

STN	Zdravotnícka informatika. Osobné komunikačné zdravotné zariadenie. Časť 10424: Špecializácia zariadenia. Dýchacia terapeutická pomôcka pri spánkovom apnoe (SABTE) (ISO/IEEE 11073-10424: 2016).	STN EN ISO 11073-10424 84 8037
------------	---	--

Health informatics - Personal health device communication - Part 10424: Device specialization - Sleep apnoea breathing therapy equipment (SABTE) (ISO/IEEE 11073-10424:2016)

Táto norma obsahuje anglickú verziu európskej normy.
This standard includes the English version of the European Standard.

Táto norma bola oznámená vo Vestníku ÚNMS SR č. 12/16

Obsahuje: EN ISO 11073-10424:2016, ISO/IEEE 11073-10424:2016

123620

Úrad pre normalizáciu, metrológiu a skúšobníctvo SR, 2017
Podľa zákona č. 264/1999 Z. z. v znení neskorších predpisov sa môžu slovenské technické normy rozmnožovať a rozširovať iba so súhlasom Úradu pre normalizáciu, metrológiu a skúšobníctvo SR.

EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN ISO 11073-10424

June 2016

ICS 35.240.80

English Version

**Health informatics - Personal health device
communication - Part 10424: Device specialization - Sleep
apnoea breathing therapy equipment (SABTE) (ISO/IEEE
11073-10424:2016)**

Informatique de la santé - Communication entre
dispositifs de santé personnels - Partie 10424:
Spécialisation de dispositif - Équipement de thérapie
respiratoire de l'apnée du sommeil (SABTE) (ISO/IEEE
11073-10424:2016)

Medizinische Informatik - Kommunikation von Geräten
für die persönliche Gesundheit - Teil 10424:
Gerätespezifikation - Schlafapnoe-Atemtherapiegeräte
(ISO/IEEE 11073-10424:2016)

This European Standard was approved by CEN on 21 February 2016.

CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and United Kingdom.



EUROPEAN COMMITTEE FOR STANDARDIZATION
COMITÉ EUROPÉEN DE NORMALISATION
EUROPÄISCHES KOMITEE FÜR NORMUNG

CEN-CENELEC Management Centre: Avenue Marnix 17, B-1000 Brussels

Contents	Page
European foreword.....	3

European foreword

This document (EN ISO 11073-10424:2016) has been prepared by Technical Committee ISO/TC 215 “Health informatics” in collaboration with Technical Committee CEN/TC 251 “Health informatics” the secretariat of which is held by NEN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by December 2016, and conflicting national standards shall be withdrawn at the latest by December 2016.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN [and/or CENELEC] shall not be held responsible for identifying any or all such patent rights.

According to the CEN-CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdom.

Endorsement notice

The text of ISO/IEEE 11073-10424:2016 has been approved by CEN as EN ISO 11073-10424:2016 without any modification.

**Health informatics — Personal health
device communication —**

Part 10424:

**Device Specialization — Sleep Apnoea
Breathing Therapy Equipment (SABTE)**

*Informatique de santé — Communication entre dispositifs de santé
personnels*

*Partie 10424: Spécialisation de dispositif — Équipement de thérapie
respiratoire de l'apnée du sommeil (SABTE)*





COPYRIGHT PROTECTED DOCUMENT

© IEEE 2014

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO or IEEE at the respective address below.

ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

Institute of Electrical and Electronics Engineers, Inc.
3 Park Avenue, New York • NY 10016-5997, USA
E-mail stds.ipr@ieee.org
Web www.ieee.org

Published in Switzerland

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

IEEE Standards documents are developed within the IEEE Societies and the Standards Coordinating Committees of the IEEE Standards Association (IEEE-SA) Standards Board. The IEEE develops its standards through a consensus development process, approved by the American National Standards Institute, which brings together volunteers representing varied viewpoints and interests to achieve the final product. Volunteers are not necessarily members of the Institute and serve without compensation. While the IEEE administers the process and establishes rules to promote fairness in the consensus development process, the IEEE does not independently evaluate, test, or verify the accuracy of any of the information contained in its standards.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

Attention is called to the possibility that implementation of this standard may require the use of subject matter covered by patent rights. By publication of this standard, no position is taken with respect to the existence or validity of any patent rights in connection therewith. ISO/IEEE is not responsible for identifying essential patents or patent claims for which a license may be required, for conducting inquiries into the legal validity or scope of patents or patent claims or determining whether any licensing terms or conditions provided in connection with submission of a Letter of Assurance or a Patent Statement and Licensing Declaration Form, if any, or in any licensing agreements are reasonable or non-discriminatory. Users of this standard are expressly advised that determination of the validity of any patent rights, and the risk of infringement of such rights, is entirely their own responsibility. Further information may be obtained from ISO or the IEEE Standards Association.

