



# NOCOSPRAY 2

User manual



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# 1. INTRODUCTION

## 1.1 General information

Nocospray 2 is a transportable device for the diffusion of a disinfectant.

This device is intended for professional use only. It can be used in hospitals, the food industry, the pharmaceutical industry or any environment where disinfection is required.

This user manual describes the operation of the NOCOSPRAY 2 diffuser and includes essential information guaranteeing safe use of the product.

The documentation refers to NOCOSPRAY 2.0 diffusers in their original state and which have not been modified.

Read this manual before using the NOCOSPRAY® 2 diffuser. It has been written to ensure that you master all the features.



- Read this manual carefully before using the NOCOSPRAY 2 diffuser. It has been created so that you can use all of its functions effectively.
- Illustrations are provided to help you understand the various functions of the NOCOSPRAY 2. They are not binding.

## 1.2 Classification/Regulations

The NOCOSPRAY 2 diffuser complies with the following standards:

Electrical classification IEC 60950-1 / 60417-5019 standard	Electrical protection Class I	
Electromagnetic compatibility (EMC) Standard EN 60601-1-2 (2007)	The product does not emit electromagnetic interference that could affect other devices. It has appropriate immunity to electromagnetic interference.	

## 1.3 Warranty

The guarantee certificate must be presented to the retailer or to the company OXY'PHARM - Z.A. des Grands Godets - 829, rue Marcel Paul – 94508 Champigny-sur-Marne, France, in order to provide proof of the guarantee.

### Guarantee Clauses

The following provisions do not exclude the benefit of the legal guarantee provided by articles 1641 et seq. of the French Civil Code relating to hidden defects.

### Purpose of the guarantee

The guarantee consists of the gratuitous provision of parts which have been recognised as defective by our technical services.

Repairs made as a result of incorrect handling, abnormal use, negligence or the overloading of the device, as well as those required by variations in the power supply network, voltage surges or defective installations, etc., cannot be covered by the guarantee.

Furthermore, if any abnormal corrosion is observed on the flexible parts, hoses, joints, etc., indicating the use of a liquid other than those produced by OXY'PHARM, the guarantee shall also be rendered null and void.

### Transportation

The equipment is to be transported at the user's own risk: in the event of damage occurring during transport, the receiving party must communicate any reservations to the carrier before accepting the delivery of the device.

The device must be stored in a dry location and transported in its original packaging.

### Guarantee period

The NOCOSPRAY 2 diffuser is guaranteed for a period of 2 years, effective as of the date of the invoice.

## 2. SAFETY INFORMATION

The manufacturer is in no case liable for any incident affecting the equipment or a person caused by incorrect or abnormal use of the device in violation of the recommended precautions in the following non-exhaustive list:

	<p><i>To avoid any risk of electrical shock, the device must be connected to an electrical power supply network equipped with a protective earthing system.</i></p> <p><i>Ensure that the electrical power supply is compliant with the standards in force.</i></p>
	<p><i>This device is intended solely for professional use.</i></p> <p><i>This appliance may be used by children of 8 years old or over and by individuals with reduced physical, sensory or mental capabilities or who lack experience or knowledge, if they are properly supervised or if they have been given instructions on how to use the appliance safely and if the risks involved have been understood. Children must not play with the appliance. User cleaning and maintenance must not be carried out by unsupervised children.</i></p> <p><i>Any modification to the EM device is strictly prohibited.</i></p> <p><i>Portable and mobile RF communication devices may affect EM devices.</i></p> <p><i>According to the requirements of the standard NF EN 60601-1, the device may cause radio-electric disturbances and/or disturb the operation of a nearby device. The use of accessories, transducers and cables other than those specified, with the exception of equipment sold by the manufacturer as replacement parts of internal components, may result in an increase in emissions or a reduction in the device's electromagnetic immunity.</i></p> <p><i>The device must not be used next to or stacked with other devices. If this condition cannot be respected, test to ensure correct operation in this configuration.</i></p> <p><i>Portable radio frequency communication devices (use of remote control) may affect other electrical equipment and devices and should not be used within 30 cm of Nocospray 2.</i></p> <p><i>To avoid any risk of electrical shock, this appliance must not be connected to an electrical power supply equipped with a protective earthing system.</i></p> <p><i>Ensure that the electrical power supply is compliant with the usual standards.</i></p> <p><i>If the power cable is damaged, it must be replaced by the manufacturer, their after-sales service or similarly qualified persons in order to avoid a hazard.</i></p>
	<p><i>Use only with products from the NOCOTECH range.</i></p>
	<p><i>Do not insert any object or liquid into the nose of the diffuser whether it is switched on or off.</i></p> <p><i>Check for the presence of the jet and the jet support (reference 4 on page &lt;AU&gt;).</i></p> <p><i>To avoid any risk of accidental spillage of product inside the machine, the cylinder must be removed from the diffuser when it is being transported.</i></p> <p><i>Handle the tank with care.</i></p>
	<p><i>The user is not authorised to replace any part of the device.</i></p> <p><i>In the event of a fault, call your retailer/approved technician.</i></p>
	<p><i>Any electrical work carried out on the device must be performed by an approved technician. OXY'PHARM provides retailers with all technical information required for troubleshooting.</i></p>

