

26 June 2026

Patient recruitment completed for Queensland Children Hospital clinical study in patients with unilateral cleft lip

Highlights

- Osteopore is delighted to announce that the recruitment of all 5 patients to the study has been completed.
- As previously announced, Osteopore and Queensland Children's Hospital (QCH) have launched and commenced a clinical trial for paranasal augmentation in 5 paediatric patients with unilateral (one-sided) cleft lip – which is the augmentation of the nose and surrounding upper jaw structure.
- Cleft lip and palate affects approximately 1 in 1,000 births globally¹, making it one of the most prevalent congenital conditions worldwide.
- The global cleft lip and palate repair market was valued at approximately USD 1.2b (~AUD 1.7b) in 2024 and is forecast to reach USD 1.9b (~AUD 2.7b) by 2033, growing at a CAGR of 5.5% ².

Australian-Singaporean regenerative medicine company **Osteopore Limited** (ASX: **OSX**; **Osteopore** or **the Company**) – a global leader in 3D-printed biomimetic and bioresorbable implants – is delighted to announce that the recruitment of all 5 patients to the clinical trial for paranasal augmentation with unilateral (one-sided) cleft lip has been completed.

¹ DOI: 10.15761/OHNS.1000246

² <https://www.verifiedmarketreports.com/product/cleft-lip-and-palate-repair-market/>



As previously announced on 2 June 2026³, QCH will conduct a single-arm feasibility trial, with the recruitment of up to 5 patients expected by end of 2026. Patient follow-up will continue for 12 months after surgery.

The trial, led by Dr. Yun Phua, a Plastic and Reconstructive Surgeon based at QCH and Mater Hospital, is the second paediatric study led by Dr. Phua – the first study was for the treatment of temporal hollowing, announced on 23 May 2025⁴.

The trial utilises Osteopore's medical-grade polycaprolactone-tricalciumphosphate (mPCL-TCP) scaffolds, manufactured using additive manufacturing (3D printing) to produce patient-specific implants precisely matched to each patient's anatomy.

The scaffolds are 3D printed from a bioresorbable composite (PCL-TCP) that gradually degrades as the body regenerates bone, mimicking the mechanical properties of trabecular bone throughout the healing process. Implants are designed using each patient's CT imaging data and manufactured under Osteopore's established Patient Specific Implant (PSI) workflow, already in clinical use at QCH.

The current standard of care for this condition — cancellous bone grafting harvested from the hip — is associated with poor long-term graft retention and unpredictable outcomes for nasal form.

Osteopore's scaffold is designed to replace this graft entirely, providing a structurally accurate, reproducible alternative that supports bone regeneration and nasal base support without adding surgical time. This represents a meaningful clinical advancement over existing practice.

³ ASX announcement, "OSX & QCH launch trial for nasoalveolar augment in children", 2 June 2026.

⁴ ASX announcement, "OSX partners QCH in trial for temporal hollowing in children", 23 May 2025.



Cleft lip and palate affects approximately 1 in 1,000 births globally⁵, making it one of the most prevalent congenital conditions worldwide. Patients typically undergo multiple reconstructive procedures throughout childhood and adolescence.

The global cleft lip and palate repair market was valued at approximately USD 1.2b (~AUD 1.7b) in 2024 and is forecast to reach USD 1.9b (~AUD 2.7b) by 2033, growing at a CAGR of 5.5%⁶. Growth is being driven by rising surgical volumes, increasing awareness of treatment options, and the adoption of advanced biomaterials and surgical techniques.

This trial, conducted at one of Australia's leading children's hospitals, provides a strong clinical and commercial foundation for broader market adoption.

By delivering a more predictable and durable outcome, this technology has the potential to reduce revision surgeries, improve facial symmetry, and meaningfully enhance quality of life for patients — outcomes being measured using the validated Cleft-Q patient-reported outcome tool.

ENDS

This announcement has been authorised for release to the ASX by the Board of Osteopore Limited.

For more information, please contact:

Dr Yujing Lim
Chief Executive Officer
Osteopore Limited
E: lim_yujing@osteopore.com

Mark Leong
Executive Chairman
Osteopore Limited
E: mark_leong@osteopore.com

⁵ DOI: 10.15761/OHNS.1000246

⁶ <https://www.verifiedmarketreports.com/product/cleft-lip-and-palate-repair-market/>



About Osteopore Limited

Osteopore Ltd. is a global medical technology company founded in Singapore and listed in Australia that commercialises products designed to enable natural bone healing across multiple therapeutic areas. Osteopore's patented technology fabricates specific micro-structured scaffolds for bone regeneration through 3D printing and bioresorbable material.

Osteopore's patent-protected scaffolds are manufactured using a proprietary manufacturing technique with a polymer that naturally dissolves over time to only allow natural and healthy bone tissue, significantly reducing the post-surgery complications commonly associated with permanent bone implants. Our 3D printing technology is unique to Osteopore.

Forward-Looking Statements

Some of the statements appearing in this announcement may be similar to forward-looking statements. You should be aware that such statements are only predictions and are subject to inherent risks and uncertainties. Those risks and uncertainties include factors and risks specific to the industries in which the Company operates and proposes to operate as well as general economic conditions, prevailing exchange rates and interest rates and conditions in the financial markets, among other things.

Actual events or results may differ materially from the events or results expressed or implied in any forward-looking statement. No forward-looking statement is a guarantee or representation as to future performance or any other future matters, which will be influenced by several factors and subject to various uncertainties and contingencies, many of which will be outside the Company's control. The Company does not undertake any obligation to update publicly or release any revisions to these forward-looking statements to reflect events or circumstances after today's date or to reflect the occurrence of unanticipated events.

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