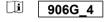
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Electronic Instruction For Use: version



ORG 906G Anti-Chlamydia trachomatis IgG

INTENDED PURPOSE

Anti-Chlamydia trachomatis IgG for Alegria[®] is an ELISA-based test system for the quantitative measurement of IgG class antibodies against *Chlamydia trachomatis* in human serum or plasma samples. This product is intended for professional in vitro diagnostic use only.

After primary infection, IgM, IgA, and IgG antibodies can be detected successively in serum samples. IgG antibodies are generally considered as markers for any contact with the pathogen irrespective of disease stage. Several years after the infection, they may disappear again. IgM antibodies are characteristic for acute *Chlamydia* infection, and IgA antibodies indicate ongoing progression of a persistent infection.

SYMBOLS USED

IVD	In vitro diagnostic medical device		
***	Manufacturer	ALEGRIA TEST STRIPS	Alegria [®] Test Strips
REF	Catalogue number	WASH SYSTEM FLUID	Wash Buffer System Fluid
24	Sufficient for determinations	RTU	Ready to use
LOT	Batch code		
\square	Use by		
2°C	Temperature limitation		
ĺ	Consult instructions for use		
*	Keep away from sunlight		
8	Do not reuse		
M	Date of manufacture		

CE marked according to 98/79/EC

PRINCIPLE OF THE TEST

C. trachomatis antigens MOMP, TARP, and CPAF are bound to reaction wells.

The Alegria[®] assay features barcoded 8-well-microstrips, called Alegria[®] Test Strips. Each strip is designed for a single determination of one patient sample. The Alegria[®] Test Strip holds a complete set of reagents. Included are enzyme conjugate, enzyme substrate, sample buffer and a test specific control. Furthermore each strip has two antigen-coated wells which serve as reaction wells for one control and one patient sample.

The determination is based on an indirect enzyme linked immune reaction with the following steps: Antibodies present in positive samples bind to the antigen coated on the surface of the two reaction wells forming an antibody antigen complex. After incubation, a first washing step removes unbound and unspecific bound molecules. Subsequently added enzyme conjugate binds to the immobilized antibody-antigen complex. After incubation, a second washing step removes unbound enzyme conjugate. Addition of enzyme substrate solution results in hydrolisation and color development during incubation. The intensity of the blue color correlates with the concentration of the antibody-antigen-complex and can be measured photometrically at 650 nm.

The Alegria[®] Test Strip is based on the proprietary SMC[®]-Technology (Sensotronic Memorized Calibration): information about the assay, analysis and evaluation, and the lot-specific expiry date is contained on the barcode printed on each Alegria[®] Test Strip.

The Alegria[®] Test Strip can be used with the diagnostic instrument Alegria[®] - a fully automated Random Access Analyser. By means of SMC[®]-Technology data encoded on the barcode are transferred from the Alegria[®] Test Strip to the instrument and the assay is automatically processed and evaluated. The instrument reads the date of expiry and rejects further processing if the Alegria[®] Test Strip is out of date.

WARNINGS AND PRECAUTIONS

- · All reagents of this kit are intended for professional in vitro diagnostic use only.
- Components containing human serum were tested and found negative for HBsAg, HCV, HIV1 and HIV2 by FDA approved methods. No test can guarantee the absence of HBsAg, HCV, HIV1 or HIV2, and so all human serum based reagents in this kit must be handled as though capable of transmitting infection.
- Bovine serum albumin (BSA) used in components has been tested for BSE and found negative.
- Avoid contact with the substrate TMB (3,3',5,5'-Tetramethyl-benzidine).
- · System fluid contains acid, classifiaction is non-hazardous. Avoid contact with skin.
- Control, sample buffer and wash buffer contain sodium azide 0.09% as preservative. This concentration is classified as non-hazardous
- Enzyme conjugate, control and sample buffer contain ProClin 300 0.05% as preservative. This concentration is classified as non-hazardous.

During handling of all reagents, controls and serum samples observe the existing regulations for laboratory safety regulations and good laboratory practice:

First aid measures: In case of skin contact, immediately wash thoroughly with water and soap. Remove
contaminated clothing and shoes and wash before reuse. If system fluid comes into contact with skin,
wash thoroughly with water. After contact with the eyes carefully rinse the opened eye with running
water for at least 10 minutes. Get medical attention if necessary.

• Personal precautions, protective equipment and emergency procedures:

Observe laboratory safety regulations. Avoid contact with skin and eyes. Do not swallow. Do not pipette by mouth. Do not eat, drink, smoke or apply makeup in areas where specimens or kit reagents are handled. When spilled, absorb with an inert material and put the spilled material in an appropriate waste disposal.

