TRANSMITTER

ZM-920PA/930PA

0614-007205

Model: ZM-920PA/930PA

Manual code no.: 0614-007205

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GENERAL HANDLING PRECAUTIONS

This device is intended for use only by qualified medical personnel. Use only Nihon Kohden approved products with this device. Use of non-approved products or in a non-approved manner may affect the performance specifications of the device. This includes, but is not limited to, batteries, recording paper, pens, extension cables, electrode leads, input boxes and AC power.

Please read these precautions thoroughly before attempting to operate the instrument.

1. To safely and effectively use the instrument, its operation must be fully understood.

2. When installing or storing the instrument, take the following precautions:

- (1) Avoid moisture or contact with water, extreme atmospheric pressure, excessive humidity and temperatures, poorly ventilated areas, and dust, saline or sulphuric air.
- (2) Place the instrument on an even, level floor. Avoid vibration and mechanical shock, even during transport.
- (3) Avoid placing in an area where chemicals are stored or where there is danger of gas leakage.
- (4) The power line source to be applied to the instrument must correspond in frequency and voltage to product specifications, and have sufficient current capacity.
- (5) Choose a room where a proper grounding facility is available.

3. Before Operation

- (1) Check that the instrument is in perfect operating order.
- (2) Check that the instrument is grounded properly.
- (3) Check that all cords are connected properly.
- (4) Pay extra attention when the instrument is in combination with other instruments to avoid misdiagnosis or other problems.

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- (5) All circuitry used for direct patient connection must be doubly checked.
- (6) Check that battery level is acceptable and battery condition is good when using battery-operated models.

4. During Operation

- (1) Both the instrument and the patient must receive continual, careful attention.
- (2) Turn power off or remove electrodes and/or transducers when necessary to assure the patient's safety.
- (3) Avoid direct contact between the instrument housing and the patient.

5. To Shutdown After Use

- (1) Turn power off with all controls returned to their original positions.
- (2) Remove the cords gently; do not use force to remove them.
- (3) Clean the instrument together with all accessories for their next use.
- 6. The instrument must receive expert, professional attention for maintenance and repairs. When the instrument is not functioning properly, it should be clearly marked to avoid operation while it is out of order.
- 7. The instrument must not be altered or modified in any way.

8. Maintenance and Inspection:

- (1) The instrument and parts must undergo regular maintenance inspection at least every 6 months.
- (2) If stored for extended periods without being used, make sure prior to operation that the instrument is in perfect operating condition.
- (3) Technical information such as parts list, descriptions, calibration instructions or other information is available for qualified user technical personnel upon request from your Nihon Kohden distributor.

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- 9. When the instrument is used with an electrosurgical instrument, pay careful attention to the application and/or location of electrodes and/or transducers to avoid possible burn to the patient.
- 10. When the instrument is used with a defibrillator, make sure that the instrument is protected against defibrillator discharge. If not, remove patient cables and/or transducers from the instrument to avoid possible damage.

WARRANTY POLICY

Nihon Kohden Corporation (NKC) shall warrant its products against all defects in materials and workmanship for one year from the date of delivery. However, consumable materials such as recording paper, ink, stylus and battery are excluded from the warranty.

NKC or its authorized agents will repair or replace any products which prove to be defective during the warranty period, provided these products are used as prescribed by the operating instructions given in the operator's and service manuals.

No other party is authorized to make any warranty or assume liability for NKC's products. NKC will not recognize any other warranty, either implied or in writing. In addition, service, technical modification or any other product change performed by someone other than NKC or its authorized agents without prior consent of NKC may be cause for voiding this warranty.

Defective products or parts must be returned to NKC or its authorized agents, along with an explanation of the failure. Shipping costs must be prepaid.

This warranty does not apply to products that have been modified, disassembled, reinstalled or repaired without Nihon Kohden approval or which have been subjected to neglect or accident, damage due to accident,

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fire, lightning, vandalism, water or other casualty, improper installation or application, or on which the original identification marks have been removed.

In the USA and Canada other warranty policies may apply.

CAUTION

United States law restricts this device to sale by or on the order of a physician.

Equipment Authorization Requirement

Operation of this equipment requires the prior coordination with a frequency coordinator designated by the FCC for the Wireless Medical Telemetry Service.

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EMC RELATED CAUTION

This equipment and/or system complies with the International Standard IEC60601-1-2 for electromagnetic compatibility for medical electrical equipment and/or system. However, an electromagnetic environment that exceeds the limits or levels stipulated in the IEC60601-1-2, can cause harmful interference to the equipment and/or system or cause the equipment and/or system to fail to perform its intended function or degrade its intended performance. Therefore, during the operation of the equipment and/or system, if there is any undesired deviation from its intended operational performance, you must avoid, identify and resolve the adverse electromagnetic effect before continuing to use the equipment and/or system.

The following describes some common interference sources and remedial actions:

- 1. Strong electromagnetic interference from a nearby emitter source such as an authorized radio station or cellular phone: Install the equipment and/or system at another location if it is interfered with by an emitter source such as an authorized radio station. Keep the emitter source such as cellular phone away from the equipment and/or system.
- 2. Effect of direct or indirect electrostatic discharge: Make sure all users and patients in contact with the equipment and/or system are free from direct or indirect electrostatic energy before using it. A humid room can help lessen this problem.
- 3. Electromagnetic interference with any radio wave receiver such as radio or television:
 If the equipment and/or system interferes with any radio wave receiver, locate the equipment and/or system as far as possible from the radio wave receiver.

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If the above suggested remedial actions do not solve the problem, consult your Nihon Kohden Corporation subsidiary or distributor for additional suggestions.

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Conventions Used in this Manual and Instrument

Warnings, Cautions and Notes

Warnings, cautions and notes are used in this manual to alert or signal the reader to specific information.

WARNING

A warning alerts the user to the possible injury or death associated with the use or misuse of the instrument.

CAUTION

A caution alerts the user to possible injury or problems with the instrument associated with its use or misuse such as instrument malfunction, instrument failure, damage to the instrument, or damage to other property.

NOTE

A note provides specific information, in the form of recommendations, prerequirements, alternative methods or supplemental information.

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Explanations of the Symbols in this Manual and Instrument

The following symbols found in this manual/instrument bear the respective descriptions as given.

Symbol	Description	Symbol	Description
	Power On	\wedge	Attention, consult operator's manual
0	Power Off	+	Nurse call
I ¥ I	Defibrillation proof type BF applied part		Replace battery
l ● ŀ	Defibrillation proof type CF applied part	P	Check electrode
	Direct current		

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Introduction

The ZM-920PA/930PA transmitter transmits ECG and other data from a patient to a Nihon Kohden monitor for continuous monitoring. Available parameters and functions vary between the models. Read the operator's manual for the monitor before operation.

Model	Parameters	Functions	
		The following information is	
714 02004	• ECG	indicated by LED.	
ZM-920PA	Impedance respiration	 Check ECG electrodes 	
		 Replace batteries 	
		The following information is	
		indicated on LCD.	
	• ECG	• SpO2 value	
7M 020DA	• ECO	 Pulse wave amplitude 	
ZM-950FA		Replace batteries	
	• SpO2	The following information is	
		indicated by LED.	
		Check ECG electrodes	

The transmitter channel can be changed by the QI-901PK Channel Writer. To change the channel number, refer to the channel writer manual.

