

ALBUWELL™ M ELISA



DESCRIPTION

Intended Use

Albuwell™ M is an enzyme linked immunosorbent assay (ELISA) for the quantitative determination of albumin in mouse urine.

Technical Background

Albuwell M is an *in vitro* tool for assessing kidney function in mice. It is simple to perform and highly specific for mouse albumin. It is a competitive antibody capture ELISA completed in a direct mode (See Appendix, Fig. 1). To that end, the Anti-albumin antibody is conjugated to horseradish peroxidase (Anti-mouse Albumin Ab-HRP), i.e., directly labeled.

To complete the assay, sample and Anti-mouse Albumin Ab-HRP are added to the mouse albumin-coated well. The antibody interacts and binds with the albumin immobilized onto the stationary phase or with that in the fluid phase, hence the notion of competitive binding. Washing removes unbound Anti-mouse Albumin Ab-HRP and other reactants of the fluid phase from the well. Only the antibody-conjugate that was bound to the albumin of the stationary phase remains, and this is detected using tetramethylbenzidine (TMB) in a chromogenic reaction. The reaction is stopped with acid, and absorbance is measured at 450 nm. Absorbance is inversely proportional to the logarithm of albumin concentration in the fluid phase.

SPECIMEN COLLECTION AND STORAGE

Collect samples without preservative and clarify them by centrifugation if necessary. Mouse urine is often contaminated by food and fecal material, and these contaminants present potential sources of error. Store clarified urine at 4°C for up to 1 week or at -60°C for up to 2 months. Prior to assay, allow the samples to come to room temperature. Do not apply heat to thaw frozen samples.

MATERIALS NEEDED

Kit Contents

- 2 Albuwell M Test Plates
- 2 NHEBSA (Diluent)
- Mouse Serum Albumin (MSA) Standard
- 2 Anti-mouse Albumin Ab-HRP Conjugate

- 2 Color Developer
- 2 Color Stopper
- Instructions

MSA Standard, NHEBSA and Anti-mouse Albumin Ab-HRP Conjugate preparations contain 0.05% Proclin® 300 (active components isothiazolones) as preservative. Color Stopper contains dilute (2.0 N) sulfuric acid.

Albuwell M plates are pre-coated and ready to use. All other kit reagents are supplied in ready to use liquid form. A plate wash procedure using tap water has been shown to be suitable in experimental and quality control contexts. However, where tap water is unavailable or proves unsuitable, we recommend using the Ethos Dilurex™ TEA Wash Buffer (pH 6.8) available at 10x and 1x concentrations.

Additional Supplies Needed

Micropipettors capable of delivering 10, 50, 100, 120 and 200 µL are required. Multi-channel pipettors capable of delivering 50 and 100 µL are recommended. In addition, small test tubes are required to complete dilutions (microfuge tubes work well in this application). Finally, a microplate reader equipped to determine absorbance at 450 nm is required.

ASSAY PROCEDURE

Allow reagents and samples to come to room temperature before running the assay.

Dilute Standard and Samples carefully. For the standards, a single tip may be used to prepare the dilution series. For the experimental samples, a fresh tip should be used for each urine specimen.

Standard Dilutions

This procedure describes the preparation of seven (7) two-fold dilutions of standard (See Appendix, Fig. 2).

1. Prepare 8 microfuge tubes with 200 µL of NHEBSA per tube.
2. Label the tubes C and 1 to 7.
3. Transfer 200 µL of MSA Standard to tube 1.
4. Mix contents by aspirating and expelling the fluids 5 times.
5. Transfer 200 µL of solution from tube 1 to tube 2.

6. Mix as before.
7. Continue this procedure through tube 7.
8. Tubes 1 to 7 now contain concentrations of 10.0, 5.0, 2.5, 1.25, 0.625, 0.313 and 0.156 µg MSA/mL respectively.

Preparation of Urine Sample Dilutions

Accurate determination of urinary albumin depends upon proper sample dilution. In most cases, a 1:21 dilution is sufficient, but collection methods and animal kidney function (or dysfunction) may lead to exceptionally high or exceptionally low concentrations. For initial studies it is wise to complete the analysis at three concentrations, such as undilute, 1:21, and 1:81, to determine the dilution that falls within the linear range of this assay.

