## Chapter 9 EC DECLARATION OF CONFORMITY

The following EC declaration of conformity exemplifies the required content according to Directive 93/42/EEC.

Doc: 906-00522 rev.: A01 date: 2021-04-20

## EC DECLARATION OF CONFORMITY

| Name and address of the<br>manufacturer:<br>Name and address of the<br>European Representative<br>We declare under our sole responsibil   | SONOSCAPE MEDICAL CORP.<br>Room 201 & 202, 12th Building, Shenzhen Software Park Phase<br>II, 1 Keji Middle 2nd Road, Yuehai Subdistrict, Nanshan District,<br>Shenzhen, 518057, Guangdong, China<br>Shanghai International Holding Corp. GmbH (Europe)<br>Eiffestrasse 80, 20537 Hamburg, Germany<br>lity that   |
|---|---|
| the medical device:   | Digital Color Doppler Ultrasound System<br>Model: P60 Exp/P60 Pro/P60/P60 CV/ P70T/P70S/P60S/P60 VO/<br>P55/P55 Elite/ P55S/P50T/P50 Elite/P50E/P40T/P40 Elite/P40E/<br>P30T/P30 Elite/P30E/P25S/P22S<br>(Supported Probes: 3C-A, C1-6, C1-6A, C2-9, 12L-A, 12L-B,<br>13L-A, 9L-A, 18L-A, 10I2, 4P-A, S1-5, 7P-A, 8P1, VE9-5, VC6-<br>2, VC2-9, 6V3, C3-10V, 6V3A, 6V7, EC9-5, 6V1, BCC9-5,<br>BCL10-5, C322, C613, 12LT-A, 12LI-A, 6CT-A, 6CI-A,<br>CWD2.0, MPTEE, MPTEE mini, LAP7, L741, L3-9, 3P-A, 7P-<br>B) |
| of class: /   | IIa<br>according to annex IX of directive 93/42/EEC   |
| meets the provisions of the directive 93/42/EEC and its transpositions in national laws which apply to it. The declaration is valid in connection with the "final inspection report" of the device. |   |

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

Registration No.:

HD 2027206-1

Notified Body:

TÜV Rheinland LGA Products GmbH Tillystraße 2 90431 Nürnberg Deutschland CE 0197

Zhou Wenpin

Name and function

Vice President

Shenzhen, April 20, 2021

Place, date /

906-00522