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): October 6, 2020

AVROBIO, INC.

(Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation)

001-38537 (Commission File Number) 81-0710585 (I.R.S. Employer Identification No.)

One Kendall Square Building 300, Suite 201 Cambridge, MA 02139 (Address of principal executive offices, including zip code)

(617) 914-8420 Registrant's telephone number, including area code

Not Applicable (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:

- ☐ Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- □ 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, \$0.0001 par value per share	AVRO	Nasdag 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 $\ oxtimes$

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. \Box

tem 7.01 Regulation FD Disclosure.

On October 6, 2020, AVROBIO, Inc. updated its corporate presentation for use in meetings with investors, analysts and others. A copy of the presentation is furnished as Exhibit 99.1 to this Current Report on Form 8-K.

The information in this Form 8-K shall not be deemed "filed" for purposes 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, or the Exchange Act, except as expressly set forth by specific reference in such a filing.

Item 9.01. Financial Statements and Exhibits.

(d) Exhibits

- 99.1 <u>AVROBIO, Inc. slide presentation, dated October 2020.</u>
- $104 \qquad \text{The cover page from this Current Report on Form 8-K, formatted in Inline XBR} \\$

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.

AVROBIO, INC.

Date: October 6, 2020

By: /s/ Geoff MacKay Geoff MacKay President and Chief Executive Officer



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 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 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 expected benefits and results of our implementation of the 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 runway. Any such statements in this presentation that are not statements of historical fact may be deemed to be forward-looking statements.

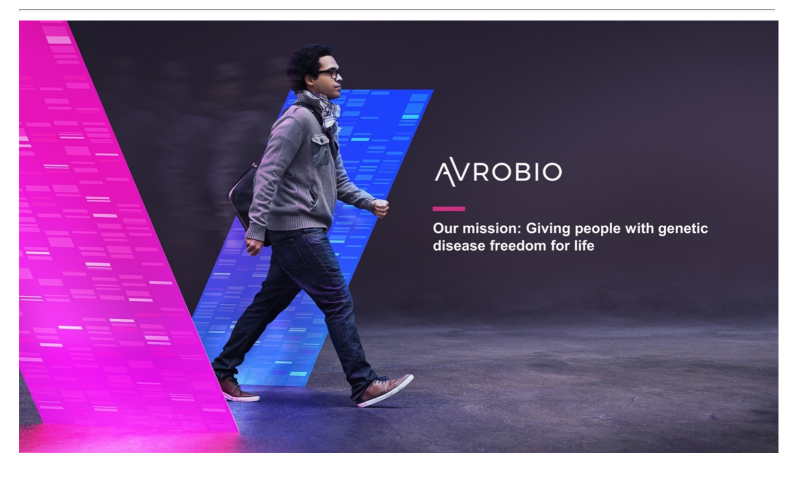
Any forward-looking statements in this presentation are based on our current expectations, estimates and projections about our industry as well as management's current beliefs and expectations of future events only as of today and are subject to a number of risks and uncertainties that could cause actual 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 of procedures in connection with the administration thereof 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 forwardlooking statements, see the section entitled "Risk Factors" in AVROBIO's most recent Quarterly Report on Form 10-Q, 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 evets, 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.





Multiple programs in the clinic 12 patients dosed to date across three indications



AVROBIO Plat

	Proof-of-Concept	IND-Enabling	Phase 1/2	Commercial Rights
Fabry AVR-RD-01	Phase 2			AVROBIO
Cystinosis AVR-RD-04	Phase 1/2			AVROBIO
Gaucher AVR-RD-02	Phase 1/2			AVROBIO
Hunter AVR-RD-05	Preclinical			AVROBIO
Pompe AVR-RD-03	Preclinical	-		AVROBIO

IND: Investigational New Drug

Addressing multi-billion dollar market opportunity



CURRENT STANDARD OF CARE COSTS

Disease	Est. Cost Per Patient Per Year	Approx. 2019 Net Sales	Selected Companies
Fabry	\$320k	\$1.4B	SANOFI GENZYME Shire
Gaucher	\$250k-400k	\$1.4B	SANOFI GENZYME Shire
Pompe	\$500k	\$1.0B	SANOFI GENZYME 🧳
Cystinosis	\$625k-700k*	\$0.2B	## HORIZON III Mylan° N RECORDATI

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

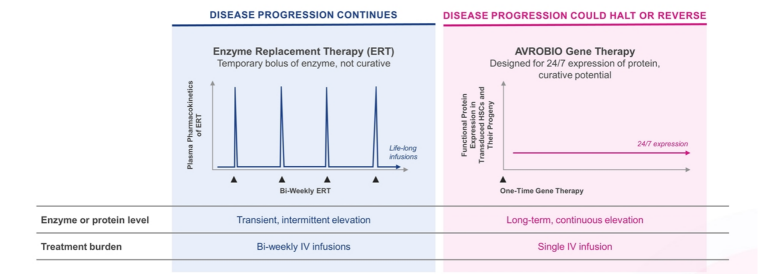
for Horizon's Procysbi oral therapy (delayed release cysteamine bitartrate,

Vote: Shire acquired by Takeda in 201



Lifelong treatments vs. potential single-dose therapy

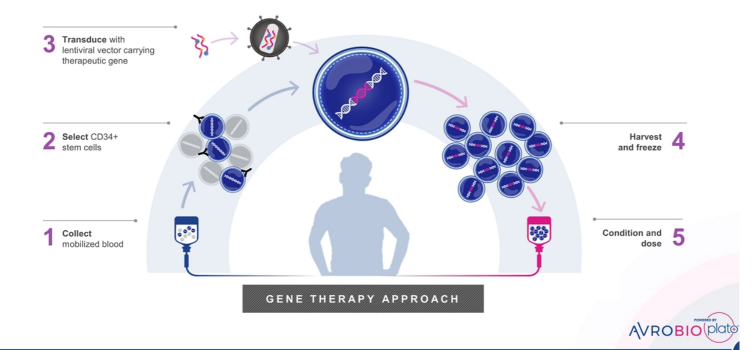






Established ex vivo lentiviral approach









Goals for gene therapy in Fabry disease

UNMET NEEDS:



Kidney function Unmet needs: proteinuria, polyuria, kidney failure



Cardiac function

Unmet needs: left ventricular hypertrophy, fibrosis, heart failure



Neuropathic pain

Unmet needs: pain and burning sensations in hands and feet, pain crises



CNS complications

Unmet needs: TIA/stroke, depression, impaired executive function, white matter



Everyday burden of illness and life expectancy Unmet needs: fatigue, inability to sweat, joint pain, abdominal pain, diarrhea, vomiting, cloudy vision, hearing loss, tinnitus, rash, angiokeratomas, biweekly infusions, shortened lifespan



Sources: Wanner C et al, Med Genetics and Metab, 2018; Burlina A, JIEMS, 2016 CNS: Central Nervous System; TIA: Transient Ischemic Attack

Two AVR-RD-01 Fabry clinical trials



9 patients dosed across Phases 1 and 2



PHASE 1

Investigator-Sponsored Trial*

PHASE 2

AVRO – FAB-201 Trial

Patients

n = 5 (fully enrolled) On ERT > 6 months prior to enrollment 18 - 50 year-old males

