Chapter 10

Maintenance

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10 Maintenance

- Handling

After Use / Display Panel

This section describes precautions for handling the equipment.

Handling After Use

- ∠Do not apply excessive force when disconnecting the cables. Always pull on the connector housing and not on the cable.
- ∠Always check for adequate supply of disposable accessories such as ECG electrodes. If any shortage, contact our service representative and supply as necessary.

Handling the Display Panel

- ∠The display panel utilizes exclusive fluorescent light for the backlight.

 As this fluorescent light tube has product life cycle, it needs to be replaced periodically. If the display becomes dark, scintillates, or does not light, contact your nearest service representative.
- ∠The LCD used for the display panel utilizes highly accurate picture elements of pixels over 99.99%, but there may be an absence (less than 0.01%) or constant lighting of pixels.

Storage -

Device / Recording Paper

This section describes about the storage of the device and recording paper.

Storing the Device

- ≤Store in a place where the device will not be exposed to splashing water.
- Store in a place where the device will not be adversely affected by atmospheric pressure, temperature, humidity, ventilation, sunlight, dust or atmosphere containing salt or sulfur.
- ∠The following environmental conditions should be observed when storing the device.

Storage Temperature : - 10 ~ 60?C Storage Humidity : 10 ~ 95% (at 60?C)

Storing the Recording Paper

The DS-7100 system utilizes heat sensitive recording paper. If placed in a high temperature for long period of time, the print may become indistinct, and unable to read. When storing, follow the precautions below.

- ≤Store in a place where light is shut off and avoid direct sunlight.

- Avoid using adhesive agents other than water based glue.

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Storage

Cleaning

Display Panel and Housing

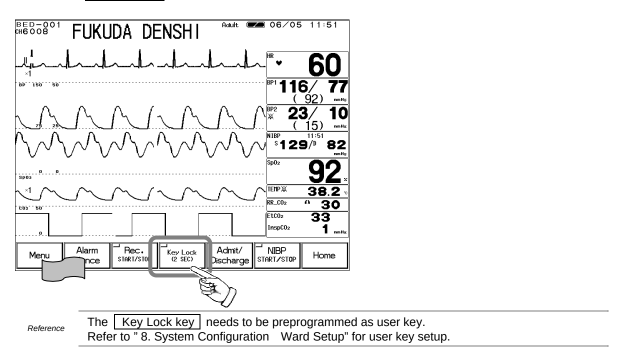
This chapter explains about the cleaning of the device and sensors.

Cleaning the Display Panel

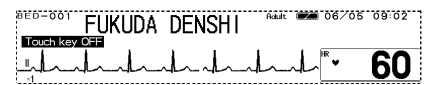
Since this device incorporates a touch screen, finger prints and other stains are likely to appear on the display panel.

Follow the procedure below to clean the display panel.

1. Press the Key Lock key for more than 2 seconds.



2. Clean the touch panel.



While the "Touch key OFF" message is displayed, the touch panel key will be deactivated. If "LEAD OFF" or other message is displayed, the key lock message will not be displayed.

- 3. Wipe the touch panel using cleaning cloth.
- **4.** Press again the Key Lock key for more than 2 seconds. The message will disappear and the keys will be activate again.

∆ CAUTION	 Do not clean the touch panel using strong acid. A special coating is applied to the surface of the touch panel. surface with the soft cleaning cloth provided as optional access with commercially available eyeglass cleaning cloth. 	•
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Cleaning the Housing

Clean the housing using tightly squeezed gauze or an absorbent cotton cloth dampened with alcohol or a neutral cleanser

alconol of a fleutral	cieanser.
^ CAUTION	 Clean the equipment frequently so stains can be removed easily. To prevent injury, it is recommended to wear gloves when cleaning the equipment. Do not allow liquids or cleaning solution to enter the monitor or connectors. Do not use organic solvents, thinner, toluene and benzene to avoid damaging the resin case. Do not polish the housing with abrasive or chemical cleaner. When sterilizing the entire room using a spray solution, pay close attention not to have liquids get into the monitor or connectors. Use only neutral detergent to clean the housing. Do not use chemical cloth, scrub brush, abrasive, polishing powder, hot water, volatile solvent and chemicals (cleanser, thinner, benzine, benzol, and synthetic detergent for house and furniture), or sharp-edged tools. The surface resin coating may be damaged, resulting in discoloration, scratches, and other problems.

Disinfecting the Blood Pressure Transducers

Disinfect the blood pressure transducers according to the manufacturer's guidelines.

Disinfecting the Temperature Probes

Disinfect the Temperature Probes according to the manufacturer's guidelines.

Disinfecting the CO₂ Filter Line

Disinfect the Filter Line/Capno Line according to the manufacturer's guidelines.

Cleaning and Disinfecting the SpO₂ Transducer

Cleaning and Disinfecting the NIBP Cuff

Cleaning and Disinfecting the ECG lead Cable

∠Do not soak the sensor in water or antiseptic solution.

Wipe the ECG lead cable with disinfectant such as 70% alcohol. Do not disinfect by applying radioactive rays, steam, or ethylene oxide.

- Battery -

Handling the Battery Pack

This section describes about the handling and storage of the battery pack.

Handling the Battery

- ∠The battery pack can be continually used for more than 300 times (or about 1 year) under normal temperature, but the continuous use will degrade the battery and shortens the usable time.
- When the battery operation time becomes short even after it is fully charged, the battery pack needs to be replaced.
- Mhen the charge time of the battery pack becomes short, the battery pack needs to be replaced.
- Mhen the battery pack level becomes low, charge the battery well in advance for the next use.

Storing the Battery

To take advantage of the characteristic of battery pack, pay attention to the following when storing.

Storage Temperature and Humidity

Store in an environment specified below without corrosive gas.

Storage Period	Storage Temperature	Storage Humidity
Within 30days	- 20 ~ 60?C	
30 days ~ 90 days	- 20 ~ 45?C	65?20%
90 days ~ 1year	- 20 ~ 35?C	

[∠]Do not store in an environment outside the specified temperature range or excessive high
humidity. This may result in leakage caused by expansion/contraction inside the battery pack,
or rusting of the metal part.

Long-term Storage

If left installed in the monitor for long period of time, the electrolyte may leak, or inactivate the battery which degrades the capacity recovery after storage. Therefore, always remove the battery from the monitor when storing for long period of time. Contact our service representative when removing the battery.

This section explains the daily check and periodic check items of the device.

About the Maintenance Check

Periodic inspection must be performed. When reusing the device which was left unused for a while, always check that the device operates properly and safely before use.

To ensure safety, reliability, and high performance, a "Daily Check" and "Periodic Inspection" must be performed. We are not liable for any accident arising from lack of maintenance.

▲ CAUTION	✓ Do not open the housing of this device.✓ Avoid alcohol or other liquids from getting into the equipment.
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Daily Check

Perform daily inspection using the "Daily Check List" on the next page.

Periodic Check

Periodic inspection of medical electronic equipment is mandatory to prevent failures and accidents and to ensure safety and reliability.

Periodic maintenance may be performed by each medical institution or by a third party by concluding a "Maintenance Contract".

For more details, contact your nearest service representative.

Time-Change Components

To ensure reliability of safety, function, and performance of this device, the time-change components must be replaced periodically. When replacing, contact our service representative.

EtCO₂ Unit Replacing Period: 7,000 hours (approx. 10 months of

continuous use)

LCD Unit, Inverter UnitReplacing Period: 50,000 hours or 6 years

NIBP Unit Replacing Period: 100,000 times of use or 6 years

Short Term Backup Battery Replacing Period: 4 years ~ 6 years according to the used

frequency

Long Term Backup Battery Replacing Period: 6 years

Battery Pack Replacing Period: 1 year or 300 times of charging /

discharging.

