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Company presentation

April 2024

Forward looking statements

This presentation contains forward-looking statements that provide our expectations or forecasts of future events such as new product developments and regulatory approvals and financial performance.

Camurus is providing the following cautionary statement. Such forward-looking statements are subject to risks, uncertainties and inaccurate assumptions. This may cause actual results to differ materially from expectations and it may cause any or all of our forward-looking statements here or in other publications to be wrong. Factors that may affect future results include currency exchange rate fluctuations, delay or failure of development projects, loss or expiry of patents, production problems, unexpected contract, patent, breaches or terminations, government-mandated or market-driven price decreases, introduction of competing products, Camurus' ability to successfully market products, exposure to product liability claims and other lawsuits, changes in reimbursement rules and governmental laws and interpretation thereof, and unexpected cost increases.

Camurus undertakes no obligation to update forward-looking statements.

Camurus snapshot

Rapidly growing commercial stage company

Leader in opioid dependence treatment with Buvidal[®] weekly and monthly depots

Strong financial performance

Entered profitability in 2022



Advancing late-stage pipeline with blockbuster potential

Prospects for multiple new approvals in coming years in CNS and rare disease indications



Unique FluidCrystal[®] technology platform

Commercially validated, with a broad range of applications

LISTED ON NASDAQ STOCKHOLM TICKER CAMX; EMPLOYEES: 215+

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Successful 2023 lays foundation for continued profitable growth

Strengthened leadership in opioid dependence treatment

Brixadi[™] launched in the US for treatment of opioid use disorder

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- Positive results and progress in three Phase 3 studies
- NDA submission for Oclaiz[™] (CAM2029) in acromegaly
- Strong financials and operating performance



Outlook 2024

Total revenue SEK 1,740 – 1,860 million + 33 – 42% excl. one-time milestones 2023

Profit before tax **SEK 330 – 450 million** +131 – 215% excl. one-time milestones 2023



Positive financial development

Revenues





Operating results

FluidCrystal® extended-release technology

- $\checkmark\,$ Easy and convenient administration
- ✓ Rapid onset & long-acting release
- Controlled by composition, liquid crystal phase structure and biodegradation
- ✓ Applicable across substance classes
- Compatible with prefilled syringes, pen-injectors, and other advanced devices
- ✓ Manufacturing by standard processes



Sources: Tiberg F, et al. Chapter in Long Acting Injections and Implants, Advances in Delivery Science and Technology 2012; Tiberg F, et al. OnDrugDelivery 2010; Tiberg F, et al. Drug Del. Sci. Tech., 21 (1) 101-109 2011.

Broad and diversified product portfolio and pipeline





"Buvidal became my way out"

Justin, Buvidal patient in Australia

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Buvidal – game changing opioid dependence treatment

Weekly and monthly, subcutaneous buprenorphine for individualized treatment of opioid dependence within a framework of medical, social and psychological treatment in adults and adolescents 16 years or over¹

Demonstrated benefits to patients and society

- Superior treatment outcome and patient satisfaction²⁻⁵
- Blockade of subjective opioid effects from first dose³
- Reduced treatment burden and improved quality of life^{5,6}
- Decreased risk of diversion, misuse and pediatric exposure^{7,8}
- Reduced treatment costs⁹

¹ SmPC Buvidal May 2021; ²Lofwall et al. JAMA Int. Med. 2018;178(6); 764-773; ³Walsh et al, JAMA Psychiatry 2017;74(9):894-902; ⁴ Frost, M, et al. Addiction. 2019;114(8):1416-1426. <u>doi:10.1111/add.14636</u>; ⁵Lintzeris, N, et al. JAMA Network Open. 2021;4(5):e219041. <u>doi:10.1001/jaman.etworkopen.2021.9041</u>, ⁶Barnett et al Drug and Alcohol Dependence 2021; <u>https://doi.org/10.1016/j.drugalcdep.2021.108959</u>; ⁷EPAR for Buvidal; ⁸Dunlop, A. J., et al. Addiction. 2021. <u>https://doi.org/10.1111/add.15627</u>; ⁹Dunlop, A. Oral presentation at CPDD June 2020.

