Intra.Ox™ Handheld Tissue Oximeter
Instructions for Use

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ViOptix part number for this IFU: OXY-2-DUR-IFU-1
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1 General Information

1.1 Overview

The ViOptix Intra.Ox™ 2.0 Handheld Tissue Oximeter consists of two components: a reusable main unit that shows a digital readout of %StO2 when the system is in contact with tissue, and a disposable kit that contains a non-sterile single-use battery pack, sterile single-use disposable sheath, and QC Optical Target. The Intra.Ox™ 2.0 is abbreviated as Intra.Ox in this document. This sheath is placed over the reusable main unit with the battery pack attached.

The Intra.Ox™ non-invasively estimates the percent oxygen saturation (StO2) in a volume of tissue. The device uses spatially-resolved optical measurements at five wavelengths. The device performs measurements on the patient by direct physical contact to the patient's tissue and displays the StO2 estimate on the built-in screen. The ViOptix Intra.Ox™ Handheld Tissue Oximeter is constructed from biocompatible materials that can tolerate bodily fluids and other liquids such as disinfectants and marking materials.

This manual has been prepared to assist medical personnel in the operation of the ViOptix Intra.Ox™ Handheld Tissue Oximeter. Prior to operating this device, all personnel must read this manual and gain a thorough understanding of its proper operation. Special attention should be directed to all cautions and warnings regarding the use of the product.

ViOptix cannot, and does not intend within this manual, to give medical advice.

1.2 Indications For Use

The Intra.Ox™ 2.0 Handheld Tissue Oximeter is intended to non-invasively estimate the percent oxygen saturation (StO2) in a volume of tissue.

The Intra.Ox™ 2.0 Handheld Tissue Oximeter is indicated for use in monitoring patients during circulatory or perfusion examinations.

The Intra.Ox™ 2.0 Handheld Tissue Oximeter is intended to be used by physicians, surgeons, nurses, or other skilled users in a medical environment.

The Intra.Ox™ 2.0 Handheld Tissue Oximeter should only be used on adult patients.
1.3 Intended User
This device is intended to be used by physicians, surgeons, nurses, or other skilled users in a medical environment.

1.4 References

Trademarks

Intra.Ox™ Handheld Tissue Oximeter is a trademark of ViOptix, Inc.

References
References to ”ViOptix” in this manual shall imply ViOptix, Inc.

The information in this manual has been carefully checked and is believed to be accurate. In the interest of continued product development, ViOptix reserves the right to make changes and improvements to this manual and the products it describes at any time, without notice or obligation.

Caution: Federal law (US) restricts this device to sale by or on the order of a physician.

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6,516,209
9,398,870

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2  Safety

2.1  Contraindications, Warnings, and Cautions

2.1.1  Contraindications
There are no known contraindications for the use of the Intra.Ox™ Handheld Tissue Oximeter.

2.1.2  Warnings
Warnings alert the operator to potential serious outcomes to the patient or operator.

- Never make direct patient contact between the main unit and the patient. All measurements should be taken through the sheath to maintain sterility.
- Once inserted into the sheath, the reusable main unit should not be removed during the course of the procedure.
- Carefully follow all instructions for transfer of the reusable main unit into the sterile field to ensure that sterility is maintained.
- The Intra.Ox™ Handheld Tissue Oximeter comes packaged with a reusable main unit that shows a digital readout of %StO2 and a disposable kit that contains a battery pack and sterile single use disposable sheath, which is placed over the reusable main unit with the battery pack attached.
- Please note that the Intra.Ox™ Handheld Tissue Oximeter is NOT intended to be used in an MR environment
- Do not look directly at the light-emitting distal tip of the Intra.Ox™.
- Inspect the sensor before each use for visible damage. Do not use the instrument if the sensor has visible damage.
- To prevent damage, do not bend or apply torque to the sensor face.
- Hard knocks, particularly at the distal end of the device, may result in damage to the delicate fiber-optic cables, which could affect instrument performance. Do not use if there is visible damage to the Handheld Tissue Oximeter.
- Avoid extreme changes in temperature and/or humidity.
- To reduce the risk of electrical shock, do not open the equipment’s inner housing. Refer servicing to qualified personnel only.
- This device is not to be used in the presence of combustible or flammable gases, anesthetics, or cleaners/disinfectants.
- Dispose of used Intra.Ox™ Handheld Tissue Oximeter, disposable sheath, and battery pack using appropriate biohazard precautions.
- This equipment has been tested and found to comply with the limits of the standard for medical devices, IEC 60601-1-2 for Class A equipment. The limits are designed to provide reasonable protection against harmful interference in a typical medical installation. This equipment generates, uses and can radiate radio frequency energy,
and, if not installed and used in accordance with the manufacturer’s instructions, may cause harmful interference to other devices in the vicinity. Portable and mobile RF communications equipment can affect medical electrical equipment. There is no guarantee that interference will not occur in a particular installation. If this equipment causes interference with other devices, which may be determined by turning the equipment off and on, the user is encouraged to try and correct the interference by one or more of the following measures:

