nexus

BoneScalpel® AccessTM Handpiece Instructions for Use

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1. Overview

This Misonix neXus BoneScalpel AccessTM Instructions for Use Manual describes how to use the handpiece and handpiece accessories. Misonix recommends that you read and understand the instructions in this manual before using the handpiece. Misonix also recommends that you read and understand the separate Instructions for Use Manuals for the neXus Ultrasonic Surgical Aspiration System.

This Instructions for Use Manual includes indications for use, contraindications, general safety statements, adverse effects, handpiece assembly and disassembly, cleaning and sterilization, specifications, service, repair and technical correspondence, and replacement parts.

Table 1 Conventions on Warning and Cautions

Conventions on Warnings and Cautions		
WARNING	Denotes potentially dangerous situation that could result in death or serious injury to patient, operator, or staff.	
CAUTION		

Refer to Section 4.5 for symbols used on product labeling and in the Misonix neXus® IFUs.

1.1. Principle of Operation

The Misonix neXus® Ultrasonic Surgical Aspirator System is comprised of a generator which converts mains voltage and frequency to a 22.5 kHz (neXus® Standard Handpiece) or 23.0 kHz (neXus BoneScalpel Access™ Handpiece, neXus SonaStar® Long Handpiece and neXus SonaStar® Short Handpiece) electrical signal depending upon the handpiece and accessories that are connected to the console. The generator feeds the electric signal to a piezoelectric transducer comprised of a ceramic crystal stack in the handpiece. The crystals vibrate at the output frequency translating the electrical energy into mechanical vibration. A titanium horn amplifies the vibration and transmits the amplified vibration to a titanium probe tip. The titanium probe tip is the applied part that comes into contact with patient tissue. An integrated irrigation pump delivers an irrigation solution to the surgical site. An integrated aspiration system removes the fragmented, emulsified material and waste liquids from the area. Accessories include various horn/probe tips, irrigation & aspiration tubing sets, wrenches, and cleaning brushes. The system, with the Misonix neXus SonaStar® Long and Short Handpieces may also be combined with electrosurgery using optional RF surgery interface components.

2. Indications and Contra Indications

2.1. Indications

The indications for use for the neXus **BoneScalpel**® **AccessTM Handpiece** with the approved neXus **BoneScalpel**® **AccessTM Probe and Tubeset** single-use disposable kits are listed below.

The neXus BoneScalpel® AccessTM Handpiece is indicated for use in the fragmentation and aspiration of soft and hard (e.g., bone) tissue in the following surgical specialties:

- Neurosurgery
- Gastrointestinal and Affiliated Organ Surgery
- Urological Surgery
- Plastic and Reconstructive Surgery
- General Surgery
- Orthopedic Surgery
- Gynecology
- External genitalia condyloma benign tumors (lipomas, fibromas, and leiomyomas) malignant primary and metastatic tumors of all types and the following cystic lesions: Bartholin's cysts, Vestibular adenitis, Inclusion cysts, Sebaceous cysts
- Abdominal area any abnormal growth, cystic or solid, benign, or malignant, involving the ovary, fallopian tube, uterus, or the supporting structures of the uterus except as contraindicated for uterine fibroids.
- Thoracic Surgery
 Limited pulmonary reception such as segmetectomies, nonanatomical subsegmentectomies and metastatectomies.

2.2. Contraindications

- The neXus Ultrasonic Surgical Aspirator System probe tips are not indicated for and should not be used for direct contact with cardiac tissue (direct cardiac application).
- 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 neXus Ultrasonic Surgical Aspirator System device is not indicated for and should not be used for the fragmentation, emulsification, and aspiration of uterine fibroids.

3. General Safety Statements

WARNING

The neXus® Ultrasonic Surgical Aspirator 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 to ensure maximum safety during operation. This manual shall be kept in close proximity to the system for easy referral when needed.

WARNING

The neXus® Ultrasonic Surgical Aspirator 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

Special Skills Training Requirements

- Caution: United States Federal law restricts this device to sale by or on the order of a physician or health care practitioner licensed by the law of the State in which the practitioner practices to use or order the use of the device. Not applicable in the European Union.
- The neXus® Ultrasonic Surgical Aspirator System is to be used by an appropriately trained and licensed healthcare practitioner.
- All health care institution personnel are to be trained in the healthcare institution's procedures for universal precautions for bloodborne pathogens and the use of appropriate PPE.

3.1. Summary of Safety Notices

Please read this section of the manual carefully. It contains a summary of all 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, Inc. There are no service controls accessible to the user.

Table 2: Conventions on Warnings and Cautions

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

3.2. List of Warnings

- The neXus® Ultrasonic Surgical Aspirator 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.
- The neXus® Ultrasonic Surgical Aspirator System is intended to be used in various types of 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.
- Explosion Hazard: Never use the neXus[®] Ultrasonic Surgical Aspirator System in the presence of a flammable or explosive atmosphere, such as flammable anesthetics.
- Only use the BoneScalpel Access® Handpiece with probe accessory kit configurations for the indications for use charted in **Section 2.1**

• Potential Burn Hazard

- o neXus probes have a silicone or hard plastic sheath. Compressing or bending the sheath may cause the sheath to contact the vibrating surface of the probe along the length of the probe or at the probe tip and may cause excessive heating, which may burn user or patient tissue at the surgical site.
- Excessive loading of neXus probes at the surgical site may induce heating due to vibration and friction as target tissue is fragmented and emulsified. It is critical to manage the temperature of the probe by adjusting the irrigation, aspiration, and ultrasound settings, and surgical technique. Tissue necrosis may result if probe tip is not moved relative to tissue. A continuous, lateral sweeping motion is recommended in order to minimize contact duration with the ultrasonic probe tip and minimize heat build-up. When lateral motion is not possible withdraw and re-insert probe tip frequently.
- Contact to vibrating elements like the extension and ultrasonic tip may cause burns and should be avoided by all means. The handpiece should only be held at the black handpiece housing or sheath.
- A protective silicone sleeve, included with certain probe tips, reduces the risk of thermal damage but
 does not eliminate it. Contact of non-target patient tissue 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.
- Contact of the rigid or silicone sheaths with patient tissue under pressure, may create a burn hazard.
 Avoid contact of sheath elements with patient tissue under pressure.
- O Probe Tip temperatures may exceed the tissue necrosis point if insufficient irrigant is present at the probe tip-tissue interface. For hard tissue removal, always use the maximum irrigation flowrate that does not affect the surgical field of view, or impact surgical technique. 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.
- o For hard tissue applications, a minimum Irrigation setting of 20 is recommended to minimize or prevent thermal injury and/or tissue necrosis.
- 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.

