

**OCHRATOXIN-A IN WINE
FLOW-THROUGH ASSAY**
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A flow-through enzyme immunoassay for
the detection of Ochratoxin-A in wine

**EUROPROXIMA OCHRATOXIN-A IN WINE
FLOW-THROUGH RAPID TEST**

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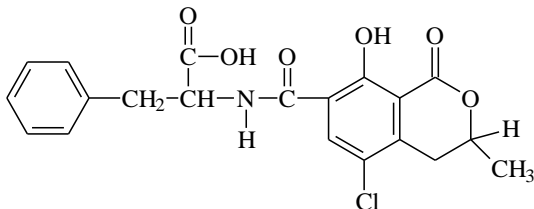
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BRIEF INFORMATION

The Ochratoxin-A (OTA) Flow-Through Rapid Test is a competitive membrane-based enzyme immunoassay for the detection of OTA in red, white and rosé wines. The test is based on mouse monoclonal antibodies against OTA. With this kit 10 analyses can be performed. The kit contains all the reagents required to perform the tests. The cut-off value of this test is 1 µg/l of OTA in wine.

1. INTRODUCTION



OCHRATOXIN A

Ochratoxin A (OTA) is a nephrotoxic and hepatocarcinogenic mycotoxin produced by *Penicillium verrucosum* and *Penicillium viridicatum* in temperate and cold climates and by a number of *Aspergillus* species such as *A. ochraceus* in warmer and tropical areas of the world. OTA has been shown to occur in various cereals and plant products such as coffee and grapes. Therefore, OTA also occurs in wine. In the European Union the maximum limit for OTA in wine is 2 µg/L in accordance with Commission Regulation 1881/2006.

2. PRINCIPLE OF THE OCHRATOXIN-A FLOW-THROUGH ASSAY

The membrane-based assay kit consists of 10 devices each pre-coated with OTA-BSA (test line) anti-mouse IgG (control line). Specific antibodies (mouse anti-OTA) labelled with horseradish peroxidase (HRP-antibody conjugate) are supplied lyophilised in 10 separate vials. Wine extract is added to these reaction vials and pre-incubated for 5 minutes. During this incubation specific antibodies bind any OTA present in the sample. Then the mixture is transferred onto the membrane where any unbound antibodies bind to OTA-BSA on the test line and bound/unbound antibodies bind to the anti-mouse IgG on the control line. After the sample extract has been completely absorbed through the membrane the unbound HRP-antibody conjugate is removed by a washing step. A chromogen substrate (tetramethylbenzidine, TMB) is then added. Bound HRP transforms the chromogen substrate into a blue coloured product and this appears as a line. Two minutes after the addition of the substrate the background membrane colouration is washed away with a final washing step. The results are then visually interpreted.

3. HANDLING AND STORAGE

- Store the kit at 2°C to 8°C in a dark place.
- After the expiry date has passed, the kit cannot be used.
- Store the remaining devices in the resealable zip lock bag and refrigerate. Before opening the kit, let it reach ambient temperature.
- Avoid direct light on the substrate solution.
- If a blue colouring of the substrate solution is observed, it may indicate a degeneration and the component cannot be used for the test.

4. KIT CONTENTS

- 2 x 5 membrane devices
- 10 filters
- 10 syringes
- 20 Pasteur pipettes (1 ml)
- 10 vials with wine extraction buffer (Extract)
- 10 reaction vials (React, black cap)
- 1 vial with wash buffer (Wash, white cap)
- 1 vial with substrate solution (Colour, blue cap)

5. EQUIPMENT AND MATERIALS REQUIRED BUT NOT PROVIDED

- Pipette 100 – 1000 µl (recommended) (alternatively use 1 ml Pasteur pipettes provided in the kit)
- Vials for sample filtrate collection (glass or polypropylene tube, vial, cup, container; volume at least 3 ml)
- Timer/clock
- Camera/smartphone for taking photos of the results (if required)
- Gloves

6. PRECAUTIONS

- This kit may contain hazardous substances. For hazard notes please refer to the appropriate safety data sheets (SDS).
- Avoid contact of all biological materials with skin and mucous membranes.
- Do not pipette by mouth.
- Do not eat, drink, smoke, store or prepare foods, or apply cosmetics within the designated work area.
- Do not use components past expiration date and do not use components from different lots.
- Optimal results will be obtained by strict adherence to this protocol. Careful pipetting and washing throughout this procedure are necessary to maintain good precision and accuracy.

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7. PREPARATION OF REAGENTS

Before starting the test, the reagents should be brought up to ambient temperature. Any reagent not used should be put back into storage immediately at 2°C to 8°C. All the reagents are provided ready to use.

8. SAMPLE PREPARATION AND ASSAY PROCEDURE

Sample preparation

1. Add 1 ml of wine to an extraction buffer vial (Extract). Use a Pasteur pipette provided in the kit to measure 1 ml or use a single channel pipette (100 μ l – 1000 μ l). Shake by hand for 1 minute.
2. Remove a plunger from a syringe and attach a filter to the syringe.
3. Mix the content of the extraction vial well and pour quickly into the syringe. Place the plunger back onto the syringe.
4. Filter the sample and collect the filtrate into a clean sample tube/container. The sample extract is ready to use in the test (stable for at least 1 hour).

Assay procedure

5. Add 0.75 ml of the filtered sample extract to a reaction vial (React). Use a Pasteur pipette provided in the kit to measure this 0.75 ml or use a single channel pipette (100 μ l – 1000 μ l). Mix by swirling and incubate for 5 minutes at room temperature.
6. Pour the content of the reaction vial onto the membrane window of a FTR device. Allow liquid to flow-through completely.
7. Add 5 drops of wash buffer (Wash). Allow liquid to flow-through completely.
8. Add 3 drops of substrate (Colour) and incubate for 2 minutes.
9. Add 2 drops of wash buffer (Wash, white cap) onto the device. Allow the liquid to flow-through completely.
10. Read the result.

R-Biopharm Nederland presents a step-by-step photo instruction of the sample preparation and assay procedure of this test at www.europroxima.com,

Scan the QR code below



9. INTERPRETATION OF RESULTS

Short explanation

If the sample is negative for OTA (<1 µg/l) then two blue coloured lines appear (C + T).

If the sample is positive for OTA (≥1 µg/l) then only one blue coloured line (control line) appears (C).

The test is invalid when no blue coloured line appears. The sample should be retested.

Further considerations

The test gives positive result (only control line visible) if a sample contains at least 1 µg/l of OTA.

Test line	Control line	Result interpretation
yes	yes	sample contains no OTA
yes/no (weak)	yes	sample can contain low level of OTA (below 1 µg/l) – please see example result on our website(OTA 0.5 µg/l)
no	yes	sample contains ≥1 µg/l of OTA

If the sample is found to be non-compliant, the results shall be verified by re-analysis of the sample using a confirmatory method.

For pictures of examples of results, please look at www.europroxima.com.

Scan the QR code below



10. LITERATURE

Commission Regulation (EC) No 1881/2006 of 19 December 2006 setting maximum levels for certain contaminants in foodstuffs, OJ L 364.

11. ORDERING INFORMATION

For ordering the Ochratoxin A wine FTD kit, please use cat. code 5127OTAW.

12. REVISION HISTORY

Not applicable