Argutus Medical Urinary Collagen IV EIA – BIO83 – Instructions for use





# **Urinary Collagen IV EIA**

# **Enzyme Immunoassay**

**Instructions for Use** 

FOR RESEARCH USE ONLY Not for use in Diagnostic Procedures

Manufactured by: Daiichi Fine Chemical Co., Ltd. Japan

# TABLE OF CONTENTS

INTENDED USE	3
BACKGROUND	3
ASSAY PRINCIPLE	4
COMPONENTS	4
PRECAUTIONS	5
STABILITY AND STORAGE	6
ADDITIONAL MATERIALS REQUIRED	6
PREPARATION OF REAGENTS	7
SAMPLE COLLECTION AND STORAGE	7
ASSAY PROCEDURE	8
CALCULATION OF RESULTS	8
LIMITATIONS OF USE	8
PERFORMANCE CHARACTERISTICS	9
EXAMPLE OF CALIBRATION CURVE	11
WARRANTY	11
OTHER ARGUTUS MEDICAL ASSAYS	12
REFERENCES	12

# INTENDED USE

The Argutus Medical Urinary Collagen IV EIA provides a method for the quantitative determination of collagen IV in human urine. Please contact Argutus Medical for further information regarding the assay of collagen IV in other tissue fluids. The Argutus Medical Urinary Collagen IV EIA is for research use only. Not for use in diagnostic procedures.

# BACKGROUND

Type IV collagen ( $IV \bullet C$ ), is a major component of the basement membrane (BM), and it is considered to constitute its basic framework. Urinary collagen IV levels are elevated in a variety of renal pathologies<sup>1</sup>, particularly diabetic nephropathy<sup>2-8</sup>. Urinary collagen IV is significantly higher in patients with non-insulin dependent diabetic mellitus (NIDDM) than in normal subjects and urinary collagen IV levels correlate with the deposition of collagen IV in the kidney<sup>2</sup>. In diabetic subjects, urinary collagen IV is significantly increased in patients with microalbuminuria or overt proteinuria as compared to those with normoalbuminuria<sup>3</sup> (Figure 1). Moreover, in diabetics with normoalbuminuria, those with elevated urinary collagen IV were at an increased risk for progression to microalbuminuria<sup>4</sup>. Intensive therapy of diabetic nephropathy can slow the temporal increase in urinary collagen IV in diabetics indicating the potential of urinary collagen IV for studying the renal effects of new therapies<sup>5</sup>. These results suggest that the measurement of UIV•C might provide a useful biomarker for studying diabetic nephropathy.

In the field of renal transplantation, renal collagen IV levels are increased<sup>9</sup> and increased urinary collagen IV levels are found in acute renal rejection<sup>10</sup>, indicating the potential value of urinary collagen IV in studying these conditions.



<sup>#</sup> p<0.05 vs Micro.

# <u>ASSAY PRINCIPLE</u>

The Argutus Medical Urinary Collagen IV EIA is designed for the assay of urinary collagen IV. It is a solid phase one-step sandwich EIA. Collagen IV in the sample is bound simultaneously by a solid phase monoclonal antibody and a monoclonal antibody-enzyme conjugate directed at different antigenic sites. This results in the collagen IV molecule being sandwiched between the solid phase and enzyme labelled antibodies. After removing unbound enzyme labelled antibody and sample, the plate is then incubated with enzyme substrate, resulting in the development of a colour. The colour developed is directly proportional to the amount of collagen IV in the sample.

# **COMPONENTS**

- 1. Antibody coated Microassay plate: 12x8 well strips coated with IgG directed against human collagen IV. **READY TO USE**
- 2. Collagen IV Calibrators: Purified collagen IV in CAL 0.8 1X phosphate buffer (pH 6.0) with Bovine Serum Albumin (BSA). CAL 3.2 1X 0.8, 3.2, 12.5 and 50 µg/L. (1 mL each). Contains 30 mg/L Proclin 300 as preservative. CAL 12.5 1X **READY TO USE** CAL 50
  - BUF DIL 3. Assay Buffer: Phosphate buffer (pH 7.0) 1X containing bovine serum albumin and horse serum (10 mL). Contains 30 mg/L Proclin 300 preservative. **READY TO USE**
  - 4. Conjugate: Anti-collagen IV mouse Fab' conjugated to horseradish peroxidase (20 mL). Contains 30 mg/L Proclin 300 as preservative. **READY TO USE**
  - 5. Wash Concentrate: 10x Conc. Phosphate buffer BUF WASH 10X with Tween 20 (PBT), (2 bottles of 50 mL). Contain 30 mg/L Proclin 300 as preservative. CONCENTRATE
  - 6. Substrate: Stabilised liquid TMB solution (15 mL) SUBS ТМВ 1X **READY TO USE**
  - 7. Stop Solution: 1M Sulphuric Acid (15 mL). **READY TO USE**
  - 8. Plate Seal: 1 sheet
  - 9. Instructions for use

