DECLARATION OF CONFORMITY

TO COUNCIL DIRECTIVE 93/42/EEC: 2007/47/EC CONCERNING MEDICAL DEVICES

Manufacturer: Name:JiangSu YuYue Medical Equipment & Supply CO., LTD.

Address: No.1 Baisheng Road, Development Zone, 212300, Danyang, Jiangsu,

PEOPLE'S REPUBLIC OF CHINA

European Representative:

Name: Metrax GmbH

Address: Rheinwaldstr. 22, D-78628 Rottweil, Germany

VAT: DE 166 892 350

Product Name: Oxygen Concentrator Model: 8F-5AW, 8F-3A, 8F-5A

Classification (MDD, Annex IX): IIa (Rule 11)
Conformity Assessment Route: MDD Annex V.3

This Declaration is issued under the sole responsibility of the manufacturer, and the above mentioned products meet the transposition into national law, the provisions of Council Directive 93/42/EEC: 2007/47/EC concerning medical devices. All supporting documentations are retained under the premises of the manufacturer.

DIRECTIVES

General applicable directives:

Medical Device Directive: COUNCIL DIRECTIVE 93/42/EEC: 2007/47/EC concerning medical devices (MDD 93/42/EEC).

STANDARDS APPLIED: SEE ATTACHED LIST OF (HARMONISED - EN) STANDARDS FOR WHICH DOCUMENTED EVIDENCE OF COMPLIANCE CAN BE PROVIDED.

Notified Body: TÜV SÜD Product Service GmbH, Ridlerstrabe 65, 80339, MÜnchen, Germany

Identification number: CE0123

(EC) Certificate(s): G2 055329 0025 REV.00

Start of CE Marking: Date CE mark was affixed: 2020-03-30

Expire date of the Certificate: 2024-05-26

Place, Date of Issue: DanYang, JiangSu , P.R.CHINA 2022-06-01

Name: Bill Wang databat to the suitable to the

Position: Management Representative 江苏鱼跃医疗设备股份有限公司



LIST OF EU HARMONISED AND INERNATIONAL STANDARDS

S/N	Document No.	Edition	Title
1	93/42/EEC	2007/47/EC	Medical device directives of EU
2	ISO 13485	2016	Medical devices-Quality management systems-Requirements for regulatory purposes
3	ISO 14971	2007	Medical devices - Application of risk management to medical devices
4	EN ISO 10993-1	2009/AC:2010	Biological evaluation of medical devices – Part 1: Evaluation and testing
5	EN ISO 10993-5	2009	Biological evaluation of medical devices – Part 5: Tests doe in vitro cytotoxicity
6	EN ISO 10993-10	2013	Biological evaluation of medical devices – Part 10: Tests for irritation and delayed-type
7	EN ISO 15223-1	2016	Medical devices - Symbols to be used with medical device Labels, labeling and information to be supplied-Part1: general requirements
8	EN 1041	2008+A1:2013	Information supplied by the manufacturer with medical devices
9	EN 60601-1	2012	Medical electrical equipment - Part 1 : General requirements for basic safety and essential performance
10	EN 60601-1-2	2015	Medical electrical equipment – Part 1-2: General requirements for safety – Collateral standard: Electromagnetic compatibility – Requirements and tests
11	EN ISO 80601-2-69	2014	Oxygen concentrators for medical use – Safety requirements
12	EN 60601-1-6	2010	Medical electrical equipment -Part I-6: General requirements for basic safety and essential performance -Collateral Standard: Usability

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