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Investment Highlights

Key player in advanced protein sciences with novel pipeline addressing 45B EUR markets



Leader in production of complex proteins with the advantageous ExpreS² technology



Co-Founder of AdaptVac ApS, owner of a unique Virus Like Particle (VLP) technology



Pipeline of therapeutics/vaccines, addressing high-need and attractive markets



Revenue of 15M SEK / ~1.5M EUR with >10% growth from legacy service contract business



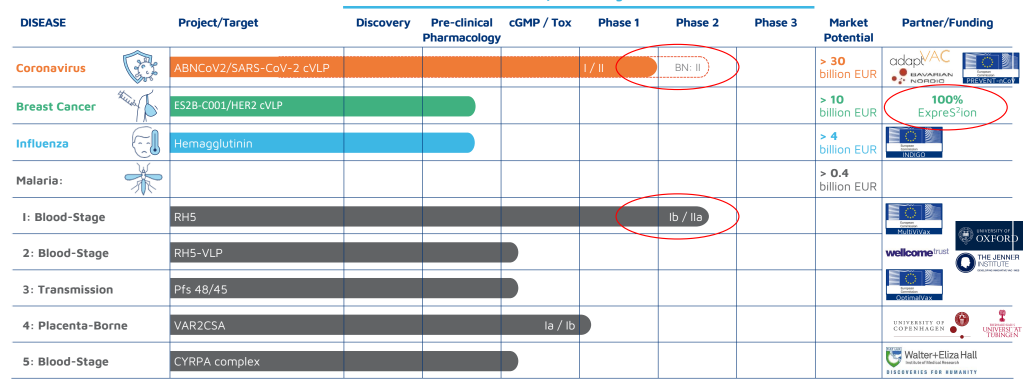
NASDAQ First North GM Stockholm [EXPRS2]. >12x increase in share price since 01/2020

Market Cap: >1.4B SEK / >135M EUR



Deep Pipeline for Value Creation

Development Progress



AdaptVac is a joint venture between ExpreS2ion (34% owned) and NextGen Vaccines (66% owned)

Significant events in 2021



Management Team

Expanded team in 2021 brings skills to build our pipeline-focused business



Bent U. Frandsen, CEO

- MSc. In Finance/Strategic Management, Copenhagen Business School Denmark
- Born 1967, Danish citizen
- >25 years industry finance, business dev and management experience













Dr. Mette Thorn, VP Preclinical Development

- PhD in Immunology, and a MSc in Chem Eng., Tech. Univ of Denmark
- Born 1972, Danish citizen
- 20 years industrial research experience















Keith Alexander, CFO

- MBA, The Wharton School and the University of Pennsylvania, USA
- Born 1975, American citizen with Danish permanent residence
- >20 years of equity research, corporate strategy, asset management and consulting experience

Danske Bank J.P.Morgan accenture



Prof. Lars Petersen, Medical Dir., Oncology

- MD, DMSc in immuno-pharmacology, from Univ of Copenhagen, and CBA from AVT Business School
- Born 1960, Danish citizen
- >30 years academic and clinical development experience











Max Soegaard, VP of R&D and Technology

- PhD in Biochem., UCL, UK, and MSc in Molecular Biology; AU, Denmark
- Born 1970. Danish citizen
- 20 years academic and industrial research experience



MOLECULAR BIOPHYSICS SUITE DEPARTMENT OF BIOCHEMISTRY



Eske Rvgaard-Hialsted. VP Business Dev.

- MSc in Molecular Biology from Technical Univ. of Denmark (DTU)
- Born 1965, Danish citizen
- > 25 years across business dev, sales and marketing in life sciences









Board of Directors

Expanded the Board in 2021 in support of the transition to a pipeline-focused business



Dr. Martin Roland Jensen, Chairman

- PhD. in Molecular and Cell Biology, Univ. of Copenhagen, Denmark
- Born 1960. Danish citizen
- >35 years biotech industry management and co-founder experience, incl. scientific work in immunology and cancer vaccine development













Dr. Karin Garre, Board Member

Elected in 2021

- MD, from University of Copenhagen, Denmark
- Born 1957, Danish citizen
- >25 years bio-industry management and drug development experience from early to late-stage phases and registration















Dr. Allan Rosetzsky, Board Member

- Doctor of Medicine (MD), from University of Copenhagen, Denmark
- Born 1948, Danish citizen
- >40 years of healthcare and biopharma experience, including founding, running, and successfully selling the clinical CRO KLIFO



RHÔNE-POULENC





Sara Sande, Board Member

Elected in 2021

- MSc in Economics, from University of Copenhagen , Denmark
- Born 1975, Danish citizen
- 20 years leadership experience in high-tech B2B companies, incl. sales excellence, strategy and commercial development













