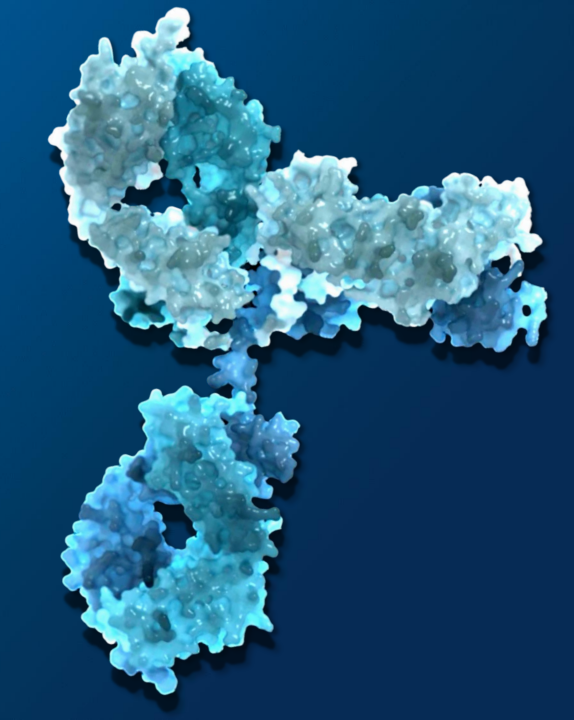


Sutro Biopharma

July 2025 NASDAQ: STRO



Forward-Looking Statements

This presentation and the accompanying oral presentation contain "forward-looking" statements that are based on our management's beliefs and assumptions and on information currently available to management. Forward-looking statements include all statements other than statements of historical fact contained in this presentation, including information concerning our future financial performance; business plans and objectives; anticipated preclinical and clinical development activities, including enrollment and site activation; timing of announcements of clinical results, trial initiation, and regulatory filings; outcome of regulatory decisions; and our expectations about our cash runway; potential benefits of our product candidates and platform; potential expansion into other indications and combinations, including the timing and development activities related to such expansion; potential growth opportunities, financing plans, potential future milestone and royalty payments, competitive position, industry environment and potential market opportunities for our product candidates.

Forward-looking statements are subject to known and unknown risks, uncertainties, assumptions and other factors, including risks and uncertainties related to our cash forecasts, our and our collaborators' ability to advance our product candidates, the receipt, feedback and timing of potential regulatory submissions, designations, approvals and commercialization of product candidates and the design, timing and results of preclinical and clinical trials and our ability to fund development activities and achieve development goals. It is not possible for our management to predict all risks, nor can we assess the impact of all factors on our business or the extent to which any factor, or combination of factors, may cause actual results to differ materially from those contained in any forward-looking statements we may make. These factors, together with those that may be described in greater detail under the heading "Risk Factors" contained in our most recent Annual Report on Form 10-K, Quarterly Report on Form 10-Q and other reports the company files from time to time with the Securities and Exchange Commission, may cause our actual results, performance or achievements to differ materially and adversely from those anticipated or implied by our forward-looking statements.

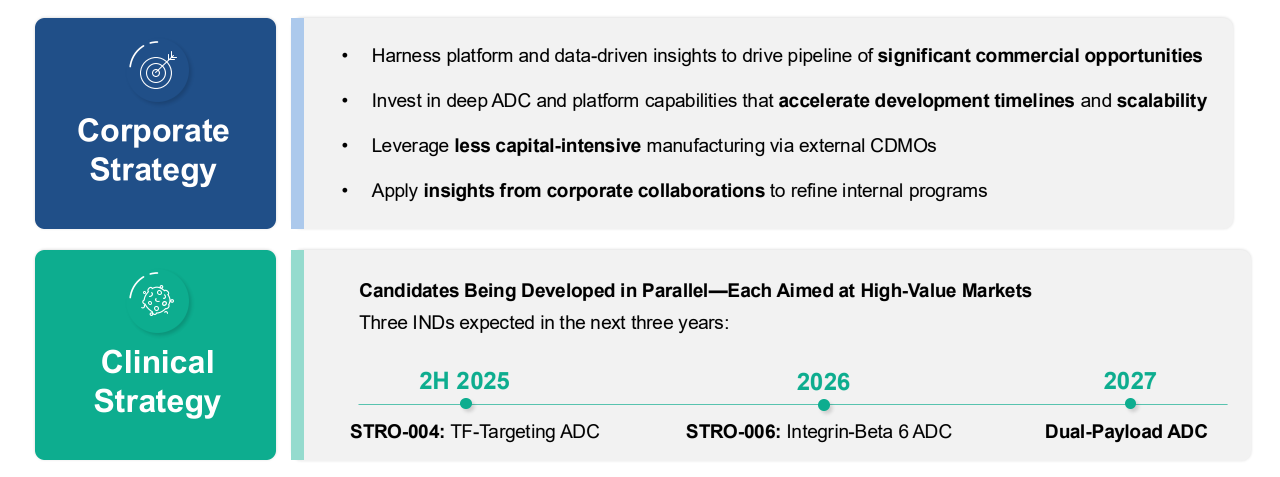
You should not rely upon forward-looking statements as predictions of future events. Although our management believes that the expectations reflected in our forward-looking statements are reasonable, we cannot guarantee that the future results, levels of activity, performance or events and circumstances described in the forward-looking statements will be achieved or occur. Moreover, neither we nor our management assume responsibility for the accuracy and completeness of the forward-looking statements. We undertake no obligation to publicly update any forward-looking statements for any reason after the date of this presentation to conform these statements to actual results or to changes in our expectations, except as required by law.

This presentation also contains estimates and other statistical data made by independent parties and by us relating to market size and growth and other data about our industry. This data involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. In addition, projections, assumptions, and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk.