WARNING

The following actions must be taken to properly receive the transmitter signal of the correct patient on the receiving instrument, otherwise, there may be signal loss or signals may mix causing a serious accident, such as monitoring a different patient.

• Assign a channel administrator in the hospital and only he or she should manage channel assignments.

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- The channel administrator must manage the channels in the facility so that there is no signal interference.
- When the transmitter channel is changed, the channel administrator must check that the channel on the receiving monitor is also changed and the signal is properly received.

• The channel administrator must replace the channel number label on the transmitter with the new one after changing the channel.

CAUTION

- Do not use the same channel for different patients, otherwise, two patients' data will be lost due to mutual modulation interference, or the wrong patient's data may appear on the receiving monitor screen.
- Do not use transmitters of adjacent channels in a hospital, otherwise, radio waves from one transmitter affect the receiver of the adjacent channel's in the transmitter and there may be interference.

NOTE

- Use Nihon Kohden parts and accessories to assure maximum performance from your instrument.
- It is recommended to use a diversity antenna system on the receiving monitor for stable signal reception, otherwise, spike noise from transient fading of electric field strength (for example, people moving) may interfere with the transmitter signal and may be mistaken as an arrhythmia on the receiving monitor.

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Panel Description

Top Panel



WARNING

- Before performing defibrillation, check that the electrode leads and SpO₂ probe attached to the patient are properly connected to the transmitter. Touching the metal parts of disconnected leads and probes may cause serious electrical shock or injury by discharged energy.
- When performing defibrillation, all persons must keep clear of the bed and must not touch the patient, any equipment connected to the patient or the metal parts of leads and probes connected to the patient. Failure to follow this warning may result in serious electrical burn, shock or other injury.

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Front Panel

ZM-920PA



WARNING

Close the battery case cover during operation. If the transmitter is used with the battery case cover open, the patient may get an electrical shock when defibrillation is performed, and electrostatic discharge by the patient may intermittently interfere with the waveform or data.

CAUTION

Only use your finger to press the CALL key. Do not press the key with a sharp object, otherwise the key may be damaged.

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ZM-930PA



When the detected pulse is too small to measure, "---" is displayed.

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Important Safety Information

General

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WARNING

- Never use this transmitter in the presence of any flammable anesthetic gas or high concentration oxygen atmosphere. Failure to follow this warning may cause explosion or fire.
- Never use this transmitter in a high-pressure oxygen medical care tank. Failure to follow this warning may cause explosion or fire.
- Never take this transmitter into an MRI test room.
- Before performing defibrillation, check that the electrode leads and SpO₂ probe attached to the patient are properly connected to the transmitter. Touching the metal parts of disconnected leads and probes may cause serious electrical shock or injury by discharged energy.
- When performing defibrillation, all persons must keep clear of the bed and must not touch the patient, the equipment connected to the patient, nor the metal parts of leads and probes connected to the patient. Failure to follow this warning may result in serious electrical burn, shock or other injury.
- When performing defibrillation, discharge as far as possible from electrodes on the patient. If there is a possibility that the defibrillator paddle could touch electrodes, remove electrodes from the patient. If the defibrillator directly contacts the electrodes, the discharged energy may cause serious electrical burn to the patient.
- When using this transmitter with an electrosurgery unit, its return plate and the electrodes for monitoring must be firmly attached to the patient. If the return plate is not attached correctly, it may burn the patient's skin where the electrodes are attached. Refer to the instruction manual for the ESU.
- Close the battery case cover during operation. If the transmitter is used with the battery case cover open, the patient may get an electrical shock when defibrillation is performed, and electrostatic

discharge by the patient may intermittently interfere with the waveform or data.

CAUTION

- Use Nihon Kohden specified electrode leads and SpO₂ probes to assure maximum performance from your instrument.
- Do not reuse disposable products.
- Do not shake or swing the transmitter holding the leads/cables connected to the transmitter. The transmitter may come off and cause injury to a person or damage surrounding instruments.
- Attach a strap to the transmitter to prevent the transmitter from falling.
- Turn off the power of cellular telephones, small wireless devices and other devices which produce strong electromagnetic interference around a patient. Radio waves from devices such as cellular telephones or small wireless devices may be mistaken as pulse waves and incorrect data may be displayed.
- Do not use the same channel for different patients. This could produce a mutual modulation interference resulting in loss of data from both patients or the incorrect patient's data can appear on the receiving monitor screen.
- Do not use transmitters of adjacent channels in a hospital, otherwise, radio waves from one transmitter may affect the receiver of the adjacent channel's transmitter and can cause interference.

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Battery

WARNING

- Do not dispose of the battery in fire, or it may explode.
- Do not disassemble the battery. The contents of the battery are harmful and flammable.
- Never short-circuit the + and terminals. This can produce overheating and with its flammable capabilities can produce a fire.
- Make sure that the patient does not touch the batteries.

CAUTION

Battery replacement must be performed by medical staff. When replacing batteries in the transmitter currently used for a patient, disconnect electrode leads from the transmitter before replacing batteries. Do not touch the patient during replacement.

Transmitter Channel Management

WARNING

The following actions must be taken to properly receive the transmitter signal of the correct patient on the receiving instrument, otherwise, there may be signal loss or signals may mix causing a serious accident, such as monitoring a different patient.

- Assign a channel administrator in the hospital and only he or she should manage channel assignments.
- The channel administrator must manage the channels in the facility so that there is no signal interference.
- When the transmitter channel is changed, the channel administrator must check that the channel on the receiving monitor is also changed and the signal is properly received.
- The channel administrator must replace the channel number label on the transmitter with the new one after changing the channel.



For Patients Using Implantable Pacemaker

WARNING

Interaction Between Minute Ventilation Rate-Adaptive Pacemakers and Cardiac Monitoring and Diagnostic Equipment. The bioelectric impedance measurement sensor of a minute ventilation rate-adaptive implantable pacemaker, may be affected by the transmitter which is connected to the same patient. If this occurs, the pacemaker may pace at its maximum rate and the transmitter may give incorrect data to the monitor. If this occurs, disconnect the electrode leads from the patient or change the setting on the pacemaker by referring to the pacemaker's manual. For more details, contact your pacemaker distributor or Nihon Kohden distributor.

Output Signal

CAUTION

Do not use the output signal from the receiving monitor as the synchronization signal for other equipment such as IABP, MRI, echocardiography or defibrillator, there may be time delay between the monitor and the other equipment and spike noise may interfere with the output signal and be mistaken as a trigger.

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ECG Monitoring

CAUTION

- Use Nihon Kohden specified consumables. With electrodes other than specified ones, the CHECK ELECTRODE message appears and monitoring may stop.
- When the "ELECTRODE OFF" or "CHECK ELECTRODE" message is displayed on the receiving monitor, check electrodes and electrode leads. While "ELECTRODE OFF" or "CHECK ELECTRODE" message is being displayed, there is no ECG monitoring or alarms.

