

American National Standard

ANSI/AAMI ST79:2006 and
A1:2008, A2:2009
(Consolidated Text)



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Comprehensive guide to steam sterilization and sterility assurance in health care facilities



Association for the Advancement
of Medical Instrumentation

Objectives and uses of AAMI standards and recommended practices

It is most important that the objectives and potential uses of an AAMI product standard or recommended practice are clearly understood. The objectives of AAMI's technical development program derive from AAMI's overall mission: the advancement of medical instrumentation. Essential to such advancement are (1) a continued increase in the safe and effective application of current technologies to patient care, and (2) the encouragement of new technologies. It is AAMI's view that standards and recommended practices can contribute significantly to the advancement of medical instrumentation, provided that they are drafted with attention to these objectives and provided that arbitrary and restrictive uses are avoided.

A voluntary *standard* for a *medical device* recommends to the manufacturer the information that should be provided with or on the product, basic safety and performance criteria that should be considered in qualifying the device for clinical use, and the measurement techniques that can be used to determine whether the device conforms with the safety and performance criteria and/or to compare the performance characteristics of different products. Some standards emphasize the information that should be provided with the device, including performance characteristics, instructions for use, warnings and precautions, and other data considered important in ensuring the safe and effective use of the device in the clinical environment. Recommending the disclosure of performance characteristics often necessitates the development of specialized test methods to facilitate uniformity in reporting; reaching consensus on these tests can represent a considerable part of committee work. When a drafting committee determines that clinical concerns warrant the establishment of *minimum* safety and performance criteria, referee tests must be provided and the reasons for establishing the criteria must be documented in the rationale.

A *recommended practice* provides guidelines for the use, care, and/or processing of a medical device or system. A recommended practice does not address device performance *per se*, but rather procedures and practices that will help ensure that a device is used safely and effectively and that its performance will be maintained.

Although a device standard is primarily directed to the manufacturer, it may also be of value to the potential purchaser or user of the device as a frame of reference for device evaluation. Similarly, even though a recommended practice is usually oriented towards healthcare professionals, it may be useful to the manufacturer in better understanding the environment in which a medical device will be used. Also, some recommended practices, while not addressing device performance criteria, provide guidelines to industrial personnel on such subjects as sterilization processing, methods of collecting data to establish safety and efficacy, human engineering, and other processing or evaluation techniques; such guidelines may be useful to health care professionals in understanding industrial practices.

In determining whether an AAMI standard or recommended practice is relevant to the specific needs of a potential user of the document, several important concepts must be recognized:

All AAMI standards and recommended practices are *voluntary* (unless, of course, they are adopted by government regulatory or procurement authorities). The application of a standard or recommended practice is solely within the discretion and professional judgment of the user of the document.

Each AAMI standard or recommended practice reflects the collective expertise of a committee of health care professionals and industrial representatives, whose work has been reviewed nationally (and sometimes internationally). As such, the consensus recommendations embodied in a standard or recommended practice are intended to respond to clinical needs and, ultimately, to help ensure patient safety. A standard or recommended practice is limited, however, in the sense that it responds generally to perceived risks and conditions that may not always be relevant to specific situations. A standard or recommended practice is an important *reference* in responsible decision-making, but it should never *replace* responsible decision-making.

Despite periodic review and revision (at least once every five years), a standard or recommended practice is necessarily a static document applied to a dynamic technology. Therefore, a standards user must carefully review the reasons why the document was initially developed and the specific rationale for each of its provisions. This review will reveal whether the document remains relevant to the specific needs of the user.

Particular care should be taken in applying a product standard to existing devices and equipment, and in applying a recommended practice to current procedures and practices. While observed or potential risks with existing equipment typically form the basis for the safety and performance criteria defined in a standard, professional judgment must be used in applying these criteria to existing equipment. No single source of information will serve to identify a particular product as "unsafe". A voluntary standard can be used as one resource, but the ultimate decision as to product safety and efficacy must take into account the specifics of its utilization and, of course, cost-benefit considerations. Similarly, a recommended practice should be analyzed in the context of the specific needs and resources of the individual institution or firm. Again, the rationale accompanying each AAMI standard and recommended practice is an excellent guide to the reasoning and data underlying its provision.

In summary, a standard or recommended practice is truly useful only when it is used in conjunction with other sources of information and policy guidance and in the context of professional experience and judgment.

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ANSI/AAMI ST42:1998,
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Developed by
Association for the Advancement of Medical Instrumentation

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Amendment 1 Approved **27 August 2008** by
Amendment 2 Approved **10 July 2009** by
American National Standards Institute Inc.

Abstract: This recommended practice covers steam sterilization in health care facilities. The recommendations are intended to promote sterility assurance and to guide health care personnel in the proper use of processing equipment. Included within the scope of the recommended practice are functional and physical design criteria for sterilization processing areas (decontamination, preparation, sterilization, and sterile storage areas); staff qualifications, education, and other personnel considerations; processing procedures; installation, care, and maintenance of steam sterilizers; quality control; and quality process improvement.

Keywords: cleaning, continuous quality improvement, decontamination, moist heat sterilization, packaging, quality control, quality system, saturated steam, sterile storage, surgical instruments, ambulatory care facilities, dentist office, flash sterilization, sterilization containers, table-top sterilizers

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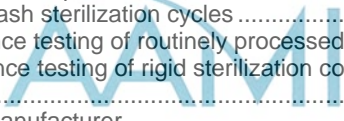
Contents

