



RETRIEVE[®]

Bone Graft Harvest System



Our mission is to deliver cutting-edge technology, research, and innovative solutions to promote healing in patients with musculoskeletal disorders.

Life moves us 

The Surgical Technique shown is for illustrative purposes only. The technique(s) actually employed in each case always depends on the medical judgment of the surgeon exercised before and during surgery as to the best mode of treatment for each patient. Additionally, as instruments may occasionally be updated, the instruments depicted in this Surgical Technique may not be exactly the same as the instruments currently available. Please consult with your sales representative or contact Globus directly for more information.

SURGICAL TECHNIQUE GUIDE

RETRIEVE[®]

Overview	4
Instrument Overview	6
Surgical Technique	
1. Preparation and Guide Wire Placement	8
2. Driver Assembly and Insertion	9
3. Driver Rotation and Removal	11
4. Disassembly	13
Optional Technique: Using the Driver (691.020)	14
RETRIEVE [®] Bone Graft Harvest System Sets	18
Important Information	20

RETRIEVE®

Bone Graft Harvest System

The RETRIEVE® Bone Graft Harvest System is a minimally invasive option for harvesting autogenous bone from the iliac crest. These cores can be used for general bone grafting or to fill cervical and lumbar interbody fusion devices.

The system removes the graft through an incision slightly larger than that required for collecting bone marrow aspirate. Due to less tissue damage and minimal resection, this can help to reduce postoperative pain. Additionally, ease of use of the instruments may help reduce the time typically required to harvest iliac crest bone.

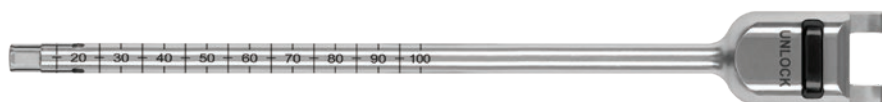


The harvested autogenous bone cores can be used alone as a general bone graft or to fill interbody fusion devices. For the posterolateral spine, autograft cores can be mixed with bone graft extenders and bone void fillers to enhance the potential for fusion.

