

Extension of the highest dose level in the knee osteoarthritis study

Agreement with Region Östergötland for GMP manufacturing of cell therapy product

Expansion of Targinta's patent portfolio



The Group -

Second quarter 2024

- » Income amounted to TSEK 4 (0).
- » Loss before tax totalled TSEK 10,400 (loss: 18,401).
- » Loss per share* was SEK 0.02 (loss: 0.06).

First half year 2024

- » Income amounted to TSEK 303 (0).
- » Loss before tax totalled TSEK 21,773 (loss: 34,244).
- » Loss per share* was SEK 0.04 (loss: 0.11).

The Parent company

Second quarter 2024

- » Income amounted to TSEK 4 (0).
- » Loss before tax totalled TSEK 8,565 (loss: 11,151).
- » Loss per share* was SEK 0.02 (loss: 0.04).

First half year 2024

- » Income amounted to TSEK 303 (0).
- » Loss before tax totalled TSEK 18,254 (loss: 21,553).
- » Loss per share* was SEK 0.03 (loss: 0.07).
- » At June 30, 2024, the equity/assets ratio** was 44 % (17).

Significant events in the second quarter of 2024

» Xintela and EQGen Biomedical to collaborate to develop EQSTEM stem cell treatment for horses. (May 23, 2024)

Significant events after the end of the period

- » Xintela extends clinical study with XSTEM on knee osteoarthritis patients. (August 23, 2024)
- » Xintela signs agreement with Region Östergötland for GMP process development of cell therapy for burn patients. (August 29, 2024)

Note to the reader

The "company" refers to Xintela AB (publ), corporate registration number 556780-3480. All figures are given in TSEK unless otherwise stated. Amounts in parentheses: Comparative period of the preceding year.

Trademark

In addition to patents, the IP portfolio also currently includes seven trademarks - the company names XINTELA® and TARGINTA®, XINMARK® which is the name of Xintela's technology platform, and XSTEM® which is the name of Xintela's stem cell platform. EQSTEM® and CANISTEM® which are the company's brands for stem cell treatment for horses and dogs and XACT® which is the name of an analytical test for chondrocytes.

- * Earnings/loss per share: Profit/loss for the period divided by 567,190,598 shares, which was the average number of shares at June 30, 2024. In the year-earlier period, the number of average shares was 307,573,263.
- ** Equity/assets ratio: Equity divided by total capital.



From the CEO

Aiming for results from clinical studies with XSTEM and license agreement with EQSTEM

Our knee osteoarthritis study is nearing the end of the 18-month follow-up after treatment with XSTEM® and recruitment of patients continues in our difficult-to-heal leg ulcers study. The license agreement work for EQSTEM® progresses very well and, in addition, we have signed an agreement with Region Östergötland for isolation and quality assurance of skin cells from burn patients. In our subsidiary, Targinta, we are strengthening our patents for our candidate drugs.

Extension of the highest dose level in the osteoarthritis study

In our clinical study in patients with knee osteoarthritis, three dose levels of XSTEM are being evaluated. At present, patients at the two lower dose levels have completed or will soon complete the study, 18 months after dosing, as planned. Patients at the highest dose level will continue the study for an additional six months for evaluation at 24 months after dosing. The reason for this change in the protocol is that 24 months is a common time for investigating early regenerative effects on cartilage and other joint tissues. It is therefore a good time to compare our results with published results from other clinical studies. We will nevertheless perform an interim analysis of study data up to 18 months after dosing for all dose levels in Q1 2025. The possibility to expand the study with an additional 30 patients has not yet been activated.

We can report that the study progresses very well and as planned. The study's Safety Review Committee has previously assessed all three dose levels as safe at a three-month follow-up, and patients at all dose levels have reported reduced pain and improved joint function after 12 months. As we approach the end of our knee osteoarthritis study, we are also getting closer to possible partnerships that will take XSTEM further in clinical studies.

Recruitment of patients with difficult-to-heal leg ulcers continues

Recruitment work is back on track at the clinics after a long summer break. We are now collaborating with two clinics, in Gothenburg and in Stockholm. As we have previously communicated, recruitment to the leg ulcer study is a challenge because the patients are older and often have other diseases and complications, which makes participation difficult. The inclusion criteria are particularly tough because the study is first and foremost a safety study. We have seen great interest in participating in the study, for example after advertising, but most of the screened patients could not be included for various reasons. Three patients who have completed the study after four months have not shown any safety concerns. We continue to work intensively with the clinics to get the remaining patients in. In the meantime, we have been approached to start an investigator-initiated study with XSTEM for a wider group of wound patients and we are now investigating this interesting possibility.

Intensive collaboration with EQGen Biomedical to land license agreement

As previously announced, Xintela has signed a non-binding term sheet for a license agreement with EQGen Biomedical Inc. in the US, where EQGen Biomedical will receive global rights to Xintela's stem cell product EQSTEM for horses. Xintela will receive upfront payments, covered development costs and revenues from commercialization. The license agreement is contingent on, among other things, EQGen Biomedical securing financing for the continued development of EQSTEM. Xintela will participate, among other things, through process development and production of EQSTEM, in an arrangement that is fully financed by EOGen Biomedical. Work on the license agreement is ongoing at the same time as EQGen Biomedical is building up a fantastic team and is enthusiastically working to land financing for the development work. We have a very good collaboration with EQGen Biomedical and look forward to landing the license agreement and getting started with the development.







Agreement with Region Östergötland

We have recently announced that Xintela has signed an agreement with its first major customer, Region Östergötland. The agreement is worth SEK 3.6 million. The assignment involves Xintela developing and establishing a GMP process for isolating and quality assuring autologous (patient-own) skin cells for the treatment of burns. The process will form the basis for an approval from the Medical Products Agency for the Burn Centre, Linköping University Hospital, to start a clinical study on burn patients. Xintela will produce skin cell preparations from patient biopsies in Xintela's GMP-approved production facility. It is very positive that through the collaboration with Region Östergötland, we are now broadening the use of our GMP facility and our expertise in process development and production of other advanced drugs, so-called ATMPs (Advanced Therapy Medicinal Products) and generating revenues for Xintela.

