

Interim report, Jan-Jun 2022

- Recruitment target for PHSU05 was achieved according to plan in March
- Last PHSU05 clinic visit concluded in June



Promore Pharma AB (publ)

Interim report January - June 2022

April to June

- Net sales amounted to MSEK 0 (0)
- Net loss was MSEK -6.1 (-7.9), corresponding to earnings per share of SEK -0.10 (-0.22)
- Cash flow after financing activities amounted to MSEK -6.8 (-5.5)
- Cash amounted to MSEK 29.6 (13.1), compared to MSEK 45.3 on 31 December 2021

January to June

- Net sales amounted to MSEK 0 (0)
- Net loss was MSEK -14.5 (-15.0), corresponding to earnings per share of SEK -0.24 (-0.41)
- Cash flow after financing activities amounted to MSEK -15.7 (-11.2)

Significant events during January – June

- In January 2022, warrants corresponding to a dilution of 0.2% of the number of outstanding shares were deregistered.
- In February 2022, the first trial person was enrolled in PHSU05 (ensereptide).
- The recruitment target for the study was achieved according to plan in March.
- In June 2022, the last clinic visit in PHSU05 occurred (ensereptide).
- At the AGM in May, Marianne Dicander Alexandersson was elected new chairman of the board. Also, Candice Jung was elected new member of the board.

Events after the reporting period

- In August 2022, the company received a granted patent in the European Union for the use of the candidate drug ropo-campide (LL-37) for the treatment of chronic wounds.

“I am pleased to say that Promore Pharma, despite the current situation in the world, so far, has been able to follow the operating plan”

Jonas Ekblom, President and CEO of Promore Pharma

Financial overview for the Company

<i>Amounts in MSEK</i>	Apr-Jun		Jan-Jun	
	2022	2021	2022	2021
Net sales	-0.0	0.0	-0.0	0.0
Operating loss	-6.1	-7.8	-14.5	-14.9
Profit/Loss for the period	-6.1	-7.9	-14.5	-15.0
Earnings per share, SEK	-0.10	-0.22	-0.24	-0.41
Cash flow after financing activities	-6.8	-5.5	-15.7	-11.2
Cash and cash equivalents at the end of the period	29.6	13.1	29.6	13.1

Promore Pharma in brief:

Promore Pharma is a biopharmaceutical company that develops pharmaceutical product candidates for bioactive healing of wounds. The company has two drug candidates in late clinical development stages, that are based on endogenous peptides, and thus have a strong safety profile. These two products are intended for treatment of chronic wounds, and prevention of scarring on the skin and other tissues. The company is listed on the Nasdaq First North Growth Market.

Statement of the CEO

The first half of 2022 has been intense and successful. I am pleased to say that Promore Pharma, despite the current situation in the world, so far, has been able to follow the operating plan, and thereby conduct the development at the scheduled rate.

We have made important progress in our project with ensereptide which is aimed as a new treatment to prevent skin scarring. It is very satisfactory to note that the clinical part (recruitment, treatment and follow-up) in our clinical trial PHSU05 has been successfully completed according to plan during the first six months of the year. We have achieved the goal of finalizing the treatment and follow-up of 24 patients in accordance with the clinical study protocol. The project has four fundamental components: (i) a single treatment event with ensereptide or placebo, (ii) a three-month follow-up period, (iii) an extensive histopathological analysis of biopsies collected, and finally, (iv) an analysis phase of the study when the data is quality assured and analyzed in detail by the reviewers defined in the study plan. We expect to be able to unblind the study results and present final data from the study in the first quarter of 2023. At present, work on the histopathological analysis is where we put our largest attention.



If the analysis of data from the ongoing clinical study demonstrates a clearcut treatment effect of ensereptide, we make the assessment that we have very good opportunities to create great values as we address a great medical need, and a gigantic marketplace.

We have also made important progress in our project with ropocamptide which relates to a new treatment of venous leg ulcers, the most common type of chronic wounds. The technological development is currently aimed at creating an improved dosage form that is more user-friendly. Although this work will continue for the rest of 2022, we can already say that a number of significant risks have been removed. We have created a preliminary manufacturing process that seems to be robust, cost-effective and scalable. We are now working with external service providers to validate that this manufacturing process results in an end-product that meets all our specifications.

We also continue the planning work to enable a Phase III trial with ropocamptide in the EU. The company has not yet made any decisions regarding timing of study initiation or financing.

