

Dear customer,

As an accredited certification body for ISO 13485 / EN ISO 13485, we require up-to-date information on your company for the preparation of quotations and for the planning and preparation of certification, extension and re-certification audits. This is required by the German Accreditation Body (DAkkS) at the beginning of each certification period and in case of significant changes in the scope of your certification. We kindly ask you to complete the questionnaire and to attach the required evidence.

1. General information

Company with legal form			
Street			
City		Postal code	
Country			
Contact person Title	☐ Mr ☐ Ms ☐ Diverse		
Contact person First- / Surname			
Function/Role		E-Mail	
Phone		Mobile	
Commercial register number		Industry	
Tax number non-EU countries		VAT ID	
Homepage			

2. Role(s) of the company (multiple answers possible)

☐ Manufacturer ☐ Authorised representative ☐ Importer	☐ System & Procedure Pack Producer ☐ Trader / Sales partner ☐ other (please describe):
--	---

3. Information on the number of employees at the main location / head office / parent company

Number of employees at the main location in full-time equivalents (FTE): For multi-site procedures: Full-time equivalents (FTE) in the scope of certification across all sites:		Number of shifts:	
		Please provide further information on locations in P11F006.	

4. What certifications are you aiming for?

Activity	Standard / service	
☐ Initial certification	☐ ISO 13485	☐ EN ISO 9001
	☐ EN ISO 13485	
☐ Recertification	☐ MDSAP (via TUV USA)	☐ MDR (separate questionnaire)
☐ Transfer	☐ Ukraine registration	
☐ Extension	☐ DIN EN ISO 13485 / KRINKO (Germany only; additional questionnaire P11F501)	
☐ Change (please use additional P11F002)	☐ Other:	

5. Scope of certification

e.g.: "Development, production and distribution of...".

Desired scope ISO 13485 / EN ISO 13485	Exclusions, sections not applicable
Desired scope ISO 9001 / EN ISO 9001	Exclusions, sections not applicable

6. Factors that could have an influence on the audit effort (multiple selection possible)

No Design / development	☐
Products with low risk or low process risk (e.g. simple processes, no validations necessary)	☐
Sophisticated management system	☐
High number of employees with the same, simple job	☐
Exclusively transport/trade within the scope of certification	☐
Products with high risk or high process risk (e.g. processes requiring validation)	☐
Large location with low employee density	☐
Installation activities at the customer	☐
High level of automation	
Identical activities in all shifts	
High proportion of outsourced activities	
High proportion of employees in the field	
Combined audit with several standards / guidelines / regulations	
Complicated logistics with multiple buildings, locations	
Interpreter needed	
In-house sterilisation	Method(s):
Have you been supported by a consultant when setting up your management system?	☐ Yes ☐ No
Consultancy	
Contact	
Have you received in-house training from a TÜV NORD company?	☐ Yes ☐ No
Training providers	
Training content	
When are you planning the audit?	

7. What type of certification is being sought? (Multiple selection possible)

Single-site certification (All locations are independently certified)	
Multi-site certification (Sites are certified in one group)	
Combined / Integrated Certification (Several management systems are audited simultaneously)	
Do you want a simultaneous audit of all management systems?	☐ Yes ☐ No
Would you like a remote audit (maximum 50% of the audit time)?	☐ Yes ☐ No
Do you have the necessary infrastructure for the remote audit?	☐ Yes ☐ No

8. Degree of integration when auditing several standards at the same time

In case of a simultaneous certification procedure with several standards, please fill in the following items:

Integrated management system documentation including procedural and work instructions	☐ Yes ☐ No
Management reviews that take into account the overall business strategy and corporate plan	☐ Yes ☐ No
An integrated approach to internal audits	☐ Yes ☐ No
An integrated approach to the organisation's policies and goals	☐ Yes ☐ No
An integrated approach to system processes (process descriptions)	☐ Yes ☐ No
An integrated approach to improvement mechanisms (corrective and preventive actions; measurement and continuous improvement)	☐ Yes ☐ No
Integrated management support and responsibilities (joint management representatives)	☐ Yes ☐ No

9. Information on the transfer of a certification from other certification bodies

Are the audit reports from the last certification period available?	☐ Yes ☐ No
Were there any non-conformities during the last audit?	☐ Yes ☐ No
Have all non-conformities from the last audit been closed?	☐ Yes ☐ No
Please attach the current certificates to be transferred in electronic form here:	
Why do you want to change the certifier?	

Note: In the case of an order to take over a certification, please enclose all certificates issued by the last certification body and relevant for transfer, all audit reports and reports on non-conformities of the last certification period.

10. Existing certifications

Please enter your existing certifications here.

