Nicolet Tendon (Reflex) Hammer

Instructions for Use:

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EC REP EU Representative / Importer
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IDA Business Park
Gort, Co. Galway, Ireland

Associated product part number:

842-116700

Description:

The Nicolet Tendon (Reflex) Hammer can be used as an accessory to Electrodiagnostic testing systems during mechanical stimulation procedures. The Tendon Hammer, P/N 842-116700, is a device used to mechanically present stimuli to patients while simultaneously triggering data acquisition on the Electrodiagnostic system.

Intended Use:

The Tendon Hammer is used in conjunction with nerve conduction studies to manually stimulate the patient while simultaneously triggering the initiation of the data acquisition. It is used to assist in the qualification of central and peripheral physiological reflexes of the body. The product is intended to be used by trained medical healthcare professionals.

Intended Users and Patient Target Group:

The Tendon Hammer is intended for use by skilled physicians and technologists trained in the specialty of Electrodiagnostic testing.

The target patient population is the pediatric and adult patient population requiring NCS/EMG/EP procedures.
Clinical Benefits:
The Tendon Hammer is a useful accessory during reflex studies and mechanical stimulation procedures. Usage of the Tendon Hammer during such diagnostic procedures is at the discretion of the clinical provider.

Contraindications and Side Effects:
There are no known contraindications or side effects for procedures performed with the Tendon Hammer.

Operating Instructions:
• Connect the 1/8” mono mini-phone plug of the Tendon Hammer to the matching 1/8” mono mini-phone receptacle of the Electrodiagnostic system.

Cleaning Instructions:
• Clean with a commercial wipe such as CaviWipes™ or Sani-Cloth® to remove visible soil.
• Wipe the article using a lint-free cloth and air dry.
• The cleaning procedure must be in accordance with your local facility’s guidelines. The user/operator shall clean the device after every use.
• Tendon Hammer cannot be sterilized.

Understanding Warnings and Cautions Statements:

WARNING
Refers to a hazardous situation that could result in death or serious injury if not avoided.
• Information on how the hazardous situation is avoided.

CAUTION
Refers to a hazardous situation that could result in minor or moderate injury or material damage if not avoided.
• Information on how the hazardous situation is avoided.

Warnings and Precautions:

WARNING
System components immersed or in contact with liquids may cause electrical shock.
• Do not immerse, drip, or spray liquids onto the device.
CAUTION

Device dropped or damaged in transit/ use could lead to loss of function or delayed diagnosis.
• Inspect the device prior to each use and do not use if damaged.

Device when used by untrained user could lead to patient injury, incorrect diagnosis or delayed diagnosis.
• This device is intended to be used by qualified healthcare professionals.

Environmental Specifications:

Operating Conditions:
• Temperature: +15.6°C (+60°F) to +32.2°C (+90°F)
• Relative Humidity: 20% to 80% (non-condensing)
• Altitude: 0 to 10,000 ft (0 to 3 km)

Storage Conditions:
• Temperature: -17.7°C (0°F) to +55°C (+132°F)
• Relative Humidity: 10% to 90% (non-condensing)
• Altitude: 0 to 35,000 ft (0 to 10.668 km)

Compliance Standards:
• ISO 10993-1: 2018 Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process
• ETS 300 019-2-1 Environmental Engineering (EE); Environmental conditions and environmental tests for telecommunications equipment; Part 2-1: Specification of environmental tests; Storage
• ETS 300 019-2-2 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
• IEC 60601-1:2005+A1:2012+Cor1:2014 - General Safety Ed. 3.1
• IEC 60601-1-2:2014 – EMC Fourth Edition
• IEC 60601-2-40:2016 – Particular requirements for the basic safety and essential performance of electromyography and evoked response equipment
• IEC 60601-1-6:2013 – Collateral Usability

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:
Natus Medical Incorporated is not responsible for injury, infection or other damage resulting from the use of this product.

Any serious incident that has occurred in relation to the device should be reported to Natus Medical Incorporated 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.
## Glossary of Symbols:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Standards Reference</th>
<th>Standard Title of Symbol</th>
<th>Symbol Title as per Referenced Standard</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medical Device</td>
<td>-</td>
<td>-</td>
<td>An indication of Medical device</td>
<td>This product is a medical device.</td>
</tr>
<tr>
<td><strong>Rx only</strong></td>
<td>21 CFR Part 801.109(b)(1)</td>
<td>Labeling-Prescription devices.</td>
<td>Prescription only</td>
<td>Indicates the product is authorized for sale by or on the order of a licensed healthcare practitioner.</td>
</tr>
<tr>
<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>
<td></td>
</tr>
<tr>
<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>
<td></td>
</tr>
<tr>
<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>
<td></td>
</tr>
<tr>
<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 or Lot code</td>
<td>Indicates the manufacturer's batch code so that the batch or lot can be identified.</td>
<td></td>
</tr>
<tr>
<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>
<td></td>
</tr>
<tr>
<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>
<td></td>
</tr>
<tr>
<td>ISO 60601-1 Table D.2 #10</td>
<td>Medical electrical equipment — Part 1: General requirements for basic safety and essential performance</td>
<td>Follow instructions for use</td>
<td>Refer to instruction manual/Booklet.</td>
<td></td>
</tr>
<tr>
<td><em>NOTE on ME EQUIPMENT</em></td>
<td><em>Follow instructions for use</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ISO 15223-1 Symbol 5.4.4</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied.</td>
<td>Caution: Read all warnings and precautions in instructions for use</td>
<td>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.</td>
<td></td>
</tr>
<tr>
<td>ISO 60601-1 Table D.1 #10</td>
<td>Medical electrical equipment — Part 1: General requirements for basic safety and essential performance.</td>
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<td></td>
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<tr>
<td>ISO 60601-1 Table D.2 #2</td>
<td>Medical electrical equipment — Part 1: General requirements for basic safety and Essential performance.</td>
<td>General warning sign</td>
<td>Indicates a hazard of potential personal injury to patient or operator.</td>
<td></td>
</tr>
<tr>
<td>ISO 15223-1 Symbol 5.3.7</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied.</td>
<td>Temperature limit</td>
<td>Indicates the (storage) temperature limits to which the medical device can be safely exposed.</td>
<td></td>
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<tr>
<td><img src="image1" alt="Symbol" /></td>
<td>ISO 15223-1 Symbol 5.3.8</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied.</td>
<td>Humidity limitation</td>
<td>Indicates the range of (storage) humidity to which the medical device can be safely exposed.</td>
</tr>
<tr>
<td><img src="image2" alt="Symbol" /></td>
<td>ISO 15223-1 Symbol 5.2.8</td>
<td>Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied.</td>
<td>Do not use if package is damaged</td>
<td>Indicates a medical device that should not be used if the package has been damaged or opened.</td>
</tr>
<tr>
<td><img src="image3" alt="Symbol" /></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 a medical device that is not made with natural rubber latex.</td>
</tr>
<tr>
<td><img src="image4" alt="Symbol" /></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>
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