



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.

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Camurus snapshot



Rapidly growing commercial stage company

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



Advancing late-stage pipeline with blockbuster potential

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



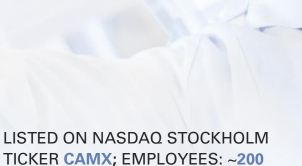
Strong financial performance

Entered profitability in 2022



Unique FluidCrystal® technology platform

Commercially validated, with a broad range of applications



Significant recent progress



Positive financial development

- ✓ High double-digit year-on-year revenue growth
- ✓ Sustained profitability
- ✓ Robust cash position SEK 1.15 B end Q3 2023 – no debt
- ✓ Raised Full Year 2023 guidance



Commercial development

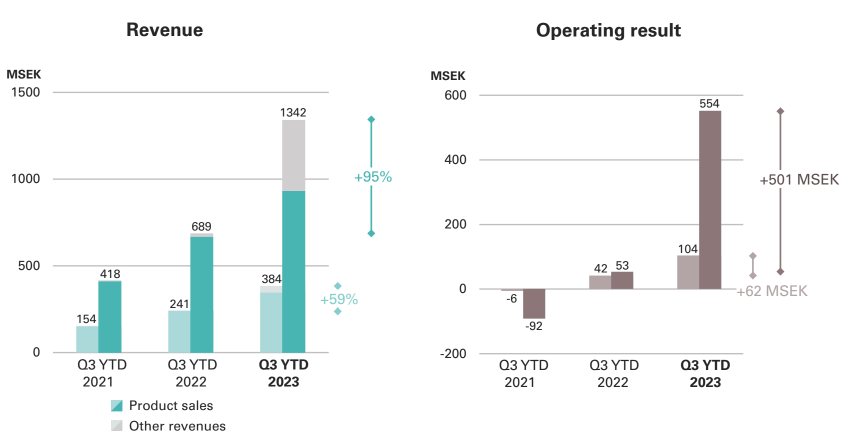
- ✓ Strengthened leadership in LAI treatment of opioid dependence
- ✓ High growth of Buvidal sales to SEK 346 million, up 44% YoY
- ✓ Brixadi™ launched in the US by Braeburn for the treatment of opioid use disorder



Pipeline progress

- ✓ Positive ACROINNOVA 2 Phase 3 results for CAM2029 in acromegaly
- ✓ CAM2029 pre-NDA meeting for acromegaly with the US FDA
- ✓ Recruitment in SORENTO Phase 3 trial GEP-NET nearing completion

Third quarter 2023 – Strong revenue growth and result

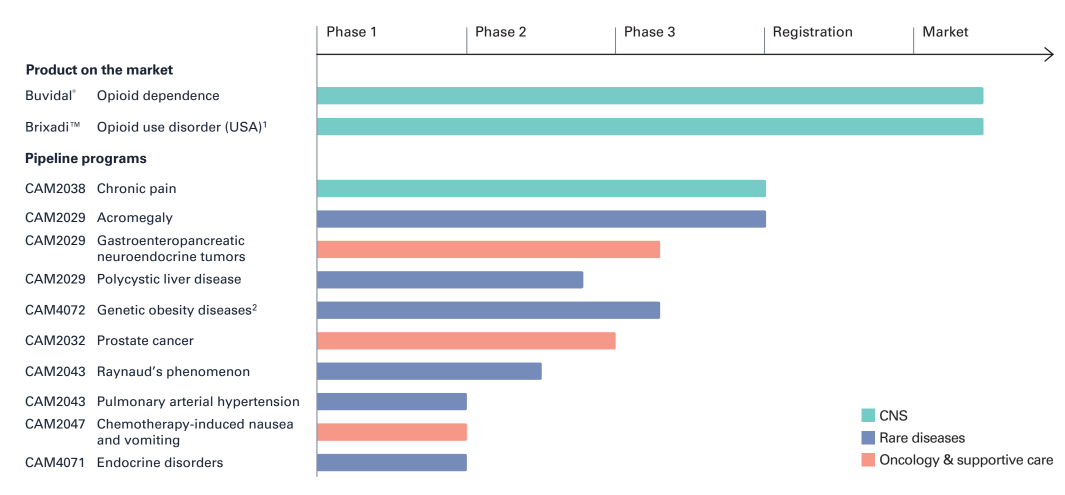


Cash position
SEK 1,154 million
+122% vs Q3 2022



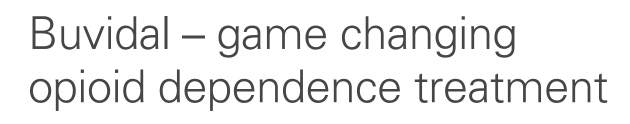


Broad and diversified product portfolio and pipeline



¹Licensed to Braeburn in North America; ²Licensed to Rhythm Pharmaceuticals worldwide





