

CORPORATE OVERVIEW

Q2 2024

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FORWARD LOOKING STATEMENTS



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WHO WE ARE



Pioneering Novel Gene Therapeutics for a Longer, Healthier Life

MISSION: aging is underlying risk factor for disease, and our aim is to deliver gene therapeutics that potentially halt or slow the aging process in humans and dogs

PROMISING PRECLINICAL RESULTS: lead drug candidate GF-1002 delivers a centenarian variant of the SIRT6 gene for treatment of MASH (formerly known as NASH), which has the greatest potential to progress to cirrhosis, liver failure, and liver cancer

SIRT6 GENE: longevity protein that widely regulates aging and immunity and is considered to be a potential therapeutic target for the treatment of diseases

SEASONED TEAM: experienced and proven management with extensive experience in public & private pharma and biotech

ROBUST PIPELINE: advancing multiple development programs over 24 months



MARKET

genflow biosciences

Aging Is One of Our Greatest Societal & Economical Challenges

Increasing life expectancy,
decreasing healthspan, rising
healthcare costs – all highlight the
urgent need for age-related
disease treatments



Source: Morgan AE, Davies TJ, Mc Auley MT. The role of DNA methylation in ageing and cancer. Proc Nutr Soc. 2018 Nov;77(4):412-422. doi: 10.1017/S0029665118000150. Epub 2018 Apr 30. PMID: 29708096

88 yrs*

LIFE EXPECTANCY



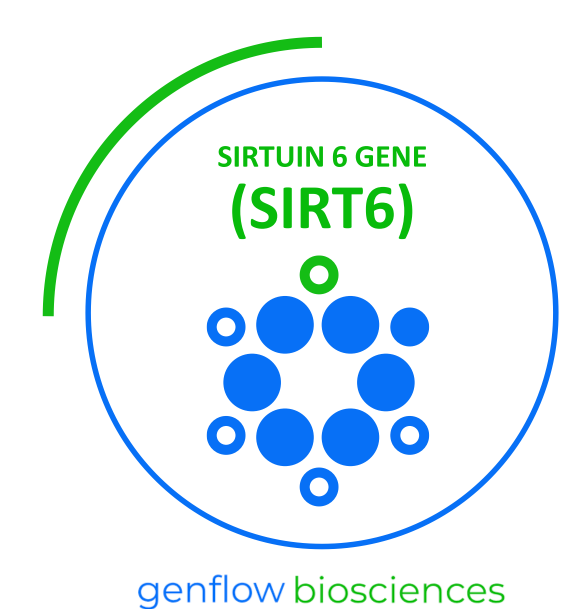


GENE REGULATION IN AGING



Aging is a function of overworked epigenetic regulator genes unable to respond to cellular DNA damage

MANY GENES REGULATE AGING.
OUR FOCUS IS THE **SIRT6 GENE**



Aging is driven by 9 interlinked Hallmarks, all rooted in DNA damage. Targeting one individual factor is unlikely to be effective



SIRT6: REPAIRING DNA



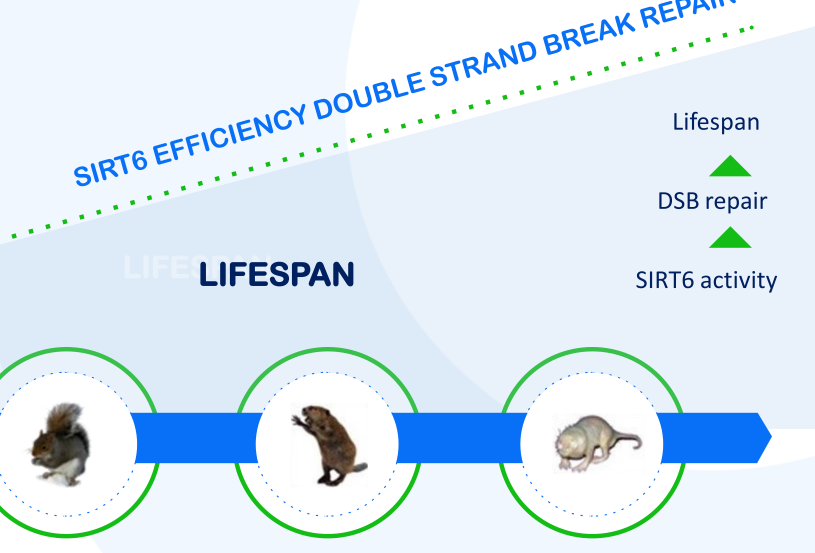
SIRT6 gene/protein repairs DNA damage (especially double strand breaks (DSB)) and prevents senescence of our cells

