

LDH 21 FS*

Order Information

Cat. No.

1 4251 99 10 920

Kit size



800 (4 x 200)

Intended Use

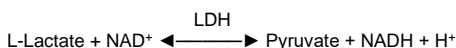
Diagnostic reagent for quantitative in vitro determination of lactate dehydrogenase activity in human serum or heparin plasma on automated respons[®]920.

Summary

Lactate dehydrogenase (LDH) is an enzyme, consisting of five different isoenzymes, which catalyze the interconversion of L-lactate and pyruvate with concomitant interconversion of NADH and NAD⁺. LDH is present in the cytoplasm of all human tissues with higher concentrations in liver, heart and skeletal muscle and kidney and lower values in erythrocytes [1]. Increased LDH activities are found in a variety of pathological conditions such as myocardial infarction, cancer, diseases of liver, blood or muscle [1,2]. However, because of the lack of organ specificity, determination of its isoenzymes or other enzymes such as alkaline phosphatase or ALAT/ASAT is necessary for differential diagnosis [1,2].

Method

Optimized UV-test according to IFCC (International Federation of Clinical Chemistry and Laboratory Medicine) [modified].



One unit of LDH is the amount of enzyme required to produce 1.0 μmol of pyruvate per minute under enzyme specific conditions.

Reagents

Components and Concentrations

R1:	N-Methyl-D-Glucamine	pH 8.4	420 mmol/L
	L-Lactate		65 mmol/L
R2:	NAD ⁺		50 mmol/L

Storage and Stability

Reagents are stable up to the date of expiry indicated on the kit, if stored at 2 – 8°C and contamination is avoided. Do not freeze and protect from light.

The open-vial stability of the reagent is 24 months until expiry date.

Warnings and Precautions

1. Reagent 1 contains sodium azide (0.95 g/L) as preservative. Do not swallow! Avoid contact with skin and mucous membranes.
2. Reagent 1 contains material of biological origin. Handle the product as potentially infectious according to universal precautions and good clinical laboratory practice.
3. In very rare cases, samples of patients with gammopathy might give falsified results [3].
4. To avoid carryover interference, please take care of efficient washing especially after use of interfering reagents. Please refer to the DiaSys respons[®]920 Carryover Pair Table. Carryover pairs and automated washing steps with the recommended cleaning solution can be specified in the system software. Please refer to the user manual.
5. In case of product malfunction or altered appearance that could affect the performance, contact the manufacturer.
6. Any serious incident related to the product must be reported to the manufacturer and the competent authority of the Member State where the user and/or patient is located.
7. Please refer to the safety data sheets (SDS) and take the necessary precautions for the use of laboratory reagents. For diagnostic purposes, the results should always be assessed with the patient's medical history, clinical examinations and other findings.
8. For professional use only.

Waste Management

Refer to local legal requirements for chemical disposal regulations as stated in the relevant SDS to determine the safe disposal.

Warning: Handle waste as potentially biohazardous material. Dispose of waste according to accepted laboratory instructions and procedures.

Reagent Preparation

The reagents are ready to use. The bottles are placed directly into the reagent rotor.

Materials Required

General laboratory equipment

Specimen

Human serum or heparin plasma

Only use suitable tubes or collection containers for specimen collection and preparation.

When using primary tubes, follow the manufacturer's instructions.

Stability [4]:

7 days	at	20 – 25°C
4 days	at	4 – 8°C
6 weeks	at	-20°C

Only freeze once. Discard contaminated specimens.

Calibrators and Controls

DiaSys TruCal U is recommended for calibration. Calibrator values have been standardized against the original IFCC formulation. Use DiaSys TruLab N and P for internal quality control. All target values of the controls are traceable to DiaSys reagent/calibrator system. Quality control must be performed after calibration. Control intervals and limits have to be adapted to the individual requirements of each laboratory. Results must be within the defined ranges. Follow the relevant legal requirements and guidelines. Each laboratory should establish corrective action in case of deviations in control recovery.

	Cat. No.	Kit size
TruCal U	5 9100 99 10 063	20 x 3 mL
	5 9100 99 10 064	6 x 3 mL
TruLab N	5 9000 99 10 062	20 x 5 mL
	5 9000 99 10 061	6 x 5 mL
TruLab P	5 9050 99 10 062	20 x 5 mL
	5 9050 99 10 061	6 x 5 mL

Performance Characteristics

Measuring range from 18 U/L up to 1500 U/L. Linearity < 15 U/L is given with ± 5.4 U/L, between 15 U/L to 30 U/L within ± 10%, at > 30 U/L within ± 5%. In case of higher activities re-measure samples after manual dilution with NaCl solution (9 g/L) or use rerun function.	
Limit of detection**	18 U/L
Limit of quantitation**	18 U/L
Onboard stability	12 weeks
Calibration stability	12 weeks

Interference by	Interferences ≤ 10% up to	Analyte concentration [U/L]
Ascorbic acid	30 mg/dL	166
	30 mg/dL	243
Bilirubin (conjugated)	60 mg/dL	154
	60 mg/dL	235
Bilirubin (unconjugated)	50 mg/dL	162
	50 mg/dL	252
Lipemia (triglycerides)	2000 mg/dL	168
	2000 mg/dL	240
Sulfapyridine	30 mg/dL	157
	30 mg/dL	238
Sulfasalazine	30 mg/dL	172
	30 mg/dL	253

Hemoglobin interferes at low concentrations.

For further information on interfering substances, refer to the literature [5-7].