At present, the company has not suffered any extensive consequences of the ongoing conflict in Ukraine. The perceptions of the crisis vary, but one thing experts in various functions seem to agree upon; there are a large number of uncertainties in nearly all sectors of society. For us as a company, it may, for example, mean capacity restrictions in the availability of certain raw materials, changed priorities at other companies that affect the interest in entering license agreements and strategic alliances, as well as uncertainties in the capital markets that may have implications for future capital raises. Our overall plans have so far not been significantly affected, but naturally, we are constantly working to assess how this may affect our situation. We have a stable cash position, and I believe that we have the resources to conduct our work according to plan, at least for a year ahead.

All in all, our progress within the company's research programs gives me great hope for an exciting time ahead of us, awaiting results from our clinical trial of ensereptide, PHSU05, where we expect to have results around the turn of the year. My colleagues and I are hopeful that our future end results will benefit all our stakeholders, not least the patients.

Solna, August 30, 2022

Jonas Ekblom

President & CEO

Interview with Marianne Dicander Alexandersson

You have been a part of the company's board since 2017. Can you describe in a few words what is unique with Promore Pharma?

MDA: Firstly, not many small development companies have two relatively advanced projects in the clinical phase, which Promore Pharma has. Secondly, Promore Pharma works with substances occurring naturally in the body, which leads to a low risk for side effects. Thirdly, the company has very limited competition in the pharmaceutical market within our application areas: medicinal preparation development and scar formation/confluency. The company's product candidates are intended to reach global markets which represent hundreds of billion Swedish crowns in annual turnover.



You took up the position as chairwoman of the board as of this year's shareholders' general meeting. What are your thoughts about this?

MDA: It is a great pleasure to be chairwoman of the board in this interesting company which has great potential to ease the lives of many people as well as to save funds in the health care system. Some remarkable aspects of the company are its professional management, great structural capital, a board with broad experience, and a considerable commitment to the company. The time of exchange of the chairwoman of the board was well planned. Promore Pharma has a stable business plan and capital for the next 12 to 18 months. We intend to have data from an ongoing phase II clinical study of ensereptide around the turn of the year, and the task of improving the ropocamptide product will advance as planned.

Is there anything in particular that you would like to focus on during the next years?

MDA: Today, the company has a firmly established strategy, and the business follows the plan made out by the board of directors very well. The company currently has its main focus on two projects: ropocamptide and ensereptide. I would like to see a further focus on studies in these two areas, while we in the meantime are open to collaborations and eventual expansions to further increase the spreading of risks.

What global trends do you see will affect the conditions for small-scale innovative companies in the biotechnical sector during the coming years?

MDA: The current state of the world, with apprehension for poorer trade conditions and rising rates of interest, is a great challenge. I estimate that the competition for capital will increase. There is a great number of companies, both public and private, that will need to secure procurement of capital in the next two years. The challenge in this type of company is always to acquire sufficient capital.

Summary of facts

Marianne Dicander Alexandersson (MDA) is the new chairwoman of the board of Promore Pharma AB as well as chairwoman of Saminvest AB, Occlutech AB, and Sahlgrenska Science Park AB. She is a member of the board at Linc AB and Oblique AB.

MDA has experience as CEO of Kronans Droghandel AB, Sjätte AP fonden and GHP Global Health Partner AB, as well as vice CEO and accountable for the private customer business at Apoteket AB.

Marianne is a civil engineer and has held office at Volvo, Pharmacia, and ICI AB within sales, market, and quality development.

Previous duties as part of boards of committees has been in Camurus AB, Praktikertjänst AB, Recipharm AB, Enzymatica AB, Mölnlycke Healthcare, Addera Care AB, Chalmers Tekniska Högskola, and Svenskt Näringsliv amongst others.

Overview of activities

Promore Pharma is a biopharmaceutical company that develops peptide-based product candidates aimed at the bioactive wound care market. Ropocamptide (LL-37) has recently passed clinical Phase IIb trial on patients with venous leg ulcers, and ensereptide (PXL01), which is developed for the treatment of post-surgery scars, is undergoing a Phase II proof-of-concept study for the treatment of post-surgery skin scars.

