

CuringPen-E
Dental Curing Light USER MANUAL
Changzhou Sifary Medical Technology Co.,Ltd.

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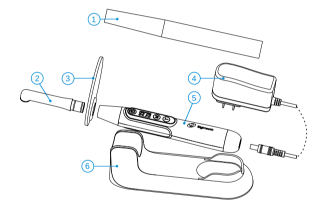
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1.Scope of CuringPen-E

1.1 Parts Identification

- 1 Disposable sleeve
- 2 Light source head
- 3 Protective light shield
- (4) Adapter
- (5) Handpiece
- (6) Base



1.2 Components and accessories

nz componente ana accessories		
Handpiece (1pcs) 6551021	Light source head (1pcs) 6551020	
Disposable sleeves (100 pcs) 6531034	Base (1pcs) 6551022	
Protective light shield (1pcs) 6551005	Manual(1pcs) 6535021	

For different regions, there are several different adapter options to be selected as follows.

Standard	Adapter	Power plug
European standard	Adapter (1pcs) 6516007	1
American standard	Adapter (1pcs) 6516008	American standard power plug (1pcs) 6316008
Multi- standard	Adapter (1pcs) 6516008	British standard power plug (1pcs) 6316006
		Australian standard power plug (1pcs) 6316007
		Argentina standard power plug (1pcs) 6316011

2. Symbols Used

Warning	If the instructions are not followed properly, operation may lead to hazards for the product			
<u> </u>	or the user/patient. Additional information, explanation of			
NOTE	operation and performance.			
SN	Serial number			
REF	Catalogue number			
	Manufacturer			
<u>~</u>	Date of manufacture			
	Class II equipment			
∱	Type B applied part			
===	Direct current			
A	Dispose of in accordance with the WEEE directive			
*	Keep dry			
2	Do not reuse			
₿	Consult instructions for use			
EC REP	Authorized Representative in the European Community			
-20°C 55°C	Temperature limitation			
20%	Humidity limitation			
70kPa 106kPa	Atmospheric pressure limitation			
Eighteeth	Manufacturer's LOGO			
LOT	Lot number			
MD	Medical Device			

3. Before Use

3.1 Scope of application

CuringPen-E is intended to cure dental resins and composites.

This device must only be used in hospital environments, clinics or dental offices by qualified dental personnel and not used in the oxygen-rich environment.

The Turbo Mode (P3) must only be used for direct restorations in the posterior region (Class I & II). Do not use the Turbo Mode in cases of deep cavities (caries profunda).

3.2 Contraindications

Materials, the polymerization of which is activated outside the wavelength range of 380 - 515 nm (no materials known to date). If you are not sure about certain products, please ask the manufacturer of the corresponding material.

Do not use the device for non dental procedure.

Safety and effectiveness have not been established in pregnant women and children



Warning

Read the following warnings before use:

- The device must not be placed in humid surroundings or anywhere where it can come into contact with any type of liquids.
- Do not expose the device to direct or indirect heat sources. The device must be operated and stored in a safe environment.
- 3. The device requires special precautions with regard to electromagnetic compatibility (EMC) and must be installed and operated in strict compliance with the EMC information. In particular, do not use the device in the vicinity of fluorescent lamps, radio transmitters, remote controls, portable or mobile RF communication devices and do not use this system near the active HF Surgical Equipment in the hospital. Please charge at least 3 hours before first use .Do not charge, operate or store at high temperatures. Comply with the specified operating and storage conditions.
- Protective light shield and a disposable sleeve are compulsory during treatment.

- 5. If the light emission window cannot be optimally placed in relation to the composite restoration, the restoration must be polymerized using a conventional method. If soft tissue exposure to the curing light cannot be avoided, the Turbo Mode must not be used, as exposure may result in damage of the soft tissues.
- If irregularities occur in the device during treatment, switch it off. Contact the agency.
- No modification of this equipment is allowed. Never open or repair the device yourself, otherwise, void the warranty.

4.Setting up the CuringPen-E

4.1 Install the light source head

Make sure the light source head align to the slots of the handpiece. Push gently until there is a "click" sound which indicates that the light source head is securely installed into the handpiece.



The light source head can be 360 degrees rotated without being taken off, which makes it easy to watch the LCD during the treatment.



