



RESEARCH ARTICLE OPEN ACCESS

Control of Physical and Biochemical Parameters Influencing Exogeneous Cargo Protein Association to Extracellular Vesicles Using Lipid Anchors Enables High Loading and Effective Intracellular Delivery

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ABSTRACT

Despite biomolecule delivery is a natural function of extracellular vesicles (EVs), low loading of exogenous macromolecules such as proteins into EVs limits their interest as convincing protein delivery systems for health applications. In this context, lipid-anchorage of exogenous cargo into EV membrane recently emerged as a promising option to enable their vectorisation into cells. Nevertheless, this option was not explored for protein intracellular delivery, and further characterisation of critical parameters governing the association of a lipid-anchored cargo protein to EVs is still needed to confirm the relevance of this anchorage strategy. Therefore, we sought to identify these parameters in a precise and quantitative manner, using bulk and single nanoparticle analysis methods to identify protein loading capacity and subsequent intracellular delivery. We identified incubation temperature, cargo concentration, lipid anchor (LA) structure (lipid nature, linker) and EV origin as critical factors influencing maximal EV loading capacity. Precise control of these parameters enabled to load cargo protein close to EV saturation without hindering cellular delivery. The structural properties of LA influenced not only cargo protein/EV association but also intracellular delivery into different carcinoma cell lines. By thoroughly characterising Lipid-PEG-protein anchorage, this study evidences the interest of this tunable and controllable approach for efficient EV protein delivery.

1 | Introduction

Therapeutic proteins are powerful and fast-acting drugs that have proved their efficacy for many years in the treatment of a wide

range of pathologies such as cancers (Tanaka and Rabbitts 2008, Gulfidan et al. 2020, Ivanov et al. 2013), infectious or genetic diseases (HIV infection; Maes et al. 2012), cystic fibrosis (Naren et al. 1998, etc.) or auto-immune pathologies (Poluri et al. 2023,

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Andrews et al. 2022). By contrast to small molecules, proteins have pharmacological effects at very low concentration and are highly specific for their targets (Gurevich et al. 2014). As a consequence, development of therapeutic proteins has soared over the two last decades leading proteins such as antibodies to be top-tier drugs (Lu et al. 2020) and this trend is expected to continue (Ebrahimi and Samanta 2023). However, despite their very interesting intrinsic properties, proteins also struggle with important limitations hindering their therapeutic potential. Delivery is one of the main challenges to overcome, as these fragile macromolecules face numerous barriers before reaching their target, especially if intracellular delivery is sought. For molecules of this size, crossing the cell membrane is impossible, and no delivery system capable of fully resolving this problem has yet been developed. As a result, most of currently marketed therapeutic proteins target extracellular sites such as membrane proteins (Strohl 2018), limiting effective therapeutic applications.

Among promising options, natural nanoparticles such as extracellular vesicles (EVs) have been recently considered (Le Saux et al. 2021). EVs are produced by the totality of known cells and organisms and have the ability to vectorise biomolecules from nucleic acids to proteins, making them a highly attractive alternative to synthetic nanoparticle-based therapies (Herrmann et al. 2021). EVs have been successfully used to deliver several proteins (Wang et al. 2022, Zhang et al. 2020, Lainšček et al. 2018, Ilaibaks et al. 2023, Heath et al. 2019, Mizrak et al. 2013, Haney et al. 2019) relying either on endogenous (e.g., using cell engineering for protein fusion (Heath et al. 2019, Silva et al. 2021, Yim et al. 2016, Wang et al. 2018)) or exogenous (e.g., using isolated EV submitted to membrane permeabilisation processes (Haney et al. 2019, Lamichhane et al. 2015)) loading methods. Using endogenous methods, reported loading efficiencies are usually limited below 200 proteins/EV (Silva et al. 2021) and engineering efficiency (i.e., the proportion of EV loaded as compared to total EV population) is generally below 30% (Silva et al. 2021, Corso et al. 2019, Ivanova et al. 2023, Rankin-Turner et al. 2021). Moreover, these strategies imply time-consuming cell genetic manipulations and restrict the loading to certain EV sub-populations (e.g., CD63 positive EV). In contrast, exogenous methods rely on the physical destabilisation of EV membranes. These approaches have the disadvantage of producing protein aggregates that form during this harsh process and may be confused with EVs. Moreover, the volume occupied by EVs, even at high concentration ($<5.10^{11}$ EV/mL), is too low to enable an efficient loading only based on cargo diffusion into EVs. Finally, it is difficult to compare all these approaches due to the lack of characterisation and comparable metrics. Thus, the development and characterisation of efficient protein-loading approaches for EVs is urgently needed.

In this context, the addition of a lipid anchor (LA) to cargo is an appealing option to increase EV loading by exploiting hydrophobic interactions with the phospholipid EV membrane in mild conditions. Moreover, this strategy can be considered more versatile, especially compared to fusion with EV proteins, as all EV sub-populations are enclosed by a hydrophobic membrane. This approach has been successfully used for some types of cargos such as small chemicals (Jing et al. 2021, Zheng et al. 2023, Choi et al. 2019, Pitchaimani et al. 2023) or nucleic acids (Didiot et al. 2016, Haraszti et al. 2018, Stremersch et al. 2016,

Zhang et al. 2019, Gong et al. 2019). However, cargo proteins were barely reported and never for intracellular delivery and parameters able to influence cargo protein association to EVs remain to be investigated. Some studies suggested the impact of the lipid nature of LA but only for small cargos (Zheng et al. 2023, Biscans et al. 2018) and potential effects from other parameters (e.g., linker) were never investigated, especially for intracellular administration of a protein.

