



SOFTOX SOLUTIONS AS

# Interim Report

## H1 2026

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TICKER: SOFTX • Euronext Growth Oslo

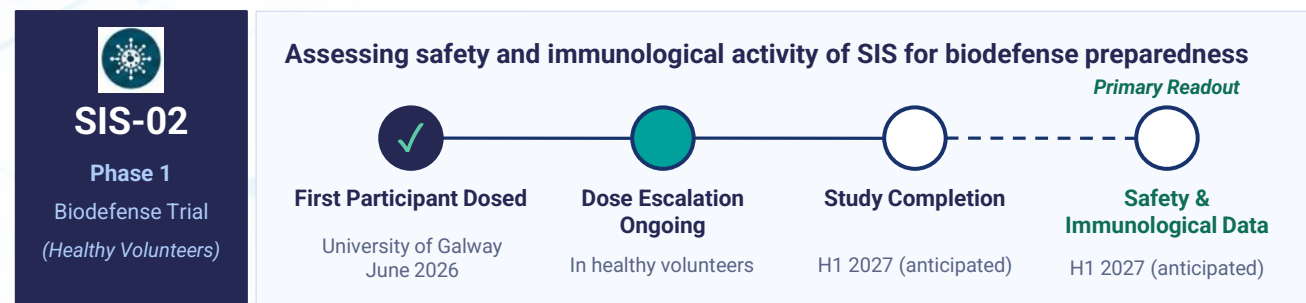
Clinical-stage pharmaceutical company developing  
a novel anti-infective therapy for airway infections

The SoftOx Solutions Group (SoftOx) comprises the holding company SoftOx Solutions AS, company org. no. 998 516 390, and the subsidiaries Water Innovation AB and SoftOx Defense Solutions AS. SoftOx Solutions Group is based in Oslo, Norway, with a subsidiary in Malmö, Sweden, and Clinical Operations in Copenhagen, Denmark.

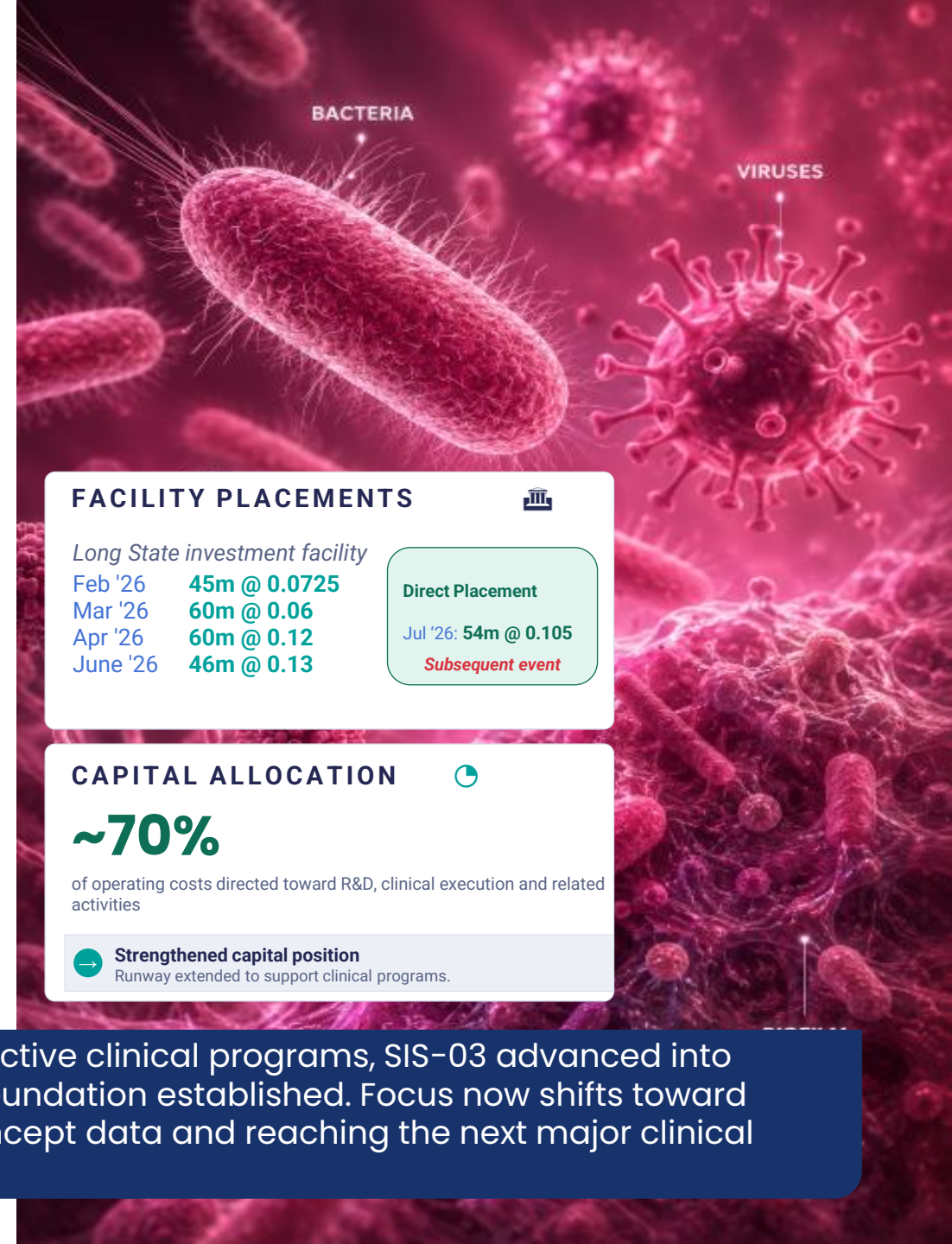


# Two clinical programs advancing toward key value inflection points

Strong clinical execution in H1 2026, positive safety foundation established, first patient dosed in SIS-03, disciplined capital allocation, and preparation for upcoming value inflection milestones.



✓ Value Inflection Point



## FACILITY PLACEMENTS

Long State investment facility

Feb '26 45m @ 0.0725  
Mar '26 60m @ 0.06  
Apr '26 60m @ 0.12  
June '26 46m @ 0.13

Direct Placement

Jul '26: 54m @ 0.105

*Subsequent event*

## CAPITAL ALLOCATION

~70%

of operating costs directed toward R&D, clinical execution and related activities

→ Strengthened capital position  
Runway extended to support clinical programs.

SoftOx enters H2 2026 with two active clinical programs, SIS-03 advanced into patients, and a positive safety foundation established. Focus now shifts toward generating patient proof-of-concept data and reaching the next major clinical value inflection point.

# Chairman's Reflections



**Ulrik Spork**

**Chairman of the Board**

SoftOx Solutions AS

The first half of 2026 marked an important transition for SoftOx as the Company moved from clinical preparation into active execution across both of its clinical programs.

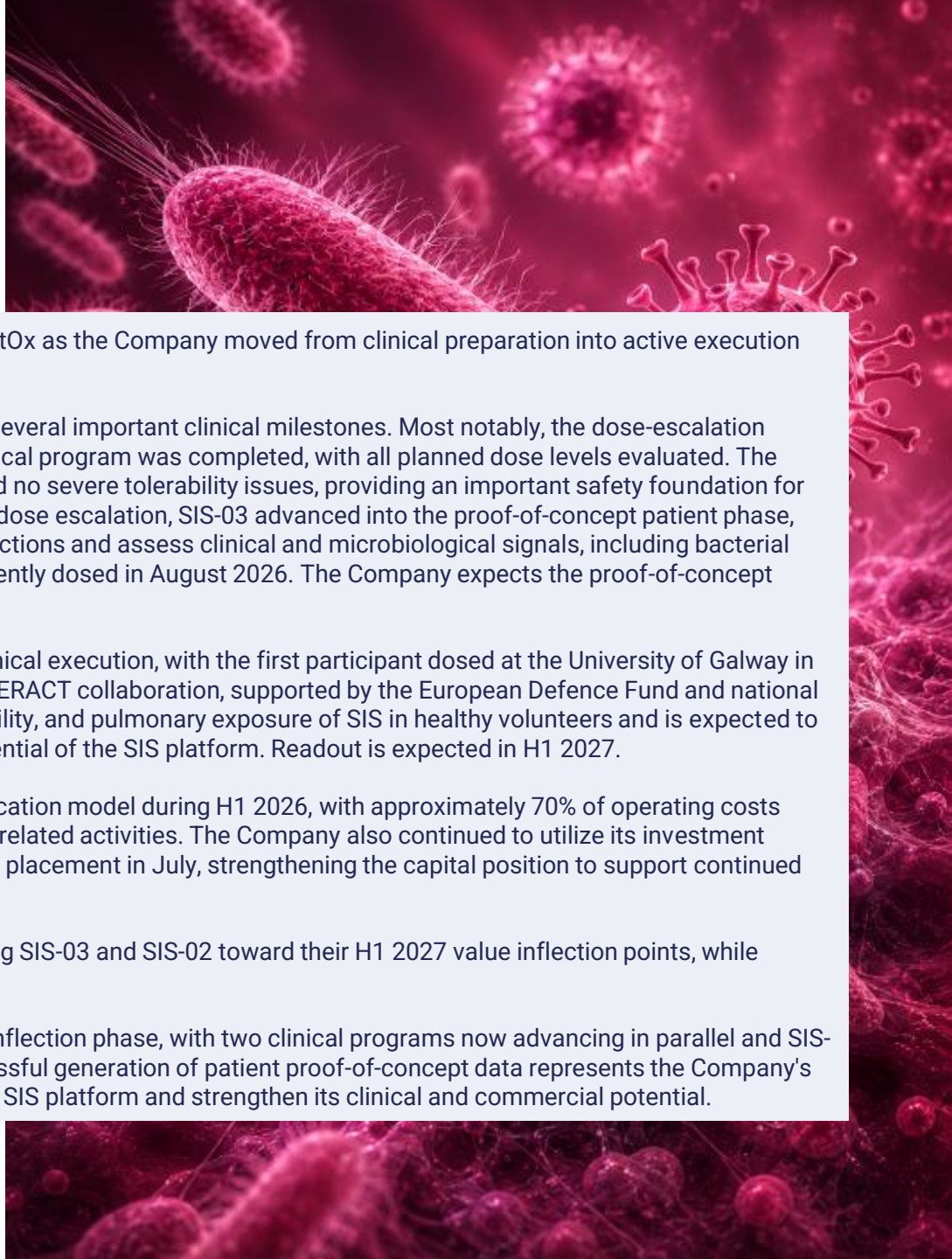
During the period and subsequent events, SoftOx achieved several important clinical milestones. Most notably, the dose-escalation phase of the **SIS-03 Phase 2a Chronic Airway Infection** clinical program was completed, with all planned dose levels evaluated. The topline safety review reported no serious adverse events and no severe tolerability issues, providing an important safety foundation for the continued development of SIS. Following completion of dose escalation, SIS-03 advanced into the proof-of-concept patient phase, designed to evaluate SIS in patients with chronic airway infections and assess clinical and microbiological signals, including bacterial load reduction in the airways. The first patient was subsequently dosed in August 2026. The Company expects the proof-of-concept phase to conclude in H1 2027.

