FibroGen, Inc. Corporate Presentation

June 2022



Forward-Looking Statements

This presentation contains "forward-looking" statements that involve substantial risks and uncertainties. All statements other than statements of historical facts contained in this presentation, including statements regarding our future financial condition, business strategy, and plans, and objectives of management for future operations, are forward looking statements. These forward-looking statements can generally be identified by terminology such as "believe," "will," "may," "estimate," "continue," "anticipate," "contemplate," "intend," "target," "project," "should," "plan," "expect," "predict," "could," "or potentially," or by the negative of these terms or other similar expressions. Forward-looking statements appear in a number of places throughout this presentation and include statements regarding our intentions, beliefs, projections, outlook, analyses, or current expectations concerning, among other things, our ongoing and planned development and clinical trials, the timing of and our ability to make regulatory filings and obtain and maintain regulatory approvals for roxadustat, pamrevlumab, and our other product candidates, the potential safety, efficacy, reimbursement, convenience, or clinical and pharmaco-economic benefits of our product candidates, including in China, the potential markets for any of our product candidates, our ability to develop commercial functions, results of commercial operations, or our ability to operate in China, expectations regarding clinical trial data, our results of operations, cash needs, spending of proceeds from our public offerings, financial condition, liquidity, prospects, growth, and strategies, the industry in which we operate, and the trends that may affect the industry or us.

Forward-looking statements involve known and unknown risks, uncertainties, assumptions, and other factors that may cause our actual results, performance, or achievements to be materially different from any future results, performance, or achievements expressed or implied by the forward-looking statements, including the other risks and uncertainties that are described in the Risk Factors section of our most recent annual report on Form 10-K or quarterly report on Form 10-Q filed with the Securities and Exchange Commission. Forward-looking statements represent our management's beliefs and assumptions only as of the date of this presentation. Except as required by law, we assume no obligation to update these forward-looking statements publicly, or to update the reasons why actual results could differ materially from those anticipated in the forward-looking statements, even if new information becomes available in the future.



Company Overview

Mission

Developing innovative, first-in-class medicines for the treatment of chronic and life-threatening or debilitating conditions

Employees

~575 · ~300 US · ~275 ex-US worldwide

Cash as of March 31, 2022

\$ 565.4 million

Estimated 2022 ending cash to be in the range of \$310-\$340 million



Strategic Objectives: Three Areas of Focus

01

Developing pamrevlumab in three high value indications: Idiopathic pulmonary fibrosis (IPF), locally advanced pancreatic cancer (LAPC), and Duchenne muscular dystrophy (DMD)

02

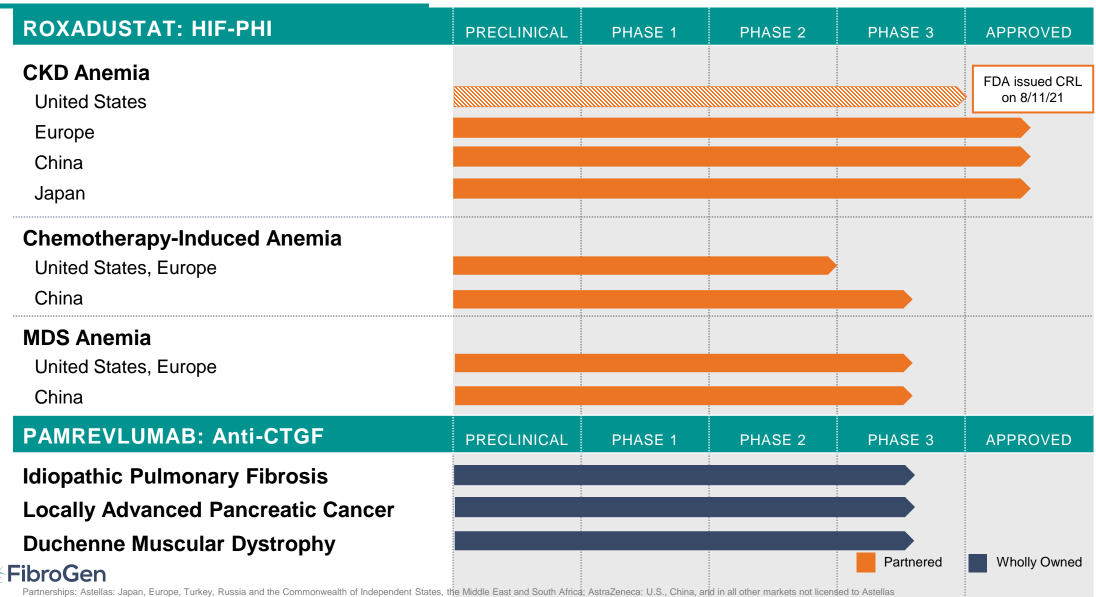
Ensuring regulatory and commercial success of roxadustat in chronic kidney disease (CKD) and other indications