PLA

1X

1X

1X

INS

CONJ

STP

SOLN

# SUPPLIED IN A SEPARATE PACKAGE

Argutus Medical Urinary Collagen IV EIA Sample Collecting Tubes coated with urinary collagen IV stabiliser (45 tubes)

# PRECAUTIONS

## SAFETY

- The Argutus Medical Urinary Collagen IV EIA kit is intended for use by qualified laboratory staff only.
- The kit contains material of human origin that has been tested and found to be negative for Hepatitis B surface antigen, Hepatitis C and HIV antibodies. However, since no test can provide complete assurance, treat all materials as potentially infectious.
- The Stop Solution contains sulphuric acid which is corrosive. Avoid contact with the skin and eyes. If contact occurs, rinse off immediately with water and seek medical advice.
- The Substrate contains TMB that may irritate the skin and mucous membranes. Any substrate which comes in contact with the skin should be rinsed off with water.
- Dispose of all clinical specimens, infected or potentially infected material in accordance with good laboratory practice. All such materials should be handled and disposed of as though potentially infectious.
- Residues of chemicals and kit components are generally considered as hazardous waste. All such materials should be disposed of in accordance with established safety procedures.
- Wear protective clothing, disposable latex gloves and eye protection while handling specimens and performing the assay. Wash hands thoroughly when finished.
- Do not pipette materials by the mouth and never eat or drink at the laboratory workbench.

# PROCEDURAL

- Do not use kit or individual reagents past their expiry date.
- Do not mix or substitute reagents from different kit lot numbers.
- Deviation from the protocol provided may cause erroneous results.
- Performing the assay outside the time and temperature ranges specified may produce invalid results. Assays not falling within the established time and temperature ranges must be repeated.
- Reagent delivery should be aimed at midpoint of the side of the wells, taking care not to scratch the side with the pipette tip.
- Do not allow the wells to dry at any stage during the assay procedure.
- Care must be taken not to contaminate components and always use fresh pipette tips for each sample and component.
- Do not use reagents that are cloudy or that have precipitated out of solution.

- Ensure Wash Concentrate is mixed thoroughly and no crystals remain before reconstitution.
- High quality distilled or deionised water is required for the Wash Solution. The use of poor quality or contaminated water may lead to background colour in the assay.
- Allow all reagents to come to room temperature (20-27°C) and mix well prior to use.
- Avoid leaving reagents in direct sunlight and/or above 2-8°C for extended periods.
- Always use clean, preferably disposable, glassware for all reagent preparation.
- Ensure that the bottom surface of the plate is clean and dry before reading.
- Before commencing the assay, an identification and distribution plan should be established.

# STABILITY AND STORAGE

- 1. All kit reagents should be stored at 2-8°C and are stable as supplied until the expiry date shown.
- 2. Prepared Wash Solution (PBT) is stable for up to one month at 2-8°C.
- 3. Plate assay wells should be stored in sealed bags with desiccants at 2-8°C until required for use. Return unused wells to the storage bag together with desiccant.
- 4. Argutus Medical Urinary Collagen IV Sample Collecting Tubes are stable at room temperature until the expiry date shown. Return unused tubes to the storage bag.

# ADDITIONAL MATERIALS REQUIRED

- 1. Pasteur pipette
- 2. 50 µL Micropipette and a 100-150 µL multichannel pipette
- 3. Microassay strip washing system
- 4. ELISA plate reader capable of measuring at 450nm with reference at 630nm if available
- 5. 1 L beaker
- 6. Timer
- 7. Liquid trough
- 8. Deionised/Distilled water
- 9. Graduated cylinder (500mL)

# PREPARATION OF REAGENTS

#### WASH SOLUTION (PBT)

Perform a 1/10 dilution of Wash Concentrate adding, for example, 10mL Wash Concentrate to 90 mL deionised water as required. Prepare only the volume of Wash Solution required for the assay. Each row of assay wells requires 15 mL of Washing Solution.