SIRT6 gene codes for SIRT6 protein

Stronger SIRT6: Longer lifespan

The Ability of SIRT6 to stimulate DSB repair corelates with maximum lifespan (MLS) in rodents

5 Amino Acids determine the differential activities of SIRT6



LSE: GENF - OTCQB: GENFF

Source: Tian et al., 2019, Cell 177, 622–638 April 18, 2019



FOCUS ON CENTENARIAN SIRT6



SIRT6 centenarian variant gene has more efficient DNA repair properties

HOMOLOGOUS

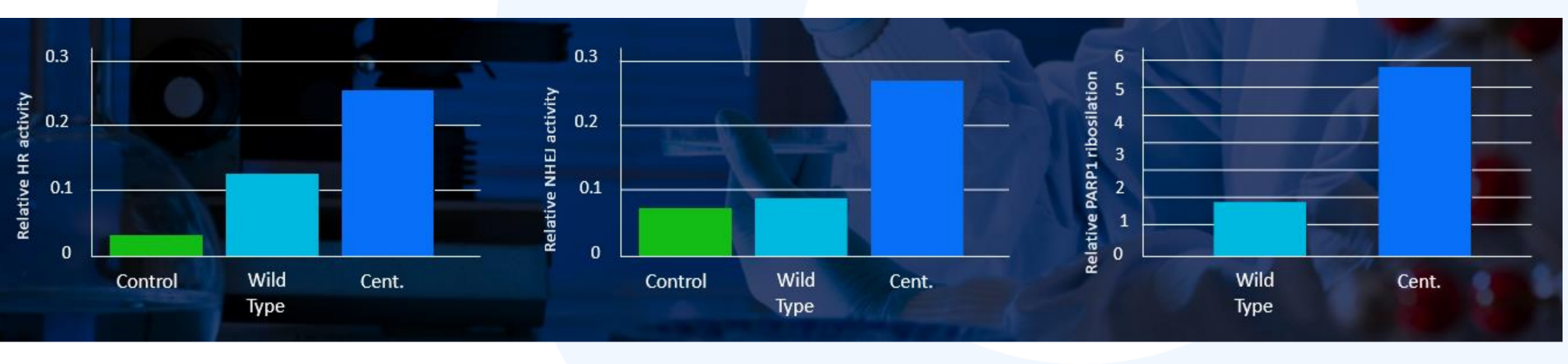
Recombination Repair

NON-HOMOLOGOUS End

Joining Repair

RELATIVE PARP1

Ribosilation



DEVELOPMENT PIPELINE



GF-1002 (Pre-Clinical) –Exo-AAV vector for intravenous infusion, expressing cDNA of centenarian variant of SIRT6 in liver

GF-1003 (Pre-Clinical/Development) Suspension of exosomes, expressing mRNA of centenarian variant of SIRT6 in fibroblasts

GF-1005 (Pre-Clinical/Development) Mitochondrial disfunction: Myoblast progenitors loaded by photoporation with centSIRT6

GF-1004 (Pre-Clinical) Suspension of AAV vector for intravenous infusion, obtaining cDNA of centenarian variant of SIRT6

