



KEY FIGURES & FINANCIAL HIGHLIGHTS

TDKK	Q2 2024	Q1-Q2 2024	Q2 2023	Q1-Q2 2023	FY 2023
Income Statement					
Operating loss	-9,359	-19,458	-11,318	-23,292	-45,357
Net finance income/cost	225	418	66	131	654
Loss before tax	-9,133	-19,040	-11,253	-23,160	-44,704
Net loss	-7,164	-14,733	-8,774	-18,062	-39,204
Total comprehensive loss	-7,164	-14,733	-8,774	-18,062	-39,204
Balance Sheet					
Total non-current assets	4,949	4,949	7,154	7,154	897
Total current assets	34,037	34,037	53,032	53,032	33,664
Hereof Cash and Cash equivalents	26,952	26,952	45,709	45,709	26,520
Total Assets	38,987	38,987	60,186	60,186	34,560
Total Equity	30,727	30,727	52,265	52,265	31,122
Cash Flow					
From Operating activities	-3,893	-13,455	-14,280	-31,505	-50,668
From Investing activities	0	88	0	0	288
From Financing activities	13,889	13,799	-196	-391	-705
Net cash flow for the period	9,996	432	-14,476	-31,895	-51,085
Key ratios					
Equity ratio	79%	79%	87%	87%	90%
Earnings per share (EPS)	-0.03	-0.06	-0.22	-0.44	-0.96
Earnings per share (EPS-D)	-0.03	-0.06	-0.22	-0.44	-0.96
Shareholder EQT per share	0.13	0.13	1.28	1.28	0.76
Employees					
Average number of FTE	4	4	9	10	7
Number of FTE end of period	4	4	5	5	4
Shares, Outstanding end of period	231,928,544	231,928,544	40,706,972	40,706,972	40,706,972

HIGHLIGHTS DURING Q2 2024

ON APRIL 19, Scandion Oncology announced the intention, subject to authorizations by the annual general meeting of the Company on 6 May 2024, to carry out a Rights Issue with preferential rights for the Company's existing shareholders. The Rights Issue of potentially SEK 60 million is secured up to SEK 30.6 million.

ON MAY 13, Scandion announced final data from the Phase Ib open-label PANTAX trial which confirms the good safety profile of SCO-101 and shows good signs of efficacy in hard-to-treat pancreatic cancer.

ON JUNE 3, Scandion announced, that the company has entered into an agreement with Vator Securities AB regarding the service as a Certified Adviser. Vator Securities will be appointed Certified Adviser (CA) on September 1, 2024.

ON JUNE 25, Scandion announced the outcome of the rights issue of units. The Rights Issue was subscribed to a total of approximately 50.3 percent. Through the Rights Issue, Scandion will initially receive approximately SEK 30.6 million before issue costs and in the event of exercise of warrants of series TO 2 and TO 3, in November 2024 and April 2025, respectively, the Company will receive additional proceeds.

HIGHLIGHTS AFTER THE END OF THE REPORTING PERIOD

ON JULY 1, Scandion board member Michel Ducreux stepped down due to ESMO scientific society's guidelines prohibiting such board positions. He joined the advisory board.

ON AUGUST 16, Scandion Oncology achieved Maximum Tolerated Dose (MTD) for CORIST part 3. The established MTD for a 4-Days schedule of SCO-101 in combination with FOLFIRI was found to be 250 mg daily SCO-101, 50% irinotecan and 100% Leucovorin and 5-FU.

ON AUGUST 19, Scandion announced that the top priority following the very encouraging part 3 CORIST data is business development and partnering activities. As part of these efforts, Scandion is working together with Back Bay Life Science Advisors LLC, a prominent life sciences investment banking firm, to explore and evaluate actionable strategic and financial alternatives.





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In this document, the following definitions shall apply unless otherwise specified: *"the Company"* or *"Scandion"* refers to **Scandion Oncology A/S**, CVR No. 38613391.

CEO LETTER

POSITIVE CORIST DATA AND AN INCREASED FOCUS ON **BUSINESS DEVELOPMENT**

Scandion continued to advance the company's lead compound SCO-101 as a combination treatment with FOLFIRI chemotherapy in metastatic colorectal cancer (mCRC) during the second quarter.

Scandion is in a unique position to help patients with SCO-101, our innovative drug efflux pump inhibition treatment using biomodulation capabilities to revert drug resistance, which remains a significant problem in cancer treatment and in the development of new medicines as cancer tumors rapidly adapt and resist many of the drugs used to battle cancer.

Maximum tolerated dose reached in CORIST Phase IIa Part 3 continuation trial

Main focus for the second guarter and the period after has been to finalize the continuation trial of the CORIST Phase IIa Part 3 trial evaluating SCO-101 as a combination treatment with FOLFIRI chemotherapy in mCRC patients, with the primary endpoint being to establish the maximum tolerated dose (MTD). Recently, in August, we were happy to announce topline data in the continuation trial and that MTD was achieved, thus the goal of the trial was reached ahead of schedule.

The best dose level for a 4-day schedule of SCO-101 in combination with FOLFIRI was found to be 250 mg daily of SCO-101, 50% irinotecan and 100% Leucovorin and 5-FU. Of the 3 patients included in the continuation trial, 2 experienced a dose-limiting toxicity of neutropenia at 65% irinotecan, which was expected based on previous data. No new safety signals were detected, confirming SCO-101 remains a safe compound to use.

