



An innovative pharmaceutical
company with a broad and strong
portfolio of projects for the treatment
of cancer and infections

Q3

INTERIM REPORT JANUARY– MARCH 2026

The "Company" or "Hamlet BioPharma" refers to Hamlet BioPharma AB,
corp. reg. no. 556568-8958

OPENING REMARKS

The third quarter of 2025-2026 has been extremely productive for Hamlet BioPharma, with high-profile Phase II trial publications, establishment of clinical Phase III trial infrastructure, drug production and partnering of the tuberculosis project with a Korean drug formulation specialist company. Ongoing, positive partnering discussions with a major player in the bladder cancer field, under a letter of intent, specifically address the commercialization of the bladder cancer asset Alpha1H. In parallel, we are securing and extending the IP portfolio.

Hamlet BioPharma is driving a shift in infectious disease management. The infectious disease treatment assets target the disease response of the host rather than the bacteria, affecting both antibiotic-sensitive and resistant pathogens.

- A successful Phase II clinical study compared host directed therapy to antibiotics. The IL-1 receptor antagonist was shown to inhibit the overactive immune response that drives disease in patients with acute cystitis, with similar efficacy as antibiotics. Clinical data show a reduction in symptoms and recurrences and increase in quality of life.
- A leading international journal, Nature Microbiology has chosen to publish the study (<https://www.nature.com/articles/s41564-026-02262-1>).
- The trials of anti-infective therapies will be extended to include more severe infections.
- Additionally, immunotherapy has shown positive effects in patients with bladder pain syndrome. Clinical data from the Phase II study show a reduction in pain scores and an improved quality of life in patients treated with anakinra.

In the cancer area, the Company's immediate goal is to conduct Phase III studies, obtain market approval for Alpha1H as a treatment for bladder cancer and develop clinical programs for other indications. The investigative new drug Alpha1H kills cancer cells by inducing apoptosis and has shown therapeutic efficacy in Phase II trials without significant side effects. Hamlet BioPharma has received feedback from the FDA for a first in class neo-adjuvant therapy for patients with low risk non-muscle invasive bladder cancer (NMIBC). Intense development activities, regulatory contacts with the FDA and interactions with study sites are ongoing.

Hamlet BioPharma recently signed a partnering agreement with the South Korean ImmunoForge. The Korean partner develops technology for drug release, which is suitable for prolonged delivery of the antibacterial peptide to infected tissues. Tuberculosis is a concern in South Korea and world-wide, creating a special need for novel antibacterial agents to be used against tuberculosis

Hamlet BioPharma has further signed a Letter of Intent outlining the terms of a potential collaboration and a commercial agreement concerning Alpha 1H. The dialogue with our potential partner has been very constructive and forward-looking. At this stage, the focus is on the completion of clinical development, manufacturing scale-up and establishing the right commercial framework to plan market access. This intended co-development would position us to deliver long-term value to patients, healthcare systems and shareholders, subject to successful study outcomes and market approval.

To date, the company has delivered three successful Phase II studies on a budget of less than SEK 300 million, including the cancer project in Phase III and the successful development of alternatives to antibiotics. This track record should maintain the trust of investors regarding the company's ability to deliver on strategic goals at low cost and high quality. We would like to thank our multi-national team, as well as our external partners and collaborators, who are behind the successes and remain a resource for the future. We are grateful for the strong investor support and will continue to use our capital effectively for optimal development and commercialization of our drugs.

Jakob Testad
CEO

Catharina Svanborg
Chairman of the Board

SIGNIFICANT EVENTS

Novel treatments of bacterial infections

On January 16th – Hamlet BioPharma welcomed interested parties to the Symposium “**Novel treatments for Infections and Cancer**” that took place on the 21st of January. The invited speakers are leading international scientists from the U.S. with a background in infection biology and scientists from Lund University, sharing cutting-edge research and future perspectives on novel therapies for bacterial infections and cancer.

On February 23rd – Nature Microbiology published the paper “Targeted innate immune inhibition therapy compared with antibiotics for recurrent acute cystitis: a randomized, open-label phase 2 trial.” the paper was accepted on February 12th.

This groundbreaking study introduces a new approach to managing recurrent urinary tract infections. Instead of killing the bacteria, the treatment focuses on targeting and calming the disease response. Clinical efficacy of this approach is demonstrated in this study, showing effects comparable to standard antibiotic therapy.

This discovery could also mark a major conceptual breakthrough in the fight against antibiotic resistance and the future treatment of infections.

On February 3rd – Hamlet BioPharma signed a collaboration agreement with the Korean biotech company *ImmunoForge*, to develop novel drug delivery technology for the peptide drug NZX, for use in patents with tuberculosis. *ImmunoForge* based in Seoul, South Korea, contacted Hamlet BioPharma to initiate this collaboration. The company who are developing novel drug delivery techniques for clinical use, will develop novel administration technology for the NZX peptide, specifically a slow-release technology that would prolong the effect of each treatment. The future rights to the results of the collaboration will be jointly owned by the two companies, as well as future patent rights.

Cancer Therapy

On March 6th – Hamlet BioPharma signed a Letter of Intent to outline terms of a potential collaboration and a commercial agreement concerning Alpha 1H.

Hamlet signed a non-binding Letter of Intent (LoI) with an undisclosed company specializing in uro-oncology based in northern Germany. The purpose is to outline terms and conditions for the completion of development and for global commercialization of Alpha 1H in the field of bladder cancer.

Company and Finance

On March 15th – a new supplementary subscription period was announced, from 16-23 of March 2026. The Board of Directors had decided to adjust the terms and conditions for warrants of series TO5B ahead of the next subscription period, which runs from 20 July to 31 July 2026.

