



Vectra[®] Neo Clinical Therapy System User Manual

Operator and
Installation
Instructions



Rx Only

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FOREWORD

This manual is intended for users of Vectra® Neo Clinical Therapy System. It contains general information on operation, precautionary practices, and maintenance. In order to maximize use, efficiency, and the life of the system, please read this manual thoroughly and become familiar with the controls, as well as the accessories before operating the system.

In addition to the above information, this manual contains care and installation instructions for the optional Cart, Channel 1/2 Electrotherapy module, Channel 1/2 Electrotherapy module + sEMGmodule, Channel 3/4 Electrotherapy module, Laser module, and Ultrasound module for the users of the Vectra® Neo Clinical Therapy System.

Specifications put forth in this manual were in effect at the time of publication. However, owing to DJO's policy of continual improvement, changes to these specifications may be made at any time without notification on the part of DJO.

Before administering any treatment to a patient, the users of this equipment should read, understand, and follow the information contained in this manual for each mode of treatment available, as well as the indications, contraindications, warnings, and precautions. Consult other resources for additional information regarding the application of electrotherapy, ultrasound, and laser.

PRECAUTIONARY INSTRUCTIONS

The precautionary instructions found in this section and throughout this manual are indicated by specific symbols. Understand these symbols and their definitions before operating this equipment. The definition of these symbols are as follows:

 **CAUTION**

Text with a "CAUTION" indicator explains possible safety infractions that have potential to cause minor or moderate injury or damage to the equipment.

 **WARNING**

Text with a "WARNING" indicator explains possible safety infractions that will potentially cause serious injury and equipment damage.

 **DANGER**

Text with a "DANGER" indicator will explain possible safety infractions that are imminently hazardous situations that would result in death or serious injury.



Text with a "DANGEROUS VOLTAGE" indicator serves to inform the user of possible hazards resulting in the electrical charge delivered to the patient in certain treatment configurations of TENS waveforms.



Warning; Laser beam



Explosion Hazard - Text with an "Explosion Hazard" indicator will explain possible safety infractions if this equipment is used in the presence of flammable anesthetics, mixture with air, oxygen, or nitrous oxide.



Wear eye protection

NOTE: Throughout this manual, "NOTE" indicators provide helpful information regarding the particular area of function being described.

GENERAL TERMINOLOGY

The following are definitions for the terminology used throughout this manual. Study these terms to become familiar with them for ease of system operation and control functionality of the Vectra® Neo Clinical Therapy System.

SYSTEM SOFTWARE SYMBOLS

| | |
|---|--|
|  | Back Arrow |
|  | Home |
|  | Increase/Decrease Parameter |
|  | Scroll Up or Down in a text box |
|  | Select |
|  | Page Up |
|  | Page Down |
|  | Customize |
|  | Save Data |
|  | When pressed, will print the screen contents or Patient Treatment Results Report to the USB flash drive |
|  | Indicates a USB flash drive is inserted |
|  | Patient Remote/Laser Shut Off Icon Indicates the Remote is plugged in |

| | |
|--|----------------------------|
|  | Stim |
|  | Electrode Placement |
|  | Ultrasound |
|  | Combo |
|  | sEMG |
|  | Laser |
|  | CPS |
|  | Custom Protocols |
|  | Patient Data |
|  | Anatomical Library |

DESCRIPTION OF DEVICE MARKINGS

The markings on the unit are assurance of its conformity to the highest applicable standards of medical equipment safety and electromagnetic compatibility. One or more of the following markings may appear on the device:

- Refer to Instructional Manual Booklet 
- Equipment capable of delivering output values in excess of 10 mA r.m.s. or 10V r.m.s. averaged over any period of 5s 
- Testing Agency 
- Dangerous Voltage 
- Electrical Type B 
- Electrical Type BF 
- Laser 
- Ultrasound 
- Stim 
- Start 
- Stop 
- Pause 
- Intensity 
- Lock/Unlock 
- ON/OFF 
- Laser Stop Switch 

This unit is considered to be a Class 3B laser product and thus emits visible and invisible laser radiation (IR). Avoid direct eye exposure to the Laser beam. The symbol to the right is located on the back of the applicator and indicates the active radiant surface (the area on the applicator that emits infrared laser energy and the direction of the beam of light)



MRI Unsafe (device, its components, and accessories are not to be present in an MRI or CT environment) 

ELECTROTHERAPY INDICATIONS

Indications

For VMS (Pulsed Mode, Burst Mode, or FR Mode), Russian, Monophasic Hi-Volt (NMES) & Interferential, and Premodulated (IFS):

- Relaxation of muscle spasms
- Prevention or retardation of disuse atrophy
- Increasing local blood circulation
- Muscle re-education
- Maintaining or increasing range of motion
- Immediate postsurgical stimulation of calf muscles to prevent venous thrombosis

Additionally for Microcurrent, Interferential, Premodulated (IFS), VMS (Pulsed Mode, Burst Mode, or FR Mode), Asymmetrical Biphasic (TENS), and HANS:

- Symptomatic relief or management of chronic, intractable pain
- Post-traumatic acute pain
- Post-surgical acute pain

For DC (Direct Current) Continuous Mode:

- Relaxation of muscle spasm

For FES:

- Stimulation of the muscles of the leg and ankle of partially paralyzed patients to provide flexion of the foot and thus improve the patient's gait.

Contraindications

The Vectra® Neo Clinical Therapy System should NOT be used under the following conditions:

- Do not use for symptomatic local pain relief unless etiology is established or unless a pain syndrome has been diagnosed.
- Do not use when cancerous lesions are present in the treatment area.
- Do not apply stimulation over swollen, infected, inflamed areas or skin eruptions (e.g., phlebitis, thrombophlebitis, varicose veins, etc.).
- Do not use when patient is suspected or known to have infectious disease and/or disease where it is advisable, for general medical purposes, to suppress heat or fevers.
- Do not place electrode placements to the carotid sinus region (anterior neck) or transcerebrally (through the head).
- Do not use on pregnant women. Safety has not been established for the use of therapeutic electrical stimulation during pregnancy.

- Do not use powered muscle stimulators or TENS waveforms on patients with cardiac demand pacemakers.
- Do not use Vectra® Neo Clinical Therapy System on patients who have or have had implantable neurostimulating cardiac demand pacemakers, ICD, or other implantable electronic devices.
- Do not use Vectra® Neo Clinical Therapy System on patients with body worn electro mechanical medical devices, i.e. insulin pump.
- Do not use this system in an MRI or CT environment. The Vectra® Neo Clinical Therapy System, its components, and accessories are not to be present in an MRI or CT environment.

Additional Precautions

- Use caution for patients with suspected or diagnosed heart problems.
- Use caution for patients with suspected or diagnosed epilepsy.
- Use caution in the presence of the following:
 - When there is a tendency to hemorrhage following acute trauma or fracture
 - Following recent surgical procedures when muscle contraction may disrupt the healing process
 - Over a menstruating or pregnant uterus
 - Over areas of the skin that lack normal sensation
- Some patients may experience skin irritation or hypersensitivity due to the electrical stimulation or electrical conductive medium. The irritation can usually be reduced by using an alternative conductive medium or an alternative electrode placement.
- Electrode placement and stimulation settings should be based on the guidance of the prescribing practitioner.
- Powered muscle stimulators should be used only with the lead wires and electrodes recommended for use by the manufacturer.
- With TENS waveforms, isolated cases of skin irritation may occur at the site of electrode placement following long-term application.
- The effective management of pain by TENS waveforms is highly dependent upon patient selection by a person qualified in pain management.

Adverse Effects

- Skin irritation and burns beneath the electrodes have been reported with the use of powered muscle stimulators.
- Potential adverse effects with TENS are skin irritation and electrode burns.

SEMG & STIM INDICATIONS

Indications

For EMG triggered Stim:

- Stroke rehab by muscle re-education
- Relaxation of muscle spasms
- Prevention or retardation of disuse atrophy
- Increase local blood circulation
- Muscle re-education
- Maintaining or increasing range of motion

For EMG:

- To determine the activation timing of muscles for:
 - Retraining of muscle activation
 - Coordinating of muscle activation
- Any indication of the force produced by muscle for control and maintenance of muscle contractions
 - Relaxation muscle training
 - Muscle re-education

Contraindications

The Vectra® Neo Clinical Therapy System should not be used under the following conditions:

- Do not use for symptomatic local pain relief unless etiology is established or unless a pain syndrome has been diagnosed.
- Do not use when cancerous lesions are present in the treatment area.
- Do not apply stimulation over swollen, infected, inflamed areas or skin eruptions (e.g., phlebitis, thrombophlebitis, varicose veins, etc.).
- Other contraindications are patients suspected of carrying serious infectious disease and/or disease where it is advisable, for general medical purposes, to suppress heat or fevers.
- Do not place electrode placements to the carotid sinus region (anterior neck) or transcerebrally (through the head).
- Safety has not been established for the use of therapeutic electrical stimulation during pregnancy.
- Do not use powered muscle stimulators or TENS waveforms on patients with cardiac demand pacemakers.
- Do not use Vectra® Neo Clinical Therapy System on patients who have or have had implantable neurostimulating cardiac demand pacemakers, ICD, or other implantable electronic devices.

- Do not use Vectra® Neo Clinical Therapy System on patients with body worn electro mechanical medical devices, i.e. insulin pump.
- Do not use this system in an MRI or CT environment. The Vectra® Neo Clinical Therapy System, its components, and accessories are not to be present in an MRI or CT environment.

Additional Precautions

- Use caution for patients with suspected or diagnosed heart problems.
- Use caution for patients with suspected or diagnosed epilepsy.
- Use caution in the presence of the following:
 - When there is a tendency to hemorrhage following acute trauma or fracture
 - Following recent surgical procedures when muscle contraction may disrupt the healing process
 - Over a menstruating or pregnant uterus
 - Over areas of the skin that lack normal sensation
- Some patients may experience skin irritation or hypersensitivity due to the electrical stimulation or electrical conductive medium. The irritation can usually be reduced by using an alternative conductive medium or an alternative electrode placement.
- Electrode placement and stimulation settings should be based on the guidance of the prescribing practitioner.
- Powered muscle stimulators should be used only with the lead wires and electrodes recommended for use by the manufacturer.
- With TENS waveforms, isolated cases of skin irritation may occur at the site of electrode placement following long term application.
- The effective management of pain by TENS waveforms is highly dependent upon patient selection by a person qualified in the management of pain patients.

Adverse Effects

- Skin irritation and burns beneath the electrodes have been reported with the use of powered muscle stimulators.
- Potential adverse effects with TENS are skin irritation and electrode burns.

ULTRASOUND INDICATIONS

Indications

Application of therapeutic deep heat for the treatment of selected sub-chronic and chronic medical conditions such as:

- Relief of pain, muscle spasms, and joint contractures
- Relief of pain, muscle spasms, and joint contractures that may be associated with:
 - Adhesive capsulitis
 - Bursitis with slight calcification
 - Myositis
 - Soft tissue injuries
 - Shortened tendons due to past injuries and scar tissues
- Relief of sub-chronic and chronic pain and joint contractures resulting from:
 - Capsular tightness
 - Capsular scarring

Contraindications

- This device should not be used for symptomatic local pain relief unless etiology is established or unless a pain syndrome has been diagnosed.
- This device should not be used when cancerous lesions are present in the treatment area.
- Other contraindications are patients suspected of carrying serious infectious disease and disease where it is advisable for general medical purposes to suppress heat or fevers.
- This device should not be used over or near bone growth centers until bone growth is complete.
- This device should not be used over the thoracic area if the patient is using a cardiac pacemaker.
- This device should not be used over a healing fracture.
- This device should not be used over or applied to the eye.
- This device should not be used over a pregnant uterus.
- Tissue necrosis might result if the device is used on ischemic tissues in individuals with vascular disease, where the blood supply would not keep up with the metabolic demand.
- Do not use Vectra® Neo Clinical Therapy System on patients who have or have had implantable neurostimulating cardiac demand pacemakers, ICD, or other implantable electronic devices.

- Do not use Vectra® Neo Clinical Therapy System on patients with body worn electro mechanical medical devices, i.e. insulin pump.
- Do not use this system in an MRI or CT environment. The Vectra® Neo Clinical Therapy System, its components, and accessories are not to be present in an MRI or CT environment.

Additional Precautions

Additional precautions should be used when ultrasound is used on patients with the following conditions:

- Over an area of the spinal cord following a laminectomy, i.e., when major covering tissues have been removed
- Over anesthetic areas
- On patients with hemorrhagic diatheses

LASER INDICATIONS

Indications

To provide topical heating for the following:

- Temporary increase in local blood circulation
- Temporary relief of minor muscle and joint aches, pains, and stiffness
- Relaxation of muscles
- Temporary relief of muscle spasms
- Temporary relief of minor pain and stiffness associated with arthritis

Contraindications

The Vectra® Neo Clinical Therapy System Laser should NOT be used:

- Where analgesia may mask progressive pathology, and where the practitioner would normally avoid the use of any other analgesia in order to retain the beneficial aspects of pain
- For direct aim into the eyes of humans over areas injected with steroids in the past 2-3 weeks
- Over areas that are suspicious or contain potentially cancerous tissue
- Over areas of active hemorrhage
- Over a pregnant uterus
- Over the neck (thyroid or carotid sinus region) or chest (vagus nerve or cardiac region of the thorax)
- Directly over areas with open wounds, unless covered with a clear protective barrier
- Treatment over sympathetic ganglia
- For symptomatic local pain relief unless etiology is established or unless a pain syndrome has been diagnosed
- On patients suspected of carrying serious infectious disease and/or disease where it is advisable, for general medical purposes, to suppress heat or fevers
- Over or near bone growth centers until bone growth is complete
- Over the thoracic area if the patient is using a cardiac pacemaker
- Over or applied to the eye
- On ischemic tissues in individuals with vascular disease where the blood supply would be unable to follow the increase in metabolic demand and tissue necrosis might result
- Do not use Vectra® Neo Clinical Therapy System on patients who have or have had implantable neurostimulating cardiac demand pacemakers, ICD, or other implantable electronic devices.

- Do not use Vectra® Neo Clinical Therapy System on patients with body worn electro mechanical medical devices, i.e. insulin pump.
- Do not use this system in an MRI or CT environment. The Vectra® Neo Clinical Therapy System, its components and accessories, are not to be present in an MRI or CT environment.

Additional Precautions

Additional precaution should be used when the Laser is used on patients with the following conditions:

- Over an area of the spinal cord following a laminectomy, i.e., when major covering tissues have been removed
- Over anesthetic areas
- On patients with hemorrhagic diatheses

Preventing Adverse Effects

Perform the following procedures to avoid the negative effects of Laser therapy:

- Inspect the treatment area during and following treatment, and discontinue if an adverse reaction does occur.
- Higher output levels have a greater potential for patient discomfort. Choose a lower dosage to reduce output or select a pulsed duty cycle to decrease patient discomfort.

Factors that Affect Treatment

The following factors may affect Laser treatment:

- Color of skin (light or dark)
- Age of lesion
- Depth of lesion
- Sensitivity of patient
- Type of tissue
- Medications that increase sensitivity to light

PRODUCT DESCRIPTION

The Vectra® Neo Clinical Therapy System is a modular system used with or without an optional Cart, allowing for the inclusion of Channel 1/2 Electrotherapy module with or without sEMG, Channel 3/4 Electrotherapy module, Laser module, and Ultrasound module.

To maximize functionality and life of Vectra® Neo, be sure to:

- Stay current with the latest clinical developments in the field of electrotherapy, ultrasound, laser therapy, sEMG, and sEMG + electrotherapy.
- Observe all applicable precautionary measures for treatment.
- Keep informed of appropriate indications and contraindications for the use of the Vectra® Neo Clinical Therapy System.

NOTE: This equipment is to be used only under the prescription and supervision of a licensed medical practitioner.

OPERATOR INTERFACE

The Vectra® Neo Clinical Therapy System Operator Interface contains all the functions and controls necessary for operator access to all operator utilities, modalities, and parameters for modification and system set up.

1. Color Display
2. Intensity Dial (Gray outer ring)
3. Start/Pause button
4. Stop button
5. ON/OFF switch
6. Ultrasound Applicator holder, left and right sides
7. Laser Applicator holder, left and right sides
8. Patient Remote/Laser Interrupt Switch port
9. Mains Power Cord
10. Rear Access Panel
11. Serial Label
12. USB Flash Drive Port (USB flash drive not included)
13. Tilt Screen
14. Swivel function
15. Laser Interlock Port and Icon
16. Lead wire holders

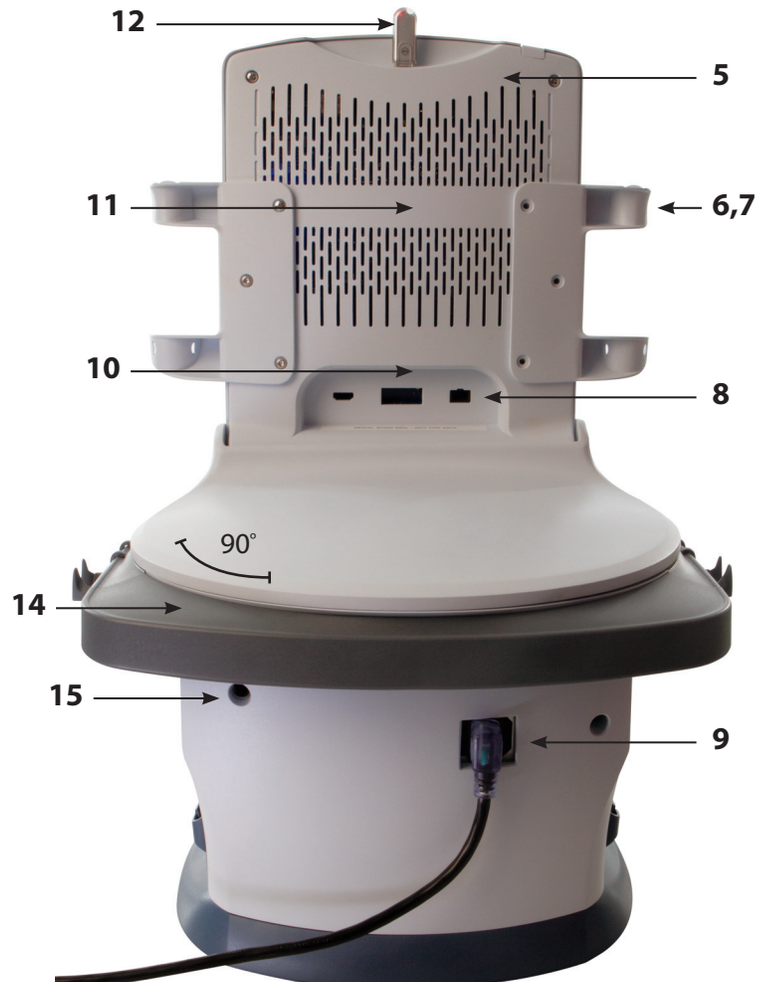
Side Holders



Front Controls



Rear Access Panel



**CAUTION**

- Read, understand, and practice the precautionary and operating instructions. Know the limitations and hazards associated with using any electrical stimulation, Laser device, or ultrasound device. Observe the precautionary and operational decals placed on the unit.
- All modalities should be routinely checked before each use to determine that all controls function normally, especially that the intensity control does properly adjust the intensity of the ultrasonic power output in a stable manner. Also, determine that the treatment time control does actually terminate ultrasonic power output when the timer reaches zero.
- DO NOT use sharp objects such as a pencil point or ballpoint pen to operate the buttons on the control panel.
- This unit should be operated at 10°C to 45°C and 0% to 90% relative humidity. The unit should be transported and stored at 0°C to 60°C and 0% to 95% relative humidity.
- Handle Ultrasound Applicator and Laser Applicator with care. Inappropriate handling may adversely affect its characteristics.
- Before each use, inspect Ultrasound Applicator for cracks, which may allow the ingress of conductive fluid.
- Inspect Applicator cables and associated connectors before each use.
- Device is designed to comply with electromagnetic safety standards. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with instructions, may cause harmful interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. Harmful interference to other devices can be determined by turning this equipment on and off. Try to correct the interference using one or more of the following:
 - Reorient or relocate the receiving device.
 - Increase the separation between the equipment.
 - Connect the equipment to an outlet on a different circuit from that to which the other device(s) are connected and consult the factory field service technician for help.
 - Consult your authorized DJO dealer for help.
- Do not operate this unit when connected to any unit other than DJO devices or accessories specifically described in user or service manuals.
- Use of controls, adjustments or performance of procedures other than those specified herein may result in hazardous exposure to Laser energy.
- DO NOT disassemble, modify, or remodel the unit or accessories. This may cause unit damage, malfunction, electrical shock, fire, or personal injury.

**CAUTION**

- Failure to use and maintain the Vectra® Neo Clinical Therapy System, its modules, and its accessories in accordance with the instructions outlined in this manual will invalidate the warranty.
- DO NOT permit foreign materials, liquids, or cleaning agents (including, but not limited to, inflammables, water, and metallic objects) to enter the unit to prevent unit damage, malfunction, electrical shock, fire, or personal injury.
- If you have difficulty operating the unit after carefully reviewing this user manual, contact your DJO dealer for assistance.
- DO NOT remove the cover. Doing so may cause unit damage, malfunction, electrical shock, fire, or personal injury. There are no user-serviceable parts inside the unit. If a malfunction occurs, discontinue use immediately and consult dealer for repair service.
- Use of parts or materials other than DJO's can degrade minimum safety.
- The Vectra® Neo Clinical Therapy System is not designed to prevent the ingress of water or liquids. Ingress of water or liquids could cause malfunction of internal components of the system and therefore create a risk of injury to the patient.
- DO NOT operate the Vectra® Neo Clinical Therapy System within the vicinity or environment as any microware and RF shortwave diathermy system.
- DO NOT operate the Vectra® Neo Clinical Therapy System within the vicinity or environment as an ultrasonic diathermy system. The Ultrasound (diathermy) Module of the Vectra® Neo Clinical Therapy System does not require separation distance.
- DO NOT use electrodes with an active area less than 7.92 cm², as there will be a risk of suffering a burn injury. Always exercise caution with current densities more than 2mA/cm². Refer to the CURRENT DENSIT CALCULATIONS FOR DJO ELECTRODES table in Appendix 3.