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Buvidal continues to grow in Europe, Australia and MENA

Sales growth across all markets

- Net sales 2023: SEK 1.3 billion; +39% vs 2022
 - Strong performance in key markets in UK, Nordics, Australia
 - Germany, Spain, France growing well from a lower base
- Est. 48,000 patients in treatment with Buvidal end 2023

Market expansion and LCM

- Reecent market authorizations in Kuwait and New Zealand (160 mg)
- Four market authorization applications under review
- Several pricing and reimbursement submissions under review
- New launches planned



Quarterly product sales

Brixadi[™] launch gains momentum in the US

Braeburn responsible for US commercialization

- Focused commercial organization of over 100 people

Wide access to Brixadi for the treatment of OUD

- Available in all 50 US states
- High payer coverage on par with competition

Accelerated uptake

- SEK 8.3 million royalty vs SEK 1.2 million in Q3
- Est. more than 2,000 US patients in treatment with Brixadi at end-2023¹

Peak market potential est. >USD 1 billion¹

- Brixadi has unique and competitive product profile
- Supportive market dynamics, and increasing awareness of LAI treatment options



US drug overdose deaths per 100,000 residents²

Opioid crisis in the US continues



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Significant treatment gap



Jan 19

High medical need in the US All drugs ••••• Opioids 40 000

181-22

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Jan 20 Jan 21

Brixadi – well differentiated in the US market

Convenient and flexible administration

- Weekly and monthly dosing
- Multiple dose strengths (four weekly, three monthly)
- Choice of multiple injection sites
- Thin needle and small dose volumes
- Room temperature stability (no cold chain required)

Strong scientific evidence base

 Superior efficacy and patient reported treatment satisfaction vs daily standard of care

Competitive label¹

- Switch from daily sublingual buprenorphine using conversion table for dose equivalency
- Direct initiation of treatment following a single dose of transmucosal buprenorphine

LAI features ²	Sublocade	Vivitrol	Buvidal. Brixadi
Weekly dosing	-		✓
Monthly dosing	\checkmark	✓	\checkmark
Multiple doses	_	_	\checkmark
Choice of inj. sites	_	_	\checkmark
Smallest needle	(19G)	(20G)	🗸 (23G)
Lowest dose volume	0.5–1.5mL	3.4mL	✓ 0.16–0.64mL
Room temp. storage	_	_	✓
Day one initiation	_	_	✓
Clin. data vs active contro	I _	_	✓
Launched	US, CAN, AUS,SE, FI, IL	US	US, EU, UK, AUS

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Communicating a growing evidence base

Active scientific conference agenda



Recent key publications

American Col Neuropsychop	kgar of harmacelogy	www.nature.com/
blockade by participants	N netic-pharmacodynamic a / buprenorphine subcutar with opioid use disorder n Come ² , Auf Aguar Zdove ² , Celine Sar ² , Marcus nd Fredrik Tiberg ⁴⁴⁵	neous depot (CAM2038)
ELSEVIER	Contents lites available at Sectore International Journal of D journal homepage: www.elicylar.comd	Drug Policy
buprenorphine longitudinal q Stephen Parkin ^{3,1} , Jos "National Addition Garry, Iunica"	g retention in treatment with (for opioid use disorder) as ualitative study une Norde A. ³ , John Strang ^{3,4} (Revery Norder Assentiate, RC) Objectades, 83 Alls, (Thereby (Norder Assentiate, Strands) (Stranger View and Mills, STR Jacobs)	a journey: Findings from a
Research report		
agonist trea	not entering opioid atment: A survey arisk opioid users	Nordic Studies on Alcohol and Drug O The Audit Ori O The Audit Ori Sector Studies of Audit Ori DOI: 10.1177/16/2523120477 journabaagepub conthornetha
Tuire Prami 🗓 Oriola, Espoo, Finland		

³Prami T et al, Nordic studies on drug and alcohol. 2023



Octreotide SC depot, CAM2029

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CAM2029 under development for three serious rare disease indications

- Acromegaly
- Gastroenteropancreatic neuroendocrine
 tumors (GEP-NET)
- Polycystic liver disease (PLD)

Designed for enhanced efficacy and patient convenience

Somatostatin receptor ligands established treatment

Wide use of somatostatin receptor ligands (SRLs)

- Antisecretory, antiproliferative, and immunomodulatory activity
- First-line medical treatment of acromegaly (ACRO) and neuroendocrine tumors (NET)¹
- SRLs also used in other fields of endocrinology and oncology, as well as in gastrointestinal, kidney and liver diseases²

