



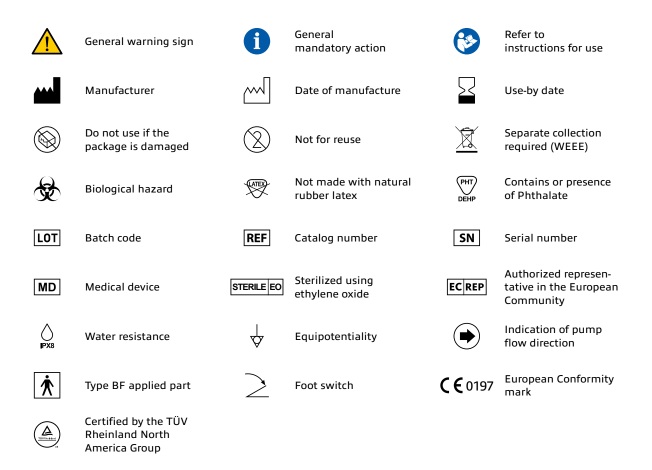


CONGRATULATIONS ON YOUR PURCHASE OF A PRODUCT FROM NOUVAG.

We are pleased that you have chosen a quality product from NOUVAG and thank you very much for the trust you have placed in us.

These instructions for use will familiarize you with the device and its functions so that you can apply and use them correctly.

SYMBOLS



CONTENT

INTENDED PURPOSE	4
Medical indications	
Contraindications	
Side effects	
Intended users	
Target population	
Ambient conditions	
SAFETY INFORMATION	5
Modifications and misuse	
Essential requirements	
During use	
SCOPE OF DELIVERY	6
DEVICE OVERVIEW	7
Front view	
Rear view	
SETUP	8
Device and accessories setup	
Connection to the power supply	
Potential equalization connection according to DIN 42801	
Device preparation	
OPERATION	10
Switching the device on and off	
Regulation of the infiltration process	
Peristaltic pump	
Functional check	
CLEANING AND DISINFECTION	10
Control unit and foot switch	
Tubing set REF 6022	
MAINTENANCE	11
Replacing the control unit fuses	
Safety inspections	
MALFUNCTIONS AND TROUBLESHOOTING	12
ACCESSORIES AND SPARE PARTS	13
Information on disposal	
TECHNICAL DATA	13
WARRANTY COVERAGE	14
Post market surveillance	
Service points	
APPENDIX	15

INTENDED PURPOSE

MEDICAL INDICATIONS

The Dispenser DP 30 LipoPlus is an infiltration pump, that is used in combination with a foot pedal and a sterile single use tubing set (independent medical devices) to infiltrate tumescent solution (Klein solution) in the subcutaneous fat layer. The Dispenser DP 30 LipoPlus is used in the following medical indication:

// Liposuction

CONTRAINDICATIONS

General contraindications concern patient health status, such as severe cardiopulmonary disease, local inflammation, sepsis, and coagulation disorders.

Specific contraindications include:

Infectious wounds Liposuction may only be performed after the treatment of the infection and necrotic tissue. **Liposuction shortly after a strict diet of the patient**.

Morbid obesity (obesity) Large suction volumes increase the risk of death due to fluid shifts.

Intravascular infusion of liquids.

Infiltration of excessive volume of tumescent solution (mutiple liters).

Treatment of a excessive surface area.

SIDE EFFECTS

Side effects of infiltration of tumescent solution for liposuction are generally infrequent and minor.

Clinical literature reports the following side effects:

- Inflammation (local/systemic)
- Bleeding
- Haematomas
- Oedema
- Fat/Pulmonary embolysm

INTENDED USERS

The device is designed to be used by professional and trained users only, in professional contexts (e.g. hospital, clinic). The device is not to be used by patients or by untrained users.

TARGET POPULATION

Adults (>18 years old). See contraindications for limiting health factors.

