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### Orexo in brief



#### **US Pharma**

Commercial US Pharma platform since 2013, incl. market access team and sales representatives who on a daily basis visit physicians, medical clinics and minor hospitals.

#### **HQ & Pipeline**

Development of improved drugs based on well-known substances combined with innovative proprietary Drug Delivery technologies, such as amorphOX<sup>®</sup>.



Evidence-based digital therapies grounded in cognitive behavioral therapy techniques, offer better treatment access for patients and improve their outcomes.





#### **ZUBSOLV®** for the treatment of opioid use disorder

#### ZUBSOLV® short facts

API Buprenorphine/naloxone

**Technology** Sublingual

**Indication** Opioid use disorder

Market approvals US, EU and Australia

US launch 2013

**Commercial rights** Orexo owns global rights, ex EU

Partner EU Accord Healthcare

Patent protection US, EU, Australia and

New Zealand until 2032

#### **Product advantages include:**

- Higher bioavailability
- Fast dissolve time
- Preferred menthol flavor
- Broadest range of dose strengths





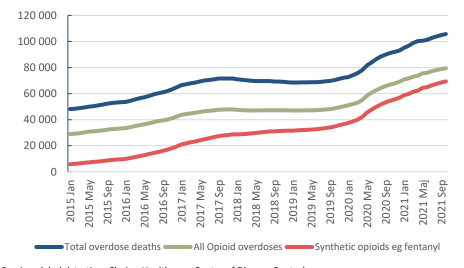
#### The opioid crisis - one of the largest health crises ever in the US

Americans misusing opioids

In need of treatment million

Under treatment million

Overdose deaths have surpassed 107 000 in 2021, fueled by use of synthetic opioids such as fentanyl





"Giana made the switch to heroin, and it was all downhill from there."



Elise discovered her daughter's opioid addiction months before she died from an overdose.

Read more at the Orexo blog

## ZUBSOLV® is an important cash generator

#### Key operational highlights in Q1 2022

- ✓ ZUBSOLV® added to preferred NY Medicaid formulary from March 22
- ✓ Commercial market share in NY in Q1 was 10 percent¹ compared to Medicaid with ~0.5 percent
- ✓ Increased investment in field force from April in NY
- ✓ Field force continue with MODIA™ awareness campaign and initiated modiaONE trial campaign to ZUBSOLV® customers in February

#### **Accumulated net revenues**

Since US launch 2013 - Q1 2022

4.2

US Pharma net revenues<sup>2</sup>

535 ×

#### ZUBSOLV® EBIT (SEK M) and EBIT margin, LTM Q1 2022



 $<sup>^{</sup>m 1}$  Includes UHG and Humana, 7.2% marketshare if these are excluded

<sup>&</sup>lt;sup>2</sup> LTM, Last Twelve Months (Q2 2021-Q1 2022)

## Multiple drivers for growth

1

Covid-19 effects likely to diminish improving patient access to care and Q2-Q4 historically with improving growth rates

2

Multiple comprehensive activities on-going on federal and state levels to enable more patients access to treatment

3

Improved market access for ZUBSOLV® with Public access increasing to 48%(42%) and a slight decrease in Commercial at 98% (99%)

4

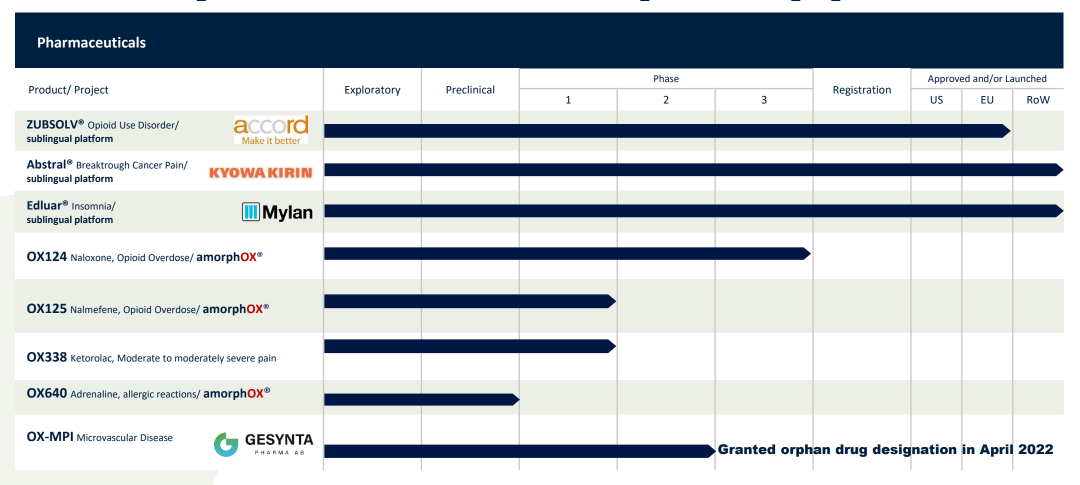
The launch of MODIA™ will open up new market segments and is highly complementary to ZUBSOLV®







