

CK-MB FS*

Order Information

Cat. No.

1 1641 99 10 921

Kit size

 480 (4 x 120)

Intended Use

Diagnostic reagent for quantitative in vitro determination of CK-MB in human serum or lithium heparin plasma on automated DiaSys respons[®]920.

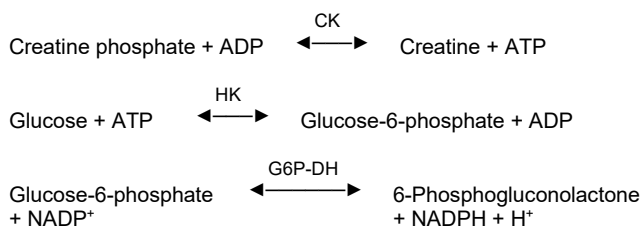
Summary

Creatine kinase (CK) is an enzyme, which consists of isoenzymes mainly of the muscle (CK-M) and the brain (CK-B). CK exists in the human body in dimeric forms as CK-MM, CK-MB, CK-BB and as macro-enzyme. Measurement of CK-MB is a specific test for the detection of cardiac muscle damage and, therefore, is used for diagnosis and monitoring of myocardial infarction. [1,2,3]

Method [4]

Optimized UV test according to DGKC (German Society of Clinical Chemistry) and IFCC (International Federation of Clinical Chemistry and Laboratory Medicine) for CK with inhibition of CK-M isoenzymes by monoclonal antibodies.

CK-MB consists of the subunits CK-M and CK-B. Specific antibodies against CK-M inhibit the complete CK-MM activity (main part of the total CK activity) and the CK-M subunit of CK-MB. Only CK-B activity is measured, which is half of the CK-MB activity.



Reagents

Components and Concentrations

R1:	Imidazole/Good's buffer	120 mmol/L
	Glucose	25 mmol/L
	N-Acetylcysteine (NAC)	25 mmol/L
	Magnesium acetate	12.5 mmol/L
	EDTA-Na ₂	2 mmol/L
	NADP	2.5 mmol/L
	Hexokinase (HK)	≥ 5 kU/L
	Monoclonal antibodies against human CK-M (mouse); inhibiting capacity	≥ 2500 U/L
R2:	Imidazole/Good's buffer	90 mmol/L
	ADP	10 mmol/L
	AMP	28 mmol/L
	Glucose-6-phosphate dehydrogenase (G6P-DH)	≥ 15 kU/L
	Diadenosine pentaphosphate	50 μmol/L
	Creatine phosphate	150 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. Protect from light.

Warnings and Precautions

- ⚠ Reagent 1 and 2: Danger. Contains: Imidazole. H360D May damage the unborn child. P201 Obtain special instructions before use. P280 Wear protective gloves/protective clothing/eye protection. P308+P313 If exposed or concerned: Get medical advice/attention.
- The reagents contain sodium azide (0.95 g/L) as preservative. Do not swallow! Avoid contact with skin and mucous membranes.
- The reagents contain animal and biological material. Handle the product as potentially infectious according to universal precautions and good clinical laboratory practice.
- 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.
- In very rare cases, samples of patients with gammopathy might give falsified results [5].
- Sulfasalazine medication may cause false results in patient samples. Blood collection must be performed prior to drug administration.
- Heterophile antibodies in patient samples may cause falsified results.
- Please refer to the safety data sheets 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.
- For professional use only.

Waste Management

Refer to local legal requirements.

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 lithium heparin plasma

Stability [6]:

2 days	at	20 – 25°C
7 days	at	4 – 8°C
4 weeks	at	–20°C

Only freeze once. Discard contaminated specimens.

Calibrators and Controls

DiaSys TruCal CK-MB is recommended for calibration. TruCal CK-MB calibrator values have been made traceable to the molar extinction coefficient. Control sera and calibrators containing non-human CK-MB fractions are not suitable to be applied with this test due to the monoclonal antibody used in the reagent. Only use controls and calibrators containing exclusively human CK-MB. Use DiaSys TruLab N and P for internal quality control. Each laboratory should establish corrective action in case of deviations in control recovery.

	Cat. No.	Kit size
TruCal CK-MB	5 9450 99 10 074	6 x 1 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

Exemplary data mentioned below may slightly differ in case of deviating measurement conditions.