SpO₂ Monitoring

WARNING

- Measurement may be incorrect in the following cases.
 - When the oxyhemoglobin or methemoglobin (HbCO, Met Hb) increases abnormally.
 - When dye is injected in the blood.
 - · When using an electrosurgical unit.
 - During CPR.
 - · When measuring at a site where there are venous pulses.
 - · When there is body movement.
 - · When the pulse wave is small.
- Check the circulation condition by observing the skin color of the measuring site and pulse waveform. Change the measuring site every 8 hours for disposable probes and every 4 hours for reusable probes. The skin temperature may increase at the attached site by 2 or 3°C (4 or 5°F) and cause a burn or pressure necrosis. When using the probe on the following patients, take extreme care and change the measurement site more frequently according to symptoms and degree.
 - · A patient with a fever
 - · A patient with peripheral circulation insufficiency
 - · Neonate or low birth weight infant with delicate skin
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- To avoid poor circulation, do not wrap the tape too tight. Check the blood circulation condition by observing the skin color and congestion at the skin peripheral to the probe attachment site. Even for short-term monitoring, there may be burn or pressure necrosis from poor blood circulation, especially on neonates or low birth weight infants whose skin is delicate. Accurate measurement cannot be performed on a site with poor peripheral circulation.
- When not monitoring SpO₂, disconnect the SpO₂ probe cord from the transmitter. Otherwise, noise may interfere from the probe sensor and cause incorrect data to be displayed on the transmitter and receiving monitor.

CAUTION

- Do not pull or bend the probe cable or put caster feet on the probe cable. Do not immerse the probe cable in detergents or water. Failure to follow these cautions may cause cable discontinuity, short circuit, skin burn on the patient or incorrect measurement data. Replace any broken probe with a new one.
- When the attachment site is wet with blood or when the patient has nail polish on, remove the dirt and nail polish before attaching the probe. The transmitted light may decrease due to the blood or nail polish and the measurement data may be incorrect.
- If the skin gets irritated or redness appears on the skin by the probe, change the attachment site.
- When the probe is attached to an appropriate site with sufficient circulation and an error message confirming the probe attachment repeatedly appears, the probe may be deteriorated. Replace it with a new one.
- Do not use the probe over its stated lifetime. Otherwise the SpO₂ measurement accuracy cannot be guaranteed.
- Use the disposable probe only for one patient. Never reuse the disposable probe for another patient because it causes cross infection.
- Do not use damaged or disassembled probe. Measured data may be incorrect.

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- When measuring under strong light (surgical light, bilirubin light, sunlight, etc.), cover the probe with a blanket or cloth. Otherwise, noise may interfere.
- When any of the following occurs, the probe may be broken. Replace it with a new one and check the probe.
 - The transmitter generates "pip" sound every 0.25 seconds.
 - SpO₂ data is 85% and blinking.

Disposable SpO₂ Probes

CAUTION

- Replace the probe with a new one as specified in the probe manual. If the probe is deteriorated, correct SpO₂ monitoring cannot be performed.
- When using a disposable probe, be careful when removing the adhesive tape from neonatal skin.
- When removing a disposable probe taped to the skin, do not pull from the cable of the probe because this can damage the probe's cable connection.

TL-260T Multi-site Y Probe

CAUTION

- Before use, be sure to attach the probe to the sponge attachment tape S or L. Do not use the probe without the sponge attachment tape attached. It causes incorrect measurement and may damage the attachment site on the skin.
- When fixing the probe with the sponge attachment tape, confirm that the adhesive part of the tape is not on the skin. The adhesive may cause oversensitive symptoms on the skin such as redness or itch. If the adhesive touches the skin, remove it carefully and slowly because neonatal skin is very delicate.
- Do not use a dirty sponge attachment tape. The measurement value may be incorrect.



• Do not pull from the cable when removing the probe from the sponge attachment tape because this can damaged the cable.

Maintenance

WARNING

If detergents or dirty liquid spills into the transmitter, clean it and dry it completely before use. If the wet transmitter is used, the patient or anyone in contact with the transmitter may receive an electric shock.

CAUTION

Do not disassemble the transmitter when performing maintenance and inspection. If there is a problem with the transmitter after maintenance and inspection, contact your Nihon Kohden distributor.

Reusable Probes

CAUTION

- Do not soak the probe in cleaning solution. It is not waterproof.
- Do not use creosol soap, glutaraldehyde, sodium hypochlorite, or benzalkonium chloride, as these substances may damage the probe.

Disposable Probes

CAUTION

- Do not soak the probe in cleaning solution. It is not waterproof.
- Do not use any disinfecting alcohol. It can damage the probe.

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Preparation

Installing (Replacing) Batteries

Use two AA type alkaline dry cell batteries, manganese dry cell batteries, NiCd rechargeable batteries or NiMH batteries.

With new alkaline batteries, the transmitter can continuously measure ECG, respiration and SpO_2 for approximately 3 days or ECG and respiration for approximately 4 days. Operation time depends on the thickness of SpO_2 probe attachment site.

NOTE

The capacity of manganese, NiCd and NiMH batteries is less than that of alkaline batteries, therefore the lifetime of the battery is shorter.

Туре	Lifetime	
Manganese	About 1/2 of alkaline batteries	
NiCd	About 1/3 of alkaline batteries (when fully charged)	
NiMH	About 1/2 of alkaline batteries (when fully charged)	

CAUTION

Battery replacement must be performed by medical staff. When replacing batteries in the transmitter currently used for a patient, disconnect electrode leads from the transmitter before replacing the batteries. Do not touch the patient during replacement.

If electrode leads are attached to the patient and the person replacing the batteries touches the patient, the patient leakage current over the amount allowed may occur.

CAUTION

- Replace both batteries at the same time.
- Do not use different types of batteries together.



NOTE Insert the batteries with the correct polarity (+ and –).



Procedure

- 1. Open the battery case cover.
- Insert two dry cell batteries (LR6) into the battery case observing the correct polarity.
- 3. Close the cover and press it gently until it clicks.

WARNING and CAUTION for Battery Handling

WARNING

- Do not dispose of the battery in fire, or it may explode.
- Do not disassemble the battery. The contents of the battery are harmful and flammable.
- If the contents of the battery contacts the skin or clothes, wash immediately and thoroughly with running water.
- Never short-circuit the + and terminals. This can produce overheating and with its flammable capabilities can produce a fire.
- Make sure that the patient does not touch the batteries.

CAUTION

When the transmitter is not in use, remove batteries or turn the power OFF. With the power ON, battery power is consumed even if measurement is not performed. The batteries may become unusable from overdischarge, and leakage from the battery may damage the transmitter.

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NOTE

- When using rechargeable NiCd batteries or NiMH batteries, shallow charging/discharging shortens battery capacity. For details, refer to the battery operator's manual.
- Remove the batteries from the transmitter before disposing of the transmitter.

Situations Requiring Battery Replacement

Replace the batteries when any of the following occurs.

- The "a" LED lights (ZM-920PA) or the "a" mark is displayed on the LCD (ZM-930PA) on the transmitter.
- The transmitter generates a constant alarm (continuous "peep" sound).
- The monitor displays the battery replacement message on the screen.
- When the power of the LCD transmitter is turned on, no message or icon is displayed. (Only the ZM-930PA).

Attaching a Strap to the Transmitter



To open the clip, firmly pull out the tab in direction of the arrow.

NOTE

- Attach a strap to the transmitter to prevent the transmitter from falling.
- Do not attach the clip to hard objects such as thick cloths or zippers, or the clip may break.

Attach a strap to the transmitter and fasten the clip to the patients' clothes or bed sheets.

If the transmitter falls off, the battery cover may be opened. If the patient touches the terminals of the batteries, patient leakage current over the allowable amount can occur.