The following example illustrates a 1:21 dilution protocol.

1. Prepare and label a microfuge tube for each sample.
2. Add 200 µL NHEBSA to each tube.
3. Use a dry fresh tip to transfer 10 µL of sample to the appropriate tube, wash out the tip by repeated aspiration and expellation in the tube.
4. Vortex the tube briefly.
5. Continue this procedure for the rest of the samples.
6. Each sample is now diluted 1:21 in NHEBSA.

Addition of Controls, Standard MSA Dilutions, and Samples to the Plate

Label the strips with an indelible marker, 1-12. This will allow reconstruction of the plate should strips fall out during the washing procedures. The diluted standards and samples may be added directly to the dry plate. A plate map is provided to assist with this section, see Appendix, Figure 3.

The plate design includes two controls. Well A1 is a negative control, and well A2 is a positive control. All other wells in column 1 and 2 are diluted MSA standards. The balance of the plate is available for diluted samples.

1. Add an 100 µL aliquot of NHEBSA from tube C to well A1. This will be the negative control. Do NOT add any Anti-mouse Albumin Ab-HRP conjugate during the primary incubation.
2. Add 50 µL of NHEBSA from tube C to well A2. This will be the positive control.
3. With a fresh tip, pre-wet the tip in standard dilution number 7, and transfer 50 µL aliquots to wells H1 and H2.
4. With the same tip, pre-wet/rinse the tip in standard dilution number 6, and transfer 50 µL aliquots to wells G1 and G2.
5. Continue transferring diluted standard to the plate in this fashion, i. e. pre-wetting/rinsing the tip and transferring aliquots of standard in order.
6. Taking care to change the tip and to pre-wet it each time, add diluted samples to the plate. Samples can be added in duplicates starting at A3 and A4 and continuing through the rest of the plate.
7. The plate now contains controls and diluted standards in columns 1 and 2, and experimental samples in duplicate in the balance of the plate.

Primary Incubation: Reaction with Anti-mouse Albumin Ab-HRP conjugate.

1. Add 50 µL of Anti-mouse Albumin Ab-HRP conjugate to any well except A1, which is the negative control.
2. Cover and incubate the plate for 30 minutes.
3. Wash plate using a plate washer or wash plate by hand as follows: Remove fluids from the well, ie. aspirate off fluids or flip them out into a sink.
4. Fill wells to over-flowing with water or wash buffer.
5. Remove fluids as before.
6. Steps 4 and 5 constitute a wash cycle.
7. Repeat the process to yield a total of 10 wash cycles.
8. Invert the plate on a paper towel and tap gently to remove excess fluids.

Color Development and Measurement

1. Add 100 µL of Color Developer to each well.
2. Develop 5-10 minutes.
3. Add 100 µL of Color Stopper to each well.
4. Examine the plate. The negative control in well A1, should have little or no color, but the positive control in well A2, should be the most intensely colored well on the plate. The rest of the wells should show absorbances intermediate between these extremes.
5. Use a plate reader to determine the absorbance at 450 nm. Use well A1 as the blank.

ANALYSIS

This analysis assumes that computer and analysis software, e.g. MS Excel®, is available.

1. Prepare a spreadsheet entering appropriate data including standard dilution, concentration, sample dilution, and absorbance data. Determine the mean for replicate wells.
2. Prepare a semi-logarithmic plot of standard dilutions with the \log_{10} [MSA] on the x-axis and mean absorbance on the y-axis. This is the standard curve.
3. The data that fall into the linear portion of the standard curve constitute the usable portion of the assay.
4. Subject these data to semi-logarithmic analysis to yield a mathematical model, of the form:
 $\log_{10} [\text{MSA}] = m A450 + b$
5. MSA concentration is determined by taking the anti-log of the calculated values from this equation.
6. Multiply by 21 (or inverse dilution factor) to correct for the dilution.

