# MILLIPLEX<sup>™</sup> MAP

# HUMAN CARDIOVASCULAR DISEASE (CVD) SINGLE PLEX HAPTOGLOBIN KIT 96 Well Plate Assay

#### # HCVD2-67BK-1HAP

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

#### INTRODUCTION

Patients with diabetes are at two to four times higher risk for cardiovascular disease than non-diabetics. One of the key factors in the development of arteriosclerosis, an underlying cause of CVD, is the oxidative modification of LDL molecules by reactive oxygen species (ROS). A plasma glycoprotein synthesized in the liver, haptoglobin's primary function is the binding of free or extracellular hemoglobin, a strong oxidizing agent released by red blood cells. Haptoglobin inhibits extracellular hemoglobin's ability to cause oxidative tissue damage by preventing the release of heme iron. Once hemoglobin is bound to haptoglobin, the complex is rapidly cleared from the bloodstream by either hepatocytes or monocytes.

Studies have shown that the severity of CVD risk in diabetics can be determined by the genetic alleles that encode the haptoglobin protein. The three different haptoglobin protein phenotypes inhibit hemoglobin-induced oxidation at different rates, with the weakest phenotype linked to the greatest risk of CVD. In addition, a diabetic hyperglycemic environment is essential for the different expressions of these proteins.

Millipore provides important tools to understand better the role of such biomarkers as haptoglobin in the relationship between diabetes and cardiovascular disease. Based on the Luminex xMAP platform, the MILLIPLEX MAP Haptoglobin Single Plex, together with other Millipore multiplex and single plex panels, assists in the investigation of the modulation and expression of dozens of analytes, giving you the advantage of speed and sensitivity.

This kit is for research purposes only.

Please read entire protocol before use.

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

#### **PRINCIPLE**

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

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

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

#### STORAGE CONDITIONS UPON RECEIPT

- Recommended storage for kit components is 2 8 ℃.
- Once the standards and controls have been reconstituted, immediately transfer contents into polypropylene vials. DO NOT STORE RECONSTITUTED STANDARDS OR CONTROLS IN GLASS VIALS. For long-term storage, freeze reconstituted standards and controls at ≤ -20 °C. Avoid multiple (>2) freeze/thaw cycles.
- DO NOT FREEZE Antibody-Immobilized Beads, Detection Antibodies, and Streptavidin-Phycoerythrin.

# **REAGENTS SUPPLIED**

Note: Store all reagents at 2-8 °C

REAGENTS SUPPLIED	CATALOG NUMBER	VOLUME	QUANTITY
Human CVD Panel 2 Standard	HCVD2-8067-2	lyophilized	1 vial
Human CVD Panel 2 Quality Controls 1 and 2	HCVD2-6067-2	lyophilized	2 vials
Set of one 96-Well Filter Plate with 2 Sealers	MX-PLATE		1 plate 2 sealers
Assay Buffer	L-AB	30 mL / bottle	2 bottles
10X Wash Buffer Note: Contains 0.05% Proclin	L-WB	30 mL	1 bottle
Human CVD Panel 2 Detection Antibodies	HCVD2-1067-2	3.2 mL	1 bottle
Streptavidin-Phycoerythrin	L-SAPE6	3.2 mL	1 bottle
Bead Diluent	LBD	3.5 mL	1 bottle
Mixing Bottle			1 bottle

Bead / Analyte Name	Luminex Bead (20X concentration, 20)		icentration, 200μL)
beau / Analyte Name	Region	Available	Cat. #
Anti – Human Haptoglobin Beads	51	<b>√</b>	HHPTGN