## INSERT THE CONTACT DETAILS OF YOUR RETAILER HERE

Mr(s) .....

Address .....

Telephone No .....

The device diffuses a chemical substance. As a consequence, the following recommendations must be respected:

	<p><i>Users are required to read the accompanying documents. The decontamination process should be monitored.</i></p>
	<p><i>The device must only be used indoors.</i></p>
	<p><i>Do not inhale the product.</i></p>
	<p><i>Do not smoke near the device</i></p>
	<p><i>Do not direct the device towards an incandescent area.</i></p>
	<p><i>Do not use in the presence of an inflammable anaesthetic mixture.</i></p>
	<p><i>The decontamination process should be monitored. Evacuate the room during diffusion. Respect the contact times specified in the usage protocols. Respect a minimum delay of 30 minutes after treatment before returning into the treated area. In case of emergency, if a person has to enter the treated area, they must be equipped with Personal Protective Equipment: ear muffs and PPE according to the SDS of the product used. To ensure the safety of personnel not protected by PPE, an H2O2 detector can be used to measure any residual gas. The room may only be entered when the concentration (ppm) of H2O2 is below the levels required by local laws on permitted exposure times in the workplace.</i></p>

### 3. DESCRIPTION

#### 3.1 Description of the NOCOSPRAY 2



- 1 Handle
- 2 Diffuser release button
- 3 Diffuser nose (venturi)
- 4 Jet and support jet
- 5 Bottle
- 6 Control keypad
- 7 Power switch
- 8 Fuses
- 9 Power cable plug
- 10 Mini-USB Port

#### 3.2 Overall dimensions



## 4. USE

### 4.1 Positioning of the device

The Nocospray should be placed on a flat, sturdy surface, ideally in the corner of the room to be treated and raised off the ground if possible.



*Before beginning a treatment, check that there is no obstacle within 1.5 metres of the nozzle tip.*

### 4.2 Fitting the cartridge

The installation of the bottle is carried out in the manner below.

Consider dating the product bottle before installing it onto the NOCOSPRAY 2.



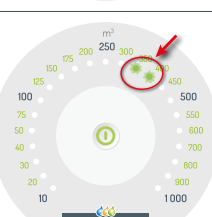
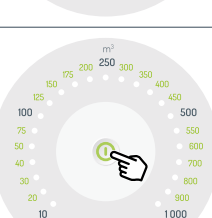
*Always check that the bottle contains a sufficient amount of product for the desired treatment (the device does not indicate this).*

*Please refer to the attached table on page <AU>.*

Steps	Action	Illustration
1	Remove the diffuser nose by pressing the release button.	
2	Screw the product bottle (after dating it) onto the diffuser nose.  Do not screw it too tight in order to avoid irreversibly damaging the joint.	
3	Place the diffuser nose back onto the device.  You should hear a "click".	

