

...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 our current expectations and beliefs regarding future events and are subject to significant uncertainties and risks since they relate to events and depend on circumstances that will occur in the future. Some of these forward-looking statements, by their nature, could have an impact on Hansa Biopharma's business, financial condition and results of operations [or that of its parent, affiliate, or subsidiary companies]. Terms such as "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 statements. There are a number of factors that could cause actual results and developments to differ materially from those projected, whether expressly or impliedly, in a forward-looking statement or affect the extent to which a particular projection is realized. Such factors may include, but are not limited to, changes in implementation of Hansa Biopharma's strategy and its ability to further grow; risks and uncertainties associated with the development and/or approval of Hansa Biopharma's product candidates; ongoing clinical trials and expected trial results; the ability to commercialize imlifidase if approved; changes in legal or regulatory frameworks, requirements, or standards; 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.

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Q2'20 Business Update

# Positive CHMP Opinion in EU; Gene therapy partnership with Sarepta; USD 121m raise in direct placement

### Highlights for the second quarter 2020

- Imlifidase in kidney transplantation
  - CHMP/EMA adopted a positive opinion for imlifidase in highly sensitized kidney patients in EU; Approval expected in Q3'20
  - New US study: The proposed study protocol for a randomized controlled study was submitted in June; Initiation expected in Q4'20
- Clinical Pipeline
  - Anti-GBM: Expect first data read-out from the Phase 2 trial in Q3'20
  - GBS/AMR: No patients enrolled during Q2 due to the impact from COVID-19. Both programs expected to be reinitiated in Q3'20
- Hansa continues to expand the organization
  - Achim Kaufhold, M.D. appointed as new Chief Medical Officer
  - Katja Margell appointed as new Head of Corporate Communications

#### Events after the reporting period

- New Gene therapy partnership with Sarepta Therapeutics
- SEK 1.1bn / USD 121m direct placing of new ordinary shares to fund R&D programs and commercial build-up



# EMA: The positive CHMP opinion serves as a validation of Hansa's proprietary enzyme technology

## EU: Imlifidase in kidney transplantation

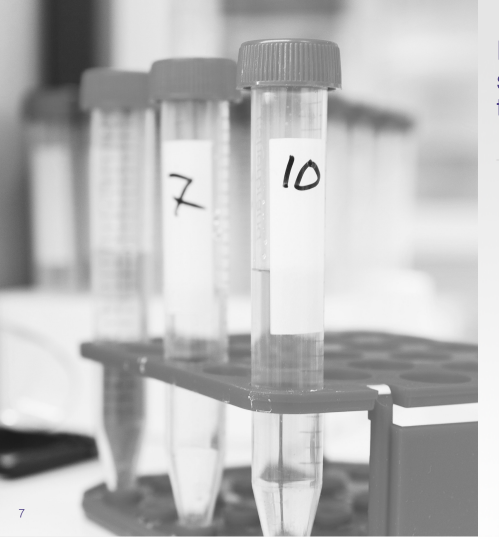
#### Europe

- CHMP/EMA recommends conditional approval of imlifidase for "the desensitization treatment of highly sensitized adult kidney transplant patients with a positive crossmatch against an available deceased donor".
- An approval by the EU Commission is expected in Q3 2020.
- The MAA for imlifidase in kidney transplant was accepted for review by EMA back in 2019 based on data from four completed phase 2 studies across Sweden, France and the US.
- Imlifidase was supported through EMA's PRIority MEdicines (PRIME) scheme, which was granted in May 2017.

#### Expected launch in Q4'20 in leading transplantations centers

- Our launch strategy involves targeting leading kidney transplantation centers with the potential to become early adopters and centers of reference
- A post approval study will run in parallel with the launch





# FDA: The proposed study protocol was submitted to the FDA in June; The new trial is expected to be initiated in Q4

## US: Imlifidase in kidney transplantation

U.S. (FDA)

- Hansa Biopharma submitted the proposed study protocol for the randomized controlled study with imlifidase in kidney transplant to the FDA on June 17, 2020.
- The new trial is expected to be initiated in Q4 this year. Potential reprioritization of activities by the FDA due to COVID-19 may however impact the timeline for the initiation of our new US trial
- The proposed trial would include 45 patients with a cPRA score of 99.9% or above. eGFR (kidney function) will be used as a surrogate endpoint to demonstrate a clinical benefit of imlifidase therapy vs. patients being waitlisted



# First read-out in the Anti-GBM study in Q3'20. Recruitment in AMR & GBS expected to be reinitiated in Q3'20

## Ongoing Phase 2 programs

# Enrollment status end Q2'2020



#### Anti-GBM (investigator-initiated study)

- 15/15 patients enrolled in anti-GBM across 5 European countries
- First data read-out expected in Q3 2020



#### Antibody Mediated Rejection

- 4/30 patients enrolled in AMR study.
- Recruitment is expected to be reinitiated in Q3 2020\*
- Enrollment is expected to be completed H1 2021



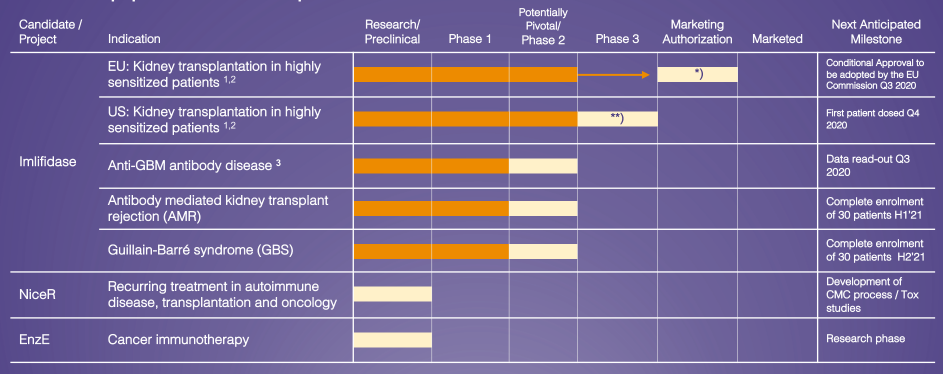
#### Guillain-Barré Syndrome

- 4/30 patients enrolled in GBS study
- Recruitment is expected to be reinitiated in Q3 2020\*
- Enrollment is expected to be completed in H2 2021
- Patients enrolled
- Patients left



<sup>\*</sup> Recruitment process was been impacted in Q2 following the COVID-19 virus pandemic causing a 3-6 months delay

# Broad pipeline in transplantation and auto-immune diseases



<sup>&</sup>lt;sup>1</sup> Results from the Phase 1 study have been published, Winstedt el al. (2015) PLOS ONE 10(7)



<sup>&</sup>lt;sup>2</sup> Lorant et al American Journal of Transplantation and 03+04 studies (Jordan et al New England Journal of Medicine)

<sup>&</sup>lt;sup>3</sup> Investigator-initiated study by Mårten Segelmark, Professor at the universities in Linköping and Lund

<sup>\*)</sup> EMA: Positive CHMP opinion received June 2020 for a conditional approval – Formal adoption by the EU Commission expected Q3 2020, while a post-approval study will commence in parallel with the launch

<sup>\*\*)</sup> FDA: Agreement with the FDA on a regulatory path forward in the US. New clinical study could support BLA submission by 2023. Safety review of an Investigational New Drug application (IND) expected in Q3 2020, while the study is expected to be initiated Q4 2020

# 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

- Preclinical/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 in milestones plus royalties and incremental imlifidase sales





#### Sarepta's key competences

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



#### Upfront payment

Hansa to receives a USD 10m upfront payment from Sarepta for accessing Hansa's unique IgG antibody-cleaving enzyme technology (imlifidase)

#### Milestones

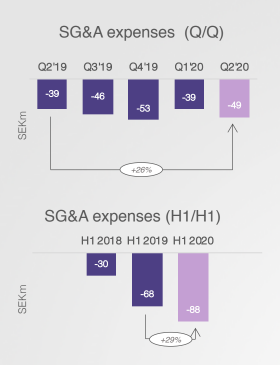
Hansa is eligible for a total of up to USD 397.5m in development, regulatory and sales milestone payments

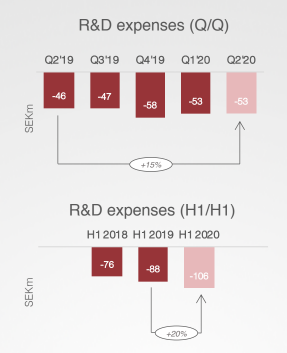
#### Royalties & Sales

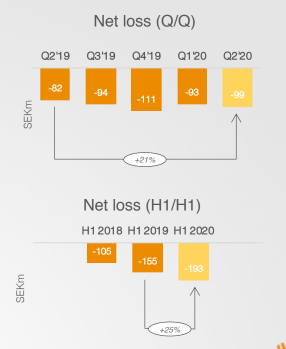
Hansa to receive high single-digit to mid-teens royalties on Sarepta's gene therapy sales enabled with imlifidase treatment in Nabs positive patients and book all sales of imlifidase



