



For research use only

Protocol

CD9 Fab-TACS® Exosome Agarose Column Starter Kit

Cat. no. 6-3319-002

human, for cell culture supernatant, serum and plasma

1. GENERAL INFORMATION & TECHNICAL SPECIFICATIONS

Kit components:

Cat. no.	Product	Quantity	Required/isolation
6-6310-300	Strep-Tactin® TACS Agarose Column, 0.3 ml	2	1
6-8019-150	CD9 Fab-Strep, human, lyophilized, 50 µg	1	20 μg
6-6996-001	Biotin stock solution, 100 mM, 250 µl	1	~20 µl
6-3333-001	TACS Column Adapter (0.3 ml column)	1	1

Required: Buffer with pH 7.4 (e.g. PBS, TBS or HEPES buffer depending on downstream application).

Column Capacity: 2-3 x 10⁹ targeted exosomes per column (sample-dependent)

specifications: Reservoir volume: 3 ml; **For single use only!**

Storage: Store all components at 2 - 8 °C. Store reconstituted Fab-Strep at -80 °C.

Stability: 6 months after shipping.

Shipping: Room temperature

Hazards: Products are not classified as hazardous according to (EC) No 1272/2008 [CLP].

Material Safety Data Sheets are provided.

2. INITIAL PREPARATIONS

2.1. Reagent preparation

Allow the reagents to equilibrate to room temperature (RT) prior to use. For a sterile isolation, work under a safety cabinet. **The following volumes will be sufficient for one selection process**.

- **2.1.1** Filtrate approximately **15 ml** buffer to remove interfering particles (recommended: 0.2 μm cellulose acetate filter).
- **2.1.2.** Dissolve lyophilized Fab-Strep in **1 ml** filtrated buffer by carefully pipetting up and down (avoid foam formation). **Do not vortex!**



Required per column: **20 µg** Fab-Strep in **400 µl** buffer. Store remaining solution in aliquots at **-80 °C** (stable for 6 months) if not required immediately. Avoid multiple freeze-thaw cycles.

2.1.3. Prepare 1 mM Biotin Elution Buffer by adding 20 μI of the 100 mM Biotin stock solution to 2 mI filtrated buffer (2.1.1.). Mix thoroughly.

2.2. Sample preparation

2.2.1. Cell culture supernatants: Centrifuge cell culture supernatant at 3000 x g for 10 min or 2000 x g for 30 min in advance. Filtrate supernatant (recommended: 0.22 μm polyethersulfone filter. Do not use cellulose acetate filters!).



If cells need to be collected as well, first centrifuge supernatant at 300 x g for 10 min and continue with further centrifugation steps using the supernatant.

2.2.2. Serum and plasma: Sediment blood for 30 min at room temperature. Centrifuge serum/plasma twice at 3000 x g for 10 min. Filtrate supernatant (recommended: 0.22 μm polyethersulfone filter. Do not use cellulose acetate filters!)

2.3. Column preparation



- **2.3.1. Remove** the caps at the top and at the bottom of the column. Allow the storage solution (contains sodium azide) to drain. Place the Strep-Tactin® TACS Agarose Column into the TACS Column Adapter.
- **2.3.2. Wash** the Strep-Tactin[®] TACS Agarose Column by applying **1 ml** buffer and allow the buffer solution to enter the packed bed completely.

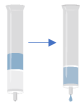


- **2.3.3. Load** the **400 μl** Fab-Strep solution (2.1.2.) onto the Strep-Tactin[®] TACS Agarose Column. Let the Fab-Strep solution enter the packed bed completely. Incubate for **2 min**.
- **2.3.4. Wash** the Strep-Tactin[®] TACS Agarose Column with **1 ml** buffer. Discard effluent and change collection tube. Strep-Tactin[®] TACS Agarose Column is now ready for exosome isolation.



Do not interrupt the procedure for more than 60 min.

3. PROTOCOL



3.1.1. Load

Apply prepared sample (2.2.) in steps of **3 ml** (max.: 7 ml in total). Collect flow-through containing unwanted material.



3.1.2. Wash

Apply 3 x 3 ml buffer. (In each step: Let the buffer solution enter the gel bed completely). The agarose bed should now be white again.



3.1.3. Elute

From this step on your effluent contains your target exosomes. Use a **new collection tube**. Apply **400 \muI** Biotin Elution Buffer (2.1.3.) and incubate for **5 min**. Elute target cells by applying **3 x 400 \muI** Biotin Elution Buffer.



Optional: Use size exclusion chromatography or hydrostatic filtration dialysis as an additional step to remove biotin and Fab-Streps for an ultra-pure exosome suspension.



Check our Downloads page

https://www.iba-lifesciences.com/resources/download-area/

for the latest version of this protocol



Info on warranty / licensing and trademarks available at:

www.iba-lifesciences.com/patents-licenses-trademarks/



If you have any questions, please contact

strep-tag@iba-lifesciences.com

We are here to help!