

IntelliVue MP2 Patient Monitor

Philips M8102A Technical data sheet

The IntelliVue MP2 portable patient monitor is compact in size, ergonomic, and modular in design. It provides an easy-to-use touchscreen user interface, is highly customizable and shares a technological platform with the Philips IntelliVue MP5-MP90 patient monitors.

The IntelliVue series offers a complete monitoring solution that is flexible and modular, designed to suit a broad spectrum of monitoring needs.

Measurement Features

- Compact, rugged, lightweight monitor with built in measurements.
- ECG monitoring using any combination of three to 10 electrodes.

- 12-lead ECG monitoring with five electrodes using the EASI method or with 10 electrodes using the conventional method.
- Multi-lead arrhythmia and ST segment analysis at the bedside on all available leads.
- Mainstream or Sidestream CO₂
- Choice of FAST SpO_2 or $Nellcor^{TM}$ $OxiMax^{TM}$ SpO_2^{-1}
- Invasive Pressure and Temperature measurement
- The monitor can operate using battery power for up to 3 hours with basic monitoring configuration to let you safely and easily monitor patients during



¹ The following are trademarks of Covidien AG and/or its affiliates: Nellcor $^{\text{TM}}$, Durasensor $^{\text{TM}}$, Dura-Y $^{\text{TM}}$, Oxiband $^{\text{TM}}$, OxiCliq $^{\text{TM}}$, OxiMax $^{\text{TM}}$, MAXFAST $^{\text{TM}}$.

in-hospital transfer. AC power is provided by an external power supply.

- Telemetry devices can be connected via short range radio to monitor telemetry data (ECG/SpO₂) on the MP2 screen (Telemetry as a parameter (TAAP)).
- IntelliVue Cableless Measurement Devices can be connected via short range radio to monitor data from the IntelliVue CL SpO₂ Pod or IntelliVue CL NBP Pod on the MP2 screen. The Cableless Measurement Devices can also be controlled from an assigned MP2 via short range radio.

Usability Features

- Touchscreen and hardkeys as input device.
- · Intuitive user interface.
- Simple menu hierarchy gives fast access to all basic monitoring tasks.
- Patient data management with tabular and graphic trends.
- Settings "Profiles" for rapid case turnover.
- Patented automatic alarm limits help clinicians provide care more efficiently.
- 3.5" TFT flat panel display with QVGA (320 x 240) resolution, wide viewing angle, large numerics, permanently visible alarm limits, and up to three real-time waves.
- Capable of functioning in a wireless infrastructure (IIT)

Intended Use

The monitor is intended to be used for monitoring and recording of, and to generate alarms for, multiple physiological parameters of adults, pediatrics, and neonates in a hospital environment and during patient transport inside and outside of hospitals. The MP2 when used with the TRx4841A/TRx4851A IntelliVue Telemetry System Transceiver is intended for use in a hospital environment and during patient transport inside the hospital environment.

The monitor is intended for use by health care professionals. The monitor is only for use on one patient at a time. It is not intended for home use. Not a therapeutic device.

Rx only: U.S. Federal Law restricts this device to sale by or on the order of a physician.

ST segment monitoring is intended for use with adult patients only and is not clinically validated for use with neonatal and pediatric patients. The ECG measurement is intended to be used for diagnostic recording of rhythm and detailed morphology of complex cardiac complexes (according to AAMI EC 11).

Hospital Environment:

The monitor is suitable for use in all medically used rooms which fulfill the requirements regarding electrical installation according to IEC60364-7-710 "Requirements for special installations or locations - Medical locations", or corresponding local regulations.

Upgradability

The MP2 monitor allows new capabilities to be added in the future as your monitoring requirements evolve. This upgradability gives the security of knowing that the monitors can be enhanced and updated as practices and technologies advance, and it protects long-term investments.

Main Components

Monitor

The monitor has a color LCD TFT display with a wide viewing angle, providing high resolution waveform and data presentation.

The display, processing unit and measurements are integrated into one device. An external power supply provides power for the monitor.

User Interface

The user interface is designed for fast and intuitive operation. The color graphical user interface ensures that clinicians quickly feel at ease using the monitor.

Configurable SmartKeys with intuitive icons allow monitoring tasks to be performed quickly and easily, directly on the monitor screen. Waves and numerics are color-coded.

The monitor displays up to three measurement waves simultaneously. For 12-lead ECG monitoring it can display 12 real-time ECG waves, with a rhythm strip and all ST values.

Flexible screen layout allows optimal use of the available display space, for example, waves can be overlapped or wave size can adjust dynamically depending on the number of waves configured for the space.

The MP2 monitor is supplied with a resistive touchscreen.

Simulated Keyboard

If alpha or numeric data entry is required, for example to enter patient demographics, an on-screen keyboard will automatically appear on the screen.

Mounting

The mounting options available enable flexible, space saving placement of the monitors for an ergonomic work space. The monitor is shipped with a low cost mounting plate if not specified otherwise.

Application Features

Critical and Cardiac Care Features

 The monitor performs multi-lead arrhythmia detection analysis on the patient's ECG waveform at the bedside. It analyzes for ventricular arrhythmias, calculates heart rate, and generates alarms, including asystole, bradycardia, and ventricular fibrillation.

- Up to 12 leads of ST segment analysis can be performed on adult patients at the bedside, measuring ST segment elevation and depression and generating alarms and events. The user can trend ST changes, set high and low alarm limits, and set both ST and isoelectric measurement points. Using ST Snippets, one-second wave segments can be compared with a baseline segment for each measured ST lead.
- optional ST Map application shows ST changes over time in two multi-axis spider diagrams.
- QT/QTc interval monitoring provides the measured QT interval, the calculated heart-rate corrected QTc value and a Δ QTc value, which tracks variation in the QT interval in relation to a baseline value.
- optional 12-lead ECG data can be measured, using either the EASI placement method with five standard electrodes or conventional electrode placement with 10 electrodes.¹
- 12 realtime ECG waveforms can be displayed simultaneously. Diagnostic 12-lead ECG can be captured, reviewed and stored on the patient monitor before it is sent to the Information Center. Local printout is available, in harmonized layout.
- High performance pulse oximetry technologies perform accurately even in cases with low perfusion.
- Choice of sidestream or mainstream CO₂ monitoring for high quality measurements with intubated and non-intubated patients.
- Telemetry devices (TRx4841A/TRx4851A TRx/TRx+ IntelliVue Transceiver) can be connected via short range radio to the MP2 to monitor telemetry data (ECG/SpO₂) on the MP2 screen.

Ease of Use

- Screen layouts are easily adjustable, allowing flexible display of measurement information.
- Temperature, height, and weight can be configured either in metric or imperial units. Pressure measurements can be displayed in kPa or mmHg. Gases can be displayed in kPa, mmHg.

Trends

- The trend database stores patient data from up to 16 measurement numerics. The measurement information can be sampled every
 seconds, one minute, or five minutes, and stored for a period ranging from four to 48 hours.
- Each NBP measurement generates a column in the Vital Signs trend table. The values for the other measurements are added to provide a complete vital signs set for the NBP measurement time.
- Horizon Trends show the deviation from a stored baseline.

EASI-derived 12-lead ECGs and their measurements are approximations to conventional 12-lead ECGs. As the 12-lead ECG derived with EASI is not exactly identical to the 12lead conventional ECG obtained from an electrocardiograph, it should not be used for diagnostic purposes.

Transport Features

- The monitor's portable design means it can be used for in and out-of-hospital transport: a basic monitor weighs 1.5 kg.
- The monitor can operate using battery power for up to 3 hours, to let you safely and easily monitor patients during procedures or inhospital transfer.
- Specially-designed mounting solutions let you quickly disconnect the monitor for transport and reconnect to the mount after transport.
- The Universal Admit, Discharge and Transfer (ADT) feature means that all ADT information is shared between the networked monitor and the Information Center. Information need only be entered once.

Patient Data Documentation

- An extensive range of Patient Reports can be printed:
 - 12-lead ECG Reports
- Alarm Limit Reports
- Vital Signs
- Graphic Trends
- Realtime Wave Reports

Report templates can be defined in advance, enabling print-outs tailored to each hospital's specific requirements to be started quickly. Reports can be printed on centrally-connected printers or via the IntelliVue PC Printing Solution, and they can be initiated manually or automatically at user-defined intervals.

 The IntelliVue PC Printing Solution allows printing of reports, waveform captures and trends from the MP2 to a standard off-the shelf printer or to an electronic file.

Alarms

The alarm system can be configured to present either the traditional HP/Agilent/Philips alarm sounds or sounds compliant with the IEC 60601-1-8 Standard.

Alarm limits are permanently visible on the main screen. The Alarm Limits page provides a graphic depiction of alarm limits in relation to the currently monitored measurement values and lets you adjust alarm limits. It also lets you preview wide and narrow automatic alarm limits before you apply them.

When an alarm limit is exceeded, it is signalled by the monitor in the following ways:

- an alarm tone sounds, graded according to severity
- an alarm message is shown on the screen, color-coded according to severity
- the numeric of the alarming measurement flashes on the screen
- alarm lamps flash for red and yellow alarms and are illuminated for technical INOPs

A "SmartAlarm Delay" algorithm helps to reduce the number of pulse oximetry nuisance alarms.²

If the monitor is connected via a network to a central monitoring station, alarming is simultaneous at the monitor and at the Information Center.

Alarms are graded and prioritized according to severity:

- Red Alarms*** identify a potentially life threatening situation for a
 patient.
- Yellow Alarms** indicate conditions violating preset vital signs limits.
- Technical Alarms (INOPS) are triggered by signal quality problems, equipment malfunction or equipment disconnect.
 The Silence/Pause Alarms function (equivalent to Silence/Suspend with previous monitor generations) allows the user to switch off alarm tones with one touch.

All alarms can be paused for a period of one, two, three, five, or 10 minutes or turned off indefinitely.

Alarm strip recordings are available on a centrally-connected recorder or via the IntelliVue PC Printing Solution.

Patented automatic alarm limits automatically adapt the alarm limits to the patient's currently measured vital signs within a safe margin defined individually for each patient.

Visual and/or audible latching and non-latching alarm handling is available.

Profiles

Profiles are predefined configuration settings for Screens, measurement settings, and monitor settings. Each Profile can be designed for a specific application area and patient category, for example OR adult, or ICU neonatal. Profiles enable a quick reaction to patient and care location changes: activating a Profile with a particular patient category (Adult, Pediatric or Neonatal) automatically applies suitable alarm and safety limits and saves time usually spent carrying out a complete set-up procedure.

Profiles can be created directly on the monitor or remotely on a personal computer and transferred to the monitor using the IntelliVue Support Tool. A selection of Profiles for common monitoring situations is provided with the monitor. These profiles can be changed, added to, renamed, or deleted.