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

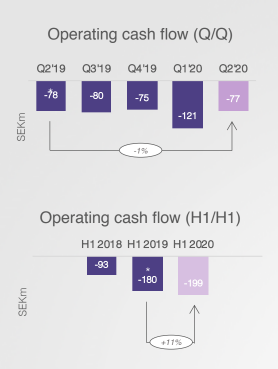


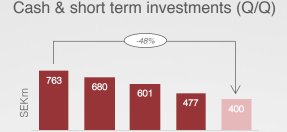






# Capital injection from the issue of 4.4m new shares (SEK 1.1bn) and upfront payment from Sarepta (SEK ~100m) will finance Hansa into 2023







Q4'19

Q1'20

Q2'20

Q2'19

Q3'19

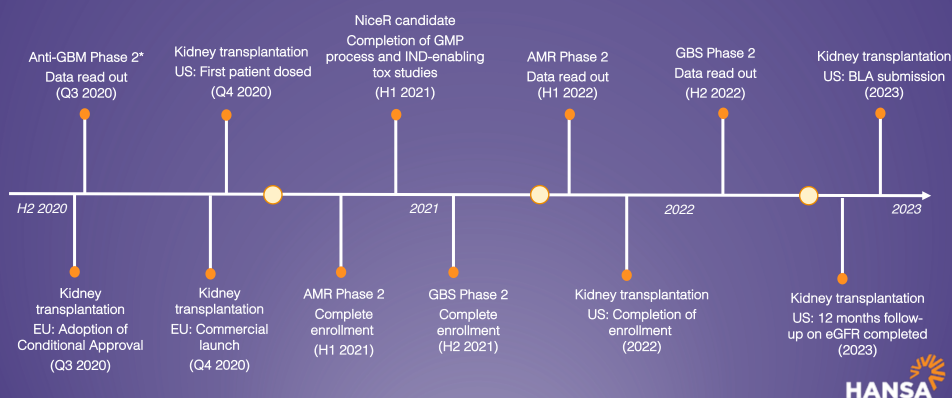


Cash position post Sarepta and capital raise



<sup>\*</sup> Excl. positive impact from sale of Genovis shares of SEK 89m in Q2'19

# Upcoming milestones





# Hansa Biopharma at a glance



#### Company background

- · Founded 2007 with HQ in Lund. Sweden
- Søren Tulstrup, CEO Ulf Wiinberg, Chairman
- ~80 employees (~2/3 in R&D) at June 30, 2020
- · Operations in Sweden, US & across Europe
- Market cap: SEK ~10bn (USD ~1bn) July 2020
- · Listed on Nasdag 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
- EU: Positive CHMP opinion received June 2020, EU approval expected Q3 2020
- US: Study protocol submitted June 2020, study expected to be initiated Q4 2020
- 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.
 H1'20\* SEK 400m (H1'19 SEK 763m)
 FY'19 SEK 601m
 Operating Profits/Loss
 H1'20\* SEK -193m (H1'19 SEK -156m)
 FY'19 SEK -360m
 FY'19 SEK -360m
 FY'19 SEK -355m

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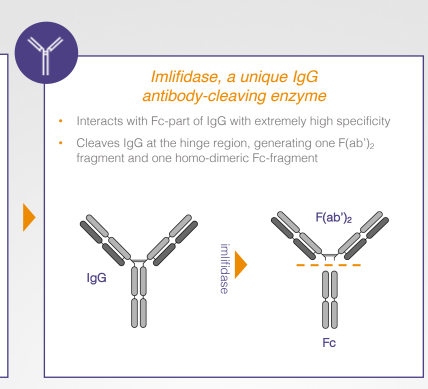


# Imlifidase – a novel approach to eliminate pathogenic IgG

# Origins from Streptococcus pyogenes

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

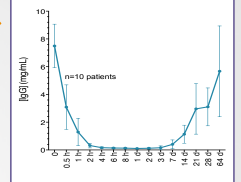






# Imlifidase inactivates IgG in 2 hours

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





# **Our Equity Story**



#### Targeting rare diseases with a high unmet medical need

 Imlifidase is a unique IgG antibody-cleaving enzyme with a rapid onset of action and high specificity for inactivation of IgG in patients with rare immunologic diseases.



#### Preparing for commercialization

- Preparing for potential European launch of imlifidase following potential conditional approval in 2020. Positive CHPM opinion received June 2020. Formal adoption by the EU Commission expected in Q3 2020.
- Imlifidase to be launched through Hansa's own medical and commercial organization, while we expected to pursue a partnership strategy outside core markets.
- In the US, a randomized controlled trial is planned to be initiated in Q4 this
  year, which could support a future BLA submission in the US by 2023. The
  study protocol was submitted to the FDA June 2020.
- Broad technology protection with patent coverage throughout 2035 in key markets and orphan drug designation in both the US/EU in our lead indications.



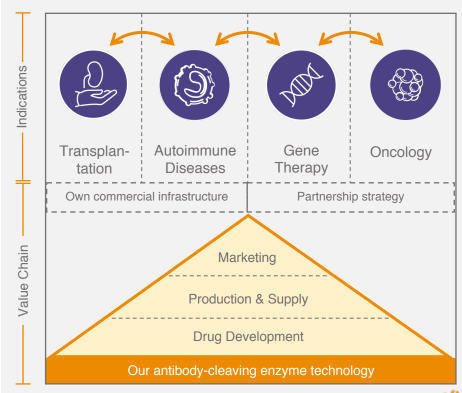
#### Evolution into a fully integrated biopharmaceutical company

 Controlling the full value chain from early discovery through commercialization to maximize the value creation and capture.



#### Leveraging our proprietary antibody cleaving enzyme technology

- Advancing our pipeline with three Phase 2 programs in transplantation and acute autoimmune diseases.
- New set of modified enzymes under development (NiceR program) for repeat dosing; potentially enabling treatment in relapsing diseases and oncology.
- Exploring potential combination therapies in oncology with IgG-modulating enzymes and gene therapy in patients with neutralizing antibodies through potential partnerships.

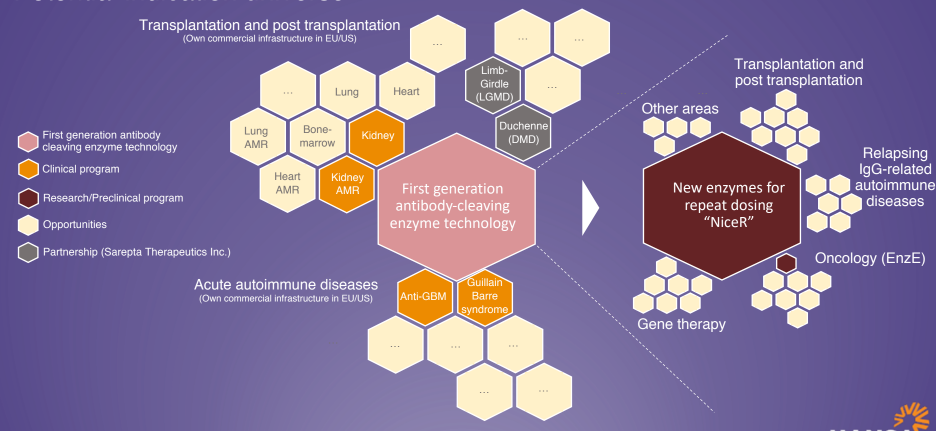




# Potential indication universe

#### Gene therapy pre-treatment

(partnership opportunity)



# Our unique enzyme technology platform offers significant potential for growth and expansion

## Our strategic priorities



Establish a commercial and medical infrastructure in Europe



Attain marketing authorization in Europe for imlifidase as a treatment for highly sensitized patients to enable kidney transplantation. Conduct a new randomized, controlled study in the US in the context of KAS to support a BLA filing by 2023



Investigate the potential of imlifidase in autoimmune indications and post transplantation

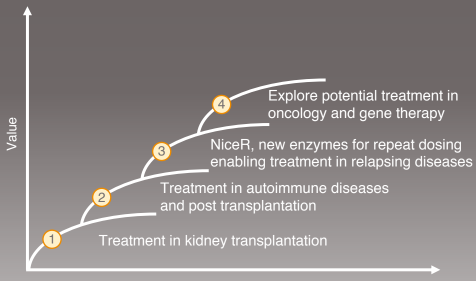


Advance a new set of immunomodulatory enzymes designed for repeat dosing in relapsing diseases (NiceR) into clinical development



Explore potential combination therapies with imlifidase in oncology and in gene therapy

## Our road map for growth and expansion







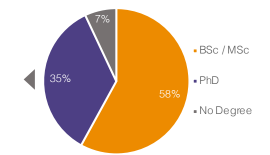
# The Hansa team has extensive experience from international life science industry and academia

Highly educated team with more than 1,100 "man years" in the life science industry and academia

More than 1/3 of the team holds a relevant life science PhD

#### PhD specializations include

- Applied Microbiology
- Biotechnology
- Cell and Molecular Biology
- Clinical Infection Medicine
- Engineering
- Experimental Clinical Chemistry
- Experimental Medicine
- Immune Technology
- Medical Microbiology
- Medical Science
- Physiological Chemistry

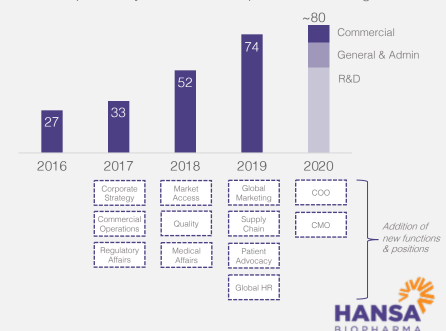


Vast experience from life science; +50% has worked in Big Pharma

Biotech Commercial-stage/Big pharma **MERCK** • Biolnvent AstraZeneca 2 novo nordisk **FERRING** AnaMar santaris