Warning

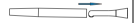
 Only the original light source head can be used.
 Check the light source head and handpiece before

- installation. Do not use damaged light source head and handpiece.
- After installing the light source head, pull it gently to make sure the connection is good, otherwise, it may cause unexpected fault, even hurt the patients.

4.2 Install the disposable sleeve

Apply a disposable sleeve over the entirety of the light source head

and handpiece before beginning a procedure.



4.3 Install the protective light shield

Make sure the light source head align to the slots of the protective light shield, plug them together.



Warning

- Disposable sleeves must be discarded after each use.
- The light source head, protective light shield , Base and handpiece should be cleaned and disinfected after each treatment.



Plug the round connector of the adapter into the charging hole at the rear of handpiece, and then the buzzer makes beeps twice, and then the display symbol of charging cycle will show on the screen,and then place the handpiece on the Base.





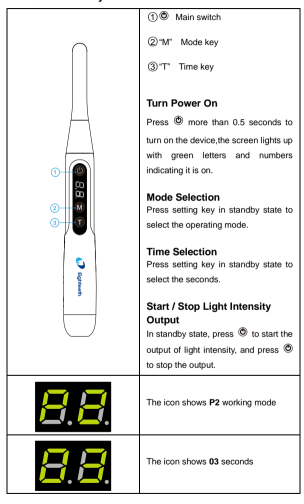


NOTE

- Only the original adapter can be used.Otherwise, the device may be damaged.
- The round connector of the adapter must be pluged into the charging hole at the rear of handpiece in the right way.

5.Use Interface

5.1 Panel key



6.Setting

6.1 Selecting memory mode

Memory mode setting

There are 5 built-in memory programs, namely P1 Normal Mode,P2 High Power Mode,P3 Turbo Mode,P4 Pulse Mode,P5 Ramp Mode. Press setting key to enter the mode setting menu. In the menu, press setting key gently again to change the memory.

Curing Time setting

In the curing time setting menu, press "T" to select different times. The time selection is different under different light intensity:

light intensity mode	time selection(sec)	
P1 Normal Mode	05,10,15,20,25,30,35,40	
P2 High Power Mode	01,03,05,10	
P3 Turbo Mode	01,03,05	
P4 Pulse Mode	05,10,15,20	
P5 Ramp Mode	05,10,15,20	

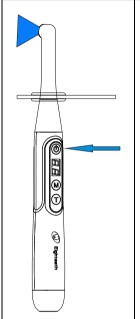


NOTE

 The light intensity of P1 to P5 memory modes are built-in, and the user cannot modify the settings.

7.Operation

7.1 Handpiece operation



In the off state, short press to turn on the device, and then select one mode, and then short press to start the light intensity output, and the time will start counting down. Press to stop the output during the light intensity output.

There is a beep prompt after finishing treatment, or every 5 seconds while working, and the output is automatically turned off after the countdown ends.

In the on state, press 1, then press 6 to shut down.



Warning

- When the device is working, do not directly illuminate the eyes, otherwise it will cause injury.
- Do not directly illuminate the skin, otherwise high temperature burns may occur.
- Before using, please try it outside the oral cavity to ensure that there is no problem with the function of the device.
- Do not disassemble the light source head during treatment.



NOTE

- To prevent the lamp from injurying patients by overheating, after the device is used continuously 300 seconds at P2 High Power Mode and P3 Turbo Mode, it will be prohibited to use the highest light intensity output within 60 seconds.
- When using, the light should be directly irradiated onto the curing dental resins and composites to avoid improper exposure.
- The disposable sleeve and protective light shield are highly recommended.
- If there is any abnormal functioning, stop using the device and report to the distributor.
- Gloves are compulsory during treatment.
- Always disinfect the handpiece and light source head after each treatment.

