Characterization of extracellular vesicles and synthetic nanoparticles with four orthogonal single-particle analysis platforms

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Abstract
We compared four orthogonal technologies for sizing, counting, and phenotyping of extracellular vesicles (EVs) and synthetic particles. The platforms were: single-particle interferometric reflectance imaging sensing (SP-IRIS) with fluorescence, nanoparticle tracking analysis (NTA) with fluorescence, microfluidic resistive pulse sensing (MRPS), and nanoflow cytometry measurement (NFCM). EVs from the human T lymphocyte line H9 (high CD81, low CD63) and the promonocytic line U937 (low CD81, high CD63) were separated from culture conditioned medium (CCM) by differential ultracentrifugation (dUC) or a combination of ultrafiltration (UF) and size exclusion chromatography (SEC) and characterized by transmission electron microscopy (TEM) and Western blot (WB). Mixtures of synthetic particles (silica and polystyrene spheres) with known sizes and/or concentrations were also tested. MRPS and NFCM returned similar particle counts, while NTA detected counts approximately one order of magnitude lower for EVs, but not for synthetic particles. SP-IRIS events could not be used to estimate particle concentrations. For sizing, SP-IRIS, MRPS, and NFCM returned similar size profiles, with smaller sizes predominating (per power law distribution), but with sensitivity typically dropping off below diameters of 60 nm. NTA detected a population of particles with a mode diameter greater than 100 nm. Additionally, SP-IRIS, MRPS, and NFCM were able to identify at least three of four distinct size populations in a mixture of silica or polystyrene nanoparticles. Finally, for tetraspanin phenotyping, the SP-IRIS platform in fluorescence mode was able to detect at least two markers on the same particle, while NFCM detected either CD81 or CD63. Based on the results of this study, we can draw conclusions about existing single-particle analysis capabilities that may be useful for EV biomarker development and mechanistic studies.

KEYWORDS
etosomes, exosomes, extracellular vesicles, microvesicles, nanoflow cytometry, nanoparticle tracking analysis, resistive pulse sensing, single particle interferometric reflectance imaging sensing
1 | INTRODUCTION

Classification of extracellular vesicles (EVs) into subtypes has been proposed based on size, biogenesis pathway, separation procedure, cellular or tissue origin, and function, among others (Cocucci & Meldolesi, 2015; Lázaro-Ibáñez et al., 2019; Russell et al., 2019; Théry et al., 2018; Witwer & Théry, 2019; Yáñez-Mó et al., 2015). However, reproducible classification of EV subtypes will require single-particle characterization techniques including phenotyping by surface molecules or molecular signatures (Lässer et al., 2018; Tkach et al., 2018). In this sense, current knowledge of EV subtypes could be compared with knowledge of immune cells in the 1970s and early 1980s. Around that time, multiplexed flow cytometry capabilities and cell sorting were developed, allowing more precise identification, characterization, and molecular and functional profiling of immune cell subsets (Lanier et al., 1983). Single-particle technologies for much smaller biological entities will be needed to divide heterogeneous EV populations into well-defined and easily recognized subgroups.

In this study, we evaluated several particle types and single-particle characterization platforms. For input, we used a selection of biological and synthetic particles. EVs were separated from culture medium of H9 T lymphocytic cells and U937 promonocytic cells using several methods. These two cell lines were chosen because they display different levels of the tetraspanins CD63 and CD81. H9 cells have high CD81 and low CD63 levels, while U937 produce little CD81 but abundant CD63. Mixtures of distinct sizes of synthetic silica and traceable polystyrene beads were also tested, not because they mimic EVs or can serve as EV reference materials, but precisely because of their known size and composition, creating a ‘best case scenario’ to assess ability to measure particles. The technology platforms (Text Box 1) were: single-particle interferometric reflectance imaging sensing (SP-IRIS, NanoView) (Daaboul et al., 2016; Lopez et al., 2011; Reddington et al., 2013) with fluorescence, nanoparticle tracking analysis (NTA, ParticleMetrix) (Dragovic et al., 2011; Giebel & Helmbrecht, 2017; Sokolova et al., 2011) with fluorescence, microfluidic resistive pulse sensing (MRPS, Spectradyne) (Anderson et al., 2013; Fraikin et al., 2011; Giebel & Helmbrecht, 2017) (which does not have fluorescence capabilities), and nanoflow cytometry measurement (NFCM, NanoFCM) (Tian et al., 2018, 2020) with fluorescence.

2 | MATERIALS AND METHODS

Please see Table 1 for manufacturer, part number, and (where applicable) dilution of reagents.

Certain commercial equipment, instruments, and reagents are identified in this paper to foster understanding. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology or any other entity, nor does it imply that the materials or equipment identified are necessarily the best available for the purpose.

2.1 | Particle Preparation

Human cells lines H9 (T lymphocytic) and U937 (pro-monocytic) were obtained from the American Type Culture Collection (ATCC). Cells were maintained in Roswell Park Memorial Institute (RPMI) 1640 Medium supplemented with either replete or EV-depleted 10% heat-inactivated foetal bovine serum, with 1% HEPES buffer, 1% Penicillin-Streptomycin, and 1% L-Glutamine.
### Table 1 Reagents: manufacturer, part number, and dilution (where applicable)

| Antibodies                        | Manufacturer                  | Cat #       | Dilution |
|-----------------------------------|-------------------------------|-------------|----------|
| Primary Anti-CD81                 | Santa Cruz, Dallas, TX        | sc-23962    | 1:500    |
| Primary Anti-CD63                 | BD Pharmigen, San Diego, CA   | 556019      | 1:1000   |
| Primary Anti-CD9                  | BioLegend, San Diego, CA      | 31202       | 1:1000   |
| Primary Anti-TSG101               | AbCam, Cambridge, MA          | ab125011    | 1:1000   |
| Primary Anti-Calnexin             | AbCam Cambridge, MA           | ab22595     | 1:2000   |
| Primary Anti-BiP/GRP78            | BD Pharmigen, San Diego, CA   | 619078      | 1:500    |
| Primary Anti-GMI30                | AbCam, Cambridge, MA          | ab52649     | 1:400    |
| Secondary Mouse Anti-Rabbit IgG BP-HRP | Santa Cruz, Dallas, TX  | sc-2357    | 1:5000   |
| Secondary Rabbit Anti-Mouse IgG BP-HRP | Santa Cruz, Dallas, TX  | 516102     | 1:5000   |
| PE-Conjugated Mouse Anti-Human CD81 | BD Biosciences, Franklin Lakes, NJ | 555676 | n/a      |
| AF488-Conjugated Mouse Anti-Human CD63 | Novus Biologicals, Littleton, CO | NBP2-42225 | n/a      |

