

The Power of Biomarker Analysis

Multiplex assay tips from the makers of MILLIPLEX[®] MAP



Why just multiplex when you can MILLIPLEX®?

For over ten years, Merck Millipore has offered the benefits of MILLIPLEX® MAP multiplexed assay panels—containing all the components and reagents you need to detect multiple analytes simultaneously. The benefits of multiplex protein detection assays are endless, but navigating a protocol can be challenging. We're so confident in the benefits of MILLIPLEX® MAP kits that we've compiled this book of tips and tricks, straight from the experts, to eliminate any doubt in your ability to multiplex like a pro.

Every year, thousands of your colleagues experience the benefits of MILLIPLEX® MAP kits, publishing in scientific journals around the world. We hope this guide enhances the power of your research with multiplexing.



NOTE: Alternate methods presented in this guide may deviate from the protocol. These methods have either been tried by Merck Millipore or our end users using our MILLIPLEX® MAP kits. We cannot guarantee methods presented will work in all cases. These procedures have not been validated.

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Introduction

The Luminex xMAP® Technology:

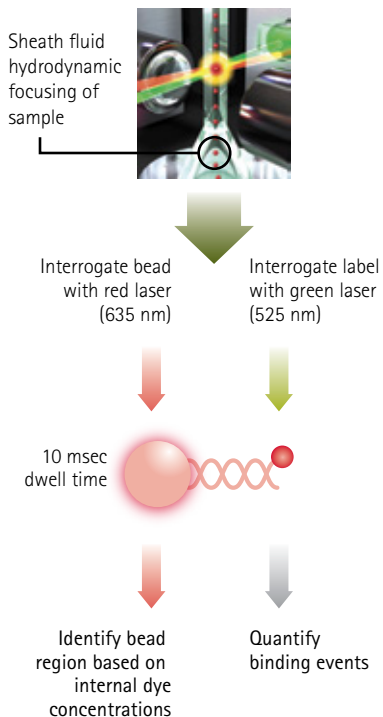
- MILLIPLEX® MAP is based on the Luminex xMAP® bead-based assay platform—one of the fastest-growing and most respected multiplex technologies, supporting applications throughout the life sciences. This platform is capable of performing a variety of bioassays, including immunoassays, on the surface of fluorescent-coded magnetic (MagPlex®) and nonmagnetic (MicroPlex®) bead microspheres.
- Luminex® uses proprietary techniques to internally color-code microspheres with multiple fluorescent dyes. Through precise concentrations of these dyes, distinctly colored bead sets of 500 5.6 µm non-magnetic or 80 6.45 µm magnetic polystyrene microspheres can be created, each of which is coated with a specific capture antibody.
- After the target protein from a test sample is captured by the bead, a biotinylated detection antibody is introduced.
- The reaction mixture is then incubated with Streptavidin-PE conjugate, the reporter molecule, to complete the reaction on the surface of each microsphere.
- Merck Millipore provides three Luminex® instruments to acquire and analyze data using two detection methods (see Figure 1):
- The Luminex 200™ and FLEXMAP 3D® systems are flow cytometry-based instruments that integrate key xMAP® detection components, such as lasers, optics, advanced fluidics and high-speed digital signal processors.
- The MAGPIX® analyzer is a CCD-based instrument that integrates key xMAP® capture and detection components with the speed and efficiency of magnetic bead processing.
- Each individual microsphere is identified by its "bead signature", and the result of its bioassay is quantified based on fluorescent reporter signals. Merck Millipore combines the streamlined data acquisition power of Luminex® xPONENT® acquisition software with sophisticated analysis capabilities of the new MILLIPLEX® Analyst 5.1 software, integrating data acquisition and analysis seamlessly with all Luminex® instruments.

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

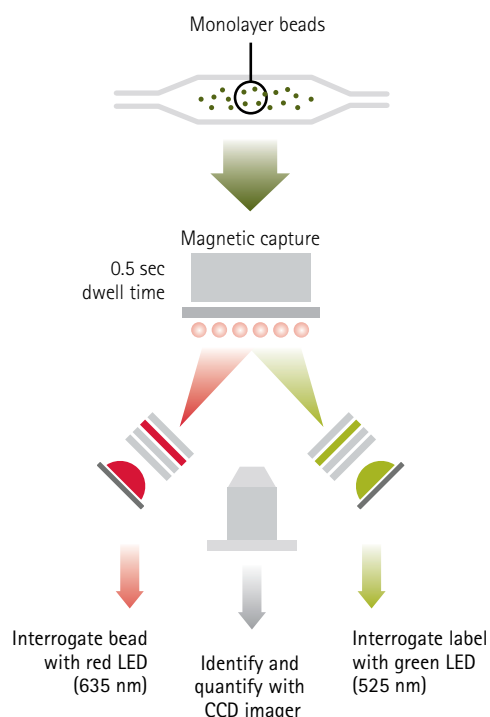
Figure 1.

Two different fluorescence detection methods for acquiring and analyzing bead-based assay data.

Flow cytometry-based analysis



LED-based analysis



MILLIPLEX[®] MAP: Bring your biomarkers to life

MILLIPLEX[®] MAP kits offer multiplex detection of biomarkers in key research focus areas:

- Bone Metabolism
- Cancer Biomarkers
- Cardiovascular Disease
- Immunology/Immune Response
- Intracellular
 - Cell Signaling
 - Cellular Metabolism
- Metabolism/Endocrinology
- Neuroscience
- Toxicity

Our kits offer:

- The broadest selection of analytes across a wide range of disease states for both circulating and intracellular biomarkers.
- All the components and reagents you need to detect multiple analytes simultaneously.
- Quality controls provided to qualify assay performance.
- Analytically validated panels yielding consistent analyte profiles within panels.
- Comparison of standard and QC lots to a reference lot to ensure lot-to-lot consistency.
- Panels that meet stringent manufacturing criteria to ensure batch-to-batch reproducibility.



Deciding Which MILLIPLEX® Assay is Best for Your Research

- Merck Millipore's ELISAs and MILLIPLEX® assays for the same analyte commonly use the same antibody pairs and conditions.
 - In Method Comparison tests, while the absolute values are not exactly the same, the results do correlate. Hence when switching from one assay platform to another, a correlation factor may be used when comparing with past data. In most situations the dynamic range and the sensitivity of the assay may be better with the MILLIPLEX® kits.
 - Please contact Technical Support for more information on correlation factors.

- To locate protocols and technical documents for a specific panel:
 - Search the website for the catalog number.
 - The link to the protocol can be found using:
 - The "Documents" tab.
 - The right-hand side of the product name and catalog number.
 - The top of the Product Description page.

- The easiest way to find a panel that contains the analytes you want to measure would be to search online:
 - The MILLIPLEX® Analyte Kit Finder located on the MILLIPLEX® home page.
 - Search the latest edition of the Analyte Quarterly: www.merckmillipore.com/milliplex
- To find publications citing a specific panel or analyte:
 - Search the website for the catalog number. Links to some of the references using the product can be found at the top of the Product Description page.
 - For a more complete list, contact Technical Support.

- To determine cross-reactivity for other species for a panel or analyte:
 - See the Species Cross-reactivity Tables in Appendix 1.
 - For intracellular assay kits, we analytically validate the assay with human cell/tissue culture samples. However, we provide the species homology for each analyte in a table on the product detail page on our website. We have also compiled a list of this information on our website:
 - Kits: Kit Species Cross Reactivity: www.merckmillipore.com/kits_species
 - MAPmate™ assays: MAPmate™ Species Cross Reactivity & Buffer Table: www.merckmillipore.com/mapmates_sp
 - Search the latest edition of the Analyte Quarterly: www.merckmillipore.com/milliplex



General Assay Information

- All kits are for Research Use Only.

 - Always read the entire protocol before proceeding.

 - For questions or issues with Luminex® instruments, contact Luminex® at:
 - **All Regions:**
www.merckmillipore.com/lmx_contact
 - **Technical Support:**
Phone: 512-381-4397
Toll-free: 1-877-785-2323
Fax: 512-219-5114
 - **Email:** support@luminexcorp.com

 - For questions or issues with BioTek washers, contact BioTek at:
 - **All Regions:**
www.merckmillipore.com/biotek_contact
 - **Technical Support:**
In North America Call:
(800) 242-4685
Outside the U.S. Call:
(802) 655-4740
 - **Email:** TAC@biotek.com

 - For questions or issues with MILLIPLEX® assays, please contact Merck Millipore Technical Support or your Sales Specialist.
- How to design a "customized" kit:
 - Select your panel of interest: for example, Human Cytokine/Chemokine Panel 1 (Cat. No. HCYTOMAG-60K).
 - Choose only the analytes you want from that panel: for example, you may need only five analytes: IL-2, IL-6, IL-10, GM-CSF, VEGF-A.
 - Add the number of analytes you chose to the catalog number: HCYTOMAG-60K-05 and list the specific analytes.

 - How to design and order a customized kit online:
 - From the Product Description page:
 - Click "Design & Purchase Your Own Kit".
 - Make your choices; add to the cart and/or save to your favorites and go to "Checkout."
 - From the MILLIPLEX® MAP website:
 - www.merckmillipore.com/milliplex
 - Click "Design & Purchase Your Own Kit".
 - Make your choices; add to the cart and/or save to your favorites and go to "Checkout."
 - In "Quick Purchase":
 - Click on the "Begin" icon located within "Design and Purchase your MILLIPLEX® MAP Kits."
 - Make your choices; add to the cart and/or save to your favorites and go to "Checkout."

 - Do not use a kit beyond its expiration date.

 - The expiration date for a kit is that of the component with the shortest expiration date. This date is printed on the box label.

 - Kits will ship with a minimum of 3 months until expiration.

 - Longer expiration dates can be requested. Please contact your Sales Specialist.



Materials Required But Not Provided

- Adjustable pipettes with tips capable of delivering 25 μ L to 1000 μ L.

 - Multichannel pipettes capable of delivering 5 μ L to 50 μ L or 25 μ L to 200 μ L.

 - Laboratory vortex mixer.

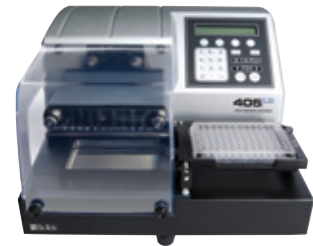
 - Water bath sonicator (Branson Ultrasonic Cleaner Model #B200 or equivalent).
 - Sonicator probes are not recommended.

 - Orbital titer plate shaker (Lab-Line Instruments Model #4625 or equivalent).
 - Very important: For incubating assays overnight a power supply must be available for the orbital shaker in a refrigerator or cold room.
 - The orbital titer plate shaker should be set at a speed to provide maximum orbital mixing without the 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.
 - Note that orbital shakers vary. Your shaker can be calibrated by pre-wetting the plate with buffer and slowly increasing the speed until splashing occurs. Then lower the speed slightly. The shaker should be set at the highest speed allowable without splashing the liquid.

 - Luminex 200™, MAGPIX®, or FLEXMAP 3D® analyzer instruments, available from Merck Millipore and manufactured by Luminex® Corporation.

 - Sheath fluid (Luminex 200™ or FLEXMAP 3D® systems) or drive fluid (MAGPIX® instrument)
 - Sheath fluid and drive fluid are specific for use in the appropriate instrument and cannot be used interchangeably.
 - Sheath fluid or drive fluid can be reordered directly from Merck Millipore:
 - Sheath Fluid (Cat. No. SHEATHFLUID)
 - MAGPIX® Drive Fluid 4PK (Cat. No. MPXDF-4PK)
- Before you open a MILLIPLEX® MAP kit, check your instrument.
 - Is the instrument calibrated?
 - Has it been maintained?

 - Bead washer (either automated or manual):
 - Automated magnetic bead plate washers:
 - BioTek® 405 LS Magnetic 96-23II Washer (Cat. No. 40-0944).
 - BioTek® 405 LS Magnetic/Vacuum Filtration 96-well Washer (Cat. No. 40-095).
 - BioTek® 405 TS Magnetic 96-well Washer Complete with Touch Screen and Ultrasonic Cleaning (Cat. No. 40-096).
 - BioTek® 405 TS Magnetic/Vacuum Filtration 96-well Washer Complete with Touch Screen and Ultrasonic Cleaning (Cat. No. 40-097).
 - Please see our Luminex® Technology Brochure (Lit. No. DS0012ENEU) for more information.
 - Handheld Magnetic Separator Block for 96-well Flat Bottom or Conical Well Plates (Cat. No. 40-285).



BioTek® plate washer model 405 LS (top) and models 40-094 and 40-095 (bottom).



Sample Collection and Preparation

General Information

- Proper and consistent pipetting technique is key to consistent and accurate data, especially if multiple users will be generating data in collaboration. Improper or inconsistent technique can affect delivery volumes and impact data reliability. Training or best practices for pipetting, including that pipettors are properly calibrated can substantially increase pipetting precision. For tips on proper pipetting techniques visit: www.merckmillipore.com/mlo
- Before you collect samples to run an assay, it's important to read the entire protocol. If you have any questions, contact Technical Support or your Sales Specialist.
- If the protocol states that the kit can be used in either serum or plasma, and you have the option, choose serum because it tends to be cleaner.
- Be consistent with the use of sample types within a study/project.
- If you are trying to decide whether to collect serum or plasma samples, ask yourself what you have observed from preliminary data, publications or collaborators.
 - Still unsure? Contact Technical Support.
- Freeze/thaw limits:
 - Multiple freeze/thaw cycles may reduce the stability of the analytes.
- Vortexing samples and beads:
 - To get a homogeneous prep, sample bead vortexing is recommended, especially after the sample has been centrifuged and the supernatant separated.
- Tips on using tissue culture media as assay buffer:
 - If the medium used to grow the cells is used as the matrix, take care that there are no proteases or any supplements that may interfere with the assay or generate inaccurate results (e.g., cytokines, human serum, etc.).
- Some kits for metabolism biomarkers require adding protease inhibitors to samples. Others may require a sample extraction or acidification.
 - Consult the protocol of the appropriate kit.
 - See Sample Preparation outlines for kits requiring sample extraction, acidification, etc. in Appendix 2.

Preparation of Serum/Plasma Samples

- For serum or plasma samples that require a dilution instead of "Neat", use the serum matrix provided in the kit as the diluents.
- Hemolysis can result in increased proteolytic activity and analyte degradation primarily due to enzymes released from lysed cells.
- Trace hemolysis in samples collected with protease inhibitors may be acceptable, but gross hemolysis will probably interfere with assay performance.
- Hemoglobin (at >10 mg/mL) is known to interfere with antigen/antibody interactions.
- If you want to run a MILLIPLEX® kit using samples other than serum or plasma, we have protocols to address tissue lysates, urine, blood spots, gingival fluid, nasal lavage fluid, tears, cerebrospinal fluid (CSF), bronchoalveolar fluid, saliva and cervical/vaginal secretions, as well as protocols that are modified for use with small volume samples.
 - Please contact Technical Support or refer to Appendix 3.