## 4.3 Operation in standard mode

### 4.3.1 Use without remote control

Steps	Action	Illustration
1	Plug in the NOCOSPRAY 2 power cable.  Set the power switch to "1".	
2	Select the volume to be treated by sliding your finger over the pale part of the keypad (*).  The selection range is between 10 m³ and 1 000 m³.  <i>(* In standard mode, the volume selected by default is 10 m³.</i>	
3	The level is progressive: if two LEDs are lit, the intermediate value is selected.  For example, if LEDs 10 and 20 are illuminated, the selected volume is 15 m³.	
4	To launch the device, press the button at the centre of the key pad (press and hold).	
5	A beep will sound for 15 seconds before the machine begins to function.  Leave the room before the end of the 15 seconds.  At the end of the diffusion process, there is a beep.  The device will stop automatically.	

	<p>If necessary, the device may be stopped during treatment by pressing the button at the centre of the keypad again.</p> <p><i>Respect the safety information provided on page &lt;EX&gt;.</i></p>	
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### 4.3.2 Use with remote control

The remote control allows the device to be launched from outside of the room.

#### 4.3.2.1 Initial pairing with the remote control

Before using the remote control for the first time, it needs to be paired with the NOCOSPRAY 2 device. Follow the steps below:

Steps	Action	Illustration
1	Plug in the NOCOSPRAY 2 power cable. Set the power switch to "0".	
2	Press the button on the remote control (approx. 5 seconds) in direct proximity to the NOCOSPRAY 2 with which you wish to pair it, hold the button until the LED on the remote control begins to blink.	
3	Keep the button on the remote control held down. Turn on the NOCOSPRAY 2 to be paired by setting the power switch to "1".	

Steps	Action	Illustration
4	The LED on the remote control will begin to blink rapidly.  The paired NOCOSPRAY 2 will beep several times to signal that the pairing has been made.	

#### 4.3.2.2 Re-pairing a remote control

The remote control can only be paired with one device at a time.  
However, a remote control which has already been paired can be paired with another device.

Steps	Action	Illustration
1	Ensure that all NOCOSPRAY 2 devices in proximity to the remote control are switched off.  Switches set to "0".	
2	Plug in the power cable of the new NOCOSPRAY 2 to be paired.	
3	Press and hold the button on the remote control (approx. 30 seconds directly next to the NOCOSPRAY 2 to be paired).	
4	For 30 seconds, the LED of the remote control will blink, then it will stop before beginning to blink again.  This phase is repeated 3 times.	

Steps	Action	Illustration
5	Once the LED of the remote control blinks regularly and without stopping, turn on the NOCOSPRAY 2 to be paired using its power switch whilst holding down the button on the remote control.	
6	The LED on the remote control will begin to blink rapidly.  The paired NOCOSPRAY 2 will beep several times to signal that the pairing has been made.	

#### 4.3.2.3 Operation with remote control

Steps	Action	Illustration
1	Plug in the NOCOSPRAY 2 power cable.  Set the power switch to "1".	
2	Select the volume to be treated by sliding your finger over the pale part of the keypad.  The selection range is between 10 m³ and 1 000 m³.	
3	The level is progressive: if two LEDs are lit, the intermediate value is selected.  For example, if LEDs 10 and 20 are illuminated, the selected volume is 15 m³.	

Steps	Action	Illustration
4	Press the button on the remote control to launch\treatment.	
5	A beep will sound for 15 seconds before the machine begins to function.  Leave the room before the end of the 15 seconds.  AT the end of the diffusion process, there is a beep.  The device will stop automatically.	
	If necessary, the device may be stopped during treatment by pressing the button on the remote control again.  <i>Respect the safety information provided on page &lt;EX&gt;.</i>	

## 4.4 Operation in programmer mode

### 4.4.1 Modus operandi

It is possible to use the NOCOSPRAY 2 with a programmer (not provided) in order to differ or repeat treatments. Follow the procedure outlined below to select the programmer mode.

Steps	Action	Illustration
1	Plug in the NOCOSPRAY 2 power cable.  Set the power switch to "1".	