- 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 break. 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 biohazardous sharps container.
- Immediately suspend operation if Electrical Fault appears on the console display and/or an Electrical Fault
 audible indicator sounds. Remove ultrasonic tip from surgical site. Turn Mains Power OFF on the console. Do
 not touch any metallic parts of handpiece, extension, ultrasonic tip, or generator while fault is indicated.
- Improper connection of the handpiece cable may present a shock hazard. Confirm that handpiece connector is dry prior to plugging it in.
- Remove black sheath and ultrasonic tip from the handpiece prior to cleaning and/or sterilization; otherwise, proper cleaning/sterilization may be inhibited. Refer to **Section 5: Handpiece Assembly and Disassembly** and **Section 6: Cleaning and Sterilization**.
- Single-use disposable kits are marked with the international symbol for "do not reuse single use only" (②). Discard these items following each surgical procedure in accordance with the health care institution protocol for biohazardous waste. Tips are to be disposed of in a biohazardous sharps container.
- The reusable handpiece and accessories (counter wrench, torque wrench, and sterilization tray) must be properly decontaminated, cleaned and sterilized before each use. Refer to **Section 6: Cleaning and Sterilization**. Failure to do so may lead to infections, which can ultimately cause patient death.
- All Misonix reusable items must be sterilized by moist heat (steam sterilization/autoclaved) after manual cleaning.
- Automated cleaning-disinfection is not the final step prior to use. All Misonix reusable items must be sterilized by moist heat (autoclaved) after automated cleaning and disinfection.
- Misonix Inc. has validated all cleaning and sterilization cycles given in this manual. Refer to Section 6: Cleaning and Sterilization. To prevent transmission of disease or malfunction of the neXus® system, Misonix recommends that the procedures given in this manual for cleaning and sterilizing the neXus® Ultrasonic Surgical Aspirator System and related accessories be followed. However, other end-user validated cleaning agents, cleaning procedures, and/or sterilization cycles may be used.

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 in accordance with applicable local laws and regulations on cleaning and sterilization of reusable medical devices in the healthcare environment.

- For all sterilization protocols listed in **Section 6: Cleaning and Sterilization**, always assure the cap is placed securely on the cable connector to protect the connector during sterilization.
- The single-use disposable kits are intended for one procedure only. Do not attempt to reuse, clean or re-sterilize single-use disposable kit components.
- The neXus® Ultrasonic Surgical Aspirator System and its accessories may emit harmful acoustic pressure if exposure exceeds recommended limits. Refer to **Section 4: Adverse Effects** for the recommended limits.

- Modifications to the equipment are NOT allowed, except as noted for cleaning and sterilization instructions. The user should return the equipment to Misonix or an authorized service center if the equipment malfunctions or requires service.
- The neXus® Ultrasonic Surgical Aspirator System, including all accessories and components, is MR Unsafe. It must not be brought into the MR environment.

3.3. List of Cautions

- Special Skills Training Requirements
 - o Caution: United States Federal law restricts this device to sale by or on the order of a physician or health care practitioner licensed by the law of the State in which the practitioner practices to use or order the use of the device. Not applicable in the European Union.
 - o The neXus® Ultrasonic Surgical Aspirator System is to be used by an appropriately trained and licensed healthcare practitioner.
 - o All health care institution personnel are to be trained in the healthcare institution's procedures for universal precautions for bloodborne pathogens and the use of appropriate PPE.
- The use of accessories, transducers, and cables other than those specified may result in increased RF emissions or decreased immunity to RF of the device. Use only Misonix branded equipment and accessories.
- 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.
- Insufficient irrigation and high tip pressure (loading) under extended exposure, e.g., in tight cavities, are to be avoided while removing hard tissue. It is recommended to withdraw and re-insert the ultrasonic tips (e.g., Blades & Shavers) repeatedly to re-establish adequate cooling and lubrication. Additional external irrigation, e.g., by administering sterile saline with a syringe over the distal tip portion, may be necessary when removing very dense, hard osseous structures.
- All reusable system components like handpiece, counter wrench, and torque 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 in Section 6: Cleaning and Sterilization before the first clinical use and before every
 subsequent clinical use.
- The single-use disposable kits are intended for one procedure only. Do not attempt to reuse, clean or re-sterilize the single-use disposable kit components.
- Prime the irrigation tubing prior to use. At all times ensure that the irrigation flows towards the handpiece when footswitch is depressed. If no irrigation 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 patient in case of system malfunction.
- Ensure all connections and mating surfaces of handpiece, extension and ultrasonic tip are clean and dry before assembly.
- Do not use ultrasonic cleaners to clean the handpiece as this method could damage handpiece.
- Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder sterilization of units during autoclaving. Refer to the Pre-cleaning step below.
- Do not immerse the ultrasonic handpiece or the handpiece cable. These items are not sealed against liquids and damage to equipment will result.
- 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.

