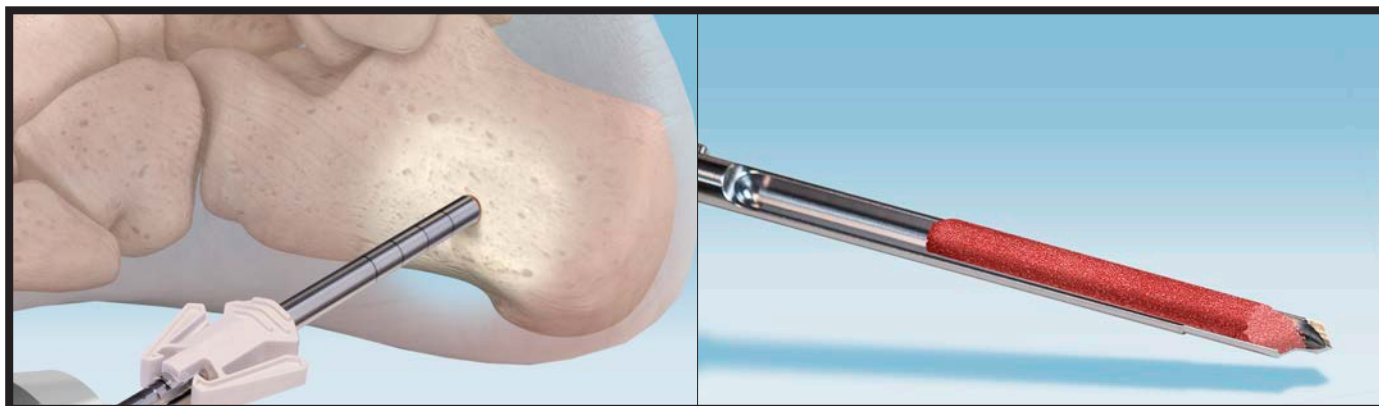


MINIMALLY INVASIVE

CoLink® Bone Graft Harvester



Delivered in sterile, single-use kit for rapid harvest

Morselizes bone to facilitate void filling

The In2Bones CoLink® Bone Graft Harvester is a single-use, pre-assembled bone graft harvesting device provided sterile that may be used to harvest bone from various sites in the body including the calcaneus, iliac crest, proximal tibia, distal tibia, distal radius, and distal femur. The Bone Graft Harvester is intended to morselize cancellous bone for enhanced bone healing in fusion and fracture stabilization procedures. The Bone Graft Harvester is available in two sizes, 6mm and 8mm outer diameter.

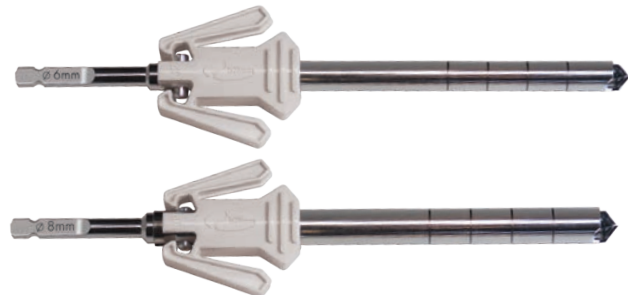


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CoLink® Bone Graft Harvester

CoLink® Bone Graft Harvester / Sterile, Single-use

CAT NO.	DESCRIPTION	SIZE
G05 S1006	CoLink® Bone Graft Harvester	6mm
G05 S1008	CoLink® Bone Graft Harvester	8mm



INDICATIONS

The CoLink® Bone Graft Harvester is a general single-use, sterile instrument. The In2Bones single-use, sterile instruments are indicated for assisting surgeons in the manipulation of soft tissue and bone, during surgical procedures.

CONTRAINDICATIONS

Contraindications for the instruments include general contraindications for orthopedic surgery such as:

- Bone, musculature, tendons, or adjacent soft tissue compromised by disease, infection or prior instrumentation;
- Metal allergy;
- Active infection in the joint.
- Skeletal immaturity.
- Diabetes.

RECOMMENDATION

It is recommended to carefully read the instructions for use available in the package insert.

DEVICES

CE Classification (Directive MDD 93/42/EC)
Single use instruments and instruments connected to a power driver: Class CE IIa - CE2797

REIMBURSEMENT

Reimbursement may vary from countries to countries. Check with local authorities.

MANUFACTURER

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EC REP

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Prinsessegracht 20
2514 AP The Hague
The Netherlands

DOCUMENT

Reference: BR-DIG-COLINKBGH-EN-012021

All content contained herein is furnished for informational purposes only. In2Bones does not recommend a particular surgical product or procedure suitable for all patients. Each surgeon must evaluate the appropriateness of a device and corresponding techniques based on medical training, clinical judgment and surgical experience. The proper surgical technique and/or procedure are the responsibility of the medical professional. Indications, contraindications, warnings, and precautions are listed in the implant package insert and should be reviewed carefully by the physician and operating room personnel prior to any proposed procedure. Availability of these products might vary from a given country or region to another as a result of specific local regulatory approval or clearance requirements for sale in such country or region.

CAUTION: Federal law (USA) restricts this device to sale and use by, or on the order of a physician.



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