SRL market dominated by long-acting injectables

- Key products: Sandostatin[®] LAR[®] (octreotide LAR) and Somatuline[®] Autogel[®] (lanreotide ATG)
- Market size approximately US\$ 3 billion³



Key limitations of current SSA therapies



First approved 1998

POSOLOGYMonthly intramuscular injectionDOSAGE FORM19-gauge 38mm needleDOSE10-40mg per month, 2.5mL

Limitations:

- Complex reconstitution
- Refrigerated storage
- · Large injection needle
- IM injection
- Dosing by trained HCP
- Limited exposure, and efficacy with incomplete symptom control^{1,2}

Somatuline[®] Autogel[®]



First approved 2007

POSOLOGYMonthly deep subcutaneous injectionDOSAGE FORM18-gauge 20mm needleDOSE60-120mg per month, 0.2-0.5mL

Limitations:

- · Refrigerated storage
- Large injection needle
- Deep SC injection
- Dosing by HCP (US)
- Limited efficacy with incomplete symptom control^{1,2}





First approved 2020

POSOLOGYTwice daily (BID)DOSAGE FORMOral capsuleDOSE40-80mg per day

Limitations^{3,4}:

- Significant food effect requiring dosing under fasting conditions twice daily
- Multiple DDIs
- Modest efficacy 42% of patients in pivotal trial lost biochemical control (IGF-1) after switch from injectable SSAs
- Not approved in NET

CAM2029 designed to address key limitations

Differentiating features

- ✓ Ready-for-use FluidCrystal[®] technology
- ✓ Rapid onset and long-acting octreotide release¹
- ✓ 5-fold octreotide bioavailability vs Sandostatin LAR with potential for improved efficacy^{1,2,3}
- ✓ State-of-the-art, pre-filled pen injector enabling convenient patient self-administration
- ✓ Subcutaneous administration with thin needle (22-gauge, 12.5mm)
- ✓ Room temperature storage



Source: ¹ Tiberg F, et al., Br J Clin Pharmacol. 2015; 80(3): 460-472; ² Constant dose; ³ Pavel M, et al., Cancer Chemotherapy and Pharmacology 2019; 83: 375-383; ⁴ Adelman D et al. Adv Ther. 2020;37(4):1608-19.



CAM2029 provides high SRL exposure

~5x higher octreotide plasma exposure for CAM2029 vs. Sandostatin LAR CAM2029 octreotide plasma levels in the range of immediate release octreotide



SRL – somatostatin receptor ligand; PK – pharmacokinetic; IR – immediate release; LAR – long-acting release; TID – three times per day; q4w – every 4 weeks Data on file

Status overview of CAM2029 programs by indication



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NDA submission in acromegaly following positive ACROINNOVA Phase 3 study results

Key milestones achieved for CAM2029

- ✓ Positive ACROINNOVA 1 Phase 3 results¹
 - Demonstrating superior biochemical control vs placebo
 - Improved convenience and quality of life vs SoC
 - Safety profile consistent with 1st generation SRLs
- Positive ACROINNOVA 2 interim Phase 3 results² \checkmark
 - · Reinforcing long-term safety and effectiveness
 - Improved symptom control, treatment satisfaction and quality of life scores vs SoC at baseline
- ✓ Population PK and PKPD models developed
- Positive pre-NDA meetings \checkmark
- NDA submission of Oclaiz[™] in acromegaly3 \checkmark
 - Submission date 21 December 2023

Efficacy demonstrated in ACROINNOVA 1 & 2^{1,2}





Proportion of responders over time (IGF-1≤ULN)



CAM2029 has an attractive product profile in acromegaly



Once-monthly self-administration with prefilled pen



- Improved convenience and treatment satisfaction^{1,2}
- Long-acting release with ~5X octreotide bioavailability^{3,4}



High rates of biochemical control¹



Improved symptom control & quality of life²





* The Acromegaly Index of Severity (AIS) score was calculated as the sum of the scores for the six symptoms of headache, sweating, fatigue, joint pain, paresthesia and soft tissue swelling. The AIS score ranges from 0 (no symptoms) to 18 (severe symptoms)

ACROINNOVA 1 Phase 3 RCT efficacy and safety trial

ACROINNOVA 1 trial design

 24-week, randomized, double blind, placebo-controlled trial

Key eligibility criteria:

- Patients with acromegaly on treatment with a stable dose of octreotide LAR or lanreotide ATG for at least 3 months with
- IGF-1 levels ≤1xULN at screening

