

# **INSTRUCTION MANUAL**

CAUTION: United States Federal Law restricts this device to sale by or on the order of a physician



# This manual is valid for the Quattro<sup>™</sup> II

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Declaration of conformity:

Current Solutions<sup>™</sup>, LLC. declares that the Quattro<sup>™</sup> II complies with following normative documents:

IEC60601-1, IEC60601-1-2, IEC60601-2-10, ISO 7010 ISO14971, ISO10993-1, ISO10993-5, ISO10993-10

Quattro<sup>®</sup> II

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1. FOREWORD	
1.1 General information	Thank you for purchasing the Quattro <sup>™</sup> II. The microprocessor controlled Quattro <sup>™</sup> II provides interferential (4-pole), premodulated (2-pole interferential), medium frequency (Russian), EMS and TENS waveform. You can choose between several different amplitude modulation options. The interferential and premodulated modes offer frequency modulation as well as a static frequency option.
1.2 Introduction to This Manual	This manual has been written for the users of Quattro <sup>™</sup> II. It contains general information on the operation, precautionary practices, and maintenance information. In order to maximize its 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.

#### 2. SAFETY INFORMATION

#### 2.1 Precaution definitions

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 is as follows;

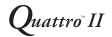


**Caution:** Text with a "CAUTION" indicator will explain possible safety infractions that could have the potential to cause minor to moderate injury or damage to equipment.



**Warning:** Text with a "WARNING" indicator will explain 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.



#### 2.2 CAUTION

## Caution:

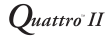
- Federal law (USA) restricts this device to sale by or on the order of a physician.
- Keep yourself informed of the contraindications.
- Read, understand, and practice the warnings, cautions and operating instructions. Know the limitations and hazards associated with using any device. Observe the precautionary and operational decals placed on the unit. Always follow the operating instructions prescribed by your healthcare practitioner.
- The long-term effects of chronic electrical stimulation are unknown. Electrical stimulation devices do not have any curative value.
- Stimulation is not effective for pain of central origin, including headache.
- Stimulation is not a substitute for pain medications and other pain management therapies.
- This stimulator not intended for unattended, personal use by patients who have noncompliant, emotionally disturbed, dementia, or low IQ.
- Stimulation delivered by this device may be sufficient to cause electric shock. Electrical current of this magnitude must not flow through the thorax or across the chest because it may cause a cardiac arrhythmia.
- Do not place electrodes on the front of the throat as spasm of the Laryngeal and Pharyngeal muscle may occur. Stimulation over the carotid sinus (neck region) may close the airways, make breathing difficult, and may have adverse effects on the heart rhythm or blood pressure.
- Do not place electrodes on your head or at any sites that may cause the electrical current to flow transcerebrally (through the head).
- Patients with heart disease, epilepsy, cancer or any other health condition should not use this device without first consulting a physician.
- Some patients may experience skin irritation or hypersensitivity due to the electrical stimulation or electrical conductive medium (gel).
- Use caution when the patient has a tendency to bleed internally, such as following an injury or fracture;
- Use caution following recent surgical procedures when stimulation may disrupt the patient's healing process;
- Use caution if stimulation is applied over the menstruating or pregnant uterus;
- Use caution if stimulation is applied over areas of skin that lack normal sensation.
- Electrode placement and stimulation settings should be based on the guidance of prescribing practitioner.
- Effectiveness is highly dependent upon patient selection by a practitioner qualified in the management of pain patients.

- The electrodes are only to be placed on healthy skin. Avoid skin irritation by ensuring that good contact is achieved between electrodes and skin.
- If the stimulation levels are uncomfortable or become uncomfortable, reduce the stimulation Intensity to a comfortable level.
- Never use the device in rooms where aerosols (sprays) are used or pure oxygen is being administered.
- Do not use this device at the same time as other equipment which sends electrical pulses to your body.
- Do not confuse the electrode cables and contacts with your headphones or other devices, and do not connect the electrodes to other devices.
- Turn the device off before applying or removing electrodes.
- Electrical stimulators should be used only with the leads and electrodes recommended for use by the manufacturer.
- Keep the stimulator out of reach of children.

#### 2.3 WARNING

# A Warning

- This device should be used only under the continued supervision of a licensed physician.
- TENS is a symptomatic treatment and, as such, suppresses the sensation of pain, which would otherwise serve as a protective mechanism.
- Safety has not been established for the use of therapeutic electrical stimulation during pregnancy. Do not use during pregnancy unless directed by your physician.
- Electrical stimulation is not effective for pain of central origin.
- Electronic monitoring equipment (such as ECG monitors and ECG alarms) may not operate properly when electrical stimulation is in use.
- Stimulation should not be applied over the carotid sinus nerves, particularly in patients with a known sensitivity to the carotid sinus reflex.
- Do not apply stimulation over the patient's neck because this could cause severe muscle spasms resulting in closure of the airway, difficulty in breathing, or adverse effects on heart rhythm or blood pressure.
- Stimulation should not be applied transthoracically in that the introduction of electrical current into the heart may cause cardiac arrhythmias.
- Stimulation should not take place while the user is connected to high-frequency surgical equipment, it may cause burn injuries on the skin under the electrodes, as well as problems with the stimulator.
- Do not use the stimulator in the vicinity of shortwave or microwave



therapy equipment, since this may affect the output power of the stimulator.

- Never use in environments with high humidity such as in the bathroom or when having a bath or shower.
- Do not apply stimulation while the patient is sleeping;
- Caution should be used in applying electrical stimulation to patients suspected of having heart disease. Further clinical data is needed to show there are no adverse results.
- Never use near the heart. Stimulation electrodes should never be placed anywhere on the front of the thorax (marked by ribs and breastbone), but above all not on the two large pectoral muscles. Here it can increase the risk of ventricular fibrillation and lead to cardiac arrest.
- Electrodes should not be placed over the eyes, in the mouth, near the genitals or internally.
- Keep electrodes separate during treatment, electrodes in contact with other could result in improper stimulation or skin burns.
- Apply stimulation only to normal, intact, clean, healthy skin.
- Avoid use over or near bone growth centers until bone growth is complete.

# 2.4 Danger 7

## Danger

Patients with an implanted neurostimulation device must not be treated with or be in close proximity to any shortwave diathermy, microwave 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."

2.5 Skin irritation, inflammation, and electrode burns beneath the electrodes are potential adverse reactions.

#### 2.6 Contraindications

- Do not use this device on patients who have a cardiac pacemaker, implanted defibrillator, or other implanted metallic or electronic device, because this may cause electric shock, burns, electrical interference, or death.
- Do not use this device on patients whose pain syndromes are undiagnosed.
- This device should not be used when cancerous lesions are present in the treatment area.
- This device should not be used over or near bone growth centers until

bone growth is complete.