- Use softened, filtered, or deionized water for diluting cleaning agents and for the final equipment rinse. Deionized water is recommended for the final rinse, if available. Mineral residues from hard water in the final rinse step can cause water stains and/or affect cleaning and disinfection.
- The tethered handpiece cable cap should be placed on the handpiece cable connector immediately after the cable is disconnected from the console to prevent damage to the connector pins and remain on during precleaning, manual cleaning, automated cleaning/disinfection, and sterilization procedures.
- Misonix does not recommend "FLASH" sterilization. Misonix has not validated "FLASH" sterilization.
- Poor steam quality may impair the sterilization process. For this reason, various norms (European standard EN 285 and the United States standard ANSI/AAMI ST79) recommend maximum impurity levels for steam feed water of autoclaves and sterilizer used in the medical field. Misonix recommends using water of a quality that conforms to the norms, the health care institution validated specifications for water quality, or otherwise using deionized water to generate steam for moist heat sterilization.
- Ensure all connections and mating surfaces of handpiece and ultrasonic tip are clean and dry before assembly.
- 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.
- 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.
- 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 torque wrench when tightening or untightening the tip. Never apply a pipe or strap wrench to the handpiece case. Do not over-tighten the tip or the extension.
- Always tighten or un-tighten the black sheath by hand and without using any wrenches. Do not over- tighten the probe cover.
- Always hold the handpiece at its metallic endcap when attaching or removing the irrigation tubing.
- Allow reusable, autoclavable items to gradually return to room temperature after steam sterilization and prior to usage.
- The specified reuse life considers wear and tear due to cleaning and sterilization. Damage or wear caused by misuse in treatments will affect life of components.

3.4. Symbol Definition Chart

Table 3: Symbol Definitions

Table 5: Symbol Definitions		
(B)	Caution: Consult accompanying documents	
R _X ONLY	Caution: United States Federal law restricts this device to sale by or on the order of a physician or health care practitioner licensed by the law of the State in which the practitioner practices to use or order the use of the device; Not applicable in the European Union.	
~	Manufacturer	
$\prod i$	Consult Instructions for Use	
2	Do Not Re-Use	
STERILE EO	Sterilized using Ethylene Oxide	
LOT ABC123	Lot or Batch Code	
	Warning: Hearing Protection	
	Disposal to be compliant with EN 50419 (WEEE Directive)	
REF	Catalog number	
MR	MR Unsafe	

4. Adverse Effects

4.1. Airborne Acoustic Exposure

Table 4: Limits for Airborne Acoustic Exposure

Limits for Airborne Acoustic Exposure		
Distance from operator's or patient's ear		Maximum Exposure Period Within a 24-hour period
3" - 24" 8 cm - 60 cm		28 minutes
> 24"	> 60 cm	287 minutes

WARNING

The neXus® Ultrasonic Surgical Aspirator System and its accessories may emit harmful acoustic pressure if exposure exceeds recommended limits.

4.2. Probe Tip Temperature

WARNING

Probe tip temperatures may exceed the tissue necrosis point if insufficient irrigant is present at the probe tip-tissue interface. For hard tissue removal, always use the maximum irrigation flowrate that does not affect the surgical field of view, or impact surgical technique. Additional external irrigation, e.g., by administering sterile saline with a syringe over the distal probe tip portion, may be necessary for removal of very dense, hard osseous structures.

WARNING

Tissue necrosis may result if probe tip is not moved relative to tissue. A continuous, lateral sweeping motion is recommended in order to minimize contact duration with the ultrasonic probe tip and minimize heat build-up. When lateral motion is not possible withdraw and reinsert probe tip frequently.

WARNING

neXus probes have a silicone or hard plastic sheath. Compressing or bending the sheath may cause the sheath to contact the vibrating surface of the probe along the length of the probe or at the probe tip and may cause excessive heating, which may burn user or patient tissue at the surgical site.

WARNING

Excessive loading of neXus probes at the surgical site may induce heating due to vibration and friction as target tissue is fragmented and emulsified. It is critical to manage the temperature of the probe by adjusting the irrigation, aspiration, and ultrasound settings, and surgical technique. Tissue necrosis may result if probe tip is not moved relative to tissue. A continuous, lateral sweeping motion is recommended in order to minimize contact duration with the ultrasonic probe tip and minimize heat build-up. When lateral motion is not possible withdraw and re-insert probe tip frequently.

WARNING

Contact to vibrating elements like the extension and ultrasonic tip may cause burns and should be avoided by all means. The handpiece should only be held at the black handpiece housing or sheath.

WARNING

A protective silicone sleeve, included with certain probe tips, reduces the risk of thermal damage but does not eliminate it. Contact of non-target patient tissue 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

Contact of the rigid or silicone sheaths with patient tissue under pressure, may create a burn hazard. Avoid contact of sheath elements with patient tissue under pressure.

5. Handpiece Assembly and Disassembly

Handpiece assembly in the sterile field should be performed by trained and authorized OR staff only. Once the handpiece has been assembled, refer to neXus Console Instructions for Use for connectivity with system.

CAUTION	Ensure all connections and mating surfaces of handpiece and ultrasonic tip are clean and dry before assembly.
CAUTION	Single-use disposable kits are marked with the international symbol for "do not reuse - single use only" (②). Discard these items following each surgical procedure in accordance with the health care institution protocol for biohazardous waste. Tips are to be disposed of in a biohazardous sharps container.
CAUTION	The reusable handpiece and accessories (counter wrench, torque wrench, and sterilization tray) must be properly decontaminated, cleaned and sterilized before each use. Refer to Section 6: Cleaning and Sterilization . Failure to do so may lead to infections, which can ultimately cause patient death.
CAUTION	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.
CAUTION	Always tighten or un-tighten the black sheath by hand and without using any wrenches. Do not over-tighten the probe cover.
CAUTION	Always hold the handpiece at its metallic endcap when assembling or removing the irrigation tubing. Always assemble or remove the irrigation tubing by hand and without using any wrenches. Do not over- tighten the tubing connector.

