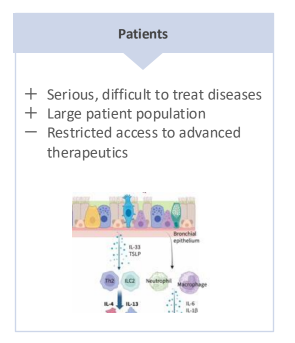
ZW1528: IL-4Rα x IL-33 Bispecific Blocker for COPD

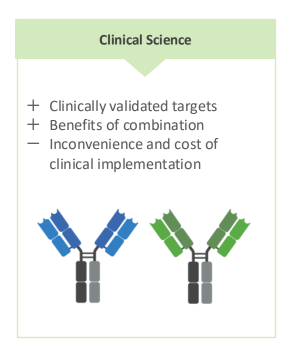
Alex Berezhnoy, PhD Director, Immunology

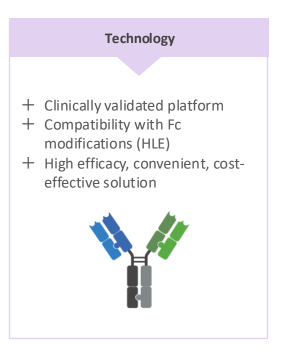


Bispecific Antibody Therapeutics as the Potential Answer to Complex Biology of Autoimmune and Inflammatory Diseases







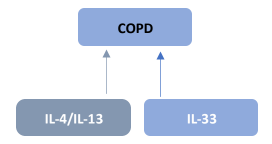


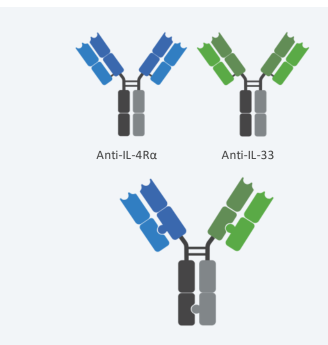
Zymeworks brings a wealth of experience in this area, both within our R&D team, as well as with previous collaborations in this space.

Rationale for Anti-IL-4Rα as an Anchor Arm



- Dupixent®/dupilumab is a highly successful mAb targeting IL-4Rα
 - Approved for multiple atopic and inflammatory diseases
 - Generated revenues >\$11Bn in 2023
- Blocking IL-4Rα inhibits both IL-4 and IL-13 signaling
 - Two key cytokines responsible for driving Type II inflammation
- Multiple cytokines drive pathology of respiratory inflammation
 - Add inhibition of an additional inflammatory pathway to augment or improve on monotherapy effects
 - ZYME opportunity to develop more efficacious molecules



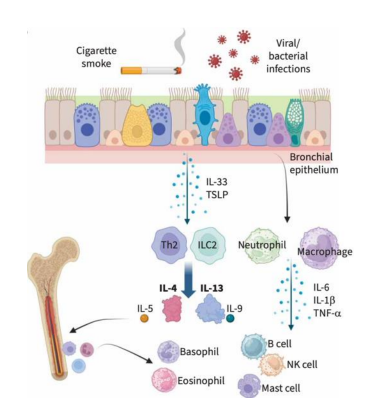


- Aim at complete, prolonged blockade of IL-4Rα
- Utilize potential advantages of local retention
- Take advantage of IgG-like geometry (PK, CMC)

IL-33 as a Bispecific Arm for COPD and Other Respiratory Diseases



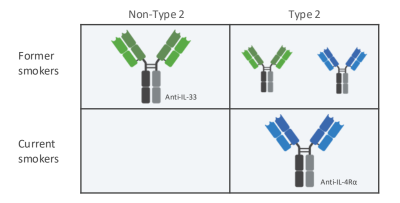
- IL-33 is a tissue alarmin released in response to epithelial damage
 - Acts on a range of cells e.g., neutrophils,
 Th2 cells, eosinophils, and mast cells
- Initiates and amplifies inflammatory response Perpetuates chronic immune response
 - May also drive tissue remodelling in chronic inflammatory diseases e.g., COPD and asthma
- Clinical proof-of-concept for targeting IL-33
 - In former smokers with COPD, and in asthma
 - Phase III trials underway for anti-IL-33 mAbs Itepekimab
 [Regeneron / Sanofi] and Tozorakimab [Astra Zeneca]



IL-4Rα x IL-33 Bispecific Provides Opportunity to Treat Broader Set of COPD Patients with Single Molecule



Anti-IL4Rα (Dupixent®) and anti-IL-33 (itepekimab) are being developed to treat different COPD populations

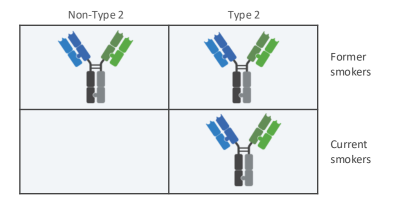


Anti-IL4Rα effective in Type 2 COPD (those with eosinophilia)

Anti-IL-33 may prove to be effective in former smokers

Post-hoc analyses of phase II data (NCT03546907)

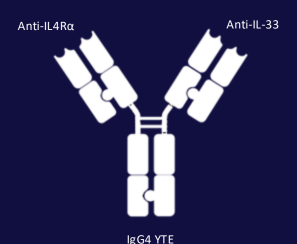
IL-4Rα x IL-33 bispecific provides opportunity to treat broader set of COPD patients with single molecule



IL-4R α x IL-33 bispecific to combine the effects of two mAbs

Potential for increased efficacy in monotherapyresponsive patients





ZW1528 IL-4Rα x IL-33 Bispecific

Inhibits multiple pathways within complex pathophysiology of inflammation



Design

- In-house antibody discovery of novel anti-IL4Rα and IL-33 paratopes
- Native IgG-like geometry; highly manufacturable, compatible with half-life extending Fc modifications
- Clinically-validated targets; core arm mediates complete, prolonged IL-4Rα blockade. Second arm adds inhibition of IL-33 - an upstream cytokine involved in perpetuating chronic inflammation.



