

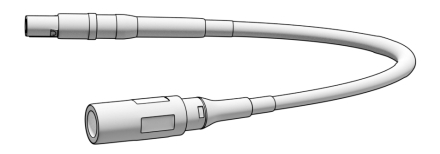
# **iCONNECT**

# **NAVISTAR ADAPTER CABLE**

INSTRUCTIONS FOR USE

REF

001-009081-1



 $\mathbf{R}_{\scriptscriptstyle\mathsf{ONLY}}$ 

Caution: Federal Law (USA) restricts this device to sale by or on the order of a physician.

Made in USA

DSP-0304; Rev. F

Effective date: 10 November 2023

	SYMBOL	LEGEND	
EC REP	Authorized Representative in the European Community	REF	Catalogue Number
$\triangle$	Caution		Consult Operating Instructions
	Date of Manufacture		Do not use if package is damaged and consult instruction for use
类	Keep Away from Sunlight		Keep Dry
	Importer	LOT	Lot Number
	Manufacturer	CE	Marking for devices entering the European market
MD	Medical Device	NON STERILE	Non-Sterile
	Packaging Unit	Ronly	Prescription Only
SN	Serial Number	1	Temperature Limit
<b>*</b> *	Usable Length		

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# **ENGLISH**

CAUTION: Read the instructions carefully before using the product.

# **DEVICE DESCRIPTION**

The Navistar® Adapter System Cable is a component of the iCONNECT™ System. The adapter cable offers the ability to use the *NAVISTAR* RMT Catheter or *NAVISTAR* THERMOCOOL® RMT Catheter in conjunction with the *iCONNECT* System.

# ADDITIONAL REQUIRED COMPONENTS

- iCONNECT Electronics Hub
- Biosense Webster Interface Cable C5MHNAVMHS
- NAVISTAR RMT Catheter or
- NAVISTAR THERMOCOOL RMT CATHETER

# **INDICATIONS**

The Navistar Adapter System Cable is designed to connect a NAVISTAR RMT Catheter or NAVISTAR THERMOCOOL RMT CATHETER to the appropriate equipment for use with the iCONNECT System.

# CONTRAINDICATIONS

The Navistar Adapter System Cable has no known contraindications.

#### **HOW SUPPLIED**

This cable is provided non-sterile. Inspect the packaging for damage prior to use. Examine the cable prior to use or reuse for any damage or defects (i.e. bent pins, kinks, nicks, crushed or elongated sections). Do not use if any damage or defects are identified.

#### HANDLING AND STORAGE

The product may only be used in a medical treatment facility that has been set up for the appropriate application and by trained staff. Store the product in its original packaging in a cool, dry place away from light at between  $10^{\circ}$ C and  $25^{\circ}$ C. When stored appropriately, the product can be used up to the use before date indicated on the packaging.

#### **CLEANING**

The product may be contaminated after use and should be cleaned with a cloth or swab dampened in pH neutral, EPA-approved hospital grade solutions.

#### **DISPOSAL**

If damaged, dispose of the product and its residual elements or waste items in accordance with hospital regulations or local government policy.

# WARNINGS

- The Navistar Adapter System Cable should only be used by trained physicians.
- Dispose of product packaging in accordance with hospital, administrative and/or local government policy.
- If product damage or defects are identified, dispose of the product in accordance with hospital, administrative and/or local government policy.
- No modification of this device is allowed.
- Patient or operator injury can result from improper handling of the cable.
- Failure to abide by the above warnings might result in damage to the product, or result in serious adverse events.

# **CAUTIONS**

- Do not immerse cable connector(s) in fluids.
- If the cable is used in the presence of electrical equipment, noise may be induced into the cable.
- Do not attempt to repair any damage. In case of doubt, discard the cable and do not use or reuse.

# **INSTRUCTIONS FOR USE**

Refer to the *iCONNECT* User Guide for cable connection instructions and operation of the *iCONNECT* System.

#### TRADEMARKS

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# STEREOTAXIS TECHNICAL SUPPORT

For technical support, please contact Stereotaxis TeleRobotic Support Team (TST) at 1-866-269-5268 or 1-314-678-6200 or email tst@stereotaxis.com.

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### NOTICE TO THE USER AND/OR PATIENT

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

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