

Disclaimer



This presentation has been prepared by AVROBIO, Inc. ("AVROBIO") for informational purposes only and not for any other purpose. Certain information contained in this presentation and statements made orally during this presentation relate to or are based on studies, publications, surveys and other data obtained from third-party sources and AVROBIO's own internal estimates and research. While AVROBIO believes these third-party sources to be reliable as of the date of this presentation, it has not independently verified, and AVROBIO makes no representation as to the adequacy, fairness, accuracy or completeness of any information obtained from third-party sources. While AVROBIO believes its internal research is reliable, such research has not been verified by any independent source.

This presentation may contain forward-looking statements made pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995. These statements may be identified by words and phrases such as "aims," "anticipates," "believes," "could," "designed to," "estimates," "expects," "forecasts," "goal," "intends," "may," "plans," "possible," "potential," "seeks," "will," and variations of these words and phrases or similar expressions that are intended to identify forward-looking statements. These forward-looking statements include, without limitation, statements regarding our business strategy for and the potential therapeutic benefits of our current and prospective product candidates; the design, commencement, enrollment and timing of ongoing or planned clinical trials and regulatory pathways; the timing of patient recruitment and enrollment activities, clinical trial results, and product approvals; the timing and results of our ongoing preclinical studies; the anticipated benefits of our gene therapy platform including the potential impact on our commercialization activities, timing and likelihood of success; the anticipated benefits and safety profile of busulfan as a conditioning agent; the expected benefits and results of our manufacturing technology, including the implementation of our plato® platform in our clinical trials and gene therapy programs; the expected safety profile of

our investigational gene therapies; the potential impact of the COVID-19 outbreak on our clinical trial programs and business generally, as well as our plans and expectations with respect to the timing and resumption of any development activities that may be temporarily paused as a result of the COVID-19 outbreak; the market opportunity for and anticipated commercial activities relating to our investigational gene therapies; and statements regarding our financial and cash position and expected cash reserves. Any such statements in this presentation that are not statements of historical fact may be deemed to be forward-looking statements.

Any forward-looking statements in this presentation are based on our current expectations, estimates and projections about our industry as well as management's current beliefs and expectations of future events only as of today and are subject to a number of risks and uncertainties that could cause actual results to differ materially and adversely from those set forth in or implied by such forward-looking statements. These risks and uncertainties include, but are not limited to, the risk that any one or more of our product candidates will not be successfully developed or commercialized; the risk of cessation or delay of any ongoing or planned clinical trials of AVROBIO or our collaborators; the risk that we may not successfully recruit or enroll a sufficient number of patients for our clinical trials; the risk that we may not realize the intended benefits of our gene therapy platform, including the features of our plato[®] platform; the risk that our product candidates or procedures in connection with the administration thereof, including our use of busulfan as a conditioning agent, will not have the safety or efficacy profile that we anticipate; the risk that prior results, such as signals of safety, activity or durability of effect, observed from preclinical or clinical trials, will not be replicated or will not continue in ongoing or future studies or trials involving our product candidates: the risk that we will be unable to obtain and maintain regulatory approval for our product candidates; the risk that the size and growth potential of the market for our product candidates

will not materialize as expected; risks associated with our dependence on third-party suppliers and manufacturers; risks regarding the accuracy of our estimates of expenses and future revenue; risks relating to our capital requirements and needs for additional financing; risks relating to clinical trial and business interruptions resulting from the COVID-19 outbreak or similar public health crises, including that such interruptions may materially delay our development timeline and/or increase our development costs or that data collection efforts may be impaired or otherwise impacted by such crises; and risks relating to our ability to obtain and maintain intellectual property protection for our product candidates. For a discussion of these and other risks and uncertainties, and other important factors, any of which could cause AVROBIO's actual results to differ from those contained in the forward-looking statements, see the section entitled "Risk Factors" in AVROBIO's most recent Annual Report on Form 10-K, as well as discussions of potential risks, uncertainties and other important factors in AVROBIO's subsequent filings with the Securities and Exchange Commission. AVROBIO explicitly disclaims any obligation to update any forward-looking statements except to the extent required by law.

Note regarding trademarks: plato[®] is a registered trademark of AVROBIO. Other trademarks referenced in this presentation are the property of their respective owners.

Note regarding future updates: The statements contained in this presentation reflect our current views with respect to future events, which may change significantly as the global consequences of the COVID-19 pandemic rapidly develop. Accordingly, we do not undertake and specifically disclaim any obligation to update any forward-looking statements.

Copyright© 2021 AVROBIO, Inc. All rights reserved.





Leading lysosomal disorder gene therapy pipeline



14 patients dosed to date across three indications

	Proof-of-Concept	IND-Enabling	Phase 1/2
Fabry AVR-RD-01			
Gaucher type 1 AVR-RD-02			
Cystinosis AVR-RD-04			
Hunter AVR-RD-05			
Gaucher type 3 AVR-RD-06			
Pompe AVR-RD-03			



Multi-billion dollar market opportunity

+

Over 50,000 patients across target indications

Disease	Approx. 2020 Global Net Sales [†]	Five-Year SOC Cost per U.S. Patient*	Selected Companies w/ Marketed Therapies
Fabry	\$1.4B	\$1.7M	SANOFI GENZYME Shire
Cystinosis	\$0.2B	\$4.3M	#HORIZON [‡]
Gaucher	\$1.5B	\$2.3M	SANOFI GENZYME Shire
Hunter	\$0.6B	\$2.4M	Takeda Shire
Pompe	\$1.1B	\$3.2M	SANOFI GENZYME 🎝
	Total: \$4.0D		

Total: \$4.8B

Sources: Rombach S et al., Orphanet J Rare Dis, 2013; van Dussen L et al., Orphanet J Rare Dis, 2014

Note: Shire acquired by Takeda in 2019

SOC: Standard of Care



^{*} WAC pricing from Redbook using standard dosing assumptions

^{† 2020} Net Sales from company annual and other reports

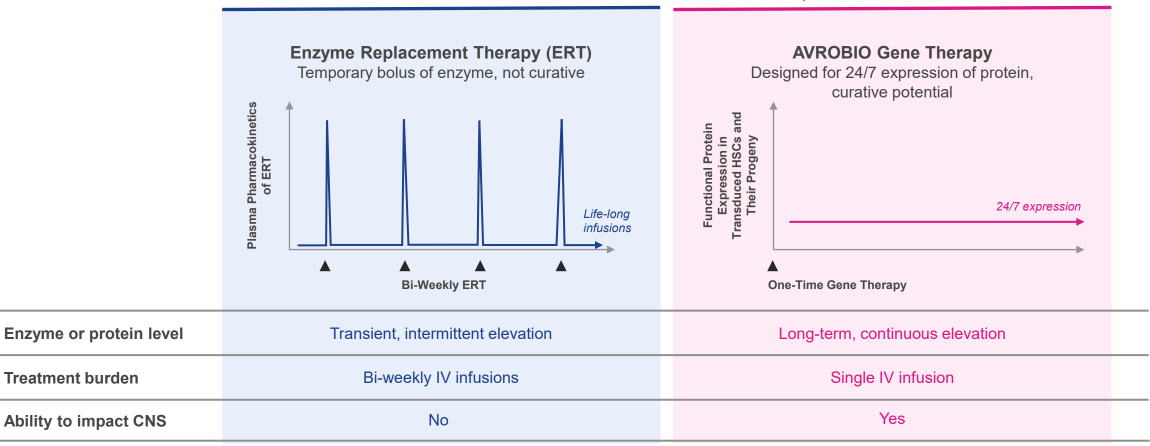
[‡] Horizon's Procysbi oral therapy (delayed release cysteamine bitartrate); midpoint between avg. adult and pediatric

Lifelong treatments vs. potential single-dose therapy



DISEASE PROGRESSION CONTINUES

COULD HALT, PREVENT OR REVERSE DISEASE

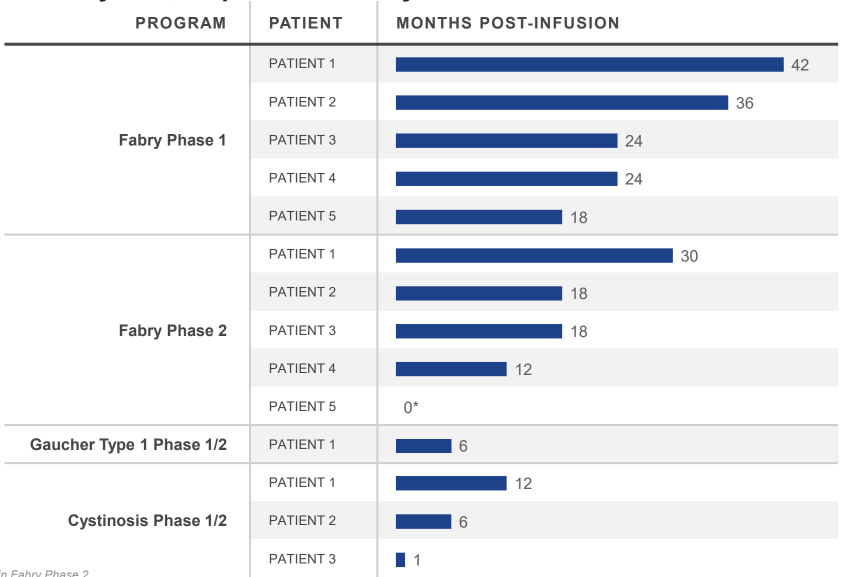




Durability demonstrated across clinical programs



First patient out 3.5 years; 10 patients out 1 year or more

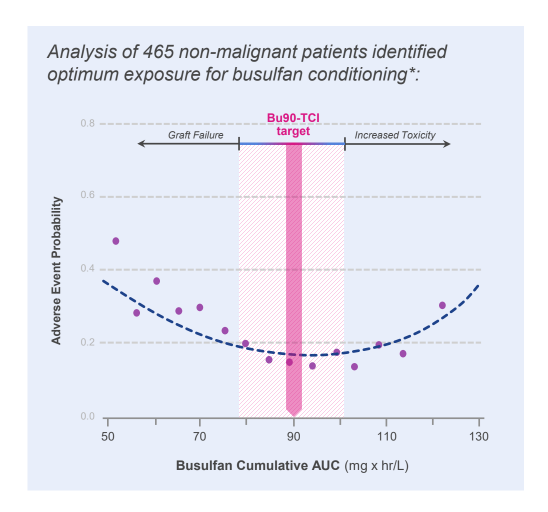


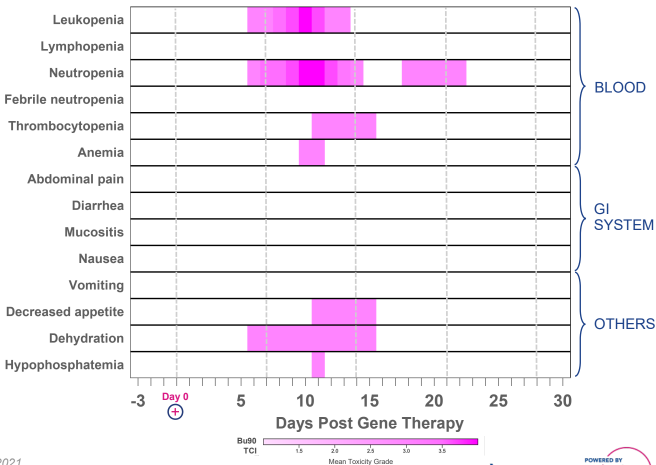


Bu90-TCI conditioning-related side effects have been predictable and transient in first two plato® patients



