**INTENDED USE**

For the quantitative determination of human alpha-1 microglobulin in human, urine, serum, and plasma by immunoturbidimetric assay. FOR IN VITRO DIAGNOSTIC USE.

**INTRODUCTION AND SUMMARY**

Alpha-1 microglobulin is a low molecular weight glycoprotein of 24 - 31 kDa that was initially isolated from the urine of patients with renal tubular disorders in 1975. It is mainly synthesized in the liver and is widely distributed in various body fluids. The measurement of alpha-1 microglobulin in serum and urine has been considered to be useful for the diagnosis of functional renal disorders and the assessment of the progress and the prognosis of diseases. The K-ASSAY® Alpha-1 Microglobulin assay is intended for the quantitative determination of human alpha-1 microglobulin in urine, serum, and plasma by immunoturbidimetric assay.

**PRINCIPLE OF TEST**

When an antigen-antibody reaction occurs between alpha-1 microglobulin in a sample and alpha-1 microglobulin antibody that has been adsorbed to latex particles, agglutination results. This agglutination is detected as an absorbance change, with the magnitude of the change being proportional to the quantity of alpha-1 microglobulin in the sample. The actual concentration is then determined by interpolation from a calibration curve prepared from calibrators of known concentration.

**KIT COMPOSITION**

Reagents (Liquid Stable)

- R1: Buffer Reagent 1 x 50 mL
- R2: Latex Reagent 1 x 50 mL
  0.25 w/v% suspension of latex particles sensitized with anti-human alpha-1 microglobulin antibodies (rabbit).

**WARNINGS AND PRECAUTIONS**

FOR IN VITRO DIAGNOSTIC USE. Rx only.

Not to be used internally in humans or animals. Normal precautions exercised in handling laboratory reagents should be followed. Do not mix or use reagents from one test kit with those from a different lot number. Do not use reagents past their expiration date stated on each reagent container label.

Do not pipette by mouth. Avoid ingestion and contact with skin.

Reagents in this kit contain sodium azide as a preservative. Sodium azide may form explosive compounds in metal drain lines. When disposing of reagents through plumbing fixtures, flush with copious amounts of water.

**REAGENT PREPARATION**

Reagents are ready to use and do not require reconstitution.

To prepare physiological saline, dissolve 0.9 g sodium chloride in distilled water and bring to a final volume of 100 mL.

**STORAGE AND HANDLING**

All reagents should be stored refrigerated (2-8°C). Return all reagents to 2-8°C promptly after use. The reagents can be used until the expiration date.

**REAGENT STABILITY**

Discard reagents if they become contaminated. Evidence of obvious precipitation in reagent 2 (R-2) solution is cause to discard. Opened reagents can be used for 1 month if stored at 2-8°C. It is recommended that R-2 be gently inverted once a week after it is opened.

**SPECIMEN COLLECTION AND PREPARATION**

The sample may be urine, serum or plasma (depending on calibrator used).

It is recommended that specimen collection be carried out in accordance with NCCCLS document M29-A2 and H2-A2. No method can offer complete assurance that human blood samples will not transmit infection. Therefore, all blood samples should be considered potentially infectious.

Testing should be performed immediately. If this is not possible, the sample should be placed in a tightly sealed container and stored at 20°C. Avoid freeze/thaw cycles.

**AUTOMATED ANALYZER APPLICATION**

Suitable for two-reagent clinical chemistry analyzers that can measure a rate reaction at an absorbance of 570 nm primary and 800 nm secondary. Refer to the instrument manual from the manufacturer regarding the following:

a) Use or function
b) Installation procedures and requirements
c) Principles of operation
d) Performance characteristics, operating instructions
e) Installation procedures including materials and / or equipment to be used
f) Service and maintenance information

**PROCEDURE**

**Materials Supplied**

- KAI-056 Reagent 1 (R-1) Buffer Reagent 1 x 50 mL
- KAI-056 Reagent 2 (R-2) Antiserum Reagent 1 x 50 mL

**Calibrators**

**Urine samples:**

Urine Alpha-1 Microglobulin Calibrator, Cat. No. KAI-067C

**Serum or plasma samples:**

Serum / Plasma Alpha-1 Microglobulin Calibrator, Cat. No. KAI-066C

Clinical chemistry analyzer capable of accurately reading at 570 and 800 nm, accurately dispensing the required volumes, and maintaining 37°C.