Mechanism

- Inhibition of 3 cytokines in single asset
- Potential advantages of local retention

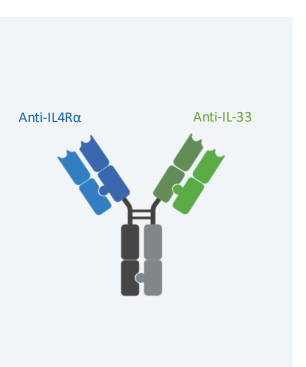


Profile

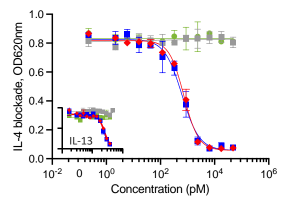
ZW1528 potently blocks of two complementary pathways of respiratory inflammation

ZW1528 Effectively Blocks of IL-4/13 and IL-33 Signaling

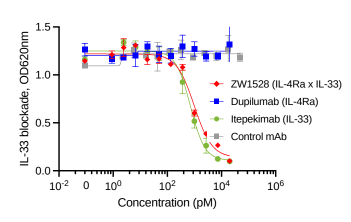




Blockade of IL-4/13



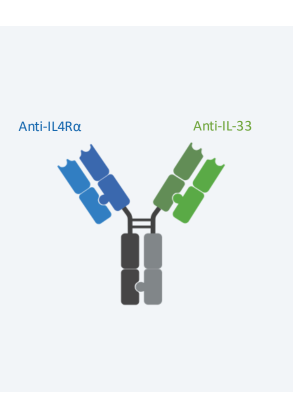
Blockade of IL-33



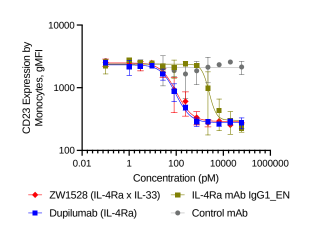
- Potency of ZW1528 similar to (bivalent) benchmark mAbs
- ZW1528 blocks both targets

ZW1528 Exhibits Favorable *In Vitro* Potency in Primary Cells



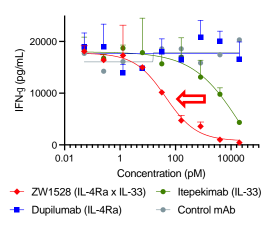


Blockade of IL-4-driven Monocyte activation



Superior potency vs IgG1 effector-negative IL-4R α mAb

Blockade of IL-33 induced IFN-γ secretion in PBMCs

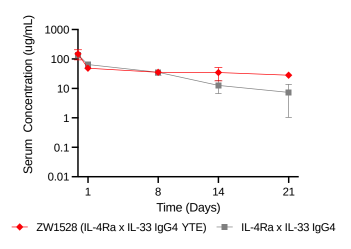


Superior potency vs Itepekimab in PBMC (blocking IFN- γ)

ZW1528 Demonstrate IgG-like PK and Block IL-4Rα In Vivo

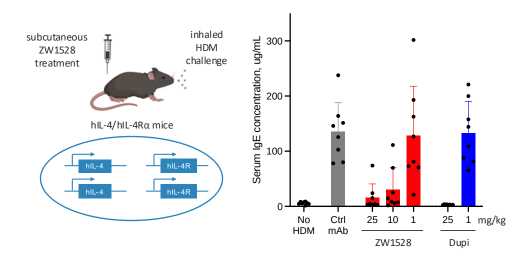


IgG-like PK (Tg32 mice)



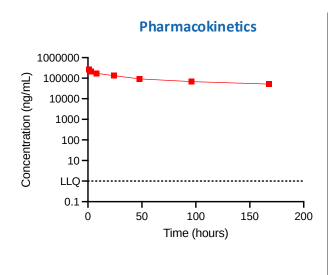
- IgG-like pharmacokinetics
- Suppression of IgE after inhaled allergen challenge

Suppression of IgE after inhaled allergen challenge

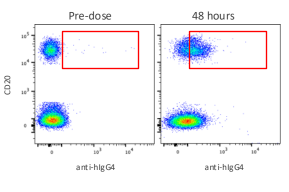


ZW1528 Demonstrates Biomarkers of IL-4R α /IL-33 Blockade in NHP

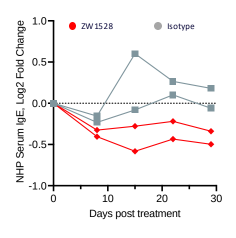








Reduction of Serum IgE

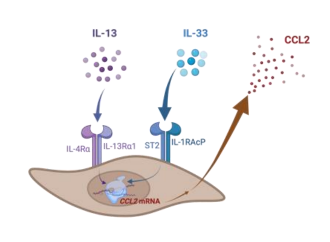


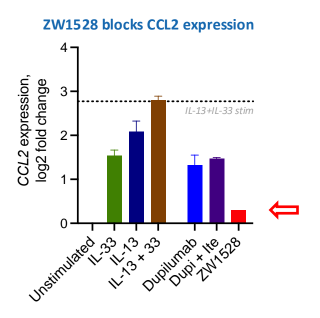
- IgG-like pharmacokinetics in non-human primates (NHP)
- Biomarkers of IL-4Rα/IL-33 blockade up to **6 weeks** after single administration

ZW1528-mediated Blockade of Primary Cell Activation is Superior to Dupilumab and Itepekimab



IL-33 and IL-13 activate human keratinocytes





- IL-13 and IL-33 treatment induces disease-relevant genes in keratinocytes
- ZW1528-mediated blockade is superior to dupilumab, itepekimab and the combination of two antibodies

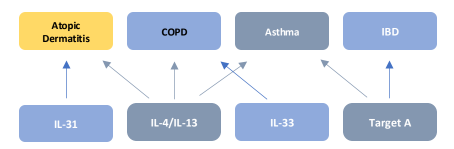
Molecules generated by Zymeworks from published sequences.

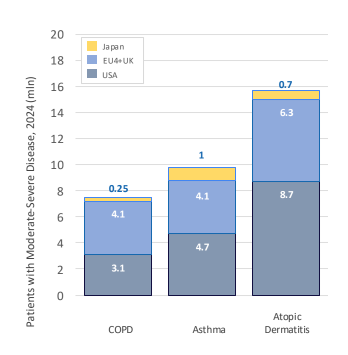
CCL2 is also known as macrophage chemoattractant protein (MCP-1) and has a role in attracting pro-inflammatory or

Expansion to AIID: Broadens Portfolio using Validated IL-4Rα Blocker



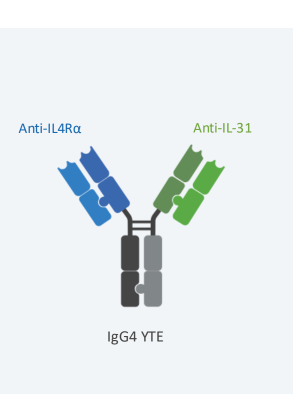
Program	Target Pair	Target Validation
ZW1528 (IND-enab)	IL4Rα x IL-33	Anti-IL4Rα approved in COPD Anti-IL33 in pivotal COPD phase 3 studies
ZW1572 (Candidate nomination)	IL4Rα x IL-31	Anti-IL4Rα approved in Atopic Dermatitis Anti IL-31 validated clinically for itch control
Next gen (<u>AD</u> VAN <u>CE</u>)	IL4Rα x Target A	Anti-IL4Rα approved in Asthma Target A efficacious in multiple AIIDs



