

		产品名称	说明书	文件编号	250117
				要 求	骑马钉
产品名称	经皮神经刺激器	产品型号	KTR-2401	版 本	A/1
编制		日期		尺 寸	L130 x W120mm
审核		日期		材 质	80G书写纸/骑马钉
批准		日期		颜 色	4C



**PL009B Penguin Electronic Pulse TENS EMS**  
Transcutaneous Electrical Nerve Stimulator



**User Manual** **KTR-2401**

Thanks for choosing this product, please read this User Manual carefully before use!

# **Content**

<b>1. Foreword.....</b>	<b>2</b>
<b>2. Warnings.....</b>	<b>2</b>
<b>3. Intended use.....</b>	<b>2</b>
<b>4. Product Introduction.....</b>	<b>3</b>
<b>5. Primary Structure of Product.....</b>	<b>3</b>
<b>6. Product Main Features.....</b>	<b>4</b>
<b>7. Product Require ments and Major Parametric Description.....</b>	<b>4</b>
<b>7.1 Product Power Requirement.....</b>	<b>4</b>
<b>7.2 Product Main Technical Parametric description ....</b>	<b>4</b>
<b>7.3 Product Environmental Requirement.....</b>	<b>5</b>
<b>a) Normal work environmental requirements:.....</b>	<b>5</b>
<b>b) Storage environment requirements:.....</b>	<b>5</b>
<b>c) Transport environment requirements:.....</b>	<b>5</b>
<b>8. Method of application.....</b>	<b>5</b>
<b>9. Safety Precautions.....</b>	<b>7</b>
<b>10. Product maintenance.....</b>	<b>10</b>
<b>11. Product Scrap Processing.....</b>	<b>10</b>
<b>12. Accessories and Parts.....</b>	<b>11</b>
<b>13. The Paraphrase of Graphic Symbol.....</b>	<b>12</b>
<b>14. Executive Standards.....</b>	<b>13</b>
<b>Appendix EMC Declaration.....</b>	<b>13</b>

## 1. Foreword

Thank you very much for your trust and for your purchase of Transcutaneous Electrical Nerve Stimulator. Our objective is to help you live a more enjoyable life with less pain. Before use, please read the User's manual carefully, so that you can operate it correctly for best results. The User's manual should be kept for future reference.

## 2. Warnings

- a) The long-term effects of chronic electrical stimulation to the human body are not known.
- b) Patients with cardiac pacemaker are disabled!
- c) Patients with suspected or confirmed have epilepsy, heart disease, pregnancy or menstruating women are prohibited.
- d) Treatment parts have the haemorrhage after acute trauma or fractures, disabling wound healing stage after the operation.
- e) Treatment is not sensitive to heat or electrical stimulation of the skin patients disabled.
- f) Disturbance of consciousness or disabled children.
- g) Treatment of swelling, skin infection or with skin disease, phlebitis patients disabled.
- h) With wet skin after shower, sweating and disabled when you go to sleep.
- i) Driving traffic tools or disable during movement.
- j) Metal allergies are disabled.

## 3. Intended use

To be used for temporary relief of pain associated with sore and aching muscles in the shoulder, waist, back, back of the neck, arm, leg and foot, due to strain from exercise or normal household and work activities.

## 4. Product Introduction

Transcutaneous Electrical Nerve Stimulator (Models: KTR-2401) is a portable and battery powered multifunctional device.

Transcutaneous Electrical Nerve Stimulator has 3 operation modes, which can give certain electrical pulse through electrode pads placed on the skin to help users to enjoy body massage.

The electronic stimulatory module has the operating elements of ON/OFF Key and Intensity Modification keys.

The device is equipped with accessories of electrode pads and rechargeable batteries.

The electrode pads are complying with the biocompatibility standards ISO 10993-5 (Cytotoxicity) and ISO 10993-10 (Irritation and Sensitization), are interchangeable.