03

Increasing research productivity to advance novel programs that leverage internal expertise and access external innovation

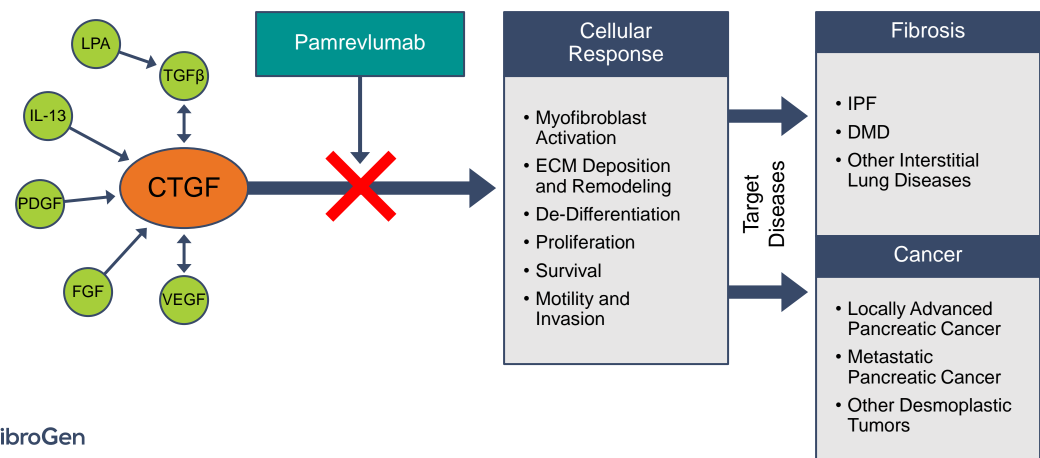


FibroGen Marketed and Late-Stage Portfolio



Pamrevlumab: Innovative Treatment for Fibrosis and Fibroproliferative Disease

 PAMREVLUMAB – Fully human monoclonal antibody targeting activity of connective tissue growth factor, a central factor in fibrosis





Current Status of Pamrevlumab Development

PAMREVLUMAB

Idiopathic Pulmonary Fibrosis

ZEPHYRUS-1 Phase 3 Study

ZEPHYRUS-2 Phase 3 Study

Enrollment Complete







 LAPIS Phase 3 Study Enrollment Complete

Pancreatic Cancer

Metastatic Pancreatic Cancer

Precision PromiseSM Platform Phase
 2/3 Sponsored by Pancreatic Cancer
 Network - Enrolling



Duchenne Muscular Dystrophy

- LELANTOS-1 (non-ambulatory)
 Phase 3 Study
 Enrollment Complete
- LELANTOS-2 (ambulatory)
 Phase 3 Study Enrollment
 Complete

