# The Creative Approach to Bioscience

# SP-NORMOTROL

Control serum normal/borderline, for Control of Accuracy and Precision in quantitative In vitro Diagnostics

REF: 326 001 1 x 5 ml **REF:** 326 002 (5 vials) 5 x 5 ml REF: 326 003 (6 vials) 6 x 5 ml

## Description

SP-NORMOTROL is a lyophilized human-based control serum.

#### Storage and stability

Unopened bottles must be stored at 2 - 8°C and are stable up to the expiration date printed on the labels

After reconstitution the serum can be used, stored at 2 - 8°C and protected from light , in between 5 days. Bilirubin is stable 1 day ,refrigerated .

The frozen serum is stable for at least 30 days at < -20°C.

### Warnings and Precautions

Each individual blood donation used for this serum was found negative when tested with FDA-approved methods on HBsAg, anti-HIV 1+2 and anti-HCV. Nevertheless the serum should be treated for safety reasons always as a potentially infectious material

#### Preparation

Reconstitute carefully by adding exactly 5 ml of distilled water. Allow the vial to stand for 30 minutes ,swirling occasionally. Avoid foaming i.e. do not shake! For alkaline phosphatase the control serum should stand for 2

hours at +25 °C before use

### **Procedure**

Refer to the package inserts of the reagent kits

### **Assay Values and Ranges**

Assay values for analytes for which approved reference methods are available were determined according to Guidelines of the German Federal Medical Council [Bundesaerztekammer] from 1987 (reference method values) [3]. Ranges of acceptance were calculated as assigned value  $\pm$  three

times the maximum tolerable deviation of a single value according to the Guidelines of the German Federal Medical Council [4].

#### **Expected Values**

- 1. The expected value of specific assays are provided on the assay value sheet accompanying each kit, and are lot specific.
- 2. The expected values are obtained using replicate assay of each manufactured lot of SP-Normotrol.
- 3. The individual laboratory values should fall within the expected values
- 4. It must however be noted that each laboratory should establish its own normal values and reference range according to GLP.

#### SYMBOLS IN PRODUCT LABELLING

Authorised Representative 

Use by/Expiration Date Batch Code/Lot number REF Catalogue Number Consult instructions for use X (Xi) - Irritant

Temperature Limitation

For in-vitro diagnostic use / CAUTION. Consult instructions for use

Manufactured by

#### Literature

- 1. Röhle G, Siekmann L. Quality assurance of quantitative determination. In: Thomas L, editor. Clinical laboratory diagnostics. 1st ed. Frankfurt:TH-Books Verlagsgesellschaft; 1998. p. 1393-
- 2. Biosafety in Microbiological and Biomedical Laboratories. U.S. Department of Health and Human Services, Washington 1993 (HHS Publication No. [CDC] 93-8395).
- 3. Richtlinien der Bundesärztekammer zur Qualitätssicherung in medizinischen Laboratorien. Deutsches Ärzteblatt 1988;85: B519–B532.
- 4. Richtlinie der Bundesärztekammer zur Qualitätssicherung quantitativer laboratoriumsmedizinischer Untersuchungen. Deutsches Ärzteblatt 2002;98:A 2747-59.

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