Conditioning-related grade 3-4 AEs in first two plato® patients





Note: FAB-GT, f-k-a FAB-201, safety data cut-off December 7, 2020; Gaucher safety data cut-off January 4, 2021

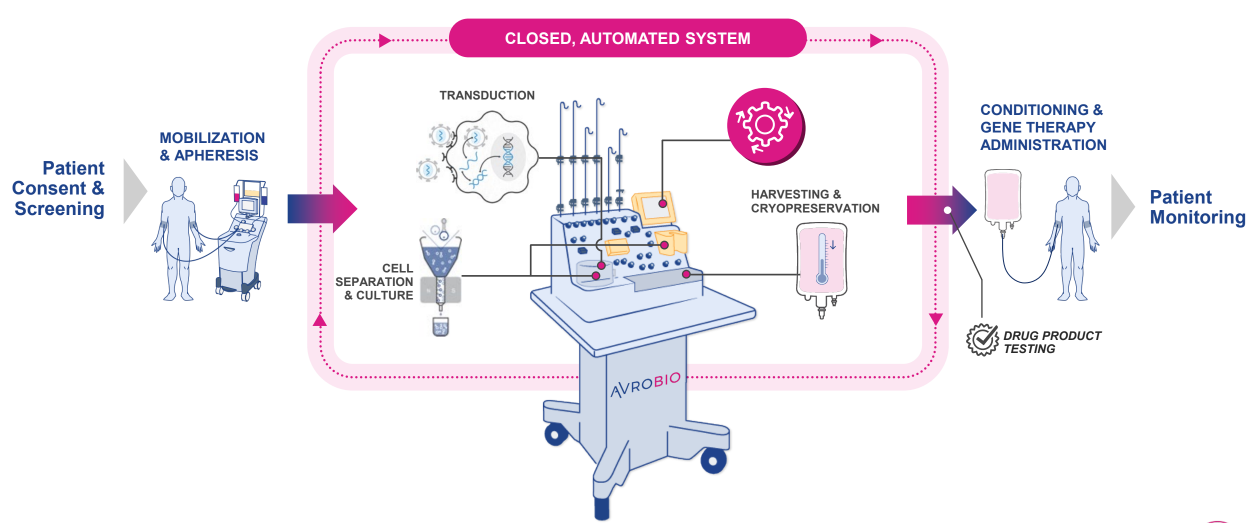
* Source: Bartelink IH et al., Lancet Haematol, 2016

Bu90-TCI: Busulfan 90-Target Concentration Intervention; AUC: Area Under the Curve; TCI: Target Concentration Intervention



Unrivaled commercial-scale platform in plato®





"First Wave" Programs

Fabry, Gaucher Type 1, cystinosis

Fabry disease opportunity



Caused by mutation in gene encoding for alpha-galactosidase A enzyme

Standard of care (SOC): ERT

- Not curative, relentless progression of disease continues
- Burdensome and expensive bi-weekly ERT infusions required; 5-year treatment cost of ERT = ~\$1.7 million*

Unmet needs with SOC:



Kidney function

Proteinuria, polyuria, kidney failure



Cardiac function

Left ventricular hypertrophy, fibrosis, heart failure



Neuropathic pain

Pain and burning sensations in hands and feet, pain crises



Everyday burden of illness, and life expectancy

Not curative, relentless progression of disease, shortened lifespan



CNS complications

TIA/stroke, depression, executive function deficit, white matter lesions

Fabry Disease Target Product Profile:**

- Prevents, halts or reverses disease; extends/normalizes lifespan
- Addresses all patient segments all genetic mutations, male and female, all ages
- Lifelong durability single infusion; off ERT
- Impacts hard-to-reach organs e.g., brain, heart, kidney
- Well tolerated

Affects ~ 1:40,000 males and 1:118,000 females in U.S.



^{*} WAC pricing from Redbook using standard dosing assumptions

^{**} Note: these are target attributes for a first-line therapy

Two AVR-RD-01 Fabry clinical trials



10 patients dosed across Phase 1 and 2



PHASE 1

Investigator-Sponsored Trial*

FULLY ENROLLED





PHASE 2

AVROBIO FAB-GT Trial **

ACTIVELY RECRUITING







OBJECTIVES

Safety and tolerability

Preliminary efficacy

PATIENTS

- n = 5 patients
- 18 59 year-old males
- On ERT >6 months prior to enrollment

OBJECTIVES

- Safety and tolerability
- Efficacy

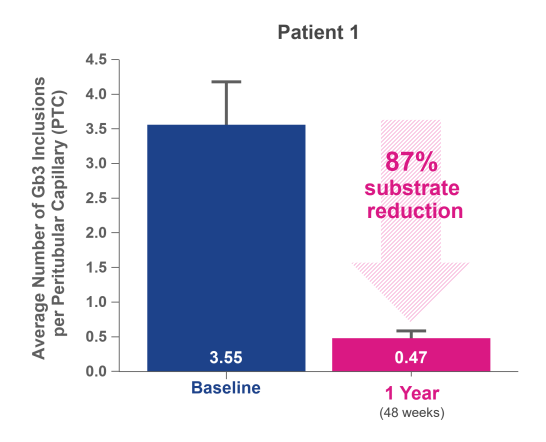
PATIENTS

- n = 8-12 patients (5 dosed to-date)
- 16 50 year-old males
- Treatment naïve



^{*} Sponsored by FACTs team (Fabry Disease Clinical Research and Therapeutics) in Canada ** FAB-GT fka FAB-201

Clinically meaningful and statistically significant reduction in substrate in first two evaluable kidney biopsies





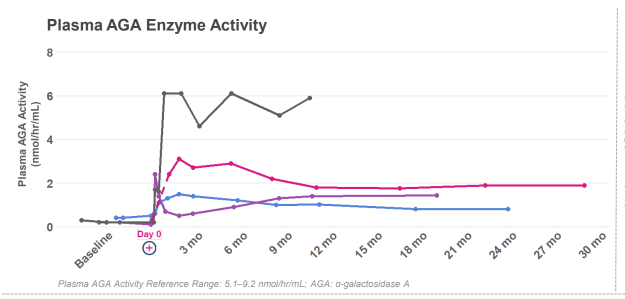
Two-sample t-test for difference between average PTCs at Baseline vs. 48 weeks; p < 0.0001; Error bar represents the standard error at Baseline (n=48 PTCs) and 48 weeks (n=101 PTCs). Scored by 2 independent, blinded pathologists

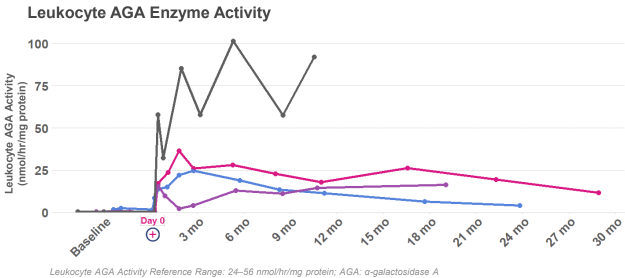
Two-sample t-test for difference between average PTCs at Baseline vs. 48 weeks; p < 0.0001; Error bar represents the standard error at Baseline (n=103 PTCs) and 48 weeks (n=99 PTCs). Scored by 2 independent, blinded pathologists

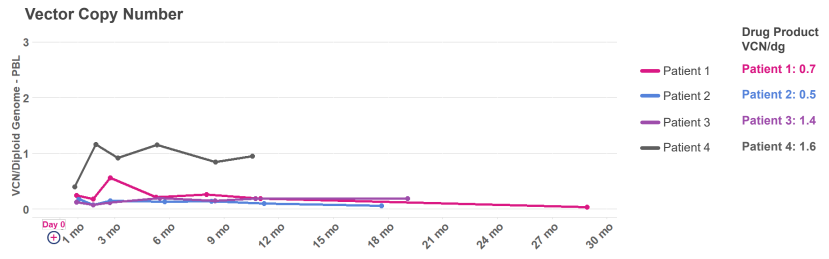


Durability demonstrated over multiple measures up to 2.5 years (+) Patient 4 dosed using plato®



