

## List of Standard Fees According to Regulation (EU) 2017/745

This document aims to provide general guidance on the fee structure of SZUTEST Konformitätsbewertungsstelle GmbH for (EU) 2017/745 (MDR) services. Once the pre-application forms are provided, we would be happy to calculate pricing and provide you with a detailed quotation. For any inquiries, you may send us an email via [mdsales@szutest-germany.de](mailto:mdsales@szutest-germany.de).

### A) APPLICATION FEES

Product Risk Class	Fees	Type	Remarks
Class Is, Im, Ir and Sterile Systems or Procedure Packs	1000 Euro	FLAT	If more than one device is involved, the fee for the highest product risk class is applied.
Class IIa	1500 Euro		
Class IIb non-implantable	2000 Euro		
Class IIb implant	2500 Euro		
Class IIb active devices intended to administer and/or remove a medicinal product	3000 Euro		
Class III	3500 Euro		
Class III implantable	4000 Euro		

### B) ANNUAL CERTIFICATE USAGE FEE

Product Risk Class	Fees	Type	Remarks
Class Is, Im, Ir, and Sterile Systems or Procedure Packs	5000 Euro	FLAT	Annual certificate usage fee covers routine annual administrative maintenance of valid certificates, including certificate record upkeep, certificate status management, routine administrative follow-up, and maintenance of related certification data.  If more than one device is involved, the fee for highest product risk class is applied.  This fee is calculated for calendar year.
Class IIa	7000 Euro		
Class IIb non-implantable	9000 Euro		
Class IIb implantable	11000 Euro		
Class IIb active devices intended to administer and/or remove a medicinal product	13000 Euro		
Class III	15000 Euro		
Class III implantable	17000 Euro		

### C) AUDIT MAN/DAY FEE

Location	Fees	Type	Remarks
EU Countries	2500 Euro	DAILY (MAN/DAY)	The number of man-days required for the audit is calculated mainly based on IAF MD 9 by applying several increasing and decreasing factors. One man-day is typically calculated as 8 hours.
Others	2500 Euro		

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**D) TECHNICAL DOCUMENTATION REVIEW MAN/DAY FEE**

Location	Fees	Type	Remarks
EU Countries	2500 Euro	DAILY (MAN/DAY)	Time spent will be calculated based on the product risk class. The following factors will increase the duration per device, - PSUR, PMCF, SSCP, PMS Reviews - Routine reviews on Technical Documentation changes - Devices in sterile condition and the number of applied sterilization methods - Devices requiring biocompatibility review - Devices incorporating software - Devices that are absorbable or locally dispersed - Pre-market clinical investigation review - Medicinal Product Authority Consultation - Clinical Evaluation Consultation Procedure - Consultation procedure for devices that are systemically absorbed
Others	2500 Euro		

**E) ADMINISTRATIVE AND OTHER FEES**

Task	Fees	Type	Remarks	
Initial review on changes	Class Is, Im, Ir and Sterile Systems or Procedure Packs	100 Euro	FLAT	If more than one device is involved in the certification scope, the fee for the highest product risk class is applied.
	Class IIa	150 Euro	FLAT	
	Class IIb non-implantable	200 Euro	FLAT	
	Class IIb implantable	250 Euro	FLAT	
	Class IIb active devices intended to administer and/or remove a medicinal product	250 Euro	FLAT	
	Class III	300 Euro	FLAT	
	Class III implantable	350 Euro	FLAT	
Administrative task for outgoing transfers	500 Euro	FLAT	Fees to be paid to the authorities are to be invoiced separately based on the rates available during the consultation process.	
Preparation and follow-up activities for authority consultations	100 Euro	HOURLY		
Assessment on appeals	100 Euro	HOURLY		
Travel Time (excluding travel and accommodation expenses)	50 Euro	HOURLY		

**F) FEES FOR REPEATING NON-CONFORMITY CORRECTION REVIEW**

Task	Fees	Type	Remarks
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Repeating Non-conformity Correction Review	350 Euro	FLAT (Per each repeating non-conformity review)	The contract will include a one-time review of the non-conformities. For repeating non-conformity reviews within a defined deadline, the repeating review fee will be invoiced separately. Repeating reviews will not be conducted once the deadlines are reached.
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**G) FEE CALCULATION PARAMETERS**

Type of Assessment	Application Fee	Annual Certificate Usage Fee	Audit Fee	Technical Documentation Review Fee
Initial Assessment	+	+	+	+
Surveillance Assessment	-	+	+	+
Re-Assessment	+	+	+	+
Transfer Assessment (From another Notified Body to SZUTEST)	+	+	+	0
Transfer Assessment (From SZUTEST to another Notified Body)	***	-	-	-
Change Assessment	**	-	0	0
Scope Extension Assessment	+	*	0	0
Unannounced Site Audit	-	-	+	-
Follow-Up Audit	-	-	+	-

'o': Optional    '+': to be calculated    '-': not to be calculated.