AMBIENT CONDITIONS

	TRANSPORT AND STORAGE	DURING USE
Relative humidity	max. 90%	max. 80%
Temperature	0°C-60°C	10°C-30°C
Atmospheric pressure	700–1'060 hPa	

SAFETY INFORMATION

Every use of the Dispenser DP 30 LipoPlus different to the [INTENDED PURPOSE >4] causes risks for patients and trained personnel. If physical examinations and therapies are carried out without use of the devices, then the devices must be removed from the place of treatment. Avoid any connection or close adjacency to other devices.

MODIFICATIONS AND MISUSE

Modifications/manipulations on the Dispenser DP 30 LipoPlus and its accessories are not permitted. Failure to follow these instructions can have unpredictable consequences for the user, the patient or third parties. For consequential complications, resulting from illicit modifications/manipulations the manufacturer assumes no responsibility and the guarantee is void.

NOUVAG recommends the use of Klein tumescent anesthesia solution. The use of other solutions is on the responsibility of the surgeon. When infiltrating tumescent anesthesia solution, do not exceed 0.05% w/w anesthetic concentration.

ESSENTIAL REQUIREMENTS

Do not use the device if the shipping box has holes/cracks on the flat surfaces, and/or if the Styrofoam protective packaging is broken.

The Dispenser DP 30 LipoPlus may only be operated by qualified and trained personnel!

The use of third-party products is the responsibility of the operator. Functionality and patient safety cannot be guaranteed with third-party accessories.

Repairs may only be performed by authorized NOUVAG service technicians!

Improper use or repair of the device, or failure to observe these instructions, relieves NOUVAG from any obligation arising from warranty provisions or other claims.

Prior to using the device, before startup, and before operation, the user must always ensure that the device and accessories are in good working order and are clean, sterile and operational.

Ensure that the operating voltage setting corresponds to the local mains voltage.

The Dispenser DP 30 LipoPlus may only be operated under constant supervision of medical personnel. The absence of a warning buzz to indicate malfunctions of the device requires the permanent control of the volumetric displacement of the pump.

DURING USE

The device is not sterile on delivery. Please observe the instructions [CLEANING AND DISINFECTION >10].

Do not use device in the vicinity of flammable mixtures!

The use of the Dispenser DP 30 LipoPlus other than that for which it was designed (see [MEDICAL INDICA-TIONS >4]) is not permitted. The responsibility is solely carried by the operator.

SCOPE OF DELIVERY

REF	DESCRIPTION	QUANTITY
4181	Dispenser DP 30 LipoPlus control unit	1
6022	Disposable tubing set, sterile, 4m	1
1770	Stand for irrigation fluid bottle	1
31628	Dispenser DP 30 LipoPlus instructions for use	1

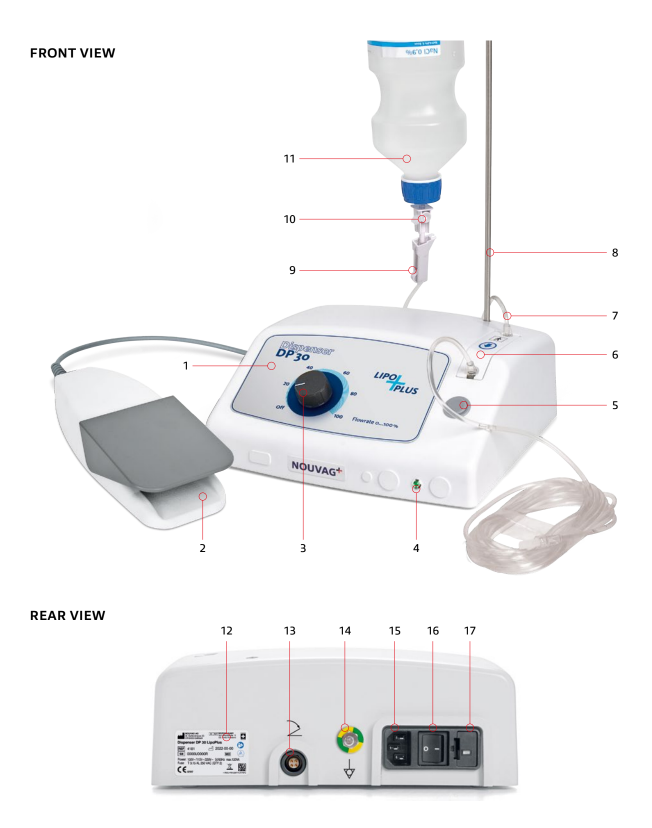
SELECTIVELY: SET NO. 4161 – DISPENSER DP 30 LIPOPLUS CONTROL UNIT WITH ON/OFF FOOT SWITCH