**WARNING**

- U.S.A. Federal Law restricts these devices to sale by, or on the order of, a physician or licensed practitioner. This device should be used only under the continued supervision of a physician or licensed practitioner.
- Be sure to read all instructions for operation before treating patient.
- Make certain the unit is electrically grounded by connecting only to a grounded electrical service receptacle conforming to the applicable national and local electrical codes.
- Care must be taken when operating this equipment around other equipment. Potential electromagnetic or other interference could occur to this or to the other equipment. Try to minimize this interference by not using other equipment in conjunction with it.
- The safety of TENS waveforms for use during pregnancy or birth has not been established.
- TENS is not effective for pain of central origin. (This includes headache.)
- TENS waveforms have no curative value.
- Electronic monitoring equipment (such as ECG monitors and ECG alarms) may not operate properly when electrical stimulation is in use.
- TENS is a symptomatic treatment, and as such, suppresses the sensation of pain which would otherwise serve as a protective mechanism.
- Inspect the plastic lens of the laser head for blemishes, deformation, pitting, scratches, discoloration, and cleanliness before each use.
- Do not drop the applicator or unit on hard surfaces or submerge in water. These actions will damage the applicator and unit. Damage resulting from these conditions is not covered under the warranty.
- Use of controls or adjustments or performance of procedures other than those specified herein may result in hazardous exposure to Laser energy.
- This device should be kept out of the reach of children.
- Use of other accessories other than those specified in this User Manual may increase electrical emissions and decrease electrical immunity of the device.
- Contaminated sponges, electrodes, lead wires, and gel can lead to infection.
- Use of electrode with degraded hydrogel can result in burn to the skin.
- DO NOT operate this unit in an environment where other devices are being used that intentionally radiate electromagnetic energy in an unshielded manner.
- Use of electrode on multiple patients can lead to infection.
- Clean applicators after each use, otherwise it can lead to cross contamination and infection.
- When the Laser Module is not in use, it should be protected against unqualified use.

**WARNING**

- Do not treat through clothing.
- Stop treatment immediately if patient experiences discomfort or pain.
- Do not apply laser on an area of skin that has lotion or ointments applied as burns may occur.
- Do not use laser on or over a tattoo.
- The laser head must be cleaned with a disinfectant cleaner (i.e. Virex® II 256) or germicidal cloth (i.e. PDI Sani-Cloth® Plus/Hb) between each therapy session. Ensure no liquids enter into the laser head while cleaning. Do not use any chlorine-based cleaners on the laser head.
- The color of skin, age of lesion, depth of lesion, sensitivity of the patient, tissue type, and medications that increase sensitivity to light may affect therapy.
- Powered muscle stimulators should be used only with the leads and electrodes recommended for use by the manufacturer.
- In the event of all 300-Level or a 200-Level error message that cannot be resolved, immediately stop all use of the system, and contact the dealer or DJO for service. Errors and Warnings in these categories indicate an internal problem with the system that must be tested by DJO or a Trained Technician before any further operation or use of the system.
 - Use of a system that indicates an Error or Warning in these categories may pose a risk of injury to the patient, user, or extensive internal damage to the system.
- Use of controls or adjustments or performance of procedures other than those specified herein may result in hazardous exposure to ultrasonic energy.
- Before administering any treatment to a patient you should become acquainted with the operating procedures for each mode of treatment available, as well as the indications, contraindications, warnings, and precautions. Consult other resources for additional information regarding the application of each mode of treatment.
- Disconnect the system from the power source before attempting any maintenance, installation, removal, or replacement procedures to prevent electrical shock and possible damage to system.
- Keep electrodes separated during treatment. Electrodes in contact with each other could result in improper stimulation or skin burns.

**WARNING**

- The Vectra® Neo Clinical Therapy System may be susceptible to Electro-Static Discharge (ESD) at greater than ± 4 kV when first grasping either the Ultrasound or Laser Applicator. In the event of such a discharge, the Vectra® Neo Clinical Therapy System may experience communication loss with the installed modules. The Vectra® Neo Clinical Therapy System will terminate all active outputs (stim, ultrasound, laser), automatically place the unit in a safe state, and issue an error message 301 or 307.
 - To recover from an error message 301 or 307, turn the unit off and on using the ON/OFF switch located at the top of the display. Once the system restarts, re-initiate all treatments that were interrupted.
- To prevent Electro-Static Discharge (ESD) at greater than ± 4 kV:
 - Grasp and hold the Ultrasound or Laser Applicator prior to starting treatment. If the applicator must be put down prior to completion of treatment, stop the current treatment first and then place the applicator in the holder.
 - Maintain humidity in the use environment to at least 50% relative humidity.
 - Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, DJO recommends implementing additional controls to maintain relative humidity to at least 50%.
 - Communicate these ESD-precautionary procedures to healthcare staff, contractors, visitors, and patients.
- Long term effects of chronic electrical stimulation are unknown.
- Stimulation should not be applied over the anterior neck or mouth. Severe spasm of the laryngeal and pharyngeal muscles may occur and the contractions may be strong enough to close the airway or cause difficulty in breathing.
- Stimulation should not be applied transthoracically in that the introduction of electrical current into the heart may cause cardiac arrhythmia.
- Stimulation should not be applied over swollen, infected, and inflamed areas or skin eruptions, e.g., phlebitis, thrombophlebitis, varicose veins, etc.
- Stimulation should not be applied over, or in proximity to, cancerous lesions.
- Electrotherapy output current density is related to electrode size. Improper application may result in patient injury. If any question arises as to the proper electrode size, consult a licensed practitioner prior to therapy session.
- The Vectra® Neo Clinical Therapy System optional modules and associated accessories are designed for use only with the Vectra® Neo Clinical Therapy System.
- Remove the Ultrasound or Laser Applicator by pulling the cable connector only. DO NOT remove by pulling the cable.

**WARNING**

- Output current density is related to electrode size. Improper application may result in patient injury. If any question arises as to the proper electrode size, consult a licensed practitioner prior to therapy session.
- Do not apply the Ultrasound Applicator to the patient during the Head Warming period. Applicator must remain in Applicator Hook during the Head Warming period.
- Some patients are more sensitive to Laser output (i.e., patients taking medications that increase sensitivity to light) and may experience a reaction similar to a heat rash.
- Before each Laser use, clean the plastic lens with a clean cloth. Make certain to apply with a clean cloth. Failure to clean the lens between patient therapy sessions could cause beam fragmentation, which may reduce the effectiveness of the treatment.
- Medical electrical equipment needs special precautions regarding EMC. Portable and mobile RF communication equipment can be affected by other medical electrical devices. If you believe interference is occurring, please consult the ELECTROMAGNETIC COMPATIBILITY (EMC) section on page 73, to assist in removing the interference.
- Common RF emitting devices (e.g., RFID) and electromagnetic security systems (e.g., metal detectors) may interfere with the operation of the Vectra® Neo Clinical Therapy System. The Vectra® Neo Clinical Therapy System has been tested in the presence of these types of devices and while no adverse event occurred, the device should not be operated within the vicinity or environment as another RF emitting device.

**DANGER**

- Stimulus delivered by the TENS waveforms of this device, in certain configurations, will deliver a charge of 25 microcoulombs (μC) or greater per pulse and may be sufficient to cause electrocution. Electrical current of this magnitude must not flow through the thorax because it may cause a cardiac arrhythmia.
- Patients with an implanted neurostimulation device must not be treated with or be in close proximity to any shortwave diathermy, therapeutic ultrasound diathermy, or laser diathermy anywhere on their body. Energy from diathermy (shortwave, microwave, ultrasound, and laser) can be transferred through the implanted neurostimulation system, can cause tissue damage, and can result in severe injury or death. Injury, damage, or death can occur during diathermy therapy even if the implanted neurostimulation system is turned off.
- Handle, clean, and dispose of components and accessories that have come in contact with bodily fluids according to National, Local, and Facility rules, regulations, and procedures.



- This unit is considered to be a Class 3B Laser product and thus emits visible and invisible Laser radiation (IR). Avoid direct eye exposure to the Laser beam. The symbol to the left is located on the back of the applicator and indicates the active radiant surface (the area on the applicator that emits infrared Laser energy and the direction of the beam of light). When the unit is on, not all wavelengths are visible to the naked eye. Therefore, when performing any operational or functional check, always wear Chattanooga laser protective eyewear.
- The solvents of adhesives and flammable solutions used for cleaning and disinfecting should be allowed to evaporate before the unit is used.
- DO NOT connect the unit to an electrical supply without first verifying that the power supply is the correct voltage. Incorrect voltage may cause unit damage, malfunction, electrical shock, fire, or personal injury. Your unit was constructed to operate only on the electrical voltage specified on the Voltage Rating and Serial Number Plate. Contact your DJO dealer if the unit is not properly rated.



- Laser protective eyewear should be worn during laser treatment by the operator and patient to block infrared light energy from the eyes during treatment.
- DO NOT point the Laser beam directly into human or animal eyes. The lens of the eye does not detect the invisible, coherent Laser beams, potentially resulting in permanent retinal damage.



- Device is not designed to be used in oxygen rich environment. Explosion hazard if the device is used in the presence of flammable anesthetic mixture with air, oxygen, or nitrous oxide.

COMPONENTS

Throughout these instructions the terms “left” and “right” referring to the machine sides are from the perspective of a user standing in front of the unit.

The Vectra® Neo Clinical Therapy System allows installation of optional modality modules by the user. Specifically designed for use with the Vectra® Neo Clinical Therapy System, these modules configure the system to meet virtually every therapeutic need that a clinician may have. The components of the Vectra® Neo Clinical Therapy System are shown below.

NOTE: The Vectra® Neo Clinical Therapy System, when ordered as a Tabletop System, without cart, is assembled with Base (shown below). The only assembly required is the installation of the desired Modules described in the MODULE INSTALLATION section on page 23.

Head



Cart



Modules

- Stimulation Channel 1/2
- Stimulation Channel 1/2 + sEMG
- Stimulation Channel 3/4
- Laser
- Ultrasound



Lead Wires

The available lead wires are shown below. If the user orders Stimulation Channel 1/2 module, the box will include the blue and green lead wires. Stimulation Channel 3/4 is the cranberry and orange lead wires. If both modules are ordered, the box contains all four colored lead wires. Stimulation modules channel 1/2 with sEMG includes blue and green sEMG lead wires.



Lead Wire Holders



Powercord



MODULE SLOTS

- 1. Laser
- 2. Stimulation (1 & 2) / Stimulation (1 & 2) + sEMG
- 3. Ultrasound
- 4. Stimulation (3 & 4) – opposite side

Side Module Slots – Left Side



Side Module Slots – Right Side



To remove module, take right side off face plate and push module from right side.

MODULE KIT CONTENTS

Electrotherapy Module Channels 1/2 – PN 70000

- Stimulation module
- Lead wires
- DURA-STICK® 2 in (5 cm) Round Disposable Electrodes (1 pack of 4)
- Faceplates (to cover module after inserted into main unit)

Ultrasound Module – PN 70002

- Ultrasound module
- Faceplates (to cover module after inserted into main unit)

Electrotherapy Module Channels 3/4 – PN 70003

- Stimulation module
- Lead wires
- DURA-STICK® 2 in (5 cm) Round Disposable Electrodes (1 pack of 4)
- Faceplates (to cover module after inserted into main unit)

Electrotherapy Module Channels 1/2 + sEMG – PN 70004

- Stimulation module (2 channel Stimulation with sEMG)
- sEMG Lead Wires
- DURA-STICK® 2 in (5 cm) Round Disposable Electrodes (2 packs of 4)
- Faceplates (to cover module after inserted into main unit)

Laser Module – PN 70005

- Laser module
- Protective Eyewear, 2 Pair
- Interlock
- Patient Remote/Laser Interrupt Switch
- Faceplates (to cover module after inserted into main unit)

ULTRASOUND APPLICATOR

1. Applicator Head

The component of the applicator that makes contact with the patient during Ultrasound or Combination therapy.

2. Applicator

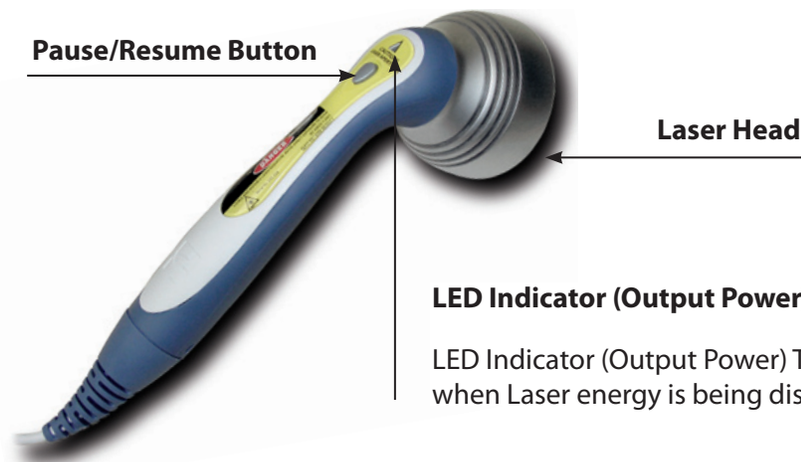
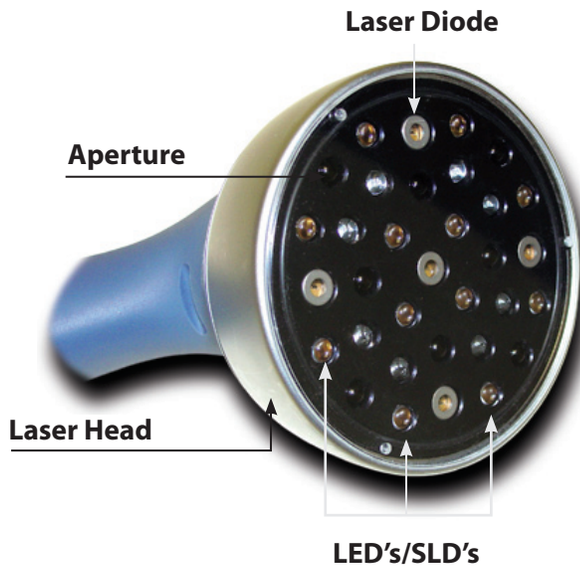
The assembly that connects to the system and incorporates the Applicator.

3. LED

The component of the applicator that indicates if the Applicator is coupled or uncoupled on the treatment area. Coupling is not available on the 1 cm² applicator.



LASER APPLICATOR



LED Indicator (Output Power) This orange light illuminates when Laser energy is being distributed by the applicator.

PATIENT REMOTE/LASER INTERRUPT SWITCH

The Vectra® Neo Patient Remote/Laser Interrupt Switch buttons are described in the table below. By default, the remote is not assigned to any treatment. When assigned, the buttons function as follows:

| | |
|-------------|---------------------------------|
| ▲ | Increase Intensity (1) |
| ▼ | Decrease Intensity (2) |
| STOP | STOP/Pause Treatment (3) |
| M | Manual Stimulation (4) |

Increase Intensity (1) (Electrical Stimulation Treatments Only) - Increases the intensity of the assigned stim treatment; button is not active if stim treatment is unassigned. Button is not active and has no function for Ultrasound or Laser treatments.

Decrease Intensity (2) (Electrical Stimulation Treatments Only) - Decreases the intensity of the assigned stim treatment; button is not active if stim treatment is unassigned. Button is not active and has no function for Ultrasound or Laser treatments.

STOP/Pause Treatment (3) (All Treatments) - Pauses treatment

M (Manual Stimulation) (4) (Electrical Stimulation Treatments Only) - Provides one cycle of stimulation. Can only be operated when the clinician enables manual mode on the base unit (head). This mode is clinician monitored and is not for use when the patient is unattended. Button is not active and has no function for Ultrasound or Laser treatments.



HEAD TO CART ASSEMBLY

The optional Therapy System Cart, PN 70001, is designed for use with the Vectra® Neo Clinical Therapy System only and allows the user to easily transport the System from patient to patient within the clinic as well as store all necessary accessories, supplies, and applicators used for the various modalities of the System.

Tools required (not included): #2 Phillips screwdriver and standard slotted screwdriver.

Remove the Vectra® Neo Clinical Therapy System from the shipping carton. Visually inspect for damage. Report any damage to the carrier immediately.

To assemble the Neo Head to the Cart, follow these steps:

1. Remove the top drawer from the Cart. Pull the drawer open. Press the plastic tabs on both drawer slides simultaneously in opposite directions, as shown. Pull the drawer completely out.



2. Remove the base from the head unit prior to placing it on the cart. Do this by removing the four screws from the underside of the base where they secure to the Neo Head. Retain for use when attaching the Neo Head to the Cart.



3. Place the Neo Head on the cart facing toward the drawers.



4. Fasten the Neo Head to the cart using four screws to connect the base to the Neo Head.



5. If desired, replace the closed Handles with the open Handles. Each Handle is attached with four screws as shown.



6. Reinstall the drawer.

NEO LEG TO CART ASSEMBLY/ADJUSTMENT

The Neo Cart is shipped without the legs attached.

Tools Required:

- 3/16" Hex Key Wrench (provided)
- Flat Washer 1/4" Internal diameter, quantity 6 (provided)
- Socket Head Cap Screw 1/4-20 x 1-1/4", quantity 6 (provided)

To install or adjust the leg assemblies onto the Neo Cart, follow these steps:

1. Remove the bottom drawer from the Cart. Pull the drawer open. Press the plastic tabs on both drawer slides simultaneously in opposite directions, as shown. Completely pull the drawer out.



2. There are two Cart height adjustments. Standard shown on the left and lowered, shown on the right. For initial installation, determine the desired height. Locate three Allen-style bolts for each leg, left and right and insert, by hand, in their respective slots. Use the Allen wrench to secure the legs.

NOTE: To adjust height at a later time, simply remove the Allen-style bolts, re-position the legs, and re-insert the bolts.



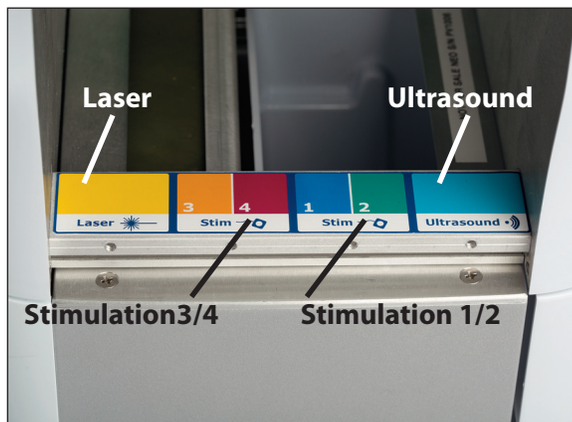
3. Reinstall the Bottom Drawer.

MODULE INSTALLATION

All modules are installed from the left side (when facing the screen) of the Neo Head unit and are each installed in the same manner. Each has color-coded lead wires that correspond to the appropriate colored labeling on the modules. Module-specific installation instructions are shown after the generic instructions. To install the modules in the Vectra® Neo Clinical Therapy System, follow the steps shown.

Tools required (not included): #2 Phillips screwdriver and standard slotted screwdriver.

The System is programmed to automatically recognize the new Module(s), therefore, no software installation is required.



1. Ensure that the power cord is removed from the device.
2. Remove the blank faceplate over the slot from the left and right sides of the Neo head. (The example displays the Ultrasound module.)



3. Insert a standard slotted screwdriver (not provided) into the top slot, pressing down with slight pressure. Pull the faceplate off (in this example showing Electrotherapy Channel 3/4).



4. The module is inserted on the left side of the Neo Head in the slot as shown in this example (with the Ultrasound module).



5. Carefully insert the module into the slot, with 32 pins (2x16) in first. Secure the module in place with gentle pressure until you feel the module is seated.



MODULE INSTALLATION (CONTINUED)

- Secure the module with a screw provided at the bottom as shown (using channel 3/4 as example).



- In this example showing the Laser faceplate, insert the faceplate at the bottom and snap into place at the top, as shown on the left and right sides (Vectra® Neo Clinical Therapy System allows Laser access on the left and right sides.)



- Plug in the unit and press the power button. Allow the unit to initialize, and then verify that the newly installed module is shown as available on the Home screen.

MODULE-SPECIFIC INFORMATION

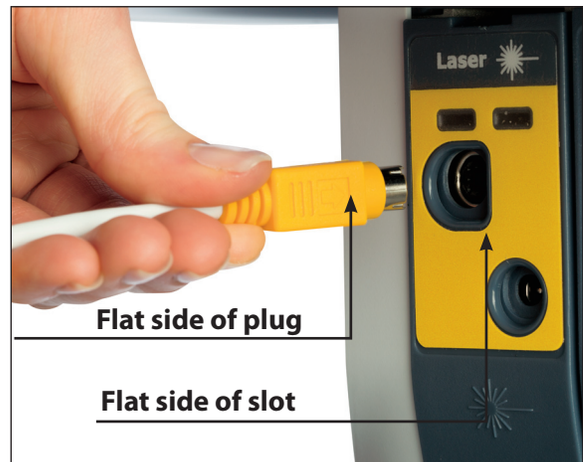
Ultrasound Cable Insertion

Shown below is the Ultrasound Cable Insertion location.



INSERTING PLUGS

When inserting the plugs for the Ultrasound and Laser modules, be sure to align the flat side of the plug with the flat side of the slot and push in gently. This is to avoid bending the pins in the plug.



PATIENT REMOTE/LASER INTERRUPT SWITCH INSTALLATION

To operate the Patient Remote/Laser Interrupt Switch, plug the remote into the device on the Rear Access Panel receptacle, as shown below:



2. Press the Remote ON/OFF toggle icon to assign or unassign the remote to the selected treatment. The remote can be assigned to only one treatment at a time, however the remote can be reassigned as needed.

When not in use, the Patient Remote/Laser Interrupt Switch can be stored by hooking it onto the lead wire holder clips in the same manner as lead wires and cables (shown below).



⚠ CAUTION

- Patient Remote/Laser Interrupt Switch is to be used under supervision of a physician or licensed practitioner only.

Complete the following steps to assign the remote to a treatment:

1. When the remote is plugged into the unit, a Remote ON/OFF toggle icon is displayed on the Treatment review screen in the upper right corner (shown below).



