

The management system of

MED.E.COM s.a.r.l. trading as MEDECOM

9 bis - rue de Kerbrat, 29470 Plougastel Daoulas, France

has been assessed and certified as meeting the requirements of

Directive 93/42/EEC on medical devices, Annex II (excluding Section 4)

For the following products

**Digital diagnostic software and digital acquisition software for radiology.
Medical Image Processing and Communication Software.**

Where the above scope includes class III medical device(s), a valid EC Design Examination Certificate according to Annex II (Section 4) is a mandatory requirement for each device in addition to this certificate to place that device on the market.

This certificate is valid from 28 August 2018 until 28 June 2023
and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 16 May 2021

Issue 7. Certified since 29 June 2012

Certification is based on reports numbered FR/MD 217815

Authorised by

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