



## DexTech Medical AB Interim report July 1- September 30, 2025

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By "Company" or "DexTech" is meant DexTech Medical AB with organization number 556664-6203.

### Summary of the first quarter (2025-07-01 – 2025-09-30)

- Net sales amounted to MSEK 0.0 (0.0)
- Operating profit/loss amounted to MSEK -1.3 (-1.3)
- Earnings per share\* SEK -0.07 (-0.06)
- Cash and cash equivalents at the end of the interim period amounted to MSEK 14.1 (14.7).

*\* Before and after dilution. Earnings per share: Profit for the period divided by the average number of shares 18,485,857. For the comparison period, the average number of shares was 18,485,857. Amounts in brackets refer to the corresponding period last year.*

### Comments from the CEO

The ongoing myeloma study is being conducted at Uddevalla Hospital, PI (principal investigator) Dr Dorota Knut and at Karolinska University Hospital, Huddinge, PI Dr Katarina Uttervall dept. Hematology/HERM. Biomarkers are analysed at the Central Laboratory, Karolinska University Hospital, NKS, Solna.

The primary objective is to confirm safety and tolerability and as a secondary objective to determine indications of treatment response. The patients who can be included in the study (inclusion criteria), must have relapsed/treatment-resistant disease and received 1-5 prior lines of therapy.

Treatment with ODX is given in a maximum of 7 doses, one dose every two weeks. Patients are divided into 3 dose groups, 3mg/kg, 6mg/kg and 9mg/kg, 4 patients in each group.

An addition to the study protocol enables follow-up of patients who responded to ODX treatment and refers to the time to new disease progression after completion of ODX treatment. No other anti-cancer treatment is given during the follow-up period.

Dose group 2 (6mg/kg) has been fully recruited. Two of the patients had progressive disease after completion of treatment and the other 2 patients are expected to complete their treatment in early December.

Follow-up of all patients who had so far achieved stable disease shows that the disease-inhibiting effect lasted at most just over six months without initiation of other cancer treatment.

A meeting of the DMC (Independent Data Monitoring Committee) was held in early October and approved the continuation to dose group 3. One patient for dose group 3 has been screened and received his first dose at the end of October.

The results so far, i.e. for patients with relapsed/treatment-resistant disease, must be seen as promising with a majority achieving stable disease, and, without serious side effects.

*Anders R Holmberg*

### Significant events during the interim period (July - September 2025)

On August 20, DexTech Medicals announced that Dose Group 2 (6mg/kg) in the myeloma study is expected to be fully recruited in August/September. Two patients in dose group 2 had progressive disease after completion of treatment. This means that so far, 67% of patients have responded positively to ODX treatment (transition from progressive disease to stable disease). No significant, ODX-related, side effects have been noted. Follow-up of all patients who have achieved stable disease is done to determine how long the disease-inhibiting effect persists after the ODX treatment has been

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discontinued. Follow-up of patients from dose group 1 shows that the disease-inhibiting effect continued, even after discontinuation of ODX. At most, the disease-inhibiting effect lasted just over six months without the initiation of other cancer treatment.

A meeting of the DMC (the independent data monitoring committee) was held in early October, which approved the continuation to dose group 3. One patient for dose group 3 was screened.

### Events after the end of the interim period

On October 16, DexTech Medicals announced that Dose Group 2 (6mg/kg) had been fully recruited and that DMC (the independent data monitoring committee) had given the OK for dose group 3. Two patients in dose group 2 had progressive disease after completion of treatment. The other 2 patients are expected to be completed in early December. The DMC approved continuation to dose group 3. No ODX related serious side effects had been noted. A new patient for dose group 3 had been screened.

Follow-up of all patients who have achieved stable disease is done to determine how long the disease-slowng effect persists after the ODX treatment has been discontinued, i.e., until new progress.

