

D1104A
 (REV. 08/2023)

SECURE®-C CERVICAL ARTIFICIAL DISC



GLOBUS MEDICAL, INC.
 Valley Forge Business Center
 10000 Antietam Avenue
 Auburn, PA 19403 USA
 Customer Service:
 Phone 1-866-GLOBUS1 (OR)
 1-866-456-2873
 Fax 1-866-GLOBUS3 (OR)
 1-866-456-2873

IMPORTANT INFORMATION ON THE SECURE®-C CERVICAL ARTIFICIAL DISC

ENGLISH

IMPORTANT INFORMATION ON THE SECURE®-C CERVICAL ARTIFICIAL DISC

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

How Supplied

Implant Components – Sterile
Surgical Instruments – Non-Sterile (unless otherwise noted on the package label)

DESCRIPTION

The SECURE®-C Cervical Artificial Disc (SECURE®-C) is an articulating intervertebral device comprised of two endplates and a central core, and is inserted using an anterior cervical approach. The superior and inferior cobalt-chrome alloy (CoCrMo) per ISO 5832-12, ASTM F1537) endplates feature multiple serrated keels and a commercially pure titanium plasma sprayed (CoCrMo) per ISO 5832-2, ASTM F1537, F1147, and Co-6353) on the bone contacting surfaces. The sliding core is composed of ultra-high molecular weight polyethylene (UHMWPE per ISO 5834-2, ASTM F648), with a spherical superior interface and a cylindrical inferior interface articulating with the endplates. SECURE®-C implants are offered in a variety of configurations to accommodate varied patient anatomy. Implant footprints are as follows (AP depth x ML width): 11x2mm, 13x16mm and 15x16mm. SECURE®-C provides 7° or 0° lordosis options in the neutral position. Implant heights range from 9mm to 12mm, in 1mm increments. A list of SECURE®-C implants is provided in **Table 1**.

The SECURE®-C Cervical Artificial Disc is designed to allow motion in flexion and extension up to 30° (+15°), and in lateral bending to 20° (+10°). The design is intended to also allow unlimited axial rotation, and is constrained by ligaments and posterior elements. The device is also designed to permit translation of a ±1.25mm in the sagittal plane.

Part #	Description	Part #	Description
414.107S	SECURE®-C Core, 11x12, 7mm	414.312S	SECURE®-C Core, 14x16, 12mm
414.108S	SECURE®-C Core, 11x12, 8mm	714.100S	SECURE®-C Endplate Assembly, 11x12, 0°
414.109S	SECURE®-C Core, 11x12, 9mm	714.106S	SECURE®-C Endplate Assembly, 11x12, 6°
414.110S	SECURE®-C Core, 11x12, 10mm	714.166S	SECURE®-C Endplate and Core Assembly, 11x12, 6mm, 6°
414.111S	SECURE®-C Core, 11x12, 11mm	714.176S	SECURE®-C Endplate and Core Assembly, 11x12, 6mm, 0°
414.112S	SECURE®-C Core, 11x12, 12mm	714.200S	SECURE®-C Endplate Assembly, 13x14, 0°
414.207S	SECURE®-C Core, 13x14, 7mm	714.206S	SECURE®-C Endplate Assembly, 13x14, 6°
414.208S	SECURE®-C Core, 13x14, 8mm	714.266S	SECURE®-C Endplate and Core Assembly, 13x14, 6mm, 6°
414.209S	SECURE®-C Core, 13x14, 9mm	714.276S	SECURE®-C Endplate and Core Assembly, 13x14, 6mm, 0°
414.210S	SECURE®-C Core, 13x14, 10mm	714.300S	SECURE®-C Endplate Assembly, 14x16, 0°
414.211S	SECURE®-C Core, 13x14, 11mm	714.306S	SECURE®-C Endplate Assembly, 14x16, 6°
414.212S	SECURE®-C Core, 13x14, 12mm	714.366S	SECURE®-C Endplate and Core Assembly, 14x16, 6mm, 6°
414.307S	SECURE®-C Core, 14x16, 7mm	714.376S	SECURE®-C Endplate and Core Assembly, 14x16, 6mm, 0°
414.308S	SECURE®-C Core, 14x16, 8mm	714.400S	SECURE®-C Endplate Assembly, 15x16, 0°
414.309S	SECURE®-C Core, 14x16, 9mm	714.406S	SECURE®-C Endplate Assembly, 15x16, 6°
414.310S	SECURE®-C Core, 14x16, 10mm	714.466S	SECURE®-C Endplate and Core Assembly, 15x16, 6mm, 6°
414.311S	SECURE®-C Core, 14x16, 11mm	714.476S	SECURE®-C Endplate and Core Assembly, 15x16, 6mm, 0°

SECURE®-C devices are implanted using instruments specific to the device, as well as manual surgical instruments. Instruments specifically designed for implanting SECURE®-C consist of trays, milling guides, broaching chisels, keel chisels, chisel endcap, implant holding block, implant holders, and endplate positioners. Manual surgical instruments include instruments for cervical dissection, discectomy preparation, and milling.

INDICATIONS FOR USE

The SECURE®-C Cervical Artificial Disc is indicated in skeletally mature patients for reconstruction of the disc at one level from C3-C7 following single-level discectomy. Irretractable radiculopathy (arm pain and/or neurological deficit with or without neck pain), or myelopathy due to a single-level abnormally localized to the disc space and at least one of the following conditions confirmed by radiographic imaging (CT, MRI, X-ray): herniated nucleus pulposus, spondylosis (defined by the presence of osteophytes), and/or visible loss of disc height as compared to adjacent levels. The SECURE®-C Cervical Artificial Disc is implanted using an anterior approach. Patients should have failed at least six weeks of conservative treatment prior to implantation of the SECURE®-C Cervical Artificial Disc.