#### MATERIALS REQUIRED BUT NOT PROVIDED

#### Reagents

1. Luminex Sheath Fluid (Luminex Catalogue #40-50000)

## Instrumentation / Materials

- 1. Adjustable Pipettes with Tips capable of delivering 5 μL to 1000 μL
- 2. Multichannel Pipettes capable of delivering 5 µL to 50 µL or 25 µL to 200 µL
- 3. Reagent Reservoirs
- 4. Polypropylene Microfuge Tubes
- 5. Rubber Bands
- 6. Absorbent Pads
- 7. Laboratory Vortex Mixer
- 8. Sonicator (Branson Ultrasonic Cleaner Model #B200 or equivalent)
- 9. Titer Plate Shaker (Lab-Line Instruments Model #4625 or equivalent)
- 10. Vacuum Filtration Unit (Millipore Vacuum Manifold Catalog #MSVMHTS00 or equivalent with Millipore Vacuum Pump Catalog #WP6111560 or equivalent)
- 11. Luminex 100™ IS, 200™, or HTS by Luminex Corporation
- 12. Plate Stand (Millipore Catalog # MX-STAND)

#### SAFETY PRECAUTIONS

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

#### **TECHNICAL GUIDELINES**

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

- FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES.
- Do not use beyond the expiration date on the label.
- Do not mix or substitute reagents with those from other lots or sources.
- The Antibody-Immobilized Beads are light sensitive and must be protected from light at all times. Cover the assay plate containing beads with opaque plate lid or aluminum foil during all incubation steps.
- It is important to allow all reagents to warm to room temperature (20-25°C) before use in the assay.

- The bottom of the Microtiter Filter Plate should not be in direct contact with any surface during assay set-up or incubation times. The plate can be set on a plate stand or on the non-flat side of the plate cover or any other plate holder to raise the plate from the surface. A plate stand can be purchased separately from Millipore (Millipore Catalog #MX-STAND).
- Incomplete washing can adversely affect the assay outcome. All washing must be performed with the Wash Buffer provided.
- After the wash steps, keep the bottom of the Microtiter Filter Plate clean by blotting on paper towels or absorbent pads to prevent any leakage due to capillary action.
- Keep the vacuum suction on the plate as low as possible. It is recommended to have a vacuum setting that will remove 200 µL of buffer in ≥ 5 seconds (equivalent to < 100 mmHg).
- After hydration, all Standards and Controls must be transferred to polypropylene tubes.
- The Standards prepared by serial dilution must be used within 1 hour of preparation.
   Discard any unused standards except the standard stock which may be stored at
   ≤ -20°C for 1 month and at ≤ -80°C for greater than one month.
- If samples fall outside the dynamic range of the assay, further dilute the samples with the appropriate diluent and repeat the assay.
- Any unused mixed Antibody-Immobilized Beads may be stored in the Mixing Bottle at 2-8°C for up to one month.
- During the preparation of the standard curve, make certain to mix the higher concentration well before making the next dilution. Use a new tip with each dilution.
- The plate should be read immediately after the assay is finished. If, however, the plate cannot be read immediately, seal the plate, cover with aluminum foil or an opaque lid, and store the plate at 2-8°C for up to 24 hours. Prior to reading, agitate the plate on the plate shaker at room temperature for 10 minutes. Delay in reading a plate may result in decreased sensitivity for some analytes.
- The titer plate shaker should be set at a speed to provide maximum orbital mixing without splashing of liquid outside the wells. For the recommended plate shaker, this would be a setting of 5-7 which is approximately 500-800 rpm.
- Ensure that the needle probe is clean. This may be achieved by sonication and/or alcohol flushes. Adjust probe height according to the protocols recommended by Luminex to the kit filter plate using 3 alignment discs prior to reading an assay.
- For cell culture supernatants or tissue extraction, use the culture or extraction medium as the matrix solution in background, standard curve and control wells. If samples are diluted in Assay Buffer, use the Assay Buffer as matrix.
- For serum/plasma samples, use the Serum Matrix provided in the kit.
- For cell/tissue homogenate, the final cell or tissue homogenate should be prepared in a buffer that has a neutral pH, contains minimal detergents or strong denaturing detergents, and has an ionic strength close to physiological concentration. Avoid debris, lipids, and cell/tissue aggregates. Centrifuge samples before use.
- Vortex all reagents well before adding to plate.

