

MEDIVIR AB – INTERIM REPORT JANUARY – JUNE 2024

“Our study shows a superior profile with the combination of fostrox and Lenvima in advanced liver cancer”

April – June

Financial summary for the quarter

- Net turnover amounted to SEK 1.1 (2.0) million.
- The loss before interest, tax, depreciation and amortization (EBITDA) amounted to SEK -36.7 (-26.3) million. Basic and diluted earnings per share amounted to SEK -0.32 (-0.47).
- Cash flow from operating activities amounted to SEK -26.3 (-17.9) million.
- Cash and cash equivalents at the end of the period amounted to SEK 126.7 (82.8) million.

Significant events during the quarter

- In April it was announced that Medivir's partner Vetbiolix, a veterinary biotechnology company based in France, reported positive results from a proof-of-concept clinical trial in canine periodontitis with its drug candidate VBX-1000, formerly known as MIV-701.
- In April it was announced that Medivir completed a so-called Type C meeting with the FDA and that the company's preparations for the planned phase 2b study in the fostrox program are progressing according to plan, with a few adjustments in study design that have limited impact on timeline and study size.
- In April, MIV-711 was granted Rare Pediatric Disease Designation (RPDD) as well as Orphan Drug Designation (ODD) from the FDA for the treatment of Legg-Calvé-Perthes Disease (LCPD), an unusual hip disease that affects children between the ages 2 and 12.
- The AGM in May re-elected board members Uli Hacksell, Lennart Hansson, Bengt Westermark and Yilmaz Mahshid, and elected Angelica Loskog and

Anna Törner as new board members. Uli Hacksell was re-elected as Chairman of the Board.

- In June it was announced that Medivir has selected a global CRO partner for the planned phase 2b study evaluating fostrox+ Lenvima® compared to Lenvima alone in second-line liver cancer/hepatocellular cancer (HCC).
- On June 26 new positive data showing further improved effect with longer time to progression in Medivir's ongoing phase 1b/2a trial of fostrox + Lenvima in advanced HCC, were presented at the ESMO GI Cancer Congress in Munich.

January – June

Financial summary for the period

- Net turnover amounted to SEK 1.6 (2.4) million.
- The loss before interest, tax, depreciation and amortization (EBITDA) amounted to SEK -63.3 (-45.2) million. Basic and diluted earnings per share amounted to SEK -0.55 (-0.81).
- Cash flow from operating activities amounted to SEK -61.3 (-34.1) million.
- Cash and cash equivalents at the end of the period amounted to SEK 126.7 (82.8) million.

Events after the end of the period

- In July it was announced that Medivir will present updated clinical data from the phase 1b/2a study with fostrox in advanced HCC, at the ESMO Cancer Congress in Barcelona in September.

Medivir in brief

Medivir develops innovative drugs with a focus on cancer where the unmet medical needs are high. The drug candidates are directed toward indication areas where available therapies are limited or missing and there are great opportunities to offer significant improvements to patients. Medivir is focusing on the development of fostroxacitabine bralpamide (fostrox), a drug candidate designed to selectively treat cancer cells in the liver and to minimize side effects. Collaborations and partnerships are important parts of Medivir's business model, and the drug development is conducted either by Medivir or in partnership. Medivir's share (ticker: MVIR) is listed on Nasdaq Stockholm's Small Cap list. www.medivir.com.

CEO's message

Medivir is working decisively to ensure that the combination of fostrox and Lenvima® becomes the first approved treatment alternative after first line standard-of-care in advanced liver cancer. At the ESMO-GI congress in Munich in June, we presented new data from the ongoing phase 1b/2a study. These data indicate that fostrox + Lenvima provides a substantially better effect than previously shown in second-line liver cancer treatment, which generated significant positive attention, both from analysts and clinical experts.

Our ongoing phase 1b/2a study with fostrox + Lenvima continues to show increasingly promising data and we see the opportunity to become the first approved medical treatment in a market worth ~\$2.5 billion annually.

The data presented at the ESMO-GI congress in Munich at the end of June showed good tolerability during longer treatment and that the clinical effect has continued to improve. The Objective Response Rate (ORR) was 24%, higher than the 5–10% ORR seen in other published studies in second-line HCC. The estimated median time to progression at the time of ESMO-GI was 10.8 months, which is substantially better than what's been shown in other studies in second-line HCC. It is tremendously encouraging that the patient in the study who has benefited the longest is still responding to treatment after 2 years.

We now look forward to presenting detailed and mature data highlighting the combination's clinical value in second-line liver cancer at the ESMO Congress in Barcelona in mid-September.

The strong and continuously improving data strengthen our belief in the combination's potential as the first approved treatment option in second-line liver cancer. Preparations for our planned phase 2b study continue based on the feedback we received at our Type C meeting with the FDA.

For the phase 2b study, we have recently chosen a CRO partner with a global presence and a strong track record in oncology studies and in particular HCC studies. We are now initiating the next study phase to identify investigators and hospitals for the study and to complete the study protocol. This will lead us to opening an IND in the US, which is expected to take place in H2 2024.

Regarding the projects out-licensed to collaborators, our partner Vetbiolix, a veterinary biotechnology company based in France, was in April able to report positive results from a clinical Proof-of-Concept study in periodontitis (tooth loss) in dogs with its candidate drug VBX-1000 (MIV-701), which was out-licensed to Vetbiolix in 2019. Vetbiolix is now preparing to evaluate VBX-1000 in a phase 2/3 study in dogs. Tooth loss is an immense problem for dogs and today there is no approved treatment. Vetbiolix estimates that the global market for oral care in pets amounts to approximately SEK 3 billion.

