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**Quick Start Guide**

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1. Insert batteries

2. Attach belt clip and close battery compartment

3. Connect the electrodes to the leadwire

4. Apply the electrodes.
The electrode placement depends on the indication, see also chapter 2.2 / page 11

5. Connect the electrode leadwire to the device
QUICK START GUIDE...

6. Switch the therapy unit on

7. Select a program (only possible when intensity = 0)
The program depends on the indication to treat, see also chapter 2.2 / page 11

8. Set intensity for selected channel

9. After approx. 10 seconds the keys are automatically locked to prevent the treatment parameters from being changed inadvertently. To unlock the keys, press either key (channel 1 or 2).

10. To terminate the treatment simply turn off the device with the ON/OFF key. When the therapy timer is activated, the stimulator switches automatically off at the end of the programmed interval.

Note: The Belt Clip can be attached or removed as required. You can find the procedure to attach/remove the belt clip on page 14, chapter 3.1.
GENERAL INFORMATION...

- The product Direct TENS™ bears the CE marking CE-0473 (Notified Body: AMTAC Certification Services Limited) showing that it complies with the Council Directive 93/42/EEC as amended concerning medical devices and fulfills the essential requirements of Annex I of this directive. It has an internal power source and is classified as IIa equipment (MDD).

- The device has a type BF applied part.

- The device fulfills the requirements of the standard EN 60601-1 "Medical electrical equipment, Part 1: General requirements for safety" as well as the immunity requirements of the standard EN 60601-1-2 “Electromagnetic compatibility - medical electrical equipment”.

- This manual is an integral part of the device and should be kept near the device at all times. Close observance of the information given in this manual is a prerequisite for using the device as intended and for correct operation to ensure user’s safety. Please note that information pertinent to several chapters is given only once. Therefore, carefully read the manual once in its entirety.

- Using the device for purposes other than those described in this manual is not permitted.

- The safety information given in this manual is classified as follows:

  - **Warning**

    Indicates a hazard. If not avoided, the hazard can result in death or serious injury.

  - **Caution**

    Indicates a potential hazard. If not avoided, the hazard may result in minor injury and/or product/property damage.

- No part of this manual may be reproduced without written permission from DJO.

- Key to symbols used on the equipment

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Warning" /></td>
<td>Caution, consult accompanying documents</td>
</tr>
<tr>
<td><img src="image" alt="Caution" /></td>
<td>Patient part type BF - Body floating</td>
</tr>
<tr>
<td><img src="image" alt="Device" /></td>
<td>Device complies with the Council Directive 93/42/EEC about medical devices, tested and approved by AMTAC Certification Services Limited</td>
</tr>
<tr>
<td><img src="image" alt="Disposal" /></td>
<td>Do not dispose with unsorted domestic waste</td>
</tr>
</tbody>
</table>

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Storage temperature: from -20˚C to 45˚C
Max. relative humidity: 75%
Atmospheric pressure: from 700hPa to 1060hPa
1. POINTS TO NOTE BEFORE USE...

Intended Use

Direct TENS™ is a transcutaneous electrical nerve stimulator. Transcutaneous electrical nerve stimulation (TENS) uses electrical pulses that are delivered through the skin to the cutaneous (outer) and afferent (deeper) nerves to alleviate pain. Contrary to medication and cream used on the skin, there are no known side effects resulting from TENS therapy. Use Direct TENS™ only as described in this manual. Other uses of the stimulator are not permitted.

Indications

- Direct TENS™ can be used to alleviate different types of acute and chronic pain such as
  - Joint pain (e.g. knee, hip arthrosis)
  - Chronic pain originating in the spine
  - Degenerative diseases of the musculoskeletal system
  - Tension headache
  - Radiating pain (e.g. back pain, cervicobrachial syndrome)
  - Amputation stump/phantom limb pain
  - Pain from rheumatic diseases

Contraindications

Do not use Direct TENS™ in the following situations:

- If you have an implanted demand pacemaker, intracardiac defibrillator or other active implants
- Undiagnosed pain until the cause has been ascertained
- Epilepsy
- During pregnancy (unless approved by your referring gynaecologist)

Treatment should never be applied near the area of an implant, such as cochlear, pacemakers, skeletal or electrical.

