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# PM-900

Multi-parameter  
Patient Monitor



## Features:

- 12.1 inches high resolution color TFT screen
- User friendly and configurable interface design
- Intuitive human voice alarms
- Powerful data review:
  - 480 hours graphic and tabular trends of all parameters
  - 1000 pieces of NIBP record storage
  - 700 pieces of alarm review
- Built-in rechargeable Li-ion battery for up to 5 hours of continuous work
- VGA output for external display (optional)
- Wire/wireless central monitoring system solution (optional)

Standard Parameters: 3/5-lead ECG, PR, RESP, NIBP, SpO<sub>2</sub>, TEMP  
 Optional: Recorder, 2-IBP, EtCO<sub>2</sub>(mainstream, sidestream),  
 central monitoring system

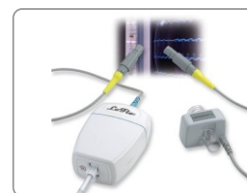
## Respironics EtCO<sub>2</sub> (optional)



CAPNOSTAT5  
mainstream module



LoFlo sidestream module



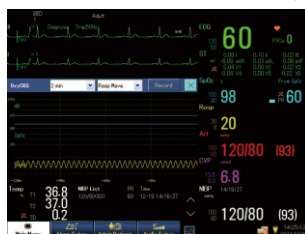
EtCO<sub>2</sub>(Respironics)  
"Plug and Play"

## Display Modes

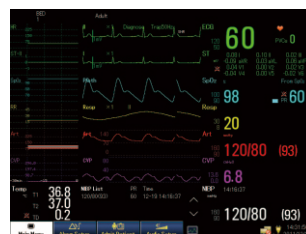
- Flexible display solutions to meet different needs in clinical application scenarios



Big font interface



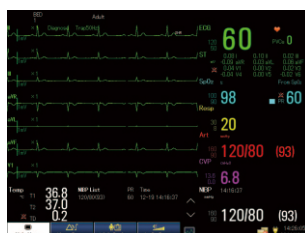
OxyCRG interface



Mini-trend interface



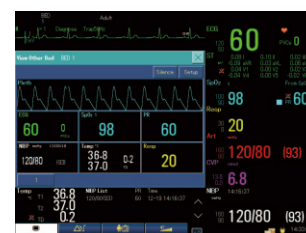
Drug dose calculation interface



7-lead full screen interface

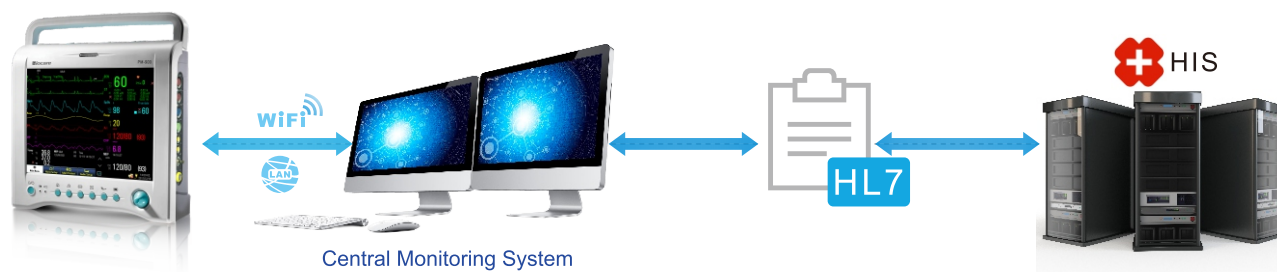


7-lead half screen interface



Bed to bed view interface

## HL7 supports the patient data to be transferred to HIS



## Shenzhen Biocare Bio-Medical Equipment Co., Ltd.

**Add:** #16-1, Jinhui Road, Jinsha Community, Kengzi Sub-District, Pingshan New District, 518122 Shenzhen, P.R. CHINA

**Tel:** +86-755-3661 5333

**Fax:** +86-755-2796 0643

**E-mail:** sales@biocare.com.cn

**Website:** www.biocare.com.cn

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 ENG-CATA-V1.5-20170829

## Appendix B Product Specifications

### B.1 Safety Specifications

#### B.1.1 Product Classification

For classification of this series of monitors comply with IEC60601-1, please refer to Table B.1.

Table B.1 Module Classification

Components	Type of Protection Against Electric Shock	Degree of Protection Against Electric Shock	Degree of Protection Against harmful ingress of water	Degree of Protection Against hazards of Explosion	Mode of Operation
Main unit	I	Not marked	IPX1	Not suitable	Continuous
ECG (Resp) Module	NA	CF(*)			
IBP Module (Optional)					
NIBP Module					
Temp Module					
SpO <sub>2</sub> Module					
CO <sub>2</sub> Module (Optional)		BF(*)			

#### ATTENTIONS:

- I: Class I Equipment
- BF: Type BF applied part (The symbol ‘\*’ indicates the availability of defibrillation-proof function).
- CF: Type CF applied part (The symbol ‘\*’ indicates the availability of defibrillation-proof function).
- NA: Not applicable.
- IPX1: Liquid intake protection grade is first level.
- Not suitable: the equipment is not suitable for use in an environment with air, oxygen or nitrous oxide mixed with flammable anesthetic gas.