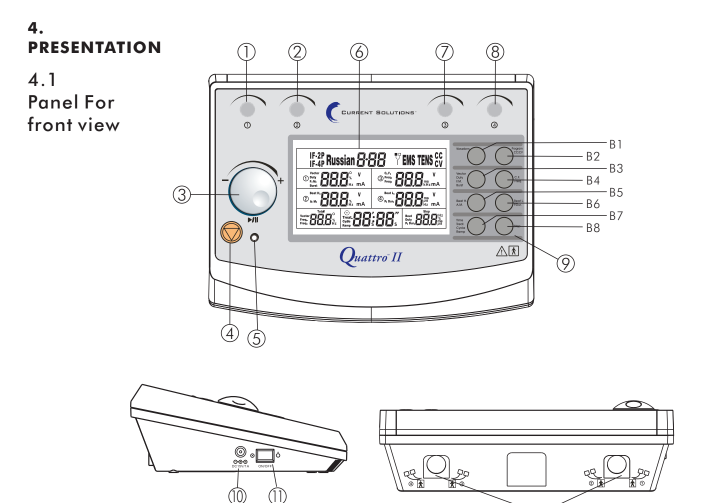
- This device should not be used over a healing fracture.
- This device should not be used over or applied to the eye.
- This device should not be used over a pregnant uterus.
- This device should not be used on ischemic tissues in individuals with vascular disease where the blood supply would be unable to follow the increase in metabolic demand and tissue necrosis might result.

#### 3. INDICATIONS FOR USE

# For TENS, EMS, Russian, Interferential and premodulated waveforms:

- Relaxation of muscle spasms
- Prevention disuse atrophy
- Increase local blood circulation
- Muscle re-education
- Maintaining or increasing range of motion
- Symptomatic relief and management of chronic, Intractable pain
- Post-traumatic acute pain
- Post-surgical acute pain

**J**uattro<sup>®</sup> II



- 1) Adjust the output intensity of channel 1.
- 2) Adjust the output intensity of channel 2.
- 3) Parameters control knob and pause button.
- 4) Stop treatment button.

(11)

- 5) Power indicator.
- 6) LCD display: Shows the current information of the device.

(12)

- 7) Adjust the output intensity of channel 3.
- 8) Adjust the output intensity of channel 4.
- 9) Eight parameters selection buttons, see below for details:
  - B1: Toggle the therapeutic waveform.
  - B2: Toggle the therapeutic program and select the output mode (CC/CV).
  - B3: Toggle the parameter Vector/Duty/F.M./Burst
  - B4: Toggle the parameter Freq./C.F.
  - B5: Toggle the parameter Beat H./A.M.
  - B6: Toggle the parameter Beat L./P.Dur.
  - B7: Toggle the parameter Treat./Cycle/Ramp time
  - B8: Step button

## Remark:

- CC Constant current output mode.
- CV Constant voltage output mode.
- F.M. Frequency Modulation
- Burst— Burst Frequency
- Freq. Frequency
- C.F. Carrier Frequency
- Duty Duty Cycle for Russian waveform
- Beat H. Sweep High Beat Frequency
- A.M. Amplitude Modulation
- Beat L. Sweep High Beat Frequency
- P.Dur. Pulse Duration
- Treat. Treatment time
- Cycle—Cycle time
- Ramp— Ramp time
- 10) Adapter receptacle
- 11) ON/OFF switch
- 12) Output connector: connect with connector of cable



Symbol definitions	Symbol definitions				
IF-4P IFC-Interferential (Traditional 4 Pole)	IF-2P Premodulated (Traditional 2 Pole IFC)				
Electrical stimulation	🕥 Time indicator				
D Electrical output channel 1 indicator	Electrical output channel 2 indicator				
3 Electrical output channel 3 indicator	Electrical output channel 4     indicator				
<b>CC</b> Constant current control	<b>CV</b> Constant Voltages control				

## 5. INSTALLATION

5.1 Remove the equipment and all accessories from shipping carton and box. Visually check if there is any damage or missing parts or accessories. If yes, please report to local dealer or retailer where you purchase this unit. Your Quattro<sup>TM</sup> II equipment contains the following accessories.



	Part	Quantity
1	Rubber Electrodes,60x90mm	4pcs
2	Rubber Electrodes,70x110mm	4pcs
3	Electrode Sponges,70x100mm	4pcs
4	Electrode Sponges,80x120mm	4pcs
5	Self-adhesive Electrodes,50x50mm	8pcs
6	Self-adhesive Electrodes,50x100mm	8pcs
7	Elastic Wrap,75x1200mm	2pcs
8	Elastic Wrap ,75x600mm	2pcs
9	Electrode wires (black/red)	4pcs
10	Adapter 100-240V/50-60Hz	lpc
11	Power cord	lpc
12	Connector of cable	2pcs
13	Use Manual	lpc

5.2

• Connect the power cord to the power adapter.

#### Connection • Connect the power adapter to the device connector. • Connect the power adapter to a wall socket.

of the power adapter

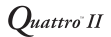
#### **Caution:**

- Prior to connecting this apparatus to the power supply, check that the voltage and frequency stated on the rating label match with the available power supply.
- The power adapter is a part of the supply circuit on which the device's safety partly depends. The approvals for Quattro<sup>™</sup> II are only valid if used in combination with this type of adapter.

5.3 Switch on the device, using ON/OFF switch (1)).

Switching on

5.4 • Switch off the device by switching the ON/OFF switch from [] to Switching off [O] position. and • Pull out the power adapter from the wall socket. disconnect • Pull out the power adapter from device. power adapter



### 6. OPERATION

6.1 Measures with regard to treatments

6.1.1 Electrotherapy Before the treatment

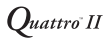
6.1.2 Electrode Placement

- Ensure there are no contraindications to treatment.
- ElectrotherapyInspect the treatment area skin seriously for any abrasions,Beforeinflammation, surface veins etc.
  - Clean the skin of the treatment area with soap or alcohol (70%).
  - If the skin is hairy, shaving can get optimal treatment.
  - Test the heat sensibility of the treatment area.
  - 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.
  - Follow electrode manufacturer instructions.
  - To avoid skin irritation due to high current density, do not use electrodes smaller in surface area than 25cm<sup>2</sup> self-adhesive electrode.

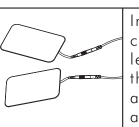
# ⚠ Caution

- Keep electrodes separated during treatment. Electrodes in contact with each other could result in improper stimulation or skin burns.
- 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.
- Powered muscle stimulators should be used only with the leads and electrodes recommended by the manufacturer.

6.1.3 This device is supplied with 8 pieces 50mm×50mm and 8 pieces 50mm×100mm adhesive electrodes. You can select the right adhesive electrodes according to treatment area and output current density. It is recommended that manufacturer's 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. Properly dispose of used Electrodes upon completion of the therapy session.