Patients

n = 8-12 (4 patients dosed to-date) Treatment-naive 16 - 50 year-old males

Key Objective

Safety and preliminary efficacy

Key Objectives

Safety and efficacy

FAB-201 = AVRO-RD-01-201 Study
* Sponsored by FACTs team (Fabry Disease Clinical Research and Therapeutics) in Canada ERT: Enzyme Replacement Therapy





Fabry FAB-201 • Patient Characteristics

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
Primary disease signs and symptoms	Kidney disease Chronic pain GI symptoms Decreased cold sensation	Cardiac disease Peripheral neuropathy Chronic pain Increased tiredness GI symptoms Intermittent tinnitus Mild high frequency hearing loss Raynaud's syndrome	Kidney disease GI symptoms Peripheral neuropathy Bilateral deafness Tinnitus Peripheral edema Decreased cold sensation	Chronic pain Peripheral neuropathy Neuropathic shuffling gait Lethargy Temperature intolerance Tinnitus Hearing loss GI symptoms
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***
Comment	IgA deposits in kidney biopsy	Cardiac variant, not a classic Fabry male		

^{*} Mayo Lab, ref range ≥23.1 nmol/hr/mg protein

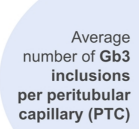


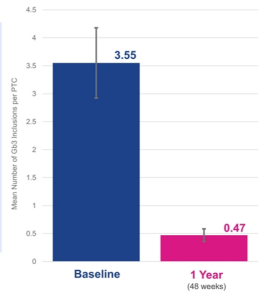
^{***} Reference value ≤ 2.4 nM

AGA: q-galactosidase A: Lvsn-Gb3: Globotriaosylsphingosine: GI: Gastrointestinal: IgA: Immunoglobulin-A



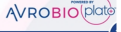
Patient 1: 87% substrate reduction in kidney biopsy at 1 year





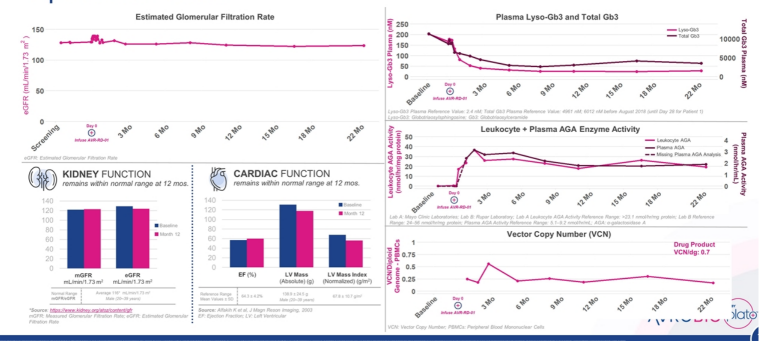
- Unpaired t-test for difference between n=55 PTCs at baseline vs. n=101 PTCs at 1 year; p < 0.0001
- Error bar represents the standard deviation

Baseline: The last available, non-missing observation prior to AVR-RD-01 infusion
Note: With respect to Fabry disease, Gb3 inclusions per PTC is interchangeable with GL-3 inclusions per KIC
FAB-201-1: First patient in FAB-201 clinical trial
PTC: Peritubular Capillary; Gb3: Globotriaosylceramide; GL-3: Globotriaosylceramide; KIC: Kidney Interstitial Capillary



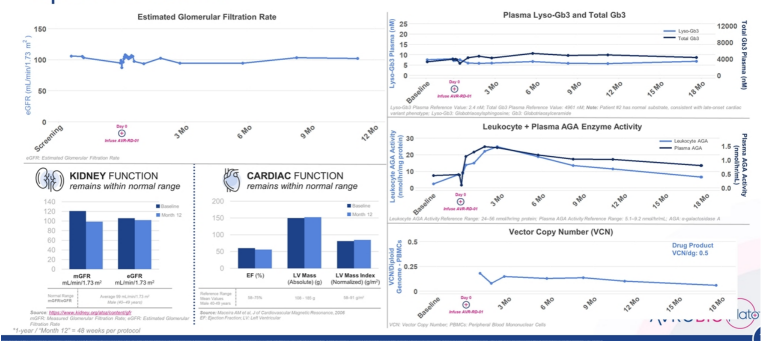
(+)

Patient 1: Sustained response across multiple measures up to 22 months



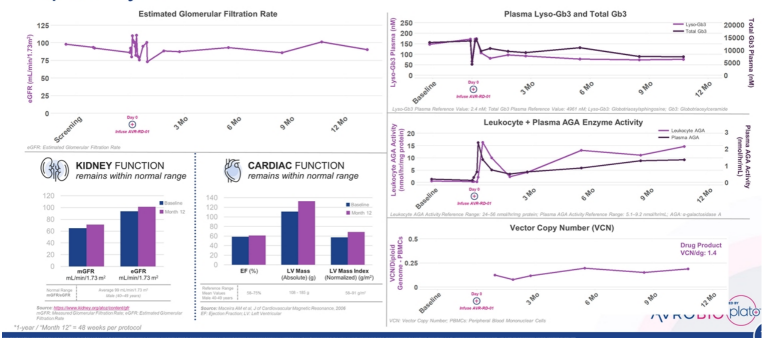
(+)

Patient 2: Sustained response across multiple measures up to 18 months



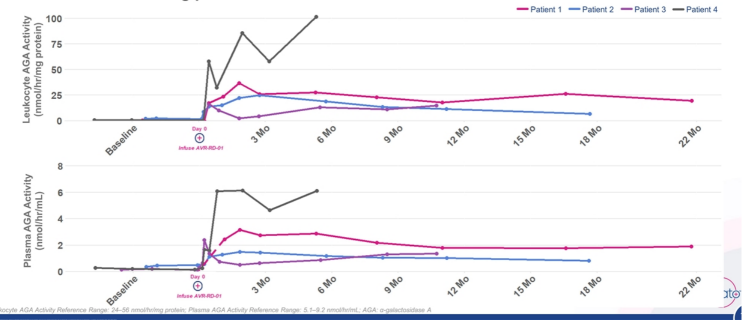
\bigoplus

Patient 3: Sustained response across multiple measures up to 1 year*





Patients 1-4: Leukocyte and plasma enzyme activity sustained up to 22 months Patient #4 dosed using plato®





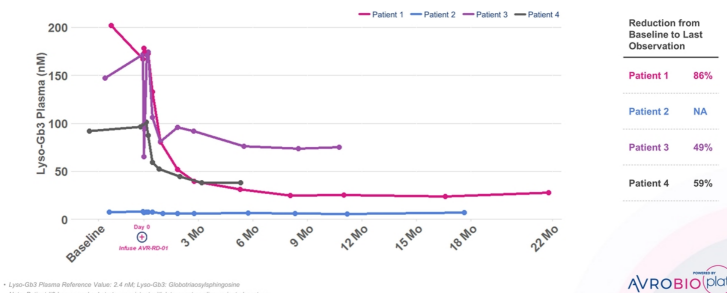
86%

NA

49%

59%

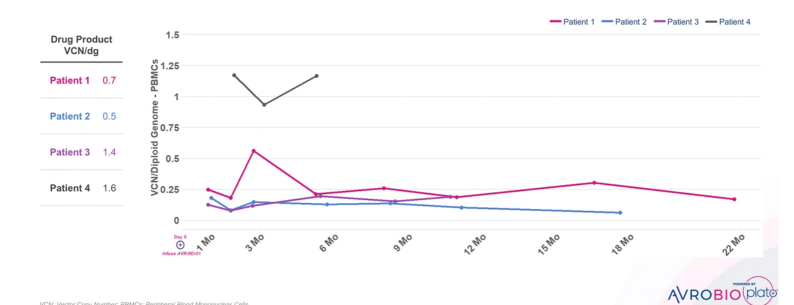
Patients 1-4: Plasma lyso-Gb3 reduction sustained up to 22 months





