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PBMC- 02 - CD4+ T cell Isolation from PBMC with "Dynabeads CD4 Positive Isolation Kit" V.2

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1 Works for me

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ABSTRACT

List of published works using this protocol:

- Kustrimovic N., Comi C., Magistrelli L., Rasini E., Legnaro M., Bombelli R., Aleksic I., Blandini F., Minafra B., Riboldazzi G., Struchio A., Mauri M., Bono G., Marino F., Cosentino M. Parkinson's disease patients have a complex phenotypic and functional Th1 bias: cross-sectional studies of CD4+ Th1/Th2/T17 and Treg in drug-naïve and drug-treated patients (2018). Journal of neuroinflammation, 15(1), 205. https://doi.org/10.1186/s12974-018-1248-8
- Kustrimovic, N., Rasini, E., Legnaro, M., Bombelli, R., Aleksic, I., Blandini, F., Comi, C., Mauri, M., Minafra, B., Riboldazzi, G., Sanchez-Guajardo, V., Marino, F., & Cosentino, M. (2016). Dopaminergic Receptors on CD4+ T Naive and Memory Lymphocytes Correlate with Motor Impairment in Patients with Parkinson's Disease. Scientific reports, 6, 33738. https://doi.org/10.1038/srep33738
- Cosentino M., Ferrari M., Kustrimovic N., Rasini E., Marino F. (2015). Influence of dopamine receptor gene polymorphisms on circulating T lymphocytes: A pilot study in healthy subjects. Human immunology, 76, 10, 747-752. https://doi.org/10.1016/j.humimm.2015.09.032

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MATERIALS TEXT

MATERIALS

Fisher Catalog #11331D

(FBS) EuroClone Catalog #ECS0180L-500 ml

⊠ RPMI

1640 EuroClone Catalog #ECM 0495L- 500 ml

⊠BSA **Sigma**

Aldrich Catalog #A2153

⊠ BD tubes **Becton**-

Dickinson Catalog #352054

Instrumentation required:

a.Magnet (DynaMag[™]) b.Sample Mixer with rotation c.Laminar flow hood

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BEFORE STARTING

If you need to obtain CD4+ T cell for subsequent cell culture, make sure you are using sterile buffers and sterile plastic disposables as well. Moreover, work under laminar flow hood when you are processing samples (from the beginning to the end of the following procedure). Otherwise, use non-sterile Buffers and disposables, and process samples in a cell isolation laboratory.

IMPORTANT NOTE: the isolation protocol is calibrated for using $25\mu L$ of beads for $10x10^6$ PBMCs resuspended in 1mL. For lower or higher cell number than 10x106, resize the volumes, accordingly. (See also Table 1on the data sheet of the kit).

ALL REAGENTS MUST BE AT ROOM TEMPERATURE WHEN USED!!!

1	Isolate PBMCs according either to the standard protocol from fresh blood or from buffy coat (PBMC- 01a - Isolation of Human PBMC from Buffy Coat, PBMC- 01b - Isolation of Human PBMC from Whole Blood).
2	Count the cells with Cellometer machine or by manual count, using either Trypan Blue or Türk solutions accordingly.
	For automatic cell count with Cellometer machine use Trypan Blue. Follow protocol CELL COUNT- 03
	SOLUTION- 09 - Trypan Blue solution by Farmacologia Medica
	For manual cell count use Türk solution for checking purity. Follow protocol CELL COUNT- 02
	SOLUTION- 08 - Türk solution by Farmacologia Medica
	Cellometer Auto T4 Automated cell counter
	Nexcelom Bioscience EuroClone
3	Resuspend Dynabeads in the vial using a vortex for >30 sec.
4	Transfer the desired volume of Dynabeads to a 5mL-tube (use BD tubes cat. n. 352054) following this proportion: $25\mu L$ of beads for $10x10^6$ cells.
5	Add 22 µl of Solution-11 (found in the kit materials as Buffer 1), resuspend and place the tube into the magnet: beads will attach to the magnet very quickly (few seconds).
	Discard then the supernatant by using a glass Pasteur pipette.

Remove the tube from the magnet.