Promore Pharma's product candidates are based on innate peptides, which are a part of the human defense and healing system and have a strong safety profile since they are quickly degraded in the blood stream and are therefore unlikely to contribute to severe systemic adverse events. This is supported by the results from prior clinical studies, where both ensereptide and ropocamptide showed strong tolerability and safety as well as efficacy. The product candidates are protected by several international patent families offering protection until 2030 and longer. The patents provide protection in several dimensions, such as therapeutic use, formulation, and dosage ranges.

Promore Pharma's product candidates represent first-in-category therapeutics for several patient groups, segments where patients experience pain, reduced mobility, and lowered quality-of-life. When Promore Pharma's product candidates in clinical development receive market authorization and are established as treatment for chronic wounds and for preventing adhesions and scars, it would mean shorter treatment times for patients and lower costs for society.

Promore Pharma is a small and cost-effective company without its own laboratories or research facilities, using a network of high-quality contract research organizations and contract manufacturing organizations. The company has experienced advisors in all critical aspects of the strategic planning process, including product development, regulatory affairs, design, and execution of clinical trials. Promore Pharma's overall strategy is to take the product candidates through clinical development to market authorization or to a point when a license agreement, alternatively a commercial deal with a larger pharmaceutical company with global presence, can be realized. Such transactions may include out-partnering/licensing, strategic partnerships, joint ventures, or asset sales.

About ensereptide (PXL01)

Ensereptide is derived from a human anti-bacterial protein (lactoferrin), which is part of the innate immune system. This protein and its peptide fragments have several modes of action, including immunomodulation and enhancement of fibrinolytic activity. It is well established that inflammation and fibrin formation after surgery or trauma are two pivotal mechanisms that strongly contribute to scar formation.

Ensereptide is aimed at local administration, and the development of the product is focused on preventing different kinds of scarring after surgery. In a Phase II clinical study that has been completed by the company in several countries of the European Union (EU), it has been demonstrated that ensereptide is efficacious and safe. Promore Pharma is conducting a clinical Phase II trial in the EU to explore the efficacy of the product for prevention of skin scarring. The study was initiated according to plan in the beginning of 2022.

Every year, more than 300 million surgical procedures are performed worldwide, and a proportion of these procedures result in disfiguring skin scars, for example after plastic and trauma surgery. Today, there are no drug products for prevention skin scarring after surgery. The addressable market is estimated to exceed SEK 100 billion. In other types of surgical procedures, there is a risk for occurrence of internal scars, which can cause adhesions (unfavorable attachments of tissues). This is a major medical problem, for example after surgical repair of injured tendons in the hand.

About ropocamptide (LL-37)

Ropocamptide is based on a human antimicrobial peptide, which stimulates several processes in wound healing. In a clinical Phase IIa study conducted by the company in patients with venous leg ulcers (VLUs), ropocamptide showed, in the most effective dose, an increase in the healing rate of relative wound area reduction of close to 70% after one month's treatment, suggesting a significantly higher efficacy than what has been reported for any other treatment in chronic wounds. No serious adverse events that were deemed to be caused by the investigational product occurred in the trial. The product candidate can be easily combined with the standard wound care treatments and given by a nurse or the patient.

The development of ropocamptide is initially focused on venous leg ulcers and the company has recently concluded a clinical Phase IIb study (HEAL LL-37) on patients with VLUs in Europe. VLUs constitute the largest category of all chronic or hard-to-heal ulcers and represent significant challenges to patients and healthcare systems since they are frequent, costly to manage, recurring, and may persist for months or years.

The development of ropocamptide focuses initially on VLUs but the company sees good potential in also developing ropocamptide for diabetic foot ulcers.

Significant events during January – June 2022

Deregistration of warrants

In January 2022, warrants related to program 1 & 2, corresponding to a dilution of 0.2% of the number of outstanding shares, were deregistered.

First patient in PHSU05 enrolled

In the middle of February, the first patient of approx. 24 was enrolled in PHSU05, the company's Phase II study for the prevention of scars in conjunction with surgery.

Recruitment goal reached in clinical trial of ensereptide

In March it was announced that the recruitment goal has been accomplished according to plan in the company's Phase II study (PHSU05) with the company's drug candidate ensereptide for the prevention of skin scarring.

The last clinic visit in the company's Phase II study on ensereptide

In June it was announced that the last clinic visit was carried out for the subjects who participated in the company's Phase II study PHSU05 with ensereptide against skin scarring.

Changes in the Board of Directors

At the AGM in May, Marianne Dicander Alexandersson was elected new chairman of the board. Also, Candice Jung was elected new member of the board.

