

Vaccibody's approach to cost-effective personalized cancer neoantigen vaccines inducing a unique CD8-dominated T cell response

August 30, 2018

Agnete Fredriksen, PhD
President & CSO
Vaccibody AS

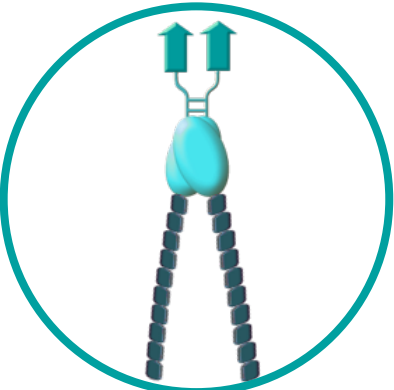
abfredriksen@vaccibody.com

Agenda

1. Background Cancer Neoantigens



2. Vaccibody's Cancer Vaccine Strategy



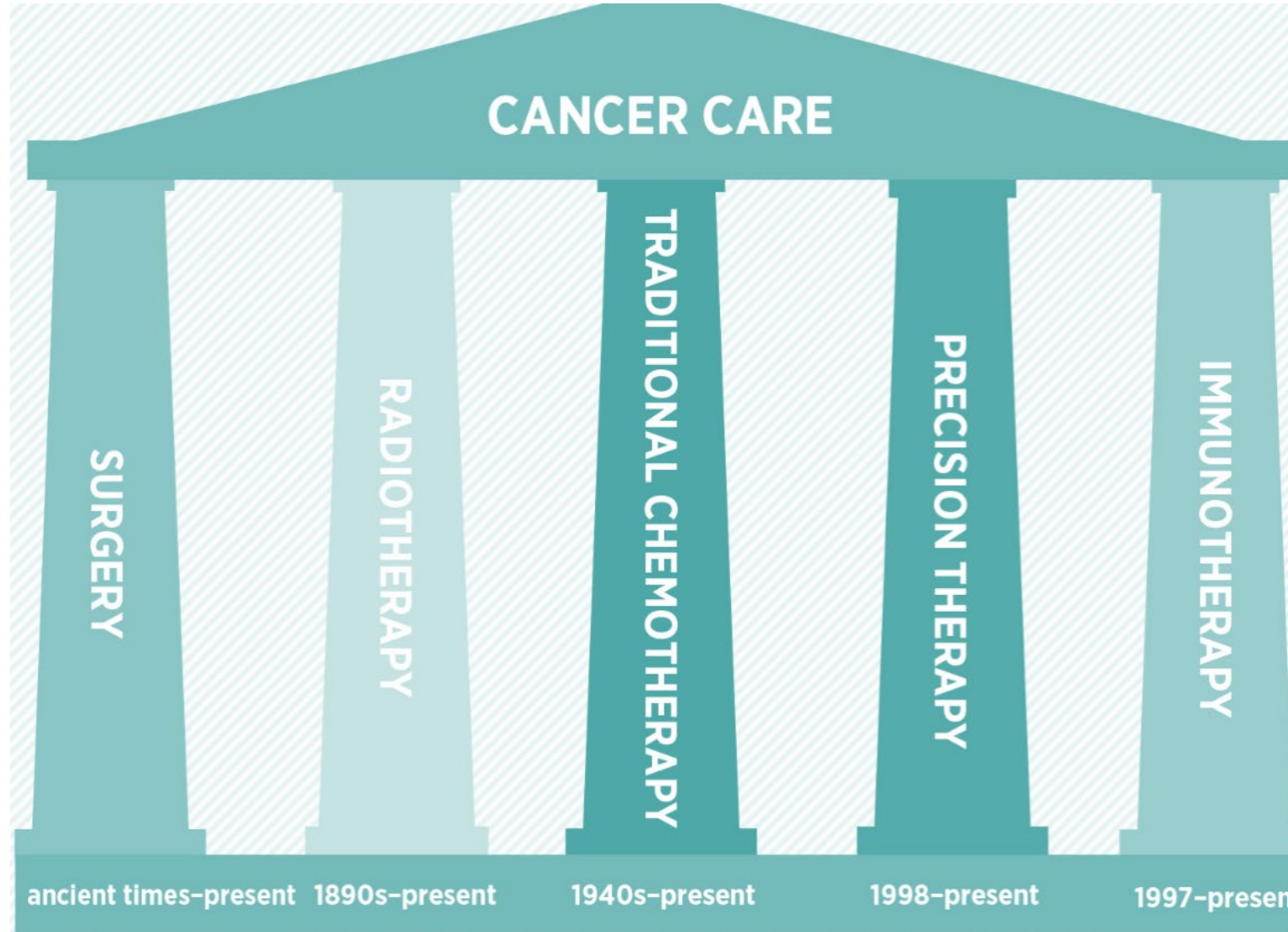
3. Neoantigen Prediction Tools
NeoSELECT™



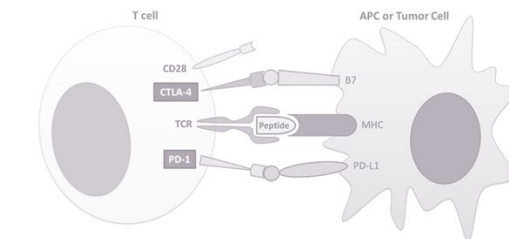
4. Vaccibody's Clinical Trial Experience
and Future Plans



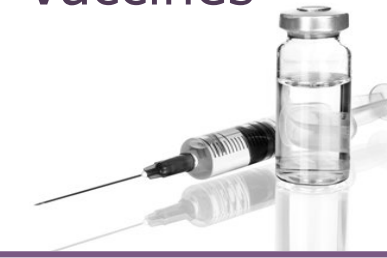
Immunotherapy: The next Wave of Cancer Therapy



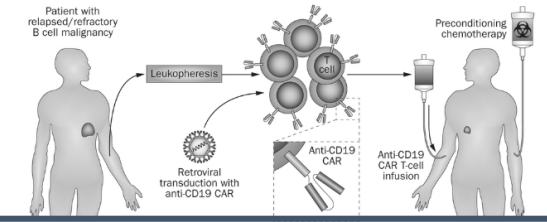
Checkpoint Inhibitors



Vaccines



Cell Therapies

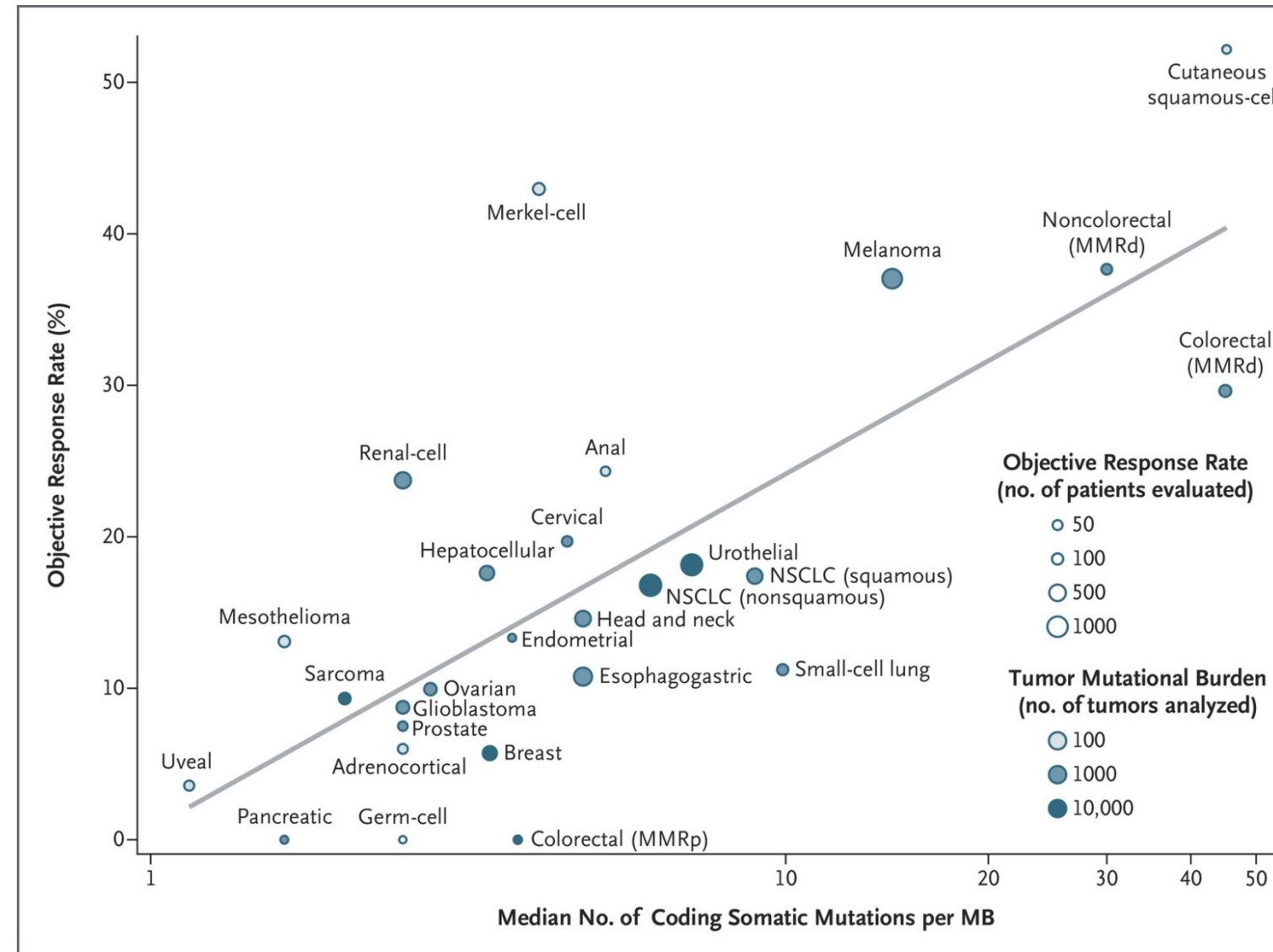


Others, e.g.