Next-Generation ADCs Designed to Address Critical Treatment Gaps in Oncology

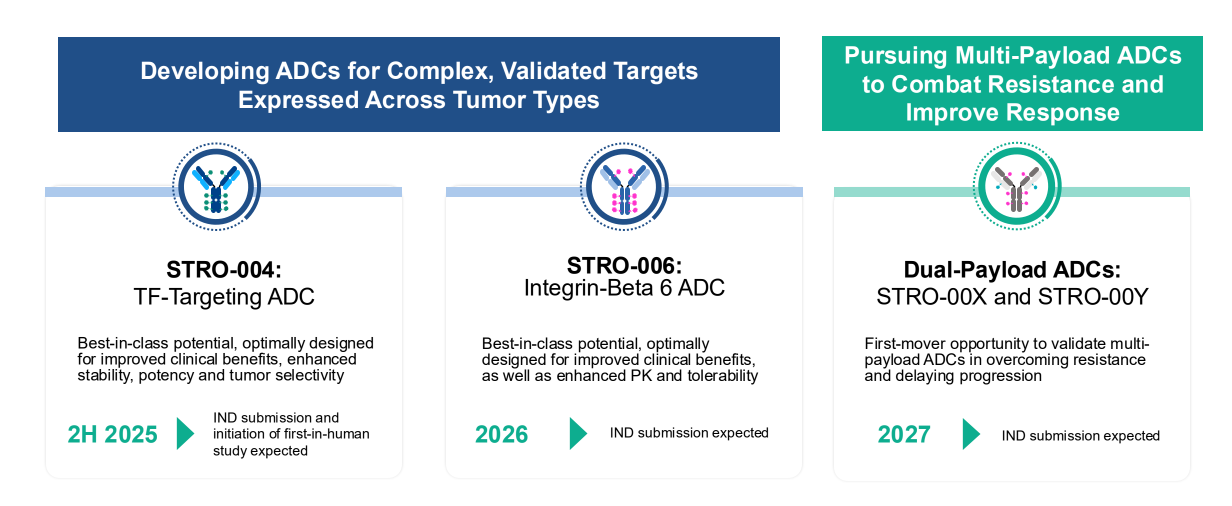
Well-capitalized with runway into early 2027; partnership milestone payments expected within 12 months





ADCs Designed to Engage Hard-to-Reach Targets with Single- and Dual-Payloads

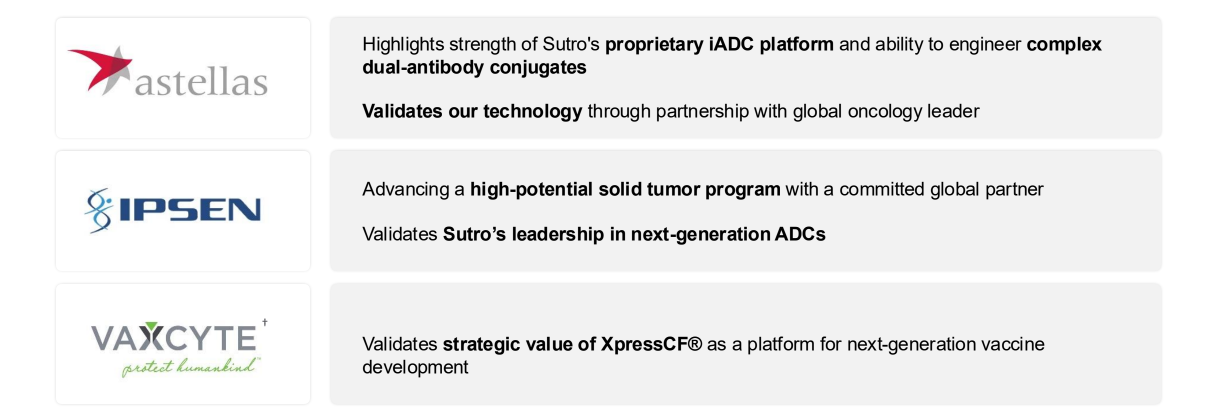
Pursuing complex targets enabled by our proprietary technology, differentiating from conventional ADCs



TF – Tissue Factor; IND – Investigational New Drug; PK – Pharmacokinetics

Unlocking Capital and Expertise Through Strategic XpressCF® Partnerships*

Partnerships provide up to \$2B in potential milestones and royalties



*Vaxcyte is advancing vaccines using advanced chemistry and the XpressCF® platform, exclusively licensed from Sutro †Blackstone purchased 4% royalties on potential future net sales of Vaxcyte's PCV products; Potential future payments to Sutro

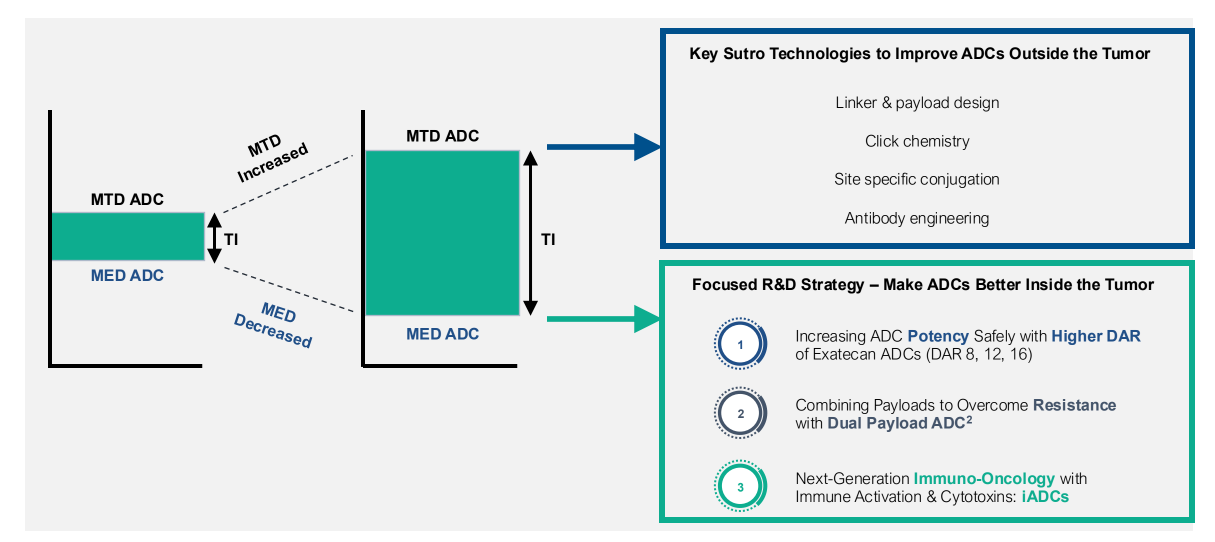




A Leader in Next-Gen ADCs

Pursuing Complex, High-Value Targets Enabled by XpressCF[®]