ISO/IEEE 11073-10424 was prepared by the IEEE 11073 Standards Committee of the IEEE Engineering in Medicine and Biology Society (as IEEE Std 11073-10424-2014). It was adopted by Technical Committee ISO/TC 215, *Health informatics*, in parallel with its approval by the ISO member bodies, under the “fast-track procedure” defined in the Partner Standards Development Organization cooperation agreement between ISO and IEEE. IEEE is responsible for the maintenance of this document with participation and input from ISO member bodies.

Abstract: Within the context of the ISO/IEEE 11073 family of standards for device communication, a normative definition of the communication between sleep apnoea breathing therapy equipment (SABTE) devices (agents) and managers (e.g., cell phones, personal computers, personal health appliances, set top boxes), in a manner that enables plug-and-play interoperability, is established in this standard. It leverages appropriate portions of existing standards including ISO/IEEE 11073 terminology, information models, application profile standards, and transport standards. It specifies the use of specific term codes, formats, and behaviors in telehealth environments restricting optionality in base frameworks in favor of interoperability. This standard defines a common core of communication functionality for SABTE. In this context, SABTE is defined as a device that is intended to alleviate the symptoms of a patient who suffers from sleep apnoea by delivering a therapeutic breathing pressure to the patient. SABTE is primarily used in the home health-care environment by a lay operator without direct professional supervision.

Keywords: IEEE 11073-10424™, medical device communication, personal health devices, sleep apnoea breathing therapy equipment (SABTE)

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

Copyright © 2014 by The Institute of Electrical and Electronics Engineers, Inc.
All rights reserved. Published 29 September 2014. Printed in the United States of America.

IEEE is a registered trademark in the U.S. Patent & Trademark Office, owned by The Institute of Electrical and Electronics Engineers, Incorporated.

PDF: ISBN 978-0-7381-9316-8 STD98794
Print: ISBN 978-0-7381-9317-5 STDPD98794

IEEE prohibits discrimination, harassment, and bullying.

For more information, visit <http://www.ieee.org/web/aboutus/whatis/policies/p9-26.html>.

No part of this publication may be reproduced in any form, in an electronic retrieval system or otherwise, without the prior written permission of the publisher.

Important Notices and Disclaimers Concerning IEEE Standards Documents

IEEE documents are made available for use subject to important notices and legal disclaimers. These notices and disclaimers, or a reference to this page, appear in all standards and may be found under the heading “Important Notice” or “Important Notices and Disclaimers Concerning IEEE Standards Documents.”

Notice and Disclaimer of Liability Concerning the Use of IEEE Standards Documents

IEEE Standards documents (standards, recommended practices, and guides), both full-use and trial-use, are developed within IEEE Societies and the Standards Coordinating Committees of the IEEE Standards Association (“IEEE-SA”) Standards Board. IEEE (“the Institute”) develops its standards through a consensus development process, approved by the American National Standards Institute (“ANSI”), which brings together volunteers representing varied viewpoints and interests to achieve the final product. Volunteers are not necessarily members of the Institute and participate without compensation from IEEE. While IEEE administers the process and establishes rules to promote fairness in the consensus development process, IEEE does not independently evaluate, test, or verify the accuracy of any of the information or the soundness of any judgments contained in its standards.

IEEE does not warrant or represent the accuracy or content of the material contained in its standards, and expressly disclaims all warranties (express, implied and statutory) not included in this or any other document relating to the standard, including, but not limited to, the warranties of: merchantability; fitness for a particular purpose; non-infringement; and quality, accuracy, effectiveness, currency, or completeness of material. In addition, IEEE disclaims any and all conditions relating to: results; and workmanlike effort. IEEE standards documents are supplied “AS IS” and “WITH ALL FAULTS.”

