UNITED STATES SECURITIES AND EXCHANGE COMMISSION WASHINGTON, D.C. 20549

FORM 8-K

CURRENT REPORT Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): April 5, 2022

FREQUENCY THERAPEUTICS, INC.

(Exact name of Registrant as Specified in Its Charter)

Delaware (State or Other Jurisdiction of Incorporation)

001-39062 (Commission File Number)

47-2324450 (IRS Employer Identification No.)

75 Hayden Avenue, Suite 300 Lexington, MA 02421 (Address of principal executive offices) (Zip Code)

(781) 315-4600

(Registrant's telephone number, include area code)

 $$\mathbf{N}/\mathbf{A}$$ (Former Name or Former Address, if Changed Since Last Report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions (see General Instructions A.2 below):

Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

П Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

	Trading	Name of each exchange
Title of each class	Symbol(s)	on which registered
Common stock, par value \$0.001 per share	FREQ	The Nasdaq Stock Market LLC (The Nasdaq
		Global Select Market)

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§ 230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).

Emerging growth company 🗵

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 2.05. Costs associated with Exit or Disposal Activities

On April 8, 2022, Frequency Therapeutics, Inc. (the "Company") announced a reduction in force (the "Reduction") of approximately 30% of its workforce. The purpose of the Reduction, which was approved by the Board of Directors (the "Board") of the Company on April 5, 2022, is to better align the Company's workforce with the needs of its business and focus more of its capital resources on its clinical program for its lead candidate for hearing restoration (FX-322); a second pre-clinical program for hearing restoration (FX-345); and a pre-clinical program for remyelination in Multiple Sclerosis. These changes will preserve capital, ensuring that the Company is appropriately resourced to advance its pipeline of potential first-in-class treatments through key development milestones. These milestones are the completion of the Phase 2b study of FX-322, a Phase 1b study of FX-345, and a Phase 1 study in the Multiple Sclerosis program.

A majority of the Reduction has already taken place, and the remainder will be completed by July 31, 2022, though all impacted employees have been notified. The total costs related to the Reduction are estimated to be approximately \$1.2 million in future cash outlays primarily related to severance costs and related expenses.

Item 5.02. Departure of Directors or Certain Officers; Election of Directors; Appointment of Certain Officers; Compensatory Arrangements of Certain Officers.

On April 5, 2022, the Board appointed Richard M. Mitrano, the Company's Vice President of Finance and Operations, as the Company's principal financial officer and principal accounting officer, succeeding Peter P. Pfreundschuh, the Company's former Chief Financial Officer.

Richard M. Mitrano, 51, has served as the Company's Vice President of Finance and Operations since July 2016. From 2012 to 2015, Mr. Mitrano served as the Director of Finance and Operations of Semprus, where he oversaw all accounting and finance operations and provided strategic direction and oversight. Prior to Semprus, Mr. Mitrano was a contract Accounting Manager for Predictive Biosciences, Inc. ("Predictive"), a diagnostics company, from 2010 to 2012. Prior to Predictive, from 2008 to 2010, Mr. Mitrano served as Corporate Controller of Pioneer Behavioral Health, a company providing behavioral health services. Mr. Mitrano holds a B.A. in Accounting from Bentley University.

Item 7.01. Regulation FD Disclosure.

On April 8, 2022, the Company posted an updated corporate slide presentation in the "Investors & Media" portion of its website at www.frequencytx.com. A copy of the slide presentation is attached as Exhibit 99.1 to this Current Report on Form 8-K (the "Current Report").

The information in Item 7.01 of this Current Report, including Exhibit 99.1 attached hereto, is intended to be furnished and shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liabilities of that section, nor shall it be deemed incorporated by reference in any filing under the Securities Act of 1933, as amended (the "Securities Act"), or the Exchange Act, except as expressly set forth by specific reference in such filing. The Company undertakes no obligation to update, supplement or amend the materials attached hereto as Exhibit 99.1.

Item 9.01. Financial Statements and Exhibits.

(d) Exhibits

The following Exhibit 99.1 relates to Item 7.01, and shall be deemed to be furnished, and not filed:

Exhibit No.	Description
99.1	Frequency Therapeutics, Inc. Corporate Slide Presentation as of April 8, 2022
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

FREQUENCY THERAPEUTICS, INC.