This study intends to characterise the LA-cargo protein association to EVs by using biochemical readouts, state-of-the-art single-particles analysis methods and algorithmic pixel-classifying analysis of confocal microscopy images. We assumed that control of fundamental parameters governing this loading could turn EVs into effective intracellular protein delivery systems whilst maintaining both protein and EV properties. We investigated the influence of physical parameters (cargo concentration and incubation temperature), and LA structure (lipid nature and PEG linker molecular weight). All these parameters appeared crucial for the maximal loading capacity of EVs, the stability of the association or the efficiency of intracellular delivery. By controlling this process, we obtained a very high loading capacity equal to ≈ 1000 cargo proteins per EVs. Our results, therefore, evidenced this strategy as a new interesting option for protein intracellular delivery, opening promising opportunities for various therapeutic applications.

2 | Materials and Methods

2.1 | Chemicals

Chemicals were purchased from (i) Biopharma PEG (USA): DSPE-PEG₁₀₀₀-SH, DSPE-PEG₂₀₀₀-SH, DSPE-PEG₅₀₀₀-SH, CLS-PEG₂₀₀₀-SH (structures are shown in Figure S1A); (ii) Sigma-Aldrich (France): sepharose CL-2B cross-linked, sulfuric acid 95%–97%; formic acid 98%–100%; acetonitrile 99%, trehalose, Tris-HCl; (iii) Thermofisher (France): EZ-Link Maleimide Activated Horseradish Peroxidase, ELISA 1-Step Ultra TMB, M-PER Mammalian Protein Extraction Reagent, Pierce Protease Inhibitor Mini Tablets, CellMask Plasma Membrane Stains, DAPI, ProLong Gold Antifade Mountant; (iv) Gibco Life Technologies (USA): Dulbecco's Modified Eagle Medium (DMEM), FBS, penicillin/streptomycin, L-glutamine, trypsin-EDTA, opti-MEM, Dulbecco's Phosphate-Buffered Saline (DPBS).

2.2 | Murine Mesenchymal Stem Cell Derived EVs (mMSC-EVs)

mMSC-EV were obtained from Everzom (Paris, France, products #230801-G5 and #230608-G12). Briefly, EVs from Murine C3H/10T1/2, Clone 8 (ATCC CCL-226) cells were produced and isolated using Everzom's optimised 3D biomanufacturing and isolation processes, followed by an additional purification step with Size Exclusion Chromatography (SEC; Captopore 700). The column volume was adapted to the protein concentration measured after Tangential Flow Filtration (TFF). A 1:10 addition of 250 mM trehalose (for a final concentration of 25 mM) was done before freezing the sample in cryotubes. For some additional experiments, a second EV product was used. These mMSC-

EV#2 were produced following a well-established protocol in our laboratory (Le Saux et al. 2020). The final EV pellet was resuspended in DPBS buffer containing 25 mM of trehalose and stored at -80°C .

2.3 | Synthesis of HRP Conjugates

2.3.1 | Synthesis of Lipid-PEG-HRP

Stock solutions of Lipid-PEG-SH (structures and conjugation process are described in Figure S1A,B) were obtained by dissolving 20 g/L of powder in ultrapure water. EZ-Link Maleimide Activated Horseradish Peroxidase (mal-HRP) was rehydrated in H_2O for a final concentration of 2 g/L before the addition of 20 molar equivalents of Lipid-PEG-SH (Lipid-Anchor abbreviated as LA). Depending on the LA MW and the final volume needed, 45–122 μL of lipid-PEG-SH were added to 500–1000 μL of mal-HRP. Water was added according to the volume of added LA to obtain identical final concentration in all mixes, equal to 7.8×10^{-4} M of Lipid-PEG-SH (1.4–4.5 g/L according to the MW of the LA) with 3.9×10^{-5} M (1.6 g/L) of mal-HRP. The mix was then incubated at 37°C overnight under gentle agitation. A dialysis step (Slide-A-Lyzer Dialysis Cassette 20,000 MWCO, 0.5 mL, Thermo Fisher Scientific) was performed against DPBS under gentle agitation to eliminate unconjugated Lipid-PEG-SH. The dialysis buffer was frequently changed (2 h at RT, 2 h at 30°C and finally 20 h at 4°C). MW of DSPE-PEG₁₀₀₀-HRP, DSPE-PEG₂₀₀₀-HRP, DSPE-PEG₅₀₀₀-HRP and CLS-PEG₂₀₀₀-HRP were observed using HPLC-MS (Figure S2C) and mass gains of respectively 1392, 2362, 4815 and 2154 g/mol as compared to unconjugated HRP were measured. Enzymatic assay revealed that Lipid-PEG-HRP retained 80%–95% of its biological activity regardless of the type of Lipid-PEG after conjugation and dialysis (Figure S2D).

2.3.2 | Synthesis of Fluorescent Lipid-PEG-HRP-A488

Alexa-Fluor 488 was conjugated with Lipid-PEG-HRP to obtain Lipid-PEG-HRP-A488 for single-particles analysis and cell internalisation microscopic observations using the Alexa Fluor 488 Microscale Protein Labelling Kit (Thermo Fisher Scientific) following the manufacturer's protocol. Fluorescent conjugates were purified using Zeba Dye and Biotin Removal Spin Columns (Thermo Fisher Scientific) and 50 mM of Tris-HCl were added to quench the reactivity of residual dye and avoid further artefactual conjugation of the dye with EV proteins.

2.4 | Lipid-PEG-HRP and EVs Association

5.10^{10} EV/mL (assessed by NanoTracking Analysis, NTA) were mixed with different final concentrations of Lipid-PEG-HRP (from 0.5 to 75 mg/mL). Depending on the requirements of the experiment, volumes of preparation were comprised between 50 μL and 1 mL. Samples were incubated overnight at 25°C or 37°C , depending on the experiment, under gentle agitation.

