



Liberty II

Dynamic Low Air Loss Mattress System

UPRA367804-2

User Manual

Issue 8 – 17/12/2020 UPRA367804-2.UM-8

Contents

1.	Introduction	4
2.	Intended Use	4
	2.1 Contraindications	4
3.	About the Product	4
4.	Symbols and Statements	6
5.	Important Safety Information	7
6.	Technical Specification	8
	6.1 Power unit	8
	6.2 Mattress	8
7.	Installation and Set-Up	9
	7.1 Setting up mattress	9
	7.2 Setting up power unit	9
	7.3 Cable management system	10
	7.4 Connecting mattress to the power unit	10
8.	Control Panel Operation Guide	11
	8.1 Comfort level	12
	8.2 Bariatric	12
	8.3 Control panel lockout	12
	8.4 Function mode switch	12
	8.4.1 Alternate	12
	8.4.2 Pulsate	13
	8.4.3 Static	13
	8.5 Auto-Firm	13
	8.6 Operate / standby	13
	8.7 Alternate / pulsate cycle time selection	13
	8.8 Alarm mute	14
9.	CPR Mode	14
10.	Transport Mode	14
11.	Alarms & Fault Findings	15
	11.1 Low pressure alarm	15
	11.2 Power failure alarm	15
	11.3 Service (alternating failure alarm)	15
12.	Troubleshooting	17
13.	Cleaning and Decontamination	18
	13.1 Basic cleaning information	18
	13.2 Mattress and cover cleaning	
	13.3 Power unit cleaning	19
	Storage	
15.	Service and Maintenance	19
	15.1 General	19
	15.2 Fuse replacement	19
	15.3 Air filter replacement	
16.	EMC Information	20

1. Introduction

This User Manual contains instructions for the installation, use and maintenance of the Ultimate Healthcare Liberty II dynamic low air loss mattress system. You must read and fully understand this manual before using the system.



Caution: Ultimate Healthcare shall not be liable for any damage or injury caused by failure to follow the proper instructions as described in this User Manual.



Caution: Before using the dynamic low air loss mattress system all staff must familiarise themselves thoroughly with the various parts and controls as detailed in this User Manual.

Note: Ultimate Healthcare reserves the right to modify the information in this User Manual at any time. The information in this User Manual may vary slightly with respect to the basic design of the product.

2. Intended Use

This product is intended to help and reduce the incidence of pressure ulcers while optimising patient comfort. It is also indicated for the following purposes:

 Individual home care setting, long-term care & Acute settings of whom suffering from pressure ulcers

2.1 Contraindications

• The mattress is not suitable for use on patients with unstable fractures.