Events after the reporting period

Patent for ropocamptide in Europe

In August the company received a granted patent in the European Union for the use of the candidate drug ropocamptide (LL-37) for the treatment of chronic wounds.

Financial information

Net sales and result for the second quarter 2022

The company has no revenues from products sales.

The company's costs for Commodities and supplies are mainly related to development costs, such as costs for clinical trials, patents, products for the clinical trials and consultants working with the development of the company's candidate drugs. In the quarter, these costs decreased according to plan to MSEK 3.5 (5.1). The reason to the decrease is that the major costs for PHSU05 were taken by Q1 2022, and that those costs has levelled out.

Other external expenses amounted to MSEK 1.1 (1.6), where the decrease is mainly due to the re-classification of remuneration to the board of directors and lower consultancy costs.

Personnel costs were MSEK 1.5, which is MSEK 0.4 higher compared to the same period last year. The whole increase is due to the re-classification of remuneration to the board of directors from Other external expenses to Personnel costs.

The operating loss for the period amounted to MSEK -6.1, compared to MSEK -7.8 in the same period last year. Net loss for the period amounted to MSEK -6.1 (-7.9), corresponding to earnings per share of SEK -0.10 (-0.22).

Net sales and result for the first half year 2022

The company has no revenues from products sales.

The company's costs for Commodities and supplies are mainly related to development costs, such as costs for clinical trials, patents, products for the clinical trials and consultants working with the development of the company's candidate drugs. In the period, these costs amounted to MSEK 8.7 (8.9), of which MSEK 2.0 of last year's costs were related to the closing of HEAL LL-37 in Q1.

Other external costs amounted to MSEK 2.6 (3.7). The difference is largely explained by the re-classification of remuneration to the board of directors (see above).

Personnel expenses costs were MSEK 3.2, which is MSEK 0.9 higher compared to the same period last year, again explained by the re-classification of remuneration to the board of directors.

The operating loss for the period amounted to MSEK -14.5, compared to MSEK -14.9 in the same period last year. Net loss for the period amounted to MSEK -14.5 (-15.0), corresponding to earnings per share of SEK -0.24 (-0.41).

Cashflow, liquidity and financing during the first half year 2022

The cash flow from operating activities during the period amounted to MSEK -11.2 (-60.9) (where last year's numbers were affected by the fact that the funds from the new issue not yet had been transferred to the company by June 30). A change in the working capital of MSEK +3.3 (-46.0) during primarily the first quarter explains the difference to the net result.

The cash flow from investment activities amounted to MSEK 0.0 (+1.2), where the last year's result is related to the sale of the final shares in Herantis Pharma Oyj.

The cash flow from financing activities was MSEK -0.2 (0.0) during the period, where is related to a paid dept to Karolinska Development as a consequence of the sale of shares in Herantis Pharma Oy.

The company's cash and cash equivalents amounted to MSEK 29.6, compared to MSEK 36.4 by 31 of March, 45.3 by 31 December 2021 and MSEK 13.1 by 30 June 2021. The net proceeds of MSEK 44.7 from last year's new issue were transferred to the company in July 2021.

Group, MSEK	Q2'21	Q3'21	Q4'21	Q1'22	Q2'22
Cash and cash equivalents	13.1	52.1	45.3	36.4	29.6
Working capital	57.3	47.4	41.6	33.0	26.8

Auxiliary information

Risks and uncertainties

Regarding the outbreak of coronavirus and COVID-19, Promore Pharma has taken relevant measures to minimize the impact on the company's business and is following the guidelines from "Folkhälsomyndigheten" (The Public Health Agency of Sweden) and other authorities. Until now, COVID-19 has only had minor effects on Promore Pharma's operations.

The ongoing war in Ukraine and the related sanctions against Russia has so far only had limited effect on Promore Pharma's operations but the company is following the development closely to be able to handle any changed prerequisites. The largest individual effects from the war for Promore Pharma's operations are expected

Further information about risks and uncertainties can be obtained from the company's website, www.promorepharma.com.

Group structure

The Promore Pharma Group comprises, except for the parent company Promore Pharma AB (reg. nr. 556639-6809), also the wholly owned subsidiaries Pergamum AB (reg. nr. 556759-9203) and Pergasus AB (reg. nr. 559349-7695).