INSTALLING THE LASER INTERLOCK (DOOR INTERRUPT SWITCH)

The Laser Interlock is an optional safety device designed to interrupt Laser therapy if the door to the therapy room is opened. The Laser Interlock Kit consists of a switch resistor and a jack. Customers must supply the necessary cable that complies with local and international codes. Use only qualified electricians to install the Laser Interlock Kit.

The diagrams to the right provide installation guidelines for therapy room with single and multiple doors.

Operation of the Laser Interlock

Laser Interlock works as an interrupt switch once it is installed and connected to the Vectra® Neo Clinical Therapy System with Laser module.

Laser Interlock monitors the state of the door(s) of the therapy room and only allows start of Laser treatment if all of the doors are closed.

If any door is open, it will not allow user to start the Laser treatment; and if treatment is already started and someone opens the door, it interrupts the system to stop the Laser treatment.



WARNING

- **Disconnect the system from the power source before attempting any maintenance, installation, removal, or replacement procedures to prevent electrical shock and possible damage to system.**
- **The Laser Interlock must be installed by a professional or qualified electrician. Serious eye injury can result if the device is not properly installed. Also, when installing the device for multiple doors, the resistance total may not exceed 4800 ohm.**

Diagram for Therapy Room with One Door

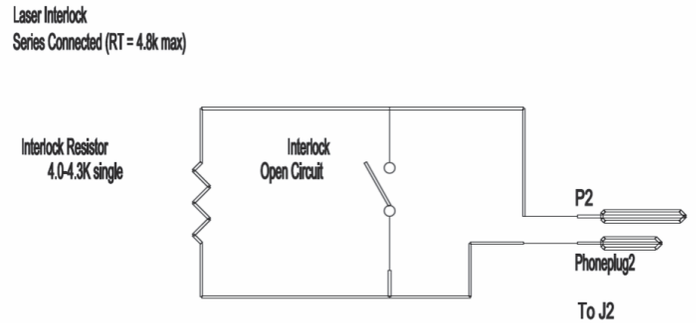
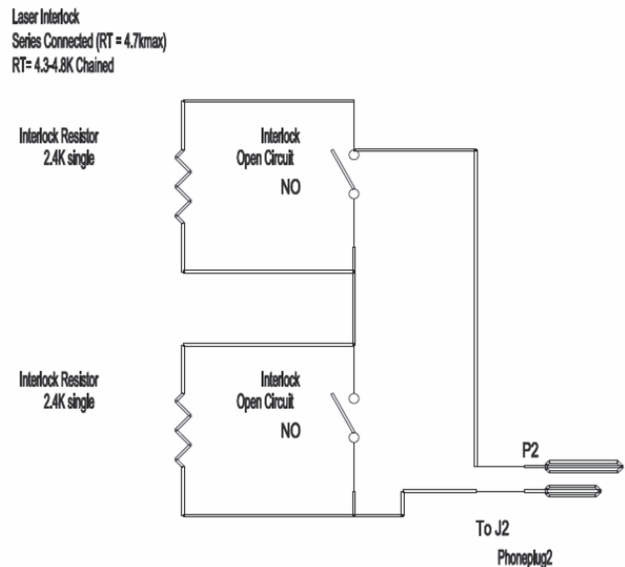


Diagram for Therapy Room with Multiple Doors



THERAPY SYSTEM START-UP

Complete the following steps for initial setup of the Vectra® Neo Clinical Therapy System:

1. Plug the Power cord into the back of device. Plug the other end of the cord into an electrical outlet.

NOTE: The Power Cord may be unplugged from the back of the cart in an emergency situation.



2. Press the Power button located on the top left portion of the LCD casing (shown below):



3. Select desired function on the Home screen (shown below).



SYSTEM SPECIFICATIONS AND DIMENSIONS

| | Width | Depth | Height | Weight |
|--|---------------------|---------------------|---------------------|----------------------|
| Module | 11.12" (28.2448 cm) | 6.34" (16.1036 cm) | 1.43" (3.6322 cm) | 1lb (0.453592 kg) |
| Head @ 45 degree with Base (Tabletop) | 15.89" (40.3606 cm) | 15.89" (40.3606 cm) | 22.05" (56.007 cm) | 20.7lb (9.389362 kg) |
| Cart Lowered (with casters) | 23.94" (60.8076 cm) | 26.19" (66.5226 cm) | 27.41" (69.6214 cm) | 29.4lb (13.33562 kg) |
| Cart Raised (with casters) | 23.94" (60.8076 cm) | 26.19" (66.5226 cm) | 30.15" (76.581 cm) | |
| Head and raised cart with screen @ 90deg | 23.94" (60.8076 cm) | 26.19" (66.5226 cm) | 52.85" (134.239 cm) | 48.9lb (22.18067 kg) |

POWER (COMBINATION AND ELECTROTHERAPY UNITS)

Input 100 - 240 V AC, 2.5A to 1.25A, 50/60 Hz
 Electrical Class CLASS I
 Mode of Operation Continuous

Electrical Type (Degree of Protection)

- UltrasoundTYPE B 
- LaserTYPE B 
- ElectrotherapyTYPE BF 
- Electrotherapy & sEMGTYPE BF 
- Ultrasound & ElectrotherapyTYPE B 

NOTE: All waveforms except High Voltage Pulsed Current (HVPC) have been designed with a 200 mA current limit. VMS™, VMS™ Burst, and all TENS waveform output intensities are measured, specified, and listed to peak, not peak to peak.

GENERAL SYSTEM OPERATING AND STORAGE TEMPERATURE

Operating Conditions

The device will meet its requirement under the following conditions:
 Temperature: 10° C to 45° C
 Relative Humidity: 0% to 90%
 Atmospheric Pressure: 700hPa to 1060hPa

Transport and Storage Conditions

The device will remain in proper condition under the following conditions:
 Temperature: Above 0° C freezing to +60°C
 Relative Humidity: max 95%
 Atmospheric Pressure: 700hPa to 1060hPa

ULTRASOUND SPECIFICATIONS

Frequency..... 1 MHz, ±5%; 3.3 Mhz, ±5%
 Duty Cycles 10%, 20%, 50%, Continuous
 Pulse Repetition Rate 100 Hz ±20%
 Pulse Duration 1mSec, ±20%; 2mSec, ±20%; 5mSec, ±20%

Output Power

10 cm² Crystal..... 0-15 W at 1 MHz, 0-10 W at 3.3 MHz
 5 cm² Crystal..... 0-6W @ 1 and 3.3 MHz
 2 cm² Crystal..... 0-3 W @ 1 and 3.3 MHz
 1 cm² Crystal..... 0-1.5 W @ 3.3 MHz

Amplitude.....0-2.5 W/cm² in continuous and pulsed modes
 Output accuracy..... ±20%, 10% of maximum
 Temporal Peak to Average Ratio:..... 2:1, ±20%, at 50% Duty Cycle
 5:1, ±20%, at 20% Duty Cycle
 9:1, ±20%, at 10% Duty Cycle
 Beam Nonuniformity Ratio..... 5:1 maximum
 Beam Type..... Collimating
 IPXX Rating for Unit..... IPX0
 IPXX Rating for Applicator..... IPX7
 Treatment Time 1 to 30 min

| Effective Radiating Areas (ERA) | | | | | |
|---------------------------------|------------------------|-----------------|------|-----------------|------|
| Description | ERA (cm ²) | ERA High | | ERA Low | |
| | | cm ² | % | cm ² | % |
| 10 cm ² Crystal | 8.5 | 10 | +18% | 7 | -18% |
| 5 cm ² Crystal | 4 | 5 | +25% | 3 | -25% |
| 2 cm ² Crystal | 1.8 | 2 | +11% | 1.4 | -22% |
| 1 cm ² Crystal | 0.9 | 1 | +11% | 0.4 | -55% |

Head Warming Feature

The Head Warming feature of an Vectra® Neo Clinical Therapy System utilizes Ultrasound output, resulting in warming of the Applicator to increase patient comfort.

With Head Warming enabled, ultrasound is emitted without pressing the Start button while an ultrasound treatment is being set up. The Applicator LED will not illuminate during the Head Warming period. US Channel will indicate “Head Warming”.

Output..... 0-50% Cycling of maximum power
 Frequency..... 3.3 Mhz
 Applicator Temperature..... 29.4 °C - 43.3 °C (85 °F - 110 °F)

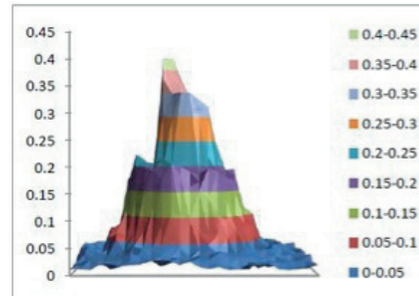
ULTRASOUND SPATIAL PATTERN

The following charts represent the distribution of the ultrasonic radiation field and the orientation of the field with respect to each applicator (Y-plane represents voltage in Vrms and X-plane represents applicator head surface in 1mm resolution).

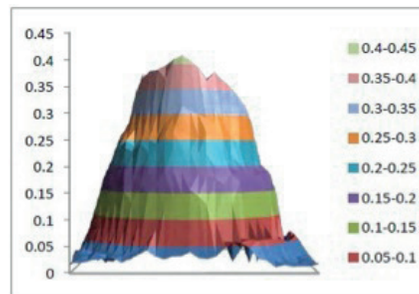
1 cm² Crystal (model 27333)



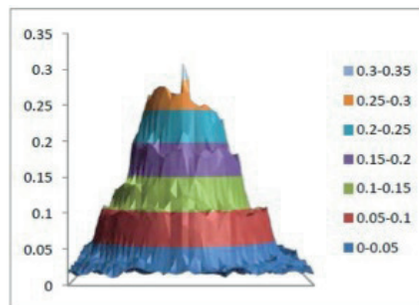
2 cm² Crystal (model 27334)



5 cm² Crystal (model 27335)



10cm² Crystal (model 27336)



LASER SPECIFICATIONS

Power

Output Type Infrared Lamp (Laser)
Laser Class..... 3B

Laser Technical Specifications

Pulse Frequencies..... 8 Hz - 10000 Hz and continuous
Wavelengths..... 670-950 nm (dependent on applicator)
Output..... 100-1440 mW (dependent on applicator)
Output accuracy..... $\pm 20\%$ of nominal

LASER APPLICATOR TECHNICAL SPECIFICATIONS

For all single diode and cluster laser and LED applicators, the expected increase in the measured quantities after manufacture added to the values measured at the time of manufacture is $\pm 20\%$.

The software incorporates a cooling function that forces the user to cool the laser cluster prior to the next treatment.

The software will calculate the cooling time needed when treatment times exceed 3 minutes per application. For a 3 minute treatment, it will force a 15 second cool down period before the next treatment can begin. For a 4 minute treatment, it will force a 2 minute cool down period before the next treatment can begin. The software extrapolates for times between 3 and 4 minutes.

A message will display on the screen informing the user that the probe is cooling down and the time period required. After 5 seconds, this message will disappear. If the user attempts to use the probe before the cool down period is completed, the message will re-display to signify that the applicator is still in cool down mode. After the cool down period is complete, a message displays that informs the user that the unit is ready for use.

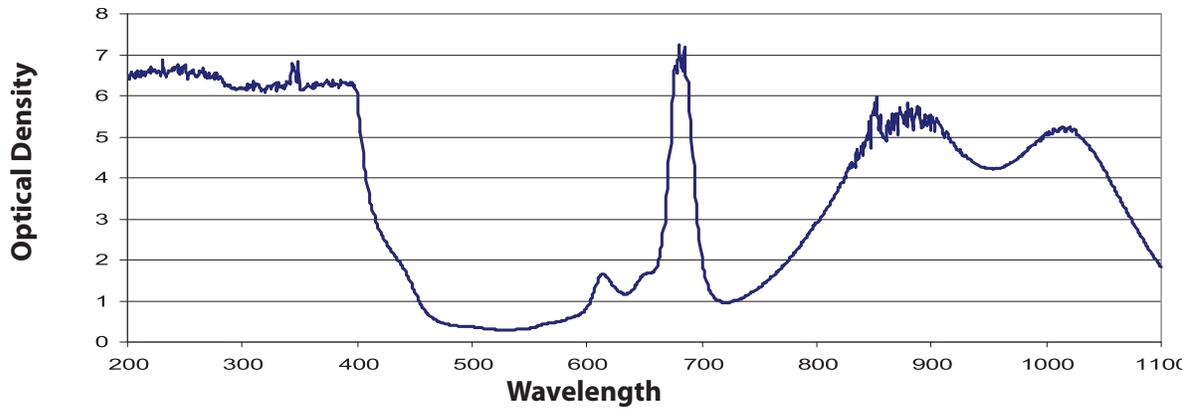
| Model # | Description | Diode Type | Wavelength(s) (nm) | Output Power (mW) | Power Density (W/cm ²) | Treatment Area (Spot Size) (cm ²) | Nominal Ocular Hazard Distance (NOHD-in meters) | Divergence a1 (rad) | Divergence a1 (rad) |
|---------|----------------------------|------------|--|-------------------|------------------------------------|---|---|---------------------|---------------------|
| 27799 | Single 670nm LED | LED | 670 | 10 | 0.012 | 0.785 | 0.386 | 0.698 | N/A |
| 27802 | 33-Diode cluster Laser/LED | GaAlAs LED | 670 nm (10mW) LED x12 880 nm (25mW) LED x8 950 nm (15mW) LED x8 850 nm (50mW) Laser x5 | 690 | 0.022 | 31.2 | 3.110 | 0.097 | 0.543 |
| 27803 | Single 850nm Laser | GaAlAs | 850 | 40 | 0.05 | 0.785 | 2.488 | 0.097 | 0.543 |
| 27804 | Single 850nm Laser | GaAlAs | 850 | 150 | 0.191 | 0.785 | 8.800 | 0.097 | 0.543 |
| 27805 | Single 820nm Laser | GaAlAs | 820 | 300 | 0.382 | 0.785 | 15.240 | 0.097 | 0.543 |
| 27807 | 33-Diode cluster Laser/LED | GaAlAs LED | 670 nm (10mW) LED x12 880 nm (25mW) LED x8 950 nm (15mW) LED x8 850 nm (100mW) Laser x5 | 940 | 0.03 | 31.2 | 6.221 | 0.097 | 0.543 |
| 27808 | 33-Diode cluster Laser/LED | GaAlAs LED | 670 nm (10mW) LED x12 880 nm (25mW) LED x8 950 nm (15mW) LED x8 850 nm (200mW) Laser x5 | 1440 | 0.045 | 31.2 | 12.443 | 0.097 | 0.543 |
| 27809 | 33-Diode cluster LED | LED | 670 nm (10mW) LED x12 880 nm (25mW) LED x13 950 nm (15mW) LED x8 | 565 | 0.018 | 31.2 | 0.386 | 0.698 | N/A |

LASER APPLICATOR TECHNICAL SPECIFICATIONS (CONTINUED)

| Model # | Description | Diode Type | Wavelength(s) (nm) | Output Power (mW) | Power Density (W/cm ²) | Treatment Area (Spot Size) (cm ²) | Nominal Ocular Hazard Distance (NOHD-in meters) | Divergence a1 (rad) | Divergence a1 (rad) |
|---------|----------------------------|------------|---|-------------------|------------------------------------|---|---|---------------------|---------------------|
| 27810 | 9-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x4 850 nm (50mW) Laser x5 | 290 | 0.038 | 7.55 | 3.110 | 0.097 | 0.543 |
| 27811 | 9-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x4 850 nm (100mW) Laser x5 | 540 | 0.071 | 7.55 | 6.221 | 0.097 | 0.543 |
| 27812 | 9-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x4 850 nm (200mW) Laser x5 | 1040 | 0.137 | 7.55 | 12.443 | 0.097 | 0.543 |
| 27813 | 13-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x7 950 nm (15mW) LED x3 850 nm (50mW) Laser x3 | 265 | 0.035 | 7.55 | 3.110 | 0.097 | 0.543 |
| 27814 | 13-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x7 950 nm (15mW) LED x3 850 nm (100mW) Laser x3 | 415 | 0.054 | 7.55 | 6.221 | 0.097 | 0.543 |
| 27815 | 19-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x6 880 nm (25mW) LED x7 950 nm (15mW) LED x6 | 325 | 0.043 | 7.55 | 0.386 | 0.698 | N/A |
| 27816 | 13-Diode cluster Laser/LED | GaAIAs LED | 670 nm (10mW) LED x7 950 nm (15mW) LED x3 850 nm (200mW) Laser x3 | 715 | 0.094 | 7.55 | 12.443 | 0.097 | 0.543 |
| 27840 | Single 850nm Laser | GaAIAs | 850 | 100 | 0.127 | 0.785 | 6.221 | 0.097 | 0.543 |
| 27841 | Single 85 nm Laser | GaAIAs | 850 | 200 | 0.254 | 0.785 | 12.440 | 0.097 | 0.543 |

LASER PROTECTIVE EYEWEAR SPECIFICATIONS

The graph below illustrates optical density in relation to wavelength. Each unit is shipped with laser protective eyewear that is L3 rated and approved, as well as EN207 compliant.



Useful Range

- Optical Density 5+..... 190-400 nm
- Optical Density 3+..... 625-830 nm
- Optical Density 3+..... 815-1050 nm

LASER LABELS

| | |
|---|--|
| <p>This serial decal label is affixed at the rear of the system.</p> | |
| <p>The device is labeled with the date of manufacture.</p> | |
| <p>The Laser Applicator handle label includes important safety information.</p> | |
| <p>The LED Applicator handle label includes important safety information.</p> | |

WAVEFORMS

For additional information on Waveforms and output energy, refer to APPENDIX 3 on page 78.

CC: Constant Current
CV: Constant Voltage



IFC (Interferential) Traditional (4 Pole)

Interferential Current is a medium frequency waveform. Current is distributed through two channels (four electrodes). The currents cross each other in the body at the area requiring treatment. The two currents interfere with each other at this crossing point, resulting in a modulation of the intensity (the current intensity increases and decreases at a regular frequency).

| | |
|--------------------------------|--|
| Output Mode | Electrodes |
| Carrier Frequency | 2000-10,000 Hz |
| Beat Frequency..... | 1-200 Hz |
| Sweep Time..... | 15 sec |
| Sweep Low Beat Frequency | 1-200 Hz |
| Sweep High Beat Frequency..... | 1-200 Hz |
| Scan Percentage..... | Static, 40%, 100%, Manual |
| Amplitude..... | 0-100 mA (CC with carrier freq ≤ 5000 kHz) |
| | 0-90 mA (CC with carrier freq > 5000 kHz) |
| | 0-64 V (CC with carrier freq ≤ 5000 kHz) |
| | 0-45 V (CC with carrier freq > 5000 kHz) |
| Treatment Time | 1-60 Minutes |
| Available on Channel..... | 1&2, 3&4 Option |
| Mode Selection | CC or CV |



TENS-Symmetrical Biphasic

The Symmetrical Biphasic waveform has a short pulse duration and is capable of strong stimulation of nerve fibers in the skin and in muscle. This waveform is often used in portable muscle stimulation units, and some TENS devices.

| | |
|----------------------------|------------------------------|
| Output Mode | Electrodes |
| Output Intensity..... | 0-80 mA (CC) 0-80 V (CV) |
| Phase Duration | 20-1,000 µsec |
| Frequency | 1-250 Hz |
| Mode Selection | CC or CV |
| Burst Frequency..... | 0-31 bps |
| Frequency Modulation | 0-250 Hz |
| Amplitude Modulation | Off, 40%, 60%, 80%, and 100% |
| Treatment Time | 1-60 min |
| Ramp..... | 0-5 sec |



TENS-Asymmetrical Biphasic

The Asymmetrical Biphasic waveform has a short pulse duration. It is capable of strong stimulation of the nerve fibers in the skin as well as of muscle tissue. This waveform is often used in TENS devices. Because of its short pulse, the patient typically tolerates the current well, even at relatively high intensities.

| | |
|----------------------------|------------------------------|
| Output Mode | Electrodes |
| Output Intensity..... | 0-110 mA (CC) 0-110 V (CV) |
| Phase Duration..... | 20-1,000 µsec |
| Frequency | 1-250 Hz |
| Mode Selection | CC or CV |
| Burst Frequency | 0-31 Hz |
| Frequency Modulation | 0-250 Hz |
| Amplitude Modulation | Off, 40%, 60%, 80%, and 100% |
| Treatment Time | 1-60 minutes |
| Ramp..... | 0-5 sec |



TENS - HAN

The HAN Waveform provides optimal parameters with a precisely controlled sequence of Dense-and-Disperse (DD) modes of stimulation where 2 Hz is alternating with 15 or 70 Hz, each lasting for 3 seconds.

| | |
|----------------------------|--|
| Output Mode | Electrodes |
| Output Intensity..... | 0-100 mA |
| Phase Duration | 180 µsec |
| Mode Selection | CC |
| Burst Frequency..... | 0-2 Hz |
| Frequency Modulation | 80 Hz |
| Cycle Time .. | Burst of 8 pulses at 80 Hz (at a frequency of 2 Hz) for 3 seconds to |
| | .80 Hz continuous (no burst) for 3 seconds, repeated |
| Treatment Time | 1-60 min |
| Available on Channels..... | 1, 2, 3, or 4 |

DANGER

- Stimulus delivered by the TENS waveforms of this device, in certain configurations, will deliver a charge of 25 microcoulombs (µC) or greater per pulse and may be sufficient to cause electrocution. Electrical current of this magnitude must not flow through the thorax because it may cause a cardiac arrhythmia.**

WAVEFORMS (CONTINUED)



VMS™ is a symmetrical biphasic waveform with a 100 µsec interphase interval. Because the pulse is relatively short, the waveform has a low skin load, making it suitable for applications requiring high intensities, such as in muscle strengthening protocols.

| | |
|-----------------------------|---|
| Output Mode | Electrodes |
| Output Intensity | 0-200 mA (CC) 0-200 V (CV) |
| Channel Mode | Single, Reciprocal, Co-Contract |
| Phase Duration | 20-1,000 µsec |
| Mode Selection | CC or CV |
| Anti-Fatigue | Off or On |
| Set Intensity | Individual Channel Intensity Setting in Reciprocal and Co-Contract modes |
| Cycle Time | Continuous or User Defined |
| Frequency | 1-200 pps |
| Ramp | 0-5 sec |
| Treatment Time | 1-60 min |
| Available on Channels | 1, 2, 3, or 4 |



