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I S O 17665-1:2006
Sterilization of health care products — Moist heat — Part 1 : Requirements for
development, validation and routine control of a sterilization process for medical
devices
(IDT)



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1. » (ISO 17665-1:2006 «Sterilization of health care
products — Moist heat — Part 1: Requirements for development, validation and routine control of a sterilization
process for medical devices»).

5 13683—2000. 11134—2000

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Sterilization of health products. Moist heat. Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices

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(ISO 10012. Measurement management systems — Requirements for measurement processes and measuring equipment)

11138-1:2006

1.

(ISO 11138-1:2006. Sterilization of health care products — Biological indicators — Part 1: General requirements)

11138-3:2006

3.

(ISO 11138-3:2006, Sterilization of health care products — Biological indicators — Part 3: Biological indicators for moist heat sterilization processes)

11140-1

1.

(ISO 11140-1, Sterilization of health care products — Chemical indicators — Part 1: General requirements)

11140-3

3.

(ISO 11140-3. Sterilization of care products — Chemical indicators — Part 3: Class 2 indicators for steam penetration test sheets)

11140-4

4.

(ISO 11140-4. Sterilization of health care product — Chemical indicators — Part 4: Class 2 indicators for steam penetration test packs)

11140-5

5.

(ISO 11140-5, Sterilization of health care products — Chemical indicators — Part 5: Class 2 indicators for air removal test sheets and packs)

11607-1

1.

(ISO 11607-1. Packaging for terminally sterilized medical devices — Part 1: Requirements for materials, sterile barrier systems and packaging systems)

11607-2

2.

(ISO 11607-2. Packaging for terminally sterilized medical devices — Part 2: Validation requirements for forming, sealing and assembly processes)

11737-1

1.

(ISO 11737-1. Sterilization of medical devices — Microbiological methods — Part 1: Determination of a population of microorganisms on products)

11737-2

2.

(ISO 11737-2. Sterilization of medical devices — Microbiological methods — Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process)

13485:2003
 (ISO 13485:2003. Medical devices — Quality management systems — Requirements for regulatory purposes)
 17664
 (ISO 17664. Sterilization of medical devices — Information to be provided by the manufacturer for the processing of resterilizabile medical devices)

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(air detector):

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(automatic controller): (),

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(bioburden):

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(biological indicator):

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(calibration):

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indicator): () (chemical indicator) (non-biological

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(contained product):

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(correction):

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(corrective action):

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(development):

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(environmental control):

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3.13 (equilibration time):

3.14 (establish):

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(exposure time):

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(fault):

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3.17 F_0 (F_0 value):

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(health care product(s):

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3.19 (holding time):

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(installation qualification):

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3.21 (load configuration):

3.22 (maintenance):

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(medical device):

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3.24 (measuring chain):

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(microorganism):

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[11139:2006, 2.26]

3.26 (moist heat):

3.27 (non-condensable gas): /

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(operational qualification OQ):

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(operating cycle):

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(packaging system):

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(performance qualification PQ):

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(preventive action):

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3.33 « » (plateau period):

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(process challenge device PCD):

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(process parameter):

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(process variable):

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(product):

[11139:2006, 2.36)

3.36	(product family):	()	,
3.39	(reference challenge device):		-
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3.41	(reference measuring point):		-
3.42	(reference microorganism):		-
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3.43	(requalification):		-
[11139:2006,	2.40]		
3.44	(saturated steam):		-
3.45	(services):		-
[11139:2006,	2.41]		
3.46	(specification):		-
[9000:2005,	3.7.3]		
3.47	(specify):		-
[11139:2006,	2.42]		
3.46	(sterile):		-
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3.49	(sterility):		-
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3.50	(sterility assurance level SAL):		-
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(sterilization):

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() (sterilization toad):

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(sterilization process):

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(sterilization temperature):

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(sterilization temperature band):

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(sterilizer chamber):

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(sterilizing agent):

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(thermal energy):

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(test of sterility):

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(validation):

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9.2 (Installation Qualification, IQ)

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Halverson Zetgler 1932(37).