Our project for partnership MIV-711 received Rare Pediatric Disease Designation and Orphan Drug Designation for the treatment of Legg-Calvé-Perthes Disease from the FDA. It creates new opportunities for collaborations and future income.

At the annual general meeting, our board of directors was strengthened with two new members, Angelica Loskog and Anna Törner. Their competence and experience will contribute strongly to Medivir's success, where the clinical development of fostrox is our focus.

We are convinced of the potential of fostrox to become a valuable treatment option that makes a real difference to patients with liver cancer. There is a clear need and an obvious place for fostrox in the treatment landscape. Our goal is to become the first approved treatment alternative in second-line for patients with primary liver cancer. I look forward to keeping you informed of Medivir's continued development.



Jens Lindberg
Chief Executive Officer

Proprietary project



PROPRIETARY PROJECT

Fostroxacitabine bralpamide (fostrox) – for the treatment of liver cancer.

Fostrox is Medivir’s proprietary drug for the treatment of liver cancer. Fostrox is a liver-targeted inhibitor of DNA replication that selectively kills cancer cells in the liver, while the concentration in the rest of the body is lower to minimize possible side effects.

Fostrox’s mechanism of action, inhibition of cancer cells’ DNA replication and induction of DNA damage and cell death, is well proven in cancer therapy. This type of prodrug has successfully proven its ability to deliver the active substance to the liver in anti-viral drugs for hepatitis C. Fostrox has received Orphan Drug Classification (ODD), both in the US and in the EU, for the treatment of HCC.

Primary liver cancer, where the most common form HCC originates from liver cells, is the third leading cause of cancer-related deaths worldwide¹. Although existing treatments for HCC can extend the lives of patients, far from all patients respond to treatment and mortality remains at a high level.

Phase 1a/1b monotherapy study

In the first study with fostrox, phase 1a, safety and tolerability were evaluated at different doses to establish dose levels for the phase 1b study. The results were positive with a good safety and tolerability profile. Thereby the starting dose could be determined for the initial part of the phase 1b/2a study, where fostrox is given in combination with Keytruda® or Lenvima®.

In the monotherapy study, a total of nineteen patients with various types of advanced liver cancer were included and evaluated. These patients had exhausted all possible approved treatments prior to being included in the study.

A positive sign of efficacy was that four out of seven patients with primary liver cancer showed stable disease in the liver. In addition, liver biopsies from patients confirmed delivery of fostrox to the liver, and a selective effect of fostrox on cancer cells in different cancer types.

Ongoing combination study in phase 1b/2a

In December 2021, the phase 1b/2a combination study was initiated with fostrox in combination with two other medicines, either with Lenvima, a tyrosine kinase inhibitor that inhibits blood vessel formation in the tumor, or with Keytruda, an anti-PD-1 checkpoint inhibitor that stimulates the immune system, to patients with HCC for whom current first-line treatment had shown to be ineffective or intolerable. The aim of the study is to evaluate safety, tolerability and to get an indication of the efficacy of fostrox in each combination. The study was initiated at 15 clinics in the UK, Spain and South Korea and is still ongoing. Interest in participating in the study has been great. The dose escalation part (phase 1b) for the combination with Lenvima was completed in February 2023. The preliminary results were positive with a good safety and tolerability profile with no dose-limiting toxicity observed. The recommended phase 2 dose could thereby be determined for the first combination arm, and shortly thereafter the expansion part (phase 2a) for the first combination arm was started. The expansion part of the study is designed for an initial evaluation of safety and efficacy.

In March 2023, the first patient in the phase 2a study was dosed with fostrox in combination with Lenvima and in August the last patient in the phase 2a study was included in this combination. Data reviewed by investigators and local radiologists showed promising tumor control and good tolerability.

The dose escalation part (phase 1b) for the combination with Keytruda was completed in June 2023, establishing a safe dose for treatment with fostrox in combination with Keytruda. However, Medivir is focusing on the combination of fostrox and Lenvima in the expansion part of the ongoing phase 2a study and intends to explore the possibility of fostrox in triple combination with immunotherapy in the earlier treatment-line.

In January 2024, Medivir presented clinical data at the ASCO Gastrointestinal Cancers Symposium in San Francisco and an update of the data was given in June at the European Society for Medical Oncology (ESMO) Gastrointestinal (GI) Cancers Congress in Munich.