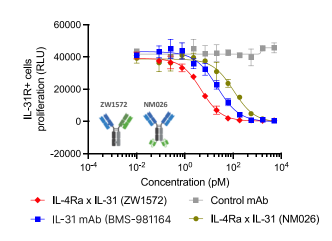


ZW1572: Bispecific Inhibitor of IL-4Rα and IL-31 for Atopic Dermatitis



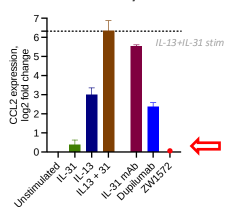


Enhanced blockade of IL-31



Superior IL-31 blockade vs. (bivalent) clinical benchmarks

Suppression of CCL2 induction in keratinocytes



Superior potency vs. individual mAbs in primary cells

Summary: ZW1528, an IL-4Rα x IL-33 Bispecific Antibody



IL-33 bispecific antibody has the potential to be a significant new treatment option for patients with COPD

ZW1528 potently blocks two complementary pathways of respiratory inflammation

- Repression of Th2 driven inflammation via blockade of IL-4Rα and inhibition of (non-Th2) IL-33-driven inflammation
- Favourable PK profile and biomarkers of IL-4R α /IL-33 inhibition in non-human primates up to 6 weeks after administration
- Preliminary evidence of advantages of bispecific blockade in disease-relevant cell types

ZW1528 aligns with requirements for successful AIID therapeutics

- Disease-modifying advantages of co-localized dual target blockade
- Stable IgG-like bispecific molecule (easy-to-manufacture, expected COGS advantages)
- Designed to allow for less frequent dosing (patient convenience and compliance)

Zymeworks' AIID opportunity

- Potential to benefit mixed-type COPD patients, and increase benefits for other COPD patient populations
- Clinical potential in other atopic diseases (asthma and atopic dermatitis) with established IL-4Rα and/or IL-33 role
- Spearhead development of other Zyme AIID therapeutics



ADVANCE PROGRAM

Next Generation Technologies



ADVANCE PROGRAM

Antibody-Drug Conjugate Technologies



Jamie Rich, PhD
Senior Director
Technology
ADC Therapeutics

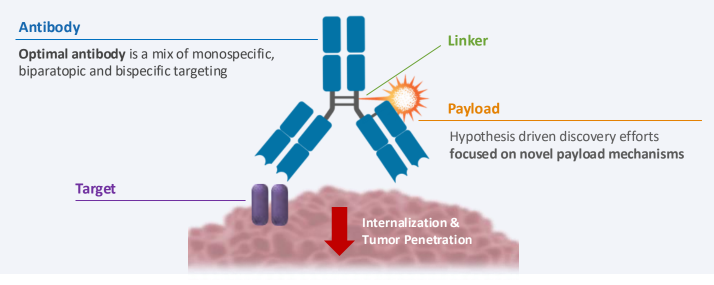


Stuart Barnscher
Senior Director
Preclinical Programs
ADC Therapeutics

ADVANCE: Novel Payloads and Optimized Antibody Formats for Next Generation ADCs



We are developing **novel payload mechanisms** & **antibody formats** to suit disease and target biology



ADVANCE ADCs will support our 2027+ IND pipeline

Therapeutic Applications

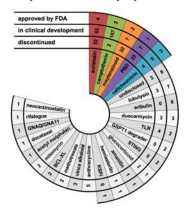
- Solid Tumors: current focus GI tumors including CRC, PDAC, esophageal, HNSCC
- Heme Onc: current focus AML

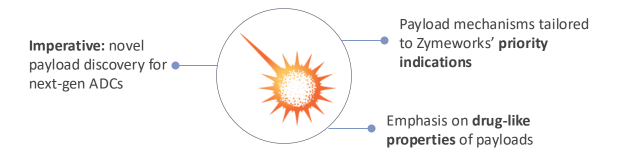
CRC: Colo rectal cancer, PDAC: Pancreatic ductal adeno carcinoma; HNSCC: Head and neck squamous cell carcinoma; AML: Acute myeloid leukemia

Novel Payloads to Help Drive Innovation in the ADVANCE ADC Portfolio



40 years of ADC payloads:1





Challenging nature of ADC Platform development necessitates multiple approaches

Cytotoxic ADCs

- + Validated approach
- + Broad utility
- Normal tissue toxicity

Conditionally Cytotoxic ADCs

- + Reduced normal tissue tox
- + Built in patient biomarker
- + Restricted patient group

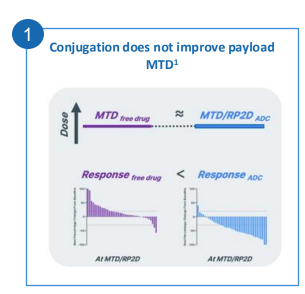
Protein Degradation

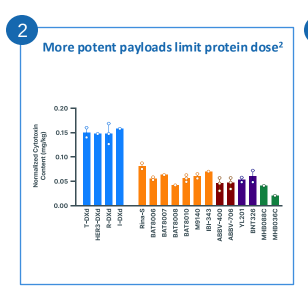
- + New and existing targets
- + Potential tissue selectivity
- + Limited validation
- + Restricted patient group

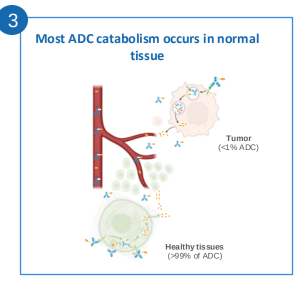
1. R. Colombo et al. Cancer Discov 2024, 14(11):2089-2108.

