#### AUTHORS

Valentine de Puyraimond†, Patrick Rowe†, Juntao (Matt) Mai<sup>†</sup>, Phillipe Pouliot, Kate Caldwell, Lauren Clifford, Allie Goodman, Patrick Farber, Harveer Dhupar, Davide Tortora, Peter Bergqvist, Janice Reimer, Karine Herve, Vivian Li, Irene Yu, John Marwick, Stefania Carrarra, Nathalie Blamey, Melissa Cid, Cindy-Lee Crichlow, Lena M. Bolten, Tova Pinsky, Gesa Volkers, Jessica Fernandes Scortecci, Girija Bodhankar, Caitlyn De Jong, Sohpie Cullen, Ping Xiang, Courtenay Lai, Ahn Lee, Gabrielle Conaghan, Antonios Samiotakis, Stefan Hannie, Rhys Chappell Emma Lathouwers, Lauren Chong, Kate Gibson, Stephanie K. Masterman, Kirstin Brown, Kelly Bullock, Raffi Tonikian, Lindsay DeVorkin, Sherie Duncan, Aaron Yamniuk, Kush Dalal, Tim M. Jacobs, Bryan C. Barnhart\*

†these authors contributed equally \* presenter

AUTHOR AFFILIATION AbCellera, Vancouver, Canada



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#### OUR PLATFORM

### Diverse CD3-binding antibodies for TCE development

Effective CD3 T-cell engagers (TCEs) must balance potency with potential toxicities by maximizing tumor-cell killing while fine-tuning cytokine release. However, most TCEs in clinical development are derived from a small number of CD3-binding antibodies, such as SP34-2<sup>1</sup>, limiting the number of antibodies available for optimal immunological synapse formation.

To address these barriers, AbCellera developed a complete TCE platform: a discovery engine to identify diverse tumor-binding antibodies; a portfolio of fully human, CD3-binding antibodies that are distinct from SP34-2; a clinically validated multispecific engineering platform; and a high-throughput strategy to rapidly identify TCE candidates<sup>2-4</sup>.

#### THE CHALLENGE

## CD3-binding antibodies result in diverse TCE functional profiles

Previously, we demonstrated that identical CD3-binders result in variable function when paired with different tumor-binding arms<sup>4</sup>. However, due to the limited number of CD3-binding antibodies that have been assessed in the field, the full extent to which CD3-binding parameters impact TCE function is not well understood.

To address this, we designed and engineered our largest panel of TCEs to date. We paired hundreds of CD3-binders with a single tumor-binding paratope and assessed bispecifics in high-throughput functional assays. We performed a clustering analysis to identify sets of CD3-binders that result in similar function as bispecifics, and developed a process that leverages these functional clusters to streamline selection of CD3-binders for different tumor targets.

