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Kateena Addae Konadu, Morehouse School of Medicine
Ming Bo Huang, Morehouse School of Medicine
William Roth, Morehouse School of Medicine
Wendy Armstrong, Emory University
Michael Powell, Morehouse School of Medicine
Francois Villinger, Emory University
Vincent Bond, Morehouse School of Medicine

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**Isolation of Exosomes from the Plasma of HIV-1 Positive Individuals**

Kateena Addae Konadu¹, Ming Bo Huang¹, William Roth¹, Wendy Armstrong², Michael Powell¹, Francois Villinger³,4, Vincent Bond¹

¹Department of Microbiology, Biochemistry, Immunology, Morehouse School of Medicine
²Department of Medicine, Emory University School of Medicine
³Department of Pathology and Laboratory Medicine, Emory University School of Medicine
⁴Yerkes National Primate Research Center

Correspondence to: Vincent Bond at vbond@msm.edu

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**Abstract**

Exosomes are small vesicles ranging in size from 30 nm to 100 nm that are released both constitutively and upon stimulation from a variety of cell types. They are found in a number of biological fluids and are known to carry a variety of proteins, lipids, and nucleic acid molecules. Originally thought to be little more than reservoirs for cellular debris, the roles of exosomes regulating biological processes and in diseases are increasingly appreciated.

Several methods have been described for isolating exosomes from cellular culture media and biological fluids. Due to their small size and low density, differential ultracentrifugation and/or ultrafiltration are the most commonly used techniques for exosome isolation. However, plasma of HIV-1 infected individuals contains both exosomes and HIV viral particles, which are similar in size and density. Thus, efficient separation of exosomes from HIV viral particles in human plasma has been a challenge.

To address this limitation, we developed a procedure modified from Cantin et. al., 2008 for purification of exosomes from HIV particles in human plasma. Iodixanol velocity gradients were used to separate exosomes from HIV-1 particles in the plasma of HIV-1 positive individuals. Virus particles were identified by p24 ELISA. Exosomes were identified on the basis of exosome markers acetylcholinesterase (AChE), and the CD9, CD63, and CD45 antigens. Our gradient procedure yielded exosome preparations free of virus particles. The efficient purification of exosomes from human plasma enabled us to examine the content of plasma-derived exosomes and to investigate their immune modulatory potential and other biological functions.

**Video Link**

The video component of this article can be found at http://www.jove.com/video/53495/

**Introduction**

The HIV-1 epidemic continues to have a significant impact throughout the world. As of 2013, approximately 35 million people worldwide were living with HIV, and 2.1 million of these were newly infected individuals¹. Prevention strategies and increased access to antiretroviral therapy have been helpful in reducing the overall acquisition of HIV. However, individual populations are still experiencing rises in the acquisition of HIV¹. Thus, there is a need for continued efforts to address this epidemic.

One of the strongest predictors of HIV disease progression is chronic immune activation (CIA)²-15. Defined by persistently high levels of detectable cytokines and elevated expression markers on the surface of T lymphocytes, CIA has been attributed to: (i) continuous dendritic cell production of Type I IFN¹ⁱ; (ii) direct immune activation driven by HIV proteins Tat, Nef and gp120¹²; (iii) translocation of bacterial proteins in gut associated immune cells⁶. However, the exact mechanism(s) underlying chronic, systemic immune activation in HIV infection remain to be fully elucidated.

Our group and others have demonstrated a role of exosomes in HIV pathogenesis¹⁶-¹⁸. Our group has determined that the Nef protein is excreted from infected cells in exosomes¹⁵, and exosomal Nef (exNef) is present in the plasma of HIV-infected individuals at nanogram levels¹⁸. We have shown that bystander CD4+ T-cells exposed to exNef resulted in activation-induced cell death dependent on the CXCR4 pathway¹⁹, ²⁰. Alternatively, monocyte/macrophages were refractory to exNef-induced apoptosis, but exhibited altered cellular functions and cytokine expression. Most recently, our group has shown exosomes isolated from the plasma of HIV-infected individuals contain a variety of pro-inflammatory cytokines. Further, naïve peripheral blood mononuclear cells exposed to plasma-derived exosomes from HIV-infected patients induced expression of CD38 on naïve and central memory CD4+ and CD8+ T cells. This likely contributes to systemic inflammation and viral propagation via bystander cell activation¹⁴, and suggests that exosomes play a significant role in HIV pathogenesis.