Phase I/II MASH

PoC in 36 patients
IND-Enabling Phase
18 months to first-in-human

Werner Syndrome

36 months to first-inhuman COLLABORATIONS

Sarcopenia

Pre-clinical

Anti-aging for dogs w/ Vet Partner









MASH PROGRAM



Affects est. 35 million people globally

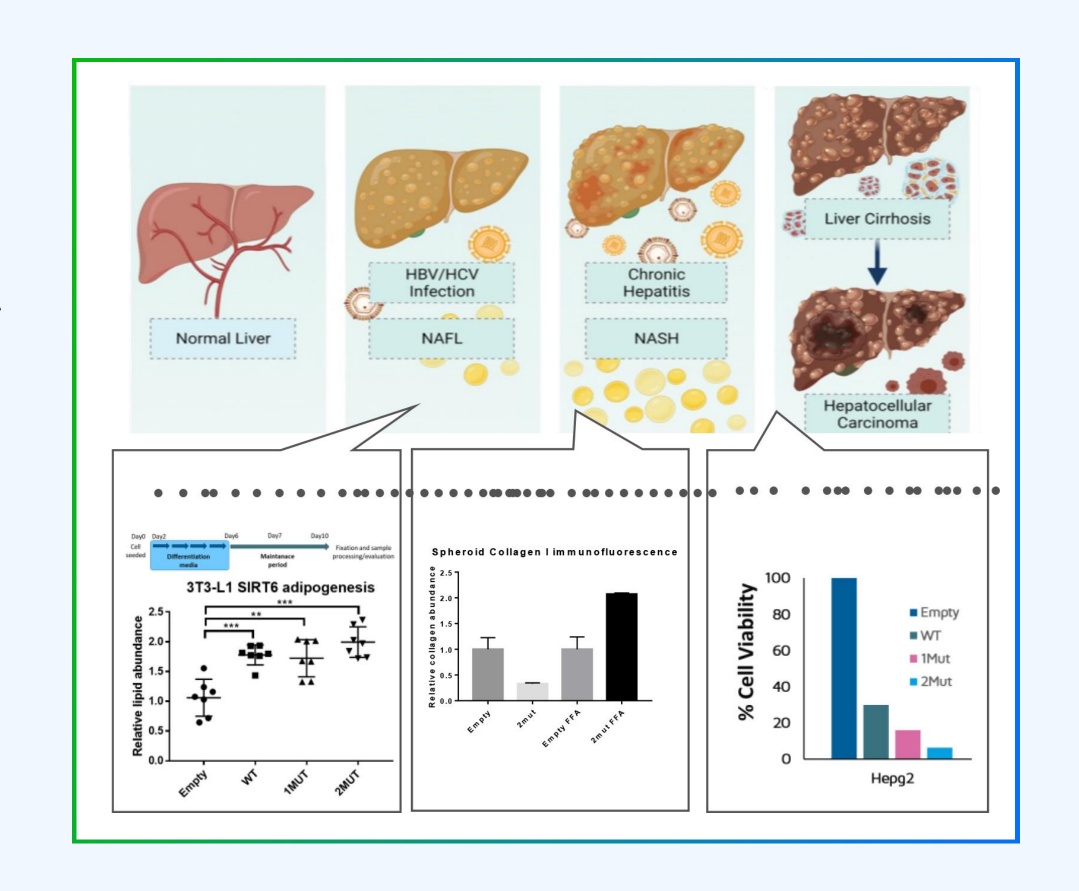
- Increasing prevalence
- Leading cause of chronic liver disease and liver transplant

Significant unmet medical need

Clear regulatory accelerated development pathway. EMA and FDA guidelines accept:

- Key surrogate outcomes for therapeutic trials: regression of fibrosis or resolution of MASH
- ✓ Histological changes are achievable within a 12-18-month time-frame
- ✓ Placebo control
- ✓ Conditional fast-track approval

Pais R, Barritt AS 4th, Calmus Y, Scatton O, Runge T, Lebray P, Poynard T, Ratziu V, Conti F. NAFLD and liver transplantation: Current burden and expected challenges. J Hepatol. 2016 Dec;65(6):1245-1257. Vlad Ratziu, Sven Francque, Arun Sanyal, Breakthroughs in therapies for NASH and remaining challenges, Journal of Hepatology, Volume 76, Issue 6, 2022



EXOSOME DELIVERY SYSTEM: SAFE AND COST-EFFECTIVE



Genflow's patent-pending technology has already been tested in several preclinical studies



