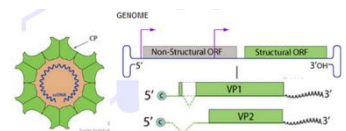


## INTENDED USE

The Mouse Anti-Minute Virus of Mice (MVM) IgG ELISA Kit is an indirect ELISA suitable for detecting and quantifying IgG antibody activity specific for MVM-capsid protein (VP2) in serum or plasma or other biological fluids. The kit uses recombinant MVM-VP2 antigen and has no virus or viral proteins to avoid contamination. For research use only (RUO).

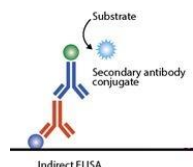
## GENERAL INFORMATION



The Minute Virus of Mice (MVM) is the prototype virus of the Protoparvovirus genus within the Parvoviridae family of viruses.

MVM exists in two variant forms: MVMp, which is the prototype strain, infects cells of fibroblast origin, while MVMi, the immunosuppressive strain, infects T lymphocytes. MVM is a common infection in laboratory mice due to its highly contagious nature. Minute virus of mice (MVM) and mouse parvovirus (MPV or MPV-1) are among the most prevalent infectious agents detected in contemporary laboratory mouse colonies, with approximately 45% of USA research institutions harboring these infectious agents and MPV being among the most prevalent viruses detected in research mice. Various clinical disease syndromes in mice have been associated with MVM infection and both MVM and MPV can have deleterious effects on research due to in vitro and in vivo immunomodulatory effects and contamination of cell cultures and tissues originating from mice. As a result, murine parvovirus infections comprise one of the most significant infectious disease problems encountered in contemporary laboratory animal research facilities.

## PRINCIPLE OF THE TEST



The Mouse Anti-LCMV IgG ELISA kit is based on the binding of mouse anti-MVM IgG in samples to MVM antigen immobilized on the microwells, and anti-MVM IgG antibody is detected by anti-mouse IgG-HRP conjugate. After a washing step, chromogenic substrate (TMB) is added and color (blue) is developed by, which is directly proportional to the amount of anti-MVM IgG present in the sample. Stopping Solution is added to terminate the reaction (blue converts to yellow), and A450nm is then measured using an ELISA reader. The presence of mouse IgG antibody in samples is determined relative to mouse anti-MVM IgG Controls.

## PRODUCT SPECIFICATIONS

### Specificity

Purified recombinant MVM-VP2 protein is used as antigen; thus the assay is specific for antibodies directed to MVM-VP2. The Antibody-HRP conjugate reacts specifically with mouse IgG class antibodies; IgA, IgM and IgE antibody would not be measured above background signals. The Anti-Mouse IgG conjugate is blended to equally quantify various IgGs (IgG1-3) subclasses, important when considering difference in subclass emphasis between natural infection and MAP immunization.

### Assay Sensitivity

The MVM-VP2 antigen coating level, HRP conjugate concentration, and Low NSB Sample Diluent are optimized to differentiate anti-MVM IgG from background (non-antibody) signal with mouse serum samples diluted 1:100.

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## KIT CONTENTS

The microtiter well plate and all other reagents, if unopened, are stable at 2-8°C until the expiration date printed on the box label. Stabilities of the working solutions are indicated under Reagent Preparation.

**To Be Reconstituted:** Store as indicated.

Component	Preparation Instructions
<b>Wash Solution Concentrate (100x)</b> Cat. No. WB-100, 10ml	Dilute the entire volume 10ml + 990ml with distilled or deionized water into a clean stock bottle. Label as <b>Working Wash Solution</b> and store at ambient temperature until kit is used entirely.
<b>Sample Diluent Concentrate (20x)</b> Cat. No. SD-20T, 10ml	Dilute the entire volume, 10ml + 190ml with distilled or deionized water into a clean stock bottle. Label as <b>Working Sample Diluent</b> and store at 2-8°C until the kit lot expires or is used up.
<b>Anti-Mouse IgG-HRP Conjugate Concentrate (100x)</b> Part: H-MsG-112b, 0.15ml	Peroxidase conjugated anti-Mouse IgG in buffer with detergents and antimicrobial as stabilizers. Dilute fresh as needed; 10ul of concentrate to 1ml of <b>Working Sample Diluent</b> is sufficient for 18-well strip. Use within the working day and discard. Return 100X to 2-8°C storage.

