MEXPO INTERNATIONAL INC.

2828 Faber Street Union City, CA 94587-1204, USA www.blossom-disposables.com

EU Declaration of Conformity- Face Mask

PRODUCT DESCRIPTION

1. Product Name: Face Mask (non-woven products)

 Product Classification: Class I under Medical Device Regulation (EU) 2017/745 Annex VIII Rule 1

ADDRESS:

Mexpo International Inc.

2828 Faber Street

Union City, CA 94587-1204, USA

Tel: +1 (510) 489-6800; Fax: +1 (510) 489-3111

E-mail: blossomglo@aol.com; tim@mexpo-glove.com

Single Registration Number (SRN):

US-MF-000032548

Brand Owner

white-med GmbH Marburger Straße 251 35396 Gießen, Germany

AUTHORIZED REPRESENTATIVE:

EC Representative:

Blossom Europe, S.L. C/ Zurbano 45, 1st floor,

28010 Madrid, Spain

CH Representative:

CMC Medical Devices GmbH

Rigistrasse 3 6300 Zug,

Switzerland

Single Registration Number (SRN):

ES-AR-000019689

We, **Mexpo International, Inc.** as the legal manufacturer declare under our sole responsibility that the medical devices listed below conform to the requirement of the Medical Device Regulation (EU) 2017/745.

Face Mask Product:

1) N770 Peppler Care 3 PLY Latex Free Earloop Face Mask (Type II R) COLOR: ITEM #: UDI #:

Blue N770BG 00723860011886
Green N770GG 00723860011893
Lila N770LG 00723860011916
Pink N770PG 00723860011923
White N770WG 00723860011930

2) N770 Peppler Care 3 PLY Latex Free Tie-On Face Mask (Type II R)

COLOR: ITEM #: UDI#: N770BB Blue 00723860015860 Green N770GB 00723860015877 Lila N770LB 00723860015884 N770PB Pink 00723860015891 White N770WB 00723860015907

Basic UDI-DI: 0723860351778E

INTENDED USE: When properly worn, the surgical face masks are intended to protect both patient and healthcare workers from transfer of microorganisms, body fluids and airborne particles. This device is non sterile and for single use only.

MEXPO INTERNATIONAL INC.

2828 Faber Street Union City, CA 94587-1204, USA www.blossom-disposables.com

It is declared that above devices meet the requirement of the Medical Device Regulation (EU) 2017/745.

PPE Regulation (EU) 2016/425- Cat III and, where such is the case, with the following standards. DIN EN 14683, ASTM F2299/F2299M-03(2017), ASTM F2101, ASTM F2100, ASTM F1862/F1862M-17, ASTM F1494-14.

In accordance with Annex VIII, Medical Device Regulation (EU) 2017/745, listed above are non-invasive transient devices and are Class I devices under Rule 1 as Rules 2, 3, and 4 do not apply.

The following person is responsible for signing of this document:

Name and Address: Mexpo International Inc., 2828 Faber Street, Union City, California 94587-1204, USA

Authorized Signature:

Date: March 5, 2024

Name of responsible Person: Tim Thai

Position: President