NATIONAL STANDARDS

Recently Published


AAMI Call for Comments

If you would like to comment on one of the draft documents listed below, contact the individual indicated by e-mail to receive a PDF copy of the draft. These copies are free.

Published documents proposed for reaffirmation can be purchased from the AAMI Store.
Comments due March 2, 2020

AAMI/ISO 11140-1, *Sterilization of health care products — Chemical indicators — Part 1: General requirements* (reaffirmation of an American National Standard). Specifies general requirements and test methods for indicators that show exposure to sterilization processes by means of physical and/or chemical change of substances, and which are used to monitor the attainment of one or more of the process parameter(s) specified for a sterilization process. Contact: Amanda Benedict

Comments due March 12, 2020

AAMI AT6, *Autologous transfusion devices* (reaffirmation of an American National Standard). Establishes labeling and performance requirements, test methods, and terminology that will help define a reasonable level of safety and efficacy for autologous transfusion devices. Specifically, it includes requirements for sterile, disposable systems and associated electromechanical hardware designed to collect and filter or process, or both, extravasated blood for reinfusion of erythrocytes or filtered whole blood into the patient’s circulation. Aspects of these systems related to collection, anticoagulation (systemic and device), storage, processing and filtration, and reinfusion are within the scope of this standard. Contact: Cliff Bernier


AAMI BF64, *Leukocyte reduction filters* (reaffirmation of an American National Standard). Contains labeling requirements, performance requirements, test methods, and terminology for disposable filters used for the reduction of leukocytes from blood or blood components. Contact: Cliff Bernier

Comments due March 23, 2020


Comments due April 13, 2020

AAMI CN27, General requirements for Luer activated valves (LAVs) incorporated into medical devices for intravascular applications (new American National Standard). Covers Luer activated valves (LAVs) for intravascular applications, which open and permit access to the fluid conduit when a male Luer connector is inserted. This standard applies only to the valve end of LAVs. This standard applies to LAVs as stand-alone devices or as components of a medical device. Contact: Colleen Elliott.

AAMI/ISO 10993-15, Biological evaluation of medical devices — Part 15: Identification and quantification of degradation products from metals and alloys (new American National Standard). Specifies general requirements for the design of tests for identifying and quantifying degradation products from final metallic medical devices or corresponding material samples finished as ready for clinical use. Applicable only to those degradation products generated by chemical alteration of the final metallic device in an in vitro degradation test. Because of the nature of in vitro tests, the test results approximate the in vivo behaviour of the implant or material. The described chemical methodologies are a means to generate degradation products for further assessments. Applicable to both materials designed to degrade in the body as well as materials that are not intended to degrade. Contact: Colleen Elliott.

AAMI/ISO 10993-18, Biological evaluation of medical devices — Part 18: Chemical characterization of medical device materials within a risk management process (new American National Standard). Specifies a framework for the identification, and if necessary, quantification of constituents of a medical device, allowing the identification of biological hazards and the estimation and control of biological risks from material constituents, using a generally stepwise approach to the chemical characterization which can include one or more of the following: — the identification of its materials of construction (medical device configuration); — the characterization of the materials of construction via the identification and quantification of their chemical constituents (material composition); — the characterization of the medical device for chemical substances that were introduced during manufacturing (e.g. mould release agents, process contaminants, sterilization residues); — the estimation (using laboratory extraction conditions) of the potential of the medical device, or its materials of construction, to release chemical substances under clinical use conditions (extractables); — the measurement of chemical substances released from a medical device under its clinical conditions of use (leachables). Contact: Colleen Elliott.

AAMI/ISO 14155, Clinical investigation of medical devices for human subject – Good clinical practice. Addresses good clinical practice for the design, conduct, recording and reporting of clinical investigations carried out in human subjects to assess the clinical performance or effectiveness and safety of medical devices. For post-market clinical investigations, the principles set forth in this document can be followed as far as relevant, considering the nature of the clinical investigation (see Annex I). This document specifies general requirements intended to — protect the rights, safety and well-being of human subjects, — ensure the scientific conduct of the clinical investigation and the credibility of the clinical investigation results, — define the responsibilities of the sponsor and
principal investigator, and — assist sponsors, investigators, ethics committees, regulatory authorities and other bodies involved in the conformity assessment of medical devices. Contact: Colleen Elliott.