Use of an IEEE standard is wholly voluntary. The existence of an IEEE standard does not imply that there are no other ways to produce, test, measure, purchase, market, or provide other goods and services related to the scope of the IEEE standard. Furthermore, the viewpoint expressed at the time a standard is approved and issued is subject to change brought about through developments in the state of the art and comments received from users of the standard.

In publishing and making its standards available, IEEE is not suggesting or rendering professional or other services for, or on behalf of, any person or entity nor is IEEE undertaking to perform any duty owed by any other person or entity to another. Any person utilizing any IEEE Standards document, should rely upon his or her own independent judgment in the exercise of reasonable care in any given circumstances or, as appropriate, seek the advice of a competent professional in determining the appropriateness of a given IEEE standard.

IN NO EVENT SHALL IEEE BE LIABLE FOR ANY DIRECT, INDIRECT, INCIDENTAL, SPECIAL, EXEMPLARY, OR CONSEQUENTIAL DAMAGES (INCLUDING, BUT NOT LIMITED TO: PROCUREMENT OF SUBSTITUTE GOODS OR SERVICES; LOSS OF USE, DATA, OR PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY OF LIABILITY, WHETHER IN CONTRACT, STRICT LIABILITY, OR TORT (INCLUDING NEGLIGENCE OR OTHERWISE) ARISING IN ANY WAY OUT OF THE PUBLICATION, USE OF, OR RELIANCE UPON ANY STANDARD, EVEN IF ADVISED OF THE POSSIBILITY OF SUCH DAMAGE AND REGARDLESS OF WHETHER SUCH DAMAGE WAS FORESEEABLE.

Translations

The IEEE consensus development process involves the review of documents in English only. In the event that an IEEE standard is translated, only the English version published by IEEE should be considered the approved IEEE standard.

Official statements

A statement, written or oral, that is not processed in accordance with the IEEE-SA Standards Board Operations Manual shall not be considered or inferred to be the official position of IEEE or any of its committees and shall not be considered to be, or be relied upon as, a formal position of IEEE. At lectures, symposia, seminars, or educational courses, an individual presenting information on IEEE standards shall make it clear that his or her views should be considered the personal views of that individual rather than the formal position of IEEE.

Comments on standards

Comments for revision of IEEE Standards documents are welcome from any interested party, regardless of membership affiliation with IEEE. However, IEEE does not provide consulting information or advice pertaining to IEEE Standards documents. Suggestions for changes in documents should be in the form of a proposed change of text, together with appropriate supporting comments. Since IEEE standards represent a consensus of concerned interests, it is important that any responses to comments and questions also receive the concurrence of a balance of interests. For this reason, IEEE and the members of its societies and Standards Coordinating Committees are not able to provide an instant response to comments or questions except in those cases where the matter has previously been addressed. For the same reason, IEEE does not respond to interpretation requests. Any person who would like to participate in revisions to an IEEE standard is welcome to join the relevant IEEE working group.

Comments on standards should be submitted to the following address:

Secretary, IEEE-SA Standards Board
445 Hoes Lane
Piscataway, NJ 08854 USA

Laws and regulations

Users of IEEE Standards documents should consult all applicable laws and regulations. Compliance with the provisions of any IEEE Standards document does not imply compliance to any applicable regulatory requirements. Implementers of the standard are responsible for observing or referring to the applicable regulatory requirements. IEEE does not, by the publication of its standards, intend to urge action that is not in compliance with applicable laws, and these documents may not be construed as doing so.

Copyrights

IEEE draft and approved standards are copyrighted by IEEE under U.S. and international copyright laws. They are made available by IEEE and are adopted for a wide variety of both public and private uses. These include both use, by reference, in laws and regulations, and use in private self-regulation, standardization, and the promotion of engineering practices and methods. By making these documents available for use and adoption by public authorities and private users, IEEE does not waive any rights in copyright to the documents.

Photocopies

Subject to payment of the appropriate fee, IEEE will grant users a limited, non-exclusive license to photocopy portions of any individual standard for company or organizational internal use or individual, non-commercial use only. To arrange for payment of licensing fees, please contact Copyright Clearance Center, Customer Service, 222 Rosewood Drive, Danvers, MA 01923 USA; +1 978 750 8400. Permission to photocopy portions of any individual standard for educational classroom use can also be obtained through the Copyright Clearance Center.