Certificate number	Standard	Certification Body	Date of certification audit	Valid until

Questions mandatory to be answered by companies providing parts and/or services	
Is the product a nearly finished and assembled medical device? (i.e., it is intended to be used for a medical purpose and only needs packaging and/or labelling)	☐ Yes ☐ No
Is the product intended to be a component/part of a medical device?	☐ Yes ☐ No
Is the organization contracted to carry out any activities that are regulated by a medical device regulation (e.g., relabelling, remanufacturing of other medical devices)?	☐ Yes ☐ No
Is the product supplied sterile?	☐ Yes ☐ No
Does the product contain software developed by the client organization or a supplier?	☐ Yes ☐ No
Is "Design and Development" in the scope of the ISO 13485 / EN ISO 13485 certification?	☐ Yes ☐ No
Is the product (Raw Materials, Parts, Components, Subassemblies, Maintenance Services, or Other Services) intended to support associated medical devices?	☐ Yes ☐ No

11. Do you want to tell us something?

12. Documents required by TÜV NORD CERT for the preparation of the quotation and for the preparation for the (Re-)Certification or extension audit

- Extract from a professional or commercial register (or comparable evidence), if applicable
- Organisational chart/evidence of the organisational structure

**13. Note for audit planning of a (re)certification or extension audit:
Documents to be made available to the auditor or the audit team in advance**

- Extract from a professional or commercial register (or comparable evidence, if applicable)
- Management system documentation
(e.g.: Table of contents or presentation of the structure of the management system documentation)
- Organisational chart/evidence of the organisational structure
- Corporate policy
- Management review (e.g.: cover sheet or table of contents with date and signature)
- Current annual planning of internal audits and evidence of audit report(s) (e.g.: cover sheet with date and signature)
- As applicable: List of critical suppliers, certificates, quality agreements and evidence for supplier evaluation.
- Standard-specific documents, if applicable
- Procedural instructions in German or English language

We confirm all information and agree that our information will be stored as part of the offer preparation and process / order processing.

 Place and date

 Name

 Signature

Please email the completed form to: medical@tuev-nord.de

TÜV NORD CERT GmbH
 Certification Body and
 Notified Body for medical devices
 Am TÜV 1
 45307 Essen
 Germany

Phone: +49(0)201-825 2236
 E-mail: medical@tuev-nord.de

Appendix 1: Information on products, manufacturing technologies and services

Products Manufacturing Technologies and Services		
Classification of Main technical Areas (IAF MD 9) for ISO 13485 / EN ISO 13485 will be made by applying (EU) 2017/745 (MDR) codes. Please note, that this is not an MDR application and select as applicable: - Companies with Class I or higher products please select all applicable MDA/MDN and MDS product codes. - Companies manufacturing components or providing services, please select the most applicable MDT / MDS / S- codes for the most appropriate product technology or service(s).		
MDA, MDN, MDS, MDT		
	MDA	Active Implantable Products
☐	MDA 0101	Active implantable products for stimulation/inhibition/monitoring*
☐	MDA 0102	Active implantable devices delivering drugs or other substances
☐	MDA 0103	Active implantable devices substituting or replacing organ functions
☐	MDA 0104	Active implantable devices utilising radiation and other active implantable devices
	MDA	Active non-implantable devices for imaging, monitoring and/or diagnostic purposes
☐	MDA 0201	Active non-implantable imaging devices utilising ionizing radiation
☐	MDA 0202	Active non-implantable imaging devices utilising non-ionizing radiation
☐	MDA 0203	Active non-implantable devices for monitoring of vital physiological parameters
☐	MDA 0204	Other active non-implantable devices for monitoring and/or diagnosis
	MDA	Active non-implantable therapeutic products and general active non-implantable products
☐	MDA 0301	Active non-implantable devices utilising ionizing radiation
☐	MDA 0302	Active non-implantable devices utilising non-ionizing radiation
☐	MDA 0303	Active non-implantable devices utilising hyperthermia / hypothermia
☐	MDA 0304	Active non-implantable devices for shock-wave therapy (lithotripsy)
☐	MDA 0305	Active non-implantable devices for stimulation or inhibition
☐	MDA 0306	Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemopheresis
☐	MDA 0307	Active non-implantable respiratory devices
☐	MDA 0308	Active non-implantable devices for wound and skin care
☐	MDA 0309	Active non-implantable ophthalmologic devices
☐	MDA 0310	Active non-implantable devices for ear, nose and throat
☐	MDA 0311	Active non-implantable dental devices
☐	MDA 0312	Other active non-implantable surgical devices
☐	MDA 0313	Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport
☐	MDA 0314	Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitro fertilisation (IVF) and assisted reproductive technologies (ART)
☐	MDA 0315	Standalone software including software design for medical devices
☐	MDA 0316	Medical gas supply systems and parts thereof
☐	MDA 0317	Active non-implantable devices for cleaning, disinfection and sterilisation
☐	MDA 0318	Other active non-implantable devices