**B.1.2 Environment Specifications**

Equipment Environment			
Item	Temperature	Humidity	Atmospheric Pressure
Operating	0°C ~ 40°C (32°F ~ 104°F) If the machine includes CO <sub>2</sub> module, the operating temperature is 5 °C ~ 40 °C (41 °F ~ 104 °F )	15% ~ 80%, Non-Condensing	442.5 mmHg ~ 805.5 mmHg (59 kPa ~ 107.4 kPa)
Storage&Transport	-20°C ~ +55°C (-4°F ~ 140°F)	≤93%, Non-Condensing	525 mmHg ~ 795 mmHg (70 kPa ~ 106 kPa)

**B.1.3 Power Specifications**

(AC) Input Voltage	100 V ~ 240 V
Input Power	160 VA
Frequency	50 Hz/60 Hz (Allowable frequency error ±1Hz)
Fuse	3.15A/250V
Safety Classification	Class I, Type BF, CF

**B.2 Physical Specifications**

Host	
Weight	Approx. 4.0 kg
Size (L×W×H)	312 mm×139 mm×305 mm

**B.3 Hardware Specifications**

Display	
Type	TFT LCD Screen
Dimensions	12.1 inches
Resolution	800×600 pixels (12 inches monitor)
Screen Brightness	10-level, adjustable
LCD View Angle	Horizontal / vertical view angle at least 150°/120°
Recorder (Optional)	
Type	Thermal array recorder
Horizontal Resolution	16 dots/mm (Paper Speed: 25.0 mm/s)
Vertical Resolution	8 dots/mm
Printing Paper Size	50 mm×20 mm
Paper Speed	12.5 mm/s; 25.0 mm/s; 50.0 mm/s
Waveform	Max. 3 waveforms
Battery	
Dimensions	147.5 mm×72.5 mm×19.5 mm



Weight	0.38 kg
Type	Rechargeable lithium battery
Rated voltage	14.8 V
Battery Capacity	4.4 Ah
Length of Power Supply	In environment temperature 25 °C and in standard configuration (the SpO <sub>2</sub> sensor connects ,the ECG cable and Temp cable disconnect, the “Measure Mode” of NIBP is “Auto” and the “Interval” is 15 minutes), the continuous working time of the battery is not less than <b>5 hours</b> .
Time for recharging battery to 90% from zero power state	The charging time is not more than 12 hours to charge the battery to 90%.
Shutdown Delay	0 s, 0.5 s, 1 s, 1.5 s, 2 s
<b>Host LED</b>	
Physiological Alarm Indicator Lamp	1 (Dual color yellow & red)
Battery Power Indicator Lamp	1 (Green)
Speaker	Give out alarm sound (45 dB~85 dB), keystroke sound and QRS sound. Alarm sound complies with IEC 60601-1-8
<b>Interface</b>	
Power	1 AC power port
Network	Standard RJ45 network port, which can network with the central monitoring system and transmit all the patient monitored data to the central monitoring system.
USB	USB disk supported. For the manufacturer to upgrade and service the application software, and export data (Structurally 1 USB host interfaces supported)
VGA	Supported, for connection of external display
Equipotential Terminal Port	1 piece
<b>ECG Analog Signal Output</b>	
Bandwidth (-3 dB, reference 10Hz)	Surgery mode: 1 Hz~15 Hz Monitor mode: 0.5 Hz~40 Hz Diagnose mode: 0.05 Hz~150 Hz
Max. Transmission Delay	25ms (Wave filter closed under diagnose mode)
Sensitivity	1 V/mV ±5%
Accuracy of input signal reproduction	Using the method described in 4.2.7.1 of AAMI EC11 to test the overall system error, which is within ±5%; Using method A and D described in 4.2.7.1 of AAMI EC11 to test frequency response. Because of sampling characteristics and the asynchronism between sample rate and signal rate of the ECG module, digital systems may produce a noticeable modulating effect from one cycle to the next, particularly in pediatric recordings. This phenomenon, which is not physiologic, shall be clearly described in the operator's and service manuals.
<b>IBP Analog Signal Output</b>	

## Product Specifications

Bandwidth (-3 dB, reference 10Hz)	0 Hz~50 Hz
Max. Transmission Delay	30 ms (Filter closed)
Sensitivity	0.01 V/mmHg $\pm$ 5%

## B.4 Data Storage

Trend Data	Short Trend (Trend Window Time 4 min, 40 min, 2 h) Resolution of Trend Chart 5 s, 30 s, 1 min, 10 min): Max. storage time: 72h. Long trend (Trend Window Time 4 h, 16 h, 32 h, 48 h) Resolution of Trend Chart 15 min, 30 min, 1 h, 2 h, 3 h): Max. storage time: 480h.
Parameter Alarm Event	700 parameter alarm events and manual events, as well as the parameter waveform related to the occurring time, wave length 10s
NIBP Measuring Result	Max. 1000 groups
Single-Channel ECG Waveform	Max. 2h
Holographic Waveform	Max. 2 min (Power cutoff storage not supported)

## B.5 Wireless Network (Optional)

Applicable Standard	IEEE 802.11b/g/n (2.4G)	IEEE 802.11a/n (5G)
Frequency Range	2.412 GHz~2.472 GHz	4.9 GHz~5.975 GHz
Band Width	20~40MHz	20~40MHz
Radiated Power	+18dBm	+13.5dBm
Signal Path	1-13 (China)	
Type and Frequency Characteristics of the Modulation	CCK/DSSS/OFDM/MCS7/MCS0	

## B.6 Measuring Specifications

### B.6.1 ECG Monitoring

Input Mode	3-Lead ECG input (Optional) 5-Lead ECG input (Standard)
Lead Selection	I , II , III(Optional) I , II , III, aVR, aVL, aVF, V
Lead Standard	AHA, IEC
Measuring Range of Heart Rate	Adult: 15 bpm~300 bpm Pedi: 15 bpm~350 bpm Neonate: 15 bpm~350 bpm
Heart Rate Display Tolerance	$\pm$ 1% or $\pm$ 1 bpm, whichever is higher

