

Tactra™ Malleable Penile Prosthesis



Offered in three size ranges:

9.5 mm diameter width x 14 cm – 23 cm length
11 mm diameter width x 16 cm – 25 cm length
13 mm diameter width x 18 cm – 27 cm length

General Information

Name of Product: Tactra Malleable Penile Prosthesis

Product Description: Malleable Penile Prosthesis

Manufacturer: Boston Scientific

Manufacturer Federal Tax ID: 04-269-5240

Supplement/Replace current in-house products: Yes

Clinical Improvements

Tactra Penile Prosthesis is the next-generation malleable prosthesis with enhanced ease of implant. It is designed for durability, offering both excellent rigidity and dependable concealment in a device that is natural to the touch.

Regulatory

Is this product FDA cleared for this intended use? FDA cleared

What Class of device under the FDA is this considered? IIB

Does the product/device have an FDA investigational device exemption (IDE)? No

Utilization

Is this item/technology on contract with GPOs and/or IDNs?
Please speak to your Boston Scientific Sales Representative for the contract status of specific GPOs and IDNs

Ship unit: Each

Mode of transportation: FedEx™ Delivery

Lead time in working days? 2 days

What are the dimensions of the package? 1.19 x 12.39 x 5.19” – Cardboard Box

Method of purchase: Direct purchase or bill upon use

Does this item require special storage considerations?
Per the DFU, store in a clean, dry, dark area at room temperature.

Is this a dated product? Product contains expiration date on package label.

What specific departments/clinical areas will use the product/procedure?
Urology Operating Room (OR)

What department(s) will use and/or be affected by this product?
Urology OR, and Purchasing

Is there a requirement for staff training? No

Will there be additional implementation costs, such as installation, cost of education, impact on equipment or additional space? No

Does the product/procedure require a company representative to be present to operate equipment or to provide assistance to the physicians?
Per institution’s penile implant procedure protocol

Is there any other equipment involved with the use of this product that will need to be leased, purchased, consigned or rented? No

What is the average length of procedure time to use this product/perform this procedure (surgery minutes)? 60 minutes

Will this equipment interface with any other equipment/supplies currently utilized at this facility? Yes, penile implant procedure instruments



Material/Environment

Does this product contain metal substances that may affect tests and or procedures performed on patients? No

If yes, is this product MRI safe? MRI conditional, refer to the DFU for further description.

Is this considered an implantable device? Yes

Does this item and its packaging contain latex? No

Is this a pharmaceutical or contain any pharmaceutical product? No

Does the product require a Material Safety Data Sheet? No

Is this product reusable? No

What additional waste or recycle costs are anticipated? After use, dispose of product and packaging in accordance with hospital, administrative, and/or local government policy.

Does the product contain: Mercury? No
PVC? No

Halogenated flame retardants/halogenated organic chemicals (HOCs)? No

UPN	720080-01	720081-01	720082-01
GTIN Number	08714729979340	08714729979357	08714729979364
Catalog Number	720080-01	720081-01	720082-01
Product Name	Tactra Malleable Penile Prosthesis	Tactra Malleable Penile Prosthesis	Tactra Malleable Penile Prosthesis
Product Description	Malleable Penile Prosthesis	Malleable Penile Prosthesis	Malleable Penile Prosthesis
Size (cm)	9.5mm x 14cm – 23cm	11mm x 16cm – 25cm	13mm x 18cm – 27cm

Reimbursement

Is this product reimbursable by insurance? The procedures for which it is used are reimbursable when deemed medically necessary. For additional coding and reimbursement information, contact your local Territory Manager or the Urology and Pelvic Health Reimbursement Help Desk at 866-367-2796.

What is the Medicare Pass-Through Code (aka C-code or HCPCS)? C-Codes are tracking codes established by the Centers for Medicare & Medicaid Services (CMS) to assist Medicare in establishing future APC payment rates. C-Codes only apply to Medicare hospital outpatient claims. They do not trigger additional payment to the facility. The C-code for this product is C2622.

Is this a patient-chargeable product? “Patient chargeable” is a colloquial term used to convey that a device/supply is appropriately charged to the patient’s account (i.e., as a distinct line item on the patient’s claim) in the hospital/facility’s patient accounting or AR system. It does not mean that the patient is actually charged directly for the device/supply nor would an insured patient ever pay an additional amount “out of pocket” for the device/supply. The fact that a hospital/facility chooses to designate certain devices/supplies (e.g., single-use devices) as “patient chargeable” will not in and of itself result in immediate increased reimbursement for the hospital/facility. It will allow CMS to better factor the true cost of the procedure into future Medicare reimbursement rate setting. It may also help in negotiations with private payers by more clearly demonstrating novel device costs that have been introduced to a procedure.

