FRUCTOSAMINE



Intended use. System for fructosamine determination by fixed-time kinetic method in serum samples.

Professional use.

[For in vitro diagnostic use.]

Test principle. Glucose binds to amino groups of proteins yielding a Schiff's base (aldimine) that after a molecular rearrange, transforms to a stable ketoamine generally known as fructosamine¹.

In alkaline pH fructosamine is converted to an enolic form that reduces nitro blue tetrazolium to a "purple formazan". The absorbance difference. after incubation in 10 minutes and 15 minutes, is proportional to fructosamine concentration in the sample². The system calibration is performed with bovine matrix calibrator, calibrated with glycated polylysine. The results are presented as micromoles/liter (µmol/L).

Summary . Fructosamine determination, vielded from the bind of glucose and plasmatic proteins, is based on its reduction ability in alkaline medium. Other reducer agents may be present in serum sample and interferes in the assay. Labtest incorporated uricase and a clarifier agent based on detergents in the Fructosamine system in order to minimize the interferences of uric acid and lipemia.

The assay calibration with the glycated albumin calibrator in vitro, calibrated by glycated polylysine containing glucose labeled with ¹⁴C, makes that small variations of test condition do not importantly interfere². The system benefits are more consistent results with more reproducibility and accuracy.

The system uses a Work Reagent stable for 30 days, if stored at 2 - 8 °C. The system also allows preparing the volume of the Work Reagent needed to one measure of the Fructosamine concentration.

It is easily applied to most automatic equipments which are able to measure, in a kinetic method, absorbances difference at 530 nm⁵

Methodology . NBT reduction.

Reagents

1. RI - Reagent 1 - Store at 2 - 8°C.

Reagent label bears expiration date. Buffer pH 7.3 (83 mmol/L); nitro blue tetrazolium (NBT) (967 µmol/L); uricase (≥5000 U/L); sodium azide (14.6 mmol/L) stabilizer and surfactants.

2. R 2 - Reagent 2 - Store at 2 - 8 °C.

Reagent label bears expiration date. Buffer pH 10.4 (625 mmol/L): sodium azide (14.6 mmol/L).

3. CAL - Calibrator - Store at 2 - 8 °C.

Reagent label bears expiration date and calibrator concentration. Glycated Bovine albumin, Buffer pH 7.4 (50 mmol/L); and sodium azide (14.6 mmol/L).

Precautions and warnings

Disposal of all waste material should be in accordance with local auidelines

The usual security cares should be applied on the reagent handling. Do not pipette the reagent with mouth.

The reagents contain sodium azide as preservative. Avoid ingestion. In case of eyes contact, immediately flush eyes with plenty of water and get medical assistance.

Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. On disposal, flush with a large volume of water to prevent azide accumulation.

Temperature and incubation time controls during the measurement must be rigorous. 1 °C difference in the temperature produces 5% error, while 1 minute difference during the ΔA measurement produces a 20% error.

Storage and stability. Unopened reagents, when stored at indicated temperature, are stable up to expiration date shown on the label.

Deterioration. The Working Reagent is an alkaline solution (pH= 10.3) and, as such, is instable when expose to ambient atmosphere. In order to conserve its performance, the Working Reagent must be opened only the time needed to get the volume desired. Keep the bottle tightly closed.

Specimen collection and preparation

Use serum or plasma (EDTA and heparin) without hemolysis. The analite is reportedly stable for about 7 days at 2 - 8 °C and 3 months at -20 °C.

No known test method can offer complete assurance that human blood samples will not transmit infectious diseases. Therefore, all blood derivatives should be considered potentially infectious.



Interference

Bilirubin up to 8.0 mg/dL, and glucose up to 1000 mg/dL do not interfere significantly.

High ascorbic acid concentrations over 3.0 mg/dL and hemoglobin concentrations over 100 mg/dL result in significant negative interference.

Triglycerides concentrations up to 1000 mg/dL and uric acid up to 14 mg/dL do not significantly interfere.

Materials required not provided

- 1. A constant temperature water bath (37 °C).
- 2. Photometer capable of measuring absorbance at 510 550 nm.
- 3. Pipettes to measure reagents and samples.
- 4. Timer

Preparing the reagents

Working reagent. Use one bottle of Reagent 1 and Reagent 2 for preparing Working Reagent. Transfer all the contents of one Reagent 2 bottle to one Reagent 1 bottle and mix by inversion.

The Working Reagent is stable 30 days at 2 - 8 °C, when no chemistry or microbiological contamination occurs (See Calibration).