Preparation of Tissue Culture Supernatants

- For cell culture supernatants or tissue extracts, use the culture or extraction medium as the matrix solution in the blank, standard curves and controls.
 - If samples are diluted in assay buffer, use the assay buffer as the matrix.

- For cell/tissue homogenates, the final cell or tissue homogenate should be prepared in a buffer that has a neutral pH, contains minimal detergents or strongly denaturing agents and has an ionic strength close to physiological concentrations.
 - Avoid debris, lipids, and cell/tissue "aggregates."
 - Centrifuge samples before use.

Preparation of Cell Lysates for Intracellular Assays in 96-well Plates

- For adherent cell lines: seed ~40,000 cells/well and allow growth for 48 hours.
- For suspension cell lines: seed ~250,000 cells/well and collect at desired time.
- For cell lysis: add 30 μ L lysis buffer per well and pipet up and down thoroughly without creating too many bubbles. For a more detailed protocol, request information from Technical Support.

- Add protease inhibitors and/or phosphatase inhibitors to "home-brew" lysis buffers.

- Lysis buffer selection:
 - Lysis buffer can be found in the Cell Signaling Buffer & Detection Kit (Cat. No. 48-602MAG) or sold separately (Cat. No. 43-040).
 - Non-ionic detergents (NP40, Tergitol, IPEGAL) are recommended in lysis buffers for solubilizing cytoplasmic proteins.
 - Partially ionic detergents (Triton® X-100) are recommended in lysis buffers for cytoplasmic or membrane-bound proteins.
 - Ionic detergents (sodium dodecyl sulfate, SDS) are recommended in lysis buffers for membrane-bound, nuclear or mitochondrial proteins. If using SDS in the lysis buffer (i.e., Radioimmunoprecipitation assay (RIPA) buffer), then cell lysate must be diluted to less than 0.05% SDS for assays to detect intracellular proteins, such as cell signaling proteins.

- NOTE: to solubilize nuclear/mitochondrial proteins, you must use either SDS or another method (such as ultrasonication) to puncture the tough nuclear/mitochondrial membranes.
- Reducing agents, like β -mercaptoethanol or dithiothreitol, are not recommended.

- For more information about the compatibility of buffers with MILLIPLEX® MAP Signaling kits, contact Technical Support.

- Perform all dilutions with lysis buffer (not assay buffer or phosphate-buffered saline (PBS)).

Table 1. Detergent compatibility with MILLIPLEX® intracellular assays.

Type of detergents	Protein localization	Maximum allowed protein concentration	MILLIPLEX® assay compatibility
Non-ionic detergents	Cytoplasm	5 mg/mL	Yes
Partially ionic detergents	Cytoplasm, Membrane-bound	5 mg/mL	Yes
Ionic detergents	Membrane-bound, Nucleus, Mitochondria	5 mg/mL	Requires dilution

- Total protein concentration limits*:
 - Do not collect lysates at greater than 5 mg/mL protein concentration.
 - At protein concentrations higher than 5 mg/mL, not all proteins will be solubilized equally by the lysis buffer. Some proteins can be solubilized at a given detergent concentration, while other proteins are not as affected. For example, β -tubulin signal decreases with increasing total protein concentration (signal decrease occurs at 5 to 6 mg/mL for Jurkat cell and peripheral blood mononuclear cell (PBMC) lysates).
 - Total protein concentrations should be within a specific range, which is outlined in each protocol.
 - A starting protein amount is 10 μ g per well (10 μ g protein in the final 25 μ L that is loaded into each assay well) is recommended.
 - Working backwards from there:
 - 10 μ g/25 μ L = 0.4 μ g/ μ L (mg/mL).
 - Diluting the cell/tissue lysates 1:1 in the assay buffer provided in the intracellular kit is recommended.
 - Consequently, all samples need to be brought to a protein concentration of 0.8 μ g/ μ L in lysis buffer.
 - Take 30 μ L of each lysate sample and add it to 30 μ L of assay buffer, bringing the final concentration down to 0.4 mg/mL.
 - Then load 25 μ L of diluted samples into each well, usually in duplicate.

*The Direct Detect® spectrometer enables you to determine protein sample concentrations quickly and accurately. For more information visit: www.merckmillipore.com/directdetect

Preparation of Reagents

General Information

- Deliver extremely precise volumes of solvent when reconstituting lyophilized products. Variations of even a few microliters will significantly affect quantitation.
- Do not mix or substitute assay reagents with those from other lots or sources.
- If leftover reagent lots match and the reagents have been kept at the appropriate storage conditions, they can be used in combination until the expiration dates.
- Serum matrix, bead diluents and wash and assay buffers from other kits can be used/combined if the catalog numbers of these components match in the protocols for the kits in question.

Use of Bead Diluent

- Approximately 10% of a normal population of samples, especially human serum or plasma samples, has heterophilic antibodies that can nonspecifically bind to the capture and detection antibodies simultaneously, thus generating a false positive signal.
 - Bead diluents contain a cocktail of proprietary reagents that significantly reduce this false signal without reducing the true analyte measurement.
 - Bead diluents may also contain factors for detection (e.g., insulin in mouse kits).
- If assay buffer was erroneously added instead of bead diluent, transfer samples to a clean, clear centrifuge tube, spin down and remove buffer; replace with bead diluents and proceed with the assay protocol.

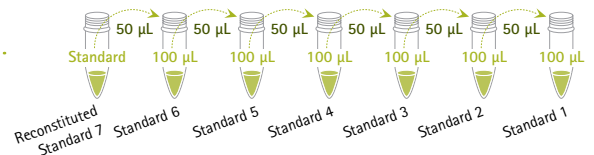
Antibody-Immobilized Beads

- The antibody-immobilized beads are light-sensitive and must be protected from light at all times.
 - Cover the assay plate containing beads with an opaque plate lid or aluminum foil during all incubation steps.
- Any unused mixed antibody-immobilized beads may be stored in the Bead Mix Bottle at 2-8 °C for up to one month.

Standards

- After hydration/reconstitution, all standards and controls must be transferred to polypropylene tubes.
- During the preparation of standard curves, thoroughly mix each higher concentration before making the next dilution.
- Use a new pipette tip with each dilution.
- The standards prepared by serial dilution must be used within one hour of preparation.
 - Discard any unused standards except the standard stock.
 - The standard stock can be stored at ≤ -20 °C for one month or at ≤ -80 °C for more than one month.

Example of Standards Preparation



- The quality of the standard curves can be determined by the % recovery of the standards and the QC values.

Quality Controls

- We include Quality Controls (QCs) to qualify assay performance.
 - QC values are based on a minimum of six assays run by at least three different operators. The midpoint and +/- 35% of the high/low value (mean) are reported.
 - When a customer contacts Technical Support with a concern related to assay performance, the customer is usually first asked if the QC values are in a specific range. This tells the Technical Support Specialist whether or not the kit is performing correctly.
-

- Individual labs can qualify their own assay performance by including a high and low QC that may better reflect their unique experimental samples.
-

- QCs are important for translational studies that require more validation, ensuring that the data are reproducible across various kit lots.
-

- QCs are also important when comparing data for multi-site studies or to compare assay results from multiple technicians.
-

Effect of Serum Matrix

- If the recovery of analytes spiked into sample wells in an assay using a buffer standard curve falls outside our acceptance criteria (80% – 120%), this indicates that there is a nonspecific matrix effect from the samples.
 - To compensate for this effect, a serum matrix with a similar effect is added to the standard curve wells to shift the actual curve so that it matches the recovery in the sample wells.
 - Serum matrix is usually a similar sample with all the endogenous and cross-reacting analytes extracted.

- Because blood is a complex matrix, which contains large numbers of proteins that may interfere with the accurate measurement of desired analytes, using an optimized serum matrix in the standard curves when measuring analytes secreted in serum/plasma:
 - Significantly improves accuracy of measurement.
 - More accurately simulates the conditions of the native analyte present in serum or plasma compared to a standard curve generated by spiking an analyte into a buffer solution.
 - Mimics the environment of native analytes in serum or plasma.
-

- Other commercial multiplex kits add a serum matrix to sample wells. With some exceptions, we do not do this for the following reasons:
 - While this method does effectively show good recoveries, in most cases, adding serum matrix to sample wells can mask the matrix effect, likely affecting the sensitivity of the actual analyte measurement.
 - It is very difficult to predict the effect of mixing serum matrix with samples from a randomly sampled population.
-

- Kits designed for non-serum/plasma samples (e.g., urine, CSF) or samples that require a significant dilution (at least 1:20) do not require serum matrix.
-

- For non-serum/plasma samples, the appropriate medium (e.g., cell culture medium) should be added instead of serum matrix.
 - In the absence of appropriate medium or when using a blank, assay buffer can be used.
 - For cell/tissue homogenates, the final cell or tissue homogenate should be prepared in a buffer that has a neutral pH, contains minimal detergents or strongly denaturing agents and has an ionic strength close to physiological concentrations.
 - An approximately 1:20 dilution would be required to eliminate the need for using serum matrix in an assay.

Enough Assay Buffer?

- Additional assay buffer that is required for samples requiring higher dilutions may need to be ordered before running the assay. See Protocol sections "Reagents Supplied" or "Replacement Reagents" for the appropriate catalog number.

Wash Buffer

- Incomplete washing can adversely affect the assay outcome.
- All washing must be performed with the wash buffer provided.
- If more is required, see Protocol sections "Reagents Supplied" or "Replacement Reagents" for the appropriate catalog number.



Immunoassay Procedure

Tips for Reducing Variability

- Ensure proper sample collection.
- To avoid low bead counts, thaw, vortex and centrifuge all samples for 5-10 minutes at a minimum of 3,000 x g. Avoid or remove any fat layers that may develop.
- Centrifuge samples after thawing or if they appear turbid. This is especially recommended for plasma samples.
- Ensure the proper mixing of samples and controls.
- Use appropriate pipetting technique:
 - Hold the pipette at the same angle each time.
 - Use pipettes calibrated for values in the middle range (not extremes).
- Warm reagents to room temperature (20–25 °C) before mixing. For assays requiring overnight incubation in a cold room, warm reagents to room temperature on the second day as well.
- Cover the plate with a plate sealer before shaking.
- The plate shaker speed should be increased to agitate the plate at the highest speed that does not lead to splashing on the sealer.

Immunoassay Procedure

- Before running an assay, check the instrument the night before.
 - Is the instrument calibrated?
 - Has it been maintained?
 - Have fresh water prepared, calibrated and accurate pipettes, multichannel pipettes and an orbital shaker or alternative.
 - Confirm availability of a cold room or refrigerator with power access for the orbital shaker.
- When running samples in duplicate, a maximum of 38 samples can be run per kit.

	1	2	3	4	5	6	7	8	9	10	11	12
A	Standard 0 (Background)	Standard 4	QC-2 Control									
B	Standard 0 (Background)	Standard 4	QC-2 Control									
C	Standard 1	Standard 5	Sample 1									
D	Standard 1	Standard 5	Sample 1									
E	Standard 2	Standard 6	Sample 2									
F	Standard 2	Standard 6	Sample 2									
G	Standard 3	QC-1 Control	Etc.									
H	Standard 3	QC-1 Control										

96-Well Plate Map. Sample map showing placement of standards, QCs, background and samples.

- To pre-wet the plate, use 150 μ L wash buffer or assay buffer.
-
- If you accidentally use wash buffer instead of assay buffer for your assay, and if sample has not yet been loaded, remove wash buffer and replace with assay buffer.
 - If sample has been added to the plate with wash buffer, there is a potential for low recovery as it may not have the required protein concentration or protease inhibitors.
-
- Vortex all reagents well before adding them to the plate.
-
- When using frozen samples, it is recommended to thaw the samples completely, mix well by vortexing at high setting and centrifuge at a minimum of 3000 x g prior to use in the assay to remove particulates.
-
- Be precise when adding samples, standards and QCs to the plate.
 - Pipette to the sides of the wells.
 - Be sure all fluid is expressed out of the pipette tips.
-
- For incubating assays overnight, a power supply must be available for the orbital shaker in a refrigerator or cold room.
 - If the plate shaker has been turned off during the night, shake again at room temperature for one hour before proceeding with the assay protocol.
-
- After overnight incubation of assays, remember to allow all reagents to warm to room temperature (20 $^{\circ}$ C – 25 $^{\circ}$ C) before use in the assay.
-
- Detection antibody cocktail and SAPE incubation times are critical. Do NOT exceed the dictated times as this will result in higher background signals.
-
- If the detection antibody has been accidentally aspirated off or poured off before adding SAPE to the well, it is possible to recover the assay:
 - Add 20 μ L to 50 μ L of detection antibody and continue to follow the protocol.
 - Replace the detection antibody cocktail volume with assay buffer, add SAPE and continue to follow the protocol.
 - If no detection antibody is available, add SAPE and continue to follow the protocol, keeping in mind that the signal may be lower.
-
- The plate should be read immediately (within 4 hours) after the assay is finished. If the plate cannot be read immediately, seal the plate, cover with aluminum foil or an opaque lid and store the plate at 2–8 $^{\circ}$ C for up to 72 hours, with samples brought up in sheath/drive fluid. There may be a loss of sensitivity after 24 hours.
-
- Before reading the plate, agitate the plate on the plate shaker at room temperature for 10 minutes.
-
- Do not store processed samples in wash buffer.
-
- It is possible to run a portion of a plate initially, then reuse the plate with other samples later.
 - Cover the wells that are not being used.
 - Use precise volumes of reagents to ensure that enough remains to run the remaining wells at a later time.
 - Store reagents at appropriate conditions quickly after the first use (e.g., stock standard at -20 $^{\circ}$ C or lower).
 - Remake standards for subsequent batches. Be sure to run a standard curve for each batch.
 - When running subsequent batches, cover the previously used wells.
 - The mix of beads may be used for one month if stored at 2–8 $^{\circ}$ C; stock standards should be stored at \leq -20 $^{\circ}$ C for one month and at \leq 80 $^{\circ}$ C for more than one month.
 - If using the same plate, keep the plate very clean. Alternatively, use a second plate for the remaining samples.

Plate Washing

Tips for Reducing Variability

- Orbital Titer Plate Shaker (Lab-Line Instruments Model #4625 or equivalent)
 - Very important: For incubating assays overnight, a power supply must be available for the orbital shaker in a refrigerator or cold room.
 - The orbital 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.
 - However, orbital shakers vary. Your shaker can be calibrated by pre-wetting the plate with buffer and slowly increasing the speed until splashing occurs. Then lower the speed slightly. The shaker should be set at the highest speed allowable without splashing of the liquid.
- Handheld Magnetic Separation Block (Merck Millipore Cat. No. 40-285)
 - When getting ready to decant the liquid from the plate, the plate **MUST** be firmly attached to the magnet.
 - Grip the handheld separation block firmly.
 - During the wash steps, decant with a good shake while the plate is attached to the magnet over a sink.
 - When using a new magnet, check for space between the plate and magnet. Adjustments require a US Allen (hex) key to adjust the screws (not provided).



