



SLOVENSKI STANDARD
SIST EN ISO 13485:2016/oprA1:2019
01-december-2019

Medicinski pripomočki - Sistemi vodenja kakovosti - Zahteve za zakonodajne namene (ISO 13485:2016)

Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016)

Medizinprodukte - Qualitätsmanagementsysteme - Anforderungen für regulatorische Zwecke (ISO 13485:2016)

Dispositifs médicaux - Systèmes de management de la qualité - Exigences à des fins réglementaires (ISO 13485:2016)

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Ta slovenski standard je istoveten z: EN ISO 13485:2016/prA1

ICS:

03.100.70	Sistemi vodenja	Management systems
11.020.01	Vodenje kakovosti in ravnanje z okoljem v zdravstvu	Quality and environmental management in health care

SIST EN ISO 13485:2016/oprA1:2019 **en**

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EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

DRAFT
EN ISO 13485:2016
prA1

October 2019

ICS 03.100.70; 11.040.01

English version

Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016)

Dispositifs médicaux - Systèmes de management de la
qualité - Exigences à des fins réglementaires (ISO
13485:2016)

Medizinprodukte - Qualitätsmanagementsysteme -
Anforderungen für regulatorische Zwecke (ISO
13485:2016)

This draft amendment is submitted to CEN members for enquiry. It has been drawn up by the Technical Committee CEN/CLC/JTC 3.

This draft amendment A1, if approved, will modify the European Standard EN ISO 13485:2016. If this draft becomes an amendment, CEN and CENELEC members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for inclusion of this amendment into the relevant national standard without any alteration.

This draft amendment was established by CEN and CENELEC in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN and CENELEC member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

CEN and CENELEC members are the national standards bodies and national electrotechnical committees of Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and United Kingdom.

Recipients of this draft are invited to submit, with their comments, notification of any relevant patent rights of which they are aware and to provide supporting documentation. Recipients of this draft are invited to submit, with their comments, notification of any relevant patent rights of which they are aware and to provide supporting documentation.

Warning : This document is not a European Standard. It is distributed for review and comments. It is subject to change without notice and shall not be referred to as a European Standard.



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European foreword

The text of ISO 13485:2016 has been prepared by Technical Committee ISO/TC 210 “Quality management and corresponding general aspects for medical devices” of the International Organization for Standardization (ISO) and has been taken over as EN ISO 13485:2016/prA1:2019 by Technical Committee CEN/CLC/TC 3 “Quality management and corresponding general aspects for medical devices” the secretariat of which is held by NEN.

This document is currently submitted to the CEN Enquiry.

This document supersedes EN ISO 13485:2016, incorporating corrigenda March 2016, December 2016 and 2018, with a revised European Foreword and European Annexes ZA, ZB and ZC, and additional European Annexes ZD and ZE.

This document has been prepared under a mandate given to CEN by the European Commission and the European Free Trade Association, and supports requirements of EU Directive(s) and Regulation(s).

For relationship with EU Directive(s) and Regulation(s), see informative Annex ZA, ZB, ZC, ZD and ZE, which are an integral part of this document.

The following referenced documents are indispensable for the application of this document. For undated references, the edition of the referenced document (including any amendments) listed below applies. For dated references, only the edition cited applies. However for any use of this standard within the meaning of Annex ZA, ZB, ZC, ZD or ZE, the user should always check that any referenced document has not been superseded and that its relevant contents can still be considered the generally acknowledged state-of-art.

When an IEC or ISO standard is referred to in the ISO standard text, this should be understood as a normative reference to the corresponding EN standard, if available, and otherwise to the dated version of the ISO or IEC standard as listed below.

NOTE The way in which these referenced documents are cited in normative requirements determines the extent (in whole or in part) to which they apply.

Table – Correlation between normative references and dated EN and ISO standards

Normative references as listed in Clause 2 of the ISO standard	Equivalent dated standard	
	EN	ISO
ISO 9000:2015	EN ISO 9000:2015	ISO 9000:2015

Endorsement notice

The text of ISO 13485:2016 has been approved by CEN as EN ISO 13485:2016/prA1:2019 without any modification.

Annex ZA (informative)

Relationship between this European Standard and the conformity assessment requirements of EU Directive 90/385/EEC on active implantable medical devices [OJ L 189] (as amended) aimed to be covered

This European standard has been prepared under a Commission's standardisation request M/023 to provide one voluntary means of conforming to essential requirements of Council Directive 90/385/EEC of 20 June 1990 on the approximation of the laws of the Member States relating to active implantable medical devices [OJ L 189] (as amended).