#### SAMPLE COLLECTION AND STORAGE

## A. Preparation of Serum Samples:

- Allow the blood to clot for at least 30 minutes before centrifugation for 10 minutes at 1000xg. Remove serum and assay immediately or aliquot and store samples at ≤ -20 °C.
- Avoid multiple (>2) freeze/thaw cycles.
- When using frozen samples, it is recommended to thaw the samples completely, mix well by vortexing and centrifuge prior to use in the assay to remove particulates.
- Serum sample dilution: Concentrations of acute phase proteins can change dramatically depending upon specific treatments or pathological conditions. Customers may need to determine the optimal dilution factors for their samples depending on the expected biological range. In general, samples from normal subjects should be diluted 1:40,000 for quantification of Haptoglobin. L-AB Assay Buffer provided in the kit should be used as the sample diluent.
- Sample dilutions can be done in two steps. For samples diluted at 1:40,000: Step 1: add 2.5 μL serum to 497.5 μL Assay Buffer (1:200); Step 2: add 2.5 μL of 1:200 diluted sample to 497.5 μL of Assay Buffer (1:40,000).

## B. Preparation of Plasma Samples:

- Plasma collection using EDTA as an anticoagulant is recommended.
   Centrifuge for 10 minutes at 1000xg within 30 minutes of blood collection.
   Remove plasma and assay immediately or aliquot and store samples at < -20°C.</li>
- Avoid multiple (>2) freeze/thaw cycles.
- When using frozen samples, it is recommended to thaw the samples completely, mix well by vortexing and centrifuge prior to use in the assay to remove particulates.
- Plasma sample dilution: Concentrations of acute phase proteins can change dramatically depending upon specific treatments or pathological conditions. Customers may need to determine the optimal dilution factors for their samples depending on the expected biological range. In general, samples from normal subjects should be diluted 1:40,000 for quantification of Haptoglobin. L-AB Assay Buffer provided in the kit should be used as the sample diluent.
- Sample dilutions can be done in two steps. For samples diluted at 1:40,000: Step 1: add 2.5 μL plasma to 497.5 μL Assay Buffer (1:200); Step 2: add 2.5 μL of 1:200 diluted sample to 497.5 μL of Assay Buffer (1:40,000).

## C. Preparation of Tissue Culture Supernatant:

- Centrifuge the sample to remove debris and assay immediately or aliquot and store samples at ≤ -20 °C.
- Avoid multiple (>2) freeze/thaw cycles.
- Tissue culture supernatant may require a dilution with an appropriate control medium prior to assay. Users need to provide the control medium as the sample diluent.

#### NOTE:

- A maximum of 25 µL per well of tissue extract, cell / tissue culture supernatant, diluted serum, or diluted plasma samples can be used. Samples may require appropriate dilutions before the assay. Dilution factor varies with sample types and treatment.
- All samples must be stored in polypropylene tubes. DO NOT STORE SAMPLES IN GLASS.
- Avoid debris, lipids and cells when using samples with gross hemolysis or lipemia.
- Care must be taken when using heparin as an anticoagulant since an excess of heparin will provide falsely high values. Use no more than 10 IU heparin per mL of blood collected.

#### PREPARATION OF REAGENTS FOR IMMUNOASSAY

## A. <u>Preparation of Antibody-Immobilized Beads</u>

Prepare bead dilution right before setting up the assay. Early preparation of the bead dilution (>1 hour) may result in high background signal for haptoglobin.

Sonicate the antibody-bead vial for 30 seconds; vortex for 1 minute. Add 150  $\mu$ L from the antibody bead vial to the Mixing Bottle and bring final volume to 3.0 mL with Bead Diluent. Vortex the mixed beads well. Any remaining unused portion of the bead mixture should be discarded.

## B. Preparation of Quality Controls

Before use, reconstitute Quality Control 1 and Quality Control 2 with 250  $\mu$ L deionized water. Invert the vial several times to mix and vortex. Allow the vial to sit for 5-10 minutes and then transfer the controls to appropriately labeled polypropylene microfuge tubes. Unused portion may be stored at  $\leq$  -20°C for up to one month.