### OUR APPROACH

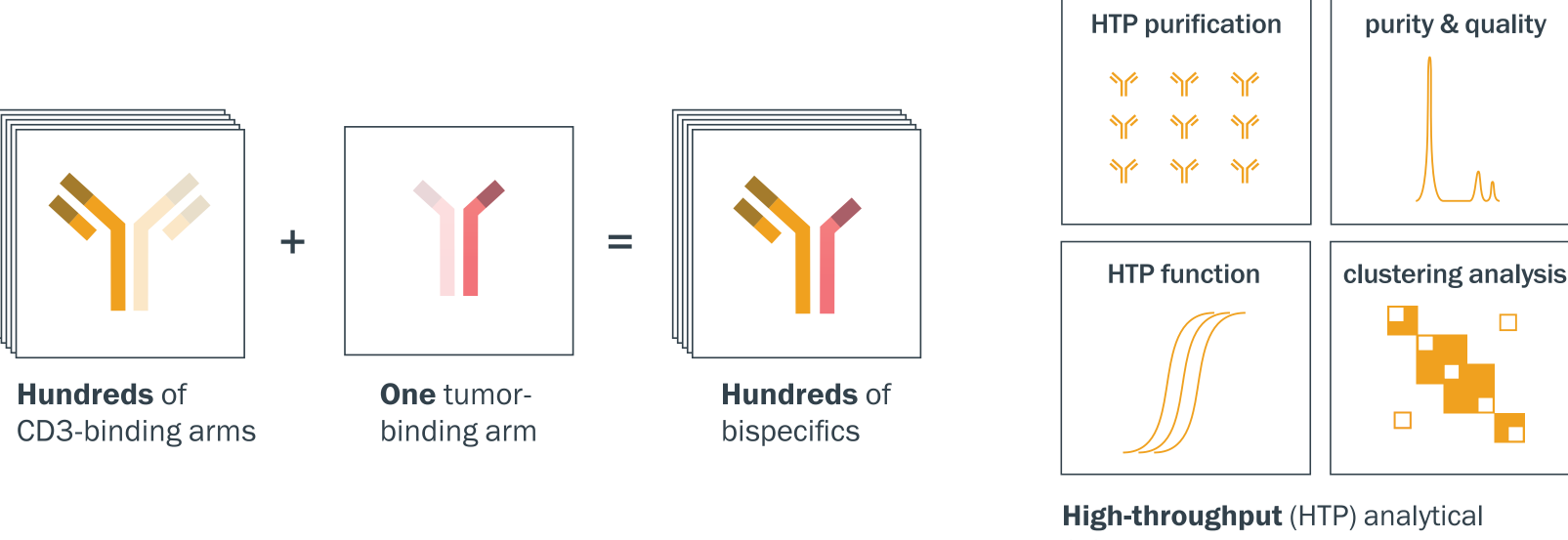
### A streamlined approach for identifying the optimal CD3-binder for any tumor target

We integrated large antibody datasets to gain insights into parameters that impact TCE function. We combined these with our previous data demonstrating the impact of the tumor-binding arm on TCE function and developed a strategy to rationally select tumor- and CD3-binding antibodies for each target. We leveraged these data to select TCEs for further assessment for multiple programs, two of which are shown here.

# Multiple CD3-binding parameters impact TCE function

### CD3-binding kinetics impact tumor-cell killing and cytokine release

(A) We engineered a panel of TCEs to assess the impact of the CD3-binding arm on function



