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**Microbiology of food and animal feed —
Horizontal method for determination
of hepatitis A virus and norovirus in
food using real-time RT-PCR —**

Part 1:
Method for quantification

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*Microbiologie des aliments — Méthode horizontale pour la recherche
des virus de l'hépatite A et norovirus dans les aliments par la
technique RT-PCR en temps réel —*

ISO/TS 15216-1:2013

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Partie 1: Méthode de quantification



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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

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.

In other circumstances, particularly when there is an urgent market requirement for such documents, a technical committee may decide to publish other types of document:

- an ISO Publicly Available Specification (ISO/PAS) represents an agreement between technical experts in an ISO working group and is accepted for publication if it is approved by more than 50 % of the members of the parent committee casting a vote;
- an ISO Technical Specification (ISO/TS) represents an agreement between the members of a technical committee and is accepted for publication if it is approved by 2/3 of the members of the committee casting a vote.

An ISO/PAS or ISO/TS is reviewed after three years in order to decide whether it will be confirmed for a further three years, revised to become an International Standard, or withdrawn. If the ISO/PAS or ISO/TS is confirmed, it is reviewed again after a further three years, at which time it must either be transformed into an International Standard or be withdrawn.

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.

ISO/TS 15216-1 was prepared by the European Committee for Standardization (CEN), in collaboration with Technical committee ISO/TC 34, *Food products*, Subcommittee SC 9 *Microbiology*.

This corrected version of ISO/TS 15216-1:2013 incorporates the following corrections.

- Throughout, textual references have been updated to take reordering of the annexes into account. [Annex B](#) was formerly Annex E; [Annex C](#) was formerly Annex D; [Annex D](#) was formerly Annex G; [Annex E](#) was formerly Annex C; [Annex F](#) was formerly Annex B; [Annex G](#) was formerly Annex H; [Annex H](#) was formerly Annex I; [Annex I](#) was formerly Annex F.
- Many cross-references to reagents or apparatus subclauses are added.
- Where units of shaking operations are mentioned, “oscillations min⁻¹” replaces “min⁻¹”.
- A phrase citing [Annex A](#) is added to the end of the introduction.
- The definitions for “food surface” (formerly 3.2 and 3.3) are combined and expanded in a redrafted [3.2](#); in consequence, the following terms in [Clause 3](#) are renumbered.
- In [3.4](#), Note 2, “There is only one serotype” is transposed to the end of Note 1. Also, “group 2 biological agent by the European Union and as a risk group 2 human aetiological agent by the United States National Institutes of Health” replaces “UK Advisory Committee on Dangerous Pathogens (ACDP) hazard group 2 pathogen”.

- In 3.5, Note 2, “group 2 biological agents by the European Union and as risk group 2 human aetiological agents by the United States National Institutes of Health” replaces “ACDP hazard group 2 pathogens”.
- In 3.6 and 3.7, “estimation of number of copies” replaces “quantification”.
- In 3.13, “used in” replaces “used as template in”.
- In 5.2.11, “from *Aspergillus niger* or *A. aculeatus*” is inserted after “Pectinase”.
- In 6.1, “Aerosol resistant tips should be used unless unobstructed tips are required, e.g. for aspiration.” is inserted.
- In 6.5, “ $37 \pm 1,0$ ” replaces “ 37 ± 10 ”.
- A redrafted 6.10 on centrifuge(s) and rotor(s) replaces the former 6.10 and 6.11, with consequent renumbering of the following subclauses.
- In 6.19, the square brackets are deleted.
- In 6.27, “**Real-time PCR machine(s)**, i.e. thermal cycler(s),” replaces “**Thermal cycler(s)**”.
- In 6.28, “selected real-time PCR” replaces “selected PCR”.
- In 8.1, “Samples arriving already frozen should be defrosted prior to testing.” is inserted as the second sentence.
- 8.2.3 Is redrafted.
- In 8.2.4, paragraph 2, “buffer (5.3.5) (for soft fruit samples, add 30 units pectinase from *A. niger*, or 1 140 units pectinase from *A. aculeatus* to the buffer) and” replaces “buffer (for soft fruit samples, add 30 units pectinase to the buffer) and”
- In 8.2.6, paragraph 2, “and the animal is supported with a rubber block” is added.
- In 8.2.6, last paragraph, “min at room temperature, decant” replaces “min, decant”
- In 8.4.2.3, paragraph 1, “using a real-time PCR machine (6.27)” is added.
- In 9.3, Note 1, “For a dsDNA standard curve with an idealized slope of $-3,32$, if the C_q value of the sample RNA + EC RNA well is $<2,00$ greater than the C_q value of the water + EC RNA well, the amplification efficiency is $>25\%$ and therefore acceptable; if the C_q value of the sample RNA + EC RNA well is $>2,00$ greater than the C_q value of the water + EC RNA well, the amplification efficiency is $<25\%$ and therefore not acceptable.” is added.
- In 9.4, Note 1 “a process control virus recovery (equal to the extraction efficiency in matrices other than BMS) of 100 %. For a process control virus RNA standard curve with an idealized slope of $-3,32$, if the C_q value of an undiluted sample RNA well is $<6,64$ greater than the C_q value of the undiluted process control virus RNA, the process control virus recovery for that sample is $>1\%$ and therefore acceptable” replaces “an extraction efficiency of 100 %”.
- The title of Annex B has been expanded to read, “Real-time RT-PCR mastermixes and cycling parameters”.
- In Table B.1, footnote a, “real-time PCR machines” twice replaces “real-time machines”.
- In C.1, “This primer set amplifies a product of 173 bp corresponding to nucleotides 68–240 of HAV isolate HM174 43c (GenBank accession number M59809).” is added as paragraph 2.
- In C.2, “This primer set amplifies a product of 86 bp corresponding to nucleotides 5291–5376 of Norwalk virus (GenBank accession number M87661).” is added as paragraph 2.”
- In C.3, “This primer set amplifies a product of 89 bp corresponding to nucleotides 5012–5100 of Lordsdale virus (GenBank accession number X86557).” is added as paragraph 2.”