Equipment Settings



FLEXMAP 3D® System



Luminex® 100/200™ System



MAGPIX® System

- For the MAGPIX® system, choose the "enhanced startup" setting instead of the common startup.
 - This will ensure proper calibration and cleaning prior to running the assay.
 - Working with serum is "dirtier" than other samples and can affect the performance of the instrument unless it is properly cleaned.
 - Luminex® can provide a recommended protocol for maintenance.
- Be sure the needle probe is clean. This may be achieved by sonication and/or alcohol flushes.
- Probe height:
 - When reading an assay on a Luminex 200™ instrument, adjust the probe height according to the protocols recommended by Luminex® to the kit solid plate using 3 alignment discs.
 - When reading an assay on a FLEXMAP 3D® system, adjust probe height according to the protocols recommended by Luminex® to the kit solid plate using 1 alignment disc.
 - When reading an assay on a MAGPIX® system, adjust the probe height according to the protocols recommended by Luminex® to the kit solid plate using 2 alignment discs.
- Annotating control wells on the instrument can be tedious, with a lot of manual typing.
 - It is possible to enter the wells as unknowns instead of controls to avoid typing in the annotations for controls, then comparing with your chart later.
- Use the manufacturer's gate settings.
- The Luminex 200™ system's xPONENT® 3.1 acquisition software has two functions: one for magnetic (MagPlex®) and one for nonmagnetic beads (MicroPlex®).
 - Be sure to select the correct setting in the protocol for your bead type.
 - If the wrong type is selected, the plate does not need to be reread. The batch can be replayed with the corrected protocol setting.
- The link for xPONENT® software templates:
 - www.merckmillipore.com/lmx_xponent (Life Science Research → Protein Detection and Quantification → Luminex® Multiplexing Instruments → Multiplex Assay Analysis Software → xPONENT®)
- If a plate cannot be run immediately (within 4 hours) (e.g., it needs to be taken to another site to run the assay), suspend your sample in sheath or drive fluid or assay buffer.
- 10-12 plates can be run with one bottle of drive fluid for the MAGPIX® system.
- To change a standard curve from, for example, a 7-point curve to an 8-point curve, simply make a new protocol and replay the batch.

Running MILLIPLEX[®] MAP Kits on Other Luminex[®] Instruments

- A Luminex 100™ system with IS software, Luminex 200™, FLEXMAP 3D[®] or MAGPIX[®] instrument is required to run a MILLIPLEX[®] MAP assay.
 - If you want to try a kit before purchasing an instrument, ask your Sales Specialist to provide a demonstration using the Luminex[®] technology and MILLIPLEX[®] MAP kits.
 - Magnetic bead assays cannot be run on any instruments using Luminex[®] IS 2.3 or Luminex[®] 1.7 software.

-
- Since all Luminex[®] machines (Luminex 200™, FLEXMAP 3D[®] and MAGPIX[®] instruments) are built by the Luminex Corporation, MILLIPLEX[®] MAP kits can be run on any of these machines, regardless of the name given to the machine by a Luminex[®] business partner.

- If using Luminex[®] instruments with software other than xPONENT[®] software (Bio-Plex Manager™, MasterPlex[®], SStarStation, LiquiChip, LABScan™ 100), follow instrument instructions for gate settings and additional specifications from the software vendors for reading assays using Luminex[®] magnetic beads.

-
- To read a MILLIPLEX[®] MAP kit on a Bio-Plex[®] machine, select 5K-25K for magnetic beads, depending on the version of Bio-Plex Manager™ software.

Overview of Instrument Considerations During a MILLIPLEX[®] MAP Assay

Starting up and shutting down your system correctly will ensure its longevity. The instructions for the MAGPIX[®] and Luminex 200™ systems are located within the systems user manual*. Short-term cleaning will prevent sample induced clogging, while long-term cleaning is important to ensure that drive fluid does not evaporate and crystallize.

Preparation

- Check probe and insert into reader; set probe height.
- Fill reservoirs (Milli-Q[®] water, 70% EtOH, 0.1M NaOH).
- Revive instrument: revive from storage, daily start-up.
- Calibrate and verify instrument: system installation.
- Read the entire kit protocol.
- Acquire "Materials Required But Not Supplied."
- Confirm accuracy of pipettes.

Assay

- Follow the kit protocol.
- Set up experimental design on acquisition software.
- Run assay.
- Run "Post Batch Routine."

Shutdown

- Daily shutdown (overnight):
 - Run "Clean Routine".
 - Run "Daily Shutdown Routine".
 - Remove probe and clean in a sonicating water bath.
- Long-term shutdown (longer than one week):
 - Run "Clean Routine" multiple times.
 - Run "Prepare for Storage" part 1.
 - Prime multiple times with Milli-Q[®] water (use an empty sheath fluid container).
 - Run "Prepare for Storage" part 2.
 - Remove probe and clean in a sonicating water bath.

* Luminex 200™ User Manual, Section 3, page 17. MAGPIX[®] User Quick Guide 4.2

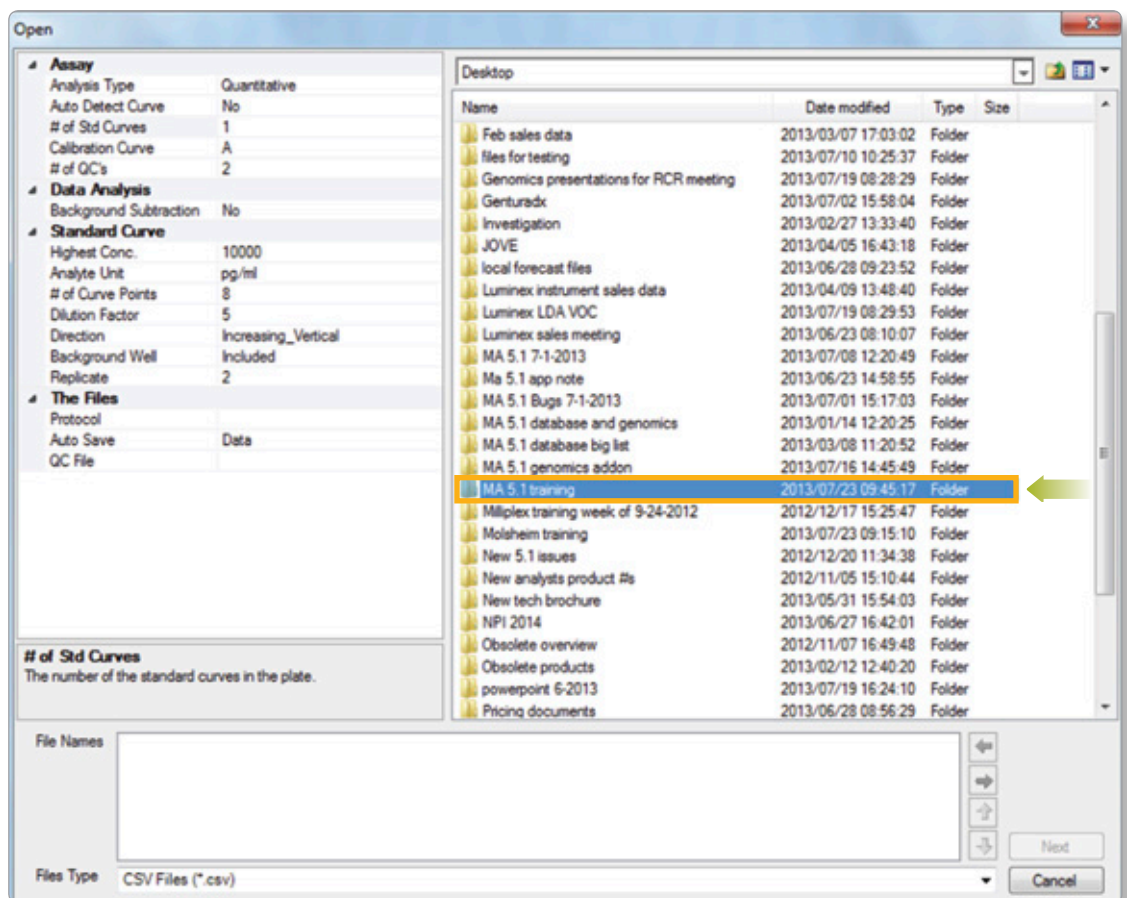
MILLIPLEX® Analyst 5.1 Software

Merck Millipore offers the most powerful combination software package, including best-in-class multiplex data analysis MILLIPLEX® Analyst 5.1 software coupled with data acquisition using the Luminex xPONENT® software. MILLIPLEX® Analyst 5.1 software enables you to manage, track and analyze your multiplex assays rapidly and efficiently, giving you more time to focus on advancing your research.

Data acquisition and analysis integrates seamlessly with all Luminex® instruments, including FLEXMAP 3D®, Luminex 200™ and MAGPIX® systems. MILLIPLEX® Analyst 5.1 software is available in one- and five-seat licenses, enabling complete flexibility for small, medium and large laboratories.

Step 1:

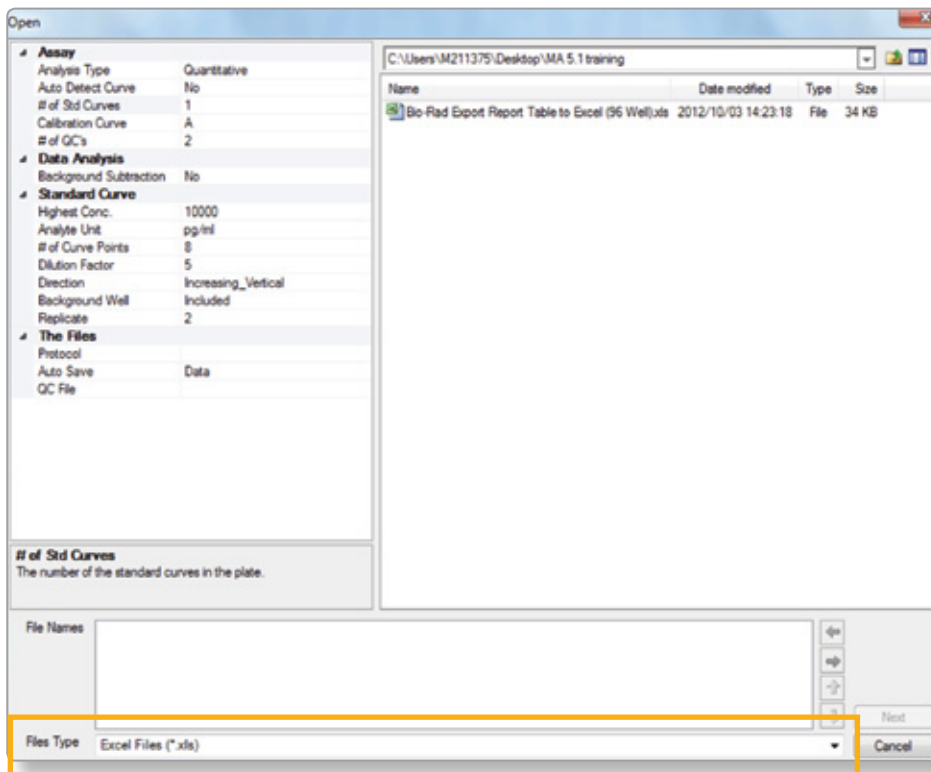
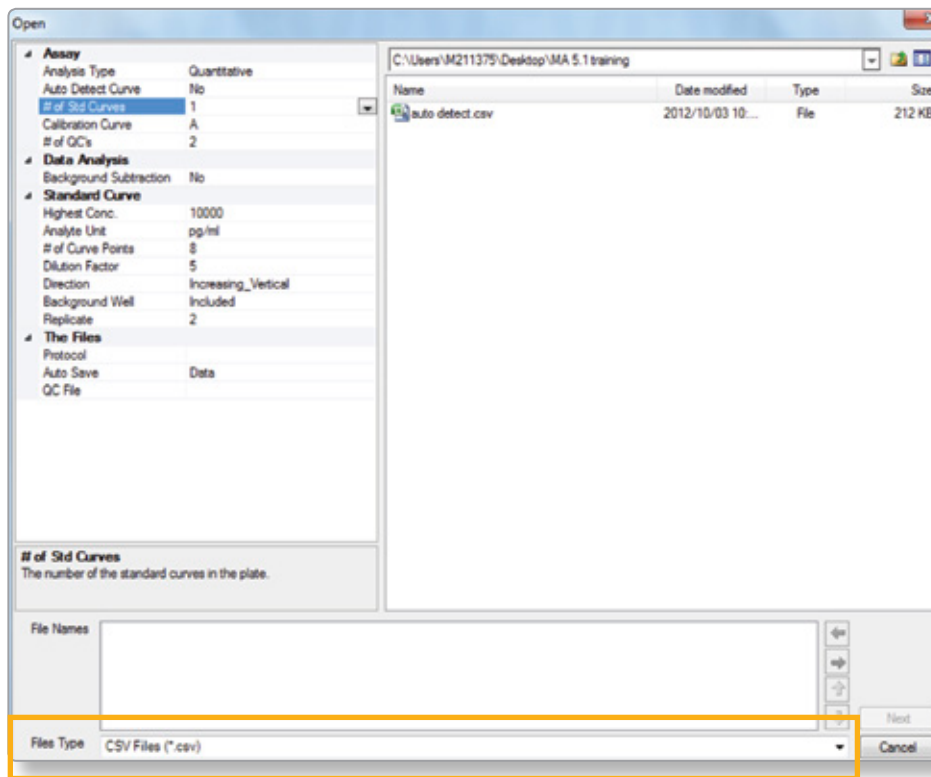
Choose the folder containing the exported data files.