- Oncolytic viruses
- Cytokines
- Bi-specific antibodies
- Small molecules
- Adjuvants

Various Immuno-Therapy Modalities

CheckPoint Inhibitors – relationship to neoantigens



Strong relationship between mutational burden and response to CPI

Limits response to already existing neoantigen-specific T cell repertoire

Reveals an important role of immune responses to neoantigens in cancer immunotherapy

Cancer neoantigen vaccines are the **optimal tool** to activate a truly specific, strong and broad neoantigen specific T cell responses

Proof of Concept published in Nature Letters July 2017

LETTER

doi:10.1038/nature22991

An immunogenic personal neoantigen vaccine for patients with melanoma

Patrick A. Ott^{1,2,3*}, Zhuting Hu^{1*}, Derin B. Keskin^{1,3,4}, Sachet A. Shukla^{1,4}, Jing Sun¹, David J. Bozym¹, Wandi Zhang¹, Adrienne Luoma⁵, Anita Giobbie-Hurder⁶, Lauren Peter^{7,8}, Christina Chen¹, Oriol Olive¹, Todd A. Carter⁴, Shuqiang Li⁴, David J. Lieb⁴, Thomas Eisenhaure⁴, Evisa Gjini⁹, Jonathan Stevens¹⁰, William J. Lane¹⁰, Indu Javeri¹¹, Kaliappanadar Nellaippan¹¹, Andres M. Salazar¹², Heather Daley¹, Michael Seaman⁷, Elizabeth I. Buchbinder^{1,2,3}, Charles H. Yoon^{3,13}, Maegan Harden⁴, Niall Lennon⁴, Stacey Gabriel⁴, Scott J. Rodig^{9,10}, Dan H. Barouch^{3,7,8}, Jon C. Aster^{3,10}, Gad Getz^{3,4,14}, Kai Wucherpfennig^{3,5}, Donna Neuberg⁶, Jerome Ritz^{1,2,3}, Eric S. Lander^{3,4}, Edward F. Fritsch^{1,4†}, Nir Hacohen^{3,4,15} & Catherine J. Wu^{1,2,3,4}

- 6 patients with melanoma (stage III/IV)
- 97 neoepitopes delivered as long-peptides with polyICLC (SC)
- CD4 dominated responses

LETTER

doi:10.1038/nature23003

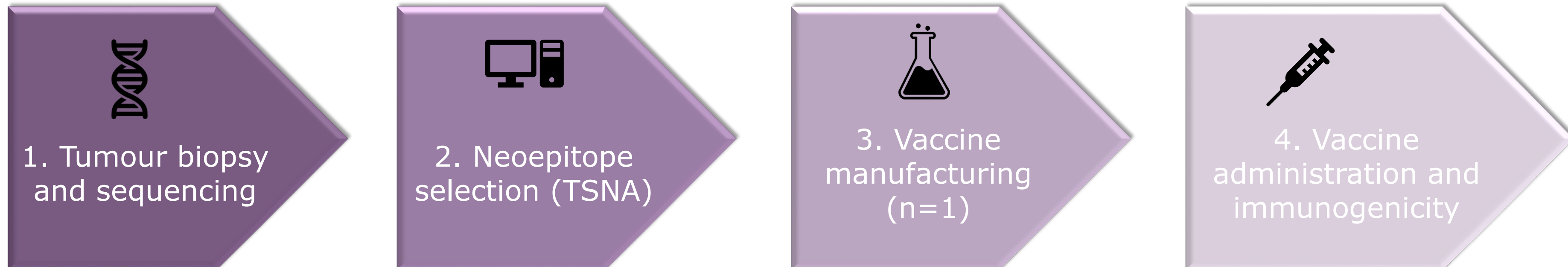
Personalized RNA mutanome vaccines mobilize poly-specific therapeutic immunity against cancer

Ugur Sahin^{1,2,3}, Evelyn Derhovanessian¹, Matthias Miller¹, Björn-Philipp Kloke¹, Petra Simon¹, Martin Löwer², Valesca Bukur^{1,2}, Arbel D. Tadmor², Ulrich Luxemburger¹, Barbara Schrörs², Tana Omokoko¹, Mathias Vormehr^{1,3}, Christian Albrecht², Anna Paruzynski¹, Andreas N. Kuhn¹, Janina Buck¹, Sandra Heesch¹, Katharina H. Schreeb¹, Felicitas Müller¹, Inga Ortseifer¹, Isabel Vogler¹, Eva Godehardt¹, Sebastian Attig^{2,3}, Richard Rae², Andrea Breitkreuz¹, Claudia Tolliver¹, Martin Suchan², Goran Martic², Alexander Hohberger³, Patrick Sorn², Jan Diekmann¹, Janko Ciesla⁴, Olga Waksman⁴, Alexandra-Kemmer Brück¹, Meike Witt¹, Martina Zillgen¹, Andree Rothermel², Barbara Kasemann², David Langer¹, Stefanie Bolte¹, Mustafa Diken^{1,2}, Sebastian Kreiter^{1,2}, Romina Nemecek⁵, Christoffer Gebhardt^{6,7}, Stephan Grabbe³, Christoph Höller⁵, Jochen Utikal^{6,7}, Christoph Huber^{1,2,3}, Carmen Loquai^{3,*} & Özlem Türeci^{8*}

- 13 patients with melanoma (stage III/IV)
- 125 neoepitopes delivered as ivt-RNA (intranodal)
- CD4 dominated responses

- Vaccinating with neoepitopes elicits a broad and strong tumour-specific immune response
- Both peptide and RNA neoantigen based vaccines elicits predominantly CD4 T cell responses

The Workflow of Personalised Cancer Treatment

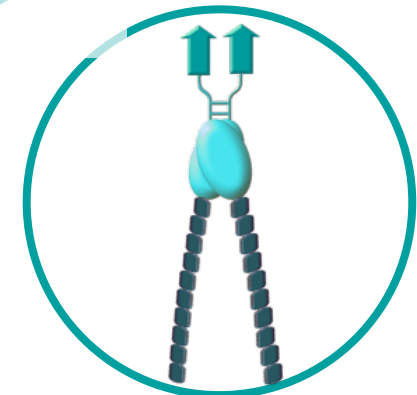


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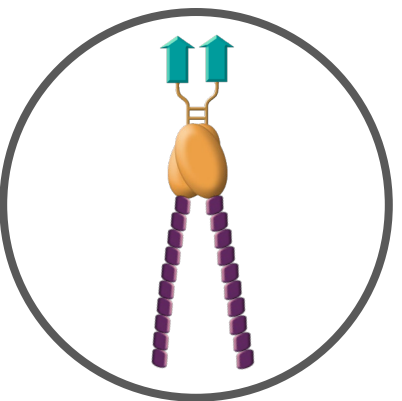
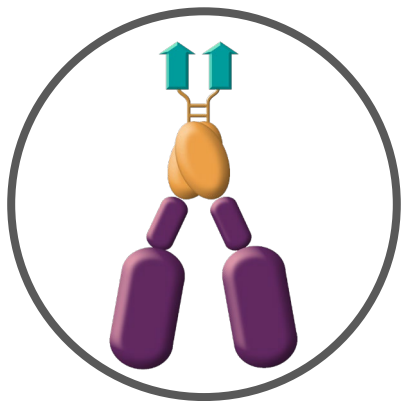
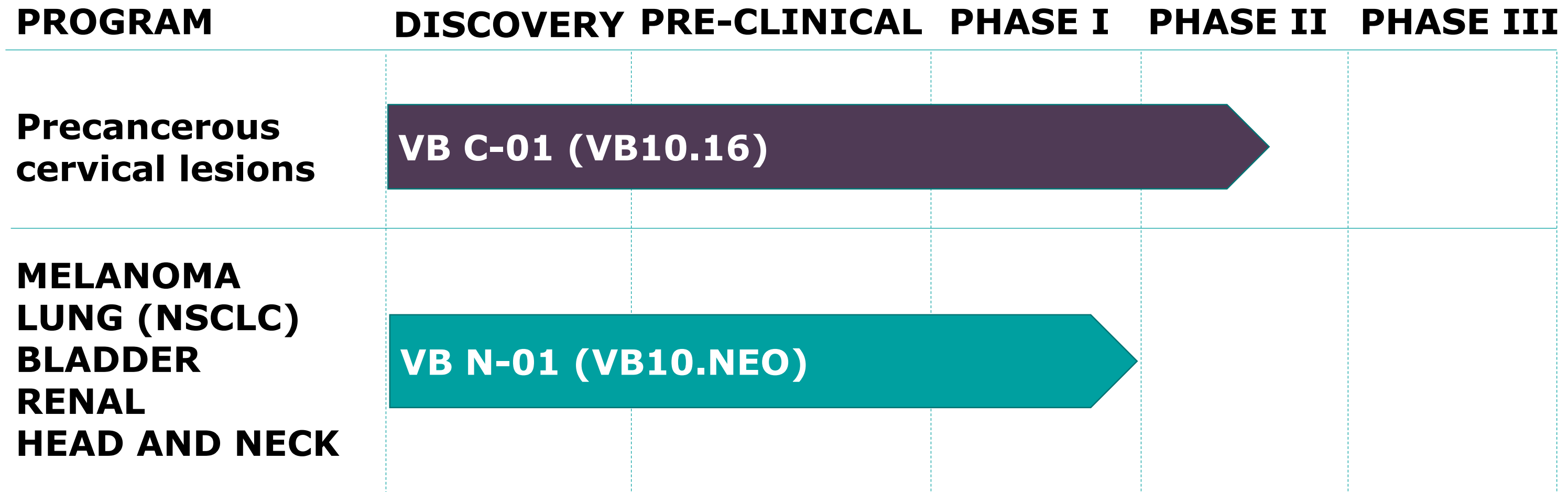
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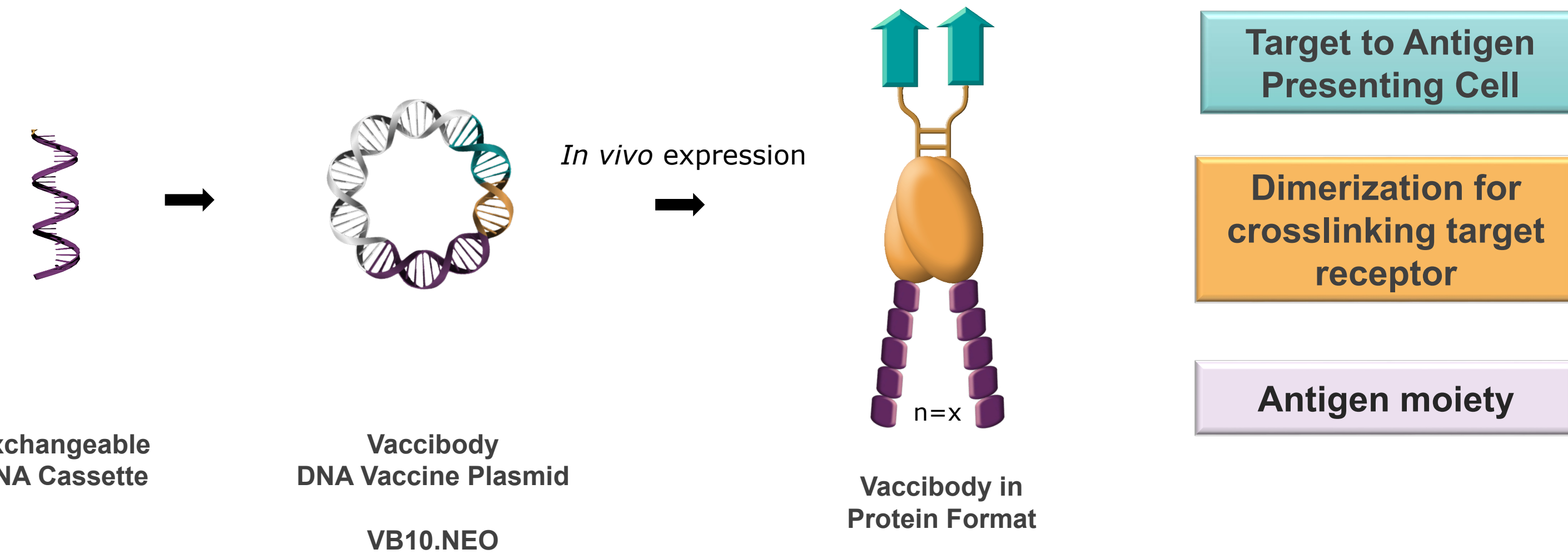