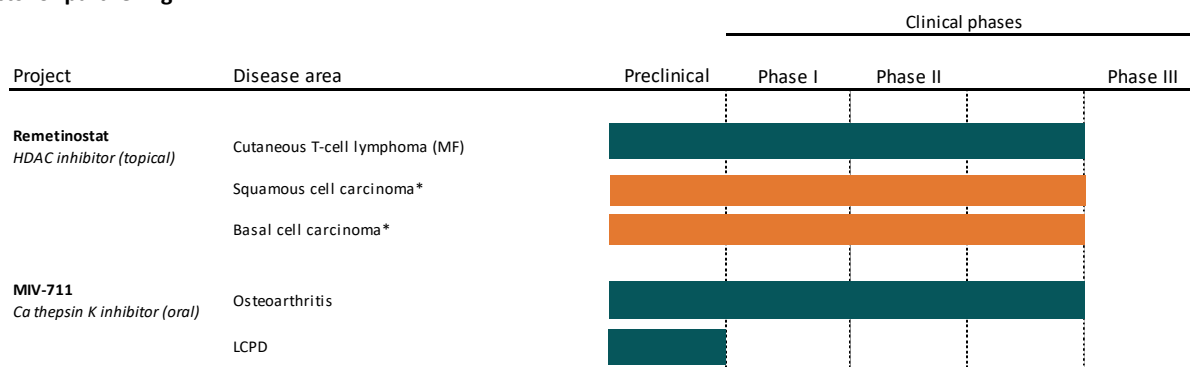
Efficacy data have continuously improved, and data presented at ESMO GI show an ORR of 24%, with disease control (DCR) of 81%, where the estimated median time to progression (TTP) was 10.8 months when approximately 25% of patients remained on

treatment. The patient who has benefited the longest is still responding to treatment after 2 years. Biopsies confirm selective DNA damage to tumor cells without impact on normal liver function based on measured liver enzyme values (ALT/AST) along with stable ALBI values (which measure liver function) over time. The update also showed continued good tolerability without any new unexpected side effects.

Taken together, these data provide strong support for accelerating the fostrox development program in second-line HCC in 2024 with a focus on initiating a randomized phase 2b trial in second-line HCC patients comparing the combination of fostrox and Lenvima to Lenvima monotherapy.

- 1) <https://gco.iarc.fr/today/data/factsheets/cancers/11-Liver-fact-sheet.pdf>

Projects for partnering



* Conducted by Stanford University, USA

█ Investigator sponsored study

PROJECTS FOR PARTNERING

Medivir has two projects for licensing/partnerships:

Remetinostat – *histone deacetylase inhibitor for the treatment of different types of cancers in the skin.*

MIV-711 – *cathepsin K inhibitor with the potential to become the first disease-modifying treatment for, among other things, osteoarthritis, but also for some rare, bone-related, diseases in children.*

Currently Medivir does not conduct any active clinical development for these projects, but instead evaluates the possibilities of concluding a license or collaboration agreement for the continued development of each project.

Remetinostat for cancer in the skin

Three phase II studies with remetinostat have been conducted, one in cutaneous T-cell lymphoma (MF) and two investigator-initiated studies in basal cell carcinoma and cutaneous squamous cell carcinoma. Remetinostat has shown positive clinical efficacy and acceptable tolerability without systemic side effects in these three types of cancer.

Project descriptions

Full descriptions of all of Medivir’s development projects, including their current status and ongoing studies, can be found on the Medivir website: <http://www.medivir.com/our-projects>

MIV-711

Medivir has conducted a phase II study with positive effects on both bone and cartilage in joints in osteoarthritis patients after only six months of treatment with MIV-711.

In February 2022, a subgroup analysis of Medivir's phase II study with MIV-711 for osteoarthritis was published, showing a significant reduction in osteoarthritis-related pain.

In April 2024, MIV-711 was granted Rare Pediatric Disease Designation (RPDD) and Orphan Drug Designation (ODD) from the FDA for the treatment of Legg-Calvé-Perthes disease (LCPD), a rare hip disorder that affects children ages 2- 12 years. A disease for which there are currently no effective treatment options.

Outlicensed projects

Project	Disease area	Partner	Preclinical development	Phase I	Phase II	Phase III	Market
Xerclear	Labial herpes	GSK					
Birinapant (9427) + IGM-8444 <i>SMAC mimetic (intravenous)</i>	Solid tumors	IGM Biosciences					
USP-7	Cancer	Ubiquigent Limited					
MBLI/MET-X	Infection	INFEX Therapeutics					
MIV-701/VBX-1000	Periodontal (veterinary)	Vetbiolix					

Ongoing study

OUTLICENSED PROJECTS

Xerclear® - In 2009, Xerclear® (Zovido®) was approved for the treatment of labial herpes. The marketing rights to Xerclear® in the USA, Canada and Mexico were divested in 2010, while the corresponding rights in Europe and the rest of the world have been out-licensed to GlaxoSmithKline, with the exception of China, where Medivir has out-licensed the rights to Shijiazhuang Yuanmai Biotechnology Co Ltd. (SYB), and Israel and South America where Medivir has retained the rights.

Medivir receives royalties on Xerclear® (Zovido®) sales from GlaxoSmithKline. In addition, Medivir would receive milestones when Zovido® is approved as an over the counter product in new markets.

After marketing approval and production in China, Medivir will receive a fixed royalty from SYB for each unit sold and the agreement guarantees a minimum sale during the first three years on the market amounting to single-digit million SEK.

Birinapant – for the treatment of solid tumors.

In January 2021, Medivir entered into a licensing agreement with IGM Biosciences regarding the global and exclusive rights to develop birinapant.

Medivir received a payment of USD 1 million upon signing the agreement, which was followed by an additional USD 1.5 million when IGM in November 2021 initiated a clinical phase I study in solid cancers with birinapant in combination with its DR5-agonist antibody IGM-8444 now called aplitabart.