Step 2:

Select the appropriate File Type from the pull-down menu:

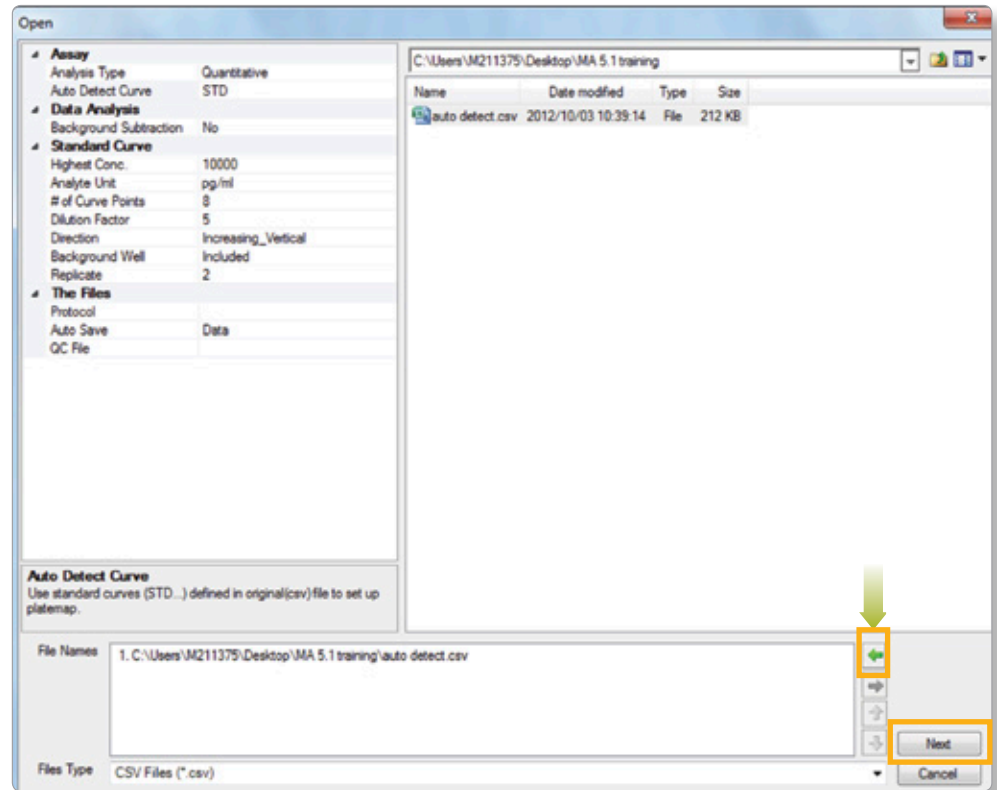
- CSV Files (*.csv) from xPONENT® software.
- Excel® files (*.xls) from Bio-Plex® software.



Step 3:

Importing data from xPONENT® software using Auto Detect Curve:

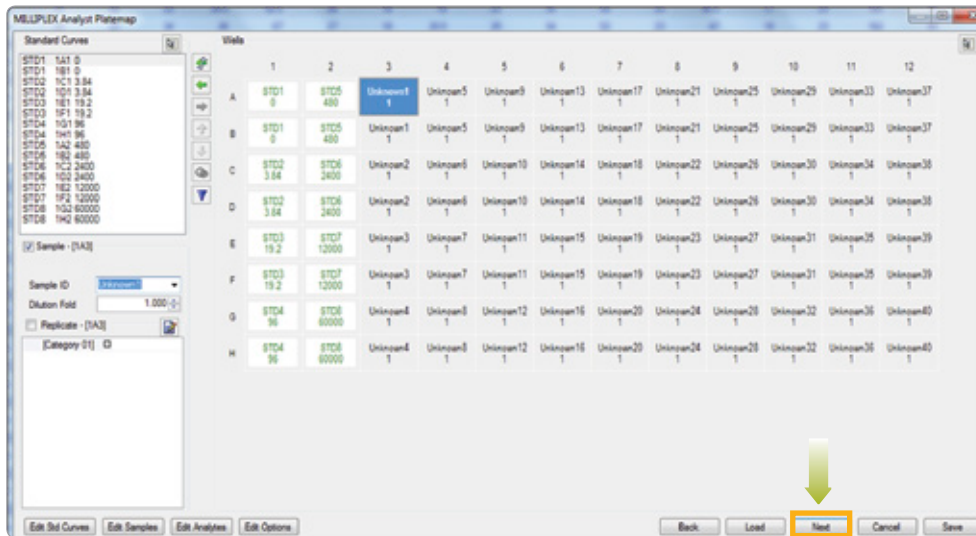
- Set Auto Detect Curve to STD and highlight the file (auto detect.csv).
- Press the green arrow/icon.
- Loaded files are shown in the File Names window.
- The Standard Curve settings will be ignored and the software will use the Standard Curves as set up in xPONENT® software.
- Click "Next" to continue to the Plate Map screen.



Step 4:

Importing Data from xPONENT® software using Auto Detect Curve – Plate Map:

- Assign analytes, dilutions and sample names as needed.
- If the kit and Plate Layout will be used again, save as a "Protocol".
- When ready for Analysis, click "Next".



Step 5:

Data Analysis with MILLIPLEX® Analyst 5.1 software:

- Concentration of each analyte is calculated in pg/mL using the Standard Curve.
- Best Fitting is a 5-P log curve fitting algorithm.
- Click on different analytes to see where that sample is on the Standard Curve.
- All graphs are updated in real time.
- Detailed Reports can be saved as an Excel® file.



Data Analysis

Bead Counts

- Merck Millipore recommends counting 50 beads.
 - According to Luminex®, a minimum of 35 beads per region need to be counted.
 - Fewer than 35 beads could cause a shift in the MFI (Median Fluorescence Intensity) value of the bead population.
 - However, MFI will not change for bead counts greater than or equal to 35.
 - Therefore, don't worry if there is a 35 bead count on one bead region and 400 for others. MFIs will not be affected.

How to Correct or Prevent Low Bead Counts

- Be sure to specify MagPlex® in the kit protocol for xPONENT® software or use the correct gate setting on Bio-Plex® software.
- Sample preparation: Thaw, vortex and centrifuge samples at a minimum of 3,000 x g. Avoid or remove any fat layers that may develop.
- For samples known to be challenging (e.g., synovial fluid, saliva), one may increase wash steps after incubation with primary antibody.
- Resuspend beads in wash buffer instead of sheath/drive fluid. However, the plate must be read within four hours.
- Add 1x wash buffer, which contains Tween® 20, to keep the beads from clumping or sticking.
- Store only in sheath or drive fluid.

- When using a handheld magnet, blot the plate gently.

- When using a plate washer, check the settings to make sure the plate is soaking for 60 seconds and the aspiration is not all the way down in the well.

- Warm the plate to room temperature after an overnight 4 °C capture antibody incubation step. Let the plate shake at room temperature for one hour.

- For MAGPIX® users, cleaning the instrument is critical.

- Special care should be taken to use the enhanced startup or washing procedures.
- There is an advanced cleaning method that includes sodium hydroxide (NaOH) and bleach.
- Washing between wells can also be selected during the plate reading.
- Cleaning the instrument regularly is important even if the instrument is not being used.

Percent Coefficient of Variation (%CV)

- High %CVs for standards or samples can be due to bead count.

- For assays that have a standard curve, our target inter-assay %CV is <15% and our target intra-assay %CV is <10%.

- For qualitative assays (no standard curves) the MFI target inter-assay %CV is <20% and the target intra-assay %CV is <15%.

Calculating Lower Limit of Quantification (LLOQ) and Upper Limit of Quantification (ULOQ)

- Defining LLOQ and ULOQ requires a tight standard curve (e.g., a 1:2 or 1:3 serial dilution to the point that you achieve saturation at both ends).
- Choose the lowest and highest standard curve points that have a recovery of +/- 20%.
- Verify that this is the LLOQ and ULOQ by running 5 assays with the LLOQ and ULOQ as samples against a curve using the assay serial dilution factor where the lowest standard is below LLOQ and the highest standard is above ULOQ.
- Inter-assay precision should be within 20% for LLOQ and ULOQ samples.

Curve Performance/Fit

- Standard point %CVs should be <15%.
 - High %CVs here indicate improper technique was used when making standard curve dilutions. Examples of poor technique include:
 - Not vortexing between tubes.
 - Not vortexing while loading the plate.
 - Not pipetting equal amounts into the plate.
 - The lower the concentrations of analytes, the higher the %CVs tend to be. With new users, this improves with time and practice.
 - For any standard points that have high %CVs, samples in that range of the curve should be interpreted with caution.
 - Alternatively, a standard point or one of the replicate wells can be flagged/masked, although it can be difficult to decide which well to flag if only duplicates are run.

Recovery

- Percent recovery should be 100% +/- 30% (industry minimum), although some customers will have their own acceptance criteria.
- Percent recovery is usually worse at either extreme of the curve, but this also improves with time and practice.
 - For curve statistics, focus on the R² value, which approach, but never equal unity. (Note that a R² value of "1" is seen with software rounding of 0.9999).

Minimum/Maximum Detectable Concentration (minDC/maxDC)

- For many assays, the minDC/maxDC will be outside the standard points (extrapolated) due to good curve performance and fit.
- To avoid seeing extrapolated data, set the desired range of detection in MILLIPLEX® Analyst 5.1 software.
 - Deciding whether to use the "Best Fit" vs. 5-parameter lot option depends on your comfort level to determine how appropriate it is to "play" with curve fit to find the best one.
 - If samples fall outside the dynamic range of the assay, dilute the samples further with the appropriate matrices/media and repeat the assay.

How Merck Millipore Monitors/Avoids Lot-to-lot Drift

- Lot-to-lot drift is monitored and mitigated using full-curve comparison and comparing the relative potency of each analyte against a reference lot.
- All data are compiled in a single database, and trend charts are maintained in our records.
- MILLIPLEX® MAP standard points maintain consistent values from lot to lot, unlike other kits that may have values that vary lot to lot.

Comparison of MILLIPLEX® Analyst 5.1, StatLIA and Bio-Plex® Analysis Software

Analyte: IFN γ , Kit: Rat Cytokine/Chemokine

Units: pg/mL

	MILLIPLEX® Analyst 5.1*	StatLIA®	Bio-Plex®
Standard1	14.7	3.0	
Standard2	57.4	67.0	53.7
Standard3	241.6	245.0	248.6
Standard4	932.1	897.0	908.1
Standard5	3683.0	3858.0	3820.0
Standard6	15184.0	14769.0	14824.5
Standard7	59874.0	61392.0	60975.2
Unknown1	<3.40 ↓	<1.0e-07	OOR <
Unknown2	16.8	8.0	OOR <
Unknown3	51.3	60.0	45.0
Unknown4	197.1	205.0	205.8
Unknown5	844.1	809.0	821.3
Unknown6	3412.0	3564.0	3531.3
Unknown7	14639.0	14296.0	14339.3
Unknown8	70718.0	82002.0	78697.5
Unknown9	<3.40 ↓	<1.0e-07	OOR <
Unknown10	<3.40 ↓	<1.0e-07	OOR <
Unknown11	<3.40 ↓	<1.0e-07	OOR <
Unknown12	3.7	<1.0e-07	OOR <
Unknown13	<3.40 ↓	<1.0e-07	OOR <
Unknown14	<3.40 ↓	<1.0e-07	OOR <
Unknown15	5.0	<1.0e-07	OOR <
Unknown16	16.8	8.0	OOR <
Unknown17	24.0	22.0	OOR <
Unknown18	28.6	29.0	1.1
Unknown19	<3.40 ↓	<1.0e-07	OOR <
Unknown20	<3.40 ↓	<1.0e-07	OOR <
Unknown21	<3.40 ↓	<1.0e-07	OOR <
Unknown22	14.7	3.0	OOR <
Unknown23	<3.40 ↓	<1.0e-07	OOR <
Unknown24	<3.40 ↓	<1.0e-07	OOR <
Unknown25	6.8	<1.0e-07	OOR <
Unknown26	<3.40 ↓	<1.0e-07	OOR <
Unknown27	<3.40 ↓	<1.0e-07	OOR <
Unknown28	5.9	<1.0e-07	OOR <
Unknown29	4.5	<1.0e-07	OOR <
Unknown30	8.2	<1.0e-07	OOR <
Unknown31	<3.40 ↓	<1.0e-07	OOR <
Unknown32	<3.40 ↓	<1.0e-07	OOR <

Green: Extrapolated value
 Orange: Extrapolated value
 OOR<: Out of Range Below
 <3.40 ↓ : Out of Range Below
 <1.0e-07: Out of Range Below
 *Best Fitting, 5P Log

Table 2.

Significantly more IFN γ concentrations could be calculated at the low end of the curve in the Rat Cytokine/Chemokine Magnetic Bead Panel by MILLIPLEX® Analyst 5.1 software compared to the Bio-Plex® and StatLIA® software packages.

Intracellular Assays

Differences between multiplex assays for circulating analytes vs. intracellular analytes

Circulating Analyte Assay	Intracellular Analyte Assay
Quantitative	Qualitative (fold change)
Serum, plasma, tissue culture, urine, CSF, etc.	Cells must be lysed
Analytes analytically validated within panel	Fixed kits and individual MAPmates™ are analytically validated
Kit includes standards and QCs	Kits and MAPmate™ assays include positive and negative control cell lysates
Most panels are customizable	Kits are fixed; create custom kits by combining Cell Signaling MAPmate™ assays

Using Cell Signaling MAPmate™ Assays

- All magnetic MAPmate™ assays require the Cell Signaling Buffer & Detection Kit (Cat. No. 48-602MAG).
 - This kit contains all necessary reagents except the MAPmate™ assays. Both a filter and flat-bottom plate are included for convenience.
- To select the appropriate buffer for your MAPmate™ assays, please refer to the protocols or the buffer selection tables on the website, in the Analyte Quarterly or Appendix 5. www.merckmillipore.com/cellsignaling_assays
- The following MAPmate™ assays should not be plexed together:
 - Phospho-specific and total MAPmate™ pairs (e.g., total GSK3β and phospho-GSK3β (Ser9)).
 - Pan Tyr and site-specific MAPmate™ assays (e.g., phospho-EGF Receptor (pan Tyr) and phospho-STAT1 (Tyr701)).
 - More than 1 phospho-specific MAPmate™ assay for a single target (e.g., pAkt (Ser473) and pAkt (Thr308)).
 - GAPDH and β-Tubulin assays cannot be plexed with kits or MAPmate™ assays containing pan Tyr assays.

Preparation of Cell Lysates for Intracellular Assays

- 96-well plates:
 - For adherent cell lines: seed ~40,000 cells/well and allow growth for 48 hours.
 - For suspension cell lines: seed ~250,000 cells/well and collect at desired time.
 - For cell lysis: add 30 μL lysis buffer per well and pipet up and down thoroughly without creating too many bubbles. For a more detailed protocol, request info from Technical Support.
 - Unbroken cells/parts can be cleared by either filtration or by centrifugation.
- MAPmate™ assays can also be added to existing intracellular assay kits to enhance the panel or serve as controls.
 - Refer to the guidelines provided in the kit protocol.
- MAPmate™ assays can also be added to existing intracellular assay kits to enhance the panel or serve as controls.
 - Refer to the guidelines provided in the kit protocol.
- Add protease inhibitors and/or phosphatase inhibitors to "home-brew" lysis buffers.