Optional Networking Capabilities

The monitor can operate as part of a wired or wireless hospital network system, using the Philips IntelliVue Clinical Network interface. This includes:

- DHCP protocol support (as an alternative to BootP in certain network designs)
- 802.1x basic support on wireless networks
- $^2\,$ The "SmartAlarm Delay" is not available in the U.S.A. and territories relying on FDA Market clearance.
- The Smart Alarm Delay functionality is currently not available in China or in clinical environments under SFDA control.

- WMM on wireless networks
- QoS Tagging

Service Features

- The Support Tool helps technical personnel to
- carry out configuration, upgrades and troubleshooting via the network, or on an individual monitor
- share configuration settings between monitors
- back up the monitor settings.
- A password-protected Service Mode ensures that only trained staff can access service tests and tasks.
- The Configuration Mode is password-protected and allows trained users to customize the monitor configuration.

Device Connections

The monitor can be connected to:

- an Information Center (for example M3150B)
- a PC
- MMS Extensions (M3012A, M3014A, M3015A/B,)¹

Network Interface

The network interface provides the system with networking capability via a wired or wireless network connection.

Wireless Network

The monitor can function within a wireless infrastructure based on an IEEE 802.11 a/g network in the 2.4 GHz / 5 GHz bands (ISM). Additionally, the monitor can function within a telemetry infrastructure compatible with the Philips Cellular Telemetry System (CTS) in the WMTS and ISM bands. Additional components are required to complete the system. Please refer to the M3185A IntelliVue Clinical Network Technical Data Sheet for further information.

A short range radio interface for an IEEE 802.15.4 network in the 2.4 GHz (ISM) band is also available. This allows a telemetry device with a short range radio adapter or IntelliVue Cableless Measurement Devices to be assigned to the monitor.

Monitor Specifications

Safety Specifications

The monitor complies with the Medical Device Directive 93/42/EEC (CE₀₃₆₆) and with IEC 60601-1:1988 + A1:1991 + A2:1995; EN60601-1:1990 + A1:1993 + A2:1995; UL 60601-1:2003; CAN/CSA

¹ The MMS Extensions will only function when they are connected to the Philips Battery Extension, or the monitor is connected to external power.

C22.2#601.1-M90 + Suppl. No 1-94 + Am.2; JIS T 0601-1:1999; IEC 60601-1-1:2000; EN 60601-1-1:2001.

All applied parts are Type CF unless otherwise specified. They are protected against damage from defibrillation and electrosurgery. The possibility of hazards arising from software errors was minimized in compliance with ISO 14971:2000, EN60601-1-4:1996 + A1:1999 and IEC 60601-1-4:1996 + A1:1999.

The monitor complies with the EMC standards

IEC 60601-1-2:2001; EN 60601-1-2:2001

This ISM device complies with Canadian ICES-001. Cet appareil ISM est conforme à la norme NMB-001 du Canada.

The MP2 patient monitor with measurements and interfaces other than those listed below cannot be used for patient transport outside of the hospital environment.

The MP2 patient monitor_with the following measurements and interfaces:

- ECG/Respiration, NBP, SpO₂, Pressure, Temperature, CO₂¹
- LAN, Battery

can be used in a transport environment such as road ambulance, airplane or helicopter. For this purpose, the monitor fulfills the following additional mechanical, EMC and environmental requirements:

- **Shock Tests** according to IEC TR 60721-4-7, Class 7M3. Test procedure according to IEC/EN 60068-2-27 (peak acceleration up to 100g).
- *Random Vibration* according to IEC TR 60721-4-7, Class 7M3. Test procedure according to IEC/EN 60068-2-64 (RMS acceleration 5g).
- Sinusoidal Vibration according to IEC TR 60721-4-7, Class 7M3.
 Test procedure according to IEC/EN 60068-2-6 (acceleration up to amplitude 2g).
- Bump Test according to IEC/EN60068-2-29 (peak acceleration 15g, 1000 bumps).
- Free Fall Test according to EN1789 (covers also IEC TR 60721-4-7 and Class 7M3). Test procedure according to EN 60068-2-32 (height 0.75 m).
- Specification for degrees of protection provided by enclosures according to IEC/EN 60529: IP 32
- EN 1789 +A1:2003 Medical vehicles and their equipment Road ambulances (chapter 6 Medical Devices).
- EN13718-1 Air, water and difficult terrain ambulances. Medical devices interface requirements for the continuity of patient care. For Ambulances, Patient transport equipment, Emergency vehicles, Ambulance services, Rough-terrain vehicles, Water transport, Air transport, Medical equipment, Medical instruments, Interfaces, Performance.

- Radiated susceptibility 20 V/m according to EN ISO 9919 (SpO_2) and EN ISO 21647 (CO_2).
- **Altitude Range** from -500 to 3000 m operating and -500 to 4600 m storage and transportation.
- Extended radiated susceptibility tests

The MP2 patient monitor with its out-of-hospital parameter set provides a general immunity level of 20 V/m with only few restrictions. Details are as listed below:

- GSM 900: Immunity at 900 MHz (uplink mobile phone), 20 V/m, duty cycle 1:8
- GSM 1800: Immunity at 1800 MHz (uplink mobile phone), 20 V/m, duty cycle 1:8.
- DECT: Immunity at 1800 MHz (digital cordless phone), 20 V/m, duty cycle 1:24
- AM: 1 kHz Immunity from 80 MHz to 2.5 GHz (any radio communication unit, broadcasting and TV transmitter), 20 V/m, modulation factor 80%. (ECG: 20 V/m except 0.8-1.2 GHz where it is 10 V/m)
- Operating ambient temperature testing over the range from 0°C to 40°C (32°F to 104°F).
- Operating ambient humidity testing up to 95% RH at 40°C (104°F), non condensing.

US Army Airworthiness Certification Granted

- U.S. Army Airworthiness Certification and Evaluation (ACE) program of U.S. Army Aeromedical Research Laboratory (USAARL)
- Tests performed in accordance with the following standards:
- MIL-STD-461E Electromagnetic interference characteristics requirements and limits.
- MIL-STD-810F Department of Defense test method standard for environmental engineering considerations and laboratory tests.
- MIL-STD-1472F Human Engineering.
- ANSI/AAMI HE48-1993 HF Engineering guidelines & preferred practices for the design of medical devices.
- ANSI/AAMI ES1-1993 Safe current limits for electromedical apparatus.

¹ The MP2 Patient Monitor in combination with the M2741A LoFlo CO₂ Sensor meets CISPR 11, Group 1, Class A emission limits. Thus it is suitable only for use in all establishments other than domestic and those connected to a low power network which supplies buildings used for domestic purposes.

Physical Specifications

Product	Max Weight	WxHxD
M8102A	<1.25 kg	< 188 x 99 x 86 mm
IntelliVue MP2 (without	(2.8 lb)	$(7.4 \times 3.9 \times 3.4 \text{ in})$
handle and options)		

Environmental Specifications

Item	Condition	Range
Temperature	Operating	0°C to 40°C
Range		(32°F to 104°F)
	Storage	-20°C to 60°C
	(incl. Transport)	(-4°F to 140°F)
Temperature	Operating	0°C to 35°C
Range when		(32°F to 95°F)
charging the		
battery		
Humidity	Operating	15% to 95% Relative
Range		Humidity (RH) (non
		condensing)
	Storage and	5% to 95% Relative
	Transport	Humidity (RH)
Altitude	Operating	-500 m to 3000 m
Range		(10000 ft)
	Storage and	-500 m to 4600 m
	Transport	(15000 ft) ^a
Ingress	Monitor	IP32 (protected against
Protection		the ingress of solid
		foreign objects 2.5 mm
		in diameter or larger,
		and the ingress of water
		when the water is
		dripping vertically and
		the monitor is tilted up
	5 10	to 15°).
	External Power	IP31(protected against
	Supply (M8023A)	the ingress of solid
		foreign objects 2.5 mm
		in diameter or larger,
		and the ingress of water when the water is
		dripping vertically) when
		rested on its rubber feet
		on a flat, level surface.
		IP32 when mounted
		with the connectors
		facing downwards
		lacing downwards

Performance Specifications

Monitor Perform	mance Specifications		
Power	Power	<40 W average, <65 W	
Specifications	consumption	peak	
	Line Voltage	100 to 240 V ~	
	Current	1.3 to 0.7 A	
	Frequency	50/60 Hz	
Battery	Operating Time	Basic monitor	
S pecifications	(with new, fully	configuration: 3 hours	
	charged battery at		
	25°C)		
	Charge Time	When MP2 is off: 2 h	
		When MP2 is in use and	
		connected to the	
		external power: 12 h	
		approx.	
Indicators	Alarms Off	red LED	
	Alarms	red/yellow/cyan LED	
	On/Standby/Error	green/red LED	
	AC Power	green LED	
	Battery	yellow (charging)/red	
		blinking (empty) LED	
	External Power	green LED	
Sounds	Audible feedback for user input. Prompt tone.		
	QRS tones, or SpO ₂ modulation tone. Four		
	different alarm sounds	different alarm sounds.	

Trends:

12, or 16 numerics @ 12 sec, 1 minute, 5 minute resolution.

Multiple choices of number of numerics, resolution and duration depending on trend option and application area.

Alarm Signal	System delay	less than 3 seconds
	Pause duration	1,2,3 minutes or infinite,
		depending on
		configuration
	Extended alarm	5 or 10 minutes
	pause	
Review	Information: all alarm	s / inops, main alarms on/
Alarms	off, alarms acknowledged and time of	
	occurrence	
	capacity	500 items
Real Time	Range: from: January 1, 1997, 00:00 to:	
Clock	December 31, 2080, 23:59	
	Accuracy: < 4 seconds per day (typically)	
	Hold Time: infinite if powered by host monitor or external power supply; otherwise at least 48 hours	

Monitor Performance Specifications		
Buffered	Contents: Active settings, trends, patient data,	
Memory	realtime reports, review alarms	
	Hold Time: infinite if powered by external	
	power supply; otherwise at least 48 hours	
Restart time: After power interruption, an ECG wave will be shown		
on the display after 30 seconds maximum.		