Vaccibody Product Pipeline

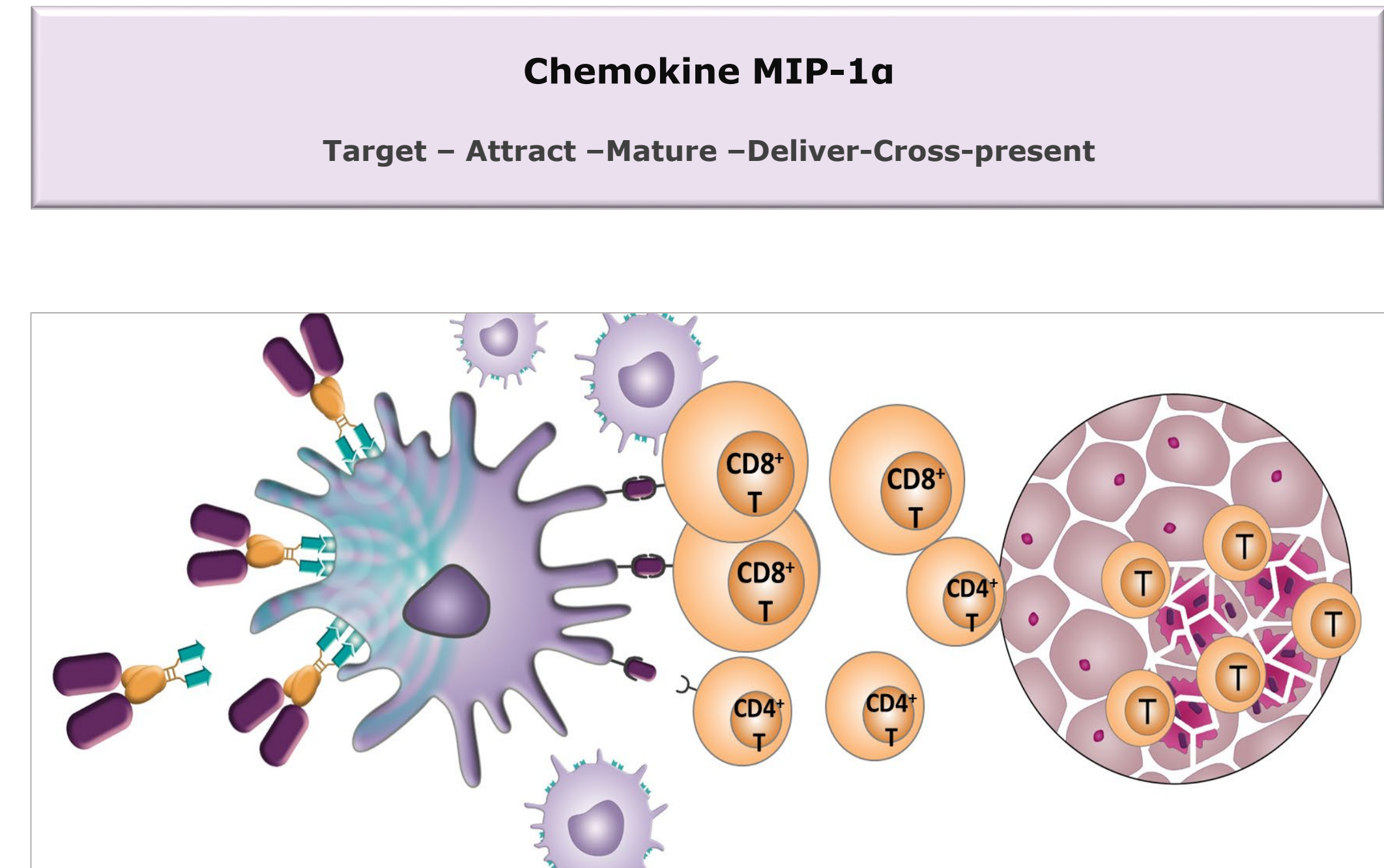
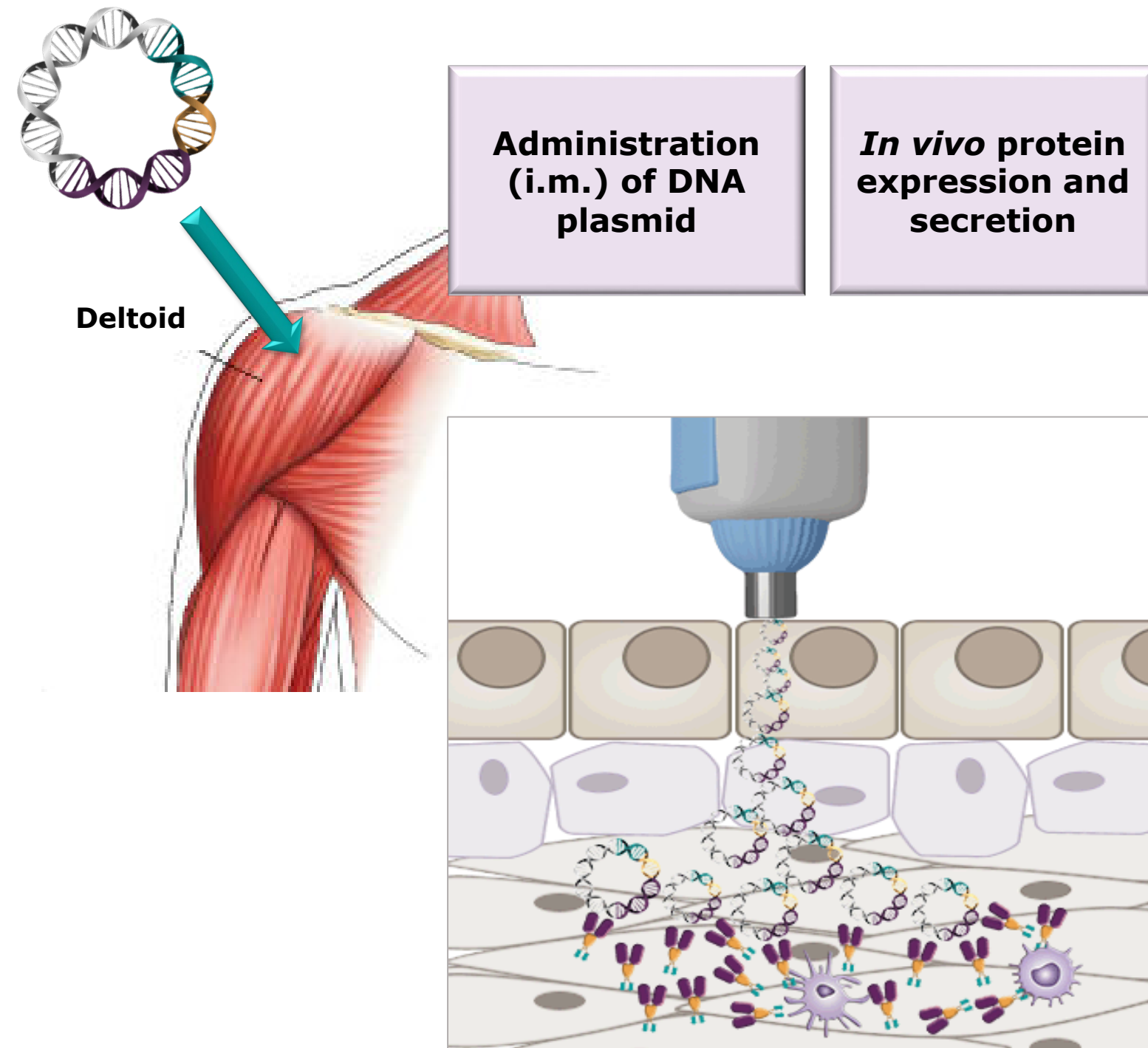


Vaccibody – Proprietary Vaccine Technology Platform

The Vaccibody Technology Platform was developed based on the concept of **targeting antigen to APC** in order to create more efficacious vaccines.