**ASSAY PROCEDURE**

An example of automated applications (Hitachi 717):

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
<th>Temperature</th>
<th>Absorbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>urine sample</td>
<td>7 µL</td>
<td>37°C, 5 min.</td>
<td>570/800 nm</td>
</tr>
<tr>
<td>serum sample</td>
<td>3 µL</td>
<td>37°C, 5 min.</td>
<td>570 nm</td>
</tr>
<tr>
<td>2-point endpoint</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**STORAGE AND HANDLING**

Room temperature. Mix all reagents gently before using.

**Chemistry Parameters for Automatic Analyzer:**

**Institution**

Hitachi 717

**Temperature**

37°C

**Test**

A1IMG

**Sample Volume**

(2 POINT): (28): (-50)

**R1 Volume**

(125): (100): (NO)

**R2 Volume**

(125): (100): (NO)

**Wavelength**

800: (570)

**Calib. Method**

NONLINEAR (4) / (5)

**STD(1) CONC.**

0.00: (1)

**STD(2) CONC.**

0.2: (2)

**STD(3) CONC.**

0.3: (3)

**STD(4) CONC.**

0.4: (4)

**STD(5) CONC.**

0.5: (5)

**STD(6) CONC.**

999

**SD Limit**

10000

**DUPLICATE LIMIT**

Sensitivity Unit (0)

ABS LIMIT (INC/DEC) (-32000) / (INC/DEC)

**PROZONE LIMIT**

32000 / (LOWER)

**EXPECTED VALUE**

99999 / (99999)

**PANIC VALUE**

99999 / (99999)

**INSTRUMENT FACTOR**

1.0

Use isotonic saline as STD (1)

* 2-5 Input concentration of calibrators

Parameters for other automated analyzers are available.
It is recommended that a multi-point calibration curve be made using either the K-ASSAY® Serum / Plasma Alpha-1 Microglobulin Calibrator or K-ASSAY® Urine Alpha-1 Microglobulin Calibrator. Please be sure to use the proper instrument application for the calibrator you are using. It is recommended that the user determine calibration frequency as this will depend on the instrument and type/number of other assays being run. Initially, calibration should be performed each day.

QUALITY CONTROL

Normal and abnormal controls of known concentration should be included with every assay performed. The value determined for the controls should fall within the stated limits of the values assigned to the controls. The validity of the assay is in question if the values for the controls generated by the assay's calibration curve do not fall within this range. Recalibrate if the values determined for the controls fall outside the stated range.

CALCULATIONS

Alpha-1 microglobulin levels are determined by the analyzer using the prepared calibration curve.

LIMITATIONS OF PROCEDURE

The Alpha-1 microglobulin test is suitable for measuring in the range of:

- Serum and Plasma: 1.0 - 137.0 mg/L
- Urine: 0.2 - 34.0 mg/L

Reagents should not be used after the expiration date indicated on the kit label. Do not mix reagents with different lot numbers.

If the alpha-1 microglobulin concentration is greater than highest calibrator value, dilute one part sample with four parts isotonic saline and re-assay. Multiply results by 5 to compensate for the dilution.

PERFORMANCE

Sensitivity

Serum and Plasma: When a calibrator containing 17.1 mg/L alpha-1 microglobulin is measured, the range of the absorbance change per minute at a wavelength of 572 nm is 0.016 - 0.030.

Urine: When a calibrator containing 3.4 mg/L alpha-1 microglobulin is measured, the range of the absorbance change per minute at a wavelength of 572 nm is 0.010 - 0.020.

Specificity

When a sample containing a known level of alpha-1 microglobulin was measured, the value obtained for the sample was in the range of the known concentration, ± 10%.

Precise Assay:

When a sample containing a known level of alpha-1 microglobulin is tested 10 times, the CV of the test should typically be under 10%.

