# **\sonic**one O.R. Instructions For Use

SonicOne O.R. Powered by Misonix



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For additional information not contained in this manual, please visit www.misionix.com or contact your local sales representative.

# 1. General Safety Statements

- WARNING 1.1 The SonicOne O.R. system is an electro-mechanical device, which under certain circumstances could present an electrical shock hazard to the operator and/or patient. Please read manual thoroughly and follow directions stated herein to assure maximum safety during operation. This manual shall be kept in close proximity to the system for easy referral when needed.
- WARNING 1.2 The SonicOne O.R. system is intended to be used in various types of invasive, surgical procedures. There may be indirect danger to the patient should the device fail during the procedure. It is recommended that the facility follows its back-up equipment protocols.
- CAUTION 1.1 Special Skills and Training Requirements
  - 1. U.S. federal law restricts this device to sale by, or on the order of a licensed healthcare practitioner.
  - 2. The SonicOne O.R. system is to be used by an appropriately trained and licensed healthcare practitioner.

#### 1.1. EMC Statement

The SonicOne O.R. system is designed and tested to comply with FCC regulations for conducted and radiated emissions under Part 18 Subchapter J. and to comply with IEC EN60601-1-2:2007 guidelines for EMC.

- CAUTION 1.2 This device is considered medical electrical equipment. Medical electrical equipment needs special precautions regarding electromagnetic compatibility (EMC) and needs to be installed and put into service according to the EMC information provided in this operator's manual.
- WARNING 10.2 Portable and mobile RF communication equipment (including peripherals such as antennas) should be no closer than 30 cm (12 inches) to any part of the console, including the cables supplied with the console Otherwise degradation of the performance of this equipment could result.
- WARNING 10.3 The use of accessories, transducers and cables other than those specified or provided by Misonix may result in increased electromagnetic emissions or decreased immunity of the device and may result in improper operation. Use only Misonix branded equipment and accessories.
- WARNING 10.4 Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.

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## Electromagnetic Compatibility Guidance (in accordance with UL/EN/IEC 60601-1)

#### Guidance And Manufacturer's Declaration - Electromagnetic Emissions (Table 201) The SonicOne O.R SYSTEM is intended for use in the electromagnetic environment specified below. The customer or the user of SonicOne O.R SYSTEM should ensure that it is used in such an environment. **Emissions test** Compliance Electromagnetic environment – guidance The SonicOne O.R SYSTEM uses RF energy only for its internal RF emissions Group 1 function. Therefore, its RF emissions are very low and are not likely CISPR 11 to cause any interference in nearby electronic equipment. RF emissions Class A CISPR 11 The SonicOne O.R SYSTEM is suitable for use in all establishments Harmonic emissions Class A other than domestic and those directly connected to the public IEC 61000-3-2 low-voltage power supply network that supplies buildings used Voltage fluctuations/ for domestic purposes. flicker emissions Complies

Table 1.1 Guidance & manufacturer's declaration on electromagnetic emissions (EN table 201)

IEC 61000-3-3

NOTE: The EMISSIONS characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A).

NOTE: The emissions characteristics of this equipment make it suitable for use in industrial areas and hospitals. If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.

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## Guidance And Manufacturer's Declaration – Electromagnetic Immunity (Table 202)

The SonicOne O.R. SYSTEM is intended for use in the electromagnetic environment specified below. The customer or the user of the SonicOne O.R. SYSTEM should assure that it is used in such an environment.

IEC 60601 test level	Compliance level	Electromagnetic environment – guidance
o ±8 kV contact o ±2kV, ±5kV, ±8kV, ±15 kV air	o ±8 kV contact o ±2kV, ±5kV, ±8kV, o ±15 kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
<ul> <li>±0.5 kV, ±1 kV</li> <li>line to line</li> <li>±0.5 kV, ±1 kV, ±2 kV</li> <li>line to ground</li> </ul>	<ul> <li>±0.5 kV, ±1 kV</li> <li>line to line</li> <li>±0.5 kV, ±1 kV, ±2 kV</li> <li>line to ground</li> </ul>	Mains power quality should be that of a typical commercial or hospital environment.
o ±1 kV differential mode o ±2 kV common mode	o ±1 kV differential mode o ±2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.
<0 % U <sub>T</sub> (100 % dip in U <sub>T</sub> for 0,5 cycle 70 % U <sub>T</sub> (30 % dip in U <sub>T</sub> ) for 25 cycles 0 % U <sub>T</sub> (100 % dip in U <sub>T</sub> ) for 1 cycles 0 % U <sub>T</sub> (100 % dip in U <sub>T</sub> ) for 5 sec	0 % Uτ (100 % dip in Uτ) for 0,5 cycle 70 % Uτ (30 % dip in Uτ) for 25 cycles 0 % Uτ (100 % dip in Uτ) for 1 cycles 0 % Uτ (100 % dip in Uτ) for 5 sec	Mains power quality should be that of a typical commercial or hospital environment. If the user of the SonicOne O.R SYSTEM requires continued operation during power mains interruptions, it is recommended that the powered from an uninterruptible power supply.
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.
	test level  0 ±8 kV contact  0 ±2kV, ±5kV, ±8kV,  ±15 kV air  0 ±0.5 kV, ±1 kV line to line  0 ±0.5 kV, ±1 kV, ±2 kV line to ground  0 ±1 kV differential mode  0 ±2 kV common mode  <0 % UT (100 % dip in UT) for 25 cycles  0 % UT (100 % dip in UT) for 1 cycles  0 % UT (100 % dip in UT) for 5 sec	test level  0 ±8 kV contact  0 ±2kV, ±5kV, ±8kV,  ±15 kV air  0 ±0.5 kV, ±1 kV  line to line  0 ±0.5 kV, ±1 kV, ±2 kV  line to ground  0 ±1 kV differential  mode  0 ±2 kV common mode  <0 ±1 kV differential  mode  0 ±2 kV common mode  <0 % UT  (100 % dip in UT for  0,5 cycle  70 % UT  (30 % dip in UT) for 25  cycles  0 % UT  (100 % dip in UT) for 1  cycles  0 % UT  (100 % dip in UT) for 5  sec   Compliance level  0 ±8 kV contact  0 ±2 kV, ±5kV, ±8kV,  ±15 kV air  0 ±0.5 kV, ±1 kV  line to line  0 ±0.5 kV, ±1 kV, ±2 kV  line to ground  0 ±1 kV differential  mode  0 ±2 kV common  mode  70 % UT  (100 % dip in UT) for 25  cycles  0 % UT  (100 % dip in UT) for 1  cycles  0 % UT  (100 % dip in UT) for 5  sec

Table 1.2 Guidance & manufacturer's declaration on electromagnetic immunity (EN table 202)

List of Cables			
Item	Cable Length	Туре	
Handpiece cable	15 ft   4.6 m	shielded 2-conductor	
Power cord	10 ft   3.0 m	unshielded 3-conducter	
Footswitch cable	14 ft   4.3 m	shielded 2-conductor	

Table 1.3 List of cables

#### Guidance And Manufacturer's Declaration – Electromagnetic Immunity (Table 204)

The SonicOne O.R. SYSTEM is intended for use in the electromagnetic environment specified below. The customer or the user of the SonicOne O.R. SYSTEM should assure that it is used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment – guidance
			Portable and mobile RF communications equipment should be used no closer to any part of the SonicOne O.R SYSTEM, 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 Vrms 150 kHz to 80 MHz	3 V	d = 1.2√P
			d = 1.2VP 80 MHz to 800 MHz
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2. <b>7</b> GHz	3 V/m	d = 2.3vP 800 MHz to 2.7 GHz
			where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m).
			Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.
			Interference may occur in the vicinity of equipment marked with the following symbol:
			$((\bullet))$

NOTE 1 At 80 MHz and 800 MHz, 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.

Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the SonicOne O.R SYSTEM is used exceeds the applicable RF compliance level above, the SonicOne O.R SYSTEM should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the SonicOne O.R SYSTEM.

<sup>b</sup>Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Table 1.4 Guidance & manufacturer's declaration on electromagnetic immunity (EN table 204)

# Recommended Separation Distances Between Portable And Mobile RF Communications Equipment And The SonicOne O.R. System (Table 206)

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

Rated maximum output power	Separation distance according to frequency of transmitter m			
of transmitter W	150 kHz to 80 MHz d = 1.2√P	80 MHz to 800 MHz d = 1.2VP	800 MHz to 2,7 GHz d = 2.4VP	
0.01	0.12	0.12	0.24	
0.1	0.38	0.38	0.76	
1	1.2	1.2	2.4	
10	3.8	3.8	7.6	
100	12	12	24	

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.

NOTE 1 At 80 MHz and 800 MHz, the separation distance for 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.

Table 1.5 Recommended separation distances (EN table 206)

# 1.2. Electrical Safety Statement

The SonicOne O.R. system is designed and tested to comply with IEC 60601-1, UL 60601-1 and BSEN 60601-1.

- WARNING 1.3 The SonicOne O.R system generates high voltages within the console itself and the connected handpiece. To avoid injury, the console should never be operated before ensuring that its cover is properly closed and not tampered with. Do not attempt to remove or disassemble the cover. There are no user-serviceable parts inside the console. All service should only be performed by an authorized Misonix representative. No modification of this equipment is required.
- WARNING 1.4 Proper system grounding can only be ensured when an approved, hospital-grade receptacle and matching power cord are used. To avoid the risk of electric shock, this equipment must only be connected to a supply with protective earth. Install plug and receptacles as per local regulations before operating the unit. Power cord, plug and receptacle should be examined to verify that they are in good working condition before connecting the console. Never pull on the power cord to remove it from the receptacle.
- WARNING 1.5 Connecting the console to a power outlet with inadequate voltage or frequency may cause the unit to malfunction or to create a shock or fire hazard. Confirm that the voltage selector switch on the console rear panel is set to the local voltage setting and ensure that the correct fuses are being used. Refer to section 12.1 on instructions for fuse replacement.

#### 1.3. Environmental Statement

This equipment consists of materials that may be recycled if disassembled by a specialized company. Please observe local and federal regulations regarding the disposal of packing materials and old equipment.

#### Important Environmental Information for Users within the European Economic Area

The European Parliament did enforce new regulations developed in 2005 concerning the disposal medical electrical and electronic equipment. The regulations, called Directives, place responsibilities on the supplier and you, the purchaser/user. One of the actions required is to inform users of their obligations.

The device has been assessed in accordance with the European Parliament Directive 2002/96/EC on Waste Electrical and Electronic Equipment, usually referred to as WEEE Directive.

The WEEE Directive requires that the device be disposed of at the end of its useful life in an environmentally responsible manner. Similar requirements have been applied to refrigerators for some time.