Mechanism of Action – Intrinsic Adjuvant



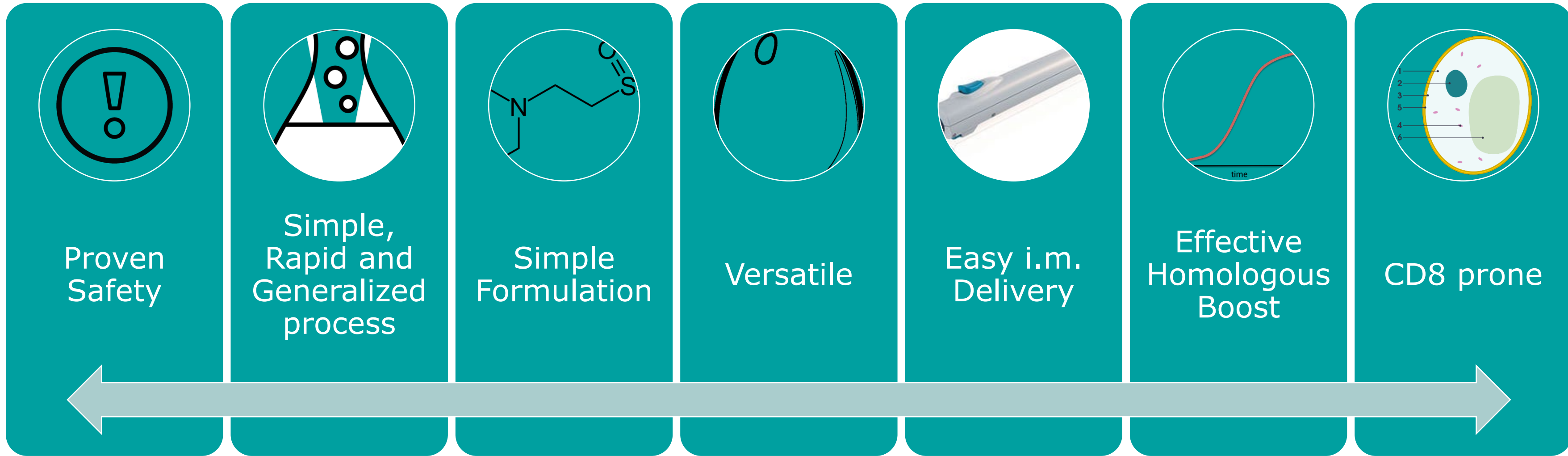
Patient Friendly, simple Vaccine Delivery

PharmaJet®



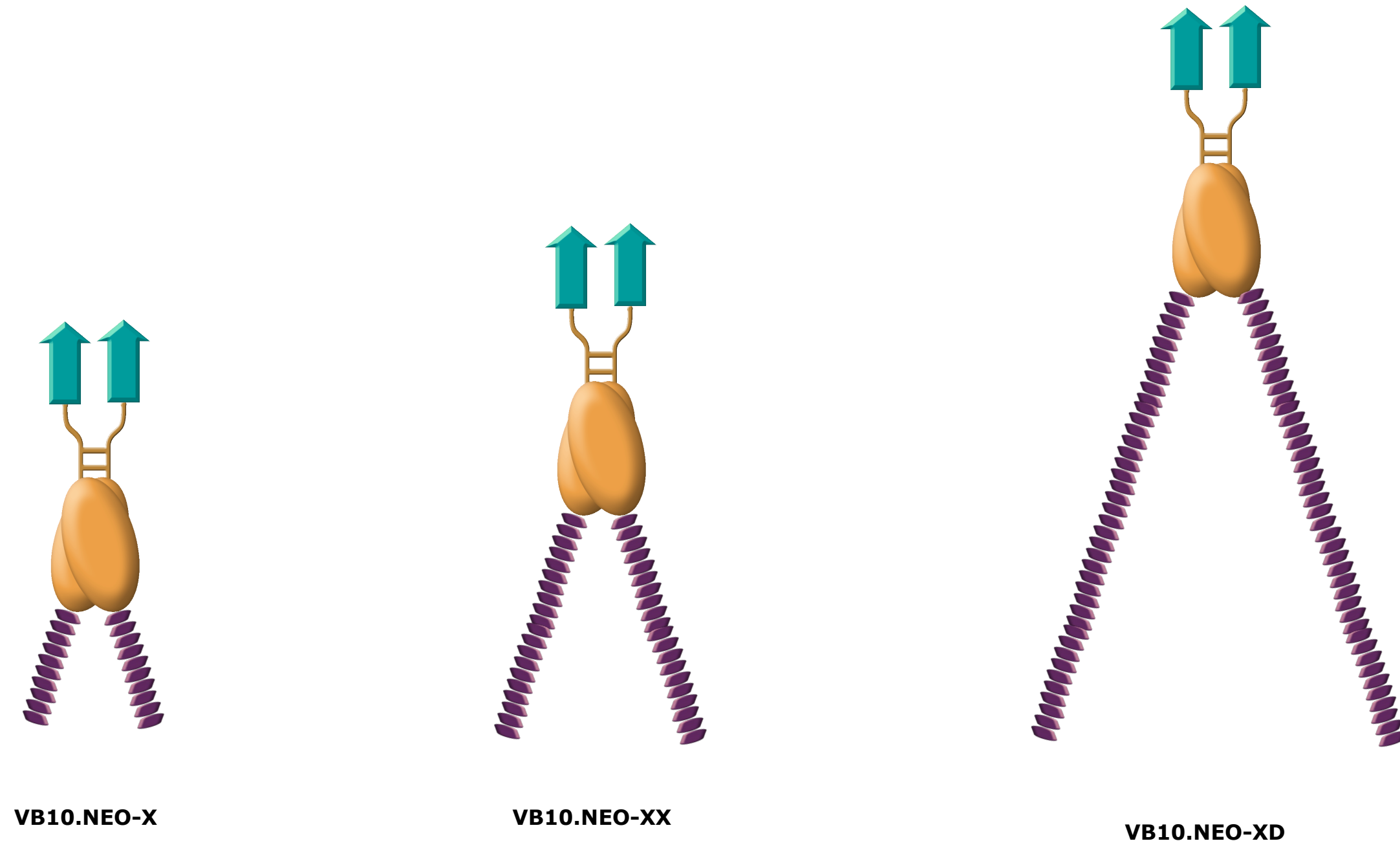
- Needle free injection**
- Small, handy, easy to use**
- Minimal pain compared to electroporation**
- Cost effective**
- Applicable for multiple immunizations**
- High patient compliance**

Naked DNA plasmid as IMP



DNA plasmid is an ideal platform for bringing individualized neoantigen vaccines to the market as a viable product