⚠CAUTION The time-change components must be replaced at specified period.

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Maintenance Check

Daily Check List

		No.		
nspected Date	Inspected by	Location		
Device Type	Serial No.	Date of Purchase	Date of Purchase	
ltem	Details	Criteria	Judgement	
Appearance	Visually check the exterior for scratches, cracks, deformation,	No abnormality should be found.	OK / NG	

Item	Details	Criteria Judgement	
Appearance	Visually check the exterior for scratches, cracks, deformation, and rust.	No abnormality should be found.	OK / NG
	Check whether the unit is installed on a level surface.	The installation area must be level and free from vibration and shock.	OK / NG
Installation	Check whether the unit is installed in a place susceptible to adverse environment.	The environmental condition (ex. temperature, humidity) of the installed place should be as specified. The unit should not be subjected to splashing water.	OK / NG
Functions	Turn ON the monitor, and check whether it operates normally.	The home display appears, and the lamp located at the right side of the display panel lights.	OK / NG
		The date and time should be correct.	OK / NG
Cables	Visually check all cables for any damage.	No damage should be found.	OK/ NG
Periodic Inspection	Check the date of previous periodic inspection.	Should be within 1 year.	OK / NG
CO₂ Calibration (DS-7141)	Check the date of previous calibration date. Previous Date Day Year Month	Should be within 6 months.	OK / NG

Troubleshooting

This section explains the troubleshooting for each case.

ECG

The "LEAD OFF" message is displayed.

Cause : The electrode is detached, or is not making good electrical contact with the skin.

Solution: • Check if the electrodes are properly attached.

· Replace the electrode, or check the lead cable.

The "ECG failed" message is displayed.

Cause 1 : The ECG amplitude is 0.25mV or below for the waveform size of \times 1, \times 1/2, \times 1/4, and 0.150mV or below for the waveform size of \times 2, \times 4.

Solution: Change the electrode attachment site, or select the lead with higher QRS amplitude.

Note: Using 4-electrode or 5-electrode instead of 3-electrode allows more accurate QRS detection.

Cause 2: The electrode contact is poor.

Electrical blanket or other noise source is near the patient.

Solution: Attach the electrode firmly.

- · Replace the lead cable if defective.
- If any noise source is near the patient, locate it away from the patient as much as possible.

ECG waveform contains noise.

The "Artifact" message is displayed.

Cause 1: The electrode contact is poor.

Electrical blanket or other noise source is near the patient.

Solution: Attach the electrodes firmly.

- · Replace the lead cable if defective.
- If any noise source is near the patient, locate it away from the patient as much as possible.

Cause 2 : EMG is interfering.

Solution: • Change the electrode site to a location where EMG will less likely to interfere.

· Select ESIS mode for the filter mode.

Note : Selecting a ESIS mode for the filter mode will decrease the QRS amplitude and may result in not counting the heart rate.

The "Check electrode" message is displayed.

Cause : The electrode contact with the skin is poor. There is substantial contact resistance

between the electrodes.

Solution: Replace all the electrodes.

Use the electrodes of the same type.

The "ECG unit error" message is displayed.

Cause : A communication error with the ECG measuring unit exists.

Solution: The breakage of wire or failure of the ECG unit can be considered.

Contact our service representative.

The measured data is displayed as " $\times \times \times$ ".

Cause : The heart rate is outside the measurement range.

Solution: • Check the electrode application.

· Replace the electrode, or check the lead cable.

Heart rate is not counted. Heart rate is low.

Cause : The ECG waveform amplitude is below the QRS detection level (0.3mV). Solution : Change the electrode site, or select a lead with higher QRS amplitude.

Note : Using 4-electrode or 5-electrode instead of 3-electrode allows more accurate QRS

detection.

Also, if large amount of noise is interfering, the noise may be erroneously detected as QRS. It is recommended to change the electrode site and increase the ECG

amplitude.

Heart rate is not counted, and "LEAD OFF" message is displayed.

Cause : The electrode of the displayed lead type is detached, or is not making good electrical

contact with the skin.

Solution: • Check the electrode application.

· Replace the electrode, or check the lead cable.

Artificial pacemaker is not displayed.

Cause : On the admit / discharge menu, Not used is selected for the pacemaker use.

Solution: Select Used for the pacemaker use.

The "Pacemaker error" message is displayed.

Cause : The pacemaker pulse is detected 16 pulses or more per second.

Solution 1 : Attach the electrodes firmly.

- · Replace the lead cable if defective.
- If any noise source is near the patient, locate it away from the patient as much as possible.

Solution 2: If the patient is not wearing a pacemaker, set to Not used for the pacemaker use in the patient admit/discharge menu.

The "ECG not connected" message is displayed.

Cause : When the ECG relay cable is disconnected during ECG monitoring, this message will be displayed.

Solution 1 : To cease monitoring, press the Alarm Silence key to clear the message and silence the alarm.

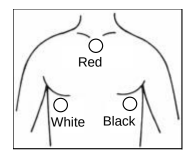
Solution 2 : To continue monitoring, plug in the ECG relay cable. This will clear the message and silence the alarm.

Respiration

The "CVA detected" message is displayed.

Cause : Heartbeat is interfering and superimposed on the respiration waveform.

Solution: Place the electrode as shown below where the heartbeat will be less likely to interfere.



"0" is displayed for respiration rate, or apnea alarm is generated.

Cause : The respiration waveform amplitude is below the detection level (0.20).

Solution: Change the electrode site.

The respiration waveform and respiration rate is not displayed.

Cause 1: The ECG relay cable designed for electrosurgical knife is used.

Solution: The impedance respiration can not be measured if the cable designed for

electrosurgical knife is used. Use the standard ECG relay cable if not using the

electrosurgical knife.

Cause 2: The impedance respiration measurement is ceased.

Solution: Turn ON the impedance respiration measurement on the admit / discharge menu or

RESP configuration menu.

Note : If the pacemaker with the minute ventilation measuring function is used, turn OFF the

impedance respiration measurement. Otherwise, both the pacemaker and the monitor

will not be able to perform accurate measurement.

The measured data is displayed as " $\times \times \times$ ".

Cause : The respiration rate is outside the measurement range.

Solution: • Check the electrode application.

• Replace the electrode, or check the lead cable.

Invasive Blood Pressure

The "BP1 Transducer OFF", "BP2 Transducer OFF" message is displayed.

Cause : The transducer for BP1 or BP2 is not connected.

Solution: Connect the transducer.

The "BP1 not zero balanced", "BP2 not zero balanced" message is displayed.

Cause : The BP zero balance has not been performed since the power is turned ON. Solution : Open the three-way cock of the transducer to air and perform zero balance.

The measured data is displayed as "- - -".

Cause : The BP zero balance has not been performed since the power is turned ON. Solution : Open the three-way cock of the transducer to air and perform zero balance.

BP value and waveform are not displayed properly.

Cause : Blood pressure line has not been zero balanced.

Solution: Open the three-way cock of the transducer to air and perform zero balance.

The measured data is displayed as " $\times \times \times$ ".

Cause : The BP value is outside the measurement range.

Solution: Perform zero balance again.

The "BP not connected" message is displayed.

Cause : When the BP interface cable or 2ch BP conversion cable is disconnected during BP

monitoring, this message will be displayed.

Solution 1 : To cease monitoring, press the Alarm Silence key to clear the message and

silence the alarm.