# We are building an organization in preparation to become a commercial-stage biopharma company

Staff has tripled in 5 years as new competences are being added

























# Experienced Board and Executive Committee with many years in the global healthcare industry

#### **Executive Committee**



Sören Tulstrup

President & CEO (2018)
+30 years in the Healthcare sector
Ex-CEO at Vifor Pharma
Ex-SVP at Shire Pharmaceuticals
Ex-CEO at Santaris Pharma



Christian Kjellman
SVP & CSO/COO (2008)
+20 years in the Healthcare sector
Ex-Head of Research at Cartela
Ex-Senior Scientist at Biolnvent,
MSc Chemical Biology, PhD in Turnour
Immunology from Lind University



Donato Spota

SVP & CFO (2019)
+20 years in the Healthcare sector
Ex-CFO Basilea Pharmaceutica
Senior Finance roles at Roche



Achim Kaufhold

CMO (2020)
+40 years in the Healthcare sector
Ex-CMO Basilea Pharmaceutica
Ex-CEO Affitech (merged with Pharmexa A/S)

Ex-CMO Chiron (acquired by Novartis)



Max Sakajja
VP, Corporate Strategy (2017)
Ex-M&A Director at SOBI
Ex-Global Product and Service
Development Manager at
Fruitotaine

Ex-independent life science industry management consultant



Henk D. van Troostwijk SVP & CCO (2016) +20 years in the Healthcare sector Ex-GM at Raptor Pharmaceuticals

Ex-BU Director at Genzyme



Anne Säfström Lanner VP, Global HR (2019) Ex-Head of HR European Spallation Source



Ulf Wiinberg



Chairman (2016) +30 years in the Healthcare sector Ex-CEO at Lundbeck (2008-14) Ex-President at Wyeth of the global consumer heath care and European Pharma business



Birgit Stattin Norinder
Board Member (2012)
Ex-CEO and Chairman at Prolifix Ltd.
Ex-SVP, Pharmacia & Upjohn
Member of Hansa Biopharma Scientific
Committee and Remuneration Committee



Anders Gersel Pedersen Board Member (2018) +30 years in the Healthcare sector Ex-EVP R&D H.Lundbeck Chairman of Hansa Biopharma's Scientific Committee



Eva Nilsagård

Board Member (2019)
interim CFO at OptiGroup AB
CEO of Nilsagård Consulting AB
Ex-CFO of Vitrolife and Plasta
Chairman of Hansa Biopharma's
Audit Committee



Mats Blom
Board Member (2019)
CFO of NorthSea Therapeutics
Ex-CFO Zealand Pharma
Member of Hansa Blopharma's Audit
Committee



Andreas Eggert

Board Member ( 2018)

Ex- SVP at H. Lundbeck A/S

Ex-VP Wyeth/Pfizer in the U.S.

Member of Hansa Biophama's Audit

Committee and Renumeration Committee



# From technology development to expected commercialisation in 13 years



Hansa Medical founded

IdeS (imlifidase) discovered and patented by Prof. Lars Björk, M.D. Lund University



Partnship with Axis-Shield for HBP-test



Imlifidase first-in-man study



Start imlifidase Phase 2 at Cedars Sinai and UUH



Imlifidase NEJMpublication: Anti-GBM initiated



Imlifidase MAA submitted Partnship with to EMA: Sarepta for gene AMR & GBS Phase 2 therapy initiated





Project IdeS (imlifidase) initiated



Imlifidase GMP process development and toxicology studies initiated



Imlifidase 1st Phase 2



Initiation of imlifidase HighdeS study



Imlifidase: 46 transplants enabled



Imlifidase: Positive CHMP opinion June 2020



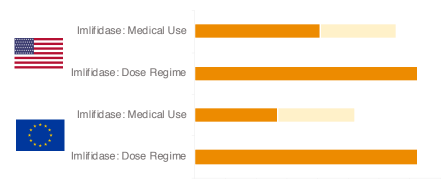


# Strong technology protection through patents and orphan drug designation

### Patent coverage out to 2035 in key markets

- Hansa Biopharma's portfolio consist of 11 separate patent families incl. 7 patent families in relations to the use of imlifidase (granted/pending)
- · Patents cover use of isolated imlifidase in:





### Orphan drug designation

- Orphan drug designation is granted to drugs intended for rare diseases (affecting max 5 patients in 10,000 persons in EU or affecting less than 200,000 patients in the US.
- Designation provides development and commercial incentives incl. 10 years market exclusivity in EU and 7 years in the US

#### **EMA**

#### Orphan drug designation

- Imlifidase for the prevention of graft rejection following solid organ transplantation (2017)
- Imlifidase for the treatment of the rare and acute disease anti-GBM (2018)

#### FDA

#### Orphan drug designation

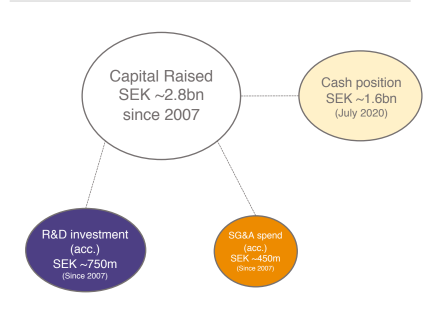
- Imlifidase for the prevention of antibody-mediated organ rejection in solid organ transplantation (2015)
- Imlifidase for the treatment of Guillian-Barré Syndrome (2018)
- Imlifidase for the treatment of the rare and acute disease anti-GBM (2018)



2021 2023 2025 2027 2029 2031 2033 2035 2037

# 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



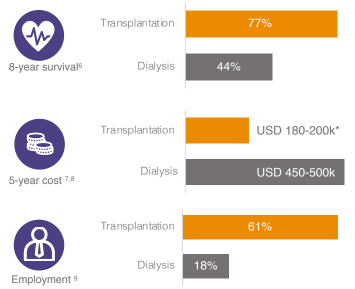


# Kidney transplantation saves lives, reduce costs and increase quality of life incl. gains for the society

## Several complications and risks with dialysis

- Undergoing dialysis treatment is associated with many complications and side effects incl. cardiovascular diseases<sup>1</sup>. In the long term, patients may also eventually lose access to dialysis as a result of failed ports, bad veins, and other factors<sup>2</sup>
- In general, patients on the kidney transplant waiting list and who are on dialysis have a lower quality of life than non-dialysed patients or patients who have been transplanted<sup>3</sup>
- First study in Europe on labor market outcomes demonstrates societal gains of enabling transplantation with three times as many transplant patients employed compared to dialysis patients.
- Lastly, extended dialysis is also a high-risk factor for removal from the transplant wait list<sup>6</sup>

### Transplantation leads to better outcomes





<sup>&</sup>lt;sup>1</sup> Cozzolino et al., 2018

<sup>&</sup>lt;sup>2</sup> Sinnakirouchenan and Holley, 2011 Shenoy, 2017

<sup>&</sup>lt;sup>3</sup>Wyld et al., 2012

<sup>&</sup>lt;sup>4</sup> Jarl et al. Transplantation, 2018, 102:1375-1381

<sup>&</sup>lt;sup>5</sup> NHS blood and transplant, 2018.

<sup>&</sup>lt;sup>6</sup> Orandi et al. N Engl J Med 2016;374:940-50

<sup>&</sup>lt;sup>7</sup> www.usrds.org

<sup>&</sup>lt;sup>8</sup> Shehata et al, Transfus Med Rev 201, 24 Suppl 1: S7-S27

<sup>&</sup>lt;sup>9</sup> Jarl et al. Transplantation, 2018, 102:1375-1381

# Approximately 10-15% of patients on wait list are highly sensitized

## Highly sensitized patients are difficult to match

Causes of sensitization include





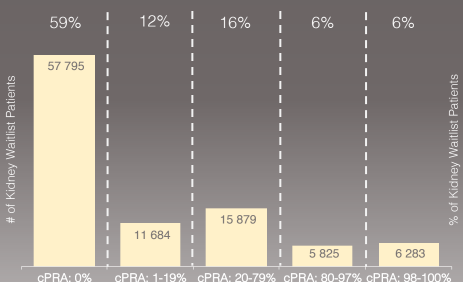


Pregnancy

Blood transfusion

Previous transplantations

- Calculated Panel Reactive Antibodies (cPRA) is a measure for **HLA-sensitization**
- Inability to match or effectively desensitize patients remains a barrier for transplantation in highly sensitized patients
- Allocation Systems such as KAS and Eurotransplant rely on cPRA score to characterize patients for transplant





# Imlifidase may enable transplantation in highly sensitized kidney patients

### Creating equity for highly sensitized patients

- Allocation systems increase transplantation rates, however the rates for highly sensitized patients are still very low compared with average or non-sensitized patients
- If approved, imlifidase may potentially:
  - Complement allocation systems (e.g. KAS, Euro-transplant) to reduce time to transplant in highly sensitized patients
  - Reduce the need for antibody matching and give sensitized patients access to a larger pool of organs
  - Reduce the risk for co-morbidities and mortality associated with dialysis and waiting time
  - · Increase transplant rates in highly sensitized patients
  - Help reduce the number of discarded kidneys
     (1,000 donated kidneys are discarded in the U.S. alone every year<sup>3</sup>)

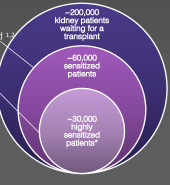


Delilah, a 23 years old highly sensitized kidney transplant patient from California

#### U.S. + EU Kidney Transplant Waitlist Breakdown

>30% of waitlist patients are sensitized

- 15% moderately sensitized
- 15% highly sensitized1,2 \*



~40,000 transplants done annually in the US and EU.