Date: April 8, 2022

/s/ David L. Lucchino

 By:
 /s/ David L. Lucchino

 Name:
 David L. Lucchino

 Title:
 President and Chief Executive Officer

Pioneering a New Category in Regenerative Medicine

Frequency Therapeutics Corporate Presentation April 2022

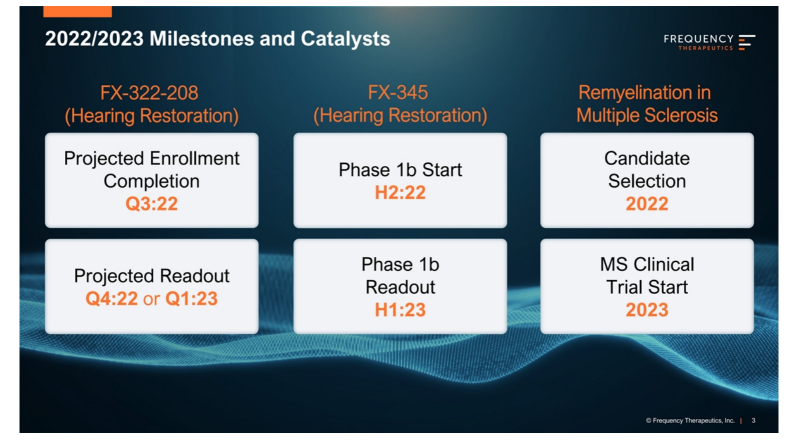


Forward-Looking Statements and Other Disclaimers

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements contained in this presentation that do not relate to matters of historical fact should be considered forward-looking statements, including without limitation statements regarding the timing and design of Frequency Therapeutics' (the "Company") new Phase 2b trial of FX-322, including the type of SNHL that the enrolled patients will have and the ability of design features to reduce bias, the interpretation and implications of the results and learnings of other FX-322 clinical studies, the acceptance by the FDA of particular endpoints in the Company's trials, the treatment potential of FX-322, FX-345, and the novel approach for remyelination in multiple sclerosis, the timing and progress of the FX-345 and remyelination programs, the sufficiency of the size of the hearing [oss population and population at risk for hearing loss, estimates of the size of the population with multiple sclerosis, estimates of the potential application of the PCA platform to provide patient benefit, and paradigms, the ability of our technology platform to orther diseases.

These forward-looking statements are based on management's current expectations. These statements are neither promises nor guarantees, but involve known and unknown risks, uncertainties and other important factors that may cause actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements, including, but not limited to, the following: the impact of COVID-19 on the Company's ongoing and planned clinical trials, research and development and manufacturing activities, the Company's business and financial markets; the Company has incurred and will continue to incur significant losses and is not and may never be profitable; need for additional funding to complete development and commercialization of any product candidate; the Company's dependence on the development of FX-322; the unproven approach of the PCA platform; the lengthy, expensive and uncertain process of clinical drug development and regulatory approval; limited experience successfully obtaining marketing approval for and commercializing product candidates; the results of earlier clinical trials not being indicative of the results from later clinical trials; differences between preliminary or interim data and final data; adverse events or undesirable side effects; disruptions at the FDA and other regulatory agencies; failure to identify additional product candidates; new or changed legislation; failure to maintain Fast Track designation for FX-322 and such designation failing to result in faster development or regulatory review or approval; costly and damaging litigation, including related to product liability, intellectual property or brought by stockholders; dependence on Astellas Pharma Inc. for the development and commercialization of FX-322 outside of the United States; misconduct by employees or independent contractors; reliance on third parties, including to conduct clinical trials and manufacture product candidates; compliance with laws and regulations; failure to obtain, maintain and enforce protection of patents and other intellectual property; security breaches or failure to protect private personal information; attracting and retaining key personnel; and ability to manage growth.

These and other important factors discussed under the caption "Risk factors" in the Company's Form 10-K filed with the Securities and Exchange Commission (SEC) March 15, 2022 and its other reports filed with the SEC could cause actual results to differ materially from those indicated by the forward-looking statements made in this presentation. Any such forward-looking statements represent management's estimates as of the date of this presentation. While the Company may elect to update such forward-looking statements at some point in the future, it disclaims any obligation to do so, even if subsequent events cause its views to change. These forward-looking statements should not be relied upon as representing the Company's views as of any date subsequent to the date of this presentation.



A Vision Built on Regeneration

Since 2014, Frequency has been developing *small molecule therapeutics* to activate a person's innate regenerative potential, within the body, to repair tissue and restore human function.



Power of the Progenitor Cell Activation (PCA) Approach

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No Change to Genome

Activating native programs, reducing safety concerns

Harnessing Innate Biology

Progenitors already located within the target tissue

Ease of Manufacturing

Use of small molecules: no need to remove or grow cells *ex vivo*

A Series of Firsts in Hearing Restoration

First PK/PD shown for a hearing therapeutic candidate First clinical studies to show hearing improvements

First speech perception improvements measured First to show sustained improvements and continued improvements over time

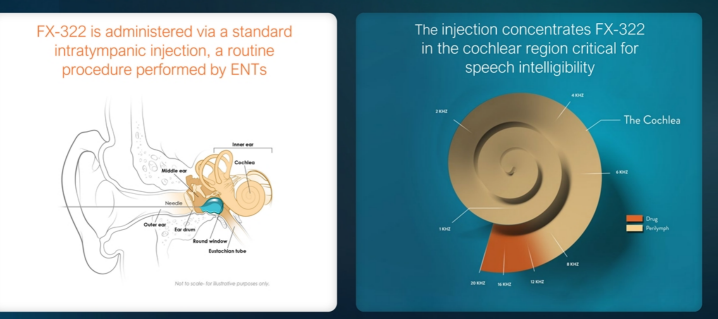
FX-322: A Small Molecule Candidate to Address the Underlying Pathology

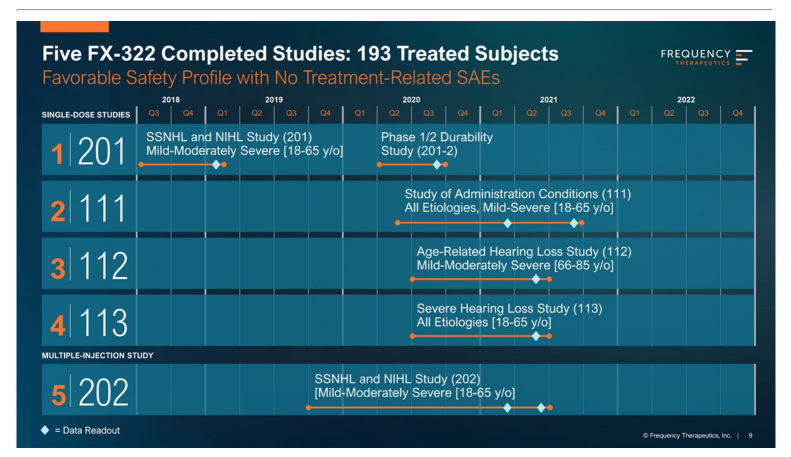
Synergy between pathways aims to activate progenitor cells and regenerate sensory cells in the cochlea