Establishing the MTD was a final step, giving us the recommended dose that we can use in the next steps of SCO-101's development. We expect final data in the CORIST part 3 continuation trial in the first half of 2025.

PANTAX Phase IB data confirms the safety of SCO-101

During the quarter we also reported final data from our PANTAX phase Ib trial, an open-label international multi-center trial evaluating SCO-101 in combination with standard of care chemotherapies gemcitabine and nab-paclitaxel in patients with advanced pancreatic cancer. It was very encouraging to see that the data confirmed the good safety profile of SCO-101.

Increased focus on business development and partnering opportunities

During the quarter we concluded a rights issue, raising about SEK 31 million, before issue costs. In the event of exercise of warrants of series TO 2 and TO 3, in November 2024 and April 2025, respectively, the company will receive additional proceeds. With positive data and new financing, we announced after the end of the second guarter that the top priority is business development and partnering activities.

As part of these efforts, Scandion is working together with Back Bay Life Science Advisors LLC, a prominent life sciences investment banking firm, to explore and evaluate actionable strategic and financial alternatives. This will be an important step for Scandion to reach its future potential.



Francois R. Martelet

We are grateful to our shareholders for their continued support during difficult market conditions.

Scandion truly has a unique mechanism of action and SCO-101 continues to show positive data, encouraging us to continue its development.

Thanks for following Scandion Oncology.

Francois Martelet, M.D.

CEO

Scandion Oncology A/S – The Cancer Drug Resistance Company



SCANDION ONCOLOGY AND THE THERAPY

THE COMPANY

Scandion Oncology is a clinical-stage biotechnology company developing first-in-class medicines aimed at treating cancer which is resistant to current treatment options.

One of the most significant challenges in modern oncology is how to treat tumors that are or have become resistant to prescribed anti-cancer drugs. Scandion Oncology's most advanced innovative drug, SCO-101, is an oral drug that in preclinical studies has been documented to reverse resistance towards some of the most commonly used anti-cancer drugs.

The uniqueness of SCO-101 lies in its specific and dual-targeting mechanism of action. Unlike traditional single-target therapies, SCO-101 specifically targets the protein ABCG2 and the enzyme UGT1A1 simultaneously.

Cancer cells often exhibit redundancy and compensatory mechanisms and targeting only a single protein may lead to acquired resistance. SCO-101 addresses this challenge by simultaneously inhibiting a key enzyme and protein, leading to a more profound impact on exposure of cancer cells to cancer therapy.

SCO-101 represents a novel approach in targeted therapy. By concurrently addressing a key enzyme and protein important for exposure and effect of cancer therapeutics, it aims to maximize therapeutic efficacy while minimizing the risk of resistance development.

SCO-101 is currently being tested in a clinical phase Ib and a phase IIa trial in cancer patients.

Scandion Oncology has additionally other products in its pipeline targeting cancer drug resistance as future development opportunities. All with the aim to be the Cancer Drug Resistance Company.

THE THERAPY

All cancer patients with metastatic disease fail their cancer treatment – largely due to their cancer cells either being resistant already from the time of the primary diagnosis or because the cancer cells acquire resistance during anti-cancer treatment.

As a result, the cancer continues to grow despite treatment and without any other effective drugs, the patients are left to fight the growing cancer on their own.

Therefore, drug resistance is a major threat to cancer patients and a huge burden on the health care systems. As such, it also presents a significant commercial opportunity for Scandion Oncology.

The Global Cancer Chemotherapy Market Size accounted for USD 41 Billion in 2021 and is estimated to garner a market size of USD 106 Billion by 2030 rising at a CAGR of 11.5% from 2022 to 2030.

An add-on therapy such as SCO-101 would be able to tap into a share of this market and reach adoption fast.

At Scandion Oncology we are not aware of any drugs that are registered for blocking anti-cancer drug resistance.

SCANDION ONCOLOGY IN BRIEF

MISSION

To bring new medicines to patients in order to overcome cancer drug resistance and improve lives for cancer patients and their families

7,263

SHAREHOLDERS JUNE 30, 2024

27 MDKK

CASH POSITION JUNE 30, 2024

67 MSEK

MARKET CAP JUNE 30, 2024



2 CLINICAL PROGRAMS

CORIST currently in Phase IIa, (~100 subjects dosed), PANTAX in Phase Ib



PIPELINE

SCO-101 SCO-201 800 analogues



CANCER INDICATIONS

Colorectal, Pancreatic, Gastric and others



PEOPLE

Current, permanent staff of 4 employees as of June 30, 2024 Office in Copenhagen, Denmark



LISTED STOCK EXCHANGE

Nasdag First North Stockholm



O2 2024 REPORT



PIPELINE AND STRATEGY

CLINICAL PIPELINE

Developing First-in-class Medicines for Personalized Therapy

Scandion Oncology is currently developing a unique First-in-class lead compound SCO-101 – an oral add-on therapy to standard anti-cancer treatment. The most advanced program, CORIST, is a clinical phase IIa study for the treatment of drug resistant metastatic colorectal cancer (mCRC). The second program, PANTAX, is a clinical phase Ib study for the treatment of unresectable or metastatic pancreatic cancer.

