

**Mouse Immunoglobulin
Isotyping Magnetic Bead
Panel**

96-Well Plate Assay

MGAMMAG-300K

MILLIPLEX[®] MAP

Mouse Immunoglobulin Isotyping Magnetic Bead Panel 96-Well Plate Assay

MGAMMAG-300K

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By purchasing this product, which contains fluorescently labeled microsphere beads authorized by Luminex Corporation ("Luminex"), you, the customer, acquire the right under Luminex's patent rights, if any, to use this product or any portion of this product, including without limitation the microsphere beads contained herein, only with Luminex's laser based fluorescent analytical test instrumentation marketed under the name of Luminex 100[™] IS, 200[™], HTS.

INTRODUCTION

Produced by plasma cells and lymphocytes, immunoglobulins (antibodies) are critically involved in immune response, attaching to antigens and playing a role in their destruction. Immunoglobulins (Ig) can be classified by isotype, classes that differ in function and antigen response due to structure variability. Five major isotypes have been identified in placental mammals: IgM, IgG, IgA, IgE and IgD (B-cell receptor) – all found in normal individuals. Immunoglobulin-deficiency disorders, such as autoimmune disease, some GI conditions and malignancies, are characterized by specific isotype deficiencies or varying concentrations of one or more isotypes. Disease states can range from the absence of one isotype class or subclass to a total deficiency of immunoglobulin classes. In addition, isotyping applications include analyzing hybridomas during antibody development.

EMD Millipore recognizes the need to provide you with the ability to quantitate immunoglobulin classes and subclasses simultaneously. Our MILLIPLEX MAP Mouse Isotyping Kit has been designed to enable you to measure accurately mouse IgG subclasses (1, 2a, 2b, and 3), IgM, and IgA– all in one well. In addition, this configurable kit enables the customer to customize which subclasses are measured. The xMAP multiplex technology is ideal for measuring levels of these isotypes, which not only decreases the number of assays as well as the amount of sample required, but also greatly reduces the possible inaccuracies that result from performing multiple assays.

Magnetic Beads can make the process of automation and high throughput screening easier with features such as walk-away washing. Beyond automation, other advantages of using magnetic bead technology include:

- More flexible plate and plate washer options
- Improved performance with turbid serum/plasma samples
- Assay results equivalent to non-magnetic beads
- Automated washing eliminates technical obstacles (i.e. clogging of wells that contain viscous samples) which may result during vacuum manifold/manual washing

Coupled with the Luminex xMAP® platform in a **magnetic bead** format, you receive the advantage of ideal speed and sensitivity, allowing quantitative multiplex detection of dozens of analytes simultaneously, which can dramatically improve productivity.

EMD Millipore's MILLIPLEX MAP Mouse Immunoglobulin Isotyping Panel can be used for the simultaneous quantification of the following 6 analytes: IgM, IgG1, IgG2a, IgG2b, IgG3 and IgA.

This kit may be used for the analysis of all or any combination of the above analytes in serum and tissue/cell lysate and culture supernatant samples.

For Research Use Only. Not for Use in Diagnostic Procedures.

Please read entire protocol before use.

It is important to use same assay incubation conditions throughout your study.

PRINCIPLE

MILLIPIX® MAP is based on the Luminex® xMAP® technology — one of the fastest growing and most respected multiplex technologies offering applications throughout the life sciences and capable of performing a variety of bioassays including immunoassays on the surface of fluorescent-coded beads known as microspheres.

- Luminex® uses proprietary techniques to internally color-code microspheres with two fluorescent dyes. Through precise concentrations of these dyes, 100 distinctly colored bead sets can be created, each of which is coated with a specific capture antibody.
- After an analyte from a test sample is captured by the bead, a detection antibody conjugated to PE, the reported molecule, is introduced.
- The microspheres are allowed to pass rapidly through a laser which excites the internal dyes marking the microsphere set. A second laser excites PE, the fluorescent dye on the reporter molecule.
- Finally, high-speed digital-signal processors identify each individual microsphere and quantify the result of its bioassay based on fluorescent reporter signals.

The capability of adding multiple conjugated beads to each sample results in the ability to obtain multiple results from each sample. Open-architecture xMAP® technology enables multiplexing of many types of bioassays reducing time, labor and costs over traditional methods.

