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RESEARCH ARTICLE



Comparative and integrated analysis of plasma extracellular vesicle isolation methods in healthy volunteers and patients following myocardial infarction

Daan Paget ^{1,2} Antonio Checa ³ Benedikt Zöhrer ^{4,5} Raphael Heilig ⁶
Mayooran Shanmuganathan ^{1,7} Raman Dhaliwal ⁸ Errin Johnson ⁸
Maléne Møller Jørgensen ^{9,10} Rikke Bæk ⁹ Oxford Acute Myocardial Infarction
Study (OxAMI) ^{1,7} Craig E. Wheelock ^{3,5,11} Keith M. Channon ^{1,7} Roman Fischer ⁶
Daniel C. Anthony ² Robin P. Choudhury ^{1,7} Naveed Akbar ¹

¹Division of Cardiovascular Medicine, Radcliffe Department of Medicine, University of Oxford, Oxford, UK

³Unit of Integrative Metabolomics, Institute of Environmental Medicine, Karolinska Institute, Stockholm, Sweden

⁴Respiratory Medicine Unit, Department of Medicine Solna and Center for Molecular Medicine, Karolinska Institutet, Stockholm, Sweden

⁵Department of Respiratory Medicine and Allergy, Karolinska University Hospital, Stockholm, Sweden

⁶Target Discovery Institute, Centre for Medicines Discovery, Nuffield Department of Medicine, University of Oxford, Oxford, UK

⁷Acute Vascular Imaging Centre, Radcliffe Department of Medicine, University of Oxford, UK

⁸Sir William Dunn School of Pathology, University of Oxford, Oxford, UK

⁹Department of Clinical Medicine, Aalborg University, Aalborg, Denmark

¹⁰Department of Clinical Immunology, Aalborg University Hospital, Aalborg, Denmark

¹¹Gunma University Initiative for Advanced Research (GIAR), Gunma University, Showa-machi, Maebashi, Gunma, Japan

Correspondence

Naveed Akbar, Division of Cardiovascular Medicine, Radcliffe Department of Medicine, John Radcliffe Hospital, Level 6, West Wing, OX3 9DU, United Kingdom. Email: Naveed.Akbar@cardiov.ox.ac.uk

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Abstract

Plasma extracellular vesicle (EV) number and composition are altered following myocardial infarction (MI), but to properly understand the significance of these changes it is essential to appreciate how the different isolation methods affect EV characteristics, proteome and sphingolipidome. Here, we compared plasma EV isolated from platelet-poor plasma from four healthy donors and six MI patients at presentation and 1-month post-MI using ultracentrifugation (UC), polyethylene glycol precipitation, acoustic trapping, size-exclusion chromatography (SEC) and immunoaffinity capture. The isolated EV were evaluated by Nanoparticle Tracking Analysis (NTA), Western blot, transmission electron microscopy (TEM), an EV-protein array, untargeted proteomics (LC-MS/MS) and targeted sphingolipidomics (LC-MS/MS). The application of the five different plasma EV isolation methods in patients presenting with MI showed that the choice of plasma EV isolation method influenced the ability to distinguish elevations in plasma EV concentration following MI, enrichment of EV-cargo (EV-proteins and sphingolipidomics) and associations with the size of the infarct determined by cardiac magnetic resonance imaging

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²Department of Pharmacology, University of Oxford, Oxford, UK

6 months post-MI. Despite the selection bias imposed by each method, a core of EV-associated proteins and lipids was detectable using all approaches. However, this study highlights how each isolation method comes with its own idiosyncrasies and makes the comparison of data acquired by different techniques in clinical studies problematic.

KEYWORDS

acoustic trapping, human, immunoaffinity capture, omics, plasma, precipitation, size exclusion chromatography, ultracentrifugation

1 | INTRODUCTION

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Plasma extracellular vesicles (EV) are increased in number and carry altered protein, lipid and RNA cargo in the peripheral blood in many pathologies (Akbar et al., 2017, 2022; Burrello et al., 2020; de Miguel Pérez et al., 2020; Dong et al., 2016; Kalani et al., 2020). Analysis of plasma EV by omics approaches may provide unparalleled insight into multiple disease mechanisms and disease monitoring in patients for personalized medicine. However, it is unclear how different plasma EV isolation methods influence the plasma EV-profile or the so-called "EV-signature" in patients.

Current methods for the isolation of heterogeneous plasma EV include: ultracentrifugation (UC), density UC, field-flow fractionation, size-exclusion chromatography (SEC), precipitation, acoustic trapping (Evander et al., 2015) and immunoaffinity capture (Kowal et al., 2016). However, unsurprisingly, laborious protocols that yield pure EV from plasma are less well favoured in large cohorts (Nordin et al., 2015; Onódi et al., 2018) than protocols that are easier, lower cost, and more convenient. These methodological predilections are further impacted by the availability of stored biobank plasma, which often carry contaminating erythrocytes, immune cells, and platelets (Geyer et al., 2019). Irrespective of pre-storage processing, all plasma is a rich source of lipoproteins (apolipoprotein A and B), albumin, globulins and fibrinogens, which can co-isolate with plasma EV. The proportion of these cell-derived and non-cellular contaminants in the purified sample is method-dependent (Brennan et al., 2020) and they may obscure EV associated cargo (Théry et al., 2018; Van Deun et al., 2014).

Comparative isolation studies for plasma EV often assess EV size and concentration by Nanoparticle Tracking Analysis (NTA) and morphology by transmission electron microscopy (TEM). EV markers are usually evaluated using western blot or flow cytometry (Gidlöf et al., 2019; Menezes-Neto et al., 2015; Tian et al., 2020; Veerman et al., 2021). These standard analyses are often driven by the requirements of scientific bodies and consensus statements, which might be considered too prescriptive and refractory to change as our understanding of EV biology evolves (Théry et al., 2018). However, additional assessments of how isolation methods influence the plasma EV preparation have been explored with a range of techniques, including proteomics (Askeland et al., 2020; Cao et al., 2019; Gidlöf et al., 2019; Palviainen et al., 2020; Veerman et al., 2021), profiling of cytokines (Jung et al., 2020), RNA integrity (García et al., 2020; Stranska et al., 2018), lipidomics (Peterka et al., 2020), flow cytometry (Shtam et al., 2020; Tian et al., 2020) or a combination of these methods (Veerman et al., 2021). However, the wide range of plasma EV isolation and characterization techniques has evolved largely in the absence of systematic method characterization comparisons. As a consequence, the absence of an understanding of the impact of isolation techniques has led to uncertainty in relation to the interpretation of EV discoveries in clinical samples.

Here, we used platelet-free plasma from the same healthy volunteers to compare five different plasma EV isolation methods including: UC, precipitation, acoustic trapping, SEC and immunoaffinity capture using tetraspanins CD9, CD63, and CD81 (Figure 1). We then sought to determine how plasma EV isolation methods influence EV characteristics in a set of clinically well characterised individuals. Acute myocardial infarction (MI) is an important pathology; it is also an example of sterile inflammation where plasma EV number and composition are acutely altered (Akbar et al., 2017, 2022). Plasma EV from MI patients at two different time points were isolated using the five different EV isolation methods (Figure 1). Integrated unsupervised analysis of all the acquired characterization data from the five different methods was used to identify the similarities and differences between each method and to highlight how each technique might influence the protein and sphingolipid composition.

2 | MATERIALS AND METHODS

2.1 | Healthy volunteers and acute MI patients

All human investigations were conducted in accordance with the Declaration of Helsinki. The Oxfordshire Research Ethics Committee (Ref: 08/H0603/41 and 11/SC/0397) approved the human clinical protocols. All healthy volunteers and MI patients provided informed written consent for inclusion in the study.