Our Hypothesis on ADC Mechanism Guides our Approach to Novel Payload Discovery









Criteria for novel payloads:

- 1. Moderate potency to enable higher protein dose
- 2. Bystander activity
- 3. Evidence of activity in disease indication of interest

1. R. Colombo and J.R. Rich, Cancer Cell 2022, 40(11):1255-1263; 2. R. Colombo et al, Cancer Discov 2024, 14(11):2089-2108

Four Novel Payload Mechanisms in Discovery to Help Drive Innovation Beyond TOPO1i and Auristatin Platforms



Challenging nature of ADC Platform development necessitates multiple approaches

Cytotoxic ADCs

Translation Inhibitor
Platform

Targeting protein synthesis pathway. Novel payloads with optimized potency

Conditionally Cytotoxic ADCs

Oncogene Targeting
Platform

Inhibitor of key oncoprotein suited to application in ZW target indications

Synthetic Lethality
Platform

First of kind synthetic lethal ADC approach

Protein Degradation

4 Protein Degrader Conjugates

Degrader antibody conjugate modality holds significant promise across diverse diseases

Current focus in diverse GI tumor indications

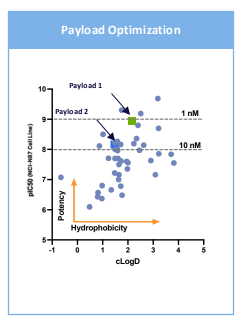
Biomarker driven application in solid and liquid tumors

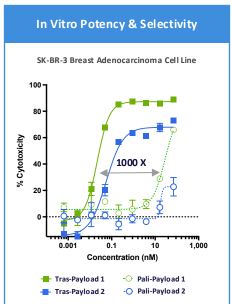
Interest in AML, PDAC, and CRC targets

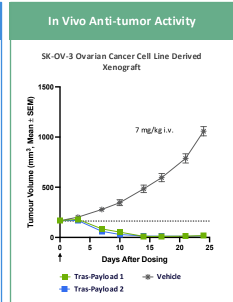
Protein Synthesis Inhibitor ADCs Demonstrate Preclinical Promise

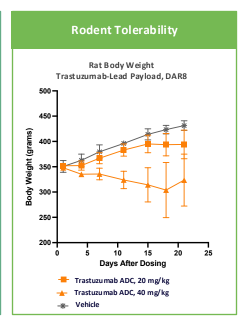


- Targeting a pathway critical to cancer cell growth enables potential first in class ADC opportunity
- · Optimized ADCs demonstrate potency, selectivity, strong anti-tumor activity, and promising rodent tolerability
- Evaluation in target indications is ongoing (H&N, PDAC, CRC); planned non-human primate toxicology studies





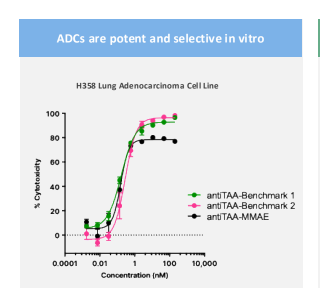


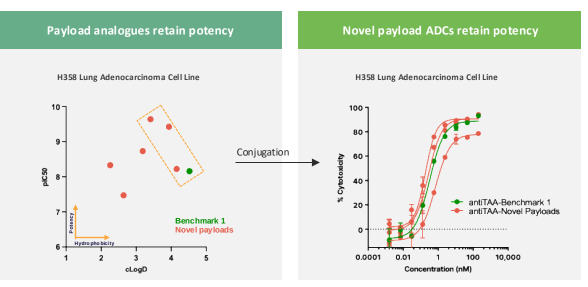


Inhibitors of Key Oncoprotein Serve as Novel ADC Payloads



- First in class ADC opportunity with cytotoxic mechanism that is well validated in our priority indications
- ADC approach may improve on small molecule drugs via increased exposure, efficacy, and tolerability
- Currently evaluating in vivo efficacy

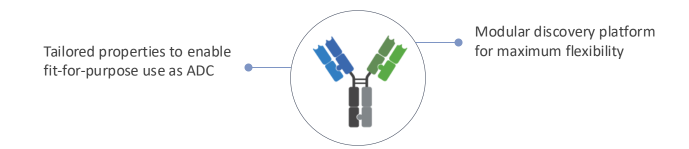




Multi-Pronged Approach to Antibody Discovery:



Seeking Optimal Antibodies for Development of Differentiated ADCs



Animal

Discovery Technology

Characterization

Developability

Engineering

Use of wildtype and transgenic animal platforms for maximum diversity

Use of **internal** B-cell culture & **external** antibody discovery technologies

Robust, high throughput screening to evaluate optimal ADC properties

Comprehensive
evaluation of
developability risks early
in discovery process

Optimized format

Selection of Desired Antibody Format for ADC Assets



- Generation of differentiated fit-for-purpose ADCs is enabled by a flexible antibody discovery workflow and the Azymetric™ platform
- Target expression and biology may dictate the use of one format over another

Monospecific



Binds to single epitope on a single target

- Suitable for targets with some normal tissue expression
- May be only format available for targets with restricted epitope space
- Internalization dependent on target biology and specific epitope

Biparatopic



Binds to two distinct epitopes on a single target

- Suitable for targets with limited normal expression
- Targets with large epitope space may be most suitable
- Internalization can be enhanced via increased surface decoration and antigen clustering

Bispecific



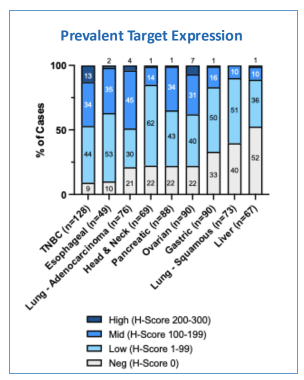
Binds to two distinct targets

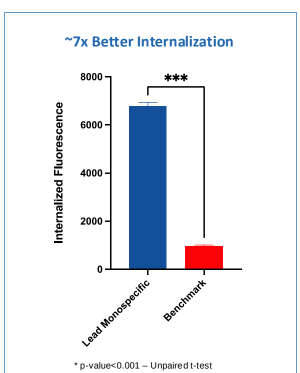
- Suitable for targets with biologic association or targets with high 'tumor coverage'
- Valency can be tuned to suit tumor and normal expression of each target
- Internalization may be enhanced due to target co-engagement

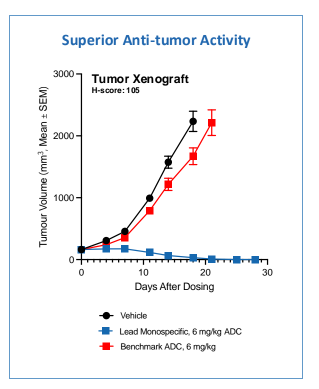
Monospecific Targeting: ADVANCE Solid Tumor ADC#1



A monospecific format was selected for this program due to the small extracellular domain of the target, limited epitope diversity, and the superior characteristics of the lead monospecific over the clinical benchmark