- Lysis buffer can be found in the Cell Signaling Buffer & Detection Kit (Cat. No. 48-602MAG) or it is sold separately (Cat. No. 43-040).
-
- Other lysis buffer selections:
 - Non-ionic detergents (NP40, Tergitol, IPEGAL) are recommended in lysis buffers for solubilizing cytoplasmic proteins.
 - Partially ionic detergents (Triton® X-100) are recommended in lysis buffers for cytoplasmic or membrane-bound proteins.
 - Ionic detergents (sodium dodecyl sulfate, SDS) are recommended in lysis buffers for membrane-bound, nuclear or mitochondrial proteins. If using SDS in the lysis buffer (i.e., Radioimmunoprecipitation assay (RIPA) buffer), then cell lysate must be diluted to less than 0.05% SDS for assays to detect intracellular proteins, such as cell signaling proteins.
 - Note: to solubilize nuclear/mitochondrial proteins, you must use either SDS or another method (such as ultrasonication) to puncture the tough nuclear/mitochondrial membranes.
 - Reducing agents, like β -mercaptoethanol or dithiothreitol, are not recommended.

Type of detergents	Protein localization	Maximum allowed protein concentration	Luminex® assay compatibility
Non-ionic detergents	Cytoplasm	5 mg/mL	Yes
Partially ionic detergents	Cytoplasm, Membrane-bound	5 mg/mL	Yes
Ionic detergents	Membrane-bound, Nucleus, Mitochondria	5 mg/mL	Requires dilution

- Perform all dilutions with lysis buffer (not assay buffer or phosphate-buffered saline (PBS)).

- Total protein concentration limits:
 - Do not collect lysates at greater than 5 mg/mL protein concentration.
 - At protein concentrations higher than 5 mg/mL, not all proteins will be solubilized equally by the lysis buffer. Some proteins can be solubilized at a given detergent concentration, while other proteins are not as affected. For example, β -tubulin signal decreases with increasing total protein concentration (signal decrease occurs at 5 to 6 mg/mL for Jurkat cell and peripheral blood mononuclear cell (PBMC) lysates).
 - Total protein concentrations should be within a specific range, which is outlined in each protocol.
 - A starting protein amount is 10 μ g per well (10 μ g protein in the final 25 μ L that is loaded into each assay well) is recommended.
 - Working backwards from there:
 - 10 μ g/25 μ L = 0.4 μ g/ μ L (mg/mL)
 - Diluting the cell/tissue lysates 1:1 in the assay buffer provided in the intracellular kit is recommended.
 - Consequently, all samples need to be brought to a protein concentration of 0.8 μ g/ μ L in lysis buffer.
 - Take 30 μ L of each lysate sample and add it to 30 μ L of assay buffer, bringing the final concentration down to 0.4 mg/mL.
 - Then load 25 μ L of diluted samples into each well, usually in duplicate.

Appendix 1: Species Cross-reactivity

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Canine

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	Eotaxin	FGF-2	Fractalkine	G-CSF	GM-CSF	GRO	IFN α 2
Human Cytokine/Chemokine Panel 1	4	2	1	1	1	1	1	1
	IFN γ	IL-1 α	IL-1 β	IL-1ra	IL-2	IL-3	IL-5	IL-6
	1	2	3	2	2	2	1	1
	IL-7	IL-8	IL-9	IL-10	IL-12(p40)	IL-13	IL-15	IL-17A
	1	1	1	2	1	2	2	1
	IP-10	MCP-1	MCP-3	MDC	MIP-1 α	MIP-1 β	sCD40L	TGF α
	1	1	2	2	2	1	2	2
	TNF α	TNF β	VEGF-A	PDGF-AB/BB				
	0	2	1	3				

Panel Name	SDF-1 α + β	EOTAXIN-3	CTACK	IL-23	TPO	TSLP	IL-33
Human Cytokine/Chemokine Panel 2	1	1	1	2	1	1	1

Panel Name	IL-17F	GM-CSF	IL-10	MIP3 α	IL-15	IL-17A	IL-22	IL-9
Human TH17	1	4	3	1	3	1	3	3
	IL-1 β	IL-33	IL-2	IL-4	IL-23	IL-17E	IL-27	IL-31
	3	1	3	3	3	1	2	1
	TNF β							
	3							

Panel Name	GM-CSF	sCD137	IL-10	IL-13	Granzyme B	IL-2	IL-4	MIP-1 α
Human CD8+	1	2	1	1	1	4	1	1
	MIP-1 β							
	1							

Panel Name	GM-CSF	IFN γ	IL-10	IL-13	IL-17A	IL-1 β	IL-2	IL-4
Human High Sensitivity T Cell	2	1	3	1	2	1	1	3
	IL-23	IL-7	IL-8	MIP1 α	MIP1 β			
	3	1	3	2	1			

Panel Name	Eotaxin	G-CSF	GM-CSF	IFN γ	IL-1 α	M-CSF	IL-1 β	IL-2
Mouse Cytokine/Chemokine Panel 1	4	1	4	3	2	4	4	2
	IL-3	IL-4	IL-5	IL-7	IL-10	IL-12(p40)	IL-13	IL-15
	2	1	3	3	1	1	3	2
	IL-17A	IP-10	MIP-2	KC	LIF	LIX	MCP-1	MIP-1 α
	2	4	2	4	2	4	4	4
	MIP-1 β	MIG	RANTES	TNF α	IL-12(p70)	VEGF-A	IL-9	
	1	3	3	4	2	4	3	

Cytokine kits (continued)

Panel Name	GM-CSF	IFN β	MIP-3 α	IL-1 β	IL-2	IL-4	IL-5	IL-6
Mouse TH17	4	3	1	2	3	1	1	1
	IL-21	IL-22	IL-28B	IL-10	IL-23	IL-12(p70)	IL-27	IL-13
	4	4	3	2	1	1	0	2
	IL-15	IL-17F	IL-33	IL-31	TNF β	TNF α	sCD40L	
4	4	4	1	4	3	4		
Panel Name	GM-CSF	TGF α	G-CSF	IFN γ	IL-2	IL-10	IL-15	sCD40L
Non-Human Primate Cytokine/ Chemokine	1	4	3	2	4	0	2	1
	IL-17A	IL1ra	IL-13	IL-1 β	IL-4	IL-5	IL-6	IL-8
	4	3	0	2	2	4	2	1
	MIP-1 α	TNF α	IL-12	VEGF-A	IL-18			
1	3	1	4	2				
Panel Name	G-CSF	Eotaxin	GM-CSF	IL-1 α	Leptin	MIP-1 α	IL-1 β	IL-2
Rat Cytokine/Chemokine	1	1	3	2	4	4	4	4
	IL-6	EGF	IL-13	IL-10	IL-12(p70)	IFN γ	IL-17A	IL-18
	2	4	3	4	2	2	1	4
	MCP-1	IP-10	GRO α /KC	VEGF-A	Fractalkine	LIX	MIP-2	TNF α
	3	4	3	4	4	4	3	2
	RANTES							
1								
Panel Name	IL-1 α	IL-1 β	IL-1ra	IL-2	IL-4	IL-6	IL-10	IL-12
Porcine Cytokine/Chemokine	1	4	4	4	4	2	4	4
	IL-18							
4								

Cardiovascular Disease (CVD) kits

Panel Name	NT proBNP	CK-MB	CXCL6	Endocan-1	FABP4	LIGHT	Oncostatin	Troponin-I
Human CVD1	4	4	1	1	1	1	1	1
Panel Name	ADAMTS13	D-Dimer	FABP5	GDF-15	Myoglobin	sP-Selectin	sVCAM-1	SAA
Human CVD2	4	3	4	4	3	4	3	1
Panel Name	α -2-Macroglobulin	AGP	Fibrinogen	sL-Selectin	Haptoglobin	Platelet Factor 4	von Willebrand Factor	
Human CVD3	4	3	1	4	4	2	4	
Panel Name	Follistatin	dPAPP-A	sPECAM-1	Pentraxin-3	Tissue Factor	Thrombomodulin	Troponin T	
Human CVD4	3	4	2	4	1	1	3	
Panel Name	MMP-9							
Mouse CVD1	1							

Angiogenesis kits: 8 samples run

Panel Name	EGF	ANGPT-2	Leptin	FGF-1	IL-8	HGF	HB-EGF	VEGF-C	VEGF-D	FGF-2	VEGF-A
Human Angiogenesis/ Growth Factor	3	8	1	8	1	2	8	2	6	2	2

Feline

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	Eotaxin	FGF-2	Flt-3L	Fractalkine	G-CSF	GRO	IFN α 2
Human Cytokine/Chemokine Panel 1	1	1	1	1	2	2	2	2
	IFN γ	IL-1 α	IL-1 β	IL-1ra	IL-2	IL-3	IL-5	IL-6
	1	2	3	2	2	2	1	1
	IL-7	IL-8	IL-9	IL-10	IL-12(p40)	IL-13	IL-15	IL-17A
	1	1	1	2	1	2	2	1
	IP-10	MCP-1	MCP-3	MDC	MIP-1 α	MIP-1 β	sCD40L	TGF α
	1	1	2	2	2	1	2	2
TNF α	TNF β	VEGF-A	PDGF-AB/BB					
0	2	1	3					

Panel Name	SDF-1 α + β	I-309	IL-23	TPO	IL-33
Human Cytokine/Chemokine Panel 2	3	1	3	2	3

Panel Name	GM-CSF	IL-2	MIP-1 α	Perforin
Human CD8+	1	2	1	1

Panel Name	IL-17E	GM-CSF	MIP-3 α	IL-1 β	IL-2	IL-4	IL-5	IL-21
Human TH17	3	3	1	1	1	2	2	3
	IL-22	IL-28B	IL-10	IL-23	IL-12(p70)			
	4	4	3	2	4			

Panel Name	GM-CSF	TGF α	G-CSF	IFN γ	IL-2	IL-15	IL-17A	IL-1ra
Non-Human Primate Cytokine/ Chemokine	1	3	3	2	3	1	2	3
	IL-13	IL-1 β	IL-4	IL-5	IL-6	IL-8	MIP-1 α	TNF α
	2	3	3	3	2	3	1	2
	MIP-1 β	IL-12/23 (p40)	VEGF-A	Fractalkine	LIX	MIP-2	TNF α	RANTES
0	1	3	4	4	4	2	2	

Panel Name	G-CSF	Eotaxin	GM-CSF	IL-1 α	Leptin	MIP-1 α	IL-4	IL-1 β
Rat Cytokine/Chemokine	2	3	4	3	4	4	3	4
	IL-2	IL-6	EGF	IL-13	IL-10	IL-12(p70)	IFN γ	IL-5
	3	2	4	2	4	3	2	2
	IL-17A	IL-18	MCP-1	IP-10	GRO α /KC			
2	3	3	4	4				

Panel Name	GM-CSF	IFN γ	IL-2	IL-6	IL-7	IL-8	IL-15	IP-10
Canine Cytokine/Chemokine	4	2	4	4	3	1	3	3
	KC-like	IL-10	IL-18	MCP-1	TNF α			
	3	1	4	4	1			

Panel Name	GM-CSF	IL-1 α
Porcine Cytokine/Chemokine	1	1

CVD kits

Panel Name	FABP3	FABP4	Troponin-I
Human CVD1	3	1	4

Panel Name	ADAMTS13	FABP5	GDF-15	Myoglobin	sP-Selectin
Human CVD2	4	4	4	4	1

Panel Name	α -2-Macroglobulin	AGP	sL-Selectin	SAP	Haptoglobin	Platelet Factor 4	von Willebrand Factor
Human CVD3	4	1	4	1	3	1	4

Panel Name	dPAPP-A	Tissue Factor
Human CVD4	2	1

Panel Name	MMP-9	sP-Selectin
Mouse CVD1	1	4

Angiogenesis kits: 6 samples run

Panel Name	EGF	G-CSF	ET-1	FGF-1	Follistatin	HB-EGF	VEGF-A
Human Angiogenesis/ Growth Factor	5	2	1	5	3	5	2

Guinea Pig

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	IL-3	MDC
Human Cytokine/Chemokine Panel 1	1	2	4

Panel Name	SDF-1 α + β	BCA-1	IL-16
Human Cytokine/Chemokine Panel 2	4	1	1

Panel Name	GM-CSF	MIP3 α
Human TH17	4	3

Panel Name	IL-2
Human CD8+	1

Panel Name	Eotaxin	IFN γ	IL-1 β	IL-12(p40)	IL-13	IL-15	IL-17A	IP-10
Mouse Cytokine/Chemokine Panel 1	2	2	2	4	4	1	1	4

Panel Name	MIP-2	LIX	RANTES
Human CD8+	4	4	2

Cytokine kits (continued)

Panel Name	IL-17E	GM-CSF	IFN γ	IL-2	IL-4	IL-5	IL-22	IL-28B
Mouse TH17	4	3	4	2	1	4	4	4
	IL-10	IL-23	IL-12(p70)	IL-15	IL-17A	IL-17F	IL-33	TNF β
	1	4	4	2	4	1	4	2
	sCD40L							
	3							

Panel Name	GM-CSF	G-CSF	IL-2	VEGF-A
Non-Human Primate Cytokine/ Chemokine	1	4	4	2

Panel Name	Eotaxin	GM-CSF	IL-1 α	Leptin	IL-4	IL-1 β	IL-2	IL-6
Rat Cytokine/Chemokine	1	2	2	4	3	4	2	1
	IL-13	IL-10	IL-12(p70)	IL-18	MCP-1	IP-10	GRO α /KC	VEGF-A
	2	4	1	1	3	2	2	2
	LIX	MIP-2	TNF α	RANTES				
	4	4	2	2				

Panel Name	GM-CSF	IL-15	IP-10	IL-18
Canine Cytokine/Chemokine	4	1	4	2

Panel Name	GM-CSF	IL-1 β	IL-1ra	IL-2	IL-4	IL-6	IL-10	IL-12
Porcine Cytokine/Chemokine	3	4	1	2	3	1	1	2
	IL-18							
	2							

CVD kits

Panel Name	ADAMTS13	D-Dimer	FABP5	GDF-15	Myoglobin	sP-Selectin
Human CVD2	4	3	4	4	3	1

Panel Name	AGP	Haptoglobin	von Willebrand Factor
Human CVD3	2	4	4

Panel Name	MMP-9	PAI-1 (total)	sP-Selectin
Mouse CVD1	2	3	3

Angiogenesis kits: 6 samples run

Panel Name	ANGPT-2	BMP-9	FGF-1	Follistatin	HB-EGF
Human Angiogenesis/Growth Factor	5	4	1	5	5

Hamster

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	FGF-2	Fractalkine	IL-1 α	IL-3	MIP-1 β	PDGF-AA	PDGF-AB/ BB
Human Cytokine/Chemokine Panel 1	2	1	3	1	4	1	1	

Panel Name	SDF-1 α + β	CTACK	IL-21
Human Cytokine/Chemokine Panel 2	4	1	1

Panel Name	GM-CSF	MIP3 α	IL-13	TNF- α
Human TH17	4	1	4	4

Panel Name	G-CSF	GM-CSF	IFN γ	IL-1 α	M-CSF	IL-7	IL-12(p40)	IL-13
Mouse Cytokine/Chemokine Panel 1	1	1	2	1	1	1	1	4
	IL-15	IL-17A	IP-10	LIX	MIP-1 α	MIP-1 β	MIG	RANTES
	2	1	1	1	1	2	2	3
	IL-12(p70)	VEGF-A	IL-9					
	1	4	3					