- Reorient or relocate the device receiving the interference
- Increase the separation between the equipment
- Consult the manufacturer for help

2.1.3 Precautions

Precautions alert the operator to conditions that could lead to tissue irritation or erroneous results.

- Clean tissue if colored disinfectants are used on or near the measurement location.
- Clean blood or other colored liquids off of tissue before measurements.
- Check the sensor application site frequently to assess positioning, circulation, and tissue sensitivity of the patient. If required, reposition the sensor to a new site or if redness or irritation is noted. If irritation continues, discontinue use.
- Avoid placement directly over bony prominences, scar tissue, dark birthmarks or other visibly non-homogenous tissue, as it could provide improper readings.
- If tissue is uneven, gently flatten tissue or move to a new location.
- When repositioning device, pick up sensor and replace; do not drag.
- Use extra caution when placing on thin or delicate tissue
2.1.4 Disposal

2.1.4.1 Sheath and Optical QC Target
The sheath and the optical QC target should be considered biohazardous waste after the procedure. They should be disposed of along with ordinary biohazard waste.

2.1.4.2 Re-usable Main Unit
The re-usable main unit contains a variety of electronics. It should be disposed of with electrically hazardous medical waste and in accordance with all local and hospital disposal procedures.

2.1.4.3 Battery Pack
The Intra.Ox™ Handheld Tissue Oximeter battery pack contains lithium batteries and should only be disposed of with electrically hazardous medical waste and in accordance with all local and hospital disposal procedures. Contact the hospital or local environmental control agency for additional instructions. **The battery pack must not be incinerated.**

2.2 Target Population
The Intra.Ox™ Handheld Tissue Oximeter should only be used on adult patients.
3 Installation and Setup

The Intra.Ox™ Handheld Tissue Oximeter is a prescription-only device. The reusable main unit comes fully assembled and packaged in a protective box along with storage bags and wipes for cleaning the sensor tip between uses. The disposable kit contains one disposable sheath in a sterile double-pouched configuration and one optical QC target in a sterile double-pouched configuration, along with a non-sterile battery pack. The Intra.Ox™ is handheld, battery powered, and requires no external power source.

Before using, inspect the device and packaging for any sign of damage. Do not use if unit has been compromised. No installation is required.

4 Device Configuration and Interface Elements

Figure 1: Device Configuration and Interface Elements

4.1 Display

When the battery pack is connected to the reusable main unit, the OLED displays a splash screen, followed by a series of instructions related to sheathing procedure.

When in use during surgery, the OLED display shows the following:

- Current tissue oxygen saturation value, in percent
- Current measurement quality 'Q' on 1-5 scale, represented by a series of ascending bars
- Total powered-up duration during the current surgery
- A battery life indicator
- The remaining lifetime number of uses for the main unit
4.2 Sensing Surface (Sensor Face)

The sensor face, which takes measurements through the sterile sheath, includes:

- Light sources
- Light detectors

5 Operating the ViOptix Intra.Ox™

5.1 Device Setup

The procedure for Intra.Ox™ setup requires two operators:

- A ‘sterile operator,’ inside the sterile field. This operator handles the disposable kit, including the sterile sheath.
- A ‘non-sterile operator,’ outside of the sterile field. This operator handles the non-sterile main unit.

