## Chapter 9

## EC DECLARATION OF CONFORMITY

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

Doc: 906-01683 rev.: A01 date: 2022-2-21

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## EC DECLARATION OF CONFORMITY

Name and address of the manufacturer: Name and address of the European Representative	SONOSCAPE MEDICAL CORP. Room 201 & 202, 12th Building, Shenzhen Software Park Phase II, 1 Keji Middle 2nd Road, Yuehai Subdistrict, Nanshan District, Shenzhen, 518057, Guangdong, China Shanghai International Holding Corp. GmbH (Europe) Eiffestrasse 80, 20537 Hamburg, Germany
We declare under our sole responsibility that	
the medical device:	Digital Color Doppler Ultrasound System Model: P25 Exp/P25 Elite/P20 Exp/P20 Elite/P22 Elite/P22 Exp/P15 Exp/P15 Elite (Supported Probes: L741,10L-I,9L-A,12L-B,10I2,6CI-A,6CT- A,12LI-A,12LT-A,3C-A,C322,C613,BCC9-5,BCL10-5,VC6-2, C1-6,6V1,6V3A,6V7,6V3,EC9-5,3P-A,S1-5,LAP7,MPTEE, MPTEE mini,L742,2P1,CWD2.0,C1-5,C1-6A,C2-9,6V1A,12L- A,L752,4P-A,7P-A,8P1,VE9-5
of class: /	IIa
	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

Zhow Wenping Vice President

Shenzhen, Feb 21, 2022

Place, date /

Name and function