CONTRAINDICATIONS

The SECURE®-C Cervical Artificial Disc should not be implanted in patients with the following conditions:

- Active systemic infection or localized infection at the surgical site
- Unretractable radiculopathy or myelopathy due to pathology at more than one level and/or pathology not localized to the disc space
- Skeletally immature patients, pediatric or adolescent children (<21 years old), or those over the age of 60;
- Allergy or sensitivity to cobalt, chromium, molybdenum, titanium or polyethylene
- Marked cervical instability on neutral resting lateral or flexion/extension radiographs; translation >3mm and/or >11° rotational difference from that of either adjacent level
- Severe spondylosis at the level to be treated, characterized by bridging osteophytes, loss of disc height >50%, an absence of motion (<2°) as this may lead to a limited range of motion and may encourage bone formation (e.g., heterotopic ossification, fusion)
- Severe facet joint arthropathy
- Significant cervical anatomical deformity or clinically compromised vertebral bodies at the affected level due to current or past trauma (e.g., by radiographic appearance of fracture callus, malunion or nonunion) or disease (e.g., ankylosing spondylitis, rheumatoid arthritis)
- Any conditions attributed to more than one vertebral level

WARNINGS

The SECURE®-C Cervical Artificial Disc should only be used by surgeons who are experienced with anterior cervical spinal procedures and have undergone hands-on training in the use of this device. Only surgeons who are familiar with the implant components, instruments, procedure, clinical applications, biomechanics, adverse events, and risks associated with the SECURE®-C Cervical Artificial Disc should use this device. A lack of adequate experience and/or training may lead to a higher incidence of adverse events, including neurological complications.

Correct selection of the appropriate implant size and correct placement of the device are essential to ensure optimal performance and function of the device. Please refer to the SECURE®-C Cervical Artificial Disc Surgical Technique manual for step-by-step instructions on the required surgical technique, including determining the correct implant size.

Due to the proximity of vascular and neurological structures to the implantation site, there are risks of serious or fatal hemorrhage and risks of neurological damage with the use of this device. Care should be taken to identify and protect these structures during surgery.

Heterotopic ossification (HO) is a potential complication associated with cervical total disc replacement devices, which could result in reduced motion. It is recommended that bone wax is used following removal of HO bone formation. The short-term postoperative use of non-steroidal anti-inflammatory drugs (NSAIDs), is recommended to possibly reduce the chance of developing HO.

Anterior or posterior extension/migration of the SECURE®-C core or endplates have been reported in patients outside of the clinical study (<0.5%). These events have occurred in patients with instability, poor implant placement or sizing, or postoperative trauma. Proper patient selection, surgical technique and implant sizing should be used to help prevent extensor/migration.

PRECAUTIONS

The safety and effectiveness of this device has not been established in patients with the following conditions:

- Unretractable radiculopathy or myelopathy due to pathology at more than one level and/or pathology not localized to the disc space;
- Skeletally immature patients, pediatric or adolescent children (<21 years old), or those over the age of 60;
- Prior fusion at an adjacent vertebral level;
- Prior surgery at the level to be treated;
- Progressive symptoms and signs of spinal cord/nerve root compression with less than six weeks of conservative treatment;
- Facet joint disease or degeneration at the involved level;
- Neck or arm pain of unknown etiology;
- Neck pain alone;
- Page's disease, osteomalacia, or other metabolic bone disease;
- Rheumatoid arthritis or other autoimmune disease;
- Neuromuscular disorders such as muscular dystrophy, spinal muscular atrophy, amyotrophic lateral sclerosis;
- Severe insulin dependent diabetes;
- Systemic disease including AIDS, HIV, and Hepatitis;
- Taking medications known to potentially interfere with bone/soft tissue healing (e.g., steroids);
- Active malignancy (including spinal metastases);
- Acute mental illness or substance abuse; and
- Pregnancy.

Pre-operative

Patient selection is extremely important. In selecting patients for a cervical total disc replacement, the following factors can be of extreme importance to the success of the procedure: the patient's occupation or activity level; a condition of senility, mental illness, alcoholism or drug abuse; and certain degenerative diseases (e.g., degenerative scoliosis or ankylosing spondylitis) that may be so advanced at the time of implantation that the expected useful life of the device is substantially decreased. As in order to minimize the risk of perioperative vertebral fractures, surgeons must consider all co-morbidities, past and present medications, previous treatments, etc., and screening questionnaires for osteoporosis or osteopenia. SCORE (Simple Calculated Osteoporosis Risk Estimation), may be used to screen patients to determine if a DEXA bone mineral density measurement is necessary. If DEXA is performed, the patient should be excluded from receiving the device if the DEXA bone density measured T score is < -1.0, as the patient may be osteoporotic or osteopenic.

The patient should be informed of the potential adverse effects (risks/complications) contained in this insert (see POTENTIAL ADVERSE EFFECTS OF THE DEVICE ON HEALTH).

Preoperative planning may be used to estimate the required implant size, and to assure that the appropriate range of sizes is available for surgery. The procedure should not take place if the appropriate range of sizes will not be available.

Examine all instruments prior to surgery for wear or damage. Instruments which have been used excessively may be more likely to break. Replace any worn or damaged instruments.

Intra-operative

The SECURE®-C Cervical Artificial Disc implant should not be used with components or instruments of spinal systems from other manufacturers. Refer to the SECURE®-C surgical technique manual for step-by-step instructions.

Use aseptic technique when removing the SECURE®-C Cervical Artificial Disc implants from the innermost packaging. Carefully inspect each component and its packaging for any signs of damage, including damage to the sterile barrier. Do not use SECURE®-C implants if the packaging is damaged or the implant shows signs of wear.

Use care when handling the SECURE®-C Cervical Artificial Disc implant to ensure that it does not come in contact with objects that could damage the implant. Exercise care to ensure that implantation instruments do not contact the highly polished articulating surfaces of the endplates. Damaged implants are no longer functionally reliable.

To help prevent extension or migration of the SECURE®-C core or endplates, ensure that the SECURE®-C device is properly placed and correctly sized to fit the patient's anatomy.

To prevent unnecessary damage to the bearing surfaces, ensure that tissue or other debris is not trapped within the device.

Surgical implants must never be re-used or re-implanted. Even though the device appears undamaged, it may have small effects and internal stress patterns that may lead to early breakage.

When preparing the disc space, remove anterior or posterior osteophytes as needed, taking care to minimize bone removal. Avoid excessive bone removal as this may compromise the vertebral endplates or vertebral body. Care should be taken to perform careful cuts. Care should be taken to correctly position the trial during this step. Ensure proper alignment and placement of device components as misalignment may cause excessive wear and/or early failure of the device. Bone wax should be used to fill any exposed bleeding bone.

