



Please send two PDF versions of this form to [med.certification.de@dekra.com](mailto:med.certification.de@dekra.com):

1. By "save-as" function
2. With signature

<b>1. Applicant</b>
Registered company name
House no., street
Further address, if necessary
Town (and region if necessary)
ZIP code/Postcode
Country

<b>2. Please enter your company details below.</b>	
Website address	
E-mail address for general enquiries	
Managing director/owner	
First name	Last name

Contact person		
	First name	Last name
Contact person's job title	Head of quality assurance other:	Executive management
Contact person's e-mail address		
Contact person's phone number		
Contact person's fax number		
Tax code number/VAT ID		

**NOTE: Please include a copy of your current trade register excerpt.**

**3. Please click on the box next to the certification basis according to which you need certification.**

EN ISO 13485:2016

ISO 9001:2015 (only possible in combination with one of the displayed certification bases)

ISO 13485:2016 TCP, for acceptance in Taiwan

ISO 13485:2016 MDSAP (Medical Device Single Audit Program)\*

Please complete the relevant sheets in F-091-57 (see 11. Details of the medical devices which are covered by the certification).

\* Certification is done through DEKRA B.V., The Netherlands

(EU) 2017/745 MDR Annex IX, chapter I EU-Quality Management System

(EU) 2017/745 MDR Annex IX, chapter II EU technical documentation assessment certificate

(EU) 2017/745 745 MDR Annex XI part A production quality assurance applicable to class IIa devices

(EU) 2017/745 MDR Article 120

(EU) 2017/745 MDR Article 16

(EU) 2017/746 IVDR Annex IX, chapter I EU Quality Management System

(EU) 2017/746 IVDR Annex IX, chapter II EU technical documentation assessment certificate

**4. Please detail your company's activities below.**

What does your company do?  
In which areas are you active?  
Please give a short description.

Which of the following functions  
does your company carry out?  
Please click all the relevant boxes.

Design and development of:

Production of:

Distribution of:

Installation of:

Servicing of:

Requested Scope EN ISO 13485:2016
Requested Scope ISO 9001:2015
Requested Scope EN ISO 13485:2016 TCP
Requested Scope ISO 13485:2016 MDSAP

<b>5. Do you hold any of the following certificates?</b>			
Yes, as follows:	ISO 9001 Directive 93/42/EWG (EU) 2017/745 MDR ISO 13485:2016 TCP Other:	EN ISO 13485 Directive 98/79/EG (EU) 2017/746 IVDR ISO 13485:2016 MDSAP	No

**NOTE: Please attach a copy/copies of your certificate(s) and the audit report(s) covering the current certification period.**

<b>6. Have you used consultancy services to help you establish the QM System which you are applying for?</b>	
Yes, and their name is:	No

<b>7. Is your production split into different shifts?</b>	
Number of shifts in production:	Do all the shifts carry out the same processes? Yes                      No
	How are the shifts established? Rolling                      Fixed night shift

When would you like the certification audit to take place?	
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9. Languages - please click the relevant boxes.		
What language is your QM documentation in?	German	English
What language is/are your technical documentation(s) in?	German	English
What language should the audits be conducted in?	German	English
What language should the reports be written in?	German	English

**NOTE: Documents will only be accepted in German or English.**

10. Please give details below of the headquarters and its employees.	
Location/Site (A1)	
Main site/Headquarters (A1)	
Company name	
House no., street	
Town (and region if necessary)	
ZIP code/Postcode	
Country	
Number of shifts	



10a. Please give details below about the number of employees at this location.				
	Employees who work between 21 and 40 hours per week on average	Employees who work between 11 and 20 hours per week on average	Employees who work between 1 and 10 hours per week on average	Trainees
Design and development				
Production and warehousing				
Administration, purchasing & miscellaneous				
Quality management, regulatory affairs				

10b. Are some of your services carried out at your customer’s premises (projects)?		
No	Yes, as follows:	Number:

**NOTES:**

- Please attach a copy of your company’s current organization chart.
- For documentation of additional sites, please use form C-031-07 and submit with this application.

11. Details of the medical devices which are covered by the certification.
See F-091-57 dated (YYYY-MM-DD):      JJJJ-MM-TT



**NOTE:**

- In order for us to create your individually tailored offer according to MDR/IVDR we also need you to submit the completed form F-09 1-57 "Customer data Sheet" (CDS).
- This CDS is the basis for the certification for which you are applying.
- The date entered in section 11 must correspond to the date of revision in the CDS.

**The following documents must be submitted together with this application:**

- Extract from the trade register excerpt
- Organization chart
- Data sheet F-09 1-57 "Customer Data Sheet" (if relevant)
- Current reports/certificates (only for new customers)

**12. Confirmation**

Date

Name in CAPTIAL LETTERS

Digital ID/Signature