## 5. Primary Structure of Product

The Transcutaneous Electrical Nerve Stimulator (Model: KTR-2401) The primary structure and components as shown below:



Fig 1. Primary structure and components of KTR-2401



## 6. Product Main Features

- 1.The device is convenient to carry, and is suitable for personal health care;
- 2.Electrode patch intelligent detection and protection, users use more secure;
- 3.Electric pulse combination, 16 level strength can be adjusted, according to personal preferences to adjust the need;
- 4.Integrated fuselage design, functional operation, simple and clear;
- 5.Rechargeable lithium design, built-in power protection circuit, easy to use, safe;
- 6.Contains a variety of massage procedures, can meet different massage needs, suitable for a wider range of people;

## 7. Product Requirements and Major Parametric Description

### 7.1 Product Power Requirement

- a) Power source: Rechargeable lithium battery3.7v/100mAh
- b) Device safety classes: class II type BF

### 7.2 Product Main Technical Parametric description

- a)Impulse frequency:  
TENS:20-100Hz
- b)Pulse width:  
TENS: 120μs
- c)Impulse waveform: square wave
- d)The product output has no the DC component.
- e)Influence of output end open circuit and short circuit: it is able to support the influence of output end open and short circuit, and its performance will not be allowed to weaken.
- f)Adjustment of output amplitude: 0-16 levels.
- g)The timer of therapeutic equipment: 15 minutes.
- h)Safety classification: internal electric source class and type BF equipment
- i)Boundary dimension:  
Main unit : Model KTR-2401: Φ46.3\*12.07;
- j)Product software version No.: A/0
- k)Service life: the shelf life of subject device is 2 years; the use life of the Electrode Pad is 80 times and shelf life is 2 years

### 7.3 Product Environmental Requirement

(Transportation and storage environment between secondary uses)

- a)Normal work environmental requirements:  
Environment temperature: +5 ℃~+40 ℃;  
Environment humidity: 0%-80%RH;  
Atmospheric environment conditions: 700hPa-1060hPa.
- b)Storage environment requirements:  
Environment temperature: -25 ℃~+55 ℃;  
Environment humidity: 0-93%RH;  
Atmospheric environment conditions: 700hPa-1060hPa.
- c)Transport environment requirements:  
Environment temperature: -25 ℃~+55 ℃;  
Environment humidity: 0-93%RH;  
Atmospheric environment conditions: 700hPa-1060hPa.



When the ambient temperature is 20 ℃, the time required for the product to be ready for its intended use from the minimum storage temperature after use is about 2 hours; At an ambient temperature of 20 ℃, the time required for the product to go from the maximum storage temperature after use to the time the product is ready for its intended use is approximately 2 hours.

## 8. Method of application

### (1)Preparation before use

Use the electrode plate for the first time, install the electrode plate to the master machine of the massage machine, and then align the metal buckle of the main machine to remove the film on the electrode sheet,The electrode film is affixed to the main body of the massage instrument,Attach the main body to the use part;

### (2)Switch machine

Long press  key, the host boot starts, at the same time will send the "drop" sound prompt, the strength output default is 0, the host starts to work; the default is the first mode of operation. Press the  key again, then turn off the machine.

### (3)Charge


If you can't turn on the machine, or turn it off automatically, or when the buzzer is alerted for 30s, it indicates that the product needs to change the battery.

#### (4)Regulating intensity

Users can press the "+" key, set the strength, press the "-" key, you can reduce the strength; short press the "+" key or "-" key strength will increase or lower level, at the same time will send out a "drop", the user can choose their own strength, in order to the most comfortable; (a total of 0 ≤ 16 levels of strength adjustment)

The host computer has the intelligent detection function, when the electrode piece and the skin contact is good, the main opportunity moderates the force, reaches the set strength directly.