REF	DESCRIPTION	QUANTITY
1527nou	ON/OFF foot switch	1

SELECTIVELY: SET NO. 4163 - DISPENSER DP 30 LIPOPLUS CONTROL UNIT WITH VARIO FOOT SWITCH

REF	DESCRIPTION	QUANTITY
1511nou	VARIO foot switch	1

DEVICE OVERVIEW



1 Operating panel with pump displacement scale 2 Pedal 3 Control dial to set pump displacement volume 4 Indicator light for Power ON/OFF 5 Release key for tubing set bracket 6 Peristaltic pump 7 Tubing set 8 Stand for irrigation fluid bottle 9 Roller clamp 10 Venting valve 11 Irrigation fluid container 12 Type plate with type designation, reference number, serial number, information on power supply and device fuse 13 Pedal socket 14 Potential equalization 15 Power plug socket 16 Main switch 17 Fuse compartment

SETUP

DEVICE AND ACCESSORIES SETUP

- Place the Dispenser DP 30 LipoPlus and all required accessories and instruments on an even, non-slip surface and make sure you have good access to all controls.
- Do not allow the operating range of the device and accessories to be compromised by limiting factors.
- ¬ The system display must be always fully visible.
- ¬ It must be explicitly ensured that no objects can fall on the pedal.
- ¬ The power plug at the rear of the device must be always accessible.

CONNECTION TO THE POWER SUPPLY

To prevent of risks of an electric shock, the device may only be connected to a power network with a PE protective ground conductor!

If the voltage shown does not correspond to the local mains voltage, the grey fuse holder must be set to the correct voltage:



- 1 Unplug the power cable.
- 2 Use a screwdriver to open the fuse slot.
- 3 Remove the fuse holder.
- 4 Remove the grey fuse holder and reinsert it so that the local mains voltage setting is shown in the small window.
- 5 Slide fuse holder back in and close the fuse slot.
- 6 Check the mains voltage shown on the fuse slot.
- 7 Plug the power cable back into the device.

POTENTIAL EQUALIZATION CONNECTION ACCORDING TO DIN 42801

At the back of the device a potential equalization plug is installed, according to DIN 42801. The additional potential equalization has the task of equalizing potentials between different parts of conductive materials that can be touched at the same time, or reducing potential differences. This connection must be used, to protect the patient, the user and third parties from touch voltages.

The equipotential plug is marked with the following symbol: $\frac{1}{\nabla}$

SETUP

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DEVICE PREPARATION

- 1 Insert the stand for the irrigation fluid into the stand holder.
- 2 Plug the foot switch plug into the foot switch socket at the rear of the control unit. >
- 3 Assemble the tubing set (see images).
 - Check the expiry date of the tubing set and ensure that the packaging is not damaged. Using non-sterile tubing sets can result in serious infection and, in extreme cases, can cause death.
 - When inserting the tubing set, notice the arrow on the cover of the pump compartment. It indicates the flow direction of the cooling liquid.

Use only NOUVAG tube sets, otherwise the correct function cannot be guaranteed.

Do not regulate the amount of irrigation fluid using the roller clamp on the tube set; with the Dispenser DP 30 LipoPlus, this is regulated instead using the control dial and the foot pedal. For this reason,

make sure to open the roller clamp as far as it will go.

The container of infiltration fluid may weigh a maximum of 2 kg. Heavier containers can cause the device to tip over



- A Press the release key for tubing set bracket to open the pump.
- B The compartment with the integrated tubing bracket opens.
- C Place the tubing set into the tubing bracket provided in such a way that the end of the tubing set with the spike exits the pump to the rear of the control unit. Check that the tubing is secure.
- D With the tubing set inserted, press the compartment downwards until it clicks into place.