Glossary of equivalent standards	ix
Committee representation	xi
Acknowledgments	xiii
Background on Amendments 1 and 2	xiii
Foreword	xiv
Introduction: Need for the recommended practice	1
1 Scope	5
1.1 General	5
1.2 Inclusions	5
1.3 Exclusions	5
2 Definitions and abbreviations	6
3 Design considerations	15
3.1 General rationale	15
3.2 Work area design and functional workflow	15
3.2.1 Definitions of work areas	15
3.2.2 Design criteria	16
3.2.3 Functional workflow patterns	18
3.2.4 Traffic control	19
3.3 Physical facilities	21
3.3.1 Space requirements	21
3.3.2 Mechanical systems	21
3.3.3 Electrical systems	21
3.3.4 Steam for sterile processing	21
3.3.5 Utility monitoring and alarm systems	22
3.3.6 General area requirements	22
3.3.7 Special area requirements and restrictions	27
3.3.8 Emergency eyewash/shower equipment	30
3.4 Housekeeping procedures	30
4 Personnel considerations	31
4.1 General rationale	31
4.2 Qualifications	31
4.2.1 Supervisory personnel	31
4.2.2 Sterile processing personnel	31
4.3 Training and continuing education	32
4.3.1 Sterile processing personnel	32
4.3.2 Service personnel	32
4.3.3 Other personnel	33
4.4 Health and personal hygiene	33
4.5 Attire	33
4.5.1 General considerations	33
4.5.2 Decontamination area	34
4.5.3 Sterilization area (flash sterilization)	34
4.5.4 Service personnel	35
4.6 Standard/transmission-based (enhanced) precautions	35
5 Receiving	36
5.1 General rationale	36
5.2 Receiving of purchased or loaner items	36
5.2.1 General considerations	36
5.2.2 Newly purchased reusable items and repaired reusable items	36
5.2.3 Rigid sterilization container systems	36
5.2.4 Disposable items	37
5.3 Disposition of sterile items (issued but not used)	37

6	Handling, collection, and transport of contaminated items	38
6.1	General rationale	38
6.2	Separation of waste and reusable items at point of use.....	38
6.3	Care and handling of contaminated reusable items at point of use.....	38
6.4	Containment	39
6.5	Transport	40
6.5.1	Transportation scheduling and routes.....	40
6.5.2	Transportation equipment.....	40
6.5.3	Hand transport.....	40
6.5.4	Dedicated lifts.....	40
6.5.5	Transport between buildings.....	40
6.5.6	Off-site transportation	41
7	Cleaning and other decontamination processes	43
7.1	General rationale	43
7.2	Policies, procedures, and manufacturers' instructions	43
7.2.1	Policies and procedures	43
7.2.2	Manufacturers' instructions.....	43
7.3	Presoaking.....	44
7.4	Disassembly.....	44
7.4.1	Sorting and disassembly of instrumentation	44
7.4.2	Disassembly of rigid sterilization container systems	44
7.5	Cleaning.....	45
7.5.1	General considerations.....	45
7.5.2	Cleaning agents.....	45
7.5.3	Methods of cleaning	46
7.5.4	Rinsing.....	47
7.5.5	Verification of the cleaning process.....	48
7.5.6	Cleaning of instruments.....	48
7.5.7	Utensils.....	49
7.5.8	Reusable textiles	49
7.5.9	Rigid sterilization container systems.....	49
7.6	Microbicidal processes.....	50
7.6.1	General considerations.....	50
7.6.2	Processes to decontaminate devices so that they are safe to handle	51
7.6.3	Terminal sterilization processes to prepare devices for the next patient use.....	53
7.7	Servicing and repair of devices in the health care facility.....	54
7.7.1	General considerations.....	54
7.7.2	Potential for exposure.....	54
7.7.3	Protective measures for service personnel.....	55
7.7.4	Postexposure program	55
7.7.5	Devices that cannot be repaired in-house	55
8	Packaging, preparation, and sterilization	57
8.1	General rationale	57
8.2	Selection of packaging materials	57
8.3	Package configurations and preparation.....	57
8.3.1	General considerations.....	57
8.3.2	Package labels	58
8.3.3	Package closures	58
8.3.4	Paper-plastic pouches	63
8.3.5	Textile packs.....	63
8.3.6	Basins and basin sets.....	64
8.3.7	Surgical supplies	64
8.3.8	Devices with lumens	65
8.4	Preparation and assembly of surgical instrumentation.....	65
8.4.1	General considerations.....	65
8.4.2	Weight and density of sets.....	65
8.4.3	Inspection	66
8.4.4	Instrument placement	66
8.4.5	Use of tray liners or containment devices	67
8.5	Loading the sterilizer.....	68
8.5.1	General considerations.....	68

8.5.2	Paper-plastic pouches	68
8.5.3	Instrument sets	68
8.5.4	Textile packs.....	68
8.5.5	Utensils and glassware.....	68
8.5.6	Rigid sterilization container systems	70
8.5.7	Liquids and solutions	71
8.5.8	Powders and oils	71
8.6	Sterilization parameters	71
8.6.1	Sterilization parameters for wrapped or containerized items	71
8.6.2	Flash sterilization parameters.....	73
8.6.3	Specialty instruments.....	74
8.7	Monitoring sterilization cycles	74
8.8	Unloading the sterilizer	74
8.8.1	Large-chamber sterilizers	74
8.8.2	Table-top sterilizers	75
8.8.3	Open-tray flash cycles	75
8.8.4	Flash cycles with single wrappers or other textile packaging.....	75
8.8.5	Flash cycles with sealed containment devices.....	75
8.8.6	Handling and inspection	76
8.9	Sterile storage.....	76
8.9.1	Sterility maintenance covers.....	76
8.9.2	Storage facilities	76
8.9.3	Shelf life	77
8.10	Distribution (general).....	77
8.10.1	Handling and inspection	77
8.10.2	Distribution containers	77
8.11	Transport of sterile packaged items.....	78
8.11.1	General considerations.....	78
8.11.2	Tables and carts (open or closed)	78
8.11.3	Hand transport.....	78
8.11.4	Dedicated lifts	78
8.11.5	Off-site transportation.....	78
8.11.6	Policies and procedures	78
8.12	Aseptic presentation	79
8.12.1	Opening sterile packages	79
8.12.2	Removing items from sterile packaging and transferring them to the sterile field	79
9	Installation, care, and maintenance of sterilizers.....	80
9.1	General rationale	80
9.2	Instruction manuals.....	80
9.3	Installation.....	80
9.4	Routine care	80
9.5	Preventive maintenance	80
9.5.1	General considerations.....	80
9.5.2	Scheduled maintenance	81
9.6	Calibration.....	81
9.7	Record-keeping.....	81
10	Quality control	83
10.1	General rationale	83
10.2	Monitoring of mechanical cleaning equipment.....	83
10.3	Product identification and traceability	83
10.3.1	Lot control numbers	83
10.3.2	Sterilizer records.....	84
10.3.3	Expiration dating.....	84
10.4	Overview of sterilization process monitoring.....	85
10.5	Sterilization process monitoring devices.....	88
10.5.1	Physical monitors	88
10.5.2	Chemical indicators (CIs).....	88
10.5.3	Biological indicators	90
10.5.4	Process challenge devices (PCDs).....	91
10.6	Routine load release.....	92
10.6.1	Process monitoring devices.....	92
10.6.2	Release criteria for nonimplants	92