- Exposure controls / personal protection: Wear protective gloves of nitril rubber or natural latex.
 Wear protective glasses. Used according to intended use no dangerous reactions known.
- Conditions to avoid: Since substrate solution is light-sensitive. Store Alegria® strips in the dark.
- · For disposal of laboratory waste the national or regional legislation has to be observed.

Observe the guidelines for performing quality control in medical laboratories by assaying controls and/or pooled sera.

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CONTENTS OF THE K	т
🗑 24 ORG 906G	Sufficient for 24 determinations
ALEGRIA TEST STRIPS 24	Alegria® Test Strips are modules of 8 wells each composed of: Wells 1 + 2: empty and not coated (wells for the sample dilution) Wells 3 + 4: coated with antigen (reaction wells)
	Well 5: Control; yellow; containing C. trachomatis antibodies, PBS, BSA, detergent, preservative sodium azide 0.09% and ProClin 300 0.05%.
1^2 4^3	Well 6: Enzyme Conjugate; light red; containing anti-human IgG antibodies, HRP labelled; PBS, BSA, detergent, preservative ProClin 300 0.05%.
	Well 7: Sample Buffer: yellow; containing PBS, BSA, detergent, preservative sodium azide 0.09% and ProClin 300 0.05%.
	Well 8: TMB Substrate: clear; containing 3,3', 5,5'- Tetramethylbenzidin.
	Code on barcode: Chlamy tracho IgG on printout: Ctrac-G
WASH 1x 20	nl Wash Buffer, containing Tris, detergent, preservative sodium azide 0.09%; 50 x conc.
SYSTEM FLUID 1x 2.5	ml System Fluid, contains acid; 1000 x concentrate

Certificate of Analysis

STORAGE AND STABILITY

- Store test kit at 2-8°C in the dark.
- Do not expose reagents to heat, sun, or strong light during storage and usage.
- Store Alegria[®] Test Strips sealed and dessicated in the clip bag provided.
- Shelf life of the unopended test kit is 15 months from day of production. Unopened reagents are stable until expiration of the kit. See labels for individual batch.
- Diluted Wash Buffer and System Fluid are stable for at least 30 days when stored at 2-8°C. Once transferred to the reagent container we recommend consumption on the same day.

MATERIALS REQUIRED

- Vortex mixer
- Pipettes for 10 µl
- Measuring cylinder for 1000 ml and 2500 ml
- · Distilled or deionized water

SPECIMEN COLLECTION, STORAGE AND HANDLING

- Collect whole blood specimens using acceptable medical techniques to avoid hemolysis.
- · Allow blood to clot and separate the serum or plasma by centrifugation.
- · Test serum should be clear and non-hemolyzed. Contamination by hemolysis or lipemia should be avoided, but does not interfere with this assav.
- Specimens may be refrigerated at 2-8°C for up to five days or stored at -20°C up to six months.
- · Avoid repetitive freezing and thawing of serum or plasma samples. This may result in variable loss of antibody activity.
- · Testing of heat-inactivated sera is not recommended.

PROCEDURAL NOTES

- · Do not use kit components beyond their expiration dates.
- All materials must be at room temperature (20-28°C) prior to use.
- To avoid carryover or contamination, change the pipette tip between samples.

PREPARATION OF REAGENTS

WASH

Dilute the content of the Wash Buffer concentrate (50x) with distilled or deionized water to a final volume of 1000 ml prior to use. Transfer the diluted Wash Buffer into the instrument reagent container. If only one Alegria run is to be performed on one day we recommend transferring only 500 ml diluted Wash Buffer.

SYSTEM FLUID

Dilute the content of the System Fluid concentrate (1000x) with distilled or deionized water to a final volume of 2500 ml prior to use. Transfer the diluted System Fluid into the instrument reagent container. ALEGRIA TEST STRIPS

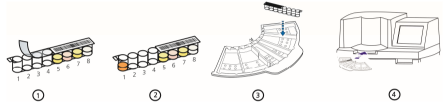
Take the required number of Alegria® Test Strips out of the clip bag and let them reach room temperature (20-28°C). Do not remove foil covering the empty wells until you are ready to start the assay.

TEST PROCEDURE

Alegria[®] Test Strips with SMC[®] technology are used with the diagnostic instrument Alegria[®]. Detailed information about operating the instrument can be taken from the Instrument User Manual. (1) Remove the foil from the empty wells 1 to 4 of the Alegria[®] Test Strip.

Do not remove foil with printed barcode, covering wells 5 to 8.

- (2) Pipette 10 µl undiluted sample at the bottom of well 1.
- (3) Insert the strip into the SysTray.
- (4) Place loaded SysTrays into the correct position in the Alegria® instrument and start run. All further steps will be done automatically. The test run is completed when the instrument starts printing the results.



