



**Quality management—Guidelines for
quality plans**

STANDARDS
Australia



Currently in preview, click buy full version

AS ISO 10005:2018

This Australian Standard® was prepared by QR-008, Quality Systems. It was approved on behalf of the Council of Standards Australia on 23 October 2018.

This Standard was published on 6 November 2018.

The following are represented on Committee QR-008:

Association of Accredited Certification Bodies
Australian Industry Group
Australian Institute of Petroleum
Australian Organisation for Quality
Australian Petroleum Production and Exploration Association
Australian Prudential Regulation Authority
Chemistry Australia
Consumers Federation of Australia
Department of Defence (Australian Government)
Department of Infrastructure and Regional Development (Australian Government)
Energy Networks Australia
Engineers Australia
ISACA
Joint Accreditation System of Australia and New Zealand
Master Builders Australia
Materials Australia
Royal Australian Chemical Institute
University of Wollongong

This Standard was issued in draft form for comment as DR AS/NZS ISO 10005:2018.

Keeping Standards up-to-date

Ensure you have the latest versions of our publications and keep up-to-date about Amendments, Rulings, Withdrawals, and new projects by visiting:

www.standards.org.au

www.saiglobal.com (sales and distribution)

ISBN 978 1 76072 219 7



Quality management—Guidelines for quality plans

Originally AS/NZS ISO 9004.5(Int.):1995.
Jointly revised and designated AS/NZS 9004.5:1998.
Jointly revised and re-designated AS/NZS ISO 10005:2006.
Revised and re-designated AS ISO 10005:2018.

COPYRIGHT

© ISO — All rights reserved
© Standards Australia Limited 2018

All rights are reserved. No part of this work may be reproduced or copied in any form or by any means, electronic or mechanical, including photocopying, without the written permission of the publisher, unless otherwise permitted under the Copyright Act 1968 (Cth).

Published by SAI Global Pty Limited under licence from Standards Australia Limited, GPO Box 476, Sydney, NSW 2001, Australia.

Preface

This Standard was prepared by the Australian committee members of the Joint Standards Australia/Standards New Zealand Committee QR-008, Quality Systems, to supersede AS/NZS ISO 10005:2006, *Quality management systems — Guidelines for quality plans*.

After consultation with stakeholders in both countries, Standards Australia and Standards New Zealand decided to develop this Standard as an Australian Standard rather than an Australian/New Zealand Standard.

The objective of this Standard is to provide guidelines for establishing, reviewing, accepting, applying and revising quality plans. This document is applicable to quality plans for any intended output, whether a process, product, service, project or contract, and any type or size of organization. It is applicable whether or not the organization has a management system in conformity with AS/NZS ISO 9001.

This document provides guidance and does not specify requirements. It is focused primarily on the provision of outputs and is not a guide to the planning of quality management system development.

NOTE To avoid undue repetition of “process, product, service, project or contract”, this document uses the term “specific case”.

This Standard is identical with, and has been reproduced from, ISO 10005:2018, *Quality management — Guidelines for quality plans*.

As this document has been reproduced from an International Standard, the following applies:

- (a) In the source text “this document” should read “this Australian Standard”.
- (b) A full point substitutes for a comma when referring to a decimal marker.

Australian or Australian/New Zealand Standards that are identical adoptions of international normative references may be used interchangeably. Refer to the online catalogue for information on specific Standards.

The terms “normative” and “informative” are used in Standards to define the application of the appendices or annexes to which they apply. A “normative” appendix or annex is an integral part of a Standard, whereas an ‘informative’ appendix or annex is only for information and guidance.

NOTE Any feedback or questions on this document should be directed to the user’s national standards body.