\* For higher product classes

\*\* The application fees to be invoiced for change assessment are given in section E.

\*\*\* The expenses to be invoiced for administrative work in case of a transfer from SZUTEST to another notified body is 500 EUR.

**H) SPECIAL CONDITIONS FOR MANUFACTURERS BELONGING TO SMEs AS DEFINED IN RECOMMENDATION 2003/361/EC**

%3 of discount is applied for SMEs from the total initial and re-certification contract amount.

**i) HOW TO CALCULATE AUDIT DURATION**

Effective Number of Employees	Total Duration (man/day)
1-5	3
6-10	4
11-15	4,5
16-25	5
26-45	6
46-65	7
66-85	8
Refer to IAF MD-9 for a greater number of effective employees	

Once the total audit duration is calculated, 1/3 will be allocated to surveillance audits and 2/3 to re-certification audits. For unannounced audits, the audit duration is 2 man/days.

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Increase and decrease factors are applied in line with the Team-NB Code of Conduct.

### J) HOW TO CALCULATE TECHNICAL DOCUMENTATION REVIEW DURATION

Normal Clinical Evaluation Duration		3 man/day				
Type	Code	Normal	If systems (man/day)	If systems cover more than 5 devices (man/day)	If WET	If similar devices are combined under similar intended use
Active (Diagnostic)	MDA0201-0204	3	5	+2	N/A	+2 For TD Review +1 For Clinical
Active (Therapeutic)	MDA0301-0318	3	5	+2	N/A	+2 For TD Review +1 For Clinical
Implantable	MDN1101-1104	4	6	+2	-1	+2 For TD Review +1 For Clinical
Non-Implantable	MDN1201-1214	3	6	+2	N/A	+2 For TD Review +1 For Clinical
<b>Notes</b> 1- Systems: Systems consist of several components, which are medical devices on their own right; however, every component of the system shall be reviewed at the same time.						

#### Additional Conditions Applicable for Class IIa and Higher Devices (Applies together with the above table)

Condition	Type/Code	Result	Remarks
Class IIb active device administering/removing medicinal products	Rule 12	+1 man/day	Not whole rule 12 class IIb devices
Sterile Devices	MDS1005	+0,5 man/day per sterilization type	N/A
Devices requiring biocompatibility review	N/A	+1 man/day	N/A
Software	MDS1009	+1 man/day	N/A
Locally dispersed absorbed or biological coating	MDS1008	+1 man/day	Is not applicable if the device is systemically absorbed
Machinery	MDS1004	+0,5 man/day	N/A
CECP- For new class III and rule 12 devices	Article 54	+2 man/day	Only for new devices
Medicinal product consultation	Rule 14	+2 man/day per medicinal product	N/A

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<b>Devices that are systemically absorbed</b>	Rule 21	+2 man/day per substance	N/A
<b>New devices with pre-market clinical investigation</b>	N/A	+2 man/day	Not applicable for legacy devices and devices with PMCF investigation
<b>Orphan Devices</b>	N/A	+2 man/day	N/A
<b>Nanomaterial</b>	MDS1007	+1 man/day	N/A
<b>Annex XVI Devices</b>	N/A	+1 man/day	N/A

### Fixed Technical Documentation Review Durations for Class Is, Im, Ir Devices

Type	Duration (man/day)
Class Is	2
Class Im	2
Class Ir	2

### Fixed Technical Documentation Review Durations for Sterile Systems/Procedure Packs

Normal Duration	If large range of diverse devices covered	If more than 1 sterilization type
2 man/day	4 man/day	+1 man/day per additional sterilization type

For surveillance review of Technical Documentation, 0,5 to 1 day review may be applicable for certain devices and conditions.

## K) PRICING CALCULATION EXAMPLE

Scenario 1
<b>Product Name:</b> Surgical Instrument (Class Ir), Surgical Instrument Bone Cutting System (Class IIa), Plate and Screw System (Class IIb Implantable WET)
<b>Product Count:</b> 3
<b>Product MDA/MDN Code:</b> MDN1208, MDN1102
<b>Number of Employees:</b> 86
<b>Sampling Applied:</b> No
<b>Critical Supplier:</b> 2
<b>Clinical Evaluation Assessment:</b> Yes
<b>Biocompatibility Review:</b> Yes

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**Sterilization Review:** N/A