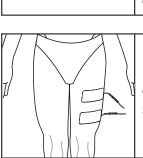


Electrode Instructions Connecting Lead Wires



Insert the lead with the Red (+) electrode connector into one adhesive Electrode. Insert the lead with the Black (-) electrode connector into the other electrode. Make certain the lead wires are seated completely into the electrodes, there are no bare metal of the pins exposed.

Securing Electrodes



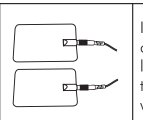
Remove the adhesive Electrodes from the protective backing and apply to the treatment area as prescribed. Ensure the entire electrode surface is in contact with patient skin by pressing into place.

#### 6.1.4. Rubber electrodes

If used for delivery of electrotherapy, there are two conductive mediums for you to select, the first one is use electrode sponges as conductive mediums, another is use other conductive medium such as Transmission Gel.

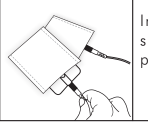
These Rubber Electrodes should be secured to the treatment area using the Nylon Wraps shipped with the Therapy System.

Reusable rubber Electrodes Connecting Lead Wires



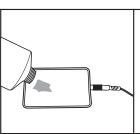
Insert the lead with the Red (+) electrode connector into one rubber electrode. Insert the lead with the Black (-) electrode connector into the other rubber electrode. Make certain the lead wires are seated completely into the electrodes.

Conductive Medium 1

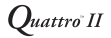


Inserted the Rubber Electrodes into the electrode sponges moistened with distilled water prior to placement on the patient.

Conductive Medium 2



Liberally apply Transmission Gel to electrode prior to placement on patient. Please note: Please purchase the Transmission gel with CE mark or FDA approved.



Securing Electrodes		Use Nylon Wrap to secure each rubber electrode in position on the patient.
6.2 Quick Set-up for Electrical Stimulation	⊙∏ċ	<ol> <li>In order to turn on the device, please press ON/OFF switch to [⊙] icon which is located on the side of the device</li> <li>When you turn the Ougsttre<sup>™</sup>II on the</li> </ol>
	IF4P         P:0         I         V         CC           "***"         YS"         """         YUD           "***"         I         IO         """         IOO           "***"         I         IO         """         IOO           "***"         ISTON"         """         IOO	<ol> <li>When you turn the Quattro<sup>™</sup>II on, the device will get down to self- check about 6~8 seconds, and then the default parameters are displayed the last treatment mode.</li> </ol>
		<ul> <li>3. Connect the electrode wires to the cable; please note the color of the wires and the color marks on the cable, they should be corresponding.</li> <li>Caution: If you want to use 4 channels, please connect all electrode wires to two cables.</li> </ul>
		4. Two connectors are individual, each connector output two channels through a cable. So you can plug the cable(s) into one or two output conne- ctors (12)connectors)according to patient's need.
	A A A	5. Connect the electrodes to electrode wires.
		<ol> <li>6. Place the electrodes on the patient according to section 6.1.</li> </ol>
	Waveform	7. There are 5 therapeutic waveforms for you to select. Press the "Waveform" button to toggle the therapeutic waveform, and then rotating the Parameters control knob (③) to select waveform like IF-4P, IF-2P, TENS, EMS and Russian.

Program CC/CV	8. Each therapeutic waveform has 10 programs. The details parameters for each program please refer to section 6.3 in this manual. Press the "Program" button to toggle the therapeutic program, and then rotating the Parameters control knob to select the therapeutic programs in corresponding waveform.
CC	9. Press "B2" Program button to select "CC" or "CV" control mode.
	10. Adjust the output intensity and start electrical treatment that you are using by rotating the output intensity adjustable knobs on the control panel. 0.5mA/step or 0.5V/step.
1 <b>1 1 1 1 1 1 1 1 1 1</b>	<ul> <li>11. For safety using, load detection was designed in this device after the output intensity surpass 10.0mA/10.0V. If there are no electrodes stuck on patient' skin, an alarm buzzer sound will appear and the intensity value flashing.</li> </ul>
	<ol> <li>Press the " <sup>©</sup> " button to stop treatment if any emergency happened.</li> </ol>
++	13. Press the "▶/II " button to pause treatment; you can press it again to restart the treatment.

6.3 Each therapeutic waveform has 10 programs. They have default
 Programs
 Programs
 Each therapeutic waveform has 10 programs. They have default
 parameters for each program, but you can set and save the parameters
 according to patient's need. the default parameters for each program
 please refer to below:

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4-Pole Interferential preset	Waveform	Prog- ram	Phase	CC/ CV		Vector (Manual)	C.F.	Beat. H	Beat. L	Treat. Time
programs			1	СС	0	45°	4kHz	110Hz	100Hz	15min
		1	2	СС	0	45°	4kHz	110Hz	100Hz	Omin
			3	СС	0	45°	4kHz	110Hz	100Hz	Omin
			1	СС	0	45°	4kHz	150Hz	100Hz	10min
		2	2	СС	0	45°	4kHz	150Hz	100Hz	Omin
			3	СС	0	45°	4kHz	150Hz	100Hz	Omin
			1	СС	0	45°	4kHz	50Hz	50Hz	15min
		3	2	СС	0	45°	4kHz	50Hz	50Hz	Omin
			3	СС	0	45°	4kHz	50Hz	50Hz	Omin
			1	СС	0	45°	4kHz	150Hz	90Hz	15min
		4	2	СС	0	45°	4kHz	150Hz	90Hz	Omin
	Interferential Traditional		3	СС	0	45°	4kHz	150Hz	90Hz	Omin
		5	1	СС	0	45°	4kHz	110Hz	100Hz	15min
			2	СС	0	45°	4kHz	110Hz	100Hz	Omin
			3	СС	0	45°	4kHz	110Hz	100Hz	Omin
	(4P)		1	CC	0	45°	4kHz	110Hz	100Hz	15min
	IF-4P	6	2	СС	0	45°	4kHz	110Hz	100Hz	15min
	11 - 41		3	СС	0	45°	4kHz	110Hz	100Hz	15min
			1	СС	0	45°		110Hz		
		7	2	СС	0	45°	4kHz	110Hz	100Hz	15min
			3	СС	0	45°		110Hz		
			1	СС	0	45°		110Hz		
		8	2	СС	0	45°		110Hz		
			3	СС	0	45°		110Hz		
			1	CC	0	45°		110Hz		
		9	2	СС	0	45°		110Hz		
			3	CC	0	45°		110Hz		
			1	CC	0	45°		110Hz		
		10	2	СС	0	45°		110Hz		
			3	СС	0	45°	4kHz	110Hz	100Hz	15min

