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Congratulations on the addition of the Epic Pro diode laser system to your practice!

This User Manual is designed to help you become familiar with the operation and functions of the laser system. Read it carefully before using the laser clinically, follow all safety instructions and cautions, and always have it accessible for use as a reference.

We appreciate your commitment to better care and know your investment in this technology is a sound foundation for the successful growth of your practice.

BIOLASE Team
Overview

The Epic Pro diode laser system is a surgical device designed for a wide variety of surgical and oral soft-tissue procedures, and dental whitening.

Epic Pro utilizes a solid state diode as a semiconductor source for invisible infrared radiation. The energy is delivered to the treatment site via a flexible fiber connected at one end to the laser source and the other end to the Handpiece. Various types of single use, disposable Tips are designed and optimized for different applications. The device is activated by means of a wireless Footswitch.

This prescription device is indicated for professional use only by licensed dental practitioners. The use of this device requires proper clinical and technical training. This manual provides instructions for those professionals that have completed the appropriate training.

When used and maintained properly, the Epic Pro will prove a valuable addition to your practice. Please contact BIOLASE Customer Service at 1-800-321-6717 in the U.S. for any service needs. If you are located outside the USA, please contact your BIOLASE-authorized distributor.
The Epic Pro device with surgical laser operation (Automatic Power Control), used in contact or non-contact mode, is indicated for dental soft-tissue indications including incision, excision, vaporization, ablation, hemostasis, or coagulation of intraoral and extra-oral soft-tissue (including marginal and interdental gingiva and epithelial lining of free gingiva); examples include:

- Frenectomy
- Debridement of diseased epithelial lining
- Frenotomy
- Incisions and draining of abscesses
- Biopsy
- Tissue retraction for impressions
- Operculectomy
- Papillectomy
- Implant recovery/ uncovered
- Vestibuloplasty
- Gingivectomy
- Excision of lesions
- Gingivoplasty
- Exposure of unerupted teeth/partially erupted teeth
- Gingival troughing
- Leukoplakia
- Crown lengthening
- Removal of hyperplastic tissues
- Removal of granulation tissue
- Laser-assisted flap surgery
- Sulcular debridement (removal of diseased or inflamed soft-tissue in the periodontal pocket)

The Epic Pro device with surgical laser operation, used in contact or non-contact mode, is intended for use in general surgery for incision/excision, vaporization, ablation and coagulation of soft-tissue.
2 Indications for Use

The Epic Pro device with dental laser operation, used in contact or non-contact mode, is intended for incision, excision, vaporization, ablation, hemostasis, or coagulation of intraoral and extra-oral soft-tissue (including marginal and interdental gingiva and epithelial lining of free gingiva); examples include:

- Frenectomy
- Debridement of diseased epithelial lining
- Frenotomy
- Incisions and draining of abscesses
- Biopsy
- Tissue retraction for impressions
- Operculectomy
- Papillectomy
- Implant recovery/uncovery
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- Gingivectomy
- Excision of lesions
- Gingivoplasty
- Exposure of unerupted teeth/partially erupted teeth
- Gingival troughing
- Leukoplakia
- Crown lengthening
- Removal of hyperplastic tissues
- Removal of granulation tissue
- Laser-assisted flap surgery
- Hemostasis of donor site
- Pulpotomy
- Treatment of aphthous ulcers
- Pulpotomy as an adjunct to root canal therapy
- Sulcular debridement (removal of diseased or inflamed soft-tissue in the periodontal pocket)
- Light activation of bleaching materials for teeth whitening
# 3 Specifications

## 3.1 GENERAL
- **Dimension (W x L x H)**: 7.25in x 4.5in x 6.5in (18.4cm x 11.4cm x 16.5cm)
- **Weight**: 4lbs (1.8kg)

## 3.2 ELECTRICAL
- **Operating Voltage**: 100V-240V ~ at 1.2A/0.5A
- **Frequency**: 50/60 Hz
- **External Fuses**: None
- **Main Control**: Main Power Switch
- **Remote Interruption**: Remote Interlock Connector
- **Disable Control**: Emergency Stop Button

## 3.3 LASER
- **Laser Classification**: IV (4)
- **Medium**: Semi-Conductor Diode
- **Wavelength**: 980 ± 20nm
- **Max Power Output**: 25W
- **Power Accuracy**: ± 20%
- **Power Modes**: Continuous, Pulse Modulation
- **Fiber Tips Diameter**: 300µm, 400µm
- **Pulse Duration**: 10µs-100ms
- **Pulse Energy**: 0.01ms – 20ms
- **Pulse Repetition Rate**: Up to 20kHz (for reference)
- **NOHD**: 4.77 meters
- **Beam Divergence**: 0.22
- **Standard Fiber Cable Length**: 6 feet (1.8 meters)

## 3.4 OTHER LIGHT SOURCES
- **Aiming Beam**: Laser Diode, Max 5mW 450-680nm, Class 3R
4.1 SYSTEM COMPONENTS

The Epic Pro laser system includes the following:

1. Laser Console
2. Fiber Management Module
3. Fiber Optic Cable
4. Surgical Handpiece (2)
5. Assorted Surgical Tips (non-initiated and pre-initiated)
6. Wireless Footswitch
7. Protective Eyewear (3)
8. Power Supply/Adapter
10. Disposable Screen Protectors (30)
11. Remote Interlock Connector
12. AAA Batteries (2-pack)
13. Welcome Kit (Includes User Manual and Laser Warning Sign)

WARNING: No modification of this equipment is allowed.

4.2 GENERAL

The Epic Pro laser system consists of three components:

1. Console and fiber management module
2. Reusable fiber delivery system (Fiber Optic Cable, reusable surgical Handpiece, fiber Tips)
3. Wireless Footswitch
CONSOLE
The console, supported by a snap-on table base, is portable, contains the diode laser and all the elements which power the laser. A touch screen Graphical User Interface (GUI) allows the user to navigate from screen to screen, select procedures, enter and/or change settings, view system status, etc. (Figure 4.1)

FIBER MANAGEMENT MODULE
This module is a separate component that clips onto the console. It provides a storage channel for the fiber and a holder for the Handpiece (Figure 4.1).

FIBER OPTIC CABLE
The Fiber Optic Cable transmits the laser radiation from the laser console through the Handpiece and fiber Tips to the target tissue.

SURGICAL HANDPIECE
Part of the delivery system, the surgical Handpiece is re-useable and detachable from the fiber. It is the conduit through which the laser energy flows from the Fiber Optic Cable to the Tip.

FIBER TIPS
Fiber Tips deliver the energy generated by the laser and transmitted through the Fiber Optic Cable to the treatment site.
4 Equipment Description

WIRELESS FOOTSWITCH
Epic Pro is activated by a Footswitch designed to work using wireless technology. It is powered by two (2) AAA batteries (Refer to section 11 for instructions on how to install/replace the Footswitch batteries).

The Epic Pro laser will only emit laser energy when the Footswitch is pressed while the laser is in **Ready** mode.

The Footswitch and laser are shipped from the factory already paired.

The Footswitch is protected by a metal cover. To access, press down on the cover (Figure 4.2).

Three separate LED indicator lights are displayed on the rear panel of the Footswitch.

<table>
<thead>
<tr>
<th>LED Indicator</th>
<th>Status</th>
<th>Status Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amber</td>
<td>Blinking or Solid</td>
<td>A critical error or short has occurred; the relevant error message will appear on the tablet screen.</td>
</tr>
<tr>
<td>Blue</td>
<td>Solid</td>
<td>The Footswitch is connecting to the laser, or has successfully paired with the laser.</td>
</tr>
<tr>
<td>Green</td>
<td>Blinking</td>
<td>The Footswitch batteries are low; replace the batteries.</td>
</tr>
<tr>
<td>Green</td>
<td>Solid</td>
<td><strong>ON:</strong> the Footswitch is pressed while paired to the laser; <strong>OFF:</strong> the footswitch is not pressed, or the system is in <strong>Sleep</strong> mode, or the batteries are dead.</td>
</tr>
</tbody>
</table>
5 Contraindications, Warnings, and Precautions

5.1 CONTRAINDICATIONS
All clinical procedures performed with Epic Pro must be subjected to the same clinical judgment and care as with traditional techniques. Patient risk must always be considered and fully understood before clinical treatment. The clinician must completely understand the patient’s medical history prior to treatment and exercise caution for general medical conditions which might contraindicate a local procedure; such conditions may include, but are not limited to, allergy to local or topical anesthetics, heart disease, lung disease, bleeding disorders, sleep apnea or an immune system deficiency or any medical conditions or medications that may contraindicate use of certain light/laser type sources associated with this device. Medical clearance from the patient’s physician is advisable when doubt exists regarding treatment. Epic Pro treatment may be contraindicated for patients suffering from photodermatosis, photosensitive patients, and patients who are taking photosensitive drugs. Conditions such as pregnancy or pacemakers are not known as contraindications for laser light energy.

5.2 WARNINGS AND PRECAUTIONS

PRESCRIPTION STATEMENT
U.S. Federal Law restricts this device to sale by or on the order of a dentist or dental specialist.

TRAINING
Only licensed professionals who have reviewed and understood this User Manual should use this device. BIOLASE assumes no responsibility for parameters, techniques, methods, or results. Physicians must use their own clinical judgment and professionalism in determining all aspects of treatment, technique, proper power settings, interval, duration, etc.

EYEWEAR
Doctor, patient, assistant and all others inside or entering the operatory must wear appropriate laser protection eyewear for the 980nm wavelength, OD 5 or greater (OD5+) whenever the laser is in user. Prior to use, inspect eyewear for pitting and/or cracking. Do not use if damaged.