QUALITY CONTROL

Record Keeping: It is good laboratory practice to record the lot numbers and dates for the kit components and reagents used for each assay.

LIMITATIONS

1. Samples must not contain inhibitors for HRP, i.e., sodium azide. These will affect results.
2. It is the responsibility of the investigator to determine if the presence of experimental compounds or their metabolites in the urine will affect the assay results.
3. Gross microbiological contamination may affect assay results.
4. Bloody urine specimens are unsuitable for use, even if clarified by centrifugation, since blood flow is a sign of contamination and since albumin concentrations in the blood are approximately 2000 times those normally found in urine. Semen contains significant levels of albumin and is also a potential source of contamination.

TROUBLE SHOOTING

1. No color appears after adding Color Developer: One or more reagents may have been adversely affected by storage above 8°C. One or more reagents may not have been added. Repeat assay. Be sure to store the kit appropriately.
2. Color in wells too light: Longer incubation with Color Developer may be required. If the color is still too light after 10 minutes development, repeat the assay but increase the primary incubation with Anti-mouse Albumin Ab-HRP conjugate to 1 hour.
3. Color in wells is too dark: Decrease the development time. If a 5-minute development is still too dark, repeat the assay and reduce the primary incubation with Anti-mouse Albumin Ab-HRP conjugate to 15 minutes.
4. If color is dark and the standard dilutions fail to show the appropriate linearity, Color Developer may have been contaminated with conjugate or the plate was poorly washed. Repeat the assay and take care in the pipetting and in the washing operations.
5. Color in sample well(s) is darker or lighter than lowest or highest concentrations of the standard curve. Change sample dilution protocol appropriately.
6. Poor agreement between duplicate wells. This is almost always due to pipetting error. Repeat the assay.
7. Microplate ELISAs may be prone to edge effects wherein the outer rows and columns show a darker response than the inner ones. This effect may be minimized by incubating the plate in a closed humid container. A plastic food storage container with a tight-fitting lid and a water moistened paper towel work well in this respect. Place the moistened towel in the bottom of the container and place the plate upon it. Position the cover and incubate as described.

PRODUCT INFORMATION

CAT. #	DESCRIPTION
1011	Albuwell™ M ELISA
1012	Creatinine Companion Assay

Order today at [EthosBiosciences.com](https://www.ethosbiosciences.com)

