

SLOVENSKI STANDARD oSIST prEN ISO 21285:2020

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Kakovost tal - Zaviranje razmnoževanja Hypoaspis aculeifer zaradi onesnaževal v tleh (ISO 21285:2019)

Soil quality - Inhibition of reproduction of the soil mite (Hypoaspis aculeifer) by soil contaminants (ISO 21285:2019)

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Qualité du sol - Inhibition de la reproduction de l'acarien prédateur (Hypoaspis aculeifer) par des contaminants du sol (ISO 21285:2019)

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Soil quality — Inhibition of reproduction of the soil mite (*Hypoaspis aculeifer*) by soil contaminants

Qualité du sol — Inhibition de la reproduction de l'acarien prédateur (Hypoaspis aculeifer) par des contaminants du sol

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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).

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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 190, *Soil quality*, Subcommittee SC 4, *Biological characterization*.

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.

Introduction

Ecotoxicological test systems are applied to obtain information about the effects of contaminants in soil and are proposed to complement conventional chemical analysis (see ISO 15799 and ISO 17616). ISO 15799 includes a list and short characterization of recommended and standardized test systems and ISO 17616 gives guidance on the choice and evaluation of the bioassays. Aquatic test systems with soil eluate are applied to obtain information about the fraction of contaminants potentially reaching the groundwater by the water path (retention function of soils), whereas terrestrial test systems are used to assess the habitat function of soils.

Mites (Acari) are a world-wide and diverse group of arthropods belonging to the class Arachnida with over 40 000 species recorded, divided into two super-orders (Acariformes and Parasitiformes). Due to their relative small size (a few μ m to a few cm), they occupy specific ecological niches on plants as well as in soils (see Reference [13]).

Among soil-inhabiting mites, the role of predation is ensured by, for example, *Hypoaspis* sp. (Laelapidae). Because they are exposed to chemical contamination, mites are already considered in the environmental risk assessment of pesticides, as non-target organisms (see Reference [10]). Indeed, among the data required for active substances of pesticides, effects on predatory mites are assessed, i.e. for the plant-inhabitant *Typhlodromus pyri* (Phytoseiidae) and the soil-inhabitant *Hypoaspis aculeifer* (Laelapidae) (see Reference [6]).

The first authors introducing *H. aculeifer* as a test organism in ecotoxicological studies [23][17] were later proposed a two-species test system in the European project SECOFASE (Sublethal Effects of Chemicals on Fauna in the Soil Ecosystem), including the collembolan *Folsomia fimetaria* as prey. In the context of the development of an ecotoxicological test for the assessment of plant protection products on nontarget arthropods (see References [5][6]), a protocol on soil predatory mites using *H. aculeifer* was further proposed. More recently, a standard test protocol for the assessment of chemicals was developed for this species by OECD in 2008 and revised in 2016. The results of the associated international ringtest were published in Reference [25].

Among mites, the predator *Hypoaspis aculeifer* is the most studied species in laboratory. The reproduction end point was found in general to be more sensitive than mortality and avoidance. Compared to other soil meso-fauna invertebrates, mites were found in general less sensitive than or as sensitive as other test species, depending on the end points and chemicals studied. Considering semi-field studies, *H. aculeifer* was used as a top predator whereas other soil invertebrates, mainly springtails, were ranked in the grazer group. In these studies, mites showed to be quite tolerant towards anthropogenic contamination. This statement was also corroborated by field surveys. However, the applicability of laboratory test methods for the assessment of environmental samples (contaminated soils, wastes etc.) with mites is emphasized, as to date a limited number of studies are available.

This document describes a method that is based on the determination of lethal and sublethal effects of contaminated soils to adult predatory mites of the species *Hypoaspis aculeifer*. This species is considered to be representative of predatory soil arthropods. Background information on the ecology of these mites and their use in ecotoxicological testing is available in Reference [14].

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Soil quality — Inhibition of reproduction of the soil mite (*Hypoaspis aculeifer*) by soil contaminants

1 Scope

This document specifies a chronic test method for evaluating the habitat function of soils and determining effects of soil contaminants and substances on the reproduction of *Hypoaspis aculeifer* by – mainly – alimentary uptake. This method is applicable to soils and soil materials of unknown quality, e.g. from contaminated sites, amended soils, soils after remediation, industrial, agricultural or other sites under concern and waste materials (e.g. dredged material, municipal sludge from a wastewater treatment plant, composed material, or manure, especially those for possible land disposal). The reproduction (= number of juveniles) is the measured parameter of the test. The test reflects the bioavailability of a mixture of contaminants in natural soils (contaminated site soils) to a species which represents a trophic level which is not covered by other ISO standards. This test is not intended to replace the earthworm (see ISO 11268-2) or Collembola (see ISO 11267) reproduction tests since this species belongs not only to a different trophic group but also a different taxonomic group (= mites; i.e. arachnids) than those used usually.

Effects of substances are assessed using a standard soil, preferably a defined artificial soil substrate. For contaminated soils, the effects are determined in the soil to be tested and in a control soil. Depending on the objective of the study, the control and dilution substrate (dilution series of contaminated soil) are either an uncontaminated soil comparable to the soil to be tested (reference soil) or a standard soil (e.g. artificial soil).