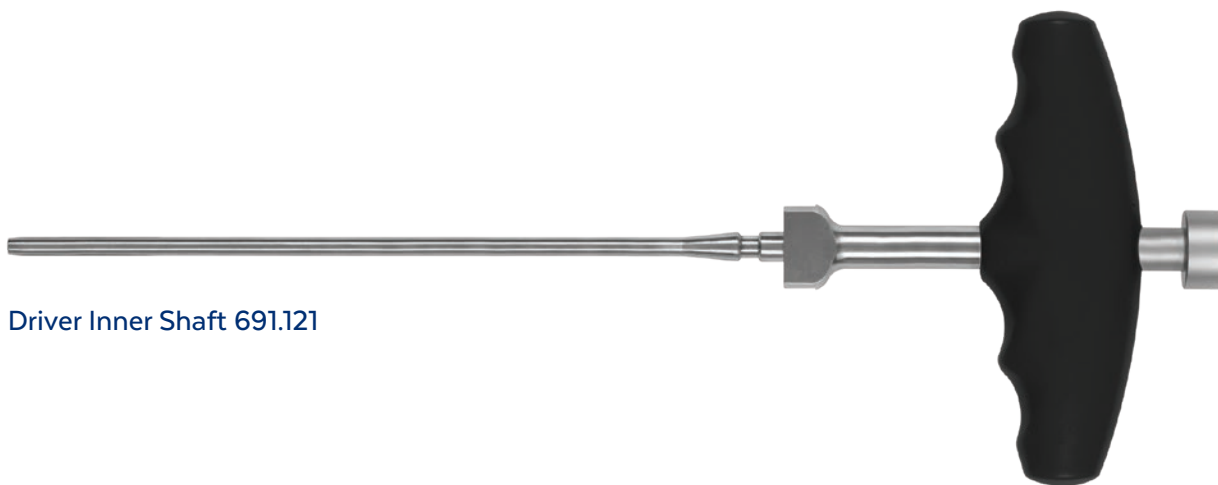


INSTRUMENT OVERVIEW

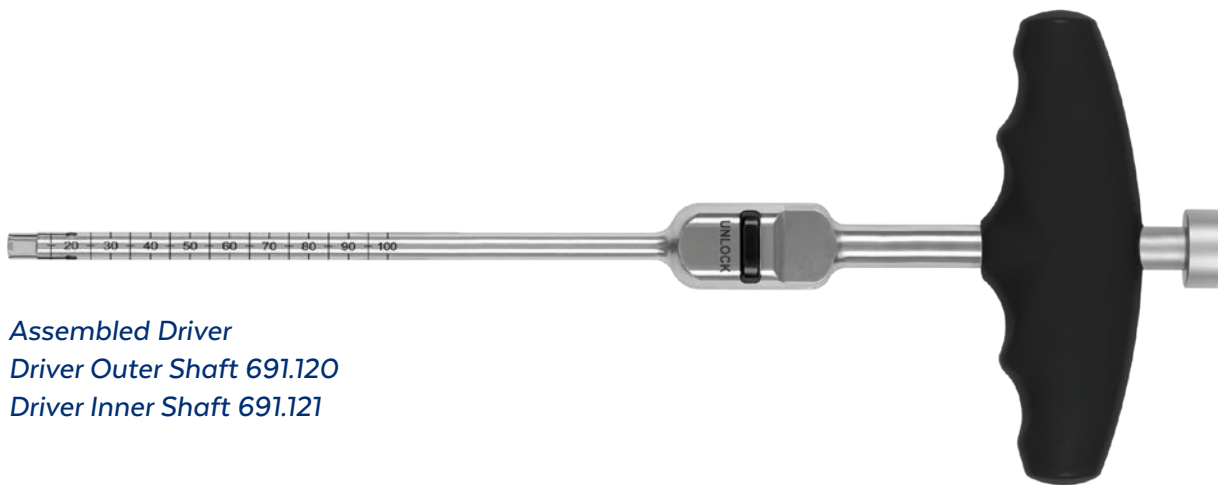
REUSABLE INSTRUMENTS



Driver Outer Shaft 691.120



Driver Inner Shaft 691.121



Assembled Driver
Driver Outer Shaft 691.120
Driver Inner Shaft 691.121



Driver 691.020



Hammer 691.024



Graft Remover 691.025



Slide Hammer, Small 622.410

DISPOSABLE INSTRUMENTS - STERILE, SINGLE-USE



RETRIEVE® Tip, 7x30mm 691.002S



RETRIEVE® Tip, 12x30mm 691.004S



RETRIEVE® Tip, 10x30mm 691.003S



RETRIEVE® Tip, 15x30mm 691.005S



Guide Wire 691.022S

SURGICAL TECHNIQUE

RETRIEVE®

Refer to the device insert (also printed at the back of this guide) for important information on the intended use/indications, device description, contraindications, precautions, warnings, and potential risks associated with this system.

STEP 1 PREPARATION AND GUIDE WIRE PLACEMENT

After locating the iliac crest, palpate either the posterior or anterior crest. For the anterior crest, position the **Guide Wire** 2cm posterior to avoid the lateral femoral cutaneous nerve. The wire is provided with 5mm depth marks near the insertion tip to prevent over-penetration into the crest. These marks should be used to insert the wire 15mm into the crest. Using the flat side of the Hammer, gently tap the wire into the iliac crest. The shoulder at the bottom of the wire can provide tactile feedback when the guide wire is fully docked on the crest.

Note: The shoulder of the guide wire must be resting on the iliac crest.