FX-322:

Directly Targeting the Regeneration of Sensory Hair Cells in the Cochlea





FX-322-201, FX-322-111, FX-322-113

Single-Dose Safety Studies with Hearing Improvement Signal



FREQUENCY

Data from Controlled Studies (FX-322-201, FX-322-111)



Improvement Shown in Speech Perception in Quiet with Single Dose

Phase 1/2 Study FX-322-201 Overview

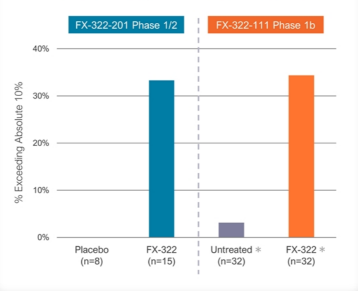
- Placebo-controlled, multi-center, randomized study
- Mild to moderately severe subjects, age 18-65 (n=23)
- NIHL/SSNHL

Study Results

- 33% of subjects achieved 10% or greater absolute improvement in word recognition in treated ear
- Statistically significant and clinically meaningful improvements in WR
- No meaningful changes in placebo group

Favorable safety profile

Day-90 Word Recognition Scores Across Studies



Phase 1b Study FX-322-111 Overview

- Compared different FX-322
- administration conditions • Open-label, multi-center, randomized study
- Mild to severe subjects, age 18-65 (n=33)

Study Results

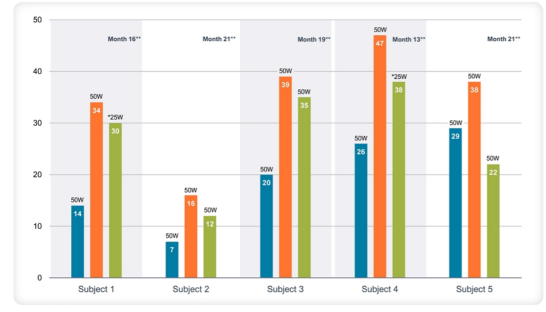
- 34% of subjects achieved 10% or greater absolute improvement in word recognition (WR) in treated ear
- Statistically significant and clinically meaningful improvements in WR
- Favorable safety profile

*Total of 33 patients enrolled in study, 32 subjects completed 90-day clinical assessment period

FX-322 Phase 1/2 Durability Data:



Patients Show Sustained Hearing Improvements 13-21 Months After Initial Dosing



Key Findings

Preliminary evidence indicating a durable benefit of hearing clarity

Baseline - Correct words out of 50

Day 90 - Correct words out of 50

1-2 Years - Correct words out of 50

Three patients who had durable improvements in intelligibility also had pure tone audiometry improvements of 10 - 15 dB at the highest frequency tested (8k Hz)

* 25W = 25 Word test performed outside an official study site at 13-18 months after dosing; results scaled to 50 words 50W = 50 Word test performed under a formal protocol at original study site at 18-21 months after dosing **Since FX-322 dosing

Subjects in FX-322-111 Study Show Additional Hearing Improvements at Later Time Points

Conducted longer-term, follow-up of FX-322-111 study subjects

• 25 of 33 study subjects evaluated at 8-12 months following FX-322 dosing

Results show some FX-322 dosed subjects accumulated hearing benefits over time

• 3 subjects that had shown improvement trends in word recognition scores at day 90, achieved statistically significant scores when tested at the later time points

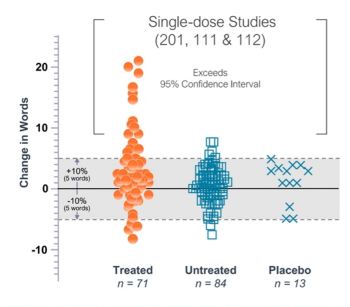
To date, 8 of 32 evaluated study subjects have shown statistically significant improvements in speech perception scores in treated ears between 90 days and 1 year

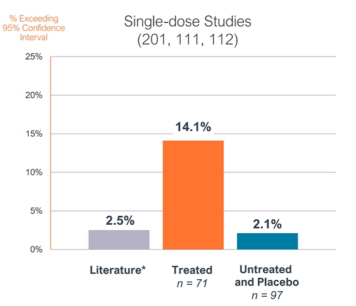
No change observed in untreated ears

Pooled FX-322 Data Shows Patterns of Response



Single-dose Studies (201, 111, 112) Exceed 95% Confidence Interval





95% confidence intervals established by Thornton & Raffin (1978) and modified by Carney & Schlauch (2007)

FX-322-113: Hearing Signal and Speech Perception Improvements Observed in Subjects with Severe SNHL

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Double-blind, placebo-controlled study of 31 individuals randomized 4:1

- Pure tone average deficit between 71-90 decibel hearing level (dBHL)
- Potential cochlear implant candidates

