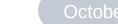


Unlocking the potential of advanced therapies for hearing loss



October 2024

SENSORION

Our vision is to help people with inner ear hearing disorders to live life with unlimited connections



DISCLAIMER

- This document has been prepared by Sensorion (the "Company") and is provided for information purposes only. This document does not purport to contain comprehensive or complete information about the Company and is qualified in its entirety by the business, financial and other information that the Company is required to publish in accordance with the rules, regulations and practices applicable to companies listed on Euronext Paris. No reliance may be placed for any purposes whatsoever on the information or opinions contained in this document or on its accuracy or completeness.
- This presentation does not constitute an offer to sell, a solicitation of, or an invitation to subscribe for or to buy, securities of Sensorion in any jurisdiction.
- The information and opinions contained in this document are provided as of the date of this document only and may be updated, supplemented, revised, verified or amended, and thus such information may be subject to significant changes. The Company is not under any obligation to update the information or opinions contained herein which are subject to change without prior notice.
- The information contained in this document has not been subject to independent verification. No representation, warranty or undertaking, express or implied, is made as to the accuracy, completeness or appropriateness of the information and opinions contained in this document. The Company, its subsidiaries, its advisors and representatives accept no responsibility for and shall not, under any circumstance, be held liable for any loss or damage that may arise from the use of this document or the information or opinions contained herein.
- This document contains information on the Company's markets and competitive position, and more specifically, on the size of its markets. This information has been drawn from various sources or from the Company's own estimates which may not be accurate and thus no reliance should be placed on such information.
- This document contains certain forward-looking statements. These statements are not guarantees of the Company's future performance. These forward-looking statements relate to the Company's future prospects, developments and marketing strategy and are based on analyses of earnings forecasts and estimates of amounts not yet determinable. Forward-looking statements are subject to a variety of risks and uncertainties as they relate to future events and are dependent on circumstances that may or may not materialize in the future. Forward-looking statements cannot, under any circumstance, be construed as a guarantee of the Company's future performance and the Company's actual financial position, results and cash flow, as well as the trends in the sector in which the Company operates, may differ materially from those proposed or reflected in the forward-looking statements contained in this document. Important factors that could cause actual results to differ materially from the results anticipated in the forward-looking statements include those discussed or identified in the "Risk Factors" section of our 2023 Annual Report published on March 14, 2024, and available on our website (www.sensorion.com). Even if the Company's financial position, results, cash-flows and developments in the sector in which the Company operates were to conform to the forward-looking statements contained in this document, such results or developments cannot be construed as a reliable indication of the Company's future results or developments. The Company does not undertake any obligation to update or to confirm projections or estimates made by analysts or to make public any correction to any prospective information in order to reflect an event or circumstance that may occur after the date of this document.
- Certain figures and numbers appearing in this document have been rounded. Consequently, the total amounts and percentages appearing in the tables may not necessarily equal the sum of the individually rounded figures, amounts or percentages.
- All persons accessing this document must agree to the restrictions and limitations set out above.

Sensorion: Establishing Global Leadership In Hearing Loss With Strong And Diversified Pipeline



Untapped opportunity with 1,5 bn people affected by hearing loss (HL); 0,5 bn suffer from disabling HL Multiple causes: genetic, environmental, idiopathic



Modality agnostic approach leveraging unparalleled understanding of the inner ear and world-leading, differentiated and exclusive partnerships (Institut Pasteur, Necker Hospital, Cochlear Ltd, Sonova)



Two gene therapies (GT): SENS-501 (OTOF-GT – First patient injected); **GJB2-GT** (preclinical IND/CTA enabling activities) Prospective Natural History Studies ongoing



Small molecule, SENS-401, for the treatment and prevention of HL caused by Cochlear Implantation CI (POC Ph2a completed), Cisplatin-Induced Ototoxicity CIO (POC Ph2a ongoing), and Sudden Sensorineural HL (Ph2b completed)

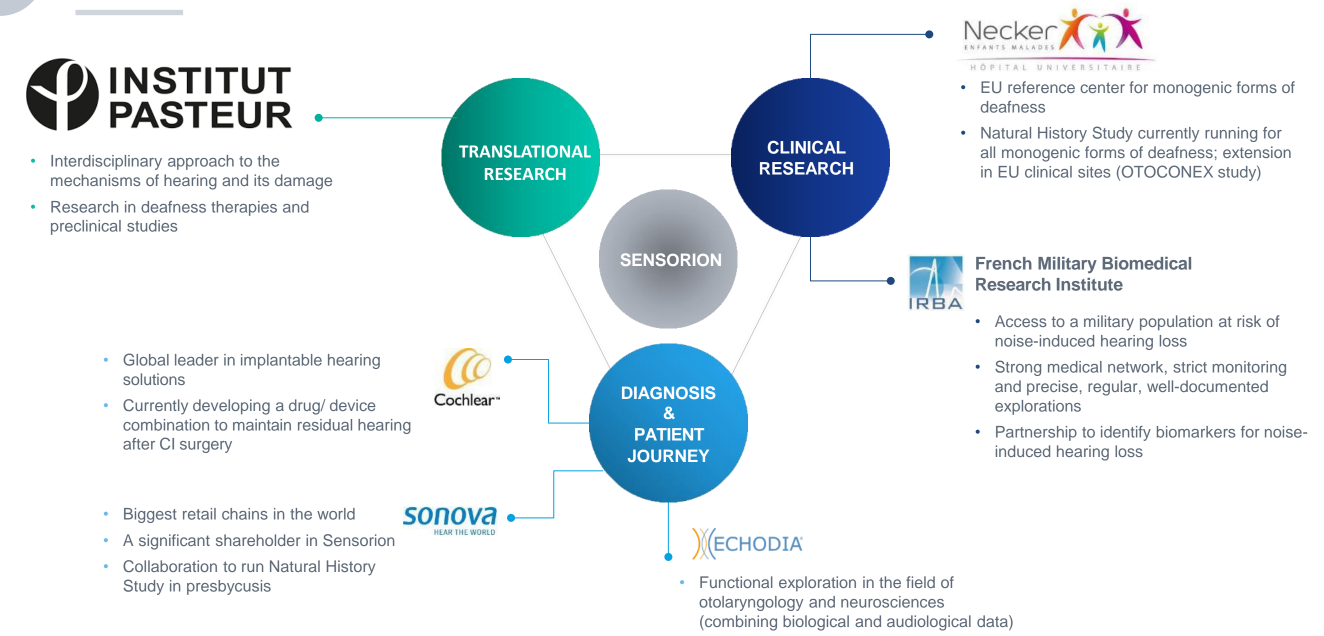


Multiple **upcoming milestones** across the GT and small molecule pipeline, including completion of patient recruitment of the first cohort in the Ph1/2 GT trial of SENS-501 in H2 2024; the completion of patient recruitment in Ph2a SENS-401 CIO in H1 2025; CTA submission for GJB2-GT in H2 2025



Experienced and visionary management team Strong shareholder base backed by leading blue-chip life sciences investors; €100m raised since Aug 2023 giving cash runway until the end of 2025

Together With Best-In-Class Partners We Can Transform The Current Standard of Care



Sensorion's Portfolio Of Advanced Hearing Loss Therapies

	Product	Indication	Discovery	In-vivo POC	Preclinical	Phase 1	Phase 2	Phase 3	Upcoming Milestones (estimated)
	SENS-501*	Otoferlin Deficiency				Phase	1/2		1 st cohort completed by end of 2024
RESTORE	GJB2-GT*	Adult Onset (presbycusis)							Preclinical CTA/IND enabling activities
REST	GJB2-GT*	Pediatric Progressive					, 		Preclinical CTA/IND enabling activities
	GJB2-GT*	Congenital Onset							Preclinical CTA/IND enabling activities
PREVENT	SENS-401	Hearing Preservation after CI						Cochlear"	Clinical Study Report completed
PREV	SENS-401	Cisplatin-Induced Ototoxicity							Recruitment Completed H1 2025
REAT	SENS-401	SSNHL					1		Exploring Partnering Opportunities

