## sequanamedical



# Innovators in the management of fluid overload

liver disease – malignant ascites – heart failure

Investor presentation – November 2020

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### **Disclaimers**

#### Regulatory disclaimer:

- The alfapump® system has not yet received regulatory approval in the United States and Canada. Any statement in this presentation about safety and efficacy of the alfapump® system does not apply to the United States and Canada. In the United States and Canada, the alfapump® system is currently under clinical investigation (POSEIDON Study) and is being studied in adult patients with refractory or recurrent ascites due to cirrhosis. For more information regarding the POSEIDON clinical study see <a href="https://www.poseidonstudy.com">www.poseidonstudy.com</a>.
- The DSR therapy is still in development and it should be noted that any statements regarding safety and efficacy arise from ongoing pre-clinical and clinical investigations which have yet to be completed. The DSR therapy is not currently approved for clinical research in the United States or Canada. There is no link between the DSR therapy and ongoing investigations with the alfapump® system in Europe, the United States or Canada.

#### COVID-19 disclaimer:

- Sequana Medical is closely following the evolution of the COVID-19 global health crisis and is in constant dialogue with its partners to assess the impact and adapt its operations as necessary.
- Sequana Medical has put in place mitigation plans to minimise delays. The impact of increased demands on the healthcare systems, restrictions on non-essential hospital visits and procedures, social-distancing and travel restrictions may result in further delays to execution of clinical studies and impact sales.
- Sequana Medical will continue to update the market as needed and whenever possible.

## **Company Overview**

Founded in 2006

Gent, Belgium (HQ): corporate, clinical, commercial

Zurich, Switzerland: manufacturing, engineering, QA/RA

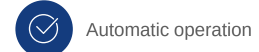
Euronext Brussels: SEQUA



## alfapump® platform

Using the bladder to manage fluid overload

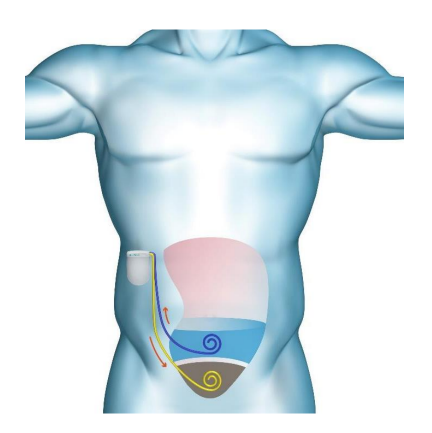


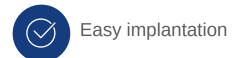








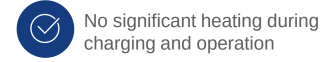












### One platform - two products



### **alfa**pump<sup>®</sup>

### **Liver Disease (NASH)**

Proven step change in liver refractory ascites and malignant ascites

Over 800 devices implanted

> €3 Bn / year market opportunity<sup>(1)</sup>



**POSEIDON** pivotal study ongoing

**Self-commercialisation** 

### alfapump® DSR

## 1

#### **Heart Failure**

Breakthrough approach to fluid overload in heart failure

Clinical proof-of-concept of Direct Sodium Removal (DSR)

> €5 Bn / year market opportunity<sup>(2)</sup>



**RED DESERT repeated dose study ongoing** 

Partnering after US efficacy study

### Built upon proven European clinical & commercial experience

### **NASH drives US market attractiveness**

Stronger competitive position in a much larger and dynamic market



alfapump® market potential

Underlying disease

Patient characteristic

Average age

alfapump competitive positioning

~€0.4 Bn / year

Alcoholic Liver Disease, Hepatitis

"Outside mainstream"

40-50 yr

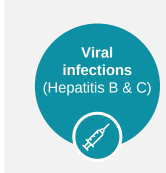






### Liver cirrhosis and refractory ascites

A key complication of liver cirrhosis, with a dramatic impact on quality of life

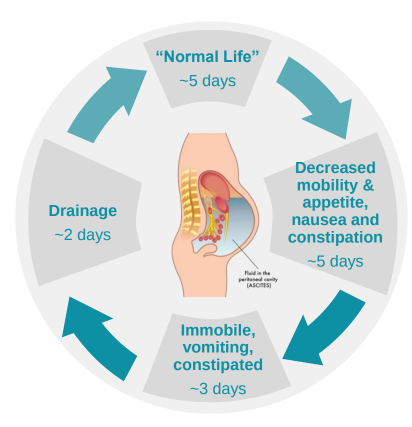


Alcoholic Liver Disease

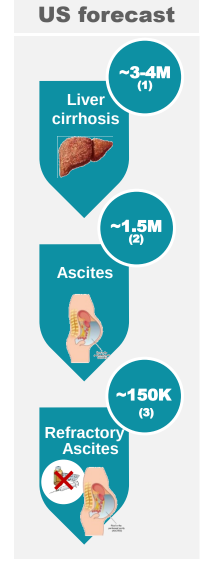


Non-Alcoholic Steatohepatitis (NASH)





