Dia Sure

Allergen-E1

Allergen (Cat) Rapid Test Device

INTENDED US

The Allergen (Cat) rapid test is a rapid test for the qualitative determination of cat dander specific Immunoglobulin E (sIgE) in human serum, plasma or whole blood.

The test, in conjunction with other clinical observations, is intended to identify the patient whose allergic symptoms maybe mediated by cat dander-specific immunoglobulin E (IgE) Type I hypersensitivity.

INTRODUCTION

Allergy is a common health problem, affecting approximately 20-25% of people with immediate-type hypersensitivity reactions that manifest in the form of rhinitis, urticaria, dermatitis, gastrointestinal illness, wheezing and rarely anaphylactic shock. The term allergy is often used for type I hypersensitivity reactions (immediate type reactions), whose symptoms generally occur within 30-60 minutes after contact with the allergen.

The allergens causing type I hyper sensitivity reactions are mostly proteins derived from the natural environment e.g. plant pollen, animal hair, food, mites, and insect venoms. A characteristic of type I allergies is the involvement of allergen specific immunoglobulins(antibodies) of class E (sIgE). Hence, the detection of sIgE is an important tool of modern allergy diagnostics.

Cat dander is formed on the pet's skin and is shed through the coat. People are allergic to a certain protein that is present in the cat dander. The same protein is present in the cat's saliva also. These allergens are airborne and when inhaled, cause allergic reactions in sensitive people. Cat dander may stick easily to clothes, rugs or couches and may resist for up to 6 months in a household, even after the cat is gone.

The most frequent allergy symptoms caused by cat dander are: skin rashes, sneezing, itchy eyes, itchy skin and runny nose. In some cases, cat dander can cause asthma episodes and even chronic asthma.

PRINCIPLE

The Allergen (Cat) rapid test Device has been designed to detect cat dander slgE through visual interpretation of color development in the internal strip. The membrane was immobilized with streptavidin on the test region, the conjugate pad was pre-coated with colored anti-IgE antibody colloidal gold conjugates and the sample pad was pre-coated with biotinylated allergen. After specimens were added, the gold-conjugates move along the membrane chromatographically by capillary action and antibodies get to the test region. If sufficient cat dander slgE is present in the sample, it will react with biotinylated allergen in sample pad, the mixture then migrates through conjugate pad by capillary action and interact with colored anti-IgE antibody colloidal gold conjugates, form a complex. Then the complex moves to the membrane, and combine with streptavidin. As a result, a colored band will form at the test region of the membrane.

If no cat dander sIgE is present in the sample, biotinylated allergen pre-coated on the sample pad will bind to streptavidin immediately, so there is no colored line at the test region of the membrane.

Therefore, the colored band on the test region indicates a positive result. And appearance of a colored band at the control region serves as a procedural control. This indicates that proper volume of specimen has been added and membrane wicking has occurred.

MATERIALS

Materials Provided

Individually packed test devices

· Package insert

Disposable pipettes

Whole blood buffer

Materials Required but Not provide

· Specimen collection container

• Timer

PRECAUTIONS

- · For professional in vitro diagnostic use only.
- Do not use after the expiration date indicated on the package. Do not use the test if the foil pouch or canister is damaged. Do not reuse tests.
- This kit contains products of animal origin. Certified knowledge of the origin and/or sanitary state
 of the animals does not completely guarantee the absence of transmissible pathogenic agents. It is,
 therefore, recommended that these products be treated as potentially infectious, and handled by
 observing usual safety precautions (e.g., do not ingest or inhale).
- Avoid cross-contamination of specimens by using a new specimen collection container for each specimen obtained.
- Read the entire procedure carefully prior to testing.
- Do not eat, drink or smoke in the area where specimens and kits are handled. Handle all specimens
 as if they contain infectious agents. Observe established precautions against microbiological hazards
 throughout the procedure and follow standard procedures for the proper disposal of specimens.
 Wear protective clothing such as laboratory coats, disposable gloves and eye protection when
 specimens are assayed.
- Humidity and temperature can adversely affect results.
- Used testing materials should be discarded in accordance with local regulations.

STORAGE AND STABILITY

• The kit should be stored at 2-30°C until the expiry date printed on the sealed pouch or canister.

- The test must remain in the sealed pouch or closed canister until use.
- Do not freeze.
- · Kits should be kept out of direct sunlight.
- Care should be taken to protect the components of the kit from contamination. Do not use if there is
 evidence of microbial contamination or precipitation. Biological contamination of dispensing
 equipment, containers or reagents can lead to false results.