STORAGE CONDITIONS UPON RECEIPT

- Recommended storage for kit components is 2 - 8°C.
- Once the standards and controls have been reconstituted, immediately transfer contents into polypropylene vials. **DO NOT STORE RECONSTITUTED STANDARDS OR CONTROLS IN GLASS VIALS.** For long-term storage, freeze reconstituted standards and controls at $\leq -20^{\circ}\text{C}$. Avoid multiple (>2) freeze/thaw cycles.
- **DO NOT FREEZE Antibody-Immobilized Beads or PE Antibodies.**

REAGENTS SUPPLIED

Note: Store all reagents at 2 – 8°C

REAGENTS SUPPLIED	CATALOG NUMBER	VOLUME	QUANTITY
MILLIPLEX™ MAP Anti-Mouse κ Light Chain, PE	44-029	50 µL	1 tube
MILLIPLEX™ MAP Anti-Mouse λ Light Chain, PE	44-030	26 µL	1 tube
MILLIPLEX™ MAP Mouse Multi-Immunoglobulin Standard	47-300	0.5 mL	1 vial
MILLIPLEX™ MAP Mouse Immunoglobulin Positive Control	43-008L	0.25 mL	1 vial
MILLIPLEX® MAP Assay Buffer	L-AB	30 mL	1 bottle
MILLIPLEX® MAP Wash Buffer, 10X	L-WB	30 mL	2 bottles
Set of one 96-Well Plate with 2 Sealers	-----	-----	1 plate 2 sealers

Mouse Immunoglobulin Isotyping Antibody-Immobilized Beads:

Bead/Analyte Name	Luminex Bead Region	Customizable 6 Analytes (20X concentration, 200 µL) Available Cat. #	
Anti-Mouse IgA Bead	18	✓	MIGA-MAG
Anti-Mouse IgG1 Bead	21	✓	MIGG1-MAG
Anti-Mouse IgG2a Bead	36	✓	MIGG2A-MAG
Anti-Mouse IgG2b Bead	51	✓	MIGG2B-MAG
Anti-Mouse IgG3 Bead	54	✓	MIGG3-MAG
Anti-Mouse IgM Bead	72	✓	MIGM-MAG

MATERIALS REQUIRED BUT NOT PROVIDED

Reagents

1. Luminex Sheath Fluid (Luminex Catalog #40-50000) or Luminex Drive Fluid (Luminex Catalog # MPXDF-4PK)

Instrumentation / Materials

1. Adjustable Pipettes with Tips capable of delivering 25 μ L to 1000 μ L
2. Multichannel Pipettes capable of delivering 5 μ L to 50 μ L or 25 μ L to 200 μ L
3. Reagent Reservoirs
4. Polypropylene Microfuge Tubes
5. Rubber Bands
6. Aluminum Foil
7. Absorbent Pads
8. Laboratory Vortex Mixer
9. Sonicator (Branson Ultrasonic Cleaner Model #B200 or equivalent)
10. Titer Plate Shaker (Lab-Line Instruments Model #4625 or equivalent)
11. Luminex 200™, HTS, or FLEXMAP 3D™, or MAGPIX® with xPONENT software by Luminex Corporation, or Bio-Plex with Bio-Plex Manager software
12. Automatic Plate washer for magnetic beads (Bio-Tek ELx405, EMD Millipore catalog #40-015 or equivalent) or Hand held Magnetic Separation Block (EMD Millipore catalog # 40-285 or equivalent)

Note: If a plate washer or hand held magnetic separation block for magnetic beads is not available, one can use Microtiter filter plate (MX-PLATE) to run the assay with the use of Vacuum Filtration Unit (EMD Millipore Vacuum Manifold Catalog #MSVMHTS00, or equivalent. EMD Millipore Vacuum Pump Catalog #WP6111560 or equivalent.)

SAFETY PRECAUTIONS

- All blood components and biological materials should be handled as potentially hazardous. Follow universal precautions as established by the Centers for Disease Control and Prevention and by the Occupational Safety and Health Administration when handling and disposing of infectious agents.
- Sodium Azide or Proclin has been added to some reagents as a preservative. Although the concentrations are low, Sodium Azide and Proclin may react with lead and copper plumbing to form highly explosive metal azides. On disposal, flush with a large volume of water to prevent azide build up.

TECHNICAL GUIDELINES

To obtain reliable and reproducible results, the operator should carefully read this entire manual and fully understand all aspects of each assay step before running the assay. The following notes should be reviewed and understood before the assay is set up.

- FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES.
- Do not use beyond the expiration date on the label.
- Do not mix or substitute reagents with those from other lots or sources.
- The Antibody-Immobilized Beads are light-sensitive and must be protected from light at all times. Cover the assay plate containing beads with opaque plate lid or aluminum foil during all incubation steps.
- It is important to allow all reagents to warm to room temperature (20-25°C) before use in the assay.
- Incomplete washing can adversely affect the assay outcome. All washing must be performed with the Wash Buffer provided.
- The Standards prepared by serial dilution must be used within 1 hour of preparation. Discard any unused standards except the standard stock which may be transferred to polypropylene tubes and stored at $\leq -20^{\circ}\text{C}$ for 1 month and at $\leq -80^{\circ}\text{C}$ for greater than one month.
- If samples fall above the dynamic range of the assay, further dilute the samples with the appropriate diluent and repeat the assay.
- During the preparation of the standard curve, make certain to mix the higher concentration well before making the next dilution. Use a new tip with each dilution.
- The plate should be read immediately after the assay is finished. If, however, the plate cannot be read immediately, seal the plate, cover with aluminum foil or an opaque lid, and store the plate at 2-8°C for up to 24 hours. Prior to reading, agitate the plate on the plate shaker at room temperature for 10 minutes.
- The titer plate shaker should be set at a speed to provide maximum orbital mixing without splashing of liquid outside the wells. For the recommended plate shaker, this would be a setting of 5-7, which is approximately 500-800 rpm.
- Ensure that the Luminex needle probe is clean. This may be achieved by sonication and/or alcohol flushes.
- When reading the assay on Luminex 200™, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 4 alignment discs. When reading the assay on FLEXMAP 3D™, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 1 alignment disc. When reading the assay on MAGPIX, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 2 alignment discs.
- For cell culture supernatants, use the culture medium as the diluent in background, standard curve, and control wells. In assays using serum samples, all samples, standards, and controls should be diluted in Assay Buffer. In all cases, use Assay Buffer for resuspension steps.
- Vortex all reagents well before adding to plate.

SAMPLE COLLECTION AND STORAGE

A. Preparation of Serum Samples:

- Allow the blood to clot for at least 30 minutes before centrifugation for 10 minutes at 1000xg. Remove serum and assay immediately or aliquot and store samples at $\leq -20^{\circ}\text{C}$.
- Avoid multiple (>2) freeze/thaw cycles.
- When using frozen samples, it is recommended to thaw the samples completely, mix well by vortexing and centrifuge prior to use in the assay to remove particulates.
- Serum samples should be diluted 1:25,000 in the Assay Buffer and a standard curve diluted in Assay Buffer should be used.
- To achieve a 1:25,000 dilution, dilute 5 μL of sample into 1245 μL of ultrapure water (1:250). Immediately dilute 5 μL of the 1:250 dilution into 495 μL of Assay Buffer (1:100) dilution.
- For data analysis, multiply the final concentration of each sample by the dilution factor.
- If prepared samples yield assay results outside of the standard curve, adjustment of the assay buffer dilution step may be necessary.

B. Preparation of Tissue Culture Supernatant:

- Centrifuge the sample to remove debris and assay immediately or aliquot and store samples at $\leq -20^{\circ}\text{C}$.
- Avoid multiple (>2) freeze/thaw cycles.
- Dilute the sample to approximately 1 $\mu\text{g}/\text{mL}$ Ig in Assay Buffer. [Cell culture supernatants samples approximately (1:5); bioreactor supernatants (1:100)]. Note: Cell culture supernatant concentrations are cell-line dependent and range from 5-50 $\mu\text{g}/\text{mL}$. Bioreactor supernatants may be as concentrated as 1 mg/mL .

NOTE:

- A maximum of 50 μL per well of diluted serum or supernatant can be used.
- All samples must be stored in polypropylene tubes. **DO NOT STORE SAMPLES IN GLASS.**
- Avoid debris, lipids and cells when using samples with gross hemolysis or lipemia.
- Care must be taken when using heparin as an anticoagulant since an excess of heparin will provide falsely high values. Use no more than 10 IU heparin per mL of blood collected.

PREPARATION OF REAGENTS FOR IMMUNOASSAY

A. Preparation of Antibody-Immobilized Beads

Sonicate each antibody-bead vial for 30 seconds; vortex for 1 minute. Add 150 μ L from each antibody bead vial to the Mixing Bottle and bring final volume to 3.0 mL with Assay Buffer. Vortex the mixed beads well. Unused portion may be stored at 2-8°C for up to one month. (Note: Due to the composition of magnetic beads, you may notice a slight color in the bead solution. This does not affect the performance of the beads or the kit.)

Example 1: When using 6 antibody-immobilized beads, add 150 μ L from each of the 6 bead sets to the Mixing Bottle. Then add 2.1 mL Assay Buffer.

B. Preparation of Standards

Resuspend MILLIPLEX MAP Mouse Multi-Immunoglobulin (IgG1, IgG2a, IgG2b, IgG3, IgA, and IgM) Standard (Catalog # 47-300) in 0.5 mL Assay Buffer (Catalog # L-AB). Vortex at high speed for 15 seconds. Place on ice for 15 minutes. This is Standard 7. Note: Standards are of kappa light chain isotype and therefore will react only with anti-Mouse Kappa-PE detection reagent.