In parallel, the **SIS-02 Phase 1 biodefense study** entered clinical execution, with the first participant dosed at the University of Galway in June 2026. SIS-02 is being conducted as part of the COUNTERACT collaboration, supported by the European Defence Fund and national defence partners. The study is evaluating the safety, tolerability, and pulmonary exposure of SIS in healthy volunteers and is expected to provide further clinical evidence supporting the broader potential of the SIS platform. Readout is expected in H1 2027.

SoftOx maintained a disciplined operational and capital-allocation model during H1 2026, with approximately 70% of operating costs directed toward R&D, manufacturing, clinical execution, and related activities. The Company also continued to utilize its investment facility during the period and completed a subsequent direct placement in July, strengthening the capital position to support continued clinical execution and upcoming value-driving milestones.

Looking ahead, the Company's primary focus is on advancing SIS-03 and SIS-02 toward their H1 2027 value inflection points, while preparing for future financing and potential partnerships..

The Board believes SoftOx has entered an important value inflection phase, with two clinical programs now advancing in parallel and SIS-03 progressing from healthy volunteers into patients. Successful generation of patient proof-of-concept data represents the Company's most important near-term opportunity to further validate the SIS platform and strengthen its clinical and commercial potential.



## SOFT-OX INHALED PLATFORM

# A non-antibiotic inhaled anti-infective therapy

*Solving high unmet needs in Airway & Lung Infections*



### THE PROBLEM



Respiratory infections are a massive global burden with limited treatment options.

- Rising antimicrobial drug resistance
- No broad-spectrum, non-antibiotic therapies delivered locally to the lungs
- High unmet need in cystic fibrosis, non-cystic fibrosis bronchiectasis, etc.

### OUR SOLUTION: – THE SIS THERAPY



SIS is a novel, non-antibiotic, broad anti-infective therapy delivered by inhalation.

- ✓ Broad activity against bacteria, viruses, fungi, and biofilms
- ✓ Local delivery – high lung concentration, low systemic exposure
- ✓ Potential across multiple indications with significant market opportunities

### MARKET OPPORTUNITY



#### CHRONIC AIRWAY INFECTIONS

~USD 5-8 BILLION+

Large and growing market with persistent unmet need



#### ACUTE RESPIRATORY INFECTIONS

~USD 10-15 BILLION+

High unmet need across a broad range of severe respiratory infections



#### BROAD ANTI-INFECTION OPPORTUNITIES

~USD 5-10 BILLION+

Addressing a wide spectrum of respiratory pathogens and emerging threats

### KEY INVESTMENT HIGHLIGHTS



#### Large market opportunity

Multi-billion USD markets across cystic fibrosis (CF), non-cystic fibrosis bronchiectasis (NCFB), ventilator-associated pneumonia (VAP), hospital-acquired pneumonia (HAP), and other respiratory and airway infections



#### Strong differentiation

Non-antibiotic mechanism of action (MoA) with broad anti-infective activity and local delivery



#### Clear value inflection

Proof-of-Concept (CF) readout H1 2027



#### Strategic optionality

Partner or advance independently – controlled path to value

### DEVELOPMENT & VALUE CREATION TIMELINE



Topline Safety Data SIS-03 Chronic Airway Infections (H1 2026)  
✓



FSFD SIS-02 Biodefense (H1 2026)  
✓



First Patient-Dosed SIS-03 Chronic Airway Infections (August 2026)  
✓



PoC Readout SIS-03 Chronic Airway Infections (H1 2027)



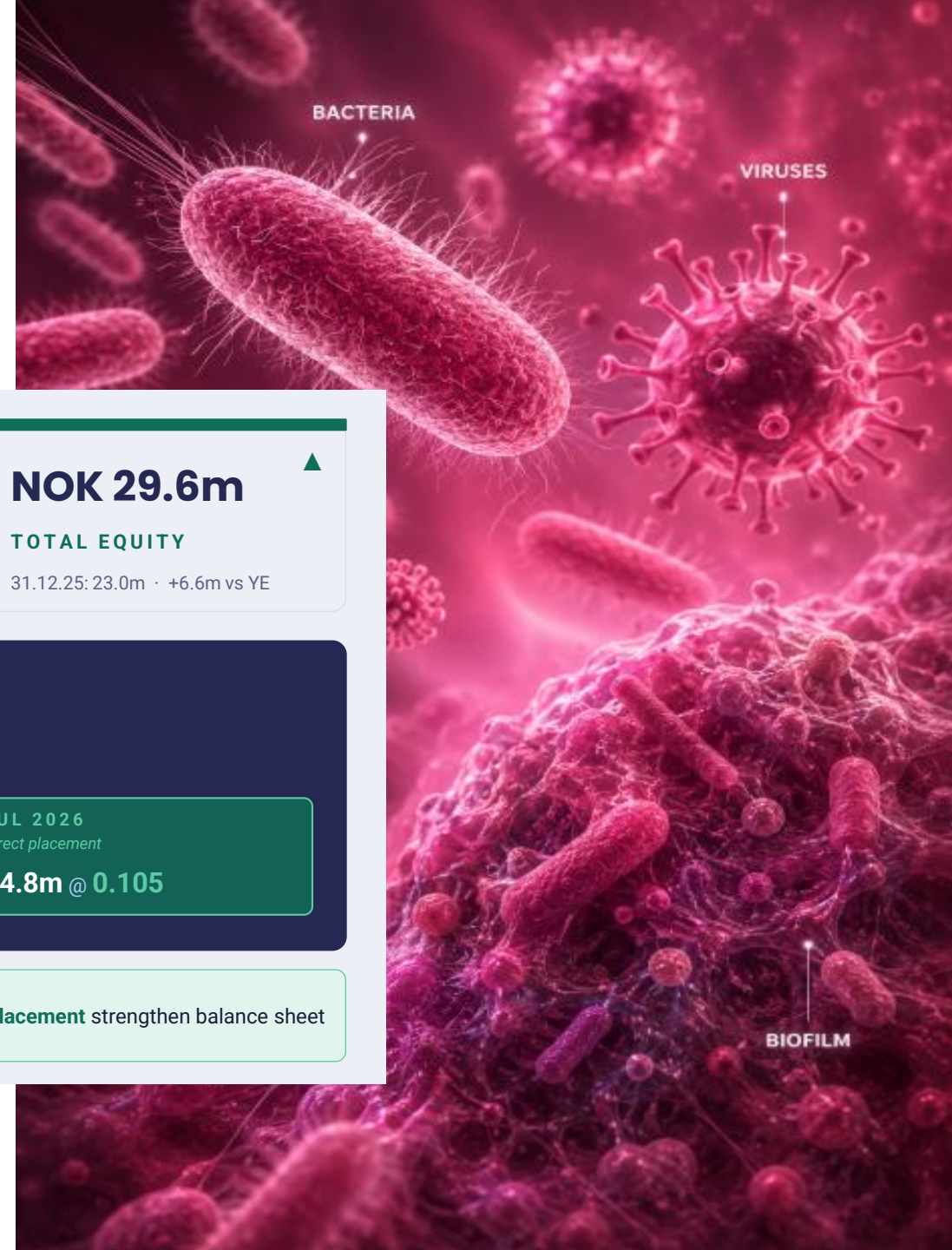
Readout SIS-02 Biodefense (H1 2027)



Value Realization Post PoC (Partnership or Phase II advancement)

# H1 2026 & subsequent financing highlights

Financial position and funding activity at a glance



**NOK 5.8m**

OPERATING REVENUE

H1 2025: 7.5m · EDF project timing

**NOK (13.6m)**

PRE-TAX LOSS

H1 2025: (1.4m) · SIS-02 & SIS-03 study ramp-up

**NOK 21.4m**

CASH POSITION

31.12.25: 19.8m · +1.6m in H1

**NOK 29.6m**

TOTAL EQUITY

31.12.25: 23.0m · +6.6m vs YE

## FUNDING ACTIVITY | H1 2026 + JULY 2026

Long State facility **up to NOK 50m** / 24m · extendable to **NOK 80m** / 36m

Equity line of credit + supplemental direct placements.