First-in-class medicine

There are currently no drugs on the market targeting cancer drug resistance, and SCO-101 has the potential to be first in mCRC of treatments and become the defining drug for a group of patients in very high need of medical innovation.

Personalized therapy

Scandion Oncology is developing predictive biomarkers in conjunction with the ongoing CORIST and PANTAX studies, to enable a personalized medicine approach for the use of SCO-101.

Scandion Oncology's Clinical Pipeline

Program	Compound	Indication	Discovery / Pre-clinical	Phase I	Phase II	Phase III
CORIST	SCO-101	Colorectal cancer	SCO-101 + FOLFIRI			
PANTAX	SCO-101	Pancreatic cancer	SCO-101 + nab-paclitaxel and ger	ncitabine		

ACHIEVED MILESTONES

- **CORIST:** Final data from the phase IIa, part 2 trial released Q4, 2023
- **CORIST:** Topline results from part 3 released January 2024
- PANTAX: Final data from the phase lb trial released May 2024

UPCOMING KEY EVENTS

- **CORIST:** Final data from part 3 is expected in H2, 2024
- **CORIST:** Final data from the part 3 continuation trial is expected in H1, 2025



CORIST

For the Treatment of Patients with Metastatic Colorectal Cancer

In the CORIST phase IIa study, patients with chemotherapy resistant metastatic colorectal cancer (mCRC) receive SCO-101 treatment together with the standard chemotherapy drug combination FOLFIRI. All patients enrolled in the trial have previously demonstrated FOLFIRI resistance.

The first part of the CORIST phase IIa study, which aimed at establishing a safe dose of SCO-101 when given together with FOLFIRI, has been successfully completed and positive interim results were presented in June 2021.

The interim results led Scandion to continue the second part of the CORIST phase IIa study (part 2) in RAS wild-type patients. This second part of the CORIST phase IIa study has completed recruitment of 25 patients, and continues the focus on safety, tolerability, and efficacy parameters, to establish initial proof-of-concept for SCO-101 in mCRC on a schedule combining SCO-101 and FOLFIRI.

Topline data from CORIST part 2 have been released end of Q3, 2022. The topline results confirmed the safety and tolerability of SCO-101 in this indication and combination. Further, tumor reductions were observed in some patients, however below the 30% threshold defined as the trial's primary endpoint. Also, indication of prolonged progression free survival and stable disease (secondary endpoints) were observed in this hard-to-treat refractory patient population.

The final results from the part 2 analysis are highly positive as data show impressive overall survival for the patients participating in the trial. Further, four out of the 25 patients had shrinkage of their tumors, and the Clinical Benefit Rate evaluated after 8 weeks was 42%. Also, a potential biomarker for identifying patients most likely to respond to the treatment was identified in the trial. As already communicated last year, the data also confirmed the safety and tolerability of SCO-101.

Specifically, the data shows a median Overall Survival (mOS) of 10.4 months in CORIST part 2 with historical data for placebo or best supportive care having been reported in the range of 5-7 months in large international, multicenter, randomized, double-blinded phase III trials. A subset of patients (17 out of 25) had mOS of 13.4 months. This impressive data from CORIST is important, since mOS is the gold standard in oncology trials and an important regulatory endpoint. It is encouraging to see tumor reductions in four patients, a high proportion in this group of refractory hard-to-treat patients.

In January 2024, positive topline phase IIa data from the CORIST part 3 trial was reported, and impressive tumor reduction of more than 30% (partial response) was observed in one patient in the last cohort (out of 21 evaluated patients).

In March 2024 another partial response was reported in the last trial cohort, meaning that two of the six total patients have had a partial response, i.e. tumor reduction of more than 30%.

Median Progression Free Survival (PFS) was 4.6 months in Part 3, superior to the PFS reported in CORIST part 2, and Clinical Benefit Rate (CBR) was 76% after eight weeks of treatment, a significant increase from the 42% CBR from CORIST part 2.

In August 2024 Scandion achieved Maximum Tolerated Dose (MTD) for CORIST part 3. The established MTD for a 4-Days schedule of SCO-101 in combination with FOLFIRI was found to be 250 mg daily SCO-101, 50% irinotecan and 100% Leucovorin and 5-FU. The continuation study of CORIST part 3 included 3 patients. The dose of SCO-101 was the same as in the previous cohort, i.e., 250 mg per day for four days. Folinic acid and 5-FU were administered as per standard of care. The dose of irinotecan was increased from 50% to



65% of the normal standard dose. Of the 3 patients, 2 experienced a dose-limiting toxicity of neutropenia, which was expected based on previous data. No new safety signals were detected.

