## GOT/AST FL IFCC

GO F080 CH	4 x 20 ml
GO F245 CH	12 x 20 ml
GO F400 CH	8 x 50 ml
GO F500 CH	5 x 100 ml
GO F600 CH	5 x 120 ml

#### **INTENDED USE**

Reagent for quantitative in vitro determination of GOT in biological fluids.

#### **SUMMARY OF TEST**

The aminotransferases (transaminases) constitute a group of enzymes that catalyze the interconversion of amino acids and -oxo-acids by transfer of amino group. Transaminases are widely distributed in animal tissues. Both AST and ALT are normally present in human plasma, bile, cerebrospinal fluid, and saliva, but none is found in urine unless a kidney lesion is present.

## PRINCIPLE OF THE METHOD

The enzyme aspartate aminotransferase (EC 2.6.1.1; L-Aspartate:2-Oxoglutarate Aminotransferase, AST or AspAT; Glutamate Oxaloacetate Transaminase, GOT) catalyzes the transaminase reaction between L-Aspartate and 2-Oxoglutarate. The 2-Oxaloacetate formed, is reduced to malate in the presence of MDH. As the reactions proceed, NADH is oxidized to NAD. The disappearance of NADH per unit time is followed by measuring the decrease in absorbance at 340 nm.

The present method has been made according to IFCC (2002).

## MATERIALS REQUIRED BUT NOT SUPPLIED

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

## KIT COMPONENTS

# For in vitro diagnostic use only.

The components of the kit are stable until expiration date on the label.

Keep away from direct light sources.

GOT R1	F080: 4 x 16 ml (liquid) blue cap
	F245: 12 x 16 ml (liquid) blue cap
	F400: 8 x 40 ml (liquid) blue cap
	F500: 4 x 100 ml (liquid) blue cap
	F600: 4 x 120 ml (liquid) blue cap

GOT R2 F080: 1 x 16 ml (liquid) red cap F245: 3 x 16 ml (liquid) red cap

F400: 2 x 40 ml (liquid) red cap F500: 1 x 100 ml (liquid) red cap F600: 1 x 120 ml (liquid) red cap

Composition in the test: Tris buffer 80 mM pH 7.65, L-aspartate 240 mM, 2-Oxoglutarate 12 mM, NADH 0.18 mM, MDH  $\geq$  600 U/I, LDH  $\geq$  900U/I.

Store all components at 2-8°C.

## REAGENT PREPARATION

# Serum as starter procedure:

Codes F080/F245: add 4 ml of reagent R2 to a bottle of reagent R1.

Code F400: add 10 ml of reagent R2 to a bottle of reagent R1.

Code F500/F600/100F: mix 1 part of reagent R2 with 4 parts of reagent R1.

Stability of working reagent: preferably within 30 days at 2-8°C, away from light sources.

## Reagent as starter procedure:

use separate reagents ready to use.

Stability: up to expiration date on labels at 2-8°C;

Stability since first opening of vials: preferably within 60 days at  $2\text{-}8^{\circ}\text{C}$ .

## **PRECAUTIONS**

GOT R1: Warning. Causes serious eye irritation (H319).

Causes skin irritation (H315). Wear protective gloves. Eye protection (P280). IF ON SKIN: Wash with plenty of water (P302+P352). IF IN

EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing (P305+P351+P338). If eye irritation persists: get medical advice (P337+P313).

GOT R2: It is not classified as hazardous.

#### **SPECIMEN**

Serum, plasma,

add sample:

Collect blood with a minimum of venous stasis. GOT is stable up to 4 days at 2-8°C or 1 month at -20°C.

Of its stable up to 4 days at 2-6 C of 1 month at -20 C

## TEST PROCEDURE (sample as starter)

Wavelenght: 340 nm
Ligthpath: 1 cm
Temperature: 37°C

dispense in cuvette working reagent: 1 ml

preincubate at 37°C for 5 minutes.

Mix, execute a first reading of absorbance after 90 seconds, incubating at  $37^{\circ}$ C. Perform other 3 readings at 60 seconds intervals. Calculate the  $\Delta$ A/min.

100 սl

## **TEST PROCEDURE** (reagent as starter)

Wavelenght: 340 nm
Ligthpath: 1 cm
Temperature: 37°C

dispense in cuvette reagent R1: 1 ml
add sample 125 μl
incubate at 37°C for 5 minutes.

Mix, execute a first reading of absorbance after 90 seconds, incubating at 37°C. Perform other 3 readings at 60 seconds intervals. Calculate the  $\Delta A/min$ .

## **RESULTS CALCULATION**

Perform calculation in units per litre, multiplying the  $\Delta A/min$  by the factor as it is indicated.

Activity in  $\mu$ kat/l: U/I x 0.0167 =  $\mu$ kat/l

## **EXPECTED VALUES**

Men: < 35 U/I ( $< 0.58 \mu \text{kat/I}$ ) Women: < 31 U/I ( $< 0.52 \mu \text{kat/I}$ )

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

## QUALITY CONTROL AND CALIBRATION

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

## QUANTINORM CHEMA

with normal or close to normal control values

# QUANTIPATH CHEMA

with pathological control values.

If required, a multiparametric, human based calibrator is available:

## **AUTOCAL H**

Please contact Customer Care for further information.

# TEST PERFORMANCE

## Linearity

the method is linear up to 440 U/I.

If a  $\Delta A$ /min of 0.200 is exceeded, it is suggested to dilute sample 1+9 with saline and to repeat the test, multiplying the result by 10.

## Sensitivity/limit of detection (LOD)

the limit of detection is 0.463 U/l.

## Interferences

no interference was observed by the presence of: hemoglobin ≤ 200 mg/dl

hemoglobin  $\leq$  200 mg/dl bilirubin  $\leq$  40 mg/dl lipids  $\leq$  500 mg/dl

## Precision

intra-assay (n=10)	mean (U/I)	SD (U/I)	CV%
sample 1	46.19	0.31	0.67
sample 2	137.25	0.92	0.67
:	(1.10)		
inter-assay (n=20)	mean (U/I)	SD (U/I)	CV%
sample 1	mean (U/I) 46.18	SD (U/I) 2.04	CV% 4.41

## Methods comparison

a comparison between Chema and a commercially available product gave the following results:

GOT Chema = x GOT competitor = y n = 83

y = 1.003x - 0.560 U/I  $r^2 = 0.990$ 

## **WASTE DISPOSAL**

This product is made to be used in professional laboratories.

P501: Dispose of contents according to national/international regulations.

## REFERENCES

J. Clin.Chem.Clin.Biochem 8 (1970) 658; 10 (1972) 182 Tietz Textbook of Clinical Chemistry, Second Edition, Burtis-Ashwood (1994).

HU Bergmeyer - Methods of enzymatic analysis, (1987). CCLM 2002; 40(7):725-733, Schumann et al. - IFCC reference procedure for aspartate aminotransferase.

#### **MANUFACTURER**

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### **SYMBOLS**

IVD in vitro diagnostic medical device

LOT batch code

use-by date

[]i

consult instructions for use