Do not apply stimulation in the vicinity of metal. Remove jewellery, body piercings, buckles or any other removable metallic product or device in the area of stimulation.

Do not attempt to place electrodes on any part of the body not directly visible without assistance.

Do not stimulate at the front or side of the neck to avoid a drop in blood pressure. Furthermore it is not permitted to attach electrodes to the head.

This device should not be used for symptomatic local pain relief unless diagnosis is established or unless a pain syndrome has been diagnosed.

Biocompatibility

Those parts of the Direct TENS™ that come into contact with the user when the device is used as intended, are designed to fulfil the biocompatibility requirements of the applicable standards.
• Magnetic and electrical fields are capable of interfering with the proper performance of the device. Do not use the Direct TENS™ device in the vicinity of equipment that emits high levels of electromagnetic radiation, such as X-ray equipment, MRI devices, radio systems and mobile telephones. These devices may affect the Direct TENS™ output power. Keep the device away from such equipment and verify its performance before use (section “Treatment”). Not for use in presence of shortwave therapy device.

• Disconnect the Direct TENS™ stimulation electrodes before using electrosurgical equipment or defibrillators. Otherwise skin burns may be caused below the electrodes and the Direct TENS™ device may be destroyed.

• Avoid the simultaneous use of Direct TENS™ and electronic patient monitoring systems. Direct TENS™ may interfere with the proper functioning of these systems, compromising the monitoring quality.

• Do not use more than one stimulator at a time.

• Avoid the simultaneous use of Direct TENS™ and electronic patient monitoring systems. Direct TENS™ may interfere with the proper functioning of these systems, compromising the monitoring quality.

• Do not apply electrodes around the throat area or the anterior neck as this could cause severe muscle spasms resulting in closure of your airway, difficulty in breathing, or adverse effects on heart rhythm or blood pressure.

• Do not use the stimulator when in the bath, shower or humid environment (sauna, hydrotherapy, pools, etc.).

• Do not use the stimulator while driving, operating machinery, or during any activity in which electrical stimulation can put you at risk of injury.

• Stimulation should not be carried out over swollen, infected, and inflamed areas or skin eruptions, e.g., phlebitis, thrombophlebitis, varicose veins, etc.

• Stimulation should not be carried out over, or in proximity to, cancerous lesions.

• Never carry out the first 5 minutes of any stimulation session standing. Make sure you are seated or lying down. In rare instances, people of a nervous disposition may experience a vasovagal reaction. This is a psychological response triggered by fear of the procedure.

• Never connect the stimulation cables to an external power source due to risk of electric shock.

• Sudden temperature changes can cause condensation to build up inside the stimulator. To prevent this, allow it to reach ambient temperature before use.

• Never use the electrodes contra-laterally, i.e. do not use two poles connected to the same channel on opposite segments of the body.

• Stimulate with precaution while treating angina pectoris and the thoracic region on patients with cardiac arrhythmia.
User's hazard and comfort

- Inspect the stimulator and its accessories for integrity before use. If you detect signs of damage, do not use the stimulator.
- Use only original accessories (electrodes, cables).
- Do not use the stimulator during sleep.
- Do not open the battery compartment while the stimulator is operating.
- Use caution if you have a tendency to bleed internally, such as following an injury or fracture.
- Consult your physician prior to using the device after a recent surgical procedure, because stimulation may disrupt the healing process.
- Always switch off the stimulator before disconnecting any stimulation cables during a session.
- TENS is a symptomatic treatment and, as such, suppresses the sensation of pain that would otherwise serve as a protective mechanism. TENS devices have no curative value.
- Use caution if stimulation is carried out over areas of skin that lack normal sensation.
- Attach the electrodes in such a way that their entire surface is in contact with the skin.

