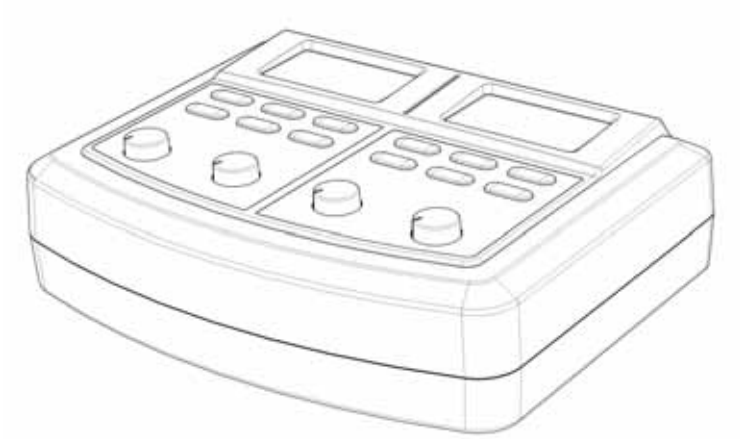


Powered Muscle Stimulator (EMS)

Read before using



QUATTRO EMS

Model: GM380E (4CH EMS)

FDA 510k



Operation Manual

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GENERAL DESCRIPTION

Electric stimulation has proven its high value as a method of pain therapy and is a great help to the experienced therapist.

With some indications, physicians can prescribe a unit to patients for the use at home.

This series of device is the combination of two units with dual-Channel electric stimulator for active treatment application, which is equipped with a Liquid Crystal Display indicating operation modes and output as well as an 8-bit micro computer for controlling the system.

The electronics of the unit create electric impulses, the intensity duration, frequency per second and modulation of these impulses can be adjusted through the button or knob.

SYSTEM COMPONENTS

Your device may include the following components or accessories:

- Unit
- Carrying case
- Lead wires / Electrodes
- 6 x UM-3/AA size 1.5V batteries.
- Adaptor
- Operation Manual

WARRANTY

This device carries a one-year warranty from the date of purchase. The warranty applies to the device and necessary parts and labor relating thereto. The distributor reserves the right to replace or repair the unit at their discretion.

The warranty does not apply to electrode, batteries, lead wires, carrying case, damage resulting from failure to follow the operating instructions, accidents, abuse, alterations or disassembly by unauthorized individuals.

INDICATIONS AND CONTRAINDICATIONS

Read the operation manual before using the device.

Federal law (USA) restricts this device for sale by or on the order of a physician.

Observe your physician's precise instructions and let him show you where to apply the electrodes. For a successful therapy, the correct application of the electrodes is an important factor. Carefully write down the settings your physician recommended.

Indications for use

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

Contraindications

Powered muscle stimulators should not be used on patients with cardiac demand pacemakers.

WARNINGS AND PRECAUTIONS** Warnings**

- The long-term effects of chronic electrical stimulation are unknown.
- Stimulation should not be applied over the carotid sinus nerves, particularly in patients with a known sensitivity to the carotid sinus reflex.
- Stimulation should not be applied over the 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 trans-thoracically in that the introduction of electrical current into the heart may cause cardiac arrhythmias.
- Stimulation should not be applied transthoracically.
- Stimulation should not be applied over swollen, infected, or inflamed areas or skin eruptions, e.g. phlebitis, thrombophlebitis, varicose veins, etc.
- Stimulation should not be applied over, or in proximity to, cancerous lesions.
- Stimulation should not be applied trans-cerebrally (across the head).
- EMS devices should be used only under the continued supervision of a physician.

 Precautions/adverse Reactions

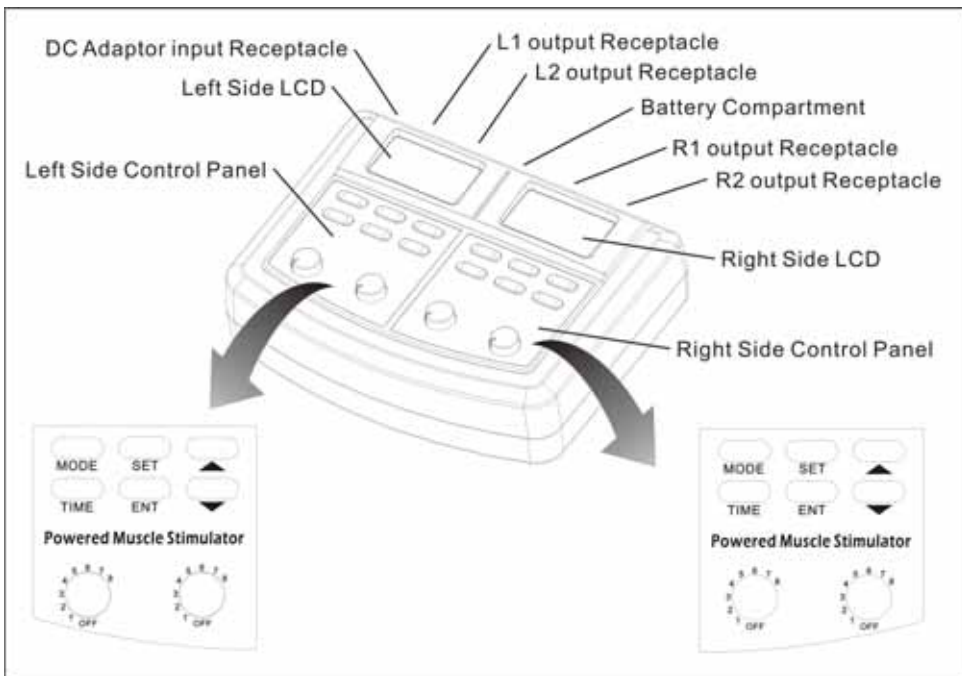
- Safety of powered muscle stimulators for use during pregnancy has not been established.
- Caution should be used for patients with suspected or diagnosed heart problems.
- Caution should be used for patients with suspected or diagnosed epilepsy.
- Caution should be used in the presence of the following:
 - a. When there is a tendency to hemorrhage following acute trauma of fracture;
 - b. Following recent surgical procedures when muscle contraction may disrupt the healing process;

