|   | IFU-002-05<br>Date Issued: 2024.04.30  |   | INSTRUCTION FOR USE  |  |                    |                              |        |  |  |  |   |   |  |  |  |
|---|--|---|--|--|--------------------|------------------------------|--------|--|--|--|---|---|--|--|--|
| SOUTHERN MEDICAL  |  |   | PEEK CERVICAL FUSION CAGES   | Page 1 of 1  |                    |                              |        |  |  |  |   |   |  |  |  |
| Manufactured by:<br>Southern Medical (Pty) Ltd.<br>55 Regency Drive,<br>Route 21 Corporate Park,<br>reap. Conturingo, OR2, South Africa.  | O Box 17198<br>yttleton, 0140<br>outh Africa<br>el: +27 12 667 6243/4<br>mail: <u>info @southmed.co.za</u>                 | <ul> <li>RISK &amp; UNDESIRA</li> <li>Damage to the view of surgery.</li> <li>Soft tissue injury.</li> </ul>  | BLE SIDE-EFFECTS:<br>ertebral endplates and/or fracture of the vertebrae.<br>s pulposus, disc disruption, or degeneration at, above, or below the  | Autograft or allogenic bone graft should be used in combination with the device. Failure to use bone graft may result in non-union. In the absence of osseous fusion, mechanical failure of the implant can be expected as a result of everyday mechanical stresses.   |                    |                              |        |  |  |  |   |   |  |  |  |
| IMPORTANT: PLEASE READ<br>For detailed information on correct & safe device use, please consult the<br>Surgical Manual.   |  | <ul> <li>Esophageal injury/perioration.</li> <li>Dural tears, persistent CSF leakage, or meningitis.</li> <li>Bone loss or decrease in bone density.</li> <li>Graft site complications (if autograft is used).</li> <li>Hemorrhage, hematoma, occlusion, seroma, edema, embolism, stroke, excessive bleeding, phlebitis, damage to blood vessels, or cardiovascular system compromise.</li> <li>Infection.</li> <li>Foreign body (allergic) reaction to implant.</li> <li>Post-operative change in spinal curvature, loss of correction, height and/or reduction.</li> <li>Loss of neurological function including paralysis, radiculopathy, and/or the development or continuation of pain, numbness, spasms, or sensory loss.</li> <li>Transient or permanent neurological deficits, reflex deficits, irritation, and/or muscle loss.</li> <li>Scar formation possibly causing neurological compromise around nerves and/or pain.</li> <li>Non-union (or pseudarthrosis), delayed union or mal-union.</li> <li>Disassembly, bending, and/or breakage of the implant.</li> <li>Implant loosening and/or migration.</li> <li>Subsidence of the device into the vertebral body(ies).</li> <li>Transient or persistent dysphagia (swallowing difficulties).</li> <li>Sore throat and/or hoarseness.</li> <li>Development of respiratory problems.</li> <li>Loss of spinal mobility or function.</li> <li>Wound necrosis or wound dehiscence.</li> <li>Inability to perform the activities of daily living.</li> </ul> |  | There are no special storage instructions. Storage conditions must not lead to degradation or contamination of the packaging or its contents. Handle with care.  POST-OPERATIVE CARE INSTRUCTIONS The surgeon/physician should provide appropriate postoperative care instructions to the patient and should ensure that the patient understands the potential consequences of non-compliance. To achieve the desired clinical outcome, the patient should be instructed to minimize physical activity as far as possible and avoid activities that require lifting, bending, twisting or excessive movement. Follow-up consultations should be arranged with the patient at appropriate intervals, as deemed necessary by the physician. The patient should only resume their normal activities once cleared to do so by their physician. The patient should only resume their normal activities once cleared to do so by their physician. The spleCCA devices have not been evaluated for safety in the MR environment. The safety of the SPICCA devices in the MR environment is unknown. Performing an MR exam on a person who has this medical device may result in injury or device malfunction.  DEVICE REMOVAL Surgical removal of the device is possible. A decision to remove the device must be made by a healthcare professional after taking into consideration the patient's general medical condition and the potential risk of a subsequent surgical procedure. Refer to the Surgical Manual for removal instructions. |                    |                              |        |  |  |  |   |   |  |  |  |
| <b>DESCRIPTION:</b><br>The SPICCA (Southern Cervical Fusion Cages for Anterior Placement) devices are<br>vedge-shaped, PEEK (ASTM F2026) anterior cervical interbody fusion cages. The<br>devices are made available with Tantalum (ASTM F560) radiographic markers and some<br>device variants are made available with a Titanium Coating (ASTM F1580). The cages<br>have at least one central "bone pocket" that is intended to be filled with autograft or<br>allogenic bone graft. The SPICCA, SPICCA-2 and SPICCA-F devices are intended to be<br>used in combination with the AXIS anterior cervical plate system, while the SPICCA-S2<br>and SPICCA-SP devices are stand-alone devices that do not require supplemental<br>ixation.<br><b>NDICATIONS FOR USE:</b><br>The SPICCA, SPICCA-F Cervical Fusion Cages are cervical interbody fusion<br>devices intended for spinal fusion procedures at one or two levels from the C2/C3 disc<br>space to the C7/T1 disc space in skeletally mature patients with degenerative disc disease<br>defined as neck pain of discogenic origin with degeneration of the disc confirmed by<br>history and radiographic studies) of the cervical spine. Implants are to be implanted via an<br>open, anterior approach and packed with autograft or allogenic bone graft comprised of<br>ancellous and/or corticocancellous bone graft. Patients must have undergone a regimen<br>of at least six (6) weeks non-operative treatment prior to being treated with these devices.<br>The implant is intended to be used in combination with an anterior cervical plating system.  |  |   |  |  |                    |                              |        |  |  |  |   |   |  |  |  |
