ADMA (human) ELISA Kit

Manufactured by Immundiagnostik AG.

ALX-850-323-KI01

96 wells (~80 tests)

(Version 05: December 9, 2009)

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For laboratory use only. Not for human or diagnostic use.
ADMA ELISA Kit

For the determination of ADMA in human serum, citrate- and EDTA-plasma

Valid from 09.12.2009

REF K 7828

Valid from 09.12.2009
1. INTENDED USE

The ADMA ELISA Kit is intended for the quantitative determination of asymmetric dimethylarginine (ADMA) in human serum, citrate- and EDTA-plasma. It is for in vitro diagnostic use only.

2. INTRODUCTION

Asymmetric dimethylarginine (ADMA) is an endogenous inhibitor of NO-synthase. It is formed during proteolysis of methylated proteins and removed by renal excretion or metabolic degradation by the enzyme dimethylarginine dimethylaminohydrolase (DDAH). Several celltypes, including human endothelial and tubular cells are capable of synthesizing and metabolizing ADMA. Elevated ADMA concentrations in the blood are found in numerous diseases associated with endothelial dysfunction. For example, elevated ADMA levels in blood of dialysis patients correlate significantly with the degree of arteriosclerosis and cardiovascular risk. Furthermore, elevated ADMA levels are found in patients with hypercholesterolemia, hypertension, arteriosclerosis, chronic renal failure and chronic heart failure, and are associated with restrictions in endothelial vasodilatation.

During the last years, the important clinical relevance of the regulation of vascular tone and structure by nitric oxide (NO) has been shown. Moreover, there were reports that human endothelial cells produce ADMA as well as nitric oxide, which points to an endogenous endothelial NO-regulation by ADMA. Therefore it was assumed that hypertension, arteriosclerosis and immunological dysfunction in patients with chronic renal failure are connected to a dysfunction of the L-arginin/NO-metabolism and to ADMA accumulation. The reasons for the deregulation of the L-arginin/NO-metabolism could only partially be elucidated. Certainly, there are multiple factors involved in the L-arginin/NO-metabolism regulation as for example elevation of free superoxide radicals (O$_2^{-}$), ADMA accumulation and reduced NO-synthase activity.

Prospective clinical studies of the last years demonstrate the increased importance of ADMA as a novel cardiovascular risk factor.
Indication

- Arteriosclerosis
- Hypertension
- Chronic heart failure
- Coronary artery disease
- Hypercholesterolemia
- Chronic renal failure
- Diabetes mellitus
- Peripheral arterial occlusive disease

3. Principle of the Test

This assay is based on the method of competitive enzyme linked immunoassays. The sample preparation includes the addition of a derivatization-reagent for ADMA coupling. Afterwards, the treated samples and the polyclonal ADMA-antiserum are incubated in wells of microplate coated with ADMA-derivative (tracer). During the incubation period, the target ADMA in the sample competes with the tracer immobilized on the wall of the microtiter wells for the binding of the polyclonal antibodies. The ADMA in the sample displaces the antibodies out of the binding to the tracer. Therefore the concentration of the tracer-bound antibody is inverse proportional to the ADMA concentration in the sample. During the second incubation step, a peroxidase-conjugated antibody is added to each microtiter well to detect the anti-ADMA antibodies. After washing away the unbound components, tetramethylbenzidine (TMB) is added as a substrate for peroxidase. Finally, the enzymatic reaction is terminated by an acidic stop solution. The color changes from blue to yellow and the absorbance is measured in the photometer at 450 nm. The intensity of the yellow color is inverse proportional to the ADMA concentration in the sample; this means high ADMA concentration in the sample reduces the concentration of tracer-bound antibodies and lowers the photometric signal.