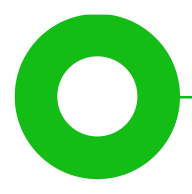
ADVANATGES: EXOSOME DELIVERY



Exo-AAV can mediate efficient, specific, and more durable SIRT6 expression in liver compared to conventional AAV

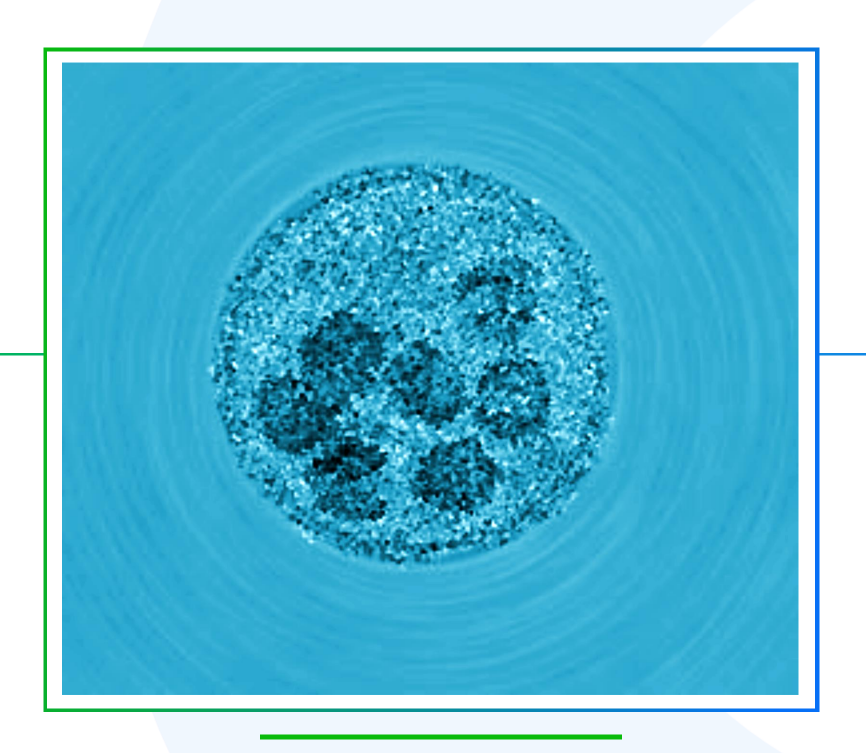
NO IMMUNOGENICITY

Lack of Local Systemic Immunogenicity



TARGETED DELIVERY

Engineered Exosome
To Direct to Specific
Cell Types



POTENCY ADVANTAGE

Improved Transduction
Verses Free AAVs Rapid Uptake
Sustained

THERAPEUTIC WINDOW

Potency Improvement, Local Retention, Lack of Systemic Leakage

Hudry E, Martin C, Gandhi S, György B, Scheffer DI, Mu D, Merkel SF, Mingozzi F, Fitzpatrick Z, Dimant H, Masek M, Ragan T, Tan S, Brisson AR, Ramirez SH, Hyman BT, Maguire CA. Exosome-associated AAV vector as a robust and convenient neuroscience tool. Gene Ther. 2016 Apr;23(4):380-92. doi: 10.1038/gt.2016.11. Epub 2016 Feb 2. Erratum in: Gene Ther. 2016 Nov;23 (11):819. PMID: 26836117; PMCID: PMC4824662.