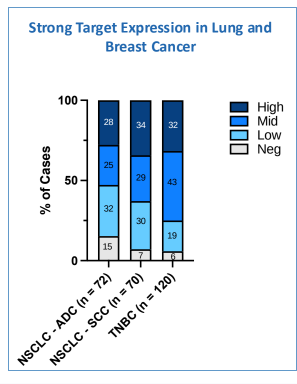


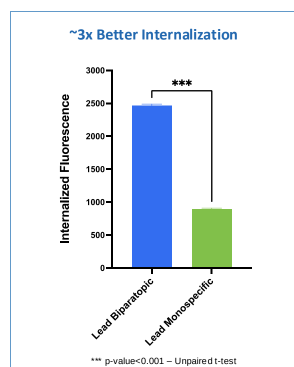


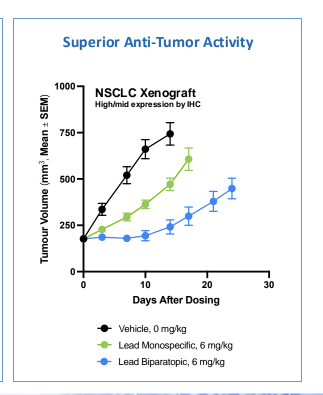
Biparatopic Targeting: ADVANCE Solid Tumor ADC#2



A biparatopic format is being evaluated for this program due to broad epitope diversity, enhanced internalization profile, and superior anti-tumor activity compared to a lead monospecific antibody





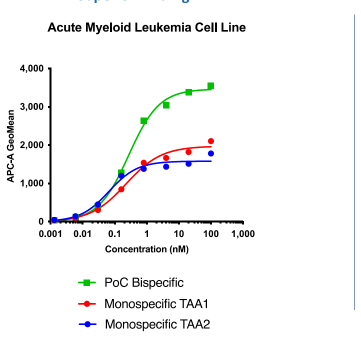


Bispecific Targeting: ADVANCE AML ADC

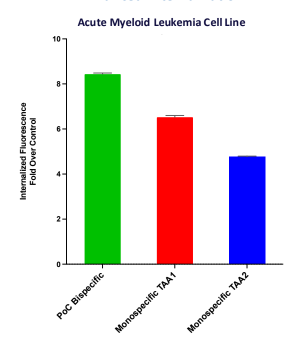


A bispecific format is being evaluated for this program to overcome individual target heterogeneity and to enhance functionality by co-targeting cells that express both targets

Superior Binding



Enhanced Internalization



AML: acute myeloid leukemia; ADC: antibody drug conjugate; PoC: proof of concept; TAA: tumor associated antigen





Novel Payload Discovery

- Our hypothesis on ADC mechanism guides our approach to novel payload discovery
- Four novel payload mechanisms in discovery to help innovation beyond TOPO1i and auristatin platforms



Optimal antibody formats

- Multi-pronged strategy to discover antibodies with enhanced ADC properties
- Antibody format (monospecific, biparatopic, or bispecific) dictated by target characteristics



Therapeutic Application

 Target, payload mechanism, and antibody format selected for enhanced activity in disease indication



For solid tumors (GI tract cancers) and heme-onc will help to support our 2027+ investigational new drug application pipeline



ADVANCE PROGRAM

Multispecific Antibody Technologies



Nina Weisser, PhD
Senior Director
Multispecific Antibody
Research

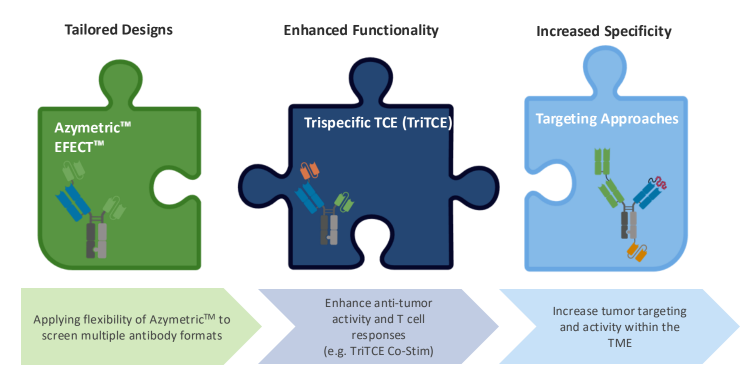


Thomas Spreter Von Kreudenstein, PhD Senior Director Protein Engineering

Driving the Forefront of Next Generation T cell Engagers

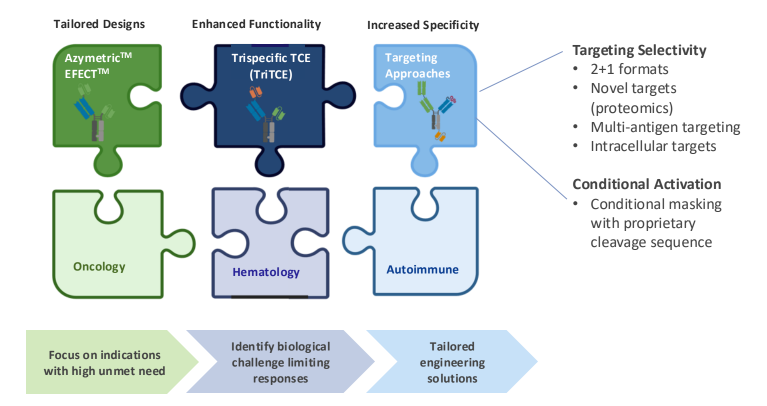


Plug and Play Platforms to Potentially Build Differentiated Therapeutic Cell Engagers



ADVANCE: Enhancing Functionality and Specificity to Help Improve Responses Across Diverse Therapeutic Areas





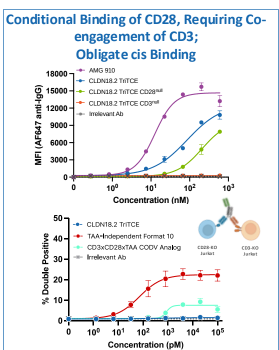


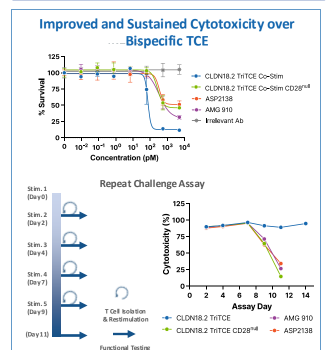
ZW239: CLDN18.2 Targeted TriTCE Co-stim to Help Improve Depth and Durability of Responses in CLDN18.2 Expressing Tumors