**Ready For Use:** Store as indicated on labels.

Component	Part	Amt	Contents
<b>MVM Ag Microwell Strip Plate</b>	300201	8-well strips (12)	Coated with MVM VP2 antigen, and post-coated with stabilizers.
<b>Mouse Anti-MVM Sensitivity Control</b>	300202	0.65 ml	Low level Mouse Anti-MVM, in buffer with detergents and antimicrobial.
<b>Mouse Anti-MVM Positive Control</b>	300203	0.65 ml	High level Mouse Anti-MVM, in buffer with detergents and antimicrobial.
<b>Low NSB Sample Diluent</b>	TBTm	30 ml	Buffer with protein, detergents and antimicrobial. Use as is for sample dilution
<b>TMB Substrate</b>	80091	12 ml	Chromogenic substrate for HRP containing TMB and peroxide.
<b>Stop Solution</b>	80101	12 ml	Dilute sulfuric acid.

### Materials Required But Not Provided:

- Pipettors and pipettes that deliver 100ul and 1-10ml. A multi-channel pipettor is recommended.
- Disposable glass or plastic 5-15ml tubes for diluting samples and Anti-Mouse IgG HRP Concentrate.
- Stock bottle to store diluted Wash Solution; 0.2 to 1L.
- Distilled or deionized water to dilute reagent concentrates.
- Microwell plate reader at 450 nm wavelength.

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## ASSAY DESIGN AND SET-UP

### Sample Collection and Handling

Serum and other biological fluids may be used as samples with proper dilution to avoid solution matrix interference. For serum, collect blood by venipuncture, allow clotting, and separate the serum by centrifugation at room temperature. For other samples, clarify the sample by centrifugation and/or filtration prior to dilution in Sample Diluent. If samples will not be assayed immediately, store refrigerated for up to a few weeks, or frozen for long-term storage.

### Antibody Stability

Initial dilution of serum into **Working Sample Diluent** (WSD) is recommended to stabilize antibody activity. This enhances reproducible sampling, and stabilizes the antibody activity for years, stored refrigerated or frozen. Further dilution into **Low NSB Sample Diluent** (LNSD), which provides the lowest assay background, should be at least 10 times the initial dilution and performed the same day as the assay.

Example:

Initial (1/5): 10ul serum + 40ul WSD [or 0.1ml + 0.4ml]  
Further (1/100): 10ul initial (1/5) + 190ul LNSD

### Assay Design

Review Calculation of Results (p5-7) and Limits of the Assay (above) before proceeding:

- Select the proper sample dilutions accounting for expected potency of positives and minimizing non-specific binding (NSB) and other matrix effects; for example, net signal for non-immune samples should be lower than the **Sensitivity Control**. This is usually 1/100 or greater dilution for mouse sera with normal levels of IgG and IgM.

- Run a Sample Diluent **Blank**. This signal is an indicator of proper assay performance, especially of washing efficacy, and is used for net OD calculations, if required. Blank OD should be <0.3.

- Run the **Positive** and **Sensitivity Controls**, which validate that the assay was performed to specifications: the **Positive Control** should give a high signal (>1.5 OD); the **Sensitivity Control** should give a low signal which can be used to discriminate at the Positive/Negative threshold (see Interpretation of Results, p. 5).

### Plate Set-up

Bring all reagents to room temperature (18-30° C) equilibration (at least 30 minutes).

- Determine the number of wells for the assay run. Duplicates are recommended, including 4 Control wells and 2 wells for each sample and internal control to be assayed.

- Remove the appropriate number of microwell strips from the pouch and return unused strips to the pouch. Reseal the pouch and store refrigerated.

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- Add 200-300ul Working Wash Solution to each well and let stand for about 5 minutes. Aspirate or dump the liquid and pat dry on a paper towel before sample addition.

## Assay Procedure

ALL STEPS ARE PERFORMED AT ROOM TEMPERATURE. After each reagent addition, gently tap the plate to mix the well contents prior to beginning incubation.

### 1. 1<sup>st</sup> Incubation [50 or 100ul – 60 min; 4 washes]

- Add 100ul of sample diluent (blank) calibrators, samples and controls each to pre-determined wells.
- Tap the plate gently to mix reagents and incubate for 60 minutes.
- Wash wells 4 times and pat dry on fresh paper towels. As an alternative, an automatic plate washer may be used. Improper washes may lead to falsely elevated signals and poor reproducibility.

### 2. 2<sup>nd</sup> Incubation [100ul – 30 min; 5 washes]

- Add 100ul of diluted Anti-Mouse IgG HRP to each well.
- Incubate for 30 minutes.
- Wash wells 5 times as in step 2.

### 3. Substrate Incubation [100ul – 15 min]

- Add 100ul TMB Substrate to each well. The liquid in the wells will begin to turn blue.
- Incubate for 15 minutes in the dark, e.g., place in a drawer or closet.