Once this standard is cited in the Official Journal of the European Union under that Directive, compliance with the normative clauses of this standard given in Tables ZA.1 or ZA.2 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding conformity assessment requirements of that Directive and associated EFTA regulations.

The Conformity Assessment Annexes 2 and 5 of the Directive include description of the regulatory process and activities undertaken by the Notified Body, which both are outside of the scope of this European Standard and therefore not covered by this European Standard. Furthermore, the requirements of the Directive refer to an application to a Notified Body not to the requirement for a quality system as such. Accordingly, coverage of legal requirements can only be presumed to the extent listed in Tables ZA.1 and ZA.2 if an application to a Notified Body:

- contains the necessary quality system documentation;
- has been reviewed and approved by a Notified Body,

and the undertakings listed in the application are correctly executed by the manufacturer.

NOTE 1 Where a reference from a clause of this standard to the risk management process is made, the risk management process needs to be in compliance with 90/385/EEC, as amended by 2007/47/EC. This means that risks have to be reduced 'as far as possible', 'to a minimum', 'to the lowest possible level', 'minimized' or 'removed', according to the wording of the corresponding essential requirement.

NOTE 2 The manufacturer's policy for determining **acceptable risk** must be in compliance with essential requirements 1, 4, 5, 8, 9 and 10 of the Directive.

NOTE 3 This Annex ZA is based on normative references according to the table of references in the European foreword, replacing the references in the core text.

NOTE 4 When a requirement does not appear in Tables ZA.1 or ZA.2, it means that it is not addressed by this European Standard.

NOTE 5 This annex uses the term "quality system" as used in the Directive whereas this European Standard uses the term "quality management system" in accordance with ISO terminology.

ZA.1 Relationship with Annex 2 of Directive 90/385/EEC (as amended)

Compliance with this European Standard does not provide presumption of conformity with all the aspects of Annex 2, as outlined in Table ZA.1. Therefore, a manufacturer or a Notified Body has to take additional provisions to ensure conformity, and claim or certify conformance, with Annex 2 of this Directive. The legal requirements must be examined, applied and verified one by one and the solutions adopted must become part of the quality system in the meaning of the Directive.

Table ZA.1 — Correspondence between this European Standard and Annex 2 of Directive 90/385/EEC [OJ L 189] (as amended)

Paragraph of Directive 90/385/EEC, Annex 2	Clauses of this EN	Qualifying remarks/Notes
3.1, 1 st sentence		Not covered.
3.1, 2 nd sentence, 1 st indent		Not covered.
3.1, 2 nd sentence, 2 nd indent	4.1.1, 4.1.2, 4.1.3, 4.1.4, 4.1.6, 4.2.1, 4.2.2, 4.2.3, 4.2.4, 4.2.5	Covered. The documentation required in this European Standard covers the quality system documentation meant in 3.2 of Annex 2 when the explicit legal requirements are incorporated into the quality system documentation. See also coverage of 3.2 below.
3.1, 2 nd sentence, 3 rd indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered in part. This European Standard requires top management commitment to implementation of the quality system and that documented procedures are implemented but does not require a signed undertaking.
3.1, 2 nd sentence, 4 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered in part. This European Standard requires maintenance of the approved quality system but does not require a signed undertaking.
3.1, 2 nd sentence, 5 th indent		Not covered. This European Standard includes requirements on post-market surveillance, and reporting adverse events and field safety corrective actions to authorities but does not cover all the details required by the Directive including timescales for reporting.
3.2, 1 st paragraph		Not covered. The application of this European Standard does not by itself ensure the fulfilment of all regulatory requirements of the Directive. The legal requirements must be examined, applied and verified one by one and the solutions adopted become part of the quality system in the meaning of the Directive.
3.2, 2 nd paragraph, 1 st sentence	4.1, 4.2	Covered.

