



Investor Presentation

*Bryan Garnier Healthcare Conference
November 17, 2020*



*...at Hansa Biopharma we envision a world where all patients
with rare immunologic diseases can lead long and healthy lives...*

Forward-looking statement

This presentation may contain certain forward-looking statements and forecasts based on uncertainty, since they relate to events and depend on circumstances that will occur in the future and which, by their nature, will have an impact on Hansa Biopharma's business, financial condition and results of operations. The terms "anticipates", "assumes", "believes", "can", "could", "estimates", "expects", "forecasts", "intends", "may", "might", "plans", "should", "projects", "will", "would" or, in each case, their negative, or other variations or comparable terminology are used to identify forward-looking statement. There are a number of factors that could cause actual results and developments to differ materially from those expressed or implied in a forward-looking statement or affect the extent to which a particular projection is realized. Factors that could cause these differences include, but are not limited to, implementation of Hansa Biopharma's strategy and its ability to further grow, risks associated with the development and/or approval of Hansa Biopharma's products candidates, ongoing clinical trials and expected trial results, the ability to commercialize imlifidase, technology changes and new products in Hansa Biopharma's potential market and industry, the ability to develop new products and enhance existing products, the impact of competition, changes in general economy and industry conditions and legislative, regulatory and political factors.

No assurance can be given that such expectations will prove to have been correct. Hansa Biopharma disclaims any obligation to update or revise any forward-looking statements, whether as a result of new information, future events or otherwise.

Hansa Biopharma at a glance



Company background

- Founded 2007 with HQ in Lund, Sweden
- Søren Tøulstrup, CEO – Ulf Wiinberg, Chairman
- ~80 employees (~2/3 in R&D) end of Q3 2020
- Operations in Sweden, US & across Europe
- Market cap: SEK ~11bn (~1.25 bn USD) end of September 2020
- Listed on Nasdaq OMX Stockholm (HNSA)



Leader in immunomodulatory enzymes for rare IgG-mediated diseases

- Imlifidase is a unique IgG antibody-cleaving enzyme. If approved, imlifidase may have the potential to meet a large unmet need and preserve and transform the lives of people with rare diseases
- Imlifidase has been studied in five clinical studies in kidney transplantation
- Imlifidase has been published in peer-reviewed journals (e.g. New England Journal of Medicine and the American Journal of Transplantation)



Broad pipeline in transplantation and autoimmune diseases

- Lead indication in kidney transplantation in highly sensitized patients
 - The European commission has granted conditional approval for Idefixir™ (imlifidase) in highly sensitized kidney transplant patients in the European Union
 - US: Study protocol for RCT submitted June 2020, discussions with FDA ongoing
- Anti-GBM antibody disease (Phase 2)
- Antibody mediated kidney transplant rejection (AMR) (Phase 2)
- Guillain-Barré syndrome (GBS) (Phase 2)
- NiceR - Recurring treatment in autoimmune disease, transplantation and oncology (Preclinical)
- EnzE – Cancer immunotherapy (Preclinical)



Key financials*

| | | |
|--------------------------|------------------------------------|-----------------|
| • Cash & short-term inv. | 9M'20* SEK 1.5bn (9M'19 SEK 680m) | FY'19 SEK 601m |
| • Operating Profits/Loss | 9M'20* SEK -317m (9M'19 SEK -250m) | FY'19 SEK -360m |
| • Operating cash flow | 9M'20* SEK -194m (9M'19 SEK -260) | FY'19 SEK -335m |

* Unaudited

*...at Hansa Biopharma we envision
a world where all patients with rare
immunologic diseases can lead
long and healthy lives...*

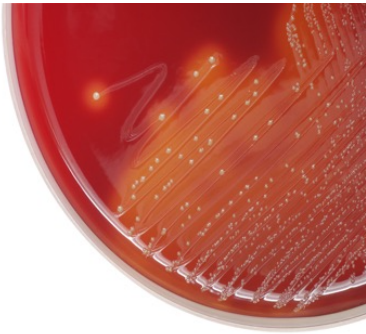


Imlifidase – a novel approach to eliminate pathogenic IgG



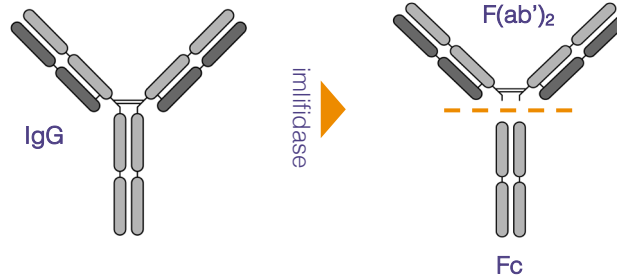
Origins from *Streptococcus pyogenes*

- Species of Gram-positive, spherical bacteria in the genus *Streptococcus*
- Usually known from causing a strep throat infection