Overview of the CORIST phase IIa study

	CORIST Part 1		CORIST Part 2	CORIST Part 3				
Primary endpoint	MTD			Objective response		MTD		
Patients (N)		18 patients		25 patients (gCSF mandated)			tients mmended)	
Populations (mCRC)		All-comers		K-Ras wild type		All-co	mers	
SCO-101 (mg) and Patients (N)	150mg (4)	150mg (8)	100mg (6)	150mg (25)	150mg (7) 200mg (4)		200mg (7)	250mg (10)
Dose IRI (%)	80%	65%	50%	50%	50%			
Dose FOL and 5-FU (%)	80%	65%	50%	50%	100%			
Schedule		D-101: Days LFIRI: Days 5		SCO-101: Days 1-6 FOLFIRI: Days 5-7	SCO-101: Days 1-6 FOLFIRI: Days 2-4 FOLFIRI: Days 2-4			
Main outcome	• RP2D used in part 2 decided by the DSMB		Impressive OS Potential biomarker 6 patients with tumor reduction	 MTD established for 4 day schedule at 250 mg Potential biomarker associated with a longer PFS and OS Two patients had a partial response (i.e., 30% or more tumor reduction was observed) Meaningful improvements to PFS and CBR compared to Part 2 Awaiting final OS data; follow up ong 		with onse on		

ABOUT THE DISEASE

Colorectal cancer (CRC) is one of the most common cancers worldwide with over 0.5 million new cases every year in the US and EU. Unfortunately, a large proportion of patients diagnosed with CRC will develop metastatic disease (mCRC) despite prior adjuvant treatment and approximately 20% of newly diagnosed CRC patients have already developed metastatic disease at the time of diagnosis. The standard of care for patients with mCRC is either surgery and/or chemotherapy and targeted therapy with monoclonal antibodies.

For incurable patients, standard drugs are 5-FU and derivatives, oxaliplatin, irinotecan, bevacizumab and panitumumab or cetuximab. The anti-cancer agent irinotecan is most often prescribed in combination with 5-FU and leucovorin (FOLFIRI). One major problem in the treatment of mCRC is the frequent development of drug resistance. In practical terms, this means that the cancer continues to either grow during the anti-cancer treatment (de novo resistance) or re-grow after an initial response to the anticancer treatment (acquired resistance).

PANTAX

For the Treatment of Patients with Unresectable or Metastatic Pancreatic Cancer

In the PANTAX phase Ib study, patients with unresectable or metastatic pancreatic cancer receive SCO-101 treatment in combination with nab-paclitaxel and gemcitabine which is standard first- or second-line therapy.

The PANTAX phase Ib dose-finding study was initiated in Q4, 2020 and patients were enrolled from clinical sites in Denmark and Germany. In August 2022, Scandion announced that due to good tolerability the dosing was escalated to higher levels than expected based on the initial findings in the CORIST trial, which prompted the amendment of the PANTAX trial design communicated in January 2021. The continued dose escalation extended the PANTAX trial and enrollment was completed in H1, 2023.

Topline data from the PANTAX phase Ib study were released on March 31, 2023. The primary endpoint was achieved, as the maximum tolerated dose of Scandion's lead compound SCO-101 in combination with standard of care chemotherapies gemcitabine and nab-paclitaxel in patients with advanced pancreatic cancer was established at 200 milligrams given for 6 consecutive days every 2 weeks.

In May 2024 final data was published confirming the MTD of 200 mg. Further PK data demonstrated that the exposure of SCO-101 was in line with the expectations. 15 patients were evaluable for response and 1 had a PR resulting in an ORR of 6.7%. Amongst the 15 evaluable patients CBR was 53%. Progression-free survival (PFS) was 2.5 months and overall survival (OS) was 9.5 months.

About the PANTAX study

In the PANTAX study, patients with unresectable or metastatic pancreatic cancer receive SCO-101 treatment in combination with nab-paclitaxel and gemcitabine which is standard first- or second-line chemotherapy.

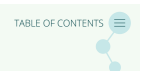
The aim of the phase Ib study is to establish a safe dose (maximum tolerated dose) of SCO-101 in combination with nab-paclitaxel and gemcitabine.

ABOUT THE DISEASE

Approximately 150,000 patients in the US and EU are newly diagnosed with pancreatic cancer each year. Pancreatic cancer has a very high unmet need, with poor prognosis and high treatment failure rates. Despite the comparably low incidence, it is the 3rd leading cause of cancer death in the US and 7th worldwide. Approximately 70% of diagnosed patients have a life expectancy of less than 1 year without adequate treatment and patients with metastatic disease (50-55%) have a limited survival of only 3 to 6 months.

The treatment paradigm for pancreatic cancer is predominantly composed of chemotherapies, most notably FOLFIRINOX or gemcitabine and nab-paclitaxel.

Pancreatic cancer has a high frequency of primary (de novo) resistance against chemotherapy, but also fast development of secondary (acquired) resistance is a major problem. This means that most patients who initially experience a positive effect of the chemotherapy, will experience disease progression relatively fast.



PRE-CLINICAL PIPELINE

Building Future Value

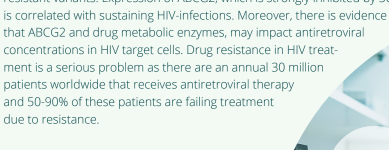
Scandion Oncology's Pre-clinical Pipeline

Program	Compound	Indication	Discovery / Pre-clinical	Phase I	Phase II	Phase III
101	SCO-101	Other cancer indications				
201	SCO-201	Solid tumors/ HIV				

Scandion has completed pre-clinical studies confirming that the company's lead compound, SCO-101, could potentially be an effective treatment for gastric cancer. SCO-101 is currently being clinically developed as a combination treatment for metastatic colorectal cancer and pancreatic cancer, presenting gastric cancer as an appealing new opportunity for Scandion.