### Financial overview

|                                                | <i>First quarter</i>     |                          |
|------------------------------------------------|--------------------------|--------------------------|
|                                                | 2025-07-01<br>2025-09-30 | 2024-07-01<br>2024-09-30 |
| Net sales, TSEK                                | –                        | –                        |
| Profit after net financial items, SEK thousand | -1 256                   | -1 139                   |
| Earnings per share SEK*                        | -0,07                    | -0,06                    |
| * Before and after dilution                    |                          |                          |
|                                                | 2025-09-30               | 2025-06-30               |
| Cash and cash equivalents SEK thousand         | 14 075                   | 14 709                   |
| Balance sheet total TSEK                       | 24 253                   | 25 100                   |
| Equity ratio %                                 | 97                       | 99                       |

### Results, The first quarter, July – September 2025

#### Turnover and earnings

The company had no sales during the first quarter. Operating profit amounted to MSEK -1.3 (-1.3). During the first quarter, costs of MSEK 0.8 (0.6) were capitalized on drug development and patents. Operating expenses amounted to MSEK 2.2 (1.9) and consist of personnel costs MSEK 0.1 (0.2), other external expenses MSEK 1.0 (0.7) and depreciation costs MSEK 1.1 (0.9). Other external costs include hospital costs MSEK 0.4, costs for patents MSEK 0.1 and regulatory control MSEK 0.3 regarding the MM-study. Profit after tax amounted to MSEK -1.3 (-1.1).

#### Liquidity and financing

Cash and cash equivalents at the end of the interim period amounted to MSEK 14.1 (14.7).

Cash flow for the period amounted to MSEK -0.6 (-1.1).

Financing is done with equity. Equity at the end of the interim period amounted to MSEK 23.5 (24.8) million, corresponding to SEK 1.27 (1.34) per share. The equity/assets ratio was 97 (99) percent.

#### Working capital

In December 2021, DexTech carried out a rights issue that raised SEK 46.3 million before issue costs. Net proceeds of SEK 37.1 million to DexTech after issue costs of SEK 9.2 million. The Rights Issue 2021 ensured continued operations until the end of 2026. The goal is for license revenues to finance operations thereafter.

### Activities

DexTech Medical, org.nr 556664-6203 based in Stockholm, Sweden, develops drug candidates with applications in urological oncology, primarily prostate cancer. Operations commenced on August 9, 2004 and the Company was listed on the Spotlight Stock Market on June 19, 2014.

The company has a strong clinical foundation with valuable specialist expertise, from research laboratory and manufacturing to clinical oncology. Research and development is conducted cost-effectively through collaboration in a global network.

Based on a proprietary patented technology platform, GuaDex, the Company has developed four different drug candidates, OsteoDex, SomaDex, CatDex & GuaDex and a PSMA-binding conjugate, with patents/patent applications in several key markets.

- The company's lead candidate, *OsteoDex*, for the treatment of bone metastases in castration-resistant prostate cancer, CRPC, has after extensive preclinical studies shown strong tumoricidal effect and potent inhibition of bone degradation. After a successful phase I/IIa study where the results show high tolerability with only mild side effects and a clear effect in the highest dose group, a clinical phase IIb study (efficacy study) was initiated in the autumn of 2014. The full Phase IIb clinical study report (CSR) from the Phase IIb study for OsteoDex was completed in December 2018 and in June DexTech's Phase IIb study was completed with the receipt of 2-year follow-up results from the last patients. The study, which was conducted in Sweden, Finland, Estonia and Latvia, included 55 well-defined patients with castration-resistant prostate cancer with bone metastases (mCRPC).  
OsteoDex's efficacy for the treatment of multiple myeloma is ongoing (phase 1-2).
- *SomaDex* for the treatment of acromegaly, neuroendocrine tumors and palliative care in advanced prostate cancer. SomaDex is a drug candidate based on an endogenous hormone, somatostatin, for the treatment of acromegaly, neuroendocrine tumors and palliative treatment in advanced prostate cancer. SomaDex has undergone a Phase I clinical study (in Sweden/Finland) and a Phase II pilot study in Mexico. The studies showed that SomaDex has few and mild side effects (phase I) and has a palliative effect in advanced prostate cancer (pilot study).
- *CatDex & GuaDex*: *GuaDex* is the so-called technology platform and is a charge-modified dextran molecule with tumor-toxic properties (kills tumor cells) and is a development of CatDex.
- *PSMA-binding conjugate*, for target-specific treatment of mCRPC that overexpresses PSMA (prostate specific membrane antigen). The compound is based on the platform, GuaDex.

DexTech's goal is to out-license each drug candidate no later than after completion of the phase II study.

The technology platform, which can be likened to a "Lego box" with multiple opportunities to build new molecules, can also be out-licensed.