In investigating the role of exosomes in HIV pathogenesis, one challenge is developing techniques to efficiently separate exosomes from HIV particles while maintaining the exosomal content as well as their functional immune modulatory capability. Several methods have been described...
for isolating exosomes from cell culture and biological fluids. Because of their small size and low density (exosomes float at a density of 1.15 - 1.19 g/ml), differential ultracentrifugation and/or ultrafiltration are the most commonly used techniques for exosome isolation. However, HIV-infected cell culture supernatants and patients' plasma contain both exosomes and HIV-1 viral particles. Exosomes and HIV-1 particles are very similar in both size and density. Alternatively, taking advantage of the expression of unique exosomal markers such as CD63, CD45, and CD81, exosomes have been isolated using immunoaffinity capture methods. This procedure can separate virus from exosomes. However, the drawback of this technique is the tight attachment of antibodies to the purified exosomes, which could interfere with assessment of the immunomodulatory potential of exosomes in culture.

To address these limitations, we developed a procedure for purification of exosomes from HIV particles in human plasma modified from Cantin and coworkers using iodixanol velocity gradients. Exosomes were found to segregate in the low-density/upper fractions of iodixanol gradients, whereas virus particles segregated in the high-density/lower fractions. Virus particles were identified by p24 ELISA and exosomes were identified using exosome markers AChE, CD9, CD63, and CD45. The upper low-density fractions collected contained exosomes which were negative for HIV-1 p24 contamination. The efficient purification and separation of exosomes from HIV particles in human plasma allows for accurate examination of the content of exosomes derived from human plasma as well as the investigation of their immune modulatory potential and the diagnostic and prognostic value of exosomes in HIV-1 pathogenesis.

A general diagram of the exosome isolation and purification procedure is provided in Figure 1. Whole blood was obtained from healthy volunteer donors and from HIV-positive individuals not receiving antiretroviral therapy attending the Hope Clinic of Emory University and the Infectious Disease Program of Grady Health System in Atlanta, Georgia. This study was approved by the institutional review boards of Emory University and Morehouse School of Medicine. All persons participating in the study gave written and informed consent.

### 1. Preparation of Exosomes from Blood Plasma

#### 1. Human Blood Collection and Processing

1. Collect 10 ml of peripheral blood by venipuncture in EDTA blood collection tubes (containing 18 mg potassium EDTA), and gently invert five times to mix.
2. Centrifuge the blood collection tubes at 1,000 x g for 20 min at RT to pellet blood cells. Use a sterile pipette to transfer the plasma fraction (4-5 ml) to a 25 ml conical tube. Dilute the plasma with 10 ml of 1 X PBS. Discard blood cells (red blood cells and white cells, also known as PBMC) in an appropriately marked container for biohazard waste.

**NOTE:** In the case of uninfected donor blood samples, the PBMC can be reserved for further use (see procedure IV.2, below).

3. Store the plasma samples at 4 °C for short term (2-3 days) or at -80 °C for longer term storage.

**NOTE:** Bring any frozen plasma samples to 4 °C before further processing.

#### 2. Preparation of Exosome Fraction from Plasma

1. Centrifuge plasma at 10,000 x g for 30 min at 4 °C to remove cellular debris. Use a sterile serological pipette to transfer the cleared plasma supernatant to a clean 25 ml ultracentrifuge tube. Discard the pellet in the biohazard container.

2. Centrifuge the cleared plasma at 100,000 x g for 2 hr at 4 °C to remove large vesicles. Remove the 100,000 x g supernatant carefully by pipetting, and discard in biohazard waste. Resuspend the 100,000 x g pellet in 1 ml of 1X PBS in a clean tube and incubate at RT for 30 min, swirling gently to dislodge and separate particles.

3. Wash the suspended 100,000 x g exosome/virus pellet by adding 25 ml of PBS. Gently invert the tube five times to mix, then centrifuge again at 100,000 x g for 2 hr at 4 °C. Discard the PBS wash solution in the biohazard waste container.

4. Resuspend the 100,000 x g pellet in 1 ml of 1 X PBS and incubate at RT for 30 min swirling gently to dislodge and dissolve the pellet.

**NOTE:** If it is not possible to proceed directly to the iodixanol gradient step, store the pellet, containing exosomes and virus particles, at 4 °C for 1-2 days until the iodixanol gradient step.