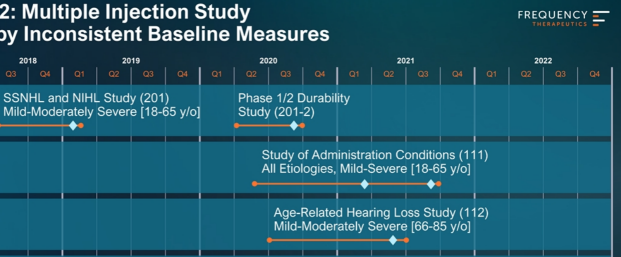
Improvements in Bamford-Kowal-Bench Sentence-in-Noise exam (BKB-SIN) observed in treated ears

- BKB-SIN measures signal-to-noise ratios required for subjects to correctly repeat words in sentences
- Four FX-322 treated subjects show improvement, two with a 6 dB response
- A single placebo subject showed a 3.6 dB change
- No improvements observed in words-in-quiet

Favorable safety profile

No treatment-related SAEs

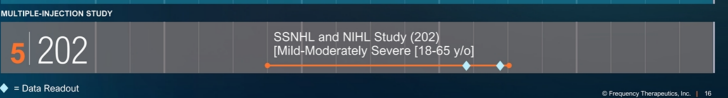
FX-322-202: Multiple Injection Study Impacted by Inconsistent Baseline Measures



Severe Hearing Loss Study (113)

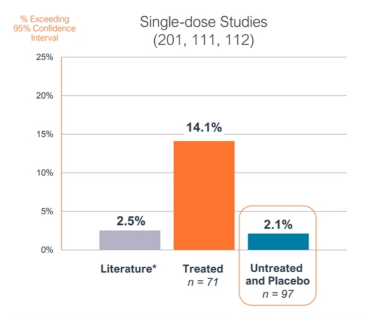
All Etiologies [18-65 y/o]

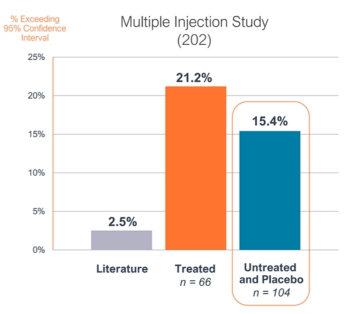
SINGLE-DOSE STUDIES



Comparing Pooled Data to Multiple-Injection Study FX-322-202 FREQUENCY

Placebo-Treated and Untreated Ears are Outside 95% Confidence Interval





95% confidence intervals established by Thornton & Raffin (1978) and modified by Carney & Schlauch (2007)



Clinical Study Data Informs New FX-322 Phase 2b Study

New Clinical Study FX-322-208 Designed to Advance Drug Candidate to Pivotal Trials

Built upon insights from trials with hearing restoration signal

Etiology, severity, baseline speech perception Sufficient sample size to demonstrate efficacy

Approach based on pooled data

Primary endpoint of speech perception

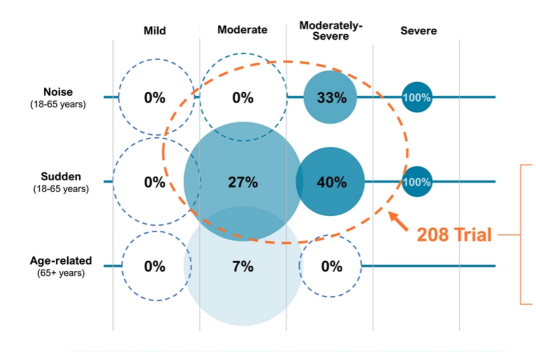
Reduce potential for bias

Multiple baseline measures

Multiple speech perception tests

Pooled Single-Dose Studies (201, 111, 112)





71 Treated with single-dose of FX-322 The size of each circle represents the

number of people tested per group The color of the circle represents the percentage of responders

208 Trial: Target Population 7-10 Million U.S. patients

FX-322: Extended Population

15+ Million U.S. patients

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Multiple Design Features Have Been Added to Mitigate Bias



And Demonstrate Greater Separation Between Signal and Placebo

 \checkmark

Lead-in phase with multiple baseline measures

Sites and patients masked to qualifying test results



All sessions recorded and monitored

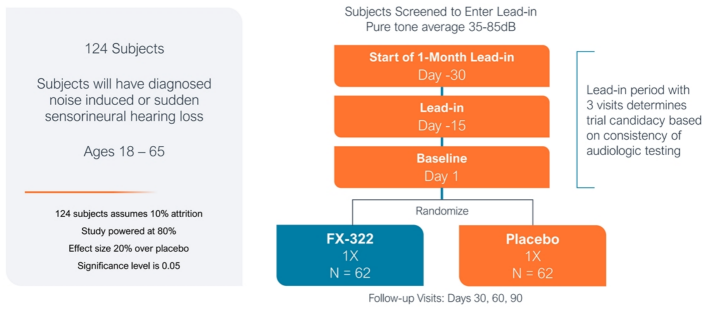
Ability to disqualify subjects based on symptom stability

Start of 1-Month Lead-in Day -30
Lead-in
Day -15
Baseline Day 1

New FX-322 Placebo-Controlled Phase 2b Study Commenced



First patient dosed in FX-322-208 Study in October 2021



FDA Type C Meeting Held to Gain Alignment



ALIGNMENT

Primary Endpoint

Gained alignment with FDA on speech perception as the primary endpoint

208 Study Design

FDA reviewed and commented on 208 study, comments were incorporated into study protocol

