# Chapter 3 Vital Application

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**3** Vital Application

# - To Acquire ECG Waveform -



### **Before Attaching the Electrodes**

1. Clean the electrode sites with an alcohol swab or other skin preparation. If necessary, shave the electrode sites to remove excessive hair.



2. Peel off the backing of disposable electrode.



Pay attention not to touch the electrode jelly.

### **Electrode Placement**

There are 3-electrode, 4-electrode, 5-electrode application depending on the cable type. Using the 4-electrode or 5-electrode application allows simultaneous monitoring of 2 ECG waveforms, and high accuracy of arrhythmia analysis can be attained. Also, the displayed lead type can be changed.

For 3	For 3-electrode lead (1 waveform monitoring)			
				RALA
	Symbol	Color	Electrode Site	White Black
	RA	White	On the right infraclavicular fossa	
	LA	Black	On the left infraclavicular fossa	
	LL	Red	On the left midclavicular line, near the supracrestal line.	Red
For 4	<b>4-electro</b> Lead Typ	d <b>e lead</b> ( e /	(Simultaneous 2 waveforms monitoring)	VF
For 4	<b>4-electro</b> Lead Typ	e /	(Simultaneous 2 waveforms monitoring) / / / _aVR / _aVL / _a	
For 4	<b>4-electro</b> Lead Typ <b>Symbol</b>	e /	(Simultaneous 2 waveforms monitoring) / / / _aVR / _aVL / _a\ Electrode Site	
For 4	<b>1-electro</b> Lead Typ Symbol RA	o <b>de lead</b> ( e / 	(Simultaneous 2 waveforms monitoring) / / / _aVR / _aVL / _a` Electrode Site On the right infraclavicular fossa	VF RA White Black
For 4	<b>4-electro</b> Lead Typ Symbol RA LA	o <b>de lead</b> ( e / 	(Simultaneous 2 waveforms monitoring) / / / _aVR / _aVL / _av Electrode Site On the right infraclavicular fossa On the left infraclavicular fossa	VF RA White Black
For 4	<b>4-electro</b> Lead Typ Symbol RA LA LL	i <b>de lead</b> ( ie / 	(Simultaneous 2 waveforms monitoring) / / / AVR / AVL / <i>Electrode Site</i> On the right infraclavicular fossa On the left infraclavicular fossa On the left midclavicular line, near the supracrestal line.	VF RA White Black Black

For 5	-electro Lead Typ	<b>de lead</b> ( e /	(Simultaneous 2 waveforms monitoring)	VF / V
	Symbol	Color	Electrode Site	a li a
	RA	White	On the right infraclavicular fossa	RA S Black
	LA	Black	On the left infraclavicular fossa	
	LL	Red	On the left midclavicular line, near the supracrestal line.	Brown
	RL	Green	On the right midclavicular line at the same height as F.	Jan 695
	V	Brown	Chest Lead (V1 ~ V6)	Green

#### **Connection to the Patient Monitor**

1. Connect the lead cable to the electrode.



2. Connect the lead cable to the relay cable.



3. Plug in the relay cable to the ECG input connector (green) of the patient monitor.



4. Verify that the ECG waveform is displayed on the monitor.



Adjust the waveform size and position. The monitoring lead can be also changed.





compares the waveform (QRS pattern) and RR interval for each heartbeat to determine the VPC. It compares the parameters such as QRS amplitude, QRS width, QRS polarity, RR interval, and selects abnormal QRS. Then the QRS with suspected VPC is pattern matched to distinguish the noise and VPC. This will finally determine the VPC and generate the arrhythmia alarm.

#### **QRS Classification**

Each heartbeat will be classified to the following patterns according to the QRS judgement.

N (Normal)	Normal QRS beat
V (VPC)	Ventricular Extrasystole
S (SVPC)	Supraventricular Extrasystole
P (Pacing Beat)	Pacing beat
F (Fusion Beat)	Fusion beat of pacing and spontaneous beat
? (Undetermined Beat)	Learning arrhythmia, or beat not matching the pattern

#### Arrhythmia Type

With the above QRS judgement, the following 12 types of arrhythmia alarm can be generated.

