Nyxoah November 2022



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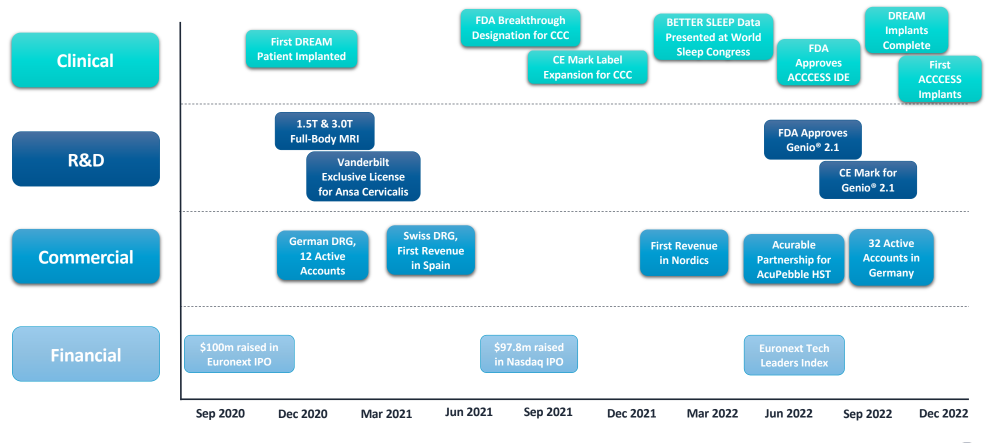
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Investment Highlights

Significant Underpenetrated Global OSA Market Opportunity	 425m moderate-to-severe OSA patients globally eligible for treatment >1m OSA patients eligible for HGNS treatment annually in US and Europe HGNS treatment solution now fully accepted by physicians and payors
Innovative Patient-Centric Technology	 Single-incision procedure with a scalable technology platform Bilateral stimulation with external power source and no lead tunneling Full-body 1.5T and 3.0T MRI compatibility
Growing Body of Strong Clinical Data	 BETTER SLEEP demonstrated safety and efficacy in both non-CCC and CCC patients DREAM IDE pivotal trial to secure US regulatory approval in H1 2024 ACCCESS IDE study to secure US approval for CCC patients in H1 2025
Targeted Commercial Strategy	 CE Mark with first European sales in 2021 Commercial proof of concept through expected market leadership in Germany in Q4 2022 US pre-commercial activities focused on market access and payor coverage
Healthy Financial Position	 Dual-listed (NYXH) company trading on Nasdaq and EuroNext stock exchanges €115 million cash balance after two successful IPOs, which brought in top tier investors Ample liquidity to fund key R&D, clinical, and commercial priorities into 2024

Executing on Our Strategic Priorities Euronext IPO – Present



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Experienced Board and Management Team



Robert Taub Founder, Chairman

- Serial entrepreneur in the pharmaceutical and medical fields
- Co-founded and co-managed Octapharma Human plasma protein company
- Founded and managed Omrix Biopharmaceuticals NASDAQ IPO, followed by the acquisition by J&J
- Early investor and chairman of Neuroderm, a Parkinson's disease pharmaceutical company – IPO on NASDAQ and later sale to Mitsubishi-Tanabe



Olivier Taelman Chief Executive Officer

- Experienced Medtech leader
- 7 years in pharmaceutical healthcare at Eli Lilly and Sanofi Aventis leading specific Business Units
- 18 years within the field of Medtech neuromodulation at Medtronic, managing EMEA at Stryker NeuroVascular and serving a neuromodulation company Nevro where he was responsible for building the European business during the successful NASDAQ IPO
- Joined Nyxoah in July 2019 as Chief Operating and Commercial Officer, subsequently being appointed as CEO in November 2019.



Loïc Moreau Chief Financial Officer

- Experienced Finance leader
- 13 years in Pharmaceutical healthcare at GSK with roles in Corporate Development/ M&A (UK), Finance R&D (UK) or Country CFO where he notably led and structured the various support functions for GSK business in France (€1bn+ turnover)
- Started his career at EY (external audit) followed by PwC (Corporate Finance)
- Joined Nyxoah in 2022 to take the leadership of the finance department.

