miniPURE-EVs

Size Exclusion Chromatography columns for Exosome and Microvesicle isolation Product Code: HBM-mPEV-##. Quantity: 10 or 20 SEC columns



About miniPURE-EVs Columns.

Size Exclusion Chromatography (SEC) is a very efficient method for separating EVs from the circulating proteins not affecting the original shape and functionality of the vesicles. miniPURE-EV is a SEC column designed for isolating EVs in a fast and easy way form small volume amount of different fluids. Additionally the column can be used for removal of small molecules from purified EVs, as the eccess of a dye after EV labeling procedure.

Fluid	Volume amount	
Plasma	100 µl up to 500 µl	
Serum	100 µl up to 500 µl	
Cell medium	100 µl up to 500 µl, pre-concentrated 10 folds.	
Urine	100 µl up to 500 µl, pre-concentrated 10 folds.	
Other samples	100 µl up to 500 µl	

Procedure for EV Isolation.

1. Sample preparation.

Prepare the sample by centrifugation steps as suggested in the table below:

Table Delow.			
Fluid	Suggested	Optional	
Plasma	10 min at 300 g (save super) 20 min at 1200 g (save super)		
Serum	10 min at 300 g (save super) 20 min at 1200 g (save super)	30 min at 10000 g (to eliminate particleses > 200 nm)	
Urine	10 min at 300 g (save super) Concentrate 10 folds*.		
Cell media	10 min at 300 g (save super) 20 min at 1200 g (save super) Concentrate 10-fold*.		

^{* **}It is recommended the use of TFF-Easy for concentrating the diluted fluids. Alternatilvely MWCO concentrators (100K) can be used.

2. Column preparation.

- miniPURE-EVs columns are provided with layer of preservative buffer.
- Open <u>first the upper</u> and <u>then the lov</u> cap of the column and let to flow almost the buffer throught the column, avoiding dry the surface of the gel.
- Wash the column with 3 volumes of PBS buffer (3 \times 4 ml) to eliminate preservative buffer residues.

3- Sample loading.

- Rinse the column with 100 μl up to 500 μl of sample containing EVs.
- Collect 100 μl fractions.
- When the sample is inside the gel matrix rinse the column with PBS 1x.

PBS 1x is the mobile phase of SEC column, do not let the column to get dried.

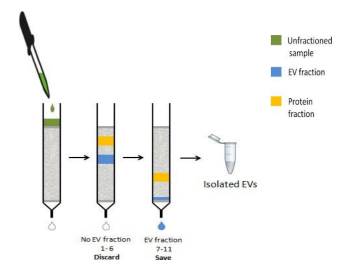
4. EV isolation.

Separation of EV and circulating proteins proceeds as indicated in the figure 1 (collecting 100 µl fractions).

5. Column washing.

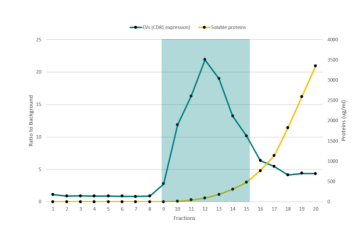
- After all fractions are collected wash the column from the residues of sample with approximately 10 ml of PBS. Never get the column dried. After the last washing step add to the column 0.5 ml of PBS 1x and close the caps.

Columns can be stored at 4°C and reused up to 5 times.



EV Separation.

miniPURE-EVs column was filled with 500 μ l of medium from HCT116 cells, previously concentrated by TFF-Easy. 20 fractions (100 μ l each one) have been collected and analyzed by ELISA ExoTEST^M assay (CD81 marker) and by BCA test for determining respectively vesicle and total protein content. EVs are eluted in fractions 9 - 14 (turnaround time approximately 10 min), whereas circulating proteins corresponded to the fractions 16 - 20.





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