Note: If your microplate reader does not register optical density (OD) above 2.0, incubate for less time, or read OD at 405-410 nm (results are valid).

### 4. Stop Step [Stop: 100ul]

- Add 100ul of Stop Solution to each well.
- Tap gently to mix. The enzyme reaction will stop; liquid in the wells will turn yellow.

### 5. Absorbance Reading

- Use any commercially available microplate reader capable of reading at 450nm wavelength. Use a program suitable for obtaining OD readings, and data calculations if available.
- Read absorbance of the entire plate at 450nm using a single wavelength within 30 minutes after Stop Solution addition. If available, program to subtract OD at 630nm to normalize well background.

## PRECAUTIONS AND SAFETY INSTRUCTIONS

Controls, Sample Diluent, and Antibody HRP contain bromonitrodioxane (BND: 0.05%, w/v). Stop Solution contains dilute sulfuric acid. Follow good laboratory practices, and avoid ingestion or contact of any reagent with skin, eyes or mucous membranes. All reagents may be disposed of down a drain with copious amounts of water. MSDS for TMB, sulfuric acid and BND can be requested or obtained from the ADI website: [http://4adi.com/commerce/info/showpage.jsp?page\\_id=1060&catqory\\_id=2430&visit=10](http://4adi.com/commerce/info/showpage.jsp?page_id=1060&catqory_id=2430&visit=10)

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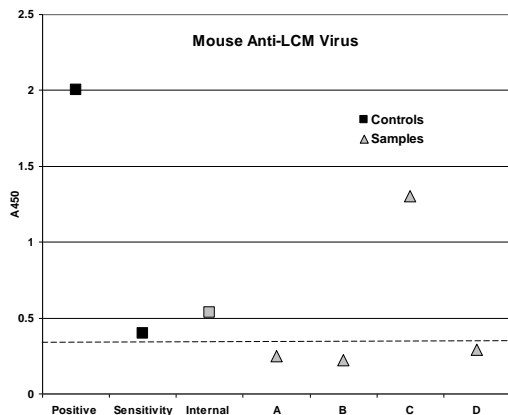
## INTERPRETATION OF RESULTS

### A. Antibody Activity Threshold Index

Compare Samples to **Sensitivity Control** or **Internal Control**

= **Positive/Negative Cut-off.**

#### Example:



#### Results

The **sensitivity** of the assay to detect anti-MVM IgG, from either natural infection or MAP, is controlled so that the **Sensitivity Control** represents a threshold OD for most true positives in mouse serum diluted in the Low NSB Sample Diluent at 1:100 or greater. Visual inspection of the data in the above graph shows the following:

**Positive Control** – clearly positive, shows the OD range of the assay; high value indicates optimal sensitivity of the assay.

**Sensitivity Control** – a 'Cut-off' line has been drawn to indicate a threshold distinguishing between **Positive/Negative**. This is not a clear-cut threshold, rather a low OD area that could represent either low positives or high background negatives.

**Internal Control** – a true positive from an infected animal that represents the lab's experience in distinguishing low positive from negative samples. This should be run in each assay to supplement the Sensitivity Control for Positive/Negative discrimination purposes.

**Samples A,B,C,D** – 3 samples (A, B, D) are negative: below the threshold; 1 sample (C) is positive: clearly above the threshold.

The Sensitivity Control can be used to calculate a **Threshold Index** that numerically discriminates Positive/Negative, as follows:

- ❖ Divide each Sample net OD by the Sensitivity Control net OD. Values above 1.0 are a measure of **Positive** Antibody Activity; below 1.0 are **Negative** for antibody.

This calculation was used to represent Assay Precision, page 7.

## INTERPRETATION OF RESULTS (cont)

### B. Positive Index

Experimental sample values may be expressed relative to the values of Control or Non-immune samples, by calculation of a **Positive Index**. One typical method is as follows:

1. Calculate the net OD mean + 2 SD of the Control/Non-immune samples = **Positive Index**.
2. Divide each sample net OD by the Positive Index. Values above 1.0 are a measure of **Positive** Antibody Activity; below 1.0 are **Negative** for antibody.

A sample value would be **Positive** if significantly above the value of the pre-immune serum sample or a suitably determined non-immune panel or pool of samples, tested at the same sample dilution.

This calculation also **quantifies** the positive Antibody Activity level, assigning a higher value to samples with higher Antibody Activity, and vice versa.