Туре	Meaning	Detection Criteria
ASYSTOLE	Cardiac Arrest	Cardiac arrest is detected for more than preprogrammed time.
VF	Ventricular Fibrillation	A random, rapid electrical activity of the heart is detected.
VT	Ventricular Tachycardia	9 or more continuous ventricular beats are detected. (HR: 140bpm / 120bpm or over)
SLOW_VT		9 or more continuous ventricular beats are detected. (HR: under 140bpm / 120bpm)
TACHY	Tachycardia	HR is over the upper alarm limit.
BRADY	Bradycardia	HR is below the lower alarm limit.
RUN	Consecutive VPC	Continuous VPC exceeding the preprogrammed value is detected.
COUPLET	Couplet Ventricular Extrasystole	2 continuous beats of VPC is detected.
PAUSE		Cardiac arrest of 1.5 seconds and over is detected.
BIGEMINY	Ventricular Bigeminy	QRS pattern of V-N-V-N-V-N is detected.
TRIGEMINY	Ventricular Trigeminy	QRS pattern of V-N-N-V-N-N is detected.
FREQUENT	Frequent VPC	VPC exceeding the preprogrammed value is detected within 1 minute.

#### **Filter Selection**

#### **Filter Mode Setup**

The waveform frequency characteristic can be selected from Monitor Mode, ESIS Mode, or ST Display Mode according to the monitoring purpose.

**1. Monitor Mode** Frequency Characteristic Adult / Pediatric : 0.5 ~ 40Hz Neonate : 1.6 ~ 40Hz This is the standard mode for ECG monitoring. The upper frequency is set to 40Hz to reduce artifact caused by EMG, etc.

**2. ESIS Mode** Frequency Characteristic Adult / Pediatric : 1.6 ~ 15Hz Neonate : 1.6 ~ 15Hz By selecting this mode when using electrosurgical instrument, electrical noise can be largely reduced. Do not select this mode unless using electrosurgical instrument.

**3. ST Display Mode** Frequency Characteristic Adult / Pediatric : 0.05 ~ 40Hz Select this mode if ST measurement is the main purpose of ECG monitoring.





Reference

Refer to " 6. Parameter Setup ECG" for details of filter mode.

#### **Procedure for Filter Mode Selection**

- 1. Press the ECG parameter key and display the ECG setup menu.
- 2. Press the Config. key.

Configuration 1/2		Page do	wn Prev. Disp.
Filter	Monitor	ESIS	ST Display
HR Average	Instant	Average	
HR sync Indicator	ON	OFF	
ECG Source	ECG	SpO₂	BP1
	Auto		

3. Select the filter mode from 3 selections.

#### **AC Filter**

If the ECG waveform is interfered with AC noise, the AC filter cuts off the frequency component (50Hz or 60Hz).

r is always set to ON
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Reference

Refer to "8. System Configuration Hospital Setup AC Filter" for AC filter setup (50Hz or 60Hz).

### Lead Cable Types

There are various combinations of lead cable connecting type and electrode material. Contact our service representative for details and select the appropriate electrode.

[for 3-electrode]	ECG Relay Cable (defibrillation-proof)	CI-700D-3
	ECG Relay Cable (electrosurgery-proof)	CI-700E-3
	ECG Lead Cable	3380.0648.13
[for 4-electrode]	ECG Relay Cable (defibrillation-proof)	CI-700D-4
	ECG Relay Cable (electrosurgery-proof)	CI-700E-4
[for 5-electrode]	ECG Relay Cable (defibrillation-proof)	CI-700D-5
	ECG Relay Cable (electrosurgery-proof)	CI-700E-5
	ECG Lead Cable	3380.0661.13

# - Respiration (Impedance Measurement)

HEDOOBOT FUKUDA DENSHI 60 T . . ý 116/ ( 92) 77 10 **23**/ (15) <sup>BP</sup> 129<sup>/3:43</sup> 82 92 38.2 11\_ A 30 Alarm silence Bec. Lead• Size Admit/ Discharge Home Menu

1. Verify that the ECG waveform is properly acquired.

The respiration waveform is detected from lead

-

of ECG mentioned in the previous section. Therefore if stable ECG is acquired, the respiration waveform can be acquired at the same time.

2. Verify that the respiration waveform and respiration rate is displayed on the home display.



Adjust the waveform size, baseline position and sweep speed.

Reference

Refer to "6. Parameter SetupRespiration" for scale / baseline setup.Refer to "8. System ConfigurationSweep Speed" for sweep speed setup.