2-Pole Interferential	Waveform	Prog- ram	Phase	CC/ CV	C.F.	Beat. H	Beat. L	Cycle	Treat. Time
programs			1	СС	2500Hz	110Hz	100Hz	continuous	15min
		1	2	СС	2500Hz	110Hz	100Hz	continuous	Omin
			3	СС	2500Hz	110Hz	100Hz	continuous	Omin
			1	СС	2500Hz	150Hz	100Hz	continuous	10min
		2	2	СС	2500Hz	150Hz	100Hz	continuous	Omin
			3	СС	2500Hz	150Hz	100Hz	continuous	Omin
			1	СС	2500Hz	50Hz	50Hz	continuous	15min
		3	2	СС	2500Hz	50Hz	50Hz	continuous	Omin
			3	СС	2500Hz	50Hz	50Hz	continuous	Omin
			1	СС	2500Hz	150Hz	90Hz	continuous	15min
		4	2	СС	2500Hz	150Hz	90Hz	continuous	Omin
			3	СС	2500Hz	150Hz	90Hz	continuous	Omin
	IFC Premod. (2P)		1	СС	2500Hz	110Hz	100Hz	continuous	15min
		5	2	СС	2500Hz	110Hz	100Hz	continuous	Omin
	IF-2P		3	СС	2500Hz	110Hz	100Hz	continuous	Omin
		6	1	СС	2500Hz	110Hz	100Hz	continuous	15min
			2	СС	2500Hz	110Hz	100Hz	continuous	15min
			3	СС	2500Hz	110Hz	100Hz	continuous	15min
			1	СС	2500Hz	110Hz	100Hz	continuous	15min
		7	2	СС	2500Hz	110Hz	100Hz	continuous	15min
			3	СС	2500Hz	110Hz	100Hz	continuous	15min
			1	СС	2500Hz	110Hz	100Hz	continuous	15min
		8	2	СС	2500Hz	110Hz	100Hz	continuous	15min
			3	СС	2500Hz	110Hz	100Hz	continuous	15min
			1	СС	2500Hz	110Hz	100Hz	continuous	15min
		9	2	СС	2500Hz	110Hz	100Hz	continuous	15min
			3	СС	2500Hz	110Hz	100Hz	continuous	15min
			1	СС	2500Hz	110Hz	100Hz	continuous	15min
		10	2	СС	2500Hz	110Hz	100Hz	continuous	15min
			3	СС	2500Hz	110Hz	100Hz	continuous	15min

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TENS programs	Waveform	Prog- ram	Phase	CC/ CV	F.M.	Burst	Freq	A.M.	P. Dur.	Cycle	Treat. Time
-			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		1	2	СС	0	0	120Hz	0%	70µs	continuous	Omin
			3	СС	0	0	120Hz	0%	70µs	continuous	Omin
			1	СС	0	0	200Hz	0%	60µs	continuous	20min
		2	2	СС	0	0	200Hz	0%	60µs	continuous	Omin
			3	СС	0	0	200Hz	0%	60µs	continuous	0min
			1	СС	0	0	10Hz	0%	180µs	continuous	14min
		3	2	СС	0	0	10Hz	0%	180µs	continuous	Omin
			3	СС	0	0	10Hz	0%	180µs	continuous	Omin
			1	СС	0	0	80Hz	0%	100µs	continuous	30min
		4	2	СС	0	0	80Hz	0%	100µs	continuous	Omin
	TENO		3	СС	0	0	80Hz	0%	100µs	continuous	Omin
	TENS		1	СС	50Hz	0	180Hz	0%	30µs	continuous	16min
		5	2	СС	50Hz	0	180Hz	0%	30µs	continuous	Omin
			3	СС	50Hz	0	180Hz	0%	30µs	continuous	Omin
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		6	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		7	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		8	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		9	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
		10	1	СС	0	0	120Hz	0%	70µs	continuous	14min
			2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min

EMS programs	Waveform	Prog- ram	Phase	CC/ CV	F.M.	Burst	Freq	A.M.	P. Dur.	Cycle	Treat. Time
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		1	2	СС	0	0	120Hz	0%	70µs	continuous	Omin
		3	СС	0	0	120Hz	0%	70µs	continuous	Omin	
			1	СС	0	0	200Hz	0%	60µs	continuous	20min
		2	2	СС	0	0	200Hz	0%	60µs	continuous	Omin
			3	СС	0	0	200Hz	0%	60µs	continuous	Omin
			1	СС	0	0	10Hz	0%	180µs	continuous	20min
		3	2	СС	0	0	10Hz	0%	180µs	continuous	Omin
			3	СС	0	0	10Hz	0%	180µs	continuous	Omin
			1	СС	0	0	80Hz	0%	100µs	continuous	30min
		4	2	СС	0	0	80Hz	0%	100µs	continuous	Omin
			3	СС	0	0	80Hz	0%	100µs	continuous	Omin
			1	СС	50Hz	0	180Hz	0%	30µs	continuous	16min
	EMO	5	2	СС	50Hz	0	180Hz	0%	30µs	continuous	Omin
	EMS		3	СС	50Hz	0	180Hz	0%	30µs	continuous	Omin
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		6	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		7	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
			1	СС	0	0	120Hz	0%	70µs	continuous	14min
		8	2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min
		1	СС	0	0	120Hz	0%	70µs	continuous	14min	
		9	2	СС	0	0	120Hz	0%	70µs	continuous	14min
		3	СС	0	0	120Hz	0%	70µs	continuous	14min	
		10	1	СС	0	0	120Hz	0%	70µs	continuous	14min
			2	СС	0	0	120Hz	0%	70µs	continuous	14min
			3	СС	0	0	120Hz	0%	70µs	continuous	14min

Ramp

1s

ls

Treat.

Time

10min

0min

Russian programs	Waveform	Prog- ram	Phase	CC/ CV	C. F.	Freq.	Duty	Cycle
			1	СС	2500Hz	50Hz	50%	10s/10s
		1	2	СС	2500Hz	50Hz	50%	10s/10s
			3	СС	2500Hz	50Hz	50%	10s/10s
			1	СС	2500Hz	50Hz	50%	4s/12s
		2	2	СС	2500Hz	50Hz	50%	4s/12s
			3	СС	2500Hz	50Hz	50%	4s/12s
			1	СС	2500Hz	50Hz	50%	4s/12s
		3	2	СС	2500Hz	50Hz	50%	4s/12s
			3	СС	2500Hz	50Hz	50%	4s/12s
			1	СС	2500Hz	50Hz	50%	10s/10s
		4	2	СС	2500Hz	50Hz	50%	10s/10s
			3	СС	2500Hz	50Hz	50%	10s/10s
			1	СС	2500Hz	50Hz	50%	5s/5s
	Russian	5	2	СС	2500Hz	50Hz	50%	5s/5s
			3	СС	2500Hz	50Hz	50%	5s/5s
			1	СС	2500Hz	50Hz	50%	10s/10s
		6	2	СС	2500Hz	50Hz	50%	10s/10s
			3	СС	2500Hz	50Hz	50%	10s/10s
			1	СС	2500Hz	50Hz	50%	10s/10s
		7	2	CC	2500Hz	50Hz	50%	10s/10s