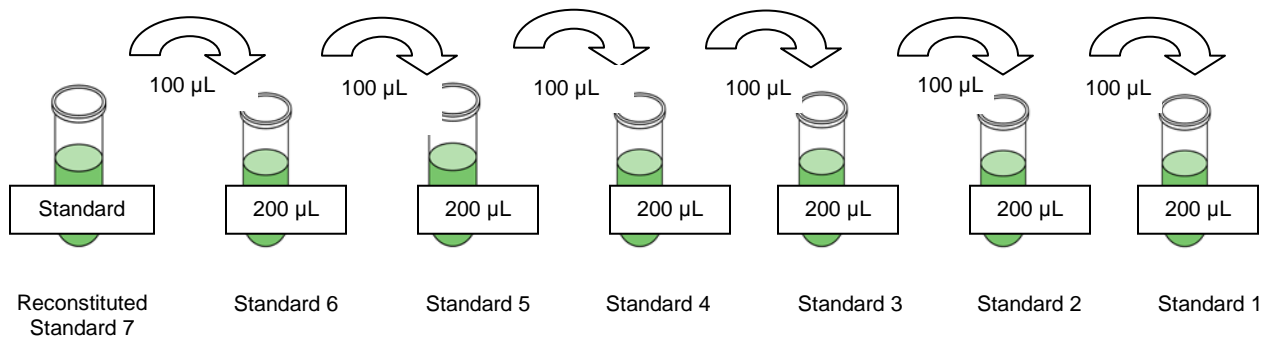
Preparation of Working Standards (Standard):

Label six polypropylene microfuge tubes Standard 6, Standard 5, Standard 4, Standard 3, Standard 2 and Standard 1. Add 200 μ L of Assay Buffer to each of the six tubes. Prepare 3-fold serial dilutions by adding 100 μ L of the reconstituted Standard 7 to the Standard 6 tube, mix well and transfer 100 μ L of the Standard 6 to the Standard 5 tube, mix well and transfer 100 μ L of the Standard 5 to the Standard 4 tube, mix well and transfer 100 μ L of the Standard 4 to Standard 3 tube, mix well and transfer 100 μ L of the Standard 3 to the Standard 2 tube, mix well and transfer 100 μ L of the Standard 2 to the Standard 1 tube and mix well. The Standard 0 (Background) will be Assay Buffer.

Standard	Volume of Assay Buffer to Add	Volume of Standard to Add
Original (Standard 7) (Refer to analysis sheet for exact concentration)	500 μ L	0

Standard	Volume of Assay Buffer to Add	Volume of Standard to Add
Standard 6	200 μ L	100 μ L of Standard 7
Standard 5	200 μ L	100 μ L of Standard 6
Standard 4	200 μ L	100 μ L of Standard 5
Standard 3	200 μ L	100 μ L of Standard 4
Standard 2	200 μ L	100 μ L of Standard 3
Standard 1	200 μ L	100 μ L of Standard 2

Preparation of Working Standards:



C. Preparation of Positive Control

Resuspend MILLIPLEX MAP Mouse Multi-Ig Positive Control (Catalog # 43-008L) in 0.25 mL Assay Buffer (Catalog #L-AB) (or cell culture medium if running with cell culture supernatants). Vortex at high speed for 15 seconds. Place on ice for 15 minutes.

D. Preparation of Detection Reagent

To prepare 100X detection reagent, dilute anti-Mouse κ Light Chain, PE to working solution (1:100) with Assay Buffer (for a full plate, use 25 μ L of the 100X anti-Mouse kappa-PE in 2.475 mL assay buffer). [Note: 95% of mouse antibodies have κ light chains, so in most instances, use of only the κ Light Chain Detection Reagent will be necessary. If determination of light chain is desired or samples show no signal with the κ light chain detection, vacuum plate and prepare anti-Mouse λ , PE the same as above. Repeat detection step with anti-mouse λ , PE and reread plate.

E. Preparation of Wash Buffer

Bring the 10X Wash Buffer to room temperature and mix to bring all salts into solution. Dilute 30 mL of 10X Wash Buffer with 270 mL deionized water. Store unused portion at 2-8°C for up to one month.

IMMUNOASSAY PROCEDURE

- Prior to beginning this assay, it is imperative to read this protocol completely and to thoroughly understand the Technical Guidelines.
- Allow all reagents to warm to room temperature (20-25°C) before use in the assay.
- Diagram the placement of Standards [0 (Background) through 7], Positive Control, and Samples on Well Map Worksheet in a vertical configuration. (Note: Most instruments will only read the 96-well plate vertically by default.) It is recommended to run the assay in duplicate.
- If using a filter plate, see special instruction on the next page.

