

Forward-looking statement

This presentation contains forward-looking statements within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. The words "target," "believe," "expect," "hope," "aim," "intend," "may," "might," "anticipate," "contemplate," "continue," "estimate," "plan," "potential," "predict," "project," "will," "can have," "likely," "should," "would," "could," and other words and terms of similar meaning identify forward-looking statements. Actual results may differ materially from those indicated by such forward-looking statements as a result of various factors, including, but not limited to, risks related to: our financial condition and need for additional capital; our development work; cost and success of our product development activities and preclinical and clinical trials; commercializing any approved pharmaceutical product developed using our AI platform technology, including the rate and degree of market acceptance of our product candidates; our dependence on third parties including for conduct of clinical testing and product manufacture; our inability to enter into partnerships; government regulation; protection of our intellectual property rights; employee matters and managing growth; our ADSs and ordinary shares, the impact of international economic, political, legal, compliance, social and business factors, including inflation, and the effects on our business from other significant geopolitical and macro-economic events; and other uncertainties affecting our business operations and financial condition. For a further discussion of these risks, please refer to the risk factors included in our most recent Annual Report on Form 20-F and other filings with the U.S. Securities and Exchange Commission (SEC), which are available at www.sec.gov. We do not assume any obligation to update any forward-looking statements except as required by law.

This presentation includes statistical and other industry and market data that we obtained from industry publications and research, surveys and studies conducted by third parties or us. Industry publications and third-party research, surveys and studies generally indicate that their information has been obtained from sources believed to be reliable, although they do not guarantee the accuracy or completeness of such information. All of the market data used in this presentation involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. While we believe these industry publications and third-party research, surveys and studies are reliable, we have not independently verified such data. The industry in which we operate is subject to a high degree of uncertainty, change, and risk due to a variety of factors, which could cause results to differ materially from those expressed in the estimates made by the independent parties and by us.

EVAXION

A pioneer in fast and effective AI-powered development of new medicines



Multidisciplinary capability set and state of the art

wetlab and animal facilities



Broad pipeline
Preclinical and clinical
programs across cancer

and infectious diseases



Platform and pipeline provides unique avenues for value creation



Multi-partner approach to value realization with several partnerships in place



Al-Immunology™ Clinically validated and leading Al platform

Deriving value from both our platform and pipeline

PLATFORM



Enables high value/low risk-partnerships on target discovery, design and development agreements PLUS fast and cost-effective replenishing of internal pipeline

- One stop-shop for pharmaceutical and biotech companies
- Full capability set
- Modality agnostic
- Scalable to multiple disease areas

PIPELINE



Advancing select high value programs into preclinical and early clinical development, retaining value and leveraging multidisciplinary capability base

- Fast and effective preclinical validation
- Preclinical and clinical development capabilities
- Flexible in-house development to maximize value and monetization of assets

A transformative partnership with MSD



Option and license agreement covering EVX-B2 and EVX-B3, offers fast and effective development to address serious unmet needs, no approved vaccines available today



Significant financial and strategic value to Evaxion, both short- and long-term



Upfront payment of \$3.2 million received in October 2024 and up to \$10 million in 2025, contingent upon MSD* exercising its option to license either one or both candidates



Milestone payments of up to potentially \$592 million per product plus royalties on sales, providing a very important source of income and funding for the years ahead



Massive validation of Al-Immunology™ and pipeline from the world leader in vaccine development and commercialization



Further solidifies our collaboration with MSD, who is already our single largest shareholder with close to 20% equity stake





Targeting significant unmet needs and large markets

Major challenges

- 1 in 5 develop cancer in a lifetime¹. Lack of effective treatments for many cancer patients
- No approved vaccines against S. aureus, Gonorrhoea or Cytomegalovirus (CMV)
- Antibiotics resistance and healthcare burden continues to increase: 4.7 million deaths associated with resistance in 2021, expected to grow to 8.2 million in 2050⁶

