Flow cytometry is a powerful method, which is widely used for high-throughput quantitative and qualitative analysis of cells. However, its straightforward applicability for extracellular vesicles (EVs) and mainly exosomes is hampered by several challenges, reflecting mostly the small size of these vesicles (exosomes: \(\sim 80-200\) nm, microvesicles: \(\sim 200-1,000\) nm), their polydispersity, and low refractive index. The current best and most widely used protocol for beads-free flow cytometry of exosomes uses ultracentrifugation (UC) coupled with floatation in sucrose gradient for their isolation, labeling with lipophilic dye PKH67 and antibodies, and an optimized version of commercial high-end cytometer for analysis. However, this approach requires an experienced flow cytometer operator capable of manual hardware adjustments and calibration of the cytometer. Here, we provide a novel and fast approach for quantification and characterization of both exosomes and microvesicles isolated from cell culture media as well as from more complex human samples (ascites of ovarian cancer patients) suitable for multiuser labs by using a flow cytometer especially designed for small particles, which can be used without adjustments prior to data acquisition. EVs can be fluorescently labeled with protein- (Carboxyfluoresceinsuccinimidyl ester, CFSE) and/or lipid- (FM) specific dyes, without the necessity of removing the unbound fluorescent dye by UC, which further facilitates and speeds up the characterization of microvesicles and exosomes using flow cytometry. In addition, double labeling with protein- and lipid-specific dyes enables separation of EVs from common contaminants of EV preparations, such as protein aggregates or micelles formed by unbound lipophilic styryl dyes, thus not leading to overestimation of EV numbers. Moreover, our protocol is compatible with antibody labeling using fluorescently conjugated primary antibodies. The presented methodology opens the possibility for routine quantification and characterization of EVs from various sources. Finally, it has the potential to bring a desired level of control into routine experiments and non-specialized labs, thanks to its simple bead-based standardization.

**Keywords:** exosomes; microvesicles; extracellular vesicles; quantification; flow cytometry; fluorescent labeling; CFSE; lipophilic styryl dye; ascites

*Correspondence to: Vitezslav Bryja, Faculty of Science, Institute of Experimental Biology, Masaryk University, Kamenice 753/5, building A36, CZ-625 00 Brno, Czech Republic, Email: bryja@sci.muni.cz

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RNA, protein) [reviewed in (2,3)]. EVs, and particularly exs, are an important, yet only recently appreciated, mode of intercellular communication (4) and thus have attracted growing interest of the basic researcher community. Moreover, they have also gained lots of attention from clinical researches as increasingly recognized high-potential biomarkers of several diseases including cancer (5), as well as possible therapeutic vehicles (6).

The exosomal field suffers from the lack of an easy, fast, and reliable method for quantification and characterization of these nanosized vesicles on a single particle level. Currently, characterization of exs is mainly based on the level of total amount of proteins and/or lipids in bulk isolates from biological samples. However, the methods used for bulk isolates, such as mass spectrometry or western blotting (WB), are rigid because they cannot dissect, whether the observed difference in protein/lipid abundance reflects the change in number of exs or in their composition mostly due to the absence of invariant “household” exs markers (7). Recent developments of nanoparticle tracking analysis (NTA) (8) allow for enumerating and also sizing of individual exs on the basis of Brownian motion. However, this light scattering–based approach also harbors several inherent disadvantages, for example, the ultimate need for multiple measurements of samples at different dilutions and different settings due to low dynamic range of the camera, which does not cover the range of 5 order of magnitude difference between the light scattered by smallest and largest EVs (9). Moreover, NTA technology is nowadays often used to detect only size and not to discriminate the composition of the particles (such as protein aggregates, micelles and lipoprotein particles, or abundance of a particular protein) thus often leading to overestimation of EV numbers. These potential errors in enumeration of EVs can only be partially limited by standardization of NTA measurements (10).

To ease the flow cytometry (FC; in further text also refers to “flow cytometer,” that is, instrument used in flow cytometry) analyses, exs are often bound to beads that provide larger surface and more scattered light. However, this approach is inconvenient for subsequent use of exs for functional studies. Moreover, the detection is largely dependent on the abundance/availability of antigen on the exs, which is recognized by the antibody coupled to the beads reflecting both pro et contra associated with high specificity of antibodies (11). Currently, the only reliable protocol available for free exs and direct FC analysis is a protocol for multiparameter FC analysis developed by research group of Prof. Marca Wauben (7,12). It allows for a high-resolution quantitative and qualitative analysis of individual cell-derived vesicles using an optimized configuration of the commercially available high-end FC (BD Influx). This extremely useful and exemplary protocol, though, demands an experienced FC operator capable of manual hardware adaptations with adjustments and calibration of the FC before use. Moreover, the exs labeling part of the experimental procedure, which yields multiple highly purified populations of exs, is also rather time and labor extensive and therefore as such may be discouraging for users in clinical labs or labs that do not highly specialize in the exs research.

Here, we present a simplified protocol for FC analysis of EVs using UC followed by a sucrose cushion purification step and simple fluorescent labeling step without additional washes requiring prolonged UC. The FC characterization is performed using a dedicated FC, where no adjustments prior to measurements are needed. By term dedicated FC [adopted from (9)], we specifically refer to Apogee A50 Micro (Apogee Flow Systems, Hertfordshire, UK), a FC specially developed for analysis of small particles (9,13,14).

