

Unlocking the Full Potential of the Immune System Against Cancer

Corporate Presentation

April 9, 2024



This presentation contains forward-looking statements, which are subject to numerous risks and uncertainties, which could cause actual results to differ materially from those anticipated. There can be no guarantee that (i) the results of pre-clinical work and prior clinical trials will be predictive of the results of the clinical trials currently under way, (ii) regulatory authorities will agree with the Company's further development plans for its therapies, or (iii) the Company will find development and commercialization partners for its therapies in a timely manner and on satisfactory terms and conditions, if at all. The occurrence of any of these risks could have a significant negative outcome for the Company's activities, perspectives, financial situation, results and development.

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Innovative Clinical-Stage Immunotherapy Portfolio Based on Viral Vectors

Cutting-edge individualized neoantigen cancer vaccine (TG4050)

- Proof of principle obtained in randomized
 Phase I study (H&N adjuvant)
- Randomized Phase II trial to start in Q2 2024

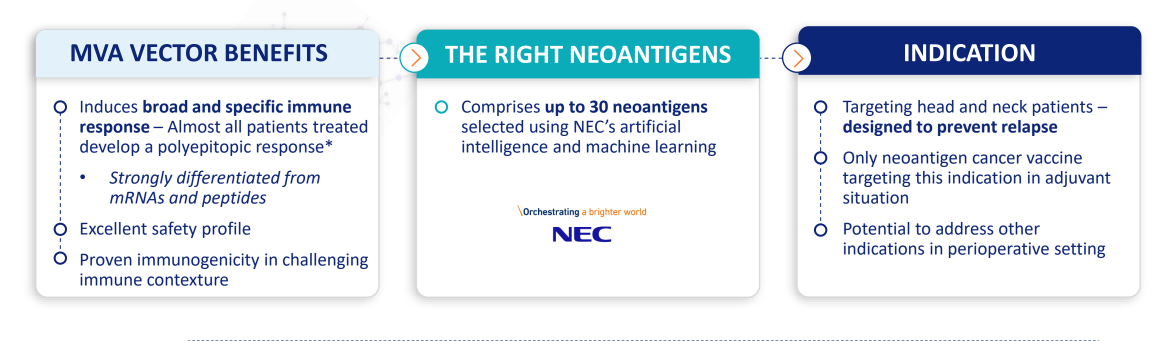
Additional immuno-oncology programs with clinical proof of principle

- Shared antigens vaccines (HPV16)
- Oncolytic viruses

Significant --- value creation catalysts expected in 2024

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TG4050 – A Novel Individualized Cancer Immunotherapy





Building upon proof of principle: Randomized Phase II trial to begin in Q2 2024 based on promising Phase I data

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Our pipeline – poised to deliver important data

Product	Indication	Collaboration	Discovery	Phase I	Phase II	Key upcoming catalysts		
INDIVIDUALIZED NEOANTIGEN CANCER VACCINES								
TG4050 myvac	Head and neck cancer (adjuvant)	\Orchestrating a brighter world	•		•	24-month median follow up (H2 2024) Ph. II trial to start (Q2 2024)		
	Other indication					Additional Ph. I trial to start (2025)		
SHARED ANTIGENS CANCER VACCINES								
TG4001	Anogenital HPV+ cancers					Randomized Phase II trial results (H2 2024)		
Internal myvac	Shared driver mutations							
ONCOLYTIC VIRUSES (OVs)								
TG6050 invir	Lung cancer (IV*)					First data (H2 2024)		
BT-001 invir	Solid tumors (IT*)	BioInvent				First data in combination with pembrolizumab (H2 2024)		
Internal	Synthetic OV (IV*)							





Cancer Therapeutic Vaccines

Focused on delivering the promise of individualized cancer vaccine

myvac[®] - TG4050 | Combines Unique Know How and Expertise

ransgene

MVA viral vector: a powerful platform for vaccine development

Strongly immunogenic vector

- Demonstrated capability to express complex antigen structures and have them presented by APCs
- Ability to elicit strong, durable and specific immune response
- Established safety profile

Optimal neoantigen display

- VacDesignR[™] for optimal design of the recombinant cassettes
- Selection of best promoter sequences



one patient • one genomeone vaccine

Artificial Intelligence to identify up to 30 potent neoantigens

- NEC's machine learning environment based on multiple parameters to classify most immunogenic neoantigens from whole tumor genome analysis*
 - Takes in account multiple parameters
 - NEC covers 50% of the clinical development costs of TG4050 in head and neck cancer