2.5 | Purification of EV~Lipid-PEG-HRP

SEC separation was used for all samples except in indicated microscopy experiments. Columns were packed following Tulkens et al. (2020) protocol. 50 μL of EV/HRP mixture at 5.10^{10} EV/mL were diluted 11-fold in DPBS and 500 μL of the sample were deposited on the top of the column. 500 μL elution fractions were collected and analysed (NTA, enzymatic activity). Depending on the experiment and the purity level needed, only the fourth fraction or the combination of third, fourth and fifth fractions was collected. Samples used for microscopy observations in Figure 7 were purified using tangential ultrafiltration with Amicon 100 kDa 0.5 mL (Merck Millipore) in order to avoid the dilution resulting from SEC. 500 μL of samples previously diluted by 10 in DPBS were deposited in an Amicon filter before centrifugation at $2000 \times g$ during 15 min at 4°C (Centrifuge 5804 R, Eppendorf). Then, retentates were diluted by 10 in DPBS once again to obtain 500 μL before a second identical centrifugation. Final EV-containing retentates were analysed using NTA and TMB enzymatic assay.

2.6 | Characterisation

2.6.1 | NanoTracking Analysis (NTA)

EVs were analysed with a NanoSight NS300 (Malvern Panalytical Instruments, UK). Samples were diluted with particle-free DPBS in order to obtain a concentration range between 1.10^8 and 5.10^8 particles/mL. Measurements were performed with a 405 and 488 nm laser. Data were collected using the NanoSight NTA 3.4 software following a tailored script: temperature was set at 25°C , syringe pump at 40 arbitrary units and 5 videos of 60s were recorded with a camera level set to 15 and analysed with a detection threshold set to 5. All groups of compared data were measured and analysed using identical conditions. Zetaview (Particle Metrix, USA) was used for ZP and particle fluorescence measurements. For ZP measurements, 2-cycle measurements on 2 positions (SL1 and SL2) using a 488 nm laser were used with a sensitivity set at 80 and a shutter at 100. Samples were diluted to obtain a final EV concentration equal to 5.10^7 EV/mL in 1:10 DPBS (1.5 mM). Sensed conductivity was equal to $145\mu\text{S}/\text{cm}$. For particle fluorescence measurement, two successive measurements were done in fluorescent and scatter mode. Fluorescence measurements were performed with a sensitivity of 90 and a shutter of 50 using a 488 nm laser and a 500 nm filter, whilst scatter measurements were carried out with a sensitivity of 80 and a shutter of 100 without filter. Concentration correction factor was calculated following the manufacturer's guidelines by using reference fluorescent beads YG-488 and was equal to 1.1. Data were analysed using Software ZetaView (8.05.16 SP3 version).

2.6.2 | HPLC-MS Experiments

ElectroSpray Ionisation Mass Spectrometry (ESI/MS) and MS/MS analyses were performed on a Synapt G2-S instrument equipped with Ultra Performance Liquid Chromatography (Waters Corporation, UK). For the separation, a gradient with H_2O containing 0.1% of formic acid (FA) as eluent A and acetonitrile containing 0.1% of FA as eluent B was applied as follow: from 95% A to

70% in 8 min and from 70% until 50% between 8 and 25 min before going back to 95% A in a total run of 35 min, at a flow rate of 0.45 mL/min on a polyphenyl column (BioResolve, 150 × 2.1 mm, 2.7 µm particle size), designed for intact proteins and antibodies analysis. UV detection was done at 220 and 280 nm. Mass spectrum acquisition was performed with a mass range of m/z 800–5000, a scan time of 0.5 s in a positive ion mode with capillary voltage 3 kV, 40°C and 350°C for the source and desolvation temperatures, respectively. MassLynx software was used for data acquisition and processing, with the MaxEnt1 module for the deconvolution process.

2.6.3 | Enzymatic Assay and Determination of EV-Associated HRP Amount

The amount of EV-associated HRP was measured by enzymatic activity using tetramethylbenzidine (TMB) after SEC separation. For each SEC-purified sample, an individual standard curve was generated using the pre-SEC sample (of known HRP concentration) to account for any possible variation due to the LA-HRP lot. 50 µL of sample were added to 50 µL of TMB and incubated 30 min at RT in 96-well plate (Nunc MaxiSorp, Invitrogen). Then, 50 µL of sulfuric acid (2 M) were added to stop reaction. Absorbance at 450 nm was then measured to quantify the HRP activity using a MultiScan Go (Thermo Fisher Scientific). The association capacity was calculated by dividing the HRP concentration (expressed as the number of proteins/mL) by the EV concentration (number of particles/mL) and is expressed as the number of HRP/EV. Association efficiency was defined as the proportion of HRP remaining with EVs after SEC as compared to the initial amount during incubation with EVs and before SEC.

2.6.4 | MACSPlex Assay

EV surface epitopes determination was performed using the MACSPlex Exosome Kit mouse (Miltenyi Biotec). The test was done according to the manufacturer's protocol. Briefly, 1.10^9 EVs were diluted in 120 µL MACSPlex Buffer. A detection cocktail [MACSPlex Exosome Capture Beads and Detection Reagent CD9, CD63 and CD81] was then added to the EV suspension and incubated for 1 h at RT under gentle agitation (450 rpm) protected from light. Capture beads contained 39 populations of dyed beads coated with monoclonal antibodies able to recognise 37 potential EV surface antigens along with 2 internal isotype negative controls. MACSPlex Exosome Detection reagent contained APC-labelled detection antibodies against the three tetraspanins CD9, CD63 and CD81. After the washing steps, samples were analysed using a BD Accuri C6 (BD Biosciences, USA (software version 1.0.34.1 Build 20180111.34.1)). For each sample, 7000–12,000 single-bead events were measured. Bead populations were gated based on their fluorescence intensity according to the manufacturer's guidelines. The obtained median fluorescence intensity (MFI) of 37 exosomal surface epitopes and two isotype controls were subtracted from the corresponding MFI of the blank. This blank was the EVs buffer diluted in the same way as EV samples, mixed with capture beads and detection reagent without EVs.