<sup>&</sup>lt;sup>1</sup> Jordan et al. British Medical Bulletin, 2015, 114:113–125

<sup>&</sup>lt;sup>2</sup> Orandi et al. N Engl J Med 2016;374:940-50

<sup>&</sup>lt;sup>3</sup> Organ Procurement and Transplantation Network (OPTN)

<sup>&</sup>lt;sup>4</sup> Jordan et al. British Medical Bulletin, 2015, 114:113-125

# Focused launch strategy targeting leading kidney transplantation centers to ensure positive experience

### Potential EU launch following conditional approval

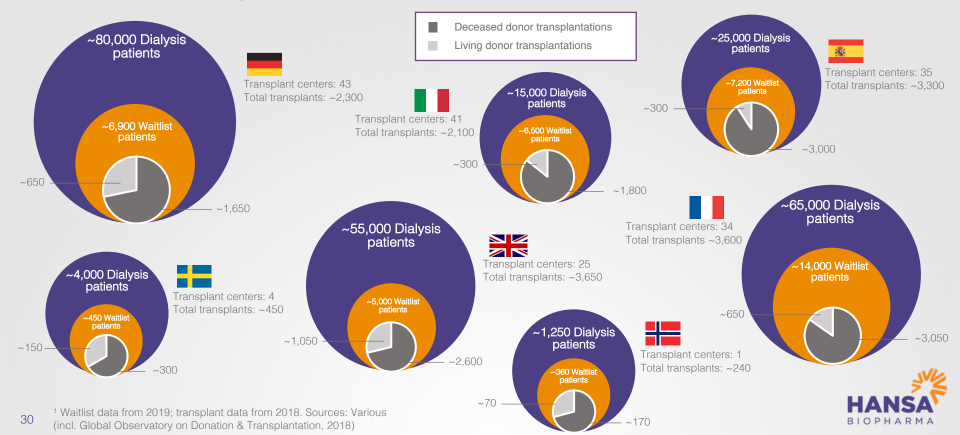
- A sequenced and focused strategy to launch imlifidase
  - Well defined and concentrated target audience
  - Center-focused launch strategy targeting leading clinics with the potential to become early adopters
  - Key to secure early positive experience in right patients; sales ramp-up as leading centers and clinicians gain experience
- Building awareness and Key Opinion Leader advocacy through Medical Science Liaisons (MSLs) in key European markets
- Post-approval study to be initiated following potential marketing authorization - an opportunity to generate relevant experience and broaden out the experience with imlifidase

### EU launch will focus on leading transplantation centers



# European transplantation landscape

Approximately 16,000 kidney transplants in EU5 plus Sweden and Norway<sup>1</sup> with 70-80% performed at leading transplantation centres in each country

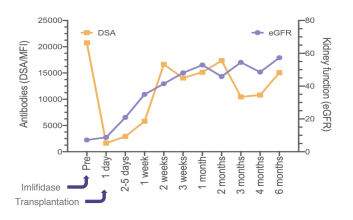




# Imlifidase has enabled kidney transplantation in 46 highly sensitized patients

### Pooled analysis from four Phase 2 trials

- Analysis included 46 patients
  - 50% had a cPRA of 100% (Average 99%)
  - 85% were crossmatch positive
  - 70% were retransplanted
- Donor Specific Antibody (DSA) levels rapidly decreased and all crossmatches were converted to negative, thus enabling transplantation in all patients
- At study completion, all patients alive and graft survival at 94% six months post transplantation



## Study design of our four Phase 2 trials

Study 02 Phase 2

8 patients Subjects



Design

Single-center, single-arm, open-label

Main objective

Efficacy defined as Imlifidase dosing scheme resulting in HLA antibody levels acceptable for transplantation, within 24 hours



Subjects

10 patients



Design

Single-center, single-arm, open-label, no prior desensitization

Main objective

Safety in the transplantation setting and efficacy defined as

HLA antibody levels acceptable for transplantation



Subjects

17 patients



Design

Investigator initiated, single-center, single-arm, open-label. All patients had prior desensitization with IVIG and/or PLEX

Main objective

Safety in combination with Cedars Sinai's "standard protocol"

for desensitization of highly sensitized patient



Subjects

18 patients



Design

Multicenter, multinational, single-arm, open-label

Main objective

Efficacy in creating a negative crossmatch test

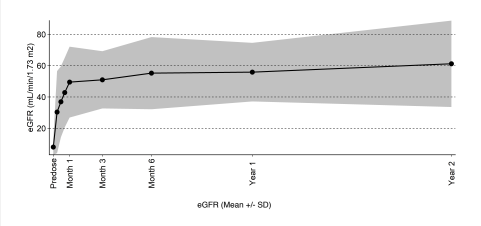


# Two year follow-up data show graft survival of 90% and well functioning kidneys in 92% of these patients

### AMR frequency in line with less sensitized patients

- Two-year follow-up data post imlifidase treatment and transplantation show 90% graft survival for 31 patients
- Of the patients with data at two years, 92% had a well functioning kidney with median eGFR of 61ml/min/1.73 m<sup>2</sup>
- 33% of the patients experienced active antibody mediated rejections (AMR) within the first six months, which compares with 25-60% of patients in the literature for this group of highly sensitized patients<sup>1</sup>
- Only one patient experienced an AMR episode later than six months after transplantation
- The analysis concludes that the AMR frequency was comparable with other studies with less sensitized patients in crossmatch positive patients

### Median eGFR at 61ml/min/1.73 m² after year 2







#### CLINICALTRIALS.GOV ID

NCT01802697 (2013/2014)

#### **SUBJECTS**

29 (20 active plus 9 placebo) health subjects (Sweden)

#### DOSES/FOLLOW UP TIME

The starting dose was 0.01 mg/kg BW and the highest dose group received 0.24 mg/kg BW

#### MAIN OBJECTTIVES

 The objectives were to assess safety, efficacy in IgG cleavage, pharmacokinetics and immunogenicity of imilifidase following intravenous administrati

#### STUDY DESIGN

 Randomized placebo-controlled dose-escalation study with 29 (2 active plus 9 placebo) healthy subjects

#### **STATUS**

#### Camalakad

 The 01 study showed complete removal of IgG and that Imlifidas was considered safe to use

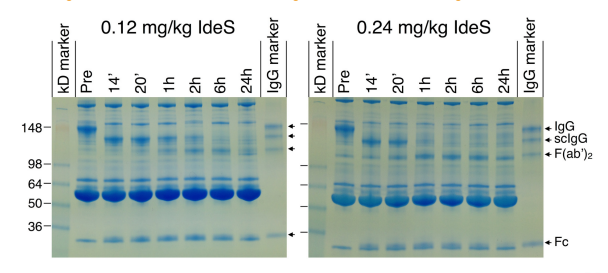
# The 01 study showed complete removal of IgG and a good tolerability profile

Efficacy

✓ Rapid degradation of IgG in serum in subjects dosed with 0.12 and 0.24 mg/kg imlifidase. Imlifidase had full effect within 6 hours. The entire IgG pool was converted into F(ab')<sub>2</sub> and Fc-fragments. Maximal effect was accomplished 2-6 hours after dosing.

Safety

✓ Newly synthesized intact IgG was clearly detectable in all subjects after 1-2 weeks after dosing. After 3 weeks the level of intact IgG constituted the main IgG fraction in serum





#### CLINICALTRIALS.GOV ID

NCT02224820

#### **SUBJECTS**

8 Patients with chronic kidney disease (Sweden)

#### DOSES/FOLLOW UP TIME

0.12 & 0.25 mg/kg BW given once or twice within 48 hours

#### MAIN OBJECTTIVES

- Efficacy defined as Imlifidase dosing scheme resulting in HLA antibody levels acceptable for transplantation, within 24 hours from dosing
- Safety

#### STUDY DESIGN

- Single-center, Single arm with ascending doses, open label.
- Transplantation not part of protoco

#### **STATUS**

#### Completed

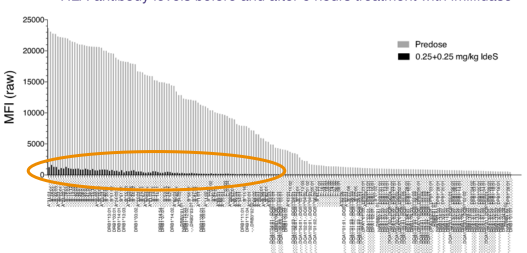
- Primary efficacy endpoint reache
- · Sale allu well luleralet

#### 35

# The 02 study showed that 1-2 doses of imlifidase at 0.25 mg/kg BW resulted in HLA antibody levels acceptable for transplantation<sup>1</sup>