CAUTION: Always check eyewear specifications as etched on the glasses themselves to ensure they offer the protection required by the laser wavelength.

ANESTHESIA
In soft-tissue cases anesthesia may not be required, but patients should be closely monitored for signs of pain or discomfort at all times. If such signs are present, adjust settings, apply anesthesia, or cease treatment if required.
TREATMENT, TECHNIQUE, AND SETTINGS
Use your clinical judgment to determine all aspects of treatment including, but not limited to, the laser treatment protocol, technique, power settings, pulse duration and interval settings, mode of operation, as well as the accessories (e.g. tip type) and other procedural requirements. Closely observe and monitor clinical effects and use your judgment to determine the clinical parameters and approach for the treatment; make appropriate power, pulse length, and interval adjustments to compensate for varying tissue composition, density, and thickness. Always start treatment at the lowest power setting for the specific indication and increase as required. BIOLASE assumes no responsibility for parameters, techniques, methods, or results.

ADJACENT STRUCTURES
Epic Pro is designed to remove soft tissues. Therefore, always be aware of adjacent structures and substructures during use. Be extremely careful not to inadvertently penetrate or ablate underlying or adjacent tissues. Do not direct energy towards hard tissue such as tooth or bone. Do not direct energy towards amalgam, gold or other metallic surfaces. Do not direct energy towards cements or other filling materials. Exercise extreme caution when using this device in areas such as pockets, cavities or channels such as third molar sockets, where critical structures (i.e. nerves, vessels) could be damaged. Do not proceed with using the laser if visibility is limited in these areas.

SUCTION
Use high-speed suction as required to maintain a clear field of vision during treatment. Do not use the Epic Pro if you cannot clearly see the treatment site.

PLUME REMOVAL

CAUTION: Laser plume may contain viable tissue particulates.

Special care must be taken to prevent infection from the laser plume generated by vaporization of virally or bacterially infected tissue. Ensure that appropriate protective equipment, including high-speed suction to remove the plume, appropriately filtered masks, and other protective gear is used at all times during the laser procedure.

CLINICAL ENVIRONMENT
Only use this device in clinical environments that observe proper standard aseptic techniques with all oral procedures.
6 Safety with the Epic Pro

Failure to comply with precautions and warnings described in this user manual may lead to exposure to dangerous optical radiation sources. Please comply with all safety instructions and warnings.

**WARNING:** No modification of this equipment is allowed.

### 6.1 SAFETY INSTRUCTIONS

**LASER WARNING:** Emits invisible and/or visible laser radiation. Avoid eye or skin exposure to direct or scattered radiation. Do not use the Laser System in any manner other than as described in this User Manual; do not use the Laser System if you suspect it is not functioning properly; do not use procedures other than those specified herein as it may result in hazardous radiation exposure.

Follow these safety instructions before and during treatments:

- When the laser is in use, all operatory entrances must be marked with an appropriate warning sign (included)

- Do not operate in the presence of explosive or flammable materials. Flammable anesthetics or oxidizing gases such as nitrous oxide (N2O) and oxygen should be avoided. Solvents of adhesives and flammable solutions used for cleaning and disinfecting should be allowed to evaporate before the laser is used. Attention should also be drawn to the danger of ignition of endogenous gases

- Doctor, patient, assistant, and all others in or entering the operatory must wear appropriate laser protection eyewear. Always ensure that the laser protective eyewear is appropriate for the 980nm wavelength (OD 5+). Prior to use always inspect the eyewear for pitting and/or cracking; do not use if damaged

- Do not look directly into the beam or at specular reflections

- Never direct or point the beam at a person’s eyes

- Always place the system into **Standby** mode before exchanging the fiber delivery system, Handpieces or disposable Tips

- Always ensure the proper laser parameters are set before the Epic Pro is used in a clinical procedure

- Before leaving the unit unattended, return to the Home screen so the device cannot accidentally emit laser energy. The laser can also be turned off at the power switch

- Do not use the Epic Pro if the Fiber Optic Cable is damaged or broken
6.2 SAFETY CLASSIFICATION

The following safety classifications are applicable to the device:

- Laser Radiation – Class 4
- Aiming Beam – Class 3R
- Type of protection against electrical shock – Class 2
- Degree of protection against electrical shock – Type BF Class 1
- Not protected against water ingress – Ordinary Equipment
- Not suitable for use in presence of flammable anesthetic mixture
- Operation Mode – Continuous Wave and Pulse Modulation
- Wireless Footswitch – IPX6

Laser Warning: Do not open the console housing at any time. Danger from optical radiation may exist.

Laser Warning: Do not aim the laser at metallic or reflective surfaces, such as surgical instruments or dental mirrors. If aimed directly at these surfaces the laser beam will be reflected and create a potential hazard.

Caution: This laser system has been designed and tested to meet the requirements of electromagnetic, electrostatic, and radio frequency interference standards. However, the possibility of electromagnetic or other interference may still exist. Relocating the device may help to eliminate the interference.

Caution: High temperatures produced in the normal use of this laser equipment may ignite some materials (e.g., cotton wool when saturated with oxygen); solvents or adhesive and flammable solutions used for cleaning and disinfecting should be allowed to evaporate before the laser is activated.

Caution: Be aware that the metal / cannula on the Tips may become hot during use. Avoid touching any tissue with the cannula.
6.3 SAFETY FEATURES

SYSTEM MONITOR
The system monitors the emergency stop switch, remote key, wireless Footswitch connection, and output power. An error in any one of these will stop the system. The text display will indicate the type of error. Operation will not resume until the error is cleared.

POWER SWITCH
The laser console can be switched ON (I) or OFF (O) using the Power Switch on the side of the console (Figure 6.1).

CAUTION: Use only the power supply module and power cord supplied with the Epic Pro laser system.
6 Safety with the Epic Pro

ACCESS KEY CODE
The Access Key Code (888) prevents unauthorized use of the system. It is activated every time the system is turned ON with the Power Switch.

WIRELESS FOOTSWITCH
The Epic Pro will not emit laser energy until the user presses down on the Footswitch while the laser is in Ready mode. The Footswitch is designed to work using wireless technology.

REMOTE INTERLOCK
The Epic Pro console is equipped with a remote interlock feature that links a door switch (optional) with the Epic Pro console and will prevent the laser from firing when the sensor is triggered. To utilize this safety feature, wire the remote interlock plug to a multi-conductor wire and insert it into the opening at the side of the laser console (Figure 6.3). Attach the other end of the wire to a door switch/sensor; if the closed door is opened, the connection to the switch is deactivated, immediately stopping the laser from firing. To override this feature, don’t connect the plug to a door switch.

EMERGENCY STOP
Press the red Emergency Laser Stop button located on the bottom left of the front panel to instantly turn off the laser console (Figure 6.4). The laser will immediately stop, the screen will go dark, and the system will need to be completely powered OFF and then rebooted to resume operation.
7 Installation and Set-Up

7.1 FACILITY REQUIREMENTS

- Electrical Supply (100-240V ~): 1.2A/0.5A, 50/60Hz
- Environmental Requirements:
  Temperature: 15-30 °C
  Humidity: 10-90%, Non-condensing

**CAUTION:** Do not position this equipment so that it is difficult to pull the plug from the power source. To avoid the risk of electric shock, this equipment must only be connected to a supply mains with protective earth.

**CAUTION:** Do not stack this device with other equipment, as it could affect proper operation. Always monitor this and any adjacent equipment to ensure they are operating normally.

7.2 SYSTEM SETUP

The Epic Pro is designed to be transportable between operatories using the built-in handle. Prior to use, the laser must be properly set up as described in this user manual to ensure safe and efficient functionality.

- Attach the base to the console by sliding it from front to back until it clicks into place
- Press the lever of the fiber management module and push it onto the right side of the console until it clicks into place (Figure 7.1)
7 Installation and Set-Up

- Place the unit on a flat surface in a clean, dry, and well-ventilated area
- Verify the power switch is in the OFF (O) position
- Connect the power supply/adapter to the laser console by orienting the power supply plug as indicated by the symbol on the connector panel and inserting it fully into the console’s power jack until snug (Figure 7.2). Plug the power cord into the power supply and insert the distal end of the cord into a wall outlet. Install the remote interlock, if used.

Figure 7.2

CAUTION: Never touch the exposed pins and the patient simultaneously whenever the fiber management module is not attached to the console, but the laser is ON (I), to avoid electric shock to the patient.

7.3 SURGICAL DELIVERY SYSTEM

CAUTION: The Fiber Optic Cable is detachable from the console; it cannot be autoclaved but must always be disinfected between patients by following the procedure outlined in section 10.3.

- Verify the power switch is in the OFF (O) position
- Remove the protective cap on the end of the fiber connector (the cap is reusable and needed when storing the fiber)
- Attach the Fiber Optic Cable to the console by aligning the center of the fiber SMA connector with the center of the port at the rear of the console, gently inserting the connector into the port until snug, and turning it clockwise until secure (Figure 7.3)
- To disconnect the Fiber Optic Cable from the console, turn the SMA connector counterclockwise; when loose, pull gently from the connector port

Figure 7.3

CAUTION: Do not connect or disconnect the fiber while the laser is turned ON. Only connect or disconnect the Fiber Optic Cable when the laser is turned OFF.

CAUTION: Never bend the Fiber Optic Cable at a sharp angle, as it can break the internal optical fiber.

