

Declaration of Conformity

Manufacturer	ArjoHuntleigh AB Hans Michelsensgatan 10 211 20 Malmö, Sweden		
Single Registration Number	SE-MF-00000696		
Declaration	ArjoHuntleigh AB as the manufacturer of the following medical devices, takes sole responsibility and declares conformity with the applicable provisions of Medical Device Regulation (EU) 2017/745 concerning medical devices, by Annex IX.		
Device Family Name	D900/SD2/MD2		
Intended Purpose	The product is intended to monitor assess blood flow.		
Basic UDI-DI	5060693520365WJ		
Risk Glass and Rule	Class IIa, Rule 10		
Additional Information	Manufactured and distributed on behalf of ArjoHuntleigh AB by: Huntleigh Healthcare Ltd 35 Portmanmoor Road Cardiff CF24 5HN United Kingdom Also complies with the following EU Legislation: RoHS Directive 2011/65/EU WEEE Directive 2012/19/EU		
Notified Body Name and Number	BSI Group The Netherlands B.V. Number: 2797 2797 CE Certificate Number MDR 718928		

	APPROVED BY	
Title: QRE Compliance Director	Signature:	Wind
Name: Steve Monks	Date:	19/12/2023

On behalf of ArjoHuntleigh AB; Place: Cardiff