		3	CC	2500Hz	50Hz	50%	10s/10s	ls	Omin
	2	1	CC	2500Hz	50Hz	50%	4s/12s	1s	10min
		2	CC	2500Hz	50Hz	50%	4s/12s	ls	Omin
		3	CC	2500Hz	50Hz	50%	4s/12s	ls	Omin
	3	1	CC	2500Hz	50Hz	50%	4s/12s	1s	10min
		2	CC	2500Hz	50Hz	50%	4s/12s	1s	Omin
		3	CC	2500Hz	50Hz	50%	4s/12s	ls	Omin
		1	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
	4	2	CC	2500Hz	50Hz	50%	10s/10s	]s	Omin
		3	CC	2500Hz	50Hz	50%	10s/10s	ls	Omin
		1	CC	2500Hz	50Hz	50%	5s/5s	]s	20min
sian	5	2	CC	2500Hz	50Hz	50%	5s/5s	1s	Omin
		3	CC	2500Hz	50Hz	50%	5s/5s	ls	Omin
		1	CC	2500Hz	50Hz	50%	10s/10s	]s	10min
	6	2	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		3	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
	7	1	CC	2500Hz	50Hz	50%	10s/10s	1s	10min
		2	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		3	CC	2500Hz	50Hz	50%	10s/10s	1s	10min
		1	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
	8	2	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		3	CC	2500Hz	50Hz	50%	10s/10s	1s	10min
	9	1	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		2	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		3	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
		1	СС	2500Hz	50Hz	50%	10s/10s	ls	10min
	10	2	СС	2500Hz	50Hz	50%	10s/10s	ls	10min
		3	CC	2500Hz	50Hz	50%	10s/10s	ls	10min
									21

#### 6.4 Each stimulation set-up procedure

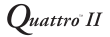
6.4.1 4-Pole Interferentia Stimulation Set-up Procedure

⊙∏Ò	<ol> <li>In order to turn on the device, please press ON/OFF switch to [⊙] icon which is located on the side of the device.</li> </ol>
PO         V         CC           Very         45°         5°         400mm           berk         1 10mm         berth         100mm           J         100mm         15'000"         31'	<ol> <li>When you turn the Quattro<sup>™</sup> II, the device will get down to self- check about 6~8 seconds, and then the default parameters are displayed the last treatment mode.</li> </ol>
IF-4P	3. Press "B1" Waveform button to toggle the therapeutic waveform, then rotating the parameters control knob (③) to select "IF-4P" waveform.
P-[] {	<ul> <li>4. Press "B2" Program button to toggle the therapeutic program, and then rotating the parameters control knob (③) to select the therapeutic programs from PO1 to P10. Each program has 3 treatment phases, you can set and save the parameters according to patient's need.</li> <li>5. Press "B2" Program button to select "CC" or</li> </ul>
UU	"CV" control mode.
<u>™</u> 15′00″ 1	6. There are two modes in Quattro <sup>™</sup> II, press and hold ""B8" button to switch Normal mode and Professional mode. In professional mode, each program has 3 treatment phase, the LCD display like left figure. Rotating the parameters control knob (③) to select phase program from 1 to 3 when the device enter into professional mode. The parameters of each phase program can be set according to following methods.
Vector	7. Press "B3" button to toggle Vector parameter, then rotating the parameters control knob (③) to set the vector (manual) parameter from 0°to 90°, 15°/step.
Vector %	8. Press "B3" button again, the vector parameter change to auto mode, the LCD display "0%" like left figure. rotating the parameters control knob (③) to set the vector (auto) parameter from 0 % to 100%, 20%/step.
	IF-4P       P·O I V       CC         WT       45°       Sr       40         IF-4P       IF-4P       IF-4P         CCC       CC       CC         Image: IS' 000"       Y       Y         IF-4P       IF-4P       IF-4P         Vector       IF-500"       Y         Image: IS' 000"       Y       Y         IF-4P       IF-4P       IF-4P         IF-4P       IF-4P       <



Beat H.	<ol> <li>Press "B5" button to toggle Beat H. parameter, then rotating the parameters control knob (③) to set the parameter from (Beat. L) Hz to 150Hz, 1Hz/step.</li> </ol>	
Beat L.	10. Press "B6" button to toggle Beat L. parameter, then rotating the adjust parameters contorl knob (③) to set the parameter from 1Hz to (Beat. H)Hz, 1Hz/step.	
• Treat. 5'00"	11. Press "B7" button to toggle Treat. time parameter, then rotating the parameters control knob (③) to set the treatment time from 1 min to 60min, 1 min/step.	
0	12. Stick the electrodes on the patient. You will need two electrodes for each channel, four in total.	
1	13. Adjust the output intensity and start electrical treatment that you are using by rotating the output intensity adjustable knobs on the control panel. 0.5mA/step or 0.5V/step. For safety using, load detection was designed in this device after the output intensity surpass 10.0mA/10.0V. If there are no electrodes stuck on patient' skin, an alarm buzzer sound will appear and the intensity value flashing.	
	<ol> <li>Press the "♥" button to stop treatment if any emergency happened.</li> </ol>	
	15. Press the " ►/II " button to pause treatment; you can press it again to restart the treatment.	

6.4.2 2-Pole Interferential Stimulation	⊙∏ċ	<ol> <li>In order to turn on the device, please press ON/OFF switch to[⊙] icon which is located on the side of the device.</li> </ol>
Set-up Procedure	IF-2P         P-0         1         Y         CC           """         45°         ""         2.5           """         1         10         ""         100           """         1         10         ""         100           """         1         10         ""         100	<ol> <li>When you turn the Quattro<sup>™</sup> II on, the device will get down to self- check about 6~8 seconds, and then the default parameters are displayed the last treatment mode.</li> </ol>
	IF-2P	3. Press "B1" Waveform button to toggle the therapeutic waveform, then rotating the parameters control knob (③) to select " <b>IF-2P</b> " waveform.
	₽-0 ¦	4. Press "B2" Program button to toggle the therapeutic program, and then rotating the parameters control knob (③) to select the therapeutic programs from P01 to P10. Each program has 3 treatment phases, you can set and save the parameters according to patient's need.
	CC	5. Press "B2" Program button to select "CC" or "CV" control mode.
	<u>™</u> 15′00″ "i	6. There are two modes in Quattro <sup>™</sup> II, press and hold "B8" button to switch Normal mode and Professional mode. In professional mode, each program has 3 treatment phase, the LCD display like left figure. Rotating the parameters control knob (③) to select phase program from 1 to 3 when the device enter into professional mode. The parameters of each phase program can be set according to following methods.
	Beat H.	8. Press "B5" button to toggle Beat H. parameter, then rotating the parameters control knob (③) to set the parameter from (Beat. L) Hz to 150Hz, 1Hz/step.
	Beat L.	9. Press "B6" button to toggle Beat L. parameter, then rotating the parameters control knob (③) to set the parameter from 1Hz to (Beat. H)Hz, 1Hz/step.
	<sup></sup> € Treat. <b>15'00</b> "	10. Press "B7" button to toggle Treat. time parameter, then rotating the parameters control knob (③) to set the treatment time from 1 min to 60min, 1 min/step.