4 Insert the spike at the end of the tubing set into the infiltration fluid bottle and hang the bottle onto the stand.



5 Open the roller clamp on the tubing set as far as it will go.



- 6 Open the vent valve at the spike.
- 7 Connect the control unit to the power socket.

OPERATION

SWITCHING THE DEVICE ON AND OFF

The main switch «I/O» at the back of the unit is used to switch the control unit on and off. Switching off can be done at any time and is not dependent on a switch-off procedure.

The green LED light at the top left of the control panel lights up when the main switch has been activated and the unit is ready for operation.

REGULATION OF THE INFILTRATION PROCESS

Control dial in conjunction with ON/OFF foot switch The desired volumetric displacement is set with the control dial. The pumping process is started by actuating the ON/OFF foot switch. The volumetric displacement can be varied at any time using the control dial.

Control dial in conjunction with VARIO foot switch The maximum volumetric displacement can be varied at any time using the control dial, even while the foot switch is being pressed. Control using the VARIO foot switch regulates the volumetric displacement of the pump up to the set maximum value.

PERISTALTIC PUMP

Turn control dial clockwise from the OFF position. Pump starts, liquid emerges from the open tube end. Turning the dial up to the maximum value controls the increase in volumetric displacement. The pump stops immediately when the release button of the pump compartment is pressed.

FUNCTIONAL CHECK

Prior to Dispenser DP 30 LipoPlus startup or use of accessory equipment, the user must always ensure that each individual component is in good working order, free of defects, clean, sterile and operational. The tube set has to correspond with the correct flow direction and the pump has to function. The green LED is on after the device is switched on.

CLEANING AND DISINFECTION



Clean and disinfect the devices after every treatment!

CONTROL UNIT AND FOOT SWITCH

Wipe the outside using tested surface disinfectant or 70% of isopropyl alcohol. The front plate of the control unit is sealed and can be wiped clean.

TUBING SET REF 6022

Single-use tubing sets may not be reused!		
Used tubing sets must be disposed of properly.		
Don't use tube set when pack is already opened or damaged!		
Do not use tubing set if expired.		
Use only NOUVAG tubing sets with REF 6022.		
Sterility cannot be guaranteed by reusing and re-sterilization of tubing sets. The characteristics of the device		
may change resulting in serious infections or, in worst case, the death of the patient.		

MAINTENANCE

REPLACING THE CONTROL UNIT FUSES

Users can replace faulty control unit fuses themselves. These are located at the rear of the device in the fuse slot beside the power switch:

- 1 Switch off device.
- 2 Unplug the power plug.
- 3 Open the fuse slot using a screwdriver.
- 4 Replace the faulty fuse T 3,15 A, 250 V AC.
- 5 Slide the fuse holder back in and close the fuse slot.
- 6 Check the mains voltage shown on the fuse slot.
- 7 Plug in the power plug again.



1 Fuse slot locking mechanism 2 Display window for voltage setting 3 Fuse slot 4 Fuse 1 5 Fuse 2

SAFETY INSPECTIONS

The essential requirements have been defined and assessed within the risk analysis. The results of the analysis are stored in the risk management file of the manufacturer.

The performance of safety inspections on medical devices is required by law in several countries. The safety inspection is a regular safety check that is compulsory for those operating medical devices. The objective is to ensure that device defects and risks to patients, users or third parties are identified in time.

The STI (Safety Technical Inspection) for the Dispenser DP 30 LipoPlus shall be executed every 2 years by authorized experts. Results shall be documented. The service manual, wiring diagrams, and descriptions are available upon request from the manufacturer.

NOUVAG offers a safety inspection service for its customers. Addresses can be found in the appendix of this instructions for use under [Service POINTS >14]. For further information please contact our technical service department.