10 million

annual deaths from cancer¹

7.8 million

annual deaths from infectious diseases²

Global market forecast

\$277bn

Cancer immunotherapy market estimated to grow to \$277 billion by 2030³

\$7.4bn

Melanoma market estimated to grow to \$7.4 billion by 20294

\$67.5bn

Infectious disease vaccines market expected to reach \$67.5 billion by 2031⁵

The power of Al-Immunology™ in vaccine target discovery, design and development

VALIDATED

clinical trials confirm predictive capabilities and link to outcomes

SCALABLE

different diseases can

AGNOSTIC

different modalities (protein, DNA, mRNA, peptides) validated



Saving and improving lives with Al-Immunology™

ACCURATE

FAST

24hrs to complete target discovery

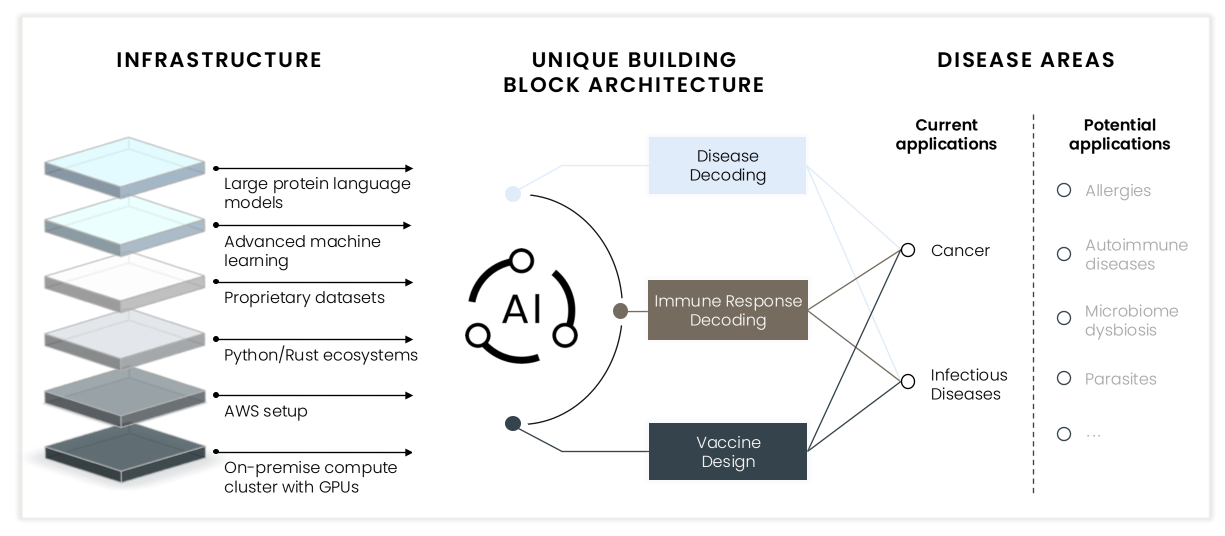
TARGETED

80% fewer targets needs testing

COST EFFECTIVE

on average cheaper target discovery versus reverse vaccinology

The unique architecture of Al-Immunology™



EVAXION

© Evaxion A/S. All rights reserved.