M8023A External Power Supply Performance Specifications

M8023A External Power Supply Performance Specifications		
Power	Power	< 12 W average
S pecifications	Consumption	< 30 W peak
	Line Voltage	100 to 240 V ~
	Current	0.7 to 0.4 A
	Frequency	50/60 Hz ~
Indicators	AC Power	green LED

Interface Specifications

MP2 (M8102A)	Interface Spec	ifications
Measurement	Connectors	Female ODU (Proprietary)
Link (MSL)	Power	30 V to 60 V input
` '	Power Sync	RS-422 compliant input
		78.125 kHz (typical
	LAN signals	IEEE 802.3 10-Base-T
		complaint
	Serial signals	RS-422 compliant
	Local signals	Provided for connecting MMS
		extensions
ECG Sync	Cable	Yes
Pulse	Detection	
Output ^a	Marker In	No
·	Wave	No
	Output	
	Connector	Binder Series 709/719
	Output	Output low <0.8V @ I =
	Levels	-4 mA
		Output high >2.4 V @ I =
		4 mA
	Isolation	None
	Pulse Width	100 +/- 10 ms (high)
	Delay from	20 ms maximum per AAMI
	R-wave peak	EC13
	to start of	
	pulse	
	Minimum	0.5 V
	required R-	
	required K- wave	
802.11	wave	IEEE 802.11 a/b/g
802.11 Bedside	wave amplitude	IEEE 802.11 a/b/g
	wave amplitude Wireless	IEEE 802.11 a/b/g 2.4 GHz and
Bedside	wave amplitude Wireless Technology	·
Bedside	wave amplitude Wireless Technology Frequency	2.4 GHz and
Bedside Adapter	wave amplitude Wireless Technology Frequency Band	2.4 GHz and 5 GHz ISM
Bedside Adapter Internal	wave amplitude Wireless Technology Frequency Band	2.4 GHz and 5 GHz ISM compatible with Philips
Bedside Adapter Internal WMTS	wave amplitude Wireless Technology Frequency Band	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System
Bedside Adapter Internal WMTS Adapter (US	wave amplitude Wireless Technology Frequency Band Technology	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure
Bedside Adapter Internal WMTS Adapter (US	wave amplitude Wireless Technology Frequency Band Technology	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure WMTS, 1395-1400 MHz and
Bedside Adapter Internal WMTS Adapter (US only)	wave amplitude Wireless Technology Frequency Band Technology Frequency Band	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure WMTS, 1395-1400 MHz and 1427-1432 MHz
Bedside Adapter Internal WMTS Adapter (US only) Internal ISM	wave amplitude Wireless Technology Frequency Band Technology Frequency Band	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure WMTS, 1395-1400 MHz and 1427-1432 MHz compatible with Philips
Bedside Adapter Internal WMTS Adapter (US only) Internal ISM	wave amplitude Wireless Technology Frequency Band Technology Frequency Band	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure WMTS, 1395-1400 MHz and 1427-1432 MHz compatible with Philips Cellular Telemetry System
Bedside Adapter Internal WMTS Adapter (US only) Internal ISM	wave amplitude Wireless Technology Frequency Band Technology Frequency Band Technology	2.4 GHz and 5 GHz ISM compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure WMTS, 1395-1400 MHz and 1427-1432 MHz compatible with Philips Cellular Telemetry System (CTS) cellular infrastructure

MP2 (M8102A) Interface Specifications		
Short Range	Туре	Dual internal SRR Interface
Radio	Technology	IEEE 802.15.4
Interface ^b	Frequency	2.4 GHz ISM
	Band	(2,400 - 2,483 GHz)
	Modulation	DSSS (O-QPSK)
	Technique	
	Effective	max. 0 dBm (1 mW)
	Radiated	
	Power	

- a ECG Sync Pulse not available if ECG is sourced from the telemetry device b The short range radio interface is compatible with the IntelliVue Cableless Measurements and the following telemetry devices: TRx4841A/TRx4851A IntelliVue Telemetry System Transceiver.

M8023A Extern	al Power Supply Inte	erface Specifications
Measure-	Connectors	Male ODU (Proprietary)
ment Link	Power	48 V output
(MSL)	Power Sync.	RS-422 compliant
		output 78.125 kHz
		(typical)
	LAN signals	IEEE 802.3 10-Base-T
		compliant
	Serial signals	RS-422 compliant
		output
	Local signals	Not connected

Display Specifications		
Integrated	Sweep Speeds	6.25, 12.5, 25 and
QVGA		50 mm/s;
Display	Resolution	320 x 240
	Refresh frequency	60 Hz
	Useful screen	72 x 54 mm
		$(2.8 \times 2.1 \text{ in})$
	Pixel size	0.22 x 0.22 mm

MP2 (M8102A) Compatible Devices		
IntelliVue Instrument Telemetry Wireless Network		
(USA only)		
Internal	Technology	compatible with Philips
WMTS		Cellular Telemetry System
Adapter		(CTS), cellular infrastructure
	Frequency	WMTS, 1395-1400 MHz and
	Band	1427-1432 MHz
IntelliVue Instrument Telemetry Wireless Network		
(except USA)		
Internal ISM	Technology	compatible with Philips
Adapter		Cellular Telemetry System
·		(CTS), cellular infrastructure
	Frequency Band	2.4 GHz ISM

M4607A Battery Specifications		
Physical Specifications		
WxDxH	66 mm (2.36 in) x 80 mm (3.15 in) x	
	20 mm (0.79 in)	
Weight	160 g ±5%	
Performance Specification	tions	
Nominal Voltage	10.8 Volt	
Rated Capacity at	1000 mAh (typical)	
discharge C/5		
Environmental Specific	ations	
Temperature Range	Discharge 0°C to 60°C (32°F to 140°F)	
	Charge 0°C to 60°C (32°F to 140°F)	
	Storage and Transportation: -20°C to	
	65°C (-4°F to 149°F)	
Humidity Range	Operating: 15% to 95% Relative	
	Humidity (RH)	
	Storage and Transportation: 5% to 95%	
	Relative Humidity (RH)	
Battery Type	Lithium Ion Mangan, 10.8 V, 1000 mAh,	
Safety	complies with UL 2054 (UL	
	Recognized)	
Communication	complies with the SMBus specification	
Standard	v1.1	

M4605A Battery Specif	fications
Physical Specifications	
W×D×H	149 mm (5.866 in) x 89 mm (3.504 in) x 19.8 mm (0.78 in)
Weight	490 g (1.08 lb)
Performance Specificat	
Nominal Voltage	10.8 Volt
Rated Capacity at discharge C/5	6000 mAh (typical)
Continuous Discharge Capability	6.5 A
Environmental Specific	ations
Temperature Range	Discharge 0°C to 50°C (32°F to 122°F) Charge 0°C to 50°C (32°F to 122°F) Storage and Transportation: -20°C to 65°C (-4°F to 149°F)
Humidity Range	Operating: 15% to 95% Relative Humidity (RH) Storage and Transportation: 5% to 95% Relative Humidity (RH)
Battery Type	Smart Battery 10.8 V, 6000 mAH, Lithium Ion
Safety	complies with UL 2054 (UL Recognized)
Communication Standard	complies with the SMBus specification v1.1

Measurement Specifications

ECG/Arrhythmia/ST/QT

Complies with IEC 60601-2-25:1993 + A1:1999 /EN60601-2-25:1995 + A1:1999, IEC 60601-2-27:2005/EN60601-2-27:2006, IEC 60601-2-51:2003 /EN 60601-2-51:2003 and AAMI EC11/EC13:1991/2002.

ECG/Arrhythm	ia/ST Performance	Specifications	
Cardiotach	Range	Adult/pedi:	
	6-	15 to 300 bpm	
		Neo range:	
		15 to 350 bpm	
	Accuracy	±1% of range	
	Resolution	1 bpm	
	Sensitivity	≥200 µV _{peak}	
PVC Rate	Range	0 to 300 bpm	
	Resolution	1 bpm	
ST Numeric	Range	-20 to +20 mm	
	Accuracy	±0.5 mm or 15%,	
		whichever is greater	
	Resolution	0.1 mm	
QT Numeric	Range	200 to 800 ms	
	Accuracy	±30 ms	
	Resolution	8 ms	
QTc Numeric	Range	200 to 800 ms	
	Resolution	1 ms	
Δ Q Tc	Range	-600 to +600 ms	
Numeric	Resolution	1 ms	
QT-HR	Range - adult	15 to 300 bpm	
Numeric	Range - pediatric	15 to 350 bpm	
	and neonatal		
Sinus and SV	Brady	Adult: 15 to 60 bpm	
Rhythm		Pedi: 15 to 80 bpm	
Ranges		Neo: 15 to 90 bpm	
	Normal	Adult: 60 to 100 bpm	
		Pedi: 80 to 160 bpm	
		Neo: 90 to 180 bpm	
	Tachy	Adult: >100 bpm	
		Pedi: >160 bpm	
		Neo: >180 bpm	

ECG/Arrhythm	ia/ST Performance S	pecifications
Bandwidth	Diagnostic Mode	Adult/neo/pedi: 0.05 to 150 Hz
	Extended Monitoring Mode	Neo/pedi: 0.5 to 150 Hz
	Monitoring Mode	Adult: 0.5 to 40 Hz Neo/pedi: 0.5 to 55 Hz
	Filter Mode	Adult/neo/pedi: 0.5 to 20 Hz
Bandwidth when ECG is	Diagnostic Mode	Adult/neo/pedi: 0.05 to 40 Hz
transmitted from a	Extended Monitoring Mode	Neo/pedi: 0.5 to 40 Hz
telemetry device via	Monitoring Mode	Adult: 0.5 to 40 Hz Neo/pedi: 0.5 to 40 Hz
short range radio	Filter Mode	Adult/neo/pedi: 0.5 to 20 Hz
Differential Inp	ut Impedance	 >2 MΩ RA-LL leads (Resp) >5 MΩ at all other leads (at 10 Hz including patient cable)
Common Mode	Rejection Ratio	Diagnostic mode: >86 dB (with a 51 k Ω / 47 nF imbalance). Filter mode: >106 dB (with a 51 k Ω /47 nF imbalance).
Electrode Offse Tolerance	et Potential	±500 mV
Auxiliary Curre		Active electrode:
(Leads off Dete	cuon)	Reference electrode:
Input Signal Ra	nge	±5 mV

E001		
ECG/		
Arrhythmia/	Range	Adjustment
ST Alarm	, and the second se	
Specifications		
HR	15 to 300 bpm	Adult:1 bpm steps (15 to
	maximum delay:	40 bpm)
	10 seconds	5 bpm steps (40 to
	according to AAMI	300 bpm)
	EC 13-1992	Pedi/Neo:1 bpm steps
	standard	(15 to 50 bpm)
		5 bpm steps (50 to
		300 bpm)
Extreme	Difference to high	5 bpm steps
Tachy	limit 0 to 50 bpm	
	Clamping at 150 to	5 bpm steps
	300 bpm	
Extreme	Difference to low	5 bpm steps
Brady	limit 0 to 50 bpm	
	Clamping at 15 to	5 bpm steps
	100 bpm	
Run PVCs	2 PVCs	Not adjustable by user
PVCs Rate	1 to 99 PVCs/	1 PVC
	minute	
Vent Tach HR	20 to 300 bpm	5 bpm
Vent Tach	3 to 99 PVCs/	1 PVC
Run	minute	
Vent Rhythm	2 to 99 PVCs/	1 PVC
Run	minute	
SVT HR	120 to 300 bpm	5 bpm
SVT Run	3 to 99 SV beats	1 SV beat
ST High	-19.8 to +20 mm	0.2 mm
ST Low	-20 to +19.8 mm	0.2 mm
QTc High	200 ms to 800 ms	10 ms steps
∆QTc High	30 ms to 200 ms	10 ms steps
- 0		