FEB 2026

Long State

**45m @ 0.0725**

MAR 2026

Long State

**60m @ 0.06**

APR 2026

Long State

**60m @ 0.12 ↑**

JUNE 2026

Long State

**46m @ 0.13 ↑**

JUL 2026

Direct placement

**54.8m @ 0.105**

## OPERATIONAL ACTIVITY

Higher operating expenses reflect increased clinical and development activity across SIS-02 and SIS-03.

## LIQUIDITY POSITION

H1 raises of **NOK 20.1m + July direct placement** strengthen balance sheet for continued clinical execution.

H1 2026 figures unaudited. Comparable figures: H1 2025 (P&L lines) or 31.12.2025 (balance sheet lines). Values rounded to NOK 0.1m for presentation.



# ABOUT SOFT-OX

## SIS: A THERAPY WITH SIGNIFICANT POTENTIAL

# Multi-pathogen therapy for respiratory infections with clear Proof-of-Concept path

SoftOx Solutions AS is developing SIS, a novel inhaled anti-infective therapy for respiratory infections in the airways and lungs. Delivered by nebulizer, SIS is being developed to address bacterial, viral, and fungal pathogens through local delivery directly at the site of infection, without systemic exposure to the remainder of the human body, with a mechanism of action (MoA) not expected to drive antimicrobial resistance (AMR). Its broad pathogen relevance positions SIS as a distinct therapeutic in respiratory infections.

A favorable safety profile has already been demonstrated in first-in-human studies, supporting progression into patient trials. The next key step is to establish a clinical proof-of-concept to support further clinical progression and partnering.

### THREE SIGNIFICANT UNMET MEDICAL NEEDS

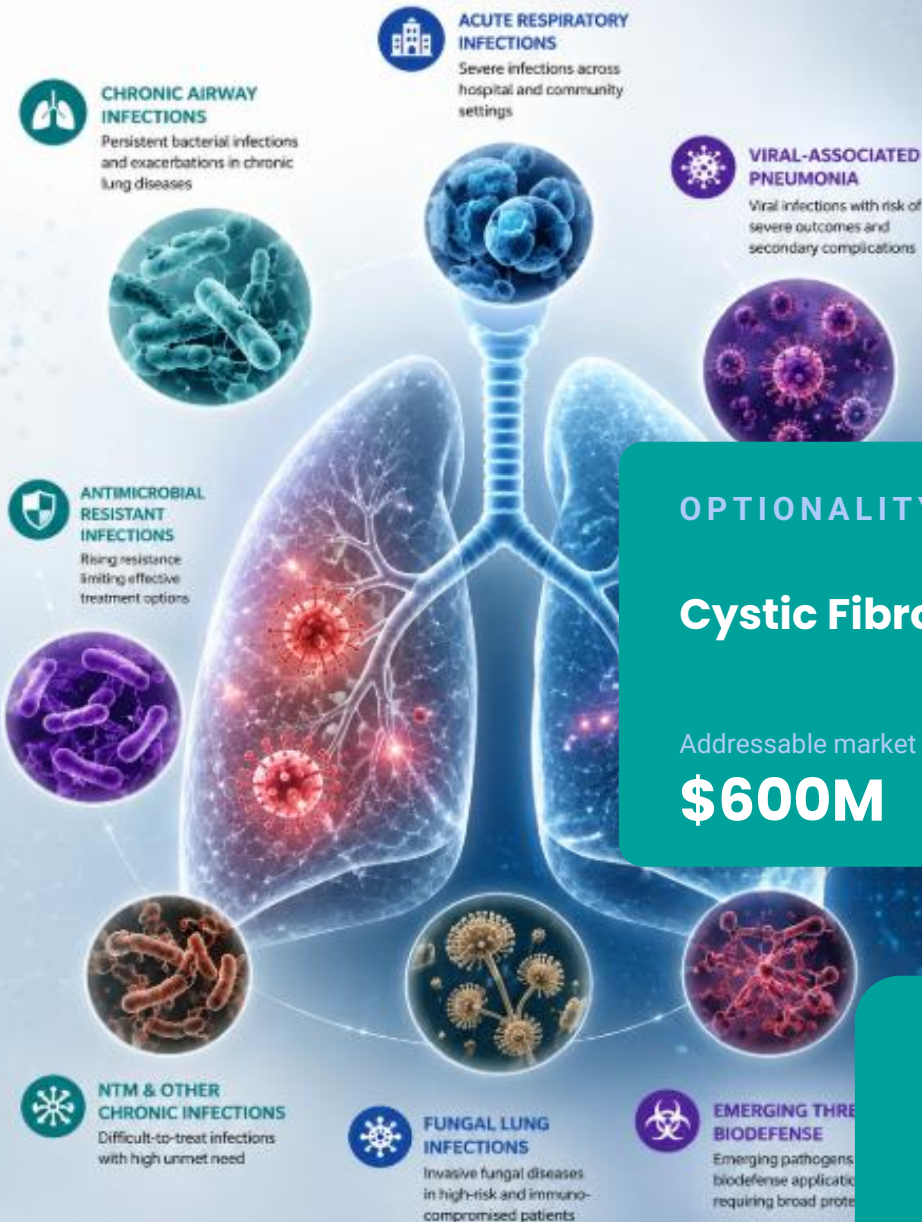
- 1 Chronic respiratory infections**  
CF, NCFB, and biofilm-driven infections with limited treatment options.
- 2 Multi-drug-resistant pathogens**  
As AMR emerges, the existing standard-of-care becomes less effective
- 3 Biological threat countermeasures**  
Biodefence: Preparedness potential against airborne biological threats

Strategy: clinical validation → Proof-of-Concept → partnering for late-stage development



# Respiratory infections represent a multi-billion-dollar market

*SoftOx is targeting well-defined respiratory segments with high unmet medical need, using chronic lung disease in Cystic Fibrosis as the initial proof-of-concept (PoC) setting to support broader future expansion across multiple respiratory indications.*



## OPTIONALITY A

### Cystic Fibrosis (CF)

Addressable market

**\$600M**

## OPTIONALITY B

### Non-CF Bronchiectasis (NCFB)

Addressable market

**\$5B**

## OPTIONALITY C .. X

### Biodefense, Pneumonia, VAP

Addressable market

**Multi-billion**

Successful Proof-of-Concept in Cystic Fibrosis (CF) unlocks expansion into significantly larger respiratory markets and multiple high-value indications

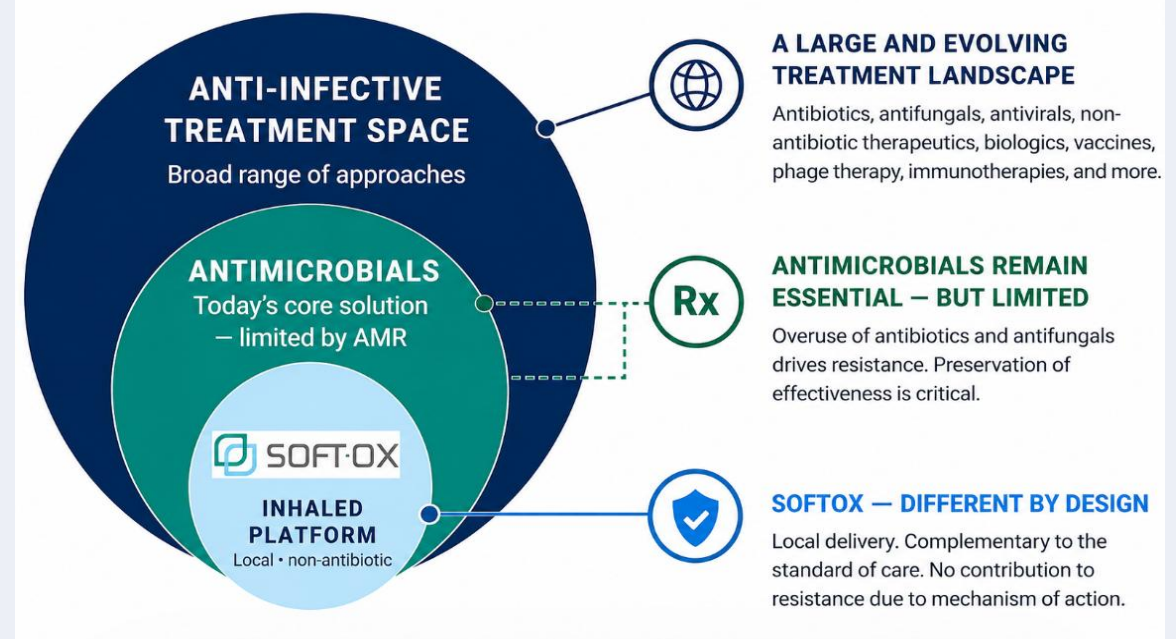
# Antimicrobial resistance (AMR) drives need for complementary anti-infective therapies

Antibiotics, antifungal, and antivirals have been the foundation of infection treatment for decades, but rising antimicrobial resistance is reducing effectiveness in important settings and increasing the need for complementary treatment approaches.

This is highly relevant in respiratory infections, where treatment limitations, recurrent infections, and resistant pathogens create demand for non-antibiotic options that can work alongside standard of care.