Materials and methods

Cell culture and labeling by CFSE

HEK293 cells were cultured in DMEM supplemented with 10% FBS, 2 mM L-glutamine, 50 units/ml penicillin, and 50 units/ml streptomycin. Kuramochi and Ovsaho cell lines were cultured in RPMI 1640 supplemented with 10% FBS, 2 mM L-glutamine, 50 units/ml penicillin, and 50 units/ml streptomycin. For isolation of exs, cells were harvested with 0.05% trypsin/EDTA, washed twice in PBS and plated at 50% confluence onto fresh plates into RPMI 1640 or DMEM supplemented with vesicle-free 10% FBS, 2 mM L-glutamine, 50 units/ml penicillin, and 50 units/ml streptomycin. Supernatants were collected 24–48 h later.

Vesicle-free medium was prepared as described in Current Protocol in Cell Biology (15). In brief, medium was supplemented by 20% FBS, 4 mM L-glutamine, 100 units/ml penicillin, and 100 units/ml streptomycin and centrifuged at 100,000 × g, 4°C in SW28 rotor overnight (16–18 h). Supernatant was filter sterilized using 0.22 μm PVDF filter and 1 time diluted with DMEM or RPMI 1640 to reach the 10% FBS, 2 mM L-glutamine, 50 units/ml penicillin, 50 units/ml streptomycin concentrations. To harvest fluorescently labeled EVs, CFSE (Carboxyfluorescein succinimidyl ester, e-Bioscience) (or DMSO as diluent control) was added to final concentrations of 2–20 μM into vesicle-free cell culture medium at the time of seeding of the cells, which was typically onto 150 mm dish in 35 ml of culture media.

Ascites collection

Primary ascites of ovarian cancer patients were collected at the Department of Obstetrics and Gynecology, Faculty Hospital Brno, Czech Republic, under the informed consent of patients and IRB protocol of Vitezslav Bryja.
Isolation of cells and EVs

Exs and MVs were isolated according to slightly modified Current Protocol in Cell Biology (15). A scheme of the isolation protocol is given in Fig. 2. In detail, cell culture media or ascites were collected and transferred to 50-ml centrifugation tubes. Live cells were pelleted upon 5 min centrifugation at 200 × g at 4°C (designated P 0.2 – P, pellet, 0.2 – relative centrifugation force (RCF) in thousands of g), P 0.2 fraction from ascites usually contains red blood cells (RBC), which were removed by washing with ACK lysis buffer (150 mM NH₄Cl, 10 mM KHCO₃, 1 mM EDTA) and subsequent pelleting. The RBC removal step was repeated several times if necessary until both the pellet and the supernatant were clear (without the red traces of contaminating hemoglobin). Supernatants after live cell pelleting (S 0.2) were transferred to new 50-ml centrifugation tubes and centrifuged at 1,500 × g at 4°C for 10 min to remove cell debris (P 1.5). S 1.5 was transferred to 38-ml thinwall polycarbonate SW28 UC tubes (Beckman Coulter) and centrifuged at 14,000 × g (8,800 rpm for SW28 rotor – Beckman Coulter) for 1:10 h at 4°C to isolate MVs (P 14).

The P 14 was resuspended in filtered (0.22 mm PVDF filter) PBS and the centrifugation step was repeated to wash MVs from non-specifically adhered proteins. S 14 was optionally filtered using 0.22 μm PVDF filter to remove any contaminating MVs (larger than 220 nm) and the S 14 filtrate or S 14 was transferred to fresh 38-ml thinwall polycarbonate SW28 UC tubes and centrifuged for 100,000 × g (23,500 rpm for SW28 rotor – Beckman Coulter) at 4°C for 3:10 h using Optima XPN 90K Ultracentrifuge (Beckman Coulter). Sucrose cushion (4 ml) (30% sucrose in 20 mM Tris pH 7.6 in D₂O, filter sterilized) was pipetted into new SW28 tubes and centrifuged at 100,000 × g at 4°C (23,500 rpm for SW28 rotor) at 4°C in the dark. Specifically, for measurements in triplicate, 500-μl samples were prepared by diluting the isolated exs in 495 μl of filtered PBS and supplementing it with 5 μl of 10 to 1,000 μg/ml FM dye (or HBSS as a diluent control), prepared according to manufacturer’s instructions. After its cleavage of acetate groups by esterases (16), which are present in cells as well as in secreted EVs, CFSE has a peak excitation of 494 nm and peak emission of 521 nm (17), which is ideally measured using 488-nm (blue) laser excitation and 535/35 band pass filter for detection. Alternatively, for fluorescent labeling of exs lipids, the isolated exs were incubated in 100 ng/ml to 10 μg/ml FM 1-43FX or FM 4-64FX (Life Technologies) for 10 min at 37°C in the dark.