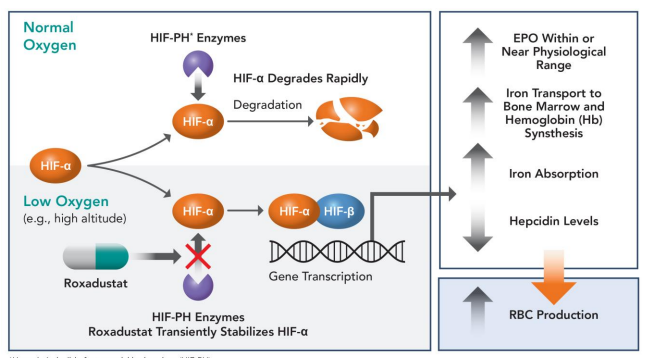


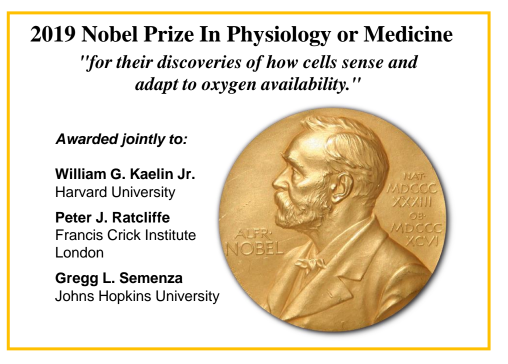
Enrolling

Roxadustat: Novel, First-in-Class Treatment for CKD Anemia

ROXADUSTAT – Oral hypoxia-inducible factor prolyl hydroxylase (HIF-PH) inhibitor based on 2019 Nobel Prize-winning science

- Increases hemoglobin (Hb) by mimicking the body's natural response to low oxygen
- Studied for treatment of anemia in Stage 3-5 CKD patients, both on and not on dialysis





^{*}Hypoxia-inducible factor prolyl hydroxylase (HIF-PH)



Roxadustat Update

Anemia of Chronic Kidney Disease (CKD)







Received European Commission
Approval for First-in-Class
EVRENZO™ (roxadustat) for the
treatment of symptomatic anemia of
chronic kidney disease (CKD) in
adult patients.

 Astellas has launched EVRENZO™ in Germany, the United Kingdom, the Netherlands, Austria, and the Nordic countries.



Continued strong roxadustat performance in China.

Roxadustat is the number one brand based on value share in the anemia of CKD market in China.



Pre-Clinical Pipeline:

Licensed programs in transformative partnership with HiFiBiO Therapeutics

- <u>Galectin-9</u>: antibody designed to inhibit target driven cancer stem cell self-renewal in acute myeloid leukemia (AML) and immune resistance in many solid tumors.
- <u>CCR8</u>: antibody designed to deplete suppressive T regulatory cells in the tumor microenvironment with broad potential activate immune responses in solid tumors.

FibroGen fully owned proprietary assets

 Undisclosed: additional drug development programs leveraging FibroGen expertise in HIF and CTGF biology



Pamrevlumab

Fibrosis

Pamrevlumab: A Unique Phase 3 Investigational Drug



Novel, differentiated antifibrotic MOA

- First-in-class CTGF-targeting mAb
- In vivo efficacy in multiple preclinical models
- Potential disease-modifying mechanism across fibrotic diseases



Phase 2 outcomes target serious unmet needs

- IPF: Observed benefit in lung function preservation with a differentiated safety profile
- PDAC: Safe and well tolerated with trend for improved resection rate and increased completion of chemotherapy cycles
- DMD: Safe and well tolerated with favorable outcomes vs historical data
 - Pamrevlumab can potentially be used in a broad range of DMD patients regardless of specific mutations



Significant commercial potential

- Phase 3 programs in areas of high unmet medical need with limited late-stage competitive intensity
- Highly differentiated target indications with fibrotic components but independent risk profiles
- Esbriet and Ofev combined 2021 sales ~\$4.0B



Pamrevlumab: Potential First in Class Antibody Targeting High Need Medical Indications

Idiopathic Pulmonary Fibrosis (IPF)

- FDA Fast Track and Orphan Drug Designation
- ZEPHYRUS-1 Phase 3 Study Enrollment Complete
- ZEPHYRUS-2 Phase 3 Study Enrolling

Pancreatic Cancer (LAPC)

- FDA Fast Track and Orphan Drug Designation
- LAPIS Phase 3 Study Enrollment Complete

Metastatic Pancreatic Cancer

Precision PromiseSM Platform Phase 2/3
 Sponsored by Pancreatic Cancer Network
 Enrolling

Duchenne Muscular Dystrophy (DMD)

- FDA Orphan Drug, Fast Track and Rare Pediatric Disease Designation
- EMA Orphan Medicinal Product Designation
- LELANTOS-1 Phase 3 Enrollment Complete
- LELANTOS-2 Phase 3 Study Enrollment
 Complete



Pamrevlumab Commercial Opportunity

Idiopathic Pulmonary Fibrosis

Diagnosed Prevalence (US, EU, CN, JP)	~330k
2021 Branded Category Revenue	~\$4.0B; +11% YoY
Current Standard of Care	Ofev (nintedanib – BI); Esbriet (pirfenidone – Genentech / Roche)
SoC Limitations	Disease progression; poor tolerability / adherence
Late-Stage Competitive Intensity	PRM-151 (Roche), BI-1015550 (BI)