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Large and Underpenetrated Global OSA and HGNS Market Opportunity

Worldwide Obstructive Sleep Apnea Prevalence 936 Million 936M individuals (30-69 year) are estimated to suffer from OSA¹ 425 Million 425M suffer from moderate to severe OSA, requiring therapy¹ Increasing prevalence of OSA due to rise in obesity Undiagnosed OSA 80% 20% Only 20% OSA patients are diagnosed²

Benjafield, Adam V et al. Estimation of the global prevalence and burden of obstructive sleep apnoea: a literature-based analysis. Lancet Respir Med 2019
 Harvard Medical School Division of Sleep Medicine, The Price of Fatigue - The surprising economic costs of unmanaged sleep apnea, December 2010
 Presents annual revenue growth for Inspire Medical. Inspire Medical corporate presentation – February 2022

Hypoglossal Nerve Stimulation Market Opportunity

>1 million eligible annually in key geographies

- US: 510,000 eligible patients annually
- Europe: 500,000 eligible patients annually

< 25,000 received HGNS as treatment

- First HGNS CE-Mark approval in 2010 FDA authorization in 2014
- Low awareness on neurostimulation as an OSA solution
- Limited reimbursement

+70% CAGR HGNS revenue 2016 - 20213

- Endorsed by the global sleep and ENT medical communities
- Accepted by US/EU payors
- Embraced by OSA patient association groups

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A Disruptive Patient-Centric Solution TODAY...







27.5 mm

Weight 2,96 grams



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Our Vision: Improving Lives Through Restful Nights

Connectivity & Remote Care

- Connectivity with remote smartphone application
- Cloud based access for patients and physicians

Patient Engagement

• Feedback to / from the patient for optimized compliance & comfort

Predictable Outcomes & Auto-Titration

- Patient acclimation through amplitude adjustment @ home
- Sensors for adaptive therapy & auto-titration











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The Genio[®] Duty Cycle Approach to Hypoglossal Nerve Stimulation

The Genio[®] proprietary Duty Cycle algorithm



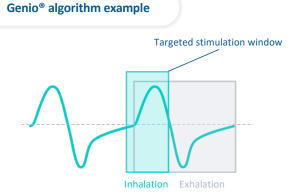
 The patient breathing frequency is used as a reference to adjust the stimulation cycle

2 Ensures a timely and efficient stimulation, through Duty Cycle

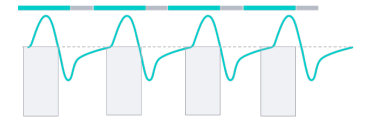
- Genio[®] Duty Cycle algorithm: alternance of ON/OFF stimulation phases in a cyclical pattern
- The cyclical "ON" phase overlaps with the target window (end of expiration + early inspiration) to deliver a timely stimulation
- The "OFF" phase prevents the occurrence of nerve and muscle fatigue

3 Validated by clinical evidence

- Data presented in a poster at the 26th Congress of the European Sleep Research Society in Sep 2022
- Optimal programming allows for a stimulation overlap with inspiration phase as high as 82% of cycles
- The study demonstrated that "customized patient programming strategies to maximize therapeutic effect of bilateral HGNS can be achieved by adjusting stimulation
- parameters without the need for a sensing lead"



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Stimulation "ON"

Targeted stimulation window

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The Genio[®] System – Addressing Unmet Patient and Physician Needs

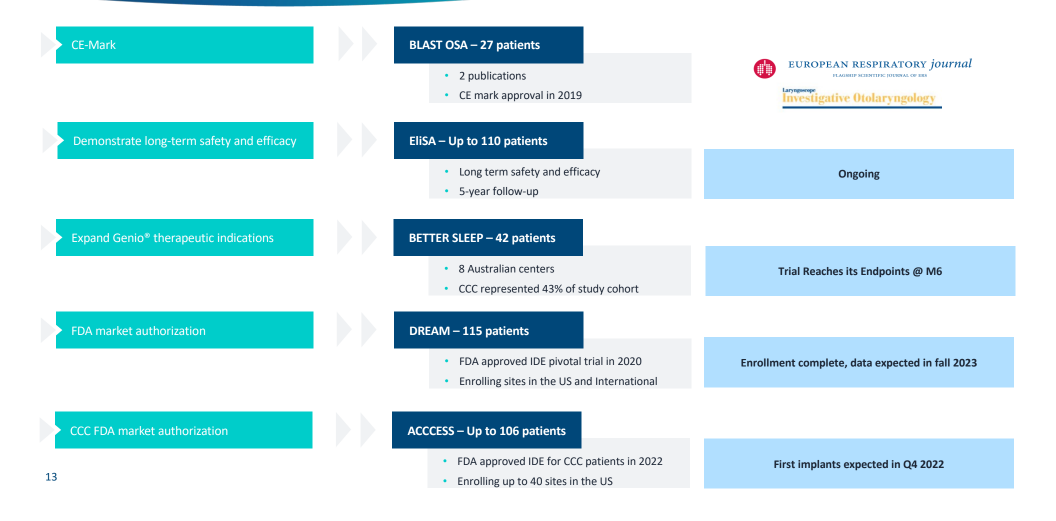


- 60% MRI scans are performed in the thorax/abdominal area¹⁷
- 40% MRI scans are 3T

