

## **Declaration of Conformity**

| Manufacturer                     | ArjoHuntleigh AB<br>Hans Michelsensgatan 10<br>211 20 Malmö, Sweden  |
|----------------------------------|--|
| Single Registration              | SE-MF-000000696  |
| 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               | SF1 Toco Transducer (ACC-OBS-053)  |
| Intended Purpose                 | The product is intended to monitor fetal heart and vital signs of pregnant woman.  |
| Basic UDI-DI                     | 5060693520389WY  |
| Risk Class and Rule              | Class IIa, Rule 10   |
| Additional Information           | Manufactured and distributed on behalf of ArjoHuntleigh AB by:<br>Huntleigh Healthcare Ltd<br>35 Portmanmoor Road<br>Cardiff<br>CF24 5HN<br>United Kingdom<br>Also complies with the following EU Legislation:<br>RoHS Directive 2011/65/EU<br>WEEE Directive 2012/19/EU |
| Notified Body Name<br>and Number | CEBSI Group The Netherlands B.V.Number: 27972797CE Certificate Number MDR 718928   |

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

On behalf of ArjoHuntleigh AB: Place: Cardiff