1. Add 100 μ L Assay Buffer to each well of the Assay Plate. Mix on a plate shaker for 10 minutes at room temperature.
2. Decant Assay Buffer and remove the residual amount from all wells by inverting the plate and tapping it smartly onto absorbent towels several times.
3. Add 50 μ L of Assay Buffer to background wells. Add 50 μ L of standard, control or diluted sample to appropriate wells.
4. Vortex the MILLIPLEX MAP Anti-Mouse Multi-Immunoglobulin Beads at medium speed for 15 seconds, and then sonicate for 15 seconds using a sonication bath. Add 25 μ L of bead solution to each well.
5. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
6. Wash plate 2 times with 200 μ L/well of Wash Buffer, removing Wash Buffer by aspiration between each wash. To avoid washing/aspiration related bead loss, allow approximately 60 seconds between dispensing of the Wash Buffer and subsequent aspiration. Note: If using the recommended plate washer for magnetic beads (Bio-Tek ELx405) follow the appropriate equipment settings outlined on Page 13.
7. Add 25 μ L per well of prepared Anti-Mouse κ Light Chain, PE.
8. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
9. Gently remove fluid by aspiration. (NOTE: DO NOT INVERT PLATE.) To avoid aspiration related bead loss, allow the plate to soak on the magnet of the plate washer for 60 seconds prior to aspiration.
10. Resuspend in 150 μ L/well of Sheath Fluid (or Drive Fluid if using Magpix® instrument).

Add 100 μ L Assay Buffer per well



Shake 10 min, RT
Decant

- Add 50 μ L Assay Buffer, Standard or Control to appropriate wells
- Add 50 μ L prepared samples to sample wells
- Add 25 μ L Beads to each well



Incubate 15 minutes at RT with shaking; dark
Wash plate 2X with 200 μ L Wash Buffer

Add 25 μ L κ light chain, PE solution per well



Incubate 15 minutes at RT with shaking; dark

Aspirate fluid from plate and add 150 μ L Sheath Fluid (or Drive Fluid) per well. Read results using an appropriate Luminex® instrument.

11. Proceed to reading results on an appropriate Luminex[®] instrument.
12. **OPTIONAL:** If no results are observed, or if determination of light chain is desired, aspirate plate and re-assay with Anti-Mouse λ Light Chain, PE.
13. Add 25 μ L Anti-Mouse λ Light Chain, PE per well.
14. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
15. Remove fluid by aspiration and resuspend in 150 μ L/well of Sheath Fluid (or Drive Fluid if using Magpix[®] instrument).
16. Proceed to reading results on an appropriate Luminex[®] instrument.

Continue only with samples where no results were observed



Aspirate Sheath Fluid

Add 25 μ L λ light chain PE solution per well



Incubate 15 minutes at RT with shaking; dark

Aspirate plate and add 150 μ L Sheath Fluid (or Drive Fluid) per well. Read results using an appropriate Luminex[®] instrument.

FILTER PLATE PROCEDURE

* Set the filter plate on a plate holder at all times during reagent dispensing and incubation steps so that the bottom of the plate does not touch any surface.

1. Pre-wet filter plate by pipetting 25 μ L of Assay Buffer into each well. Seal and mix on a plate shaker for 10 minutes at room temperature. Remove Assay Buffer by vacuum. Blot bottom of plate with absorbent pad or paper towel.
2. Add 50 μ L of Assay Buffer to background wells. Add 50 μ L of standard, control or diluted sample to appropriate wells.
3. Vortex the MILLIPLEX MAP Anti-Mouse Multi-Immunoglobulin Beads at medium speed for 15 seconds, and then sonicate for 15 seconds using a sonication bath. Add 25 μ L of bead solution to each well.
4. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
5. Remove fluid by vacuum. Wash plate 2 times with 150 μ L/well of Wash Buffer, removing Wash Buffer by vacuum filtration between each wash. Blot bottom of plate with absorbent pad or paper towel. **Do not over-dry.**
6. Add 25 μ L per well of diluted Anti-Mouse κ Light Chain, PE.
7. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
8. Remove fluid by vacuum. Blot bottom of plate with absorbent pad or paper towel and resuspend in 150 μ L/well of Sheath Fluid.
9. Proceed to reading results on an appropriate Luminex[®] instrument.
10. **OPTIONAL:** If no results are observed, or if determination of light chain is desired, vacuum plate and re-assay with Anti-Mouse λ Light Chain, PE.
11. Add 25 μ L Anti-Mouse λ Light Chain, PE per well.
12. Cover with opaque plate cover and incubate 15 minutes with agitation on plate shaker at room temperature.
13. Remove fluid by vacuum and resuspend in 150 μ L/well of Sheath Fluid (or Drive Fluid if using Magpix[®] instrument).
14. Proceed to reading results on an appropriate Luminex[®] instrument.

Add 25 μ L Assay Buffer per well



Incubate 10 mins at RT
Vacuum Plate

- Add 50 μ L Assay Buffer, Standard or Control to appropriate wells
- Add 50 μ L diluted samples to sample wells
- Add 25 μ L Beads to each well



Incubate 15 minutes at RT with shaking; dark
Vacuum and wash 2X with 150 μ L Wash Buffer

Add 25 μ L κ light chain, PE solution per well



Incubate 15 minutes at RT with shaking; dark

Vacuum plate and add 150 μ L Sheath Fluid (or Drive Fluid) per well. Read results using an appropriate Luminex[®] instrument.