On March 24th – Hamlet BioPharma announced that the company had received approximately 9 million SEK through subscription of warrants series TO5B, before issue costs.

On February 13th – Hamlet published the Q2 interim report for October to December 2025.

Investor Relations

During the fiscal year 2025/2026, Hamlet BioPharma has continued its series of digital investor meetings held on July 25th, August 28th, September 25th, October 31st November 14th, January 21st, February 13th, March, 18th and April 23rd.

On the 30th of March, it was announced that Hamlet BioPharma presented at the Financial Stockholm event on March 26, 2026.

Additional significant event after the third quarter




On May 18th, Hamlet Biopharma announced the successful granting of new patents in Japan and India and a further patent allowance in Japan. This continued strengthening of Hamlet BioPharma’s international intellectual property portfolio reaffirms its position at the forefront of technological innovation.

COMPANY OVERVIEW

HAMLET BIOPHARMA TRANSLATES INNOVATION INTO CLINICAL SUCCESS

Hamlet BioPharma is a pharmaceutical company focused on developing innovative treatments for cancer and infectious diseases. With a mission to address large, unmet medical needs, the company has built a robust pipeline of therapeutic candidates targeting malignant tumors and antibiotic-resistant infections. These advances underscore Hamlet BioPharma's potential in innovative drug development that addresses important medical needs.

Translating Innovation Into Clinical Success: Three positive Phase II studies

Drug Candidate	Indication	Discovery	Pre-clinical	Clinical	Phase II	Phase III
Alpha1H	Bladder Cancer					FDA Fast track
IL-1 receptor antagonist (anakinra)	Recurrent Urinary tract infection					
	Bladder Pain Syndrome					

- Strong clinical pipeline with advanced assets in bladder cancer, infection, and pain.
- Powerful discovery platform driving innovations in oncology and antibacterial treatments.
- Proven drug development expertise, offering valuable long-term partnership opportunities.

Successful Phase II studies in three clinical indications, driven by research originating from Hamlet's laboratories, with findings published in top journals like Nature.

- 1. Bladder Cancer (Alpha1H)** – Placebo controlled study demonstrated 88% tumor reduction at higher doses. Striking induction of apoptosis, immune activation and down-regulation of cancer genes.
- 2. Recurrent Urinary Tract Infections (Anakinra)** – Two arm study showed that anakinra was as effective as antibiotics, reducing symptoms and recurrence rates.
- 3. Bladder Pain Syndrome (Anakinra)** – Significant reduction in pain and improved quality of life for patients with severe bladder pain.

NEW CANCER TREATMENTS

Alpha1H – Successful Phase II Completed in Bladder Cancer

Alpha1H – Proven Clinical Anti-tumor Effect

Dose-dependent efficacy, with 88% of patients treated with the higher dose experiencing partial or complete responses. Cancer specific effect. Healthy tissues do not take up the drug.

Tumor cell **death by apoptosis** – striking difference between tumor and healthy tissue and low toxicity.

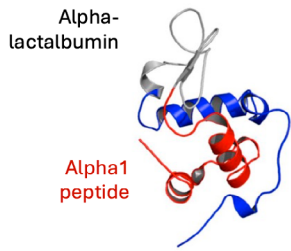
Strong **immune activation** profile similar to BCG treatment.

Down-regulation of cancer genes in the tumor – return towards a non-cancerous healthy profile.

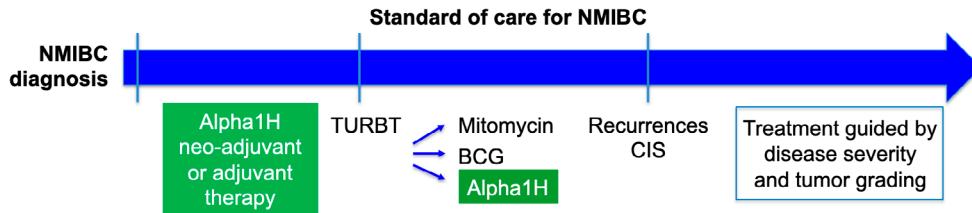
Commercial opportunity, bladder cancer

- ✓ Fast Track FDA designation with strong clinical data completed Phase 2.
- ✓ Positive FDA discussions ongoing for Phase III.
- ✓ GMP-ready for Phase III & commercialization.
- ✓ First-in-Class Neoadjuvant Non-Muscle Invasive Bladder Cancer Therapy.
- ✓ Strong IP positions with long lifetime.
- ✓ Manufacturing & CMC established.

Bladder Cancer treatment with Alpha 1



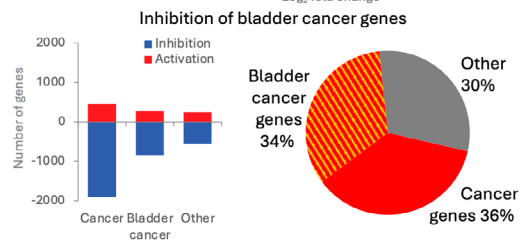
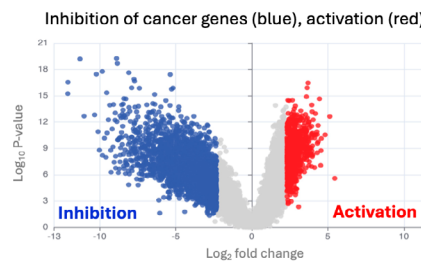
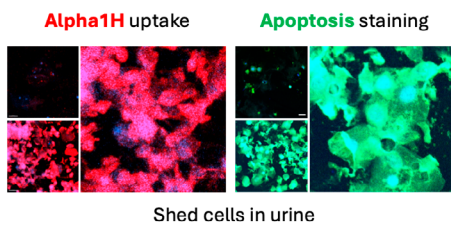
- Alpha1H – broad tumoricidal effects
- Synthetic, peptide-based GMP manufacturing
- Documented long-term stability
- International patents valid >2038
- Therapeutic effects in animal models of bladder cancer
- Documented response in patients with bladder cancer
- Low toxicity