|   |  |   |  |  |                    |                              |        | he SPICCA-SP & SPICCA-S2 Stand-Alone Cervical Fusion Cages are stand-alone<br>terbody fusion devices intended for spinal fusion procedures at one or two levels from<br>e C2/C3 disc space to the C7/T1 disc space in skeletally mature patients with<br>agenerative disc disease (defined as neck pain of discogenic origin with degeneration of<br>e disc confirmed by history and radiographic studies) of the cervical spine. Implants are<br>be implanted via an open, anterior approach and packed with autograft or allogenic<br>one graft comprised of cancellous and/or corticocancellous bone graft. Patients must<br>ave undergone a regimen of at least six (6) weeks non-operative treatment prior to being<br>eated with these devices. The implant is designed to accommodate two screws. Two<br>crews should be used to ensure adequate fixation of the implant. |  | Charge in mental status.     Death.     Additional surgical intervention may be required to correct/prevent some of these     possible adverse events.     WARNINGS & PRECAUTIONS: STERILITY, PACKAGING & RE-USE |   | DISPOSAL<br>Single-use devices that have been in contact with blood or bodily fluids/tissues should be<br>disposed of in accordance with the hospital's procedure for disposal of hazardous and/or<br>biological waste. Users must wear surgical gloves and take care to avoid sharp edges.<br>Devices that cannot be used because the packaging has been damaged, but have not<br>— been in contact with blood or bodily fluids/tissues, should be returned to the manufacturer. |  |  |  |
|   |  |   |  |  |                    |                              |        |  |  | STERILE R Ma   | e devices are sterilized via gamma irradiation and are provided<br><b>ERILE</b> . Do not re-sterilize the device. Re-sterilization could cause<br>terial degradation and could result in mechanical failure of the device,<br>st rejection and/or post-operative infection. | <b>REPORTING</b><br>Any device-related adverse events must be reported to the manufacturer as soon as possible.   |  |  |  |
| DNTRAINDICATIONS:<br>Patients with known or probable intolerance to the third double  | he materials used in the manufacture   |   | not use implants if the packaging has been damaged or previously<br>ened, or if the expiration date on the label has been exceeded.  | DESCRIPTIONS OF SYME<br>MANUFACTURER   |                    | IN PACKAGING<br>USE BY       | $\sum$ |  |  |  |   |   |  |  |  |
| <sup>a</sup> atients with infection, inflammation, fever, tumor<br>regnancy, mental illness and other medical<br>peneficial surgical outcome  | rs, elevated white blood count, obesity, conditions which would prohibit a   | The imp<br>in c   | e devices are <b>SINGLE-USE</b> . Do not re-use implants. An explanted<br>blant must never be re-implanted. Re-use or re-implantation may result<br>cross-contamination or infection.  | CONSULT INSTRUCTIONS<br>FOR USE  |                    | CATALOGUE NUMBER             | REF    |  |  |  |   |   |  |  |  |
| Patients resistant to following post-operative re<br>thletic and occupational activities.<br>Jse with components from other manufacturers.<br>Grossly distorted anatomy caused by congenital  | estrictions on movement especially in _ abnormalities.   | WARNINGS & PRE<br>Thi<br>kno  | CAUTIONGS: CORRECT & SAFE USE<br>e device should only be implanted by qualified surgeons with<br>bwledge of the correct and safe use of the device and associated  | DO NOT RE-STERILIZE  |                    | LOT NUMBER                   |        |  |  |  |   |   |  |  |  |
| Rapid joint disease, bone absorption or oste  | eopenia. Osteoporosis is a relative he degree of obtainable correction or  | ins<br>Me<br>Re   | trumentation. Refer to the Surgical Manual provided by Southern<br>dical (Pty) Ltd. If uncertain, please contact a Southern Medical<br>presentative.   | DATE OF MANUFACTURE<br>DO NOT USE IF PACKAGING<br>IS DAMAGED   |                    | STERILIZED USING IRRADIATION |        |  |  |  |   |   |  |  |  |
| ontraindication since this condition may limit the tabilization, and/or the amount of mechanical five neurophysical five components select the implant components select s                      | xation.<br>ted for use would be too large or too   | 110   |  |  | <u> </u>           |                              |        |  |  |  |   |   |  |  |  |
| contraindication since this condition may limit the<br>stabilization, and/or the amount of mechanical fix<br>Any case where the implant components select<br>small to achieve a successful result.<br>Any patient having inadequate tissue coverage<br>pone stock or quality.<br>Any patient in which implant utilization would in<br>supported having language for the support of t | kation. ted for use would be too large or too over the operative site or inadequate nterfere with anatomical structures or | The<br>in use<br>not  | e SPICCA, SPICCA-2 & SPICCA-F devices are intended to be used<br>combination with the AXIS anterior cervical plate system. Failure to<br>e supplemental fixation may lead to mechanical failure of the device,<br>n-union and/or cage migration/expulsion. | CAUTION: FEDERAL LAW<br>RESTRICTS THIS DEVICE TO<br>SALE BY OR ON THE ORDER<br>OF A PHYSICIAN  | R <sub>Xonly</sub> |                              |        |  |  |  |   |   |  |  |  |