IPF Patients Need New Therapeutic Options



- One in 200 over the age of 70 are living with IPF¹
- U.S. prevalence of 200,000+1
- U.S. incidence of 50,000¹ cases per year



- Leads to irreversible loss of lung function with high morbidity and mortality rates
- Median survival of 3-5 years following diagnosis



- Slow pulmonary function loss
- Modest effect on slowing disease progression
- Require side effect management
- Esbriet and Ofev combined 2021 sales ~\$4.0B

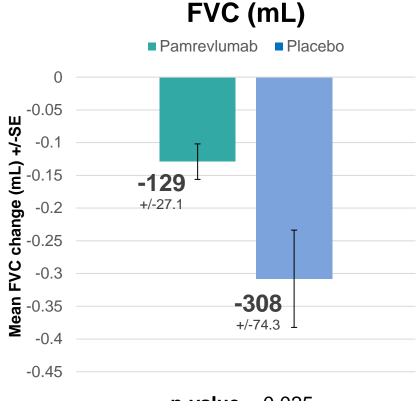


PRAISE Phase 2: Efficacy Analysis, Primary Endpoint; Mean Change from BL at Week 48 in FVC

FVC%-Predicted ■ Pamrevlumab ■ Placebo 0 Mean FVC% change +/-SE -2.85 -5 +/-0.79 -7.17 +/-1.86 -10

p-value = 0.033

FVC%-Predicted Difference: 4.33% Relative Difference: 60%

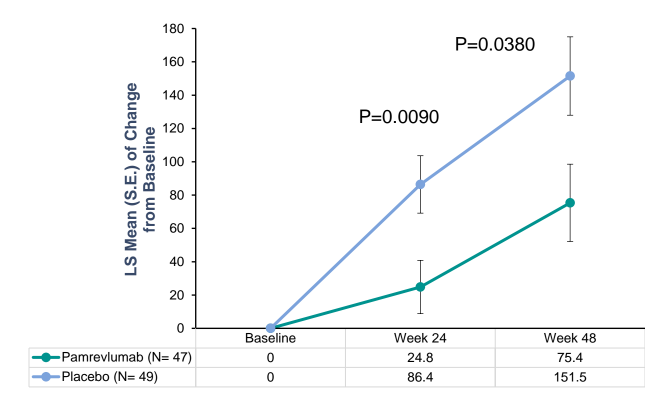


p-value = 0.025

Absolute FVC Difference: 178mL Relative Difference: 58%



PRAISE Phase 2: Pamrevlumab Attenuated Progression of Fibrosis



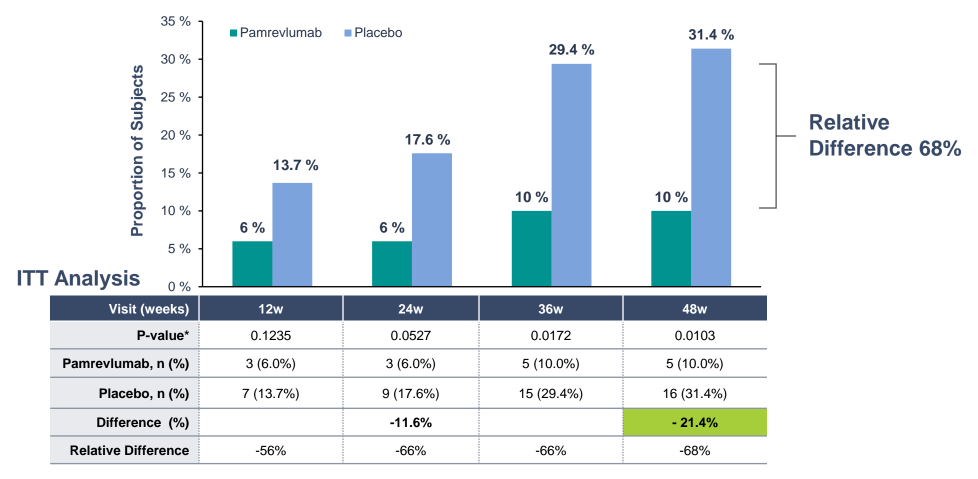
- Absolute differences in Quantitative Lung Fibrosis (QLF) changes from baseline to Weeks 24-48 between pamrevlumab arm and placebo arm were 61.6 ml and 76.2 ml, respectively
- First reported statistically significant results for attenuation of fibrosis by qHRCT
- Change in fibrosis (lung structure)
 correlates with change in FVC% predicted
 (lung function), primary endpoint of study
 (Spearman's correlation coefficient of
 -0.64, p=0.0001)

Richeldi, et al. Lancet Respir Med 2020 Jan;8(1):25-33.