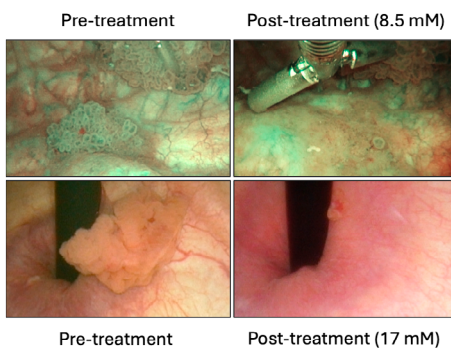
Rapid treatment response (2 hours)

Acute tumor response measured in urine Pre vs. post treatment samples

- Tumor cell shedding into the urine
- Alpha1H uptake by tumor cells
- Apoptosis in tumor cells
- Inhibition of cancer gene expression, urine RNA



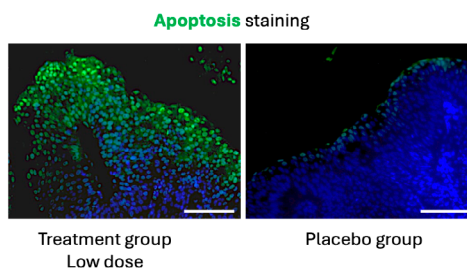
Treatment response (1 month) – effects on the tumor

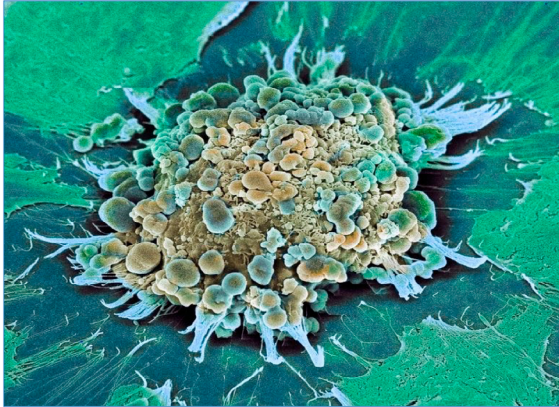


Treatment resulted in a complete or partial response in 82% of the tumors treated with the higher dose and in 45% treated with the lower dose of Alpha1H.

Analysis of treated tumors and tumor tissue

- Reduction in tumor number and size
- Alpha1H uptake by the tumor
- Apoptosis in tumor tissue
- Inhibition of cancer gene expression





The molecules

- Kill tumor cells broadly
- Remove tumor tissue
- Can be used for many cancer types
- Act without damaging healthy tissues

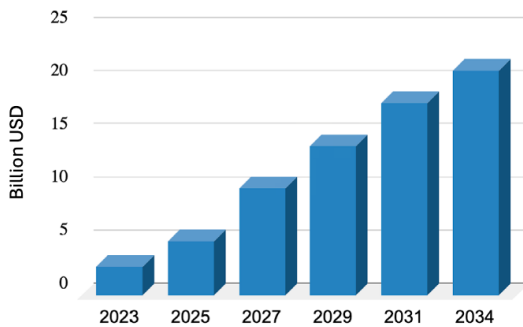
Examples of future indications

- Bladder cancer
- Colon cancer
- Skin tumors
- Brain tumors

MARKET POTENTIAL BLADDER CANCER

The need for effective new drugs against cancer and infections is huge globally and is constantly increasing. Hamlet BioPharma's drug candidates are well positioned to capture significant shares in these fast-growing markets and to address critical needs and position itself for sustainable growth.

Market size, Bladder Cancer 2023-2034

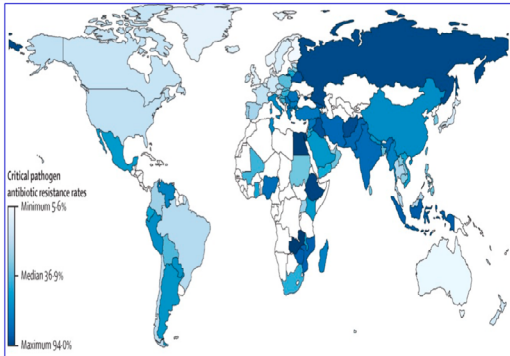


Source: Transparency Market Research <https://www.transparencymarketresearch.com/non-muscle-invasive-bladdercancer-market.html>

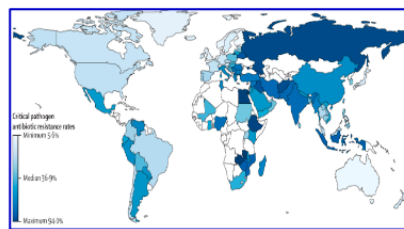
Alpha1H, with its promising clinical results and immune activation profile, is well suited to address the need for novel NMIBC treatment and to become a valuable asset for the global bladder cancer market. Its innovative mechanism of action – targeting cancer cells while sparing healthy tissue – could make Alpha1H a preferred alternative to traditional treatments, which are often associated with high toxicity and frequent relapses.

TREATING BACTERIAL INFECTIONS WITHOUT ANTIBIOTICS

New Non-antibiotic Infection treatment Paradigm



New Infection Treatment Paradigm: Immunotherapy as an Equally- Efficacious Alternative to Antibiotics
 Harnesses the immune response rather than directly targeting bacteria.
 Offers a proven, non-antibiotic approach for clearing antibiotic-resistant pathogens.