- ✓ Imlifidase is well tolerated in patients with chronic kidney disease
- Efficacy results strongly support further development in the patient population
- ✓ The first HLA-incompatible transplantation ever after desensitization with imlifidase was performed in one of these patients (2014)

#### HLA-antibody levels before and after 6 hours treatment with imlifidase



<sup>&</sup>lt;sup>1</sup> Lorant et al (2018) American Journal of Transplantation (2018)



# Study 03 Phase 2

#### CLINICALTRIALS.GOV ID

NCT0247555

#### **SUBJECTS**

10 Patients (Sweden)

#### DOSES/FOLLOW UP TIME

0.25 and 0.50 mg/kg during 180 days

#### MAIN OBJECTTIVES

- · Safety in the transplantation setting
- Efficacy defined as HLA antibody levels acceptable for transplantation

#### STUDY DESIGN

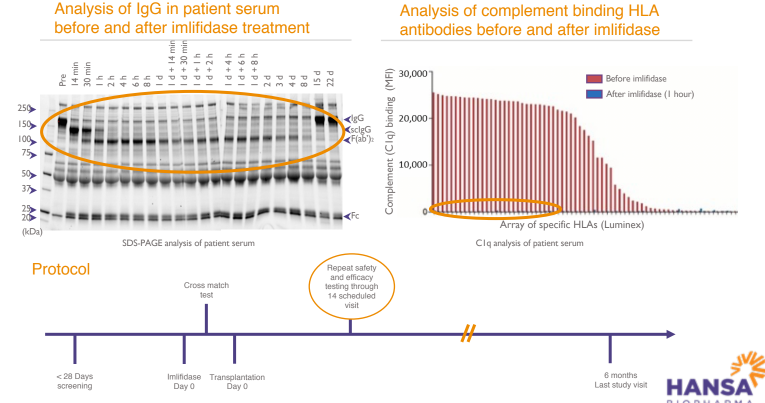
- Single-center, single-arm, openlabel, no prior desensitization
- Similar design as 13-HMedIdeS-0 but transplantation part of protoco
- · In deceased and living donors

#### **STATUS**

#### Completed

Proofed safety and efficacy with HLA antibodies at acceptable levels; enabling transplantation in all patients

# The 03 study proved safety and efficacy with HLA antibodies at acceptable levels; enabling transplantation in all patients





#### CLINICALTRIALS.GOV ID

NCT024226684

#### **SUBJECTS**

17 Patients (US)

#### DOSES/FOLLOW UP TIME

0.24 mg/kg 180 days

#### MAIN OBJECTTIVES

- Safety in combination with Cedars Sinai's "standard protocol" for desensitization of highly sensitized patients
- · Efficacy in preventing AMF

#### STUDY DESIGN

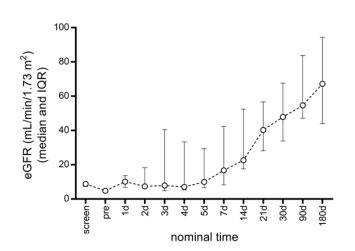
- Investigator initiated study
- · Investigator sponsored IND
- Imilifidase to desensitize patient previously treated with rituximal and IMa
- Deceased donors onl

#### **STATUS**

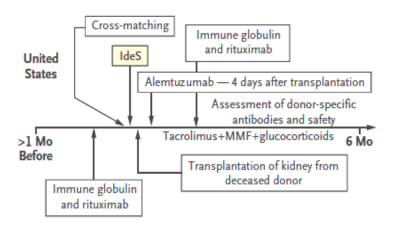
Completed

## The 04 study proved safety and efficacy with Cedar Sinai's standard protocol (rituximab and IVIg)

## Graft function (eGFR) post six months



## Cedar's desensitization protocol in combination with imlifidase







#### CLINICALTRIALS.GOV ID

NICTO2790437

#### **SUBJECTS**

18 Patients (US+Sweden+France) 19 safety set, 18 efficacy set

#### DOSES/FOLLOW UP TIME

0.25 mg/kg 180 days

#### MAIN OBJECTTIVES

Efficacy in creating a negative crossmatch test

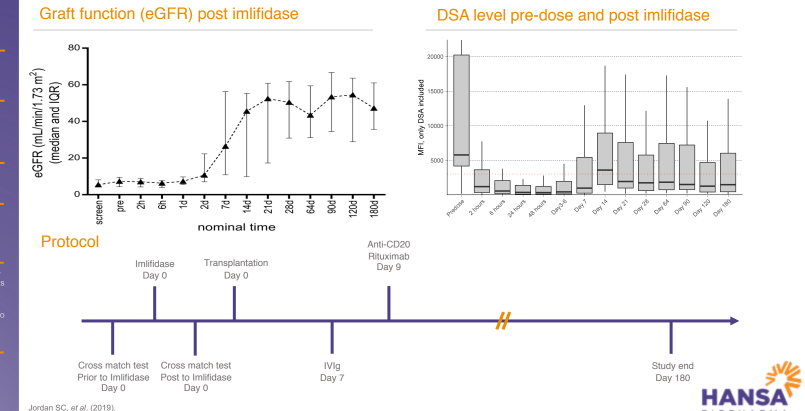
#### STUDY DESGIN

 Multicenter, multinational, singlearm, open-label Included patients who may have had prior unsuccessful desensitization or patients in whom it was unlikely to be effective

#### STATUS

Complete

# The 06 study showed proved safety and efficacy in making highly sensitized patients eligible for kidney transplantation



## Overview of all completed studies with imlifidase in transplantation

STUDY	SUBJECTS/ COUNTRY	STUDY DESIGN	PRIMARY ENDPOINT	SECONDARY ENDPOINTS	STATUS/ PUBLICATION
Study 01 Phase 1	29 subjects	<ul> <li>Randomized placebo-controlled dose- escalation study with 29 (20 active plus 9 • placebo) healthy subjects</li> </ul>	Safety and tolerability	Efficacy in IgG cleavage, the pharmacokinetics (PK) and immunogenicity of imlifidase	Complete PLOS ONE (2015) <sup>1</sup>
Study 02 Phase 2	8 subjects	Single-center, single-arm, open-label	Dosing resulting in HLA-antibody reduction (MFI<1100)	Efficacy: HLA antibody reduction acceptable for transplantation (MFI <1100 as measured in SAB assay)	Complete Lorant et al (2018) American Journal of Transplantation <sup>2</sup>
Study 03 Phase 2	10 subjects	<ul> <li>Single-center, single-arm, open-label</li> <li>No prior desensitization</li> </ul>	Safety: AEs, clinical laboratory tests, vital signs, ECGs	Efficacy: HLA antibody reduction acceptable for transplantation (MFI <1100 as measured in SAB assay)	Complete The New England Journal of Medicine (2017) <sup>3</sup>
Study 04 Phase 2	17 subjects	<ul> <li>Investigator initiated study, Single-center, single-arm, open-label</li> <li>All patients had prior desensitization with IVIG and/or plasmapheresis</li> </ul>	Assessment of efficacy in eliminating DSAs in DSA and flow cytometry positive, highly sensitized patients Assessment of safety Assessment of efficacy/kidney function	<ul> <li>Serum creatinine (0-6 months)</li> <li>Proteinuria (0-6 months)</li> <li>DSA at multiple timepoints posttransplant (day 0, D30, D90, D180)</li> </ul>	Complete The New England Journal of Medicine (2017) <sup>3</sup>
Study 06 "Highdes" Phase 2	18 subjects	Multicenter, multinational, single-arm, open-label Included pts who may have had prior unsuccessful desensitization or pts in whom it was unlikely to be effective	Crossmatch conversion in DSA+ patients who have a positive XM test to their available LD or DD	<ul> <li>DSA reduction at multiple timepoints (2, 6, 24, 48 h after imlifidase)</li> <li>Time to create negative CDC XM test and/or flow cytometry (FACS) XM test</li> <li>Safety</li> </ul>	Complete Annals of Surgery (Lonze et al, only New York patients) Montgomery et al ATC abstract (2019) <sup>4</sup>
Long-term follow-up	Up to 46 subjects	<ul> <li>A prospective, observational long-term follow-up study of patients treated with</li> </ul>	Long-term graft survival in patients who have undergone kidney transplantation	<ul> <li>Patient survival, kidney function, comorbidity, treatments and QoL</li> <li>Safety</li> </ul>	Ongoing

after imlifidase administration

Immunogenicity

Jointain et al., "Selfety And Efficacy Of Imitifidase In Highly Sensitized Kidney Transplant Patients: Results From A Phase 2 Study" ATC Abstract, 2019

imlifidase prior to kidney transplantation

study

<sup>1</sup> Winstedt el al., "Complete Removal of Extracellular IgG Antibodies in a Randomized Dose Escalation Phase I Study with the Bacterial Enzyme IdeS – A Novel Therapeutic Opportunity", PLOS ONE 2015, 10(7)

<sup>&</sup>lt;sup>2</sup> Lorant et al., "Safety, immunogenicity, pharmacokinetics and efficacy of degradation of anti-HLA antibodies by IdeS (imlifidase) in chronic kidney disease patients" Am J Transplant. 2018 Nov;18(11):2752-2762 Jordan et al., "IgG Endopeptidase in Highly Sensitized Patients Undergoing Transplantation", N Engl J Med 2017;377:442-53.