| Reagents                          | Manufacturer                  | Cat #       |
|-----------------------------------|-------------------------------|-------------|
| 3K MWCO Centricon Plus-70         | Millipore Sigma               | UFC700308   |
| 3K MWCO Amicon Ultra-15           | Millipore Sigma               | UFC900396   |
| Blotting Grade Blocker            | Bio-Rad                       | 170-6404    |
| Carbon Coated 400 Mesh Copper Grids | Electron Microscopy Science  | CF400-Gu-UL |
| Criterion TGX Stain-Free Precast Gel | Bio-Rad                       | 5678084    |
| Distilled Water                   | Gibco                         | 15230-162   |
| FluoSpheres Carboxylate-Modified Microspheres, 0.1 µm, Yellow Green Fluorescent | Thermo Scientific     | F8803       |
| H9 Cell Line                      | American Type Culture Collection | HTB-176     |
| Heat-Inactivated Fetal Bovine Serum | GE Healthcare               | SH30396.03  |
| Heat-Inactivated Fetal Bovine Serum, Exosome-Depleted | Gibco                  | A2720801    |
| HEPES buffer                      | Gibco                         | 15630080    |
| Immuno-Blot PVDF Membrane         | Bio-Rad                       | 620177      |
| L-Glutamine                       | Gibco                         | 25030081    |
| Open-Top Thin Wall Ultra-Clear Tubes | Beckman Coulter             | 344091      |
| Penicillin-Streptomycin           | Gibco                         | 15140122    |
| Phosphatidylserine Beads          | NanoFCM                       | S16M-Exo    |
| Phosphate-Buffered Saline (PBS)   | Gibco                         | 14190-144   |
| Pierce BCA Protein Assay Kit      | Thermo Scientific             | 23225       |
| Polypropylene Ultracentrifugation (UC) Tubes | Sorvall              | 03-141      |
| Polystyrene Spheres 150 nm        | Thermo Scientific             | 3150A       |
| Polystyrene Spheres 125 nm        | Thermo Scientific             | 3125A       |
| Polystyrene Spheres 90 nm         | Thermo Scientific             | 3090A       |
| Polystyrene Spheres 70 nm         | Thermo Scientific             | 3070A       |
| Protease Inhibitor Cocktail       | Millipore Sigma               | 11697498001 |
| RIPA                              | Cell Signaling Technology     | 9806        |
| Roswell Park Memorial Institute (RPMI) 640 Medium | Gibco                     | I875093     |
| Silica Nanosphere Cocktail        | NanoFCM                       | n/a         |
| SuperSignal West Pico PLUS Chemiluminescent Substrate | Thermo Scientific     | 34577       |
| Swinging Bucket Rotor AH-629      | Thermo Scientific             | 54284       |
| Tris Buffered Saline (TBS)        | Bio-Rad                       | 1706435     |
| Tween-20                          | Millipore Sigma               | P7949       |
| U937 Cell Line                    | American Type Culture Collection | CRL-1593.2  |
| Ultra-Pure Distilled Water        | Invitrogen                    | 10977015    |
| Uranyl Acetate                    | Polysciences                  | 2144725     |
Text Box 1: Evaluated Technologies

Single-particle interferometric reflectance imaging sensing (SP-IRIS) captures particles (e.g., EVs) onto a chip by affinity reagents, usually antibodies, to surface antigens. Particles are imaged by interferometric reflectance for sizing and counting, and fluorescence detection may be done for up to three channels for surface antigens or internal molecules following fixation and permeabilization. Website for the platform we used: https://www.nanoviewbio.com/

Nanoparticle tracking analysis (NTA) is an optical method to track single particles and assign sizes and counts. Measuring Brownian motion allows calculation of a hydrodynamic sphere-equivalent radius of each tracked particle. Additionally, fluorescence filters can be used for detection of particle-associated fluorescence moieties channels. Website for the platform we used: https://www.particle-metrix.de/en/particle-metrix

Microfluidic resistive pulse sensing (MRPS) counts and sizes particles as they pass through a pore between microfluidic chambers. Occlusion of the pore results in a measurable change in electrical signal (defining an event) that is proportional to the volume of the particle. Often, this technique uses different disposable cartridge pore sizes to detect particle populations within specific size ranges. As a non-optical technology, fluorescence detection is not available. Website for the platform we used: https://nanoparticleanalyzer.com/

Nanoflow cytometry measurement (NFCM) is a flow-based technique that detects nano-sized particles by scatter and/or fluorescence. Compared with traditional flow cytometry, a smaller flow channel reduces background signal, and lower system pressure increases dwell time of particles for enhanced signal integration. Website for the platform we used: http://www.nanofcm.com/products/flow-nanoanalyzer

Cells were cultured at 37°C in 5% CO2. Silica spheres (SS, NanoFCM, Nottingham, England) were a premixed combination of diameters 68 nm, 91 nm, 113 nm and 151 nm. Individual polystyrene spheres (PS, Thermo Fisher) of diameters 70 nm, 90 nm, 125 nm, and 150 nm were purchased. Nominal equal concentrations (1 x 10^12 particles/ml) of beads were mixed.

2.2 Size-exclusion chromatography (SEC)

A total of 60 ml of culture-conditioned medium (CCM) from each cell line was centrifuged at 1000 x g for 5 min at 4°C to remove cells and cellular debris. 3 kDa molecular weight cut off (MWCO) Centricon Plus-70 centrifugal filters (Millipore Sigma) were used to concentrate the initial volume to 1.5 ml. Size exclusion chromatography (SEC) was done with qEV Automated Fraction Collectors (AFC; Izon Science, Cambridge, MA) and qEV original 70 nm columns (Izon Science, Cambridge, MA). Columns were left at room temperature for 30 min and washed with phosphate-buffered saline (PBS). 0.5 ml of concentrated CCM was loaded onto each of three columns, and 0.5 ml fractions were collected by adding additional PBS to the column. EV-enriched fractions (SEC; fractions 7–9) were pooled altogether from the three columns used for each sample and further concentrated using 3 kDa MWCO Amicon Ultra-15 Centrifugal Filters to a final volume of 1 ml. 50-μl aliquots were stored at –20°C for downstream assays.

2.3 Differential ultracentrifugation (dUC)

60 ml of CCM from each cell line was centrifuged at 1000 x g for 5 min at 4°C to remove cells and cellular debris and 2000 x g for 10 min at 4°C to remove additional debris. The supernatant was transferred to polypropylene thin-wall ultracentrifugation (UC) tubes and centrifuged at 10000 x g for 30 min at 4°C using a swinging bucket rotor (Thermo Scientific rotor model AH-629, k-factor 242, acceleration and deceleration settings of 9) to pellet large EVs. Supernatant was transferred into new polypropylene thin wall UC tubes and centrifuged at 100,000 x g for 70 min at 4°C using the same swinging bucket rotor. The 100K pellets containing small EVs were resuspended in 1 ml of PBS, vigorously vortexed, and placed on ice for 20 min. 50-μl aliquots were stored at –20°C for downstream assays.