NEC

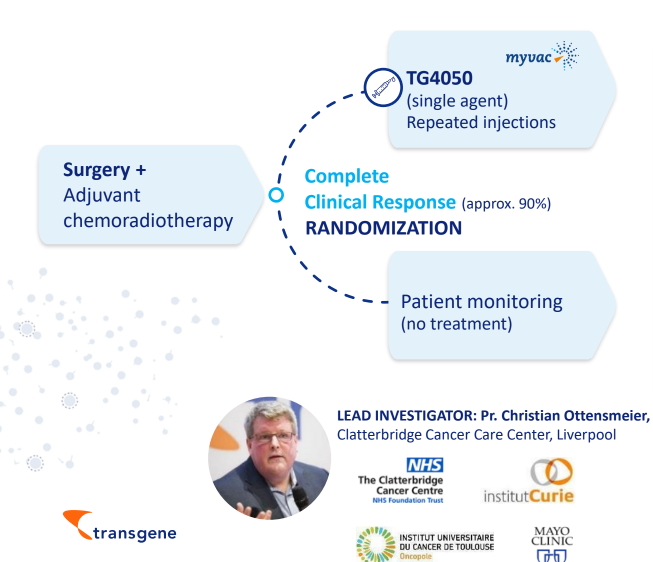


Technology well suited to demonstrate benefit in minimal residual/molecular disease

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*Source: Mallone et al., "Performance of neoantigen prediction for the design of TG4050, a patient-specific neoantigen cancer vaccine". AACR, June 2020, Poster presentation

TG4050 | HPV-Negative Head and Neck Cancer - Trial after Surgery and Adjuvant Therapy

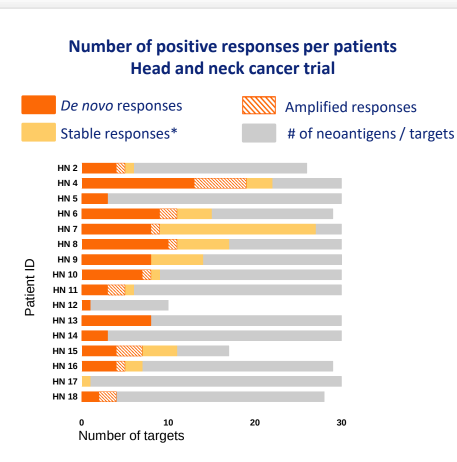


Need to prevent or delay relapse Clinical situation where checkpoint blockers have failed (ie. KN412, Javelin 100) Approx. 30% patients relapse within 24 months after adjuvant therapy*

Randomized Phase I trial – Promising initial immunological and clinical data presented at AACR 2024 32 patients (NCT: 04183166)

- → All treated patients remaining disease-free
- ➔ Trial extended to Phase I/II, patient inclusions to restart in Q2 2024

TG4050 | Generates and/or Expands Tumor Specific T Cells



Despite low mutational burden, immunogenic targets could be selected for all patients

CD4+ or CD8+ responses were detected in all but one patients vaccinated patients

80% of immunoreactivities detected after vaccination were not detectable at baseline.

*Immunoreactive T-cells detected at baseline but not amplified by vaccine

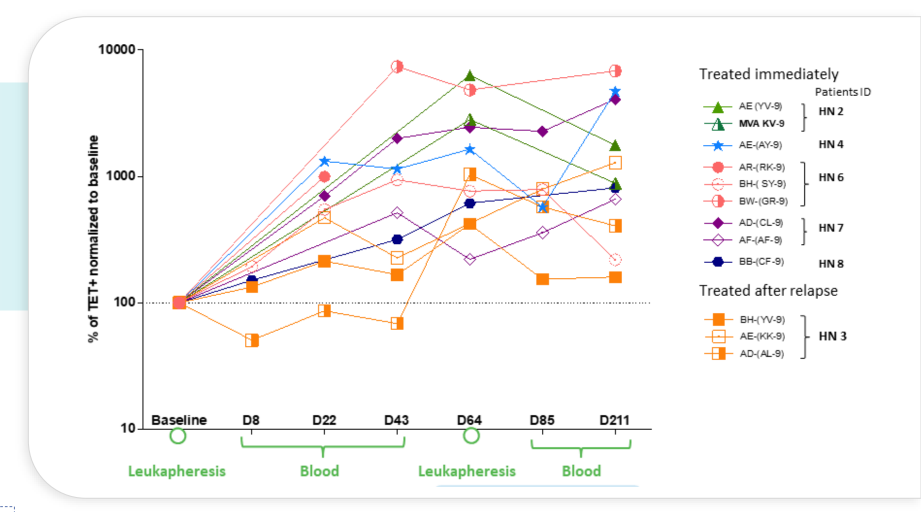
American Association for Cancer Research *Source: A. Lalanne *et al.*, "Personalized vaccine TG4050 induces polyepitopic immune responses against private neoantigens in resected HPV negative head and neck cancers." <u>AACR 2024</u>, April 10, Poster presentation