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





Buvidal continuing to grow in Europe, Australia and MENA

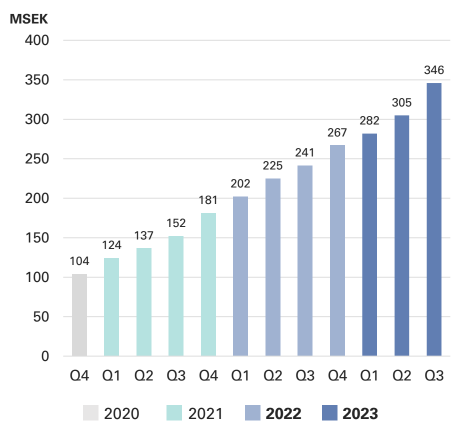
Continued market penetration

- Robust double-digit YoY sales growth
 - Q3 net sales: SEK 346 million; +44% YoY, +13% QoQ
- Strong performance across markets including the UK, Nordics, Germany, Austria and Spain
- Est. 45,000 patients in treatment with Buvidal end Q3
- Target more than 100,000 in 2027

Regulatory and market expansion processes

- Buvidal launched in Italy
- Four regulatory and four reimbursement submissions progressing
- New markets planned

Quarterly product sales





US launch of Brixadi in opioid use disorder

Braeburn responsible for US commercialization

Focused commercial organization of over 100 people

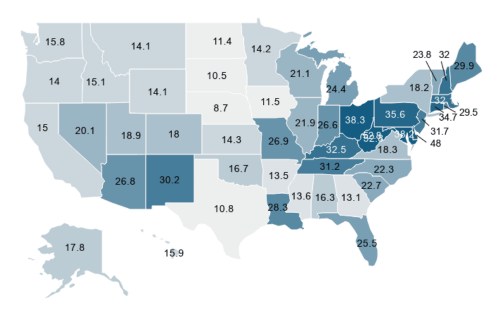
Launch initiated 5 September 2023

- Brixadi available in all 50 US states; in several cases with unrestricted access through Medicaid
- Increasing coverage through private payers
- First royalty revenue received by Camurus

High peak market potential est. >USD 1 billion¹



US drug overdose deaths per 100,000 residents⁴

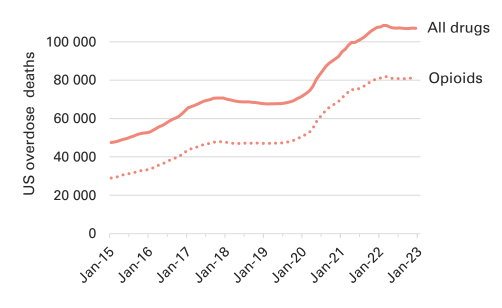




Opioid crisis in the US continues

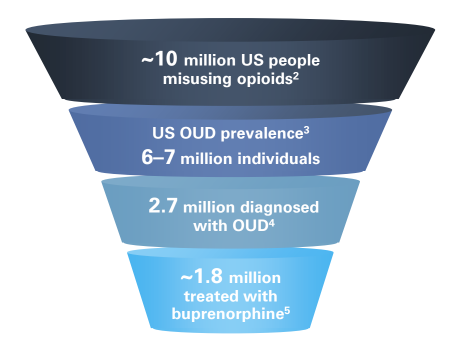
High medical need in the US

~80,000 annual deaths in opioid overdoses¹



12 month-ending period

Significant treatment gap





Positive market dynamics in the US

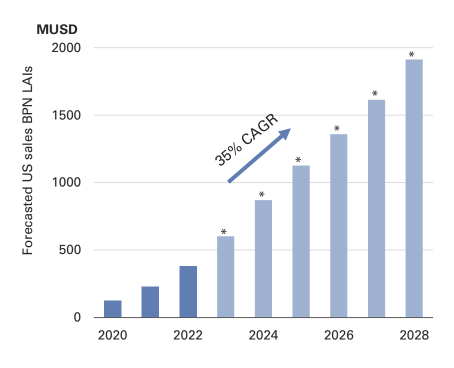
Recent initiatives to address treatment hurdles