Sensitivity	1.25 mm/mV ( $\times 1/8$ ), 2.5 mm/mV ( $\times 1/4$ ), 5.0 mm/mV ( $\times 1/2$ ), 10.0 mm/mV ( $\times 1$ ), 20.0 mm/mV ( $\times 2$ ), 40.0 mm/mV ( $\times 4$ ), Auto. Error: $\pm 5\%$
Resolution Stability	The resolution change 1 minute after the instrument is powered on does not exceed 0.66% per minute. The total change within 1h does not exceed any available fixed gain setting by $\pm 10\%$ .
Sweep Speed	6.25 mm/s, 12.5 mm/s, 25.0 mm/s, 50.0 mm/s. Error: $\pm 10\%$
Noise Level	$\leq 30 \mu V_{p-p}$
Input Circuit Current	$\leq 0.1 \mu A$
Input Impedance	$\geq 2.5 M\Omega$
Patient Leakage Current	$< 10 \mu A$
ESU Proof	Cutting Mode: 300 W Coagulation Mode: 100 W Recovery Time: $\leq 10$ s
ESU Noise Inhibition	Tested acc. to 5.2.9.14 of ANSI/AAMI EC 13:2002: 1) The ECG signal track does not disappear; 2) Change in heart rate does not exceed 10% of the heart rate when the electrosurgical knife is not activated.
CMRR	Diagnose Mode: $\geq 89$ dB Surgery & Monitor Mode: $\geq 100$ dB
Time Constant	Monitor Mode: $\geq 0.3$ s Diagnose Mode: $\geq 3.2$ s
Frequency Response	Surgery Mode: 1 Hz-15 Hz; Monitor Mode: 0.5 Hz-40 Hz; Diagnose Mode: 0.05 Hz-150 Hz.
ECG Parameter Frequency Characteristics	Surgery Mode: Meet ( $+0.4$ dB $\sim$ $-3.0$ dB)) requirements at 15 Hz. Monitor Mode: Meet ( $+0.4$ dB $\sim$ $-3.0$ dB)) requirements at 0.5 Hz $\sim$ 40 Hz. Diagnose Mode: Meet ( $+0.4$ dB $\sim$ $-1.0$ dB)) requirements at 0.05 Hz $\sim$ 60 Hz.  Meet ( $+0.4$ dB $\sim$ $-3.0$ dB)) requirements at 61 Hz $\sim$ 150 Hz.
Notch	Monitor & Surgery Mode: notch filter automatically activated at 50 Hz/60 Hz Diagnose Mode: Notch filter manually activated or deactivated at 50 Hz/60 Hz
Range of Electrode Polarized Voltage	$\pm 300$ mV d.c.
Lead Fall Testing Current	Measuring Electrode: $< 0.1 \mu A$ Drive Electrode $< 1 \mu A$
<b>Pacemaker Pulse</b>	
Pacemaker Pulse Display Capacity	Pace-making mark can be displayed for the following pacemaker pulses: Pulse Amplitude: $\pm 2$ mV $\sim$ $\pm 100$ mV Pulse Width: 0.1 ms $\sim$ 2 ms Pulse Rise Time: 10 $\mu s$ $\sim$ 100 $\mu s$ Pacemaker pulse should be no overshoot
Pacemaker Pulse Suppression Capacity	The monitor can inhibit the pacemaker pulse that conforms to the following conditions: Pulse Amplitude: $\pm 2$ mV $\sim$ $\pm 100$ mV Pulse Width: 0.1 ms $\sim$ 2 ms

## Product Specifications

	Pulse Rise Time: 10 $\mu$ s $\sim$ 100 $\mu$ s Pacemaker pulse should be no overshoot
Pacemaker suppression to quick ECG signal	$\leq 5$ V/s
<b>Alarm Limit Specifications</b>	<b>Range</b>
Upper Limit of ECG Heart Rate	Alarm upper limit for adult: (Lower limit+2) bpm $\sim$ 300 bpm Alarm upper limit for pedi: (Lower limit+2) bpm $\sim$ 350 bpm Alarm upper limit for neonate: (Lower limit+2) bpm $\sim$ 350 bpm
Lower Limit of ECG Heart Rate	Alarm lower limit for adult: 15 bpm $\sim$ (Upper limit-2)bpm Alarm lower limit for pedi: 15 bpm $\sim$ (Upper limit-2)bpm Alarm lower limit for neonate: 15 bpm $\sim$ (Upper limit-2)bpm
Resolution	$\pm 1$ bpm
Accuracy	The tolerance of alarm limit setting is $\pm 1$ bpm. In addition, the ECF signal alarm below the publicized lower limit of the alarm will not fail. If the alarm is not disabled, the alarm will not fail if you enter the ECG input signal higher than the upper limit of alarm up to 300 bpm (350 bpm for neonate and pedi).
<b>HR</b>	
Heart Rate Testing Amplitude	$\pm 0.3$ mV $\sim$ $\pm 5$ mV
Resolution	1 bpm
Alarm Time for Tachycardia	Acc. to ANSI/AAMI EC13:2002 Part 4.1.2.1 g). 4ah-Range: 11 s 4a-Range: 11 s 4ad-Range: 11 s 4bh-Range: 11 s 4b-Range: 11 s 4bd-Range: 11 s
Heart Rate Average	Acc. to ANSI/AAMI EC13:2002 Part 4.1.2.1 g). The average heart rate is obtained by the method below: If the interval of the last continuous 3 RR is higher than 1200ms, the heart rate is averaged based on the most recent 4 RR intervals; otherwise, the heart rate is averaged based on the most recent 12 RR intervals. The heart rate displayed on the screen is refreshed every second.
Response to Irregular Rhythm of the heart	Acc. to ANSI/AAMI EC13:2002 Part 4.1.2.1 e). The heart rate displayed after 20s stabilizing period is: 3a (Ventricular bigeminy) $\sim$ 80 $\pm$ 1bpm 3b (Slow alternating ventricular bigeminy) $\sim$ 60 bpm $\pm$ 1 bpm 3c (Rapid alternating ventricular bigeminy) $\sim$ 120 bpm $\pm$ 1 bpm 3d (Bidirectional systoles) $\sim$ 90 bpm $\pm$ 6 bpm
Response Time to Heart Rate Change	Acc. to ANSI/AAMI EC13:2002 Part 4.1.2.1 f). Increase of heart rate: response time $\leq 11$ s Decrease of heart rate: response time $\leq 11$ s
High T-wave Suppression Capacity	Acc. to ANSI/AAMI EC13:2002 Part 4.1.2.1 c). The heart rate monitor inhibits all T-waves with amplitude lower than 1.2 mV, 100msQRS wave groups, T-wave period 180 ms and QT period 350ms.