Typical patient life<sup>(4)</sup>



Source 3: Ginès et al., NEJM 2004: refractory ascites occurs in 5-10% patients with ascites

### **Cancer and malignant ascites**

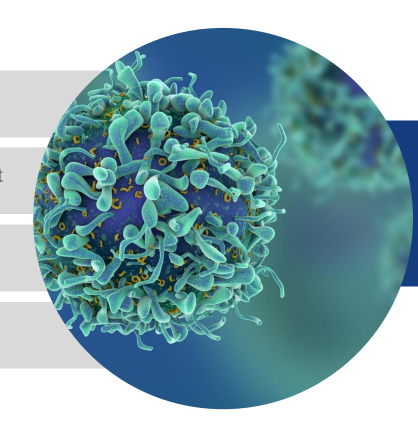
Severe complication of late-stage cancers

Fluid accumulation in the abdomen due to **drainage of lymph system** 

**Breast and ovarian cancer** have longest survival with ascites<sup>(1)</sup>

Severe impact on quality of life

Reduces ability to undergo anti-cancer treatment



Malignant ascites due to breast and ovarian cancer<sup>(2)</sup>:

EU5: ~18K

US: ~16K



Clear unmet need for improving Quality of Life and the ability to increase cancer treatment intensity

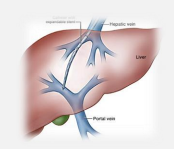
## Severe limitations of existing therapies

### **Diet / Diuretics**



Resistance, Complications

## **Transjugular Intrahepatic Portosystemic Shunt (TIPS)**



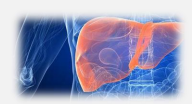
Complications, Contraindications

## **Drainage ("Large Volume Paracentesis / LVP")**



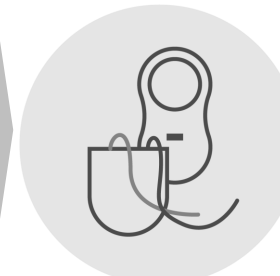
Painful, Poor Quality of Life, Short Term Benefit

### **Liver transplantation**



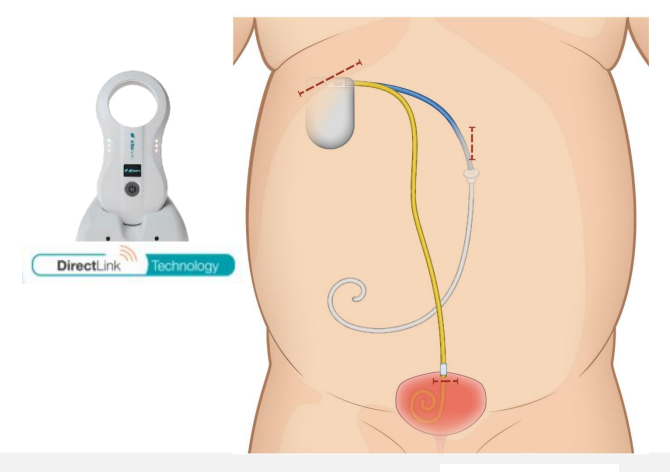
High Cost, Limited Availability





## alfapump® for long-term treatment

Over 800 implants and hundreds of years of patient experience













## Strong clinical and economic rationale

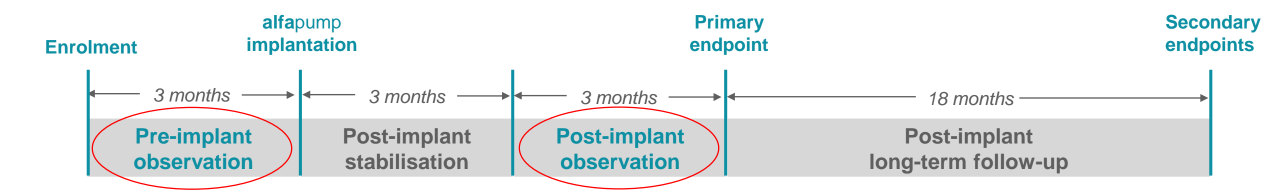
### Significant reduction in regular drainage leads to:

- Reduced burden of disease
- Improved patient QoL
- Cost savings for hospitals and payers

<sup>\*</sup> Management estimate of US treatment costs, assuming no complications

## North American Pivotal Study (POSEIDON) underway

Pivotal Cohort of up to 50 patients implanted; Roll-In ("training") cohort of up to 30 patients



Primary efficacy: 1) 50% reduction in average monthly frequency of Therapeutic Paracentesis ("TP") post-implant vs. pre-implant

2) 50% of patients achieve a 50% reduction in the requirement for TP post-implant vs. pre-implant

**Primary safety:** Rate of **alfa**pump related re-interventions adjudicated by the Clinical Events Committee (CEC)

**Secondary:** QoL (SF36, Ascites-Q), nutritional status, health economics, safety (device and/or procedure-related AEs), survival

# Positive outcomes against all primary endpoints in first 13 Roll-In patients

Substantial reduction in therapeutic paracentesis (TP) and safety profile in line with expectations

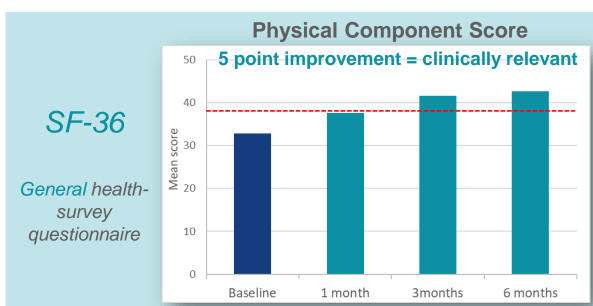
### **EFFICACY**

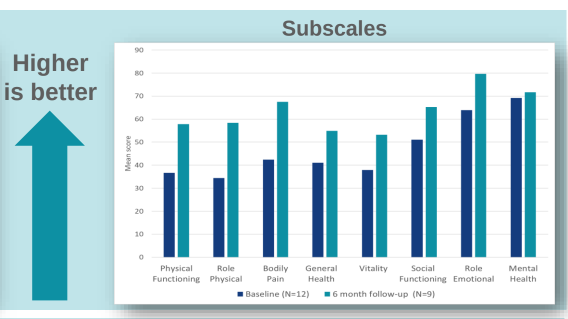
Mean values post-implant vs. pre-implant	N = 13
% reduction in frequency of TP	> 90%
% patients with >50% reduction in TP	100%

#### **SAFETY**

- Safety profile of the alfapump consistent with previously reported data
- Adjudication process by the Clinical Events Committee for two alfapump® explants ongoing