Optionally, a lower volume of the Working Reagent may be prepared by using the volume proportion 3:2 of the Reagent 1 and Reagent 2, respectively.

Calibrator. Dissolve the Calibrator bottle (nº 3) contents in 2.0 mL water reagent. Wait 30 minutes. Mix by inversion. Homogenize before using. It is stable 60 days at 2 - 8 °C and 6 months at -10 °C.

Manual procedure

This procedure is not applied to automated and semi-automated equipments only with flow cuvette system. It is available application procedures to automated and semi-automated equipments.

See notes 1, 2 and 3.

Set up two tubes and proceed as follows:

	Unknown	Calibrator
Working Reagent	1.0 mL	1.0 mL

Incubate 2 minutes at 37 °C.

Sample	0.050 mL	
Calibrator		0.050 mL

Mix and incubate exactly 10 minutes in a water bath at 37 $^{\circ}$ C. Measure the absorbance (A₁) of the Unknown and Calibrator against water at 530 nm (510 - 550). Continue the reaction for more exactly 5 minutes and determine the absorbance (A₂) of the Unknown and Calibrator against water at 530 nm (510 - 550).

Calibration . The concentration is traceable to a proteic Calibrator calibrated with glycated polylysine sample with glucose ¹⁴C labeled.

Manual calibrations

Two points calibration.
Calibrator included (Ref.: 97.3).

Calibration frequency

Perform a new calibration weekly and in the following situations:

When the internal quality control indicates.

After reagent lot change.

When a new Working Reagent is used.

Automatic Systems

Blank of reagents: water or 0.85% NaCl (150 mmol/L);

Two points calibration

Calibrator included (Ref. 97.3).

Calibration frequency

Perform a new calibration weekly and in the following situations:

When the internal quality control indicates.

After reagent lot change.

When a new Working Reagent is used.

Quality control. For quality control use Qualitrol H Level 1 and Qualitrol H Level 2 or other suitable control material. The limits and control interval must be adapted to the laboratory requirements. Each laboratory should establish corrective measures to be taken if values fall outside the control limits

Calculations

Determine the difference of the Calibrator and Unknown absorbance: $\Delta A = A_2 \cdot A_1$.

Fructosamine (μ mol/L) = (Δ A Unknown/DA Calibrator) x Calibrator concentration

Due the great reproductive results of the assay system, it is possible to use the factor method:

Calibration factor = Calibrator concentration/ $\triangle A$ Calibrator

Fructosamine (μ mol/L) = Δ A Unkown x Factor



Measurement/reportable range

The measurement result is linear between 20 µmol/L and 800 µmol/L.

If fructosamine concentration exceeds $800\,\mu\text{mol/L}$, the sample must be diluted with 0.85% NaCl. Multiply the result by the appropriate dilution factor.

Expected range. Each laboratory should evaluate the transferability of the expected values to its own patient population and, if necessary, estimate its own reference interval.

For non-diabetic individuals (all ages): 205 to 285 μ mol/L³.

Performance characteristics8

Recovery Studies . In two samples with fructosamine concentrations of 175 and 253 μ mol/L were added known quantities of fructosamine. Subsequent analyses provided recoveries around 102 %. The mean proportional systematic error was 3.5 μ mol/L or 2.0 %.

Method Comparison . A group of 40 sera were assayed by the proposed method and similar technique. Serum fructosamine values ranged from 33 - 775 μ mol/L. The comparisons yielded a correlation coefficient of 0.998 and regression equation was y = 1.029x - 1.346.

Imprecision - Within Run

	N	Mean (μmol/L)	SD (µmol/L)	(%) CV
Sample 1	20	302	2.51	0.83
Sample 2	20	388	3.24	0.84

Imprecision - Run-to-Run

	N	Mean (μmol/L)	SD (µmol/L)	(%) CV
Sample 1	20	302	7.58	2.50
Sample 2	20	388	8.21	2.12

Analytical sensitivity. Detection limit: $5~\mu$ mol/L. The detection limit represents the lowest measurable fructosamine concentration that can be distinguished from zero. It is calculated as two standard deviations of 20 replicates of one sample without fructosamine. It was verified that the photometric detection limit (1 cm light path cuvette) was $3.2~\mu$ mol/L, what corresponds to an absorbance equal to 0.001.

Effects of matrix dilution. A sample with a value equal to $658 \mu mol/L$ was used to evaluate the system response to the matrix dilutions with 150 mmol/L NaCl (0.85%). Using a variety of dilution factors an average recovery of 101% was found.