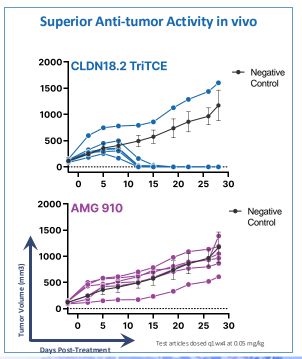


Focus indication of high unmet need
Gastric and Pancreatic

Biological challenge limiting responses Low T cell Infiltration and T cell Dysfunction Tailored engineering solutions
CLDN18.2 TriTCE Co-stim









Novel Tumor Targeting to Help Improve Treatment Responses Across Diverse Solid Tumors



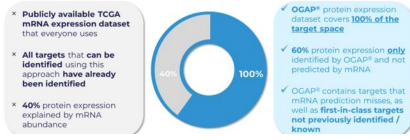
Focus indication of high unmet need Multiple Solid Tumor Indications **Biological challenge limiting responses** Low T cell Infiltration & T cell Dysfunction Tailored engineering solutions
Novel Target TriTCE Co-Stim

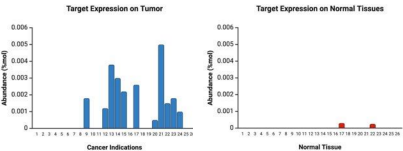


Multi-indication TriTCE Co-stim to novel target

Novel proteomic approach¹ used to identify **unique membranous tumor-specific targets** with ideal TCE target profile

- No/minimal normal tissue expression
- Supportive biology and expression profile
- Novel target with multi-indication potential





1. Houghton et al. 2024, 6th Annual Targeted Radiopharmaceuticals Summit Europe



Overcoming Antigen Escape and T cell Dysfunction to Help Improve Treatment Responses in AML



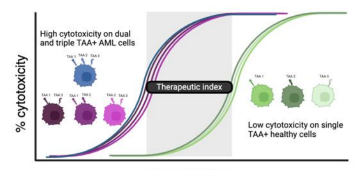
Focus indication of high unmet need AML

Biological challenge limiting responsesHealthy Tissue Expression, Antigen Escape & T cell Dysfunction

Tailored engineering solutions Multi-targeted Co-Stim

Biological Challenge	Limitation of Mono-antigen Targeted Therapies	Solution	
Heterogeneous intertumoral antigen expression	Antigen escape and treatment failure ¹	Multi-antigen targeting with co-stimulation	
Lack of a clean single target between AML blasts, LSCs and healthy cells	Narrow therapeutic window		
T cell dysfunction ²	Bispecific TCE resistance and lack of long-term responses ²	Repres entative potential format	

Selective tumor cytotoxicity in presence of 3 or 2 target antigens



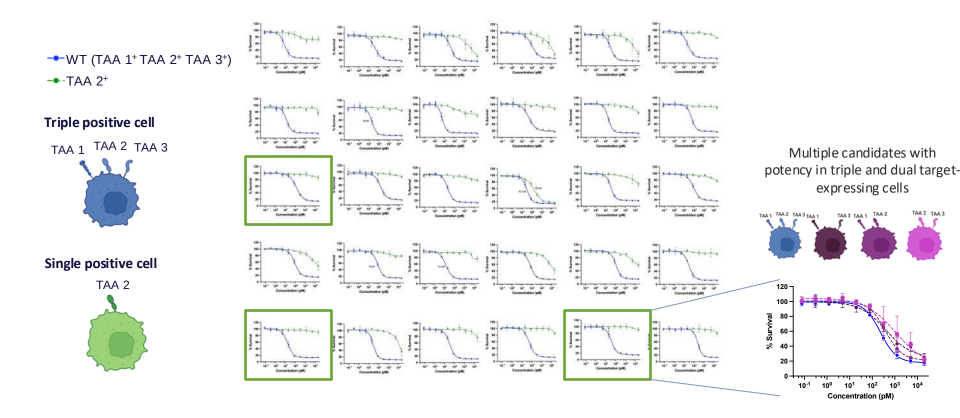
Concentration

1. Atar 2024 Leukemia, 38; 2.Kazerani et al. 2024 Leukemia, 38. AML, Acute myeloid leukemia; LSC, leukemic stem cell; TCE, T cell engager



Multiparameter Screening using Azymetric™ Identifies Candidates with the Desired Biology







Advanced Protein Engineering Solutions

- FDA approved, clinically validated, and novel engineering solutions enable plug and play building blocks to address complex biological challenges
- Flexibility of Azymetric[™] facilitates high throughput multiparameter antibody screening to identify molecules with the desired biology

Addressing Biological Challenges in Indications with High Unmet Need

- Focusing on indications with high unmet need and complex biological hurdles
- Designing next generation T cell engagers to overcome biological challenges not addressed with traditional bispecific T cell engagers

Driving the Forefront of Next Generation T cell Engagers

- Enhancing functionality and specificity to drive deep and durable responses in difficult to treat tumors
- Plug and play platforms enable fast development to rapidly address patient need

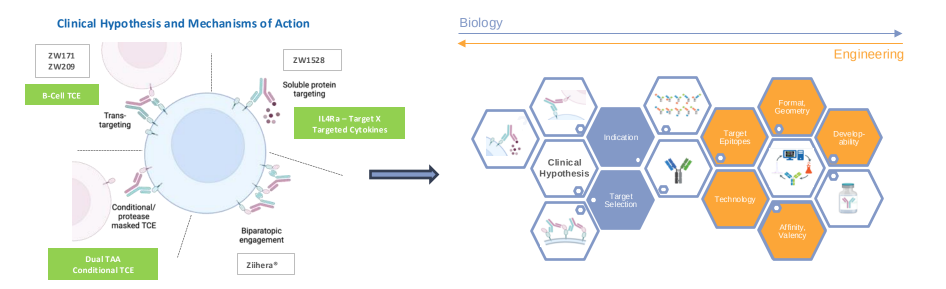


For difficult to treat tumors with complex disease biologies in oncology and hematology to support our 2027+ investigational new drug application pipeline

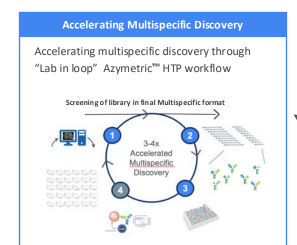
Zymeworks' Technology Development Approach: Integration of Engineering and Drug Discovery Driving Differentiation

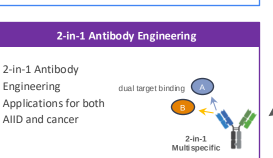


- Cross-functional approach critical to enable 'Technology Driven' innovation and differentiation
- 10+ years of structure-based Protein Engineering and Design and in-house tool development
- Critical learnings for Multispecific Engineering and further in-house Technology Development



