PANCREATIC ISOAMYLASE EPS FL

5 x 16 ml

INTENDED USE

Reagent for quantitative in vitro determination of pancreatic isoamylase in biological fluids.

SUMMARY OF TEST

Two types of amylase may be distinguished, the pancreatic type (P-amylase) and the salivary type (S-amylase). Assays of amylase activity in serum and urine are largely of use in the diagnosis of diseases of the pancreas and in the investigation of pancreatic function. For such reason could be useful to have a distinct response for P-amylase.

PRINCIPLE OF THE METHOD

The enzyme α -amylase (EC 3.2.1.1, 1,4 α -D-glucose glucanohydrolase) hydrolizes the EPS to release several different fragments. The fragments so formed are completely hydrolyzed to 4-nitrophenol and glucose by α -glucosidase. The selective inhibition of S-amylase is performed by use of two different monoclonal antibodies.

The 4-nitrophenol formed is detected spectrophotometrically at 405 nm to give a measurement of α -amylase activity in the sample.

The present method has been made according to IFCC.

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. **DO NOT PIPETTE BY MOUTH!**

AMY-P R1 F080: 4 x 16 ml (liquid) blue cap

AMY-P R2 F080: 1 x 16 ml (liquid) red cap

Composition in the test: Hepes buffer 50 mM pH 7.10, NaCl 70 mM, calcium acetate 1.0 mM, 4,6-Ethylidene(G1)-4-nitrophenyl(G7)- α -(1->4)-D-maltoheptaoside 5.0 mM, α -glucosidase 6 kU/l, monoclonal antibodies (mouse) \geq 25 mg/l.

Store all components at 2-8°C.

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.

REAGENT PREPARATION

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 2-8°C.

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" (GLP) guidelines.

SPECIMEN

Serum, plasma (heparinate only).

Amylase is stable in serum and plasma sample up to 2 months at 2-8°C.

TEST PROCEDURE

Wavelenght: Ligthpath: Temperature:	405 nm 1 cm 37°C		
dispense in cuvette reagent R1:			1 ml
add sample:			25 μl
incubate at 37°C for 5 minutes.			
dispense in cuvette reagent R2:			250 μl

Mix, execute a first reading of absorbance after 1 minute, 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 ΔA /min by the factor as it is indicated.

Calculation in U/I: $\Delta A/\min x 6280$

Calculation in µkat/l: U/l x 0.0167 = µkat/l

EXPECTED VALUES

Serum - plasma:	13 - 53 U/l	(0.22 - 0.88 μkat/l)
Random urine:	≤ 350 U/l	(≤ 5.84 μkat/l)

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 2500 U/l. If a ΔA /min of 0.500 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 2 U/I.

Interferences

Precision	
lipids	interfer in low values
bilirubin	≤ 25 mg/dl
hemoglobin	≤ 500 mg/dl
no interference was of	oserved by the presence of:

intra-assay (n=10) sample 1 sample 2	mean (U/I) 38.00 103.00	SD (U/I) 0.67 1.41	CV% 1.80 1.40
inter-assay (n=20)	mean (U/I)	SD (U/I)	CV%
sample 2	102.61	1.62	2.50

Methods comparison

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

Isoamylase Chema = x
Isoamylase competitor = y
n = 108

y = 1.02x - 0.605 U/l r² = 0.997

WASTE DISPOSAL

This product is made to be used in professional laboratories.

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

REFERENCES

Clin.Chem. 33, 1158-1162 (1987)

Lab.Med. 12 110-113 (1989) Clin.Chem.Lab.Med. 1998; 36(3):185-203

Junge W, Waldenström J, Bouman A et al. Evaluation of the Assays for Total and Pancreatic α -Amylase based on 100% Cleavage of Et-G7-PNP at 6 European Clinical Centres (Poster Medlab 97). Basel, Switzerland: 12th IFCC Euro-

pean Congress of Clinical Chemistry, 17–22 August 1997. MANUFACTURER

Chema Diagnostica Via Campania 2/4 60030 Monsano (AN) - ITALY - EU phone +39 0731 605674 fax +39 0731 605672 e-mail: mail@chema.com website: http://www.chema.com

SYMBOLS

IVD	in vitro diagnostic medical device
LOT	batch code
REF	catalogue number
X	temperature limit
Σ	use-by date

▲ caution

consult instructions for use