Steps	Action	Illustration
2	Turn your finger in a clockwise direction on the pale part of the keypad up to the value 1000, then carry out at least 3 supplementary rotations.	
3	<p>The programmer mode is engaged with 3 LEDs are illuminated in the following manner:</p> <ul style="list-style-type: none"> <li>the LEDs 100 m<sup>3</sup> and 500 m<sup>3</sup> are illuminated,</li> <li>the LED 250 m<sup>3</sup> is blinking.</li> </ul>	
4	<p>Select the desired volume by turning in an anticlockwise direction (*).</p> <p>(* In programmer mode, the volume selected by default is 1000 m<sup>3</sup>. The selected volume is displayed only when you touch the keypad.)</p>	
5	<p>Set the power switch to "0".</p> <p>Unplug the NOCOSPRAY 2.</p>	
6	<p>Connect the NOCOSPRAY 2 to the programmer.</p> <p>Plug the programmer into a mains socket.</p> <p>Set the power switch to "1".</p>	
7	<p>Refer to the programmer's notice to program it.</p> <p>NB: It is possible to program it beforehand.</p>	

Steps	Action	Illustration
8	<p>The programmer will launch and stop the NOCOSPRAY 2 according to the performed programming.</p> <p>A beep will sound for 15 seconds before the machine begins to function.</p> <p>Leave the room before the end of the 15 seconds.</p> <p>At the end of the diffusion process, there is a beep.</p> <p>The device will stop automatically.</p>	 

#### 4.4.2 Special features of programmer mode

	<p><i>In programmer mode, the device cannot be stopped by pressing the centre of the keypad.</i></p> <p><i>In case of emergency, set the power switch to "0" or unplug the device.</i></p>	
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When the treatment in programmer mode is finished, it is highly recommended to configure the device in standard mode before switching it off.

Refer to § 4.4.3 below.

	<p><i>If the device is in programmer mode and you switch the device off and on again, you have 1 minute to change the volume or return to standard mode.</i></p> <p><i>At the end of this minute, the device will begin to beep in order to signal the beginning of the treatment.</i></p> <p><i>Leave the room before the end of the 15 seconds.</i></p>	 
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Steps	Action	Illustration
1	<p>At the end of treatment, if the device is still turned on, the volume may be modified.</p>	

#### 4.4.3 Return to standard mode

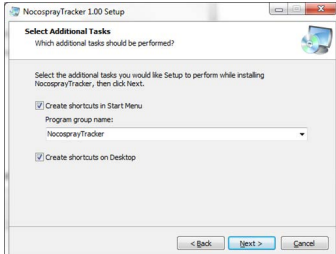
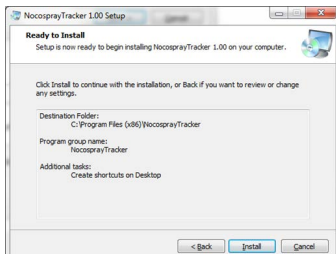
Steps	Action	Illustration
1	<p>To exit programmer mode and return to standard mode: Turn your finger in a clockwise direction on the pale part of the keypad up to the indication 10, then carry out at least 3 supplementary rotations.</p>	 <p>m<sup>3</sup></p> <p>OXY PHARM®</p>

## 5. TRACKING SYSTEM

The tracking system records the 125 latest treatments performed by the NOCOSPRAY 2. The 126<sup>th</sup> treatment erases the 1<sup>st</sup> and so forth.

### 5.1 Software installation

- Insert the CD Rom or the USB key provided into your PRC and open the folder named "Nocospray Tracker".
- Launch "NocosprayTracker\_Setup.exe".
- Authorise the software installation if Windows asks you to.

Steps	Action	Illustration
1	Click "Next".	
2	Click "Next" again.	
3	Click "Install".	

Steps	Action	Illustration
4	Click "Finish".	
5	If the following window appears, click "Close".	
6	The icon opposite may be displayed to indicate that the peripheral driver has been correctly installed.	
7	The software "NocosprayTracker" is launched automatically after installation.	

If your computer does not allow automatic installation, seek advice from your IT manager.

## 5.2 History transfer

Steps	Action	Illustration
1	Plug the USB cable into the back of the NOCOSPRAY 2 and into the USB Port of your computer (*).	
2	During data transfer: <ul style="list-style-type: none"><li>the LEDs 10 and 1000 blink,</li><li>the LED 250 is illuminated.</li></ul>	

(\*) No treatment may be performed whilst the USB cable is plugged in.