It has been well documented in scientific literature that the protein ABCG2 is overexpressed in gastric cancer cells and that high ABCG2-expression is associated with poor clinical outcome (i.e., reduced survival). Scandion's pre-clinical studies have confirmed that ABCG2, which SCO-101 specifically inhibits, is overexpressed in gastric cancer cells, meaning that gastric cancer cells will be sensitive to SCO-101 treatment. SCO-101 works synergistically with chemotherapy in ABCG2-positive cells. This is similar to colorectal cancer in which we have seen impressive overall survival (OS) for patients when SCO-101 is combined with chemotherapy.

SCO-201 is a potent anti-viral molecule blocking early stages of viral replication. The anti-viral effect has been demonstrated in vitro and in vivo for Picornaviridae, especially Rhino and Enterovirus, and in drug resistant variants. Expression of ABCG2, which is strongly inhibited by SCO-201,





SCANDION ONCOLOGY INTELLECTUAL PROPERTY

Scandion Oncology is diligently expanding and strengthening the Company's portfolio of intellectual property rights providing valuable long term commercial exclusivities.

At the end of Q2, 2024, Scandion Oncology owned a portfolio of twelve patent families, taking effect in commercially relevant countries.

Changes to Scandion Oncology's patent portfolio will be updated continuously and will be summarized in the Company's quarterly reports.

IP related events of high strategic value for the Company will be announced through press releases.

IP PORTFOLIO **NEWS**

- ON NOVEMBER 23, 2023, SCANDION WAS GRANTED NEW COMPOSITION OF MATTER-PATENT ON LEAD COMPOUND SCO-101 EXTENDING IT'S EXCLUSIVITY **UNTIL AT LEAST 2042**
- ON JANUARY 5, 2024, SCANDION RECEIVED NOTICE OF ALLOWANCE FOR PATENT TO **ENHANCE US PATENT EXCLUSIVITY ON** SCO-101. WHEN GRANTED, THE PATENT WILL OFFER A VERY BROAD INTELLECTUAL PROTECTION UNTIL AT LEAST 2037.





FINANCIAL REVIEW

Results of operations

Total operating expenses in Q2, 2024 reached 9.4 MDKK (11.8), a decrease of 2.4 MDKK compared to Q2, 2023, which reflects savings in study costs due to reduced activity level as studies progress.

Operating expenses can be divided into two main cost groups, Research & Development and General & Administration expenses. Research & Development expenses in Q2, 2024 of 5.7 MDKK (8.3), relate to the two clinical studies, CORIST and PANTAX. General & Administration expenses in Q2, 2024 amounted to 3.7 MDKK (3.5).

Operating loss for Q2, 2024 was 9.4 MDKK (11.3).

In Q2, 2024, net financial items amounted to 0.2 MDKK (0.1), which mainly derives from interest and currency adjustments.

The total comprehensive loss for the period is 7.2 MDKK (8.8).

Financial position

Total assets as of June 30, 2024, were 39.0 MDKK (60.2). Hereof, cash and cash equivalents amounted to 27.0 MDKK (45.7).

Receivables amounted to 7.1 MDKK (7.3) which mainly relates to income tax receivables in the amount of 5.5 MDKK (5.5), other receivables of 1.2 MDKK (1.1) and prepayments of 0.4 MDKK (0.7).

The equity ratio as of June 30, 2024 was 79% (87%), and equity was 30.7 MDKK (52.3).

Cash flow and Cash Position

The cash flow from operating activities in Q2, 2024 was an outflow of 3.9 MDKK (14.3) and is explained mainly by the loss before tax and change in working capital. The cash flow from investing activities was 0.0 MDKK (0.0). The cash flow from financing activities was an inflow of 13.9 MDKK (outflow of 0.2) mainly related to the Rights Issue carried out in June 2024.

Hence, the total net cash flow for Q2, 2024 was a net cash inflow of 10.0 MDKK (outflow of 14.5) leaving the company with a cash position of 26.9 MDKK as of June 30, 2024.

With the cash position as of June 30, 2024, and with the net proceeds from the Rights Issue carried out in June 2024, Scandion Oncology is sufficiently capitalized to fund ongoing activities in to Q4 2025.

(Numbers in brackets represent the corresponding reporting period last year)



SHAREHOLDER INFORMATION

The share

The shares of Scandion Oncology A/S are listed on Nasdaq First North Growth Market Sweden.

Scandion Oncology's share capital amounts to 17,047 TDKK divided into 231,928,544 shares of nominal value 0.0735 DKK each. There is only one class of shares, and each share represents one vote.

As of June 30, 2024, the number of shares was 231,928,544 (40,706,972).

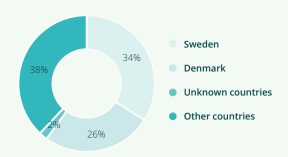
Shareholders

There are no individual shareholders that own 5% or more of the shares in Scandion Oncology as of June 30, 2024.

