

Medical Devices



Fees for Conformity Assessment Activities (EUR)

Medical Devices Regulation (MDR and IVDR)

Effective 1 January 2022



Medical Devices

Conformity assessment activities and their fees

Quality System Audit Fees:

Medical Device QMS Audit (Exclusive of travel time and expenses)

€245 per hour

Technical Documentation Review Fees

The review of technical documentation requires the highest levels of technical expertise. The time spent on these reviews is dependent on a number of critical factors:

- The quality and completeness of the submission
- Type of the device
- Whether the device is novel and/or high risk
- Level of review required

Regulation (MDR/IVDR) Technical Documentation Review*

€420 per hour

Unannounced Audit Fees:

BSI is required to perform routine unannounced audits of manufacturers and/or their critical sub-contractors or crucial suppliers. Duration of unannounced audits is at least one day and typically performed by two auditors, at least once during the 5-year certification cycle with some exceptions.

Unannounced Audit (performed by 2 auditors for 1 day) (Exclusive of travel time and expenses)

€6,665 per day

Fees may vary slightly due to currencies and different travel policies that may apply to some specific geographies.

BSI Group EMEA

Kitemark Court Davy Avenue Knowlhill Milton Keynes MK5 8PP United Kingdom

T: +44 (0)345 080 9000

 $\hbox{\bf E: eu.medical devices@bsigroup.com}\\$

BSI Group The Netherlands B.V.

Say Building John M. Keynesplein 9 1066 EP Amsterdam The Netherlands

T: +31 (0)20 346 07 80

E: eu.medicaldevices@bsigroup.com

For external use

...making excellence a habit."

^{*}Clinical Documentation Review forms part of the Technical Documentation Review.