Panel Name	IL-17E	GM-CSF	IFN γ	IL-2	IL-4	IL-5	IL-6	IL-21
Mouse TH17 Panel	2	3	1	2	2	2	2	1
	IL-22	IL-28B	IL-10	IL-23	IL-27	IL-13	IL-15	IL-17A
	4	4	4	2	1	3	2	1
	IL-33	TNF β	sCD40L					
	2	4	1					

Panel Name	Leptin	IL-1 β	IL-2	IL-13	IL-10	IL-4	MCP-1	IP-10
Rat Cytokine/Chemokine	4	4	1	1	3	2	1	4
	GRO α /KC	VEGF-A	Fractalkine	LIX	MIP-2	RANTES		
	4	3	2	1	3	2		

Panel Name	GM-CSF	IL-1 β	IL-1ra	sCD40L	IL-17A	IL-12	IL-18
Porcine Cytokine/Chemokine	3	3	1	3	2	4	1

Panel Name	TGF α	G-CSF	IFN γ	IL-2	IP-10	IL-13	IL-5	IL-6
Non-Human Primate Cytokine/ Chemokine	1	2	3	4	4	2	4	2
	IL-8	MIP-1 α	MCP-1	TNF α	IL-12/23 (p40)	VEGF-A		
	1	1	3	1	1	3		

Panel Name	IL-18
Canine Cytokine/Chemokine	2

CVD kits

Panel Name	FABP3	Troponin-I					
Human CVD1	4	4					

Panel Name	ADAMTS13	D-Dimer	FABP5	GDF-15	MPO	sP-Selectin	
Human CVD2	4	4	4	4	4	1	

Panel Name	dPAPP-A	Tissue Factor	Troponin T				
Human CVD4	3	1	3				

Panel Name	MMP-9	PAI-1 (total)	sP-Selectin				
Mouse CVD1	1	1	4				

Angiogenesis kits: 6 samples run

Panel Name	BMP-9	FGF-1	Follistatin				
Human Angiogenesis/Growth Factor	3	3	2				

Horse

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	Eotaxin	Fractalkine	G-CSF	GM-CSF	GRO	IFN α 2	IFN γ
Human Cytokine/Chemokine Panel 1	4	4	1	4	4	4	4	2
	IL-1 α	IL-1 β	IL-1ra	IL-3	IL-4	IL-5	IL-6	IL-7
	4	4	4	4	4	4	4	4
	IL-8	IL-9	IL-10	IL-12(p40)	IL-12(p70)	IL-13	IL-15	IL-17A
	4	4	4	4	1	4	4	1
	IP-10	MCP-1	MCP-3	MDC	MIP-1 α	MIP-1 β	sCD40L	TGF α
	1	1	4	3	4	1	4	4
	TNF α	VEGF-A	PDGF-AB/BB					
	0	1	4					

Panel Name	MCP-4	SDF-1 α + β	IL-16	MIP-1 δ	6Ckine	CTACK	IL-23	LIF
Human Cytokine/Chemokine Panel 2	1	2	3	2	2	3	3	1
	TPO	TSLP	IL-28A	IL-33				
	3	2	2	2				

Panel Name	IL-17F	GM-CSF	IFN γ	IL-10	MIP-3 α	IL-13	IL-15	IL-17A
Human TH17	2	2	4	4	1	1	2	4
	IL-22	IL-9	IL-1 β	IL-33	IL-2	IL-21	IL-4	IL-23
	1	3	4	3	4	1	1	2
	IL-5	IL-6	IL-17E	IL-27	IL-31	TNF α	TNF β	IL-28A
	1	1	1	2	1	1	4	1

Cytokine kits (continued)

Panel Name	GM-CSF	sCD137	IFN γ	IL-10	Granzyme A	IL-13	Granzyme B	IL-2
Human CD8+	2	1	4	1	2	3	1	4
	IL-4	IL-5	IL-6	MIP-1 α	MIP-1 β	TNF α	Perforin	
	1	1	2	3	1	1	1	
Panel Name	Eotaxin	G-CSF	GM-CSF	IFN γ	IL-1 α	M-CSF	IL-1 β	IL-2
Mouse Cytokine/Chemokine Panel 1	4	4	4	4	4	4	4	4
	IL-3	IL-4	IL-5	IL-6	IL-7	IL-10	IL-12(p40)	IL-13
	4	4	4	4	4	4	4	4
	IL-15	IL-17A	IP-10	MIP-2	KC	LIF	LIX	MCP-1
	4	4	4	4	4	4	4	4
	MIP-1 α	MIP-1 β	MIG	RANTES	TNF α	IL-12(p70)	VEGF-A	IL-9
4	4	4	4	4	4	4	4	
Panel Name	EPO	Exodus-2	Fractalkine	IL-16	IL-21	IL-22	IL-17E	IL-28B
Mouse Cytokine/Chemokine Panel 2	+	+	+	+	+	+	+	+
	MCP-5	MIP-3 α	MIP-3 β	TARC				
	+	+	+	+				
Panel Name	MDC	IL-23	IL-27	TIMP-1	IL-20	IL-33		
Mouse Cytokine/Chemokine Panel 3	+	+	+	+	+	+		
Panel Name	GM-CSF	TGF α	G-CSF	IFN γ	IL-2	IL-15	sCD40L	IL-17A
Non-Human Primate Cytokine/ Chemokine	1	4	3	2	4	4	1	4
	IL-1ra	IL-13	IL-1 β	IL-4	IL-5	IL-6	IL-8	MIP-1 α
	4	1	4	4	1	4	4	4
	TNF α	MIP-1 β	IL-12/23(p40)	VEGF-A	IL-18			
2	1	1	3	4				
Panel Name	G-CSF	Eotaxin	GM-CSF	IL-1 α	Leptin	MIP-1 α	IL-4	IL-1 β
Rat Cytokine/Chemokine	1	1	1	2	2	2	2	2
	IL-2	IL-6	EGF	IL-13	IL-10	IL-12(p70)	IFN γ	IL-5
	2	2	2	2	2	2	2	1
	IL-17A	IL-18	MCP-1	IP-10	GRO α /KC	VEGF-A	Fractalkine	LIX
	1	2	2	2	2	2	2	2
MIP-2	TNF α	RANTES						
2	2	2						
Panel Name	GM-CSF	IFN γ	IL-2	IL-6	IL-7	IL-8	IL-15	IP-10
Canine Cytokine/Chemokine	3	4	4	4	4	2	4	4
	KC	IL-10	IL-18	MCP-1	TNF α			
	3	2	4	4	2			

CVD kits

Panel Name	CK-MB	FABP3	Troponin-I				
Human CVD1	4	1	1				

Panel Name	α -2-Macroglobulin	Fetuin A	AGP	Fibrinogen	sL-Selectin	SAP	Haptoglobin
Human CVD3	4	1	4	3	4	2	3
	Platelet Factor	von Willebrand Factor					
	4	3					

Panel Name	sE-Selectin	dPAPP-A	sPECAM-1	Pentraxin-3	Tissue Factor	Troponin T	
Human CVD4	1	4	1	4	4	4	

Angiogenesis kits: 6 samples run

Panel Name	EGF	ANGPT-2	ET-1	FGF-1	Follistatin	IL-8	HB-EGF
Human Angiogenesis/ Growth Factor	1	3	3	3	3	3	3
	VEGF-D	VEGF-A					
	2	3					

Porcine

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	Eotaxin	FGF-2	Fractalkine	GM-CSF	GRO	IFN α 2	IFN γ
Human Cytokine/Chemokine Panel 1	1	2	4	3	3	3	3	2
	IL-1 α	IL-1 β	IL-1ra	IL-4	IL-5	IL-6	IL-7	IL-8
	3	4	4	3	2	2	3	3
	IL-9	IL-10	IL-12(p40)	IL-12(p70)	IL-13	IL-15	IL-17A	MCP-3
	3	4	4	0	3	3	2	3
	MDC	MIP-1 α	MIP-1 β	sCD40L	TGF α	TNF β	VEGF-A	RANTES
	3	4	3	4	4	4	1	1
	PDGF-AB/BB							
	4							

Panel Name	SDF-1 α + β	IL-16	IL-23	TPO	IL-20	IL-28A	IL-33
Human Cytokine/Chemokine Panel 2	4	1	2	1	1	1	1

Panel Name	IL-17F	GM-CSF	IL-10	IL-15	IL-22	IL-9	IL-1 β	IL-33
Human TH17	3	3	1	3	4	3	3	3
	IL-2	IL-4	IL-23	IL-17E	IL-27	IL-31	TNF β	IL-28A
	4	1	3	2	3	3	2	1

Panel Name	GM-CSF	sCD137	IFN γ	IL-10	Granzyme A	IL-13	Granzyme B	IL-2
Human CD8+	2	2	2	1	2	3	2	2
	IL-4	IL-5	IL-6	sFasL	MIP-1 α	MIP-1 β	TNF α	Perforin
	2	2	2	2	2	2	2	2

Cytokine kits (continued)

Panel Name	GM-CSF	IFN γ	IL-10	MIP3 α	IL-13	IL-17A	IL-1 β	IL-2
Human High Sensitivity T Cell	1	1	2	1	2	3	2	2
	IL-4	IL-23	IL-5	IL-6	IL-8	MIP1 α	MIP1 β	
	2	3	1	1	4	1	1	

Panel Name	IL-17E	GM-CSF	IFN γ	MIP-3 α	IL-1 β	IL-2	IL-4	IL-5
Mouse TH17	2	3	2	2	2	3	2	3
	IL-6	IL-21	IL-22	IL-28B	IL-10	IL-23	IL-12(p70)	IL-27
	2	4	4	3	3	3	3	2
	IL-13	IL-15	IL-17A	IL-17F	IL-33	IL-31	TNF β	TNF α
	2	4	2	4	4	2	4	2
sCD40L								
4								

Panel Name	GM-CSF	TGF α	G-CSF	IFN γ	IL-2	IL-10	IL-15	sCD40L
Non-Human Primate Cytokine/ Chemokine	0	4	2	3	4	0	4	1
	IL-17A	IL-1ra	IL-13	IL-1 β	IL-4	IL-5	IL-6	IL-8
	4	4	0	4	3	2	3	4
	MCP-1	TNF α	MIP-1 β	IL-12	VEGF-A	IL-18		
	1	1	2	1	2	3		

Panel Name	Eotaxin	GM-CSF	IL-1 α	Leptin	MIP-1 α	IL-4	IL-1 β	IL-2
Rat Cytokine/Chemokine	2	2	2	4	4	3	3	3
	IL-6	EGF	IL-13	IL-10	IL-12(p70)	IL-18	MCP-1	IP-10
	1	2	2	3	3	3	2	4
	GRO α /KC	VEGF-A	Fractalkine	LIX	MIP-2	TNF α	RANTES	
3	4	4	3	4	2	2		

CVD kits: 8 samples run

Panel Name	NT proBNP	CK-MB	CXCL6	Endocan-1	Oncostatin	Troponin-I
Human CVD1	1	8	2	2	2	5

Panel Name	ADAMTS13	D-Dimer	FABP5	GDF-15	Myoglobin	sP-Selectin	Lipocalin-2
Human CVD2	8	3	3	8	7	8	1
	SAA						
4							

Panel Name	α -2-Macroglobulin	CRP	Fetuin A	AGP	Fibrinogen	sL-Selectin	Haptoglobin
Human CVD3	4	1	1	4	2	4	4
	Platelet Factor 4	von Willebrand Factor					
1	4						

Panel Name	Tissue Factor	Troponin T
Human CVD4	5	7

Panel Name	sP-Selectin
Mouse CVD1	1

Angiogenesis kits: 6 samples run

Panel Name	EGF	ANGPT-2	ET-1	FGF-1	IL-8	HB-EGF	PLGF
Human Angiogenesis/ Growth Factor	1	5	5	5	1	5	1
	VEGF-C	VEGF-D	FGF-2	VEGF-A			
	1	4	5	1			

Rabbit

- Four serum or plasma samples were usually run in each kit. Exceptions are noted.
- Data below are the number of samples that showed signal above background.
- For more information contact Technical Support.

Cytokine kits

Panel Name	EGF	Eotaxin	FGF-2	G-CSF	GRO	IFN γ	IL-1 α	IL-1 β
Human Cytokine/Chemokine Panel 1	4	4	4	2	4	4	3	4
	IL-1ra	IL-3	IL-4	IL-5	IL-6	IL-7	IL-8	IL-9
	2	1	1	2	2	2	2	3
	IL-10	IL-12(p40)	IL-12(p70)	IL-13	IL-15	IL-17A	IP-10	MCP-1
	3	2	0	2	2	3	1	1
	MCP-3	MDC	MIP-1 α	MIP-1 β	sCD40L	TGF α	TNF β	VEGF-A
	2	4	4	2	4	4	4	3
	PDGF-AA							
	4							

Panel Name	MCP-2	MCP-4	ENA-78	SDF-1 α + β	I-309	TARC	6CKine	Eotaxin-3
Human Cytokine/Chemokine Panel 2	0	0	0	3	2	1	3	3
	CTACK	IL-23	LIF	TSLP	IL-33			
	1	3	3	3	3			

Panel Name	GM-CSF	IL-17A
Human TH17	4	4

Panel Name	GM-CSF	sCD137	IFN γ	IL-10	IL-6
Human CD8+	2	4	4	2	4

Panel Name	IFN γ	IL-1 α	M-CSF	IL-1 β	IL-2	IL-3	IL-4	IL-5
Mouse Cytokine/Chemokine Panel 1	1	2	2	1	1	1	1	1
	IL-12(p40)	IL-15	IL-17A	IP-10	MIP-2	LIF	LIX	MIP-1 α
	4	1	1	2	3	1	3	1
	MIP-1 β	RANTES	TNF α	VEGF-A	IL-9			
	2	4	1	4	2			

Cytokine kits (continued)

