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Oppenheimer Healthcare Conference – 17 March 2022

Ian Crosbie, CEO

Innovators in the treatment of diuretic-resistant fluid overload

liver disease malignant ascites heart failure

Disclaimers

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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 visit www.poseidonstudy.com.
- DSR® therapy is still under 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. DSR® therapy is currently not approved for clinical research in the United States or Canada. There is no link between 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 operations accordingly.
- Sequana Medical has put in place mitigation plans to minimise delays. The impact of increased demands on the healthcare systems, limitations 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.

Note:

alfapump® is a registered trademark. DSR® and alfapump DSR® are registered trademarks in the Benelux, China, the EU, United Kingdom, and Hong Kong.

Uniquely positioned in two large markets



- alfapump[®] in liver disease over €3 Bn / year ⁽¹⁾
 - NASH changing liver cirrhosis market and driving growth
 - Approved in EU / FDA breakthrough designation in US
 - US pivotal study fully enrolled / positive interim data / primary endpoint Q4 '22
 - Direct commercialisation in US

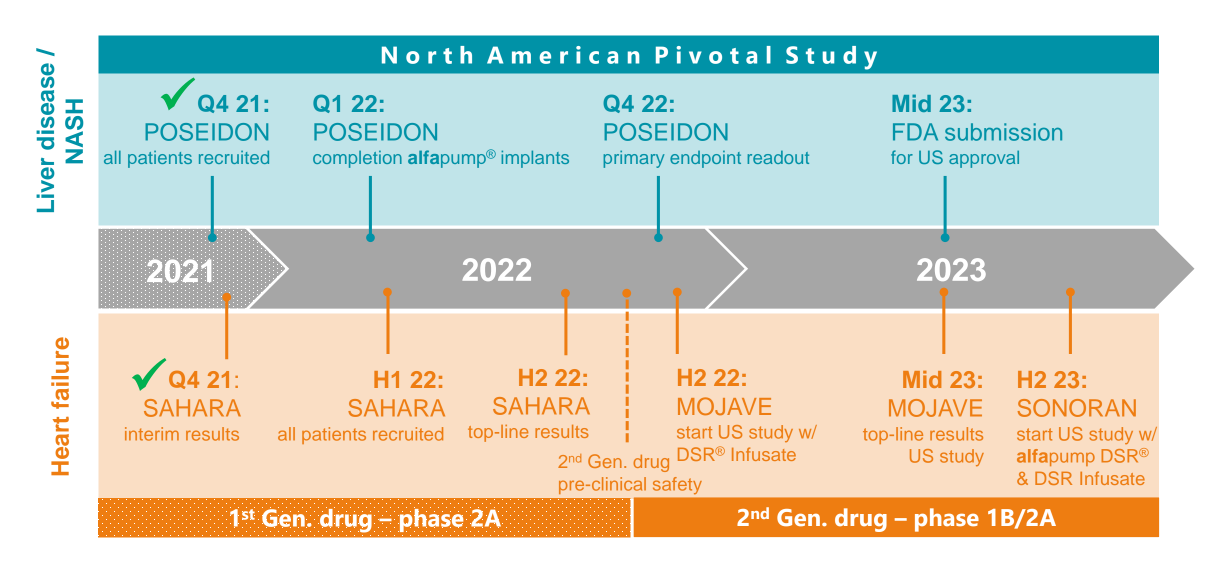


- DSR® in heart failure over €5 Bn / year (2)
 - Clinical proof-of-concept established
 - Ph. 2A 1st Gen. drug positive interim data
 - Low-risk proprietary 2nd Gen. drug in development
 - Partnering after US efficacy study



- Proprietary technologies treating diuretic-resistant fluid overload
 - Key clinical problem in liver disease, heart failure, renal failure and cancer
 - Diuretic-resistance is common alternatives have significant disadvantages
- Strong granted IP portfolio

