



PRESTIGE 24i DIGITOXIN

DIAGNOSTIC KIT FOR DETERMINATION OF DIGITOXIN CONCENTRATION

INTRODUCTION

Digitoxin is a cardiac glycoside used to treat congestive heart failure and certain arrhythmias. It increases the force and excitability of the heart muscle, slows down conduction in atrioventricular node and also reduces heart rate. Digitoxin is a glycoside obtained from *Digitalis lanata* and *Digitalis purpurea*. It is characterized by a high absorption from the alimentary tract and very long half-life. It is metabolized in the liver (over 80%). Digitoxin concentration monitoring is recommended because of a narrow therapeutic range and serious side effects: arrhythmia, disturbance of conduction. Using digitoxin under the control of concentration enables detection of dangerous side effects before clinical symptoms appear, also it allows to avoid underdosage - "sham therapy".

METHOD PRINCIPLE

Immunoturbidimetric method; inhibition of agglutination. Increase of absorbance measured at $\lambda = 700$ nm is inversely proportional to the concentration of digitoxin in the sample. Digitoxin which is present in the sample forms immune complexes with specific antibody contained in the first reagent. After adding a second reagent containing polystyrene latex particles coated with digitoxin, agglutination is inhibited in proportion to digitoxin concentration in the sample.

REAGENTS

Package

	Cat. No 4-250 (24-TRAY)	Cat. No 4-450 (36-TRAY)
1-Reagent	2 x 11.5 ml	2 x 10 ml
2-Reagent	2 x 7 ml	2 x 6.5 ml

Unopened reagents stored at 2-8°C are stable up to the expiry date printed on the package. After opening the reagents are stable for 21 days on board the analyser at 2-10°C. Protect from light!

Reagents composition

Bis-Tris buffer, monoclonal antibodies to digitoxin, polystyrene latex particles coated with digitoxin, sodium azide (< 0.1%).

Warnings and notes

- Products for in vitro diagnostic use only.
- The reagents should be used by suitably qualified laboratory personnel only in accordance with intended purpose.
- The reagents contain sodium azide as a preservative (< 0.1%). Avoid contact with skin and mucous membranes. Sodium azide can form high explosive metal azide combinations with lead and copper. Drains should be flushed well with a large amount of water when discarding the solution.
- Mix well Reagent 2 before first use. Avoid foam formation.
- For optimal reagents stability it is recommended to remove them from the system and store in tightly closed bottles at 2-8°C.

SPECIMEN

Serum. Samples may be stored up to 3 days at 2-8 °C. or longer at -20°C. Nevertheless it is recommended to perform the assay with freshly collected samples!

Avoid repeated freezing and thawing. Mix well the samples before analysis.

PROCEDURE

These reagents may be used in automatic analysers Prestige 24i, Biolis 24i and Sapphire 400.

1-Reagent and 2-Reagent are ready to use.

1-Reagent put on basic position in reagent tray.

2-Reagent put on start position in reagent tray.

For reagent blank 0.9% NaCl is recommended.

APPLICATION

Item name	DIGIT
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Data information

Units	ng/ml
Decimals	2

Analysis

Type	END
Main W.Length1	700
Sub W.Length2	
Method	Immunoturbidimetric

Corr

	Slope		Inter
Y=	1.000	X+	0.000

Calibration

Type	Linear
Factor	

Standard

#1	*	#4	*
#2	*	#5	*
#3	*	#6	*

Normal Range

	Male		Female	
	Low	High	Low	High
Serum				
Urine				
Plasma				
CSF				
Dialysis				
Other				

Aspiration

Kind	Double	
	Volume	
Sample	4.5	μl
Reagent1	150	
Reagent2	90	

Third Mix.	OFF
R1 Blank	Water-B

Monitor

0 Level Point	1
Span	3.000

Data Process

Read			Absorbance Limit	
	Start	End	Low	High
Main	53	54	0.000	
Sub	34	35	3.000	

Factor		Endpoint Limit	2.000
Blank correction:	1.0000	Linear Check (%)	90%

Dilution

Diluent	100 : Dil2
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Prozone Check

	Start	End	Limit (%)
First			
Second			Low
Third			Low

Auto Rerun SW

ON

Auto Rerun Range (Result)

	OFF Lower	OFF Higher
Serum		
Urine		
Plasma		
CSF		
Dialysis		
Other		

Auto Rerun Condition (Absorbance)

Absorbance Range		
	Lower	OFF
	Higher	OFF
Prozone Range		OFF

THERAPEUTIC RANGE

therapeutic concentration 10 – 30 ng/ml (13 – 39 nmol/l)

Some patients achieve the desired therapeutic response at levels outside this range, so it is recommended to consider the need to establish an individual therapeutic ranges for each patient.

QUALITY CONTROL

For internal quality control it is recommended to use the CORMAY DIGITOXIN CONTROLS (Cat. No 5-108) with each batch of samples. For the calibration of automatic analysers systems the CORMAY DIGITOXIN CALIBRATORS (Cat. No 5-114) is recommended.

The calibration curve should be prepared every 7 days, with change of reagent lot number or as required e.g. quality control findings outside the specified range.

PERFORMANCE CHARACTERISTICS

Metrological characteristics may differ from values presented below with the instrument used.

- **Sensitivity / Limit of Detection:** 0.90 ng/ml (1.18 nmol/l).
- **Linearity:** up to 80 ng/ml (104.8 nmol/l) If the digitoxin concentration exceeds 80 ng/ml, dilute the sample 1:5 with saline solution and repeat the assay. The dilution take into account when making the results.
- **Specificity / Interferences**
Haemoglobin up to 12 g/dl, bilirubin up to 19 mg/dl, triglyceride up to 1.1 g/dl do not interfere with the test.

Precision

Reproducibility (day to day) n = 80	Mean [ng/ml]	SD [ng/ml]	CV [%]
level 1	23.7	0.4	2.2
level 2	51.8	1.5	2.6
level 3	36.9	0.7	2.3

Method comparison

A comparison between CORMAY reagent (y) and commercially available assay (x) using 51 samples gave following results:

$$y = 0.95x + 3.32 \text{ ng/ml;}$$

$$R = 0.9797$$

(R – correlation coefficient)

WASTE MANAGEMENT

Please refer to local legal requirements.

LITERATURE

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