During the fourth quarter, the fifth dose-escalation cohort was completed, and no dose-limiting toxicity has been observed to date. In December, IGM communicated a strategic pipeline prioritization in order to save costs, and it is currently unclear how it affects the future development of aplitabart in combination with birinapant. The terms of the agreement entitles Medivir to milestone payments up to a total of approximately USD 350 million, given that birinapant is successfully developed and approved, and tiered royalties up to "mid-teens" on net sales. A portion of all revenue is shared with Tetralogic

Pharmaceuticals Corporation, but the main part goes to Medivir.

USP-1/TNG348

In the first quarter of 2020 Medivir entered a licensing agreement with the US-based company Tango Therapeutics for Medivir's preclinical research program USP-1. In September, Tango received IND approval from the FDA and in January 2024, Tango Therapeutics announced that the company dosed the first patient in a phase 1/2 study with TNG348, a USP-1 inhibitor from Medivir's preclinical research program. In May, Tango announced that the phase 1/2 study of TNG348 is being terminated due to toxicity observed in the first study cohorts. Tango maintains the preclinical USP-1 program and is evaluating potential options moving forward.

MIV-701

Medivir's selective cathepsin-K inhibitor MIV-701 was discovered to have properties suitable for use in animals and was out-licensed to France's Vetbiolix in 2019. In April 2024, Vetbiolix reported positive results from a Proof-of-Concept clinical study in canine periodontitis with its drug candidate VBX-1000 (MIV-701). A disease for which there are currently no approved treatments and where the global market for oral care in pets is estimated at SEK 3 billion annually. The company is now preparing a phase 2/3 study to further strengthen the documentation of the effects of VBX-1000. The agreement entitles Medivir to minor development and regulatory milestone payments with value upside potential coming from future royalty payments on net sales and/or share of payments that Vetbiolix receives in the event of a future partnering agreement with VBX-1000.

Preclinical projects

USP-7

In February 2021 a licensing agreement with Ubiquigent was signed for the preclinical research program USP-7. The agreement grants Ubiquigent an exclusive global

license to develop and commercialize all of the program's related substances in all therapeutic indications in exchange for agreed revenue sharing with Medivir upon successful development or commercialization.

MBLI/MET-X

Medivir's Metallo Beta Lactamase (MBLI) program aimed at addressing the threat of resistant bacteria was out-licensed in 2017 to the AMR Centre (today INFEX Therapeutics) in England.

In 2022, INFEX presented additional preclinical data, received patent approval for the substance in the United States. In January 2023, MET-X received QIDP-designation (Qualified Infectious Disease Product) from the FDA and in August patent approval was obtained in Europe. INFEX has communicated its intention to initiate a phase I program for MET-X in 2024. Medivir is entitled to a share of potential future revenue.

In the event of any discrepancies between the Swedish and the English Interim Report, the former should have precedence.

Financial overview, April – June 2024

Summary of the Group's figures

(SEK m)

	Q2		Q1 - Q2		Full Year
	2024	2023	2024	2023	2023
Net turnover	1.1	2.0	1.6	2.4	7.6
Operating profit before depreciation and amortization (EBITDA)	-36.7	-26.3	-63.3	-45.2	-88.7
Operating profit (EBIT)	-37.3	-27.0	-64.7	-46.6	-91.4
Profit/loss before tax	-36.0	-26.6	-62.0	-45.5	-89.3
Basic earnings per share, SEK	-0.32	-0.47	-0.55	-0.81	-1.48
Diluted earnings per share, SEK	-0.32	-0.47	-0.55	-0.81	-1.48
Net worth per share, SEK	1.54	2.61	1.54	2.61	2.07
Return on equity, %	-74.3	-66.1	-63.0	-53.4	-43.5
Cash flow from operating activities	-26.3	-17.9	-61.3	-34.1	-59.7
Cash and cash equivalents at period end	126.7	82.8	126.7	82.8	169.5

Revenues

Net turnover for the period from April – June was SEK 1.1 million (2.0 m), corresponding to a decrease of SEK 0.9 million. The decrease refers to lower royalty income.

Operating expenses

Other external costs totaled SEK -30.3 million (-21.2 m), corresponding to an increase of SEK 9.1 million which relates to higher cost for clinical studies.

Personnel costs amounted to SEK -7.6 million (-7.4 m), corresponding to an increase of MSEK 0.2 which relates foremost to costs for the share savings program that was implemented during Q2, 2024. The total overheads amounted to SEK -38.6 million (-29.5 m), an increase of 9.0 million.

Operating profit/loss

The operating loss totaled SEK -37.3 million (-27.0 m), SEK 10.4 million lower result compared to previous year. The lower result mainly relates to higher clinical costs.

Cash flow, investments, and financial position

Liquid assets, including short-term investments amounted to SEK 126.7 million (82.8 m) at the end of the period, corresponding to an increase of SEK 43.8 million. The opening balance 2024 was SEK 169.5 million (117.4 m).

Cash flow from operating activities totaled SEK -26.3 million (-17.9 m), with changes in working capital accounting for SEK 9.1 million (8.5 m) of this total.

The period's investments in tangible and intangible fixed assets totaled SEK 0.0 million (-0.3 m).

Cash flow from financing activities totaled SEK -0.4 million (0.2 m).

Financial overview, January – June 2024

Revenues

Net turnover for the period from January – June was SEK 1.6 million (2.4 m) corresponding to a decrease of SEK 0.7 million. The decrease refers to lower royalty income in Q2 2024

Operating expenses

Other external costs totaled SEK -51.0 million (-34.3 m), corresponding to an increase of SEK 16.7 million which relates to higher cost for clinical studies.