10.6.3	Release criteria for implants	92
10.6.4	Sterilization process failures	92
10.7	Routine sterilizer efficacy monitoring	93
10.7.1	General considerations.....	93
10.7.2	Routine biological monitoring of sterilizers larger than 2 cubic feet	94
10.7.3	Routine biological monitoring of table-top sterilizers.....	96
10.7.4	Routine biological monitoring of flash sterilization cycles.....	97
10.7.5	Actions to take when PCDs (BI challenge test packs or CI challenge test packs) indicate failure.....	98
10.7.6	Routine Bowie-Dick testing of dynamic-air-removal sterilizers.....	100
10.8	Qualification testing	105
10.8.1	General considerations.....	105
10.8.2	Qualification testing of sterilizers larger than 2 cubic feet	106
10.8.3	Qualification testing of table-top sterilizers	108
10.8.4	Qualification testing of flash sterilization cycles	109
10.9	Periodic product quality assurance testing of routinely processed items	110
10.10	Periodic product quality assurance testing of rigid sterilization container systems.....	111
10.10.1	General considerations.....	111
10.10.2	Responsibilities of the manufacturer.....	111
10.10.3	User responsibilities.....	113
10.11	Product recalls	118
10.11.1	General considerations.....	118
10.11.2	Recall procedure.....	118
10.11.3	Recall order	118
10.11.4	Recall report	118
11	Quality process improvement.....	119
11.1	General rationale	119
11.2	Quality process	119
11.2.1	General considerations.....	119
11.2.2	Risk analysis.....	119
11.2.3	Decontamination.....	119
11.2.4	Rigid sterilization container systems.....	121
11.2.5	Flash sterilization.....	122
11.3	Functional areas for product and process improvements	122
11.3.1	Workplace design	122
11.3.2	Processing policies and procedures	123
11.3.3	Product use	123
11.4	Implementing product and process improvements	123
Annexes		
A	Examples of workplace design.....	125
B	Infection transmission	133
C	Processing CJD-contaminated patient care equipment and environmental surfaces.....	137
D	User verification of cleaning processes.....	142
E	Selection and use of chemical disinfectants.....	147
F	Thermal disinfection.....	153
G	Devices returned to the manufacturer	154
H	Occupational exposure to blood-borne pathogens (29 CFR Part 1910.1030).....	161
I	Development of a prepurchase evaluation protocol for rigid sterilization container systems.....	175
J	Effect of containerized packaging on load heat-up time.....	180
K	Development and qualification of the 16 towel PCD (biological-indicator challenge test pack)	182
L	Example of documentation of premature release of implants	191
M	Steam quality	193
N	Toxic anterior segment syndrome (TASS) and the processing of intraocular surgical instruments.....	195


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O	Bibliography	199
----------	--------------------	-----

Tables

1	Saturated steam pressure conversion units at sea level	12
2	Ventilation requirements for functional areas	24
3	IES-recommended illuminance levels for work environments	26
4	Minimum cycle times for gravity-displacement steam sterilization cycles	72
5	Minimum cycle times for dynamic-air-removal steam sterilization cycles	72
6	Sterilization process monitoring recommendations	86
7	Types and applications for use of sterilization monitoring devices	87
8	Checklist for identifying reasons for steam sterilization process failures	101
9	Summary of test configurations for prepurchase evaluation of rigid sterilization container systems	115
D.1	In-use tests available to assess efficacy of cleaning of medical devices	145
D.2	In-use tests available to assess efficacy of washer-disinfectors used for medical device reprocessing	146
E.1	Levels of disinfection according to type of microorganism	149
E.2	Occupational exposure limits for some chemical sterilants and disinfectants	152
K.1	16 towel pack survey	183
K.2	Biological-indicator results from 121 °C (250 °F) gravity cycle	185
K.3	Biological-indicator results from 132 °C (270 °F) deep-vacuum cycle	186
K.4	Biological-indicator results from 132 °C (270 °F) pulsing vacuum cycle	187
K.5	Comparison of the 16 towel pack with the 12 × 12 × 20 inch pack by Most Probable Number and sterility assessment of spore strips (121 °C [250 °F] gravity cycle)	188
K.6	Fraction-negative results in a 121 °C (250 °F) gravity cycle	188
K.7	Biological-indicator results from 121 °C (250 °F) steam-flush pressure-pulse cycle	189
K.8	Biological-indicator results from 132 °C (270 °F) steam-flush pressure-pulse cycle	189

Figures

1	Functional work areas of a sterile processing department	16
2	Workflow	20
3	Microbicidal processes and use of PPE	51
4	Sequential double-wrapping: envelope fold	59
5	Sequential double-wrapping: square fold	60
6	Simultaneous double-wrapping: envelope fold	61
7	Simultaneous double-wrapping: square fold	62
8	Example of single- and double-packaging with paper/plastic pouches	63
9	Loading the sterilizer	69
10	Preparation of the 16 towel PCD (BI challenge test pack)	95
11	Placement of the 16 towel PCD (BI challenge test pack) for routine biological monitoring of sterilizers larger than 2 cubic feet	95
12	Decision tree for conducting investigations of steam sterilization process failures	100