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Turning On/Off the Transmitter

Check Items Before Turning On the Power

To use the instrument in a safe and optimum condition, before turning on the transmitter power switch, check the following.

Appearance

- There is no damage or dirt on the outside of the transmitter. (Power switch, LED, LCD, CALL key, junction, battery case cover, battery case, etc.)
- The transmitter is completely dry.
- The electrode lead is not broken.
- There is no damage or dirt on the SpO₂ probe or on the disposable electrodes. **Battery**
- The battery polarity is correct.
- The battery case spring is firmly fixed and the battery is not loose.
- The battery case cover is firmly closed.

Channel Setting

- The transmitter channel corresponds to those of the receiving monitor.
- The same channel is not being used on a different transmitter in the surrounding area.

Turning On/Off the Power



To turn on the power, turn the power switch to the right. After a "peep" sound for about one second, the power is turned on. (There is no "peep" sound when the "a" LED light or the "a" are blinking on the LCD).

To turn off the power, turn the power switch to the left.

Check Items After Turning On the Power

After turning on the power, check the following items.

Power On

- The power switch is not damaged.
- The transmitter generates a "peep" sound for about one second.

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- All LEDs light and values are displayed on the LCD for about one second.
- The transmitter does not generate a continuous "pip" sound.
- The transmitter does not liberate excessive heat.
- The ", LED does not light or the ", mark is not displayed on the LCD.
- The transmitter does not interfere with the operation of medical instruments used near it.

Basic Operation

- The "signal loss" message is not displayed on the monitor when the transmitter is inside the receiving range of the monitor.
- A "peep" sounds at the transmitter and "CALL" message appears at the receiving monitor when the CALL key is pressed and the transmitter is inside the receiving range of the monitor.
- The battery replacement message is not displayed on the monitor.

Check Items After the Power Off

- ECG electrode leads and SpO₂ probe are cleaned and disinfected.
- When the transmitter gets wet, liquid is wiped off and the transmitter is thoroughly dried.
- There are enough consumables, such as disposable electrodes.
- The power is turned off.
- The batteries are removed from the transmitter when it will not be used for a long time.
- Dead batteries are disposed of properly.

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ECG Monitoring

When 6 leads are used on this transmitter, up to 8 lead (I, II, III, aVR, aVL, aVF, Va and Vb) of ECG waveforms can be displayed on the receiving monitor. The heart rate is also measured. When 3 leads are used, one channel ECG waveform of lead II can be displayed on the receiving monitor. Refer to the operator's manual of the monitor for details.

WARNING

Interaction Between Minute Ventilation Rate-Adaptive Pacemakers and Cardiac Monitoring and Diagnostic Equipment* The bioelectric impedance measurement sensor of a minute ventilation rate-adaptive implantable pacemaker may be affected by the transmitter which is connected to the same patient. If this occurs, the pacemaker may pace at its maximum rate and the transmitter may give incorrect data to the monitor. If this occurs, disconnect the electrode leads from the patient or change the setting on the pacemaker by referring to the pacemaker's manual. For more details, contact your pacemaker distributor or Nihon Kohden distributor.

* Minute ventilation is sensed in rate-adaptive pacemakers by a technology known as bioelectric impedance measurement (BIM). Many medical devices in addition to pacemakers use this technology. When one of these devices is used on a patient with an active, minute ventilation rate-adaptive pacemaker, the pacemaker may erroneously interpret the mixture of BIM signals created in the patient, resulting in an elevated pacing rate.

For more information, see the FDA web site. http://www.fda.gov/cdrh/safety.html

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WARNING

When using the transmitter with an ESU, the ESU return plate and the electrodes for monitoring must be firmly attached to the patient. If the return plate is not attached correctly, it may burn the patient's skin where the electrodes are attached. Refer to the instruction manual for the ESU.

NOTE

- This transmitter is not protected against noise generated from an electrosurgery unit.
- If an electric blanket is used and incorrect heart rate is displayed on the monitor, turn off the pacing pulse detection on the monitor.

ECG Measurement Procedure

- 1. Select the type of electrode lead and disposable electrode according to the purpose.
- 2. Connect the electrode lead to the ECG/RESP socket.
- 3. Connect disposable electrodes to the electrode lead and attach electrodes to the patient.

After steps 1 to 3 are finished, ECG monitoring automatically starts.

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Selecting Electrode Lead and Disposable Electrode

CAUTION

Use Nihon Kohden specified consumables. With electrodes other than specified ones, the CHECK ELECTRODE message appears and monitoring may stop.

Option



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Connecting the Electrode Lead to the Transmitter

Connect the electrode lead to the ECG/RESP socket on the transmitter.

CAUTION

- Do not shake or swing the transmitter holding the leads/cables connected to the transmitter. The transmitter may come off and cause injury to a person or damage surrounding instruments.
- Hold the connector of the electrode lead when connecting/ disconnecting the electrode lead. If you disconnect the electrode lead holding the lead, it damages the electrode lead.



Selecting the Electrode Position

Follow the physician's instructions for electrode placement when available. For ECG monitoring, electrodes are attached only on the chest to allow patient movement and obtain continuous stable ECG. Following leads are examples. When also monitoring respiration, refer to the "Electrode Position for Respiration Monitoring" section.

NOTE

The optimum electrode positions for ECG measurement of a patient are not always optimum for respiration measurement of the patient. Select positions suitable for both ECG and respiration measurements, or positions which have priority for one measurement.

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Six Electrodes Electrode Position

The 6-electrode method with lead II and lead V5 is effective for monitoring myocardial ischemia. You can improve monitoring accuracy considerably by adding lead V4 to this combination. Va and Vb can be at any position of the standard 12 leads V1 to V6, but V4 and V5 are most appropriate for myocardial ischemic monitoring.



Symbol	Lead Color (Clip Color)	Electrode Position	
RA	White (White)	Right infraclavicular fossa	
LA	Black (Black)	Left infraclavicular fossa	
LL	Red (Red)	Lowest rib on the left anterior axillary line	
N (RL)	Green (Green)	Right anterior axillary line at the same level as LL	
Va	Va Brown (Brown) (BR-906PA) Brown-blue (BR-916PA) Fifth intercostal space midclavicular line. (V standard 12 leads)		
VbBrown (Brown) (BR-906PA) Brown-orange (BR-916PA)		Left anterior axillary line at the same level as Va. (V5 position of standard 12 leads)	

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Lead Position Standard limb leads Lead I







Monopolar limb leads aVR lead







Monopolar chest leads V1 to V6 leads



When Using 4 to 6 DIN Type Leads to Monitor 6 Lead ECG



When the BR-906PA/916PA electrode leads are not used, the transmitter is fixed to 3 lead ECG monitoring. To monitor 6 lead ECG using 4 to 6 DIN type leads, the transmitter must be fixed to 6 lead monitoring. To fix transmitter to the 6 lead ECG monitoring, turn off the transmitter power, press and hold the CALL key and turn on the transmitter power.

When the transmitter power is turned off and on again, the monitoring mode returns to the original mode.



Three Electrodes

By using the optional BR-913PA/903PA electrode lead, 3 lead ECG monitoring is available.

