

EUROLINK (EUROPE) LTD

DECLARATION OF CONFORMITY

Manufacturer:

The Lebanon Corporation
1700 N. Lebanon Street
Lebanon
Indiana 46052
USA

Conforming Apparatus:

Intraocular Pressure Reducer

Classification:

Class I - according to Annex IX Rule 5

**Medicines and Healthcare products
Regulatory Agency Apparatus Description:**

Pressure Relief Device and Accessories

**Medicines and Healthcare products
Regulatory Agency Registration
Reference:**

CA 008803

Technical Document Reference No:

TF-0001

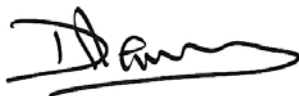
**Harmonised EMC Standard(s) and
Documents Referenced:**

BS EN ISO 14971:2001 Medical Devices –
Application of Risk Management to Medical Devices
Medical Devices Directive – Annex I, Essential
Requirements

EU Authorised Representative:

D.R.M. Green
Eurolink (Europe) Ltd.
Avalon House
Marcham Road
Abingdon
Oxon OX14 1UD
United Kingdom
Tel: (44) 1793 784545
Fax: (44) 1793 784551

We certify that the apparatus identified above conforms to the requirements of Council Directive 93/42/EEC, on the approximation of the laws of the member state relating to medical devices.



Signed:

D. R. M. Green

Date: 11 March 2009
