



SLOVENSKI STANDARD

oSIST prEN ISO 11350:2025

01-april-2025

Kakovost vode - Določanje genotoksičnosti za vodo in odpadno vodo - Salmonella/mikrosomski fluktuacijski preskus (Amesov fluktuacijski preskus)

Water quality - Determination of the genotoxicity of water and waste water -
Salmonella/microsome fluctuation test (Ames fluctuation test) (ISO 11350:2012)

Wasserbeschaffenheit - Bestimmung der Gentoxizität von Wasser und Abwasser -
Verfahren mittels Salmonella/Microsomen-Fluktuationstest (Ames-Fluktuationstest) (ISO
11350:2012)

Qualité de l'eau - Évaluation de la génotoxicité des eaux résiduaires - Essai de
Salmonella/microsome (essai d'Ames-fluctuation) (ISO 11350:2012)

Ta slovenski standard je istoveten z: prEN ISO 11350

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ICS:

13.060.70	Preiskava bioloških lastnosti vode	Examination of biological properties of water
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Water quality — Determination of the genotoxicity of water and waste water — Salmonella/microsome fluctuation test (Ames fluctuation test)

*Qualité de l'eau — Évaluation de la génotoxicité des eaux résiduaires —
Essai de Salmonella/microsome (essai d'Ames-fluctuation)*

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ISO 11350:2012(E)**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.

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 11350 was prepared by Technical Committee ISO/TC 147, *Water quality*, Subcommittee SC 5, *Biological methods*.

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Water quality — Determination of the genotoxicity of water and waste water — *Salmonella*/microsome fluctuation test (Ames fluctuation test)

WARNING — Persons using this International Standard should be familiar with normal laboratory practice. This standard does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user to establish appropriate safety and health practices and to ensure compliance with any national regulatory conditions.

IMPORTANT — It is absolutely essential that tests conducted according to this International Standard be carried out by suitably trained staff.

1 Scope

This International Standard specifies a method for the determination of the genotoxic potential of water and waste water using the bacterial strains *Salmonella enterica* subsp. *enterica* serotype Typhimurium TA 98 and TA 100 in a fluctuation assay. This combination of strains is able to measure the genotoxicity of chemicals that induce point mutations (base pair substitutions and frameshift mutations) in genes coding for enzymes that are involved in the biosynthesis of the amino acid, histidine.

NOTE 1 ISO 13829^[8] applies for the measurement of genotoxicity of samples containing DNA-crosslinking agents.

This method is applicable to:

- fresh water;
- waste water;
- aqueous extracts and leachates;
- eluates of sediments (fresh water);
- pore water;
- aqueous solutions of single substances or of chemical mixtures;
- drinking water.

NOTE 2 When testing drinking water, extraction and pre-concentration of water samples can prove necessary.

2 Normative references

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 3696, *Water for analytical laboratory use — Specification and test methods*

ISO 7027, *Water quality — Determination of turbidity*

ISO 11350:2012(E)

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

3.1

cofactor solution

aqueous solution of chemicals (e.g. NADP, glucose-6-phosphate, and inorganic salts) needed for the activity of the enzymes in the S9 fraction

[Source: ISO 21427-2:2006,^[10] definition 3.2]

3.2

culture medium

nutrients presented in a form and phase (liquid or solidified) which support microbiological growth

[Source: ISO 6107-6:2004,^[6] definition 24]

3.3

dilution level

D

denominator of the dilution coefficient (using the numerator 1) of a mixture of water or waste water with dilution water as integral number

NOTE 1 to entry: For undiluted water or waste water, this coefficient per definition is 1→1. [In this International Standard, the arrow indicates the transition from initial total volume to final total volume.] The corresponding and smallest possible value of *D* is 1.

[Source: ISO 6107-6:2004,^[6] definition 28]

3.4

lowest ineffective dilution

LID

lowest dilution within a test batch which does not show any effect, i.e. no statistically significant increase in the number of revertant wells compared with the negative control

NOTE 1 to entry: LID is determined for each incubation condition (strain, $\pm$ S9 mix). The highest LID value is decisive for the overall assessment.

3.5

induction rate

difference between the mean value of wells with revertant growth counted on the plates treated with a dose of the test sample or with a positive control, and the mean value of the corresponding wells treated with the negative control using the same strain under identical conditions

[Source: ISO 6107-6:2004,^[6] definition 43, modified: “wells with revertant growth” replaces “mutant colonies”; “corresponding wells” replaces “corresponding plates”]

3.6

inoculum

fraction of a culture of microorganisms used to start a new culture, or an exponentially growing preculture, in fresh medium

[Source: ISO 6107-6:2004,^[6] definition 44]

3.7

negative control

dilution water without test sample

[Source: ISO 6107-6:2004,^[6] definition 51]

3.8**revertant growth**

visible mutant colonies on the microplate at the end of the respective test

3.9**overnight culture**

culture started late in the afternoon and incubated overnight (usually about 16 h) to be ready during the following morning for purposes such as the inoculation of a preculture

[Source: ISO 6107-6:2004,^[6] definition 54]

NOTE 1 to entry For specification, see 9.1.