**1.27 million** deaths per year attributable to AMR globally (Lancet, 2022) – projected to rise sharply by 2050.



AMR increases the relevance of complementary, non-antibiotic approaches in respiratory infections.

## THE SOFTOX TECHNOLOGY

# A non-antibiotic inhaled anti-infective therapy



### HYPOCHLOROUS ACID

Documented broad antimicrobial effect



### ORGANIC ACID

Antimicrobial stabilizer & biofilm eradicator



#### SYNERGY

### SoftOx Inhalation Solution (SIS)

*Unique ability to eradicate infections at the site*



### SIS

A non-antibiotic inhaled anti-infective therapy

- **Broad pan-spectrum effect**

Virucidal and bactericidal activity across diverse pathogens

- **Effective against tolerant bacteria**

Targets biofilms where traditional antibiotics fail

- **No induction of resistance**

No evidence of contributing to AMR development

- **Favorable safety profile**

No systemic side effects in first-in-human study

- **Stabilized formulation**

Maintains efficacy throughout shelf life

- **Completed all pre-clinical Studies**

All necessary preclinical studies have been completed.

- **No severe adverse effects in Phase 1 trial**

27.9% adverse effects (SIS) vs 21.4% (placebo) – all mild

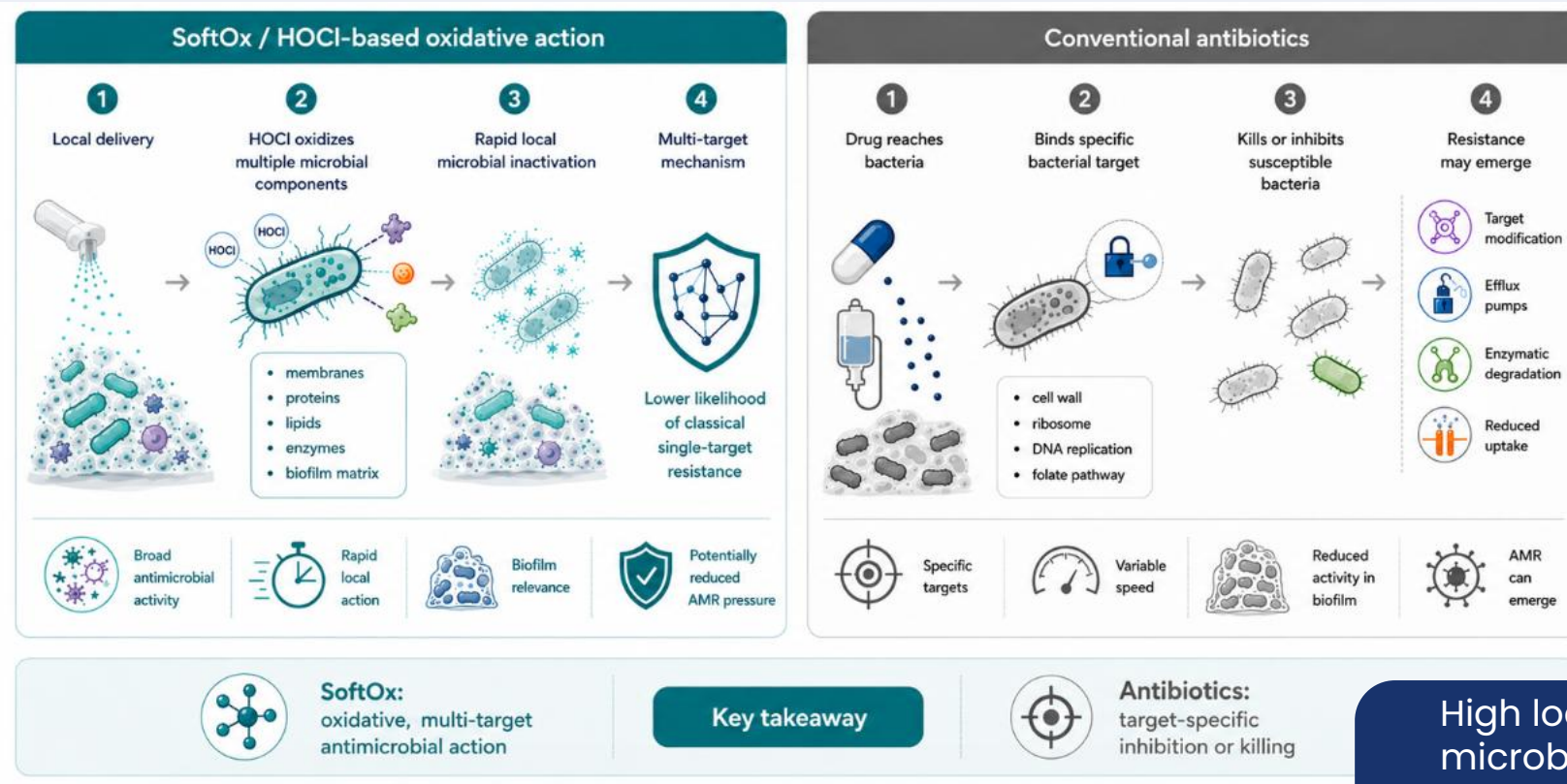
#### OUR POSITIONING

- **Anti-infective, not antibiotic**  
Novel mechanism and delivery.
- **Complement, not compete**  
Designed to work alongside existing therapies.
- **Focus where need is high**  
Respiratory infections – major driver of mortality.
- **Large market potential**  
Growing market, room for differentiated solutions.
- **Long-term value**  
Better outcomes and antibiotic stewardship.

A non-antibiotic therapy designed to deliver effective infection control without contributing to antimicrobial resistance.

## SoftOx vs Conventional Antibiotics

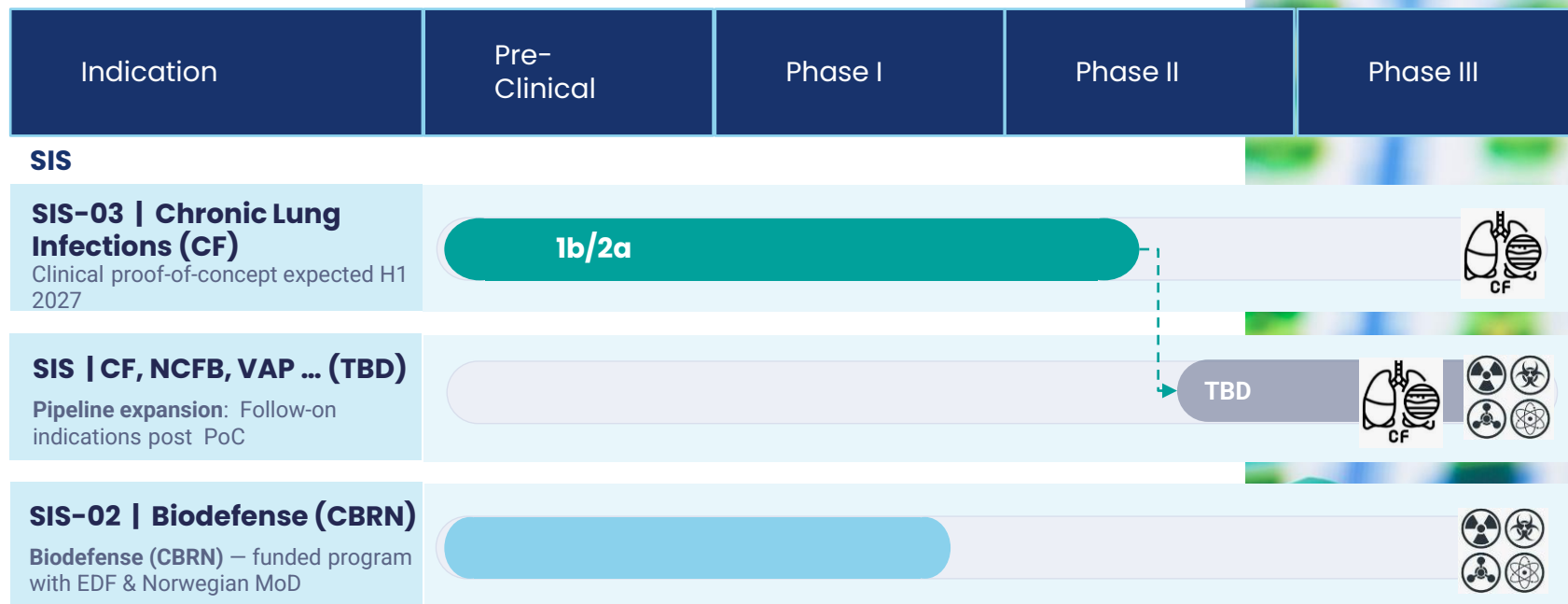
*A hypochlorous acid (HOCL) based oxidative antimicrobial approach differs from target-specific antibiotic therapy*



High local exposure and broad oxidative microbial inactivation may increase the potential for clinical benefit while reducing the selective pressure driving antimicrobial resistance.