FIGURE 1 A methodological overview. Platelet-poor plasma was obtained from healthy volunteers (n = 4) and patients presenting with myocardial infarction (MI) (n = 6) (and from the same patients 1-month post-MI) and plasma extracellular vesicles (EV) were isolated using five different methods: ultracentrifugation (UC), precipitation, acoustic trapping, size exclusion chromatography (SEC) and immunoaffinity capture with a matched vehicle phosphate buffered saline (PBS) or IgG control. The plasma EV were analysed using Nanoparticle Tracking Analysis, protein concentration, Western blot, transmission electron microscopy, a targeted EV-protein array for EV-markers CD9, CD63, CD81, ALIX, TSG101, flotillin, Annexin V, and 18 other cell associated markers, untargeted proteomics (LC-MS/MS) and targeted sphingolipidomics (LC-MS/MS). The data were analyzed in isolation and following integrated hierarchical clustering and principal component analysis. Room temperature (RT).

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2.2 | Generation of platelet poor plasma

Platelet-poor plasma was generated from healthy volunteers (n = 4), patients presenting with MI (n = 6) and from the same patients 1-month post-MI. 10–20 mL of whole blood was collected in EDTA (Greiner Bio-One, Stonehouse, United Kingdom) coated tubes and centrifuged at 1,000 × g for 25 min. The plasma was collected and centrifuged again for 10 min at 5,000 × g to produce platelet-poor plasma. The platelet-poor plasma was stored in 500 μ L aliquots at -80°C for future use.

2.3 | Isolation of plasma EV using UC

Plasma aliquots were thawed at room temperature and EV were isolated by UC by transferring 500 μ L of platelet-poor plasma from healthy volunteers or 100 μ L from platelet-poor plasma from MI patients to a 13.2 mL QuickSeal tube (Beckman Coulter, California, United States). The tubes were filled with 16 G hypodermic needles (Microlance, VWR, Pennsylvania, United States) fitted to a 10 mL syringe (VWR, Pennsylvania, United States) and after plasma was injected into the tube, 13 mL of phosphate buffered saline (PBS, Thermo Fischer Scientific, Massachusetts, United States) was added. The tubes were sealed using a soldering iron (Zacro, 60 W) and centrifuged using an Optima MAX-XP ultracentrifuge (Beckman Coulter, California, United States) at 120,000 × g for 120 min at 4°C with a MLA55 fixed-angle rotor (Beckman Coulter, California, United States) (Akbar et al., 2017, 2022). The pelleted plasma EV were resuspended in 100 μ L PBS or lysed in 100 μ L of RIPA buffer (Thermo Fischer Scientific, Massachusetts, United States) uses achusetts, United States) for subsequent analysis.

2.4 | Isolation of plasma EV using precipitation

Plasma EV were isolated by precipitation by using the Total Exosome Isolation Kit (Invitrogen, Massachusetts, United States). 500 μ L of platelet-poor plasma from healthy volunteers or 100 μ L of platelet-poor plasma from MI patients was thawed at room temperature. After vortexing, 20–100 μ L of the Exosome Precipitation Reagent (Total Exosome Isolation Kit (for plasma), ThermoFisher Scientific, Massachusetts, United States) was added and mixed by repeated pipetting. Samples were incubated for 30 min on ice, the mixture was centrifuged at 10,000 × g for 10 min by a Haraeus Fresco 17 benchtop centrifuge (ThermoFisher Scientific, ibid) at room temperature. The supernatant was removed and samples were centrifuged again at 2,000 × g for 2 min to remove residual supernatant. The pelleted plasma EV were resuspended in 100 μ L PBS or lysed in 100 μ L of RIPA buffer for downstream analysis by repeated pipetting.

2.5 | Isolation of plasma EV using acoustic trapping

Acoustic trapping of plasma for the isolation of EV was achieved by diluting plasma 1:1 with PBS, as previously described (Gidlöf et al., 2019). Briefly, samples were loaded onto a Costar 96-well plate (Corning, New York, United States) and inserted in the AcouSort device (Version 2.0) (AcouSort AB, Lund, Sweden). Acoustic waves were produced by a waveform generator (Keysight 33210A, Keysight, California, United States) with a 9.2 V output and were directed into a borosilicate capillary acoustic trapping unit (AcouSort AB, Lund, Sweden). Following activation of the waveform generator and the acoustic trapping unit was initialised, 50 μ L of 12 μ m polystyrene beads (AcouSort AB, Lund, Sweden) were loaded into the acoustic trapping unit using a syringe pump (Tricontinent C2400 (Tricontinent, Fürstenfeldbruck, Germany) set to 50 μ L/min. Each run consisted of 100 μ L of 1:1 diluted plasma and were loaded with a syringe pump speed of 20 μ L/min. After acoustic trapping, the samples were washed by aspirating 15 μ L of PBS at 20 μ L/min and dispensing 50 μ L at 20 μ L/min. After the PBS wash, the samples were eluted in 50 μ L of PBS and used for downstream analysis.

2.6 | Isolation of plasma EV using size exclusion chromatography (SEC)

Size exclusion chromatography (SEC) isolation of plasma EV was achieved by using the Exo-spinTM 96 (Cell Guidance Systems, Missouri, United States) (Welton et al., 2015). Columns were equilibrated at room temperature for 15 min prior use. Afterwards, the columns were washed twice with 250 μ L of PBS. Plasma was thawed at room temperature and 100 μ L of plasma from healthy volunteers or MI patients was loaded into each column. Plasma EV were eluted by adding 200 μ L of PBS to the top of the column eluted under gravity. SEC derived plasma EV were isolated by collecting the third SEC fraction (Figure S1). Plasma EV were collected and stored for subsequent analysis.



2.7 | Isolation of plasma EV using immunoaffinity capture

Prior to isolation, the plasma was passed through the SEC method as described above. Exosome-Human CD9 Isolation Reagent, Exosome-Human CD63 Isolation/Detection Reagent and Exosome-Human CD81 Isolation Reagent (all Invitrogen, Massachusetts, United States) were used to capture plasma EV. The mixture was created by resuspending each bead solution and mixing by repeated pipetting. Control IgG-isotype (10400C, Thermo Fischer Scientific, Massachusetts, United States) matched beads were conjugated according to the instructions of the Dynabeads Antibody Coupling Kit (Invitrogen, Massachusetts, United States). After SEC pre-isolation, the samples were incubated with 80 μ L of the combined CD9, CD63 and CD81 beads (end concentration 1 × 10⁷/mL, equal quantity of each bead mixture) or equally concentrated IgG control beads at 4°C for 18 hours under continual rotation by a vertical rotor (Grant Bio, Essex, United Kingdom). After incubation, the beads were pelleted for 5 min at room temperature using a Dynal Magnet (Invitrogen, Massachusetts, United States) and the supernatant was collected for subsequent analysis. Following pelleting, the samples were washed three times with 200 μ L of PBS and magnetic beads were pelleted as described above. Following the washes, EV were lysed in 100 μ L of RIPA buffer and supernatant was separated by using a 2,000 × g centrifugation and collected for subsequent analysis.

2.8 | Nanoparticle Tracking Analysis (NTA)

Plasma EV particle size distribution and concentration were determined by NTA using a Zetaview device (Particle Metrix, Inning am Ammersee, Germany) as previously described (Akbar et al., 2022). Prior to injection into the sample chamber, samples were diluted in PBS. The Zetaview measured the sample chamber from 11 positions in two cycles. The settings were set at sensitivity 80, frame 30 and shutter speed 100. Silica 100 nm microspheres (Polysciences Inc., Philadelphia, United States) were used to quality check the instrument performance routinely. The particle concentration per method was calculated as the change (Δ) compared to the control sample, in which PBS replaced the plasma sample. For immunoaffinity plasma EV, the particle concentration was calculated by measuring the change (Δ) in the initial concentration and unbound fraction.