2.4 Transmission electron microscopy (TEM)

A total of 10 μl freshly thawed aliquots were adsorbed to glow-discharged carbon-coated 400 mesh copper grids by flotation for 2 min. Grids were quickly blotted and rinsed by flotation on three drops (1 min each) of 1X Tris-buffered saline. Grids were negatively stained in 2 consecutive drops of 1% uranyl acetate (UAT) with tylose (1% UAT in deionized water (dH2O), double
filtered through a 0.22 µm filter), blotted, then quickly aspirated to cover the sample with a thin layer of stain. Grids were imaged on a Hitachi 7600 TEM operating at 80 kV with an AMT XR80 CCD (8 megapixel). SS and PS were absorbed to grids as above, but with initial flotation for 5 min and imaging on a Phillips CM-120 TEM operating at 80 kV with an AMT XR80 CCD (8 megapixel).

2.5 Western blot (WB)

H9 and U937 cell pellets and isolated EVs were lysed in 1× radioimmunoprecipitation assay buffer (RIPA) supplemented with protease inhibitor cocktail. Protein quantification of cell and EV lysates was done using a bicinchoninic acid assay (BCA) (Pierce BCA Protein Assay Kit). A total of 5 µg of lysates were resolved using a 4% to 15% Criterion TGX Stain-Free Precast gel, then transferred onto an Immuno-Blot PVDF membrane. Blots were probed using primary antibodies in PBS-T and 5% Blotting Grade Blocker. Primary antibodies were against CD81, CD63, CD9, TSG101, calnexin, BiP/GRP78 and GM130. Secondary antibodies were rabbit anti-mouse IgGk BP-HRP and mouse anti-rabbit IgGk BP-HRP. SuperSignal West Pico PLUS Chemiluminescent Substrate was used for detection and blots were visualized with an iBright Western Blot (Thermo Fisher, Waltham, MA) imaging system.

2.6 Single particle interferometric reflectance imaging (SP-IRIS)

Measurements were performed largely as described previously (Ahn et al., 2020; Gámbaro et al., 2019). A total of 35 µl of H9 and U937 EVs isolated by SEC or dUC were diluted 1:1 in incubation buffer (IB) and incubated at room temperature on ExoView R100 (NanoView Biosciences, Brighton, MA) chips printed with anti-human CD81 (JS-81), anti-human CD63 (H5C6), anti-human CD9 (H9a), and anti-mouse IgG1 (MOPC-21). After incubation for 16 h, chips were washed with IB 4 times for 3 min each under gentle horizontal agitation at 500 rpm. Chips were then incubated for 1 h at room temperature with a fluorescent antibody cocktail of anti-human CD81 (JS-81, CF555), anti-human CD63 (H5C6, CF647), and anti-human CD9 (H9a, CF488A) at a dilution of 1:1200 (v:v) in a 1:1 (v:v) mixture of IB and blocking buffer. The buffer was then exchanged to IB only, followed by 1 wash with IB, 3 washes with wash buffer, and 1 wash with rinse buffer (3 min each at 500 rpm). Chips were immersed twice in rinse buffer for approximately 5 s each and removed at a 45-degree angle to allow the liquid to vacate the chip. All reagents and antibodies were supplied by NanoView Biosciences (Brighton, MA, Cat # EV-TETRA-C). Both SS and PS were diluted in dH2O to load 10,000 particles, nominally, per antibody capture spot on the ExoView chips. A total of 35 µl of diluted spheres were incubated on ExoView chips and allowed to fully dry. All chips were imaged in the ExoView scanner (NanoView Biosciences, Brighton, MA) by interferometric reflectance imaging and fluorescent detection. Data were analyzed using NanoViewer 2.8.10 Software (NanoView Biosciences). Fluorescent cutoffs were as follows: CF555 channel 230, CF488 channel 475, CF647 channel 250 (biological particles) and CF555 channel 675, CF488 channel 600 and CF647 channel 375 (SS and PS).

2.7 Nanoparticle tracking analysis (NTA)

ZetaView QUATT-NTA Nanoparticle Tracking-Video Microscope PMX-420 and BASIC NTA-Nanoparticle Tracking Video Microscope PMX-120 (Particle Metrix, Inning am Ammersee, Germany) instruments were used for particle quantification in both scatter and fluorescence (488 nm) modes. Calibration beads and biological samples were diluted in distilled water and PBS, respectively, to a final volume of 1 ml. Calibration was done for both scatter and fluorescence measurements. For scatter-mode calibration, 100 nm PS beads were diluted 1:250,000 (v:v). Capture settings were: sensitivity 65, shutter 100, minimum trace length 10. Cell temperature was maintained at 25°C for all measurements. For fluorescence calibration, 488 nm yellow-green FluoSpheres were diluted 1:250,000 (v:v), and both scatter and fluorescence were measured. Scatter was recorded as above, and fluorescence was measured at sensitivity 80, shutter 100, and minimum trace length 15. To measure SS and PS mixtures and individual size populations of PS, samples were diluted such that at least 200 particles could be counted per frame. Technical triplicates were measured for each sample. A washing step was done between each measurement using dH2O. For H9 and U937 EVs separated by SEC or dUC, one cycle was performed by scanning 11 cell positions. Capture was done at medium video setting, corresponding to 30 frames per second. ZetaView software 8.5.10 was used to analyze the recorded videos with the following settings: minimum brightness 30, maximum brightness 255, minimum area 10, and maximum area 1000. Since subpopulations of particles might also be identified based on signal intensity, we used manual and population distribution gates in the ZetaView software to assess this possibility for SS and PS mixtures. PE-conjugated mouse anti-human CD81 and AF488-conjugated mouse anti-human CD63 were used for fluorescence detection of EVs. Antibodies were mixed 1:9 (v:v) with PBS, incubated 2 h at room temperature, and diluted to a final volume of 1 ml. Supplementary Table 2 lists all antibodies tested with this platform.
2.8 | Microfluidic resistive pulse sensing (MRPS)

Microfluidics resistive pulse sensing measurements were conducted using the nCSI instrument (Spectradyne, Torrance, CA) as described previously (Ahn et al., 2020). For biological particles, sample volumes of a few µl of H9 and U937 EVs isolated by SEC or dUC were diluted with an equal volume of 1% polysorbate 20 (Tween 20) in 1× PBS (PBST) and further diluted as indicated with 1× PBS, and loaded onto polydimethylsiloxane cartridges (diameter range 65 nm to 400 nm). A different cartridge was used for each sample and replicate. Approximately 5 µl of the diluted sample was used and about 25,000 events were recorded for each analyte. For synthetic nanoparticles, SS and PS were diluted 100-fold by volume in dIH2O, then 10-fold by volume with equal volumes of PBST and the remainder with 1× PBS and loaded onto TS-400 polydimethylsiloxane cartridges. Approximately 3,000 events were obtained for each SS and PS repeat. All acquired results were analyzed using the nCSI Data Analyzer (Spectradyne, Torrance, CA). For all samples, user-defined filtering was applied by defining 2D polygonal boundaries based on transition time and diameter to exclude false positive signals, similar to gating commonly used in analyzing flow cytometry data. Effects of Tween 20 on EV integrity or counts were assessed by diluting samples to a final concentration of Tween 20 (in PBS) ranging from 0.1% to 0.9%.