Personnel costs amounted to SEK -14.1 million (-13.6 m), corresponding to an increase of MSEK 0.5 which relates foremost to the cost of the share savings program that was implemented during Q2, 2024. The total overheads amounted to SEK -66.6 million (-49.9 m), an increase of 16.7 million.

Operating profit/loss

The operating loss totaled SEK -64.7 million (-46.6 m), SEK 18.2 million lower result compared to previous year. The lower result mainly relates to higher clinical costs.

Cash flow, investments, and financial position

Liquid assets, including short-term investments amounted to SEK 126.7 million (82.8 m) at the end of the period, corresponding to an increase of SEK 43.8 million. The opening balance 2024 was SEK 169.5 million (117.4 m).

Cash flow from operating activities totaled SEK -61.3 million (-34.1 m), with changes in working capital accounting for SEK -0.3 million (10.8 m) of this total.

The period's investments in tangible and intangible fixed assets totaled SEK 0.0 million (-0.3 m). Cash flow from financing activities totaled SEK 18.5 million (-0.3 m)

Other disclosures, January – June 2024

Employees

Medivir had 10 (10) employees (FTEs) at the period end, 60% (60%) of whom were women.

Share and related incentive plans

In January 2024, the company carried out a directed issue of 7,547,170 ordinary shares to Hallberg Management AB, resulting in Medivir receiving approximately 20 million SEK before transactions costs. At the annual general meeting on May 7, 2024, it was decided to adopt a new long-term incentive program in the form of a share matching program (LTIP 2024). Against the background of LTIP 2024, a new issue of 1,700,000 C shares has taken place during the second quarter and of these 114,587 have been converted into ordinary shares. The converted shares as well as 11,413 existing ordinary shares held by the company, a total of 126,000, have been transferred to the participants in LTIP 2024 as Investment Shares.

Number of shares	Ordinary		Total Shares
	Shares	C shares	
No. of shares 1/1-2024	104 506 048	864 750	105 370 798
Direct issue shares	7 547 170	-	7 547 170
LTIP 2024	114 587	1 585 413	1 700 000
No. of shares 30/6-2024	112 167 805	2 450 163	114 617 968

Medivir's holdings amount to 2,450,163 own C shares in the company.

Warrants - At the beginning of the period, there were 1,060,000 outstanding warrants in the ongoing incentive programs. There was no change during the

period. The total number of outstanding warrants at the end of the period amounted to 1,060,000.

In May 2021, the Board of Directors proposed, and the AGM approved a new long-term incentive program. During the second quarter 2021, Medivir employees bought 230 000 warrants at a market value of 1.00 each with an exercise price of SEK 13.79 per share. In the fourth quarter 2021, Medivir employees bought an additional 305,000 warrants of which incoming CEO bought 240,000. These warrants were issued at a market value of SEK 1.71 each with an exercise price of SEK 13.79 per share. The warrants may be exercised to subscribe for new ordinary shares during the period from 1 December 2024 up to and including 15 December 2024. The valuation calculation for 2021 was based on the following figures: term, 3.60 years; strike price, SEK 13.79; VWAP, SEK 7.88; risk-free interest rate, 0.4 percent; volatility, 41 percent. After recalculation caused by the rights issue in quarter 4 2023, each such warrant entitles the holder to subscribe for 1.06 new ordinary shares in the company at a subscription price of SEK 12.98.

In May 2022, the Board of Directors proposed and the AGM approved a new long-term incentive program with similar terms to the program in 2021. In the fourth quarter 2022, Medivir employees bought 525,000 warrants of which CEO bought 250,000. These warrants were issued at a market value of SEK 0.77 each with an exercise price of SEK 14.13 per share. The warrants may be exercised to subscribe for new ordinary shares during the period from 1 December 2025 up to and

including 15 December 2025. The valuation calculation for 2022 was based on the following figures: term, 3.12 years; strike price, SEK 14.13; VWAP, SEK 8.07; risk-free interest rate, 2.14 percent; volatility, 36 percent. After recalculation caused by the rights issue in quarter 4 2023, each such warrant entitles the holder to subscribe for 1.06 new ordinary shares in the company at a subscription price of SEK 13.30.

Share savings program – At the beginning of the period, there were 105,750 investment shares in ongoing share savings programs. In Q2 2024, a new share savings program was implemented, and participants acquired a total of 126,000 investment shares. Total outstanding investment shares at the end of the period amounted to 231,750.

In May 2023, the board and the annual general meeting approved a new long-term incentive program in the form of a share matching program. For each investment share, participants have the opportunity, provided that certain conditions are met, to receive one (1) ordinary share free of charge within the framework of LTIP 2023 ("matching shares") and in addition, provided that certain performance conditions are met, a maximum of five (5) additional ordinary shares ("performance shares") free of charge according to the terms of the program. As of December 30, Medivir's employees have purchased 105,750 investment shares at a price of SEK 7.34. The earned period is until the publication of the interim report for January-March 2026. After recalculation due to rights issue during quarter 4 2023, each investment share entitles to 1.22 ordinary shares.

In May 2024, the board and the annual general meeting approved a new long-term incentive program in the form of a share matching program. For each investment share, participants have the opportunity, provided that certain conditions are met, to receive one (1) ordinary share free of charge within the framework of LTIP 2024 ("matching shares") and in addition, provided that certain performance conditions are met, a maximum of five (5) additional ordinary shares ("performance shares") free of charge according to the terms of the program. As of June 30, 2024, Medivir's employees have purchased 126,000 investment shares at a price of SEK 2.94. The earned period is up to and including publication of the interim report for January-March 2027.