Number of shares

Promore Pharma's share is listed on Nasdaq First North (now Nasdaq First North Growth Market) in Stockholm since 6 July 2017 with the ticker PROMO and ISIN code SE0009947740.

The average number of shares, as well as the number of shares at the end of the period, amounted to 60,713,936, while the corresponding number for the same period last year was 36,428,362.

Number of shares	Apr-Jun		Jan-Jun	
	2022	2021	2022	2021
Average number of shares	60,713,936	36,428,362	60,713,936	36,428,362
Number of shares by the end of the period	60,713,936	36,428,362	60,713,936	36,428,362

After the new issue, the main owners Corespring New Technology AB* and PharmaResearch Co. Ltd together own just below 50% of the shares in the company.

Ownership Promore Pharma per 2022-06-30	number	share
Corespring New Technology AB*	22,710,730	37.4%
PharmaResearch Co. Ltd.	7,468,132	12.3%
Nordnet Pensionsförsäkring AB	4,267,278	7.0%
Daniel Johnsson	3,695,444	6.2%
Exceca Allocation & Assoc.	3,332,584	5.5%
Arne Andersson	3,293,254	5.4%
Avanza Pension	1,802,154	2.0%
Other	14,144,360	24.3%
TOTAL	60,713,936	100.0%

*Formerly Midroc New Technology AB

Warrants – external partners

The company announced in March 2021 that, as a consequence of the changed priority for ensereptide, a total of 72,755 warrants (1,091,325 after split) in programs 3-7 issued in 2016 with a dilution effect of approximately 3.0% had been de-registered. After this, 54,599 warrants (818,985 after split) remain related to programs 1, 2 and 8, with a dilution effect of approximately 1.3%. During Q1 2022, another 9 144 warrants (137,160 after split), corresponding to 0.2% of the shares, related to programs 1 & 2 were deregistered. Consequently, 681,825 warrants were outstanding on 30 June 2022, corresponding to a maximum dilution of 1.1%.

Warrants – LTI 2020

It was resolved at the Annual General Meeting in 2020 to adopt a performance-based stock savings program (LTI 2020) for certain employees and contractors in Promore Pharma. A maximum of 1,400,000 Performance Share Rights may be allotted under LTI 2020, corresponding to approximately 3.7 percent of the shares in the company.

In accordance with the Board's proposal, it was resolved that a directed issue of 1,800,000 warrants with the right to subscribe for new shares in the company be used to implement LTI 2020. For those who are offered to join LTI 2020 and previously participated in the company's old bonus program, the old bonus agreements will be terminated without any awards.

Holding of shares in Herantis Pharma Oyj

The company has held shares in the Finnish biotech company Herantis Pharma Oyj. This is a consequence of a passive historic holding in the Finnish company Biocis Oy since the formation of Pergamum AB in 2010. Biocis has since then undergone a number of corporate mergers and ownership restructurings which has resulted in a holding of shares in Herantis Pharma Oyj, a company that executed an IPO in 2015. The last part of the shares were divested in Q1 2021.

Personnel

Promore Pharma has a small and cost-effective organization that is primarily focused on business development, project coordination as well as management of intellectual property and core development documentation. All personnel except the CEO operate on a consultancy basis. Per 30 June 2022, the company consequently had one employee.

Transactions with related parties

The company has not had any transactions with related parties during the period.

Accounting principles

The report has been drawn up in accordance with the Swedish Annual Accounts Act (1995:1554) and the Swedish Accounting Standards Board's (BFNAR) General Recommendation 2012:1: Annual Report and Consolidated Accounts ("K3").

Financial calendar 2022

Q3 report	29 November
Q4 report	28 February 2023
AGM 2022	23 May 2023

Review by auditor

This report has not been reviewed by the Company's auditor.

The Board's declaration

The Board of Directors and the CEO assure that this report provides a fair overview of the company's operations, position, and results.