# Medical Advisory Board in kidney transplantation



### Professor Stanley Jordan

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## Professor Kathryn Wood

Ph.D. Fellow of the Academy of Medical Sciences, Professor of Immunology in the Nuffield Department of Surgical Sciences, University of Oxford, England, runs the Transplantation Research Immunology Group

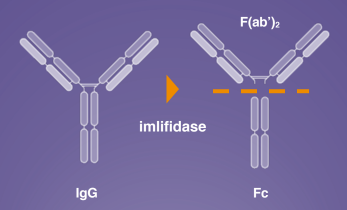




# Imlifidase, a novel approach to effectively eliminate pathogenic IgIG

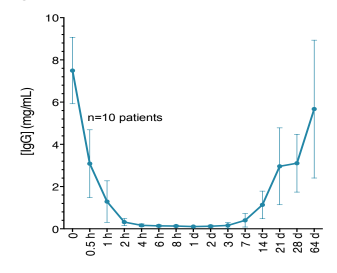
## Proven mechanism of action

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



## Inactivation of IgG in human serum

- Rapid onset of action that takes down IgG below detectable level in 2 hours post 15 min infusion
- IgG antibody-free window for approximately one week

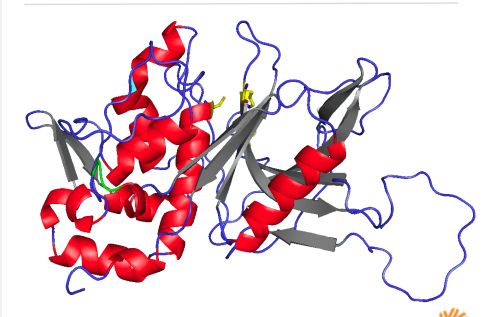


## Our IgG antibody-cleaving enzyme

## Origins from Streptococcus pyogenes

- Cysteine protease derived from an Immunoglobulin G (IgG)degrading enzyme of Streptococcus pyogenes
- Contains only one cysteine no disulfide bridges
- Monomeric protein with a molecular mass of 35 Kilo Dalton
- Isoelectric point of 6.1
- The coding gene for imlifidase is cloned and expressed in Escherichia coli

## Imlifidase consists of 311 amino acids





# Imlifidase is a lyophilized product formulation with a shelf life of 12 months at 2-8° Celsius storage

### Imlifidase will be infused in 15 minutes

- The product for commercial supply will be a lyophilized (cold chain) powder concentrate (11 mg solution) for infusion currently with a claimed shelf life of 12 months at 2-8°C storage. Ongoing stability studies indicate a shelf life of at least 24 months.
- Each vial is filled with 1.2 mL of a 10 mg/mL solution before freeze drying (=12 mg). Extractable volume after reconstitution with 1.2 mL sterile water is 1.1 mL of 10 mg/mL solution resulting in 11 mg product
- The protein concentration,10 mg/mL, has desirable characteristics with respect to not form aggregates
- Continuous stability programs ongoing to study changes in protein characteristics and performance.
- Imlifidase dose is clinically set to 0.25 mg/kg bodyweight (11 mg / 0.25 mg/kg = 44 kg (BW) / vial content) 2R vial size is suitable for the content





## Supply Chain for imlifidase in kidney transplantation



## Baxter



Drug product manufacturer (upscaling)



# Manufacturing will be done in close collaboration with highly experienced European based third party CMOs

Drug substance production process (API)

#### **Biotechpharma**



#### Fermentation/ harvesting

- Working Cell Bank
- Pre-Cultivation
- Main Cultivation
- Cell Harvest

#### Protein purification

- · Cell Disruption
- · Protein Release
- Ceramic Hydroxy Apatite Chromatography

Protein purification cont.

Chromatography

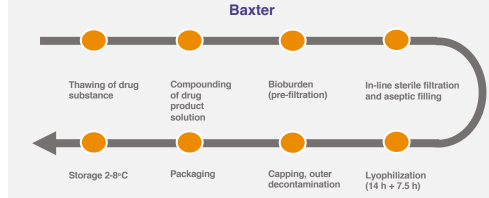
Ion Exchange

- Hydrophobic Interaction Chromatography
- · Ultrafiltration/ Diafiltration

#### Filling

 Formulation, filtration, filling and storage (-80°C)

## Drug product production process (upscaling)





#### <u>Facts</u>

- Based in Vilnius, Lithuania
- · Start-up Year: 2004
- Capacity: 300 L fermentor (1000 L fermentor in 2020)
- Certifications: GMP compliance, Manufacturing authorization license
- Inspections: National regulatory agency (EU), EU/US customer inspections,
   FDA mock inspection



#### Facts

1000年7月

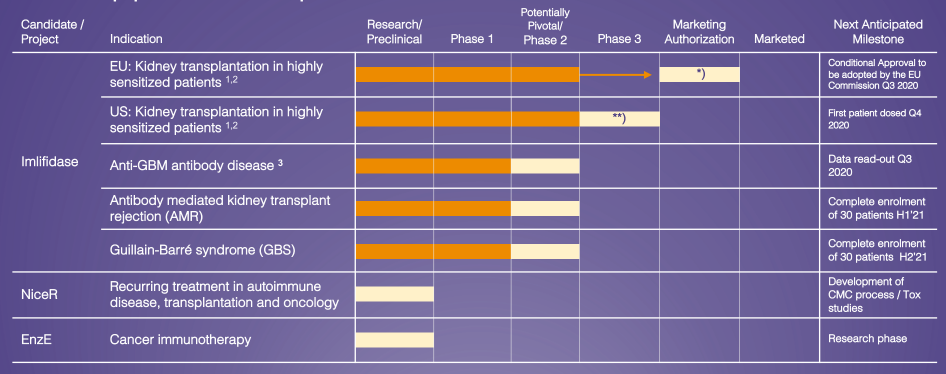
- Based in Halle/Westfalen Germany
- Start-up Year: 2001 (contract manufacturing)
- Capacity: 6-35 L drug product solution per batch (5,000-30,000 vials)
- · Certifications: GMP compliance, Manufacturing authorization license
- Inspections: National regulatory agency (EU), FDA, EU/US customer inspections

HANSA

Clinical development programs



## Broad pipeline in transplantation and auto-immune diseases



<sup>&</sup>lt;sup>1</sup> Results from the Phase 1 study have been published, Winstedt el al. (2015) PLOS ONE 10(7)



<sup>&</sup>lt;sup>2</sup>Lorant et al American Journal of Transplantation and 03+04 studies (Jordan et al New England Journal of Medicine)

<sup>&</sup>lt;sup>3</sup> Investigator-initiated study by Mårten Segelmark, Professor at the universities in Linköping and Lund

<sup>\*)</sup> EMA: Positive CHMP opinion received June 2020 for a conditional approval – Formal adoption by the EU Commission expected Q3 2020, while a post-approval study will commence in parallel with the launch

<sup>\*\*)</sup> FDA: Agreement with the FDA on a regulatory path forward in the US. New clinical study could support BLA submission by 2023. Safety review of an Investigational New Drug application (IND) expected in Q3 2020, while the study is expected to be initiated Q4 2020

# First read-out in the Anti-GBM study in Q3'20. Recruitment in AMR & GBS expected to be reinitiated in Q3'20

## Ongoing Phase 2 programs

## Enrollment status end Q2'2020



#### Anti-GBM (investigator-initiated study)

- 15/15 patients enrolled in anti-GBM across 5 European countries
- First data read-out expected in Q3 2020



#### Antibody Mediated Rejection

- 4/30 patients enrolled in AMR study.
- Recruitment is expected to be reinitiated in Q3 2020\*
- Enrollment is expected to be completed H1 2021



#### Guillain-Barré Syndrome

- 4/30 patients enrolled in GBS study
- Recruitment is expected to be reinitiated in Q3 2020\*
- Enrollment is expected to be completed in H2 2021
- Patients enrolled
- Patients left



<sup>\*</sup> Recruitment process was been impacted in Q2 following the COVID-19 virus pandemic causing a 3-6 months delay

# Anti-GBM, a rare acute autoimmune disease affecting kidneys and lungs; Enrollment completed in Q1 2020

## 2/3 of Anti-GBM patients lose kidney function<sup>2</sup>

- Indication: Antibodies are directed against an antigen intrinsic to the glomerular basement membrane (GBM) causing acute injury of kidney and/or lung
- Anti-GBM affects 1.6 in a million people annually with majority of patients losing their kidney function<sup>1,2</sup>, requiring chronic dialysis and kidney transplantation.
- The study is an open label investigator-initiated Phase 2 with Professor Mårten Segelmark at Linköping- and Lund University Hospital as the sponsor and principal investigator
- The study is designed to evaluate the safety and tolerability of imlifidase in patients with severe anti-GBM disease on top of standard care consisting of plasmapheresis, steroids and cyclophosphamide.
- 15/15 patients enrolled in anti-GBM across 5 European countries. First data read-out expected in Q3 2020.
- Our Anti-GBM program obtained Orphan Drug designation from both FDA and European Commission (2018)



## Anti-GBM Phase 2

#### CLINICALTRIALS.GOV ID

NCT03157037 (Since March 2017)

#### **SUBJECTS**

15 patients targeted. Patients will be monitored for six months Recruitment at 15 clinics