Solution 2: To continue monitoring, plug in the BP interface cable or 2ch BP conversion cable.

This will clear the message and silence the alarm.

The "Incorr. BP cable" message is displayed.

Cause : The cable other than 2ch BP conversion cable is plugged in to the BP connector.

Solution: Use the 2ch BP conversion cable.

SpO₂

The "Check SpO₂ sensor" message is displayed.

Cause : Sensor is detached from the patient.

Solution 1: Check if the sensor part is properly attached to the patient.

Solution 2: Check if the light emitting part and light receiving part of the sensor LED is aligned.

The "Pulse search" message is displayed.

Cause : The amplitude of the pulse waveform is low, or the sensor is not positioned correctly. Solution : Check if the light emitting part and light receiving part of the sensor LED is aligned.

The "No pulse detect" message is displayed.

Cause : The amplitude of the pulse waveform is low, or the sensor is not positioned correctly. Solution : Check if the light emitting part and light receiving part of the sensor LED is aligned.

The "Motion Artifact" message is displayed.

Cause : There is excessive body motion of the patient.

Solution: Change the sensor position where the body motion will have less effect.

The pulse waveform is not displayed, or interrupted

Situation: "Check SpO₂ sensor" is displayed.

Cause 1: The amplitude of the pulse waveform is low, or the sensor is not positioned correctly. Solution: Check if the light emitting part and light receiving part of the sensor LED is aligned.

Cause 2 : Sensor is defective. Solution : Replace the sensor.

Cause 3 : SpO_2 sensor is not firmly connected to the SpO_2 input connector.

Solution: Make sure the SpO_2 sensor is securely connected.

Cause 4: Sensor is exposed to light.

Solution: Place a black or dark cloth over the sensor to avoid direct sunlight. Also when not used, avoid placing the sensor in light or unplug the sensor from the connector.

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Troubleshooting

The SpO₂ measurement is unstable.

Cause : There is excessive body motion of the patient which disables correct measurement.

Solution: 1. Have the patient lie still as much as possible.

2. Relocate the sensor, or change the sensor to which the body motion will have less influence.

The "SpO₂ unit error" message is displayed.

Cause 1: There is a failure of communication with the SpO₂ measurement unit.

Solution: Breaking of wire or SpO₂ unit failure can be considered.

Contact our service representative.

Cause 2 : Sensor is defective. Solution : Replace the sensor.

The "SpO₂ sensor fault" message is displayed.

Cause : Sensor is defective. Solution : Replace the sensor.

The "SpO₂ not connected" message is displayed.

Cause : When the SpO₂ relay cable is disconnected during SpO₂ monitoring, this message will

be displayed.

Solution 1: To cease monitoring, press the Alarm Silence key to clear the message and

silence the alarm.

Solution 2: To continue monitoring, plug in the SpO₂ relay cable. This will clear the message

and silence the alarm.

Non-Invasive Blood Pressure

The cuff is not inflated although the pump is operating.

Cause1: The air hose is not firmly connected, and the air is leaking.

Solution: Check if the air hose is properly connected.

Cause 2: The cuff size is not corresponded to the selected patient type. Solution: Check if the cuff size is corresponded to the selected patient type.

The monitor repeats the measurement, or "- - -" is displayed for the numeric data.

Cause 1: The measurement accuracy is not reliable due to body motion artifact.

Solution: Have the patient stay still as much as possible during the measurement.

Cause 2: The pulse is too small to acquire reliable measurement accuracy.

Solution: Check if the cuff application is proper, and if the cuff size is corresponded to the

selected patient type.

The "Check NIBP hose" message is displayed.

Cause : The applied pressure to the cuff has exceeded the maximum limit. The measurement time has exceeded the maximum limit.

Solution: Check if the cuff application is proper, if the cuff size is corresponded to the selected patient type, or if the air hose is not bent. After checking the above, perform the measurement again.

If the same message is displayed again, a failure of the equipment can be considered. Cease the measurement, and contact our service representative.

The "NIBP unit error" message is displayed.

Cause : The zero balancing before the measurement has failed, and measurement could not be

started.

Solution: The body movement or other artifact may cause zero balance failure. During the

measurement, have the patient stay still as much as possible.

If the same message is displayed again, the failure of the equipment can be considered.

Cease the measurement, and contact our service representative.

The time of measurement disappears and the numeric data is displayed as "- - -".

Cause : The NIBP data will be erased when the preprogrammed NIBP erase time has elapsed. Solution : Select the appropriate time for NIBP data erase time from 10min, 30min, 60min, 24hrs

which best fits the monitoring purpose.

Temperature

The "Wrong Temp Probe" message is displayed.

Cause 1: The YSI-700 is used.

Solution: Use the YSI-400 temperature probe for measurement. The YSI-700 can not be used

with the DS-7100 series.

Cause 2: There is a contact failure of the temperature probe.

Solution: Check if the temperature probe is properly inserted.

The numeric data is displayed as " $\times \times \times$ ".

Cause : The temperature measurement is outside the measurement range.

Solution: Check if the temperature probe is properly inserted.

The "TEMP not connected" message is displayed.

Cause : When the temperature sensor is disconnected during temperature monitoring, this

message will be displayed.

Solution 1 : To cease monitoring, press the Alarm Silence key to clear the message and

silence the alarm.

Solution 2: To continue monitoring, plug in the temperature sensor. This will clear the message

and silence the alarm.

The "TEMP auto check" message is displayed. The numeric data is displayed as "- - -".

Cause : The temperature is calibrated once every hour on this monitor. During calibration, the

numeric data will be displayed as "- - -".

Solution: The calibration will complete in 10 seconds. If the calibration does not complete within

10 seconds, cease the measurement and contact our service representative.

The "TEMP unit check" message is displayed.

Cause : Error is detected during temperature calibration.

Solution: A unit failure can be considered. Cease the measurement and contact our service

representative.

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CO₂ Concentration

The "Check filter line" message is displayed.

Cause : The sampling tube is clogged. Solution : Replace the sampling tube.

The "Self-diag CO₂" message remains displayed.

Cause : An error has occurred to the self-check procedure at power ON.

Solution: The CO₂ unit failure can be considered.

The "Initializing CO₂" message does not disappear.

Cause : An error has occurred during the initialization at power ON.

Solution: The CO₂ unit failure can be considered.

The "Check CO₂ unit" message is displayed.

Cause 1: The exhaust connector is clogged.

Solution: After checking the exhaust system and removing the clog, press the "Restart CO2" key

on the CO₂ configuration menu.

Cause 2: The sampling tube or nasal prong is clogged.

Solution: After checking the inhalation system and removing the clog, press the "Restart CO2"

key on the CO₂ configuration menu.

Cause 3 : The CO_2 unit needs to be replaced. Solution : Contact our service representative.

The "CO₂ unit error" message is displayed.

Cause $\,\,$: There is a communication error with the CO_2 unit.

Solution: The break of wire or CO₂ unit failure can be considered.

Contact our service representative.

There is substantial measurement error.

Cause 1: 20 minutes have not yet elapsed since the power is turned ON.

Solution: For 20 minutes from turning ON the power, there will be a substantial measurement

error.

Cause 2: The calibration is not properly performed.

Solution: Perform CO₂ calibration again.

The "CO₂ not connected" message is displayed.

displayed.

Solution 1 : To cease monitoring, press the Alarm Silence key to clear the message and

silence the alarm.

Solution 2: To continue monitoring, plug in the filter line. This will clear the message and silence

the alarm.

Recorder

No recording is performed.