Jakob Knudsen, Board Member

- Law Degree from Univ. of Copenhagen, and MBA, Imperial College, UK
- Born 1968. Danish citizen
- >25 years commercial experience from international biotech industry



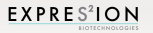




Board update and expansion at AGM May 2021

 Combined more than 140 years of deep professional experience that supports ExpreS²ion's vision of leadership in the infectious diseases and cancer fields





The Most Common Cancer

1 in 8

women will be diagnosed with invasive breast cancer in her lifetime

~25%

have overexpression of HER2 receptors, associated with more aggressive tumors and reduced survival²

685,000

deaths worldwide in 2020 due to breast cancer¹

BREAST CANCER

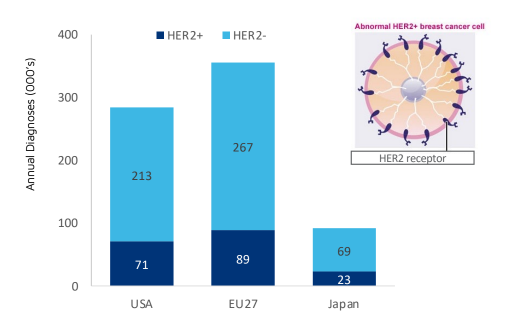




HER2+ Breast Cancer Overview

The ES2B-C001 vaccine can offer significant benefits compared to current treatment options

Over 180,000 people diagnosed with HER2+ breast cancer per year across US, EU, & Japan^{1,2}



Monoclonal antibodies are the cornerstone of treatment for HER2+ breast cancer (>\$7B USD sales)

 Target the HER2 receptor on tumor cells to reduce proliferation and induce tumor cell destruction





However, serious drawbacks exist with these therapies

- <u>Resistance</u> to monoclonal antibodies may develop
- Potential for cardiac toxicity
- Repeated administration required: 28 day half-life requires administration every 3rd week until remission or resistance develops, costs \$30-\$50k USD



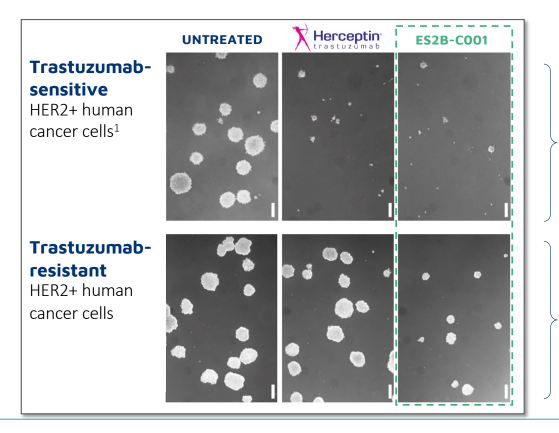
^{1.} US: BreastCancer.org: https://www.breastcancer.org/symptoms/understand_bc/statistics; EU27: Information System (Oct 2020) (https://ecis.jrc.ec.europa.eu/pdf/Breast_cancer_factsheet-Oct_2020.pdf); Japan: https://gco.iarc.fr/today/data/factsheets/populations/392-japan-fact-sheets.pdf.

2. Mitri Z et al. The HER2 Receptor in Breast Cancer: Pathophysiology, Clinical Use, and New Advances in Therapy. Chemother Res Pract. 2012; 2012: 743193



ES2B-C001 overcomes Herceptin resistance

The soft agar human cancer cell growth inhibition assay provides in vitro evidence



Both Herceptin (trastuzumab) and ES2B-C001 inhibited growth in the trastuzumab-sensitive cells

Only ES2B-C001 inhibited growth in the trastuzumab-resistant cells; cells were unresponsive to Herceptin

BREAST CANCER



Strong Preclinical Data for VLP Approach

ES2B-C001 has demonstrated animal proof-of-concept, and on track to repeat in vivo PoC

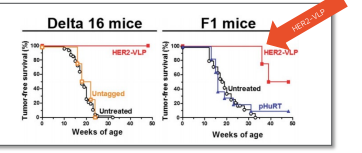
- Prevention of 50-100% of spontaneous mammary carcinogenesis
- Strong tumor growth inhibition in therapeutic studies (mice transplanted with tumor cells/fragments)

Preclinical *in vivo* studies are underway in collaboration with University of Bologna; proof-of-concept data expected primo 2022.

On path for clinical trial application submission before end of 2022.