	Nyxoah	Sleep Aprice Innovation	LivaNova 🐠 Imthera
Implantable parts	1 – Genio [®] IS	3 – Pulse generator + 2 leads	2 – Pulse generator + lead
Number of incisions	1 incision	2 incisions	2 incisions
Lead(s)	0 – No lead	2 – Stimulation and breathing	1 – Stimulation
Tunneling	No tunneling	At least 1 step	1 step
Power source	External	Implanted battery	Implanted battery
Estimated implantation time	1 hour	2.5 hours	2 hours
MR conditional labeling	Full-body 1.5T and 3.0T	1.5T only	No



Clinical Strategy *Building Robust Clinical Evidence*



BETTER SLEEP Study – Demonstrated Efficacy for Both Non-CCC and CCC Patients

Population and key inclusion criteria	22 < Age < 75	Multicenter, prospective, open-label, two group study	With and without complete concentric collapse (CCC)
	$BMI < 32 \text{ kg} / \text{m}^2$	Moderate to Severe OSA – AHI 15-65	Failed, did not tolerate, or refused PAP
	Target: 42 patients		
Safety and Performance			

- Safety: Incidence of device-related serious adverse events (SAEs)
- Efficacy Endpoints
 - Primary: Mean reduction in AHI (4% oxygen desaturation, AHI_{4%}) for the entire cohort
 - Exploratory: Mean reduction in AHI4 for the CCC subgroup
 - Secondary: Mean reduction in oxygen desaturation index scored at 4% desaturation (ODI_{4%}) for the entire cohort



BETTER SLEEP *Responder Rate – Sher Criteria*

- 36 patients completed polysomnography at month 6
- Responder defined according to the Sher criteria²² (\geq 50 % reduction in AHI and AHI less than 20)
- Strong responder rates help de-risk DREAM U.S. IDE study

	AHI _{4%} Responder Rate — Sher Criteria —
CCC & Non-CCC – n=36	64%
CCC – n=15	60%
Non-CCC – n=21	67%

Results subject to final review and validation

15 22. Sher AE, Schechtman KB, Piccirillo JF (1996) The efficacy of surgical modifications of the upper airway in adults with obstructive sleep apnea syndrome. Sleep 19(2):156–177



- >70% reduction in AHI_{4%} in responders in both CCC and non-CCC population
- Presence of "Super Responders" in both CCC and non-CCC cohorts provides additional validation

AHI responders	– AHI _{4%} – Mean % reduction at 6 months
Entire cohort (n=23)	73.4%
CCC cohort (n=9)	71.1%
Non-CCC cohort (n=14)	74.9%

BETTER SLEEP – Data from the entire cohort



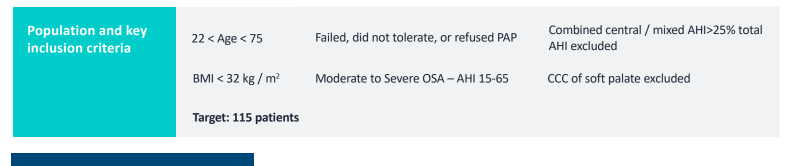
BETTER SLEEP *CCC Market Expansion Opportunities*

Europe	 CE Mark label expansion to include CCC patients in Europe CE-Mark Instructions for use: "The Genio® system is indicated to treat patients suffering from moderate to severe OSA with and without Complete Concentric Collapse (CCC) at the soft palate level" No DISE required to assess presence of CCC at the soft palate level
United States	 Genio[®] received "Breakthrough Device Designation" in the US for CCC patients ACCCESS IDE approved, first patients expected to be implanted in Q4 2022 FDA approval expected in the first half of 2025

17 Source: Lewis R. et al. Bilateral Hypoglossal Nerve Stimulation Improves Moderate to Severe Obstructive Sleep Apnoea in Participants With and Without Complete Concentric Collapse (BETTER SLEEP) – World Sleep Congress 2022 Abstract