High antibiotic resistance rates for critical pathogens, world-wide

- Acute cystitis is a common infection of the urinary bladder, frequently caused by resistant bacteria
- **Phase II study – Blocking the IL1 receptor is as Effective as Antibiotics – IL-1RA shows comparable outcomes to antibiotics in treating bacterial infections (recurrent cystitis)**
- Evidence of effects against multi-resistant bacteria in animal models.
- **Breakthrough New Infection Treatment Paradigm – Balances the immune response instead of killing bacteria with antibiotics.**

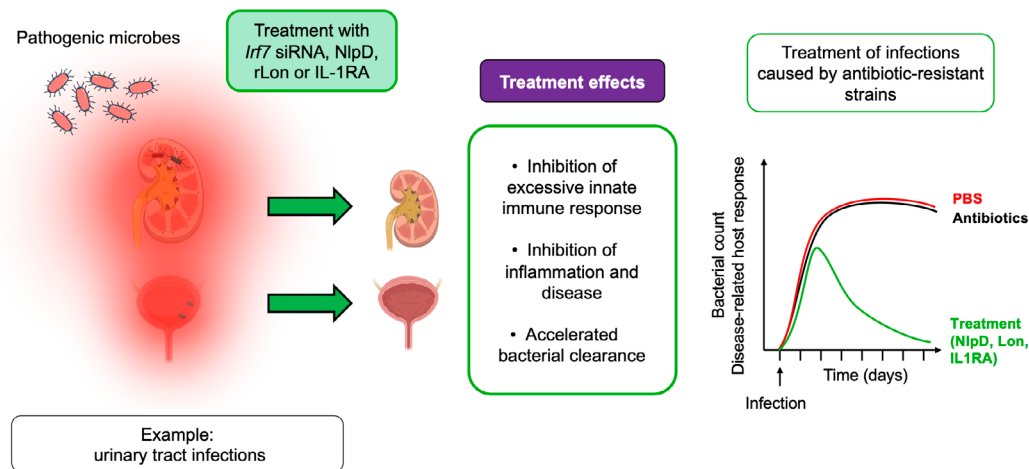
Outcome of two-armed Phase II study

- **Reduced symptoms**
- **Reduced recurrence rates**
- **improved quality of life**
- **Inhibition of immune hyperactivation**

Advantages of acute immunotherapy

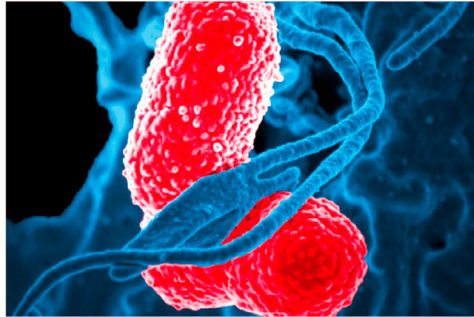
- Provides alternatives to antibiotics
- Reduces selective pressure on resistance
- Reduces effects on the normal flora in individual patients and in the population

Our molecules target the disease rather than the bacteria – new non-antibiotic treatments



Clinical Phase II data supports the efficacy of non-antibiotic therapy of bacterial infections

- Acute cystitis is a common infection of the urinary bladder, frequently caused by resistant bacteria
- Phase II study – shows comparable outcomes to antibiotics in treating bacterial infections (recurrent cystitis)
- Outcome of two-armed Phase II study
 - Reduced symptoms
 - Reduced recurrence rates
 - improved quality of life
 - Inhibition of immune hyperactivation
- Advantages of acute immunotherapy
 - Provides alternatives to antibiotics
 - Reduces selective pressure on resistance
 - Reduces effects on the normal flora in individual patients and in the population



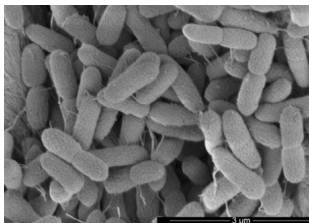
Infections - a major cause of disease and mortality

TREATMENT OF MYCOBACTERIUM TUBERCULOSIS

Tuberculosis is a major cause of mortality in all parts of the world and treatment options are becoming more limited, due to escalating antibiotic resistance. The antimicrobial peptide NZX is being developed as a drug candidate for pulmonary tuberculosis treatment, in collaboration with Hamlet BioPharma. The new, peptide-based drug has shown promising treatment effects against lung tuberculosis in animal models, both against antibiotic sensitive and antibiotic-resistant Mycobacterium tuberculosis bacteria.

The peptide NZ2114 has been identified as a promising candidate for future treatment of tuberculosis. NZ2114 can cross the complex, lipid-rich membrane of Mycobacterium and kill the bacteria in the laboratory and in animal models. NZ2114 is stable in serum and is not toxic to human cells. Antimicrobial effects were also observed against several clinical isolates of Gram-positive bacteria, including methicillin-resistant Staphylococcus aureus (MRSA). NZ2114 is active as a stand-alone treatment and has synergistic effects with established anti-tuberculosis treatments. The peptide eliminated M. tuberculosis in an animal model of tuberculosis with an 81.14% reduction after three doses, compared to untreated controls.

Peptide induced *M. bovis* membrane changes



Untreated



NZ2114 treated

Figure legend. Mycobacteria treated with 6.3µM NZ2114 for 24 hours and visualized with scanning electron microscopy. The image shows how the bacterial membrane is destroyed by NZ2114.

