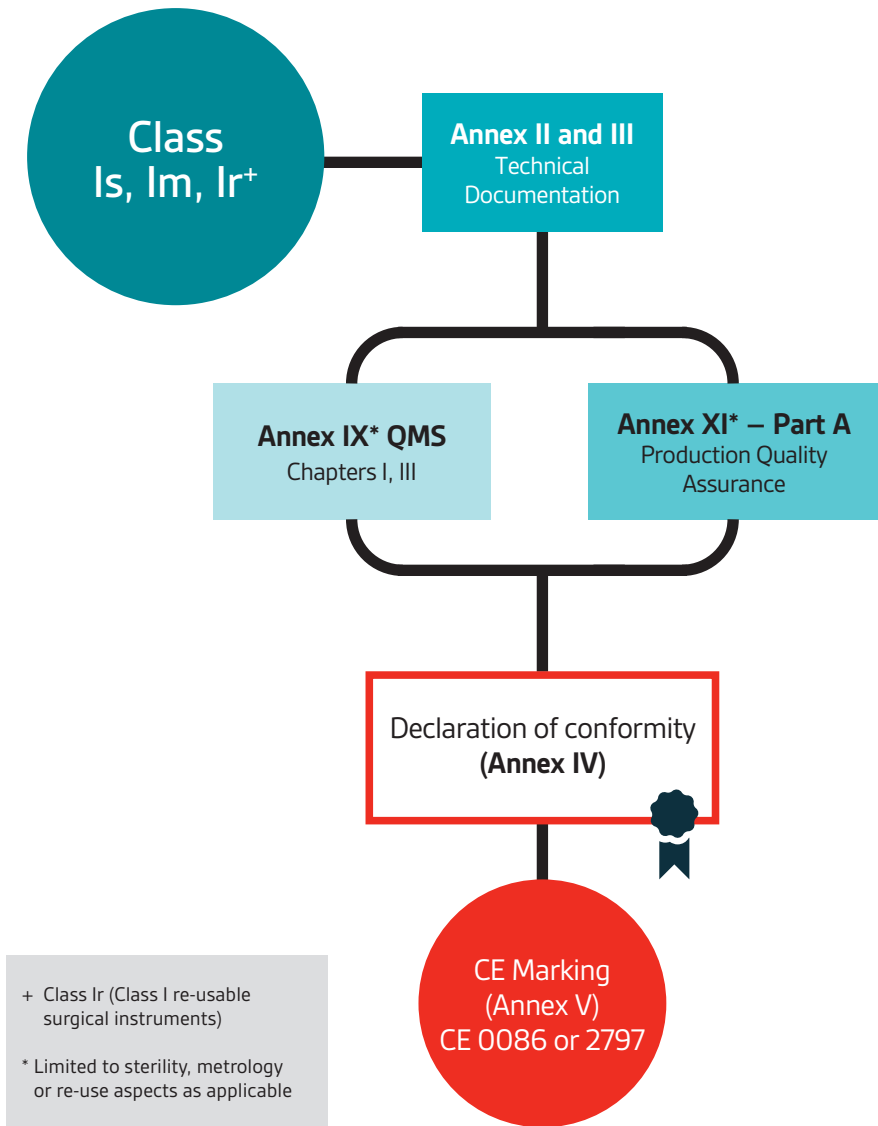


Class Is/Im/Ir devices



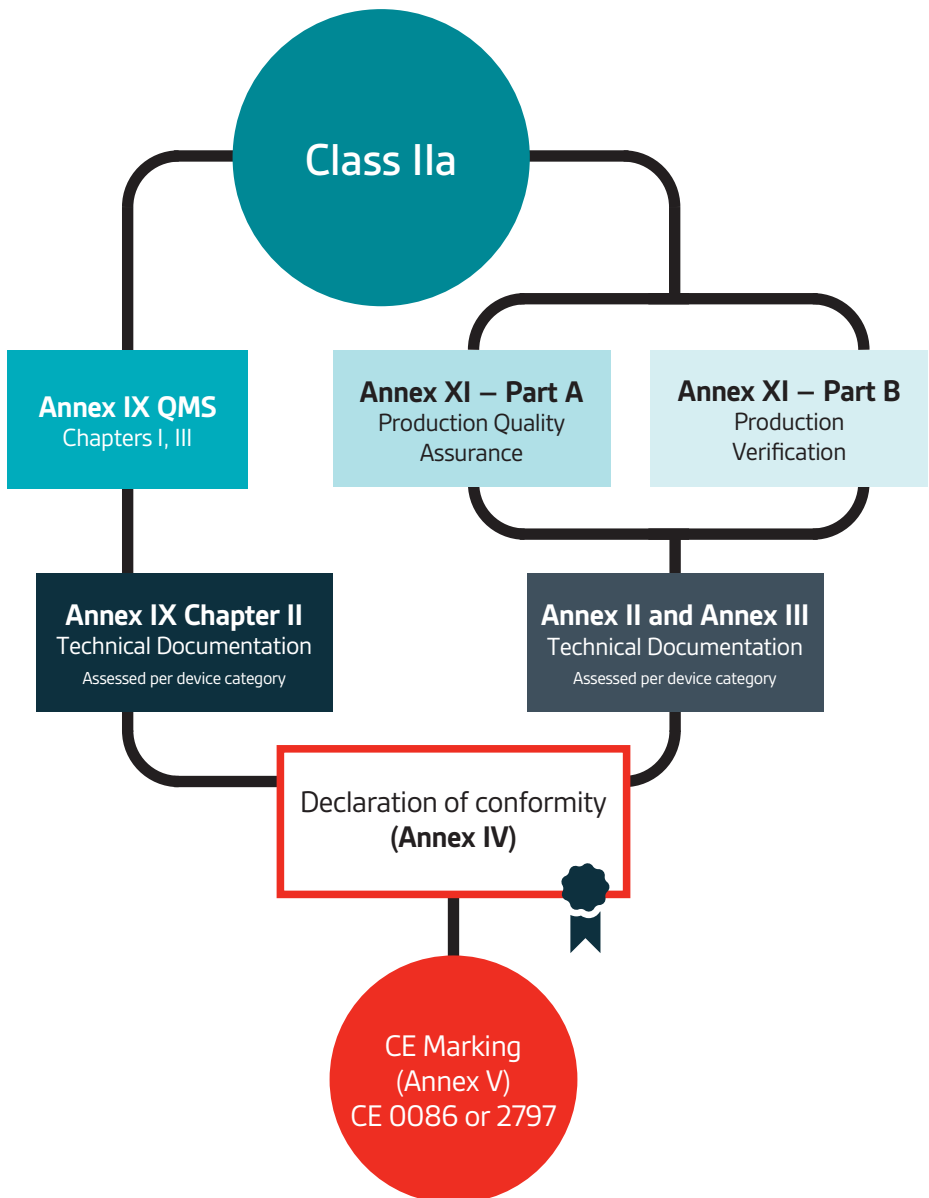
CLASS Is/Im/Ir DEVICES

| Class Is/Im/Ir devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|---|-------------------------------|--------------|------|--------|------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes* | N/A | Yes* | N/A | Yes* | N/A |
| Technical Documentation Assessment | N/A | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | N/A | N/A | N/A | N/A | N/A | N/A |

*if sterile or re-usable surgical instruments

| | | | | | |
|---|--|-----|-----|-----|-----|
| Clinical Evaluation Report updates | Updated as per Manufacturer's clinical evaluation plan | | | | |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated as per Manufacturer's PMS, PMCF plans; NB QMS audits to verify implementation of the plan by sampling complaints, vigilance information etc. | | | | |
| Periodic Safety Update Report (Article 86) | N/A | N/A | N/A | N/A | N/A |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 5 years | | | | |

Class IIa devices



CLASS IIa NON-IMPLANTABLE DEVICES

| Class IIa non-implantable devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|---|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Sample per category of devices | As per the Technical Documentation Sampling Plan | | | | |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | N/A | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Report updates | Updated as per Manufacturer's clinical evaluation plan; NB to review as per Technical Documentation Sampling Plan | | | | | |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated as per Manufacturer's PMS, PMCF plans; NB to review as per Technical Documentation Sampling Plan | | | | | |
| Periodic Safety Update Report (Article 86) | PSUR update required at least once every 2 years; NB to review as per Technical Documentation Sampling Plan | | | | | |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 5 years | | | | | |

Continued on page 6

Class IIa devices continued

CLASS IIa IMPLANTABLE DEVICES

| Class IIa implantable devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|--------------------------------------|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Sample per category of devices | As per the Technical Documentation Sampling Plan | | | | |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | Yes | Updated at least annually "if indicated"; NB to review as per Technical Documentation Sampling Plan or at the time of PSUR assessments | | | | |
| Clinical Evaluation Report updates | | Updated as per Manufacturer's clinical evaluation plan; NB to review updates as per Technical Documentation Sampling Plan or at the time of PSUR assessments | | | | |
| Post Market Clinical Follow-Up Update Report (Article 61) | | Updated at least annually; NB to review as per Technical Documentation Sampling Plan or at the time of PSUR assessments | | | | |
| Periodic Safety Update Report (Article 86) | | Updated when necessary and at least every two years; submitted to NB via EUDAMED for NB review | | | | |
| Unannounced Audits (BSI policy as of Feb 2019) | | At least once every 3 years | | | | |



General Safety and Performance Requirements (Annex I) in the New Medical Device Regulation