The device setup procedure must be followed strictly to ensure sterility of the sterile field.

Figure 2: On overview of the device setup process
5.1.1 Step 1: Passing components into the sterile field

**IMPORTANT:** The outer pouches are non-sterile and should only be handled by the non-sterile operator. The inner pouches are sterile and should only be handled by the sterile operator.

1. The non-sterile operator obtains the double-bagged sheath and double bagged optical QC target from disposable kit. The outer pouches are non-sterile, and the inner pouches are sterile.
2. The outer pouches should be opened by the non-sterile operator, and the sterile contents of the inner pouches slid into the sterile field.

5.1.2 Step 2: Removal of Sterile Contents

3. The inner, sterile pouches should be opened by the sterile operator in the sterile field. The optical QC target and the sheath can be removed from each bag and set aside.
5.1.3 **Step 3: Battery Installation**

4. The non-sterile operator holds up the reusable main unit and attaches the battery pack to the back of the device. It will attach to the device with magnets.

5. As soon as the battery pack is connected, the OLED screen of the device will turn on. The non-sterile operator will see the Test Pattern, followed by the ViOptix logo.

![Figure 3: Left: The test pattern. Right: The ViOptix logo](image)

After a few seconds, the device will authenticate the battery and display the following screen:

![Figure 4: Battery and configuration authentication screens](image)

After battery authentication, the both operators will be prompted to work together install the reusable main unit into the single-use sheath. The device will display an animation of the process.

5.1.4 **Step 4: Tab and Foam Removal**

6. The sterile operator obtains the sheath, and removes the tab protecting the window by pulling upwards on the exposed plastic component.

7. The sterile operator removes the protective foam tab from the device latch by pulling downwards on the latch.
5.1.5 Step 5: Installation into Sheath

8. The non-sterile operator drops the combined reusable main unit and battery pack into the sterile sheath at the edge of the sterile field.

5.1.6 Step 6: Closing the Sheath

9. The sterile operator closes the lid of the sheath until it latches.

At this point, the non-sterile operator has completed their part of the installation process.

Once the sheath is latched closed, it should not be opened again for the duration of the procedure.

After installation, the system will verify the sheath installation process as it goes through the below screens.

The device warm-up may take up to ten minutes. Keep the device away from heat sources (such as bright lights) and from cold circulating air during the warm-up.

If the error ‘Missing Calibration’ appears on-screen during device warm-up, the device cannot be used. Please use a different reusable main unit for this procedure and contact ViOptix.
5.1.7 Optical Target Quality-Control Check

To verify the sheath installation process, the sterile operator uses the enclosed optical QC target.

10. First, the distal tip of the device is firmly pressed against the black hemispherical side of the optical QC target. This ensures good contact between the optics and the window.

![Image of optical QC target with tip against a surface]

11. Next, the device prompts the operator to press the device against the other side of the optical QC target.

![Image indicating to adjust position until Q level reaches max]

The sterile operator will hold the distal tip of the assembled device in firm, parallel contact against the plastic. After this check, the device transitions into measurement mode, and the Intra.Ox™ is ready for use.

5.2 Holding the Device

Hold the tip of the Intra.Ox™ device similarly to a pen, with the wrist in a neutral position. Lightly rest hand on the patient and use fingers to brace the device for maximum stability. Sensing face should be parallel to tissue and in gentle contact. Do not apply pressure to the sample site as blanching may occur, resulting in local ischemia and a misleading measurement.

Avoid non-parallel/angled contact with the tissue as doing so reduces device measurement accuracy.
5.3 Measuring Percent Oxygen Saturation

Ensure the tissue is clean and the distal tip of the sheath is dry. Place the sensor face on the patient in gentle contact with the tissue of interest. Continue to hold the device against the tissue for a few seconds until an oxygen saturation percentage appears on the display. The display is organized as depicted below:

**Figure 5: Normal Display Layout**

5.3.1 Powered Up Time Display

The total elapsed time that the device has been powered up is displayed in hours, minutes and seconds in the lower left of the display after the device has warmed up.