#### 3. Purification of Exosomes

1. Generate 6%-18% velocity gradients of iodixanol using a dual-chamber gradient former apparatus.

   - Prepare 6% and 18% solutions of the iodixanol reagent, supplied as a 60% solution in water, by dilution in PBS. Pipet 5.5 ml of the 18% solution into the stirred chamber, and pipet 5.5 ml of the 6% solution into the reservoir chamber. Turn on the stirrer, open the stopcock, and allow each gradient to flow into a 14 ml ultracentrifuge tube.

   **NOTE:** Either use immediately or store prepared gradients O/N at 4 °C prior to use.

   2. Carefully layer 1 ml of the exosome/virus solution onto the top of each 11 ml gradient. Centrifuge gradients at 250,000 x g for 2 hr at 4 °C, using a SW40Ti swinging bucket rotor.

   3. Label twelve (12) 1.5 ml microcentrifuge tubes. Remove 1.0 ml from the top of the gradient and transfer to tube #1. Transfer the remaining 1ml fractions to tubes 2-12 in sequential order.

   **NOTE:** The topmost fraction will thus be #1, the bottom fraction, #12. Store the gradient fractions at 4 °C. The purified exosomes, which should be in the fractions 1-3 at the top of the gradient, are stable for 3-4 weeks after stored at 4 °C.

#### 4. Exosome Characterization

1. **Acetylcholinesterase (AChE) Activity Assay**

   - Prepare 100 mM substrate stock by mixing acetylthiocholine iodide 28.9 mg in 1 ml of 1X PBS. Store substrate stock at -20 °C up to 1 month.
2. Prepare 10 mM color indicator stock by mixing benzoic acid 39.6 mg and sodium bicarbonate 15 mg in 10 ml of 1X PBS. Store color indicator stock at 4 °C up to two weeks.

3. Prepare assay reagent by mixing 1X PBS, substrate, and color indicator in a ratio of 100:2:5 (for example, 10 ml 1X PBS + 200 μl substrate + 500 μl color indicator).

4. Transfer 50 μl each from 1.5 ml gradient fraction tube (labeled 1-12, from procedure III.3) to the wells of a 96-well microtiter plate. Prepare duplicate wells for each gradient sample.

5. Prepare a set of standards by first making a 2000 μU/ml AChE stock in PBS. Make eleven (11) 2-fold serial dilutions of this stock and add 50 μl of each dilution to a single well of a microtiter plate, so that standard #1 = 2,000 μU/ml, #2 = 1,000 μU/ml, #3 = 500 μU/ml, etc. until #12 = 0.98 μU/ml. The twelve AChE standards will thus occupy a single row of a 96-well assay plate.

6. Add 200 μl of assay reagent mixture to each well and incubate 20 min (in the dark) at RT to allow development of the colored reaction product. Measure AChE activity at a wavelength of 450 nm using a fluorescent microplate reader.

NOTE: The percentage of starting material remaining after exosome purification is assessed by AChE assay of the unfraccionated plasma.

2. **Immunoblot Analysis**

1. Separate proteins in gradient fractions 1-12 (from procedure 2.1.3) by SDS-PAGE on 4-20% Tris-HCl precast gels at 100V x 1 hr. Transfer proteins in the gel to a nitrocellulose membrane using an electro-blotting transfer cell according to the manufacturer's instructions. Allow the transfers to blot for 12-16 hr (O/N) at 350V.

2. Remove the membrane from the blotting apparatus and wash in Tris-Buffered Saline (TBS) for 5 min. Block with 5% nonfat milk in TTBS (TBS with 0.1% Tween 20) for 1 hr by shaking at RT.

3. Incubate the membrane with primary antibodies specific for exosomes (CD9, CD45, CD63) or HIV-1 capsid protein (p24) in 5% nonfat milk with shaking at 4 °C for 12-16 hr (O/N). Dilutions of primary antibodies for immunoblots range from 1:2,000 -1:5,000.

4. Wash the blot in TTBS for 20 min, followed by incubation with a 1: 2,000 dilution of horseradish peroxidase (HRP)-conjugated to anti-IgG (secondary antibody), in 5% nonfat milk for 1 hr at RT. Wash the blot 3 times with TTBS, 10 min per wash.

5. Incubate the blot with luminol substrate reagent according to the manufacturer's directions, to generate a chemiluminescent signal from the HRP-labeled proteins Detect the chemiluminescent signal using an electronic imaging system with CCD camera.