Expands and strengthens Targinta's patent portfolio

We continue to work on finding financing and/or partners for our subsidiary Targinta and it's unique drug candidates TARG9 and TARG10 for the treatment of aggressive cancers. Targinta has developed First-in-Class antibodies and antibody-drug conjugates (ADCs) that target a new oncology target and have shown strong preclinical results in cancer models of invasive and difficult-to-treat solid cancers. In addition, Targinta has a strong patent portfolio with several approved patents. Now, we are further strengthening it by protecting Targinta's antibodies and ADCs in large and important markets.

Continued financing of our operations

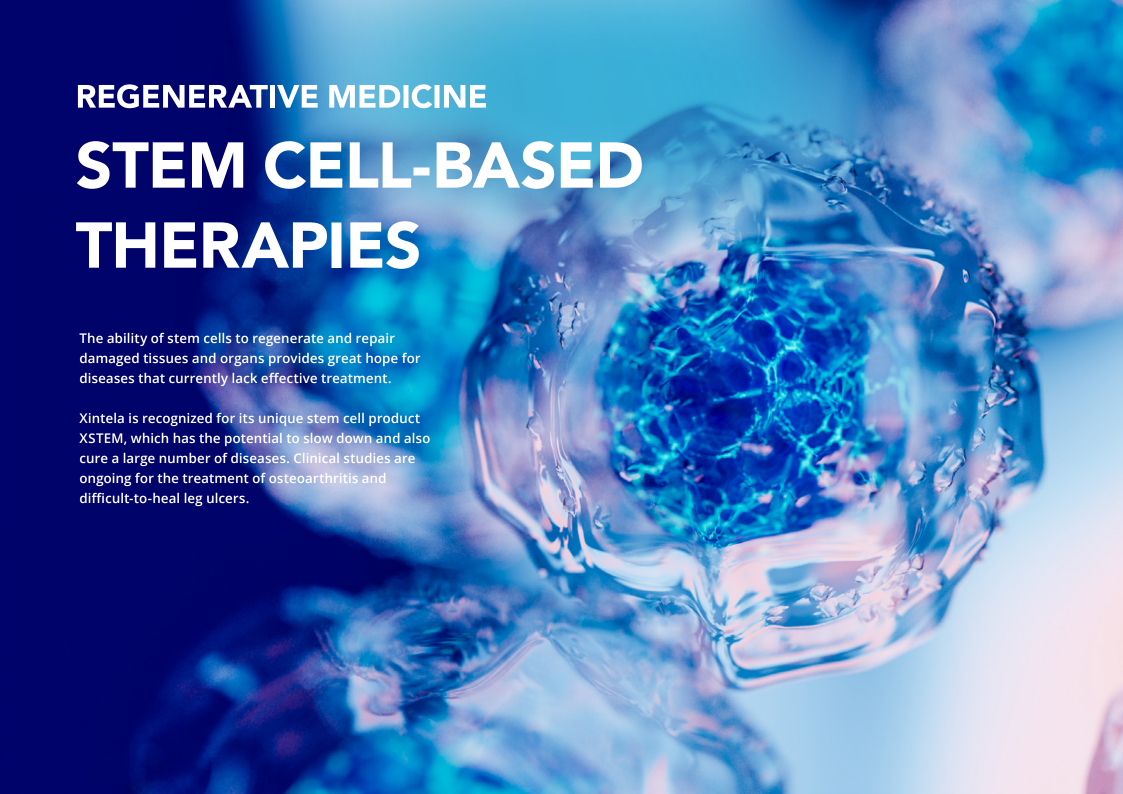
Our ambition is that financing of our development projects going forward will come mainly from development milestone revenues from collaborations, partnerships and licensing, such as from EQGen Biomedical and Region Östergötland. We are working in parallel with other financing solutions for Xintela and Targinta, such as capital raising, grants and loans, which can be carried out either jointly or separately.

We are now looking forward to completing our clinical studies and landing partnerships and commercial agreements.

Evy Lundgren-Åkerlund

CEO, Xintela AB (publ)





Xintela is strongly positioned to develop and commercialize safe and effective stem cell treatments

Xintela has developed the competitive stem cell product XSTEM, which consists of integrin α10β1-selected mesenchymal stem cells. Through the unique selection step in the production process, homogeneous stem cells of high and reproducible quality can be produced. XSTEM is manufactured in Xintela's own GMP facility and is patented both as a product and for therapeutic uses in all indications.







Mesenchymal stem cells have therapeutic properties

Xintela develops stem cell-based treatments from allogeneic (donated) mesenchymal stem cells isolated from adipose tissue from healthy adult donors. Stem cells from a donor can treat a large number of patients, which not only significantly reduces the cost of XSTEM compared to autologous (patient's own) stem cells but will also give physicians an off-the-shelf therapy. An important property of mesenchymal stem cells is their ability to transform into different cell types to regenerate and repair damaged tissues and organs. They also have the ability to stimulate damaged cells to self-repair. Another important property is that stem cells secrete various substances that can regulate the immune system and thus have anti-inflammatory effects.

Stem cell selection – a critical step in the production of XSTEM

Stem cell preparations produced from tissues are heterogeneous, i.e. they contain contaminating cells that are not stem cells. When developing a stem cell product, this is both a regulatory and functional problem.

Xintela solves the problem by selecting (purifying) stem cells using an antibody that binds to the company's stem cell marker, integrin $\alpha 10\beta 1$. In this way, homogeneous stem cell preparations of high quality can be produced that are reproducible between different donors.

