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 October 5, 2020

AAMI/ISO 80369-7, Small-bore connectors for liquids and gases in healthcare applications – Part 7: Connectors for intravascular or hypodermic applications (revision of an American National Standard). Specifies dimensions and requirements for the design and functional performance of small-bore connectors intended to be used for connections in intravascular applications or hypodermic connections in hypodermic applications of medical devices and accessories. Contact: celliott@aami.org

Comments due October 26, 2020

AAMI PC76, Active implantable medical devices – Requirements and test protocols for safety of patients with pacemakers and ICDs exposed to magnetic resonance imaging (proposed new American National Standard). Provide requirements and test protocols for implantable pacemakers and ICDs exposed to magnetic resonance imaging. Physicians are increasingly using magnetic resonance imaging as tool for differential diagnostic, thus exposing pacemakers and ICD patients to such equipment. Current product standards for implantable pacemakers and ICDs do not include requirements and test protocols for implantable pacemakers and ICDs, which would ensure patient safety during such procedures. Contact: jmoyer@aami.org

Comments due November 9, 2020

AAMI HIT1000-1, AAMI HIT1000-1/Ed 1, Health IT Software and Systems – Part 1: Fundamental concepts and principles (proposed new American National Standard from a provisional standard). Identifies the fundamental concepts and principles needed to maintain safe, secure and effective health IT software and systems. Identifies the roles, and defines the responsibilities, activities and best practices that are necessary for managing safety, security and effectiveness of health IT software and systems. Applies throughout the whole lifecycle of health IT software and systems and to all sizes and type of actors involved with that system. Contact: wvargas@aami.org

Comments due November 16, 2020

AAMI/IEC 60601-2-20, Medical electrical equipment - Part 2-20: Particular requirements for the basic safety and essential performance of infant transport incubators (proposed revised American National Standard). Applies to the basic safety and essential performance of transport incubators. This standard does not apply to heating devices intended for physiotherapy, baby incubators, radiant warmers. Contact: jmoyer@aami.org

New Work

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: Amanda Benedict.

AAMI QM-WG01, Application of Quality Systems to Medical Devices Working Group. The working group is working on the development of AAMI CR510, Consensus Report on Compliant Use of Cloud Computing for Quality Systems and Medical Devices. This Consensus Report will provide guidance to
multiple stakeholders regarding the appropriate and compliant use of cloud computing both as a component of medical devices and in support of quality systems. Contact: Wil Vargas.

AAMI II, Infant Incubators. The committee will be working on the identical expedited adoption of the following documents:


- AAMI/IEC 60601-2-50, Medical electrical equipment - Part 2-50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment. Applies to the basic safety and essential performance of infant phototherapy equipment.

To get involved in the development of these documents, please contact Jennifer Moyer.

**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 PC/WG 3, Pacemaker & MRI Compatibility Working Group. The working group is seeking user, regulatory, and general interest members to participate in the development of AAMI PC76, Active implantable medical devices – Requirements and test protocols for safety of patients with pacemakers and ICDs exposed to magnetic resonance imaging. 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: Wil Vargas.

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/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/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/HIT, Health IT Committee. The working group is seeking industry, general interest, and regulator members to participate in the development of AAMI HIT1000-1/Ed., *Health IT Software and Systems — Part 1: Fundamental concepts and principles*. Contact: Wil Vargas

AAMI/HIT-WG02, Health IT Quality Systems Working Group. The working group is seeking industry, general interest, and regulator members to participate in the development of AAMI HIT1000-2/Ed., *Health IT software and systems — Part 2: Application of quality systems principles and practices (Provisional Standard)*. Contact: Wil Vargas

AAMI/II, Infant Incubators. The committee is seeking users, regulatory, and general interest members to participate in the development of AAMI/IEC 60601-2-20, *Medical electrical equipment - Part 2-20: Particular requirements for the basic safety and essential performance of infant transport incubators*. Contact: Jennifer Moyer

**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.
October 2020

AAMI Sterilization Standards Week - details below (open web meetings). 13-16 and 26-29 October 2020, virtual meetings.  
Contact: Amanda Benedict, sterilization@aami.org.

October 2020 Sterilization Standards Opening Plenary (open web meeting). 13 October 2020, 09:00h to 11:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

October 2020 Sterilization Standards New Participant Orientation (open web meeting). 13 October 2020, 11:00h to 12:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Reusable surgical textiles processing Working Group (open web meeting). 13 October 2020, 13:00h to 17:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Assurance of Sterility Working Group (open web meeting). 14 October 2020, 08:00h to 12:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Industrial EO sterilization Working Group (open web meeting). 14 October 2020, 13:00h to 17:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Microbiological methods Working Group (open web meeting). 15 October 2020, 09:00h to 12:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Water quality for reprocessing medical devices Working Group (open web meeting). 15 October 2020, 13:00h to 17:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Compatibility of materials subject to sterilization Working Group (open web meeting). 15 October 2020, 09:00h to 12:00h ET. Virtual meeting.  
Contact: Cliff Bernier, sterilization@aami.org.

Quality systems for device processing Working Group (open web meeting). 26 October 2020, 14:00h to 16:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Endoscope reprocessing Working Group (open web meeting). 27 October 2020, 09:00h to 12:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

Chemical sterilants hospital practices Working Group (open web meeting). 28 October 2020, 13:00h to 17:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.

AAMI October 2020 Sterilization Standards Closing Plenary and US TAG to ISO/TC 198 Web Meeting (open web meeting). 29 October 2020, 09:00h to 11:00h ET. Virtual meeting.  
Contact: Amanda Benedict, sterilization@aami.org.
November 2020

Vaporized hydrogen peroxide sterilization Working Group (open web meeting). 10 November 2020, 11:00h to 14:00h ET. Virtual meeting. Contact: Amanda Benedict.

Cardiac Rhythm Management Devices (open remote meeting). 12 November 2020, 11:00 to 13:30 h. Contact: Jennifer Moyer

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.

October 2020

ISO/TC 150/SC 6/JWG 1, Cardiac pacemakers and implantable defibrillators (closed meeting). 1 October 2020, 10:00 to 12:00 h ET, Zoom meeting. Contact: Jennifer Moyer

ISO/TC 198/WG 8, Microbiological methods (closed meeting), 5 October 2020, 09:00 h to 11:00 h ET, Zoom meeting. Contact: Amanda Benedict

ISO/TC 198/WG 8, Microbiological methods (closed meeting), 22 October 2020, 09:00 h to 11:00 h ET, Zoom meeting. Contact: Amanda Benedict

ISO/TC 150/SC 6, Active implants (closed meeting), 22 October 2020, 11:00 to 13:00 h ET, Zoom meeting. Contact: Jennifer Moyer