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CD34+ isolation from human bone marrow V.2

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Human Islet Research Ne...



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Abstract

This protocol details the steps for isolating CD34+ cells from human bone marrow. The CD34+ cells isolated from this protocol can be used for generating humanized mice through reconstitution of immune cells via IV injection after bone marrow ablation. These cells can also be used for mixed lymphocyte reaction experiments.

Note

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Materials

Required material

- Bone Marrow Medium
- MACS buffer(degassed)
- Sterile flask for BM rinse
- 15 and 50 mL Falcons
- Histopaque
- Sterile pipetts
- CD34+ MACS kit (<u>130-046-702</u>)
- Human serum
- MACS supply
- antibodies

Required Buffers

- BM Medium (\bot 500 mL Media 199, \bot 5 mL Hepes, \bot 5 mL DNAse, \bot 40 μ L Gentamycin) ■ MACS buffer (△ 500 mL PBS, △ 5 g BSA, △ 2 mL EDTA, sterile filtrated and degassed)
- Cryomedium (☐ 90 mL PBS, ☐ 10 mL FBS, ☐ 10 mL DMSO)

Before start

Human bone marrow is a rich source for CD34+ hematopoietic stem cells. CD34+ cells can be easily isolated and further processed.



- 1 Transfer the content of the collection bag into a sterile flask
- 2 Add <u>A</u> 250 mL BM Medium to the bag and rinse it thoroughly
- 3 Transfer the content of the collection bag into the sterile flask
- 4 Layer 🚨 35 mL of the suspension over 🚨 15 mL of Histopaque
- 5 Centrifuge the tubes for 30 minutes 4 500 g without brake at RT
- 6 Collect the leukocyte ring in 4 50 mL Falcons and fill up with BM medium
- Wash once by centrifuging 6 minutes 500 g
- 8 Resuspend the cells in MACS buffer and count (take \perp 50 µL for FACS confirmation = PRE)
- Wash down again and resuspend the pellet according to the protocol (130-046-702 MACS Human CD34+ kit)
- 10 Add \perp 300 μ L of MACS buffer per 10^8 cells
- 11 Add \perp 100 μ L of FcR-B reagent per 10^8 cells, mix it and incubate in fridge for 15 minutes
- 12 Add $\underline{\bot}$ 100 μ L of CD34 beads per 10^8 cells to the suspension and mix and let it sit for 30 min (fridge)
- 13 Fill up with 4 50 mL MACS buffer and strain through a blue strainer (40 uM)



- 14 Wash cells (\$\rm 500 g 6 min) and resuspend in MACS buffer 500 uL/200,000,000 cells. If you have more cells, increase volume accordingly. E.g 3 *10^9 cells = 7,5mL. Aliquot this volume to more than one (with 4 3 mL prerinsed) MACS column.
- 15 Wash with buffer \bot 3 mL 3 times and keep negative fraction (*Take* \bot 50 μ L *for FCM* = POST neg)
- 16 Put the column out of the magnet and push out positive fraction with 4 5 mL Buffer and the plunger
- 17 Collect the positive fractions. (Take \perp 50 µL for FCM = Post pos)
- 18 Process the cells as desired (injection in mouse or cryopreservation)
- 19 Check the puritiy with FACS

20

FACS pane	<u>el</u>	
CD45	FITC	5 ul
CD3	PercpC5.5	5 ul
CD14	PacBlue	5 ul
CD19	APC	5 ul
CD34	PE	5 ul
CD38	PeC7	5 ul
Human Serum		5 ul
FACSbuffer		15 ul
Total		50 ul/sample