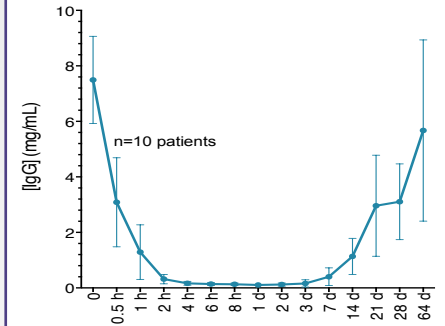
Imlifidase, a unique IgG antibody-cleaving enzyme

- Interacts with Fc-part of IgG with extremely high specificity
- Cleaves IgG at the hinge region, generating one F(ab')₂ fragment and one homo-dimeric Fc-fragment

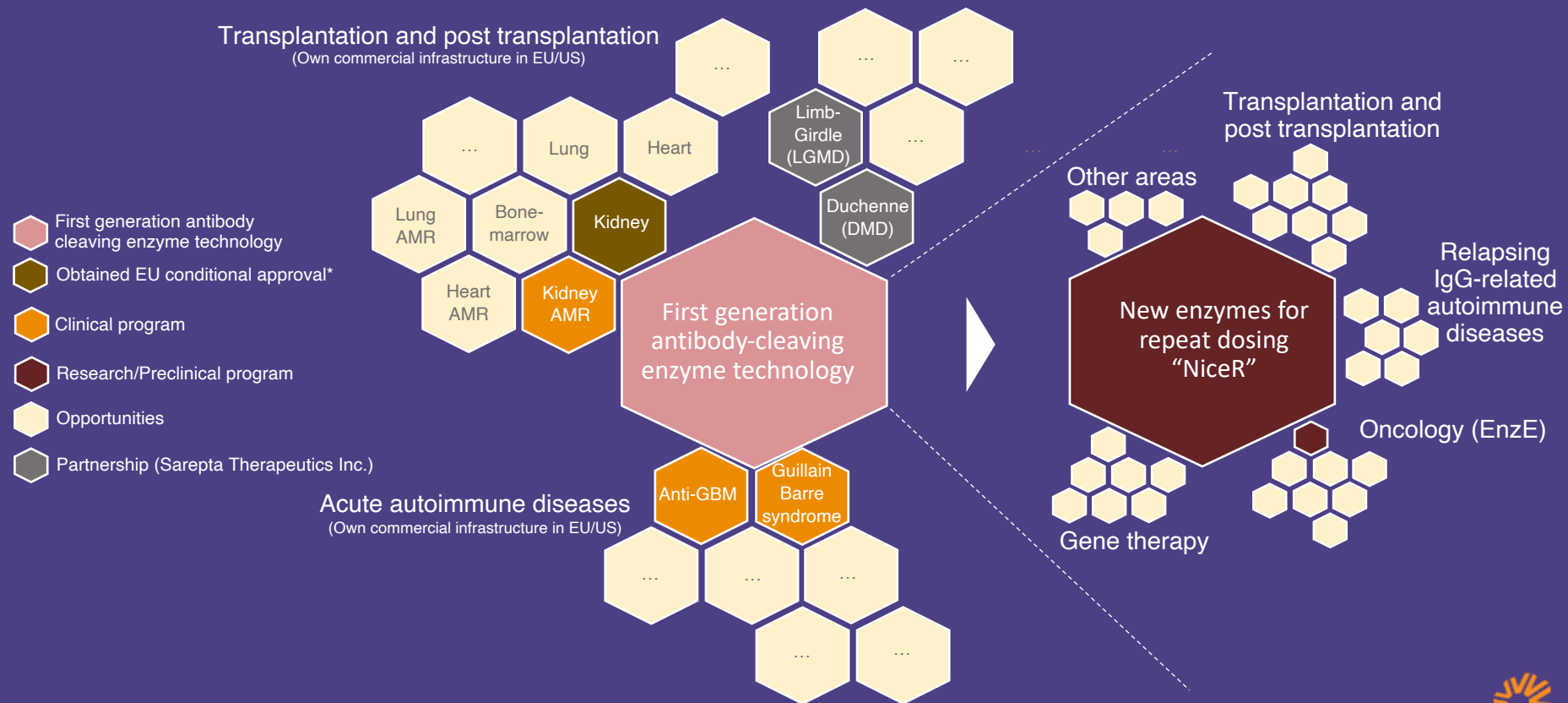


Imlifidase inactivates IgG in 2 hours

- Rapid onset of action that inactivates IgG below detectable level in 2 hours
- IgG antibody-free window for approximately one week



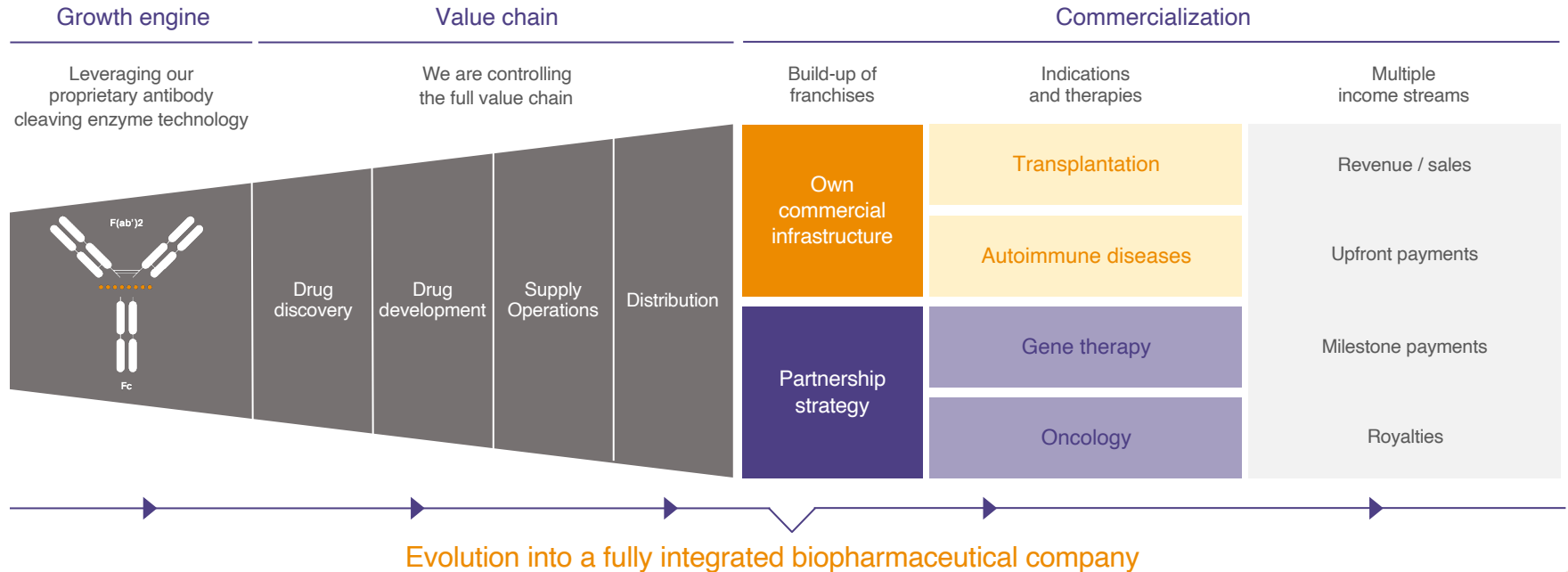
Potential indication universe



* US: Study protocol submitted June 2020, study expected to be initiated H1 2021. The new clinical study could support BLA submission by 2023

Leveraging our technology platform

Developing new therapies targeting rare diseases with unmet medical need across a range of indications



Exclusive agreement with Sarepta Therapeutics to develop and promote imlifidase as pre-treatment ahead of gene therapy in select indications

A unique opportunity to combine efforts...