The designation of a given device/supply as “patient chargeable” is entirely up to the discretion and policy of the individual hospital/facility. Section 2202.8 of the Medicare Provider Reimbursement Manual dealing with Ancillary Services (e.g., operating room) does not specifically address which items are part of the basic (routine) charge and which are charged in addition to the basic charge (non-routine). Medicare is on record that it is up to the individual hospital to determine whether to and how to itemize the charge for a specific device/supply or alternatively incorporate it into overhead (e.g., via the OR charge).

However, Medicare does require that charges billed on the CMS-1450 form (aka UB-04) be aggregated under the appropriate Revenue Code.

The appropriate Revenue Code is 272 – Medical/Surgical Supplies and Devices-Sterile Supply.

Relevant Reimbursement Codes: Payer policies will vary and should be verified prior to treatment for limitations on diagnosis, coding or site of service requirements. The coding options listed within this guide are commonly used codes and are not intended to be an all-inclusive list. We recommend consulting your relevant manuals for appropriate coding options.

The coding information listed within this guide have been included solely for informational purposes only. It is not intended to imply, promote, or guarantee coverage and/or payment from a payer. The Health Care Provider (HCP) is solely responsible for selecting and accurately reporting coding based on the patient’s medical condition and the treatment needs of that patient, and the independent medical judgment of the HCP.

Procedure Name	Malleable Penile Implant	Malleable Penile Implant Removal and Replacement
CPT Code	54400	54416
CPT Code Description	Insertion penile prosthesis; non-inflatable (semi-rigid)	Removal and replacement of non-inflatable (semi-rigid) or inflatable (self-contained) penile prosthesis at the same operative session
ICD-10 Diagnosis Code	F52.21Male erectile disorder N52.01Erectile dysfunction due to arterial insufficiency N52.02Corporo-venous occlusive erectile dysfunction N52.03Combined arterial insufficiency and corporo-venous occlusive erectile dysfunction N52.2Drug-induced erectile dysfunction N52.31Erectile dysfunction following radical prostatectomy N52.32Erectile dysfunction following radical cystectomy N52.33Erectile dysfunction following urethral surgery N52.34Erectile dysfunction following simple prostatectomy N52.35Erectile dysfunction following radiation therapy N52.36Erectile dysfunction following interstitial seed therapy N52.37Erectile dysfunction following prostate ablative therapy N52.39Other and unspecified postprocedural erectile dysfunction N52.8Other male erectile dysfunction N52.9Male erectile dysfunction, unspecified T83.410ABreakdown (mechanical) of implanted penile prosthesis, initial encounter T83.420ADisplacement of implanted penile prosthesis, initial encounter T83.490AOther mechanical complication of implanted penile prosthesis, initial encounter T83.61Infection and inflammatory reaction due to implanted penile prosthesis T83.61XAInfection and inflammatory reaction due to implanted penile prosthesis, initial encounter T83.69Infection and inflammatory reaction due to other prosthetic device, implant and graft in genital tract T83.69XAInfection and inflammatory reaction due to other prosthetic device, implant and graft in genital tract, initial encounter T83.728AExposure of other implanted mesh into organ or tissue, initial encounter T83.82XAFibrosis due to genitourinary prosthetic devices, implants and grafts, initial encounter T83.83XAHemorrhage due to genitourinary prosthetic devices, implants and grafts, initial encounter T83.84XAPain due to genitourinary prosthetic devices, implants and grafts, initial encounter T83.85XAStenosis due to genitourinary prosthetic devices, implants and grafts, initial encounter T83.86XAThrombosis due to genitourinary prosthetic devices, implants and grafts, initial encounter T83.89XAOther specified complication of genitourinary prosthetic devices, implants and grafts, initial encounter T83.9XXAUnspecified complication of genitourinary prosthetic device, implant and graft, initial encounter	
ICD-10 Procedure Code	0VUS0JZSupplement Penis with Synthetic Substitute, Open Approach 0VPS0JZRemoval of Synthetic Substitute from Penis, Open Approach 0VUS0JZSupplement Penis with Synthetic Substitute, Open Approach	
Possible MS-DRG Assignment	709Penis procedures with CC/MCC 710Penis procedures without CC/MCC	

Caution: U.S. Federal law restricts this device to sale by or on the order of a physician.

CAUTION: The law restricts these devices to sale by or on the order of a physician. Indications, contraindications, warnings and instructions for use can be found in the product labelling supplied with each device. Information for use only in countries with applicable health authority registrations. This material not intended for use in France.

Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, and potential adverse events.

The Tactra[®] Malleable Penile Prosthesis is intended for use in the treatment of erectile dysfunction (impotence) in adult males. Implanting a penile prosthesis will damage or destroy any remaining natural ability to have a spontaneous erection, as well as make other treatment options impossible.

Men with diabetes, spinal cord injuries, or skin infections may have an increased risk of infection. Implantation may result in penile shortening, curvature or scarring.

Potential adverse events may include device malfunction/failure leading to additional surgery, device/tissue erosion, infection, and pain/soreness. MH-611819-AA

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Products shown for INFORMATION purposes only and may not be approved or for sale in certain countries. Please check availability with your local sales representative or customer service.

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