Guide wire preparation



Guide wire placement
in iliac crest

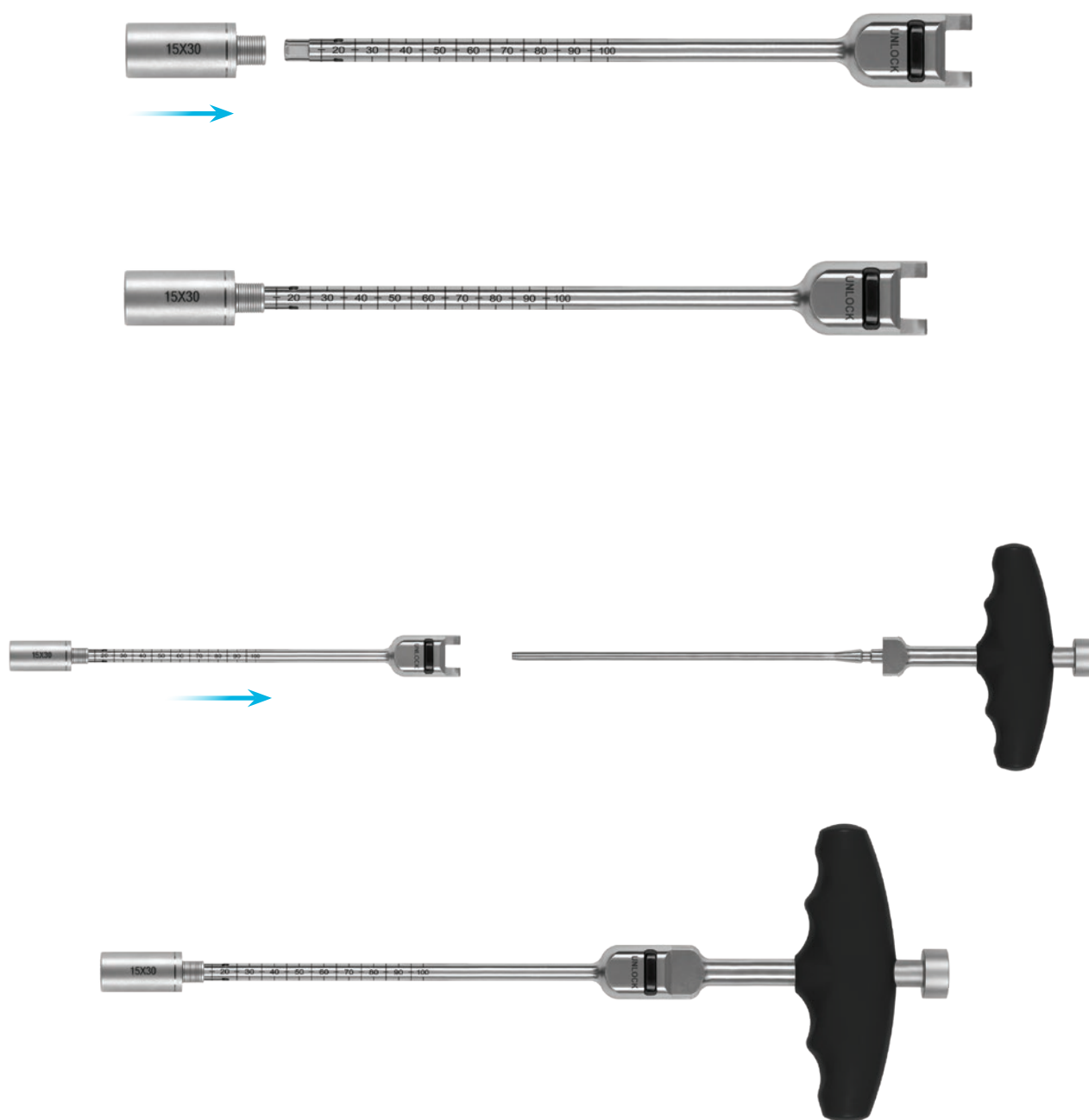
STEP

2

DRIVER ASSEMBLY AND INSERTION

Note: For instruction on use of the first generation Driver (691.020), please refer to the Appendix.

Determine the desired bone plug diameter. Place the selected **RETRIEVE® Tip** (7, 10, 12, or 15mm) onto the end of the **Driver Outer Shaft**. Slide the outer shaft over the **Driver Inner Shaft** until it locks into place and an audible click is heard.



DRIVER ASSEMBLY AND INSERTION (CONT'D)

Insert the driver into the iliac crest by placing the driver over the guide wire. Use the **Hammer** to advance the driver into the crest. Use the depth marks at every 5mm as a guide to penetrate 30mm into the iliac crest. It is important not to sink the driver below the surface of the iliac crest.



STEP

3

DRIVER ROTATION AND REMOVAL

Rotate the driver a quarter to half turn to remove the core from the surrounding tissue. Using a twisting motion, ease the driver from the iliac crest. If twisting does not suffice, the **Slide Hammer, Small** can be used to help release the driver.



DRIVER ROTATION AND REMOVAL (CONT'D)

Remove the bone core by inserting the **Graft Remover** into the back of the driver. Using the Hammer, gently tap on the end of the Graft Remover to expose the Guide Wire with the attached bone core. Remove the bone core from the wire.