2.9 | Nano-flow cytometry measurement (NFCM)

The nFCM flow nano-analyzer was used to measure concentration and size of particles following the manufacturer's instructions and as described previously (Huang et al., 2020). Briefly, two single photon-counting avalanche photodiodes (APDs) were used for the simultaneous detection of side scatter (SSC) and fluorescence of individual particles. The instrument was calibrated separately for concentration and size using 200 nm PE- and AF488 fluorophore-conjugated PS beads and a Silica Nanosphere Cocktail, respectively. 20 µl of each EV preparation was incubated with 20 µl PE-conjugated CD81 and 5 µl AF488-conjugated CD63 antibodies at 37°C for 30 min. After incubation, the mixture was washed twice with PBS and centrifuged at 110,000 × g for 70 min at 4°C (TH-641 rotor, k-factor 114, Thermo Fisher, using thin-wall polypropylene tubes with 13.2 ml capacity and acceleration and deceleration settings of 9). The pellet was resuspended in 50 µl PBS. Events were recorded for 1 min. Using the calibration curve, the flow rate and side scattering intensity were converted into corresponding particle concentrations and size.

2.10 | Dynamic light scattering (DLS)

To check the nominal size values of PS beads, particle diameter was measured by dynamic light scattering using a Malvern Zetasizer Nano-ZS90. Each suspension was diluted 10 × in ultrapure water, and measurements were carried out in triplicate at 25°C. A single peak was observed for each individual run.

2.11 | Data and methods availability

We have submitted all relevant data of our experiments to the EV-TRACK knowledgebase (EV-TRACK ID: EV200090) (Van Deun et al., 2017). Reporting for NFCM was submitted to FlowRepository as ID:FR-FCM-Z2U3 (Welsh et al., 2020). Any and all data are available on reasonable request.

3 | RESULTS

3.1 | Production, separation and characterization of input materials

Supernatants were collected from cultured human cell lines: H9 (T-lymphocytic) and U937 (pro-monocytic). Our goal was to obtain EV-enriched or -depleted biological material from cells with different tetraspanin expression. EVs were partially separated by size exclusion chromatography and ultrafiltration or differential ultracentrifugation (Figure 1a). Marker expression and morphology were assessed by WB (Figure 1b and Supplementary Figure 1) and TEM. WB revealed characteristic cellular CD63 and CD81 expression patterns, with CD81 above the limit of detection only for H9 and CD63 predominating for U937 (Figure 1b). CD81 was apparently enriched in EV fractions from H9, while CD63 appeared to be present, but not enriched, in EVs from U937, suggesting relatively inefficient release. Please note, however, that protein amount was used to normalize WB input, so per-particle content cannot be easily compared across sample types, and see also additional blots in Supplementary Figure 1B-D. Calnexin was detected in cell lysates, with little or no signal in EV fractions (Figure 1b). For EVs concentrated and
FIGURE 1  Methodology and EV separation. [(a) EVs were separated from H9 and U937 culture-conditioned media by a combination of ultrafiltration and size exclusion chromatography (SEC EVs) or by differential ultracentrifugation (100K EVs). (b) Immunoblots of cell lysates from H9 and U937, EVs separated by ultracentrifugation (100K EVs) and SEC (SEC EVs), and later fractions of SEC (enriched for protein; SEC-P). Antibodies are specified in Table 1; see also Supplementary Figure 1. (c) Electron micrograph of SEC EVs and 100K EVs from both cell lines. As indicated for each subpanel, leftmost scale bars represent 500 nm at magnification 40,000x; rightmost scale bars are 100 nm at magnification 100,000x. (d) EM of SS and PS. Leftmost scale bars are 500 nm at magnification 17,500x; rightmost scale bars are 100 nm at magnification 65,000x.]
separated by each method, TEM showed heterogeneous populations (particles ranging from approximately 50 to approximately 500 nm in diameter) including EVs with the typical ‘cup-shaped’ fixation artefact (Figure 1c and Supplementary Figure 2). UC pellets displayed higher background and apparent non-EV particles than SEC EV fractions, possibly consistent with proteinaceous material that that elutes in later, relatively EV-depleted fractions of SEC (Figure 1c and Supplementary Figure 2).

Silica spheres (SS) and polystyrene spheres (PS) of known size were obtained from commercial sources. These artificial nanoparticles were measured here not as reference materials for EV studies, but simply because they have known diameters and composition, along with higher refractive indices (RIs) than EVs. Note that such particles can be used as reference materials in EV studies if the RI of the material is accounted for, for example with several available software packages (Exometry, 2020; Welsh & Jones, 2020; Welsh et al., 2020). For PS, we used National Institute of Standards and Technology (NIST)-traceable size standards. These beads are among the most commonly used size calibrants for materials in their size ranges and are compared with a known standard maintained by NIST. A Certificate of Calibration and Traceability allows labs to show compliance with various ISO and GMP standards and regulations. Additionally, uncertainty of measurement is indicated on a certificate of analysis for each bead lot. We nevertheless confirmed size and purity of SS and PS mixtures using TEM (Figure 1d). Beads corresponding to all four sizes in each mixture were clearly present on the grids, with little or no contaminating material. Bead diameters as measured by TEM were consistent with the nominal diameters and data sheet specifications (Supplementary Table 1, n = at least 30 per population over 4 TEM frames). We also measured individual PS bead populations by dynamic light scattering (DLS), a method best suited for measurement of monodisperse populations (three bead preparations each, measured thrice each). Results showed a single peak for each individual run and polydispersity indices consistent with monodispersity (Supplementary Table 1).

3.2 Artificial nanoparticle sizing

Mixed silica spheres (SS) with nominal diameters of 68 nm, 91 nm, 113 nm and 151 nm were measured with the four platforms. SP-IRIS identified four distinct populations with diameters around 75 nm, 100 nm, 120 nm and 150 nm (Figure 2a). Since the SP-IRIS technology uses affinity to capture particles, particle mixtures were dried onto the SP-IRIS chips before imaging. NTA detected a broad population distribution with a mode around 105 nm diameter (Figure 2b). MRPS resolved four distinct peaks for each individual chip, but this distinction was masked somewhat by averaging all results (Figure 2c; see inset for an example of an individual reading and also Supplementary Figure 3). NFCM resolved four populations with distinct peaks at diameters of approximately 66 nm, 85 nm, 112 nm and 154 nm (Figure 2d). Polystyrene spheres (PS) with nominal diameters 70 nm, 90 nm, 125 nm, and 150 nm were mixed to a nominal concentration of 1 × 10^12 particles/ml. SP-IRIS detected four distinct peaks around 80 nm, 110 nm, 140 nm and 170 nm (Figure 2e). NTA returned a broad population distribution centred around 105 nm (Figure 2f). MRPS identified distinct peaks at diameters 71 nm, 92 nm, 123 nm and 150 nm (Figure 2g). Nano-flow showed four populations around 85 nm, 120 nm, 170 nm and 225 nm in diameter, as well as a possible smaller population around 60 nm (Figure 2h). We also measured several dilutions of SS and PS particles using MRPS and NFCM, with results similar to those described above. Raw and dilution-corrected data are presented in Supplementary Figure 4.