2023 KEY HIGHLIGHTS & 2024 PRIORITIES



GROWING IP PORTFOLIO

Provisional patent application in 2023 for editing SIRT6 gene, linked to longevity and age-related diseases

EXPANDED MARKET

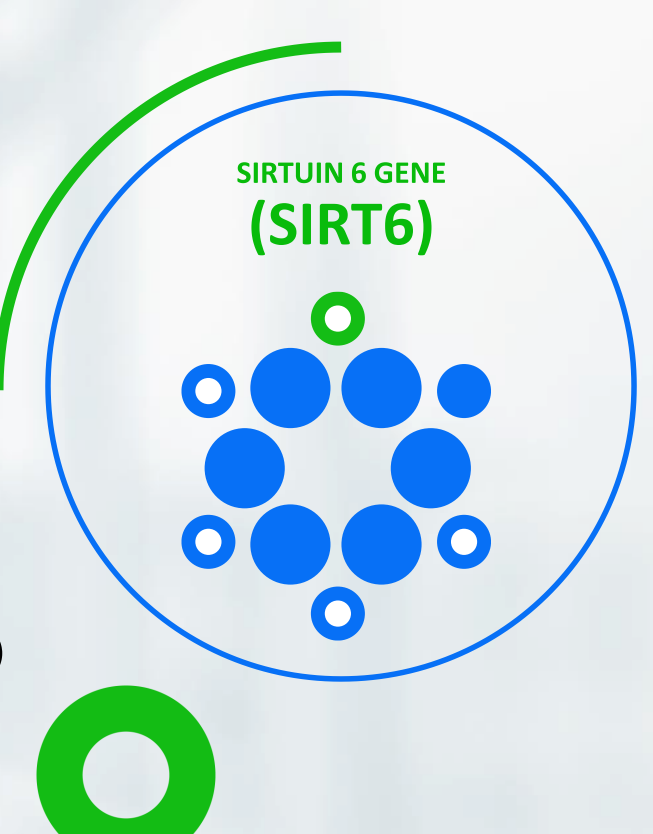
Awarded to new grants in 2023 & 2024 expanding research pipeline and size of therapeutic markets

GF-1002

Undertaking key Investigational New Drug (IND) -enabling development activities to help define pharmacological and toxicological properties and potential benefit to MASH patients

GF-1003

Commencing preliminary discussions with the European Medicines Agency (EMA) on Mechanism of Action (MoA) data for Orphan Drug Application (ODA) targeting Werner Syndrome



2023 KEY HIGHLIGHTS & 2024 PRIORITIES



LARGE MARKET OPPORTUNITY

MASH: 35 million globally. Increasing prevalence. Door opener to even broader anti-aging indication

LONG LIFE IP

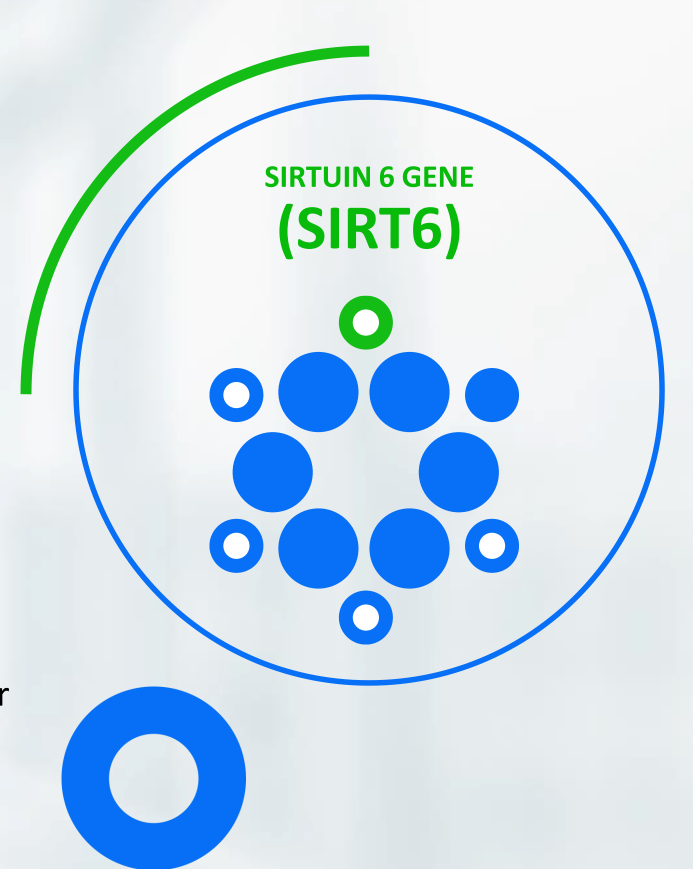
2 patent families SIRT6 centenarian and gene delivery (entering National Phase); Additional upcoming patent applications (entering PCT)