3.10**positive control**

any well characterized material and/or substance, which, when evaluated by a specific test method, demonstrates the suitability of the test system to yield a reproducible, appropriate positive or negative response in the test system

[Source: ISO 10993-12:—,^[7] definition 3.12]

NOTE 1 to entry The positive controls mentioned in this International Standard are dissolved in dimethyl sulfoxide (DMSO) prior to use. For the purposes of this International Standard, the positive controls are known mutagens which are suitable for the verification of the sensitivity of the method and/or the activity of the S9 mix.

3.11**S9 fraction**

supernatant at 9 000g of a tissue homogenate in 0,15 mol/l KCl, obtained from livers of male rats (200 g to 300 g) pretreated with a substance or substance combination appropriate for enzyme induction

[Source: ISO 6107-6:2004,^[6] definition 74]

3.12**S9 mix**

mixture of S9 fraction and cofactor solution

[Source: ISO 6107-6:2004,^[6] definition 75]

3.13**stock culture**

culture of a strain of organisms maintained under conditions to preserve original features such as nucleotide sequences

[Source: ISO 6107-6:2004,^[6] definition 87]

3.14**test sample**

undiluted, diluted or otherwise prepared portion of a sample to be tested, after completion of all preparation steps such as centrifugation, filtration, homogenization, pH adjustment and determination of ionic strength

[Source: ISO 6107-6:2004,^[6] definition 92]

4 Interferences

Bacteriotoxic effects of the test sample can lead to a reduction of viable bacteria and to a reduction of wells with revertants due to a repression of revertant growth.

This method includes sterile filtration of water and waste water prior to the test. Due to this filtration, solid particles are separated from the test sample. Thus, there is a possibility that genotoxic substances adsorbed on particles are not detected.

5 Principle

The bacteria are exposed under defined conditions to various concentrations of the test sample and incubated for 100 min at $37\text{ °C} \pm 1\text{ °C}$ in 24 well plates. Due to this exposure, genotoxic agents enclosed in the test sample can induce mutations in one or both marker genes of the bacterial strains used (hisG46 for TA 100 and hisD3052 for TA 98) in correlation with the applied concentrations. Induction of mutations causes a concentration-related increase in the number of mutant colonies.

After exposure of the bacteria, reversion indicator medium (7.40), containing the pH indicator dye bromocresol purple (7.7), is added to the wells. Subsequently, the batches are distributed to 384 well plates (48 wells for each parallel) and incubated for 48 h to 72 h (9.3.2, 9.3.3).

Mutagenic activity of the test sample is determined by counting the number of purple to yellow shifted wells (per 48 wells of each parallel), treated with the undiluted or the diluted test sample, compared to the negative control.

The lowest dilution ($1 \rightarrow N$) of the test sample which induces no mutagenic effect under all experimental conditions (if any mutagenic effect is induced by the test sample) is the criterion for evaluating the mutagenic potential. Sample dilutions above this ($1 \rightarrow A$, $A < N$) shall induce a mutagenic effect according to the criteria of this International Standard in at least one strain under at least one activation condition (with or without addition of S9 mix). The respective LID value is N . If no mutagenic effect is observed under all experimental conditions, this dilution is $1 \rightarrow 1$ and the respective LID value is 1.

6 Apparatus and materials

6.1 Temperature- and time-controlled incubator, $37\text{ °C} \pm 1\text{ °C}$.

6.2 pH meter.

6.3 Analytical balance.

6.4 Steam sterilizer.

6.5 Dry sterilizer.

6.6 Magnetic stirrer.

6.7 Rotary mixer.

6.8 Freezer, capable of being maintained at $\leq -18\text{ °C}$ and at $\leq -70\text{ °C}$.

6.9 Pipettes, 0,1 ml, 0,5 ml, 1 ml, 2 ml, 5 ml, 10 ml and 25 ml, of glass or plastics.

6.10 Storage bottles, 250 ml and 1 000 ml.

6.11 Measuring cylinders, 100 ml and 200 ml.

6.12 Volumetric flasks, 20 ml, 200 ml and 500 ml.

6.13 Sterile filters, 0,2 μm and 0,45 μm .

6.14 Erlenmeyer flasks, 50 ml, 100 ml and 250 ml.

6.15 Inoculating loops.

6.16 Eight-channel multistepper pipette (repeater pipette).

6.17 Eight-channel pipettes, 5 µl to 50 µl and 50 µl to 300 µl.

6.18 Spectrophotometer.

6.19 Transparent sterile polystyrene 24 well and 384 well plates with flat bottom and lid.

6.20 Microplate photometer for 24 well plates and optionally for 384 well plates, filters: 420 nm ± 15 nm and 595 nm ± 10 nm.

