

GIBRALT® Surgical Technique Cervical Thoracic Spinal System

The GIBRALT Spinal System is a comprehensive solution for posterior stabilization and fusion of the cervical and thoracic spine. Offering exceptional versatility and ease-of-use, the GIBRALT Spinal System features top-loading polyaxial screws, hooks, offset connectors, rod-to-rod connectors, and occipital plates which can be constructed into a multitude of configurations based on individual patient anatomy.

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# **OPERATIVE TECHNIQUE OVERVIEW**







Screw Placement







Tap to Desired Depth

8 Insert Screw







Cut and Contour the Rod

Place Rod into Tulip Heads

Insert Set Screw





Tighten Set Screws

13 Compression/Distraction



Final Tightening



Additional Options – Cross Connectors and Rod-to-Rod Connectors

# **DETAILED OPERATIVE TECHNIQUE**

# PRE-OPERATIVE PLANNING

When using the Gibralt Spinal Screw System, the patient should be positioned prone, lying flat on the table. A radiolucent frame or chest rolls may be used, but the kneeto-chest position should be avoided.

Using fluoroscopic imaging, it should be verified that the true views of both anterior-posterior (A/P) and lateral images of the spine (views which adequately delineate pedicle morphology and geometry) are obtainable. It is recommended that pre-operative planning is used to help determine a proper entry point and trajectory.



Hooks are available for use in the cervical spine. Select the appropriate Hook size and configuration for the anatomy. There are five different types of Hooks available: Straight, Left and Right Offset and Left and Right Angle Hooks (See page 12 for Hook offering).

Clamp the desired Hook with the **Hook Holder (05-009-40-0000)** making sure that the prongs of the instrument interface with the indentations on the outside head of the Hook (Figure 1).

Place Hooks as needed under the superior or inferior lamina (Figure 2).

The Hooks may be oriented either in a cranial or caudal position.

Repeat the steps above to place remaining hooks as determined in the pre-operative plan.

Once the Hooks have been inserted, utilize the **Set Screw Starter (05-009-20-0000)** to insert the appropriate inner Set Screw and provisionally tighten in a clockwise motion (Figure 3).





Fig. 2



Fig. 3









Fig. 6



# PLACEMENT OF POLYAXIAL SCREWS

Determine the ideal entry point for the Polyaxial Screw and penetrate the cortical bone to initiate an entry point using the **Awl (05-009-02-0000)** (Figure 4).

The **Probe (05-009-03-0000)** can then be used to cannulate the pedicle (Figure 5).

After the pedicle has been cannulated, it may be tested to ensure the integrity of the pedicular wall by using the **Sounding Probe (05-009-04-0000)** (Figure 6).

Determine the desired diameter and depth of the drill penetration. There are two drill options available, Fixed and Adjustable. Fixed drills are available in either a 10mm, 12mm or 14mm depth. Adjustable Drill (05-009-94-00XX) and Drill Guide Stop (05-009-93-0000) offer a drilling depth range from 14mm to 28mm in 2mm increments. The depth is determined by the position of the drill guide stop on the adjustable drill.

Attach the Drill Bit to the desired handle. Align the **Drill Guide (05-009-15-0000)** with the appropriate screw trajectory. Insert the Drill Bit through the **Drill Guide (05-009-15-0000)** and proceed with drilling to the desired depth (Figure 7).

Confirm depth and containment within the pilot hole with the **Depth Gauge (05-009-10-0000)** or probe. Tap the pilot hole using the **3.5mm Tap (05-009-05-0000)** or **4.0mm Tap (05-009-07-0000)** while maintaining the appropriate trajectory (Figure 8).

Note: The Taps are undersized by approximately 0.15mm. Continue to drill and tap the remaining pilot holes in the same manner.

Fig. 7

## **SCREW INSERTION**

After selecting the appropriate screw size, insert the hexalobe tip of the **Polyaxial Screwdriver (05-009-85-0000)** into the screw. Rotate the outer knob of the Screwdriver clockwise until the head of the screw is secured on the Driver (Figure 9). Insert the screw into the prepared pilot hole to the desired depth (Figure 10).

To disengage the screw from the driver, turn the knob counterclockwise and pull straight out of the internal hexalobe on the screw. To back out or adjust the screw, insert the hexalobe tip of the screwdriver into the screw. Rotate the outer knob of the screwdriver clockwise until the screwdriver is secured to the screw. Once the screw is engaged, back out the screw by turning the handle counterclockwise. Continue to insert all remaining Polyaxial Screws in the same manner.



# Fig. 9 Fig. 10

# **ROD PLACEMENT**

The rods are provided in pre-contoured, precut lengths, however a **Rod Cutter (05-009-31-0000)** is provided if other sizes are needed. The final length of the rod should extend 2mm beyond the margin of the screw housing so the screw locking mechanism engages correctly (Figure 11). To contour the rods, secure the rod with the **Rod Bender (05-009-22-0000)** and contour to achieve the desired curvature (Figure 12).