Patients 1-4: VCN stable up to 22 months Patient #4 dosed using plato®

VCN: Vector Copy Number; PBMCs: Peripheral Blood Mononuclear Cells



Two AVR-RD-01 Fabry clinical trials



9 patients dosed across Phases 1 and 2



PHASE 1

Investigator-Sponsored Trial*

Patients

n = 5 (fully enrolled) On ERT > 6 months prior to enrollment 18 - 50 year-old males

n = 8-12 (4 patients dosed to-date) Treatment-naive 16 - 50 year-old males







Key Objectives

Safety and preliminary efficacy

Key Objectives

AVROBIO (plat

FAB-201 = AVRO-RD-01-201 Study
* Sponsored by FACTs team (Fabry Disease Clinical Research and Therapeutics) in Canada ERT: Enzyme Replacement Therapy



Fabry Phase 1 Patient Characteristics

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)
Primary disease signs and symptoms	Kidney disease Cardiac disease GI pain GI diarrhea Angiokeratoma Insomnia	Kidney disease Cardiomyopathy Hypohidrosis Corneal verticillata Peripheral neuropathy Gl symptoms Angiokeratoma Lymphedema Acroparesthesia	Cardiac Disease Tinnitus Headaches Dizziness Acroparesthesia	Cardiac Disease Hypohidrosis Tinnitus Corneal verticillata Angiokeratoma GI symptoms	Kidney disease Hypertension Hypohidrosis Tinnitus Migraines Impaired hearing Angiokeratoma Sleep apnea Asthma Depression
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**
ERT discontinuation status	18 months after gene therapy dose		Did not resume ERT after gene therapy dose	6 months after gene therapy dose	

* Rupar Lab, ref range 24-56 nmol/hr/mg protein
** Reference value ≤ 2.4 nM protein
Note: AGA: a-galactosidase A; ERT: Enzyme Replacement Therapy; GI: Gastrointestinal; Lyso-Gb3: Globo





Patients 1-5: Plasma lyso-Gb3 reduction sustained up to 32 months

All patients who have discontinued ERT remain off ERT*



\bigoplus

Leukocyte and plasma enzyme activity sustained up to 32 months with consistent trend across all other patients

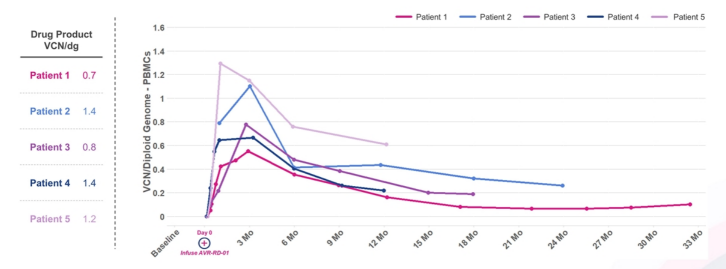
All 5 patients now out 1 year or more





Patients 1-5: VCN stable at 32 months with consistent trend across all other patients

All 5 patients now out 1 year or more



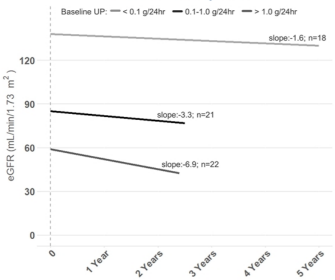
Note: 0.1 VCN is indicative of approx. 5-10% of all nucleated cells having an average of 1-2 copies of the transgene

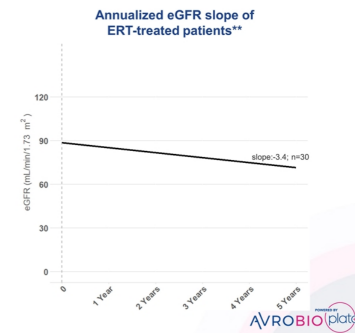




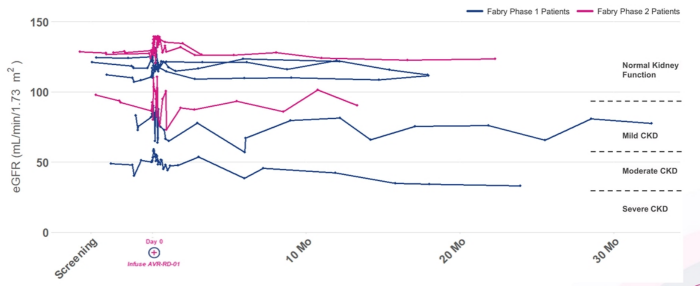
eGFR declines in natural history and on ERT Classic Fabry male literature eGFR data







Kidney function stable across Phase 1 and Phase 2 trials, up to 32 months*



* Eight of nine patients stable; other patient entered trial with more advanced kidney disease and a baseline eGFR level < 50. As expected, this patient has not stabilized, and the patient remains on Eagers: Estimated Glomerular Filtration Rate. Patient #2 from the Phase 2 trial, who is a cardiac variant and as expected has stable eGFR, has been excluded above.





Phase 1 Fabry (5 patients) and FAB-201 (4 patients)

No unexpected safety events or trends identified



No SAEs related to AVR-RD-01 drug product



AEs and SAEs reported

Phase 1 AEs (n = 100):

 Generally consistent with myeloablative conditioning, underlying disease or pre-existing conditions

FAB 201 AEs (n = 91):

- Generally consistent with myeloablative conditioning, underlying disease or pre-existing conditions
 - Grade 1 or 2 (n = 74)
 - Grade 3 or 4 (n = 17)

Phase 1 SAEs (n = 2):

- · Febrile neutropenia (grade 3)
- Thrombophlebitis (grade 2)

FAB 201 SAEs: (n = 6)

Pre-treatment and prior to conditioning

Seizure (grade 2)

Post-treatment

- Dehydration, nausea, vomiting (grade 3)
- Febrile neutropenia (2 patients, grade 3 & 4)
- Culture negative fevers (grade 2)
- Mucositis (grade 2)



Anti-AGA antibodies

Anti-AGA antibody titers observed in 4 patients in the Phase 1 trial and 2 patients in FAB-201. We believe none of these are of clinical significance.

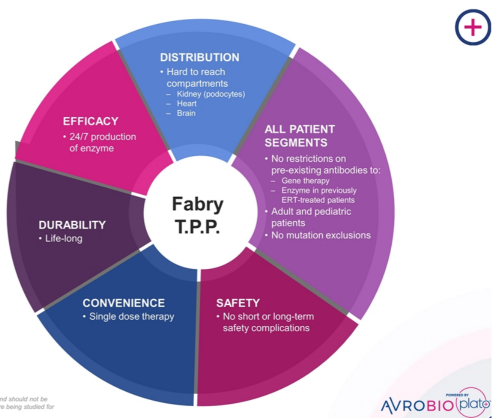


Note: Safety data cut off April 23, 2020 AE: Adverse Event; SAE: Serious Adverse Event NOTE: AVR-RD-01 is an investigational gene therapy



Fabry disease target product profile (T.P.P.)

