



Liquick- Cor UA

Kit name	Cat. No
Liquick Cor-UA mini	2-224
Liquick Cor-UA 30	2-235
Liquick Cor-UA 60	2-208
Liquick Cor-UA 120	2-209

INTENDED USE

Diagnostic kit for determination of uric acid concentration used both for manual assay (Sample Start and Reagent Start method) and in several automatic analysers.

The reagents must be used only for in vitro diagnostic, by suitably qualified laboratory personnel, only for the intended purpose, under appropriate laboratory conditions.

INTRODUCTION

Uric acid is a product of purine catabolism. It is produced in the liver and excreted in the urine. Both, the amount of uric acid production and the efficiency of renal excretion, affect serum urate level. Elevated serum uric acid level is caused usually by gout, leukemia, diabetes mellitus, hyperfunction of parathyroid and thyroid, renal failure, renal calculosis. Urate concentration in serum depends on glomerular filtration, thus is useful for renal function monitoring.

METHOD PRINCIPLE

Enzymatic, colorimetric method with uricase and peroxidase.

uric acid + 2
$$\text{H}_2\text{O}$$
 + O_2 $\xrightarrow{\text{uricase}}$ allantoine + CO_2 + H_2O_2

ADPS + 4-aminoantipyrine + 2 H_2O_2 $\xrightarrow{\text{POD}}$ quinoneimine dye + $\text{4H}_2\text{O}$

(coloured compound)

The colour intensity is proportional to the uric acid concentration.

REAGENTS Package

g.	Liquick Cor-UA mini	Liquick Cor-UA 30	Liquick Cor-UA 60	Liquick Cor-UA 120
1-UA	2 x 24 ml	5 x 24 ml	5 x 48 ml	5 x 96 ml
2-UA	1 x 12 ml	1 x 30 ml	1 x 60 ml	1 x 120 m
3-STANDARD	1 x 1 ml	1 x 2 ml	-	-

3-STANDARD is uric acid standard solution: 300 $\mu mol/l$ (5.05 mg/dl).

The reagents when stored at 2-8°C are stable up to expiry date printed on the package. The reagents are stable for 12 weeks on board the analyser at 2-10°C.

Working reagent preparation and stability

Assay can be performed with use of separate 1-UA and 2-UA reagents or with use of working reagent. For working reagent preparation mix gently 4 parts of 1-UA with 1 part of 2-UA. Avoid foaming.

Stability of working reagent: 3 months at 2-8°C 2 weeks at 15-25°C

Concentrations in the test

Concenti auons in the test	
buffer PIPES (pH 7.0)	100 mmol/l
4-aminoantipyrine	0.78 mmol/l
ADPS	0.67 mmol/l
ferricyanide potassium	3.8 µmol/l
Liquick Cor-UA	

peroxidase (POD) > 38.34 μkat/l uricase > 1.65 μkat/l

Warnings and notes

- Protect from light and avoid contamination!
- The reagents and standards contain sodium azide (< 0.1%) as a preservative. Avoid contact with skin and mucous membranes.
- 1-UA meeting the criteria for classification in accordance with Regulation (EC) No 1272/2008.

Warning



H315 Causes skin irritation.

H319 Causes serious eye irritation.

P280 Wear protective gloves/protective clothing/eye protection/face protection.

P302+P352 IF ON SKIN: Wash with plenty of soap

and water.

P305+P351+P338 IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

 Kit Liquick Cor-UA PLUS (Cat. No 2-225, 2-260, 2-258, 2-259) is recommended for determination of uric acid concentration in urine. This reagent contains ascorbic acid oxidase which eliminates interference of ascorbic acid in the assay.

ADDITIONAL EQUIPMENT

- automatic analyzer or photometer able to read at 546 nm (Hg 530-550 nm);
- thermostat at 25°C or 37°C;
- general laboratory equipment;

SPECIMEN

Serum, heparinized plasma free from hemolysis.

Do not use EDTA and fluoride as anticoagulants.

Specimen can be stored 3-5 days at 2-8°C or 6 months at -20°C. Nevertheless it is recommended to perform the assay with freshly collected samples!

PROCEDURE

Applications for them are available on request.