Post-operative

Patients should be instructed in postoperative care procedures and should be advised of the importance of adhering to these procedures for successful treatment with the device. Patients are recommended to wear a cervical collar for a few weeks following surgery, follow a therapy program for active range of motion exercises, and to avoid lifting, bending, twisting, or strenuous activities. The time period of these recommendations is managed by the treating physician. Patients should be instructed to avoid heavy lifting, bending, twisting, or strenuous activities. The time period of these recommendations is managed by the treating physician, taking into consideration the individual patient's condition as well as the stability and functioning of the implant. A two week postoperative course of non-steroidal anti-inflammatories (NSAIDs) is recommended to potentially reduce the incidence of heterotopic ossification (HO).

MR Safety Information

Non-clinical testing has demonstrated that the SECURE®-C implant is MR Conditional. A patient with this device can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5Tesla or 3Tesla only
- Maximum spatial gradient magnetic field of 4.000 gauss/cm (40T/m) or less
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2W/kg (Normal Operating Mode)

Under the scan conditions defined above, the SECURE®-C implant is expected to tolerate a maximum temperature rise of 1.4° after 15 minutes of continuous scanning. In non-clinical testing, the image artifact caused by the device extends approximately 20mm from the device when imaged with a gradient echo pulse sequence and a 3Tesla MR system.

POTENTIAL ADVERSE EFFECTS OF THE DEVICE ON HEALTH

The potential adverse effects (risks/complications) associated with the use of the SECURE®-C Cervical Artificial Disc include: (1) those associated with any surgical procedure; (2) those associated with anterior cervical spine surgery; and (3) those associated with a cervical artificial disc device, including the SECURE®-C Cervical Artificial Disc. However, the presence of these events does not exclude the device from being used. In addition to these risks, listed below, there is also the risk that surgery may not be effective in relieving symptoms, or may cause worsening of symptoms. Additional surgery may be required to correct some of the adverse effects.

- Risks associated with any surgical procedure include: abscess, cellulitis; wound dehiscence; wound, local, and/or systemic infection; wound necrosis; edema; hematoma; heart and vascular complications; hypertension; thrombosis; ischemia; embolism; thromboembolism; hemorrhage; thrombophlebitis; adverse reactions to anesthesia; pulmonary complications; organ, nerve, or muscular injury; gastrointestinal or genitourinary compromise; seizures, convulsion, or changes in mental status; complications of pregnancy including miscarriage and fetal birth defects; inability to resume activities of daily living; and death.
- Risks associated with anterior cervical spine surgery include: dysphagia; dysphasia; dysphonia; hoarseness; vocal cord paralysis; laryngeal palsy; sore throat; recurring epistaxis; nerve deficits or damage; tracheal, esophageal, or pharyngeal perforation; airway obstruction; external dysrhythmia; warmth or tingling in the extremities; deficit or damage to the spinal cord, nerve roots, or nerves possibly resulting in paralysis or pain; dorsal tears or leaking; cerebrospinal fistula; discitis, arachnoiditis, and other types of inflammation; loss of disc height; loss of proper curvature, correction; hemorrhage; or reduction of the spine; vertebral spilling; scarring, herniation or degeneration of adjacent discs; surrounding soft tissue damage, spinal stenosis; spondylolysis; otitis media; fistula; vascular damage and/or rupture; and headache.
- Risks associated with a cervical artificial disc device, including the SECURE®-C Cervical Artificial Disc, include: early or late loosening of the components; disassembly; bending or breakage of any or all of the components; implant migration; implant malpositioning; loss of purchase; sizing issues with components; anatomical or technical difficulties; implant fracture; bone fracture; bone fracture, spinal stenosis, or spinal cord injury; or pain; tissue or nerve damage caused by improper positioning or placement of implants or instruments; bending or breakage of a surgical instrument; and, as well as the possibility of a fragment of a broken instrument remaining in the patient; loss of neurological function; decreased strength of extensors; decreased reflexes; cord or nerve root injury; loss of bowel and/or bladder control or other types of urological system compromise; and interference with radiographic imaging because of the presence of the implant.

For the specific adverse events that occurred in the clinical study of the SECURE®-C Cervical Artificial Disc, please see the Safety Results in the CLINICAL STUDY section below. Some of the most common adverse events experienced by study patients were: neck and/or arm pain, dysesthesia, back and/or leg pain, musculoskeletal events (excluding spinal events), and difficulty swallowing.

CLINICAL STUDY

The clinical investigation of the SECURE®-C Cervical Artificial Disc was conducted under an approved IDE (G050075). The study was a prospective, multi-center, two-arm, randomized, unmasked, concurrently controlled, non-inferiority trial to assess the safety and effectiveness of the SECURE®-C Cervical Artificial Disc compared to anterior discectomy and fusion (ACDF) using a plate and structural allograft for reconstruction of the disc at one level from C3-C7 following single-level discectomy for irretractable radiculopathy or myelopathy due to a single-level abnormally localized to the disc space. The first five patients at each site were treated with the SECURE®-C device; subsequent patients were randomized in a 1:1 ratio to each site to receive either the SECURE®-C or control treatment. The purpose of the study was to determine whether the SECURE®-C Cervical Artificial Disc was non-inferior to the control.

Patients were treated between July 7, 2005 and April 25, 2008. A total of 380 patients were enrolled at 18 sites. Of these patients, 89 were non-randomly assigned to SECURE®-C. Of the randomized patients, 151 patients were randomized to SECURE®-C and 140 to control ACDF treatment. Final analysis was conducted after all patients had reached the two year time point; additional analysis was conducted after all patients had reached the seven year time point. The database for the PMA reflects data collected through July 9, 2015.