Labeling of exs with antibodies

Isolated exs were labeled according to (7). In short, isolated exs were incubated for 60 min with 10 μl of mouse anti CD63-PE (MEM-259, Exbio, IP-343). Prior to data acquisition, the samples were diluted with 480 μl 0.22 μm filtered PBS (final dilution 1:25). PE conjugated antibody of unknown specificity (MOPC-21, Exbio, IP-632) was used as negative IgG1 isotype control for background fluorescence. For double labeling with CFSE and anti EpCAM antibody, exs were incubated for 45 min at 37°C with anti EpCAM-APC (e-Bioscience, 17-5791-80) or isotype control (e-Bioscience, 17-4321-81) antibody and 1 μM CFSE.

Dedicated FC instrument settings

Data were obtained using 2 individual Apogee A50/Micro flow FCs equipped with 50 mW 405-nm (violet), 488-nm (blue) and 638-nm (red) lasers. Parameters in the Control panel were set to sheath pressure of 150 mbar and number of flush cycles to 4. Peak is a preferred parameter to area for light scatters (LSs) and fluorescence channels, as it corresponds better to submicron particles than the area. Particles with size several times smaller than the focused laser beam are being exposed to the light continuously across their path through the beam. Thus, the area of the pulse corresponds more to the length of the path (or the exposure time), than to the size of the particle. The peak value of the pulse should remain consistent across the particle path through the focused laser beam and should only reflect the scattering properties of the particle, with small or no regard to the length of the path (or exposure time) (18).
On each day of measurements, the FC performance was first verified using a reference bead mix. The reference bead mix (ApogeeMix, Apogee Flow Systems, cat#1493) composed of a mixture of plastic spheres with diameters of 180 nm, 240 nm, 300 nm, 590 nm, 880 nm, and 1,300 nm with a refractive index (RI) of 1.42, and 110-nm and 500-nm green fluorescent (excited by blue laser) beads with RI of 1.59 (latex) were used. These beads were used to assess the FC’s light scattering and fluorescence performance (both sensitivity and resolution). This can easily and quickly inform the user, whether the FC at current settings is capable of quality measurements of EVs. These beads were run several times during the acquisitions and at the end of the day to control the quality of recorded data among samples and stable FC performance during all measurements. Reference bead mix was used to set the PMT voltages and the thresholds for light scattering. Furthermore, as 2 populations of these beads are fluorescent in green channel, they were used to set the PMT voltages and the thresholds for fluorescent threshold triggering in parallel. All measurements were performed in log mode. The noise levels in PMT panel were kept below 1 at all parameters, ideally below 0.5.

Use of 405-nm laser is preferable for determination of light scattering, as it results in more scattered light and thus better resolution and sensitivity at light scattering detectors than 488-nm laser. However, as we used fluorescence threshold triggering in the blue part of spectrum, we measured the samples with both 405-nm and 488-nm lasers on. Importantly, we have not recorded photobleaching of the fluorescent particles with 405-nm laser when we compared the cytograms recorded with 405 nm, 488 nm, or both lasers. It is highly advisable to check regularly on Live camera window, whether the stream of particles is directed to the middle of laser beam. This indicates that there is no air bubble stuck in the tubing. In case of distorted particle stream, extensive cleaning can be applied to the tubing by selecting Clean valve button in the Control panel. If cleaning the valve does not correct the stream flow, it is preferable to replace the sheath fluid (dH₂O water).

**FC data acquisition**

After FC calibration by reference bead mix, the FC tubing was washed by 10% bleach to remove any particles which might have had adhered within the tubing. This wash was performed at sample flow rate 10.5–15 μl/min with 130-μl sample volume for 120 s. To remove the bleach, further wash was performed similarly with 0.22 μm filtered PBS. Such bleach and PBS washes were included after every positive sample, to exclude carryover of fluorescently positive events (EVs/fluorescent dye) between samples. The cleanness of tubing was verified by subsequent acquisition of filtered PBS sample at sample flow rate 0.75 or 1.5 μl/min with 130-μl sample volume for 120 s. This very low sample flow rate 0.75 or 1.5 μl/min (0.75 μl/min is the minimum) was used for all measurements and the time of acquisition was held constant for all samples, at least 120 s (or until the data buffer was full – up to 5,000,000 events), to yield enough events. Samples (500-μl) were just enough for measurements of 130-μl per sample in triplicates, leaving approximately 20-μl sample volume for data acquisition (more than 10 min at flow rate 1.5 μl/min), as roughly 110 μl encompass the dead volume of tubing of the FC.

**Western blotting**

Parental cells were lysed in NP40 lysis buffer (50 mM Tris pH 7.4, 150 mM NaCl, 1 mM EDTA, 0.5% NP40) supplied with 1 × Complete protease inhibitor mix (Roche), and 1 × Phosphatase inhibitor cocktail (Calbiochem). Lysates were spun down (16,000 × g, 10 min, 4°C). The total protein concentration was determined using DC™ Protein Assay (Bio-Rad). The protein lysates and isolated exs were mixed with 2 × Laemmli buffer and separated according to their molecular mass on 10–15% SDS-PAGE and transferred to Immobilon-P Membrane (Millipore). WB and antibody detection was performed as described previously (19). The antibodies used were rabbit anti Alix (gift from Pascale Zimmermann), mouse anti CD63 (MEM-259, Exbio, 11-343), mouse anti Golgin A1 (Invitrogen, A21270), mouse anti Hsc70 (Santa Cruz, sc-24), and rabbit anti Tsg101 (Sigma Aldrich, HPA006161).