Battery life is expected to last 8 hours of operating time including time in standby mode

5.3.2 Remaining Device Uses

The fraction at the top of the display indicates the remaining number of procedures for which the re-usable durable device can be used. The re-usable durable device can be used for a total of 25 procedures.
5.3.3 Quality Measure

An approximate metric of measurement quality is displayed on the bottom of the screen above the powered-up time. The metric is on a 1-5 scale, 5 being the most desirable. A low number on this scale does not necessarily represent poor contact, as the Quality Measure incorporates both contact quality and tissue homogeneity.

The operator should seek to have both a number displayed for the Quality Measure and to have that number stay constant for several seconds while in gentle contact with the tissue. If at least a '2' cannot be achieved, try taking a measurement on an adjacent measurement site.

5.3.4 Percent Oxygen Saturation Display

The estimated oxygen saturation is indicated in percent in large text, centered on the OLED screen.

![Image of 62% oxygen saturation]

If no sample is detected by the Intra.Ox™ (ie, the device is being held in air), the digits for Percent Oxygen Saturation and Quality Measure are replaced by dashes:

![Image of dashes for oxygen saturation]

Figure 6: Invalid Measurement Display

Other warning and error messages are addressed in Section 5.

If the Intra.Ox™ is positioned on tissue, and no percent oxygen saturation value appears, and no warnings are indicated on the screen, gently wipe the sensor face with a soft cloth moistened with saline solution to remove any contaminants and dry thoroughly before use.
5.3.5 Low Battery Indicator Display

The approximate remaining battery life is provided on the top left of the device screen.

![Battery Indicator]

Figure 7: Low Battery Indicator

The Intra.Ox™ tissue oximeter can continue to be used as long as battery life remains. The battery is not rechargeable.

5.4 Other Device Modes

5.4.1 Standby Mode

If there are no valid measurements for 2 minutes, the Intra.Ox™ will automatically go into Standby mode. In this mode, the following screen will appear.

![Standby Mode]

The device will return to normal use mode when it is picked up.
5.4.2 Device Dropped Mode

The Intra.Ox™ contains an accelerometer that detects if the system is dropped. **If it is dropped on a non-sterile surface, the device cannot continue to be used.** However, the Intra.Ox may continue to be used if dropped onto a sterile surface, such as a table within the sterile field in the operating room.

In this mode, the device will display the following guidance:

![Device Dropped]

It will then ask the user to re-do the optical QC target checks outlined in Section 0. Assuming that the optics are still working properly after the drop, the device will return to its typical measurement mode.

5.4.3 Sheath Open Mode

If at any time during the procedure the sheath is opened, the device will display the following:

![Sheath Open]

The device should not be used for patient contact while the sheath is open. The device can be returned to measurement mode by re-closing the sheath.
5.5    After the procedure

5.5.1    Disassembly

Once the sterile field is broken, the Intra.Ox™ device may be removed from the sheath by pushing upwards on the metal latch on the front of the device. This will result in the 'Sheath Open Mode' warning appearing on-screen. The re-usable main unit and battery pack can be pulled out of the sheath.

To power off the device, pull apart the battery pack and the re-usable main unit. (see figure below)

The battery pack and sheath should be disposed-of in accordance with Section 2.1.4.3.

5.5.2    Preparing the re-usable main unit component for the next procedure

After removing the battery pack, the re-usable main unit component requires preparation to get ready for the next procedure. This requires a two-step process

5.5.2.1    Cleaning the sensor face

Immediately after the procedure, the sensor face at the distal tip of the device should be wiped down with a wet cloth or gauze pad to remove any traces of adhesive from the window. These may interfere with device performance if not removed from the device prior to the next device use.

