

# **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: 2019

EN ISO 15223-1: 2016

EN 1041:2008+A1:2013

ISO 10993-1: 2018

EN ISO 10993-5: 2009

EN ISO 10993-10: 2013

#### Remark

The declaration of conformity is valid in connection with the release technical document CE/MDR-TNQ-02.

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.

## Manufacturer

Name: TNQ INDUSTRIAL CO., LTD

**Address:** 301, No.20, Aimin Road, NanCun Town, Panyu District, Guangzhou City, Guangdong P.R.,

China

SRN NO.: CN-MF-000008950

### **Product Information**

Name: TOURNIQUET

Model: TNQ-T0A00, TNQ-T0B00, TNQ-T0C00,

TNQ-T0D00, TNQ-T0K00, TNQ-YP00, TNQ-T0E00, TNQ-T0E01, TNQ-T0E02, TNQ-T0E03, TNQ-T0F00, TNQ-TDT00,

TNQ-TDT01, TNQ-TDT02, TNQ-TDT03

**GMDN**: 58128

Basic UDI-DI: 697463766TNQ-T0A001MG

Classification: Class I, According to Rule 1, Annex

VIII, Regulation (EU) 2017/745

### Declaration

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

Signature:

Position: GM

Date: 2021/7/6

Place Guanddogg/China