

Certificate

Quality Management System EN ISO 13485:2016 + AC:2018 + A11:2021 ISO 13485:2016

Registration No.: SX 2680897-1
Certificate Holder: DG Technologies GmbH
Ober der Mühle 39
42699 Solingen
Germany
Scope: Design and Development, Manufacture, Distribution and Service of analyzing systems used in in-vitro diagnostic applications.

The Certification Body of TÜV Rheinland LGA Products GmbH certifies that the organization has established and applies a quality management system for medical devices.
Proof has been furnished that the requirements specified in the abovementioned standard are fulfilled. The quality management system is subject to yearly surveillance.

Report No.: 1199657-100
Effective date: 2026-06-19
Expiry date: 2029-06-18
Issue date: 2026-06-19



Christina Suntrup
TÜV Rheinland LGA Products GmbH
Tillystraße 2 · 90431 Nürnberg · Germany

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The scope of certification also covers the following sites:

No.	Facility	Scope
/01	c/o DG Technologies GmbH Ober der Mühle 39 42699 Solingen Germany	Legal address
/02	c/o DG Technologies GmbH Bodenseeallee 20 78333 Stockach Germany	Design and Development, Manufacture, Distribution and Service of analyzing systems used in in-vitro diagnostic applications.

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