

# IRRAflow® Laser Leveler (REF ICLS 010)

# **User Manual**

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IRRA*flow* Laser Leveler 7000494 Rev H, USA

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#### WARNING

Read all instructions and warnings prior to use.
Users of the IRRAflow Laser Leveler shall be trained medical personnel with training and experience in neurosurgical medical care.

Please note the components included in this package are meant to be used in conjunction with the IRRAflow Control Unit and IRRAflow instructions for use. The Laser Leveler is attached to the IRRAflow Control Unit for height positioning, activated after pushing on the switch and automatically turns off after 20 seconds.

IRRAS only accepts responsibility for the equipment's safety, usability and performance if:

- the equipment is used in accordance with its intended use, and
- the equipment is used in accordance with the product documentation.

#### 1. Scope

This manual describes the usage of the IRRAflow Laser Leveler model number ICLS 010. The IRRAS Laser Leveler is intended to be used with the IRRAflow CNS System. The Laser Leveler is used to target the tip of the IRRAflow catheter in use.

Users must read this manual carefully prior to using the IRRAflow Laser Leveler for the first time so that the features are thoroughly understood.



Warning - Failure to follow the instructions in this manual may endanger the patient and/or the operator!

# 2. IRRAflow Laser Leveler Accessory Device Description

The IRRAflow Laser Leveler is a reusable accessory to the IRRAflow CNS System and consists of a laser leveler as shown in **Figure 1 below**.

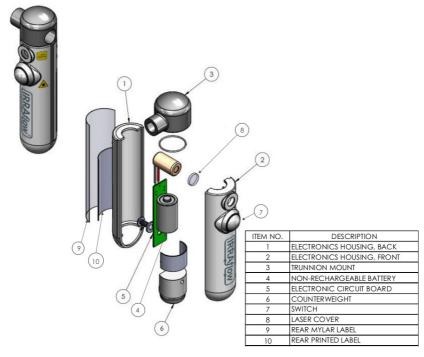


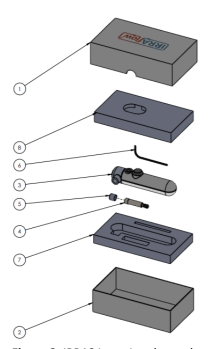
Figure 1. IRRAflow Laser Leveler.

The laser leveler secures to the side of the control unit with the laser beam aligned to the control unit's zero point. When the control unit is attached to the vertical IV pole and the laser leveler button is pressed, the laser leveler generates a horizontal laser beam that extends the zero point of the control unit and assists the user in setting the height of the control unit to the auditory meatus (theoretical tip of the catheter in use). Note: after pressing the button to operate the laser, the laser will automatically shut off after approximately 20 seconds.

The laser leveler is packaged with accessories and an IFU (Instructions for Use) in a product box as show in **Figure 2 below**. This product box includes:

- Laser Leveler
- Shoulder screw
- Push on cap
- Allen wrench
- Instructions for use

The device does not include software. It is non-sterile, can be re-used multiple times based on battery life, and has no patient contact.



ITEM NO.	DESCRIPTION
1	PACKAGING, LID, LL
2	PACKAGING, BASE, LL
3	ASSEMBLY, LASER LEVELER
4	screw, shoulder
5	CAP, PUSH ON, LASER LEVELER
6	WRENCH, ALLEN 4MM
7	insert, foam, bottom ll
8	INSERT, FOAM, TOP LL

Figure 2. IRRAS Laser Leveler packaged with accessories and IFU.

#### 2.1 Intended Use

The IRRAflow Laser Leveler is intended to be used with the IRRAflow CNS System. The Laser Leveler is used to adjust the height of the IRRAflow Control Unit such that the zero point of the Control Unit is at the same elevation as the patient's auditory meatus (or ear canal) prior to beginning treatment.

#### 3. Labels

#### 3.1 IRRAflow Laser Leveler Symbols

The IRRA <i>flow</i> Laser Leveler has been labeled with the following symbols:		
Symbol and text	Meaning	
	Warning	
[]i	Refer to the manual	
	Manufacturer	
EC REP	European Authorized Representitive	
~~ <u> </u>	Date of Manufacture	
IP22	Ingress Protection Level	
	Do not dispose of device in trash	
REF	Catalogue number	