#### **B** We compared CD3-binding kinetics to TCE function

Colored by CD3-binding antibody clonal family

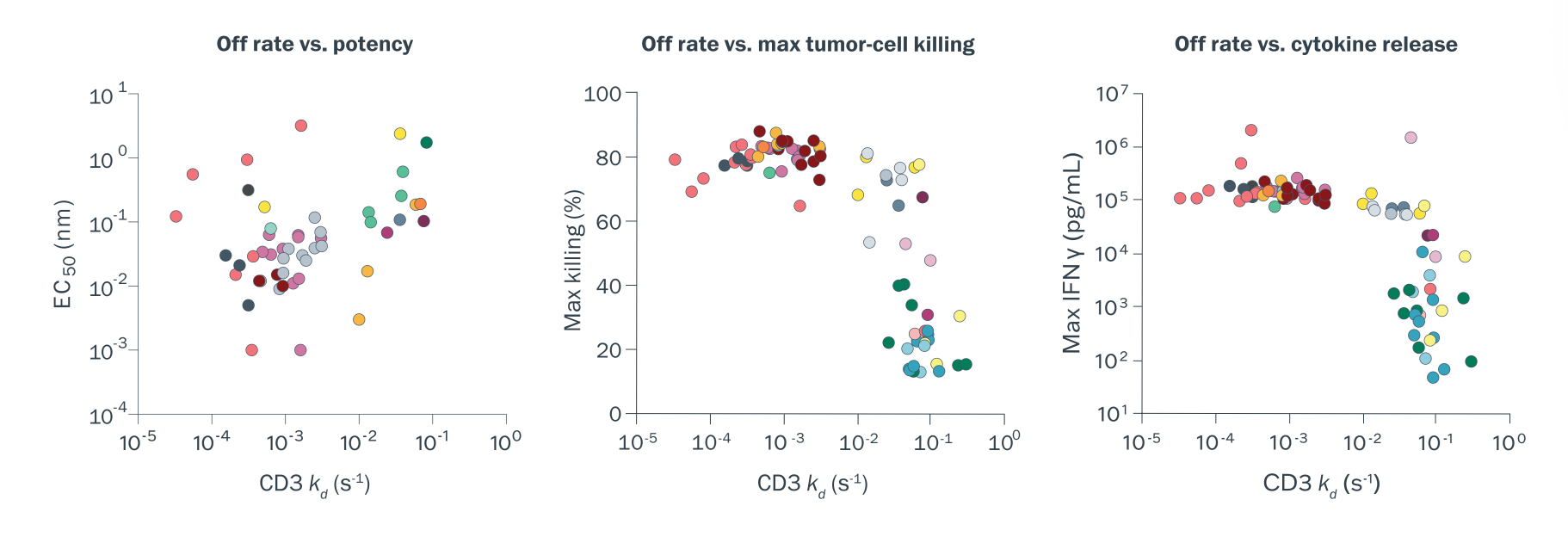


Figure 1. Hundreds of bispecifics were engineered to assess the impact of the CD3-binding arm on **TCE function. (A)** AbCellera's CD3-binding antibodies<sup>2-4</sup> were paired with a single tumor-binding arm. The resulting bispecifics were expressed and analyzed using high-throughput functional assays. (B) Binding kinetics for the human CD3  $\epsilon\delta$  subunit were compared to TCE potency, tumor-cell killing, and cytokine release. Because off rate was found to be the primary driver of affinity, and release of TNFα and IL-2 was consistent with that of IFNγ, these parameters are shown. CD3-binding antibody clonal family (defined as antibodies sharing the same V gene, J gene, CDR3 length, and animal origin for both heavy and light chains) is overlaid using color. Off rate was found to be a significant contributor to cytokine release and tumor-cell killing, but was not sufficient to fully describe these properties, particularly at faster off rates. In contrast, potency was not tightly correlated with either off rate or clonal family.

