



## EC CERTIFICATE

Certificate No 1934/MDD

### Full Quality Assurance System Approval Certificate

On the basis of our examination carried out according to Annex II, excluding section 4, of the Directive 93/42/EEC and its revised version, we hereby certify that:

#### NOVAX PHARMA SAM

98000 MONACO - 20,Av. De Fontvieille,Le Coronado,7° Etg, 3 (MCO) - Monaco

manages in the factory of:

98000 MONACO - 20,Av. De Fontvieille,Le Coronado,7° Etg, 3 (MCO) - Monaco

a quality assurance system ensuring the conformity of the following products:

**Lubricant ophthalmic solutions with sodium hyaluronate**

**Sterile wipes for ophthalmic use**

**Ophthalmic solutions with sodium hyaluronate and sodium chloride 5%**

**Wetting and lubricating ophthalmic solutions**

**Lubricant ophthalmic solution with sodium hyaluronate spray**

**Eyelid foams**

series and type refs in the Annex

with the relevant essential requirements of the aforementioned directive (from design to final inspection and testing) and it is subject to surveillance as specified in section 5 of Annex II. For class III devices, this certificate is valid only with the relevant EC Design-Examination Certificate of Annex II.4.

Reference to IMQ files Nos:

DM16-0004410-01; DM17-0012269-01; DM17-0016717-01; DM17-0019462-01; DM18-0023521-01; DM18-0029039-01; DM18-0030362-01; DM19-0038813-01; DM19-0042511-01; DM19-0046044-01; DM19-0047010-01; DM20-0048282-01; DM20-0048500-01; DM20-0055276-01; DM20-0057459-01; DM21-0060709-01.

**This Approval Certificate is issued by IMQ S.p.A. as Notified Body for the Directive 93/42/EEC and its revised version. Notified Body notified to European Commission under number: 0051.**

Date: 2017-03-13  
 Updated: 2021-03-11  
 Substitution Date: 2020-11-02  
 Expiry Date: 2024-05-26

IMQ



## EC CERTIFICATE

Certificate No 1869/MDD

### Full Quality Assurance System Approval Certificate

On the basis of our examination carried out according to Annex II, excluding section 4, of the Directive 93/42/EEC and its revised version, we hereby certify that:

#### NOVAX PHARMA SAM

98000 MONACO - 20,Av. De Fontvieille,Le Coronado,7° Etg, 3 (MCO) - Monaco

manages in the factory of:

98000 MONACO - 20,Av. De Fontvieille,Le Coronado,7° Etg, 3 (MCO) - Monaco

a quality assurance system ensuring the conformity of the following products:

#### Contact lens care solutions

#### Wetting and lubricating ophthalmic solutions

Type ref. as to document "CODE PRODUCTS LIST Ref TF 01" rev. 04 dated 07/04/2020"; valid only if provided with IMQ stamp.  
Trade mark NOVAX PHARMA

with the relevant essential requirements of the aforementioned directive (from design to final inspection and testing) and it is subject to surveillance as specified in section 5 of Annex II. For class III devices, this certificate is valid only with the relevant EC Design-Examination Certificate of Annex II.4.

Reference to IMQ files Nos:

DM15A0483599-01; DM17-0015589-01; DM18-0026876; DM19-0038813-01; DM19-0042511-01; DM19-0046044-01; DM19-0047010-01.

**This Approval Certificate is issued by IMQ S.p.A. as Notified Body for the Directive 93/42/EEC and its revised version. Notified Body notified to European Commission under number: 0051.**

Date: 2016-04-19  
Updated: 2020-04-24  
Substitution Date: 2019-11-18  
Expiry Date: 2024-05-26

IMQ



www.imq.it

**CERTIFICATO N.  
CERTIFICATE N. 9124.NVXP**

SI CERTIFICA CHE IL SISTEMA DI GESTIONE PER LA QUALITA' DI  
WE HEREBY CERTIFY THAT THE QUALITY MANAGEMENT SYSTEM OPERATED BY

**NOVAX PHARMA SAM**

20 AVENUE DE FONTVIEILLE - MC 98000 MONACO

UNITA' OPERATIVE / OPERATIVE UNITS

20 AVENUE DE FONTVIEILLE - MC 98000 MONACO

E' CONFORME ALLA NORMA / IS IN COMPLIANCE WITH THE STANDARD

**ISO 13485:2016**

PER LE SEGUENTI ATTIVITA' / FOR THE FOLLOWING ACTIVITIES

Gestione della progettazione e della fabbricazione ed immissione  
in commercio di dispositivi oftalmici non attivi

*Management of the design and of manufacturing and placing on the market  
of non-active ophthalmic devices*

Ulteriori informazioni riguardanti l'applicabilità dei requisiti ISO 13485:2016 possono essere ottenute consultando l'organizzazione  
*Further clarifications regarding the applicability of ISO 13485:2016 requirements may be obtained by consulting the organization*

IL PRESENTE CERTIFICATO E' SOGGETTO AL RISPETTO DEL  
REGOLAMENTO PER LA CERTIFICAZIONE DEI SISTEMI DI GESTIONE  
*THE USE AND THE VALIDITY OF THE CERTIFICATE SHALL SATISFY THE  
REQUIREMENTS OF THE RULES FOR CERTIFICATION OF MANAGEMENT SYSTEMS*

<b>DATE:</b>	<b>PRIMA CERTIFICAZIONE</b> <i>FIRST CERTIFICATION</i> 2016-04-07	<b>EMISSIONE CORRENTE</b> <i>CURRENT ISSUE</i> 2022-01-19	<b>SCADENZA</b> <i>EXPIRY</i> 2025-04-07
--------------	---	---	--

IMQ S.p.A. - VIA QUINTILIANO, 43 - 20138 MILANO ITALY  
Management Systems Division - Flavio Ornago



*IQNet, the association of the world's first class certification bodies, is the largest provider of management System Certification in the world. IQNet is composed of more than 30 bodies and counts over 150 subsidiaries all over the globe.*



SGQ N° 005 A  
Membro degli Accordi di Mutuo Riconoscimento EA, IAF e ILAC  
*Signatory of EA, IAF and ILAC Mutual Recognition Agreements*

La validità del certificato è subordinata a sorveglianza annuale e riesame completo del Sistema di Gestione con periodicità triennale  
*The validity of the certificate is submitted to annual audit and a reassessment of the entire Management System within three years*



Organismo di Certificazione Federato CISQ  
[www.imq.it](http://www.imq.it)



[www.cisq.com](http://www.cisq.com)

CISQ è la Federazione Italiana di Organismi di Certificazione dei sistemi di gestione aziendale.  
*CISQ is the Italian Federation of management system Certification Bodies.*