Strong outlook for value drivers



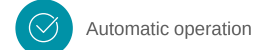


alfapump®

Eliminating fluid from the peritoneal cavity – working in partnership with the bladder

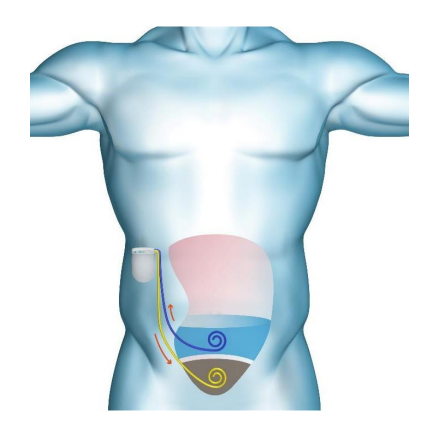








- Settings wirelessly adjusted
- Remote data monitoring

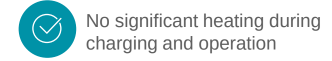








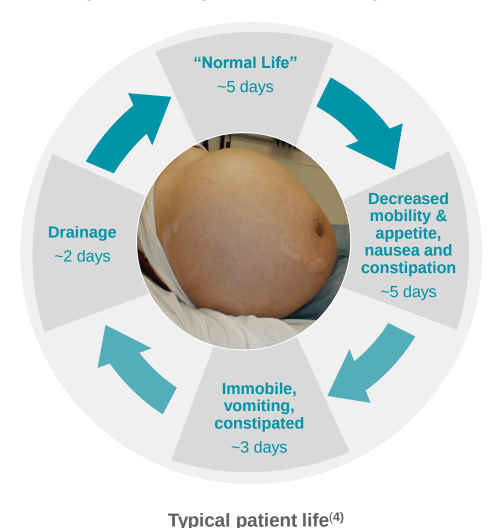


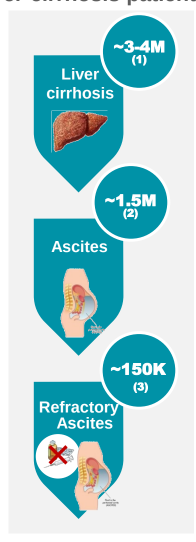


Refractory ascites - key complication of liver cirrhosis

Fatty liver disease / NASH is driving dramatic growth and change in attitudes to liver cirrhosis patients







NASH transforming the face of liver cirrhosis

In US, liver cirrhosis is transitioning to a mainstream disease requiring modern treatment options

alfapump® market potential

Underlying disease

Patient characteristic



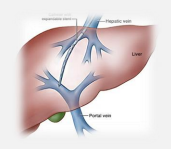
Limitations of existing therapies

Drainage ("Large Volume Paracentesis / LVP")



Painful, Poor Quality of Life, Short Term Benefit

Transjugular Intrahepatic Portosystemic Shunt (TIPS)



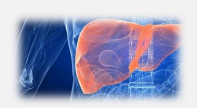
Complications, Contraindications

Permanent Catheter System



External Catheter, Risk for Infections / Blockage

Liver transplantation



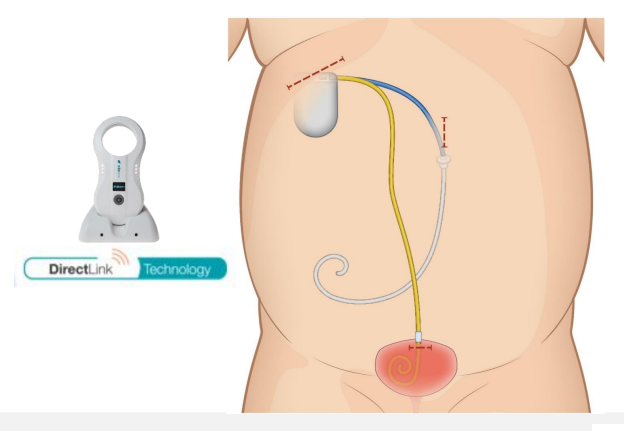
High Cost, Limited Availability

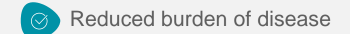




alfapump® strong clinical and economic rationale

Over 900 implants and hundreds of years of patient experience







Cost savings for hospitals and payers

Estimated treatment cost / patient*:

LVP: ~\$54K alfapump®: ~\$35K

~\$1.8K / LVP⁽¹⁾
2 LVP / month
15 months

~\$25K / **alfa**pump

~\$10K / implantation

* Management estimate of US treatment costs, assuming no complications QoL: Quality of Life; LVP: Large Volume Paracentesis





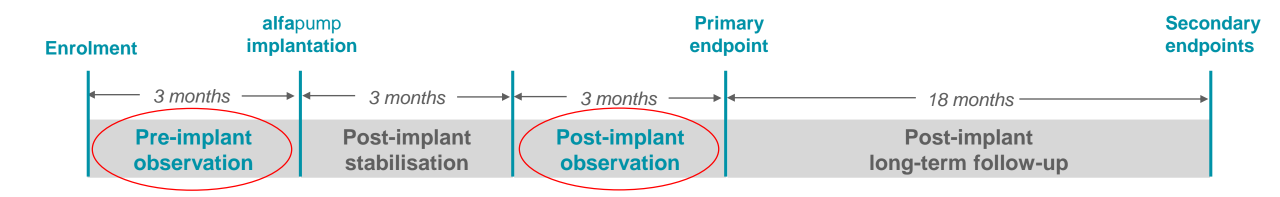






North American Pivotal Study (POSEIDON) underway

Pivotal Cohort of up to 50 implanted patients; Roll-In ("training") cohort of up to 40 patients



POSEIDON Study Endpoints

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

Interim POSEIDON: Positive for primary endpoints

Data from 26 Roll-In patients

EFFICACY

- ✓ Over 90% reduction in mean Therapeutic Paracentesis (TP) frequency (primary endpoint >50% reduction)
- ✓ 100% patients with > 50% reduction in mean TP frequency per month (primary endpoint >50% of patients)

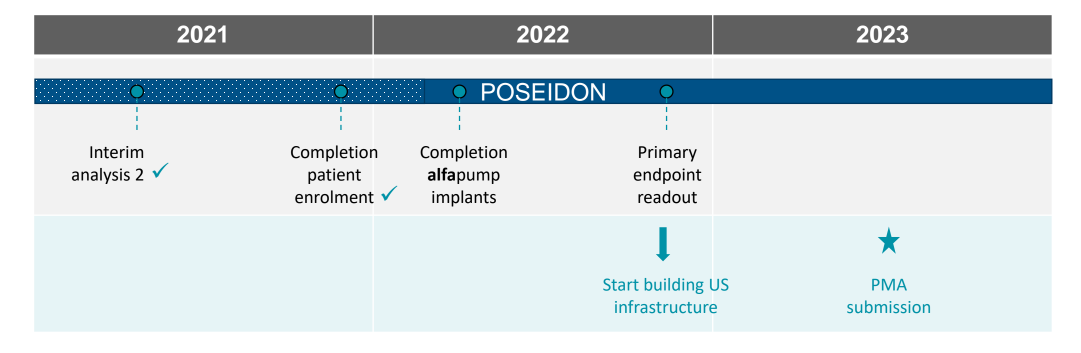
SAFETY

✓ In line with expectations – 3 composite primary safety events

QUALITY OF LIFE

✓ Clinically important improvement maintained for up to 12 months post-implantation

Pursuing North American alfapump® approval



NTAP for breakthrough devices de-risks reimbursement in key Medicare population



US commercialisation through our specialty salesforce





Initial focus on key

transplant centres

~50-person team:

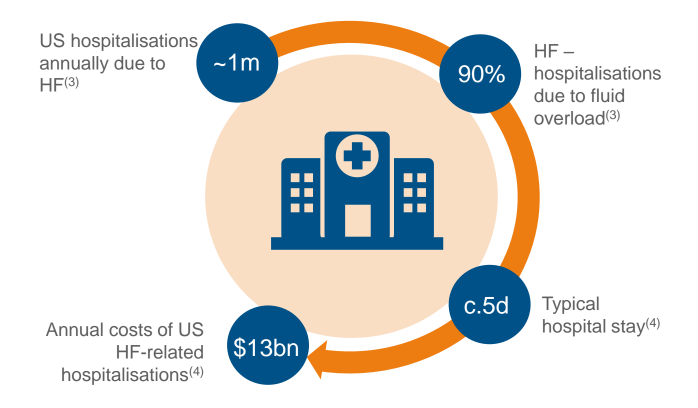
35 sales reps, 10 clinical,

5 corporate



Diuretic-resistant congestion in heart failure

Clear unmet clinical need and driver of costs for heart failure patients



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

Direct Sodium Removal (DSR®) platform

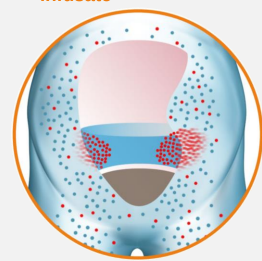
Eliminating fluid spread across the body – working in partnership with the kidneys



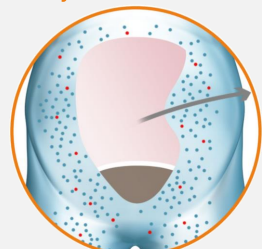




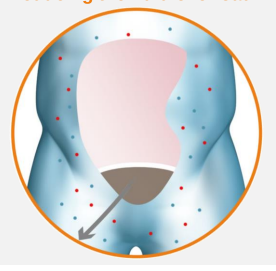
- 1 Sodium-free DSR infusate administered to peritoneal cavity
- 2 Sodium diffuses from body into DSR infusate



3 DSR infusate + extracted sodium removed from the body



4 Body eliminates free water to restore sodium balance, reducing the fluid overload

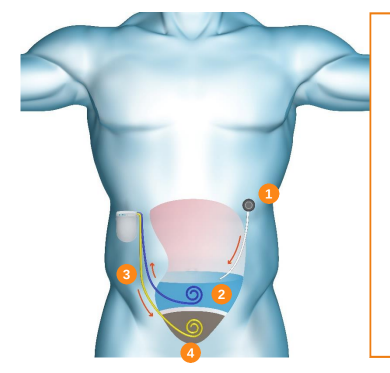


water



alfapump DSR® leveraging proven alfapump® platform

Fully implanted system for long-term DSR® therapy – keeping patients out of the hospital



- Sodium-free DSR infusate administered to peritoneal cavity via implanted subcutaneous port
- 2 Sodium diffuses into DSR infusate
- 3 alfapump pumps sodium-rich DSR infusate into the bladder
- Body eliminates excess fluid through osmotic ultrafiltration and urination

RED DESERT: Successful Proof-of-Concept Study

8 euvolemic heart failure patients on high dose diuretics treated with DSR 3x per week up to 6 weeks

Highly effective management of fluid and sodium balance

Generally safe and well tolerated; no clinically relevant hyponatremia

Presented as Late-Breaker and Highlight at Heart Failure 2021

Significant improvement in cardio-renal function

- 30% decrease* in NT-proBNP** (p<0.001)
- 22% increase* in eGFR** (p<0.001) / 22% decrease* in creatinine** (p<0.001)