Wider Therapeutic Index Achieved with Sutro's Cell-Free ADC Platform



Adapted from Gerber et al, mAbs, 2023 MTD – Maximum tolerated dose; MED – Minimum effective dose



Our XpressCF[®] Platform has Unique ADC Performance Capabilities Over Other Topo1 ADC Platforms

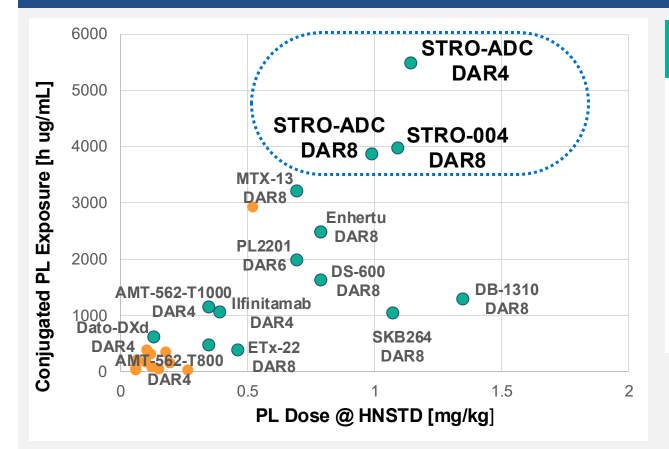
| | DAR>8 | Beta-Glu Linker | ADC ² / Dual LPs | iADC/ iSAC | Site Specific | Fc Silent | Bispecific | HT Screening |
|--------------------|------------|--------------------|--------------------------------|---------------|------------------|------------|------------|-----------------|
| SUTR: BIOPHARMA | \bigcirc | \bigcirc | \bigcirc | \bigcirc | \bigcirc | \bigcirc | \bigcirc | \bigcirc |
| Abbvie | | | | \oslash | | \oslash | \oslash | |
| AstraZeneca | | | | | \oslash | \oslash | \bigcirc | |
| Daiichi Sankyo | | | | | | | | |
| Dualitybio | | | | \oslash | | \oslash | \oslash | |
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| Medilink | | | | | | | | |
| Merck KGaA | | \oslash | | | | | \bigcirc | |
| Pfizer | | \oslash | | \oslash | | | | |

LP – Linker payloads; iSAC – Immune stimulating antibody conjugate; HT – High throughput; Comparison of Topo1i ADC platforms (selected)





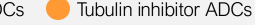
Comparison of Exposure Levels in NHPs at Highest Non-Severely Toxic Dose (HNSTD) Levels in DAR Equivalents

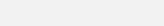


Why does it matter?

- For ADCs, exposure drives efficacy
- Based on PK data, our exatecan ADCs are positioned to be differentiated on safety and efficacy versus on-market ADCs

Exatecan/Topo1i ADCs





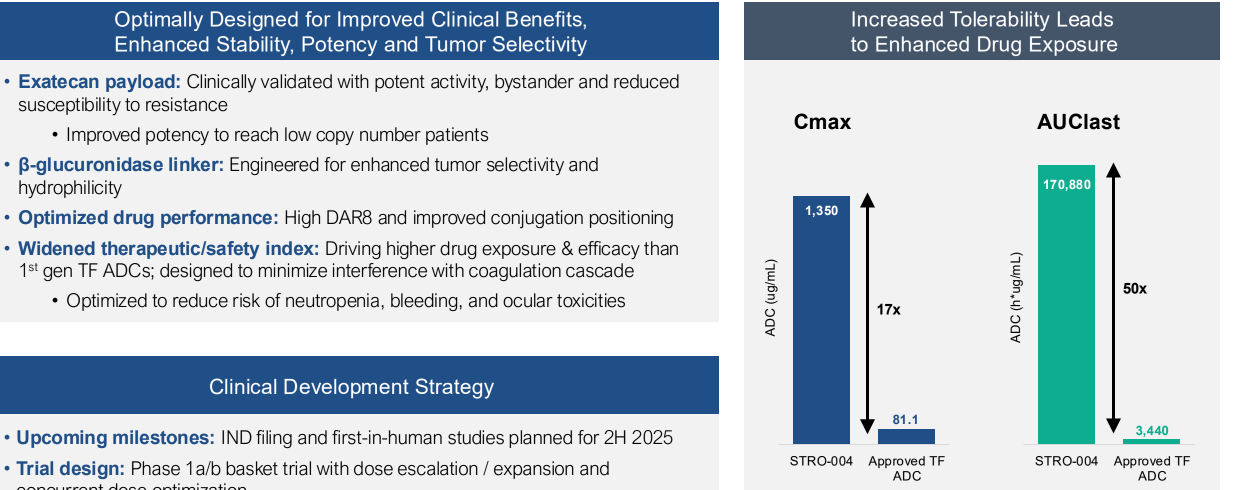




STRO-004

Potential Best-in-Class Exatecan ADC Targeting TF

STRO-004: Next-Generation TF-Targeting Exatecan/Topo1 ADC with Enhanced **Therapeutic Potential**



| 1 st gen TF ADCs; designed to minimize interference with coagulation cascade | |
|--|---|
| Optimized to reduce risk of neutropenia, bleeding, and ocular toxicities | |
| | 2 |