Own GMP production of stem cells

Our stem cells are produced in bioreactors in the company's own GMP-approved facility and stored frozen until used in the treatment of patients. Through its in-house, production facility, Xintela has full control over the stem cell production which significantly reduces risks such as unexpected costs and delays. The company's strategy is to establish Xintela as a manufacturer of stem cell products developed in collaboration with partners and to also offer development and production of other advanced therapy products (ATMP).



OSTEOARTHRITIS

Osteoarthritis is a joint disease characterized by degradation of the articular cartilage and impaired function of the cartilage cells. It is the most common chronic joint disease, especially in the knees, hips and hands, as well as the most common cause of disability in the elderly. The main symptoms are severe pain, inflammation, stiffness in the joint and reduced mobility. The disease affects about 25 percent of all individuals over the age of 60 and is increasing in extent due to an increasing elderly population. Drugs offered today are primarily pain-relieving and anti-inflammatory, which treat the symptoms but not the actual cause of the disease. [1,2]





DIFFICULT-TO-HEAL LEG ULCERS

Difficult-to-heal leg ulcers in the elderly, including venous leg ulcers, are a major medical problem that results in pain and reduced quality of life for the patient, as well as large costs for healthcare systems.

The incidence increases with age and is estimated to be about 4 percent among people over 65 years of age. Today's treatments for difficult-to-heal leg ulcers include compression techniques and various surgical techniques, but there is a lack of effective drugs. [1,2]

Steady progress in XSTEM clinical studies

XSTEM in clinical study for the treatment of knee osteoarthritis

Xintela is conducting a clinical study (Phase I/IIa) with XSTEM in Australia, in patients with moderate knee osteoarthritis (Kellgren-Lawrence grade II-III). Three different dose levels of XSTEM are being evaluated in up to 54 patients and each patient is followed for 18 months with safety evaluation and preliminary efficacy evaluation every six months. The highest dose level has recently been extended by six months to obtain additional efficacy data for XSTEM 24 months after dosing. XSTEM have been dosed at all dose levels in a total of 24 patients and all dose levels have been judged safe by the study's Safety Review Committee after three months. All patients at the low and mid dose level have completed, or will shortly complete the study, 18 months after treatment. The primary goal of the study is to show that XSTEM is safe, but also to obtain preliminary efficacy results that show that the product has DMOAD (Disease Modifying Osteoarthritis Drug) properties and can slow down cartilage and joint degradation as well as restore damaged articular cartilage and improve joint function. Xintela's earlier results from preclinical osteoarthritis models, support the possibility that XSTEM may have a positive disease-modifying effect.

An interim analysis will be performed in Q1 2025 on 18-month data for all dose levels. The dose escalation part of the study

will end in June 2025 after the 24-month follow-up of the highest dose level. The possibility to expand the study with an additional 30 patients has not yet been activated. In parallel with the clinical study being conducted, discussions with potential partners and licensees for further clinical development and commercialization are ongoing.

XSTEM in clinical study for the treatment of difficultto-heal venous leg ulcers

Xintela's clinical study (Phase I/IIa), in patients with difficult-to-heal venous leg ulcers, is being conducted in Sweden. Twelve patients with difficult-to-heal venous leg ulcers will be treated with XSTEM or placebo. XSTEM/placebo will be applied to the wound and patients will then be followed weekly for ten weeks as well as at four months after treatment. The primary goal of the study is to show that XSTEM is safe, but also to evaluate wound healing efficacy. The first three patients have completed the study and recruitment and screening of patients is ongoing. A major part of the study is funded by a grant from Vinnova.

Xintela has previously shown in a preclinical wound model that XSTEM has excellent wound healing capacity, which gives great hope that XSTEM will show effective healing on patients difficult-to-heal leg ulcers.

Market

Osteoarthritis

The global market for osteoarthritis is mainly driven by an increase in an aging population, as well as a significant increase in obesity, but osteoarthritis can also affect young and middle-aged individuals. The market for drug treatment of osteoarthritis was estimated to be USD 7.3 billion in 2020 and is expected to grow by approximately 9 percent annually until 2025, when the market is estimated at USD 11.0 billion.[3]

Venous leg ulcers

In 2018, the global market for the treatment of venous leg ulcers was estimated at USD 2.95 billion, a figure that is expected to increase to USD 4.84 billion by 2026 with an average annual growth rate of 6.4 percent. The increase is partly due to the expectation that the incidence of venous leg ulcers will increase in line with an aging population.[4]

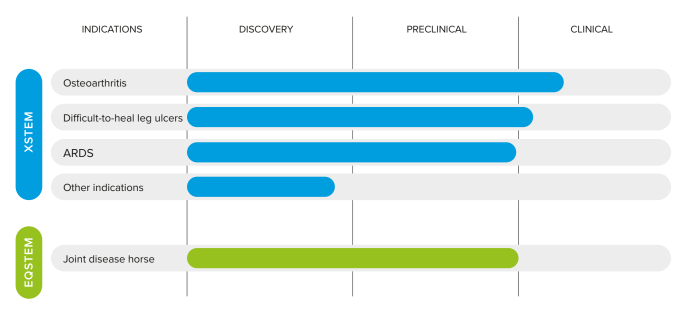
Commercialization strategy for XSTEM

The company's overall strategy is to take the stem cell projects to Proof of Concept, by clinical Phase I/Ila studies, and then enter into partnerships and commercial agreements for continued clinical development and global commercialization. Xintela is very active in business development and has ongoing dialogue with potential partners and licensees within the pharmaceutical industry.



A product platform for the treatment of several diseases

Xintela has two clinical studies ongoing with the stem cell product XSTEM, one in osteoarthritis and one in difficult-to-heal leg ulcers, as well as a project for the treatment of acute respiratory distress syndrome (ARDS) in preclinical phase. In addition, Xintela has carried out preclinical development with the stem cell product EQSTEM for the treatment of joint disease in horses.