Because the NTA platform did not appear to resolve different populations, we also assessed individual PS bead sizes and also tried to use intensities to resolve individual bead populations. Individually, the four bead sizes returned measurements (arithmetic mean +/- SD) of 109.0 nm +/- 0.4 nm (70 nm PS), 105.2 nm +/- 0.3 nm (90 nm PS), 126.5 nm +/- 0.4 nm (125 nm PS), and 148.7 nm +/- 3.6 nm (150 nm PS). Mixed beads again produced a single broad peak averaging 124.2 nm. Using the NTA software to assign gates based on intensity, we assessed the possibility that individual bead sizes could be resolved. For PS beads, the most intense signals skewed slightly towards larger returned sizes (Supplementary Figure 5A); however, there was no apparent size or distribution difference between medium- and low-intensity populations (Supplementary Figure 5A,B). Similar results were obtained for SS beads (Supplementary Figure 5C). Gating on intensity might, however, be a useful tool in some settings.

3.3 Counting of synthetic nanoparticles

In addition to particle size, we also assessed counts. For SP-IRIS, a mean of around 3000 SS particles were detected per printed antibody spot (Figure 3a), with no overall differences between groups of antibody spots (i.e., three spots per chip each of three tetraspanins and an isotype control; note that no differences would be expected, since particles were dried onto the chips). However, per-spot events ranged considerably from <2000 SS particles per spot to >4500 SS particles per spot (Figure 3a). SP-IRIS performed similarly for PS. There were no differences between spots printed with different antibodies, with a mean of around 1400 events/antibody spot (Figure 3b), but events per spot ranged from <1000 PS particles/spot to 3000 PS particles/spot. Based on the nominal PS bead concentration and the surface area of the chips, 10,000 particles per spot would have been expected (Figure 3b, dotted line); however, it is possible that beads may not have dried evenly, for example, if they were relatively repelled by antibody-printed surfaces. Following SP-IRIS measurements, chips were probed with three fluorescently labelled antibodies (anti-CD81, anti-CD63, and anti-CD9) to assess background binding. Background binding was negligible for both SS and PS.
FIGURE 2 SS and PS size distribution. [Size distributions for SS (n = 3) with standard deviation for (a) SP-IRIS, (b) NTA, (c) MRPS, and (d) NFCM. Nominal SS diameters are indicated by vertical dotted lines: 68 nm, 91 nm, 113 nm, and 151 nm. Size distributions for PS (n = 3; with SD) for (e) SP-IRIS, (f) NTA, (g) MRPS, and (h) NFCM. Nominal PS diameters are indicated by vertical dotted lines: 70 nm, 90 nm, 125 nm, and 150 nm. Inset in Figure 2c shows a single MRPS measurement of the size distribution; see also Supplementary Figure 3 for individual readings. 5-nm bin sizes were used for all graphs.]
SS and PS quantification. [(a) SP-IRIS label-free capture for SS and PS using four capture spots (n = 3 per group; mean particle count per spot with SD). b) SS quantification (n = 3; mean particles/ml with SD). c) PS quantification (n = 3; mean particles/ml with SD). In panels b and d, nominal PS concentration is indicated by a horizontal dotted line (1.0 × 10^{12} particles/ml).]

(Supplementary Figure 6A and B, respectively). Some outliers were observed for CD9 (SS) or CD63 (PS); however, none exceeded 1000 events. Particle concentrations were also measured by NTA, MRPS, and NFCM. For SS (Figure 3c), MRPS estimated a concentration approximately one log higher than NTA (5.1 × 10^{11} particles/ml vs. 5.4 × 10^{10} particles/ml, respectively), with NFCM in the middle (1.7 × 10^{11} particles/ml). For PS, all three methods were in close agreement (Figure 3d). The difference for SS, but not for PS, likely reflects the sensitivity of the optical techniques to the refractive index of the particles being counted. Furthermore, the measured concentration was very close to the nominal PS concentration of 1 × 10^{12} particles/ml (Figure 3d, dotted line).

### 3.4 Biological particle sizing

EV preparations from H9 and U937 cell supernatants enriched by ultrafiltration and SEC (SEC EVs) or by differential ultracentrifugation (100K EVs) were next measured using each platform. For H9-derived materials, SP-IRIS returned an almost identical size distribution profile for both EV enrichment methods (Figure 4a). In contrast, NTA, MRPS, and NFCM returned data indicating particles at smaller diameters for the 100K EVs compared with the SEC EVs, with roughly similar particle size distributions (Figure 4b-d). However, substantial variation between replicates might limit the conclusions that can be drawn from this observation; we also expect that the polydisperse nature of EVs will naturally lead to greater CVs. For U937-derived materials, SP-IRIS and NTA (Figure 4e,f) detected more particles at smaller diameters from the 100K EVs compared with the SEC-EVs, again with roughly similar particle size distribution. MRPS produced equivalent particle size distribution and particle number between the two enrichment techniques (Figure 4g). In contrast, NFCM detected a higher particle count of smaller particle diameters from the SEC EVs than the 100K EVs, with the particle size distributions significantly different (Figure 4h). Please see Supplementary Figure 7 for plots drawn without error bars for clarity. Again, variability between replicates limits conclusions. Overall, the results are broadly consistent with the reported power-law size distribution of EVs (Paulaitis et al., 2018; Van Der Pol et al., 2016) and the expectation that UC pellets may contain non-EV extracellular particles (EPs) around the same size as EVs (Théry et al., 2018).