Situation: The "Paper Out" message is displayed on the upper left of the screen.

The "Paper Out" message is displayed on the Rec. START/STOP key.

Cause : There is no recording paper in the recorder magazine. Solution : Install a new pad of paper into the paper magazine.

Situation: The "Magazine Open" message is displayed.

Cause: The paper magazine is open.

Solution: Close the magazine.

Situation: The "Paper jammed" message is displayed.

Cause: The paper is jammed.

Solution: Open the magazine and install the paper correctly.

Situation: No message is displayed, but recording can not be performed.

Cause : The recording paper is not correctly installed. The front and backside of the paper is

set oppositely.

Solution: The "END" printed side of the paper should be facing down in the magazine.

The second waveform and third waveform are not recorded.

Situation: The second waveform and third waveform are not recorded for manual recording or

alarm recording.

Cause : The second waveform and third waveform are not set on the recording setup menu. Solution : Set the second waveform and the third waveform on each recording setup menu.

The "Recorder error" message is displayed.

Cause : The thermal head temperature has increased.

Solution: A damage to the thermal head can be considered.

Contact our service representative.

Telemetry

The "Telemetry unit error" message is displayed.

Cause : There is a communication error with the telemetry transmission unit.

Solution: The breaking of wire or telemetry transmission unit failure can be considered.

Contact our service representative.

There is no reception at the telemetry center.

Cause : The channel ID or group ID is not corresponded with the telemetry receiver.

Solution: Set the correct channel ID and group ID.

The BP waveform of 100mmHg or above can not be properly received.

Cause : The BP waveform and scale is not corresponded.

Solution: When BP waveform is above 100mmHg, set the BP scale above 100mmHg.

Cause : A system error has occurred.

Solution: Turn off the power, unplug the power cable, and contact our service representative.

The "Adjusting" message is displayed. Numbers are displayed large on the display.

Cause : This is the test mode. Stop using the device immediately. Solution : Restart the system. The test mode will be cancelled.

If the same situation is observed again, contact our service representative.

Turn off the DIP switch No.1.

The data is initialized each time the power is turned ON.

Cause 1: The internal switch is set to initialize.

Solution: The internal switch setting needs to be changed. Contact our service representative.

Set the rotary switch to 0.

Cause 2: The battery for backup memory is depleted.

Solution: The battery needs to be replaced. Contact our service representative.

The display is not clear.

Cause 1: The display brightness is not adjusted.

Solution: Due to the LCD display characteristic, the visible range is limited. Adjust to the

appropriate brightness.

Cause 2: The monitor is set to the night mode.

Solution: Cancel the night mode.

The system does not start although the power switch is turned ON.

Cause 1: The power cable is not connected.

The battery is not charged.

Solution: Turn off the power and connect the power cable. If the battery is not charged, use the

power cable until the battery charging is complete.

Cause 2: Incorrect IC card is inserted. Solution: Turn off the DIP switch No.8.

The clock is often delayed.

Cause : The battery for the backup memory is depleted. Check if the time is delayed when the

power is turned off.

Solution: The battery needs to be replaced. Contact our service representative.

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Troubleshooting

Battery

The operation time is short although the battery is charged.

Cause 1: The battery life has expired.

Solution: The battery pack is a consumable product. Replace it once a year.

Cause 2: The ambient temperature is too high or too low.

Solution: For safety, the charging operation will be in a standby mode when the battery pack

temperature becomes excessively high or low.

The charging will automatically resume when appropriate temperature is reached.

Charge the battery in an ambient temperature of 10 ~ 30?C.

The charge lamp on the patient monitor does not light.

Cause 1: The AC power cable is disconnected.

Solution: Plug in the AC power cable.

The battery pack can be charged only during the AC operation.

Cause 2: The battery pack is not installed.

Solution: The battery pack is optional. If a battery pack is required, contact our service

representative and install the battery pack.

Cause 3: The battery life has expired.

Solution: The battery pack is a consumable product. Replace it once a year.

During the charging procedure, the charge lamp (orange) does not switch to charge complete status (green) and extinguishes.

Cause 1: The battery pack temperature is too high or too low.

Solution: For safety, the charging operation will be in a standby mode when the battery pack

temperature becomes excessively high or low.

The charging will automatically resume when appropriate temperature is reached.

Cause 2: The breakdown of battery pack can be considered.

Solution: If the charging operation does not complete within the specified charging time, the

charging operation will cease for safety purpose.

Contact our service representative and replace the battery pack.

Cause 3: The battery life has expired.

Solution: The battery pack is a consumable product. Replace it once a year.

The "Charge battery" message is displayed.

Cause : The AC power cable is disconnected.

Solution: Plug in the AC power cable.

The battery pack can be charged only during the AC operation.

11 Technical Information

Chapter 11 Technical Information

-	Specification/Performance - · · · · · · · · · · · · · · · · · ·
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	Patient Admit / Discharge
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- Specification/Performance -

This section states the specification and performance of this equipment.

Specification

Size

260 (W) × 197 (D) × 264 (H) mm (not including the protrusion)

Weight (not including the battery)

DS-7141 : 5.3kg DS-7101L : 5.1kg DS-7101LT : 5.2kg

Environmental Condition

Operating Temperature : 10 ~ 40 °C

Operating Humidity : 30 ~ 85% (without condensation)

Transport / Storage Temperature : - 10 ~ 60 °C

Transport / Storage Temperature : 10 ~ 95% (at 60 °C)

Safety

General Standard : EN60601-1:1990

(Medical electrical equipment - Part 1: General requirements for

safety)

Amendment A1 to EN 60601-1:1993 Amendment A2 to EN 60601-1:1995

EMC Standard : IEC 60601-1-2:1993

(Medical electrical equipment - Part 1: General requirements for safety

- 2.Collateral standard:

Electromagnetic compatibility - requirements and

tests. IEC 601-1-2:1993)

Withstand Voltage : Class I, Type CF

Power Requirements

Voltage	AC100 ~ 240V ?10%	DC14V ?10%
Frequency	50/60Hz	?
Power Consumption	82VA	60W

Performance

Device : 8.4 inch TFT Color LCD Control : Touch Screen Type Waveform Trace : Stationary Trace

Waveform Speed : ECG / SpO₂ / BP (6.25mm/s, 12.5mm/s, 25mm/s)

RESP / CO₂ (6.25mm/s, 12.5mm/s, 25mm/s)

Parameter : ECG, RESP, TEMP, SpO₂ (Arterial Oxygen Saturation), BP1, BP2,

NIBP, CO_2 concentration

Recording

Recording Method : Thermal Array Type

Recording Speed : 25mm/s

Recording Waveform : Max. 3 waveforms

Recording Width : 50mm

Operation

Touch Screen : Eight-Wire Resistive Analog Touch Screen

ECG

Lead Type : Wired 3-electrode, 4-electrode, 5-electrode

Frequency Characteristic: 40Hz / 15Hz Input Impedance : 5M? or above Max. Input Voltage : ±10mV

Polarization Voltage : ±825mV or above

Common Mode

Rejection Ratio : 80 dB or above

HR Measurement Range: Adult $0, 12 \sim 300 \text{bpm} \pm 3\% \text{ or } \pm 5 \text{bpm}$

Neonate $0, 30 \sim 300$ bpm $\pm 3\%$ or ± 5 bpm

Waveform Size Selection: 1/4, 1/2, 1, 2, 4 **Defibrillation Proof** : Provided

Respiration

Method : Impedance Method

Frequency Characteristic: 1.5Hz (adult, child) / 2.5Hz (neonate)