GENE DELIVERY SYSTEMS

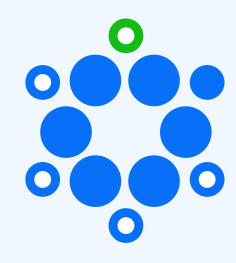
Proprietary innovative gene delivery systems: exo-AAV, mRNA exosomes, LNPs Centenarian variant of SIRT6 gene

GF-1002 and **GF-1003**

Multiple key clinical and regulatory milestones expected in next 18 months Undervalued stock opportunity; Potential acquisition by pharmaceutical partner



MANAGEMENT











TAMARA JOSEPH Chairperson

- Seasoned healthcare leader with extensive experience in both early-stage and commercial biotech companies
- Supported Nasdaq financings of over \$800m
- Currently serving as Chief Legal Officer at Spero Therapeutics Inc. (NASDAQ:SPRO)
- Served as an adviser to the boards of five US publicly traded biotechs, including Cubist Pharmaceuticals Inc.
- BA in Economics from Duke, a JD from the University of Michigan, and LLM degrees from Belgium and the University of Paris



DR ERIC LEIRE MD MBA

Founder & CEO

- MD and MBA, Eric has been involved in biotech for over 30 years
- Held senior positions including CEO of publicly traded biotech companies (Nasdaq, OTC.QB, OMX.Nasdaq)
- Inventor of several patents and author of medical peer-reviewed publications





















SCIENTIFIC ADVISORY BOARD





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Buck Institute

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UCSF School Of Medicine



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Assistant Professor

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Principle Investigator

University of Liverpool

Affiliated With

UCL



PROF. DR. SVEN FRANCQUE, PHD

MASH Expert

University of Antwerp



DR. MARY
RINELLA, MD
MASH Expert

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THANK YOU

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HARBOR ACCESS INVESTOR RELATIONS

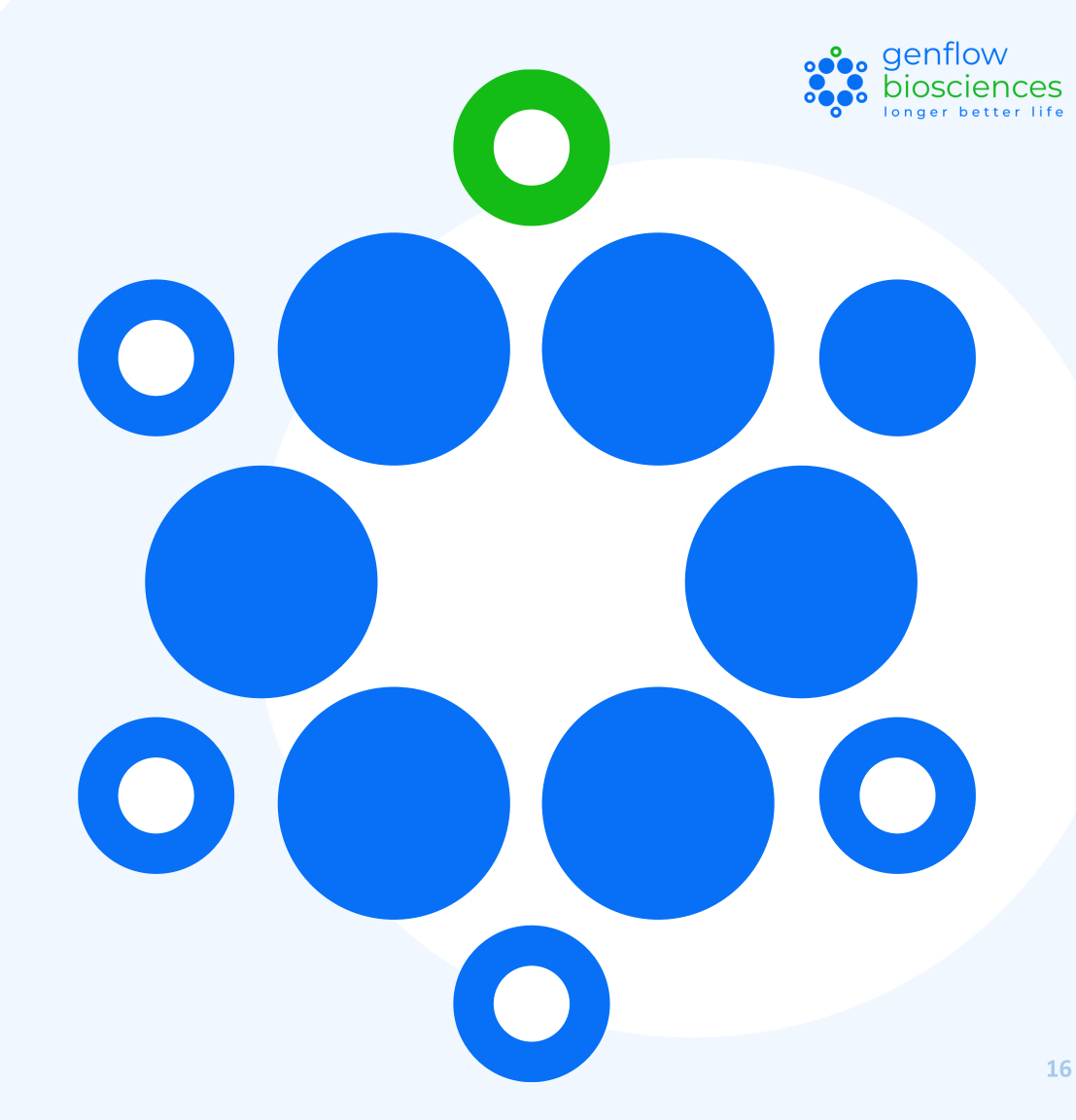
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INTELLECTUAL PROPERTY