The WEEE Directive requires that if replacing the device with a new equivalent product, that supplier shall collect the old item without cost to the user.

If you wish to dispose of the device without replacing it then the device must not be mixed with unsorted municipal waste. The crossed-out wheeled bin symbol on the unit label or packaging, and repeated below, indicates this requirement.



Disposal Symbol, disposal to be compliant with EN 50419

You must ensure that the device is disposed of at an authorized treatment facility; details can be obtained from your local council.

Table 1.3 Environmental statement

# 1.4. Summary Of Safety Notices

Please read this section of the manual carefully. It contains a summary of all precaution, warning and caution statements contained in the manual. However, the user is advised to read the entire manual and operate the device only in accordance with all of the instructions contained herein.

Servicing of this device should only be performed by qualified technicians authorized by Misonix, LLC. There are no service controls accessible to the user.

Conventions on Warnings, Cautions and Notes		
WARNING	Denotes potentially dangerous situation that could result in death or serious injury to patient, operator or staff.	
CAUTION	A caution contains information regarding any special care to be exercised by the practitioner and/or patient for the safe and effective use of the device.	
NOTE	Indicates potential hazard that may result in product damage.	

Table 1.4 Conventions on warnings, cautions and notes

### **List Of Warnings**

- WARNING 1.1 The SonicOne O.R. system is an electro-mechanical device, which under certain circumstances could present an electrical shock hazard to the operator and/or patient. Please read manual thoroughly and follow directions stated herein to assure maximum safety during operation. This manual shall be kept in close proximity to the system for easy referral when needed.
- WARNING 1.2 The SonicOne O.R. system is intended to be used in various types of invasive, surgical procedures. There may be indirect danger to the patient should the device fail during the procedure. It is recommended that the facility follows its back-up equipment protocols.
- WARNING 1.3 The SonicOne O.R. system generates high voltages within the console itself and the connected handpiece.

  To avoid injury, the console should never be operated before ensuring that its cover is properly closed and not tampered with. Do not attempt to remove or disassemble the cover. There are no user-serviceable parts inside the console. All service should only be performed by an authorized Misonix representative. No modification of this equipment is required.
- WARNING 1.4 Proper system grounding can only be ensured when an approved, hospital-grade receptacle and matching power cord are used. To avoid the risk of electric shock, this equipment must only be connected to a supply with protective earth. Install plug and receptacles as per local regulations before operating the unit. Power cord, plug and receptacle should be examined to verify that they are in good working condition before connecting the console. Never pull on the power cord to remove it from the receptacle.
- WARNING 1.5 Connecting the console to the power outlet with inadequate voltage or frequency may cause the unit to malfunction or to create a shock or fire hazard. Confirm that the voltage selector switch on the console rear panel is set to the location voltage setting and ensure that the correct fuses are being used. Refer to section 12.1 on instructions for fuse replacement.
- WARNING 1.7 The SonicOne O.R. system, including all accessories and components, is MR unsafe. It must not be brought into the MR environment.
- WARNING 3.1 Tip and irrigant temperatures may exceed the tissue necrosis point if insufficient irrigation flow rates are used. For hard tissue removal, set the irrigation flowrate to a setting no less than the comparable vibration setting. For example, if the vibration setting is 7, a minimum flow setting of 70% should be used. Additional external irrigation, e.g. by administering sterile saline with a syringe over the distal tip portion, may be necessary for removal of very dense, hard osseous structures.
- WARNING 3.2 Tissue necrosis may result if tip is not moved relative to tissue. A continuous, lateral sweeping motion is recommended in order to minimize contact duration with the ultrasonic tip and minimize heat build-up. When lateral motion is not possible withdraw and re-insert tip frequently.
- WARNING 3.3 The SonicOne O.R. system and it's accessories may emit harmful acoustic pressure if exposure exceeds recommended limits.
- WARNING 3.4 For the Wide Hatch Probe: Less than 50" (127 cm) tall patients to wear ear protection during debridement.

  All other patients, to wear ear protection only for upper body debridement.
- WARNING 4.1 Contact to vibrating elements like extension and ultrasonic tip may cause burns and should be avoided by all means. The handpiece should only be held at the black housing area. An optional, protective silicone sleeve, included with certain tips, reduces the risk of thermal damage but does not eliminate it. Contact with the silicone sleeve should be avoided or kept brief with minimal amount of contact pressure. Pressure and extended exposure can still result in excessive frictional heat and cause burns.
- WARNING 4.2 Ultrasonic tips can break under excessive use in extreme conditions, e.g. when cutting for extended / duration in tight cavities with limited lateral motion. The tip could break into two or more fragments with the main fragment remaining attached to the handpiece. All fragments must be retrieved immediately from the surgical site. The fragments should be checked to ensure that no further pieces are missing. It is possible that a fragment is propelled outside of the surgical cavity. Diagnostic imaging, such as X-ray, must be used if a fragment cannot be found to confirm that the broken piece is outside of the surgical cavity.
- WARNING 4.3 Breakage of ultrasonic tips will result in sharp edges that can be harmful to soft tissue even without activation of ultrasound. Tips can bend or deform before they actually brake. Tips showing signs of deformation or cracking should be replaced immediately since tip breakage is otherwise imminent. Do not bend or twist the ultrasonic tips since it reduces the structural integrity and can result in tip breakage during use. Dispose of deformed or broken tips immediately in a sharps container.

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- WARNING 6.1 Immediately suspend operation if Electrical Fault appears on display and/or an Electrical Fault audible indicator sounds. Remove ultrasonic tip from surgical site. Turn Mains Power OFF. Do not touch any metallic parts of handpiece, extension, ultrasonic tip or generator while fault is indicated.
- WARNING 7.1 Improper connection of the handpiece cable may present a shock hazard. Confirm that handpiece connector is dry prior to plugging it in.
- WARNING 7.2 Do not operate pump with pump cover in raised position. Rollers might pinch loose clothing or fingers.

  Personal injuries may result. Cooling of the ultrasonic tip and irrigation at the treatment site will be inhibited.
- WARNING 7.3 Tip and irrigation temperatures may exceed the tissue necrosis point with SonicOne O.R. accessories for hard tissue removal if insufficient irrigation flow rates are used. Always set the irrigation flowrate for hard tissue removal to a setting no less than the comparable vibration setting. For example, if the vibration setting is a 7, a minimum flow setting of 70% should be used.
- WARNING 7.4 Heat is being generated at the tip/tissue interface. A continuous, lateral sweeping motion is recommended for general bone/tissue removal in order to minimize contact duration with the ultrasonic tip and minimize the temperature increase.
- WARNING 8.1 Remove probe cover, ultrasonic tip and extension from the handpiece prior to cleaning and/or sterilization; otherwise proper cleaning/sterilization may be inhibited.
- WARNING 9.1 Single-use items should be discarded following each surgical procedure according to hospital protocol for disposal of bio contaminated wastes. Do not attempt to reuse or re-sterilize any single-use items

  Dispose ultrasonic tips in a sharps container.
- WARNING 9.2 All reusable handpiece parts and accessories must be properly decontaminated, cleaned and sterilized before each use as per instructions contained in this manual. Failure to do so may lead to infections, which can ultimately cause patient death.
- WARNING 9.3 Misonix LLC has validated all cleaning and sterilization cycles given in this manual. It is highly recommended that the procedures given in this manual for cleaning and sterilizing the SonicOne O.R.system and related accessories be followed. It is the responsibility of the user of this device or any accessories used with it to validate procedures for cleaning and/or sterilization if they differ from the procedures as outlined in this manual.
- WARNING 9.4 The disposable items are intended for one procedure only (single use). Do not attempt to reuse or re-sterilize.
- WARNING 10.1 If a Mains Power fuse fails after replacement when the unit is reactivated, discontinue use of the device and contact an authorized Misonix representative.
- WARNING 10.2 Portable and mobile RF communication equipment (including peripherals such as antennas) should be no closer than 30 cm (12 inches) to any part of the console, including the cables supplied with the console Otherwise degradation of the performance of this equipment could result.
- WARNING 10.3 The use of accessories, transducers and cables other than those specified or provided by Misonix may result in increased electromagnetic emissions or decreased immunity of the device and may result in improper operation. Use only Misonix branded equipment and accessories.
- WARNING 10.4 Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.
- WARNING 12.1 Connecting the console to a power outlet with inadequate voltage or frequency may cause the unit to malfunction or to create a shock or fire hazard. Confirm that the voltage selector switch in the console rear is set to the local voltage setting and ensure that the correct fuses are being used. Refer to section 12 on fuse replacement.
- WARNING 12.2 No Modifications of this equipment is allowed except as noted for cleaning and sterilization. The user should return to Misonix or an authorized service center for servicing.

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#### **List Of Cautions**

CAUTION 7.8

CAUTION 8.1

- **CAUTION 1.1** Special Skills Training Requirements 1. Federal law restricts this device to sale by, or on the order of a licensed healthcare practitioner. The SonicOne O.R. system is to be used by an appropriately trained and licensed healthcare practitioner. **CAUTION 1.2** This device is considered medical electrical equipment. Medical electrical equipment needs special precautions regarding electromagnetic compatibility (EMC) and needs to be installed and put into service according to the EMC information provided in this operator's manual. CAUTION 4.1 Ultrasonic energy is inhibited if excessive physical force is applied to the ultrasonic tip; use only enough force to guide the tip to the surgical site and to advance it through the tissue. Do not force the tip; allow the ultrasonic action to do the work. CAUTION 4.2 Insufficient irrigation and high tip pressure (loading) under extended exposure, e.g. in tight cavities, are to be avoided in SonicOne O.R. hard tissue removal. It is recommended to withdraw and re-insert the ultrasonic tip repeatedly to re-establish adequate cooling and lubrication. CAUTION 4.3 Additional external irrigation, e.g. by administering sterile saline with a syringe over the distal tip portion may be necessary for removal of very dense, hard osseous structures of the skull, when using the SonicOne O.R. accessories. CAUTION 7.1 All reusable system components like handpiece, probe covers, counter wrench, and T-wrench are supplied industrially cleaned, but NON-STERILE. All items intended for use in the sterile field must be cleaned and sterilized as per the indicated instructions before first clinical use and before every subsequent clinical use. CAUTION 7.2 All items intended for use in the sterile field must be cleaned and sterilized as per indicated instructions before each clinical use. CAUTION 7.3 The disposable items are intended for one procedure only (single use). Do not attempt to reuse or re-sterilize. Do not place the soft silicone tube behind or in front of the rollers (latch removed in illustrations) CAUTION 7.4 CAUTION 7.5 Do not pinch the soft silicone tube when the latch is locked. CAUTION 7.6 Do not pinch barb fittings when closing the latch. **CAUTION 7.7** Prime the irrigation tubing prior to use. At all times ensure that the irrigant flows towards the handpiece
- before assembly.
- CAUTION 9.1 Do not use ultrasonic cleaners to clean the handpiece as this method could damage handpiece.
- CAUTION 9.2 Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder sterilization of units during autoclaving.

when footswitch is depressed. If no irrigant is flowing, cease use until flow is restored.