#### Ensure salt crystals are dissolved prior to dilution.

Gentle warming of Wash Concentrate at 37°C for 30 minutes will aid dissolution of salt crystals.

# SAMPLE COLLECTION AND STORAGE

Collagen IV precipitates out of urine upon standing leading to falsely low results. This can be prevented by the addition of a stabilising buffer to the urine after which the urine can be stored<sup>11</sup>. The Argutus Medical Urinary Collagen IV Sample Collecting Tubes provide a simple and reproducible means of collecting urine samples for the assay of collagen IV.

Transfer urine to the collection tube using a Pasteur pipette. Fill the tube to the line indicated, then mix thoroughly. As collagen IV is absorbed by urinary precipitates that form during storage, collect fresh urine in the Argutus Medical Urinary Collagen IV EIA Urine Collection Tubes. Urine samples must be transferred to the collection tubes on the day of collection. Samples must be transferred to the collection tubes, even if they are not to be stored. To facilitate compensation for dieresis, it is recommended that a simultaneous sample be taken for urinary creatinine.

After addition to the urine tubes, samples can be stored at  $2-8^{\circ}$ C for one week or nine months at  $-20^{\circ}$ C. If samples have been frozen, it is essential to mix thoroughly to dissolve any precipitates. Repeated freeze thawing of samples should be avoided.

# ASSAY PROCEDURE

**NOTE:** All reagents should be allowed to reach room temperature prior to commencement of assay.

#### 1. IMMUNOREACTION

- 1.1 Prepare Wash Solution as described in "Preparation of Reagents".
- 1.2 Place required number of Microassay wells in the assay plate (10 for the Calibrators plus two for each sample).
- 1.3 Add **150 µL** Conjugate to each well using a multichannel pipette.
- 1.4 Add Calibrators (0 (Assay buffer), 0.8, 3.2, 12.5 and 50 μg/L) and samples (50 μL/well), in duplicate, to the Microassay plate.
- 1.5 Cover the Microassay plate with the plate seal and incubate at room temperature (20-27°C) for 24 hours.
- 1.6 Remove plate seal cover and wash each strip five times **(350 μL/well)** with Wash Solution. When complete, firmly tap the plate against a paper towel to ensure complete removal of wash fluid from wells.

# 2. COLOUR DEVELOPMENT

2.1 Add **100µL** Substrate/well using a multichannel pipette and incubate at room temperature (20-27°C) for exactly one hour.

# 3. STOP

- 3.1 Add **100 µL** Stop Solution/well using a multichannel pipette. Ensure complete mixing of Substrate and Stop Solution.
- 3.2 Read **immediately** at 450nm using 630nm as reference (if available).

# **CALCULATION OF RESULTS**

- 1. Calculate the mean absorbance for each calibrator and sample.
- 2. Plot a Calibration curve of  $A_{450/630nm}$  versus collagen IV (µg/L) on a log-log scale.
- 3. Read the collagen IV (µg/L) indicated by the mean absorbances of the samples from the calibration curve.
- 4. If the samples have been diluted, multiply the calculated [collagen IV] by the appropriate dilution factor in order to obtain the actual [collagen IV].

# PERFORMANCE CHARACTERISTICS

#### NORMAL RANGE

Based on healthy Japanese volunteers, 95% confidence limits for urinary collagen IV are:

		Collagen IV µg/g creatinine	Ν
Early morning urine	30-39 years	<4.0	122
	>40 years	<4.9	64
Spot urine	<21 years	<7.3	390

#### LIMIT OF DETECTION

The detection limit of Argutus Medical Urinary Collagen IV EIA is 0.8 µg/L.

#### MEASURING RANGE

The calibration curve range covers the range 0.8-50  $\mu$ g/L. This range may be extended by increasing sample dilution.

#### SPECIFICITY

The Argutus Medical Urinary Collagen EIA IV is highly specific for the detection of collagen IV. Cross reactivity is less than 2% with Collagen II and less than 0.5% with other forms of collagen.

#### SENSITIVITY

When reading from the standard curve the  $A_{450nm}$  value of the 0.8 µg/L standard should be 0.01 – 0.06, and the 50 µg/L standard should be >0.8.