### Indication of fast and persistent improvement in Quality of Life



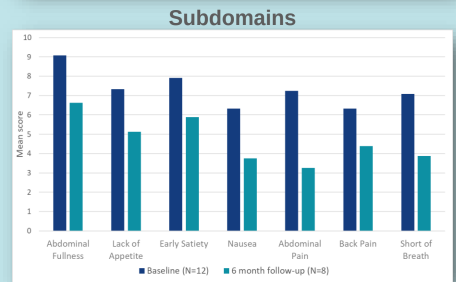


## Ascites Q

Specific healthsurvey questionnaire for ascites

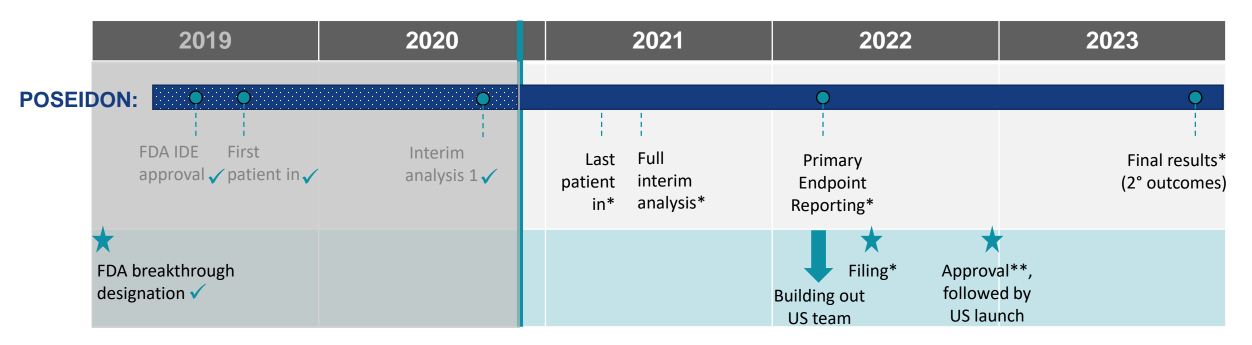






## Strong progress towards alfapump® US approval

**Targeting Announcement of Primary Endpoint in Q1 2022** 



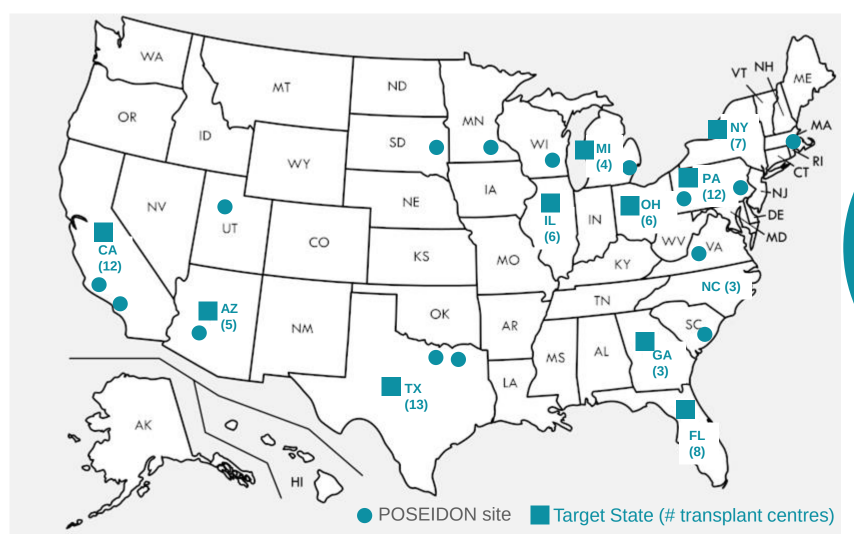


Proposed CMS rule for automatic Medicare coverage of breakthrough devices for four years post-approval

<sup>\*</sup> Subject to further developments related to the ongoing COVID-19 pandemic

<sup>\*\*</sup> Subject to FDA review

## **Self-commercialisation in US through specialty salesforce**

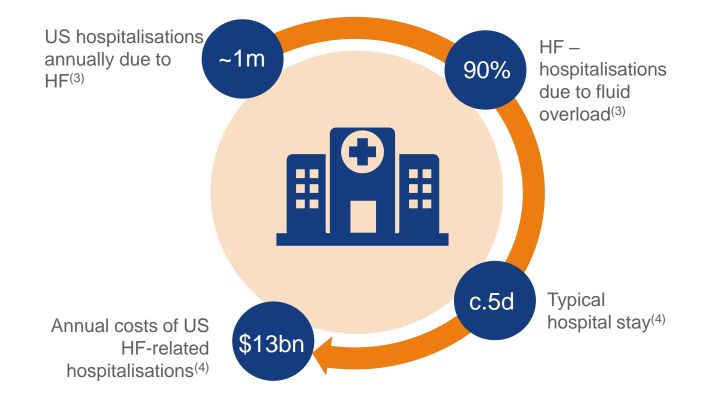






# Fluid overload in heart failure – major clinical problem and key driver of costs





- 40% of heart failure patients on IV loop diuretics have a poor response<sup>(1)</sup>
- 24% re-admission rate at 30 days<sup>(2)</sup>

## **Direct Sodium Removal (DSR)**

Proprietary approach to fluid overload – supported by interim RED DESERT clinical data

We remove the sodium and then the body "does the math" to maintain serum sodium balance