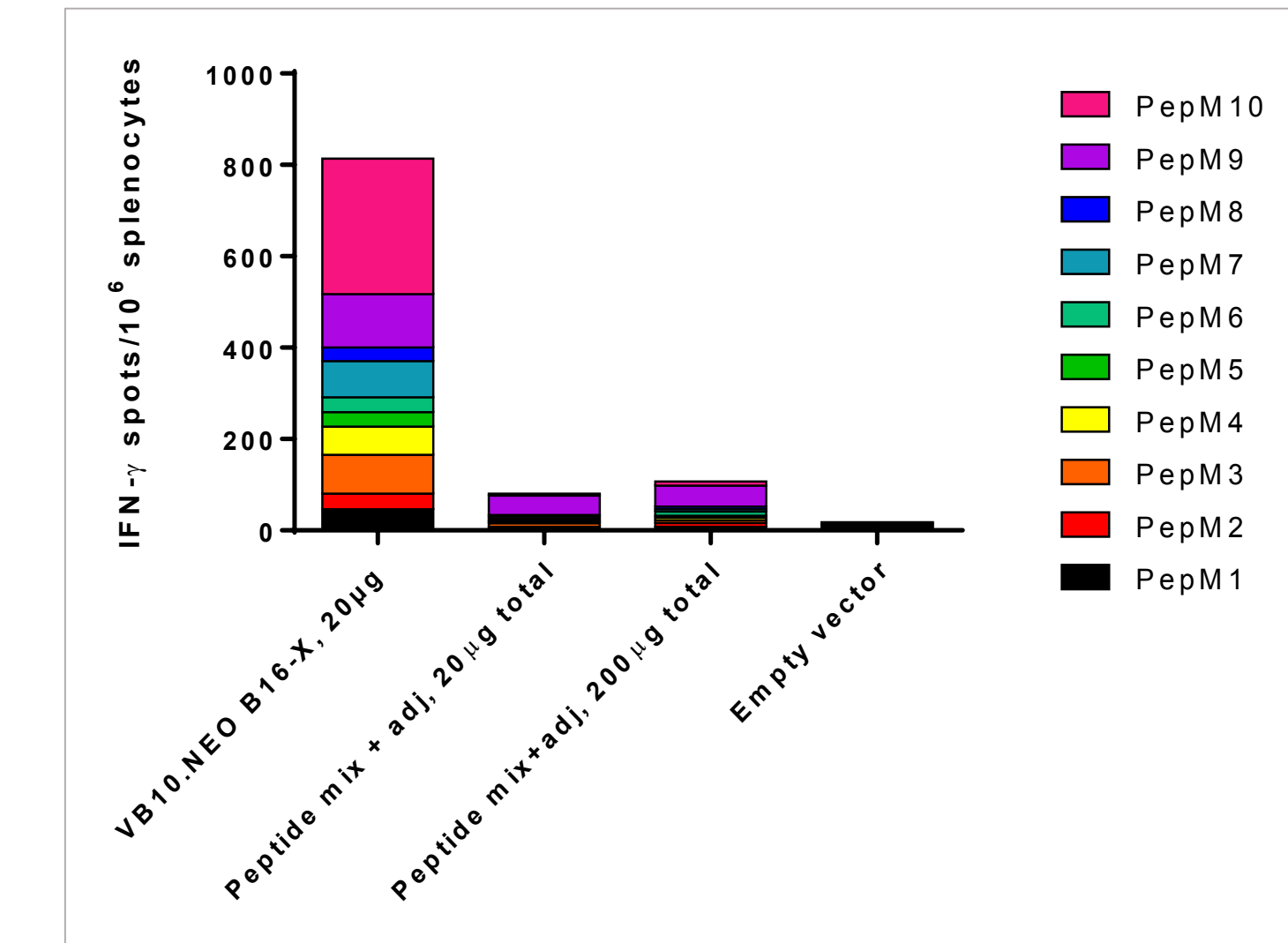
VB10.NEO – A Robust Vaccine Format



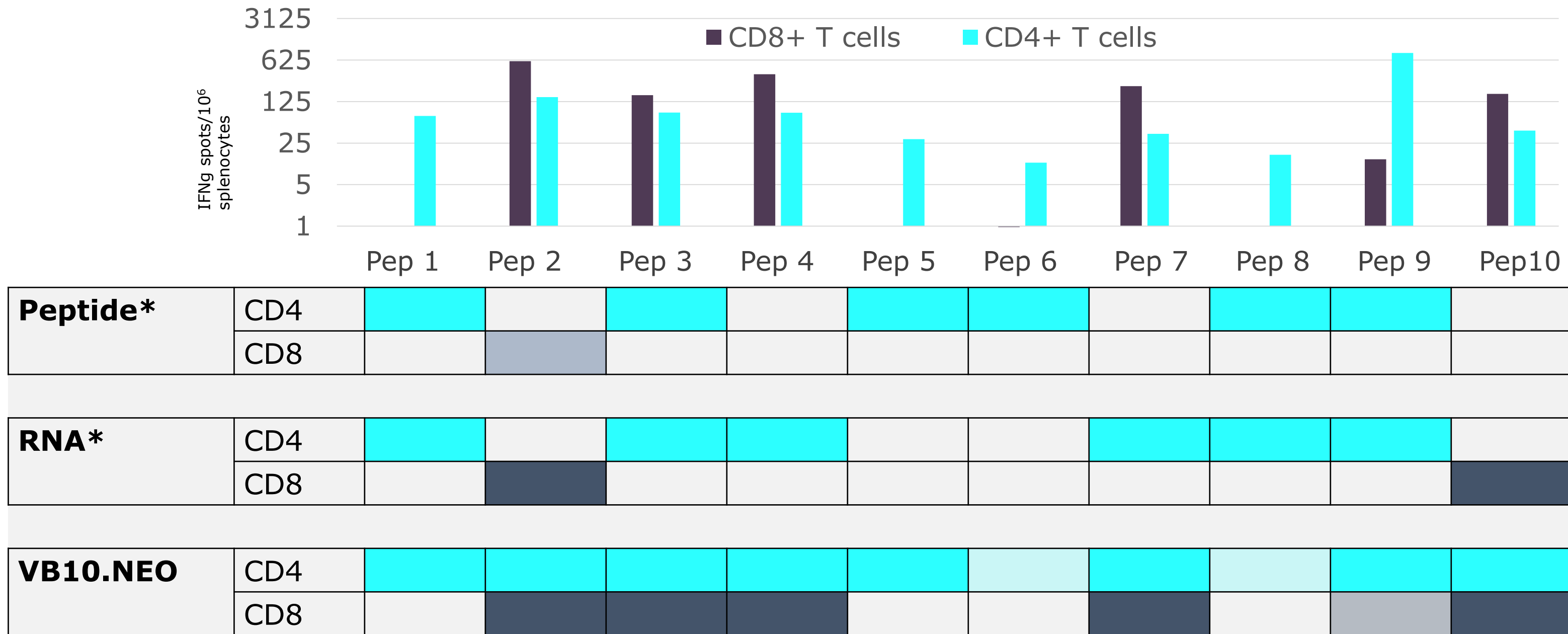
>80 different VB10.NEO constructs with >250 neoepitopes constructed to date with up to 40 neoepitopes

VB10.NEO induces Rapid, Broad and Strong responses to multiple Neoepitopes by single Vaccination

- VB10.NEO induces a broader and stronger response than Peptide + Poly (I:C) Adjuvant vaccines after a single immunization.
 - VB10.NEO vaccinated animals respond to all 10 neoepitopes after a single immunization.
 - Immunodominant neoepitopes differ between delivery vehicles



VB10.NEO generates a broader immune response profile dominated by CD8⁺ T cells than competing technologies



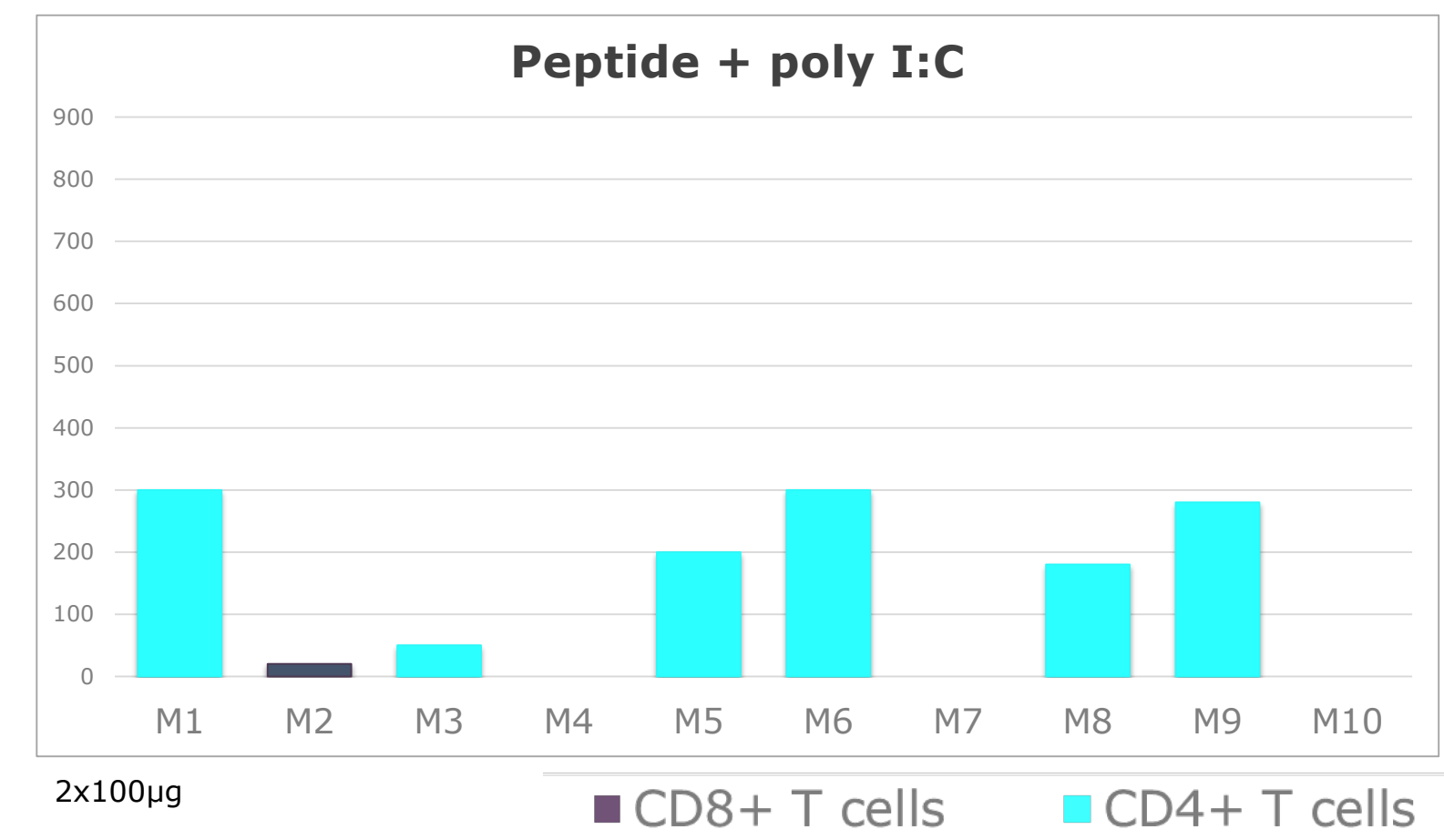
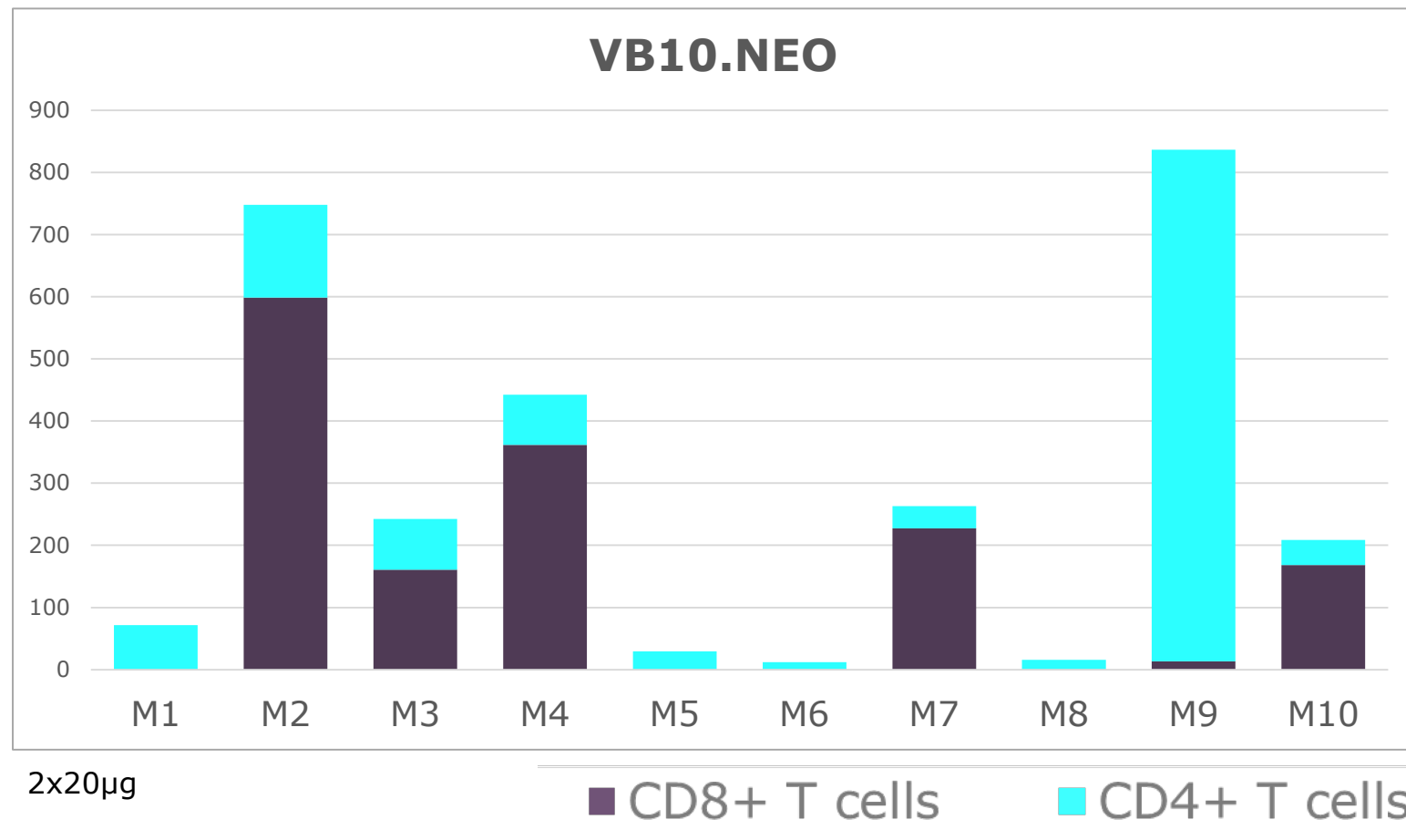
* Tested IFN- γ CD4 and CD8 T cell response against 10 identical neoepitopes from B16 melanoma