Updating of IEEE Standards documents

Users of IEEE Standards documents should be aware that these documents may be superseded at any time by the issuance of new editions or may be amended from time to time through the issuance of amendments, corrigenda, or errata. An official IEEE document at any point in time consists of the current edition of the document together with any amendments, corrigenda, or errata then in effect.

Every IEEE standard is subjected to review at least every ten years. When a document is more than ten years old and has not undergone a revision process, it is reasonable to conclude that its contents, although still of some value, do not wholly reflect the present state of the art. Users are cautioned to check to determine that they have the latest edition of any IEEE standard.

In order to determine whether a given document is the current edition and whether it has been amended through the issuance of amendments, corrigenda, or errata, visit the IEEE-SA Website at <http://ieeexplore.ieee.org/xpl/standards.jsp> or contact IEEE at the address listed previously. For more information about the IEEE-SA or IEEE's standards development process, visit the IEEE-SA Website at <http://standards.ieee.org>.

Errata

Errata, if any, for all IEEE standards can be accessed on the IEEE-SA Website at the following URL: <http://standards.ieee.org/findstds/errata/index.html>. Users are encouraged to check this URL for errata periodically.

Patents

Attention is called to the possibility that implementation of this standard may require use of subject matter covered by patent rights. By publication of this standard, no position is taken by the IEEE with respect to the existence or validity of any patent rights in connection therewith. If a patent holder or patent applicant has filed a statement of assurance via an Accepted Letter of Assurance, then the statement is listed on the IEEE-SA Website at <http://standards.ieee.org/about/sasb/patcom/patents.html>. Letters of Assurance may indicate whether the Submitter is willing or unwilling to grant licenses under patent rights without compensation or under reasonable rates, with reasonable terms and conditions that are demonstrably free of any unfair discrimination to applicants desiring to obtain such licenses.

Essential Patent Claims may exist for which a Letter of Assurance has not been received. The IEEE is not responsible for identifying Essential Patent Claims for which a license may be required, for conducting inquiries into the legal validity or scope of Patents Claims, or determining whether any licensing terms or conditions provided in connection with submission of a Letter of Assurance, if any, or in any licensing agreements are reasonable or non-discriminatory. Users of this standard are expressly advised that determination of the validity of any patent rights, and the risk of infringement of such rights, is entirely their own responsibility. Further information may be obtained from the IEEE Standards Association.

Participants

At the time this IEEE standard was completed, the Personal Health Device Working Group had the following membership:

Daidi Zhong, Chair
Michael J. Kirwan, Chair
Christoph Fischer, Vice Chair

Charles R. Abbruscato
 Nabil Abujbara
 Maher Abuzaid
 Manfred Aigner
 Jorge Alberola
 Karsten Alders
 Murtaza Ali
 Rolf Ambuehl
 David Aparisi
 Lawrence Arne
 Diego B. Arquillo
 Serafin Arroyo
 Muhammad Asim
 Merat Bagha
 Doug Baird
 David Baker
 Anindya Bakshi
 Ananth Balasubramanian
 Sunlee Bang
 M. Jonathan Barkley
 Gilberto Barrón
 David Bean
 John Bell
 Rudy Belliardi
 Daniel Bernstein
 George A. Bertos
 Chris Biernacki
 Ola Björnsne
 Thomas Blackadar
 Marc Blanchet
 Thomas Bluethner
 Douglas P. Bogia
 Xavier Boniface
 Shannon Boucousis
 Julius Broma
 Lyle G. Bullock, Jr.
 Bernard Burg
 Chris Burns
 Anthony Butt
 Jeremy Byford-Rew
 Satya Calloji
 Carole C. Carey
 Santiago Carot-Nemesio
 Randy W. Carroll
 Simon Carter
 Seungchul Chae
 Rahul Chauhan
 James Cheng
 Peggy Chien
 Chia-Chin Chong
 Saeed A. Choudhary