2.9 | Protein concentration

Protein concentration was determined by bicinchoninic assay (BCA) (Thermo Fischer Scientific, Massachusetts, United States). A standard curve with Bovine Serum Albumin (Thermo Fischer Scientific, Massachusetts, United States) was used to calculate the protein concentration. Isolated EV were lysed in RIPA at a ratio of 1:2 or 1:6 and needle sonicated by a SonoPuls HD2070 (Bandelin, Berlin, Germany) at 40% power for 10 seconds. 25 μ L of sonicated sample or standard was incubated in duplicate with 175 μ L of a 25:1 ratio between Reagent A and Reagent B (Thermo Fischer Scientific, Massachusetts, United States) and incubated at 37°C for 30 min. After incubation, absorbance was measured at 562 nm using a plate reader (FLUOstar Omega plate reader, BMG Labtech, Aylesbury, United Kingdom).

2.10 | Transmission electron microscopy (TEM)

TEM of the isolated EV was conducted as previously described (Akbar et al., 2022). Briefly, grids (300 mesh Cu carbon film) were glow discharged for 20 seconds at 15 mA (Leica EM ACE 200). The isolated plasma EV samples were added to the grid for 2 min, blotted, stained with 2% uranyl acetate for 20 seconds, blotted and allowed to air dry. Images were acquired on a 120 kV Tecnai 12 TEM (Thermo Fischer Scientific, Massachusetts, United States) equipped with a OneView digital camera (Gatan, California, United States). TEM images of control samples, in which the isolation method was run with PBS in the place of the plasma, were obtained for each method. Immunoaffinity-based isolated EV were fixed with 1.6% glutaraldehyde in PBS for 1 hour at 4°C. Following fixation, the beads were fixed in 4% agarose with PBS. The agarose was cut into small cubes (1–2 mm³). Sections were collected onto 200 mesh Cu grids and imaged using a Gatan OneView camera with a FEI Tecnai 12 TEM at 120 kV.

2.11 | Proteomics

Isolated plasma EV were processed for proteomics as previously described (Thompson et al., 2020). In short, the samples were reduced in 5 mM dithiothreitol for 30 min at room temperature. Subsequently, the samples were alkylated with 20 mM iodacetamide for 30 min and precipitated using chloroform-methanol precipitation. Quantified protein groups with \geq 2 unique peptides were included in the comparison of proteomes. The LFQ values for each protein group were deducted by the LFQ from



control samples per isolation method. Protein abundance was normalised by log-transformation. Gene Ontology (GO) analysis of the protein groups was conducted by using Genontology.org (Mi et al., 2019) and the data were extracted on 19-07-2021. *p*-Values were corrected for false discovery rate (FDR) and significance was set at p < 0.05. Fisher exact tests were conducted using R on published databases EVpedia (Kim et al., 2013), Vesiclepedia (Kalra et al., 2012), and Exocarta (Simpson et al., 2012).

2.12 | EV-Array

Isolated plasma EV were analysed by a targeted EV-Array, which has been described previously (Akbar et al., 2022; Bæk & Jørgensen, 2017; Jørgensen et al., 2013). In short, a protein microarray plate was generated with the following antibodies: CD146 (PIH12), Flotillin-1, TSG101 (Abnova, Taiwan), CD9, CD81 (Ancell corporation, Minnesota, United States); CD16 (3G8, BD Biosciences, California, United States); ALIX (3A9), VEGFR2 (7D4-6; Biolegend, California, United States); CD63 (Bio-Rad, California, United States); ICAM-1 (R6.5, eBioscience, California, United States); Endoglin (LSbio, Washington, United States); Tissue factor (323,514), VCAM-1 (HAE-2Z), Thrombomodulin (501733), CD31 (AF806, R&D Systems, Minnesota, United States), VE-Cadherin (AF938, R&D Systems, Minnesota, United States). After incubation with the blocking buffer (50 mM ethanolamine, 100 mM Tris, 0.1% SDS, pH 9.0) for 30 min, the wells were emptied, and the plate was dried for 5 hour and sealed. The samples were incubated in the antibody coated microarray plate overnight at 4°C. Following a wash, each well was incubated with a 100 μ L of a detection antibody cocktail (biotinylated anti-human-CD9, -CD63 and -CD81 (Ancell, Minnesota, United States). After another wash, 100 μ L streptavidin-Cy3 Life Technologies, Massachusetts, United States) diluted 1:3,000 was added to each well and incubated for 30 min. The plate was scanned using a sciREADER FL2 microarray scanner (Scienion AG, Berlin, Germany), at 535 nm and an exposure time of 2,000 msec. For each protein the control, PBS sample value was subtracted from the result.

2.13 | Western blot

Isolated plasma EV or controls were lysed using RIPA buffer with protease and phosphatase inhibitors PhosSTOP (Roche, Basel, Switzerland) and cOmplete (Roche, Basel, Switzerland) and were needle sonicated by a SonoPuls HD2070 (Bandelin, Berlin, Germany) at 40% power for 10 seconds as previously described (Akbar et al., 2017, 2022). Following sonication, samples were incubated at 95°C for 5 min. Eight microgram of protein was combined with NuPage LDS sample buffer (4×) agent (Invitrogen, Massachusetts, United States). The samples were loaded onto a 4%-12% Bis-Tris gradient gel (NuPAGE 4%-12% Bis-Tris Protein Gel; 1.5 mm (ThermoFisher Scientific, Massachusetts, United States) with Amersham ECL Full Range ladder (Cytiva Life Sciences, Massachusetts, United States). Separated samples were transferred to a nitrocellulose membrane (Amersham Proton 0.2 μ m, GE Healthcare, Illinois, United States) and blocked for non-specific binding in 5% skimmed milk powder (Marvel Original, New York, United States) in 0.5% PBS-Tween 20 (Sigma-Aldrich, Missouri, United States) for 1 hour. Membranes were incubated with primary antibodies overnight: ALIX (ab117600, Abcam, Cambridge, United Kingdom) (1/1,000 dilution), CD63 (EXOAB-KIT-1, System Biosciences, California, United States) (1/1,000 dilution), ApoB (Ab139401, Abcam, Cambridge, United Kingdom) (1/20,000 dilution), albumin (MAB1455, R&D Systems, Minneapolis, Canada) (1/8,000 dilution), ApoA-I (Mab36641, R&D systems, Minneapolis, Canada) (1/8,000 dilution) and H3 (D1H2, Cell Signalling Technology, Massachusetts, United States) (1/1,000 dilution) 5% milk in PBS-Tween (PBS-T). Membranes were washed three times with PBS-T and incubated with secondary-horse radish peroxidase (HRP) conjugated antibodies (1/20,000 α -mouse W402B or 1/50,000 α -rabbit W401B, Promega, Wisconsin, United States) for 1 hour. The membranes were triple washed once again with PBS-T before incubating them with enhance chemiluminescence substrate (Pierce ECL, ThermoFisher Scientific, Massachusetts, United States) for imaging (Bio-Rad ChemiDoc MP Imaging system, California, United States).

2.14 | Sphingolipidomics

Sphingolipids were determined as previously described (Akawi et al., 2021). 25 μ L of each plasma EV sample were combined with 10 μ L solution containing labelled sphingolipid internal standards in LC-MS Methanol (Honeywell, North Carolina, United States). This was followed by the addition of 100 μ L of methanol to each sample. Samples were then vortexed for 30 s and sonicated for 15 min in a Fisherbrand Ultrasound bath S60 (Fischer-Scientific, Massachusetts, United States) with ice. Next, samples were centrifuged at 12,000 × g during 15 min at 6°C. Finally, 80 μ L of the supernatant were transferred to an LC-MS amber vial (Waters, Wilmslow, United Kingdom) equipped with a 150 μ L insert. Samples were randomized by time point of the individual within an extraction method. Samples were analyzed on an Acquity UPLC coupled to a Xevo TQ-S Mass Spectrometer (Waters, Wilmslow, United Kingdom) as previously described (Akawi et al., 2021).



