-57	IFU-014-T02-03	INSTRUCTION FOR USE	: SOUTHERN AXIS™ 2 CERVICA	AL PLATE SYSTEM		
SOUTHERN MEDICAL	Date Issued: 2018.08.30	CE0086				
Manufactured by:     European R       Southern Medical (Pty) Ltd     Southern Imp       Building 10,     Building 3, C       Southern Implants Office Park,     566 Chiswidi       1 Albert Road,     Irene, 0062       South Africa     Tel: +27 12 667 6243/4		I Representative: Implants UK, Inc. , Chiswick Park vick, High Road /4 5YA, United Kingdom		Point of use Containment and transportation Preparation for cleaning	Thorough cle     These practic     surfaces, join     No particular     Devices capa     Dried-on soil     The removal	aning and rinsing should begin as soon as po zes include keeping devices moist after use ts, and all other hard-to-clean design features requirements. Ible of disassembly must be disassembled pri is difficult to remove with automatic washing of gross soil from these areas prior to washing
IMPORTANT: PLEASE READ For detailed information on the Southern AXIS™ 2 Description: The AXIS™ 2 Cervical Plate and the fixed and variab RADIOACTIVITY WARNING: No radioactivity substance or radioac Intended purpose: The Southern AXIS™ 2 Anterior Cervical Plate thus intended for immobilization and stabilization of the cervical spi level plates with lengths from 19.5 mm up to 83 mm. It is intended	Cervical Plate and Screw System, ple le angle screws are manufactured from ctivity. e System (ACP) arthrodesis is intended ine. The plate is used in conjunction wit to be used for treatment of degenerative	ase consult the Surgical Manual Grade 23 Titanium (ASTM F136). for cervical fusions thus relieving painfu n a bone graft, interbody or vertebrector e disc disease and trauma in the cervica	I motion of the vertebrae. The AXIS™ 2 is ny cage. The range consists of one to four I soine. The device consists of a plate that	Cleaning: Automated	Pre-cleaning     Rinse with wa     Load the inst retain liquid c     Run the auto     Phase     2 Rinse col     5Wash I de     13 Rinse w     16 Disinfec	should be through soaking in enzymatic deter arm water ruments in the washer such that all design fe an drain (for example, hinges should be open matic wash cycle: Minimum cycle parameters: d + warm water tergent arm water tion temperature is set at A0=600: 90°C (+1°C
<ul> <li>is fixated to the vertebral body by means of anchoring screws.</li> <li><i>Indications for the use of arthrodesis cervical plates and screw</i></li> <li>Deformity (defined by kyphosis, lordosis, or scoliosis)</li> <li>Degenerative spondylosis (as defined by neck pain of discogeni</li> <li>Failed previous fusions</li> <li>Pseudarthrosis</li> <li>Trauma including fractures</li> <li>Tumors</li> <li><i>Contraindications for the use of arthrodesis cervical plates and</i></li> <li>Failure of anterior plate fixation</li> <li>Infection</li> <li>Injury to the jugular vein, carotid artery, trachea, laryngeal nerve</li> <li>Injury to the oesophagus and pharynx from retractor traction perforation can lead to rapid infection, sepsis, hypotension, shood</li> </ul>	<i>vs are:</i> ic origin with degeneration of the disc co <i>d screws are:</i> e, thyroid gland or direct injury resulting in dysphagia ck and death.	nfirmed by patient history and radiograp	hic studies.) ses frank pharangeal and oesophageal	Cleaning: Manually	<ol> <li>Check instrur</li> <li>Long narrow</li> <li>Repeat clean</li> <li>Thermal disir</li> <li>Immerse inst</li> <li>Blind holes si</li> <li>Long narrow</li> <li>Use cleaning</li> <li>Scrub interfaa</li> <li>open the part</li> <li>Scrub inside ensure that fit</li> <li>Scrub arounce</li> <li>Scrub arounce</li> <li>Rinse thorout</li> </ol>	nents for visible soil cannulations and blind holes require particula ing if soil is visible and re-inspect. ifection Minimum cycle parameters: one (1) m rument and soak for a minimum of ten (10) m hould be repeatedly filled and emptied cannulations and blind holes require particula brushes/pipe cleaners to remove additional s zes several times using a twisting action. If co in order to access and clean these areas. holes with a tight-fitting brush or pipe cleaner all depth of the feature is reached I hinged surface areas with a brush or pipe cleaner ghly with warm water, making sure to wet the
Faitigenic risk to structures encountered during surgical approact     Severe osteophytes     Titanium/Titanium alloy allergy or intolerance     Risks Associated with surgery     Allergic or other reaction to anesthesia     Approach related injury     Blood loss or hemorrhage     Cerebrospinal fluid leak (CSF)     Death     Disease progression	<ul> <li>Infection</li> <li>Injury to oeso</li> <li>Injury to the ju</li> <li>Malnutrition</li> <li>Morbidity</li> <li>Osteolysis or</li> </ul>	ohagus and pharynx igular vein, carotid artery, trachea, laryn vertebral inflammation	geal nerve, thyroid gland	Disinfection: Drying Maintenance, Inspection and Testing of Instrumentation	10. Check instrum         11. Repeat clean         Disinfectant s         3% hydrogen         When drying         Visually inspe         Hinged instru         Locking mecl         All surfaces s         Reamer/drill	nents for visible soil ing if soil is visible solution (EndoZyme®) may be used in accord peroxide may be used on difficult to reach ar is achieved as part of a washer disinfector cyr act for damage or wear ments should be check for smooth movement hanisms should be checked for action hould be smooth and free of cracks and deep bits should be inspected for deformities and di