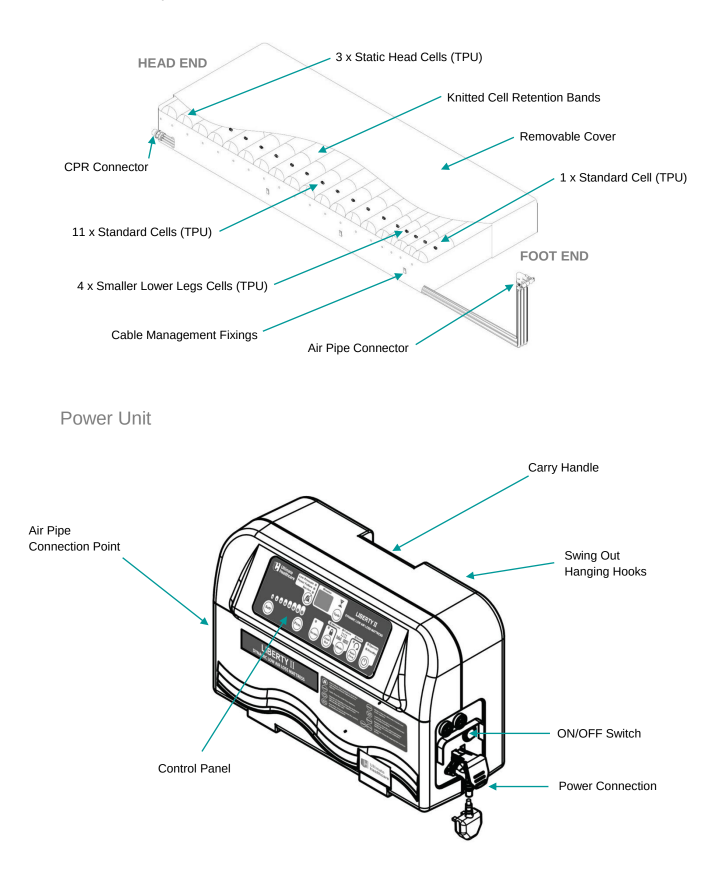
3. About the Product

The Liberty II dynamic Low Air Loss mattress replacement provides continuous low pressure and is designed for use with patients who are at Very High Risk of developing pressure ulcers. The system is particularly suitable for use with terminally ill and frail patents together with the treatment of burns and patients with lymphedema.

The Liberty II delivers *true Low Air Loss* therapy and limits both skin warming and moisture accumulation. *Constant low pressures* provide a gentle support surface that allows patient immersion, increasing the surface area of the skin that is in contact with the mattress resulting in reduced interface pressures.

The excellent *microclimate control* helps to reduce the build-up of heat and moisture ensuring that it is ideal for patients who need special protection from the damaging effects of excess moisture.

Mattress Replacement



4. Symbols and Statements



Note: Indicates tips and advice for the correct use of this product.

Caution: Indicates correct operating or maintenance procedures in order to prevent damage to or destruction of the product or other property.

Warning: Indicates potential danger that requires correct procedures or practises in order to prevent personal injury.

B symbol, indicates this product is according to degree of protection against electric shock for type B equipment



Class 1 electrical device.



Caution, read the instructions before use.

IP21

Water and dust protection classification.



Disposal of electrical and electronic equipment (WEEE): This product should be handed over to an applicable collection point for the recycling of electrical and electronic equipment.

CE certified.

Issue 8 - 17/12/2020

5. Important Safety Information

Please read all instructions prior to using any Ultimate Healthcare supplied product. The Liberty II dynamic low air loss mattress system must be used in accordance with this User Manual.



The mattress must only be operated by personnel who have been properly trained or have suitable experience with products of this nature.



Ensure a clinical Risk Assessment is conducted, which should take account of the suitability of use of this product, patient's condition, any ancillary equipment in use and the surrounding environment. This should include assessing the use of side rails, head and footboards etc. Pressure settings should be advised or prescribed by a medical practitioner.



Only personnel trained or formally approved by Ultimate Healthcare in operation and maintenance of Ultimate Healthcare products may perform maintenance; modification or repair work on any Ultimate Healthcare supplied product.



Ensure the power cable is not trapped or twisted and is routed suitably to avoid crushing or entrapment when connected to the product.



Do not use your mattress system power unit in the presence of flammable gases. This excludes oxygen cylinders.



Avoid hazards caused by inappropriate handling of the power cable e.g. by kinking, shearing or other mechanical damages.



The power cable for this product must be unplugged from the mains power outlet socket and suitably stowed before moving, cleaning or maintenance activities.

Do not secure mattress straps to removable head or footboards or any fixed (non-moving) parts of a profiling bedframe.

Disconnect from mains (power supply) before cleaning the power unit.

When cleaning do not immerse the power unit in water.



Use only the cleaning and disinfectant agents recommended in this User Manual.



When connecting product after transportation or storage, inspect the power cable visually for any signs of damage. If evident, do not use product and contact Ultimate Healthcare or your local distributor for repair.



When storing, ensure the product is stored away from direct sunlight and extreme cold conditions.



Never block the air openings of this product or place it on a soft surface, such as a bed or couch, where their openings may be blocked. Keep the air opening free of lint, hair, and other similar particles. Air ventilation through the power unit is vital for correct and safe operation.



If this product is used for any activity other than detailed within this User Manual then personal risk to the end user or patient may occur. Ultimate Healthcare shall not be held liable/responsible for such an event.