Clinical Inclusion and Exclusion Criteria

Patients were enrolled in the study according to the following inclusion/exclusion criteria:

Clinical Inclusion Criteria

Enrollment in the SECURE®-C study was limited to patients who met the following inclusion criteria:

- Symptomatic cervical disc disease (SCDD) in one vertebral level between C3-C7, defined as neck or arm (radicular) pain, or functional or neurological deficit and radiographic confirmation (CT, MRI, X-ray, etc.) of any of the following:
 - Herniated nucleus pulposus;
 - Radiculopathy or myelopathy;
 - Spondylosis (defined by the presence of osteophytes) or;
 - Loss of disc height.
- Age between 18 and 60 years
- Failed at least 6 weeks of conservative treatment
- Neck Disability Index (NDI) Questionnaire score of at least 30 (as percentage of 50 point total)
- Unable to understand and sign informed consent form
- Psychosocially, mentally and physically able to fully comply with this protocol including adhering to follow-up schedule and filling out forms
- Unable to meet the proposed follow-up schedule at 6 weeks, 3 months, 6 months, 12 months and 24 months
- Unable to follow postoperative management program

Clinical Exclusion Criteria

Patients were not permitted to enroll in the SECURE®-C study if they met any of the following exclusion criteria:

- More than one vertebral level requiring treatment
- Prior fusion surgery adjacent to the vertebral level being treated
- Prior surgery at the level to be treated
- Clinically compromised vertebral bodies at the affected level due to current or past trauma
- Prior interest in becoming pregnant in the next 2 years
- Active systemic or local infection
- Osteoporosis, osteopenia, Page's disease, osteomalacia or any other metabolic bone disease
- Pregnant or interested in becoming pregnant in the next 2 years
- Active systemic or local infection

- Known allergy to titanium, polyethylene, cobalt, chromium or molybdenum
- Taking medications or any drug known to potentially interfere with bone/soft tissue healing (e.g., steroids)
- Rheumatoid arthritis or other autoimmune disease
- Systemic disease including AIDS, HIV, Hepatitis
- Active malignancy (any type of cancer, including but not limited to non-melanoma skin cancer), unless he/she has been treated with curative intent and there has been no clinical signs or symptoms of the malignancy for at least 5 years
- Neuromuscular disorders such as muscular dystrophy, spinal muscular atrophy, amyotrophic lateral sclerosis, etc.;
- Acute mental illness or substance abuse
- Use of bone growth stimulator within past 30 days
- Participation in other investigational device or drug clinical trials within 30 days of surgery
- Pregnancy.

Postoperative Care

The recommended postoperative care included use of an external orthosis for 3 weeks postoperatively, followed by physical therapy program for active range of motion exercises. Patients were instructed to avoid lifting above shoulders for 3 months, and to avoid athletic activities and repetitive bending or lifting for 6 months. Smokers were encouraged not to smoke with their device. Patients were not treated with NSAIDs postoperatively in either treatment group. Some reports in the literature show that short-term postoperative use of NSAIDs may reduce the incidence of heterotopic ossification at some total disc replacement patients.

Follow-up Schedule

All patients were scheduled to return for follow-up examinations at 6 weeks (±2 weeks), 3 months (±2 weeks), 6 months (±1 month), 12 months (±2 months), 24 months (±2 months), and annually thereafter (±2 months), as shown in **Table 2**. All adverse events (device-related or not) were monitored over the course of the study and radiographic assessments were done by an independent core laboratory. All adverse events were independently adjudicated (for adverse event code, severity, and relationship to the device and/or procedure) by a Clinical Events Committee (CEC). At each evaluation time point, the following clinical and radiographic outcomes were evaluated per the clinical evaluation schedule below: the Neck Disability Index (NDI) to assess pain and function, Visual Analog Scale (VAS) to assess neck and arm pain, neurological status, general health status as assessed based on the SF-36, medication usage, patient satisfaction with surgery, radiographic parameters, and overall success. Radiographic included range of motion (flexion and translation), range of motion (flexion and translation), device displacement, radiolucency, fusion status, and heterotopic ossification. Overall success was determined based on data collected during the initial 24 months of follow-up.

Evaluation	Pre-op	Surgery/ Hospital Discharge	6 wks	3 mo	6 mo	12 mo	24 mo & annually
Neck Disability Index	X		X	X	X	X	X
Neck and Arm Pain (VAS)	X		X	X	X	X	X
Health Status (SF-36)	X		X	X	X	X	X
Neurological Status	X		X	X	X	X	X
Adverse Events*	X	X	X	X	X	X	X
Demographic/Baseline Data	X						
Operative Data	X						
Medication Use	X				X	X	X
Imaging Studies:							
AP & lateral	X	X	X	X	X	X	X
Lateral flexion/extension	X			X	X	X	X
CT and/or MRI	X						
Radiographic Outcomes:							
Range of motion	X			X	X	X	X
Disc height	X			X	X	X	X
Device migration			X	X	X	X	X
Fusion status				X	X	X	X
Radiolucency				X	X	X	X
Patient Satisfaction						X	X

*Adverse events with complications were recorded at all visits (both scheduled and unscheduled).

Clinical Endpoints

The safety of the SECURE®-C was assessed by comparing the nature and frequency of adverse events (overall and in terms of severity and relationship to the device and/or procedure) and secondary surgical procedures as well as maintenance or improvement in neurological status to the ACDF control group.

The effectiveness of the SECURE®-C was assessed by evaluating improvement in the Neck Disability Index (NDI), neck and arm pain based on a Visual Analog Scale (VAS), majority of life using the short-form 36 questionnaire (SF-36) as well as patient satisfaction compared to the ACDF control group.

In addition, several radiographic endpoints were considered in evaluating both safety and effectiveness, including degree of motion, disc height, device displacement or migration, radiolucency, spinal fusion status, and heterotopic ossification.

Per the protocol, an individual patient was considered a success if the following criteria were met at 24 months:

- Paradoxically improvement of at least 25% in NDI compared to baseline;
- No device failures requiring revision, removal, replacement, or supplemental fixation;
- Absence of major complications defined as major vessel injury, neurological damage, or nerve injury; and
- For control fusion patients only, radiographic fusion, as defined by the presence of bridging trabecular bone, without evidence of pseudarthrosis (defined radiographically as no apparent bridging trabecular bone and range of motion >3mm in translation and >2° in rotation).