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TG4050 | Persistent Specific Cellular Response Following Vaccination

Patients display **persistent specific CD4+ and CD8+ responses** against multiple selected targets, 7 months after treatment induction



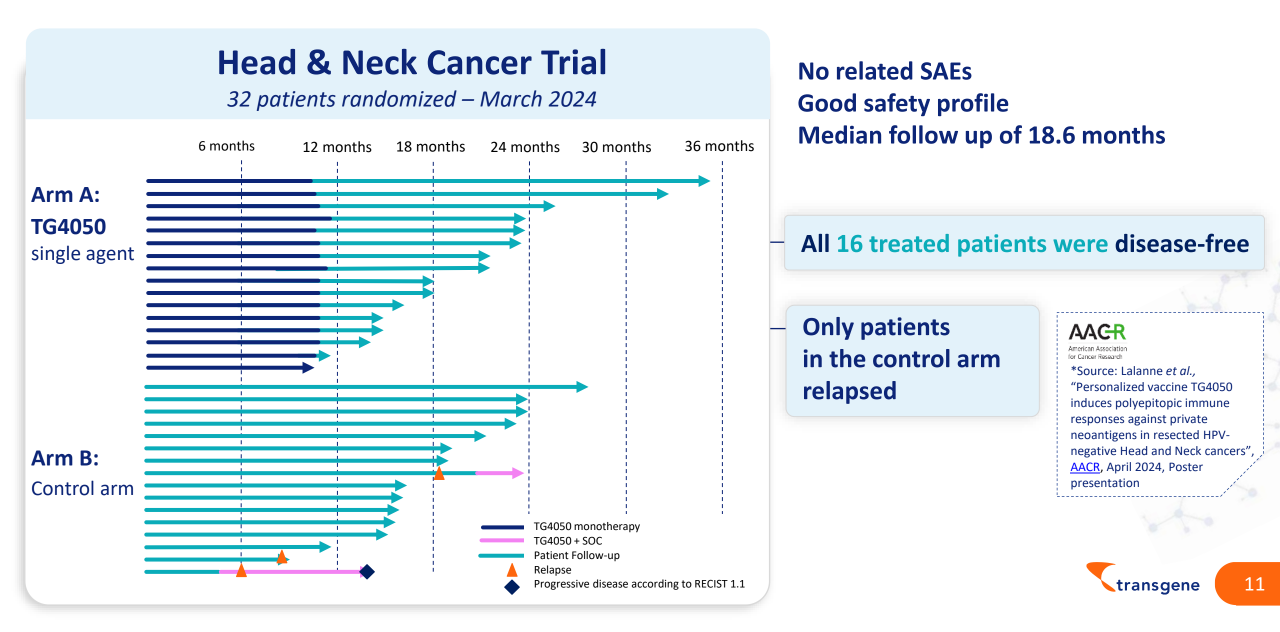


Source: A. Lalanne *et al.*, "Personalized vaccine TG4050 induces polyepitopic immune responses against private neoantigens in resected HPV negative head and neck cancers." <u>AACR 2024</u>, April 10, Poster presentation

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Promising Signals of Clinical Activity in Adjuvant Setting