Jinhan Chung
 Malcolm Clarke
 John A. Cogan
 John T. Collins
 Cory Condek
 Todd H. Cooper
 David Cornejo
 Douglas Coup
 Nigel Cox
 Hans Crommenacker
 Tomio Crosley
 David Culp
 Allen Curtis
 Ndifor Cyril Fru
 Eyal Dassau
 David Davenport
 Russell Davis
 Ed Day
 Sushil K. Dekka
 Pedro de-las-Heras-Quiros
 Jim DelloStritto
 Matthew d'Entremont
 Lane Desborough
 Kent Dicks
 Hyoungho Do
 Xiaolian Duan
 Brian Dubreuil
 Jakob Ehrensvar
 Fredrik Einberg
 Roger M. Ellingson
 Michihiro Enokida
 Javier Escayola Calvo
 Leonardo Estevez
 Roger Feeley
 Bosco T. Fernandes
 Morten Flintrup
 Joseph W. Forler
 Russell Foster
 Eric Freudenthal
 Matthias Frohner
 Ken Fuchs
 Jing Gao
 Marcus Garbe
 John Garguilo
 Rick Geimer
 Igor Gejdos
 Ferenc Gerbovics
 Nicolae Goga
 Julian Goldman
 Raul Gonzalez Gomez
 Chris Gough

Channa Gowda
 Charles M. Gropper
 Amit Gupta
 Jeff Guttmacher
 Rasmus Haahr
 Christian Habermann
 Michael Hagerty
 Jerry Hahn
 Robert Hall
 Rickey L. Hampton
 Sten Hanke
 Jordan Hartmann
 Kai Hassing
 Marc Daniel Haunschild
 Wolfgang Heck
 Charles Henderson
 Jun-Ho Her
 Takashi Hibino
 Timothy L. Hirou
 Allen Hobbs
 Alex Holland
 Arto Holopainen
 Robert Hoy
 Frank Hsu
 Anne Huang
 Sen-Der Huang
 Zhiqiang Huang
 Ron Huby
 Robert D. Hughes
 David Hughes
 Jiyoung Huh
 Hugh Hunter
 Hitoshi Ikeda
 Yutaka Ikeda
 Philip O. Isaacson
 Atsushi Ito
 Michael Jaffe
 Praduman Jain
 Danny Jochelson
 Chris Johnson
 Phaneeth Junga
 Akiyoshi Kabe
 Steve Kahle
 Tomio Kamioka
 Kei Kariya
 Andy Kaschl
 Junzo Kashiwara
 Kohichi Kashiwagi
 Ralph Kent
 Laurie M. Kermes
 Ikuko Keshi

Junhyung Kim
 Min-Joon Kim
 Minho Kim
 Taekon Kim
 Tetsuya Kimura
 Alfred Kloos
 Jeongmee Koh
 Jean-Marc Koller
 John Koon
 Patty Krantz
 Alexander Kraus
 Ramesh Krishna
 Geoffrey Kruse
 Falko Kuester
 Rafael Lajara
 Pierre Landau
 Jaechul Lee
 JongMuk Lee
 Kyong Ho Lee
 Rami Lee
 Sungkee Lee
 Woojae Lee
 Yonghee Lee
 Joe Lenart
 Kathryn A. Lesh
 Qiong Li
 Ying Li
 Patrick Lichter
 Jisoon Lim
 Joon-Ho Lim
 John Lin
 Jiajia Liu
 Wei-Jung Lo
 Charles Lowe
 Don Ludolph
 Christian Luszick
 Bob MacWilliams
 Srikanth Madhurbootheswaran
 Romain Marmot
 Sandra Martinez
 Miguel Martínez de Espronceda
 Cámara
 Peter Mayhew
 Jim McCain
 László Meleg
 Alexander Mense
 Ethan Metsger
 Yu Miao
 Jinsei Miyazaki
 Erik Moll
 Darr Moore
 Piotr Murawski
 Soundharya Nagasubramanian
 Jae-Wook Nah
 Alex Neefus
 Trong-Nghia Nguyen-Dobinsky
 Michael E. Nidd
 Tetsu Nishimura
 Jim Niswander
 Hiroaki Niwamoto