	MDN	Non-active implants and surgically invasive products for long-term use
☐	MDN 1101	Non-active cardiovascular, vascular and neurovascular implants
☐	MDN 1102	Non-active osteo- and orthopaedic implants
☐	MDN 1103	Non-active dental implants and dental materials
☐	MDN 1104	Non-active soft tissue and other implants
	MDN	Non-active non-implantable devices
☐	MDN 1201	Non-active non-implantable devices for anaesthesia, emergency and intensive care
☐	MDN 1202	Non-active non-implantable devices for administration, channelling and removal of substances, including devices for dialysis
☐	MDN 1203	Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters and related tools
☐	MDN 1204	Non-active non-implantable medical devices for wound and skin care
☐	MDN 1205	Non-active non-implantable orthopaedic and rehabilitation devices
☐	MDN 1206	Non-active non-implantable ophthalmologic devices
☐	MDN 1207	Non-active non-implantable diagnostic devices
☐	MDN 1208	Non-active non-implantable instruments
☐	MDN 1209	Non-active non-implantable dental materials
☐	MDN 1210	Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases
☐	MDN 1211	Non-active non-implantable devices for disinfecting, cleaning and rinsing
☐	MDN 1212	Non-active non-implantable devices for processing and preservation of human cells, tissue or organs including in vitro fertilisation (IVF) and assisted reproductive technologies (ART)
☐	MDN 1213	Non-active non-implantable devices composed of substances to be introduced into the human body via a body orifice or the dermal route
☐	MDN 1214	General non-active non-implantable devices used in health care and other non-active non-implantable devices
	MDS	Products with special properties
☐	MDS 1001	Devices incorporating medicinal substances
☐	MDS 1002	Devices manufactured utilising tissues or cells of human origin, or their derivatives
☐	MDS 1003	Devices manufactured utilising tissues or cells of animal origin, or their derivatives
☐	MDS 1004	Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC (from 2027: EU Machinery Regulation (EU) 2023/1230)
☐	MDS 1005	Devices in sterile condition: ☐ EO ☐ Radiation (gamma, electron, X-rays) ☐ Moist heat ☐ Hydrogen peroxide ☐ Aseptic filling ☐ Formaldehyde incl. low-temperature steam-formaldehyde sterilization ☐ Thermal sterilization processes, dry heat ☐ Plasma
☐	MDS 1006	Reusable surgical instruments
☐	MDS 1007	Devices incorporating or consisting of nanomaterial

☐	MDS 1008	Devices utilising biological active coatings and / or materials or being wholly or mainly absorbed or locally dispersed in the human body or are intended to undergo a chemical change in the body.
☐	MDS 1009	Devices incorporating software / utilising software / controlled by software, including devices intended for controlling, monitoring or directly influencing the performance of active or active implantable devices
☐	MDS 1010	Devices with a measuring function
☐	MDS 1011	Devices in systems or procedure packs
☐	MDS 1012	Products without an intended medical purpose listed in Annex XVI of Regulation (EU) 2017/745
☐	MDS 1013	Class III custom-made implantable devices
☐	MDS 1014	Devices incorporating as an integral part an in vitro diagnostic medical device
	MDT	Products for which special technologies or processes are used
☐	MDT 2001	Metal processing
☐	MDT 2002	Plastic processing
☐	MDT 2003	Non-metal mineral processing including glass, ceramics
☐	MDT 2004	Non-metal non-mineral processing including textiles, rubber, leather, paper
☐	MDT 2005	Biotechnology
☐	MDT 2006	Chemical processing
☐	MDT 2007	Production of pharmaceuticals
☐	MDT 2008	Clean room production
☐	MDT 2009	Processing of materials of human or animal origin (no accreditation for human material).
☐	MDT 2010	Manufacture or processing of electronic components including communication devices
☐	MDT 2011	Packaging, including labelling
☐	MDT 2012	Installation, refurbishment
☐	MDT 2013	Reprocessing of medical devices
	IVD	In vitro diagnostics Reagents and reagent products, calibrators, and control materials for
☐	IVD 0001	Clinical Chemistry
☐	IVD 0002	Haematology, haemostaseology, immunohaematology
☐	IVD 0003	Immunochemistry, immunology
☐	IVD 0004	IVD instruments and software
☐	IVD 0006	IVD medical devices other than specified above: specimen receptacles
	S	S-Codes
☐	S 0010	Raw materials (Raw metals, plastic, wood, ceramic)
☐	S 0011	Components (Electrical components, fasteners, shaped raw materials, machined raw materials, and molded plastic)
☐	S 0012	Subassemblies (Electronic subassemblies mechanical subassemblies, made to drawings and/or work instructions)
☐	S 0013	Distribution services (Distributors providing storage and delivery of medical devices, not acting as a 'legal manufacturer' for medical devices)
☐	S 0014	Maintenance services (Electrical or mechanical repair services, facility cleaning and maintenance services, uniform cleaning and testing of ESD smocks)
☐	S 0015	Transportation services (Trucking, shipping, air transportation service in general)
☐	S 0016	Custom made devices (dental technology, orthopaedics and orthopaedic shoe technology, rehabilitation technology)

☐	S 0017	Activities of economic operators
☐	S 0018	Administrative, supporting activities for manufacturers
☐	S 0019	Consultancy services on general medical device regulatory topics
☐	S 0020	Reprocessing of medical devices up to and including critical B
☐	S 0021	Reprocessing of medical devices up to and including critical C (TÜV NORD CERT Certificate will be issued)

For Scopes highlighted in grey, TÜV NORD CERT does not have an accreditation.