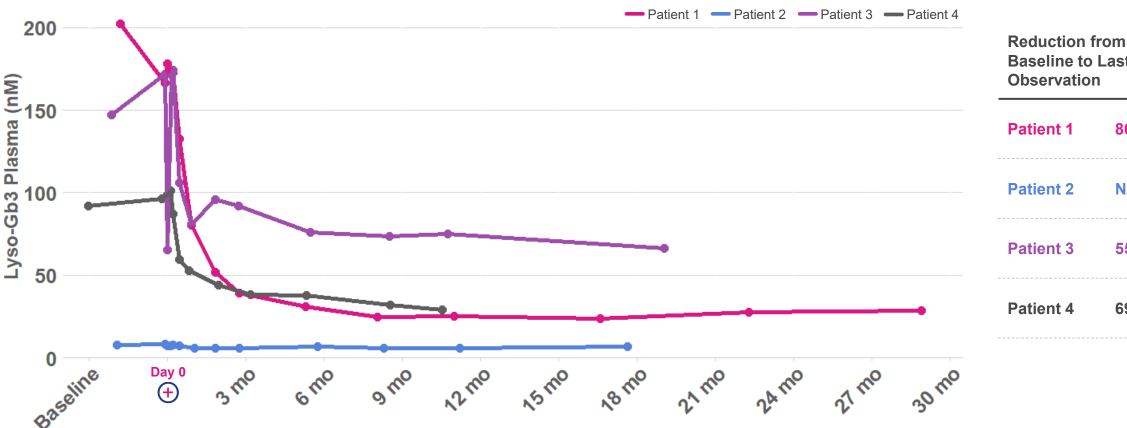






70% average plasma lyso-Gb3 reduction





Baseline to Last Observation		
Patient 1	86%	
Patient 2	N/A	
Patient 3	55%	

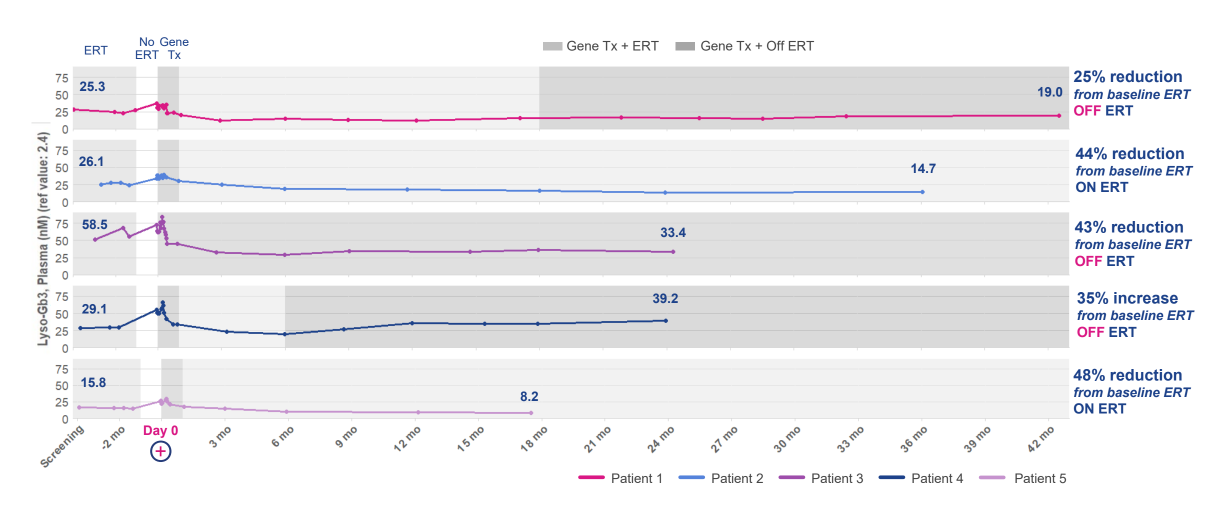


69%

25% average plasma lyso-Gb3 reduction below baseline ERT



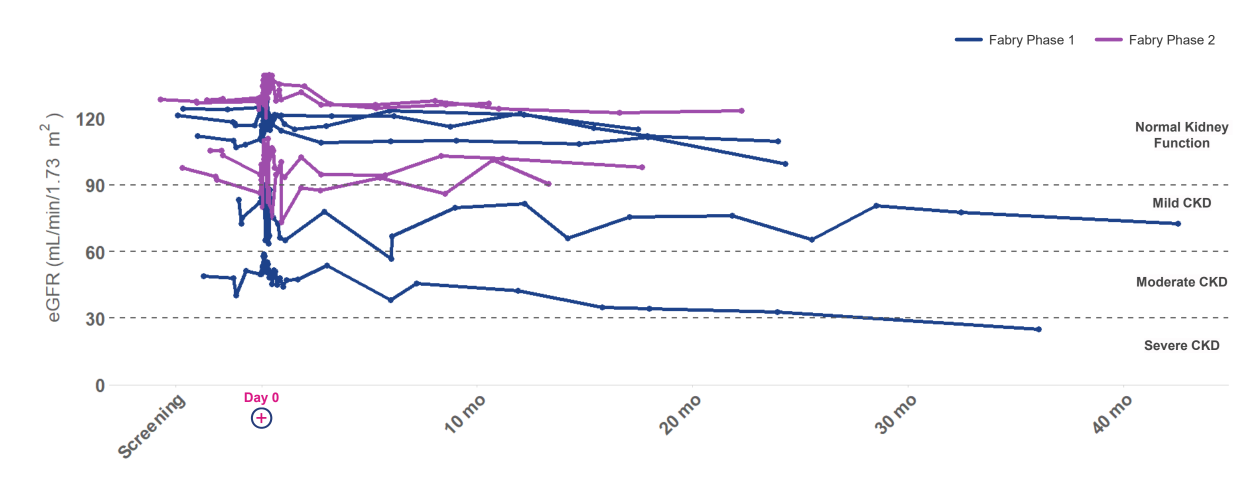
All patients who have discontinued ERT remain off ERT*





Kidney function (eGFR) stable up to 3.5 years*





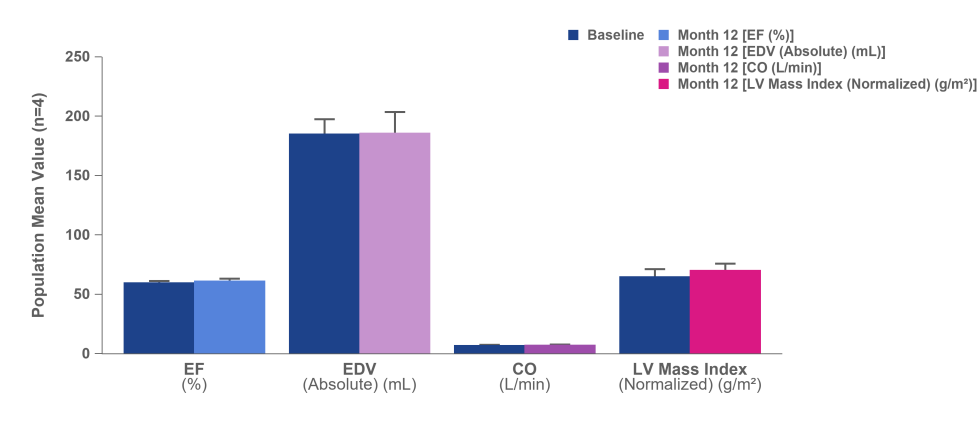
^{*} Eight of nine patients stable; other patient entered trial with more advanced kidney disease and a baseline eGFR level <50 mL/min/1.73m2; as expected, this patient has not stabilized, and the patient remains on ERT

Note: eGFR was calculated using the CKD-EPI formula
eGFR: Estimated Glomerular Filtration Rate; CKD: Chronic Kidney Disease; CKD-EPI: Chronic Kidney Disease Epidemiology Collaboration



Cardiac function and mass stable across multiple measures up to 1 year





Abbreviations: EF=Ejection Fraction; EDV=End Diastolic Volume; LV=Left Ventricular. Error bar represents the standard error of the population mean (n=4).



^{*}Reference Range Mean Values Male 20-39 yrs; EF: 64.3 ± 4.2%; EDV: 178.6 ± 30.1 mL; CO: 4-8 L/min; LV Mass Index: 67.8 ± 10.7 g/m²

^{**}Reference Range Mean Values Male 40-49 yrs; EF: 58-75 %; EDV: 117-200 mL; CO: 4-8 L/min; LV Mass Index: 58-91 g/m²

No unexpected safety events identified

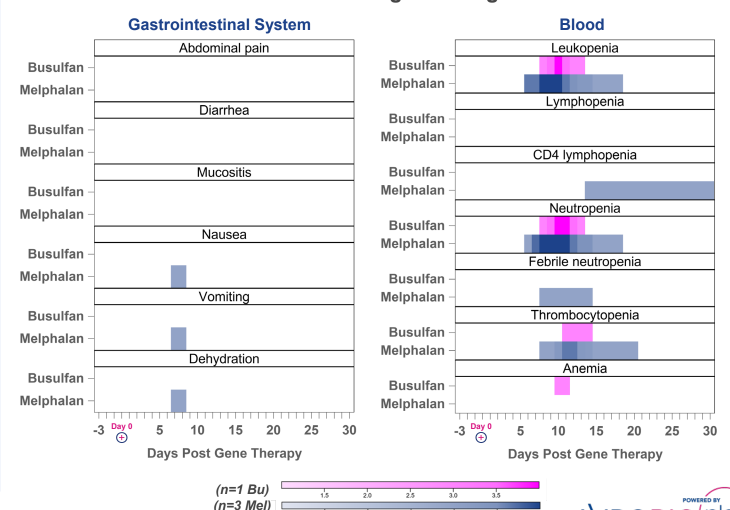


Conditioning-related side effects have been manageable and transient

Phase 1 & 2 AEs and SAEs

- No AEs or SAEs related to AVR-RD-01 drug product
- AEs across trials generally consistent with myeloablative conditioning, underlying disease or pre-existing conditions
- Phase 1 AEs (n=94)
 - Grade 3 or 4 (n=14)
- Phase 1 SAEs (n=2) resolved without clinical sequelae
 - Post-AVR-RD-01 treatment: febrile neutropenia; thrombophlebitis
- Phase 2 AEs (n=111)
 - Grade 3 or 4 (n=22)
- Phase 2 SAEs (n=6) resolved without clinical sequelae
 - Post-AVR-RD-01 treatment: dehydration; nausea; vomiting; febrile neutropenia

Phase 2 conditioning-related grade 3/4 AEs



0 2.5 3 Mean Toxicity Grade

Note: Phase 2 safety data cut-off December 7, 2020; Phase 1 safety data cut-off November 26, 2020 AE: Adverse Event; Bu: Busulfan; Mel: Melphalan