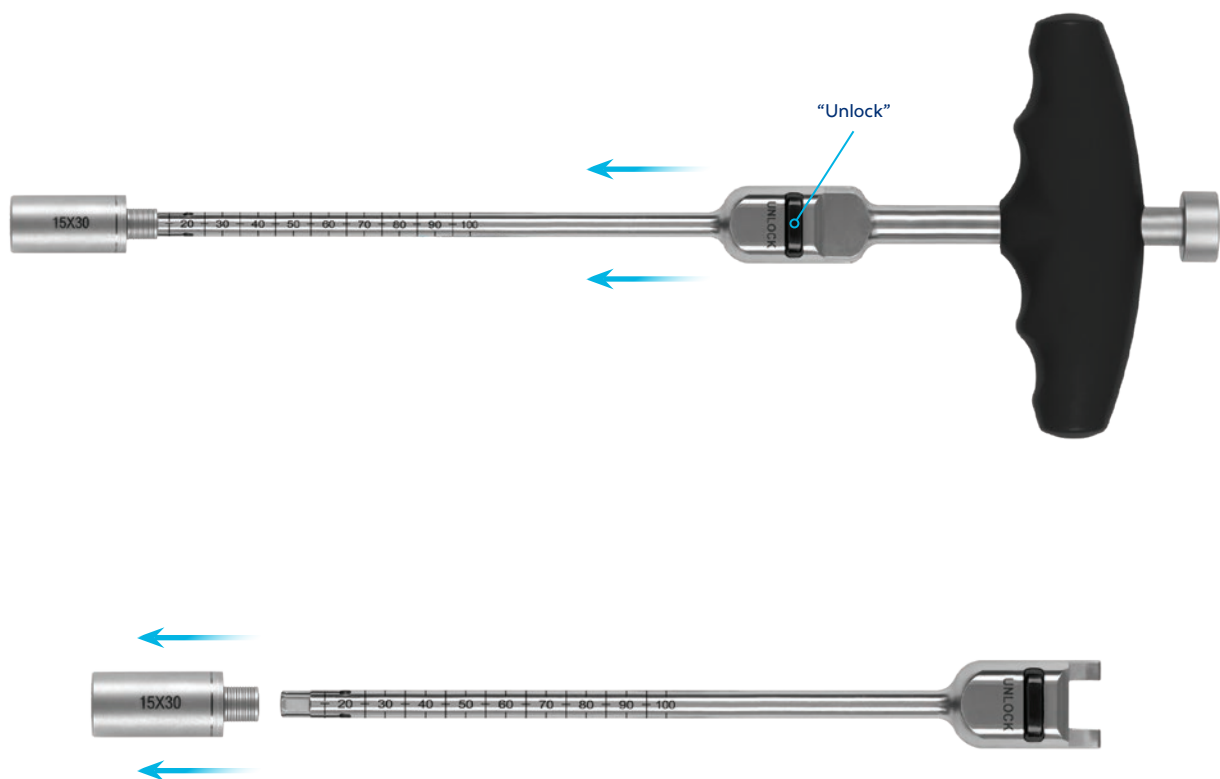
Note: Perform this step over a sterile table to prevent exposure to the non-sterile field.



Technique Tip: The incision may be injected with anesthetics and/or steroids to provide postoperative analgesia, hemostasis, and potentially reduce swelling.

STEP**4****DISASSEMBLY**

After the bone core is removed from the guide wire, dispose of the guide wire. To disassemble the driver, press the “Unlock” button to release the outer sleeve from the inner sleeve. Slide the outer sleeve distally from the handle to separate. Once the driver is disassembled, remove the tip from the outer sleeve and dispose.

