Instructions for Blood Pressure cuff

Introduction

Product name: Blood Pressure Cuff Product model and configuration: KM series (reusable cuff):

Model	Applicable	Application	Limb
INIOUEI	to	site	circumference
KM-221	Infant	Arm	6cm - 11cm
KM-222	Infant		8cm - 13cm
KM-232	Child		10cm - 19cm
KM-233	Child		18cm - 26cm
KM-241	Adult		21cm - 35cm
KM-242	Adult		27cm - 42cm
KM-243	Adult		40cm - 48cm
KM-244	Adult	Thigh	46cm - 66cm
KM-333	Child	Arm	18cm - 26cm
KM-341	Adult		21cm - 35cm
KM-342	Adult		27cm - 42cm
KM-343	Adult		40cm - 48cm

Service life: 3 years

KN series (reusable cuff):

Model	Applicable to	Application site	Limb circumference
KN-221	Infant		6cm - 11cm
KN-231	Child		10cm - 19cm
KN-233	Child	Arm	18cm - 26cm
KN-241	Adult		25cm - 35cm
KN-243	Adult		33cm - 47cm
KN-244	Adult	Thigh	46cm - 66cm

Intended use: The blood pressure cuff is indicated for use in manual measurement and automatic non-invasive blood pressure monitoring. It's applicable to be used with a compatible monitor.

Instructions for use

- ▲ Appropriate cuff should be selected according to the age and arm/thigh circumference of the subject. Its width should be 2/3 of the length of the upper arm/thigh. The inflatable part should be long enough to permit wrapping approximately 80% of the limb. When cuff sizes overlap for a specified circumference, choose the larger size.
- ▲ Check the cuff before use, replace the cuff when aging, tearing or weak closure is apparent. Do not use a damaged cuff.
- \triangle Select the appropriate blood pressure measurement site. Inspect patient's limb prior to application.
- M When applying the cuff, unfold and wrap around the upper arm/thigh evenly to the appropriate tightness. The cuff should be tightened to a degree where insertion of one finger is allowed
- ▲ Locate the cuff in such a way that the artery mark " **I** " is at a location where the clearest pulsation of brachial artery is observed.
- \triangle Remember to empty any residual air in the cuff before the measurement is commenced.

Operating Environment

Ambient temperature range: -10°C- 40°C; Relative humidity: 10% - 85%;

Atmospheric pressure: 50kPa - 106.0kPa

The cuff should be stored and used within the specified temperature and humidity range, or it may cause damage to the cuff or inaccurate measurement results.

Cleaning and Disinfection

- 1. Prepare the enzymatic detergent or equivalent and distilled water, and 10% bleach solution in separate spray bottles.
- 2. Spray detergent liberally on cuff, tubing and hose. If dirt is dried on, allow the detergent to soak in to the cuff for one minute.
- 3. Wipe smooth surface with a soft cloth. Use a soft-bristle brush on visibly stained areas and irregular surfaces. Note: Take particular care when cleaning the bulb and control valve knob on a complete inflation system. Do not allow fluid to enter back valve or saturate control valve knob. Remove visible contaminants from the periphery and the underside of the control valve knob.
- 4. Rinse with copious amounts of distilled water.
- 5. To disinfect, spray 10% bleach solution on cuff until saturated and allow to soak for five minutes.
- 6. Wipe away excess solution and rinse again with distilled water. Allow cuff to air dry.

Warnings and Precautions

- A Blood pressure measurement is prohibited to those who have severe hemorrhagic tendencies or with sickle cell disease, as partial bleeding may be caused.
- ▲ Continuous measurement may result in purpura, neuralgia and lack of blood.
- \triangle Do not place the cuff on limbs with transfusion tubes, intubations or skin lesions on the area, as damage may be caused to the limbs.
- \triangle Avoid compressing or restricting the connection tubing.
- Minimize limb movement and cuff motion during measurement.
- \triangle Check the site and limb frequently, especially when monitoring at frequent intervals and/or over extended periods of time.
- \triangle Remove the cuff from the patient when the measurement has been taken.
- \triangle Use cuff only under direct supervision by trained healthcare professional when attached to automated monitors without alarms.
- A Before use, empty the cuff until there is no residual air inside. Do not allow the cuff to twist or bend.
- \triangle Do not twist the cuff hose or put heavy things on it.
- A Please hold the connector of the hose while connecting and disconnecting it to the device.
- ⚠ If arrhythmia or auricular fibrillation occurs, take the measurement again.
- Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

CE	Medical Device compliant with Regulation (EU) 2017/745	×	Keep away from sunlight
	Manufacturer	EC REP	Authorised representative in the European Community
MD	Medical Device	\triangle	Caution
8	Refer to the accompanying documents	YYYY-MM-DD	Country of manufacture (CN stands for Made in China) and Date of manufacture
-10°C	Temperature range	10%	Relative humidity
50kPa	Atmospheric pressure	Ť	Keep in a cool, dry place

Manufacturer: Shenzhen Creative Industry Co., Ltd.

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