Dramatic and sustained improvement in diuretic response

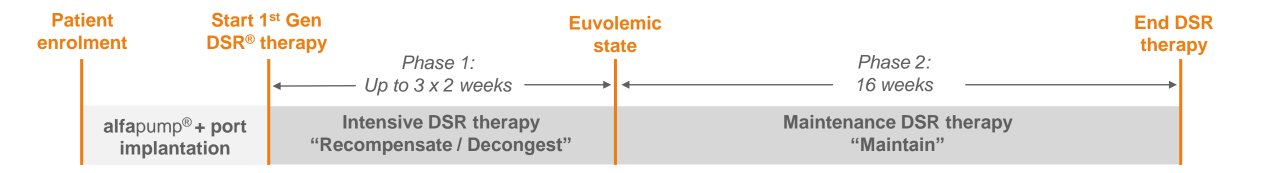
- End of 6-week study: over 150% increase** in diuretic response***
- Long-term follow-up (9-19 months after study completion): 40-96% reduction in diuretic dose at last visit during follow-up

"Simultaneous normalisation of diuretic response and improvement in cardio-renal status is a never before seen treatment effect" – Dr. Testani, Yale

^{*} Paired statistical analysis of patients with baseline and D42 value (N=7); ** mean value; ***assessed by 6-hour excretion of sodium after IV administration of 40mg furosemide NT-proBNP: N-terminal-pro hormone B-type Natriuretic Peptide (analysed in local lab); eGFR: estimated glomerular filtration rate

SAHARA DESERT: Ph. 2A in target patient population

20 decompensated heart failure patients with persistent congestion on high dose diuretics – ongoing



Study Endpoints

- **Primary:** safety and tolerability of **alfa**pump DSR[®] therapy
- Secondary: feasibility of DSR therapy to restore and maintain euvolemia without additional loop diuretics
- Exploratory: evaluate potential impact of SGLT-2 inhibitors on DSR therapy*

SAHARA DESERT: Ph. 2A positive interim data

Indication of strong safety and efficacy in 6 patients

Safe, effective and rapid elimination of persistent congestion and restoration of euvolemia without any loop diuretics

Mean weight loss of ~6kg (=7% of body weight) vs. baseline

Considerable benefit in cardio-renal status

- Reduction* in NT-proBNP >30% vs. baseline
- eGFR* and creatinine* similar to baseline
 - Worsening in kidney function is normally expected during significant volume removal

Dramatic improvement in diuretic responsiveness for months post-treatment

- End of phase 1 (n=6***): more than doubling* of sodium excretion** (near normal levels)
- 3 months* after end of Phase 1 (n=4): less than 10% of their baseline loop diuretic dose

"These interim results are highly encouraging and could potentially provide a course of therapy for severely ill diuretic-resistant heart failure patients with persistent congestion where alternative treatment options are currently exceedingly limited" – Dr. Testani

Proprietary heart failure drug development programme

Red Desert PoC

alfapump

8 stable HF

· Pos. outcome

patients

1st Gen. Infusate +



Sahara Desert Ph. 2A ongoing



- Ph. 2A ongoing
- 1st Gen. Infusate + alfapump
- Up to 20
 decompensated
 HF patients
- · Pos. interim data

2nd Gen. Infusate Development

Animal GLP

in H2 '22

and CMC data



Mojave Desert Ph. 1B/2A (US)



Sonoran Desert Ph. 2A (US)

- 2nd Gen. Infusate (no alfapump)
- Decompensated HF patients
- 2nd Gen. Infusate
 - + alfapump
- Decompensated HF patients

