



# CERTIFICATE

Quality Austria - Trainings, Zertifizierungs und Begutachtungs GmbH awards this **qualityaustria** certificate to the following organisation:

This **qualityaustria** certificate confirms the application and further development of an effective

**HIMEDIA**

**HiMedia Laboratories Pvt. Ltd.**  
Plot NO. C40, Road - 21Y, Wagle Industrial Estate,  
Thane (West) - 400604 Maharashtra, INDIA

**QUALITY MANAGEMENT SYSTEM**  
complying with the requirements of standard  
**ISO 9001:2015**

Design, Development & Testing of Microbiology, Animal Cell Culture, Plant Tissue Culture & Molecular Biology products

Registration No.: 27302/0  
Date of initial issue: 28 February 2022  
Valid until: 27 February 2025

The validity of the **qualityaustria** certificate will be maintained by annual surveillance audits and one renewal audit after three years.

Vienna, 28 February 2022

Quality Austria - Trainings, Zertifizierungs und Begutachtungs GmbH,  
AT-1010 Vienna, Zelinkagasse 10/3



Mag. Christoph Mondl  
General Manager



Mag. Dr. Werner Paar  
General Manager



Mag. Dr. Anni Koubeck  
Specialist representative



 **qualityaustria**



Quality Austria - Trainings, Zertifizierungs und Begutachtungs GmbH is accredited according to the Austrian Accreditation Act by the BMWFVW (Federal Ministry of Science, Research and Economy).

Quality Austria is accredited as an organisation for environmental verification by the BMLFUW (Federal Ministry of Agriculture, Forestry, Environment and Water Management).

Quality Austria is authorized by the VDA (Association of the Automotive Industry).

For accreditation registration details please refer to the applicable decisions or recognition documents.

Quality Austria is the Austrian member of IQNet (International Certification Network).

Dok. Nr. FO\_24\_028

Digitally signed by Iurcu Nicolae  
Date: 2019.11.14 12:02:41 EET  
Reason: MoldSign Signature  
Location: Moldova



## DECLARATION OF CONFORMITY

1. We declare that our products, specified under following categories as listed below, comply to and are manufactured according to the requirements of the International Standards as specified in the In Vitro Diagnostic Medical Devices Directive 98/79/EC.

- Dehydrated Culture Media and Supplements
- Culture Media Bases
- Antimicrobial Susceptibility Systems
- Bacteriological Differential Aids
- Cell Culture Media
- Lymphocyte and Granulocyte Separation Media
- Latex Agglutination Test Kits
- Epidemiological Screening Kit

2. Company or its authorized representative :

Name : Mr. Federico Pontigia, (Company – Neomed S.R.L.)  
Address : Via G.DI Vittorio, 2-A, 20017 Mazzo Di Rho, MILANO,  
ITALY

Phone : 00-39-02-93900652/93902434

Fax : 00-39-02-93900968

Email : [neomed@neomed.it](mailto:neomed@neomed.it)

shall fulfill the obligations imposed by in vitro medical device directives as applicable.

3. Company undertakes to keep upto date a systematic procedure to review experience gained during post production phase and to implement appropriate means to apply any necessary corrective action taking account of the nature and risk in relation to the product.
4. Company undertakes to notify immediately any malfunction/ deterioration of the performance of the product to the appropriate authority and shall recall such products already placed in the market.

For HiMedia Laboratories Private Limited, India

CEO, Dr. G.M. Warke

Dated : 15<sup>th</sup> November, 2011

15<sup>th</sup> November, 2011**AGREEMENT & REPRESENTATION**

With this letter of agreement and representation, HiMedia Laboratories Private Limited, India, appoints an authorized representative in the EC to represent the company. The details of the authorized representative are given below:

**Name** : Mr. Federico Pontigia, (Company-Neomed S.R.L.)  
**Address** : Via G. DI Vittorio, 2-A, 20017 Mazzo Di Rho, MILANO, ITALY  
**Phone** : 00-39-02-93900652/93902434  
**Fax** : 00-39-02-93900968  
**Email** : [neomed@neomed.it](mailto:neomed@neomed.it)

HiMedia Laboratories Private Limited, India shall, through this authorized representative, fulfill all the obligations imposed by the Directive 98/79/EC of the European parliament and of the Council of the European Union on *In Vitro* Diagnostic Medical devices and ensure that the products of the company meet all provisions of the directives as applicable from time to time.


HiMedia Laboratories Private Limited, India will be responsible for all legal and insurance matters pertaining to our products.

For HiMedia Laboratories Private Limited, India



CEO, Dr. G.M. Warke

I agree to represent HiMedia Laboratories Private Limited, India and be appointed as their EU Representative as per the tenets above.



EU Representative  
Name: Mr. Federico Pontigia



**NEOMED**

s.r.l.

Via G. DI VITTORIO,  
20017 MAZZO DI RHO (MI)  
Tel. 02/939.00.652-939.01.463-939.02.434  
Fax 02/939.00.968  
C.F./P. I. 09580650159  
C.C.I.A.A. 1304819 - TRIB. MILANO 291273