The system check should always be done in advance of preparing patient for surgery to minimize risk to

Ensure all connections and mating surfaces of handpiece, extension and ultrasonic tip are clean and dry

- CAUTION 9.3 Do not immerse ultrasonic console, handpiece, irrigation pump, remote footswitch or electric cables. These items are not sealed against liquids and damage to equipment will result.
- CAUTION 12.1 Use only genuine replacement parts from Misonix. Use of parts furnished by other sources may result in patient or operator injury or system malfunction and will void any applicable warranty.
- CAUTION 12.2 Before using loose packing materials, such as foam pellets, shredded paper or similar, be sure to wrap the component(s) separately in plastic bags, film or other protective wrapping.

patient in case of system malfunction.

#### **List Of Notes**

NOTE 4.1 After extended periods of operation, the bottom of the console housing may become warm to the touch. This is normal. Do not touch the bottom of the console housing while in operation or shortly after operation. NOTE 4.2 Loose tip/tissue contact upon an initial bone incision can cause a thin tip to resonate not only longitudinally but also transversely. This can cause a thin tip to break. It is necessary to engage bone actively and with a minimal tip pressure greater than zero in order to prevent the shattering. NOTE 4.3 Contact of the ultrasonic tip or the exposed extension with metal, surgical instruments or other objects during ultrasound use must be avoided. Such contact can damage the ultrasonic components very easily and may result in compromised performance, including failure. Discard any extensions or tips that show signs of damages like gouges, nicks or fractures. External aspiration may be used but it is recommended that a plastic suction tip should be used when in proximity with the probe tip. **NOTE 7.1** Adequate air circulation is needed to cool electronic components inside of the unit. Do not block the cooling fan at the console rear or the air vents on the console bottom. Do not place the unit on a towel, foam or other soft surface since the material may block the air vents. Blocking these vents may cause unit to overheat and malfunction or create a shock hazard. A clear drape can be used to protect the console front panel but do not cover the pump housing or other console portions. **NOTE 8.1** The handpiece must be placed into the counter wrench. Do not attempt to tighten or loosen handpiece components by holding the handpiece case or endcap. Always use the T-wrench wrench when tightening or un-tightening the tip or an extension. Never apply a pipe or strap wrench to the handpiece case. Do not over-tighten the tip or the extension. **NOTF 8.2** Always tighten or un-tighten the probe cover by hand and without using any wrenches. Do not overtighten the probe cover. **NOTE 8.3** Always hold the handpiece at its metallic endcap when tightening or un-tightening the irrigation tubing. Always tighten or un-tighten the irrigation tubing by hand and without using any wrenches. Do not overtighten the tubing connector. **NOTE 9.1** Allow reusable, autoclavable items to gradually return to room temperature after steam sterilization and prior to usage. NOTE 9.2 The reuse life given takes into account wear and tear due to cleaning and sterilization only. Damage or wear caused by actual use in treatments will affect life of components.

#### 1.5. Trademark Information

Misonix® and SonicOne® are registered trademarks of Misonix, LLC, Farmingdale, NY ASP Enzol® and Prolystica® are registered trademarks of STERIS Corporation, Mentor, OH

# 1.6. Explanation Of Symbols

## **Console Related Symbols**

Symbol	Description	Symbol	Description	Symbol	Description
•	Enable / Standby Ultrasound	4	Caution: Dangerous voltage		Mains Power ON
	Scroll through menu pages		Caution: Consult accompanying documents	0	Mains Power OFF
3)))	Amplitude setting		Caution: Pinch hazard	=	Protective earth ground
	Pulse setting	<b>†</b>	Type B equipment	$\Diamond$	Equipotentiality connection
<u>\</u>	Flow setting	STERILE EO	Sterilized using Ethylene Oxide		Disposal to be compliant with EN 50419 (WEEE directive)
2	Do not reuse	YYYY- MM-DD	Use by date indicated	R <sub>X</sub> ONLY	Restricted to sale by or on the order of a physician only
<b>(S)</b>	Do not use if packaging is damaged	LOT ABC12	Lot or batch code	EC REP	Authorized representative
LATEX	Contents are latex-free	-	Fuse	REF	Catalog number
_°C Max	Do not expose to temperatures greater than indicated	C UL US	Classified by UL	~	AC Voltage
<b>WH</b>	Must use hospital grade powercord only		Vacuum source		Manufacturer
<b>C C</b> 0482	Misonix CE number				Footswitch connector
PHT	Contains DEHP and/or Phthalates			*	Irrigation Source

Symbol	Description
	Warning: Hearing Protection
MR	MR Unsafe
UDI	Unique Device Identification is specific to a manufacturer and a device
MD	Identifies product as a medical device
	Date of Manufacture
STEELE	Do not re-sterilize
NON	Non sterile medical device
	Single sterile barrier system with protective packaging inside

## 2. Indications And Contra Indications

#### 2.1. Indications

The SonicOne O.R.® is indicated for use in debridement of wounds, such as, but not limited to burn wounds, diabetic ulcers, bedsores and vaginal ulcers, soft tissue debridement and cleansing of the surgical site in application, in which, in the physician's judgement would require the use of an ultrasonic aspirator with sharp debridement.

It is also indicated for use in the fragmentation and aspiration of both soft and hard (e.g.: bone) tissue as used in the following surgical specialties. Wound Care, Orthopedic Surgery, Plastic and Reconstructive Surgery, Thoracic Surgery, NeuroSurgery and General Surgery.

The SonicOne O.R. system to be operated by medical professionals justified to practice in the fields covered by the stated indications for use.

CAUTION 1.1 Special Skill and Training requirements. Federal law restricts this device to sale by or on the order of a licensed healthcare practitioner. The SonicOne O.R. is to be used by an appropriately trained and licensed healthcare practitioner.

#### 2.2. Contra Indications

The SonicOne O.R. ultrasonic surgical aspirator system is not indicated for and should not be used for cardiac surgery and any procedure in the proximity of the heart.

The irrigation pump is not indicated for and should not be used for the administration of parenteral fluids, infusion of drugs, or for any life sustaining purposes.

This SonicOne O.R. ultrasonic surgical aspirator device is not indicated for and should not be used for the fragmentation, emulsification, and aspiration of uterine fibroids.

# 3. Adverse Effects

Limits For Airborne Acoustic Exposure			
Distance fr or pat	Maximum Exposure Period Within a 24 hour period		
3" - 24" 8 cm – 60 cm		28 minutes	
> 24"	> 60 cm	287 minutes	

WARNING 3.1	Tip and irrigant temperatures may exceed the tissue necrosis point if insufficient irrigation flow rates are used. For hard tissue removal, set the irrigation flowrate to a setting no less than the comparable vibration setting. For example, if the vibration setting is 7, a minimum flow setting of 70% should be used. Additional external irrigation, e.g. by administering sterile saline with a syringe over the distaltip portion, may be necessary for removal of very dense, hard osseous structures.
WARNING 3.2	Tissue necrosis may result if tip is not moved relative to tissue. A continuous, lateral sweeping motion is recommended in order to minimize contact duration with the ultrasonic tip and minimize heat build-up. When lateral motion is not possible withdraw and re-insert tip frequently.
WARNING 3.3 Wear	The SonicOne O.R system and its accessories may emit harmful acoustic pressure if exposure exceeds recommended limits.
vveai	hearing protection or protect patient hearing if not within the exporsure limits.
WARNING 3.4	For the Wide Hatch Probe: Less than 50" (127 cm) tall patients to wear ear protection during debridement. All other patients

# 4. Considerations During Clinical Use

WARNING 4.1 Contact to vibrating elements like extension and ultrasonic tip may cause burns and should be avoided by all means. The handpiece should only be held at the black housing area. An optional, protective silicone sleeve, included with certain tips, reduces the risk of thermal damage but does not eliminate it. Contact with the silicone sleeve should be avoided or kept brief with minimal amount of contact pressure. Pressure and extended exposure can still result in excessive frictional heat and cause burns.

NOTE 4.1 After extended periods of operation, the bottom of the console housing may become warm to the touch.

This is normal. Do not touch the bottom of the console housing while in operation or shortly after operation.

#### 4.1. SonicOne O.R. Use

#### **Recommended Settings**

The following settings are general guidelines and should be adjusted based on indication, anatomy, pathology and surgeon's preference.

	Amplitude	Pulse	Flow
Highest	10	100%	100%
Very High	9	100%	90%
High	8	100%	80%
Standard (Default)	7	100%	70%
Moderate	6	100%	60%
Low	5	100%	50%

Table 4.1 Recommended settings

- · A high amplitude setting results in more aggressive tissue removal, a low setting in less aggressive tissue removal.
- A higher amplitude setting in combination with lower irrigation could result in increased tissue necrosis. A lower amplitude setting in combination with higher irrigation would minimize or eliminate tissue necrosis.
- Bone shaving tips tend to require a lower amplitude than cutting blades.

CAUTION 4.1 Ultrasonic energy is inhibited if excessive physical force is applied to the ultrasonic tip; use only enough force to guide the tip to the surgical site and to advance it through the tissue. Do not force the tip; allow the ultrasonic action to do the work.

#### Tip Limitations During Bone Removal

Both the ultrasonic tip and the extension are vibrating at high frequency and are thus exposed to extreme mechanical stresses, especially when cutting bone.

- WARNING 4.2 Ultrasonic tips can break under excessive use in extreme conditions, e.g. when cutting for extended duration in tight cavities with limited lateral motion. The tip could break into two or more fragments with the main fragment remaining attached to the handpiece. All fragments must be retrieved immediately from the surgical site. The fragments should be checked to ensure that no further pieces are missing. It is possible that a fragment is propelled outside of the surgical cavity. Diagnostic imaging, such as X-ray, must be used if a fragment cannot be found to confirm that the broken piece is outside of the surgical cavity.
- WARNING 4.3 Breakage of ultrasonic tips will result in sharp edges that can be harmful to soft tissue even without activation of ultrasound. Tips can bend or deform before they actually brake. Tips showing signs of deformation or cracking should be replaced immediately since tip breakage is otherwise imminent. Do not bend or twist the ultrasonic tips since it reduces the structural integrity and can result in tip breakage during use. Dispose of deformed or broken tips immediately in a sharps container.
- CAUTION 4.2 Insufficient irrigation and high tip pressure (loading) under extended exposure, e.g. in tight cavities, are to be avoided in SonicOne O.R. hard tissue removal. It is recommended to withdraw and re-insert the ultrasonic tip repeatedly to re-establish adequate cooling and lubrication.
- CAUTION 4.3 Additional external irrigation, e.g. by administering sterile saline with a syringe over the distal tip portion, may be necessary for removal of very dense, hard osseous structures of the skull, when using the SonicOne O.R. accessories.
- NOTE 4.2 Loose tip/tissue contact upon an initial bone incision can cause a thin tip to resonate not only longitudinally but also transversely. This can cause a thin tip to break. It is necessary to engage bone actively and with a minimal tip pressure greater than zero in order to prevent the shattering.
- NOTE 4.3 Contact of the ultrasonic tip or the exposed extension with metal, surgical instruments or other objects during ultrasound use must be avoided. Such contact can damage the ultrasonic components very easily and may result in compromised performance, including failure. Discard any extensions or tips that show signs of damages like gouges, nicks or fractures. External aspiration may be used but it is recommended that a plastic suction tip should be used when in proximity with the probe tip.