6.21 Clean bench.

6.22 Petri dishes with venting ribs, diameter approximately 94 mm, height approximately 16 mm.

6.23 Cryogenic vials, sterile, 1 ml, 10 ml.

7 Reagents, media and dilutions

7.1 General. As far as possible, use “reagent grade” chemicals. If hydrates of anhydrous compounds or hydrates different from those specified are used, ensure that the appropriate mass of the main compound is employed.

When necessary, autoclave for 20 min at 121 °C ± 2 °C. Cover vessels loosely (e.g. with aluminium foil). Never seal air-tight.

7.2 Water, grade 1, as defined in ISO 3696, or water with a conductivity of ≤5 µS/cm.

If sterile water is needed, sterilize by sterile filtration (0,2 µm) or autoclaving. Water as specified here is also used for the stepwise dilution of the test sample.

7.3 Tester strains. Use mutant strains of *Salmonella* Typhimurium LT2, which enable detection of point mutations, to determine the mutagenic potential of a test sample. Since point mutations can be subdivided into two classes (frameshift mutations and base pair substitutions), the two tester strains TA 98 and TA 100 are used. TA 98 contains as a marker the frameshift mutation (+2 type) hisD3052, whereas TA 100 bears the base pair substitution hisG46.

In addition, both strains shall have the following genetic properties:

- they contain the plasmid pKM101, coding for ampicillin resistance;
- they are all deep rough, e.g. partly deficient in lipopolysaccharide side chains, enabling also larger molecules to penetrate the bacterial cell wall and to cause mutations;
- due to a mutation in *uvrB*, the capability of the tester strains to repair DNA-damage is limited and the likelihood that DNA-damage results in mutations is increased.

NOTE The use of additional tester strains is described in Annex K.

7.4 2-Aminoanthracene (2-AA), C₁₄H₁₁N, CAS No: 613-13-8.

7.5 Ampicillin sodium salt, C₁₆H₁₈N₃NaO₄S, CAS No: 69-52-3.

7.6 D-Biotin, C₁₀H₁₆N₂O₃S, CAS No: 58-85-5.

7.7 Bromocresol purple, sodium salt, CAS No: 62625-30-3.

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- 7.8 Citric acid monohydrate**, $C_6H_8O_7 \cdot H_2O$, CAS No: 5949-29-1.
- 7.9 Dimethylsulfoxide**, DMSO, C_2H_6SO , CAS No: 67-68-5.
- 7.10 D-Glucose**, anhydrous, $C_6H_{12}O_6$, CAS No: 50-99-7.
- 7.11 D-Glucose-6-phosphate disodium salt hydrate**, G-6-P- Na_2 , $C_6H_{11}Na_2O_9P \cdot 2H_2O$ CAS No: 3671-99-6.
- 7.12 Hydrochloric acid solution**, HCl, $c(HCl) = 1 \text{ mol/l}$.
- 7.13 Magnesium chloride hexahydrate**, $MgCl_2 \cdot 6H_2O$, CAS No: 7791-18-6.
- 7.14 Magnesium sulfate heptahydrate**, $MgSO_4 \cdot 7H_2O$, CAS No: 10034-99-8.
- 7.15 Potassium chloride**, KCl, CAS No: 7447-40-7.
- 7.16 Dipotassium hydrogenphosphate**, K_2HPO_4 , CAS No: 7758-11-4.
- 7.17 Sodium ammonium hydrogenphosphate tetrahydrate**, $NaNH_4HPO_4 \cdot 4H_2O$, CAS No: 7583-13-3.
- 7.18 Sodium chloride**, NaCl, CAS No: 7647-14-5.
- 7.19 Sodium dihydrogenphosphate**, anhydrous, NaH_2PO_4 , CAS No: 7558-80-7.
- 7.20 Disodium hydrogenphosphate**, anhydrous, Na_2HPO_4 , CAS No: 7558-79-4.
- 7.21 Sodium hydroxide solution**, $c(NaOH) = 1 \text{ mol/l}$.
- 7.22 β -Nicotinamide adenine dinucleotide phosphate sodium salt**, $NADP \cdot H_2O$, $C_{21}H_{27}N_7NaO_{17}P_3 \cdot H_2O$, CAS No: 698999-85-8.
- 7.23 Nitrofurantoin (NF)**, CAS No: 67-20-9.
- 7.24 4-Nitro-o-phenylenediamine (4-NOPD)**, CAS No: 99-56-9.
- 7.25 Nutrient broth powder**.¹⁾
- 7.26 S9 fraction** (liver homogenate; induced by phenobarbital/ β -naphthoflavone).¹⁾
- 7.27 L-Histidine**, $C_6H_9N_3O_2$, CAS No: 71-00-1.
- 7.28 Phosphate buffer.**
- 7.28.1 Sodium dihydrogenphosphate buffer**, $c(NaH_2PO_4) = 0,2 \text{ mol/l}$.
- Dissolve 14,39 g NaH_2PO_4 (or 16,55 g $NaH_2PO_4 \cdot H_2O$) in 600 ml of water (7.2).

1) This reagent is commercially available. This information is given for the convenience of users of this International Standard and does not constitute an endorsement by ISO of these products.