The following parameters have been important for DexTech's positive development to date:

- modified generics with well-documented mechanisms of action that are patented, entailing a lower risk in clinical development;
- early proof-of-concept data;
- strong clinical foundation with daily contact in clinical oncology;
- networked, academically and commercially;
- minimized fixed costs;
- Capital has been dedicated to drug development and patents.

### Prostate cancer

- Prostate cancer is the most common form of cancer in men in the Western world.
- About 25% of those who have prostate cancer develop incurable castration-resistant prostate cancer (CRPC) with bone metastases.
- Today, there are only a handful of approved drugs that can prolong the lives of these patients. All of these drugs have more or less serious side effects. Each of these drugs currently has, or is expected to achieve, sales of over USD 1 billion annually, so-called blockbusters.

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- After a limited time, CRPC becomes resistant to the respective drug, which means that the need for new complementary life-prolonging drugs is great.
- DexTech's lead candidate, OsteoDex, has the potential to become such a complementary drug.

### Fas IIb Studies

The original study protocol with ID ODX-002, was approved by the Swedish and Danish Medical Products Agencies in October 2014 (a placebo-controlled randomized multicenter phase II study) regarding OsteoDex for the treatment of castration-resistant prostate cancer with bone metastases (CRPC). On October 27, 2015, DexTech decided to change the study design and give all study patients active substance (OsteoDex). This is as a result of discussions with the Medical Products Agency in Uppsala and advice from "BigPharma". The study design was changed to active treatment for all patients. DexTech thus gains faster knowledge about the tumor-inhibiting effect in relation to dose, the efficacy parameter requested by prospective licensees. DexTech also responded to patients' requests for access to the active substance and thus avoid the risk of randomization to the placebo group. A decision to approve the new study protocol with ID ODX-003 was given by the Medical Products Agency in Uppsala on February 28, 2016.

The primary objective of the Phase II study was to document the efficacy of OsteoDex in the treatment of CRPC. The study included 55 well-defined CRPC patients. Patients were divided between three treatment arms (blinded distribution, 3 ascending dose levels of OsteoDex). The treatment was given for 5 months where OsteoDex was administered every two weeks. The study was conducted in Sweden (Norrländ University Hospital in Umeå, Södersjukhuset in Stockholm and University Hospital in Örebro), in Finland (Tampere University Hospital), in Estonia (East Tallin Central Hospital and Tartu University Hospital) and in Latvia (Riga East University Hospital and Daugavpils Regional Hospital). The first patient received his first treatment in September 2016 at Södersjukhuset in Stockholm.

In connection with these changes, the company chose to change the study organization by recruiting Crown-CRO Oy as GCP manager (good clinical practice) for the OsteoDex study. Crown-CRO Oy specializes in oncology studies in the Nordic and Baltic countries. Crown-CRO Oy replaces the company's former partner SynteractHCR.

In June 2018, the last patients in DexTech's phase IIb study for OsteoDex were completed. The work was then focused on the completion of the formal study report.

At the beginning of October 2018, DexTech was able to present the first results from the completed phase IIb study for Osteodex. The results obtained meet the primary objective of the protocol.

The Clinical Study Report (CSR) shows that 51 percent of patients completed the treatment (5 months, dose every two weeks). Of these, 52% showed stable disease (improved/unchanged) regarding bone metastasis. 35% of patients who completed treatment had a reduced tumour burden in the bones. The majority of the patients who had a reduced tumour burden in the skeleton had been treated with and no longer responded to two or more of the currently available drugs (docetaxel, cabazitaxel, abiraterone, enzalutamide, radium-223 dichloride) prior to enrolment in the study. This finding is of great importance for the continued clinical development of OsteoDex as the current patient group represents a significant so-called "unmet medical need". The results show that OsteoDex has a significant inhibitory effect on the vicious cycle ("vicious circle") in the skeleton, i.e. the biological process that drives this disease and thus also to shortened survival. More than 50% of patients showed marked decreases in the levels of markers related to bone metabolism and a particularly marked decrease was noted in 67% of patients for the marker CTX, which reflects bone degradation. The effect on this marker and other markers related to bone metastasis reflects the biological effect of the OsteoDex molecule. Tolerability was remarkably good with only few side effects. No patients needed to discontinue treatment due to adverse events and no OsteoDex-related serious adverse events (SAEs) were noted. The three dose arms in the protocol show equivalent treatment effect. The interpretation is that even the lower doses are sufficient to saturate the metastasis areas in the bones.