6. Save the images as TIFF files which can be viewed in Adobe Photoshop. Perform densitometry analysis of bands using ImageJ software (National Institutes of Health, Bethesda, MD).

3. **Cytokine Assay**

1. Prior to the cytokine assay, disrupt immune complexes that are usually present in human plasma by acid dissociation and lyse the exosomes by detergent treatment. Assay both starting plasma samples and purified exosome preparations.

   a. To 100 μl of human plasma add 100 μl 0.33 N HCL and incubate at 37 °C for 1 hr. Add 100 μl 0.33 N NaOH to neutralize acid-treated plasma. Add Triton X-100 to both the neutralized plasma and purified exosome samples, to a final concentration of 1%, to cause lysis of exosomes.

2. For this assay (a fluorescent bead-based procedure), use magnetic beads pre-coated with antibodies by the manufacturer. Use a custom-designed panel of anti-human cytokine antibody-coated beads; listed in Table 1. Vortex the antibody-coated magnetic beads from the cytokine assay kit for 30 sec and add 50 μl beads to each well of a 96-well plate to be used in the assay.

3. Prepare a dilution series of standards in plasma standard buffer (both contained in the cytokine assay kit) as described in the instruction manual for the cytokine assay kit, to generate an 8-point standard curve.

4. Add 150 μl of 1X wash buffer (from kit) to each well and wash the beads using a magnetic bead washer, as described in the manual. Add 25 μl plasma assay buffer (from kit) to each well. Add 25 μl of standards and samples to all wells used in the assay. Add 25 μl of plasma sample buffer to negative control wells. Seal and shake the plate at 700 rpm for 60 min at RT and incubate O/N at 4 °C.

5. Add 150 μl of 1X wash buffer to each well and wash the plate (as in step 3.3.4, above). Add 25 μl of detection antibodies into each well, seal and shake plate at 700 rpm for 30 min at RT. Repeat the washing step.

6. Add 200 μl of assay reagent mixture to each well of a 96-well microtiter plate. Prepare duplicate wells for each gradient sample.

7. Add 120 μl of reading buffer (from kit) into each well and shake on a laboratory table top shaker at 700 rpm for 5 min at RT to allow the fluorescent signal to develop. Read plate on the plate reader as per the manufacturer's instructions.

8. In order to account for the amount of exosomes lost during the isolation procedure, use the following equation: [original reading / plasma volume] x 66.6= adjusted cytokine reading.

### 4. Assay for Immunomodulatory Potential

1. **Preparation of Culture Medium with Exosome-depleted Fetal Bovine Serum**

   - Centrifuge 500 ml of fetal bovine serum (FBS) at 100,000 x g for 20 hr at 4 °C to pellet contaminating exosomes and microvesicles. Carefully remove and save the exosome-depleted FBS supernatant. Discard the pellet in the biohazard waste.

   - Combine 20% of the exosome-depleted FBS supernatant with 500 ml of Roswell Park Memorial Institute 1640 (RPMI 1640) medium. Filter the medium through a 0.45 μm membrane filter.

   - Add streptomycin (100 U/ml), penicillin (100 U/ml), L-glutamine (2 mM), HEPES-buffered saline solution (10 μM), and IL-2 (20 U/ml) to the filtered medium.

2. **Cell Culture and Exosome Exposure**

   - Expose exosomes to donor peripheral blood mononuclear cells (PBMC) obtained from healthy clinic volunteers. Suspend PBMC in prepared culture medium and count the cells using a cell/particle counter (according to the manufacturer's standard method).

   NOTE: The PBMC are obtained during the plasma preparation described above (step 1.2).

   - Centrifuge 500 ml of fetal bovine serum (FBS) at 100,000 x g for 20 hr at 4 °C to pellet contaminating exosomes and microvesicles. Carefully remove and save the exosome-depleted FBS supernatant. Discard the pellet in the biohazard waste.

   - Add streptomycin (100 U/ml), penicillin (100 U/ml), L-glutamine (2 mM), HEPES-buffered saline solution (10 μM), and IL-2 (20 U/ml) to the filtered medium.
2. Using prepared medium from procedure 4.1.3, co-culture $3.0 \times 10^6$ (PBMC) with 1 μg/ml of pooled exosomes from three (3) HIV-1 seropositive or from three (3) HIV-1 seronegative individuals in a total volume of 1 ml in the wells of a 12-well culture plate (with lid). Incubate cultures for 48 hr at 37 °C.