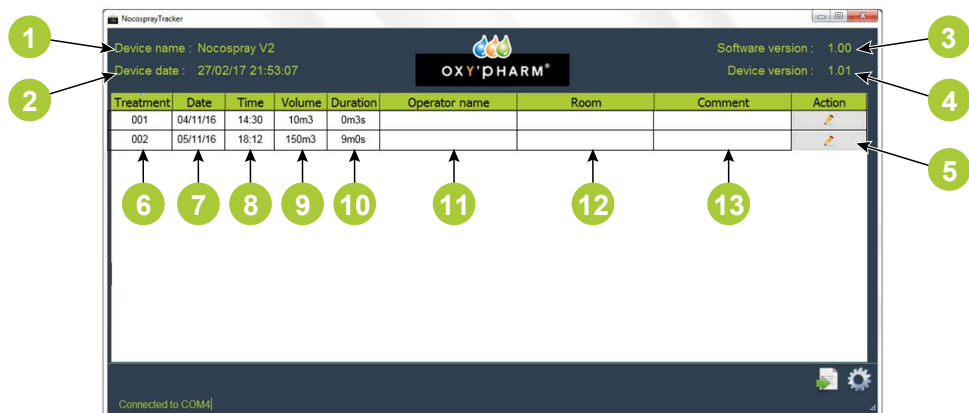
It is recommended to keep the USB cable connected during data modification  
(see § 5.3).

## 5.3 History display

### 5.3.1 Screen description

Open the software "NocosprayTracker" (in the programmes on your computer). The screen below shows an example of a history. Each line of the table represented a treatment.

Note: The indication in the lower left, "Connected to COM4", indicates that the NOCOSPRAY 2 is connected.



- 1 Device name (modifiable parameter)
- 2 Device date and time
- 3 Nocospray Tracker software version
- 4 Nocospray internal software version
- 5 Input and "Operator name", "Room" and "Comment" parameters (\*)
- 6 Treatment number
- 7 Treatment date
- 8 Treatment time
- 9 Nocospray setting volume for the desired treatment
- 10 Treatment duration (this parameter is calculated automatically, not measured)
- 11 Operator name
- 12 Name of the room in which the treatment was performed
- 13 Other comments

It is possible to class treatments according to the number, date, time, volume or the duration, operator name, room name or comments.

(\*) Once the "Operator name", "Room" and "Comment" parameters have been entered and validated, it is no longer possible to change them.

### 5.3.2 Modification of general settings

Steps	Action	Illustration
1	To modify the general settings, click on the  icon at the bottom of the screen.	
2	The following window will be opened:  This window allows the modification of the device name and the date and time of the device.	
3	Proceed to modifications.	
4	Click "Save".	

### 5.3.3 Export the history in PDF format

Steps	Action	Illustration
1	To export the history in PDF format, click on the  icon at the bottom of the screen.	
2	The following window will be opened:  Click on the  icon to select the folder in which you want to save the history file.	
3	Enter a file name in the "PDF Name" field.	
4	Next, click on "Export" to save the file.	

### 5.3.4 Exit USB mode

Steps	Action	Illustration
1	To exit USB mode, you simply have to: <ul style="list-style-type: none"><li>• Unplug the USB cable from the computer and the device (*)</li><li>• Close the software.</li></ul>	

(\*) No treatment may be performed whilst the USB cable is plugged in.

## 6. APPENDICES

### 6.1 Duration of diffusion and consumption

Volume to be treated (m <sup>3</sup> )	Consumption (ml)	Duration of diffusion (hh:mm:ss)
10	10	00:00:36
15	15	00:00:54
20	20	00:01:12
25	25	00:01:30
30	30	00:01:48
35	35	00:02:06
40	40	00:02:24
45	45	00:02:42
50	50	00:03:00
62.5	62.5	00:03:45
75	75	00:04:30
87.5	87.5	00:05:15
100	100	00:06:00
112.5	112.5	00:06:45
125	125	00:07:30
137.5	137.5	00:08:15
150	150	00:09:00
162.5	162.5	00:09:45
175	175	00:10:30
187.5	187.5	00:11:15
200	200	00:12:00
225	225	00:13:30
250	250	00:15:00
275	275	00:16:30
300	300	00:18:00
325	325	00:19:30
350	350	00:21:00
375	375	00:22:30
400	400	00:24:00
425	425	00:25:30
450	450	00:27:00
475	475	00:28:30
500	500	00:30:00
525	525	00:31:30
550	550	00:33:00
575	575	00:34:30
600	600	00:36:00
650	650	00:39:00
700	700	00:42:00
750	750	00:45:00
800	800	00:48:00
850	850	00:51:00
900	900	00:54:00
950	950	00:57:00
1000	1000	01:00:00

