





Titan X300

(US Patent 9,772,094)

Xenon Illuminator with Manual Dimming Control **Operator Manual**

CE



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1. INTRODUCTION

Congratulations on the purchase of your new Titan X300 Xenon Illuminator!

This user-friendly Xenon Illuminator is a high efficiency light source utilizing state-of-the-art illumination technology. It offers a variety of features such as:

- 5600 K daylight brightness for perfect color definition
- Quiet operation
- Compact and light weight
- Turret which adapts to various types of light guides
- Mechanical Iris
- Easy lamp replacement
- Lamp life display indication

In short, you have chosen the best and we would like to make sure you receive the optimal results with your new illuminator by using it correctly.

This Operator Manual will help you to install the device and optimally integrate it with other components of your system. It will also instruct you how to operate the illuminator and how to keep it clean. It will give you maintenance and service guidelines as well as recommendations for best performance results.

1.1 Indications for Use

The intended use of this device is to provide light for fiberoptic cables and instruments – providing light for instrumentation via fiberoptic cables for use in surgical fields.

The xenon illuminator is intended to be used in a controlled operating room environment by qualified medical personnel. The illuminator is provided non-sterile and is not intended to be sterilized. The system has an expected, but not limited, three-year service life.

The illuminator is not intended to be used for monitoring, diagnostic, or other life support functions. This device does not sustain nor support life. The device is not intended to compensate for injury, handicap, replacement or modification of anatomy, or control of conception. No special intervention is necessary in the event of device failure. As such, this device has no essential performance as defined by IEC 60601-1, and should this device fail to operate, a suitable backup should be available for any procedure where it may be used.

There are no contraindications.

1.2 Functions of Design

The xenon illuminator comprises a standalone, mains-powered illuminator that produces 235klux (minimum) with a standard headlight at 16-inch (40cm) distance in the visible spectrum range. The illuminator is typically used with a fiberoptic light guide that connects to a light port located on the front panel. Light intensity is controlled via dimming knob located on the front panel of the console.

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2. WARNINGS AND CAUTIONS

Use of this equipment may present hazards to the user and/or patient. Before operating this device, please read this operating manual thoroughly and follow all warnings, cautions, and instructions for use. The words warning, caution, and note carry special meaning and should be carefully reviewed:



WARNING: Indicates risks to the safety of the patient or user. Failure to follow warnings may result in injury to the patient or user.



CAUTION: Indicates risks of improper use and/or damage to the equipment. Failure to follow cautions may result in loss of function or product damage.



NOTE: Indicates special information to clarify instructions or present additional useful information.

The appropriate "WARNING", "CAUTION" or "NOTE" symbol in this manual is intended to alert the user to the presence of important operating and maintenance instructions in the manual.



2.1 Warnings

- Federal law restricts this device to sale by or on the on the order of a licensed practitioner.
- The illuminator produces highly concentrated light. Avoid shining light beam into eyes or looking directly into the light beams at the ends of connected instruments and/or light guides. When not using the device, it is advised to fully dim the illuminator.
- Qualified personnel must determine a safe working distance between the ends of connected instruments and/or light guides and the patient for each application. There is a risk of patient injury if a light guide or instrument connected to the light source comes to close to the patient.
- User is responsible for determining if interruption of light output will create an unacceptable risk. Having a backup illuminator is advised.
- Not suitable for use in presence of flammable anesthetic mixture with air or with oxygen or nitrous oxide.
- For endoscopic procedures: the illuminator should only be used with type BF endoscopic instruments which have been certified to IEC 60601-1 and IEC 60101-2-18.



This symbol indicates type BF equipment.

- All devices and/or instruments connecting to the illuminator must be classified as medical
 equipment. It is the user's responsibility to ensure that all equipment used with this device
 meets all applicable standards such as IEC 60601-1.
- To prevent fire and/or electric shock, do not open or expose the illuminator to liquids.
- The illuminator-side light guide connection may become hot during use. Allow adequate time to for end tip to cool before removal from illuminator.
- Instruments and/or light guides connected to the illuminator must be NON-CONDUCTIVE.
 There should be no conductive shielding or any conductive connection between the illuminator and the patient. Such connections present a risk to patient safety.
- Instruments and/or light guides should be clean and dry before being connected to the illuminator
- **DO NOT** modify the equipment without authorization from the manufacturer.
- The illuminator is provided non-sterile and is not intended to be sterilized.
- Use only the power cord supplied with illuminator or medically approved power cords with less than 200mΩ of ground impedance and less than 16ft (<5m) of length. If unauthorized