#### Pharma products and development pipeline



### OX124 & OX125 – overdose rescue medications

- Significant health issue in the US with >107.000 deaths from overdose in 2021
- Current US market exceeding USD 400 million
- Based on amorphOX® and designed to treat overdoses caused by synthetic opioids, such as fentanyl
- OX124 clinically differentiated to market leader and GX of market leader
- OX124 on track to be filed with FDA in H2 2022



## OX640 – emergency treatment of allergic reactions

- First line treatment today: intramuscular auto-injectors
- OX640 offers:
  - Needle free alternative based on amorphOX®
  - Improved handling and storage (doubling shelf life of existing products)
  - No bisulfite antioxidant and preservative free
  - Improved convenience and acceptability vs injection products increases likelihood of timely use
- OX640 with excellent stability data compared to other products<sup>1</sup>
- First clinical trial on track to be initiated in Q3 2022



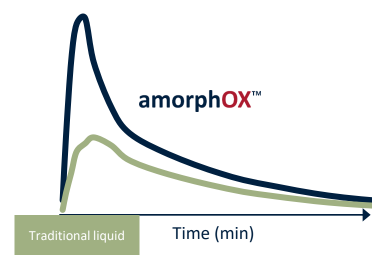


## Orexo internal platform building on the amorphOX® technology

#### Validated in humans

✓ Superior pharmacokinetic properties with more rapid onset, higher peak and overall exposure, lower variability

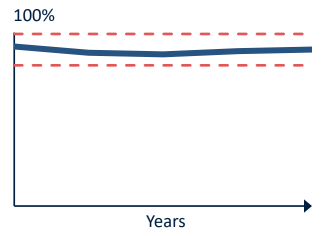
Plasma concentration from clinical trial



#### **Excellent stability**

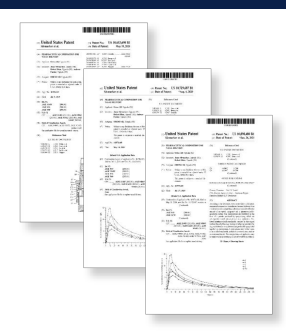
 Excellent stability even under accelerated conditions and proven to work on a broad scope of API's

Amount of API



#### Patent protected

✓ Three granted US patents and several patent applications have been filed with potential protection until 2042



## amorphOX® works for a broad scope of drugs

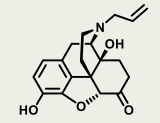
#### **Adrenaline**

0.5% after 6 months

#### **Apomorphine**

0.2% after 9 months

#### **Naloxone**



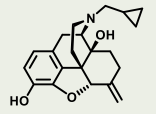
≤0.1% after 12 months

#### Loxapine

#### Ketorolac

0.8% after 6 months

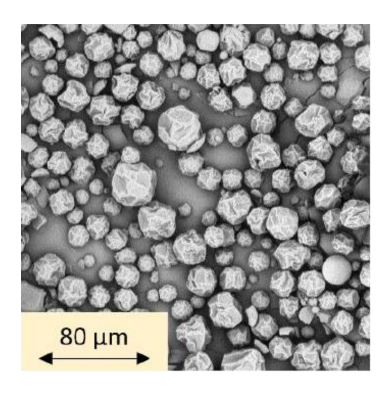
#### **Nalmefene**



≤0.1% after 15 months

Chemical degradation after accelerated stability studies at 40°C/75% RH

### amorphOX® and macromolecules



#### **Recent protein example**

- ✓ Enzyme of 464 kDa (~1000 amino acids)
- ✓ Successfully formulated in the **amorphOX**® platform
- ✓ High yield
- ✓ Free-flowing powder
- ✓ Narrow particle size distribution
- ✓ Enzymatic activity fully retained after:
  - manufacturing
  - 1M storage @ 40°C/75%RH