DexTech has previously reported promising follow-up results from the company's Phase IIb study of OsteoDex for the treatment of castration-resistant metastatic prostate cancer (mCRPC). Patients were then followed for survival for 24 months after completion of OsteoDex treatment. The results as of October 14, 2020 showed the following: of the patients who had stable (unchanged) bone metastasis disease at the end of treatment, 58% were alive, of the patients who had discontinued or discontinued treatment with progressive disease (progressive disease progression) were alive, and of the patients who had objective regression of bone metastases (reduction of existing bone metastases) at the end of treatment, 86% were alive. The results indicate prolonged survival after OsteoDex treatment.

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DexTech announced on June 12, 2020 that the randomized phase IIb study for the treatment of bone metastatic castration-resistant prostate cancer (mCRPC) was completed with the receipt of 2-year follow-up results from the last patients.

The study's primary endpoints regarding markers of bone metabolism had been well achieved. A clear majority of patients showed a reduction in their bone markers in blood from the given treatment with OsteoDex. The treatment was very well tolerated (few and mild side effects) and good disease-inhibiting effect was seen even in the lowest doses. Slowing and regression of the disease was also seen in patients whose disease progressed after treatment with several of the other available medications for castration-resistant prostate cancer.

The study's secondary endpoints include overall survival that was studied through 24 months of follow-up after completion of treatment. Of the patients who responded to treatment, with slowing or stabilization of the disease, the median survival had not yet been achieved ( $> 27$  months, thus a positive result), compared to 14 months for the other patients (significance,  $p < 0.05$ ). The 2-year survival rate after study entry was 65% for patients who responded to treatment, with disease slowing or stabilization, compared to 28% for other patients (significance,  $p < 0.05$ ).

The results of the study were very positive and show that OsteoDex effectively slows down the tumor disease. Data regarding overall survival should be seen as an indication, as these data, for natural reasons, need to be confirmed in a significantly larger, so-called Phase-III study.

None of the modern drugs are curative for castration-resistant prostate cancer and there is therefore a great need (unmet need) for new potent and tolerable drugs. OsteoDex has a clear potential to fill this need.

The continued clinical development of OsteoDex will be carried out by or together with a prospective licensee.

### Preclinical research

OsteoDex has a mechanism of action against cancer cells that is general and therefore also has other cancer diseases have been investigated as possible indications in addition to mCRPC, i.e. breast cancer, lung cancer and multiple myeloma.

#### Breast cancer

There are significant similarities between castration-resistant prostate cancer and advanced breast cancer in terms of propensity to metastasize to the bones. DexTech's preclinical studies conducted so far have clearly shown that OsteoDex has promising potential for the treatment of this form of cancer as well. The global market value of breast cancer drugs (total sales) is estimated to be USD 26 billion by 2025 (<https://www.precedenceresearch.com/breast-cancer-market>). The expanded preclinical program is part of the company's strategy to demonstrate OsteoDex's potential beyond the indication of castration-resistant prostate cancer.

#### Lung cancer

DexTech has previously announced data from preclinical studies regarding the effect of OsteoDex on the most common form of lung cancer, so-called non-small cell lung cancer (NSCLC). In conducted in vitro trials at Karolinska Institutet, OsteoDex shows a robust cell-killing effect in non-small cell lung cancer (NSCLC). The cell-killing effect was found to be completely on par with that seen in castration-resistant prostate cancer and breast cancer.

Lung cancer is divided into two main groups; non-small cell lung cancer and small cell lung cancer. Approximately 80 percent of all lung cancer cases are non-small cell lung cancer (NSCLC), which in turn is divided into several subgroups. Globally,  $>1.5$  million people are diagnosed with lung cancer every year, and the vast majority of these die from it. The lack of active and well-tolerated drugs is strikingly great.

#### Multipelt Myelom

DexTech has conducted an extensive preclinical program regarding the effect of OsteoDex on multiple myeloma. In vitro experiments conducted at Karolinska Institutet show that OsteoDex has a robust cell-killing effect on myeloma cells. The cell-killing effect has been shown to be superior to the standard drug Melphalan.



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MM is a form of blood cancer that starts in the bone marrow and causes the breakdown of the bones. The disease is incurable and the treatments that are currently available are used to, as far as possible, slow down the progression. The treatments often have severe side effects.