Notes

- **1.** The material cleaning and drying are fundamental factors to the reagent stability and to obtain correct results.
- 2. The deionized or distilled water in the laboratory to prepare reagents, use in the measurements and for final glass washing must have resistivity ≥1 megaohm.cm, or conductivity ≤1 microsiems/cm and silicates concentration must be <0.1mg/L.
- **3.** It is suggested to consult "www.fxol.org" In order to review physiopathological source and drugs interference in results and methodology.

References

- 1. Baker JR, Metcalf PA, Johnson RN, Newman D, Rietz P. Clin Chem 31:1550-1554. 1985.
- Johnson RN, Metcalf PA, Baker JR, Clin Chem Acta 127:87-95, 1983.
- 3. Kruse-Jarres JD, Jarausch J, Lehmann P, Vogt BW, Rietz P. Lab Med 13:245-253, 1989.
- 4. Lim YS, Staley MJ. Clin Chem 32:403-404, 1986.
- 5. Lloyd D, Marples J. Clin Chem 30:1686-1688, 1984.
- 6. Schleicher ED, Vogt BW. Clin Chem 36:136-139, 1990.
- Westgard JO, Barry PL, Hunt MR, Groth T. Clin Chem. 1981, 27:493-501.
- Sociedad Española de Bioquímica Clínica y Patología Molecular, Base de Datos de Variación Biológica. Disponível em:http://www.seqc.es/article/articleview/330/1/170 (acesso em 04/2006).
- Basques JC. Especificações da Qualidade Analítica. Labtest Diagnóstica 2005.
- 10. Labtest: data on file.

Presentation

Product	Reference	Contents	
Fructosamine	97-6/15	R1 6 X 9 mL	
		R 2 6 X 6 mL	
		CAL 1 X 2 mL	
Function		R 1 4 X 9 mL	
Fructosamine Labmax 560/400	97-4/15	R 2 4 X 6 mL	
		CAL 1 X 2 mL	

^{*} The number of tests in automated application procedures depends on the programmed parameters.



Application procedures using fructosamine are available for various automated instruments.

Customer information

[Warranty conditions]

Labtest Diagnóstica warrants the performance of this product under the specifications until the expiration date shown in the label since the application procedures and storage conditions, indicated on the label and in this insert, have been followed correctly.

CE

Labtest Diagnóstica S.A.

CNPJ: 16.516.296 / 0001 - 38

Av. Paulo Ferreira da Costa, 600 - Vista Alegre - CEP 33400-000 Lagoa Santa . Minas Gerais Brasil - **www.labtest.com.br**

Customer Service e-mail: customerservice@labtest.com.br

Revision: December, 2014 Ref.: 260117 Copyright by Labtest Diagnóstica S.A. Reproduction under previous autorization



Símbolos utilizados com produtos diagnósticos in vitro

Símbolos usados con productos diagnósticos in vitro Symbols used with ivd devices

$\sum_{}$	Conteúdo suficiente para < n > testes Contenido suficiente para < n > tests Contains sufficient for < n > tests	₩	Risco biológico Riesgo biológico Biological risk
	Data limite de utilização (aaaa-mm-dd ou mm/aaaa) Estable hasta (aaaa-mm-dd o mm/aaaa) Use by (yyyy-mm-dd or mm/yyyy)	C€	Marca CE Marcado CE CE Mark
CAL	Material Calibrador Material Calibrador Calibrator Material	₩	Tóxico Tóxico Poison
CAL	Material Calibrador Material Calibrador Calibrator Material	R	Reagente Reactivo Reagent
-	Limite de temperatura (conservar a) Temperatura limite (conservar a) Temperature limitation (store at)		Fabricado por Elaborado por Manufactured by
EC REP	Representante Autorizado na Comunidade Europeia Representante autorizado en la Comunidad Europea Authorized Representative in the European Community	LOT	Número do lote Denominación de lote Batch code
Ţį	Consultar instruções de uso Consultar instrucciones de uso Consult instructions for use	CONTROL	Controle Control Control
REF	Número do catálogo Número de catálogo Catalog Number	CONTROL -	Controle negativo Control negativo Negative control
	Adições ou alterações significativas Cambios o suplementos significativos Significant additions or changes	CONTROL +	Controle positivo Control positivo Positive control
IVD	Produto diagnóstico in vitro Dispositivo de diagnóstico in vitro In vitro diagnostic device	CONTROL	Controle Control Control
LYOPH	Liofilizado Liofilizado Lyophilized		Corrosivo Corrosivo Corrosive
	Período após abertura Período post-abertura Period after-opening	③	Uso veterinário Uso veterinario Veterinary use
N	Instalar até Instalar hasta Instala before		Ref.: 140214