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- cables are used, the device may have increased electromagnetic emissions and/or decreased electromagnetic immunity which may result in improper operation.
- This device meets CISPR 11 Class A limits and is suitable for use in a hospital and industrial environments. If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radiofrequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.
- Performance of this device may be affected in proximity of another device and/or equipment capable of producing high levels of RF emissions. The device should be used no closer than 12 inches (30 cm) to any part of RF equipment including cables. In the event performance of this device is affected due to high levels of RF emissions, relocation of the suspected device and/or equipment producing high levels of RF emissions, or the headlight system may reduce or eliminate the problem.



2.2 Cautions

- Before each procedure, carefully check the illuminator for damage. DO NOT use a damaged illuminator.
- The user should verify the light guide end tip and the active illumination port are of the same type before insertion. DO NOT attempt to force an end tip into an incorrect port.
- All servicing and repair must be performed by the manufacturer or qualified service technicians.
- Ensure that the air vents located on the illuminator are not obstructed to allow the device to receive the necessary cooling to prevent an overheating.
- Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

3. SPECIFICATIONS

PARAMETER	VALUE
Light Source Type	Ceramic 300W Xenon
Color Temperature	5600 K (typ.)
Lamp Life	1000 hours (typ.)
Lamp Replacement	Cartridge replacement
Brightness Control	Mechanical iris control
Light guide adaptor	Rotating turret with STORZ, ACMI, WOLF, and OLYMPUS
Input Power	100-240 VAC, 50/60 Hz
	450W (max.)
Circuit Protection	Resettable
Operating Conditions	68 to 104°F (20 to 40°C), 30 to 85% RH non-condensing, 700 to 1060 hPa
Storage Conditions	-4 to 140°F (-20 to 60°C), 0 to 95% RH non-condensing, 700 to 1060 hPa
Dimensions	13.3" x 6.1" x 18" (W x H x D)
	33.8 x 15.5 x 45.7 cm (W x H x D)
Weight	22 lbs / 10.0 kg

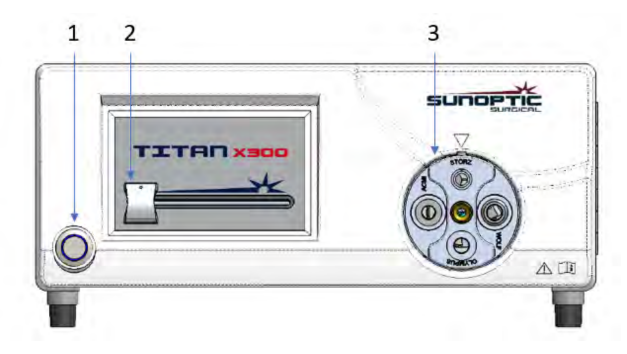
4. CLASSIFICATION

PARAMETER	VALUE
System Classification	FDA Class I, 510(k) exempt
	EU Class I, Active device per Annex IX, rule 1
Isolation	Type BF
EMC Certifications	CISPR 11 Class A, IEC 60601-1-2 4th Edition
	Electrostatic discharge: ±8 kV contact, ±15 kV air
	Radiated RF EM Fields: 3 V/m, 80 – 2700 MHz
	EFT / Burst: ±2 kV, ±1 kV signal lines, 100 kHz
	SURGE: ±0.5, ±1 kV
	Conducted disturbance: 3 V 150 kHz – 80 MHz and 6 V in ISM
	bands
	Power frequency magnetic fields: 30 A/m
	Voltage dips: 0% Un / 0.5 cycles at 0°, 45°, 90°, 135°, 180°,
	225°, 270°, 315°, 40 % Un / 5 cycles, 70 % Un / 25 cycles
	Voltage interruptions: 0% for 5000 ms
	Proximity fields: in accordance with EN 60601-1-2: 2015 table 9
CE Marking	Regulation (EU) 2017/745
Degree of protection against	IPX-0; no protection.
harmful ingress of water	
Degree of safety in the	Equipment is NOT suitable for use in the presence of flammable
presence of Flammable	anesthetics.
Anesthetics	
Mode of operation	Continuous