ECG/Arrhythmia/ST Supplemental Information as required by AAMI EC11/13		
$\begin{tabular}{ll} \textbf{Respiration Excitation} & Sinusoidal signal, 260 μA, 40.5 kHz \\ \textbf{Waveform} & \end{tabular}$		
Noise Suppression	RL drive gain 44 dB max., max. voltage 1.8 Vrms	

		emental Information as
	AMI EC11/13	
Time to	Vent	Gain 0.5, Range 6.5 to 8.4 seconds,
Alarm for	Tachycardia	Average 7.2 seconds
Tachy-	1 mV _{pp} ,	Gain 1.0 Range 6.1 to 6.9 seconds,
cardia	206 bpm	Average 6.5 seconds
		Gain 2.0, Range 5.9 to 6.7 seconds,
		Average 6.3 seconds
	Vent	Gain 0.5, Range 5.4 to 6.2 seconds,
	Tachycardia	Average 5.8 seconds
	2 mV _{pp} ,	Gain 1.0, Range 5.7 to 6.5 seconds,
	195 bpm	Average 6.1 seconds
		Gain 2.0, Range 5.3 to 6.1 seconds,
		Average 5.7 seconds
Tall T-Wave F	Rejection	Exceeds ANSI/AAMI EC 13 Sect.
Capability		3.1.2.1(c)
		minimum recommended 1.2 mV T-
		Wave amplitude
Heart Rate A	veraging	Three different methods are used:
Method		Normally, heart rate is computed
		by averaging the 12 most recent
		RR intervals.
		For runs of PVCs, up to 8 RR
		intervals are averaged to compute
		the HR.
		If each of 3 consecutive RR
		intervals is greater than 1200 ms
		(that is, rate less than 50 bpm),
		then the 4 most recent RR
		intervals are averaged to compute
		the HR.
Response Tin	ne of Heart	HR change from 80 to 120 bpm:
Rate Meter to	o Change in	Range: [6.4 to 7.2 seconds]
Heart Rate	J	Average: 6.8 seconds
		HR change from 80 to 40 bpm:
		Range: [5.6 to 6.4 sec] Average:
		6.0 seconds
Heart Rate M	leter	Ventricular bigeminy: 80 bpm
Accuracy and	l Response	Slow alternating ventricular
to Irregular R	-	bigeminy: 60 bpm
	•	Rapid alternating ventricular
		bigeminy: 120 bpm
		Bidirectional systoles: 90 bpm
Accuracy of I	nput Signal	Methods A and D were used to
Reproduction		establish overall system error and
		frequency response.
		1 1 1 2 2 2

Respiration

Respiration Performance Specifications		
Respiration	Range Adult/pedi: 0 to 120 rpn	
Rate		Neo: 0 to 170 rpm
	Accuracy	at 0 to 120 rpm ±1 rpm
		at 120 to 170 rpm
		±2 rpm
	Resolution	1 rpm
Bandwidth		0.3 to 2.5 Hz (-6 dB)
Noise		Less than 25 m Ω (rms)
		referred to the input

Alarm Range Adjustment Delay Specifications High Adult/pedi: under max. 10 to 20 rpm: 14 seconds 100 rpm 1 rpm steps Neo: 30 to over 20 rpm: 150 rpm 5 rpm steps Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above 20 rpm: max.	Respiration			
High Adult/pedi: under max. 10 to 20 rpm: 14 seconds 100 rpm 1 rpm steps Neo: 30 to over 20 rpm: 150 rpm 5 rpm steps Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above	Alarm	Range	Adjustment	Delay
10 to 20 rpm: 14 seconds 100 rpm 1 rpm steps Neo: 30 to over 20 rpm: 150 rpm 5 rpm steps Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above	Specifications			
100 rpm 1 rpm steps Neo: 30 to over 20 rpm: 150 rpm 5 rpm steps Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above	High	Adult/pedi:	under	max.
Neo: 30 to over 20 rpm: 150 rpm 5 rpm steps Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: as. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above		10 to	20 rpm:	14 seconds
Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above		100 rpm	1 rpm steps	
Low Adult/pedi: 0 under for limits to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above		Neo: 30 to	over 20 rpm:	
to 95 rpm 20 rpm: from 0 to Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above		150 rpm	5 rpm steps	
Neo: 1 rpm steps 20 rpm: max. 0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above	Low	Adult/pedi: 0	under	for limits
0 to 145 rpm over 20 rpm: 4 seconds 5 rpm steps for limits above		to 95 rpm	20 rpm:	from 0 to
5 rpm steps for limits above		Neo:	1 rpm steps	20 rpm: max.
above		0 to 145 rpm	over 20 rpm:	4 seconds
			5 rpm steps	for limits
20 rpm: max.				above
				20 rpm: max.
14 seconds				14 seconds
Apnea Alarm 10 to 5 second	Apnea Alarm	10 to	5 second	
40 seconds steps		40 seconds	steps	

Philips FAST SpO₂

Complies with EN ISO 9919:2005 (except alarm system; alarm system complies with IEC 60601-2-49:2001).

Measurement Validation: The SpO_2 accuracy has been validated in human studies against arterial blood sample reference measured with a CO-oximeter. Pulse oximeter measurements are statistically distributed, only about two-thirds of the measurements can be expected to fall within the specified accuracy compared to CO-oximeter measurements. Display Update Period: Typical: 2 seconds, Maximum: 30 seconds. Max. with NBP INOP suppression on:

60 seconds. For ${\rm SpO_2}$ specifications of a connected telemetry device, see the specifications document of the telemetry device.

SpO ₂ Performa	nce Specifications	
SpO ₂ *	Range	0 to 100%
	Accuracy	Philips Reusable
		Sensors:
		M1191A, M1191AL,
		M1191B, M1191BL,
		M1192A: 2% (70% to
		100%)
		M1193A, M1194A,
		M1195A, M1196A:
		3% (70% to 100%)
		Philips Reusable
		Sensors with
		M1943A(L):
		M1191T, M1192T,
		M1193T (Adult),
		M1196T:
		3% (70% to 100%)
		M1193T (Neonate):
		4% (70% to 100%)
		Philips Disposable
		Sensors with
		M1943A(L):
		M1132A, M1133A,
		M1134A (adult/infant): 2%
		M1131A, M1133A,
		M1134A (neonate),
		M1901B, M1902B,
		M1903B, M1904B: 3%
		(70% to 100%)
		Nellcor™ Sensors
		with M1943A(L):
		MAXA, MAXAL, MAXP,
		MAXI, MAXN, D-25, D-
		20, I-20, N-25, OxiCliq
		A, P, I, N:
		3% (70% to 100%)

SpO ₂ Performa	nce Specifications	
SpO ₂ *	Accuracy	Masimo Reusable
· -		Sensors [®] with LNOP
		MP12 or LNC MP10:
		LNOP DCI, LNOP
		DCIP, LNOP YI, LNCS
		DCI, LNCS DCIP:
		2% (70% to 100%)
		LNOP TC-I, LNCS TC-I:
		3.5% (70% to 100%)
		Masimo Disposable
		Sensors® with LNOP
		MP12 or LNC MP10:
		LNOP Adt, LNOP Adtx,
		LNOP Pdt, LNOP Pdtx,
		LNOP Inf-L, LNCS
		Adtx, LNCS Pdtx, LNCS
		Inf-L: 2% (70% to 100%)
		LNOP Neo-L, LNOP
		NeoPt-L, LNCS Neo-L,
		LNCS NeoPt-L: 3%
		(70% to 100%)
	Resolution	1%
Pulse	Range	30 to 300 bpm
	Accuracy	±2% or 1 bpm,
		whichever is greater
	Resolution	1 bpm
Sensors		Wavelength range: 500
		to 1000 nm
		Emitted Light Energy:
		≤15 mW
		Information about the
		wavelength range can be
		especially useful to
		clinicians (for instance,
		when photodynamic
		therapy is performed)
Pulse Oximete	r Calibration Range	70 - 100%

*The specified accuracy is the root-mean-square (R	RMS) difference between the measured
values and the reference values	

SpO ₂ Alarm	Range	Adjustment	Delay
Specifications			
SpO ₂	Adult: 50% to 100% Pedi/Neo: 30 to 100%	1% steps	(0, 1, 2, 3, 30) + 4 seconds
Desat	Adult: 50% to Low alarm limit Pedi/Neo: 30% to Low alarm limit	1% steps	
Pulse	30 to	Adult:	max.
Tachycardia	300 bpm Difference to	1 bpm steps (30 to 40 bpm) 5 bpm steps (40 to 300 bpm) Pedi/Neo: 1 bpm steps (30 to 50 bpm) 5 bpm steps (50 to 300 bpm) 5 bpm steps	14 seconds max.
	high limit 0 to 50 bpm Clamping at	5 bpm steps	14 seconds
	150 to 300 bpm		
Bradycardia	Difference to low limit 0 to 50 bpm	5 bpm steps	max. 14 seconds
	Clamping at 30 to 100 bpm	5 bpm steps	

Nellcor OxiMax SpO₂ Specifications (M8102A #SP4)

Pulse Oximetry Perfor	mance Specifications
SpO ₂	
Measurement Range	1 to 100%
Resolution	1%
Accuracy	see table below
Low Perfusion	2% (70 - 100%)
Accuracy ^a	
Pulse	
Measurement Range	25 to 300 bpm
Resolution	1 bpm
Accuracy	+/- 3 bpm (25 to 250 bpm)
Low Perfusion	+/- 3 bpm (25 to 250 bpm)
Accuracy ^a	
Sensors	
Sensors	Wavelength range: 500 to 1000 nm
	Emitted Light Energy: ≤15 mW
	Information about the wavelength
	range can be especially useful to
	clinicians (for instance, when
	photodynamic therapy is performed)
Numeric Update Rate	,, ,
Numeric Update	typical 1 second, max <= 60 sec
Rate	y picar i second, max - oo see
nace	

a Specification applies to Monitor performance. Reading accuracy in the presence of low perfusion (detected IR pulse modulation amplitude 0.03% - 1.5%) was validated using signals supplied by a patient simulator. SpO₂ and pulse rate values were varied across the monitoring range over a range of weak signal conditions and compared to the known true saturation and pulse rate of the input signals.

The specified accuracy is the root-mean-square (RMS) difference between the measured values and the reference values.

