



# AlerTox Sticks Egg

Rapid immunochromatographic test for the qualitative detection of ovalbumin in food, kitchens and production facilities.

**REF** KIT3025, KIT3026





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#### 1. Intended Use

AlerTox® Sticks Egg is a rapid, immunochromatographic, lateral flow test for the qualitative detection of ovalbumin in food, kitchens and production facilities. Samples that are prepared following the instructions below can be tested using test strips (sticks) from the AlerTox Sticks Casein, Beta-Lactoglobulin, Total Milk and Egg Kits, but not with other AlerTox Sticks kits. Please read all the instructions before beginning the assay.

#### 2. Introduction

Egg protein is considered one of the major food allergens. Egg allergies often cause severe and even fatal immune reactions, including anaphylaxis, acute bronchial asthma, severe atopic dermatitis and gastroenterocolitis.

Eggs are widely used in food products such as ice cream, noodles, pasta, dressings and wine. Egg traces may remain on surfaces used for food processing. Furthermore, some vaccines may also contain trace amounts of egg proteins, posing a hazard when injected into allergic individuals.

In the US, the Food Allergen Labeling and Consumer Protection Act (FALCPA) identified egg allergies as one of the major food allergies, and the presence of egg must be labeled on packages. The European Food Safety Authority (EFSA) established a list of allergens, including egg, whose presence in foods must be indicated according to Regulation (EU) No. 1169/2011 Annex II.

# 3. Test Applications, Specificity and Sensitivity

AlerTox Sticks Egg is based on a lateral flow immunoassay and combines antibodies specific to ovalbumin. AlerTox Sticks Egg detects ovalbumin residue in a wide variety of food matrices and in environmental samples. This kit is suitable for the following applications:

- Food samples
- Rinse water testing
- Surface testing

The limit of detection (LOD) for solid and liquid samples is 1.25 ppm of ovalbumin (1.25 mg of ovalbumin per kg or L of sample). The range of detection (ROD) is 1.25 - 8,000 ppm of ovalbumin (mg/kg or mg/L). Overloading (signal decrease) may be seen at 100 - 8,000 ppm of ovalbumin. A total hook effect (i.e., false negative) is observed at 16,000 ppm and above. If a false negative due to the hook effect is suspected, repeat the test using a diluted sample.

The LOD of AlerTox Sticks Egg for surface analysis is approximately  $0.031 \, \mu g$  of ovalbumin/16 cm<sup>2</sup> on a model, dry surface (stainless steel), sampled with a wet swab. View the LOD for surface testing on the Certificate of Analysis (search by lot number at <a href="https://www.hygiena.com/documents">www.hygiena.com/documents</a>).

Section 12 contains the list of matrices currently validated for the kit using an LOD of 1.25 ppm of ovalbumin.

AlerTox Sticks Egg is a qualitative assay. To quantify the amount of antigen, use the AlerTox ELISA Egg Kit (based on ovomucoid, KIT3046) or the AlerTox ELISA Ovalbumin Kit (KIT3045).

#### NOTE:

- AlerTox Sticks Egg *cannot* detect ovalbumin in heat-treated matrices (i.e., heated over 100 °C or 212 °F for more than 30 min).
- Samples that are very viscous, dense or have a high fat content can migrate incorrectly along the
  chromatography membrane, affecting the assay results (weakening or suppressing of test and control lines).
   Contact us for more information, as these sample extractions may require larger dilutions that affect the LOD
  (www.hygiena.com/support).





#### 4. Kit Contents

Component	KIT3026	KIT3025
Ovalbumin immunochromatographic test strips in a sealed container	25 (1 container)	10 (1 container)
Extraction solution, ready to use, 60 mL	3	1
Small yellow pipettes, 1 mL	25	10
Large transparent pipettes, 3 mL	25	10
Empty tubes for extraction procedure	25	10
Swabs (for surface sampling)	25	10
Microtiter 8-well strips	4	2
Microtiter tray	1	1

# 5. Other Materials Not Supplied

- Grinder, mortar or any other manual or automatic homogenization system to crush the sample
- Vortex mixer/shaker (recommended, not required)
- Pipette or syringe to transfer 0.5 mL (only for liquid samples)
- Scissors (only for surface sampling)
- Digital scale to weigh 0.5 g (sensitive to 0.1 g)

#### 6. Precautions

- All kit components should be stored at 10 to 30 °C (50 to 86°F).
- When opening the container to use test strips, only remove the necessary number of test strips and close the container immediately.
- Do **NOT** touch the white end of the test strip.
- Use the test strip within 10 minutes after removal from the container.
- Do not use the test strip if it is broken or damaged.
- Do not use the test strips beyond the expiration date.
- Do not combine components from different kits.
- All test kit components are disposable; do not reuse them.