2.6.5 | NanoFlowCytometry (nanoFCM)

NanoFCM was used in 40–200 nm range mode. Alexa-Fluor 647 anti-mouse IgG1 (isotype control), APC anti-mouse CD63 Alexa-Fluor 647, APC anti-mouse CD9 Alexa-Fluor 647 and APC anti-mouse/rat CD81 Alexa-Fluor 647 (Biolegend) were diluted at 1:50,000 before incubation with the sample diluted at 5.10^8 EV/mL at 4°C overnight (ON).

2.6.6 | Cryo-Transmission Electron Microscopy (Cryo-TEM) Imaging

Samples were imaged using cryo-TEM. Carbon film coated EM grids (Ted Pella Inc.) were glow-discharged at 30 mA, for 30s (Pelco easiGlow, Ted Pella Inc.). 4 µL of each sample solution was applied to the carbon-coated side of the TEM grid, blotted for 10s, before plunge-freezing in liquid ethane using the Vitrobot Mark MkIII (FEI). Vitrified samples were then imaged in a Glacios cryo-transmission electron microscope (ThermoFisher) equipped with a K3 direct electron detector, SerialEM software (D. Mastronarde, Boulder Lab) was used in acquisition.

2.6.7 | Super-Resolution Microscopy Imaging

Single molecule localisation microscopy was performed using an ONI Nanoimager (ONI Bio) following manufacturer guidelines. Native EV samples were incubated with CD9 Alexa-Fluor 488, CD63 Alexa-Fluor 568 and CD81 Alexa-Fluor 647 monoclonal antibodies from the EV Profiler kit (ONI Bio). EV~LA-HRP samples were incubated with CD9 Alexa-Fluor 647 (Clone MZ3, BioLegend) and Vybrant DiI (Invitrogen, Thermo Fisher). Unbound antibodies and dyes were removed after overnight incubation using SEC (70 nm qEV, IZON). 10 µL droplet was added between a 1 mm TOMO adhesion glass slide (Avantor) and a #1 glass cover slip (Corning) for immediate imaging. Briefly, the instrument was calibrated for direct stochastic optical reconstruction microscopy with bead calibration (ONI Bio) slide using NimOS version 1.19.7. Three-color super-resolution microscopy was then performed with 500 frames taken per channel. Laser powers were adjusted according to company standard operating procedures. Raw image files were uploaded to the cloud software CODI via CODI Desktop Uploader version 0.30.0. Image analysis was performed on CODI. Briefly, drift correction, filtering, clustering and counting algorithms were performed using the predefined "EV Profiling" workflow.

2.7 | Cell Internalisation Experiments

2.7.1 | Quantitative Cell Lysis Experiments

1.10^5 PANC-1 (ATCC, CRL-1469), SKBR3 (ATCC, HTB-30) and A549 (ATCC, CCL-185) cells in 1 mL of DMEM supplemented with 10% FBS, 2 mM L-glutamine, 100U/mL penicillin and 100 µg/mL streptomycin were plated in 24-well plate and incubated at 37°C and 5% CO₂ prior to cell internalisation experiments. After 24 h, the medium was discarded and replaced by samples to be tested diluted in fresh identical medium for a total volume of 400 µL. Cells and samples were then incubated at 37°C with different

amounts of HRP (from 1 to 15 ng of HRP/well) for different time periods (2, 4 or 24 h) depending on the experiment. The sample-containing medium was afterwards removed, and cells were washed with cold DPBS 3 times before the addition of 200 μ L of M-PER supplemented with protease inhibitor and incubated 5 min at RT under gentle shaking. To ensure optimal lysis, cells were then pipetted in the well and collected in microtubes before a 10 s vortex. Lysates were finally centrifuged for 15 min at 4°C and $14,000 \times g$ to keep only the supernatant. HRP activity was measured with TMB following the protocol described in 5.6.3. except that all dilutions and standard curves were diluted in lysis buffer instead of DPBS to avoid any bias on the peroxidase activity from surfactants present in lysate. Blank was done with untreated cells control lysate.

2.7.2 | Microscopy Observations and Quantifications

A549 cells were seeded at 17,500 cells per well in 8-well Nunc Lab-Tek II Chamber Slide plates (ThermoFisher) and incubated for 24 h at 37°C in a humidified 5% v/v CO₂ atmosphere. Then, 150 μ L of samples diluted with 150 μ L of DMEM supplemented with 10% FBS, 2 mM L-glutamine, 100U/mL penicillin and 100 μ g/mL streptomycin were added and incubated for 5 h with cells. Cells were washed with DPBS, labelled with CellMask (1:1000 dilution in fresh medium) and incubated for 30 min. Finally, cells were fixed in DPBS with 4% formaldehyde (36% w/w) for 10 min at room temperature and washed with DPBS. DAPI (1 μ g/mL in DPBS) was added and left for 5 min before washing. The slide was mounted with Prolong mounting medium for microscopy observations. Images were acquired using a full-field epifluorescence microscope Nikon Ti2 (Nikon Instrument Inc), with a CMOS back-illuminated Prime95B (1200*1200 pixels, 11 μ m pixel, Teledyne Photometrics) camera, Lumencor system as a source of illumination and 40X objective (oil, 1.3 numerical aperture). Images were treated with Fiji (ImageJ) software with identical adjustment of brightness and contrast. The set parameters minimum/maximum displayed values were equal to 3000/62,000 for HRP-488, -140/17,500 for CellMask and 6800/34,500 for DAPI staining. For quantitative analysis, images were acquired on the same samples using a spinning-disk confocal microscope with a Yokogawa CSU-W1 spinning head (Yokogawa Electric Corporation) on a sCMOS back-illuminated Prime95B (1200*1200, 11 μ m pixel) camera and Nikon 60X objective (Plan Apo, 1.4 numerical aperture). Images were taken with a Z-step of 210 nm using Oxixus lasers (excitation at 488 and 561 nm). Image processing for quantitative analysis was performed using Fiji. Cells were segmented into a binary mask using the random-forest pixel-classifier algorithm (Labkit ImageJ plugin) to attribute a class ("inside cell" or "background") to every pixel. Spot detection was done by applying a Laplacian of Gaussian filter before local minima detection (using FeatureJ Log filter and MorphoLibJ local minima detector) to extract a 3D point per spot. The list of coordinates generated based on CellMask were used to differentiate cells from the background in order to count HRP-A488 fluorescent spots inside cells. This number of internalised fluorescent spots was normalised per the total number of intracellular voxels in order to normalise the difference in cell number per image.