5.1. Items Required for Handpiece Assembly

Table 5: Items Required for Handpiece Assembly

Part Number	Description	
100-22-0001 BoneScalpel Access TM (BSA) Handpiece		
100-64-0000	Access Handpiece Counter Wrench	
100-63-0000	Handpiece Torque Wrench	
100-71-0000	Sterilization Tray	

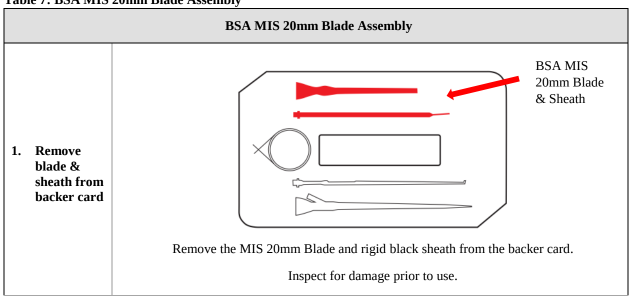
5.2. Handpiece Inspection

Table 6: Handpiece inspection

Handpiece Inspection: Perform Inspection Prior To Use		
Inspect Handpiece	Inspect the handpiece housing and front cover for damage and signs of wear such as scratches, cracks, and chips.	
	Inspect the handpiece cable to assure it is not cut or frayed.	
	Inspect the handpiece cable connector, connector pins, and the tethered cap to assure they are not damaged.	
	Place the tethered cap on the connector after inspection and leave in place until connection to the generator.	
	Do not use damaged handpieces or handpiece components. Contact Misonix Customer Service if damage is noted.	
	The handpiece should be fully inspected for loose or missing components and tested for proper operation prior to each procedure.	
Inspect Mating Surface	Inspect mating face of handpiece and ultrasonic tip to verify that it is clean and dry.	

5.3. neXus BoneScalpel® AccessTM Handpiece Assembly

Table 7: BSA MIS 20mm Blade Assembly



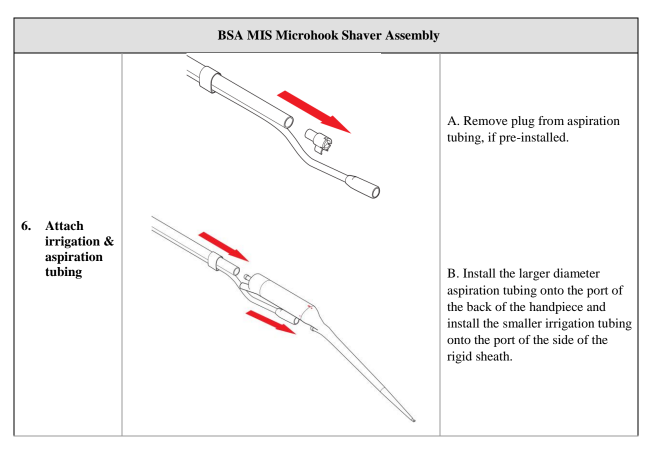
	BSA MIS 20mm Blade Assembly			
2.	Install the tip	Insert the tip into the handpiece and hand tighten turning clockwise.		
3.	Insert handpiece into the Counter Wrench	Counter Wrench	Insert the BoneScalpel Access Handpiece into the Counter Wrench as indicated on the drawing. The handpiece should be angled upward as indicated in the drawing.	
4.	Tighten the ultrasonic tip		Slide the torque wrench over the tip, ensuring that etched text "Handpiece this side" faces the handpiece. Rotate clockwise until one click is heard. Remove the wrench.	

	BSA MIS 20mm Blade Assembly		
5.	Install the housing		A. Slide the rigid sheath onto the probe and align the small triangular mark on the rigid sheath to the small triangular lower mark on the handpiece.
			B. Tighten by turning clockwise until the small triangular mark on the rigid sheath aligns with the upper mark, resembling a lock, on the Handpiece.
			Attach the smaller diameter irrigation tube onto the port at the back of the handpiece.
6.	Attach irrigation tubing		NOTE: The larger diameter aspiration tubing is NOT to be connected to the handpiece and remains free standing.

Table 8: BSA MIS Microhook Shaver Assembly

BSA MIS Microhook Shaver Assembly BSA MIS 1. Remove MicroHook shaver & Shaver & sheath from Sheath backer card Remove the MIS Microhook Shaver and the rigid and soft black sheath from the backer card. Inspect for damage prior to use. Install the tip Insert the tip into the handpiece and hand tighten turning clockwise. Insert the BoneScalpel Access Handpiece into the Counter Wrench as indicated on the 3. Insert handpiece drawing. into the The handpiece should be angled counter upward as indicated in the wrench drawing. **Counter Wrench**

	BSA MIS Microhook Shaver Assembly		
	thten the rasonic	1	The Torque Wrench is etched "Handpiece this side". Slide the Torque Wrench over the tip, ensuring that etched text "Handpiece this side" faces the handpiece. Rotate clockwise until one click is heard. Remove the wrench.
5. Ins	tall the		A. Slide the assembled sheath onto the probe and align the small triangular mark on the rigid sheath to the small triangular lower mark on the handpiece.
bla	black sheath		B. Tighten by turning clockwise until the small triangular mark on the rigid sheath aligns with the upper mark, resembling a lock, on the Handpiece.



The BoneScalpel Access® Handpiece is now ready for use and can be connected to the neXus System. Refer to the neXus Console Instructions for Use for connectivity with system.



Figure 1: Fully assembled neXus® BoneScalpel Access Handpiece

neXus BoneScalpel® AccessTM Handpiece Disassembly **5.4.**

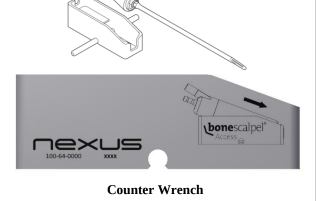
Table 9: BSA MIS 20mm Blade Disassembly **BSA MIS 20mm Blade Disassembly** Note: After disconnection of the cable connector from the generator, place the tethered cap onto the cable connector. Leave the cable connector on during cleaning, disinfection, and sterilization. Remove the tubeset from the Remove the handpiece and dispose. irrigation tubeset A. Turn the housing counter clockwise until the small triangular mark on the rigid sheath aligns with the small triangular mark on the Handpiece. Remove the housing B. Remove housing and dispose of the housing in the appropriate biohzardous sharps container.

	BSA MIS 20mm Blade Disassembly			
3.	Place handpiece into torque fixture	Counter Wrench	Insert the BoneScalpel Access Handpiece into the Counter Wrench as indicated on the drawing. The handpiece should be angled upward as indicated in the drawing.	
4.	Loosen the ultrasonic tip		Slide the torque wrench over the MIS 20mm Blade, ensuring that etched text "Handpiece this side" faces the handpiece. Rotate counterclockwise until the MIS 20mm Blade is loosened. Once the MIS 20mm Blade is loosened, remove the wrench.	
5.	Remove the tip	Hand loosen and remove the tip by turning counterclockwise.	Hand loosen the MIS 20mm Blade by turning couterclockwise. Once fully loosened, remove the MIS 20mm Blade and dispose in the appropriate biohzardous sharps container.	

