

Bio-Plex Pro Mouse Cytokine Assays

Quick Guide

For Use with	Instruction Manual #
Bio-Plex Pro Mouse Cytokine, Chemokine, and Growth Factor Assays	10000142118

This guide can be used to prepare and run a full 1 x 96-well assay plate. Refer to the complete instruction manual for more information on a given step. New users can go to **bio-rad.com/bio-plex** and download the manual, which includes detailed instructions and a list of kit components.

IMPORTANT! Pay close attention to **vortexing**, **shaking**, and **incubation** instructions. Deviation from the protocol may result in low assay signal and assay variability.

Initial Preparation

- 1. Plan the plate layout.
- Start up/warm up the Bio-Plex Multiplex Immunoassay System (30 min). Bring assay buffer, wash buffer, and sample diluent to room temperature (RT) and keep other items on ice until needed. Begin to thaw frozen samples.
- 3. After thawing samples, prepare them according to the following guidelines.

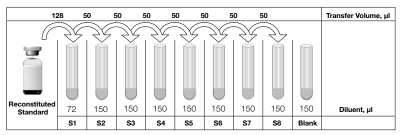
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Assay	Dilution	Diluent	Dilution	Diluent	Dilution	Diluent
Mouse cytokine	1:4	Bio-Plex sample diluent	User optimized	Cell culture medium or buffer similar to sample*	User optimized (1:2 lysates at 200–900 µg/ml protein)	Bio-Plex sample diluent

* If samples are serum-free, add bovine serum albumin (BSA) to 0.5% final w/v.

- Prime the wash station for a flat bottom plate. Prepare 1x wash buffer. Mix 10x stock by inversion to ensure all salts are in solution. Then dilute 1 part 10x wash buffer (60 ml) with 9 parts distilled water (540 ml).
- 5. Calibrate the Bio-Plex System by following the prompts in Bio-Plex Manager Software. This can be done now or during an assay incubation step.
- Reconstitute a single vial of standards in 500 μl of a diluent similar to the final sample type or matrix. Vortex for 5 sec and incubate on ice for 30 min.

Sample Type	Diluent for Standards	Add BSA
Serum and plasma	Standard diluent	None
Culture media with serum	Culture media	None
Culture media, serum-free	Culture media	To 0.5% final (w/v)

 Prepare a fourfold standard dilution series and blank as shown. Vortex for 5 sec between liquid transfers.



Note: Change tips between each dilution.

8. Vortex the 10x coupled beads for **30 sec** and dilute to 1x in Bio-Plex Assay Buffer as shown. Protect from light.

Mouse Cytokine Assays

Number of Wells	10x Beads, µl	Assay Buffer, µl	Total Volume, µl
96	575	5,175	5,750

Running the Assay

- Vortex the diluted (1x) beads for 10–20 seconds. Add 50 μl to each well of the assay plate.
- 2. Wash the plate two times with 100 µl Bio-Plex Wash Buffer.
- 3. Vortex samples, standards, and blank. Add 50 µl to each well.
- Cover the plate with sealing tape and incubate on shaker at 850 ± 50 rpm at RT. See the following table for incubation times for sample, detection antibody, and streptavidin-phycoerythrin (SA-PE).

I	Incubation Time	
Detection		
Sample	Antibody	SA-PE
30 min	30 min	10 min
	Sample	Detection Sample Antibody

 With 10 min left in the incubation, vortex the 10x detection antibody for 5 sec and quick-spin to collect liquid. Dilute to 1x as shown.

Number of Wells	10x Detection Antibody, µl	Detection Antibody Diluent HP, µl	Total Volume, µl
96	300	2,700	3,000

- 6. Wash the plate three times with 100 µl wash buffer.
- 7. Vortex the diluted (1x) detection antibody. Add $25 \ \mu l$ to each well.
- 8. Repeat step 4. See table for incubation time. Meanwhile, prepare the Bio-Plex Manager Software protocol; enter standard S1 values provided in the assay kit.
- **9.** With 10 min left in the incubation, **vortex** the 100x SA-PE for **5 sec** and quick-spin to collect liquid. Dilute to 1x as shown and protect from light.

Number of Wells	100x SA-PE, μΙ	Assay Buffer, µl	Total Volume, µl
96	60	5,940	6,000

- 10. Wash the plate three times with 100 μI wash buffer.
- 11. Vortex the diluted (1x) SA-PE. Add 50 µl to each well.
- **12.** Repeat step 4. See table for incubation time.

13. Wash the plate three times with 100 μI wash buffer.

- 14. Resuspend the beads in $125 \ \mu$ I assay buffer. Cover the plate as in step 4 and shake it at $850 \pm 50 \ rpm$ for $30 \ sec$.
- 15. Remove the sealing tape and read plate using the following settings for optimal sensitivity. Use low photomultiplier tube (PMT) RP1 setting for the Bio-Plex 100, 200, 3D, or similar system. Use default instrument settings for the Bio-Plex MAGPIX Multiplex Reader.

Assay	Low PMT, RP1	High PMT, RP1
Bio-Plex Pro Mouse Cytokine	•	-

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