In addition, FDA requested an additional analysis in which individual patient was considered a success if the following criteria were met at 24 months:

- Paradoxically improvement of at least 15 points in NDI compared to baseline;
- No secondary surgery at the index level including revision, removal, replacement or supplemental fixation;
- No potentially device-related adverse events;
- Maintenance or improvement in all components of neurologic status; and
- No SECURE®-C intraoperative changes in treatment.

Overall study success criteria were based on a comparison of individual patient success rates, such that the patient success rate for the SECURE®-C investigational group must be non-inferior to that of the ACDF fusion control group. The IDE study was approved using an off-informity major protocol (data) of 15% with an advisory that a non-inferiority test would be used to evaluate the success of the device's effectiveness. According to statistical analysis plan, if non-inferiority was demonstrated, then superiority would be evaluated as defined more specifically in the analysis plan. Of note, the statistical analysis plan pre-specified that the analysis technique would involve predicting 24 months outcomes for those without them, based on interim 6 month and 12 month observed outcomes, and integrating over the predictions to obtain posterior probabilities of non-inferiority and superiority.

Secondary effectiveness evaluations specified in the protocol included comparisons of the following:

- Components of the primary
 - Pain/Disability improvement (NDI)
 - No device failures requiring revision, re-operation or removal
 - Absence of major complications
- Neck Disability Index: 25% improvement from baseline
- VAS pain scales (neck, right, and left arm): 20mm improvement from baseline
- Health Status Survey SF-36 (mental and physical composite scores): 15% improvement from baseline
- Neurological status (maintenance, or improvement): proportion of patients maintained or improved
- Mean range of motion (angulation and translation)
- Disc height on standard lateral radiograph: 2mm changes compared to baseline
- No significant radiolucency for the SECURE®-C device; proportion of patients
- Spinal fusion in the control arm
- Patient satisfaction (definitively/improved): proportion of patients
- Device displacement or migration (>3mm)
- Operative time
- Operative blood loss
- Return to work

Accountability of PMA Cohort

A total of 380 patients at 18 sites were enrolled and treated in the IDE clinical trial: 236 received SECURE®-C (88 non-randomized, 148 randomized) and 144 received control treatment. All of the 380 patients enrolled in the PMA study have reached the 84 month post-operative visit: 331 (87.1%) had 24 month data available for analysis and 288 (75.8%) had 84 month data available for analysis at the completion of the study. Complete primary endpoint data (including fusion status for control group patients) was available for 215 investigational (77 non-randomized, 138 randomized) and 88 control patients at 24 months, and for 185 investigational (61 non-randomized, 124 randomized) and 101 control patients at 84 months. A total of 10 investigational (4 non-randomized and 6 randomized) and 22 control patients had failures at or prior to the 84 month visit. A summary of annual patient accountability data is provided in **Table 3**.

Number of Patients	12 Months (±2 Months)			24 Months (±2 Months)			36 Months (±2 Months)		
	NR SEC	R SEC	R ACDF	NR SEC	R SEC	R ACDF	NR SEC	R SEC	R ACDF
Enrolled	89	151	140	89	151	140	89	151	140
Theoretical	89	151	140	89	151	140	89	151	140
Deaths ¹	0	0	0	1	0	0	1	0	0
Failures ²	1	1	6	2	3	10	2	4	11
Not yet overdue	0	0	0	0	0	0	0	0	0
Expected ³	87	150	134	86	148	130	86	147	129
Actual, efficacy ⁴ (% Follow-up)	82	140	114	77	138	96	61	69	45
Actual, efficacy in window ⁵ (% Follow-up)	81	127	99	70	110	83	40	49	31
Actual, any data ⁶ (% Follow-up)	100	101	73	81.4%	74.3%	63.8%	46.5%	33.3%	24.0%
Actual, any data ⁷ (% Follow-up)	87	140	121	79	138	102	61	70	47
Actual, any data ⁸ (% Follow-up)	82	140	114	77	138	96	61	69	45
Actual, efficacy in window ⁹ (% Follow-up)	19	30	14	12	35	27	15	38	32
Actual, any data ¹⁰ (% Follow-up)	32	36	31	30	42	35	18	70	48
Actual, any data ¹¹ (% Follow-up)	37	46	24	36	48	28	18	70	48

</

Table 12. Statistical Comparison of Adverse Events (Randomized Patients As Treated)

