

## TD2 Adhesive Clip

H100 – 34015-002

H200 – 34015-001

H300 – 34015-003

### DESCRIPTION

The single-use adhesive clip is designed to hold the TD2 Therapy Disc in place over the Implantable Pulse Generator location. The adhesive clip is not intended to be reused.

### INDICATIONS FOR USE

#### Spinal Cord Stimulation (SCS)

This system is indicated as the sole mitigating agent, or as an adjunct to other modes of therapy used in a multidisciplinary approach for chronic, intractable pain of the trunk and/or limbs, including unilateral or bilateral pain.

The trial devices are solely used for trial stimulation (no longer than 30 days) to determine efficacy before recommendation for a permanent (long term) device.

#### Peripheral Nerve Stimulation (PNS)

This system is indicated for pain management in adults who have severe intractable chronic pain of peripheral nerve origin, as the sole mitigating agent, or as an adjunct to other modes of therapy used in a multidisciplinary approach. The Nalu Neurostimulation System for PNS is not intended to treat pain in the craniofacial region.

The trial devices are solely used for trial stimulation (no longer than 30 days) to determine efficacy before recommendation for a permanent (long term) device.

### CONTRAINDICATIONS AND WARNINGS

For a complete list of contraindications and warnings associated with this product, see the Nalu Neurostimulation System User instructions for use.

Carefully examine all product packaging before use; do not use product if packaging is damaged or open.

### HANDLING (H100, H200, H300)

Single use only; DO NOT REUSE.

Do not tamper with Adhesive Clip; it is designed to work as intended without modification.

### INSTRUCTIONS FOR USE

For complete instructions, please refer to the Nalu Neurostimulation System User instructions for use.

#### Preparing Your Skin

1. Make sure your skin is clean and dry.
2. If necessary, trim hair with scissors or small beard trimmer. Do not shave since this can lead to irritation.
3. If desired, commercially available skin barrier wipes and sprays may be used to prepare the skin.

#### Applying the Clip

1. Remove the adhesive liner
2. Grasp the clip between your thumb and finger and place your index finger through the center hole.
3. Use your inserted index finger and locate the implant site.
4. Once the implant site has been located, press the clip against your skin, ensuring the heel (where the therapy disc sits) is towards the bottom.

#### Placing the Therapy Disc into the Adhesive Clip

1. Slide the Therapy Disc into the open side of the clip until it clicks into place. Ensure that the nalu logo is facing away from your skin.

### CUSTOMER SERVICE – USA

In USA, to order replacement parts, or product inquiries, please contact Nalu's Customer Service.



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

**R<sub>x</sub> Only**

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Refer to the following table for an explanation of applicable product label symbols:

Symbol	Title	Explanation	Standard	Reference
	Manufacturer	Medical device manufacturer, as defined in EU Directive 93/42/EEC	ISO 15223-1	5.1.1
	Date of manufacture	Date when the medical device was manufactured.	ISO 15223-1	5.1.3
	Use-by date	Date after which the medical device is not to be used.	ISO 15223-1	5.1.4
	Batch code	Manufacturer's batch code so that the batch or lot can be identified.	ISO 15223-1	5.1.5
	Catalogue number	Manufacturer's catalogue number so that the medical device can be identified.	ISO 15223-1	5.1.6
	Do not use if package is damaged	Medical device should not be used if the package has been damaged or opened.	ISO 15223-1	5.2.8
	Best Before	This device is best used before this date	N/A	N/A
	Keep away from sunlight	A medical device that needs protection from light sources.	ISO 15223-1	5.3.2
	Keep dry	A medical device that needs to be protected from moisture.	ISO 15223-1	5.3.4
	Temperature limit	The temperature limits to which the medical device can be safely exposed.	ISO 15223-1	5.3.7
	Do not re-use	A medical device that is intended for one use, or for use on a single patient during a single procedure.	ISO 15223-1	5.4.2
	Consult instructions for use	Indicates the need for the user to consult the instructions for use.	ISO 15223-1	5.4.3
	Refer to instruction manual/booklet.	Indicates the instruction manual/booklet must be read. (This symbol is blue and white on the device label)	ISO 7010:2011	M002
	Caution	User to consult the instructions for use for important cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device itself.	ISO 15223-1	5.4.4
	MR Conditional	Medical device demonstrated safety in the MR environment within defined conditions.	ASTM F2503-13	Fig. 6
	MR Unsafe	Medical device poses unacceptable risks to the patient, medical staff or other persons within the MR environment.	ASTM F2503-13	Fig. 9
	Prescription use only	Caution: Law prohibits dispensing without prescription	21 CFR 801.109	N/A
	Quantity	Indicates the total number of products provided in a package.	N/A	N/A

Symbol	Title	Explanation	Standard	Reference
	Serial Number	Manufacturer's serial number so that a specific medical device can be identified.	ISO 15223-1	5.1.7
	Radio Transmitter	Device contains a radio-frequency (RF) transmitter, which may cause RF interference with other devices near this device	IEC 60601-1-2:2007	5.1.1
	Authorized Representative in the European Community	Authorized representative in the European Community	ISO 15223-1	5.1.2
	Protected against electric shock	Device contains a type BF applied part to protect you from shock. The device is internally powered and is intended for continuous operation	IEC 60417	5333
	Protected against access to certain hazardous parts	Protected against solid objects over 12.5mm (e.g., a finger) and protected against falling drops of water, if the case is disposed at any angle up to 15 degrees from vertical	IEC 60529	N/A
	Protected from dust and temporary immersion in water	Protected from dust and against effects of immersion in water up to 1m depth for 30 mins	IEC 60529	N/A
	FCC	This symbol indicates that this equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules.	47 CFR 2.926	N/A
	Do not throw in the trash	This product shall not be treated as household waste. Instead it is the user's responsibility to return this product to Nalu Medical for reprocessing. By ensuring that this product is disposed of properly, you will help prevent potential negative consequences for the environment and human health, which could be caused by inappropriate waste handling of this product. The recycling of materials will help to conserve natural resources. For more information about how to return this product for recycling, please contact Nalu Medical.	BS EN 50419 Marking of Electrical and Electronic Equipment in Accordance with Article 11(2) of Directive 2002/96/EC (WEEE)	Fig. 1
	Sterilized using ethylene oxide	Indicates a medical device that has been sterilized using ethylene oxide.	ISO 15223-1	5.2.3
	Do Not Resterilize	Indicates a medical device that is not to be resterilized.	ISO 15223-1	5.2.6
	Stimulator length	Indicated the length of the device	N/A	N/A
	Humidity limitation	Indicates the range of humidity to which the device can be safely exposed	ISO 15223-1	5.3.8
	Atmospheric pressure limitation	Indicates the range of atmospheric pressure to which the device can be safely exposed	ISO 15223-1	5.3.9