TG4050 | Additional Data Expected in 2024

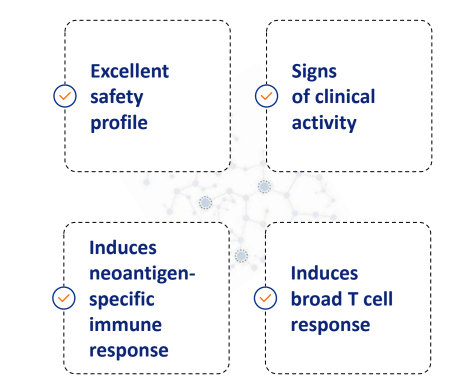
one patient • one genome • one vaccine

- → 24-month median follow up expected in H2 2024
- → Phase II part of the trial to start in coming weeks in head and neck cancer (adjuvant setting)

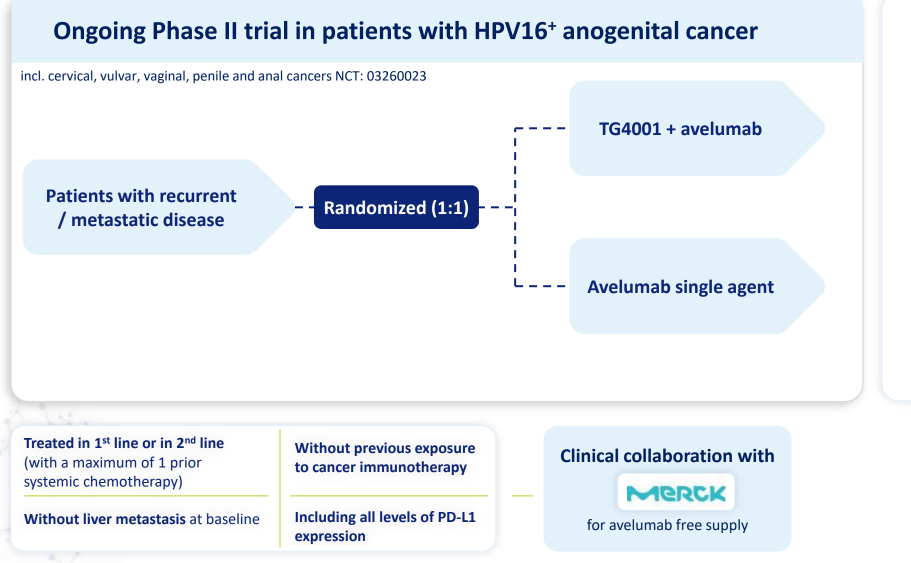
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Potential to extend remission period and address a significant market (head and neck cancer – adjuvant)

Could address other solid tumors in perioperative settings w or w/o ICIs – Additional trial to start in 2025



TG4001 | Ongoing Proof of Concept Trial to Read out in H2 2024



Topline data expected in H2 2024

- To deliver PoC data in significant patient population
- To further validate MVA platform

Clear path to approval in recently changed landscape

- Phase III in 1st line HPV+ cervix cancer in combin. with SoC (CT + ICI)
- Pivotal 2nd line after ICI

Objective

Sign partnership or licensing agreement based on Ph. II data

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Oncolytic Viruses

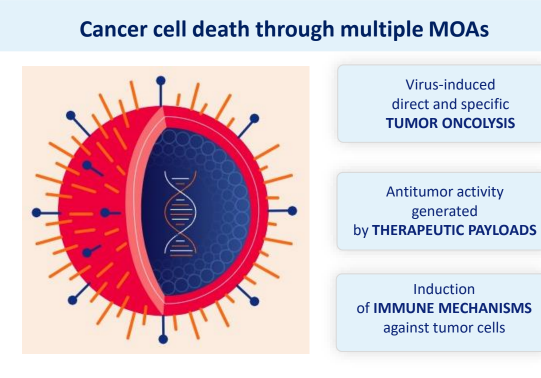
Rapidly Generating Multiple Virus-PoweredOff-the-Shelf Drug Candidates Targeting Solid Tumors





Our Oncolytic Viruses (OV) – Combined Effects of Vector, Payload and Immune Stimulation Compelling Clinical Data Support Intravenous (IV) Route of Administration

invir



Patented Backbone VV_{cop}TK⁻RR⁻ vector with multiple competitive advantages:

Encode numerous and various payloads

Multiple routes of administration (IV, IT, locoregional) and extend OV market beyond IT administration

- Potential to target multiorgan lesions and warm up TME
- O Address broad range of solid tumors

