EasySep™ Direct Human PBMC Isolation Kit

For processing 100 mL leukapheresis

Catalog #19654

Negative Selection

Document #10000012561 | Version 00



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Description

Isolate highly purified peripheral blood mononuclear cells (PBMCs) directly from leukapheresis samples by immunomagnetic negative selection. This kit can also be used to isolate PBMCs from other sample types (see Table 1).

- 99.9% RBC depletion without the need for density gradient centrifugation, sedimentation or lysis
- Fast, easy-to-use and column-free
- Isolated cells are untouched

This kit targets granulocytes, platelets, and red blood cells (RBCs) for removal with antibodies recognizing specific cell surface markers. Unwanted cells are labeled with antibodies and magnetic particles, and separated using an EasySep[™] magnet. PBMCs are simply collected into a new tube and are immediately available for downstream applications such as flow cytometry, culture, or DNA/RNA extraction.

• This is the Product Information Sheet (PIS) for isolating PBMCs from leukapheresis samples. If isolating PBMCs from other sample types, refer to the applicable PIS Document Number (see Table 1), available at www.stemcell.com or contact us to request a copy.

Table 1. Applicable Document Number for Other Sample Types

SAMPLE TYPE	PIS DOCUMENT NUMBER		
Whole blood	1000003347		
Buffy coat	10000012559		
Bone marrow	10000012544		
Cord blood	10000012560		
Leukoreduction system chamber (LRSC)	10000012562		

Component Descriptions

COMPONENT NAME	COMPONENT #	QUANTITY	STORAGE	SHELF LIFE	FORMAT
EasySep™ Direct Human PBMC Isolation Cocktail	19654C	2 x 2.5 mL	Store at 2 - 8°C. Do not freeze.	Stable until expiry date (EXP) on label.	A combination of monoclonal antibodies in PBS. Includes an Fc receptor blocking antibody.
EasySep™ Direct RapidSpheres™ 50300	50300	4 x 2.5 mL	Store at 2 - 8°C. Do not freeze.	Stable until expiry date (EXP) on label.	A suspension of magnetic particles and monoclonal antibodies in PBS.

PBS - phosphate-buffered saline

Components may be shipped at room temperature (15 - 25°C) but should be stored as indicated above.

Sample Preparation

For available fresh samples, see www.stemcell.com/primarycells.

LEUKAPHERESIS

For best recovery, use an unprocessed leukapheresis pack. Recovery of the desired isolated cells decreases with samples that are older than 24 hours. To avoid loss of monocytes, EDTA must be added to the leukapheresis sample to a final concentration of 6 mM prior to labeling and separation (see step 2, Tables 2 - 4). An EDTA stock solution greater than 0.05 M is recommended to avoid overdiluting the start sample.

Recommended Medium

EasySep[™] Buffer (Catalog #20144), RoboSep[™] Buffer (Catalog #20104), D-PBS (Without Ca++ and Mg++; Catalog #37350), or PBS containing 2% FBS and 1 mM EDTA. Medium should be free of Ca++ and Mg++.



Directions for Use – Manual EasySep™ Protocols

See page 1 for Sample Preparation and Recommended Medium. Refer to Tables 2 and 3 for detailed instructions regarding the EasySep[™] procedure for each magnet.

Table 2. EasySep™ Direct Human PBMC Isolation Kit Protocol for LEUKAPHERESIS

		EASYSEP™ MAGNETS				
STEP	INSTRUCTIONS	EasySep™ (Catalog #18000)	"The Big Easy" (Catalog #18001)			
	Prepare sample within the volume range.	0.5 - 1.5 mL	1 - 6 mL			
1	Add sample to required tube.	5 mL (12 x 75 mm) polystyrene round-bottom tube (e.g. Catalog #38007)	14 mL (17 x 95 mm) polystyrene round-bottom tube (e.g. Catalog #38008)			
2	Add EDTA to sample.	At a final concentration of 6 mM EDTA	At a final concentration of 6 mM EDTA			
3	Add Isolation Cocktail to sample. NOTE: Do not vortex cocktail.	50 μL/mL of sample	50 μL/mL of sample			
	Mix and incubate.	RT for 5 minutes	RT for 5 minutes			
4	Add recommended medium to top up the sample to the indicated volume. Mix by gently pipetting up and down 2 - 3 times.	Top up to double the original sample volume	Top up to double the original sample volume			
5	Vortex RapidSpheres™. NOTE: Particles should appear evenly dispersed.	30 seconds	30 seconds			
6	Add RapidSpheres™ to sample and mix.	50 µL/mL of original sample volume NOTE: No incubation, IMMEDIATELY proceed to next step	50 µL/mL of original sample volume NOTE: No incubation, IMMEDIATELY proceed to next step			
7	Place the tube (without lid) into the magnet and incubate.	RT for 5 minutes	RT for 5 minutes			
8	Pick up the magnet, and in one continuous motion invert the magnet and tube,* pouring the enriched cell suspension into a new tube.	Use a new 5 mL tube	Use a new 14 mL tube			
9	Add RapidSpheres™ to the new tube containing the enriched cells and mix.	Use same volume as in step 6 NOTE: No incubation, IMMEDIATELY proceed to next step	Use same volume as in step 6 NOTE: No incubation, IMMEDIATELY proceed to next step			
10	Remove the tube from the magnet; Place the tube from step 9 (without lid) into the magnet and incubate for a second separation.	RT for 5 minutes RT for 5 minutes				
11	Pick up the magnet, and in one continuous motion invert the magnet and tube,* pouring the enriched cell suspension into a new tube.	Isolated cells are ready for use	Isolated cells are ready for use			

RT - room temperature (15 - 25°C)

* Leave the magnet and tube inverted for 2 - 3 seconds, then return upright. Do not shake or blot off any drops that may remain hanging from the mouth of the tube.