#### C. Preparation of Wash Buffer

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

#### D. Preparation of Standard

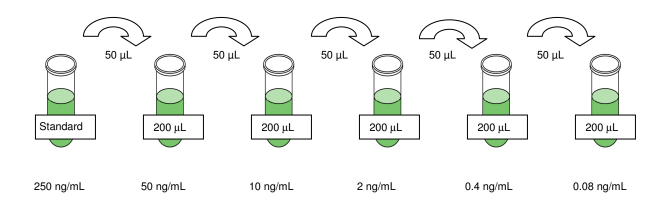
1.) Prior to use, reconstitute the Standard with 250 µL Deionized Water to give a 250 ng/mL final concentration. Invert the vial several times to mix. Allow the vial to sit for 5-10 minutes to make sure that the standard is completely reconstituted, and then transfer the standard to an appropriately-labeled polypropylene microfuge tube. This will be used as the stock standard (250 ng/mL). The unused portions of this stock may be stored at ≤ -20°C for up to one month.

## 2.) Preparation of Working Standards

Label five polypropylene microfuge tubes 50, 10, 2, 0.4, and 0.08 ng/mL. Add 200  $\mu$ L of Assay Buffer to each of the five tubes. Prepare serial dilutions by adding 50  $\mu$ L of the 250 ng/mL reconstituted standard to the 50 ng/mL tube, mix well and transfer 50  $\mu$ L of the 50 ng/mL standard to the 10 ng/mL tube, mix well and transfer 50  $\mu$ L of the 10 ng/mL standard to the 2 ng/mL tube, mix well and transfer 50  $\mu$ L of the 2 ng/mL standard to 0.4 ng/mL tube, mix well and transfer 50  $\mu$ L of the 0.4 ng/mL standard to the 0.08 ng/mL tube and mix well. The 0 standard (Background) will be the Assay Buffer.

Standard Dilution	Volume of Deionized Water to Add	Volume of Standard to Add
250 ng/mL	250 μL	0

Standard Dilution (ng/mL)	Volume of Assay Buffer to Add	Volume of Standard to Add
50	200 μL	50 μL of 250 ng/mL
10	200 μL	50 μL of 50 ng/mL
2	200 μL	50 μL of 10 ng/mL
0.4	200 μL	50 μL of 2 ng/mL
0.08	200 μL	50 μL of 0.4 ng/mL



#### **IMMUNOASSAY PROCEDURE**

- Prior to beginning this assay, it is imperative to read this protocol completely and to thoroughly understand the Technical Guidelines.
- Allow all reagents to warm to room temperature (20-25°C) before use in the assay.
- Diagram the placement of Standards [0 (Background), 0.08, 0.4, 2, 10, 50 and 250 ng/mL], Controls 1 and 2, and Samples on Well Map Worksheet in a vertical configuration. (Note: Most instruments will only read the 96-well plate vertically by default.) It is recommended to run the assay in duplicate.
- Set the filter plate on a plate holder at all times during reagent dispensing and incubation steps so that the bottom of the plate does not touch any surface.
  - 1. Prewet the filter plate by pipetting 200 μL of 1X Wash Buffer into each well of the Microtiter Filter Plate. Seal and mix on a plate shaker for 10 minutes at room temperature (20-25°C).
  - Remove Wash Buffer by vacuum. (NOTE: DO NOT INVERT PLATE.) Blot excess Wash Buffer from the bottom of the plate with an absorbent pad or paper towels.
  - 3. Add 25 µL of each Standard or Control into the appropriate wells. Assay Buffer should be used for the 0 ng/mL standard (Background).
  - 4. Add 25 μL of Assay Buffer to the sample wells.
  - 5. Add 25 µL of Sample into the appropriate wells.
  - 6. Add 25 μL of appropriate matrix solution to the background, standards, and control wells. When assaying tissue/cell extract or tissue/cell culture medium samples, use identical extraction buffer or control medium respectively as the matrix solution and sample diluent. When assaying diluted serum or plasma samples, use the Assay Buffer provided in the kit as the matrix solution and sample diluent.
- Vortex Mixing Bottle and add 25 μL of the Mixed Beads to each well. (Note: During addition of Beads, shake bead bottle intermittently to avoid settling.)
- 8. Seal the plate with a plate sealer, cover it with the lid. Wrap a rubber band around the plate holder, plate and lid and incubate with agitation on a plate shaker for 1 hour at room temperature (20-25°C).