- President Biden's Unity Agenda¹
- Improved funding²
- Removal of DATA 2000 waiver and number of patients
 HCPs can treat³
- Expanded access to treatment in criminal justice system⁴
- Est, six million people with opioid use disorder⁵

Long-acting injectable buprenorphine growing

- Currently low patient share (<5%6) but rapidly growing
- Brixadi entering market with competitive and well differentiated product profile

Positive outlook on BPN LAI market growth⁷



* Forecast based on analyst consensus data



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	✓	✓	✓
Multiple doses	_	_	✓
Choice of inj. sites	_	_	✓
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



Octreotide SC depot

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





CAM2029 targeting USD 3-billion SRL market

SRLs established treatment with limitations

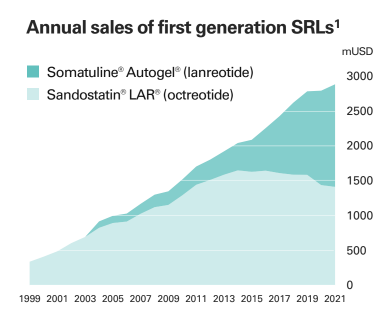
- First-line treatment of acromegaly and neuroendocrine tumors (NET)
- Established safety and efficacy profile
- Potential for significant improvements of efficacy and patient convenience

CAM2029 best-in-class treatment potential

- Convenient self-administration with state-of-the-art pen device



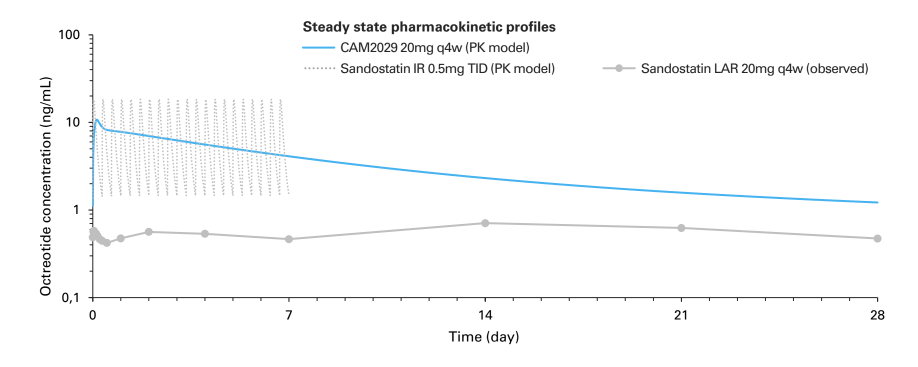
- 5-fold increase of octreotide plasma exposure (dose adjusted)
- Potential for improved disease control and treatment outcomes





CAM2029 provides high SSA exposure

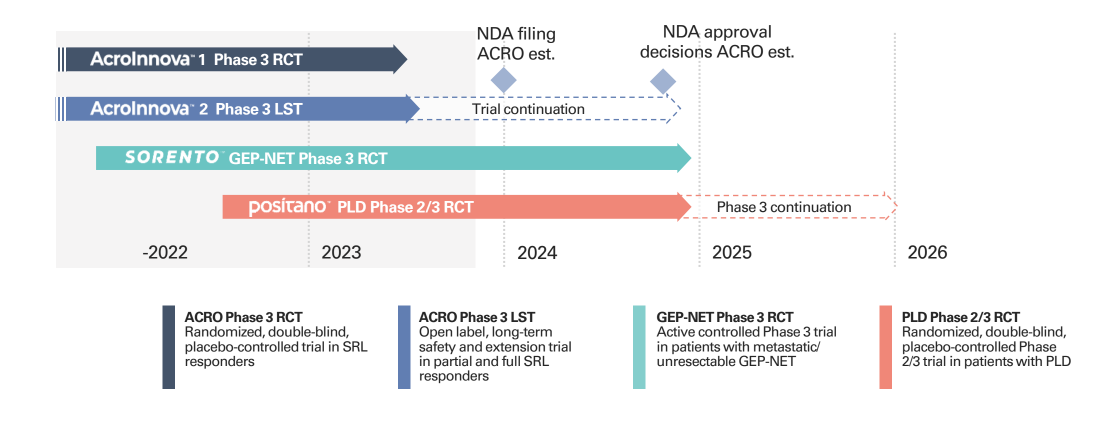
~5x higher octreotide plasma exposure for CAM2029 vs. Sandostatin LAR CAM2029 octreotide plasma levels in the range of immediate release octreotide