#### DOSES/FOLLOW UP TIME

Dosage 0.25mg/kg 180 days follow up

#### MAIN OBJECTTIVES

 Primary objective is to evaluate the safety and tolerability of imlifidase on background of standard of care and assess efficacy based on rena function at six months after treatment

#### STUDY DESIGN

- Open label, multicenter, single arn Phase 2 study with adverse renal prognosis
- Investigator initiated study

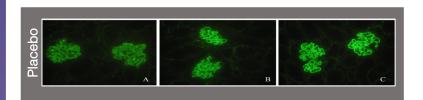
#### STATUS

Ongoin

#### 51

## Favourable pre-clinical studies show that imlifidase degrades IgG bound to the GBM in vivo; preventing renal damage in animals

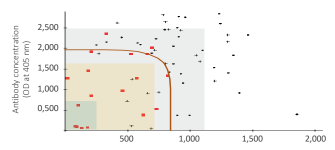
### Mouse anti-rabbit IgG (Fc specific)





### Anti-GBM creatinine and antibody concentration

 Both creatinine and levels of antibodies predict outcome and we expect that imlifidase can treat the disease by degrading IgG bound to the GBM



Serum creatinine concentration at diagnosis (µmol/l)

#### Inclusion criteria

Inclusion: Toxic anti-GBM antibodies level as considered by the investigator. eGFR < 15 ml/min/1.73 m2 or if the patient is non-responsive to standard treatment, and has lost >15 ml/min/1.73 m2 after start of treatment

**Exclusion:** Anuria for more than 2 days (less than 200 ml during last 48 hours); Dialysis dependency for more than 5 days



Yang et al. Favorable pre-clinical studies: "Imlifidase degrades IgG bound to the GBM in vivo, thereby preventing renal damage in this animal model. Nephrology Dialysis Transplantation. 2010;25(8): 2479-86.

# Long term graft survival is challenged by antibody mediated rejection post transplantation

## There is no approved treatment for AMR

- Active antibody mediated rejection after transplantation occurs in 10-15% of kidney transplants<sup>1</sup> or ~ 3,200<sup>2,3</sup> new patients annually<sup>4</sup> and is a significant challenge to long term graft survival
- Today's standard of care include plasma exchange, and treatment with steroid and IVIg. AMR patients not treated successfully risk graft failure, dialysis and return to the waitlist
- The AMR Phase 2 study is a randomized, open-label, multicenter, active control study designed to evaluate the safety and efficacy of imlifidase in eliminating donor specific antibodies (DSAs) in the treatment of active episodes of acute AMR in kidney transplant patients.
- 4/30 patient treated with imlifidase in AMR. 6/8 sites have been initiated to recruit patients in the US, Europe and Australia.
- Enrollment is expected to be completed H1 2021



<sup>&</sup>lt;sup>1</sup> Puttarajappa et al., Journal of Transplantation, 2012, Article ID 193724.

<sup>&</sup>lt;sup>2</sup> Jordan et al., British Medical Bulletin, 2015, 114:113-125.

<sup>&</sup>lt;sup>3</sup> http://www.irodat.org.

<sup>&</sup>lt;sup>4</sup> Seven major markets – US, Germany, UK, France, Spain, Italy, and Japan



## New AMR Phase 2 study initiated to test imlifidase ability to reduce the amount of DSA in AMR patients post transplantation

#### CLINICALTRIALS.GOV ID

NCT03897205 (2019

#### SUBJECTS

30 patients targeted (20 patients will be treated with imlifidase and 10 with Plasma exchange). Recruitment from 8 sites in the U.S., EU and Australia.

#### DOSES/FOLLOW UP TIME

1 dose of imlifidase (0.25 mg/kg) o 5-10 sessions of plasma exchange

#### MAIN OBJECTTIVES

- Imlifidase ability to reduce the amount of DSA in comparison with plasma exchange in patients who have an active AMR post transplantation
- Ensure safety for patients

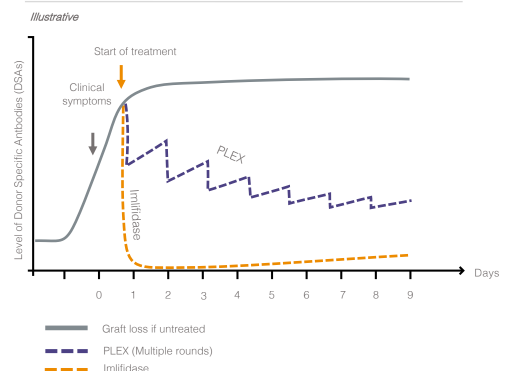
#### STUDY DESIGN

 Randomized, open-label multicenter, active control study, designed to evaluate the safety and efficacy of imilifidase in eliminating DSA in active AMR

#### STATUS

Ongoing 53

## Potential of using imlifidase vs. PLEX in AMR





# Guillain-Barré syndrome is an acute autoimmune attack on the peripheral nervous system

## GBS can affect anyone at any age

- GBS is an acute autoimmune attack on the peripheral nervous system, which rapidly and progressively weakens extremities.
- Only parts of the patients fully recover from GBS, thus a high unmet medical need for new treatments; 40% lose strength and have pain while mortality is 3-7%
- Addressable population of ~ 11,000¹ per year in 7MM²
- Current Standard of Care is treatment with IVIG or PLEX
- The new Phase 2 study is an open-label, single arm, multi-center study evaluating the safety, tolerability and efficacy of imlifidase in GBS patients in combination with standard of care intravenous immunoglobulin (IVIg)
- 4/30 patients enrolled. 6/10 sites are recruiting patients across France, UK and the Netherlands. Enrollment is expected to be completed in H2 2021
- In 2018, the FDA granted Orphan Drug Designation to imlifidase for the treatment of GBS



<sup>&</sup>lt;sup>1</sup> McGrogan et al. Neuroepidemiology 2009;32(2):150-63.

<sup>&</sup>lt;sup>2</sup> 7MM = Seven major markets – US, Germany, UK, France, Spain, Italy, and Japan



## New Phase 2 study initiated in GBS to evaluate safety, tolerability and efficacy of imlifidase in GBS patients

#### CLINICALTRIALS.GOV ID

#### SUBJECTS

#### DOSES/FOLLOW UP TIME

Dosage 0.25mg/kg follow up 180

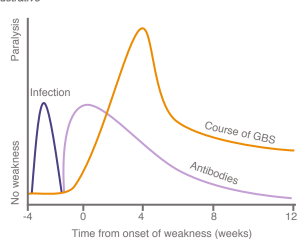
#### MAIN OBJECTTIVES

#### STUDY DESIGN

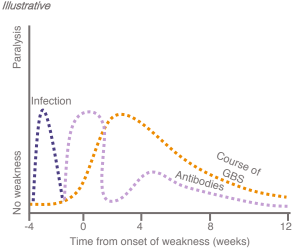
#### STATUS

### Today's Standard of Care IVIg or PLEX





### Potential with imlifidase





Pre-clinical programs

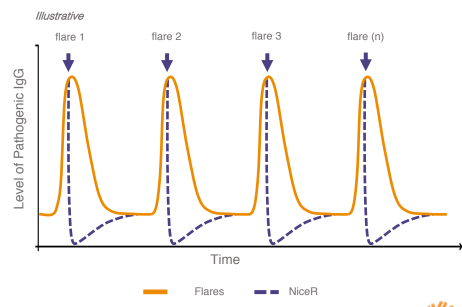


# "NiceR" – new set of enzymes for repeat dosing; potentially enabling treatment of relapsing diseases

## IgG-cleaving enzyme with lower immunogenicity

- Potential application for a broad array of indications, including reoccurring AMR, relapsing autoimmune diseases and oncology
- The first selected promising new drug candidate from the NiceR program is an IgG-cleaving enzyme (cysteine peptidase) with characteristics based on a homolog to imlifidase, but with lowered immunogenicity
- Development of a GMP-manufacturing process has been initiated

## NiceR can potentially inactivate flares



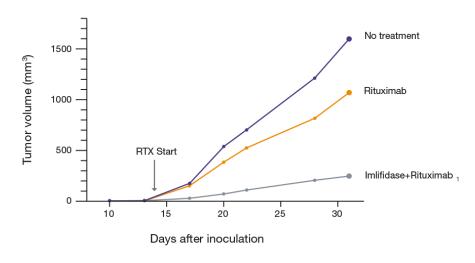


## Imlifidase / NiceR can potentially improve the therapeutic effect in oncology (EnzE)

## Proof of concept demonstrated in vivo for mice

- Enzyme based antibody enhancement through pre-treatment
- The abundance of normal IgG in blood interferes with therapeutic monoclonal antibodies
- Pre-treatment with imlifidase / NiceR has potential to significantly potentiate antibody-based cancer therapies
- Suppressive effect of IVIg on effector cell function abrogated by imlifidase
- Imlifidase can significantly improve the therapeutic effect of

## Mice with human IgG (~9mg/mL)



<sup>&</sup>lt;sup>1</sup> Järnum et al. Mol Cancer Ther 2017:16:1887-1897



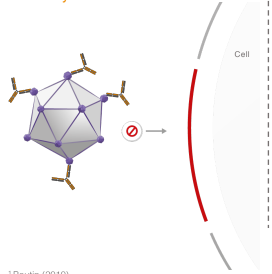
Gene Therapy

## Neutralizing antibodies (Nabs) are immunological barriers in gene therapy

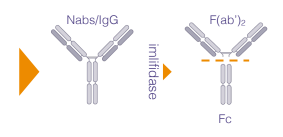
Between approximately 5% and 70%<sup>1,2</sup> of patients considered for gene therapy treatment carry neutralizing anti-AAV antibodies forming a barrier for treatment eligibility