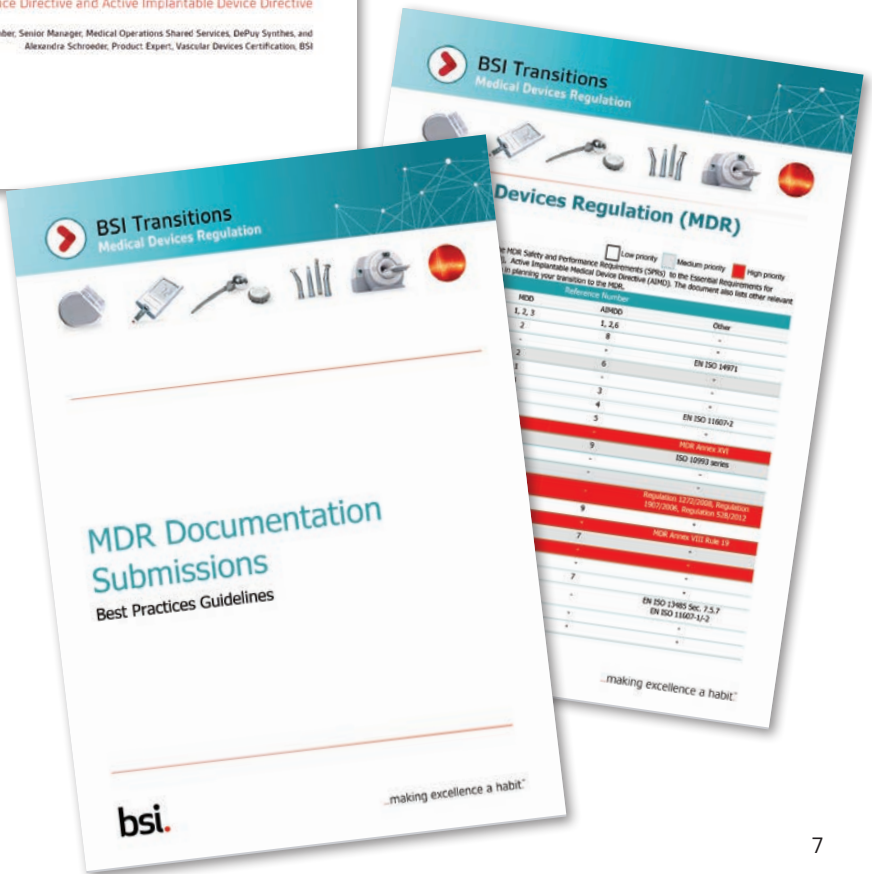
Comparison with the Essential Requirements of the Medical Device Directive and Active Implantable Device Directive

Laurel Macomber, Senior Manager, Medical Operations Shared Services, DePuy Synthes, and Alexandra Schroeder, Product Expert, Vascular Devices Certification, BSI

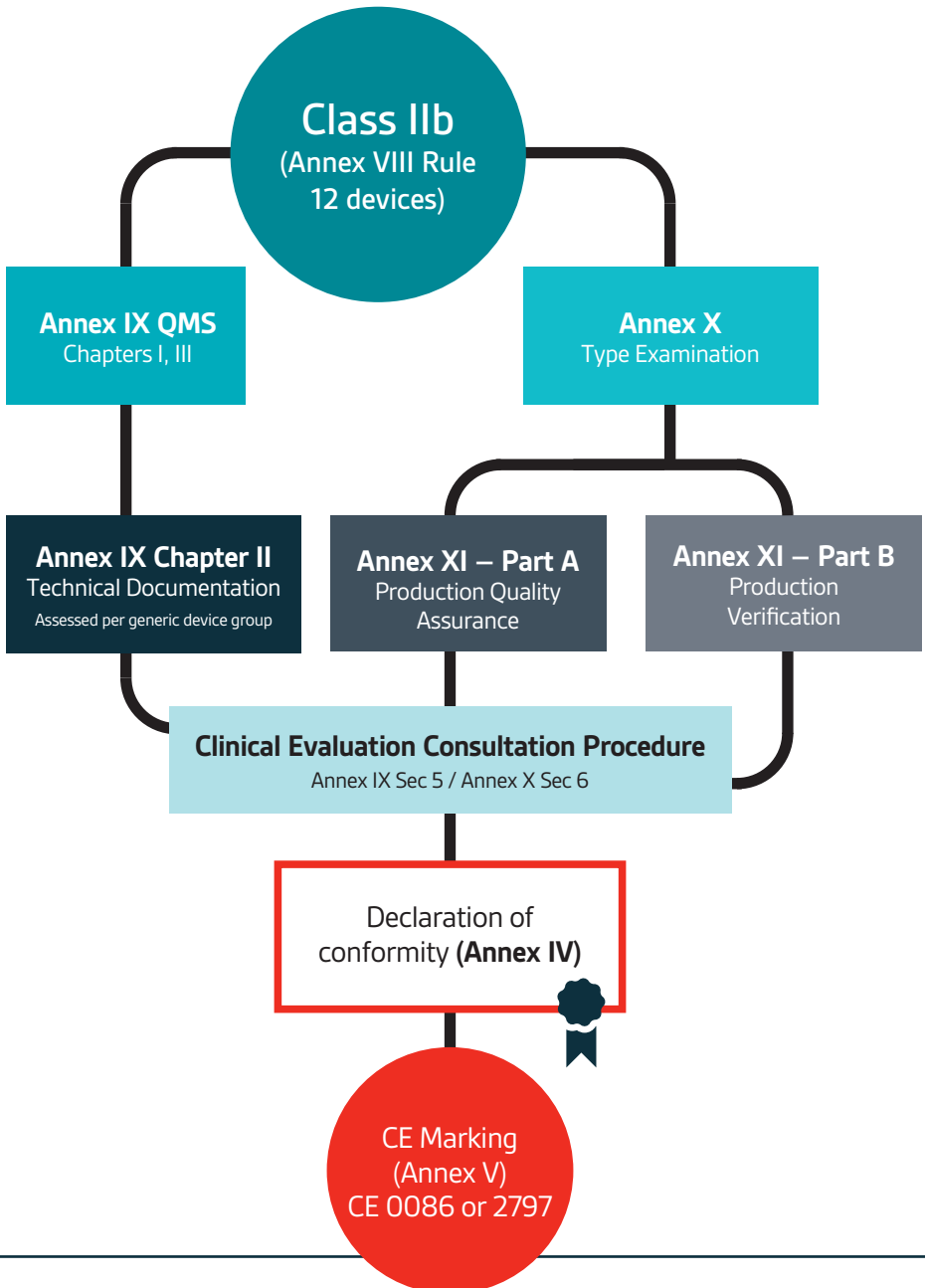
Our website offers a wealth of useful resources including white papers, guidance documents and webinars.

