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Profenofos pesticides — Specification —

Part 3; Ultra low volume liquids (ULV)

ICS 65.100.10

Reference number

DRS 618-3: 2025

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Foreword

Rwanda Standards are prepared by Technical Committees and approved by Rwanda Standards Board (RSB) Board of Directors in accordance with the procedures of RSB, in compliance with Annex 3 of the WTO/TBT agreement on the preparation, adoption and application of standards.

The main task of technical committees is to prepare national standards. Final Draft Rwanda Standards adopted by Technical committees are ratified by members of RSB Board of Directors for publication and gazettment as Rwanda Standards.

DRS 618-3 was prepared by Technical Committee RSB/TC 64, Pesticides.

In the preparation of this standard, reference was made to the following standard:

ES 771-3: Pesticides — Profenofos — Part 3: Ultra low volume liquids (ULV)—Specification

The assistance derived from the above source is hereby acknowledged with thanks.

DRS 618 consists of the following parts, under the general title *Profenofos pesticides* — *Specification*:

- Part 1: Technical material
- Part 2: Emulsifiable concentrate (EC)
- Part 3: Ultra low volume liquids (ULV)

Committee membership

The following organizations were represented on the Technical Committee on *Pesticides* (RSB/TC 64) in the preparation of this standard.

Rwanda Food and Drugs Authority

Rwanda Forensic Institute

University of Rwanda/College of Sciences and Technology (UR/CST)

University if Rwanda/College of Agriculture, Animal Sciences and Veterinary Medicine (UR/CAVM)

Standards of Sustainability

CYIRA Ltd

Rwanda Inspectorate, Competition and Consumer Protection Authority (RICA)

Rwanda Investigation Bureau (RIB)

Rwanda Agriculture and Inputs Organization (RAIDO)

VeR Alka Group Ltd

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Introduction

Profenofos is an organophosphate insecticide. It is a liquid with a pale yellow to amber color and a garlic-like odor. Profenofos can be used on a variety of crops including cotton and vegetables such as maize, potato, soybean, and sugar beet. Mixed with phoxim, cypermethrin, beta-cypermethrin imidacloprid and deltamethrin, profenofos can be used against Cotton MealyBug, cabbage caterpillar, Plutella xylostella and asparagus caterpillars, as well as against wheat and cabbage aphids.

Like other organophosphates, the profenofos mechanism of action is via the inhibition of the acetylcholinesterase enzyme. Although it is used in the form of a racemate, the S(-) isomer is a more potent inhibitor. Profenofos can be synthesized by reacting phosphorus oxychloride with sodium ethoxide and

the S(-) isor, with sodium of the solution of

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Profenofos pesticides — Specification — Part 3: Ultra low volume liquids (ULV)

1 Scope

This Draft Rwanda Standard specifies the requirements, sampling and test methods for technical material of profenofos pesticides in form of Ultra Low Volume liquids (ULV) meant for plant protection purpose.

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.

RS 405, Pesticides - Sampling

RS 406, Pesticides — Terminology

DRS 618-1, Profenofos Pesticides — Specification — Part 1: Technical material

RS 565-2, Packaging of Pesticides — Requirements — Part 2: Liquid pesticides

RS 578, Pesticides — Guidelines on good labelling practices

RS 579, Pesticides — Guidelines for retail, distribution, storage and handling

RS 210, Safety procedures for the disposal of surplus pesticides and associated toxic waste — Code of practice

RS ISO 2719, Determination of flash point — Pensky-Materns closed cup method

3 Terms and definitions

For the purposes of this standard, the terms and definitions given in RS 406 apply.

4 Requirements

4.1 General requirements

4.1.1 The product shall consist of technical profenofos, complying with the requirements on DRS 618-1, together with other necessary formulants.

4.1.2 The profenofos content shall be declared in (g/kg or g/l at 20 \pm 0.5 °C and, when determined in accoradance with annex A, the content obtained shall not differ from that declared by more than the following amounts:

Table 1—Permitted tolerance on declared content

Declared content	Permitted tolerance
Above 100 up to 250 g/kg or g/l	± 6% of the declared content
Above 250 up to 500 g/kg or g/l	± 5% of the declared content
Above 500 g/kg or g/l	± 2.5% of the declared content

NOTE If the buyer requires both g/kg and g/l at 20 °C, then in case of dispute the analytical result shall be calculated as g/kg.

- 4.1.3 The product shall be in the form of a stable liquid, free from visible suspended sediment.
- **4.1.4** The product shall be readily for use through ULV equipment.

4.2 Specific requirements

The product shall comply with the requirements given in Table 1 when tested in accordance with the test methods prescribed therein.

