

**SLOVENSKI STANDARD**  
**oSIST prEN ISO 11073-10700:2024**  
**01-junij-2024**

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**Zdravstvena informatika - Interoperabilnost naprav - 10700. del: Komunikacija medicinskih naprav na mestu oskrbe - Standard za osnovne zahteve za udeležence v storitvi - Sistem usmerjenega povezovanja naprav (SDC) (ISO/IEEE FDIS 11073-10701:2024)**

Health informatics - Device interoperability - Part 10700: Point-of-Care Medical Device Communication - Standard for Base Requirements for Participants in a Service-Oriented Device Connectivity (SDC) System (ISO/IEEE FDIS 11073-10700:2024)

Medizinische Informatik - Interoperabilität von Geräten - Teil 10700: Kommunikation persönlicher Gesundheitsgeräte - Standard für Basisanforderungen für Teilnehmer an einem serviceorientierten Gerätekonnektivitätssystem (SDC) (ISO/IEEE FDIS 11073-10700:2024)

Informatique de santé - Interopérabilité des dispositifs - Partie 10700: Titre manque (ISO/IEEE FDIS 11073-10700:2024)

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# FINAL DRAFT International Standard

## ISO/IEEE FDIS 11073-10700

### Health informatics — Device interoperability —

Part 10700:

### Point-of-Care Medical Device Communication –Standard for Base Requirements for Participants in a Service-Oriented Device Connectivity (SDC) System

ISO/TC 215

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Institute of Electrical and Electronics Engineers, Inc  
3 Park Avenue, New York  
NY 10016-5997, USA

Email: [stds.ipr@ieee.org](mailto:stds.ipr@ieee.org)  
Website: [www.ieee.org](http://www.ieee.org)

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ISO/IEEE 11073-10700 was prepared by the IEEE 11073 Standards Committee of the IEEE Engineering in Medicine and Biology Society (as IEEE Std 11073-10700) and drafted in accordance with its editorial rules. It was adopted, under the “fast-track procedure” defined in the Partner Standards Development Organization cooperation agreement between ISO and IEEE, by Technical Committee ISO/TC 215, *Health informatics*.

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Any feedback or questions on this document should be directed to the user’s national standards body. A complete listing of these bodies can be found at [www.iso.org/members.html](http://www.iso.org/members.html). **Abstract:** Medical devices that offer a communication interface as specified by the IEEE 11073 Service-oriented Device Connectivity (SDC) standards can be integrated into a health IT system to jointly execute system functions. However, implementing the IEEE 11073 SDC communication protocol is not sufficient to demonstrate safety, effectiveness, and security of system functions resulting from the combination of system function contributions from two or more medical devices. SDC participant key purposes (PKPs) are sets of requirements that allow for manufacturers to have certain expectations about BICEPS participants from other manufacturers. This common understanding enables the manufacturers to perform risk management, verification, validation, and usability engineering for the safe use of system functions. This standard specifies requirements for the allocation of responsibilities to SDC base participants.

**Keywords:** base PKP; BICEPS; communication protocol specification; documentation and process responsibilities; dynamic medical device interoperability; IEEE 11073-10700™; integrated clinical environment; participant key purpose; point-of-care medical device communication; risk management; SDC; service-oriented device connectivity; safety, effectiveness, and security; system function; system function contribution; usability engineering

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At the time this IEEE standard was completed, the Point-of-Care Devices Working Group had the following membership:

**Stefan Schlichting, Chair**  
**Martin Kasparick, Subgroup Chair**

Björn Andersen	Christian Haye	Erik Moll
Fabian Baumeister	Werner Hoelzl	Karen Moniz
Jin-Woo Choi	Martin Hurrell	Jody Paul
Malcolm Clarke	Andy Iverson	Craig Reister
Paul Close	Jennifer Jacobs	John Rhoads
Todd Cooper	Sven Kämmer	Sean Rocke
Sandra Costanzo	Anton Keller	Martin Rosner
Steven Dain	Tobias Klotz	Enrico Rudolf
Kurt Eliason	Satoshi Kobayashi	Gilani Sadeghi
Javier Espina	Anil Kochhar	Paul Schluter
Michael Faughn	Peter Kranich	Elliot Silver
Ken Fuchs	Ray Krasinski	Tulasi Sivanesan
John Garguilo	Jithin Krishnan	Isabel Tejero
Frank Golatowski	Sungkee Lee	James Vollmer
David Gregoczyk	Konstantinos Makrodimitis	Brian Witkowski
Steve Griffiths	Koichiro Matsumoto	Ravi Sekhar Yarrabothu
Peter Gunter	Jörg-Uwe Meyer	Greg Zeller
Bob Harbort	Madhu Mohan	Daidi Zhong

The following members of the individual Standards Association balloting group voted on this standard. Balloters may have voted for approval, disapproval, or abstention.