CAUTION: Avoid direct contact with the proximal end of the Fiber Optic Cable SMA connector. Contamination of the connector may result in laser damage.
7.4 SURGICAL HANDPIECE ASSEMBLY

**CAUTION:** Handpieces are not sterile when sold and must be sterilized prior to initial use, and cleaned and sterilized between patients. Refer to Section 10 for complete instructions on cleaning and sterilization.

- Remove the protective red cap from the distal end of the fiber shaft
- Connect the Handpiece to the Fiber Optic Cable assembly by pushing the Handpiece onto the fiber shaft until it clicks on and is secured (Figures 7.4, 7.5), without twisting

![Diagram of Fiber Shaft and Protective Cap](image1)

![Diagram of Handpiece and Collar](image2)

Figure 7.4: Connecting the Handpiece to the Fiber Optic Cable assembly

Figure 7.5: Surgical Handpiece Assembly fully assembled
7 Installation and Set-Up

Disconnect the Handpiece from the Fiber Optic Cable assembly (Figure 7.6):

1. Take the Handpiece body in one hand and the shaft in the other,
2. Push the two buttons on the fiber shaft,
3. Gently pull the Handpiece by the collar to separate it from the fiber shaft without twisting.

![Figure 7.6: Disconnecting the Handpiece from the Fiber Optic Cable](image)

7.5 SINGLE-USE TIPS

Fiber Tips are single-use accessories available in 300μm and 400μm core diameters and in different lengths; Tips are available non-initiated, for procedures that do not involve cutting tissue, such as hemostasis, or pre-initiated, for all ablative procedures. The choice of which Tip to use is based on the type of procedure/treatment mode. Refer to Section 9 for a details on procedures and related presets and recommended Tips.

**CAUTION:** Tips are for one-time use only to avoid cross-contamination and are designed to withstand only a single sterilization cycle. Always visually inspect the Tip prior to use to make sure it is free of debris or damage

**CAUTION:** Be aware that the metal / cannula on the Tips may become hot during use. Avoid touching any tissue with the cannula.

To connect the Tip, **first connect the Handpiece to the Fiber Optic Cable**, then insert the Tip firmly into the distal end of the Handpiece as far as it will go, and tighten by turning clockwise until snug (Figure 7.7). Bend the metal cannula according to the specific procedure requirements (Figure 7.10).

Remove the fiber Tip by twisting it counterclockwise (Figure 7.8).
**7 Installation and Set-Up**

**CAUTION:** To deliver proper laser operation, do not install Tips when the Handpiece is not connected to the Fiber Optic Cable.

Figure 7.7: Insert the fiber Tip into the Handpiece *(only when the Handpiece is connected to the Fiber Optic Cable)* and twist clockwise until snug.

Figure 7.8: Remove the fiber Tip by twisting the Tip counterclockwise.

Correct

Incorrect

Figure 7.9: Installing the Tip

Correct

Incorrect

Figure 7.10: Bending the Tip cannula
7. Installation and Set-Up

7.6 WHITENING HANDPIECE (OPTIONAL ACCESSORY)

The Whitening Handpiece (Figure 7.11) is a re-usable accessory intended for light activation of bleaching materials in tooth whitening procedures. It is equipped with disposable non-sterile protective shields for single patient use (Figure 7.12) and requires disinfection before and after each patient treatment. **This Handpiece cannot be sterilized in the autoclave.** Doing so will damage its internal optics. For instructions on how to disinfect the Whitening Handpiece, refer to Section 10.

The disposable shield fits over the head of the Handpiece and must be discarded after one-time use to avoid cross-contamination between patients. Always wipe with alcohol prior to use.

The area of Laser Energy Output for the Whitening Handpiece is 35mm x 8mm = 2.8cm² Spot Size.

To connect the Handpiece to the Fiber Optic Cable, push it onto the fiber shaft until it clicks on and is secured.

To disconnect the Handpiece from the Fiber Optic Cable:

- Take the Handpiece body in one hand and the shaft in another
- Press both buttons at the base of the fiber shaft
- Gently pull the Handpiece from the collar without twisting to separate

### 7.6.1 WHITENING GEL

The Whitening Handpiece is designed to work with LaserWhite™ 20 Whitening Gel to achieve teeth whitening. This item is sold separately, as a kit in packs of five. Detailed step-by-step instructions, contraindications, precautions, and warnings for teeth whitening are provided with the LaserWhite 20 Whitening Gel Kit. Please read the instructions carefully before proceeding with the procedure.
7.7 PAIRING THE FOOTSWITCH AND LASER

Epic Pro ships with the laser and Wireless Footswitch already paired. However, if the wireless connection should fail at any time, pairing will need to be re-established by taking the following steps:

1. Make sure two (2) AAA fully charged batteries are installed in the Footswitch.

2. Go to the System Settings Menu by pressing the settings icon at the bottom right of the screen.

3. Select **FOOT PEDAL** and follow the instructions displayed on the next series of screens.
   - Place the Footswitch on a flat surface and release the Footswitch cover (Figure 7.13)
   - Use a coin or a flat-head screw driver to remove the battery compartment cover on the side of the Footswitch (Figure 7.14)
   - Press down on the Footswitch and hold. While continuing to press down the Footswitch, screw the battery compartment cover back in place (Figure 7.15); continue to hold down the Footswitch for an additional 5 seconds, or until both the blue and green lights on the Footswitch light up, then release
   - Both the blue and green LEDs will be on indicating the system is ready to pair. Select “Begin Pairing” to initiate the pairing process (Figure 7.16). If the pairing operation times out, the screen will display a message “Pairing Timed Out” and offer the option to “Try Again”

If pairing is successful, a “Pairing Successful” message appears on the screen and only the blue LED will light up on the Footswitch; press “Return to Home” to begin using the laser.

If replacing an existing Footswitch, first insert two (2) new AAA batteries in the battery compartment, then follow the pairing steps noted above.
8 Operating Instructions

8.1 EPIC PRO INTERACTIVE DISPLAY
The Epic Pro interactive display allows for touch screen operation. All functions are activated by gently touching the appropriate button on the display. Apply the disposable screen protector provided before use to protect the screen from water and debris.

8.2 ICONS - DEFINITIONS
The Epic Pro Graphic User Interface (GUI) is the main part of the system control. It communicates with the user through the interactive touchscreen display and is designed to provide easy and intuitive interaction with the laser system while performing clinical procedures. Buttons on each screen allow easy navigation of the GUI.

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>💘</td>
<td>Favorites</td>
</tr>
<tr>
<td>🏡</td>
<td>Home</td>
</tr>
<tr>
<td>🛠️</td>
<td>Settings</td>
</tr>
<tr>
<td>↩️</td>
<td>Back Button (Return to Previous Screen)</td>
</tr>
<tr>
<td>⚛️</td>
<td>Advanced Mode Indicator</td>
</tr>
<tr>
<td>📰</td>
<td>Signal Strength Indicator (Wireless FS)</td>
</tr>
<tr>
<td>🚨</td>
<td>Settings modified and saved</td>
</tr>
<tr>
<td>🔒</td>
<td>Settings modified but not saved</td>
</tr>
<tr>
<td>📈</td>
<td>FS Battery Strength Indicator</td>
</tr>
<tr>
<td>🔧</td>
<td>Decrease setting/Increase setting</td>
</tr>
<tr>
<td>🛠️</td>
<td>Service</td>
</tr>
</tbody>
</table>

Figure 8.1
8 Operating Instructions

8.3 SYSTEM START-UP AND FUNCTIONS

Turn the power switch at the side of the console to the ON (I) position. The system will boot up and display two screens in quick succession – the first displaying the “BIOLASE” logo followed by one showing the Epic Pro name; after 3 seconds, the system will display the “Welcome” screen.

8.3.1 LOGIN

Enter the three digit default access code (888) using the keypad on the touchscreen. If the incorrect code is keyed, an “X” appears in the window; wait 3 seconds for the “X” to clear, then re-enter the correct code (Figure 8.3).

After logging in, the Epic Pro will display the Home Screen (Figure 8.4).

While in the Login Screen, the Epic Pro cannot emit laser energy.

CAUTION: Always check the aiming beam before using the laser; the shape of the beam on a surface should be symmetrical. This indicates that the delivery system (Fiber Optic Cable, Handpiece) is working properly. This check should be performed with a non-initiated Tip only.
8.3.2 HOME SCREEN

The HOME screen offers the user four operational options (Figure 8.9): 

1. **Quick Start**: accesses the main treatment screen; select the treatment parameters and start treatment immediately.

2. **Procedures**: displays the Procedures menu; select from a series of factory pre-programmed dental treatments and settings.

3. **Favorites**: presents a list of the user’s favorite procedures/settings; select the chosen procedure/settings and begin treatment.

4. **Settings**: allows modification of user options and settings.

8.3.3 BASIC AND ADVANCED

The Epic Pro laser system offers two levels of operation, **Basic and Advanced**; usage is based on specific procedure requirements and the user’s level of training and experience with the laser system.

- **Basic**: Only some of the pre-set parameters for any given procedure may be modified by the user.
- **Advanced**: Allows the user to modify all the pre-set values for a procedure, even from a pre-set procedure screen.

The user may switch between levels by pressing the Advanced mode Indicator on the bottom right of any procedure screen, or through the Settings menu.
8.3.4 TREATMENT MODES OF OPERATION

The Epic Pro soft-tissue laser is a versatile system that can be used for a variety of surgical and dental soft-tissue procedures. It offers four treatment modes of operation:

- Continuous Wave (CW) Mode (Figure 8.5)
- Automatic Power Control (APC™) (Figure 8.6).
- Pulse Mode (Figure 8.7)
- Super Thermal Pulse (STP™) (Figure 8.8)

While on the main operating screen, the user has the option to toggle between treatment modes of operation, i.e., APC, CW, Pulse and STP modes. Once a selection is made, the screen will display the treatment parameter adjustment controls for the corresponding treatment mode.