According to the shareholder register maintained by Euroclear Sweden AB, Scandion Oncology had 7,263 (7,842) shareholders as of June 30, 2024.

Listing	First North Growth Market Sweden
Number of shares	231,928,544 (40,706,972)
Share price (June 30, 2024)	0.29 SEK (1,26 SEK)
Market capitalization (June 30, 2024)	67 MSEK (51 MSEK)
Ticker	SCOL
ISIN	DK0061031895

Shareholders by country, June 30, 2024



Source: Monitor by Modular Finance AB.

Share-based incentive schemes

At the Annual General meeting on April 27, 2022 a warrant program was approved, authorizing the Board of Directors to issue up to 4,177,620 warrants which carry the right to subscribe for an equal number of shares in Scandion Oncology A/S. As of June 30, 2024 a total of 417,762 warants has been issued to the Board of Directors and a total of 1,882,033 warrants has been issued to the Executive Management and Employees – giving 2,299,795 warrants issued in total.

Warrant scheme related to Rights Issue

At the Annual General meeting on May 6, 2024 the board of directors were authorized to issue warrants (TO 2) and (TO 3) and any related capital increase(s) coinciding with the Rights Issue. The number of issued warrants in the TO 2 series was 143.416.179 respectively 47.805.393 in th TO 3 series which potentially, if utilized in full, will add TDKK 14,055 to the share capital. Please see note 5 for more details.

Share price

The Scandion Oncology share price on June 30, 2024 was 0.29 SEK (1.26), equivalent to a market capitalization of 67 MSEK (51 MSEK).

Relative to Q2, 2023, the average, daily turnover of Scandion Oncology shares was 0.3 MSEK in Q2, 2024 equivalent to an daily turnover increase of 0.2 MSEK.

(Numbers in brackets represent the corresponding reporting period last year)



12 month share price development and trading volume, July 1, 2023 to June 30, 2024



PUBLIC PRESENTATIONS

Date

Event

Sep 18, 2024

Nordic Life Science Days 2024, Stockholm, SE



ANALYST COVERAGE

Scandion Oncology is covered by the following analysts:

Redeye AB

(Christian Binder)



CORPORATE MATTERS

FINANCIAL CALENDAR

November 27, 2024 Q3 report 2024 February 28, 2025 Year-end report 2024



Forward looking statements

This financial report includes statements that are forward-looking, and actual future results may differ materially from those stated. In addition to the factors explicitly commented upon, other factors that may affect the actual future results are for example development within research programs, including development in pre-clinical and clinical trials, the impact of competing research programs, the effect of economic conditions, the effectiveness of the company's intellectual property rights and preclusions of potential second party's intellectual property rights, technological development, exchange rate and interest rate fluctuations and political risks.

For further information, please contact

Johnny Stilou, CFO

T: +45 29 60 35 32

E: jos@scandiononcology.com

The information was provided by the contact person above for publication on August 29, 2024 at 07.00 CET.

Certified Advisor

Västra Hamnen Corporate Finance AB.

From September 1, 2024 Vator Securities AB is appointed Certified Advisor

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STATEMENT BY THE BOARD OF DIRECTORS

The Board of Directors provides their assurance that the Q2 2024 report provides a fair and true overview of the Company's operations, financial position, and results.

Copenhagen, August 29, 2024 The Board of Directors of Scandion Oncology A/S

Martin Møller Chairman of the Board

Alejandra Mørk Deputy chairman of the Board

Keld Flintholm Jørgensen *Member of the Board of Directors*

Per Pfeiffer Member of the Board of Directors

The interim report has not been audited or reviewed by the company's auditors.





STATEMENT OF COMPREHENSIVE INCOME

TDKK	Q2 2024	Q1-Q2 2024	Q2 2023	Q1-Q2 2023	FY 2023
Other operating income	0	0	510	684	446
Other operating costs	0	0	0	0	-220
Research and development expenses	-5,676	-11,990	-8,321	-17,111	-31,631
General and administration expenses	-3,683	-7,468	-3,507	-6,865	-13,952
Operating loss	-9,359	-19,458	-11,318	-23,292	-45,357
Financial items					
Financial income	279	604	335	670	1,640
Financial expenses	-54	-186	-269	-539	-987
Loss before tax	-9,133	-19,040	-11,253	-23,160	-44,704
Tax	1,969	4,307	2,478	5,098	5,500
Net loss for the period	-7,164	-14,733	-8,774	-18,062	-39,204
Other comprehensive					
income for the period	0	0	0	0	0
Total comprehensive loss	-7,164	-14,733	-8,774	-18,062	-39,204



BALANCE SHEET

TDKK	Q1-Q2 2024	Q1-Q2 2023	FY 2023
Assets			
Non-current assets			
Equipment	199	568	151
Right of use assets	179	1,198	497
Deposits	249	290	249
Income Tax receivables	4,323	5,098	0
Total Non-current assets	4,949	7,154	897
Current Assets			
Prepaid expenses and accrued income	397	712	612
Other receivables	1,189	1,111	1,032
Income Tax receivables	5,500	5,500	5,500
Cash and cash equivalents	26,952	45,709	26,520
Total current assets	34,037	53,032	33,664
Total Assets	38,987	60,186	34,560
Equity and liabilities			
Equity			
Share capital	17,047	2,992	2,992
Share premium reserved	233,291	233,008	233,008
Retained earnings	-219,611	-183,735	-204,878
Total Shareholders equity attributable to Shareholders	30,727	52,265	31,122
Non-current liabllities			
Lease liabilities	0	603	0
Other non-current liabilities	0	941	0
Total non-current liabilities	0	1,544	0
Current liabilities			
Lease liabilities	180	603	499
Account liabilities	1,939	2,361	1,381
Other current liabilities	6,141	3,412	1,558
Total current liabilities	8,260	6,377	3,438
Total equity and liabilities	38,987	60,186	34,560