#### **DTx** in brief

- ✓ Subsection of digital health
- ✓ Evidence-based therapeutic intervention
- ✓ Prevent, manage, or treat a medical disorder or disease
- ✓ Particularly applicable in the mental illness & addiction space
- ✓ Standalone or along with pharma treatment
- ✓ Available 24/7



## Strong underlying trends will force the DTx market to gain traction

#### Healthcare systems challenged

Aging population and sky-rocketing costs are forcing the healthcare providers to rethink how to deliver healthcare to increase efficiency and value.

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Widespread technology acceptance

Covid-19 has further pushed forward the ongoing tech revolution and the use of telemedicine is pervasive.

Consumers (patients) in the center

Patients want to be seen as consumers and requires holistic and customized treatments with access 24/7.



Value-based care

Providers will be rewarded based on the ability to add patient value. Analyzing RWE data pave the way for efficient allocation of resources.

# Orexo is at the leading edge of digitalization in the pursuit to take DTx from its infancy to become a natural part of healthcare

"In less than a decade, DTx companies have completely disrupted the healthcare scene for the better."

The Future of Digital Therapeutics and The Impact On Care, The Linus Group, May 2021

- ✓ Establishing Reimbursement
  Orexo working in tight collaboration
  with world leaders in digital health to
  make DTx accessible to all patients.
  However, universal reimbursement
  processes still to be established
- ✓ **Disruptive technology**Through pilot programs, trials and real world evidence collection, Orexo is working with payers and leading healthcare organizations to build confidence in the value of our DTx to healthcare.

## Clinically proven DTx in collaboration with GAIA AG

Rooted in cognitive behavioral therapy techniques and based on AI technology offering a highly individualized intervention.

	modia	VORV!DA	deprexis
Instructions for use	Opioid dependence	Alcohol misuse	Depression
Clinical evidence	Ongoing randomized clinical trial, 400 patients	Evaluated in 1 randomized clinical trial, > 600 patients	Evaluated in 13 randomized clinical trials, > 2.800 patients
Length of treatment	6 months	6 months	3 months
Treatment method	Along with current standard of care including medication	Standalone or as a complement to current standard of care	Standalone or as a complement to current standard of care
FDA clearance	Will apply for a 510 k clearance, meanwhile launched under FDA's Public Health Emergency Use Authorization (EUA)	FDA cleared under the EUA	FDA cleared under the enforcement discretion



## Q1 2022 LTM (USD) - A transformative period building for future growth

- Significant investments in establishing digital therapeutics business and development of OX124
- Recurring business is well financed from ZUBSOLV® profit contribution

**Group net revenues** 

63

USD N

**US Pharma net revenues** 

**57** 

USD N

**EBITDA** 

14

**US Pharma EBIT** 

31

USD M

**Cash position** 

46 s

US Pharma EBIT margin

**55** 

6

## Legal update – no changes in Q1 2022

#### ZUBSOLV® patent dispute vs Sun Pharma

#### No changes in Q1

- √ 9 patents listed in the Orange Book
- ✓ Expiring dates Dec 2027– Sep 2032
- ✓ Previously successfully managed to defend ZUBSOLV® IP rights in the US appeal court

### Subpoena with regards to ZUBSOLV®

#### No changes in Q1

- √ Very limited activities in Q1
- ✓ No additional information received since issuance of subpoena July 2020





## Strong value drivers for long-term growth

1

Product portfolio addressing large and growing markets



2

Leveraging our US commercial excellence



3

Expanding pipeline based on the novel proprietary technology platform amorphOX®

4

Strong cash flow generation from the US Pharma

5

Entering digital therapeutics, a new evidence-based frontier in patient care

#### orexo