Accelerating enrollment by adding international referrals



ONE Fabry patient from Brazil has been dosed and THREE have been enrolled in Australia







Long-term follow-up expected to take place in Brazil

Global patient recruitment

- Expands pool of potential patients
- Helps navigate COVID-19 issues
- First global center of excellence established in Australia

Cystinosis opportunity



Caused by CTNS gene defect, resulting in cystine buildup in lysosomes

Standard of care (SOC): Cysteamine pills & eye drops

- Not curative, relentless progression of disease continues; significantly shortened lifespan; kidney transplant often required
- Burdensome and expensive high pill burden and hourly eye drops; 5-year treatment cost with SOC ~\$4.3 million*

Unmet needs with SOC:



Kidney function

Renal Fanconi syndrome, proteinuria, CKD, kidney failure



Vision

Corneal cystine accumulation, photophobia, involuntary eyelid closure



Endocrine disorders

Softening & deformation of bones, hypothyroidism, diabetes, infertility



CNS complications

Myopathy, hypotonia, tremors, swallowing, neurodevelopmental issues



Everyday burden of illness, reduced life expectancy

High pill burden causes GI discomfort; sulfur body odor and breath

Cystinosis Target Product Profile:**

- Prevents, halts or reverses disease; extends/normalizes lifespan
- Addresses all patient segments male & female; kidney transplant independent; all ages
- Lifelong durability single infusion; off cysteamine pills and eye drops
- Impacts hard-to-reach organs e.g., eye, endocrine organs, brain
- Well tolerated

Affects ~ 1:170,000 people



^{*} WAC pricing from Redbook using standard dosing assumptions

^{**} Note: these are target attributes for a first-line therapy

Steady enrollment in AVR-RD-04 IST trial in cystinosis





PHASE 1/2

AVR-RD-04

ACTIVELY RECRUITING:

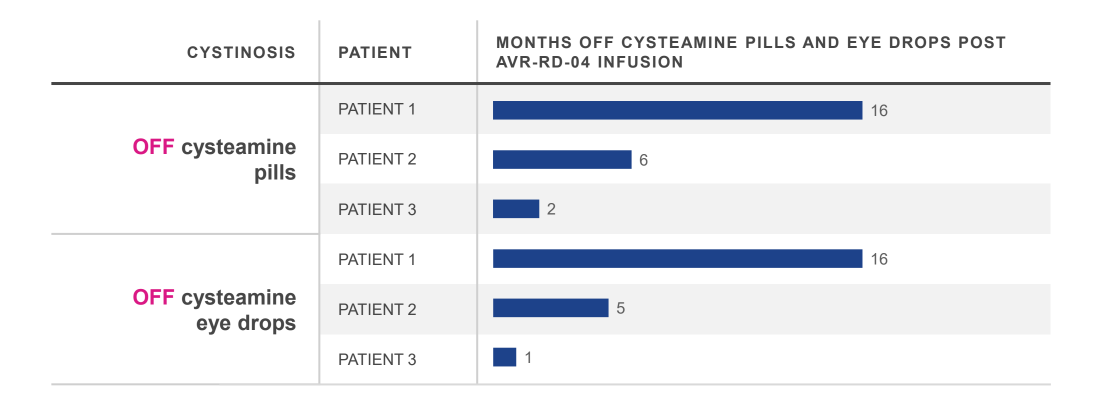


OBJECTIVES	PATIENTS
 Safety and tolerability Hypothesis generation of endpoints 	 Up to 6 patients (3 patients enrolled to-date) Adults and adolescents Cohorts 1-2 >18 years; Cohort 3 >14 years Male and female Oral and ophthalmic cysteamine



All patients continue to be cysteamine-independent

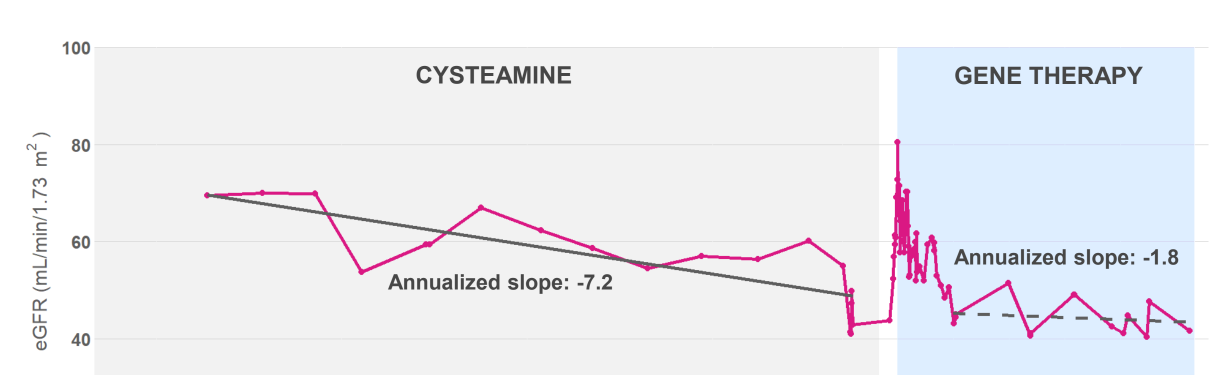






20

eGFR data at 16 months suggest renal function stabilization to post-gene therapy after years of pathological decline



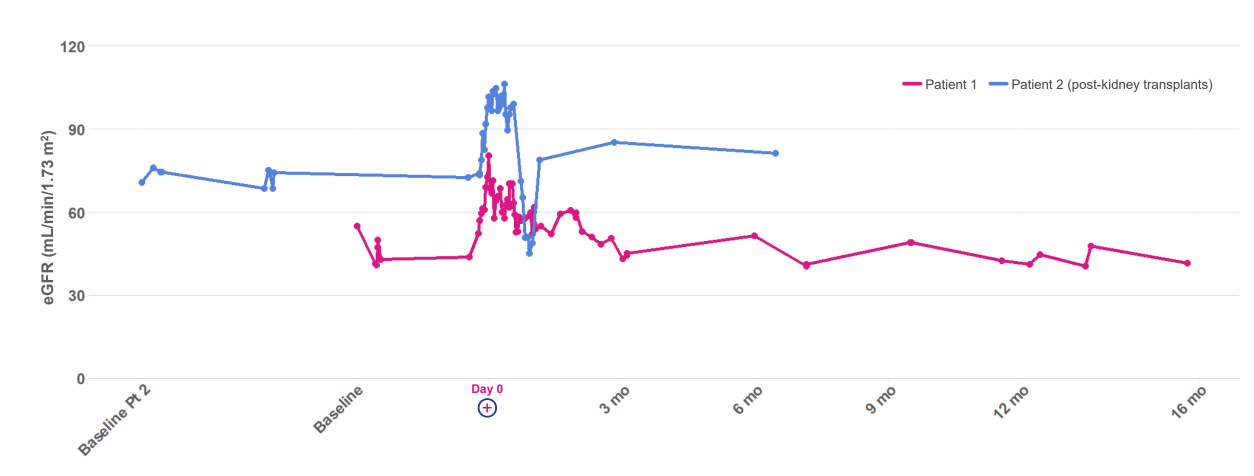
JOMO



+

Trial designed to demonstrate broad applicability across cystinosis patient population

Positive eGFR trends independent of kidney transplant status







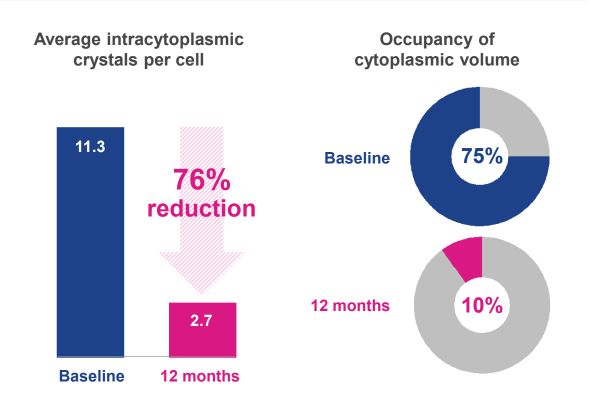
Sharp drop in the number and size of cystine crystals in skin and rectal biopsies



SKIN BIOPSY

Average intracytoplasmic crystals per cell 4.6 440% reduction 2.6 Baseline 12 months Occupancy of cytoplasmic volume 90% 12 months

RECTAL BIOPSY





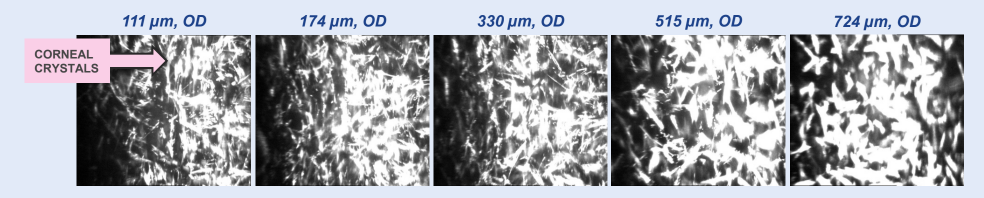
Substantial decline in corneal crystals observed at 1 year



Front of cornea Back of cornea

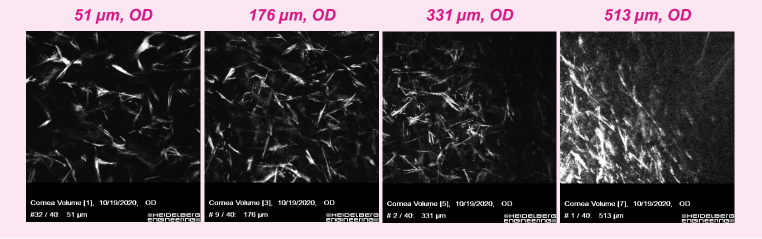
Baseline

IVCM images from Nidek Confoscan



12 months post-gene therapy

IVCM images from Heidelberg HRT3 w/ Rostock Corneal Module





Photophobia improved meaningfully at 1 year

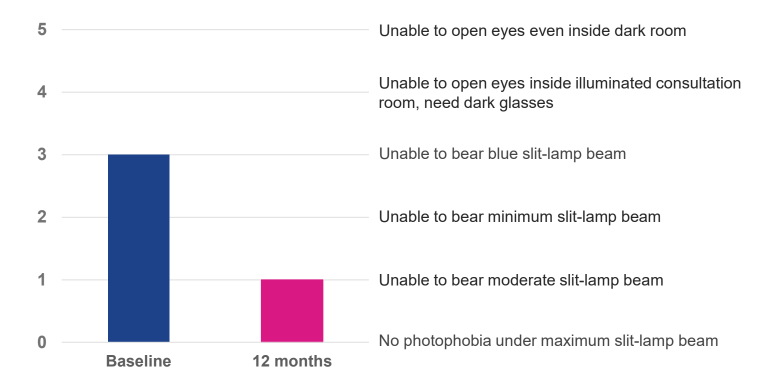


Photophobia, or extreme sensitivity to light, is a hallmark of cystinosis

Cystinosis photophobia intensity associated with:

- Crystal density (light scattering)
- Inflammatory cell infiltration
- Corneal nerve damage

Clinician-Assessed Photophobia Grade (Patient 1)





Darker pigmentation may be a sign of multi-functional cystinosin activity post-gene therapy



Cystinosin is located in melanosomes and regulates melanin synthesis

Patient 1 appears to exhibit progressively darkening skin, eyebrows and hair color post-infusion, suggesting a possible impact of cystinosin protein on melanin





VCN trending as expected across patients



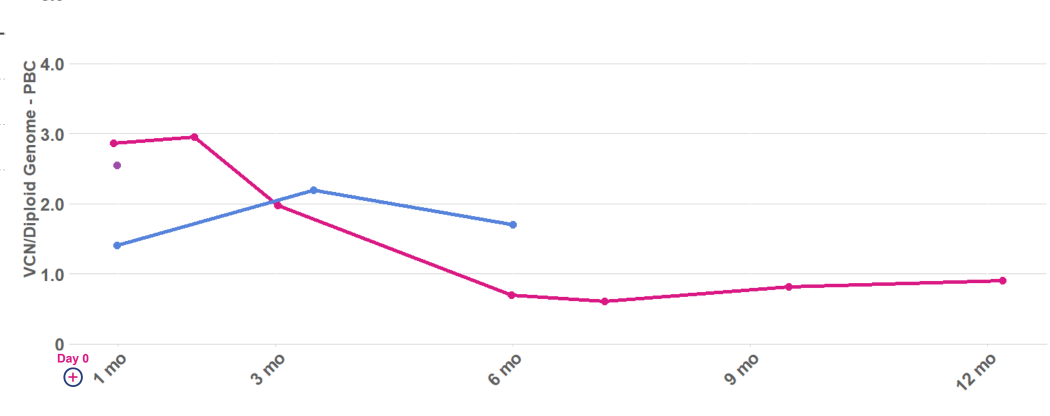
Patient 1 reached VCN therapeutic plateau

5.0



Patient 2 1.3*

Patient 3 1.6



AVROBIO Plate

- Patient 1 - Patient 2 - Patient 3

No unexpected safety events



Conditioning-related side effects have been manageable and transient

No SAEs or AEs related to AVR-RD-04 drug product

AEs & SAEs reported

- AEs (n=48)
 - Majority of AEs are mild or moderate and resolved
- SAE (n=1)
 - Post AVR-RD-04 treatment: appendicitis unrelated to study treatment or procedures

 AEs are generally consistent with myeloablative conditioning or underlying disease:

Pre-AVR-RD-04 treatment and prior to conditioning (not all events listed)

- Diarrhea, hypokalemia, dizziness
- Dehydration, vomiting

Post-AVR-RD-04 treatment (not all events listed)

- Alopecia, intermittent diarrhea, vomiting, loss of appetite
- Mucositis, intermittent febrile neutropenia, intermittent epistaxis
- Intermittent blurry vision, intermittent hypokalemia, mucoceles
- Thrombocytopenia



Planned global regulatory strategy for cystinosis

Planned

POTENTIAL REGISTRATION

- · Adults and pediatrics, males and females
- Mutation-independent, kidney transplant-independent
- · Efficacy, durability, safety
- · Ophthalmology, kidney, and other undisclosed
- Multiple crystal measures
- Quality of life

50% Enrolled

PHASE 1/2 - INVESTIGATOR SPONSORED TRIAL

- n ≤6
- Adults and adolescents, males and females
- Mutation-independent, kidney transplant-independent
- Safety, durability, preliminary efficacy
- Biomarker data, kidney function, vision
- · Quality of life

Anticipated Next Steps:

- Complete Phase 1/2 enrollment in 2021
- Engage with FDA on registration trial design
- Identify global sites for registration trial
- Prepare plato[®] CMC / analytics requirements

Gaucher disease type 1 opportunity



Caused by mutation in the gene encoding for glucocerebrosidase (GCase) enzyme

Standard of care (SOC): ERT

- Not curative, relentless progression of disease continues, including bone crisis and fatigue
- Burdensome and expensive bi-weekly infusions required; 5-year treatment cost with ERT = ~\$2.3 million*

Unmet needs with SOC:



Bone-related manifestations

Skeletal abnormalities, avascular necrosis, osteoporosis



Hemoglobin levels and platelet counts

Anemia, thrombocytopenia, easy bruising, bleeding



Hepatosplenomegaly

Enlarged liver, enlarged spleen



Everyday burden of illness, and life expectancy

Fatigue, pain, lung disease, biweekly infusions, shortened lifespan



CNS complications

Increased risk of GBA-Parkinson's disease

Gaucher Disease Type 1 Target Product Profile**:

- Prevents, halts or reverses disease; extends/normalizes lifespan
- Addresses all patient segments all GD1 genetic mutations, all ages, male & female
- Lifelong durability single infusion; off ERT/chaperone therapy
- Impacts hard-to-reach organs e.g., brain, bone and bone marrow
- Well tolerated

Affects ~ 1:44,000 people worldwide

^{*} WAC pricing from Redbook using standard dosing assumptions

^{**} Note: these are target attributes for a first-line therapy

Guard1: Phase 1/2 study in Gaucher disease type 1





PHASE 1/2

AVR-RD-02

An adaptive, open-label, multinational phase 1/2 study of the safety and efficacy of ex vivo, lentiviral vector-mediated gene therapy AVR-RD-02 for patients with Gaucher disease type 1

ACTIVELY RECRUITING:





RECRUITING PLANNED 2021:





OBJECTIVES

- Safety
- Efficacy
- Engraftment

PATIENTS

- Enrollment goal 8-16 patients
- 18-45-year-old males and females
- Have a confirmed diagnosis of GD1 based on:
 - Deficient glucocerebrosidase enzyme activity
 - Clinical features consistent with GD1

Gaucher disease type 1 patients who are:

- ERT-stable for >24 months *or*
- Treatment-naïve or
- Have not received ERT or SRT in the last 12 months

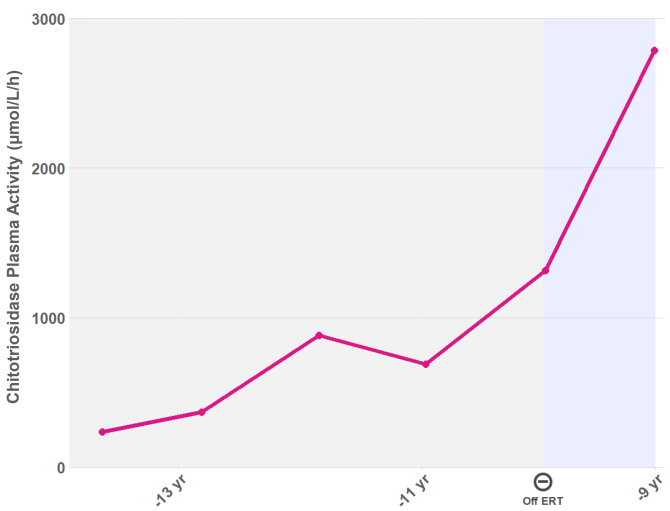


First patient's plasma chitotriosidase levels spike off ERT



Personal history documents response to intermittent and halted ERT use

Chitotriosidase is a marker of inappropriately activated macrophages (Gaucher cells)

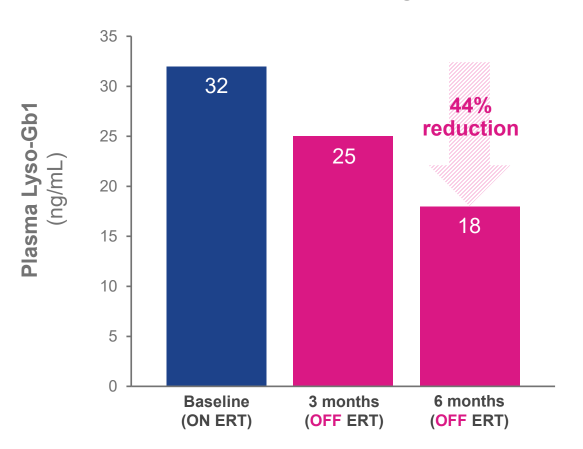




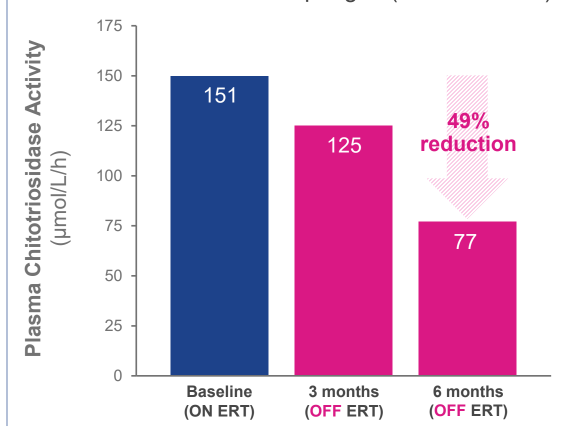
Key biomarkers below ERT baseline at 6 months



Lyso-Gb1 is a sensitive and specific marker of toxic metabolite accumulation in Gaucher disease



Chitotriosidase is a marker of inappropriately activated macrophages (Gaucher cells)