Add 200 µL Wash Buffer per well



Shake 10 min, RT

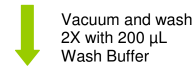
Vacuum

- Add 25 µL Standard or Control to appropriate wells
- Add 25 µL Assay Buffer to background and sample wells
- Add 25 μL Samples to sample wells
- Add 25 µL Matrix to background, standards and control wells
- Add 25 μL Beads to each well



Incubate 1 hour at RT with shaking

- Gently remove fluid by vacuum. (NOTE: DO NOT INVERT PLATE.)
- 10. Wash plate 2 times with 200 μL/well of Wash Buffer, removing Wash Buffer by vacuum filtration between each wash. Blot excess Wash Buffer from the bottom of the plate with an absorbent pad or paper towels.
- 11. Add 25 μL of Detection Antibodies into each well. (Note: Allow the Detection Antibodies to warm to room temperature prior to addition.)
- 12. Seal, cover with lid, and incubate with agitation on a plate shaker for 30 minutes at room temperature (20-25°C). **DO NOT VACUUM AFTER INCUBATION.**
- 13. Add 25 μL Streptavidin-Phycoerythrin to each well containing the 25 μL of Detection Antibodies.
- 14. Seal, cover with lid and incubate with agitation on a plate shaker for 30 minutes at room temperature (20-25°C).
- 15. Gently remove all contents by vacuum. (NOTE: DO NOT INVERT PLATE.)
- 16. Wash plate 2 times with 200  $\mu$ L/well Wash Buffer, removing Wash Buffer by vacuum filtration between each wash. Wipe any excess buffer on the bottom of the plate with a tissue.
- 17. Add 100 μL of Sheath Fluid to all wells.
  Resuspend the beads on a plate shaker for 5 minutes.
- 18. Run plate on Luminex 100™ IS, 200™, or HTS.
- 19. Save and analyze the Median Fluorescent Intensity (MFI) data using a weighted 5-parameter logistic or spline curve-fitting method for calculating concentrations in samples. Note: remember to multiply the results by the sample dilution factor.



Add 25 µL Detection Antibodies per well



Incubate 30 minutes at RT

Do Not Vacuum

Add 25 µL Streptavidin-Phycoerythrin per well



Incubate for 30 minutes at RT

Vacuum and wash 2X with 200 µL Wash Buffer

Add 100 μL Sheath Fluid per well

Read on Luminex (50 μL, 50 beads per bead set)

#### **EQUIPMENT SETTINGS**

These specifications are for the Luminex 100<sup>™</sup> IS v.1.7 or Luminex 100<sup>™</sup> IS v2.1/2.2, Luminex 200<sup>™</sup> v2.3, xPONENT, and Luminex HTS. Luminex instruments with other software (e.g. MasterPlex, StarStation, LiquiChip, Bio-Plex, LABScan100) would need to follow instrument instructions for gate settings and additional specifications from the vendors.

Events:	50, per bead	
Sample Size:	50 μL	
Gate Settings:	8,000 to 15,000	
Reporter Gain:	Default (low PMT)	
Time Out:	60 seconds	
Bead Set:	Haptoglobin	51

#### **QUALITY CONTROLS**

The ranges for each analyte in Quality Control 1 and 2 are provided on the card insert or can be located at the MILLIPORE website <a href="https://www.millipore.com/techlibrary/index.do">www.millipore.com/techlibrary/index.do</a> using the catalog number as the keyword.

#### **ASSAY CHARACTERISTICS**

## Assay Sensitivities (minimum detectable concentrations, ng/mL)

MinDC: Minimum Detectable Concentration is calculated by the StatLIA® Immunoassay Analysis Software from Brendan Technologies. It measures the true limits of detection for an assay by mathematically determining what the empirical MinDC would be if an infinite number of standard concentrations were run for the assay under the same conditions.