In the knee osteoarthritis study all patients at the lowest dose level have completed the study

The clinical study (Phase I/IIa), conducted in Australia, is evaluating XSTEM for the treatment of patients with knee osteoarthritis. All patients in the dose escalation part of the study, 24 patients in total, have been dosed. Safety and efficacy readings will be evaluated every six months up to 18 months (the two lower dose levels) or 24 months (highest dose level) after treatment. The patients at the low and mid dose level have completed, or will shortly complete the study 18 months after treatment.

In the difficult-to-heal leg ulcers study, the first three patients have completed the study

The placebo-controlled clinical study (Phase I/IIa) is evaluating XSTEM for the treatment of difficult-to-heal venous leg ulcers. The first three patients have completed the study and recruitment of additional patients is ongoing at clinics in Sweden. A total of twelve patients will be recruited. Safety and efficacy readings are performed weekly for ten weeks as well as four months after treatment.

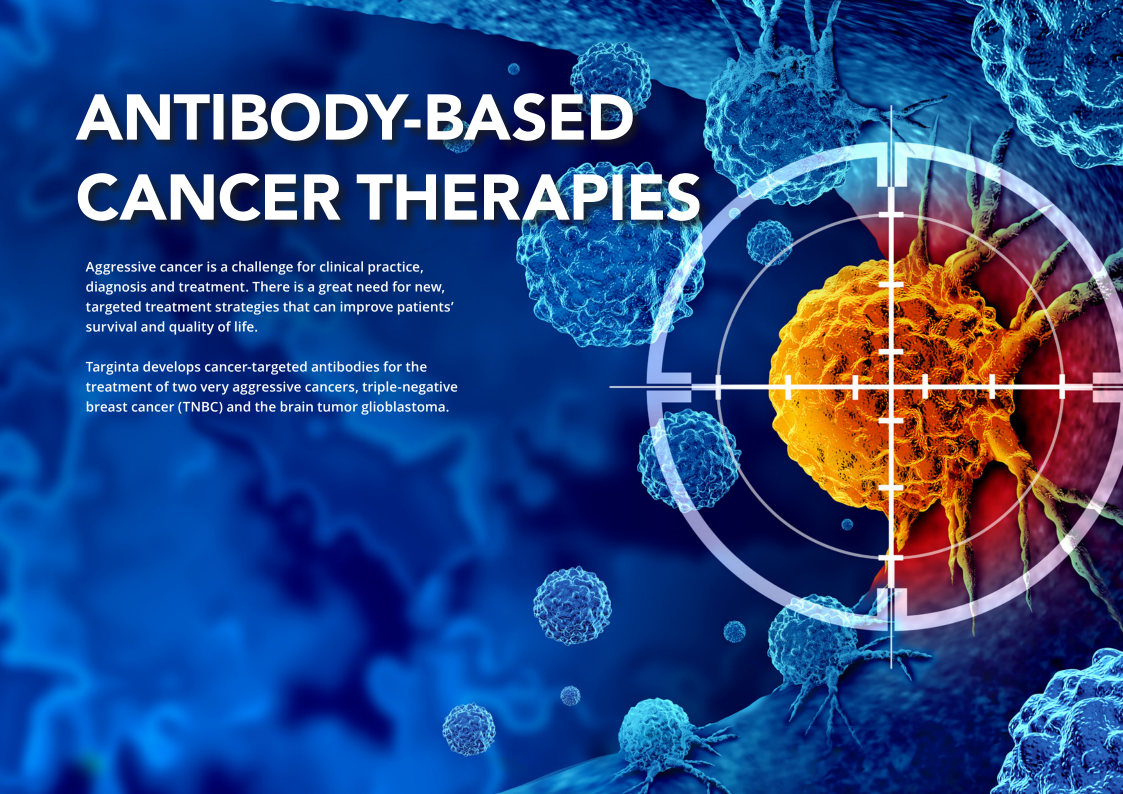
Preclinical study on ARDS show therapeutic effect with XSTEM

ARDS, acute respiratory distress syndrome, is a form of acute severe lung failure that can occur as a result of, for example, pneumonia, trauma or blood poisoning. The condition means that the lung function collapses. The mortality is high and there is currently no effective treatment for ARDS. Xintela has successfully conducted preclinical studies for the treatment of ARDS with XSTEM in collaboration with Skåne University Hospital and plans to carry out clinical development in collaboration with a partner.

EQSTEM® show disease modifying effect in preclinical horse models for osteoarthritis

Xintela has developed the stem cell product EQSTEM for the treatment of joint diseases in horses. Results from two preclinical studies in horses with post-traumatic osteoarthritis show disease modifying effect with reduces lameness and improved cartilage and bone structure. Xintela has ongoing partnering and licensing discussions with EQGen Biomedical (US) for clinical development and commercialization of EQSTEM.







TRIPLE-NEGATIVE BREAST CANCER

Triple-negative breast cancer, i.e. breast cancer that responds neither to hormone therapy nor to targeted treatment with HER2 antibodies, constitutes 10-15 percent of all breast cancer diagnoses and corresponds to approximately 300,000 new cases per year globally. It spreads and recurs to a greater extent and has a worse prognosis compared to other forms of breast cancer. The five-year survival rate for metastatic triple-negative breast cancer is about 12 percent. [5,6]

GLIOBLASTOMA

Glioblastoma (glioblastoma multiforme) is the most common and aggressive brain tumor in adults. Glioblastoma is characterized by the tumor cells rapidly spreading into the adjacent normal brain tissue, which contributes to the difficulty of removing the entire tumor without damaging the surrounding tissue. Glioblastoma cells are often resistant to both radiation and cytostatics and, as a result, the prognosis for patients is very poor. Approximately 55,000 people are estimated to be diagnosed with the disease annually in the 8 largest markets (USA, France, Germany, Italy, Spain, UK, Japan and China). [7,8,9]

New cancer target and effective First-in-Class antibodies

Cancer target with unique properties

Xintela's subsidiary Targinta is developing new targeted anti-body-based drugs (First-in-Class) for the treatment of aggressive cancer. The company has been founded on its own discovery that Xintela's stem cell marker, integrin $\alpha 10\beta 1$, is also expressed in aggressive cancers such as triple-negative breast cancer (TNBC) and the brain tumor glioblastoma.