SpO ₂ Accuracy Table			
	SaO ₂ Range: 70-100%		SaO ₂ Range: 60%-80%
Sensor	Adult/Infant	Neonate	Adult
M1901B ^a	Identical to OxiMax MAXN		
M1902B	Identical to OxiMax MAXI		
M1903B	Identical to OxiMax MAXP		
M1904B	Identical to OxiMax MAXA		
MAXA,	2%	NA	3%
MAXAL			
MAXN ^a	2%	2%	3%
MAXP	2%	NA	3%
MAXI	2%	NA	3%

SpO ₂ Accuracy Table			
	SaO ₂ Range 70-100%		SaO ₂ Range: 60%-80%
MAXFAST	2%	NA	3%
MAXR ^b	3.5%	NA	NA
SC-A	2%	NA	NA
SC-PR-I ^c	NA	2%	NA
SCNEO-I ^c	NA	2%	NA
OxiCliq A	2.5%	NA	NA
OxiCliq P	2.5%	NA	NA
OxiCliq N ^d	2.5%	3.5%	NA
OxiCliq I	2.5%	NA	NA
D-YS ^d	3%	4%	NA
D-YS & D- YSE	3.5%	NA	NA
D-YSPD	3.5%	NA	NA
DS-100A-1	3%	NA	NA
OXI-A/N ^d	3%	4%	NA
OXI-P/I	3%	NA	NA

a M1901B/MAXN:

Clinical functionality has been demonstrated on a population of hospitalized neonate patients. The observed SpO_2 accuracy was 2.5% in a study of 42 patients with ages of 1 to 23 days, weight from 750 to 4100 grams, and 63 observations made spanning a range of 85 to 99% SaO₂ while monitored with Nellcor OxiMax N-595 pulse oximeters.

- b The accuracy specification has been determined between saturations of 80%–100%. c SoftCare SC-PR-I, SCNEO-I:

Clinical functionality has been demonstrated on a population of hospitalized neonate and infant patients. The observed SpO₂ accuracy was 3.0% in a study of 57 patients with ages of 24 to 40 weeks, weight from 710 to 5,000 grams, and 185 observations made spanning a range of 63 to 100% SaO_2 while monitored with Nellcor OxiMax N-

595 pulse oximeters.
d Neonatal accuracy: When sensors are used on neonatal subjects as recommended, the specified accuracy range is increased by $\pm\,1$ digit, as compared to adult usage, to account for the theoretical effect on oximeter measurements of fetal hemoglobin in neonatal blood. For example, OxiCliq N accuracy on neonates is ± 3.5 digits, rather than ± 2.5.

NBPComplies with IEC 60601-2-30:1999/EN60601-2-30:2000.

Measure-ment Ranges Systolic Adult: 30 to 270 mmHg (4 to 36 kPa) Pedi: 30 to 180 mmHg (4 to 24 kPa) Neo: 30 to 130 mmHg (4 to 17 kPa) Diastolic Adult: 10 to 245 mmHg (1.5 to 32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to 34 kPa)
Pedi: 30 to 180 mmHg (4 to 24 kPa) Neo: 30 to 130 mmHg (4 to 17 kPa) Diastolic Adult: 10 to 245 mmHg (1.5 to 32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
24 kPa) Neo: 30 to 130 mmHg (4 to 17 kPa) Diastolic Adult: 10 to 245 mmHg (1.5 to 32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
17 kPa) Diastolic Adult: 10 to 245 mmHg (1.5 to 32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
Diastolic Adult: 10 to 245 mmHg (1.5 to 32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
32 kPa) Pedi: 10 to 150 mmHg (1.5 to 20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
20 kPa) Neo: 10 to 100 mmHg (1.5 to 13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
13 kPa) Mean Adult: 20 to 255 mmHg (2.5 to
- `
34 kPa)
,
Pedi: 20 to 160 mmHg (2.5 to 21 kPa)
Neo: 20 to 120 mmHg (2.5 to
16 kPa)
Pulse Rate Adult:40 to 300
Pedi: 40 to 300
Neo: 40 to 300
Accuracy Max. Std. Deviation: 8 mmHg (1.1 kPa)
Max. Mean Error: ±5 mmHg
(±0.7 kPa)
Pulse Rate Measurement 40 to 100 bpm: ± 5 bpm
Accuracy 101 to 200 bpm: ± 5% of
reading
201 to 300 bpm: ± 10% of
reading (average over NBP measurement cycle)
Heart Rate Range 40 to 300 bpm
Measurement Time Typical at HR > 60 bpm
Auto/manual: 30 seconds (adult)
25 seconds (neonatal)
Stat: 20 seconds
Maximum time: 180 seconds
(adult/pediatric) 90 seconds (neonates)
Cuff Inflation Time Typical for normal adult cuff:
Less than 10 seconds
Typical for neonatal cuff: Less
than 2 seconds

NBP Performance Specifications		
Initial Cuff Infla	ation	Adult: 165 ±15 mmHg
Pressure		Pedi: 130 ±15 mmHg
		Neo: 100 ±15 mmHg
Auto Mode Rep	petition	1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45,
Times		60 or 120 minutes
STAT Mode Cycle Time		5 minutes
Venipuncture Mode Inflation		1
Inflation	Adult	20 to 120 mmHg (3 to 16 kPa)
Pressure	Pediatric	20 to 80 mmHg (3 to 11 kPa)
	Neonatal	20 to 50 mmHg (3 to 7 kPa)
Automatic	Adult/	170 seconds
deflation	pediatric	
after	Neonatal	85 seconds

Measurement Validation: In adult and pediatric mode, the blood pressure measurements determined with this device comply with the American National Standard for Electronic or Automated Sphygmomanometers (ANSI/AAMI SP10 - 1992) in relation to mean error and standard deviation, when compared to intra-arterial or auscultatory measurements (depending on the configuration) in a representative patient population. For the auscultatory reference the 5th Korotkoff sound was used to determine the diastolic pressure. In neonatal mode, the blood pressure measurements determined with this device comply with the American National Standard for Electronic or Automated Sphygmomanometers (ANSI/AAMI SP10 - 1992 and AAMI/ANSI SP10A -1996) in relation to mean error and standard

deviation, when compared to intra-arterial measurements in a representative patient population.

NBP Alarm Specifications	Range	Adjustment
Systolic	Adult: 30 to 27 mmHg	10 to 30 mmHg:
	(4 to 36 kPa)	2 mmHg (0.5 kPa)
	Pedi: 30 to 180 mmHg	> 30 mmHg:
	(4 to 24 kPa)	5 mmHg (1 kPa)
	Neo: 30 to 130 mmHg	
	(4 to 17 kPa)	
Diastolic	Adult: 10 to 245 mmHg	
	(1.5 to 32 kPa)	
	Pedi: 10 to 150 mmHg	
	(1.5 to 20 kPa)	
	Neo: 10 to 100 mmHg	
	(1.5 to 13 kPa)	
Mean	Adult: 20 to 255 mmHg	
	(2.5 to 34 kPa)	
	Pedi: 20 to 160 mmHg	
	(2.5 to 21 kPa)	
	Neo: 20 to 120 mmHg	
	(2.5 to 16 kPa)	

	Transducer		Load Impedance: 200 to
			2000 Ω (resistive)
djustment			Output Impedance:
			\leq 3000 Ω (resistive)
to 30 mmHg:	Frequency Resp	oonse	dc to 12.5 Hz or 40 Hz
mmHg (0.5 kPa)	Zero	Range	±200 mmHg (±26 kPa)
30 mmHg:	Adjustment	Accuracy	±1 mmHg (±0.1 kPa)
nmHg (1 kPa)		Drift	Less than 0.1 mmHg/°C
			(0.013 kPa/°C)
	Gain	Accuracy	±1%
	Accuracy	Drift	Less than 0.05%/°C
		Non linearity	Error of \leq 0.4% FS (@CAL
		and Hysteresis	200 mmHg)
	Overall	(including	± 4% of reading or ±
	Accuracy	transducer)	4 mmHg
			(± 0.5 kPa), whichever is
			greater
	Volume displac	ement of	0.1 mm ³ /100 mmHg
	CPJ840J6		

Invasive Pressure Performance Specifications

NBP Overpressure Settings		
Adult	> 300 mmHg (40 kPa)	not user adjustable
	> 2 sec	
Pedi	> 300 mmHg (40 kPa)	
	> 2 sec	
Neo	> 150 mmHg (20 kPa)	
	> 2 sec	

Invasive Pressure and Pulse

Complies with IEC 60601-2-34:2000/EN60601-2-34:2000.

Invasive Pressure Performance Specifications		
Measurement Range		-40 to 360 mmHg
Pulse Rate	Range	25 to 350 bpm
	Accuracy	±1% Full Range
	Resolution	1 bpm
Input Sensitivi	ty	Sensitivity: 5 μ V/V/mmHg
		(37.5 μV/V/kPa)
		Adjustment range: ±10%

Invasive Pressure Alarm Specifications	Range	Adjustment	Delay
Pressure	-40 to 360 mmHg (-5.0 to 48 kPa)	-40 to 30 mmHg 2 mmHg (0.5 kPa) > 30 mmHg 5 mmHg (1 kPa)	max. 12 seconds
Extreme High	Difference to high limit 0 to 25 mmHg Clamping at -40 to 360 mmHg	5 mmHg steps (0.5 kPa) 5 mmHg steps (1.0 kPa)	
Extreme Low	Difference to low limit 0 to 25 mmHg Clamping at 40 to 360 mmHg	5 mmHg steps (0.5 kPa) 5 mmHg steps (1.0 kPa)	

Invasive			
Pressure	Pango	Adjustment	Dolov
Alarm	Range	Adjustment	Delay
S pecifications			
Pulse	25 to	Adult:	
	300 bpm	1 bpm steps	
		(25 to	
		40 bpm)	
		5 bpm steps	
		(40 to	
		300 bpm)	
		Pedi/Neo:	
		1 bpm steps	
		(25 to	
		50 bpm)	
		5 bpm steps	
		(50 to	
		300 bpm)	
Tachycardia	Difference to	5 bpm steps	max.
	high limit 0 to		14 seconds
	50 bpm		
	Clamping at	5 bpm steps	
	150 to		
	300 bpm		
Bradycardia	Difference to	5 bpm steps	max.
	low limit 0 to		14 seconds
	50 bpm		
	Clamping at	5 bpm steps	
	25 to		
	100 bpm		

Temp

Complies with EN 12470-4:2000

Temp Performance Specifications		
Range	-1°C to 45 °C	
	(30°F to 113 °F)	
Resolution	0.1 °C (0.2 °F)	
Accuracy	±0.1 °C (±0.2 °F)	
Constant	Less than 10 seconds	
Range	-1°C to 45 °C (30°F to 113 °F)	
Adjustment	-1 to 35 °C (30 to 95 °F):	
	0.5 °C (1.0 °F) steps	
	35 to 45 °C (95 to 113 °F):	
	0.1 °C (0.2 °F) steps	
	Range Resolution Accuracy Constant Range	

Temp Alarm Specifications	Range	Adjustment
Temp High/	-1°C to 45°C (30°F	-1°C to 35°C (30°F to
Low Alarms	to 113°F)	95°F),
		0.5°C (1.0°F) steps
		35°C to 45°C (95°F to
		113°F), 0.1°C (0.2°F)
		steps

CO_2

The ${\rm CO_2}$ measurement in the monitor, M3014A and M3015A/B complies with EN ISO 21647:2004 + Cor.1:2005 (except alarm system; alarm system complies with IEC 60601-2-49:2001).