Handheld Rod Benders (05-009-23-0001 & 05-009-23-0002) are also available and can be used to provide additional leverage when contouring the rod. Utilize the removal screwdriver to adjust the A-P height of the screws as needed. Adjust the alignment of the Polyaxial Screws using the Head Adjuster (05-009-59-0000) so that the rod openings are in alignment. Once adjusted, they will easily stay in the correct alignment due to the unique EZ Set Tulip Design.



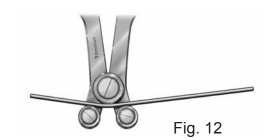




Fig. 13



Fig. 14



Fig. 15

Place the contoured rod into Polyaxial Screw (Figure 13). If necessary, there are two options for reducing the rod into the Hook or screw heads.

- 1) The **Rod Persuader (05-009-33-0000)** can be placed over the head of the hook or screw and the handles compressed to reduce the rod (Figure 14).
- 2) There are also multiple Inline Rod Persuaders (05-009-65-0000) which fit over the individual screw heads. The Inline Rod Persuader Knob (05-009-66-0000) is inserted over the top of the Inline Rod Persuader (05-009-65-0000) and rotated clockwise to reduce the rod (Figure 15). The Rod Persuader Knob can then be removed and placed over the next Inline Rod Reducer to reduce the rod into the next screw head sequentially.

## SET SCREW INSERTION

Determine the appropriate inner set screw for each Polyaxial Screw or Hook, depending on whether a cross connector will be used at the indicated level. Utilizing the **Set Screw Starter (05-009-20-0000)**, insert the appropriate inner set screw into the Hooks and Polyaxial Screws and provisionally tighten in a clockwise motion (Figure 16).



Fig. 16

## COMPRESSION/DISTRACTION

After the construct has been properly assembled, segmental compression and/or distraction can be accomplished using the **Compressor (05-009-28-0000)** or **Distractor (05-009-29-0000)** while tightening the Set Screws sequentially (Figure 17).

## FINAL TIGHTENING

To perform final tightening of the construct, attach the **Set Screw Driver (05-009-19-0000)** to the **Torque Limiting Handle (05-009-78-0000)** (22 in/lb) slide the **Inline Counter Torque (05-009-88-0000)** over the screw head and seat it against the rod. Insert the **Set Screw Driver (05-009-19-0000)** and turn the **Torque Limiting Handle (05-009-78-0000)** clockwise until an audible/tactile click is achieved (Figure 18). Repeat in the same manner on all remaining components to secure the construct.

## **ADDITIONAL OPTIONS**

# **ROD TO ROD CROSS CONNECTOR PLACEMENT**

Choose the appropriate size Rod to Rod Cross Connector and contour as needed, using the Cross Connector Benders (05-009-41-0000) provided. Capture the Cross Connector Nut with the Cross Connector Nut Starter (05-009-53-0000) to hold the cross connector and place onto the rod (Figure 19). Insert the Set Screwdriver (05-009-19-0000) Attach to the Torque Limiting Driver (05-009-88-0000) through the Torque Limiting Nut Driver (05-009-53-0000) over the rod to rod cross connector nut into the set screw. Rotate the Torque Limiting Nut Driver (05-009-53-0000) counterclockwise until the Torque handle breaks over (Figure 20). Repeat the procedure on the opposite side to final tighten the construct.

# SCREW HEAD TO SCREW HEAD CROSS CONNECTOR PLACEMENT

Choose the appropriate size Screw Head to Screw Head Cross Connector and contour as needed, using the Cross Connector Benders (05-009-41-0000) provided (Figure 21). Note: It is critical that the Cross Connector be contoured in most situations so that both rings of the cross connector are flush with the base of the tulip. If not contoured correctly, the application of torque to the Cross Connector Nut may cause issues with the inner set screw. Place the Hook Holder (05-009-40-0000) around the outside of the Polyaxial Screw where the Cross Connector will be inserted. Capture the retaining nut with the Cross Connector Nut Starter (05-009-53-0000) and turn clockwise to thread onto the extended Set Screw (Figure 22). Attach the Cannulated Palm Torque Nut Driver (05-009-88-0000) on the cross connect nut, then put the Set Screw Driver with torque T-handle (05-009-78-0000) through cannulation palm torque handle to engage the T-15 set screw drive feature. Hold the Torque T-handle stationary, then rotate the cannulated palm torque handle clockwise until it audibly clicks to secure the Cross Connecter Retaining Nut (Figure 23).

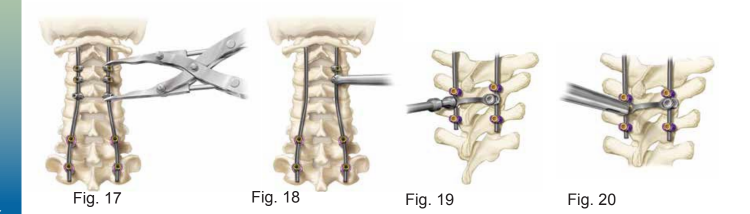
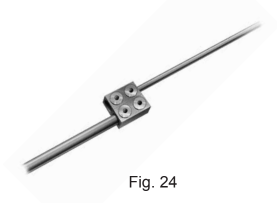




Fig. 22



Fig. 23



## TRANSITIONAL RODS

Transitional Rods and Rod Connectors are available to link to other ChoiceSpine Spinal Systems. The Gibralt System offers two different types of transitional rods, which can be linked to thoracic components.