Analyte	MinDC (ng/mL)
Haptoglobin	0.052

#### **Precision**

Intra-assay precision is generated from the mean of the %CV's from 8 reportable results across two different concentrations of analytes represented by serum samples in one experiment. Inter-assay precision is generated from the mean of the %CV's from two reportable results each for two different concentrations of analytes across 5 different experiments.

Analyte	Intra-Assay (CV%)	Inter-Assay (CV%)
Haptoglobin	4.4	21.0

## Accuracy

Spike Recovery: The data represent mean percent recovery of 3 levels of spiked standards in serum matrices (0.4, 2.0, and 10.0 ng/mL for each analyte).

Analyte	Serum (1:40,000)
Haptoglobin	93.5

## **Cross-Reactivity**

There is no detectable crossreactivity with other major acute phase proteins in the serum.

Note: Haptoglobin showed high cross-reactivity in <u>non-human primate</u> samples (not fully validated).

# TROUBLESHOOTING GUIDE

Problem	Probable Cause	Solution
Filter plate will not	Vacuum pressure is	Increase vacuum pressure such that 0.2mL
vacuum	insufficient	buffer can be suctioned in 3-5 seconds.
	Samples have insoluble particles	Centrifuge samples just prior to assay set-up and use supernatant.
		If high lipid concentration, after centrifugation, remove lipid layer and use supernatant.
	Sample too viscous	May need to dilute sample.
Insufficient bead count	Vacuum pressure too high	Adjust vacuum pressure such that 0.2mL buffer can be suctioned in 3-5 seconds.
	Bead mix prepared incorrectly	Sonicate bead vials and vortex just prior to adding to bead mix bottle according to protocol. Agitate bead mix intermittently in reservoir while pipetting into the plate.
	Samples cause interference due to particulate matter or viscosity	See above. Also sample probe may need to be cleaned with alcohol flush, backflush and washes; or, if needed, probe should be removed and sonicated.
	Probe height not adjusted correctly	Adjust probe to 3 alignment discs in well H6.
Plate leaked	Vacuum pressure too high	Adjust vacuum pressure such that 0.2mL buffer can be suctioned in 3-5 seconds. May need to transfer contents to a new (prewetted) plate and continue.
	Plate set directly on table or absorbent towels during incubations or reagent additions	Set plate on plate stand or raised edge so bottom of filter is not touching any surface.
	Insufficient blotting of filter plate bottom causing wicking	Blot the bottom of the filter plate well with absorbent towels after each wash step.
	Pipette touching plate filter during additions	Pipette to the side of well.
	Probe height not adjusted correctly	Adjust probe to 3 alignment discs in well H6.
Background is too high	Background wells were contaminated	Avoid cross-well contamination by using sealer appropriately and by pipeting with multichannel pipets without touching reagent in plate.
	Matrix used has endogenous analyte or interference	Check matrix ingredients for crossreacting components (e.g. interleukin modified tissue culture medium).
	Insufficient washes	Increase number of washes.