Potential attributes intended to support first-line use



ERT: Enzyme Replacement Therapy

Note: Potential attributes represent desired target product profile, and are not intended, and should not be interpreted, to be attributes of AVROBIO's current investigational gene therapies, which are being studied for safety and efficacy and have not been approved by the FDA or any other regulatory body.

Building commercial capabilities

44+ product launches, including 1 gene therapy







- Led commercial teams for LSDs at SanofiGenzyme
- Head of Commercial at SOBI, a rare disease company







- Led global strategic marketing and global market access functions at AveXis
- Led market access, global strategic pricing and reimbursement functions for LSDs at Shire





Sean Ring VP, Head of Commercial Operations



- Extensive orphan drug commercial strategy and launch expertise
- Built out market development and commercial ops at Editas, Zafgen, Cubist, Shire, Biogen



Ramesh Arjunji VP, Global Health Economics and Outcomes Research / Value

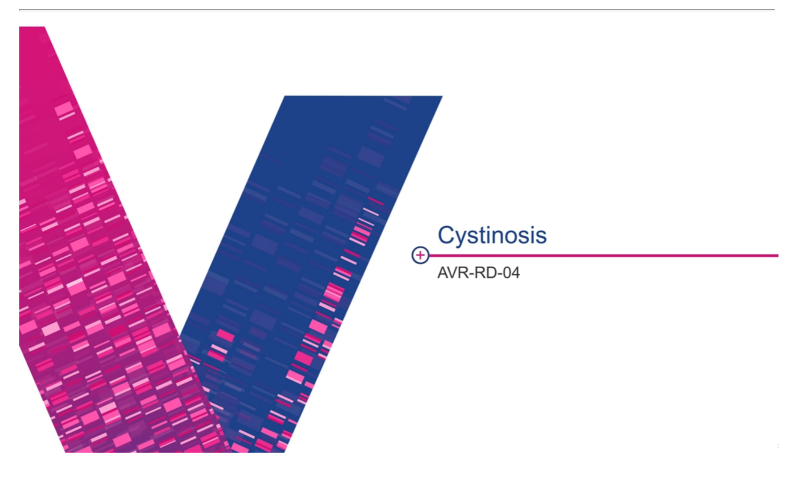


- Led value demonstration for insurers globally at AveXis
- Significant access and reimbursement responsibilities at leading global biotech companies











Goals for gene therapy in cystinosis

UNMET NEEDS:



Kidney function

Unmet needs: renal Fanconi syndrome, proteinuria, chronic kidney disease, kidney failure



Vision

Unmet needs: corneal cystine accumulation, photophobia, involuntary eyelid closure



Endocrine disorders

Unmet needs: softening/weakening of bones, bone pain, rickets, long bone deformations, hypophosphatemia, delayed growth, hypothyroidism, pancreatic insulin insufficiency, diabetes, infertility



CNS complications

Unmet needs: myopathy, hypotonia, tremors, difficulty swallowing, neurodevelopmental issues (speech and walking delay and cognitive impairment)



Everyday burden of illness and life expectancy

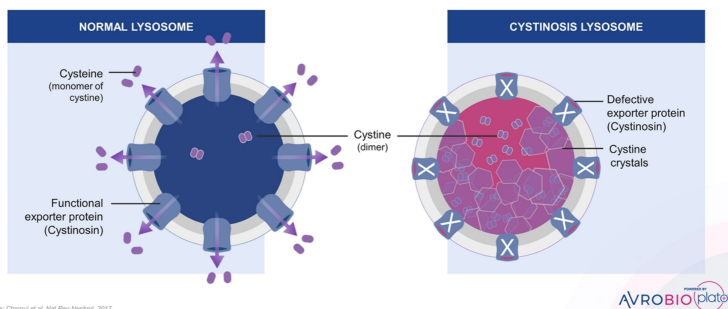
Unmet needs: medications multiple times per day that cause GI discomfort and sulfur body and breath smell, shortened lifespan





Cystinosis caused by defective gene that encodes cystinosin, an exporter protein

Cystine crystals build up in lysosomes causing tissue and organ damage



Source: Cherqui et al, Nat Rev Nephrol. 2017

Drug product-derived macrophages restore normal cystine recycling

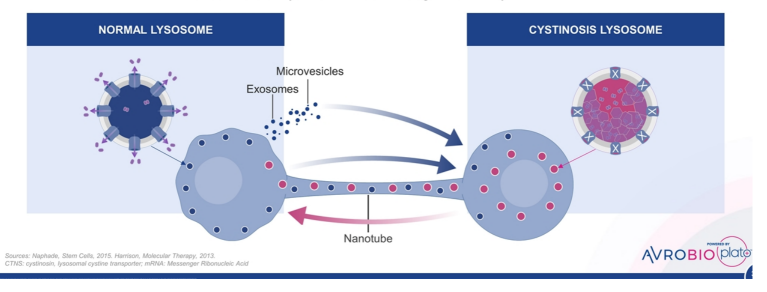


Mechanisms of action

Macrophages with CTNS transgene restore cystine recycling to CTNS-ve cells via:

- 1. Tunneling nanotubes transfer of corrected lysosomes, cystinosin, CTNS mRNA
- 2. Exosomes / Microvesicles transfer of cystinosin, CTNS mRNA

Net result: Corrected lysosomes in cells throughout the body



Allogeneic transplant demonstrated stabilized renal function, corrected polyuria and improved photophobia



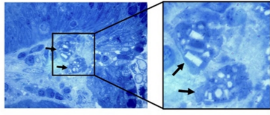
Allogenic HSC Transplant

University Hospital Leuven

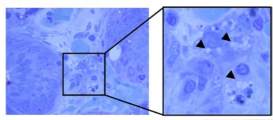
- 16 year old male
- · Diagnosed at 2.7 years old, started on cysteamine
- Age 15 years cysteamine toxicity
- Age 16 years fully matched HLA transplant
- Acute GvHD
- · First few months
- Kidney function stabilized
- Polyuria resolved
- · 6 months
 - Photophobia score reduced from 5 (unable to open eyes even inside dark room) to 0 (no photophobia)

Cystine crystal reduction (31%) in macrophages in gastric mucosa at 30 months post transplant

BEFORETRANSPLANT



30 MONTHS POST TRANSPLANT



Arrows/arrowheads point to tissue macrophages



lmonem M.A. et al. Am. J. Transplant. 2018: HSC: Hematopoietic Stem Cell: HLA: Human Leukocyte Antigen: GvHD: Graft vs Host Disease



Investigator-sponsored* study of AVR-RD-04 in cystinosis patients

Two patients dosed



Patients

Up to 6 patients
Adults and adolescents
Cohorts 1-2 ≥18 years; Cohort 3 ≥14 years
Male and Female
On oral and ophthalmic cysteamine



Key Objectives

Safety and efficacy



* Sponsored by University of California, San Diego Note: AVR-RD-04 aka CTNS-RD-04





Cystinosis
AVR-RD-04
Phase 1/2
Patient
Characteristics

	PATIENT 1	
Age of symptom onset / diagnosis	0 year / 8 months	
Age dosed with AVR-RD-04	20 years	
Gender	Male	
Mutation	Allele 1: 57-kb deletion Allele 2: c.696dupC, p.Val233Argfs*63	
Primary disease signs and SoC treatment related symptoms, including	Fanconi syndrome Polyuria Corneal abnormalities Mild photophobia Vomiting	
Granulocyte Cystine levels at baseline (nmol half cystine per mg protein)*	7.8	
Comments	NO kidney transplant; stage 3 (moderate CKD) renal failure	
	Cysteamine 1125 mg p.o. every 12 h/day since 2009; discontinued prior to AVR-RD-04 influsion Cysteamine eyedrops 4-5x/day Concomitant medications not listed	