The VMS™ FR version of the VMS™ waveform is a physiologically based channel interaction in which one channel stimulates the agonist and the other the antagonist of the muscle group that is being exercised. VMS is a symmetrical biphasic waveform with a 100 µsec interphase interval. Because the pulse is relatively short, the waveform has a low skin load, making it suitable for applications requiring high intensities, such as in muscle strengthening protocols.

| | |
|-----------------------------|---|
| Output Mode | Electrodes |
| Output Intensity | 0-150 mA (CC) 0-150 V (CV) |
| Burst Duration | 200 - 5000 ms |
| Phase Duration | 20-400 µsec |
| Mode Selection | CC or CV |
| Channel Intensity | Setting in Reciprocal and Co-Contract modes |
| Cycle Time | Continuous, 5/5, 4/12, 10/10, 10/20, 10/30, 10/50 |
| Frequency | 20-80 pps |
| Treatment Time | 1-60 min |
| Available on Channels | 1&2, 3&4 |



VMS™ Burst is a symmetrical biphasic waveform delivered in a burst format. Because the pulse is relatively short, the waveform has a low skin load, making it suitable for applications requiring high intensities, such as muscle strengthening protocols.

| | |
|-----------------------------|---|
| Output Mode | Electrodes |
| Output Intensity | 0-200 mA (CC) 0-200 V (CV) |
| Channel Mode | Single, Reciprocal, Co-Contract |
| Phase Duration | 20-700 µsec |
| Mode Selection | CC or CV |
| Anti-Fatigue | Off or On |
| Set Intensity | Individual Channel Intensity Setting in Reciprocal and Co-Contract modes |
| Cycle Time | Continuous or User Defined |
| Frequency | 1-200 bps |
| Ramp | 0-5 sec |
| Treatment Time | 1-60 min |
| Available on Channels | 1, 2, 3, or 4 |



Premodulated Current is a medium frequency waveform. Current comes out of one channel (two electrodes). The current intensity is modulated: it increases and decreases at a regular frequency (the Amplitude Modulation Frequency).

| | |
|---------------------------------|--|
| Output Mode | Electrodes |
| Output Intensity | 0-100 mA (CC) 0-96 V (CV) (with Carrier Frequency ≤ 5,000 kHz) 0-100 mA (CC) 0-68 V (CV) (with Carrier Frequency > 5,000 kHz) |
| Carrier Frequency | 2,000-10,000 Hz |
| Beat Fixed (Sweep Off) | 1-200 Hz |
| Sweep Low Beat Frequency | 1-200 Hz |
| Sweep High Beat Frequency | 2-200 Hz |
| Cycle Time | Continuous or User Defined |
| Mode Selection | CC or CV |
| Treatment Time | 1-60 Min |
| Available on Channel | 1, 2, 3, or 4 |

WAVEFORMS (CONTINUED)



DC (Direct Current)

DC Current is a direct current flowing in one direction only. The current can be continuous or interrupted.

| | |
|--|---|
| Output Mode | Electrodes |
| Output Intensity | 0-72 mA |
| Polarity Reversal | On or Off |
| With Polarity Reversal On, Polarity will change every 5 minutes. | |
| Cycle Time | Continuous, 5/5, 4/12, 10/10, 10/20, 10/30, 10/50 |
| Mode Selection | CC |
| Treatment Time | 1-60 min |
| Available on Channels | 1, 2, 3, or 4 |



Russian

Russian Current is a sinusoidal waveform, delivered in bursts or series of pulses. This method was claimed by its author (Kots) to produce maximal muscle strengthening effects without significant discomfort to the patient.

| | |
|--|---------------------------------|
| Output Mode | Electrodes |
| Output Intensity | 0-100 mA (CC) 0-90 V (CV) |
| Channel Mode | Single, Reciprocal, Co-Contract |
| Duty Cycle | 10%, 20%, 30%, 40%, 50% |
| Mode Selection | CC or CV |
| Anti-Fatigue | Off or On |
| Cycle Time | Continuous or User Defined |
| Carrier Frequency | 2,500 Hz |
| Burst Frequency (Anti-Fatigue Off) | 20-100 bps |
| Ramp | 0-5 sec |
| Treatment Time | 1-60 min |
| Available on Channels | 1, 2, 3, or 4 |



High Voltage Pulsed Current (HVPC)

The High Voltage Pulsed Current (HVPC) has a very brief pulse duration characterized by two distinct peaks delivered at high voltage. The waveform is monophasic (current flows in one direction only).

| | |
|-----------------------------|----------------------------|
| Output Mode | Electrodes |
| Output Intensity | 0-500 V |
| Polarity | Positive or Negative |
| Ramp | 0.5-5 sec |
| Display | Peak Current or Volts |
| Sweep High Frequency | 20-120 pps |
| Sweep Low Frequency | 10-110 pps |
| Frequency | 10-120 pps |
| Cycle Time | Continuous or User Defined |
| Treatment Time | 1-60 Min |
| Available on Channels | 1, 2, 3, or 4 |
| Mode Selection | CV |



Microcurrent

Microcurrent is a monophasic waveform of very low intensity. The physiological working mechanism of this effect is as yet not clearly understood. It is thought to stimulate tissue healing by stimulating the 'current of injury', a current which naturally occurs in healing tissue.

| | |
|-----------------------------|------------------------------------|
| Output Mode | Electrodes |
| Output Intensity | 0-1,000 µA |
| Polarity | Positive, Negative, or Alternating |
| Treatment Time | 1-60 Min |
| Available on channels | 1, 2, 3, or 4 |
| Duty Cycle | 50% |
| Frequency | 0.1-1,000 Hz |
| Mode Selection | CC |

ELECTRODE PLACEMENT

- Examine the skin for any wounds and clean the skin.
- Apply the electrodes to the treatment area.
- Ensure the electrodes are applied securely to the skin.
- Ensure good contact between each electrode and the skin.
- Check the electrode contact regularly during the treatment.
- Examine the skin again after the treatment.
- Choose electrodes that fit the anatomy.
- View the Electrode Placement recommendations in the Treatment Review screen for the particular modality being used for treatment as a reference point only prior to administering treatment.
- Follow electrode manufacturer instructions.
- DO NOT use electrodes with an active area less than 7.92 cm², as there will be a risk of suffering a burn injury. Always exercise caution with current densities more than 2mA/cm². Refer to CURRENT DENSITY CALCULATIONS FOR DJO ELECTRODES table in Appendix 3.

DURA-STICK® ELECTRODES

DURA-STICK® Electrodes are a self adhesive, disposable product designed specifically for use with Vectra® Neo Clinical Therapy System.

It is recommended that DURA-STICK® Electrodes be used whenever possible to ensure the highest level of contact with the treatment area and most uniform delivery of the prescribed electrotherapy treatment.



For ELECTROTHERAPY OPERATION, refer to page 48.

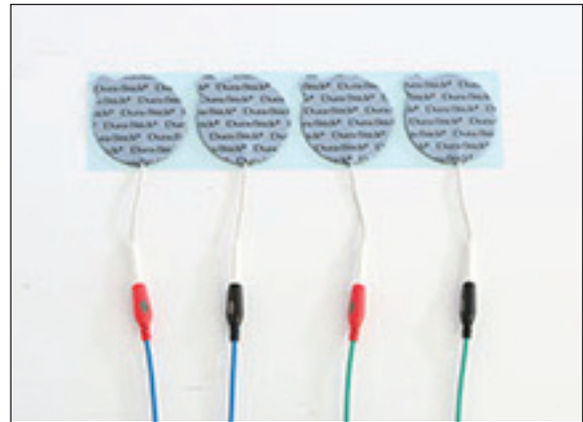
ELECTROTHERAPY PATIENT PREPARATION

DURA-STICK® Electrode Instructions

Connecting Lead Wires

1. Insert the lead with the Red (+) electrode connector into one DURA-STICK® Electrode.
2. Insert the lead with the Black (-) electrode connector into the other electrode.
3. Make certain the lead wires are seated completely into the electrodes.

NOTE: Use of conductive medium or sponges is not required or recommended. DURA-STICK® Electrodes are manufactured to ensure the optimum conductivity during therapy when properly applied.



Securing Electrodes

1. Remove the DURA-STICK® Electrodes from the protective backing.
2. Apply to the treatment area as prescribed.
3. Ensure the entire electrode surface is in contact with patient skin by pressing into place.



SEMG & STIM PATIENT PREPARATION

Install DURA-STICK® Electrodes

1. Connect a DURA-STICK® 2 in (5 cm) disposable electrode to each lead. These electrodes are designed for use with DJO equipment and will provide an accurate reading of sEMG activity. Active (Red) Lead, Active (Black) Lead, and Reference (Green) Lead are shown below.



2. Leave protective backing on electrodes until treatment area has been prepared.

Electrode Placement by Body Area

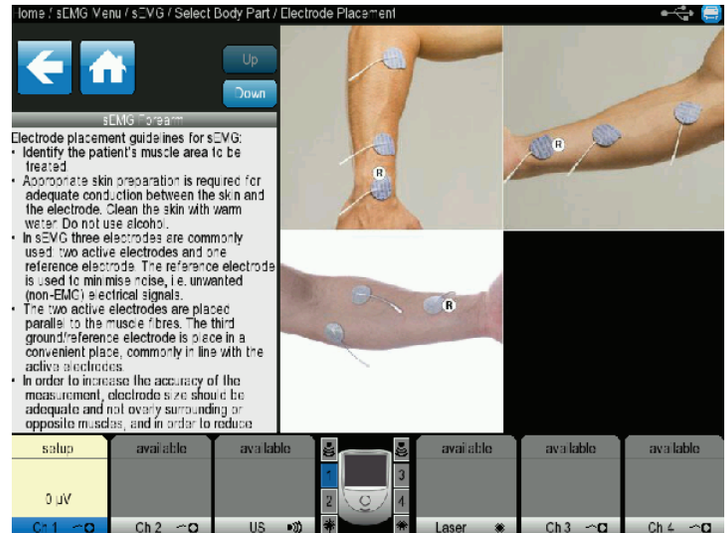
1. From the Home screen, select sEMG.
2. Choose one of the following modalities:
 - sEMG (Ch 1)
 - sEMG (Ch 2)
 - sEMG (Ch 1 + 2)
 - sEMG + Stim VMS
 - sEMG + Stim Sym Biph

Connect the sEMG Lead Wires

1. Connect the sEMG Lead wires to the channel(s) desired for use with the selected modality.



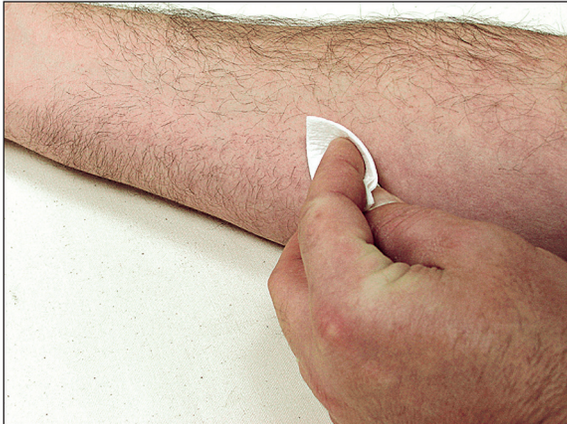
2. Select the Customize icon.
3. Select the Electrode Placement icon to view electrode placement according to body area.
4. Touch the body part you wish to treat.
5. View the electrode placement(s) for desired body area and press the Up and Down icons to scroll through text relating to specific electrode placement and typical conditions of the area.



6. Press the Back Arrow icon to return to desired screen.
7. Examine the skin for any wounds.

sEMG & STIM PATIENT PREPARATION (CONTINUED)

8. Thoroughly cleanse the skin treating area.

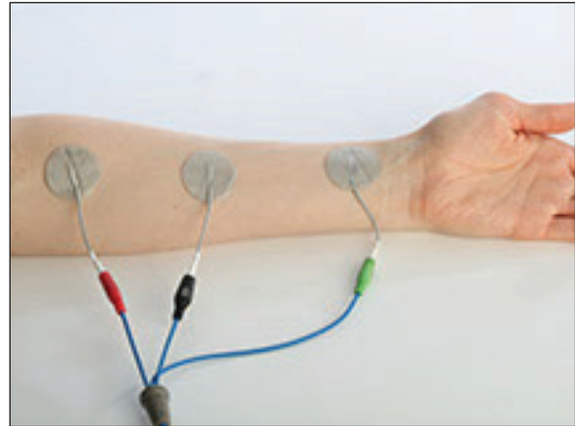


NOTE: Thorough and proper cleaning of the treatment area to remove any topical medication and cream film as well as loose skin particles from the treatment area is critical to the skin contact and reception of the Electrodes during sEMG and sEMG + Stim therapy.

Electrode Placement

1. Using DURA-STICK® 2 in (5 cm) electrodes, place the Active (red and black lead) electrodes in the center of the muscle belly and parallel with the muscle fibers.
2. Position the Reference electrode (green lead) in close proximity to the treatment area.
 - Using small electrodes and placing them closer together will render a more specific reading of muscle activity during sEMG and sEMG + Stim therapy.
 - The Active electrodes may be placed farther apart to obtain a general reading of a muscle or muscle group activity during the session.
 - DJO recommends using only DURA-STICK® electrodes to obtain the most accurate sEMG feedback.
 - Follow the electrode manufacturer instructions.
 - Trimming or cutting electrodes may interfere with the reception of sEMG data and may affect the delivery of electrical stimulation in the sEMG + Stim Modality.

3. Review the specific Electrode Placement graphic for positioning of the Reference (green lead) electrode.



NOTE: The electrodes may be placed for specific, general, and quasi-specific biofeedback muscle or muscle group activity.

For sEMG and sEMG + Electrical Stimulation operation, refer to page 55.

LASER PATIENT PREPARATION

Preparing the Patient's Skin for Laser Therapy

Before applying Laser therapy to the patient, you must first prepare the patient's skin. By properly preparing the patient's skin for Laser therapy, you will allow more Laser energy to reach the targeted areas and reduce the risk of skin irritation.

To prepare the patient's skin for Laser therapy, do the following:

1. Thoroughly wash the skin on which you intend to place the laser head with mild soap and water or alcohol wipe.
2. Dry the skin thoroughly.

For LASER OPERATION, refer to page 56.

ULTRASOUND PATIENT PREPARATION

1. Examine the skin for any wounds and clean the skin.
2. View the Applicator recommendation in the treatment.
3. Review screen for Ultrasound (as a reference point only) prior to administering treatment.

NOTE: Applicators are available in the sizes shown below:



Applicator Preparation and Use

1. Clean applicator before each therapy session with warm soapy water.
2. Liberally apply Conductor™ Transmission Gel to the treatment area on the patient.
 - Move the Applicator during therapy session in a circular motion. The area treated should be twice the diameter of the Applicator.
3. If US Coupling is On, the Applicator is properly coupled to the patient and administering ultrasound when the LED is constantly illuminated. Coupling is not available on the 1 cm² applicator.

For ULTRASOUND OPERATION, refer to page 51.

SCREEN DESCRIPTION

Each screen contains the following areas:

Title Bar

Located at the top of each screen and lists the current screen and previous screens back to the Home screen. It also contains a Print Screen icon at the top right, Patient Remote/Laser Interrupt Switch, when installed, and a USB connectivity icon when a USB flash drive is inserted.

Main Area

- Located under the Title bar, this area displays icons unique to the current screen.
- All screens (except the Home screen) will contain the Back Arrow icon to scroll to the previous screen and the Home icon to return to the Home screen.

Channel Area

Located at the bottom of each screen, this screen displays the following status information about each channel:

- **Not Installed:** Indicates the associated module is not installed in the unit
- **Available:** Indicates the channel is available for use

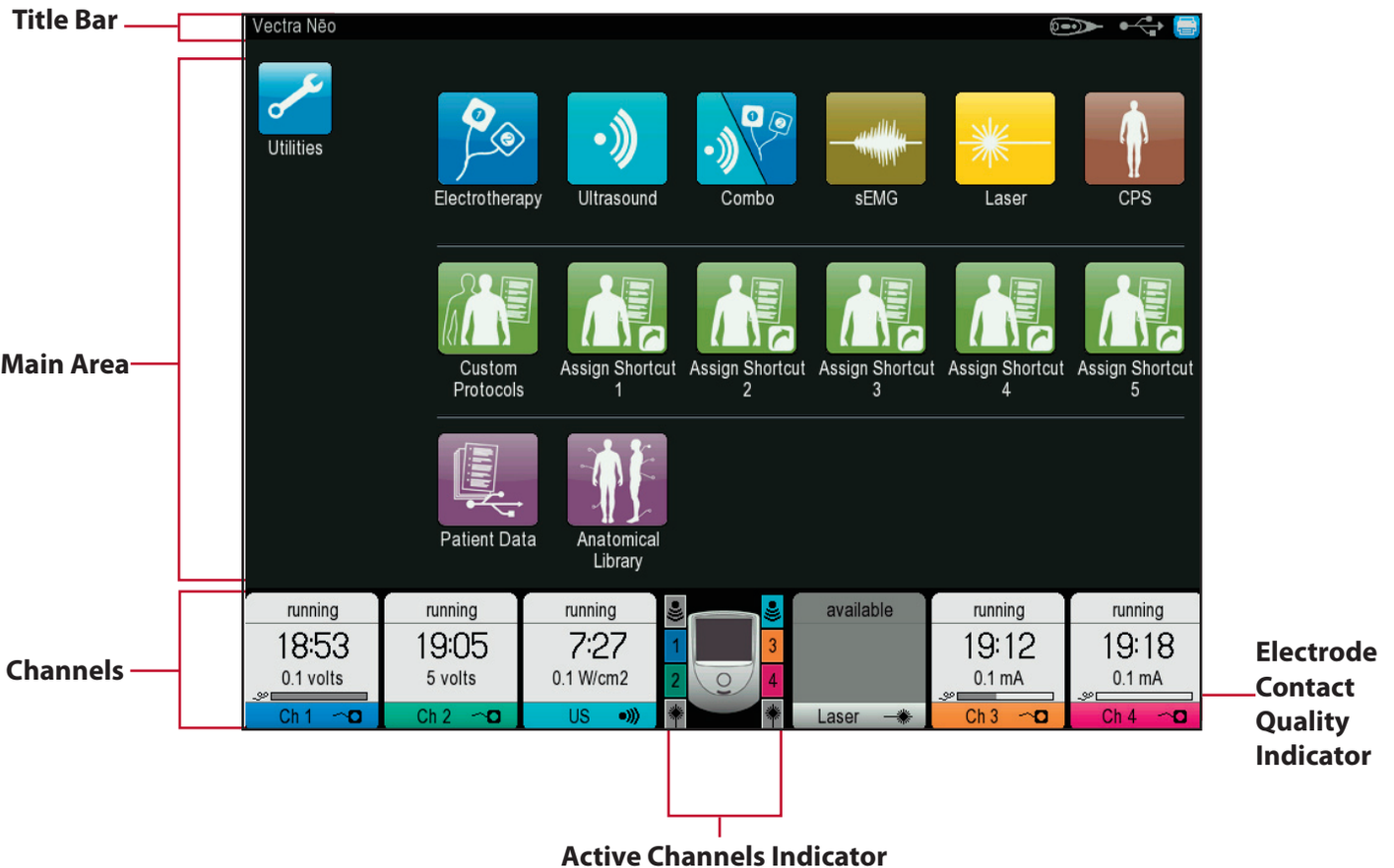
- **Setup:** Indicates a treatment for the channel is currently being set up but treatment has not yet begun
- **Running:** Indicates a treatment for the channel is currently running
- **Paused:** Indicates a treatment for the channel is currently paused
- **Completed:** Indicates a treatment for the channel has completed
- **No applicator:** Indicates there is not a valid applicator plugged into the channel's module (only valid for Ultrasound and Laser)

Electrode Contact Quality Indicator

(Electrotherapy Channels Only)

Active Channels Indicator

The image below shows the Home screen with modality and resource icons.



HOME SCREEN

The Vectra® Neo Clinical Therapy System Home screen provides access to all of the system modalities and functions. The Home screen has the following information:

1. Utilities

Modality Icons:

2. Electrotherapy
3. Ultrasound
4. Combo
5. sEMG
6. Laser
7. Clinical Protocol Set-up (CPS)
8. Custom Protocols
9. Assign Shortcut Options
10. Patient Data Storage
11. Anatomical Library



UTILITIES AND OPTIONS

The Utilities icon on the Home screen offers users the opportunity to set the following preferences::



1. **Print Screen**

Select the Printer icon in the top right corner to capture screen shots of current screen. Screen shots are saved to the USB flash drive plugged into the unit and are saved in Windows bitmap (.bmp) format. Note that a screen cannot be printed while a treatment is in process.

2. **Volume**

Select the Volume icon to set desired audio volume. The volume range is 0% (off) to 100% (loudest) and is measured in 20% increments. The default setting is 60%.

3. **LCD Brightness**

Select the Brightness icon to set the brightness of the LCD screen. The brightness ranges from 50% (dimmiest) to 100% (brightest) in 10% increments. The default setting is 100%.

4. **US Coupling**

Select the US Coupling icon to set the functionality of an ultrasound treatment when unit detects that the ultrasound applicator is uncoupled. Coupling is not available on the 1cm² applicator. The default is set to Off. The following preferences are available:

- **Off:** Nothing happens when applicator becomes uncoupled.
- **Pause and Beep:** When applicator becomes uncoupled, treatment timer will pause, unit will beep, and channel area indicates “uncoupled” state.
- **Pause and No Beep:** When applicator becomes uncoupled, treatment timer will pause, unit will not beep, and channel area indicates “uncoupled” state.
- **Beep:** When applicator becomes uncoupled, treatment timer will continue to run, unit will beep, and channel area indicates “uncoupled” state.
- **No Beep:** When applicator becomes uncoupled, treatment timer will continue to run, unit will not beep, and channel area indicates “uncoupled” state.

NOTE: Ultrasound output will continue to be emitted in all US Coupling modes even if the applicator is uncoupled.

UTILITIES AND OPTIONS (CONTINUED)

5. US Head Warming

Select the US Head Warming icon to turn on or off the ultrasound warming feature during ultrasound treatment setup. The available choices are either On or Off. The default setting is Off.