Primary endpoint:

Proportion of patients with mean IGF-1 ≤1xULN (week 22 and 24)

Key secondary endpoints:

- Proportion of patients with mean IGF- 1 levels ≤1xULN , incl. patients with decreased dose
- Proportion of patients with mean IGF-1 levels ≤1xULN and GH cycle levels <2.5 µg/L

Secondary endpoints, e.g,:

- Time to loss of IGF-1 response
- IGF-1 and GH over time and change from baseline
- Clinical signs and symptoms (AIS score)
- Patient satisfaction and treatment satisfaction (PSS and TSQM)
- Acromegaly quality of life (AcroQoL)
- Self-injection assessments (SiAQ)
- Plasma concentrations of octreotide
- Safety and tolerability



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Completed patient recruitment in Phase 3 SORENTO study of CAM2029 in GEP-NET

Enrollment across 12 countries exceeding randomization target (302)



SORENTO assessing CAM2029 superiority in PFS

Randomized, active-controlled Phase 3 trial

- Randomized, multi-center, open-label, active-controlled Phase 3 trial of CAM2029 vs. long-acting octreotide or lanreotide in patients with GEP-NET
- Single trial fulfilling regulatory requirements for safety and efficacy

Patient population

 Patients with confirmed, advanced and well-differentiated GEP-NET (grade 1 to grade 3)

Primary endpoint

- Superiority in progression free survival, PFS, vs. standard of care (first-line medical treatment)
- Assessed after 194 documented PFS events

Secondary endpoints include

- Overall survival
- PROs (e.g., treatment satisfaction, quality of life)
- Plasma concentrations of octreotide
- Safety



SORENTO

Clinical Phase 2/3 study in PLD fully recruited

POSITANO trial to assess efficacy and safety

- 53-week randomized, placebo-controlled, three-arm trial
 - Randomization of 71 patients completed in February 2024
 - Primary endpoint is liver volume change
 - Key secondary endpoint is Camurus' developed PRO, PLD-S
 - Multiple secondary endpoints, incl. quality of life, safety, etc.
- Open label extension extended to 120 weeks
 - · Offer continued treatment in patients with expected benefits

Large unmet medical need in PLD

- Severe quality-of-life implications for patients with symptomatic PLD
- No labelled option available







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CAM2029 progressing towards market with key upcoming key milestones 2024/25

AcroInnova[™]

Pivotal randomized placebo controlled and long-term safety trials in acromegaly

- ✓ Positive ACROINNOVA 1 results
- ✓ Positive ACROINNOVA 2 interim results
- ✓ NDA submission
- ✓ NDA acceptance for review
- MAA submission H1 2024
- NDA approval decision expected Q4 2024
- **US launch Q1 2025**

SORENTO

Subcutaneous Octreotide Randomized Efficacy in Neuroendocrine TumOrs

- ✓ SORENTO Phase 3 start Q4 2021
- ✓ SORENTO fully enrolled Q4 2023
- □ Topline result H1 2025
- NDA/MAA submission H2 2025

<u>posíτano</u>™

Polycystic liver Safety and efficacy TriAl with subcutaneous Octreotide

- ✓ POSITANO Phase 2/3 Q2 2022
- ✓ POSITANO fully enrolled Q1 2024
- □ Topline result H1 2025

High market potential for CAM2029 – largest opportunity in GEP-NET

Attractive specialty pharma opportunity

- Blockbuster potential in NET
- Highly concentrated target audiences
- Differentiated product features
- Switch from established first-line treatments

CAM2029 peak sales estimates from third party market research¹⁻⁴

	TERRITORY	PATIENT POPULATION	EST. PEAK PATIENT SHARE	EST. PEAK SALES
ACRO	EU/AUS	16,500 ⁴	20 – 35%	€30 – 65 million
	US	10,000	25 – 40%	\$150 – 280 million
NET ¹	EU/AUS	68,000 ⁴	30%	€300 – 400 million
	US	37,000	40%	\$1,200 – 1,500 million
PLD ¹	EU/AUS	15-18,000 ⁴	30 – 40%	€80 – 100 million
	US	12-13,000	30 – 40%	\$200 – 300 million