MALFUNCTIONS AND TROUBLESHOOTING

MALFUNCTION	CAUSE	SOLUTION	REFER TO INSTRUCTIONS FOR USE
Device is not functional (Indicator light is off)	Control unit not switched on	Set the power switch «I/O» to «I»	[Switching the device on and off >10]
	Power connection not established	Connect the control unit to the mains power supply	[CONNECTION TO THE POWER SUPPLY >8]
	Incorrect operating voltage	Check the mains voltage	[CONNECTION TO THE POWER SUPPLY >8]
	Faulty fuse	Replace fuse	[REPLACING THE CONTROL UNIT FUSES >11]
Pump doesn't work (Indicator light is on)	Infiltration quantity set too low or set to «OFF»	Raise pump performance by turning control switch up	[REGULATION OF THE INFILTRATION PROCESS >10]
	Tubing set incorrectly inserted	Insert tubing set correctly	[Device preparation >9]
	Incorrect operation	Check instructions for use	[DEVICE PREPARATION >9]
	Foot switch was not pressed	Press foot switch down, if infiltration process is controlled via the foot switch	[REGULATION OF THE INFILTRATION PROCESS >10]
	Roller clamp is closed	Open roller clamp all the way	[Device preparation >9]
Foot switch doesn't work (Indicator light is on)	Foot switch is not connected	Connect foot switch with the socket on rear of device	[Device overview >7] [Device preparation >9]
	Incorrect operation	Check instructions for use	[Device preparation >9] [Regulation of the infiltration process >10]

If the problem cannot be solved please contact your supplier or an authorized service center. Addresses can be found in the appendix of this instructions for use under [SERVICE POINTS > 14].

ACCESSORIES AND SPARE PARTS

ACCESSORIES

DESCRIPTION	REF
ON/OFF foot switch	1527nou
VARIO foot switch	1511nou
Stand for irrigation fluid bottle	1770
Disposable tubing set, sterile, 4 m	6022

To order any additional parts, please contact our customer service department.

INFORMATION ON DISPOSAL



Electrical and electronic devices that have reached the end of their service life comprise hazardous waste and may not be disposed of together with household waste. Prevailing national and local disposal regulations apply.

When disposing of the device, device components and accessories, the requirements specified in legislation must be followed. To ensure environmental protection, old devices can be returned to the dealer or manufacturer.

TECHNICAL DATA

Voltage, switchable	100 V~ / 115 V~ / 230 V~, 50 / 60 Hz
Fuse power supply	2 fuses, T 3,15A, 250 V AC
Power consumption	120 VA
Volumetric displacement	0–480 ml/min.
Maximum pressure with closed tube set	2,0 bar
Applied part	Type BF*
Protection class	Class I
Dimensions (W x D x H)	260 x 250 x 110 mm
Net weight control unit	3,4 kg
Maximum weight at the stand for the irrigation fluid bottle	2,0 kg

The mentioned volumetric displacement is only valid for aqueous solutions without any instrument connected.

* Applied part is the tube set with its attached instruments.

WARRANTY COVERAGE

NOUVAG warrants this product to be free from defects in workmanship and materials for a period of twelve (12) months from the original date of purchase. If the warranty card is returned for registration or the warranty extension is requested on our website within 4 weeks from the date of purchase, the warranty coverage is extended for a period of 6 months, wear parts are excluded from the warranty. During this warranty period, NOUVAG agrees to either repair or replace the product at its option if the product fails to function properly under normal use and service and such failure is due solely to a defect in workmanship or materials. This warranty is void if repair or service of the product is performed or attempted by anyone not authorized by NOUVAG to do so, or if a replacement part not authorized by NOUVAG is used in any repair or service.

POST MARKET SURVEILLANCE

In the event of incidents related to the use of the medical device, please contact immediately the manufacturer by email <u>complaint@nouvag.com</u> or by phone.
 To provide adequate information, please compile the incident questionnaire at the web address
 <u>Nouvag.com</u> > <u>Contact us</u> > <u>Incident questionnaire</u>.

SERVICE POINTS



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Phone +41 71 846 66 00 info@nouvag.com www.nouvag.com



A complete list of NOUVAG certified service points are found on the NOUVAG website: Nouvag.com > Service

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APPENDIX

Electromagnetic compatibility (EMC)

Remark:

The Product subsequently referred to herein always denotes the Dispenser DP 30 LipoPlus.