6. E-Stim Contact Quality

Select the E-Stim Contact Quality icon to turn on or off the electrotherapy contact quality monitoring function. The available choices are either On or Off. The default setting is On, and in the On position, the unit will monitor the contact quality of the electrodes on all waveforms, excluding High Voltage Pulsed Current (HVPC) and Microcurrent.

NOTE: If the impedance feedback exceeds 5000 ohms (tolerance $\pm 20\%$) and the set intensity is ≥ 10 mA CC or 10 V CV, then the unit will display an information message and set the intensity to zero for all channels associated with this treatment.

7. Restore Default Unit Settings

Select the Restore Default Unit Settings icon to reset all of the following settings back to their factory defaults:

- Volume
- LCD Brightness
- US Coupling
- US Head Warming

8. Restore Default Protocols

Select the Restore Default Protocols icon to reset all protocols (factory, custom, and the 5 custom protocol shortcuts) to their factory defaults.

9. Display Unit Version Information

Select the Display Unit Version Information icon to view the install status of each module. For all installed modules, the PCB information and the Software version for that module are displayed.

If a valid software upgrade USB flash drive is inserted into the unit, the version of software for each module that is on the USB flash drive is displayed.

NOTE: USB flash drive version is displayed in yellow if it is newer than the current software version for the applicable module.

10. Export Protocols to USB flash drive

Select the Export Protocols to USB flash drive icon to export all protocols (both default and custom) to a valid USB flash drive.

11. Import Protocols from USB Flash Drive

Select the Import Protocols from USB Flash Drive icon to import all protocols from a valid USB flash drive. Any protocols with the same name as an existing unit protocol will be overwritten by the imported protocol, and any protocols of a different name on the unit will remain intact.

12. Date and Time

Select the Date and Time icon to set the date and time on the unit.

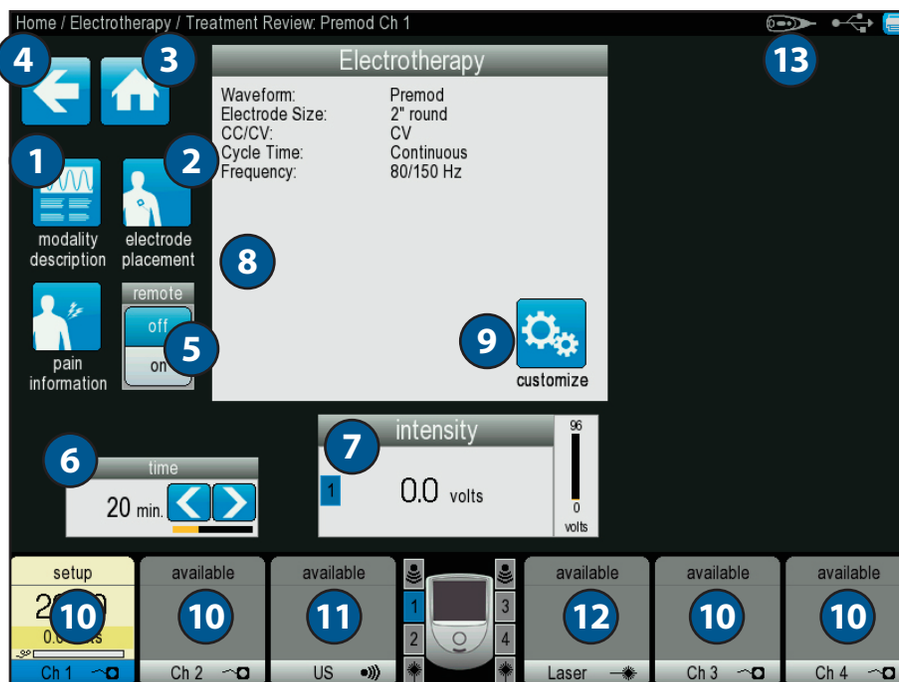
13. Clinic Name

Select the Clinic Name icon to enter the name of your clinic. The clinic displays on the Home screen and on the patient treatment reports saved to the USB flash drive. The following icons will perform the functions listed:

- **Left:** Moves the cursor to the left one space per instance pressed.
- **Right:** Moves the cursor to the right one space per instance pressed.
- **Space:** Inserts one space per instance pressed.
- **Clear:** Clears the entire entry.
- **Delete:** Once the cursor has been moved using the Left or Right icons, press the Delete icon to delete the character.
- **Caps:** Press the Caps icon to capitalize the letters or to make letters lowercase.
- **Cancel:** Press the Cancel icon to cancel the clinic name entry function and return to the Utilities screen.
- **Accept:** Press the Accept icon once entry of the clinic name is complete. The clinic name should now appear in the title bar area of the screen.

TREATMENT SCREENS

The Vectra® Neo Clinical Therapy System Treatment screens for Electrotherapy and Ultrasound include the following information:



1. Modality Description Icon

Press the Modality Description Icon to view the text explaining the rationale for the modality associated with the specific Clinical Protocol selected.

2. Electrode Placement Icon

Press the Electrode Placement Icon to view the specific electrode placement for the Clinical Protocol selected.

3. Home Icon

Press the Home Icon to return to the Home screen at any time.

4. Back Arrow Icon

Press the Back Arrow Icon to return to the previously viewed screen.

5. Remote Icon

Press either Off/On to deactivate or activate the Remote Control feature.

6. Time Icon

Press the Time icon to adjust therapy time/duration.

7. Intensity Icon

Set intensity by rotating the Intensity Control Knob to the prescribed level:

- Clockwise – Increases Intensity
- Counterclockwise – Decreases Intensity

8. Therapy Information Window

View selected Therapy information such as Waveform, Electrode Size, CC/CV, Carrier Frequency, Cycle Time, Frequency, and Vector Scan on the Therapy Information Window.

9. Customize Icon

Press the Customize icon to edit the Therapy information.

10. 4 Channel Icons

This icon shows the modalities in use. Channels are automatically assigned to the next available channel. Manual selection is done by touching the desired channel.

11. Ultrasound Icon

This icon shows the Ultrasound information when in use.

12. Laser Icon

This icon shows the Laser module information when in use.

13. Remote Icon

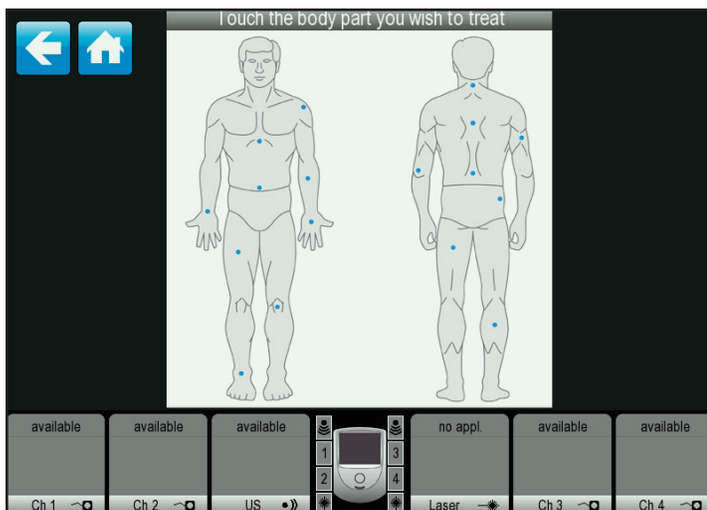
This is how the Remote Icon is viewed from the main screen.

CPS (CLINICAL PROTOCOL SETUP)

The Vectra® Neo Clinical Therapy System has a Clinical Protocol Setup (CPS) icon that is a series of protocol presets where the Body Area, Clinical Indication, Pathological Condition, and Pathological Severity are selected by the user to assist in selecting the parameter setting. These Clinical Protocols are to be used only as guidelines. Each patient should be individually assessed to determine the appropriateness of the protocol parameters prior to use.

The following information gives general instructions to access, select, and set up Clinical Protocols. Each Clinical Protocol is set up and edited in the same basic manner.

1. Select CPS from the Home screen.
2. Touch the body part you wish to treat.



3. Using the Up and Down Arrows, scroll to the appropriate indication and select by pressing the “√” icon.
4. Using the Up and Down Arrows, scroll to the appropriate condition you wish to treat and select by pressing the “√” icon.
5. Select the modality desired.
6. Begin therapy by pressing the Start button or make setting changes pressing the Customize icon.
7. To save the customized protocol, press the Save icon.
8. Save a new protocol by pressing the Save as a New Name icon and typing in a name for the protocol. Press the Save icon when finished.

9. Overwrite a previously saved protocol by using the Up and Down arrows or Previous Page/Next Page icons to select the protocol to overwrite. Press the Save icon to save the overwritten icon.
10. Return to Home screen by pressing the Home icon. Press the Back arrow to scroll back one screen.

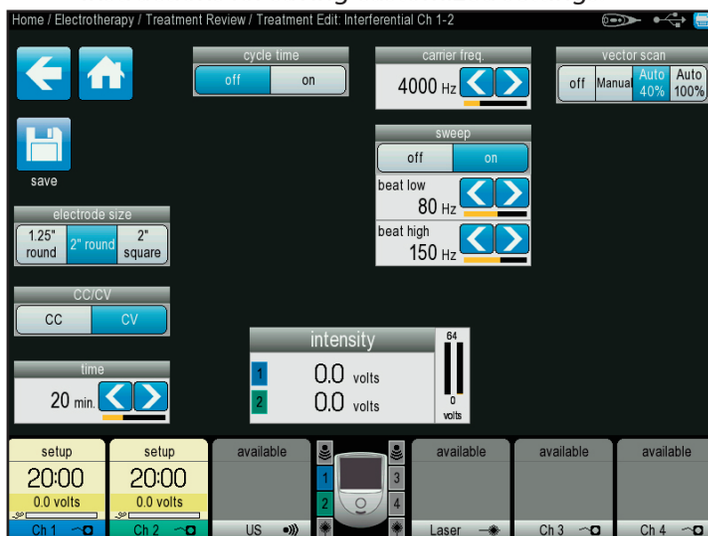
ELECTROTHERAPY OPERATION

All waveforms in the Vectra® Neo Clinical Therapy System are set up and edited in the same basic fashion. The Vectra® Neo Clinical Therapy System has an Electrotherapy Icon with the following waveforms: IFC (Interferential) Traditional (4 Pole), TENS- Asymmetrical Biphasic, TENS- Symmetrical Biphasic, TENS HAN, High Voltage Pulsed Current (HVPC), VMS™, VMS™Burst, VMS™ FR, IFC Premodulated (Traditional 2 Pole), Russian, and Microcurrent.

Complete the following steps to begin Electrotherapy treatment:

1. Prepare patient and therapy system for Electrotherapy. Refer to the ELECTROTHERAPY PATIENT PREPARATION section on page 38 for electrode selection, preparing the patient, and securing electrodes.
2. Touch the screen to activate the system. Read and carefully follow the instructions on the screen.
3. From the Home screen, press the Electrotherapy icon.
4. Select desired waveform from the listing on the screen by pressing the appropriate icon. Refer to the SYSTEM SPECIFICATIONS section on page 28 for all waveform specifications for the Vectra® Neo Clinical Therapy System. The screen shown below will appear.

- To view information explaining the waveform, press the Modality Description icon. Press the Up and Down icons to view additional text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
- To view the most commonly used electrode placement for the selected waveform, press the Electrode Placement icon. A picture will appear prompting you to select the body part you wish to treat. Press the Down icon to scroll through text. Press the Back icon to return to the previous screen or the Home icon to return to the Home screen.
- To customize waveform settings shown in the list box, press the Customize icon located in the list box, and the screen shown below will appear. Make the desired changes and press the Back icon to return to the previous screen, the Home icon to return to the home screen, or the Save icon to save the customized settings. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.



- To set pain information for the treatment, press the Pain Information icon. At this screen, you can do the following:
 - Map electrode placement by dragging the electrodes to their appropriate locations on the body and selecting either front, back, left, or right side.
 - Map pain information by touching the body area in the list box.
 - Edit pain scale information by pressing the Edit Pain Scale icon. Press the Up and Down arrows until pain is properly gauged. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.

ELECTROTHERAPY OPERATION (CONTINUED)

5. If desired, connect optional Remote Control to device.
6. Use the Intensity dial to set therapy intensity:
 - Clockwise - Increases Intensity
 - Counterclockwise - Decreases Intensity
7. Press the Start button to begin therapy, the Pause button to pause treatment, or the Stop button to terminate the treatment.

NOTE: To make setting adjustments during therapy, press the Customize icon or adjust intensity using the intensity knob.

8. When treatment has completed, the Treatment Summary screen will appear with the following options:
 - Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
 - Repeat the treatment by pressing the Run this Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions on customized settings.

SEQUENCING OPERATION

The Vectra® Neo Clinical Therapy System offers sequencing for special electrotherapy waveform treatment purposes and stores these protocols in the system memory for recall and use. Sequencing allows you to link up to 3 electrotherapy protocols together to encompass one treatment session. Each sequence can have up to 3 protocols. The Vectra® Neo Clinical Therapy System memory will accommodate up to 200 user-defined protocols, including all user protocols, sequences, and system default protocols. The Vectra® Neo Clinical Therapy System denotes default protocols with an asterisk (*) before the named sequence.

Complete the following steps to create and access customized sequences:

1. Press the Electrotherapy icon from the Home screen.
2. Press the Sequencing icon.
3. Select a sequence from the list box by pressing the Up and Down arrows to find the desired sequence and press the “√” icon. The screen shown below will appear.



4. Make any desired changes to the existing sequences by using the Up and Down arrows to find the desired sequence and press the Edit icon. This screen also allows you to:
 - Delete a sequence from the protocol by pressing the Delete icon.
 - Add a new sequence to the protocol by pressing the New icon and using the Up and Down arrows to find the desired protocol in the list box. Press the “√” icon to add the new sequence.
 - Save the newly created sequence by pressing the Save icon and following the saving a treatment protocol instructions listed in the ELECTROTHERAPY OPERATION section on page 48.
5. Set the intensity for each protocol by using the Up and Down arrows for each sequence and press the Start button to begin treatment.
6. When treatment is completed, the Treatment Summary screen will appear with the following options:
 - Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
 - Repeat the treatment by pressing the Run this Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.

ULTRASOUND OPERATION

The Vectra® Neo Clinical Therapy System Ultrasound modality allows the user to select specific Applicator recommendations and edit treatment parameters for various syndromes requiring the use of ultrasound therapy. The following information gives general instructions for the setup of Ultrasound Therapy when selecting Ultrasound from the Home screen.

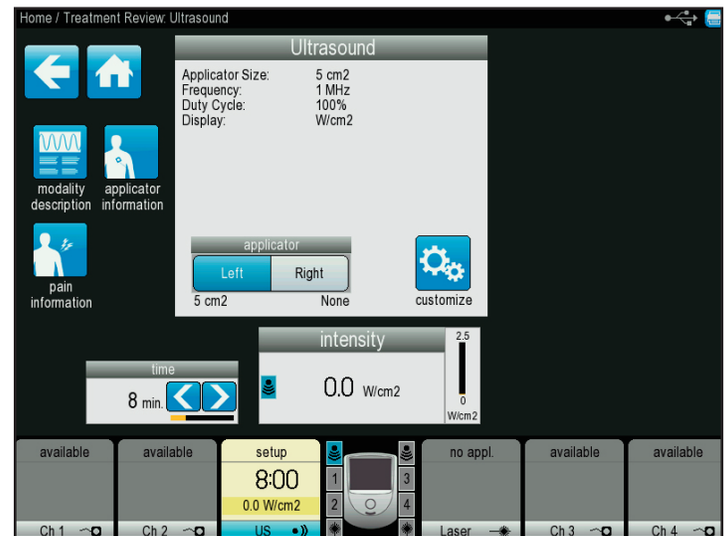
WARNING

- **The Vectra® Neo Clinical Therapy System may be susceptible to Electrostatic Discharge (ESD) at greater than ± 4 kV when first grasping either the Ultrasound or Laser Applicator. In the event of such a discharge, the Vectra® Neo Clinical Therapy System may experience communication loss with the installed modules. The Vectra® Neo Clinical Therapy System will terminate all active outputs (stim, ultrasound, laser), automatically place the unit in a safe state, and issue an error message 301 or 307.**
 - **To recover from an error message 301 or 307, turn the unit off and on using the ON/OFF switch located at the top of the display. Once the system restarts, re-initiate all treatments that were interrupted.**
- **To prevent Electrostatic Discharge (ESD) at greater than ± 4 kV:**
 - **Grasp and hold the Ultrasound or Laser Applicator prior to starting treatment. If the applicator must be put down prior to completion of treatment, stop the current treatment first and then place the applicator in the holder.**
 - **Maintain humidity in the use environment to at least 50% relative humidity.**
 - **Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, DJO recommends implementing additional controls to maintain relative humidity to at least 50%.**
 - **Communicate these ESD-precautionary procedures to healthcare staff, contractors, visitors, and patients.**

NOTE: Prior to starting treatment, replace the ultrasound faceplate on the side of the module not being used with a blank faceplate.

Complete the following steps to begin Ultrasound treatment:

1. Touch the screen to activate the system. Read and carefully follow the instructions on the screen.
 - To prepare the patient's skin for Ultrasound Therapy, prepare patient as described in the ULTRASOUND PATIENT PREPARATION section on page 41.
2. From the Home screen, select the Ultrasound icon. The screen shown below will appear:



- To view information explaining the modality, select the Modality Description icon. Press the Up and Down icons to view additional text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
- To view applicator information, press the Applicator Information icon:
 - Touch the body part you wish to treat.
 - Information about applicator sizes, frequency, and treatment directions will appear as seen on the screen above. Press the Down icon to scroll through the information.
 - Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
- Use the Intensity dial to set therapy intensity:
 - Clockwise – Increases Intensity
 - Counterclockwise – Decreases Intensity

ULTRASOUND OPERATION (CONTINUED)

3. Press the Start button to begin therapy, the Pause button to pause treatment, or the Stop button to terminate the treatment.
4. When treatment has completed, the Treatment Summary screen will appear with the following options:
 - Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
 - Repeat the treatment by pressing the Run this Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.

COMBINATION OPERATION

The Vectra® Neo Clinical Therapy System Combination modality allows the user to select and use Ultrasound therapy in combination with electrical muscle stimulation.

Combination therapy utilizes the Ultrasound modality in conjunction with High Voltage Pulsed Current (HVPC), IFC (4 pole), IFC Premodulated (2 pole), Asymmetrical Biphasic, Symmetrical Biphasic, or VMS™ to generate a therapeutic effect. In this mode of therapy, the Applicator of the Ultrasound Applicator becomes one-half of the electrical circuit. An electrode attached to the Black (-) lead wire completes the circuit.

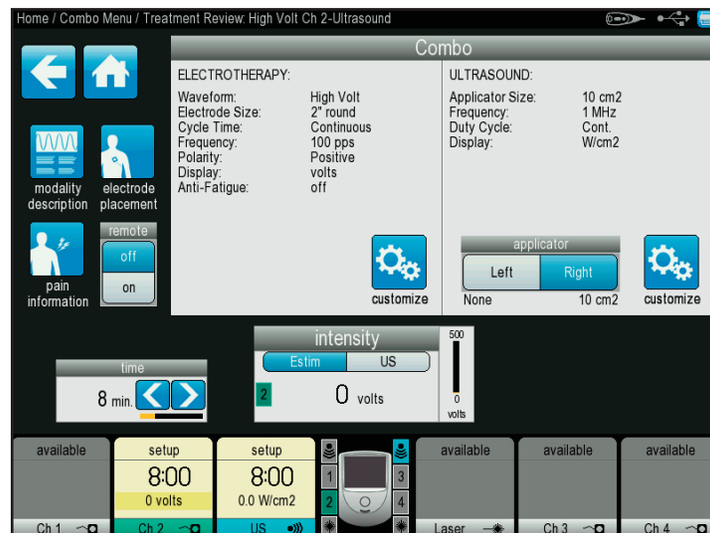
Complete the following steps to begin Combination treatment:

1. Touch the screen to activate the system. Read and carefully follow the instructions on the screen.
2. Prepare patient and therapy system. Refer to the PATIENT PREPARATION section on page 38 for electrode selection, preparing the patient, and securing electrodes. For ULTRASOUND PATIENT PREPARATION, refer to page 41.
3. Connect the Black (-) lead wire from Channel 2 to the electrode. Make certain the lead wire is completely seated in the electrode. The Red (+) lead wire is not used. The Ultrasound Applicator completes the circuit for Combination Therapy.



4. From the Home screen, select the Combo icon.

5. Select the ultrasound combination therapy desired, and press the corresponding icon. The Treatment Review screen shown below will appear:

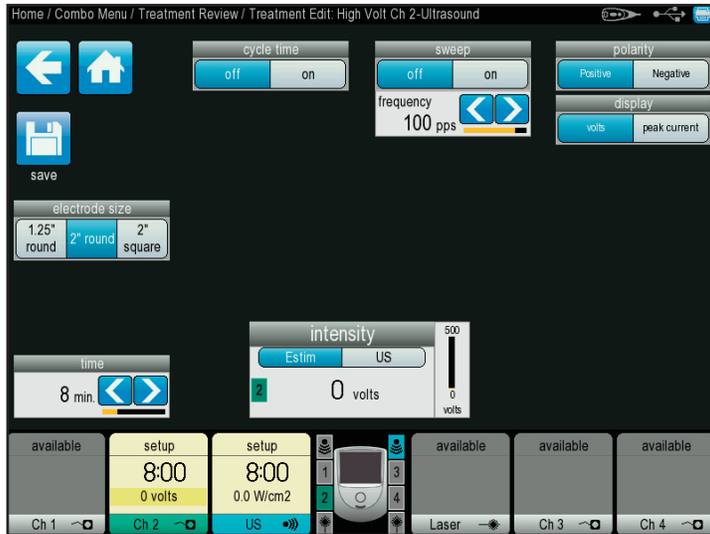


- To view information explaining the waveform, select the Modality Description icon. Press the Up and Down icons to view additional text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
- To view the most commonly used electrode placement for the selected waveform, select the Electrode Placement icon. A picture will appear prompting you to select the body part you wish to treat. Press the Down icon to scroll through text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.

COMBINATION OPERATION (CONTINUED)

- To customize settings shown in the list boxes, press the Customize icon located in the list box, and the screen shown below will appear.