Do not smoke, or allow the patient to smoke when using this product. Keep all possible ignition sources clear of this product

6. Technical Specification

6.1 Power unit

Dimensions:	185mm x 390mm x 280mm (D x W X H)
Weight:	6.2 kg
Alternating cycle time:	5-30 mins
Pulsate cycle time:	1-20 mins
Output pressure range:	20 to 50mmHg (+/-2)
Power supply:	AC 230V 50 Hz
Current:	0.12 amp
Classification:	Class I, Type B
Warranty:	2 years
Operation environment:	5°C to 40°C 15%RH ~ 93%RH
Storage environment:	-25°C~70°C ≦93%RH
Environment pressure:	70 kPa-101.3 kPa no condensation
Water & dust protection classification:	IP21

6.2 Mattress

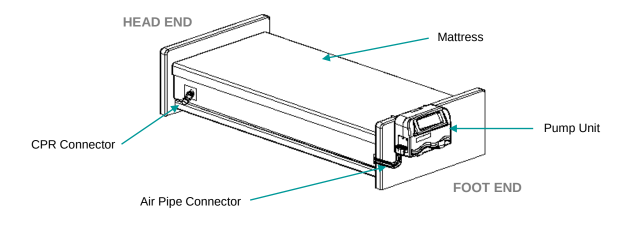
Dimensions:	2000mm x 890mm x 210mm (L x W X H)
Weight:	10.5 kg
No of cells:	19
Cover material:	2 way stretch polyester with PU Coated
Bottom material:	Polyester with PU coated
Max user weight:	320 kg / 50 stone

7. Installation and Set-Up

7.1 Setting up mattress

Secure the mattress to the moving parts of the bedframe using the securing straps, and when the system is fully inflated it is ready for the patient to be positioned onto the mattress. The mattress is designed to completely replace any existing mattress which may be in use on a bed.

Remove any existing mattress and ensure that there are no protruding parts or sharp objects on the bed, which could cause damage to the mattress. Lay the mattress on the bed patient surface ensuring that the air pipe connector is situated on the bottom left hand side of the bed foot end (as viewed from the foot of the bed).



There are security straps fitted to the base cover of the mattress, which should be fastened loosely to convenient points on the bedframe patient surface.



Caution: Ensure when fixing the mattress to the bed the security straps are only fitted to the moving parts of the mattress platform. Straps secured to the fixed parts of the mattress platform will damage the bed/ mattress when operated.

7.2 Setting up power unit

Whilst holding the power unit, unfold the hanging hooks on the rear of the power unit and hang it from the beds footboard. If required the power unit can be placed on the floor at the foot of the bed.



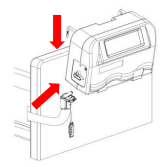
• Caution: Ensure that the power cable is routed in such a manner so that it cannot be twisted, trapped, crushed or stressed.

7.3 Cable management system

Cable management fixings are located on each side of the mattress underneath the flap of the cover. The mains power cable should be secured through the cable management fixing as follows:

- Locate each cable management fixing.
- If necessary, open the press studs.
- Run the mains power cable along the side of the mattress securing each fixing loop around the cable using the press studs.

7.4 Connecting mattress to the power unit



Unplug the cover of the air pipe connector and connect the air pipe connector to the power unit and then ensure that the air tubes are free from any kind of obstruction, and are not kinked.

Plug the power cable into a suitable electrical socket and switch 'ON' using mains power switch found at the side of the power unit. The *Standby* indicator on the control panel will light up.

Push the *Operate* button and the system will start inflation and the *Auto-Firm* indicator will flash.

Once the mattress is fully inflated, you can then set the mattress to the appropriate setting suitable for the patient. *Please refer to Table 1 for the most suitable settings.*



Caution: The power unit must only be connected to the mattress recommended by the manufacturer. Do not use it for any other purpose.



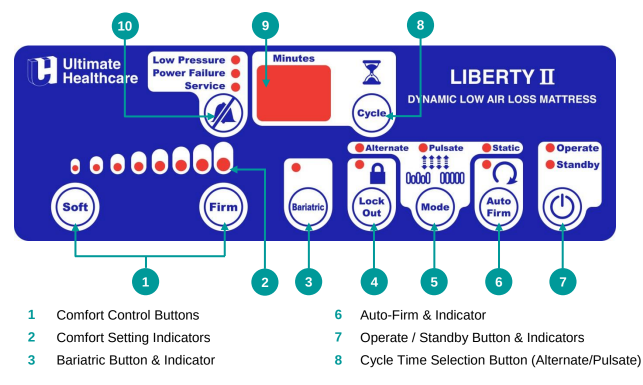
Note: Ensure that the CPR valve is set to the CLOSED position.