3SBio has a right of first refusal with respect to licensing in Greater China of SENS-401 (except in combination with cochlear implants) and SENS-501 OTOF-GT *Option to obtain a licence from the Institut Pasteur (pre-defined financial terms and other terms to be negotiated)

We Have Established Internal Capabilities To Ensure Successful Execution



GENE THERAPY PROGRAMS



Sensorion's Gene Therapy Programs Target Rare Auditory Diseases

FIRST PROGRAMS RESULTING FROM THE INSTITUT PASTEUR COLLABORATION

OTOFERLIN DEFICIENCY	GJB2-RELATED HEARING LOSS
 Pediatric patients with mutations in OTOF suffer from severe to profound sensorineural prelingual non-syndromic hearing loss Otoferlin deficiency could be responsible for up to 8% of all cases of congenital hearing loss Prevalence ~20,000 in the USA + EU Incidence ~1,100 per year in USA + EU EU and US Orphan Disease Designation , US Rare Pediatric Disease Designation Pediatric Investigational Plan Agreed in EU 	 We have identified three forms of hearing loss associated with <i>GJB2</i> gene mutations: Early onset of severe presbycusis (adult population) Childhood onset (pediatric population) Congenital onset (pediatric population) ~100,000 patients between 30- and 69-years old thought to be affected by a monogenic form of presbycusis due to <i>GJB2</i> mutations Prevalence of congenital and childhood onset forms are estimated to be around 200,000 patients as around 50% of autosomal recessive nor syndromic hearing loss cases are thought to be from <i>GJB2</i> mutations
Sources: Akil et al. 2019 (<u>link</u>), Orphanet (<u>link</u>), NIH (<u>link</u>), company estimates base & Co 2019 report, Institut Pasteur	

DELAYED DIAGNOSIS – NOT SUSPECTED AT FIRST SIGHT

GENE THERAPY HAS A LIFE-CHANGING POTENTIAL FOR THESE AUDITORY DISEASES

SENS-501 Aiming To Develop Best-In Class And First-In Class Gene Therapy

CRITERIA	SENSORION
AAV capsid selected for high-level of target cells specificity	
GT product showing high level of target cells transduction	
Limited off-target tissue biodistribution	
Surgical approach developed and mastered by ENTs surgeons	
Natural History Study preparing execution of the clinical trial	
Regular engagement with regulatory agencies	



SENS-501 development has been supported by Audinnove consortium (Institut Pasteur, Hôpital Necker- enfants malades, Fondation pour l'Audition & Sensorion) which is financed by the French State, via the National Research Agency through the "Investing for the future" program (ref: ANR-18-RHUS-0007)

SENS-501 Gene Therapy Pediatric Indications Have Blockbuster Sales Potential

SENS-501 (OTOF-GT) is the Perfect Pilot Program

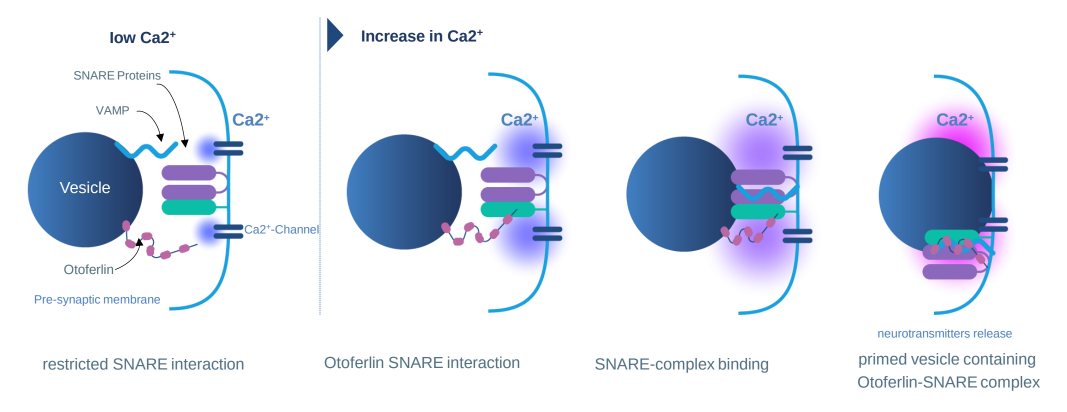
- Well understood biology and pathology of the otoferlin deficiency
- Full functionality of the remaining chain
- High specificity for the inner hair cells (IHCs), no off-target effect expected

- SENS-501 will be the pilot program demonstrating that GT is a relevant medical approach for the inner ear
- SENS-501 will establish understanding of GT in the inner ear by the Regulators and the Payers for future GT programs
- Medical plausibility and target population have been confirmed through :
 - ✓ ODD in the US and EU, RPDD with eligibility for voucher in the US
 - ✓ PIP Agreed in EU
 - ✓ Clinical Trial Application approved in France (FPI injected)



Sources: Sensorion, AT Kearney market research

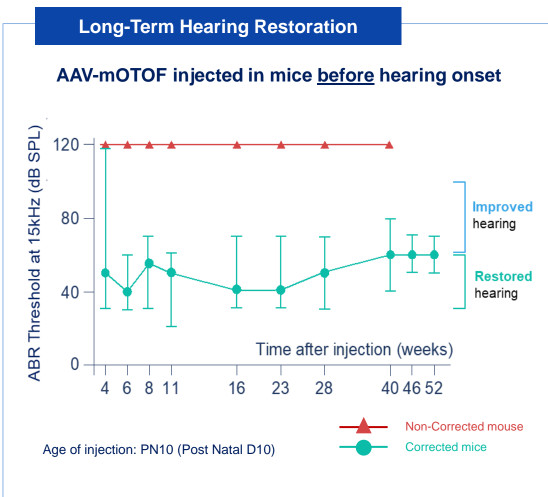
SENS-501 OTOF Gene Encodes Otoferlin, A Key Ca2⁺ Sensor Protein



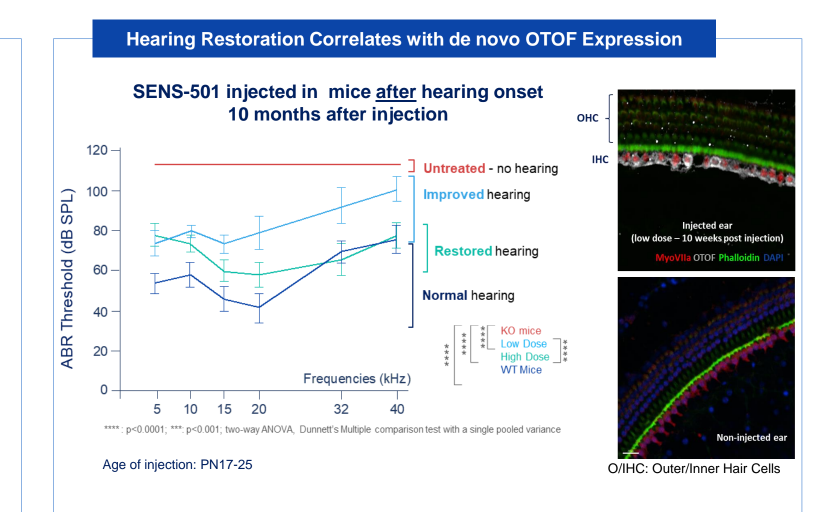
Model illustrating calcium regulation of otoferlin/SNARE interaction in the hair cell – Adapted from Ramakrishnan et al. 2014

OTOF is the gene coding for the otoferlin protein, a Ca2⁺ sensor for vesicle fusion and vesicle pool replenishment at auditory hair cell ribbon synapses

SENS-501 SENS-501 Leads to Long-Term Hearing Recovery in a Translational Model of Otoferlin Deficiency