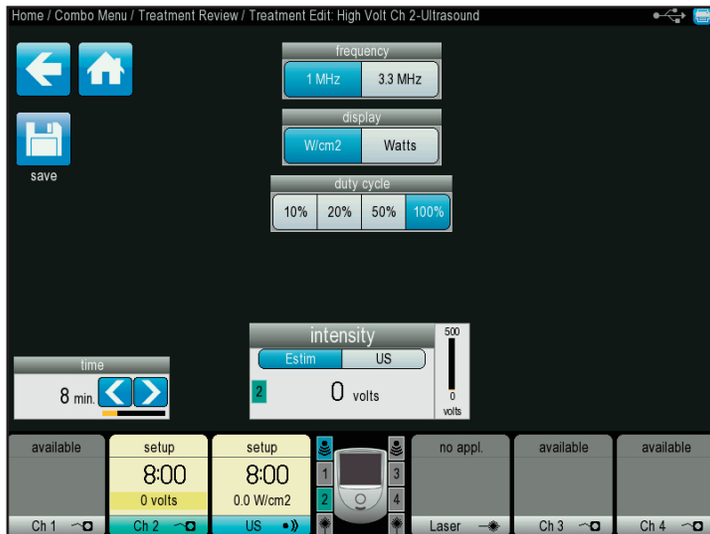
Electrotherapy Customization Screen



6. Press the Start button to begin therapy, the Pause button to pause treatment, or the Stop button to terminate the treatment.
7. When treatment has completed, the Treatment Summary screen will appear with the following options:

- Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
- Repeat the treatment by pressing the Run This Treatment icon.
- Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized setting.

Ultrasound Customization Screen



- Make the desired changes and press the Back Arrow icon to return to the previous screen, the Home icon to return to the home screen, or the Save icon to save the customized settings. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.
- To adjust treatment time, press the left or right arrows to reach desired treatment time.
- Use the Intensity dial to set therapy intensity:
 - Clockwise – Increases Intensity
 - Counterclockwise – Decreases Intensity

sEMG OPERATION

The Vectra® Neo Clinical Therapy System sEMG + Stim modality utilizes sEMG biofeedback activity coupled with triggered electrical muscle stimulation using selected electrotherapy waveforms for the maximum benefit in muscle retraining. The following options are available: sEMG (Channel 1), sEMG (Channel 2), sEMG (Channels 1 & 2), sEMG + Stim VMS, and sEMG + Stim Sym Biphasic.

The Electrical Muscle Stimulation is triggered when the muscle contraction (sEMG portion of the therapy) reaches the target, sEMG stops, and the muscle is then electrically stimulated for the pre-set period. After stimulation, the patient is given a short rest period and then repeats the muscle contraction, attempting to reach the target to again trigger the electrical stimulation. This is repeated throughout the therapy session.

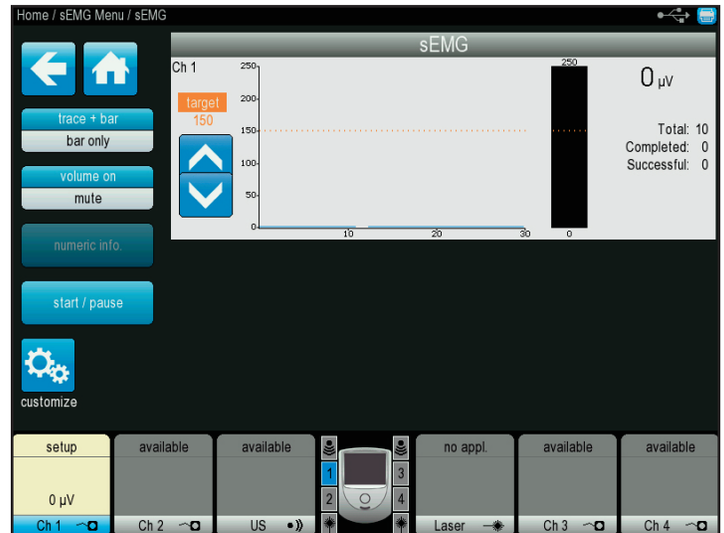
Session parameters can be stored on a USB storage drive and accessed on the Home screen under the Patient Data icon. The sEMG portion of sEMG + Stim modality is used to force the patient to contract the muscle to a prescribed target. The data cannot be recorded or stored on the Patient Data Card or the sEMG Data Card.

NOTE: This section applies when Stimulation Channels and sEMG module is installed.

Complete the following steps to begin sEMG treatment:

1. Touch the screen to activate the system.
2. Prepare patient and therapy system. Refer to the PATIENT PREPARATION section on page 38 for electrode selection, preparing the patient, and securing electrodes.
3. Select sEMG icon from the Home screen.

4. Select the prescribed channel icon (see previous for the available choices). The treatment screen will appear (the screen shown below illustrates the sEMG Channel 1 selection).



- To set the Target Threshold either Automatically or Manually, touch the CUSTOMIZE Icon.
 - Use the Intensity dial to set therapy intensity:
 - Clockwise – Increases Intensity
 - Counterclockwise – Decreases Intensity
5. Press Start/Pause button to begin therapy. Screen prompts will appear:
 - **Contract:** Instructs the patient to attempt reaching the Target Threshold. “Contract” and a green vertical line appear on the screen, indicating the patient should attempt to contract the selected muscle(s). “Contract” remains on the screen until the patient’s sEMG output reaches the Target Threshold, at which time Electrical Stimulation is delivered.
 - **Hold:** Instructs the patient to contract the selected muscle(s). When the Target Threshold is reached, the “Hold” prompt appears, instructing the patient to continue to contract the selected muscle(s) until the pre-set time for the Stimulation ends. A yellow vertical line appears at the end of the Hold period.
 - **Relax:** Instructs the patient to relax. “Relax” appears, indicating the patient should relax, stopping the contraction. Relax continues for the pre-set time. The cycle repeats when “Contract” re-appears, again indicating the patient should attempt to contract the selected muscle(s).

SEMG OPERATION (CONTINUED)

6. The following options are available under the Customize Treatment screen and accessed by pressing the appropriate icon:
 - To view information explaining the waveform, press the Modality Description icon. Press the Up and Down icons to view additional text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
 - To view the most commonly used electrode placement for the selected waveform, press the Electrode Placement icon. A picture will appear prompting you to select the body part you wish to treat. Press the Down icon to scroll through text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.

NOTE:When using sEMG + Stim, the Customization screen adds two icons to edit Stimulation settings and to set the stimulation intensity.

 - Use the Intensity dial to set therapy intensity:
 - Clockwise - Increases Intensity
 - Counterclockwise - Decreases Intensity
7. Make the desired changes and press the Back Arrow icon to return to the previous screen and begin treatment, the Home icon to return to the home screen, or the Save icon to save the customized settings. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.
8. Press the Start button to begin therapy, the Pause button to pause treatment, or the Stop button to terminate the treatment.
9. When treatment has completed, the Treatment Summary screen will appear with the following options:
 - Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
 - Repeat the treatment by pressing the Run This Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.

LASER OPERATION

Before applying Laser therapy to the patient, you must first prepare the patient’s skin as described in the PATIENT PREPARATION section on page 38. By properly preparing the patient’s skin for Laser therapy, you will allow more Laser energy to reach the targeted areas and reduce the risk of skin irritation.

⚠ DANGER



- **Laser protective eyewear should be worn during laser treatment by the operator and patient to block infrared light energy from the eyes during treatment.**

⚠ WARNING

- **The Vectra® Neo Clinical Therapy System may be susceptible to Electrostatic Discharge (ESD) at greater than ±4 kV when first grasping either the Ultrasound or Laser Applicator. In the event of such a discharge, the Vectra® Neo Clinical Therapy System may experience communication loss with the installed modules. The Vectra® Neo Clinical Therapy System will terminate all active outputs (stim, ultrasound, laser), automatically place the unit in a safe state, and issue an error message 301 or 307.**
 - **To recover from an error message 301 or 307, turn the unit off and on using the ON/OFF switch located at the top of the display. Once the system restarts, re-initiate all treatments that were interrupted.**
- **To prevent Electrostatic Discharge (ESD) at greater than ±4 kV:**
 - **Grasp and hold the Ultrasound or Laser Applicator prior to starting treatment. If the applicator must be put down prior to completion of treatment, stop the current treatment first and then place the applicator in the holder.**
 - **Maintain humidity in the use environment to at least 50% relative humidity.**
 - **Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, DJO recommends implementing additional controls to maintain relative humidity to at least 50%.**
 - **Communicate these ESD-precautionary procedures to healthcare staff, contractors, visitors, and patients.**

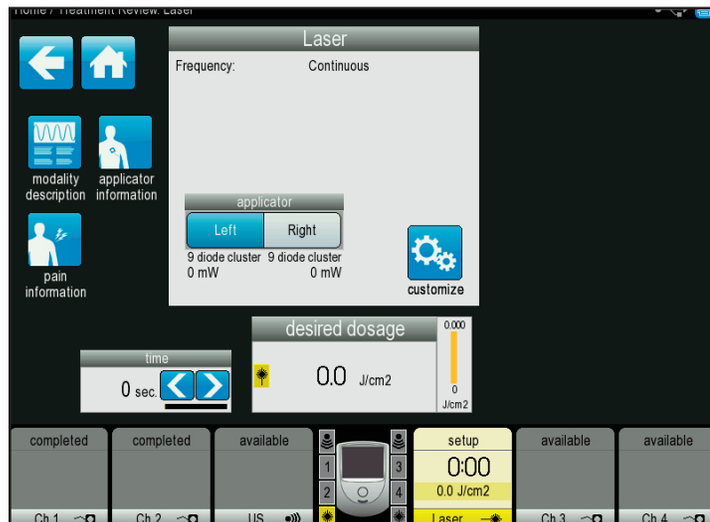
**Refer to the TREATMENT TIPS section on page 71.

NOTE: Prior to starting treatment, replace the laser faceplate on the side of the module not being used with a blank faceplate.

LASER OPERATION (CONTINUED)

Complete the following steps to begin Laser therapy treatment:

1. Touch the screen to activate the system. Read and carefully follow the instructions on the screen.
2. Select the Laser icon from the Home screen.
3. Type in a four-digit PIN and press the Accept icon to accept the entry or choose from the following options:
 - The Cancel icon to return to the Home screen
 - The Clear icon to clear the numbers entered
 - The Left and Right arrows to scroll through the numbers entered
 - The Delete icon to delete numbers entered one at a time
4. After successfully entering your PIN, the screen shown below will appear:



- To view information explaining the modality, press the Modality Description icon. Press the Up and Down icons to view additional text. Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.
5. Verify that the Patient Remote/Laser Interrupt Switch is connected and provided to the patient.
 - To view applicator information, press the Applicator Information icon:
 - Touch the body part you wish to treat.
 - Information about applicator sizes, frequency, and treatment directions will display. Press the Down icon to scroll through the information.
 - Press the Back Arrow icon to return to the previous screen or the Home icon to return to the Home screen.

- To customize settings shown in the list boxes, press the Customize icon located in the list box, and the screen shown below will appear.



- Make the desired changes and press the Back Arrow icon to return to the previous screen, the Home icon to return to the Home screen, or the Save icon to save the customized settings. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.
6. Use the Intensity dial to set therapy duration:
 - Clockwise – Increases duration
 - Counterclockwise – Decreases duration
 7. Press the Start button to begin therapy, the Pause button to pause treatment, or the Stop button to terminate the treatment.
 8. When treatment has completed, the Treatment Summary screen will appear with the following options:
 - Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
 - Repeat the treatment by pressing the Run This Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.

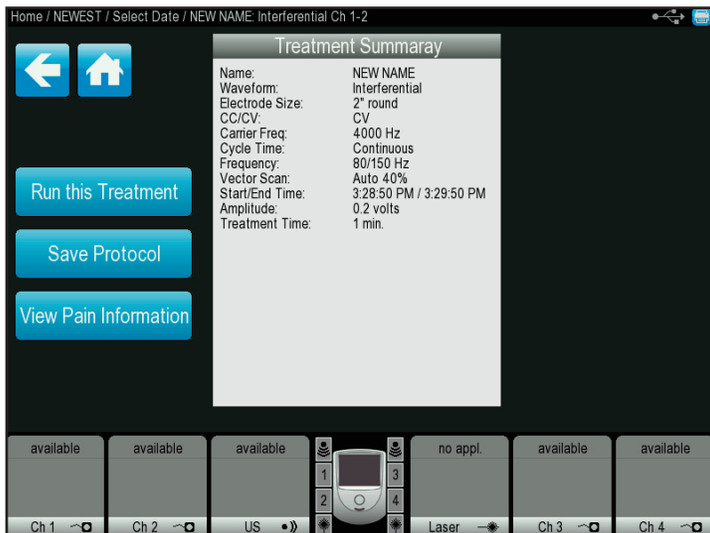
SAVING TO USB FLASH DRIVE/PATIENT DATA

Patient treatment data can be saved to the USB flash drive for viewing/printing on a PC, as well as for retrieving for later use on the unit or multiple units.

Complete the following steps to view and access patient data:

1. Press the Patient Data icon on the Home screen. The screen will display a list box of all previously saved patient data sessions found on the USB flash drive connected to the unit.
2. Select the patient ID from the list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the patient ID information. Select the patient ID you wish to view and access by pressing the “√” icon.
3. Select the treatment date desired from the list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the treatment date information. Select the treatment date you wish to view and access by pressing the “√” icon.
4. The Treatment Summary list box will appear with detailed information about the specific treatment.

5. Choose one of the following options from the Treatment Summary screen:
 - Run the treatment by pressing the Run this Treatment icon.
 - Save the treatment protocol by pressing the Save Protocol icon. Refer to the CUSTOM PROTOCOLS section on page 59 for detailed instructions for saving customized settings.
 - View the pain information by pressing the View Pain Information icon.
 - Save therapy information to USB flash drive by inserting a USB flash drive and pressing the Save Protocol icon.



CUSTOM PROTOCOLS

A new custom protocol may be saved at either the Treatment Edit or Treatment Complete screen. The Vectra® Neo Clinical Therapy System will allow a maximum of 200 custom protocols to be defined, and the Home screen will allow for 5 custom protocol shortcut assignments.

Complete the following steps to save a customized protocol created from a Treatment Summary screen:

1. Press the Save Protocol icon from either the Treatment Edit or the Treatment Complete screen.



2. Save the new protocol by pressing the Save as a New Name icon and typing in a name for the protocol, or overwrite a currently saved protocol by scrolling to the protocol using the Up and Down arrow icons. Press the Save icon when finished typing in the protocol name. A yellow text box will appear confirming the newly saved protocol and name.



3. After saving the protocol, you will return to the Treatment Summary screen. Complete one of the following actions listed on the Treatment Summary screen:

- Return to Home screen by pressing the Home icon.
- Scroll back one screen at a time by pressing the Back Arrow icon.
- Run the treatment again by pressing the Run this Treatment icon.
- Save therapy information to a USB flash drive by inserting a USB flash drive into the device and pressing the Save to USB Flash Drive icon.
- Clear the treatment from "Completed" to "Available" on the involved channel by pressing the Clear this Treatment icon.

NOTE: The treatment protocol cannot be saved or run once Clear the Treatment is selected.



CUSTOM PROTOCOLS (CONTINUED)

Complete the following steps to delete a previously saved protocol:

1. From the Home screen, press the Custom Protocols icon.
2. Find the previously saved protocol in the list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the protocol. Select the protocol you wish to delete and press the Delete icon at the top of the screen (shown below).



3. A confirmation prompt will appear asking, "Are you sure you want to delete protocol ____". Press the Yes icon to delete the protocol.
4. Return to Home screen by pressing the Home icon, or press the Back Arrow icon to scroll back one screen at a time.

Complete the following steps to assign a Home screen shortcut for a customized protocol:

1. Press one of the 5 Assign Shortcut icons on the Home screen.



2. Find the previously saved protocol in the list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the protocol. Select the protocol for which you wish to create a shortcut by pressing the "✓" icon.



3. A yellow text box will appear confirming the newly saved protocol and name, and you automatically return to the Home screen. The newly saved protocol will now appear on the Home screen as a unique icon.

CUSTOM PROTOCOLS (CONTINUED)

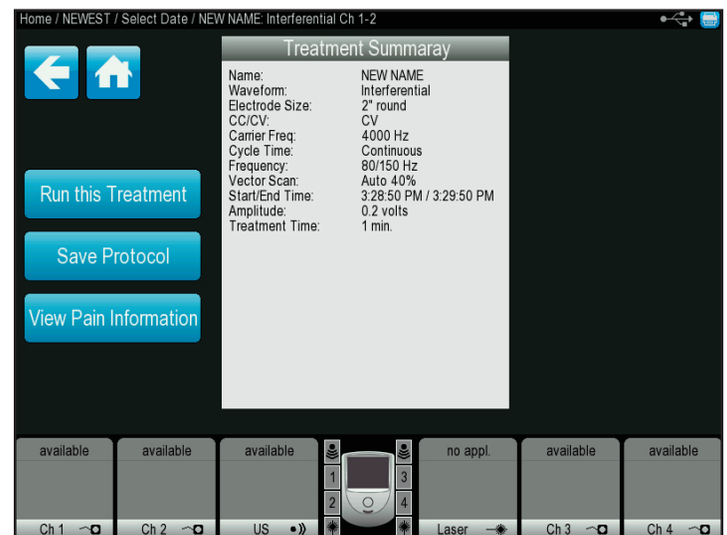
Complete the following steps to deassign a Home screen shortcut for a customized protocol:

1. From the Home screen, press and hold the shortcut icon you wish to deassign.

NOTE: You can only deassign a shortcut if that shortcut currently has a protocol assigned to it.
2. The unit will display a text box asking, "Are you sure you want to de-assign protocol _____ shortcut?"
3. Press the No icon to quit the deassignment process and return to the Home screen or press the Yes icon to continue with the deassignment process.
4. After pressing the Yes icon to continue the deassignment process, a confirmation message will display for 5 seconds and return to the Home screen. The previously assigned shortcut will no longer appear on the Home screen.

Complete the following steps to view and access patient data:

1. Press the Patient Data icon on the Home screen.
2. Select the patient ID from the list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the patient ID information. Select the patient ID you wish to view and access by pressing the "√" icon.
3. Select the treatment date desired from the list box by pressing the Up and Down arrows or Previous Page/Next Page icons to locate the patient ID information. Select the treatment date you wish to view and access by pressing the "√" icon.
4. The Treatment Summary list box will appear with detailed information about the specific treatment.



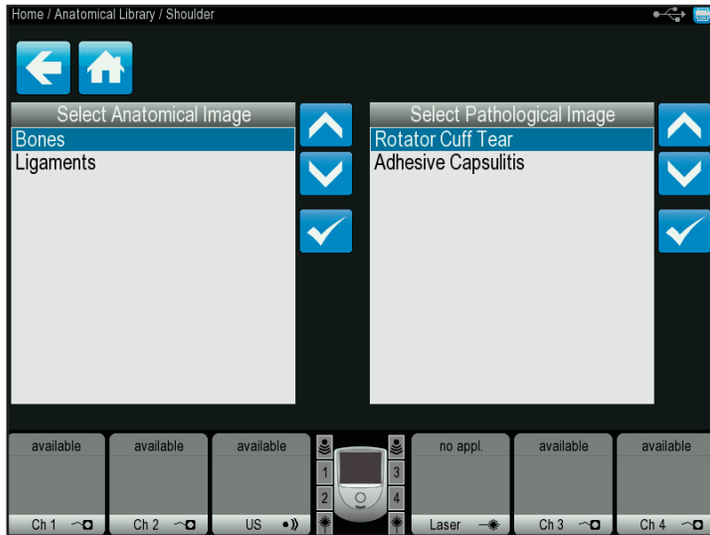
5. Choose one of the following options from the Treatment Summary screen:
 - Run the treatment by pressing the Run the Treatment icon.
 - View the pain information by pressing the View Pain Information icon.

ANATOMICAL LIBRARY

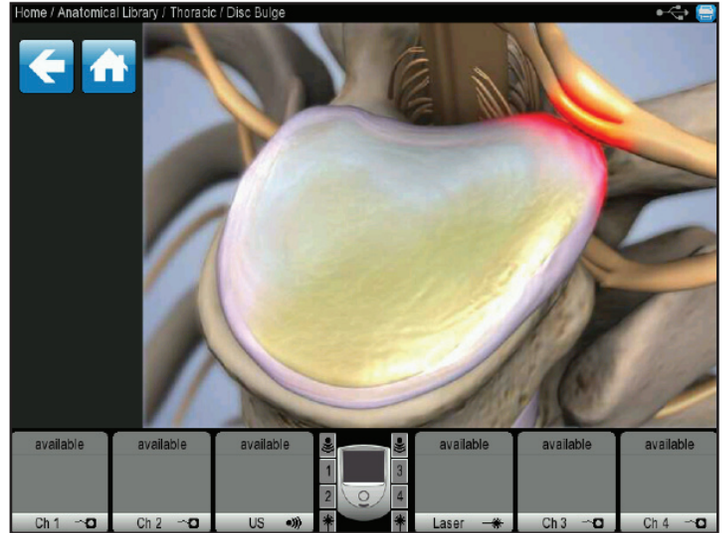
The Vectra® Neo Clinical Therapy System contains a unique Anatomical Library designed to aid the operator in visually understanding and locating specific muscle groups and commonly found problems associated with pathological conditions, as well as providing an educational tool for the clinician to use with the patient.

Complete the following steps to view the Anatomical Library:

1. Press the Anatomical Library icon on the Home screen.
2. Touch the body part for which you wish to view information. Choose either anterior (on left of screen) or posterior (on right of screen).
3. Select an image from either the Anatomical Image list box or the Pathological Image list box by using the Up and Down arrows or Previous Page/Next Page icons to locate the desired image. Select the image you wish to view and access by pressing the “√” icon.