Adverse Event	Patients Experiencing Adverse Events (%)				95% BCI (lower, upper)
	SEC (N=148)	ACDF (N=143)			
Any Adverse Event	132 (89.2%)	130 (90.9%)			(-8.7%, 5.3%)
Cancer	2 (1.4%)	4 (2.8%)			(-5.4%, 2.2%)
Cardiovascular	14 (9.5%)	4 (2.8%)			(-1.1%, 12.4%)
Carpal Tunnel Syndrome (CTS)	11 (7.4%)	13 (9.1%)			(-8.2%, 4.8%)
Cerebrovascular	1 (0.7%)	2 (1.4%)			(-3.9%, 2.2%)
Compressive Peripheral Neuropathy	4 (2.7%)	2 (1.4%)			(-2.4%, 5.1%)
Death	1 (0.7%)	2 (1.4%)			(-3.9%, 2.2%)
Dysesthesia - Lower Extremities	6 (4.1%)	4 (2.8%)			(-3.3%, 5.8%)
Dysesthesia - Other	3 (2.0%)	3 (2.1%)			(-3.9%, 3.7%)
Dysesthesia - Upper Extremities	21 (14.2%)	29 (20.3%)			(-14.7%, 2.6%)
Dysphagia	4 (2.7%)	10 (7.0%)			(-9.7%, 0.7%)
Dysphonia	1 (0.7%)	2 (1.4%)			(-3.9%, 2.2%)
Endocrinology	6 (4.1%)	5 (3.5%)			(-4.2%, 5.3%)
Gastrointestinal	12 (8.1%)	6 (4.2%)			(-1.8%, 9.7%)
Headache	14 (9.5%)	16 (11.2%)			(-8.9%, 5.4%)
Infection - Other	6 (4.1%)	3 (2.1%)			(-2.3%, 6.4%)
Infection - Superficial Wound	0	4 (2.8%)			(-6.4%, 0.1%)
Muscle Spasms	0	1 (0.7%)			(-3.9%, 1.5%)
Musculoskeletal	33 (22.2%)	21 (14.7%)			(-1.4%, 16.2%)
Neurological	4 (2.7%)	7 (4.9%)			(-7.1%, 2.4%)
Other*	11 (7.4%)	9 (6.3%)			(-4.9%, 7.1%)
Pain - Back and/or Lower Extremities	39 (26.4%)	46 (32.2%)			(-16.1%, 4.6%)
Pain - Neck	52 (35.1%)	58 (40.6%)			(-16.4%, 5%)
Pain - Neck and Upper Extremities	26 (17.6%)	36 (25.2%)			(-16.9%, 1.8%)
Pain - Neck and Upper Extremities with Dysesthesia	6 (4.1%)	8 (5.6%)			(-6.8%, 3.6%)
Pain - Neck with Dysesthesia	1 (0.7%)	6 (4.2%)			(-7.8%, 0.1%)
Pain - Other	2 (1.4%)	2 (1.4%)			(-3.4%, 3.2%)
Pain - Upper Extremities	40 (27.0%)	41 (28.7%)			(-11.9%, 8.6%)
Pain - Upper Extremities with Dysesthesia	7 (4.7%)	7 (4.9%)			(-5.4%, 0.5%)
Psychological	2 (1.4%)	1 (0.7%)			(-2.3%, 3.7%)
Respiratory	3 (2.0%)	1 (0.7%)			(-3.9%, 4.7%)
Surgery - Adjacent Level	4 (2.7%)	9 (6.3%)			(-8.6%, 1.3%)
Surgery - Index Level	6 (4.1%)	22 (15.4%)			(-18.2%, -4.7%)
Surgery - Lumbar Level	11 (7.4%)	8 (5.6%)			(-4.1%, 7.7%)
Surgery - Other Cervical	0	2 (1.4%)			(-4.4%, 1.0%)
Surgery - Thoracic Level	0	1 (0.7%)			(-3.3%, 1.5%)
Trauma	27 (18.2%)	26 (18.2%)			(-8.6%, 8.9%)
Urogenital	7 (4.7%)	2 (1.4%)			(-6.0%, 7.8%)
Weakness	4 (2.7%)	2 (1.4%)			(-2.4%, 5.1%)
Wound Issue	1 (0.7%)	4 (2.8%)			(-5.9%, 1.2%)

SEC = at SECURE-C Cervical Artificial Disc; ACDF = Anterior Cervical Discectomy and Fusion (Control)

*Other category defined in Table 2

Table 13 provides a higher level comparison of all pain adverse events that occurred in the study. There were no statistical differences between randomized groups for all pain categories listed. Rates were higher for ACDF than for SECURE-C in all categories, but the differences were not statistically significant.

Table 13. Pain Adverse Events (All Treated Subjects)

Category	NR SEC (N=88)	R SEC (N=148)	R ACDF (N=144)	95% BCI (lower, upper)*
Subjects with ≥1 pain AE	58 (65.9%)	103 (69.6%)	111 (77.1%)	(-17.9%, 2.1%)
Total pain AEs	128	279	207	
Subjects with ≥1 cervical spine related pain AE	47 (53.4%)	89 (60.1%)	102 (69.4%)	(-20.4%, 1.2%)
Total cervical spine related pain AEs	81	173	207	
Pain AEs by location:				
• Neck	• 38 (43.2%)	• 73 (49.3%)	• 82 (56.9%)	• (-19.2%, 3.5%)
• Arm	• 33 (37.5%)	• 64 (43.2%)	• 68 (47.2%)	• (-15.5%, 7.1%)
• Head and arm	• 46 (52.3%)	• 86 (58.1%)	• 96 (66.0%)	• (-3.6%, 3.4%)
• Headache	• 6 (6.8%)	• 14 (9.5%)	• 16 (11.1%)	• (-4.8%, 5.4%)
• Back and/or LE	• 39 (26.5%)	• 63 (31.9%)	• 63 (31.9%)	• (-16.1%, 4.6%)
• Other	• 2 (1.4%)	• 2 (1.4%)	• 3 (4.2%)	• (-3.4%, 3.2%)

NR SEC=Non-randomized SECURE-C; R SEC=Randomized SECURE-C; R ACDF=Control

As specified in the analysis plan, the threshold for establishing success for non-inferiority or superiority is a posteriori probability of 95.4%. Therefore, overall success results demonstrate non-inferiority of the SECURE-C group to the control group. In addition, all components of overall success for the SECURE-C group were non-inferior to the control group. Statistical conclusions at 60 months were not possible due to missing data.

Some adverse events resulted in surgical intervention at the index level, subsequent to the initial surgery. Secondary surgical interventions, classified as revisions, removals, reoperations or supplemental fusions at the index level, are study failures and are reported in **Table 14**, with details provided in **Table 15**. The percentage of patients experiencing secondary surgery at the index level was lower for the SECURE-C group (2.5%) than the ACDF group (9.7%).

Table 14. Secondary Surgical Procedures Interventions at the Index Level (All Patients As Treated)

Adverse Event	Intra-Op (0-2 days)		Peri-Op (>2days-6wks)		Short Term (>6wks-12mo)		Long Term (>12mo-24mo)		Longer Term (>24-60mo)		Longer Term (>60-84mo)		Longer Term (>84mo)		TOTAL Patients (%)	
	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC (N=236)	ACDF (N=144)
Revision	0	0	0	0	0	0	0	0	0	0	0	0	1	0	0	1 (0.7%)
Removal	0	0	0	0	1	2	4	1	4	1	8	2	2	0	1	6 (2.5%)
																(13.9%)
Reoperation	0	0	0	0	0	1	1	0	1	0	1	0	0	0	3	1 (1.3%)
Suppl Fixation	0	0	0	0	0	0	0	0	0	0	1	0	0	0	1	0 (4.4%)
Total	0	0	0	0	1	2	5	2	4	2	8	4	3	0	1	10 (4.2%)
																(15.3%)

SEC = at SECURE-C Cervical Artificial Disc; ACF = ACDF = Anterior Cervical Discectomy and Fusion (Control)