The problem with most target molecules expressed in cancer is that the expression in normal tissues is relatively high. Integrin $\alpha 10\beta 1$ is unique in this respect as it expression is very limited in normal tissue, which reduces the risk of off-target side effects. Integrin $\alpha 10\beta 1$ is thus a very promising target molecule for the development of new and more selective cancer therapies.

Targinta has an extensive patent portfolio with several approved patents that protect both the company's antibody-based drug candidates as well as antibody treatment and diagnostics directed against the target molecule integrin $\alpha 10\beta 1$. The company can thus prevent competitors from developing integrin $\alpha 10\beta 1$ targeted antibodies for the treatment of aggressive cancers.

Targinta's candidate drugs

Targinta is developing two types of antibodies, TARG9 and TARG10, for the treatment of aggressive cancer. TARG9 is a so-called Antibody-Drug Conjugate (ADC) and is armed with a powerful toxin that has a killing effect on cancer cells. TARG9 has shown significant inhibitory effect on the growth

of glioblastoma tumors in preclinical models. TARG10 is a function-blocking antibody that slows down the growth and spread of cancer cells. TARG10 has in preclinical studies shown strong inhibitory effect on growth and metastasis of triplenegative breast cancer (TNBC). Targinta has a collaboration with Abzena Ltd for cell line development and initial production of TARG9 and TARG10.

Antibodies	Research	Preclinical	Clinical
TARG9			
TARG10			

Targinta positions itselfs in the ADC field

TARG9 was selected as the company's first candidate drug in the ADC area. This antibody has been developed with the latest ADC technology, which means a more powerful toxin that is well anchored to the antibodies as long as they circulate in the bloodstream, but which is released and activated when the antibody binds to and is taken up in cancer cells with integrin $\alpha 10\beta 1$ on the surface. The interest in toxin-armed antibodies, ADCs, has increased significantly in recent years and the area is considered one of the hottest in oncology. A large number of commercial agreements have been made even at the early preclinical stage.

Partnership for further development

Targinta is seeking financing and/or partnership for further development of its candidate drugs. One possible development route is to conduct clinical Phase 0 studies (microdosing) in cancer patients to show that the antibodies are able to reach and bind to the target molecule integrin $\alpha 10\beta 1$ on tumors and thus validate our target molecule and our candidates drugs. Positive results from the Phase 0 study will significantly reduce risk in the continued clinical development and thereby increase the attractiveness to potential partners and licensees.

The market for triple-negative breast cancer and glioblastoma

The global market value for the treatment of triple-negative breast cancer is estimated to be approximately USD 2.1 billion by 2028 and for the treatment of glioblastoma to approximately USD 1.4 billion by 2026. [10,11]

Commercialization strategy

Targinta's strategy is to enter into commercial agreements with the company's drug candidates during preclinical development and clinical Phase 0 studies to accelerate future clinical development and market approval. Drug candidates against new target molecules on cancer cells, so-called First-in-Class products, are very attractive to drug development companies due to the great need for new and more effective cancer treatments.





The Group Income statement in brief

Earnings

Operating loss for the first quarter amounted to TSEK -10,393 (-17,550) for the Group.

The costs for research and development account for the largest part of the group's costs and for the period April to June amounted to TSEK -7,891 (-14,074).

Market and sales costs for the quarter amounted to TSEK -812 (-1,275) for the Group.

Administrative expenses for the period amounted to TSEK -1,686 (-2,469) for the Group.

Loss before tax for the period amounted to TSEK -10,400 (-18,401) for the Group.

Under the heading "Tax on the period's results", SEK 409,000 is booked as revenue. This refers to the estimated size of the tax refund that will be paid out by the Australian Taxation Agency to Xindu, for parts of the costs the subsidiary Xindu has for the clinical studies during the period April to June 2024.

	Quart	Half	Full year		
	4/1/2024	4/1/2023	1/1/2024	1/1/2023	1/1/2023
(TSEK)	6/30/2024	6/30/2023	6/30/2024	6/30/2023	12/31/2023
Operating income					
Net sales	4	0	303	0	78
Cost of goods sold	0	0	0	0	0
Gross profit	4	0	303	0	78
Operating expenses					
Research and development costs	-7,891	-14,074	-16,241	-26,461	-46,239
Selling costs	-812	-1,275	-1,657	-2,338	-4,871
Administrative expenses	-1,686	-2,469	-3,592	-4,615	-7,919
Other operating income	-11	268	0	657	1,729
Other operating expenses	4	0	-8	0	-15
Operating loss	-10,393	-17,550	-21,196	-32,757	-57,237
Profit/loss from financial items					
Profit/loss from financial items Financial income	3	0	3	0	6
	3 -10	0 -851	3 -580	0 -1,487	6 -1,135
Financial income		-			
Financial income Financial expenses Loss before tax	-10 - 10,400	-851 -18,401	-580 -21,773	-1,487 -34,244	-1,135 - 58,367
Financial income Financial expenses	-10	-851	-580	-1,487	-1,135
Financial income Financial expenses Loss before tax	-10 - 10,400	-851 -18,401	-580 -21,773	-1,487 -34,244	-1,135 - 58,367



The Group Balance sheet in brief

Financial position

On June 30, 2024 the group's cash and cash equivalents amounted to TSEK 445 (697). Total assets amounted to TSEK 10,933 (13,991).