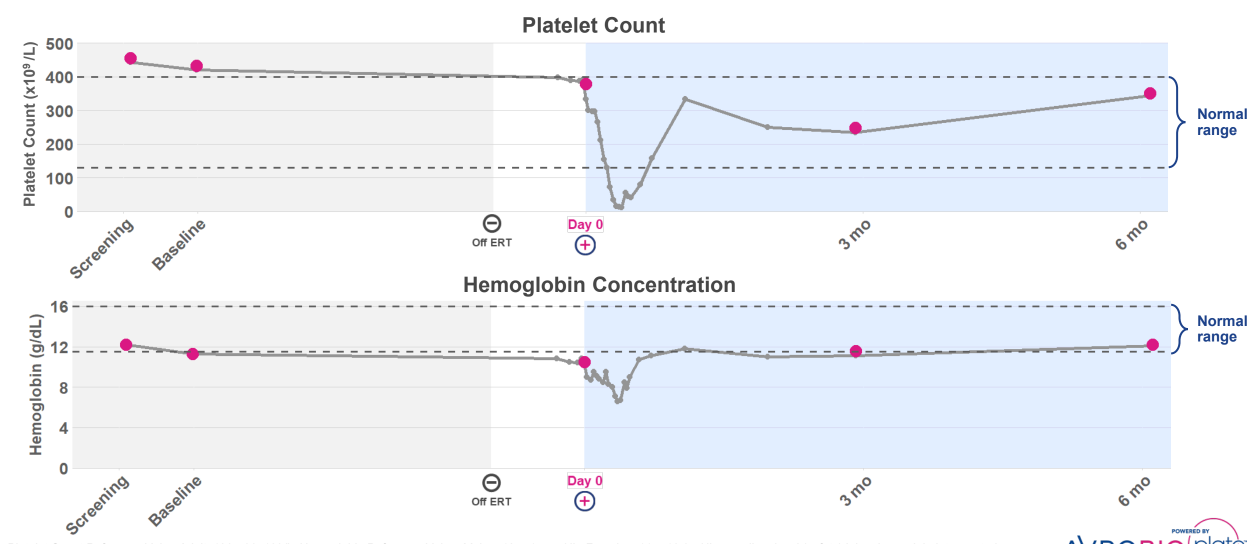
Baseline taken one month prior to gene therapy which is when ERT is discontinued Lyso-Gb1 Plasma Normal Range: 0.5 - 1.2 ng/mL Plasma chitotriosidase activity normal range: 0.0 – 44.2 µmoL/L/h

ERT: Enzyme Replacement Therapy



(+)

Platelet counts and hemoglobin in normal range at 6 months, despite being off ERT



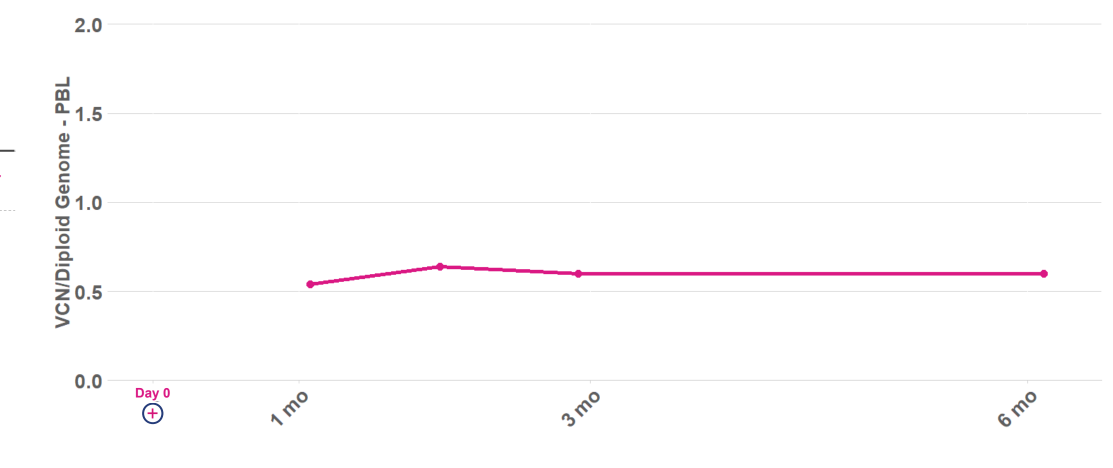
Platelet Count Reference Value Adult: 130-400x109/L; Hemoglobin Reference Value: Males: 13.5-17.5 g/dL; Females: 11.5-16.0 g/dL; grey line: local (safety) lab values; pink dots: central (efficacy) lab values; ERT: Enzyme Replacement Therapy

VCN trending as expected at 6 months





Patient 1 3.7





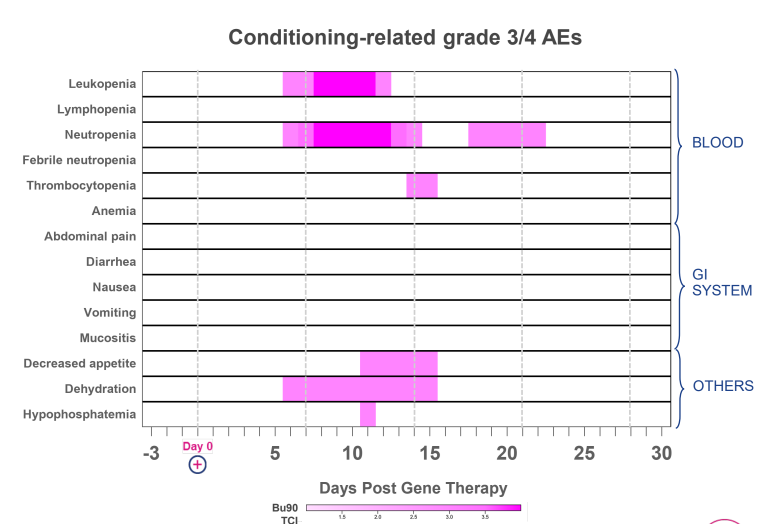
No unexpected safety events identified in first patient



Conditioning-related side effects have been predictable and transient

AEs (no SAEs reported)

- No AEs or SAEs related to AVR-RD-02 drug product
- AEs generally consistent with myeloablative conditioning, underlying disease or pre-existing conditions
- AEs n=29
 - Grade 3 (n=7)
 - Eye pain, decreased appetite, dehydration, headache, hypophosphatemia, neutropenia, thrombocytopenia
 - Grade 4 (n=2)
 - Leukopenia and neutropenia
- AEs resolved without clinical sequelae



Mean Toxicity Grade

Note: Safety database cut as of January 04, 2021

AE: Adverse Event; SAE: Serious Adverse Event; G-CSF: Granulocyte Colony Stimulating Factor G-CSF 5 μg/kg @ Days 5, 6, 7, 10, 11, and 14 post-infusion of AVR-RD-02

Bu90-TCI: Busulfan 90-Target Concentration Intervention; GI: Gastrointestinal



Planned global development strategy for Gaucher disease type 1

Planned

POTENTIAL REGISTRATION PATH

- Phase 1/2 expansion
- · Safety, efficacy, durability
- Organ volumes, hematologic measures, bone assessments, pain, and QOL

Enrolling

PHASE 1/2

- n=8-16
- Adults, males and females, ages 18-45 years old
- ERT-switch and ERT-naïve
- · Safety, efficacy, durability
- Biomarker data, organ volumes, hematologic measures, bone assessments, pain, and QOL

Anticipated Next Steps:

- Advance patient enrollment
- Advance regulatory dialogue on registration pathway

QOL: Quality Of Life; ERT: Enzyme Replacement Therapy



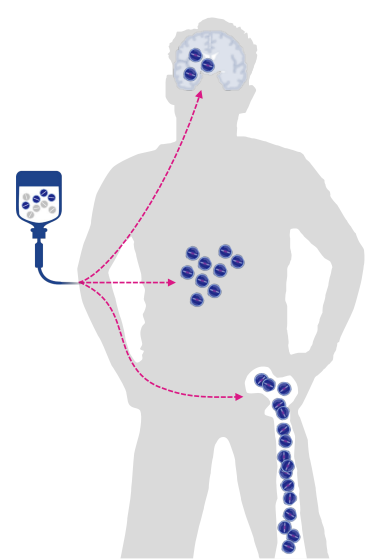
Bold expansion of our leadership in lysosomal disorders

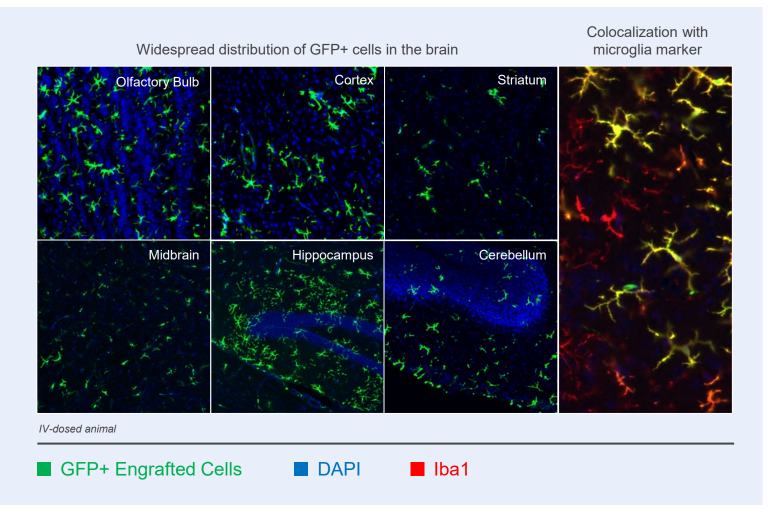


	Proof-of-Concept	IND-Enabling	Phase 1/2
Fabry AVR-RD-01			
Gaucher type 1 AVR-RD-02			
Cystinosis AVR-RD-04			
Hunter AVR-RD-05			
Gaucher type 3 AVR-RD-06			
Pompe AVR-RD-03			

Lentiviral gene therapy enables global distribution of functional enzyme to brain and bone in preclinical studies