4. View the selected image and, if desired, print the selected image by pressing the printer icon:



5. Press the Back arrow icon to scroll to the previous screen or Home icon to return to the Home screen.

TROUBLESHOOTING CODES

The Vectra® Neo Clinical Therapy System incorporates information, warning, and error messages to inform the user of problems or potential problems with the system, modality, or accessories. **The numbering sequence is: 100-level messages are general use information messages, 200-level messages are warnings, and 300-level messages are errors.** Use the following Troubleshooting Chart to define the code and locate the probable cause and possible remedies before contacting your dealer or the DJO office.

| Code Number | Type Message | Probable Cause | Possible Remedies |
|-------------|--------------|--|--|
| 100 | Information | Attempting to save a treatment to a USB flash drive with a blank Patient ID | Type in or select a Patient ID prior to saving treatment to a USB flash drive. |
| 101 | Information | Attempting to save a Custom protocol with a blank protocol name | Ensure protocol name is not blank prior to saving the protocol. |
| 102 | Information | Attempting to save a Sequence with a blank sequence name | Ensure sequence name is not blank prior to saving the sequence. |
| 103 | Information | Attempting to save the Clinic Name with a blank name | Ensure Clinic name is not blank prior to saving the Clinic name. |
| 104 | Information | Attempting to delete a factory sequence | Factory sequences (names begin with a “*”) cannot be deleted. |
| 105 | Information | Attempting to delete a factory protocol | Factory protocols (names begin with a “**”) cannot be deleted. |
| 106 | Information | Attempting to save a Custom protocol or Sequence after system memory has reached the maximum allowed (200) | Delete unused User protocols or sequences. |
| 107 | Information | Attempting to access protocols or sequences and none are found in the system | 1. Custom protocols: No Custom protocols have been saved in the system. Refer to the CUSTOM PROTOCOLS section on page 59. |
| 108 | Information | | 2. Sequences: No User sequences have been saved in the system. Refer to the SEQUENCING OPERATION section on page 50. |
| 109 | Information | | |
| 110 | Information | Ultrasound applicator became unplugged during a treatment | Ensure ultrasound applicator is securely connected to the unit prior to starting an ultrasound treatment. |
| 111 | Information | No ultrasound applicator is plugged into the unit | Ensure desired ultrasound applicator is securely connected to the unit prior to selecting an ultrasound treatment. |
| 112 | Information | Attempting to select a treatment but available channels for desired modality are currently in use | Complete existing treatment before attempting to start another treatment on the same channel. |
| 113 | Information | Attempting to select an sEMG treatment but sEMG channels are currently in use | Complete existing treatment before attempting to start another treatment on the same channel. |
| 114 | Information | Attempting to select an sEMG treatment but no sEMG is installed on the Stim 1/2 module. | 1. If sEMG is installed on the Stim 1/2 module, reset the Therapy System by turning the Therapy System Off then back On via the Power button. 2. If sEMG is installed on the Stim 1/2 module and the problem persists, discontinue use of the device and contact the dealer or factory for technical service. |
| 115 | Information | Attempting to select a Laser treatment while another treatment is running | Laser treatments cannot be run concurrently with another treatment. Complete existing treatment(s) before attempting to select a laser treatment. |
| 116 | Information | No laser applicator is plugged into the unit | Ensure desired laser applicator is securely connected to the unit prior to selecting a laser treatment. |
| 117 | Information | Laser Applicator became unplugged during a laser treatment | Ensure desired laser applicator is securely connected to the unit prior to selecting a laser treatment. |
| 118 | Information | Incorrect Laser PIN was entered | Enter the correct laser PIN when prompted prior to starting a laser treatment. |
| 119 | Information | Attempting to select a laser treatment but no laser module is installed in the unit | 1. If laser module is installed in the unit, ensure module is securely inserted. 2. If laser module is not installed, contact the dealer or factory representative to purchase one. |
| 120 | Information | Attempting to start a Laser treatment but the remote control is not plugged into the unit | Prior to starting a laser treatment, the remote control must be plugged into the unit and given to the patient to allow use as an emergency stop. |

TROUBLESHOOTING CODES (CONTINUED)

| Code Number | Type Message | Probable Cause | Possible Remedies |
|-------------|--------------|---|--|
| 121 | Information | Attempting to select an electrotherapy treatment but no Stim module is installed in the unit | <ol style="list-style-type: none"> 1. If a Stim module is installed in the unit, ensure module is securely inserted into the unit. 2. If a Stim module is not installed, contact the dealer or factory to purchase one. |
| 122 | Information | Attempting to select a Combination treatment but no Stim module #1 (channels 1 and 2) is installed in the unit | <ol style="list-style-type: none"> 1. If Stim module #1 is installed in the unit, ensure module is securely inserted into the unit. 2. If Stim module #1 is not installed, contact the dealer or factory to purchase one. |
| 123 | Information | Attempting to select an Ultrasound or Combination treatment but no Ultrasound module is installed in the unit | <ol style="list-style-type: none"> 1. If Ultrasound module is installed in the unit, ensure module is securely inserted into the unit. 2. If Ultrasound module is not installed, contact the dealer or factory to purchase one. |
| 124 | Information | Attempting to perform an action requiring the USB flash drive but no USB flash drive is plugged into the unit or USB flash drive cannot be accessed correctly | <ol style="list-style-type: none"> 1. Ensure valid USB flash drive is firmly inserted into the unit or 2. Try a different USB flash drive. |
| 125 | Information | | |
| 126 | Information | | |
| 127 | Information | Attempting to perform an action requiring a particular file on the USB flash drive but the file does not exist on the flash drive | <ol style="list-style-type: none"> 1. Ensure USB flash drive contains the correct file for the desired function. 2. Ensure valid USB flash drive is firmly inserted into the unit or 3. Try a different USB flash drive. |
| 128 | Information | | |
| 129 | Information | | |
| 130 | Information | | |
| 132 | Information | Attempting to perform a Print Screen to the USB flash drive while a treatment is running | The Print Screen function is not allowed while a treatment is running. Wait for treatment to complete and then try again. |
| 133 | Information | Attempting to import protocols from the USB flash drive while a treatment is running | The Protocol Import function is not allowed while a treatment is running. Wait for treatment to complete and then try again. |
| 203 | Warning | Error reading a protocol from internal storage | <ol style="list-style-type: none"> 1. Reset the Therapy System by turning the power switch Off and On. 2. If problem persists after resetting the unit, select the Restore Default Protocols in the Utilities screen. Refer to the UTILITIES AND OPTIONS section on page 44. 3. If problem persists, discontinue use of the device and contact the dealer or factory for technical service. |
| 210 | Warning | Laser application is out of calibration | <ol style="list-style-type: none"> 1. Discontinue use of this laser applicator and contact the dealer or factory for technical service to have the laser applicator recalibrated. 2. If problem exists on multiple laser applicators then discontinue use of laser and contact the dealer or factory for technical service. |
| 211 | Warning | | |
| 212 | Warning | | |
| 213 | Warning | | |
| 216 | Warning | Error reading a protocol from internal storage | <ol style="list-style-type: none"> 1. Reset the Therapy System by turning the power switch Off and On. 2. Restore the Default Protocols in the Utilities screen. Refer to the UTILITIES AND OPTIONS section on page 44. 3. If problem persists, discontinue use of the device contact the dealer or factory for technical service. |
| 233 | Warning | Electrotherapy - bad electrode contact, overcurrent, or shorted leads condition has been detected | <ol style="list-style-type: none"> 1. Ensure proper electrode placement use guidelines are followed. 2. Check electrode lead wire connections both at the unit and at the electrode. 3. Replace electrodes. 4. Replace lead wires. |
| 234 | Warning | | |
| 235 | Warning | | |

TROUBLESHOOTING CODES (CONTINUED)

| Code Number | Type Message | Probable Cause | Possible Remedies |
|-------------------------------------|--------------|--|-------------------|
| Any 200-level code not listed above | Warning | <p>Immediately stop all use of the system and contact the dealer or DJO for service. Warnings in these categories indicate an internal problem with the system that must be tested and corrected by DJO or a Trained Technician before any further operation or use of the system.</p> <p>Use of a system that indicates a Warning in these categories may pose a risk of injury to the patient, user, or extensive internal damage to the system.</p> | |
| Any 300-level code | Error | <p>Immediately stop all use of the system and contact the dealer or DJO for service. Errors in these categories indicate an internal problem with the system that must be tested and corrected by DJO or a Trained Technician before any further operation or use of the system.</p> <p>Use of a system that indicates an Error in these categories may pose a risk of injury to the patient, user, or extensive internal damage to the system.</p> | |

REPLACEMENT ACCESSORIES

The following provides users of the Vectra® Neo Clinical Therapy System the necessary information to order replacement accessories used with the system. This list of replacement accessories is designed for use with the Vectra® Neo Clinical Therapy System. When ordering, provide the respective part number, description, and quantity desired.

| ELECTRODES | |
|-------------------|-------------------------------------|
| Model Number | Description |
| 42045 | ELECTRODE DURA-STICK 2.75"X 5" |
| 42159 | DURA-STICK+ 1.5"X2.5" OV 40/CASE |
| 42160 | DURA-STICK+ 2"X4"OV 40/CASE |
| 42170 | DURA-STICK PREM 1.25" RND 40/CASE |
| 42171 | DURA-STICK PREM 2" RND 40/CASE |
| 42172 | DURA-STICK P 1.5"X2.5" OV 40/CASE |
| 42173 | DURA-STICK PREM 3"X5" OV 20/CASE |
| 42174 | DURA-STICK PREM 2" SQ 40/CASE |
| 42175 | DURA-STICK P 2"X3.5" REC 40/CASE |
| 42176 | DURA-STICK BG 2" SQ 40/CA |
| 42177 | DURA-STICK BG 1.5"X3.5" REC 40/CASE |
| 42178 | DURA-STICK 2" SQ 40/CASE |
| 42179 | DURA-STICK 2"X3.5" REC 40/CASE |
| 42181 | DURA-STICK+ 2"X3.5" REC 40/CASE |
| 42182 | DURA-STICK+ 2"RND 40/CASE |
| 42183 | DURA-STICK+ 2" SQ 40/CASE |

| GENERAL ACCESSORIES | |
|----------------------------|--|
| Model Number | Description |
| 70000 | Electrotherapy Module Channels 1/2 |
| 70001 | Neo Cart |
| 70002 | Ultrasound Module |
| 70003 | Electrotherapy Module Channels 3/4 |
| 70004 | Electrotherapy Module Channels 1/2 + sEMG |
| 70005 | Laser Module |
| 70008 | PATIENT REMOTE/LASER INTERRUPT SWITCH |
| 79977 | HIGH VOLT PROBE KIT- Includes Probe and Sponge Applicator Tips (15 and 8 mm) |
| 70010 | STIM CH 1/2 LEADWIRE KIT STD |
| 70011 | STIM CH 3/4 LEADWIRE KIT STD |
| 70012 | STIM CH 1/2 LEADWIRE KIT XL |
| 70013 | STIM CH 3/4 LEADWIRE KIT XL |
| 70014 | STIM CH 1/2 + EMG LEADWIRE KIT |
| 70020 | Powercord US Black 6 FT |

REPLACEMENT ACCESSORIES (CONTINUED)

| LASER APPLICATORS AND EYEWEAR ACCESSORIES | |
|--|---|
| Model Number | Description |
| 27840 | Single 850nm Laser 100mW |
| 27804 | Single 850nm Laser 150mW |
| 27841 | Single 850nm Laser 200mW |
| 27803 | Single 850nm Laser 40mW |
| 27805 | Single 820nm Laser 300mW |
| 27810 | 9-Diode cluster: 5-50mW laser diodes + 4 LED's |
| 27811 | 9-Diode cluster: 5-100mW laser diodes + 4 LED's |
| 27812 | 9-Diode cluster: 5-200mW laser diodes + 4 LED's |
| 27813 | 13-Diode cluster: 3-50mW laser diodes + 10 LED's |
| 27814 | 13-Diode cluster: 3-100mW laser diodes + 10 LED's |
| 27816 | 13-Diode cluster: 3-200mW laser diodes + 10 LED's |
| 27802 | 33-Diode cluster: 5-50mW laser diodes + 28 LED's |
| 27807 | 33-Diode cluster: 5-100mW laser diodes + 28 LED's |
| 27808 | 33-Diode cluster: 5-200mW laser diodes + 28 LED's |
| 27799 | Single 670nm LED 10mW |
| 27815 | 19-Diode LED cluster, no laser |
| 27809 | 33-Diode LED cluster, no laser |
| 27525 | Laser Protective Eyewear |
| 27904K | Laser Interlock |

| ULTRASOUND APPLICATORS AND GEL | |
|---------------------------------------|---|
| Model Number | Description |
| 27333 | 1 cm ² Sound Head Applicator |
| 27334 | 2 cm ² Sound Head Applicator |
| 27335 | 5 cm ² Sound Head Applicator |
| 27336 | 10 cm ² Sound Head Applicator |
| 4248 | Conductor™ Transmission Gel - 9 oz Bottle |

CLEANING THE VECTRA® NEO CLINICAL THERAPY SYSTEM

With the system disconnected from the power source, clean the system with a clean, lint-free cloth moistened with water and mild antibacterial soap. If a more sterile cleaning is needed, use a cloth moistened with an antimicrobial cleaner.

Do not submerge the system in liquids. Should the unit accidentally become submersed, contact the dealer or DJO Service Department immediately. Do not attempt to use a system that has been wet inside until inspected and tested by a Trained Technician.

Do not allow liquids to enter the ventilation holes in the optional modules. This could permanently damage the modules.

Cleaning the LCD Screen

Clean the Therapy System LCD with a clean, dry cloth, in the same way as cleaning the computer monitor screen. Do not use abrasive materials or chemicals or liquids.

CALIBRATION REQUIREMENTS

Annual factory calibration is required for all Ultrasound and Laser Applicators. Only the Applicators should be sent to the factory or a Trained Technician for this procedure.

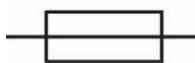
NOTE: The unit was calibrated during the manufacturing process and is ready to be placed into service upon delivery.

DEVICE DISPOSAL



Council Directive 2012/19/EU concerning Waste Electrical and Electronic Equipment (WEEE) requires not to dispose of WEEE as municipal waste. Contact your local distributor for information regarding disposal of the unit and accessories.

FUSE INFORMATION



3.15A, 250V AC

INSTRUCTION FOR SOFTWARE UPGRADE

1. Obtain a USB flash drive with the upgrade file in root directory.
2. Power on Unit with the USB flash drive inserted in the USB port. Allow initialization to complete.
3. Enter the Utilities screen by pressing the Utilities icon on the Home screen.
4. On the Utilities screen, press the Display Unit Version Information icon.
5. Press the Upgrade Unit Software from USB icon to apply any upgraded software packages to the unit.

COPY OF MANUAL

To obtain a copy of the Vectra® Neo Clinical Therapy System User Manual, Item #13-7646 (CD Version, Item #13-7647), contact your local representative or DJO Global Customer Care.

WARRANTY REPAIR/OUT OF WARRANTY REPAIR

Service

When the Vectra® Neo Clinical Therapy System or any of the accessory modules require service, contact the selling dealer or DJO Service Department.

All Therapy System and accessory modules returned to the factory for service must include the following:

1. Written statement containing the following information:
 - RA Number – Obtain from DJO
 - Therapy System or Module Model Number
 - Therapy System or Module Serial Number
 - Contact Person with Phone and Fax Numbers
 - Billing Address (for Out of Warranty Repair)
 - Shipping Address (Where to Ship Unit after Repair)
 - Detailed Description of Problem or Symptoms
2. Copy of original invoice issued at purchase of the Therapy System or Module
3. Ship the unit to address specified by an authorized Service Technician

Service to these units should be performed only by a service technician certified by the Company.

Ultrasound Applicators require annual calibration, from the date placed in service, by the Factory or a Trained Technician.

Through the purchase of a Service Manual, DJO, LLC has made available circuit diagrams, component part lists, descriptors, or other information which will assist authorized technical personnel to repair those parts of the equipment which are designated by DJO, LLC as repairable.

Expected Life

Expected life of the applicator is 5 years and the control unit is 10 years. Yearly calibration of the device would extend the life of the device for as long as servicing is available by DJO or other factory certified personnel.

WARRANTY

DJO, LLC (“Company”) warrants that the Vectra® Neo Clinical Therapy System, Channel 1/2 Electrotherapy Module, Channel 1/2 Electrotherapy+EMG Module, Channel 3/4 Electrotherapy Module, Laser Module, and Ultrasound Module (“Products”) are free of defects in material and workmanship. This warranty shall remain in effect for two years (24 months) from the date of original consumer purchase. If these Products fail to function during the two year warranty period due to a defect in material or workmanship, at the Company’s option, Company or the selling dealer will repair or replace the respective Product without charge within a period of thirty (30) days from the date on which the Product is returned to the Company or the dealer.

All repairs to the Product must be performed by a service center authorized by the Company. Any modifications or repairs performed by unauthorized centers or groups will void this warranty.

The warranty period for certain accessories is 90 days. Accessories consist of Lead Wires, Patient Remote/Laser Interrupt Switch, and Electrodes.

The warranty period for the Therapy System Cart, Laser Applicators and Ultrasound Applicators is one year (12 months).

To participate in warranty coverage, this Product’s warranty registration card (included with Product) must be filled out and returned to the Company by the original owner within ten (10) business days of purchase.

This Warranty Does Not Cover:

- Replacement parts or labor furnished by anyone other than the Company, the selling dealer, or a Company service technician
- Defects or damage caused by labor furnished by someone other than Company, the selling dealer, or a Company service technician
- Any malfunction or failure in the Product caused by product misuse, including, but not limited to, the failure to provide reasonable and required maintenance or any use that is inconsistent with the Product User’s Manual

COMPANY SHALL NOT BE LIABLE IN ANY EVENT FOR INCIDENTAL OR CONSEQUENTIAL DAMAGES.

Some states do not allow the exclusion or limitation of incidental or consequential damages, so the above limitation or exclusion may not apply to you.

To Obtain Service From Company or the selling dealer under this warranty:

1. A written claim must be made within the warranty period to the Company or the selling dealer. Written claims made to the Company should be sent to:

DJO, LLC
 1430 Decision Street
 Vista, CA 92081-8553 USA
 T: 1-800-592-7329 USA
 F: 1-760-734-5608

and

2. The Product must be returned to the Company or the selling dealer by the owner.

This warranty gives you specific legal rights and you may also have other rights which vary from location to location.

The Company does not authorize any person or representative to create for it any other obligation or liability in connection with the sale of the Product.

Any representative or agreement not contained in the warranty shall be void and of no effect.

THE FOREGOING WARRANTY IS IN LIEU OF ALL OTHER WARRANTIES, EXPRESSED OR IMPLIED, INCLUDING ANY WARRANTY OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

OVERVIEW OF LASER THERAPY

Laser therapy results in energy absorbed into the patient's tissue from light, triggering biological changes at a cellular level, resulting in the following:

- Topical heating for the temporary increase in local blood circulation
- Temporary relief of minor muscle and joint aches, pains, and stiffness
- Relaxation of muscles and relief of muscle spasm
- Temporary relief of minor pain and stiffness associated with arthritis. The dose and frequency of treatment can be adjusted to produce the desired effect.

There is no exact limit with respect to the penetration of the light. The light gets weaker the further from the surface it penetrates. There is, however, a limit at which the light intensity is so low that no biological effect of the light can be registered. This limit, where the effect ceases, is called "the greatest active depth." This depth is also contingent on tissue type, pigmentation, and dirt on the skin. For example, fat tissue is more transparent than muscle tissue and more easily penetrable by Laser than muscle tissue as blood hemoglobin is an important absorber of light, and highly vascularized muscle tissue makes Laser penetration more difficult.

NOTE: Some laser applicators may cause a noticeable heat sensation, particularly in hairy areas and on sensitive tissues such as lips.

TREATMENT TIPS

Contact

To obtain the most effective results, the applicator should be in contact with the patient's skin.

Applicator Position

Due to the characteristics of Laser, the angle at which the light enters the patient's skin is very important. Therefore, the applicator lens should always be parallel to the treatment area.

Treating Joints

If you are applying Laser therapy to a patient's joint, it is more effective to apply the Laser energy into the joint by positioning the joint in an open position (e.g., knee in flexion). However, do not attempt this method if it is uncomfortable to the patient.

Applicator Selection

If the injury you intend to treat is very small (pinpoint), you should only need to treat the area with a single diode applicator. If the area surrounding the treatment area is sensitive, it is recommended that you apply therapy with a single diode applicator first, then use the cluster applicator for the surrounding area.

Cold and Heat

If you intend to apply cold or heat as an adjunct to Laser therapy, use the following guidelines:

- Use cold before Laser therapy. This slows the flow of red blood cells and reduces the amount of energy removed from the area.
- Use heat after the treatment. This speeds the flow of red blood cells so that more energy can be removed from the area.

COMMON TERMS

Applicator – Hand held assembly that delivers Laser energy and includes laser head, diode, and related electronics

Collimating – The shape of the Laser beam. It is neither focused or dispersed and resembles a column when applied

Continuous Mode – The output of the Laser is not interrupted during the treatment time

Dosage – Intensity measurement of the Laser energy over the treatment area. It is measured in Joules or Joules/cm²

Energy – Measured in Joules, energy equals the treatment time multiplied by the power. Energy density equals the power output multiplied by the treatment time, and divided by the spot size (cm²). This gives a more specific measurement of energy delivered

Frequency – Pulsed frequencies are selectable from 8 to 10,000 Hz

Laser Head – The clear lens applicator face contacts the patient's skin. It consists of laser diodes with or without LED's or SLD's (depending on the applicator)

Power – Measured in Watts (W), power wattage is directly proportional to treatment time and penetration of the Laser energy. High-powered diodes will reduce patients' treatment times and give a higher amount of energy at a deeper depth. Power output can be either continuous or pulsed

Power Density - Ratio of power divided by treatment time

Pulsed Mode - This is the ratio of the "On" time to the "Total" time of the cycle, expressed as a percentage. The lower the percentage, the lower temporal average intensity. 100% is continuous Laser. Pulsed Mode is 90% on and 10% off

NOTE: Pulsed Mode is also equivalent to Duty Cycle.

Spot Size - Area of the LED, SLD, or laser beam when it leaves the face of the lens

Treatment Area - Area of tissue affected by LED, SLD, or laser when wavelength, divergence angles, and depth of penetration are factored. This is the area used to calculate dosage

Treatment Time – Measured in seconds, it is the suggested time per laser point that therapy is given

Wavelength – Wavelength is measured in nanometers (nm) and is the key component in obtaining effective therapy as different wavelengths cause different physiological effects. Superficial skin disorders are most effectively treated at wavelengths 600-700 nm, while deeper muscular or ligament lesions and joint conditions are best treated at higher wavelengths of 700-1000 nm.