Thomas Norgall
 Anand Noubade
 Yoshiteru Nozoe
 Abraham Ofek
 Brett Olive
 Begonya Otal
 Charles Palmer
 Bud Panjwani
 Carl Pantiskas
 Harry P. Pappas
 Mikey Paradis
 Hanna Park
 Jong-Tae Park
 Myungeun Park
 Soojun Park
 Phillip E. Pash
 TongBi Pei
 Soren Petersen
 James Petisce
 Peter Piction
 Michael Pliskin
 Jeff Price
 Harald Prinzhorn
 John Quinlan
 Arif Rahman
 Tanzilur Rahman
 Steve Ray
 Phillip Raymond
 Tim Reilly
 Barry Reinhold
 Brian Reinhold
 Melvin I. Reynolds
 John G. Rhoads
 Jeffrey S. Robbins
 Moskowitz Robert
 Timothy Robertson
 David Rosales
 Bill Saltzstein
 Benedikt Salzbrunn
 Giovanna Sannino
 Jose A. Santos-Cadenas
 Stefan Sauermann
 John Sawyer
 Guillaume Schatz
 Alois Schloegl
 Paul S. Schluter
 Lars Schmitt
 Mark G. Schnell
 Richard A. Schrenker
 Antonio Scorpiniti
 Kwang Seok Seo
 Riccardo Serafin
 Sid Shaw
 Frank Shen
 Liqun Shen
 Bozhi Shi
 Min Shih
 Mazen Shihabi
 Redmond Shouldice
 Sternly K. Simon
 Marjorie Skubic

Robert Smith
 Ivan Soh
 Motoki Sone
 Emily Sopensky
 Rajagopalan Srinivasan
 Andreas Staubert
 Nicholas Steblay
 Beth Stephen
 Lars Steubesand
 John (Ivo) Stivorik
 Raymond A. Strickland
 Hermann Suominen
 Lee Surprenant
 Ravi Swami
 Ray Sweidan
 Jin Tan
 Haruyuyki Tatsumi
 John W. Thomas
 Brad Tipler
 Jonas Tirén
 James Tomcik
 Janet Traub
 Jesús Daniel Trigo
 Gary Tschautscher
 Masato Tsuchid
 Ken Tubman
 Yoshihiro Uchida
 Sunil Unadkat
 Fabio Urbani
 Philipp Urbauer
 Laura Vanzago
 Alpo Värri
 Ciro de la Vega
 Dalimar Velez
 Naveen Verma
 Rudi Voon
 Isobel Walker
 David Wang
 Jerry P. Wang
 Yao Wang
 Yi Wang
 Steve Warren
 Fujio Watanabe
 Toru Watsuji
 Mike Weng
 Kathleen Wible
 Paul Williamson
 Jan Wittenber
 Jia-Rong Wu
 Will Wykeham
 Ariton Xhafa
 Junjie Yang
 Ricky Yang
 Melanie Yeung
 Done-Sik Yoo
 Jason Zhang
 Zhiqiang Zhang
 Thomas Zhao
 Miha Zoubek
 Szymon Zysko

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

Pieter Botman
Lyle G. Bullock, Jr.
Juan Carreon
Jianwen Chen
Keith Chow
Raul Colcher
Charles Cook
Paul Croll
Russell Davis
Sourav Dutta
Christoph Fischer
David Friscia
David Fuschi
Hector Barron Gonzalez
Randall Groves
Kai Hassing

Werner Hoelzl
Ruimin Hu
Ronald Huby
Noriyuki Ikeuchi
Akio Iso
Atsushi Ito
Mark Jaeger
Raj Jain
Shinkyō Kaku
Piotr Karocki
Stuart Kerry
Bruce Kraemer
Geoff Ladwig
Richard Lancaster
Vincent Lipsio

Greg Luri
Michael Newman
Charles Ngethe
Satoshi Oyama
Melvin I. Reynolds
Bartien Sayogo
Lars Schmitt
Shusaku Shimada
Steven Smith
Kapil Sood
Walter Struppler
Jiande Sun
Hung-Yu Wei
Jan Wittenber
Oren Yuen
Daidi Zhong

When the IEEE-SA Standards Board approved this standard on 21 August 2014, it had the following membership:

John Kulick, *Chair*
Jon Walter Rosdahl, *Vice Chair*
Richard H. Hulett, *Past Chair*
Konstantinos Karachalios, *Secretary*

Peter Balma
Farooq Bari
Ted Burse
Clint Chaplain
Stephen Dukes
Jean-Phillippe Faure
Gary Hoffman

Michael Janezic
Jeffrey Katz
Joseph L. Koepfinger*
David J. Law
Hung Ling
Oleg Logvinov
T. W. Olsen
Glenn Parsons

Ron Peterson
Adrian Stephens
Peter Sutherland
Yatin Trivedi
Phil Winston
Don Wright
Yu Yuan

*Member Emeritus

Also included are the following nonvoting IEEE-SA Standards Board liaisons:

Richard DeBlasio, *DOE Representative*
Michael Janezic, *NIST Representative*

Don Messina
IEEE-SA Content Publishing

Kathryn M. Bennett
IEEE-SA Technical Community Programs

Introduction

This introduction is not part of IEEE Std 11073-10424-2014, Health informatics—Personal Health Device Communication—Part 10424: Device Specialization—Sleep Apnoea Breathing Therapy Equipment (SABTE).

