

Produktinformation



Forschungsprodukte & Biochemikalien



Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



Laborgeräte & Service

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Lieferung & Zahlungsart

siehe unsere Liefer- und Versandbedingungen

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- Trockeneiszuschlag
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ASO

HumaFIA SR

Fluorescence immunoassay for the quantitative determination of Antistreptolysin O (ASO)

Package Size 25 tests/kit REF 16090/80

IVD

Intended Purpose

ASO HumaFIA SR is a fluorescence immunoassay for the semi-quantitative determination of ASO in human serum.

Anti-streptolysin O (ASO) is an antibody produced by the body after infection with group A streptococci. The ASO assay can be used to diagnose Group A Streptococcus infection and to judge the severity of the infection. Group A Streptococcus can cause tonsil angina, sinusitis, otitis media, pneumonia through respiratory tract infection, and can also cause scarlet fever, streptococcal allergic diseases such as acute glomerulonephritis, rheumatic fever, etc.

Elevated ASO is also seen in the acute stage of rheumatoid arthritis, so it can also be used for the diagnosis or differential diagnosis of rheumatoid arthritis in the acute stage, or combined with other rheumatoid arthritis indicators to evaluate the condition of rheumatoid arthritis. For professional in vitro diagnostic use only.

The kit uses a sandwich detection method to determine ASO in human serum. When an adequate volume of test sample is dispensed into the sample well of the test cartridge, the sample migrates by capillary action across the cartridge. ASO, if present in the sample, will bind to the ASO antibody conjugates, and flow downstream. The immunocomplex is then captured on the nitrocellulose membrane by the pre-coated another ASO antibody, forming an antibody-antigen-antibody complex. The more antigen in sample forms the more complex and leads to stronger intensity of fluorescence signal, which is processed by instrument for the test to show ASO concentration in sample.

Reagents and Contents

25 tests Reagent cartridge, labelled as ASO, individually RGT

sealed foil pouches

CAL 1 card Calibration card, for upload of calibration curve into

the analyzer

25 x 1 mL Sample Diluent DIL

Applicable Instrument

REF	
16090	HumaFIA

Storage/Stability

Storage/Stability		
Unopened at 4-30°C	Up to the stated expiration date. Do not use beyond the expiration date	
Opened pouch at 18-28°C	Test should be completed within 1 hour	

Reagent preparation **Specimens**

- 1. Sample type: Serum. Other sample types have not been evaluated.
- 2. The serum should be separated by centrifugation within 2 hours from the time of blood collection.
- 3. After clinical blood samples are collected, the test should be completed within 2 hours at room temperature of 18-28 °C. If the test cannot be carried out in time, it is recommended that the serum samples can be stored at 2-8°C for 8 hours. Stored at -20°C for 3 months. Avoid heating inactivated samples and hemolysis samples should be discarded.

Procedure

Follow the procedure exactly as described.

Procedural Notes

- 1. Before sample testing, read the instruction and applicable Instrument User Manual carefully. Before using refrigerated test cartridge, allow it to reach the operating temperature (18-28°C). Place the test cartridge on a level, horizontal surface.
- 2. Check the lot number of the test cartridge RGT and of the diluent DIL matches that of the Calibration card CAL
- 3. Insert the CAL into the CAL card port of the instrument for tests.
- 4. Refrigerated serum specimen must be allowed to reach room temperature before testing. Frozen serum should be vortexed and centrifugated after thawing. The supernatant liquid should be obtained and reach room temperature before testing.

 5. Pipette exactly 10 µL of serum to the vial of diluent. Reclose the cap of
- the vial and mix the sample mixture by inverting the vial 10 times. 6. Pipette exactly 100μ L of diluted sample into the sample port of the test
- cartridge. Start the timer, 10 minutes is needed for the reaction.
- 7. Insert the test cartridge into the applicable analyzer after reaction time is elapsed, select the accurate sample type and press 'STAT sample'. The

test cartridge can be detected automatically. The results will be displayed on the screen and printed when the analysis is completed.