Electrode Position

• Lead MII, which is similar to standard lead II, used when ECG measurement has priority



	Electrode Position	Symbol	Lead Color
	Left infraclavicular fossa	LA(N)	Black
	Right infraclavicular fossa	RA (-)	White
	Below lowest rib on the		Pad
`	left anterior axillary line	LL(+)	Reu

standard lead III

- Lead MI, which is similar to standard Lead MIII, which is similar to lead I
 - Change LL and LA of the lead MII.
 - Ш LA (N)



If the electrode position shown above is not available due to chest surgery, attach the electrodes to the root of the limbs or below the clavicles for stable ECG monitoring.

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Connecting the Electrode Lead and Disposable Electrodes

Preparing the Patient Skin

Shave off excessive body hair.

To reduce skin impedance, clean the electrode site with cream or with a cotton pad moistened with the electrode site with cream or with a cotton pad moistened with alcohol. Thoroughly dry the skin with a clean cotton pad.

NOTE

- For a patient with frequent body movement, rub the sites with Skinpure skin preparation gel. However, do not use Skinpure skin preparation gel for sensitive skin.
- Do not place electrodes on a wound or on an inflamed, wrinkled or uneven skin surface.

Attaching Electrodes to the Patient

CAUTION

Do not reuse disposable products.

NOTE

- To maintain good contact between the electrode and skin, check that the paste of the disposable electrode is not dry.
- When contact between the disposable electrode and skin becomes poor, replace electrodes with new ones immediately. Otherwise, contact impedance between the skin and the electrode increases and the correct ECG cannot be obtained.

Refer to the electrode operator's manual for details.



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1. Connect the electrode lead to the electrode.





- 2. Carefully remove the backing paper from the electrode. Avoid touching the adhesive surface.
- Place the electrode on the previously cleaned skin. Pay attention to the electrode lead color and symbol.
 - Fasten the electrode lead wire with surgical tape with an extra length of wire between the tape and the electrode. This lessens the movement of electrode leads by body movement and helps stable monitoring.

Detection and Display of Measurement Condition Electrode Detachment

The """ LED lights on the transmitter or the "CHECK ELECTRODE" message

is displayed on the screen of the monitor in the following cases.

- Electrode is detached from skin.
- Electrode lead is disconnected from the electrode.
- Polarization voltage between the electrode and skin is excessively high.

In these cases, check the cause and if necessary, replace electrodes with new ones.

CAUTION

When the "ELECTRODE OFF" or "CHECK ELECTRODE" message is displayed on the receiving monitor, check electrodes and electrode leads and remove the cause. While "ELECTRODE OFF" or "CHECK ELECTRODE" message is being displayed, there is no ECG monitoring and no alarms.

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Respiration Monitoring

Respiration is monitored by measuring changes in impedance between the RA and LL ECG electrodes. This transmitter sends the changes in impedance to the monitor as a respiration waveform. The monitor displays the respiration waveform and calculates respiration rate. Refer to the operator's manual of the monitor for details.

WARNING

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* Minute ventilation is sensed in rate-adaptive pacemakers by a technology known as bioelectric impedance measurement (BIM). Many medical devices in addition to pacemakers use this technology. When one of these devices is used on a patient with an active, minute ventilation rate-adaptive pacemaker, the pacemaker may erroneously interpret the mixture of BIM signals created in the patient, resulting in an elevated pacing rate.

For more information, see the FDA web site. http://www.fda.gov/cdrh/safety.html



Respiration Measurement Procedure

- 1. Select the electrode lead and disposable electrodes.
- 2. Connect the electrode lead to the ECG/RESP socket.
- 3. Connect disposable electrodes to the electrode lead and attach electrodes to the patient.

After steps 1 to 3 are finished, respiration monitoring automatically starts.

Electrode Position for Respiration Monitoring

Place the RA and LL electrodes so that the lungs are between the electrodes.

NOTE

The optimum electrode positions for ECG measurement of a patient are not always optimum for respiration measurement of the patient. Select positions suitable for both ECG and respiration measurements, or positions which have priority for one measurement.

Electrode Position Examples

NOTE

The following examples are when monitoring with 3 electrodes. ECG cannot be monitored correctly when electrodes are attached as the following examples when monitoring with 6 electrodes.

Position 1

In this position, respiration measurement is available; however, there is a difference in amplitude between different patients.



RA	LL
Right infraclavicular	Fifth intercostal space on the
tossa	left midclavicular line, V4

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Position 2

In this position, the waveform amplitude is usually large and the ECG lead is similar to Lead MII. This position can be generally recommended.



Position 3

In this position, the respiration waveform is optimum, but the ECG lead is unusual.



Position 4

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In this position, the respiration measurement is influenced by the impedance variation of the abdomen, so the cardiac pulse wave included in the respiration wave is reduced. Note that the waveform is inverted in phase compared with the chest movement (the waveform goes down during inspiration). It is difficult to measure the ECG at the same time.



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SpO₂ Monitoring

The SpO₂ monitoring is only available on the ZM-930PA transmitter.

This transmitter sends SpO_2 and pulse waveform to the monitor and displays SpO_2 data and pulse level bar graph on the LCD. Refer to the operator's manual of the monitor for details.

WARNING

- Measurement may be incorrect in the following cases.
 - When the oxyhemoglobin or methemoglobin (HbCO, Met Hb) increases abnormally.
 - When dye is injected in the blood.
 - When using an electrosurgical unit.
 - · During CPR.
 - · When measuring at a site where there are venous pulses.
 - · When there is body movement.
 - · When the pulse wave is small.
- Check the circulation condition by observing the skin color of the measuring site and pulse waveform. Change the measuring site every 8 hours for disposable probes and every 4 hours for reusable probes. The skin temperature may increase at the attached site by 2 or 3°C (4 or 5°F) and cause a burn or pressure necrosis. When using the probe on the following patients, take extreme care and change the measurement site more frequently according to symptoms and degree.
 - · A patient with a fever
 - · A patient with peripheral circulation insufficiency
 - · Neonate or low birth weight infant with delicate skin
- To avoid poor circulation, do not wrap the tape too tight. Check the blood circulation condition by observing the skin color and congestion at the skin peripheral to the probe attachment site. Even for short-term monitoring, there may be burn or pressure necrosis from poor blood circulation, especially on neonates or

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low birth weight infants whose skin is delicate. Accurate measurement cannot be performed on a site with poor peripheral circulation.

• When not monitoring SpO₂, disconnect the SpO₂ probe cord from the transmitter. Otherwise, noise may interfere from the probe sensor and cause incorrect data to be displayed on the transmitter and receiving monitor.

CAUTION

- Do not pull or bend the probe cable or put caster feet on the probe cable. Do not immerse the probe cable in detergents or water. Failure to follow these cautions may cause cable discontinuity, short circuit, skin burn on the patient or incorrect measurement data. Replace any broken probe with a new one.
- Turn off the power of cellular telephones, small wireless devices and other devices which produce strong electromagnetic interference around a patient. Radio waves from devices such as cellular telephones or small wireless devices may be mistaken as pulse waves and the displayed data may be incorrect.

NOTE

When monitoring SpO_2 , monitor ECG at the same time. The ECG electrode lead works as an antenna for transmitting data from the transmitter to the receiving monitor. If ECG is not measured, the telemetry signal may not be received.

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Measurement Procedure

- 1. Select the SpO_2 probe.
- 2. Connect the SpO_2 probe to the SpO_2 socket.
- 3. Attach the SpO_2 probe to the patient.

