

Certificate

Quality Management System

ecm Zertifizierungsgesellschaft für Medizinprodukte in Europa mbH, Bismarckstr. 106, 52066 Aachen, Germany, hereby declares that an examination of the undermentioned quality assurance system has been carried out following the requirements of DIN EN ISO 13485:2016.



Through an audit performed on behalf of

Alto Products Im- und Export GmbH
Paul-Thomas-Str. 48, 40599 Düsseldorf, Germany

it could be demonstrated that a quality management system

according to **DIN EN ISO 13485:2016**
"Medical devices – Quality management systems –
Requirements for regulatory purposes"

for the **manufacture and distribution of medical
devices for hospital use and distribution
of medical merchandise**

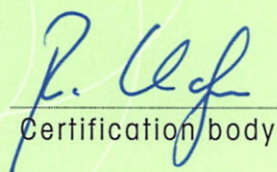
has been established and implemented.

This certificate is only valid under the conditions stated in the audit report mentioned hereafter.

Any substantial changes of the quality assurance have to be notified to ECM and are subject to a separate assessment.

Report number	Registered under	Valid until
181-21-823	Z/22/04778E	05 January 2025

Valid as of: 06 January 2022


Certification body