

## Xintela completes dosing of XSTEM second dose level in knee osteoarthritis clinical study

**Xintela's first-in-human study (Phase I/IIa) for the treatment of knee osteoarthritis, being conducted in Australia, is testing 3 different dose levels of the stem cell product XSTEM®. All patients at the second dose level have now been dosed. XSTEM, which consists of allogeneic (donated) integrin  $\alpha 10\beta 1$ -selected mesenchymal stem cells, is developed and manufactured by Xintela.**

Patients with moderate knee osteoarthritis (grade II-III) receive an injection of XSTEM into the knee joint. Three different dose levels are being evaluated in up to 54 patients and each patient will be followed for 18 months with efficacy assessments every six months. The primary goal is to show that XSTEM is safe, and also to investigate preliminary efficacy signals, such as reduced breakdown of joint cartilage, regeneration of damaged cartilage, and improved joint function. Safety data and early efficacy results are expected in 2023.

"The first 16 patients with knee osteoarthritis have now been injected with XSTEM. We are pleased that the study is progressing well and are looking forward to initiate dosing at the third and last dose level after the study's Safety Review Committee has evaluated the second dose level's one-month data", says Camilla Wennersten, Director Clinical Development.

### Contacts

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## About Xintela

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Xintela develops medical products in stem cell therapy and targeted cancer therapy based on the Company's cell surface marker integrin  $\alpha 10\beta 1$  which is found on mesenchymal stem cells and on certain aggressive cancer cells. The stem cell marker is used to select and quality-assure the patent-protected stem cell product XSTEM®, which is in clinical development for treatment of knee osteoarthritis and difficult-to-heal leg ulcers. The company produces XSTEM for the clinical studies in its GMP-approved manufacturing facility. In cancer therapy, which is run by the wholly owned subsidiary Targinta AB, therapeutic antibodies, targeting integrin  $\alpha 10\beta 1$  (First-in-Class) are being developed for the treatment of triple-negative breast cancer and the brain tumor glioblastoma. Xintela conducts its business at Medicon Village in Lund, Sweden, and is listed on Nasdaq First North Growth Market Stockholm since 22 March 2016. Xintela's Certified Adviser at Nasdaq First North Growth Market is Erik Penser Bank AB, +46 8-463 80 00, [certifiedadviser@penser.se](mailto:certifiedadviser@penser.se).

## Attachments

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