

ANSI/AAMI/ UL 2800-1: 2022

Standard for Medical Device Interoperability





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Abstract: Specifies a baseline set of requirements for assuring safe and secure interoperability for

Interoperable Medical Systems.

Keywords: interoperability requirements, medical systems, medical devices, interoperable systems

Commitment for Amendments

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Committee representation

Association for the Advancement of Medical Instrumentation

AAMI/UL Joint Committee for Medical Device Interoperability, JC 2800

The publication of AAMI/UL 2800-1:2022 as a new American National Standard was initiated by the AAMI/UL Joint Committee for Medical Device Interoperability, JC 2800.

This list represents the membership at the time the Committee balloted on the final text of this edition. Since that time, changes in the membership may have occurred.

Cochairs: Diana P. Jordan

Ovidiu Munteanu

Members: Dave Arney, CIMIT (MGH Anesthesia & Biomedical Engineering)

Oliver Christ, Prosystem AG

R Cooper, Eurofins E&E North America

Holly Drake, Dexcom Inc. Sherman Eagles, SoftwareCPR Scott Eaton, Mindray DS USA Inc

Kenneth Fuchs, Draeger Medical Systems Inc. Julian Goldman, Massachusetts General Hospital

Pamela K. Gwynn, UL LLC

John Hatcliff, Kansas State University Jacob Johnson, Kaiser Permanente

Diana Pappas Jordan (JC Cochair), Underwriters Laboratories Inc.

Edmund Kienast, National E-Health Transition Authority (NEHTA)-Australia

Todd Konieczny, Intertek Testing Services

Patty Krantz, Medtronic Inc.

Insup Lee, University of Pennsylvania Marina Lee, Staubli Electrical Connectors, Inc.

Ovidiu Munteanu JC Cochair), AAMI Steve Nichols, GE Healthcare

Geetha Rao, Springborne Life Sciences

Tracey Rausch, DocBox Inc. Daniel Rubery, NxStage Medical, Inc.

Patricia A. Sena (JC Project Manager), Underwriters Laboratories Inc. Elliot Sloane, Center for Healthcare Information Research & Policy

Erin Sparnon, ECRI

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Standard for Medical Device Interoperability

1 Introduction

- 1.1 The AAMI/UL 2800 series of standards covers the interoperability of medical products. AAMI/UL 2800-1 is the general standard that specifies a baseline set of requirements for assuring safe and secure interoperability for interoperable medical systems. The requirements in the AAMI/UL 2800-1 standard are supplemented by the requirements in additional AAMI/UL 2800 standards. These additional standards are intended to be used in conjunction with the general standard and applied as needed. While this introduction applies to all of the AAMI/UL 2800 series of standards, the scope section of each additional standard describes what is covered by that standard.
- 1.2 Multiple stakeholders may participate in the development, deployment, assembly, and operation of a medical system with interoperable elements. Such a system, referred to as an interoperable medical system, should minimize patient risks, maintain clinical effectiveness, ensure timely and adequate access to data while protecting its security, and enable adequate provision of care. In order to facilitate alignment of stakeholders around these aims, the AAMI/UL 2800 series of standards establishes a baseline set of requirements for assuring safe and secure interoperability.
- 1.3 Each stakeholder will need to determine the specific level and manner in which interoperability will be specified and assured for its interoperable medical products. However, a specific system may be developed, assembled, deployed, and operated through a range of processes undertaken by multiple stakeholders. Specific activities in these processes assure interoperability. In order for stakeholders to collectively accomplish this, the processes need to be linked effectively.
- 1.4 Effective linkage of processes across multiple stakeholders is a core focus of the AAMI/UL 2800 series of standards. This first requires that each stakeholder adequately assesses and manages safety, security and essential performance vulnerabilities of its interoperable medical products. Secondly, it requires that each stakeholder understands and conforms with interoperability aspects of disclosed specifications of an interoperable medical product which it acquires or with which it interoperates, including the consequent safety and security characteristics. Finally, it requires that each stakeholder clearly communicates to the other stakeholders the information required to assure interoperability.
- 1.5 The requirements in the AAMI/UL 2800 series of standards are intended to apply to medical devices, as well as other connected infrastructure elements, and interoperable medical systems constructed from these. The AAMI/UL 2800 series of standards is intended to be used by individual stakeholders.
- 1.6 The AAMI/UL 2800 series of standards employ a lifecycle process approach to organizing requirements. In addition to a set of broad management functions, the standards provide for a set of interoperability planning, realization, deployment, and monitoring activities. These activities also incorporate cross-cutting requirements for security and risk management. The standards recognize that a given organization may be responsible for only a part of the full range of activities required for an interoperable medical system. Furthermore, the organization's interoperable medical products may provide only a specific or limited functionality. To accommodate this, the standards provide for flexibility in the scope, sequence, and interaction of these activities. Finally, the standards provide requirements and supplementary guidance on key clinical and engineering properties of an interoperable medical system that are essential to assuring safe and secure interoperability and provide guidance on lifecycle activities.