Current : 100 µ A or lower : 0, 4 ~ 150Bpm ± 5Bpm Measurement Range

Temperature

Method : Thermistor Method : only YSI-400 series Probe Measurement Range $: 0 \sim 50$?C ± 0.2 ?C No. of Channel : 1 channel

SpO₂ (Arterial Oxygen Saturation)

Method : 2 Wavelength Pulse Wave Method

Measurement Range : 0% ~ 100%

Resolution : 1%

Accuracy : Adult at 70 ~ 100% ± 2%

at 0 ~ 69% not specified

Neonate at 70 ~ 100% ± 3%

at 0 ~ 69% not specified

Measurement Range : 20 ~ 250bpm ± 3bpm

Specification / Performance

Blood Pressure

Transducer Sensitivity : $5 \mu V / V / mmHg$ Measurement Range : $-50 \sim 300 mmHg$

Frequency Characteristic: DC ~ 6Hz / 8Hz / 12Hz / 40Hz

Accuracy : ±2% of full scale or within ±1mmHg

Zero Balance Range : within ±150mmHg

Measurement Range : Adult $20 \sim 300$ bpm $\pm 3\%$

Neonate 30 ~ 300bpm ± 3%

Channel : 2 channels

Non-Invasive Blood Pressure

Method : Oscillometric Method
Measurement Range : 10 ~ 280mmHg
Resolution : 1mmHg
Accuracy : ±4mmHg

Measurement Range : 40 ~ 240bpm ± 5%

CO₂ Concentration (DS-7141)

Method : Infra-Red Solid-State Method, Microstream Method

Measurement Range : 0 ~ 99mmHg

0-20min

0 ~ 38mmHg: ± 4mmHg 39 ~ 99mmHg: ± 12%

20min. and up

0 ~ 38mmHg: ± 2mmHg 39 ~ 99mmHg: ± 5%

RR Measurement Range: 0 ~ 150bpm

0 ~ 40bpm: ± 1bpm 41 ~ 70bpm: ± 2bpm 71 ~ 100bpm: ± 3% 101 ~ 150bpm: ± 5% : 50 ± 7.5 ml/min

Flow Rate : 50 ± 7.5 ml/min System Response Time : 2.9 seconds (Typical) Delay Time : 2.7 seconds (Typical) Rise Time : 190 msec (maximum)

Telemetry (DS-7141, DS-7101LT)

Transmission Freq. : 608~614 MHz

Exact frequency depends on the destination.

RF Output Power : -15dBm Standard, 0dBm MAX

Channel Spacing : 12.5 kHz

Modulation Mode : Digital, Frequency Shift Keying (FSK)

This section lists selection, default setting, and backup status for each setup item. Backup Item
" ": Setup item will be retained even when the power is turned OFF.

- ": Setup item will be retained even when the power is turned OFF. When discharging procedure is performed, the value will be reset to initial setting. The alarm setup will be reset to initial setting with the selected alarm mode.
- " ": Setup item will be reset to initial setting when the power is turned OFF.

Patient Admit / Discharge

Itei	n	Selection	Default	Backup
Patient Name		Numeric, Alphabet, Symbol (16 characters)	Blank	
Sex		Male, Female	Undetermined	
Age		0 ~ 150 years or 0 ~ 999 days	0 year	
Birth Date		Birth Date (Year, Month, Day)	Blank	
ID		Numeric, Alphabet, Symbol (10 characters)	Blank	
Patient Type		Adult, Child, Neonate	Adult	
Pacemaker		Used, Not used	Not used	
Impedance Me	easurement	ON, OFF	ON	
Filter Mode		Monitor, ST Display, ESIS	Monitor	
	Bed ID	0~999	0	
Room/Bed ID	Room ID	Numeric, Alphabet, Symbol (4 characters)	BED -	

Alarm Setup

Item	Selection	Default	Backup
System Alarm	Suspend, ON	Suspend	
HR	ON, OFF 20 - 300bpm	ON 40 - 120	
AYSTOLE	ON, OFF 3 - 10 sec.	ON 5 sec.	
VF	ON, OFF	ON	
VT	ON, OFF	ON	
SLOW_VT	ON, OFF	ON	
RUN	ON, OFF 2 - 8 beats	ON 3 beats	
COUPLET	ON, OFF	OFF	
PAUSE	ON, OFF 1.5 - 5 sec.	OFF 2 sec.	
BIGEMINY	ON, OFF	OFF	
TRIGEMINY	ON, OFF	OFF	
FREQUENT	ON, OFF 1 - 50 beats / min.	OFF, 10 beats	
TACHY	ON, OFF 20 - 300	ON	/
BRADY	ON, OFF 20 - 300	ON	
	ON, OFF		
ST	ST1 ± 2.0 mV / ± 20 mm	OFF	
	ST2 ± 2.0mV / ± 20mm		
		ON	
554 (11)	011 055 0 000 11	SYS 80 - 180	
BP1 (mmHg)	ON, OFF 0 - 300mmHg	DIA OFF - OFF	
		MEAN OFF - OFF	
		ON	-
	011 055 0 40 015	SYS 10.0 - 24.0	
BP1 (kPa)	kPa) ON, OFF 0 - 40.0kPa	DIA OFF - OFF	
		MEAN OFF - OFF	

Item	Selection	Default	Backup
BP2 (mmHg)	ON, OFF 0 - 300mmHg	ON SYS OFF - OFF DIA OFF - OFF MEAN OFF - OFF	•
BP2 (kPa)	ON, OFF 0 - 40.0kPa	ON SYS OFF - OFF DIA OFF - OFF MEAN OFF - OFF	
RR	ON, OFF 5 - 150Bpm	ON 5 - 30	
APNEA	ON, OFF 5 - 20sec.	ON 15sec.	
SpO ₂	ON, OFF 50 - 100%	ON 90 - OFF	
NIBP (mmHg)	ON, OFF 10 - 300mmHg	ON SYS 80 - 180 DIA OFF - OFF MEAN OFF - OFF	/
NIBP (kPa)	ON, OFF 1.5 - 40.0kPa	ON SYS 10.0 - 24.0 DIA OFF - OFF MEAN OFF - OFF	
TEMP	ON, OFF 30 - 50?C	OFF OFF - OFF	
EtCO ₂ (mmHg)	ON, OFF 1 - 100mmHg	ON 30 - 45mmHg	
EtCO ₂ (kPa)	ON, OFF 0.1 - 13.3kPa	ON 4.0 - 6.0kPa	
EtCO ₂ (%)	ON, OFF 0.1 - 13.3%	ON 4.0 - 6.0%	
InspCO ₂ (mmHg)	ON, OFF 1 - 4mmHg	ON 3mmHg	
InspCO ₂ (kPa)	ON, OFF 0.1 - 0.4kPa	ON 0.4kPa	
InspCO ₂ (%)	ON, OFF 0.1 - 0.4%	ON 0.4%	

NOTE The alarm setup will be retained even after the power is turned OFF.

If discharging procedure is performed, the alarm setup will be initialized with the selected alarm mode.