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Paragraph of Directive 90/385/EEC, Annex 2	Clauses of this EN	Qualifying remarks/Notes
3.2, 2 nd paragraph, 2 nd sentence	4.1, 4.2	Covered.
3.2, 2 nd paragraph, 3 rd sentence	4.1, 4.2, 7	Covered provided quality management system documentation makes possible a uniform interpretation of the quality policies and procedures, such as quality programs, quality plans, quality manuals and quality records, and that the applicable documentation listed in 3.2 of Annex 2 is incorporated into the quality system documentation.
3.2, 3 rd paragraph (a)	4.2.1, 4.2.3, 5.1, 5.3, 5.4.1	Covered.
3.2, 3 rd paragraph (b)	4.2.2, 5.1	Covered.
3.2, 3 rd paragraph (b), 1 st indent	4.2.2, 5.1, 5.5.1, 5.5.2	Covered.
3.2, 3 rd paragraph (b), 2 nd indent	4.1, 5.6, 7.1, 8.2.4, 8.3, 8.4, 8.5.2, 8.5.3	Covered provided that the methods and acceptance criteria chosen by the manufacturer ensure that the requirements of the Directive are fulfilled.
3.2 3 rd paragraph (b) 3 rd indent	4.1, 4.2, 7.4, 8.2.2	Covered.
3.2 3 rd paragraph (c) 1 st indent	4.2, 7.3.2, 7.3.3, 7.3.7, 7.3.9, 7.3.10	Covered provided that the applicable quality management system documentation includes design specifications identifying standards which will be applied and a description of the solutions adopted to fulfil the essential requirements which apply when harmonized standards are not applied in full.
3.2, 3 rd paragraph (c), 2 nd indent	7.3.1, 7.3.6, 7.3.7, 7.3.9	Covered.
3.2, 3 rd paragraph (c), 3 rd indent		Not covered.
3.2, 3 rd paragraph (c), 4 th indent	7.3.6, 7.3.7	Covered provided that the quality management system records include the pre-clinical evaluation.
3.2, 3 rd paragraph (c), 5 th indent		Not covered. Clause 7.3.7 does not include the details of Annex 7.
3.2, 3 rd paragraph (d), 1 st indent	4.2, 6.4, 7.1, 7.4 7.5	Covered provided that the quality management system documentation includes relevant documents and records in regards to sterilization and purchasing.
3.2, 3 rd paragraph (d), 2 nd indent	4.2, 7.5.8, 7.5.9	Covered.
3.2, 3 rd paragraph (e)	4.2, 7.1, 7.4.3, 7.5.9.1, 7.6, 8.2.6	Covered provided that the documented frequency at which tests are carried out is detailed in the quality management system documentation.
6.1		Not covered. The specific time periods in Directive are not specified in 4.2.4 or 4.2.5.

ZA.2 Relationship with Annex 5 of Directive 90/385/EEC (as amended)

Compliance with this European Standard does not provide presumption of conformity with all the aspects of Annex 5, as outlined in Table ZA.2. Therefore, a manufacturer or a Notified Body has to take additional provisions to ensure conformity, and claim or certify conformance, with Annex 5 of this Directive. The legal requirements must be examined, applied and verified one by one and the solutions adopted must become part of the quality system in the meaning of the Directive.

Table ZA.2 — Correspondence between this European Standard and Annex 5 of Directive 90/385/EEC (as amended)

Paragraph of Directive 90/385/EEC, Annex 5	Clauses of this EN	Qualifying remarks/Notes
3.1, 1 st paragraph		Not covered.
3.1, 2 nd paragraph, 1 st indent		Not covered.
3.1, 2 nd paragraph, 2 nd indent	4.1, 4.2	Covered. The documentation required in this European Standard covers the quality system documentation meant in 3.2 of Annex 5 when the explicit legal requirements are incorporated into the quality system documentation. See also coverage of 3.2 below.
3.1, 2 nd paragraph, 3 rd indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1, 2 nd paragraph, 4 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1, 2 nd paragraph, 5 th indent	4.1, 4.2	Covered in part provided that quality management system includes the technical documentation relating to the applicable approved type(s) of medical device(s). Reference to the EC type-examination certificate is not covered.
3.1, 2 nd paragraph, 6 th indent		Not covered. This European Standard includes requirements on post market surveillance, and reporting adverse events and field safety corrective actions to authorities but does not cover all the details required by the Directive including timescales for reporting
3.2, 1 st paragraph		Not covered. Reference to the EC type-examination certificate is not covered.
3.2, 2 nd paragraph	4.1, 4.2	Covered.
3.2, 3 rd paragraph (a)	4.2.1, 5.1, 5.3, 5.4.1	Covered.
3.2, 3 rd paragraph (b), 1 st indent	5.5.1, 5.5.2	Covered.