13	Composition of the Bowie-Dick test pack	104
14	Placement of the Bowie-Dick test pack.....	105
15	Placement of the 16 towel PCD (BI challenge test pack) for qualification testing	107
A.1	Example of a work area design and workflow pattern for a sterile processing department in a typical small hospital	125
A.2	Example of a work area design and workflow pattern for a sterile processing department in a typical medium-sized hospital	126
A.3	Example of a work area design and workflow pattern for a sterile processing department in a typical regional processing center.....	127
A.4	Example of a work area design and workflow pattern for a sterile processing department	128
A.5	Example of a work area design and workflow pattern for a sterile processing department	129
A.6	Example of a work area design and workflow pattern for a sterile processing department	130
A.7	Example of an ambulatory surgery facility	131
A.8	Example of a dental facility	132
B.1	The chain of infection, components of the infectious disease process.....	133
B.2	Blood-borne pathogen strike-through conversion chart.....	135
J.1	Typical rigid sterilization container system processed in a gravity-displacement cycle at 121 °C (250 °F).....	180
J.2	Muslin-wrapped, 16-pound instrument set processed in a gravity-displacement cycle at 121 °C (250 °F).....	181
J.3	Typical rigid sterilization container system processed in a prevacuum cycle at 132 °C (270 °F).....	181
K.1	Temperature profiles for two different configurations of 12 x 12 x 20 inch packs in a 121 °C (250 °F) gravity cycle.....	184
K.2	Temperature profiles for huck and absorbent 16 towel packs in a 121 °C (250 °F) gravity cycle	184
K.3	Average temperature profile for the 16 towel pack in a 121 °C (250 °F) gravity cycle	186
L.1	Implantable devices load record	191
L.2	Exception form for premature release of implantable device/tray	192



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International designation	U.S. designation	Equivalency
IEC 60601-1:2005	ANSI/AAMI ES60601-1:2005	Major technical variations
IEC 60601-1-2:2007	ANSI/AAMI/IEC 60601-1-2:2007	Identical
IEC 60601-2-2:2009	ANSI/AAMI/IEC 60601-2-2:2009	Identical
IEC 60601-2-4:2002	ANSI/AAMI DF80:2003	Major technical variations
IEC 60601-2-19:2009	ANSI/AAMI/IEC 60601-2-19:2009	Identical
IEC 60601-2-20:2009	ANSI/AAMI/IEC 60601-2-20:2009	Identical
IEC 60601-2-21:2009	ANSI/AAMI/IEC 60601-2-21:2009	Identical
IEC 60601-2-24:1998	ANSI/AAMI ID26:2004	Major technical variations
IEC 60601-2-47:2001	ANSI/AAMI EC38:2007	Major technical variations
IEC 60601-2-50:2009	ANSI/AAMI/IEC 60601-2-50:2009	Identical
IEC 80601-2-58:2008	ANSI/AAMI/IEC 80601-2-58:2008	Identical
IEC/TR 60878:2009	ANSI/AAMI/IEC TIR60878:2003	Identical
IEC/TR 62296:2009	ANSI/AAMI/IEC TIR62296:2009	Identical
IEC 62304:2006	ANSI/AAMI/IEC 62304:2006	Identical
IEC/TR 62348:2006	ANSI/AAMI/IEC TIR62348:2006	Identical
ISO 5840:2005	ANSI/AAMI/ISO 5840:2005	Identical
ISO 7198:1998	ANSI/AAMI/ISO 7198:1998/2001/(R)2004	Identical
ISO 7199:2009	ANSI/AAMI/ISO 7199:2009	Identical
ISO 8637:2004	ANSI/AAMI RD16:2007	Major technical variations
ISO 8638:2004	ANSI/AAMI RD17:2007	Major technical variations
ISO 10993-1:2003	ANSI/AAMI/ISO 10993-1:2003	Identical
ISO 10993-2:2006	ANSI/AAMI/ISO 10993-2:2006	Identical
ISO 10993-3:2003	ANSI/AAMI/ISO 10993-3:2003	Identical
ISO 10993-4:2002 and Amendment 1:2006	ANSI/AAMI/ISO 10993-4:2002/(R)2009 and Amendment 1:2006/(R)2009	Identical
ISO 10993-5:2009	ANSI/AAMI/ISO 10993-5:2009	Identical
ISO 10993-6:2007	ANSI/AAMI/ISO 10993-6:2007	Identical
ISO 10993-7:2008	ANSI/AAMI/ISO 10993-7:2008	Identical
ISO 10993-9:1999	ANSI/AAMI/ISO 10993-9:1999/(R)2005	Identical
ISO 10993-10:2002 and Amendment 1:2006	ANSI/AAMI BE78:2002/(R)2008 ANSI/AAMI BE78:2002/A1:2006/(R)2008	Minor technical variations Identical
ISO 10993-11:2006	ANSI/AAMI/ISO 10993-11:2006	Identical
ISO 10993-12:2007	ANSI/AAMI/ISO 10993-12:2007	Identical
ISO 10993-13:1998	ANSI/AAMI/ISO 10993-13:1999/(R)2004	Identical
ISO 10993-14:2001	ANSI/AAMI/ISO 10993-14:2001/(R)2006	Identical
ISO 10993-15:2000	ANSI/AAMI/ISO 10993-15:2000/(R)2006	Identical
ISO 10993-16:1997	ANSI/AAMI/ISO 10993-16:1997/(R)2009	Identical
ISO 10993-17:2002	ANSI/AAMI/ISO 10993-17:2002/(R)2008	Identical
ISO 10993-18:2005	ANSI/AAMI BE83:2006	Major technical variations
ISO/TS 10993-19:2006	ANSI/AAMI/ISO TIR10993-19:2006	Identical
ISO/TS 10993-20:2006	ANSI/AAMI/ISO TIR10993-20:2006	Identical
ISO 11135-1:2007	ANSI/AAMI/ISO 11135-1:2007	Identical

