



Headquartered in Switzerland – a biotech-hub of Europe – Idorsia is a high-potential biopharmaceutical company, specialized in the discovery, development, and commercialization of innovative small molecules, with the aim of transforming the horizon of therapeutic options.

We have a 25-year heritage of drug discovery, a broad portfolio of innovative drugs in the pipeline, an experienced team of over 750 professionals covering all disciplines from bench to bedside, and commercial operations in Europe and North America – the ideal constellation for bringing innovative medicines to patients.

Financial overview		Ni	ne Months	
in CHF millions, except EPS (CHF) and		US GAAP		Non-GAAP*
number of shares (millions)	2023	2022	2023	2022
Net revenues	131	43	131	43
Operating expenses	(275)	(653)	(517)	(621)
Operating income (loss)	(144)	(610)	(386)	(577)
Net income (loss)	(181)	(635)	(420)	(597)
Basic EPS	(1.02)	(3.58)	(2.36)	(3.36)
Basic weighted average number of shares	178.2	177.4	178.2	177.4
Diluted EPS	(1.02)	(3.58)	(2.36)	(3.36)
Diluted weighted average number of shares	178.2	177.4	178.2	177.4



Company Profile

overview



Company strategy

Innovation portfolio

Company milestones The full financial statements can be found in the Financial Report available on our corporate website.

* Idorsia measures, reports and issues guidance on non-GAAP operating performance. Idorsia believes that these non-GAAP financial measurements more accurately reflect the underlying business performance and therefore provide useful supplementary information to investors. These non-GAAP measures are reported in addition to, not as a substitute for, US GAAP financial performance.



Share Information

Idorsia was incorporated in March 2017 and listed on the SIX Swiss Exchange (ticker symbol: IDIA) in June 2017.

Idorsia Ltd is part of the following indices: SPI, SPIEX, SPI ESG, SXSLI, SXI Life Sciences, SXI Bio+Medtech, and SSIRT.

Idorsia is traded under the following symbols: Reuters IDIA.S / Bloomberg IDIA:SW.

Company Strategy

We will develop Idorsia into a leading biopharmaceutical company, with a strong scientific core. We have identified five key strategic priorities to ensure the company's success going forward.

Advance late-stage pipeline

We believe that our development compounds have the potential to significantly change treatment in their target diseases, resulting in medicines with substantial commercial potential.

Successfully launch our new products

In order to successfully bring our pioneering therapies to patients and to maximize the value of our innovations, we will continue to build and strengthen our global commercial organization.

Bring Idorsia to sustainable profitability

We are building Idorsia with a long-term focus and ambitious aspirations. By advancing our development pipeline and successfully launching our first products, we aim to bring Idorsia to sustainable profitability as soon as possible.

Fuel our pipeline with new discoveries

While launching our first marketed products and developing our late-stage clinical pipeline to bring our innovative therapies to patients, we also continue to discover new compounds.

Utilize state-of-the-art technologies to drive innovation

As we wish to remain at the cutting edge of science, it is vital that we consider innovative approaches and utilize state-of-the-art technologies at each stage of the process, from bench to bedside.

Company Profile

Financial overview

> Company strategy

Innovation portfolio

Company milestones

Innovation portfolio



Idorsia-led

Compound	Target indication	Mechanism of action	Status					
			P1	P2	P3	R	С	
QUVIVIQ™ (daridorexant)	Insomnia	Dual orexin receptor antagonist						Commercially available as QUVIVIQ in the US, Germany, Italy, Switzerland, Spain, the UK, Canada, Austria, and France; approved throughout the EU
TRYVIO™ (aprocitentan)	Systemic hypertension in combination with other antihypertensives	Dual endothelin receptor antagonist						Approved in the US, launch planned for H2 2024
JERAYGO™ (aprocitentan)	Resistant hypertension in combination with other antihypertensives	Dual endothelin receptor antagonist						Positive opinion from the European Committee for Medicinal Products for Human Use (CHMP) received in April 2024 – European Commission decision expected in approx. 2 months
Lucerastat	Fabry disease	Glucosylceramide synthase inhibitor						Phase 3 primary endpoint not met; open-label extension study ongoing
Daridorexant	Pediatric insomnia	Dual orexin receptor antagonist						Phase 2 in pediatric insomnia ongoing
ACT-1004-1239	Demyelinating diseases including MS	ACKR3/CXCR7 antagonist						Phase 2 in preparation
Sinbaglustat	Rare lysosomal storage disorders	GBA2/GCS inhibitor						Phase 1 complete
ACT-777991	Recent-onset Type 1 diabetes	CXCR3 antagonist						Phase 1 complete
ACT-1014-6470	Immune-mediated disorders	C5aR1 antagonist						Phase 1
IDOR-1117-2520	Immune-mediated disorders	Undisclosed						Phase 1 ongoing
IDOR-1134-2831	Clostridium difficile infection	Synthetic glycan vaccine						Phase 1 in preparation

Company Profile

Financial overview

Company strategy

> Innovation portfolio

Company milestones P1: Phase 1, P2: Phase 2, P3: Phase 3, R: Registration, C: Commercially available For more information about our portfolio please read our Innovation fact sheet

