

C3

For use on Diatron Pictus® series analyzers Immunoturbidimetry Method: 1419-0762, 1419-0760 Product code:

4x21.6 ml (R1) + 4x10.8 ml (R2), 4x9.6 ml (R1) + 4x4.8 ml (R2) Package:

Store at: 2°-8°C For in vitro use only

INTENDED USE

Ready to use reagents for the quantitative determination of Complement component 3 (C3) in human serum specifically for use with Diatron Pictus® series analyzers. For in vitro diagnostic use only.

CLINICAL SIGNIFICANCE

Increased C3 levels in serum are observed in several inflammatory conditions, bile duct obstruction, amyloidosis (especially during the recovery period), acute heart attack, acute rheumatoid fever, ulcerative colitis, cancer. Reduced levels appear in autoimmune diseases, glomerulonephritis, malnutrition, hepatitis, rheumatoid arthritis, anemia, Sjogren's syndrome, diseases of the autoimmune system, Gram(-) bacteremia, cirrhosis.

METHOD PRINCIPLE

The immunoturbidimetric method is applied. When sample is mixed with the appropriate buffer (R1) and antiserum solution (R2), C3 reacts selectively with C3 antibodies, leading to formation of insoluble aggregates. The absorbance of the test solution at 340/700 nm is proportional to the concentration of C3 in the sample.

METHOD LIMITATIONS

Refer to the book "Effects of Preanalytical Variables on Clinical Laboratory Tests" for possible interference of other pharmaceutical agents in this particular test. Interference of other agents is described in the "Clinical Guide to Laboratory Tests"

The reagent is designed especially for use with the Diatron Pictus® series of chemistry analyzers. For chemistry protocols and further information please contact the customer support unit at Medicon or Diatron.

REAGENT COMPOSITION

Reagent 1 (R1)

Polyethylene glycol in Tris buffer

Non reactive ingredients, preservatives.

Reagent 2 (R2)

Goat anti-human C3 protein antibodies Non reactive ingredients, preservatives.

WARNINGS - PRECAUTIONS

- This reagent is designed for in vitro diagnostic use. In vitro diagnostic reagents can be hazardous. They should be handled according to good laboratory techniques. Avoid inhalation and contact with eyes and skin.
- · Samples should be considered as potentially infectious. Handle with special caution.
- · Antisera are raised in clinically healthy animals in monitored facilities under constant surveillance.
- This reagent contains sodium azide (NaN₃) ≤ 0.1%. Avoid swallowing and contact of the reagent with skin and mucous membranes.
- · Dispose all waste according to national laws.
- . MSDS is available by Diatron or MEDICON HELLAS (manufacturer) upon request.

REAGENT PREPARATION

Reagents R1 and R2 are liquid, ready to use when placed in the corresponding positions of the analyzer. The vials bear barcodes for automatic recognition by Pictus® series analyzers.

REAGENT DETERIORATION

The reagents should not be used:

- · When they do not exhibit the specified linearity or control values lie outside the acceptable range after recalibration.
- · After prolonged exposure to sunlight or high temperature.
- . When the solution is turbid.

Unopened, reagents are stable at 2°-8°C up to the expiry date stated on the label. Once opened, they remain stable for 56 days when stored in the cooled reagent tray of the Pictus® series analyzers.

Specimens of fresh serum or plasma with EDTA must be used. No fasting or special preparation of the patient is necessary prior to sampling. Separate the sample from cellular components and measure as soon as possible. Samples may be kept at -70°C.

Diatron provides Protein Standard Set (1578-1190). Calibrate the assay every 28 days on Pictus® series analyzers. Calibration should also be repeated after major maintenance is performed on the analyzer, after a critical part is replaced, or when a significant shift in control values occurs.

QUALITY CONTROL

Diatron provides the Immunology Control Levels 1,2,3 (1578-1195-04, 1578-1196-04, 1578-1197-04 respectively). Control recovery should lie within the acceptable range. Results outside the acceptable range even after recalibration could be due to reagent deterioration, unsuitable storage conditions or control deterioration, instrument malfunction, or error during test procedure.