Proof of principle obtained

- Good safety profile
- Able to reach tumors, selectively replicate and express payload, incl. via intravenous administration

Goal: to target multiorgan lesions and reverse tumor resistance

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TG6050 Administered IV | IL-12 and anti-CTLA4 Produced Directly in the Tumor

Ongoing Phase I Trial to Assess Systemic Route of Administration



Initial goal

demonstrate potential of IV administration in "cold", non-resectable metastatic tumors

Oncolytic armed with IL-12 and anti-CTLA4 Ab

- P Triggers a powerful antitumor immune response
- Restores the immune defenses within the tumor
- O Outstanding preclinical data (strong antitumor activity) presented at AACR 2023

Phase I trial - Indication: metastatic and PD1 failed tumors

- Advanced or metastatic NSCLC after failure with available treatment options, including anti-PD1/PD-L1 Intravenous (IV) administration Inclusions ongoing (NCT: 05788926)
- Initial data (single agent) in H2 2024 Could be combined with ICIs

Potential to address a major oncology market



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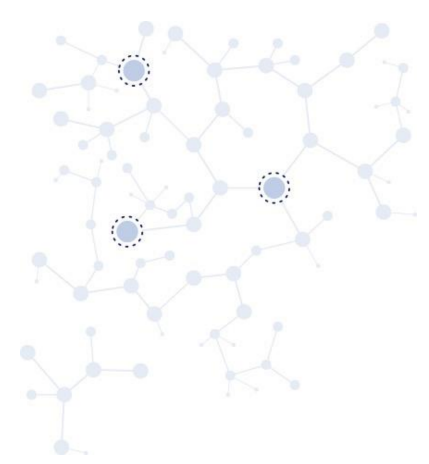
BT-001 OV Armed with Anti-CTLA4 Ab + GM-CSF 50/50 with **BioInvent Ongoing Phase I Trial Assessing IT* Route of Administration** The right virus + payload **Positive Phase I part A readout** VV_{cop}TK⁻RR⁻ oncolytic armed with → Single agent well tolerated BioInvent's potent anti-CTLA4 Ab + GM-CSF → Replicates and persists in tumor tissue Activates and increases T-effector cells → Anti-CTLA4 expressed in the tumor · Treg depleting activity with no detectable systemic exposure · Stimulates immune cells (incl. APC) → Stable injected lesion in 11/18 patients → Tumor shrinkage observed in two patients Ongoing Phase I (NCT04725331) monotherapy and combination w. anti-PD1

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(>)**Ph. I part B** (combination with pembrolizumab) – Enrolment ongoing - First data expected in H2 2024

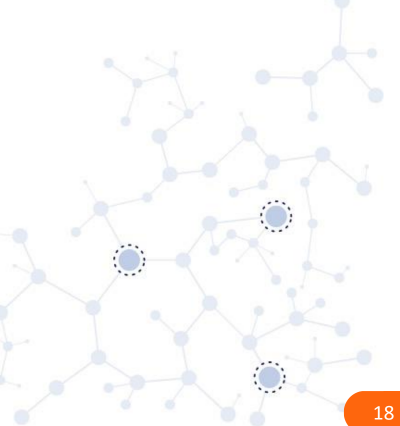
collaboration

Can be developed for multiple cancer indications lesions with high Treg infiltration



Outlook





Company Funded to Deliver Multiple Value Generating Milestones

FINANCIAL VISIBILITY secured until Q4 2025

Enables Transgene to deliver significant milestones including key PoC data for TG4050 and data on all assets





TG4050 | Neoantigen vaccine

myvac 🗸

Proof of principle

already obtained in Head and Neck cancer (adjuvant)

Ongoing randomized Phase I (head and neck cancer)

- 24-month median follow up of patients (H2 2024)

^o Randomized Phase II (head and neck cancer)

- Initiation of the trial in collaboration with NEC (Q2 2024)

Other indication

Prepare new Phase I

Shared antigens cancer vaccines

• **TG4001:** Results from ongoing randomized Phase II (H2 2024)

Oncolytic viruses

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TG6050: Initial Phase I data (**H2 2024**)

• **BT-001:** Initial data in combination with pembrolizumab (**H2 2024**)



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Investment Highlights



Unique and highly potent viral vector based immunotherapies Lead program TG4050 to deliver data in 2024 and create significant value by 2026

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> Additional programs and R&I activity to deliver news flow and fuel Transgene's portfolio in the mid term

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Appendices

New Leadership to Take Transgene to the Next Level



ALESSANDRO RIVA, MD Chairman & CEO

30+ years experience



GILEAD ...ichnos...