Equipment damage

- Remove the batteries from the Direct TENS™ device, if it is not used for a prolonged period of time (more than approx. 3 months).
- Liquids or foreign matter (soil, metal, etc.) must not enter the stimulator. If liquids have entered the stimulator or if it was accidentally immersed in liquid, stop using it and return it to DJO for inspection.
2. HOW DOES THE DIRECT TENS™ DEVICE FUNCTION...

Via electrodes attached to the skin, Direct TENS™ sends electrical pulses to the nerves. This will block the pain impulses. Four electrodes – two for each channel – can be connected to the device. Pain relief is most efficient during stimulation, but the effect can last after the treatment. Additionally, the TENS treatment increases the blood circulation. You can use Direct TENS™ at any time for pain relief and muscle relaxation. Each therapy session should last 30 minutes minimum and can be continued for several hours.
2.1. TENS therapy principle

Two pain theories play an important role in the application and parameter settings of the Direct TENS™ device:

• The **Gate Control Theory** by WALL and MELZACK (1965)
• The **Endorphine Theory** by ERIKSON and SJÖLUND (1979)

According to the Gate Control Theory, **weak** TENS impulses block the pain impulses travelling to the brain (sensor stimulation). ERIKSON and SJÖLUND found that **strong** TENS impulses increase the release of internal substances (e.g. endorphins) that also alleviate pain (motor stimulation).

<table>
<thead>
<tr>
<th>Theory</th>
<th>Gate Control Theory</th>
<th>Endorphine Theory</th>
</tr>
</thead>
<tbody>
<tr>
<td>Principle</td>
<td>Via sensory nerves</td>
<td>Via motor nerves</td>
</tr>
<tr>
<td>Intensity</td>
<td>Low, light tingling</td>
<td>High, just bearable</td>
</tr>
<tr>
<td>Impulse Width¹</td>
<td>Short, e.g. 100 µs</td>
<td>Long, e.g. 250 µs</td>
</tr>
<tr>
<td>Frequency¹</td>
<td>100 Hz</td>
<td>2-10Hz</td>
</tr>
<tr>
<td>Muscle Contraction</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Onset Pain Relief</td>
<td>Quickly</td>
<td>Slowly (20-60 minutes)</td>
</tr>
<tr>
<td>Duration of Pain Relief</td>
<td>Short (5-15 minutes)</td>
<td>Long (30 minutes-12 hours or longer)</td>
</tr>
<tr>
<td>Treatment Duration</td>
<td>Permanent</td>
<td>30-60 minutes, 3-5 times/day</td>
</tr>
</tbody>
</table>

¹ For easier operation, intensity and pulse width are combined in Direct TENS™. (low intensity = short pulse width, high intensity = long pulse width)
### 2.2. DESCRIPTION OF THE PROGRAMS AND THE CORRESPONDING INDICATIONS...