The IRRA <i>flow</i> Laser Leveler has been labeled with the following symbols:		
Symbol and text	Meaning	
LOT	Lot number	
QTY	Packaged Quantity	
CAUTION LASER RADATION LASER RADATION And Essen to Essen Wavelength: 630 - 680 nm MAX Output: Crief Class 2 Laser Product Complies with 2158 4684 and 164.11 (EC 00027-1/2014	Laser Warning/Rating Label	
1	Temperature Range Limitations (storage and shipping identifier provided below symbol)	
<b>%</b>	Humidity Range Limitations (storage and shipping identifier provided below symbol)	
	Laser symbol	
Laser Aperture	Laser aperture location	
	Unique Device Identifier (UDI) 2D Barcode, unique to each device	
	Packaging made from recycled materials	
<u>11</u>	Correct upright position of a package	
Ţ	Device that can be broken or damaged if not handled carefully	
<b>®</b>	Do not use device if package is damaged	
R <sub>X</sub> Only	For Prescription Use Only	
R <sub>X</sub> Only C €	The Device Complies with Medical Device Directive 93/42/EEC	

## 4. Safety Regulations

#### General Safety Regulations and Warnings 4.1

NOTE: This device complies with 21 CFR 1040.10 and 1040.11 except for conformance with IEC 60825-1 Ed.3., as described in Laser Notice No. 56, dated May 8, 2019.



Device is not MRI compatible.



Do not attach other pieces (other than those specified) to the system.



Do not touch parts that could result in excessive leakage currents and the patient at the same time.



All specified cleaning and disinfection procedures must be performed.



Contraindication: Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.



Contraindication: Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could increased electromagnetic emissions result or decreased electromagnetic immunity of this equipment, result in improper operation and/or hazardous radiation exposure.



Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the Laser Leveler, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.



This is not a toy, keep away from children.



Do not stare into laser. Do not point laser into someone's eyes.



Device contains lithium metal (non-rechargeable) battery.



Device is not rechargeable and should not be recharged.



Do not open, crush, or heat device.



No modification of the equipment is allowed



There are no serviceable parts. No part of this medical accessory is to be serviced or maintained while in use



Corneal Exposure – Meets the Accessible Emissions Limits for Class 2 laser products. Never expose eyes to laser.



Skin Exposure – The laser diode output is below the Acceptable Exposure Limit (AEL) Class 3B and risk of burn when skin is exposed near the aperture to the active laser is unlikely. As with any laser, use appropriate precaution and avoid exposure near the aperature of the active laser diode.



Figure 3. Laser aperture shown.

### 5. Environmental and Handling Conditions

Laser Leveler Operation Temperature range	+10 to +40°C	
Operation	15 – 90%	
Air humidity	10 00/0	
Operation	70 – 106 kPa	
Ambient pressure	70 - 100 KPa	
Storage and transport	12 to 150 °C	
Temperature range	+2 to +50 °C	
Storage and transport	15 000/	
Air humidity	15 – 90%	
Storage and transport	50 40C l.D-	
Ambient pressure	50 – 106 kPa	
IP Rating	IP22	
Power Source	3.6V Battery	

Do not place the device near heating equipment nor expose to direct sunlight. Elevated temperatures can result in shortened battery life and degrade performance.

#### 6. Installation Procedure

- This device is not intended to be used as an independent device. This device is intended and designed to be used within the ICCU020 system.
- The IRRAflow Laser Leveler device is provided with a mounting screw for use with the The IRRAflow CNS control unit.
  - Contraindication: This device is not designed to be installed or used other than indicated.
- Clean and disinfect device per Section 9.
- Remove black cap from the head of the shoulder screw. Insert shoulder screw into the device as shown.



- Using the provided allen wrench, tighten the shoulder screw into the side of the IRRAflow Control Unit. The device should swing freely with no looseness along the shoulder screw. You should not be able to see screw thread.
- Replace the black cap over the head of the shoulder screw.



#### 7. Instructions for Use

- The recommended workflow begins with verifying the patient is in a resting position, the patient's eyes are closed and shut via tape and the laser leveler is attached to the control unit. The treatment normally occurs at the bedside but anywhere with direct line of sight to the patient's head may work.
- Rotate the body of the device towards desired target location (away from eyes).
- Push button to activate laser. A red dot shall appear on or near the patient.

Note: Do not hold down the button.

- Adjust the body of the device as need.
- With the device hanging freely, adjust height of the IRRAflow Control Unit as necessary.
- Once the ICCU020 is in the correct position, tighten the pole clamp feature to secure the ICCU020. Confirm the correct location via the laser leveler. Note: Do not hold onto the device during height positioning.
- Laser automatically turns off after approximately 20 seconds.

### 8. Disposal

Do not throw in trash. Recycle per your facility's electronic waste procedure and local regulations.