## 6.2 Maintenance



*Before any work is carried out, switch off the device and unplug the mains power cable.  
Do not use abrasive products, bleach, acetone or solvents for maintenance or disinfection of the device. These aggressive products risk damaging the device's surface materials.*

To clean its external surfaces, use a slightly moist cloth or wipe.

## 6.3 Technical specifications

Transport and storage conditions		
<b>Temperature</b>	0 °C to 70 °C	
<b>Hygrometry</b>	10% to 95% RH without condensation	
<b>Atmospheric pressure</b>	700 to 1060 hPa	
Usage conditions		
<b>Temperature</b>	0 °C to 40°C	
<b>Hygrometry</b>	10% to 90% RH without condensation	
<b>Atmospheric pressure</b>	800 to 1060 hPa	
General characteristics		
<b>Device weight</b>	5.9 kg	
<b>Package weight</b>	8.6 kg	
<b>Compatible packaging</b>	1 l	
<b>Maximal treatment volume</b>	1000 m <sup>3</sup>	
<b>Average liquid flow rate</b>	1000 ml/h	
Electrical characteristics		
<b>Power supply voltage</b>	230 V single-phase	120 V single-phase
<b>Power supply frequency</b>	50 to 60 Hz	50 to 60 Hz
<b>Power</b>	1000 W	1000 to 1100 W
<b>Nominal intensity</b>	4.5 A	9 A
<b>Fuse</b>	5x20 mm 10 A Time-delay	

## 7. EMC compliance table

### 7.1 Cable length

Cables & accessories	Length max.	Type of test	In accordance with
Cables / Cords	<3 m	RF emission	CISPR 11, Class B
		Harmonic currents emission	IEC 61000-3-2
		Voltage fluctuation and flickers	IEC 61000-3-3
		Electrostatic discharge immunity	IEC 61000-4-2
		Radiated immunity Electromagnetic fields	IEC 61000-4-3
		Immunity to interference against fast transient electrical disturbance/burst	IEC 61000-4-4
		Immunity to fast transients	IEC 61000-4-5
		Conducted immunity RF conduct interference	IEC 61000-4-6
		Radiated immunity Magnetic fields	IEC 61000-4-8
		Immunity to voltage drops, short interruptions and voltage variations	IEC 61000-4-11

### 7.2 Minimum recommended distance

The medical device is intended for be used in an electromagnetic environment with controlled interference from RF radiation.

The user or the installer of the medical device can work to avoid electromagnetic interference by maintaining a minimum distance, which depends on the maximum power of the radio frequency transmission equipment. Portable RF communication devices (including peripheral device such as antenna cables and external antennas) should not be used closer than 30 cm (12 inches) to any part of the appliance, including cables specified by the manufacturer, as this may impair the performance of these devices.

### 7.3 Electromagnetic emission

Test emission	Standard	Electromagnetic environment
Electromagnetic radiation disturbances. (Radiated emission) (CISPR 11)	Group 1	The Nocospray 2 uses RF energy only for its internal function.
Terminal supply disturbance voltage (Conducted emissions) (CISPR 11)	Class B	The Nocospray 2 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.
Harmonics emissions (IEC61000-3-2)	N/A	
Voltage / Flicker test (IEC 61000-3-3)	N/A	