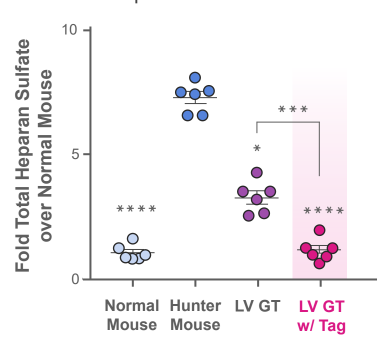


Proprietary tags deliver therapeutic protein into hard-to-reach organs



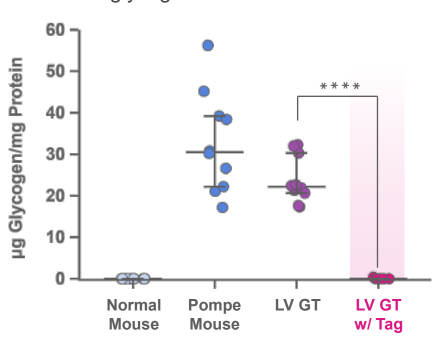
Hunter syndrome

Tag normalizes heparan sulfate in brain



Pompe disease

Tag normalizes glycogen substrate in brain







plato®

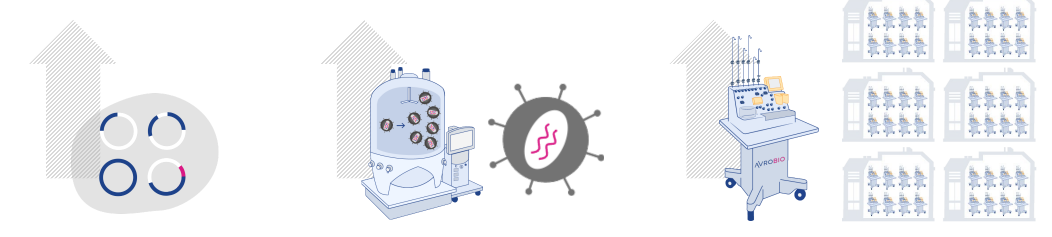
AVROBIO's platform for global gene therapy commercialization

- Redefines manufacturing best practices
- + Solves key industry challenges

Designed to be fully scalable



Common components and automation leveraged across manufacturing



PLASMID

3 of 4 component plasmids used in every vector

Each plasmid can be mass produced and stored for use

VECTOR

State of the art, largest scale, 200L vector production

Expansion through simple installation of additional units, mass produced, frozen, and stored for use

DRUG PRODUCT

Closed system automated platform

Scale out of manufacturing suites and automation units to meet patient demand



Poised to manufacture at scale

(+)

Global infrastructure already in place

VECTOR 2,400 PATIENTS ANNUALLY >50 patients per run 12 runs per year per suite (200L scale bioreactor runs (109 titer)) **4** production suites

DRUG PRODUCT

2,400 PATIENTS ANNUALLY



100 patients per unit per year

8 automated units per suite

3+ global production suites









CMC achievements have defined the plato® story



Strategic investment in technology laid the foundation for our manufacturing leadership

Manufacturing

Robust production platform

- Best-in-class LV manufacturing
- Scalable from plasmid to drug product

Global footprint

In the clinic in multiple jurisdictions

Cost effective

Intended to address key COGs issues

Analytics

Robust platform analytics

- Best-in-class VCN assay
- First-in-class transduction assay

Deep product characterization

First-in-class single cell analytics

Potency assay matrix

Intended to accelerate regulatory approvals



Key anticipated 2021 milestones



Goal:
30 patients
dosed
cumulatively
by end of
2021

Fabry AVR-RD-01 Seek agreement with regulators on approval pathway in one or more major markets

Gaucher type 1
AVR-RD-02

Execute on global phase 1/2 trial

Cystinosis AVR-RD-04 Complete phase 1/2 enrollment Engage w/ FDA on pivotal trial design

Hunter AVR-RD-05

Conduct Phase 1/2 trial initiation activities

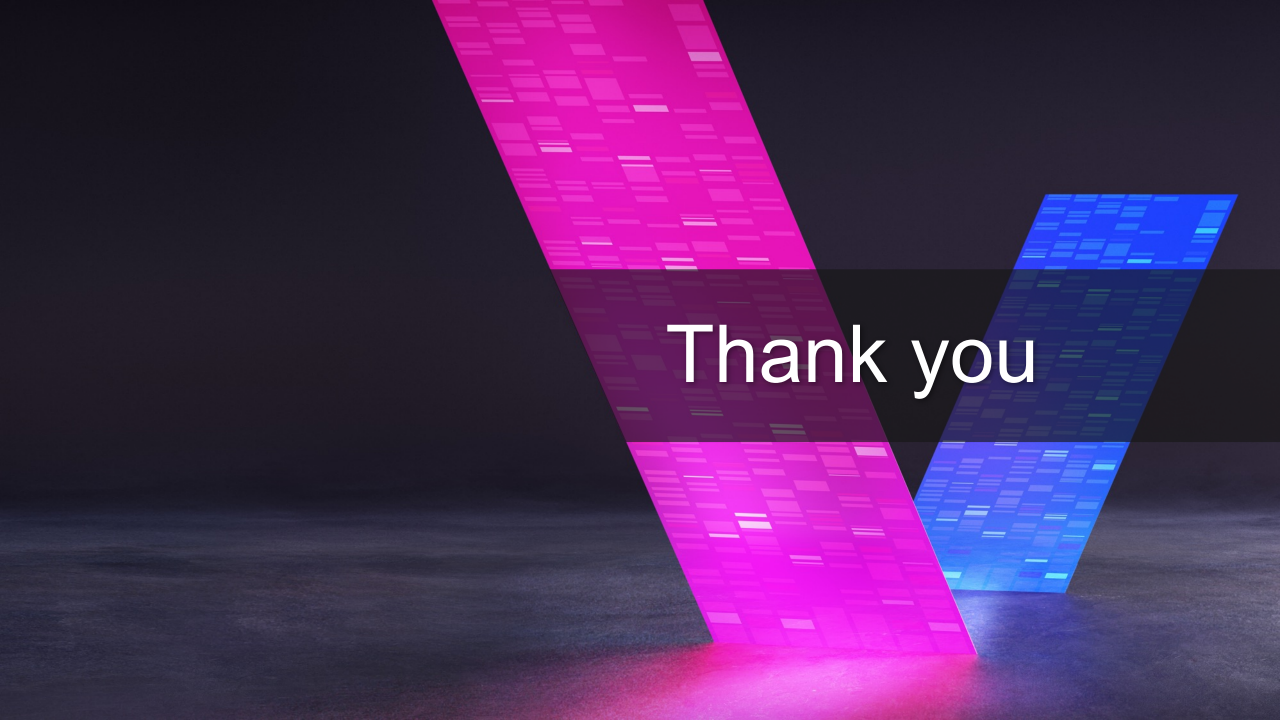
Gaucher type 3
AVR-RD-06

FDA dialogue on path to clinic

Pompe AVR-RD-03

Prepare for classic infantile-onset study







Appendix





Zero cases reported outside of sickle cell disease

SICKLE CELL DISEASE (SCD)

2 or 3 cases out of 47 patients

NON-SCD MONOGENIC DISEASES

0 cases out of >300 patients

CAR-T

0 cases out of >1,000 patients



Fabry Phase 1 & 2 Patient Characteristics



i						
	PHASE 1: ERT-Treated Fabry Patients					
	PATIENT 1	PATIENT 2	PATIENT 3	PATIENT 4	PATIENT 5	
Age of symptom onset / diagnosis	18 / 37 years	9 / 29 years	10 / 0 years	7 / 4 years	10 / 14 years	
Years on ERT	11 years	6 years	4 years	11 years	2 years	
Age dosed with AVR-RD-01	48 years	39 years	40 years	37 years	30 years	
Mutation	c.962A>G (p.Q321R)	c.1033T>C (p.S345P)	c.427G>C (p.A143P)	c.427G>C (p.A143P)	(p.Y134S)	
Leukocyte AGA activity at baseline (nmol/hr/mg protein)**	2.1	1.1	0.6	2.2	1.0	
Plasma lyso- Gb3 at baseline (nM)***	25	26	59	29	16	
eGFR (mL/min/1.73m²) at baseline****	83	49	112	124	121	
ERT discontinuation status	18 months after gene therapy dose		Did not resume ERT after gene therapy dose	6 months after gene therapy dose		

	PHASE 2: Treatment-naïve Fabry patients					
	PATIENT 1	PATIENT 2	PATIENT 3	PATIENT 4		
Age of symptom onset/diagnosis	10 / 19 years	36 / 37 years	13 / 13 years	9 / 9 years		
Age dosed with AVR-RD-01	21 years	46 years	40 years	26 years		
Mutation	c.1021G>A (p.E341K)	c.644A>G (p.N215S)	c.639+1G>T	c.833dupA		
Leukocyte AGA enzyme activity at baseline (nmol/hr/mg protein)	0.10*	2.38**	0.58**	0.46**		
Plasma lyso- Gb3 at baseline (nM)***	202	8	147	92		
eGFR (mL/min/1.73m²) at baseline****	128	106	98	129		
Comment	Few IgA deposits in kidney biopsy, no mesangial proliferation	Cardiac variant, not a classic Fabry male				

^{*} Mayo Lab, ref range ≥23.1 nmol/hr/mg protein; ** Rupar Lab, ref range 24-56 nmol/hr/mg protein; *** Reference value ≤ 2.4 nM; **** eGFR: Estimated Glomerular Filtration Rate; calculated using CKD-EPI formula