#### Example:

Sample	Assay Net OD		Calculated Antibody Activity	
	Control	Exptl	Control	Exptl
1	0.248	<b>P</b> 2.212	0.79	<b>7.04</b>
2	0.290	<b>S</b> 0.452	0.92	<b>1.44</b>
3	0.186	<b>I</b> 0.541	0.59	<b>1.72</b>
4	0.276	<b>U</b> 0.212	0.88	0.68
5	0.161	<b>U</b> 0.122	0.51	0.39
6	0.173	<b>M</b> 1.491	0.55	<b>4.75</b>
7	0.153	<b>M</b> 0.694	0.48	<b>2.21</b>
8	0.211	<b>N</b> 1.487	0.67	<b>4.74</b>
9	0.145	<b>N</b> 0.546	0.46	<b>1.74</b>
10	0.110	<b>U</b> 0.263	0.35	0.84
Mean	0.195			
SD	0.0595			
Mean +2 SD	<b>0.314</b>	<b>= Positive Index</b>		

#### Results

**Experimental Samples** are represented as follows:

**P** – Positive Control

**S** – Sensitivity Control

**I** – Internal Control; lab's threshold positive serum

**U** – Uninfected mouse sample

**M** – Mouse Antibody Production (MAP) samples represent injection of MVM antigen (same as used for plate coating) into mice; positive indicates presence of the MVM in the inoculum.

**N** – Naturally infected mouse samples.

**Note:** The prevalent IgG antibody from MAP samples can be a different subclass (often IgG1, depending on adjuvant used) from the prevalent subclass from natural infection (IgG2a). The Anti-IgG-HRP used in the kit has been balanced for equal sensitivity for the IgG1 and IgG2a subclasses, therefore avoiding a bias in assay sensitivity for the various uses of the assay.

## ASSAY PERFORMANCE

### Precision

Samples and Controls (as shown in the graph, page 5) were assayed in duplicate in 5 - 6 separate runs, to provide a measure of between-assay reproducibility.

The data are represented using the value of the Sensitivity Control in each assay to calculate a **Threshold Index** for each control and samples (as described on page 5).

Sample	Ave OD	Threshold Index (mean)	Inter-assay %CV
Positive Control	2.00	5.03	<b>7.8</b>
Sensitivity Control	0.40	<b>1.0</b>	<b>0</b>
Internal Control	0.54	1.34	<b>5.8</b>
Natural Positive	1.30	3.38	<b>14.8</b>
MAP Positive	1.29	3.33	<b>5.1</b>

#### Results

The coefficient of variation (%CV) shows the reproducibility of the assay for measuring one antibody activity (sample or control) relative to another antibody activity (**Sensitivity Control**). Variation increases in the threshold region; for this reason, consider running additional tests for borderline samples.

#### Note:

- The sensitivity of the assay may be increased to perhaps convert a borderline sample to a positive by using a lower dilution of the sample, e.g., 1/50. The values of negatives may increase, so an alternative threshold should be established using known negatives to develop a **Positive Index** (page 6), or by using known **Internal Controls** as discriminator for a **Threshold Control** (instead of the kit Sensitivity Control)

#### Limits of the Assay

- The assay detects and quantifies IgG antibodies directed to the major capsid protein VP1. It may be possible for an animal to have Lymphocytic Choriomeningitis Virus infection without producing antibodies specific to VP1.
- Anti-MVM antibody levels of an infected animal may be below detection threshold related to the time course of the infection, e.g., too early for positive titer development.

#### Specificity & Cross-reactivity

MVM-VP1 and VP2 are produced as a result of alternative splicing and the two proteins have significant overlap. MVM-VP2 shares significant sequence conservation with the related mouse parvoviruses (~74%) and rat minute virus (72%), Kilham rat virus (71%) canine parvovirus (52%), hamster parvovirus (66%), rat parvovirus (60%) and canine parvovirus 2a/b & mink enteritis virus, blue fox parvovirus, Feline panleukopenia virus (52%). Antibody crossreactivity of MVM-VP2 with various related VP2s has not been studied.

**References:** Ball-Goodrich LJ (1994) J. Virol. 68, 6476; Brownstein DG (1991) Lab. Invest. 65, 357; Astell CR (1986) J. Virol. 57, 656; sahl R (1985) Nucl. Acid., red. 13, 3617

# RecombiVirus™ Mouse Anti-Minute Virus of Mice (MVM) IgG ELISA Kit

**Cat. #. AE-300900-1, 96 Tests**

For Quantitation of Anti-MVM-VP2 IgG in Serum or plasma or other biological fluids

For in vitro research use only (RUO), not for therapeutic or diagnostic use.



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