Bjoern Andersen	David Fuschi	Javier Luiso
Pradeep Balachandran	Frank Golatowski	H. Moll
Fabian Baumeister	David Gregorczyk	Bansi Patel
Lyle Bullock	Werner Hoelzl	Dalibor Pokrajac
Pin Chang	Piotr Karocki	John Rhoads
Diego Chiozzi	Martin Kasparick	Elie Sarraf
Todd Cooper	Stuart Kerry	Stefan Schlichting
Houde Dai	Edmund Kienast	Walter Struppler
Kurt Eliason	Yongbum Kim	Maria Isabel Tejero del Rio
Michael Faughn	Raymond Krasinski	John Vergis
Immanuel Freedman	Ting Li	Yu Yuan
Kenneth Fuchs		Oren Yuen

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**Ted Burse, Vice Chair**  
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Edward A. Addy	Johnny Daozhuang Lin	Mark Siira
Ramy Ahmed Fathy	Kevin Lu	Dorothy V. Stanley
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Guido R. Hiertz	Andrew Myles	F. Keith Waters
Yousef Kimiagar	Damir Novosel	Karl Weber
Joseph L. Koepfinger*	Annette D. Reilly	Sha Wei
Thomas Koshy	Robby Robson	Philip B. Winston
John D. Kulick	Jon Walter Rosdahl	Daidi Zhong

\*Member Emeritus

## ISO/IEEE 11073-10700:2024(en)

### Introduction

This introduction is not part of IEEE Std 11073-10700-2022, Health Informatics—Device Interoperability—Part 10700: Point-of-Care Medical Device Communication—Standard for Base Requirements for Participants in a Service-Oriented Device Connectivity (SDC) System.

The IEEE 11073 Point-of-Care Medical Device Communication Standards enable communication between health IT elements in a HEALTH IT SYSTEM including MEDICAL DEVICES. They provide automatic and detailed electronic data capture of patient vital signs information and device operational data. The primary goals are to:

- Provide real-time plug-and-play interoperability for MEDICAL DEVICES. “Real-time” means that data from multiple MEDICAL DEVICES can be retrieved, temporally correlated, displayed, and processed in fractions of a second. “Plug-and-play” means that there are no recurring configuration steps necessary to enable data exchange between MEDICAL DEVICES.
- Facilitate the efficient and effective exchange of vital signs and MEDICAL DEVICE data acquired at the PoC in all health care environments. “Efficient and effective exchange of MEDICAL DEVICE data” means that data captured at the PoC, e.g., patient vital signs, can be received, parsed, and interpreted by different types of applications without the loss of safety-critical information.

The IEEE 11073 Point-of-Care Medical Device Communication Standards are targeted at surgical as well as acute and continuous care devices, such as patient monitors, ventilators, infusion pumps, ECG devices, endoscopic camera systems, insufflators, dissectors, etc. They build a family of standards that can be bound to one another to provide optimized connectivity for devices at the PoC.

Within the context of the ISO/IEEE 11073 family of standards for Point-of-Care Medical Device Communication, this standard defines the requirements for SDC BASE PARTICIPANTS in an SDC SYSTEM that comprises an IT NETWORK of MEDICAL DEVICES to enable safe and secure contribution to SYSTEM FUNCTIONS.

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ISO 20417:2021, Sections 3.2 and 3.11  
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 ISO 14971:2019, Section 3.18

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# Health Informatics—Device Interoperability

## Part 10700: Point-of-Care Medical Device Communication— Standard for Base Requirements for Participants in a Service- Oriented Device Connectivity (SDC) System

### 1. Overview

#### 1.1 Scope

This standard specifies the base set of Participant Key Purposes (PKPs) for the Service-oriented Device Connectivity (SDC) series of standards. PKPs are role-based sets of requirements for products in order to support safe, effective, and secure interoperability in medical IT networks at point-of-care environments such as the intensive care unit (ICU), operating room (OR) or other acute care settings. This standard specifies both product development process and technical requirements.

#### 1.2 Word usage

The word *shall* indicates mandatory requirements strictly to be followed in order to conform to the standard and from which no deviation is permitted (*shall equals is required to*).<sup>6-7</sup>

The word *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 (*should equals is recommended that*).

The word *may* is used to indicate a course of action permissible within the limits of the standard (*may equals is permitted to*).

The word *can* is used for statements of possibility and capability, whether material, physical, or causal (*can equals is able to*).

#### 1.3 Service-oriented Device Connectivity standards

The SDC STANDARDS are a subset of the IEEE 11073 standards and define requirements for MEDICAL DEVICES and other participants that exchange physiological or technical information or enable external control while being operated in an IT NETWORK.

The SDC STANDARDS comprise the specification of a domain and message model (IEEE Std 11073-10207) and transport technology (IEEE Std 11073-20702) that form a service-oriented MEDICAL DEVICE architecture (IEEE Std 11073-20701).<sup>8</sup> These SDC core standards constitute the technical building blocks for foundational, structural, and semantic MEDICAL DEVICE interoperability over secure data transmission. The SDC PKP STANDARDS (see 1.4) and particular SDC Device Specializations address additional levels.