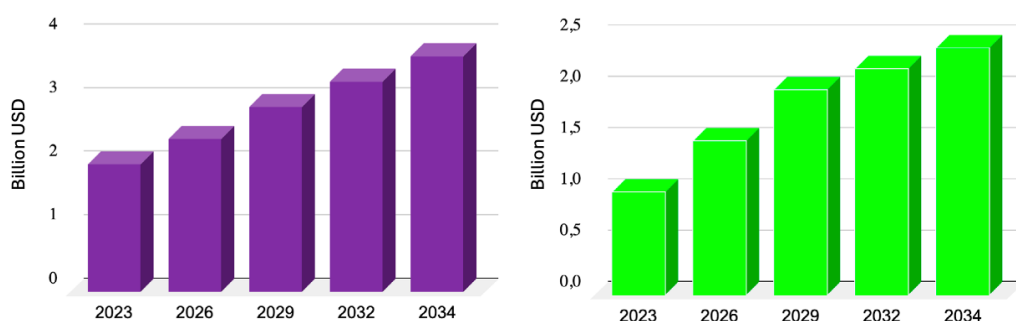
Clinical Phase II data supports therapeutic efficacy in debilitating Bladder Pain Syndrome

- **Rationale:** Bladder pain syndrome (BPS) involves chronic inflammation of the bladder wall, driven by dysregulated immune signaling. Anakinra blocks IL-1 β pathways, reducing inflammatory pain responses.
- **Clinical Proof:** Early studies showed symptom improvement (pain reduction, urinary function) in patients with chronic bladder pain treated with anakinra. Phase II trial shows positive outcome data.
- **Advantages:** Novel, targeted non-opioid approach for chronic bladder pain, where treatment options are currently poor. Repurposing an existing biologic accelerates clinical development.
- **Market:** Chronic bladder pain affects millions globally, with few satisfactory treatments. Targeting a ~\$2–3B unmet need market.

THE INFECTION TREATMENT MARKET

Hamlet BioPharma offers a groundbreaking solution to address the problem of severe bacterial infections and antibiotic resistance. By focusing on the disease rather than the bacteria, the advances could lead to a paradigm shift in the treatment of bacterial infections. The patient's health can thus be improved, and bacteria can be cleared through normal defense mechanisms, rather than directly targeting the bacteria. In this way, infections caused by antibiotic-resistant and antibiotic-sensitive organisms can be targeted in animal models. This approach also reduces the selective pressure that contributes to resistance in the environment and the general population.

Market size Recurrent Acute Cystitis and Bladder Pain Syndrome 2023-2034



Source: Data are estimated values to represent market development from reports and research from the American Urological Association (AUA) and International.

Robust Pipeline – 180 patents, patent classes

Preclinical		
Alpha1H	Brain tumor	Positive data in animal model, development of technology
Hamlet	Colon and rectal cancer	Positive data in animal model
Hamlet	Oral cancer	Preclinical evaluation
NK1R-receptor antagonist	Pain and nerve activation inhibitors	Positive data in animal model, development of technology. Preparation of substance for clinical studies
RNA Pol II inhibitor - protein	Preventive anti-inflammatory and antibacterial effects	Positive data in animal model, development of technology. Preparation of substance for clinical studies
RNA Pol II inhibitor - bacteria	Prevention of inflammation and treatment of infection	Positive data in animal model, development of technology
IRF7 inhibitor, siRNA	Inhibits severe bacterial infections	Positive data in animal model, technology development. Data to support the development of drugs for clinical trials
Anti-TBC peptide	Pulmonary tuberculosis	Positive data in animal model, development of technology for drug production

Market strategy – overall goals

Hamlet Biopharma's overall goal is to

- reach the market with Phase III ready assets in cancer and infection
- drive the clinical development of groundbreaking preclinical assets with proven therapeutic efficacy in animal models through clinical trials to partnering.

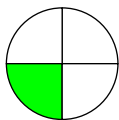
Financing

- **Phase III ready asset Alpha1H**
FDA and EMA market approval
- **Phase II validated** infection and pain indications
New indications for established drug
- **Building the company**

Partnering

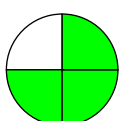
- Specialist or Big pharma, providers of market access
- Development of preclinical/clinical assets

THE PERIOD IN SUMMARY



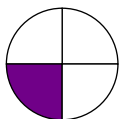
THIRD QUARTER, JAN 1, 2026-MAR 31, 2026 (THE PARENT COMPANY)

- Net sales totaled KSEK 0 (0)
- EBITDA amounted to KSEK -9,561 (-9,855)
- EBIT amounted to KSEK -11,615 (-11,992)
- Net result amounted to KSEK -11,612 (-12,007)
- Earnings per share* was SEK -0.0629 (-0.0676) and -0.0585 after dilution



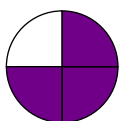
FIRST NINE MONTHS, JUL 1, 2025-MAR 31, 2026 (THE PARENT COMPANY)

- Net sales totaled KSEK 0 (0)
- EBITDA amounted to KSEK -32,212 (-29,504)
- EBIT amounted to KSEK -38,356 (-35,911)
- Net result amounted to KSEK -38,203 (-35,660)
- Earnings per share* was SEK -0.2069 (-0.2007) and -0.1923 after dilution
- On March 31, 2026, the equity/assets ratio** was 88.0 (92.4) %
- Cash amounted to KSEK 13,986 (20,002)



THIRD QUARTER, JAN 1, 2026-MAR 31, 2026 (THE GROUP)

- Net sales totaled KSEK 0 (0)
- EBITDA amounted to KSEK -9,561 (-9,855)
- EBIT amounted to KSEK -12,114 (-12,490)
- Net result amounted to KSEK -12,111 (-12,505)
- Earnings per share* was SEK -0.0656 (-0.0704) and -0.0610 after dilution



FIRST NINE MONTHS, JUL 1, 2025-MAR 31, 2026 (THE GROUP)

- Net sales totaled KSEK 0 (0)
- EBITDA amounted to KSEK -32,212 (-29,504)
- EBIT amounted to KSEK -39,853 (-37,407)
- Net result amounted to KSEK -39,699 (-37,157)
- Earnings per share* was SEK -0.2150 (-0.2091) and -0.1999 after dilution
- On March 31, 2026, the equity/assets ratio** was 86.2 (91.9) %
- Cash amounted to KSEK 14,036 (20,027)

Amounts in parentheses above and below indicate the corresponding value in the preceding year.