MATERIALS REQUIRED BUT NOT SUPPLIED WITH THE KIT

- Calibrator
- · Quality control materials
- Diatron Pictus® P400/P700/P500
- · Common laboratory equipment

REFERENCE INTERVALS

60 - 170 mg/dl Serum:

Reference values are based on current bibliography. Each laboratory should determine its own expected values as dictated by good laboratory practices.

WASTE DISPOSAL

This product contains sodium azide (NaN₃), which forms sensitive explosive compounds with copper or lead. Flush waste pipes with water after the disposal of undiluted reagent in order to avoid azide build up in the drain pipes.

SPECIFIC PERFORMANCE CHARACTERISTICS

The following values are representative of the reagent performance on Diatron Pictus® series analyzers. The reagent performance has been evaluated on other types of analyzers, covering all requirements of the 98/79 IVD Directive. A list of analyzers with the corresponding performance characteristics is available in the special leaflet accompanying the insert. The results taken in your laboratory may differ from these values

	Pictus® P400	Pictus® P700/P500
Linearity	Up to 400 mg/dL	Up to 400 mg/dL
Hook effect	> 2500 mg/dL	> 2500 mg/dL
Lowest detection limit	1.0 mg/dL	0.6 ma/dL

The lowest detection limit (LDL) is defined as the lowest concentration of analyte that is distinguishable from zero. A sample free of analyte is assayed 20 times within the assay and the LDL is calculated as the absolute mean plus three standard deviations.

Precision: Precision is estimated on two concentration levels of analyte according to CLSI protocol EP 5-A (20 consecutive days, 2 runs per day, 2 repeats per run).

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Level	Within Run		Level	Within Run	
(mg/dL)	CV%	Total CV%	(mg/dL)	CV%	Total CV%
78.9	2.11	2.74	66.7	2.68	3.69
159.7	2.24	2.74	130	2.28	3.48
220.4	2.80	3.85	194	2.22	3.79
Criterion: recovery within +20% from target value					

Dictue® D700/D500

Interference	Criterion: recovery within ±20% from target value			
	Pictus® P400	Pictus® P700/P500		
Lipemia	Insignificant up to Intralipid® 500 mg/dl	Insignificant up to 400 mg/dl Intralipid®		
Haemoglobin	Insignificant up to 500 mg/dL	Insignificant up to 500 mg/dL		
Bilirubin	Insignificant up to 20 mg/dL	Insignificant up to 20 mg/dL		
Ascorbate	Insignificant up to 3 mg/dL	Insignificant up to 3 mg/dL		

Correlation: A comparison was performed between this reagent on a Pictus® series analyzer, and a BECKMAN COULTER AU-series system. The results were as follows:

Pictus® P400

Y = 0.925X + 18.75	R=0.955	N=40	Sample range = 55 - 237 mg/dL
Pictus® P700/P500			
Y = 0.935X + 19.67	R=0.965	N=26	Sample range = 82.9 – 268 mg/dL

BIBLIOGRAPHY

- 1. Tietz, NW, ed. Clinical Guide to Laboratory Tests. 3rd. ed. Philadelphia: W.B. Saunders Company Ltd., 1995
- 2. Burtis, CA and Ashwood, ER, ed. Tietz Textbook of Clinical Chemistry. 2nd. ed. Philadelphia: W.B. Saunders Company Ltd., 1994.
- 3. Jacobs, DJ, Demott, WR, Grady, HJ, Horvat, RJ, Huestis, DW and Kasten, BL, JR, eds. Laboratory Test Handbook. 4th. ed. Ohio, Hudson: Lexi-Comp Inc., 1996.
- 4. The American Association for Clinical Chemistry, Inc. Effects of Preanalytical Variables on Clinical Laboratory Tests. 2nd. ed. Ohio, Hudson: Lexi-Comp Inc., 1997.

SYMBOLS









PICTUS®: Registered Trademark of Diatron Medical Instruments Limited, Táblás utca 39, H-1097 Budapest, Hungary, used here after contractual agreement.