Solna 30 August 2022

Marianne Dicander Alexandersson
Chairman of the Board

Hans-Peter Ostler

Göran Linder

Kerstin Valinder Strinnholm

Candice (Yujin) Jung

Consolidated income statement

<i>Amounts in SEKk</i>	Apr-Jun		Jan-Jun		Jan-Dec
	2022	2021	2022	2021	2021
Operating income					
Net sales	-	-	-	-	18
Other operating income	24	-1	47	1	417
Operating expenses					
Commodities and supplies	-3,491	-5,055	-8,722	-8,852	-15,312
Other external expenses	-1,104	-1,647	-2,584	-3,714	-7,127
Personnel costs	-1,549	-1,139	-3,189	-2,328	-4,690
Other operating expenses	-13	-5	-53	-8	-
Operating loss (EBIT)	-6,133	-7,847	-14,500	-14,901	-26,694
Financial items					
Net financial items	-4	-24	-7	-65	-78
Profit/loss after financial items	-6,136	-7,872	-14,507	-14,966	-26,772
Profit/loss before tax	-6,136	-7,872	-14,507	-14,966	-26,772
Tax	-	-	-	-	-
Profit/Loss for the period	-6,136	-7,872	-14,507	-14,966	-26,772
EPS	-0.10	-0.22	-0.24	-0.41	-0.56

Consolidated balance sheet

<i>Amounts in SEKk</i>	30 Jun		31 Dec
	2022	2021	2021
ASSETS			
FIXED ASSETS			
Financial fixed assets	1	1	1
Total fixed assets	1	1	1
CURRENT ASSETS			
Current receivables	-	49,985	328
Other receivables	1,622	151	1,555
Cash and cash equivalents	29,635	13,086	45,317
Total current assets	31,257	63,222	47,200
TOTAL ASSETS	31,258	63,223	47,201
EQUITY AND LIABILITIES			
EQUITY			
Share capital	2,429	2,429	2,429
Other equity including the result for the period	23,671	53,965	38,178
Total equity	26,100	56,394	40,607
LONG-TERM LIABILITIES			
Liabilities to credit institutions	714	714	714
Other liabilities	-	237	237
Total long-term liabilities	714	951	951
CURRENT LIABILITIES			
Accounts payable	2,874	1,807	4,002
Deferred taxes	218	140	146
Other current liabilities	1,353	3,932	1,495
Total current liabilities	4,445	5,879	5,643
TOTAL EQUITY AND LIABILITIES	31,258	63,223	47,201

Consolidated cash flow analysis

<i>Amounts in SEKk</i>	Apr-Jun		Jan-Jun		Jan-Dec
	2022	2021	2022	2021	2021
OPERATING ACTIVITIES					
Operating profit	-6,133	-7,847	-14,500	-14,901	-26,694
Adjustments for items not included in cash flow	-21	-24	-7	-27	-190
Tax paid	-	-	-	-	-
Cash flow from operating activities before changes in working capital	-6,153	-7,872	-14,507	-14,928	-26,884
Increase/decrease other current receivables	563	-48,911	260	-49,236	-982
Increase/decrease other current liabilities	-1,220	2,700	-1,198	3,271	3,035
Cash flow from operating activities	-6,810	-54,082	-15,445	-60,893	-24,831
INVESTING ACTIVITIES					
Sale of financial fixed assets	-	-	-	1,159	1,159
Cash flow from investing activities	-	-	-	1,159	1,159
FINANCING ACTIVITIES					
New share issue	-	48,571	-	48,571	44,740
Repaid loans	-	-	-237	-	-
Cash flow from financing activities	-	48,571	-237	48,571	44,740
Cash flow for the period	-6,810	-5,511	-15,682	-11,163	21,068
Cash and cash equiv. at the beginning of the period	36,445	18,597	45,317	24,249	24,249
Exchange rate difference cash and cash equivalents	-	-	-	-	-
Cash and cash equiv. at the end of the period	29,635	13,086	29,635	13,086	45,317

Change in equity for the group

<i>Amounts in SEKk</i>	Share capital	Other paid-in capital	Other equity	Total equity
Amount at the beginning of the period (1 Jan 2022)	2,429	-	38,178	40,607
New share issue	-	-	-	-
Repurchased warrants	-	-	-	-
Profit for the period	-	-	-14,507	-14,507
Amount at the end of the period (30 Jun 2022)	2,429	-	23,671	26,100
Amount at the beginning of the period (1 Jan 2021)	1,457	-	50,736	52,193
New share issue	-	-	-	-
Profit for the period	-	-	-14,966	-14,966
Amount at the end of the period (30 Jun 2021)	1,457	-	35,770	37,227