Table 10: BSA MIS Microhook Shaver Disassembly

BSA MIS Micro Hook Shaver Disassembly Note: After disconnection of the cable connector from the generator, place the tethered cap onto the cable connector. Leave the cable connector on during cleaning, disinfection, and sterilization. Remove the tubeset from the 1. Remove the handpiece and dispose. irrigation tubeset A. Turn the housing counter clockwise until the small triangular mark on the rigid sheath aligns with the small triangular mark on the Handpiece. Remove the housing B. Remove housing and dispose of the housing in the approppriate biohzardous sharps container.

3. Place handpiece into torque fixture



Insert the BoneScalpel Access Handpiece into the Counter Wrench as indicated on the drawing.

The handpiece should be angled upward as indicated in the drawing.

	BSA MIS Micro Hook Shaver Disassembly				
4.	Loosen the ultrasonic tip		Slide the torque wrench over the MIS Micro Hook Shaver, ensuring that etched text "Handpiece this side" faces the handpiece. Rotate counterclockwise until the		
			MIS Micro Hook Shaver is loosened. Once the MIS Microhook Shaver is loosened, remove the wrench.		
5.	Remove the tip		Hand loosen the MIS Microhook Shaver by turning couterclockwise. Once fully loosened, remove the MIS Micro Hook Shaver and dispose in the approppriate biohzardous sharps container.		
		Hand loosen and remove the tip by turning counterclockwise.			

CAUTION The reusable handpiece and accessories (counter wrench, torque wrench, and sterilization tray) must be properly decontaminated, cleaned and sterilized before each use. Refer to Section 6: Cleaning and Sterilization. Failure to do so may lead to infections, which can ultimately cause patient death.

CAUTION Single-use disposable kits are marked with the international symbol for "do not reuse - single

use only" (②). Discard these items following each surgical procedure in accordance with the health care institution protocol for biohazardous waste. Tips are to be disposed of in a biohazardous sharps container.

Remove black sheath and ultrasonic tip from the handpiece prior to cleaning and/or sterilization; otherwise, proper cleaning/sterilization may be inhibited. Refer to **Section 5: Handpiece Assembly and Disassembly** and **Section 6: Cleaning and Sterilization**.

CAUTION 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.

CAUTION Always tighten or un-tighten the probe cover by hand and without using any wrenches. Do not over-tighten the probe cover.

WARNING

6. Cleaning and Sterilization

6.1. Dispose of Single-Use Items

WARNING: Follow the health care institutions protocol for Universal Precautions for Blood Borne

Pathogens including the use of Personal Protective Equipment (PPE) when cleaning and

disinfecting reusable items after a clinical procedure.

WARNING Single-use disposable kits are marked with the international symbol for "do not reuse - single

use only" (②). Discard these items following each surgical procedure in accordance with the health care institution protocol for biohazardous waste. Tips are to be disposed of in a

biohazardous sharps container.

WARNING The single-use disposable kits are intended for one procedure only. Do not attempt to reuse,

clean or re-sterilize single-use disposable kit components.

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 health care institution procedures for disposal of biohazardous waste.

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

Table 11: neXus BoneScalpel® Access™ Handpiece – Reusable Items

Part Number	Description	
100-22-0001 BoneScalpel Access TM (BSA) Handpi		
100-64-0000	Access Handpiece Counter Wrench	
100-63-0000	Handpiece Torque Wrench	
100-71-0000	Sterilization Tray	

Misonix Inc. 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 The reusable handpiece and accessories (counter wrench, torque wrench, and sterilization

tray) must be properly decontaminated, cleaned and sterilized before each use. Refer to **Section 6: Cleaning and Sterilization**. Failure to do so may lead to infections, which can

ultimately cause patient death.

WARNING Misonix Inc. has validated all cleaning and sterilization cycles given in this manual. To

prevent transmission of disease or malfunction of the neXus® system, Misonix recommends that the procedures given in this manual for cleaning and sterilizing the neXus® Ultrasonic Surgical Aspirator System and related accessories be followed. However, other end-user validated cleaning agents, cleaning procedures, and/or sterilization cycles may be used.

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 in accordance with applicable local laws and regulations on cleaning and sterilization of reusable medical devices in the healthcare environment.

6.2. Point of Use Cleaning

Following use, flush the handpiece lumen with a minimum of 100ml of saline to clear the bore of biological debris. Then remove visible blood and biological debris from the surface of the handpiece and components.

• Misonix recommends the use of CaviWipesTM or equivalent quaternary ammonium compound surface disinfectant wipes to remove visible blood and biological debris from the surface of the handpiece and components. Please follow manufacturer's instructions for surface cleaning and disinfection of hard non-porous surfaces, including, without limitation, following the instructions for the use of personal Protection Equipment (PPE) for blood borne Pathogens. Dispose of the used wipes in accordance with the health care institution protocol and local regulations regarding the disposal of biological hazardous waste.

Place the handpiece into a tray and transport to the health care institution decontamination processing area.

- CAUTION: To avoid drying of biological soil:
 - Transport the neXus Standard Handpiece to the decontamination processing area as soon as practical after the clinical procedure for cleaning.
 - If transport to the decontamination processing area is delayed, cover the tray with a water dampened cloth or spray the tray and its contents with a pre-cleaning foam. The pre-water dampened cloth or cleaning foam will minimize the drying of biological soil and facilitate later decontamination processing. Transport the neXus Standard Handpiece to the decontamination area as soon as practical.
- CAUTION: DO NOT use saline to wet the tray and tray contents before transport to the decontamination processing area.
- CAUTION: DO NOT mix other heavy devices with the neXus Standard Handpiece during transportation to avoid damage to the handpiece.