# 7. Sample Handling

All samples must be at 18 to 35 °C (64.4 to 95 °F) before testing.

The test is designed to detect the target antigen in:

- Solid food
- Liquid samples:
  - Beverages
  - Wash water from cutting equipment
- Surfaces





# 8. Test Procedure for Solid Food Samples

- **8.1** Before adding the sample to the provided extraction tube, mash or crush it to obtain the finest crumbs possible. Use a mortar or a grinder, if possible.
- **8.2** Add 0.5 g of sample to the extraction tube.
- 8.3 Add 5 mL of extraction solution with the transparent pipette.
- 8.4 Shake for at least 20 seconds using a vortex mixer to ensure homogenization. Alternatively, shake the tube vigorously by hand.
- **8.5** Let it rest for 2 minutes so the solids settle.
- 8.6 Using the yellow pipette, add 10 drops of the supernatant to a clean, unused well (8-well strips provided).

**Note**: For samples with high fat content, avoid the fat layer of the supernatant.

8.7 Open the container of test strips, remove the needed number of strips by holding the RED end of the strip and close the container immediately. Then, insert the white end of the strip vertically into the well containing the sample extract.

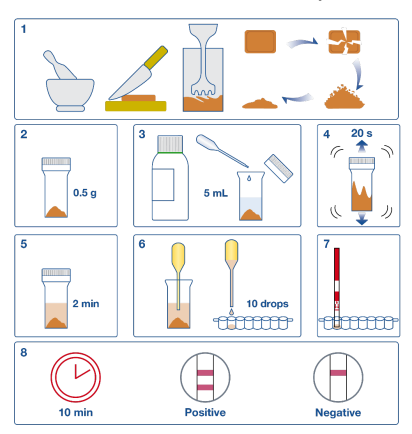
**Note**: Do NOT touch the white end of the test strip.

8.8 Wait 10 minutes to read the result.

Note: Do not read results after more than 10 minutes, as results may vary. Do not touch the test strip while waiting.

NOTE: Larger samples produce more reliable results because larger samples are more representative of the matrix than smaller samples. When extracting larger samples, maintain the 1:10 ratio of sample weight (g) to extraction solution (mL).

# **Workflow for Solid Food Samples**







# 9. Test Procedure for Liquid Samples

Liquid samples – beverages, wash water from kitchen dishes, technological surfaces or cutting machines – may be tested directly. Turbid samples should be filtered (paper or textile filter) or allowed to settle.

- 9.1 Shake the sample to ensure it is homogeneous and that you are taking a representative test portion.
- 9.2 Add 0.5 mL of the sample to the provided extraction tube using a pipette or syringe (not provided).
- 9.3 Add 4.5 mL of extraction solution with the transparent pipette.
- 9.4 Shake the sample for at least 20 seconds using a vortex mixer to ensure homogenization. Alternatively, shake it vigorously by hand.
- 9.5 If the liquid is cloudy, let it rest for 2 minutes so the solids settle.
- 9.6 Using the yellow pipette, add 10 drops of the supernatant to a clean, unused well (8-well strips provided).
  - **Note**: For samples with high fat content, avoid the fat layer of the supernatant.
- 9.7 Open the container of test strips, remove the needed number of strips by holding the RED end of the strip and close the container immediately. Then, insert the white end of the strip vertically into the well containing the sample extract.

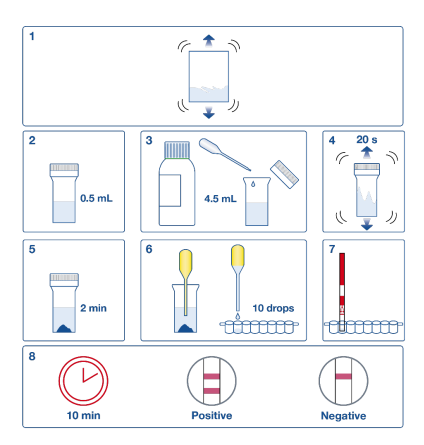
Note: Do NOT touch the white end of the test strip.

9.8 Wait 10 minutes to read the result.

Note: Do not read results after more than 10 minutes, as results may vary. Do not touch the test strip while waiting.

NOTE: Larger samples produce more reliable results because larger samples are more representative of the matrix than smaller samples. When extracting large samples, maintain the 1:10 ratio of sample volume to extraction mixture volume.

# **Workflow for Liquid Samples**







#### 10. Test Procedure for Surfaces

Collect each sample using a clean, unused swab. The swab can be used on working surfaces or equipment.

- 10.1 Add 0.5 mL of extraction solution to one of the provided extraction tubes with the transparent pipette.
- 10.2 Wet the tip of the swab with the solution. Then, firmly rub and rotate the swab on the testing surface using a zigzag pattern (at least 16 cm<sup>2</sup>/2.5 in<sup>2</sup> or a line of 40 cm/15.6 in).