Multidisciplinary capability set and state of the art facilities

Disease Biology









Target Discovery

Clinical Development





Vaccine target discovery, design & development



Antigen Design

CMC





Vaccine & Adjuvant Design

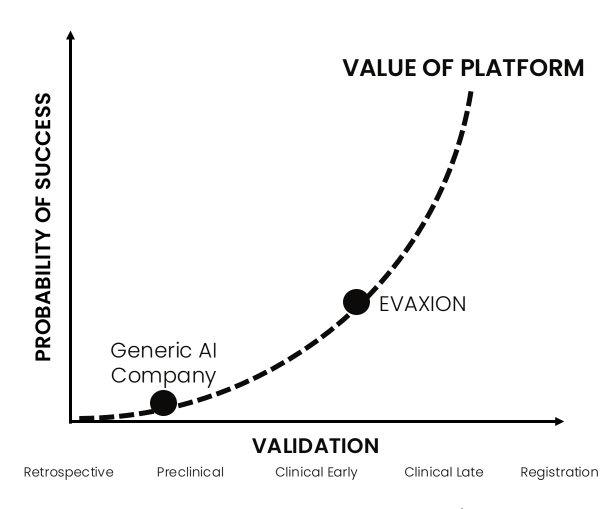






Al-Immunology™ and our multidisciplinary capability set drive differentiation

- We believe our multidisciplinary capability set allows for:
 - Continuous iterative learning loops
 - Ongoing expansion of data sets with proprietary data
 - Rapid validation of Al predictions
 - Full control of process from idea to validation
 - Continued expansion of pipeline assets
- Significantly enhancing the value of our platform



Pipeline: Demonstrating the performance and scalability of our Al-Immunology™ platform

CANDIDATE	INDICATION/ PATHOGEN	PARTNER	STAGE OF DEVELOPMENT			
			TARGET DISCOVERY	PRECLINICAL	PHASE 1	PHASE 2
CANCER						
EVX-01* (Liposomal/peptide)	Advanced melanoma					
EVX-02** (DNA)	Adjuvant melanoma					
EVX-03 (Targeted DNA)	TBD					
Multiple candidates	Undisclosed					
INFECTIOUS DI	SEASES					
EVX-B1 (Proteins)	S. aureus					
EVX-B2 (Proteins)	N. gonorrhoeae	♦ MSD				
EVX-B2*** (mRNA)	N. gonorrhoeae	Afrigen Biologica 8 Voccines In a mare North 8 OC Cansor				
EVX-B3	Bacterial pathogen	♦ MSD				
EVX-B4	Group A Streptococcus				* Pembrolizumab supply agreement	aith MSD
EVX-V1	Cytomegalovirus				(Tradename of Merck & Co., Inc., Rah) ** The data generated in the EVX-02	vay, NJ, USA)
Multiple candidates	Undisclosed				development of the second generati *** Collaboration with Afrigen for low-	on EVX-03 DNA vaccine

EVX-01: Peptide-based personalized cancer vaccine

HIGHLIGHTS

- Personalized vaccine for first-line treatment of advanced melanoma (skin cancer)
- Vaccine targets (neoantigens) identified by Al-Immunology™ based on individual tumor profile
- Combined with an anti-PDI antibody with the aim of improving clinical outcome
- Well-tolerated in all patients
- Next milestone: Two-year phase 2 clinical efficacy readout

KEY DATA

69%

Overall Response Rate (ORR) in phase 2 trial¹ 80%

of vaccine targets induced a specific immune response²

EVX-01 Phase 13



EVX-01 Phase 21



KEYTRUDA® REG. TRIAL⁴

ORR* 33%

STAGE OF DEVELOPMENT

TARGET DISCOVERY	PRECLINICAL	PHASE 1	PHASE 2

UNMET MEDICAL NEED

52%

of advanced melanoma patients survived five years with combined immunotherapy⁵ 510,000