7.2 Operation mode

	Normal Mode
	Light intensity :1200mW/cm²
	Wavelength: 380nm-515nm
	When is pressed, the set light intensity is
	output immediately. There is a beep prompt after
P1	finishing treatment, or every 5 seconds while working.
	light intensity †
	Time
	High Power Mode
	Light intensity: 2000mW/cm²
	Wavelength: 380nm-515nm
	When 🕲 is pressed, the set light intensity is
	output immediately. There is a beep prompt after
P2	finishing treatment, or every 5 seconds while working.
	light intensity
	Turbo Mode
	Light intensity: 3000mW/cm²
	Wavelength: 380nm-515nm
	When 🕲 is pressed, the set light intensity is
	output immediately. There is a beep prompt after
P3	finishing treatment.
	light intensity 100% Time

Operation Pulse Mode Light intensity: 1200mW/cm² Wavelength: 380nm-515nm When 🕲 is pressed, the set light intensity is output immediately and flashes once every 1 second. There is a beep prompt after finishing treatment, or **P4** every 5 seconds while working. light intensity 100% Ramp Mode Light intensity: 1200mW/cm² Wavelength: 380nm-515nm When (b) is pressed, the light intensity gradually increases from 0mW/cm2 to 1200mW/cm2, and then continues to output 1200mW/cm2. There is a beep prompt after finishing treatment, or every 5 seconds **P5** while working. light intensity 100%

Time

7.3 Charge



When the "E0" low battery prompt appears on the screen, it means the battery is exhausted, please charge it in time. How to plug the adapter is described in the Chapter 4.4 Plug the adapter.



During charging, the charging indication

" appears on the screen dynamically. When the battery is fully charged, the indication " appears on the screen statically.



It takes about 3 hours for full charge, depending on residual battery power and battery state.

It can be recharged 300-500 times, depending on the operating conditions of the device. If there is a significant decline in battery power, please report to the distributor, so as not to affect the curing effect.



NOTE

- When the "E0" low battery warning shows on the screen,the device must be recharged within 15 days, otherwise the battery may cause irreparable damage due to long periods of low power.
- If the device flickers on the screen in standby mode, it indicates that the battery is less than 20%, so please charge it in time.
- Only the original adapter can be used. Otherwise, the device may be damaged.
- The round connector of the adapter must be pluged into the charging hole at the rear of handpiece in the right way.

- If pluging the adapter while the device is working, other functions of the device will forcibly stop, and then the device will enter charging status.
- When the device is not used for a long time, please change the device at least once a month.
- The device automatically enters the shutdown state after 120 seconds without operation. Please press to restart the device.



Warning

 Do not change the battery, only trained technician or distributor can change the battery, the electronic parts will be damaged if use a wrong battery or install with a wrong way.

8.Maintenance

8.1 Foreword

For hygiene and sanitary safety purpose, the handpiece, light source head and protective light shield must be cleaned and disinfected even if the disposable sleeve is used. They should be cleaned and disinfected before each usage to prevent any contamination. This concerns the first use as well as the subsequent uses. Comply with your national guidelines, standards and requirements for cleaning and disinfection.

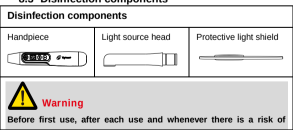
Reprocessing procedures have only limited implications to this dental instrument. The limitation of the numbers of reprocessing procedures is therefore determined by the function / wear of the device. From the processing side there is no maximum number of allowable reprocessing. The device should no longer be reused in case of signs of material degradation.

In case of damage, the device should be reprocessed before sending back to the manufacturer for repair.

8.2 General recommendations

- The user is responsible for the sterility of the product for the first cycle and each further usage as well as for the usage of damaged or dirty instruments, where applicable after sterility.
- For your own safety, please wear personal protective equipment (gloves, safety glasses, etc.).
- Use only a disinfecting solution which is approved for its efficacy (VAH/DGHM-listing, CE marking, and FDA approval) and in accordance with the DFU of the disinfecting solution manufacturer.
- The water quality has to be convenient to the local regulations especially for the last rinsing step or with a washer-disinfector.
- Do not use bleach or chloride disinfectant materials.

8.3 Disinfection components



contamination, disinfect the above components.

Reprocessing Instructions

Preparation before cleaning:

Immediately after using, remove the used disposable sleeve. Disconnect all plug connections. Put the handpiece, light source head and protective light shield in container for transportation.



Warning

Make sure that used sleeves are disposed of as infected waste which is potentially biologically hazardous.

Transportation:

Safe storage and transportation to the reprocessing area to avoid any damage and contamination to the environment.