SOLUTION- 11 - CD4+T cells isolation buffer by Farmacologia Medica

- 6 Repeat the washing step 2 or 3 times to make sure that DMSO is all washed up.
- 7 After counting, centrifuge PBMCs sample at **\$\pi 1200 x g, 00:05:00**.

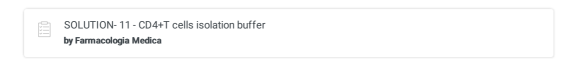
Allegra AVANTI 30 Centrifuge

Beckman Coulter Beckman Italy

- 8 Discard supernatant and resuspend pellet of $10x10^6$ cells in $\Box 1$ mL of **SOLUTION-11**.
 - SOLUTION- 11 CD4+T cells isolation buffer by Farmacologia Medica
- Q Transfer cell suspension into the tube with beads, and resuspend vigorously.
- 10 Incubate the beads with cells for **© 00:20:00** at **§ 4 °C** with gentle rotation by putting the Sample Mixer in the fridge.
- 11 After incubation place the tube on the magnet and wait for 1-2 minutes, that is until the complex beads-cells is completely attached to the magnet.
- 12 🛕

While the tube is still in the magnet, carefully remove and discard the supernatant with a glass Pasteur pipette.

13 Remove the tube from the magnet, add **2** μ**I** of **SOLUTION-11** and resuspend the cells very vigorously because of aggregates.

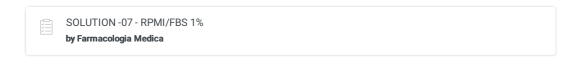


- Repeat steps 11-13 twice (in total 3 times) to wash the bead-bound CD4+ T cells. These steps are critical to obtain a high purity of isolated cells.
- 15 Resuspend cell pellet in 100 μl of **SOLUTION- 07** (found in the kit materials as Buffer 2)

 [The volume is calibrated for 10x10⁶ cells, for lower or higher number of cell resize the volume accordingly].

	SOLUTION -07 - RPMI/FBS 1% by Farmacologia Medica	
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- 16 Add 10 μl of DETACHaBEAD® CD4 for each 10x10⁶ PBMCs. (Resize this volume if the number of starting cell is different)
- 17 Add another $\Box 500 \ \mu I$ of SOLUTION- 07 to increase the volume and transfer everything in a 1.5 mL eppendorf.



- 18 Incubate © 00:45:00 at & Room temperature (RT) with gentle rotation by using a Sample Mixer.
- 19 Transfer the sample from eppendorf to BD tube, and place the tube on magnetand wait for 1-2 mins, that is until the complex beads-cells is completely attached to the magnet.
- 20 /

While the tube is still in the magnet, **transfer the supernatant** containing the released cells into a 15 mL conical tube.

To obtain residual cells, wash the beads 3 times with $\Box 500~\mu I$ of **SOLUTION- 07** and collect the supernatant each time.

	SOLUTION -07 - RPMI/FBS 1% by Farmacologia Medica	
21	Add to the detached cell suspension SOLUTION- 07 to a final volume of □5 mL and centrifuge at ③1200 x g, Room temperature, 00:05:00	
	SOLUTION -07 - RPMI/FBS 1% by Farmacologia Medica	
	Allegra AVANTI 30 Centrifuge Beckman Coulter Beckman Italy	
22	Resuspend the cells for cell counting in 1 mL of SOLUTION- 07: follow the appropriate protocol (see st protocol). Check the viability with Trypan blue.	ep 2 of this
	SOLUTION -07 - RPMI/FBS 1% by Farmacologia Medica	
	SOLUTION- 09 - Trypan Blue solution by Farmacologia Medica	

23 🙀

OPTIONAL STEP

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Check the purity of the isolated CD4+ T cells by flow cytometry.

If needed, check the purity by labeling CD4 with the appropriate CD markers, such as CD3, CD4, CD8 and CD14 Ab and analyze samples with a flow cytometer to exclude the presence of undesired subsets.

BD FACS Celesta Flow Cytometer

Becton Dickinson Milan Italy BD

24 EXPECTED RESULTS



Cell Viability: ≥95%

Cell Yield: $\pm 4.6 \times 10^6$ cells starting from 25 mL of Fresh Blood $\pm 6 \times 10^6$ cells starting from 25 mL of Buffy Coat

If checked, purity of the isolated CD4+ cells must be ≥95 %