#### (5)Mode selection

A total of three modes can be selected; in work, short press the  key to switch the working mode; when the mode is switched, the strength automatically recovers to the "1" file to prevent the intensity from being very high at the beginning of the mode switching, which makes the user feel uncomfortable. According to their own preferences, the user can choose different working modes.

#### (6)shut down

Work from start to end about 15 minutes. the host will automatic shutdown after 15 minutes;

If the host is not used in the boot state, if the host is not used within 30 seconds, the host will automatically shut down the computer.




Long press the  key, you can manually shut down the machine halfway.



### 8.1 The use of battery

- 1)The device powered by the lithium battery, which can rechargeable through the adapter.
- 2)If the indicator is flashing, showing the electric quantity of battery is not enough and please charging in time.
- 3)If it is not used for one month or long term, please fully charge it first.
- 4)Do not use the device under the environment which more than 45 °C , otherwise it will affect the performance and life of the battery.
- 5)The discarding method of the post-batteries should be deal with according to the urban environmental protection.

## 9. Safety Precautions

- The manual said that the purpose of warning sign and legend is safe and proper using the product by yours, and prevents the harm to you and others.
- The warning sign and legend as well as their meaning is as follows:

Warning Sign	Meaning
 using contraindication	It shows that it will appear the dangerous of casualties or serious injury in the error use.
 Warning	It shows that it will appear the possibility of casualties or serious injury in the error use.
 attention	It shows that it will appear the possibility of personal injury or damaged goods in the error use.

 attention	
<p>-- The therapeutic apparatus cannot be used with the HF apparatus to avoid burns or damaged apparatus</p> <p>-- If the patient himself uses the therapeutic apparatus and HF apparatus at the same time, the part of massage plate may be cause burns on apparatus, it may also damage the apparatus; If use the apparatus near (1 meter) the short wave or microwave therapeutic apparatus, the output of apparatus may be instability.</p> <p>-- It will increase a danger of heart fibrillation by using electrode pads close to the chest.</p> <p>-- Do not modify this equipment without authorization of the manufacturer</p> <p>--When need replacing the lithium batteries, please contact aftersales staff designated and authorized by the manufacturer directly, its accessories shall not be replaceable.</p> <p>-- Product battery replacement should be replaced by professional maintenance staff, otherwise it will produce risk.</p> <p>--Shall not make personnel can no longer take care of themselves, infant, or not sensitive person use.</p>	

<p>-- Simultaneous connection of a patient to a high frequency surgical ME equipment may result in burns at the site of the stimulator electrode pads and possible damage to the stimulator.</p> <p>-- Operation in close proximity (e.g. 1 m) to a shortwave or microwave therapy ME equipment may produce instability in the stimulator output.</p> <p>--Application of electrode pads near the thorax may increase the risk of cardiac fibrillation.</p> <p>--Advice that stimulation should not be applied across or through the head, directly on the eyes, covering the mouth, on the front of the neck, (especially the carotid sinus), or from electrode pads placed on the chest and the upper back or crossing over the heart.</p> <p>--Prevent inhalation or accidental swallowing of small parts (including 3*AAA LR03 battery).</p> <p>-- Prevent sharp parts from damaging the product.</p> <p>-- Do not use accessories not specified by the manufacturer.</p>	
<p> <b>using contraindication</b></p>	
<p>Pregnant women and women's menstrual period, person with sensitive skin, heart disease, abnormal blood pressure, malignant tumors, cerebrovascular patients , patients with acute disease or other person treated by doctor must consult a doctor rear can use this product.</p>	
<p>(1) It is contraindicated for use by those who has a skin perceptual disturbance or is not sensitive about the heat.</p> <p>(2)It is prohibited to use when bathing, sweating and sleeping.</p> <p>(3)The patient with cerebral hemorrhage: It should be disabled in unsteady phase; the person who has sequela must be used under supervision of doctors.</p> <p>(4)It is contraindicated for use by those who has purulent inflammation, acute blood poisoning and continuous hyperpyrexia.</p> <p>(5)It is contraindicated for use by those who has acute cardiovascular and cerebrovascular diseases.</p> <p>Please stop using it immediately to the doctor to consult when unwell felt or skin complaint in the process of use.</p>	