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EQUITY

1/1 2024 - 30/6 2024 TDKK	Share capital	Share premium	Retained earnings	Total equity
Balance at January 1, 2024	2,992	233,008	-204,878	31,122
Comprehensive loss				
Result for the period			-14,733	-14,733
Net comprehensive loss			-14,733	-14,733
Transaction with owners				
Increase of Capital	14,055	6,199		20,254
Expenses related to capital increase		-5,917		-5,917
Share-based compensation expenses				
Net transactions with owners	14,055	283	0	14,337
Balance at June 30, 2024	17,047	233,291	-219,611	30,727

1/1 2023 – 30/06 2023 TDKK	Share capital	Share premium	Retained earnings	Total equity
Balance at January 1, 2023	2,992	233,008	-165,673	70,327
Comprehensive loss				
Result for the period			-18,061	-18,061
Net comprehensive loss			-18,061	-18,061
Transaction with owners				
Increase of Capital				
Expenses related to capital increase				
Share-based compensation expenses				
Net transactions with owners	0	0	0	0

1/7 2023 - 31/12 2023 TDKK	Share capital	Share premium	Retained earnings	Total equity
Balance at July 1, 2023	2,992	233,008	-183,735	52,265
Comprehensive loss				
Result for the period			-21,143	-21,143
Net comprehensive loss			-21,143	-21,143
Transaction with owners				
Increase of Capital				
Expenses related to capital increase				
Share-based compensation expenses				
Net transactions with owners	0	0	0	0
Balance at December 31, 2023	2,992	233,008	-204,878	31,122



CASH FLOW STATEMENT

TDKK	Q2 2024	Q1-Q2 2024	Q2 2023	Q1-Q2 2023	FY 2023
Operating activities					
Result before tax	-9,133	-19,040	-11,253	-23,160	-44,704
Non-cash sharebased payments	0	0	0	0	0
Financial items, reversed	-225	-418	-66	-131	-654
Depreciation, reversed	110	219	245	490	969
Change in working capital	5,131	5,366	-3,272	-8,835	-12,432
Cash flow from operating activities before financial items	-4,118	-13,873	-14,345	-31,636	-56,821
Interest and exchange rate gains	279	604	335	670	1,640
Interest and exchange rate losses	-54	-186	-269	-539	-987
Corporate tax received	0	0	0	0	5,500
Cash flow from operating activities	-3,893	-13,455	-14,280	-31,505	-50,668
Investing activities					
Equipment	0	88	0	0	0
Sale, tangible assets	0	0	0	0	247
Financial assets	0	0	0	0	41
Cash flow from investing activities	0	88	0	0	288
Financing activities					
Contributed capital	19,895	19,895	0	0	0
Expenses related to capital increase	-5,917	-5,917	0	0	0
Lease payments	-89	-179	-196	-391	-705
Cash flow from financing activities	13,889	13,799	-196	-391	-705
Net cash flow for the period	9,996	432	-14,476	-31,895	-51,085
Cash and cash equivalents beginning of the period	16,956	26,520	60,185	77,605	77,605
Cash and cash equivalents end of the period	26,952	26,952	45,709	45,709	26,520



NOTES

NOTE 1:

GENERAL INFORMATION

Scandion Oncology A/S (the "Company"), Corporate Registration Number DK-38613391, is a limited liability company, incorporated and domiciled in Denmark. The Company is

listed at Nasdag First North Growth Market under the ticker SCOL and the ISIN code DK0061031895. The registered office is at Fruebjergvej 3, 2100 Copenhagen, Denmark.

NOTE 2:

ACCOUNTING POLICIES

Basis for Preparation

The interim financial statements have been prepared in accordance with IAS 34, Interim Financial Reporting, as adopted by EU and the additional requirements for submission of interim reports for companies listed on Nasdaq First North Growth Market Sweden.

The interim financial statements are presented in Danish kroner (DKK) which is the functional currency of the Company. All values are presented in thousand DKK and all amounts are rounded to the nearest thousand DKK

New IFRS standards & interpretations

There are no IFRS standards and interpretations issued before the end of this reporting period of relevance for the Company, which are expected to change current accounting regulation significantly.

Foreign currency translation

On initial recognition, foreign currency transactions are translated at the exchange rate at the transaction date. Receivables, liabilities and other monetary items denominated in foreign currency that have not been settled at the balance sheet date are translated at closing rates.

Foreign exchange differences between the rate of exchange at the date of the transaction and the rate of exchange at the date of payment or the balance sheet date, respectively, are recognized in the income statement under financial items.

Definitions

Earnings per share (EPS) and diluted earnings per share (EPS-D) are calculated in accordance with IAS 33.

