

SteriBridge™

SteriBridge™ allograft implants are composed of demineralized cortical bone fibers entangled and performed into a shape for use in orthopedic surgical applications. The allograft utilizes the inherent osteoconductive and osteoinductive potential of demineralized bone to provide a scaffold and facilitate bone formation, respectively

Key Features

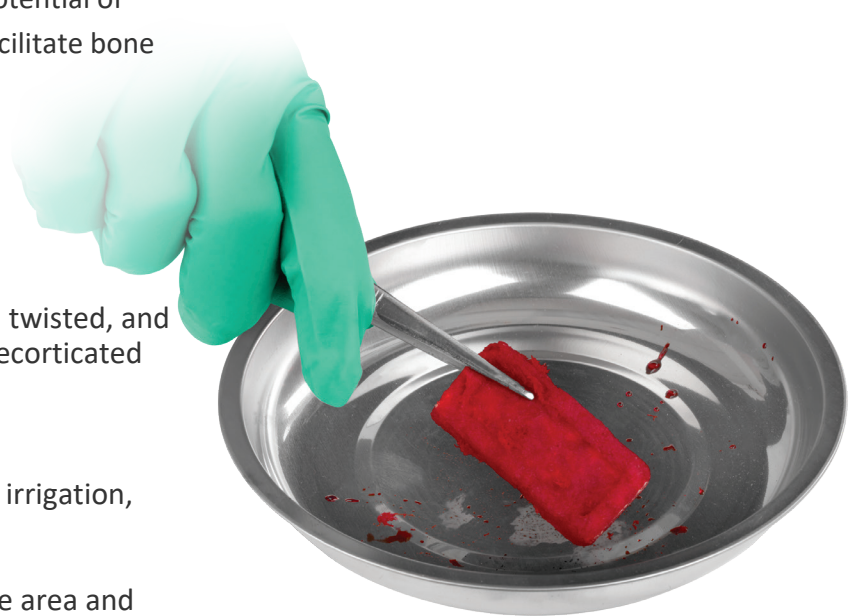
- **Flexible Handling**
Once hydrated, grafts are able to be folded, twisted, and molded to conform to the defect site and decorticated bone,
- **Migration Resistance**
grafts are preformed to stay in place during irrigation,
- **Osteoconductive**
Grafts provide a scaffold with a large surface area and porosity designed to help promote cellular attachment and proliferation.

Safety

Grafts are processed and sterilized through their proprietary GraftShield® process for improved allograft safety. The process meets all FDA human tissue regulations (CFR 1271) and applicable standards.

Ordering Information

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101.05251

SteriBridge Boat 50x25x7mm



The SteriBridge HSA™ Allograft implants are regulated by the FDA under 21 CFR Part 1271 Human Cells, Tissues and Cellular and Tissue-Based Products (HCT/Ps). Innovasis, Inc. is registered with the FDA for tissue storage and distribution. Bone Bank Allografts is registered with the FDA for tissue processing and is accredited by the American Association of Tissue Banks (AATB).

Marketed and Represented by: Innovasis, Inc. 614 E 3900 S Salt Lake City, UT 84107

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