#### Binding kinetics alone cannot predict TCE function

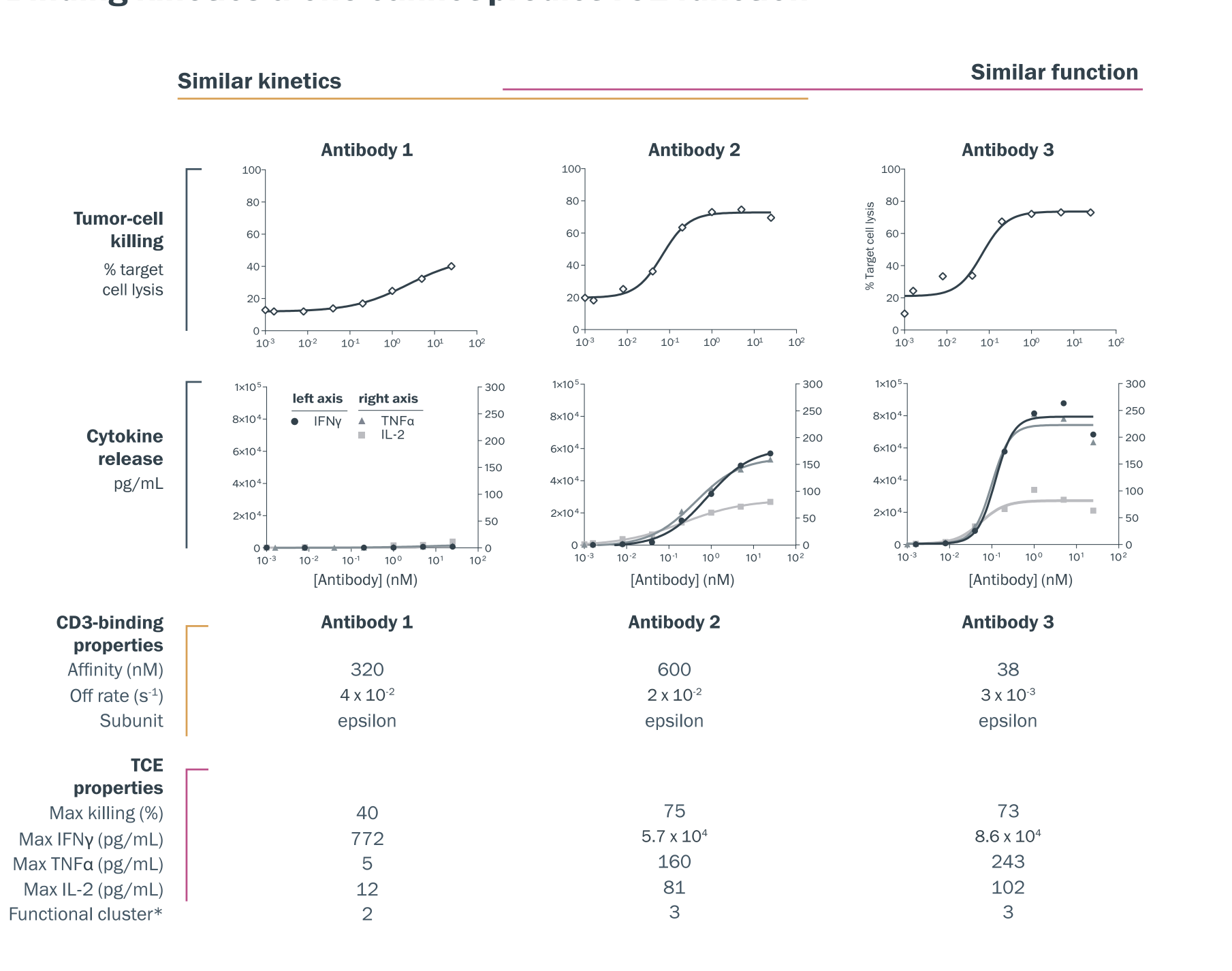


Figure 2. Diverse functional profiles were observed despite similar kinetic parameters. Tumor-cell killing and cytokine release curves for three example TCEs are shown. The CD3-binding properties and functional readouts for each are summarized in the table below. Antibodies 1 and 2 have similar binding kinetics but display distinct functional profiles, while antibodies 2 and 3 have similar functional profiles but different binding kinetics. All three antibodies are from different clonal families.