**Applied Increasing Factors For QMS Audit Duration:** Critical Supplier +1 M/D

**Applied Decreasing Factors For QMS Audit Duration:** The company is already certified -%15

**FEE FOR INITIAL CERTIFICATION**

Application Fee	2500 EUR
QMS Audit Fee	9,5 M/D X 2500 EUR
TD Review Fee	16 M/D X 2500 EUR
Annual Certificate Usage Fee	11000 EUR
<b>Total</b>	<b>77250 EUR</b>

**FEE FOR EACH SURVEILLANCE**

QMS Audit Fee	3 M/D X 2500 EUR
TD Review Fee	0,5 M/D X 2500 EUR
Annual Certificate Usage Fee	11000 EUR
<b>Total</b>	<b>19750 EUR</b>

**FEE FOR UNANNOUNCED AUDIT**

QMS Audit Fee	2 M/D X 2500 EUR
<b>Total</b>	<b>5000 EUR</b>

**Scenario 2**

**Product Name:** Feeding Catheter (Class IIa), Nasogastric Catheter (Class IIa), Nelaton Catheter (Class IIa)

**Product Count:** 3

**Product MDA/MDN Code:** MDN1202

**Number of Employee:** 16

**Sampling Applied:** Yes

**Clinical Evaluation Assessment:** Yes

**Biocompatibility Review:** Yes

**Sterilization Review:** Yes

**Critical Supplier:** 0

**Applied Increasing Factors For QMS Audit Duration:** Sterilization +0,5 M/D

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**Applied Decreasing Factors For QMS Audit Duration: Low risk product -%15**

**FEE FOR INITIAL CERTIFICATION (Sampling- Feeding Catheter (Class IIa))**

Application Fee	1500 EUR
QMS Audit Fee	5 M/D X 2500 EUR
TD Review Fee	7,5 M/D X 2500 EUR
Annual Certificate Usage Fee	7000 EUR
<b>Total</b>	<b>39750 EUR</b>

**FEE FOR SURVEILLANCE 1 (Sampling- Nasogastric Catheter (Class IIa))**

QMS Audit Fee	1,5 M/D X 2500 EUR
TD Review Fee	7,5 M/D X 2500 EUR
Annual Certificate Usage Fee	7000 EUR
<b>Total</b>	<b>29500 EUR</b>

**FEE FOR SURVEILLANCE 2 (Sampling- Nelaton Catheter (Class IIa))**

QMS Audit Fee	1,5 M/D X 2500 EUR
TD Review Fee	7,5 M/D X 2500 EUR
Annual Certificate Usage Fee	7000 EUR
<b>Total</b>	<b>29500 EUR</b>

**FEE FOR SURVEILLANCE 3**

QMS Audit Fee	1.5 M/D X 2500 EUR
TD Review Fee	0,5 M/D X 2500 EUR
Annual Certificate Usage Fee	7000 EUR
<b>Total</b>	<b>12000 EUR</b>

**FEE FOR SURVEILLANCE 4**

QMS Audit Fee	1.5 M/D X 2500 EUR
TD Review Fee	0,5 M/D X 2500 EUR
Annual Certificate Usage Fee	7000 EUR
<b>Total</b>	<b>12000 EUR</b>

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**FEE FOR UNANNOUNCED AUDIT**

<b>QMS Audit Fee</b>	2 M/D X 2500 EUR
<b>Total</b>	5000 EUR

**Scenario 3**

**Product Name:** Medical Device Disinfectant (Class IIb), Medical Device Disinfectant (Class IIa), Hemodialysis solutions and concentrates (Class IIb), Endoscope Washer and Disinfector Device (Class IIb)

**Product Count:** 4

**Product MDA/MDN Code:** MDN1211, MDN 1202, MDA0317

**Number of Employee:** 45

**Sampling Applied:** No

**Clinical Evaluation Assessment:** Yes

**Biocompatibility Review:** N/A

**Sterilization Review:** N/A

**Machinery (MDS 1004):** Yes

**Applied Increasing Factors For QMS Audit Duration:** Requiring interpreter(s) +%10, Require visiting temporary sites +0,5 M/D

**Applied Decreasing Factors For QMS Audit Duration:** Maturity of the management system -%20

**FEE FOR INITIAL CERTIFICATION**

<b>Application Fee</b>	2000 EUR
<b>QMS Audit Fee</b>	6 M/D X 2500 EUR
<b>TD Review Fee</b>	24,5 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	9000 EUR
<b>Total</b>	87250 EUR

**FEE FOR EACH SURVEILLANCE**

<b>QMS Audit Fee</b>	2 M/D X 2500 EUR
<b>TD Review Fee</b>	0,5 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	9000 EUR
<b>Total</b>	15250 EUR

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**FEE FOR UNANNOUNCED AUDIT**