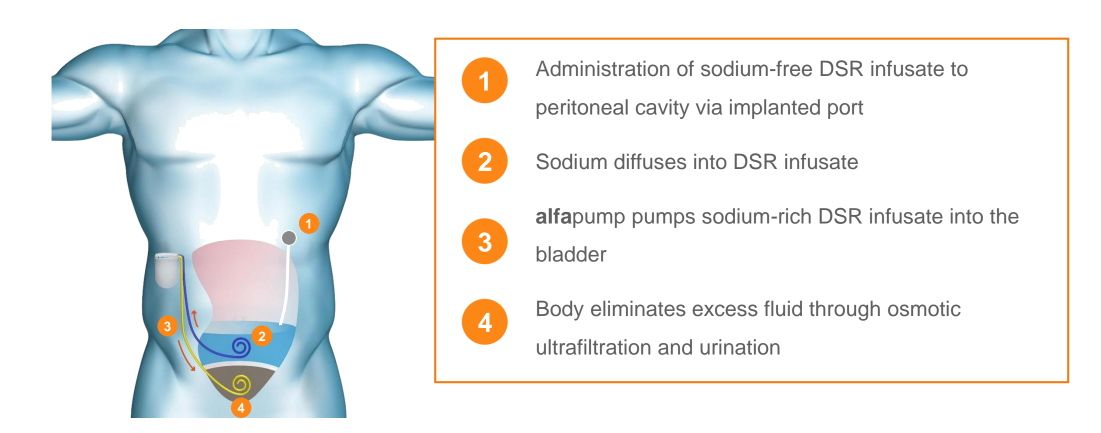
"DSR represents a new potential therapy for nonrenal sodium and fluid removal in edematous disorders such as heart failure" First in Human Experience with
Peritoneal Direct Sodium Removal Using
a Zero Sodium Solution: A New
Candidate Therapy for Volume Overload

Veena S. Rao, Jeffrey M. Turner, Matthew Griffin, Devin Mahoney, Jennifer Asher, Sangchoon Jeon, Peter S. Yoo, Nabil Boutagy, Attila Feher, Albert Sinusas, F. Perry Wilson, ... Show all Authors

Originally published 8 Jan 2020  $\mid$  https://doi.org/10.1161/CIRCULATIONAHA.119.043062  $\mid$  Circulation. ;0:null

## alfapump® DSR

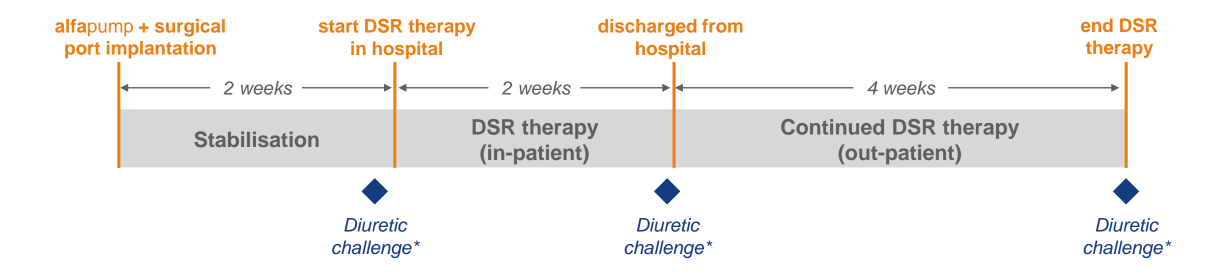
Potential chronic therapy for heart failure patients with fluid overload that are not well controlled on diuretics



Fundamental patents to reduce fluid overload in heart failure allowed in US and Europe

### **RED DESERT: Study design**

Repeated dose proof-of-concept study of alfapump® DSR in diuretic-resistant heart failure patients



√ Positive interim results (5 patients) reported

Top-line results (up to 10 patients) expected in H1 2021

<sup>\*</sup> intravenous dose of 40mg dose furosemide

## Interim RED DESERT: Strong safety & efficacy results from first 5 patients

#### **SAFETY**

- Implant procedure of alfapump® DSR and repeated dosing of DSR therapy were well-tolerated
- No clinically significant changes in serum sodium levels / no progressive hyponatremia
- Reported adverse events were manageable

#### **EFFICACY**

- No diuretics required in any of the patients during 6-week alfapump DSR treatment
- Reduced doses of DSR therapy and / or less frequent DSR dosing in majority of patients
  - ⇒ maintaining stable to lower weight and NT-proBNP compared to baseline

## Interim RED DESERT: Restore normal kidney response

Loop diuretic responsiveness restored to near normal levels in all 5 patients

- Diuretic response assessed by 6-hour excretion of fluid and sodium following intravenous administration of 40mg furosemide
  - ⇒ Baseline: objectively poor diuretic response
  - ⇒ End of 6-week study period: more than doubling of sodium excretion (near normal levels)
- Long-lasting diuretic responsiveness after completion of alfapump® DSR therapy
  - ⇒ dramatic reduction in loop diuretics requirements in majority of patients

