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**Molecular biomarker analysis —
Method for the statistical evaluation of
analytical results obtained in testing
sub-sampled groups of genetically
modified seeds and grains — General
requirements**

*Analyse moléculaire de biomarqueurs — Méthode pour l'évaluation
statistique des résultats d'analyse obtenus lors des essais de sous-
échantillons multiples de semences et de graines génétiquement
modifiées — Exigences générales*



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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).

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For an explanation of 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 www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 34, *Food products*, Subcommittee SC 16, *Horizontal methods for molecular biomarker analysis*.

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.

This corrected version of ISO 22753:2021 incorporates the following corrections:

- [Formula C.1](#) has been corrected.

Introduction

Seed and grain testing is used throughout the world to commercially define the purity of seed and grain lots.

Commercial requirements for labelling agricultural products with genetically modified organism (GMO) content at a specified threshold level both as a seed/grain contaminant and a food ingredient have become common to satisfy regulations and consumer demands. Conformance with these specifications is evaluated at various points of the supply chain, often starting with the harvested grain.

Quantitative real-time polymerase chain reaction (PCR) can be used to determine the GMO content by analysis of the ratio of GMO DNA copy numbers to plant-species specific DNA copy numbers followed by a conversion to genetically modified (GM) mass fraction.

Multiple events stacked in a crop, such as those generated by crossing two or more single events, are widely used in agricultural production. A stacked event seed or grain containing GMO DNA corresponding to two or more GM events commingled in lot cannot be differentiated by quantitative PCR alone from multiple seeds within the lot each containing a single GM event. Consequently, if the actual measured GMO arises only from GM stacked event seeds, GM content measured by quantitative real-time PCR of a single sample will lead to an overestimation of the actual number of GM seeds or grains present.

The group testing strategy described in this document provides a reliable alternative to estimate the GM content on the basis of the fact that whole seeds/grains are the sample material.

The process described in this document can provide a method to accurately estimate the percentages of GM seeds/grains in a lot irrespective of the presence of stacked event seeds/grains. GM content is determined for representative subsampled groups of seed/grain from a lot and statistically analysed.

Molecular biomarker analysis — Method for the statistical evaluation of analytical results obtained in testing sub-sampled groups of genetically modified seeds and grains — General requirements

1 Scope

This document describes general requirements, procedures and performance criteria for evaluating the content of genetically modified (GM) seeds/grains in a lot by a group testing strategy that includes qualitative analysis of sub-sampled groups followed by statistical evaluation of the results.

This document is applicable to group testing strategy estimating the GM content on a percentage seed/grain basis for purity estimation, testing towards a given reject/accept criterion and for cases where seed/grain lots are carrying stacked events.

This document is not applicable to processed products.

NOTE Description of the use of group testing strategy are available in References [1], [7], [8], [18], [19] and [20].

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 16577, *Molecular biomarker analysis — Terms and definitions*

ISO 21572, *Foodstuffs — Molecular biomarker analysis — Immunochemical methods for the detection and quantification of proteins*

ISO 24276, *Foodstuffs — Methods of analysis for the detection of genetically modified organisms and derived products — General requirements and definitions*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 16577 and the following apply.

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

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

3.1

absolute PCR limit of detection

absolute polymerase chain reaction limit of detection

absolute PCR LOD

lowest nominal (average) number of target copies in the template volume distributed to individual PCRs that would allow for an acceptable probability of detecting the target