The company sees OsteoDex as very promising for the treatment of MM and **decided to conduct** a phase 1 study regarding the effect of OsteoDex on patients with multiple myeloma. This is based on OsteoDex's dual mechanism of action, inhibition of bone-degrading cells and tumor cell toxicity as well as with mild side effects, verified in clinical results.

On August 10, 2022, the Swedish Medical Products Agency approved and granted permission to conduct the phase 1 study regarding the effect of OsteoDex on patients with multiple myeloma.

DexTech announced on March 27, 2023 that the Company's Phase 1 study regarding the effect of OsteoDex on patients with multiple myeloma (MM) was initiated and recruitment of patients was initiated. The study initially included 20 patients and is being conducted at two hospitals in Sweden: Karolinska University Hospital Huddinge and Uddevalla Hospital.

The study is expected to be completed in Q4, 2025. The Principal Investigator (PI) is Dr Katarina Uttervall, MD, PhD, Department of Hematology/HERM, Karolinska University Hospital Huddinge. Analysis of main blood markers takes place at the Central Laboratory, Karolinska University Hospital Solna, NKS. In accordance with the treatment schedule, OsteoDex is given every two weeks. The inclusion criteria include adult MM patients with relapsed/treatment-resistant disease, who received 1–5 prior lines of therapy. The primary objective is to confirm safety and tolerability. The secondary objective is to determine treatment response, change in the level of disease-related biomarkers, and documentation of quality of life.

The ongoing myeloma study is being conducted at Uddevalla Hospital, Dr Dorota Knut, and Dr Katarina Uttervall dept. Haematology/HERM, Karolinska University Hospital, Huddinge, Katarina Uttervall is principal investigator (PI). Biomarkers are analysed at the Central Laboratory, Karolinska University Hospital, NKS, Solna. The primary objective is to confirm safety and tolerability and as a secondary objective to determine indications of treatment response. The patients who can be included in the study (inclusion criteria), must have relapsed/treatment-resistant disease and received 1-5 prior lines of therapy. Treatment with ODX is given in a maximum of 7 doses, one dose every two weeks. Patients are divided into 3 dose groups, 3mg/kg, 6mg/kg and 9mg/kg, 4 patients in each group. An earlier amendment (add-on study protocol) allows for follow-up of patients who have responded to ODX treatment and intends to record time to new disease progression after completion of ODX treatment.

On April 23, DexTech Medical announced strong follow-up results from the myeloma study. The follow-up showed remarkable results. At the first follow-up after completion of treatment (after 2 weeks), all patients in dose group 1 (3 mg/kg) had stable disease. The patients have then been followed, without other cancer treatment, until new progression of the disease (according to the amendment). Time to progress was 89 days, 59 days, 188 days and 39 days. None of the patients had any significant side effects from the ODX treatment. The results indicate that ODX is effective against multiple myeloma and that the slowing effect persists over time without other treatment. It should be noted when comparing time to progress with established myeloma drugs where the drug is given continuously until treatment resistance compared to ODX where the treatment is completed after only 7 doses. The time to progress after ODX treatment is nevertheless comparable to existing drugs against relapsed/treatment-resistant disease.

On August 20, DexTech Medicals announced that Dose Group 2 (6mg/kg) in the myeloma study is expected to be fully recruited in August/September. Two patients in dose group 2 had progressive disease after completion of treatment. This means that so far, 67% of patients have responded positively to the ODX treatment (transition from progressive disease to stable disease). No significant, ODX-related, side effects have been noted. Follow-up of all patients who have achieved stable disease is done to determine how long the disease-inhibiting effect persists after the ODX treatment has been discontinued. Follow-up of patients from dose group 1 shows that the disease-inhibiting effect continued, even after discontinuation of ODX. At most, the disease-inhibiting effect lasted just over six months without the initiation of other cancer treatment.

A meeting of the DMC (the independent data monitoring committee) was held in early October, which approved the continuation to dose group 3. One patient for dose group 3 has been screened and received his first dose at the end of October.

## PSMA-binding association

In June 2016, DexTech filed a patent application for important innovation regarding companion diagnostics and target-specific treatment of prostate cancer.

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It is well known that prostate cancer cells on their surface overexpress (present in larger quantities) the protein PSMA (prostate-specific membrane antigen, i.e. PSMA is found in greater quantities on the surface of the tumor cell). Extensive international research activity is underway to produce molecules that can bind specifically to PSMA and thus be used as carriers of cancer cell-killing substances (radioactive isotopes, cytostatics, etc.) for so-called target-specific treatment of prostate cancer. Such molecules (including antibodies to PSMA) have been produced in several laboratories, but there are still challenges regarding production for clinical use, shelf life, patent protection, regulatory requirements, etc.