#### INTERFERENCE

No significant interference has been observed in this assay with creatinine, haemolytic or icteric samples.

- Creatinine: Less than 10% interference up to 3 g/L in sample.
- Haemolysis: Less than 10% interference up to 4.8 g/L haemoglobin.
- Icteris: Less than 10% interference up to 0.2 g/L bilirubin.

# **DILUTION – RECOVERY**

Sample	Dilution								
	1/2			1 / 4		1 / 8			
	Expected	Obtained	Recovery	Expected	Obtained	Recovery	Expected	Obtained	Recovery
	µg/L	µg/L	%	µg/L	µg/L	%	µg/L	µg/L	%
А	11.6	11.0	95	5.8	5.2	90	2.9	2.6	90
В	13.4	12.7	95	6.7	6.4	96	3.3	3.1	94
С	5.1	5.4	106	2.6	2.5	96	1.3	1.3	100

Dilution of samples containing high levels of collagen IV gave the following results:

#### REPRODUCIBILITY

#### Intra-assay variation of the Argutus Medical Urinary Collagen IV EIA

Sample	¯χ [Collagen IV] μg/L	SD	% CV	N
Low	2.5	0.07	2.8	8
Medium	6.2	0.13	2.1	8
High	10.8	0.29	2.7	8

#### Inter-assay variation of the Argutus Medical Urinary Collagen IV EIA

Sample	 [Collagen IV] μg/L	SD	%CV	Ν
Low	2.4	0.05	2.1	4
Medium	6.0	0.15	2.0	4
High	20.8	1.52	7.3	4

#### Inter-batch Variation of the Argutus Medical Urinary Collagen IV EIA calculated for four batches of kits

Sample	¯χ [Collagen IV] μg/L	SD	%CV	Ν
Low	3.2	0.18	5.6	4
Medium	5.8	0.11	1.9	4
High	17.0	0.52	3.1	4

# **EXAMPLE OF A CALIBRATION CURVE**



<u>Figure 2</u>: Typical Calibration curve obtained using the Argutus Medical Urinary Collagen IV EIA. Plot of A450<sub>/630nm</sub> versus [Collagen IV] µg/L. Assay range is 0.8 – 50 µg/L

# WARRANTY

The performance data presented here was obtained using the procedure described. Any change or modification of the procedure, not recommended by Argutus Medical, may affect the results, in which case Argutus Medical disclaims all warranties, expressed, implied or statutory, including implied merchantability and fitness for use. In the case of such an event, Argutus Medical shall not be liable for damages, direct or consequential.

# **OTHER ARGUTUS MEDICAL ASSAYS**

# Pancreatic Injury Testing Service

Catalogue No	Product Name	Description
TEST BBU	Trypsinogen Activation Peptide	TAP in human and mammalian urine and
	(TAP) EIA	tissue

#### **Animal Organ Damage Biomarkers**

Catalogue No	Product Name	Description
BIO64RT	Rat Alpha GST EIA	$\alpha$ GST in rat serum, urine and tissue culture
BIO76YB1	Rat Yb1 GST EIA	GSTYb1 (µGST) in rat urine
BIO89RPA1	RPA-1 EIA	Renal papillary antigen 1 in rat urine
BIO87CD	RPA-1 Antibody	Antibody to rat collecting duct
BIO88LH	RPA-2 Antibody	Antibody to rat loop of henle

#### Human Organ Damage Biomarkers

Catalogue No	Product Name	Description
BIO66NEPHA	NEPHKIT® Alpha GST EIA	αGST in human urine
BIO60HEPA	HEPKIT® Alpha GST EIA	$\alpha$ GST in human serum and plasma
BIO60HEPAS	High Sensitivity Alpha GST EIA	$\alpha$ GST in human serum and plasma
BIO85	PI GST EIA	$\pi$ GST in human urine and plasma
BIO82	Serum Collagen IV EIA	Collagen IV in human serum
BIO81DNA	OxyDNA test	Fluorescence method for the detection of
		oxidative DNA damage in cell suspensions

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Unit 9 Trinity Technology & Enterprise Campus, Pearse Street, Dublin 2, Ireland Tel: +353 1 670 8576 Fax: +353 1 670 8575 <u>info@argutusmed.com</u> <u>http://www.argutusmed.com</u>

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