Table 3. EasySep[™] Direct Human PBMC Isolation Kit Protocol for LEUKAPHERESIS

		EASYSEP™ MAGNETS				
		EasyEights™ ((Catalog #18103)		Easy 50
STEP	INSTRUCTIONS		5 mL tube	14 mL tube		Easy 50 (Catalog #18002)
	Prepare sample within the volume range.	0.5 - 2 mL		1 - 6 mL		5 - 25 mL
1	Add sample to required tube.	5 mL (12 x 75 mm) polystyrene round-bottom tube (e.g. Catalog #38007)		14 mL (17 x 95 mm) polystyrene round-bottom tube (e.g. Catalog #38008)		50 mL (30 x 115 mm) conical tube (e.g. Catalog #38010)
2	Add EDTA to sample.	At a final	concentration of 6 mM EDTA	At a final concentration of 6 mM EDTA		At a final concentration of 6 mM EDTA
3	Add Isolation Cocktail to sample. NOTE: Do not vortex cocktail.	50 µL/mL of sample		50 µL/mL of sample		50 µL/mL of sample
-	Mix and incubate.		RT for 5 minutes	RT for 5 minutes		RT for 5 minutes
4	Add recommended medium to top up the sample to the indicated volume. Mix by gently pipetting up and down 2 - 3 times.	Top up to double the original sample volume		Top up to double the original sample volume		 Top up to double the original sample volume for samples ≤ 20 mL Top up to 50 mL for samples > 20 mL
5	Vortex RapidSpheres™. NOTE: Particles should appear evenly dispersed.	30 seconds		30 seconds		30 seconds
6	Add RapidSpheres™ to sample.	50 µL/mL of original sample volume NOTE: No incubation, IMMEDIATELY proceed to next step		50 μL/mL of original sample volume NOTE: No incubation, IMMEDIATELY proceed to next step		50 μL/mL of original sample volume NOTE: No incubation, IMMEDIATELY proceed to next step
7	Place the tube (without lid) into the magnet and incubate.	RT for 5 minutes		RT for 5 minutes		RT for 5 minutes
8	Carefully pipette** (do not pour) the enriched cell suspension into a new tube. NOTE: Collect the entire clear fraction from top to bottom. For optimal recovery, also collect a small volume of RBCs (up to 10% of the starting sample volume).	Use a new 5 mL tube		Use a new 14 mL tube		Use a new 50 mL tube
9	Add RapidSpheres™ to the new tube containing the enriched cells and mix.	Use same volume as in step 6 NOTE: No incubation, IMMEDIATELY proceed to next step		Use same volume as in step 6 NOTE: No incubation, IMMEDIATELY proceed to next step		Use same volume as in step 6 NOTE: No incubation, IMMEDIATELY proceed to next step
10	Remove the tube from the magnet; Place the tube from step 9 (without lid) into the magnet and incubate for a second separation.	RT for 5 minutes		RT for 5 minutes		RT for 5 minutes
11	Carefully pipette ^{**} (do not pour) the enriched cell suspension into a new tube. NOTE: Collect only the clear fraction.	Isolated cells are ready for use		Isolated cells are ready for use		Isolated cells are ready for use

RT - room temperature (15 - 25°C)

** Collect the entire supernatant, all at once, into a single pipette (e.g. for EasyEightsTM 5 mL tube use a 2 mL serological pipette [Catalog #38002]; for EasyEightsTM 14 mL tube use a 10 mL serological pipette [Catalog #38004]).



Directions for Use – Fully Automated RoboSep[™] Protocol

See page 1 for Sample Preparation and Recommended Medium. Refer to Table 4 for detailed instructions regarding the RoboSep[™] procedure. NOTE: If using RoboSep[™]-S, ensure the software is at least v.1.2.0.2 and a carousel compatible with this product is installed. Contact us at techsupport@stemcell.com for more information.

Table 4. RoboSep[™] Direct Human PBMC Isolation Kit Protocol for LEUKAPHERESIS

STEP	INSTRUCTIONS	RoboSep™ (Catalog #21000)			
	Prepare sample within the volume range.	1 - 6 mL			
1	Add sample to required tube.	14 mL (17 x 95 mm) polystyrene round-bottom tube (e.g. Catalog #38008)			
2	Add EDTA to sample.	At a final concentration of 6 mM EDTA			
3	Select protocol.	EasySep Direct Human PBMC Isolation 19654 - WB CB BM LEUK LRSC			
4	Vortex RapidSpheres™. NOTE: Particles should appear evenly dispersed.	30 seconds			
5	Load the carousel.	Follow on-screen prompts			
-5-	Start the protocol.	Press the green "Run" button			
6	Unload the carousel when the run is complete.	Isolated cells are ready for use			

Notes and Tips

• EasySep[™] Direct Human PBMC Isolation Kit is not suitable for use with downstream magnetic positive selection.

• If performing an EDTA-sensitive downstream application, wash the isolated cell fraction in desired medium prior to use.

ASSESSING PURITY

For purity assessment of residual RBCs by flow cytometry, use the following fluorochrome-conjugated antibody clones:

- Anti-Human CD235ab (Glycophorin A/B) Antibody, Clone HIR2 (Catalog #60111), and
- · Anti-Human CD41 Antibody, Clone HIP8 (Catalog #60114), and
- Anti-Human CD45 Antibody, Clone HI30 (Catalog #60018)

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