

Instructions for Use

25ga I.D.D. Internal Delivery Device (VS0225.25)

Indications for use - Vortex Surgical 25ga I.D.D. Internal Delivery Device is intended to inject, irrigate, and/or remove fluids during vitreoretinal surgery. **Contraindications** - None

Device Description - Vortex Surgical 25ga I.D.D. Internal Delivery Device has a curved nitinol needle that can be retracted within 25ga stainless steel needle. The I.D.D. is compatible with its corresponding gauge trocar cannula system.

Known complications - Retinal Detachment, tears, holes, contusions, bleeding, inflammation, infection, and partial loss of vision.

These complications are statistically rare and it is assumed that the user is adequately trained in the treatment of these known complications and methods of avoidance.

Caution - Federal (USA) law restricts this device sale by, or on the order of, a physician.

Instructions for use: (Electronic Copy of IFU can be found at <https://www.vortexsurgical.com/instructionsforuse>)

1. Determine package integrity. 2. Affect sterile transfer of the product to the sterile field. 3. Remove the instrument from the tray. 4. Visually inspect tip surface. Do not use if damaged. 5. The instrument is now prepared for use. 6. Follow established surgical procedures.

Warnings - The device is designed to be utilized with cannulas. Ensure the curved portion of the device is fully retracted prior to insertion through the valved or non-valved cannulas. Do not use if package integrity or product compromised. Do not use if product exposed to conditions outside of indicated range. Do not use if EtO indicator does not show exposure to sterilization gas.


Device Storage - Vortex Surgical, Inc. recommends that the product is stored in a clean, dry and well-ventilated area at room temperature 15-37 °C (59-98.6 °F) away from direct sunlight.

Sterilization - This product is sterilized by Ethylene Oxide Gas (EtO) and provided with an EtO indicator. Verify indicator status color prior to use. **Re-sterilization** - Do not re-sterilize instrument. **Reuse** - Do not reuse or reprocess instrument. Reuse could lead to cross contamination, infection, or device failure.

Device Disposal - This single use surgical instrument should be considered clinical waste and should be disposed in accordance with clinical waste laws applicable in your country.

Reporting - Report any serious incident that has occurred in relation to the device to Vortex Surgical and your regulatory body including the competent authority of the Member State in which the user and/or patient is established.

Vortex Surgical, Inc. excludes all warranties, whether expressed, implied, or otherwise, to matters beyond the direct control of Vortex Surgical, Inc. and the end result of this device's use. This would include, but not be limited to handling, shipment and storage of the device and patient diagnosis and treatment. The fitness and merchantability of this device are as specified. Implied factors are specifically excluded. Vortex Surgical, Inc. is not liable for loss, whether incidental or consequential, damage and/or expense, arising wither directly or indirectly from the use of this device. There is no additional liability or responsibility assumed, other than that specified. All additional liability or responsibility is specifically disclaimed.

 **Vortex Surgical, Inc., 4 Research Park Drive, Suite 124, St. Charles, MO 63304, USA**
info@vortexsurgical.com 636-778-4350 www.vortexsurgical.com

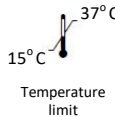
Manufacturer



Authorized representative
in the European Union



Sterilized using
ethylene oxide



Temperature
limit



Keep
dry



Keep away
from sunlight



Do not
resterilize



Do not
reuse



Do not used
if damaged



Prescription
use only



Medical
Device



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Issue Date: 2024-12-06

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Consult
Instructions
for Use



Lot
number



Catalog
number



Quantity



Use
By



Manufactured
in USA/Date of
Manufacture



Not made with
natural rubber
latex

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
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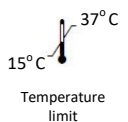
Manufacturer



Authorized representative
in the European Union



Sterilized using
ethylene oxide



Temperature
limit



Keep
dry



Keep away
from sunlight



Do not
resterilize



Do not
reuse



Do not used
if damaged



Prescription
use only



Medical
Device



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Issue Date: 2024-12-06

2200009 rev C



Consult
Instructions
for Use



Lot
number



Catalog
number



Quantity



Use
By



Manufactured
in USA/Date of
Manufacture



Not made with
natural rubber
latex