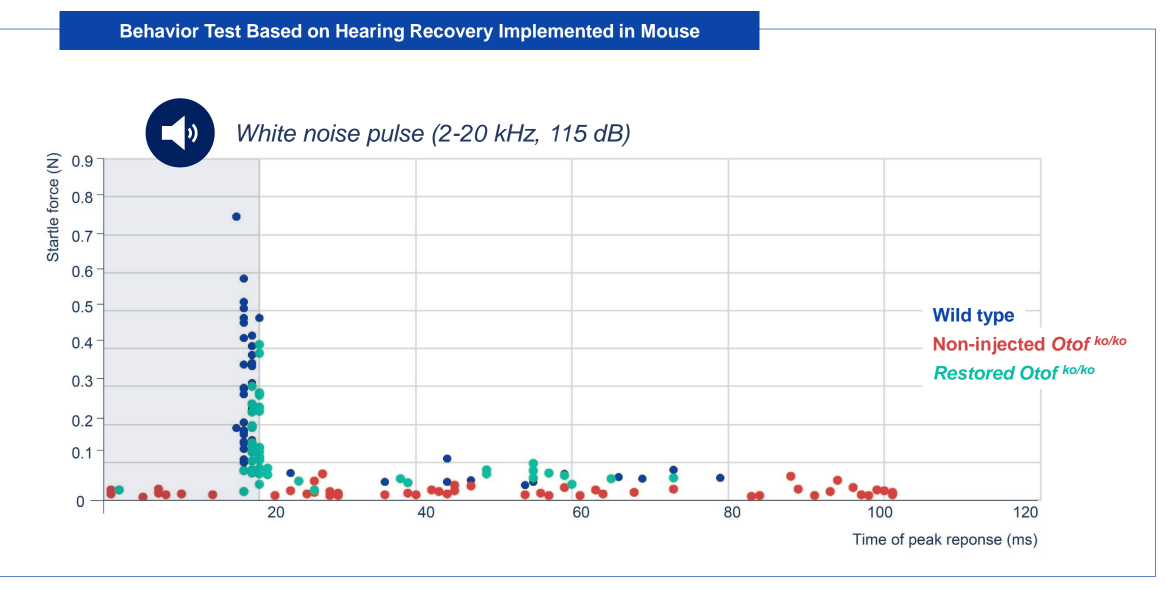
• **Durable hearing restoration** in Otof-/- mice by dual AAV-OTOF directly delivered to the inner ear up to one year post-injection



- Both doses of SENS-501 demonstrated efficacy in improving hearing in KO mice
- SENS-501 leads to otoferlin expression in Inner Hair Cells

Olivier et al. ASGCT 2023 link

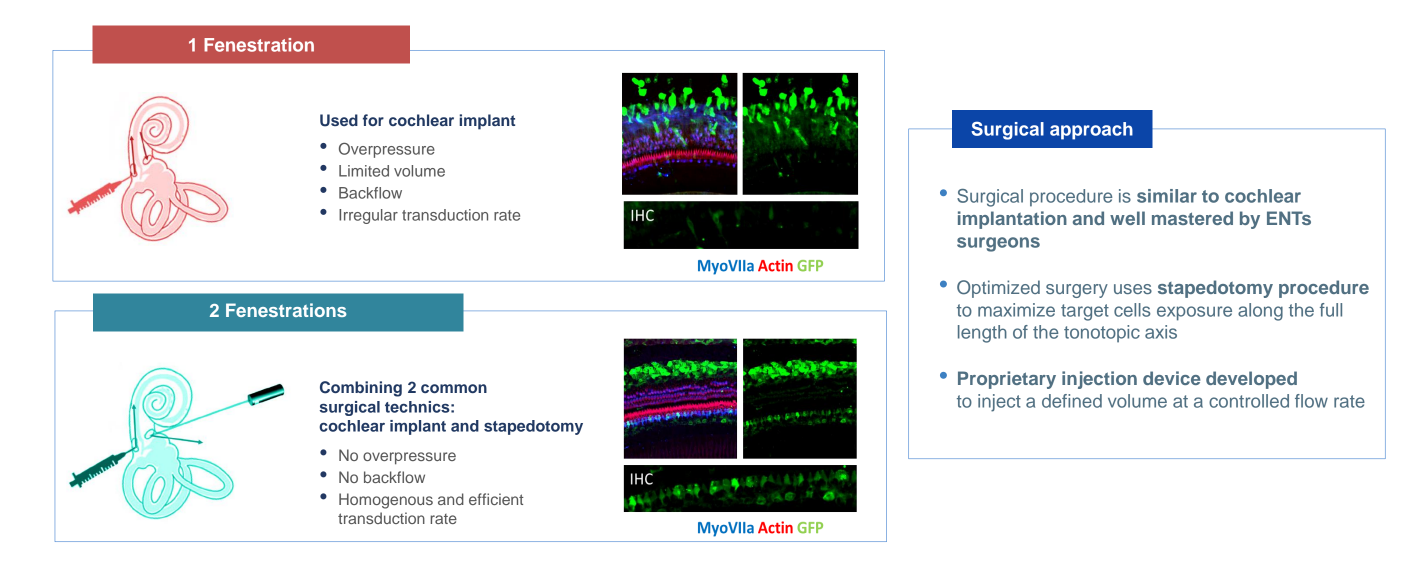
SENS-501 SENS-501 Leads to Restoration of Efficient Sound Processing in Behavioural Test



Olivier et al. ASGCT 2023 link

SENS-501 Dedicated Surgical Approach for Gene Therapy

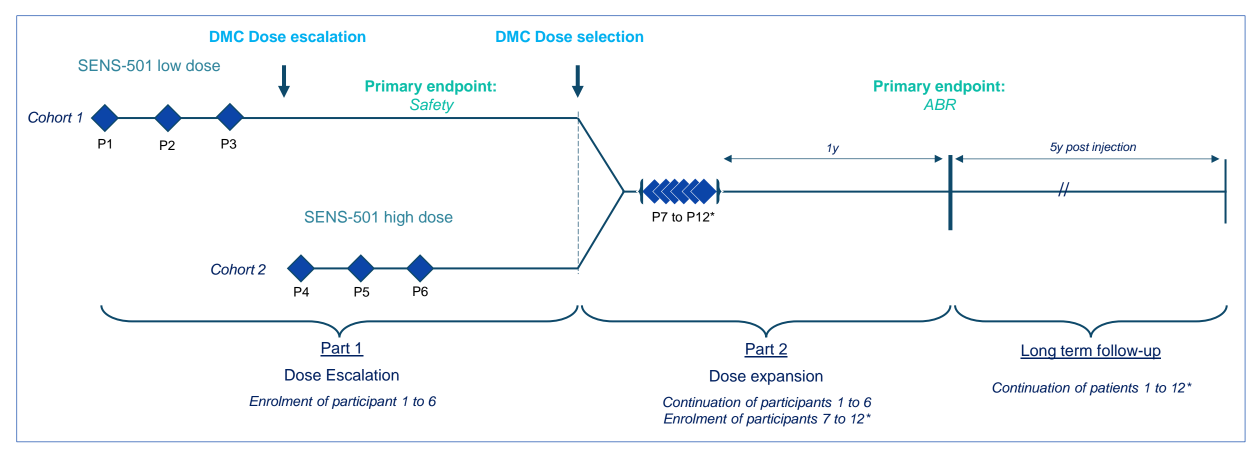
Non-Human Primates injected through the round window membrane (RWI) with or without stapedotomy (stap)



SENS-501 Phase 1/2 Audiogene Study (SENS-501)



Audiogene, a Phase 1/2 clinical trial in children aged 6 to 31 months to assess safety, tolerability, and efficacy of SENS-501 following unilateral injection into the cochlea



- Pediatric patients, aged 6 to 31 months at the time of the injection (to maximize chances of acquiring speech and language)
- Single intra-cochlear unilateral injection
- **Dose escalation** (Primary endpoint: safety and tolerability)
- **Dose expansion** (Primary endpoint: efficacy (ABR, Auditory Brainstem Response)

Px = participant number x y = year *Further participants may be recruited if required, who will be assessed in the same way as P7 to P12.