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Paragraph of Directive 90/385/EEC, Annex 5	Clauses of this EN	Qualifying remarks/Notes
3.2, 3 rd paragraph (b), 2 nd indent	4.1, 5.6, 7.1, 8.2.4, 8.3, 8.4, 8.5.2, 8.5.3	Covered provided that the methods and acceptance criteria chosen by the manufacturer ensure that the requirements of the Directive are fulfilled.
3.2, 3 rd paragraph (b), 3 rd indent	4.1, 4.2, 7.4, 8.2.2	Covered.
3.2, 3 rd paragraph (c), 1 st indent	4.2, 6.4, 7.1, 7.4, 7.5, 8.2.5	Covered provided that the quality management system documentation includes relevant documents and records in regards to sterilization and purchasing.
3.2, 3 rd paragraph (c), 2 nd indent	4.2, 7.5.8, 7.5.9	Covered.
3.2, 3 rd paragraph (d)	7.1, 7.4.3, 7.6, 8.2.6	Covered provided that the frequency at which tests are carried out is documented in the quality management system documentation.

WARNING 1 — Presumption of conformity stays valid only as long as a reference to this European Standard is maintained in the list published in the Official Journal of the European Union. Users of this standard should consult frequently the latest list published in the Official Journal of the European Union.

WARNING 2 — Other Union legislation may be applicable to the products falling within the scope of this standard.

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Annex ZB (informative)

Relationship between this European Standard and the conformity assessment requirements of EU Directive 93/42/EEC on medical devices [OJ L 169] (as amended) aimed to be covered

This European Standard has been prepared under a Commission's standardization request M/023 to provide one voluntary means of conforming to essential requirements of Council Directive 93/42/EEC of 14 June 1993 concerning medical devices [OJ L 169 (as amended)].

Once this standard is cited in the Official Journal of the European Union under that Directive, compliance with the normative clauses of this standard given in Tables ZB.1, ZB.2 or ZB.3 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding conformity assessment requirements of that Directive and associated EFTA regulations.

The Conformity Assessment Annexes II, V and VI of the Directive include description of the regulatory process and activities undertaken by the Notified Body, which both are outside of the scope of this European Standard and therefore not covered by this European Standard. Furthermore, the requirements of the Directive refer to an application to a Notified Body, not to the requirement for a quality system as such. Accordingly, coverage of legal requirements can only be presumed to the extent listed in Tables ZB.1, ZB.2 and ZB.3 if an application to a Notified Body:

- contains the necessary quality system documentation;
- has been reviewed and approved by a Notified Body,

and the undertakings listed in the application are correctly executed by the manufacturer.

NOTE 1 Where a reference from a clause of this standard to the risk management process is made, the risk management process needs to be in compliance with 93/42/EEC, as amended by 2007/47/EC. This means that risks have to be reduced 'as far as possible', 'to a minimum', 'to the lowest possible level', 'minimized' or 'removed', according to the wording of the corresponding essential requirement.

NOTE 2 The manufacturer's policy for determining **acceptable risk** must be in compliance with essential requirements 1, 2, 5, 6, 7, 8, 9, 11 and 12 of the Directive.

NOTE 3 This Annex ZB is based on normative references according to the table of references in the European foreword, replacing the references in the core text.

NOTE 4 When a requirement does not appear in Tables ZB.1, ZB.2 or ZB.3 it means that it is not addressed by this European Standard.

NOTE 5 This annex uses the term "quality system" as used in the Directive whereas this European Standard uses the term "quality management system" in accordance with ISO terminology.

ZB.1 Relationship with Annex II of Directive 93/42/EEC (as amended)

Compliance with this European Standard does not provide a presumption of conformity with all the aspects of Annex II, as outlined in Table ZB.1. Therefore, a manufacturer or a Notified Body has to take additional provisions to ensure conformity, and claim or certify conformance, with Annex II of this Directive. The legal requirements must be examined, applied and verified one by one and the solutions

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adopted must become part of the quality system in the meaning of the Directive.