# Select functionally diverse CD3-binding antibodies

### AbCellera's CD3-binding antibodies can be clustered into 19 groups based on function



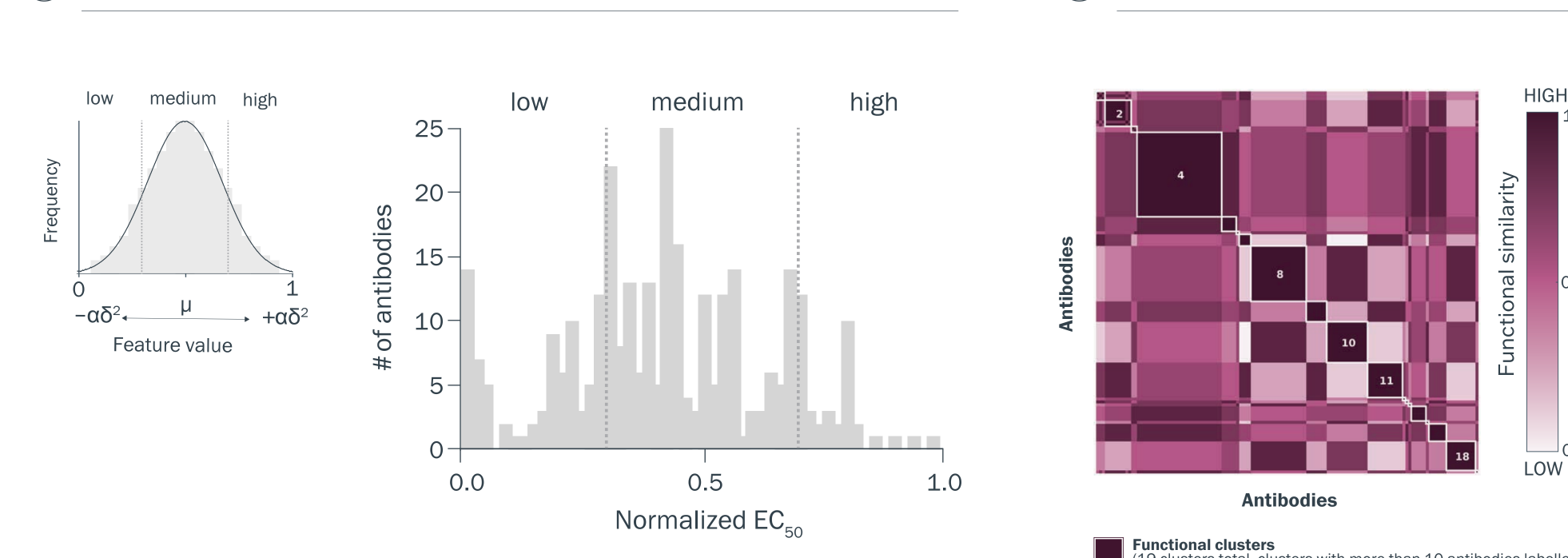
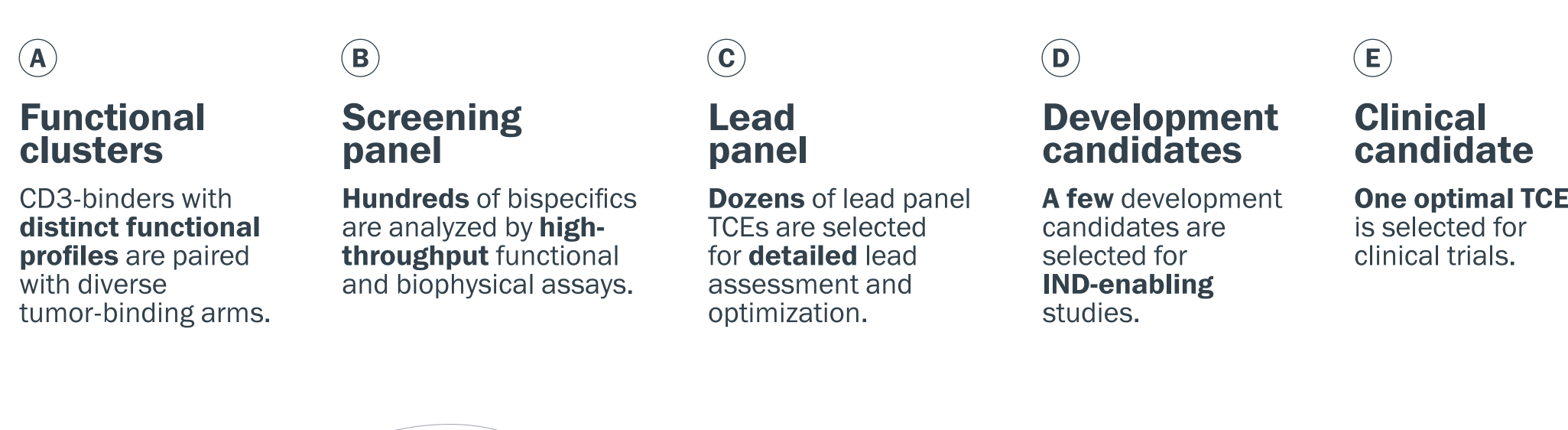


Figure 3. CD3-binding antibodies were clustered based on TCE functional profiles. (A) Tumor-cell killing and cytokine release data were normalized to range (0, 1) and antibodies were categorized as high, medium, or low based on their distance from the mean. Example normalized  $EC_{50}$  data is shown. (B) Antibodies were then clustered based on functional profiles as shown in the white boxes along the diagonal of the heatmap. Example antibodies can be seen in Figure 2, where antibodies 1 and 2 are in different clusters and antibodies 2 and 3 are in the same cluster.