8.3.5 CONTINUOUS WAVE (CW) MODE:

CW treatment mode allows the user to adjust the power and the total time of firing, which can be set to OFF to enable continuous firing with no time limit (Figure 8.5).

When CW is selected while in Basic user mode, the Epic Pro delivers continuous laser power output. This can be increased or decreased incrementally by using the corresponding arrows, or by using the slider at the bottom of the power window. The output power is adjustable from 0.1 to 25W.

The timer, or the length of time the laser will fire continuously without stopping, can either be turned off, or adjusted in increments up to 99 seconds.
8.3.6 AUTOMATIC POWER CONTROL (APC):

Epic Pro has a treatment mode of operation called Automatic Power Control (APC) in which the laser, in continuous wave mode, converts light energy to thermal power at the tip (Figure 8.6). The cutting effect in contact mode with a pre-initiated Tip is achieved not by the direct action of laser power, but rather by the thermo-mechanical action of a "hot tip." Therefore, Tip temperature is a very important parameter and is adjusted automatically to maintain a uniform temperature, resulting in constant and reliable cutting, independent of speed of movement, tissue type, etc.

In the Advanced level of operation, the peak "not to exceed" upper power limit is adjustable from 3W to 25W. Thermal Level is adjusted automatically when Power Limit is modified within these parameters.

![Figure 8.6](image.png)

**CAUTION:** APC procedures utilize only pre-initiated tips. **Do not operate the Epic Pro in APC mode with a non-initiated Tip**; the system will not be able to reach the target temperature and will temporarily operate at “Not to Exceed” power, which is much higher than the power required for treatment, before displaying an error message.

**CAUTION:** The pulse width, peak and average power, and timer duration parameters are not adjustable in the APC operating treatment mode.
8.3.7 PULSE MODE:

When the Pulse operating treatment mode is selected, Epic Pro delivers repetitive pulses. The peak laser power output can be increased or decreased by moving the power slider right or left. The peak output power is adjustable from 3W up to 150W, depending on the Pulse duration selected (Figure 8.7).

In Pulse mode, pulse width (duration), peak power, average power, and the timer, or the length of time the laser will fire continuously without stopping, are all adjustable.

For each pulse duration (10μs – 100ms), there is a corresponding set of peak powers, and for each peak power there is a corresponding average power setting.

When pulse width is adjusted, the corresponding default (minimum) peak power and (maximum) average power are automatically selected and displayed by the system software.

![Figure 8.7](image-url)
8.3.8 SUPER THERMAL PULSE:

In Super Thermal Pulse (STP) treatment mode, the device converts pulsed laser energy into pulses of thermal power at the Tip, resulting in deeper penetration, more precise cutting, and less trauma to tissue. These power bursts are emitted at a preset pulse width and repetition rate, while maintaining a precise thermal level range at the tip. If the measured Tip temperature is higher than the desired thermal level, the peak power of the laser pulses will be decreased to bring it down; if the measured Tip temperature is lower than desired, the peak power will be increased accordingly. In this way, dynamic control of the peak power ensures the stable temperature range of the Tip and, consequently, consistent thermal effect on the target tissue.

Figure 8.8

CAUTION: STP procedures utilize only pre-initiated Tips. Do not operate the Epic Pro in STP mode with a non-initiated Tip; the system will not be able to reach the target temperature and will temporarily operate at “Not to Exceed” power, which is much higher than the power required for treatment, before displaying an error message.
8.3.9 QUICK START

Selecting Quick Start on the Home screen displays the main operating screen, with APC as the default mode of operation (Figure 8.9). Quick Start allows the user to adjust treatment parameters, select the treatment mode of operation (CW, APC, Pulse, or STP), choose a Tip suitable for the procedure, save custom treatment settings, and, ultimately, fire the laser using the displayed parameters. This screen also offers the user the option to view system settings or procedure info, return to the previous screen, or return to the Home screen. A graphical indicator displays the status of the wireless Footswitch connection, while the laser defaults to Standby mode.

The main operating screen allows the user to toggle between APC, CW, Pulse, and STP modes, as needed. Once a selection is made, the screen will display the treatment parameter adjustment controls for the corresponding mode of operation for treatment.
8 Operating Instructions

8.3.10 TIP SELECTION

From the main treatment screen, the user has the option of choosing a treatment Tip by pressing the “Select Tip” button; the screen will display a selection of non-initiated and initiated Tips (Figure 8.10).

After Tip selection is completed, the system returns to the main operating screen which will now display the Tip chosen (Figure 8.11).

If an attempt is made to fire the laser without first having selected a Tip, a message stating “Please Select Tip” will appear on the screen and the laser will not fire (Figure 8.12). Press the Select Tip button and choose a Tip as described above, then press on the Footswitch to activate the laser.

In the Advanced level of operation, if a pre-initiated Tip is selected in APC or STP modes, the system software returns to the main operating screen and displays a graph that plots the set Tip temperature and real-time laser power during emission.

---

![Figure 8.10](image1)

![Figure 8.11](image2)

![Figure 8.12](image3)
The set Tip temperature and set power limit are displayed on the graph (Figure 8.13). While the laser is emitting laser energy, the program will plot the Tip temperature feedback and real-time power value to achieve the set Tip temperature.

CAUTION: Laser power output above the level shown on the touch screen may cause unwanted tissue damage. The optical power output and pulse width must remain within +/- 20% of the selected values while the laser is emitting energy.

8.3.11 TIP WARNING

In APC and STP modes, if a pre-initiated Tip inserted into the Handpiece is not properly initiated, the Tip's initiation has degraded after use, or a non-initiated Tip is inserted, a message will appear on the screen when attempting to fire the laser instructing the user to replace the Tip (Figure 8.14).
8.3.12 PROCEDURE SELECTION

When Procedures is selected at the Home screen, seven treatment categories are displayed; the user is able to select any one of these to access a specific pre-set procedure (Figure 8.15). However, settings may be adjusted by the user, as needed, on the Main operation screen.

The recommended settings for each indication are preprogrammed at the factory and cannot be deleted. However, they may be adjusted by the user, as needed, on the Main operating screen.

Once a pre-set procedure is chosen, the screen will display the treatment parameters, corresponding mode (APC, CW, Pulse, or STP), and Tip selection established for treatment (Figure 8.15).
8 Operating Instructions

8.3.13 CREATING A FAVORITES PROCEDURE

Any changes to the pre-programmed settings of a clinical procedure can be saved to Favorites by pressing the heart button on the right side of the main operating screen (Figure 8.17):

- A full keyboard and text input field are displayed; the input field is limited to 30 characters, and may be all alpha, all numeric, or a combination of the two. Type in a unique name for the procedure, or leave as the default name set by the system.

- Press Save. The settings are now stored in Favorites for use when needed. The program returns to the main treatment screen with the new saved Favorite name displayed on the banner and the Favorites button is highlighted (Figure 8.18).

- The customized procedure is now saved and can be accessed at any time from the Favorites screen.

![Select Save Favorite Button](Figure 8.17)

![User Favorite Name Displayed in Banner](Figure 8.18)
If a Favorites procedure is selected, but any of the parameters that appear on the screen are changed, the locked lock will change to an unlocked lock. The new settings may be saved by pressing the Favorites icon at the bottom right of the display; the name of the procedure will remain the same unless the modified settings are saved as a new Favorites.

To permanently delete a Favorites entry, press and hold the procedure name displayed on the screen for 4 seconds. A screen requesting confirmation will always appear before the procedure is deleted (Figure 8.19).

While in the Favorites selection screen, Epic Pro is in Standby and cannot emit laser energy.
8.3.14 SELECTING A FAVORITES PROCEDURE

To access the Favorites screen, select Favorites on the Home screen; all custom procedures previously saved will be displayed (Figure 8.21). This list is created solely by the user; it is empty by default when the system is first installed.

Select from the procedures listed. Once a selection is made, the Main operating screen is displayed with the saved parameters ready for use. Navigate through the full list of Favorites by using the up and down arrows.

8.3.15 READY MODE

Epic Pro will only emit laser energy when the Footswitch is pressed while the console is in Ready mode. Press the Standby button on the bottom left of the screen to shift from Standby to Ready.

To return to Standby, press the Ready button on the bottom right of the screen.
8 Operating Instructions

CAUTION: The aiming beam is on only when the laser is in Ready mode or when adjusting the brightness of the beam through the Settings menu. If the aiming beam is not visible in either instance, remove the Handpiece and confirm the beam is actually on by shining the end of the Fiber Optic Cable on a plain, non-reflective surface. **DO NOT** look directly at the output end of the Fiber Optic Cable. If the aiming beam is not on, turn off the laser console, then remove and re-install the Fiber Optic Cable (see Section 7.3). If the aiming beam is still not detected, turn the laser console OFF and call BIOLASE Service.

**LASER WARNING:** Never point the laser at a person's eyes. All persons present in the operatory must wear appropriate protective eyewear when the laser is in use.

8.3.16 FIRING THE LASER

Pressing the wireless Footswitch while the laser is in **Ready** mode will cause the laser to fire; a beeping sound indicates that laser energy is present.