Comfort Control	Bariatric							Pati	ent	Weig	ght (KG)						
		45	55	65	80	90	105	120	135	150	170	190	210	230	250	270	300	320

Table 1: Weight and Suggested Comfort Level Reference Table

8. Control Panel Operation Guide

The Control Panel of the power unit is used to make adjustments to the mattress and also indicates fault conditions/service requirements. These are either visual (indicator lights) or audible.



- 4 Control Panel Lockout Button & Indicator
- 5 Function Mode Selection Button
 - Alternate / Pulsate / Static

8.1 Comfort level



The **Soft** and **Firm** buttons allow carers to adjust pressures within a safe pre-set range to provide patients with enhanced comfort or support whilst maintaining a very good level of protection and therapy. Qualified clinical advice must always be taken before adjusting mattress pressures.



The required pressure is selected using the *Soft* and *Firm* buttons to move the pressure by one step at a time.

When pressing the *Firm* button, the output pressure will increase to provide a higher pressure output and thus increased support.

When pressing the *Soft* button, the output pressure will be decreased to provider a lower pressure output and thus increased comfort.

To check if the pressure is adequately supporting the patient, slide one hand between the air mattress and bed frame to feel under the patient's bottom. You should be able to slide the hand in-between and an acceptable range is approximately 25 to 40 mm (1" to 1-1/2") to ensure the patient Is not bottoming out.

8.2 Bariatric



This feature will enhance the power output of the power unit for heavier patient support. *Please refer to Reference Table 1 for the weight and comfort selection guide.*

8.3 Control panel lockout



If the control panel is not used for a period of 30 seconds it will lock out and inhibit the use of the functions. Additionally, if you wish to lock out the control panel press the *Lock Out* button and all functions will be locked. This is to prevent the system being altered accidently.

In order to unlock the control panel, simply press and hold the *Lock Out* button for 3 seconds, the control panel will now be active for use.

8.4 Function mode switch

8.4.1 Alternate

- 9 Cycle Time Display
- **10** Alarm Mute Button & Indicators



Within this mode the mattress will operate in an alternating 1-in-2 cell cycle. The alternating cycle will continue at the selected cycle time until another mode is selected.

8.4.2 Pulsate



Pressing the *Function Mode Selection* button until the *Pulsate* indicator illuminates puts the system into *Pulsate* mode.

Within this mode the pressure in each cycle is reduced to 40% lower than the surface pressure. *Pulsate* mode will continue until another mode is selected.

8.4.3 Static



Pressing the *Function Mode Selection* button until the *Static* indicator illuminates puts the system into *Static* mode.

Within this mode the mattress will maintain selected constant pressure.

8.5 Auto-Firm



Auto-Firm mode can be selected by pressing the *Auto-Firm* button. In this mode all cells inflate to a single pressure setting to provide a firm and stable surface for nursing procedures or for patient ingress/egress. The system will automatically return to the previously selected mode and comfort level after 20 minutes.

8.6 Operate / standby



Press the *Operate / Standby* button to turn the power unit ON. Press again to turn OFF/Standby the power unit.



Note: The power switch on the side of the power unit must be turned ON.

8.7 Alternate / pulsate cycle time selection



The *Alternating* and *Pulsate* cycle times can be selected to provide an individualised care program for each patient.

Within *Alternating* mode the alternating cycle time can be selected by pressing the *Cycle* button. Selections can be made from 5-30 minutes at 5 minute intervals.

Within *Pulsate* mode the pulsate cycle time can be selected by pressing the *Cycle* button. Selections can be made from 1-20 minutes at 1 minute intervals.

8.8 Alarm mute



The *Alarm Mute* button temporarily resets the audible *Low Pressure/Power Failure/Service* alarms. Should the situation not be resolved and the fault condition continues the alarm will resume notifying carer.

9. CPR Mode

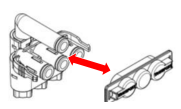


CPR (Cardio Pulmonary Resuscitation) can be performed using the red CPR valve which is situated at the head end on the left hand side of the mattress.