Table ZB.1 — Correspondence between this European Standard and Annex II of Directive 93/42/EEC [OJ L 169] (as amended)

Paragraph of Directive 93/42/EEC, Annex II	Clauses of this EN	Qualifying remarks/Notes
3.1, 1st sentence		Not covered.
3.1, 2 nd sentence, 1 st indent		Not covered.
3.1, 2 nd sentence, 2 nd indent		Not covered.
3.1, 2 nd sentence, 3 rd indent		Not covered.
3.1, 2 nd sentence, 4 th indent	4.1, 4.2	Covered. The documentation required in this European Standard covers the quality system documentation meant in 3.2 of Annex II when the explicit legal requirements are incorporated into the quality system documentation. See also coverage of 3.2 below.
3.1, 2 nd sentence, 5 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1, 2 nd sentence, 6 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1, 2 nd sentence, 7 th indent 3.1, 7 th indent (i) 3.1, 7 th indent (ii)		Not covered. This European Standard includes requirements on post market surveillance, and reporting adverse events and field safety corrective actions to authorities but does not cover all the details required by the Directive including timescales for reporting.
3.2, 1 st paragraph, 1 st sentence		Not covered. The application of this European Standard does not by itself ensure the fulfilment of all regulatory requirements of the Directive. The legal requirements must be examined, applied and verified one by one and the solutions adopted become part of the quality system in the meaning of the Directive.
3.2, 1 st paragraph, 2 nd sentence	4.1, 4.2, 7.1.	Covered.
3.2, 2 nd paragraph	4.1, 4.2, 7	Covered provided quality management system documentation makes possible a uniform interpretation of the quality policies and procedures, such as quality programs, quality plans, quality manuals and quality records, and that the applicable documentation listed in 3.2 of Annex II is incorporated into the quality system documentation.
3.2, 3 rd paragraph (a)	4.2.1, 5.1, 5.3, 5.4.1	Covered.
3.2, 3 rd paragraph (b)	4.2.2, 5.1	Covered.
3.2, 3 rd paragraph (b), 1 st indent	4.2.2, 5.1, 5.5.1, 5.5.2	Covered.

Paragraph of Directive 93/42/EEC, Annex II	Clauses of this EN	Qualifying remarks/Notes
3.2, 3 rd paragraph (b), 2 nd indent	4.1, 5.6, 7.1, 8.2.4, 8.3, 8.4, 8.5.2, 8.5.3	Covered provided that the methods and acceptance criteria chosen by the manufacturer ensure that the requirements of the Directive are fulfilled.
3.2, 3 rd paragraph (b), 3 rd indent	4.1, 4.2, 7.4, 8.2.2	Covered.
3.2, 3 rd paragraph (c)	7.1, 7.2, 7.3	Covered.
3.2, 3 rd paragraph (c), 1 st indent	4.2.3, 7.2, 7.3.3, 7.3.4, 7.3.10	Covered provided that the documentation containing a general description of the medical device includes any variants.
3.2, 3 rd paragraph (c), 2 nd indent	4.2, 7.3.3, 7.3.4, 7.3.6, 7.3.8	Covered provided that the applicable quality management system documentation includes design specifications identifying standards which will be applied and a description of the solutions adopted to fulfil the essential requirements which apply when harmonized standards are not applied in full.
3.2, 3 rd paragraph (c), 3 rd indent	7.3.1, 7.3.6, 7.3.7, 7.3.8, 7.3.9, 7.3.10	Covered.
3.2, 3 rd paragraph (c), 4 th indent	7.3.2, 7.3.3, 7.3.5, 7.3.6	Covered.
3.2, 3 rd paragraph (c), 5 th indent	4.2.3	Covered provided that the quality management system documentation includes a statement indicating whether or not the medical device incorporates, as an integral part, a substance or a human blood derivative and the data on the tests conducted in this connection required to assess the safety, quality and usefulness of that substance or human blood derivative, taking account of the intended purpose of the medical device.
3.2, 3 rd paragraph (c), 6 th indent	4.2.3	Covered provided that the quality management system documentation includes a statement indicating whether or not the device is manufactured utilizing tissues of animal origin as referred to in Commission Directive 2003/32/EC.
3.2, 3 rd paragraph (c), 7 th indent		Not covered.
3.2, 3 rd paragraph (c), 8 th indent	7.3.6, 7.3.7	Covered provided that the quality management system records include the pre-clinical evaluation.
3.2, 3 rd paragraph (c), 9 th indent		Not covered. 7.3.7 does not include the details of Annex X.