Clinical Development Strategy

- Upcoming milestones: IND filing and first-in-human studies planned for 2H 2025
- Trial design: Phase 1a/b basket trial with dose escalation / expansion and concurrent dose optimization

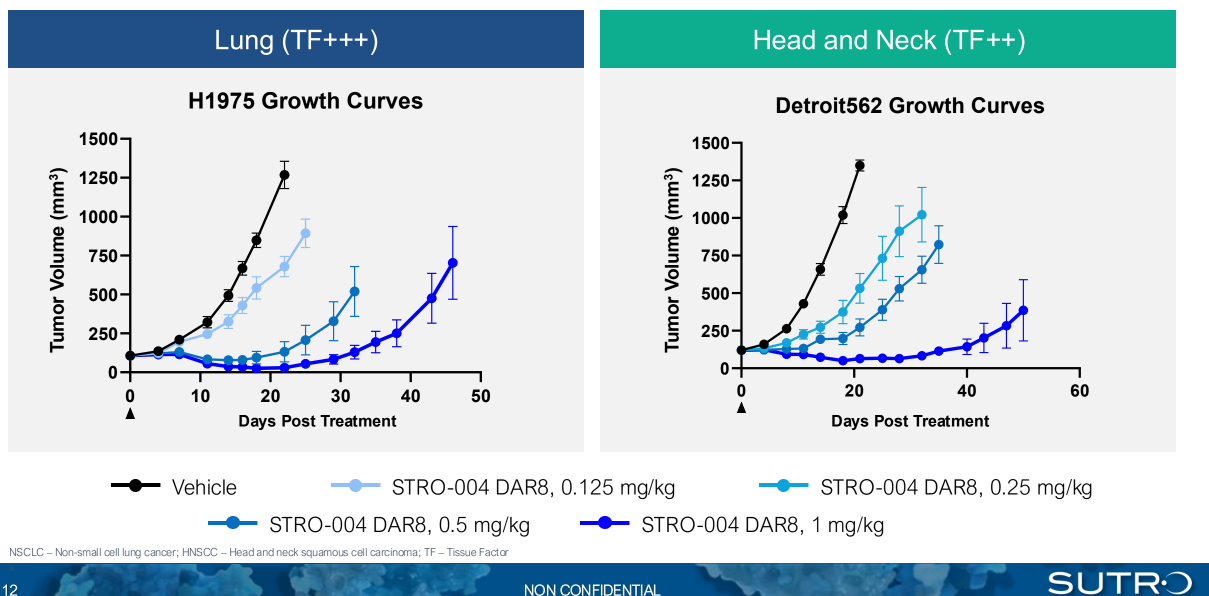
TF - Tissue factor

susceptibility to resistance

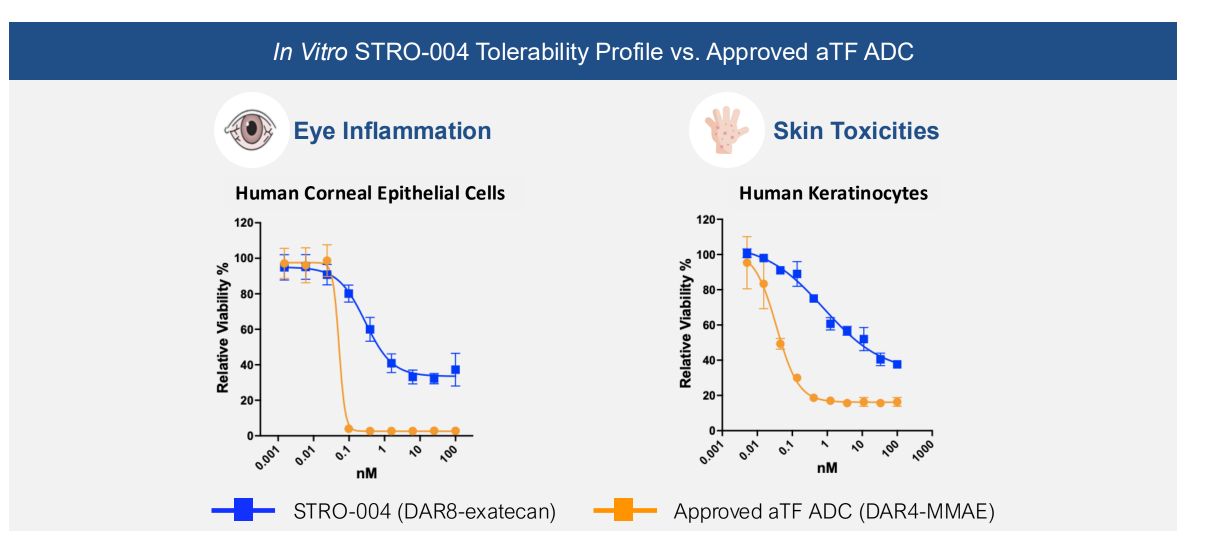
hydrophilicity



STRO-004 DAR8 Exatecan Achieved Sustained Tumor Regressions in Xenograft Models of NSCLC and HNSCC at Low Doses



STRO-004 Demonstrated Reduced Platform and On-target Toxicity Due to Site Specific Conjugation and Beta Glu Linker-Payload Technology





STRO-004: Well-Tolerated in NHP up to 50 mg/kg

Objective:

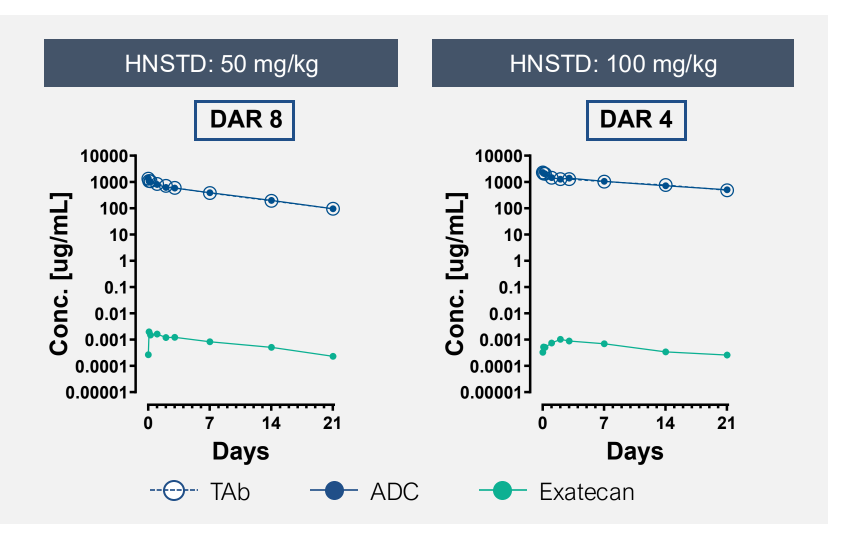
Compare nonclinical safety of DAR8 and DAR4 TF exatecan-ADC

Study:

Dosed twice, three weeks apart, payload-matched doses

Findings:

- DAR8 and DAR4 ADCs were welltolerated up to 50 and 100 mg/kg, respectively
- DAR8 50 mg/kg $t_{1/2}$ of 6.9 days
- No evidence of eye toxicity
- Mild skin toxicity observed in both DAR8 and DAR4

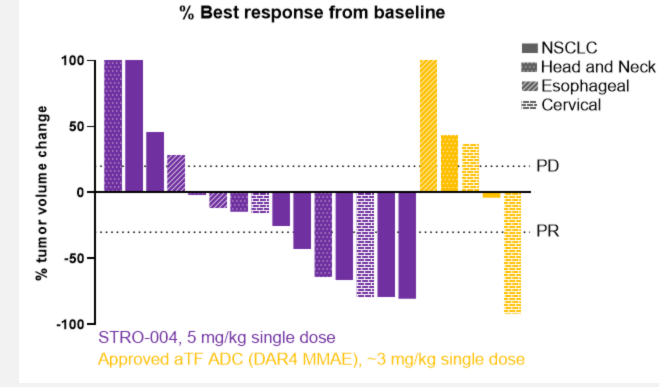


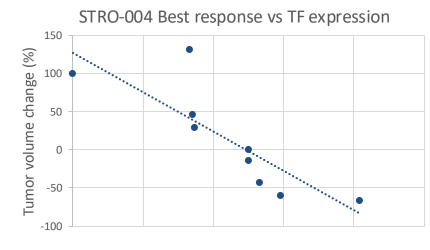
NHP – Non-human primate; TF – Tissue factor; TAb – Total antibody; HNSTD – Highest non-severely toxic dose



STRO-004 Shows Promising Anti-tumor Activity In TF Positive PDX Models of HNSCC, NSCLC, Esophageal, and Cervical Cancer

> 50% of Tumors Respond to STRO-004 at Low Dose



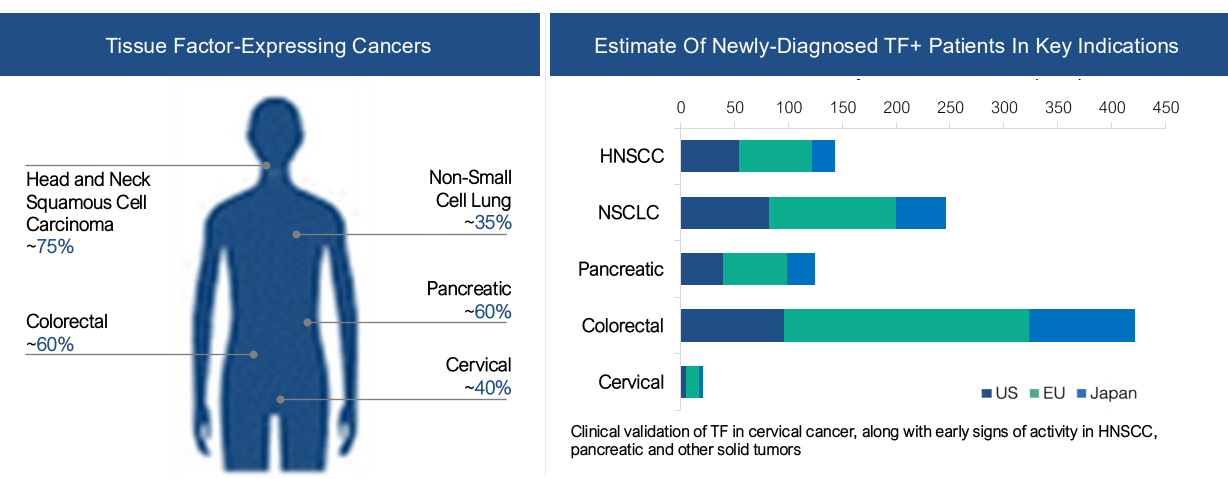


TF expression



TF is Broadly Expressed Across Multiple Solid Tumor Indications with High Unmet Need, Presenting Opportunity for Pan-Tumor Targeting

TF expression has been associated with poor disease prognosis and increased metastatic properties



