CERTIFICATE

The Certification Body TÜV Rheinland Italia S.r.l.

certifies, in accordance with the TÜV Rheinland Group procedures, that the Company

Qiagen S.r.l. Via Filippo Sassetti, 16 IT - 20124 Milano (MI)



has established and applies a quality management system for the following scope:

Trade of consumable and instrumentation for diagnostic use, both in the public and private sectors.

Trade of non-active medical devices in the urological and cytological fields, both in the public and private sectors.

Installation and assistance of diagnostic instrumentation.

Through an Audit, Report No. 7991500010MDA30, proof has been furnished that the quality management system fulfils the requirements of the standard

UNI CEI EN ISO 13485:2021

Please refer to the Quality Manual for the details about the exclusions with respect to the requirements of the standard.

Certificate Registration No. 39 05 1801510.

This Certificate is valid from 2024/09/05 to 2027/09/11.

The reference date for all the next audits is (day-month): 04/11.

Milan, 2024/09/05. First Certification: 2015/11/12

The certification responsible: Daniele Ricchi

TÜV Rheinland Italia S.r.I., Via E. Mattei, 3 - I - 20005 Pogliano Milanese (MI)



MS Nº 0083

Membro degli Accordi di Mutuo Riconoscimento EA, IAF e ILAC. Signatory of EA, IAF and ILAC Mutual Recognition Agreements.



Management EN ISO 13485:2016





the Directives 93/42/EEC, 90/385/EEC, 98/79/EC or

Regulations (UE) 2017/745, (UE) 2017/746 have been fulfilled

This certificate does not represent proof that the statutory requirements of