<ul> <li>Dyspriagia</li> <li>Failed fusion</li> <li>Hoarsness and swallowing difficulties</li> </ul>	<ul> <li>Pullionary en</li> <li>Sepsis</li> <li>Shock</li> </ul>	looism		Packaging	Metal surface     Blunt or dams     Instrument tra	Inspect for corrosion and major deformation aged instruments should be returned to sales ay should be double wrapped with Steriwrap.
<ul> <li>Hypotension</li> <li>Risks Associated with Cervical Spinal Systems <ul> <li>Annular ossification</li> <li>Bone graft migration</li> </ul> </li> <li>Degenerative changes in adjacent segment</li> <li>Dural injury</li> <li>Facet joint deterioration</li> <li>Failed back syndrome</li> <li>Fracture/bending of plate and/or screw</li> <li>Hematoma or Seroma</li> <li>Heterotopic ossification</li> <li>Hypopharyngeal screw migration</li> <li>Implant toelarsed or subsidence into adjacent vertebrae</li> <li>Implant degradation</li> <li>Myocardial infarction</li> <li>Nerve damage</li> <li>Neurologic deterioration; clumsiness, foot drop, limp, short stepslow moving gait, weakness</li> <li>Numbness</li> </ul>	<ul> <li>Inrombosis</li> <li>Perineural fibrosis</li> <li>Plate size may not be incluselection.</li> <li>Reflex sympathetic dystrophy</li> <li>Removal of the device in the p</li> <li>Reoperation at the study treat modification of any or all cor</li> <li>Revision with or without replace</li> <li>Screw back out</li> <li>Spinal instability</li> <li>Spinal stenosis</li> <li>Spondylolisthesis acquisita</li> <li>Spondylolisthesis acquisita</li> <li>Spontaneous fusion</li> <li>Sterility</li> <li>Supplemental fixation</li> <li>Transitional syndrome</li> </ul>	usive of range/incorrect plate length (RSD) ost-op or follow-up period tment level with or without removal or nponents of the device ement of a component/device	<ul> <li>Vessel damage</li> <li>Wear debris generation</li> <li>If re-used:</li> <li>Infection</li> <li>Non-oesteointegration</li> <li>Transmittable diseases</li> </ul>	Sterilization Instructions	Instruments and i Instruments and i It is importar performed pu Use Long narrow during cleani- Instrument st Sterilization t prior to sterilization t prior to sterilization t prior to sterilization t prior to sterilization t Do not stack Limitations on rej Repeated pro Recommended S Instrument se (270°-275°F) Some health Jakob diseas temperature i	nstrument trays: It that adequate cleaning of instrument case ior to sterilization. Preparation for Sterilizat cannulations and blind holes require particula ng. Iould be placed in Instrument Trays prior to st rays must be wrapped with an approved auto lization. The tray by itself does not provid a sterilization trays in the sterilizer! Drocessing: Drocessing: Drocessing has minimal effect on these instrument terilization Parameters: ats produced by Southern Medical can be steril 12 minute exposure, 30 minute vacuum dry, care authorities recommend sterilization acc e, especially of surgical instruments that could and time is required: 134°C, 20 minutes exposent
Osteophyte resorption     Osteophyte resorption     Procedure:     Refer to the surgical procedure provided by Southern Medical (Pty) Ltd.			Storage The instructions provided ab	If stored betworks the stored betworks th	veen cleaning and sterilization, dry instrumen et storage. devices can be stacked for storage. ved by Southern Medical. It remains the res	
<b>USAGE WARNING:</b> Improper technique in implant placement can result in implant failu used for the placement of the implant. Placement of this device implantable devices (screws and plates).	re. Surgical instrumentation is provided is limited to qualified surgeons. Refer t	I for specific use with the implant and n o surgical procedure and product brock	o other instrumentation is intended to be nure for more information. Do not reuse	Post Implantation: Movemen Description of Symbols Use	nt of the operation site v ed in Packaging	vill be restricted according to the discretion of
STERILITY: Plates All implant plates are supplied sterils, and	are for single use only before the label	ad expiration date. Do not re-use implor	ts. Do not use implants if the packaging	Use by		Do not reuse (implant devices)
An implaint plates are supplied stellle, and has been damaged or previously opened, degradation and could result in surgical i implanted. Re-use or re-implantation may i	or if the expiration date has passed. D rejection and/or post-operative infection result in cross-contamination or infection	o not re-sterilize implants provided sterilize implants provided sterilize implants provided sterilize implant is designed for single p . If uncertain be sure to contact a South	ile. Re-sterilization could cause material vatient use only and must never be re- ern Medical Representative	Sterilization (Gamma)		Manufacturer address
Screws Cleaned Implant screws are packed onto in is the responsibility of the hospital to ensur	strument sets as non-sterile and require	sterilization by an ISO 17665 validated	steam sterilization (autoclave) method. It aning and sterilization of the instruments	Do not resterilize	STERNUZE	Do not use if packaging is damaged