Continue only with samples where no results were observed



Vacuum plate

Add 25 μ L λ light chain PE solution per well



Incubate 15 minutes at RT with shaking; dark

Aspirate plate and add 150 μ L Sheath Fluid (or Drive Fluid) per well. Read results using an appropriate Luminex[®] instrument.

PLATE WASHING

1.) Solid Plate

If using a solid plate, use either a hand-held magnet or magnetic plate washer.

- A.) For hand-held magnet, rest plate on magnet for 60 seconds to allow complete settling of magnetic beads. Remove well contents by gently decanting the plate in an appropriate waste receptacle and gently tapping on absorbent pads to remove residual liquid. Wash plate with 200 μ L of Wash Buffer by removing plate from magnet, adding Wash Buffer, shaking for 30 seconds, re-attaching to magnet, letting beads settle for 60 seconds and removing well contents as previously described after each wash. Repeat wash steps as recommended in Assay Procedure.
- B.) For magnetic plate washer, let plate “soak” on magnet for 60 seconds to allow complete settling of the magnetic beads. Remove well contents by aspiration. Wash plate 2 times with 200 μ L/well of Wash Buffer, letting beads “soak” for 60 seconds and removing Wash Buffer by aspiration after each wash. Repeat wash steps as recommended in Assay Procedure. **Note:** If using the recommended plate washer for magnetic beads (Bio-Tek ELx405) follow the appropriate equipment settings outlined on Page 14, see **EQUIPMENT SETTINGS**.

2.) Filter Plate (EMD Millipore Cat# MX-PLATE)

If using a filter plate, use a vacuum filtration manifold to remove well contents. Wash plate 200 μ L/well of Wash Buffer, removing Wash Buffer by vacuum filtration after each wash. Repeat wash steps as recommended in the Assay Procedure.

EQUIPMENT SETTINGS (Plate Reader)

These specifications are for the Luminex 200™ xPONENT™, FlexMAP 3D™, MAGPIX® and Luminex HTS. Luminex instruments with other software (e.g. MasterPlex, StarStation, LiquiChip, Bio-Plex, LABScan100) would need to follow instrument instructions for gate settings and additional specifications from the vendors for reading Luminex Magnetic Beads.

For magnetic bead assays, the Luminex 200™ and HTS instruments must be calibrated with the xPonent 3.1 compatible Calibration Kit (EMD Millipore Cat.# 40-275) and performance verified by Verification Kit (EMD Millipore Cat. # 40-276). The Luminex FlexMAP 3D™ instrument must be calibrated with the FlexMAP 3D™ Calibration Kit (EMD Millipore Cat# 40-028) and performance verified with the FlexMAP 3D™ Performance Verification Kit (EMD Millipore Cat# 40-029). The Luminex MAGPIX® instrument must be calibrated with the MAGPIX® Calibration Kit (EMD Millipore Cat# 40-049) and performance verified with the MAGPIX® Performance Verification Kit (EMD Millipore Cat# 40-050).

NOTE: These assays cannot be run on Luminex 100™ instruments or any instruments using the Luminex IS 2.3 or Luminex 1.7 software.

The Luminex probe height must be adjusted to the plate provided in the kit. Please use Cat.# MAG-PLATE, if additional plates are required for this purpose.

Events	50 per bead
Sample Size	100 μ L
Gate Settings	8,000 to 15,000
Reporter Gain	Default (Low PMT)
Time Out	60 seconds
Bead Set	Bead #
IgA	18
IgG1	21
IgG2a	36
IgG2b	51
IgG3	54
IgM	72

EQUIPMENT SETTINGS (Plate Washer)

Bio-Tek ELx405:

The general recommended wash protocol (Link Protocol) is as follows:

Soak Program: Wash Program:

Soak _ Aspirate _ Dispense _ Soak _ Aspirate _ Dispense _ Soak _ Aspirate

1.) Soak program:

1. Soak duration: 60 sec
2. Shake before soak?: NO

2.) Wash program:

Method:

1. Number of cycles: 2
2. Soak/shake: YES
3. Soak duration: 60 sec
4. Shake before soak: NO
5. Prime after soak: NO

Dispense:

1. Dispense volume: 200 μ L/well
2. Dispense flow rate: 5
3. Dispense height: 130 (16.51 mm)
4. Horizontal disp pos: 00 (0 mm)
5. Bottom Wash first?: NO
6. Prime before start?: NO

Aspiration:

1. Aspirate height: 35 (4.445 mm)
2. Horizontal Asp Pos: 30 (1.372 mm)
3. Aspiration rate: 06 (15.0 mm/sec)
4. Aspiration delay: 0
5. Crosswise Aspir: NO
6. Final Aspir: YES
7. Final Aspir delay: 0 (0 msec)

3.) Link program: (Note: This is the program to use during actual plate washing.)