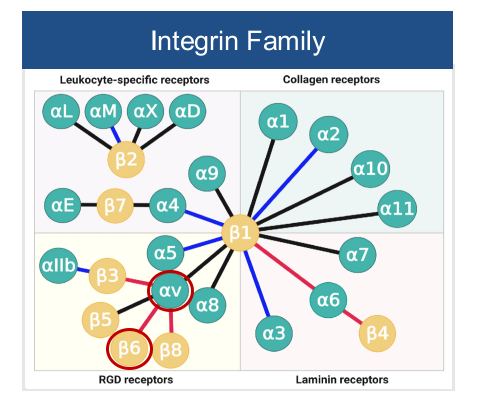
TF expression assumptions are based on a weighted average of tissue factor expression as reported in publicly available literature and triangulated with internal Sutro data on file. Does not account for subsets of tumor types (e.g., MSS vs. MSI in colorectal cancer). Sources for incidence across geographies: 1. Cancer Statistics, 2025 from CA: A Cancer Journal for Clinicians (Siegel RL et al., ACS Journal, Jan 2025), which leverages SEER data: https://acsjournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer Information System (ECIS), EU-27+EFTA data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer Information System (ECIS), EU-27+EFTA data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer Information System (ECIS), EU-27+EFTA data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer Information System (ECIS), EU-27+EFTA data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer registry incidence data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer registry incidence data, accessed Feb 2025: https://acsiournals.onlinelibrary.wiley.com/doi/epdf/10.3322/caac.21871. European Cancer registry incidence data, accessed



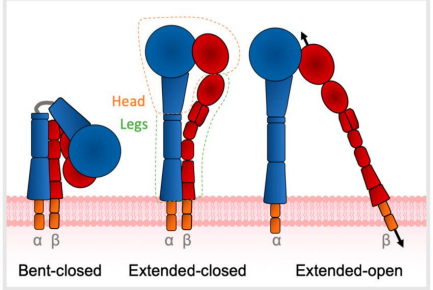


STRO-006 (Integrin ανβ6 / ITGB6) Potential Best-in-Class Integrin-Beta 6 ADC

Complex ITGB6 Biology Requires Advanced Protein Engineering Capabilities







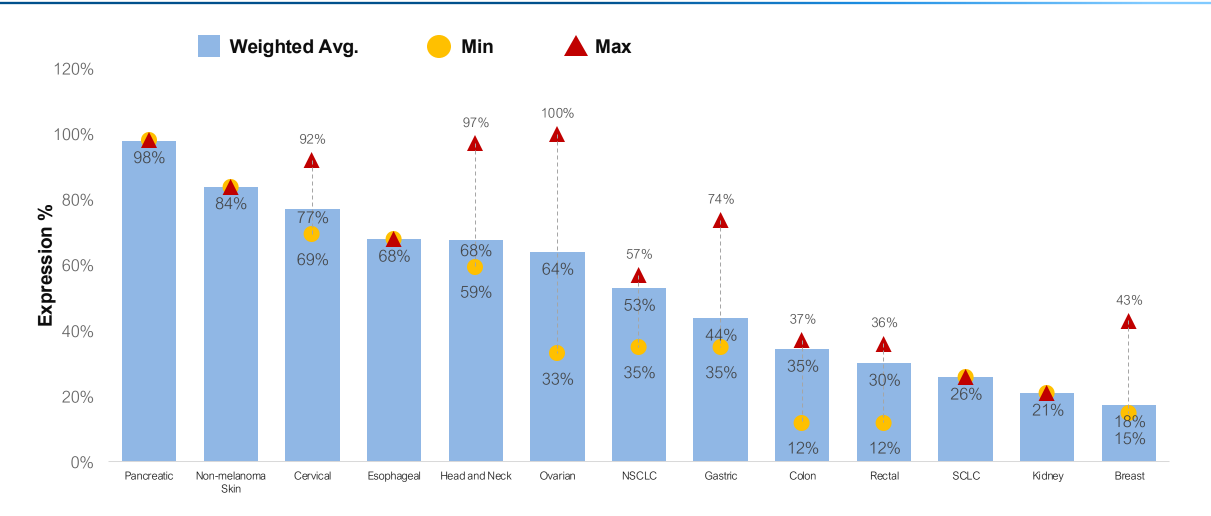
- ITGB6 belongs to integrin family of adhesion proteins, heterodimerizes with alpha-v (αvβ6)
- Exists in multiple conformations makes it a challenging protein to target







ITGB6 is Widely Expressed Across Multiple Solid Tumors

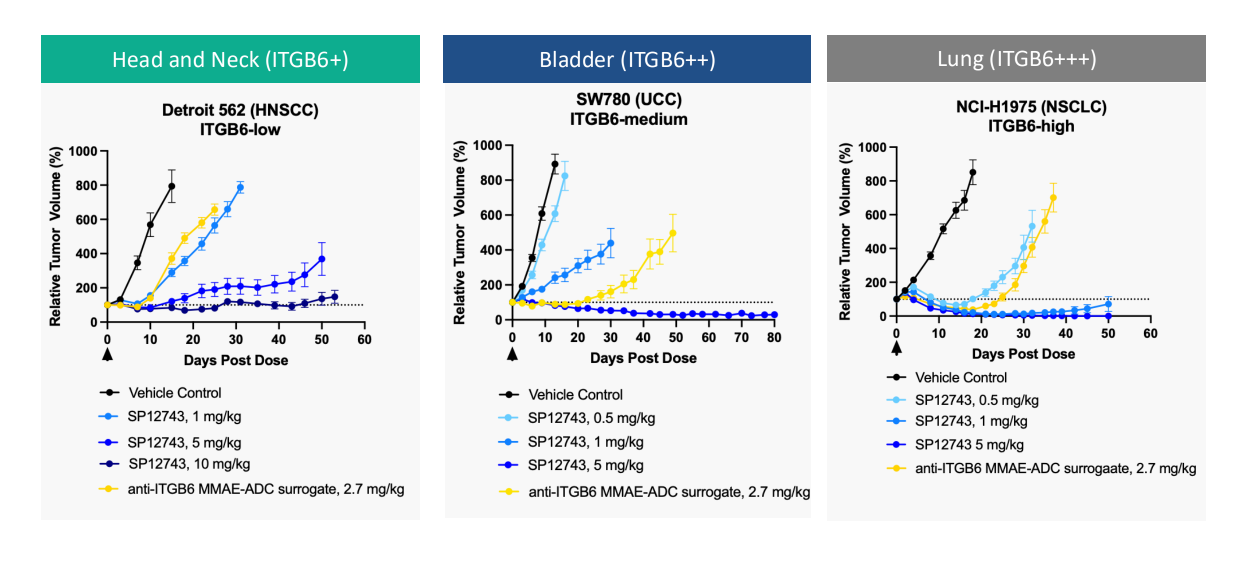


ITG66 expression assumptions are based on a weighted average of expression as reported in publicly available literature and triangulated with internal Sutro data on file. Criteria for positivity differs across studies, overall positive staining/overexpression % is used

