

DECLARATION OF CONFORMITY

ACCORDING TO (EU) 2017/745 MEDICAL DEVICE REGULATION

EU Representative

SUNGO Europe B.V.

Olympisch Stadion 24, 1076DE

Amsterdam, Netherlands SRN: NL-AR-000000247

Conformity Assessment

Conformity Assessment Procedure

Annex II+III of Regulation (EU) 2017/745

Applicable Standards

EN ISO 14971: 2012

EN ISO 15223-1: 2016

EN 1041:2008+A1:2013

ISO 10993-1: 2018

EN ISO 10993-5: 2009

EN ISO 10993-10: 2013

EN 14683:2019+AC:2019 Type IIR

Remark

The declaration of conformity is valid in connection with the release technical document CE/MDR-AMS-01.

All the supporting documentation is retained at the premises of the manufacturer.

The Declaration of Conformity is exclusively under the sole responsibility of the manufacturer.

On behalf of SUNGO Europe office, I confirmed we are EU REP of the company who issue this document.

Manufacturer

Name: Ammex-Weida (Hubei) Health and Safety

Products Co., Ltd.

Address: Southern Industrial Zone (Xinlirenkou),

Xiantao, Hubei, China.

Product Information

Name: Medical Face Mask

Model: A-31, A-05B

GMDN: 35177

Basic UDI-DI: /

Classification: Class |

Declaration

We herewith declare that the above-mentioned products meet the requirements of Medical Device Regulation (EU) 2017/745 and the applicable standards above.

2021.2,25

Place: Hube

Sungo

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