To find out more, visit

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Class IIb Annex VIII Rule 12 devices



Annex VIII Rule 12 devices – All active devices intended to administer and/or remove medicinal products, body liquids or other substances to or from the body.

CLASS IIb ANNEX VIII RULE 12 DEVICES

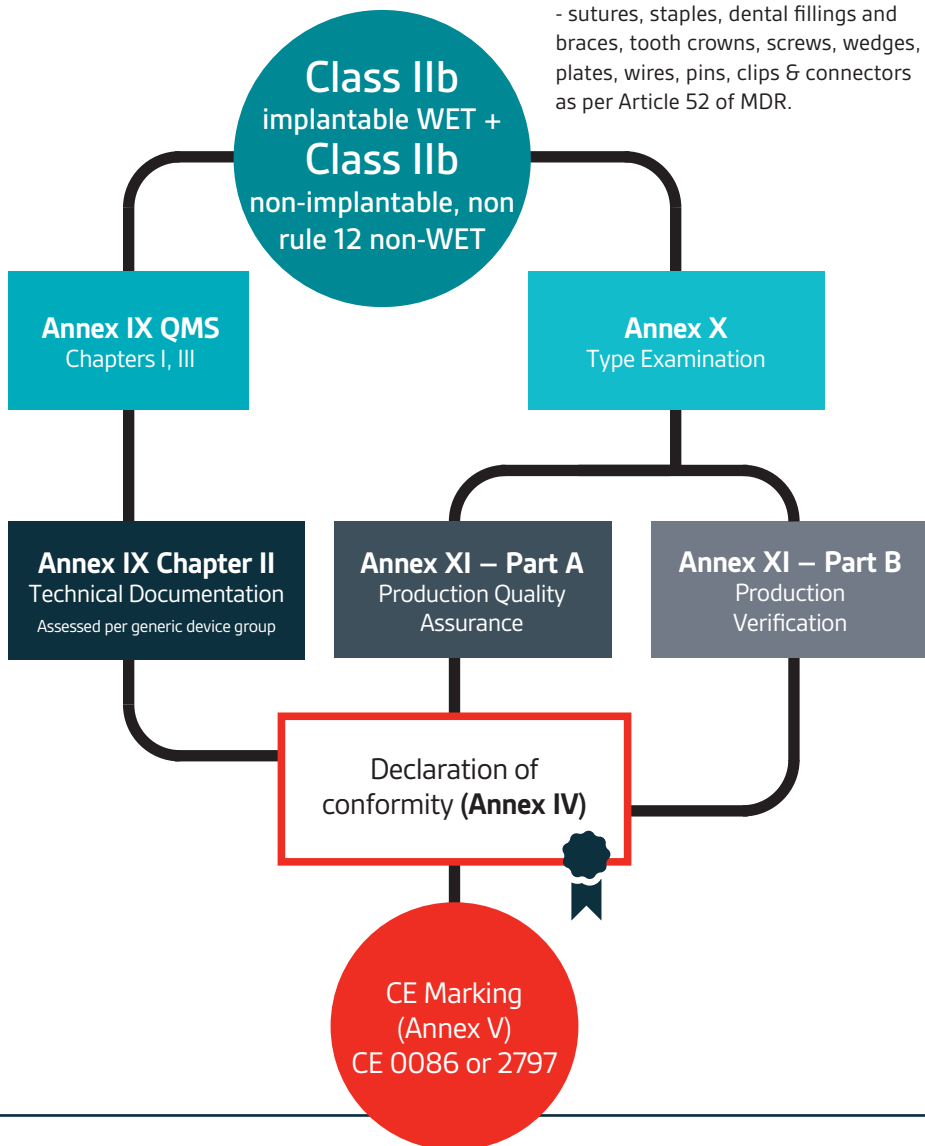
| Class IIb Annex VIII Rule 12 devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|---|---|---|---------------------|---------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert' | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Sample per Generic Device Group | As per the Technical Documentation Sampling Plan | | | | |
| Clinical Evaluation Consultation Procedure (Article 54) | Yes, but exemptions may apply as per Article 54.2 | Maybe required if any modifications to the device adversely affect the risk-benefit ratio | | | | |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | N/A | N/A | N/A | N/A | N/A | N/A |