6.3. Manual Cleaning/Washing Procedure

Table 12: Cleaning of Handpiece and Wrenches

	BoneScalpel Access TM Handpiece and Wrenches
Wash & Brush	• Disassemble the handpiece. Refer to Section 5.4 .
Diusii	 Prepare the alkaline enzymatic cleaning solution. Misonix has validated the use of ASP Enzol® or Steris Prolystica® alkaline enzymatic detergents. Misonix recommends the use of equivalent alkaline enzymatic detergents. Please follow manufacturer's instructions preparation of the detergents, including, without limitation, the use of recommended Personal Protection Equipment (PPE).
	 Misonix recommends following the health care institutions protocol for Universal Precautions for Blood Borne Pathogens including the use of Personal Protective Equipment when cleaning reusable items after a clinical procedure. Dispose of single use items in accordance with local regulations regarding the disposal of biological hazardous wipes.
	 Thoroughly wet all surfaces of the handpiece covers and wrenches with any enzymatic detergent solution, such as ASP Enzol® or Steris Prolystica® in accordance with the directions provided in the manufacturer's Instructions for Use.
	Wrenches may be fully immersed.
	• Thoroughly wet a brush with warm cleaning solution. 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. Attention should be given to hard to clean features such as crevices, channels, joints, or hard to reach areas where soil may be difficult to remove by brushing. Flush hard to reach areas using a sterile syringe filled with the enzymatic detergent in accordance with the directions provided in the manufacturer's Instructions for Use.
	• Item's exterior surface can be cleaned using a standard soft bristle cleaning brush.
Rinse	 Rinse item under warm running softened, filtered, or deionized water for a minimum of 1 minute to clear soap residue.
Dry • Drain and then dry item fully with lint-free cloth, paper, or with medic compressed air.	
	 Dispose of lint-free cloth or paper in accordance with Health care institution or Clinic practices for contaminated wastes.
Inspect	 Inspect wrenches and remove any item which shows signs of damage (cracks, gouges, fractures etc.). Mark damaged items clearly to prevent future use before disposal.
Post Cleaning	 Inspect all items for cleanliness and damage following cleaning and prior to terminal sterilization. If soil remains, repeat the cleaning and rinsing procedure using fresh warm cleaning solution.

WARNING: All Misonix reusable items must be sterilized by moist heat (steam sterilization/autoclaved)

after manual cleaning.

WARNING: Follow the health care institutions protocol for Universal Precautions for Blood Borne

Pathogens including the use of Personal Protective Equipment (PPE) when cleaning and

disinfecting reusable items after a clinical procedure.

CAUTION: Do not use ultrasonic cleaners to clean the handpiece as this method could damage the

handpiece.

CAUTION: Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder

sterilization of units during autoclaving. Refer to the Pre-cleaning step below.

CAUTION: Do not immerse the ultrasonic handpiece or the handpiece cable. These items are not sealed

against liquids and damage to equipment will result.

CAUTION: Use softened, filtered, or deionized water for diluting cleaning agents and for the final

equipment rinse. Deionized water is recommended for the final rinse, if available. Mineral residues from hard water in the final rinse step can cause water stains and/or affect cleaning

and disinfection.

Table 13: Cleaning the neXus BoneScalpel® Access™ Handpiece

BoneScalpel® Access TM Handpiece		
Wipe Cable	• Misonix recommends the use of CaviWipes® or equivalent quaternary ammonium compound surface disinfectant wipes to remove visible blood and biological debris from the surface of the handpiece and components. Please follow manufacturer's instructions for surface cleaning and disinfection of hard non-porous surfaces, including, without limitation, following the instructions for 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 waste.	
Wash & Brush	 Misonix recommends the use of ASP Enzol® or Steris Prolystica®, or equivalent alkaline enzymatic detergents. Please follow manufacturer's instructions preparation of the detergents, including, without limitation, the use of recommended Personal Protection Equipment (PPE). Misonix recommends following the health care institutions protocol for Universal Precautions for Blood Borne Pathogens including the use of Personal Protective Equipment when cleaning reusable items after a clinical procedure. Dispose of single use item in accordance with local regulations regarding the disposal of biological hazardous wipes. Wash and brush the handpiece with an enzymatic detergent such as ASP Enzol® or Steris Prolystica® in accordance with the directions provided in the manufacturer's Instructions for Use. The handpiece cannot be immersed. Brush all passages (lumen) 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. The item's exterior surface can be cleaned using a standard soft bristle cleaning brush. 	
Rinse	Rinse item under warm running water for a minimum of 1 minute to clear soap residue.	

	BoneScalpel [®] Access™ Handpiece			
Dry	• Dry item fully with absorbent towel or paper. Dispose of cloth or paper in accordance with Health care institution 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. 			
Post Cleaning	Inspect all items for cleanliness and damage following cleaning and prior to terminal sterilization.			
	 Inspect the handpiece housing and front cover for damage and signs of wear such as scratches, cracks, and chips. 			
	 Inspect the handpiece cable to ensure it is not cut or frayed. Inspect the handpiece cable connector and the tethered connector cap to ensure they are not damaged. 			
	 Do not use damaged handpieces or handpiece components. Contact Misonix Customer Service if damage is noted. 			
	 The neXus handpiece should be fully inspected for loose or missing components and tested for proper operation prior to each procedure. 			

WARNING: All Misonix reusable items must be sterilized by moist heat (autoclaved) after manual cleaning. WARNING: Follow the health care institutions protocol for Universal Precautions for Blood Borne Pathogens including the use of Personal Protective Equipment (PPE) when cleaning and disinfecting reusable items after a clinical procedure. CAUTION: Do not use ultrasonic cleaners to clean the handpiece as this method could damage the handpiece. CAUTION: Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder sterilization of units during autoclaving. Refer to the Pre-cleaning step below. CAUTION: Do not immerse ultrasonic handpiece, handpiece cable. These items are not sealed against liquids and damage to equipment will result. CAUTION: Use softened, filtered, or deionized water for diluting cleaning agents and for the final rinse of equipment. Deionized water is recommended for the final rinse if available. Mineral residues from hard water in the final rinse step can cause water stains and/or affect cleaning and disinfection. CAUTION: The tethered handpiece cable cap should be placed on the handpiece cable connector immediately after the cable is disconnected from the console to prevent damage to the connector pins and remain on during precleaning, manual cleaning, automated cleaning/disinfection, and sterilization procedures.