Arrhythmia Type	<p>a) Monitoring type: Asystole, VFib/VTac, VTac, Ventricular bradycardia, Extreme-Tachy, Extreme-Brady, Non-Sustained VT, PVC, Tachycardia, Bradycardia, VR(ventricular rhythm), V-Bigeminy, V-Trigeminy, Irr.Rhythm, PVCs/min, Run PVCs &gt; 2, Couplet, R on T, Multiform, HeartBeat Pause, Missed Beats</p> <p>b) Pace-making: Pacemaker not captured (PNC), Pacemaker not paced (PNP).</p>
<b>ST Interval Measuring</b>	
Range	(-2.0 mV)~(+2.0 mV)
Accuracy	Measuring Tolerance: measuring tolerance within (-0.8 mV)~(+0.8 mV) is $\pm 0.02$ mV or $\pm 10\%$ , whichever is higher. It not defined for other ranges.
ST Interval Updating Interval	A single heart beat interval or 1s, whichever is higher.

### B.6.2 Respiration (Resp) Monitoring

Measuring Method	Chest Impedance Method
Measuring Lead	Lead I and II for selection. Lead I defaulted.
Respiration Exciting Waveform	< 300 $\mu$ A, Sine signal, 62.8 kHz ( $\pm 10\%$ )
Range of Respiration Impedance	0.5 $\Omega$ ~3 $\Omega$
Range of Base Impedance	250 $\Omega$ -2000 $\Omega$ (Use of ECG cable with 1k $\Omega$ resistor)
Differential Input Impedance	> 2.5 M $\Omega$
Brand width	0.2 Hz~2 Hz (-3 dB)
Waveform Sensitivity	$\times 1/4$ , $\times 1/2$ , $\times 1$ , $\times 2$ , $\times 4$ , Auto
Sweep Speed	6.25 mm/s; 12.5 mm/s; 25.0 mm/s
Resolution	1 rpm
Accuracy	$\pm 2$ rpm
Asphyxia Alarm	Off, 10 s, 15 s, 20 s, 25 s, 30 s, 35 s, 40 s
<b>RR</b>	
Range	<p>Monitoring Range for adult: 0 rpm~120 rpm</p> <p>Monitoring Range for pedi: 0 rpm~150 rpm</p> <p>Monitoring Range for neonate: 0 rpm~150 rpm</p>
Resolution	1 rpm
Respiration Monitoring Tolerance	<p>Within 7 rpm~150 rpm, the measuring error is <math>\pm 2</math> rpm or <math>\pm 2\%</math>, whichever is higher.</p> <p>The tolerance is not defined for other ranges.</p>
Asphyxia Alarm Tolerance	Within 10 s~40 s (Increase/decrease by 5s for each rotation of the knob), the asphyxia alarm tolerance is $\pm 5$ s.

Alarm Limit Specifications	Range
RR Upper Limit	Alarm upper limit for adult: (Lower limit+2) rpm ~ 100 rpm Alarm upper limit for pedi: (Lower limit+2) rpm ~ 100 rpm Alarm upper limit for neonate: (Lower limit+2) rpm ~ 100 rpm
RR Lower Limit	Alarm lower limit for adult: 0 rpm ~ (Upper limit-2) rpm Alarm lower limit for pedi: 0 rpm ~ (Upper limit-2) rpm Alarm lower limit for neonate: 0 rpm ~ (Upper limit-2) rpm

**B.6.3 SpO<sub>2</sub> Monitoring**

Alarm Limit Specifications	Range
SpO <sub>2</sub> Upper Limit	(Lower limit+1)% ~ 100%
SpO <sub>2</sub> Lower Limit	80% ~ (Upper limit-1)%
Alarm Tolerance	±1% of the setting
Sensing element	Optical power <15 mW Red light wavelength: 658 nm~664 nm, infrared light: 897 nm~915 nm Information on the wavelength range is particularly useful for clinicians (e.g. in optical dynamic therapy)
Monitoring Parameters	SpO <sub>2</sub> and Pulse Rate (PR)
Range	0% ~ 100%
Resolution	1%
Data update period	1 s
Accuracy	Within 70% ~ 100%, the measuring tolerance is ±2%. Within 0% ~ 69%, the measuring tolerance is not defined.