# 5. System Overview

# 5.1. Principle Of Operation

The SonicOne O.R. system is designed to ultrasonically dissect and fragment soft and hard tissue. The system consists of an ultrasonic console with handpiece and accessories. The console features an integrated irrigation pump.





Figure 5.1 Misonix Console

Figure 5.2 SonicOne O.R Handpiece

The console produces an electrical signal that is fed into the handpiece and its piezoelectric transducer. The transducer converts the electrical signal into mechanical vibrations. The vibratory motion is amplified all the way down to the tip's distal end. Various tip shapes and sizes are available to achieve desired tissue effects.

- Wound Debridement Applications
  - Debridement probes are typically used for contact wound debridement. Tissue excision and fragmentation are achieved through cavitation and other mechanical and hydrodynamic effects.
- Applications: Specialized tips are utilized on hard tissue structures.
  - SonicOne O.R. blades, typically used for cutting bone, are usually flat and have a blunt active edge.
     A compression cut is achieved through repetitive impacts on the bone at an ultrasonic frequency.
  - Bone shaving tips are used for sculpting bone. They have an abrasive surface for bone removal through abrasion under ultrasonic oscillation.
  - SonicOne O.R. multi-function tips can have a combination of blunt and abrasive cutting surfaces.

A peristaltic pump, integrated into the SonicOne O.R. console, provides irrigant (sterile physiological saline) to the operative site during use.

# 5.2. Reusable System Components

The following system components represent the minimum requirements for performing soft and hard tissue procedures. They can be ordered as a system or individually.

Required System Components				
BCM-GN / EBC06 or BCM-GN-100V	SonicOne O.R System, includes BCM-GN (console and footswitch)	1 ea.		
BCM-GN/EBC06 or BCM-GN-100V	Misonix console	1 ea.		
SOM-HP	Includes console and footswitch  SonicOne O.R handpiece	1 ea.		
BCM-CW	Counter wrench for SonicOne O.R handpiece	1 ea.		
BCM-2W	T-Wrench	1 ea.		
BCM-SS	Hard Tissue Probe Cover	1 ea.		
BCM-H2	Soft Tissue Probe Cover	1 ea.		

Table 5.2 Required system components

Components and quantities included with the system may change over time, please check with your Misonix representative for the most current configuration.

# 5.3. Single Use, Sterile Components

At least one irrigation tubeset must be available for each surgical procedure.

	Irrigation Tubeset	
MXB-T	Irrigation Tubeset	1 ea.

Table 5.3 Irrigation tubeset

There are a variety of ultrasonic tips available for the SonicOne O.R. system. Please ask your Misonix representative for the latest catalog of available tips. Ultrasonic tips are supplied sterile and are for single use only.

SonicOne O.R. Tips		
MXC-C1	Curette Style Titanium Probe Kit	
MXC-R1	Cylindrical Titanium Probe Kit	
MXC-X1	Hatched Titanium Probe Kit	
MXC-X2	Wide Hatch Probe Kit	
MXC-C2-VAC	Debridement Probe with Aspiration	
MXC-X2-VAC	Debridement Probe with Aspiration	

## 6. Console

# 6.1. Receptacles, Controls And Indicators

1

The rear of the console features receptacles for the power cord, fuses, footswitch cable, equipotentiality connector, and IV-pole as well as a switch for mains power. The equipotentiality connector makes the connected equipment potential equal (ref IEC 60601-1).

2 3 4

5

- 1. IV-pole receptacle
- 2. Mains power on/off
- 3. Power cord receptacle with fuse block
- 4. Voltage selector switch
- 5. Cooling fan
- 6. Equipotentiality connection
- 7. Footswitch receptacle

Figure 6.1.1 Console rear

6

The front of the console features a receptacle for the handpiece cable and an irrigation pump head, in which the irrigation tubing is inserted. A large color LCD screen provides information on system status and set points for ultrasound amplitude, pulse rate and irrigant flow rate with respective controls on the panel below. Additional controls for ultrasound enable/ standby and menu access are provided on the left of the display panel. An ultrasound timer indicates the elapsed time, in which the ultrasound was on.



- 1. Amplitude setting
- 2. Pulse setting
- 3. Flow setting
- 4. Enable/standby button
- 5. Ultrasound timer
- 6. Menu button
- 7. Handpiece cable receptacle
- 8. Indicator for flow direction
- 9. Irrigation pumphead
- A-F Custom buttons

Figure 6.1.2 Console front

Buttons A-F perform various functions, depending on the information displayed on the screen. The display screen shown is the Main Screen used for all major control functions.

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The handpiece receptacle is keyed in order to facilitate connection. The red dot on top of the receptacle must be in line with the corresponding red dot on the handpiece cable.

#### 6.2. Menu Functions

The standard screen is the Main Screen. Additional screens are the Options and the Help Screen. Both the Options and Help screens can be accessed by pressing the menu button to toggle through the three main screens; Main Menu, Options and Help.

#### Main Screen

The Main Screen allows control of the main system functions such as Amplitude, Pulse and Flow.



Menu Button
 A-F Custom buttons

Figure 6.2.1 Main screen

#### **Amplitude Control**

The amplitude can be set between 0 and 10. Press A to increase and B to decrease the amplitude in increments of 1. The default setting for amplitude is 7. Refer to section 1.1 for further details on the Amplitude feature.

#### **Pulse Control**

The pulse can be set between 50% and 100%. Press C to increase and D to decrease the pulse in increments of 10%. The default setting for pulse is 100%. Refer to section 6.3 for further details on the Pulse feature.

#### Flow Control

The flow can be set between 20% and 100%. Press E to increase and F to decrease the flow. The flow is increased and decreased in increments of 5% from 20%-40% and increased and decreased in increments of 10% from 40% to 100%. The default setting is 70%. Refer to section 6.3 for further details on the Flow feature.

#### **Ultrasound Timer**

The ultrasound timer records the elapsed time, in which the ultrasound was activated with the footswitch. The timer can be re-set to zero via the secondary screen.

In the event of error, such as a Mechanical Limit or an Electrical Fault, the main screen is replaced by alert screens. Refer to section 6.4 for a description of these warnings.

#### **Options Screen**

The Options Screen allows the user to do the following; re-set the elapsed ultrasound time, save and choose presets and adjust the display contrast.



Menu Button
 A-F Custom buttons

Figure 6.2.2 Options screen

#### Ultrasound timer

The elapsed ultrasound time can be re-set to 00:00 by pressing either A or B.

#### Presets

Preferred settings for amplitude, pulse and flow can be saved as two presets. A third preset features the default settings, which cannot be customized. A sub-screen for presets 1, 2 and 3 can be accessed by pressing either C or D.

#### Contrast

The display contrast can be adjusted. Press E to increase and F to decrease contrast.

Pressing the menu button toggles from the options to the help screen.

#### Help Screen

The Help Screen provides access to a quick guide on system operation and troubleshooting.

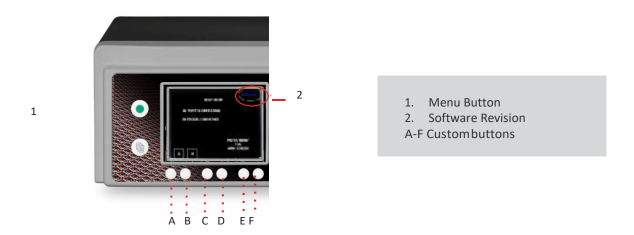


Figure 6.2.3 Help screen

#### **System Operation**

Press A to access the quick reference guide on system operation.

#### **Troubleshooting**

Press B to access the quick reference guide on troubleshooting.

The software revision may be found in the upper right corner of this screen. Pressing the menu button toggles from the help to the main screen.

#### 6.3. Main Functions

#### **Amplitude**

The ultrasonic tip engages the target area in linear strokes at a rate of approximately 22,500 cycles per second. During each cycle the tip elongates from resting to maximum position, contracts back over resting and to minimum position and elongates back to its resting point. The peak-to-peak amplitude or stroke distance can be adjusted by changing the Amplitude from setting 1-10. This is the main parameter to control the rate of tissue removal. A high amplitude setting results in more aggressive tissue removal, a low setting in less aggressive tissue removal. Amplitude and thus removal rate may alter with size and geometry of the ultrasonictip.

#### **Pulse**

The ultrasonic energy output over time can be reduced by using the pulse mode, in which a resting period is inserted within the duty cycle. This results in an active period, followed by a resting period during each duty cycle. The total period is a ¼ second (250ms). The pulse can be set between 50% and 100%.

#### Pulse Setting 100% [Continuous]

The default setting is 100% or continuous, which refers to 100% energy output or zero resting period. This is the recommended setting.

#### Pulse Setting 50-90% [Pulsed]

The Pulse function minimizes exposure to ultrasound over time.

The Pulse setting corresponds to the duration of the active period of the ultrasound output. For example, a Pulse setting of 60% corresponds to an active period of 60% of ¼ second (150ms). The resulting resting period is 40% of ¼ second (100ms). The ultrasonic energy output over time is reduced by 40% with this setting. Note that the ultrasound timer will only advance during the active period and not during the resting period.

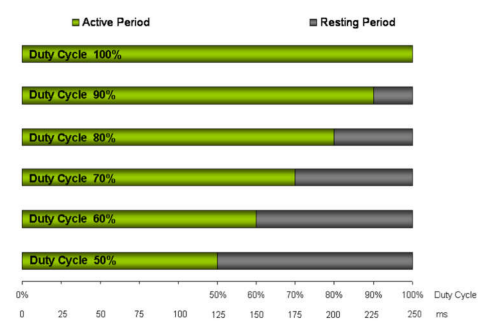


Figure 6.3.1 Illustration of pulse setting

#### Irrigation

Proper irrigation with sterile saline ensures:

- 1. Cooling of handpiece and vibrating elements
- 2. Cooling and lavage of the surgical site
- 3. Lubrication of bone/tip interface for SonicOne O.R. hard tissue removal

The active ultrasonic probe remains cold when not in contact with tissue. However, when a tip contacts tissue heat is generated. The heat increases with applied tip pressure or amplitude. Irrigant needs to be applied at the tip/tissue interface to mitigate this temperature rise.