Cleaning

Wipe the handpiece, light source head and protective light shield surface with a cloth moistened in Ethanol (70 to 80vol%), until the components are free of visible soils. Repeat this step with a new gauze soaked in Ethanol (70 to 80vol%), if necessary.

Dry the components with compressed air.

Disinfection

Wipe he handpiece, light source head and protective light shield surface with a cloth lightly moistened with Ethanol for Disinfection (Ethanol 70 to 80vol%) at least 2min, repeat for 5 times.

Remove any chemical residues by wiping the components clean and dry with a sterile cloth



NOTE

1.Do not use disinfectants other than Ethanol (70 to 80vol%) for disinfection.

2.Make sure no liquid penetrates the handpiece and light source head, otherwise, it will damage the internal parts.

3.Adapter and base should be cleaned and disinfected with a cloth moistened in Ethanol (70 to 80vol%) before first use and after each use.



- 1.Do not disinfect the handpiece and light source head in an autoclave or other sterilization container.
- 2.Do not soak or immerse any part of the handpiece and light source head in liquid.

Storage:

Store the components in a clean and dry place for the next treatment.

9.Error Warning

E0	When the indication "E0" blinks on the screen, it means the battery is exhausted, please charge it in time. If the error warning persists, please contact your local distributor.		
E1	When the indication "E1" blinks on the screen, it indicates that the light source head has failed. Please stop using and contact your local distributor.		
E2	When the indication "E2" blinks on the screen, please stop using and contact your local distributor.		
EH	When the indication "EH" blinks on the screen, it indicates that the number of continuously using High light intensity mode reaches the limit. Please stop the device working for 60 seconds before continuing to use it.		

10.Troubleshooting

When trouble is found, check the following points before contacting your distributor. If none of these are applicable or the trouble is not remedied even after action has been taken, the product may have failed. Please contact your distributor.

Problem	Cause	Solution	
The power is not	The battery is out of power.	Charge the battery.	
turned on.	Handpiece is broken.	Contact your local distributor.	
The device flickers on the screen in standby mode.	The battery is out of power.	Charge the battery.	
	There is no electricity in the outlet.	Check the connection.	
The power LED does not light up	Use a wrong adapter.	Use the original adapter.	
when charging.	The adapter is damaged.	Contact your local distributor.	
	Handpiece is broken.	Contact your local distributor.	
No sound.	Handpiece is broken.	Contact your local distributor.	
Insufficient light intensity.	There are resin or other contaminants on the surface of the lamp lens.	Cleaning the lamp head residue.	
"E0" error warning.	The battery power is too low.	Charge the battery.	
"E1" error warning.	The circuit of the light source head is open.	Check the circuit of the light source head.If the error warning persists ,please contact your local distributor.	

10 Troubleshooting

"E2" error warning.	The circuit of the light source head is short.	Contact your local distributor.
	Handpiece is broken.	Contact your local distributor.
"EH" error warning.	The number of using High light intensity mode reachs the limit.	Keep the device from working for 60 seconds before continuing.

11.Technical Data

Manufacturer	Changzhou Sifary Medical Technology Co.,Ltd		
Model	CuringPen-E		
Dimensions	19cm×16.5cm×7.5cm±1cm		
Weight	660g±10%		
Power supply	Lithium ion battery: DC 3.7V, 1400mAh, ±10%		
Charger power supply	AC 100-240 V, ±10%		
Charger power output	5V 1A		
Power Frequency	50/60Hz, ±10%		
Charger nominal power input	0.2A		
P1:1200±15%mW/cm² P2:2000±15%mW/cm² Light intensity P3:3000±15%mW/cm² P4:1200±15%mW/cm² P5:1200±15%mW/cm²			
Wavelength	380nm-515nm		
Electrical safety class	Class II		
Applied part	В		
Operation mode	Intermittent operation 5mins. ON / 1min. OFF		
	Use: in enclosed spaces		
	Ambient temperature: 5°C ~ 40 °C		
Operation conditions	Relative humidity: <80%		
Operation containons	Operating altitude < 3000m above sea level		
	Atmospheric pressure: 70kPa ~ 106kPa		
	Ambient temperature: -20 °C ~ +55 °C		
Transport and storage	Relative humidity: 20% ~ 80 %		
conditions	Atmospheric pressure: 70kPa ~ 106kPa		

12.EMC Tables

This product has no essential performance.