SPECIMEN COLLECTION AND STORAGE

- The Allergen (Cat) rapid test Device is intended for use with human whole blood, serum and plasma only.
- Only clear, non-hemolyzed specimens are recommended for use with this test. Serum or plasma should be separated as soon as possible to avoid hemolysis.
- Perform testing immediately after specimen collection. Do not leave specimens at room temperature
 for prolonged periods. Serum and plasma specimens may be stored at 2-8°C for up to 3 days. For
 long term storage, specimens should be kept below -20°C. Whole blood collected by venipuncture
 should be stored at 2-8°C if the test is to be run within 2 days of collection. Do not freeze whole
 blood specimens. Whole blood collected by finger stick should be tested immediately.
- Bring specimens to room temperature prior to testing. Frozen specimens must be completely thawed
 and mixed well prior to testing. Avoid repeated freezing and thawing of specimens.
- If specimens are to be shipped, pack them in compliance with all applicable regulations for transportation of etiological agents.

PROCEDURE

Bring tests, specimens, and/or controls to room temperature (15-30°C) before use.

Remove the test from its sealed pouch, and place it on a clean, level surface. Label the test with
patient or control identification. For best results, the assay should be performed within one hour.
 For Serum or Plasma specimens: Hold the dropper vertically and transfer 2 drops of serum or plasma

(approximately 50 μL) to the specimen well (S) of the test device, then start the timer.. For Whole Blood specimens: Hold the dropper vertically and transfer 3 drops of whole blood

For Whole Blood specimens: Hold the dropper vertically and transfer 3 drops of whole blood (approximately 75µL) to the specimen well (S) of the test device, then add 1 drop of buffer(approximately 40µL) and start the timer.

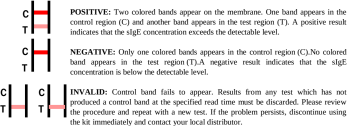
Avoid trapping air bubbles in the specimen well (S), and do not add any solution to the result area.

As the test begins to work, color will migrate across the membrane.

 Wait for the colored band(s) to appear. The result should be read at 10 minutes. Do not interpret the result after 15 minutes.
 3 drops of Whole Blood

2 drops of Serum/plasma 1 drops of buffer 10 minutes

INTERPRETATION OF RESULTS



- The intensity of color in the test region (T) may vary depending on the concentration of analytes present in the specimen. Therefore, any shade of color in the test region (T) should be considered positive. Please note that this is a qualitative test only, and cannot determine the concentration of analytes in the specimen.
- Insufficient specimen volume, incorrect operating procedure or expired tests are the most likely reasons for control band failure.

QUALITY CONTROL

 Internal procedural controls are included in the test. A colored band appearing in the control region (C) is considered an internal positive procedural control, confirming sufficient specimen volume and correct procedural technique. External controls are not supplied with this kit. It is recommended that positive and negative
controls be tested as a good laboratory practice to confirm the test procedure and to verify proper
test performance.

LIMITATIONS OF THE TEST

- The Allergen (Cat) rapid test Device is for professional in vitro diagnostic use, and should be only
 used for the qualitative detection of allergen specific Immunoglobulin E.
- The Allergen (Cat) rapid test Device will only indicate the presence of sIgE in the specimen and should not be used as the sole criteria for the diagnosis of Allergy.
- If the test result is negative and clinical symptoms persist, additional testing using other clinical methods is recommended.
- As with all diagnostic tests, a confirmed diagnosis should only be made by a physician after all clinical and laboratory findings have been evaluated.

PERFORMANCE CHARACTERISTICS

Sensitivity

The analytical sensitivity of The Allergen (Cat) rapid test is 0.7IU/mL

Accuracy

A multi-center clinical evaluation was conducted comparing results obtained using The Allergen (Cat) rapid test to another commercially available Allergen Rapid Test. The results of the study, which included 87serum specimens, demonstrated 97.7 % accuracy of The Allergen (Cat) rapid test when compared to the EIA.

Allergen (E1) Rapid Test vs. EIA						
Method		EIA		Total results		
Allergen	Results	Positive	Negative	1 otal results		
Rapid	Positive	25	1	26		
Test-E1	Negative	1	60	61		
Total Results		26	61	87		

Positive Agreement: 96.2% (81.1%-99.3%)* Negative Agreement: 98.4% (91.3%-99.7%)* Overall Agreement: 97.7% (92.0%-99.4%)*

Interference Testing

The following substances were added to cat dander sIgE free serum and whole blood samples spiked with 0.7TU/mL cat dander sIgE. None of the substances interfered with the assay at the listed concentrations.

Acetaminophen	20 mg/dL	Acetylsalicylic Acid	20 mg/dL
Ascorbic Acid	20 mg/dL	Atropine	20 mg/dL
Caffeine	20 mg/dL	Gentisic Acid	20 mg/dL
Glucose	2 g/dL	Hemoglobin	1 mg/dL

GLOSSARY OF SYMBOI

ρ	Catalog number	0	Temperature limitation
ι	Consult instructions for use	Λ	Batch code
I	In vitro diagnostic medical device	3	Use by
μ	Manufacturer	T	Contains sufficient for <n> tests</n>
σ	Do not reuse	A	Authorized representative in the European Community
Y	CE marking according to IVD Medical Devices Directive 98/79/EC		





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