Link together the Soak and Wash programs outlined above.

Note: After the final aspiration, there will be approximately 25 μ L of residual Wash Buffer in each well. This is expected when using the BioTek Plate washer and this volume does not need to be aspirated from the plate.

If using an automatic plate washer other than BioTek ELx405, please refer to the manufacturers's recommendations for programming instructions.

QUALITY CONTROLS

The ranges for each analyte in the Mouse Multi-Ig Positive Control are provided on the card insert or can be located at the EMD MILLIPORE website

www.millipore.com/techlibrary/index.do using the catalog number as the keyword.

ASSAY CHARACTERISTICS

Assay Sensitivities (minimum detectable concentrations, ng/mL)

MinDC: Minimum Detectable Concentration is calculated using Milliplex Analyst Software from EMD Millipore. It measures the true limits of detection for an assay by mathematically determining what the empirical MinDC would be if an infinite number of standard concentrations were run for the assay under the same conditions.

Analyte	MinDC + 2SD
IgA	1.1
IgG1	0.27
IgG2a	0.67
IgG2b	0.18
IgG3	0.32
IgM	1.73

N=4 assays

Precision (%CV)

Intra-assay precision is generated from the mean of the %CV's from four reportable results across 1 concentration of analytes in one assay. Inter-assay precision is generated from the mean of the %CV's from four reportable results across 1 concentration of analytes across 4 different assays.

Analyte	Intra-Assay (%CV)	Inter-Assay (%CV)
IgA	10	17
IgG1	6	11
IgG2a	8	14
IgG2b	7	16
IgG3	5	13
IgM	5	14

Accuracy (% Recovery)

Spike Recovery: The data represent mean percent recovery of 4 levels of spiked standards in diluted serum from 4 different rat samples.

<i>Isotype</i>	<i>Spike Recovery in Serum</i>
IgA	98%
IgG1	96%
IgG2a	76%
IgG2b	98%
IgG3	85%
IgM	100%

TROUBLESHOOTING GUIDE

Problem	Probable Cause	Solution
Insufficient bead count	<p data-bbox="451 184 803 239">Plate Washer aspirate height set too low</p> <p data-bbox="451 275 803 302">Bead mix prepared incorrectly</p> <p data-bbox="451 428 803 516">Samples cause interference due to particulate matter or viscosity</p> <p data-bbox="451 579 803 634">Probe height not adjusted correctly</p>	<p data-bbox="836 184 1373 239">Adjust aspiration height according to manufacturers' instructions.</p> <p data-bbox="836 275 1373 392">Sonicate bead vials and vortex just prior to adding to bead mix bottle according to protocol. Agitate bead mix intermittently in reservoir while pipetting into the plate.</p> <p data-bbox="836 428 1373 546">See above. Also sample probe may need to be cleaned with alcohol flush, backflush and washes; or, if needed, probe should be removed and sonicated.</p> <p data-bbox="836 579 1373 940">When reading the assay on Luminex 200™, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 4 alignment discs. When reading the assay on FLEXMAP 3D™, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 1 alignment disc. When reading the assay on MAGPIX, adjust probe height according to the protocols recommended by Luminex to the kit solid plate using 2 alignment discs.</p>
Background is too high	<p data-bbox="451 976 803 1031">Background wells were contaminated</p> <p data-bbox="451 1129 803 1184">Matrix used has endogenous analyte or interference</p> <p data-bbox="451 1247 803 1274">Insufficient washes</p>	<p data-bbox="836 976 1373 1094">Avoid cross-well contamination by using sealer appropriately and by pipeting with multichannel pipets without touching reagent in plate.</p> <p data-bbox="836 1129 1373 1213">Check matrix ingredients for cross reacting components (e.g. interleukin modified tissue culture medium).</p> <p data-bbox="836 1247 1373 1274">Increase number of washes.</p>
Beads not in region or gate	<p data-bbox="451 1318 803 1373">Luminex not calibrated correctly or recently</p> <p data-bbox="451 1436 803 1491">Gate settings not adjusted correctly</p> <p data-bbox="451 1589 803 1644">Wrong bead regions in protocol template</p> <p data-bbox="451 1680 803 1707">Incorrect sample type used</p> <p data-bbox="451 1803 803 1858">Instrument not washed or primed</p>	<p data-bbox="836 1318 1373 1402">Calibrate Luminex based on instrument manufacturer's instructions at least once a week or if temperature has changed by >3°C.</p> <p data-bbox="836 1436 1373 1554">Some Luminex instruments (e.g. Bio-Plex) require different gate settings than those described in the kit protocol. Use instrument default settings.</p> <p data-bbox="836 1589 1373 1644">Check kit protocol for correct bead regions or analyte selection.</p> <p data-bbox="836 1680 1373 1764">Samples containing organic solvents or if highly viscous should be diluted or dialyzed as required.</p> <p data-bbox="836 1803 1373 1921">Prime the Luminex 4 times to eliminate air bubbles. Wash 4 times with sheath fluid or water if there is any remnant alcohol or sanitizing liquid.</p>