Guidance and manufacturer's declaration electromagnetic emissions

The **CuringPen-E** is intended for use in the electromagnetic environment specified below. The customer or the user of the **CuringPen-E** should assure that it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment-guidance
RF emissions CISPR 11	Group 1	The CuringPen-E 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 IEC61000-3-2	Class A	The CuringPen-E is suitable for
Voltage fluctuations/flicker emissions IEC 61000-3-3	Complies	use in professional healthca re facility environment.

Guidance	and	and manufacturer's		declaration -	
electromagnetic immunity					
The CuringPer	The CuringPen-E is intended for use in the electromagnetic environment				
specified below	specified below. The customer or the user of the CuringPen-E should				
assure that it is used in such an environment.					
Immunity IEC 60601 Compliance Electromagnetic					
test	test le	vel	level	environment -	
				guidance	

	12	EMC Tables	
Electrostatic	+/- 8 kV	+/- 8 kV contact	Floors should be
discharge	contact		wood, concrete or
(ESD) IEC			ceramic tile. If
61000-4-2	+/- 2 kV, +/- 4	+/- 2 kV, +/- 4	floors are covered
	kV, +/- 8 kV,	kV, +/- 8 kV, +/-	with synthetic
	+/- 15 kV air	15 kV air	material, the
			relative humidity
			should be at least
			30 %.
Electrical fast	±2kV	±2kV	Mains power
Transients	100kHz	100kHz	quality should be
/bursts	repetition	repetition	that of a typical
IEC 61000-4-	frequency	frequency	commercial or
4			hospital
			environment.
Surge	Line to line:	Line to line:	Mains power
IEC 61000-4-	±0.5kV, ±1kV	±0.5kV, ±1kV	quality should be
5		·	that of a typical
			commercial or
			hospital
			environment.
Voltage dips,	0% UT; 0.5	0% UT; 0.5	Mains power
short	cycle	cycle	quality should be
interruptions	at 0°, 45°,	at 0°, 45°, 90°,	that of a typical
and	90°, 135°,	135°. 180°.	commercial or
voltage	180°, 225°,	225°, 270°, and	hospital
variations on	270°, and	315°	environment. If the
power supply	315°		user of devices
lines			require continued
IEC 61000-4-	0% UT; 1		operation during
11	cycle and	0% UT; 1 cycle	power mains
[.,	70% UT:	and 70% UT:	interruptions, it is
	25/30 cycles	25/30 cycles	recommended
	sine phase at	sine phase at 0°	that devices be
	0°	Sine priase at 0	powered form an
			uninterruptible
	0% UT:		·
	0% UT; 250/300	0% UT;	power supply or
	230/300	070 01,	a battery

12 EMC Tables

	cycle		250/300 cycle		
Power	30 A/m		30 A/m	Power frequency	
frequency	50Hz c	or	50Hz or 60Hz	magnetic field	
magnetic	60Hz			should be at levels	
field IEC				characteristic of a	
61000-4-8				typical	
				location in a	
				typical	
				commercial or	
				hospital	
				environment.	

Note: UT: rated voltage(s); E.g. 25/30 cycles means 25 cycles at 50Hz or 30 cycles at 60Hz

Guidance and manufacturer's declaration - electromagnetic immunity

The CuringPen-E is intended for use in the electromagnetic environment specified below. The customer or the user of the CuringPen-E should assure that it is used in such an environment.

Proximity	IEC 61000-4-	Compliance	Electromagnetic		
magnetic	39 test level	level	environment -		
fields			guidance		
Proximity	65A/m	65A/m	Power frequency		
magnetic	134.2kHz		magnetic field		
fields	Pulse		should be at levels		
	modulation		characteristic of a		
	2.1 kHz		typical		
Proximity	7.5A/m	7.5A/m	location in a		
magnetic	13.56MHz		typical		
fields	Pulse		commercial or		
	modulation		hospital		
	50 kHz		environment.		

Guidance and manufacturer's declaration – electromagnetic immunity

12 EMC Tables

The **CuringPen-E** is intended for use in the electromagnetic environment specified below. The customer or the user of the **CuringPen-E** should assure that it is used in such an environment.