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



CAM2029 Phase 3 programs advancing





Positive topline results for CAM2029 in two Phase 3 trials¹

Key results from ACROINNOVA 1

- Met primary and key secondary endpoints of superior IGF-1 response rate versus placebo
 - Confirmed by sensitivity and supportive analyses
- / IGF-1 and GH well-controlled over time
 - Measured pre-dose, at trough octreotide conc.
- Increased treatment satisfaction scores (TSQM) versus standard of care (SoC) at baseline
- Increased quality of life (AcroQoL) scores versus SoC at baseline
- Safety profile comparable to first-generation SRLs, octreotide LAR and lanreotide ATG

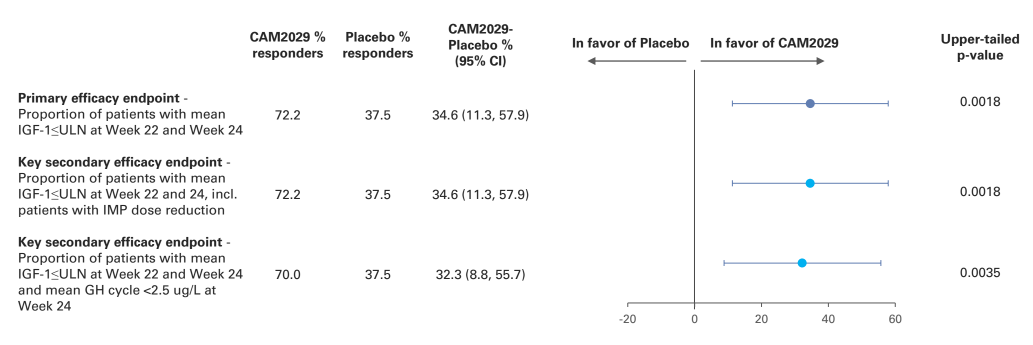
Key interim results from ACROINNOVA 2

- ✓ Affirmative safety profile over 52-weeks
 - No new or unexpected safety findings
- Increased IGF-1 response vs baseline in uncontrolled patients and treatment naïve patients after washout
- ✓ Stable IGF-1 response in controlled roll-over patients
- Decrease in symptom scores vs SoC at baseline
- Increased treatment satisfaction (TSQM) and quality of life scores (AcroQoL, EQ-5D-5L) vs SoC at baseline
- Improved injection experience by self-injection assessment questionnaire (SiAQ) scores



ACROINNOVA 1 – CAM2029 superior to placebo for IGF-1 and GH response

Primary and key secondary endpoints met with high statistical significance (ITT)



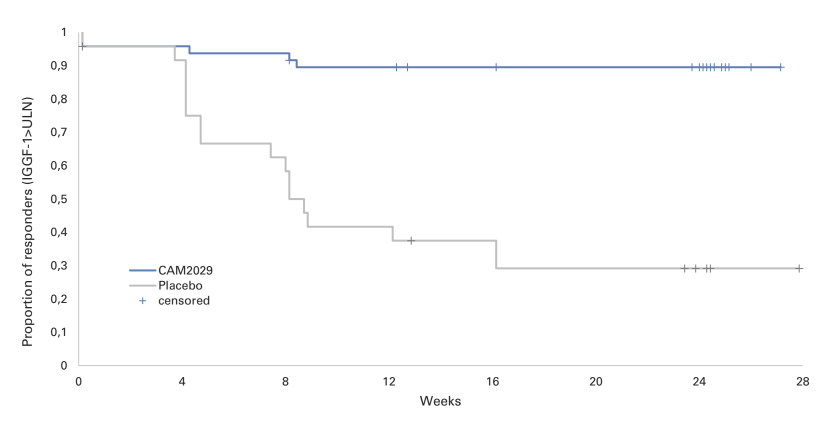
Difference in proportion (%) and 95% CI (CAM2029-Placebo)

ITT – intention-to-treat analysis set.