**TEM and immunogold labeling**

For negative contrasting 10-μl drops of exs in PBS were adsorbed at activated formvar membrane coated HF35Cu with carbon EM grids (Pyser –SGI Limited) for 15 min, stained with 2% ammonium molybdate for 20 s at RT, washed with dH₂O water and imaged using transmission electron microscopy (TEM) Morgagni 268D (FEI) equipped with Mega ViewIII (Soft Imaging System) at 70 kV.

For immunogold labeling, isolated exs were mixed 1:1 with freshly prepared 8% paraformaldehyde and fixed for 10 min at RT. Then they were resuspended in 5 ml PBS and ultracentrifuged onto activated formvar membrane coated HF35Cu with carbon EM grids (Pyser –SGI Limited) placed at the bottom of the SW55Ti rotor UC tubes (Beckman Coulter) for 1 h at 100,000 × g, 4°C. Grids were blocked by 0.5% BSA + 0.1% Tween 20 in PBS for 30 min, incubated with primary antibody anti CD63 (MEM-259, Exbio, 11-343, dilution 1:200 in 0.5% BSA in PBS) overnight at 4°C, washed 3 times for 15 min with 0.5% BSA in PBS, then incubated with secondary anti mouse antibody conjugated with 10-nm gold particles (G7652, Sigma Aldrich, dilution 1:40 in PBS) for 2 h at RT, washed 3 times for 15 min in PBS and once with dH₂O. After staining with 2% ammonium molybdate for 20 s at RT, grids were washed with dH₂O water and imaged using
TEM Morgagni 268D (FEI) equipped with Mega ViewIII (Soft Imaging System) at 70 kV.

Statistical analysis
Graph visualization, data transformation, linear regression analysis, and concentration calculations were performed using Prism (version 5.0, GraphPad Software) and Microsoft Office Excel (version 2010, Microsoft).

Results and discussion

Challenges of FC of EVs and the ways to overcome them
We have decided to use FC to precisely quantify and characterize exs and MVs. FC is a powerful method for the high-throughput analysis of small particles which, however, has its limitations. Specifically, there are 2 main issues, which complicate FC-based analysis of EVs. First, the size of the vesicles ranges from 80 up to more than 1,000 nm, with the majority formed by exs, which are smaller than ~200 nm. The vast majority of conventional FC has the minimal detection limit for light scattering in a range of 200–500 nm (20). In addition, the conventional FC cannot discriminate accurately between particles that differ by less than 100–200 nm in diameter (21). Second, LS-based detection of EVs smaller than 200–300 nm is severely compromised by overlapping noise derived from optics, electronics, and buffers used.

To overcome these limitations we used a FC, which is optimized for the detection of small particles. In the further text we refer to this instrument as the “dedicated FC.” Conventional FCs use forward scatter (FSC), which detects the amount of light scattered at small angles (0.5–15°) and side scatter (SSC), which detects the light scattered at the angle of 90°. FSC is commonly interpreted as an estimate of size for objects larger than 500 nm, for example, cells, whereas SSC is used as a measure of their internal complexity (18). The dedicated FC builds on the fact that objects with sizes close to the observing light wavelength, such as exs, scatter significantly more light at larger angles (15–150°) due to diffraction (18,22). Thus, the dedicated FC uses alternative LSs such as SALS (small angle light scatter), MALS (middle angle light scatter), and LALS (large angle light scatter) for determination of light scattering. SALS, MALS, and LALS can be used for relative particle sizing and the latter also for internal complexity of the particles to limited extent.

Dedicated FC can thus, in principle, distinguish particles with the optical properties and size similar to exs and thus MVs as well. Our testing of the reference bead mix confirmed that dedicated FC is suitable for small particle measurements giving exceptional small particle sensitivity (approximately 100-nm latex beads) and resolution (difference of latex bead size in tens of nanometers) (Fig. 1a). Silica beads were used as a size reference because they have a RI of approximately 1.42 which is closer to the RI of biological particles [approximately 1.4 for cells (23) and 1.40±0.02 for EVs (9)] than latex particles [RI of 1.59 (24)].

The reference bead mix used in our experiments is a mixture of fluorescent and non-fluorescent spheres with diameters from 110 to 1,300 nm. To assess the suitability of individual LS detectors for detection of small particles, we have used the reference bead mix to assess the coefficient of variation (CV) of the SALS, MALS, and LALS detectors (reflecting the size of particles). With the manufacturer’s default settings, we did not observe CV >10%; actually in most cases CV was below 3%. At the default manufacturer’s setting, the best resolution and sensitivity for smallest particles was obtained on SALS (Fig. 1a). However, it should be noted that the device settings can be fine-tuned by experienced FC operators for best performance at LALS and/or MALS and yield better resolution and sensitivity for the expected exs populations (we have used this setting in Fig. 6c). Nevertheless, all other measurements in this study (shown in Figs. 1, 3–5, 6d) were obtained using the default settings to better illustrate the situation in non-specialized labs.