# Functional clusters streamline selection of CD3-binders with relevant diversity



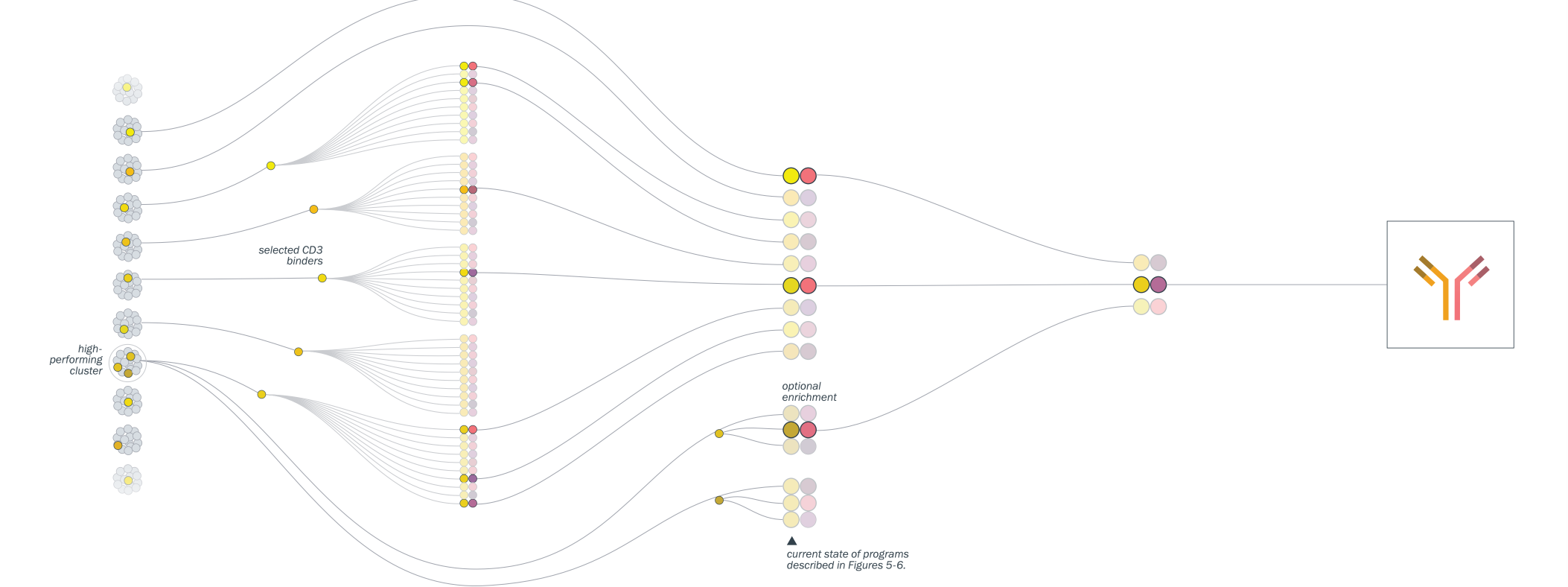


Figure 4. Functional clusters are used to select diverse CD3-binding antibodies for different tumor targets. As shown previously, the same CD3-binder can display different function depending on the tumor-binding arm it is paired with<sup>4</sup>. As a result, functional profiles for different tumor targets cannot be accurately predicted. To address this challenge, we developed a streamlined, function-first approach to identify optimal CD3- and tumor-binding pairs for each target.

(A) We maximize functional diversit upfront by pairing CD3-binders from diverse tumor-binding arms, prioritizing desired properties such as cyno cross-reactivity and

subunit specificities

where needed.

(B) We engineer and bispecifics using high-throughput functional and biophysical

assessments.

CD3-binding arm

the lead panel with

with robust functiona candidates for

IND-enabling studies.

(**D**) We analzye

selected for GMP manufacturing.