ACROINNOVA 1 High statistical difference in time to loss of response

Cox regression analysis (ITT): Hazard ratio=0.1; p<0.0001

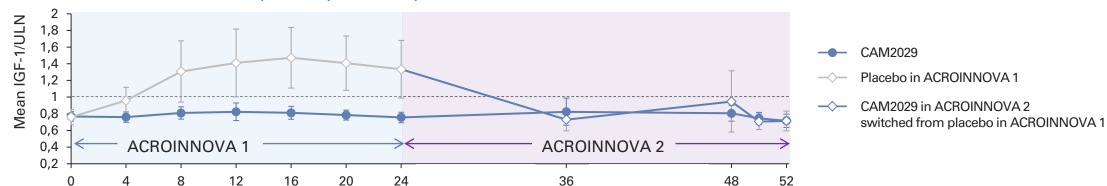




Patients retained or regained IGF-1 control with CAM2029

Weeks

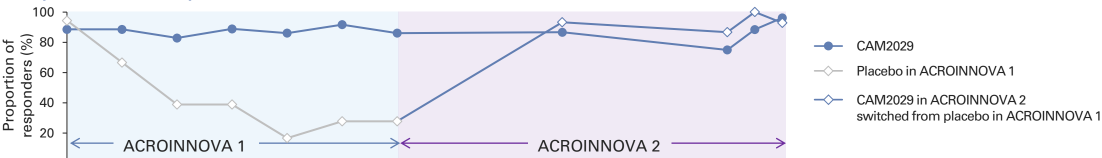
IGF-1 values over time (mean, 95% CI)



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

12

16



36

52

48

Patients with data at the cut-off timepoint for the interim analysis (N=54). All values are pre-dose and time points are nominal

20

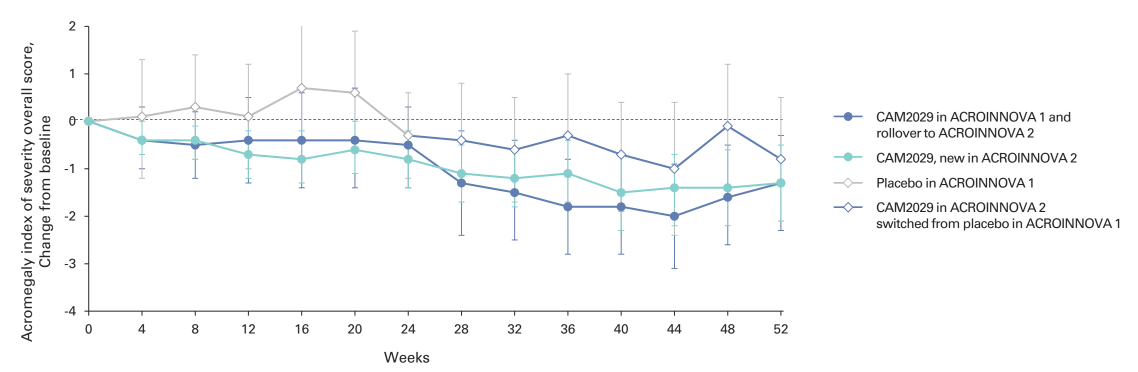
24

Weeks



ACROINNOVA 2 Decreasing acromegaly symptoms over time

Change from SoC treatment baseline in Acromegaly Index of Severity Score (6 symptoms)*



^{*} The AIS overall 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 overall score ranges from 0 (no symptoms) to 18 (severe symptoms)



Progress in three clinical programs

AcroInnova

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

- ✓ Topline results reported from two Phase 3 trials
- ✓ Positive ACROINNOVA 1 results 20 June 2023
- ✓ Positive ACROINNOVA 2 results 17 July 2023
- ✓ Pre-NDA meeting
- NDA submission in acromegaly planned around end of 2023
- MAA submission H1 2024

SORENTO

Subcutaneous Octreotide Randomized Efficacy in Neuroendocrine TumOrs

- ✓ SORENTO Phase 3 trial progressing well
- Est. completion of patient enrollment in Q4 2023 (target 302 patients)
- ☐ Primary endpoint readout after 194 PFS events
- Est. NDA/MAA GEP-NET submissions in 2025