SENS-501 Audiogene (SENS-501) Study Status

Audiogene

First Patient Injected

- First Patient Injected in Q3 2024
- Initial safety data reported at 4 weeks
 - No dose-limiting toxicities, no Serious Adverse Events
 - Vestibular function remained intact and unchanged from baseline
 - Otoacoustic Emissions (OAEs) remain present
- Surgical administration procedure was uneventful
- Medical condition of the child (upper respiratory infection unrelated to SENS-501) at the time of assessment did not allow to get reliable hearing sensitivity. ABR at Month 1 was postponed
 - From early observations, the clinical team and the patient's mother noted a change in the child's behaviour and vocalisations

Study Update

- Patient recruitment going as planned:
 - 1st cohort screened; completion expected by year-end 2024
 - Initial efficacy data to be reported by year-end 2024
 - 2nd cohort of patients to be recruited in H1 2025
- Ongoing Natural History Study Otoconex supports eligible patients' identification

SENS-501 SENS-501 (OTOF) Gene Therapy Program Status

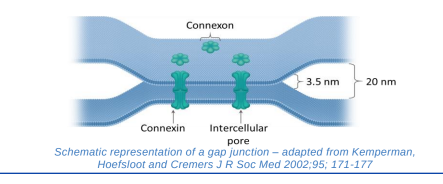


GJB2-GT

Connexin 26: A Gap-junction Protein Encoded By *GJB2* Gene and Responsible For Tissue Homeostasis

Mutations in the GJB2 Lead to Deafness

- *GJB2* is the gene encoding for the Connexin 26 protein; one of 20 known connexins
- Cx26 and Cx30 proteins are the dominating connexins in the cochlea; heteromeric or heterotypic hexamers forming Gap Junctions
- Gap Junctions are key for the intercellular exchange of molecules (miRNA, glucose, ions, etc.) hence responsible for tissue homeostasis
- More than 100 recessive mutations origin Cx26 truncation / deletion leading to non-syndromic hearing loss and deafness, most are addressable via gene replacement
- Severity of hearing loss correlates with degree of loss of GJB2 function



GJB2 Expression in the Cochlea Muse cochlea Organ of Corti top view Cochlea cross section Organ of Corti top view Image: Cochlea cross section Image: Cochlea cross section<

Phalloidin Cx26 DAPI

- Supporting cells of the organ of Corti
- Fibrocytes of the spiral limbus and the lateral wall
- Intermediate and basal cells of the stria vascularis
- Not expressed in hair cells

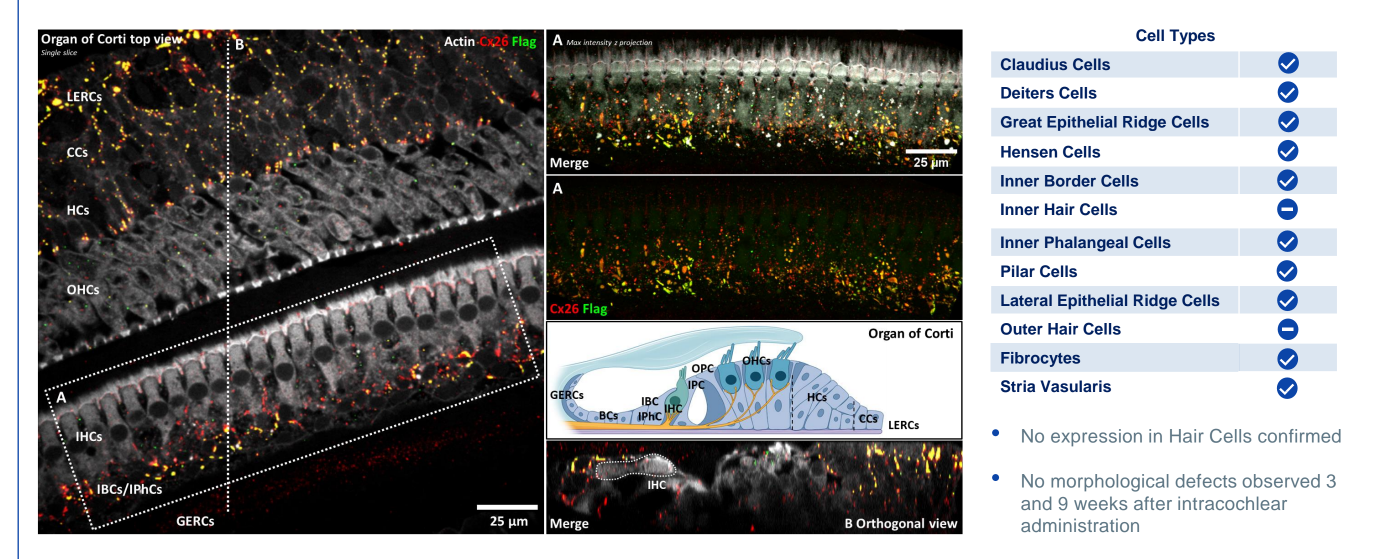
GJB2-GT Lead Candidate Was Selected to Answer Specific Development Criteria

CRITERIA	LEAD CANDIDATE
Natural and synthetic AAV capsid libraries screening for broad coverage of target cells	
Expression cassette design for high-level of target cells transduction, correct cellular localization, active gap-junctions	
Avoiding off-target expression (i.e. hair cells): promoter and regulatory sequences design	
Limited off-target tissue biodistribution	$\mathbf{\sim}$
Surgical approach developed and mastered by ENT surgeons	

Our Lead Candidate Was Designed to Ensure Broad Coverage of Relevant Cochlear Cells While Detargeting Hair Cells

GJB2-GT Lead Candidate Can Deliver Cx26 in the Appropriate Target Cells

Correct Delivery of Cx26 Using Lead Candidate Flag in Non-Human Primate Cochlea



Lead Candidate Demonstrates Adequate Safety and Biodistribution **GJB2-GT Profile - Including Long-Term Local Tolerability in Mice and NHP**

Acute toxicity in WT Mice - High dose IV injection

- Study performed in preparation of upcoming GLP-toxicity in mice after IV injection
- GT-GJB2 does not interfere with normal growth and don't elicit elevated transaminase levels 4 and 8 weeks after injection
- Behavioral evaluation (Functional Observation Battery, exploratory behavior (videotracking) 3 and 7 weeks after injection: no findings

6 months after injection No impact on ABR up to 6 months following Lead Candidate injection Normal histology maintained, WT mice - Lead Candidate FLAG BR Threshol (dB SPL) transgene expression persistence WT mice - Lead Candidate Hair cells detargeted WT mice - Not injected Clinical pathology: no findings WT mice - Vehicle

3-Month Exploratory Toxicity and Biodistribution in Non-Human Primate – Intracochlear injection

- Lead Candidate is well tolerated and did not induce any macroscopic/organ weight changes or local/systemic microscopic findings
- Normal cochlear histology

Frequency (kHz

- No lab and clinical findings
- Biodistribution: the vast majority of the vector remains in injected ears, no dissemination observed in gonads, main organs, dorsal root ganglion (DRG)

6-Month Exploratory Safety and Transgene Expression in WT Mice – Intracochlear injection