PRAISE Phase 2: Attenuation of IPF Disease Progression

IPF Disease Progression (FVC% Predicted Decline ≥10% or Death) vs. Placebo





Pamrevlumab IPF Phase 3 Program: ZEPHYRUS-1 and ZEPHYRUS-2

Patient Population

- IPF diagnosis according to ATS/ERS/JRS/ALAT guidelines*
 - ZEPHYRUS-1
 - IPF patients who have declined approved therapies
 - IPF patients previously but not now currently being treated with approved therapies
 - ZEPHYRUS-2
 - IPF patients previously but not now currently being treated with approved therapies

Study Design

- Placebo-controlled, double-blind
- Enroll ~340 patients in each study
- Randomization 1:1 pamrevlumab or placebo

Primary Endpoint

- Change in forced vital capacity (FVC) from baseline for U.S
- Disease progression defined as FVC % predicted decline of 10% or death for EU/UK

Secondary Endpoints

- Quantitative changes in lung fibrosis volume from baseline
- Patient-reported outcomes



Study Fully Enrolled



NCT04419558

Study Enrolling



Pamrevlumab Commercial Opportunity

Locally Advanced Pancreatic Cancer

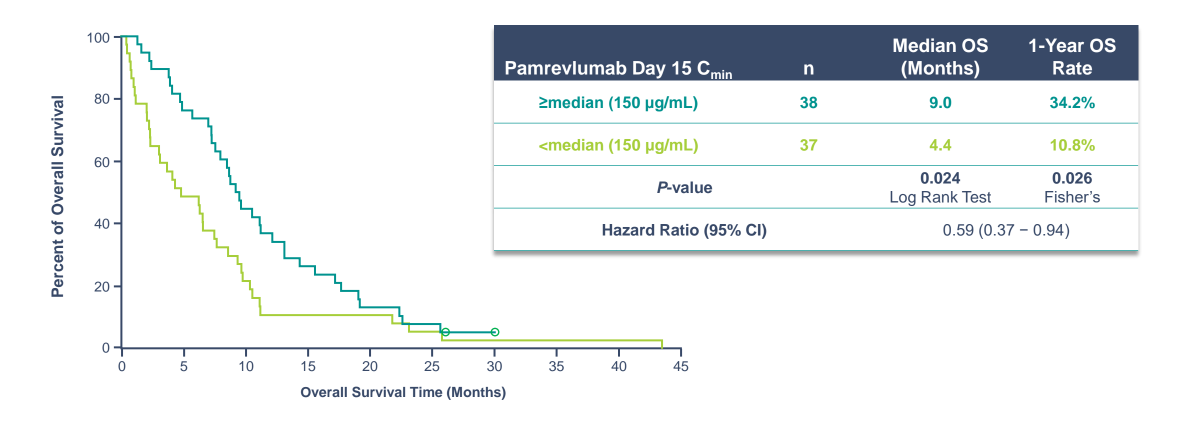
Diagnosed Prevalence (US, EU, CN, JP)	~140k (non-metastatic)
Branded Category Revenue	N/A
Current Standard of Care	gemcitabine + nab-paclitaxel; gemcitabine + folfirinox
SoC Limitations	5-year Disease-Free Survival ~10%¹; No major therapeutic advances in decades,² with major therapy classes like IOs failing to offer survival benefits
Late-Stage Competitive Intensity	Limited in non-metastatic disease

^{1.} Sources: GLOBOCAN (<u>link</u>) 2. Sources: SEER; Cancer.Net (for <u>NSCLC</u> and <u>H&N</u>); Dela Cruz, Charles S et al. "Lung cancer: epidemiology, etiology, and prevention." *Clinics in chest medicine* vol. 32,4 (2011): 605-44. doi:10.1016/j.ccm.2011.09.001 (for <u>SCLC</u>)



