29G SPOTLIGHT
DIRECTIONAL CHANDELIER

Patent Pending

Directional function

“A bright vision on illumination”
29G Spotlight Directional Chandelier
A new generation chandelier with major improvements in functionality and ease of use.

Patent Pending

The Vitreo 29G Spotlight Directional Chandelier featuring high-end specifications with consistent performance for bimanual ophthalmic endo-illumination procedures.

“The Spotlight Directional Chandelier offers an easy and quick insertion in combination with an effective directional control of the light beam.”
Developed in collaboration with Professor Claus Eckardt

Benefits
• Improved illumination / visualization due to directional illumination system.
• Directional Chandelier facilitates bimanual surgery.
• Adapters provide compatibility to most endo-illumination systems.

Features
• Directional Chandelier system designed to enhance illumination control.
• Wide view fiber tip for global endo-illumination.
• Easy insertion and fixation of the fiber with 29G valved entry system.
• 29G minimal invasive incision.
• Integrated scleral marker
The Vitreq 29G Spotlight Directional Chandelier is compatible with most endo-illumination systems in combination with Light Source Adaptors:

- Constellation® Light Source Adaptors
- Stellaris® Light Source Adaptors
- Geuder® Light Source Adaptors
- Ruck® Light Source Adaptors
- DORC® Light Source Adaptors

Art.No.: LF29.D02
Packaged box/5, Sterile

**Light Source Adaptors**

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- Constellation® Light Source Adaptor
- Stellaris® Light Source Adaptor
- Geuder® Light Source Adaptor
- Ruck® Light Source Adaptor
- DORC® Light Source Adaptor

*All product names, trademarks and registered trademarks are property of their respective owners.

**Endo-Illumination Brief Statement for Vitreq Spotlight Directional Chandelier:**

**Indications for Use:** Vitreq Spotlight Chandelier is intended for endo-illumination during ophthalmic surgery.

**Precautions:** Do not use product if package integrity has been breached or compromised; do not use product after the expiration date.

**Caution (USA):** Federal law restricts this device to sale by or on order of a licensed physician.

**Attention:** See Instructions for Use included in product packaging for complete listing of indications and warnings.
ISO Certification
VitreQ’s quality system complies with the international quality and regulatory requirements and is certified by the Notified Body DQS Medizinprodukte GmbH, according to international standard ISO 13485 Medical devices – Quality management systems – Requirements for regulatory purposes.

CE Certification
The VitreQ medical devices are certified according to the medical device directive 93/42/EEC.

FDA
VitreQ is an FDA registered manufacturer of medical devices.

European Countries
To order VitreQ products in Europe contact a Customer Service Representative via e-mail, telephone or fax.

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