	IEC 60601	Compliance	Electromagnetic			
Immunity test	test level	level	environment -			
		12.12.	guidance			
Conducted dis-	3 V	3 V	Portable and			
turbances	0.15 MHz – 80	0.15 MHz –	mobile RF			
induced by RF	MHz, 6 V in	80 MHz, 6 V	communications			
fields	ISM bands be-	in ISM bands	equipment should			
IEC 61000-4-6	tween 0.15	be-tween	be usedno closer			
	MHz and 80	0.15 MHz	to any part of the			
	MHz, 80 %	and 80 MHz,	CuringPen-E,			
	AM at 1 kHz	80 % AM at 1	including cables,			
		kHz	than the			
			recommended			
			separation			
			distance			
			calculated from			
			the equation			
Radiated RF	3 V/m, 80	3V/m	applicable to the			
	MHz – 2,7		frequency of the			
EM fields	GHz, 80 %		transmitter.			
IEC 61000-4-3	AM at 1 kHz					
			Recommended			
			minimum			
			separation			
Proximity fields	Refer to table	Complies	distances			
from RF	"Recommend		Refer to table			
wireless	ed minimum		"Recommended			
communicatio	separation		minimum			
n equipment	distances"		separation			
IEC 61000-4-3			distances"			
120 01000-4-3						

Recommended minimum separation distances

Nowadays, many RF wireless equipment have being used in various healthcare locations where medical equipment and/or systems are used. When they are used in close proximity to medical equipment and/or systems, the medical equipment and/or systems' basic safety and essential performance may be affected. The **CuringPen-E** has been tested with the immunity test level in the below table and meet the related requirements of IEC 60601-1-2:2014+A1:2020. The customer and/or user should help keep a minimum distance between RF wireless communications equipment and the **CuringPen-E** as recommended below.

Test				Maxim	Dist	Immun
freque	Band	Service	Modulation	um	anc	ity test
ncy	(MHz)	Service	iviodulation	power	е	level
(MHz)				(W)	(m)	(V/m)
385	380-390	TETRA 400	Pulse modulation 18Hz	1.8	0.3	27
450	430-470	GMRS 460 FRS 460	FM ± 5 kHz deviation 1 kHz sine	2	0.3	28
710		LTE Band	Pulse			
745	704-787	13, 17	modulation	0.2	0.3	9
780		10, 17	217Hz			
810		GSM				
870		800/900,				
930	800-960	TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse modulation 18Hz	2	0.3	28
1720		GSM				
1845		1800;				
1970	1700- 1990	CDMA 1900; GSM 1900; DECT;	Pulse modulation 217Hz	2	0.3	28

		12	LIVIC Tables			
		LTE Band 1, 3, 4, 25; UMTS				
2450	2400- 2570	Bluetooth , WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation 217Hz	2	0.3	28
5240 5500 5785	5100- 5800	WLAN 802.11 a/n	Pulse modulation 217Hz	0.2	0.3	9



Warning

 Use of accessories and cables other than those specified or provided by the manufacturer of CuringPen-E could result in increased electromagnetic emissions or decreased electromagnetic immunity of CuringPen-E and result in improper operation.

Cable information:

Cable Name	Cable Length (m)	Shielded or not	Remark
Adapter Cable	1.2	No	/

- 2. Use of CuringPen-E adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, CuringPen-E and the other equipment should be observed to verify that they are operating normally.
- 3. Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm(12 inches) to any part of the CuringPen-E, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.

13.Statement

Service Life

The service life of CuringPen-E series products is 5 years.

Maintenance

MANUFACTURE will provide circuit diagrams, component part lists, descriptions, calibration instructions to assist to SERVICE PERSONNEL in parts repair.

Disposal

The package should be recycled. Metal parts of the device are disposed as scrap metal. Synthetic materials, electrical components, and printed circuit boards are disposed as electrical scrap. The lithium batteries are disposed as special refuse. Please deal with them according to the local environmental protection laws and regulation.

Rights

All rights of modifying the product are reserved to the manufacturer without further notice. The pictures are only for reference. The final interpretation rights belong to CHANGZHOU SIFARY MEDICAL TECHNOLOGY CO., LTD. The industrial design, inner structure, etc, have claimed for several patents by SIFARY, any copy or fake product must take legal responsibilities.

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