3. Prepare untreated PBMC cultures and cultures treated with Concanavalin A (Con A; 5 μg/ml) to serve as negative and positive controls, respectively, as described above (4.2.2). Incubate all PBMC cultures for 48 hr at 37 °C.

4. Immediately prior to harvesting the cells, prepare dilutions of the following fluorochrome-conjugated monoclonal antibodies: Alexa Fluor 700-labeled anti-CD3 (1:400 dilution), allophycocyanin (APC)/cyanine 7 (Cy7)-labeled anti-CD4 (1:400), peridinin chlorophyll protein complex-labeled anti-CD4 (1:400), V450-labeled anti-CD8 (1:400), biotin-labeled anti-CD45RA (1:1,000), phycoerythrin (PE)/Cy7-labeled anti-CD62L (1:1,000), PE/cyanine 5 (Cy5)-labeled anti-CD38 (1:200), PE-Texas Red-labeled anti-streptavidin (1:2,000), and PE/Cy5-labeled mouse IgG1K isotype control (1:200).

5. After the 48 hr incubation period, harvest the PBMC. Wash PBMC in PBS to remove exosomes and stain the cells by incubation for 1 hr at 4 °C with individual fluorochrome-tagged antibodies. Analyze the stained PBMC by flow cytometry to quantify the expression of chemokines (as described).

**Representative Results**

Exosomes are efficiently purified from HIV-1 positive human plasma. Isolated exosomes, identified by acetylcholinesterase (AChE) activity, segregated in lower density fractions 1-3 at the top of the iodixanol gradients, whereas virus particles, identified by HIV-1 antigen p24, segregated in the higher-density fractions (10-12, near the bottom). The presence of exosomes was further confirmed by immunoblot identification of exosome markers AChE, CD9, CD45, and CD63, and by electron microscopy (Figure 2).

Pro-inflammatory cytokines and chemokines are associated with and significantly elevated in the exosomes of HIV-1 seropositive individuals. Purified exosomes and unfractionated plasma from HIV-1 infected and HIV-1 seronegative individuals were analyzed for 21 cytokines/chemokines by multiplex assay. All 21 cytokines/chemokines were detected in exosomes isolated from HIV-1 positive individuals. Additionally, their levels were significantly elevated as compared to plasma and exosomes from HIV-1 seronegative controls (Table 1).

CD38 expression was increased on the surface of cells exposed to exosomes from HIV-1 seropositive individuals. PBMCs from uninfected human donors were exposed for 48 hr to pooled exosomes from HIV-1 seropositive or seronegative individuals and assessed for levels of the activation marker CD38 on CD4+ and CD8+ T-cells via flow cytometry. We observed that by 48 hr post-exposure, CD38 expression on the surface of naïve and central memory CD4+ and CD8+ T cells were significantly elevated in T-cells exposed to exosomes from HIV-1 positive individuals compared to HIV-negative exosome treatment and untreated controls (Figure 3).
**Figure 1. Schematic representation of exosome isolation from HIV-1 positive human plasma.** (A) 10 ml of peripheral blood was collected from HIV-1 seropositive and seronegative individuals. (B) EDTA blood collection tubes were centrifuged at 1,000 x g for 20 min at RT. (C) Separated plasma was transferred to 50 ml conical tubes and diluted ½ with 1x PBS and (D) centrifuged 10,000 x g for 30 min. (E) The supernatant was transferred to ultracentrifuge tubes and (F) centrifuged at 100,000 x g for 2 hours, 4°C. (G) Discard supernatant and re-suspend pellet in 1ml 1X PBS. (H) Centrifuge 100,000 x g, 2 hours, 4°C. (I) Re-suspend pellet in 1ml 1X PBS and overlay on 6-18% iodixanol velocity gradient. (J) Centrifuge 250,000 x g, 2 hours, 4°C. (K) Piviot 12 (1ml) fractions into individually label tubes and prepare for AChE measurements and immunoblot analysis. (L) Combine fractions 2 & 3 and dilute with 4ml of 1X PBS. (M) Centrifuge 400,000 x g, 2 hours, 4°C. (N) Re-suspend exosome pellet in 1ml 1X PBS store at 4°C for in vitro assay and -80°C for protein characterization. Please click here to view a larger version of this figure.
Figure 2. Exosomes are efficiently purified from human plasma. Individual iodixanol velocity gradient fractions from HIV-seropositive or seronegative individuals were subjected to (A) enzymatic assay for acetylcholinesterase (AChE), and (B) Western blot analysis for exosomal markers CD9, CD45, and CD63 and viral protein p24 and were (C) immunolabeled with anti-CD63 and examined with electron microscopy to confirm preparation of purified exosomes (C). (Figure from Konadu et al, 201421, used by permission). Please click here to view a larger version of this figure.