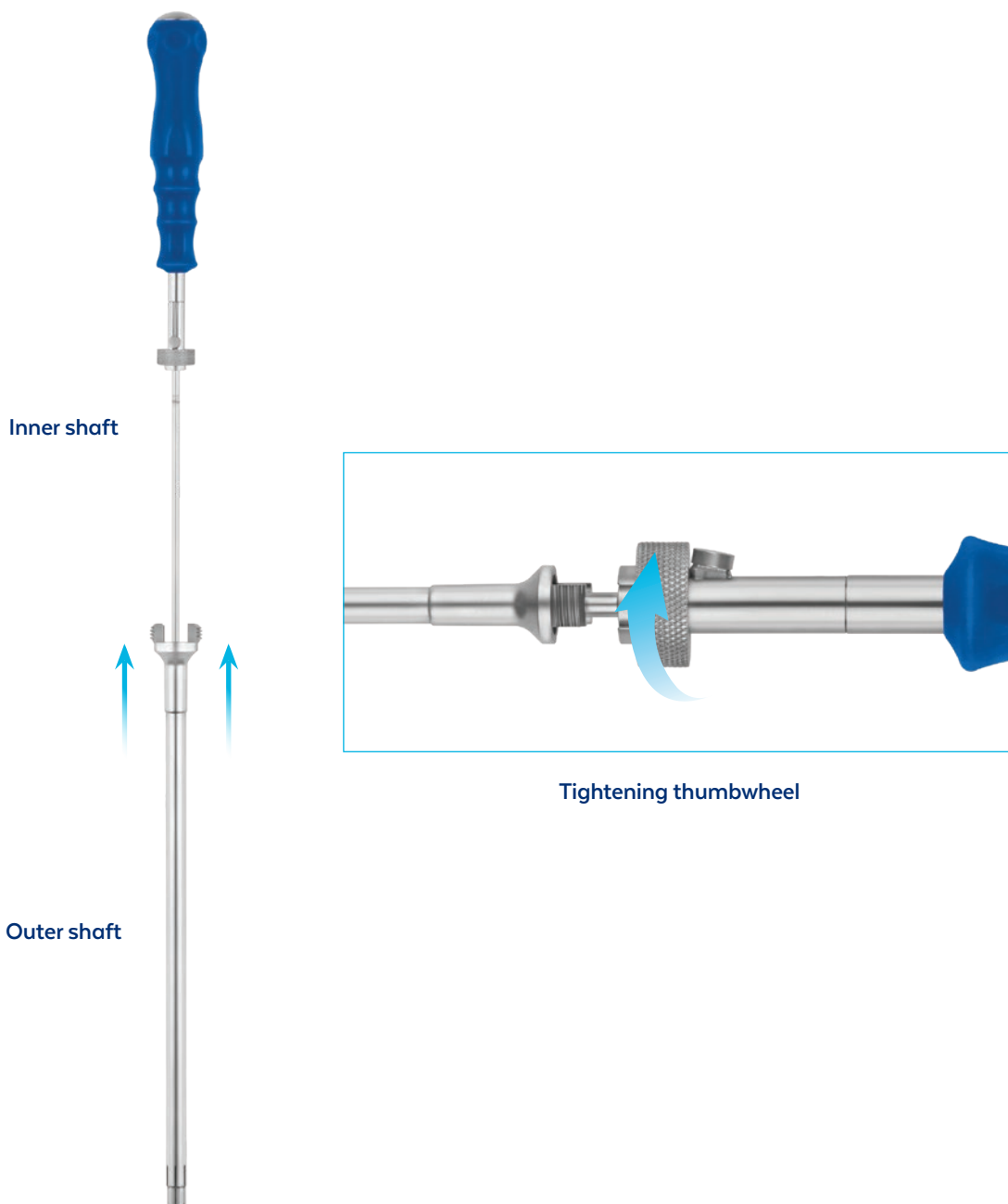


OPTIONAL TECHNIQUE: USING THE DRIVER (691.020)

Follow Step 1 for preparation and guide wire placement.

Determine the desired bone plug diameter. Select a disposable **RETRIEVE® Tip** based on the size of the graft needed. Tips are available in diameters of 7, 10, 12, and 15mm. Insert the desired tip onto the end of the outer shaft of the **Driver**. Assemble the driver by sliding the outer shaft over the inner shaft until it stops. Screw the outer shaft to the inner shaft.

Align the outer shaft tabs with the inner shaft slots and press the outer shaft into the inner shaft along its axis. Thread the thumbwheel until tightened; it must be tightened completely before use.



Insert the Driver into the iliac crest by placing the Driver over the Guide Wire. Hammer the driver into the crest. Use the depth marks at every 5mm as a guide to penetrate 30mm into the iliac crest. It is important not to sink the driver below the surface of the iliac crest.



Placing Driver over Guide Wire



Hammering Driver into crest

OPTIONAL TECHNIQUE: USING THE DRIVER (691.020) (CONT'D)

Rotate the Driver a quarter to half turn to embed the bone plug in the barrel of the tip. Press and hold the thumbwheel lock to capture the Guide Wire, and remove the Driver and wire simultaneously.



Rotating Driver

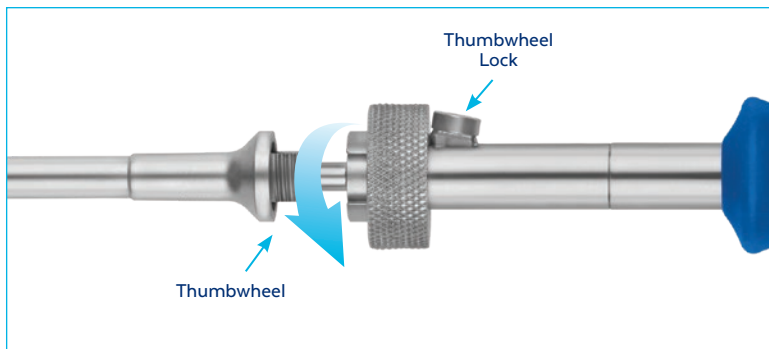


Removing Driver and
Guide Wire

Insert the **Graft Remover** into the handle of the driver until it meets with the guide wire. Lightly hammer on the graft remover until the wire and bone plug are pushed out of the driver.

Press and hold the thumbwheel lock and simultaneously unscrew the thumbwheel to disengage the shaft. Remove and dispose of the tip and wire.

Note: Do not disassemble the driver prior to bone plug removal.