After steps 1 to 3 are finished, SpO₂ monitoring automatically starts.

Selecting SpO₂ Probe

Select an appropriate probe for the patient.

CAUTION

Use Nihon Kohden specified SpO_2 probe to assure maximum performance from your instrument.

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Model	Subject (Weight)	Attachment Site
Finger Probe TL-201T	Adults, children (Weight more than 20 kg)	Finger
Finger Probe TL-101T	Adults, children (Weight more than 20 kg)	Finger or toe
Multi-site Probe TL-120T	Adults, children, infants (Weight more than 3 kg)	Finger or toe
Foot Probe TL-121T	Infants, neonates (Weight less than 3 kg)	Instep and sole

Reusable Probes

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Disposable Probes

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CAUTION

Use the disposable probe only for one patient. Never reuse the disposable probe for another patient because it causes cross infection.

Model	Subject (Weight)	Attachment Site
TL-251T	Adults (Weight more than 30 kg)	Finger or toe
TL-252T	Children (Weight from 3 to 40 kg)	Finger or toe
TL-253T	Neonates (Weight less than 3 kg)	Instep and sole
TL-260T	Adults, children (Weight more than 3 kg)	Finger or toe
	Neonates (Weight less than 3 kg)	Instep and sole

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Model	Subject (Weight)	Attachment Site
TL-051S/052S	Adults	Finger
	(Weight more than	
	50 kg)	
	Neonates	Instep and sole
Cable length TL-051S: 80 cm	(Weight less than 3	
TL-052S: 160 cm	kg)	
TL-061S/062S	Adults, children	Finger
	(Weight from 15 to	
	50 kg)	
	Children, infants	Тое
Cable length TL-061S: 80 cm	(Weight from 3 to	
TL-062S: 160 cm	15 kg)	

Connecting SpO₂ Probe to the Transmitter

Connect the probe to the SpO_2 socket on the transmitter.



CAUTION

- Do not shake or swing the transmitter holding the cables connected to the transmitter. Otherwise, the transmitter may come off and cause injury to a person or damage surrounding instruments.
- Hold the connector when connecting/disconnecting the probe. If you disconnect the SpO₂ probe holding the cable, it damages the cable.



Attaching the Probe to the Patient

For details, refer to the operator's manual of each probe.

WARNING

To avoid poor circulation, do not wrap the tape too tight. Check the blood circulation condition by observing the skin color and congestion at the skin peripheral to the probe attachment site. Even for short-term monitoring, there may be burn or pressure necrosis from poor blood circulation, especially on neonates or low birth weight infants whose skin is delicate. Accurate measurement cannot be performed on a site with poor peripheral circulation.

CAUTION

- When the attachment site is wet with blood or when the patient has nail polish on, remove the dirt and nail polish before attaching the probe. The transmitted light may decrease due to the blood or nail polish and the measurement data may be incorrect.
- If the skin gets irritated by the tape or redness appears on the skin by the probe, change the attachment site.
- When the probe is attached on an appropriate site with sufficient circulation and the error message confirming the probe attachment repeatedly appears, the probe may be deteriorated. Replace it with a new one.
- Do not use the probe over its stated lifetime. Otherwise the SpO₂ measurement accuracy cannot be guaranteed.
- Do not use damaged or disassembled probe.
- Replace the probe with a new one as specified in the probe manual. If the probe is deteriorated, correct SpO₂ monitoring cannot be performed.
- Do not attach the probe to the same limb that is used for NIBP measurement or an IBP catheter.
- When attached, make sure that the photo emitter and the detector of the probe face each other. Otherwise, SpO₂ cannot be measured properly.

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- When using a disposable probe, be careful when removing the adhesive tape from neonatal skin.
- When removing a disposable probe that is taped to the skin, do not pull the cable part of the probe because this can damage the probe's cable connection.
- Before using the TL-260T multi-site Y probe, be sure to attach the probe to the sponge attachment tape S or L. Do not use the probe without the sponge attachment tape attached. It causes incorrect measurement and may damage the attachment site on the skin.
- When fixing the TL-260T multi-site Y probe with the sponge attachment tape, confirm that the adhesive part of the tape is not on the skin. The adhesive may cause oversensitive symptoms on the skin such as redness or itch. If the adhesive touches the skin, remove it carefully and slowly because neonatal skin is very delicate.
- Do not use a dirty sponge attachment tape. The measurement value may be incorrect.
- Do not pull the cable when removing the TL-260T multi-site Y probe from the sponge attachment tape. Otherwise the cable may get damaged.
- Refer to the probe instruction manual for details.

Starting Measurement

When monitoring starts, SpO_2 and pulse waveform are sent to the monitors and SpO_2 data and pulse level bar graph are displayed on the transmitter LCD.

You can turn off the display of SpO_2 data and pulse level bar graph on the LCD. Refer to the "Turning SpO_2 Data and Pulse Level Bar Graph Display On/Off" section.

CAUTION

When the probe is attached on an appropriate site with sufficient circulation and the error message confirming the probe attachment repeatedly appears, the probe may be deteriorated. Replace it with a new one.



Turning SpO₂ Data and Pulse Level Bar Graph Display On/Off

You can turn off the display of SpO_2 data and pulse level bar graph on the LCD.



Press the CALL key for more than 3 seconds within 10 seconds after turning transmitter power on (after a "peep" sound).

"%SpO₂" is not displayed. When SpO₂ monitoring starts, SpO₂ data and pulse level bar graph are not displayed on the LCD.

To turn SpO_2 display on, turn the transmitter power off and turn the power on again.

Detecting and Displaying Measurement Condition External Light Noise Alarm

CAUTION

When measuring under strong light (surgical light, bilirubin light, sunlight, etc.), cover the probe with a blanket or cloth. Otherwise, noise may interfere.

Strong external light (surgical light or inverter type fluorescent lamp, etc.), may affect SpO₂ monitoring. When external light is too strong to correctly measure SpO₂, the transmitter generates an alarm tone ("pip" sound every 0.5 seconds). Cover the probe attachment site with blanket or cloth.

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Insufficient Light Alarm

When sufficient light cannot be obtained from the photo emitter of the probe, the transmitter generates the following sound and indication.

- (1) "peep" sound (every 1 second)
- (2) pulse level bar graph is maximum (all bars are lit.)
- (3) SpO₂ data is displayed as "- -"
- (4) %SpO2 is blinking.



In this case, change the attachment site to the appropriate site. Refer to the operator's manual of the SpO_2 probe.

Probe Malfunction Alarm

CAUTION

When any of the following occurs, the probe may be broken. Replace it with a new one and check the probe.

- The transmitter generates "pip" sound every 0.25 seconds.
- SpO₂ data is 85% and blinking.

When Measurement Condition is Unstable

 SpO_2 data blinks every 1 second when SpO_2 signal stability decreases and the transmitter cannot detect correct pulse waveform because of patient body movement, poor attachment condition or poor circulation condition at the probe attachment site.

CAUTION

 SpO_2 data blinking every second indicates an unstable pulse waveform and displayed SpO_2 value may be inaccurate. The displayed data may not reflect sudden SpO_2 changes.