A dose response curve of absorbance unit (optical density, OD at 450 nm) vs. concentration is generated using the values obtained from the standard. ADMA present in the patient samples is determined directly from this curve.
## 4. Material Supplied

<table>
<thead>
<tr>
<th>Catalog No</th>
<th>Content</th>
<th>Kit Components</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>K7828MTP</td>
<td>PLATE</td>
<td>One holder with precoated strips</td>
<td>12 x 8 wells</td>
</tr>
<tr>
<td>K7828ST</td>
<td>STD</td>
<td>Standards (diluted in reaction buffer)</td>
<td>6 x 1 vial</td>
</tr>
<tr>
<td>K7828KO</td>
<td>CTRL 1+ CTRL 2</td>
<td>Controls (diluted in reaction buffer)</td>
<td>2 x 1 vial</td>
</tr>
<tr>
<td>K7828WP</td>
<td>WASHBUF</td>
<td>Wash buffer concentrate (10 fold)</td>
<td>2 x 100 ml</td>
</tr>
<tr>
<td>K7828AK</td>
<td>AB</td>
<td>ADMA antibody (lyophilized)</td>
<td>2 x 1 vial</td>
</tr>
<tr>
<td>K7828K</td>
<td>2.AB</td>
<td>POD antibody (concentrate)</td>
<td>120 µl</td>
</tr>
<tr>
<td>K7828CSP</td>
<td>2.ABDIL</td>
<td>Conjugate stabilizing buffer</td>
<td>24 ml</td>
</tr>
<tr>
<td>K7828RP</td>
<td>DERBUF</td>
<td>Reaction buffer</td>
<td>15 ml</td>
</tr>
<tr>
<td>K7828DR</td>
<td>DER</td>
<td>Derivatization reagent</td>
<td>2 x 50 mg</td>
</tr>
<tr>
<td>K7828LM</td>
<td>DMSO</td>
<td>Dimethylsulfoxid (DMSO)</td>
<td>7 ml</td>
</tr>
<tr>
<td>K7828SL</td>
<td>CODIL</td>
<td>Dilution buffer for coupling</td>
<td>28 ml</td>
</tr>
<tr>
<td>K7828TMB</td>
<td>SUB</td>
<td>TMB substrate</td>
<td>25 ml</td>
</tr>
<tr>
<td>K7828AC</td>
<td>STOP</td>
<td>Stop solution</td>
<td>15 ml</td>
</tr>
</tbody>
</table>
5. **Material Required but Not Supplied**

- Bidistilled water (aqua bidist.)
- Precision pipettors and disposable tips to deliver 10-1000 µl
- Foil to cover the microtiter plate
- Horizontal microtiter plate shaker
- A multi-channel dispenser or repeating dispenser
- Centrifuge capable of 10000 x g
- Vortex-Mixer
- Standard laboratory glass or plastic vials, cups, etc.
- Microtiter plate reader at 450 nm (reference wave length 620 or 690 nm)

6. **Preparation and Storage of Reagents**

- To run assay more than once, ensure that reagents are stored at conditions stated on the label. **Prepare only the appropriate amount necessary for each assay.** The kit can be used up to 2 times within the expiry date stated on the label.

- Reagents with a volume less than **100 µl** should be centrifuged before use to avoid loss of volume.

- The **Wash buffer concentrate (WASHBUF)** should be diluted with aqua bidist. **1:10** before use (100 ml concentrate + 900 ml aqua bidist.), mix well. Crystals could occur due to high salt concentration in the stock solutions. The crystals must be redissolved at room temperature or at 37°C using a water bath before dilution of the buffer solutions. The **buffer concentrate** is stable at **2-8°C** until the expiry date stated on the label. Diluted **buffer solution** can be stored in a closed flask at **2-8°C for one month**.

- **Standards (STD) and Controls (CTRL1, CTRL2)** are already diluted in the **reaction buffer (DERBUF)**. Store Standards and Controls frozen at -20°C, thaw before use in the test, and re-freeze immediately after use. Standards and Controls can be re-frozen up to 3 times.
• DMSO could crystallize at 4°C. Dissolve the crystals at 20-25°C in a water bath.