2.15 | Bioinformatics and integrated analysis

Integrated analysis was conducted by combining data from the NTA, protein concentration, EV-Array, proteomic and sphingolipidomic analyses. The data was normalized per row and log transformed. Following data normalization, a principal component analysis was conducted in R 4.0.0 (R Core Team, 2021) using the *factoextra* (Kassambara & Mundt, 2020) and *FactoMineR* (Lê et al., 2008) packages. For visualization *ggplot2* (Wickham, 2016) was used. Each of the principal components were then used to create a heatmap by extracting the values from the eigenvalues for each of the principal components and processing them in *pheatmap* (Kolde, 2018). The data was clustered by using double hierarchical clustering using the *pheatmap* R package.

2.16 | Data availability

All z-scored normalised healthy volunteer data acquired in this study are accessible through: https://ora.ox.ac.uk/objects/ uuid:131e61e2-9349-45ce-8db3-377ccfd7a18e. All other data produced in the present study, including those from patients experiencing MI are available upon reasonable request to the Corresponding Author.

2.17 | Statistical analysis

Data were plotted as mean with standard deviation. Normality of the data was confirmed using QQ plot and D'Agostino-Pearson normality test. For paired analysis, at the two-time points, or for two independent groups, paired or unpaired Students *t*-test were used, respectively (GraphPad Prism 9). Correlation analysis was carried out using linear Pearson regression analysis (Graphpad Prism 9). One-way ANOVA with Bonferroni correction post-hoc tests were used for analyses with >2 independent groups. The proteomic and lipidomic data was analyzed using a Kruskal-Wallis statistical test with Bonferroni post-hoc tests. *p*-Values < 0.05 were considered significant.

3 | RESULTS

3.1 | Plasma EV number and size is influenced by the isolation method

The concentration of isolated particles (expressed as delta over a matched PBS or and IgG control sample) determined by NTA differed per method (UC $9.5 \times 10^9 \pm 1.8 \times 10^9$ particles/mL, precipitation $6.1 \times 10^{11} \pm 2.7 \times 10^{11}$ particles/mL, acoustic trapping $6.4 \times 10^9 \pm 2.8 \times 10^9$ particles/mL, SEC $5.5 \times 10^{10} \pm 1.9 \times 10^9$ particles/mL and immunoaffinity $2.8 \times 10^{10} \pm 7.1 \times 10^9$ particles/mL, Figure 2a). In agreement with previously published studies, precipitation yielded significantly higher numbers of particles/mL compared to UC (Serrano-Pertierra et al., 2019; Tian et al., 2020) and SEC (Gámez-Valero et al., 2016; Tian et al., 2020) (both p < 0.01). Particle number, isolated by precipitation, were also higher than the concentration acquired by acoustic trapping or by immunoaffinity capture (both p < 0.01) (Figure 2a). The size and concentration distribution for each biological replicate exhibited uniformity within each isolation method and the different isolation methods gave a similar size distribution profile overall, which ranged from 15 nm (Figure 2b). However, the mean size of isolated particles by UC was significantly higher compared to the other methods (UC 143.7 \pm 3.4 nm, vs. precipitation 94.2 \pm 3.9 nm, acoustic trapping 81.2 ± 3.9 nm and SEC 87.2 ± 1.5 nm, p < 0.01 all) (Table 1).

TABLE 1 Nanoparticle Tracking Analysis showing the mean and median size of plasma-EV isolated by ultracentrifugation (UC), precipitation, acoustic trapping and size-exclusion chromatography (SEC)

	Mean size (nm)	Median size (nm)
*Ultracentrifugation	143.7 ± 3.4	125.0 ± 2.9
Precipitation	94.2 ± 3.9	84.0 ± 3.7
Acoustic trapping	81.2 ± 3.9	70.7 ± 4.6
SEC	87.2 ± 1.5	75.0 ± 1.4
Immunoaffinity	-	-

Note: Immunoaffinity capture cannot be acquired. Data are group averages \pm standard deviation (SD) and were analyzed by one-way ANOVA with post-hoc Bonferroni correction. *p < 0.001 (n = 4).



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FIGURE 2 Plasma EV characterization using different isolation methods. (a) Total plasma extracellular vesicles (EV) as particles/mL concentrations and (b) size and concentration distribution profiles were obtained by Nanoparticle Tracking Analysis (NTA) using ultracentrifugation (UC), precipitation, acoustic trapping, size exclusion chromatography (SEC) and immunoaffinity capture (n = 4). Values are presented as a delta compared to a vehicle control or IgG control. Scale is logarithmic. (c) Protein concentration of plasma EV using UC, precipitation, acoustic trapping, SEC and immunoaffinity capture (n = 4). Values are presented as a delta compared to a vehicle control or IgG control. Scale is logarithmic. (c) Protein concentration of plasma EV using UC, precipitation, acoustic trapping, SEC and immunoaffinity capture (n = 4). Values are presented as a delta compared to a vehicle control or IgG control. (d) Western blot of plasma EV isolated by UC, precipitation, acoustic trapping, SEC and immunoaffinity capture versus controls using EV markers ALIX and CD63, lipoprotein contaminants apolipoprotein B (ApoB), and apolipoprotein A-I (ApoA-I), plasma contaminant albumin and cellular contaminant histone H3. Endothelial cell EV and peripheral blood mononuclear cells (PBMCs) were used for H3 positive controls. (e–j). Transmission electron microscopy (TEM) images of isolated plasma EV from UC, precipitation, acoustic trapping, SEC and immunoaffinity capture versus controls. Each sub panel contains a zoomed-in image (left image), an overview image (top right) and a control vehicle image (bottom right). For the immunoaffinity bead capture images, the red arrows indicate EV particles. The scale bar is 200 nm for the zoomed images surrounded by a dashed line and 1,000 nm for the overview images Data are group average \pm standard deviation (SD). Data was analyzed by one-way ANOVA with post-hoc Bonferroni correction. ***p < 0.001.



3.2 | Plasma EV protein concentration

The protein concentration (expressed as the change over a matched PBS or and IgG control) also differed across the EV isolation methods: UC 982 \pm 113 µg/mL, precipitation 9,478 \pm 3,174 µg/mL, acoustic trapping 1,211 \pm 141 µg/mL, SEC 382 \pm 51 µg/mL and immunoaffinity 33 \pm 12 µg/mL (Figure 2c). EV generated by precipitation had a significantly higher protein concentration compared to UC, acoustic trapping, SEC or immunoaffinity (all p < 0.01, Figure 2c). The purity of EV isolation can be estimated by calculating a ratio of EV number (EV/mL) to protein concentration (µg/mL) (Webber et al., 2013). The EV purity ratio differed between isolation method: UC 9.0 × 10⁶ \pm 1.4 × 10⁶; precipitation 4.7 × 10⁸ \pm 1.5 × 10⁸; acoustic trapping 2.1 × 10⁷ \pm 7.6 × 10⁶; SEC 6.8 × 10⁶ \pm 4.0 × 10⁶ and immunoaffinity capture 3.1 × 10⁸ \pm 1.0 × 10⁸. Precipitation achieved a significantly higher EV purity ratio when compared to UC, SEC, acoustic trapping and immunoaffinity (p < 0.001 compared to all methods) (Figure S2).

