

EC CERTIFICATE

PRODUCTION QUALITY ASSURANCE SYSTEM CERTIFICATE

Cert no. 454001

issued to

Qualisys AB

Kvarnbergsgatan 2, 41105 GÖTEBORG, SWEDEN

We hereby certify that the Quality System of Qualisys AB for production, final inspection and marketing of Quality System of Qualisys AB for production, final inspection and marketing of Quality System of Quality System of Quality System for production, final inspection and marketing of Quality System of Quality System of Quality System for production, final inspection and marketing of Quality System of Quality System of Quality System for production, final inspection and marketing of Quality System of Quality System of Quality System for production, final inspection and marketing of Quality System of Quality System for production and Quality System of Quality System for production and Quality System of Quality System for production and Quality System of Quality System for Quality System

Qualisys Clinical Systems (including Qualisys registered capture cameras, hardware and related software)

medical devices in class Im has been assessed with respect to the conformity assessment procedure according to Annex V of Council Directive 93/42/EEC on Medical Devices, as latest amended by Council Directive 2007/47/EC is implemented in Swedish Law by the national regulation LVFS 2003:11, and found to comply with the requirements

 $The \ assessment \ has \ been \ restricted \ to \ the \ aspects \ of \ manufacture \ concerned \ with \ the \ conformity \ of \ the \ devices \ with \ meteorological \ requirements.$

This certificate applies to activities performed at

Kvarnbergsgatan 2, 411 05 Göteborg, SE-411 05 Göteborg, Sweden

 Originally issued
 2016-05-20

 Decision date
 2020-06-17

 Expiry date
 2024-05-26

Issued by Notified body 0402

Helén Dahl

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Conditions

Validity

The certificate will remain valid until the expiry date, and allows the holder to use RISE notified body identification number 0402 in conjunction with the CE-mark, on products covered by this certificate, provided that the conditions stated below are fulfilled:

- that surveillance audits are performed, with approved result;
- that the company notifies RISE on all modifications on the products, and that the company does not apply the CE-mark to any new or modified products without confirmation from RISE;
- that the company notifies RISE on all significant changes in the quality system, in its activities and/or organization;
- that the certificate is not used in a misleading manner, e.g. in marketing activities;
- that the company notifies RISE about vigilance actions, if any.

Basis for certificate

- The documentation presented has been examined and assessed by RISE in accordance with LVFS 2003:11, Annex V.
- An initial audit and follow-up audits of the quality system at the company's premises in Göteborg has been performed by RISE.
- RISE file Ecert ID 68518

Surveillance

RISE will perform surveillance inspections to ensure that the company maintains and applies the quality system that is subject of the certificate.

In accordance with the EU Commission recommendations of 2013-09-24, there will also be unannounced audits once per every three years. These audits can be performed at the manufacturers as well as at selected crucial supplier's premises.

Miscellaneous

Additional conditions appear in "RISE General Terms – Assignment" and "Rules and process assessment of medical devices as notified body LVFS 2003:11".

Certificate history

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Issue	Date	Activity	
1	2016-05-20	Original certificate	
2	2018-01-04	Certificate revised, accreditation mark added	
3	2020-05-13	Certificate revised, validity extended	
4	2020-06-17	Certificate revised due to technical error with digital signature	

Register of products covered by the certificate

Product	Art.no.	Class
Qualisys Clinical systems	1504xx	lm
(including Qualisys registered motion capture cameras, compatible hardware and related software)	(xx=01 u.i.99)	

Note: New products in **bold**