This document provides information on how to use this method for testing samples (soils or substances) under temperate conditions. <u>SIST EN ISO 21285:2020</u>

This document is not applicable to substances for which the air/soil partition coefficient is greater than one, or to substances with vapour pressure exceeding 300 Pa at 25 °C.

NOTE The stability of the test substance cannot be ensured over the test period. No provision is made in the test method for monitoring the persistence of the substance under test.

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 10390, Soil quality — Determination of pH

ISO 10694, Soil quality — Determination of organic and total carbon after dry combustion (elementary analysis)

ISO 11260, Soil quality — Determination of effective cation exchange capacity and base saturation level using barium chloride solution

ISO 11277, Soil quality — Determination of particle size distribution in mineral soil material — Method by sieving and sedimentation

ISO 11465, Soil quality — Determination of dry matter and water content on a mass basis — Gravimetric method

ISO 18400-206, Soil quality — Sampling — Part 206: Collection, handling and storage of soil under aerobic conditions for the assessment of microbiological processes, biomass and diversity in the laboratory

3 Terms and definitions

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

contaminant

substance or agent present in the soil as a result of human activity

3.2

effect concentration for x % effect

 EC_X

concentration (mass fraction) of a test sample that causes x % of an effect on a given end point within a given exposure period when compared with a control

EXAMPLE An EC_{50} is a concentration estimated to cause an effect on a test end point in 50 % of an exposed population over a defined exposure period.

Note 1 to entry: The ECx is expressed as a percentage of soil to be tested (dry mass) per soil mixture (dry mass). When substances are tested, the ECx is expressed as the mass of the test substance per dry mass of soil in milligrams per kilogram.

3.3

effect rate

 ER_x

rate of a soil to be tested that causes an x % of an effect on a given end point within a given exposure period when compared with a control

3.4 https://standards.iteh.ai/catalog/standards/sist/4cabe9d5-3202-40c7-87fd

limit test

single concentration test consisting of at least four replicates each, the soil to be tested without any dilution or the highest concentration of test substance mixed into the control soil and the control

3.5

lowest observed effect concentration

LOEC

lowest test substance concentration that has a statistically significant effect (probability p < 0.05)

Note 1 to entry: In this test, the LOEC is expressed as a mass of test substance per dry mass of the soil to be tested. All test concentrations above the LOEC should usually show an effect that is statistically different from the control.

3.6

lowest observed effect rate

LOER

lowest rate of a soil to be tested in a control soil at which a statistically significant effect is observed

3.7

no observed effect concentration

NOEC

highest test substance concentration immediately below the LOEC (3.5) at which no effect is observed

Note 1 to entry: In this test, the concentration corresponding to the NOEC, has no statistically significant effect (probability p < 0.05) within a given exposure period when compared with the control.

3.8

no observed effect rate

NOER

lowest rate of a soil to be tested immediately below the *LOER* (3.6) which when compared to the control has no statistically significant effect (probability p < 0.05) within a given exposure period

3.9

reference soil

uncontaminated soil with comparable pedological properties (nutrient concentrations, pH, organic carbon content and texture) to the soil being studied

3.10

standard soil

field-collected soil or artificial soil whose main properties (pH, texture, organic matter content) are within a known range

EXAMPLE Euro soils, artificial soil, LUFA standard soil type 2.2.

Note 1 to entry: The properties of standard soils can differ from the soil to be tested.

3.11

control soil

reference or standard soil used as a control and as medium for preparing dilution series with soils to be tested or a reference substance, which fulfils the validity criteria

Note 1 to entry: In the case of natural soil, it is advisable to demonstrate its suitability for a test and for achieving the test validity criteria before using the soil in a definitive test.

3.12

test mixture

mixture of contaminated soil or the test substance (e.g. chemical, biosolid, waste) with control soil

3.13

test mixture ratio

ratio between the soil to be tested and the control soil in a test mixture

4 Principle

Adult females are exposed to the soil to be tested and the effects on reproduction measured are compared to those observed for females exposed to a control soil. If appropriate, effects based on exposure to a dilution range of contaminated soil and control soil or a range of concentrations of a test substance mixed into control soil are determined. Test mixtures are prepared at the start of the test and are not renewed within the test period. The test is started with 10 adult females per test vessel. Males are not introduced in the test, because experience has shown that females mate immediately or shortly after hatching from the deutonymph stage, if males are present. As the females are introduced into the test about 7 d after they have reached the adult stage, the females can be considered as already mated (Annex A and Annex E). The test runs until the first offspring have reached the deutonymph stage. At 20 °C the exposure time ends at day 14 after introducing the females (day 0), followed by two days of extraction. The number of surviving females and the number of juveniles per test vessel are determined. The reproductive output of the mites exposed to the test mixtures is compared to that of the controls in order to determine the concentrations which cause no effects on mortality and reproduction (NOER/NOEC) and the concentration resulting in x % reduction of juveniles hatched from eggs compared to the control (ER $_x$ /EC $_x$) respectively, depending on the experimental design (see 7.1.3).

In case there is no prior knowledge of the dilution/concentration of the soil to be tested or the test substance likely to have an effect, then it is useful to conduct the test in two steps:

 A range-finding test on reproduction is carried out, to give an indication of the effect dilution/ concentration, and the dilution/concentration giving no mortality (NOER/NOEC). Dilutions/ concentrations to be used in the definitive test can then be selected;