Innovation portfolio



Partner-led

Compound	Target indication	Mechanism of action	Partner	Status						
			Terms	P1	P2	P3	R	С		
Daridorexant	Insomnia	Dual orexin receptor antagonist	Nxera Pharma: license to develop and commercialize for Asia-Pacific region (excluding China)					NDA submitted in Japan		
Daridorexant	Insomnia	Dual orexin receptor antagonist	Simcere: license to develop and commercialize for Greater China region					Phase 3 ongoing		
Selatogrel	Acute myocardial infarction	P2Y ₁₂ inhibitor	Viatris: worldwide development and commercialization rights					Phase 3 "SOS-AMI" program ongoing		
Cenerimod	Systemic lupus erythematosus	S1P ₁ receptor modulator	Viatris: worldwide development and commercialization rights (excluding Japan, South Korea and certain countries in the Asia-Pacific region)					Phase 3 "OPUS" program ongoing		
Daridorexant	Posttraumatic stress disorder (PTSD)	Dual orexin receptor antagonist	US Department of Defense (DOD): Idorsia is supporting a clinical study sponsored by the US DOD to develop new therapies to treat PTSD					Phase 2 ongoing		
ACT-709478/ NBI-827104	Epileptic encephalopathy with continuous spike- and-wave during sleep (CSCW)	T-type calcium channel blocker	Neurocrine Biosciences: global license to develop and commercialize					Phase 2 OLE study ongoing		

Company Profile

Financial overview

Company strategy

> Innovation portfolio

Company milestones P1: Phase 1, P2: Phase 2, P3: Phase 3, R: Registration, C: Commercially available For more information about our partnerships please visit the **corporate website**



Company milestones

2024

March JERAYGO™ (aprocitentan) positive opinion from EU CHMP

March TRYVIO™ (aprocitentan) approved by the US FDA

March QUVIVIQ (daridorexant) launched in France

March Global development and commercialization agreement with Viatris for selatogrel and cenerimod

February QUVIVIQ (daridorexant) launched in Austria

2023

November QUVIVIQ (daridorexant) launched in Canada

October QUVIVIQ (daridorexant) launched in the UK

September QUVIVIQ (daridorexant) launched in Spain

June QUVIVIQ (daridorexant) launched in Switzerland

2022

December OPUS Phase 3 program to investigate cenerimod for the treatment of patients with systemic lupus erythematosus initiated

November QUVIVIQ (daridorexant) launched in Germany and Italy

November The Lancet and American Heart Association (AHA) late-breaking science session reports significant and sustained effect of aprocitentan on lowering blood pressure for patients with resistant hypertension

May Europe's first dual orexin receptor antagonist – QUVIVIQ (daridorexant) – granted approval to improve both nighttime symptoms and daytime functioning in adults with chronic insomnia disorder

May QUVIVIQ (daridorexant) 25 mg and 50 mg launched in the US for the treatment of adults with insomnia

January The Lancet Neurology reports impact of daridorexant on both nighttime symptoms and daytime functioning in adults with insomnia

Company Profile

Financial overview

Company strategy

Innovation portfolio

> Company milestones



2021

December Idorsia to further characterize lucerastat for Fabry disease by continuing the open-label extension of the Phase 3 MODIFY study

September Five Idorsia affiliates in key European markets (France, Germany, Italy, Spain, UK) established

June Initiation of Phase 3 registration study with selatogrel for the treatment of acute myocardial infarction

2020

July Establishment of Idorsia Pharmaceuticals US Inc. to perform commercial operations

May Global license agreement with Neurocrine Biosciences for the development and commercialization of ACT-709478

2019

November Collaboration Agreement with Halozyme for the development of a novel self-administered drug-device product for selatogrel

2017

June Idorsia opens its doors and is listed on SIX Swiss Stock Exchange

Company Profile

Financial overview

Company strategy

Innovation portfolio

> Company milestones

Idorsia is an independent biopharmaceutical company based on science and innovation. The company is specialized in the discovery, development, and commercialization of innovative small molecules, with the aim of transforming the horizon of therapeutic options. It is headquartered in Allschwil/Basel, Switzerland and is quoted on the SIX Swiss Exchange (tickersymbol: IDIA). All trademarks are legally protected by their respective owners.

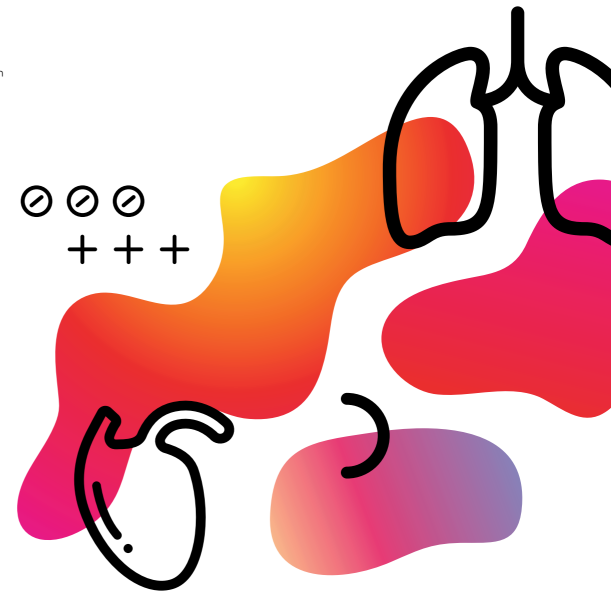
Disclaimer This fact sheet has the sole purpose to provide members of the public with general information about the activities of Idorsia. The forward-looking statements in this fact sheet are based on current expectations and belief of company management, which are subject to numerous risks and uncertainties.

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Company Profile

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