Problem	Probable Cause	Solution
	Beads were exposed to light	Keep plate and bead mix covered with dark lid or aluminum foil during all incubation steps.
Signal for whole plate is same as background	Incorrect or no Detection Antibody was added Streptavidin-Phycoerythrin was not added	Add appropriate Detection Antibody and continue. Add Streptavidin-Phycoerythrin according to protocol. If Detection Antibody has already been removed, sensitivity may be low.
Low signal for standard curve	Detection Antibody may have been removed prior to adding Streptavidin Phycoerythrin Incubations done at incorrect temperatures, timings or agitation	May need to repeat assay if desired sensitivity not achieved. Assay conditions need to be checked.
Signals too high, standard curves are saturated	Calibration target value set too high Plate incubation was too long with standard curve and samples	With some Luminex instruments (e.g. Bio-Plex) default target setting for RP1 calibrator is set at High PMT. Use low target value for calibration and reanalyze plate. Use shorter incubation time.
Sample readings are out of range	Samples contain no or below detectable levels of analyte Samples contain analyte concentrations higher than highest standard point Standard curve was saturated at higher end of curve	If below detectable levels, it may be possible to use higher sample volume. Check with tech support for appropriate protocol modifications. Samples may require dilution and reanalysis for that particular analyte. See above.
High variation in samples and/or standards	Multichannel pipet may not be calibrated Plate washing was not uniform Samples may have high particulate matter or other interfering substances Plate agitation was insufficient Cross-well contamination	Calibrate pipets. Confirm all reagents are removed completely in all wash steps. See above. Plate should be agitated during all incubation steps using a vertical plate shaker at a speed where beads are in constant motion without splashing. Check when reusing plate sealer that no reagent has touched sealer.

Problem	Probable Cause	Solution
High variation in samples and/or standards (cont.)		Care should be taken when using same pipet tips that are used for reagent additions and that pipet tip does not touch reagent in plate.

FOR FILTER PLATES ONLY

Filter plate will not vacuum	<p>Vacuum pressure is insufficient</p> <p>Samples have insoluble particles</p> <p>High lipid concentration</p>	<p>Increase vacuum pressure such that 0.2mL buffer can be suctioned in 3-5 seconds.</p> <p>Centrifuge samples just prior to assay set-up and use supernatant.</p> <p>After centrifugation, remove lipid layer and use supernatant.</p>
Plate leaked	<p>Vacuum Pressure too high</p> <p>Plate set directly on table or absorbent towels during incubations or reagent additions</p> <p>Insufficient blotting of filter plate bottom causing wicking</p> <p>Pipette touching plate filter during additions</p> <p>Probe height not adjusted correctly</p> <p>Sample too viscous</p>	<p>Adjust vacuum pressure such that 0.2mL buffer can be suctioned in 3-5 seconds. May need to transfer contents to a new (blocked) plate and continue.</p> <p>Set plate on plate holder or raised edge so bottom of filter is not touching any surface.</p> <p>Blot the bottom of the filter plate well with absorbent towels after each wash step.</p> <p>Pipette to the side of plate.</p> <p>Adjust probe to 3 alignment discs in well H6.</p> <p>May need to dilute sample.</p>

REPLACEMENT REAGENTS**Catalog #**

MILLIPLEX™ MAP Anti-Mouse κ Light Chain, PE	44-029
MILLIPLEX™ MAP Anti-Mouse λ Light Chain, PE	44-030
MILLIPLEX™ MAP Mouse Multi-Immunoglobulin Standard	47-300
MILLIPLEX™ MAP Mouse Immunoglobulin Positive Control	43-008L
Wash Buffer	L-WB
Assay Buffer	L-AB
Set of two 96-Well Plates with 4 Sealers	MAG-PLATE

Antibody-Immobilized Magnetic Beads

<u>Analytes</u>	<u>Bead #</u>	<u>Cat. #</u>
Anti-Mouse IgA	18	MIGA-MAG
Anti-Mouse IgG1	21	MIGG1-MAG
Anti-Mouse IgG2a	36	MIGG2A-MAG
Anti-Mouse IgG2b	51	MIGG2B-MAG
Anti-Mouse IgG3	54	MIGG3-MAG
Anti-Mouse IgM	72	MIGM-MAG

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WELL MAP

	1	2	3	4	5	6	7	8	9	10	11	12
A	Standard 0 Background	Standard 4	Pos Control									
B	Standard 0 Background	Standard 4	Pos Control									
C	Standard 1	Standard 5										
D	Standard 1	Standard 5										
E	Standard 2	Standard 6										
F	Standard 2	Standard 6										
G	Standard 3	Standard 7										
H	Standard 3	Standard 7										