Note: AVR-RD-01 aka CTNS-RD-04



Phase 1/2 Cystinosis

No unexpected safety events or trends identified



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



No SAEs reported



AEs reported

- Consistent with myeloablative conditioning and underlying disease
- N = 22 (moderate = 9, mild = 13)

Pre-treatment and prior to conditioning (n = 6, not all events listed)

- Diarrhea, hypokalemia, dizziness
- Dehydration, vomiting

Post-treatment (n = 16, not all events listed)

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



Note: Safety database cut as of January 27, 2020 for first patient dosed in the trial AE: Adverse Event; SAE: Serious Adverse Event



Patient 1: Initial data indicate positive trends across multiple measures



Asymptomatic Heterozygous Carrier Granulocyte Cystine Range: 0.2 – 1.9 µmol half cystine/g protein
Source: Gertsman I et al., Clinical Chemistry, 2016
VCR: Vector Copy Number, CTNS: Cystinosin, Lysosomal Cystine Transporter; mRNA: Messenger Ribonucleic Acid; eGFR: Estimated Glomerular Filtration Rate; SCr: Serum Creatinine
"Data obtained using a novel experimental methodology utilizing in vivo confocal microscopy, to image crystals in the skin behind the ear





Patient 1: Reduced treatment burden at 6 months

Number of Medications and Supplements

(max per day)

Before Gene Therapy

ON Cysteamine

52

After Gene Therapy

(at 6 months post-gene therapy)

OFF Cysteamine





NOTE: Investigational gene therapy







UNMET NEEDS:



Bone-related manifestations

Unmet needs: bone pain, avascular necrosis, bone crisis, osteoporosis, fractures, joint destruction, skeletal abnormalities



Hemoglobin levels and platelet counts

Unmet needs: anemia, thrombocytopenia, easy bruising, bleeding



Hepatosplenomegaly

Unmet needs: enlarged liver, enlarged spleen



CNS complications

Unmet needs: Increased risk of GBA-Parkinson's disease



Everyday burden of illness, and life expectancy

Unmet needs: fatigue, pain, lung disease, biweekly infusions, shortened lifespan

Sources: Grabowski G et al, Online Metabolic and Molecular Bases of Inherited Disease, 2018; Weinreb N et al, AJH, 2008; Pastores G et al, Semin Hematol, 2004 CNS: Central Nervous System: GBA; gene coding for glucocerebrosidase



Long-term follow-up study highlights significant unmet need in Gaucher Type 1



Despite standard-of-care ERT, disease progression continues and unmet need remains.

Incomplete therapeutic response is common:

- 60% of patients failed to achieve at least one of the six therapeutic goals evaluated after 4+ years of ERT¹
- A clinically significant percentage of patients continue to exhibit bone pain, organomegaly and cytopenia after 10 years of ERT²
- 25% of patients continue to suffer from physical limitations after two years of ERT, primarily due to bone disease³

Persistence after 10 years ERT †	Non-splenectomized Patients	Splenectomized Patients	
Anemia	12.4%	8.8%	
Thrombocytopenia*	22.7%	0.7%	
Splenomegaly*	38.3%	N/A	
Hepatomegaly*	14.3%	18.8%	
Bone Pain	42.9%	62.5%	
Bone Crisis	7.4%	16.7%	

^{*} Higher persistence rates observed when more severe manifestations were present at baseline

Following 10 years of treatment, \sim 26% of patients were receiving between 45-150 U/kg EOW, and 96% of these individuals were receiving doses between 45-90 U/kg EOW.

Sources: ¹Weinreb N et al. Amer J Hematol, 2008; ²Weinreb N et al. J Inherit Metab Dis, 2013; ³Giraldo P et al. Qual Life Res, 2005. GD1: Gaucher Disease Type 1; SOC: Standard of Care; ERT: Enzyme Replacement Therapy; EOW: Every Other Week



[†] Persistence refers to the presence of anemia, bone pain, bone crisis, or at least moderate thrombocytopenia, splenomegaly, or hepatomegaly, present after 10 years of ERT among those with baseline involvement of these parameters (from a registry of 757 GD1 patients; Weinreb et al., 2013).

GuardOne: Phase 1/2 study in Gaucher Type 1 patients 🖰



First patient dosed



Patients

n = 8 - 16Type 1 Gaucher Treatment naïve or on ERT 16 - 35 year-old Male and Female







Safety, Engraftment, Efficacy, ERT-independence



GAU-201: AVR-RD-02 Study; ERT: Enzyme Replacement Therapy





UNMET NEEDS:



Neurological complications

Unmet needs: cognitive deficits, seizures or behavior changes

Goals for gene therapy in Hunter syndrome



Skeletal and connective tissue issues

Unmet needs: changes in facial features, short stature, short neck, irregularly shaped and widely spaced teeth, thick skin, joint stiffness with associated restriction of movements and lump-like skin growths



Respiratory and cardiac system impacts

Unmet needs: difficulty breathing, chronic ear and sinus infections, respiratory infections and pneumonia; potential to lead to cardiac valve disease



Everyday burden of illness and life expectancy

Unmet needs: impaired vision, impaired or loss of hearing, hepatosplenomegaly, inguinal hernias, weekly infusions, significantly reduced life span

Sources: Mucopolysaccharidosis Type II - Genetics Home Reference - NIH. https://ghr.nlm.nih.gov/condition/mucopolysaccharidosis-type-liftstatistics; Scarpa M. Mucopolysaccharidosis Type II. 2007 Nov 6 [Updated 2018 Oct 4]. In: Adam MP, Ardinger HH, Pagon RA, et al., editors. GeneReviews®. https://www.ncbi.nlm.nih.gov/books/NBK1274/; Hunter syndrome. Mayo Clinic. https://www.mayoclinic.org/diseases-conditions/hunter-syndrome/symptoms-causes/syc-20350706; Mucopolysaccharidoses Fact Sheet, NINDS, NIH Publication No. 19-NS-5115. November 2019. https://www.ninds.nih.gov/Disorders/Patient-Caregiver-Education/Fact-Sheets/Mucopolysaccharidoses-Fact-Sheet; J.B. Eisengart, et al. The nature and impact of neurobehavioral symptoms in neuronopathic Hunter syndrome. Molecular Genetics and Metabolism Reports, Volume 22, 2020, 100549, ISSN 2214-4269. http://www.sciencedirect.com/science/article/bii/S2214426913301521





Planned Phase 1/2 investigator-sponsored* study in neuronopathic Hunter syndrome to evaluate safety and efficacy in CNS outcomes



PHASE 1/2 AVR-RD-05 Trial

Patients

n = 5
Early progressive form
Treatment naïve or on ERT
< 2 years old
Male



Key Objectives

Safety, Tolerability, Engraftment, Efficacy, Enzyme and Substrate biomarker response



* Sponsored by The University of Manchester, UK ERT: Enzyme Replacement Therapy





TO PREVENT OR IMPROVE:



Pulmonary function

Unmet needs: respiratory insufficiency, chronic respiratory infections, sleep apnea, artificial ventilation