Peptide and RNA vaccines induces primarily CD4 T cell responses, while VB10.NEO induces strong, dominating CD8 responses to the identical neoepitope sequences

VB10.NEO leads to a unique CD8 dominated neoepitope response

VB10.NEO induces a **strong, broad** immune response
dominated by CD8+ T cells

Peptide + poly I:C vaccination has been reported to
 induce **dominantly CD4 T cell responses**

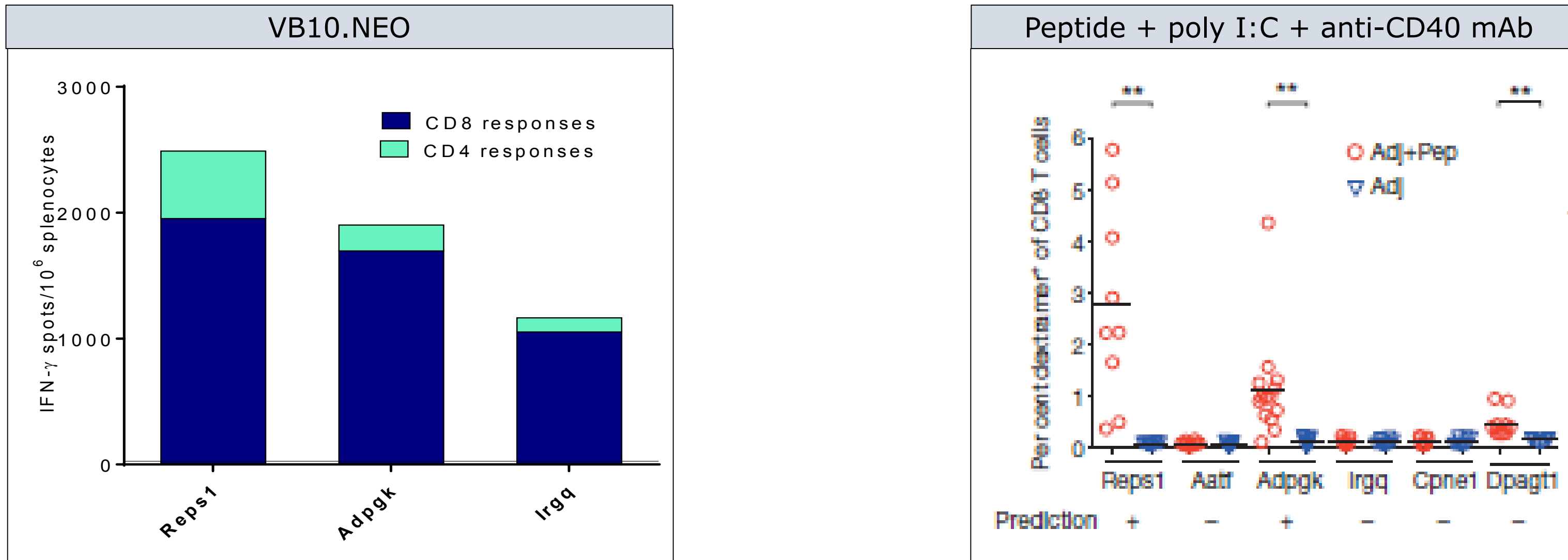


VB10.NEO induces strong, dominantly CD8+ T cell response to identical neoepitopes
 that induces **no or weak** immune response if delivered as peptide vaccine

Confirmation of VB10.NEO's unique ability to induce strong neoepitope-specific CD8 responses

Yadav et al., 2014

MC38 colon carcinoma



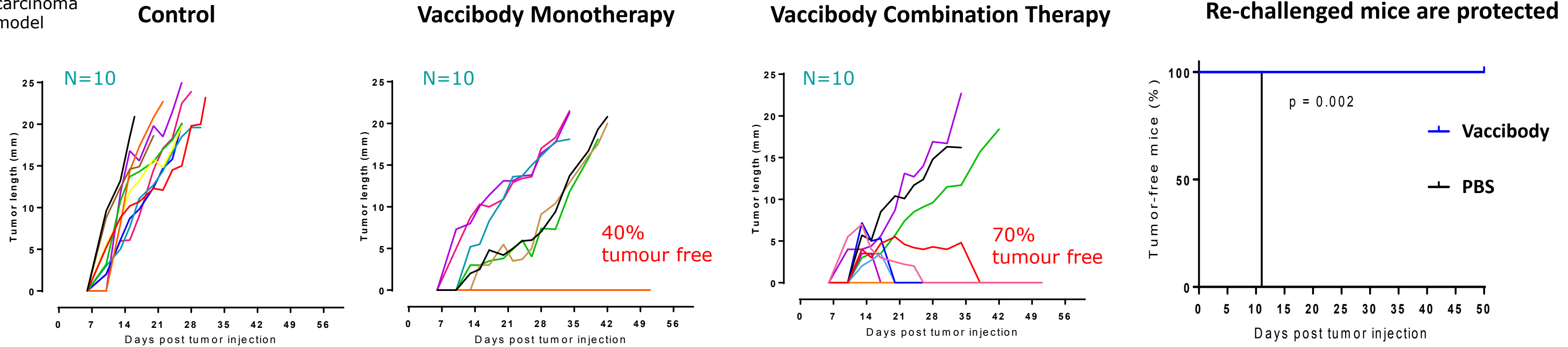
-**VB10.NEO induces a strong CD8 T cell response**, combined with a CD4 T cell response **to all** peptides tested for MC38 colon carcinoma.