Integration of AI/ML with Azymetric™ Engineering Expertise Enables Novel Technology Drivengueworks Drug Development





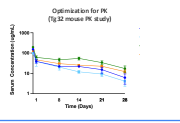
Opportunities for Combining AI/ML with Multispecific Engineering Expertise AI/ML Innovation

Physics-Based Engineering (ZymeCAD™)

Multi-parameter Antibody Optimization

Enhanced Multispecific Engineering

- Multifactorial optimization for affinity, stability, developability
- Engineering for improved PK and s.c. formulation

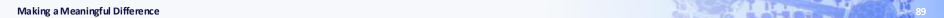


Cytokine and Receptor Engineering

Stability engineering of cytokines and natural receptors or ligands for cancer and AIID

Difficult targets (low stability, manufacturability)

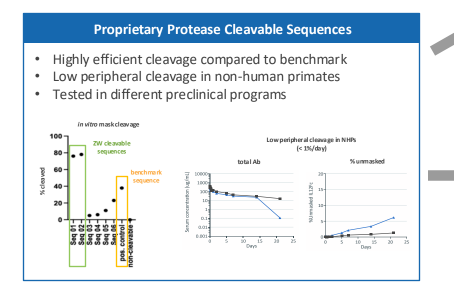




Next Gen Technologies: Conditional Masking Technologies

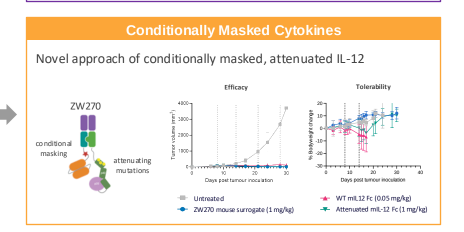


- Internal development of highly efficient tumor specific protease cleavage sequences
- Masking technology application for TCE and TriTCE
- Internal IL-12 program using attenuated IL-12 displays higher therapeutic window than clinical competitors



Expanding target space for TCE and TriTCE portfolio Potential to combine with

2+1 and CD3 engineering



Next Gen Technologies: Novel Engineering for Autoimmune and Inflammatory Indications

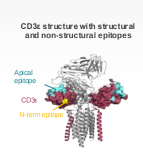


T Cell Engager Engineering for AIID

Significant opportunity for enhanced B cell depleting therapies for AIID

 $\label{lem:critical} \mbox{Critical engineering for high activity with reduced cytokine release} \mbox{syndrome potential}$

ZW has panel of anti-CD3 binders with fast Kon/off and different epitopes to fine tune activity



Name	Epitope	k _{on} (1/Ms)	k _{off} (1/s)	K _d (nM)
ZW1	Epitope 1	1.38E+05	3.12E-03	22.7
ZW2	Epitope 2	5.20E+05	4.08E-03	7.84
ZW3	Epitope 1	1.62E+05	1.10E-02	68.3
ZW4	Epitope 2	1.45E+05	1.26E-02	87.1
ZW6	Epitope 1	1.59E+05	2.54E-02	159.9

ZW fast Kon/off CD3



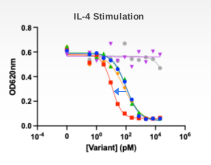
Novel Bispecific Cytokine Antagonists

Additional IL4Rα-X bispecific cytokine antagonists

Engineering for optimized PK

Bispecific engineering for superior potency to IL-4R α comparators and bispecific advantage vs. combination



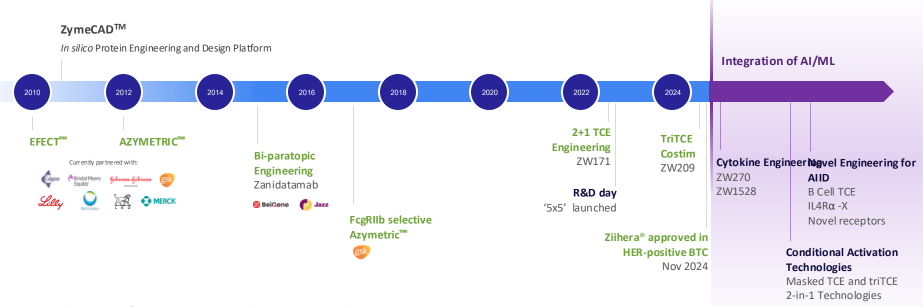


- Dupilumab
- Target A monospecific
- IL4Ra-TargetX bispecific
- Dupilumab + Target A combination
- Isotype control

Building on 10+ Years of Protein Engineering and Multispecific Development: Continued 'Technology Driven' Innovation



- Continued 'Technology Driven' Innovation and differentiation
- Integration of new AI/ML methods enable acceleration of Multispecific Discovery and novel targeting approaches
- New Technologies applied to Oncology and AIID pipeline



Integrated Approach of Protein Engineering and Technology Development →



RESEARCH & DEVELOPMENT DAY

R&D Strategy Summary: 2025-2027

Kenneth Galbraith Chair and CEO



R&D Strategy Summary for 2025-2027





Focus

- 5 x 5 solid tumor portfolio completed
 18 months ahead of original target
- Solid tumor portfolio expansion focused on mainly digestive system cancers and certain other unmet needs
- Patient populations of interest to expand beyond solid tumors to include heme cancers and AIID



Execution

- Investment in preclinical research and capabilities over 2022-24 allows for acceleration of ADVANCE strategy
- Zanidatamab experience, focused efforts and quality drives higher probability of success in portfolio
- Preclinical, clinical and TMO groups built to manage active portfolio of candidates across expanded focus areas
- R&D investment balanced across clinical candidates and preclinical research over 2025-2027



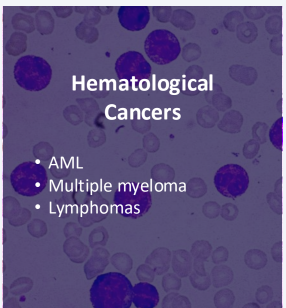
Growth

- Partnering, strong target product profiles and attrition in development will maintain a manageable portfolio of opportunities over time
- New product candidates will continue to add novelty, differentiation and value to the product portfolio through 2027

Focused Therapeutic Areas Provide Diversity to R&D Portfolio and Enhanced Optionality for Partnering and Retained Product Rights