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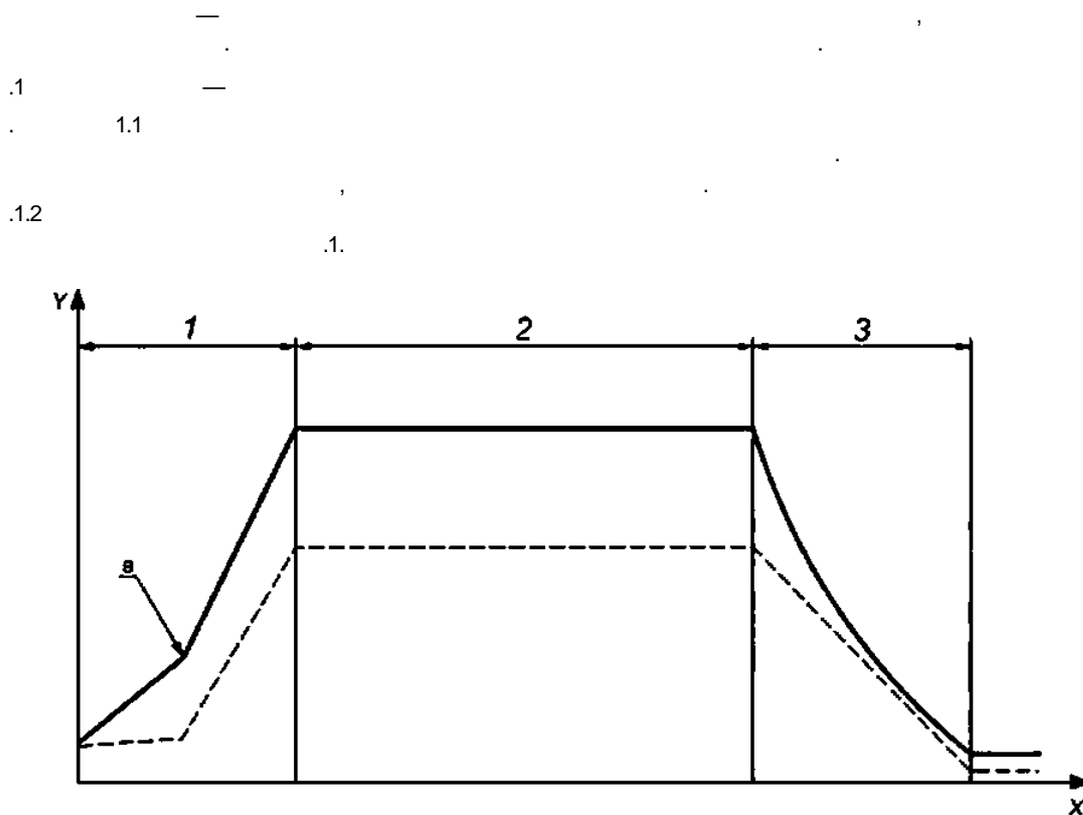
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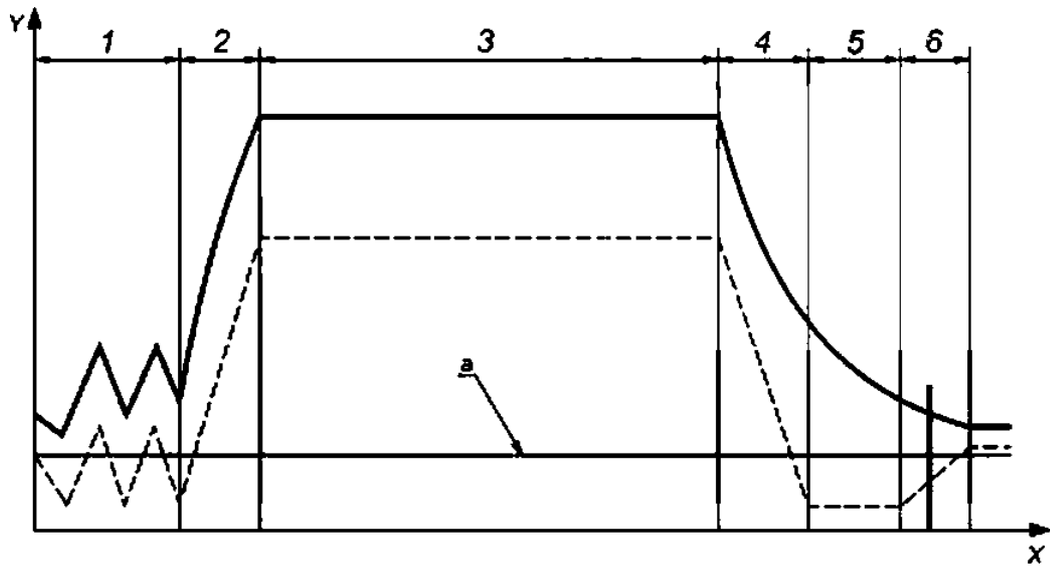
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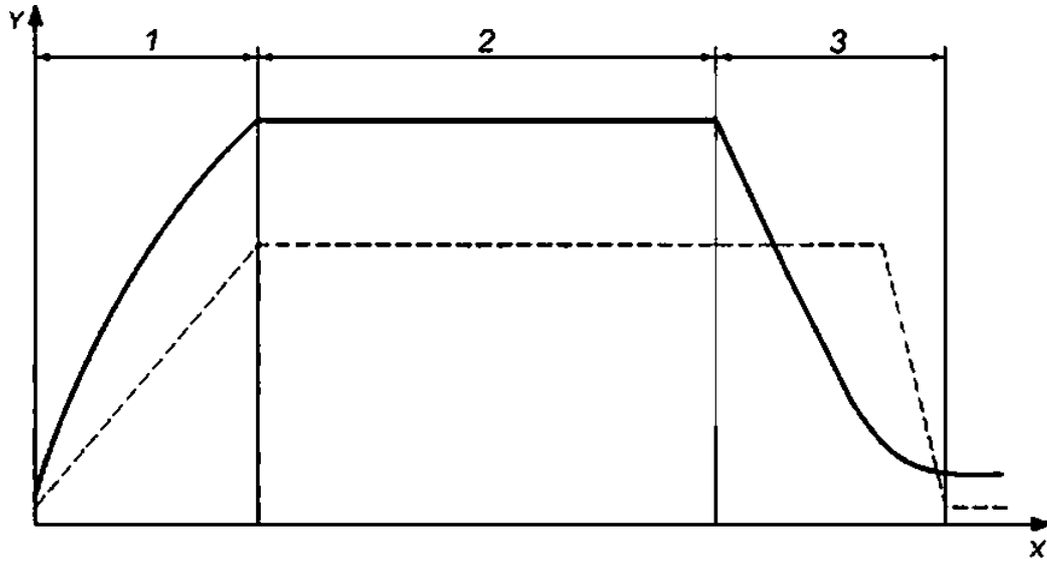
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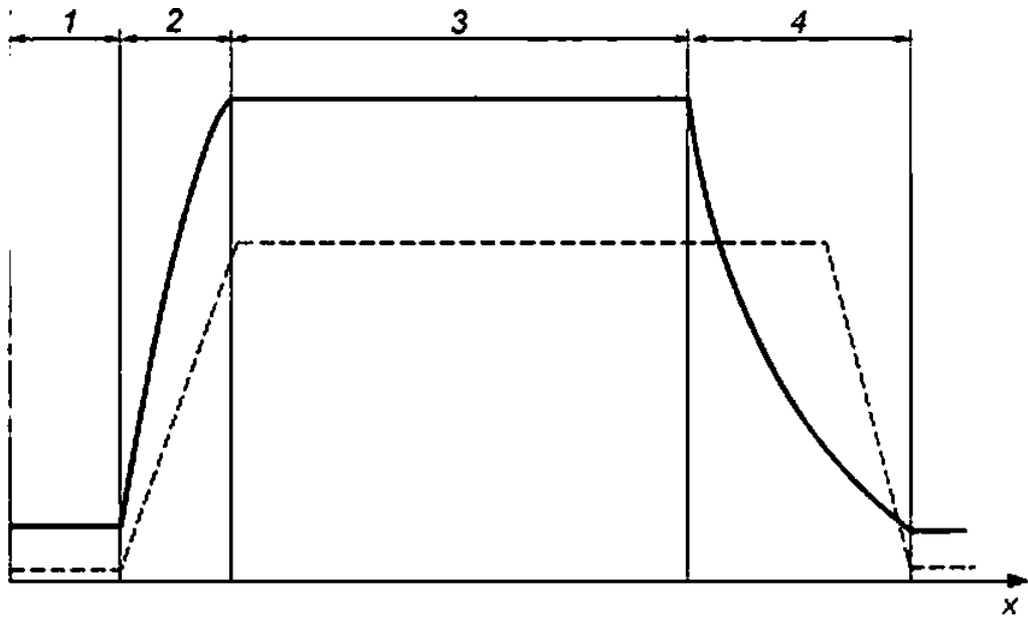
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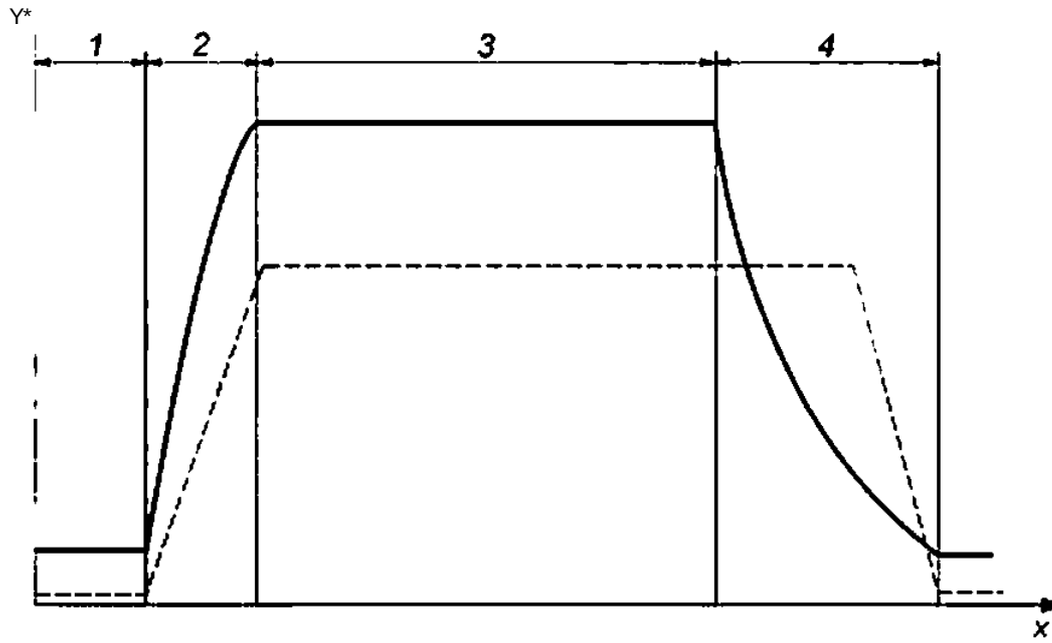
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11140-5		ISO 11140-5—2011 « 5. 2- »
11140-4		ISO 11140-4—2011 « 4. 2- »
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11138-3:2006	—	•
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9000:2005		(Quality management systems. Fundamentals and Vocabulary)	
9000-3:1991	9001	(Quality management and quality assurance standards. Part 3: Guidelines for the application of ISO 9001 to the development, supply and maintenance of computer software)	3.
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9004:2000		(Quality management systems. Guidelines for performance improvements)	-
10993-1		evaluation of medical devices. Part 1: Evaluation and testing)	1. (Biological
10993-17		Establishment of allowable limits for teachable substances)	17. -
