🐏 pluriSelect

Protocol for the use of PBMC Spin Medium pre-filled pluriMate® II Tubes

pluriMate® II - Specification

	pluriMate II - 15 ml, pre-filled	pluriMate II - 50 ml, pre-filled
Order No. 50 pcs.	44-19215-10	44-19250-10
Order No. 100 pcs.	44-19215-11	44-19250-11
Order No. 500 pcs.	44-19215-15	44-19250-15

Product Description The pluriMate[®] II centrifugation tubes pre-filled with PBMC Spin Medium[®] can be used for an optimal separation peripheral blood mononuclear cells (PBMC) from whole blood and bone marrow. The key feature of the pluriMate[®] II tubes is the mesh supported barrier. This barrier prevents you from time-consuming and laborious overlaying of the sample material. Anticoagulated blood or bone marrow can simply be poured directly from the blood sampling tube into the pluriMate[®] II tube. The porous barrier prevents mixture of the sample material with the separation medium. When separation is complete, the barrier prevents recontamination of the enriched cell fraction during harvest.

Pre-filled with	PBMC Spin Medium [®] (Catalog 60-00092-10)
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- **Enrichment of** Peripheral Blood Mononuclear Cells (PBMC)
- Age of blood < 8 hours

Directions for the use of the pluriMate® II Tube

1. Check that recommended medium, blood sample, density gradient medium and centrifuge are all at room temperature.

Preparation of the pluriMate® II Tube

2. Centrifuge at 1000 x g for 10 sec. and leave 3 - 5 mm supernatent on top.

Add Sample Material

 Fill in sample material on top of mesh (Fig. a).
Note: To reduce platelet contamination you can add pluriSpin[®] PLT Depletion (Order No. 19-00002-31)

	pluriMate® II 15 ml	pluriMate® II 50 ml
Sample material vol.	2 - 7 ml	5 - 30 ml

Spin

 Centrifuge for 15 minutes at 800 x g at room temperature with in a swing bukket rotor and the **brake on**. Using blood older than 4 hours centrifuge for 30 minutes at 1000g.

Collect

- 5. Remove plasma by pipetting until white cell layer (Fig. d).
- 6. Collect cells in the white layer in a fresh tube (Fig. e).

Wash

- 7. Fill up reaction tube with wash buffer.
- 8. Spin down cells 10 minutes with 300 x g (no or small brake) at 4°C.
- 9. Pour out supernatant, leave the reaction tube on the table for 20 sec. Wash buffer excess will run down from the tube wall and collect at the bottom.
- 10. Aspirate most of the liquid above the pellet. The liquid will look foggy, these are mostly platelets aspiration will improve washing result.
- 11. Reconstitute pellet with 1 ml of wash buffer by carefully pipetting.
- 12. Repeat steps 7 to 10.
- 13. Reconstitute pellet at your desired volume.

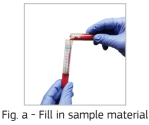




Fig. b - Before centrifugation



Fig. c - After centrifugation



Fig. d - Remove plasma



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