(E) Based on

a final clinical

these studies,

# Identify the optimal pair of antibodies for each tumor target

### Functionally diverse CD3-binders were used to engineer TCEs for two tumor targets

A AbCellera TCE program 1: PSMA x CD3

B AbCellera TCE program 2: MAGE-A4 x CD3

<b>)</b> +	<b>&gt;&gt;&gt;</b>	19 »> functional clusters	<b>10</b> CD3-parentals	180+	300+ »> CD3-binders	19 »> functional clusters	<b>19</b> CD3- parentals
<b>)</b> +	<b>&gt;&gt;&gt;</b>	92 »	9	bispecifics	200 »	45 »	6
<b>;</b>		expressed & characterized	PSMA- parentals	engineered	MAGE-A4- binders	expressed & characterized	MAGE-A4- parentals

Figure 5. (A) CD3-binding antibodies selected for our PSMA x CD3 program were prioritized based on cyno cross-reactivity. In addition to having functional diversity, antibodies had different subunit specificities and affinities ranging from 1 nM to 12 µM. We selected nine developable PSMA-binding antibodies from five different epitope bins with affinities ranging from 46 pM to 31 nM.

(B) Cyno cross-reactivity was not required for our MAGE-A4 x CD3 program, enabling selection from a broader range of functional profiles. The 19 selected CD3-binders ranged in affinity from 1 nM to 3 µM. We discovered developable antibodies with high specificity for a MAGE-A4 peptide-MHC<sup>5</sup> and selected six for bispecific engineering.

### High-throughput assessment guided selection of TCEs for further assessment

We selected nine PSMA x CD3 TCEs with functional profiles that span the range of clinical molecules

Lead pane	l antibodies	Kinetic properties		Tumor-cell killing		Cytokine release		
CD3 arm	TAA arm	CD3	PSMA	potency	max	IFNγ	IL-2	TNFα
• CD3-1	PSMA-1	10 -5 —	10 -7	102	100	106	104	104
• CD3-1	PSMA-2	8		000		105		102
O CD3-6	PSMA-6	10 -6 -		101-	<u>@</u> 80-	105-	□ 10³-	103-
<ul><li>CD3-2</li></ul>	PSMA-4	₹	\$\frac{10^{-8}}{\times^{\times}}\$	100-	60   60   60   60   60   60   60   60	(Jm/gd) [hmL] 10 <sup>4</sup> -	(Jm/gd) [Z	(Jm/gd)
OCD3-3	PSMA-3	10 <sup>-7</sup> 10 <sup>-8</sup> —		EC <sub>50</sub> (n)		(pg	8d)	] (pg
• CD3-3	PSMA-5	D3	AM2	© 10 <sup>-1</sup> −	cell	Z 10 <sup>3</sup>		Σ 10 <sup>1</sup> -
<ul><li>CD3-3</li></ul>	PSMA-2	O 10 <sup>-8</sup> □	hPSMA		40 — mpt	= XE 102	Max [IL	
• CD3-4	PSMA-2		10-9-	10 -2 —		$\stackrel{\text{xe}}{\succeq}$ 10 <sup>2</sup>	∑ 10¹−	× 10°− × × × × × × × × × × × × × × × × × × ×
OCD3-5	PSMA-3	10 -9 —	-	10-3	<u>×</u> 20−	101		10 -1 —
clinical molecules		10 -10	10 -10	10 -4 —		100	100	10-2
• TNB-585		10 10	10	10 '	0 —	10 ° –	10°—	10 -2 —
△MG-160	)							

#### We selected 12 MAGE-A4 x CD3 TCEs with desired functional profiles and peptide-MHC specificity



Figure 6. High-throughput assessment of TCEs enabled selection of molecules for further assessment. (A) 180+ PSMA x CD3 TCEs were assessed and compared to two clinical-stage molecules (TNB-585<sup>6</sup> and AMG-160<sup>7</sup>) in a high-throughput assay using unactivated human T cells incubated with PSMA-expressing LNCaP cells at a ratio of 10:1 for 48 hours. Nine selected antibodies are highlighted. (B) 200+ 1x1 MAGE-A4 x CD3 T-cell engagers were assessed and compared to Molecule R<sup>8</sup> (057D03 paired with V9 in a 2x1 bispecific format) in a highthroughput assay using unactivated human T cells incubated with wild-type or MAGE-A4 knockout A375 cells at a ratio of 10:1 for 48 hours. Twelve selected antibodies are highlighted.



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<sup>\*</sup>Functional clustering is described in Figure 3.