(TSEK)	6/30/2024	12/31/2023
ASSETS		
Fixed assets		
Intangible assets	56	195
Tangible assets	1,072	1,358
Total fixed assets	1,128	1,553
Current assets		
Tax assets	553	398
Accounts receivable	0	97
Tax receivable	4,427	4,347
Other receivables	1,739	3,066
Prepaid expenses	2,641	1,126
Cash and cash equivalents	445	7,809
Total current assets	9,806	16,843
TOTAL ASSETS	10,933	18,395
(TETIO	6/30/2024	12/31/2023
(TSEK)	6/30/2024	12/31/2023
EQUITY AND LIABILITIES Equity, the group		
Share capital	17,063	17,010
Other contributed capital	350,376	349,927
Reserve	-734	1,289
Balanced result incl. Profit for the year	-384,834	-363,846
Total equity	-18,129	4,380
Current liabilities		
Accounts payable	4,828	7,483
Tax liability	0	84
	20,052	4,214
Other liabilities	20,032	4,214
Other liabilities Accrued expenses and deferred income	4,182	2,234



The Group Cash flow statement in brief

Cash flow and investments

The group's cash flow for the period April to June 2024 was TSEK -9,940 (-2,056). Investments for the period amounted to TSEK 0 (0) for the Group.

	Quarter 2		Half	f year	Full year	
	4/1/2024	4/1/2023	1/1/2024	1/1/2023	1/1/2023	
(TSEK)	6/30/2024	6/30/2023	6/30/2024	6/30/2023	12/31/2023	
Operating activities						
Operating loss	-10,394	-17,550	-21,196	-32,757	-57,238	
Depreciation/amortisation	213	933	425	1,869	3,766	
Taxes	0	0	0	0	6,948	
Financial income	3	0	3	0	6	
Financial expenses	-10	-851	-580	-1,487	-1,135	
Cash flow from operating activities before changes in						
working capital	-10,188	-17,467	-21,348	-32,374	-47,652	
Changes in weaking conital						
Changes in working capital Increase/decrease in receivables	470	261	-831	1,011	-739	
Increase/decrease in current liabilities	-724	15.150	15,047	23.868	-4.725	
Changes in working capital	-254	15,411	14,216	24,879	-5,464	
	-	-,	, .	,	-, -	
Cash flow from operating activities	-10,442	-2,056	-7,132	-7,495	-53,116	
Investing activities						
Increase/decrease of tangible assets	0	0	0	0	-104	
Increase/decrease of intangible assets	0	0	0	0	0	
Increase/decrease of financial assets	0	0	0	0	0	
Cash flow from investing activities	0	0	0	0	-104	
Financing activities						
New share issue	0	0	0	0	45,216	
Equity, June 30, 2024	502	0	502	0	6,290	
Warrants, personnel	0	0	0	0	284	
Convertible	0	0	0	0	0	
Cash flow from financing activities	502	0	502	0	51,790	
Change in cash and cash equivalents	-9,940	-2,056	-6,630	-7,495	-1,430	
Cash and cash equivalents at the beginning of the period	10,409	2,415	7,809	8,343	8,343	
Conversion difference	-24	338	-734	-151	896	
Cash and cash equivalents at the end of the period	445	697	445	697	7,809	



The Group Change in equity in brief

		Other			
(TSEK)	Share capital	contributed capital	Reserves	Loss for the period	Total
Opening balance, January 1, 2023	9,227	305,920	393	-309,763	5,777
New share issue	7,150	39,241	0	0	46,391
New share issue, costs	0	-1,175	0	0	-1,175
New share issue, TO3	633	5,657	0	0	6,290
Warrants, personnel	0	284	0	0	284
Conversion difference	0	0	896	0	896
Loss for the period	0	0	0	-54,083	-54,083
Equity, December 31, 2023	17,010	349,927	1,289	-363,846	4,380
Opening balance, January 1, 2024	17,010	349,927	1,289	-363,846	4,380
Conversion difference	0	0	-2,023	1	-2,022
New share issue, TO3	53	449	0	0	502
Loss for the period	0	0	0	-20,988	-20,988
Equity, June 30, 2024	17,063	350,376	-734	-384,834	-18,129



The Parent Company Income statement in brief

Income

The parent company reports a net turnover of TSEK 4 (0) for the second quarter of the year. Other income amounted to TSEK 0 (268) and refer to contributions from Vinnova.

Earnings

Loss for the second quarter amounted to TSEK -8,933 (-10,614) for the Parent Company .

The costs for research and development account for the largest part of the Company's costs and amounted to TSEK -6,594 (-7,976) for the period April to June.

Market and sales costs for the quarter amounted to TSEK -813 (-1,173) for the Parent Company.

Administrative expenses for the period amounted to TSEK -1,530 (-1,733) for the Parent Company.

The financial income amounts to 373 (0) KSEK and refers to internal interest between Xintela and Xindu for the period April to June 2024.

Loss before tax and end of year dispositions for the period April to June amounted to TSEK -8,565 (-11,151) for the Parent Company.

	Quai	rter 2	Half	year	Full year	
	4/1/2024	4/1/2023	1/1/2024	1/1/2023	1/1/2023	
(TSEK)	6/30/2024	6/30/2023	6/30/2024	6/30/2023	12/31/2023	
Operating income						
Net sales	4	0	303	0	78	
Cost of goods sold	0	0	0	0	0	
Gross profit	4	0	303	0	78	
Operating expenses						
Research and development costs	-6,594	-7,976	-13,742	-16,051	-31,769	
Selling costs	-813	-1,173	-1,657	-2,121	-4,518	
Administrative expenses	-1,530	-1,733	-3,309	-3,122	-5,797	
Other operating income	0	268	0	643	1,656	
Other operating expenses	0	0	0	0	0	
Operating loss	-8,933	-10,614	-18,405	-20,651	-40,350	
Profit/loss from financial items						
Financial income	373	0	726	0	1,324	
Financial expenses	-5	-537	-575	-902	-908	
Loss before tax	-8,565	-11,151	-18,254	-21,553	-39,935	
Appropriations	0	0	0	0	-2,749	
Tax on loss for the year	0	0	0	0	0	
Loss for the period	-8,565	-11,151	-18,254	-21,553	-42,684	
Loss per share, SEK	-0.02	-0.04	-0.03	-0.07	-0.10	



The Parent Company Balance sheet in brief

Financial position

On June 30, 2024 the parent company's equity/assets ratio was 44 per cent (17) and equity amounted to TSEK 20,154 (7,247). The Parent company's cash and cash equivalents amounted to TSEK 269 (391). Total assets amounted to TSEK 45,778 (42,422).