Contents

Preface	ii
Foreword	v
Introduction	vii
1 Scope	1
2 Normative references	1
3 Terms and definitions	1
4 Using a quality plan	2
4.1 Introduction	2
4.2 Requesting external provider quality plans	2
4.3 Managing external provider quality plans	3
5 Development of a quality plan	4
5.1 Context of the quality plan	4
5.2 Inputs to the quality plan	4
5.3 Defining the scope of the quality plan	5
5.4 Preparation of the quality plan	5
5.4.1 Initiation	5
5.4.2 Defining the quality plan	5
5.4.3 Consistency and compatibility	5
5.4.4 Presentation and structure	6
6 Content of the quality plan	6
6.1 General	6
6.2 Scope of the quality plan	6
6.3 Quality plan inputs	6
6.4 Quality objectives	7
6.5 Quality plan responsibilities	7
6.6 Control of documented information	7
6.7 Resources	8
6.7.1 Provision of resources	8
6.7.2 Materials, products and services	8
6.7.3 People	8
6.7.4 Infrastructure and environment for the operation of processes	8
6.7.5 Monitoring and measuring resources	8
6.8 Customers and other interested parties communication	9
6.9 Design and development	9
6.9.1 Design and development process	9
6.9.2 Control of design and development changes	9
6.10 Externally provided processes, products and services	10
6.11 Production and service provision	10
6.12 Identification and traceability	11
6.13 Property belonging to customers or external providers	11
6.14 Preservation of outputs	11
6.15 Control of nonconforming outputs	12
6.16 Monitoring and measurement	12
6.17 Audits	12
7 Operation and control of the quality plan	13
7.1 Review and acceptance of the quality plan	13
7.2 Implementation and monitoring of the quality plan	13
7.3 Revision of the quality plan	14
7.4 Feedback and improvement	14
Annex A (informative) Examples of formats for quality plans	15

Annex B	(informative) Schematic representation of a process approach applied to quality plans	22
Annex C	(informative) Correlation matrix between the clauses in this document and those in ISO 9001:2015	23
Annex D	(informative) Correlation matrix between the clauses of this document and the quality management principles from ISO 9000:2015	24
Bibliography		27

Currently in preview, click buy full version

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 176, *Quality management and quality assurance*, Subcommittee SC 2, *Quality systems*.

This third edition cancels and replaces the second edition (ISO 10005:2005), which has been technically revised.

The main changes compared with the previous edition are as follows.

- a) It applies the terminology from ISO 9000:2015, which includes changes to key definitions, such as:
 - 1) for the definition of "quality plan" (see 3.2), which has been modified to replace the phrase "procedures and associated resources to be applied when and by whom" by "actions, responsibilities and associated resources";
 - 2) for the definition of "specific case" (see 3.3), which has been modified to make reference to "service", as ISO 9001:2015 now refers to "products and services" and no longer just to "products";
 - 3) the replacement of the terms "documentation" and "record" by the term "documented information", which is generally used in ISO management system standards to include both "procedures" and "records" which are not necessarily distinct from each other in a digital environment (documented information needed to support process operation is "maintained", which means that it is established and updated as required; documented information that provides evidence of conformity with requirements is "retained" which means that it is protected from unintended alterations).

Table 1 — Major changes to terms in this document since its previous edition

ISO 10005:2005	This document
Products	Products and services
Documentation Quality manual Documented procedures Records	Documented information
Purchased product	Externally provided processes, products and services
Supplier	External provider
Monitoring and measuring equipment	Monitoring and measuring resources

b) It is aligned to ISO 9001:2015, leading to:

- 1) a significant revision in the clause/subclause sequence, titles and the addition of new material, e.g. the inclusion of “[5.2](#) Context of a quality plan”, or the extension of [7.2](#) to also reference the monitoring of a quality plan;
- 2) the incorporation of “risk-based thinking”.

c) A new clause ([Clause 4](#)) on using a quality plan.

Introduction

0.1 General

This document was prepared to address the need for guidance on quality plans, either in the context of an established quality management system or as an independent management activity. In either case, quality plans provide a means of relating specific requirements of the process, product, service, project or contract to work methods and practices. Quality plans are most effective when they are compatible with other associated plans. The guidance in this document can also be used where quality plans are integrated with other management plans or quality management systems.

Benefits of establishing a quality plan include increased confidence that requirements will be met, greater assurance that processes are in control and the motivation it can give to those involved. It might also give insight into opportunities for innovation and improvement.

The guidance on quality plans in this document is based on the quality management principles described in ISO 9000 and the concepts used in ISO 9001 for the establishment of quality management systems. [Clause 6](#), which describes the typical contents of a quality plan, includes guidance to applying relevant ISO 9001 requirements. The guidance is limited to quality plans and does not replace guidance given in ISO 9000 on quality management concepts or ISO/TS 9002 on the application of ISO 9001 requirements within an organization.

