Natus Base Unit to Nicolet™ Cortical Stimulator Cable

Instructions for Use:

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EU Representative / Importer
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Gort, Co. Galway, Ireland

Associated Product Part Number:
016728

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CaviWipes is a trademark of Metrex Research, LLC. Sani-Cloth is a registered trademark of PDI, Inc.
Description:
The Natus Base Unit to Nicolet Cortical Stimulator Cable consists of a 197 inch (5 m) long serial cable connecting the Natus Base Unit to the Nicolet™ Cortical Stimulator serial interface connector.

Intended Use:
The Natus Base Unit to Nicolet Cortical Stimulator Cable is intended to be used as an accessory with Natus Quantum and Natus Base to connect the Nicolet Cortical Stimulator control port to the Natus Base serial port for the purpose of controlling the stimulator from NeuroWorks® software during functional mapping procedures during LTM studies.

Intended Users and Patient Target Group:
The Natus Base Unit to Nicolet Cortical Stimulator Cable is intended to be used by trained medical professionals and is designed for use in clinical environments such as hospital rooms, epilepsy monitoring units, intensive care units, and operating rooms. It can be used with patients of all ages but is not designed for fetal use.

Clinical Benefits:
The Natus Base Unit to Nicolet Cortical Stimulator Cable is a mandatory item for NeuroWorks to control the Nicolet Cortical Stimulator for the purpose of cortical mapping procedures in the Epilepsy Long Term Unit. Cortical mapping provides valuable information about functional areas in the brain in the context of brain surgery.

Contraindications and Side Effects:
There are no known contraindications or side effects for using the Natus Base Unit to Nicolet Cortical Stimulator Cable.

Operating Instructions:
The Natus Base Unit to Nicolet Cortical Stimulator Cable is connected on one end to Natus Base Unit, and on the other end to the Nicolet Cortical Stimulator. Nicolet Cortical Stimulator is not included in Natus Quantum and must be added as an option.
Cleaning Instructions:

1. Clean with a commercial wipe such as CaviWipes™ or Sani-Cloth® to remove visible soil.
2. Wipe the article using a lint-free cloth and air dry.
3. The cleaning procedure must be in accordance with your local facility's guidelines. The user/operator shall clean the device after every use.

Understanding Warning and Caution Statements:

<table>
<thead>
<tr>
<th>WARNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Refers to a hazardous situation that could result in death or serious injury if not avoided.</td>
</tr>
<tr>
<td>• Information on how the hazardous situation is avoided.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CAUTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Refers to a hazardous situation that could result in minor or moderate injury or material damage if not avoided.</td>
</tr>
<tr>
<td>• Information on how the hazardous situation is avoided.</td>
</tr>
</tbody>
</table>

Warnings and Precautions:

<table>
<thead>
<tr>
<th>WARNING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Misaligned connector pins may lead to damage of system components at power-on.</td>
</tr>
<tr>
<td>• Refer to the Natus Quantum User and Service Manual for connection diagrams.</td>
</tr>
</tbody>
</table>
**CAUTION**

Device dropped or damaged in transit or use may lead to loss of function.
- Inspect the device before each use and do not use if damaged.

Unauthorized modification or servicing could lead to loss of device safety, function or performance.
- Do not perform any unauthorized modifications.

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**Environmental Specifications:**

**Operating Conditions:**
- Temperature: +10°C to +30°C (+50°F to +86°F)
- Relative Humidity: 30% to 75%
- Atmospheric Pressure: 700 hPa to 1060 hPa

**Storage Conditions:**
- Temperature: -25°C to +60°C (-13°F to +140°F)
- Relative Humidity: 10% to 95%
- Atmospheric Pressure: 500 hPa to 1060 hPa

**Compliance Standards:**
- ISO 10993-1: 2018 Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process
- ETSI EN300 019-2-1 Environmental Engineering (EE); Environmental conditions and environmental tests for telecommunications equipment; Part 2-1: Specification of environmental tests; Storage
- ETSI EN300 019-2-2 Environmental Engineering (EE); Environmental conditions and environmental tests for telecommunications equipment; Part 2-2: Specification of environmental tests; Transportation
- ASTM D4169-16 Standard Practice for Performance Testing of Shipping Containers and Systems for Vibration

**Disposal Instructions:**

Natus is committed to meeting the requirements of the European Union WEEE (Waste Electrical and Electronic Equipment) Regulations 2014. These regulations state that electrical and electronic waste must be separately collected for the proper treatment and recovery to ensure that WEEE is reused or recycled safely. In line with that commitment Natus may pass along the obligation for take back and
recycling to the end user, unless other arrangements have been made. Please contact us for details on the collection and recovery systems available to you in your region at natus.com.