AAMI ST91:202x, *Flexible and semi-rigid endoscope processing in health care facilities* (revision of an American National Standard). Provides guidelines for point of use treatment, transporting, leak-testing (where indicated), cleaning, packaging (where indicated), high-level disinfecting and/or sterilizing, storage, and quality control procedures of flexible gastrointestinal (GI) endoscopes, flexible bronchoscopes, flexible ear, nose, and throat endoscopes, flexible urology endoscopes, and other types of reusable flexible endoscopes used in procedural and surgical settings, and semi-rigid operative endoscopes (e.g., choledochoscopes) used in health care facilities. These guidelines are intended to provide comprehensive information and direction for health care personnel in the processing of these reusable devices and accessories to render them safe for patient use. Contact: Amanda Benedict.

AAMI ST98:202x, *Cleaning validation of health care products -- Requirements for development and validation of a cleaning process for medical devices* (proposed new American National Standard). Provides requirements to validate the cleaning instructions that are provided by the medical device manufacturer for processing medical devices. Contact: Amanda Benedict.

*Comments due April 29, 2020*

AAMI/ISO 15223-1/Ed.4, *Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements* (revision and parallel adoption of an American National Standard). Provides applicable to symbols used in a broad spectrum of medical devices, which are marketed globally and therefore need to meet different regulatory requirements. Contact: Wil Vargas.
New Work

AAMI NS/WG 1, ICP Device Working Group will be reviewing ANSI/AAMI NS28:1988(R2015), *Intracranial pressuring monitoring devices*, to determine whether or not the document requires revision. Contact: Jennifer Moyer

AAMI/CN, Small Bore Connectors Committee is working on the revision of AAMI/ISO 80369-7, *Small-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for intravascular or hypodermic applications*. Contact: Colleen Elliott

AAMI EQ, Medical Equipment Management Committee. The committee is working on the development of AAMI EQ110/Ed.1, *Guidance for health care technology management education programs*. This standard seeks to provide a recommended framework for new or established education programs in health care technology management (HTM). Contact: Patrick Bernat.


AAMI ST/WG 40, Hospital Practices Steam Sterilization. The working group is working on the developments of AAMI TIR109/Ed.1, *External transport of medical devices processed by health care facilities*. This document will provide guidance for health care facilities regarding the transportation of medical devices from one facility to another; includes the safe method of transport for contaminated items and the maintenance of integrity of sterilized items. Contact: Amanda Benedict.

Consensus Body Members Needed

The committees listed below are seeking new members in the indicated stakeholder interest category to participate in the development of their documents. Definitions of the various stakeholder groups are:

**User:** An individual or organizational representative, who purchases, utilizes or receives the materials, products, systems, or services covered in the scope of technical documents developed by AAMI in the delivery of healthcare; individuals in this interest category include clinicians, employees or representatives of Healthcare Delivery Organizations, clinical consultants, patients, etc.

**Industry:** An individual or organizational representative involved in the commercial production, promotion, sale, use or distribution of materials, products, systems, or services covered in the scope of technical documents developed by AAMI; this interest category includes manufacturers, those involved in supply chains, employees of test labs or commercial labs, industry consultants, etc.
Regulatory: An individual or organizational representative involved in the regulation of the materials, products, systems, or services covered in the scope of the technical documents developed by AAMI; this interest category includes those representing federal, state, local, foreign, or other government entities.

General interest: An individual or organizational representative with a general material interest in the materials, products, systems, or services covered in the scope of the technical documents developed by AAMI and who does not fit into any of the preceding categories; this interest category can include noncommercial academicians, noncommercial researchers, patient or consumer advocates, representatives of accrediting organizations, representatives of other organizations, etc.

Please contact the staff person indicated for more information on how to join.

AAMI NS/WG 1, ICP Device Committee. The committee is seeking new members of all interest categories – users, industry, regulatory, and general interest – to participate in the review of ANSI/AAMI NS28:1988(R2015), Intracranial pressuring monitoring devices. Contact: Jennifer Moyer

AAMI/CN, Small Bore Connectors Committee. The committee is seeking user, regulatory and general interest members to participate in the development of AAMI/ISO 80369-7, Small-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for intravascular or hypodermic applications. Contact: Colleen Elliott

AAMI EQ, Medical Equipment Management Committee. The committee is seeking industry, regulatory and general interest members to participate in the development of AAMI EQ110/Ed.1, Guidance for health care technology management education programs. Contact: Patrick Bernat.