Parent company income statement

Promore Pharma AB, parent company	Apr-Jun		Jan-Jun		Jan-
	2022	2021	2021	2021	Dec
<i>Amounts in SEKk</i>					2021
OPERATING INCOME					
Net sales	-	-	-	-	18
Other operating income	15	-3	35	-4	412
OPERATING EXPENSES					
Commodities and supplies	-3,431	-4,984	-8,611	-8,715	-15,140
Other external expenses	-1,092	-1,646	-2,553	-3,696	-7,022
Personnel costs	-1,548	-1,139	-3,189	-2,328	-4,689
Depreciation and amortization of tangible assets	-	-	-	-	-
Total operating expenses	-13	-1	-53	-5	-16
Operating profit/loss (EBIT)	-6,069	-7,774	-14,371	-14,748	-26,437
FINANCIAL ITEMS					
Net financial items	-	-	-	-150	-150
Profit/Loss after financial items	-6,069	-7,774	-14,371	-14,898	-26,587
Pre-tax profit	-6,069	-7,774	-14,371	-14,898	-26,587
Tax	-	-	-	-	-
Net profit/loss for the period	-6,069	-7,774	-14,371	-14,898	-26,587

Parent company balance sheet

Promore Pharma AB, parent company	30 Jun		31 Dec
Amounts in SEKk	2022	2021	2021
NON-CURRENT ASSETS			
Share in other long-term securities holdings	10,423	10,398	10,398
Total fixed assets	10,423	10,398	10,398
CURRENT ASSETS			
Accounts receivables	-	-	328
Receivables from group companies	5,305	4,805	4,805
Current tax assets	222	157	144
Other current receivables	794	49,887	713
Prepaid expenses and accrued revenue	457	101	521
Cash and bank balances	23,299	6,857	39,330
Total current assets	30,077	61,806	45,839
TOTAL ASSETS	40,500	72,204	56,238
EQUITY			
Restricted equity			
Share capital	2,429	1,457	2,429
Reserve fund	380	380	380
Total restricted equity	2,809	1,837	2,809
Unrestricted equity			
Share premium reserve	220,462	176,693	220,462
Loss brought forward	-187,239	-133,194	-146,301
Profit/Loss for the period	-	-27,834	-26,567
Total unrestricted equity	33,224	15,665	47,595
Total equity	36,032	17,502	50,404
LONG-TERM LIABILITIES			
Other liabilities	-	237	237
Total long-term liabilities	-	237	237
CURRENT LIABILITIES			
Accounts payables	2,874	1,802	3,934
Current tax liabilities	429	832	347
Accrued expenses and deferred income	1,165	3,260	1,316
Total current liabilities	4,468	5,894	5,597
TOTAL EQUITY AND LIABILITIES	40,500	23,633	56,238

Parent company cash flow analysis

Promore Pharma AB, parent company <i>Amounts in SEKk</i>	Apr-Jun		Jan-Jun		Jan-Dec
	2022	2021	2022	2021	2021
Operating activities					
Operating loss	-6,069	-7,774	-14,371	-14,748	-26,437
Adjustments for non cash flow items	-25	1	-25	1	-149
Tax paid	-	-	-	-	-
Cash flow from operating activities before changes in working capital	-6,094	-7,773	-14,396	-14,747	-26,586
Change in accounts receivables	7	-48,892	-268	-49,257	-818
Change in accounts payable	-1,210	2,675	-1,130	3,276	2,980
Cash flow from operating activities	-7,297	-53,990	-15,793	-60,727	-24,424
FINANCING ACTIVITIES					
New share issue	-	48,571	-	48,571	44,740
Repaid loans	-	-	-237	-	-
Cash flow from financing activities	-	48,571	-237	48,571	44,740
Cash flow for the period	-7,297	-5,419	-16,030	-12,156	20,316
Cash and bank balances in the beginning of the period	30,596	12,276	19,014	19,014	19,014
Exchange rate difference cash and cash equivalents	-	-	-	-	-
Cash and bank balances at year end	23,299	6,857	2,983	6,857	39,330

For additional information, please contact

Jonas Ekblom, CEO

Phone: [+46] 736 777 540

Email: jonas.ekblom@promorepharma.com

Erik Magnusson, CFO

Phone: [+46] 708 56 52 45

Email: erik.magnusson@promorepharma.com

Street address: Fogdevreten 2, 171 65 Solna, Sweden

Website: www.promorepharma.com

Corporate registration number: 556639-6809

Promore Pharma's Certified Adviser is Erik Penser Bank AB .

Phone: [+46] 8 463 83 00

Email: certifiedadviser@penser.se