Precision			
Repeatability (n=20)	Sample 1	Sample 2	Sample 3
Mean [U/L]	107	243	977
CV [%]	1.87	1.24	2.15
Within-laboratory (n=80)	Sample 1	Sample 2	Sample 3
Mean [U/L]	104	257	982
CV [%]	3.01	3.08	2.91

Method comparison (n=210)	
Test x	Competitor LDH (cobas c 501)
Test y	DiaSys LDH 21 FS (respons [®] 920)
Slope	0.948
Intercept	0.799 U/L
Coefficient of correlation	0.998

** according to CLSI document EP17-A2, Vol. 32, No. 8

Conversion Factor

LDH [U/L] x 0.0167 = LDH [μkat/L]

Reference Range [1]

	U/L	μkat/L
Children		
0 – 1 year	196 – 438	3.27 – 7.3
1 – 3 year(s)	105 – 338	1.75 – 5.6
4 – 6 years	107 – 314	1.78 – 5.2
7 – 11 years	112 – 307	1.87 – 5.1
13 – 17 years	115 – 287	1.94 – 4.8
Adults		
Female	< 247	< 4.12
Male	< 248	< 4.13

Consensus for upper reference limits for adults: < 250 U/L (4.20 μkat/L)

Each laboratory should check if the reference ranges are transferable to its own patient population and determine own reference ranges if necessary.

Literature

1. Thomas L. Clinical Laboratory Diagnostics [Internet]. Prof. Lothar Thomas; 2024 [cited 2024 June 10]. Available from: <https://www.clinical-laboratory-diagnostics.com/>
2. Moss DW, Henderson AR. Clinical enzymology In: Burtis CA, Ashwood ER, editors. Tietz Textbook of Clinical Chemistry. 4th ed. St. Louis Missouri: Elsevier Saunders Company;2006. 601-604.
3. Bakker AJ, Mücke M. Gammopathy interference in clinical chemistry assays: Mechanisms, detection and prevention. Clin Chem Lab med 2007; 45(9): 1240-1243.

4. Guder WG, da Fonseca-Wollheim F, Heil W, Schmitt Y, Töpfer G, Wisser H, Zawta B. Quality of Diagnostic Samples. 3rd edition; 2010. p. 52-3.
5. Young DS. Effects of Drugs on Clinical Laboratory Tests. 5th ed. Volume 1 and 2. Washington, DC: The American Association for Clinical Chemistry Press 2000.
6. Young DS. Effects on Clinical Laboratory Tests - Drugs Disease, Herbs & Natural Products, <https://clinf.wiley.com/aaccweb/aacc/>, accessed in March 2021. Published by AACC Press and John Wiley and Sons, Inc.
7. Sonntag O, Scholer A. Drug interference in clinical chemistry: recommendation of drugs and their concentrations to be used in drug interference studies. Ann Clin Biochem. 2001 Jul;38:376-85.

Additions and/or changes in the document are highlighted in grey. Deletions are communicated via customer info by stating the edition no. of the package insert/instruction for use.



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* Fluid Stable

LDH 21 FS

Application for serum and plasma

Test Details		Test Volumes		Reference Ranges	
Test	: LDH21			Auto Rerun	<input type="checkbox"/>
Report Name	: LDH 21 IFCC			Online Calibration	<input type="checkbox"/>
Unit	: U/L	Decimal Places	: 1	Cuvette Wash	<input type="checkbox"/>
Wavelength-Primary	: 340	Secondary	: 405	Total Reagents	: 2
Assay Type	: RATE - A	Curve Type	: Linear	Reagent R1	: LDH21 R1
M1 Start	: 0	M1 End	: 0	Reagent R2	: LDH21 R2
M2 Start	: 21	M2 End	: 33	Consumables/Calibrators:	
Sample Replicates	: 1	Standard Replicates	: 3	Blank/Level 0	: 0
Control Replicates	: 1	Control Interval	: 0	Calibrator 1	: *
Reaction Direction	: Increasing	React. Abs. Limit	: 1.20		
Prozone Limit %	: 0	Prozone Check	: Lower		
Linearity Limit %	: 0	Delta Abs./Min.	: 0.00		
Technical Minimum	: 18.00	Technical Maximum	: 1500.00		
Y = aX + b	a = 1.00	b =	: 0.00		

* Enter calibrator value.

Test Details		Test Volumes		Reference Ranges	
Test	: LDH21				
Sample Type	: Serum				
Sample Volumes				Sample Types	
Normal	: 3.00 μ L	Dilution Ratio	: 1 X	<input checked="" type="checkbox"/> Serum <input type="checkbox"/> Urine <input type="checkbox"/> CSF <input checked="" type="checkbox"/> Plasma <input type="checkbox"/> Whole Blood <input type="checkbox"/> Other	
Increase	: 6.00 μ L	Dilution Ratio	: 1 X		
Decrease	: 2.00 μ L	Dilution Ratio	: 1 X		
Standard Volume	: 3.00 μ L				
Reagent Volumes and Stirrer Speed					
RGT-1 Volume	: 160 μ L	R1 Stirrer Speed	: Medium		
RGT-2 Volume	: 40 μ L	R2 Stirrer Speed	: High		

Test Details		Test Volumes		Reference Ranges	
Test	: LDH21				
Sample Type	: Serum				
Reference Range	: DEFAULT				
Category	: Male				
Reference Range				Sample Types	
	Lower Limit		Upper Limit	<input checked="" type="checkbox"/> Serum <input type="checkbox"/> Urine <input type="checkbox"/> CSF <input checked="" type="checkbox"/> Plasma <input type="checkbox"/> Whole Blood <input type="checkbox"/> Other	
	(U/L)		(U/L)		
Normal	: 0.00		: 248.00		
Panic	: 0.00		: 0.00		