M3015A/B N	1icrostream (CO ₂ Performance Specifications
CO ₂	Range	0 to 98 mmHg (0 to 13 kPa), or 13%
		CO ₂ , whichever is lower
	Accuracy	Up to 5 minutes during warm-up:
		±4 mmHg or 12%, whichever is
		greater
		After 5 minutes warm-up:
		0 to 40 mmHg (0 to 5.3 kPa):
		±2.2 mmHg (±0.3 kPa)
		Above 40 mmHg (5.3 kPa): ±(5.5% +
		(0.08%/mmHg above 40 mmHg)) of
		reading
		These specifications are valid for
		21% O ₂ and N ₂ balance, up to 35°C
		ambient temperature, up to 60 rpm
		in adult mode and 100 rpm in
		neonatal mode. Outside of these
		conditions the accuracy reaches at a
		minimum ± 4 mmHg or $\pm 12\%$ of the
		reading, whichever is greater.
	Resolution	Numeric: 1.0 mmHg (0.1 kPa)
		Wave: 0.1 mmHg (0.01 kPa)
	Stability	Included in Accuracy specifications
awRR	Range	0 to 150 rpm
	Accuracy	0 to 40 rpm: ±1 rpm
		41 to 70 rpm: ±2 rpm
		71 to 100 rpm: ±3 rpm
		>100 rpm: ±5% of reading
Warm-up T	ime	5 minutes for full accuracy
		specification

M3015A/B Microstream (CO ₂ Performance Specifications
Rise Time	190 ms for neonatal mode
	(measured with FilterLine H for
	neonatal)
	240 ms for adult mode
	(measured with FilterLine H for
	adult)
Sample Flow Rate	50 + 15/-7.5 ml/minute
Gas Sampling Delay	Typical:2.3 seconds
Time	Maximum:3 seconds
Sound Pressure	Acoustic noise: <45 dBA
Total System Response	The total system response time is
Time	the sum of the delay time and the
	rise time.

M3014A Ma	instream CO ₂ Pe	rformance Specifications
CO ₂	Range	0 to 150 mmHg (0 to 20.0 kPa)
4	Accuracy	after 2 minutes warmup:
	ŕ	For values between 0 and
		40 mmHg: ±2.0 mmHg
		(±0.29 kPa)
		For values from 41 to 70 mmHg:
		±5% of reading
		For values from 71 to
		100 mmHg: ±8% of reading
		The specifications are valid for
		standard gas mixtures, balance
		air, fully hydrated at 35°C, P _{abs} =
		760 mmHg, flow rate = 2 l/min.
	Resolution	Numeric: 1.0 mmHg (0.1 kPa)
		Wave: 0.1 mmHg (0.01 kPa)
	Stability:	
	Short term drift	±0.8 mmHg over four hours
	Long term drift	Accuracy specification will be
		maintained over a 120 hour
		period
awRR	Range	2 to 150 rpm
	Accuracy	±1 rpm
Warm-up T	ime	2 minutes with CO ₂ transducer
		attached for full accuracy
		specification
Response Ti	me	Less than 60 ms (with adult or
		infant reusable or disposable
		adapter)

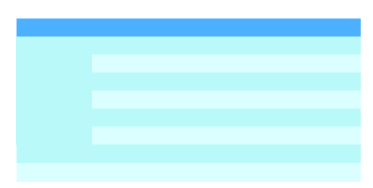
M3014A Sid	estream CO ₂ Per	formance Specifications
CO ₂	Range	0 to 150 mmHg (0 to 20.0 kPa)
	Accuracy	after 2 minutes warmup:
		For values between 0 and
		40 mmHg: ±2.0 mmHg
		(±0.29 kPa)
		For values from 41 to 70 mmHg:
		±5% of reading
		For values from 71 to
		100 mmHg: ±8% of reading
		For values from 101 to
		150 mmHg: ±10% of reading
		At respiration rates above
		80 rpm, all ranges are ±12% of
		actual. The specifications are
		valid for gas mixtures of CO_2 ,
		balance N ₂ , dry gas at
		760 mmHg within specified
		operating temperature range.
	Resolution	Numeric: 1.0 mmHg (0.1 kPa)
		Wave: 0.1 mmHg (0.01 kPa)
	Stability:	
	Short term drift	±0.8 mmHg over four hours
	Long term drift	Accuracy specification will be
		maintained over a 120 hour
		period
awRR	Range	2 to 150 rpm
	Accuracy	±1 rpm
Warm-up T	ime	2 minutes with CO ₂ sensor
		attached for full accuracy
		specification
Sample Flov	v Rate	50 ±10 ml/minute
Total System	n Response	3 seconds
Time		
Operating T	emperature	0°C to 40°C (32°F to 104°F)

M3014A Mainstream and Sidestream CO_2 Humidity Correction Factor

Either BTPS or STPD can be selected as the humidity correction factor for the ${\rm CO_2}$ readings. The formula for the correction calculation is:

$$P_{STPD} = P_{BTPS} \cdot \frac{P_{abs}}{P_{abs} - P_{H2O}}$$

Where p = partial pressure, P_{abs} = absolute pressure, and P_{H2O} = 42 mmHg @35°C and 100% RH.



Mainstream CO₂ Humidity Correction Factor

Either BTPS or STPD can be selected as the humidity correction factor for the Mainstream ${\rm CO}_2$ readings. The formula for the correction calculation is:

$$P_{STPD} = P_{BTPS} \cdot \frac{P_{abs}}{P_{abs} - P_{H2O}}$$

Where p = partial pressure, P_{abs} = absolute pressure, and P_{H2O} = 47 mmHg @37°C and 100% RH.

CO ₂ Alarm Specifications	Range	Adjustment	Delay
etCO ₂ High	20 to 95 mmHg (2 to 13 kPa)	1 mmHg (0.1 kPa)	M3014A: less than 14 seconds
etCO ₂ Low	10 to 90 mmHg (1 to 12 kPa)		M3015A/B: less than18 secon ds.
imCO ₂ High	2 to 20 mmHg (0.3 to 3.0 kPa)	steps of 1 mmHg (0.1 kPa)	M3014A: less than 14 seconds M3015A/B: less than18 secon ds.

CO ₂ Alarm Specifications	Range	Adjustment	Delay
awRR High	Adult/pedi: 10 to 100 rpm Neo: 30 to 150 rpm	under 20 rpm: 1 rpm steps over 20 rpm: 5 rpm steps	M3014A: less than 14 seconds M3015A/B: less than18 secon ds.
awRR Low	Adult/pedi: 0 to 95 rpm Neo: 0 to 145 rpm		M3015A/B: settings <20 rpm: less than 8 seconds >20 rpm: less than 18 seconds M3014Asetti ngs <20 rpm: less than 4 seconds >20 rpm: less than 14 seconds
Apnea delay	10 to 40 seconds	5 second steps	set apnea delay time + 4 seconds (M3014A) or 8 seconds (M3015A/B)

Ordering Information

Ordering information for the M8102A patient monitor is given here.

Parameters	M8102A
Order one Bxx option	
ECG, Resp, NBP, SpO ₂	B20
ECG, Resp, NBP, SpO ₂ , Press/Temp	B22
ECG, Resp, NBP, SpO ₂ , CO ₂	B23

Application OptionsXDS Connectivity

Application Options	M8102A
Full Arrhythmia Capability	C01
12-Lead ECG Application (conventional)	C12
ST Map	C13
Full Networking	C15

Options	M8102A
4-Wave XDS Connectivity	X04
6-Wave XDS Connectivity	X06 ^a
XDS Remote Control	X20
XDS Clinical Workstation	X30
a BU approval required	

SpO₂ Technology Choice

Options	M8102A
Philips FAST SpO ₂	Standard
Substitute Philips FAST SpO ₂ with Nellcor	SP4
OxiMax SpO ₂	

Hardware Options

Hardware Add-Ons	M8102A
Anti-slip pad	E18
Carrying strap	E19
MMS Mount	E20
Protective cover	E23
Add 1X Lithium-Ion battery	E24
Add 2X Lithium-Ion battery	E26
SN3 ECG Sync Cable	SN3

Interface Options

Interfaces	M8102A
IntelliVue 802.11 Bedside Adapter	J35 ^a
Instrument Telemetry 1.4 GHz	J45 ^a
Instrument Telemetry 2.4 GHz	J47 ^a
Short Range Radio Interface	J46 ^a

a May not be available in all geographies

Upgrade Options M8102AU

Options	MP2 M8102AU
Application	
Full Arrhythmia capability	<u>C01</u>
conv. 12-lead ECG	C12
Full Networking	<u>C15</u>
XDS external display solution	
4-wave XDS connectivity	X04
XDS Remote Control	X20
XDS Clinical Workstation	X30
Interfaces	
802.11 Wireless Interface	J35 ^a
Instrument Telemetry 1.4 GHz	J45 ^a
Short Range Radio	J46 ^a
Instrument Telemetry 2.4 GHz	J47 ^a
Software Upgrade	
Current Software Revision	SU <u>J</u> O

a May not be available in all geographies

Sensors and Disposables

Accessory	M8102A
3-lead Accessories Bundle ICU-AAMI	G06
Tyco low cost cable	
3-lead Accessories Bundle ICU-IEC	G07
Tyco low cost cable	
5-lead Accessories Bundle ICU-AAMI	G08
Tyco low cost cable	
5-lead Accessories Bundle ICU-IEC	G09
Tyco low cost cable	
5-lead Accessories Bundle ICU-AAMI	H06
5-lead Accessories Bundle ICU-IEC	H07
5-lead Accessories Bundle OR-AAMI	H08
5-lead Accessories Bundle OR-IEC	H09
Accessories Bundle Neonatal-AAMI	H14
Accessories Bundle Neonatal-IEC	H15
3-lead Accessories Bundle ICU-AAMI	H16
3-lead Accessories Bundle ICU-IEC	H17
3-lead Accessories Bundle OR-AAMI	H18
3-lead Accessories Bundle OR-IEC	H19
CO ₂ Mainstream Sensor	N01
Reusable Adult Airway Adapter (msCO ₂)	N02
Reusable Infant Airway Adapter (msCO ₂)	N03
Single Use Adult Airway Adapter (msCO ₂)	N04
Single Use Infant Airway Adapter (msCO ₂)	N05
CO ₂ Sidestream Sensor	N11
Non-intubated Adult Airway Adapter (ssCO ₂)	N12
Non-intubated pediatric Airway Adapter	N13
(ssCO ₂)	
Intubated Adult Airway Adapter (ssCO ₂)	N14
Intubated Pediatric Airway Adapter (ssCO ₂)	N15

Related Products

M3086A Support Tool

Mounting Information

The Intellivue MP2 Roll Stand Mounting Kit (Order No. 989803153021) is compatible with the table top mount and the standard mounting plate. For information on other mounting hardware, contact your local Philips sales representative. For GCX mounting hardware information, see www.gcx.com/philips.

