Exosome – Human CD9 Isolation (from cell culture)

\bigcirc	Package Contents	Catalog NumberSize10614D2 mL		
	Storage Conditions	 Store at 2°C to 8°C. When stored as instructed, expires two year from date of receipt unless otherwise indicated on product label. 		
	Required Materials	1 List of Materials		
	Timing	Hands-on time: 45–60 minutesIncubation time: 16–24 hours		
R	Selection Guide	Exosome Research Products Magnetic Separators Go online to view related exosome products and magnets.		
<u></u>	Product Description	 Exosome – Human CD9 Isolation (from cell culture) is intended for isolation of CD9-positive human exosome subsets from a pre-enriched exosome sample. After isolation, exosomes can be characterized by downstream applications including western blot, qRT-PCR, and sequencing. Dynabeads[®] are uniform, superparamagnetic polystyrene beads (2.7 µm dia.) coated with a primary monoclonal antibody specific for the CD9 membrane antigen expressed on most human exosomes. The Dynabeads[®] magnetic beads are incubated with your samples overnight and isolated exosomes are magnetically separated. 		
	Important Guidelines	 Follow the recommended pipetting volumes and incubation times. Avoid air bubbles (foaming) during pipetting. The western analysis is dependent on the level of exosomes present in the pre-enriched exosome sample, the protein transfer efficiency, the quality of the western blotting antibody and detection system (chromogenic detection is not recommended). 		
	Online	Visit our product pages for additional information and protocols. For support,		



Protocol outline

- 1. Pre-enrich exosomes.
- 2. CD9 positive isolation.
- 3. Protein electrophoresis.
- 4. Western blot analysis.

Pre-enriched exosome sample input

Pre-enriched exosome solution can be prepared using Total Exosome Isolation (from cell culture media) reagent, (Cat no 4478359) or ultracentrifugation.

Pre-enriched Exosome sample	Isolation Buffer	Dynabeads	Final Volume (after buffer exchange)
200 µL	0 µL	80 µL	200 µL
100 µL	0 µL	40 µL	100 µL
10 µL*	90 μL	40 µL	100 µL
1 µL	99 μL	40 µL	100 µL

Note: The protocol can be scaled up from 100 μ L to 5 mL by adjusting all volumes proportionally.

* Titration of exosome input is recommended: starting with 100 mL conditioned cell culture medium, concentrated to 2 mL after pre-enrichment (50x concentrated), use 10 μ L pre-enriched exosomes as starting sample (equals 500 μ L conditioned cell culture medium).

Ouidelines for optimal mixing conditions

Good mixing is critical to successful exosome isolation. Use a mixer that tilts and rotates to ensure that the beads do not settle in the tube.

Example of CD9 western blot analysis

Limited product warranty and disclaimer details

CD9 positive isolation

This protocol is designed for one isolation. The protocol can be scaled according to the desired number of analyses to be performed. The protocol below describes an exosome input of $10 \,\mu$ L pre-enriched exosome solution with $40 \,\mu$ L of bead solution.

	Timeline		Steps	Actions
Day 1	1		Prepare exosome – human CD9 isolation beads	 Place vial of beads on a roller for >10 minutes or vortex for 30 sec to resuspend. Transfer 40 μL bead solution to a tube containing 1 mL Isolation Buffer. Place the tube in magnetic separator for 1–2 min. Remove the buffer.
	2	10µL 90µL	Mix isolation beads with pre- enriched exosome sample	 Add 90 μL Isolation Buffer to tube containing beads. Add 10 μL pre-enriched exosome sample.
	3		Incubate beads and exosomes	Incubate at 2–8°C overnight with end-over-end mixing (tilting and rotation).
Day 2	4		Isolate bead-bound exosomes with magnetic separator	 Spin sample tube briefly 1–2 sec. Add 1 mL of Isolation Buffer and place tube in magnetic separator for 1–2 min before removing all supernatant. Remove tube from magnetic separator. Add 0.5 mL of Isolation Buffer and place tube in magnetic separator for 1–2 min before removing supernatant.
	5		Proceed to downstream analysis	 Western blot analysis qPCR Sequencing

Western blot analysis after CD9 positive exosome isolation

- Electrophoresis should be performed using a 5–15% gradient gel, or 12% homogeneous gel.
- Exosomal markers such as CD9, CD81, and CD63 should be separated under non-reducing conditions.
- For detection of proteins that are equal in size to antibody heavy- or light-chains (e.g. CD9) we recommend the Mouse TrueBlot[®] Ultra Ig HRP Secondary antibody (eBioscience Cat. no. 18-8817).

	Timeline		Steps	Actions
Day 2	1	»)) ((Lyse exosomes	 Add 10 µL 1X RIPA buffer with protease inhibitors to bead bound exosomes. Sonicate for 10 sec. Incubate on ice for 15 min.
	2	>>>	Denature protein sample	 Add 10 μL 2X sample buffer (with or without reducing agent). Add 1 μL loading buffer. Incubate at 95°C for 5 min.
	3		Load sample on polyacrylamide gel	 Spin sample tube briefly 1–2 sec. (<i>Optional</i>) Place tube on magnetic separator. Pipette sample into well.
	4		Perform electrophoresis	 Add appropriate molecular weight markers. Perform electrophoresis at 200 V for 30 min (or according to your standard protocol).
Day 3	5		Perform western blot	 Perform wet transfer at 100 V for 1 hour on ice. Incubate with primary antibody overnight. Detect protein by chemiluminescence.