2.8 | Statistical Analysis

RStudio version 1.4.1717 (2009-2021 RStudio, PBC) was used for post hoc statistical analysis. Unilateral or two-tailed Welch tests were used depending on the null hypothesis as indicated in figure legends.

3 | Results

3.1 | Characterisation and Tunability of the Association of Lipid-PEG-Cargo to EVs

3.1.1 | Cargo Protein Concentration and Incubation Temperature Are Key Parameters Controlling Maximal Association to EVs and Enable High Loading Efficiency

A commercial source of EVs (from Everzom company) produced by murine mesenchymal stem cells (mMSC) and extracted by tangential flow filtration followed by a steric exclusion chromatography (SEC) step was characterised (Figure S1). These EVs exhibited expected EV features (i.e., morphology, size and antigen markers). Horse Radish Peroxidase (HRP) was chosen as model cargo protein for its representative size (Hydrodynamic Radius \approx 3 nm and 45 kDa) and extremely accurate enzymatic readout, enabling us to validate the maintenance of protein activity during this process. Thus, Lipid-PEG-HRP (namely DSPE-PEG₁₀₀₀-HRP, DSPE-PEG₂₀₀₀-HRP, DSPE-PEG₅₀₀₀-HRP and CLS-PEG₂₀₀₀-HRP) were obtained after the reaction of maleimide-functionalised HRP and Lipid-PEG-SH anchors as described in Section 5.3.1 and summarised in Figure S2B. Lipid-PEG-HRP conjugates were characterised by HPLC-MS (Figure S2C). The conservation of enzymatic activity of HRP was controlled after conjugation (Figure S2D). SEC was used to remove all non-associated cargo protein from samples and to measure accurate association capacity (Figure 1A). Fractions 3, 4 and 5 were identified as EV-containing fractions ($13 \pm 3\%$, $28 \pm 4\%$ and $16 \pm 2\%$, respectively) whilst a negligible amount of free Lipid-PEG-HRP was found in these fractions (0%, 0% and $2 \pm 1.4\%$, respectively). Thus, the F4 fraction was selected to ensure that no free lipid-PEG-HRP could contaminate samples and lead to an overestimation of the association calculation.

Finally, two negative controls were added in association experiments to ensure that the retention of HRP with EVs in SEC fractions was the result of their association and not an artifact due to HRP and Lipid-PEG aggregates: (1) unconjugated HRP with EVs and (2) Lipid-PEG-CH₃ + HRP with EVs. For this last condition, the N-terminal methoxy function (CH₃) does not allow the covalent attachment of Lipid-PEG to mal-HRP.

We first investigated the influence of HRP concentration and incubation temperature to identify optimal association conditions. The first experiment set was done with a fixed EV concentration (5.10^{10} EV/mL) incubated with various concentrations of DSPE-PEG₂₀₀₀-HRP (from 0.5 to 75 μ g/mL) at 25°C (Figure 1B). After SEC separation, association capacity was firstly calculated based on the assumption of a homogeneous distribution of HRP among EVs in fraction F4. Association capacities of controls remained constantly below 50 HRP/EV without significant variation between 0.5 and 30 μ g/mL. Above this concentration,