## CLINICAL PROGRAMS WITH FUTURE EXPANSION POTENTIAL

# Clinical Development Roadmap

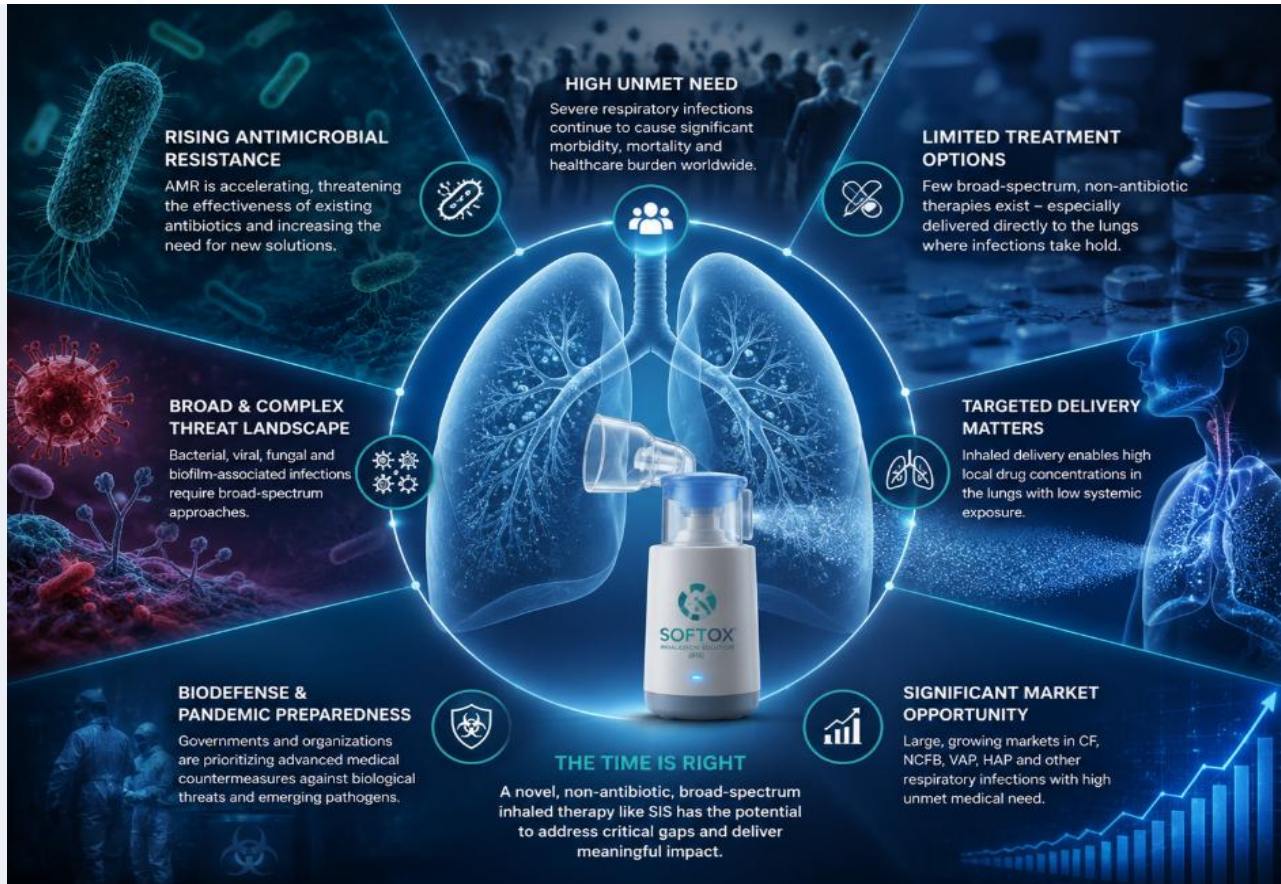


One inhaled antimicrobial platform advancing multiple respiratory indications toward clinical Proof-of-Concept and value-creating partnerships.

## SOFT-OX INHALED THERAPIES – WHY NOW?

# A Broad-Spectrum Inhaled Anti-Infective Opportunity

*Rising antimicrobial resistance, limited treatment options, and increasing focus on biodefense are driving demand for new localized anti-infective therapies.*



*SIS is being developed as a novel inhaled anti-infective therapy designed to reduce bacterial burden directly in the airways and lungs.*

High local exposure combined with broad oxidative microbial inactivation may increase the potential for clinical benefit while reducing the selective pressure driving antimicrobial resistance.



## AIRWAY & LUNG INFECTION APPLICATIONS (Lead Clinical Program SIS-03)

# Lead clinical program: SIS-03 phase 2a in Chronic Lung Infections

SIS is being developed as a novel inhaled anti-infective therapy designed to reduce bacterial burden directly in the lungs with limited risk of driving antimicrobial resistance.

## TRIAL DESIGN

### PART A • Dose Escalation

24-30 healthy volunteers | H1 2026  
*Safety & dosing foundation*



### PART B • Proof of Concept

15-25 patients | completes Q1 2027  
*Bacterial load reduction*

## REGULATORY MILESTONES

- CTA submitted Sep 2025 ✓
- Approved by Danish Medicines Agency (DKMA) Dec 2025 ✓
- First volunteer dosed Mar 2026 ✓
- Dose Escalation Readout June 2026 ✓
- Topline Safety Data June 2026 ✓
- First Patient Dosed August 2026 ✓

## STRONG STUDY DESIGN

- Well-defined patient population
- Rigshospitalet specializes in chronic airway infections
- Designed per EMA/FDA guidance

## MANUFACTURING

- GMP drug substance ✓
- GMP drug product ✓
- Clinical supply secured ✓

Clinical proof-of-concept readout expected H1 2027

Our mission is to transform the treatment of chronic airway infections by delivering a broad-spectrum inhaled anti-infective therapy designed to target infections locally in the lungs without systemic exposure.

# POSITIVE TOPLINE SAFETY DATA FROM **SIS-03** DOSE-ESCALATION PHASE

Dose escalation completed with **no serious adverse events**  
and no severe tolerability issues recorded.



## STUDY OVERVIEW

- SIS-03 dose-escalation phase conducted in healthy volunteers
- Multiple ascending dose levels evaluated
- Primary objectives: safety, tolerability and assessment of pulmonary exposure profile
- All planned dose levels successfully evaluated

## KEY SAFETY OUTCOMES



### NO SERIOUS ADVERSE EVENTS

No SAEs reported  
at any dose level.



### NO SEVERE TOLERABILITY ISSUES

No severe treatment-  
related adverse events.



### RESPIRATORY SAFETY CONFIRMED

No safety concerns  
related to pulmonary  
function or oxygenation.



### DOSE ESCALATION COMPLETED

All planned dose levels  
were well tolerated.

## TOPLINE SAFETY SUMMARY



### SAFETY FIRST

No serious adverse  
events observed.



### WELL TOLERATED

No severe tolerability  
issues recorded across  
all dose levels.



### RESPIRATORY SAFETY VERIFIED

No impact on lung  
function or  
oxygenation.



### ADVANCING WITH CONFIDENCE

SIS-03 is advancing  
to the proof-of-concept  
patient phase.



## NEXT STEP

SIS-03 proof-of-concept patient phase initiated in cystic fibrosis patients with chronic airway infections.  
Expected to dose between 15 and 25 patients. Trial expected to conclude in H1 2027.



# FIRST PATIENT DOSED IN SIS-03 PROOF-OF-CONCEPT PATIENT PHASE

First patient with chronic airway infection dosed, marking initiation of the proof-of-concept phase



## A KEY MILESTONE FOR SOFTOX



### FIRST PATIENT DOSED

First patient with chronic airway infection dosed in the SIS-03 proof-of-concept patient phase.



### PROOF-OF-CONCEPT PHASE INITIATED

Evaluating SIS in patients with chronic airway infections to generate clinical and microbiological data, including bacterial load reduction.



### BUILT ON A STRONG SAFETY FOUNDATION

Follows completion of the dose-escalation phase in healthy volunteers with positive topline safety data.

## SIS-03 PROGRAM OVERVIEW

- ✓ Novel inhaled anti-infective therapy (SIS) designed to address chronic airway infections in patients
- ✓ Targeting biofilm-associated bacteria, a major unmet medical need
- ✓ Potential to overcome limitations of existing therapies and antimicrobial resistance
- ✓ Developed using our proprietary HOCl + acetic acid platform



## SIS-03 TRIAL STATUS



### DOSE-ESCALATION PHASE COMPLETED

No serious adverse events and no severe tolerability issues observed



### POSITIVE TOPLINE SAFETY REVIEW

All planned dose levels well tolerated



### FIRST PATIENT DOSED

Proof-of-concept patient phase initiated in patients with chronic airway infections



### EXPECTED COMPLETION H1 2027

15-25 patients planned in this phase



**A MAJOR STEP FORWARD IN OUR MISSION TO TRANSFORM CARE FOR PEOPLE LIVING WITH CHRONIC AIRWAY INFECTIONS.**

# Established Copenhagen CF Center Supports SIS-03 Patient Recruitment

The Copenhagen Cystic Fibrosis Center at Rigshospitalet provides access to one of the largest and most experienced CF centers in Europe, with a long track record of research and trials in chronic lung infection.



Rigshospitalet, Copenhagen

## THE COPENHAGEN CYSTIC FIBROSIS CENTER – A LEADING EUROPEAN SITE



- ✓ One of Europe's largest and most experienced CF centers
- ✓ More than 300 CF patients cared for at the Copenhagen Cystic Fibrosis Center\*
- ✓ CF research and clinical trials for several decades
- ✓ Specialized focus on chronic lung infection and biofilm infection in CF
- ✓ Strong collaboration with the Danish Cystic Fibrosis Registry and patient organizations
- ✓ State-of-the-art infrastructure and expertise to conduct high-quality clinical studies

## STRATEGIC LOCATION & RESEARCH EXCELLENCE



### SPECIALIZED CF CENTER

More than 300 CF patients cared for at the Copenhagen Cystic Fibrosis Center\*



### FOCUS ON CHRONIC INFECTION

Decades of research experience in chronic lung infection and biofilm infection in CF



### NATIONAL COLLABORATION

Close collaboration with the Danish Cystic Fibrosis Registry and patient organizations

## THE COPENHAGEN CF CENTER PATIENT POOL



More than  
**300**  
CF patients  
cared for at the  
Copenhagen CF Center\*

A large and well-characterized CF population with extensive clinical data and long-term follow-up, providing a strong foundation for efficient and high-quality clinical research.