* Profit/loss after tax for the period divided by 184,661,871 (177,685,127), respectively 198 615 359, where 184,661,871 is the number of shares outstanding on March 31, 2026, and 198 615 359 shares constitute the number of shares that the Company will have if all subscribed units in the rights issue are paid and exercised. The comparative figure in parentheses was the number of shares on March 31, 2025.

** Equity divided by total capital.

Revenue

Hamlet Biopharma is specializing in the development of drugs with a broad and strong portfolio of projects for the treatment of cancer and infections. The company remains pre-revenue, with value creation driven by milestones in development and partnerships. Net sales amounted to KSEK 0 (0) during the quarter. Other operating income amounted to KSEK 53 (208) during the quarter.

Earnings

External costs were related to the continued drug development activities of the research team at Lund University. The team at Lund University is also responsible for the development of manufacturing methods, stability testing, and chemical and functional characterization of existing and new drug substances and plays a key role in the coordination of laboratory testing in the clinical trial. EBITDA for the quarter amounted to KSEK -9,561 (-9,855), and for the first nine months KSEK -32,212 (-29,504). The depreciations in the quarter were KSEK -2,054 (-2,136), and for the first nine months KSEK -6,145 (-6,407). EBIT for the quarter amounted to KSEK -11,615 (-11,992), and for the first nine months KSEK -38,356 (-35,911). Net result for the quarter was KSEK -11,612 (-12,007), and for the first nine months KSEK -38,203 (-35,660).

Financial position

The company was provided with approximately SEK 9 million through the subscription of warrants series TO5B, so-called short options, during March/April 2026. The issue costs were stated at SEK 35 thousand. Registration with the Swedish Companies Registration Office took place after the quarter, on April 2, 2026. At the end of the quarter, the equity/assets ratio was 88.0 (92.4)%, and the Company's cash and cash equivalents were KSEK 13,986 (20,002).

Investments

The Company does not capitalize expenses for research and development as assets, since the Company is in an R&D stage. R&D costs are therefore recognized as operating expenses in the income statement.

Depreciation

During the quarter, depreciation of equipment amounted to KSEK 33 (115), and the depreciation of patents from the merger with SelectImmune Pharma AB amounted to KSEK 2,021 (2,021).

In the group, depreciation of patents, including the acquisition of Linnane Projects AB, amounted to KSEK 2,553 (2,635) during the quarter.

Employees

The company had the equivalent of 7 (7) full-time employees during the quarter

The share

The Company's shares have been traded on Spotlight Stock Market since October 23, 2015. The share is traded under the short name "HAMLET B" with ISIN code SE0015661152.

At the extraordinary general meeting in Hamlet Pharma AB on March 2, 2021, it was decided that the company's common shares would undergo a split with relation 3:1 and would be reclassified as A- and B-shares. The B-shares will be traded on Spotlight Stock Market. The A-shares will not be listed. Each A-share entitles to ten votes and B-shares entitles to one vote. Furthermore, it is possible for shareholders to convert A-shares to B-shares, which can be traded on Spotlight Stock Market. This conversion program is ongoing with no current deadline. This means that the ratio between A- and B-shares will change over time.

As of March 31, 2026, the number of shares registered at the Swedish Companies Registration Office (Bolagsverket) totaled 184,661,871. The registered current ratio of shares was 39,947,400 A-shares and 144,714,471 B-shares.

Subscription warrants

The warrants in serie TO5B and serie TO6B give the right to subscribe for a total of 20 930 232 shares. Of these, 1,518,838 shares were subscribed for in series TO5B during March/April 2026. The issue was aimed at approximately 20 external investors. The subscription price for the short option with a term of 12 months is SEK 6 per share and the subscription price for the long option with a term of 30 months is SEK 10 per share. The options are not admitted to trading.

Transactions with related parties

During the quarter, KSEK 1,959 (1,459) was paid to Linnane Pharma AB, of which KSEK 1,839 (1,339) refers to the co-operation agreement, KSEK 120 (120) refers to patent license.

The co-operation agreement with Linnane Pharma refers to compensation for access to advanced science and cutting-edge technology for drug development. The collaboration means that Linnane Pharma's technology platform and other resources are available to Hamlet BioPharma. Hamlet BioPharma is a subsidiary company of Linnane Pharma AB, which owns 33.00% of the capital and 73.96% of the votes of Hamlet BioPharma.

Furthermore, salaries and allowances to board and management were paid during the period. Transactions with related parties are on market terms.

Significant risks and uncertainties

The Board's assessment of significant risks and uncertainties is unchanged compared with the most recent financial year and are described in the most recently published annual report (2025-06-30).

Basis of preparation for the interim report

The Company prepares its accounts in accordance with the Swedish Annual Accounts Act (Årsredovisningslagen) and the K3 framework (BFNAR 2012:1) of the Swedish Accounting Standards Board (Bokföringsnämnden).

The company's accounting principles are unchanged compared with most recent financial year and are described in the most recent published annual report (2025-06-30).

On March 31st, 2023, Hamlet BioPharma acquired Linnane Projects AB from Linnane Pharma AB and the patents and know-how regarding a new peptide-based drug against tuberculosis as well as the know-how required to develop the project. In accordance with regulations at Spotlight and the Swedish Accounting Standards Board (Bokföringsnämnden), consolidated accounts of Linnane Projects and Hamlet BioPharma are drawn up. The quarterly report is prepared with the parent company's accounting in focus. In texts, the group is only commented on if something differs significantly from the parent company.