...and to use the unique features of imlifidase to potentially enable gene therapy treatment in patients who today aren't eligible for these breakthrough therapies due to pre-existing neutralizing antibodies in two indications with a very high unmet medical need

Structure of the partnership

- Sarepta will be responsible for conducting
- Pre-clinical/clinical studies with imlifidase
 - Regulatory approvals
 - Promotion of imlifidase as a pre-treatment to Sarepta's gene therapies following potential approval

Hansa will supply product, support with know-how and involve in the regulatory approval process

Hansa's financial participation

Potential total deal value for Hansa amounts to up to USD ~400m plus royalties and incremental imlifidase sales



Hansa's key competences

- Leader in immunomodulatory enzyme technology for rare IgG mediated diseases
- Strong experience in antibody cleaving and desensitization
- Broad enzyme technology that can be used in a variety of indications





Sarepta's key competences

- Market leader within gene therapy targeted at muscular dystrophies
- Strong pre-clinical and clinical gene therapy portfolio
- Scientific approach and knowledge within gene therapy
- Experience with challenges of Nab-positive patients



Broad pipeline in transplantation and auto-immune diseases

| Candidate / Project | Indication | Research/ Preclinical | Phase 1 | Potentially Pivotal/ Phase 2 | Phase 3 | Marketing Authorization | Marketed | Next Anticipated Milestone |
|---------------------|--|-----------------------|---------|------------------------------|---------|-------------------------|----------|---|
| Imlifidase | EU: Kidney transplantation in highly sensitized patients ^{1,2} | | | | → | | *) | EU: Commercial launch Q4 2020 |
| | US: Kidney transplantation in highly sensitized patients ^{1,2} | | | | **)) | | | First patient dosed H1 2021 |
| | Anti-GBM antibody disease ³ | | | | | | | Next step is to engage with regulators and agree on a path forward toward BLA/MAA |
| | Antibody mediated kidney transplant rejection (AMR) | | | | | | | Complete enrolment of 30 patients H2'21 |
| | Guillain-Barré syndrome (GBS) | | | | | | | Complete enrolment of 30 patients H2'21 |
| | Limb-Girdle (LGMD) & Duchenne (DMD) (Pre-treatment ahead of gene therapy with Sarepta) | | | | | | | Research phase |
| NiceR | Recurring treatment in autoimmune disease, transplantation and oncology | | | | | | | Development of CMC process / Tox studies |
| EnzE | Cancer immunotherapy | | | | | | | Research phase |

Completed Ongoing

¹ Results from the Phase 1 study have been published, Winstedt et al. (2015) PLOS ONE 10(7)

² Lorant et al American Journal of Transplantation and 03+04 studies (Jordan et al New England Journal of Medicine)

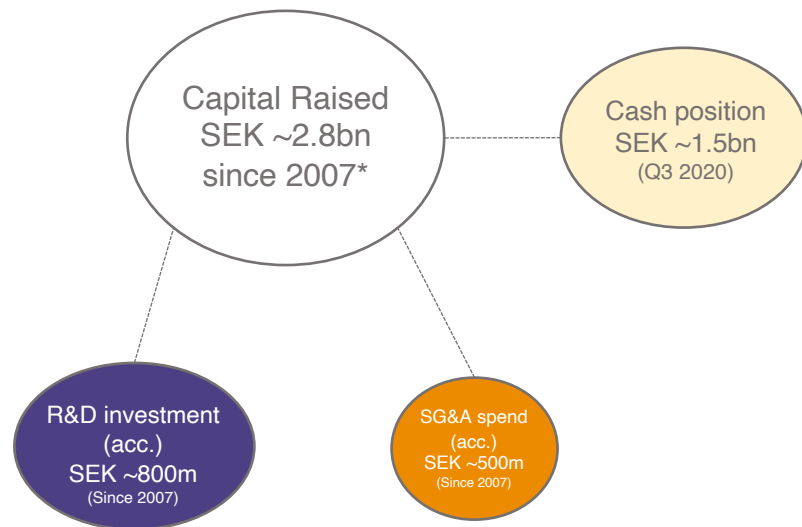
³ Investigator-initiated study by Mårten Segelmark, Professor at the universities in Linköping and Lund

*) The EU Commission has granted conditional approval for imlifidase in highly sensitized kidney transplant patients. A post-approval study will commence in parallel with the launch

**) FDA: Proposed study protocol submitted June 2020. Discussions are currently ongoing with the FDA. Once the final protocol has been agreed upon, Hansa Biopharma will proceed to set up centers in the US and start to enroll patients. Given the continued impact of the COVID-19 pandemic and the timeline for the finalization of the study protocol Hansa expect recruitment of the first patient to be in H1 2021