new melanoma cases globally by 2040⁶

EVAXION

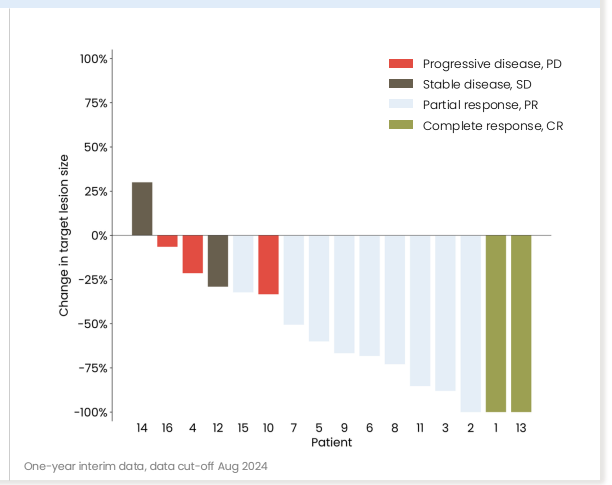
- https://www.evaxion.ai/scientific-posters/esmo-2024
- 2. https://evaxion.ai/scientific-posters/aacr-2025
- 3. https://pubmed.ncbi.nlm.nih.gov/38782542/

- 4. https://www.nejmorg/doi/full/10.1056/NEJMoa1503093. The trial is not a head-to-head evaluation of EVX-01 in combination with anti-PD-1 therapy against standard-of-care
- 5. Larkin J., New England Journal of Medicine, 2019
- 6. <u>Arnold. M. JAMA Dermatology, 2022</u>

EVX-01: Peptide-based personalized cancer vaccine

KEY DATA

- The combination of EVX-01 and anti-PD-1 therapy led to an Overall Response Rate of 69% (Objective Response in 11 out of 16 patients)
- In 15 out of the 16 patients, the tumor target lesions were reduced
- 3 out of 16 (19%) patients achieved complete remission of tumor target lesions



EVX-02: DNA-based personalized cancer vaccine

HIGHLIGHTS

- Personalized vaccine for treatment of fully resected melanoma (skin cancer) as adjuvant therapy
- Aims at preventing tumor recurrence by actively inducing tumor-specific T cells
- Carries 13 tumor-specific Al-Immunology™ identified neoantigens delivered to each patient
- Combined with an anti-PDI antibody with the aim of improving clinical outcome
- Well-tolerated in all patients

KEY DATA

100%

of patients were relapse-free at last assessment*

100%

of patients mounted an EVX-02 specific T-cell response



Immune responses were mediated by activated CD4+ & CD8+ T cells



Neoantigens with higher Al-Immunology™ scores were more likely to be immunogenic

STAGE OF DEVELOPMENT

TARGET DISCOVERY	PRECLINICAL	PHASE 1	PHASE 2

UNMET MEDICAL NEED

63%

relapse-free survival rate 12month for immunotherapy treatment¹



510,000

new melanoma cases globally by 2040²

EVAXION

- Larkin J., New England Journal of Medicine, 2019
- Arnold. M, JAMA Dermatology, 2022
- * 10 patients receiving all eight EVX-02 doses in combination with Nivolumab

EVX-03: Targeted DNA personalized cancer vaccine

HIGHLIGHTS

- Personalized vaccine for treatment of solid tumors (TBD)
- The first ever vaccine combining personalized neoantigens and endogenous retrovirus (ERV) antigens
- Contains a proprietary targeting unit that directs the antigens to the relevant immune cells
- GLP toxicology study completed without concerns
- Next milestone: Regulatory submission (CTA/IND* ready)

KEY DATA



Induces antitumor immunity with complete responses in mouse tumor model



Induces strong and durable neoantigen- and ERV-specific Tcell responses



Additive effect of combining with anti-PDI in mice



Antitumor effect obtained in both prophylactic and therapeutic models

STAGE OF DEVELOPMENT

OTAGE OF DEVELOT WILIT			
TARGET DISCOVERY	PRECLINICAL	PHASE 1	PHASE 2

UNMET MEDICAL NEED

9,7 million

global cancer associated deaths in 2022¹



35 million

new cancer cases are estimated annually worldwide by 2050¹

EVAYION.