# **ROD-TO-ROD TRANSITION CONNECTORS**

The Gibralt offers three different sizes of Rod-to-Rod Connectors for use with other ChoiceSpine Spinal Systems. See page 11 with images of Connector options.

- 1) Axial Rod-to-Rod Connectors are available to connect 3.5mm to 5.5.mm and 3.5mm to 6mm rod sizes.
- 2) Wedding Band Connectors are available to connect 3.5mm to 5.5.mm and 3.5mm to 6mm rod sizes.
- 3) Inline Rod-to-Rod Connectors are available for 3.5mm to 3.5mm rods.

To utilize a Rod-to-Rod connector, select the appropriate type and size and insert the end of the 3.5mm rod into the 3.5mm opening of the connector. Use the **Set Screw Starter (05-009-20-0000)** to engage the appropriate set screw by turning clockwise in the locking hole to secure the rod provisionally.

Note: The open side of the wedding band connector utilizes the Rod to Rod Connector Set Screw.

All other Rod to Rod Connectors utilize the Standard Set Screw. Next, insert the other rod (either 3.5mm, 5.5.mm or 6.0mm depending on the component used) into the remaining opening on the connector. Use the **Set Screw Starter (05-009-20-0000)** to engage the appropriate Set Screw by turning clockwise in the locking hole to secure the rod provisionally (Figure 24). Use the **Set Screw Driver (05-009-19-0000)** connected to the **Torque Handle (05-009-78-0000)** to tighten all Set Screws until the Torque Handle audibly clicks.

# **IMPLANT LISTING**

PART NUMBER	DESCRIPTION	
05-000-20-3510 05-000-20-3512 05-000-20-3514 05-000-20-3516	Screw, Polyaxial, 3.5X10 Screw, Polyaxial, 3.5X12 Screw, Polyaxial, 3.5X14 Screw, Polyaxial, 3.5X16	
05-000-20-3518 05-000-20-3520 05-000-20-3522	Screw, Polyaxial, 3.5X18 Screw, Polyaxial, 3.5X20 Screw, Polyaxial, 3.5X22	The state of the s
05-000-20-3524 05-000-20-3526 05-000-20-3528	Screw, Polyaxial, 3.5X24 Screw, Polyaxial, 3.5X26 Screw, Polyaxial, 3.5X28	
05-000-20-4010 05-000-20-4012 05-000-20-4014 05-000-20-4016	Screw, Polyaxial, 4.0X10 Screw, Polyaxial, 4.0X12 Screw, Polyaxial, 4.0X14 Screw, Polyaxial, 4.0X16	
05-000-20-4018 05-000-20-4020 05-000-20-4022 05-000-20-4024	Screw, Polyaxial, 4.0X18 Screw, Polyaxial, 4.0X20 Screw, Polyaxial, 4.0X22 Screw, Polyaxial, 4.0X22 Screw, Polyaxial, 4.0X24	
05-000-20-4026 05-000-20-4028 05-006-01-3501	Screw, Polyaxial, 4.0X26 Screw, Polyaxial, 4.0X28  Hook Straight (Image Shown)	
05-006-01-3502 05-006-01-3503 05-006-01-3504 05-006-01-3505	Hook Left Angle Hook Right Angle Hook Offset Right Hook Offset Left	
05-000-24-0275	Set Screw	0
05-004-00-2230 05-004-00-2636 05-004-00-3242 05-004-00-3848 05-004-00-4452	X-Connector, Rod to Rod, 22-30 X-Connector, Rod to Rod, 26-36 X-Connector, Rod to Rod, 32-42 X-Connector, Rod to Rod, 38-48 X-Connector, Rod to Rod, 44-52	<b>5</b> 30 mg
05-004-10-2230 05-004-10-2636 05-004-10-3242 05-004-10-3848 05-004-10-4452	X-Connector, Head to Head, 22-30 X-Connector, Head to Head, 26-36 X-Connector, Head to Head, 32-42 X-Connector, Head to Head, 38-48 X-Connector, Head to Head, 44-52	2 · 2n · n
05-000-25-0001	Connector, Nut	
05-000-24-0925	Connector, Set Screw	M0000000
05-002-03-3530 05-002-03-3540 05-002-03-3550 05-002-03-3560 05-002-03-3570 05-002-03-3580	Rod, Prebent, Ti, 3.5X30 Rod, Prebent, Ti, 3.5X40 Rod, Prebent, Ti, 3.5X50 Rod, Prebent, Ti, 3.5X60 Rod, Prebent, Ti, 3.5X70 Rod, Prebent, Ti, 3.5X80	
05-002-03-3590 05-004-08-3512 05-004-08-3515	Rod, Prebent, Ti, 3.5X90  Offset, Connector 3.5mm x 12mm  Offset, Connector 3.5mm x 15mm	