Phase I in Locally Advanced or Metastatic Pancreatic Ductal Adenocarcinoma Cancer Patients

Improved OS with Higher Pamrevlumab Exposure





LAPIS Pamrevlumab LAPC Phase 3 Study

Patient Population

- Locally advanced, unresectable pancreatic cancer
- ECOG 0-1 (health status of patient)
- No prior therapy

Primary Endpoint

Overall Survival (OS)

Secondary Endpoints

- Event-free survival
- Patient-reported outcomes

Study Design

- Double-blind, placebo-controlled
- Enrolled 284 subjects at sites globally
- Randomization 1:1 to pamrevlumab + gemcitabine/nab-paclitaxel/ FOLFIRINOX or to placebo + gemcitabine/nab-paclitaxel/ FOLFIRINOX
- Six cycles of neoadjuvant treatment followed by evaluation for surgical eligibility and possible resection
- Interim assessment of Event Free Survival
- Long-term overall survival follow-up for all subjects



NCT03941093

Study Fully Enrolled



Pamrevlumab Commercial Opportunity

Duchenne Muscular Dystrophy

Diagnosed Prevalence (US, EU, CN, JP)	~70k
2021 Branded Category Revenue	~\$0.75B
Current Standard of Care	corticosteroids; anti-sense oligonucleotides / exon-skipping
SoC Limitations	Continued disease progression; tolerability; need for confirmed gene mutation; biomarker benefit only (exon-skipping therapies)
Late-Stage Competitive Intensity	Dystrophin-targeted therapies (gene-based and exon-skipping); Limited anti-fibrotic therapies



Duchenne Muscular Dystrophy (DMD) Background

- Affects ~1 in every 5,000 newborn boys
- About 20,000 children are diagnosed with DMD globally each year
- A fatal disease caused by genetic mutations leading to the absence or defect of dystrophin, a protein necessary for normal muscle function:
 - The absence of dystrophin results in muscle weakness, muscle loss, fibrosis, and inflammation
- Patients with DMD are often non-ambulatory before the age of 12, and their progressive muscle weakness can lead to serious medical problems relating to diminished function of respiratory and cardiac muscles



Phase 2 MISSION Study in DMD

First Time an Antifibrotic Has Shown Potential to Slow Disease Progression in Non-ambulatory DMD

- Mechanism of action may mitigate fibrosis in non-ambulatory DMD patients, irrespective of the causative genetic mutation
- Phase 2 Study 079 (MISSION) performed in non-ambulatory DMD subjects, showed pamrevlumab may slow DMD disease progression
- Promising safety profile, with no major SAEs leading to discontinuations
- Pamrevlumab can potentially be used in a broad range of DMD patients regardless of specific mutations
- Limitations: This study was limited by lack of an internal control group



LELANTOS-1 Pamrevlumab DMD Phase 3 Study

Patient Population

 Males 12 years and older with nonambulatory DMD

Primary Endpoint

 Change in the total score of performance of upper limb (PUL) assessment, from baseline to Week 52

Secondary Endpoints

- Pulmonary function tests
- Cardiac function tests

Study Design

- Double-blind, placebo-controlled
- Enrolled 99 subjects at sites globally
- Randomized 1:1 to receive pamrevlumab plus systemic corticosteroids, or placebo plus systemic corticosteroids, for up to 52 weeks



NCT04371666

Study Fully Enrolled



LELANTOS-2 Pamrevlumab DMD Phase 3 Study

Patient Population

 Males 6-12 years old with ambulatory DMD

Primary Endpoint

 Ambulatory function assessment, measured by the change in North Star Ambulatory Assessment (NSAA) from baseline to Week 52

Secondary Endpoints

 Additional functional secondary endpoints will be assessed in the study

Study Design

- Double-blind, placebo-controlled
- Enroll ~70 subjects at 60-80 sites globally
- Randomized 1:1 to receive pamrevlumab plus systemic corticosteroids, or placebo plus systemic corticosteroids, for up to 52 weeks
- Subjects who complete the 52week study will be eligible for rollover into an open-label extension study



