

EC Certificate - Full Quality Assurance System

Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No. CE 95140
Issued To: **Vital Images, Inc.**
5850 Opus Parkway
Suite 300
Minnetonka
Minnesota
55343
USA

In respect of:

The design and manufacture of software for medical imaging applications intended to allow visualization, analysis and communication of medical images and those aspects of Annex II concerned with the metrological requirements of software for medical imaging applications.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):



Frank Lee, EMEA Compliance & Risk Director

First Issued: **12 August 2005**

Date: **23 July 2015**

Expiry Date: **10 August 2020**

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Page 1 of 1

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.

This certificate was issued electronically and is bound by the conditions of the contract.

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List of Significant Subcontractors

Recognised as being involved in services relating to the product covered by:

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Subcontractor:	Service(s) supplied
Medical Device Safety Service (MDSS) GmbH Schiffgraben 41 30175 Hannover Germany	EU Representative
Toshiba Medical Systems Corporation (TMSC) 1385, Shimoishigami, Otawara-Shi, Tochigi 324-8550 Japan	Crucial Supplier
Toshiba Medical Visualization Systems Europe, Ltd (TMVS) 2 Bonnington Bond Anderson Place Edinburgh EH6 5NP United Kingdom	Crucial Supplier

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EC Certificate - Full Quality Assurance System Certificate History

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Date	Reference Number	Action
12 August 2005		First Issue Transfer from TUV PRODUCT SERVICE, Certificate No.: G2M.05.03.39815.004
12 July 2010	7475400	Certificate renewal and the addition of MediMark as EU representative
13 November 2012	7903620	Certificate reissue due to the reclassification by manufacturer of some software for medical imaging applications from Class I with measuring function to Class IIa
23 July 2015	8312538	Certificate renewal. Upgrade from Annex V to Annex II Scope change: The design and manufacture of software for medical imaging applications intended to allow visualization, analysis and communication of medical images and those aspects of Annex II concerned with the metrological requirements of software for medical imaging applications. Change of EU representative details to 'Medical Device Safety Service (MDSS) GmbH, Schiffgraben 41 , 30175 Hannover, Germany ' Addition of 'Toshiba Medical Systems Corporation (TMSC), Tochigi, Japan' and 'Toshiba Medical Visualization Systems Europe, Ltd (TMVS), Edinburgh, UK' as crucial suppliers