Unscrewing thumbwheel



RETRIEVE®

BONE GRAFT HARVEST SYSTEM SETS

Instrument Set 991.901

	Part No.	Description	Qty
	691.020	Driver	0
1	691.024	Hammer	1
2	691.025	Graft Remover	1
3	691.120	Driver Outer Shaft	1
4	691.121	Driver Inner Shaft	1
5	622.410	Slide Hammer, Small	1

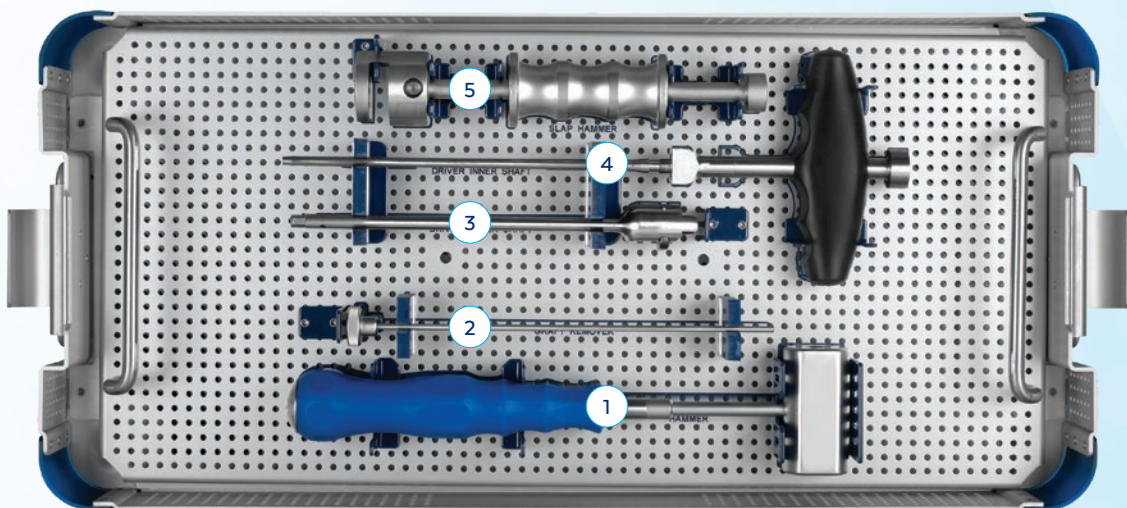
Disposable Instrument Set 991.902

Sterile, Single-Use

	Part No.	Description	Qty
	691.002S	RETRIEVE® Tip, 7x30mm	2
	691.003S	RETRIEVE® Tip, 10x30mm	2
	691.004S	RETRIEVE® Tip, 12x30mm	2
	691.005S	RETRIEVE® Tip, 15x30mm	2
	691.022S	Guide Wire	2

RETRIEVE®

BONE GRAFT HARVEST SYSTEM SETS



IMPORTANT INFORMATION ON THE RETRIEVE® UNIVERSAL BONE GRAFT HARVESTING SYSTEM

DESCRIPTION

The RETRIEVE® Universal Bone Graft Harvesting System is comprised of the RETRIEVE® Bone Graft Harvest Set and the RETRIEVE® Bone Marrow Aspiration Kit. The Bone Graft Harvest Set includes instruments for harvesting autogenous bone from the iliac crest. The instruments include guide wires, cutting tips, graft remover, and various other instruments. The Bone Marrow Aspiration Kit includes instruments for aspirating bone marrow or blood from the iliac crest. The kit includes needle(s), syringes, and mixing components.

INDICATIONS

The RETRIEVE® Bone Graft Harvest Set is used to harvesting autogenous bone from the iliac crest. The harvested bone graft can then be used for a variety of uses, including filling spinal interbody fusion devices and supporting posterolateral fusion.

The RETRIEVE® Bone Marrow Aspiration Kit is intended for aspiration of bone marrow, autologous blood, plasma or other blood components. The harvested bone marrow aspirate or blood can then be used for a variety of uses, including filling spinal interbody fusion devices and supporting posterolateral fusion. The bone marrow aspirate or autologous blood may be combined with bone graft or bone void fillers.

