

EU Declaration of Conformity

2625 Patton Road Roseville, MN 55113 800-322-0956 www.ablenetinc.com

This declaration of conformity is issued under the sole responsibility of the manufacturer. The device covered by the present declaration is in conformity with all regulations below, including compliance with related Essential Requirements.

Object of the declaration:

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Product Name:	Quicktalker 7
Product Model Designator:	Quicktalker 7
Product Part Number(s):	10003501
Basic UDI-DI	00850011150139
<required for="" mdr=""></required>	
Control Indicator:	2020W20 thru 2025W20
Global Medical Device Nomenclature Code (GMDN) and Description	
Product Options/Accessories:	N/A



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The object of the declaration described above is in conformity with the following regulations:

EU Regulation	Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices
Device Classification:	Class I based on Annex VIII and Rule 13
Conformity Assessment Path	Not Applicable – Class I device
Name/Address/ID of Notified Body:	Not Applicable – Class I device
Standards and Common Specifications	The products listed above have been tested in a typical configuration as described in the Manufacturer's accompanying documentation and are compliant with the product standards listed below.
	EN 60601-1-2:2001 + A1:2006, Class B for Emissions, Immunity for Non Life-Supporting Equipment Risk Analysis per EN ISO 14971:2007



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Additional information:

EU Authorized	EUCEREP
Representative :	Roald Dahllaan 33
_	5629MC – Eindhoven
	The Netherlands

Signature (signed for and on behalf of AbleNet):	Date of Issue: <dd month="" yyyy=""> 13 May 2020</dd>
Printed Name:	Place of Issue:
Joe Volp	AbleNet Inc – Roseville, MN
Title: Director of Marketing	Document Number: DoC_10003501_Quicktalker 7_0513320