Physical endurance and strength

Unmet needs: proximal myopathy, progressive muscle weakness in diaphragm, trunk and lower limbs, wheel-chair bound



CNS complications

Unmet needs: neuromuscular control, reduction in executive function, cognitive impairment



GI complications

Unmet needs: macroglossia (childhood onset), difficulty chewing and swallowing, GI symptoms, including irritable bowel-like symptoms



Everyday burden of illness, and life expectancy

Unmet needs: fatigue, hepatomegaly, independent living, biweekly infusions, shortened lifespan

AVROBIO (pla

Sources: Barba-Romero M et al, Rev Neurol, 2012; Dasouki M et al, Neurol Clin, 2014; Hagemans M et al, J Neurol, 2007; Musumeci O et al, Eur J of Neurol, 2018

Goals for gene

Pompe Disease

therapy in

Pompe lentiviral gene therapy program advancing



Integrated three-part approach

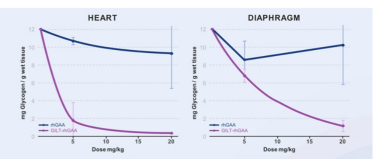
THE CHALLENGE

- Pompe requires 20x more ERT than Fabry or Gaucher
- Requires GAA activity restored to muscle and CNS

AVROBIO's APPROACH

- Potent transgene promoter
- GILT uptake tag
- Bu90-TDM for CNS impact

GILT-tagged Recombinant Human (rh)GAA impacts levels of stored glycogen compared to non GILT-tagged Recombinant Human (rh)GAA in a Pompe mouse model



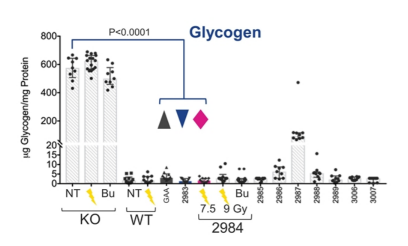


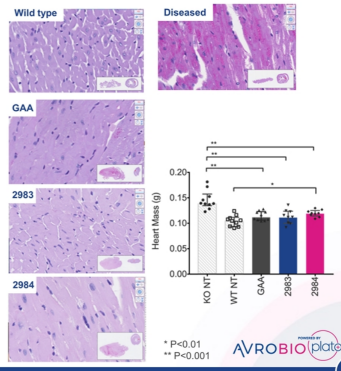
• Sources: Burton B et al, J Pediatr, 2017; Ausems M et al, Eur J Hum Genet, 1999; Gungor D et al, Orphanet J Rare Dis, 2011; Maga JA et al, J of Bio Chem, 2013; Bartelink, Lancet Haematol, 20



GILT and GILT mutant v1 reduce glycogen by >99% in heart 🕀

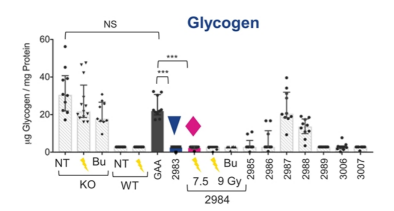


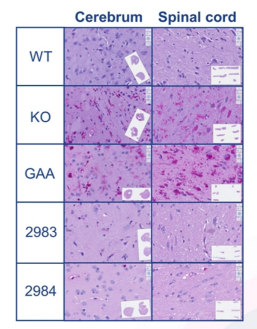




Glycogen and GILT and GILT mutant v1 similar to wildtype mice GILT tag is essential for glycogen clearance in CNS







*** P<0.001







plato®

AVROBIO's foundation designed to scale gene therapy worldwide

State-of-the-art technologies including automated manufacturing platform

Optimized for performance

+ Redefines manufacturing best practices



plato®: Three upgrades designed to optimize potency, safety and durability



ဖြင့် UPGRADES	Increase enzyme activity	Increase transduction efficiency	Increase VCN	Increase marrow space / engraftment	Increase consistency and safety
1 Vector	(+)	(+)	(+)		
2 Conditioning			(+)	(+)	(+) *
3 Automation	(+)				(+)
			to increase Vect	or Copy Number (VC	N),



TDM (therapeutic drug monitoring

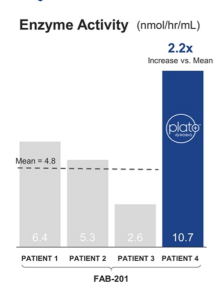


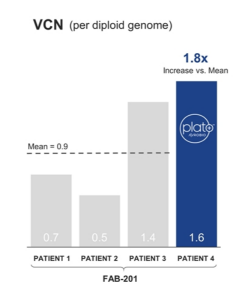
VECTOR UPGRADE:

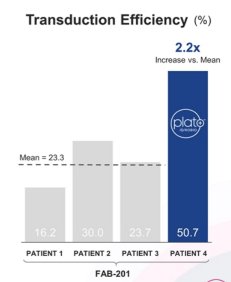
(+)

Metrics compared to academic process

FAB-201 patient #4 drug product data with plato®







AVROBIO (plate

VCN: Vector Copy Number; FAB-201: AVR-RD-01 Study NOTE: Data is from drug product

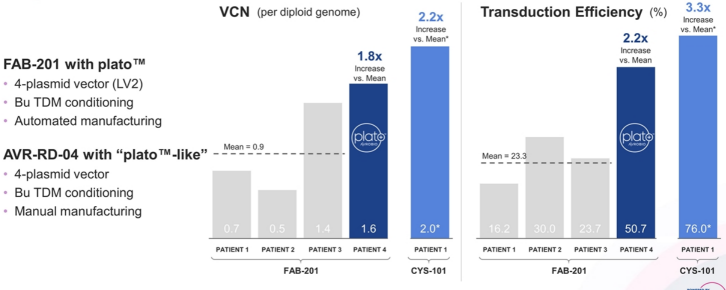


VECTOR UPGRADE:



Metrics compared to academic process

FAB-201 and AVR-RD-04 drug product data



BU TDM: Busulfan Therapeutic Drug Monitoring; VCN: Vector Copy Number; FAB-201: AVR-RD-01 Study; CYS-101: AVR-RD-04 Study; LV: Lentiviral Vector

Manufactured at UCLA using UCLA's assays and methodologies
NOTE: Data is from drug product





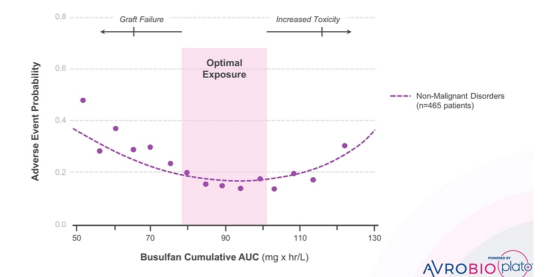


Targeted busulfan intended to balance optimal engraftment with enhanced safety

Meta-analysis of 465 patients identified optimal exposure

Optimized precision dosing designed to enhance tolerability

Lowest rate of adverse events in the Bu90 range

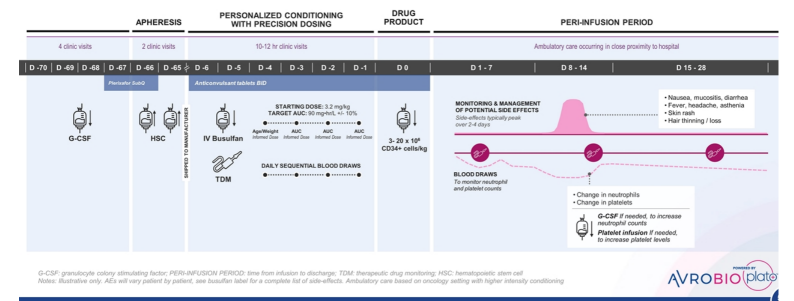


Bu: Busulfan; AUC: Area Under the Curve Sources: Bartelink IH et al, Lancet Haematol, 2016