Based on these interim data, it appears that DSR therapy is not just an alternative means to remove sodium and water, it restores kidney response to near normal levels – opening up whole new ways it can be used

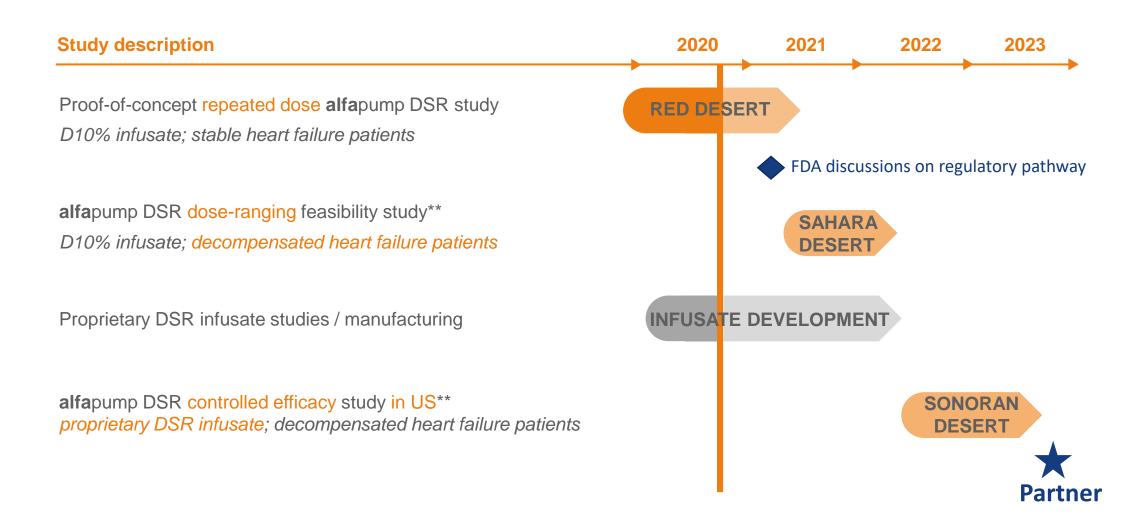
## **Developing high value proprietary DSR Infusate**

- D10% was chosen as the initial DSR infusate for fastest proof-of-concept
- We are developing our proprietary next-generation DSR infusate:



- ✓ Improved therapeutic profile compared to D10%
- ✓ IP protected
- ✓ Recurring revenue from high gross margin consumable

## alfapump® DSR development strategy\*



<sup>\*</sup> Timelines subject to further developments related to the ongoing COVID-19 pandemic

<sup>\*\*</sup> Subject to change and/or feedback from applicable regulatory authorities



### **Strong organisation**

Highly experienced leadership team supported by committed and well-reputed shareholders