DexTech has developed a new PSMA-binding compound with the help of the company's technology platform. The new substance has unique properties in that it has multiple PSMA-binding parts and can carry a larger load of cell-killing substances than has been possible with PSMA-specific molecules produced so far. The production of the new substance can be relatively easily adapted to the company's GMP platform (i.e. manufacturing that is approved for clinical use). The current patent application complements and strengthens the company's other patents. DexTech intends to seek a development partner for the pre-clinical/clinical development of the new drug candidate.

In June 2016, DexTech filed a patent application for an important innovation (patent family 4) regarding diagnosis (so-called companion diagnostics) and target-specific treatment of prostate cancer, PSMA. This application was approved for a patent in Finland in June 2018. In the autumn of 2017, DexTech filed an international patent application (so-called PCT application). Patents have now been granted in Europe, Israel, Canada and Japan.

### Patent

DexTech's patent portfolio includes four patent families and a new application for GMP manufacturing of OsteoDex (October 2023). Patents/applications provide strong protection of the Company's drug candidates and the Company's technology platform. The portfolio has a relevant geographical spread for DexTech. The Company's four patent families/patent applications are strongly related and each patent family is therefore relevant for all of the Company's drug candidates as well as for the platform, GuaDex. Patent applications are filed in countries where there is advanced pharmaceutical research and development and in countries that are larger markets for pharmaceutical products.

#### Patent family 1 - filed in 1999

Patent family 1 describes how the positively charged substance, CatDex, is selectively enriched in the tumor tissue, i.e. selectively relative to normal tissue.

Patent family 1 includes approved patents in Australia, Canada, the United States, and Europe (registered in Belgium, Switzerland, Germany, France, the United Kingdom, Italy, and Sweden). The patent was valid until October 12, 2019.

#### Patent family 2 filed in 2008

Patent family 2, the GuaDex patent, a further development of patent family 1, describes its tumor cell-killing properties against a number of different tumors, tumor cell cultures.

Patent family 2 includes approved patents in China, Finland, Israel, the United States, Mexico, Canada, Japan, and Europe (registered in Switzerland, Germany, France, the United Kingdom, Italy, and Sweden). The patent is valid until March 6, 2028.

#### Patent family 3 - filed in 2008

Patent family 3, the OsteoDex patent, is a GuaDex molecule with an additional component, a bisphosphonate, which has selectivity for the skeleton, i.e. where the metastasis is located.

Patent family 3 includes approved patents in China, Japan, Canada, Israel, Mexico, Brazil and Europe (registered in Switzerland, Germany, France, United Kingdom, Italy and Sweden). The patents are valid until April 7, 2028.

#### Patent family 4 - filed in 2016

In June 2016, DexTech filed a patent application for an important innovation (patent family 4) regarding diagnosis (so-called companion diagnostics) and target-specific treatment of prostate cancer, PSMA. This application was approved for a patent in Finland in June 2018. In the autumn of 2017, DexTech filed an international patent application (so-called PCT application). The application has been approved and patents have been granted in Europe, Israel, Canada and Japan. The patents are valid until 2036.

#### Patent family 5 - filed in 2023

The company's patent application to the European Patent Office regarding GMP manufacturing of OsteoDex (GMP= good manufacturing practice), was granted in May 2025 (unitary patent, approved in all EU countries). The patent provides patent protection until 2044. The patent is considered crucial for

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the continued clinical development of OsteoDex with future market exclusivity for a prospective partner.

### Prospects

DexTech's lead drug candidate OsteoDex has a unique dual mode of action, tumor-specific denaturation and inhibition of bone resorbing cells (osteoclasts). OsteoDex has been studied in a clinical phase II study with good results. There are significant similarities between bone metastases from mCRPC and Multiple Myeloma, such as growth site, bone degradation, and stimulation from osteoclasts.

These similarities have motivated DexTech's studies of OsteoDex's effects on Multiple Myeloma. In extensive preclinical studies conducted at Karolinska Institutet in Stockholm, the company has shown that OsteoDex has a very pronounced tumor cell-killing effect, which has been demonstrated on various Multiple Myeloma cell lines. OsteoDex shows strong efficacy even at low concentrations. Even compared to Melphalan, which is a proven standard drug for the treatment of multiple myeloma (MM), OsteoDex's effect is strikingly strong.