Clinical Trial Application/Investigational New Drug Application
Bay et al. CA. A Cancer Journal for Clinicians, 2024

EVX-B1: Staphylococcus aureus (SA) vaccine

HIGHLIGHTS

- Multi-component vaccine for prevention of Skin and Soft Tissue Infections (SSTI)
- Designed to act against SA across multiple parameters, ensuring high efficacy, high immunogenicity and targeting of functionality
- Targets essential toxins and major virulence factors widely present and conserved in relevant clinical strains
- No prophylactic SA vaccine available today
- Next milestone: CMC* and toxicology

KEY DATA



Protects against surgical site infections in minipigs



Protects against lethal sepsis and skin infections in mice



Immunized mice clear the infection from internal organs



Induces long lasting immune response in mice

STAGE OF DEVELOPMENT

TARGET DISCOVERY PRECLINICAL PHASE 1 PHASE 2

UNMET MEDICAL NEED

550,000

global deaths associated with (methicillin-resistant SA (MRSA)¹



of bloodstream infections were due to MRSA in 2021²

EVAXION

- Antimicrobial Resistance Collaborators, Lancet, 2024
- 2. <u>WHO</u>
- * Chemistry, manufacturing and controls

EVX-B2: Gonorrhea vaccine (protein and mRNA)

HIGHLIGHTS

- Multi-component prophylactic vaccine against Gonorrhea
- Contains antigens that are present in all 1,200 published N. gonorrhoeae (Ng) genomes analyzed. This implies efficacy against bacteria strains globally
- Unique Mode of Action; kills off bacteria by targeting its cell division processes
- No prophylactic Gonorrhea vaccine available today

KEY DATA



Demonstrates strong efficacy against 50 clinically relevant Ng strains



Protects against Ng in a mouse infectious model



Induces strong humoral and cellular immune responses



Established Mode of Action ->
antibody-dependent complement
mediated killing

TARGET DISCOVERY Proteins Proteins Afrigen MSD

UNMET MEDICAL NEED

82.4 million

new infections caused by *N. gonorrhoeae* in 2020 globally¹

Resistance

is escalating. *N. gonorrhoeae* has developed resistance to every class of antibiotics used to treat it²

EVAXION

U.S. Centers for Disease Control and Prevention
U.S. Centers for Disease Control and Prevention

EVX-B3: Vaccine against bacterial pathogen

HIGHLIGHTS

- Aims to address a serious global medical issue by targeting a pathogen responsible for recurrent infections, increasing incidence, and often severe medical complications, for which no vaccine currently exists
- Project initiated in September 2023 as a collaboration with MSD. In September 2024, Evaxion and MSD entered an option and license agreement for EVX-B3 and EVX-B2
- First phases of the collaboration have been successfully completed







EVX-V1: Cytomegalovirus (CMV) vaccine

HIGHLIGHTS

- Multi-component vaccine for prevention of CMV infection in connection with e.g. solid organ transplantations
- Contains novel B- and T-cell antigens and a proprietary pre-fusion glycoprotein B (gB) construct
- No CMV vaccine approved to date
- **Next milestone:** Lead antigens selection

KEY DATA



Al-Immunology™ identified novel B-cell antigens induce a strong humoral immune response



Al-Immunology™ identified T-cell epitopes induce a strong cellular immune response



Pre-fusion qB immune serum significantly neutralizes CMV infection

STAGE OF DEVELOPMENT TARGET **PRECLINICAL** PHASE 2 PHASE 1 DISCOVERY

UNMET MEDICAL NEED

1 in 3 children

in the U.S is already infected with CMV by age 51



60/90%

of adults globally are infected with CMV2 - up to millions get serious complications annually

EVAXION

U.S. Centers for Disease Control and Prevention

© Evaxion A/S. All rights reserved.