(1) Displayed as LED on ZM-920PA (2) (2) (2) (3) SpO ₂ indication (4) pulse level bar graph (3) SpO ₂ data				
Sound	Display	Cause	Countermeasure	
Single "peep" sound for 4 s		The CALL key is pressed while the key is pressed	ed. The sound lasts l.	
	(1)	The battery voltage decreases and battery charge is almost zero.	Replace the batteries with new ones.	
Continuous "peep" sound	All lights are off	Battery is completely discharged.	Replace the batteries with new ones. To stop the sound, turn off the power.	
	(2)	Electrode lead is disconnected from the electrode. Electrode lead is disconnected from the transmitter. Electrode lead discontinuity Electrode is not firmly attached to the skin. Polarization voltage is abnormally high.	Firmly connect the electrode lead to the electrode. Firmly connect the electrode lead to the transmitter. Replace the electrode lead with a new one. Replace the electrode with a new one.	
Intermittent "pip" sound every 0.5 s		SpO ₂ measurement site is under fluorescent light, surgical light, sunlight, etc.	Cover the measurement site with a blanket or cloth.	

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Alarm List

Sound	Display	Cause	Countermeasure
Intermittent "peep" sound every 1 s	(3) (4) all lit (5) blinking	Cannot receive sufficient light from the probe photo emitter.	Attach probe to a site with 6 to 14 mm thickness where sufficient light can be received.
Intermittent "pip" sound every 0.25 s	(3) 85%, blinking	Broken probe	Replace the probe with a new one.
	(3) blinking	Patient body movement	Remove the cause by checking the patient condition and changing the attachment site.
		Probe is not attached securely.	Securely attach the probe.

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Troubleshooting

If the problem still remains after checking the following, contact your Nihon Kohden distributor.

Problem	Cause	Countermeasure
The power cannot be	Batteries are not	Install the batteries correctly.
turned on.	installed correctly.	
	The battery	
	polarity is wrong.	
	Batteries are	Replace the batteries with new
	completely	ones.
	discharged.	
Nothing is displayed	SpO ₂ display is	Turn off the power, and turn on
on the LCD after	turned off.	the power again.
turning the power on.		
(ZM-930PA only)		
Nothing is displayed	The channel of the	Set the correct channel on the
on the monitor after	transmitter and	monitor.
turning the	monitor does not	
transmitter power on.	match.	
Signal receiving	Electrode lead is	Connect the electrode lead to the
condition is poor.	the transmitter	transmitter. ECG electrode lead
	the transmitter.	works as an antenna for
		monitor If ECC is not massured
		the signal may not be received
	Another	Turn the transmitter power off. If
	transmitter of the	the monitor still receives a signal
	same channel is	there is a high probability that
	used near by	another transmitter of the same
	used near by.	channel is used nearby
		Follow the instruction of your
		channel administrator and use
		another transmitter of a different
		channel.
	Signals are	Follow the instruction of your
	mixing.	channel administrator and use
	0	another transmitter of a different
		channel.
	The transmitter is	Contact your Nihon Kohden
	damaged.	distributor.

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Problem	Cause	Countermeasure
3 electrode leads	The transmitter is fixed to	Turn off and on the
are properly	6 lead monitoring.	transmitter power.
attached to the		
patient but 🎯		
LEDs light.		
Heart rate of the	Pacing pulse detection is	Turn off the pacing pulse
patient who is	set to ON on the monitor.	detection.
using an electric		
blanket cannot be		
monitored on the		
receiving monitor.		
ECG baseline is	The gel on the electrode is	Replace the electrode with
thick.	dried out.	a new one.
(Hum is	The gel on the electrode is	
overlapping)	coming off.	
	Electric blanket is used.	Cover the blanket with a
		shield cover.
	Hum filter is set to OFF on	Set the filter to ON.
	the monitor	
Respiration	The gel on the electrode is	Replace the electrode with
waveform	dried out.	a new one.
measurement is	The gel on the electrode is	
unstable.	coming off.	
SpO_2 data is	The probe size is not	Use the appropriate probe
unstable and not	appropriate for the patient.	for the patient.
reliable.	Probe is attached to the	Attach the probe to the
	same limb that is used for	opposite limb. Avoid a site
	NIBP measurement.	where blood circulation
		condition changes greatly.
	Probe attachment	Firmly attach the probe
	condition is poor. Probe is	according to the procedure
	about to detach from the	in the probe operator's
	skin. External light gets in.	manual.
	Measurement site is dirty.	Remove dirt and nail
	Patient is wearing nail	polish.
	polish.	

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Changing the Transmitter Channel

The transmitter channel can be changed by the QI-901PK Channel Writer. To change the channel number, refer to the channel writer manual.

WARNING

The following actions must be taken to properly receive the transmitter signal of the correct patient on the receiving instrument. Otherwise, there may be signal loss or signals may mix causing a serious accident, such as monitoring a different patient.

- Assign a channel administrator in the hospital and only he or she should manage channel assignments.
- The channel administrator must manage the channels in the facility so that there is no signal interference.
- When the transmitter channel is changed, the channel administrator must check that the channel on the receiving monitor is also changed and the signal is properly received.
- The channel administrator must replace the channel number label on the transmitter with the new one after changing the channel.

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Lifetime and Disposal

Disposing of Used Batteries

Replacement

When the ", LED is lit or the ", is displayed on the LCD during operation, the batteries are running out.

Replace the batteries with new ones. When using rechargeable batteries, recharge them.

Disposal

Before disposing of the batteries, check with your local solid waste officials for details in your area for proper disposal. It may be illegal to dispose of these batteries in the municipal waste stream.

Disposing of Disposable Electrodes Lifetime

Replace the disposable electrodes with new ones 48 hours after the start of usage. Otherwise, the gel on the electrode gets dry and adhesive property decreases. This increases skin electrode contact impedance and causes incorrect measurement.

Replace the electrodes with new ones even before 48 hours if the contact between skin and electrode becomes poor.

Disposal

Follow your local laws for disposing of medical waste.



Disposing of the SpO₂ Probe Lifetime

CAUTION

- Do not use the probe over its stated lifetime. Otherwise the SpO₂ measurement accuracy cannot be guaranteed.
- When the probe is attached on an appropriate site with sufficient circulation and the error message confirming the probe attachment repeatedly appears, the probe may be deteriorated. Replace it with a new one.

Reusable probe

Do not use the probe for more than one year after opening the package. If the hours are recorded, you can use the probe for up to 6,000 hours.

Warranty for reusable probe

Nihon Kohden Corporation (NKC) shall warrant this probe against all defects in materials and workmanship for a period of 6 months after the package is opened. NKC or its authorized agents will repair or replace any probe which proves to be defective during the warranty period, provided the probe is used in accordance with the operator's manual.

Disposable probe

Replace the probe with a new one as specified in the probes operator's manual. If the probe is used beyond this time, it deteriorates and correct measurement cannot be performed.

If the probe is dirty with blood or bodily fluids, replace it with a new one, regardless of the lifetime.

Disposal

Follow your local laws for disposing of medical waste.

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Cleaning, Disinfection and Sterilization

Transmitter and Electrode Lead

WARNING

If detergents or dirty liquid spills into the transmitter, clean it and dry it completely before use. If the wet transmitter is used, the patient or anyone in contact with the transmitter may receive an electric shock.

CAUTION

- Before cleaning or disinfecting the transmitter, remove the batteries from the transmitter.
- The transmitter cannot be sterilized.