While the laser is firing, the main treatment screen disables all operator options, such as mode and/or power setting adjustments, Tip selection, etc. The laser will continue to fire until the Footswitch is no longer pressed. Once the Footswitch is disengaged, the laser will stop emitting laser energy. Press the Ready button to go to **Standby**.

Figure 8.22

Laser in Ready mode, foot pedal not pressed

Foot pedal pressed/ laser firing

Figure 8.22
8.3.17 SYSTEM SETTINGS MENU

Pressing the Settings icon allows the user to make changes to the system parameters (Figure 8.23); this menu can be reached from any screen displaying this Settings icon:

- Adjust Pilot Aiming Beam
- Adjust Sleep Timeout
- Toggle Sort Order Modes
- Enter Wireless Footswitch Pairing Screen
- Enter Language Selection
- Adjust LCD Screen Brightness
- Adjust System Volume
- Import User Procedures & Settings
- Enter Service Mode
- Export User Procedures & Settings

Figure 8.23
8.3.18 SYSTEM INFORMATION AND SOFTWARE UPDATES:

System information, such as the system serial number, software revision, GUI configuration, BootLoader version, and hardware revision are displayed by pressing the system information icon located in the upper right hand corner of any screen (Figure 8.24).

This screen also allows the user to implement the following changes (Figure 8.24).

**Update Software:** From time to time, BIOLASE may provide updated software with new functionalities by supplying the user with a programmed SD card.

- Insert the card into the SD card slot and press the “Update Software” button to start the upgrade process.
- The screen will display a confirmation screen before initiating the update (Figure 8.25)
  - If “Yes” is selected, the system will begin the update
  - If “No” is selected, the system will revert to the System Information Screen
8 Operating Instructions

If a valid application software file is detected on the SD card, the update is initiated. The system screen
will temporarily turn off. When completed, the system will reboot to the splash screen logo and Welcome
screen. The system beep is audible during the update process.

If no SD card is inserted, the SD card is improperly formatted, or the application software file on the
SD card is corrupted, the program will display an error message after the Update Software button is
selected by the operator (Figure 8.26). If this occurs, replace the SD card with a known good card and
a genuine copy of the application software file to be uploaded.

Update GUI: BIOLASE will provide the user with a programmed SD card whenever an update to the
GUI is necessary.

• Insert the card into the SD card slot and press the “Update GUI” button to start the upgrade process.
• The screen will display a confirmation screen before initiating the update (Figure 8.27).
  • If “Yes” is selected, the system will begin the update.
  • If “No” is selected, the system will revert to the System Information Screen

Figure 8.26

Figure 8.27
If a valid GUI configuration file is detected on the SD card, the program will begin the update process. The system screen will temporarily turn off and then reboot to a blue screen. The system memory is erased and then replaced with the new GUI configuration file from the SD card. A status bar at the bottom of the screen measures progress, while a number value on the top right corner displays the percentage completed (Figure 8.28). When finished, the system will reboot to the splash screen logo and Welcome screen. Updating the GUI takes approximately 20-25 minutes to complete.

If no SD card is inserted, the SD card is improperly formatted, or the GUI configuration file on the SD card is corrupted, the program will display an error message after the Update GUI button is selected by the operator (Figure 8.29). If this occurs, replace the SD card with a known good card and a genuine copy of the GUI configuration file to be uploaded.
Update Procedures: BIOLASE will provide the user with a programmed SD card if any updates to the preset treatment parameters have made.

- Insert the card into the SD card slot and press the “Update Procedures” button to start the upgrade process
- The screen will display a confirmation screen before initiating the update (Figure 8.30)
  - If “Yes” is selected, the system will begin the update
  - If “No” is selected, the system will revert to the System Information Screen

If a valid factory procedures configuration file is detected on the SD card, the program will begin the software update. The system will display a blue screen briefly, then display a status of “Update Complete” when the routine is done. Upon completion, the system will reboot to the splash screen logo and Welcome screen. Updating the factory procedures takes approximately 5 seconds to complete (Figure 8.31).
If no SD card is inserted, the SD card is improperly formatted, or the procedures configuration file on the SD card is corrupted, the program will display an error message after the Update Procedures button is selected by the operator (Figure 8.32). If this occurs, replace the SD card with a known good card and a genuine copy of the factory procedures configuration to be uploaded.

CAUTION: Please contact BIOLASE Customer Care immediately if any of the above update processes results in an error message.
8.3.19 SERVICE MODE:

The Epic Pro software includes a service mode which is password-protected and may only be accessed by trained service personnel. To return to the previous screen, press the “Back” button.

8.3.20 EXPORT AND IMPORT OF SETTINGS AND FAVORITES

Exporting settings to SD card: To preserve Favorite procedures and GUI settings when returning the Epic Pro to the BIOLASE Service Center for repair, export the information to an external storage device by inserting an Epic Pro SD card into the card slot on the left side of the console. Go to Settings and select Export to begin the process. Before proceeding, a screen will appear which requires the user to confirm the export command or cancel the operation (Figure 8.33). If cancel is selected, the system will return to the Settings screen; if export is confirmed, the program will begin to download data to the SD card.

While the export operation is in process, the screen will display a busy status followed by the results of the export operation. A screen will appear either confirming the export was successful, or that it failed (Figure 8.34). In either case, press OK to return to the main treatment screen.

![Figure 8.33](image)

![Figure 8.34](image)
8 Operating Instructions

**Importing settings from SD card:** To transfer Favorites procedures and System settings to your Epic Pro, insert the Epic Pro SD card containing the information into the card reader slot on the left side of the console. Go to Settings and select Import to begin the process.

Before proceeding, a screen will appear which requires the user to confirm the import command or cancel the operation (Figures 8.35). If cancel is selected, the program will return to the Settings screen; if import is confirmed, the program will begin to download data to your Epic Pro console.

While the import operation is in process, the screen will display a busy status followed by the results of the import operation. A screen will appear either confirming the export was successful, or that it failed (Figure 8.36). In either case, press OK to return to the main treatment screen.

**8.4 TURNING THE EPIC PRO OFF**

To turn the Epic Pro off, return to the Home screen by pushing the Home icon, then press the Power Switch on the side of the console to the OFF position (O).
8.5 SYSTEM FLOW CHART

Figure 8.36
The settings chosen for a procedure will contribute to its overall success. Recommended settings have been pre-programmed into Epic Pro at the factory for different procedures in CW, APC, Pulse, and STP treatment modes. These settings may be modified, within specific parameters, and saved as Favorites.

Always use clinical judgment when making adjustments to power, pulse repetition rate, and treatment mode as may be necessary to compensate for variations in tissue composition, density, and/or thickness.

### NOTE:
N/A means the parameter is not directly modifiable by the user, but may be visible on the screen.

## 9.1 RESTORATIVE

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Treatment Regime</th>
<th>Thermal Level (Range)</th>
<th>Power Limit (Range)</th>
<th>Average Power (Range)</th>
<th>Pulse Width (Range)</th>
<th>Pulse Energy (Range)</th>
<th>Timer (Preset Value)</th>
<th>Tip Type</th>
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</thead>
<tbody>
<tr>
<td>Troughing</td>
<td>STP</td>
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<td>1.6 W (1.6-3.6W)</td>
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<td>N/A</td>
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<td>Soft-Tissue Crown Lengthening</td>
<td>APC</td>
<td>2 (1-5)</td>
<td>10W</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>PI PRO 4</td>
</tr>
<tr>
<td>Gingivoplasty</td>
<td>STP</td>
<td>4</td>
<td>5.6W (3.2-6.4W)</td>
<td>N/A</td>
<td>1 ms</td>
<td>N/A</td>
<td>N/A</td>
<td>PI PRO 4</td>
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## 9.2 AESTHETIC

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<th>Thermal Level (Range)</th>
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<th>Pulse Width (Range)</th>
<th>Pulse Energy (Range)</th>
<th>Timer (Preset Value)</th>
<th>Tip Type</th>
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<td>N/A</td>
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<td>Pre-initiated</td>
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</tr>
<tr>
<td>Gingival Contouring</td>
<td>APC</td>
<td>2 (1-5)</td>
<td>10W</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>PI PRO 4</td>
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<td></td>
<td></td>
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<td>Micro-Coagulation</td>
<td>Pulse</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>3 J (1.75-3.75)</td>
<td>Single Pulse</td>
<td>E4</td>
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<td></td>
<td></td>
<td></td>
<td></td>
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## 9.3 IMPLANT

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<th>Pulse Energy (Range)</th>
<th>Timer (Preset Value)</th>
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<td>Implant Uncovering</td>
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<td>N/A</td>
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<td></td>
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<td></td>
<td></td>
<td>Pre-initiated</td>
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<tr>
<td>Soft-Tissue Contouring</td>
<td>STP</td>
<td>3</td>
<td>5.6W (3.2-6.4W)</td>
<td>6.4W</td>
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<td>N/A</td>
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<td></td>
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## 9.4 PERIO

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<th>Tip Type</th>
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<td>Soft-Tissue Pocket Reduction</td>
<td>APC</td>
<td>2 (1-5)</td>
<td>10W</td>
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<td>N/A</td>
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<td>N/A</td>
<td>PI PRO 4</td>
</tr>
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<td></td>
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<td>Pre-initiated</td>
<td></td>
</tr>
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<td>Sulcular Debridement</td>
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<td>N/A</td>
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<td>2 ms (1-10ms)</td>
<td>N/A</td>
<td>10 s</td>
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# 9 Presets