EFS ID	1-21069	43268050			
Application Number	US 63/188,573	US 63/222,557			
Title of Invention	Variants of SIRT6 for use in preventing and/or treating age- related diseases	Method of in vivo administration of the coding sequence of the SIRT6 gene via Adeno-Associated-Virus			
First Named Inventor	Vera Gorbunova, Seluanov and Suh	Eric Leire			
Receipt Date	May 14, 2021	July 16, 2021			
Ownership	Worldwide Exclusive license from University Rochester New York / Columbia University / Albert Einstein College of medicine	Genflow Biosciences SRL			

LONGEVITY LANDSCAPE



COMPANY	OVERVIEW	TECHNOLOGY	FOCUSED ON	LOCATION	
UNITY	CLINICAL STAGE, PHASE NASDAQ (UBX) MKT CAP \$	9	mall Molecules Senolytic	Senescence	USA, San Francisco, CA
AGEX	PRE-CLINICAL STAGE NYSE (AGE) MKT CAP \$2	THE G	peutics that seek to ress human aging	Stem cells	USA, Almeda, CA
verve THERAPEUTICS	CLINICAL STAGE, PHASI NASDAQ (VERV) MKT CAP \$		Vivo LNP CRIPR Gene editing	Hypercholesterolemia	USA, Cambridge, MA
FREQUENCY THERAPEUTICS	PRE-CLINICAL STAGE NASDAQ (FREQ) MKT CAP	nro	Molecules to Activate enitor calls for MS	Stem cell exhaustion	USA, Woburn, MA
11fe BIOSCIENCES	PRE-CLINICAL STAGE PRIVATE RAISED \$124	Epiger	etic reprogramming	Mitochondrial dysfunction	USA, Boston, MA
biosplice	CLINICAL STAGE, PHASE PRIVATE RAISED \$778	to deve	ve splicing modulation lop medicines to treat ng-related diseases	Osteoarthritis	USA, San Diego, CA
REJUVENATE BIO	CLINICAL STAGE, PHASI PRIVATE RAISED \$26N		Gene Therapy	Proteostatis	USA, San Carlos, CA