<table>
<thead>
<tr>
<th>Program</th>
<th>Stimulation</th>
<th>Frequency (Hz)</th>
<th>General indications</th>
<th>Advantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1 impulse every 2 seconds</td>
<td>0.5</td>
<td>Kaada TENS (similar to acupuncture)</td>
<td>Ideal for sensitive people; Supports acupuncture treatment</td>
</tr>
<tr>
<td>2</td>
<td>Double pulse at 20 Hz (pulse separation 3 ms)</td>
<td>20</td>
<td>Cervical spine syndrome; Tense muscles</td>
<td>Muscle relaxation by double pulses</td>
</tr>
<tr>
<td>3</td>
<td>High frequency, 1000 Hz</td>
<td>1000</td>
<td>Acute, strong back pain (lumbar spine)</td>
<td>Strong analgesia; Brief, very intensive TENS treatment</td>
</tr>
<tr>
<td>4</td>
<td>Bi-modal</td>
<td>Channel 1 = 100Hz; Channel 2 = 4Hz</td>
<td>Tension headache; Neck / back pain; Radiating pain</td>
<td>Simultaneous treatment with high and low frequency; In 2-channel mode, channel 1 stimulation is superimposed on the low frequency stimulation in channel 2</td>
</tr>
<tr>
<td>5</td>
<td>Burst with alternating work and rest phases of 3 and 2 seconds respectively</td>
<td>Work = 100Hz; Rest = 0Hz</td>
<td>Tense muscles; Amputation stump/phantom limp pain; Herpes zoster; Reflex sympathetic dystrophy (RSD)</td>
<td>Easily tolerable stimulation for chronic pain conditions; Sensory as well as motor stimulation</td>
</tr>
<tr>
<td>6</td>
<td>Similar to program 5, but channels 1 and 2 alternating and longer work/rest times, 6 seconds respectively</td>
<td>Work = 100Hz; Rest = 0Hz</td>
<td>See program 5</td>
<td>Similar to program 5, but channels 1 and 2 alternating</td>
</tr>
<tr>
<td>7</td>
<td>Intensity decreases 40% in 0.5 second intervals</td>
<td>100</td>
<td>Lumbar back pain; Joint pain</td>
<td>Similar to massage; Effects both on the sensory and on the motor level; Avoids habituation</td>
</tr>
<tr>
<td>Program</td>
<td>Stimulation</td>
<td>Frequency (Hz)</td>
<td>General Indications</td>
<td>Advantages</td>
</tr>
<tr>
<td>---------</td>
<td>----------------------------------------------------------------------------</td>
<td>----------------</td>
<td>---------------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| 8       | Random modulation of intensity and frequency (down to 50% of set intensity and frequency modulation between 8 different frequencies, 2-150 Hz) | Random modulation | Chronic pain resisting therapy | Avoids habituation  
Sensory as well as motor stimulation                                                                                                      |
| 9       | Continuous                                                                 | 2 – 150        | Standard TENS       | Fast pain relief in acute pain conditions  
Fast acceptance of therapy  
Different programmable frequencies, e.g.  
100 Hz = Standard  
2 Hz = Similar to acupuncture                                                                                                                 |
| 10      | Burst with alternating work and rest phases of 2 seconds each              | Work = 2 - 150 Hz  
Rest = 0 Hz | Long-term treatment | Classic burst  
Pleasant form of stimulation  
Reduces muscle fatigue  
Prolongs battery life                                                                                                                      |
| 11      | Mixed frequency                                                            | Phase 1 = 2 - 150 Hz  
Phase 2 = 50% of work freq | Strong pain         | Pleasant stimulation also at higher intensities  
Permanent stimulation of the deep afferent nerve fibers with modulated muscle activation                                                                 |
| 12      | Multi modulation                                                           | 2 – 150         | Chronic pain        | Avoids habituation  
Simultaneous sensory and motor stimulation  
Fixed modulation pattern for intensity and frequency                                                                                          |
| 13      | Simple modulated pulse (SMP), intensity modulation diametrically opposite to frequency modulation according to a fixed 12-second cycle | 2 – 150         | Chronic pain        | Avoids habituation  
Simultaneous sensory and motor stimulation  
Intensity and frequency modulation according to fixed pattern, but diametrically opposed, i.e., when the intensity increases, the frequency decreases and vice versa. |
Frequency Selection for Programs 9 to 13

<table>
<thead>
<tr>
<th>2 – 60 Hz</th>
<th>60 – 150 Hz</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preferred in the treatment of <strong>chronic</strong> pain</td>
<td>Preferred in the treatment of <strong>acute</strong> pain</td>
</tr>
</tbody>
</table>

With the standard TENS programs 9 to 13, the frequency can be adapted manually. Possible settings: 2, 10, 20, 40, 60, 100, 125, 150 Hz

**Factory defaults for programs 9 to 12 is 100 Hz and 125 Hz for program 13.**

**Changing the frequency:**

1. Turn on the device and select one of the programs 9 to 13

2. **Simultaneously** press the program selection keys and release them.

3. Using the intensity keys , choose the set frequency.

4. Press the two program selection keys again or switch the device off to save the settings.
3. PREPARATION...

3.1. Inserting Batteries

- Adjust the belt clip until it points to the right at a 90° angle. (Figure 3-1)
- Push the battery cover down and lift. (Figure 3-2)
- Insert the batteries as shown in the illustration. Observe the correct polarity, see label in battery compartment. (Figure 3-3)
- Reinstall the battery cover and close the compartment.