Review

This interim report has not been audited.

Financial calendar

Year-end report for 2025/2026	August 28, 2026
Annual Report for 2025/2026	October 30, 2026
Interim report for Q1, 2026/2027	November 13, 2026
Annual General Meeting for 2025/2026	November 20, 2026

INCOME STATEMENT: THE PARENT COMPANY

SEK	2026-01-01 2026-03-31	2025-01-01 2025-03-31	2025-07-01 2026-03-31	2024-07-01 2025-03-31	2024-07-01 2025-06-30
Net sales	0	0	0	0	0
Other operating income	52 933	208 038	52 933	208 038	208 038
Operating income	52 933	208 038	52 933	208 038	208 038
Other external costs	-7 816 503	-8 197 565	-27 276 633	-23 768 332	-36 882 334
Employee benefit expenses	-1 773 856	-1 857 948	-4 935 209	-5 893 776	-7 935 277
Depreciation of assets	-2 054 100	-2 136 285	-6 144 651	-6 406 780	-8 467 222
Other operating expenses	-23 530	-7 970	-52 745	-50 079	-53 767
Operating loss	-11 615 056	-11 991 730	-38 356 306	-35 910 928	-53 130 563
Financial items	2 651	-15 003	153 255	250 586	472 708
Loss before tax	-11 612 405	-12 006 733	-38 203 051	-35 660 343	-52 657 855
Tax on loss for the period	0	0	0	0	0
Loss after tax	-11 612 405	-12 006 733	-38 203 051	-35 660 343	-52 657 855

BALANCE SHEET: THE PARENT COMPANY

SEK	2026-03-31	2025-03-31	2025-06-30
ASSETS			
Fixed assets			
Intangible assets	19 061 153	27 146 357	25 125 056
Tangible assets	489 412	256 301	217 160
Financial assets	10 025 000	10 000 000	10 000 000
Total fixed assets	29 575 565	37 402 658	35 342 216
Current assets			
Other receivables	3 351 728	4 898 746	1 564 911
Prepaid expenses	346 970	263 277	430 274
Cash and bank balances/financial investments	13 985 601	20 002 059	10 735 539
Total current assets	17 684 298	25 164 083	12 730 724
Total assets	47 259 863	62 566 740	48 072 940
EQUITY & LIABILITIES			
Restricted equity			
Share capital	1 846 619	1 776 851	1 776 851
Unregistered share capital	15 188	0	0
Statutory reserve	20 000	20 000	20 000
Total restricted equity	1 881 807	1 796 851	1 796 851
Non-restricted equity			
Share premium reserve	290 629 584	251 751 833	251 751 833
Retained earnings	-212 741 348	-160 083 493	-160 083 493
Loss for the period	-38 203 051	-35 660 343	-52 657 855
Total non-restricted equity	39 685 186	56 007 998	39 010 485
Total equity	41 566 993	57 804 849	40 807 337
Current liabilities			
Accounts payable	3 336 539	1 122 796	2 892 898
Tax liabilities	0	26 428	160 543
Other liabilities	348 196	353 067	358 148
Accrued expenses	2 008 136	3 259 601	3 854 014
Total current liabilities	5 692 871	4 761 891	7 265 604
Total Equity & Liabilities	47 259 863	62 566 740	48 072 940

CASH FLOW STATEMENT: THE PARENT COMPANY

SEK	2025-07-01 2026-03-31	2024-07-01 2025-03-31	2024-07-01 2025-06-30
Operating activities			
Loss after financial items	-38 203 051	-35 660 343	-52 657 855
Adjusted for non-cash items, etc.	6 144 651	6 406 780	8 467 222
Cash flow from operating activities before changes in working capital	-32 058 400	-29 253 563	-44 190 633
Cash flow from changes in working capital			
Change in current receivables	-1 703 513	-962 910	2 203 928
Change in current liabilities	-1 572 733	653 978	3 157 690
Cash flow from operating activities	-35 334 645	-29 562 495	-38 829 015
Investing activities			
Acquisition of tangible assets	-353 000	-24 922	-24 922
Acquisition of financial assets	-25 000	0	0
Cash flow from investing activities	-378 000	-24 922	-24 922
Financing activities			
Rights issue	39 113 027	26 790 008	26 790 008
Issuance costs	-150 320	-276 610	-276 610
Cash flow from financing activities	38 962 707	26 513 398	26 513 398
Cash flow for the period	3 250 061	-3 074 020	-12 340 540
Cash and cash equivalents at the beginning of the period	10 735 539	23 076 079	23 076 079
Cash and cash equivalents at the end of the period	13 985 601	20 002 059	10 735 539

EQUITY: THE PARENT COMPANY

SEK	Share capital	Unregistered share capital	Statutory reserve	Share premium reserve	Retained earnings	Loss for the period	Total
Opening balance July 1, 2025	1 776 851	0	20 000	251 751 833	-160 083 493	-52 657 855	40 807 337
Transfer of prior year's loss					-52 657 855	52 657 855	0
Rights issue	69 767			29 815 011	0	0	29 884 779
Loss for the period, Q1					-11 658 253	-11 658 253	-11 658 253
Loss for the period, Q2					-14 932 393	-14 932 393	-14 932 393
T05 emission		15 188		9 062 740			9 077 928
Loss for the period, Q3					-11 612 405	-11 612 405	-11 612 405
Equity March 31, 2026	1 846 619	15 188	20 000	290 629 585	-212 741 348	-38 203 051	41 566 993