AAMI HE, Human Factors Engineering Committee. The committee is seeking user, regulatory and general interest members to participate in the review of ANSI/AAMI HE75:2009/(R)2018 Human factors engineering—design of medical devices. Contact: Patrick Bernat.

AAMI ST/WG 95, Water Quality for Reprocessing Medical Devices. The working group is general interest, user, and regulatory stakeholders to participate in the development of AAMI ST108/Ed.1, Water for the processing of medical devices. Contact: Amanda Benedict.

AAMI ST/WG 40, Hospital Practices Steam Sterilization. The working group is seeking regulatory and general interest stakeholders to participate in the developments of AAMI TIR109/Ed.1, External transport of medical devices processed by health care facilities. Contact: Amanda Benedict.

AAMI/CV, Cardiac valves. The committee is seeking general interest/regulator members to participate in the revision of ISO 5910, Cardiovascular implants and extracorporeal systems — Cardiac valve repair devices. Contact: Cliff Bernier.

AAMI/VP, Vascular Prostheses. The committee is seeking user and general interest/regulator members to participate in the revision of ISO 25539-3, Cardiovascular implants — Endovascular devices — Part 3:

AAMI/RD, Renal Disease and Detoxification Committee. The committee is seeking general interest/regulator members to participate in the revision of the ISO 8637, Extracorporeal systems for blood purification series, including Part 1: Haemodialysers, haemodiafilters, haemofilters and haemoconcentrators; Part 2: Extracorporeal blood circuit for haemodialysers, haemodiafilters and haemofilters; and Part 3: Plasmafilters. Contact: Cliff Bernier

AAMI/CO, Cardiac Occluders. The committee is seeking industry, user, and general interest/regulator members to participate in the development of ISO 22679, Cardiovascular implants — Transcatheter cardiac occluders. Contact: Cliff Bernier

AAMI/BG, Blood/Gas Exchange Device Committee. The committee is seeking industry, user, and general interest/regulator members to participate in the development of the following Cardiovascular implants and artificial organs documents: ISO 18193, Cannulae for extracorporeal circulation; Amendment 1 to ISO 18242:2016 Centrifugal blood pumps for pulsatile pumps; and revision of ISO 7199, Blood-gas exchangers. Contact: Cliff Bernier

AAMI/VI, Cardiovascular absorbable implants. The committee is seeking industry, user, and general interest/regulator members to participate in the revision of ISO/TS 17137:2019, Cardiovascular implants and extracorporeal systems - Cardiovascular absorbable implants. Contact: Cliff Bernier

AAMI/QM-WG03, Symbols and nomenclature for medical devices Working Group. The committee is seeking user, general interest, and regulator members to participate in the revision of ISO 15223-1/Ed.4, Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements. Contact: Wil Vargas

UPCOMING MEETINGS

AAMI Committees and U.S. TAGs
Open meetings of AAMI committees and U.S. Technical Advisory Groups (TAGs) are listed here and at the AAMI website (www.aami.org). Note: If you plan to attend a meeting, please send a brief note to the AAMI Standards Department (standards@aami.org) indicating the name and date of the meeting so that staff can contact you in the event of a last-minute cancellation.

March 2020

AAMI Sterilization Standards Week (open meetings – advance registration required) 16-19 March 2020, AAMI, Arlington, VA, USA. Contact: Amanda Benedict
April 2020

Infusion Device Committee (open meeting). 28-30 April 2020. 9:00 to 17:00 h, Minneapolis, MN.  
Contact: Jennifer Moyer

May 2020

Renal Disease and Detoxification Committee (open meeting). 4 May 2020. 9:00 to 17:00h, AAMI, Arlington, VA, USA. Contact: Cliff Bernier

Transvenous Cardiac Leads Working Group (open meeting). 4-5 May 2020. 9:00 to 17:00 h, San Diego, CA. Contact: Jennifer Moyer

Cardiac Rhythm Management Devices (open meeting). 5 May 2020. 9:00 to 17:00 h, San Diego, CA. Contact: Jennifer Moyer.

