

AliveCor, Inc.  
Ph:415-795-9811

Kardia Mobile

**DECLARATION OF CONFORMITY  
MEDICAL DEVICES**

We hereby declare that the products identified above are in conformity with all relevant provisions of Council Directive 93/42/EEC, as amended September 21, 2007 (M5), concerning Medical Devices and 2011/65/EU, concerning Restrictions of Hazardous Substances. Conformity to Directive 93/42/EEC is assessed by the notified body, LNE/G-MED. Conformity to Directive 2011/65/EU is by self-declaration.

This Declaration of Conformity is made under Annex II of this directive according to EC Conformity Certificate No. 28705, issued on May 14, 2015 and delivered by LNE-GMED located at 1 rue Gaston Boissier 75724 Paris Cedex 15, France (Notified Body No. 0459).

In addition, we ensure and declare that the distributed CE marked products, as mentioned and falling within Class IIa, Rule 10, meet the provisions of the EC-Directive which apply to them, including an Authorized Representative. The Authorized Representative is Obelis SA, located at BD General Wahis 53 1030, Brussels Belgium.

This declaration is based on the application of the Quality System approved for the manufacture and final inspection of the products concerned, in accordance with Annex II, excluding Section 4, of Directive 93/42/EEC. This declaration is supported by the Quality System certification based on the harmonized standards ISO 13485:2012, certificate number 28707, and ISO 13485:2003 with CMDCAS, certificate number 28706, both issued on January 14, 2015 and delivered by LNE/G-MED (Notified Body No. 0459).


AliveCor's Notified Body is LNE/G-MED located at 1 rue Gaston Boissier 75724 Paris Cedex 15, France.

This declaration covers the Kardia Mobile and concerns the following products:

Kardia Mobile, AC-009.

This declaration is valid for all products described here above, bearing the CE marking and manufactured at the following site(s):

AliveCor, Inc.  
30 Maiden Lane Suite 600  
San Francisco, CA 94108  
USA

  
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Arvita Tripathi  
Director of Quality  
AliveCor, Inc.

March 15, 2016  
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Date