### 3.5 Biological particle counting

Particle counts were next assessed. As before, we present the SP-IRIS data separately because this platform does not provide an overall particle count, but rather a number of events detected on surfaces printed with antibodies (shown here: to CD81 and to
CD63 plus an isotype control). Consistent with expectations based on cellular tetraspanin expression and release, SP-IRIS showed that more H9 particles were captured by anti-CD81 than by anti-CD63 (Figure 5a) and that U937 particles could be captured by CD63 capture antibodies and not by CD81 capture antibodies (Figure 5b). For the remaining three platforms, which measure overall concentration, several trends were apparent (Figure 5c,d). First, for both the H9 and the U937 source, and for both EV separation methods, data were consistent with the results of SS counting in that NTA, NFCM, and MRPS measurements ordinally ranked from least particles/ml to greatest particles/ml. Second, MRPS and NFCM measured greater particle concentrations for 100K EVs than for SEC EVs (corrected for processing and dilution), although NTA results were similar. Finally, this is in contrast to results for the PS particles, where the three techniques returned roughly equivalent particle counts.
**3.6 Do low concentrations of detergent affect MRPS measurements?**

During the review process for this manuscript, a question arose about the possible effect of detergent on the reported results for biological particles measured by MRPS since a manufacturer-recommended dilution buffer contains 1% Tween 20. Specifically, it was proposed that the higher particle counts obtained for the same samples by MRPS and NTA could be due to artifactual small particle production when EVs are disrupted by detergent (see the comments section of (Mallick et al., 2020). We thus studied the effects of different concentrations of Tween 20 on MRPS measurements using archived aliquots of H9 EVs and U937 EVs. Because the maximum Tween 20 concentration used in any reported experiment was 0.5%, we conducted dilution series such that the same biological samples were measured in the presence of Tween 20 concentrations ranging from 0.1% to 0.9%. For reference, our highest concentration of Tween 20 was well below the concentrations previously reported to affect any of three classes of EVs (Osteikoetxea et al., 2015). Across Tween 20 concentrations, measured EV concentrations averaged $1.7 \times 10^{11} \pm 2.2 \times 10^{10}$ (H9 EVs) and $1.5 \times 10^{11} \pm 2.9 \times 10^{10}$ (U937 EVs), with no correlation between counts and detergent concentration (Supplementary Figure 8). Despite some variability in size profiles, there was also no evidence of a clear decrease in particle size with increasing detergent concentration (Supplementary Figure 8B,C).

**3.7 Single particle phenotyping by fluorescence**

Fluorescence measurements for biological particles were done with three platforms. The MRPS platform has no fluorescence capabilities, so it was used only for sizing and counting. SP-IRIS performs a kind of single-particle phenotyping even in label-free mode, since diameter is measured for individual particles captured by antibodies and thus putatively positive for an antigen. What is more, captured particles can additionally be probed with fluorescently labelled antibodies. For chips incubated with H9 EVs (Figure 6a,b), EVs captured by CD81 antibodies were also generally positive for CD81 by fluorescence, and some also appeared to be CD63 positive. In contrast, CD63 capture spots were largely devoid of fluorescence for H9 EVs, as were (most) control capture spots. For chips incubated with U937 EVs (Figure 6c,d), events on CD63 capture spots were also positive for CD63 by fluorescence. CD81-linked fluorescence was at background levels for all spots. Note that numbers of ‘positive’ events are higher in fluorescence mode than with label-free imaging (Figure 5a,b), likely, as discussed later, because fluorescence detection is more sensitive than reflectance imaging.
FIGURE 6  Particle phenotyping. [SP-IRIS fluorescence detection using labelled anti-CD81 and anti-CD63 after particle capture with CD81, CD63, and mouse isotype control (n = 3 per group; mean and SD) for (a) H9 SEC EVs, (b) H9 100K EVs, (c) U937 SEC EVs, and (d) U937 100K EVs. Percent of particles detected with fluorescently-labelled anti-CD81 and anti-CD63 by NTA and NFCM (n = 3 per group; mean and SD) for (e) H9 SEC EVs, (f) H9 100K EVs, (g) U937 SEC EVs, and (h) U937 100K EVs. Asterisk. An asterisk indicates that, in the authors’ view, an antibody did not perform on the instrument; it does not necessarily mean that the antibody would not perform in another context or with additional optimization.]
For the two remaining platforms with fluorescence capabilities, NTA and NFCM, results are shown as percent of total particles (Figure 6e-h). Approximately 40% to 50% of detected particles from H9 cells were positive for CD81 according to fluorescent NTA, while little to no CD81 signal was detected for U937 materials, consistent with expectations. However, we could not detect CD63-linked signal by fluorescent NTA for any sample. In contrast, NFCM detected either CD81 or CD63 on a small percentage of particles. The percentages were similar for the two tetraspanins for H9-derived particles. For U937 material, CD63-positive particles were more abundant than CD81-positive particles. No major differences between the SEC and 100K separation methods were apparent according to these data (Figure 6e-h).

3.8 Background signal: buffer and process controls

As a final note, we would like to mention the issue of background particle counts and fluorescence. It is important to regularly assess background signal obtained from buffer only and process controls (e.g., for cell culture experiments, prepared from unconditioned medium subjected to the same protocols used for EV separation). However, in our experience, if buffers and medium have been prepared correctly, results are typically below what could be considered a reasonable range of detection for each instrument. For example, when process controls were prepared from unconditioned culture medium by differential centrifugation or SEC, SP-IRIS recorded only several hundred events per affinity spot (as compared with several thousand for conditioned samples), with no differences between tetraspanin and isotype control capture (Supplementary Figure 9, top panel). In fluorescence mode, background levels were even lower, in the tens of events, with no differences between tetraspanin and isotype control capture or between controls prepared by differential centrifugation or SEC (Supplementary Figure 9, bottom panels). See Table 2 for estimates of useful range of measurement for each instrument; also, for the NTA instrument we used, there is a default minimum number of recorded tracks at each or most read positions. In sum, background signal is usually an order of magnitude or more below readings obtained with experimental samples.

4 DISCUSSION

This study evaluated the abilities of four orthogonal technology platforms to size, count, and/or phenotype biological EVs and synthetic nanoparticles. Three of the technologies—SP-IRIS, NTA, and NFCM—are optical in nature and can perform some form of phenotyping/fluorescence analysis, while the other, MRPS, is an electric sensing platform that we did not attempt to apply to particle phenotyping. Although numerous comparisons of EV characterization platforms have been published previously (Anderson et al., 2013; Bachurski et al., 2019; Corso et al., 2019; Erdbrügger & Lannigan, 2016; Van Der Pol et al., 2014; Vogel et al., 2021), this study includes NFCM and MRPS and focuses in part on single-particle phenotyping.