ELECTROMAGNETIC COMPATIBILITY (EMC)

The Vectra® Neo Clinical Therapy System has been tested and found to comply with the electromagnetic compatibility (EMC) limits for medical devices to IEC 60601-1-2. These limits are designed to provide reasonable protection against harmful interference in a typical medical installation.

The guidelines below are intended to help promote electromagnetic compatibility (EMC) in the identified use environment for the Vectra® Neo Clinical Therapy System.

- **Make use of available resources** such as EMC professionals and publications and Internet web pages on the subject of medical device EMC;
- **Assess** the electromagnetic environment of the facility (e.g., identify radio transmitters in around the facility) and identify areas where critical medical devices are used;
- **Manage** the electromagnetic environment, RF transmitters and all electrical and electronic equipment, including medical devices, to reduce the risk of medical device electromagnetic interference (EMI) and achieve EMC;
- **Coordinate** the purchase, installation, service, and management of all electrical and electronic equipment used in the facility to achieve EMC;
- **Educate** healthcare facility staff, contractors, visitors, and patients about EMC and EMI and how they can recognize medical device EMI and help minimize associated risks;
- **Establish and implement written policies and procedures** that document the intentions and methods of the healthcare institution for reducing the risk of medical device EMI and achieving EMC;
- **Report** EMI problems to the US FDA MedWatch program and communicate EMI/EMC experiences to colleagues in open forums such as medical/technical publications and conferences.

More information is contained within a comprehensive guidance document for EMC in healthcare facilities, developed, with FDA participation, by the Association for the Advancement of Medical Instrumentation (AAMI):

Technical Information Report (TIR) 18, Guidance on Electromagnetic Compatibility of Medical Devices for Clinical/Biomedical Engineers. AAMI TIR 18-1997. Arlington, Virginia: Association for the Advancement of Medical Instrumentation; 1997.

Caution:

Medical electrical equipment requires special precautions regarding EMC and must be installed and operated according to these instructions. It is possible that high levels of radiated or conducted radio-frequency electromagnetic interference (EMI) from portable and mobile RF communications equipment or other strong or nearby radio-frequency sources, could result in performance disruption of the ultrasound system. Evidence of disruption may include image degradation or distortion, erratic readings, equipment ceasing to operate, or other incorrect functioning. If this occurs, survey the site of disruption, and take the following actions to eliminate the source(s).

- Turn equipment in the vicinity off and on to isolate disruptive equipment.
- Relocate or re-orient interfering equipment.
- Increase distance between interfering equipment and your ultrasound system.
- Manage use of frequencies close to ultrasound system frequencies.
- Remove devices that are highly susceptible to EMI.
- Lower power from internal sources within the facility control (such as paging systems).
- Label devices susceptible to EMI.
- Educate clinical staff to recognize potential EMI-related problems.
- Eliminate or reduce EMI with technical solutions (such as shielding).
- Restrict use of personal communicators (cell phones, computers) in areas with devices susceptible to EMI.
- Share relevant EMI information with others, particularly when evaluating new equipment purchases which may generate EMI.
- Purchase medical devices that comply with IEC 60601-1-2 EMC Standards (3V/meter EMI immunity, limit interference level to 0.0014 V/meter).

ELECTROMAGNETIC COMPATIBILITY (EMC) TABLES

| Guidance and Manufacturer's Declaration - Electromagnetic Emissions | | |
|---|-------------------|---|
| The Vectra® Neo Clinical Therapy System is intended for use in the electromagnetic environment specified below. The customer or the user of the Vectra® Neo Clinical Therapy System should assure that it is used in such an environment. | | |
| Emissions Tests | Compliance | Electromagnetic Environment - Guidance |
| RF emissions CISPR 11 | Group 1 | The Vectra® Neo Clinical Therapy System uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. |
| RF emissions CISPR 11 | Class A | The Vectra® Neo Clinical Therapy System is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes. |
| Harmonic emissions IEC 61000-3-2 | Class A | |
| Voltage fluctuations IEC 61000-3-3 | Complies | |

ELECTROMAGNETIC COMPATIBILITY (EMC) TABLES (CONTINUED)

| Guidance and Manufacturer’s Declaration - Electromagnetic Immunity | | | |
|--|---|---|---|
| <p>The Vectra® Neo Clinical Therapy System is intended for use in the electromagnetic environment specified below. The customer or the user of the Vectra® Neo Clinical Therapy System should assure that it is used in such an environment.</p> | | | |
| Immunity Test | IEC 60601 Test Level | Compliance Level | Electromagnetic Environment - Guidance |
| Electrostatic discharge (ESD) IEC 61000-4-2 | ±6 kV contact ±8 kV air | ±6 kV contact ±8 kV air | <p>Risk assessment on the Vectra Neo Clinical Therapy System indicates the compliance levels claimed are acceptable when ESD-precautionary measures are taken.</p> <p>The Vectra Neo Clinical Therapy System may be susceptible to Electro-Static Discharge (ESD) at greater than ±4 kV when first grasping either the Ultrasound or Laser applicator. In the event of such a discharge, the Vectra Neo Clinical Therapy System may experience communication loss with the installed modules. The Vectra Neo Clinical Therapy System will terminate all active outputs (stim, ultrasound, laser), automatically place the unit in a safe state, and issue an error message 301 or 307.</p> <p>To recover from an error message 301 or 307, turn the unit off and on using the ON/OFF switch located at the top of the display. Once the system restarts, re-initiate all treatments that were interrupted.</p> <p>To prevent Electro-Static Discharge (ESD) at greater than ±4 kV:</p> <ul style="list-style-type: none"> - Grasp and hold the Ultrasound or Laser Applicator prior to starting treatment. If the applicator must be put down prior to completion of treatment, stop the current treatment first and then place the applicator in the holder. - Maintain humidity in the use environment to at least 50% relative humidity. - Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, DJO recommends implementing additional controls to maintain relative humidity to at least 50%. - Communicate these ESD-precautionary procedures to healthcare staff, contractors, visitors and patients. |
| Electrical fast transient/burst IEC 61000-4-4 | ±2 kV for power supply lines ±1 kV for input/output lines | ±2 kV for power supply lines ±1 kV for input/output lines | Mains power quality should be that of a typical commercial or hospital environment. |
| Surge IEC 61000-4-5 | +1 kV differential mode (line to line) +2 kV common mode (line to ground) | ±1 kV differential mode ±2 kV common mode | Mains power quality should be that of a typical commercial or hospital environment. |
| Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11 | <5% U _T (>95% dip in U _T) for 0,5 cycle 40% U _T (60% dip in U _T) for 5 cycles 70% U _T (30% dip in U _T) for 25 cycles <5% U _T (>95% dip in U _T) for 5 sec | <5% U _T (>95% dip in U _T) for 0,5 cycle 40% U _T (60% dip in U _T) for 5 cycles 70% U _T (30% dip in U _T) for 25 cycles <5% U _T (>95% dip in U _T) for 5 sec | Mains power quality should be that of a typical commercial or hospital environment. If the user of the Vectra® Neo Clinical Therapy System requires continued operation during power mains interruptions, it is recommended that the Vectra® Neo Clinical Therapy System be powered from an uninterrupted power supply. |
| Power frequency (50/60 Hz) magnetic field IEC 61000-4-8 | 3 A/m | 3 A/m | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment. |

NOTE: U_T is the AC mains voltage prior to application of the test level.

ELECTROMAGNETIC COMPATIBILITY (EMC) TABLES (CONTINUED)

Guidance and Manufacturer’s Declaration - Electromagnetic Immunity

The Vectra® Neo Clinical Therapy System is intended for use in the electromagnetic environment specified below. The customer or the user of the Vectra® Neo Clinical Therapy System should assure that it is used in such an environment.

| Immunity Test | IEC 60601 Test Level | Compliance Level | Electromagnetic Environment - Guidance |
|-------------------------------|-----------------------------|----------------------------------|---|
| Conducted RF IEC 61000-4-6 | 3 Vrms 150 kHz to 80 MHz | $[V_1] V$, where $V_1 = 3V$ | <p>Portable and mobile RF communications equipment should be used no closer to any part of the Vectra® Neo Clinical Therapy System, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</p> <p>Recommended separation distance</p> $d = \left[\frac{3.5}{V_1} \right] \sqrt{P}$ |
| Radiated RF IEC 61000-4-3 | 3 V/m 80 MHz to 2,5 GHz | $[E_1] V/m$, where $E_1 = 3V/m$ | $d = \left[\frac{3.5}{E_1} \right] \sqrt{P} \quad 80 \text{ MHz to } 800 \text{ MHz}$ $d = \left[\frac{7}{E_1} \right] \sqrt{P} \quad 800 \text{ MHz to } 2,5 \text{ GHz}$ <p>where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey,^a should be less than the compliance level in each frequency range.^b</p> <p>Interference may occur in the vicinity of equipment marked with the following symbol:</p>  |

NOTE 1: At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast, and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Vectra® Neo Clinical Therapy System is used exceeds the applicable RF compliance level above, the Vectra® Neo Clinical Therapy System should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Vectra® Neo Clinical Therapy System .

^b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than $[V_1] V/m$.

ELECTROMAGNETIC COMPATIBILITY (EMC) TABLES (CONTINUED)**Recommended separation distances between portable and mobile RF communications equipment and the Vectra® Neo Clinical Therapy System**

The Vectra® Neo Clinical Therapy System is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Vectra® Neo Clinical Therapy System can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Vectra® Neo Clinical Therapy System as recommended below, according to the maximum output power of the communications equipment.

| Rated maximum output power of transmitter P (W) | Separation distance according to frequency of transmitter d (m) | | |
|--|---|---|--|
| | 150 kHz to 80 MHz $d = \left[\frac{3,5}{V_1} \right] \sqrt{P}$ (where $V_1 = 3V$) | 80 MHz to 800 MHz $d = \left[\frac{3,5}{E_1} \right] \sqrt{P}$ (where $E_1 = 3V/m$) | 800 MHz to 2,5 GHz $d = \left[\frac{7}{E_1} \right] \sqrt{P}$ (where $E_1 = 3V/m$) |
| 0,01 | 0,12 | 0,12 | 0,23 |
| 0,1 | 0,38 | 0,38 | 0,73 |
| 1 | 1,2 | 1,2 | 2,3 |
| 10 | 3,8 | 3,8 | 7,3 |
| 100 | 12 | 12 | 23 |

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies

NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

CALCULATIONS FOR OUTPUT CURRENT

The Vectra® Neo Clinical Therapy System controls total output current and output energy to comply with IEC 60601-2-10. The System also controls total output current density for the selected electrode to comply with FDA requirements. The Vectra® Neo Clinical Therapy System controls output by dynamically changing the range of controllable/selectable parameters, based on the user selection of Intensity, Phase Duration, Frequency, and/or Electrode size.

The following tables provide the calculation and guidance used by the system to dynamically control the output and comply with IEC 60601-2-10 and the 250 mW/cm² power density-limit required by FDA. All calculations are based on Resistive Load = 500 ohms and Accuracy of 10% (1.1).

CC: Constant Current

CV: Constant Voltage

| Waveform | IEC current limit (in mA) | Output current equation [I _{RMS} in mA] | Intensity [I] (in mA (CC), in V (CV)) | Phase Duration [T] (in uSec) | Frequency [F] (in Hz) | Duty Cycle [D] | Number of Pulse [N] |
|----------------------------------|---------------------------|--|---------------------------------------|------------------------------|-----------------------|----------------|---------------------|
| Interferential (F <= 5 kHz) (CC) | 100 | $\frac{I * accuracy}{\sqrt{2}}$ | 0 – 100 | 50 – 250 | 2000 – 10000 | 100% | N/A |
| Interferential (F > 5 kHz) (CC) | 100 | $I * accuracy$ | 0 – 90 | 50 – 250 | 2000 – 10000 | 100% | N/A |
| Interferential (F <= 5 kHz) (CV) | 100 | $\frac{(I/R) * accuracy}{\sqrt{2}}$ | 0 – 64 | 50 – 250 | 2000 – 10000 | 100% | N/A |
| Interferential (F > 5 kHz) (CV) | 100 | $(I/R) * accuracy$ | 0 – 45 | 50 – 250 | 2000 – 10000 | 100% | N/A |
| Premodulated (F <= 5 kHz) (CC) | 100 | $\frac{I * accuracy}{2}$ | 0 – 100 | 50 – 250 | 2000 – 10000 | 66% | N/A |
| Premodulated (F > 5 kHz) (CC) | 100 | $\frac{I * accuracy}{\sqrt{2}}$ | 0 – 100 | 50 – 250 | 2000 – 10000 | 66% | N/A |
| Premodulated (F <= 5 kHz) (CV) | 100 | $\frac{(I/R) * accuracy}{2}$ | 0 – 96 | 50 – 250 | 2000 – 10000 | 66% | N/A |
| Premodulated (F > 5 kHz) (CV) | 100 | $\frac{(I/R) * accuracy}{\sqrt{2}}$ | 0 – 68 | 50 – 250 | 2000 – 10000 | 66% | N/A |
| Russian (CC) | 100 | $\frac{I * accuracy * \sqrt{D}}{\sqrt{2}}$ | 0 – 100 | 200 | 2500 | 50% | N/A |
| Russian (CV) | 100 | $\frac{(I/R) * accuracy * \sqrt{D}}{\sqrt{2}}$ | 0 – 90 | 200 | 2500 | 50% | N/A |
| VMS™ (CC) | 50 | $\frac{I * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 200 | 20 - 1000 | 1 – 200 | 100% | 2 |
| VMS™ (CV) | 50 | $\frac{(I/R) * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 200 | 20 - 1000 | 1 – 200 | 100% | 2 |
| VMS™ Burst (CC) | 50 | $\frac{I * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 200 | 20 – 700 | 1 – 200 | 100% | 6 |
| VMS™ Burst (CV) | 50 | $\frac{(I/R) * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 200 | 20 – 700 | 1 – 200 | 100% | 6 |
| VMS™ FR (CC) | 50 | $\frac{I * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 150 | 20 – 400 | 20 – 80 | 100% | 2 |
| VMS™ FR (CV) | 50 | $\frac{(I/R) * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 150 | 20 – 400 | 20 – 80 | 100% | 2 |

CALCULATIONS FOR OUTPUT CURRENT (CONTINUED)

| Waveform | IEC current limit (in mA) | Output current equation [I _{RMS} in mA] | Intensity [I] (in mA (CC), in V (CV)) | Phase Duration [T] (in uSec) | Frequency [F] (in Hz) | Duty Cycle [D] | Number of Pulse [N] |
|----------------------------------|---------------------------|--|---------------------------------------|------------------------------|-----------------------|----------------|---------------------|
| High Voltage Pulsed Current (CV) | 80 | $\frac{(I/R) * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{D}$ | 0 – 500 | 20 | 10 – 120 | 33.33% | 2 |
| Microcurrent (CC) | 80 | N/A | 0 – 1 | 500 – 500000 | 0.1 – 1000 | 50% | N/A |
| TENS Asymmetrical Biphasic (CC) | 50 | $\frac{I * accuracy * \sqrt{F * (T + 250)}}{1000}$ | 0 – 110 | 20 – 1000 | 1 – 250 | 100% | N/A |
| TENS Asymmetrical Biphasic (CV) | 50 | $\frac{(I/R) * accuracy * \sqrt{F * (T + 250)}}{1000}$ | 0 – 110 | 20 – 1000 | 1 – 250 | 100% | N/A |
| TENS Symmetrical Biphasic (CC) | 50 | $\frac{I * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 80 | 20 – 1000 | 1 – 250 | 100% | 2 |
| TENS Symmetrical Biphasic (CV) | 50 | $\frac{(I/R) * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 80 | 20 – 1000 | 1 – 250 | 100% | 2 |
| DC (Direct Current) (CC) | 80 | $I * accuracy$ | 0 – 72 | N/A | N/A | N/A | N/A |
| HAN-Stimulation | 50 | $\frac{I * accuracy * \sqrt{N} * \sqrt{F} * \sqrt{T}}{1000}$ | 0 – 100 | 180 | 80 | 100% | 2 |

MAXIMUM CURRENT OUTPUT CALCULATIONS BASED ON 250 MW/CM² POWER DENSITY LIMIT

| Electrode Part Number | Dimensions (inch) (shape) | Area (cm ²) | Maximum Power Density (mW/cm ²) | Maximum RMS Current Based on Power Density (mA) | Maximum RMS Current Allowed by System (mA) |
|-----------------------|------------------------------|-------------------------|---|---|--|
| 42045 | 5 × 2.75 (rectangle) | 88.71 | 250 | 210.61 | 90.5 |
| 42159 | 2.5 × 1.5 (ellipse) | 19.00 | 250 | 97.47 | 90.5 |
| 42160 | 4 × 2 (ellipse) | 40.54 | 250 | 142.37 | 90.5 |
| 42170 | 1.25 (circle) | 7.92 | 250 | 62.92 | 62.92 |
| 42171 | 2 (circle) | 20.27 | 250 | 100.67 | 90.5 |
| 42172 | 2.5 × 1.5 (ellipse) | 19.00 | 250 | 97.47 | 90.5 |
| 42173 | 5 × 3 (ellipse) | 76.01 | 250 | 194.95 | 90.5 |
| 42174 | 2 × 2 (square) | 25.81 | 250 | 113.60 | 90.5 |
| 42175 | 3.5 × 2 (rectangle) | 45.16 | 250 | 150.27 | 90.5 |
| 42176 | 2 × 2 (square) | 25.81 | 250 | 113.60 | 90.5 |
| 42177 | 3.5 × 2 (rectangle) | 33.87 | 250 | 130.13 | 90.5 |
| 42178 | 2 × 2 (square) | 25.81 | 250 | 113.60 | 90.5 |
| 42179 | 3.5 × 2 (rectangle) | 45.16 | 250 | 150.27 | 90.5 |
| 42181 | 3.5 × 2 (rectangle) | 45.16 | 250 | 150.27 | 90.5 |
| 42182 | 2 (circle) | 20.27 | 250 | 100.67 | 90.5 |
| 42183 | 2 × 2 (square) | 25.81 | 250 | 113.60 | 90.5 |

CURRENT DENSITY CALCULATIONS FOR DJO ELECTRODES

| Waveform | Electrode Part Number | Dimensions (inch) (shape) | Area (cm ²) | IRMS [mA] | mA/cm ² | 42170* |
|----------------------------------|-----------------------|---------------------------|-------------------------|-----------|--------------------|--------|
| Interferential (F <= 5 kHz) (CC) | 42045 | 5 x 2.75 (rectangle) | 88.7 | 70.7 | 0.8 | 42170* |
| Interferential (F > 5 kHz) (CC) | 42159 | 2.5 x 1.5 (ellipse) | 19.0 | 3.7 | 1.7 | 42170* |
| Interferential (F <= 5 kHz) (CV) | 42160 | 4x2 (circle) | 20.3 | 3.5 | 1.7 | 42170* |
| Interferential (F > 5 kHz) (CV) | 42171 | 2.5 x 1.5 (ellipse) | 19.0 | 3.7 | 1.7 | 42170* |
| Premodulated (F <= 5 kHz) (CC) | 42172 | 2.5 x 1.5 (ellipse) | 19.0 | 3.7 | 1.7 | 42170* |
| Premodulated (F > 5 kHz) (CC) | 42173 | 5 x 3 (ellipse) | 76.0 | 0.9 | 0.9 | 42170* |
| Premodulated (F <= 5 kHz) (CV) | 42174 | 2x2 (square) | 25.8 | 2.7 | 2.7 | 42170* |
| Premodulated (F > 5 kHz) (CV) | 42175 | 3.5 x 2 (rectangle) | 45.2 | 1.6 | 1.6 | 42170* |
| Russian (CC) | 42176 | 2x2 (square) | 25.8 | 3.5 | 3.5 | 42170* |
| Russian (CV) | 42177 | 3.5 x 2 (rectangle) | 33.9 | 2.1 | 2.1 | 42170* |
| WMS™ (CC) | 42178 | 2x2 (square) | 25.8 | 2.7 | 2.7 | 42170* |
| WMS™ (CV) | 42179 | 3.5 x 2 (rectangle) | 45.2 | 1.6 | 1.6 | 42170* |
| WMS™ Burst (CC) | 42181 | 3.5 x 2 (rectangle) | 45.2 | 1.6 | 1.6 | 42170* |
| WMS™ Burst (CV) | 42182 | 2 (circle) | 20.3 | 3.5 | 3.5 | 42170* |
| WMS™ FR (CC) | 42183 | 2x2 (square) | 25.8 | 2.7 | 2.7 | 42170* |
| WMS™ FR (CV) | 42184 | 2x2 (square) | 25.8 | 2.7 | 2.7 | 42170* |
| High Voltage Pulsed Current (CV) | 42185 | 2x2 (square) | 25.8 | 2.7 | 2.7 | 42170* |
| Microcurrent (CC) | 42186 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| TENS Asymmetrical Biphasic (CC) | 42187 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| TENS Asymmetrical Biphasic (CV) | 42188 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| TENS Symmetrical Biphasic (CC) | 42189 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| TENS Symmetrical Biphasic (CV) | 42190 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| DC (Direct Current) (CC) | 42191 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |
| HAN-Stimulation | 42192 | 2x2 (square) | 25.8 | 0.0 | 0.0 | 42170* |

* Due to the area of electrode 42170, the IRMS value is automatically reduced by the Vectra® Neo Clinical Therapy System.

Caution: DO NOT use electrodes with an active area less than 7.92 cm², as there will be a risk of suffering a burn injury. Caution should always be exercised with current densities more than 2 mA/cm².

ELECTRODE SIZE SELECTION

The Vectra® Neo Clinical Therapy System allows the user to select an electrode size from three different options, which allows the System to more accurately control the output current. Please refer to following table for the selection guide related to different DJO electrode part numbers.

| Device Electrode Selection | DJO Electrode Part Number |
|----------------------------|--|
| 1.25" round | 42170 |
| 2" round | 42159*, 42171, 42172*, 42182 |
| 2" square | 42045*, 42160*, 42173*, 42174, 42175*, 42176, 42177*, 42178, 42179*, 42181*, 42183 |

* The maximum output current is limited by the System to a value lower than what the electrode can deliver and therefore for the purposes of electrode selection in modality setup, all elliptical electrodes are considered as "round" and all rectangular electrodes are considered as "square". Electrodes are grouped in this manner for ease of user selection.



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