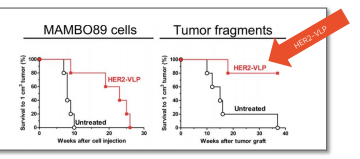


(mice with pre-disposition to spontaneous development of HER2+ tumors)



Therapeutic studies

(mice transplanted with HER2+ tumor cells or larger tumor fragments prior to vaccination)



BREAST CANCER



Influenza & Malaria



Influenza Vaccine

>4 billion EUR

The INDIGO consortium

- Led by University of Amsterdam
- Multiple research groups, incl. ExpreS²ion
- Funded by a 10 MEUR 2020 Horizon grant from the EU (0.6 MEUR awarded to ExpreS²ion)

Technologies

- Use of ExpreS² platform for antigen production
- Goal of >90% responder rate (vs <40% with current vaccines

Vaccine design completed - Lead candidate selection

• Progression towards preclinical activities – affected by the COVID-19 pandemic



Malaria Vaccine

>0.4 billion EUR

5 vaccines candidates under development that target various stages of disease & transmission

Partners

Stage/Target

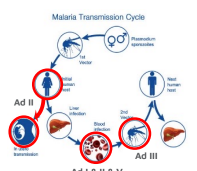
Blood stage (RH5.1)

Blood stage (RH5.2)

Transmission (Pfs48/45)

V. Blood-stage (PfRipr)





Ad I) 2021 news on RH5.1

Placenta borne (VAR2CSA)

- 04.21: Publication of Phase I/IIa data from the VAC063 study
- 07.21: The VAC080 study, a Phase Ib trial, is initiated in 60 healthy adults and infants in Tanzania to assess safety and immunogenicity

I 12 **Proteins** for Life



The 2nd Generation COVID-19 Vaccine

With **over 4.6 million deaths worldwide**, significant needs remain in the global long-term fight against the SARS-CoV-2 virus:



Uncertain duration of effect with current vaccines, expected to need repeated boosters



Storage and handling requirements for many vaccines create logistical constraints



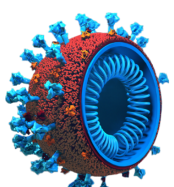
Potential mutated variants may require rapid development of new vaccines





The Best COVID-19 Vaccine

ABNCoV2 has demonstrated superior preclinical proof-of-concept, and now promising human data

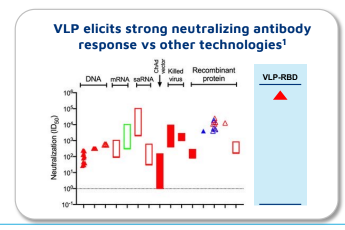


Virus

Spike proteins on surface of the coronavirus are primary target for vaccine development

Encouraging early findings:

- Durable immune response with single shot
- Strong immunogenicity vs. variants
- Well suited to rapid iteration for mutated variants if needed
- Stability at room temperature*



Phase I/II Study headline results:

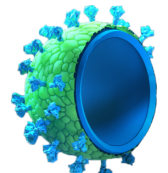
- 45 humans dosed (6-70μg)
- Aug. '21: Safe and well tolerated
- High levels of neutralizing antibodies, also for Delta/Beta VoCs

See data next slide

Bavarian Nordic holds the exclusive global license to ABNCoV2; sponsor of the on-going commercialisation



- Phase II readout within 2021
- Phase III initiation in 2022 with market launch estimated 2022/-23



Capsid VLP

Spike proteins displayed on surface but contains no genetic material

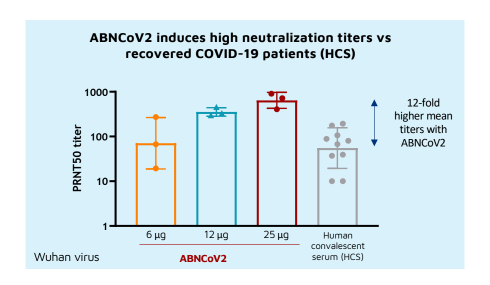
Proteins for Life

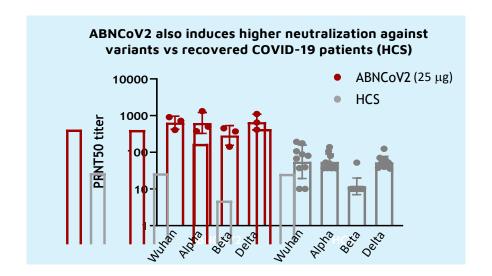
*ABNCoV2 has been proven to endure storage at room temperature for 3 months and no changes were detected even after shorter incubation at 49 degrees Celsius.