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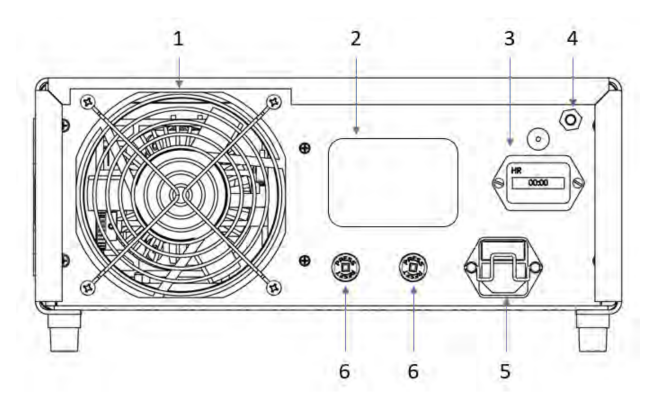
5. OVERVIEW

FRONT PANEL



No.	Name	Function
1	Lamp Switch	Turns illuminator ON and OFF. Illuminates blue when ON.
2	Intensity Control	Mechanically adjusts light intensity
3	Turret	Auto Shutter turret accepts end tip of fiber optic cable. Light is
		shuttered when no cable is present

REAR PANEL



No.	Name	Function
1	Fan	Cooling fan
2	Product Label	Device and Manufacturer information
3	Hour Meter	Allows tracking of total running time on unit and provides tracking of individual lamp running hours
4	Ground Stud	For potential equalization
5	AC Mains Inlet and Switch	Accepts AC Power cord and switches on AC power
6	Circuit Breaker	Over current protection

6. SETUP AND OPERATION

6.1 Device Setup



Place the Illuminator on a stable surface such as a cart, counter, stand, etc.

WARNING: DO NOT use the device in any environment with explosive or flammable gases.



WARNING: Avoid placing the device in an area where the illuminator may be splashed with liquids. A clear space minimum of 5 inches (12.7 cm) behind and above the cabinet is required. The illuminator should not be placed where its exhaust will influence other devices, nor where exhaust from other devices will influence the illuminator.



WARNING: DO NOT obstruct the exhaust or cooling vents of the illuminator. User must ensure that environment air temperatures surrounding the unit are within the allowable limits.

Make sure the power switch is in the OFF position.



Connect AC power cord to the appliance inlet located on the rear panel of the light source.

CAUTION: Use only power cords provided with the unit or cables approved for medical use.

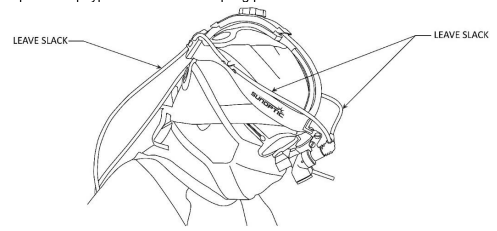


WARNING: To prevent electric shock, connect power cords of peripheral equipment through medical isolation transformers.



NOTE: When using a medical isolation transformer, ensure that the transformer has sufficient power ratings. Ensure that the power cord is connected to mains power with a three-prong plug.

Connect a fiberoptic light guide to the topmost position of the rotating turret, ensuring that the fiberoptic end tip type matches the accepting port.



6.2 Operation

After the power cable and light guide are properly connected, ensure the illuminator is in the lowest dimming position, and turn on the illuminator by pressing the mains power switch located on the front panel. The power indicator light within the switch should illuminate.

Ensuring that the light output from the attached light guide is directed to a safe area, adjust the intensity control slider until the desired brightness is achieved.



NOTE: The brightness control slider operates a progressive mechanical shutter. The lamp is running at full power irregardless of the intensity.

The hour meter located on the rear of the unit shows the total elapsed running time of the unit. When the lamp is replaced, reset the hour meter to track total lamp usage hours on the replacement lamp.

7. CLEANING AND DISINFECTION



WARNING: Ensure that the illuminator is de-energized and disconnected from mains power before attempting to clean and disinfect.

The illuminator can be wiped down with commercially available cleansers commonly used for disinfection of electronic equipment in hospitals such as ethyl or isopropyl alcohols, disinfecting sprays containing quaternary ammonium compounds, or hydrogen peroxide.



WARNING: DO NOT use strongly caustic or acidic cleansers such as "Clorox" hypochlorite bleach, ammonia, muriatic acid, or similar products. **DO NOT** use acetone, methyl ethyl ketone, or halogenated / chlorinated hydrocarbon solvents or cleansers containing any of these restricted compounds.

Apply cleaning agents by light spray or dampened towels. Do not pour liquids onto the device. Do not allow liquids to enter the device seams or ventilation openings.