Notes:
- Use only new AA type batteries.
- You may or may not use the belt clip, as preferred. Open the battery compartment. If you wish to remove the clip, pull it out towards the left. If you wish to attach the clip, push it into the holder from the left. When you close the battery compartment, the belt clip is automatically secured onto the device. (Figure 3-4)
- Dispose of the worn out batteries in accordance with local and national regulations.

3.2. Applying Electrodes, Connecting Leadwires

- First connect the electrode leadwires to the electrodes (Figure 3-5). (The colour of the electrode connectors is irrelevant.)
- Peel the electrodes off their protective paper. Keep the protective paper and the bag, because the electrodes will be reattached to the protective paper after use and stored in the bag (see also 3.3.2 “Care of the Electrodes”).
- Carefully apply the electrodes on the skin (see also 3.3.3 “Electrode Placement”).
- Connect the electrode leadwire(s) to the Direct TENS™ device (Figure 3-6).

Figure 3-1
Turning the belt clip

Figure 3-2
Opening the battery compartment

Figure 3-3
Inserting the batteries

Figure 3-4
Removing/attaching the belt clip

Figure 3-5
Connecting the electrode leadwire to the electrodes

Figure 3-6
Connecting the electrode leadwire to the device
3.3. Selection, Care and Placement of the Electrodes

3.3.1. Electrode Selection

Use large electrodes (e.g. 50 x 90 mm, to be purchased separately) for large body areas (e.g. back, leg) and for general conditions of pain.
Use small electrodes (e.g. 50 x 50 mm) for small body areas (e.g. face, hand) and for deep, local pain.

3.3.2. Care of the Electrodes

When properly handled and maintained, the supplied electrodes can be used 20 times or more.

Important for a long service life:

- Clean the skin application sites with mild soap water before attaching the electrodes. After cleaning, thoroughly rinse with water and dry the skin carefully.
- Dry electrodes with poor adhesion can be reconditioned as follows: apply a small quantity of water to the adhesive surface with your fingertip.
- If you face bad contact with the skin or repeated open lead detection, change the electrodes.
- Remove electrodes by pulling on their edges. Do not pull on the leadwire.
- After use, reattach the electrodes to their protective paper. Store the electrodes in their bags.
- Store the electrodes in a refrigerator, if possible. Do not store them in warm rooms.
- We recommend shaving skin sites where electrodes will be applied, if very hairy. Shaving irritates the skin. Therefore wait 24 hours after shaving before you attach the electrodes. Then you may start therapy.
- Do not leave the electrodes attached to your skin for a prolonged period of time. Remove the electrodes after each use. Apply the electrodes on different sites to avoid skin irritations. For the same reason clean the skin thoroughly after treatment. If you observe skin irritations, consult your physician and suspend therapy until clarification.
3.3.3. Electrode Placement

If your physician showed you the best application points, we recommend that you use them. Otherwise figures 3-7 to 3-11 show possible electrode configurations. Figures 3-12 to 3-21 show electrode configurations for different indications. However, check that the configuration is appropriate and adapt, if necessary. Depending on the site and cause of the pain, electrodes may be placed on acupuncture points or in specific dermatomic areas. In other situations we recommend applying the electrodes around the center of the pain at a distance of 3 to 5 cm (where you feel the pain).

Notes:

- Concerning the choice of indication-specific programs, please refer to the Notes in section 2.2.
- Before applying the electrodes, observe the care instructions in section 3.3.2.
- If appropriate, you can also use only two electrodes (one channel).