CALIBRATION

Since no international reference preparations for Chlamydia trachomatis IgG antibodies are available, the assay is calibrated in relative arbitrary units.

CALCULATION OF RESULTS

By means of SMC[®] Technology (Sensotronic Memorized Calibration), all test data are transferred to the system through individual barcodes on the Alegria[®] Test Strip. Calculation and interpretation of results will be performed automatically.

PERFORMANCE CHARACTERISTICS

Measuring range

The calculation range of this Alegria® assay is 0 - 200 U/ml

Expected values

The cut-off of this Alegria[®] assay is: 25 U/ml

Interpretation of results

Negative:	< 20 U/ml
Borderline:	20 - 25 U/ml
Positive:	> 25 U/ml

LIMITATIONS OF THE PROCEDURE

This assay is a diagnostic aid. A definite clinical diagnosis should not be based on the results of a single test, but should be made by the physician after all clinical and laboratory findings have been evaluated concerning the entire clinical picture of the patient. Also every decision for therapy should be taken individually.

The above reference ranges should be regarded as guidelines only. It is recommended that each laboratory establishes its own normal and pathological ranges for antibodies in patient samples.

A negative result does not rule out an infection, since the serum can be sampled too early for the antibodies to be detectable. A positive result does not rule out the presence of another infectious pathogen as the cause of disease.

Linearity

Three patient samples containing high levels of specific antibody were serially diluted in sample buffer to demonstrate the dynamic range of the assay. Activity for each dilution was calculated by means of SMC[®] Technology.

Sample	Dilution	Observed	Expected	O/E
		U/ml	U/ml	[%]
1	1:100	<mark>158.7</mark>	158.7	100
	1:200	73.9	80.0	93
	1:400	40.9	40.0	103
	1:800	21.0	20.0	105
2	1:100	113.6	113.6	100
	1:200	53.2	59.0	90
	1:400	31.1	30.0	105
	1:800	16.3	15.0	110
3	1:100	69.9	69.9	100
	1:200	37.7	40.0	94
	1:400	21.7	20.0	108
	1:800	<mark>11.1</mark>	10.0	111

Detection limit

The lowest amount of detectable antibody is: 3.9 U/ml

Reproducibility

Intra-assay precision: Coefficient of variation (CV) was calculated for each of three samples from the results of 24 determinations in a single run. Results for precision-within-assay are shown in the table below. Inter-assay precision: Coefficient of variation (CV) was calculated for each of three samples from the results of 6

determinations in 5 different runs. Results for run-to-run precision are shown in the table below.

Intra-Assay		Inter-Assay			
Sample	Mean		Sample	Mean	
	[U/ml]	% CV		[U/ml]	% CV
1	8.3	2.4	1	9.5	5.6
2	27.8	5.4	2	29.4	3.7
3	87.3	4.4	3	197.9	4.1

Interfering substances

No interference has been observed with haemolytic (up to 1000 mg/dl) or lipemic (up to 3 g/dl triglycerides) sera or plasma, or bilirubin (up to 40 mg/dl) containing sera or plasma. Nor have any interfering effects been observed with the use of anticoagulants (Citrate, EDTA, Heparine). However for practical reasons it is recommended that grossly hemolyzed or lipemic samples should be avoided.

No interference has been observed in bacterial or viral infections with *C. pneumoniae*, *Borrelia* sp., Yersinia sp., Parovirus B19, HSV-2 or acute EBV infection.

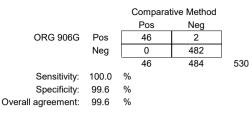
Nor have any interfering effects been observed in rheumatic diseases associated with elevated titers of autoantibodies such as rheumatoid factors or antinuclear antibodies.

In samples from patients with acute EBV infection a lower rate of seroprevalence was found possibly due to a lower average age in the study population. In samples from patients with HSV-2 infection a higher rate of seroprevalence was found possibly due to co-infection with HSV-2 and C. trachomatis. Cross-reactivity to EBV and HSV-2 are therefore not likely.

Seroprevalence

Analysis of 100 healthy blood donors from Germany showed 7 positive results equivalent to 7% seroprevalence.

Study results



*9/11 samples were confirmed positive in micro immunofluorescence assay. *5/5 samples were confirmed negative in micro immunofluorescence assay.

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Notice to the user (European Union):

Any serious incident that has occurred in relation to the device shall be reported to the manufacturer and the competent authority of the EU Member State in which the user and/or the patient is established.

Change Control

Former version: ORG 906G_IFU_EN_QM113236_2014-06-26_3 Reason for revision: Introduction electronic IFU on homepage