Figure 3. Assay for Immunomodulatory Potential. PBMCs from HIV-1 seronegative individuals were exposed to either pooled exosomes from plasma of HIV-1 seropositive (HIV+ Exo) or seronegative (HIV- Exo) individuals, left untreated, or treated with 5 µg/ml of Concanavalin A (Con A) as a positive control. After 48 hr exposure, Naïve (CD45RA+/CD62L+) , Central (TCM; CD45RA-/CD62L+) and Effector (TEM; CD45RA-/CD62L-) memory CD4+ and CD8+ T-cells were analyzed for CD38 expression by flow cytometry. Exosome concentration was normalized by total protein and added at 1 µg/ml. Error bars represent mean +/- SEM of six independent donors. Difference between groups were tested for statistical significance by the One Way ANOVA Test. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. (Figure from Konadu et al, 201421, used by permission). Please click here to view a larger version of this figure.
Table 1. Analysis of purified exosomes and whole plasma from HIV-1 seropositive and seronegative individuals for pro-inflammatory cytokine and chemokine expression. Purified exosomes from HIV-1 seropositive or seronegative individuals were analyzed for pro-inflammatory cytokine and chemokine expression. All 21 pro-inflammatory cytokines and chemokines measured were associated with and were significantly elevated in exosomes of HIV-1 seropositive individuals compared to seronegative controls. Additionally, IL-1a, IFNa2, and CXCL10 were significantly elevated in the unfractionated plasma of HIV-1 seropositive individuals compared to seronegative controls. Difference between groups were tested for statistical significance by the Mann-Whitney U Test. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. (Figure from Konadu et al., 2014, used by permission). Please click here to view a larger version of this figure.

Discussion

Chronic immune activation (CIA) and CD4+ T cell depletion are two important hallmarks of HIV-1 infection. They have been established as predictors for pathogenesis, with CIA being the best predictor. However, the underlying mechanisms driving chronic systemic immune activation and CD4+ T cell decline still have not been fully elucidated. We and other labs have developed firm evidence that exosomes secreted from HIV-1 infected cells play a role in both hallmarks.

The continuing interest in both composition and function of extracellular vesicles has led to the publication of various methods for exosome isolation from both cell culture media and biological fluids. However, a barrier to investigating the role of exosomes in HIV-1 pathogenesis has been the efficient separation of exosomes from HIV-1 particles while maintaining the ability to investigate both exosome content and functional activity. We have developed a protocol for purification of exosomes from HIV-1 particles in human plasma, utilizing iodixanol velocity gradients. The HIV-positive donors used in this study had not received antiretroviral treatment and the plasma samples used for these experiments contained from 1500 to 400,000 virus particles/mL with an average of 206,000 virus particles/ml. Thus, we demonstrate that exosomes in the plasma of HIV-1 infected individuals can be efficiently separated from HIV-1 virus particles, even when the virus loads are high. Though similar in size and density, exosomes segregated in the low-density/upper fractions of the iodixanol gradients as compared to viral particles, which segregated in the high-density/lower fractions. The exosomes prepared by the iodixanol gradient method are highly purified and free of contaminating extracellular proteins. The purity of the isolated exosome population was confirmed using p24 ELISA and Western blot analysis. The procedure does have certain pitfalls and limitations. A considerable amount of the exosomes, often a majority of the starting material as calculated by measurements of AChE activity, are lost during the process. In addition, the procedure is also time consuming and requires access to expensive equipment. Finally, the generation of the iodixanol gradients using the gradient former apparatus requires considerable practice to ensure reproducibility. An
alternative method for gradient generation might be to prepare a graded series of iodixanol solutions, followed by careful layering of the solutions into centrifuge tubes and incubating ON to allow the gradients to form. However, we have not tested this alternative.

The experimental protocol described here for isolation and separation of exosomes from HIV-1 particles in human plasma is an effective method to ensure the purity of exosomes. Use of this method has led to exciting avenues for future inquiry regarding the role of exosomes in HIV-1 pathogenesis and could equally be used in inquiries for other biological processes.

Disclosures

The authors declare that they have no competing financial interests.

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