



# Certificate

The Certification Body of  
TÜV Rheinland Product Safety GmbH

hereby certifies that the organization

**Precision Optics Corporation**  
**22 East Broadway**  
**Gardner, MA 01440**  
**USA**

has established and applies a quality management system for medical devices  
for the following scope:

**Design/development, manufacture and service of optical  
endoscope and accessories; thin film coatings and filters  
for endoscopy; optical instruments.**  
**Additional facility: see attachment**

Proof has been furnished that the requirements specified in

**EN ISO 13485:2003 + AC:2007**

are fulfilled. The quality management system is subject to yearly surveillance.

Certificate Registration No.: SX 60020945 0001

An audit was performed. Report No.: 30793222 001

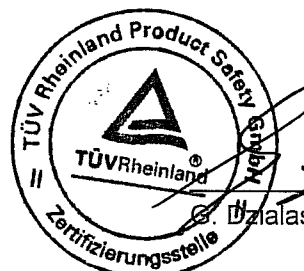
This Certificate is valid until: 04.03.2013

Certification Body



Akkreditiert durch  
Zentralstelle der Länder  
für Gesundheitsschutz  
bei Arzneimitteln  
und Medizinprodukten  
ZLG-ZQ-995.00.01-46

Cologne, 21.03.2008



**TÜV Rheinland Product Safety GmbH - Am Grauen Stein - D-51105 Köln**

Tel.: (+49/221) 806 - 1371 Fax: (+49/221) 806 - 3935 e-mail: cert-validity@de.tuv.com http://www.tuv.com/safety

# Certificate



TUV Rheinland of North America, Inc., a recognized  
CMDCAS Registrar, certifies that

**Precision Optics Corporation**  
22 East Broadway  
Gardner, MA  
USA


has established and maintained a  
**Quality Management System**  
**according to ISO 13485:2003**

Audit Report No.:	30793222.001
Certificate Registration No.:	74 500 3133
Expiry Date:	April 01, 2011

for the Design and Development, Manufacturing, and Service of  
**Optical endoscopes and accessories; thin film coatings and  
filters for endoscopy; optical instruments**

(See attachment for sites covered by this registration)



  
Gregor Dzialas - Certification Officer  
TUV Rheinland of North America, Inc.  
Newtown, Connecticut  
Effective Date: April 02, 2008



# Certificate

The Certification Body of  
TÜV Rheinland Product Safety GmbH

hereby certifies that the organization  
**Precision Optics Corporation**  
**22 East Broadway**  
**Gardner, MA 01440**  
**USA**

has established and applies a quality management system  
for the following scope:

**Design/development, manufacture and service of optical  
endoscope and accessories; thin film coatings and filters  
for endoscopy; optical instruments.  
Additional facility: see attachment**

Proof has been furnished that the requirements specified in

**EN ISO 9001:2000**

are fulfilled. The quality management system is subject to yearly surveillance.

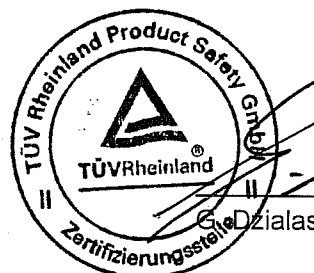
Certificate Registration No.: SY 60020946 0001

An audit was performed. Report No.: 30793222 001

This Certificate is valid until: 04.03.2013

Certification Body

Cologne, 21.03.2008



*[Handwritten signature]*  
G. Dzialas

**TÜV Rheinland Product Safety GmbH - Am Grauen Stein - D-51105 Köln**  
Tel.: (+49/221) 806 - 1371 Fax: (+49/221) 806 - 3935 e-mail: cert-validity@de.tuv.com <http://www.tuv.com/safety>



## APPROVAL

EC Directive 93/42/EEC Annex II, Article 3  
Full Quality Assurance System  
Medical Devices

Registration No.: HD 60020944 0001

Report No.: 30793222 001

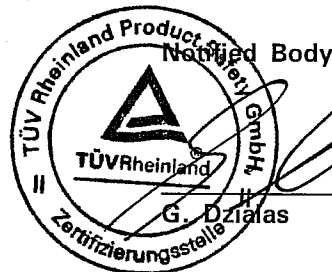
**Manufacturer:** Precision Optics Corporation  
22 East Broadway  
Gardner, MA 01440  
USA

**Scope:** Design/development, manufacture and service of optical  
endoscopes and accessories  
  
Additional facility: see attachment

**Date of Expiry:** 04.03.2013

The Notified Body hereby authorizes the quality management system established and applied by the company mentioned above. The requirements of Annex II, Article 3 of the directive have been met. This approval is subject to periodic surveillance, defined by Annex II, Article 5 of the aforementioned EC Directive, and can be used by the company with the manufacturer's declaration of conformity.

Cologne, 02.04.2008



**TÜV Rheinland Product Safety GmbH - Am Grauen Stein - D-51105 Köln**  
Accredited by Zentralstelle der Länder für Sicherheitstechnik (ZLS) and  
Zentralstelle der Länder für Gesundheitsschutz bei Arzneimitteln und Medizinprodukten (ZLG).

Notified under No. **0197** to the EC Commission.

CE The CE marking may be used if all relevant and effective EC Directives are complied with. CE