


Declaration of Conformity

Name and address of the manufacturer: **FEELLIFE HEALTH INC.**
Room 1903, Building A, No.9 Furong Road, Tantou Community,
Songgang Subdistrict, Bao'an District, Shenzhen, 518104,
Guangdong, China

Name and address of the authorized European Representative  Lotus NL B.V.
Koningin Julianaplein 10, 1e Verd,
2595AA, The Hague, Netherlands.

We declare under our sole responsibility that

the medical device: **Ultrasonic Nebulizers (Portable Mesh Nebulizer)**
mode: **Air 360+, mini Air 360+, A5, Air Pro, Air Angel, Air Force,
Air Mask, AeroCentre, Air Q+, Air Bee, Air Garden, Air Pro II,
Air Pro III, Aerogo, Air Plus, AeroCentre+, Air Kids, AirICU,
Air Pro VIII, Air Pro IX, Air Mask II, Air Plus2**
of class: **Ila, Rule 11**
according to annex IX of directive 93/42/EEC

meets the provisions of the directive 93/42/EEC and its amendments and its transpositions in national laws which apply to it.

Conformity assessment procedure: **Directive 93/42/EEC Annex II, excluding Section 4**

Registration No.: **HD 60147222 0001**
Certificate No.: 17062851 010
Issue date: 2020-07-30
Expiry date: 2024-05-26

Notified Body: **TÜV Rheinland LGA Products GmbH**
Tillystraße 2,90431 Nürnberg, Deutschland, CE 0197

This Declaration of Conformity covers all medical devices as specified in the product list belonging to this declaration and is only valid in connection with a batch specific Certificate of Compliance for all products concerned bearing the CE mark.

CE 0197

The above mentioned declaration of conformity is exclusively under the responsibility of

Company: **FEELLIFE HEALTH INC.**

Address: Room 1903, Building A, No.9 Furong Road, Tantou Community, Songgang Subdistrict, Bao'an District, Shenzhen, 518104, Guangdong, China

Shenzhen / 2020-12-17
Place, date

HUA JIAN / General Manager
Name and function 