For rapid deflation gently pull and rotate the dial of the CPR valve to *click* into the OPEN position. At the same time, disconnect the air pipe connector from the power unit to speed up the air release.

If re-inflating the mattress, ensure the dial of the CPR attachment is rotated until it '*clicks*' into the CLOSED position.

10. Transport Mode



If the patient is being moved on the mattress, or there is a power cut, general pressure can be maintained in the system for an adequate period of time whilst disconnected from the mains.

Simply disconnect air pipe connector and place the connector cover over it. The air pressures in the mattress will now equalise, but maintain a degree of comfort. This will maintain the cells in their present state for approximately 24 hours.

It is important to restore the Liberty II dynamic low air loss mattress system as quickly as possible by reconnecting the supply tubes to the power unit.

11. Alarms & Fault Findings

The Liberty II system is equipped with audible and visual alarm indicators. These alert the user to the status of the available mains supply and any mattress defect.

11.1 Low pressure alarm

Upon detection of low pressure, an audible alarm will be heard and the *Low Pressure* indicator will illuminate. The audible alarm may be cancelled by pressing the *Alarm Mute* button. The *Low Pressure* indicator will continuously illuminate until the low pressure fault condition is resolved.

This condition could be caused, for example, by incorrect fitting of the air pipe connector, opening of the CPR valve or a leak in the mattress due to a cut or puncture.

11.2 Power failure alarm

If at any time the mains power should be removed from the power unit or the power cable is unplugged without turning the power unit OFF, an audible alarm will be heard and the *Power Failure* indicator will illuminate. The audible alarm may be cancelled by pressing the *Alarm Mute* button.



Note: When the power unit has not been used for more than 3 months, it might require 60 minutes operating time (or more) for the Alarm to function correctly.

11.3 Service (alternating failure alarm)

Should your system develop a fault condition whilst in use, an audible alarm will be heard and the **Service** indicator will illuminate. The audible alarm may be cancelled by pressing the **Alarm Mute** button. Please refer to *Table 2* for error codes and call Ultimate Healthcare or your local distributor.

Table 2: Alert/Error Code Reference Table

Priority High ↓ Low	Warning Code	Indicator LED	Audible Output Mode	Condition of Output	Warning Description	Remarks	
0	N/A	N/A	ONCE	Not in System Shutdown	Key Tone	Key Tone from Functional Button	
1	5.d.	Power Failure	ONCE	Power-Off	System Shutdown	Shutdown	
2	8.8	ALL LED	ONCE	Operate or Standby	Power-On	All Indicators Light On	
3	N/A	N/A	ONCE	Operate or Standby	State / Mode Switching	No Display	
4	1.8.	Auto-Firm	ONCE	Operate	Mattress Inflation Completion	Inflation Ended	
5	R.E.	Auto-Firm	ONCE	Operate	Auto-Firm Completion	Auto-Firm Ended	
6	5.8.	S.E. Static		Operate	Static Completion	Static Ended	
7	N/A	Power Failure			Power Failure Alarm	No Display	
8	I .F.	Low Pressure	REPEAT (Cycle 4 sec)	Operate or Standby	Power-On Inflation Failure Alarm	Inflate Failure	
9	R.F.	Low Pressure	REPEAT (Cycle 4 sec)	Operate or Standby	Auto-Firm Failure Alarm	Auto-Firm Failure	
10	L.P.	Low Pressure	REPEAT (Cycle 4 sec)	Operate or Standby	Low Pressure Overtime Alarm	Low Pressure	
11	H.P.	Service	REPEAT (Cycle 4.5 sec)	Operate or Standby	High Pressure Overtime Alarm	High Pressure	
12	H.E.	Service	REPEAT (Cycle 4.5 sec)	Operate or Standby	High Ambient Temperature Alarm	High Temperature	
13	U. I	Service	REPEAT (Cycle 4.5 sec)	Operate or Standby	Air Valve 1 Positioning Failure Alarm	Air Valve 1 failure	
14	U.2	Service	REPEAT (Cycle 4.5 sec)	Operate or Standby	Air Valve 2 Positioning Failure Alarm	Air Valve 2 failure	
15	L.b.	Service	REPEAT (Cycle 15 sec)	Operate or Standby	Battery Low Alarm	Battery would need to be replaced	
16	C <u>.U</u>	NONE	NONE	Factory Calibration Mode	Calibration Not Completed	Calibration Unfinished	
17	C.C.	NONE	NONE	Factory Calibration Mode	Calibration Completed	Calibration Completed	