Caution

Failed stimulation, skin irritation, malfunction -
- Use only the original electrodes supplied with the system and replacement electrodes provided by DJO.
- Attach the electrodes on intact skin only, avoid skin areas with reduced sensitivity.
- Ensure that good contact is achieved between electrode and skin. Although the stimulator switches off when the electrode-skin contact resistance is too high, poor electrode techniques may cause skin irritations under the electrodes.
Indications:
e.g. General pain in the shoulder, bursitis

Figure 3-14 Electrode application on the cervical spine
Indications:
e.g. pain caused by intervertebral disk or vertebral arch joint problems, cervical spine syndrome, cervical syndrome, tension headache, migraine
Recommended electrode placement: Diagonal

Figure 3-15 Electrode application on the cervical spine Picture to be replaced
Indications:
e.g. for conditions of pain in the muscles or soft parts, see also Figure 3-15
Recommended electrode placement: Parallel
Figure 3-16 Electrode application on the knee joint
**Indications:**
e.g. arthrosis of the knee joint (gonarthrosis),
generalized pain in the knee, TEP
**Recommended electrode placement:**
Attach electrodes above superficial skin nerves
or acupuncture points around the knee joint.
The electrode configuration may be parallel,
medial, lateral or crosswise above or
below the knee.

Figure 3-17 Electrode application on the back
**Indications:**
e.g. lumbar spine syndrome,
lumboischialgia, pseudoradicular back pain
**Recommended electrode placement:**
paraspinal, proximal and distal to the pain
area. Channels 1 and 2 diagonal

Figure 3-18 Electrode application on the back
**Indications:**
Radicular (radiating) pain
**Recommended electrode placement:**
Channel 1 proximal and distal to the pain
area, channel 2 above the nerve.

Figure 3-19 Electrode application on the back
**Indications:**
Radicular (radiating) pain (alternative)
**Recommended electrode placement:**
Channel 1 proximal and distal to the pain
area, channel 2 above the nerve.

Figure 3-20 Electrode application for
phantom limb pain, version 1

Figure 3-21 Electrode application for
phantom limb pain, version 2
4. TREATMENT...

4.1. Starting Therapy

• Switch the stimulator on.

Once switched on, the display will briefly show the software version. A functional test where all display indicators appear for a short time is performed. The initial screen appears next (Figure 4-2).

When the initial screen is displayed, the device has successfully passed the functional test. If the letter “E” is displayed instead of the initial screen, the stimulator is defective and must be replaced. Do not use this stimulator any more.

On the initial screen you will see:
- The remaining stimulation duration
- The intensity of the stimulation (indication of the selected intensity level, adjustable in steps of 0.5 from 0 to 60)
- The selected program

- Select the program either via the quick select keys or via the program selection keys.
- Increase the stimulation intensity for a channel (1 or 2) by pressing the corresponding key.

Increase the intensity with great care and in small increments. Select a level which causes a pleasant sensation that is felt clearly.

Additionally, icons may appear on the screen:

- The keys are locked (automatic function): To prevent inadvertent activation, the keys are automatically locked 10 seconds after the current intensity has been set. The keys can be unlocked with or by switching the device off.
- The electric circuit is interrupted (see chapter 6 “What to do, if...”)
- Batteries need to be replaced. When you see this icon, replace the batteries as soon as possible.

Operation Information

- You can interrupt therapy at any time with the ON/OFF switch.
- If the stimulator is not used, it switches off automatically after approx. 5 minutes.
- When the therapy timer is activated, the device switches automatically off at the end of the programmed interval. The remaining therapy time is always indicated on the display.
- The program can only be changed when the intensity in both channels is 0.
4.2. Ending Therapy

The default setting of the Direct TENS™ device is continuous operation. If you want to end the therapy, switch off the device with the ON/OFF switch. When the therapy timer is activated, the device switches automatically off at the end of the programmed interval. The remaining time is indicated at the top of the display. Check that the Direct TENS™ stimulator is switched off before you remove the electrodes.

• If the therapy timer is not activated, switch off the Direct TENS™ stimulator with the ON/OFF switch.

• Remove the electrodes very carefully. Do not pull on the leadwires, but on the electrode.

• Reattach the electrode to its protective paper. Be sure to attach the electrode to the side marked “on”, not to the side marked “no”.

• Disconnect the electrodes from the leadwire.

• Clean the skin with a mild soap solution.