## EFFICIENT RECRUITMENT PATHWAY



**Patient Identification**  
Through clinic &  
Danish CF Registry

**Pre-screening**  
Medical record  
review

**Screening Visit**  
Eligibility confirmation  
& informed consent

**Enrollment**  
Targeted and  
efficient execution

## SITE EXPERIENCE & TRACK RECORD



**Decades**  
of CF research  
and clinical  
trials



**Proven**  
experience in  
conducting interventional  
CF studies\*\*



**Modern**  
infrastructure and  
expert team for  
trial execution



Access to an established more-than-300-patient CF center with decades of chronic lung infection and biofilm research experience provides a strong clinical setting for execution of the SIS-03 Proof-of-Concept study.



## SIS-03: Advancing into Proof-of-Concept in Chronic Airway Infections

*Dose escalation completed with positive topline safety data, advancing SIS-03 into patient Proof-of-Concept – the program's next major value inflection phase*

*SIS is being developed as a novel inhaled anti-infective therapy designed to reduce bacterial burden directly in the lungs.*

### FOUNDATION ESTABLISHED



#### Clinical & Regulatory Foundation

- Chronic airway infection PoC strategy established
- SIS-03 approved by Danish Medicines Agency



#### CMC & Manufacturing established

- New drug substance developed and GMP manufactured
- GMP drug product and clinical supply secured

### H1 2026 & SUBSEQUENT EVENTS



#### Clinical Execution Delivered

- Phase 2a initiated – first healthy volunteer dosed March 2026
- Dose escalation completed – all planned dose levels evaluated
- Positive topline safety data – no serious adverse events or severe tolerability issues
- Additional GMP drug product manufactured



#### Subsequent Event – August 2026

- First patient with chronic airway infection dosed
- Proof-of-Concept patient phase initiated

### NEXT VALUE INFLECTION



#### PATIENT PROOF-OF-CONCEPT



#### Now underway

- Evaluate clinical and microbiological signals
- Assess bacterial load reduction in patients' airways
- 15–25 patients planned



#### Next major milestone

- PoC phase expected to conclude H1 2027
- Data to inform further clinical development and potential partnering

SIS-03 has progressed from clinical preparation through dose escalation and positive topline safety to Proof-of-Concept evaluation in patients with chronic airway infections.



## MEDICAL BIODEFENCE APPLICATION (SIS-02)

# Biodefense program: SIS-02: Advancing into Phase 1

SoftOx is advancing its inhaled antimicrobial therapy as a medical countermeasure against biological threats, supported by European and Norwegian defense programs.

## COUNTERACT PROGRAM PARTNERS

### European Defence Fund

Program funding from EDF

### Norwegian MoD

National co-funding

### FFI (Norway)

Defence Research Establishment

### European Partners

EU COUNTERACT Consortium



**Demonstrated broad-spectrum activity**

### Preclinical results

- › Gram-negative bacteria (e.g. *P. aeruginosa*)
- › Gram-positive bacteria (e.g. *S. pneumoniae*)
- › Resistance pathogens (e.g. *Klebsiella pneumoniae*)
- › Viral pathogens (e.g. *influenza*)

When it matters most.  
**BREATHE. NEUTRALIZE. PROTECT.**



Rapid  
Action



Broad  
Spectrum



Protects People  
& Mission

Our mission is to develop a broad-spectrum anti-infective inhalation therapy that can rapidly neutralize biological threats, protecting warfighters and civilians when it matters most

# SIS-02: Phase 1 Clinical Study Underway

Broad-spectrum inhaled anti-infective program supporting respiratory infection and biological preparedness applications, with Phase 1 clinical evaluation now underway.

*SoftOx Inhalation Solution (SIS)*  
– designed to rapidly neutralize  
a wide range of biological  
threats at the source.

## ACHIEVED BEFORE 2026



### COUNTERACT program established

- European Defence Fund-supported collaboration established
- SoftOx leads development of the inhaled anti-infective component



### Preclinical foundation established

- In vitro and in vivo airway infection studies completed
- Activity demonstrated against relevant pathogens



### CMC & manufacturing established

- GMP drug substance successfully manufactured
- GMP drug product manufactured and released

## KEY ACHIEVEMENTS H1 2026



### Regulatory approval achieved

- SIS-02 Clinical Trial Application approved by HPRA in Ireland
- Regulatory clearance enabled initiation of the Phase 1 study



### Phase 1 clinical study initiated

- First participant dosed in June 2026
- Clinical evaluation of safety, tolerability and pulmonary exposure underway



### Clinical execution established

- Study activated at University of Galway
- Clinical operations, logistics and study supply in place



### GMP manufacturing delivered

- Additional GMP drug product manufactured to support clinical execution and continued development

## OUTLOOK



### Continue Phase 1 execution

- Complete planned healthy-volunteer clinical evaluation
- Assess safety, tolerability, and pulmonary exposure



### Generate additional clinical evidence

- Strengthen the human clinical evidence base for SIS
- Support continued development across respiratory infection and biological preparedness applications



### Next clinical milestone

- SIS-02 readout expected H1 2027
- Results to inform subsequent development activities.

SIS-02 is now in clinical execution — with the first participant dosed and Phase 1 readout expected in H1 2027

## Confirmation from the Board

### Outlook & Objectives

SoftOx has entered an important value inflection phase with two clinical programs now in active execution and key near-term milestones ahead.

**SIS-03 Phase 2a in chronic airway infections** has advanced from dose escalation to the patient Proof-of-Concept phase. The first patient was dosed in August 2026, and the program is designed to assess clinical and microbiological signals, including reduction of bacterial load in the airways. Topline PoC results are expected in H1 2027.

**SIS-02 Phase 1 biodefense study** is underway in healthy volunteers following HPRA approval in Ireland. The first participant was dosed in June 2026, and the study is evaluating safety, tolerability, and pulmonary exposure, with readout expected in H1 2027.

The upcoming clinical milestones represent important value inflection points and form the basis for potential partnering discussions and further strategic opportunities.

The Company will continue to focus on disciplined capital allocation, efficient execution of both clinical programs and maintaining a lean operating model through to key value-creating milestones.

### Confirmation from the Board of Directors

The Board confirms that, to the best of its knowledge, the financial statements for the period 1 January to 30 June 2026 have been prepared in accordance with applicable accounting standards and give a true and fair view of the Group's and the Company's assets, liabilities, financial position, and results of operations.

The Board further confirms that the report provides a true and fair overview of the development and performance of the business, as well as a description of key risks and uncertainties.

### Going Concern

The Board considers that the Company has sufficient flexibility through its existing financing arrangements, investment facility, and cash position to support planned operations in the near term.

Continued access to capital remains important to execute the Company's strategy and to advance both clinical programs through their respective milestones. The Board will actively pursue additional financing as required.

Lysaker, 22<sup>nd</sup> of September 2026

The Board of Directors of SoftOx Solutions AS

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**Ulrik Spork**

Chairman of the Board

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**Christian Vinding Thompson**

Vice Chair

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**Tore Duvold**

Board Member

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**Adrian Bignami**

Board Member



# FINANCIAL STATEMENTS

## Key Financial Figures

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

| KEY FINANCIAL FIGURES                   | SECOND QUARTER |               | FIRST HALF YEAR |               | FULL YEAR     |               |
|-----------------------------------------|----------------|---------------|-----------------|---------------|---------------|---------------|
|                                         | 2026           | 2025          | 2026            | 2025          | 2026          | 2025          |
| <b>OPERATING PERFORMANCE</b>            |                |               |                 |               |               |               |
| Total operating revenue                 | 1,906          | 4,291         | 5,807           | 7,507         | 5,807         | 15,584        |
| Total operating expenses                | 10,431         | 7,573         | 19,462          | 9,503         | 19,462        | 26,955        |
| Operating result                        | (8,526)        | (3,282)       | (13,655)        | (1,996)       | (13,655)      | (11,371)      |
| Profit before tax                       | (8,498)        | (3,343)       | (13,601)        | (1,390)       | (13,601)      | (10,813)      |
| <b>LIQUIDITY</b>                        |                |               |                 |               |               |               |
| Net proceeds from equity issues         | 13,229         | 2,925         | 20,092          | 9,055         | 20,092        | 19,089        |
| Net change in cash and equivalents      | 1,766          | (3,043)       | 1,578           | 9,260         | 1,578         | 9,281         |
| Cash and equivalents at end of period   | 21,372         | 19,773        | 21,372          | 19,773        | 21,372        | 19,794        |
| <b>EFFICIENCY</b>                       |                |               |                 |               |               |               |
| R&D costs as % of operating costs       | 66 %           | —             | 70 %            | —             | 70 %          | —             |
| <b>SHARE COUNT (in shares)</b>          |                |               |                 |               |               |               |
| Outstanding shares, beginning of period | 2,495,416,994  | 1,951,253,942 | 2,390,416,994   | 1,951,253,942 | 2,390,416,994 | 1,951,253,942 |
| Outstanding shares, end of period       | 2,601,797,517  | 2,240,416,994 | 2,601,797,517   | 2,240,416,994 | 2,601,797,517 | 2,390,416,994 |

H1 2026 figures are unaudited. Year columns show comparative full-year 2025 figures from the audited annual report.  
Outstanding shares shown in raw count.