#### **Executive team:**



lan Crosbie Chief Executive Officer



**Kirsten Van Bockstaele** Chief Financial Officer



Oliver Gödje Chief Medical Officer



**Gijs Klarenbeek** Senior Medical Advisor



Martijn Blom Chief Commercial Officer



**Timur Resch** Global VP QM/QA/RA

#### **Board of Directors:**



Pierre Chauvineau Board Chairman



Jason Hannon Director



lan Crosbie Chief Executive Officer



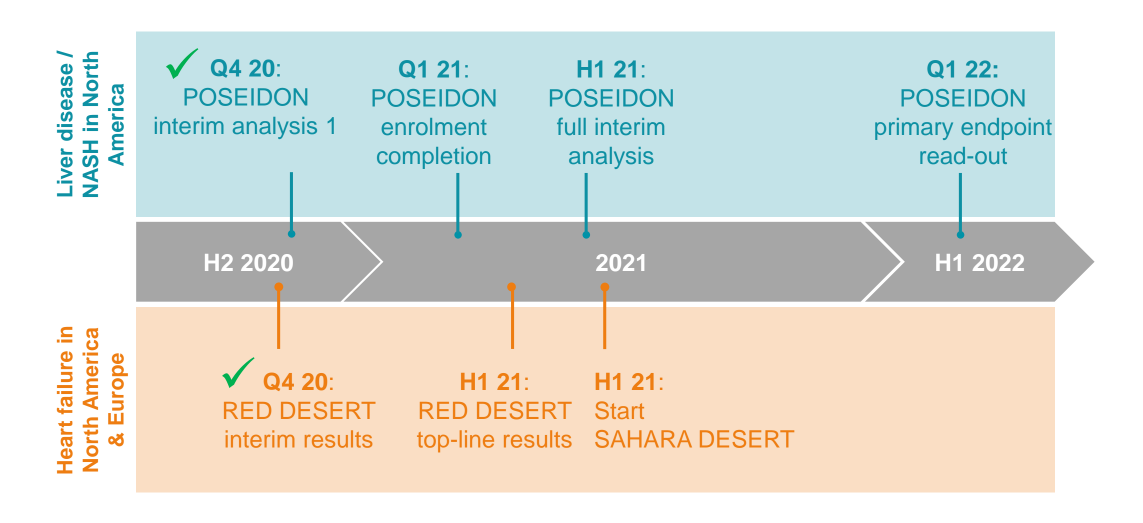


Wim Ottevaere Director



Erik Amble
Director

## **Expected core value drivers & outlook**



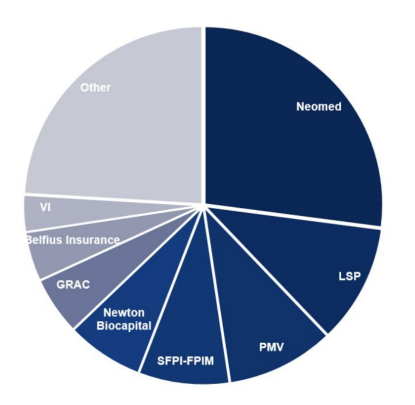


## Back-up

### **Shareholders base and financial overview**

Ticker: SEQUA - Euronext Brussels

- Outstanding shares: 15.8M
- Outstanding share options & warrants: 1.9M authorised of which 0.9M granted



- Analysts:
  - KBC Securities Lenny Van Steenhuyse
  - Kempen Ingrid Gafanhão
  - Kepler Cheuvreux Matthias Maenhaut
  - Mirabaud Daniel Jelovcan
- Cash (30 June 2020): €14.9M
- Debt financing in July 2020: €7.3M
- Cash runway into H2 2021



### **POSEIDON – study cohorts**

Patients with recurrent or refractory ascites due to liver cirrhosis in up to 20 centres across US and Canada

### Two study cohorts with the same inclusion / exclusion criteria

- 1 Pivotal Cohort
  - Up to 50 patients implanted with the **alfa**pump<sup>®</sup>
  - For primary and secondary endpoint analysis
- 2 Roll-In Cohort 📦 enables us to report interim data
  - Up to 30 patients implanted with the alfapump
  - To teach clinicians and medical teams at new centres how to use the alfapump



## Cirrhotic patients with recurrent or refractory ascites

First 13 patients in Roll-In Cohort of the POSEIDON study

Age (mean)	65 y
MELD score (mean ± SD)	10.5 ± 4.6
Cirrhosis etiology	
- Alcohol	- 61.5%
- NASH	- 23.1%
- Hepatitis C	- 7.7%
- Alcohol, Hepatitis C, and Hepatitis B	- 7.7%
TP per month prior to study (mean ± SD)	3.4 ± 1.8

Willingness to treat earlier stage patients?

NASH is already an important driver of this market

N. American patients appear to have more TP / month compared to Europe

MELD: Model for End-stage Liver Disease; SD: Standard Deviation; NASH: Non-Alcoholic Steatohepatitis; TP: Therapeutic Paracentes



## Leading experts as Heart Failure Scientific Advisors



Dr. Maria Rosa Costanzo

Medical Director of the Edward Center for Advanced Heart Failure Medical Director Heart Failure Research for the Advocate Heart Institute



**Dr. Wilson Tang** 

Professor of Medicine at Cleveland Clinic Lerner College of Medicine at Case Western Reserve University



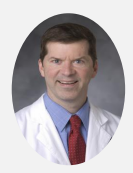
Dr. Javed Butler

Professor and Chairman of the Department of Medicine at the University of Mississippi Medical Center



**Dr. Jeffrey Testani** 

Associate Professor of Medicine and Director of Heart Failure Research at Yale University School of Medicine