¹Globe Life Science Aug 2022, data on file; ²Globe Life Science 2020, data on file; ³Assuming €10-12.5ks (EU/AUS) and \$80-70K (US) per year net pricing in acromegaly, €15-20k (EU/AUS) and \$80-100K (US) per year net pricing in NET, and €17.5k (EU/AUS) and \$80K (US) per year net pricing in PLD; ⁴Patient numbers extrapolated from 5EU estimates by assuming same prevalence across European countries and Australia



Building commercial infrastructure in the US

US launch preparations Oclaiz[™] in acromegaly

Key activities

- Camurus Inc. fully operational
- President Camurus US appointed
- In-depth market research
- Medical affairs activities
- Payor engagement
- Distribution model

Key scientific conferences for CAM2029 in 2024



Regulatory timeline:



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Strong foundation for continued value creation



Buvidal growth in Europe and Australia



Positive launch momentum for Brixadi in the US

Pipeline progress towards new approvals and launches



Establishing a US commercial organization



Strong financial position to support sustainable growth



Camurus AB | Ideon Science Park, SE-223 70 Lund, Sweden

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VALUE AND

Progress towards Camurus' Vision 2027

Status update end-2023 following one year of execution towards the five-year vision



Five-fold revenue growth (from 2022)



Establishment of US commercial infrastructure **Approvals** for four R&D pipeline programs

~50%

Operating margin around 50%

5x revenue growth in 5 years

SEK 1.7bn 2023

□ SEK 4.5 billion in 2027

Buvidal patients grew 33%

48,000 in 2023

□ >100,000 patients in 2027

Brixadi, opioid use disorder

- ✓ US launch in September 2023
- □ >\$1 billion peak sales potential

US commercial infrastructure

Preparing for Oclaiz™ launch

- ✓ Camurus Inc. fully operational
- ✓ Behshad Sheldon appointed President Camurus US
 ❑ Launch-ready Q4 2024

Accelerated commercial build-up

- ✓ Strengthened financial position
- Accelerate commercial readiness in NET and PLD

New approvals

1 of 4

Brixadi, opioid use disorder

 $\checkmark~$ US approved in May 2023

Oclaiz[™] (CAM2029) in acromegaly

- ✓ NDA submitted in December 2023
- US approval decision exp. Q4 2024

CAM2029 GEP-NET

- ✓ Completed Phase 3 recruitment in Q4 2023
- □ NDA submission est. 2025

Operating margin

31% in 2023

□ ~50% in 2027

Operational excellence

- \checkmark Increased gross margin
- Disciplined capital allocation to invest in the pipeline and commercialization

Supported by inorganic growth

- ✓ Proceeds of SEK 1.1 billion directed share issue in January 2024
- Grow and diversify revenues through partnerships and acquisition

Key milestones coming 12 months

R&D Pipeline

- ✓ Completed recruitment in POSITANO study in PLD
- □ FDA acceptance for review of Oclaiz[™]NDA
- □ MAA submission of CAM2029 in acromegaly to EMA
- □ FDA approval of Oclaiz[™] in acromegaly
- □ Topline results SORENTO study in GEP-NET
- □ Topline results POSITANO study in PLD
- □ Start new clinical program

Commercial and corporate development

- ✓ Directed share issue raising gross proceeds of SEK 1.1 billion
- □ US commercial organization fully established
- □ Business development and inorganic growth
- □ US launch of Oclaiz[™] in acromegaly





Analysts

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Shareholders and analyst coverage

Shareholders as of 28 March 2024	Number of shares	% of capital	% of votes
Sandberg Development AB	21,875,692	37.97	37.97
Fjärde AP-fonden	2,610,766	4.53	4.53
Avanza Pension	1,801,327	3.13	3.13
Swedbank Robur Fonder	1,697,879	2.95	2.95
Fredrik Tiberg, CEO	1,615,000	2.80	2.80
JP Morgan Chase Bank	1,486,936	2,58	2,58
State Street Bank and Trust	1,326,203	2.30	2.30
Handelsbankens fonder	1,283,328	2.23	2.23
The Bank of New York Mellon SA/NV	1,156,632	2.01	2.01
Afa Försäkring	646,293	1.12	1.12
The Bank of New York Mellon	590,422	1.02	1.02
CS Client Omnibus	586,293	1.02	1.02
Norges bank	560,987	0.97	0.97
SEB Investment Management	558,297	0.97	0.97
SEB	512,979	0.89	0.89
Other shareholders	19,305,584	33.51	33.51
In total	57,614,618	100.0	100.0