NCT04632940

Study Fully Enrolled



Roxadustat

Anemia



Roxadustat: An Innovative Approach to Addressing Anemia

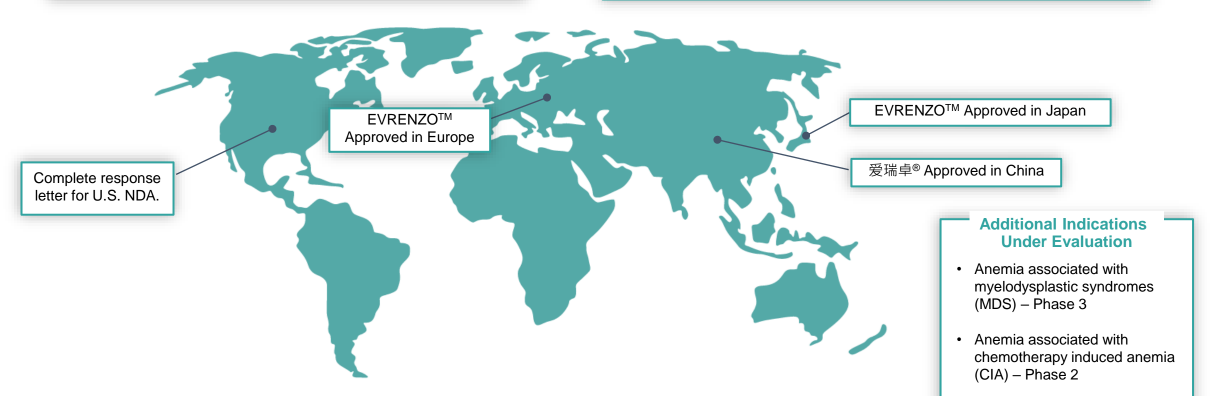
Transformational Oral Anemia Therapy Leveraging Body's Natural Response to Hypoxia

An oral hypoxia-inducible factor prolyl hydroxylase inhibitor (HIF-PHI) based on Nobel Prize winning science that stimulates a coordinated erythropoiesis response.

Strategic Partnership with Astellas and AstraZeneca

Astellas: Japan, Europe, Turkey, Russia and the Commonwealth of Independent States, the Middle East and South Africa

AstraZeneca: U.S., China, and all other markets not licensed to Astellas



Roxadustat Collaboration Economics

Royalty/Transfer Price in low 20% range in U.S./EU and ROW

China 50/50 profit split

All development costs and commercialization costs paid by partners, ex-China

Regulatory and/or commercial milestones available under our collaboration agreements



MATTERHORN Roxadustat Anemia Associated with Myelodysplastic Syndromes (MDS) Phase 3 Study

Patient Population

 Anemic Patients with Lower Risk Myelodysplastic Syndrome (MDS)

Primary Endpoint

- Transfusion independence ≥ 56 consecutive days
- Hemoglobin correction/maintenance and reduction of RBC transfusion

Secondary Endpoints

- Safety
- Quality of life parameters
- Transfusion independence

Study Design

- Randomized Double-Blind Placebo-Controlled Study
- Enroll ~160 subjects at ~72 sites globally

Topline data expected 2H 2022 / 1H 2023



NCT03263091



Roxadustat

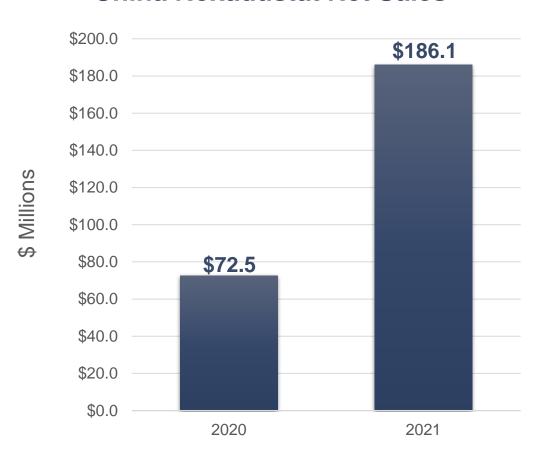
China

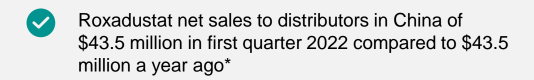




CHINA: Significant Year Over Year Roxadustat Net Sales Growth

China Roxadustat Net Sales





FibroGen net product revenue under U.S. GAAP of \$18.9 million in first quarter 2022