5.5.2.2    Cleaning the bulk device

After the procedure, the re-usable main unit component should be cleaned and disinfected using CaviWipes™. First, use a CaviWipe™ to clean the device, removing any visible debris from the surfaces of the re-usable main unit component. Then, using additional CaviWipe™ towellettes disinfect all device surfaces by ensuring all surfaces are wet for at least three minutes. After the 3 minutes of disinfection time allow the device to dry for at least 10 minutes prior to use.
5.6 Resolving Warnings and Errors

Warning and error conditions may occur that require the display of an error message on the OLED screen.

5.6.1 Warning Messages

Warning messages appear to alert the user to device conditions. Warnings will display as a yellow on-screen popup like the below.

The following table lists the conditions resulting in a warning message, the LCD message that will be displayed on screen and the recommended user action that can be taken following the display of one of these warnings.

<table>
<thead>
<tr>
<th>OLED Message</th>
<th>Recommended User Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>OVERHEATED</td>
<td>Move the device to a cooler area until condition no longer exists</td>
</tr>
<tr>
<td>Place down to cool</td>
<td></td>
</tr>
<tr>
<td>Too much light</td>
<td>Correct sensor face placement to improve tissue contact or shield the sensor face from bright light</td>
</tr>
<tr>
<td>Reposition Sensor or Reduce Ambient Light</td>
<td></td>
</tr>
<tr>
<td>Low Device Temp</td>
<td>Move the device to a warmer area until condition no longer exists</td>
</tr>
<tr>
<td>Hold device to warm</td>
<td></td>
</tr>
<tr>
<td>High Device Temp</td>
<td>Move the device to a cooler area until condition no longer exists</td>
</tr>
<tr>
<td>Allow device to cool</td>
<td></td>
</tr>
<tr>
<td>Sensor/Tissue Interference</td>
<td>Wipe to clean the sensor face or correct sensor face placement so it is flat against the tissue</td>
</tr>
<tr>
<td>Wipe sensor face or reposition</td>
<td></td>
</tr>
</tbody>
</table>
5.6.2 Critical Error Messages

Error messages represent situations that prevent the continued use of a device during the procedure. Critical errors will display as a red on-screen popup like the below.

![Critical Error Message]

The possible critical Errors that may appear are as follows:

<table>
<thead>
<tr>
<th>OLED Message</th>
<th>Recommended User Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hardware Failure</td>
<td></td>
</tr>
<tr>
<td>Replace Durable</td>
<td>The device is inoperable.</td>
</tr>
<tr>
<td>Battery Depleted</td>
<td></td>
</tr>
<tr>
<td>Replace Battery</td>
<td>The main unit must be paired with a new battery back and sheath to continue measurements</td>
</tr>
<tr>
<td>Sheath Opened</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The sheath was opened before the device was in measurement mode inside the sterile field.</td>
</tr>
<tr>
<td></td>
<td>The sheath and battery pack must be replaced</td>
</tr>
<tr>
<td>Sheath Removed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The sheath was removed entirely from the device. The sheath and battery pack must be replaced</td>
</tr>
<tr>
<td>No remaining uses</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The main unit has reached end of life, and cannot be used for further procedures. Please</td>
</tr>
<tr>
<td></td>
<td>contact ViOptix to obtain a new main unit</td>
</tr>
</tbody>
</table>
### 6 Specifications

