


ASO Kit CliniQuant

Latex Turbidimetry

 For *in vitro* diagnostic use
Read this pack insert thoroughly before use

REF ASOLIT-01	R1 ASO Reagent	R2 ASO Reagent	R3 ASO Calibrator
Pack size	1 x 40 ml	1 x 10 ml	1 x 1 ml

INTENDED USE

This reagent is intended for quantitative determination of Anti streptolysin O concentration in serum by latex turbidimetry.

CLINICAL SIGNIFICANCE

Streptolysin O (SLO) is a lethal, exocellular protein released by Group A Streptococcal bacteria. The release of SLO stimulates the production of anti-streptolysin O (ASO) antibodies to neutralise its haemolytic effect.

The ASO test is used to determine recent streptococcal infection and post streptococcal complications including rheumatic fever and glomerulonephritis. The presence and level of ASO antibodies in human serum directly reflects the extent and degree of infection. Elevated levels of ASO may also be present in other conditions including scarlet fever, acute rheumatoid arthritis, tonsillitis and various other streptococcal infections as well as in health carriers.

PRINCIPLE OF THE METHOD

Latex particles coated with streptolysin O (SLO) are agglutination when mixed with samples containing ASO. The agglutination causes an absorbance change, dependent upon the ASO contents of the patient sample that can be quantified by comparison from a calibrator of known ASO concentration.

KIT COMPONENTS

R1 - ASO Reagent
R2 - ASO Reagent
R3 - ASO Calibrator

MATERIALS REQUIRED BUT NOT PROVIDED

Laboratory instrumentation, Spectrophotometer UV/VIS with thermostatic cuvette holder or clinical chemistry analyzer: semi automated, calibrated micropipettes, glass or high quality polystyrene cuvettes, test tube/rack, heating bath, controls, saline.

REAGENT PREPARATION , STORAGE & STABILITY

Reagent is ready to use. Mix reagent R2 well before processing.

Mix reagent 1 & reagent 2 in ratio of 4:1. Keep away from direct light sources.

Stability : up to expiration date on labels at 2-8°C. Do not freeze the reagent.

WARNINGS AND PRECAUTIONS

For *in vitro* use only.

This pack insert must be completely understood prior to operation. Do not modify the test procedure or substitute reagents from other manufacturers or other lots unless the reagent is stipulated as interchangeable. It is recommended to handle carefully by entitled and professionally educated person.

Do not pipette by mouth. Use disposable gloves while performing the assay. Wash hands thoroughly when finished.

Perform the test according to the general "Good Laboratory Practice" (GLP) guidelines.

Do not use reagents beyond the expiry date.

In case of skin contact with any of the reagents, wash thoroughly with running water.

SPECIMEN

Fresh serum : Stable for 7 days at 2 - 8°C. Do not use hemolyzed or lipemic sample.

Programme Parameter for MERILYZER CliniQuant

Reading Mode	Fixed Time
Standard Conc.	lot specific
Filter- 1 (nm)	546 nm
Temperature	37 ° C
Volume (µl)	500
Delay Time (Sec)	10
Read Time (Sec)	120
Reaction Direction	Increase
Reference Low	0
Reference High	200
Linearity Limit	800

TEST PROCEDURE

Dispense	Blank	Standard	Sample
Working reagent	1ml	1ml	1ml
Distilled water	20 µl	-	-
Standard	-	20 µl	-
Sample	-	-	20 µl

Mix, incubate 10 seconds at 37°C, then record absorbance as A1. After exactly 120 seconds, record again absorbance as A2.



RESULT CALCULATION

Serum :

ASO IU/ml = $A2-A1(\text{Sample})/A2-A1(\text{Standard}) \times \text{Concentration of Standard}$

EXPECTED VALUES

Up to 200 IU/ml.

It is recommended that each laboratory verifies this range or derives reference interval for the population it serves.

QUALITY CONTROL AND CALIBRATION

It is recommended to perform internal quality control with assayed normal and assayed abnormal to confirm the validity of the test and assure the accuracy of patient result.

Using the recommended Calibrator or the Standard included, calibrate the assay:

- When using a new reagent or lot
- When QC values are out of range

PERFORMANCE CHARACTERISTICS

1. Linearity

The linearity up to 800 IU/ml

2. Sensitivity/ Limit of detection (LOD)

The limit of detection is 2.5 IU/ml.

The limit of quantification is 22.8 IU/ml.

3. Interferences

No interference has been observed for the following

Bilirubin up to 20 mg/dl.

Hemoglobin up to 500 mg/dl.

Triglycerides up to 450 mg/dl.

4. Precision

Intra-assay precision

	Mean	SD	CV
n = 20	IU/ml	IU/ml	%
Control L1	124.40	2.33	1.87
Control L3	360.41	3.15	0.87

Inter-assay precision

	Mean	SD	CV
n = 20	IU/ml	IU/ml	%
Control L1	125.26	1.38	1.10
Control L3	354.68	6.58	1.85

5. Methods Comparison

Comparison was done between ASO Reagent (y) & reference ASO Kit (x) using 20 samples gave following results:

$$y = 1.0358x + 1.1301 \quad r^2 = 0.9982$$

LIMITATIONS

Samples with values above 800 IU/ml should be diluted with 0.9% saline, re-run and multiply results by dilution factor.

WASTE DISPOSAL

This product is made to be used in professional laboratories. Please consult local regulations for correct waste disposal.

REFERENCES

- Alouf Jodeph E. Pharma Ther 1980;11:661-717.
- M Fasani et al eur J Lab Med 1994;Vol 2 no 1-67.
- Todd E W J Exp Med 1932;55-267-280.
- Data on file: Meril Diagnostics.

IFU/ASOLIT01/02

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Symbols used on Meril Diagnostics labels:

 REF	Catalogue No.		Attention See Instruction for Use
 LOT	Batch No.		In vitro Diagnostics
	Expiry Date		Consult Instruction for Use
	Manufacturer		Storage Temperature
	Keep Dry		Keep Away from Sunlight
	Manufacturing Date		Do not use if package is damaged