

GraftNet™ XL Bone Collection Device

Surgical Technique for Spine Applications



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Simplified Autologous Bone Graft Collection

The GraftNet XL device is designed to maximize autologous bone graft collection and uses a large-volume mesh collection filter to easily integrate the patient's own cells into the procedure.



Technique



Attach the GraftNet XL tubing to the desired handpiece or Frasier tip suction device.

Attach outflow tubing to the outflow adapter of the GraftNet XL device.



Continue with resection as desired. Collected autologous collected tissue will be captured in the inner filter of the GraftNet XL device.

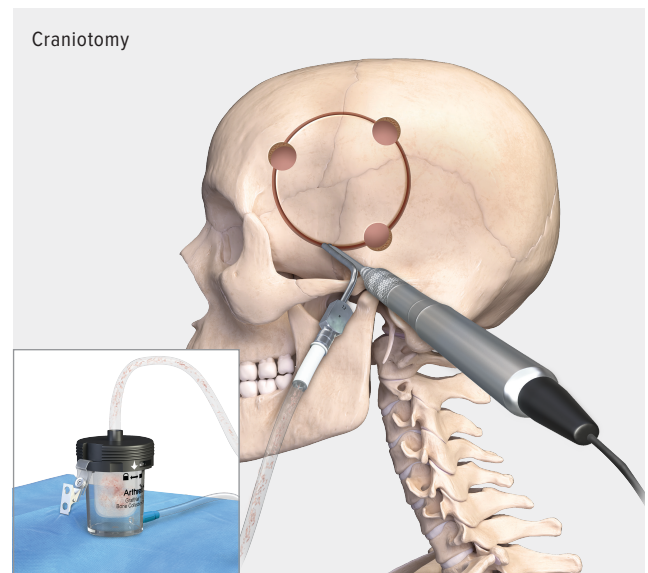
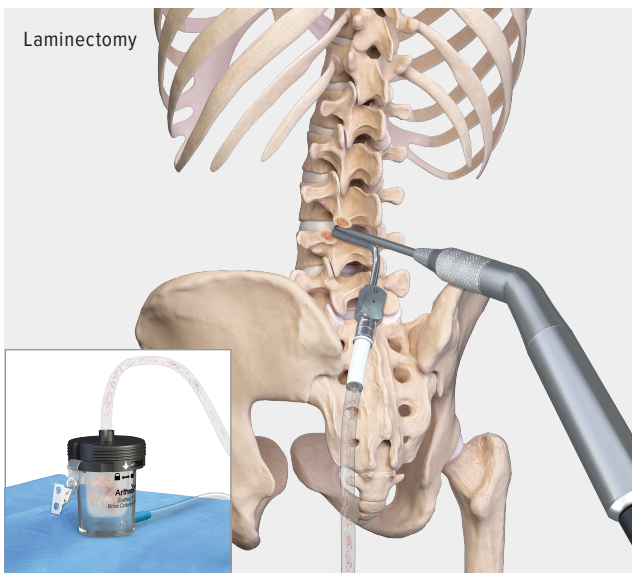


Once bone collection is complete, disconnect the device from suction and twist the cap from the locked to the unlocked position and remove the inner filter.

Excess fluid can be removed from the collected bone material by squeezing the mesh filter or using a sterile absorbent pad.

Invert the filter to remove the autologous bone and place into a sterile basin. A Freer elevator can be used to remove additional autologous bone from the filter.

Potential Applications



The GraftNet™ XL device can be used to collect bone during any procedure where there is a desire to capture and reintegrate autologous bone into the reconstruction or repair, such as a laminectomy or craniotomy. Universal adaptors at inflow and outflow ensure the GraftNet XL device can be quickly incorporated into any spine or neurosurgery procedure.

Ordering Information

Product description	Item number
GraftNet™ XL bone collection device	ABS-1052



This description of technique is provided as an educational tool and clinical aid to assist properly licensed medical professionals in the usage of specific Arthrex products. As part of this professional usage, the medical professional must use their professional judgment in making any final determinations in product usage and technique. In doing so, the medical professional should rely on their own training and experience, and should conduct a thorough review of pertinent medical literature and the product's directions for use. Postoperative management is patient-specific and dependent on the treating professional's assessment. Individual results will vary and not all patients will experience the same postoperative activity level or outcomes.



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US patent information

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