ÉRIC QUÉMÉNEUR, PharmD, PhD - Executive VP - Chief Scientific Officer



LUCIE LARGUIER VP, Chief Financial Officer



MAUD BRANDELY, MD, PhD - VP, Medical Affairs - Chief Medical Officer



JAMES WENTWORTH VP, Chief Business Officer



JOHN FELITTI VP, Legal Counsel



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CHRISTELLE SCHWOERER VP, Human Resources



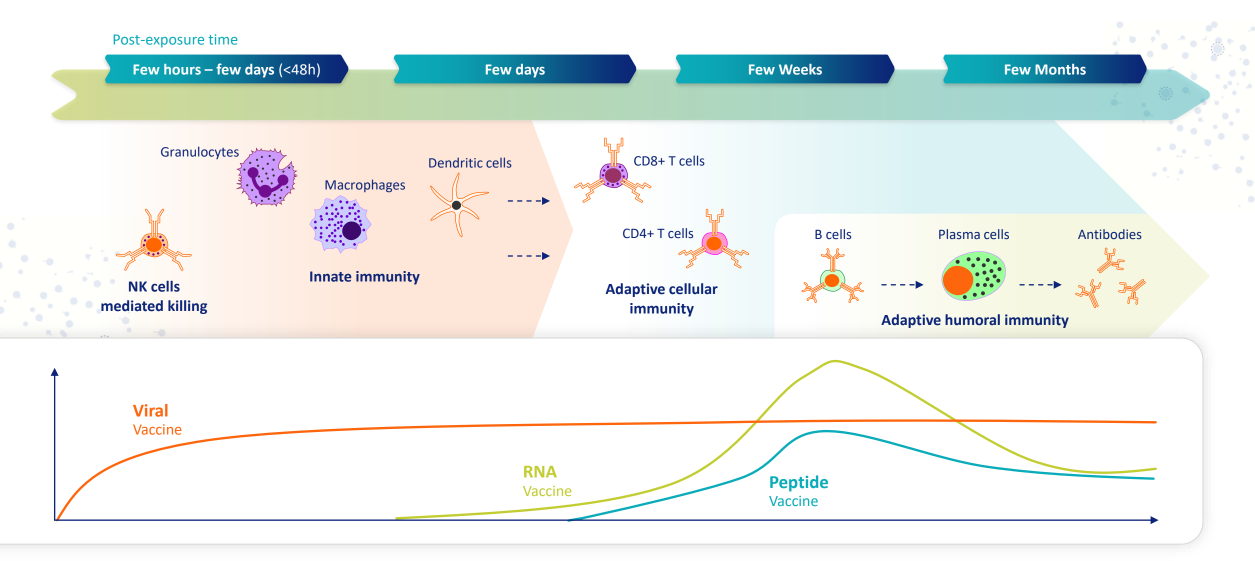
JOHN C. BELL Member of the Scientific Advisory Board



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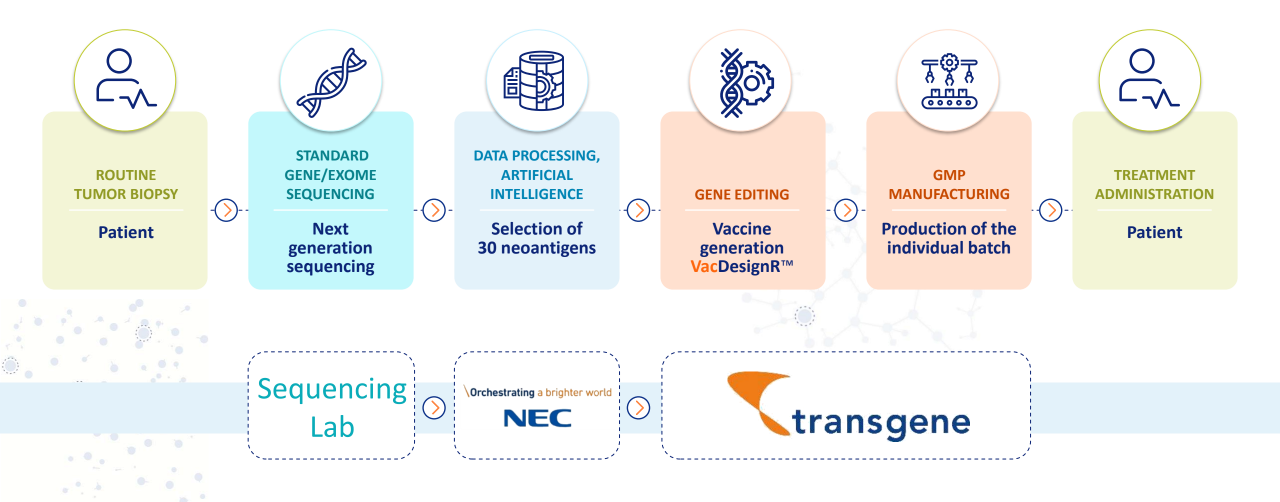