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- In C.4, “This primer set amplifies a product of 100 bp corresponding to nucleotides 110–209 of the deletant mengo virus strain MC0 used in the development of this part of ISO/TS 15216. This corresponds to nucleotides 110–270 of the non-deletant mengo virus isolate M (GenBank accession number L22089).” is added as paragraph 2.”
- In H.5, “mastermix (if the C_q difference between EC RNA stock tested with heat-treated and untreated mastermix is <10 for a dsDNA standard curve with an idealized slope of $-3,32$), the” replaces “mastermix, the”.

ISO/TS 15216 consists of the following parts, under the general title *Microbiology of food and animal feed — Horizontal method for determination of hepatitis A virus and norovirus in food using real-time RT-PCR*:

- *Part 1: Method for quantification*
- *Part 2: Method for qualitative detection*

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Introduction

Hepatitis A virus (HAV) and norovirus (NoV) are important agents of food-borne human viral illness. No routine methods exist to culture these viruses from food matrices. Detection is therefore reliant on molecular methods using the reverse-transcriptase polymerase chain reaction (RT-PCR). As many food matrices contain substances that are inhibitory to RT-PCR, it is necessary to use an extraction method that produces highly clean RNA preparations that are fit for purpose. For food surfaces, viruses are removed by swabbing. For soft fruit and salad vegetables, virus extraction is by elution with agitation followed by precipitation with PEG/NaCl. For bottled water, adsorption and elution using positively charged membranes followed by concentration by ultrafiltration is used and for bivalve molluscan shellfish, viruses are extracted from the tissues of the digestive glands using treatment with a proteinase K solution. For all matrices which are not covered by this Technical Specification, it is necessary to validate this method. All matrices share a common RNA extraction method based on virus capsid disruption with chaotropic reagents followed by adsorption of RNA to silica particles. Real-time RT-PCR monitors amplification throughout the PCR cycle by measuring the excitation of fluorescently labelled molecules. In the 5' fluorogenic nuclease real-time RT-PCR assay, the fluorescent labels are attached to a sequence-specific nucleotide probe (hydrolysis probe) that also enables simultaneous confirmation of target template. These modifications increase the sensitivity and specificity of the PCR method, and obviate the need for additional amplification product confirmation steps post PCR. Due to the complexity of the method, it is necessary to include a comprehensive suite of controls. The method described in this part of ISO/TS 15216 enables quantification of levels of virus RNA in the test sample. A schematic diagram of the testing procedure is shown in [Annex A](#).

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Microbiology of food and animal feed — Horizontal method for determination of hepatitis A virus and norovirus in food using real-time RT-PCR —

Part 1: Method for quantification

1 Scope

This part of ISO/TS 15216 describes a method for quantification of levels of HAV and NoV genogroup I (GI) and II (GII) RNA, from test samples of foodstuffs or food surfaces. Following liberation of viruses from the test sample, viral RNA is then extracted by lysis with guanidine thiocyanate and adsorption on silica. Target sequences within the viral RNA are amplified and detected by real-time RT-PCR.

This approach is also relevant for detection of the target viruses on fomites, or of other human viruses in foodstuffs, on food surfaces or on fomites following appropriate validation and using target-specific primer and probe sets.

2 Normative references

The following referenced documents are indispensable for the application 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 22174, *Microbiology of food and animal feeding stuffs — Polymerase chain reaction (PCR) for the detection of food-borne pathogens — General requirements and definitions*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 22174 and the following apply.