Patient Reported Outcomes (PRO)

FDA feedback provided on novel PRO development called **RADIAL**; special meeting granted for further discussion



Today's Hearing Loss Market Has No Restorative Treatments



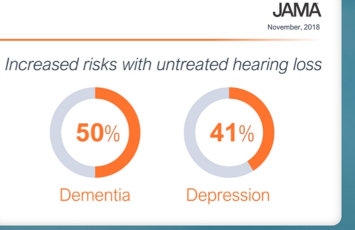
*Source: World Health Organization

Hearing Loss Can Have a Significant Impact on Overall Health

FREQUENCY



"Hearing loss is the largest potentially modifiable risk factor for developing dementia"



AMA Nov 8, 2018, Deal J, et al. Incident Hearing Loss and Comorbidity. A Longitudinal Administrative Claims Study.

Pipeline Expansion





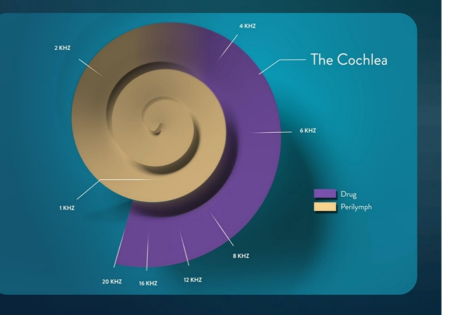
New Regenerative Program

What if we were able to get drug deeper into the cochlea?

FX-345 Working to Achieve Broad Exposure Through the Cochlea

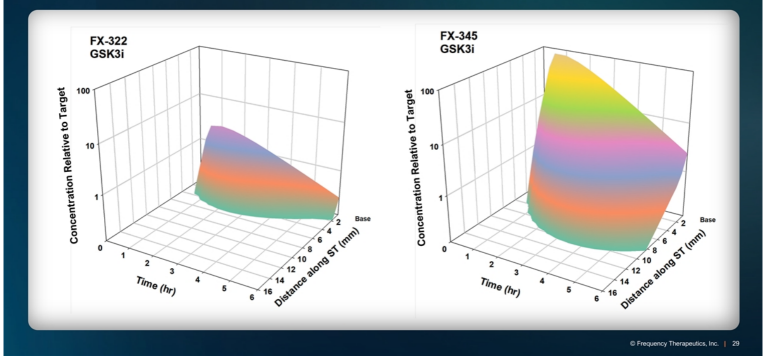


- Second clinical program focused on regrowth of sensory cells
- Enables coverage of large portion of cochlea
- Potential to address additional SNHL patient types
- Formulation enabling evaluation of a range of dose levels
- Developing in addition to FX-322. Clinical data will drive commercial positioning



FX-345 – A New Development Candidate

Creating Effective Drug Levels Through Large Portion of Cochlea

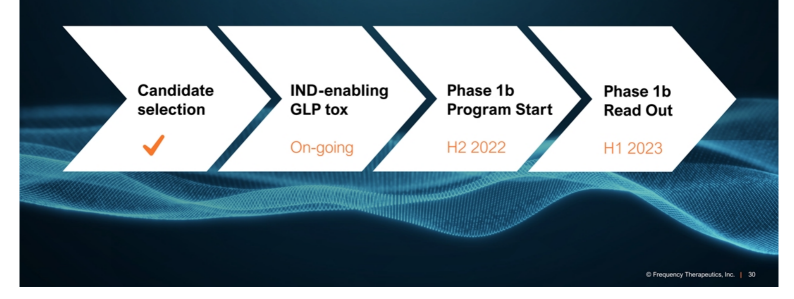


FX-345 Path to Clinic



Program start planned for H2:2022 for a Phase 1b study in patients with SNHL

Enables us to clinically evaluate increased cochlear coverage across range of doses in multiple patient populations

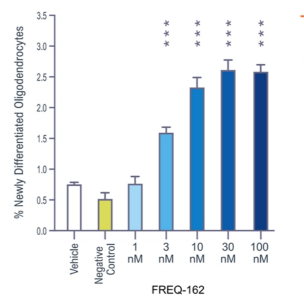




Novel Frequency Small Molecule Inhibitors Drive Oligodendrocyte Differentiation



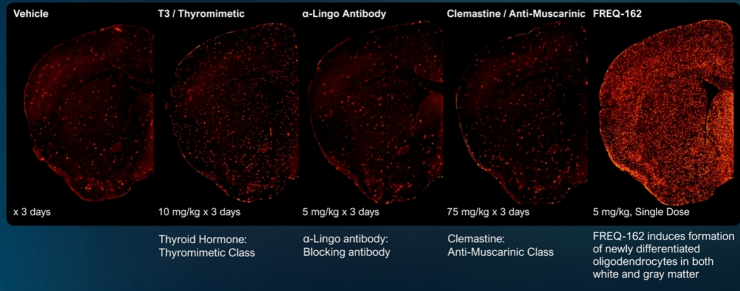
Developed novel chemical entities that are highly potent inducers of oligodendrocyte differentiation Lead Optimization generated FREQ-162



Highly potent Highly efficacious Orally bioavailable Brain penetrant Novel chemical entity Patent application filed

FREQ-162 Outperforms Literature Compounds In Vivo

Adult mice received 3 doses of comparator compounds or a single dose of FREQ-162 Brains were stained for a marker of newly generated oligodendrocytes Single dose FREQ-162 induces more OPCs to differentiate than comparator compounds