Panel Name	IL-17E	GM-CSF	IFN γ	MIP-3 α	IL-2	IL-4	IL-5	IL-6
Mouse TH17	2	4	3	1	3	1	1	4
	IL-22	IL-28B	IL-10	IL-23	IL-12(p70)	IL-27	IL-13	IL-15
	4	4	3	2	2	3	1	1
	IL-17A	IL-17F	IL-33	IL-31	TNF β	sCD40L		
	3	1	2	3	2	4		
Panel Name	GM-CSF	TGF α	G-CSF	IL-2	IL-17A	IL-13	IL-5	IL-8
Non-Human Primate Cytokine/ Chemokine	0	4	2	4	4	1	1	4
	MIP-1 α	VEGF-A						
	3	4						
Panel Name	G-CSF	GM-CSF	IL-1 α	Leptin	MIP-1 α	IL-4	IL-1 β	IL-2
Rat Cytokine/Chemokine	1	3	3	4	3	3	3	3
	IL-6	EGF	IL-13	IL-10	IL-12(p70)	IFN γ	IL-17A	IL-18
	2	2	3	4	2	2	1	4
	MCP-1	IP-10	GRO α /KC	VEGF-A	Fractalkine	LIX	MIP-2	RANTES
	2	4	4	3	3	3	4	1
Panel Name	IFN γ	IL-18	TNF α					
Canine Cytokine/Chemokine	4	3	1					
Panel Name	IL-4	IL-18						
Porcine Cytokine/Chemokine	4	1						

Appendix 2: Sample Preparation

MILLIPLEX® MAP Kits Requiring Special Sample Preparation

Kit Name	Sample Type	Cat. No.	Sample Trt Inhibitors	Inhibitor Source
Canine Gut Hormone Magnetic Bead Panel	SER / PLA / CCS	CGTMAG-98K	DPP-IV	See Note 1
			AEBSF	See Note 2
Human Gut Hormone Magnetic Bead Panel	SER / PLA / CCS	HGT-68K	DPP-IV	See Note 1
			AEBSF Protease Cocktail	See Notes 2, 4
Human IGF-1, IGF-2 Magnetic Bead Panel	SER / PLA / CCS	HIGFMAG-52K	Extraction	NONE
Human IGF Binding Protein (IGFBP) Magnetic Bead Panel	SER / PLA / CCS	HIGFBMAG-53K	Protease Inhibitor Cocktail	See Note 5
Human Metabolic Hormone Magnetic Bead Panel	SER / PLA / CCS	HMHEMAG-34K	DPP-IV	See Note 1
			Aprotinin/AEBSF Protease Cocktail	See Notes 2, 3, 4
Human Neuropeptide Magnetic Bead Panel	SER / PLA / CCS	HNPMMAG-35K	Extraction	NONE
Mouse Gut Hormone Magnetic Bead Panel	SER / PLA / CCS	MGTMMAG-78K	DPP-IV	See Note 1
Mouse Metabolic Hormone Magnetic Bead Panel	SER / PLA / CCS	MMHMAG-44K	DPP-IV	See Note 1
			Aprotinin/AEBSF Protease Cocktail	NONE
Non Human Primate Metabolic Magnetic Bead Panel	SER / PLA / CCS	NHPMMAG-45K	DPP-IV	See Note 1
			Aprotinin/AEBSF Protease Cocktail	See Notes 2, 3, 4
Rat Metabolic Hormone Magnetic Bead Panel	SER / PLA / CCS	RMHMAG-84K	DPP-IV	See Note 1
			Aprotinin/AEBSF Protease Cocktail	See Notes 2, 3, 4
Rat/Mouse Neuropeptide Magnetic Bead Panel	CSF	RMNPMAG-83K	NONE	Extraction
Human Skin Magnetic Bead Panel	SER / PLA / CCS	SKINMAG-50K	Validated for skin tape	NONE
Steroid/Thyroid Hormone Magnetic Bead Panel	SER / PLA / CCS	STTHMAG-21K	Extraction	NONE
TGFβ1 single plex Magnetic Bead Panel	SER / PLA / CCS	TGFBMAG-64K-01	Acidification	NONE
TGFβ1, -2, -3 Panel Magnetic Bead Panel	SER / PLA / CCS	TGFBMAG-64K-03	Acidification	NONE

Note:

1. DPP-IV (Merck Millipore Cat. No. DPP4-010) is used at 10 µL per mL of blood.
2. Pefabloc or AEBSF (Merck Millipore Cat. No. A8456) is used at 1 mg/mL in blood.
3. Aprotinin (Trasylol) used at 500 KIU /mL of blood, (Merck Millipore Cat. No. 7107-01).
4. Protease Inhibitor Cocktail (Merck Millipore Cat. No. P2714).
5. Active and Total cannot be run together in the same assay.

Appendix 3: Other Sample Types

Protocols Using Other Sample Types

Sample Type	Species/Kit Run	Procedure	Reference (if available)
Adipose Tissue Homogenates	Human	Approximately 4 g of adipose tissue from each subject was homogenized in 16 mL of ice-cold deoxygenated homogenization buffer containing 10% glycerol, 150 mM NaCl, 2 mM EDTA, 1 mM PMSF, 25 mM benzamidine, 10 µM leupeptin, 2.5 µmol/L pepstatin A, and 50 U/mL aprotinin in 10 mM Tris-HCl (pH 7.0), with four up/down strokes at Setting No. 3 using a Polytron (Brinkmann Instruments, Inc., Westbury NY). The crude homogenate was centrifuged at 3,000 x g for 15 minutes, and the fat cake was discarded. The infranate was made up to 1% (vol/vol). Triton® X-100 was used to solubilize PTPase enzymes from the particulate compartment into the tissue homogenate. The supernatant resulting from centrifugation at 15,000 x g for 20 minutes at 4 °C was stored in aliquots at 80 °C.	J Clin Endocrinol Metab. 2001 Dec;86(12):5973-80. PMID:11739472
Adipose Tissue Homogenate	Human/ Apolipoprotein Panel	Adipose biopsies (50–75 mg) were homogenized on ice in 1 mL of the kit assay buffer. (10 mmol/L PBS, 0.08% (wt./vol.) sodium azide, 1% (wt./vol.) BSA, pH 7.4). The homogenate was further diluted 25-fold in assay buffer to minimize assay interferences. 10 µL of dilute homogenate was incubated in a 96-well plate with 25 µL of capture antibody-conjugated beads and 65 µL assay buffer for 1 hour ambient. Beads were washed (10 mmol/L PBS, 0.05% (vol./vol.) Proclin, 0.05% (vol./vol.) Tween-20, pH 7.4) and 50 µL biotinylated detection antibody cocktail added for 30 minutes ambient, followed by further washing and incubation with 50 µL streptavidin-phycoerythrin for 30 minutes at ambient temperature. After final washing, beads were resuspended in 150 µL Luminex® sheath fluid for analysis.	Diabetologia. 2008 Nov;51(11):2041-8. doi: 10.1007/s00125-008-1126-5. Epub 2008 Aug 19 PMID:18712345
Adipose Tissue Extract	Human/CVD Panel	Approximately 100–200 mg adipose tissue (SAT and VAT) from each subject was homogenized in 250 µL of ice-cold homogenization buffer. The homogenate was centrifuged at 3,000 x g for 15 minutes at 4 °C, the fat cake was discarded and the homogenate was centrifuged again at 14,000 x g for 20 minutes at 4 °C. The supernatant was stored in aliquots at -70 °C.	Physiol Res. 2010;59(1):79-88. Epub 2009 Feb 27. PMID:19249917
Aorta Tissue Extract	Guinea Pig/ Human Cytokine/ Chemokine Panel 1	The vessel was dissected and all the surrounding tissues removed. The vessel was mixed and homogenized in a rotor-stator with 1 mL of lysis buffer (0.1 g of bovine serum albumin, 5 µL of Triton® X-100, 100 mg of gentamycin sulfate, 100 µL of HEPES buffer-1M, 23 µL of aprotinin, 18.391 mg of sodium orthovanadate and PBS to complete 1 mL). After this, 2 mL of the lysis buffer was added to the content and was homogenized in a Potter-Elvehjem tissue grinder. This was centrifuged at 400 x g for 10 minutes at 4 °C. The supernatant was analyzed.	BMC Cardiovasc Disord. 2009 Feb 17;9:7. doi: 10.1186/1471-2261-9-7. PMID:1922285
Brain Tissue Extract	Rat/Rat Cytokine	Plasma and brain tissue from injured (hyperintense tissue on DW-MRI during occlusion) and anatomically matching tissue from the contralateral hemisphere were collected from control and minocycline- or PBS-treated rat pups following 24 hours of reperfusion. The flash-frozen brain tissue was homogenized in a buffer containing 20 mmol/L Tris-HCl (pH 7.5), 150 mmol/L NaCl, 1 mmol/L PMSF, 0.05% Tween® 20, and a cocktail of protease inhibitors (Roche), and protein concentration was measured in each sample.	J Cereb Blood Flow Metab. 2005 Sep;25(9):1138-49 PMID: 15874975
Bronchoalveolar Lavage (BAL) Samples		For lavage samples, use 50 µL sample + 25 µL beads in sample wells. Set up standards using one additional lower point and dropping the highest concentration standard point. Use a buffer matrix or medium used to collect the lavage sample as the matrix, i.e., 25 µL standard/control/blank + 25 µL assay buffer /medium + 25 µL beads. The first incubation with standard/sample should be overnight, 4 °C. Final results should be divided by 2.	
Dried Blood Spot Samples		Two 3.2-mm (1/8-inch) diameter disks were punched from dried blood-spot calibrators or controls and eluted in 100 µL of 0.075 mol/L sodium barbital buffer (pH 8.6) containing 0.5 g/L anilinnaphthalenesulfonic acid and 0.5 g/L sodium azide by sonication at room temperature for 30 minutes. The volume of blood per 3.2-mm disk was 3 µL (3). The eluate was filtered in a 0.45 µm centrifugal filter unit (Merck Millipore).	
Dried Blood Spot Samples		Whole blood samples were "spotted" onto Whatman 3 mm filter paper, air-dried, and stored at 4 °C prior to extraction and testing. Areas equivalent to a 25-µL drop were punched from the filter paper and eluted in 25 µL of 0.01 M phosphate buffer, pH 7.4 prior to analysis. The protein content of each eluate was measured spectrophotometrically at 260/280 nm and the samples normalized to a standard protein content of 1 µg/mL.	J Chromatogr B Biomed Sci Appl. 1998 Sep 11;715(1):55-63. PMID: 9792497

* While these alternate methods have been tried by Merck Millipore, or our end users using MILLIPLEX® MAP kits, we cannot guarantee methods will work with all samples. These procedures have not been analytically validated.

Sample Type	Species/Kit Run	Procedure	Reference (if available)
Cervical Secretions, Vaginal Secretions, and Saliva Secretions		Saliva, cervical and vaginal secretions were collected using ophthalmic sponges (Wek-Cel, Xomed Treace, Orlando, FL) after exposure of the cervical os with the speculum. The secretions were collected by placing the ophthalmic sponge directly into the cervical os and allowing it to absorb secretions for approximately 1 minute. Vaginal secretions were collected by placing the ophthalmic sponge against the vaginal wall and allowing the sponge to collect secretions. In a similar fashion, saliva was collected by placing the ophthalmic sponge over the parotid duct and allowing the sponge to absorb saliva. All sponges were immediately placed on ice and then frozen at -20 °C. The secretions were extracted from the sponges just prior to analysis. Each individual sponge was weighed to determine the volume of secretions absorbed into the sponges. The sponges were then equilibrated in 300 µL phosphate-buffered saline (PBS) + 0.25 M NaCl with 10% fetal calf serum for 30 minutes at 4 °C. The secretions were separated using a spin-x centrifuge filter unit (Costar, Cambridge, MA), centrifuged at 12,000 x g rpm for 20 minutes. A dilution factor for the final extract was determined based on the following formula: dilution factor = [(x-0.0625 mL) + 0.3 mL buffer]/x-0.0625 mL], where x equals the volume of material collected and 0.06 equals the weight of the dry spear (mg=mL). (Note: The weight of the dry sponge is dependent on the lot number. Each lot must be weighed.) This dilution factor was used to calculate the final units of specific antibody and total immunoglobulin measured.	J Clin Immunol. 1997 Sep;17(5):370-9. PMID: 9327336
Colorectal Tissue Extracts	Human/Human Cytokine Panel 1	Normal and cancer tissue specimen weights were determined before protein extraction with Tissue Protein Extraction Reagent (T-PER; Pierce, Rockford, USA) as recommended by the manufacturer. Briefly, 20 mL of P-TER was added to 1 g of tissue and homogenized. Samples were centrifuged at 10,000 x g for 5 minutes and the supernatant (protein extract) was stored at -80 °C until cytokine/chemokine profiling.	Gut. 2009 Apr;58(4):520-9. doi: 10.1136/gut.2008.158824. Epub 2008 Nov 20. PMID: 19022917
Ear Lysates	Mouse/Mouse Cytokine Panel 1	Skin ear biopsies were pooled from four test animals. Biopsies were minced and then repeatedly homogenized with beads in phosphate buffered saline (PBS) plus Complete Protease Inhibitor Cocktail (Roche Applied Science, Indianapolis, IN) at 4 °C. Supernatant was collected and analyzed for the presence of cytokines using a Luminex® instrument with a Mouse Cytokine Panel (Merck Millipore) as per the manufacturer's protocol (MILLIPLIX® MAP Mouse Cytokine/Chemokine Magnetic Bead Panel, Cat. No. MCYTOMAG-70K).	J Invest Dermatol. 2010 Apr;130(4):1023-33. doi: 10.1038/jid.2009.358. Epub 2009 Nov 12. PMID: 19907432
Ear Lysates	Mouse/Mouse Cytokine Panel 1	Ear tissue from mice treated with vehicle or R348 (120 mg/kg) were harvested after 6 weeks of treatment and snap-frozen in liquid nitrogen. Ears were homogenized under liquid nitrogen with radioimmunoprecipitation assay buffer supplemented with protease and phosphatase inhibitors, centrifuged at 13,000 rpm, and the supernatant was collected to generate ear lysates. Ear lysates were normalized for equal protein concentration.	J Immunol. 2009; 183(3):2183-92. PMID: 19596999
Infectious Samples		For infectious samples: If washing with an automatic plate washer, add 30% bleach to the waste bottle before washing/aspirating the plate. If washing with a hand held magnetic bead separator, add 30% bleach to a container capable of catching the wash solution decanted from the plate. Then at the end of the assay, resuspend the beads in 0.1ml of 4% formaldehyde made in 10mM PBS (prepared fresh daily) instead of sheath fluid, before running the plate in the Luminex® machine. Prolonged incubation in this solution may cause bead aggregation. Consequently, after agitating the plate for 5 minutes on an orbital plate shaker, read the plate immediately.	
Jejunal Extracts	Human/Human Cytokines	Jejunal biopsy specimens were fixed with formalin or embedded in optimal-cutting-temperature (OCT) compound and snap frozen in liquid nitrogen. Protein extracts were prepared from jejunal biopsies embedded in OCT compound by washing them twice with a phosphate-buffered saline lysis buffer containing 0.05% sodium azide, 0.5% Triton® X-100, 1 mM phenylmethylsulfonyl fluoride, and protease inhibitors (Complete Mini protease inhibitor cocktail; Roche Diagnostics, Indianapolis, IN). After OCT compound removal, the tissues were minced in 1 mL of lysis buffer with a sterile disposable homogenizer on ice for 5 minutes. The homogenates from the tissues were then sonicated for 1 minute on ice. After centrifugation at 10,000 x g for 15 minutes, the supernatant was collected and stored at -80 °C or immediately assayed to determine the protein concentration with a bicinchoninic acid protein assay kit (Pierce, Rockford, IL).	Infect Immun. 2007 Jan;75(1):481-7. Epub 2006 Oct 16. PMID: 17043107
Lipemic Samples		For lipemic and plasma samples, the blood needs to be collected on ice, centrifuged in a refrigerated centrifuge, aliquoted and frozen at -20 °C for short term (<2 months) and -70 °C for long term. Prior to assay setup, thaw samples and centrifuge at 10,000 rpm for 5 minutes. Spool off the lipid layer from the surface using a cotton swab and use the supernatant below lipid layer for the assay.	
Lymph Node Homogenates	Mouse/Mouse Cytokine Panel 2	Footpad popliteal lymph nodes from mouse subjects were harvested, combined and placed in 200 µL of PBS containing 1 x protease inhibitors (Roche). The lymph nodes were mechanically homogenized with a pestle, followed by centrifugation at 4 °C. Supernatant was transferred to another tube and frozen on dry ice.	Circ Res. 1976 Nov;39(5):659-65. PMID: 184975
Saliva	Human	Add Merck Millipore protease inhibitor cocktail at 1:500 to saliva. Centrifuge at 10K rpm 10 minutes and dilute supernatant 1:2 with assay buffer prior to assay setup. This method significantly improves recovery and reduces bead aggregation. Run assay with assay buffer as matrix in standard curve. Use an overnight option if available.	