## 9.5 SURGICAL

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<th>Tip Type</th>
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<tr>
<td>Biopsy</td>
<td>STP 8</td>
<td>5.6W (3.2-6.4W)</td>
<td>N/A</td>
<td>1 ms</td>
<td>N/A</td>
<td>N/A</td>
<td>PI PRO 4 Pre-initiated</td>
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</tr>
<tr>
<td>Frenectomy</td>
<td>STP 8</td>
<td>5.6W (3.2-6.4W)</td>
<td>N/A</td>
<td>1 ms</td>
<td>N/A</td>
<td>N/A</td>
<td>PI PRO 4 Pre-initiated</td>
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<tr>
<td>Operculectomy</td>
<td>STP 8</td>
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<td>N/A</td>
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<td>Hemostatis</td>
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<td>10s</td>
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## 9.6 ENDO

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<th>Treatment Mode (Range)</th>
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<tr>
<td>Pulpotomy</td>
<td>CW N/A</td>
<td>N/A</td>
<td>0.1W (0.1-0.5W)</td>
<td>N/A</td>
<td>N/A</td>
<td>60s</td>
<td>E4 Non-initiated</td>
<td></td>
</tr>
</tbody>
</table>

## 9.7 ORAL LESIONS

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Treatment Mode (Range)</th>
<th>Thermal Level (Range)</th>
<th>Power Limit (Range)</th>
<th>Average Power (Range)</th>
<th>Pulse Width (Range)</th>
<th>Pulse Energy (Range)</th>
<th>Timer (Preset Value)</th>
<th>Tip Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aphthous Ulcers</td>
<td>Pulse</td>
<td>N/A</td>
<td>N/A</td>
<td>1.6W (1-3W)</td>
<td>1 ms</td>
<td>N/A</td>
<td>30s</td>
<td>E4 Non-initiated</td>
</tr>
<tr>
<td>Herpetic lesion</td>
<td>Pulse</td>
<td>N/A</td>
<td>N/A</td>
<td>1.6W (1-3W)</td>
<td>1 ms</td>
<td>N/A</td>
<td>30s</td>
<td>E4 Non-initiated</td>
</tr>
</tbody>
</table>
10 Cleaning, Disinfection, and Sterilization

10.1 CLEANING THE CONSOLE

Use the screen protectors provided to keep water and debris from contaminating the touchscreen, and use a disinfectant to wipe down the console and Handpiece Holder after each procedure. Do not use bleach or abrasive cleansers.

10.2 CLEANING AND STERILIZATION PROCEDURES, HANDPIECE AND TIPS

**CAUTION:** Handpiece and Tips must be cleaned and sterilized prior to first-time use.

Handpieces are reusable and must be cleaned and sterilized between patients to avoid cross-contamination.

**Tips are single-use only** to avoid cross-contamination and are designed to withstand only a single sterilization cycle; discard in a biohazard medical waste sharps container after use.

The contamination control suggested for the Epic Pro Surgical Handpiece and Tips is steam sterilization. However, before sterilization, the Epic Pro reusable Handpiece should be MANUALLY cleaned per the following procedure.

**CLEANING MUST BE PERFORMED WITHIN A MAXIMUM OF 1 HOUR AFTER THE PROCEDURE AND ALWAYS PRIOR TO STERILIZATION.**

**STEP 1 - CLEANING PROCESS (HANDPIECE)**

The cleaning process is intended to remove blood, protein and other potential contaminants from the surfaces and crevices of reusable accessories. This process may also reduce the quantity of particles, microorganisms and pathogens present. Cleaning should be performed prior to sterilization and must be conducted by qualified office personnel trained to perform the procedure.

Wear protective latex gloves when handling the contaminated delivery system.

1. After each use, carefully remove the tip from the Handpiece and discard in a biohazard medical waste sharps container.
2. Carefully disconnect the Handpiece from the Fiber Optic Cable by taking the Handpiece body in one hand and the shaft in the other, then pressing both buttons at the base of the fiber shaft and gently pulling the Handpiece by the collar to separate it from the shaft without twisting.
3. Prepare any commercially available surgical instrument detergent/enzymatic cleaning solution with a pH of 7.0, such as Enzol® or similar enzymatic presoak and cleaner, per the manufacturer’s instructions. (Follow the manufacturer’s instructions for disposal of used solution.)
4. Rinse the Handpiece under running lukewarm tap water (22 – 43°C) for a minimum of 10 seconds to remove gross soil.
5. Wrap the Handpiece in a piece of gauze that has been soaked in the cleaning solution; leave it wrapped in the gauze for a **minimum of 10 minutes**.

6. Unwrap the Handpiece from the gauze and use a soft-bristled brush dipped in the cleaning solution to gently scrub it for **at least 15 seconds**.

7. Rinse the Handpiece under running lukewarm tap water (22-43°C) for a **minimum of 10 seconds** and then dry with a lint-free cloth.

8. Visually inspect the Handpiece for any residual soil. If necessary, repeat steps 5 - 7 until all residual soil is removed.

**STEP 2 - STEAM STERILIZATION FOR SURGICAL HANDPIECE, SINGLE USE TIPS**

The steam sterilization process is intended to destroy infectious microorganisms and pathogens.

**CAUTION:** Always perform the procedure immediately after cleaning and prior to use and only use FDA-cleared (USA) or CE-marked (Europe) sterilization accessories, i.e., sterilization pouch and autoclave tray. The product packaging is not suitable for steam sterilization.

- Place the Handpiece and fiber Tips in separate single-wrap, self-seal autoclave pouches
- Place on an autoclave tray; do not stack other instruments on top of the pouches
- Place the tray inside the autoclave chamber and set the appropriate cycle as recommended in Figure 10.1
- Once the cycle is completed, remove the tray and let each sterilized item cool and dry. **The Handpiece and Tips must remain in the sterilization pouches until used in order to maintain sterility**
- Reassemble the Handpiece by pushing the Handpiece onto the fiber shaft until it clicks into place and is secured (See Section 7)

<table>
<thead>
<tr>
<th>Type of Sterilizer</th>
<th>Temperature</th>
<th>Minimum Time</th>
<th>Drying Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gravity Displacement</td>
<td>121°C (250°F)</td>
<td>30 minutes</td>
<td>15 – 30 minutes</td>
</tr>
<tr>
<td></td>
<td>132°C (270°F)</td>
<td>15 minutes</td>
<td></td>
</tr>
<tr>
<td>Dynamic-Air-Removal (Pre-Vacuum)</td>
<td>132°C (270°F)</td>
<td>4 minutes</td>
<td>20 – 30 minutes</td>
</tr>
<tr>
<td></td>
<td>134°C (EU only)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 10.1
10 Cleaning, Disinfection, and Sterilization

10.3 DISINFECTING THE FIBER OPTIC CABLE

Wear protective latex gloves when handling the contaminated delivery system.

Always disinfect the fiber between patients by wiping it completely with an appropriate disinfecting solution such as Cavicide™ or a similar quaternary ammonium compound product (containing 20% alcohol or less), and follow the manufacturer’s instructions. **DO NOT AUTOCLAVE.**

10.4 DISINFECTING THE WHITENING HANDPIECE

The Whitening Handpiece is sold with disposable non-sterile protective shields. **Neither the Handpiece nor the shields can be sterilized in the autoclave.**

To disinfect the Whitening Handpiece, wipe it down with gauze and isopropyl alcohol.

Always wipe the disposable shield with alcohol prior to use. Discard after single use; these are intended for one-time use only and should never be reused to prevent cross-contamination.
11.1 ANNUAL CALIBRATION

Epic Pro should be calibrated every 2 years in order to maintain the required accuracy of output power versus displayed power. Calibration can be performed at a certified depot repair facility. Call BIOLASE Service at 1-800-321-6717 or your Authorized Service representative for instructions on how to return the laser for service.

11.2 CHANGING THE WIRELESS FOOTSWITCH BATTERIES

The wireless Footswitch is powered by two AAA batteries. When the batteries are low, the green LED on the Footswitch cover will blink and a warning message will appear on the screen indicating that the batteries need to be replaced.

To replace the batteries:

1. Close the Footswitch cover (Figure 11.1).
2. Use a coin to remove the battery compartment cover on the side of the Footswitch by turning it counter-clockwise (Figure 11.2).
3. Remove the depleted batteries from the compartment and insert two new batteries (Figure 11.3).
4. Replace the battery compartment cover; turn clockwise to secure.

In the unlikely event that pairing of the laser console and Footswitch is disrupted when the batteries are replaced, the message "FOOTSWITCH NOT DETECTED" will appear on the touchscreen; re-establish pairing by following the instructions outlined in Section 7.7.

Replace both batteries at the same time; do not mix old and new batteries. **Old batteries must be discarded as regulated; do not throw them in a trash bin.**

To ensure the longevity of the Footswitch's battery power, only use BIOLASE-supplied batteries as replacements (P/N 6400463); these are industrial-grade and under normal use have a longer life than conventional AAA batteries.
11.3 FIBER CABLE LENS INSPECTION AND CLEANING

11.3.1 LENS INSPECTION PROCEDURE

The lens is at the distal end of the Fiber Optic Cable (Figure 11.4); this is where debris may accumulate with use. To inspect the lens, take the following steps:

1. Remove the Handpiece from the fiber shaft.
2. Inspect the distal end of the fiber using a magnifier or loupes with at least 10X magnification (Figure 11.5).
3. Inspect the lens.

11.3.2 CLEANING THE LENS:

If debris or dark spots are observed on the lens, clean the window using the procedure outlined below.

1. Always use a new, unused Microbrush to clean the lens provided (Figure 11.6).
2. Dip the Microbrush in isopropyl alcohol.
3. Gently rub the lens surface to remove debris or dark spots (Figure 11.7).
4. Repeat as needed using a new Microbrush each time until no residue appears on the swab.