Measuring range up to 2000 U/L. In case of higher activities re-measure samples after manual dilution with NaCl solution (9 g/L) or use rerun function.	
Limit of detection**	3 U/L
Onboard stability	4 weeks
Calibration stability	4 weeks

Interfering substance	Interferences ≤ 10% up to
Ascorbic acid	30 mg/dL
Bilirubin (conjugated and unconjugated)	25 mg/dL
Hemoglobin	25 mg/dL
Lipemia (triglycerides)	900 mg/dL

For further information on interfering substances refer to Young DS [7,8].

Precision			
Within run (n=20)	Sample 1	Sample 2	Sample 3
Mean [U/L]	21.0	49.0	114
CV [%]	2.49	1.35	2.00
Between day (n=20)	Sample 1	Sample 2	Sample 3
Mean [U/L]	39.5	73.0	216
CV [%]	2.50	1.78	1.27

Method comparison (n=103)	
Test x	DiaSys CK-MB FS (Hitachi 917)
Test y	DiaSys CK-MB FS (respons [®] 920)
Slope	1.05
Intercept	-1.88 U/L
Coefficient of correlation	0.998

** lowest measurable activity which can be distinguished from zero; mean + 3 SD (n = 20) of an analyte free specimen.

Conversion Factor

CK-MB [U/L] x 0.0167 = CK-MB [μkat/L]

Reference Range

Myocardial infarction: the risk of myocardial infarction is high if following three conditions are fulfilled [9]:

1. CK (Men) > 190 U/L (3.17 μkat/L)***
CK (Women) > 167 U/L (2.78 μkat/L)***
2. CK-MB > 24 U/L (0.40 μkat/L)***
3. CK-MB activity is between 6 and 25% of total CK activity.

***calculated using temperature conversion factor 2.38 (25°C → 37°C)

If myocardial infarction is suspected and the conditions are not fulfilled, the infarction may be fresh. In this case the measurements should be repeated after 4 hours with fresh samples.

In healthy individuals different values are found depending on race and age [9,10].

Each laboratory should check if the reference ranges are transferable to its own patient population and determine own reference ranges if necessary. For diagnostic purposes, CK values should always be assessed in conjunction with the anamnesis, the clinical examination and other findings.

Literature

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* Fluid Stable

CK-MB FS

Application for serum and plasma

Test Details		Test Volumes		Reference Ranges	
Test	: CKMB			Auto Rerun	<input type="checkbox"/>
Report Name	: CK-MB			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	: CKMB R1
M1 Start	: 0	M1 End	: 0	Reagent R2	: CKMB R2
M2 Start	: 26	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	: 1.5000		
Prozone Limit %	: 0	Prozone Check	: Lower		
Linearity Limit %	: 0	Delta Abs./Min.	: 0.0000		
Technical Minimum	: 3.0000	Technical Maximum	: 2000.0000		
Y = aX + b	a = 1.0000	b = 0.0000			

* Enter calibrator value

Test Details		Test Volumes		Reference Ranges	
Test	: CKMB				
Sample Type	: Serum				
Sample Volumes				Sample Types	
Normal	: 8.00 μ L	Dilution Ratio	: 1 X	<input checked="" type="checkbox"/> Serum	
Increase	: 16.00 μ L	Dilution Ratio	: 1 X	<input type="checkbox"/> Urine	
Decrease	: 4.00 μ L	Dilution Ratio	: 1 X	<input type="checkbox"/> CSF	
				<input checked="" type="checkbox"/> Plasma	
				<input type="checkbox"/> Whole Blood	
				<input type="checkbox"/> Other	
Standard Volume	: 8.00 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	: CKMB				
Sample Type	: Serum				
Reference Range	: DEFAULT				
Category	: Male				
Reference Range				Sample Types	
	Lower Limit		Upper Limit	<input checked="" type="checkbox"/> Serum	
	(U/L)		(U/L)	<input type="checkbox"/> Urine	
Normal	: 0.00		: 24.00	<input type="checkbox"/> CSF	
Panic	: 0.00		: 0.00	<input checked="" type="checkbox"/> Plasma	
				<input type="checkbox"/> Whole Blood	
				<input type="checkbox"/> Other	