- c. Over the menstruating or pregnant uterus; and
- d. Over areas of the skin which 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 alternate conductive medium, or alternate electrode placement.
- Electrode placement and stimulation settings should be based on the guidance of the prescribing practitioner.
- Powered muscle stimulators should be kept out of the reach of children.
- Power muscle stimulators should be used only with the leads and electrodes recommended for use by the manufacturer.

Skin irritation and burns beneath the electrodes have been reported with the use of powered muscle stimulators.

ABOUT THE DEVICE

Your device offers two controllable output channels for each stimulation unit. This device creates electrical impulses whose amplitude, duration, and modulation can be altered with the controls or buttons.



THE DEVICE CONTROLS

Intensity

The intensity knobs are located on the front of the unit for the strength adjustment of the stimulation and also function as ON/OFF controls.

Mode

The Mode key is used to select / set the type of treatment utilized. The available stimulation modes for each unit are as indicated in the following table:

Identification	Function as	Available mode
E	EMS	S (Synchronous), C (constant), A (Alternation)

TIME

Treatment Time of device can be pre-select / set with Time key. There are seven programs fixed duration of 5, 10, 15, 20, 30, 45, 60 minutes and continue. Press the key until engaged in position desired.

5 min : 1/4 circle icon flash.

10 min : 2/4 circle icons flash.

15 min : 3/4 circle icons flash.

20 min : 4/4 circle icons flash.

30 min : 2/4 circle icons light on.

45 min : 3/4 circle icons light on.

60 min : 4/4 circle icons light on.

Continue : the timer icon become empty.

SET/ENT

The SET/ENT keys are used to select the type of treatment utilized. The type of treatment utilized includes cycle On time, On Ramp time, cycle Off time and Pulse Rate. The selected parameter will flash during adjust.



The ▲ ▼ keys regulate the number of On, Ramp, Off and Pulse Rate value.

DANGER

The device does not have AP/APG protection. Explosion hazard is possible if used in the presence of explosives, flammable materials or flammable anesthetics.

Caution should be used when applying the device to patients suspected of having heart disease. Further clinical data is needed to show if there are adverse side effects on those with heart disease.

ATTACHING THE LEAD WIRES

The lead wires provided with the device insert into the jack sockets located on rear of the unit. Holding the insulated portion of the connector, push the plug end of the wire into one of the jacks. After connecting the wires to the stimulator, attach each wire to an electrode.

Lead wires provided with the device are compliant with mandatory compliance standards set for the by FDA.

Note: Use carefully when you plug and unplug the wires. Pulling on the lead wire instead of its insulated connector may cause wire breakage.

Caution: Never insert the plug of the lead wire into an AC power supply socket.

ELECTRODE SELECTION AND CARE

Your physician/practitioner should decide which type of electrode is best for your condition. Follow application procedures outlined in electrode packaging to maintain stimulation and prevent skin irritation. The electrode packaging will provide instruction for care, maintenance and proper storage of your electrodes. Be sure to use the electrodes provided by the manufacturer and/or the similar FDA legally marketed electrode in particular the same cross section area.

TIPS FOR SKIN CARE

Good skin care is important for comfortable use of your device.

- Always clean the electrode site with mild soap and water solution, rinse well, and blot dry thoroughly prior to any electrode application.
- Any excess hair should be clipped, not shaved, to ensure good electrode contact with the skin.
- You may choose to use a skin treatment or preparation that is recommended by your physician. Apply, let dry, and apply electrode as directed. This will both reduce the chance of skin irritation and extend the life of your electrodes.
- Avoid excessive stretching of the skin when applying electrodes, this is best accomplished by applying the electrode and smoothly pressing it in place from the center outward.
- When removing electrodes, always remove by pulling in the direction of hair growth.
- It may be helpful to rub skin lotion on electrode placement area when wearing electrodes.