Most ultrasonic tips and probes feature an integrated irrigation channel. The irrigant is expelled through a jet nozzle at the tip. Active tip surfaces are being cooled directly.

Standby Mode	Enable Mode
23:57 ▼ PULSE FLOW	80% 30% 23:57 \( \begin{array}{cccccccccccccccccccccccccccccccccccc
Amplitude setting is GREY and HOLLOW	Amplitude setting is GREEN and SOLID
Footswitch activates Irrigation only. Irrigation can be used for lavage or priming.	Footswitch activates  Ultrasound output and irrigation. A bell chime is emitted briefly.

Table 6.3.2 Enable/standby function

#### Enable/Standby

The Enable/Standby button on the console's front panel can be used to block accidental ultrasound activation during longer periods of inactivity following set-up or during surgery.

## 6.4. Alerts And Indicators

#### Mechanical Limit Alert

The console monitors the ultrasonic output at all times and alerts in cases of overload or malfunction of the vibrating elements (handpiece, extension and ultrasonic tip). A "Limit" alert is displayed together with a pulsed audible indicator as long as the footswitch is depressed. Ultrasound and Irrigation are deactivated temporarily.

Mechanical Limit Alert		
Alert Type	Alert Screen	Alert Action
Mechanical Limit	80% 30% 2357 PULSE FLOW	Displays "LIMIT" alert located above amplitude setting display.  Triggers a pulsed, audible indicator upon footswitch activation.  Temporarily deactivates ultrasound and irrigation functions.
Possible Cause	Corrective Action	
Tip overload	Release footswitch. Reduce tip pressure and/or use higher amplitude setting as required. Continue procedure.	
Loose or damaged component	Release footswitch and Set ultrasound to STANDBY. Remove silicone sleeve (if applicable) and probe cover. Inspect extension probe and ultrasonic tip for damage. Replace if necessary. Otherwise re-tighten extension probe and tip using the correct wrenches. Set ultrasound to ENABLE and Continue procedure.	
Defective Handpiece	If corrective action steps above are followed and alert continues, the handpiece may need to be replaced.	

Table 6.4.1 Mechanical limit alert and recommended corrective actions

Tip overload can occur during hard tissue removal when applying excessive tip pressure or facing strong tissue resistance, e.g. from thick cortical bone. This can lead to stalling of the ultrasonic tip. A pulsed audible signal alerts of the stalling and the ultrasound is deactivated. Release the footswitch briefly and reduce the tip pressure, e.g. by retrieving the ultrasonic tip. Depress the footswitch again and continue with reduced tip pressure. Consider using higher amplitude setting or reduced loading if stalling persists.

#### **Electrical Fault Alert**

The console monitors the electrical output at all times and alerts in cases where the handpiece is not properly connected to the console, when an output short or open circuit is detected or electrical safety is compromised.

An Electrical Fault Screen is displayed together with a steady audible indicator. Ultrasound and Irrigation are deactivated. Requires recycling of mains power switch to re-set.

WARNING 6.1 Immediately suspend operation if Electrical Fault appears on display and/or an Electrical Fault audible indicator sounds. Remove ultrasonic tip from surgical site. Turn Mains Power OFF. Do not touch any metallic parts of handpiece, extension, ultrasonic tip or generator while fault is indicated.

Electrical Fault Alert		
Alert Type	Alert Screen	Alert Action
Electrical Fault	THE CHROAL FAME & OCCURRED  CHECK HANDWICK FOR CHACKS  AND PROPER CONNECTION  TO MARK SHIPE.  CHART SHIPE.  AMERITMEN:  PARKS: MARK MUNICIPAL  YOU MUST POWER DOWN 10 PRINCE HAND THESE HALK IT	Displays Electrical Fault Screen.  Triggers steady audible indicator.  Permanently deactivates ultrasound and irrigation
Possible Cause	Corrective Action	
1. Handpiece not connected	Turn mains power OFF. Check handpiece cable connection. Restart console.	
2. Defective Handpiece	Turn mains power OFF. Replace handpiece and restart console. If problem persists replace console.	
3. Defective console	Turn mains power OFF. Replace console.	

Table 6.4.2 Electrical fault alert and recommended corrective actions

# 7. System Set-up

#### 7.1. Installation

Upon delivery perform a visual inspection of the shipping containers and all system components for obvious shipping damage. Retain the shipping container and immediately notify the shipping carrier of any damages found.

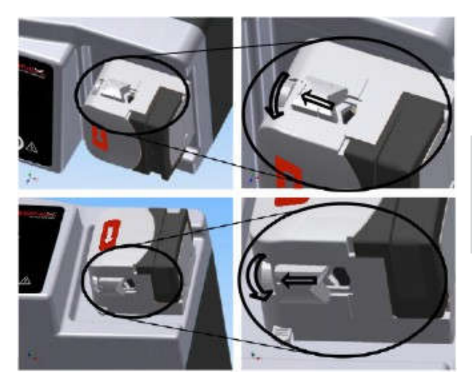
- CAUTION 7.1 All reusable system components like handpiece, probe covers, counter wrench, and T-wrench are supplied industrially cleaned, but NON-STERILE. All items intended for use in the sterile field must be cleaned and sterilized as per the indicated instructions before first clinical use and before every subsequent clinical use.
- WARNING 12.1 Connecting the console to a power outlet with inadequate voltage or frequency may cause the unit to malfunction or to create a shock or fire hazard. Confirm that the voltage selector switch on the console rear panel is set to the local voltage setting and ensure that the correct fuses are being used. Refer to section 12.1 on instructions for fuse replacement. See section 12.1 for instructions on adjusting to local electrical requirements.

Care should be taken to stay within the general operating conditions.

Operating Conditions	
Operating conditions	<ul> <li>Temperature 55-86°F (13-30°C)</li> <li>Relative humidity 20-90% (non condensing)</li> <li>-91m (-300ft) to 3000m (9840ft)</li> </ul>

Table 7.1 Operating conditions

The console can be placed on an appropriate table or cart outside of the sterile field. Ensure that the pump head on the console right is installed. Refer to section 12.2 if the pump head is not yet installed.



Adjust the grip of the V-notches

Evenly adjust the V-notches to their fully opened position by turning the adjustment wheels.

Figure 7.1.1

The console features air vents on the bottom. When installing the unit, ensure that these vents are not blocked in a way that would prevent the circulation of air around the unit.

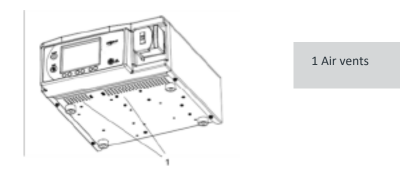


Figure 7.1.2 Underside view of console with air vents

NOTE 7.1 Adequate air circulation is needed to cool electronic components inside of the unit. Do not block the cooling fan at the console rear or the air vents on the console bottom. Do not place the unit on a towel, foam or other soft surface since the material may block the air vents. Blocking these vents may cause unit to overheat and malfunction or create a shock hazard. A clear drape can be used to protect the console front panel but do not cover the pump housing or other console portions.

## 7.2. Console Set-up – Part I (Non-sterile)

	Console	e Set-up   Part I	
Switch Mains Power OFF	Set Mains Power switch on console rear to OFF.		
Connect IV-pole	Connect IV-pole to receptacle in console rear.		
	Hang container with s	sterile physiological saline irrig	ant into IV-pole hook.
	Irrigation tubing featu	ures IV-spike and is compatible	with rigid bottles or flexible bags.
Check Mains Fuses	See section 12.1		
Check Voltage Selector	Voltage selector switch (see section 6.1 for location) shall be placed in a position that is dependent on the model and hospital voltage.		
	Model	Line Voltage	Selector Switch Position
	BCM-GN	120 VAC, 60 Hz	"115V"
		220/230/240 VAC, 50/60 Hz	"230V"
Connect Electrical Power	Connect power cord to	receptacle on console rear and	to wall outlet.
Connect Footswitch	Onnect Footswitch  Connect footswitch cable to receptacle on console rear.  Footswitch connector and receptacle are keyed to ensure proper connection. Turn cable connector until keys match. Insert connector fully into receptacle. Turn outer connector ring clockwise to lock into position.		
	Footswitch may be co	overed with clear drape during	clinical use.
Switch Mains Power ON	Set Mains Power switch	h on console rear to ON.	
	Front panel will displa	ay Main Screen upon completion	on of system start.

Table 7.2 Console set-up - part I

# 7.3. Handpiece Assembly (Sterile)

Handpiece assembly in the sterile field should be performed by trained OR staff only.

Please refer to section 8.0 for specifics on the handpiece assembly and disassembly.

Once the handpiece has been assembled, continue with part II of the Console Set Up.

CAUTION 7.2 All items intended for use in the sterile field must be cleaned and sterilized as per indicated instructions before each clinical use.

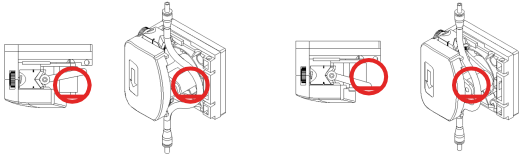
CAUTION 7.3 The disposable items are intended for one procedure only (single use). Do not attempt to reuse or re-sterilize.

# 7.4. Console Set-up — Part II (Non-sterile)

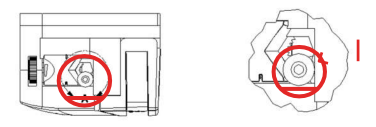
Console Set-up   Part II		
Connect Handpiece cable	Attach cable connector receptacle on console front panel.	
	Align red dot on cable connector with red dot on front panel receptacle. Push cable connector into place.	
Open pump cover	Open the latch of the irrigation pump	
	The arrow on the pump housing indicates the direction of flow.	
Insert tubing	Insert the soft silicone section by placing it over the pump rollers.	
	Verify that the tubing enters the pump from fluid container and exits to handpiece and in direction of arrow on pump housing.	
Secure tubing in V-notches	Place and hold the tubing's silicone section in the V-notches on both pump sides.	
	Apply slight pressure to ensure that tubing rests within both V-notches.	
Close pump cover	Close the latch of the irrigation pump until it locks.	
Connect Tubing to	Connect IV-Spike to fluid container following standard operating room protocol.	
Fluid Container	Irrigation tubing features vented IV-spike and is compatible with rigid bottles and flexible bags.	
Prime Tubing	Check that ultrasound is in Standby Mode. Set Flow rate to 10.  Depress footswitch until fluid discharges at ultrasonic tip.	
Adjust Settings	Set Amplitude and Flow to desired flow setting. Refer to section 4.1 for recommended settings. Enable ultrasound.	

