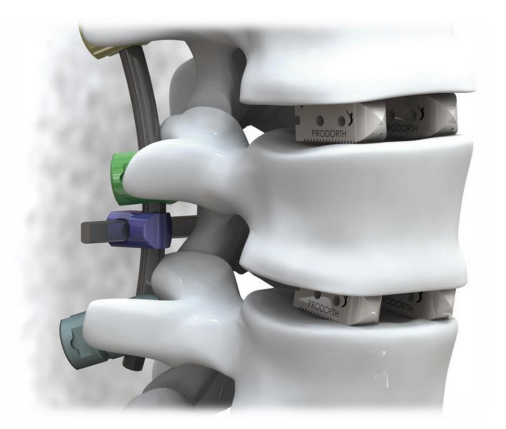


Surgical Technique Guide

PLIF PEEK CAGE

PROCORPTM





INTRODUCTION TO SURGICAL TECHNIQUE

Prodorth has developed this surgical technique document for surgeons and healthcare professionals but not for unauthorized persons. This document is a supportive source but not a complete instruction for an unexperienced surgeon to perform the entire surgery, therefore the information within this surgical technique should be considered with the previous medical experiences and education of the surgeon. Surgeon's medical judgement and decisions will be best treatment for the patient and the results will be different according to the patient's physical and mental situation.

DEVICE DESCRIPTION

PRODORTH PLIF CAGE is an implantation system used for disposing the complaints of the patients which are risen from the due to the herniation at discs, traumas or any disorders on lumbar spine.

Prodorth *PLIF* Cage Implants are long-term implants, however they are not able to withstand the forces like healthy bone structures.

- Current Status of the Device: Device is already CE marked (since 2013) and has been on the market.
- TLIF Cage GMDN No: 38161
- Product Class: Annex II of Directive 93/42/EEC) Class IIb
- Raw Materials: Ti6Al4V-ELI (ASTM F 136 / ISO 5832-3) and PEEK (ASTM F 2026)
- Biological Assesment:

Biological Assessment of Device According to TS EN ISO 10993-1 : 2011		
Category	Implant Device	
Contact Level	Bone/ Tissue	
Contact Duration	C (Permanent - > 30days)	=

• Sterilization:

Prodorth PLIF Cage is released to market as non-sterile.

They must be sterilized prior to surgical use. The validated sterilization method for PRODORTH implants is steam sterilization in autoclave. The implants which are intended to be sterilized should remain in autoclave at 134 C degree for 18 minutes. There is no other sterilization method PRODORTH recommends.

• Intended Use of the Device:

PRODORTH *PLIF* Cage is a long-term implant in order to dispose the complaints of the patients which raised because of the pain arising from the herniation at lumbar discs, traumas on lumbar spine.

- It is a single use device
- o Does not include human or animal tissue and phthalate
- \circ $\,$ Does not include any software oraccessory,
- The product is supplied as non-sterilized,
- \circ $\,$ Product does not cause any radioactive source or beam diffusing
- Population: Skelatally mature male / female patients
- Intended User(s): Healthcare professionals (Surgeons trained and experienced in the related field.)

INDICATIONS:

General criteria and principles related to instrumented spinal surgery are applied here:

Degenerative disc pathologies Herniated nucleus pulposus Grade 1 degenerative or isthmic spondylosis Visible loss of disc height compared to adjacent levels Lumbar pseudarthrosis

CONTRAINDICATIONS :

Fracture, tumor Osteoporosis. Calcium metabolism disorder Pregnancy Infection. Recognized allergies to titanium or titanium alloys Damaged lumbar vertebrae from an accident (trauma) at the level of the surgery. An unhealthy shape (deformity) of the lumbar vertebrae at the level of the surgery. Low bone mineral density, such as osteoporosis or osteopenia Mental disability

SECONDARY AND POSSIBLE SIDE EFFECTS:

Pseudarthrosis Implant penetration, migration or Implant failure Infection Allergy to materials used Dysphagia Loosening Increased neck pain Instability Hematoma c7 palsy Wound infection Hoarseness HO (heterotopic ossification) Displacement of the disc adjacent segment degeneration

WARNINGS:

Never re-use an implant even in perfect state. Any Implant which has been used, twisted, bent, implanted and then removed even If it appears intact, it must be discarded. Use new implants routinely. Similar products of competitors shall not be combined with the components of the Prodorth PLIF Cage. PRODORTH implants and instruments should only be used with PRODORTH instruments. Instruments developed by PRODORTH to be used in spinal surgeries of its spinal products. Incase of using other company's instruments, this might result in galvanic corrosion, incompatibility between the products as well. No component of the Prodorth lumbar Disc implants shall be reused.

The restricted shelf life of the device is 10 years. It should never be used after its expiration date.

Prodorth PLIF Cage Description

Unstable situation of lumbar spine causes herniation of discs and discs make a pressure to nerves. In order to solve this problem, PLIF Cage is positioned at intervertebral area after discectomy. So that 2 vertebras work as one vertebra as a result of bone fusion. By this way the pressure to nerves is eliminated and patient has a relief.

This product is manufactured from PEEK (Polyether-ether-kethon/ ASTM F2026) which is a polymer based composite material.

Prodorth PLIF Cage is made of PEEK (Polyether-ether-kethon/ ASTM F2026) which is a polymer based composite material and Ti6Al4V (grade 5) material.



PRODOLPHIN[®] PLIF Cages provides an uninterrupted guidance during operation. Circular toothed surface feature for holding the inferior and superior areas.

Made of PEEK material, originated from EVONIK Industries Germany. X-ray marker pins for the visibility.