(TSEK)	6/30/2024	12/31/2023
ASSETS		
Fixed assets		
Intangible assets	69	138
Tangible assets	696	897
Receivables from subsidiaries	28,367	23,852
Participations in subsidiaries	13,926	13,926
Total fixed assets	43,058	38,814
Current assets		
Tax assets	553	398
Accounts receivable	0	97
Tax receivable	121	63
Other receivables	766	879
Prepaid expenses	1,012	1,126
Cash and cash equivalents	269	7,092
Total current assets	2,720	9,655
TOTAL ASSETS	45,778	48,468
(TSEK)	6/30/2024	12/31/2023
EQUITY AND LIABILITIES		
Equity, parent company		
Share capital	17,063	17,010
Share premium reserve	350,376	349,927
Retained earnings	-329,031	-286,347
Loss for the period	-18,254	-42,684
Total equity	20,154	37,907
Current liabilities		
Accounts payable	3,602	4,640
Tax liability	0	(
		2.60=
Other liabilities	20,052	3,687
	20,052 1,970	
Other liabilities Accrued expenses and deferred income Total current liabilities		2,234
Accrued expenses and deferred income	1,970	3,687 2,234 10,561 48,468



The Parent Company Cash flow statement in brief

Cash flow and investments

The parent company's cash flow for the period April to June was TSEK -9,552 (-1,793) thousand. The investments for the period amounted to TSEK 1,577 (0) thousand.

	Qua	rter 2	Half	year	Full year	
	4/1/2024	4/1/2023	1/1/2024	1/1/2023	1/1/2023	
(TSEK)	6/30/2024	6/30/2023	6/30/2024	6/30/2023	12/31/2023	
Operating activities						
Operating loss	-8,933	-10,613	-18,405	-20,651	-40,350	
Depreciation/amortisation	135	856	269	1,713	3,454	
Financial income	373	0	726	0	1,324	
Financial expenses	-5	-537	-575	-902	-908	
Cash flow from operating activities before changes in						
working capital	-8,430	-10,294	-17,985	-19,840	-36,480	
Changes in working capital						
Increase/decrease in receivables	288	1,308	111	1,428	845	
Increase/decrease in current liabilities	-335	12,688	15,063	20,421	-4,194	
Changes in working capital	-47	13,996	15,174	21,849	-3,349	
Cash flow from operating activities	-8,477	3,702	-2,811	2,009	-39,829	
Investing activities						
Increase/decrease of tangible assets	0	0	0	0	-104	
Increase/decrease of receivables from subsidiaries	-1,577	-1,409	-4,514	-5,021	-5,419	
Shareholder contributions to subsidiaries	0	-4,087	0	-4,087	-4,087	
Cash flow from investing activities	-1,577	-5,495	-4,514	-9,107	-9,609	
Financing activities						
New share issue	0	0	0	0	45,216	
Equity, June 30, 2024	502	0	502	0	6,290	
Warrants, Personnel	0	0	0	0	284	
Group contribution paid	0	0	0	0	-2,749	
Cash flow from financing activities	502	0	502	0	49,041	
Change in cash and cash equivalents	-9,552	-1,793	-6,823	-7,098	-397	
Cash and cash equivalents at the beginning of the period	9,821	2,184	7,092	7,489	7,489	
Cash and cash equivalents at the end of the period	269	391	269	391	7,092	



The Parent Company Change in equity in brief

		Share	Retained	Loss for	
(TSEK)	Share capital	premium	earnings	the period	Total
Opening balance, January 1, 2023	9,227	280,920	-216,441	-44,906	28,800
Reversal of prior year's accruals	0	0	-44,906	44,906	0
Convertible	0	25,000	-25,000	0	0
New share issue	7,150	39,241	0	0	46,391
New share issue, costs	0	-1,175	0	0	-1,175
New share issue, TO3	633	5,657	0	0	6,290
Warrants, personnel	0	284	0	0	284
Loss for the period	0	0	0	-42,684	-42,684
Equity, December 31, 2023	17,010	349,927	-286,347	-42,684	37,907
Opening balance, January 1, 2024	17,010	349,927	-286,347	-42,684	37,907
Reversal of prior year's accruals	0	0	-42,684	42,684	0
New share issue, TO3	53	449	0	0	502
Loss for the period	0	0	0	-18,254	-18,254
Equity, June 30, 2024	17,063	350,376	-329,031	-18,254	20,154



Declaration by the Board of Directors and the CEO



Gregory Batcheller



Maarten de Château



Thomas Eldered



Lars Hedbys



Hans-Joachim Simons



Evy Lundgren-Åkerlund

The Board of Directors and the Chief Executive Officer certify that the interim report provides a true and fair view of the company's business, financial position, performance and describes material risks and uncertainties, to which the company is exposed.