3.1

foodstuff

substance used or prepared for use as food

Note 1 to entry: For the purposes of this part of ISO/TS 15216, this definition includes bottled water.

3.2

food surface

surface of food, food preparation surface or food contact surface

3.3

fomite

inanimate object or material on which infectious agents can be conveyed

3.4
hepatitis A virus
HAV

member of the *Picornaviridae* family responsible for infectious hepatitis

Note 1 to entry: Genetically, HAV can be subdivided into six genotypes on the basis of the VP1/2A region (genotypes 1, 2, and 3 have been found in humans, while genotypes 4, 5, and 6 are of simian origin). There is only one serotype.

Note 2 to entry: Transmission occurs via the faecal-oral route by person-to-person contact, through the consumption of contaminated foodstuffs, contact with contaminated water or food surfaces, or contact with contaminated fomites. Hepatitis A virus is classified as a group 2 biological agent by the European Union and as a risk group 2 human aetiological agent by the United States National Institutes of Health.

3.5
norovirus

member of the *Caliciviridae* family responsible for sporadic cases and outbreaks of acute gastroenteritis

Note 1 to entry: Genetically, norovirus can be subdivided into five separate genogroups.

Note 2 to entry: Three of these genogroups, GI, GII and GIV have been implicated in human gastrointestinal disease. GI and GII are responsible for the vast majority of clinical cases. Transmission occurs via the faecal-oral route by person-to-person contact, through the consumption of contaminated foodstuffs or through contact with contaminated water or food surfaces or contact with contaminated fomites. Genogroup I and II noroviruses are classified as group 2 biological agents by the European Union and as risk group 2 human aetiological agents by the United States National Institutes of Health.

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3.6
quantification of hepatitis A virus (standards.iteh.ai)

estimation of number of copies of HAV RNA in a predetermined mass or volume of foodstuff, or area of food surface

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3.7
quantification of norovirus

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estimation of number of copies of norovirus RNA in a predetermined mass or volume of foodstuff, or area of food surface

3.8
process control virus

virus added to the sample portion at the earliest opportunity prior to virus extraction to control for extraction efficiency

3.9
process control virus RNA

RNA released from the process control virus in order to produce standard curve data for the estimation of extraction efficiency

3.10
negative RNA extraction control

control free of target RNA carried through all steps of the RNA extraction and detection procedure to monitor any cross-contamination events

3.11
negative process control

target pathogen-free sample of the food matrix which is run through all stages of the analytical process

3.12
hydrolysis probe

fluorescent probe coupled with two fluorescent molecules which are sterically separated by the 5'-3'-exonuclease activity of the enzyme during the amplification process

3.13**negative RT-PCR control**

aliquot of highly pure water used in a real-time RT-PCR reaction to control for contamination in the real-time RT-PCR reagents

3.14**external control RNA**

reference RNA that can serve as target for the real-time PCR assay of relevance, e.g. RNA synthesized by *in-vitro* transcription from a plasmid carrying a copy of the target gene, which is added to an aliquot of sample RNA in a defined amount to serve as a control for amplification in a separate reaction

3.15 **C_q value**

quantification cycle; the PCR cycle at which the target is quantified in a given real-time PCR reaction

Note 1 to entry: This corresponds to the point at which reaction fluorescence rises above a threshold level.

3.16**theoretical limit of detection****tLOD**

level that constitutes the smallest quantity of target that can in theory be detected

Note 1 to entry: This corresponds to one genome copy per volume of RNA tested in the target assay, but varies according to the test matrix and the quantity of starting material.

3.17**practical limit of detection****pLOD**

lowest concentration of target in a test sample that can be reproducibly detected (95 % confidence interval) under the experimental conditions specified in the method, as demonstrated by a collaborative trial or other validation

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Note 1 to entry: The pLOD is related to the test portion, the quality or quantity of the template RNA, and the tLOD of the method.

3.18**limit of quantification****LOQ**

lowest concentration of target in a test sample that can be quantitatively determined with acceptable level of precision and accuracy under the experimental conditions specified in the method, as demonstrated by a collaborative trial or other validation

Note 1 to entry: The LOQ is related to the test portion and the quality or quantity of the template RNA.

4 Principle**4.1 Virus extraction**

The foodstuffs and food surfaces covered by this part of ISO/TS 15216 are often highly complex matrices and the target viruses can be present at low concentrations. It is therefore necessary to carry out matrix-specific virus extraction and/or concentration in order to provide a substrate for subsequent common parts of the process. The choice of method depends upon the matrix.

4.2 RNA extraction

It is necessary to extract RNA using a method that yields clean RNA preparations to reduce the effect of PCR inhibitors. In this part of ISO/TS 15216 the chaotropic agent guanidine thiocyanate is used to disrupt the viral capsid. RNA is then adsorbed to silica to assist purification through several washing stages. Purified viral RNA is released from the silica into a buffer prior to real-time RT-PCR.