Table 1 — Specific requirements for profenofos emulsifiable concentrates pesticides

S/N	Par	rameters	Requirements	Test methods
i.	Profenofos content,	% by mass	As per 4.1.2	Annex A
ii.	pH range	1,10	3.0 – 7.0	Annex B
iii.	Flash point, °C, min		Not lower that declared value	RS ISO 2719
iv.	Storage stability	At 0 ± 1°C for 7 days, ml, max.	0.3	Annex C
		At 54 ± 2°C for 14 days, % m/m, min.	96	Annex D

5 Packaging

The product shall be packaged in accordance with RS 565-2.

6 Labelling and marking

The product shall be labelled and marked in accordance with DRS 578.

7 Retail, distribution, storage and handling

The product shall be handled in accordance with DRS 579

NOTE Attention is drawn to the appropriate national and/ or international regulations on the handling and transport of flammable materials.

8 Sampling

Sampling shall be done in accordance with RS 405.

Disposal

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Annex A (normative)

Determination of profenofos content

A.1 General

Profenofos is separated from other components by gas chromatography on a packed column with flame ionization detection and internal standardization.

A.2 Reagents

- A.2.1 Acetone
- A.2.2 Profenfos of known purity; better than 980 g/kg
- A.2.3 Di-(2-ethylhexyl) adipate internal standard

A.2.4 Internal standard solution

Dissolve di-(2-ethylhexyl) adipate (980 to 1020 mg) in acetone (500 ml)

A.2.5 Calibration solution

Weight (to the nearest 0.1 mg) profenoros standard (90 to 100 mg) into a glass-stoppered conical flask (50 ml), add by pipette internal standard solution (25.0 ml), dilute to the mark with acetone and mix well. Prepare in duplicate.

A.3 Apparatus

- A.3.1 Gas chromatograph Fitted with a flame ionization detector
- A.3.2 Column Glass, 1.8 m X 2 mm (i.d.) packed with 3% OV-210 on supercoport, 80-100 mesh
- A.3.3 Electronic integrator or data system
- A.3.4 Microsyringe 10 μl

A.4 Procedure

A.4.1 Identity test

Use the GLC method below. The retention time of profenofos in the sample solution should not deviate by more than 0.2 min from that of the calibration solution.

A.4.2 Profenofos content

A.4.2.1 Operating conditions (typical):

Column

Packing

Column temperature

Injector temperature

Detector temperature

Carrier gas

Injection volume

Retention time profenofos: 6.5 min.

> Internal standard 11.4 min.

A.4.2.2 Linearity check

Check the linearity of the detector response by injecting 1 µl of solutions with profenofos with concentrations 0.5, 1 and 2 times that of the concentration of the calibration solution. Be sure that the concentrations of the solutions are in the linear range of the detector, otherwise alter the weighings or the dilutions accordingly. Inject each solution at least twice and determine the response factor (f). the single values should differ by less than 0.5 % from the mean value, otherwise repeat the calibration.

A.4.2.3 Preparation of sample

Weigh (to the nearest 0.1 mg) into a glass stoppered conical flask (50 ml) sufficient sample to contain 90 to 100 mg profenofos (w mg). add by pipette internal standard solution (25.0 ml), dilute to the mark with acetone and mix well. Prepare in duplicate.

A.4.2.4 Determination

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Inject each sample solution in duplicate into the gas chromatograph and bracket a series of sample solutions by duplicate injections of the calibration solutions as follows: calibration solution 1 (double injection) calibration solution 2 (double injection), and so on. Measure the relevant peak areas.

Average the response factors of each double injection. Calculate the mean value of each pair of response factors bracketing the injections of two samples and use this value for calculating the profenofos contents of the bracketed sample injections. mnentsonly

A.4.2.4 Calculation

$$f = \frac{Ir \, X \, s \, X \, P}{Hs}$$

Content of profenofos =
$$\frac{f \ X \ Hw}{I4 \ X \ W} \ g/kg$$

Where,

- is response factor;
- Hs is area of profenofos peak in the calibration solution;
- Hw is area of profenofos peak in the sample solution;
- is area of internal standard peak in the calibration solution;
- is area of internal standard peak in the sample solution;
- is mass of profenofos in the calibration solution (mg);
- is mass of sample taken (mg)
- is purity of profenofos reference substance (g/kg)

Annex B (normative)

Determination of pH value

B.1 Outline of the method

The pH value of a liquid is determined by means of pH meter and a glass electrode.

B.2 Reagents

- **B.2.1** Potassium hydrogen phthalate (COOH-C₀H₄-COOK) 0.05 mol/I (0.05M) Dissolve 10.21 g in freshly boiled distilled water and make up to 1000 ml. do no keep the solution for longer than one month.
- B.2.2 Disodium tetraborate (Na₂B₄O₇.10H₂O 0.05M Dissolve 19.07 g in freshly boiled distilled water and make up to 1000 ml. do no keep the solution for longer than one month.
- **B.2.3** Water Freshly boiled and cooled distilled water of pH 5.5 to 7.0

B.3 Apparatus

- B.3.1 pH meter
- B.3.2 Glass electrode and reference electrode

B.4 Procedure

Operate the pH meter and electrode system in accordance with the manufacturer's instructions. Standardize the meter and electrodes with the 0.05M phthalate (pH 4.00) when an acid solution is being measured or 0.05M borate when an alkaline solution is being measured (see Table B1). The reading should not differ by more than 0.02 pH units from the original value at which the apparatus was standardized. If the difference is greater than 0.05, then repeat the measurements.