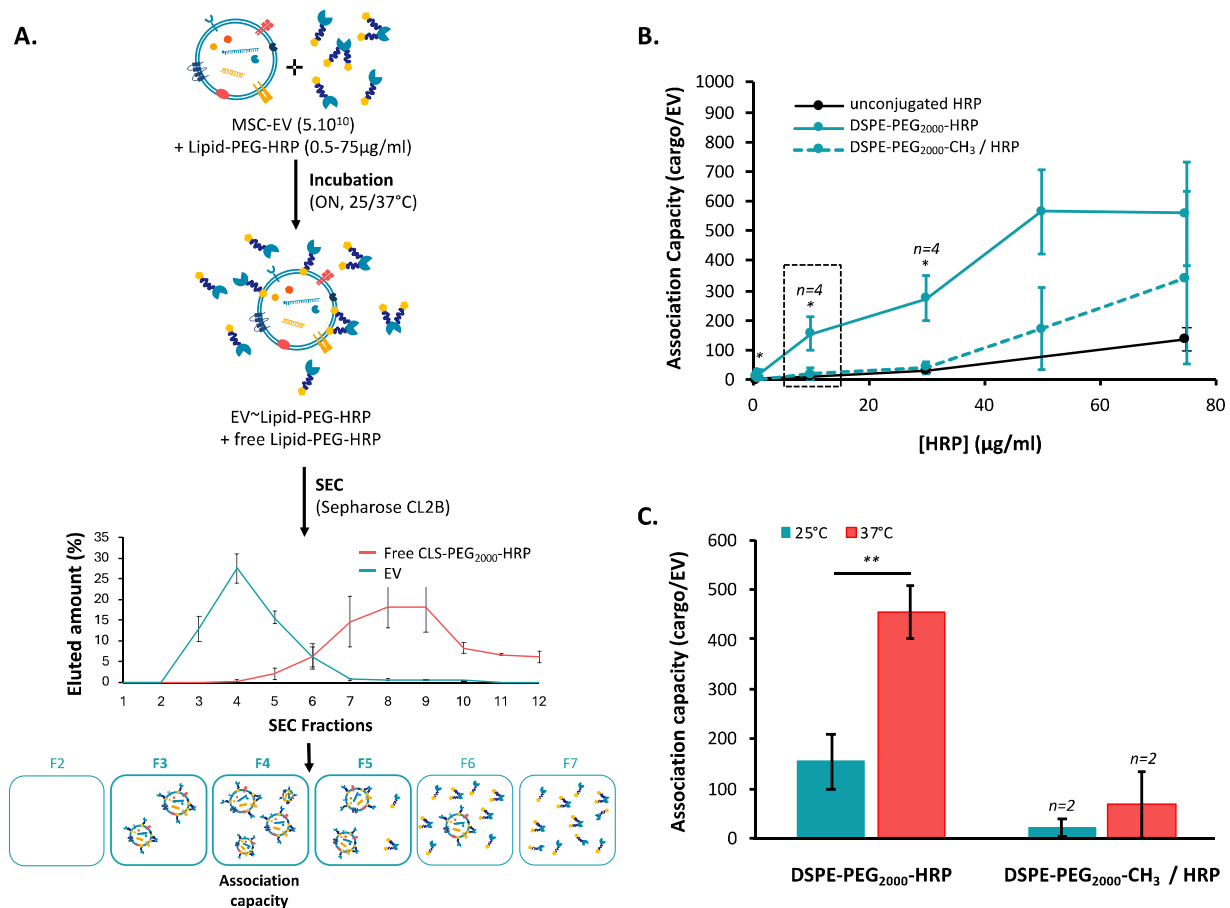


FIGURE 1 | Influence of Lipid-PEG-cargo concentration and temperature on the association with EVs. (A) Schematic representation of EV/Lipid-PEG-HRP association workflow (association, separation), not to scale. The fourth fraction of SEC was selected to calculate association capacity. (B) Number of biologically active Lipid-PEG-HRP per EV versus HRP concentration after incubation of 5.10^{10} EV/mL with unconjugated HRP, unconjugated DSPE-PEG₂₀₀₀-CH₃/HRP, and DSPE-PEG₂₀₀₀-HRP at 25°C overnight. $n = 3$ if unspecified. (C) Number of biologically active HRP conjugated per EV versus temperature during incubation of 5.10^{10} EV/mL with $10 \mu\text{g/mL}$ of DSPE-PEG₂₀₀₀-CH₃/HRP or conjugated DSPE-PEG₂₀₀₀-HRP at 25°C or 37°C overnight. p values were calculated using a two-tailed Welch test.

association capacities started to increase, reaching 341 ± 289 and 134 ± 39 HRP/EV at $75 \mu\text{g/mL}$, respectively. In contrast, DSPE-PEG₂₀₀₀-HRP association to EVs started from 8 ± 1 HRP/EV at $0.5 \mu\text{g/mL}$, and increased progressively to 21 ± 10 HRP/EV at $1 \mu\text{g/mL}$, 154 ± 56 at $10 \mu\text{g/mL}$ and 272 ± 76 at $30 \mu\text{g/mL}$ to finally reach a plateau of 564 ± 144 HRP/EV at $50 \mu\text{g/mL}$. The number of HRP associated with EVs was significantly higher than controls when HRP was conjugated with DSPE-PEG₂₀₀₀ for each tested concentration (excepted at $75 \mu\text{g/mL}$, probably due to SEC column saturation).

Then, we evaluated the influence of incubation temperature (25°C or 37°C) at $10 \mu\text{g/mL}$ of DSPE-PEG₂₀₀₀-HRP (Figure 1C), leading to a rise in association from 154 ± 56 to 456 ± 53 HRP/EV, whilst no significant increase was observed in the control. Noticeably, this value of 456 ± 53 HRP/EV was close to the EV saturation previously observed for higher DSPE-PEG₂₀₀₀-HRP concentrations ($50 \mu\text{g/mL}$) at 25°C . These results suggest that an increase in EV membrane fluidity facilitates LA-cargo insertion. This incubation at 37°C was not deleterious for EVs

based on mode, concentration and cryoTEM observations (Figure S3). Therefore, association conditions were set to a Lipid-PEG-HRP concentration of $10 \mu\text{g/mL}$ incubated at 37°C overnight with 5.10^{10} EV/mL for subsequent experiments.

3.1.2 | Lipid-PEG Anchor Structure Influences the Cargo Protein Association to EVs

We compared three different MW of PEG linkers for the same LA (DSPE) with 1000, 2000 and 5000 g/mol (Figure 2A). Similarly, cholesterol-PEG₂₀₀₀ (abbreviated in CLS-PEG₂₀₀₀) was compared to DSPE-PEG₂₀₀₀ (Figure 2B). DSPE-PEG-HRP conjugates were efficiently associated with EVs without significant differences depending on the PEG MW, with an association capacity of 373 ± 110 , 255 ± 57 and 329 ± 86 HRP/EV using PEG 1000, 2000 and 5000 g/mol, respectively. Surprisingly, an association of 898 ± 247 HRP/EV was obtained for CLS-PEG₂₀₀₀-HRP, above the EV surface saturation observed with DSPE-PEG₂₀₀₀-HRP in the same conditions ($10 \mu\text{g/mL}$ of LA-HRP incubated at 37°C overnight).