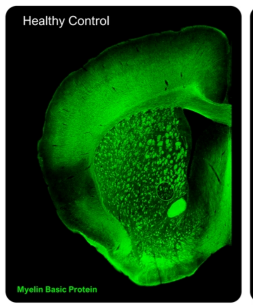


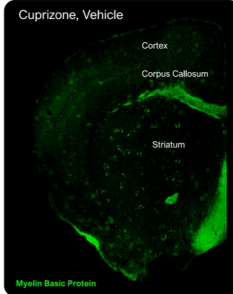
© Frequency Therapeutics, Inc. | 33

FREQUENCY

The Cuprizone Model of Chronic Demyelination







Adult mice were demyelinated via 17 months of cuprizone administration

Elderly mice with long term demyelination

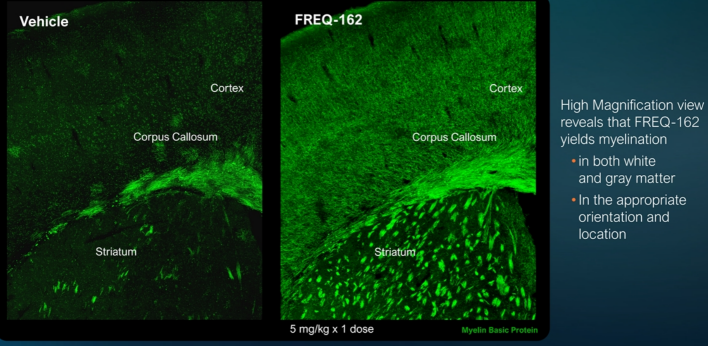
FREQ-162 Outperforms Published Compounds In Vivo

Adult mice received up to 10 daily doses of comparators or a single dose of FREQ-162 Brains were stained for Myelin Basic Protein (green) Single dose FREQ-162 induces more remyelination than comparator compounds



FREQUENCY

Frequency NCEs Outperform Competitors: High Magnification

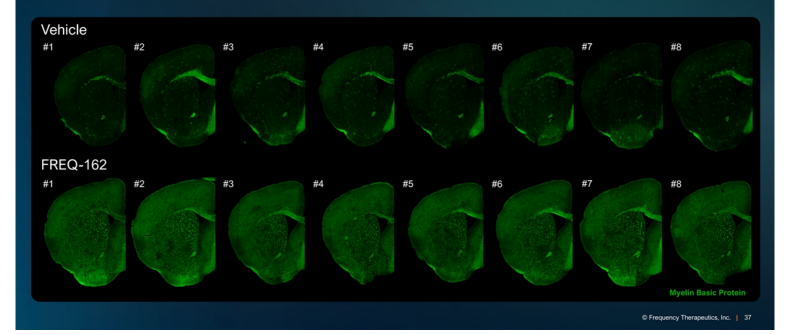


reveals that FREQ-162 yields myelination • in both white

- and gray matter
- In the appropriate orientation and location

FREQ-162: Highly Reproducible Increases in Myelination

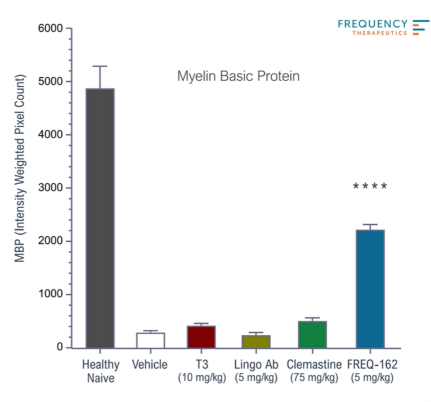
All 8 out of 8 mice treated with FREQ-162 showed robust increases in myelination in both white and gray matter tracts



Freq-162 Induces Robust Increases in Myelination

- Forebrain myelin basic protein levels quantitated
- A single dose of a Frequency compound induces robust remyelination

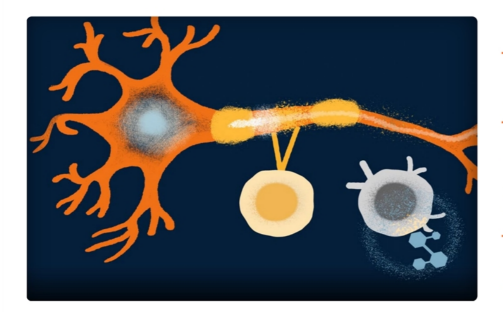
Dose (mg/kg)	# of doses	Fold change	P=
5	3	0.9 x	0.99
75	10	1.7 x	0.70
10	10	1.4 x	0.95
5	1	7.7 x	<0.0001
	(mg/kg) 5 75 10	(mg/kg) # of doses 5 3 75 10 10 10	(mg/kg) # of doses change 5 3 0.9 x 75 10 1.7 x 10 10 1.4 x



[©] Frequency Therapeutics, Inc. | 38

Remyelination: Path Forward





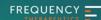
Discovered novel target

Generated multiple compounds

Induced high levels of oligodendrocyte differentiation and remyelination *in vivo*

Initiating IND enabling studies

Our Path Forward





We know characteristics of FX-322 responders.

Learnings from previous trials informed new trial design with strong controls and FDA aligned clinical endpoints.

We have a compelling new hearing program that will allow us to explore the impact of going deeper into the cochlea.

We also have an exciting remyelination program in multiple sclerosis with a novel target and a strong response *in vivo*.