PART NUMBER	DESCRIPTION	
05-004-07-3555 05-004-07-3560	Connector, WEDDING Band, 3.5-5.5 Connector, WEDDING Band, 3.5-6.0	
05-004-06-3555 05-004-06-3560	Connector, COMBINATION, 3.5-5.5 Connector, COMBINATION, 3.5-6.0	
05-004-05-3535	Connector, Inline, 3.5-3.5	
05-000-24-0002	Set Screw, Rod to Rod	
05-002-00-3512 05-002-00-3524	Rod, Straight, Ti, 3.5X120 Rod, Straight, Ti, 3.5X240	
05-002-01-0001	Rod, Transition, Ti, 3.5mm-5.5mm,420mm	

## **INSTRUMENT LISTING**

INSTRUMENT LIST	ING	
PART NUMBER	DESCRIPTION	
05-009-02-0000	Straight Awl 3.5	
05-009-03-0000	Probe 1.75	
05-009-04-0000	Straight, Sounder	
05-009-05-0000 05-009-07-0000	Tap, 3.5 Tap, 4.0	
05-009-10-0000	Depth Gauge	
05-009-10-2010 05-009-10-2012 05-009-10-2014	Drill, 2.0X10 Drill, 2.0X12 Drill, 2.0X14	
05-009-15-0000	Drill Guide	
05-009-19-0000	Set Screw, Driver, Retaining Shaft	
05-009-20-0000	Set Screw, Starter	
05-009-26-0000	Rod Holder, Forceps	
05-009-40-0000	Hook Holder	
05-009-53-0000	X-Connector Nut Starter	
05-009-56-1000	Polyaxial Screwdriver, T10	
05-009-59-0000	Poly, Head Adjuster	
		CM0=-000
05-009-78-0000	Torq, T Handle 1/4 SQ Drive, 22 IN	
05-009-85-0000	Screwdriver, Fixed Handle	
05-009-87-0000	Rod Rocker	2
05-009-88-0000	Inline Countertorque	
05-009-93-0000	Adjustable Stop	
05-009-94-0020	Drill, Adjustable, 2.0	to the state of th
05-009-95-0000	Torque Palm, Driver	
05-009-99-0000	Ratcheting AO STR Handle Straight	

D N	Programme	
<b>PART NUMBER</b> 05-009-22-0000	<b>DESCRIPTION</b> Rod Bender, 3.5	08
05-009-23-0001	IN SITU, Rod Bender, Left	•
05-009-23-0002	IN SITU, Rod Bender, Right	12000
05-009-28-0000	Compressor	
05-009-29-0000	Distractor	
05-009-31-0000	Rod Cutter, 3.5	
05-009-33-0000	Rod Persuader	
05-009-35-0000	Rod Template	
05-009-41-0000	Spine PC Bender, X-CONN	
05-009-65-0000	Inline Rod Persuader	
05-009-66-0000	Rod Persuader Knob	
05-009-91-0000	Under Rod Reducer	



## Gibralt® Spine System Instruction for Use







The ChoiceSpine Gibralt Spine System is a posterior system intended to help provide immobilization and stabilization of spinal segments as an adjunct to fusion of the cervical, and/or upper thoracic spine. The system consists of a variety of sizes of rods, hooks, poly-axial screws and connecting components, which can be rigidly locked to the rod in various configurations. The Gibral Spine System components are manufactured from titanium alloy, with rods being available in both titanium alloy and cobalt chrome alloy options.

This system can be used independently or in conjunction with ChoiceSpine 5.5mm or 6.0mm rod-based Thoraco-Lumbar Pedicle Screw Systems. The 5.5mm or 6.0mm rod-based Pedicle Screw systems are not covered by these instructions for use. Reference the instructions for use accompanying the Pedicle Screw System components for complete instructions for use

### Indications for Use:

The ChoiceSpine Gibralt Spine System is intended to provide immobilization and stabilization of spinal segments as an adjunct to fusion for the following acute and chronic instabilities of the cervical spine (C1-C7) and the thoracic spine from T.-13: ratu-matic spinal fractures and/or traumatic dislocations; instability or deformity; failed previous fusions (e.g., pseudarthrosis); tumors involving the cervical/thoracic spine; and degenerative disease including intractable radiculopathy and/or myelopathy, neck and/or arm pain of discogenic origin as confirmed by radiographic studies, and degenerative disease of the facets with instability. The Gibrat Spine System is also intended to restore the integrity of the spinal column even in the absence of fusion for a limited time period in patients with advanced stage tumors involving the cervical spine in whom life expectancy is of insufficient duration to permit achievement of fusion.

In order to achieve additional levels of fixation, the Gibralt Spine System may be connected to the Gibralt Occipital Spine System with rod-to-rod connectors. The Gibralt Spine System may also be connected to the ChoiceSpine Proliant System, ChoiceSpine Silverbolt and Mainframe Spinal Screw Systems, or ChoiceSpine Hydralok, using rod-to-rod connectors and transitional rods. Refer to the specific system package inserts for a list of their indications for use.