Precision Assay:

<table>
<thead>
<tr>
<th>Sample</th>
<th>Serum</th>
<th>Sample</th>
<th>Serum</th>
<th>Sample</th>
<th>Serum</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>N = 10</td>
<td>II</td>
<td>N = 10</td>
<td>III</td>
<td>N = 10</td>
</tr>
<tr>
<td>Mean</td>
<td>7.1 mg/L</td>
<td>Mean</td>
<td>32.1 mg/L</td>
<td>Mean</td>
<td>66.1 mg/L</td>
</tr>
<tr>
<td>SD</td>
<td>0.145</td>
<td>SD</td>
<td>0.263</td>
<td>SD</td>
<td>0.540</td>
</tr>
<tr>
<td>CV</td>
<td>2.04%</td>
<td>CV</td>
<td>0.82%</td>
<td>CV</td>
<td>0.82%</td>
</tr>
</tbody>
</table>

Accuracy / Correlation

Correlation between this test and the latex agglutination method of another company is given below:

- Serum samples:
  - y = 1.00x + 2.83
  - r = 1.00
  - n = 55
  - x = Company A's latex reagent
  - y = K-ASSAY® Alpha-1 Microglobulin

- Urine samples:
  - y = 1.00x - 0.52
  - r = 1.00
  - n = 55
  - x = Company A's latex reagent
  - y = K-ASSAY® Alpha-1 Microglobulin

Assay Range

- Serum / Plasma: 1.0 to 137.0 mg/L (or value of highest calibration point)
- Urine: 0.2 to 34.0 mg/L (or value of highest calibration point)

Lower Limit of Detection

- Serum / Plasma: 1.0 mg/L
- Urine: 0.2 mg/L

INTERFERENCE

Serum and Urine Samples:

- Hemoglobin: No interference up to 500 mg/dL
- Bilirubin C: No interference up to 30 mg/dL
- Bilirubin F: No interference up to 30 mg/dL
- Lipid (Intrafat): No interference up to 5%.

Dust particles or other particulate matter in the reaction solution may result in extraneous light scattering, which may affect the accuracy of the test.

EXPECTED VALUES

It is recommended that each laboratory establish its own expected range to reflect its patient population.

- Serum (Within Run):
  - Sample I: N = 10, Mean = 7.1 mg/L, SD = 0.145, CV = 2.04%
  - Sample II: N = 10, Mean = 32.1 mg/L, SD = 0.263, CV = 0.82%
  - Sample III: N = 10, Mean = 66.1 mg/L, SD = 0.540, CV = 0.82%

- Urine (Within Run):
  - Sample I: N = 10, Mean = 0.7 mg/L, SD = 0.029, CV = 3.93%
  - Sample II: N = 10, Mean = 3.3 mg/L, SD = 0.012, CV = 0.37%
  - Sample III: N = 10, Mean = 17.4 mg/L, SD = 0.059, CV = 0.34%

- Serum (Between Runs):
  - Sample I: N = 15, Mean = 7.5 mg/L, SD = 0.046, CV = 6.3%
  - Sample II: N = 15, Mean = 32.3 mg/L, SD = 0.04, CV = 1.2%
  - Sample III: N = 15, Mean = 67.1 mg/L, SD = 0.31, CV = 1.8%

- Urine (Between Runs):
  - Sample I: N = 15, Mean = 0.72 mg/L, SD = 0.016, CV = 2.0%
  - Sample II: N = 15, Mean = 3.3 mg/L, SD = 0.04, CV = 1.2%
  - Sample III: N = 15, Mean = 17.2 mg/L, SD = 0.54, CV = 3.1%

LIMITATIONS OF PROCEDURE

The correlation between this test and the latex agglutination method of another company is given below:

- Serum samples:
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  - n = 55
  - x = Company A's latex reagent
  - y = K-ASSAY® Alpha-1 Microglobulin

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  - n = 55
  - x = Company A's latex reagent
  - y = K-ASSAY® Alpha-1 Microglobulin

ASSAY RANGE

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EXPECTED VALUES

It is recommended that each laboratory establish its own expected range to reflect its patient population.

- Serum / Plasma:
  - 10.0 – 30.0 mg/L
- Urine:
  - 1.0 – 5.0 mg/L

Expected values were taken from the literature: Ito, Y., "Alpha-1 Microglobulin (protein HC)." Nippon-rinsho, 47:176, 1989.

LABELING SYMBOLS

- LOT: Lot Number
- ST: Reagent
- T: Expiration or 'Use By' Date
- REF: Catalog Number
- DV: For In Vitro Diagnostic Use
- 2-8°C: Temperature Limitation. Store between 2 and 8 degrees C
- H: Potential Human Biohazard
- M: Manufacturer
- P: Consulit Package Insert for Instructions for Use
- ECREP: Authorized Representative in the European Community

EU AUTHORIZED REPRESENTATIVE

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