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Paragraph of Directive 93/42/EEC, Annex II	Clauses of this EN	Qualifying remarks/Notes
3.2, 3 rd paragraph (c), 10 th indent	4.1, 4.2, 7	Covered provided that the quality management system documentation includes the label and, where appropriate, instructions for use.
3.2, 3 rd paragraph (d)	4.2, 7.1, 7.5, 7.6, 8.1, 8.2.5, 8.2.6	Covered.
3.2, 3 rd paragraph (d), 1 st indent, sterilization	4.1.1, 6.4, 7.5	Covered.
3.2, 3 rd paragraph (d), 1 st indent, purchasing	4.1.1, 7.4	Covered.
3.2, 3 rd paragraph (d), 1 st indent, relevant documents	4.2, 7.1	Covered provided that the quality management system documentation includes relevant documents and records in regards to sterilization and purchasing.
3.2, 3 rd paragraph (d), 2 nd indent	4.2, 7.5.8, 7.5.9	Covered.
3.2, 3 rd paragraph (e)	4.2, 7.1, 7.4.3, 7.5.9.1, 7.6, 8.2.6	Covered provided that the documented frequency at which tests are carried out is detailed in the quality management system documentation.
6.1		Not covered. The specific time periods in Directive are not Specified in 4.2.4 and 4.2.5

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ZB.2 Relationship with Annex V of Directive 93/42/EEC (as amended)

Compliance with this European Standard does not provide presumption of conformity with all the aspects of Annex V, as outlined in Table ZB.2. Therefore, a manufacturer or a Notified Body has to take additional provisions to ensure conformity, and claim or certify conformance, with Annex V of this Directive. The legal requirements must be examined, applied and verified one by one and the solutions adopted must become part of the quality system in the meaning of the Directive.

Table ZB.2 — Correspondence between this European Standard and Annex V of Directive 93/42/EEC (as amended)

Paragraph of Directive 93/42/EEC, Annex V	Clauses of this EN	Qualifying remarks/Notes
3.1 1 st paragraph		Not covered.
3.1 2 nd paragraph 1 st indent		Not covered.
3.1 2 nd paragraph 2 nd indent		Not covered.
3.1 2 nd paragraph 3 rd indent		Not covered.

Paragraph of Directive 93/42/EEC, Annex V	Clauses of this EN	Qualifying remarks/Notes
3.1 2 nd paragraph 4 th indent	4.1, 4.2	Covered provided quality management system documentation makes possible a uniform interpretation of the quality policies and procedures, such as quality programs, quality plans, quality manuals and quality records, and that the applicable documentation listed in 3.2 of Annex V is incorporated into the quality system documentation.
3.1 2 nd paragraph 5 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1 2 nd paragraph 6 th indent	4.1, 5.1, 5.4, 5.5, 5.6	Covered.
3.1 2 nd paragraph 7 th indent	4.1, 4.2	Covered in part provided that quality management system includes the technical documentation relating to the applicable approved type(s) of medical device(s). Reference to the EC type-examination certificate is not covered.
3.1 2 nd paragraph 8 th indent 3.1 2 nd paragraph 8 th indent (i) 3.1 2 nd paragraph 8 th indent (ii)	(standards.iteh.ai) SIST EN ISO 13485:2016/oprA1:2019 https://standards.iteh.ai/catalog/standards/sist/787ddd2d-4245-4105-bbb0-8803-1ce50d4/sist-en-iso-13485-2016-oprA1-2019	Not covered. This European Standard includes requirements on post-market surveillance, and reporting adverse events and field safety corrective actions to authorities but does not cover all the details required by the Directive including timescales for reporting.
3.2 1 st paragraph		Not covered.
3.2 2 nd paragraph	4.1, 4.2	Covered.
3.2 3 rd paragraph (a)	4.2.1, 5.1, 5.3, 5.4.1	Covered.
3.2 3 rd paragraph (b)	4.2.2	Covered.
3.2 3 rd paragraph (b) 1 st indent	5.1, 5.5.1, 5.5.2	Covered.
3.2 3 rd paragraph (b) 2 nd indent	4.1, 5.6, 7.1, 8.2.4, 8.3, 8.4, 8.5.2, 8.5.3	Covered provided that the methods and acceptance criteria chosen by the manufacturer ensure that the requirements of the Directive are fulfilled.
3.2 3 rd paragraph (b) 3 rd indent	4.1, 4.2, 7.4, 8.2.2	Covered.
3.2 3 rd paragraph (c) 1 st indent	4.2, 6.4, 7.1, 7.4, 7.5, 8.2.5	Covered provided that the quality management system documentation includes relevant documents and records in regards to sterilization and purchasing.
3.2 3 rd paragraph (c) 2 nd indent	4.2, 7.5.8, 7.5.9	Covered.