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Cell Surface Mild Acid Elution of MHC-bound Immunopeptides

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Culturing Cells

1 Culture cells. If culturing THP-1s, follow the protocol.



- 1.1 The base medium for this cell line is RMPI-1640
- 1.2 Reguired supplements:
 - [M]1 % volume L-glutamine
 - [M]10 % volume Fetal Bovine Serum
 - Most catalog numbers of RMPI-1640 contain L-glutamine, however, some do not. Ensure that it is in the media before using it for culturing.
- 1.3 Optional Supplements:
 - [M]1 % volume PenStrep
 - [M]0.05 Milimolar (mM) 2-mercaptoethanol
 - PenStrep is not required for THP-1 culturing, however, if you are having issues with bacterial contamination, it can be used at 1X.
 - 2-mercaptoethanol is stated as a required component for complete RPMI-1640 medium, however, in our laboratory it is not standard practice to add it.

4	Always store cells in liquid nitrogen. This is for both the original tube of cells from ATCC and any passages afterwards.
.5	Place the media bottle in the § 37 °C water bath at least © 00:30:00 prior to using
.6	Thaw cells at § 25 °C (room temperature) for © 00:10:00 or § 37 °C in a water bath for © 00:02:00
.7	Sanitize all items going into the Biological Safety Cabinet with 70% ethanol
.8	As soon as the cells are thawed, transfer the cells to a 15 ml conical tube and add 10 ml of complete media
	Cells are stored with 5% DMSO, which can lyse cells if they are left for too long.
.9	Pellet cells for © 00:03:00 at 500g
0	Discard supernatant
1	Resuspend cells by pipetting up and down 5X in _5 ml complete media
2	Transfer cells + media to a T-25 flask
3	Incubate cells at 8 37 °C and 5% CO2 and 80% humidity
4	THP-1 cells replicate after ~26 hours. In practice, it takes 2 days for a true doubling.
5	Once cells have doubled OR when media has begun to change colour, it is time to add media, split cells into new flasks, or to spin down to remove all media
	Adding media
6	If concerned about cell concentration, perform a cell count

- 1.17 Double the total media volume with new complete media
- 1.18 Carefully mix the new media in by rocking the flask back and forth
- 1.19 Place the flask back in the incubator
 - 2 Grow cells to a total number of 1-2x10*8 cells

Preparing Reagents

- 3 1X PBS at § 25 °C (room temperature)
 - **38 g NaCl**
 - **0.2** g KCl
 - 1.44 g NA2HPO4
 - **0.24** g KH2PO4
 - 1 L Milli-Q water
 - pH adjust to 7.4 with 37% HCl
- 4 1X PBS at 8 4 °C

This PBS solution does NOT have to be sterile, but make sure it is clearly marked.

- 5 1X saline (PBS without the phosphate) at 8 4 °C
 - ■4 g NaCl
 - **0.1** g KCl
 - **500 ml Milli-Q water**
 - no need to pH adjust, it's not a buffer
 - no need to sterilize as the mild acid elution is not done under sterile conditions
- 6 1X saline + 2% acetic acid at 4 °C
 - 245 ml of above 1X saline solution
 - **5** ml acetic acid
- 7 Each 250 ml T-175 flask will require 7 50 ml conical tubes

Pellet	Pelleting & Rinsing Cells	
8	Cells will most likely be in 250 ml complete media. Use five 50 ml conical tubes to pellet all cells from one flask at a time.	
9	Pellet cells for \bigcirc 00:03:00 at 500g	
10	Discard supernatant	
11	Resuspend each pellet by pipetting up and down 5X with 35 ml room temp PBS (sterile in the hood)	
12	Combine all five conical tubes into one	
13	If cell count is not know, perform one now to confrim a minimum of 1x10*8 cells	
14	Pellet cells for © 00:03:00 at 500g	
15	Discard supernatant	
16	Resuspend each pellet by pipetting up and down 5X with 10 ml cold PBS	
17	Pellet cells for \bigcirc 00:03:00 at 500g	
18	Discard supernatant	
19	Resuspend each pellet by pipetting up and down 5X with 10 ml cold PBS	
20	Pellet cells for $© 00:03:00$ at $500g$	
21	Discard supernatant	
22	Resuspend each pellet by pipetting up and down 5X with 10 ml cold saline	

23	Transfer cells + saline to a new 50 ml conical tube		
	This is to remove any large amounts of phosphate		
24	Pellet cells for ③ 00:03:00 at 500g		
25	Discard supernatant		
Mild Acid Elution			
26	Resuspend each pellet by pipetting up and down 5X with 10 ml cold 2% acetic acid in saline		
	EXTREMELY IMPORTANT, REMEMBER THAT THE PEPTIDES WILL BE IN THE SUPERNATANT		
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27	DO NOT DISCARD SUPERNATANT		
28	Collect the supernatant in a new 350 ml conical tube		
Drying			
29	Freeze supernatant at 8 -80 °C overnight		
30	Lyophilize sample for 48 hours		
31	Peptides are now ready to be desalted with chosen clean up method (STAGE tip, LC-UV, combination, etc)		
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