

May 15, 2019

Working

UC Davis - Lipoprotein Binding Protein (LBP)-Endotoxemia Assay 👄

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dx.doi.org/10.17504/protocols.io.yrhfv36

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ABSTRACT

Summary:

Plasma samples will be assayed for lipopolysaccharide binding protein (LBP) as surrogate for bacterial LPS/measure of endotoxemia via ELISA.

EXTERNAL LINK

https://mmpc.org/shared/document.aspx?id=299&docType=Protocol

MATERIALS

NAME V	CATALOG #	VENDOR V
Mouse LBP ELISA kit	#43	Biometec
Shaker		
Microplate spectrophotometer		
MB grade water (diluent)		

MATERIALS TEXT

Reagent Preparation: (According to the manufacturer.)

Wash Buffer (PBS/ Tween 0.05%):

Dissolve 1 Tablet Phosphate buffered saline (PBS, vial 5) in 200ml distilled water -add 100 µl Tween 20 (vial 7). (Prepared wash buffer is stable for 4 weeks at refrigerator).

Phosphate Buffered Saline (PBS):

Dilute 1 Tablet of vial 5 in 200 ml distilled water

Dilution Buffer:

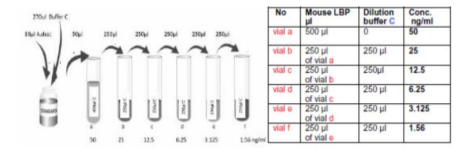
Add content of the vial 6 to 50ml PBS (Buffer C). Prepare just before use. Store remaining dilution buffer after reconstitution at -20°C

Reference serum dilution:

Add 10 µl distilled water to the vial 4. This contains 12.14 ± 3.5µg/ml LBP (! new Reference for Lot #141016). For assay dilute 1:800 (10µl serum +7990µl dilution buffer and use 100µl/well.

LBP standards:

Firstly, pipette 30 µl distilled water to the vial 3 for reconstitution and secondly add 270µl dilution buffer (C) to this vial and mix carefully, thirdly pipette 50µl from this vial to a new vial containing 450µl dilution buffer (C) and mix carefully. Finally this last vial contains 500µl standard dilution and containing 50ng/ml LBP = vial a. For standard curve prepare vial b-f and use vial a -f Prepare just before use. Store the standard at -20°C.



BEFORE STARTING

IMPORTANT: Check kit datasheet for lot-specific instructions that may modify general protocol.		
1	Prepare kit reagents.	
2	Dilute mouse plasma or serum samples 1:800.	
3	Add 100 μ l of standards (50, 25, 12.5, 6.25, 3.12, 1.56 ng/ml) or diluted samples in duplicate into the corresponding wells of the precoated modules and incubate for one hour at room temperature and shaking (300rpm).	
4	Wash 3X with Wash Buffer.	
5	Add 100 μ l detecting antibody to each well and incubate at room temperature for 1 hour at shaker.	
6	Wash 3X with Wash Buffer.	
7	Add 100 µl Substrate solution to each well. Incubate 12-14 min in the dark at room temperature without shaking. Cover with foil during incubation or place in drawer.	
8	Add 100 µl stopping solution to each well. Tap gently to mix.	
9	Read absorbance at 450 nm (reference wave length 620 nm)	
10	Calculate the LBP concentration by first plotting the OD means of standards (y-axis) and the LBP concentration (x-axis). Calculate the LBF concentration from the mean OD of samples from the standard curve and multiply with dilution factor.	

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