**B.6.4 PR Specifications**

Alarm Limit Specifications	Range
PR Upper Limit	Alarm upper limit for adult: (Lower limit+2) bpm ~ 300 bpm Alarm upper limit for pedi: (Lower limit+2) bpm ~ 300 bpm Alarm upper limit for neonate: (Lower limit+2) bpm ~ 300 bpm
PR Lower Limit	Alarm lower limit for adult: 25 bpm ~ (Upper limit-2)bpm Alarm lower limit for pedi: 25 bpm ~ (Upper limit-2)bpm Alarm lower limit for neonate: 25 bpm ~ (Upper limit-2)bpm

**PR from SpO<sub>2</sub> Module**

Range	30 bpm ~ 300 bpm
Resolution	1 bpm
Measuring Tolerance	±2 bpm
Average Time	8 s

**PR from IBP**

Range	30 bpm~350 bpm
Resolution	1 bpm
Measuring Tolerance	30 bpm~200 bpm: $\pm 1$ bpm or $\pm 1\%$ , whichever is higher; 201 bpm~350 bpm: $\pm 2\%$ .

**B.6.5 NIBP Monitoring**

Measuring Method	Automatic oscillometric method				
Safety Requirements	Acc. to ANSI/AAMI SP-10 Non-invasive Automated Blood Pressure Monitor, Part 4.4				
Work Mode	Manual, Auto, STAT Measuring				
Measuring Time under Continuous Mode	5 min				
Measuring Interval under Auto Mode	1 min, 2 min, 3 min, 4 min, 5 min, 10 min, 15 min, 30 min, 60 min, 90 min, 2 h, 4 h, 3 h, 8 h, Timer interval error: < 10 s				
Resolution	1 mmHg (0.133kPa)				
Nominal Range of Monitoring	Blood Pressure (unit)		Adult	Pedi	Neonate
	Systolic Pressure	mmHg	40~270	40~200	40~135
		kPa	5.3~35.9	5.3~26.6	5.3~18.0
	Mean Pressure	mmHg	20~230	20~165	20~110
		kPa	2.7~30.6	2.7~22.0	2.7~14.7
	Diastolic Pressure	mmHg	10~210	10~150	10~100
		kPa	1.3~27.9	1.3~20.0	1.3~13.3
Range of Initial Inflation Pressure Setting	Adult: 80 mmHg, 100 mmHg, 120 mmHg,140 mmHg,160 mmHg,180 mmHg, 200mmHg,220 mmHg,240 mmHg Pedi: 80 mmHg, 100 mmHg, 120 mmHg, 140 mmHg, 160 mmHg, 180 mmHg,200 mmHg Neonate: 60 mmHg, 80 mmHg, 100 mmHg, 120 mmHg, 145 mmHg				
Default of Initial Inflation Pressure	Adult: 160 mmHg (21.3 kPa) Pedi: 140 mmHg (18.6 kPa) Neonate: 100 mmHg (13.3 kPa)				
Measuring Tolerance of Pressure Source Testing	±3 mmHg (±0.4 kPa)				
Overpressure Protection	Adult state: when the pressure in cuff exceeds 300 mmHg (39.9 kPa)±3 mmHg (0.4 kPa), the control valve shall relieve the pressure. Pedi state: when the pressure in cuff exceeds 240 mmHg (31.9 kPa)±3 mmHg (0.4 kPa), the control valve shall relieve the pressure. Neonate state: when the pressure in cuff exceeds 147 mmHg (19.6 kPa)±3 mmHg (0.4 kPa), the control valve shall relieve the pressure.				

Alarm Limit Specifications	Range
Upper Limit of Systolic Blood Pressure	Adult: (Lower limit+5)mmHg~270 mmHg ( (Lower limit+0.7)kPa~35.9 kPa) Pedi: (Lower limit+5)mmHg~200 mmHg ( (Lower limit+0.7)kPa~26.6 kPa) Neonate: (Lower limit+5)mmHg~135 mmHg ( (Lower limit+0.7)kPa~18.0 kPa)
Lower Limit of Systolic Blood Pressure	Adult: 40 mmHg~ (Upper limit-5)mmHg (5.3 kPa~ (Upper limit -0.7)kPa) Pedi: 40 mmHg~ (Upper limit-5)mmHg (5.3 kPa~ (Upper limit-0.7)kPa) Neonate: 40 mmHg~ (Upper limit-5)mmHg (5.3 kPa~ (Upper limit-0.7)kPa)
Upper Limit of Mean Blood Pressure	Adult: (Lower limit+5)mmHg~230 mmHg ( (Lower limit+0.7)kPa~30.6 kPa) Pedi: (Lower limit+5)mmHg~165 mmHg ( (Lower limit+0.7)kPa~21.9 kPa) Neonate: (Lower limit+5)mmHg~110 mmHg ( (Lower limit+0.7)kPa~14.6 kPa)
Lower Limit of Mean Blood Pressure	Adult: 20 mmHg~ (Upper limit-5)mmHg (2.7 kPa~ (Upper limit-0.7)kPa) Pedi: 20 mmHg~ (Upper limit-5)mmHg (2.7 kPa~ (Upper limit-0.7)kPa) Neonate: 20 mmHg~ (Upper limit-5)mmHg (2.7 kPa~ (Upper limit-0.7)kPa)
Upper Limit of Diastolic Blood Pressure	Adult: (Lower limit+5)mmHg~210 mmHg ( (Lower limit+0.7)kPa~27.9 kPa) Pedi: (Lower limit+5)mmHg~150 mmHg ( (Lower limit+0.7)kPa~20.0 kPa) Neonate: (Lower limit+5)mmHg~100 mmHg ( (Lower limit+0.7)kPa~13.3 kPa)
Lower Limit of Diastolic Blood Pressure	Adult: 10 mmHg~ (Upper limit-5)mmHg (1.3 kPa~ (Upper limit-0.7)kPa) Pedi: 10 mmHg~ (Upper limit-5)mmHg (1.3 kPa~ (Upper limit-0.7)kPa) Neonate: 10 mmHg~ (Upper limit-5)mmHg (1.3 kPa~ (Upper limit-0.7)kPa)