ISO/IEEE 11073 standards enable communication between medical devices and external computer systems. Within the context of the ISO/IEEE 11073 family of standards for device communication, this standard establishes a normative definition of the communication between sleep apnoea breathing therapy equipment (SABTE) devices (agents) and managers (e.g., cell phones, personal computers, personal health appliances, set top boxes) in a manner that enables plug-and-play interoperability. It leverages appropriate portions of existing standards including ISO/IEEE 11073 terminology, information models, application profile standards, and transport standards. It specifies the use of specific term codes, formats, and behaviors in telehealth environments restricting optionality in base frameworks in favor of interoperability. This standard defines a common core of communication functionality for SABTE. In this context, SABTE is defined as a device that is intended to alleviate the symptoms of a patient who suffers from sleep apnoea by delivering a therapeutic breathing pressure to the patient. SABTE is primarily used in the home health-care environment by a lay operator without direct professional supervision.

Contents

1. Overview	1
1.1 Scope	1
1.2 Purpose	2
1.1 Context	2
2. Normative references.....	2
3. Definitions, acronyms, and abbreviations	3
3.1 Definitions	3
3.2 Acronyms and abbreviations	4
4. Introduction to ISO/IEEE 11073 personal health devices standards	5
4.1 General	5
4.2 Introduction to ISO/IEEE 11073-20601 modeling constructs	5
4.3 Compliance with other standards.....	6
5. SABTE device concepts and modalities.....	7
5.1 General	7
5.2 Compliance monitoring	10
5.3 Efficacy monitoring	11
5.4 Service monitoring	13
5.5 Device settings.....	14
5.6 Therapy settings.....	14
6. Sleep apnoea breathing therapy equipment domain information model.....	18
6.1 Overview	18
6.2 Class extensions.....	18
6.3 Object instance diagram	19
6.4 Types of configuration.....	21
6.5 Profile	21
6.6 Medical device system object.....	23
6.7 Numeric objects.....	29
6.8 Real-time sample array objects.....	48
6.9 Enumeration objects	50
6.10 PM-store objects	60
6.11 Scanner objects	65
6.12 Class extension objects	68
6.13 SABTE information model extensibility rules	69
7. SABTE service model	69
7.1 General	69
7.2 Object access services.....	69
7.3 Object access event report services	72
8. SABTE communication model.....	73
8.1 Overview	73
8.2 Communications characteristics	73
8.3 Association procedure	73
8.4 Configuring procedure.....	75
8.5 Operating procedure	78
8.6 Time synchronization	79

9. Test associations	79
9.1 Behavior with standard configuration.....	79
9.2 Behavior with extended configurations	79
10. Conformance	80
10.1 Applicability	80
10.2 Conformance specification	80
10.3 Levels of conformance	80
10.4 Implementation conformance statements	81
Annex A (informative) Bibliography	86
Annex B (normative) Any additional ASN.1 definitions	87
B.1 Efficacy annotations bit mapping.....	87
B.2 Compliance annotations bit mapping.....	88
B.3 PHD DM status bit mapping	88
B.4 Autostart/-stop bit mapping	89
Annex C (normative) Allocation of identifiers.....	90
C.1 General.....	90
C.2 Definitions of terms and codes.....	90
C.3 Systematic derivations of terms and codes	93
Annex D (informative) Message sequence examples.....	101
Annex E (informative) Protocol data unit examples	103
E.1 General	103
E.2 Association information exchange	103
E.3 Configuration information exchange.....	106
E.4 GET MDS attributes service	109
E.5 Data reporting.....	110
E.6 Disassociation	111

Health informatics—Personal health device communication

Part 10424: Device Specialization— Sleep Apnoea Breathing Therapy Equipment (SABTE)

IMPORTANT NOTICE: IEEE Standards documents are not intended to ensure safety, security, health, or environmental protection, or ensure against interference with or from other devices or networks. Implementers of IEEE Standards documents are responsible for determining and complying with all appropriate safety, security, environmental, health, and interference protection practices and all applicable laws and regulations.