| | |
|---|---|
| Clinical Evaluation Report Updates | Updated as per Manufacturer's clinical evaluation plan; NB to review updates as per Technical Documentation Sampling Plan |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated as per Manufacturer's PMCF plan; NB to review updates as per Technical Documentation Sampling Plan |
| Periodic Safety Update Report (Article 86) | Updated at least annually; NB to review updates as per Technical Documentation Sampling Plan |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 5 years |

Class IIb implantable WET

Class IIb non-implantable, non rule 12, non WET

Well-Established Technologies (WET)
- sutures, staples, dental fillings and
braces, tooth crowns, screws, wedges,
plates, wires, pins, clips & connectors
as per Article 52 of MDR.



CLASS IIb IMPLANTABLE WET

| Class IIb implantable WET devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|---|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Sample per Generic Device Group | As per the Technical Documentation Sampling Plan | | | | |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | Yes | Updated at least annually "if indicated"; NB to review updates as per Technical Documentation Sampling Plan or at the time of PSUR assessments | | | | |
| Clinical Evaluation Report updates | Updated as per Manufacturer's clinical evaluation plan; NB to review as per Technical Documentation Sampling Plan | | | | | |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated at least annually; NB to review updates as per Technical Documentation Sampling Plan or at the time of PSUR assessments | | | | | |
| Periodic Safety Update Report (Article 86) | Updated at least annually; submitted to NB via EUDAMED for NB review (assuming WET devices are implantable devices) | | | | | |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 3 years | | | | | |

Continued on page 12

CLASS IIb NON-IMPLANTABLE, NON WET, NON RULE 12

| Class IIb non-implantable, non-WET, non-Rule 12 devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|---|---------------------------------------|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Sample per Generic Device Group | As per the Technical Documentation Sampling Plan | | | | |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | N/A | N/A | N/A | N/A | N/A | N/A |

| | |
|--|---|
| Clinical Evaluation Report updates | Updated as per Manufacturer's clinical evaluation plan; NB to review updates as per Technical Documentation Sampling Plan |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated as per Manufacturer's PMCF plan; NB to review updates as per Technical Documentation Sampling Plan |
| Periodic Safety Update Report (Article 86) | Updated at least annually; NB to review updates as per Technical Documentation Sampling Plan |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 5 years |



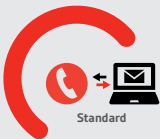
Don't delay...

excellence

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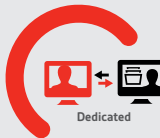
Technical Documentation Reviews for CE Marking

BSI CE-Excellence Programmes are designed for medical device manufacturers wanting to get their products to European markets efficiently and safely.



CE-Standard

The CE-Standard review service allows you to work closely with your assigned BSI Product Expert on your product certification. These reviews are conducted remotely, with communication between you and your BSI Product Expert via phone and email, as required.



CE-Dedicated

The CE-Dedicated review service allows you to book your technical documentation review in advance. The service is conducted remotely with your BSI Product Expert, who uses the time allocated to your company to conduct a focused review of your technical documentation. This allows you to interact with your BSI expert, providing them information during the review. The CE-Dedicated service improves the efficiency of the process, and provides predictability in your planning of the review.

Note: Our services do not guarantee a CE Marking certificate will be issued within a certain amount of working days, but are based on completing the review process with either a positive or negative recommendation. CE-Dedicated is not available for devices utilizing animal tissue, blood derivatives or medicinal substances.