<p> <b>precautions</b></p>	
<p>Please do not use in the parts which is nearest heart, head, eyes, front neck (especially the carotid artery), lower back, oral cavity or pudendum, skin disease.</p>	
<p>(1)It should be shut down then using again when the apparatus is moved or changed the therapeutic parts in the using process, otherwise there will be a strong stimulation.</p> <p>(2)It is not permitted to give children or people who were no ability to express consciousness to use.</p> <p>(3)If you feel unwell because of using the product, please stop using it immediately to the doctor consult.</p> <p>(4)Please pull up the power plugs when you are done or do not use the product.</p> <p>(5)Do not used in conjunction with other medical electronic devices, such as cardiac pacemaker, artificial heart and lung and other medical electronic devices with fuels life, electrocardiograph and other medical electronic devices, otherwise it will lead to danger.</p> <p>(6)Do not use the product in the place where is direct sunlight, high heat, inflammable, electromagnetic radiation and humid.</p> <p>(7)Do not disassemble, repair and transform the therapeutic apparatus, otherwise cause failure or get an electric shock accident.</p> <p>(8)The therapeutic apparatus should be placed in the position which is easy to move the plugs to use, it is easy to move the plugs in emergency circumstances.</p> <p>(9)Check the equipment before each use to avoid the exposed wires caused by accidental damages or other reasons.</p> <p>(10)Dust may affect the performance of the unit, please use a dry soft cloth to clean the device as needed.</p> <p>Please check whether the electrode is loose before each use, otherwise it may have adverse effects on performance or cause other problems.</p> <p>The patient should be the intended operator.</p> <p>Please do not maintain or repair the equipment during use, or it may cause danger.</p>	

## 10. Product maintenance

- 1) Before turn on the main unit, please check whether the battery is installed correctly, each connecting parts whether is connected in good;
- 2) If the apparatus runs normally, show key invalidly after press. To turn it off after a few seconds and then opening, if still not normal, check whether it is broken for the apparatus;
- 3) If all operation of the apparatus are normal but without output (the user without feelings) after turn it on. Please check whether the electrode pads is directly contacted with skin, the contacting parts whether has hair, clothes and so on;
- 4) While cleaning, shutdown first, let the electrode wires stay away from the electric supply socket. You can wipe by using soft cloth or towel with a little water and after dry.
- 5) After each use, please clean electrode pads, you can wipe by using soft cloth or towel with a little water and after dry, when conducting electrode pads are particularly dirt, you can wipe by using soft cloth with a little medicinal alcohol (alcohol concentration as 75%).
- 6) The apparatus should be placed in the place which is dry, ventilated and good insulation use.
- 7) When the apparatus is moved, must be handled with care, avoiding the shake.
- 8) The apparatus should be checked the batteries condition, to check whether the output is good, if has abnormal conditions, you should pay attention to it and service it in time.
- 9) The apparatus' shell board should be pay attention to protection, avoiding the abrasion.
- 10) The non-professional cannot disassemble the host to avoid getting an electric shock or damaging the apparatus, an accident can happen because of it, all at your peril.

## 11. Product Scrap Processing



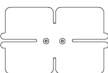





The therapeutic equipment belongs to medical device, if the complete machine is already aging and is not able to use, the process mode according to the local legislation carry through scrap processing.