## 7.4 Electromagnetic and magnetic immunity

Immunity test	IEC60601 test level	Compliance level	Electromagnetic environment
Electrostatic discharges (IEC61000-4-2)	± 8Kv contact ± 15Kv air	± 8Kv contact ± 15Kv air	Home health environment and a professional health establishment
Electrical fast Transients IEC61000-4-4)	± 2Kv for power supply lines	± 2Kv for power supply lines ± 1Kv for input/output lines	Home health environment and a professional health establishment
Surge (IEC61000-4-5)	± 1Kv line(s) to line(s) ± 2Kv line(s) to earth	± 1Kv line(s) to line(s) ± 2Kv line(s) to earth	Home health environment and a professional health establishment
Magnetic field at power frequency (IEC-61000-4-8)	30A/m	30A/m	Home health environment and a professional health establishment
Voltage dips, short interruptions and voltage variations on power supply input lines (IEC61000-4-11)	0% UT for 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270°, 315°.  0% UT for 1 cycle  70% UT For 25 cycles at 50Hz For 30 cycles at 60Hz Single-phase at 0°	0% UT for 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270°, 315°.  0% UT for 1 cycle  70% UT For 25 cycles at 50Hz For 30 cycles at 60Hz Single-phase at 0°	Home health environment and a professional health establishment
Voltage interruptions (IEC61000-4-11)	0% UT For 250 cycles at 50Hz For 300 cycles at 60Hz	0% UT For 250 cycles at 50Hz For 300 cycles at 60Hz	Home health environment and a professional health establishment

## 7.5 Electromagnetic immunity, RF

Immunity test	IEC60601 test level	Compliance level	Electromagnetic environment
RF portable communications equipment (including peripherals such as antenna cables and external antennas) should not be used closer than 30cm (12 inches) to any part of the UNIT TEST, including the cables specified by the manufacturer. Otherwise, the performance of these devices may be impaired.			
Radiated RF (IEC61000-4-3)	3V/m 80MHz to 2,7 GHz 80% MA at 1kHz  10v/m 80MHz to 2,7 GHz 80% MA at 1 kHz	3V/m 80MHz to 2,7 GHz 80% MA at 1kHz  10v/m 80MHz to 2,7 GHz 80% MA at 1 kHz	Home health environment and a professional health establishment
Proximity fields issued by RF wireless communication devices (IEC 61000-4-3 provisory method)	9v/m 710MHz, 745MHz, 780MHz, 5240MHz, 5550MHz, 5785MHz  27V/m 385MHz  28V/m 450MHz, 810MHz, 870MHz, 930MHz, 1720MHz, 1845MHz, 1970MHz, 2450MHz	9v/m 710MHz, 745MHz, 780MHz, 5240MHz, 5550MHz, 5785MHz  27V/m 385MHz  28V/m 450MHz, 810MHz, 870MHz, 930MHz, 1720MHz, 1845MHz, 1970MHz, 2450MHz	Home health environment and a professional health establishment
Conducted RF	3V 150KHz to 80MHz  6V in ISM band and band between 0.15 MHz and 80 MHz including amateur radio band.  80% MA at 1Khz	3V 150KHz to 80MHz  6V in ISM band and band between 0.15 MHz and 80 MHz including amateur radio band.  80% MA at 1Khz	Home health environment and a professional health establishment

## **8. RECYCLING**

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### **IMPORTANT INFORMATION FOR ENVIRONMENTALLY SOUND DISPOSAL**

This product is subject to the separate collection of electrical and electronic equipment in compliance with Directive 2002/96/EC (applicable in member countries of the European Union and other European countries with selective collection systems).



This symbol on the device indicates that at the end of its lifespan, the device should be disposed of separately from household waste. The device should be entrusted to a selective sorting centre for electrical and electronic devices, or returned to the retailer when purchasing a new device.

The user is responsible for the return of the device at the end of its lifespan to the appropriate sorting organisation or to a duly authorised company, under penalty of the sanctions provided for by legislation concerning waste disposal.

The selective sorting performed before recycling and the processing and disposal of the used device in an environmentally sound manner help avoid harmful consequences for the environment and health, and encourage the recycling of the materials that the device is made from.

The disposal of the used consumables must be managed in compliance with local regulations pertaining to waste management.







## OXY'PHARM

829, rue Marcel Paul – ZA des Grands Godets  
94500 Champigny sur Marne  
FRANCE

Tel. +33 145 187 870  
Email info@oxypharm.net



[www.oxypharm.net](http://www.oxypharm.net)