International designation	U.S. designation	Equivalency
ISO/TS 11135-2:2008	ANSI/AAMI/ISO TIR11135-2:2008	Identical
ISO 11137-1:2006	ANSI/AAMI/ISO 11137-1:2006	Identical
ISO 11137-2:2006 (2006-08-01 corrected version)	ANSI/AAMI/ISO 11137-2:2006	Identical
ISO 11137-3:2006	ANSI/AAMI/ISO 11137-3:2006	Identical
ISO 11138-1: 2006	ANSI/AAMI/ISO 11138-1:2006	Identical
ISO 11138-2: 2006	ANSI/AAMI/ISO 11138-2:2006	Identical
ISO 11138-3: 2006	ANSI/AAMI/ISO 11138-3:2006	Identical
ISO 11138-4: 2006	ANSI/AAMI/ISO 11138-4:2006	Identical
ISO 11138-5: 2006	ANSI/AAMI/ISO 11138-5:2006	Identical
ISO/TS 11139:2006	ANSI/AAMI/ISO 11139:2006	Identical
ISO 11140-1:2005	ANSI/AAMI/ISO 11140-1:2005	Identical
ISO 11140-3:2007	ANSI/AAMI/ISO 11140-3:2007	Identical
ISO 11140-4:2007	ANSI/AAMI/ISO 11140-4:2007	Identical
ISO 11140-5:2007	ANSI/AAMI/ISO 11140-5:2007	Identical
ISO 11607-1:2006	ANSI/AAMI/ISO 11607-1:2006	Identical
ISO 11607-2:2006	ANSI/AAMI/ISO 11607-2:2006	Identical
ISO 11737-1: 2006	ANSI/AAMI/ISO 11737-1:2006	Identical
ISO 11737-2:1998	ANSI/AAMI/ISO 11737-2:1998	Identical
ISO 13408-1:2008	ANSI/AAMI/ISO 13408-1:2008	Identical
ISO 13408-2:2003	ANSI/AAMI/ISO 13408-2:2003	Identical
ISO 13408-3:2006	ANSI/AAMI/ISO 13408-3:2006	Identical
ISO 13408-4:2005	ANSI/AAMI/ISO 13408-4:2005	Identical
ISO 13408-5:2006	ANSI/AAMI/ISO 13408-5:2006	Identical
ISO 13408-6:2006	ANSI/AAMI/ISO 13408-6:2006	Identical
ISO 13485:2003	ANSI/AAMI/ISO 13485:2003	Identical
ISO 14155-1:2003	ANSI/AAMI/ISO 14155-1:2003/(R)2008	Identical
ISO 14155-2:2003	ANSI/AAMI/ISO 14155-2:2003/(R)2008	Identical
ISO 14160:1998	ANSI/AAMI/ISO 14160:1998/(R)2008	Identical
ISO 14161:2000	ANSI/AAMI/ISO 14161:2000	Identical
ISO 14937:2000	ANSI/AAMI/ISO 14937:2000	Identical
ISO/TR 14969:2004	ANSI/AAMI/ISO TIR14969:2004	Identical
ISO 14971:2007	ANSI/AAMI/ISO 14971:2007	Identical
ISO 15223-1:2007 and A1:2008	ANSI/AAMI/ISO 15223-1:2007 and A1:2008	Identical
ISO 15225:2000 and A1:2004	ANSI/AAMI/ISO 15225:2000/(R)2006 and A1:2004/(R)2006	Identical
ISO 15674:2009	ANSI/AAMI/ISO 15674:2009	Identical
ISO 15675:2009	ANSI/AAMI/ISO 15675:2009	Identical
ISO 15882:2008	ANSI/AAMI/ISO 15882:2008	Identical
ISO/TR 16142:2006	ANSI/AAMI/ISO TIR16142:2005	Identical
ISO 17664:2004	ANSI/AAMI ST81:2004	Major technical variations
ISO 17665-1:2006	ANSI/AAMI/ISO 17665-1:2006	Identical (with inclusions)
ISO/TS 17665-2:2009	ANSI/AAMI/ISO TIR17665-2:2009	Identical
ISO 18472:2006	ANSI/AAMI/ISO 18472:2006	Identical
ISO/TS 19218:2005	ANSI/AAMI/ISO 19218:2005	Identical
ISO 22442-1:2007	ANSI/AAMI/ISO 22442-1:2007	Identical
ISO 22442-2:2007	ANSI/AAMI/ISO 22442-2:2007	Identical
ISO 22442-3:2007	ANSI/AAMI/ISO 22442-3:2007	Identical
ISO 25539-1:2003 and A1:2005	ANSI/AAMI/ISO 25539-1:2003 and A1:2005	Identical
ISO 25539-2:2008	ANSI/AAMI/ISO 25539-2:2008	Identical
ISO 81060-1:2007	ANSI/AAMI/ISO 81060-1:2007	Identical

Committee representation

Association for the Advancement of Medical Instrumentation Steam Sterilization Hospital Practices Working Group

This recommended practice was developed by the AAMI Steam Sterilization Hospital Practices Working Group under the auspices of the AAMI Sterilization Standards Committee. Approval of the recommended practice does not necessarily mean that all working group members voted for its approval.


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Acknowledgments

The AAMI Steam Sterilization Hospital Practices Working Group gratefully acknowledges the important contributions of Neal Danielson, who cochaired the working group for many years and who prepared the first draft of this comprehensive guideline on steam sterilization in health care facilities. His contribution of time, effort, and expertise to the development of AAMI recommended practices and to the field of central service generally cannot be overstated. The working group also wishes to acknowledge the contributions of former working group member Dorothy Fogg, RN, who for many years represented the Association of periOperative Registered Nurses on the working group; former working group member Gregory Stecklein of Cardinal Health; and former working group member Retta C. Sengstock, RN, Implementation Specialists for Healthcare Inc. Martin S. Favero, PhD, of Advanced Sterilization Products served as industry cochair of the working group from 2000 to 2004, and the time and expertise that he devoted to the development of this recommended practice are very much appreciated by the working group. Finally, this recommended practice could not have come to fruition without the hard work and diplomatic skills of Barbara Goodman, RN, who cochaired the working group from 1995 to 2004; the working group is extremely grateful for her commitment and her immense contributions to this project.