With the recent capital injection Hansa Biopharma is financed into 2023

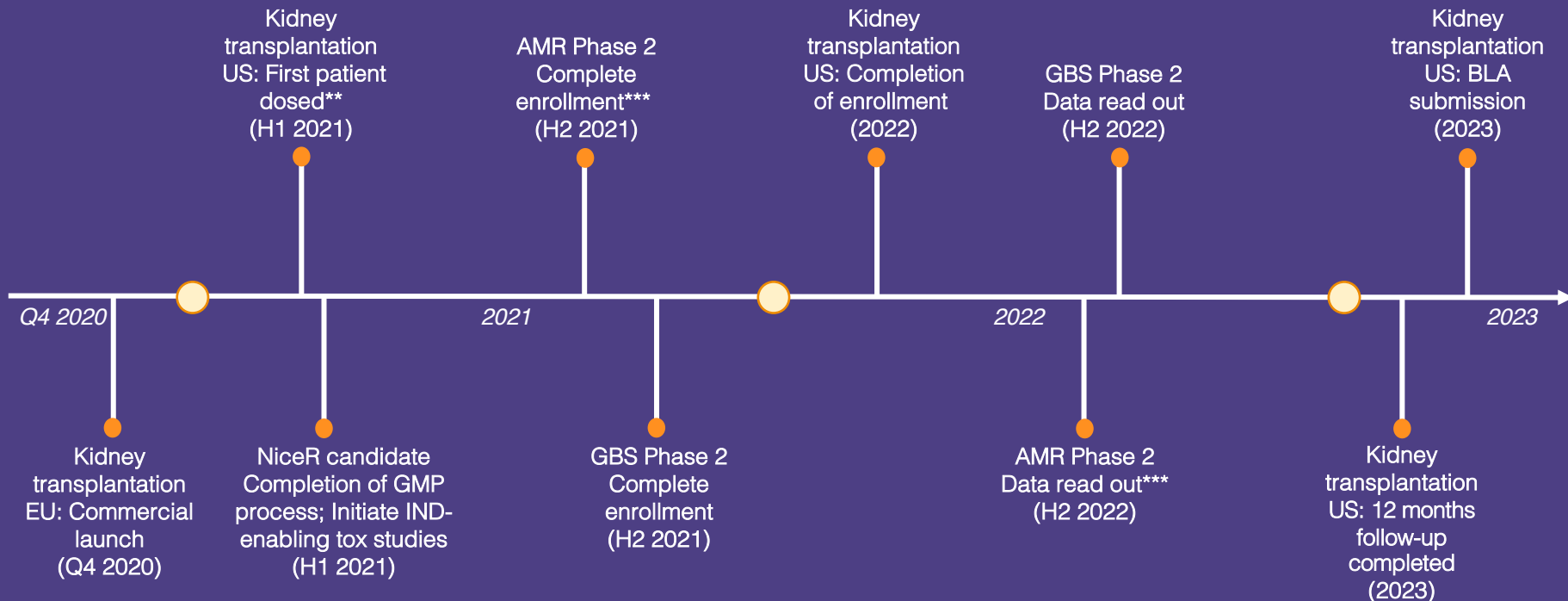
Since 2007 Hansa has mainly been backed by VCs funding the development of our enzyme platform



Capital injection from new shares (SEK 1.1bn) and Sarepta (SEK 100m) will finance Hansa into 2023



Upcoming milestones



**) FDA: Proposed study protocol submitted June 2020. Discussions are currently ongoing with the FDA. Once the final protocol has been agreed upon, Hansa Biopharma will proceed to set up centers in the US and start to enroll patients. Given the continued impact of the COVID-19 pandemic and the timeline for the finalization of the study protocol Hansa expect recruitment of the first patient to be in H1 2021

***) AMR/GBS Due to the impact from the COVID-19 pandemic, the enrollment in GBS and AMR were temporarily halted for the past six months. Hansa Biopharma expects to reinstate enrollment of these studies in Q4 2020 under a risk-based, site-by-site approach. Enrollment of patients in the AMR study is now expected to be completed in the second half of 2021, while completion of patient enrollment in the GBS study is still expected in the second half of 2021. High-level data readout for both studies are expected in the second half of 2022.