Table B1 – pH values of 0.05M disodium tetraborate Temperature, °C	10	15	20	25	30
pH	9.32	9.28	9.22	9.18	9.14

B.5 pH of aqueous dispersion

Weigh 1 g of sample, transfer to the measuring cylinder containing water (about 50 ml), make up to 100 ml with water, and shake vigorously for 1 min. allow any suspension to settle for 1 min and then measure the pH of the supernatant liquid.

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Annex C (normative)

Determination of stability of liquid formulations at 0 °C

C.1 Outline of the method

A sample is maintained at 0 °C for 1 h and the volume of any separated solid or oily matter is then recorded. Storage at 0 °C is continued for 7 days, any solid matter is settled by centrifuging and its volume recorded.

C.2 Emulsifiable concentrates and solutions

C.2.1 Apparatus

- C.2.1.1 Refrigerator Capable of maintaining a temperature at 0 ± 1 °C.
- C.2.1.2 Cone shaped centrifuge tubes, 100-mL.
- C.2.1.3 Centrifuge equipped with buckets capable of holding the specified tubes.
- C.2.1.4 Pipette, 100-mL.

NOTE A domestic refrigerator is often unsuitable because the on/off cycle covers a range greater than 2 °C.

C.2.2 Procedure

Transfer 100 \pm 1.0 mL of a sample of the product to a centrifuge tube. Cool the tube and its content to (0 \pm 1) °C in the refrigerator. Allow the tube and its contents to remain at (0 \pm 1) °C for 1 h, and during this time stir the contents of the tube at intervals of approximately 15 min, each time for approximately 30 s. After this period examine the tube and record whether any solid or oily matter is present. Replace the tube in the refrigerator and allow it to remain at (0 \pm 1) °C for a total period of 7 days.

At the end of 7 days, remove the tube from the refrigerator, and allow it to remain undisturbed at room temperature for 3h. invert the centrifuge tube once, and centrifuge for 15 min at such a speed that the relative centrifugal force (RCF) at the tips of the tubes is about $550 \times G$ (the acceleration due to gravity = 981 cm/s⁻².

Record the volume of any separated material at the bottom of the tube to the nearest 0.005 mL.

NOTE 2 : RCF =
$$\frac{(rpm)^2 d}{179000}$$
 and rpm = $\sqrt{98.45} X d^{-1} X 10^3$

Where:

RCF is relative centrifuge force;

d is diameter of swing (in cm) measured from the tips of the opposite tubes when in the position occupied during the centrifuging.

If the liquid phase is not homogenous, record the volume of each layer.

C.3 Aqueous solutions

C.3.1 Apparatus

C.3.1.1 Measuring cylinder, 100-mL.

C.3.1.2 Refrigerator, at 0 ± 1 °C

C.3.2 Procedure

Put 100 ml of the product in the measuring cylinder and then put it in the refrigerator for 48 h at 0 ± 1 °C. At the end of this time, note the amount of separated material, if any, then allow the cylinder to reach room temperature and again note the amount of separated material.

Annex D (normative)

Determination of accelerated storage stability

D.1 Outline of the method

Representative sample is stored in a screw-capped bottle in an oven at a specified temperature and time.

D.2 General method

As this is intended as a model procedure, temperature and times specified are examples only since the parameters will normally be given for individual pesticide formulations.

D.3 Apparatus

- D.3.1 Beaker 250-mL, 6 to 6.5 internal diameter.
- **D.3.2 Metal disc** Plastic coated; a loose fit in the beaker, and of such dimensions that an even pressure of 25 g/cm² can be produced on the surface of the sample in the beaker.

NOTE Alternatively, a close fitting cylinder with a flat bottom, containing lead shot, can be used, the lead shot may be sealed in with molten wax so as ti give the correct weight, and prevent the shot from being lost.

- **D.3.3** Oven Thermostatically controlled to the specified temperature (± 2 °C)
- D.3.4 Desiccator without desiccant

D.4 Procedure

Put the sample into the beaker and spread it, without using any pressure, in a smooth even layer of constant thickness. Place the disc on the surface of the solution in the beaker, and put in the oven. After the specified time remove the beaker, take out the disc, and allow the beaker to cool in the desiccator.

NOTE. Use the specified temperature and time given in the specification of method of analysis. If no temperature of time is specified, store the sample at 54 ± 0.2 °C for 14 days.

Ensure that each sample taken is truly representative of that left in the beaker. Sampling of a hard cake may be carried out conveniently by removing several cores with a small diameter (6 mm) cork borer.

Bibliography

[1] ISO/IEC Directives, Part 2, Rules for the structure and drafting of International Standards, 2016

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