/ 11139		Vocabulary)	(Sterilization of healthcare products.
14001		(Environmental management systems. Requirements with guidance for use)	-
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14161:2000		Biological indicators. Guidance for the selection, use and interpretation of results)	(Sterilization < health care products.
14937:2000		(Sterilization of health care products. General requirements for characterization of a sterilizing agent and the development, validation and routine control of a sterilization process for medical devices)	-
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15882:2003		Chemical indicators. Guidance (or selection, use and interpretation of results)	(Sterilization of health care products.
15883-1		(Washer-disinfectors. Part 1: General requirements, terms, definitions and tests)	1. -
15883-2		(Washer-disinfectors. Part 2: Requirements and tests for washer disinfectors employing thermal disinfection for surgical instruments, anaesthetic equipment, bowls, dishes, receivers. utensils, glassware, etc.)	2. -
15883-4		(Washer-disinfectors. Part 4: Requirements and tests for washer-disinfectors employing chemical disinfection for thermolabile endoscopes)	4. -
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- [22] EN 556-1 « » 1. (Sterilization of medical devices. Requirements for medical devices to be designated «STERILE». Part 1: Requirements for terminally-sterilized medical devices)
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- [24] EN 866-2:1999 2. (Packaging materials and systems for medical devices which are to be sterilized. Part 2: Sterilization wrap. Requirements and test methods)
- [25] EN 868-3:1999 3. (EN 868-4) (EN 868-5). (Packaging materials and systems for medical devices which are to be sterilized. Part 3: Paper for use in the manufacture of paper bags (specified in EN 868-4) and in the manufacture of pouches and reels (specified in EN 868-5). Requirements and test methods)
- [26] EN 868-4:1999 4. (Packaging materials and systems for medical devices which are to be sterilized. Part 4: Paper bags. Requirements and test methods)
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- [29] EN 868-9:2000 9. (Packaging materials and systems for medical devices which are to be sterilized. Part 9: Uncoated nonwoven materials of polyolefines for use in the manufacture of heat sealable pouches, reels and lids. Requirements and test methods)
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