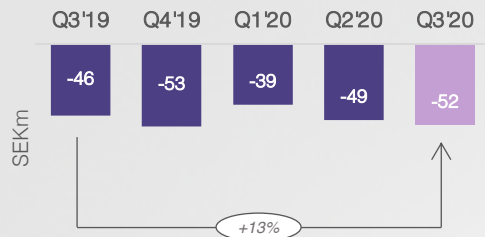


Capital Markets

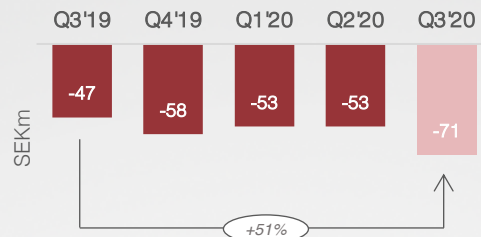


Hansa Biopharma continues to invest in the R&D pipeline and the commercial preparation towards the expected launch in Q4 2020

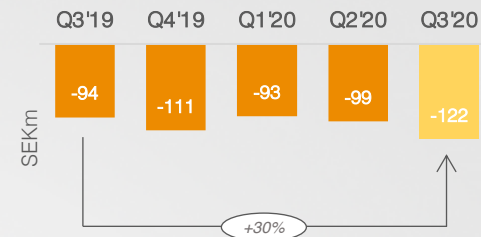
SG&A expenses (Q/Q)



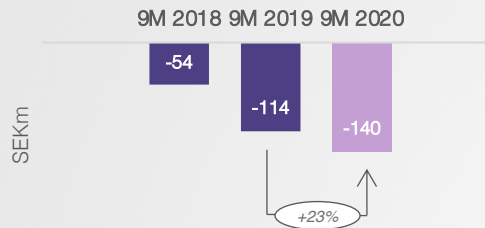
R&D expenses (Q/Q)



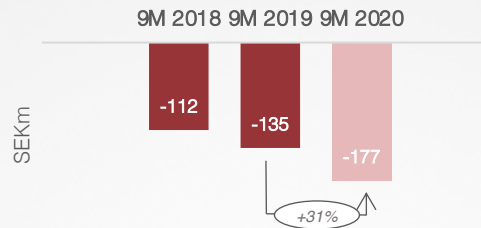
Net loss (Q/Q)



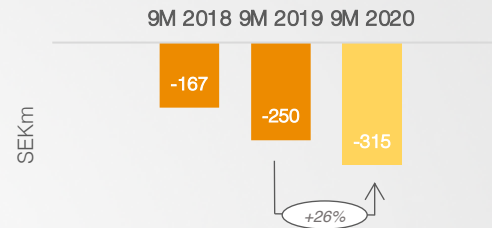
SG&A expenses (9M/9M)



R&D expenses (9M/9M)

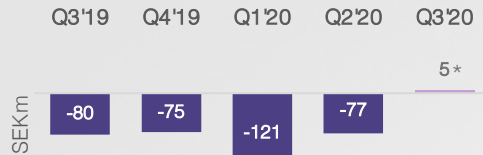


Net loss (9M/M)

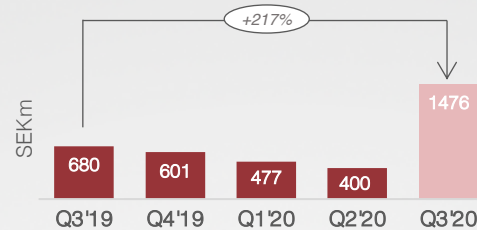


Cash position stood at SEK 1.5bn (~USD 150m) end of Q3 2020; Hansa Biopharma is financed through 2023

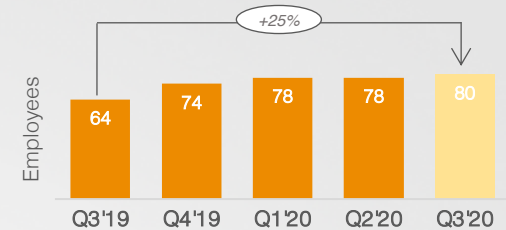
Operating cash flow (Q/Q)