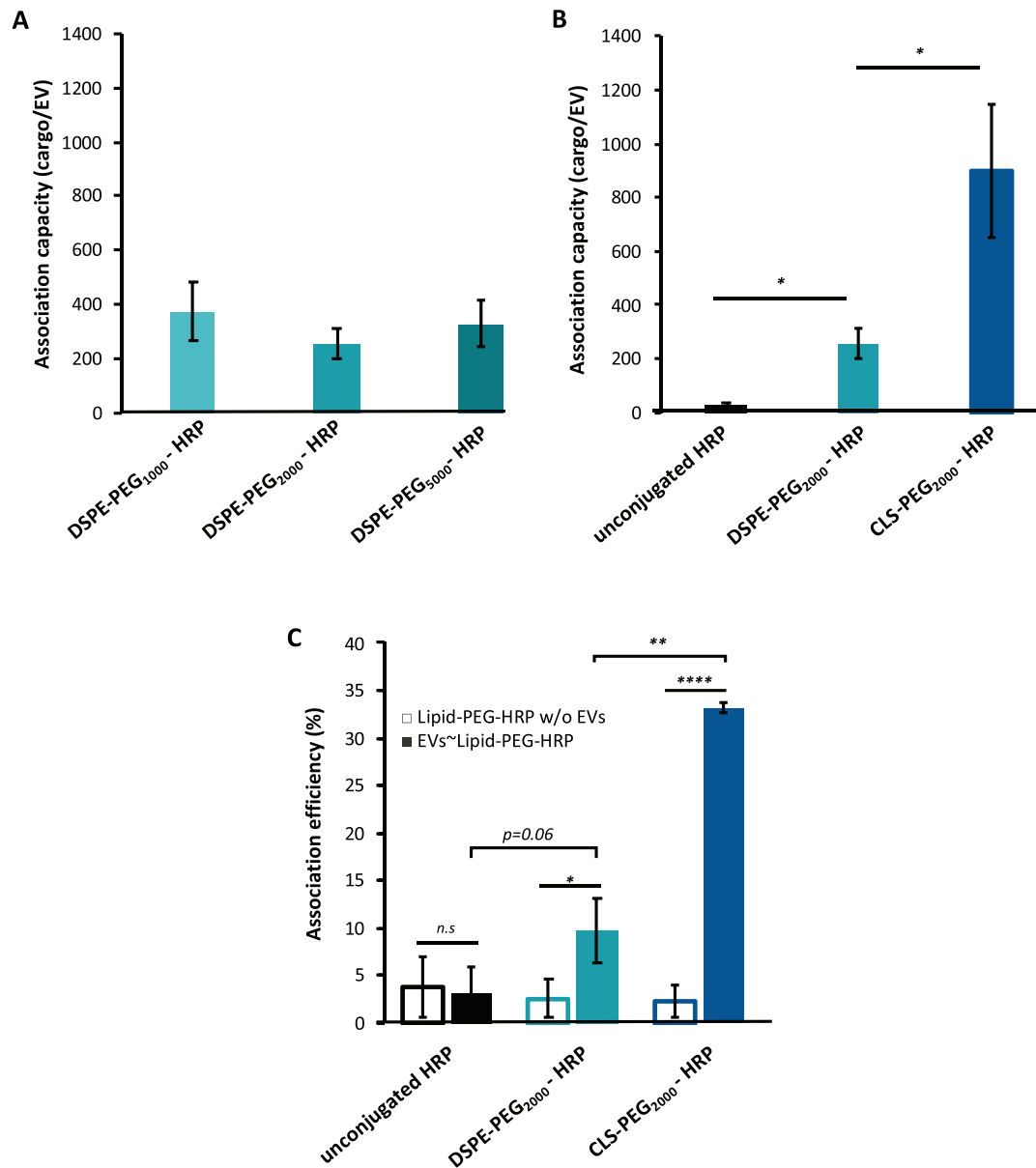


FIGURE 2 | Influence of PEG linker MW, lipid anchor nature and EV type on the association between EVs and Lipid-PEG-HRP. Association efficiency according to (A) the PEG MW of DSPE-PEG anchor or (B) the lipid nature of Lipid-PEG₂₀₀₀ anchor after incubation of 5.10^{10} EV/mL with $10 \mu\text{g/mL}$ of HRP incubated overnight at 37°C . Fourth SEC fraction was considered to calculate this association capacity ($n = 3$). (C) Loading efficiency is defined as percentage of HRP Lipid-PEG-HRP retained with EVs (all EV SEC fractions were considered) after incubation of $10 \mu\text{g/mL}$ of unconjugated HRP, conjugated DSPE-PEG₂₀₀₀-HRP and conjugated CLS-PEG₂₀₀₀-HRP with ($n = 3$) 5.10^{10} EV/mL or without ($n = 4$) EV incubated overnight at 37°C . p values were calculated using a two-tailed Welch test. $n.s > 0.05$, $* < 0.05$, $** < 0.01$, $*** < 0.001$, $**** < 0.0001$.

This high loading capacity conferred to this process an important association efficiency (i.e., the percentage of total available cargo that has been associated with EVs compared to the total amount introduced into the incubation media [Rankin-Turner et al. 2021]). After an incubation of $10 \mu\text{g/mL}$ of unconjugated HRP, DSPE-PEG₂₀₀₀-HRP or CLS-PEG₂₀₀₀-HRP with EVs, association efficiencies of $3.1 \pm 2.9\%$, $9.7 \pm 3.3\%$ and $33.2 \pm 0.5\%$ were obtained, respectively (Figure 2C). By considering that only 60% are collected after SEC, the corrected loading efficiency is superior to 50% using CLS. Therefore, we selected CLS-PEG₂₀₀₀ as a reference Lipid-PEG anchor for subsequent experiments and characterisation.

3.1.3 | Single-Particle Analysis Unveils EV~CLS-PEG₂₀₀₀-Cargo Association Heterogeneity

We fluorescently labelled HRP for single-particle analysis measurements and co-localisation observations in order to have an overview of the repartition of CLS-PEG₂₀₀₀-cargo among EVs. Unconjugated HRP and CLS-PEG₂₀₀₀-HRP were conjugated with Alexa-Fluor 488 TFP ester to obtain unconjugated HRP-A488 and CLS-PEG₂₀₀₀-HRP-A488. NTA analysis using Zetaview-Twin showed no fluorescent particles for EVs incubated with unconjugated HRP-A488, whilst $59 \pm 14\%$ of detected particles were fluorescent in EV~CLS-PEG₂₀₀₀-HRP-A488 samples (Figure 3A).