Reduction of noise
Data shown in Fig. 1a suggest that dedicated FC is able to detect particles with the properties – size and RI – very similar to MVs and exs. The detection of EVs by FC is, however, in parallel hampered by high level of noise. In the next step, we thus decided to identify and subsequently eliminate factors, which increase the noise and may interfere with the interpretation of the analysis.

First, we found it critical to acquire the data at a low flow rate (0.75 or 1.5 μl/min) for good LS resolution as well as for the accurate discrimination between non-fluorescent noise and fluorescent signals. This is likely because higher flow rates increase the diameter of the sample stream and allow the particles to move outside the center of the laser spot, yielding lower intensity of light scattering and fluorescent signals and increasing the CVs accordingly.

Second, we tested the effect of 0.22 μm filtration of PBS, which is used to dissolve the EV samples. As shown in Fig. 1b, use of filtered PBS in all steps of FC measurements greatly – 5.8 times – reduces the number of small events recorded by the FC (Fig. 1b). In the next experiments we took advantage of filtered PBS and used this solution during the calibration of the instrument for the fluorescent application in a way that the event rate was not higher than 2 events/μl for 0.22 μm filtered PBS in fluorescently positive region (quadrants A and B) (Fig. 1b).

Third, we have attempted to reduce the noise by gating events of interest using combination of 2 LSs. When using the reference bead mix, it is possible to gate on the beads
[region of interest 1 (ROI 1)] in the SALS versus LALS (Peak-Peak) cytogram and quantify the number of un-specific events in the next SALS versus 535/35 cytogram (Fig. 1b; middle panels – no gating; right panels – ROI 1 gated). We assumed that analysis of EVs may profit from such gating because of the bead size and RI resembling exs. Note that ROI 1 spreads to the noise region as well, to include any potential exs which scatter less light than 110-nm beads. However, reduction of noise was negligible (Fig. 1b) and on the opposite side restriction of events to ROI 1 significantly reduced the number of detected exs in subsequent experiments. Therefore, we present the data of ungated populations in following Figs. 3–6.

**Purification of exs from cell culture media and patient ascites for FC analysis**

The results shown in Fig. 1a suggested that dedicated FC represents a method with the potential to directly, rapidly,
and precisely quantify exs and other EVs. In the next step we decided to validate this assumption on real samples and to implement this step into possibly routine workflow for the isolation and analysis of exs and MVs.

As the first step in the workflow, we have chosen a suitable purification method – that is, a method which is relatively fast and unbiased. As such we decided to use a protocol based on UC. UC is still regarded as the gold standard for isolation of exs for basic research (25).

This is despite the increasing availability of other methods because UC-based methods collect in an unbiased way the exs, which may vary morphologically, biochemically, and functionally. Separation of exs by sedimentation only (downward displacement) during UC is not sufficient, although it is still used by many labs. Separation of exs from other particulate material can be guaranteed only by floatation (upward displacement) (26). Upward displacement can be performed either as an isopycnic

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Fig. 2 Isolation of microvesicles (MVs) and exosomes (exs) by ultracentrifugation (UC) and sucrose cushion. A schematic representation of protocol used to isolate extracellular vesicles (EVs). Black boxes describe the purpose of each centrifugation step, blue boxes contain additional instructions/explanation. Optional filtration step is indicated by italics. Green boxes depict fraction after centrifugation at relative centrifugation force (RCF) indicated by number (in thousands $\times$ g). P, pellet; S, supernatant.
Fig. 3. (Continued)
centrifugation where floatation in sucrose gradient is used or as centrifugation into the sucrose cushion. Isopycnic centrifugation yields high purity exs, but is technically more demanding and more time consuming (16–22 h) (11). In comparison to isopycnic centrifugation, sucrose cushion has several advantages for use in our workflow: (a) it is rapid (only 70 min); (b) the 30% sucrose in D$_2$O has a density of 1.19–1.2103 g/ml, which enables it to concentrate the majority of the exs (typical buoyant density 1.10–1.19 g/ml) in single low-volume fraction and to increase purity of exs, due to the formation of a continuous minigradient between 1.10 and 1.19 g/ml at the interphase between cushion and PBS (27); and (c) given the density of D$_2$O is 1.11 g/ml, significantly less sucrose is required to reach the cushion density of 1.21 g/ml than if prepared in H$_2$O, thus reducing potential hyperosmotic stress for exs (27).

Based on the arguments above we have used the UC and sucrose cushion-based protocol (schematized in Fig. 2) to isolate EVs from several sources. We purified MVs and exs from CM of 3 different cell lines – HEK293, Kuramochi, and Ovsaho (last 2 derived from high-grade serous ovarian cancer) and from primary patient samples. We have chosen samples of malignant ascites, that is, excessive fluid accumulated in the peritoneum of patients with high-grade serous ovarian cancer (28), which contains both cells, including ovarian cancer cells and leukocytes, and numerous EVs of unknown function.

MVs and exs produced by cell lines and from primary human material can be easily detected by a dedicated FC.