Background on Amendments 1 and 2

As a continuously maintained Recommended Practice, this document consolidates the text of ST79:2006, ST79:2006/A1:2008, and ST79:2006/A2:2009. Please see Amendments 1 and 2 to identify exactly what has changed. Amendment 1 shows modifications to the 2006 edition of ST79 in redline/strikeout. Amendment 2 shows modifications to the consolidated text of ST79:2006 and ST79:2006/A1:2008 in redline/strikeout. Each of these amendments is available in print or as a free PDF at <http://marketplace.aami.org>.

NOTE—(1 September 2009) The AAMI Steam Sterilization Hospital Practices Working Group is beginning a revision of ST79, therefore with publication of the 2009 Amendment, the current edition is no longer on Continuous Maintenance. If the Working Group decides to put the next edition on Continuous Maintenance, this will be announced when that document is completed and published.

Foreword

This recommended practice was developed by the Steam Sterilization Hospital Practices Working Group of the AAMI Sterilization Standards Committee. The purpose of the guidelines in this document is to help ensure the steam sterilization of products in health care facilities and the maintenance of the sterility of processed items until the point of use.

To facilitate user access to all AAMI consensus recommendations for steam sterilization in health care facilities, the committee has consolidated into one comprehensive guide the following AAMI recommended practices:

- ANSI/AAMI ST46, *Steam sterilization and sterility assurance in health care facilities*
- ANSI/AAMI ST42, *Steam sterilization and sterility assurance using table-top sterilizers in office-based, ambulatory-care medical, surgical, and dental facilities*
- ANSI/AAMI ST37, *Flash sterilization: Steam sterilization of patient care items for immediate use*
- ANSI/AAMI ST35, *Safe handling and biological decontamination of medical devices in health care facilities and in nonclinical settings*
- ANSI/AAMI ST33, *Guidelines for the selection and use of reusable rigid sterilization container systems for ethylene oxide sterilization and steam sterilization in health care facilities*

In the course of the consolidation process, the five recommended practices listed above were updated and revised to reflect current good practice. Several annexes were added to provide additional information to users. The new recommended practice serves as a comprehensive guideline for all steam sterilization activities in health care facilities, regardless of the size of the sterilizer or the size of the facility, and provides a resource for all health care personnel who use steam for sterilization.

This recommended practice reflects the conscientious efforts of health care professionals, in cooperation with medical device and equipment manufacturers, to develop recommendations for optimum performance levels in the processing of reusable medical devices to be steam sterilized. It is not intended that these recommendations be construed as universally applicable in all circumstances. Also, it is recognized that in many cases these recommendations might not be immediately achievable. Therefore, the document should be used to guide personnel towards desirable performance objectives, and all of its provisions should be considered and applied in the light of professional judgment and experience.

As used within the context of this document, “shall” indicates requirements strictly to be followed to conform to the recommended practice. “Should” indicates that among several possibilities, one is recommended as particularly suitable, without mentioning or excluding others, or that a certain course of action is preferred but not necessarily required, or that (in the negative form) a certain possibility or course of action should be avoided but is not prohibited. “May” is used to indicate that a course of action is permissible within the limits of the recommended practice. “Can” is used as a statement of possibility and capability. Finally, “must” is used only to describe “unavoidable” situations, including those mandated by government regulation.

The provisions of this recommended practice should be reviewed by departmental managers and adapted to the needs of their particular institutions. Written policies and procedures should be developed and implemented in consultation with appropriate hospital committees (e.g., safety, infection prevention and control, and hazardous materials).

The concepts incorporated in this recommended practice should be considered flexible and dynamic. The recommendations set forth in this document are reviewed and updated periodically to assimilate progressive technological developments. AAMI policies and procedures require that AAMI standards and recommended practices be reviewed and, if necessary, revised at least once every five years.

This standard is maintained under continuous maintenance procedures. AAMI has created a notification registry that will send e-mail announcements when any maintenance activity occurs to the recommended practice. To register, visit www.aami.org/standards/st79.registry. Suggestions for improving this recommended practice are invited. Comments or proposals for revisions to any part of the standard may be submitted to AAMI any time. Written comments are to be sent to: Standards Dept., AAMI, 1110 N. Glebe Road, Suite 220, Arlington, VA 22201-4795. Comments may also be e-mailed to: standards@aami.org.

NOTE—This foreword does not contain provisions of the AAMI recommended practice, *Comprehensive guide to steam sterilization and sterility assurance in health care facilities* (ANSI/AAMI ST79), but it does provide important information about the development and intended use of the document.

Comprehensive guide to steam sterilization and sterility assurance in health care facilities

Introduction: Need for the recommended practice

Overview:

Saturated steam under pressure is one of the oldest methods used in health care facilities to sterilize medical devices. Because this method has been available for so many years, it is thought to be a simple process, one that is well understood and controlled. However, the efficacy of any sterilization process, including saturated steam, depends on a consistent system for lowering and limiting bioburden before sterilization, properly preparing items for sterilization, selecting the appropriate sterilization parameters, and establishing and implementing controls to maintain the sterility of sterilized items until they are used. These four phases are critically interdependent, and each must be accomplished to produce and maintain a sterile product.

The delivery of sterile health care products for use in patient care depends not only on the efficacy of the sterilization process itself but also on the following factors:

- a) efficient facility design,
- b) proper training of personnel,
- c) good infection prevention and control practices designed to prevent health-care-associated infections,
- d) effective quality control and process improvement systems that encompass all aspects of device reprocessing from point of use through sterilization to reuse, and
- e) appropriate documentation and reporting practices that enable traceability of each facility-sterilized medical device to the patient on whom it was used.