Introduction

Strategic ambition and imperatives

Strategy

Maintain the leading Al platform

Generate positive cash flow Drive best-in-class target discovery and validation Leading **AI-powered** TechBio company Develop novel R&D pipeline Become Al partner of choice

Introduction

Al Immunology™

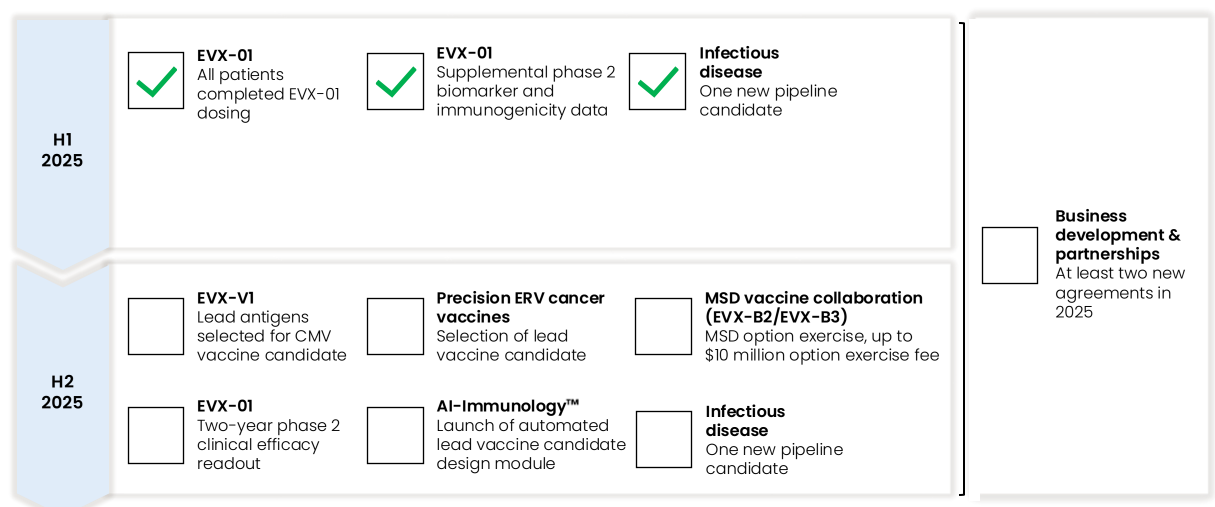
Pipeline

Leadership team

Summary

2025 milestones and value catalysts

Strategy



Strong leadership with extensive experience across all relevant fields



Interim Chief Executive Officer & Chief Scientific Officer

Birgitte Rønø

MSc Human Biology/ PhD







Chief AI Officer & Evaxion Founder

Andreas Mattsson

MSc Bioinformatics







Chief Financial

Thomas Schmidt

MSc Business Economics & Auditing





Board of directors

- Marianne Søgaard
 Chair, former tech lawyer and equity partner
- Roberto Prego
 Former Teva (head of Latin America)
- Lars Holtug
 Certified Public Accountant
- Lars Staal Wegner
 VP Business Development,
 Hengrui Pharmaceuticals, MD
- Helen Tayton-Martin
 Former Adaptimmune Therapeutics plc (COO/Co-founder)

Introduction

Al Immunology

Pipeline

Strategy Leadership team

Summary

Investment highlights

- Truly Al-first company leveraging Al-Immunology™

 a pioneering clinically validated Al platform for vaccine discovery, design and development. Its modular architecture allows for unique scalability
- Pipeline of novel clinical and preclinical vaccine candidates for cancers and infectious diseases
- Proven ability to establish and manage a range of valuecreating partnerships
- Clear strategy with strong focus on monetizing value through business development
- Very solid business development pipeline
- MSD (via its MSD GHIF venture capital arm) largest shareholder with close to 20% equity stake

Capital structure

Symbol (Nasdaq - ADS)	EVAX
Stock price (as of June 30, 2025)	\$2.45
ADS outstanding if full conversion	6.3m
Market capitalization	\$15.4m
Warrants ¹ (WAEP \$10.48)	3.1m
Average trading volume	105,017
Cash ²	\$17.8m
Debt ²	\$8m

1 Warrants convertible into ADS 2 As of Mar 31, 2025