TRADEMARKS

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APPENDIX

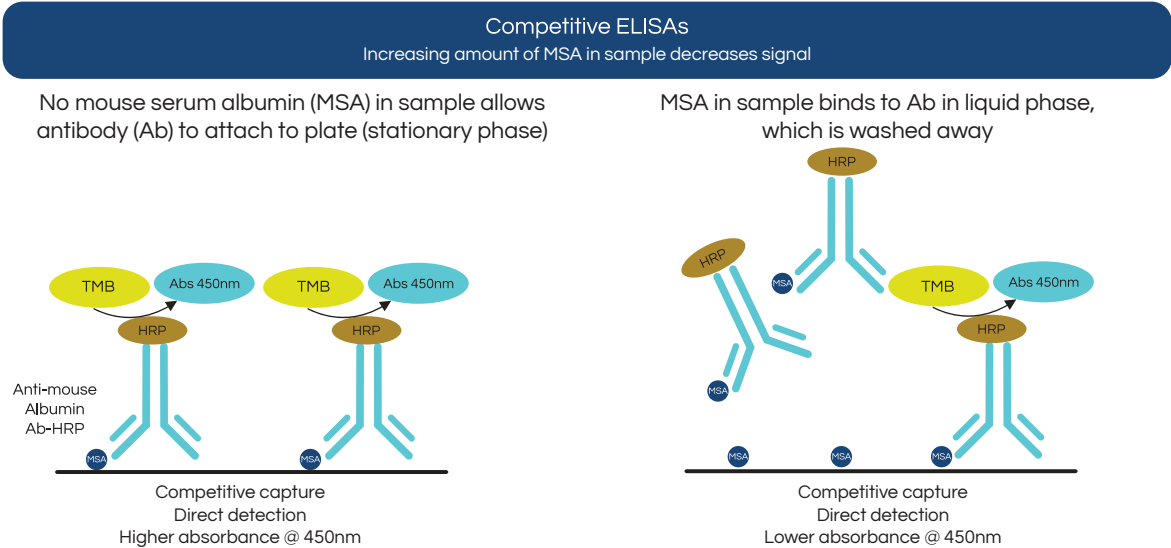


Fig. 1. Competitive ELISAs have decreasing signals when albumin is present in samples. The HRP molecule is directly attached to the Fc region of the anti-mouse albumin antibody, thus eliminating the need for adding a labeled secondary antibody. The left diagram represents what happens in well A2 of this kit. The right diagram represents the addition of MSA standards in wells B1 and B2 through H1 and H2. The more albumin in the sample, the lower the absorbance reading since the Anti-mouse Albumin Ab-HRP conjugate will wash away in the liquid phase rather than binding to the plate.

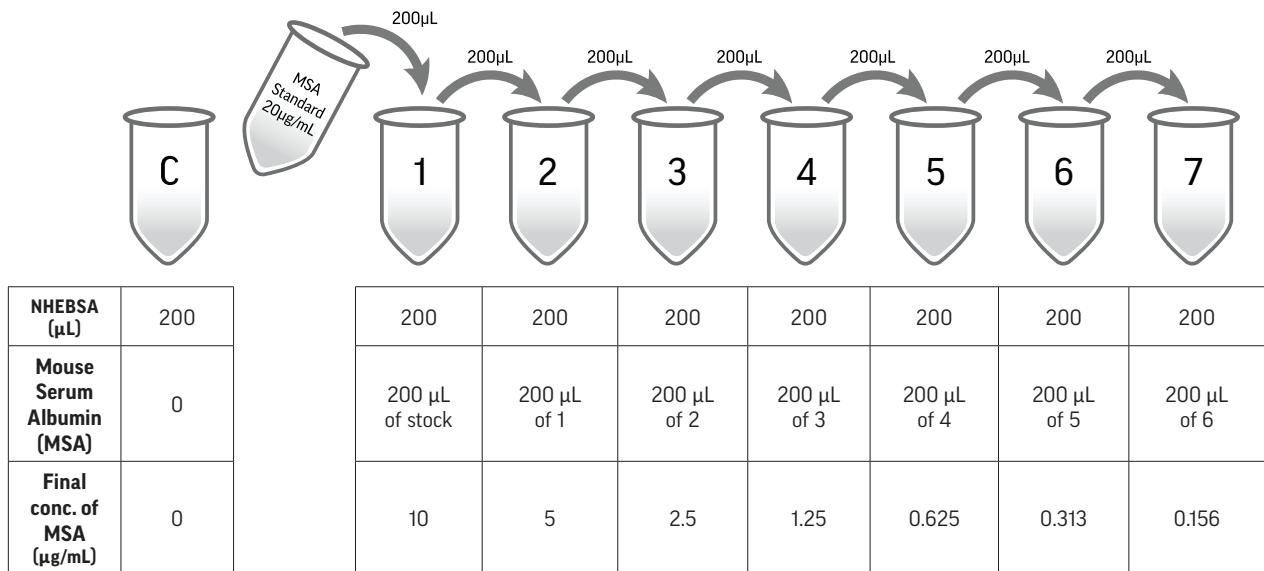


Fig. 2. Creating the mouse serum albumin (MSA) standard dilution series. MSA is supplied at 20 µg/mL stock solution that is diluted at 1:2 seven consecutive times to establish a standard curve for estimating the amount of albumin in experimental samples.