Health care facilities differ in their physical design and equipment and in the level of personnel expertise, competence, and training. This recommended practice has been developed to set forth guidelines for facility design, work practices, and process controls that will help ensure that sterile items are consistently produced using saturated steam under pressure.

This recommended practice addresses elements of a quality system, but it is not intended to provide comprehensive guidance on this subject.

Many of the activities that affect sterilization processing occur in areas separate from the location where sterilization is actually carried out. Therefore, the policies and procedures governing sterilization processing should be developed in consultation with the managers of areas that use sterile medical devices and with appropriate committees or functional groups within the facility (e.g., infection prevention and control, safety, hazardous materials, risk management). In addition, the support of the facility's administration is vital, especially in those facilities where the establishment of a quality system to implement steam sterilization process validation and parametric release is being considered.

It might not be possible for a health care facility to implement all the provisions of this recommended practice because of environmental restrictions and/or limitations in capital funding. However, it is recommended that the health care facility's administration be made aware of any current deficiencies so that the allocation of needed resources can be planned.

This recommended practice encompasses steam sterilization in all health care facilities, including ambulatory-care and office-based facilities. It covers steam sterilization by both the wrapped and unwrapped (flash) methods and

provides detailed guidance on decontamination and packaging, with special reference to rigid sterilization container systems.

Steam sterilization in office-based, ambulatory-care medical, surgical, and dental facilities:

Advances in medical, surgical, and dental practice have led to the increased use of alternative health care sites, such as offices, ambulatory-care clinics, and similar clinical settings; many such facilities use small table-top steam sterilizers. Office-based practices can differ greatly from hospitals in their physical design and in the training level of personnel. The general concepts in this recommended practice apply to these settings. In some sections, processes or equipment used most frequently within the office-based and ambulatory setting are specifically addressed.

Flash sterilization:

A flash sterilization cycle is one that has been designed to meet the following criteria:

- a) The cycle is preprogrammed to a specific time-temperature setting established by the manufacturer based on the type of sterilizer control (i.e., gravity-displacement, dynamic-air-removal) and selected by the user based on the configuration of the load (i.e., the presence or absence of porous materials).
- b) The items to be processed are usually unwrapped, although a single wrapper may be used in certain circumstances if the sterilizer or packaging manufacturer's instructions permit. Some rigid sterilization container systems have been designed and validated by the container manufacturer for use with flash cycles.
- c) Since drying time is not usually part of a preprogrammed flash cycle, the items processed are assumed to be wet at the conclusion of the cycle.
- d) The processed items(s) must be transferred immediately, using aseptic technique, from the sterilizer to the actual point of use, usually the sterile field in an ongoing surgical procedure. Regardless of whether the items are wrapped, there is no storage or shelf life of flash-sterilized items because of the higher probability of contamination after the sterilizer door is opened and the items are removed.

It is essential for health care personnel to properly carry out the complete multistep process (including decontamination and preparation) when flash sterilization is used, just as in the case of items to be processed using wrapped-goods sterilization cycles. In any method of sterilization, it is important to adhere to good processing practices. Such practices are particularly important in flash sterilization because of the difficulties associated with the aseptic delivery of devices sterilized by this method to the point of use. When performed correctly, flash sterilization is safe and effective for the sterilization of medical devices intended for use in contact with compromised tissue or the vascular system, as might occur during surgery or obstetrical delivery. The exposure times used in flash sterilization cycles are capable of producing appropriate lethality.

Several concerns stimulated the development of guidelines for flash sterilization. First, the committee was aware of inadequate cleaning and other decontamination processes in flash sterilization. Reduction of bioburden and removal of gross soil are essential steps in preparing an item for sterilization by any method. Decontamination procedures are also designed to protect the worker.

Second, documentation of the flash sterilization process is necessary and should be consistent with the requirements applicable to and the practices used in documenting the routine processing of wrapped loads.

Third, flash-sterilized items should be transported to the point of use in such a way that the potential for contamination is minimized. In deciding on transport techniques for a particular situation, personnel should consider the possible ways in which the items could become contaminated and the safety of workers handling the hot, wet, and possibly heavy trays. Contamination is an event-related process, with the probability of an event that could result in contamination increasing over time. When opened to the air, all sterile items will eventually become contaminated unless opened within and kept in a true HEPA-filtered, laminar-air-flow unit. Thus, any item that is opened and left on the back table of a surgical setup can become contaminated by particles settling on it. The longer an item is open, the greater the number of particles, with their accompanying microbiological flora.

The risk of contamination of flash-sterilized items increases if they are transported through areas where personnel are scrubbing or washing their hands, creating splashing or aerosolization. Transport through areas where air flow is not filtered to the degree present in the operating room (OR) can also increase the rate of contamination. Practitioners should examine their own situations and develop practices to minimize contamination. Some methods are placing flash sterilizers as close to the intended point of use as can be reasonably accomplished, using rigid sterilization container systems that have been specifically validated and labeled for use in flash sterilization, using the single-

wrapper technique in appropriate cycles, and aseptically placing a sterile covering completely around the sterilized item as it is removed from the sterilizer.

Finally, flash sterilization of instrumentation should be considered only if all the following conditions are met:

- a) Work practices ensure proper cleaning and decontamination, inspection, and arrangement of instruments into the recommended sterilizing trays or other containment devices before sterilization.
- b) The physical layout of the department or work area ensures direct delivery of sterilized items to the point of use (e.g., the sterilizer opens into an area either within or directly adjacent to the procedure room).
- c) Procedures are developed, followed, and audited to ensure aseptic handling and personnel safety during transfer of the sterilized items from the sterilizer to the point of use.
- d) The item is needed for use immediately following flash sterilization.