12. Troubleshooting

PROBLEM	SOLUTION
No lights on power unit	 Check the power unit is connected to the mains power supply and that the mains switch is turned ON. Check power unit for any blown fuses.
Low Pressure indicator is flashing and sounding	 Check to ensure that the CPR is in the CLOSED position. Check whether power was suddenly shut down. Check that the connection between air tube and power unit is tightly secured. Check that all tubing connections along the mattress are secured.
Power Failure Alarm Indicator is flashing and sounding	• Check the power unit is connected to the mains power supply and that the mains switch is turned ON.
Patient is bottoming out (without alarm being triggered)	 Pressure setting might be inadequate for the patient, adjust comfort level to Firm and wait for a few minutes for a better comfort.

If the problem persists, contact Ultimate Healthcare or your local service provider.

13. Cleaning and Decontamination

It is recommended that the system is cleaned regularly and after each patient use.



Warning: Disconnect the power unit from the electricity supply before carrying out cleaning/decontamination procedures.

13.1 Basic cleaning information



Caution: Only use disinfectants designed for cleaning healthcare equipment i.e. Sodium Hypochlorite or similar (up to 10,000 ppm available chlorine).

- Caution: Do not use abrasives (scouring powder), scourers or other materials/agents which could damage the mattress system.
- Caution: Do not use sodium carbonate or phenol based solutions.
- Caution: Do not use fabric softener or non-biological washing detergent.
- Caution: Do not immerse the power unit in water.



- Caution: When cleaning/disinfecting, ensure that only a damp cloth is used.
 Caution: After cleaning, dry the mattress out of direct exposure to sunlight.
- \wedge
- Caution: Using inappropriate detergents or disinfectants and not observing the manufacturer's guidelines may result in damage to the mattress which Ultimate Healthcare cannot be held liable for.



 Caution: The appropriate qualified staff must be consulted when specifying a suitable cleaning fluid. Ultimate Healthcare shall not be liable for any damages caused by the use of inappropriate detergents or disinfectants.

13.2 Mattress and cover cleaning

The Liberty II mattress, mattress cover and air pipe cover can be cleaned using the following simple procedures in accordance with your Local Infection Control Policy:

- Liberally swabbing with a damp cloth pre-soaked with hot water at 84°C containing detergent, and then drying.
- Swabbing with a solution of sodium hypochlorite (up to 10,000 parts per million available chlorine) and then drying.
- Where required mattress covers can be laundered in a pre-wash at 60°C for up to 15 minutes and in a main wash at 84°C for up to 15 minutes. This should be followed by a cold rinse and extraction.

The mattress cover and air pipe cover can also be machine washed. The mattress cover and mattress must be dry prior to refitting.

13.3 Power unit cleaning

The Power Unit can be cleaned by wiping down with a cloth dampened with hot water at 60^oC containing detergent or with sodium hypochlorite (up to 10,000 parts per million available chlorine).

14. Storage

The mattress should be loosely rolled lengthwise with the cover innermost, taking care not to strain the air pipe. It should then be placed in in a suitable protective cover with the power unit and stored in an area appropriate for electronic medical devices.

- To quickly extract air out from mattress for storage, rotate the CPR value to the OPEN position and disconnect the air hose connector to release the air.
- Lay the mattress out flat and upside down.
- Roll from the head end towards the foot end.
- The foot end strap can then be stretched around the rolled mattress to prevent unrolling.
- The power cord could be wrapped around the power unit bumper or disconnected for storage.



Caution: Do not fold, crease or stack mattresses

15. Service and Maintenance

It is recommended that this product be part of a routine preventative maintenance schedule with a planned service every 12 months regardless of product usage.

15.1 General

- (1) Check power cable and plug if there are abrasions or excessive wears.
- (2) Check mattress cover for signs of wear or damage. Ensure mattress cover and tubes are stubbed together correctly.
- (3) Check the air hoses for any kink or break. For replacement, please contact Ultimate Healthcare or your local distributor.

