



BSI Standards Publication

**Male condoms — Guidance on the use of ISO 4074 and
ISO 23409 in the quality management of condoms**

National foreword

This British Standard is the UK implementation of ISO 16038:2017. It supersedes BS ISO 16038:2005, which is withdrawn.

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A list of organizations represented on this committee can be obtained on request to its secretary.

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**Male condoms — Guidance on the
use of ISO 4074 and ISO 23409 in the
quality management of condoms**

*Préservatifs masculins — Lignes directrices sur l'utilisation de la
norme ISO 4074 et ISO 23409 sur le management de la qualité
des préservatifs en latex de caoutchouc naturel et en matériau
synthétiques*



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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.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation on the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see the following URL: www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 157, *Non-systemic contraceptives and STI barrier prophylactics*.

This second edition cancels and replaces the first edition (ISO 16038:2005), which has been technically revised, considering the revisions to ISO 4074 and the publication of ISO 23409. The modifications are as follows.

- a) The title and Scope have been expanded to include ISO 23409 and the relevant aspects of synthetic male condoms have been added in this edition. The major points incorporated are with respect to design, determination of limits for burst properties, stability studies and clinical trials.
- b) The revision to ISO 4074 and points arising out of the publication of ISO 4074:2015 have been incorporated in the guidance document.
- c) An explanation regarding the application of switching rules in sampling in accordance with ISO 2859-1 has been incorporated.
- d) The section on design has been expanded to explain significant changes to condoms, which warrant validation.
- e) The principle of estimating shelf life of natural rubber latex condoms has been revised to reflect the principles of shelf determination as given in ISO 4074:2015.
- f) The section on testing has been revised to include the modifications to test methods for determining freedom for holes.
- g) The section on dimensions has been revised to include the aspects of tolerances for thinner condoms.
- h) The aspects of condoms of smaller and larger sizes than those specified in ISO 4074 have been incorporated.
- i) The impact of new test for visibly open seals as given in ISO 4074 and potential rework has been addressed.

- j) The control of maximum storage period of naked condoms before packing them in individual sealed containers has been incorporated in accordance with ISO 4074.

Introduction

Condoms are medical devices used for contraception and for prevention of sexually transmitted infections.

ISO 4074 is a quality standard for natural rubber latex condoms and ISO 23409 for condoms made from synthetic materials. They are reference documents for standardized end product quality test protocols and a baseline specification for critical attributes that affect condom safety and effectiveness. They are applied by manufacturers, procurement agencies, regulatory bodies and testing laboratories.

The use of ISO 4074 and or ISO 23409 does not by itself ensure consistency in quality; consistent high quality at the lowest possible cost is attained only through a regime termed quality management, through which, quality is built into the product and ensured at every point in the design, planning, production and procurement processes. This document should lead to continuous improvement in manufacturing, procurement and testing processes. The special requirements of buyers and consumers should also be given due consideration when applying ISO 4074 or ISO 23409, as ISO 4074 and ISO 23409 are general by design, and will not cover all circumstances completely.

This document provides guidance to manufacturers, buyers and third-party test laboratories on implementing and applying ISO 4074 in the manufacture of condoms, and to purchasers on applying ISO 4074 or ISO 23409 and verifying that the condoms delivered conform to the specification, as appropriate.

Acceptable condoms meet or exceed the minimum requirements specified in ISO 4074 or ISO 23409, as applicable.

It is not possible, nor is it required, to subject condoms to user trials on a batch-by-batch basis. For this reason, certain evaluations are carried out only in the case of a pre-market validation; for example for new or significantly modified designs.

Design validation requirements normally include all the good manufacturing practice (GMP) validation requirements and the validation requirements of ISO 9001 and ISO 13485; these are not currently covered by ISO 4074 and ISO 23409, but are generally included by regulatory authorities as prerequisites for registering new designs of medical devices. Design considerations such as stability testing, etc., are, however, covered in ISO 4074 and evaluation of barrier properties by clinical trials and determination of burst properties are covered in ISO 23409.

ISO 4074 and ISO 23409 are mainly concerned with finished product testing carried out to monitor or to verify that the condoms have been manufactured with an adequate level of consistency in quality. For this purpose, tests have been designed that can be carried out rapidly and economically. The requirements in ISO 4074 and ISO 23409 are based on those properties which, based upon current knowledge, are believed to be relevant to the performance of condoms in normal use.

Some important properties of condoms are nevertheless difficult to define in quantitative terms because of a lack of controlled studies, the absence of practical and economical tests, and the need for different specifications to suit different users. ISO 4074 and ISO 23409 are, therefore, focused on the essential properties where limits can be clearly defined. Other properties are addressed only in general terms and are meant to be augmented through appropriate manufacturing records, certification by the manufacturer or by buyers' specifications.

This document also addresses how to deal with other related important issues not covered by ISO 4074 and ISO 23409.

It is meant to help the user of ISO 4074 and ISO 23409 to understand any risks that can be associated with the use of condoms. It also helps in deciding whether such risks are acceptable when weighed against the benefits to the condom user. ISO 4074 and ISO 23409 also help in assessing whether the products are demonstrably safe and offer protection to health. Good communication between the

buyer and the manufacturer will result in the delivery of satisfactory and safe products, thus avoiding unnecessary testing or inappropriate specifications, and thereby minimizing conformity testing costs.

NOTE In many countries, condoms, being medical devices, are subject to regulations.

The requirements for quality management are given in standards such as ISO 9001 and ISO 13485. ISO 9001 is based on the approach of achieving business excellence through quality management. For condoms, being a medical device, it is appropriate that ISO 13485 is applied for quality management as part of compliance to regulatory requirements.

Male condoms — Guidance on the use of ISO 4074 and ISO 23409 in the quality management of condoms

1 Scope

This document provides guidance on using ISO 4074 and ISO 23409 and addresses quality issues to be considered during the development, manufacture, quality verification and procurement of condoms. It encompasses the aspects of quality management systems in the design, manufacture and delivery of condoms with an emphasis on performance, safety and reliability.

Male condoms are either made from essentially natural rubber latex, in which case the requirements of ISO 4074 are applicable, or from synthetic materials and/or blends of synthetic materials and natural rubber latex, in which case the requirements of ISO 23409 are applicable. This document outlines the aspects applicable to both types of condoms with specific clarifications where appropriate.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 4074:2015, *Natural rubber latex male condoms — Requirements and test methods*

ISO 9000, *Quality management systems — Fundamentals and vocabulary*

ISO 13485, *Medical devices — Quality management systems — Requirements for regulatory purposes*

ISO 14971, *Medical devices — Application of risk management to medical devices*

ISO 23409:2011, *Male condoms — Requirements and test methods for condoms made from synthetic materials*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 4074, ISO 9000, ISO 13485, ISO 14971 and ISO 23409 apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at <http://www.electropedia.org/>
- ISO Online browsing platform: available at <https://www.iso.org/obp>

4 Quality of design

4.1 General

A condom is a single-use medical device, the performance and safety of which depends upon the design and the manufacturing process. New designs of condoms can require clinical testing, several other tests and analysis on a limited basis for validation purposes, such as shelf-life determination (type testing) and risk assessment. These requirements are generally prescribed by licensing authorities and the data generated become part of the master file for the product. Guidelines are available in ISO 13485