To distinguish EVs from the noise, we first stained them with the fluorescent dyes. Two different green fluorescent dyes – protein-specific (CFSE) and lipid-specific (FM 1-43FX) – were tested. The initial comparison was performed on MVs and exs produced by HEK293 cells into the culture media. EVs, labeled fluorescently by CFSE during cell culture (HEK293 exs+CFSE in CM) or prior to measurements by CFSE (HEK293 exs+CFSE) or FM 1-43FX dye (HEK293 exs+FM 1-43FX) were isolated using protocol outlined in Fig. 2 and subsequently analyzed by dedicated FC (Fig. 3) and by TEM in parallel (Fig. 4a).

The gates corresponding to fluorescently labeled EVs (ROI 1 in Fig. 3a–b) were set based on the acquisition of the free dyes (unbound to EVs) at highest final concentrations used for labeling and non-labeled EVs. For 10 µg/ml FM 1-43FX measurement, we allowed a maximum of 6 events/µl in ROI 1 when ungated (Fig. 3b). Using this setting, 10 µM CFSE label yielded a maximum of 8 events/µl, 0.22 µm filtered PBS ≤1 event/µl and similar events/µl were recorded for non-fluorescent EVs (Fig. 3a). EVs are clearly larger than exs, as evidenced by SALS histogram (Fig. 3c). Moreover, when S 14 was filtered prior to 100,000 × g UC step to remove all residual MVs and aggregated exs (designated as −F), it resulted in decreased number of more scattering EVs (Fig. 3) but also into significant loss of EVs detected, when compared to exs isolated from parallel samples when the 0.22 µm filtration step of S14 was not performed (designated as −NF). Thus, we included the filtration only as an optional step into the purification protocol (Fig. 2). We have noted the difference in ROI 1 population characteristics when labeled by lipophilic dye (membrane surface labeling) in comparison with luminal labeling by protein-specific dye CFSE. The surface labeling results in seemingly larger EVs as they were more shifted in SALS than if post stained with CFSE (Fig. 3). CFSE+ exs show more discrete fluorescent population when labeled during the biogenesis of exs (HEK293 exs+CFSE in CM) than population labeled by CFSE post isolation (HEK293 exs+CFSE) (Fig. 3a). Importantly, fluorescent staining of MVs and exs from HEK293 CM did not affect exosomal morphology/size as assessed by TEM (Fig. 4a).

The morphological characteristics of EVs isolated from HEK293 cells with or without CFSE in culture medium were unchanged and sizes corresponded to the expected ~80–200 nm in diameter, whereas the MVs were much larger – hundreds of nm (Fig. 4a).

Next, we tested labeling of more complex exosomal mixtures isolated from human samples – malignant ascites of ovarian cancer patients. In Fig. 4b, we also demonstrate that these exs can be labeled by CFSE or FM 1-43FX dyes prior to FC data acquisition. Filtration with 0.22 µm filter resulted again in significant loss of recorded events; however, in such exs sample isolated from a complex mixture, not all fluorescent events with high SALS value could be eliminated by filtration, as these may also represent aggregates formed during labeling. In non-filtered samples, we observed contamination of the sample by

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Fig. 3. FC analysis of EVs isolated from cell culture media. (a) EVs were labeled by protein-specific (CFSE) fluorescent dye either during their genesis in cells (HEK293 exs+CFSE in conditioned media (CM)) – middle panel, or just prior to measurements (HEK293 exs+CFSE) – lower panel. The upper panel consists of negative control samples – CFSE-only, HEK293 exs-only, HEK293 MVs-only. (b) EVs were labeled by lipid-specific (FM 1-43FX) dye prior to measurements (HEK293 exs or MVs+FM 1-43FX). FM 1-43FX-only represents negative control sample. (a–b) SALS versus CFSE or FM 1-43FX (both green fluorescence) cytoagrams of ungated populations are plotted. The ROI statistics corresponds to fluorescent EVs. Note the difference between the appearance of exs population from the same source (HEK 293 cells) stained with different dyes and/or prior or after isolation. (c) Histograms of SALS of EVs from (a). Left, histogram of EVs labeled during cell culture. Middle, histogram of differentially labeled MVs isolated from HEK293 cells. Right, histogram of differentially labeled exs isolated from HEK293 cells. F; S14 filtered through 0.22 µm filter; NF, S14 non-filtered.
Fig. 4. Quantification of EVs by dedicated flow cytometer (FC). (a) Representative images of negatively contrasted exs and MVs from HEK293 cells (used in Fig. 3) visualized by TEM. The morphological characteristics are unchanged for EVs derived from HEK293 cells in the presence of 2 mM CFSE in the CM. (b) Exs isolated from malignant ascites of ovarian cancer patients labeled by CFSE (upper middle and right panel), FM 1-43FX dye (lower left panel) and unlabeled (upper left panel). Note the difference between filtered (-F) and non-filtered (-NF) exs preparation. (c) Representative images of negatively contrasted exs from (b) visualized by TEM. Exs preparations isolated from malignant ascites of ovarian cancer patient #1 are contaminated by lipoprotein particles (smaller than exs, black asterisks) and sporadic MVs (larger than exs, black arrows), the latter are detected by FC also in (b). (d) Serial 2-fold dilutions of reference bead mix were measured using fluorescence threshold triggering. The measured values in ROI 2 (Fig. 1a) corresponding to 110-nm particles, (roughly reflecting exs) are plotted as red circles along with the calculated input amount of beads (X). Linear regression of log transformed data reveals slope $-0.9954 \pm 0.0106$, $R^2 = 0.9952$. One representative experiment out of 3 is shown. (e) Quantification of exs from CM and patient samples. Left panel, event/µl for 10-fold dilutions of exs stained with CFSE in CM (indicated by squares), exs stained with CFSE prior to FC measurement (triangles) and exs stained with FM 1-43FX prior to FC measurement (spheres) were plotted against the dilutions. Slope/Y-intercept of linear regression analysis was calculated. Pooled slopes are presented for each source of exs indicated by color. Right panel, 10-fold dilutions of control samples (negative) indicate the background level of event/µl detected by FC.
lipoprotein particles and also MVs (for TEM image see Fig. 4c). This experiment, however, provides proof of the principle that our methodology can also easily be applied to the complex samples of biological fluids and adapted for clinical use. 