Cleaned Implant screws are packed onto instrument sets as non-sterile and require sterilization by an ISO 17665 validated steam sterilization (autoclave) method. It is the responsibility of the hospital to ensure equipment and cycles are validated on site. Personnel responsible for the cleaning and sterilization of the instruments must be a fully trained hospital staff member. Screws provided separate to the screws on instrument sets are supplied sterile and are for single use only before the labeled expiration date. Unless marked sterile and clearly labeled as such in an unopened sterile package, all implants and instruments used in surgery must be sterilized by the hospital prior to use. Remove all packaging materials prior to sterilization. Sterilization must be done in time before implanting the prosthesis. Only sterile products should be placed in the operative field. Unless specified elsewhere, these products are recommended to be steam sterilized by the hospital using one of the sets of process parameters below. If uncertain contact a Southern Medical representative.

# Magnetic Resonance Imaging (MRI)

The AXIS™ 2 devices have not been evaluated for adverse effect under MRI. The AXIS™ 2 implants are manufactured from non-ferromagnetic materials. Risks of placing implants in or near a magnetic field include: (1) movement of ferromagnetic components, (2) localised heating of components caused by radio frequency induction heating and (3) image artifacts created by interaction between metallic components and the magnetic field.

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## ssible after use of the device

to prevent soil from drying and removing gross soil from the surfaces, crevices, mating s.

### ior to cleaning

g, especially at challenging design features on devices like interfaces, crevices, joint etc. ig in the automatic washer is critical for achieving adequate cleaning irgent for 5 minutes and scrub surfaces including brackets and hinges with cleaning brush

eatures of the device are accessible to cleaning and such that design features that might and cannulations and holes positioned to drain).

	Function	Duration (min)
	Rinse, mixed water	2
	Wash with detergent	7
	Rinse, warm water	2
:)/1 min	Disinfection	1 + heating

#### r attention during cleaning

#### ninute at 91° C

nutes in enzymatic detergent

## r attention during cleaning

soil from challenging design features

omponents of the instrument can be disassembled or moved, it is necessary to retract or

using a twisting action. The brush or pipe cleaner should be of an appropriate size to

#### eaner

challenging design features

### ance with label instructions.

reas <u>cle do not exceed 120°</u>C.

t of hinge without excessive "play."

nicks

listortion that might hinder insertion into a drill

representatives

	Devices Or	nly:
es/trays be ion Single-	•	Screws are supplied non-sterile unless otherwise specified on the screw packaging. Prior to sterilization of the device, remove all original
ar attention		packaging and labeling inserts. Place the device in suitable packaging for the sterilization process, i.e., central supply
erilization		wrap, autoclave pouches, etc.
clave wrap le a sterile	•	Special care should be taken to protect the device from contact with other metal or hard objects that could damage the implant
	•	Packaging should be inspected for punctures or other damage before and after sterilization

ents. End of life is normally determined by wear and damage due to use.

erilized to a sterility assurance level (SAL) of 10E-6 in a pre-vacuum, 4 pulse, 132º-135ºC steam sterilization cycle

cording to these parameters to minimize the potential risk of transmission of Creutzfeldtd come into contact with the central nervous system. Should this be required, the following sure, 30 minute drying time. Its with a low-linting, non-abrasive soft cloth to prevent microbial contamination that could

is with a low-liniting, non-abrasive sort cloth to prevent microbial contamination that could

sponsibility of the processor to ensure that the reprocessing as actually performed using rmally requires validation and routine monitoring of the process.

NON	Consult the Instructions For Use	Ĩ
$\otimes$	European Representative	EC REP
	Date Of Manufacture	M
$\bigotimes$	Caution	$\wedge$