### B.6.6 Temperature (Temp) Monitoring

Range	0°C ~ 50°C (32°F ~ 122°F)
Measuring Method	Thermal resistance method
Accuracy	The measuring tolerance is $\pm 0.1^{\circ}\text{C}$ (exclusive of probe tolerance)
Updating Interval	1 s
Nominal Resistance of Temp. Sensor	2252 $\Omega$ (25°C)
Type of Temp. Sensor	YSI400 Sensor or its Compatible Sensor (Precision $\pm 0.1^{\circ}\text{C}$ )
Channel Number	2 channels
Resolution	0.1°C
Alarm Indication	Audible & visual alarm, data and parameter blinking, alarm message displayed in the screen, 3 levels of alarm.
Alarm Limit Specifications	Range (°C)
Upper Limit	(Lower Limit +1)°C ~ 50 °C
Lower Limit	0 °C ~ (Upper Limit -1)°C

**B.6.7 IBP Monitoring**

Measuring Method		Invasive direct measuring
Volume displacement (Abbott)		<0.04 mm <sup>3</sup> /100mmHg
<b>IBP</b>		
Measuring Range		-50 mmHg~350 mmHg
Resolution		1 mmHg
Accuracy		±2% or ±1 mmHg, whichever is higher (exclusive of the sensor)
Updating Interval		1 s
<b>Alarm Limit Specifications</b>		<b>Range</b>
Art P1 P2	Upper Limit of Systolic Blood Pressure	(Lower limit+2) mmHg~350 mmHg ((Lower limit+0.3)kPa~46.7 kPa)
	Upper Limit of Mean Blood Pressure	
	Upper Limit of Diastolic Blood Pressure	
PA	Upper Limit of Systolic Blood Pressure	(Lower limit+2) mmHg~120 mmHg ( (Lower limit+0.3)kPa~16.0 kPa)
	Upper Limit of Mean Blood Pressure	
	Upper Limit of Diastolic Blood Pressure	
Art	Lower Limit of Systolic Blood Pressure	0 mmHg~(Upper limit-2)mmHg (0 kPa~(Upper limit-0.3)kPa)
	Lower Limit of Mean Blood Pressure	
	Lower Limit of Diastolic Blood Pressure	
P1 P2	Lower Limit of Systolic Blood Pressure	-50 mmHg~(Upper limit-2)mmHg (-6.7 kPa~(Upper limit -0.3)kPa)
	Lower Limit of Mean Blood Pressure	
	Lower Limit of Diastolic Blood Pressure	
PA	Lower Limit of Systolic Blood Pressure	-6 mmHg~(Upper limit-2)mmHg (-0.8 kPa~(Upper limit-0.3)kPa)
	Lower Limit of Mean Blood Pressure	
	Lower Limit of Diastolic Blood Pressure	
LAP RAP	Upper Limit of Mean Blood Pressure	(Lower limit+2)mmHg~40 mmHg ((Lower limit+0.3)kPa~5.3 kPa)
ICP CVP	Lower Limit of Mean Blood Pressure	-10 mmHg~(Upper limit-2)mmHg (-1.3 kPa~ (Upper limit-0.3)kPa)

**B.6.8 CO<sub>2</sub> Monitoring (Optional)**

Measuring Mode	Sidestream type (support 50ml/min pumping rate), mainstream type
Measuring Method	Infrared radiation absorption technique

**Respironics Sidestream LoFlo Module**

Measuring Method	Infrared Spectrum Method
Measuring Mode	Sidestream
Preheating time	Max. length of waveform is 20s. Full accuracy requirements satisfied after 2min (environment temp.: 25°C)
Range	0%~19.7% (0 mmHg ~150 mmHg) (0 kPa~20 kPa)



## Product Specifications

Resolution	0.1 mmHg 0 mmHg~69 mmHg 0.25 mmHg 70 mmHg~150 mmHg
Stability	Short-term drift: $\leq 0.8$ mmHg (0.1 kPa) within 4h Long-term drift: accuracy maintained within 120h.
Unit selection	%, mmHg, kPa
Accuracy (Gas Temp. at 25°C)	0 mmHg~40 mmHg (0 kPa~5.3 kPa), $\pm 2$ mmHg (0.27 kPa) 41 mmHg~70 mmHg (5.5 kPa~9.3 kPa), $\pm 5\%$ of the reading 71 mmHg~100 mmHg (9.4 kPa~13.3 kPa), $\pm 8\%$ of the reading 101 mmHg~150 mmHg (13.4 kPa~20 kPa), $\pm 10\%$ of the reading (When the breathing rate is $> 80$ rpm, all ranges are $\pm 12\%$ of the reading)
Total System Response Time	$< 3$ s
Range of Breathing Rate	2 rpm~150 rpm
Accuracy of Breathing Rate	$\pm 1$ rpm
Asphyxia Alarm Delay	20 s, 25 s, 30 s, 35 s, 40 s, 45 s, 50 s, 55 s, 60 s
Sampling Flow Rate	$\geq 50$ ml/min(100Hz)
Automatic Pressure Compensation	no
<b>Alarm Limit Specifications</b>	<b>Range</b>
EtCO <sub>2</sub> Upper Limit	(Lower Limit +2) mmHg~99 mmHg
EtCO <sub>2</sub> Lower Limit	0 mmHg~(Upper Limit -2) mmHg
FiCO <sub>2</sub> Upper Limit	0 mmHg~99 mmHg
awRR Upper Limit	(Lower limit+2) rpm~100 rpm
awRR Lower Limit	0 rpm~ (Upper limit-2) rpm

