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Phagocytosis Bead Conjugation V.1

Andrew Crowley¹

¹Dartmouth College, Ackerman Lab

Working

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Ackerman Lab



Andrew Crowley

Dartmouth College, Ackerman Lab



ABSTRACT

Protocols for the amine coupling of microspheres

STEPS MATERIALS

NAME ▾	CATALOG # ▾	VENDOR ▾
PBS-TBN	P0210	Teknova
PBS-TBN	P0210	Teknova

Activation

- Calculate the beads / mL from the percent solids as follows:
$$(6W \times 10^{12}) / (\rho \times \pi \times \phi^3)$$

W = grams solid per mL
ρ = density (g/mL)
φ = particle diameter (μm)

polystyrene density = 1.055 /mL
- Resuspend beads by briefly vortexing; dispense **200 μl (contains approx. 7 billion beads)**
- Pellet by centrifugation **17000 x g** **00:03:00** and remove supernatant by pipette
- Resuspend in a total of **1 ml** consisting of:
 - sulfo-NHS **50 mg/mL** made in **10 Milimolar (mM)** MES (pH 5.0) **100 μl**
 - EDC **50 mg/mL** , made in **10 Milimolar (mM)** MES (pH 5.0) **100 μl**
 - **10 Milimolar (mM)** MES (pH 5.0) **800 μl**

5 Incubate with end-over-end mixing (shielded from light) ⌚ 00:20:00 🌡 Room temperature

Coupling

6 Wash by pelleting beads and replacing supernatant with [M]10 Milimolar (mM) MES (pH 5.0) 🧴 1 ml
🌀 17000 x g ⌚ 00:02:00

7 Formulate protein for coupling @ approx. [M]25 µg/mL in [M]10 Milimolar (mM) sodium acetate (pH 5.0) 🧴 7.5 ml



n.b.

The large volume used for the step is based on observations that sufficiently high bead concentrations during this step may lead to crosslinking between microspheres (**anecdotal**)

8 Incubate with end-over-end mixing (shielded from light) ⌚ 03:00:00 🌡 Room temperature

Blocking + Washes

9 Collect the full volume in the base of the reaction tube by centrifugation 🌀 250 x g ⌚ 00:01:00

10 Split the 🧴 7.5 ml volume into multiple fractions in microcentrifuge tubes (🧴 2 ml) in order to centrifuge at a sufficiently high speed
🌀 17000 x g ⌚ 00:03:00


11 Remove supernatant from each fraction by pipette

12 Resuspend **one** fraction in PBSF (ie. 1x PBS + [M]0.1 Mass/Volume Percent BSA) 🧴 1 ml

13 Transfer the entire contents of the first fraction to the second and resuspend; continue through the fractions until as many beads as possible have been collected in a single microcentrifuge tube

14 Pellet beads via centrifuge  **17000 x g**  **00:02:00**

15  **go to step #11** for a total of 3 washes

16 Resuspend in PBSF  **500 µl**

17 Will the beads be used for an assay on the same day as production?

step case

No

no description provided

18 Block via  **16:00:00 (ie. overnight)**  **4 °C**

Counting + Qualification

19 Make serial dilutions of 1:100, 1:1000, and 1:10,000


20 Measure concentration by flow cytometry using the fluorescent dye of the microsphere as the trigger condition




Typical yield is approx. $1 - 2 \times 10^8$ beads, or around 25% of the original amount



MACS Quant
flow cytometer
Miltenyi 130-096-343

21 In a non-binding, 96-well plate, formulate serial dilutions of positive and negative control antibodies beginning at  **5 µg/mL** and diluted by 1:3 over an approx. 4 point series

22 Add 100,000 beads to each well

- 23 Incubate with shaking (shielded from light) ⌚ 01:00:00 🌡 Room temperature
- 24 Pellet by centrifugation 🌀 3500 x g ⌚ 00:05:00 and remove supernatant by decanting
- 25 Resuspend beads in PBST (ie. 1x PBS + [M] 0.05 Volume Percent Tween20) 📄 200 µl
- 26 [go to step #24](#) for a total of 2 washes
- 27 Resuspend beads in fluorescent secondary 📄 100 µl [M] 0.65 µg/mL
-  n.b.

Make sure that the fluorescent molecule chosen for the secondary does not have spectral overlap with the dye in the microspheres themselves
- 28 Incubate with shaking (shielded from light) ⌚ 00:30:00 🌡 Room temperature
- 29 Pellet by centrifugation 🌀 3500 x g ⌚ 00:05:00 and remove supernatant by decanting
- 30 Resuspend beads in PBST (ie. 1x PBS + [M] 0.05 Volume Percent Tween20) 📄 200 µl
- 31 Pellet by centrifugation 🌀 3500 x g ⌚ 00:05:00 and remove supernatant by decanting
- 32 Resuspend beads in 1x PBS 📄 100 µl
- 33 Measure fluorescent intensity by flow cytometry



MACS Quant
flow cytometer
Miltenyi 130-096-343

Washing + Blocking

step case

No

no description provided



NaN Block via 🕒 16:00:00 (ie. overnight) 🌡 4 °C

Counting + Qualification

NaN

Washing + Blocking

step case

Yes

no description provided



NaN Block via 🕒 00:30:00 🌡 Room temperature

Blocking + Washes

step case

Yes

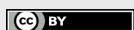
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18 Block via 🕒 00:30:00 🌡 Room temperature

Counting + Qualification

19 Make serial dilutions of 1:100, 1:1000, and 1:10,000



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