15.2 Fuse replacement

- (1) Disconnect the plug from mains power when a blown fuse is suspected.
- (2) Remove the cover of the fuse holder by means of a small screwdriver.
- (3) Insert a new fuse of the correct rating in, and replace the cover of the fuse holder back. The fuse should be rated as T1A.

15.3 Air filter replacement

(1) Replace the air filter located at the handle on the back of the power unit.

- (2) The filter is reusable and can be washed gently with a mild detergent and water. Dry the filter before use.
- (3) Check and replace air filter regularly if environment is dirty.

16. Electromagnetic Compatibility (EMC) to IEC 60601-1-2:2014

Manu	Manufacturer's declaration-electromagnetic emissions							
The <u>device(s)</u> is intended for use in the electromagnetic environment (for home and professional healthcare) specified below. The customer or the user of the <u>device(s)</u> should assure that it is used in such an environment.								
Emission test Compliance Electromagnetic environment-guidance (for home and professional healthcare environment)								
RF emissions CISPR 11	Group 1	The <u>device(s)</u> uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.						
RF emissions CISPR 11	Class B	The <u>device(s)</u> is suitable for use in all establishments, including domestic						
Harmonic emissions IEC 61000-3-2	Class A	establishments and those directly connected to the public low-voltage power supply network that						
Voltage fluctuations /flicker emissions IEC 61000-3-3								

	Manufacturer's declaration-electromagnetic immunity								
The device(s) is intended for	use in the electromagnetic environ	ment (for home and professional h	ealthcare) specified below.						
The customer or the user of the device(s) should assure that it is used in such an environment.									
Immunity test	IEC 60601	Compliance level	Electromagnetic environment-						
	test level		guidance (for home and						
			professional healthcare environment)						
Electrostatic	Contact:±8 kV	Contact: ±8 kV	Floors should be wood, concrete or						
discharge(ESD)	Air ± 2 kV, ± 4 kV, ± 8 kV, ± 15 kV	Air ± 2 kV, ± 4 kV, ± 8 kV, ± 15 kV	ceramic tile. If floors are covered with						
IEC 61000-4-2			synthetic material, the relative humidity						
			should be at least 30%						
Electrical fast	+ 2kV for power supply lines	+ 2kV for power supply lines	Mains power quality should be that of a						
transient/burst	+ 1kV for input/output lines	+ 1kV for input/output lines	typical home healthcare environment.						
IEC 61000-4-4									
Surge	+ 0.5kV, +1kV line(s) to line(s)	+ 0.5kV, +1kV line(s) to line(s)	Mains power quality should be that of a						
IEC 61000-4-5	<u>+</u> 0.5kV, <u>+</u> 1kV, <u>+</u> 2kV line(s) to earth	Not applicable	typical home healthcare environment.						
Voltage Dips, short	Voltage dips:	Voltage dips:	Mains power quality should be that of a						
interruptions and voltage	0 % <i>U</i> _T ; 0,5 cycle	0 % <i>U</i> r; 0,5 cycle	typical home healthcare environment. If						
variations on power supply	0 % <i>U</i> r; 1 cycle	0 % <i>U</i> r; 1 cycle	the user of the SD480-211 requires						
input lines	70 % <i>U</i> _T ; 25/30 cycles	70 % <i>U</i> _T ; 25 cycles	continued operation during power mains						
IEC 61000-4-11			interruptions, it is recommended that the						
	Voltage interruptions:	Voltage interruptions:	SD480-211 be powered from an						
	0 % U _T ; 250/300 cycle	0 % <i>U</i> _T ; 250 cycle	uninterruptible power supply or a battery.						
Power frequency(50, 60	30 A/m	30 A/m	The SD480-211 power frequency						
Hz) magnetic field	50 Hz or 60 Hz	50 Hz	magnetic fields should be at levels						
IEC 61000-4-8			characteristic of a typical location in a						
			typical home healthcare environment.						
NOTE UT is the a.c. mains v	voltage prior to application of the test	st level.							