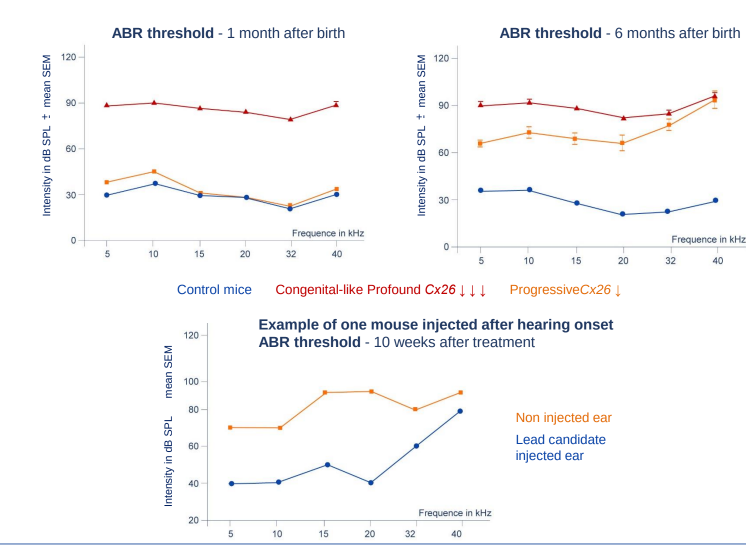


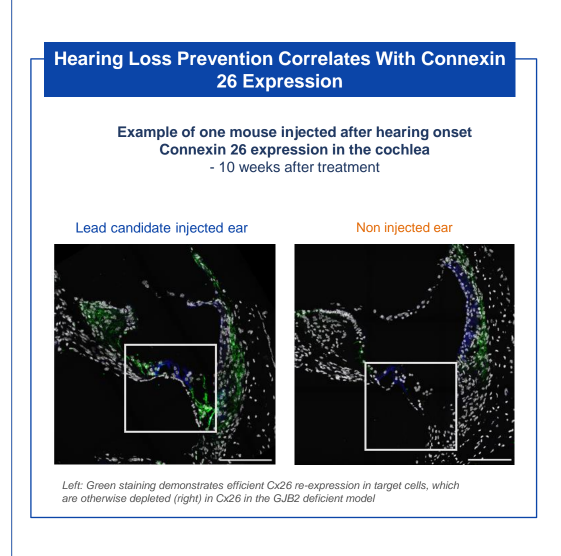
intracochlear injection in mice

GJB2-GT Lead Candidate Prevents Hearing Loss in Relevant Mouse Model



Conditional knock-out mouse model leading to 2 phenotypes





GJB2-GT GJB2 Gene Therapy Program Status

Ongoing European Natural History Study OTOCONEX

Ongoing Natural History Study in Collaboration with Sonova

Preclinical IND/CTA Enabling Studies Update on Additional PoC Efficacy and Safety Data Oct 2024 (ESGCT)

Clinical Trial Applications H2 2025

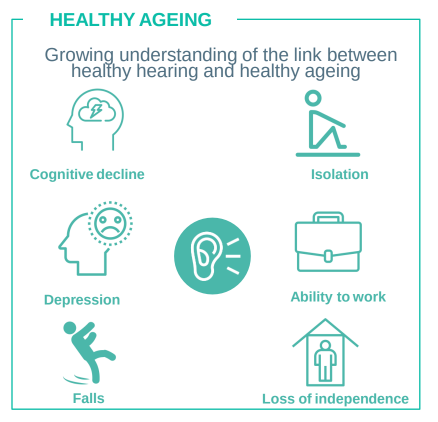
SENS-401 PROGRAMS

Multiple Indications To Treat And Prevent Hearing Loss

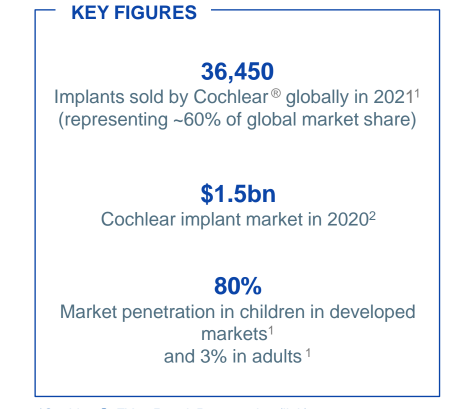


SENS-401 To Preserve Residual Hearing After Cochlear Implantation

COMBINATION OF COCHLEAR IMPLANT WITH SENS-401 TO PREVENT CELL-DEATH POST COCHLEAR IMPLANT PROCEDURE



Source: Cochlear® 2018 investor day (link)

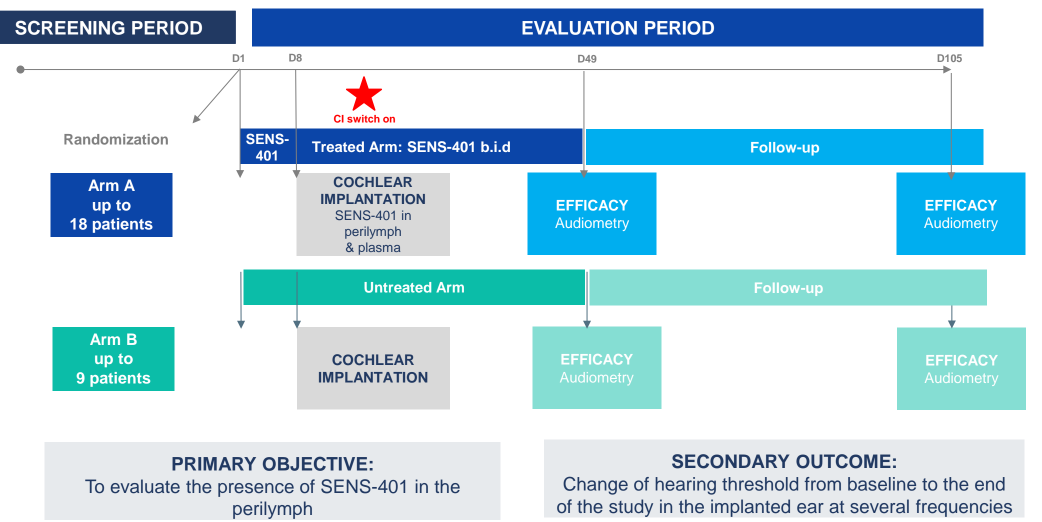


¹Cochlear[®] FY21 Result Presentation (<u>link</u>) ²Market estimates (<u>link</u>)

SENS-401 CI Study Design Study completed

A Phase 2a, Multicenter, Randomized, Controlled, Open-label Study





Primary Endpoint of the Phase 2a Clinical Study for Residual Hearing Preservation Has Been Met

Perilymph Conce	entrations Data	
		Treated with SENS-401 (n=16) n (%)
SI	ENS-401 levels ≤ LLOQ	0
SI	ENS-401 levels > LLOQ	14*(100)
*LL	nong the 16 participants who underwent surgery, 15 have a pe OQ define by a specific method developed for SENS-401 The sampling times for SENS-401 levels in the perilymph were	

- Presence of SENS-401 in the perilymph is confirmed in 100% of the patients sampled following cochlear implantation
- These results confirm that SENS-401 administered orally crosses the labyrinth barrier

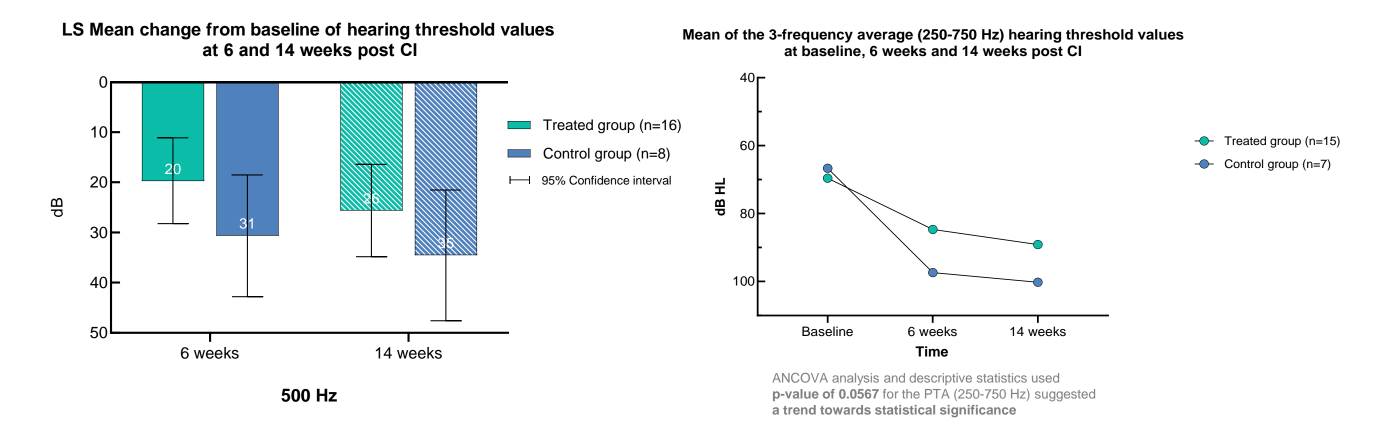
Residual Low Frequency Hearing Benefits for Cochlear Implant Users



Postoperative hearing preservation defined as: unaided air-conduction **thresholds < 85 dB HL** at 125, 250, and 500 Hz