ITGB6 – Integrin beta 6

SUTRO

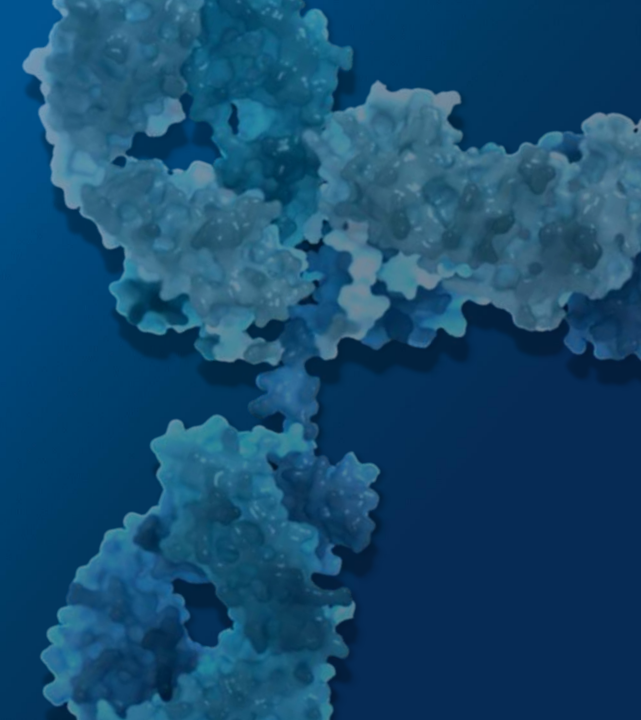
STRO-006 Has Superior Anti-Tumor Activity Compared to First-Generation ITGB6 ADCs at Clinically Relevant Dose Levels







Emerging Leader in Dual Payload ADCs



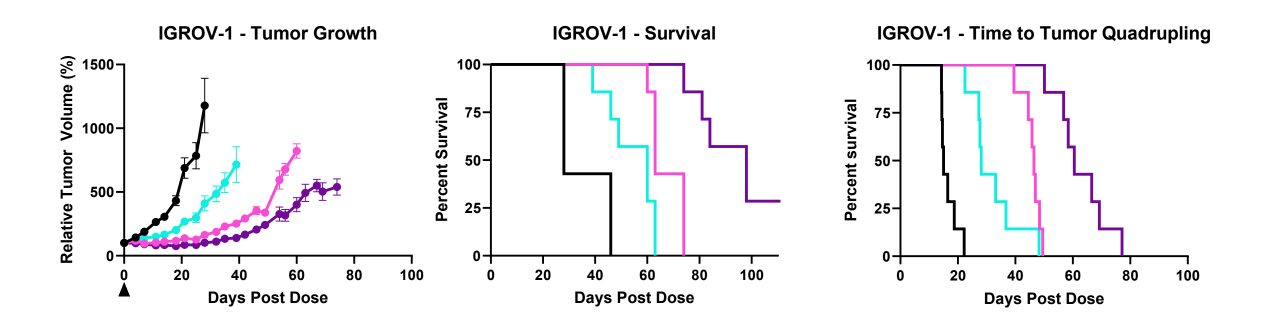
Dual-Payload ADCs: Innovative Method for Delivering Targeted Combination Therapy to Overcome Resistance and Delay Progression

| | ADC + Chemo | ADC + ADC | Dual Payload ADC | |
|---|--|---|---|---|
| | +• | | | Potential benefits of a dual- payload ADCs for targeted combination therapy |
| Safety (Compared to small molecule combinations) | Greater SAEs reported for ADC + chemo vs ADC ^{1,2} | | | Improved tolerability Through reduced systemic payload exposure |
| Efficacy (Control over delivery of drugs to same cell) | | Binding competition impacts efficiency of delivery (for same target) ³ | | Greater control over delivery Both payloads delivered to the same cell at the same time |
| Regulatory Simplicity | | | | Reduced clinical complexity Single agent regulatory data package, standard monotherapy dose escalation design |
| Combination Study Simplicity | | | Combo with modalities such as ICIs that have shown improved outcomes with ADCs ⁴ | Reduced cost Potential for combination benefit in one product |

Sources: 1. PMID: 27052654; 2. PMID: 23020162; 3. PMID: 34112795; 4. PMID: 36041086; ICI – Immune checkpoint inhibitor; TGI – Tumor growth inhibition; SAE – Severe adverse event



Dual-Payload ADC (Topo1i + anti-Tubulin) Displays Enhanced *In Vivo* Efficacy in Ovarian Cancer



Vehicle control Trastuzumab DAR4 MTI ADC (5 mg/kg) Trastuzumab DAR8 Topo1i ADC (5 mg/kg) Trastuzumab DAR8 Topo1i + DAR4 MTI dpADC (5 mg/kg)



iADC: Dual-Payload ADC Combining Tumor-Targeted Delivery of a Cytotoxin and Immune Stimulator

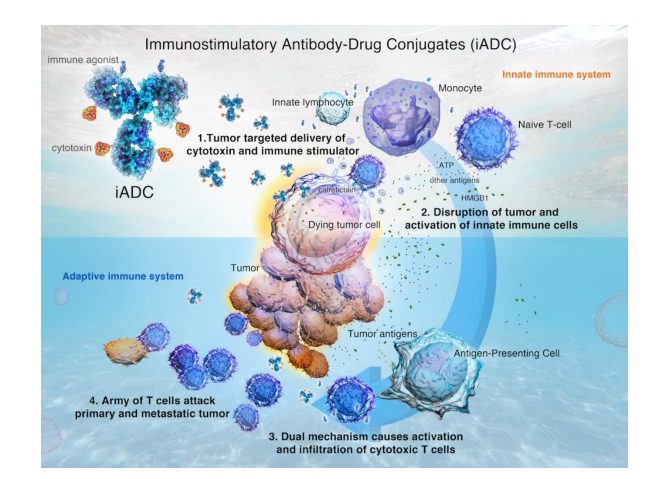
Strategic Partnership with Astellas to Deliver New Treatment Options for Cold Tumors and Patients Unresponsive to Existing Cancer Immunotherapies