⊙ Cycle S S	11. Press "B7" button again to toggle Cycle time parameter, then rotating the parameters control knob (③) to select the cycle time from "-/-(continuous)", "5/5", "4/12", "10/10", "10/20", "10/30" and "10/50".
	12. Stick the electrodes on the patient. You can use one or two channel as your needs.
	13. Adjust the output intensity and start electrical treatment that you are using by rotating the output intensity adjustable knob on the control panel. 0.5mA/step or 0.5V/ step. For safety using, load detection was designed in this device after the output intensity surpass 10. 0mA/10. 0V. If there are no electrodes stuck on patient' skin, an alarm buzzer sound will appear and the intensity value flashing.
	<ol> <li>Press the "<sup>©</sup> button to stop treatment if any emergency happened.</li> </ol>
	15. Press the " ►/II " button to pause treatment; you can press it again to restart the treatment.

6.4.3 TENS and EMS	⊙∏ċ	<ol> <li>In order to turn on the device, please press ON/OFF switch to [ ⊙ ] icon which is located on the side of the device.</li> </ol>
Stimulation Set-up Procedure	POINT         TENS <sup>CC</sup> ···<	<ol> <li>When you turn the Quattro<sup>™</sup> II on, the device will get down to self- check about 6~8 seconds, and then the default parameters are displayed the last treatment mode.</li> </ol>
	EMS / TENS	3. Press "B1" Waveform button to toggle the therapeutic waveform, then rotating the parameters control knob (③) to select TENS or EMS waveform.
	P-[] {	4. Press "B2" Program button to toggle the therapeutic program, and then rotating the parameters control knob (③) to select the therapeutic programs from P01 to P10. Each program has 3 treatment phases, you can set and save the parameters according to patient's need.
	CC	<ol> <li>Press "B2" Program button to select "CC" or "CV" control mode.</li> </ol>
	3 № 15′00″ 1	6. There are two modes in Quattro <sup>™</sup> II, press and hold "B8" button to switch Normal mode and Professional mode. In professional mode, each program has 3 treatment phase, the LCD display like left figure. Rotating the parameters control knob (③) to select phase program from 1 to 3 when the device enter into professional mode. The parameters of each phase program can be set according to following methods.
	F. M.	7. Press "B3" button to toggle F.M. parameter, then rotating the parameters control knob (③) to set the F.M. parameter from 0Hz to 249Hz, 1Hz/step. But F.M.+Freq.≤250Hz.
	Burst	8. Press "B3" button again to toggle Burst rate, then rotating the parameters control knob (③) to set the Burst rate from OHz to 10Hz, 1Hz/step. But Burst×8≤Freq.
	Freq.	<ul> <li>9. Press "B4" button to toggle Freq. parameter, then rotating the parameters control knob (③) to set the frequency from 1Hz to250Hz, 1Hz/step.</li> <li>But Freq. ≥Burst x 8 or Freq. ≤ 250-F.M.</li> </ul>



А. М.	<ol> <li>Press "B5" button to toggle A.M. parameter, then rotating the parameters control knob (③) to set the parameter from 0% to100%, 20%/step.</li> </ol>
P. Dur.	<ol> <li>Press "B6" button to toggle P.Dur. parameter, then rotating the parameters control knob (③) to set the pulse duration from 30μs to 400μs, 5μs/step.</li> </ol>
⊙         Treat.         I ⊆ Z <thi th="" z<="" ⊆=""> <thi th="" z<="" ⊆="">         I ⊆</thi></thi>	12. Press "B7" button to toggle Treat. time parameter, then rotating the parameters control knob (③) to set the treatment time from 1 min to 60 min, 1 min/step.
€ Cycle SSS	13. Press "B7" button again to toggle Cycle time parameter, then rotating the parameters control knob (③) to select the cycle time from "-/-(continuous)", "4/4", "4/8", "7/7", "5/5", "4/12", "10/10", "10/20", "10/30" and "10/50".
	14. Stick the electrodes on the patient. You can use one or two channel as your needs.
	<ul> <li>15. Adjust the output intensity and start electrical treatment that you are using by rotating the output intensity adjustable knob on the control panel. 0.5mA/step or 0.5V/ step. For safety using, load detection was designed in this device after the output intensity surpass 10. 0mA/10. 0V. If there are no electrodes stuck on patient' skin, an alarm buzzer sound will appear and the intensity value flashing.</li> </ul>
$\bigcirc$	<ol> <li>Press the "<sup>©</sup> button to stop treatment if any emergency happened.</li> </ol>
+ +/II	17. Press the " ►/II " button to pause treatment; you can press it again to restart the treatment.

6.4.4	Γ
Russian	
Stimulation	
Set-up	
Procedure	

o∏ċ	<ol> <li>In order to turn on the device, please press ON/OFF switch to [⊙] icon which is located on the side of the device.</li> </ol>
Russian ₽·0 / Ÿ         °°           ™ 50'         ™ 50           3         №. /0' 00"	<ol> <li>When you turn the Quattro<sup>™</sup> II on, the device will get down to self- check about 6~8 seconds, and then the default parameters are displayed the last treatment mode.</li> </ol>
Russian	3. Press "B1" Waveform button to toggle the therapeutic waveform, then rotating the parameters control knob (③) to select "Russian" waveform.
P-[] {	4. Press "B2" Program button to toggle the therapeutic program, and then rotating the parameters control knob (③) to select the therapeutic programs from P01 to P10. Each program has 3 treatment phases, you can set and save the parameters according to patient's need.
CC	5. Press "B2" Program button to select "CC" or "CV" control mode.
<sup>™</sup> 15′00″ 1	6. There are two modes in Quattro <sup>™</sup> II, press and hold "B8" button to switch Normal mode and Professional mode. In professional mode, each program has 3 treatment phase, the LCD display like left figure. Rotating the parameters control knob (③) to select phase program from 1 to 3 when the device enter into professional mode. The parameters of each phase program can be set according to following methods.
Freq.	7. Press "B4" button to toggle Freq. parameter, then rotating the parameters control knob (③) to set the frequency from 20Hz to100Hz,5Hz/step.
Duty 50%	8. Press "B3" button to toggle Duty parameter, then rotating the parameters control knob (③) to set the parameter from 10% to 50%, 10%/step.
• Treat.	<ol> <li>Press "B7" button to toggle Treat. time parameter, then rotating the parameters control knob (③) to set the treatment time from 1 min to 60min, 1 min/step.</li> </ol>