Table 7.4 Console set-up - part II

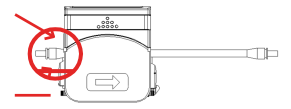
- WARNING 7.1 Improper connection of the handpiece cable may present a shock hazard. Confirm that handpiece connector is dry prior to plugging it in.
- WARNING 7.2 Do not operate pump with pump cover in raised position. Rollers might pinch loose clothing or fingers. Personal injuries may result. Cooling of the ultrasonic tip and irrigation at the treatment site will be inhibited.
- CAUTION 7.4 Do not place the soft silicone tube behind or in front of the rollers (latch removed in illustrations)



CAUTION 7.5 Do not pinch the soft silicone tube when the latch is locked.



CAUTION 7.6 Do not pinch barb fittings when closing the latch.



- CAUTION 7.7 Prime the irrigation tubing prior to use. At all times ensure that the irrigant flows towards the handpiece when footswitch is depressed. If no irrigant is flowing, cease use until flow is restored.
- WARNING 7.3 Tip and irrigation temperatures may exceed the tissue necrosis point with SonicOne O.R accessories for hard tissue removal if insufficient irrigation flow rates are used. Always set the irrigation flowrate for hard tissue removal to a setting no less than the comparable vibration setting. For example, if the vibration setting is 7, a minimum flow setting of 70% should be used.
- WARNING 7.4 Heat is being generated at the tip/tissue interface. A continuous, lateral sweeping motion is recommended for general bone/tissue removal in order to minimize contact duration with the ultrasonic tip and minimize the temperature increase.

The SonicOne O.R system is now ready for the system check.

# 7.5. Perform System Check

System Check	
Enable Ultrasound	Switch to Enable Mode using enable/standby button. Confirm that Amplitude setting is FILLED GREEN.
Depress footswitch	Direct ultrasonic tip toward suitable reservoir to collect irrigant. Depress footswitch.
Confirm Function	Console emits a bell chime. Irrigant will be pumped from console towards handpiece. Ultrasonic tip emits buzzing sound and irrigant exits tip as fine spray. Ultrasound timer counts up in 1-second increments.
Release footswitch	Release footswitch. Ultrasound and Flow output stop. Ultrasound timer freezes at last reading.
Function Confirmed	Reset ultrasound timer as desired. System is now ready for use.
Function NOT confirmed	Console alerts of Mechanical Limit or Electrical Fault or does not respond as expected. Refer to troubleshooting section for next steps.

Table 7.5 System check

CAUTION 7.8 The system check should always be done in advance of preparing patient for surgery to minimize risk to patient in case of system malfunction.

The SonicOne O.R. system is now ready for use. Refer to section 1.0 for general safety statements, indications and adverse effects and section 5.0 for use of main system functions.

# 8. Handpiece Assembly And Disassembly

The SonicOne O.R. system can accommodate different tip designs. The tip choice is determined by the surgeon's preference.

## 8.1. Handpiece Assembly

Perform an inspection of handpiece and all components prior assembly.

Handpiece Inspection	
Inspect Handpiece	Inspect the black handpiece housing for any visual cracks. Inspect the front metallic portion probe for surface damage like nicks, gouges and cracks. Replace if damaged.
Inspect Mating Surface	Inspect mating face of handpiece to verify that it is clean and dry.

Table 8.1 Handpiece inspection

- CAUTION 8.1 Ensure all connections and mating surfaces of handpiece, extension and ultrasonic tip are clean and dry before assembly.
- CAUTION 7.2 All items intended for use in the sterile field must be cleaned and sterilized as per indicated instructions before each clinical use.
- CAUTION 7.3 The disposable items are intended for one procedure only (single use). Do not attempt to reuse or re-sterilize.
- NOTE 8.1 The handpiece must be placed into the counter wrench. Do not attempt to tighten or untighten handpiece components by holding the handpiece case or endcap. Always use the T-wrench wrench when tightening or un-tightening the tip or an extension. Never apply a pipe or strap wrench to the handpiece case. Do not over-tighten the tip or the extension.
- NOTE 8.2 Always tighten or un-tighten the probe cover by hand and without using any wrenches. Do not over-tighten the probe cover.
- NOTE 8.3 Always hold the handpiece at its metallic endcap when tightening or un-tightening the irrigation tubing. Always tighten or un-tighten the irrigation tubing by hand and without using any wrenches. Do not overtighten the tubing connector.

Once the handpiece inspection is complete, please see the selected tip's (i.e., MXB-20) Instructions For Use (included in the sterile packaged product) for attachment instructions and use.

The handpiece is now ready for use and can be connected to the SonicOne O.R. system. Please refer to Section 7.0 for details.

## 8.2. Handpiece Disassembly

- WARNING 8.1 Remove probe cover, ultrasonic tip and extension from the handpiece prior to cleaning and/or sterilization; otherwise proper cleaning/sterilization may be inhibited. See section 9.
- NOTE 8.1 The handpiece must be placed into the counter wrench. Do not attempt to tighten or untighten handpiece components by holding the handpiece case or endcap. Always use the T-wrench wrench when tightening or un-tightening the tip or an extension. Never apply a pipe or strap wrench to the handpiece case. Do not over-tighten the tip or the extension.
- NOTE 8.2 Always tighten or un-tighten the probe cover by hand and without using any wrenches. Do not over-tighten the probe cover.
- NOTE 8.3 Always hold the handpiece at its metallic endcap when tightening or un-tightening the irrigation tubing. Always tighten or un-tighten the irrigation tubing by hand and without using any wrenches. Do not overtighten the tubing connector.



Table 8.2 Irrigation tube disassembly

Once irrigation tube is disconnected, please see the selected tip's (i.e., MXB-20) Instructions For Use (included in the sterile packaged product) for disassembly instructions.

# 9. Cleaning And Sterilization

Follow manufacturer's directions for preparing solutions. Misonix recommends the use of CaviWipes® or equivalent quaternary ammonium compound surface disinfectant wipe. Please follow manufacturer's instructions for surface cleaning and disinfection of hard non porous surfaces, including, without limitation, the use of personal Protection Equipment (PPE) for Blood borne Pathogens. Dispose of the used wipes in accordance with local regulations regarding the disposal of biological hazardous wipes.

## 9.1. Disassembly

Console Tear-down	
Disable Ultrasound	Switch to Standby Mode using enable/standby button. Confirm that Amplitude setting is HOLLOW GREY.
Switch Console OFF	Set Mains Power switch on console rear to OFF.
Remove Handpiece Cable	Pull cable connector from receptacle on console front.
Remove Tubing	Open pump cover. Remove tubing from pump compartment.  Disconnect tubing from irrigant container.
Wipe Down Console	Wipe down the console.

#### Table 9.1 Console tear-down

#### Handpiece Disassembly

Disassemble all handpiece components in reverse order of assembly. Please refer to Section 8.2 for disassembly after surgical procedure.

#### Dispose Of Single-Use Items

All items marked single use must not be reused. Reuse of these items could result in severe patient injury or death.

Once used, dispose of single use items in accordance with standard hospital procedures for disposal of biocontaminated wastes.

- WARNING 8.1 Remove probe cover, ultrasonic tip and extension from the handpiece prior to cleaning and/or sterilization; otherwise proper cleaning/sterilization may be inhibited.
- WARNING 9.1 Single-use items should be discarded following each surgical procedure according to hospital protocol for disposal of biocontaminated wastes. Do not attempt to reuse or re-sterilize any single-use items.

  Dispose ultrasonic tips in a sharps container.

## 9.2. Cleaning

Misonix LLC has validated the cleaning procedures outlined below.

Misonix continually updates its sterilization and cleaning instructions as required. For the latest instructions and reuse recommendations please contact your local Misonix representative.

- WARNING 9.2 All reusable handpiece parts and accessories must be properly decontaminated, cleaned and sterilized before each use as per instructions contained in this manual. Failure to do so may lead to infections, which can ultimately cause patient death.
- WARNING 9.3 Misonix LLC has validated all cleaning and sterilization cycles given in this manual. It is highly recommended that the procedures given in this manual for cleaning and sterilizing the SonicOne O.R System and related accessories be followed. It is the responsibility of the user of this device or any accessories used with it to validate procedures for cleaning and/or sterilization if they differ from the procedures as outlined in this manual.

### 9.2.1 Manual Cleaning Procedure

The following items are considered reusable items and should be cleaned as recommended:

- Console
- Footswitch
- Handpiece
- Counter wrench
- T-wrench
- Probe cover

Probe Cover and Wrenches			
Wash & Brush	<ul> <li>Wash items with hot water mixed with an enzymatic detergent such as ASP Enzol® or Steris Prolystica® according to standard hospital protocol. Follow manufacturer's directions for preparing solutions.</li> <li>Probe cover and wrenches may be fully immersed.</li> <li>Brush all passages at least four (4) times from FRONT to REAR, rotating the brushes during insertion and inserting the brushes fully. This ensures clearing of debris from the internal passages.</li> <li>Item's exterior surface can be cleaned using a standard soft bristle cleaning brush.</li> </ul>		
Rinse	Rinse item under warm running water for a minimum of 1 minute to clear soap residue.		
Dry	Dry item fully with absorbent towel or paper. Dispose of cloth or paper in accordance with Hospital or Clinic practices for contaminated wastes.		

Table 9.2.1.1 Cleaning of probe cover and wrenches

	Handpiece		
Wipe Cable	<ul> <li>Wipe cable with cloth or absorbent paper moistened with an enzymatic detergent such as ASP Enzol® or Steris Prolystica®. Follow manufacturer's directions for preparing solutions. Clean all surfaces of bloodstains and obvious signs of contamination.</li> </ul>		
Wash & Brush	<ul> <li>Wash and brush handpiece with hot water mixed with an enzymatic detergent such as ASP Enzol® or Steris Prolystica®. Follow manufacturer's directions for preparing solutions.</li> <li>The handpiece cannot be immersed.</li> <li>Brush all passages at least four (4) times from FRONT to REAR, rotating the brushes during insertion and inserting the brushes fully. This ensures clearing of debris from the internal passages.</li> <li>The item's exterior surface can be cleaned using a standard soft bristle cleaning brush.</li> </ul>		
Rinse	Rinse item under warm running water for a minimum of 1 minute to clear soap residue.		
Dry	Dry item fully with absorbent towel or paper. Dispose of cloth or paper in accordance with Hospital or Clinic practices for contaminated wastes.		
Inspect	• Inspect handpiece and cable and remove any item which shows signs of damages (cracks, gouges, fractures etc.). Mark damaged items clearly to prevent future use before disposal.		