-1/3 of these neoepitopes have been shown to be **non-immunogenic delivered as peptide + adjuvant**

-**Confirmation of VB10.NEO's ability to induce stronger CD8 responses to neoantigens**

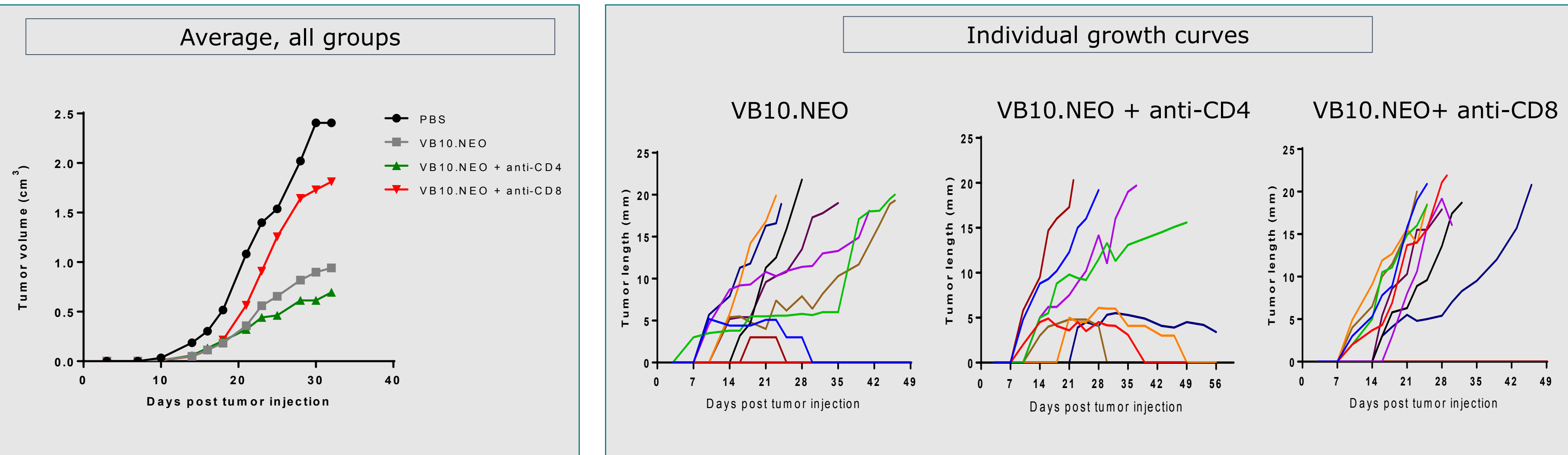
Vaccibody Induces Tumor Protection as Monotherapy

CT26 colon carcinoma model



- Vaccibody vaccination induces strong CD8+ T cell responses and **tumor protection as Monotherapy**
- Combination with anti-PD-1 immunotherapy induced enhanced anti-tumour responses in mice involving **complete tumour regression** of large, established tumours
- **Long-term memory responses** ensure effective anti-tumour responses after a 2nd tumour challenge in surviving mice with no sign of tumour growth

Neoepitope-specific CD8 T cells are crucial for tumour protection



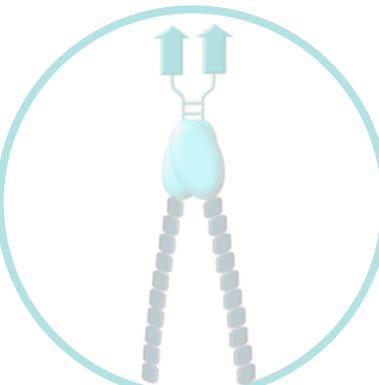
Depletion of CD8 T cells prohibit tumour protection in VB10.NEO vaccinated mice, indicating a crucial role of neoepitope-specific CD8 T cells for anti-tumour efficacy

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Why the perfect fit for individualised Vaccines?



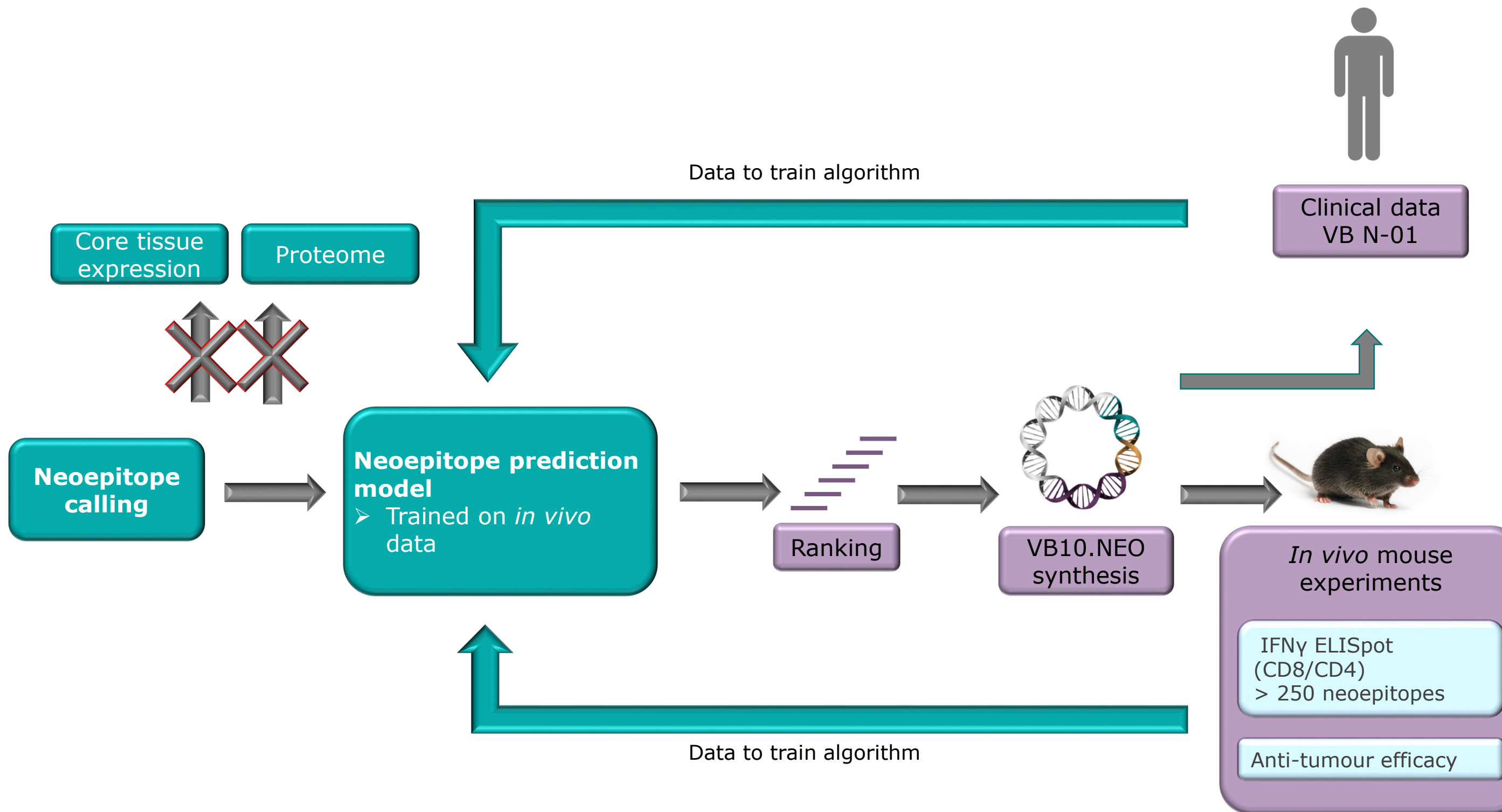
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Developing VB10.NEO specific Neoepitope Selection