Experienced and committed management team

R.	Fredrik Tiberg, PhD President & CEO, CSO In Company since 2002 Holdings: 1,615,000 shares and 102,000 employee options	Education: M.Sc. in Chem. Eng., Lund Institute of Technology, PhD and Assoc. Prof. Physical Chemistry, Lund University. Previous experience: More than 20 years executive leadership experience from the pharmaceutical industry. Professor Physical Chemistry, Lund University; Visiting Professor at Oxford University; Section Head, Institute for Surface Chemistry.	and the second s	Jon Garay Alonso Chief Financial Officer In Company since: 2022 Holdings: 1,450 shares & 57,750 employee options	Education: Bachelor in Business Administration by Universidad Comercial de Deusto. Executive MBA by IESE Business School. Previous experience: More than 20 years experience from Finance within pharmaceutical and medtech companies, incl. Baxter, Gambro, Convatec, Bristol Myers Squibb.
0	Richard Jameson <i>Chief Commercial Officer</i> In Company since : 2016 Holdings : 29, 193 shares and 57, 750 employee options	Education: B.Sc. in Applied Biological Sciences from University West of England Previous experience: General Manager, UK & Nordics for Reckitt Benckiser (2010 – 2013) and Area Director Europe, Middle East and Africa for Indivior (2013 – 2016).	(all)	Fredrik Joabsson, PhD Chief Business Dev. Officer In Company since 2001 Holdings: 50, 170 shares and 38,500 employee options	Education: M.Sc. in Chemistry, PhD in Physical Chemistry, Lund University Previous experience: More than 20 years of experience in pharmaceutical R&D, business development, alliance management and investor relations.
Ser la	Markus Johnsson Senior VP R&D In Company since: 2003-2017, 2019- Holdings: 21,000 shares & 23,500 employee options	Education: Ph.D. in physical chemistry and M.Sc. in chemistry from Uppsala University. Previous experience: More than 20 years of experience from pharmaceutical development and project management		Maria Lundqvist Head of Global HR In Company since 2021 Holdings: 38,500 employee options	Education: B.Sc: in Business and Economics, Uppsala University. Previous experience: More than 20 years of experience of leadership roles within Human Resources, including HR Director Nordics at Teva Pharmaceuticals and HR positions at Tetra Pak, Vestas and AstraZeneca.
60	Torsten Malmström, PhD Chief Technical Officer In Company since 2013 Holdings: 46,858 shares and 38,500 employee options	Education: M.Sc. in Chemistry, PhD in Inorganic Chemistry, Lund University Previous experience: More than 20 years of experience from pharmaceutical R&D including Director Pharmaceutical Development at Zealand Pharma, Director of Development at Polypeptide, Team Manager at AstraZeneca.	6	Annette Mattsson VP Regulatory Affairs In Company since: 2017 Holdings: 3,004 shares and 38,500 employee options	Education: Bachelor of Pharmacy, Uppsala University and Business Economics, Lund University Previous experience: More than 25 years of experience within regulatory affairs, including European RA Director/Global RA Lead at AstraZeneca and Global RA Lead at LEO Pharma.
20	Alberto M. Pedroncelli Chief Medical Officer In Company since 2023 Holdings: 1,000 shares and 20,000 employee options	Education: MD University of Milan. Ph. D. endocrinology post- graduate school University of London Previous experience: Head of Clinical Development and Medical Affairs Recordati, Senior Leadership positions Novartis, clinician and research fellow Dept. Endocrinology, University Hospital Bergamo, Italy		Behshad Sheldon President Camurus Inc. In Company since 2024 Holdings: 1,000 shares	Education: B.Sc. in Neuroscience from University of Rochester Previous experience: More than 25 years of experience from the international pharmaceutical industry, including President & CEO of Braeburn Pharmaceuticals and senior positions within Smithkline Beecham, Bristol-Myers Squibb and Otsuka Pharmaceuticals.
P	Agneta Svedberg VP Clinical & Regulatory Dev. In Company since: 2015 Holdings: 22,987 shares and 38,500 employee options	Education: M.Sc. In Radiophysics and B.Sc. In Medicine from Lund University, Executive MBA from Executive Foundation Lund Previous experience: More than 25 years of experience in drug development, incl. as COO at Zealand Pharma, CEO of Cantargia, Senior VP Clinical Development at Genmab.			