The project is now being developed into clinical research and a phase 1 study is underway. The study includes a maximum of 12 patients with relapsed treatment-resistant disease and is being conducted at 2 hospital centers, Karolinska University Hospital, Huddinge, and Uddevalla Hospital. The study is expected to be completed in Q4 2025.

The intention is that the study will provide proof of concept and thus further verify OsteoDex's value as a potential cancer drug. The market for the new indication is estimated to be twice as large as that for mCRPC. The global market value of multiple myeloma in 2020 was approximately USD 20 billion with a forecast value in 2026 of USD 31 billion (<https://www.fortunebusinessinsights.com/multiple-mye-loma-market-102693>).

The company's continued operations until the end of 2026 are secured.

The continued clinical development of OsteoDex with the indication mCRPC, i.e. phase III, is resource-intensive and requires a larger partner. Patent protection that provides market exclusivity after market approval is a crucial factor for a potential partner. Approval of the Company's new synthesis patent application would meet the requirement for long-term market exclusivity. The conditions for an approved patent application are considered to be good. The work on partnerships to ensure continued clinical development is ongoing.

### Organization

Anders R Holmberg is the CEO. The Board of Directors consists of Chairman of the Board Andreas Segerros and Board members Per-Olov Asplund, Peter Benson, Rolf Eriksson, and Svante Wadman.

### Share

The DexTech share was listed on the Spotlight Stock Market on June 19, 2014. Trading is done under the designation DEX.

The number of outstanding shares at the beginning and end of the interim period amounted to 18,485,857.

#### Implemented incentive program

At the Annual General Meeting of DexTech AB on October 28, 2022, it was resolved to implement an incentive program ("TO 2022/2025") for pre-selected key employees ("Option Holders") that gave the Option Holders the opportunity to subscribe for warrants in DexTech for the market value in a directed share issue. The Board of Directors of DexTech resolved on the allocation of TO 2022/2025. The subscription price for the warrants in the directed issue was set in accordance with the terms and conditions at SEK 0.13 per warrant. Warrant holders are entitled, during the period from and including 25 November 2025 up to and including 9 December 2025, or the earlier date that follows from the complete terms and conditions, to request subscription of one (1) new share in the company for each warrant at a subscription price of SEK 25. Amounts in excess of the quota value shall be added to the unrestricted share premium reserve. As a result of TO 2022/2025, the number of shares at full exercise will increase by 200,000 shares. This corresponds to, based on the Company's current share capital, a dilution of a maximum of approximately one percent of the shares and votes. The increase in the Company's share capital may, upon full exercise of the warrants, amount to a maximum of SEK 9,000.

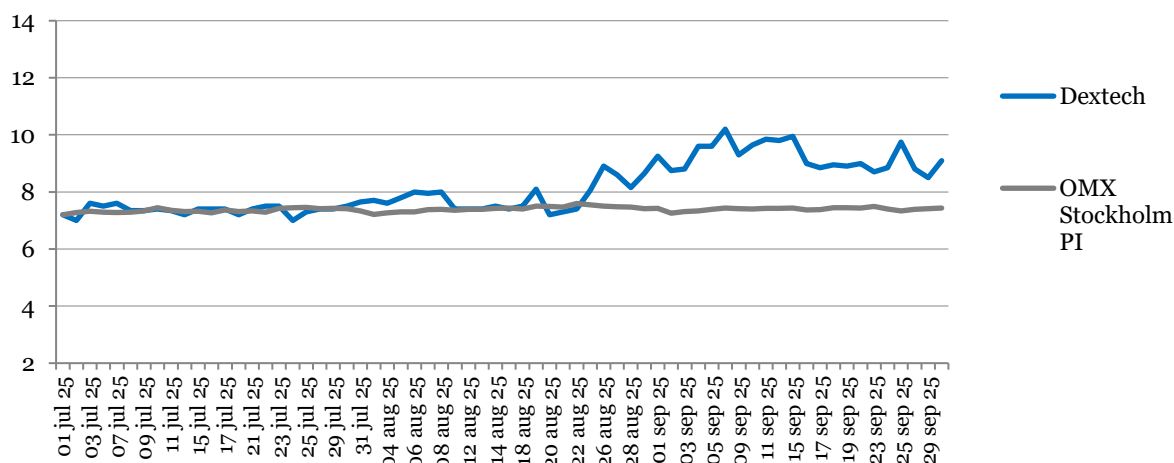
Reservations are made for such recalculations as a result of issues, etc., that may be made in accordance with the terms and conditions of the warrants.