• The content of one vial of derivatization reagent (DER) (50 mg) must be dissolved in 3 ml DMSO. Put the vial on a horizontal shaker for 5 min. After use, the rest of the reagent should be discarded. DER must be prepared immediately before use. The ELISA kit can be separated into two performances by the two DER vials. Please note: DMSO attacks all plastics but not polypropylene products and laboratory glass.

• The content of one vial with ADMA antibody (AB) must be dissolved in 5.6 ml of diluted wash buffer. Therefore, at first, the content of one AB vial is reconstituted with 0.6 ml of diluted wash buffer for 5 minutes. Then the obtained AB solution is quantitatively transferred into a separate vial and 5 ml of diluted wash buffer are added. The ELISA kit can be separated into two performances by the two AB vials. Diluted ADMA antibody (AB) is stable over a longer period of time. It can be stored at 2-8°C for 4 weeks.

• The POD antibody (2.AB) must be diluted 1:200 in conjugate stabilizing buffer (2.ABDIL) (110 µl 2.AB + 22 ml 2.ABDIL). The undiluted POD antibody (2.AB) is stable at 2-8°C until the expiry date stated on the label. Diluted POD antibody (2.AB) is not stable over a longer period. It can be stored at 2-8°C for only 5 days.

• All other test reagents are ready to use. Test reagents are stable until the expiry date (see label of test package) when stored at 2-8°C.

7. PRECAUTIONS

• For in vitro diagnostic use only.

• Human materials used in kit components were tested and found to be negative for HIV and Hepatitis B. However, for safety reasons, all kit components should be treated as if potentially infectious.

• Stop solution is composed of sulfuric acid, which is a strong acid. Even diluted, it still must be handled with care. It can cause acid burns and should be handled with gloves, eye protection, and appropriate protective clothing. Any spills should be wiped out immediately with copious quantities of water.

• Reagents should not be used beyond the expiration date shown on kit label.
8. SPECIMEN COLLECTION AND PREPARATION

Serum, citrate- and EDTA-plasma
- Venous fasting blood is suited for this test system. Samples are stable for one week at 2-8°C. For longer storage samples should be frozen at –20°C up to the measurement.
- Lipemic or hemolytic samples may give erroneous results and should not be used for analysis.
- The serum, citrate- and EDTA-plasma samples are analyzed without any dilution.*
  *If the sample volume is less than 50 µl, a 1:1 dilution in DERBUF (reaction buffer) is recommended (25 µl sample + 25 µl DERBUF). This dilution factor should be considered for data evaluation.
- Samples with visible amounts of precipitates should be centrifuged at least for 5 min at 10000 x g. The resulting supernatant is used in the assay.
- For sample preparation, a DER for coupling of ADMA is added (details are given in the sample preparation procedure).

9. ASSAY PROCEDURE

Procedural notes
- Quality control guidelines should be observed.
- Incubation time, incubation temperature and pipetting volumes of the different components are defined by the producer. Any variations of the test procedure, that are not coordinated with the producer, may influence the test results. Immundiagnostik AG can therefore not be held responsible for any damage resulting from this.
- The assay should always be performed according to the enclosed manual.

Sample preparation procedure
Coupling of standards (STD), controls (CTRL) and samples (SAMPLE) are carried out in single analysis.

1. Bring all reagents and samples to room temperature (18-26°C)

2. Add **200 µl of ready to use standards (STD), 200 µl of ready to use controls (CTRL) and 50 µl of samples (SAMPLE)** in the corresponding vial
3. Add **150 µl of reaction buffer (DERBUF)** only to the samples (SAMPLE)

4. Add **50 µl of freshly prepared derivatization reagent (DER)** into each vial (standards, controls and samples), mix well and incubate **for 45 min on a shaker (180-240 rpm) at room temperature (18-26°C)**

5. Afterwards add **250 µl of dilution buffer (CODIL)** into each vial, mix well and incubate **for 45 min on a shaker (180-240 rpm) at room temperature (18-26°C)**

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**2 x 100 µl of each treated sample (STD, CTRL, SAMPLE) are used in**
in the ELISA in duplicate.