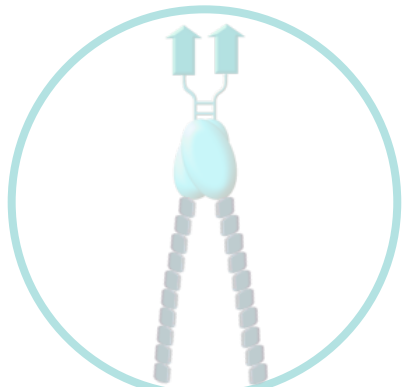


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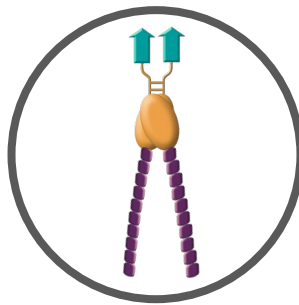
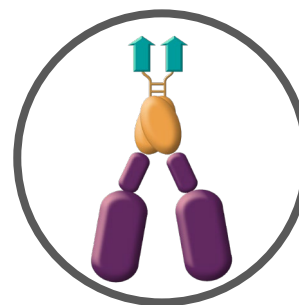
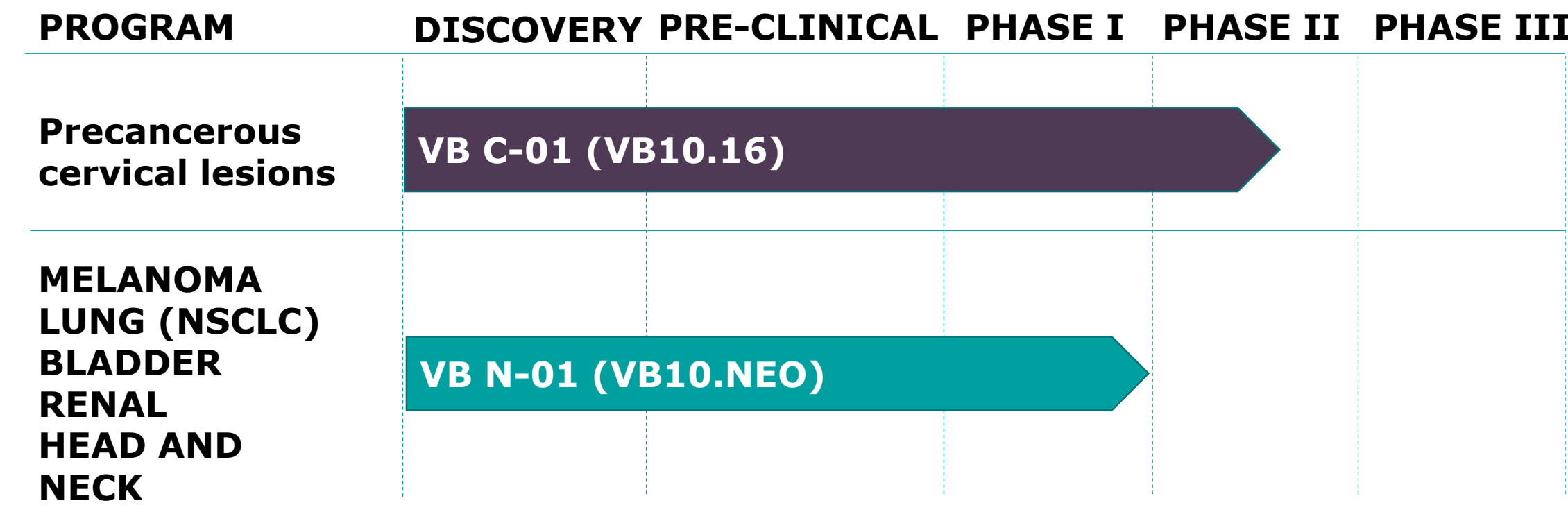
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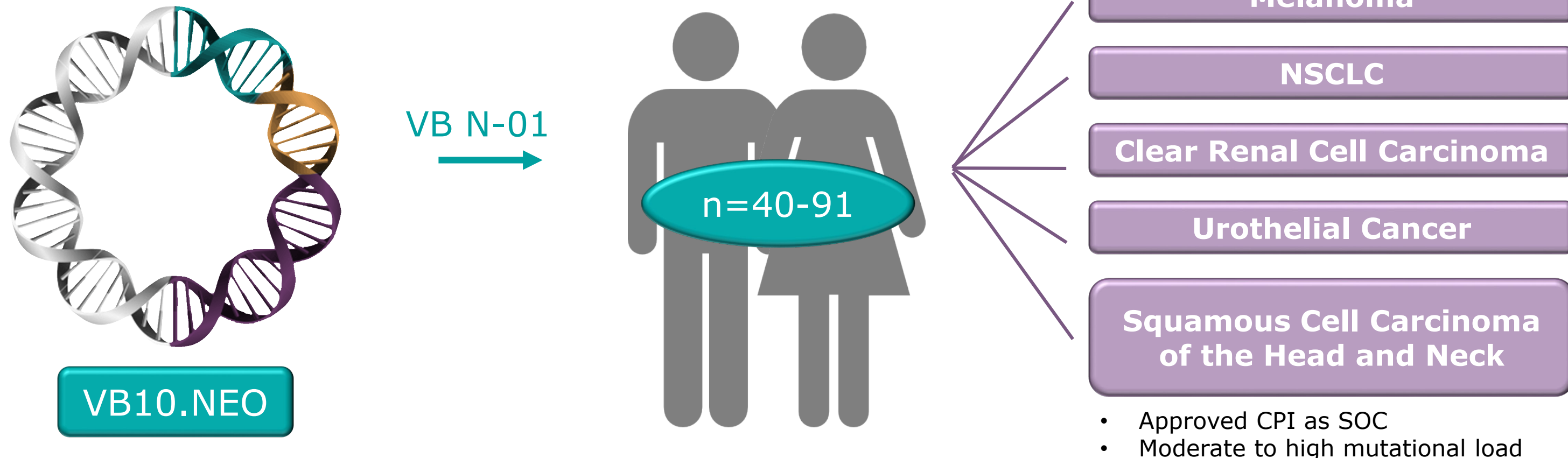
Vaccibody Vaccine Product Pipeline



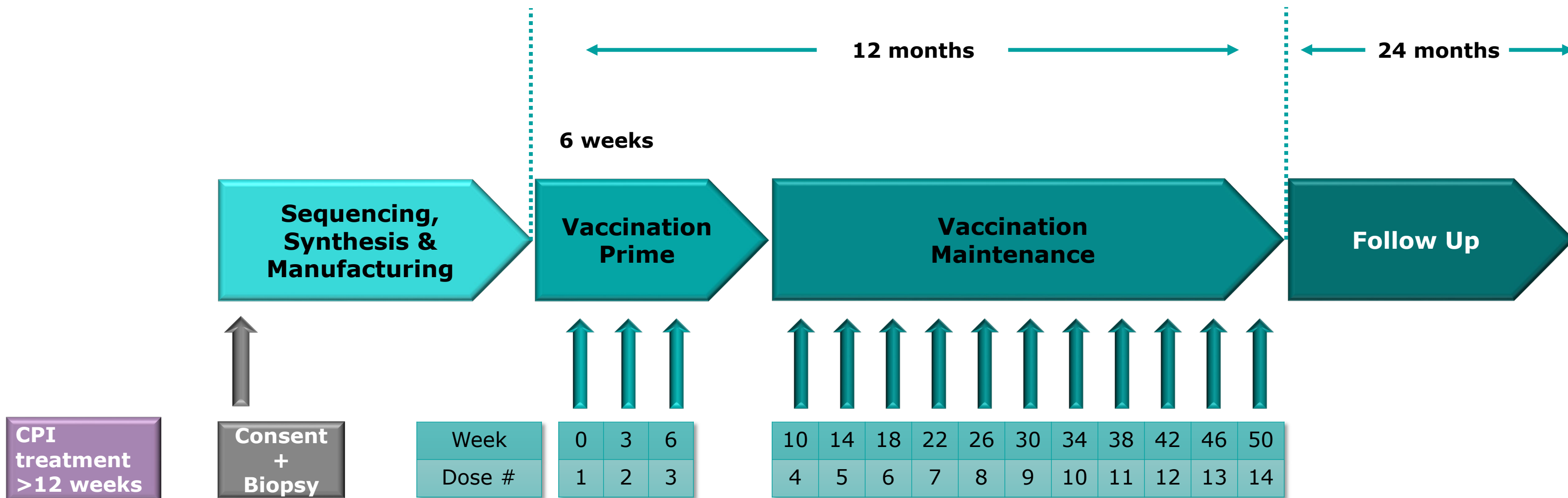
Clinical Trial VB N-01

VB N-01: An open labelled first human dose phase 1/2a study to evaluate safety, feasibility and efficacy of multiple dosing with individualised VB10.NEO immunotherapy in patients with locally advanced or metastatic melanoma, NSCLC, clear renal cell carcinoma, urothelial cancer or squamous cell carcinoma of head and neck, who did not reach complete responses with current standard of care immune checkpoint blockade

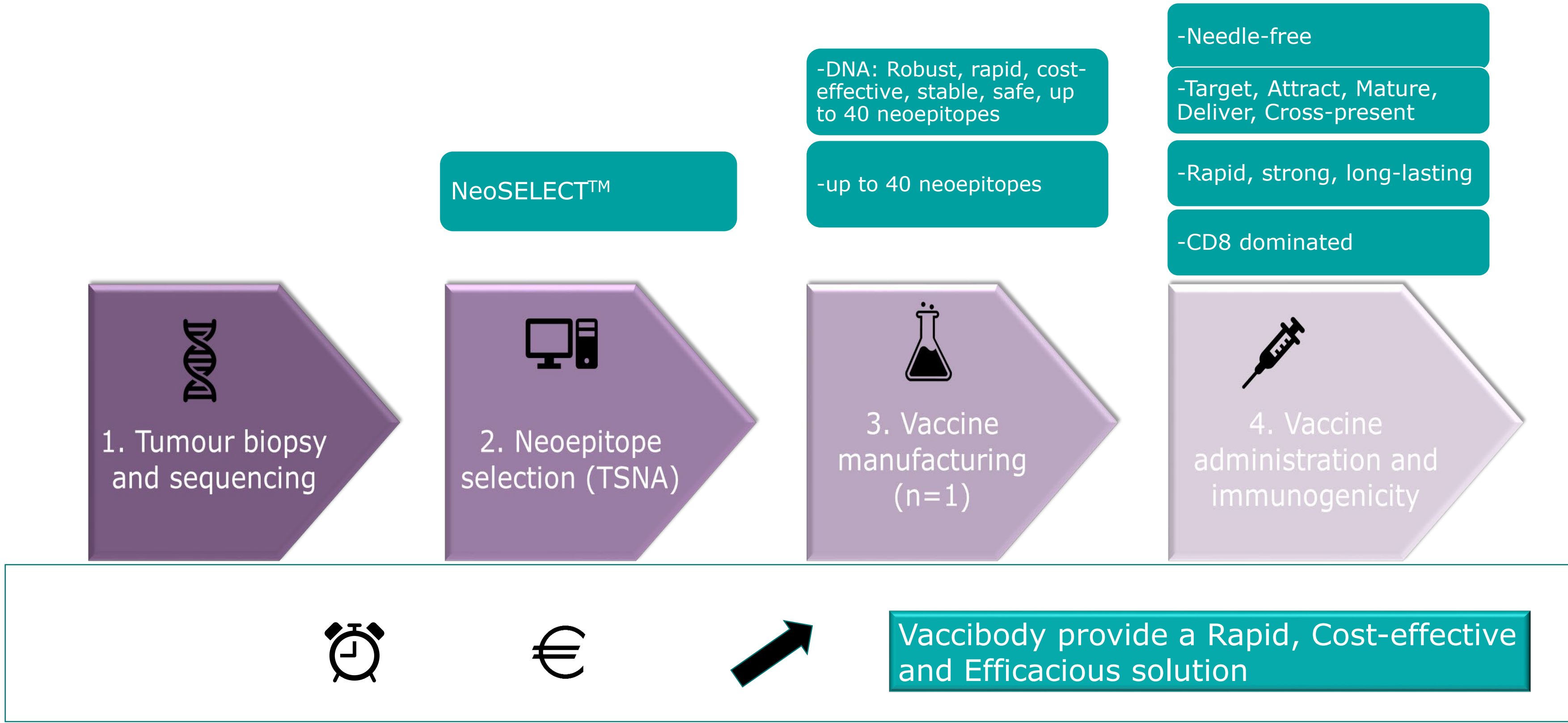
FPI April 2018



Study Design and Treatment Schedule VB N-01



Vaccibody's Solution to Personalised Cancer Treatment



vaccibody

www.vaccibody.com