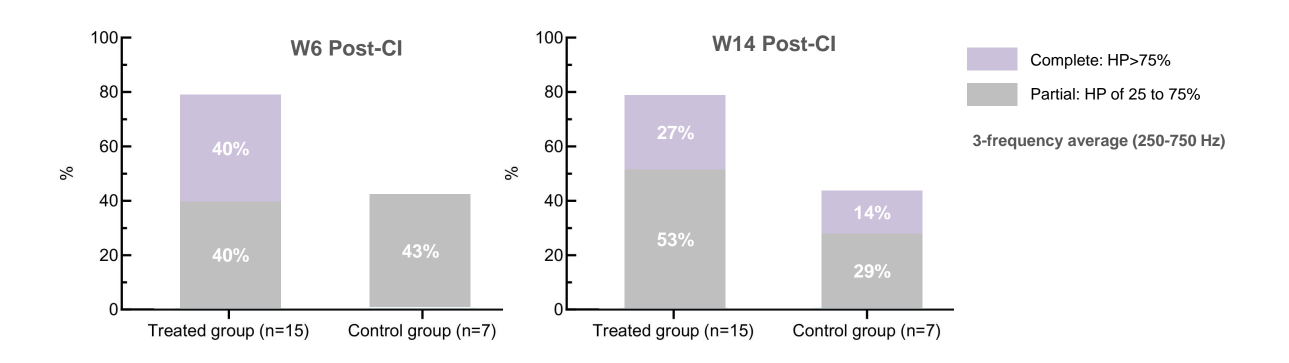
*Jensen et al., Hearing Preservation After Cochlear Implantation, 2021

SENS-401 Provides Hearing Protection 6 & 14 Weeks Post-Cochlear Implantation



- Residual hearing loss is lower in patients treated with SENS-401 compared to control group 6 weeks after cochlear implantation
- This preservation effect is maintained 8 weeks after SENS-401 discontinuation (14 weeks post-CI)

SENS-401 Provides Residual Hearing Preservation* 6 & 14 Weeks Post-Cochlear Implantation



- Patients in the SENS-401 treated group are twice as likely to show complete or partial hearing preservation compared to control group after 7 weeks of continuous treatment
- Only SENS-401 treated group show a complete hearing preservation with 40% of treated patients compared to 0% in the control group at 6 weeks post-CI
- These results are maintained 8 weeks after SENS-401 discontinuation (14 weeks post-CI)

*Skarzynski H, van de Heyning P, Agrawal S, Arauz SL, Atlas M, Baumgartner W, et al. Towards a consensus on a hearing preservation classification system. Acta Otolaryngol Suppl. 2013(564):3-13.

SENS-401 CI Final Results - Conclusion



SENS-401 can cross the labyrinthine barrier to target cochlear hair cells in all patients sampled, confirming primary endpoint is met. SENS-401, present in the perilymph fluid, reaches concentrations that are pharmacologically active.



A **complete hearing preservation is** exclusively observed in 40% of patients treated with SENS-401 at 6 weeks post cochlear implantation



Eight weeks after discontinuation of SENS-401, the hearing protective effect is maintained



Residual hearing loss is reduced in the SENS-401 treated group compared to the untreated group at 6 weeks post-cochlear implantation



SENS-401 taken for 8 weeks confirms it has a good safety profile



SENS- 401 has the potential to modify the outcome of CI while preserving residual hearing by improving speech perception in quiet and noise, music perception, spatial localization and maintaining more natural sound quality



These results support the SSNHL phase 2 data and further development of SENS-401

Cisplatin Administration For Chemotherapies Damages The Inner Ear And Leads To Hearing Loss, Tinnitus And Dizziness

WHAT IS CIO?

Hearing loss caused by cisplatin administration as chemotherapeutic treatment. Risk factors include young age as well as individual and cumulative cisplatin doses.

CIO leads to permanent inner ear problems in 50-60% of adult cases and in 90% of pediatric cases.

These complications significantly impact patients' quality of life due to:

- Hearing loss, tinnitus and dizziness impacting daily life activities
- Problems in language acquisition and learning for pediatric patients
- Difficulties in communicating, social isolation, cognitive decline

Potential treatments must not interfere with cisplatin efficacy.

Incidence of cisplatin treated patients: 500,000 patients in 2025 in G7 countries¹

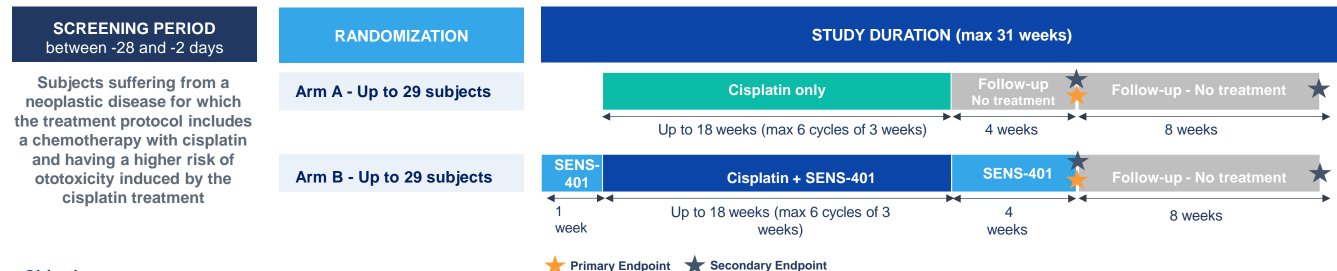


¹ Company/ estimates based on publicly available data (in the US, Japan, Germany, France, the UK, Italy and Spain)

CIO

SENS-401 Phase 2a Proof-Of-Concept Study NOTOXIS Positive Preliminary Safety Data

A Phase 2a, Multicenter, Randomized, Controlled, Open-label Study to Evaluate the Efficacy of SENS-401 to Prevent the Ototoxicity Induced by Cisplatin in Adult Subjects with a Neoplastic Disease



Objectives:

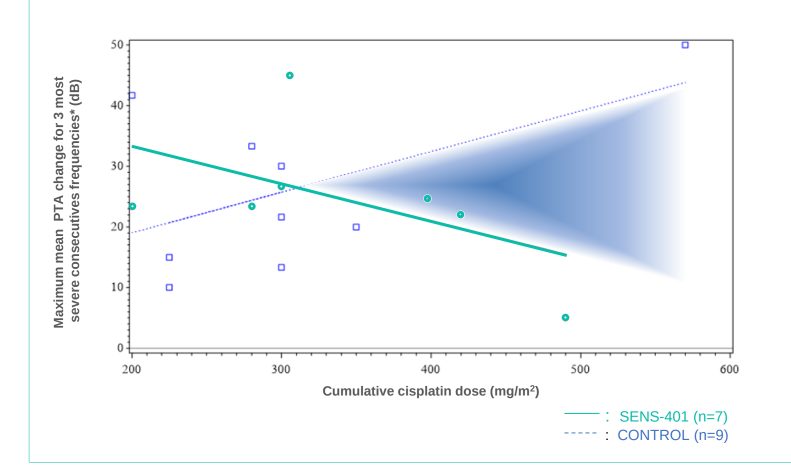
Efficacy

- Rate of ototoxicity
- High Frequency PTA
- Speech in Noise and quiet
- THI questionnaire

Safety

• AEs & SAEs incidence

Preliminary Results Show Patients with High Exposure to Cisplatin May Benefit the Most from SENS-401's Otoprotective Effects



Groups	n	Variables	Mean	SD	
Control	9	Cisplatin dose	305	110.0	
		PTA change	26	13.6	
SENS-401	7	Cisplatin dose	342	98.7	
		PTA change	24	11.7	

- Hearing loss is similar between SENS-401 and control group
- SENS-401 subjects were exposed to significantly more cisplatin than control

- As the cumulative dose of cisplatin increases, severity of ototoxicity observed in the control group escalates r=0.42
- Benefit of SENS-401 increases with higher cisplatin doses
- SENS-401 treatment group outperforms the control group at cisplatin doses > 300 mg/m²

