

## EC Declaration of Conformity

Manufacturer: **HeartSine Technologies Limited**  
Canberra House  
203 Airport Road West  
Belfast, BT3 9ED  
UK


Device: **samaritan PAD 350P**  
Model: **SAM 350P**  
Description: Automated external defibrillator.  
Medical Device Classification: Identified as **Class IIb** under rule 9 of Annex IX of Council Directive 93/42/EEC as amended by 2007/47/EC

HeartSine Technologies declares that the HeartSine samaritan PAD 350P (SAM 350P), a therapeutic medical device in the range of Automated External Defibrillators, and its associated accessories are designed and manufactured in conformity with:

- a) the essential requirements (Annex I) and provisions of the European Medical Device Directive (MDD) **European Council Directive 93/42/EEC** (as amended by 2007/47/EC)
  - And is subject to the procedure set out in Annex II (excluding section 4), Full Quality Assurance System, of Directive 93/42/EEC, as amended by Directive 2007/47/EC;
  - Under the supervision of notified body number CE0123, **TÜV SÜD Product Service GmbH, Certification Body, Ridlerstraße 65, 80339 Munich, Germany.**
- Article 4 of ROHS2 Directive (2011/65 EU), with exceptions Annex III (6c – lead in copper alloy) and Annex IV (17 – lead in solder for portable emergency defibrillators).

<b>Certification</b>	<b>TÜV Certificate Number</b>
Council Directive 93/42/EEC	<b>No. G1 067590 0002 Rev. 00</b>
ISO 13485:2016	<b>No. Q5 067590 0001 Rev. 00</b>
EN ISO 13485:2016	<b>No. Q5 067590 0001 Rev. 00</b>

Signature

  
\_\_\_\_\_

Date

15 April 2019

Rebecca Funston

Senior Manager, Regulatory Affairs & Quality Assurance  
HeartSine Technologies Ltd.