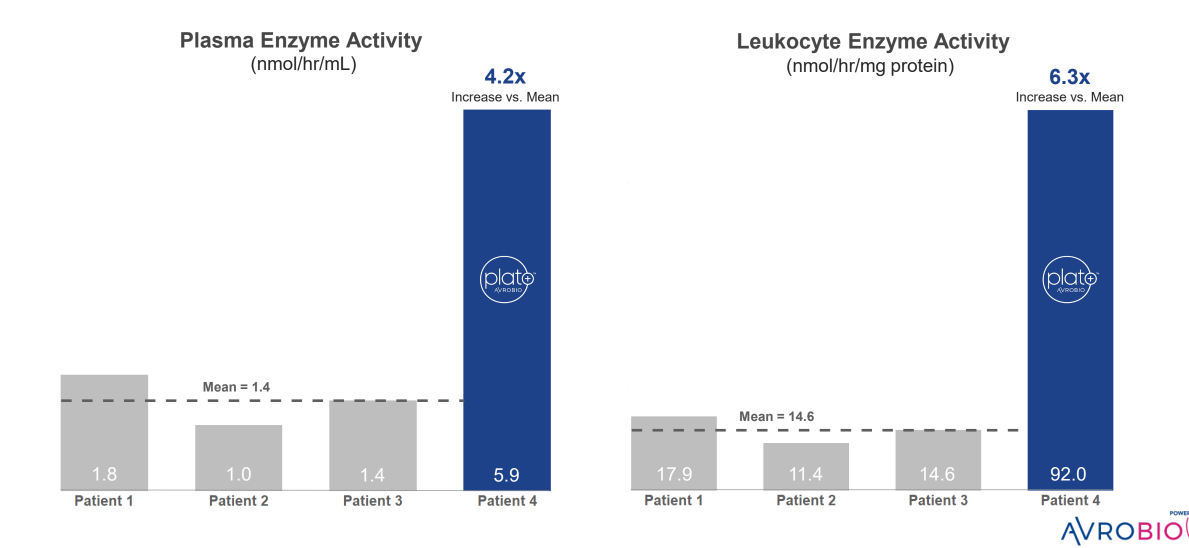
AGA: α-galactosidase A; Lyso-Gb3: Globotriaosylsphingosine;



Patient #4 is first Fabry patient dosed with plato®

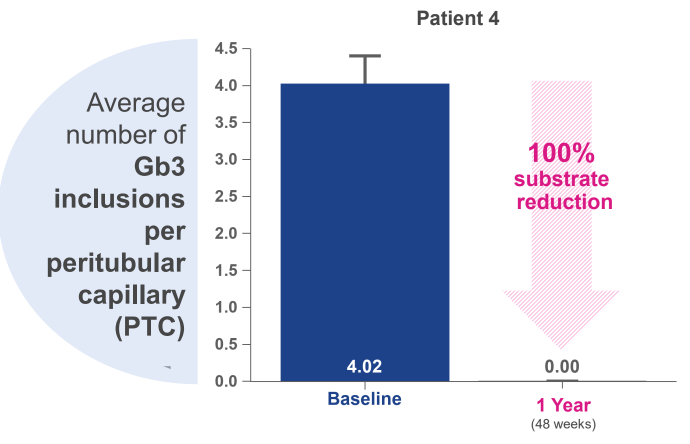


FAB-GT 12 month data for patient #4 with plato® vs. patients #1-3

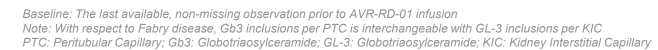


+

100% clearance of substrate in kidney biopsy at 1 year Patient dosed using plato®

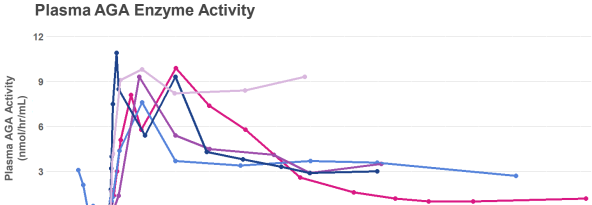


 Two-sample t-test for difference between average PTCs at Baseline vs. 48 weeks; p < 0.0001; Error bar represents the standard error at Baseline (n=103 PTCs) and 48 weeks (n=99 PTCs); scored by 2 independent, blinded pathologists



Durability demonstrated over multiple measures up to 3.5 years





Vector Copy Number

300 Leukocyte AGA Activity (nmol/hr/mg protein)

Leukocyte AGA Activity Reference Range: 24–56 nmol/hr/mg protein; AGA: α-galactosidase A

Patient 1

Patient 2

Patient 3

Patient 4

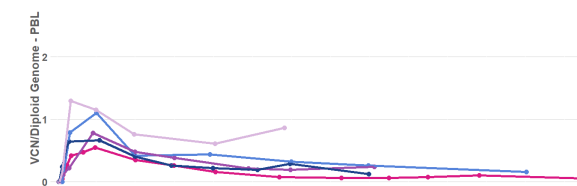
— Patient 5

Leukocyte AGA Enzyme Activity

Plasma AGA Activity Reference Range: 5.1–9.2 nmol/hr/mL; AGA: α-galactosidase A

Drug Product VCN/dg Patient 1: 0.7 Patient 2: 1.4 Patient 3: 0.8 Patient 4: 1.4

Patient 5: 1.2



AVROBIO (plate)

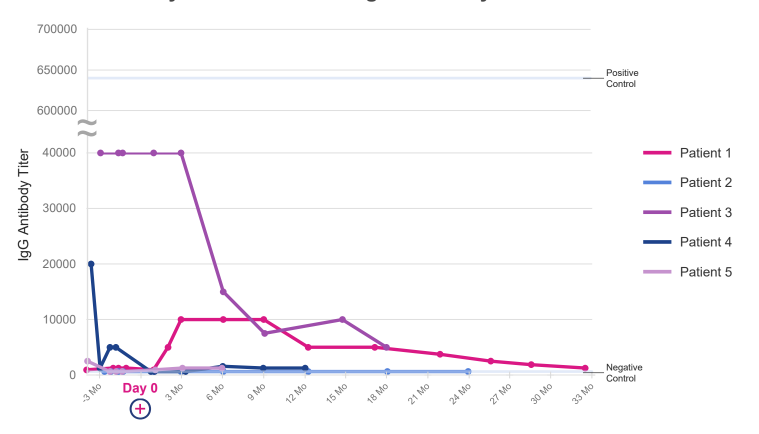
VCN: Vector Copy Number; PBL: Peripheral Blood Leukocytes; dg: Diploid Genome

Reduction of pre-existing anti-ERT drug IgG antibodies



Suggests potential as a therapeutic option independent of pre-existing antibodies

Fabry Disease Phase 1 IgG Antibody Titer



Similar results observed in other studies

San Raffaele Telethon Institute for Gene Therapy (SR-TIGET)

Change in pre-existing antibodies reported for Hurler disease (MPS-1)

- Ex vivo LV gene therapy with conditioning
- n=6
- Evaluable patients (5/6) demonstrated sustained, supraphysiologic blood IDUA activity
- 4/5 prior ERT (rhIDUA) exposure (5-28 months)
- 4/5 pre-existing ERT-induced IgG antibodies
- 6/6 anti-rhIDUA IgGs undetectable 2 months post-gene therapy

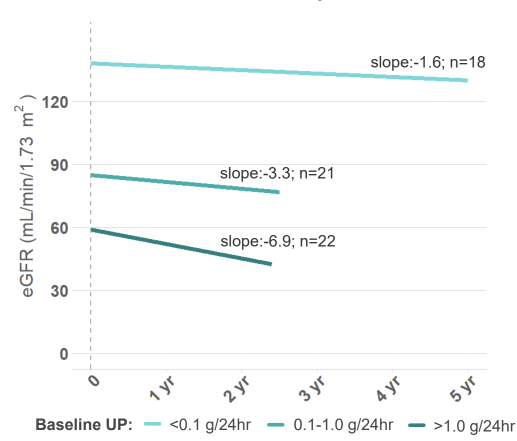


eGFR declines in natural history and on ERT

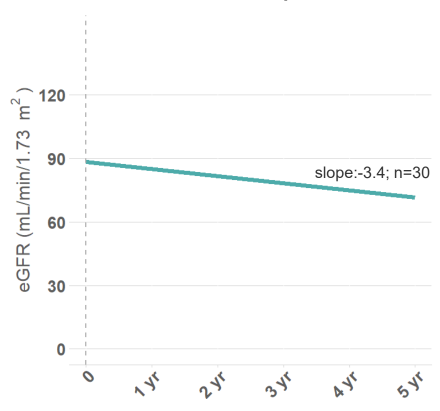


Classic Fabry male literature eGFR data

Natural history annualized eGFR slopes of treatment-naïve patients¹



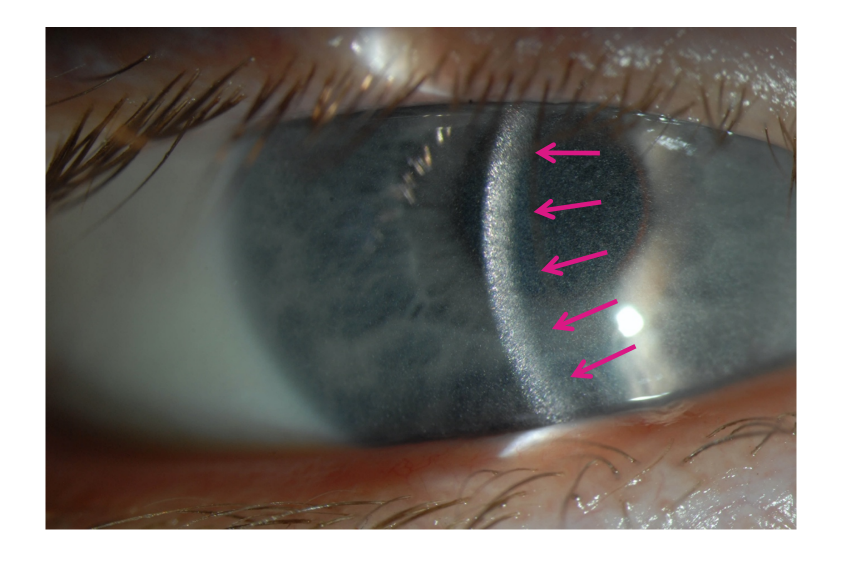
Annualized eGFR slope of ERT-treated patients²





Crystal buildup in eye clearly visible before gene therapy Patient 1 at baseline







Impact of cysteamine independence



Daily cysteamine regimen

(max per day)

Before AVR-RD-04 ON cysteamine pills

30 pills / day



ON cysteamine eye drops

Prescribed 8 drops / day



After AVR-RD-04

(16 months post-gene therapy)

OFF cysteamine pills0 pills / day

OFF cysteamine eye drops0 drops / day