## - To Measure the SpO<sub>2</sub> -

#### 1. Prepare an appropriate probe or sensor for the patient.



2. Connect the sensor to the patient monitor.



(1) Connect the SpO<sub>2</sub> relay cable (DOC-10) to the SpO<sub>2</sub> connector on the patient monitor.

(2) Insert the sensor into the SpO<sub>2</sub> relay cable connector, and lock with the transparent part.

3. Attach the sensor to the patient.



[Probe Type Sensor]



(1) Attach the probe as shown on left.The probe cable should be on the nail side.

(2) Adjust the sensor so that the light-emitting part (on cable side) touches the root of the nail, and close the probe.



(3) Press the probe lightly so that the finger and the rubber cover are appressed. This is to stabilize the probe, and to avoid ambient light.

#### [Single-use Type]

- (1) Clean the attachment site with alcohol, etc.
- (2) Align the light emitting element and light receiving element of the sensor with the measuring site in between when attaching the sensor to patient.



(3) Fixate the cable with surgical tape so that the sensor does not come off when a cable is pulled.



Attachment to the toe



Attachment to the finger

4. Verify that the SpO<sub>2</sub> is displayed.



Press the HOME key on the lower part of the display.

Verify that the  $SpO_2$  measurement and  $SpO_2$  waveform are displayed on the home display.

#### **Functional and Fractional Saturation**

The DS-7100 measures functional SpO<sub>2</sub> and may therefore produce measurements that differ from devices measuring fractional SpO<sub>2</sub>. "Functional" SpO<sub>2</sub> is the amount of oxygenated hemoglobin expressed as a percentage of the total amount of hemoglobin capable of transporting oxygen. By utilizing the light of two different wavelengths, the DS-7100 can analyze for both oxygenated and deoxygenated hemoglobin, and consequently, can determine the functional SpO<sub>2</sub>. The DS-7100 does not detect the presence of abnormal hemoglobin, such as carboxyhemoglobin or methemoglobin.

#### **Measured Versus Calculated Saturation**

When  $SpO_2$  is calculated from a blood gas measurement of the partial pressure of arterial oxygen (PaO<sub>2</sub>), the calculated value may differ from the DS-7100 SpO<sub>2</sub> measurement. This is because the calculated SpO<sub>2</sub> may not have been corrected for the effects of variables that shift the relationship between PaO<sub>2</sub> and SpO<sub>2</sub> : temperature, pH, the partial pressure of carbon dioxide(PaCO<sub>2</sub>), and the concentrations of 2, 3-DPG and fetal hemoglobin.



	<ul> <li>When measuring the SpO<sub>2</sub> of patient with high fever or peripheral circulatory insufficiency, check the sensor attachment periodically and change the attachment site. The temperature of attachment site will rise 2 ~ 3?C due to the sensor heat which may result in burn injury.</li> <li>For the following case, accurate measurement may not be possible.</li> </ul>
▲WARNING	<ul> <li>Patient with excessive abnormal hemoglobin (HbCO, MetHb)</li> <li>Patient with the pigment injected to the blood</li> <li>Patient receiving CPR treatment</li> <li>When a sensor is applied to a limb with NIBP cuff, arterial catheter, or intracatheter</li> </ul>
	When measuring at site with venous pulse
	<ul> <li>Patient with body motion</li> </ul>

	If irritation such as skin reddening or skin tit appears with the sensor
	use, change the attachment site or stop using the sensor.
	$\swarrow$ When fixating the sensor with a tape, do not wind the tape too strong.
	At the same time, check the blood flow constantly so that congestion
	is not generated at the peripheral.
	Even a short duration of attachment may inhibit the blood flow and
	generate compression necrosis and burn injury.
	Change the sensor attachment site constantly (every 4 hours). As
	the temperature of sensor attachment site normally rises $2 \sim 3$ °C.
<b><u>ZI</u>CAUTION</b>	compression necrosis and burn injury may generate
	$\propto$ As the skin of neonate / low birth weight infant is immature, change
	the concer attachment site more frequently depending on the
	Condition.
	Direct sunlight to the sensor area can cause a measurement error.
	Place a black or dark cloth over the sensor.
	Solution When not performing the measurement, unplug the relay cable and
	sensor from the SpO <sub>2</sub> connector. Otherwise, the measurement data
	may be erroneously displayed by the ambient light.