Dr. Michael Felker

Professor of Medicine in the Division of Cardiology at Duke University School of Medicine Director of Cardiovascular Research at the Duke Clinical Research Institute and Vice-Chief for Clinical Research in the Division of Cardiology



Dr. Udelson

Chief of the Division of Cardiology at Tufts Medical Center Professor of Medicine and Radiology at Tufts University School of Medicine

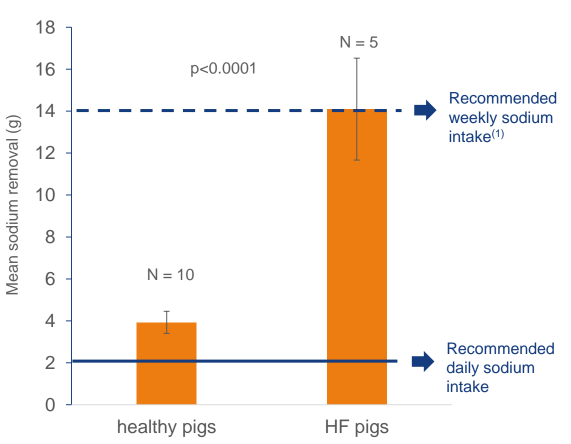


### **DSR pre-clinical Proof-of-Concept**

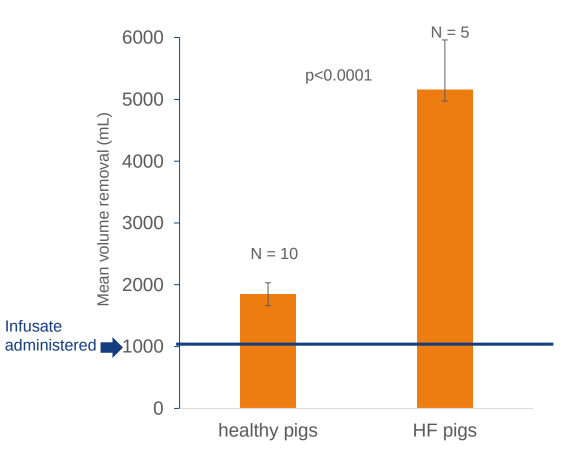
Yale

Clinically relevant sodium and fluid removal

### Clinically relevant removal of sodium



### Effective fluid removal

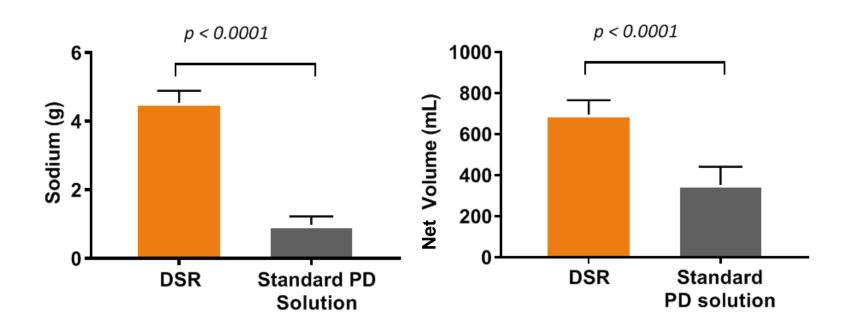




# DSR first-in-human study met primary and secondary endpoints



- $\checkmark$ 
  - DSR therapy was safe & well-tolerated with no adverse events or significant discomfort
- ✓ Substantially higher sodium removal with DSR vs standard Peritoneal Dialysis (PD) solution
- ✓ Minimal inter-patient variability



Results presented at

key Cardiac

Conferences and

published in

Circulation



### Clinical development strategy

Exciting impact on diuretic response requires additional investigation to support value of DSR therapy

### **RED DESERT – repeated dose study in stable heart failure patients**

Enrol up to five additional patients, with top-line data expected in H1 2021

### SAHARA DESERT – dose-ranging study in decompensated heart failure patients

- Move into decompensated heart failure patients with residual congestion
- Dose ranging to learn more about improvement in diuretic response and durability of effect
- Key learnings to be taken into US controlled efficacy study
- D10% as DSR infusate

### SONORAN DESERT - US study vs. stand of care with proprietary DSR infusate

- Controlled versus standard of care, in decompensated patients with residual congestion
- Treatment algorithm built upon learnings from SAHARA DESERT
- Paves the way and de-risks FDA pivotal study
- Sequana Medical proprietary DSR infusate