Documentation

All documentation is available in .pdf format on documentation CD-ROM. Additionally, a printed copy of the Instructions for Use and Quick Guide ships with each monitor.

- Instructions for Use (printed)
- Quick Guide (printed)
- Installation and Service Guide
- Configuration Guide
- Documentation CD-ROM
- Training Guide (printed)
- Computer Based Training (optional)

Carry Case & NVG Display Filter

Accessory	Part No.
MP2 Carry Case Std Red	989803163331
MP2 Carry Case Std Blk	989803163341
MP2 Carry Case Mini Red	989803163351
MP2 Carry Case Mini Blk	989803163361
MP2 Carry Case Large Blk	989803163371
MP2 Carry Case Replacement Kit	989803163631
MP2 NVG Display Filter	989803163381

Battery Extension

The Philips Battery Extension provides additional battery runtime of up to 6 hours for intra-hospital patient transport and concurrent ${\rm CO}_2$ measurement with additional invasive blood pressure and temperature measurement.

Accessory	Part No.
Philips Battery Extension	865297

ECG Accessories



This symbol indicates that the cables and accessories are designed to have special protection against electric shocks (particularly regarding allowable leakage currents), and are

defibrillator proof.

Trunk Cables

	3-Electrode Cable Set	5-Electrode Cable Set	6-Electrode Cable Set	10-Electrode Cable set (5+5)	10-Electrode Cable set (6+4)
Part No.	M1669A	M1668A	M1667A	M1663A	M1665A
Length	2.7 m	2.7 m	2.7 m	2.0 m	2.7 m

3-Electrode Cable Sets

Length	AAMI Part No.	IEC Part No.
1.0 m	M1675A	M1678A
1.0 m	M1671A	M1672A
1.0 m	M1673A	M1674A
0.45 m	M1622A	_
0.7 m	M1624A	M1626A
	1.0 m 1.0 m 1.0 m 0.45 m	Length No. 1.0 m M1675A 1.0 m M1671A 1.0 m M1673A 0.45 m M1622A

5-Electrode Cable Sets

Description	Length	AAMI Part No.	IEC Part No.
OR Grabber shielded	1.0 m/1.6 m	M1973A	M1974A
ICU Grabber shielded	1.0 m/1.6 m	M1968A	M1971A
ICU Snap shielded	1.0 m/1.6 m	M1644A	M1645A
ICU Miniclip non-shielded	0.7 m/1.3 m	M1647A	M1648A

6-Electrode Cable Sets

Description	Length	AAMI Part No.	IEC Part No.
OR Grabber	1.0 m/1.6 m	M1684A	M1685A
ICU Grabber	1.0 m/1.6 m	M1680A	M1681A
ICU Snap	1.0 m/1.6 m	M1682A	M1683A

10-Electrode (5+5)Cable Sets

Description	Length	AAMI Part	IEC Part	
Description Length	Length	No.	No.	
ICU Grabber,	1.0 m	M1976A	M1978A	
chest, shielded				
ICU Snap,	1.0 m	M1602A	M1604A	
chest, shielded				
OR Grabber,	1.0 m	M1979A	M1984A	
chest, shielded				
For Limb Leads see 5-electrode cable sets				

10-Electrode (6+4)Cable Sets

Description	Length	AAMI Part No.	IEC Part No.	
ICU Grabber,	1.0 m	M1532A	M1533A	
chest, shielded				
ICU Snap,	1.0 m	M1537A	M1538A	
chest, shielded				
OR Grabber,	1.0 m	M1557A	M1558A	
chest, shielded				
For Limb Leads see 6-electrode cable sets				

One-piece Cables

Description	Length	AAMI Part No.	IEC Part No.
3-lead Grabber,	1.0 m	989803143181	989803143171
5-lead Grabber,	1.0 m	989803143201	989803143191

Radio-translucent Cables

Pack of five single wires, radio-translucent, 0.9m, M1649A

Set Combiners and Organizers

Set combiners and organizers		Part No.
Set combiner	3-electrode	M1501A
	5-electrode	M1502A
Set organizer for shielded	3-electrode	M1503A
leadsets - grabber and snap	4-electrode	M1664A
5-electrode		M1504A
	6-electrode	M1679A
Set organizer for non-shielded	3-electrode	M1636A
lead sets - miniclip	5-electrode	M1638A
Bedsheet clip		M1509A
Replacement red cover for trunk cable (for 5-		98980814886
electrode cable sets)		1

Philips FAST SpO₂ Accessories

Philips Reusable Sensors

Part Number	Description	Connector Type
M1191A/B M1191AL/BL	Adult Sensor (2 m cable) Adult Sensor (3 m cable)	Philips 8-pin
M1191T	Adult Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable)	Generic D-Sub
M1192A	Small Adult/Pediatric sensor (1.5 m cable)	Philips 8-pin
M1192T	Small Adult Pediatric sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable)	Generic D-Sub
M1193A	Neonatal Hand/Foot Sensor (1.5 m cable)	Philips 8-pin
M1193T	Neonatal Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable)	Generic D-Sub

Part Number	Description	Connector Type
M1194A	Adult/Pediatric Clip Sensor (ear) (1.5 m cable)	Philips 8-pin
M1195A	Infant Sensor (1.5 m cable)	Philips 8-pin
M1196A	Adult Clip Sensor (3 m cable)	Philips 8-pin
	(3 III Cable)	
M1196T	Adult Clip Sensor (requires M1943A	Generic D-Sub
	(1.1 m) or M1943AL (3 m) adapter cable)	

Philips Disposable Sensors

M1131A Adult/Pediatric Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1132A Infant Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1133A Adult/Infant/Neonatal Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1134A Adhesive-free Generic D-Sub Neonatal/Infant/Adult Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable)	Part Number	Description	Connector Type
(requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1133A Adult/Infant/Neonatal Generic D-Sub Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1134A Adhesive-free Generic D-Sub Neonatal/Infant/Adult Sensor (requires M1943A (1.1 m) or	M1131A	Adult/Pediatric Sensor (requires M1943A (1.1 m) or M1943AL	
Sensor (requires M1943A (1.1 m) or M1943AL (3 m) adapter cable) M1134A Adhesive-free Generic D-Sub Neonatal/Infant/Adult Sensor (requires M1943A (1.1 m) or	M1132A	(requires M1943A (1.1 m) or M1943AL	Generic D-Sub
Neonatal/Infant/Adult Sensor (requires M1943A (1.1 m) or	M1133A	Sensor (requires M1943A (1.1 m) or M1943AL (3 m)	Generic D-Sub
adapter cable)	M1134A	Neonatal/Infant/Adult Sensor (requires M1943A (1.1 m) or M1943AL (3 m)	Generic D-Sub

Nellcor Accessories

NELLCOR™ Disposable Sensors¹:

Purchase Nellcor sensors directly from Nellcor.

Product Number	Description	Philips Part Number
OxiMax MAXA ^a	Adult SpO ₂ Sensor	M1904B ^b
OxiMax MAXALa	Adult XL SpO ₂	n/a
	Sensor	
OxiMax MAXPa	Pediatric SpO ₂	M1903B ^b
	Sensor	
OxiMax MAXI ^a	Infant SpO ₂ Sensor	M1902B ^b
OxiMax MAXN ^a	Neonatal-Adult	M1901B ^b
	SpO ₂ Sensor	
Oxisensor II D-25 ^a	Adult Sensor	n/a
Oxisensor II D-20 ^a	Pediatric Sensor	n/a
Oxisensor II I-20 ^a	Infant Sensor	n/a
Oxisensor II N-25 ^a	Neonatal Sensor	n/a
OxiCliq A ^c	Adult SpO ₂ Sensor	n/a
OxiCliq P ^c	Pediatric SpO ₂	n/a
	Sensor	
OxiCliq I ^c	Infant SpO ₂ Sensor	n/a
OxiCliq N ^c	Neonatal-Adult	n/a
	SpO ₂ Sensor	

- a Requires M1943 A(L) adapter cable b not available from Philips in the U.S.A. c Requires M1943 A(L) **and** Nellcor OC3 adapter cables

Masimo Accessories

Adapter cables are available from Philips and also from Masimo. Sensors are available directly from Masimo.

MASIMO LNOP®2 Reusable Sensors:

Product Number	Description
DCI	Reusable Finger Sensor
DCIP	Reusable Pediatric Finger Sensor
YI	Reusable Multi-Site Sensor
TC-I	Reusable Ear Sensor

MASIMO LNCS®1 Reusable Sensors:

Product Number	Description
LNCS DCI	Adult Sensor
LNCS DCIP	Pediatric Sensor
LNCS-TCI	Reusable Ear Sensor

MASIMO LNOP® Disposable Adhesive Sensors:

Description
Adult Adhesive Sensor
Adult Adhesive Sensor
Pediatric Adhesive Sensor
Pediatric Adhesive Sensor
Infant Adhesive Sensor
Neonate Adhesive Sensor
Sensitive Skin Neonate Adhesive Sensor

MASIMO LNCS® Disposable Adhesive Sensors:

Product Number	Description
Adtx	Adult Adhesive Sensor
Pdtx	Pediatric Adhesive Sensor
INF-L	Infant Toe Sensor
Neo-L	Neo Foot Sensor or Adult Finger Sensor
NeoPt-L	Neo Pre-Term Sensitive Skin Adhesive
	Sensors



The Philips M8102A with Philips FAST SpO₂ technology uses Masimo certified pulse oximetry for reduced noise and low perfusion performance with Masimo Sensors under the Masimo NR&LP protocol available from Masimo.