Sample Type	Species/Kit Run	Procedure	Reference (if available)
Skin Extracts	Human/Human Cytokine Panel 1	Quantitation of multiple proteins in extracts of D-Squame® tape samples of human scalp using a multiplex skinMAP (multiple analyte profile) immunoassay: D-Squame® tape strip samples of human scalp skin were extracted with PBS containing 0.2% SDS and 0.5% propylene glycol (PG) for 30 minutes with sonication on ice. The extracts were then centrifuged for 5 minutes at 2,100 × g to remove skin solids that might interfere in the assay. Subsequently, the extracts of D-Squame® tape samples were transferred into 96-well polypropylene deep-well plates and frozen at -80 °C for Human Skin Panel (Cat. No. SKINMAG-50K) and soluble protein analyses as previously reported.	Int J Dermatol. 2011 Jan;50(1):102-13. doi: 10.1111/j.1365-4632.2010.04629.x. PMID: 21182510
Tears	Human/Human Cytokines	Polyurethane minisponges were obtained commercially (PeleTim; VOCO GmbH, Cuxhaven, Germany). Specialized nurses carried out the procedures. A single polyurethane minisponge was laid on the outer third of the lower eyelid margin. After 5 minutes of tear collection, the sponge was recovered and placed in the narrow end of a truncated Gilson micropipette tip adapted to a 1.5 mL tube (Eppendorf, Fremont, CA) and centrifuged at 6,000 rpm for 5 minutes. Tear samples from both eyes were pooled and immediately stored at -80°C until they were used for the immunoassay.	
Tears	Human	Tear collection was performed before any other test and with a minimum of 10 minutes after the patient answered the two symptom questionnaires. Unstimulated tear samples were collected non-traumatically from the external canthus of open eyes, avoiding additional tear reflex as much as possible. Glass capillary micropipettes (Drummond, Broomall, PA) were used to collect 1 µL of tears. Each sample was then diluted 1:10 in a sterile collection tube containing ice-cold Cytokine Assay Buffer (Merck Millipore). Tubes with tear samples were kept cold (4 °C) during collection, and stored at -80 °C until assayed.	Mol Vis. 2010 May 19;16:862-73. PMID: 20508732
Cell or Tissue Extraction		Protocol varies depending on tissue types and/or analytes of interest. Generally, most protocols that are used in ELISAs can be used, but here are some guidelines in selecting a method. 1) Homogenize cells or tissues mechanically (eg. ultrasonication) in a PBS based buffer containing protease inhibitors (like aprotinin or an inhibitor cocktail) and low (< 0.2%) non-ionic detergent concentration. 2) Extraction medium should not contain any organic solvents like DMSO, etc. 3) Centrifuge the extract and freeze supernatant at <-20 °C. 4) Use the extraction medium as matrix in blank, standard curve and QCs.	
Tumor Homogenates	Mouse/Mouse Cytokines	Tumors were treated with DMXAA. After harvesting at 6, 24, and 48 hours, the tumors were sonicated for 30 seconds in 1 mL of complete buffer (50 mL PBS containing one tablet of antiprotease cocktail, Roche, Indianapolis, IN). Tissues were then spun at 3,000 rpm for 10 minutes and filtered through a 1.2 µm syringe filter unit. Total protein in each sample was determined.	Cancer Res. 2005 Dec 15;65(24):11752-61. PMID: 16357188
Urine		Typically, measurement of analytes in urine requires either a 24 hours urine collection or second morning void collection. For the second morning void urine, the analyte value is normalized against creatinine, i.e., the analyte is expressed as units/mg of creatinine. Mix urine samples 1:1 with assay buffer and incubate on the plate approximately 20 minutes on a shaker prior to addition of the beads. Use assay buffer as matrix for standard curve, controls and blank. The assumption is that this helps neutralize the sample, thereby improving recovery.	

* While these alternate methods have been tried by Merck Millipore, or our end users using MILLIPLEX® MAP kits, we cannot guarantee methods will work with all samples. These procedures have not been analytically validated.

Appendix 4: Cell Signaling MAPmate™ Buffer Table

Species Cross-Reactivity and Buffer Compatibility

Magnetic Bead MAPmate™ Kits	Cat. No.	Species Homology	AB1	AB2
β-Tubulin (Total)	46-713MAG	H,M,R	●	●
GAPDH (Total)	46-667MAG	H	○	●
Akt/PKB (assay buffer 1) (Ser473)	46-601MAG	H,M,R	●	○
Akt/PKB (assay buffer 1) (Thr308)	46-645MAG	H,M,R	●	○
Akt/PKB (assay buffer 1) (Total)	46-605MAG	H,M,R	●	○
Akt/PKB (assay buffer 2) (Ser473)	46-677MAG	H,M,R	●	●
Akt/PKB (assay buffer 2) (Total)	46-675MAG	H,M,R	●	●
BAD (Ser112)	46-694MAG	H,M	●	●
BAD (Total)	46-695MAG	H,M	●	●
Caspase 3 (Active)	46-604MAG	H,M	●	●
c-Jun (Ser73)	46-622MAG	H,M,R	●	●
c-Kit (pan Tyr)	46-619MAG	H	●	●
c-Kit (Total)	46-620MAG	H	●	●
c-Met/HGFR (pan Tyr)	46-651MAG	H	●	●
c-Met/HGFR (Total)	46-650MAG	H	●	●
CREB (Ser133)	46-631MAG	H,M,R	●	●
CREB (Total)	46-632MAG	H,M,R	●	●
EGF Receptor (pan Tyr)	46-603MAG	H	●	●
EGF Receptor (Total)	46-606MAG	H	●	●
ERK/MAPK 1/2 (Thr185/Tyr187)	46-602MAG	H,M,R	○	●
ERK/MAPK 1/2 (Total)	46-609MAG	H,M,R	●	●
GSK3β (Ser9)	46-690MAG	H,M,R	●	●
GSK3β (Total)	46-689MAG	H,M,R	●	●
H2A.X (Ser139)	46-692MAG	H,M	●	●
HIF-1α (Total)	46-665MAG	H,M,R	●	●
HSP27 (Ser78)	46-607MAG	H	●	●
HSP27 (Total)	46-608MAG	H	●	●
IRS1 (pan Tyr)	46-627MAG	H,M,R	●	●
IRS1 (Total)	46-628MAG	H,M,R	●	●
IκBα (Ser32)	46-643MAG	H	●	●
IκBα (Total)	46-644MAG	H	●	●
JNK/SAPK1 (Thr183/Tyr185)	46-613MAG	H,M	●	●
JNK/SAPK1 (Total)	46-618MAG	H,M	○	●
Lck (Tyr394)	46-712MAG	H,M,R	●	●
MEK1 (Ser222)	46-670MAG	H,M,R	●	●
MEK1 (Total)	46-669MAG	H,M	●	●
mTOR (Ser2448)	46-686MAG	H,M,R	●	●
mTOR (Total)	46-685MAG	H,M,R	●	●
NFκB (Ser536)	46-702MAG	H	●	●
NFκB (Total)	46-701MAG	H	●	●
p21 (Total)	46-621MAG	H	●	●
p38α/SAPK2 (Thr180/Tyr182)	46-610MAG	H,M,R	○	●
p38α/SAPK2 (Total)	46-612MAG	H,M,R	○	●
p53 (Ser15)	46-663MAG	H	●	●
p53 (Total)	46-662MAG	H	●	●
p70S6K (Thr389/412)	46-629MAG	H,M,R	●	●
p70S6K (Total)	46-630MAG	H,M,R	●	●
Cleaved PARP (Total)	46-656MAG	H	●	●
PTEN (Ser380)	46-679MAG	H,M,R	●	●
PTEN (Total)	46-678MAG	H,M,R	●	●
RPS6 (Ser235/Ser236)	46-714MAG	H,M,R	●	●
RPS6 (Total)	46-715MAG	H,M,R	●	●
Src (Tyr419)	46-710MAG	H,M,R	●	●
Src (Total)	46-709MAG	H,M,R	●	●
STAT1 (Tyr701)	46-655MAG	H,M	●	●
STAT1 (Total)	46-654MAG	H,M	●	●
STAT3 (Tyr705)	46-623MAG	H,M,R	●	●
STAT3 (Ser727)	46-624MAG	H,M,R	●	●
STAT3 (Total)	46-625MAG	H,M,R	●	●
STAT5A/B (Tyr694/Tyr699)	46-641MAG	H,M,R	●	●
Tie2 (pan Tyr)	46-716MAG	H	●	●
Tie2 (Total)	46-717MAG	H	●	●
VEGFR2 (pan Tyr)	46-718MAG	H	●	●
VEGFR2 (Total)	46-719MAG	H	●	●

●	Recommended
●	Acceptable
○	Not Recommended
H	Human
M	Mouse
R	Rat
AB1:	Assay Buffer 1
AB2:	Assay Buffer 2

Glossary*

Accuracy: Data representing mean percent recovery of spiked standards ranging from low, medium and high concentrations in serum matrices for a defined number of samples (see Spike Recovery).

Analyte: A chemical substance that is the subject of chemical analysis.

Configurable/customizable kit: A type of MILLIPLEX® MAP kit that enables the customer to choose the analytes within a specific panel that best meet her research needs (e.g., Cat. No. HCYTOMAG-60K, Human Cytokine/Chemokine Panel 1).

Drive fluid: Luminex® Drive Fluid is the delivery medium to transport the sample to the optic components of the MAGPIX® system.

Fixed kit: A type of MILLIPLEX® MAP kit that is not configurable. All of the analytes are sold together with the capture beads already premixed (e.g., Cat. No. 48-611, Akt/mTOR Phosphoprotein – 11 Plex).

FLEXMAP 3D® system: A Luminex® instrument that combines differentially dyed fluorescent microsphere sets with an innovative instrument design to enable precise, rapid multiplexing of up to 500 unique assays within a single sample. Other features include an automated probe height adjustment, simplified routine maintenance operations and an intuitive software interface.

Inter-assay precision (%CV): Precision generated across two different concentrations of analytes across a defined number of different assays.

Intra-assay precision (%CV): Precision generated across two different concentrations of analytes in a single assay.

Linearity: The ability (within a given range) to obtain test results which are directly proportional to the concentration (amount) of analyte in the sample.

Luminex 200™ system: A Luminex® instrument that provides a complete mid- to high-range solution for rapid, accurate biomarker quantification. The Luminex XY Platform™ (Luminex XYP™) complements this instrument system by automating the sequential positioning of each well of a microtiter plate.

Magnetic beads (MagPlex®): Similar to MicroPlex® microspheres, MagPlex® microspheres are carboxylated polystyrene micro-particles, or "beads," that have been dyed into spectrally distinct sets, or "regions"; allowing them to be individually identified by a Luminex® instrument. These uniquely coded beads provide a user an addressable substrate on which to perform multiple bio-analytical reactions in a single well.

MAGPIX® system: A Luminex® instrument based on CCD imaging technology, which allows for a compact, more robust system. Streamlined startup and shutdown protocols and minimal maintenance requirements make the system easy to operate and maintain.

MFI: Median fluorescence intensity.

MILLIPLEX® 5.1 Analyst software: Merck Millipore analysis software package that is able to automatically import data from Luminex® instruments, providing better data from the low and high ends of standard curves and comprehensive, detailed reports and enhanced visualization.

MILLIPLEX® MAP: A broad portfolio of multiplex immunoassays that includes immunology, metabolism, cardiovascular disease, cancer, neuroscience, toxicity, cellular metabolism and cell signaling pathways for a variety of species.

MinDC (minimum detectable concentration): The lowest concentration at which an analyte can be reliably detected.

Precision: Generated from the mean of the %CVs from a defined number of reportable results.

Premixed kit: A MILLIPLEX® kit in which the capture beads have been mixed together. There is often an option of choosing either a premixed kit or a configurable kit (e.g., FCYTOMAG-20K-PMX, Feline Cytokine/Chemokine 19-plex Premix).

Quality controls (QCs): Medium-low and medium-high standard points used to qualify assay performance.

SAPE: Streptavidin-phycoerythrin.

Serum matrix: An appropriately selected matrix found only in MILLIPLEX® kits that is added to the standard wells to mimic the environment in which native analytes are present in serum/plasma. The selected matrix most often consists of serum/plasma pool with all endogenous and cross-reacting proteins extracted.

Sheath fluid (Luminex® catalog number 40-50000): Luminex® Sheath Fluid is intended for use as the delivery medium of the sample to the optics component of the Luminex 100/200™ system and the FLEXMAP 3D® system.

Spike recovery: Data representing mean percent recovery of spiked standards ranging from low, medium and high concentrations in serum matrices for a defined number of samples (see Accuracy).

Stability: Resistance or the degree of resistance to chemical change or disintegration.

Standard curve/Calibration curve: A graphic plot of median fluorescence intensity versus the known concentration of test substances in a set of standards usually prepared by serial dilution or incremental addition.

xMAP® (Multi-Analyte Profiling) technology: Flexible, open-architecture design that can be configured to perform a wide variety of bioassays developed by the Luminex Corporation.

xPONENT® software: Luminex® acquisition software with analysis capabilities.

* Sources include: thefreedictionary.com; merriam-webster.com; luminexcorp.com; scientistsolutions.com; regulatory.com; dictionary.com; and MILLIPLEX® protocol.

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