<table>
<thead>
<tr>
<th>Component</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tissue Oxygen Saturation Range</strong></td>
<td>1 – 99%</td>
</tr>
<tr>
<td><strong>Wavelengths</strong></td>
<td>730, 760, 810, 845, 895 nm LED</td>
</tr>
<tr>
<td><strong>System Control</strong></td>
<td>System will perform self-test when power is turned on</td>
</tr>
<tr>
<td><strong>Alarms (visual)</strong></td>
<td>Display conditions such as</td>
</tr>
<tr>
<td></td>
<td>• Low battery</td>
</tr>
<tr>
<td></td>
<td>• Measurement errors</td>
</tr>
<tr>
<td><strong>Battery Life</strong></td>
<td>8 hours of continuous use</td>
</tr>
<tr>
<td><strong>Operating Mode</strong></td>
<td>Spot Measurement Mode</td>
</tr>
<tr>
<td><strong>Operating Conditions</strong></td>
<td>• Device shall operate normally at ambient temperatures of 18°C to 24°C</td>
</tr>
<tr>
<td></td>
<td>• Device shall operate normally in a humidity environment of 20% to 80% (non-condensing)</td>
</tr>
<tr>
<td><strong>Transport and Storage</strong></td>
<td>• Device (packaged) shall operate normally after storage at -18°C to 60°C</td>
</tr>
<tr>
<td></td>
<td>• Device (packaged) shall operate normally after storage at 10% to 85% (non-condensing).</td>
</tr>
<tr>
<td><strong>Dimensions</strong></td>
<td>3” x 3” x 7”</td>
</tr>
<tr>
<td><strong>Weight</strong></td>
<td>Less than 1 lb</td>
</tr>
<tr>
<td><strong>Sensor Specifications</strong></td>
<td>IEC 60601-1. Any surface of the system that comes into contact with a patient for a time of 1 minute shall not exceed 41°C. The device complies with ANSI/IESNA RP-27.1-15 (Recommended Practice for Photobiological Safety for Lamps and Lamp Systems – General Requirements) for light output.</td>
</tr>
<tr>
<td><strong>IPX Rating</strong></td>
<td>• Main unit Only (No Battery Pack): IPX2</td>
</tr>
<tr>
<td></td>
<td>• Sealed Sheath (with Main unit and battery pack installed): IPX2</td>
</tr>
<tr>
<td></td>
<td>• Battery Pack: IPX0</td>
</tr>
</tbody>
</table>
7 Labels

7.1 Box and Pouch label (Sterile Sheath)
7.2 Box label (Re-Usable Main unit)
7.3 Box and label (. Battery)

Contents: (1) Disposable Intra.Ox™ Battery Pack

ViOptix P/N: OXY-2-BAT-1

1 Consult instructions for use

LOT # 34105

2 Do not re-use

ViOptix, Inc.
39655 Eureka Drive
Newark, CA 94560 USA
www.vioptix.com

Rx only CAUTION: Federal law (US) restricts this device for sale by or on the order of a physician.

Manufactured in the USA for ViOptix, Inc.
39655 Eureka Drive, Newark, CA 94560 USA

U.S. Patents: www.vioptix.com/patents
Label P/N: 89712 Rev B

Part number: OXY-2-DUR-IFU-1 Rev A
7.4 Box and Pouch label (Sterile Optical QC Target)

7.5 Device label

(Placed on the back of the re-usable main unit)

(Placed on the front of the re-usable main unit)
### 7.6 Glossary of Symbols

<table>
<thead>
<tr>
<th>Symbol/Phrase</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Symbol" /></td>
<td>Do not re-use</td>
</tr>
<tr>
<td><img src="image2" alt="Symbol" /></td>
<td>Sterile Disposable Intra.Ox™ Sheath is intended for single-patient one-time use. DO NOT REUSE, DO NOT RE-STERILIZE</td>
</tr>
<tr>
<td><img src="image3" alt="Symbol" /></td>
<td>Sterilized by Ethylene Oxide</td>
</tr>
<tr>
<td><img src="image4" alt="Symbol" /></td>
<td>Use by expiration date stamped</td>
</tr>
<tr>
<td><img src="image5" alt="Symbol" /></td>
<td>Attention, consult accompanying documents</td>
</tr>
<tr>
<td>ViOptix P/N:</td>
<td>Model Number (Catalog Number)</td>
</tr>
<tr>
<td><img src="image6" alt="Symbol" /></td>
<td>Serial Number</td>
</tr>
<tr>
<td><img src="image7" alt="Symbol" /></td>
<td>Manufactured By</td>
</tr>
<tr>
<td><img src="image8" alt="Symbol" /></td>
<td>Applied part type BF</td>
</tr>
<tr>
<td><img src="image9" alt="Symbol" /></td>
<td>Do not use this equipment in the MRI scan room</td>
</tr>
</tbody>
</table>