For more information on our CE-Excellence services call BSI on
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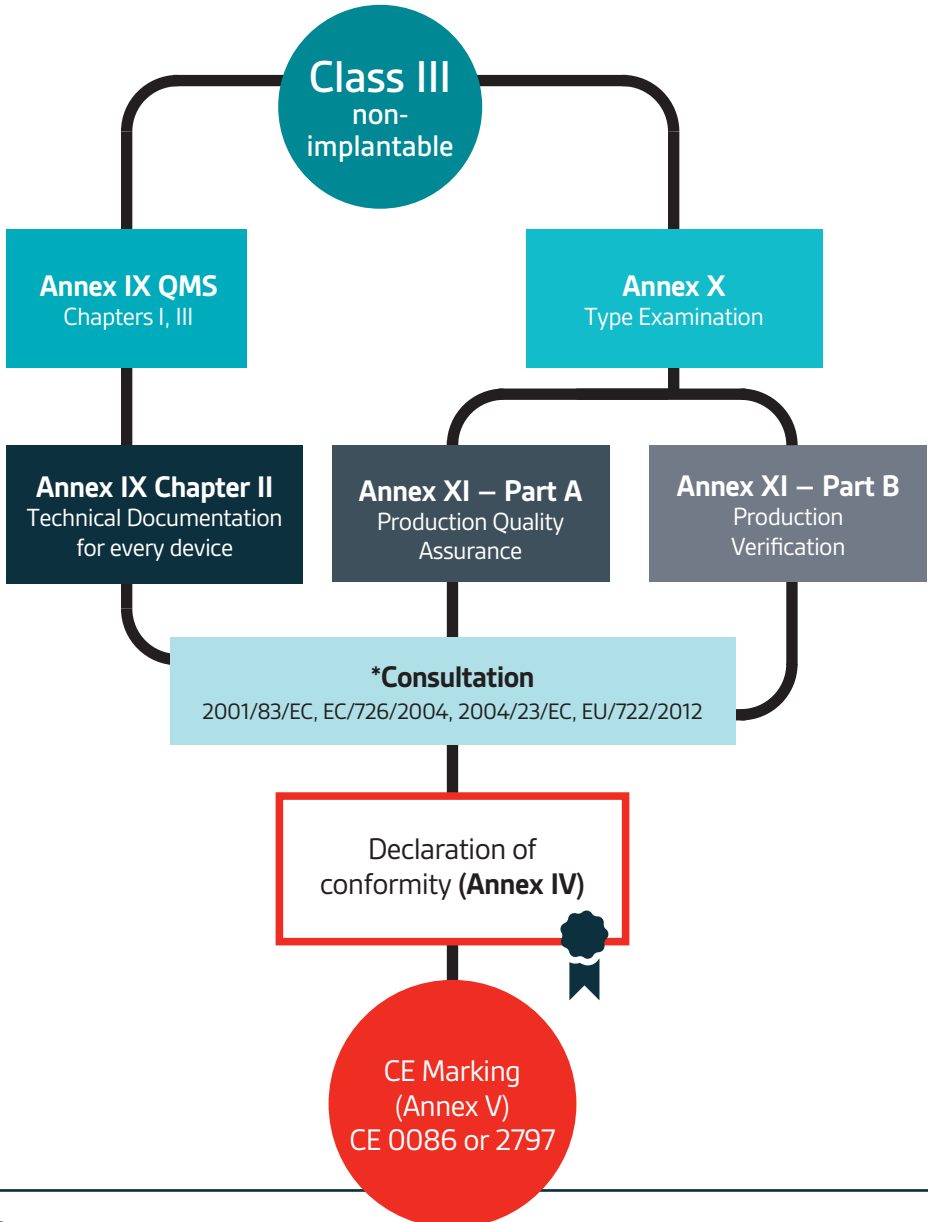
CLASS IIb IMPLANTABLE NON-WET DEVICES

| Class IIb implantable non-WET devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|-------------------------------------|---|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Review for every device | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | Yes | Updated at least annually "if indicated"; NB to review at the time of PSUR reviews or substantial change reviews | | | | |

| | |
|--|--|
| Clinical Evaluation Report updates | Updated as per Manufacturer's clinical evaluation plan; NB to review at the time of PSUR reviews or substantial change reviews |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated at least annually; NB to review at the time of PSUR reviews or substantial change reviews |
| Periodic Safety Update Report (Article 86) | Updated at least annually; submitted to NB via EUDAMED for NB review |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 3 years |

Class III non-implantable devices

(including devices with medicinal substances, human tissue or animal tissue derivatives with TSE risk, Class III Rule 21 devices).



