Core D Procedure Protocol ATP, Kdo2 Lipid A Treatment 5-12-2009

Overview

- 1. One large experiment will be completed to provide samples to all cores.
- Untreated cells will be harvested at 0 hours. Treated cells will be harvested at 15 m, 30 m, 1 hr, 2 hr, 4 hr, 8 hr and 20 hr. See Figure 1 for an overview of the treatment time course.
- 3. Table 1 for an overview of the treatment.
- 4. Only one replicate will be made for each time and treatment. 10 aliquots will be generated from each replicate. Cores E, G, and J will receive one aliquot, cores H, I and K will receive two aliquots and core G will receive 20 mL of medium from each treatment.
- 5. Data will be normalized to DNA content (pmol lipid per μ g DNA). DNA and TNFa data will be generated by core D. DNA and TNFa assay data will be emailed to each core.

Setup

- Seed 2 100 mm TC plates with 1 x 10⁷ cells in 10 mL Bone Marrow-derived Macrophage Growth Medium 2 (PS000003201) each.
- 2. Incubate 16 hours at 37°C.

Reagent Preparation

- 1. ATP (PS000004000)
- 2. Kdo2 Lipid A working solution prepare from Kdo2 Lipid A stock solution (PS0000001401)

Treatment

1. Treat each plate with the appropriate reagents:

Pretreatment (priming)	10 μL 100 ug/mL Kdo2 (FC-100
4h Kdo2 or DPBS followed	ng/mL)
by	10 μL DPBS
ATP or 10 mM Hepes/10 mM DTT	250 μL 200 mM/mL ATP (FC-5
time course: 15 m, 30 m, 1 hr, 2 hr,	mM/mL)
4 hr, 8 hr and 20 hr	250 μL 10 mM Hepes/10 mM DTT

- 2. Note the time and return the plates to the incubator.
- 3. For 0 hours, do not treat with anything. Proceed with harvest.

Harvest

- 1. At the appropriate time, remove the plates from the incubator.
- 2. Immediately remove medium from each plate and place in labeled 50 mL polypropylene tubes for Core G.

- 3. Remove .5 mL of medium from each treatment and place in labeled Eppendorf tubes for the TNF α assay.
- 4. Place the plates on ice and wash twice with 5 mL of cold DPBS.
- 5. Add 5 mL of cold DPBS to the first plate and 5.5 mL to the second plate of each treatment and scrape the cells with a scraper.
- 6. Transfer the cells to a labeled 50 mL conical tube on ice.
- 7. Gently suspend cells by swirling or lightly vortexing for 10 secs.
- 8. Remove 200 µL of the cell suspension to an Eppendorf on ice for the DNA assay.
- 9. Divide the 10 mL cells into aliquots of $1 \text{ mL/2 x } 10^6 \text{ cells each}$, placing into 13x100mm glass tubes with Teflon-lined screw caps.
- 10. Freeze and store cells at -80°C.

Medium Processing for TNFa Assay

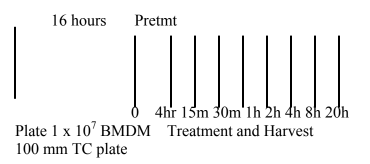
- 1. Centrifuge the TNF α aliquots at top speed in refrigerated microfuge for 2 min, collect an aliquot of supernatant (~0.4 mL) and place in a new labeled Eppendorf tube. Freeze the aliquots at -20°C.
- 2. TNFα is to be assayed by Jay Westcott at ELISA Tech (Aurora, CO).

Cell Processing for DNA Assay

- 1. Store at -20° C.
- 2. DNA is to be assayed by core D according to the LIPID MAPS DNA Assay (PP0000002700).

All samples will be shipped to their cores on dry ice

Figure 1



Sample	Pretmt	Tmt	<u>Rx</u>
<u>#</u>	Hour	Hour	
1	0	0	-K-A
2	4	15 m	-K-A
3	4	15 m	+K-A
4	4	15 m	-K+A
5	4	15 m	+K+A
6	4	30 m	-K-A
7	4	30 m	+K-A
8	4	30 m	-K+A
9	4	30 m	+K+A
10	4	1	-K-A
11	4	1	+K-A
12	4	1	-K+A
13	4	1	+K+A
14	4	2	-K-A
15	4	2	+K-A
16	4	2	-K+A
17	4	2	+K+A
18	4	4	-K-A
19	4	4	+K-A
20	4	4	-K+A
21	4	4	+K+A
22	4	8	-K-A
23	4	8	+K-A
24	4	8	-K+A
25	4	8	+K+A
26	4	20	-K-A
27	4	20	+K-A
28	4	20	-K+A
29	4	20	+K+A

 Table 1: Number labeling of samples, time and treatment

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