WARNINGS AND PRECAUTIONS

- This device is designed to be used by a physician.
- These instructions are not meant to define or suggest any medical or surgical technique. The individual practitioner is responsible for the proper procedure and technique to be used with this device.
- Possible allergic reactions should be considered.
- Check if the sterile inner package is unopened and damaged. In case of damaged inner package, do not use the product.
- Check the expiration date on sterile packaging.
- Store in a cool and dry place, protect from light.
- Use of the device is restricted only to physicians.
- Sterility and integrity are guaranteed only if observed with the prescribed conditions.
- Each sterile unit is intended for single patient use only.
- Do not use the product if the package has been damaged.
- All components should be properly and safely discarded after use.
- DO NOT RE-STERILIZE ANY STERILE PACKED COMPONENTS, INCLUDING THE KIT OR DISPOSABLES WITHIN THE KIT.
- Universal precautions should be observed by all hospital personnel that work with contaminated or potentially contaminated instrumentation. Caution should be exercised when handling devices with sharp points or cutting edges.
- Personal Protective Equipment (PPE) should be worn when handling or working with contaminated or potentially contaminated materials, devices and equipment. PPE includes gown, mask, goggles or face shield, gloves and shoe covers.
- Do not place heavy instruments on top of delicate devices.
- During the manual cleaning procedures do not use metal brushes or scouring pads. These materials will damage the surface and finish of instruments. Soft bristled, nylon brushes and pipe cleaners should be used.
- Do not use high acidic (pH<4) or high alkaline (pH>10) products for disinfection or cleaning, since these can corrode metal, cause discoloration or stress fractures.
- Do not allow contaminated instruments to dry prior to reprocessing. All subsequent cleaning and sterilization steps are facilitated by not allowing blood, body fluid, bone and tissue debris, saline, or disinfectant to dry on used instrumentation.
- Follow the instructions and warnings issued by the suppliers of any cleaning and disinfection agents and equipment used.
- Saline and cleaning/disinfection agents containing aldehyde, mercury, active chlorine, bromine, bromide, iodine or iodide are corrosive and should not be used. Instruments should not be placed or soaked in Ringers Solution.
- Mineral oil or silicone lubricants should not be used because they: 1) coat microorganisms; 2) prevent direct contact of the surface with steam; and 3) are difficult to remove.

SURGICAL GUIDELINES

These instructions are intended as guidelines for the use of the RETRIEVE® Universal Bone Graft Harvesting System as part of established surgical techniques. They are not intended to replace or change standard surgical procedures. Proper surgical procedures and techniques are the responsibility of the medical professional.

RETRIEVE® BONE GRAFT HARVEST SET

1. After suitable anesthesia is achieved, place the patient in the ventral supine position.
2. Using sterile technique, prepare the skin with antiseptic and drape.
3. After locating the iliac crest, palpate the anterior crest. Prepare the surgical site and create an incision.
4. Insert the guide wire into the posterior or anterior iliac crest.
5. Insert the driver over the guide wire and into the iliac crest .
6. Impact the driver into the crest and rotate to obtain the bone graft.
8. Remove the guide wire and remove the bone plug from the driver.
9. The bone graft can be applied directly to the surgical site.

RETRIEVE® BONE MARROW ASPIRATION KIT

1. After suitable anesthesia is achieved, place the patient in the ventral supine position.
2. Using sterile technique, prepare the skin with antiseptic and drape.
3. After locating the iliac crest, palpate the anterior crest. Prepare the surgical site and create an incision.
4. Remove the plastic guard from the aspiration needle.
5. Advance the needle with steady pressure and a slight twisting motion to the center of the anterior or posterior iliac prominence.
6. Advance needle using forward pressure through the cortex until all aspiration holes are engaged in the bone.
7. Remove the trocar needle and attach the syringe to the luer fitting on the aspiration needle.
8. Apply suction by withdrawing the syringe plunger. Remove the syringe with the harvested marrow.
9. Repeat until an appropriate amount of marrow is obtained.
10. The bone marrow aspirate can be applied directly to the surgical site or to the graft material.
11. The same procedure may be used to aspirate blood or other blood products.

PACKAGING

These instruments may be supplied pre-packaged and sterile, using gamma irradiation. The integrity of the sterile packaging should be checked to ensure that sterility of the contents is not compromised. Packaging should be carefully checked for completeness and all components should be carefully checked to ensure that there is no damage prior to use. Damaged packages or products should not be used, and should be returned to Globus Medical. During surgery, after the correct size has been determined, remove the products from the packaging using aseptic technique.

The instrument sets are provided nonsterile and are steam sterilized prior to use, as described in the STERILIZATION section below. Following use or exposure to soil, instruments must be cleaned, as described in the CLEANING section below.

HANDLING

All instruments should be treated with care. Improper use or handling may lead to damage and/or possible malfunction. Products should be checked to ensure that they are in working order prior to surgery. All products should be inspected prior to use to ensure that there is no unacceptable deterioration such as corrosion, discoloration, pitting, cracked seals, etc. Non-working or damaged instruments should not be used, and should be returned to Globus Medical.

CLEANING

All instruments that can be disassembled must be disassembled for cleaning. All handles must be detached. Instruments may be reassembled following sterilization. The instruments should be cleaned using neutral cleaners before sterilization and introduction into a sterile surgical field or (if applicable) return of the product to Globus Medical.

