**Human Sample Processing and Isolation of Extracellular Vesicles with size exclusion chromatography**

J. Nathaniel Diehl1, Amelia Ray1, Andrew Peterson2, John S. Ikonomidis2,3, and Adam W. Akerman2

1University of North Carolina School of Medicine, Chapel Hill, North Carolina.

2Department of Surgery, University of North Carolina – Chapel Hill, Chapel Hill, North Carolina.

3Division of Cardiothoracic Surgery, University of North Carolina – Chapel Hill, Chapel Hill, North Carolina.

Abstract:

This protocol details the steps necessary to isolate circulating plasma extracellular vesicles (EVs) from human peripheral blood samples. This protocol utilizes an automated fraction collector (AFC) and qEV size-exclusion chromatography columns from IZON Science. This is intended to serve as the first step in the workflow for EV quantification along with “Measurement of extracellular vesicles with tunable resistance pulse sensing (TRPS).”

Keywords:

Extracellular vesicles, EVs, exosomes, TRPS, tunable resistive pulse sensing, human plasma, automatic fraction collector, AFC, qEV, size exclusion chromatography, qNano Gold

Guidelines:

* All human blood samples should be handled according to the Centers for Disease Control (CDC) universal blood and body fluid collection guidelines. Additionally, sample handling and processing should follow the Occupational Safety and Health Administration (OSHA) blood borne pathogens procedures to prevent possible pathogen transmission.
* This protocol details the extraction of EVs from human peripheral blood samples. All human subjects research should be reviewed and approved by the site-specific Institutional Review Board (IRB) prior to proceeding.
* qEVsingle columns are used in this protocol. Newer generation multi-use columns can also be used if column-specific instructions are followed.

Before Start:

* Ensure that you have an adequate quantity of buffer (1x sterile-filtered PBS)
* If you already have frozen human plasma samples, skip to “EV Isolation”.

Safety Warnings:

Appropriate personal protective equipment (PPE) should be worn (nitrile gloves, safety goggles, and lab coat). Store all organic solvents in a flammable storage cabinet in accordance with institution policy. Utilize biohazard waste containers for sample waste.

Materials:

* Sterile alcohol prep pads, Fisher Scientific, Cat#22-363-750
* 25-gauge Safety-Lok needle and collection set, BD, SKU#367285
* BD Vacutainer EDTA tube, Fisher Scientific, Cat#23-021-013
* Cryogenic tubes with screw cap, Fisher Scientific, Cat#03-337-7Y
* Microcentrifuge tubes, Eppendorf, Cat#022364111
* qEVsingle 35 nm Legacy column, IZON, PC#SP6
* 200 μL filtered pipet tips, Millipore Sigma, Cat#CLS4823-960EA
* 1000 μL filtered pipet tips, Millipore Sigma, Cat#CLS4809-1000EA
* 10 mL Sterile syringe with Luer lock, Fisher Scientific, Cat#14-955-460
* 0.22 μm Luer lock inlet filters, Thomas Scientific, Cat#1176G49
* Sterile filtered (0.22 μm) phosphate-buffered saline (PBS), ThermoFisher, Cat#18912014
* Filtered (0.22 μm) deionized water

Equipment:

* Ultra-low temperature (-80°C) freezer
* Standard manual defrost laboratory (-20°C) freezer
* Refrigerated microcentrifuge, Eppendorf, Cat# 022620700
* Clinical Centrifuge, Globe Scientific, Item# GCC-E
* qEV Automatic fraction collector (AFC)
* 20-200 μL pipette
* 100-1000 μL pipette

Procedural Steps:

*Human plasma collection*

1. 5 mL peripheral venous blood is collected by a trained nurse or phlebotomist into prelabeled EDTA-coated evacuated tubes.
	1. EDTA anticoagulant is recommended for isolation of EVs
2. Immediately after the sample is collected, the tube should be thoroughly mixed and stored at room temperature (< 2 hours).
3. Centrifuge whole blood at 2,500 x g for 15 minutes at room temperature.
4. Collect the topmost layer (plasma) of supernatant into a 15 mL conical tube.
	1. Carefully avoid disruption of the next layer (buffy coat). Leave 1 cm of plasma above the buffy coat.
5. Centrifuge the plasma again at 2,500 x g for 15 minutes at room temperature.
6. Avoiding the bottom ~100 μL of plasma, collect the topmost plasma into a new 15 mL conical tube.
7. Aliquot desired volumes into labeled freezer-safe microcentrifuge tubes.
	1. We recommend a minimum aliquot volume of 250 μL.
8. Snap freeze plasma fractions and store at -81°C.

*EV Isolation*

1. Thaw human plasma at room temperature and centrifuge plasma 2,500 x g for 15 minutes at 4°C.
2. Power on IZON automated fraction collector (AFC).
3. Select SETUP > Calibration. Follow the on-screen prompts provided by the AFC to calibrate the machine using the provided 10 g weight.
4. Insert a new 35 nm qEVsingle column into the column mount and allow the machine to register it (the display will indicate specific column features once it has registered).
	1. Column settings: qEV = 35 nm, count = 4, size = 0.2 mL, void = 0.8 mL, sample = 0.15 mL
5. Follow on-screen prompts, click OK to proceed to next step. Permit existing buffer + 2 mL additional 1x sterile filtered PBS to flush the column.
6. Once flush has completed, click OK to proceed to next step. Load 150 μL human plasma sample in a drop-wise manner into the center of the column.
7. Click OK to proceed to next step. AFC should move from flush position to collect column buffer into the central well.
8. Once the sample has completely entered the column, carefully load 2 mL 1x PBS into the top of the column in a drop-wise manner. Continue to add buffer as needed until the extraction is complete.
	1. If the collection stops at this point, ensure adequate buffer remains above the beads in the column and click OK to proceed.
9. Once the AFC has finished the extraction, combine the first three fractions and discard or store the fourth.
	1. The fourth fraction contains primarily protein and a very low concentration of EVs.
10. Discard the used qEVsingle column and clean the central well with a Kimwipe delicate task wipe.
	1. Ensure the central well is completely dry before next use.
11. Repeat steps 4-10 for additional samples.
12. Remove peristaltic pump tubing from below the column mount by removing the plastic cover on the peristaltic pump. Flush plastic tubing with 10 mL syringe and 5 mL filter water and replace plastic tubing on AFC.
	1. This will prevent buildup of salt crystals in the tubing between uses.
13. Combined fractions 1-3 can be immediately processed for use on TRPS, or stored short term (1-2 weeks) at -20°C or long term (> 2 weeks) at -81°C.

Citations:

1. IZON AFC quick start guide 🡪 https://files.izon.com/hubfs/Manuals,%20Technical%20Notes%20and%20Customer%20Support/AFC/AFC%20Quick%20Start%20Guide.pdf
2. IZON Full Manual 🡪 https://f.hubspotusercontent30.net/hubfs/4136435/Manuals,%20Technical%20Notes%20and%20Customer%20Support/AFC/rotational-afc-user-manual-RA1-OQ-001.pdf

Additional Notes:

* If the AFC continuously stops running in the middle of an extraction and it is not due to running out of buffer, check that the tubing is not obstructed by removing it from the tubing apparatus and confirming that liquid can be pushed through. Tubing can be replaced if necessary.
* For further troubleshooting or information, please refer to the IZON AFC Quick Start Guide and the IZON AFC Full Manual.