## Respironics Mainstream CAPNOSTAT5 Module

Measuring Method	Infrared Spectrum Method
Measuring Mode	Mainstream
Preheating time	Max. length of waveform is 15s. Full accuracy requirements satisfied after 2min (environment temp.: 25°C)
Range	0%~19.7% (0 mmHg~150 mmHg) (0 kPa~20 kPa)
Resolution	0.1 mmHg 0 mmHg~69 mmHg 0.25 mmHg 70 mmHg~150 mmHg
Stability	Short-term drift: $\leq 0.8$ mmHg (0.1 kPa) within 4h Long-term drift: accuracy maintained within 120h.
Rise Time	$< 60$ ms
Unit selection	%, mmHg, kPa
Accuracy (Environment Temp. at 35°C)	0 mmHg~40 mmHg (0 kPa~5.3 kPa), $\pm 2$ mmHg (0.27 kPa) 41 mmHg~70 mmHg (5.5 kPa~9.3 kPa), $\pm 5\%$ of the reading 71 mmHg~100 mmHg (9.4 kPa~13.3 kPa), $\pm 8\%$ of the reading 101 mmHg~150 mmHg (13.4 kPa~20 kPa), $\pm 10\%$ of the reading

Range of Breathing Rate	0 rpm~150 rpm
Accuracy of Breathing Rate	±1 rpm
Asphyxia Alarm Delay	20 s, 25 s, 30 s, 35 s, 40 s, 45 s, 50 s, 55 s, 60 s
Sampling Flow Rate	100 Hz
Automatic Pressure Compensation	no
<b>Alarm Limit Specifications</b>	<b>Range</b>
EtCO <sub>2</sub> Upper Limit	(Lower Limit +2)mmHg~99 mmHg
EtCO <sub>2</sub> Lower Limit	0 mmHg~(Upper Limit -2)mmHg
FiCO <sub>2</sub> Upper Limit	0 mmHg~99 mmHg
awRR Upper Limit	(Lower limit+2) rpm~100 rpm
awRR Lower Limit	0 rpm~ (Upper limit-2) rpm

**Kingst KM7002-V33/KM7003-V40 Sidestream Module**

Measuring Method	Non-scattering Infrared Gas Analysis
Measuring Technology	Non-dispersive Infrared Gas Analysis (NIDR)
Range	0%~20% (0 mmHg~150 mmHg) (0 kPa~20 kPa)
Protection Level / Type	BF
Preheating time	2 min at 25 °C
Response Time	50 ml/min
Delay Time	50 ml/min
Fully-automatic Drift Calibration	Automated according to the time and temperature. Time 5 s~8 s
Airway Leakage	< 0.1% (within the flow range above)
Accuracy	When < 5.0%: ±0.3% (±2.0 mmHg) (0.27 kPa) When ≥5.0%: < 6% of the reading
Range of Breathing Rate	3 rpm~150 rpm
Accuracy of Breathing Rate	1% or ±1 rpm, whichever is higher.
Asphyxia Alarm Delay	30 s, 35 s, 40 s, 45 s, 50 s, 55 s, 60 s
Automatic Pressure Compensation	yes
<b>Alarm Limit Specifications</b>	<b>Range</b>
EtCO <sub>2</sub> Upper Limit	(Lower Limit +2)mmHg~99 mmHg
EtCO <sub>2</sub> Lower Limit	0 mmHg~(Upper Limit -2)mmHg
FiCO <sub>2</sub> Upper Limit	0 mmHg~99 mmHg
awRR Upper Limit	(Lower limit+2) rpm~100 rpm
awRR Lower Limit	0 rpm~ (Upper limit-2) rpm

**B.6.9 Recorder Specifications (Optional)**

Recorder	To record the patient information, the hospital information, waveform, parameters and others displayed in the screen
Method	Thermal array recorder
Printing Paper	Thermal paper
Print Resolution	8 dots/mm on Y-Axis
Delay Characteristics	$\leq 0.5$ mm
Amplitude-frequency Characteristics	Monitor Mode: 0.5 Hz $\sim$ 40 Hz; Diagnose Mode: 0.05 Hz $\sim$ 150 Hz.
Time Constant	$\geq 0.3$ s

## Mobile Trolley for Patient Monitor

### *Suitable for BIOCARE Patient Monitors*

(Model: V-Cart, Part Number: 90530000-01)

#### Features

- Easy, flexible height, rotation, tilting adjustment
- Elegant Integrated design, quick installation, fine workmanship
- Excellent durable, environmentally friendly material, with high resistance to corrosion
- 2x baskets for accessories / consumables placement
- Stable and easy-moving 5 universal wheels of 2 with braker
- Aluminum alloy tray with rail for flexible support of different models



#### Specifications

Size: 70cm x 70cm x H (92 to 115cm adjustable)

Weight: 8.6kg

Max. Load: 25kg

Rotation angle (horizontal): 360 degree

Tilting angle: +/- 30 degree

Wheel diameter: 70cm

Packing size / Weight: 71\*71\*26cm / 11.4kgs

Aluminium alloy for top supporter plates  
Stainless steel for pole  
Material: ABS for locker and wheels  
Nylon for bottom base supporter  
White coated steel for baskets



**Shenzhen Biocare Bio-Medical Equipment Co., Ltd.**

Add: #16-1, Jinhui Rd, Jinsha Community, Kengzi, Pingshan District,  
518122 Shenzhen China

Tel: +86-755-27960888

Fax: +86-755-2796 0643

Web: www.biocare.com.cn

Email: sales@biocare.com.cn

## DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 93/42/EEC CONCERNING MEDICAL DEVICES



*SHENZHEN BIOCARE BIO-MEDICAL EQUIPMENT CO., LTD.*  
*#16-1, JINHUI ROAD, JINSHA COMMUNITY, KENGZI SUB-DISTRICT, PINGSHAN NEW DISTRICT, 518122*  
*SHENZHEN, PEOPLE'S REPUBLIC OF CHINA*