Currency exposure

In accordance with Medivir's financial policy, a large part of the euro flow is currency hedged. For other currencies, the group has not used currency hedging, which means that income and costs have been affected by fluctuations in foreign exchange rates. All trading in foreign currency has taken place at the best exchange

rate that could be obtained at each time of exchange. Many of Medivir's contracts involve payment in EUR, CHF, USD and GBP, which means that accounts payable and accounts receivable have a currency exposure.

The Parent Company in brief

Medivir AB (publ.), corporate ID no. 556238-4361, is the Parent Company of the Group. Its operations consist of pharmaceutical development, administrative and company management functions. All operations in the group are carried out in the parent company.

The Parent Company's total turnover amounted to SEK 1.6 million (2.4 m).

Combined operating expenses totaled SEK -66.9 million (-50.1 m), an increase with SEK 16.8 million.

The operating loss was SEK -65.0 million (-46.8 m), corresponding to a decrease in the result of SEK 18.2 million.

Net financial items totaled SEK 3.1 million (1.5 m), corresponding to an increase of SEK 1.5 million.

The tax for the period totaled SEK 0.0 million (0.0 m).

The net loss for the period was SEK -62.0 million (-45.3 m), corresponding to a decrease of SEK 16.7 million. The lower result mainly relates to higher clinical costs.

Liquid assets, including short-term investments with a maximum term of three months, amounted to SEK 126.6 million (82.2 m).

Transactions with related parties

During the period, no transactions with related parties were carried out except for board fees.

Significant risks and uncertainty factors

The process of pharmaceutical research and development, all the way up to regulatory market approval, is both high-risk and capital-intensive. The majority of projects initiated will never achieve market authorization. If competing pharmaceuticals take market shares, or competing research projects achieve better efficacy and reach the market more quickly, the future value of Medivir's product and project portfolio may be lower than expected. Medivir's success in developing medicines, to enter into partnerships and to secure funding for its operations, are decisive in terms of the company's future.

In addition to industry-specific risk factors, there is an added uncertainty in our surrounding world, both due to Russia's invasion war in Ukraine, unrest in the Middle East, and the conflict surrounding Taiwan. Although central banks currently appear to have inflation under control, there is still a risk that political and geopolitical conflicts may negatively impact the economy and inflation.

A more detailed description of the exposure to risk, and of the ways in which Medivir manages it, is provided in the 2023 Annual Report, see pages 23-25 and 32 and in Note 7 on pages 47-49. The Annual

Report is available on the company's website:
www.medivir.com.

Outlook

Medivir's future investments will mainly be in clinical pharmaceutical projects within oncology.

It is the assessment of the Board and management that existing cash and cash equivalents are sufficient to cover the company's needs to complete the ongoing combination arm in phase 2a. The existing cash and cash equivalents are estimated to meet the company's

liquidity needs until Q1 2025 according to current plans and assumptions. The company is evaluating different financing options and the board and management make the assessment that the group has good conditions to carry out a financing within 12 months to ensure the group's continued operation and continue the development of the fostrox program.

Attestation

The Board of Directors and the President & CEO hereby affirm that the Interim Report constitutes a faithful representation of the company's and the Group's operations, position and profit/loss, and that it describes the significant risks and uncertainty factors faced by the company and the companies that make up the Group.

Huddinge, August 22, 2024

Uli Hacksell
Chairman of the Board

Lennart Hansson
Member of the Board

Angelica Loskog
Member of the Board

Yilmaz Mahshid
Member of the Board

Anna Törner
Member of the Board

Bengt Westermark
Member of the Board

Jens Lindberg
Chief Executive Officer

This report has not been subject to auditors' review.

The information was submitted for publication at 08.30 CET on August 22, 2024.

For further information, please contact

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Conference call for investors, analysts and the media

The Interim Report January - June 2024 will be presented by Medivir's CEO, Jens Lindberg.

Time: Thursday, August 22, 2024, at 14.00 (CET).

To access the webcast and find information about the teleconference, please click [HERE!](#)

The conference call will also be streamed via a link on the website: www.medivir.com/investors/calendar.

The presentation will be available on Medivir's website after completion of the conference.

Financial calendar:

Interim Report (January – September 2024)

November 6, 2024

Year End Report (January – December 2024)

February 18, 2025

Interim Report (January – March 2025)

April 29, 2025

Interim Report (January – June 2025)

August 21, 2025

Notes

Accounting principles

Medivir prepares its Consolidated Accounts in accordance with IFRS, International Financial Reporting Standards, as endorsed by the EU. In addition to the stated IFRS, the Group also applies the Swedish Financial Reporting Board's recommendation, RFR 1 Supplementary Accounting Rules for Groups, and applicable statements from the Swedish Financial Reporting Board. The Group utilizes the acquisition value for Balance Sheet item valuation, unless otherwise indicated.

The interim report has been prepared in accordance with IAS 34. IFRS are under constant development, and new standards and interpretations are published on an ongoing basis. No new standards that are expected to affect the period's earnings and financial position have entered into force. See pages 39-44 of the 2023 Annual Report for a full presentation of the accounting principles applied by the Group. There have been no changes in the accounting principles since the annual report for 2022 was submitted. Rounding off may mean that certain tables do not add up.