Follow all applicable bloodborne pathogen procedures as required by OSHA and/or your hospital when cleaning and disinfecting the product.



WARNING: The illuminator is not sterilizable. DO NOT attempt to autoclave the device.

8. MAINTENANCE, SERVICING, REPAIR & WARRANTY

Performance of preventative maintenance is not essential. Defective items or equipment are to be serviced and repaired exclusively by persons authorized by the manufacturer. All repair work shall employ original manufacturer's parts only.



8.1 Lamp Replacement

WARNING: Ensure that the illuminator is unplugged from mains power and de-energized before attempting to replace the lamp.



WARNING: Ceramic Xenon lamps are at high internal pressure when cold and at operating temperature. Ceramic Xenon lamps may unexpectedly rupture resulting in discharge of hot fragments of quartz and/or glass and metal. Only handle lamps with protective covers in place.



WARNING: Do not change lamp while touching patient.



NOTE: This illuminator is designed for exclusive use with the Sunoptic Surgical P/N: SSX0300 lamp module. Failure to use this lamp could void warranty.

Turn unit off and unplug from mains power. Open lamp door located on side of illuminator.

Move the lamp positioning lever (found to the right of the lamp) clockwise, from horizontal to vertical, and pull the lamp cartridge out.

Grasp only the horizontal top and bottom grip flanges of the plastic housing of the lamp, pull straight out to remove the lamp. Insert a new one (lamp pointed to the left), pushing firmly to assure full engagement to the power connectors. Lower the lamp engagement lever counterclockwise, from vertical to horizontal, and observe that the lamp rises about 3mm as it moves to its proper operating position.

Log the serial number and hours on the hour meter. Close the lamp compartment door. Re-connect the power cord and turn the light source on according to 6.1. Reset hour meter to zero by depressing hour meter reset button on rear panel.

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8.2 Warranty

The illuminator carries a 3-year warranty from the date of shipment on workmanship and all defects of material.

Should your product prove to have such defects within three years of shipment, Sunoptic Technologies will repair or replace the product or component part without charge. Should your product(s) need servicing under this warranty, please contact Sunoptic Technologies or a local distributor for return authorization documentation.

Please carefully pack the unit in a sturdy carton and ship it to the factory. Please include a note describing the defects, your name, telephone number and a return address. Warranty does not cover equipment subject to misuse, accidental damage, normal wear, and tear or if transferred to a new owner without authorization from Sunoptic Technologies. This warranty gives you specific legal rights and you may also have other rights that vary from state to state.

8.3 Repair

You may return your product(s) for repair, shipping prepaid to the factory. Your product will be inspected, and an estimate of repair charges will be submitted to you for approval.

PHONE: +1 (877) 677-2832 INTERNATIONAL: +1 (904) 737-7611

FAX: +1 (904) 733-4832

8.4 Troubleshooting

Problem	Solutions
The power indicator (refer to section 5) is not lit.	Check that the AC power cord is properly connected
	Check the circuit breakers. If necessary, reset.
	Power input switch is in the off position
	Check lamp connection.
The power indicator is lit but	Check that the lamp door cover is secured.
the lamp will not ignite.	Check hour meter if lamp hours exceed the rated lamp life
	Replace the lamp (refer to 8.1).

9. END OF PRODUCT LIFE

In accordance with the European Waste from Electrical and Electronic Equipment (WEEE) directive, we encourage our customers to recycle this product whenever possible. Disposal of this unit must be performed in accordance with the applicable local environmental regulations.

In the US a list of recyclers in your area can be found at: http://www.eiae.org/.

Please contact customer service to issue a return authorization to return product to manufacturer at the end of product life.



10. SYMBOLOGY

	Manufacturer
	Date of manufacture (YYYY-MM-DD)
EC REP	Symbol for Authorized Representative in the European Community
\triangle	Caution, consult accompanying documents

\bigcap i	Consult Instructions for Use
CE	CE mark
©	Do not use if package is damaged.
X	Not for disposal in general waste
MD	Medical device
	Caution: Hot Surface
A	Caution: Dangerous Voltage
<u></u>	Storage / Shipping Humidity
6.6	Barometric Pressure
UDI	Unique Device Identifier
Ť	Keep Dry
\sim	AC Current
C US 220/02 Medical Endowent	Product Safety Mark
\Diamond	Equipotentiality
	Power on
0	Power off
*	Type BF
	Protective Earth (Ground)