<b>QMS Audit Fee</b>	2 M/D X 2500 EUR
<b>Total</b>	5000 EUR

**Scenario 4**

**Product Name:** Dental implant (Class IIb Implantable WET), Surgical Instrument Set (Class IIa)

**Product Count:** 2

**Product MDA/MDN Code:** MDN1103, MDN 1208

**Number of Employee:** 175

**Sampling Applied:** No

**Clinical Evaluation Assessment:** Yes

**Biocompatibility Review:** Yes

**Sterilization Review:** Yes

**Applied Increasing Factors For QMS Audit Duration:** Critical Supplier +0,5 M/D, Sterilization +0,5 M/D, Highly complex processes +%10

**Applied Decreasing Factors For QMS Audit Duration:** Company is already certified -%15

**FEE FOR INITIAL CERTIFICATION**

<b>Application Fee</b>	2500 EUR
<b>QMS Audit Fee</b>	11,5 M/D X 2500 EUR
<b>TD Review Fee</b>	15 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	11000 EUR
<b>Total</b>	79750 EUR

**FEE FOR EACH SURVEILLANCE**

<b>QMS Audit Fee</b>	4 M/D X 2500 EUR
<b>TD Review Fee</b>	0,5 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	11000 EUR
<b>Total</b>	22250 EUR

**FEE FOR UNANNOUNCED AUDIT**

<b>QMS Audit Fee</b>	2 M/D X 2500 EUR
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<b>Total</b>	5000 EUR
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Scenario 5	
<b>Product Name:</b> Ablation Catheter (Class III)	
<b>Product Count:</b> 1	
<b>Product MDA/MDN Code:</b> MDN1203	
<b>Number of Employee:</b> 10	
<b>Sampling Applied:</b> No	
<b>Clinical Evaluation Assessment:</b> Yes	
<b>Biocompatibility Review:</b> Yes	
<b>Sterilization Review:</b> Yes	
<b>Pre-Market Clinical Investigation Review For New Devices:</b> Yes	
<b>Applied Increasing Factors For QMS Audit Duration:</b> Audit scope including Class III devices +%10 M/D, Sterilization +0,5 M/D	
<b>Applied Decreasing Factors For QMS Audit Duration:</b> N/A	
<b>FEE FOR INITIAL CERTIFICATION</b>	
Application Fee	3500 EUR
QMS Audit Fee	5 M/D X 2500 EUR
TD Review Fee	9,5 M/D X 2500 EUR
Annual Certificate Usage Fee	15000 EUR
<b>Total</b>	<b>54750 EUR</b>
<b>FEE FOR EACH SURVEILLANCE</b>	
QMS Audit Fee	1,5 M/D X 2500 EUR
TD Review Fee	1 M/D X 2500 EUR
Annual Certificate Usage Fee	15000 EUR
<b>Total</b>	<b>21250 EUR</b>
<b>FEE FOR UNANNOUNCED AUDIT</b>	
QMS Audit Fee	2 M/D X 2500 EUR
<b>Total</b>	<b>5000 EUR</b>

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**Scenario 6**

**Product Name:** Antibacterial Absorbable Suture (Class III incorporating medicinal product), Absorbable Suture (Class III)

**Product Count:** 2

**Product MDA/MDN Code:** MDN1104

**Number of Employee:** 75

**Clinical Evaluation Assessment:** Yes

**Biocompatibility Review:** Yes

**Sterilization Review:** Yes

**Clinical Evaluation Consultation Procedure (Article 54) CECP For new Class III and rule 12 devices:** Yes

**Competent Authority Medicinal Product Consultation:** Yes

**Pre-Market Clinical Investigation Review For New Devices:** Yes

**Absorbable-Locally Dispersed-Biological Coating Product Review:** Yes

**Applied Increasing Factors For QMS Audit Duration:** Audit scope including Class III devices +%10 M/D, Sterilization +0,5 M/D, Critical Supplier +0,5 M/D

**Applied Decreasing Factors For QMS Audit Duration:** N/A

**FEE FOR INITIAL CERTIFICATION**

<b>Application Fee</b>	4000 EUR
<b>QMS Audit Fee</b>	10 M/D X 2500 EUR
<b>TD Review Fee</b>	29 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	17000 EUR
<b>Total</b>	118500 EUR

**FEE FOR EACH SURVEILLANCE**

<b>QMS Audit Fee</b>	3,5 M/D X 2500 EUR
<b>TD Review Fee</b>	2 M/D X 2500 EUR
<b>Annual Certificate Usage Fee</b>	17000 EUR
<b>Total</b>	30750 EUR

**FEE FOR UNANNOUNCED AUDIT**

<b>QMS Audit Fee</b>	2 M/D X 2500 EUR
<b>Total</b>	5000 EUR