Table 15. Secondary Surgical Procedure Details

Group	Cause/Adverse Event	Action	Postop Days
NR SEC	Neck and shoulder pain	Removal C5-6, fusion same level	183
NR SEC	Arm and paraspinal pain	Removal C5-6, fusion same level	880
NR SEC	Neck and upper extremity pain	Removal C5-6, fusion same level	1945
NR SEC	Neck and upper extremity pain	Removal C6-7, fusion same level	1912
R SEC	Neck pain	Removal C6-7, fusion same level	370
R SEC	Neck pain, C5-6 disc herniation	Removal C4-5, plate and allograft inserted C5-6	507
R SEC	C5-7 radiculopathy	Posterior decompression C5-7	575
R SEC	C5-6 stenosis	Posterior decompression C5-6	942
R SEC	Left C4-5 and left C5-6 stenosis	Posterior decompression C4-5, C5-6	2235
R SEC	Neck and left arm pain	Posterior decompression and supplemental fixation at C6-7	2358
ACDF	Left C5 radiculopathy	Removal C5-6, 2-level plate† spacer inserted C4-6	54
ACDF	Device revision (non-study surgery)	Removal C4-5, 2-level plate† spacer inserted	215
ACDF	Myelopathy	Removal C4-5, plate same level (non-study surgery)	263
ACDF	Neck pain and right arm pain	Removal C6-7, plate and cage inserted C6-7	266
ACDF	Right shoulder pain	Removal C6-7, plate and allograft inserted same level	410
ACDF	Neck pain and thumb paresthesia	Posterior decompression C6-7 (non-study surgery)	408
ACDF	Neck pain and numbness	Removal C5-6, plate and spacer inserted same level	441
ACDF	C4-5, C6-7 disc herniation	Removal C5-6, 3-level plate†spacers inserted C4-7	623
ACDF	Neck and left arm pain	Removal C6-7, plate and cage inserted C5-6	735
ACDF	Neck pain, C5-6 disc herniation	Removal C4-5, plate and allograft inserted C5-6	1058
ACDF	Left arm pain, numbness	Removal C6-7, plate and allograft at C5-6	1162
ACDF	Neck/shoulder pain, C4-5 disc degeneration	Removal C5-6, plate and spacer inserted C4-5	1216
ACDF	Neck pain and right upper extremity pain	Removal C5-6, plate and cage inserted C6-7	1407
ACDF	Adjacent segment disease	C4-6 extension of fusion	1575
ACDF	Neck and arm pain, C5-6 disc degeneration	Removal C6-7, cervical arthroplasty inserted at C5-6	1576
ACDF	Neck and left arm pain	Removal C6-7, 2-level plate† spacers inserted C4-6	1750
ACDF	Left arm pain	Removal C5-6, plate and allograft inserted same level	1804
ACDF	Neck and shoulder pain	Removal of plate and screws C5-6, corpectomy C5-6 and ACDF C4-C7	1924
ACDF	Cervical spondylosis and myelopathy	Removal of plate at C6-7 and adjacent level surgery at C4-5 and C5-6	2156
ACDF	Neck and left arm pain	Removal C4-5 (partial), plate† spacer inserted C4-5 and C6-7	2623
ACDF	Neck and arm pain and numbness	Removal of plate C5-6, allograft spacer inserted at C4-5 and C6-7 and anterior plate from C4-7	3155

NR SEC=Non-randomized SECURE-C; R SEC=Randomized SECURE-C; ACDF=Control

The number and percentage of patients who experienced a device-related adverse event as determined by the Clinical Events Committee (CEC) is provided in **Table 16**. Device-related events were identified as having etiology, temporal association, or causes that is related to the device, such as: revision, removal, reoperation, or supplemental fusion at the index level, surgery at the index level to remove or alter the device, fracture or mechanical failure of device(s), pseudarthrosis, radiolucency around device(s), migration, subsidence, loosening, etc. The CEC felt that it was not appropriate to broadly classify events such as neck or arm pain as potentially device related, as these are commonly reported symptoms for patients entering a study with preoperative neck and/or arm pain.

Based on the CEC's classification, the device-related adverse event profile is lower for the randomized SECURE-C (4.2%) group than the ACDF (15.3%) group because there were fewer secondary surgeries at the index level in the SECURE-C group.

Table 16. Device-Related Adverse Events

Adverse Event	Intra-Op (0-2 days)		Peri-Op (>2days-6wks)		Short Term (>6wks-12mo)		Long Term (>12mo-24mo)		Longer Term (>24-60mo)		Longer Term (>60-84mo)		Longer Term (>84mo)		Patients (%)	
	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC	ACF	SEC (N=236)	ACDF (N=144)
Surgery - Adjacent Level	0	0	1	4	2	4	2	3	3	9	2	2	0	0	10	22 (15.3%)

SEC = at SECURE-C Cervical Artificial Disc; ACDF = Anterior Cervical Discectomy and Fusion (Control)

The change in neurologic status at each study timepoint is provided in **Table 17** for all patient groups. If any one of the four neurologic assessments deteriorated, then the overall neurologic status is considered deteriorated. For overall neurologic status and for each of the four individual assessments, the percentage of patients with stable or improved status was similar for both groups. The randomized SECURE-C group demonstrated numerically greater percentages of patients with stable/improved neurologic status than the control ACDF group at each time point; statistical comparisons of 24 month and 84 month neurologic status success demonstrate non-inferiority with a posterior probability of 100%.