below.		sgriotio on monitori (ior i	nome and professional healthcare) specified
The customer or the use	er of the <u>device(s)</u> should IEC 60601 test level	assure that it is used in s Compliance level	Electromagnetic environment-guidance (for home and professional healthcare environment)
Conducted RF IEC 61000-4-6	3 Vrms: 0,15 MHz – 80 MHz 6 Vrms: in ISM and amateur radio bands between 0,15 MHz and 80 MHz 80 % AM at 1 kHz	3 Vrms: 0,15 MHz – 80 MHz 6 Vrms: in ISM and amateur radio bands between 0,15 MHz and 80 MHz 80 % AM at 1 kHz	Portable and mobile RF communications equipment should be used no closer to any part of the <u>device(s)</u> including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
Radiated RF IEC 61000-4-3	10 V/m 80 MHz – 2,7 GHz 80 % AM at 1 kHz	10 V/m 80 MHz – 2,7 GHz 80 % AM at 1 kHz	Recommended separation distance: $d = 1, 2 \sqrt{P}$ $d = 1, 2 \sqrt{P}$ 80MHz to 800 MHz $d = 2, 3 \sqrt{P}$ 800MHz to 2,7 GHz Where <i>P</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in metres (m). Interference may occur in the vicinity of equipment marked with the following symbol: ((()))

Recommended separation distance between portable and mobile RF communications equipment and the device(s)

The <u>device(s)</u> is intended for use in an electromagnetic environment (for home and professional healthcare) in which radiated RF disturbances are controlled. The customer or the user of the <u>device(s)</u> can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the <u>device(s)</u> as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of	Separation distance according to frequency of transmitter m						
transmitter W	150 kHz to 80 MHz d =1,2√ <i>P</i>	80 MHz to 800 MHz d =1,2√P	800 MHz to 2,7 GHz d =2,3√P				
0,01	0,12	0,12	0,23				
0,1	0,38	0,38	0,73				
1	1,2	1,2	2,3				
10	3,8	3,8	7,3				
100	12	12	23				

For transmitters rated at a maximum output power not listed above, the recommended separation distance *d* in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where p is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies. NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Manufacturer's declaration-electromagnetic immunity

Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipment

The <u>device(s)</u> is intended for use in the electromagnetic environment (for home and professional healthcare) specified below. The customer or the user of the <u>device(s)</u> should assure that it is used in such an environment.

Test frequency (MHz)	Band ^{a)} (MHz)	Service ^{a)}	Modulation ^{b)}	Maximum power (W)	Distance (m)	IMMUNITY TEST LEVEL (V/m)	Compliance LEVEL (V/m) (for home and professional healthcare)
385	380 –390	TETRA 400	Pulse modulation b) 18 Hz	1,8	0,3	27	27
450	430 – 470	GMRS 460, FRS 460	FM c) ±5 kHz deviation 1 kHz sine	2	0,3	28	28
710			Pulse				
745	704 – 787	7 LTE Band 13, 17	modulation b) 217 Hz	0,2	0,3	9	9
780							
810		GSM 800/900,					
870	800 – 960	TETRA	Pulse modulation b) 18 Hz	2	0,3	28	28
930		CDMA 850,					
1 720		GSM 1800; CDMA					
1 845	1 700 – 1 990	1 990 1900;	Pulse modulation b) 217 Hz	2	0,3	28	28
1 970		DECT; LTE Band 1, 3					
2 450	2 400 - 2 570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation b) 217 Hz	2	0,3	28	28
5 240			Pulse				
5 500	5 100 – 5 800	WLAN 802.11 a/n	modulation b)	0,2	0,3	9	9
5 785			217 Hz				

NOTE If necessary to achieve the IMMUNITY TEST LEVEL, the distance between the transmitting antenna and the ME EQUIPMENT or ME SYSTEM may be reduced to 1 m. The 1 m test distance is permitted by IEC 61000-4-3.

^{a)} For some services, only the uplink frequencies are included.

^{b)} The carrier shall be modulated using a 50 % duty cycle square wave signal.

c) As an alternative to FM modulation, 50 % pulse modulation at 18 Hz may be used because while it does not represent actual modulation, it would be worst case. Ultimate Healthcare Ltd Calmore Industrial Estate, Nutwood Way, Totton Southampton, Hampshire, SO40 3WW Tel: 0333 321 8996 Fax: 023 8066 2388 Email: info@ultimatehealthcare.co.uk www.**ultimatehealthcare**.co.uk