Our hypothesis is that imlifidase has the potential to eliminate neutralizing antibodies as a pre-treatment, prior to the introduction of gene therapy

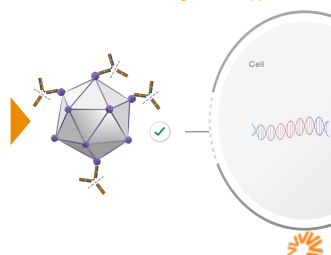
1 Antibodies prevent effective transfer of healthy gene sequence and can be a safety concern



2 Imlifidase is a unique IgG antibodycleaving enzyme that cleaves IgG at the hinge region with extremely high specificity



The idea is to eliminate the neutralizing antibodies as a pretreatment to enable gene therapy



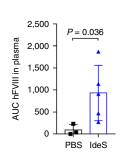
## Imlifidase (IdeS) was highlighted in Nature Medicine<sup>1</sup> with encouraging outcome

Results from preclinical studies with imlifidase (ideS) in gene therapy demonstrate imlifidase as a potential solution to overcome pre-existing antibodies to AAV-based gene therapy

## medicine Imlifidase tested in a IgG-cleaving endopeptidase enables in vivo gene hemophilia mouse model therapy in the presence of anti-AAV neutralizing Imlifidase decreased anti-AAV antibodies and enabled efficient gene transfer 10<sup>7</sup> GLuc activity (RLUs) 10<sup>6</sup> 10<sup>5</sup> 10<sup>4</sup> 10<sup>3</sup> Days post-AAV8 injection

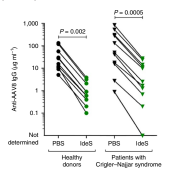
## Imlifidase tested in NHP ahead of AAV vector infusion

 Pre-treatment with imlifidase in anti-AAV positive nonhuman primates (NHP) ahead of AAV vector infusion was safe and resulted in enhanced liver transduction and hFVIII plasma levels



## Imlifidase tested in human plasma samples (GT patients)

 Imlifidase reduced anti-AAV antibody levels from human plasma samples in vitro, incl. plasma from prospective gene therapy trial participants



<sup>1</sup> Nature Medicine <a href="https://doi.org/10.1038/s41591-020-0911-7">https://doi.org/10.1038/s41591-020-0911-7</a> Leborgne et al. Nat Med (2020)

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

Upfront payment

Hansa to receives a USD 10 million

upfront payment from Sarepta for

accessing Hansa's unique IgG antibody-

cleaving enzyme technology (imlifidase)

#### 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



Milestones

Hansa is eligible for a total of

up to USD 397.5 million in

development, regulatory and

sales milestone payments.

**Royalties & Sales** 

Hansa to receive high single-digit to mid-teens

royalties on Sarepta's gene therapy sales

enabled with imlifidase treatment in Nabs

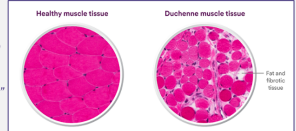
positive patients and book all sales of imlifidase

# Sarepta obtains a global and exclusive license to imlifidase in DMD and LGMD in gene therapy

## About Duchenne muscular dystrophy (DMD)

- Duchenne muscular dystrophy is a rare genetic disease caused by mutation in the DMD gene, encoding for the protein dystrophin
- Muscles in the body become weak and most patients use wheelchair by the age of 12
- Affects one in 3,500 to 5,000 males born worldwide (approximately 400-500 annual male cases in the US) of which approximately 15-20% are estimated to have pre-existing antibodies to AAV-based gene therapy which prevents the patients from being treated with gene therapy

"On average, every day DMD takes the life of a child in the United States..."

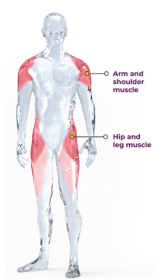


Source: Sarepta Therapeutics

https://investorrelations.sarepta.com/static-files/0c4aca61-9419-45a5-afb1-ff2092644627

## About Limb-girdle muscular dystrophy (LGMD)

- Limb-girdle muscular dystrophy is a group of diseases that cause weakness and wasting of the muscles
- May be caused by a single gene defect affecting specific proteins within muscle cells
- Global prevalence of 1.63 per 100,000 individuals (of which approximately 15-20% are estimated to have pre-existing antibodies to AAV-based gene therapy which prevents patients from being treated with gene therapy



Source: Sarepta Therapeutics

https://investorrelations.sarepta.com/static-files/0c4aca61-9419-45a5-afb1-ff2092644627



## Emerging landscape in gene therapy

## Examples of big pharma and specialized players targeting rare diseases in gene therapy

The list is non-exhaustive



Limb-Girdle (LGMD)

Sarepta Therapeutics



Hemophilia A & Fabry Sangamo Therapeutics Phenylketonuria
Homology
Medicines

LCA (Luxturna)
Spark Therapeutics

Hemophilia A & B

Takeda

OTC-deficiency
Ultragenyx

pharmaceutical

Recessive Dystrophic Epidermolysis Bullosa Abeona Therapeutics

Hemophilia A
Biomarin (BLA)

Pompe AskBio

Hemophilia A & B
Spark Therapeutics

Hemophilia B & Fabry

Hemophilia B uniQure

Freeline Therapeutics

Duchenne (DMD) Duchenne (DMD)
Sarepta Therapeutics

SMA (Zolgensma)
Novartis

type A & B
Abeona Therapeutics

Sanfilippo Syndrome

Gene therapy programs in Clinical phase

73 programs

Market Authorization
2 programs

Today experimental protocols are used based on plasmapheresis, or with immunosuppressants; however these protocols have not demonstrated sufficient efficacy and safety

187 *in vivo* programs are ongoing in gene therapy including 73 clinical stage programs<sup>1</sup>

Two *in vivo* gene therapy products have been approved by FDA: *Luxturna* from Sparks/Roche and *Zolgensma* from Novartis

<sup>1</sup> Alacrita Consulting 2019 estimate based on publicly available data





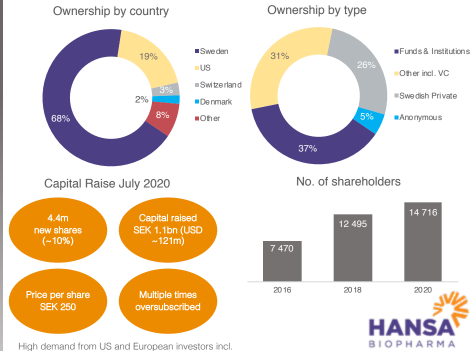
## Ownership in Hansa Biopharma

## Top 10 ownership as per June 30, 2020

Name	No. of shares	Ownership in pct.
NXT2B	5 755 379	14.4
Consonance Capital Management LP	2 478 177	6.2
Invesco	1 999 188	5.0
Thomas Olausson	1 713 474	4.3
Avanza Pension	1 396 176	3.5
Gladiator	1 260 631	3.1
Fourth Swedish National Pension Fund	1 112 044	2.8
Third Swedish National Pension Fund	1 066 470	2.7
Vanguard	938 933	2.3
ClearBridge, LLC	741 306	1.9
Other	21 564 329	54.0
Outstanding A shares in total	40 026 107	100.0

## Classification of ownership

Redmile, Consonance, HBM and Fonden TIN Ny Teknik



## Hansa Biopharma - Market data and share price development

## Market data (July 2020)

Stock Exchange: Nasdaq, Stockholm since Nov 2015

(First North Oct 2007- Nov 2015)

Ticker HNSA

Market Cap: SEK ~10bn (USD ~1bn)

**52-week range**: SEK 59-282 per share

Avg. Daily Turnover: vol ~400k shares

Shares outstanding: ~40m (pre-raise) ~45m (post-raise)

Shareholders ~14,700

Top 5 Shareholders: NXT2B 14.4% As per June 30, 2020 (pre-raise)

Consonance 6.2%

Invesco 5.0%

Thomas Olausson 4.3%

Avanza Pension 3.5%

## 12 months Share price development (July 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
Zoe Karamanoli	RBC (2017)	London	Zoe.Karamanoli@rbccm.com	+44 7834 765119
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

Jul 16, 2020	Interim Report Jan-Jun 2020
Aug 14, 2020	Nordea Small & Mid Cap Seminar, Stockholm (virtual)
Sep 1, 2020	Kempen Road Show, Benelux, Paris & Tel Aviv (virtual)
Sep 3, 2020	Pareto Healthcare Conference, Stockholm (virtual)
Sep 15, 2020	Morgan Stanley Global Healthcare Conference, NYC (virtual)
Sep 16, 2020	BofAML Global Healthcare Conference, London (virtual)
Sep 23, 2020	ABG Small & Mid Cap Seminar, Copenhagen
Oct 22, 2020	Interim Report Jan-Sep 2020
Oct 29, 2020	Hansa Biopharma Capital Markets Day ("SAVE-THE-DATE")
Nov 25, 2020	Ökonomisk Ugebrev Life Science Conference, Copenhagen