• Electrodes no longer fit for use can be disposed of with the normal domestic waste.
5. SPECIAL DIRECT TENS™ FUNCTIONS...

5.1. Therapy Timer

The Direct TENS™ stimulator comes with a therapy timer function. When the timer is activated, the remaining therapy time appears on the display. When this interval has elapsed, the device switches off automatically.

You can set therapy times of:

- 1 to 59 minutes in 1-minute increments
- 1 to 24 hours in 30-minutes increments
- Continuous operation, displayed as CONT. (this is the default setting), the timer is not activated

Follow these steps to activate the timer:

- Press the quick select program key
  low back/hip and hold it depressed.
- Then press the ON/OFF key to switch the device on.
- The display indicates “Time” and the currently set therapy time (default setting: CONT).
- Press the intensity control keys for channel 2 to set the therapy time. With you select a longer time, with , a shorter time. Go back to 00:00 to get the Continuous mode (CONT).
- To save the timer settings, turn off the stimulator. When you turn the stimulator on again, the timer setting you saved is displayed.

5.2 Factory Defaults Settings

- Hold the program selection key depressed.
- Then press the ON/OFF key to switch the device on.
- The display shows DATA 000 000 flashing.
- Next press the two intensity decrease keys simultaneously for channels 1 and 2

  to reset the device to the factory defaults:

  - All stored data will be deleted.
  - The timer function is deactivated.
  - The program lock is deactivated.
  - The frequency for all programs is reset to the respective default value.
  - Program 9 is selected.
  - The initial screen is displayed.
6. WHAT TO DO, IF...

...the stimulation feels unpleasant or different from previous sessions

• Check whether
  - The device settings were changed.
  - The electrodes are correctly placed and applied on the skin. Electrodes with poor adhesion must be replaced.
  - The skin is irritated.

...the stimulation is weak or not felt

• Check whether
  - The electrodes are properly applied to the skin. Electrodes with poor adhesion must be replaced.
  - The battery replacement symbol is displayed and the batteries need to be replaced (see the “Battery Replacement” section in chapter 7).
  - The “electric circuit interrupted” icon is displayed. This icon appears when the resistance between electrode and skin is too high. Reasons may be a poor electrode attachment or an interrupted electric circuit.
     When the electric circuit is interrupted, the intensity drops to 0. In this case check whether the electrode leadwire is correctly connected to the device and whether the electrodes are properly connected. If the problem persists, the electrode leadwire is probably broken and must be replaced.

...the letter “E” is displayed instead of the initial screen

In this case the device is defective and must not be used. Return the device to DJO for replacement.

...the device cannot be switched on

• Check whether
  - The batteries are inserted
  - The batteries are correctly inserted
  - The batteries are charged.
7. CARE, STORAGE, BATTERY REPLACEMENT, DISPOSAL...

Care, Maintenance

- Switch off the device before cleaning. Then clean it with a moist cloth. Liquids must not enter the device. If necessary, apply a drop of dishwashing liquid to remove stubborn stains.

- Do not use abrasive substances or other domestic cleaning agents.

- To clean the leadwire, disconnect it from the device and wipe it down with a moist cloth. Never immerse the leadwire in liquids.

- Direct TENS™ does not require regular maintenance.

Storage

- Always store the Direct TENS™ stimulator in its original pouch.

- Keep the device and the pouch out of reach of children.

- Store the device in a cool and dry place. Do not expose the device to direct sunlight.

- Remove the batteries from the device, if it is not used for a prolonged period of time (more than approx. 3 months).

Battery Replacement

Use only Alkaline batteries 15 A LR6 size AA 1.5 Volt or NiMH rechargeable batteries size AA 1.2 Volt (rechargeable batteries and charger not provided)

- Switch off the Direct TENS™ device.

- Turn the belt clip 90° to the right (see also section 3.1).

- Push the battery cover down, then up (see also section 3.1).

- Insert the batteries. Observe the correct polarity (see also section 3.1).

- Reinstall the battery cover and close the compartment.

- Dispose of the old batteries, observing the local regulations.