Combining a cytotoxin and immune modulator gives potential to:

- Act alone by stimulating the immune system and priming new populations of immune cells
- **Synergize with other immune therapies** that remove inhibitory signals on the immune system (e.g. checkpoint inhibitors)
- Address hard-to-treat cancers by activating a robust anti-tumor immune response

Two programs ongoing, with one in IND-enabling toxicology study





Novel Mechanism of Action Differentiates iADCs from Other Immunotherapies

| Sutro iADCs bridge | | | | | | | |
|--|--|--|---------------------------------|-------------------------|---|-----------------|---------------------------|
| adaptive immunity to provide broad protection in a single molecule | | Sutro iADC | STING / TLR | ISAC | PD-1 / PDL-1 | CAR-T Cells | Vaccine |
| | Molecule | Targeted and homogeneous | Chemo | Mixed ADC | Ab | Biologic | Biologic |
| | Opportunity: risk | Combine ICD with innate agonists (TLR, STING, etc.) | Non-targeted, issues with TI | Requires Fc effector | Limited tumor types, small tumors | Safety concerns | Ag selection challenge |
| | FcγR meditated uptake into myeloid | | | × | | | |
| | Direct tumor cell killing | ~ | | | | \checkmark | |
| Mechanisms | Tumor antigen presentation | ~ | | \checkmark | | | ~ |
| to achieve anti-tumor immunity | Priming and activation of antigen presenting cells | ~ | ~ | ~ | | | ~ |
| | T-cell recruitment to tumor | ~ | \checkmark | \checkmark | \checkmark | \checkmark | |

STING – Stimulator of interferon genes; TLR- Toll-like receptor; Immunogenic cell death X – Undesirable



XpressCF[®] Enables Development of Differentiated Dual-Payload ADCs, Leveraging **Unique Combinations of Validated Targets**

Dual-Payload ADCs Have the Potential to Become Future Standard of Care Topo1 x Tubulin Topo1 x IO Topo1 x PARPi Selected Indications Selected Indications Selected Indications Breast "Hot" Tumors • NSCLC (EGFR wild type & mutant) Ovarian Breast "Cold" Tumors ٠ • Prostate Bladder ٠ Pancreas Head & Neck ٠ ٠ **Clinical Evidence for Success Clinical Evidence for Success Clinical Evidence for Success** Improved clinical activity when Based on approved PARPis in BRCA1/2 Activity of STING agonists after mutant tumors, and early clinical activity intertumoral administration in combining Topo1 and Tubulin ADCs when combining Topo1 ADC with PARPi solid tumors small molecule

NSCLC - Non-small cell lung cancer



| Sutro's Wholly-Owned Programs | | | | | | | |
|---|---|--|--|--|--|--|--|
| STRO-004 Exatecan ADC Targeting Tissue Factor | 2H 2025: IND filing and first-in-human studies planned 2026: Phase 1a/b dose escalation data expected 2027: Phase 1a/b dose expansion data expected (initial response data anticipated 1H 2027) | | | | | | |
| STRO-006 Integrin-Beta 6 ADC | Mid-2026: IND filing planned 2027: Dose escalation data expected | | | | | | |
| Dual-Payload | 2027: STRO-00X IND filing planned | | | | | | |
| Corporate Updates | Year-End 2025: Expected to complete restructuring, divestiture of manufacturing facility, potential platform collaboration deal | | | | | | |

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Pipeline of Next-Generation ADCs Designed to Engage Hard-to-Reach Targets with Single- and Dual-Payloads

| PROGRAM | MODALITY/TARGET | INDICATION | DISCOVERY | PRECLINICAL | PHASE 1/1B | PHASE 2 | PHASE 3/ REGISTRATIONAL | WORLDWIDE OR GEOGRAPHIC PARTNER |
|----------|-------------------|--------------|-----------|-------------|------------|----------|----------------------------|---------------------------------------|
| WHOLLY | -OWNED PROG | RAMS | | | | | | |
| STRO-004 | Tissue Factor ADC | Solid Tumors | | • | | | | |
| STRO-006 | Integrin αvβ6 | Solid Tumors | | -• | | | | |
| STRO-00X | Dual Payload ADC | Solid Tumors | • | | | | | |
| STRO-00Y | Dual Payload ADC | Solid Tumors | • | | | | | |
| PARTNE | R PROGRAMS | • | | | | - Sandra | (| |
| | | Invasive | | Add | | | | |

| VAX-24 | 24-Valent Conjugate Vaccine | Invasive Pneumococcal Disease | να <mark>χ</mark> сүте |
|-------------------------|-----------------------------------|--|----------------------------|
| VAX-31 | 31-Valent Conjugate Vaccine | Invasive Pneumococcal Disease | größeit humankind |
| STRO-003 | ROR1 ADC | Solid Tumors & Hematological Cancers | SIPSEN |
| Undisclosed Programs | Immunostimulatory ADCs (iADCs) | Cancers | Astellas |



Sutro Team Comprised of Industry Leaders



Jane Chung, RPh Chief Executive Officer



Greg Chow, MBA Chief Financial Officer



Hans-Peter Gerber, PhD Chief Scientific Officer



David Pauling, JD, MA Chief Administrative Officer and General Counsel



Barbara Leyman, PhD Chief Business Officer



Venkatesh Srinivasan, PhD Chief Technical Operations Officer

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