Consolidated Income Statement, summary

(SEK m)	Q2		Q1 - Q2		Full year
	2024	2023	2024	2023	2023
Net turnover	1.1	2.0	1.6	2.4	7.6
Other operating income	0.1	0.6	0.3	1.0	1.4
Total income	1.2	2.6	1.9	3.3	9.0
Other external expenses	-30.3	-21.2	-51.0	-34.3	-68.9
Personnel costs	-7.6	-7.4	-14.1	-13.6	-27.4
Depreciations and write-downs	-0.7	-0.7	-1.4	-1.4	-2.7
Other operating expenses	0.0	-0.2	-0.1	-0.6	-1.4
Operating profit/loss	-37.3	-27.0	-64.7	-46.6	-91.4
Net financial items	1.4	0.4	2.7	1.1	2.1
Profit/loss after financial items	-36.0	-26.6	-62.0	-45.5	-89.3
Tax	-	-	-	-	-
Net profit/loss for the period	-36.0	-26.6	-62.0	-45.5	-89.3
Net profit/loss for the period attributable to:					
Parent Company shareholders	-36.0	-26.6	-62.0	-45.5	-89.3
Earnings per share, calculated from the net profit/loss attributable to Parent Company shareholders during the period					
Earnings per share (SEK per share)					
- Total operations, basic earnings	-0.32	-0.47	-0.55	-0.81	-1.48
- Total operations, diluted earnings	-0.32	-0.47	-0.55	-0.81	-1.48
Average number of shares, '000	114 051	56 383	113 485	56 059	60 438
Average number of shares after dilution '000	114 051	56 383	113 485	56 059	60 438
Number of shares at period end, '000	114 618	56 706	114 618	56 706	105 371

Consolidated Statement of Comprehensive Income

(SEK m)	Q2		Q1 - Q2		Full year
	2024	2023	2024	2023	2023
Net profit/loss for the period	-36.0	-26.6	-62.0	-45.5	-89.3
Other comprehensive income					
Exchange rate differences	-	0.1	-	0.1	-0.1
Total other comprehensive income	-	0.1	-	0.1	-0.1
Total comprehensive income for the period	-36.0	-26.5	-62.0	-45.4	-89.4

Consolidated Balance Sheet, summary (SEK m)

	30-jun 2024	30-jun 2023	31-dec 2023
Assets			
Intangible fixed assets	96.3	96.3	96.3
Tangible fixed assets	11.0	13.7	12.4
Current receivables	5.2	7.0	9.7
Short-term investments	117.0	76.5	144.0
Cash and cash equivalents	9.6	6.3	25.6
Total assets	239.2	199.8	287.9
Shareholders' equity and liabilities			
Shareholders' equity	176.0	147.9	217.9
Long-term liabilities	10.1	12.5	11.3
Current liabilities	53.1	39.4	58.7
Total shareholders' equity and liabilities	239.2	199.8	287.9

Consolidated Statement of Changes in Equity (SEK m)

	Share capital	Other paid-in capital	Exchange rate difference	Accum. loss	Total equity
Opening balance, 1 January 2023	27.9	805.3	-3.2	-637.2	192.8
Total comprehensive income for the period	-	-	-	-45.4	-45.4
Stock dividend issue	0.5	0.3	-	-	0.8
Transaction costs	-	-	-	-0.3	-0.3
Closing balance, 30 June 2023	28.4	805.6	-3.2	-682.9	147.9
Opening balance, 1 January 2023	27.9	805.3	-3.2	-637.2	192.8
Total comprehensive income for the period	-	-	-0.1	-89.3	-89.4
Stock dividend issue	24.3	104.6	-	-	129.0
Share savings program	0.5	0.3	-	0.5	1.2
Transaction costs	-	-	-	-15.7	-15.7
Closing balance, 31 December 2023	52.7	910.3	-3.3	-741.7	217.9
Opening balance, 1 January 2024	52.7	910.3	-3.3	-741.7	217.9
Total comprehensive income for the period	-	-	-	-62.0	-62.0
Share issue	3.8	16.2	-	-	20.0
Share savings program	0.9	-0.5	-	0.4	0.8
Transaction costs	-	-	-	-0.7	-0.7
Closing balance, 30 June 2024	57.3	926.0	-3.3	-804.0	176.0

Consolidated Cash Flow Statement, summary (SEK m)

	Q2		Q1 - Q2		Full Year
	2024	2023	2024	2023	2023
Cash flow from operating activities before changes in working capital	-35.4	-26.4	-61.1	-44.8	-86.1
Changes in working capital	9.1	8.5	-0.3	10.8	26.4
Cash flow from operating activities	-26.3	-17.9	-61.3	-34.1	-59.7
Investing activities					
Acquisition/sale of fixed assets	-	-0.3	-	-0.3	-0.3
Cash flow from investing activities	-	-0.3	-	-0.3	-0.3
Financing activities					
Other changes in longterm receivables/liabilities	-0.6	-0.3	-1.2	-0.8	-2.0
Right issue	0.4	0.8	20.4	0.8	129.7
Transaction costs	-0.2	-0.3	-0.7	-0.3	-15.7
Cash flow from financing activities	-0.4	0.2	18.5	-0.3	112.1
Cash flow for the period	-26.8	-18.0	-42.9	-34.6	52.1
Cash and cash equivalents at beginning of period	153.4	100.8	169.5	117.4	117.4
Exchange rate difference, liquid assets	-	0.0	-	0.0	-0.1
Cash and cash equivalents at end of period	126.7	82.8	126.7	82.8	169.5