DREAM Study – IDE Pivotal Trial



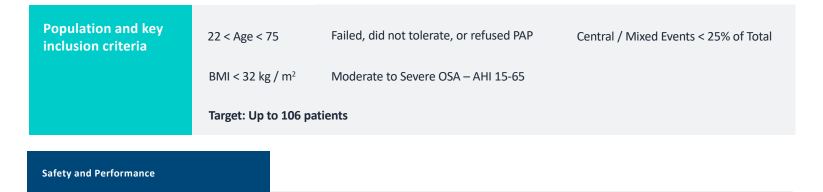
Safety and Performance

- Safety: Incidence of device-related serious adverse events at 12 months
- Co-primary effectiveness endpoints: Percentage of responders at 12 months based on Apnea Hypopnea Index (AHI4%) using the Sher criteria and percentage of responders at 12 months based on ODI4% (25% reduction of ODI between baseline and 12-month visit)





ACCCESS Study – IDE Trial to Gain US Approval for CCC Patients



- Safety: Incidence of device-related serious adverse events through 12 months
- Co-primary effectiveness endpoints: Percentage of responders at 12 months based on Apnea Hypopnea Index (AHI4%) using the Sher criteria and percentage of responders at 12 months based on ODI4% (25% reduction of ODI4 between baseline and 12-month visit)

	• FDA approved IDE pivotal trial in July 2022 – up to 40 participating sites
Status	Mix of large US academic centers & private practices, all HGNS experienced
Status	First patients expected to be implanted in Q4 2022
	FDA approval expected in the first half of 2025



Commercial Leadership Through Focused Execution

Go Deep versus Go Wide

Recognized by medical community for expertise and high level of care Center of Excellence Clinical & Administrative leadership

Stakeholders

- ENT implant surgeonQuality implant
- Sleep physician
 - Patient phenotyping
 - Patient outcomes
- Referral network
 - GPs, cardiologists

EU Market Access

Nyxoah – EU reimbursed countries

Nyxoah – EU reimbursement opportunities

Germany – 2021: Dedicated DRG
Switzerland – 2021: Dedicated DRG
Spain – 2021: Funding through hospital

budget

Belgium

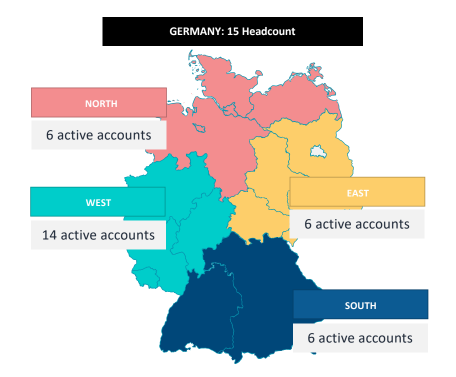
Nordic Countries

Italy

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• Finland -- 2022

Germany - Proof of concept



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Acurable Home Sleep Test German Partnership AcuPebble Aligns with Nyxoah's Patient-First Mission Statement

A Next-Generation Home Sleep Test

- CE Mark and FDA clearance
- High specificity and sensitivity for AHI and ODI, which is validated by a randomized study published in the BMJ Open

Patient-Centric Design

- *Highly accurate* >93% and >91% specificity and sensitivity for AHI and ODI compared to gold-standard polygraphy
- Easy to use patients download APP and follow simple instructions
- Comfortable small system worn at the base of neck at night
- Reusable can be used for routine monitoring, including on multiple consecutive nights
- Novel form factor aligns with trend towards wearable technology

With AcuPebble and Genio, Nyxoah can now offer patients and clinicians the most cutting edge OSA home sleep diagnosis and treatment solutions

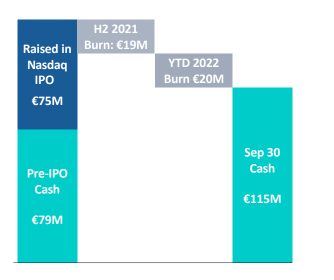


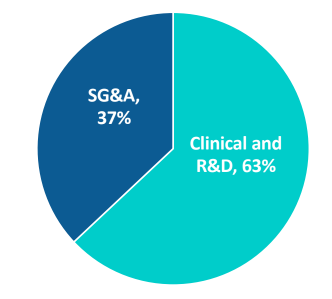
High Quality and Diverse International Shareholder Base



Strong Cash Position Supports Clinical and Commercial Priorities in 2022 and Beyond

Strong Cash Position - €115M





- R&D/Clinical to represent >60% of 2022 expenses driven by DREAM enrollment completion, EliSA study progress, and ACCCESS study commencement
- SG&A drivers include increased commercial efforts in Germany to achieve market leadership and scale-up of corporate functions



Thank you

Q&A

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