

EU Declaration of Conformity

According to Art. 19 of Regulation (EU) 2017/745 on Medical Devices

Name: Shanghai Greejoy Industry Co., Ltd.
Manufacturer: Add: 6 Building, Songhuang Road 999, Qingpu District, 201706. Shanghai, China
SRN: CN-MF-000035370
European Name: Prolinx GmbH
Representative: Add: Brehmstr. 56, 40239, Duesseldorf , Germany
SRN: DE-AR-000005129
Product Name: Dental Suction Unit
Trade Name: /
Basic UDI-DI : 697654080GS44
Product Models: GS-01 GS-02 GS-03 GS-03F GS-05 GS-10
GS-20 GS-30 GS-50 GS-M300 GS-M400 GS-E1000
GS-DC100
Classification acc. to MDR Ax. VIII: Class I, Rule13
Conformity assessment procedure: Regulation (EU) 2017/745 Annex XI Part A
Applied Common EN ISO 13485:2016 EN ISO 15223-1:2021 EN ISO 14971:2019/A11:2021
Specification/ standard: EN ISO 20417:2021 IEC 60601-1:2005 +A1:2012+A2:2020
IEC 60601-1-2:2014+A1:2020 EN 62366-1: 2015+A1:2020
IEC60601-1-6: 2010+A1: 2013+A2: 2020

We, the manufacturer, herewith declare under our sole responsibility that the above-mentioned products meet the provisions of the Regulation (EU) 2017/745 on Medical Devices (MDR), any other relevant Union legislation that provides for the issuing of an EU declaration of conformity, any CS used and in relation to which conformity is declared and other relevant regulations. All supporting documentations are retained under the premises of the manufacturer.

Place of Issue: Shanghai, China
Date of Issue: July 25, 2024
Signature:

Name: Mr.Mingbao Lu
Position: General Manager