Implantables should not be flash-sterilized (Garner and Favero, 1985; CDC, 2003a, 2003b). The possible consequences to the patient from placing even a minimally contaminated device in an essentially avascular environment and leaving it there at the conclusion of the procedure are potentially severe. Although the risk of an unrecognized sterilization failure can be minimized if the physical parameters of time, temperature, and pressure are monitored and recorded and the results examined after each cycle, it is recommended that health care personnel quarantine implantable devices and await the outcome of biological monitoring of the cycle before releasing these items for use. Current technology allows for release of loads, even those containing implants, upon obtaining results from the early readout mechanism of a BI designed and labeled for such use. However, this technology does not solve the problems with using flash sterilization for implants. Concerns about aseptic transfer remain, especially if the sterilizer does not open directly into the room containing the sterile field where the device will be used or into an area either within or directly adjacent to the procedure room. Careful planning, appropriate packaging (e.g., packaging that allows the user to see the device for sizing and verification of features), and inventory management in cooperation with suppliers can eliminate the need to flash sterilize implantable items. This is a goal that all institutions should strive to achieve.

This recommended practice incorporates guidelines that are specifically applicable to flash sterilization.

Decontamination:

All microorganisms in health care facilities should be considered potentially pathogenic. Their ability to produce an infection or disease process depends on several factors, including the number and virulence of infectious organisms, the presence of a portal of entry, and the susceptibility of the host (see Annex B). Medical devices, instruments, and equipment used in patient care become contaminated with microorganisms and must be decontaminated.

Decontamination is the process by which medical devices, instruments, and equipment are rendered safe for personnel to handle. In some cases, the decontamination process is sufficient to render the items safe for reuse in patient care. The type and level of decontamination required is determined by the circumstances of device use, the type of patient contact, and the likelihood of biological hazard to personnel.

Infection prevention is enhanced when (a) soiled supplies and equipment are correctly and safely handled, and (b) reusable medical items are thoroughly cleaned. Whenever cleaning is not sufficient to render an item safe for personnel handling, the item is subjected to a subsequent microbicidal process that has been designed to provide an appropriate level of microbial lethality (kill). This process could be a disinfection process or a sterilization process. The microbicidal process might not be effective if soil has not been first removed by cleaning. When used for decontamination purposes, a microbicidal process does not necessarily make an item safe for patient use, because the level of microbial kill might not be sufficient for the intended use (as in the case of surgical instruments needed for sterile procedures).

Adherence to the principles of infection prevention and control will help prevent the spread of potentially infectious or disease-producing microorganisms from one person to another and will help ensure that all items are safe for handling during inspection, assembly, preparation, and packaging. In addition, adherence to these principles is one of the essential factors in achieving effective terminal sterilization processing, when appropriate for a particular reusable item.

The selection of an appropriate decontamination method is complex because of the huge variety of reusable items and the wide range of processes for achieving various levels of decontamination. There are diverse and often conflicting recommendations for handling supplies and equipment and for controlling biological hazards through decontamination methods. These diverse recommendations have been provided to health care personnel by professional organizations, government agencies, manufacturers of decontamination products and equipment,

medical device manufacturers, consultants, and educational speakers. There is clearly a need for consensus guidelines, with supporting rationale, for decontamination processing techniques.

The objectives of the guidelines provided in this recommended practice are to (a) help reduce the risk of cross-infection by pathogenic microorganisms to patients, personnel, and other persons; (b) assist in the development of decontamination procedures based on knowledge and scientific data; and (c) help ensure that all reusable medical devices are handled, transported, cleaned, biologically decontaminated, and reprocessed or examined under the best possible conditions for maximum safety.

Rigid sterilization container systems:

The packaging section of this recommended practice provides detailed guidelines on the selection and use of rigid sterilization container systems intended for use in steam sterilization. These systems serve as packaging for items before, during, and after sterilization. They may also be used to contain and transport contaminated items after use. Special considerations apply to these packaging systems to ensure adequate sterilant penetration and air removal.



This is a preview edition of an AAMI guidance document and is intended to allow potential purchasers to evaluate the content of the document before making a purchasing decision.

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1 Scope

1.1 General

This recommended practice provides guidelines for decontamination and steam sterilization processing in hospitals and other health care facilities. These guidelines are intended to promote sterility assurance and to assist health care personnel in the proper use of processing equipment.

NOTE—For purposes of this recommended practice, “health care facilities” means hospitals, nursing homes, extended-care facilities, free-standing surgical centers, clinics, and medical and dental offices. For convenience, the term “hospital” is sometimes used in this recommended practice; in all instances, this term should be taken to encompass all other health care facilities.

1.2 Inclusions

This recommended practice specifically addresses

- a) functional and physical design criteria for sterilization processing areas;
- b) staff qualifications, education, and other personnel considerations;
- c) processing recommendations;
- d) installation, care, and maintenance of steam sterilizers;
- e) quality control; and
- f) quality process improvement.

Definitions of terms, a bibliography, and informative annexes also are provided in this recommended practice.

1.3 Exclusions

This recommended practice does not cover

- a) specific construction and performance criteria for steam sterilizers (see ANSI/AAMI ST8 and ANSI/AAMI ST55), rigid sterilization container systems (see ANSI/AAMI ST77), or rigid, protective organizing cases that require wrapping prior to sterilization (see ANSI/AAMI ST77);
- b) the use of containment devices for packaging items other than instrument sets or procedural trays;
- c) procedures and techniques for handling and laundering contaminated reusable surgical textiles (see ANSI/AAMI ST65), reusable laboratory items, food service items, and items assigned to a patient for the length of stay (e.g., bedpans, thermometers);
- d) decontamination of hemodialysis machines, hemodialyzers, and hemodialyzer blood tubing (see ANSI/AAMI RD5, ANSI/AAMI RD47, and AAMI RD17, respectively);
- e) the use of dry heat for decontamination purposes or for terminal sterilization of reusable medical devices (see ANSI/AAMI ST40);
- f) guidelines for safe and effective ethylene oxide sterilization (see ANSI/AAMI ST41);
- g) the reprocessing of devices labeled for single use only (see Food and Drug Administration [FDA], 2000c);

NOTE—For more information on the subjects excluded from the scope of this recommended practice, and for additional background information on the inclusions, refer to the references listed in Annex O.