CIO

Key Takeaways from Preliminary Study Data



CIO

Cumulative dose of cisplatin is a key factor of ototoxicity severity



SENS-401 has a **favorable safety profile** when administered continuously for up to **23 weeks** in adult patients undergoing cisplatin-based chemotherapy



Recruitment is progressing well



Based on preliminary data, no significant difference observed on ototoxicity measured by PTA change or CTCAE grading, however SENS-401 treated group received higher cumulative dose of cisplatin compared to control



Patients with higher exposure to cisplatin may benefit the most from SENS-401's otoprotective effect



The preliminary results suggest a trend toward an otoprotective effect of SENS-401 beyond a cisplatin dose of 300 mg/m2

SENS-401 Programs Status

SENS-401 with cochlear implants – Full Data Readout Sept 20, 2024



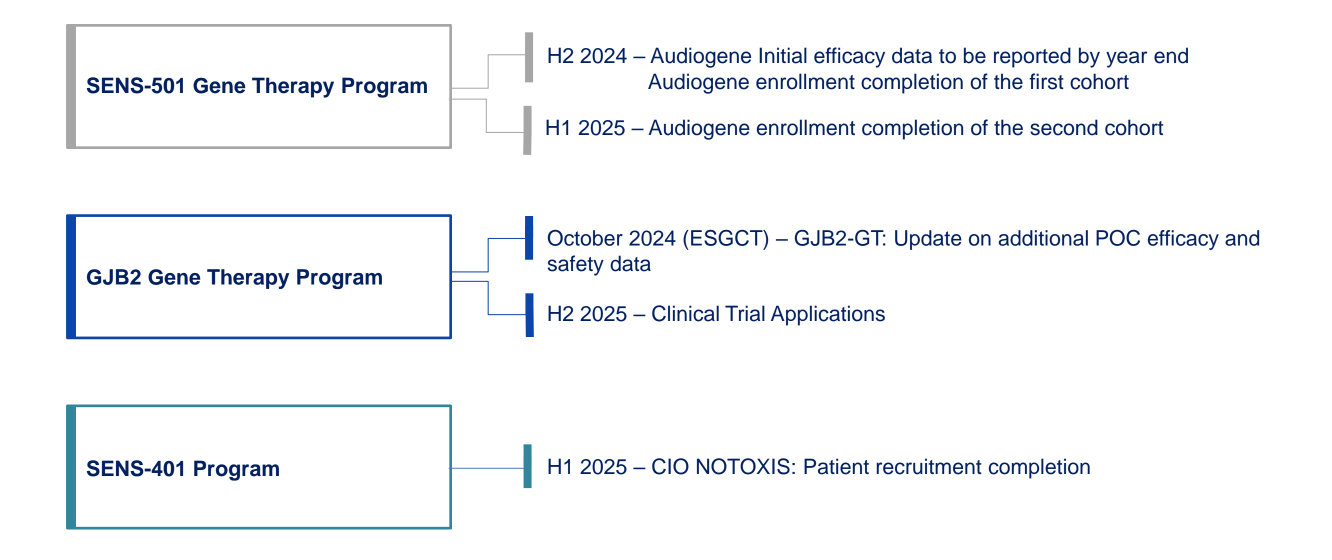
SENS-401 CIO NOTOXIS - Preliminary Results S2 2023

SENS-401 CIO NOTOXIS
- Preliminary Safety and Efficacy Data
Sep 20, 2024

SENS-401 with cochlear implants – Final Results

SENS-401 NOTOXIS – Completion of Enrollment H1 2025 \checkmark

Sensorion Newsflow [Estimated Timelines]



THANK YOU

E: contact@sensorion-pharma.com



Hearing Loss



Access And Clarity Are Mandatory For Optimal Outcomes

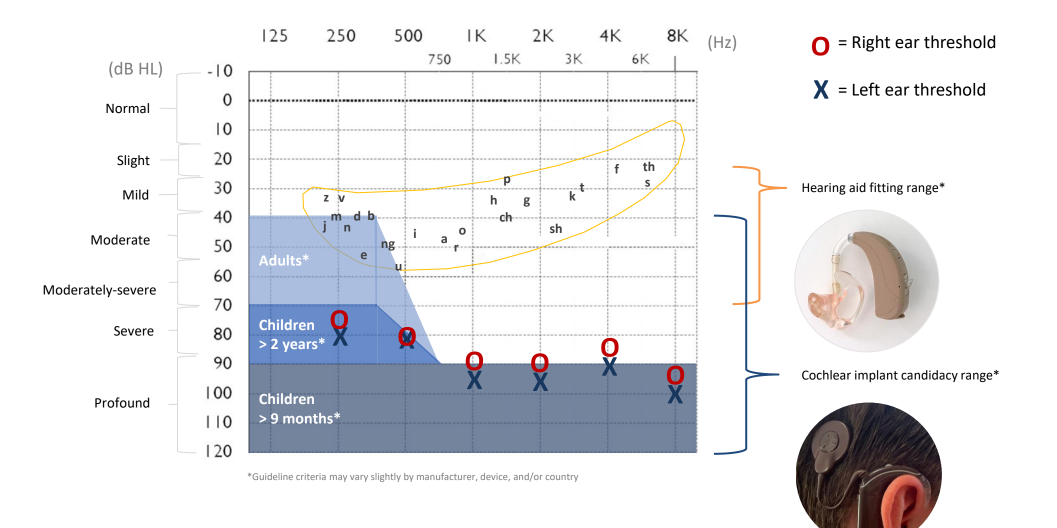
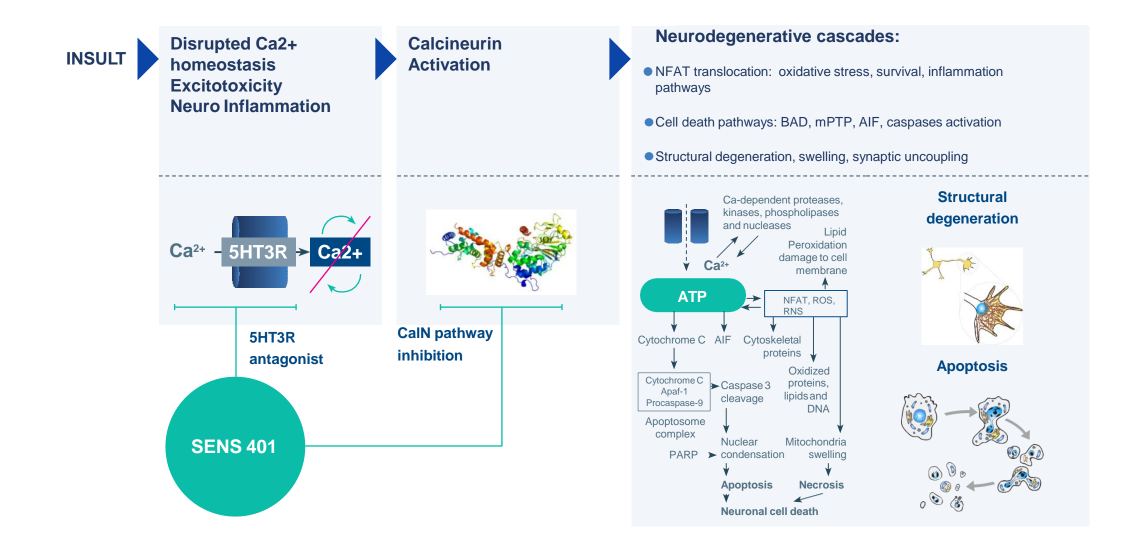


Image of hearing aid: https://commons.wikimedia.org/wiki/File:Unitron_Ziel_photo_2.jpg Image of cochlear implant sound processor on ear: https://commons.wikimedia.org/wiki/File:Cochlear_Nucleus%C2%AE_7_Sound_Processor.jpg

SENS-401



SENS-401 Mechanism Of Action



SSNHL



SSNHL

Sudden Sensorineural Hearing Loss (SSNHL) is a Severe Disease Affecting more than 200,000 Patients Per Year

WHAT IS SSNHL?

The sudden onset of a significant hearing loss due to dysfunction of the cells of the cochlea and central auditory structures.

