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CE DECLARATION OF CONFORMITY

iCAD, Inc.
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Declares under sole responsibility that the product **SecondLook® Digital (Product No: DSC002-01, DSC002-02, DSC002-03, DSC002-04, DSC002-05, DSC002-07, DSC002-08, DSC002-09, DSC002-10, DSC002-11, DSC002-12, DSC002-13 and DSC002-19)** to which this declaration relates meets the essential health and safety requirements and is in conformance with the relevant EC directives listed below using the relevant section of the EC standards and other normative documents:

IEC 60950-1 (2001) First Edition
EN 55022:1994 +A1:1995 +A2:1997 Class A
EN 55024:1998 +A1:2002 +A2:2003
Medical Device Directive 93/42/EEC

MDD Class: Class I. The classification is based on the requirements of Rule 12 of Annex IX, of the Medical Device Directive.

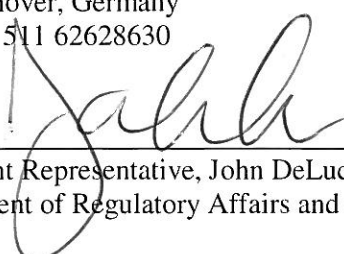
Quality System Certification: ISO 13485:2003 as indicated on certificate number 9213, granted by Intertek.

CE Mark: Article 17 of the Medical Device Directive 93/42/EEC.

Date: June 4, 2010

iCAD, Inc., Authorized Representative in the European Community (as defined in Article 14 of the medical Device Directive: 93/42/EEC):

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Management Representative, John DeLucia
Vice President of Regulatory Affairs and Quality Assurance