3.3 | Plasma EV protein characterization by Western blot

As NTA is unable to distinguish between plasma EV and similarly sized protein aggregates and lipoproteins, we analysed the isolated plasma EV obtained from each method for a number of markers by Western blotting. ALIX and CD63 were included as the EV markers. For the plasma lipoprotein contaminants, we measured ApoB and ApoA-I. Histone H3 for cellular contaminants and albumin was also included in the set. Each isolation method showed the presence of EV markers ALIX and CD63 (Figure 2d). Apolipoproteins (ApoA-I and ApoB) were also present in all isolation methods. All methods were negative for markers of cellular contamination by histone H3. Surprisingly, the IgG isotype control for immunoaffinity capture using CD9, CD63, and CD81 also showed the presence of EV markers ALIX, CD63, ApoB, albumin and ApoA-I, but histone H3 was absent (Figure 2d).

3.4 | Plasma EV morphology by TEM

To determine the morphology of isolated plasma EV, we undertook TEM for the five different plasma EV isolation methods versus a matched PBS control or an IgG control for immunoaffinity capture beads. TEM analysis showed EV-like particles for each isolation method and an absence of EV-like particles in their respective PBS isolation controls (Figure 2e-h). Immunoaffinity beads CD9, CD63 and CD81 were embedded in agar and sectioned to visualise the bead surface for EV-like structures versus the IgG control. IgG control showed no EV-like particles present (Figure 2i). Immunoaffinity capture using CD9, CD63 and CD81 beads showed intact EV-like particles captured on the bead surface (Figure 2j).

3.5 | Plasma EV compositional analysis using a high throughput protein EV-Array

The results detailed above provide evidence for the relative success of each method to yield plasma EV and assessment of relative plasma EV purity (Théry et al., 2018). However, these techniques do not readily distinguish the abundance of specific plasma EV populations carrying, for instance, specific cell-associated markers, or easily allow quantitative assessment of the abundance of contaminating lipoproteins. Thus, we probed the composition of the plasma EV isolated from each method for general EV markers CD9, CD63, CD81, ALIX, TSG101, Flotillin 1, Annexin V, and a panel of 18 cell associated markers, which may distinguish EV from platelets, endothelial cells, immune cells and muscle; simultaneously, lipoprotein contaminants apolipoprotein E (ApoE) and apolipoprotein H (ApoH) were a validated high throughput EV-protein antibody array (Figure 3) (Jørgensen et al., 2013). We utilised a matched PBS control for UC, precipitation, acoustic trapping and SEC and an IgG control for immunoaffinity capture. For the EV-associated proteins, CD9 and CD81, they were significantly higher in plasma EV isolated by UC compared to the other methods (p < 0.01 for both). Annexin V was significantly higher in the acoustic trapping samples compared to SEC and immunoaffinity capture (p < 0.01 and p < 0.05, respectively).

Hierarchical clustering of the EV-Array acquired data for the different isolation methods indicates that there are significant method-dependent differences for cell associated EV markers (Figure 3). CD31 was significantly higher in precipitation isolated plasma EV versus acoustic trapping (p < 0.05). CD41 was significantly higher in UC isolated plasma EV compared to precipitation isolated EV (p < 0.05) and CD16 content was significantly higher in the precipitation isolated plasma EV samples compared to SEC (p < 0.01). There were no differences between immunoaffinity capture using CD9, CD63, and CD81 and IgG controls (Figure 3).

3.6 | Unbiased proteomic analysis of plasma EV

We next determined the proteomic profile of each isolation method using unbiased LC-MS/MS versus their respective PBS or IgG controls for each method. The proteomic profile following plasma EV isolation showed a significantly higher number of

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FIGURE 3 Heatmap of plasma EV derived from different isolation methods using the EV-protein-Array. The heatmap contains extracellular vesicles (EV) markers CD9, CD81, CD63, ALIX, TSG101, Flotillin 1 and Annexin V for ultracentrifugation (UC), precipitation, acoustic trapping, size exclusion chromatography (SEC) and immunoaffinity capture, lipid contaminants apolipoprotein H (ApoH) and apolipoprotein E (ApoE) and cell associated markers. Values are derived from 500 μ L of platelet poor plasma per sample, per technique. Values are presented as a delta compared to a vehicle control or IgG control and are log normalized (*n* = 4 per isolation method). Data was analyzed by Kruskal-Wallis test with post-hoc Bonferroni correction. **p* < 0.05, ****p* < 0.001.

quantified protein groups compared to their respective controls for all isolation methods, except immunoaffinity capture, which displayed similar results to the IgG control beads (UC p < 0.01, precipitation p < 0.05, acoustic trapping p < 0.01 and SEC p < 0.01, Figure 4a). The choice of plasma EV isolation method influenced the overall EV-proteome, but nine protein groups were common across all methods (Figure S3). These were: immunoglobulin heavy constant gamma 1, alpha-2-macroglobulin, haptoglobin, immunoglobulin heavy constant μ , immunoglobulin kappa constant, serpin family A member 1, albumin, fibrinogen alpha chain and Apo-A1.

We used hierarchical clustering of the top 30 quantified protein groups across the different plasma EV isolation methods and found a distinct separation between isolation methods. UC and SEC isolated plasma EV clustered together and precipitation, acoustic trapping and immunoaffinity capture formed a distinct separate cluster (Figure 4b). Hierarchical clustering indicates that these differences between UC/SEC and precipitation, acoustic trapping and immunoaffinity capture by an abundance of protein groups with common plasma proteins (ALB, A2M, ApoB, C3, TF, HP, Apo-A1, SERPINA1, CP and ITIH2), fibrinogens (FGG and FGB) and immunoglobulins (IGHG1, IGKC, IGHG3, IGHM, IGK, IGHG2).

To better understand the nature of the proteomic profile for each isolation method, we conducted unbiased GO pathway analysis of the protein groups associated with each isolation method. All methods showed a significant association with EV pathways: Blood Microparticle GO: 0072562 and Extracellular Exosome GO:0070062 (all p < 0.01, Figure S4A). However, there

FIGURE 4 Proteomic comparison of plasma EV isolation methods. (a) The number of protein groups quantified by unbiased proteomics for plasma extracellular vesicles (EV) derived by: ultracentrifugation (UC), precipitation, acoustic trapping, size exclusion chromatography (SEC) and immunoaffinity capture, versus control vehicle (PBS) or an IgG control. Protein groups were only included as quantified if they had ≥ 2 unique peptides (n = 3-4). One-way ANOVA with post-hoc Bonferroni correction. Data are group averages \pm standard deviation (SD). **p* < 0.05, ***p < 0.001. (b) A heatmap of the top 30 quantified protein groups across all isolation methods. Values from control samples were subtracted to account for background and the values were log normalised. Hierarchical clustering of the isolation methods was conducted using a complete clustering method (n = 3-4).



was no clear separation between the different isolation methods using this pathway analysis approach. To further scrutinize the EV proteomic profile obtained per plasma EV isolation method we undertook a statistical comparison using a Fisher's exact test with published EV-databases: EVpedia (Kim et al., 2013), Vesiclepedia (Kalra et al., 2012), and Exocarta (Simpson et al., 2012). This determined the similarity between plasma EV proteomic profiles obtained from the five different isolations methods to those published previously by showing the size of the intersect. There was a significant overlap between the five different isolation methods and the archived EV databases (p < 0.05, all methods, Figure S4B). UC, SEC and immunoaffinity capture showed the greatest similarity with published databases. Whereas precipitation and acoustic trapping showed less similarity. Furthermore,





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FIGURE 5 Sphingolipidomic analysis of plasma EV from different isolation methods. (a) Number of sphingolipids quantified in plasma extracellular vesicles (EV) isolated by ultracentrifugation (UC), precipitation, acoustic trapping, size-exclusion chromatography (SEC) and immunoaffinity capture and subjected to targeted sphingolipid analysis versus control vehicle (PBS) or an IgG control. Data are group averages \pm standard deviation (SD) and were analysed by One-way ANOVA with post-hoc Bonferroni correction. ***p < 0.001 (n = 4). (b) Heat map of plasma EV sphingolipids. Values from control samples were subtracted to account for background and the values were log normalised. Hierarchical clustering of the isolation methods was conducted using a complete clustering method (n = 3-4). Data were analysed by Kruskal-Wallis test with

we determined whether our plasma EV acquired proteomic profiles from the five different methods were similar to previously published plasma EV proteomic data for Exospin, SEC, ExoQuick, IZON35, IZON70, Optiprep and Exo-easy (Menezes-Neto et al., 2015; Veerman et al., 2021) and found a significant overlap for all five plasma EV isolation methods (p < 0.01) (Figure S5).