CAUTION: If blemishes appear below or inside the lens, contact BIOLASE or your authorized service representative to order a replacement lens.
If the system detects a system fault while on the main treatment screen, an Error button appears in the bottom left hand corner of the screen. Pressing the button will display information related to the error (Figure 12.1). It will not be possible to enter **Ready** or **Standby** modes until the fault is resolved.

If after following the directions on the screen the error does not clear, please call your local BIOLASE Service Representative for assistance.

<table>
<thead>
<tr>
<th>Error</th>
<th>Message</th>
<th>Reason(s)</th>
<th>Fix</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Door interlock disconnected</td>
<td>Error Code: 1 Insert door interlock</td>
<td>Remote interlock connector not inserted on left side panel of system</td>
<td>Insert remote interlock</td>
</tr>
<tr>
<td>2 Laser overheating</td>
<td>Error Code: 2 Laser overheating. Wait until cools down. Reduce room temperature if possible</td>
<td>Laser temperature detected greater than 50°C</td>
<td>Allow system to cool for 5-10 minutes. If problems persist, contact the manufacturer for evaluation</td>
</tr>
<tr>
<td>3 Laser temperature too low</td>
<td>Error Code: 3 Laser temperature sensor malfunction. Please contact the manufacturer</td>
<td>Laser temperature detected less than 10°C</td>
<td>Allow system to reach ambient room temperature. If problems persist, contact the manufacturer for evaluation</td>
</tr>
<tr>
<td>4 Power transistor overheating</td>
<td>Error Code: 4 Device overheating. Please contact the manufacturer</td>
<td>Power transistor temperature detected greater than 75°C</td>
<td>Allow system to cool for 5-10 minutes. If problems persist, contact the manufacturer for evaluation</td>
</tr>
</tbody>
</table>
## Troubleshooting

<table>
<thead>
<tr>
<th>Error</th>
<th>Message</th>
<th>Reason(s)</th>
<th>Fix</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>Power transistor temp too low</td>
<td>Error Code 5: Internal temperature sensor malfunction. Please contact the manufacturer</td>
<td>Power transistor temperature detected less than 10°C</td>
</tr>
<tr>
<td>6</td>
<td>Optical Feedback Out of Range</td>
<td>Error Code 6: Power Calibration Failed. Please contact manufacturer</td>
<td>Optical feedback detected a value out of the calibrated range</td>
</tr>
<tr>
<td>7</td>
<td>Voltage Reference</td>
<td>Error Code 7: Voltage reference failure</td>
<td>Voltage feedback out of range</td>
</tr>
<tr>
<td>8</td>
<td>Footswitch pressed while not in ready state</td>
<td>Error Code 8: Release footswitch. If it is not pressed, replace the Footswitch</td>
<td>Footswitch pressed down while entering <strong>Ready</strong> Mode from <strong>Standby</strong> Mode</td>
</tr>
<tr>
<td>9</td>
<td>Fiber Not Detected</td>
<td>Error Code 9: Fiber not detected. Insert treatment fiber to resume</td>
<td>Fiber not inserted into system</td>
</tr>
<tr>
<td>10</td>
<td>Laser current exceeds set value</td>
<td>Error Code 10: Internal error with current setting</td>
<td>Current feedback detected current out of acceptable range</td>
</tr>
<tr>
<td>11</td>
<td>APC Temperature Feedback</td>
<td>Error Code 11: Cannot achieve set temperature</td>
<td>Temperature feedback setting was not able to be verified by software within 5 seconds of emission</td>
</tr>
</tbody>
</table>
## Troubleshooting

<table>
<thead>
<tr>
<th>Error</th>
<th>Message</th>
<th>Reason(s)</th>
<th>Fix</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 Calibration Data Corrupted</td>
<td>Error Code 12: System calibration data stored in internal memory has been compromised</td>
<td>System calibration data could not be verified for integrity</td>
<td>Contact the manufacturer</td>
</tr>
<tr>
<td>13 Optical module overheating</td>
<td>Error 13: Optical module overheating</td>
<td>Optical temperature detected greater than 50°C</td>
<td>Allow system to cool for 5-10 minutes. If problems persist, contact the manufacturer for evaluation</td>
</tr>
<tr>
<td>14 Optical module temperature too low</td>
<td>Error 14: Optical module temperature too low</td>
<td>Optical temperature detected less than 10°C</td>
<td>Allow system to reach ambient room temperature. If problems persist, contact the manufacturer for evaluation</td>
</tr>
<tr>
<td>16 Factory Procedure Data Corrupted</td>
<td>Error 16: Factory procedure data corrupted. Please contact the manufacturer</td>
<td>System memory corrupted</td>
<td>Copy valid procedures file onto SD card and attempt to Update Procedures through System Information screen</td>
</tr>
<tr>
<td>17 System Clock Failure</td>
<td>Error 17: System Clock Failure</td>
<td>Main controller board system clock not operating correctly</td>
<td>Contact the manufacturer</td>
</tr>
<tr>
<td>18 Cap Voltage Error</td>
<td>Error 18: Main controller board capacitor voltage out of range</td>
<td>Main controller board capacitor's voltage is out of acceptable range</td>
<td>Contact the manufacturer</td>
</tr>
</tbody>
</table>

Figure 11.2
Appendix A Labels

PRODUCT IDENTIFICATION LABEL:

LOCATION:
Bottom of laser console

LASER APERTURE WARNING:

LOCATION:
Back of console

FCC AND IC ID LABEL:

LOCATION:
Bottom of wireless Footswitch
# Appendix A Labels

<table>
<thead>
<tr>
<th>MANUFACTURER</th>
<th><img src="image1.png" alt="Image" /></th>
</tr>
</thead>
<tbody>
<tr>
<td>CATALOG/PART NUMBER</td>
<td><img src="image2.png" alt="Image" /></td>
</tr>
<tr>
<td>PRODUCT SERIAL NUMBER</td>
<td><img src="image3.png" alt="Image" /></td>
</tr>
<tr>
<td>DATE OF MANUFACTURE</td>
<td><img src="image4.png" alt="Image" /></td>
</tr>
</tbody>
</table>

**LASER WARNING SYMBOL:** Indicates the system contains a laser

**LOCATION:** Back of laser console

**FCC DECLARATION OF CONFORMITY:** Electromagnetic interference is under limits approved by the Federal Communications Commission

**RX ONLY**

(US) Federal Law restricts this device to sale by or on the order of a dentist or other licenced dental practitioner

**LASER EXPLANATORY AND CERTIFICATION LABEL:**

**LOCATION:** Back of console

**ELECTRICAL SHOCK HAZARD**
Appendix A Labels

ETL LABEL:
Meets Product Safety Standards

LOCATION:
Back of console

NON-IONIZING RADIATION:
Device includes an international radiator (Bluetooth)

REFER TO USER MANUAL

TYPE BF APPLIED PART:
The applied part is not conductive to the patient

FCC COMPLIANCE NOTICE:

LOCATION:
Bottom of wireless Footswitch

FOOTSWITCH PRODUCT ID LABEL:

LOCATION:
Bottom of Footswitch

INGRESS PROTECTION CODE:
The Footswitch is water-resistant, protected splashes of water

IPX6
### Appendix A  Labels

<table>
<thead>
<tr>
<th>Label</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INVISIBLE AND VISIBLE LASER RADIATION</strong></td>
<td></td>
</tr>
<tr>
<td><strong>SINGLE USE ONLY – DO NOT REUSE</strong></td>
<td></td>
</tr>
<tr>
<td><em>(SINGLE-USE ACCESSORIES)</em></td>
<td></td>
</tr>
<tr>
<td><strong>WEEE (WASTE ELECTRICAL AND ELECTRONIC):</strong></td>
<td></td>
</tr>
<tr>
<td>Recycle lithium ion battery as regulated. Do not throw in trash bin.</td>
<td></td>
</tr>
<tr>
<td><strong>THIS END UP</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
<tr>
<td><strong>FRAGILE: HANDLE WITH CARE</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
<tr>
<td><strong>KEEP DRY</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
<tr>
<td><strong>TEMPERATURE LIMITATIONS</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
<tr>
<td><strong>HUMIDITY LIMITATIONS</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
<tr>
<td><strong>ATMOSPHERIC PRESSURE LIMITATIONS</strong></td>
<td>Outer carton (shipping box)</td>
</tr>
</tbody>
</table>

**Temperature Limitations**
-20°C to +60°C

**Pressure Limitations**
106kPa to 50kPa
## Appendix B: Tip Guide

### NON-INITIATED

<table>
<thead>
<tr>
<th>Tip</th>
<th>Name</th>
<th>Diameter</th>
<th>Length (mm)</th>
<th>Application</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="E4-4" /></td>
<td>E4-4</td>
<td>400µm</td>
<td>4</td>
<td>Surgical</td>
<td>7400016</td>
</tr>
<tr>
<td><img src="image" alt="E4-7" /></td>
<td>E4-7</td>
<td>400µm</td>
<td>7</td>
<td>Perio</td>
<td>7410003/7400019</td>
</tr>
<tr>
<td><img src="image" alt="E4-9" /></td>
<td>E4-9</td>
<td>400µm</td>
<td>9</td>
<td>Perio</td>
<td>7410019</td>
</tr>
<tr>
<td><img src="image" alt="E3-4" /></td>
<td>E3-4</td>
<td>300 µm</td>
<td>4</td>
<td>Surgical</td>
<td>7400017</td>
</tr>
<tr>
<td><img src="image" alt="E3-7" /></td>
<td>E3-7</td>
<td>300µm</td>
<td>7</td>
<td>Perio</td>
<td>7410002/7400020</td>
</tr>
<tr>
<td><img src="image" alt="E3-9" /></td>
<td>E3-9</td>
<td>300µm</td>
<td>9</td>
<td>Perio</td>
<td>7400020</td>
</tr>
</tbody>
</table>