1st Generation Infusate

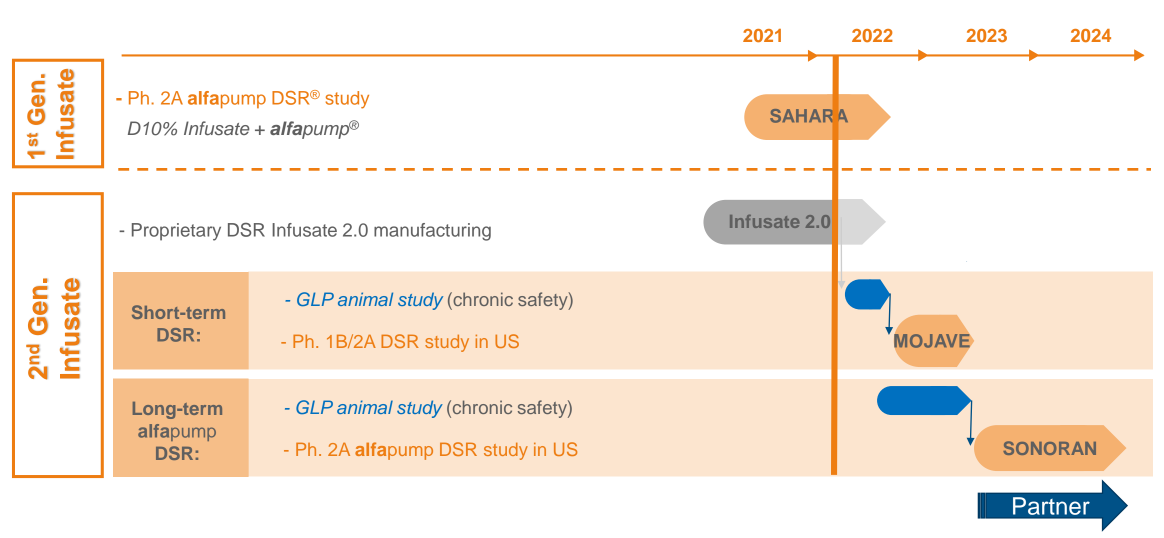
- ✓ Sodium-free D10% (off-the-shelf)
- ✓ Clinical Proof-of-Concept
- ✓ Rapid clinical path

2nd Generation Infusate

- ✓ Sodium-free dextrose / icodextrin (proprietary)
- ✓ Improved therapeutic profile
- ✓ Favorable safety profile
- ✓ IP protection drives recurring revenue from high gross margin consumable

DSR® – plan to partner after US efficacy study

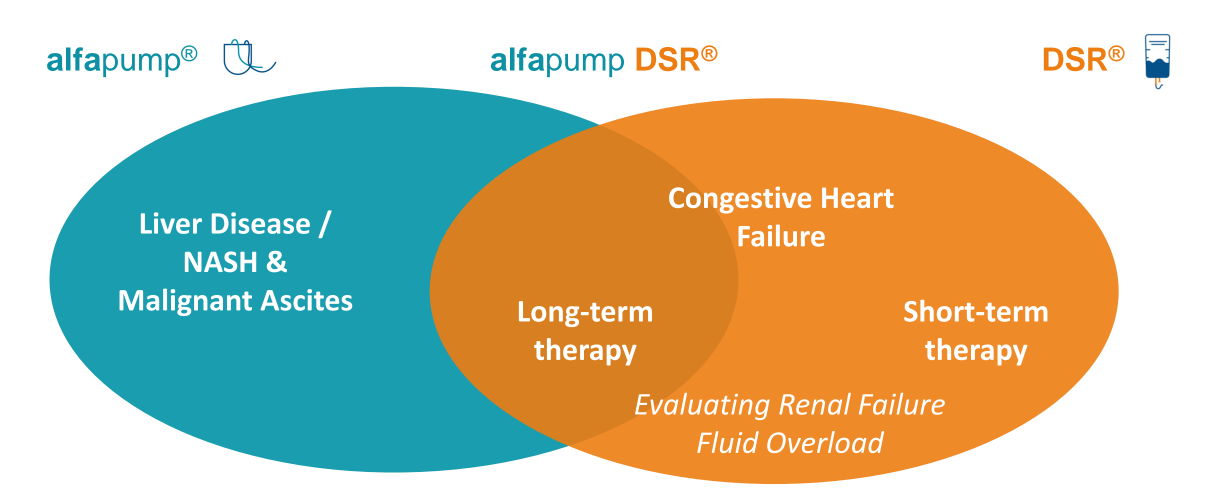
Step-by-step approach to introduction of breakthrough therapy in decompensated HF patients





Building on our two proprietary platforms

Complementary approaches to diuretic-resistant fluid overload



Strong outlook for value drivers