Cleaning

Wipe the transmitter and electrode leads with a soft cloth moistened with disinfecting alcohol or neutral detergent diluted with water. After cleaning, dry them completely.

Disinfection

CAUTION

- Do not immerse the electrode lead connector in liquid.
- Do not disinfect with hypochlorous acid.
- Use the recommended concentration.

Wipe the outside surface of the transmitter and electrode lead with a non-abrasive cloth moistened with any of the disinfectants listed on the next page. Use the recommended concentration.

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Disinfectant	Concentration (%)
Glutaraldehyde solution	2.0
Hydrochloric alkyl diaminoethylglycine	0.5
Benzalkonium chloride	0.2
Benzethonium chloride solution	0.2
Chlorohexidine gluconate solution	0.5

SpO₂ Probe

Refer to the probe manual.

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Replacing the Battery Case Cover

When the battery case cover is damaged, replace it with a new one. Battery case cover, code no. 6113-046365C.

Other than the battery case cover, there are no serviceable parts for the transmitter.

Repair Parts Availability Policy

Nihon Kohden Corporation (NKC) shall stock repair parts (parts necessary to maintain the performance of the instrument) for a period of 6 years after NKC announces discontinuation of the instrument. In that period, NKC or its distributors will repair the instrument. This period may be shorter than 6 years if the necessary board or part is not available. For discontinuation announcements, contact your Nihon Kohden distributor or representative.

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Specifications

ECG measurement

Channels:
Input range:
DC offset:
Input impedance:
Pacing pulse detection:

±5 mV or more ±500 mV or more 5 MΩ or more (5 Hz) ANSI/AAMI EC13 Based upon Pacemaker pulse rejection Capability

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Respiration measurement

Measuring method: Impedance range:

SpO₂ measurement

Measuring range:0 to 100%, in 1% stepsMeasuring accuracyWhen the measuring accuracy of theSpO2 probe is not considered: ± 1 digit (80% \leq SpO2 \leq

 $\begin{array}{l} \pm 1 \mbox{ digit } (80\% \le \mbox{SpO}_2 \le 100\%) \\ \pm 2 \mbox{ digit } (50\% \le \mbox{SpO}_2 \le 80\%) \\ \mbox{Less than } 50\% \mbox{ is not specified.} \end{array}$

Impedance method 0 to 2 k Ω

When the measuring accuracy of the SpO₂ probe is considered:

 $\pm 2 \text{ digit } (80\% \le \text{SpO}_2 \le 100\%)$ $\pm 3 \text{ digit } (70\% \le \text{SpO}_2 \le 80\%)$ Less than 70% is not specified.

Transmitter

FCC regulation:

Field strength limits: Undesired emission: FCC part 95 Subpart-H Wireless Medical Telemetry Service (WMTS) <200 mV/m (at 3 m) below 960 MHz: 200 μV/m (at 3 m) above 960 MHz: 500 μV/m (at 3 m)

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Antenna: Transmission channel: Transmission frequency range: Channel spacing: Modulation scheme: Occupied bandwidth: Effective radiated power:	ECG electrode lead indicated on the transmitter 608.0250 to 613.9750 MHz 50 KHz (25 KHz when interleaved) FSK (Frequency Shift Keying) <20 KHz 1.0 mW (conducted)
Safety standards	
Safety standard:	CSA C22.2 No. 601-1 M90 (1994) IEC 60601-1 (1988) IEC 60601-1 Amendment1 (1991) IEC 60601-1 Amendment2 (1995) IEC 60601-1-2 (1993) IEC 60601-2-27 (1994)
According to the type of protection	
against electrical shock:	INTERNALLY POWERED EQUIPMENT
According to the degree of protection	
against electrical shock:	
ECG/impedance method respiration:	DEFIBRILLATION-PROOF
	TYPE CF APPLIED PART
SpO ₂ :	DEFIBRILLATION-PROOF
	TYPE BF APPLIED PART
According to the degree of protection	
against harmful ingress of water: According to the degree of safety of application in the presence of a FLAMMABLE ANAESTHETIC MIXTURE WITH AIR, OR WITH	Ordinary equipment
OXYGEN OR NITROUS OXIDE:	Equipment not suitable for use in the presence of FLAMMABLE ANAESTHETIC MIXTURE WITH AIR, OR WITH OXYGEN OR NITROUS OXIDE
According to the mode of operation:	CONTINUOUS OPERATION

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Water resistance

Water does not get inside the transmitter except for the battery case when immersed in water up to 30 cm deep for 3 minutes.

Power requirements			
Battery type:	two AA type alkaline dry cell batteries,		
	manganese dry cell batteries, NiCd		
	rechargeable batteries, NiMH batteries		
Battery lifetime:			
ZM-930PA:	approximately 3 days		
	(with alkaline batteries, measuring ECG,		
	respiration and SpO ₂ of approximately 60		
	kg weight adult male patient at the pointing		
	finger)		
	approximately 4 days		
	(with alkaline batteries, measuring ECG		
	and respiration only)		
ZM-920PA:	approximately 4 days		
Environment			
Operating environment			
Operating temperature:	5 to 40°C, 41 to 104°F		
Operating humidity:	30 to 85% (non-condensing)		
Operating atmospheric pressure:	70 to 106 kPa		

(Op ng a sp Operating voltage:

Storage environment

Storage temperature: Storage humidity: Storage atmospheric pressure:

Dimension and Weight

Dimension: Weight (without batteries): 1.6 to 3.2 V

-20 to 65°C, -4 to 149°F 15 to 95% (non-condensing) 70 to 106 kPa

78 W \times 122 H \times 26 D (mm) ZM-930PA: about 180 g ZM-920PA: about 165 g

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Standard Accessories



Name	Q'ty	Supply code
Strap	1	Y233

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Options

CAUTION

Use only Nihon Kohden electrodes, electrode leads and SpO₂ probes to assure maximum performance from your instrument.

ECG/RESP

Name	Application	Model	Q'ty	Supply code
Electrode lead	3 electrodes, clip type, lead length 80 cm	BR-903PA	1	K911A
	3 electrodes, snap type, lead length 80 cm	BR-913PA	1	K910B
	6 electrodes, clip type, lead length 80 cm	BR-906PA	1	K912A
	6 electrodes, snap type, lead length 80 cm	BR-916PA	1	K915A

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SpO₂

Name	Cable length	Model/ Code No.	Q'ty	Supply code
Finger probe	1.6	TL-201T	- 1	P225F
		TL-101T		P224A
Multi-site probe		TL-120T		P225C
Foot probe		TL-121T		P225D
SpO ₂ probe (disposable, for adult, BLUPRO)		TL-251T	5	P201A
SpO ₂ probe (disposable, for child, BLUPRO)		TL-252T		P201B
SpO ₂ probe (disposable, for neonate, BLUPRO)		TL-253T		P201C
Multi-site Y probe (disposable, for adult, child, neonate, BLUPO)		TL-260T		P205A
SpO ₂ probe (disposable,	0.8	TL-051S	5	P228A
for adult, neonate)	1.6	TL-052S		P228B
SpO ₂ probe (disposable,	0.8	TL-061S		P229A
for child, infant)	1.6	TL-062S		P229B
COTTONY tape		340703	20	P259
Sponge attachment tape S for TL-260T			24	P260A
Sponge attachment tape L for TL-260T				P260B
Foam tape			4×25 package	P260

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