This document does not replace the guidance given in industry-specific documented information. Where quality plans are required for project applications, the guidance provided in this document is intended to be complementary to the guidance provided in ISO 10005. Some terms used in this document have been changed with respect to its previous edition to improve alignment with ISO 9001:2015 and other management system standards. There is no need for the terms used by an organization, whether in specifying quality plan requirements or developing a quality plan, to be replaced by the terms used in this document.

In this document, the following verbal forms are used:

- “should” indicates a recommendation;
- “may” indicates a permission;
- “can” indicates a possibility or a capability.

Information marked as “NOTE” is for guidance in understanding or clarifying the associated text.

NOTE See <https://www.iso.org/home/tc176sc2> for guidance on the topics in this Introduction.

0.2 Using this document

This Introduction explains some underlying concepts and changes to terms used in the previous edition of this document.

[Clauses 1 to 3](#) provide basic information (Scope, Normative references, and Terms and definitions).

[Clause 4](#) summarizes how quality plans can be used.

[Clause 5](#) describes the process of developing a quality plan.

[Clause 6](#) describes the typical contents of a quality plan.

[Clause 7](#) describes the operation and control of a quality plan.

[Annex A](#) provides examples of simple quality plans.

[Annex B](#) provides a schematic representation of a process approach applied to a quality plan

[Annex C](#) provides a correlation matrix between the clauses of this document and those of ISO 9001:2015.

[Annex D](#) provides a correlation matrix between the clauses of this document and the quality management principles from ISO 9000:2015.

The Bibliography includes a list of standards and other relevant information.

0.3 Process approach

The process approach means the systematic management of processes and their interactions to achieve intended results. Applying the process approach to quality plans assists organizations to manage the inputs, activities and outputs of each process within a coherent system of interrelated processes.

Processes referenced in a quality plan can interact with:

- each other (interactions among quality plan processes);
- other processes operated within the organization's management system;
- processes operated within other organizations (such as customers and external providers).

When considering how to manage its processes and their interactions, the organization can address these through a quality plan whether or not it has a quality management system.

[Annex B](#) provides a schematic representation of a process approach applied to quality plans.

0.4 Risk-based thinking

Risk-based thinking means applying a systematic approach to consider risk (the effect of uncertainty) so that risks can be understood and managed appropriately.

The application of risk-based thinking to the development and use of a quality plan enables an organization to determine the importance of particular issues and take appropriate actions to manage both risks and opportunities.

A customer requesting that a provider prepares a quality plan can apply risk-based thinking to determine the minimum requirements for the type and extent of the monitoring activities.

When developing a quality plan, the organization can apply risk-based thinking in deciding the processes, resources and control methods to be used. Particularly where an organization uses a standard model or template for different quality plans, risk-based thinking can assist those involved to make each quality plan fit for its intended purpose.

Australian Standard®

Quality management—Guidelines for quality plans

1 Scope

This document gives guidelines for establishing, reviewing, accepting, applying and revising quality plans.

This document is applicable to quality plans for any intended output, whether a process, product, service, project or contract, and any type or size of organization.

It is applicable whether or not the organization has a management system in conformity with ISO 9001.

This document provides guidance and does not specify requirements.

It is focused primarily on the provision of outputs and is not a guide to the planning of quality management system development.

NOTE To avoid undue repetition of “process, product, service, project or contract”, this document uses the term “specific case”.

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 9000:2015, *Quality management systems — Fundamentals and vocabulary*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 9000:2015 and the following apply.

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

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

3.1

documented information

information required to be controlled and maintained by an organization and the medium on which it is contained

Note 1 to entry: Documented information can be in any format and media and from any source.

Note 2 to entry: Documented information can refer to:

- the management system, including related *quality plans* (3.2) and processes;
- information created in order for the organization to operate (documentation);
- evidence of results achieved.

[SOURCE: ISO 9000:2015, 3.8.6, modified — In Note 2 to entry, the first list item has been modified, and Note 3 to entry has been deleted.]