Polycystic liver Safety and efficacy TriAl with subcutaneous Octreotide

- ✓ Orphan drug designation (US)
- New PROs developed and aligned with FDA
- ✓ Phase 2/3 trial ongoing
- Est. completion of enrollment around end of year 2023
- ☐ Topline results end 2024/early 2025



Preparing for commercialization of CAM2029

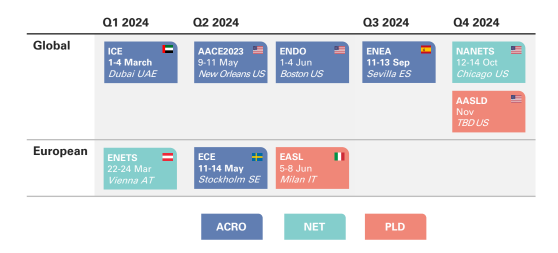
Setting up US commercial infrastructure

- ✓ Camurus Inc. operational
- ☐ Establishment of US commercial operations
 - Distribution model
 - Medical affairs
 - Commercial development
 - including market research
 - Compliance framework
- ☐ Launch ready Q4 2024

Manufacturing and devices

- √ Process validation completed
- ✓ Stability studies completed for submissions
- ✓ Human factor engineering studies

Key scientific conferences 2024



Significant market potential for CAM2029

Attractive opportunity

- Block buster potential in NET
- Highly concentrated target audiences
- Differentiated product features
- Switch opportunity 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 ¹	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

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Key takeaways Strong revenue growth and profitability YTD 2023 Brixadi launched in the US for the treatment of opioid use disorder Continued growth of Buvidal in Europe and Australia Advancing pipeline with positive Phase 3 results in acromegaly Camurus Inc. operational in the US





Reported Q3 profit and loss

MSEK	Jul – Sep 2023	Change vs. 2022	CER Change vs. 2022	YTD Jan – Sep 2023	Change YTD vs. 2022	CER Change YTD vs. 2022
Total revenues out of which CAM2038 milestones	384 36	+59%	+49%	1 342 406	+95%	+86%
Gross margin % GM Product Sales	352 <i>90,8%</i>	+162bps <i>+75bps</i>	+165bps <i>+79bps</i>	1 253 <i>90,4%</i>	+425bps <i>+135bps</i>	+473bps <i>+97bps</i>
Marketing and distribution costs	-94	+41%	+34%	-264	+35%	+29%
Administrative expenses	-10	0%	-6%	-32	+22%	+17%
Research and development costs	-148	+39%	+32%	-408	+20%	+15%
Other operating expenses	5	-	-	5	-	_
Operating result	104	+63 MSEK	+46 MSEK	554	+501 MSEK	+453 MSEK

Key milestones in 2023

Advancing the pipeline

- ✓ Topline ACROINNOVA 1 Phase 3 efficacy results in acromegaly
- ✓ Positive ACROINNOVA 2 Phase 3 long-term safety study results
- ✓ Pre-NDA meeting for CAM2029 in acromegaly
- □ NDA submission of CAM2029 in acromegaly
- □ Completed recruitment in SORENTO study in GEP-NET
- □ Completed recruitment in POSITANO study in PLD
- □ Topline Phase 3 PK results for weekly setmelanotide by Rhythm

Commercial and corporate development

- ✓ US approval and launch of Brixadi in opioid use disorder
- ✓ Establishment of US commercial infrastructure
- Business development and inorganic growth



Shareholders and analyst coverage

Shareholders as of 31 October 2023	Number of shares	% of capital	% of votes
Sandberg Development AB	21,875,692	39.4	39.4
Fjärde AP-fonden	3,116,100	5.6	5.6
Avanza Pension	2,166,414	3.9	3.9
Fredrik Tiberg, CEO	1,600,000	2.9	2.9
Swedbank Robur Fonder	1,280,000	2.3	2.3
State Street Bank and Trust	1,249,300	2.2	2.2
JP Morgan Chase Bank	1,042,313	1.9	1.9
Handelsbankens fonder	910,522	1.6	1.6
The Bank of New York Mellon SA/NV	886,453	1.6	1.6
Afa Försäkring	792,708	1.4	1.4
Svenskt Näringsliv	650,000	1.2	1.2
Öhman Fonder	555,490	1.0	1.0
Lancelot Avalon Master	494,847	0.9	0.9
Backahill Utveckling	487,359	0.9	0.9
Camurus Lipid Research Foundation	486,350	0.9	0.9
Other shareholders	17,945,270	24.2	24.2
In total	55,538,818	100.0	100.0