Parent company income statement, summary

(SEK m)	Q2		Q1 - Q2		Full year
	2024	2023	2024	2023	2023
Net turnover	1.1	2.0	1.6	2.4	7.6
Other operating income	0.1	0.6	0.3	1.0	1.4
Total income	1.2	2.6	1.9	3.3	9.0
Other external expenses	-31.1	-22.0	-52.6	-35.9	-72.0
Personnel costs	-7.6	-7.4	-14.1	-13.6	-27.4
Depreciations and write-downs	0.0	0.0	-0.1	-0.1	-0.1
Other operating expenses	0.0	-0.2	-0.1	-0.6	-1.4
Operating profit/loss	-37.5	-27.1	-65.0	-46.8	-91.9
Profit/loss from participation in Group companies	-	-	-	-	0.5
Net financial items	1.6	0.6	3.1	1.5	3.0
Profit/loss after financial items	-35.9	-26.5	-62.0	-45.3	-88.4
Tax	-	-	-	-	-
Net profit/loss for the period (=comprehensive income)	-35.9	-26.5	-62.0	-45.3	-88.4

Parent company balance sheet, summary

(SEK m)	30-jun	30-jun	31-dec
	2024	2023	2023
Assets			
Intangible fixed assets	96.3	96.3	96.3
Tangible fixed assets	0.1	0.3	0.2
Shares in subsidiaries	0.1	0.1	0.1
Current receivables	6.2	7.6	10.5
Short-term investments	117.0	76.5	144.0
Cash and bank balances	9.6	5.7	25.5
Total assets	229.4	186.6	276.6
Shareholders' equity and liabilities			
Shareholders' equity	176.5	147.5	218.3
Liabilities to Group companies	1.8	1.8	1.8
Current liabilities	51.1	37.2	56.5
Total shareholders' equity and liabilities	229.4	186.6	276.6

Key ratios, share data

	Q2		Q1 - Q2		Full year
	2024	2023	2024	2023	2023
Return on:					
- shareholders' equity, %	-74.3	-66.1	-63.0	-53.4	-43.5
- capital employed, %	-69.3	-60.0	-58.7	-48.6	-40.2
- total capital, %	-56.6	-50.7	-46.8	-41.5	-33.9
Number of shares at beginning of period, '000	112 918	55 736	105 371	55 736	55 736
Number of shares at period end, '000	114 618	56 706	114 618	56 706	105 371
- of which class A shares	112 168	55 841	112 168	55 841	104 506
- of which class B shares	-	-	-	-	-
- of which repurchased B shares	2 450	865	2 450	865	865
Average number of shares, '000	114 051	56 383	113 485	56 059	60 438
Share savings program (investment shares), '000	232	106	232	106	106
Outstanding warrants, '000	1 060	1 587	1 060	1 587	1 060
Share capital at period end, SEK m	57.3	28.4	57.3	28.4	52.7
Shareholders' equity at period end, SEK m	176.0	147.9	176.0	147.9	217.9
Earnings per share, SEK					
- Total operations, basic earnings	-0.32	-0.47	-0.55	-0.81	-1.48
- Total operations, diluted earnings	-0.32	-0.47	-0.55	-0.81	-1.48
Shareholders' equity per share, SEK	1.54	2.61	1.54	2.61	2.07
Net worth per share, SEK	1.54	2.61	1.54	2.61	2.07
Cash flow per share after investments, SEK	-0.23	-0.32	-0.54	-0.61	-0.99
Equity/assets ratio, %	73.6	74.0	73.6	74.0	75.7
EBITDA	-36.7	-26.3	-63.3	-45.2	-88.7
EBIT	-37.3	-27.0	-64.7	-46.6	-91.4

Key ratio definitions

Average number of shares. The unweighted average number of shares during the period.

Basic earnings per share. Profit/loss after tax divided by the average number of shares.

Capital employed. Balance Sheet total less non-interest-bearing liabilities including deferred tax liabilities.

Cash flow per share after investments. Cash flow after investments divided by the average number of shares.

Diluted earnings per share. Profit/loss after tax divided by the average number of shares and outstanding warrants adjusted for any dilution effect.

EBIT (Earnings before interest and taxes). Operating profit/loss after depreciation and amortization.

EBITDA (Earnings before interest, taxes, depreciation and amortization). Operating profit/loss before depreciation and amortization.

Equity/assets ratio. Shareholders' equity in relation to the Balance Sheet total.

Net worth per share. Shareholders' equity plus hidden assets in listed equities divided by the number of shares at the period end.

Operating margin. Operating profit/loss as a percentage of net turnover.

Return on capital employed. Profit/loss after financial items plus interest expenses as a percentage of the average capital employed.

Return on shareholders' equity. Profit/loss after tax as a percentage of the average shareholders' equity.

Return on total assets. Profit/loss after financial items plus interest expenses as a percentage of the average Balance Sheet total.

Shareholders' equity per share. Shareholders' equity divided by the number of shares at the period end.

The above key ratios are deemed to be relevant for the type of operations conducted by Medivir and to contribute to an increased understanding of the financial report.