Electrical and electronic equipment (EEE) contains materials, components and substances that may be hazardous and present a risk to human health and the environment when WEEE is not handled correctly. Therefore, end users also have a role to play in ensuring that WEEE is reused and recycled safely. Users of electrical and electronic equipment must not discard WEEE together with other wastes. Users must use the municipal collection schemes or the producer/importers take-back obligation or licensed waste carriers to reduce adverse environmental impacts in connection with disposal of waste electrical and electronic equipment and to increase opportunities for reuse, recycling and recovery of waste electrical and electronic equipment.

Equipment marked with the below crossed-out wheeled bin is electrical and electronic equipment. The crossed-out wheeled bin symbol indicates that waste electrical and electronic equipment should not be discarded together with unseparated waste but must be collected separately.

Disclaimer:

Any serious incident that has occurred in relation to the device should be reported to Natus Medical Incorporated DBA Excel-Tech Ltd. (Xltek) and the competent authority of the Member State in which the user and/or patient is established.

Refer to the Natus website for an electronic copy of this document.

Instructions to Access the eIFU:

A copy of the Instructions for Use in PDF format is in the associated product area:

- Neurology: https://neuro.natus.com/neuro-support

Search for “Natus Base Unit to Nicolet Cortical Stimulator Cable IFU” (refer to the Product Part Number) and choose the version for your local language for the instructions to use.

The files can be printed, saved, or searched using Adobe Reader. A copy of Adobe Reader can be downloaded directly from Adobe Systems (www.adobe.com).
# Glossary of Symbols:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Standard Reference</th>
<th>Standard Title</th>
<th>Symbol Title</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medical Device</td>
<td>Not applicable</td>
<td>Not applicable</td>
<td>An indication of Medical Device</td>
<td>This product is a medical device.</td>
</tr>
<tr>
<td>Rx only</td>
<td>21 CFR Part 801.109(b)(1)</td>
<td>Labeling-Prescription devices</td>
<td>Prescription only</td>
<td>Indicates that the product is authorized for sale by or on the order of a licensed healthcare practitioner.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.4.5 (Reference Annex B for the general prohibition symbol)</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Not made with natural rubber latex</td>
<td>Indicates that the medical device is not made with natural rubber latex.</td>
</tr>
<tr>
<td></td>
<td>2012/19/EU</td>
<td>Waste Electrical and Electronic Equipment (WEEE)</td>
<td>Disposal at end of operating life instructions</td>
<td>Indicates that electrical and electronic equipment waste should not be discarded together with unseparated waste but must be collected separately.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.1.1</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Manufacturer</td>
<td>Indicates the medical device manufacturer.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.1.2</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Authorized representative in the European Community</td>
<td>Indicates the Authorized representative in the European Community.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.1.3</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Date of manufacture</td>
<td>Indicates the date when the medical device was manufactured.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.1.5</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Batch code</td>
<td>Indicates the manufacturer's batch code so that the batch or lot can be identified.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.1.6</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Catalogue number</td>
<td>Indicates the manufacturer's catalogue number so that the medical device can be identified.</td>
</tr>
<tr>
<td></td>
<td>ISO 15223-1 Symbol 5.4.3 Annex A #A.15</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied</td>
<td>Consult instructions for use</td>
<td>Indicates an instruction to consult an electronic instructions for use (eIFU).</td>
</tr>
<tr>
<td>Symbol</td>
<td>Standard Reference</td>
<td>Standard Title</td>
<td>Symbol Title</td>
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</table>
| ![IEC 60601-1 Table D.2 #10](image) | IEC 60601-1 Table D.2 #10 | Medical electrical equipment — Part 1: General requirements for basic safety and essential performance | Follow instructions for use | Refer to instruction manual/booklet.  
NOTE on ME EQUIPMENT “Follow instructions for use” |
| ![ISO 15223-1 Symbol 5.4.4](image) | ISO 15223-1 Symbol 5.4.4 | Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied | Caution | Indicates the need for the 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. |
| ![IEC 60601-1 Table D.1 #10](image) | IEC 60601-1 Table D.1 #10 | Medical electrical equipment — Part 1: General requirements for basic safety and Essential performance | General warning sign | Indicates a hazard of potential personal injury to patient or operator. |
| ![MDR 2017/745](image) | IEC 60601-1 Table D.2 #2 | Medical electrical equipment — Part 1: General requirements for basic safety and essential performance | General warning sign | Indicates a hazard of potential personal injury to patient or operator. |
| ![ISO 15223-1 Symbol 5.3.7](image) | ISO 15223-1 Symbol 5.3.7 | Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied | Temperature limit | Indicates the (storage) temperature limits to which the medical device can be safely exposed. |
| ![ISO 15223-1 Symbol 5.3.8](image) | ISO 15223-1 Symbol 5.3.8 | Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied | Humidity limitation | Indicates the range of (storage) humidity to which the medical device can be safely exposed. |
| ![ISO 15223-1 Symbol 5.3.9](image) | ISO 15223-1 Symbol 5.3.9 | Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied | Atmospheric pressure limitation | Indicates the acceptable upper and lower limits of atmospheric pressure for transport and storage.