IMPORTANT INFORMATION ON THE RETRIEVE® UNIVERSAL BONE GRAFT HARVESTING SYSTEM

Cleaning and disinfecting of instruments can be performed with aldehyde-free solvents at higher temperatures. Cleaning and decontamination must include the use of neutral cleaners followed by a deionized water rinse. Note: certain cleaning solutions such as those containing formalin, glutaraldehyde, bleach and/or other alkaline cleaners may damage some devices, particularly instruments; these solutions should not be used.

The following cleaning methods should be observed when cleaning instruments after use or exposure to soil, and prior to sterilization:

1. Immediately following use, ensure that the instruments are wiped down to remove all visible soil and kept from drying by submerging or covering with a wet towel.
2. Disassemble all instruments that can be disassembled.
3. Rinse the instruments under running tap water to remove all visible soil. Flush the lumens a minimum of 3 times, until the lumens flush clean.
4. Prepare Enzol® (or a similar enzymatic detergent) per manufacturer's recommendations.
5. Immerse the instruments in the detergent and allow them to soak for a minimum of 2 minutes.
6. Use a soft bristled brush to thoroughly clean the instruments. Use a pipe cleaner for any lumens. Pay close attention to hard to reach areas.
7. Using a sterile syringe, draw up the enzymatic detergent solution. Flush any lumens and hard to reach areas until no soil is seen exiting the area.
8. Remove the instruments from the detergent and rinse them in running warm tap water.
9. Prepare Enzol® (or a similar enzymatic detergent) per manufacturer's recommendations in an ultrasonic cleaner.
10. Completely immerse the instruments in the ultrasonic cleaner and ensure detergent is in lumens by flushing the lumens. Sonicate for a minimum of 3 minutes.
11. Remove the instruments from the detergent and rinse them in running deionized water or reverse osmosis water for a minimum of 2 minutes.
12. Dry instruments using a clean soft cloth and filtered pressurized air.
13. Visually inspect each instrument for visible soil. If visible soil is present, then repeat cleaning process starting with Step 3.

CONTACT INFORMATION

Globus Medical may be contacted at 1-866-GLOBUS1 (456-2871). A surgical technique manual may be obtained by contacting Globus Medical.

STERILIZATION

These instruments may be available sterile or nonsterile.

Sterile instruments are sterilized by gamma radiation, validated to ensure a Sterility Assurance Level (SAL) of 10^{-6} . Sterile products are packaged in a heat sealed, double foil pouch. The expiration date is provided in the package label. These products are considered sterile unless the packaging has been opened or damaged.

Nonsterile instruments have been validated to ensure an SAL of 10^{-6} . The use of an FDA-cleared wrap is recommended, per the Association for the Advancement of Medical Instrumentation (AAMI) ST79, *Comprehensive Guide to Steam Sterilization and Sterility Assurance in Health Care Facilities*. It is the end user's responsibility to use only sterilizers and accessories (such as sterilization wraps, sterilization pouches, chemical indicators, biological indicators, and sterilization cassettes) that have been cleared by the FDA for the selected sterilization cycle specifications (time and temperature).

When using a rigid sterilization container, the following must be taken into consideration for proper sterilization of Globus devices and loaded graphic cases:

- Recommended sterilization parameters are listed in the table below.
- Only FDA-cleared rigid sterilization containers for use with pre-vacuum steam sterilization may be used.
- When selecting a rigid sterilization container, it must have a minimum filter area of 176 in² total, or a minimum of four (4) 7.5in diameter filters.
- No more than one (1) loaded graphic case or its contents can be placed directly into a rigid sterilization container.
- Stand-alone modules/racks or single devices must be placed, without stacking, in a container basket to ensure optimal ventilation.

- The rigid sterilization container manufacturer's instructions for use are to be followed; if questions arise, contact the manufacturer of the specific container for guidance.
- Refer to AAMI ST79 for additional information concerning the use of rigid sterilization containers.

For instruments provided NONSTERILE, sterilization is recommended (wrapped or containerized) as follows:

Method	Cycle Type	Temperature	Exposure Time	Drying Time
Steam	Pre-vacuum	132°C (270°F)	4 Minutes	30 Minutes

These parameters are validated to sterilize only this device. If other products are added to the sterilizer, the recommended parameters are not valid and new cycle parameters must be established by the user. The sterilizer must be properly installed, maintained, and calibrated. Ongoing testing must be performed to confirm inactivation of all forms of viable microorganisms.

CAUTION: Federal (USA) Law Restricts this Device to Sale by or on the order of a Physician.

SYMBOL TRANSLATION			
	CATALOGUE NUMBER		STERILIZED BY IRRADIATION
	LOT NUMBER		AUTHORISED REPRESENTATIVE IN THE EUROPEAN COMMUNITY
	CAUTION		MANUFACTURER
	SINGLE USE ONLY		USE BY (YYYY-MM-DD)
	QUANTITY		

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NOTES

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