⊙       Cycle       S	10. Press "B7" button again to toggle Cycle time parameter, then rotating the parameters control knob (③) to select the cycle time from "-/-(continuous)", "5/5", "4/12", "10/10", "10/20", "10/30" and "10/50".
↔ Ramp	11. Press "B7" button again to toggle Ramp time parameter, then rotating the parameters control knob (③) to select the ramp time from 1s, 2s and 5s.
	12. Stick the electrodes on the patient. You can use one or two channel as your needs.
1 (mA) 2 (mA) (mA)	13. Adjust the output intensity and start electrical treatment that you are using by rotating the output intensity adjustable knob on the control panel. 0.5mA/step or 0.5V/ step. For safety using, load detection was designed in this device after the output intensity surpass 10. 0mA/10. 0V. If there are no electrodes stuck on patient' skin, an alarm buzzer sound will appear and the intensity value flashing.
$\bigcirc$	<ol> <li>Press the "<sup>™</sup> ©<sup>™</sup> button to stop treatment if any emergency happened.</li> </ol>
	15. Press the "▶/II" button to pause treatment; you can press it again to restart the treatment.

#### 7. MAINTENANCE

#### 7.1 Cleaning of the device

Switch off the device and disconnect it from the power supply. The apparatus can be cleaned with a damp cloth. Use lukewarm water and a non-abrasive liquid household cleaner (no abrasive, no alcohol content solution). If a more sterile cleaning is needed, use a cloth moistened with an antimicrobial cleaner.



Do not submerse the apparatus in liquids. Should the unit accidentally become submersed, contact the dealer or Authorized Service center immediately.Do not attempt to use a system that has been wet inside until inspected and tested by a Service Technician Certified by Authorized Service center. Do not allow liquids to enter the ventilation holes.

#### 7.2 Cleaning the electrodes

- Apply the protective backing to the tacky side of the electrode. Place the electrode on the side of the protective backing that is labeled with the word on.
- It may be helpful to improve repeated application by spreading a few drops of cold water over the adhesive and turn the surface up to air dry. Over Saturation with water will reduce the adhesive properties.
- Between uses, store the electrodes in the reusable bag in a cool dry place.

# Caution

- The electrodes are intended for single patient use only.
- If irritation occurs, discontinue use and consult your clinician.

wires will damage the insulation and dramatically shorten their life.

• Always use the electrodes with CE mark, or are legally marketed in the US under 510(K) procedure.

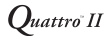
7.3 Cleaning the lead wires and cables

7.4 Maintenance • Maintenance and all repairs should only be carried out by an authorized agency. The manufacturer will not be held responsible for the results of maintenance or repairs by unauthorized persons.

Periodically wipe the lead wires clean with a cloth dampened in a mild soap

solution, and then gently wipe them dry. Use of rubbing alcohol on the lead

• Opening of the equipment by unauthorized agencies is not allowed and will terminate any claim to warranty.



#### 8. TROUBLESHOOTING

For optimal

use:

- Replace lead wires annually.
- Please follow the directions on the electrode packaging for the care of electrodes. The life of the electrodes varies, depending on skin conditions, skin preparation, storage and climate. Replace electrodes that no longer stick.
- NOTE: If the following measures fail to alleviate the problem, please call your dealer.

Problem	Possible Cause	Solution
Displays fail to light up	Adapter contact failure	Ensure adapter is connect. Check the following contacts: • All contacts are in place. • All contacts are not broken.
Stimulation weak	Electrodes 1. Dried out or contaminated 2. Placement	<ol> <li>Replace.</li> <li>Electrodes must be a minimum of 2 inches apart.</li> </ol>
	Lead wires Old/worn/damaged	Replace.
Stimulation stops	Poor electrode contact	Reapply electrodes, secure firmly.
	Damaged or worn electrodes or lead wires	Replace
Stimulation is	Intensity is too high	Decrease intensity.
uncomfortable.	Electrodes are too	Reposition the electrodes.
	close together	Electrodes must be a minimum of 2 inches apart.
	Damaged or worn electrodes or lead wires	Replace.
	Electrode active area size is too small.	Replace electrodes with ones that have an active area no less than 25.0cm <sup>2</sup> .
Stimulation is	Improper electrode	Reposition electrode
ineffective.	Unknown	Contact clinician.
"E1″ displays on LCD	Communication failure is detected	Restart the device, if the problem is still exist, please contact the
"E2" displays on LCD	The device detected the temperature transmitter was open circuit or short circuit	manufacturer or distributor

	limitative temperature	The device will stop treatment automatically, please wait
"E4" displays on LCD		several minutes before using again.
	Nemorizer failure is	Restart the device, if the problem is still exist, please contact the manufacturer or distributor

### 9. **SPECIFICATIONS**

9.1 General

Specifications:

Adapter supply voltage:	100V-240V, 50Hz-60Hz, 0.5A	
Adapter output:	15V1A Max.	
Adapter Dimensions:	110mm(L)*54mm(W)*33mm(H)	
Dimensions:	250mm(L)*185mm(L)*82mm(H)	
Operating Environmental:	Temperature:10°C(50°F) to 40°C(104°F),	
	Relative humidity: 30%-85%	
Storage Environmental:	Temperature: -20°C(-4°F) to 55°C(131°F),	
	Relative humidity: 20%-90%	
Maximum Treatment Time:	60 minutes	

## 9.2

Waveform Specifications:

#### 4-Pole Interferentia

Mode

al	Waveform Type	Bi-phasic square	
	Mode Selection	CC (Constant Current) or CV (Constant Voltage)	
	Vector	Auto: 0%-100% Manual: 0°–90°	
	Carrier Frequency (C.F.)	4.0kHz	
	Sweep Low Beat Frequency (Beat H.)	(Beat L.) -150 Hz	
	Sweep High Beat Frequency (Beat L.)	1-(Beat H.) Hz	
Output Intensity		0-50mA(CC, at 1k ohm load) 0-50V(CV, at 1k ohm load)	
	Treatment time	1-60 minutes	

2-Pole

2-Pole Interferential	Waveform Type	Bi-phasic square	
Mode	Mode Selection	CC (Constant Current) or CV (Constant Voltage)	
	Carrier Frequency (C.F.)	2.5kHz	
	Sweep Low Beat Frequency (Beat H.)	(Beat L.) -150 Hz	
	Sweep High Beat Frequency (Beat L.)	1-(Beat H.) Hz	
	Output Intensity	0-50mA(CC, at 1k ohm load) 0-50V(CV, at 1k ohm load)	
	Treatment time	1-60 minutes	
	Cycle time (cycle)	Continuous, 5/5, 4/12, 10/10, 10/20, 10/30, 10/50	
	Ramp time (Ramp)	2 seconds	