APPENDIX, CONTINUED

Key
 #1-7 = MSA dilutions
 αAlb-HRP = αMouse Albumin Ab

PLATE MAP
 Add 50 µL of each of the following to each well EXCEPT A1

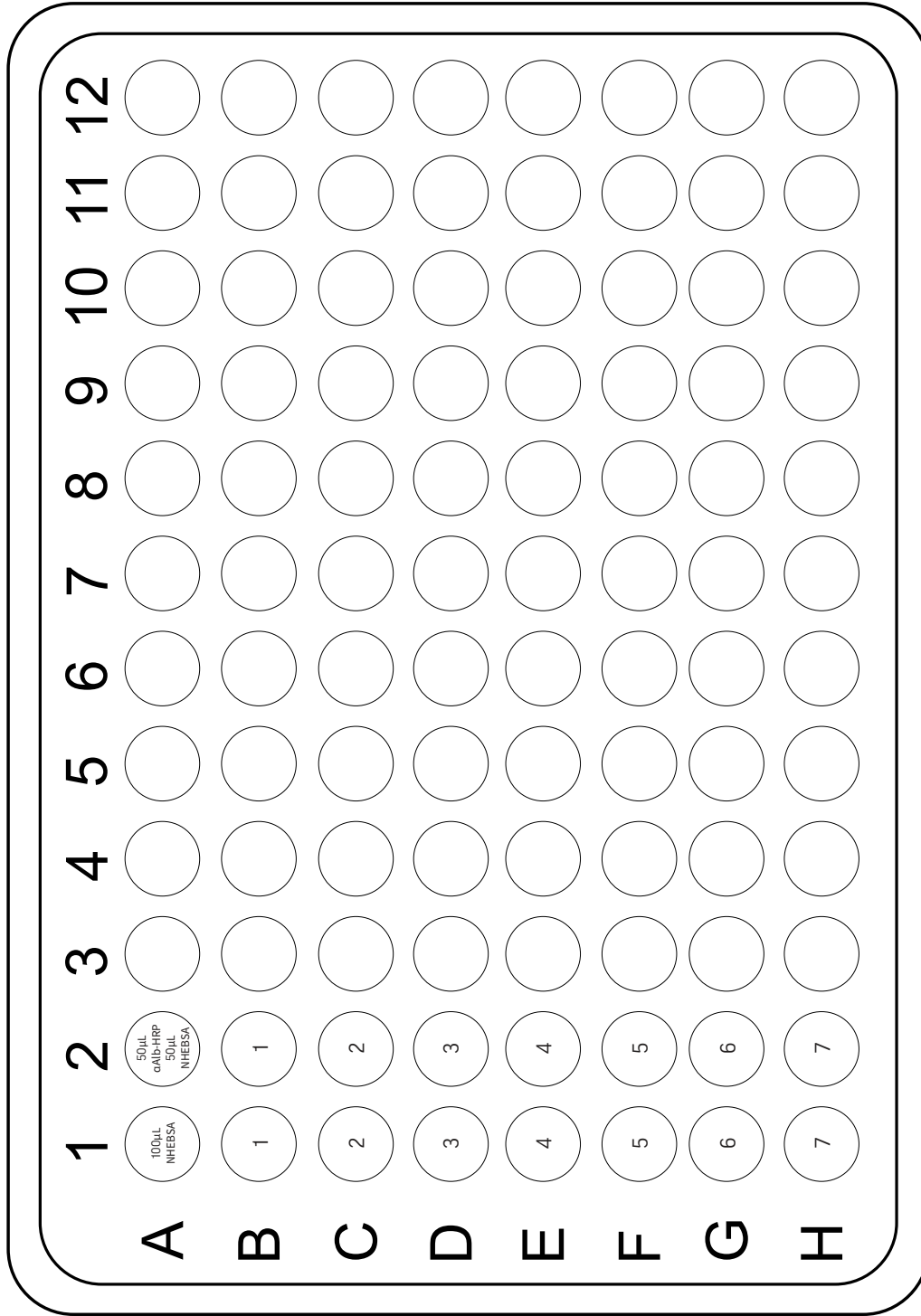


Fig. 3. Plate map for the Addition of Controls, MSA, and Samples. Well A1 has 100 µL of NHEBSA buffer only, which will be the negative control. Well A2 has 50 µL of NHEBSA buffer + 50 µL of αMouse Albumin Antibody-HRP conjugate as a positive control. The remaining wells in column 1 and 2 have 50 µL of each MSA serial dilution in duplicate to create the standard curve. Columns 3 through 12 are available for up to 40 samples tested in duplicate.