

# **GLUCOSE**

# Enzymatic colorimetric determination of glucose in serum and plasma

### **TEST SUMMARY**

Glucose become oxidize means by Glucose Oxidise creating distilled water and Gluconic Acid. The distilled water reacts with aminophenation and phenol creating a red compound. Intensity of colour measures at 510 nm and is proportional to the quantity of Glucose present in the sample.

#### **SAMPLES**

Serum, plasma, urine, liquor.

Separated and unhemolysed sample from the corpuscolated part are stable 8 hours at 25°C or 3 days at 2-8°C. Variable stability is observed with longer storage periods. In non-centrifuged sample glycolysis decreases

In non-centrifuged sample glycolysis decreases serum glucose by approximately 5-7% in an hour (5-10 mg/dl) at room temperature. The rate of in vitro glycolysis is higher in presence of leukocytosis or bacterial contamination.

Plasma, if removed from the cells after moderate centrifugation, contains leukocytes that also metabolize glucose, although cell-free sterile plasma has no glycolytic activity.

Glycolysis can be inhibited and glucose stabilized for as long as 3 days at room temperature by adding sodium idoacetate or sodium fluoride to the sample, even if it has no influence in the glycolysis during the first hour from the drawing.

The liquor can be contaminated with bacteria or other cells and may be analyzed immediately. If it's not possible making immediately the analysis the sample have to be centrifuged and stored at 4°C or -20°C.

In 24 hours collection of urine, glucose may be preserved adding 5 ml of acetic acid to the container before starting the collection. The final pH of urine, is usually between 4 and 5 which inhibits bacterial activity. Urine samples may loose as much as 40 % of glucose after 24 hours at room temperature.

## REAGENTS

Sole reagent: Phosphate buffer pH 7.00 200 mM,

GOD ≥ 15000 U/I, 4-AAP 1 mM, fenol 10 mM, surfactants.

Standard: glucose 100 mg/dl.

# MATERIALS REQUIRED BUT NOT SUPPLIED

Current laboratory instrumentation. Spectrophotometer UV/VIS with thermostatic cuvette holder. Automatic micropipettes. Glass or high quality polystyrene cuvettes.

# PRECAUTIONS

Reagent may contain some non-reactive and preservative components. It is suggested to handle carefully it, avoiding contact with skin and swallow. Perform the test according to the general "Good Laboratory Practice" (GPL) guidelines.

## **SAMPLES PREPARATION**

Reagents are supplied in liquid form and ready to use. Reagents are stored at 2-8°C until expiration date on the label, away from light sources or 60 days after first opening.

## **PROCEDURE**

 Kind of analysis:
 Final point

 Reading time:
 5 minutes

 Wavelength:
 510 nm (480-520)

 Temperature:
 37°C

 Lightpath:
 1 cm

 Zero:
 Blank reagent

Reagents	Blank	Standard	Sample
Distilled water	10 μl		
Standard		10 μl	
Sample			10 µl
Reagent	1 ml	1 ml	1 ml

#### CALCULATION

Serum/Plasma/ Spontaneous Urine Glucose (mg/dl)

(A Sample / A Standard) x 100

24 hours urine Glucose (mg/24h)

(A Sample / A Standard) x 100 x diuresis (dl)

### **EXPECTED VALUES**

Urine (fasting patient)

 $\begin{array}{ll} \mbox{Random urine} & < 300 \mbox{ mg/dl} \\ \mbox{24h urine} & < 500 \mbox{ mg/24h} \end{array}$ 

Each laboratory should establish appropriate reference intervals related to its population.

### **NOTES**

- If the results are incompatible with clinical presentation, they have to be evaluated within a total clinical study.
- Only for IVD use.

## CALIBRATION/QUALITY CONTROL

It is suggested to perform an internal quality control. For this purpose the following control sera on human base are available on request:

QN 0050 CH 10 x 5 ml

Control Sera normal values

QP 0050 CH 10 x 5 ml

Control Sera pathological values

## TEST PERFORMANCE

## Precision

Intra-assay (n = 25)	Mean (mg/dl)	SD (mg/dl)	CV%
Sample 1	87.64	0.8602	0.98
Sample 2	226.6	0.8660	0.38

Inter-assay (n = 25)	Mean (mg/dl)	SD (mg/dl)	CV%
Sample 1	87.64	1.1860	1.35
Sample 2	227.64	1.1503	0.51

## Sensibility/limit of detection

The method is able to discriminate until 1 mg/dl.

# Methods comparison

A comparison with a commercial available product gave the following results in a comparison on 36 samples of serum:

Glucose LTA = x
Glucose competitor = y

y = 0.97801x + 2.41589

A comparison with a commercial available product gave the following results in a comparison on 21 samples of urine:

Glucose LTA = x Glucose competitor = y

y = 0.97651x + 0.37795 r = 0.99825

#### Linearity

n = 21

The method is able to discriminate until 500 mg/dl. If the value is exceeded, is suggested to dilute sample 1+9 with saline and to repeat the test, multiplying the results bv10.

#### Interferences

No interference was observed by the presence of: hemoglobin  $\leq$  400 mg/dl bilirubin  $\leq$  20 mg/dl lipids  $\leq$  400 mg/dl

#### WASTE DISPOSAL

Product is intended for professional laboratories. Waste products must be handled as per relevant security cards and local regulations.

### **PACKAGING**

 CODE CC01900
 (400 TESTS)

 Sole reagent
 4 x 100 ml
 (liquid)

 Standard
 1 x 5 ml
 (liquid)

#### **REFERENCES**

Trinder P., - J. Clin. Path. 22. 158 (1969). Tietz Textbook of clinical Chemistry, Second Edition, Burtis – Ashwood (1994).

# MANUFACTURER

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## SYMBOLS

LOT

IVD Only for IVD use

REF Code number

LEF Code number

Storage temperature interval
Expiration date (year, month)

Lot of manufacturing

Marning, read enclosed documents

Warning, read enclosed documents

Read the directions

Biological risk

Mod. 01.06 (ver. 4.5 – 12/02/2009)



r = 0.9756