Manual procedure

wavelength 546 nm (Hg 530-550 nm) temperature 25°C / 37°C

cuvette 1 cm

Sample Start method

Pinette into the cuvettes:

	reagent blank	test	standard
	(RB)	(T)	(S)
working reagent	1000 μ1	1000 μ1	1000 μ1
Bring up to the tempe	erature of determinati	on. Then add	1:
standard	-	-	20 μ1
sample	-	20 μ1	-

Mix well, incubate for 10 min. at 25°C or 5 min. at 37°C. Read the absorbance of the test A(T) and standard A(S) against reagent blank (RB).

Reagent Start method

The determination can be also performed with use of separate 1-UA and 2-UA reagents.

	reagent blank (RB)	test (T)	standard (S)	
1-UA	1000 μ1	1000 μ1	1000 μ1	
Bring up to the temperature of determination. Then add:				
standard	-	-	20 μ1	

2-U	A		250 μ1		250	μl	250	μl
Mix	well:	perform	measurement a	des	scribed	for	Sample	Start

method.

uric acid	_	A(T)	**	standard / calibrator
concentration	=	A(S)	х	concentration

REFERENCE VALUES 5

Mix well, incubate for 5 min. Then add:

serum / plasma	mg/dl	μmol/l
females	2.5 - 6.8	149 – 405
males	3.6 – 7.7	214 – 458

It is recommended for each laboratory to establish its own reference ranges for local population.

QUALITY CONTROL

For internal quality control it is recommended to use the CORMAY SERUM HN (Cat. No 5-172) and CORMAY SERUM HP (Cat. No 5-173) with each batch of samples.

For calibration when using <u>manual methods</u> URIC ACID STANDARD 5 (Cat. No 5-125) or 3-STANDARD attached to the kit is recommended.

For calibration of the <u>automatic analysers systems</u> CORMAY MULTICALIBRATOR LEVEL 1 (Cat. No 5-174 and 5-176) and LEVEL 2 (Cat. No 5-175 and 5-177) is recommended.

The calibration curve should be prepared every 12 weeks, with change of reagent lot number or as required e.g. quality control findings outside the specified range.

PERFORMANCE CHARACTERISTICS

These metrological characteristics have been obtained using automatic analyser Biolis 24i Premium. Results may vary if a different instrument or a manual procedure is used.

- Sensitivity: 0.31 mg/dl (18.44 µmol/l).
- Linearity: up to 23 mg/dl (1368 µmol/l).

Specificity / Interferences

Haemoglobin up to 1.25 g/dl, ascorbate up to 31 mg/l, bilirubin up to 20 mg/dl and triglycerides up to 1000 mg/dl do not interfere with the test.

Precision

Repeatability (run to run)	Mean	SD	CV
n = 20	[mg/dl]	[mg/dl]	[%]
level 1	4.85	0.05	1.03
level 2	8.99	0.09	1.01

Reproducibility (day to day)	Mean	SD	CV
n = 80	[mg/dl]	[mg/dl]	[%]
level 1	4.83	0.07	1.39
level 2	9.03	0.16	1.78

Method comparison

A comparison between uric acid values obtained on Biolis 24i Premium (y) and at ADVIA 1650 (x) using 100 samples gave following results:

y = 0.9936 x + 0.1225 mg/dl;

R = 0.9965 (R – correlation coefficient)

TRACEABILITY

URIC ACID STANDARD 5 is traceable to the SRM 1950/909C reference material.

WASTE MANAGEMENT

Please refer to local legal requirements.

LITERATURE

- 1. Thefeld C. et al.: Dtsch. Med. Wschr. 98, 380-384 (1973).
- 2. Barham D., Trinder P.: Analyst 97, 142-145 (1972).
- 3. Fossati P., Prencipe L., Berti G.: Clin. Chem. 26/2, 227-231 (1980)
- Henry R.J.: Clinical Chemistry, Harper & Row Publishers Inc., New York (1974).
- Kaplan L.A., Pesce A.J., ed. Chemistry Theory, Analysis, and Correlation, 3rd ed. St Louis, MO: Mosby, 501-2 (1996).
- Tietz N.W., ed. Clinical Guide to Laboratory Tests, 3rd ed. Philadelphia, PA: WB Saunders, 624, (1995).

Date of issue: 05. 2018.

str./ page/ crp. 3/6 Liquick Cor-UA str./ page/ crp. 4/6 51 03 08 061 02 51 03 08 061 02