Table 17. Neurological Status

Timepoint	Status	Non-Randomized SECURE-C (N=88)		Randomized SECURE-C (N=148)		Randomized ACDF (N=144)		95% BCI* (lower, upper)	
		Improved	Stable	Improved	Stable	Improved	Stable		
6 months	Improved	51/63 (61.4%)	82/139 (59.0%)	73/130 (56.2%)				(0.7%, 12.6%)	
	Stable	27/63 (32.5%)	57/139 (41.1%)	45/130 (34.6%)					
	Deteriorated	5/63 (6.0%)	4/139 (2.9%)	12/130 (9.2%)					
12 months	Improved	49/61 (60.5%)	80/137 (58.4%)	70/124 (56.5%)				(-1.5%, 11.1%)	
	Stable	26/61 (32.1%)	51/137 (37.2%)	43/124 (34.7%)					
	Deteriorated	6/61 (7.4%)	6/137 (4.4%)	11/124 (8.9%)					
24 months	Improved	47/76 (61.8%)	74/123 (60.2%)	58/101 (57.4%)				(-2.1%, 10.4%)	
	Stable	25/76 (32.9%)	45/123 (36.6%)	36/101 (35.6%)					
	Deteriorated	4/76 (5.3%)	4/123 (3.3%)	7/101 (6.9%)					
36 months	Improved	36/63 (67.9%)	43/63 (68.3%)	29/54 (53.7%)				(-1.4%, 18.9%)	
	Stable	13/63 (24.5%)	18/63 (28.6%)	19/54 (35.2%)					
	Deteriorated	4/63 (7.5%)	2/63 (2.9%)	6/54 (11.1%)					
48 months	Improved	20/29 (68.9%)	25/63 (75.8%)	21/65 (60.0%)				(-11.3%, 16.6%)	
	Stable	8/29 (27.6%)	6/63 (18.2%)	11/65 (31.4%)					
	Deteriorated	1/29 (3.4%)	2/63 (6.1%)	3/65 (6.6%)					
60 months	Improved	16/27 (59.3%)	33/41 (80.5%)	28/38 (73.7%)				(-12.5%, 8.3%)	
	Stable	8/27 (29.6%)	6/41 (44.9%)	9/38 (23.7%)					
	Deteriorated	3/27 (11.1%)	2/41 (2.9%)	1/38 (2.6%)					
72 months	Improved	13/20 (65.0%)	49/69 (71.0%)	33/60 (66.0%)				(-6.5%, 11.7%)	
	Stable	6/20 (30.0%)	17/69 (24.6%)	14/60 (28.0%)					
	Deteriorated	1/20 (5.0%)	3/69 (4.3%)	3/60 (6.0%)					
84 months	Improved	42/62 (67.7%)	87/121 (71.9%)	72/109 (66.1%)				(-0.4%, 15.1%)	
	Stable	16/62 (25.8%)	27/121 (22.3%)	29/109 (21.1%)					
	Deteriorated	4/62 (6.5%)	7/121 (5.8%)	14/109 (12.8%)					

*95% Credible Interval on difference (SECURE-C Randomized - ACDF) between proportions of improvable/stable vs. deteriorated

Effectiveness Results

Primary Effectiveness Analysis

The analysis of effectiveness was based on the as-randomized cohort of 380 total patients (89 non-randomized SECURE-C patients, 151 randomized SECURE-C patients, and 140 ACDF patients). The individual patients' success rates are success divided by the number of patients evaluated at 24, 60 and 84 months. The success rates for each of the individual success components and the overall success is provided in **Table 18** for both randomized groups at 24, 60 and 84 months.

Posterior probabilities of non-inferiority and superiority were calculated using Bayesian statistics. The statistical analysis plan pre-specified that the analysis technique would involve predicting 24 month outcomes for those without them, based on interim 6 month and 12 month observed outcomes, and integrating over the predictions to obtain posterior probabilities of non-inferiority and superiority.

The study was approved using a non-inferiority margin (delta) of 15%. FDA advised that additional analysis be performed with a margin of 10% at the time of PMA. Only the 10% delta analysis is presented. 15% non-inferiority is always met at all variables demonstrating non-inferiority at 10%. According to the statistical analysis plan, if non-inferiority was determined, then superiority would be evaluated; these results are also provided.

In addition to the protocol-defined overall success criteria, FDA requested an additional alternative definition of overall success to include improvement in NDI of 15 points, maintenance or improvement in neurologic status, absence of device-related adverse events, exclusion of fusion criteria, and no intraoperative changes in SECURE-C treatment (i.e., intraoperative conversion to fusion). Analysis using the alternative FDA-defined definition is provided in **Table 19**.

Table 18. Overall Success Rates at 24, 60 and 84 Months Using Protocol-Specified Definition

Component	Non-Randomized SECURE-C (N=88)	SECURE-C (N=151)	ACDF (N=140)	Posterior Probability		95% BCI* (lower, upper)
				Non-Inferiority	Superiority	
24 months						
NDI (≥25% impr)	67/78 (85.9%)	127/139 (91.4%)	101/116 (87.1%)	100.0%	87.8%	(-3.2%, 12.6%)
No removals etc.	84/86 (97.7%)	144/147 (98.0%)	127/139 (92.5%)	100.0%	98.2%	(0.3%, 10.8%)
N complications	85/85 (100%)	145/145 (100.0%)	132/132 (100%)	100.0%	92.9%	(-2.0%, 2.3%)
Fusion (control)	N/A	N/A	90/101 (89.1%)	N/A	N/A	N/A
Overall Success	67/79 (84.8%)	127/141 (90.1%)	81/114 (71.1%)	100.0%	100.0%	(8.2%, 27.1%)
60 months						
NDI (≥25% impr)	27/30 (90.0%)	41/42 (97.6%)	32/37 (86.5%)	N/A	N/A	(-3.0%, 19.1%)
No removals etc.	66/68 (97.1%)	128/133 (96.2%)	107/124 (86.3%)	N/A	N/A	(2.3%, 21.1%)
No complications	66/66 (100%)	127/127 (100%)	112/112 (100%)	N/A	N/A	(-4.3%, 4.7%)
Fusion (control)	N/A	N/A	35/36 (97.2%)	N/A	N/A	N/A
Overall Success	27/32 (84.4%)	40/45 (87.0%)	28/51 (54.9%)	N/A	N/A	(15.6%, 42.7%)
84 months						
NDI (≥25% impr)	53/63 (84.1%)	113/125 (90.4%)	92/107 (86.0%)	100.0%	86.8%	(-3.6%, 13.3%)
No removals etc.	61/65 (93.8%)	124/131 (94.7%)	102/121 (84.3%)	100.0%	99.6%	(2.4%, 16.9%)
No complications	63/63 (100.0%)	125/125 (100%)	107/105 (100%)	100.0%	93.9%	(-2.2%, 2.7%)
Fusion (control)	N/A	N/A	101/104 (97.1%)	N/A	N/A	N/A
Overall Success	53/65 (81.5%)	113/131 (86.3%)	84/120 (70.0%)	100.0%	99.9%	(6.3%, 25.6%)