Other key ratios are calculated in accordance with the online version of "Recommendations and Ratios" issued by The Danish Finance Society and CFA Society Denmark.

EQUITY RATIO:

Equity (end of year) * 100

Total assets

EARNINGS PER SHARE BASIC (EPS):

Net result

Average number of shares in circulation

DILUTED EARNINGS PER SHARE (EPS-D):

Net result

Diluted average number of shares in circulation

SHAREHOLDERS' **EQUITY PER SHARE:**

Equity

Number of shares, year end

25



NOTE 3:

CRITICAL ACCOUNTING ESTIMATES AND JUDGEMENTS

In preparing the interim financial statements, management makes various accounting judgements and estimates and define assumptions, which form the basis of recognition, measurement and presentation of the company's assets and liabilities.

The estimates and assumptions applied are based on historical experience, the most recent information available at the reporting date, and other factors that management considers reasonable under the circumstances.

The basis for judgements and information can by nature be inaccurate or incomplete, and the Company is subject to uncertainties, which can result in an actual outcome that deviates

from estimates and defined assumptions. It may be necessary in the future to change previous estimates and judgements as a result of supplementary information, additional knowledge and experience or subsequent events.

In applying the Company's accounting policies described in note 2, management has exercized critical accounting judgements and estimates, which significantly influence on the amounts recognized in the financial statements.

NOTE 4:

RISK MANAGEMENT

Various risk factors may have an adverse impact on Scandion Oncology's operations and therefore the Company's results and financial position. For Scandion Oncology the main operational impact is potential delays in clinical trials as sites could be restricted from patient enrollment, or changes in requirements from authorities.

A description of Scandion Oncology's risk exposure and risk management is included in the Annual Report 2023, note 18, page 49-50 (please see www.scandiononcology.com).

NOTES



NOTE 5:

WARRANT PROGRAMS

Share-based incentive scheme

At the Annual General meeting on April 27, 2022 a warrant program was approved, authorizing the Board of Directors to issue up to 4,177,620 warrants which carry the right to subscribe for an equal number of shares in Scandion Oncology A/S.

As of June 30, 2024 a total of 257,084 warrants has been issued to the Board of Directors and a total of 1,822,033 warrants has been issued to the Executive Management and Employees - giving 2,139,117 warrants issued in total.

Exercise price/strike price for the warrants is SEK 22.00. The fair value of the warrant program is zero and calculated in accordance with the Black-Scholes option pricing model.

Outstanding at June 30, 2024	2,139,117
Cancelled	-160,678
Granted	600,000
Outstanding at January 1, 2024	1,699,795

Warrant scheme related to Rights Issue

At the Annual General meeting on May 6, 2024 the board of directors were authorized to issue 346,153,848 TO 2 warrants (corresponding to a total nominal value of DKK 25,442,307.828 in the Company) and 115,384,616 TO 3 warrants (corresponding to a total nominal value of DKK 8,480,769.276 in the Company) and any related capital increase(s) coinciding with the Rights Issue.

The TO 2 and TO 3 Warrants were issued for the benefit of investors in connection with the subscription of Units in the Company's rights issue in June 2024 (the "Rights Issue") and the Company's existing shareholders thus have a right of preemption. Each Warrant in both series entitles (but does not obligate) the Warrant Holder to subscribe for one new share in the Company of nominally DKK 0.0735 at an exercise price equal to the volume-weighted average price of the Company's share on Nasdaq First North Growth Market.

The TO 2 warrants will be priced in the period 16-29 October 2024 (the "Pricing Period"), however not less than the shares' nominal value and not higher than 125 % of the subscription price per share in the Rights Issue (the "Exercise Price"). The excercise period for the TO 2 warrants is from 4-18 November 2024.

The TO 3 warrants will be priced in the period 14-27 March 2025 (the "Pricing Period"), however not less than the shares' nominal value and not higher than 125 % of the subscription price per share in the Rights Issue (the "Exercise Price"). The excercise period for the TO 3 warrants is from 2-16 April 2025.

The number of issued warrants in the TO 2 series was 143.416.179 (hereof 1,387,353 to Management and Board of Directors) respectively 47.805.393 in the TO 3 series (hereof 462,451 to Management and Board of Directors) which potentially, if fully utilized, will add TDKK 14,055 to the share capital. All Warrants are considered vested as of the time of allocation according to section 4.1 in appendix 4.4 and 4.5 in the Company's Article of Association.

Outstanding at June 30, 2024	191,221,572
TO 3 warrants	47,805,393
TO 2 warrants	143,416,179

27



NOTE 6:

CONTINGENT ASSETS AND LIABILITIES

License and Collaboration Agreements

Scandion own all rights to assets but are not yet entitled to potential milestone payments and royalties on successful commercialization of products developed under license and collaboration agreements with potential partners.

Pending commercial litigation

Scandion is not involved in commercial litigations arising out of the normal conduct of its business.

NOTE 7:

RELATED PARTIES

Apart from salaries and warrants there were no significant transactions with Management or Board of Directors.

NOTE 8:

SIGNIFICANT EVENTS AFTER THE BALANCE SHEET DATE

No significant events have occured after the end of the reporting period.