MEDICAL DEVICE: *PATIENT MONITOR*  
*TYPE: PM-900*  
*GMDN CODE: 33586*

CLASSIFICATION - ANNEX IX: *CLASS IIb, RULE 10*  
CONFORMITY ASSESSMENT ROUTE: *ANNEX II EXCLUDING(4)*

**WE, THE MANUFACTURER, HEREWITH DECLARE THAT THE STATED MEDICAL DEVICES  
MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE  
93/42/EEC CONCERNING MEDICAL DEVICES;  
INCLUDING, AT 21 MARCH 2010, THE AMENDMENTS BY COUNCIL DIRECTIVE 2007/47/EEC.  
ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER.  
THE MANUFACTURER IS EXCLUSIVELY RESPONSIBLE FOR THE DOC.**

NOTIFIED BODY: TÜV SÜD PRODUCT SERVICE GMBH  
Ridlerstraße 65 · 80339 Munich · Germany

IDENTIFICATION NUMBER



(EC) CERTIFICATE(S): *G1 065758 0004 REV.01*

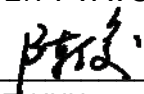


EUROPEAN REPRESENTATIVE: *SHANGHAI INTERNATIONAL HOLDING CORP. GMBH*  
*(EUROPE)*  
*Eiffestraße 80, 20537 Hamburg, GERMANY*

START OF CE-MARKING: *2012-08-02*

PLACE, DATE OF DECLARATION: *SHENZHEN P.R.C., 2019-09-19*

SIGNATURE:

  
NAME: *CHEN JUN*  
POSITION: *GENERAL MANAGER*





Benannt durch/Designated by  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln und  
Medizinprodukten  
www.zlg.de  
ZLG-BS-244.10.08



Product Service

# EC Certificate

Full Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)  
(Devices in Class IIa, IIb or III)

**No. G1 065758 0004 Rev. 01**

## Manufacturer:

**Shenzhen Biocare Bio-Medical  
Equipment Co., Ltd.**

#16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District,  
Pingshan New District  
518122 Shenzhen  
PEOPLE'S REPUBLIC OF CHINA

## Facility(ies):

Shenzhen Biocare Bio-Medical Equipment Co., Ltd.  
#16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District,  
Pingshan New District, 518122 Shenzhen, PEOPLE'S REPUBLIC  
OF CHINA

**Product Category(ies): Digital Electrocardiograph, Patient Monitor,  
B-Ultrasonic Diagnostic Equipment,  
Doppler Fetal Heart Rate Detector, Infusion  
Pump, Syringe Pump, Fingertip Pulse  
Oximeter, Handheld Pulse Oximeter,  
Fetal/Maternal Monitor, Fetal Monitor, Color  
Doppler Ultrasound System, Central  
Monitoring System, Ambulatory  
Electrocardiographs, Ambulatory blood  
pressure recorders, and associated  
software.**

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

**Report No.:** BJ1989607  
**Valid from:** 2019-09-11  
**Valid until:** 2024-05-26

**Date,** 2019-09-11

Stefan Preiß  
Head of Certification/Notified Body

ZERTIFIKAT • CERTIFICATE • 認證證書 • CERTIFICADO • CERTIFICAT



# Certificate

No. Q5 065758 0005 Rev. 01

**Holder of Certificate:** **Shenzhen Biocare Bio-Medical Equipment Co., Ltd.**

#16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District  
Pingshan New District  
518122 Shenzhen  
PEOPLE'S REPUBLIC OF CHINA

**Facility(ies):** Shenzhen Biocare Bio-Medical Equipment Co., Ltd.  
#16-1 Jinhui Road, Jinsha Community, Kengzi Sub-District,  
Pingshan New District, 518122 Shenzhen, PEOPLE'S REPUBLIC  
OF CHINA

**Certification Mark:**



**Scope of Certificate:** Design and Development, Production and Distribution of Digital Electrocardiograph, B-Ultrasonic Diagnostic Equipment, Patient Monitor, Fetal Monitor, Doppler Fetal Heart Rate Detector, Infusion Pump, Syringe Pump, Fingertip Pulse Oximeter, Handheld Pulse Oximeter, Ambulatory electrocardiographs, Ambulatory blood pressure recorder and associated software, Fetal/Maternal Monitor, Color Doppler Ultrasound System, Central Monitoring System.

**Applied Standard(s):** EN ISO 13485:2016  
Medical devices - Quality management systems -  
Requirements for regulatory purposes  
(ISO 13485:2016)  
DIN EN ISO 13485:2016

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). See also notes overleaf.

**Report No.:** BJ20089601

**Valid from:** 2020-04-01

**Valid until:** 2023-03-31

**Date,** 2020-03-17

Christoph Dicks

Head of Certification/Notified Body



REGISTRUL DE STAT AL DISPOZITIVELOR MEDICALE

Tip	Denumir	Введите текст для поиска...										
I.3. Certificat CE	Certificat CE	Nr	Denumire	Den.comerc.	Model	Nr. catalog	Tara	Producatorul	Reprezentant	Ordin	Data	Cod vamal
I.2. Declarat de conform CE	Declarat de conform CE				PM-900			biocare				
		DM000041944	MONITOR DE PACIENT		PM-900		China	SHENZHEN BIO CARE BIO-MEDICAL EQUIPMENT CO., LTD.	RAILSMED S.R.L.	A07.PS-01.Rg04-8	10-01-2018	
✓ 🔍 Содержит('Denumire','.') И Содержит('Producatorul','biocare') И Содержит('Model'),...												
Очистить												

Очистить