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

Lund August 30, 2024

Gregory BatchellerChairman

Maarten de Château
Board member

Thomas ElderedBoard member

Lars Hedbys
Board member

Hans-Joachim Simons Evy Lundgren-Åkerlund
Board member CEO



Other information

The share

Xintela AB (publ) was listed on Nasdaq First North Growth Market in Stockholm on 22 March 2016 under the ticker symbol "XINT." First North Growth Market is an alternative marketplace, operated by an exchange within the NASDAQ OMX Group. Companies on First North Growth Market are not subject to the same rules as companies on the regulated main market. They are subject to a less regulated framework, adapted for small growth companies. A company listed on First North Growth Market may therefore entail a higher investment risk than a company listed on the main market. All companies listed on First North Growth Market have a Certified Adviser to oversee their compliance with the rules. The exchange assesses applications for admission to trading. Xintela's Certified Adviser on Nasdaq First North Growth Market is Carnegie Investment Bank AB.

On June 30, 2024, the number of shares was 568,760,509. The Company has only one class of shares. Each share carries identical rights to the Company's assets and earnings, and one vote at General Meetings.

Financial statements in accordance with K3

This report has been prepared in accordance with BFNAR 2012: 1 Annual Report and Consolidated Financial Statements (Q3) and the accounting principles are unchanged compared with those applied in the Annual Report for 2023. For complete accounting principles, see the Annual Report 2023.

	Jan - Jun 2024	Jan - Jun 2023	Jan - Dec 2023
No. of shares before full dilution	568,760,509	307,573,263	567,006,473
No. of shares after full dilution	704,809,082	307,573,263	704,809,082
Loss per share before full dilution	-0.10	-0.07	-0.10
Average no. of shares before full dilution	567,190,598	307,573,263	419,869,354
Average no. of shares after full dilution	703,239,171	307,573,263	557,671,963

Group accounts

The consolidated accounts include the companies in which the parent company directly or indirectly holds more than half of the votes for all shares, or otherwise has a controlling influence according to ÅRL 1:4. The company's earnings are included in the group's earnings from and including the acquisition date until it is divested. The financial statements of foreign subsidiaries have been recalculated according to the current rate method. All items in the balance sheet have been converted to the balance sheet exchange rate. All items in the income statement have been converted to average exchange rates during the financial year. Differences that arise are reported directly in equity.

Review by auditors

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

Financial calendar

Interim report Q3 2024: November 22, 2024 Interim report Q4 2024: February 28, 2025

Risks and uncertainties

Limited resources

Xintela is a small company with limited resources in terms of management, administration, and capital. The implementation of any major strategies requires optimization of the Company's resource appropriation. There is a risk that the Company's resources could be insufficient, and lead to financial and operational problems. The company's ability to continue its operations depends on the ongoing work with the company's financing being successful. Focused work is underway to secure the company's future financing and the Board's assessment is that we will successfully secure future financing needs.

Dependence on key individuals and employees

Xintela's success is based on the knowledge, experience, and creativity of a few specific individuals. The Company's future is dependent on being able to recruit qualified employees. The Company works hard to reduce this dependency by maintaining proper documentation of procedures and working methods.

Earning capacity and capital requirements

Drug development is both expensive and time-consuming. It may take longer than expected before the Company can generate a positive cash flow. To cover these costs, Xintela may need to raise new capital. There is no guarantee that such capital can be obtained on terms that are favorable to shareholders. Failure to generate sufficient profits may impact the Company's market value.

Sales risk

There is no certainty that the products developed by the Company will gain the market acceptance reflected in this interim report. The quantity of products sold may be lower, and the period required for market establishment may be longer, than the Company currently has reason to believe.



Sources:

- [1] Global Data 2018
- [2] Markets and Markets 2020
- [3] Markets and Markets: https://www.marketsandmarkets.com/Market-Reports/osteoarthritis-therapeutics-market-209565994.html
- [4] Fortune Business Insights: https://www.fortunebusinessinsights.com/venous-leg-ulcer-vlu-treatment-market-102370
- [5] https://www.cancer.org/cancer/breast-cancer/understanding-a-breast-cancer-diagnosis/types-of-breast-cancer/triple-negative.html#:~:tex-t=Triple%2Dnegative%20breast%20cancer%20(TNBC,of%20the%20protein%20called%20HER2
- [6] American Cancer Society https://www.cancer.org/cancer/breast-cancer/understanding-a-breast-cancer-diagnosis/types-of-breast-cancer/triple-negative.html
- [7] WebMD: https://www.webmd.com/cancer/brain-cancer/what-is-glioblastoma#1
- [8] American Association of Neurological Surgeons: https://www.aans.org/en/Patients/Neurosurgical-Conditions-and-Treatments/Glioblastoma-Multiforme
- [9] Global Data: Epidemiology and Market size Database
- [10] American Cancer Society https://www.cancer.org/cancer/breast-cancer/understanding-a-breast-cancer-diagnosis/types-of-breast-cancer/triple-negative.html
- [11] GlobalData: Glioblastoma Multiforme (GBM) Opportunity Analysis and Forecast to 2027



Xintela – for life in motion

Xintela develops stem cell-based treatments focusing on osteoarthritis and difficult-to-heal leg ulcers and, through its wholly owned subsidiary Targinta, targeted antibody-based treatments for aggressive cancer. The focus is on diseases that cause great suffering and lack effective medical treatment options.

Xintela has ongoing clinical studies with the stem cell product XSTEM for the treatment of knee osteoarthritis and difficult-to-heal venous leg ulcers. The goal is to show that stem cell treatment is safe, but also investigate XSTEM's ability to repair damaged articular cartilage and improve joint function and to heal venous leg ulcers, thereby reducing pain and suffering for patients. Preclinical studies have shown that XSTEM has regenerative properties.

Within oncology, tumor-targeting and armed antibodies are developed for aggressive cancers such as triple negative breast cancer and the brain tumor glioblastoma. Results from preclinical models have shown that the antibodies have an inhibitory effect on both the growth and metastasis of cancer cells. The drug candidates TARG9 and TARG10 are in preclinical development.