INCOME STATEMENT: THE GROUP

SEK	2026-01-01 2026-03-31	2025-01-01 2025-03-31	2025-07-01 2026-03-31	2024-07-01 2025-03-31	2024-07-01 2025-06-30
Net sales	0	0	0	0	0
Other operating income	52 933	208 038	52 933	208 038	208 038
Operating income	52 933	208 038	52 933	208 038	208 038
Other external costs	-7 816 503	-8 197 565	-27 276 633	-23 768 332	-36 882 334
Employee benefit expenses	-1 773 856	-1 857 948	-4 935 209	-5 893 776	-7 935 277
Depreciation of assets	-2 552 850	-2 635 035	-7 640 901	-7 903 030	-10 462 222
Other operating expenses	-23 530	-7 970	-52 745	-50 079	-53 767
Operating loss	-12 113 806	-12 490 480	-39 852 556	-37 407 178	-55 125 563
Financial items	2 651	-15 003	153 255	250 586	472 708
Loss before tax	-12 111 155	-12 505 483	-39 699 301	-37 156 593	-54 652 855
Tax on loss for the period	0	0	0	0	0
Loss after tax	-12 111 155	-12 505 483	-39 699 301	-37 156 593	-54 652 855
Attributable to					
The parent company's shareholders	-12 111 155	-12 505 483	-39 699 301	-37 156 593	-54 652 855
Holdings without controlling influence	0	0	0	0	0

BALANCE SHEET: THE GROUP

SEK	2026-03-31	2025-03-31	2025-06-30
ASSETS			
Fixed assets			
Intangible assets	23 051 153	33 131 357	30 611 306
Tangible assets	489 412	256 301	217 160
Financial assets	0	0	0
Total fixed assets	23 540 565	33 387 658	30 828 466
Current assets			
Other receivables	3 351 728	4 898 746	1 564 911
Prepaid expenses	346 970	263 277	430 274
Cash and bank balances/financial investments	14 035 601	20 027 059	10 760 539
Total current assets	17 734 298	25 189 083	12 755 724
Total assets	41 274 863	58 576 740	43 584 190
EQUITY & LIABILITIES			
Equity			
Share capital	1 846 619	1 776 851	1 776 851
Unregistered share capital	15 188	0	0
Other contributed capital	290 649 584	251 771 833	251 771 833
Other equity including loss for the period	-256 929 399	-199 733 835	-217 230 098
Total equity attributable to the parent company's shareholders	35 581 993	53 814 849	36 318 587
Holdings without controlling influence	0	0	0
Total equity	35 581 993	53 814 849	36 318 587
Current liabilities			
Accounts payable	3 336 539	1 122 796	2 892 898
Tax liabilities	0	26 428	160 543
Other liabilities	348 196	353 067	358 148
Accrued expenses	2 008 136	3 259 601	3 854 014
Total current liabilities	5 692 871	4 761 891	7 265 604
Total Equity & Liabilities	41 274 863	58 576 740	43 584 190

CASH FLOW STATEMENT: THE GROUP

SEK	2025-07-01 2026-03-31	2024-07-01 2025-03-31	2024-07-01 2025-06-30
Operating activities			
Loss after financial items	-39 699 301	-37 156 593	-54 652 855
Adjusted for non-cash items, etc.	7 640 901	7 903 030	10 462 222
Cash flow from operating activities before changes in working capital	-32 058 400	-29 253 563	-44 190 633
Cash flow from changes in working capital			
Change in current receivables	-1 703 513	-962 910	2 203 928
Change in current liabilities	-1 572 733	653 978	3 157 690
Cash flow from operating activities	-35 334 645	-29 562 495	-38 829 015
Investing activities			
Acquisition of tangible assets	-353 000	-24 922	-24 922
Acquisition of financial assets	0	0	0
Cash flow from investing activities	-353 000	-24 922	-24 922
Financing activities			
Rights issue	39 113 027	26 790 008	26 790 008
Issuance costs	-150 320	-276 610	-276 610
Cash flow from financing activities	38 962 707	26 513 398	26 513 398
Cash flow for the period	3 275 061	-3 074 020	-12 340 540
Cash and cash equivalents at the beginning of the period	10 760 539	23 101 079	23 101 079
Cash and cash equivalents at the end of the period	14 035 601	20 027 059	10 760 539

EQUITY: THE GROUP

SEK	Share capital	Unregistered share capital	Other contributed capital	Other equity incl profit for the period	Total
Opening balance July 1, 2025	1 766 851		251 771 833	-217 230 098	36 318 587
Transfer of prior year's loss					0
Rights issue	69 767		29 815 011		29 844 779
Loss for the period, Q1				-12 157 003	-12 157 003
Loss for the period, Q2				-15 431 143	-15 431 143
Rights issue		15 188	9 062 740		9 077 928
Loss for the period, Q3				-12 111 155	-12 111 155
Equity March 31, 2026	1 846 619	15 188	251 771 833	-256 929 398	35 581 993

The Board of Directors and the Chief Executive Officer assure that the interim report provides a true and fair view of the Company's operations, position, and results.

Malmö, May 22, 2026

Catharina Svanborg
Chairperson of the Board

Jakob Testad
CEO

Bill Hansson
Board member

Magnus Nylén
Board member

Henrik Sundin
Board member

Hamlet BioPharma

Hamlet BioPharma AB
Klinikgatan 32
222 42 Lund
hamletbiopharma.com

For further information:

Catharina Svanborg, Chairman of the Board
Tel: +46 (0)709 42 65 49
E-mail: catharina.svanborg@hamletpharma.com

Jakob Testad, CEO
Tel: +46 (0)708 48 42 10
E-mail: jakob.testad@hamletbiopharma.com