ABNCoV2: Positive Phase I/II Outcomes

Exceptional safety & tolerability, as well as high neutralizing effect against variants





Results support initiation of 210-subject Phase II booster study (results Q4 2021) and parallel ramp-up for Phase III in early 2022 (with up to DKK 800 million funding by Danish Ministry of Health)



COVID-19 License and JV Economics

ABNCoV2 is already out-licensed with near-term revenue streams supporting ExpreS²ion

AdaptVac's Economics

Paid by Bavarian Nordic

- 4 MEUR upfront (paid in July 2020)
- Up to 136 MEUR in development and sales milestones

 Single- to double-digit-% royalties of Bavarian revenues

ExpreS²ion's Economics

Paid by AdaptVac

- 34% ownership of AdaptVac
- Up to 2 MEUR in commercial milestone payments
- Lower double-digit percentage of AdaptVac royalties



Carnegie

Estimated COVID-19 + Adapt Vac value¹

Nordea

Analysguiden

Danske Bank

2,800

2,500

2,200

1,900

1,600

1,000

SEK millions

COVID-19 Value to ExpreS²ion

Institutional analysts have higher sales and approval assumptions



Pareto: SEK 68 target COVID-19 + AdaptVac value: **SEK 1,322 mn** (60.9% of company valuation)

Retail

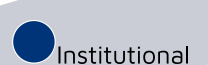


SEK 2,610 mn

Carnegie

SEK 2,596 mn

SEK 1,942 mn



Estimated COVID-19 + AdaptVac value¹

Nordea

Danske Bank





CORONAVIRUS

Analysguiden: SEK 55 target COVID-19 + AdaptVac value: SEK 1,183 mn (64.4%)





Exercise of Warrant Programme TO5

Window open during September 6-20 – Strike price determined to be 25 SEK / share

- 5.5 million TO5 warrants, part of the October 2020 successfully oversubscribed rights issue
- Exercise window September 6-20, 2021
- Strike price equal to 70% of VWAP during 10 trading days prior to exercise window
- Strike price must be within window of SEK 6-25 per share – determined to be 25 SEK
- 3 warrants equal 1 share
- Potential SEK 45 million cash inflow in gross proceeds



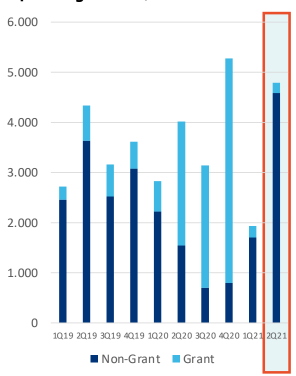


Proteins for Life VWAP = Volume-Weighted Average Price

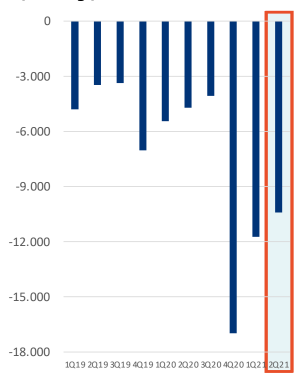


2Q21 - Key Financial Developments

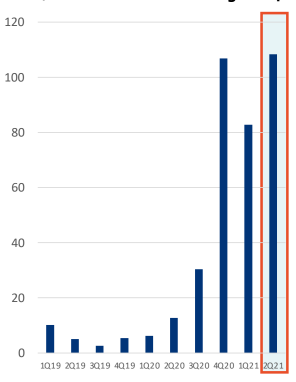
Operating income, SEK '000s



Operating profit (loss), SEK '000s



Cash, SEK millions - excluding TO5 proceeds





2021 – 2023 Outlook

On track to deliver shareholder value

	2021 CORONAVIRUS (ABNCoV2)								2023		
Care L											
₩.	♥ Phase I/II trial, COUGH-1 initiated		○ COUGH-1 full safety & efficacy results (Q3)	Ø BN Phase II trial initiation (Q3)	BN Phase II trial readout	BN Phas trial initiation		BN Phase II initial readout		t launch t to ory	
The state of the s	BREAST CA	ANCER (ES2B	-C001)								
1 //	Executed in-licensing (Feb 2021)	Preclinical animal stu initiated (6)	dies		Preclinic proof-of- results		GMP manufacti batch & to	uring clin	ng of ical trial blication	Initiation of first human clinical trial	Outlicensing window opens pending human data
	INFLUENZ	A									
		in pred	n INDIGO progress clinical animal s in (H2)	developme	upport further ent of one or idates in 2021						
	MALARIA										
V	Phase IIa results from the Rh5.1 vaccine published in 2021			Additional phase I trial in a malaria endemic region in Africa launched during 2021, with alternative adjuvant						Rh5 phase I readout	trial