4.1 Detected particles: size-range sensitivity and refractive index matter

Whereas NTA, MRPS, and NFCM accurately and consistently measured the concentration of a known mixture of polystyrene particles, estimates of the number of silica particles varied substantially. NTA measured approximately 10-fold fewer SS particles than MRPS, while NFCM measured ~ 3-fold fewer SS particles than MRPS. Since SS have a lower refractive index (nSS ∼ 1.42 (García-Santamaría et al., 2002)) than PS (nPS ∼ 1.59 (Kasarova et al., 2007)), one might predict that a mixture of EVs, with an even lower refractive index than silica (Gardiner et al., 2014; Van Der Pol et al., 2014), would have even larger variability between methods. Indeed, for EV preparations, average counts by NTA and MRPS differed by between one and two orders of magnitude. These outcomes emphasize that each platform has an effective range of measurement. MRPS is not sensitive to refractive index, but cartridges may clog (although we did not see evidence of this). In contrast, optical methods are quite sensitive to refractive index, and r\(^6\) variation of scattered intensity limits dynamic range for a single instrument setting. Thus, differences in output in part reflect different or overlapping particle populations that can be detected by the specific technologies, as indeed reported previously for several of these technologies (Van Der Pol et al., 2014). That is, NTA and MRPS are similarly capable to detect a wide range of PS particle sizes. However, NTA may detect a more limited range of biological particles (Defante et al., 2018) than the MRPS platform using a small pore-size cartridge, in that MRPS may detect more of the smaller EVs along the power-law distribution. Signal for NTA and NFCM scales with radius to the 6th power, which is general for light scattering in the Rayleigh approximation, whereas signal scales for MRPS and SP-IRIS with radius to the 3rd power. Thus, because of finite dynamic range, NTA will be biased to detecting fewer of the small particles in a sample compared with MRPS. Of course, this will also depend on how the NTA the instrument and analysis settings are configured. One might over-expose the large particles in order to see the small ones, for example, or to increase sensitivity and maximize counts, but the outcome of this adjustment may be limited by glare from the large particles.
| Technique | Estimated biological particle size detection range (diameter) | Concentration Detection Range | Input volume (µl; may be a dilution) | Operation Time: Hands-on (HO)/Read time (R) (does not include optional labelling) | Instrument Cost | Costs for disposables (USD) | Performance | Quantification (across all particles, biological and synthetic) | Fluorescence (channels; for instruments we evaluated) |
|-----------|-------------------------------------------------------------|-------------------------------|-------------------------------------|--------------------------------------------------------------------------------|----------------|-----------------------------|------------|-------------------------------------------------|-----------------------------------------------|
| SP-IRIS   | ~50 nm to ~200 nm (label-free imaging)                      | n/a depends on capture Ab, but usually >> 1 x 10^7 particle/ml preferred | 35                                  | Day 1: 10 min sample prep (HO) Day 2: 2 h (HO); 10 min to 30 min per sample (R) | $$            | $$$; Chip/antibody kits (starting at $50/sample to >$100 for custom; ordered from manufacturer) | ++         | N/A (unless a universal capture agent is available) | Up to 3                                           |
| NTA       | ~70 nm to several hundred nm                                | 5 x 10^6 to 1 x 10^8 particle/ml (in PBS) | 1000 (slightly smaller volumes are possible in our experience) | Optional 20 min calibration (HO/P); 5 min to 15 min per sample (HO/R) | $$            | $: <1 per sample: 1 ml syringes (~$25 per 100, multiple suppliers; Ab not included) | +          | +                                               | Up to 4                                           |
| MRPS      | ~50 nm to 2 µm (cartridge-dependent)                        | 1 x 10^7 to 1 x 10^10 particle/ml (in PBS/PBST) | 5                                   | Optional 20 min calibration (HO/R); 15 min to 30 min per sample (HO/R) | $             | $$: Cartridges ($8 to $12 each, ordered from manufacturer) | ++         | +++                                            | N/A: No fluorescence capability                 |
| NFCM      | ~40 nm to ~200 nm                                           | 1 x 10^7 to 1 x 10^9 particle/ml (in PBS) | 20                                  | 20 min to 120 min calibration (HO/P); 5 min per sample (HO/P) | $$$$          | $: <1 per sample: 0.5 ml Tubes (~$25 per 1000, multiple suppliers; Ab not included) | ++         | +++                                            | Up to 2 (more now available)                   |
4.2 Is it important to resolve different particle size populations?

SP-IRIS, MRPS, and NFCM could resolve up to four populations of synthetic nanoparticles with different diameters. We note that distinct populations were somewhat obscured when MRPS results were averaged for SS, but not for PS—see also Supplementary Figure 6—which may reflect aggregation of the SS due to the electrolyte solution (PBS) required for MRPS and the convolution of experimental uncertainties in particle concentration and size measurements. Alternatively, as suggested by an astute reviewer, a PS standard could be run later on the same sample for scaling of size and concentration. Also noteworthy is that the NFCM platform distinguished subpopulations of SS particles quite well, but that this is likely because the same beads are used to calibrate the instrument. While detecting the expected concentration of high refractive index PS particles, NTA was unable to resolve individual particle populations and instead characterized the SS particles as a broad population distribution centred on an ‘average’ size. To be sure, it may be possible to resolve discretely sized particle populations using NTA with mixtures at different ratios of sizes. We could not do so with the mixtures we used. Whether this matters for biological particles is unclear. It does not seem that biological samples would contain unique EV subpopulations with exquisitely defined sizes, except perhaps for samples from sources infected with specific enveloped viruses. NTA does seem to be capable of detecting shifts in population distributions, and this capability might be more important for biological particles than resolving subpopulations.

4.3 On counting by SP-IRIS/fluorescence

In this study, neither SP-IRIS label-free measurements nor subsequent fluorescence detection could be used directly to estimate overall particle concentration. Instead, SP-IRIS is best used to understand ratios within populations and for single-particle phenotyping. Only a subset of EVs bind to any given affinity reagent ‘spot’. Binding is determined by diffusion (which is slow for EVs), presence and density of recognized surface markers, and affinity characteristics of antibody-to-antigen binding. The bound population of particles remaining after wash steps is only a small proportion of the total in the input material and cannot be used to determine overall concentration. Interestingly, fluorescence results often indicated higher particle concentrations than returned by label-free counting, even though particles positive for a particular antigen are expected to be only a subset of the captured population (different antigens) or to approach equality (if the capture antigen is targeted and antigen is abundant). Counts are higher because fluorescence detection is more sensitive than label-free. That is, fluorescence detects positive particles that may be below the limit of label-free detection. Also of note, capturing EVs onto the chip via surface markers may render those markers less available to subsequent binding by fluorescently-labelled antibodies. For example, in Figure 6a and b, CD81-captured particles that also display CD63 appear to have more CD63 available for binding by fluorescent antibodies compared with CD63+ particles that have had at least some portion of their CD63 sequestered by the surface-bound antibodies.

4.4 Did any platforms identify differences between EV separation technologies?

For both biological sources of EVs, we used two methods of EV separation: dUC (100K EVs), which has been the most common method for EV separation (Gardiner et al., 2016; Royo et al., 2020; Théry et al., 2006), and a combination of filtration and size exclusion chromatography (SEC EVs) (Böing et al., 2014). According to some evidence in the literature, dUC leads to more protein contamination and aggregation and damage of EVs (Arab et al., 2019; Linares et al., 2015; Van Deun et al., 2014). It should be noted that alternative viewpoints can also be found (Tian et al., 2020). However, protein particle contamination might be expected to introduce more and smaller particles. This outcome is indeed observed based upon TEM background and particle profile shifts towards smaller particles for several of the platforms. On the other hand, evidence of aggregation by dUC is not apparent in the data presented here. We cannot rule out aggregation, however, only that the techniques used here did not appear to detect it; we also do not wish to put too fine a point on these comparisons, which are based on limited data.