### Contraindications:

Contraindications include, but are not limited to:

- Presence of overt infectious process or significant risk of infection (immunocompromise)
- Fever or leukocytosis
- Morbid obesity
- Pregnancy
- Mental illness
- Grossly distorted anatomy caused by congenital abnormalities Any other medical or surgical condition which would preclude the potential benefit of spinal implant surgery, such as the presence of congenital abnormalities, elevation of sedimentation rate unexplained

by other diseases, elevation of white blood count, or a marked left shift in the white blood count differential count

- Suspected or documented metal allergy or intolerance Rapid joint disease, bone absorption, osteopenia, osteomalacia and/or osteoporosis. Osteoporosis or osteopenia is a relative contraindication since this condition may limit the degree of obtainable correction, stabilization, and/or the amount of mechanical fixation

  Any patient unwilling to follow postoperative instructions
- Any case not needing a bone graft and fusion
- Any case where the implant components selected for use would be too large or too small to achieve a successful result
  Any case that requires the mixing of metals from two different
- components or systems Any patient having inadequate tissue coverage over the operative site
- or inadequate bone stock or quality

  Any patient in which implant utilization would interfere with anatomical
- structures or expected physiological performance Presence of any neural or vascular deficit or other compromising
- pathology, which may be further injured by device interventio Any case not described in the indications

## Warnings and Precautions:

The Gibral Spine System should only be implanted by experienced spine surgeons with specific training in the use of this spine system because this is a technically demanding procedure presenting a risk of serious injury to the patient. In addition, the surgeon should consider the levels of implantation, patient weight, patient activity level, and other patient conditions (e.g., smoking, occupation), which may impact on the performance of the system

The Gibralt Spine System has not been evaluated for safety and compatibility in the MR environment. Gibral Spine System has not been tested for heating, migration, or image artifact in the MR environment. The safety of the Gibralt Spine System in the MR environment is unknown. Scanning a patient who has this device may result in patient injury

A successful result is not always achieved in every surgical case, especially in spinal surgery where many extenuating circumstances may compromise the results. Preoperative planning and operating procedures, including knowledge of surgical techniques, proper reduction, and proper selection and placement of the implant are critical considerations in achieving a

Use of posterior cervical pedicle screw fixation at the C3 through C6 spinal levels requires careful consideration and planning beyond that required for lateral mass screws placed at these spinal levels, given the proximity of the vertebral arteries and neurologic structures in relation to the cervica pedicles at these levels.

Pre-operative planning prior to implantation of posterior cervical lateral mass and pedicle screw spinal systems should include review of cross-sectional imaging studies (e.g., CT and/or MRI imaging) to evaluate the patient's certical anatomy including the transverse foramen and the course of the vertebral arteries. If any findings would compromise the placement of lateral mass or pedicle screws, other surgical methods should be

considered. In addition, use of intraoperative imaging should be considered to guide and/or verify device placement, as necessary.

Only patients that meet the criteria described in the indications should be ed. Patient conditions and/or predispositions such as mentioned in the contraindications should be avoided

The type of construct to be assembled for the case should be determined prior to beginning the surgery. An adequate inventory of sizes should be available at the time of surgery, including sizes larger and smaller than th

Since mechanical parts are involved, the surgeon should be familiar with the rrious components before using the equipment and should personally semble the devices to verify that all parts and necessary instruments are present before the surgery begins.

All components and instruments must be cleaned and sterilized prior to use

Additional sterile components should be available in case of unexpected need.

This device system is not intended to be the sole means of spinal support. Use of this product without a bone graft or in cases that develop into a nonunion will not be successful.

No spinal implant can withstand body loads without the support of bone. In this event, bending, loosening, disassembly and/or breakage of the device(s) will eventually occur.

Prior to surgery, the patient must be informed of all potential risks and adverse effects contained in the present instructions for use.

The selection of the proper size, shape and design of the implant for each patient is crucial to the success of the procedure. Metallic surgical implants are subject to repeated stresses in use, and their strength is limited by the need to adapt the design to the size and shape of human bones. Unless great care is taken in patient selection, proper placement of the implant, and postoperative management to minimize stresses on the implant, such stresses may cause metal fatigue and consequent breakage, bending or loosening of the device before the healing process is complete, which may result in further injury or the need to remove the device prematurely.

### Intraoperative

The surgeon should follow established practices and specific instructions for implant of the system. Whenever possible or necessary, an imaging system should be utilized to verify proper component placement.

Extreme caution should be used around the spinal cord and nerve Damage to the nerves will cause loss of neurological functions.

If screws are bent or damaged during insertion or adjustment, they may not be implanted and must be replaced. Rods should only be contoured with the proper contouring instruments. The rods should not be repeatedly or excessively bent. The rods should not be reverse bent in the same location. Use great care to ensure that the implant surfaces are not scratched or notched, since such actions may reduce the functional strength of the construct. If the rods are cut to length, they should be cut in such a way as to create a flat, non-sharp surface perpendicular to the midline of the rod. Cut the rods outside the operative field and whenever possible, use pre-cut rods if available.

The rod-to-rod connecting components must be sized correctly for the diameter of the rods used.

Do not over-tap or use a screw that is either too long or too large. Overtapping or using an incorrectly sized screw hemorrhage, or other possible adverse events.

Bone grafts must be placed in the area to be fused.