Analysts

Carnegie Erik Hultgård

DNB Potrik Lin

Patrik Ling

Handelsbanken Suzanna Queckbörner Mattias Häggblom

Jefferies
James Vane-Tempest

Nordea Viktor Sundberg

Pareto Dan Akschuti

Bryan Garnier Alex Cogut



Experienced and committed management team



Fredrik Tibera, PhD President & CEO, CSO In Company since 2002 Holdings: 1,600,000 shares. 15.000 subscription warrants & 102,000 employee options

Education: M.Sc. in Chem. Eng., Lund Institute of Technology, Ph.D. 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.



Jon Garav 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.



Maria Lundqvist Head of Global HR In Company since 2021 Holdings: 38,500 employee options

Education: B.Sc.: in Business and Economics, Uppsala

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.



Richard Jameson Chief Commercial Officer In Company since: 2016 Holdings: 29, 193 shares, 8,000 subscription warrants 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).



Fredrik Joabsson, PhD Chief Business Dev. Officer In Company since 2001 Holdings: 50,170 shares & 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 and alliance management.



Markus Johnsson Senior VP R&D In Company since: 2003-2017, 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



Torsten Malmström, PhD Chief Technical Officer In Company since 2013 Holdings: 46.858 shares & 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.



Annette Mattsson VP Regulatory Affairs In Company since: 2017 Holdings: 2.004 shares, 1.000 subscription warrants &

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.



Alberto M. Pedroncelli Chief Medical Officer In Company since 2023 Holdings: 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



Agneta Svedberg VP Clinical & Regulatory Dev. In Company since: 2015 Holdings: 22.987 shares & 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.



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

ACROINNOVA 1 CAM2029 once monthly (HS-18-633) Screening Possibility to roll over R Placebo once monthly to ACROINNOVA 2 Stable dose (HS-19-647) and octreotide or N=72, 2:1 continue CAM2029 lanreotide Rescue with standard of care Double-blind treatment phase 4-8 weeks Day 1 Week 24

Statistical assumption primary endpoint:

 90% power to show treatment difference with 80% response for CAM2029 vs 40% response for placebo, based on Chi-squared test (with continuity correction)



ACROINNOVA 2 Phase 3 long-term safety and extension trial

ACROINNOVA 2 trial design

- 52-week, open-label, long-term safety and extension trial

Patient population

- New patients in trial; IGF-1<2xULN (n=81)
- Roll-over CAM2029 patients; IGF-1≤1xULN (n=36)
- Roll-over placebo patients; IGF-1≤1xULN (n=18) from ACROINNOVA 1

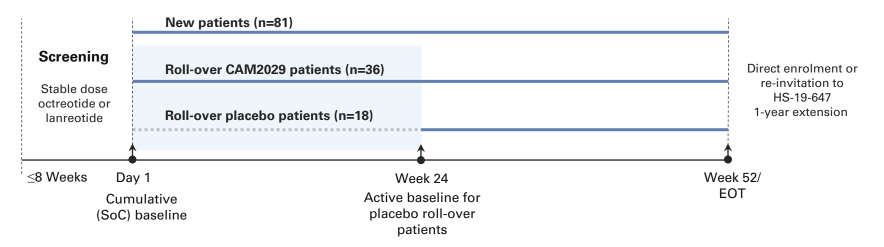
ACROINNOVA 2 (HS-19-647)

Primary endpoint:

Long-term safety and tolerability

Secondary endpoints:

- Biochemical response (IGF-1, GH)
- Mean IGF-1 and GH over time
- Clinical signs and symptoms (AIS)
- Patient and treatment satisfaction (TSQM)
- Quality of life (AcroQoL, EQ-5D-5L)
- Self-Injection Assessment Questionnaire (SiAQ)
- Octreotide concentrations





SORENTO: Largest Phase 3 trial of SSA in NET

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 (unresectable and/or metastatic), 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 progression events

Secondary endpoints include

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