## 12. Accessories and Parts

### 12.1 Electrode pads

Electrode Pads of Transcutaneous Electrical Nerve Stimulator

Please see all the electrode pads in the table below. The electrode pads here are for customers to choose

NO.	Model	Picture	Size	Effective Area (cm <sup>2</sup> )	Material
1	EPAD-H01		125×55mm	26	Hydrogels
2	EPAD-H02		130×60mm	22	Hydrogels
3	EPAD-F01		176×105mm	73	Hydrogels
4	EPAD-F02		166×166mm	80	Hydrogels
5	EPAD-F03		192×160mm	78	Hydrogels
6	EPAD-B01		240×65mm	50	Hydrogels
7	EPAD-T01		393×110mm	92	Hydrogels
8	EPAD-Z01		380×268mm	140	Silver paste

## 13. The Paraphrase of Graphic Symbol

Symbol	Explanation
	Production Batch
	Product catalogue reference code
	Manufacturer (Regulation (EU) 2017/745 for Medical Devices)
	DATE OF MANUFACTURE. This symbol shall be accompanied by a date to indicate the date of manufacture
	Caution
	Applied part of type BF
	European Authorized Representative
	Symbol for CE Mark. This symbol certifies that a product has met European Union consumer safety, health, or environmental requirements..
	CE marking with the Registration Number of the Notified Body. Regulation (EU) 2017/745 for Medical Devices
	Level of protection against the insertion of solid bodies of size/diameter $\geq 12$ mm and liquids in the presence of dripping water when tilted at $15^\circ$ compared with product.s.
	Refer to instruction manual/ booklet
	Medical device

## 14. Executive Standards

The product conforms to the following standards and laws:

1. IEC 60601-1:2005+A1: 2012 Medical electrical equipment-Part 1: General requirements for basic safety and essential performance
2. IEC 60601-1-11: 2015 Medical Electrical Equipment - Part 1-11: General Requirements For Basic Safety And Essential Performance - Collateral Standard: Requirements For Medical Electrical Equipment And Medical Electrical Systems Used In The Home Healthcare Environment
3. IEC 60601-2-10: 2013 Medical electrical equipment - Part 2-10: Particular requirements for the safety of nerve and muscle stimulators
4. IEC 60601-1-2:2014 Medical Electrical Equipment - Part 1-2: General Requirements For Basic Safety And Essential Performance - Collateral Standard: Electromagnetic Disturbances - Requirements And Tests

## Appendix EMC Declaration

The equipment is intended for use in the electromagnetic environment specified below.

The customer or the user of the EQUIPMENT should assure that it is used in such an environment.

The Transcutaneous Electrical Nerve Stimulators is suitable for use in a professional health care environment, not including areas where there are sensitive equipment or sources of intense electromagnetic disturbances, such as the RF shielded room of an imaging system magnetic resonance imaging, in operating rooms near active AF surgical equipment, electrophysiology laboratories, armored rooms or areas where short wave therapy equipment is used.



- Do not use the system around strong electric field, electromagnetic field (e.g. MRI scan room) and mobile wireless communication devices. Using the device in an improper environment may cause malfunction or damage.



- The compliance with EMC and EMI regulation cannot be guaranteed by the use of modified cables or those which does not comply with the same standards under what the equipment was validated.
- The system must not be used adjacent or supported by other equipment. The recommendations of this manual must be followed.
- Do not use accessories, transducers, internal parts of components and other cables other than those previously specified by the manufacturer. This may result in increased emission or decreased electromagnetic immunity and result in improper operation.
- Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30cm to any part of the ultrasound system, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.
- To maintain basic safety in relation to electromagnetic disturbances during the expected service life, always use the system in the specified electromagnetic environment and follow the maintenance recommendation described in this manual.

