



IgE Control

For precision control in the determination of immunoglobulin E (IgE) by immunoturbidimetric methods

USE

IgE Control Turbitest AA is designed for precision control of **IgE Turbitest AA** kit from Wiener lab.

Refer to the table of assigned values, since they are lot-specific.

PROVIDED REAGENTS

Control level 1: lyophilized human serum containing IgE values within the range of normal values in buffered saline solution with appropriate preservatives.

Control level 2: lyophilized human serum containing IgE values within the range of pathological values in buffered saline solution with appropriate preservatives.

NON-PROVIDED REAGENTS

Distilled water.

INSTRUCTIONS FOR USE

Reconstitute each vial with the volume of distilled water stated on the vial label.

Let the vial stand for 20 minutes at room temperature and then invert gently several times until complete dissolution.

WARNINGS

Controls are for "in vitro" diagnostic use.

Controls have been prepared from non-reactive material for HBsAg, HCV and HIV. Controls and all samples should be handled as if they were infectious material. Use the reagents keeping the usual work precautions in the clinical laboratory. All reagents and samples should be discarded according to local regulations.

STABILITY AND STORAGE INSTRUCTIONS

The controls are stable at 2-10°C until the expiration date stated on the box. Once reconstituted they are stable for 2 weeks at 2-10°C. Controls can be aliquoted and stored frozen (-20°C) for 5 months. Freeze and thaw only once.

PROCEDURE


Controls should be used in the same way as an unknown sample, according to the instructions supplied with the **IgE Turbitest AA** kit from Wiener lab.

WIENER LAB PROVIDES

2 x 3 ml (1 x 3 ml level 1 + 1 x 3 ml level 2) (Cat. N° 1950300)

SYMBOLS

The following symbols are used in the packaging for Wiener lab. diagnostic reagents kits.

 This product fulfills the requirements of the European Directive 98/79 EC for "in vitro" diagnostic medical devices

 Authorized representative in the European Community

 "In vitro" diagnostic medical device

 Contains sufficient for <n> tests

 Use by

 Temperature limitation (store at)

 Do not freeze

 Biological risks

 Volume after reconstitution


 Contents

 Batch code


 Manufactured by:

 Harmful

 Corrosive / Caustic

 Irritant

 Consult instructions for use


 Calibrator

 Control

 Positive Control

 Negative Control

 Catalog number

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PM-1102-137



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