CLASS III NON-IMPLANTABLE DEVICES

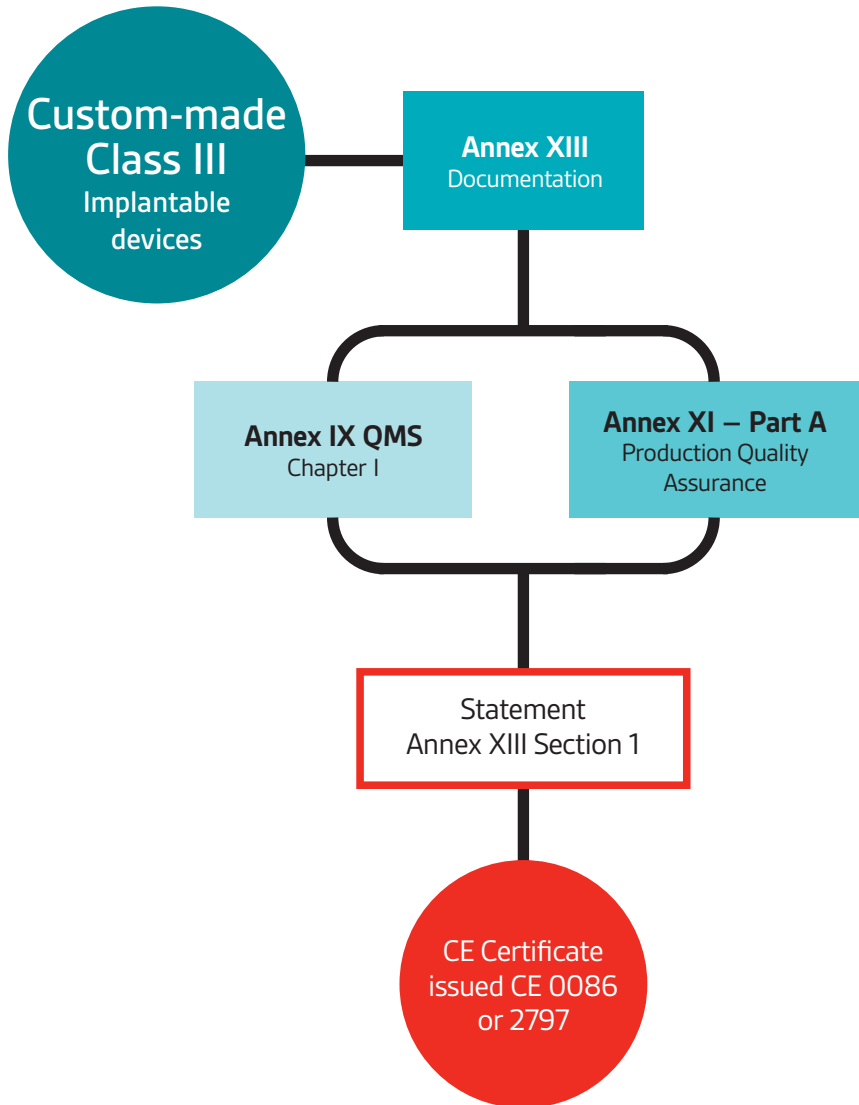
| Class III non-implantable devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|-------------------------------------|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Review for every device | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | If applicable | Modifications to the devices may need supplementary consultations; determined on a case-by-case basis taking into account the nature of the changes proposed | | | | |
| Summary of Safety and Clinical Performance (Article 32) | Yes | Updated at least annually "if indicated"; NB to review at the time of PSUR reviews or substantial change reviews | | | | |

| | |
|---|--|
| Clinical Evaluation Report Updates | Updated as per Manufacturer's clinical evaluation plan; NB to review at the time of PSUR reviews or substantial change reviews |
| Post Market Clinical Follow-Up Update Report (Article 61) | Updated at least annually; NB to review at the time of PSUR reviews or substantial change reviews |
| Periodic Safety Update Report (Article 86) | Updated at least annually; submitted to NB via EUDAMED for NB review |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 3 years |

CLASS III IMPLANTABLE DEVICES

| Class III implantable devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|--|--|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | Review for every device | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Consultation Procedure (Article 54) | Yes, but exemptions may apply as per Article 54.2 | May be required if any modifications to the device adversely affect the risk-benefit ratio | | | | |
| Consultations (Rule 14, Rule 18, Rule 21) | If applicable | Modifications to the devices may need supplementary consultations; determined on a case-by-case basis taking into account the nature of the changes proposed | | | | |
| Summary of Safety and Clinical Performance (Article 32) | Yes | Updated at least annually "if indicated"; NB to review at the time of PSUR assessments or substantial change reviews | | | | |
| Clinical Evaluation Report Updates | | Updated as per Manufacturer's clinical evaluation plan; NB to review at the time of PSUR reviews or substantial change reviews | | | | |
| Post Market Clinical Follow-Up Update Report (Article 61) | | Updated at least annually; NB review at the time of PSUR reviews or substantial change reviews | | | | |
| Periodic Safety Update Report (Article 86) | | Updated at least annually; submitted to NB via EUDAMED for NB review | | | | |
| Unannounced Audits (BSI policy as of Feb 2019) | | At least once every 3 years | | | | |