Precision dosing via state-of-the-art patient therapeutic drug monitoring (TDM)





Busulfan used in chemotherapy has a different purpose and side-effect profile than busulfan used in cell therapy

Chemotherapy

- to eradicate cancer cells
- · Used in combinations
- Intensive high-dose chemo*
- Multiple cycles (palliative)
- · Weight-based dosing
- *Requires rescue HSC Tx

Busulfan | S the therapy

Cell Therapy

- create space in bone marrow and CNS
- · Used as a single agent
- · Less intensive
- Single cycle
- · Precision TDM dosing

Busulfan **IS NOT** the therapy





Lysosomal disorder patient characteristics are typically favorable compared to oncology patients and other gene therapy indications

Typical characteristics	Cancer patients	Other LV GT patients (eg. SCD, TDT)	AVROBIO LD patients (Fabry, Gaucher*, cystinosis, Hunter*, Pompe)
Healthy bone marrow	×	×	✓
Healthy immune systems	×	✓	✓
Healthy livers	*	ж	✓
Fewer co-morbidities	×	✓	✓
Younger	×	✓	✓

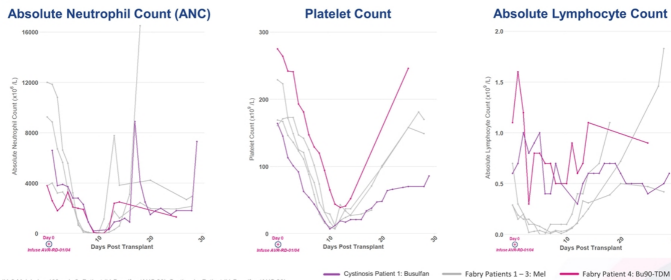
^{*} Potentially excludes treatment-naïve Gaucher disease Type 1 and treatment-naïve Hunter syndrome LV GT: Lentiviral Gene Therapy; SCD: Sickle Cell Disease; TDT: Transfusion-Dependent β-Thalassemia; LDs: Lysosomal Disorder.





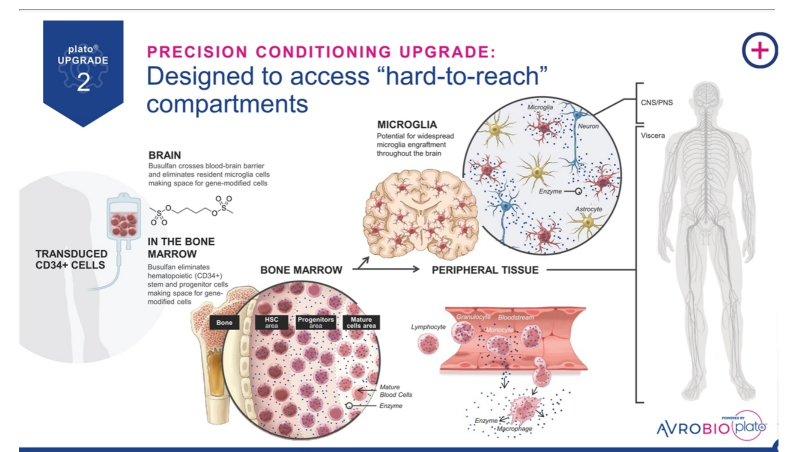


Rapid neutrophil and platelet recovery with minimal lymphocyte depletion using Busulfan TDM



bry: Patients #1-3 Melphalan 100mg/m2; Patient #4 Busulfan 'AUC 90'; Cystinosis: Patient #1 Busulfan 'AUC 90'
reshold levels for prophylactic supportive care in HSC Tx; ANC <0.5 x 10" per liter (AABB); Platelets <10 X 10" cellis/L (AABB)
DTE: Neutrophil counts - G-CSF administration post gene therapy: Pt 1: 7 Doses, Day 7 — 14, Pt 2: 11 Doses, Day 7 — 17, Pt 3: 6 Doses, Day 7 — 12, Pt 4: 5 Doses, Day 8 — 1:
DTE: Platelet counts - Platelet Transfusion: Pt 1: Day 10; Pt 2, 3: Day 11, Pt 4: no transfusion

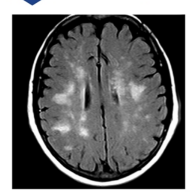




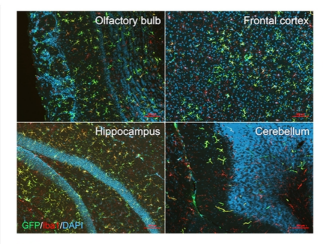




Designed to access "hard-to-reach" compartments, including the brain



MRI: 54 year old with Fabry disease demonstrating white matter lesions (WMLs)



GFP: Marker of engrafted cells lba1: Marker of microglia cells

DAPI: Nuclear stain irrespective of cell type

Global microglial coverage of mouse brain at 4 months post gene-modified HSC transplantation

- Widespread engraftment in regions critical for cognitive, motor, olfactory, and visual function
- Engrafted microglia/microglia-like cells have comparable morphology and classical lineage markers with endogenous microglia



Source: Buechner S, J. Neurol, Neurosurg, Psychiatry, 2008
MRI: Magnetic Resonance Imaging; ERT: Enzyme Replacement Therapy; WMLs: White Matter Lesions; HSC: Hematopoietic Stem Cell





Designed to elevate quality and overcome historic CMC bottlenecks





Expanded Scale

Potential to reach thousands of patients per year



Broader Reach

Portable platform designed for flexible global production using low grade clean rooms



High Quality

Automated, closed system designed to improve quality and consistency



Enhanced Convenience

Cryopreservation simplifies logistics and patient scheduling



Lower Costs

Designed to create efficiencies in vector design / scalable cell and vector production