- 8. After the assay is completed, remove the cartridge from the analyzer.
- 9. Do not re-use cartridge and diluent.
- 10. Dispose of the used cartridge, diluent, and pipette tips in accordance with any applicable regulations.

Calibration

Master curve calibration: Every HumaFIA SR reagent kit has a calibration card CAL, containing lot specific information for the calibration. After upload of calibration curve from the card, the calibration of this lot is stored at the instrument. All subsequent samples may be tested without further upload.

Calculation of Results

The HumaFIA system automatically calculates the analyte concentration of each sample. The results are given in U/mL.

Quality Control

For checking the correct functioning of the system, a suitable control material should be used, according to the legal requirements of the laboratory.

Reference Value

Reference value: <200 U/mL

It is recommended that each laboratory establish its own expected values for the population it serves.

Interpretation of Results

If the concentration in the sample exceeds the linear range, please dilute the sample with 5% BSA normal saline solution before testing, and report the result multiplied by dilution factor. The maximum dilution of the sample was 3 times.

The test results should not be used as the sole basis for diagnosis and should be combined with other clinical and laboratory data for clinical diagnosis. If the test results do not match the clinical assessment, further tests should be performed.

Limitations

Interference may be encountered with certain samples containing antibodies directed against reagent components. For this reason, assay results should be interpreted taking into consideration the patient's clinical history, and the results of any other tests performed.

When the sample contains high concentration of triglyceride, cholesterol, bilirubin or hemolysis, the test result will be affected. Do not use samples containing any of the following interfering substances for testing: Severe lipemia: triglyceride concentration exceeds 15 mg/mL; High cholesterol: cholesterol concentration of more than 400 mg/dL; Jaundice: the concentration of bilirubin exceeds 40 mg/L; Severe hemolysis: the concentration of hemoglobin exceeds 6 mg/mL; As with any assay employing mouse antibodies, the possibility exists for interference by human anti-mouse antibodies (HAMA) in the sample. The test has been formulated to minimize this interference; however, specimens from patients who have been routinely exposed to animals or to animal serum products may contain heterophile antibodies which may

cause erroneous results. Analytical Performance Characteristics

Analytical Performance Characteristics		
Appearance	The inner packaging is tightly sealed without leakage. The label content is complete, and the text is clear and easy to recognize. The surface of the test card should be smooth, without burrs, and uniform in color.	
Film strip width	The width of the test strip in the display window should be ≥3.7mm.	
Liquid migration speed	≥10mm / min.	
Samples that exceed the displayed range are marked as >higher or <lower measurement="" range<="" td="" than="" the=""></lower>		
Accuracy	Tested with accuracy reference material, the total agreement rate is >90%	
Within-run Precision	≤15%	
Between-run Precision:	≤15%	
Limit of detection	<50 U/mL	
Linearity range	50~600 U/mL	

As with all diagnostic tests, the results should be interpreted with due consideration of other laboratory findings and of the clinical status of the

Safety Notes

All patient specimens and controls should be handled as potentially infectious. All materials of animal origin avoid many risks associated with the use of human serum (e.g. Hepatitis B and C, HIV). Nevertheless, all material of human or animal origin should still be treated as potentially infectious material.

For users in the European Union only: Please report any serious incident that has occurred in relation to the device to the manufacturer and the competent authority of the Member State in which the user and/or the patient is established.

P234 Keep only in original container.
P260 Do not breathe dust/fume/gas/mist/vapors/spray.
P262 Do not get in eyes, on skin, or on clothing.
P281 Use personal protective equipment as required.
P303+P361+P353 IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water/shower.
P305+P351+P338 IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

P337+P313 If eye irritation persists: Get medical advice/attention. P401 Store in accordance with local/regional/national/international regulations.

P501 Dispose of contents/container in accordance with local/regional/national/international regulations.



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