We are a well capitalized company with resources to deliver innovation for patients and value for investors.

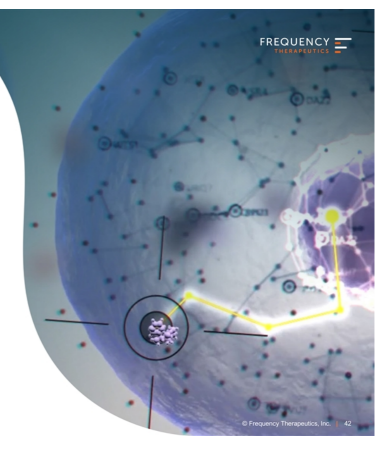
• \$142.4m in cash and cash equivalents*, runway into 2024

• Ex-US partnership with Astellas, significant milestones and royalties

*Number reflects unaudited Cash, Cash Equivalents, and Marketable Securities as of December 31, 2021, and does not include Restricted Cash



Broad Potential of Progenitor Cell Activation Approach



Origin of Frequency Therapeutics

Tissue-Specific, Pre-programmed Stem Cells



Decoding Intestinal Regeneration

Langer and Karp publish small molecules activate intestinal progenitors



Niche-independent high-purity cultures of Lgr5+ intestinal stem cells and their progeny

Enabling Cochlear Regeneration

Same cues reactivate normally inactive progenitors in the cochlea

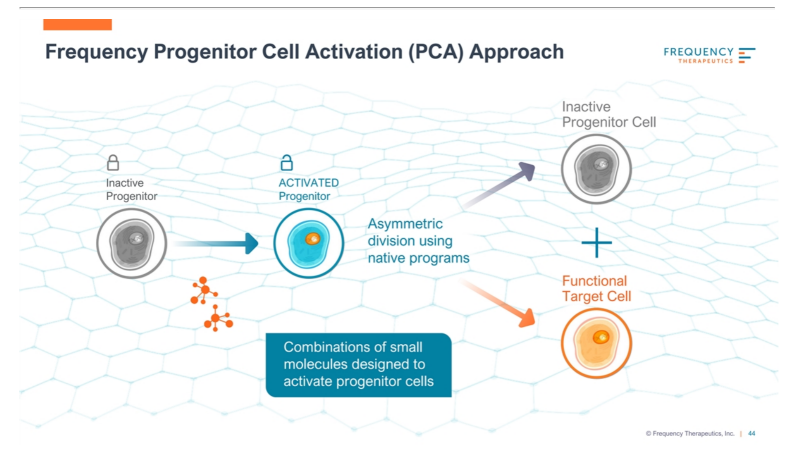


Clonal Expansion of Lgr5-Positive Cells from Mammalian Cochlea and High-Purity Generation of Sensory Hair Cells

Frequency Therapeutics

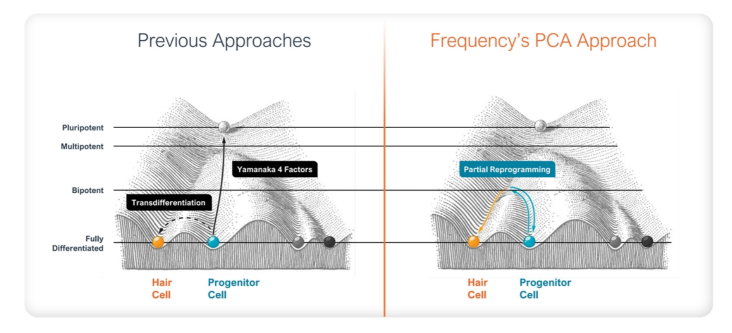
Small molecule therapeutics show clinical proof of concept





Uniqueness of Our PCA approach

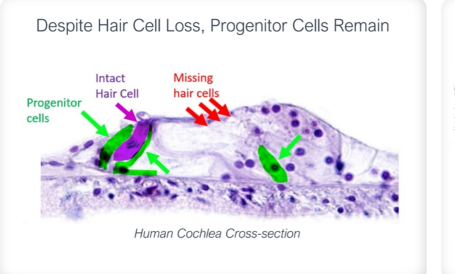


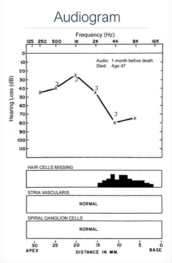


Our Approach:

Activation of Progenitors to Replace Hair Cell Loss





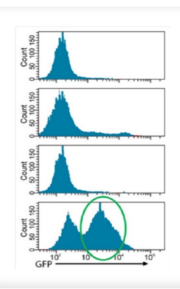


47 Year Old Male with Occupational Noise Deafness

Profound Synergy Between Pathways to Regenerate Cells

Cochlear Progenitor Proliferation (Lgr5+ – GFP)

HDAC = Histone deacetylase NCE = new chemical entity In vitro mouse model testing



Culture Media
Wnt Activation (glycogen synthase kinase-3
(GSK3) Inhibitor; NCE)
HDAC Inhibition
(sodium valproate)
Wnt Activation + HDAC inhibition
PROFOUND SYNERGY

FX-322 Agents Induce Protein Expression Consistent with Fully Functional Sensory Hair Cells

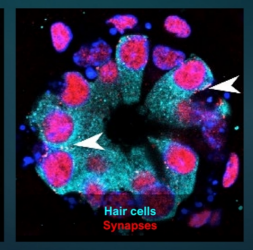


Sensing Sound Generating intricate hair bundles

McLean et al., 2017, Cell Reports 18, 1917–1929 February 21 http://dx.doi.org/10.1016/j.celrep.2017.01.066