Hearing loss develops over less than 72 hrs, hearing sensitivity is reduced by at least 30 dB (1,000 fold) in the affected ear(s).

>70% of cases are idiopathic, known causes include noise/head trauma, ischemia, infection.

>50% of patients suffer from permanent disabling hearing loss, mostly those with initial severe/profound hearing loss.

Complications significantly impact quality of life due to:

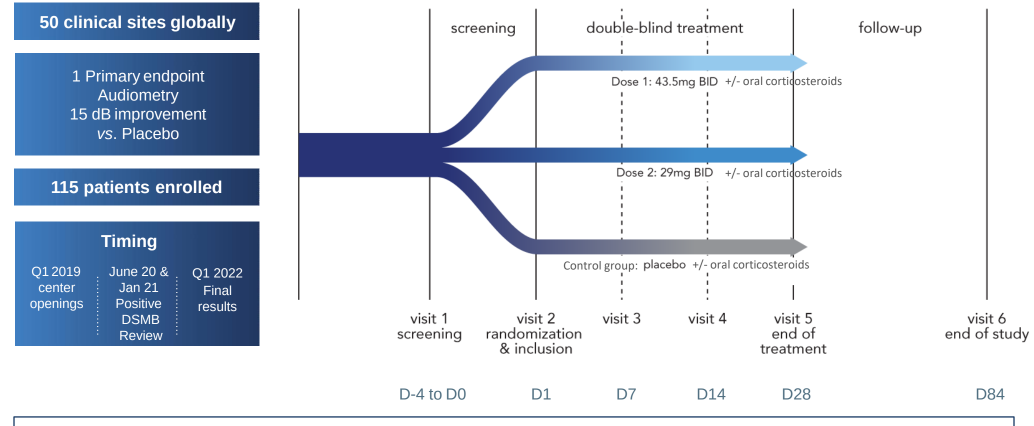
- Difficulties in communicating, social isolation, cognitive decline
- Accompanying tinnitus

Incidence: 27-35 per 100,000 (218,000 patients in 2017 in G7 countries)¹

¹ Company/ estimates based on publicly available data (in the US, Japan, Germany, France, the UK, Italy and Spain)

SENS-401 SSNHL Program: AUDIBLE-S Phase 2 Design

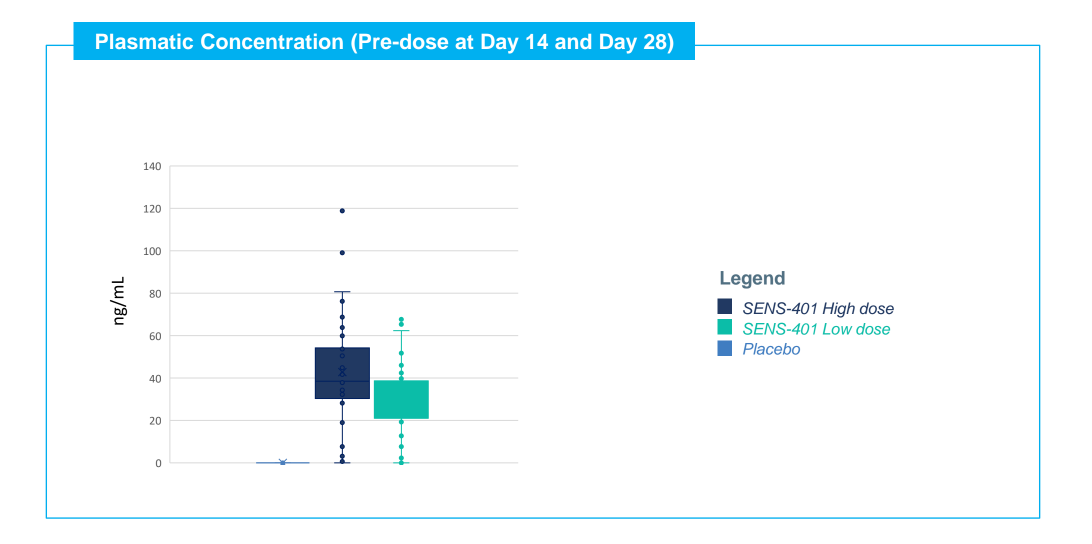
A RANDOMIZED, MULTICENTER, DOUBLE-BLIND, PLACEBO-CONTROLLED TRIAL



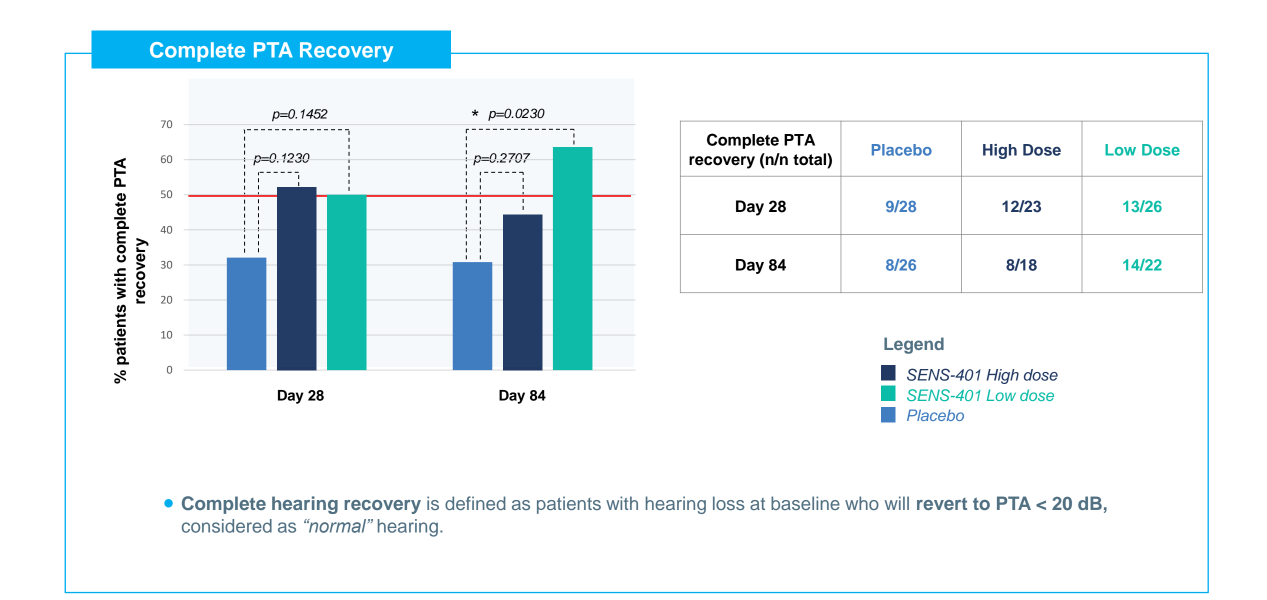
Primary endpoint definition:

"...change in pure tone audiometry (PTA); average of the hearing threshold of 3 contiguous most affected hearing frequencies in decibels in the affected ear from baseline to the end of treatment visit (Visit 5/D28±3)"

SENS-401 Plasmatic Exposure



SENS-401 Induces Complete PTA Recovery In 50% Of Patients



SENS-401 SSNHL Phase 2 Results Summary

Seeking Partners For Late-Stage Development And Commercialization

AUDIBLE-S SECONDARY ENDPOINT RESULTS

- Complete PTA recovery is achieved in 50% of the SENS-401 treated patients
- SENS-401 shows a clinically meaningful and statistically significant effect on PTA change (at least 10 dB) over time in a large homogeneous idiopathic population of patients treated with corticosteroids
- SENS-401 induces a significative PTA change of at least 19 dB at day 28 and up to 25 dB at Day 84 allowing a reduction of the hearing loss degree from profound to mild, in large profound hearing loss sub-group
- A better response was observed in both treatment groups with a **continuous improvement between Day 28 and Day 84**
- The change in PTA translates into functional improvement evidenced with speech audiometry tests
- Safe and well tolerated in 115-patient SSNHL study; although primary endpoint not met data supports and informs further clinical development
- Responder rate is always better in the treated group compared to Placebo and difference with Placebo increases over time