Prodorth PLIF CAGES

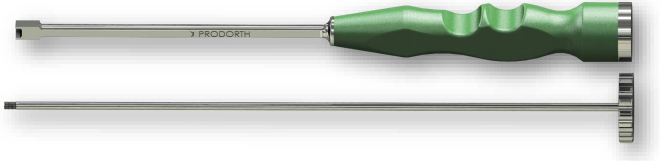
Sizes & Ref. Codes

Product Name / Sizes	Ref. Code
PRODORTH PLIF PEEK CAGE 25x7 mm	102.03 002507
PRODORTH PLIF PEEK CAGE 25x7 mm	102.03 002508
PRODORTH PLIF PEEK CAGE 25x9 mm	102.03 002509
PRODORTH PLIE PEEK CAGE 25x9 mm	102.03 002510
PRODORTH PLIF PEEK CAGE 25x10 mm	102.03 002511
PRODORTH PLIF PEEK CAGE 25x12 mm	102.03 002512
PRODORTH PLIF PEEK CAGE 25x12 mm	102.03 002513
PRODORTH PLIF PEEK CAGE 25x13 mm	102.03 002513
	102.03 002515
PRODORTH PLIF PEEK CAGE 25x15 mm	102.03 002313
	102 02 002007
PRODORTH PLIF PEEK CAGE 28x7 mm	102.03 002807
PRODORTH PLIF PEEK CAGE 28x8 mm	102.03 002808
PRODORTH PLIF PEEK CAGE 28x9 mm	102.03 002809
PRODORTH PLIF PEEK CAGE 28x10 mm	102.03 002810
PRODORTH PLIF PEEK CAGE 28x11 mm	102.03 002811
PRODORTH PLIF PEEK CAGE 28x12 mm	102.03 002812
PRODORTH PLIF PEEK CAGE 28x13 mm	102.03 002813
PRODORTH PLIF PEEK CAGE 28x14 mm	102.03 002814
PRODORTH PLIF PEEK CAGE 28x15 mm	102.03 002815

PRODORTH PLIF INSTRUMENTS

Prodorth offers different designs of instruments for each step of the surgical procedure. They have been designed as simple as possible and user-friendly in order to provide ease of use.

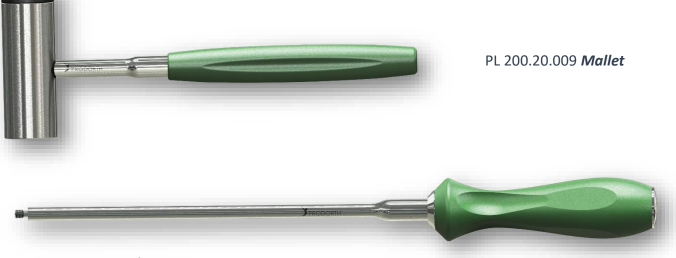
Stainless chrome nickel steel, aluminum, and silicone raw materials are used for Prodorth instruments.



PL 200.20.002 PLIF Cage Insertion







PL 200.20.010 Trial Insertion

PL 200.20.011 *T Handle Locking* ¹/₄"



*|8|8|8|____

PL 200.20.012 *Shaver 1/4'' Adaptör 7 mm* PL 200.20.013 *Shaver 1/4'' Adaptör 8 mm* PL 200.20.014 *Shaver 1/4'' Adaptör 9 mm* PL 200.20.015 *Shaver 1/4'' Adaptör 10 mm* PL 200.20.016 *Shaver 1/4'' Adaptör 11 mm*

SURGICAL PROCEDURE

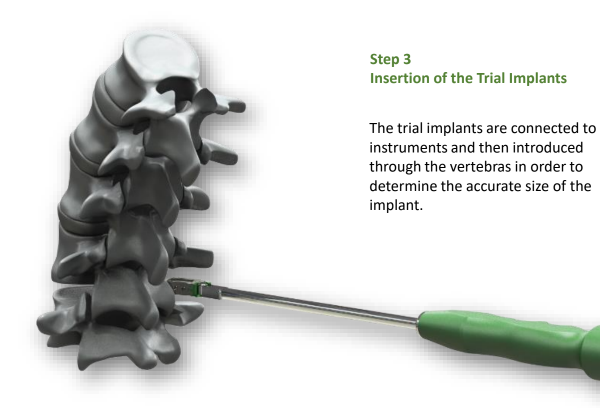
Step 1

The patient is positioned properly in the adequate position. The general essentials of lumbar surgery is applied.

Step 2 Preparation of the Disc Spaces

Use Prodorth Curettes in order to actualize the discectomy process. Remove the disc as required.





Step 4 Connection of the PLIF Cage with its inserter

The appropriately sized PLIF PEEK Interbody Cage is chosen. And it is firmly attached to the Inserter. In order to fix it accurately, the wheel behind the inserter is rotated clock-wise until assuring the implant is completely connected. And it's introduced into the intervertebral area.

Care should be taken to ensure the interbody spacer is aligned properly.



Step 5 Releasing the PLIF Cage from its inserter

After the PLIF Interbody Cage is placed, the extradural space and foramina are probed to ensure adequate decompression of the neural elements. And once it's decided the cage is positioned accurately, it's released by rotating the wheel of the inserter anti-clockwise.



RD Medikal Tıbbi Ürünler San. Tic. A.Ş. Karacaoğlan Mah. Bornova Cad. No:9G/1 Bornova İzmir / TURKEY Phone: 0090232 348 4950 (pbx) E-Mail: info@prodorth.com



See the IFU prior to use for additional information.



FR.4.2.3.9-2.3/02/04.02.2019