Changes or modifications to this product not expressly approved by the manufacturer may result in increased emissions or decreased immunity performance of the product and could cause EMC issues with this or other equipment. This product is designed and tested to comply with applicable regulations regarding EMC and shall be installed and put into service according to the EMC information stated as follows.

WARNING

Use of portable phones or other radio frequency (RF) emitting equipment, including accessories (antennas e.g.) in distances below 30 cm (12 inches) to the product, may cause unexpected or adverse operation.

WARNING

The product is suitable for use in hospitals other than in the vicinity of active devices of the HF surgical devices or except in HF screening rooms used for magnetic resonance imaging.

WARNING

The product shall not be used adjacent to, or stacked with, other equipment. If adjacent or stacked use is necessary, the product shall be tested to verify normal operation in the configuration in which it is being used.

Essential Performance

The essential performance is that the infiltration of tumescent solution in the fat tissue taking into account the infiltration flow rate and pressure is maintained. The maximum infiltration flow rate deviation is ± 25%, the infiltration flowrate is between 60 and 480ml/min and the maximum pressure is 2.5bar.

Compliant Cables and Accessories

WARNING

The use of accessories, transducers and cables other than those specified may result in increased emissions or decreased immunity performance of the product.

The table below lists cables, transducers, and other applicable accessories for which the manufacturer claims EMC compliance. **NOTE:** Any supplied accessories that do not affect EMC compliance are not listed.

Description	Length max.
Power supply cord REF 22261 / 22262 / 22264 / 22266	3.0m
Foot pedal IPX8 REF 1511nou / 1527nou	2.9m

Guidance and manufacturer's declaration – electromagnetic emissions			
	The Product is intended for use in the electromagnetic environment specified below. The customer or the user of the Product should assure that it is used in such an environment.		
Emissions test Compliance Electromagnetic environment - guidance			
RF emissions CISPR 11	Group 1	The Product 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 Product is suitable for use in all establishments, including domestic establishments and those directly connected to the	
Harmonic emissions IEC 61000-3-2	Class A	public low-voltage power supply network that supplies buildings used for domestic purposes.	
Voltage fluctuations/flicker emissions IEC 61000-3-3	complies		

Guidance and manufacturer's declaration – electromagnetic immunity					
	The Product is intended for use in the electromagnetic environment specified below. The customer or the user of the Product should assure that it is used in such an environment.				
Immunity tests	IEC 60601 Test level	Compliance level	Electromagnetic environment - guidance		
Electrostatic discharge (ESD)	+/- 8 kV contact	+/- 8 kV contact	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic		
IEC 61000-4-2	+/- 2 kV, +/- 4 kV, +/- 8 kV, +/- 15 kV air	+/- 2 kV, +/- 4 kV, +/- 8 kV, +/- 15 kV air	material, the relative humidity should be at least 30 %.		
Electrical fast transient/burst	+/- 2 kV with 100kHz for power supply lines	+/- 2 kV with 100kHz for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.		
IEC 61000-4-4	+/- 1 kV with 100kHz for input/output lines	+/- 1 kV with 100kHz for input/output lines			
Surge	+/- 0.5 kV, +/- 1 kV differential mode	+/- 0.5 kV, +/- 1 kV differential mode	Mains power quality should be that of a typical commercial or hospital environment.		
IEC 61000-4-5			1		
	+/- 0.5 kV, +/- 1 kV, +/- 2 kV	+/- 0.5 kV, +/- 1 kV, +/- 2 kV			
	common mode	common mode			