Dedicated FC represents a reliable method for the quantification of exs from different sources

Based on the results in Figs. 1, 3, and 4 we concluded that our simplified isolation and labeling protocol combined with the dedicated FC (a) can easily distinguish...
fluorescently labeled exs from non-fluorescent background, (b) does not affect the morphological characteristics of exs, and (c) is applicable for analysis of exs from both relatively uniform cell line CM as well as complex clinical samples. So far, however, we did not evaluate the suitability of our approach for the quantitative analysis. To address this point, we first performed serial 2-fold dilutions of the reference bead mix and plotted the measured values of 110-nm fluorescent beads (events in ROI 2 in Fig. 1a) together with the calculated input based on the specified concentration of beads in the original solution of reference bead mix. The number of beads in µl recorded in a fixed time frame (120 s) was plotted against the dilution factor of these beads and by linear regression we verified that the fluorescent microparticles of the approximate size of exs can be credibly quantified across wide range of concentrations (Fig. 4d). To determine the inter-sample variation, 3 individual tubes with identical reference bead mix dilutions were prepared and the mean ± standard error of the mean of 110-nm beads acquired within 120 s was calculated. The intra-sample variation was determined by quadruplicate measurements of the same beads dilution in 120-s time intervals. The inter- and intra-sample variations were relatively low, 7.0
and 1.4%, respectively, which demonstrates that dedicated FC has the potential to reliably quantify microparticles across several orders of magnitude.

Using serial dilutions, we also assessed performance of our protocol for exosomal preparations isolated from HEK293, other cell lines (Kuramochi and Ovsaho), and furthermore for the exs isolated from primary malignant ascites of 2 ovarian cancer patients. The results of this analysis are summarized in Fig. 4e. By linear regression and comparison of slopes and intercepts of the 2 or 3 lines of exs derived from the same source, we showed that numbers of recorded events/μl in dilutions corresponded well to the expected calculated ratios, as slopes of data from all individual samples were very close to the theoretically expected value -1.

This analysis also allowed us to compare individual labeling methods. First, the original amount of cells/CM and the dilution was the same for the exs labeled during the cell culture as was for the exs labeled prior to measurements. This indicates, that the short time labeling by CFSE prior to measurements (post-labeling) is as effective as the long-term cell labeling during their biogenesis in cell culture (pre-labeling). The prolonged exposure to CFSE, by which exs are believed to incorporate more CFSE, prior to and also after the release from cells, yielded equally brightly labeled vesicles as the typical CFSE labeling protocol, when the cells are labeled for 10 min in the tube and washed prior to seeding (data not shown). Second, the amount of events/μl recorded for exs + CFSE in CM were comparable to numbers of exs + CFSE and exs + FM 1-43FX with the maximal CV of 15.5% when different labeling approaches were compared. The fact that lipid-specific dye (FM 1-43FX) and protein-specific dye (CFSE) yield comparable numbers of fluorescently labeled exs isolated from CM of several cell lines (equal slopes and Y-intercepts) strongly indicates that the labeling is specific for exs, which contain both proteins and lipids. Slight differences between individual labeling techniques – with more exs consistently detected in CFSE-labeled samples (in comparison to FM 1-43FX) – were found in ascites-derived exs (CV of 33 and 22%). Such a discrepancy might be addressed by double labeling of EVs.

Thus, as a next step, we investigated whether we can double label the EVs with protein-specific and lipid-specific dyes. This approach would enable the separation of EVs from common contaminants of EV preparations, such as protein aggregates or micelles formed by unbound lipophilic styryl dyes, and explain the differences in EVs numbers obtained upon different labeling. We used CFSE and a variant of FM dye fluorescent in red spectrum – FM 4-64FX.

First, the EVs labeled during cell culture by CFSE were post-labeled with FM 4-64FX dye. Surprisingly, only 44% of CFSE + MVs were also positive for FM 4-64FX, whereas 63% of FM 4-64 + MVs were also positive for CFSE. Nevertheless, the total amount of double positive events recorded was the same regardless of the gating strategy – CFSE + first, FM 4-64FX + first, or double positive (Fig. 5a). We also noticed, that incubation of these CFSE pre-labeled EVs with FM 4-64FX dye greatly reduced their green fluorescence, resulting in non-detectable green fluorescence of HEK 293 exs (data not shown) and diminished green fluorescence of MVs, which was however still detectable and could be used for FC analysis (Fig. 5a). Thus, we post-labeled EVs with both CFSE and FM 4-64FX for 45 min at 37°C and showed, that in the case of exs isolated from malignant ascites, 56% of CFSE + exs were also positive for FM 4-64FX, whereas 54% of FM 4-64 + MVs were also positive for CFSE. Similarly as in Fig. 5a, the total amount of double positive events was the same regardless of the gating strategy – CFSE + first, FM 4-64FX + first, or double positive (Fig. 5b).