Before closing the soft tissues, all the set screws should be tightened firmly. Recheck the tightness of all set screws after finishing to ma loosened during the tightening of the other set screws. Failure to do so may cause loosening of the other components.

Some degree of corrosion occurs on all implanted metal and alloys. Mixing of dissimilar metals can accelerate the corrosion process. Stainless steel and titanium implants must NOT be used together in building a construct. Stainless steel and cobalt chrome implants must NOT be used together in building a construct.

Different manufacturers use different materials, varying tolerances and design configurations. Components of the Gibralt Spine System must not be used with components from any other system or manufacturer.

## Postoperative:

Postoperative. Doubling and care are important. It is recommended that regular, long-term postoperative follow-up be undertaken to detect early signs of component failure, and to consider the course of action to be taken if such events occur.

The physician's postoperative directions and warnings to the patient and the corresponding patient compliance are extremely important.

Detailed instructions on the use and limitations of the device should be given to the patient. The patient should be instructed to limit and restrict physical activities, especially lifting and twisting motions and any type of sport participation.

If partial weight-bearing is recommended or required prior to firm bony union, the patient must be warned that bending, loosening and/or breakage of the device(s) are complications which may occur as a result of excessive or early weight-bearing or muscular activity. The risk of bending, loosening, or breakage of a temporary internal fixation device during postoperative rehabilitation may be increased if the patient is active, or if the patient is debilitated or demented. The patient should be warned to avoid falls or sudden jolts in spinal position. The patient should be advised of their inability to bend or rotate at the point of spinal fusion and taught to compensate for this permanent physical restriction in body motion.

patient should be advised not to smoke tobacco or utilize nicotine products, or to consume alcohol or non-steroidal or anti-inflammatory medications such as aspirin during the bone graft healing process.

non-union develops or the components loosen, bend, and/or break, the device(s) should be revised and/or removed immediately before serious injury occurs. Failure to immobilize a delayed or nonunion of bone will result in excessive and repeated stresses on the implant. By the mechanism of fatigue these stresses can cause eventual bending, loosening or breakage of the device(s). It is important that immobilization of the spinal surgical site be maintained until firm bony union is established and confirmed by examination.

After the spine is fused, these devices serve no functional purpose and should be removed. If the device is not removed following completion of its intended use, one or more of the following complications may occur: (1) Corrosion, with localized tissue reaction or pain, (2) migration of implant osition possibly resulting in injury, (3) risk of additional injury from

postoperative trauma, (4) bending, loosening and breakage, which could make removal impractical or difficult, (5) pain, discomfort, or abnormal sensations due to the presence of the device, (6) potential increased risk of infection, (7) bone loss due to stress shielding; and (8) potential unknown and/or unexpected long term effects due to wear particles such as and/or interpreted to light enhanced of the wear particles such as carcinogenesis. The surgeon should carefully weight her fisk versus benefits when deciding whether to remove the implant. Implant removal should be followed by adequate postoperative management to avoid fracture, refracture, or other complications.

Implants must not be reused. Any implant, once used, should be discarded even though it may appear undamaged.

### Potential Complications and Adverse Effects:

Potential complications and adverse effects for this system are similar to those of other spinal instrumentation systems, and include, but are not

- Early or late loosening of any or all the componen
- Disassembly, bending, and/or breakage of any or all the components Foreign body (allergic) reaction to implants, debris, corrosion products
- from crevice, fretting, and/or general corrosion products (from crevice, fretting, and/or general corrosion products) Pressure on the skin from component parts in patients with inadequate tissue coverage over the implant possibly causing skin penetration, irritation, fibrosis, necrosis, and/or pain; Bursitis
- Tissue or nerve damage caused by improper positioning and placement of implants or instruments.
- Post-operative change in spinal curvature, loss of correction, height, and/or reduction
  Infection
- 8. Dural tears, pseudomeningocele, fistula, persistent CSF leakage, meningitis
- Nerve damage due to surgical trauma or presence of device and temporary or permanent loss of neurologic function, including paralysis Urinary retention or loss of bladder control or other types of urological
- system compromise
- 11. Scar formation possibly causing neurological compromise or
- reacture, micro fracture, resorption, damage, or penetration of any spinal bone (including the vertebral body) and/or bone graft or bone graft harvest site at, above, and/or below the level of surgery
- Herniated nucleus pulposus, disc disruption or degeneration at, above, or below the level of surgery

  Non-union (or pseudarthrosis), delayed union or mal-union
  Loss of or increase in spinal mobility or function
- 16. Bone loss or decrease in bone density, possibly caused by stresses shielding
- 17. Graft donor site complications including pain, fracture, or wound
- healing problems.

  18. Ileus, gastritis, bowel obstruction or loss of bowel control or other types
- of gastrointestinal system compromise

  19. Hemorrhage, hematoma, occlusion, seroma, edema, hypertension,
  embolism, stroke, excessive bleeding, phlebitis, wound necrosis, wound
  dehiscence, damage to blood vessels, or other types of cardiovascular system compromise
- 20. Reproductive system compromise, including sterility, loss of consortium, and sexual dysfunction
- Development of respiratory problems, (pulmonary embolism, atelectasis, bronchitis, pneumonia)
   Death

Additional surgery may be necessary to correct some of these potential

## How Supplied:

The Gibralt Spine System devices are provided clean but non-sterile and must be sterilized prior to use. Implants are intended for single use only. Instruments can be reprocessed using the recommended cleaning instructions.