#### 9. Cleaning and Disinfection

The Laser Leveler shall be cleaned and disinfected after each treatment.

If fluids spill onto the Laser Leveler during treatment, pause or stop treatment and wipe off the spillage immediately.

The recommended method for cleaning is to wipe the parts with a soft cloth dampened with soapy water until all foreign material has been removed.

The recommended method for surface disinfectant is to wipe the parts with a soft cloth dampened with the following intermediate level disinfectant agents or an intermediate level disinfectant wipe.

Only the following intermediate level disinfectant liquid may be used with a minimum dwell time of one minute exposure:

• 70% Isopropyl alcohol or 70% ethyl alcohol solution



No liquid may be allowed to be directly poured or sprayed onto the Laser Leveler during cleaning or disinfecting as this may damage the equipment.



No liquid may be allowed to dispel from the cloth during cleaning, as this may damage the equipment.



Never use any type of tools or brush when cleaning, as this may damage the equipment.



Device is supplied non sterile. No components, parts or accessories to the IRRAflow Laser Leveler may be sterilized.

IRRAS AB does not make any claims for or representation as to the performance characteristics of this product if it is used in conjunction with other accessories or other than intended. If there are doubts about how to install or clean the Laser Leveler, the effect of cleaning, the functions and/or safety of the unit, then the unit should be withdrawn from service and the distributor and/or manufacturer consulted (see section 11 for contact information).

# 10. Appendix

#### 10.1 Laser Technical Specifications

ITEM	SPECIFICATION
Wavelength	630-680 nm
Beam Divergence (half-angle)	0.5 mRad
Pulse Pattern	Dot (5 mm)
Max Output	< 1 mW
Laser Product Classification	Class 2
Maximum Permissible Exposure, Eye [MPE (E)]*	6.36 J/m <sup>2</sup>
Eye Maximum Permissible, Exposure [MPE (H)]*	25.5 W/m <sup>2</sup>
Nominal Ocular Hazard Distance, Eye [NOHD]*	12m maximum
Assumed Exposure Duration	≤ 20 seconds

<sup>\*</sup>The maximum anticipated exposure duration, intra-beam viewing for Continuous Wave Lasers is 0.25 seconds. This assumes unintentional viewing and is based on natural reactions times. Reference IEC 60825-1 Ed. 3, §C.2.4 for Class 2 Lasers.

#### 10.2 Electromagnetic Compatibility

Electromagnetic compatibility tests have been performed with a 2.5 m power cable.

#### Guidance and manufacturer's declaration - electromagnetic emissions

The IRRAflow Control Unit, and IRRAflow Tube Set is intended for use in the electromagnetic environment specified below. The customer or the user of the IRRAflow CNS system should assure that it is used in such an environment.

Standard	Description	IEC 60601-1-2 4 <sup>th</sup> ed
Cispr 11/EN55011	RF Emissions	Group 1, Class B
IEC61000-4-2	ESD	8 kV contact, 15 kV air

IEC61000-4-3	Radiated RF Immunity	80-2700MHz :3 V/m ordinary, 10V/m home healthcare, 80% AM, 1 kHz 385MHz :27 V/m, PM,18Hz 450MHz :28 V/m, FM+/-5kHz dev, 1 kHz sine 710, 745, 780MHz :9 V/m, PM, 217 Hz 810, 870, 930MHz :28 V/m, PM, 18Hz 1720, 1845, 1970MHz :28 V/m, PM, 217 Hz 2450MHz :28 V/m, PM, 217 Hz 5240, 5500, 5785MHz :9 V/m, PM, 217 Hz
IEC61000-4-8	Magnetic Immunity	30 A/m, 50 Hz

Table B-1 Electromagnetic compatibility

#### 10.3 Laser Levelers and Manual

Please contact your local distributor if you wish to order Laser Levelers or a replacement manual.

- IRRAflow Laser Leveler (Ref no. ICLS 010)
- Laser Leveler User manual, English edition (catalogue no. 7000494)

#### 11 Contact

#### Manufacturer:



Address: USA

IRRAS USA, Inc. 10965 Via Frontera,

San Diego, CA 92127

**URL:** http://www.irras.com **Phone:** Tel: 1-800-946-0458

E-Mail address: US.customerservice@irras.com

#### **Re-Ordering Information:**

Region: USA

E-mail address: <u>US.customerservice@irras.com</u>

**Phone:** 1-800-213-4604

Region: Global

E-mail address: global.customerservice@irras.com

**Phone:** 31 20-210-1098

URL: <a href="http://www.irras.com">http://www.irras.com</a>

#### **EC Representative:**

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