**Note:** When possible, swab an approximately 4 cm x 4 cm (1.6 in x 1.6 in) square area. For irregular surfaces, ensure the swabbing technique remains consistent for each test. The area selected for analysis must be representative of the total area of interest.

10.3 Place the swab in the tube and press it against the inside walls to facilitate sample extraction into the buffer. Then, trim the swab with scissors.

**Note**: The swab should fit in the tube when the cap is closed.

10.4 Shake the sample for at least 20 seconds using a vortex mixer to ensure homogenization.

Alternatively, shake the tube vigorously by hand.

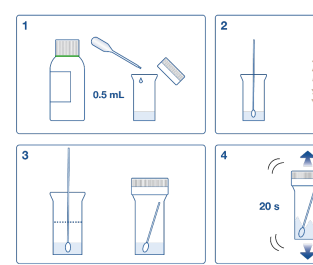
- **10.5** Open the tube and remove the swab.
- 10.6 Open the container of test strips, remove the needed number of strips by holding the RED end of the strip and close the container immediately. Then, insert the white end of the strip vertically in the tube.

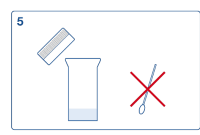
**Note**: Do NOT touch the white end of the test strip.

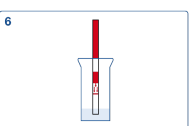
10.7 Wait 10 minutes to read the result.

Note: Do not read results after more than 10 minutes, as results may vary. Do not touch the test strip while waiting.

# **Workflow for Surfaces**

















#### 11. Interpretation of Results

The test result is POSITIVE if TWO colored lines appear: one in the control zone (C) and one in the test zone (T). The color intensity of the test line may vary, but it is not necessarily proportional to the concentration of the ovalbumin antigen in the sample.



The test result is NEGATIVE if only ONE colored line is clearly visible in the control zone (C).



If NO red line appears in the control zone (C), the test is INVALID.



If the test is invalid, check for the following and repeat the test with another strip:

- Correct specimen handling
- Correct test procedure
- Expiration date
- Correct storage conditions

For further assistance, contact Hygiena at <a href="https://www.hygiena.com/support">www.hygiena.com/support</a>.

#### **IMPORTANT NOTE!**

AlerTox Sticks is a qualitative test intended to screen samples for internal quality control. Under no circumstances can it replace laboratory analysis testing for quantification.

#### 12. Validation

AlerTox Sticks Egg has been validated for the following matrices:

Validated Matrices				
Alcoholic drinks (including wine, Irish cream)	Cereal products	Non-alcoholic drinks		
Baked products (including chocolate cookies)	Chocolate	(including soy milk, soy drinks)		
Biscuits	Chocolate cereals	Sauces		
Cereal	Dehydrated food	Snacks		

Matrices must be validated before use with AlerTox Sticks Egg. For additional information about matrix validation, contact Hygiena at <a href="https://www.hygiena.com/support">www.hygiena.com/support</a>.





#### 13. Disclaimer

Field of use: Use the Hygiena product for research and development, quality assurance and quality control under the supervision of technically qualified persons. The information generated from the Hygiena product is only to be used in conjunction with the user's regular quality assurance program. The Hygiena product should not be used as the sole basis for assessing the safety of products for release to consumers. Data obtained from the Hygiena product must not be used for human diagnostic or human treatment purposes. Before using the product, read the Limitation of Warranty and Liability (available in the Hygiena General Terms and Conditions at www.hygiena.com/terms-and-conditions).

These products are made from high-quality raw materials. No warranty of any kind is made, either expressed or implied, as to their suitability other than to measure the target antigen content when used exactly in accordance with these instructions, except regarding the quality of these materials.

Use of the kit for any other purpose is outside its intended use. For matrices that have not been previously validated, Hygiena cannot guarantee that the kit is fit for purpose and that the results obtained for these matrices are accurate. Customers may choose to use the product on unvalidated food or surface matrices; however, Hygiena strongly recommends that users perform their own fit-for-use testing to confirm suitability and performance in their specific application. Any damages, including consequential or special damage or expense arising directly or indirectly from using this product, are limited to the replacement value of the kit.

For additional information or assistance with matrix validation, contact Hygiena at www.hygiena.com/support. All Hygiena Terms and Conditions apply and can be found at: www.hygiena.com/terms-and-conditions.

#### 14. Contact Information

For more information, visit www.hygiena.com/contact. For technical support, visit www.hygiena.com/support.

# 15. Change Index

INS3014 REVC, January 2020

Included minor editorial updates. Updated the Validation section.

INS-KIT3025-3026-001-REVA, July 2025

Updated LOD and ROD information. Standardized branding, wording, some graphic workflows and document ID number.





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