All instruments and implants are supplied to the hospital clean but nonsterile and require sterilization before each use. All instruments that have been previously taken into a sterile surgical field must first be decontaminated and cleaned before sterilization and reintroduction into a sterile surgical field. Implants that have been implanted and then removed must be discarded. Cleaning and disinfecting of instruments can be performed with alkali aldehyde-free solvents at high temperatures. Cleaning and decontamination can include the use of neutral cleaners follow by a deionized water rinse.

**Note:** Certain cleaning solutions such as those containing formalin, glutaraldehyde, bleach and/or alkaline cleaners may damage some devices, particularly instruments; these solutions should not be used.

All products should be treated with care. Improper use and handling may lead to damage and possible improper functioning of the device

These devices are packaged in a convenience caddy/case. All devices must be removed from the case, inspected and cleaned via one of the appropriate methods below. Where applicable, instruments should be disassembled prior to cleaning and reassembled prior to sterilization. All devices must be placed back into the caddy and case prior to steam sterilization

Recommended Cleaning
The terms "Steris 444", "Enzol®" and "Prolystica®" are tradenames of ultrasonic equipment and detergents utilized in the recommended cleaning instructions. Any ultrasonic washer or equivalent ultrasonic detergent car be utilized when used in accordance to the manufacturer's instructions and

- Rinse instrument(s) under cool running tap water (< 35 °C) to remove gross soil. Use a sterile syringe to flush water through and around cracks, crevices, and hard to reach areas.
- Use a soft bristle brush as needed to remove soil, paying close attention to threads, crevices, and hard to areas.
- Transfer instrument(s) into a STERIS 444 washer with the following parameters. Incline the instrument(s) to assist in drainage. Moto



Phase	Time (min)	Temperature	Detergent
Pre-Wash 1	1:00	Cold Tap Water	N/A
Enzyme Wash	1:00	Hot Tap Water	Enzol® at 1 oz per 1 gal water
Wash 1	2:00	60°C	Prolystica® 2x Conc. Neutral at 1/8 oz per 1 gal water
Rinse 1	1:00	Hot Tap Water	N/A
Drying	7:00	115°C	N/A

Remove instrument(s) from washer & visually inspect for soil. Repeat if necessary

- Mechanical Cleaning (Ultrasonic):

  1. Rinse instrument(s) under cool running tap water (< 35 °C) to remove gross soil. Use a sterile syringe to flush water through & around cracks, crevices. & hard to reach areas
- Prepare Enzol® solution of one (1) ounce per one (1) gallon of warm tap water (< 55 °C).

  Surgical Technique Manual: The Gibralt Spine System
- Fully immerse instrument(s) in the detergent for at least one (1)
- Use a soft bristle brush as needed to remove soil, paying close attention
- to threads, crevices, & hard to reach areas.
- to threads, crevices, & hard to reach areas.
  Use a sterile syringe to flush detergent through & around cracks, crevices, & hard to reach areas.
  Remove instrument(s) from detergent & rinse with cool tap water (< 35°C) for at least one (1) minute.
  Prepare the ultrasonic cleaner with an Enzol® solution of one (1) ounce

- 7. Prepare the uttrasonic cleaner with an enzor' solution of one (1) ounce per one (1) gallon of warm tap water (< 55°C).

  8. Load instrument(s) into the cleaner & sonicate for ten (10) minutes.

  9. Remove instrument(s) from cleaner & thoroughly rinse using reverse osmosis/denionized (R/O/D) water for at least one (1) minute.

  10. Dry instrument(s) using a clean, soft towel & filtered, pressurized air
- 11 Visually inspect for soil Repeat if necessary

- Rinse instrument(s) under cool running tap water (< 35 °C) to remove gross soil. Use a sterile syringe to flush water through & around cracks, revices. & hard to reach areas.
- Prepare Enzol<sup>®</sup> solution of one (1) ounce per one (1) gallon of warm tap water (< 55 °C).

  Fully immerse instrument(s) in the detergent for at least one (1) minute.
- Use a soft bristle brush as needed to remove soil, paying close attention to threads, crevices, & hard to reach areas.
- attention to Interests, crevices, & nature to reach areas.
  Use a sterile syringe to flight detergent through & around cracks, crevices, & hard to reach areas.
  Remove instrument(s) from detergent & thoroughly rinse with reverse osmosis/deionized (RO/DI) water for at least one (1) minute. Use a
- sterile syringe to aid in rinsing.

  Dry instrument(s) using a clean, soft cloth & filtered, pressurized air (20
- psi). Visually inspect for soil. Repeat if necessary.