#### 1.4 Participant key purposes

MEDICAL DEVICES that offer a communication interface as specified by the SDC STANDARDS can be integrated into a HEALTH IT SYSTEM on behalf of the SYSTEM OWNER, establishing an SDC SYSTEM to be used by the HEALTHCARE DELIVERY ORGANIZATION.

The SYSTEM FUNCTIONs made available in an SDC SYSTEM depend on the individual SYSTEM FUNCTION CONTRIBUTIONs of its BICEPS PARTICIPANTs. Accordingly, the MANUFACTURER of a BICEPS SERVICE

<sup>6</sup> The use of the word *must* is deprecated and cannot be used when stating mandatory requirements; *must* is used only to describe unavoidable situations.

<sup>7</sup> The use of *will* is deprecated and cannot be used when stating mandatory requirements; *will* is only used in statements of fact.

<sup>8</sup> Information on references can be found in Clause 2.

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PROVIDER can only specify its INTENDED SYSTEM FUNCTION CONTRIBUTIONS, whereas the MANUFACTURER of a BICEPS SERVICE CONSUMER can specify the intended SYSTEM FUNCTIONS as well as the SYSTEM FUNCTION CONTRIBUTIONS required from BICEPS SERVICE PROVIDERS in the SDC SYSTEM.

But to verify the safety, effectiveness, and security of these SYSTEM FUNCTIONS, only implementing the communication protocol based on the SDC STANDARDS is not sufficient. The safety, effectiveness, and security of the SDC SYSTEM is based on allocating responsibilities to the individual BICEPS PARTICIPANTS according to the requirements of the SDC PARTICIPANT KEY PURPOSES (PKPs) they assume.

The responsibility for the individual products as BICEPS PARTICIPANTS in an SDC SYSTEM remains with the MANUFACTURERS whereas the SYSTEM OWNER is responsible for integration of the products into a HEALTH IT SYSTEM and the ADMINISTRATOR is responsible for operation and maintenance of the HEALTH IT SYSTEM (see ISO 81001-1:2021, Clause 4.5 [B16]).<sup>9</sup> In addition, the SYSTEM OWNER and ADMINISTRATOR take the responsibilities placed on them by declarations in the ACCOMPANYING INFORMATION of the individual products that are to be integrated, e.g., pertaining to configuration, NETWORK BANDWIDTH, etc.

The SDC PKP STANDARDS specify the allocation of responsibilities and allow for MANUFACTURERS to have certain expectations about BICEPS PARTICIPANTS from other MANUFACTURERS. Conformity to SDC PKP STANDARDS and indication of this conformity creates confidence in these expectations and enables MANUFACTURERS to take the responsibilities for SYSTEM FUNCTION CONTRIBUTIONS of their BICEPS PARTICIPANTS in an SDC SYSTEM. These responsibilities pertain to technical design, implementation, verification, validation, RISK MANAGEMENT, USABILITY ENGINEERING, and labeling of BICEPS PARTICIPANTS.

This standard defines the SDC BASE PROVIDER and the SDC BASE CONSUMER PKPs. They comprise the base requirements for MANUFACTURERS to support safe, effective, and secure operation of their SDC BASE PARTICIPANTS in an SDC SYSTEM.

MANUFACTURERS of SDC BASE PROVIDERS can assess and specify which requirements need to be fulfilled by SDC BASE CONSUMERS for the safe use of SYSTEM FUNCTION CONTRIBUTIONS. Based on conformity of SDC BASE CONSUMERS to this and other SDC PKP STANDARDS, SDC BASE PROVIDERS can restrict access to BICEPS SERVICES in the HEALTH IT SYSTEM.

For exchanging metric data, ALERT information, and external control commands, conformity with further SDC PKP STANDARDS is recommended. Requirements that relate to specific SYSTEM FUNCTIONS or SYSTEM FUNCTION CONTRIBUTIONS can be specified in additional SDC PARTICIPANT KEY PURPOSES.

## 2. Normative references

The following referenced documents are indispensable for the application of this document (i.e., they must be understood and used, so each referenced document is cited in text and its relationship to this document is explained). For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments or corrigenda) applies.

IEEE Std 11073-10101<sup>TM</sup>, Health informatics—Point-of-care medical device communication—Part 10101: Nomenclature.<sup>10, 11</sup>

IEEE Std 11073-10207<sup>TM</sup>, Health informatics—Point-of-care medical device communication—Part 10207: Domain Information and Service Model for Service-Oriented Point-of-Care Medical Device Communication.

IEEE Std 11073-20701<sup>TM</sup>, Health informatics—Point-of-care medical device communication—Part 20701: Service-Oriented Medical Device Exchange Architecture and Protocol Binding.

<sup>9</sup> The numbers in brackets correspond to those of the bibliography in Annex D.

<sup>10</sup> The IEEE standards or products referred to in this annex are trademarks owned by The Institute of Electrical and Electronics Engineers, Incorporated.

<sup>11</sup> IEEE publications are available from The Institute of Electrical and Electronics Engineers (<https://standards.ieee.org/>).