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Test procedure

6. Mark the positions of **standards (STD)/controls (CTRL)/ samples (SAMPLE) in duplicate on a protocol sheet**

7. Take as many microtiter strips (PLATE) as needed from kit. Store unused strips covered at 2-8°C. Strips are stable until the expiry date stated on the label

8. Wash each well **5 times** by dispensing **250 µl of diluted wash buffer** into each well. After the final washing step, the inverted microtiter plate (PLATE) should be firmly tapped on absorbent paper to remove excess solution

9. For the analysis in duplicate, take **2 x 100 µl of standard (STD)/control (CTRL)/samples (SAMPLE)** out of the vial and add into the respective well of the microtiter plate (PLATE)

10. Add **100 µl diluted ADMA antibody (AB)** into each well. Cover the plate tightly
11. Incubate overnight (15-20 hours) at 2-8°C

12. Aspirate the contents of each well. Wash each well 5 times by dispensing 250 µl of diluted wash buffer into each well. After the final washing step, the inverted microtiter plate (PLATE) should be firmly tapped on absorbent paper to remove excess solution.

13. Add 200 µl diluted POD antibody (2. AB) into each well.

14. Cover plate tightly and incubate for 1 hour at room temperature (18-26°C) on a horizontal shaker (180-240 rpm).

15. Aspirate the contents of each well. Wash each well 5 times by dispensing 250 µl of diluted wash buffer into each well. After the final washing step, the inverted microtiter plate (PLATE) should be firmly tapped on absorbent paper to remove excess solution.

16. Add 200 µl of TMB substrate (SUB) into each well.

17. Incubate for 6-10 min at room temperature (18-26°C) in the dark*.

18. Add 100 µl of stop solution (STOP) into each well, mix thoroughly.

19. Determine absorption immediately with an ELISA reader at 450 nm against 620 nm (or 690 nm) as a reference. If no reference wavelength is available, read only at 450 nm. If the extinction of the highest standard exceeds the range of the photometer, absorption must be measured immediately at 405 nm against 620 nm as a reference.

*The intensity of the color change is temperature sensitive. We recommend to observe the color change and to stop the reaction upon good differentiation.
10. EVALUATION OF RESULTS

If the test is performed in strict compliance with the manufacturer’s instructions, e.g. with the exact volumes for standards, controls and samples/sample treatment, standards, controls and samples are equally diluted. Therefore, no dilution factor is required for calculation of the results. **

**At a 1:1 dilution, the dilution factor should be considered.

The following algorithms can be used alternatively to calculate the results. We recommend to use the "4-parameter-algorithm".

1. 4-parameter-algorithm
   
   It is recommended to use a linear ordinate for optical density and a logarithmic abscissa for concentration. When using a logarithmic abscissa, the zero calibrator must be specified with a value less than 1 (e.g. 0.01).

2. Point-to-point-calculation
   
   We recommend a linear ordinate for optical density and a linear abscissa for concentration.

3. Spline-algorithm
   
   We recommend a linear ordinate for optical density and a logarithmic abscissa for concentration. When using a logarithmic abscissa, the zero calibrator must be specified with a value less than 1 (e.g. 0.01).

   The plausibility of the pairs of values should be examined before the automatic evaluation of the results. If this option is not available with the used program, a control of the paired values should be done manually.

Expected values

Based on internal studies of evidently healthy persons (n=70) a mean value of 0.45 µmol/l was estimated.