Custom-made Class III implantable devices



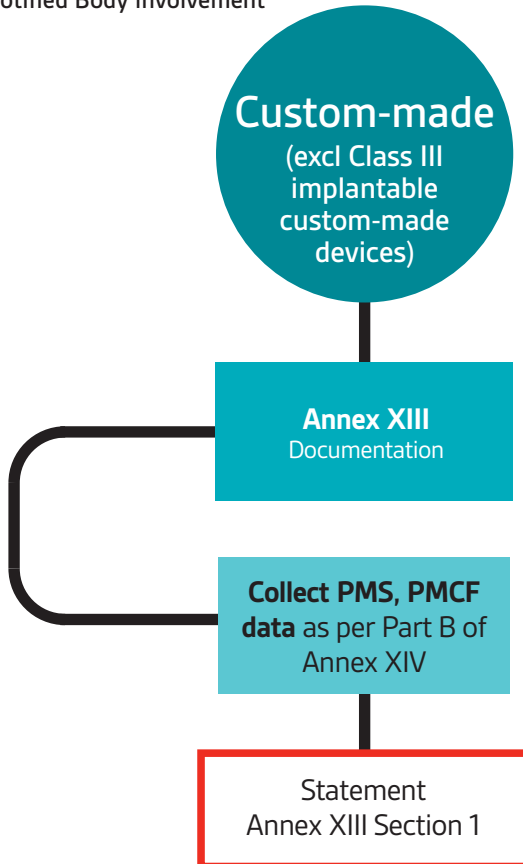
CUSTOM-MADE CLASS III IMPLANTABLE DEVICES

| Custom-made Class III implantable devices | Initial Conformity Assessment | SURVEILLANCE | | | | |
|--|---|--------------|---------------------|--------|---------------------|-----|
| | | Y1 | Y2 | Y3 | Y4 | Y5 |
| QMS Audits | Yes | Yes | Yes | Recert | Yes | Yes |
| Microbiology Audits | Yes (if sterile) | N/A | Yes (if sterile) | N/A | Yes (if sterile) | N/A |
| Technical Documentation Assessment | N/A | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Consultation Procedure (Article 54) | N/A | N/A | N/A | N/A | N/A | N/A |
| Consultations (Rule 14, Rule 18, Rule 21) | N/A | N/A | N/A | N/A | N/A | N/A |
| Summary of Safety and Clinical Performance (Article 32) | N/A | N/A | N/A | N/A | N/A | N/A |
| Clinical Evaluation Report Updates | | N/A | N/A | N/A | N/A | N/A |
| Post Market Clinical Follow-Up Update Report (Article 61) | As per Manufacturer's PMS, PMCF plans; NB QMS audits to verify implementation of the plan | | | | | |
| Periodic Safety Update Report (Article 86) | Updated at least annually; unclear whether to be submitted to EUDAMED for NB review or not; NB to verify updates at the time of surveillance QMS audits | | | | | |
| Unannounced Audits (BSI policy as of Feb 2019) | At least once every 3 years | | | | | |

Custom-made devices

(excluding custom-made Class III implantable devices)

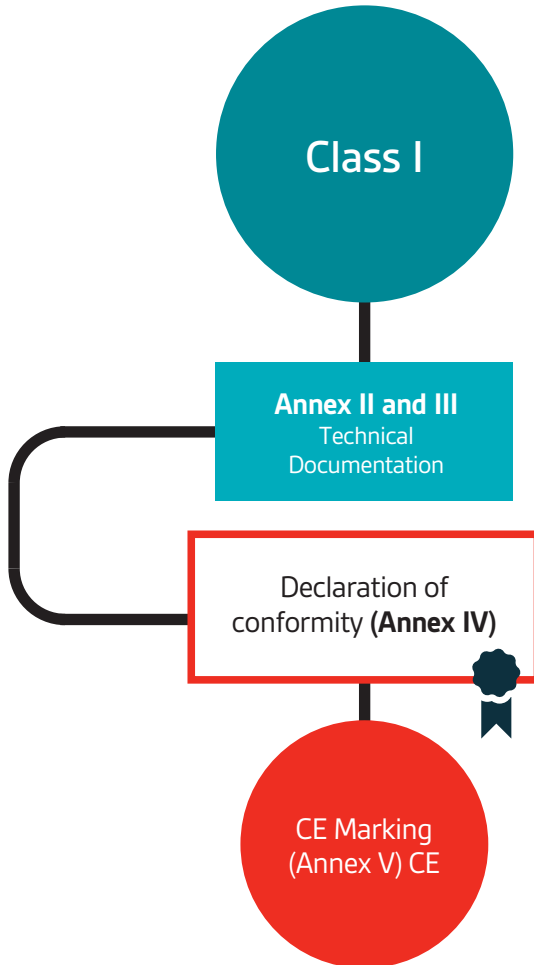
Note: No Notified Body involvement



Class I devices

(excluding Class Is/Im/Ir devices)

Note: No Notified Body involvement



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