6.4. Automated Cleaning/Washing Procedure

Table 14: Cleaning of Reusable Components

	Handpiece and Wrenches		
Pre-Cleaning	The following should be performed on a disassembled handpiece:		
	 Remove the probe and all housing components. Refer to Section 5.4 or instructio on disassembly of the handpiece and components. 		
	 Prepare neodisher[®] MediClean forte in accordance with the directions provided the manufacturer's Instructions for Use. 		
	• Use a tight-fitting brush dipped in the prepared cleaning solution to clean the lumen of the hand- piece 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.		
	 Rinse all residual soap from the handpiece under warm running water for a minimum of one minute. 		
	 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. 		
Automated Wash and Disinfection	Attempt to align the lumen in the general direction of the water jet flow in the washer but at a		
	Process the handpiece and all reusable components and accessories using the cycle parameters, in the table below. *Durations listed are minimums acceptable. Longer durations than those specified for cleaning and disinfection are acceptable.		
Post- Cleaning	Inspect all items for cleanliness and damage following cleaning and prior to terminal sterilization.		
	 Inspect the handpiece housing and front cover for damage and signs of wear such as scratches, cracks, and chips. 		
	 Inspect the handpiece cable to ensure it is not cut or frayed. Inspect the handpiece cable connector and the tethered connector cap to ensure they are not damaged. 		
	 Do not use damaged handpieces or handpiece components. Contact Misonix Customer Service if damage is noted. 		
	The neXus handpiece should be fully inspected for loose or missing components and tested for proper operation prior to each procedure.		

must be sterilized by moist heat (autoclaved) after automated cleaning and disinfection.

WARNING: Follow the health care institutions protocol for Universal Precautions for Blood Borne Pathogens including the use of Personal Protective Equipment (PPE) when cleaning and disinfecting reusable items after a clinical procedure.

CAUTION: Do not use ultrasonic cleaners to clean the handpiece as this method could damage the

Be certain to clear debris from all internal passages by brushing. Failure to do so may hinder sterilization of units during autoclaving. Refer to the Pre-cleaning step below.

Automated cleaning-disinfection is not the final step prior to use. All Misonix reusable items

WARNING:

CAUTION:

CAUTION: Do not immerse ultrasonic handpiece, handpiece cable. These items are not sealed against

liquids and damage to equipment will result.

CAUTION: Use softened, filtered, or deionized water for diluting cleaning agents and for the final rinse of

equipment. Deionized water is recommended for the final rinse if available. Mineral residues from hard water in the final rinse step can cause water stains and/or affect cleaning and

disinfection.

WASHER-DISINFECTOR:

Note 1: Misonix recommends using a washer-disinfector designed and labeled for washing and

disinfecting medical devices or meeting local regulations or regulatory standards and

guidance.

Note 2: For health care institutions and health care practitioners in the EEU, Misonix recommends the

use of a washer-disinfector meeting the requirements of the ISO 15883 Washers-

Disinfectors, Parts 1-5.

CAUTION: The cable tethered cap should be on the cable connector during precleaning and automated

cleaning and disinfection procedures.

Table 15: Automated Wash 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), or equivalent
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

(*Durations listed are minimum acceptable. Longer durations than those specified for cleaning and disinfection are acceptable.)

6.5. Sterilizing by Steam Autoclave

Sterilization Methods and terminology are based on current editions of ANSI/AAMI ST81 and EN ISO 17664 standards.

CAUTION:

For all sterilization protocols listed below, always ensure the tethered cap is placed securely on the cable connector to protect the connector during sterilization.

WARNING:

Follow the health care institution protocol for using a chemical or biological indicator with every sterilization load to ensure proper sterilization conditions of time, temperature, and saturated steam penetration.

WARNING:

Misonix Inc. has validated all cleaning and sterilization cycles given in this manual. To prevent transmission of disease or malfunction of the neXus® system, Misonix recommends that the procedures given in this manual for cleaning and sterilizing the neXus® Ultrasonic Surgical Aspirator System and related accessories be followed. However, other end-user validated cleaning agents, cleaning procedures, and/or sterilization cycles may be used.

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 in accordance with applicable local laws and regulations on cleaning and sterilization of reusable medical devices in the healthcare environment.

CAUTION:

Misonix does not recommend "FLASH" sterilization. Misonix has not validated "FLASH" sterilization.

CAUTION:

Water Quality for Steam Generation:

Poor steam quality may impair the sterilization process. For this reason, various norms (European standard EN 285 and the United States standard ANSI/AAMI ST79) recommend maximum impurity levels for steam feed water of autoclaves and sterilizer used in the medical field. Misonix recommends using water of a quality that conforms to the norms, the health care institution validated specifications for water quality, or otherwise using deionized water to generate steam for moist heat sterilization.

Table 16: neXus BoneScalpel® AccessTM Handpiece – Reusable Items

Part Number	Description	
100-22-0001 BoneScalpel Access™ (BSA) Handpiece		
100-64-0000	Access Handpiece Counter Wrench	
100-63-0000	Handpiece Torque Wrench	
100-71-0000	Sterilization Tray	

Table 17: Items Required for Sterilization

Items for Sterilization		
Item	Comment	
Autoclave	Misonix has validated several autoclave cycles for the sterilization of the Standard Handpiece reusable components. However, the specific autoclave design and performance can affect the efficacy of the process. Health care institutions should verify the process used, employing the actual equipment and personnel in place. Responsibility for verification of the sterilization process lies directly with the health care institution.	
Chemical or Biological Steam Sterilization IndicatorsFollow the health care institution protocol for using a chemical or biolo indicator with every sterilization load to ensure proper sterilization cond time, temperature, and saturated steam penetration.		
Sterilization Wrap	Misonix has validated several autoclave cycles with sterilization wrap for maintenance of package integrity post sterilization. Misonix has validated the cycles using Kimberly Clark KC 300 KIMGUARD or Kimberly Clark KC 600 KIMGUARD. Equivalent base weight 300 or 600 sterilization wraps, including Halyard Quick Check 300 or 600, can be used. The chart on sterilization parameters indicates the specific wrap used for the cycle. Follow the Association for the Advancement of Medical Instrumentation (AAMI) and the Association of perioperative Registered Nurses (AORN or EORNA) recommended guidelines for appropriate wrapping configurations.	