Parameter Setup

	Item	Selection	Default	Backup
	Lead	, , , aVR, aVL, aVF, V	ECG1 Lead ECG2 Lead V	/
	Waveform Size	×1/4, ×1/2, ×1, ×2, ×4	ECG1 ×1 ECG2 ×1	
	Filter Selection	Monitor, ST Display, ESIS	Monitor	
	HR Source	Auto, ECG, SpO ₂ , BP	Auto	
ECG	Automatic Lead Switch	ON, OFF	OFF	
	Pacemaker Pulse	ON, OFF	ON	
	HR Average	ON, OFF, Instant, Average	Average	
	HR Sync. Indicator	ON, OFF	ON	
	Pace Pulse Mask Time	Auto, 10ms, 20ms, 40ms, OFF	Auto	
	ECG Drift Filter	ON, OFF	OFF	
	Waveform Size	×1/4, ×1/2, ×1, ×2, ×4	×1	
RESP	RR Sync. Indicator	ON, OFF	ON	
	CVA	ON, OFF	OFF	
	RR Source	Auto, Impedance, CO ₂	Auto	
	Impedance Meas.	ON, OFF	ON	

	Item	Selection	Default	Backup
	Waveform Size	×1/4, ×1/2, ×1, ×2, ×4	×1	-
SpO ₂	SpO ₂ SEC Alarm	OFF, 10, 25, 50, 100	OFF	
	Ignore NIBP	ON, OFF	ON	
	Auto Mode	ON, OFF	OFF	
	End Tone	ON, OFF	ON	
	Quick SYS List	ON, OFF	ON	
NIBP	PR	ON, OFF	OFF	
	Quick SYS	3, 5, 10 min.	10 min.	
	1-Min. Auto	10, 20 min.	20 min.	
	Mean	ON, OFF	OFF	
	Scale	20, 50, 75, 100, 150, 200, 250, 300mmHg	150mmHg	/
BP1		4, 8, 12, 16, 20, 24, 32, 40kPa	20kPa	
	Filter	6, 8, 12, 40Hz	12Hz	
	Mean	ON, OFF	OFF	
	Scale	20, 50, 75, 100, 150, 200, 250, 300mmHg	50mmHg	/
BP2	Scale	4, 8, 12, 16, 20, 24, 32, 40kPa	8kPa	/
	Filter	6, 8, 12, 40Hz	12Hz	
	Mean	ON, OFF	OFF	
	Meas. Unit	mmHg, kPa, %	mmHg	
	EtCO ₂ Average	10, 20, 30 sec., OFF	10 sec.	/
CO ₂	Scale	50, 100mmHg 4, 8, 10kPa 4, 8, 10%	50mmHg 4kPa 4%	/

Review Function Setup

	Item	Selection	Default	Backup
	Parameter	HR, PR, VPC, ST1/ST2, RR, APNEA, SpO ₂ , BP, NIBP, TEMP, EtCO ₂ / InspCO ₂ , EVENT1, EVENT2	HR	
	Duration	1, 2, 4, 8, 12, 24 hours	4 hours	
Graphic Trend	Scale	HR : 100, 200, 300bpm ST : ±0.2, ±0.5, ±1.0, ±2.0mV ±2, ±5, ±10, ±20mm VPC : 20, 50, 100 beats BP1 : 20, 50, 100, 150, 200, 300mmHg 4, 8, 16, 20, 24, 40kPa BP2 : 20, 50, 100, 150, 200, 300mmHg 4, 8, 16, 20, 24, 40kPa NIBP : 100, 150, 200, 300mmHg 16, 20, 24, 40kPa TEMP : 20-45, 30-40?C SpO ₂ : 0-100, 50-100, 80-100% RR : 100, 200, 300bpm APNEA : 15, 30 sec. CO ₂ : 50, 100mmHg 4, 8, 10kPa 4, 8, 10kPa 4, 8, 10%	HR : 100bpm ST : ±0.5mV	

Item		Selection	Default	Backup
Tabular Trend	Duration	1, 5, 10, 15, 30, 60 min.	60 min.	
OCRG	Display Time	4, 8 min.	8 min.	
OUNO	Waveform	Impedance Resp., CO ₂	Impedance Resp.	
	Parameter	ECG1, ECG2, BP, SpO ₂ , RESP, CO ₂	ECG1, ECG2	
	Alarm Factor	HR/PR/BPR : ON, OFF ST : ON, OFF RR : ON, OFF APNEA : ON, OFF SpO ₂ : ON, OFF BP1 : ON, OFF BP2 : ON, OFF NIBP : ON, OFF TEMP : ON, OFF CO ₂ : ON, OFF	HR/PR/BPR : ON ST : ON RR : ON APNEA : ON SpO ₂ : ON BP1 : ON BP2 : ON NIBP : ON TEMP : ON CO ₂ : ON	
Recall	Arrhythmia Factor	ASYSTOLE: ON, OFF VF : ON, OFF VT : ON, OFF SLOW_VT : ON, OFF RUN : ON, OFF COUPLET: ON, OFF PAUSE: ON, OFF BIGEMINY: ON, OFF TRIGEMINY: ON, OFF TRIGEMINY: ON, OFF FREQUENT: ON, OFF TACHY: ON, OFF BRADY: ON, OFF	ASYSTOLE : ON VF : ON VT : ON SLOW_VT : ON RUN : ON COUPLET : ON PAUSE : ON BIGEMINY : ON TRIGEMINY : ON FREQUENT : ON TACHY : ON BRADY : ON	
ST	Meas. Point	0 ~ 560ms	120ms	
Meas.	Ref. Point	0 ~ - 240ms	- 80ms	

NOTE

- The graphic trend data, tabular trend data will be retained even after the power is turned OFF.
 The ST data, OCRG data, recall data will be retained until 5 minutes after the power is turned OFF.

System Configuration Setup

Volume Setup

	Item	Selection	Default	Backup
	Pulse	16 levels	Level 8 from left	
	Key	16 levels	Level 10 from left	
Volume	Alarm	16 levels	Level 10 from left	
	Other Bed	16 levels	Level 10 from left	
	Others	16 levels	Level 8 from left	

Display Configuration

	Item	Selection	Default	Backup
	No. of waveforms	0~6 waveforms	3 waveforms	
	No. of numeric data	0 ~ 7 numeric data	4 numeric data	
Display Configuration	Displayed waveform	ECG1, ECG2, BP, SpO ₂ RESP, CO ₂	ECG1, SpO ₂ , RESP	/
Comiguration	Displayed numeric data	HR, BP, NIBP, SpO ₂ RESP, RR, CO ₂ , VPC/ST	HR, NIBP, SpO ₂ , RR	
	Enlarged Display	ON, OFF	OFF	
	Short Trend	ON, OFF, Overlap	OFF	

NOTE

By selecting ON for backup at discharge, the setup item will be stored even after discharge.
Selecting OFF will initialize the display configuration to the initial setting of the selected display mode.

System Configuration Menu

	Item	Selection	Default	Backup
Manual	Wave Select	ECG1, ECG2, BP, SpO ₂ RESP, CO ₂	ECG1, BP, RESP	
Recording	Rec. Duration	24 sec., Continuous	24 sec.	
	Delay Time	None, 8sec., 16 sec.	8 sec.	
	Alarm Record	ON, OFF, Center	OFF	
	Wave Select	ECG1, ECG2, BP, SpO ₂ RESP, CO ₂ , Alarm Factor	ECG1, Alarm Factor	
	Rec. Duration	12, 24 sec.	12 sec.	
Alarm	Alarm Factor	HR (HR / PR / BPR) Numeric Data, Arrhythmia	HR (HR / PR / BPR) Arrhythmia	
Recording	Arrhythmia Record	ASYSTOLE, VF, VT, SLOW_VT, RUN, COUPLET, PAUSE, BIGEMINY, TRIGEMINY, FREQUENT, TACHY, BRADY	ASYSTOLE, VF, VT, SLOW_VT, RUN, TACHY, BRADY	
	Periodic Record	ON, OFF, Center	OFF	
Periodic	Wave Select	ECG1, ECG2, BP, SpO _{2,} RESP, CO ₂	ECG1, BP, RESP	
Recording	Rec. Duration	6, 12, 24 sec.	12 sec.	
	Periodic Interval	5, 10, 15, 30, 60 min.	60 min.	
	Paper Feed to Top	ON, OFF	OFF	
Rec. Operation	Paper Feed to End	ON, OFF	ON	
Operation	QRS Classification	ON, OFF	ON	
Sweep	ECG, BP, SpO ₂	25, 12.5, 6.25mm/s	25mm/s	
Speed	RESP, CO ₂	25, 12.5, 6.25mm/s	6.25mm/s	
	Mode	Manual, Auto	Manual	
Night Mode	Start Time	00:00 ~ 23:59	22:00	
	Complete Time	00:00 ~ 23:59	7:00	
	Display	Time Disp. Only, Slightly Dark, Dark	Dark	
	Volume	No change, Quiet, Very quiet, Silence	Very quiet	
	Alarm Pole	ON, OFF	OFF	