Table 9.2.1.2 Cleaning of handpiece

- CAUTION 9.1 Do not use ultrasonic cleaners to clean the handpiece as this method could damage the handpiece.
- CAUTION 9.2 Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder sterilization of units during autoclaving.
- CAUTION 9.3 Do not immerse ultrasonic console, handpiece, irrigation pump, remote footswitch or electric cables. These items are not sealed against liquids and damage to equipment will result.

Console and Footswitch		
Wipe Surfaces	<ul> <li>Follow manufacturer's directions for preparing solutions. Misonix recommends the use of CaviWipes1 ® or an equivalent quaternary ammonium compound surface cleaning and disinfection wipe. Wipe footswitch and console, including irrigation unit, with cloth or absorbent paper moistened with an enzymatic detergent such as ASP Enzol® or Steris Prolystica®. Follow manufacturer's directions for preparing solutions. Clean all surfaces of bloodstains and obvious signs of contamination.</li> <li>Dispose of cloth or paper with contaminated waste.</li> </ul>	

Table 9.2.1.3 Cleaning of console and footswitch

- Caution 9.4 The console and footswitch should never be immersed in liquids, and liquid disinfectants should not be poured directly onto the equipment as irreparable damage or electrical hazards may result. Use disinfecting products on wipe substrates.
- Caution 9.5 The only external surfaces of the console should be cleaned. Do not attempt to remove any panels in order to clean or disinfect internal surfaces.

## 9.2.2 Automated Wash Procedure

		Handpiece,	Probe Cover and Wr	renches
Point of Use	Immediately following procedure perform the following:  Flush handpiece lumen with minimum 100 mL of saline to clear the bore of debris.  Wipe all reusable devices to remove visible blood and debris.			
	CAUTION: DO NOT use saline to wet the tray and tray contents before transport to the decontamina processing area.			
	NOTE: If transport to the decontamination processing area is delayed, cover the tr or spray the tray and its contents with a pre-cleaning foam. The pre-cleaning foan drying of soil and facilitate later decontamination processing.			. The pre-cleaning foam will minimize the
Pre-Cleaning	<ul> <li>The following should be performed on a disassembled handpiece:</li> <li>Remove the probe and all housingcomponents.</li> <li>Prepare neodisher® MediClean forte at 3.9 mL per liter of water (5/8 oz. per gallon water). Water should be lukewarm (&lt;40°C, &lt;104°F).</li> <li>Use a tight-fitting brush dipped in the prepared cleaning solution to clean the lumen of the handpiece by inserting the brush fully through the lumen until visible from the other side a minimum of four times, rotating the brush as it is inserted.</li> <li>Rinse all residual soap from the handpiece under warm running water for a minimum of one minute.</li> <li>Visually inspect internal and external surfaces of the handpiece including the pin cavity and repeat the above steps as required until all visible debris and staining are removed.</li> </ul>			
Automated Wash and Disinfection	to align the lumen in the general direction of the water jet flow in the washer but at a slight angle to facilitate draining during the drying cycle.			
	- Trocess the nanapi	ece and an reas	sable components and acc	cessories using the following cycle parameters
	Phase	Time*	Parameters	Detergent Type and Concentration
	Pre-Wash 1	2 minutes	Cold tap or purified water	None
	Wash 1	2 minutes	≥65.5°C (150°F)	neodisher® MediClean forte 2mL/L (¼ oz. / gallon)
	Rinse 1	1 minute	Hot tap water	None
	Disinfection	1 minute	≥90°C (194°F)	None
	Drying	6 minutes	≥98.8 °C (210°F)	None
Post-Cleaning	*Durations listed are minimum acceptable. Longer durations than those specified for cleaning and disinfection are acceptable.  Inspect all items for cleanliness and damage following cleaning and prior to terminal sterilization.			

Table 9.2.2.1 Cleaning of Handpiece, Probe Cover and Wrenches

## 9.3. Sterilizing By Steam Autoclave

Sterilization Methods and terminology are based on ANSI/AAMI ST81 AND EN ISO 17664:2004 standards.

#### 9.3.1 Reusable, Autoclavable Components

BCM-HP	Handpiece
BCM-BW	T-Wrench
BCM-CW	Handpiece/Counter Wrench
BCM-SS	Hard Tissue Probe Cover
BCM-H2	Soft Tissue Probe Cover

#### 9.3.1 Validated Steam Sterilization Cycles

#### With Sterilization Tray

	132°C (270°F)	134-137°C (274-279°F)
Configuration	Items placed in Misonix Sterilization Tray E-SYSTRAY¹ or E-SYSTRAY-2¹	Items placed in Misonix Sterilization Tray E-SYSTRAY <sup>2</sup> or E-SYSTRAY-2 <sup>2</sup>
Cycle	Prevacuum	Prevacuum
Preconditioning Pulses	4	4
Minimum Exposure Time	4 minutes*	3 minutes*
Minimum Dry Time	30 minutes	30 minutes

#### Wrapped, No Tray

	132°C (270°F)	134-137°C (274-279°F)	134°C/273°F
Configuration	Items wrapped, NO TRAY <sup>3</sup>	Items wrapped, NO TRAY <sup>3</sup>	Items wrapped, NO TRAY <sup>3</sup>
Cycle	Prevacuum	Prevacuum	Gravity
Preconditioning Pulses	4	4	None
Minimum Exposure	4 minutes*	3 minutes*	20 minutes
Minimum Dry Time	45 minutes	30 minutes	5 minutes

#### Table 9.3.2 Steam sterilization cycles

NOTE 9.1 Allow reusable, autoclavable items to gradually return to room temperature after steam sterilization and prior to usage.

<sup>&</sup>lt;sup>1</sup>Tray wrapped in Kimberly Clark KC300 KIMGUARD sterilization wrap.

<sup>&</sup>lt;sup>2</sup>Tray wrapped in Kimberly Clark KC300 or KC600 KIMGUARD sterilization wrap.

<sup>&</sup>lt;sup>3</sup>Items wrapped in Kimberly Clark KC300 or KC600 KIMGUARD sterilization wrap.

<sup>\*</sup>Exposure time can be increased up to a maximum of 18 minutes to comply with local requirements and/or recommendations of the World Health Organization (WHO), Robert Koch Institute (RKI), etc. Misonix LLC reusable medical devices are able to sustain such sterilization cycles.

## 9.4 Expected Life, Reusable Components

All handpiece components need to examined regularly, prior each use and be replaced if damaged.

The estimated sterilization life of handpiece components is listed below. All sterilization life estimates are approximate and may be affected by rough handling, damage, wear due to vigorous cleaning, etc.

Estimated Sterilization Life		
Item	Number Of Steam Sterilization Cycles	
Handpiece with attached cable	~200 cycles	
Probe covers	~300 cycles	
Wrenches: Handpiece/counter wrench and T-wrench	~300 cycles	

Table 9.4 Estimated re-use life

**NOTE 9.4** 

The reuse life given takes into account wear and tear due to cleaning and sterilization only. Damage or wear caused by actual use in treatments will affect life of components.

## 9.5 Deviations From Decontamination, Cleaning And Sterilization Instructions

Misonix LLC has validated all cleaning, disinfection and sterilization cycles given in this manual. It is highly recommended that the procedures given in this manual for cleaning and sterilizing the SonicOne O.R. system and related accessories be followed. It is the responsibility of the user of this device or any accessories used with it to validate procedures for cleaning and/or sterilization if they differ from the procedures as outlined in this manual.

#### Technical Assistance

Should the user wish further information or instructions regarding any aspect of cleaning or sterilizing procedures, please contact Misonix LLC or an Authorized Representative.

## 10. Troubleshooting

The SonicOne O.R. system provides both visual and audible alert signals when the system is not functioning properly.

Mechanical Limit Alert			
Alert Type	Alert Screen	Alert Action	
Mechanical Limit	BO% 30% 2357 PULSE FLOW	Displays "LIMIT" alert located above amplitude setting display.  Triggers a pulsed, audible indicator upon footswitch activation.  Temporarily deactivates ultrasound and irrigation functions.	

Possible Cause	Corrective Action	
Tip overload	Release footswitch. Reduce tip pressure and/or use higher amplitude setting as required. Continue procedure.	
Loose or damaged component	Release footswitch. Set ultrasound to STANDBY. Remove silicone sleeve (if applicable) and probe cover. Inspect extension probe and ultrasonic tip for damage. Replace if necessary. Otherwise re-tighten extension probe and tip using the correct wrenches. Set ultrasound to ENABLE. Continue procedure.	
Defective Handpiece	If corrective action steps above are followed and alert continues, the handpiece may need to be replaced.	

Table 10.1 Mechanical limit alert and recommended corrective actions

Tip overload can occur during hard tissue removal when applying excessive tip pressure or facing strong tissue resistance, e.g. from thick cortical bone. This can lead to stalling of the ultrasonic tip. A pulsed audible signal alerts of the stalling and the ultrasound is deactivated. Release the footswitch briefly and reduce the tip pressure, e.g. by retrieving the ultrasonic tip. Depress the footswitch again and continue with reduced tip pressure. Consider using higher amplitude setting or reduced loading if stalling persists.

Electrical Fault Alert			
Alert Type	Alert Screen	Alert Action	
Electrical Fault	FI FCTRICAL FAMILE OCCURRED  CHECK HANDPIECE FOR CRACKS AND PROPER CONNECTION TO MAIN HIEL.  CHERKE SHEMPA.  AMELITATIO: 1 PIL SS - MICH.  ELOW. 60P1 YOU MUST FROM FROM NI TO HESS! I HIELE! THIS FALL.	Displays Electrical Fault Screen.  Triggers steady audible indicator.  Permanently deactivates ultrasound and irrigation. Requires recycling of mains power switch to re-set.	
Possible Cause	Corrective Action		
Handpiece not connected	Turn console OFF. Check handpiece cable connection. Restart console.		
Defective Handpiece	Turn console OFF. Replace handpiece and restart console. If problem persists replace console.		
Defective console	Turn console OFF. Replace console.		

Table 10.2 Electrical fault alert and recommended corrective actions

WARNING 6.1 Immediately suspend operation if Electrical Fault appears on display and/or an Electrical Fault audible indicator sounds. Remove ultrasonic tip from surgical site. Turn Mains Power OFF. Do not touch any metallic parts of handpiece, extension, ultrasonic tip or generator while fault is indicated.