Handpiece **DISASSEMBLED** using Misonix Sterilization Tray: Probe, Tubing, and Housing should be REMOVED.

Table 18: Sterilization Parameters for Handpiece DISASSEMBLED using Misonix Sterilization Tray

	132°C (270°F)	134-137°C (274-279°F)	
	Items placed in Misonix Sterilization Tray #100-71-0000	Items placed in Misonix Sterilization Tray #100-71-0000	
Configuration	Tray wrapped in base weight 300 Sterilization Wrap (Kimberly Clark / KIMGUARD KC300, and Halyard Quick Check H300, or equivalent).	Tray wrapped in base weight 300 Sterilization Wrap (Kimberly Clark / KIMGUARD KC300, and Halyard Quick Check H300, or equivalent).	
Cycle Prevacuum		Prevacuum	
Preconditioning Pulses	4	4	
Minimum Exposure Time 4 minutes*		3 minutes*	
Minimum Dry Time 30 minutes		30 minutes	

^{*}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 Inc reusable medical devices are able to sustain such sterilization cycles.

Handpiece **DISASSEMBLED** without Sterilization Tray, Items Wrapped: Probe, Tubing, and Housing should be REMOVED.

Table 19: Sterilization Parameters for Handpiece DISASSEMBLED without Sterilization Tray, Items

Wrapped

Wiappeu	132°C (270°F)	134-137°C (274-279°F)	134°C (273°F)
Configuration	Items wrapped, NO TRAY Items wrapped in base weight 300 or 600 Sterilization Wrap (Kimberly Clark / KIMGUARD KC300 or KC600, and Halyard Quick Check H300 or H600, or equivalent).	Items wrapped, NO TRAY Items wrapped in base weight 300 or 600 Sterilization Wrap (Kimberly Clark / KIMGUARD KC300 or KC600, and Halyard Quick Check H300 or H600, or equivalent).	Items wrapped, NO TRAY Items wrapped in base weight 300 or 600 Sterilization Wrap (Kimberly Clark / KIMGUARD KC300 or KC600, and Halyard Quick Check H300 or H600, or equivalent).
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

^{*}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 Inc reusable medical devices are able to sustain such sterilization cycles.

CAUTION

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

6.6. Deviations to Cleaning, Sterilization, Decontamination Instructions

Misonix Inc. has validated all cleaning and sterilization cycles given in this manual. To prevent transmission of disease or malfunction of the neXus® system, Misonix recommends that the procedures given in this manual for cleaning and sterilizing the neXus® Ultrasonic Surgical Aspirator System and related accessories be followed. However, other enduser validated cleaning agents, cleaning procedures, and/or sterilization cycles may be used. 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 in accordance with applicable local laws and regulations on cleaning and sterilization of reusable medical devices in the healthcare environment.

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

6.7. Transportation, Storage, and Handling Prior to Use

- Transport wrapped equipment to storage in a manner to prevent damaging the sterile barrier.
- Refer to the sterilization wrap manufacturer's Instructions for Use for maximum shelf-life information.
- Store wrapped equipment in a controlled environment to avoid temperature and moisture extremes.
- Avoid excessive handling or wrapped equipment to avoid damage to the wrapping and cause a breach in the sterile barrier.
- Inspect the wrapping for openings, cuts, pinholes, and other damage that would indicate a possible breach in the sterile barrier prior to use. Do not use the equipment if the wrapping is damaged. Clean and sterilize the equipment again.

6.8. Expected Life, Reusable Components

The sterilization life of handpiece components is listed below is based on cleaning and sterilization in accordance with the instructions in this manual. Life estimates may be affected by rough handling, damage, wear due to vigorous cleaning, or using alternative cleaning and sterilization procedures.

Table 20: Reusable Component Estimated Re-Use Life

Estimated Sterilization Life		
Item Number of Steam Sterilization Cycles		
Handpiece with attached cable	200 cycles	
Wrenches: Handpiece/counter wrench and torque wrench	300 cycles	

CAUTION The specified reuse life considers wear and tear due to cleaning and sterilization. Damage or wear caused by misuse in treatments will affect life of components.

WARNING The single-use disposable kits are intended for one procedure only. Do not attempt to reuse, clean or resterilize single-use disposable kit components.

7. Handpiece Specifications

Table 21: Handpiece Specifications

neXus BoneScalpel [®] Access™ Handpiece	
Operating frequency	Nominal: 23 kHz (22.0 – 24.5 kHz)
Cable length	15' 4.57 m
Dimensions	4.6" L (without probe) x 0.9" D 13 cm x 2.0 cm
Weight with tip	4.6 oz. 130 g

8. 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.

WARNING

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

9. Important Notice

Please contact Misonix with any questions regarding the specifications, use, sterilization, limitations, or maintenance of the NeXus® Ultrasonic System:

Misonix, Inc.

Web: www.misonix.com

Phone: +1.631.694.9555 / 1-800-694-9612

Fax: +1.631.694.9412 **Address:** 1938 New Hwy

Farmingdale, NY 11735,

U.S.A.

By returning any material to Misonix, Inc. 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, Inc. The correct return address should read as follows:

Misonix, Inc.
Medical Service Department
RMA #_____
1938 New Hwy
Farmingdale, NY 11735
U.S.A.

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

9.1. Trademark Information

- Misonix[®], neXus[®], and neXus BoneScalpel[®] are registered trademarks of Misonix, Inc.
- BoneScalpel AccessTM is a trademark of Misonix, Inc.
- ASP Enzol® and Prolystica® are registered trademarks of STERIS Corporation.
- CaviWipesTM is a trademark of Metrex Research LLC.