Disposal at the End of Its Service Life

The symbol indicates that the waste of electrical and electronic equipment must not be disposed as unsorted domestic waste and must be collected separately. Please contact DJO for information about the possible recycling of the product or comply with your local regulation.
8. ORDERING INFORMATION, SPECIFICATIONS...

Ordering Information

Electrodes used up?
Questions about the product?
We at DJO would be happy to help!

To avoid wrong deliveries, please indicate the part numbers when ordering products:

<table>
<thead>
<tr>
<th>Part Number</th>
<th>Product</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>42190</td>
<td>Electrodes, Square 5x5 cm Durastick, Wire, pkg. of 4 electrodes</td>
<td></td>
</tr>
<tr>
<td>42191</td>
<td>Electrodes, Rectangular 5x9 cm Durastick, Wire, pkg. of 4 electrodes</td>
<td></td>
</tr>
<tr>
<td>193068-100</td>
<td>Lead wire, 100cm (40 inches)</td>
<td></td>
</tr>
</tbody>
</table>

To reorder, please contact:

DJO France
Centre Européen de Fret
64990 Mouguerre
France
Phone: +33 (0)5 59 52 86 90
Fax: +33 (0)5 59 52 86 91
E-mail: sce.cial@DJOglobal.com
Internet: www.DJOglobal.eu

8.1. Information related to electromagnetic compatibility (EMC)

The Direct TENS™ is designed to be used in typical domestic or clinical environments and approved according to the EMC safety standard of EN 60601-1-2.

This device emits very low levels in the radio frequency (RF) interval and is therefore not likely to cause any interference with nearby electronic equipment (radios, computers, telephones etc.).

The Direct TENS™ is designed to withstand foreseeable disturbances originating from electrostatic discharges, mains supply magnetic fields or radio frequency transmitters (such as mobile telephones).

For more detailed information regarding electromagnetic emission and immunity, please contact DJO.

Specifications

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of channels</td>
<td>2</td>
</tr>
<tr>
<td>Constant Voltage</td>
<td>Up to a resistance of 1000 ohms</td>
</tr>
<tr>
<td>Output Intensity</td>
<td>0...60, adjustable in 0.5 steps</td>
</tr>
<tr>
<td>Maximum Output</td>
<td>40 mA @ 1000 ohms, 250 μs</td>
</tr>
<tr>
<td>Waveform</td>
<td>Asymmetrical biphasic square impulse</td>
</tr>
<tr>
<td>Frequency Range</td>
<td>0,5 – 1000 Hz</td>
</tr>
<tr>
<td>Impulse Duration</td>
<td>Determined by intensity setting, 0 – 250 μs</td>
</tr>
<tr>
<td>Power Supply</td>
<td>2 x 1.5 V AA disposable batteries or 2 x 1.2 V AA rechargeable batteries</td>
</tr>
<tr>
<td>Current consumption</td>
<td>50 mA</td>
</tr>
<tr>
<td>for 1 channel, 200 μs</td>
<td></td>
</tr>
<tr>
<td>impulse duration, 100 Hz</td>
<td></td>
</tr>
<tr>
<td>and 40 V</td>
<td></td>
</tr>
<tr>
<td>Ambient conditions (operation)</td>
<td>Temperature 10°C to 40°C</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>Relative humidity 30...75 %, no condensation</td>
</tr>
<tr>
<td>Atmospheric pressure</td>
<td>Atmospheric pressure 700 to 1060 hPa</td>
</tr>
<tr>
<td>Ambient conditions (storage, transport)</td>
<td>Temperature 10°C to 40°C</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>Relative humidity 30...75 %, no condensation</td>
</tr>
<tr>
<td>Atmospheric pressure</td>
<td>Atmospheric pressure 500 to 1060 hPa</td>
</tr>
<tr>
<td>Dimensions (HxWxD)</td>
<td>110 x 70 x 30 mm</td>
</tr>
<tr>
<td>Weight</td>
<td>165 g (including batteries)</td>
</tr>
</tbody>
</table>