Beads not in region or gate	Luminex not calibrated correctly or recently	Calibrate Luminex based on instrument manufacturer's instructions at least once a week or if temperature has changed by >3°C.
	Gate settings not adjusted correctly	Some Luminex instruments (e.g. Bio-Plex) require different gate settings than those described in the kit protocol. Use instrument default settings.
	Wrong bead regions in protocol template	Check kit protocol for correct bead regions or analyte selection.
	Incorrect sample type used	Samples containing organic solvents or if highly viscous should be diluted or dialyzed as required.
	Instrument not washed or primed	Prime the Luminex 4 times to eliminate air bubbles. Wash 4 times with sheath fluid or water if there is any remnant alcohol or sanitizing liquid.
	Beads were exposed to light	Keep plate and bead mix covered with dark lid or aluminum foil during all incubation steps.
Signal for whole plate is same as background	Incorrect or no Detection Antibody was added	Add appropriate Detection Antibody and continue.
-	Streptavidin-Phycoerythrin was not added	Add Streptavidin-Phycoerythrin according to protocol. If Detection Antibody has already been vacuumed out, sensitivity may be low.
Low signal for standard curve	Detection Antibody may have been vacuumed out prior to adding Streptavidin Phycoerythrin	May need to repeat assay if desired sensitivity not achieved.
	Incubations done at incorrect temperatures, timings or agitation	Assay conditions need to be checked.
Signals too high, standard curves are saturated	Calibration target value set too high	With some Luminex instruments (e.g. Bio- Plex) default target setting for RP1 calibrator is set at High PMT. Use low target value for calibration and reanalyze plate.
	Plate incubation was too long with standard curve and samples	Use shorter incubation time.
Sample readings are out of range	Samples contain no or below detectable levels of analyte	If below detectable levels, it may be possible to use higher sample volume. Check with tech support for appropriate protocol modifications.
	Samples contain analyte concentrations higher than highest standard point	Samples may require dilution and reanalysis for that particular analyte.
	Standard curve was saturated at higher end of curve	See above.
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High variation in samples and/or standards	Multichannel pipet may not be calibrated	Calibrate pipets.
- Cta. 1541 55	Plate washing was not uniform	Confirm all reagents are vacuumed out completely in all wash steps.
	Samples may have high particulate matter or other interfering substances	See above.
	Plate agitation was insufficient	Plate should be agitated during all incubation steps using a vertical plate shaker at a speed where beads are in constant motion without splashing.
	Cross-well contamination	Check when reusing plate sealer that no reagent has touched sealer.
		Care should be taken when using same pipet tips that are used for reagent additions and that pipet tip does not touch reagent in plate.

REPLACEMENT REAGENTS	Catalog #
Human CVD Panel 2 Standard Human CVD Panel 2 Quality Controls	HCVD2-8067-2 HCVD2-6067-2
Human CVD Panel 2 Detection Antibodies Bead Diluent Streptavidin-Phycoerythrin Assay Buffer Set of two 96-Well Filter Plates with Sealers 10X Wash Buffer Antibody-Immobilized Bead	HCVD2-1067-2 LBD L-SAPE6 L-AB MX-PLATE L-WB

<u>Analyte</u>	Bead #	Cat. #
Human Haptoglobin	51	HHPTGN

#### ORDERING INFORMATION

## To place an order:

To assure the clarity of your custom Human CVD Single Plex Haptoglobin kit order, please FAX the following information to our customer service department:

- Your name, telephone and/or fax number
- Customer account number
- Shipping and billing address
- Purchase order number
- Catalog number and description of product
- · Quantity of kits

FAX: (636) 441-8050 Toll-Free US: (866) 441-8400

(636) 441-8400

Mail Orders: Millipore Corp.

6 Research Park Drive

St. Charles, Missouri 63304 U.S.A.

#### For International Customers:

To best serve our international customers in placing an order or obtaining additional information about MILLIPLEX<sup>TM</sup> MAP products, please contact your multiplex specialist or sales representative or email our European Customer Service at <a href="mailto:customerserviceEU@Millipore.com">customerserviceEU@Millipore.com</a>.

#### **Conditions of Sale**

All products are for research use only. They are not intended for use in clinical diagnosis or for administration to humans or animals. All products are intended for *in vitro* use only.

#### **Material Safety Data Sheets (MSDS)**

Material Safety Data Sheets for Millipore products may be ordered by fax or phone or through our website at www.millipore.com/techlibrary/index.do.

# **WELL MAP**

	1	2	3	4	5	6	7	8	9	10	11	12
Α	0 Standard (Background)	10 ng/mL Standard	QC-2 Control									
В	0 Standard (Background)	10 ng/mL Standard	QC-2 Control									
С	0.08 ng/mL Standard	50 ng/mL Standard	Sample 1									
D	0.08 ng/mL Standard	50 ng/mL Standard	Sample 1									
E	0.4 ng/mL Standard	250 ng/mL Standard	Sample 2									
F	0.4 ng/mL Standard	250 ng/mL Standard	Sample 2									
G	2 ng/mL Standard	QC-1 Control	Etc.									
Н	2 ng/mL Standard	QC-1 Control										