

INTERNATIONAL STANDARD

ISO
13408-1

Second edition
2008-06-15

AMENDMENT 1
2013-05-01

Aseptic processing of health care products —

Part 1: General requirements

AMENDMENT 1

Traitement aseptique des produits de santé —

Partie 1: Exigences générales

AMENDEMENT 1

ISO 13408-1:2008/Amd 1:2013

<https://standards.iteh.ai/catalog/standards/iso/9227ace2-cad0-4d8e-9854-2b8470a8c62d/iso-13408-1-2008-amd-1-2013>



Reference number
ISO 13408-1:2008/Amd.1:2013(E)

© ISO 2013

iTeh Standards
(<https://standards.iteh.ai>)
Document Preview

ISO 13408-1:2008/Amd 1:2013

<https://standards.iteh.ai/catalog/standards/iso/9227ace2-cad0-4d8e-9854-2b8470a8c62d/iso-13408-1-2008-amd-1-2013>



COPYRIGHT PROTECTED DOCUMENT

© ISO 2013

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

Published in Switzerland

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

Amendment 1 to ISO 13408-1:2008 was prepared by Technical Committee ISO/TC 198, *Sterilization of health care products*.

iTeh Standards
(<https://standards.iteh.ai>)
Document Preview

[ISO 13408-1:2008/Amd 1:2013](https://standards.iteh.ai/catalog/standards/iso/9227ace2-cad0-4d8e-9854-2b8470a8c62d/iso-13408-1-2008-amd-1-2013)

<https://standards.iteh.ai/catalog/standards/iso/9227ace2-cad0-4d8e-9854-2b8470a8c62d/iso-13408-1-2008-amd-1-2013>

Aseptic processing of health care products —

Part 1: General requirements

AMENDMENT 1

Page vi, Introduction

In the second paragraph, replace the second sentence with the following:

ISO/TC 198 has prepared standards for terminal sterilization of health care products by irradiation (ISO 11137 series), by moist heat (ISO 17665 series), by dry heat (ISO 20857), by ethylene oxide (ISO 11135) and by liquid chemical sterilants (ISO 14160).

Page vii, Introduction

At the end of the last sentence of the penultimate paragraph, add the word “component” so that it reads:

“... of which process simulation studies are an essential component.”

Page 1, Normative references

Delete the following reference:

ISO 9001, *Quality management systems — Requirements*

Page 2, Normative references

Delete footnote 1 and renumber footnote 2 accordingly.

Page 3, 3.7

Delete the following:

[ISO 13408-6:2005, definition 3.1]

Page 4, 3.14

Correct the spelling of the term to read “depyrogenation”.

Page 5, 3.24

Replace the note with the following:

NOTE The required grade of cleanliness of the indirect support zone depends on the aseptic processing technologies and activities performed.

Page 7, 4.1.1

In the first sentence, replace “over all activities affecting aseptic processing” with “over all activities affecting aseptic processing (e.g. ISO 9001 and/or ISO 13485)”.

Delete the second sentence.

Page 7, 4.3.2

Replace the text with the following:

The accuracy and tolerance of all measuring instruments shall be adequate for the parameters to be measured.

Page 8, 5.2.1.2

At the end of the subclause, insert the following note:

NOTE Assessment of risk to condone poor or improper practice during aseptic processing is not appropriate.

Page 10, 5.2.4.4

Replace item a) with the following:

a) microbiological quality of the product at defined stages during the manufacturing process, alert and action levels shall be established;

Page 10, 6.1.2

Replace the second sentence of the note with the following:

Where highly potent, cytotoxic or radioactive health care products are to be processed, protection of personnel and the environment is considered an ancillary element of aseptic processing design.

Page 36, Table 1, fifth column

In the first, third and fifth rows, replace “restart validation” with “repeat initial performance qualification” to be in line with the title of Table 1.

Page 36, Table 2, fourth column

In the first, third and fifth rows, replace “revalidation” with “repeat initial performance qualification” to be in line with the title of Table 1.