This suggests that single labeling with CFSE may lead to overestimation of EV numbers; however, this overestimation is in the range of maximally tens of %, which is not ideal, but still much less than has been reported for other methods, such as NTA, which may overestimate the numbers of EVs in range of orders of magnitude (9,10).

**Dedicated FC is suitable for exosome characterization by antibody labeling**

For exs research it is not only important to enumerate the vesicles on single particle level but it is of vital importance to describe their further characteristics such as quantitatively and qualitatively assessing their protein composition and possible changes in protein abundance under various conditions. FC is ideal for precise and multiparameter characterization of EVs by staining with antibodies and detection by a fluorochrome. As a proof of principle of the suitability of the dedicated FC for this type of assay, we decided to evaluate expression of typical exs markers, tetraspanin CD63 (29), and EpCAM (30) on the exs derived from human ascites.

First, we analyzed biochemical composition of the exosomal sample by WB. This analysis showed positivity for typical exosomal markers (Alix, Hsp70, Tsg101, CD63) and negativity for “antimarkers” (Golgin A1), which are present in cells of origin but absent from isolated exs (Fig. 6a) (15,31,32). To further confirm presence of CD63 in the exosomal fractions of human ascites, we used immunolabeling with gold particles and were able to detect CD63 on the surface of exs (Fig. 6b).

Knowing that the exs from malignant ascites are CD63-positive, we decided to quantify the proportion of CD63 + exosomes. We have used R-Phycoerythrin-(PE)-labeled primary antibody directed against CD63 antigen. PE and APC were chosen as fluorochromes due to their high quantum yields and extinction coefficients. Using this approach, we were able to show by the PE-labeled primary
antibody that exs isolated from malignant ascites of ovarian cancer patient #2 were CD63 positive (Fig. 6c). Based on the total amount of exs stained by CFSE in parallel tube, CD63+ exs from malignant ascites made 38% of all CFSE+ exs detected. Similarly, when we double stained the exs derived from ascites of patient #1 with CFSE and anti EpCAM antibody, we revealed that only 11% of CFSE+ exs were positive for the cell adhesion molecule EpCAM (Fig. 6d), which has also been widely reported as an exs marker (30). Importantly, taken into account that single labeling with CFSE may overestimate the numbers of EVs as much as by 79%, the actual proportion of CD63+ exs and EpCAM+ exs may be up to 68 and 20%, respectively. This observation confirms previously anticipated, though poorly studied, variability among individual exosomal types and is consistent with our observations made by TEM, when only a fraction of ascites-derived exs stained with antimarker antibodies (Fig. 6b).

Taken together, our data demonstrate that exs and MVs from variant sources can be isolated, enumerated, and characterized via a standardized protocol and dedicated FC. Our protocol provides a fast – 10–12 h (Fig. 6e) – way to purify, quantify, and characterize EVs for both experimental as well as diagnostic purposes.

Summary and conclusions
In the current study, we show that fluorescently labeled exs can be directly quantified and characterized using dedicated FC specifically designed for the analysis of small particles. The usefulness of dedicated FC for EV research was already demonstrated by others (13–15). Here we go further and present an analytical pipeline that does not require prior hardware adjustments of the device, which were needed for high-end conventional FC with the existing protocols (7,12). We show that EVs can be fluorescently labeled by combination of unspecific protein- and lipid-specific dyes and/or primary antibodies; staining can be performed either during cell culture or just prior to measurements. In the latter case, there is no need to remove the unbound fluorescent dye or fluorescently labeled antibody and after short incubation (5–45 min for dyes, 60 min if antibodies are used) data can be readily acquired. These properties allow implementing FC analysis directly into a routine, approximately 12 h workflow for purification and characterization of exs from both experimental and diagnostic samples.

We are aware of the fact that our approach is inferior to the existing exemplary FC protocol (7,12) in terms of purity and uniformity of obtained vesicular populations and their characteristics. Main advantages of our protocol are (a) its simplicity, (b) versatility, (c) minimal requirements on FC operator, and (d) short time required for analysis. As such it is suitable for and aimed at multiuser and clinical lab, where it has the potential to bring desired level of control to the experiments involving exs and other small EVs. We believe that the ever-growing exosomal basic research may benefit from our protocol and the use of dedicated FC.

Authors’ contributions
VP and VB conceived and designed the research. VP, ZD, AK, and KK isolated the EVs and characterized them by WB. VP and JS acquired the FC data. DK, LI, and AH performed the electron microscopy experiments. IC, LM, and VW collected and EJ pathologically assessed the ascites samples. VP and VB interpreted the data and wrote the manuscript. Each author reviewed and made critical comments to the manuscript.

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