## DexTech Medical

At the end of the interim period, the share price for DexTech Medical was SEK 9.10 and the reported equity per share was SEK 1.27. The market value amounted to MSEK 168.2. The number of shareholders was 1,106.

*Development of share price per share during the financial year 2025/2026*



## Related party transactions

Apart from remuneration to the CEO and CFO, there are no related party transactions to report.

## Accounting principles

This report has been prepared in accordance with the Annual Accounts Act and BFNAR 2012:1 Annual Report and Consolidated Financial Statements (C3). The accounting principles are unchanged compared with the last annual report.

## Financial information

|                            |                   |
|----------------------------|-------------------|
| AGM*                       | 4 november 2025   |
| Half-year report 2025/2026 | February 25, 2026 |
| Q3 - Report 2025/2026      | 5 May 2026        |
| Year-end report 2025/2026  | August 31, 2026   |

\* The Annual General Meeting will be held in Stockholm on November 4, 2025.

## Contacts

Anders Holmberg, CEO, +46 73 324 27 82  
Gösta Lundgren, CFO, +46 70 710 47 88

This information is information that DexTech Medical AB is obliged to make public pursuant to the EU Market Abuse Regulation. The information was submitted for publication, through the agency of the contact persons set out above, on November 3, 2025.

Stockholm, November 3, 2025  
DexTech Medical AB

## Board

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

DexTech Medical AB  
Dag Hammarskjölds Väg 34A, Uppsala  
Mailing address:  
Box 389, 752 23 Uppsala  
www.dextechmedical.com

## SUMMARY OF INCOME STATEMENTS

|                                                               | <i>First quarter</i> |               |
|---------------------------------------------------------------|----------------------|---------------|
|                                                               | 2025-07-01           | 2024-07-01    |
| KSEK                                                          | 2025-09-30           | 2024-09-30    |
| Net sales                                                     | -                    | -             |
| Work performed by the company for its own use and capitalized | 827                  | 585           |
| Operating expenses                                            | -2 170               | -1 878        |
| <b>Operating profit/loss</b>                                  | <b>-1 343</b>        | <b>-1 293</b> |
| Net financial profit/loss                                     | 87                   | 154           |
| <b>Profit/loss before tax</b>                                 | <b>-1 256</b>        | <b>-1 139</b> |
| Tax                                                           | -                    | -             |
| <b>Net profit/loss</b>                                        | <b>-1 256</b>        | <b>-1 139</b> |
| Earnings per share, SEK *                                     | -0,07                | -0,06         |
| Average number of shares, thousand *                          | 18 485 857           | 18 485 857    |

## SUMMARY BALANCE SHEETS

| KSEK                                | 2025-09-30    | 2025-06-30    |
|-------------------------------------|---------------|---------------|
| <b>Assets</b>                       |               |               |
| Intangible assets                   | 9 680         | 9 917         |
| Financial assets                    | 1             | 1             |
| Receivables                         | 497           | 473           |
| Cash and cash equivalents           | 14 075        | 14 709        |
| <b>Total assets</b>                 | <b>24 253</b> | <b>25 100</b> |
| <b>Equity and liabilities</b>       |               |               |
| Equity                              | 24 763        | 29 608        |
| Current liabilities                 | 337           | 980           |
| <b>Total equity and liabilities</b> | <b>25 100</b> | <b>30 588</b> |

## SUMMARY CASH FLOW ANALYSIS

|                                                           | 2025-07-01    | 2024-07-01    |
|-----------------------------------------------------------|---------------|---------------|
| KSEK                                                      | 2025-09-30    | 2024-09-30    |
| Cash flow from operating activities                       | 193           | -466          |
| Cash flow from investing activities                       | -827          | -585          |
| <b>Cash flow for the period</b>                           | <b>-634</b>   | <b>-1 051</b> |
| Cash and cash equivalents at the beginning of the period  | 14 709        | 19 043        |
| <b>Cash and cash equivalents at the end of the period</b> | <b>14 075</b> | <b>17 991</b> |