### PRE-INITIATED

<table>
<thead>
<tr>
<th>Name</th>
<th>Diameter</th>
<th>Length</th>
<th>Application</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>PI PRO 3-3</td>
<td>300µm</td>
<td>4mm</td>
<td>Surgical</td>
<td>7430001</td>
</tr>
<tr>
<td>PI PRO 4-4</td>
<td>400µm</td>
<td>4mm</td>
<td>Surgical</td>
<td>7430002</td>
</tr>
</tbody>
</table>

**NOTE:** All BIOLASE Tips for diode lasers are sold non-sterile and are for single-use only. See Section 10 in this User Manual for cleaning and sterilization instructions.
## Appendix C  Accessories/Spare Parts

<table>
<thead>
<tr>
<th>PART NO.</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>6400105</td>
<td>Surgical Handpiece</td>
</tr>
<tr>
<td>2400276/2400278</td>
<td>Laser Safety Glasses</td>
</tr>
<tr>
<td>6400629</td>
<td>Fiber Optic Cable</td>
</tr>
<tr>
<td>6400622</td>
<td>Disposable Screen Protector (20/Pack)</td>
</tr>
<tr>
<td>6400616</td>
<td>Remote Interlock Plug</td>
</tr>
<tr>
<td>2400043</td>
<td>Power Cord (US)</td>
</tr>
<tr>
<td>2400055</td>
<td>Power Cord (Int'l)</td>
</tr>
<tr>
<td>6201667</td>
<td>Wireless Footswitch</td>
</tr>
<tr>
<td>7400022</td>
<td>Whitening Handpiece</td>
</tr>
<tr>
<td>6400463</td>
<td>Battery Pack, Footswitch (2 x AAA)</td>
</tr>
<tr>
<td>6400180</td>
<td>Disposable Shields for Whitening Handpiece (30/Pack)</td>
</tr>
<tr>
<td>7400030</td>
<td>Laserwhite20 Whitening Gel Kit (5/Pack)</td>
</tr>
<tr>
<td>7400063</td>
<td>Laserwhite20 Whitening Gel Kit (5/Pack)-Canada Only</td>
</tr>
<tr>
<td>7400017</td>
<td>Surgical EZ Tip, E3-4mm (non-initiated), 30-Pack</td>
</tr>
<tr>
<td>7400016</td>
<td>Surgical EZ Tip, E4-4mm (non-initiated) 30-pack</td>
</tr>
<tr>
<td>7400020</td>
<td>Perio EZ Tip Combo Pack, E3-7mm, E3-9mm (non-initiated) 30-pack</td>
</tr>
<tr>
<td>7400019</td>
<td>Perio EZ Tip Combo Pack, E4-7mm, E4-9mm (non-initiated) 30-pack</td>
</tr>
<tr>
<td>7410002</td>
<td>Perio EZ Tip, E3-7mm (non-initiated) 30-pack</td>
</tr>
<tr>
<td>7410003</td>
<td>Perio EZ Tip, E4-7mm (non-initiated) 30-pack</td>
</tr>
<tr>
<td>7430001</td>
<td>Pi Pro Tip, Surgical, A3-4mm (pre-initiated) 20-pack</td>
</tr>
<tr>
<td>7430002</td>
<td>Pi Pro Tip, Surgical, A4-4mm (pre-initiated) 20-pack</td>
</tr>
<tr>
<td>5400602</td>
<td>Danger Warning Sign</td>
</tr>
</tbody>
</table>
Appendix D Electromagnetic Compatibility

APPENDIX D: ELECTROMAGNETIC COMPATIBILITY

CAUTION: 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 the following tables.

Portable and mobile Radio Frequency (RF) communications equipment can affect medical electrical equipment.

Accessories: Medical grade power cord, maximum length 3ft (1 meter), BIOLASE p/n 2400043 (US), 2400055 (Int'l).

Footswitch: Wireless, BIOLASE p/n 6201667.

WARNING: The use of accessories, other than those specified, except those supplied or sold by BIOLASE, Inc. as replacement parts for internal or external components, may result in increased EMISSIONS or decreased IMMUNITY of the Epic Pro diode laser system.

GUIDANCE AND MANUFACTURER’S DECLARATION – ELECTROMAGNETIC EMISSIONS

The Epic Pro diode is intended for use in the electromagnetic environment specified below. The customer or the user of the Epic Pro diode should assure it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>The Epic Pro diode laser uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RF emissions</td>
<td>Class A</td>
<td>The Epic Pro diode laser is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/</td>
<td>Class A</td>
<td></td>
</tr>
<tr>
<td>flicker emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Appendix D  Electromagnetic Compatibility

### GUIDANCE AND MANUFACTURER’S DECLARATION – ELECTROMAGNETIC IMMUNITY

The Epic Pro diode laser is intended for use in the electromagnetic environment specified below. The customer or the user of the Epic Pro diode should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Continuous level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>± 6 kV contact ± 8kV air</td>
<td>± 6 kV contact ± 8kV air</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, relative humidity should be at least 30%.</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>± 2 kV for power supply lines ± 1 kV for input/output lines</td>
<td>± 2 kV for power supply lines N/A</td>
<td>Main power quality should be that of a typical commercial or hospital environment. Input/output that does not apply because the Footswitch cable length is less than 3 meters.</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>± 1 kV differential mode ± 2kV common mode</td>
<td>± 1 kV differential mode ± 2kV common mode</td>
<td>Main power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines.</td>
<td>&lt;5% Ur (&gt;95% dip in UT) for 0.5 cycle 40% Ur (60% dip in UT) for 5 cycles 70% Ur (30% dip in Ur) for 25 cycles</td>
<td>&lt;5% Ur (&gt;95% dip in UT) for 0.5 cycle 40% Ur (60% dip in UT) for 5 cycles 70% Ur (30% dip in Ur) for 25 cycles</td>
<td>Main power quality should be that of a typical commercial or hospital environment. If the user of the Epic Pro diode requires continued operation during power main interruptions, it is recommended that the Epic Pro diode be powered from an uninterrupted power supply.</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power frequency (50-60 Hz) magnetic field</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**NOTE:** Ur is the A.C. mains voltage prior to applications of the test level.
## GUIDANCE AND MANUFACTURER’S DECLARATION – ELECTROMAGNETIC IMMUNITY

The Epic Pro is intended for use in the electromagnetic environment specified below. The customer or the user of the Epic Pro should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Continuous level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>3 Vrms</td>
<td>3 V</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Epic Pro diode laser, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>150 kHz to 80 GHz</td>
<td>3Vm</td>
<td>Recommended separation distance</td>
</tr>
<tr>
<td>Radiated RF</td>
<td>3V/m</td>
<td></td>
<td>d = 1.2√P</td>
</tr>
<tr>
<td>IEC 61000-4-3</td>
<td>80 MHz to 2.5 GHz</td>
<td></td>
<td>d = 1.2√P  80 MHz to 800 MHz</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>d = 2.3√P  800MHz to 2.5GHZ</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>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).</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
</tbody>
</table>

### 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.
**Appendix D** Electromagnetic Compatibility

**GUIDANCE AND MANUFACTURER’S DECLARATION – ELECTROMAGNETIC IMMUNITY**

The Epic Pro diode laser is intended for use in the electromagnetic environment specified below. The customer or the user of the Epic Pro diode should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Continuous level</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
</table>

A. Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephone 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 Epic Pro diode laser is used exceeds the applicable RF compliance level above, the Epic Pro diode laser should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Epic Pro diode laser.

B. Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [V1] V/m.

**RECOMMENDED SEPARATION DISTANCES BETWEEN PORTABLE AND MOBILE RF COMMUNICATIONS EQUIPMENT AND THE EPIC DIODE LASER**

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

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter W</th>
<th>Separation distance according to frequency of transmitter M</th>
</tr>
</thead>
<tbody>
<tr>
<td>150kHz to 80MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>d = 1.2√P</td>
<td>d = 1.2√P</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>23</td>
</tr>
</tbody>
</table>

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.
APPENDIX E: WIRELESS EQUIPMENT COMPLIANCE STATEMENT

This statement applies only to the wireless portion of the device:

This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

- Reorient or relocate the receiving antenna
- Increase the separation between the equipment and receiver
- Connect the equipment into an outlet on a circuit different from that to which the receiver is connected
- Consult the dealer or an experienced radio/TV technician for help

<table>
<thead>
<tr>
<th>FCC ID</th>
<th>OA3MRF24J40MA</th>
</tr>
</thead>
<tbody>
<tr>
<td>IC ID</td>
<td>7693A-24-24J40MA</td>
</tr>
<tr>
<td>Type of Radio Equipment</td>
<td>Low Power Device (2400-2483.5 MHz)</td>
</tr>
<tr>
<td>Equipment Type</td>
<td>Radio Equipment (RE)</td>
</tr>
<tr>
<td>Min Power</td>
<td>475 uW</td>
</tr>
<tr>
<td>Max Power</td>
<td>475 uW</td>
</tr>
<tr>
<td>Field Strength</td>
<td>92dBuV</td>
</tr>
</tbody>
</table>