4.4.1 Single-particle phenotyping

For the three techniques with single-particle phenotyping capabilities (SP-IRIS, NTA, and NFCM), each has advantages and challenges. SP-IRIS was able to achieve true ‘multiplexed’ detection, in that signal could be obtained above background for up to three fluorescent channels. At the time of our evaluations, the NTA platform we used could not perform simultaneous multi-channel measurements and thus was not a true single-particle multiplexing platform. Instead, sequential filter switches were required, such that the same particles could not be tracked in different channels. Finally, although the nano-flow technology may be capable of multiplexed phenotyping, we did not explore this capability here.
4.5 Summary of findings: Table 2

In Table 2, we attempt to summarize our findings and views about the four investigated techniques. **Detectable size ranges** for biological particles: these should be considered to be rough estimates. If we accept the assumption that EVs follow a power-law size distribution (the smaller, the more abundant, with lower bounds defined by membrane curvature constraints), then no evaluated platform effectively detects the very smallest particles. However, SP-IRIS, MRPS, and NFCM appear to detect slightly smaller particles than NTA under the conditions and settings we tested. For NTA, MRPS, and NFCM, linear ranges for particle concentration for all instruments begin around $1 \times 10^7$ particles/ml (or slightly lower) and extend from about one order of magnitude (NTA) to multiple orders of magnitude (MRPS). This spread is important, since the wider the range, the fewer time-consuming concentrations or dilutions must be done to place an unknown particle population into the measurable range. SP-IRIS is a special case, since particles are captured by affinity, and overall concentration cannot easily be estimated. In our hands, particle concentrations must be high ($> 1 \times 10^7$ particles/ml) even for abundant antigens. Furthermore, the optimal captured particle counts are roughly several thousand per antibody spot (although this may vary). To hit a tight ‘sweet spot’, trial dilutions may be needed. Furthermore, the optimal dilution may differ for different antibodies on the chip because of different percentages of EVs positive for a particular antigen, per-EV antigen abundance, and antibody performance. Hence, dilutions are usually most important and potentially time-consuming for SP-IRIS. Related to dilution is the volume of input material required for a single reading. Assuming each platform is provided with a suspension at $1 \times 10^7$ particles/ml, the required volume of a dilution at this concentration ranges from $5 \mu l$ (MRPS) to around 1 ml (NTA). Of course, the actual volume/number of EVs needed will also depend on the number of concentrations/dilutions required to reach the measurable concentration range. The input volume difference is also inconsequential for highly abundant materials, but may be important for low-abundance EV samples. **If done, optional calibration steps** are rapid for NTA and MRPS (around 20 min). For NFCM, we find that calibration can be as short as 20 min but can sometimes take longer. Time for sample dilutions is most difficult to estimate, but is expected to correlate inversely with the range of measurement for each platform. **Read time** ranges from 5 min to about half an hour per sample. Note that the times we indicate are for sizing and counting only. Optional fluorescence measurements for the relevant platforms would in some cases add processing time for antibody incubations and removal, as well as for read times (except for NFCM). For SP-IRIS, we should also note that, although the total hands-on and read time is longer than for other techniques, each reading includes on-chip replicates, multiple capture antibodies, and up to three fluorescence readouts per capture antibody.

**Costs** for the platforms include initial outlay, disposable costs, and maintenance costs. For acquisition, the MRPS system is most economical, while the NFCM platform is the most expensive. For basic counting and sizing, operating costs for NTA and NFCM are negligible. Adding optional fluorescence increases these costs by amounts that are antibody-dependent. The MRPS system uses disposable cartridges that currently cost USD 8 to USD 12 each. The SP-IRIS platform has the highest disposable costs, with each sample requiring at least one chip that costs from USD 50 to > USD 100 each. Since optimal dilutions must be made and may be different for different capture materials on the same chip, multiple chips may be needed for the same sample. Chips also cannot be chemically stripped and re-used, at least not in our hands (Mallick and Witwer, unpublished data). Shelf-life of the chips is also a consideration. However, the company’s development of chips with extended shelf-life may overcome this potential hindrance. As noted, though, under optimal conditions, the platform provides multi-dimensional information that may justify costs and logistical challenges. We should also mention that chips for the SP-IRIS and MRPS instruments are currently available only from the instrument manufacturer for that particular measurement technique. As for maintenance costs, we are unable to estimate them at this time for any platform.

There are several limitations of our study as well as questions that might be investigated in the future. Since several of the platforms easily distinguished multiple populations of synthetic particles, but did not always identify the expected size for each population, normalization strategies including spiked-in standards could be useful. However, a question we have not addressed here is how components of the suspension medium might affect spiked-in synthetic material. For example, if synthetic beads are spiked into biological fluids, will they acquire ‘coronas’ that change their measured properties? Another question that might be addressed with these platforms is the effect of freeze-thaw and other storage conditions on EV integrity and membrane protein topology. Our study, by examining only several distinct sizes of synthetic particles, also does not rigorously define the range of size detection for biological particles for each platform. Likewise, our estimates of optimal range of concentration measurement for each instrument may be overly simplistic. In theory, a ‘single particle’ detection instrument is capable of using a single particle as a substrate, although measurement noise, contaminants, and the time required to ‘find’ the single particle are real-world considerations that might challenge this search.

**In conclusion:**

- For any platform and configuration, particle counting is accurate only within a certain range. Sensitivity for particles of different sizes and refractive indices should be considered. Recall that signal for light scattering methods like NTA and NFCM scales with radius to the sixth power, while signal for SP-IRIS and MRPS scales with radius to the third power. However, in our hands, the NFCM platform is more sensitive than NTA for small and low refractive index particles.
Different size populations within a mixture of synthetic nanoparticles can be identified by SP-IRIS, MRPS, and NFCM, but not, in our hands, by NTA. The individual sizes are not always accurately assigned, however, emphasizing the importance of calibration.

SP-IRIS, NTA, and NFCM offer fluorescent particle phenotyping, while MRPS does not. Multiplexed biological particle phenotyping of tetraspanins was easily achieved with the SP-IRIS platform (one-antibody capture and up to three-antibody fluorescence detection).

Appropriate reference materials are needed for better evaluation of single particle phenotyping capabilities, including multiplexed phenotyping.

Rather than relying on a single platform, consider using orthogonal technologies.

Both acquisition and recurring costs should be considered before choosing a platform.

No evaluated platform is necessarily ‘better’ or ‘worse’ than others; rather, it is important to be aware of the capabilities of each platform with respect to each particle population of interest and the population attributes that are of greatest interest.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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