## Care and Handling:

- All products should be treated with care. Improper use and handling may lead to damage and possible improper functioning of the device.
- Refer to ASTM standard F1744-96, "Standard Guide for Care and Handling of Stainless Steel Surgical Instruments" for additional information.
- Before use, instruments should be visually inspected, and function should be tested to ensure instruments are functioning properly. If instruments are discolored, have loose screws/pins, are out of alignment, cracked, show excessive wear, or have other irregularities,
- Lubricate instruments to protect instruments during sterilization and storage. This should be done with a water soluble, preserved lubricant after each cleaning. The lubricant should contain a chemical preservative to prevent bacterial growth and be made with distilled water. Excess lubricant should be wiped off prior to storage and sterilization

The implants should be inspected after processing, prior to sterilization. Any implant with damage, corrosion, discoloration, scratches, residue, or debris should be discarded.

## Sterilization

ChoiceSpine instruments are provided non-sterile and must be sterilized prior to use. All packaging materials must be removed prior to sterilization. Instruments are recommended to be steam sterilized by the hospital using the following process parameters (Alternative methods or cycles may be used, but should be validated according to hospital practices and procedures):

Steam Sterilizer Type: Pre-Vacuum Temperature: 132°C Duration: 4 minutes Drying Time: 40 minutes

All devices are to be wrapped in two-layer of 1-ply polypropylene wrap (Kimguard KC600 or equivalent) using various wrapping techniques per ANSI/AAMI ST79.

This steam sterilization cycle is not considered by the FDA to be a standard sterilization cycle. It is the end user's responsibility to use only sterilizers and accessories (such as sterilization wraps or pouches, chemical or biological indicators, and sterilization cassettes) that have been cleared by the FDA for the sterilization cycle specifications (time and temperature) Alternative sterilization methods or cycles may be used, but should be validated according to hospital practices and procedures. The use of an FDA cleared wrap is recommended to ensure devices remain sterile prior to implantation

Storage and Handling: Implants should be stored in their original, sealed packaging in clean, dry conditions. The packaging should not be exposed to direct sunlight, ionizing radiation, extreme temperatures, or particulate contamination. Prior to use, inspect the packaging and labeling for integrity. If the device has been opened, damaged or adulterated in any way, it must not be used. In order to ensure sterility, please observe aseptic surgical procedures when removing the implant from its packaging.

## Limitations and Restrictions:

Repeated sterilization according to these instructions has a minimal effect on ChoiceSpine devices. Sterilization equipment varies in performance characteristics and must be validated accordingly. The sterilizing facility is responsible for the routine validation and monitoring of all equipment,

materials and personnel used in their facility to ensure the desired results

These instructions have been validated as being capable of sterilizing these ChoiceSpine implants. Any deviations from these procedures must be evaluated for efficacy by the sterilizing facility.

It is essential to provide preoperative instructions to the patient. The It is essential to provide preoperative instructions to the patient. The patient should be made aware of the potential risks of the surgery and the implant limitations. The patient should be instructed to limit postoperative activity, as this will reduce the risk of bent, broken or loose implant components. The patient must be made aware that implant components may bend, break, or loosen even though restrictions in activity are followed.

Device Retrieval Efforts:
Should it become necessary to remove any or all the Gibralt® Spine System
"Challeding at the number helow to receive components, please call ChoiceSpine at the number below to receive instructions regarding data collection, including histopathological, mechanical, and adverse event information.

The Gibralt Spine System Surgical Technique Manual is available by contacting ChoiceSpine Customer Service

### Product Complaints:

Any dissatisfaction with the product quality, labeling, or performance should be reported to ChoiceSpine immediately by the customer or health care provider. Furthermore. ChoiceSpine should be notified immediately of an implant malfunction by telephone, fax, or written correspondence.

When filing a complaint, the name, part number, and lot number of the part should be provided along with the name and address of the person filing the complaint.

Some components may not be currently available. Please contact your ChoiceSpine representative for additional information. The products discussed herein may be available under different trademarks in different countries. All copyrights, and pending and registered trademarks are property of ChoiceSpine. For more information on a specific product or trademark, please contact your local ChoiceSpine representative.

Federal Law (USA) restricts this device to sale by or on the order of a physician.

### Information:

See choicespine.com for more information.

See choicespine.com/patents/ for patent information.

For product complaints please contact: ChoiceSpine, LLC Quality/Regulatory Department 400 Erin Drive

Knoxville, TN 37919 Phone: 865-246-3333; Fax: 865-588-4045

## For additional product information please contact:

MedPass SAS

ChoiceSpine, LLC Customer Service Department 400 Erin Drive Knoxville, TN 37919 Phone: 865-246-3333; Fax: 865-588-4045 customerservice@choicespine.com



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Symbol	Definition
2	Do not reuse
$\triangle$	Caution, consult instructions for use for warnings and precautions
[]i	Consult instructions for use
<b>®</b>	Do not use if package is damaged
LOT	Lot number
REF	Reference number
SN	Serial Number
STERILE R	Sterilized by irradiation
$\square$	Use by
***	Manufacturer
~~	Date of Manufacture
Ronly	Federal law (USA) restricts this device to sale by or on the order of a physician
NON	Non-Sterile
C€	European Medical Devices
EC REP	Authorized representative in the European Community



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