Normal range:

Serum/Plasma mean value ± 2 standard variations: 0.45 ± 0.19 µmol/l

We recommend each laboratory to develop its own normal range. The values mentioned above are only for orientation and can deviate from other published data.
Controls

Control samples or serum pools should be analyzed with each run. Results, generated from the analysis of the control samples, should be evaluated for acceptability using appropriate statistical methods. The results for the patient samples may not be valid, if within the same assay one or more values of the quality control sample are outside the acceptable limits.

The concentration of controls and patient samples can be determined directly from calibration curve. In the following an example of a calibration curve is given.

Example of calibration curve
11. PERFORMANCE CHARACTERISTICS

Cross reactivity

SDMA < 0,5 %  NMMA < 0,5 %  L-Arginin < 0,02 %

Precision and reproducibility

<table>
<thead>
<tr>
<th>Sample</th>
<th>ADMA [µmol/l]</th>
<th>Standard variation (SD) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0,27</td>
<td>0,021</td>
</tr>
<tr>
<td>2</td>
<td>0,78</td>
<td>0,041</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample</th>
<th>ADMA [µmol/l]</th>
<th>Standard variation (SD) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0,33</td>
<td>0,029</td>
</tr>
<tr>
<td>2</td>
<td>0,79</td>
<td>0,045</td>
</tr>
</tbody>
</table>

Sensitivity
The sensitivity was set as B₀ + 1SD. The zero-standard was measured 6 times.

<table>
<thead>
<tr>
<th>Sample</th>
<th>ADMA mean value [OD]</th>
<th>Standard variation (SD) [%]</th>
<th>Detection limit [µmol/l]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>2,78</td>
<td>0,12</td>
<td>0,05</td>
</tr>
</tbody>
</table>
Recovery

One sample was spiked with different ADMA concentrations and measured using this assay. The analytical recovery rate was determined by the expected and measured ADMA levels. The expected levels were calculated as the sum of the measured ADMA concentration in the original sample and the spiked ADMA amount. The mean recovery rate for all concentrations was 104 % (n=5).

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>x</td>
<td>x=0,419</td>
<td>100</td>
</tr>
<tr>
<td>0,25</td>
<td>0,25+x=0,669</td>
<td>0,694</td>
<td>104</td>
</tr>
<tr>
<td>0,5</td>
<td>0,50+x=0,919</td>
<td>0,948</td>
<td>103</td>
</tr>
</tbody>
</table>

Linearity

The linearity of the ELISA was determined by the dilution of a spiked patient sample. The mean linearity was 103%.

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>original</td>
<td>0,468</td>
<td>0,468</td>
<td>100</td>
</tr>
<tr>
<td>1+1</td>
<td>0,255</td>
<td>0,234</td>
<td>109</td>
</tr>
<tr>
<td>1+3</td>
<td>0,114</td>
<td>0,117</td>
<td>97</td>
</tr>
</tbody>
</table>

12. LIMITATIONS

Strong hemolytic and lipemic samples often show wrong concentrations. Do not to measure hemolytic and lipemic samples.
13. REFERENCES


14. General Notes on the Test and Test Procedure

- This assay was produced and put on the market according to the IVD guidelines of 98/79/EC.
- Test components contain organic solvents. Contact with skin or mucous membranes must be avoided.
- All reagents in the test package are for in-vitro-diagnostic use only.
- Reagents should not be used after the date of expiry stated on the label.
- Single components with different lot numbers should not be mixed or exchanged.
- Guidelines for medical laboratories should be observed.
- Incubation time, incubation temperature and pipetting volumes of the components are defined by the producer. Any variation of the test procedure, which is not coordinated with the producer, may influence the results of the test. Immundiagnostik AG can therefore not be held responsible for any damage resulting from wrong use.

18.12.2009

Used symbols:

- **Temperature limitation**
- **Catalogue Number**
- **In Vitro Diagnostic Medical Device**
- **Contains sufficient for <n> tests**
- **Manufacturer**
- **Use by**
- **Lot number**