## Profit & Loss Statement

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

| PROFIT & LOSS                           | SECOND QUARTER |                | FIRST HALF YEAR |                | FULL YEAR       |                 |
|-----------------------------------------|----------------|----------------|-----------------|----------------|-----------------|-----------------|
|                                         | 2026           | 2025           | 2026            | 2025           | 2026            | 2025            |
| <b>OPERATING REVENUE</b>                |                |                |                 |                |                 |                 |
| Other operating revenues                | 1,906          | 4,291          | 5,807           | 7,507          | 5,807           | 15,584          |
| <b>Total operating revenues</b>         | <b>1,906</b>   | <b>4,291</b>   | <b>5,807</b>    | <b>7,507</b>   | <b>5,807</b>    | <b>15,584</b>   |
| <b>OPERATING EXPENSES</b>               |                |                |                 |                |                 |                 |
| Personnel expenses                      | 908            | 1,838          | 2,503           | 3,241          | 2,503           | 6,087           |
| Other operating expenses                | 8,609          | 4,955          | 15,160          | 4,704          | 15,160          | 17,677          |
| Depreciation                            | 914            | 779            | 1,799           | 1,558          | 1,799           | 3,192           |
| <b>Total operating expenses</b>         | <b>10,431</b>  | <b>7,573</b>   | <b>19,462</b>   | <b>9,503</b>   | <b>19,462</b>   | <b>26,955</b>   |
| <b>Operating result</b>                 | <b>(8,526)</b> | <b>(3,282)</b> | <b>(13,655)</b> | <b>(1,996)</b> | <b>(13,655)</b> | <b>(11,371)</b> |
| <b>FINANCIAL ITEMS</b>                  |                |                |                 |                |                 |                 |
| Net financial items                     | 28             | (62)           | 54              | 606            | 54              | 558             |
| <b>Profit before tax</b>                | <b>(8,498)</b> | <b>(3,343)</b> | <b>(13,601)</b> | <b>(1,390)</b> | <b>(13,601)</b> | <b>(10,813)</b> |
| Tax                                     | —              | —              | —               | —              | —               | —               |
| <b>Net profit (loss) for the period</b> | <b>(8,498)</b> | <b>(3,343)</b> | <b>(13,601)</b> | <b>(1,390)</b> | <b>(13,601)</b> | <b>(10,813)</b> |

H1 2026 figures are unaudited. Year columns: 2026 = year-to-date through Q2; 2025 = full audited year. H1 2026 = Q1 + Q2 cumulative. Figures may differ from earlier interim reports as a result of subsequent adjustments.

## Balance Sheet

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

| BALANCE SHEET                       |               | AS AT          |               |
|-------------------------------------|---------------|----------------|---------------|
|                                     | 30.06.2026    | 30.06.2025     | 31.12.2025    |
| <b>ASSETS</b>                       |               |                |               |
| Other intangible assets             | 13,300        | 14,039         | 14,137        |
| Deferred tax asset                  | —             | 84,203         | —             |
| <b>Total non-current assets</b>     | <b>13,300</b> | <b>98,242</b>  | <b>14,137</b> |
| Cash and cash equivalents           | 21,372        | 19,773         | 19,794        |
| <b>Total current assets</b>         | <b>21,372</b> | <b>19,773</b>  | <b>19,794</b> |
| <b>Total assets</b>                 | <b>34,672</b> | <b>118,015</b> | <b>33,931</b> |
| <b>EQUITY &amp; LIABILITIES</b>     |               |                |               |
| Share capital                       | 52,036        | 44,808         | 47,808        |
| Share premium reserve               | (8,648)       | 63,238         | (24,494)      |
| Other equity                        | (13,791)      | (3,745)        | (361)         |
| <b>Total equity</b>                 | <b>29,598</b> | <b>104,301</b> | <b>22,954</b> |
| Public duties payable               | (394)         | (160)          | (243)         |
| Other current liabilities           | 2,484         | 10,386         | 10,223        |
| Accounts payable                    | 2,984         | 3,488          | 997           |
| <b>Total current liabilities</b>    | <b>5,074</b>  | <b>13,714</b>  | <b>10,977</b> |
| <b>Total equity and liabilities</b> | <b>34,672</b> | <b>118,015</b> | <b>33,931</b> |

## Cash Flow Statement

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

| CASH FLOW STATEMENT                            | SECOND QUARTER  |                | FIRST HALF YEAR |                 | FULL YEAR       |                 |
|------------------------------------------------|-----------------|----------------|-----------------|-----------------|-----------------|-----------------|
|                                                | 2026            | 2025           | 2026            | 2025            | 2026            | 2025            |
| <b>OPERATING ACTIVITIES</b>                    |                 |                |                 |                 |                 |                 |
| Net result before taxes                        | (8,498)         | (3,343)        | (13,601)        | (1,390)         | (13,601)        | (10,813)        |
| Depreciation                                   | 914             | 779            | 1,799           | 1,558           | 1,799           | 3,192           |
| Change in current assets                       | 274             | —              | —               | 13              | —               | 13              |
| Change in current liabilities                  | (3,770)         | (4,423)        | (5,903)         | (12,242)        | (5,903)         | (14,979)        |
| Conversion of debts / dividend                 | —               | 609            | —               | 1,066           | —               | 3,721           |
| <b>Net cash from operating activities</b>      | <b>(11,080)</b> | <b>(6,379)</b> | <b>(17,705)</b> | <b>(10,995)</b> | <b>(17,705)</b> | <b>(18,866)</b> |
| <b>INVESTING ACTIVITIES</b>                    |                 |                |                 |                 |                 |                 |
| Investments in non-current assets              | (572)           | —              | (962)           | 10,815          | (962)           | 9,083           |
| <b>Net cash from investing activities</b>      | <b>(572)</b>    | <b>—</b>       | <b>(962)</b>    | <b>10,815</b>   | <b>(962)</b>    | <b>9,083</b>    |
| <b>FINANCING ACTIVITIES</b>                    |                 |                |                 |                 |                 |                 |
| Proceeds from equity issues                    | 13,229          | 2,925          | 20,092          | 9,055           | 20,092          | 19,089          |
| Other financing activities                     | —               | 410            | —               | 410             | —               | —               |
| Translation differences                        | 188             | —              | 154             | (23)            | 154             | (22)            |
| <b>Net cash from financing activities</b>      | <b>13,417</b>   | <b>3,335</b>   | <b>20,246</b>   | <b>9,442</b>    | <b>20,246</b>   | <b>19,066</b>   |
| <b>Net change in cash &amp; equivalents</b>    | <b>1,766</b>    | <b>(3,043)</b> | <b>1,578</b>    | <b>9,260</b>    | <b>1,578</b>    | <b>9,281</b>    |
| Cash & equivalents at beginning of period      | 19,606          | 22,816         | 19,794          | 10,513          | 19,794          | 10,513          |
| <b>Cash &amp; equivalents at end of period</b> | <b>21,372</b>   | <b>19,773</b>  | <b>21,372</b>   | <b>19,773</b>   | <b>21,372</b>   | <b>19,794</b>   |

## Key Financial Figures

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

### P&L

### Profit & Loss — H1 2026

#### Operating revenue

Recognized **NOK 5.8m (7.5m)** from European Defence Fund (EDF) project funding, reflecting project milestone timing.

#### Personnel costs

NOK 2.5m (3.2m). **Significant portion relates to the SIS-02/EDF project.**

#### Other operating costs

NOK 15.2m (4.7m). Driven by **SIS-03 study execution and the SIS-02 /EDF project**. R&D/Operating Costs in % 70 %.

#### Net financial items

NOK 0.05m (0.6m).

#### H1 pre-tax loss

**NOK 13.6m loss** (NOK 1.4m loss H1 2025)

### BS / CF

### Balance Sheet & Cash Flow

#### Capitalized assets

IP and patent costs **NOK 13.3m (14.0m)**, depreciated over 5 years.

#### Cash position

**NOK 21.4m** at end of H1 2026 (31.12.2025: NOK 19.8m). Net H1 cash change **+NOK 1.6m** despite the widened R&D burn.

#### Financing activity

H1 raises of **NOK 20.1m** from three Long State placements (Feb, Mar, Apr 2026). **July 2026: additional direct placement of NOK 5.75m** (54.8m shares @ NOK 0.105) further strengthens the balance sheet.

PLACEMENTS   Feb '26: **45m @ 0.0725** · Mar '26: **60m @ 0.06** · Apr '26: **60m @ 0.12** · Jul '26: **54.8m @ 0.105 (direct)**

H1 2026 figures are unaudited. Year columns show comparative full-year 2025 figures from the audited annual report.