### **TENS** and EMS Mode

Waveform Type	Mono- or Bi-phasic square	
Mode Selection	CC (Constant Current) or CV (Constant Voltage)	
Frequency	1 - 250 Hz	
Frequency Modulation (F.M.)	0-249Hz	
Burst rate (Burst)	0-10Hz (7 pulse)	
Phase duration (P.Dur.)	30-400µs	
Amplitude Modulation (A.M.)	0%-100%	
Output Intensity	0–100mA(CC, at 1k ohm load) 0–100V(CV, at 1k ohm load)	
Cycle time (Cycle)	Continuous,4/4, 4/8,7/7, 5/5, 4/12, 10/10, 10/20, 10/30, 10/50	
Treatment time	1-60 minutes	
Ramp time	1 second	

#### Russian Mode [

Waveform Type	Bi-phasic square	
Mode Selection	CC (Constant Current) or CV (Constant Voltage)	
Carrier Frequency (C.F.)	2 .5kHz	
Burst frequency (Freq.)	20-100 Hz	
Output Intensity	0-50mA(CC, at 1k ohm load) 0-50V(CV, at 1k ohm load)	
Duty cycle	10%, 20%, 30%, 40%, and 50%.	
Cycle time	Continuous, 5/5, 4/12, 10/10, 10/20, 10/30, 10/50.	
Treatment time	1-60 minutes	
Ramp time	1s, 2s, and 5s	

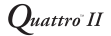
#### 10 STORAGE

For a prolonged pause in treatment, store the device with the adapter in a dry room and protect it against heat, sunshine and moisture. Store the machine in a cool, well-ventilated place. Never place any heavy objects on the machine.

#### 11 DISPOSAL



Please dispose of the device in accordance with the directive 2002/96/EC – WEEE (Waste Electrical and Electronic Equipment). Contact your local distributor for information regarding disposal of the unit and accessories.



12 EMC TABLE	Guidance and manufacturer's declaration - electromagnetic emissions			
	The Quattro <sup>TM</sup> II device is intended for use in the electromagnetic environment specified below. The customer or th user of the Quattro <sup>TM</sup> II should assures that it is used in such an environment.			
	Emissions test	Compliance	Electromagnetic environment - guidance	
	RF emissions CISPR 11	Group 1	The Quattro <sup>™</sup> II device 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 CISPR11	Class B		
	Harmonic emissions IEC 61000-3-2	Not applicable	The Quattro <sup>™</sup> II device is suitable for use in all establishments other than domestic and those directly	
	Voltage fluctuations / flicker emissions IEC 61000-3-3	Not applicable	connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.	

Guidance and manufacturer's declaration — electromagnetic immunity			
The Quattro <sup>TM</sup> II device is intended for use in the electromagnetic environment specified below. The customer or the user of the Quattro <sup>TM</sup> II should assure that it is used in such an environment.			
Immunity testIEC 60601 test levelCompliance levelElectromagnetic environ - guidance			
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	E6 kV ±6 kV contact contact contact	

Guidance and- manufacturer's declaration. Electromagnetic
immunity

The Quattro<sup>™</sup> II device is intended for use in the electromagnetic environment specified below. The customer or the user of the Quattro<sup>™</sup> II should assure that it is used in such an environment.

an environment.				
Immunity test	IEC 60501 test level	Compliance level	Electromagnetic environment - guidance	
			Portable and mobile RF communications equipment should be used no closer to any part of the E-Stim Basic Quattro <sup>™</sup> II device, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. <b>Recommended</b> separation distance	
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	3 Vrms	d=1.2√P	
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m	$d = 1.2\sqrt{P}$ 80MHz to 800MHz	
			<b>d=2.3√P</b> 80MHz to 2.5MHz	
			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 meters (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	

	Interference may occur In the vicinity of equipment marked with the following symbol:		
NOTE I At 80 MHz ends 800 MHz. th NOTE 2 These guidelines may not a Electromagnetic propagation is affe from structures, objects and people.	pply in all situations. acted by absorption and reflection		
<ul> <li>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 Quattro<sup>™</sup> II device is used exceeds the applicable RF compliance level above, should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Quattro<sup>™</sup> II.</li> <li>b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [Vi] V/m.</li> </ul>			
Recommended separation distances between portable and mobile RF communications equipment and the Quattro <sup>™</sup> II device			
The Quattro <sup>™</sup> II device is intended electromagnetic environment in whi controlled. The customer or the user device can help prevent electromag a minimum distance between portab equipment (transmitters) and the Qu	ch radiated RF disturbances are r of the Quattro <sup>™</sup> II netic interference by maintaining ole and mobile RF communications		

recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separation distance according to frequency of transmitterm			
output power of transmitter	150 kHz to 80 MHZ	80 MHz to 800 MHZ	800 MHz to 2,5 GHz	
W	d=1.2√P	d=1.2√P	d=2.3√P	
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) accordable to the transmitter manufacturer.

NOTE I At 80 MHz and 800 MHz. the separation distance for the higher frequency range applies.

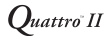
NOTE 2 These guidelines may not apply in all situations.

Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

**13.** Please contact your dealer or the device center in case of a claim under the warranty. If you have to send in the device, enclose a copy of your receipt and state the defect.

A. The following warranty terms apply:

- The warranty period for Quattro<sup>™</sup> II products is one year from date of purchase. In case of a warranty claim, the date of purchase has to be proven by means of the sales receipt or invoice.
- Defects in material or workmanship will be removed free of change with in the warranty period.
- Repairs under warranty do not extend the warranty period either for the device or for the replacement parts.
- B. The following is excluded under the warranty:
  - All damage which has arisen due to improper treatment, e.g. nonobservance of the user instruction.
- All damage which is due to repairs or tampering by the customer or unauthorized third parities.
- Damage which has arisen during transport from the manufacturer to the consumer or during transport to the service centre.
- Accessories which are subject to normal wear and tear.



• Liability for direct or indirect consequential losses caused by the unit is excluded even if the damage to the unit is accepted as a warranty claim.





Type BF Applied Part

Power polarity



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 $\ominus$   $\bullet$   $\bullet$ 



Type of protection against electric shock: Class II Equipment

Refer to Instruction Manual.

Electrical devices are recyclable material and should not be disposed of with household waste after their useful life! Help us to protect the environment and save resources and take this device to the appropriate collection points. Please contact the organization which is responsible for waste disposal in your area if you have any questions.



Equipment capable of delivering output values in excess of 10 mA r.m.s. or 10V r.m.s. averaged over any period of 5 s



Stop treatment

►/**||** 

Start/Pause the treatment



Manufactured for: Current Solutions<sup>™</sup> LLC 3814 Woodbury Drive Austin,TX 78704 Ph:(800)871-7858 www.currentsolutionsnow.com