4.3 Real-time reverse transcription polymerase chain reaction (real time RT-PCR)

This part of ISO/TS 15216 uses one step real-time RT-PCR using hydrolysis probes. In one step real-time RT-PCR, reverse transcription and PCR amplification are carried out consecutively in the same tube.

Real-time PCR using hydrolysis probes utilizes a short DNA probe with a fluorescent label and a fluorescence quencher attached at opposite ends. The assay chemistry ensures that as the quantity of amplified product increases, the probe is broken down and the fluorescent signal from the label increases proportionately. Fluorescence can be measured at each stage throughout the cycle. The first point in the PCR cycle at which amplification can be detected for any reaction is proportional to the quantity of template, therefore analysis of the fluorescence plots enables determination of the quantity of target sequence in the sample.

Due to the low levels of virus template often present in foodstuffs and the strain diversity in the target viruses, selection of fit-for-purpose one step real-time RT-PCR reagents and PCR primers and hydrolysis probes for the target viruses is important. Guidelines for their selection are given in [5.2.17](#) and [5.2.18](#). Illustrative details of reagents, primers, and probes (used in the development of this part of ISO/TS 15216) are provided in [Annexes B](#) and [C](#).

4.4 Control materials

4.4.1 Process control virus

Losses of target virus can occur at several stages during sample virus extraction and RNA extraction. To control for these losses, samples are spiked prior to processing with a defined amount of a process control virus. The level of recovery of the process control virus shall be determined for each sample.

The virus selected for use as a process control shall be a culturable non-enveloped positive-sense ssRNA virus of a similar size to the target viruses to provide a good morphological and physicochemical model. The process control virus shall exhibit similar persistence in the environment to the targets. The virus shall be sufficiently distinct genetically from the target viruses that PCR assays for the target and process control viruses do not cross-react, and shall not normally be expected to occur naturally in the foodstuffs under test.

An example of the preparation of process control virus (used in the development of this part of ISO/TS 15216) is provided in [Annex D](#).

4.4.2 Double-stranded DNA (dsDNA) control

For quantification of a target virus, results shall be related to a standard of known concentration. A dilution series of double-stranded DNA carrying the target sequence of interest ([5.3.8](#)) and quantified using spectrophotometry shall be used to produce a standard curve in template copies per microlitre. Reference to the standard curve enables quantification of the sample in detectable virus genome copies per microlitre.

4.4.3 External amplification control (EC) RNA control

Many foodstuffs contain substances inhibitory to RT-PCR, and there is also a possibility of carryover of further inhibitory substances from upstream processing. In order to control for RT-PCR inhibition in individual samples, external control (EC) RNA (an RNA species carrying the target sequence of interest, [5.3.9](#)) is added to an aliquot of sample RNA and tested using the RT-PCR method. Comparison of the results of this with the results of EC RNA in the absence of sample RNA enables determination of the level of RT-PCR inhibition in each sample under test.

Alternative approaches for RT-PCR inhibition control that can be demonstrated to provide equivalent performance to the use of EC RNA are permitted.

4.5 Test results

This method provides a result expressed in detectable virus genome copies per millilitre, per gram or per square centimetre. In samples where virus is not detected, results shall be reported as “not detected; <z detectable virus genome copies per millilitre, per gram or per square centimetre” where z is the limit of detection (LOD) for the sample.

5 Reagents

5.1 General

Use only reagents of recognized analytical grade, unless otherwise specified.

For current laboratory practice, see ISO 7218.^[10]

5.2 Reagents used as supplied

5.2.1 **Molecular biology grade water.**

5.2.2 **Polyethylene glycol (PEG)**, mean relative molecular mass 8 000.

5.2.3 **Sodium chloride (NaCl).**

5.2.4 **Potassium chloride (KCl).**

5.2.5 **Disodium hydrogenphosphate (Na₂HPO₄).**

5.2.6 **Potassium dihydrogenphosphate (KH₂PO₄).**

5.2.7 **Tris base.**

5.2.8 **Glycine.**

5.2.9 **Beef extract powder.**

5.2.10 **Proteinase K (30 U/mg).**

5.2.11 **Pectinase** from *Aspergillus niger* or *A. aculeatus*.

5.2.12 **Chloroform.**

5.2.13 **Butanol.**

5.2.14 **Sodium hydroxide (NaOH).**

5.2.15 **Hydrochloric acid (HCl).**

5.2.16 **Silica, lysis, wash, and elution buffers** for extraction of viral RNA. Reagents shall enable processing of 500 µl of extracted virus, using lysis with a chaotropic buffer containing guanidine