	Item	Selection	Default	Backup
	ECG		Green	
	ST		Green	
	VPC		White	
	PACE		White	
	BP1		Red	
Color	BP2	32 colors	Cyan	
Color	NIBP		Cyan	
	SpO ₂		Yellow	
	TEMP		Orange	
	RESP		White	
	CO ₂		Cyan	
	ST2 (Trend)		Orange	Fixed
Alarma Catura	Alarm Suspend	1, 3, 5 min.	3 min.	
Alarm Setup	Alarm Silence	1, 3, 5 min.	3 min.	

Hospital Setup

1	ltem	Selection	Default	Backup
AC filter		50, 60Hz	According to Dip_SW	
Date		07/19, Jul.19, 19.Jul	Jul.19	
Alarm Mute		ON, OFF	OFF	
Home Key Fund	ction	Home / Enlarge, Home	Home	
Night Mode Ca	ncel	Any Key, Night Mode Key	Any Key	
Asystole,VF,VT	-	ON,ON/OFF	ON	
	BP	mmHg, kPa	mmHg	
Unit	TEMP	?C, ?F	?C	
	ST	mm, mV	mV	
Telemeter	Channel	0801~0879, 0900~0979, 1000~1079, 1100~1179, 1200~1279, 1300~1379	1100	
	Group	00 ~ 63	00	

Ward Setup

	Item	Selection	Default	Backup
Trend Clip		ON, OFF	ON	
BP Record S	cale	20, 40mm	40mm	
HR Low Limit	for VT	120, 140bpm	120bpm	
Password		ON, OFF	OFF	
Discharge Mo	ode	Admit, Cease	Admit	
Event Key		ON, OFF	ON	
Mean Calcula	ation	Waveform, Calculation	Waveform	
	Menu	All Key (excluding system config.)	All Key	
Key Mask	System Config.	All Key (excluding pre-set)	All Key	
	Pre-Set Menu	All Key (excluding ward setup)	All Key	

ı	tem	Selection	Default	Backup
User Key	Selection	Rec. START/STOP, Alarm Silence, Alarm, Key Lock, NIBP List, Graphic Trend, Tabular Trend, Recall, OCRG, Freeze, NIBP Auto Mode, Size/Lead, HR Source, BP Zero, Admit/Discharge, Night Mode, Display Config., Record, Tone/Volume, Other Bed, ST Display, Cease, Enlarged Display	Alarm Silence, Rec. START/STOP, Size/Lead, Admit/Discharge (from left of display)	
	Alarm Pole	ON, OFF	ON	
Alawa Dala	Alarm Level	Level 1, Level 1 and 2, Level 1, 2, and 3	Level 1	
Alarm Pole			Level 1 : Pattern 1	
	Pattern Setup	Pattern 1 ~ 10	Level 2 : Pattern 10	
			Level 3 : Pattern 4	
NIBP Data Eras	se Time	10, 30, 60min, 24hourr	60min	

Monitor Setup

Item		Selection	Default	Backup
Battery Opera	eration Normal, Power Save Normal		Normal	
Message Icon		ON, OFF	OFF	
Parameter Ke	y Frame	ON, OFF	ON	
Wide AC Filte	r	ON, OFF	OFF	
Check Discha	rge at Power ON	ver ON ON, OFF ON		
Backup at Dis	charge	ON, OFF	OFF	
	0 0	HR, RR	HR	
	Sync. Sygnal Output	Positive Logic, Negative Logic	Positive Logic	
Status		OFF, APNEA, Level 1,		
Output Setup		Level 1 and 2,	Level 1	
	Alarm Output	Level 1, 2 and 3		
		Positive Logic, Negative Logic, Pulse	Positive Logic	

Display Mode Setup

	Item	Default	Backup
Mode Select	tion	1	
	No. of Waveforms	3 Waveform	
	No. of Numeric Data	4 Numeric Data	
	Displayed Waveforms	ECG1, SpO ₂ , RESP	
Mode 1	Displayed Numeric Data	HR, NIBP, SpO ₂ , RR	
	Enlarged Display	OFF	
	Short Trend	OFF	
	Comment	CONFIG. 1	
	No. of Waveforms	3 Waveforms	
	No. of Numeric Data	4 Numeric Data	
	Displayed Waveforms	ECG1, SpO ₂ , RESP	
Mode 2	Displayed Numeric Data	HR, NIBP, SpO ₂ , RR	
	Enlarged Display	ON	
	Short Trend	OFF	
	Comment	CONFIG. 2	
	No. of Waveforms	4 Waveforms	
	No. of Numeric Data	6 Numeric Data	
	Displayed Waveforms	ECG1, BP1/2 (overlap), SpO ₂ , RESP	
Mode 3	Displayed Numeric Data	HR, BP1,BP2, NIBP, SpO ₂ , TEMP, RR	
	Enlarged Display	OFF	
	Short Trend	OFF	
	Comment	CONFIG. 3	
	No. of Waveforms	4 Waveforms	
	No. of Numeric Data	6 Numeric Data	
	Displayed Waveforms	Cascade, BP1/2 (overlap), SpO ₂ , RESP	
Mode 4	Displayed Numeric Data	HR, BP1, BP2, NIBP, SpO ₂ , TEMP, RR	
	Enlarged Display	OFF	
	Short Trend	OFF	
	Comment	CONFIG. 4	
	No. of Waveforms	6 Waveforms	
	No. of Numeric Data	7 Numeric Data	
	Displayed Waveforms	ECG1, BP1/2 (overlap),, SpO ₂ , CO ₂	
Mode 5	Displayed Numeric Data	HR, BP1, BP2, NIBP, SpO ₂ , TEMP/ RR, CO ₂	
	Enlarged Display	OFF	
	Short Trend	OFF	
	Comment	CONFIG. 5	