Viral Vaccines Prime Both Long Lasting Innate and Adaptive Immunity



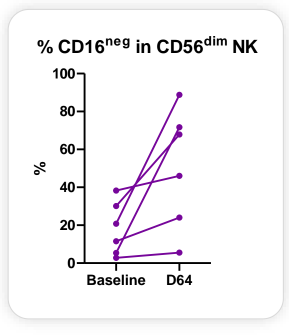


TG4050, an Individualized Neoantigen Vaccine Combining Unique Capabilities Combines Bioengineering and Digital Transformation

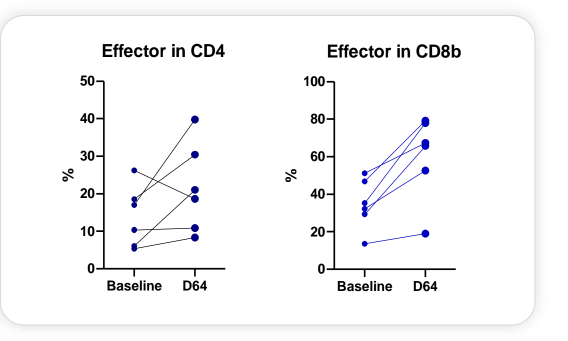


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Profound Remodelling of Immune Cells Consistent with Anti Tumor Response Suggests that the Vaccine Effectively Primes the Immune System



Priming of innate immunity: Loss of CD16 on CD56^{dim} NK cells suggests ongoing antitumor activity



- Maturation and differentiation of CD4 and CD8 into effector cells
 Consistent with the development of an active adaptive response
- Seffector subgroups of CD4 and CD8 T-cells are increased

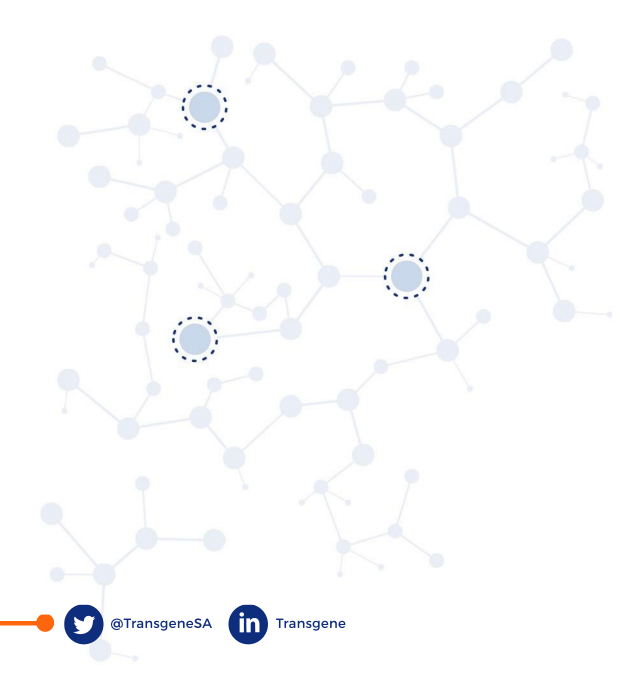


Source: Block et al, "Phase I trials of personalized cancer vaccine TG4050 in surgically treated high-risk head and neck squamous cell carcinoma (HNSCC) and relapsing ovarian cancer (OvC) patients" <u>AACR 2022</u>, April 12, 2022, Poster presentation











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