R&D Organization



Early Research & Development

- Full internal capabilities for multi-functional therapeutics development
- 150-person group located in Vancouver laboratories
- Capacity to sustain current R&D productivity with core focus on internal development vs. in-licensing
- Building novel biology/target capabilities

Early-stage Clinical Development

- Geographically dispersed capabilities located between West Coast North America, Dublin, and Singapore
- Strong outsourcing model
- 35-person group
- Capacity to manage 5-6 early-stage clinical candidate portfolio
- Specialization between solid tumors, heme cancers and AIID

TMO & Quality

- · Strong outsourcing model
- Benefit from zanidatamab experience
- 30-person group located on West Coast NA
- Capacity to manage 10-12 product candidates

Over the past 18 months we have built an internal organization globally to actively manage ~5-6 early-stage wholly-owned clinical programs

Strong Balance Sheet to Support Meaningful Catalysts in R&D Portfolio



Current Financial Status:

- Cash resources¹ of approx. \$375M (as of Sept 30, 2024)
- Additional development activities in AIID and ADVANCE, along with nomination of ZW1528 for COPD, was executed while maintaining cash runway
- Anticipated cash runway into 2H 2027, which includes certain anticipated regulatory milestone payments

Potential sources to extend cash runway beyond 2H 2027:

- Additional regulatory approval and commercial milestones for zanidatamab from Jazz and BeiGene
- Tiered royalties between 10-20% from Jazz and 10-19.5% from BeiGene sales (up to 20% when royalty reduction of 0.5% reaches cap in the low double-digit millions of dollars)
- Additional payments from legacy technology platform collaborations
- Optionality to monetize royalties from existing partnerships
- Potential new partnerships/collaborations to provide upfront payments and committed R&D funding

Differentiated Development of Multifunctional Therapeutics



Program	Technology	Target	Indication	Discovery	Preclinical	Phase 1	Phase 2	Phase 3
Solid Tumor Oncology: Antibody-Drug	Solid Tumor Oncology: Antibody-Drug Conjugates (ADC)							
ZW191 Topo1i ADC DAR 8 Fc WT	ZD06519 Payload	Fra	Gynecological Thoracic	NCT06555744				
ZW220 Topo1i ADC DAR 4 Fc Mut	ZD06519 Payload	Napi2b	Gynecological Thoracic			IND	1H 2025	
ZW251 Topo1i ADC DAR 4 Fc WT	ZD06519 Payload	GPC3	Digestive System (HCC, PDAC)			IND	2H 2025	
Solid Tumor Oncology: Multipecifics								
Za ni datamab Bis pe ci fic	Azymetric™	HER2	Multiple indications					
ZW171 Trivalent TCE 2+1 Format	Azymetric [™] Novel anti-CD3	MSLN x CD3	Gynecological Thoracic	NCT06523803				
ZW209 Trispecific TCE Tri-TCE Costim	Azymetric [™] Novel anti-CD3 Conditional CD28	DLL3xCD3xCD2 8	Thoracic			IND 1H 2026	ō	
ZW239 Trispecific TCE Tri-TCE Costim	Azymetric™ Novel anti-CD3 Conditional CD28	CLDN18.2x CD3	Digestive System					
AllD								
ZW1528 Dual Cytokine Blocker	Azymetric [™] Hetero-Fab YTE	IL4RaxIL33			IN	ID 2H 2026		
ZW1572 Dual Cytokine Blocker	Azymetric [™] Hetero-Fab YTE	IL4Rα x IL-31						

Meaningful Catalysts Expected Throughout 2025 & 2026



1H 2025	2H 2025	2026
PIPELINE EVENTS		
 Pivotal Phase 3 top-line data readout in 1L GEA for zanidatamab targeted by our partner Jazz in 2Q 2025 Potential China approval for zanidatamab in 2L BTC in Q2 2025 Initial royalty revenue from Jazz Pharmaceuticals Expected IND submission for ZW220 (NaPi2b) in 1H 2025 Potential EU approval for zanidatamab in 2L BTC in Q2 2025 	 Initial royalty revenue from BGNE Pharmaceuticals Expected IND submission for ZW251 (GPC3) in 2H 2025 Potential Phase 1 clinical data readouts on ZW171 and ZW191 as soon as 2025 Jazz Pharmaceuticals may file a sBLA for zanidatamab in 1L GEA Jazz potential for further development of zanidatamab in neoadjuvant/adjuvant GEA 	 Expected IND submission for ZW209 (DLL3) in 1H 2026 Expected IND submission for ZW1528 (IL4R x IL-33) in 2H 2026 Jazz to potentially launch expanded market strategy for zanidatamab in GEA in 2026

 ${\sf CASH\,RUNWAY^1\,FORECAST\,INTO\,2H\,2027\,WITH\,RECEIPT\,OF\,CERTAIN\,ANTICIPATED\,REGULATORY\,MILESTONE\,PAYMENTS}$

Illustrative. Key news flow only. AACR: American Association for Cancer Research; ASCO: American Society of Clinical Oncology; SASCO 6t ASCO Gastrointestinal Cancers Symposium; BIA: biologics license application; EORTC-NCI-AACR: EORTC-NCI-AACR: Symposium on Molecular Targets and Cancer Thera peutics; EMA: European Medicines Agency; ESMO: European Society for Medical Oncology; ISMO: Japanese Society of Medical Oncology; MAA: marketing authorization application; PEGS: Protein Engineering Summit; SABCS: San Antonio Breast Cancer Symposium; SITC-Society for Immunotherapy of Cancer; T-Dxxl: trastuzumab deruxtecan; World ADC: World Antibody Drug Conjugates Summit; WCGI: World Congress on Gastrointestinal Cancer.

¹ cash, cash equivalents, and marketable securities

Key Takeaways From ZYME R&D Day





With nomination of ZW209, 5x5 solid tumor portfolio construction is 18 months ahead of schedule with expected initial clinical data disclosures potentially starting in 2025



Recent approval of zanidatamab demonstrates our experience and abilities to develop unique and differentiated therapeutics with clinically meaningful benefits for patients



ADVANCE portfolio broadly diversified into hematological cancers and AIID in additional to solid tumors with initial IND planned for 2H-2026 for ZW1528 and more in 2027 and beyond



Clear decision-making processes to advance or cease development activities on product candidates based on clinical data generated



Enhanced optionality for partnerships and collaborations to share capital and development risk



Strong financial position to provide opportunity for retaining certain product rights



R&D organizational structure in place to drive continued progress in both '5x5' and ADVANCE portfolios



Additional solid tumor research focused on digestive system cancers, including CRC and PDAC



ADVANCE PROGRAM

Q&A Session #2