APPENDIX

Voltage dips, short	0 % U _{T;} for 0,5 cycle	0 % U _{T;} for 0,5 cycle	Mains power quality should bet hat of a typica
interruptions and voltage variations on power	with 0, 45, 90, 135, 180, 225, 270, 315 degree	with 0, 45, 90, 135, 180, 225, 270, 315 degree	·
supply input lines IEC 61000-4-11	0 % $U_{T;}$ for 1 cycle	0 % U _{T;} for 1 cycle	If the user of the Product requires continued operation during power mains interruptions, it is recommended that the Product be powered
	70 % U_T ; for 25/30 cycles	70 % U_T ; for 25/30 cycles	from an uninterruptible power supply or a battery.
	0 % U _{T;} for 5 sec	0 % U _{T;} for 5 sec	
Power frequency (50/60Hz) magnetic field IEC 61000-4-8	30 A/m	30 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
	voltage prior to application of the		unity for not life support equipment
The Product is intended for	r use in the electromagnetic env		customer or the user of the Product should
assure that it is used in su Immunity tests	ch an environment. IEC 60601	Compliance level	Electromagnetic environment - guidance
-	Test level		
			Portable and mobile RF communications equipment should be used no closer to any part of the Product, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
			Recommended separation distance:
Conducted RF IEC 61000-4-6	3 V rms 0.15 MHz to 80 MHz	3 V rms 0.15 MHz to 80 MHz	$d = 0.35 \sqrt{P}$
	6 V rms inside ISM bands between 150 kHz to 80 MHz	6 V rms inside ISM bands between 150 kHz to 80 MHz	
	80% AM bei 1 kHz	80% AM bei 1 kHz	
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.7 GHz	3 V/m 80 MHz to 2.7 GHz	$d = 0.35 \sqrt{P}$ 80 MHz to 800 MHz
	80% AM bei 1 kHz	80% AM bei 1 kHz	<i>d</i> = 0,7 \sqrt{P} 800 MHz to 2,7 GHz
			Where P is the maximum output power rating in the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m).
			Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ^a , should be less than the compliance level in each frequency range ^b .
			Interference may occur in the vicinity of equipment marked with the following symbol:
			(())
Note 1: At 80 MHz and	800 MHz, the higher frequency	range applies.	
Note 2: These guideline structures, obje		. Electromagnetic propagation	is affected by absorption and reflection from
a Fixed strengths amateur radio, a electromagnetic	from fixed transmitters, such as AM and FM radio broadcast and c environment due to fixed RF tra	l TV broadcast cannot be predi ansmitters, and electromagneti	r/cordless) telephones and land mobile radios, cted theoretically with accuracy. To access the c site survey should be considered. If the ne applicable RF compliance level above, the
Product should		ration. If abnormal performance	e is observed, additional measures may be

APPENDIX

Electromagnetic immunity against high-frequency wireless communication devices								
Test frequency MHz	Frequency band MHz	Communication service	Modulation	Maximum Performance W	distance m	Test level V/m		
385	380 to 390	TETRA 400	Pulse modulation 18 Hz	1.8	0.3	27		
450	430 to 470	GMRS 460, FRS 460	FM ± 5 kHz Hub 1 kHz Sinus	2	0.3	28		
710 745 780	704 to 787	LTE Band 13, 17	Pulse modulation 217 Hz	0.2	0.3	9		
810 870 930	800 to 960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse modulation 18 Hz	2	0.3	28		
1720 1845 1970	1700 to 1990	GSM 1800, CDMA 1900, GSM 1900, DECT, LTE Band 1, 3,	Pulse modulation 217 Hz	2	0.3	28		
2450	2400 to 2570	4, 25; UMTS Bluetooth, WLAN 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation 217 Hz	2	0.3	28		
5240 5500 8785	5100 to 5800	WLAN 802.11 a/n	Pulse modulation 217 Hz	0.2	0.3	9		

Recommended separation distances between portable and mobile RF communications equipment and the not life support equipment

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

	Separation distance according to frequency of transmitter					
Rated maximum output power	m					
of transmitter	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.5 GHz			
w	$d = 0.35 \sqrt{P}$	$d = 0,35 \sqrt{P}$	$d = 0.7 \sqrt{P}$			
0,01	0,04	0,04	0,07			
0,1	0,11	0,11	0,22			
1	0,35	0,35	0,7			
10	1,1	1,1	2,2			
100	3,5	3,5	7			
For transmitters rated at a maximum o	utput power not listed above, the r	ecommended separation distance d	in metres (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 higher frequency range applies.

 Note 1:
 At 80 MHz and 800 MHz, the separation distance fort the higher frequency range applies.

 Note 2:
 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.



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