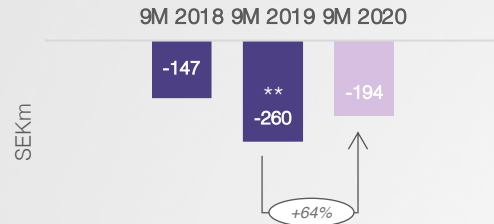
Cash & short term investments (Q/Q)



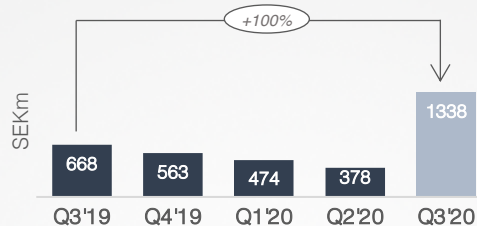
Number of employees (Q/Q)



Operating cash flow (9M/9M)



Shareholders equity (Q/Q)



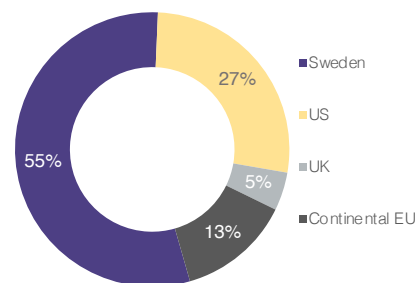
Ownership in Hansa Biopharma

Top 10 ownership as per September 30, 2020

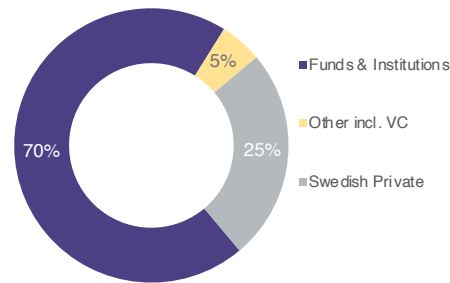
| Name | No. of shares | Ownership in pct. |
|--------------------------------------|---------------|-------------------|
| Consonance Capital Management | 2 655 009 | 6.0 |
| Redmile Group | 2 323 708 | 5.2 |
| NXT2B | 2 155 379 | 4.8 |
| Invesco | 1 938 841 | 4.4 |
| Thomas Olausson | 1 750 474 | 3.9 |
| Fourth Swedish National Pension Fund | 1 536 624 | 3.5 |
| Avanza Fonder AB | 1 387 380 | 3.1 |
| Handelsbanken Fonder AB | 1 329 744 | 3.0 |
| Gladiator | 1 025 000 | 2.3 |
| ClearBridge, LLC | 1 012 786 | 2.3 |
| Other | 27 358 507 | 61.5 |
| Outstanding A shares in total | 44 473 452 | 100.0 |

Classification of ownership

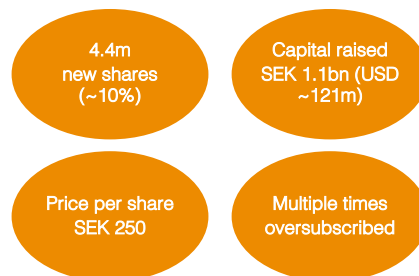
Ownership by country



Ownership by type



Capital Raise July 2020



No. of shareholders



Hansa Biopharma - Market data and share price development

Market data (Sep 2020)

Stock Exchange: Nasdaq, Stockholm since Nov 2015
(First North Oct 2007- Nov 2015)

Ticker HNSA

Market Cap: SEK ~11bn (USD ~1.25 bn)