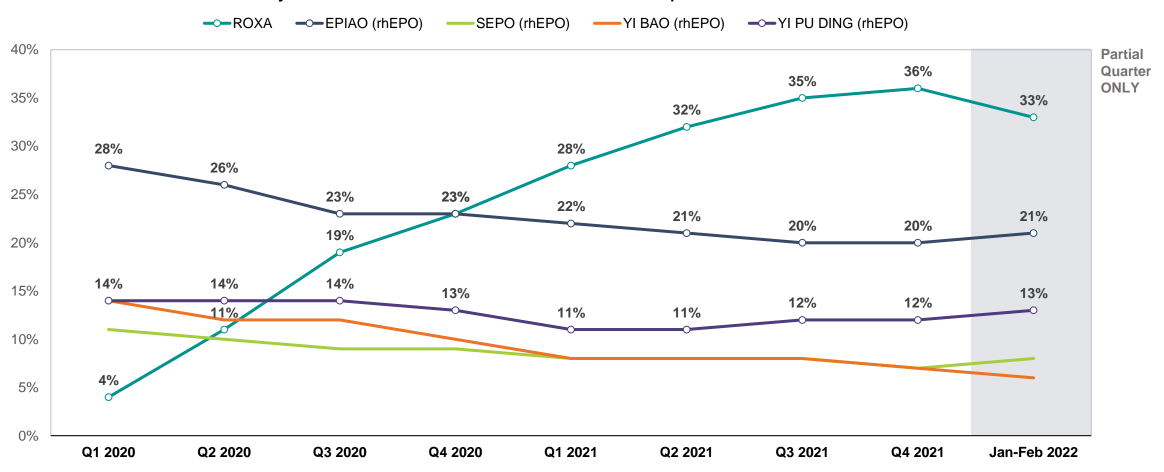
^{*}Total roxadustat net sales in China includes sales made by the distribution entity as well as FibroGen China's direct sales, each to its own distributors. The distribution entity jointly owned by AstraZeneca and FibroGen is not consolidated into FibroGen's financial statements.





CHINA: Roxadustat Maintains ESA + HIF Category Leadership Based On \$ Sales

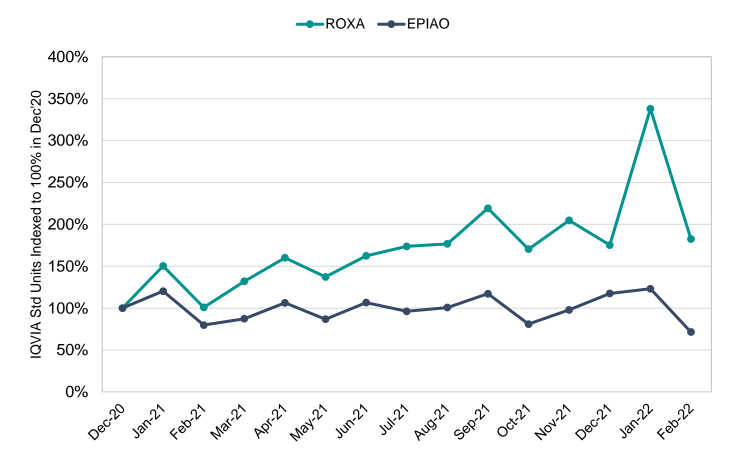
Quarterly Brand Share based on \$ Sales - Top 5 of ESA+HIF Market





CHINA: Roxadustat Unit Volume in last 3 months doubled versus prior year; EPIAO volume flat

IQVIA MIDAS Standard Units indexed to 100% in Dec '20



IQVIA Standard Units – Roxadustat vs EPIAO

	Quarter Ending Feb '21	Quarter Ending Feb '22	% Growth
Roxadustat	2,872,530	5,690,745	98%
EPIAO	3,418,561	3,556,255	4%

Clinical Trial Timelines

Study Phase	Indication	Study Name	Enrollment Target	Topline Data		
PAMREVLUMAB						
3	LAPC	LAPIS	284*	1H 2024 - OS endpoint		
3	DMD (non-ambulatory)	LELANTOS-1	99*	1H 2023		
3	DMD (ambulatory)	LELANTOS-2	73*	Mid-2023		
3	IPF	ZEPHYRUS-1	356*	Mid-2023		
3	IPF	ZEPHYRUS-2	340	TBD		
ROXADUSTAT						
3	MDS	MATTERHORN	160	1H 2023		





Thank You

For more information contact ir@fibrogen.com