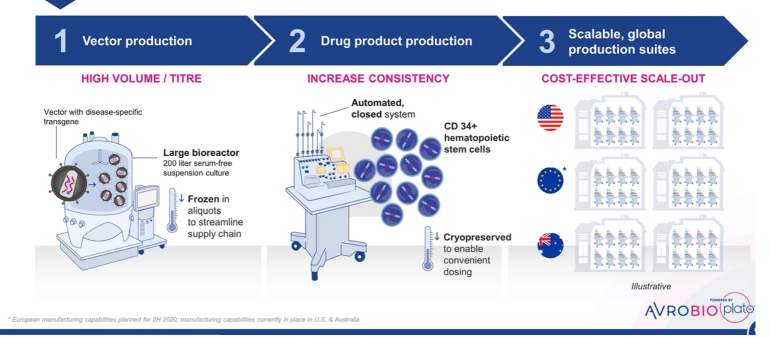






Designed to deliver large-scale manufacturing

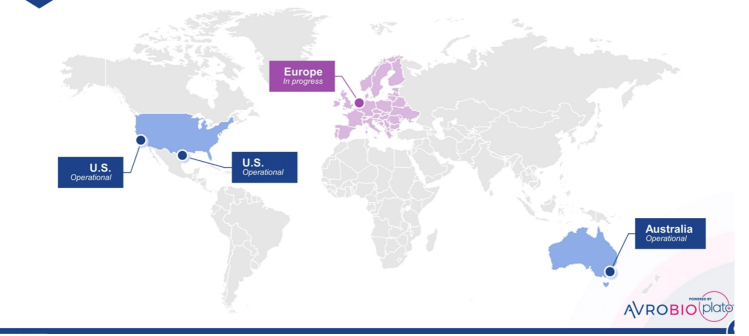
Differentiated, cost-effective approach









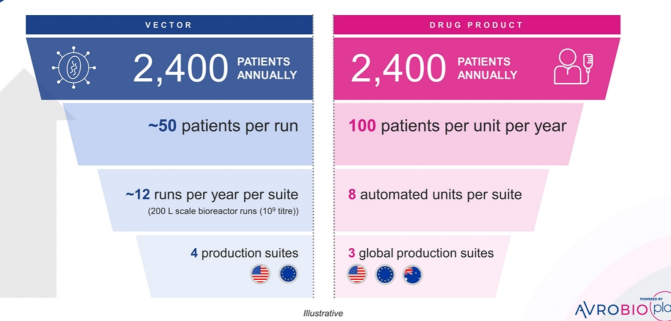






Poised to manufacture at scale

Designed to optimize potency and safety, and overcome historic CMC bottlenecks





3 UPGRADES IN PLACE:

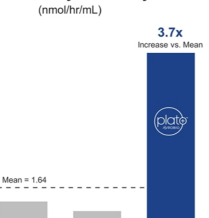


plato® metric compared to academic process

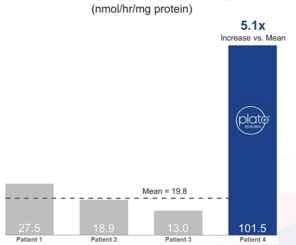
FAB-201 SIX MONTH data for patient #4 with plato® vs. patients #1-3

6.1





Leukocyte Enzyme Activity





FAB-201: AVR-RD-01 Study

Milestones anticipated across the pipeline in 2020



Clinical activities and timelines subject in all respects to ongoing and evolving COVID-19 pandemic*



FABRY

GAUCHER



CYSTINOSIS

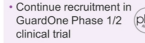


POMPE

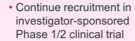
• Continue recruitment in FAB-201 Phase 2 clinical trial



 Continue to report patient data across Phase 1 and Phase 2 clinical trials



 Report initial patient data in H2 2020



 Continue to report patient data Complete preclinical IND-enabling activities

AVROBIO to hold first R&D Day on November 17, 2020

* For additional information, see the Company's Quarterly Report on Form 10-Q filed with the SEC on August 6, 2020.



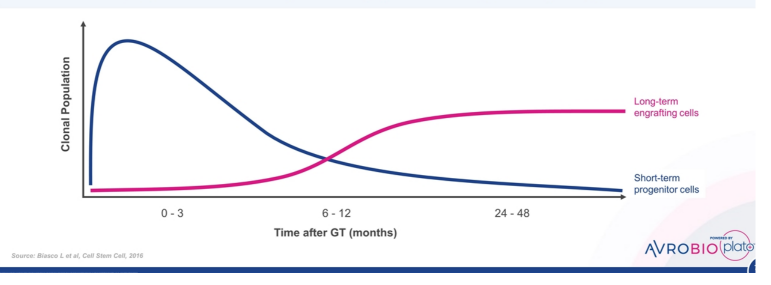




Appendix

Hematopoietic reconstitution occurs in two distinct phases A few thousand long-term engrafting cells stably sustain levels of transgene product

First wave of short-term progenitor cells start to exhaust with progressive takeover by a smaller population of long-term engrafting cells



Precedent for use of kidney biopsy data for FDA approval of drug candidate for Fabry disease



Migalastat approved on % reduction in GL-3 inclusions per KIC as compared to placebo



45 Amenable patients* (16 males / 29 females) Group | Migalastat (BL -M6) | Placebo (BL -M6) Males (N=16) | 5/7 (71%) | 4/9 (44%) | -1.10 (-1.94, -0.02) | -0.03 (-1.00, 1.69) | -0.03 (-1.00, 1.69) | -0.03 (-1.00, 1.69) | -0.01 (-1.94, 0.19) | -0.01 (-1.94, 0.19) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69) | -0.02 (-1.00, 1.69

Treatment Group	n	Baseline Median (min, max)	Month 6 Median (min, max)	Change from Baseline Median (min, max)
Average number of GL-3 inclusions per KIC (N=13)				
Galafold	7	3.6 (0.2, 6.0)	2.6 (0.1, 6.0)	-0.7 (-1.7, 1.2)
Placebo	6	1.8 (0.1, 2.8)	2.0 (0.05, 4.3)	-0.04 (-0.5, 1.5)

7/9 males ≥ 50% reduction

(at 6 months from baseline)

28% average reduction (at 6 months from baseline)



Classic Fabry patient level data 0-6 months randomized clinical trial and 6-12 months open label extension

| PTC GL-3 | Inclusions From BL/MeP to Mark | Mark

46% average reduction

(average of patients with 12 month data)

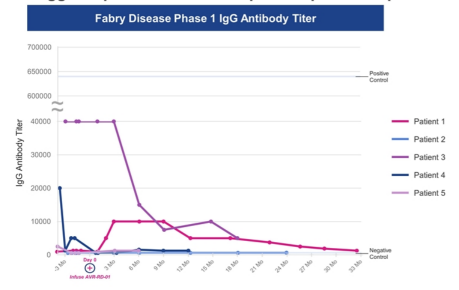


- Classic Fabry disease (AGA activity <1%)
 NOTE: For informational purposes; differen
- NOTE: For informational purposes; differences exist between trial designs and subject populations; AVROBIO has not conducted any head-to-head trials comparing migalastat to AVR-RD-to-head trials comparing m



Reduction of pre-existing anti-ERT drug IgG antibodies following AVR-RD-01

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



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-CD34+ 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

Source: Gentner B et al., Blood, 2019



ERT: Enzyme Replacement Therapy; IgG: Immunoglobulin G; MPS-1: Mucopolysaccharidosis Type 1; IDUA: Iduronidase: SR-TIGET: San Raffaele Telethon Institute for Gene Therapy; LV: Lentiviral; rhIDUA: Recombinant Human alpha-L-Iduronidase

New collaborations advancing leadership in lentiviral gene therapy













Fully Automated Bu-TDM Immunoassay

- AVROBIO and Saladax announced agreement to develop and validate Saladax's fully automated nanoparticle immunoassay kit
- Designed to work on most automated hospital analyzers
- Regular technician, 24/7 analysis possible with results anticipated in minutes using only microLs of blood
- Designed to analyze 10-100s samples per machine/hour
- Expected to eliminate Bu degradation errors as assay conducted in real-time at the point of care
- Scalable

Antibody-Drug Conjugate

- AVROBIO and Magenta announced research & clinical collaboration agreement to evaluate Magenta's preclinical CD117-targeted antibody conjugate to amanitin (MGTA-117) in conjunction with AVROBIO investigational gene therapies
- Designed to deplete only hematopoietic stem and progenitor cells
- Has shown promising data in non-human primates
- MGTA-117 currently in IND-enabling studies
- P Each party retains commercial rights to its own programs

 AVROBIO