This IEEE document is made available for use subject to important notices and legal disclaimers. These notices and disclaimers appear in all publications containing this document and may be found under the heading “Important Notice” or “Important Notices and Disclaimers Concerning IEEE Documents.” They can also be obtained on request from IEEE or viewed at <http://standards.ieee.org/IPR/disclaimers.html>.

1. Overview

1.1 Scope

Within the context of the ISO/IEEE 11073 family of standards for device communication, this standard establishes a normative definition of the communication between sleep apnoea breathing therapy equipment and managers (e.g., cell phones, personal computers, personal health appliances, set top boxes) in a manner that enables plug-and-play interoperability. It leverages appropriate portions of existing standards including ISO/IEEE 11073 terminology, information models, application profile standards, and transport standards. It specifies the use of specific term codes, formats, and behaviors in telehealth environments restricting optionality in base frameworks in favor of interoperability. This standard defines a common core of communication functionality for sleep apnoea breathing therapy equipment. In this context, sleep apnoea breathing therapy equipment are defined as devices that are intended to alleviate the symptoms of a patient who suffers from sleep apnoea by delivering a therapeutic breathing pressure to the patient. Sleep apnoea breathing therapy equipment are primarily used in the home health-care environment by a lay operator without direct professional supervision.

1.2 Purpose

This standard addresses a need for an openly defined, independent standard for controlling information exchange to and from personal health devices (agents) and managers (e.g., cell phones, personal computers, personal health appliances, and set top boxes). Interoperability is key to growing the potential market for these devices and to enabling people to be better informed participants in the management of their health.

1.1 Context

See IEEE Std 11073-20601a™-2010 for an overview of the environment within which this standard is written.¹

This standard defines the device specialization for the SABTE, being a specific agent type, and it provides a description of the device concepts, its capabilities, and its implementation according to this standard.

This standard is based on IEEE Std 11073-20601a-2010, which in turn draws information from both ISO/IEEE 11073-10201:2004 [B9] and ISO/IEEE 11073-20101:2004 [B10].² The medical device encoding rules (MDERs) used within this standard are fully described in IEEE Std 11073-20601a-2010.

This standard reproduces relevant portions of the nomenclature found in ISO/IEEE 11073-10101:2004 [B8] and adds new nomenclature codes for the purposes of this standard. Between this standard and IEEE Std 11073-20601a-2010, all required nomenclature codes for implementation are documented.

NOTE 1—IEEE Std 11073-20601a-2010 is an amendment to ISO/IEEE 11073-20601:2010. It contains new material and corrections and does not copy the content of ISO/IEEE 11073-20601:2010. Throughout this standard, a reference to IEEE Std 11073-20601a-2010 refers to the document that is obtained after applying this new material and corrections to ISO/IEEE 11073-20601:2010.³

NOTE 2—In this standard, ISO/IEEE 11073-104zz is used to refer to the collection of device specialization standards that utilize IEEE Std 11073-20601a-2010, where zz can be any number from 01 to 99, inclusive.

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.

ISO/IEEE 11073-20601:2010, Health informatics—Personal health device communication—Part 20601: Application profile—Optimized Exchange Protocol.⁴

IEEE Std 11073-20601a™-2010, Health informatics—Personal health device communication—Part 20601: Application profile—Optimized Exchange Protocol—Amendment 1.^{5, 6}

See Annex A for all informative material referenced by this standard.

¹ Information on references can be found in Clause 2.

² The numbers in brackets correspond to those of the bibliography in Annex A.

³ Notes in text, tables, and figures are given for information only and do not contain requirements needed to implement the standard.

⁴ ISO/IEEE publications are available from the ISO Central Secretariat (<http://www.iso.org/>). ISO/IEEE publications are also available in the United States from The Institute of Electrical and Electronics Engineers (<http://standards.ieee.org/>).

⁵ IEEE publications are available from The Institute of Electrical and Electronics Engineers (<http://standards.ieee.org/>).

⁶ The IEEE standards or products referred to in this clause are trademarks of The Institute of Electrical and Electronics Engineers, Inc.