52-week range: SEK 59-288 per share

Avg. Daily Turnover: vol ~565k shares

Shares outstanding: 44 473 452

Shareholders ~18,000

Top 5 Shareholders: Consonance Capman 6.0%
As per September 2020

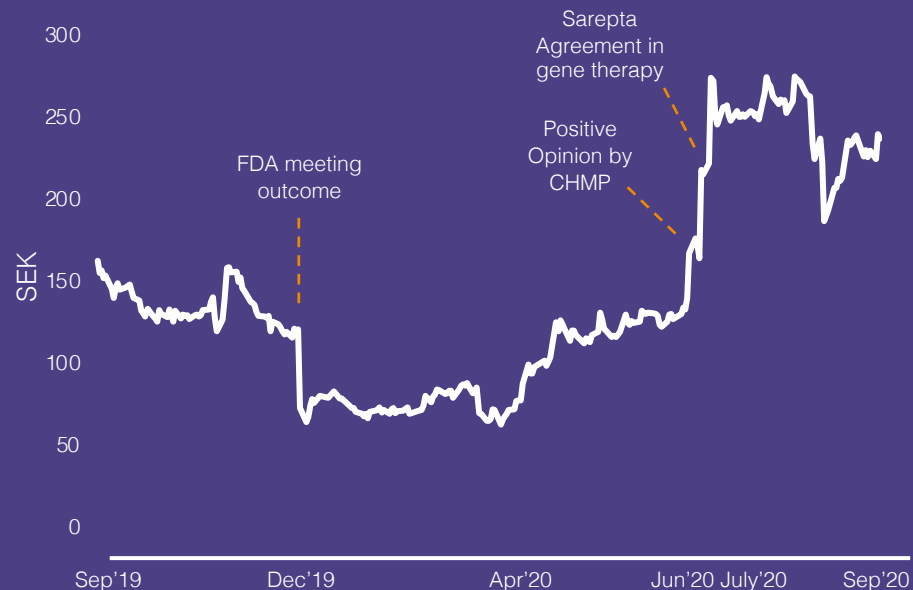
Redmile Group 5.2%

NXT2B 4.8%

Invesco 4.4%

Thomas Olausson 3.9%

12 months Share price development (Sep 2020)



Analysts covering Hansa Biopharma (ticker: HNSA, NASDAQ Stockholm)

| Analyst | Bank / Research institution (year of initiation) | Location | Email | Phone |
|---------------------|--|---------------|--|---------------------|
| Christopher Uhde | SEB (2016) | Stockholm | christopher.uhde@seb.se | +46 (0) 876-385 53 |
| Viktor Sundberg | ABG Sundal Collier (2018) | Stockholm | viktor.sundberg@abgsc.se | +46 (0) 856-628 641 |
| Charles Weston | RBC (2017) | London | Charles.Weston@rbccm.com | +44 7935 202349 |
| Ingrid Gafanhão | Kempen (2019) | Amsterdam | ingrid@gafanhao@kempen.com | +31 689 937 525 |
| Naresh Chouhan | Intron Health Research (2020) | London | naresh@intronhealthresearch.com | +44 7939 224 322 |
| Maneka Mirchandaney | Evercore (2018) | New York City | maneka.mirchandaney@evercoreisi.com | +1 646 740 1482 |
| Erik Hultgård | Carnegie (2019) | Stockholm | erik.hultgard@carnegie.com | +46 (0) 858-869 237 |
| Ludvig Svensson | Redeye (2008) | Stockholm | ludvig.svensson@redeye.se | +46 (0) 704-962 535 |
| Joseph Hedden | RX Securities (2016) | London | joseph@rxsecurities.com | +44 773 061 8803 |
| Lars Hatholt | Ökonomisk Ugebrev (2020) | Copenhagen | hatholt@outlook.com | +45 22 23 78 15 |

Contact our Investor Relations and Corporate Communications



Klaus Sindahl

Head of Investor Relations

Mobile: +46 (0) 709-298 269

Email: klaus.sindahl@hansabiopharma.com



Katja Margell

Head of Corporate Communications

Mobile: +46 (0) 768-198 326

Email: katja.margell@hansabiopharma.com

Visit our web site
www.hansabiopharma.com



Calendar

| | |
|-----------------|---|
| Nov 17, 2020 | Bryan Garnier Healthcare Conference, Paris (virtual) |
| Nov 18, 2020 | Jefferies Healthcare Conference, London (virtual) |
| Nov 25, 2020 | Ökonomisk Ugebrev Life Science Conference, Copenhagen |
| Nov 26, 2020 | Redeye Life Science Day, Stockholm (virtual) |
| Jan 11-14, 2021 | JP Morgan Week (virtual) |
| Jan 11-14, 2021 | H.C. Wainwright BioConnect Conference (virtual) |
| Jan 18, 2021 | SEB Nordic Healthcare seminar (virtual) |
| Feb 2, 2021 | Interim report Jan-Dec 2020 |
| Mar 9, 2021 | Carnegie Nordic Healthcare seminar (virtual) |
| April 22, 2021 | Interim report for Jan-Mar 2021 |
| May 5, 2021 | Kempen Life Sciences Conference (virtual) |
| July 15, 2021 | Interim report for Jan-Jun 2021 |
| Oct 21, 2021 | Interim report for Jan-Sep 2021 |