3.7 Targeted sphingolipidomic of plasma EV

EV membranes are largely composed of lipids (Skotland et al., 2020) and lipoproteins are a predominant contaminant in plasma EV samples (Karimi et al., 2018), which are influenced by the choice of isolation method (Brennan et al., 2020; Yuana et al., 2014). We compared the lipidomic profile of the different plasma EV isolations methods by undertaking targeted sphingolipidomic analysis. The sphingolipidomic analysis showed a significantly higher number of sphingolipids compared to the respective controls for all isolation methods except immunoaffinity capture (UC p < 0.01, precipitation p < 0.05, acoustic trapping p < 0.01and SEC p < 0.01, Figure 5a). However, there were distinct sphingolipidomic differences between the other methods. Eleven of the quantified sphingolipids were common to all methods, which were DhCer(d18:0/24:0), Cer(d18:1/22:0), Cer(d18:1/24:1),



Cer(d18:1/24:0), SM(d18:1/16:0), SM(d18:1/18:0), SM(d18:1/24:1), SM(d18:1/24:0), HexCer(d18:1/16:0), HexCer(d18:1/24:1) and Lac-Cer(d18:1/24:1) (Figure S6). Hierarchical clustering analysis of the sphingolipidomic profile for each of the isolation methods showed that there are three distinct clusters (Figure 5b). Immunoaffinity capture and precipitation formed two separate individual clusters, whilst UC, acoustic trapping and SEC clustered together. Hierarchical clustering indicates that these group differences were driven by an abundance of sphingomyelins (16:0, 18:0, 24:0 and 24:1), ceramides (22:0, 24:0 and 24:1), hexosylceramides (24:1) and lactosylceramides (16:0) in the precipitation group and a lack of these sphingolipids in the immunoaffinity capture group.

3.8 | Integrated comparison of plasma EV isolation methods

These data shows that plasma EV isolation methods have divergent impact on the yield of EV, their purity, the type of EV isolated from plasma that are associated with a particular cell source and the overall proteomic and sphingolipidomic profile. To better understand how these individual method associated differences influenced the plasma profile, we undertook integrated analysis of all the different plasma EV isolation methods with all of the acquired data. This included the EV particle concentration, protein concentration, EV-protein-Array data, proteomic and sphingolipidomic data. To condense the multiple different variables, we converted the data into principal components. Principal component analysis with two components (PC1 and PC2) accounted for 47.2% of the variance and showed clear separation between the different isolation methods (Figure 6a,b). UC, SEC and precipitation form distinct clusters, whereas acoustic trapping and immunoaffinity capture clustered together. Specific principal component analysis indicated that PC1 is driven by proteomics acquired data and PC2 is driven by sphingolipidomic data (Figure \$7). These integrated data show that the EV isolation method influences the omic and integrated-based plasma EV-profile, which may influence interpretation of EV-acquired data for clinical biomarker discovery and precision diagnostics.

3.9 | Plasma EV isolation methods influence the diagnostic potential of plasma EV from patients following MI

We next determined whether the choice of plasma EV isolation method impacts the ability to detect changes in EV-profile in a disease state. We obtained plasma at time of presentation with MI, but prior to percutaneous coronary intervention (PCI), and a matched control plasma sample was obtained at 1-month post-MI from the same patients. The clinical patient characteristics are detailed in Table 2.

TABLE 2 Clinical characteristics of the MI patients

	Average \pm SD
Age	67.0 ± 11.4
Sex (male/female)	6/0
Glucose (mmol/L)	$5.7 \pm 0.5 \ (n = 3)$
White blood cell count ($\times 10^9$)	7.6 ± 2.2
Troponin (peak ng/L) at presentation	358.7 ± 401.2
Cholesterol (mmol/L)	4.8 ± 1.2
Diabetic status (diabetes/non)	0/6
Smoker status (smoker/non)	1/5
Infarct size at 6-months	15 ± 7.8
LVEF at follow-up (%)	47 ± 5.0

Note: Age, sex (M/F), glucose, white blood cells counts, troponin peak, cholesterol, diabetes status, smoker status, infarct size determined by late gadolinium enhancement MRI 6 months post-AMI and left ventricle ejection fraction (LVEF %) 6 months post-MI. Data are group averages \pm standard deviation (SD) (n = 6).

3.10 | Plasma EV concentration in MI is influenced by the isolation method

To mitigate any potential variability in bio-banked plasma samples, we used three technical plasma replicates per patient, per time-point and per method. There were significantly more plasma EV at time of presentation with MI versus the 1 month control follow-up when plasma EV were isolated by UC, precipitation and acoustic trapping (p < 0.05 for all) (Figure 7a), but not by SEC and immunoaffinity capture (UC presentation: 2.8×10^9 particles/mL vs. UC follow-up: 2.1×10^9 particles/mL; precipitation



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FIGURE 6 Principal component analysis of plasma EV characteristics following data integration. (a) A principal component (PC) analysis of plasma extracellular vesicles (EV) isolated by ultracentrifugation (UC), precipitation, acoustic trapping, size-exclusion chromatography (SEC) and immunoaffinity capture including: plasma EV concentration, protein concentration, EV-protein-Array, proteomics and sphingolipidomics. The integrated data was condensed to PC1 and PC2. (b) A heatmap with the various principal components to compare the different isolation plasma EV isolation methods. Hierarchical clustering of the isolation methods was conducted using a complete clustering method (n = 3-4).

presentation: 1.5×10^{12} particles/mL vs. precipitation follow-up: 7.9×10^{11} particles/mL; acoustic trapping presentation: 4.3×10^{10} particles/mL vs. acoustic trapping follow-up: 1.9×10^{10} particles/mL; SEC presentation: 2.5×10^{11} particles/mL vs. SEC follow up: 9.2×10^{10} particles/mL and immunoaffinity presentation: 2.0×10^{11} particles/mL vs. immunoaffinity capture follow-up: 9.8×10^{10} particles/mL, Figure 7a). The technical variance between the three independent isolations from each patient, and at each time, point showed that SEC gave significantly less variance compared to UC (10.0 ± 3.1 % vs. 22.7 ± 13.0 %, p < 0.01) (Figure S8A). The mean plasma particle size was similar between the time of presentation and 1 month follow-up control for the five different plasma EV isolation methods (Figure S8B) and the size and concentration distribution profile showed no distinct differences between time-points or dependent on the plasma EV isolation method (Figure S8C).