¹ Nellcor[™], Durasensor[™], Dura-Y[™], Oxiband[™], OxiCliq[™], OxiMax[™], MAXFAST[™], are trademarks of Covidien AG and/or its affiliates

² LNOP and LNCS are federally registered trademarks of Masimo Corporation

Extension/Adapter Cables:

Part Number	Description
M1941A	Extension Cable (2m) (8-pin to 8-pin)
M1943A	Adapter Cable (1.1 m) for Philips and
	Nellcor disposable sensors
	(8-pin to 9-pin D-Sub)
M1943AL	Adapter Cable (3m) for Philips and Nellcor
	disposable sensors
	(8-pin to 9-pin D-Sub)
Nellcor OC3	Adapter cable for OxiCliq Sensors (available
	from Nellcor only)
LNOP MP12	LNOP MP Series Patient Cable (3.6 m)
(451261000761)	Adapter Cable for Masimo LNOP Sensors
LNC MP10	LNCS MP Series Patient Cable (3.0 m)
(989803148221)	Adapter Cable for Masimo LNCS Sensors

Nellcor OxiMax SpO₂ Accessories (for M8102A #SP4)

Product Number Description a/ Preferred Application Site Philips Disposable Sensors M1904B b Adult Sensor Requires M1903B b Pediatric Sensor M1943NL M1902B b Infant Sensor adapter cable M1901B b Neonatal/Adult Sensor (included with #A02). NELLCOR OxiMax Sensors ^C MAXA Adult SpO ₂ Sensor Requires MAXAL Adult XL SpO ₂ sensor M1943NL MAXP Pediatric SpO ₂ Sensor adapter cable. MAXI Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. Additionally the Nellcor OC3 adapter cable is needed.			
M1904B b Adult Sensor Requires M1903B b Pediatric Sensor M1943NL M1902B b Infant Sensor adapter cable M1901B b Neonatal/Adult Sensor (included with #A02). NELLCOR OxiMax Sensorsc MAXA Adult SpO2 Sensor Requires MAXAL Adult XL SpO2 sensor M1943NL MAXP Pediatric SpO2Sensor adapter cable. MAXI Infant SpO2Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. Additionally the Nellcor OC3 adapter cable is		Preferred Application	Comments
M1903B b Pediatric Sensor M1943NL M1902B b Infant Sensor adapter cable M1901B b Neonatal/Adult Sensor (included with #A02). NELLCOR OxiMax Sensors MAXA Adult SpO2 Sensor Requires MAXAL Adult XL SpO2 sensor M1943NL MAXP Pediatric SpO2Sensor adapter cable. MAXI Infant SpO2Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. Additionally the Nellcor OC3 adapter cable is	Philips Disposable	Sensors	
M1902B b Infant Sensor adapter cable M1901B b Neonatal/Adult Sensor (included with #A02). NELLCOR OxiMax Sensors MAXA Adult SpO2 Sensor Requires MAXAL Adult XL SpO2 sensor M1943NL MAXP Pediatric SpO2Sensor adapter cable. MAXI Infant SpO2Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. Additionally the Nellcor OC3 adapter cable is	M1904B ^b	Adult Sensor	Requires
M1901B b Neonatal/Adult Sensor (included with #A02). NELLCOR OxiMax Sensors MAXA Adult SpO2 Sensor Requires MAXAL Adult XL SpO2 sensor M1943NL MAXP Pediatric SpO2Sensor adapter cable. MAXI Infant SpO2Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	M1903B ^b	Pediatric Sensor	M1943NL
#A02). NELLCOR OxiMax Sensors MAXA Adult SpO ₂ Sensor Requires MAXAL Adult XL SpO ₂ sensor M1943NL MAXP Pediatric SpO ₂ Sensor adapter cable. MAXI Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	M1902B ^b	Infant Sensor	adapter cable
MAXA Adult SpO ₂ Sensor Requires MAXAL Adult XL SpO ₂ sensor M1943NL adapter cable. MAXI Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A OxiCliq P Pediatric OxiCliq I Infant Neonatal Adult Adult Adult Adult Adult Adult Aditionally the Nellcor OC3 adapter cable is	M1901B ^b	Neonatal/Adult Sensor	`
MAXAL Adult XL SpO ₂ sensor M1943NL MAXP Pediatric SpO ₂ Sensor MAXI Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A OxiCliq P Pediatric M1943NL Adult Requires M1943NL adapter cable. Additionally the Nellcor OC3 adapter cable is	NELLCOR OxiMa	x Sensors ^c	
MAXP Pediatric SpO ₂ Sensor Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A OxiCliq P Pediatric OxiCliq I Infant OxiCliq I Neonatal Additionally the Nellcor OC3 adapter cable is	MAXA	Adult SpO ₂ Sensor	Requires
MAXI Infant SpO ₂ Sensor MAXN Neonatal-Adult Sensor OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	MAXAL	Adult XL SpO ₂ sensor	M1943NL
MAXN Neonatal-Adult Sensor OxiCliq A OxiCliq P Pediatric OxiCliq I Infant OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	MAXP	Pediatric SpO ₂ Sensor	adapter cable.
OxiCliq A Adult Requires OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	MAXI	Infant SpO ₂ Sensor	
OxiCliq P Pediatric M1943NL OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	MAXN	Neonatal-Adult Sensor	
OxiCliq I Infant adapter cable. OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	OxiCliq A	Adult	Requires
OxiCliq N Neonatal Additionally the Nellcor OC3 adapter cable is	OxiCliq P	Pediatric	M1943NL
the Nellcor OC3 adapter cable is	OxiCliq I	Infant	adapter cable.
	OxiCliq N	Neonatal	the Nellcor OC3 adapter cable is

Product Number	Description ^a / Preferred Application Site	Comments
MAXR	Adult SpO ₂ Nasal	Requires
	Sensor	M1943NL
MAXFAST	Forehead SpO ₂ Sensor	adapter cable.
Oxiband OXI-A/N	Adult-Neonatal SpO ₂	
	Sensor with Wraps	
Oxiband OXI-P/I	Pediatric-Infant SpO ₂	
	Sensor with Wraps	
SoftCare SC-A	Adult SpO ₂ Sensor	
SoftCare SCNEO-	Neonatal SpO ₂ Sensor	
1		
SC-PR-I	Preemie SpO ₂ Sensor	
Durasensor	Adult SpO ₂ Sensor	
DS100A-1		
Dura-Y D-YS	SpO_2 Sensor	
Extension/Adapte	r Cables	
M1943NL	Adapter Cable (3 m)	
Nellcor OC3	Adapter Cable for	Available from
	Nellcor OxiCliq sensors	Nellcor only.

a For application site, please refer also to the Instructions for Use provided with the sensors.

Non Invasive Blood Pressure Accessories



These cuffs and tubings are designed to have special protection against electric shocks (particularly regarding allowable leakage currents), and are defibrillator proof.

Multi-Patient Comfort Cuffs and Disposable Cuffs			
Patient Category	Disposable cuff	Reusable cuff	
Adult (Thigh)	M1879A	M1576A	
Large Adult	M1878A	M1575A	
Adult	M1877A	M1574A	
Small Adult	M1876A	M1573A	
Pediatric	M1875A	M1572A	
Infant	M1874A	M1571A	
Tubing: Use M1598B or M1599B			

Reusable Cuff Kits	Part No.
Infant, pediatric, small adult, adult	M1577A
Small adult, adult, large adult, thigh	M1578A

sensors.
b Philips disposable sensors M1901B, M1902B, M1903B and M1904B are not available in the USA.

c Can only be ordered from Nellcor.

Reusable Cuff Kits	Part No.
Infant, pediatric, small adult, adult, large	M1579A
adult, thigh	

Adult/Pedia	tric Antimicro	bial Coated F	Reusable cuffs
Cuff Size	Circumfer	Bladder	Single-Hose Part
(color)	ence (cm)	Width	No.
Infant	9.0 - 14.8	5.4 cm	M4552A
(orange)		2.1 inches	
Pediatric	13.8 - 21.5	8.0 cm	M4553A
(green)		3.1 inches	
Small Adult	20.5 - 28.5	10.6 cm	M4554A
(royal blue)		4.2 inches	
Adult	27.5 - 36.5	13.5 cm	M4555A
(navy blue)		5.3 inches	
Adult	27.5 - 36.5	13.5 cm	M4556A
X-long		5.3 inches	
(navy blue)			
Large Adult	35.5 - 46.0	17.0 cm	M4557A
(burgundy)		6.7 inches	
Large Adult	35.5 - 46.0	17.0 cm	M4558A
X-long		6.7 inches	
(burgundy)			
Thigh	45 - 56.5	21.0 cm	M4559A
(grey)		8.3 inches	

Tubing: Use M1598B or M1599B

Adult/Pediatric Soft Single Patient Single-Hose Disposable Cuffs			
Patient Category	Limb Circumference	Bladder Width	Disposable cuff Part No.
Adult (Thigh)	45.0 - 56.5 cm	20.4 cm	M4579A
Large Adult X-long	35.5 - 46.0 cm	16.4 cm	M4578A
Large Adult	35.5 - 46.0 cm	16.4 cm	M4577A
Adult X-long	27.5 - 36.5 cm	16.4 cm	M4576A
Adult	27.5 - 36.5 cm	13.1 cm	M4575A
Small Adult	20.5 - 28.5 cm	10.4 cm	M4574A
Pediatric	15.0 - 21.5 cm	8.0 cm	M4573A
Infant	9.0 - 15.0 cm	5.6 cm	M4572A

Tubing: Use M1598B or M1599B

Neonatal/Infant Cuffs (Disposable, non-sterile)			
Cuffs	Limb	Bladder	Part No.
Cuiis	Circumference	Width	FAIL INU.
Size 1	3.1 to 5.7 cm	2.2 cm	M1866A
Size 2	4.3 to 8.0 cm	2.8 cm	M1868A
Size 3	5.8 to 10.9 cm	3.9 cm	M1870A
Size 4	7.1 to 13.1 cm	4.7 cm	M1872A

Tubing: Use M1596B or M1597B

Cuff Tubing		
Adult	1.5 m /4.9'	M1598B
	3.0 m/9.8'	M1599B
Neonatal	1.5 m /4.9'	M1596B
	3.0 m/9.8'	M1597B

Temperature Accessories

Temperature Probes	Part No.
Reusable	
General purpose probe	21075A
Small flexible vinyl probe	21076A
(Infant/Pediatric)	
Attachable surface probe	21078A
Disposable	
General purpose probe	M1837A
Skin probe	21091A
Esophageal/Stethoscope Probe	21093A
(12 French)	
Esophageal/Stethoscope Probe	21094A
(18 French)	
Esophageal/Stethoscope Probe	21095A
(24 French)	
Foley Catheter Probe	M2255A
(12 French)	
Foley Catheter Probe	21096A
(16 French)	
Foley Catheter Probe	21097A
(18 French)	
Adapter cable 1.5 m/4.9'	21082B
Adapter cable 3.0 m/9.8'	21082A

PRESS Accessories

These transducers and accessories are designed to have special protection against electric shocks (particularly regarding allowable leakage currents), and are defibrillator proof.

Pressure Transducers and Accessories	Part No.
Reusable	
Reusable pressure transducer	CPJ840J6
$5 \mu V/V/mmHg$ sensitivity	
Sterile disposable pressure domes for	CPJ84022
CPJ840J6 (pack of 50)	
Transducer holder for CPJ840J6	CPJ84046
(pack of 4)	
IV pole mount for CPJ840J6	CPJ84447
Disposable (EU/EFTA only. Not available	e in USA)
Single channel disposable sensor kit (20)	M1567A
Dual channel disposable sensor kit (20)	M1568A
Transducer holder for M1567/8A	M2271A
IV pole mount for M1567/8A	M2272C
Adapter cable for disposable sensor kit,	M1634A
3.0 m, for M1567/8A	

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M8102A complies with the requirements of the Council Directive 93/42/EEC of 14 June 1993 (Medical Device Directive).

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