June 2020

AAMI Medical Device Particulates Committee (open meeting) 4-5 June 2020. 9:00 to 17:00h, AAMI, Arlington, VA, USA. Contact: Cliff Bernier

October 2020

AAMI Sterilization Standards Week - details to come! (open meetings – advance registration will be required) 12-16 October 2020, AAMI, Arlington, VA, USA. Contact: Amanda Benedict

INTERNATIONAL STANDARDS

Information on draft international standards under ballot can be found in ANSI Standards Action.

International Committee and Working Group Meetings

Call or email the indicated staff person at AAMI for more information about upcoming international standards meetings.

March 2020

IEC/SC 62D/MT 20, Haemodialysis equipment (closed meeting), 2-3 March 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Hae Choe

ISO/TC 150/SC 6/WG 1, Fundamental standard (closed meeting), 2-3 March 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Jennifer Moyer

ISO/TC 198/WG 7, Packaging (closed meeting), 3 March 2020, 07:00 h to 10:00 h, Zoom meeting. Contact: Amanda Benedict
ISO/TC 150/SC 6/JWG 2, Effects of magnetic resonance imaging on active implantable medical devices (closed meeting), 4-6 March 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Jennifer Moyer

ISO/ 198/WG 13, Washer-disinfectors (closed meeting), 5 March 2020, 08:00 h to 11:00 h, Zoom meeting. Contact: Amanda Benedict

ISO/ 198/WG 1, Industrial EO sterilization (closed meeting), 12 March 2020, 08:00 h to 11:00 h, Zoom meeting. Contact: Amanda Benedict

ISO/TC 215, Health informatics and related WGs (closed meetings), 25 March – 3 April 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia. Contact: Joe Lewelling

April 2020

ISO/ 198/WG 13, Washer-disinfectors (closed meeting), 2 April 2020, 08:00 h to 11:00 h, Zoom meeting. Contact: Amanda Benedict

ISO/TC 150/SC 6/WG 5, Implantable neurostimulators (closed meeting), 15-17 April 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia. Contact Jennifer Moyer

IEC/SC 62A/WG 33, CAG - Chairman Advisory Group (closed meeting), 23-24 April 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia. Contact: Hae Choe

IEC/SC 62A/WG 20, Environmental protection (closed meeting), 27-28 April 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia. Contact: Hae Choe

ISO/ 198/WG 13, Washer-disinfectors (closed meeting), 30 April 2020, 08:00 h to 11:00 h, Zoom meeting. Contact: Amanda Benedict

May 2020

IEC/SC 62D - ISO/TC 121/SC 3/JWG 7, Non-invasive sphygmomanometers (closed meeting), 11-15 May 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Hae Choe

ISO/TC 121, Anaesthetic and respiratory equipment and affiliated groups (closed meetings), 18-22 May 2020, 9:00h to 17:00h, BSI, London, UK. Contact: Colleen Elliott

ISO/TC 194, Biological and clinical evaluation of medical devices and affiliated groups (closed meetings), 25-29 May 2020, 9:00h to 17:00h, Qingdao, China. Contact: Colleen Elliott CANCELED

June 2020

ISO/TC 198/WG 11, General criteria for sterilization processes and sterilizing equipment (closed meeting), 3-4 June 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Amanda Benedict

IEC/SC 62D - ISO/TC 173/JWG 4, Medical beds (closed meeting), 8-11 June 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Hae Choe
ISO/TC 198/WG 8, Microbiological methods (closed meeting), 15-16 June 2020, 9:00 h to 17:00 h, DIN, Berlin, Germany. Contact: Amanda Benedict

ISO/TC 150/SC 6/JWG 2, Effects of magnetic resonance imaging on active implantable medical devices (closed meeting), 17-19 June 2020, 9:00 h to 17:00 h, AAMI, Arlington, Virginia, USA. Contact: Jennifer Moyer

MISCELLANEOUS

2020 INTERNATIONAL CONFERENCE ON MEDICAL DEVICE STANDARDS AND REGULATION
April 20–23, 2020 • Arlington, VA
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