**The following tables provide information on compliance of the equipment according to the standard EN 60601-1- 2:2015.**

**Table 1 Compliance class**

Emissions Test	Compliance	Electromagnetic Environment and Guidance
RF emissions CISPR 11	Group 1	The equipment 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 B	

Harmonic emissions IEC 61000-3-2	Class A	The equipment is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	

**Table 2- Compliance standards**

Phenomenon	Basic Standard of	Immunity Test	Level of
Electrostatic discharge	IEC 61000-4-2	±8 KV contact ±2 KV, ±4 KV, ±8 KV, ±15KV air	±8 KV contact ± 2 K V, ± 4 K V, ± 8 KV, ±15KV air
Radiated RF EM fields1	IEC 61000-4-3	3V/m 80 MHz-2.7 GHz 80% AM at 1 KHz	3V/m 80 MHz-2.7 GHz 80% AM at 1 KHz
Proximity fields from RF wireless communication equipment	IEC 61000-4-3	See table	See table
Electrical Fast/Transients bursts	IEC 61000-4-4	±1 KV 100 KHz repetition frequency	±1 KV 100 KHz repetition frequency
Conducted disturbances induced by RF fields.	IEC 61000-4-6	3V 0.15 MHz-80 MHz 6 Vm in ISM bands between 0. 15 MHz and 80 MHz 80% AM at 1KHz	3V 0.15 MHz-80 MHz 6 Vm in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1KHz
Rated power frequency magnetic fields	IEC 61000-4-8	30 A/m 50 Hz or 60 Hz	30 A/m 50 Hz or 60 Hz


**Table 3- Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipments**

Test Frequency (MHz)	Band (MHz)	Service	Modulation	Maximum Power (W)	Distance (m)	Immunity Test Level (V/m)
385	380-390	TETRA 400	Pulse modulation 18 Hz	1.8	0.3	27
450	430-470	GMRS 460, FRS 460	FM $\pm 5$ KHz deviation 1KHz sine	2	0.3	28
710	704-787	LTE 13, 17 Band	Pulse modulation 217 Hz	0.2	0.3	9
745						
780						
810	800-960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE 5 Band	Pulse modulation 18 Hz	2	0.3	28
870						
930						
1720	1700-1990	GSM 1800, CDMA 1900, GSM 1900, DECT, LTE 1, 3, 4, 25 Band, UMTS	Pulse modulation 217 Hz	2	0.3	28
1845						
1970						
2450	2400-2570	Bluetooth, WLAN 802.11 b/g/n, RFID 2450, LTE 7 Band	Pulse modulation 217 Hz	2	0.3	28
5240	5100-5800	WLAN 802.11 a/n	Pulse modulation 217 Hz	0.2	0.3	9
5500						
5785						

## CONSUMER WARRANTY

Prospera offers a 90-day standard warranty on massage products. Please provide the model number and dated proof of purchase (sales receipt) when contacting Prospera for warranty service. Non-warranty service is provided on a “per incident” basis. The consumer will verify that the product has failed and provide information for servicing the unit. All applicable repairs, parts, shipping, handling, local tax and “per incident” fees will be charged for nonwarranty repairs and support calls.

## WARRANTY CARD

 **WARRANTY REGISTRATION FORM**

Product Name: \_\_\_\_\_ Purchase Price: \_\_\_\_\_  
 Product Model: \_\_\_\_\_ Retailer: \_\_\_\_\_  
 Date of Purchase: \_\_\_\_\_ First Name: \_\_\_\_\_ Last Name: \_\_\_\_\_  
 Address: \_\_\_\_\_  
 City: \_\_\_\_\_ State: \_\_\_\_\_ Zip: \_\_\_\_\_  
 Phone: \_\_\_\_\_ Email: \_\_\_\_\_  
 Signature and date: \_\_\_\_\_

Please mail to: \_\_\_\_\_ Tel: (925)292-5978  
 Prospera Corporation  
 392 West Larch Rd Suite 31  
 Tracy, CA 95304-1644  
 Email: sales@prosperacorp.com

 Shenzhen Kentro Medical Electronics Co., Ltd.  
 2nd Floor, No. 11, Shanzhuang Road, Xikeng Village, Yuanshan Street,  
 Longgang District, Shenzhen City, Guangdong Province, China

 2862

**RoHS**