- WARNING 1.4 Proper system grounding can only be ensured when an approved, hospital-grade receptacle and matching power cord are used. To avoid the risk of electric shock, this equipment must only be connected to a supply with protective earth. Install plug and receptacles as per local regulations before operating the unit. Power cord, plug and receptacle should be examined to verify that they are in good working condition before connecting the console. Never pull on the power cord to remove it from the receptacle.
- WARNING 10.1 If a Mains Power fuse fails after replacement when the unit is reactivated, discontinue use of the device and contact an authorized Misonix representative.

### Lack of Irrigant

#### Symptoms

- No spray from tip when ultrasound is engaged
- No flush fluid available
- Unexpected temperature rise at operative site
- Unexpected temperature rise of handpiece

Alert Type	Alert Screen	Alert Action
None	None	None
Possible Cause	Ultrasound Mode	Corrective Action
1. Closed or empty fluid bag	Set ultrasound to STANDBY.	Check fluid bag and tubing clamp. Replace fluid bag if necessary.
2. Tubing not connected	Set ultrasound to STANDBY.	Check tubing connections.
		Check mounting in pump head. Close pump cover until locked.
3. Tubing obstructed or defective	Set ultrasound to STANDBY.	Check tubing for kinking, restrictions or leaks. Replace tubing if necessary.
		Check mounting in pump head. Close pump cover until locked.
4. Tubing installed in reverse	Set ultrasound to STANDBY.	Open pump cover. Reposition the tubing in direction of flow. Close pump cover until locked.
5. Tubing slides through pump	Set ultrasound to STANDBY.	Open pump cover. Adjust the grip of the tubing by turning the adjustment wheel underneath the front and back of the pump assembly.
6. Pump defect	Set ultrasound to STANDBY.	Open pump cover. Check if pump rollers are rotating when depressing footswitch. Replace console if they don't.

Table 10.3 Troubleshooting – Insufficient Irrigation

- WARNING 3.1 Tip and irrigant temperatures may exceed the tissue necrosis point if insufficient irrigation flow rates are used. For hard tissue removal, set the irrigation flowrate to a setting no less than the comparable vibration setting. For example, if the vibration setting is 7, a minimum flow setting of 70% should be used.
- WARNING 7.1 Improper connection of the handpiece cable may present a shock hazard. Confirm that handpiece connector is dry prior to plugging it in.
- WARNING 7.2 Do not operate pump with pump cover in raised position. Rollers might pinch loose clothing or fingers.

  Personal injuries may result. Cooling of the ultrasonic tip and irrigation at the treatment site will be inhibited.

For all other malfunctions please contact Misonix or a Misonix authorized representative for service.

# 11. Specifications

Console Specifications		
Power input (BCM-GN)	<ul> <li>120VAC, 4 Amps, 60Hz</li> <li>220/230/240 VAC, 2.5 Amps, 50/60 Hz</li> </ul>	
Power input (BCM-GN-100V)	<ul> <li>100VAC, 6.3 Amps, 50/60Hz</li> <li>200 VAC, 3.15 Amps, 50/60Hz</li> </ul>	
Operating frequency	22.5 kHz	
Ground leakage current	300 μA (max)	
Output power	130 Watts (nom)	
Mode of Operation	<ul><li>Continuous Wave</li><li>Pulse Wave</li></ul>	
Controls	<ul> <li>Mains Power on/off switch (rear panel)</li> <li>Footswitch control for ultrasonic and irrigation on/off</li> <li>Ultrasound enable/standby button</li> <li>Amplitude control</li> <li>Pulse control</li> <li>Flow control</li> <li>Ultrasonic timer with reset</li> <li>Menu button</li> <li>Six screen-specific buttons</li> </ul>	
Irrigation pump	Peristaltic pump	
Pump flow rate	Max flow > 67 ml/min.	
Irrigation tubing	<ul> <li>Dedicated tubeset, sterile, single-use</li> <li>Vented IV-spike, compatible with fluid bags and bottles</li> <li>Dedicated handpiece connection</li> </ul>	
Handpiece cable	• 15 ft   4.6m	
Footswitch cable	• 14 ft   4.3m	
Footswitch	• IP 68	
Generator	• IPX 0	
Power cord	• 10 ft   3.0m	
Operating conditions	<ul> <li>Temperature 55-86°F (13-30°C)</li> <li>Relative humidity 20-90% (non condensing)</li> <li>-91m (-300ft) to 3000m (9840ft)</li> </ul>	
Shipping/storage conditions	<ul> <li>Temperature: -4-122°F (-20-50°C)</li> <li>Relative humidity: 15-90% (non condensing)</li> </ul>	
Dimensions	7" H x 16" W x 19" D   180mm H x 410 mm W x 485mm D	
Weight	25.6 Lb.   11.6 kg	

Table 11.1 Console specifications

System Contents		
BCM-GN	SonicOne O.R console	1 ea.
SOM-HP	SonicOne O.R handpiece	2 ea.
BCM-CW	Counter wrench for SonicOne O.R handpiece	2 ea.
BCM-2W	T-Wrench	2 ea.
BCM-SS	Hard Tissue Probe cover	2 ea.
BCM-H2	Soft Tissue Probe Cover	2 ea.

Table 11.2 System contents

Components and quantities included with the system may change over time, please check with your Misonix representative for the most current configuration.

UL (IEC) 60601-1 Classification

Class 1 Equipment

Type B Equipment

**Ordinary Equipment** 

## 12. Service, Repair And Technical Correspondence

WARNING 1.4 Proper system grounding can only be ensured when an approved, hospital-grade receptacle and matching power cord are used. To avoid the risk of electric shock, this equipment must only be connected to a supply with protective earth. Install plug and receptacles as per local regulations before operating the unit. Power cord, plug and receptacle should be examined to verify that they are in good working condition before connecting the console. Never pull on the power cord to remove it from the receptacle.

WARNING 12.1 Connecting the console to a power outlet with inadequate voltage or frequency may cause the unit to malfunction or to create a shock or fire hazard. Confirm that the voltage selector switch in the console rear is set to the local voltage setting and ensure that the correct fuses are being used. Refer to section 12 on fuse replacement

## 12.1. Fuse Replacement

Model BCM-GN / E-BC06	Fuse Specifications			
Line Voltage	Manufacturer	Manufacturer P/N	Rating	Description
120 VAC, 60 Hz	Cooper/Bussman	GDB-4	250V @ 4 A	Fast Acting, Low Break- ing
220/230/240 VAC, 50/60 Hz	Littlefuse	021702.5	250V @ 2.5A	Fast Acting

Table 12.1.1 Console fuse specifications model BCM-GN

Model BCM-GN-100V	Fuse Specifications			
Line Voltage	Manufacturer	Manufacturer P/N	Rating	Description
100 VAC, 50/60 Hz	Cooper/Bussman	GDB-6.3	250V @ 6.3 A	Fast Acting, Low Break- ing
200 VAC, 50/60 Hz	Cooper/Bussman	GDB-3.15	250V @ 3.15 A	Fast Acting, Low Break- ing

Table 12.2 Console fuse specifications model

Fuse Replacement (The fuse holder is located on the console rear)			
Disable Ultrasound	Switch to Standby Mode using enable/standby button. Confirm that Amplitude setting is HOLLOW GREY.		
Switch Console OFF	Switch console OFF and disconnect pow	ver cord.	
Remove Fuse Holder			
	Pinch tab on fuse holder.	Pull fuse holder out.	
Replace Fuses	Replace both fuses as specified above.		
Mount Fuse Holder	Push fuse holder back into receptacle.		
Switch Console ON	Connect power cord and switch console ON		
Check Function	Confirm that console powers up and that Main Settings respond to activation of buttons A-F.		

Table 12.1.2 Fuse replacement

## 12.2 Pump Head Replacement

The pump head may not be connected to the unit for shipping purposes.

Mount Pump Head				
Destrice Description	Shaft recess and bayonet fitting on pump head rear	Pump drive shaft on console front		
Position Pump Head	Turn pump head 45°clockwise.	Align drive shaft on console front and shaft recess on pump head rear. Drive shaft and recess must engage easily. Rotate pump head slightly back and forth to check engagement.		
Lock pump head in place	Turn pump head clockwise until it locks in place. Arrow should be in the vertical position pointing down.			

Table 12.2.1 Assembly of pump head

Remove Pump Head				
Disable Ultrasound	Switch to Standby Mode using enable/standby button. Confirm that Amplitude setting is HOLLOW GREY.			
Switch Console OFF	Switch console OFF and disconnect pov	Switch console OFF and disconnect power cord.		
Remove Tubing	Open pump cover. Remove tubing. Close pump cover.			
Release Pump Head	Press and hold lock lever on pump head bottom.	Turn pump head 45° counter clockwise.		
Remove pump head	Pull pump head away from console until pump drive shaft clears. Release lock lever.			

Table 12.2.2 Disassembly of pump head

## 12.3 Repair, Service and Replacement Parts

All requests for repairs and replacement parts should be directed to Misonix or an authorized Misonix representative. Always provide model and serial number of malfunctioning items.

When returning items include model, serial and RMA number as well as purchase order number on all documents. Always prepay return shipping and specify method of shipment.

- CAUTION 12.1 Use only genuine replacement parts from Misonix. Use of parts furnished by other sources may result in patient or operator injury or system malfunction and will void any applicable warranty.
- CAUTION 12.2 Before using loose packing materials, such as foam pellets, shredded paper or similar, be sure to wrap the component(s) separately in plastic bags, film or other protective wrapping.
- WARNING 12.3 No Modifications of this equipment is allowed except as noted for cleaning and sterilization. The user should return to Misonix or an authorized service center for servicing.

### 12.4 Important Notice

Please contact Misonix with any questions regarding the specifications, use, sterilization, limitations or maintenance of the SonicOne O.R. System:

Misonix, LLC

Web <u>www.misonix.com</u> Email sales@misonix.com

Phone +1.631.694.9555 / 1-800-694-9612

Fax +1.631.694.9412 Address 1938 New Highway

Farmingdale, NY 11735

U.S.A.

Any serious incident occurring in relation to the SonicOne O.R. System should be reported to Misonix, LLC (using the contact information listed above) and the competent authority of the Member State in which the user is established.

By returning any material to Misonix, LLC the customer or the customer's agent must certify that any and all materials so returned are or have been rendered free of any hazardous or noxious matter or radioactive contamination and are safe for handling under normal repair shop conditions.

Do not return any material for which such certification cannot be made without prior approval from Misonix, LLC.

The correct return address should read as follows:

MISONIX (Misonix, LLC)
Medical Service Department
RMA # \_\_\_\_\_
1938 New Highway
Farmingdale, New York 11735
U.S.A.

Please contact Misonix for a list of other authorized service centers.

The authorized EC representative is:

EC REP

(E

Emergo Europe Westervoortsedijk 60 6827 AT Arnhem The Netherlands



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