FIGURE 7 Plasma EV analysis using different isolation methods in patients following presentation with myocardial (MI) infarction compared to samples from the same patients after a 1-month follow-up. (a) Comparison of total concentration of plasma extracellular vesicles by Nanoparticle Tracking Analysis (EV) from patients following presentation with MI and after a 1-month follow-up using: ultracentrifugation (UC), precipitation, acoustic trapping, size exclusion chromatography (SEC) and immunoaffinity capture (n = 6 per timepoint). Data are group average \pm standard deviation (SD). Paired *t*-test analysis *p < 0.05. (b) A Pearson correlation analysis of plasma EV as particles/mL derived from each isolation method at time of presentation versus the infarct size determined by cardiac MRI using late gadolinium enhancement 6-months post-infarct (n = 5). (c) Heatmap of plasma EV-Array analysis. The top section of the heatmap contains the different EV markers CD9, CD81, CD63, ALIX, TSG101, Flotillin 1 and Annexin V. Values are presented as fold over the respective matched follow-up control sample. (d) Heatmap showing the plasma EV sphingolipidomic profiles in MI patents at presentation versus a 1-month follow-up for UC, precipitation, acoustic trapping, SEC and immunoaffinity capture. Values are presented as fold over the respective matched follow-up.

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3.11 | The plasma EV isolation method influences plasma EV concentration association with infarct size

We have previously reported that the total concentration of plasma EV in the peripheral blood isolated by UC at the time of presentation correlates with the size of myocardial injury and scar, determined by late gadolinium enhanced (LGE) MRI 6 months post-MI (Akbar et al., 2017, 2022). Therefore, we determined whether plasma EV concentrations from the five different plasma EV isolation methods influenced the ability to determine this important clinical association. Infarct size at 6-months post-MI significantly correlated with the plasma EV particles concentration at time of presentation for UC samples ($R^2 = 0.89$, p = 0.02), but not for precipitation, acoustic trapping, SEC or immunoaffinity capture ($R^2 = 0.10$ p = 0.60, $R^2 = 0.16$ p = 0.50, $R^2 = 0.05$ p = 0.71 and $R^2 = 0.10$, p = 0.78, respectively) (Figure 7b).

3.12 | Cell associated plasma EV are not influenced by the isolation method in MI

We determined whether the isolation methods influenced cell associated plasma EV-markers in MI patients at presentation versus the respective 1 month follow-up sample. The high throughput EV-Array data was expressed as fold change over the matched follow-up control samples per patient. Plasma EV markers were increased at time of presentation vs. follow-up control (Figure 7c). However, the pattern of change in plasma EV-markers were not consistent across the different isolation methods (Figure 7c). Hierarchical clustering of the plasma EV marker response at the time of presentation with MI showed method-dependent clustering. Similarly, cell associated markers on plasma EV were differentially enriched following MI (Figure 7c). Once again, unbiased clustering showed method-dependent groupings. Together these data suggest that changes in plasma-EV protein are influenced by the choice of plasma EV isolation method following MI in isolation and when clustered together.

3.13 \parallel Plasma EV sphingolipidomic profiles are altered following MI and influenced by the isolation method

Next, we profiled EV-sphingolipids in the acute phase following MI (calculated as fold over matched follow-up control samples) using the different plasma EV isolation methods. Plasma EV-sphingolipidomic profiles for samples generated by precipitation, acoustic trapping and SEC contained significantly more quantified lipid groups at time of presentation with MI versus samples obtained by UC and immunoaffinity capture (p < 0.01, all). We ranked plasma EV-lipid profiles based on their overall abundance of lipid groups and found that ceramides were significantly higher in the precipitation isolated plasma EV samples, followed by SEC, acoustic trapping and UC/immunoaffinity capture (p < 0.01). Whereas the sphingomyelins were significantly higher in the precipitation group versus all other isolation methods (p < 0.001) (Figure 7d). The clustered plasma EV-sphingolipid profile following MI was influenced by the choice of isolation method. Several ceramides and sphingomyelins were significantly higher in the plasma EV isolated by precipitation or acoustic trapping compared to UC (Figure S9). In addition, the fold increase of Cer(d18:1/22:0), SM(d18:1/18:1) and sphinganine (d18:0) in plasma EV isolated by precipitation at time of presentation versus 1-month follow-up significantly correlated with the infarct size of the patients at 6-months post-MI ($R^2 = 0.91 p = 0.01, R^2 = 0.78 p = 0.05$, respectively) (Figure S10). This result further confirmed that plasma EV isolation methods, using targeted sphingolipidomic methods, are associated with unique method dependent profiles in patients following MI.

3.14 | Integrated-omics comparison of plasma EV isolation methods in MI patients

Finally, we integrated all the acquired plasma EV data from the MI patients in the acute presentation and follow-up time point. A principal component analysis was constructed to assess if the plasma EV profile at time of presentation differed from follow-up using a 95% confidence level. Integrated analysis showed that plasma EV profiles were only distinguishable at time of presentation with MI for precipitation and acoustic trapping, but not for those plasma EV isolated by UC, SEC and immunoaffinity capture (Figure 8). These findings indicate that different plasma EV isolation methods influence the diagnostic potential of plasma liberated EV following MI when multiple datasets are integrated to determine plasma EV profiles for potential panel biomarker discovery.

4 | DISCUSSION

Plasma EV-cargo may provide unparalleled insight into tissue homeostasis and pathological processes to facilitate identification of patients for focused therapies, but the methods to isolate plasma EV may impact the EV characteristics. Here, we found that; (I) the choice of plasma EV isolation method affected the plasma EV concentration, sphingolipid and proteomic profile, (II) but the five different plasma EV isolation methods shared a common EV protein and sphingolipid profile. (III) Plasma EV isolation



FIGURE 8 Principal component analysis of integrated plasma EV characterisation at time of presentation with myocardial infarction (MI) versus 1-month follow-up in the same patients using different plasma EV isolation methods. A principal component (PC) analysis of plasma EV isolated by ultracentrifugation (UC), precipitation, acoustic trapping, size-exclusion chromatography (SEC) and immunoaffinity capture from patients presenting with MI versus a 1-month follow-up control. Plasma EV characteristics including plasma EV concentration, EV-protein-Array and sphingolipidomics were integrated. The integrated data was condensed to PC1 and PC2 (*n* = 5–6). The eclipses indicate the 95% confidence interval.

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by immunoaffinity capture using anti-CD9, -CD63 and -CD81 coated antibody beads with the current protocol yields a similar profile to IgG control beads. (IV) The isolation method affected the ability to detect alterations in plasma EV sphingolipids and proteins in MI patients and (V) their association with infarct size determined by cardiac MRI 6 months post-MI. For example, the plasma EV concentration at time of presentation obtained using UC provides prognostic information, but this method is less suitable as a tool to generate EV with higher purity for use in mechanistic studies.

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Previous studies have sought to determine the influence of plasma EV isolation methods, but have often focused on comparing the net influence on the EV-proteome (Burrello et al., 2020; Gidlöf et al., 2019; Menezes-Neto et al., 2015; Veerman et al., 2021) or RNA profiles (Veerman et al., 2021) after isolation. These approaches can neglect the increased dimensionality of EV components, which carry numerous biologically active molecules such as proteins, lipids, RNA/DNA and metabolites. Furthermore, there is varied assessment of contaminations in EV preparations for use in "omics" studies, which may mask or skew the interpretation of datasets. The approach employed here utilizes the same independent biological replicates across multiple methods for plasma EV isolation and characterization, in conjunction with a control PBS/vehicle sample or an IgG for immunoaffinity capture beads. Here, the repeated analysis of the same plasma samples across multiple methods has enabled individual biological variability and potential plasma EV isolation irregularities to be explored. By using individual and integrated analysis, we showed that each method of plasma EV isolation was internally consistent; in so much that the samples clustered according to their isolation method in unsupervised hierarchical clustering and principal component analysis of all the acquired data.