Alarm Mode Setup

	Item	Default	Backup
Alarm Mode		1	
/ Harri Mode	HR	ON 40 - 120	
	AYSTOLE	ON 5 sec.	
	VF	ON 5 SCC.	
	VT	ON	
		ON	
	SLOW_VT		
	RUN	ON 3 beats	
	COUPLET	OFF	
	PAUSE	OFF 2 sec.	
	BIGEMINY	OFF	
	TRIGEMINY	OFF	
	FREQUENT	OFF 10 beats	
	TACHY	ON	
	BRADY	ON	
	ST	OFF	
		ON	
	BP1 (mmHg)	SYS 80 - 180	
	Bi i (iiiiiiig)	DIA OFF - OFF	
		MEAN OFF - OFF	
	BP1 (kPa)	ON	
		SYS 10.0 - 24.0	
		DIA OFF - OFF	
		MEAN OFF - OFF	
Alarm Mode 1 ~ 5		OFF	
	BP2 (mmHg)	SYS OFF - OFF	
		DIA OFF - OFF	
		MEAN OFF - OFF	
		OFF	
	BP2 (kPa)	SYS OFF - OFF	
	Di 2 (Ki a)	DIA OFF - OFF	
		MEAN OFF - OFF	
	APNEA	ON 15 sec.	
	SpO ₂	ON 90 - OFF	
		ON	
	NIBP (mmHg)	SYS 80 - 180	
	(DIA OFF - OFF	
		MEAN OFF - OFF	
		ON	
	NIBP (%)	SYS 10.0 - 24.0	
	\\(\)	DIA OFF - OFF	
		MEAN OFF - OFF	
	EtCO ₂ (mmHg)	ON 30 45	
	EtCO ₂ (kPa)	ON 4.0 6.0	
	EtCO ₂ (%)	ON 4.0 6.0	
	InspCO ₂ (mmHg)	ON 3	
	InspCO ₂ (kPa)	ON 0.4	
	InspCO ₂ (%)	ON 0.4	
L	1113pco ₂ (/0)	ON 0.4	

This section explains the connector pin assignments.

Serial Connector Output Signal

No.	Signal Type	Description	Signal Level
1	RESET	Port Reset	TTL Hi Level Reset
2	RSV	Reserved	
3	TxD	Serial Transmit Data Output	RS232C
4	SG	Signal GND	
5	RxD	Serial Receive Data Input	RS232C
6	+5V	+5V	+ 5V power supply (150mA)
7	RSV	Reserved	
8	NC	No Connection	

Status I/O Signal

No.	Signal Type	Description	Signal Level
1	QRS SYNC	QRS SYNC Output	Logic TTL
2	ALM_OUT+	Alarm Output + (Isolation)	Photo MOS Relay Contact
3	RSV	Reserved	
4	RSV	Reserved	
5	RSV	Reserved	
6	RSV	Reserved	
7	+ 5V	+5V	+ 5V power supply (150mA)
8	ALM_OUT-	Alarm Output - (Isolation)	Photo MOS Relay Contact
9	GND	Power Supply Digital GND	

As the serial connector and status I/O connector uses the same isolation power supply, the total power supply capacity for +5V should be up to 200mA.

Chapter 12

Accessories

	Accessories 2
-	Optional Accessories - · · · · · · 3
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	Invasive Blood Pressure Measurement · · · · · · 3
	Non-Invasive Blood Pressure Measurement · · · · · 3
	Temperature Measurement · · · · · · 4
	SpO ₂ Measurement · · · · · · · · 4
	CO ₂ Concentration Measurement · · · · · · 4
	Others 1

12 Accessories

Accessories

This section lists the accessories for the DS-7100 system.

▲CAUTION

∠ Use only the accessories specified for this device. Otherwise, proper function cannot be executed.

For quality improvement, specifications are subject to change without prior notice.

Power Cable : CS-34 (3m)

ECG Lead Cable (5-lead) : #3380.0661.13

ECG Relay Cable (5-lead) : CI-700D-5

2ch BP Conversion Cable : CJ-7546

NIBP Air Hose : OA-7109B (3.5m)

Adult Cuff (Medium) : CUF-7102A

SpO₂ Interface Cable : DOC-10

Recording Paper : OP-124TE

This Operation Manual

- Optional Accessories -

The following products are available as optional accessories for the DS-7100 system. Purchase them as required.

▲CAUTION

- ∠ Use only the accessories specified for this device. Otherwise, proper function cannot be executed.
- For quality improvement, specifications are subject to change without prior notice.

ECG, Impedance Respiration Measurement

Item	Model Type	Q'ty	Note
ECG Lead Cable	3380.0648.13	1	3-electrode (hook type)
ECG Lead Cable	3380.0661.13	1	5-electrode (hook type)
ECG Lead Cable	500308800	1	4-electrode (hook type)
ECG Relay Cable	CI-700D-3	1	3-electrode (defibrillation-proof)
ECG Relay Cable	CI-700E-3	1	3-electrode (electrosurgery-proof)
ECG Relay Cable	CI-700D-4	1	4-electrode (defibrillation-proof)
ECG Relay Cable	CI-700E-4	1	4-electrode (electrosurgery-proof)
ECG Relay Cable	CI-700D-5	1	5-electrode (defibrillation-proof)
ECG Relay Cable	CI-700E-5	1	5-electrode (electrosurgery-proof)

ACAUTION

When using the electrosurgery-proof type ECG relay cable, respiration measurement can not be performed.

Invasive Blood Pressure Measurement

Item	Model Type	Q'ty	Note
Interface Cable (for Gambro)	CJ-369	1	
Interface Cable (for Becton Dichinson)	CJ-410	1	
2ch BP Conversion Cable	CJ-7546	1	

Non-Invasive Blood Pressure Measurement

Item	Model Type	Q'ty	Note
Adult Cuff (Large)	CUF-7101	1	
Adult Cuff (Medium)	CUF-7102A	1	
Adult Cuff (Small)	CUF-7103	1	
Pediatric Cuff	CUF-7104	1	
Infant Cuff	CUF-7105	1	
NIBP Air Hose (1.5m)	OA-7109A	1	
NIBP Air Hose (3.5m)	OA-7109B	1	
NIBP Extension Hose (1.5m)	OA-7110A	1	
NIBP Extension Hose (3.5m)	OA-7110B	1	
BP Conversion Socket	CUFJ-MO1	1	for connection to neonate cuff

Temperature Measurement

Item	Model Type	Q'ty	Note
Rectal Temperature Probe (for adult)	401J	1	
Rectal Temperature Probe (for pediatric)	402J	1	
Body Surface Temperature Probe	409J	1	
Probe Cover	70 14 616	10	

SpO₂ Measurement

Item	Model Type	Q'ty	Note
SpO ₂ DURASENSOR®	DS-100A	1	
SpO ₂ OXISENSOR®	D-25	1	24 per box
SpO ₂ OXISENSOR®	D-20	1	24 per box
SpO ₂ OXISENSOR®	I-20	1	24 per box
SpO ₂ OXISENSOR®	N-25	1	24 per box
SpO ₂ OXISENSOR®	R-15	1	24 per box
SpO ₂ Relay Cable	DOC-10	1	
MAX-FAST	MAX-FAST	1	24 per box

CO₂ Concentration Measurement

ltem	Model Type	Q'ty	Note
FilterLine H Set (Adult / Pediatric)	XS04624	1	with Nafion, adapter (25 per box)
FilterLine H Set (Infant / Neonate)	006324	1	with Nafion, adapter (25 per box)
Capno Line H (Adult)	008177	1	for nasal, with Nafion (25 per box)
Capno Line H (Pediatric)	008178	1	for nasal, with Nafion (25 per box)
CapnoLine H (Baby / Neonate)	008179	1	for nasal, with Nafion (25 per box)
CapnoLine H / O ₂ (Adult)	008180	1	for nasal, with Nafion, oxygen delivery (25 per box)
CapnoLine H / O ₂ (Pediatric)	008181	1	for nasal, with Nafion, oxygen delivery (25 per box)
Calibration Gas Kit (5% CO ₂)	GR08081	1	

Others

Item	Model Type	Q'ty	Note
Ground Cable	CE-12	1	
Battery Pack	T4UR18650F-2-4644	1	
DS-7100 Mount Kit	OA-451	1	
Recording Paper	OP-124TE	1	
Cleaning Cloth	OA-57	1	