CONNECTING THE DEVICE

1. Prepare the Skin

Prepare the skin as previously discussed and according to instructions provided

with your electrodes. Before attaching the electrodes, identify the area in which your physician/practitioner has recommended for electrode placement.

2. Connect lead wires to the electrodes

Connect the lead wires to the electrodes before applying the electrodes to the skin.

Note: Be sure all intensity controls for every Channel 1 and 2 of each unit are turned to the "OFF" position.

3. Place Electrodes on Skin

Place the electrodes on the skin as recommended by your clinician.

4. Insert Lead Wire Connector to device

Plug end of lead wire into the channel output receptacle to be used, pushing plug in as far as it will go.

5. Select Treatment Settings

Check and be sure your unit is still set to the proper settings recommended by your physician/practitioner.

6. Adjusting Channel Intensity Control

Locate the intensity control knob at the front of the unit. Slowly turn the intensity control knob for the channels to be used clockwise until you reach the intensity recommended by your medical professional. Always start with the lowest step and increase slowly.

If the stimulation levels are uncomfortable or become uncomfortable, reduce the stimulation amplitude to a comfortable level or cease stimulation and contact your physician/practitioner if problems persist.

BATTERY INFORMATION

The 6 x 1.5-volt UM-3/AA size disposable batteries are provided with your unit. When the low battery mark light on the LCD the battery has become too weak to power the unit and it is time to change the battery. At this point, the unit will shut off until a fresh battery is inserted.

CHANGING THE BATTERIES

When the low battery mark light is on the LCD panel, the batteries should be replaced with the fresh batteries.

1. Remove the battery cover by pressing this cover and sliding down until it is completely removed from the unit this will reveal the battery compartment.
2. Remove the discharged batteries from the device.
3. Place new batteries in compartment. Note the proper polarity alignment indicated on the battery and the compartment.

9-Volt DC Adaptor

Because the current consumption is large for the device, it is recommended to use the 9V DC adaptor. Plug the DC plug into the DC jack of the unit, and plug the adaptor into your 110V wall outlet. Please be sure you are using the correct polarity of DC plug.

CLEANING FOR YOUR DEVICE

Your device may be cleaned by wiping gently with a damp cloth moistened with mild soap and water. Never immerse the device in water or other liquids.

Wipe lead wires with a damp cloth as above if they become soiled.

To properly store the device an extended period of time, remove the battery from the unit. Put the unit and accessories in the carrying case and store in a cool dry location.

TROUBLESHOOTING

If the device does not function properly:

1. Make sure the battery is properly installed or replace battery. Be sure to observe proper polarity markings when replacing the battery. If the low battery mark light is on the LCD pane when the unit is turned on, replace the battery and check again.
2. If the intensity has been adjusted and there is no stimulation, check that the lead wires are properly connected and the electrodes are in place. If the unit appears to be functioning and no stimulation, the lead wires or electrodes may need to be replaced.
3. If the battery appears to be charged and the unit is not functioning, **turn both intensity Control Knobs of each unit to the OFF position (counter clockwise) for about 7 sec.** Then gradually turn the intensity Control Knob clockwise until stimulation is felt. If the device is still not working, turn off your device and contact your distributor.

Note : For any problem, please contact your distributor. Do not try to repair this device by yourself.

TECHNICAL SPECIFICATIONS

Channel:	Four, isolated between channels
Pulse amplitude:	Adjustable 0-100mA peak into 500 Ω load each channel, constant current
Pulse Rate:	1Hz-160Hz (adjustable), 1Hz/step
Pulse Width:	300 μ s fixed.
Software ramp up feature:	Pulse width ramp up when change mode
Patient Compliance counter:	Shows the treatment times. Account by hours.
Patient Lock System:	To prevent the user from changing any fixed parameters set by the physician.
Timer:	5,10,15,20,30,45,60 minutes or Continue selectable
LCD:	Shows modes, pulse rate, pulse width, timer, CH1/CH2 of each unit.
ON Ramp:	Adjustable 1-8 seconds. 1sec/step.
Cycle ON Time:	Adjustable 1-30 seconds. 1 sec/step.
Cycle OFF Time:	Adjustable 1-30 seconds. 1 sec/step.

Function Modes:

S: Synchronous.

C: Constant.

A: Alternation.

Wave Form: Asymmetrical Bi-Phasic square pulse.

Max charge per pulse: 30 micro-coulombs maximum

Voltage: 0-110 Volt (Open Circuit).

Power Source: 6 pieces of 1.5V UM-3/AA size batteries or 9-Volt adaptor

Dimension: 287 x 224 x 79 mm

Weight: Approx. 1250g (without batteries)

All electrical specifications are $\pm 10\%$ 500 Ω load.



Remark:

Printed in Taiwan

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