Consistent with previous reports, our plasma EV-proteome was dependent on the plasma EV isolation method (Menezes-Neto et al., 2015; Veerman et al., 2021). However, our data shows that there are common plasma proteins, such as albumin and apolipoproteins ApoA-I and ApoB, present in all five isolation methods tested, with significant similarities to archived EV-databases (Kalra et al., 2012; Kim et al., 2013; Simpson et al., 2012) and published studies (Menezes-Neto et al., 2015; Veerman et al., 2021). In particular, the plasma EV-proteome indicated that UC/SEC cluster together based on the top 30 protein groups quantified, due to higher quantities of fibrinogens and immunoglobulins compared to the other methods. This is, potentially, a methodological constraint of employing untargeted mass spectrometry on plasma EV, which favors the most abundant peptides.

To better determine the protein profile of known plasma EV markers and cell associated markers, we utilized a high throughput EV-Array (Jorgensen, 2017) following plasma EV isolation from the five different methods. We found that CD9 and CD81 were the most abundant in plasma EV isolated by UC versus precipitation and SEC in healthy volunteers. Additionally, we found that cell-associated markers such as CD41 (platelets) and CD16 (immune cells) were enriched in UC and precipitation compared to other methods only in healthy volunteers. This high throughput antibody EV-Array circumvents issues of non-EV associated protein and lipid contaminants in plasma EV preparations. However, the five different plasma EV isolation methods used here may have influenced the EV-protein corona. Proteins such as albumin and apolipoproteins (ApoA-I, ApoB, ApoC-III and ApoE) show an association with the surface of EV (Tóth et al., 2021), which may influence antibody mediated binding to the array or subsequent detection of EV-associated proteins in this sandwich ELISA like technique. Protein coronas found on EV are affected by the isolation method (Tóth et al., 2021) and our data shows that the plasma EV-proteome and EV-Array acquired data are also influenced by the isolation method. Indeed, our western blot and proteomic data reported considerable variation in the amount of albumin across the methods, which is a predominant protein in the EV protein corona. In particular, precipitation isolated preparations was associated with lower albumin, which could be due to the presence of PEG and high salt concentrations in the isolation buffer that might be expected to influence albumin associations with the EV protein corona. The temporal dynamics of the EV-protein and possibly EV-lipid corona are poorly understood, and further elucidation of these interactions will be essential to understand the data from these preparations.

Immunoaffinity based methods, such as magnetic and polystyrene bead conjugations to specific antibodies, such as tetraspanins (CD9, CD63 and CD81), have been heralded as an important advancement in the capture of highly pure EV populations from the plasma. Immunoaffinity based methods could circumvent issues of lipoprotein and protein contamination in plasma EV yields, which are common in methods that use EV physical characteristics such as size and density for isolation such as UC and SEC (Tian et al., 2020; Veerman et al., 2021). To the best of our knowledge this is the first study to compare magnetic beads coated with antibodies for tetraspanins (CD9, CD63, and CD81) with a matched IgG control for plasma EV isolation using detailed integrated analyses. The plasma EV-profile of proteins and lipids for the immunoaffinity beads and matched IgG control was similar by Western blot, EV-Array, the number of peptides groups in proteomic profiles, GO pathway analysis and targeted sphingolipidomics. TEM imaging of IgG beads showed no visible EV-like particles or membranous structures, indicative of whole or sheared EV-particles, captured to the bead surface. However, we detected CD9, CD63 and CD9 on the IgG beads. The underlying factor giving rise to the disparity remains unknown, but we hypothesize that soluble EV-proteins from lysed or sheared EV during extraction and non-EV-associated proteins (such as C3, A2M, ALB and fibrinogens A and B) and lipids (such as ApoB, ApoE and ApoA-I) are interacting with the IgG beads, which have become associated with EV-profiles in omics studies and in archived databases (Kalra et al., 2012; Kim et al., 2013; Simpson et al., 2012). Studies using less complicated matrices, like conditioned cell culture media, show a more robust distinction between anti-CD9, CD63 and CD81 beads versus IgG control beads (Kowal et al., 2016; Mathieu et al., 2021), indicating that the similarity between tetraspanin and IgG antibody beads here might be matrix specific. Future plasma EV studies using antibody-bead isolation should include a matched IgG control when using omic-approaches.

Having determined plasma EV isolations method similarities and differences on the EV-characteristics and cargo in healthy volunteers, we assessed the influence of plasma EV isolation methods in a clinical situation where plasma EV number and composition are altered. We have previously shown that plasma EV are altered following MI to mediate long range signalling and induce the mobilization of splenic-neutrophils, splenic-monocytes and orchestrate their transcriptional programming (Akbar et al., 2017, 2022). The choice of plasma EV isolation method determined whether there was a higher concentration of EV immediately following MI compared to a 1-month follow-up control sample from the same patients. The new data here support our previous observations that the concentration of plasma EV isolated by UC from the time of presentation with MI correlates with the infarct size (Akbar et al., 2017). However, plasma EV concentrations derived from the other isolation methods in the same patients did not associate with infarct size. One possible explanation is the modest sample size utilised in this multi-method comparison (n = 6), which is less than we used in previous publications that explored the relationship between EV concentration and infarct size (n = 15-22) (Akbar et al., 2017, 2022).

Plasma EV sphingolipids are predictive of MI (Burrello et al., 2020). However, the influence of plasma EV isolation methods on the plasma EV-sphingolipidomic profile were not reported. We found a distinct sphingolipid profile between the plasma EV isolation methods. Precipitation-based EV isolation yields a significantly higher lipid content, principally due to higher proportions of sphingomyelins and ceramides. Sphingomyelins are found in cell membranes and associated with high- and low-density lipoproteins (Martínez-Beamonte et al., 2013; Sódar et al., 2016). The high presence of sphingomyelins and ceramide in the plasma EV profile of the precipitation-based method is likely due to the co-isolation of lipoproteins. PEG-based solutions co-isolate lipoproteins (Izzo et al., 1981). Lipoproteins contain ApoA-I and ApoB, which were higher in plasma EV derived by precipitationbased isolation when compared to the other methods by Western blot. Lipoproteins can also masquerade as EV-like particles in dynamic light scattering in techniques such as NTA (Brennan et al., 2020). Precipitation isolated plasma EV had the highest concentration of particles/mL when compared to other methods. EV Cer(d18:1/20:0) only showed elevation in precipitation-based isolation compared other methods. Similarly, precipitation isolated EV showed sphinganine(d18:0) correlated with infarct size at 6-months, but this was not the case for the other methods. Analysis of plasma EV-sphingolipid remains challenging due to constraints due to lipoparticle remnants (Simonsen, 2017); however, we show here for the first time there are common sphingolipids for different plasma EV isolation methods. In particular, plasma EV-protein and EV-sphingolipids clustered uniquely based on the isolation method, which may impact plasma EV diagnostics where a panel of proteins and sphingolipids are used for differentiation of clinical disease or outcome.

In summary, our data show that the choice of plasma EV isolation method influences the concentration of plasma EV, the EV-proteome and EV-sphingolipid profile in healthy volunteers and MI patients, where methodological differences determined associations with infarct size. Precipitation based plasma EV isolation gives the highest particle concentration and enriches for more sphingolipids, but co-isolate large quantities of apolipoproteins. In addition, immunoaffinity capture by using antibodies against tetraspanins yields similar EV-protein and EV-sphingolipid profiles to IgG control beads. Unbiased integrated analysis shows that plasma EV isolation methods cluster independently, but despite the selection predispositions imposed by each method, a core of EV associated proteins and lipids was detectable using all the approaches.

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CONFLICT OF INTEREST

The Author(s) declare that there is no conflict of interest.

ORCID

Naveed Akbar D https://orcid.org/0000-0003-4620-6373

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SUPPORTING INFORMATION

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