## Statement of Changes of Equity

SoftOx Solutions Group | All amounts in NOK 1,000 unless stated otherwise |  
(X,XXX) = negative figures

| CHANGES IN EQUITY                       | SECOND QUARTER |         | FIRST HALF YEAR |         | FULL YEAR |          |
|-----------------------------------------|----------------|---------|-----------------|---------|-----------|----------|
|                                         | 2026           | 2025    | 2026            | 2025    | 2026      | 2025     |
| Equity at end of prior period           | 24,678         | 103,700 | 22,954          | 95,185  | 22,954    | 95,185   |
| Share issues (equity / debt conversion) | 13,229         | 3,991   | 20,092          | 10,531  | 20,092    | 22,810   |
| Loss for the period                     | (8,498)        | (3,343) | (13,601)        | (1,390) | (13,601)  | (10,813) |
| Other changes in equity                 | 188            | (47)    | 152             | (25)    | 152       | (84,227) |
| Equity at end of period                 | 29,598         | 104,301 | 29,598          | 104,301 | 29,598    | 22,954   |

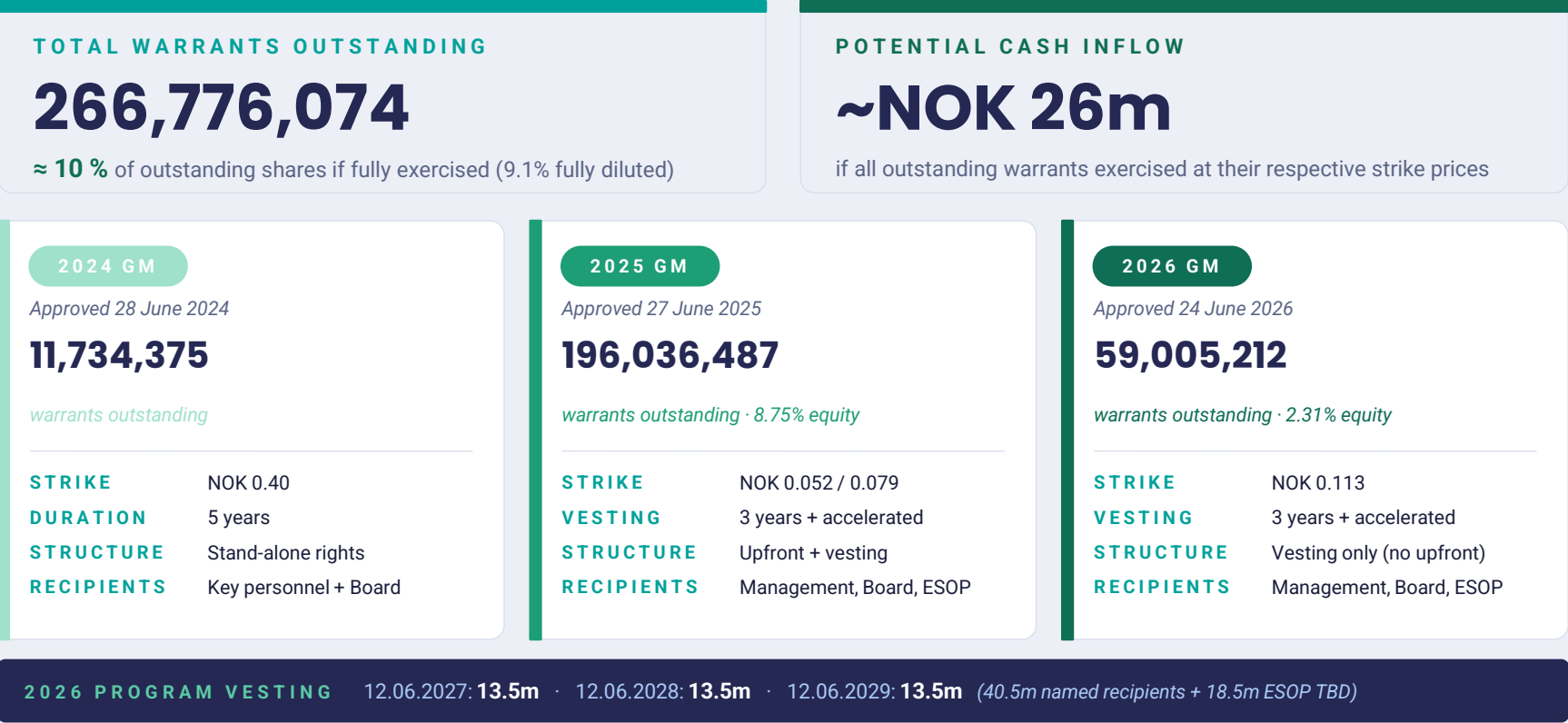
### EQUITY MOVEMENT H1 2026

Net equity change of **+6,644** driven by NOK 20,092 in new equity proceeds (Long State facility placements), partially offset by the H1 loss of NOK 13,601. The 2025 comparable reflects the NOK 84.2m deferred tax write-off recognised in "Other changes".

H1 2026 figures are unaudited. "Other changes in equity" for 2025 full year reflects the de-recognition of the deferred tax asset (NOK 84.2m, see Note 4 in the 2025 annual report).

## Warrants & Incentive Program

Three warrant programs are in force following the 24 June 2026 General Meeting.



# General Notes

*Basis of preparation, accounting principles, and material disclosures for the interim period*

## NOTE 1

### Basis of preparation

The interim financial statements have been prepared in accordance with the **Norwegian Accounting Act/Norwegian GAAP** for other enterprises ("Øvrige foretak"), applying the same accounting policies as the 2025 annual report.

*Figures are unaudited.*

## NOTE 2

### Currency & presentation

Functional and presentation currency: **Norwegian Krone (NOK)**.

All amounts in **NOK 1,000** unless otherwise stated. Foreign currency transactions are translated at the exchange rate on the transaction date; FX gains/losses are recognised in the income statement.

## NOTE 3

### Going concern

The financial statements have been prepared on a **going concern basis**, supported by the cash position at quarter-end and the Long State financing facility (up to NOK 80m available, see Note 5).

Management continues to assess additional financing to fund operations through 2027.

## NOTE 4

### Significant estimates

- **IP & patent costs** — straight-line over 5 years.
- **Deferred tax** — disclosed in notes only (see 2025 AR Note 4).
- **EDF grant** — recognised as project costs are incurred.

## NOTE 5

### Equity & financing

Long State Investments facility (Aug 2025): up to **NOK 50m / 24m**, extendable to **NOK 80m / 36m**. Equity line of credit; Company not obliged to draw.

Placements completed in Q2 2026 and after period-end disclosed in the changes-in-equity statement.

## NOTE 6

### Subsequent events

Events after the balance sheet date:

- **July 2026** — Direct Placement: 54.8 m shares @ NOK 0.105.
- **Clinical milestones** — First patient with chronic airway infection dosed in the SIS-03 Proof-of-Concept patient phase (August 2026), marking the initiation of patient PoC evaluation.

## NOTE 7

### Legal matters

Per stock notice 27 Feb 2025, a former-consultant bonus claim was settled: MNOK 1.5 + MNOK 0.8 (ex VAT, latter due 30.06.2026). The claim is fully paid per H1 26. An IP-rights matter was settled by issuing 16.5m shares to the counterpart.

*No other material proceedings pending or threatened.*

## NOTE 8

### Patent strategy

The Company pursues an **active patent strategy** — including improvements, pruning of the existing portfolio, and filing of new applications to further protect the SoftOx technology platform.

*The Company is advised by a qualified external IP / patent advisory team.*

## FINANCIAL STATEMENTS

# General Accounting Principles

*Prepared in accordance with the Norwegian Accounting Act /GAAP for other enterprises ("øvrige foretak").*

### CONSOLIDATION

Includes subsidiaries where control exists (>50%). Intercompany transactions eliminated. Acquisition method for business combinations; equity method for 20–50% associates.

### USE OF ESTIMATES

Management has applied estimates and assumptions affecting reported amounts of assets, liabilities, income, and expenses.

### FOREIGN CURRENCY

Transactions translated at transaction-date rates. Monetary items at year-end rates; differences recognized in the income statement.

### REVENUE RECOGNITION

Recognized when control and risk transfer, or in line with project completion, where outcomes are reliably measurable.

### RESEARCH & DEVELOPMENT

Development costs are capitalized only when the criteria are met; otherwise, expensed. Research costs are expensed as incurred.

### CASH FLOW STATEMENT

Prepared using the indirect method. Cash and cash equivalents include cash, bank deposits, and short-term liquid investments.

### BALANCE SHEET CLASSIFICATION

Assets and liabilities are classified as current or non-current based on maturity. Assets are measured at historical cost unless otherwise required by accounting principles.

### PP&E

Recognized at cost and depreciated over useful life. Impairment is recognized when the carrying value exceeds the recoverable amount.

### INCOME TAX

Tax expense includes current and deferred tax. Deferred tax assets recognized only to the extent probable; otherwise disclosed in notes.

### INVENTORIES

Measured at the lower of cost (FIFO) and net realizable value.

### RECEIVABLES & PENSIONS

Receivables at nominal value, less provisions for expected losses. Defined contribution pension schemes.

### SUBSIDIARIES

Investments are recorded at cost less impairment.

# Risk Factors & Risk Management

SoftOx's risk management is an integral part of operations, with ongoing monitoring by management and the Board. Risks are grouped across four principal categories.

## OPERATIONAL RISK

- Clinical trial outcomes and timelines
- Regulatory requirements and approvals
- CRO and partner dependency

## FINANCIAL RISK

- Liquidity and access to capital
- FX exposure (DKK, EUR, USD)
- Interest rate risk on cash

## COMMERCIAL RISK

- Market access, pricing, reimbursement
- Competition in respiratory infection
- Strength of partnering proposition

## LEGAL & IP RISK

- Patent prosecution and enforcement
- Regulatory compliance
- No material proceedings pending (2025)

Risks are mitigated through study design, CRO partnerships, diversified IP, disciplined cash management, and continuous board-level oversight.



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## LEGAL

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