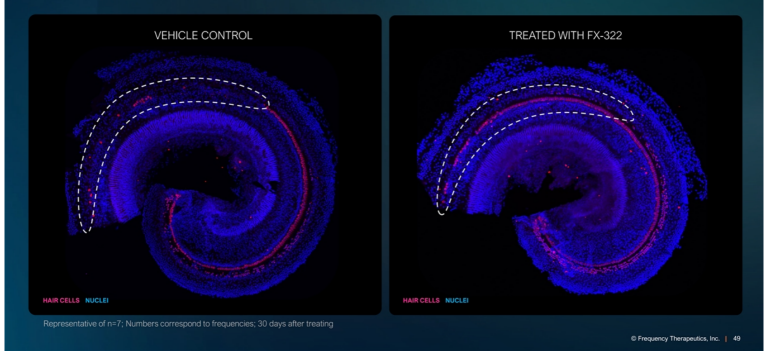
Creating Signal Producing functional ion channels



Transmitting Signal Synaptic proteins to communicate with nerve are present

Images Showing Cellular Regeneration

In Vivo Hearing Loss Model







Astellas Collaboration:

Ex-US Development and Commercialization of FX-322

- Development and commercialization collaboration for FX-322, including lifecycle improvements
- Astellas has ex-US rights; Frequency retains US rights to FX-322
- Payments of up to \$625mm which included \$80mm upfront
 - Development milestone payments to Frequency of \$65.0 million and \$25.0 million upon the first dosing of a patient in a Phase 2b clinical trial for SNHL in Europe and Asia, respectively
 - \$100.0 million and \$40.0 million upon the first dosing of a patient in a Phase 3 clinical trial for SNHL in Europe and Asia, respectively

Development & commercialization:

Astellas responsible for execution and costs of ex-US clinical development and commercialization





Strategic commitment to invest in ENT as a therapeutic area

Research focus in regenerative medicine

Global footprint in major markets and distributorship model in Africa/ME and LATAM

Proven Leadership Team



David Lucchino President, CEO & Co-Founder

Former CEO of Entrega Bio (PureTech). Co-founder / CEO of Semprus BioSciences (acquired), Polaris Partners. MIT Sloan Fellow.



Dana Hilt, M.D. Chief Medical Officer

Neurologist and neuroscientist with two decades in biopharma and CNS drug development. Amgen, Lysosomal, Forum Pharma.



Quentin McCubbin, Ph.D. Chief Manufacturing Officer

Led pharmaceutical sciences and process chemistry at Takeda / Millennium and headed technical operations Cerevel Therapeutics.





Co-founder/CTO of Semprus BioSciences through FDA / CE clearance and acquisition. Princeton, MIT, Hertz Fellow and Yale Faculty.

Sue Stewart, J.D., LLM Chief Regulatory Officer

CRO at numerous biopharma companies including Kaleido Biosciences, Candel Therapeutics, and regulatory leadership roles at Tokai Pharma, Transmolar and Genzyme Corp.



Carl Lebel, Ph.D. Chief Development Officer

Chief Scientific Officer of Otonomy (2009 to 2016). Executive Director, Amgen. Scientific fellow of the American Academy of Otolaryngology.



HR leader with extensive life science experience including senior leadership roles at Kaleido Biosciences, Moderna, Celgene Avilomics

Research, and Inotek Pharmaceuticals

Scientific Advisory Board

Robert Langer,

SC.D.



Jeff Karp, Ph.D.

Associate Professor at Brigham and Women's Hospital, Harvard Medical School





Siddhartha Sean J. Morrison, Ph.D. Director of the Children's Medical Center Research Institute, UT Southwestern

Mukherjee, M.D., D.Phil. Assistant Professor of Medicine, Columbia University Medical Center



Robin Franklin, Ph.D. Professor of Stem Cell Medicine, Wellcome Trust-MRC Cambridge Stem Cell Institute David H. Koch Institute Professor at the Massachusetts Institute of Technology

Senior Investigator, Gladstone Institute of Cardiovascular Disease



Amy Wagers, Ph.D.

Forst Family Professor of Stem Cell and Regenerative Biology, Harvard University



Sheng Ding, Ph.D.



Dan Lee, M.D.

Director, Pediatric Otology and Neurotology, Mass Eye and Ear



Chris Runge, Ph.D.

Chief of the Division of Communication Sciences, Medical College of Wisconsin

Rene Gifford, Ph.D.

Clinical Advisory Board







MD, Ph.D.

Assistant Professor of Clinical Otolaryngology-Head and Neck Surgery, Keck School of Medicine of USC.





Ruth Litovsky, Ph.D.

Professor, Communications Sciences and Disorders and Surgery Division of Otolaryngology, University of Wisconsin



David Friedland,

M.D., Ph.D.

Julie Arenberg, MS, Ph.D.

Associate Director of Clinical Audiology for Research and Education, Mass Eye and Ear Vice-Chair of the Department of Otolaryngology and Communications Sciences, Medical College of Wisconsin

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Director, Vestibular Division, Medical Director, Mass. Eye and Ear Balance and Vestibular Center

Steve Rauch, M.D.



Pioneering a New Category in Regenerative Medicine

Frequency Therapeutics Corporate Presentation April 2022