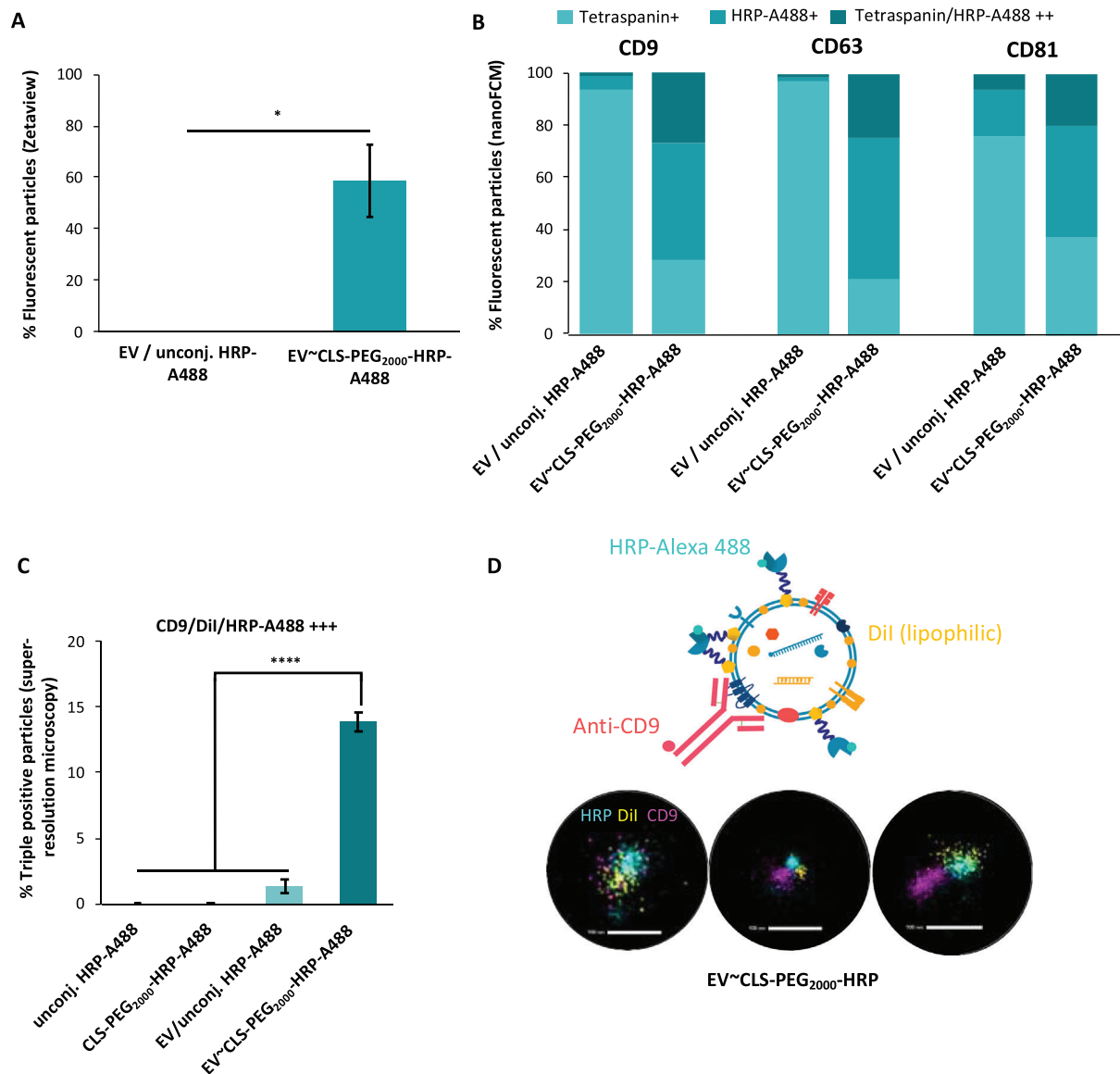


FIGURE 3 | Heterogeneity in CLS-PEG₂₀₀₀-HRP-A488 loading. (A) Proportion of fluorescent detected particles using Zetaview ($n = 3$). (B) NanoFCM analysis after incubation with anti-CD9, anti-CD63, or anti-CD81 antibodies coupled with AlexaFluor647 ($n = 1$). Only fluorescent particles have been considered. (C) Proportion of triple positive clusters using STORM. Samples were incubated with DiI dye and anti-CD9 antibody, whilst HRP was labelled with Alexa-Fluor 488 ($n = 3$). (D) Representative images of triple positive clusters observed using STORM. p values were calculated using a two-tailed Welch test. $n.s > 0.05$, * <0.05 , ** <0.01 , *** <0.001 , **** <0.0001 .

Then, NanoFlowCytometry (nanoFCM) analysis was carried out using Alexa647-coupled antibodies anti-CD9, -CD63 and -CD81, three EV markers (Figure 3B, details are provided in Figure S4). We did not purify samples by SEC to reach an EV concentration compatible with nanoFCM sensitivity. No A488+ particles were detected in the control buffer containing CLS-PEG₂₀₀₀-HRP-A488 alone (Figure S4), demonstrating that no fluorescent CLS-PEG₂₀₀₀-HRP-A488 aggregates were present. This control rules out the possibility that such aggregates of HRP could be responsible for positive artifacts in EV samples. Overall, 72%, 79% and 63% of fluorescent particles were A488+ in EV~CLS-PEG₂₀₀₀-HRP-A488 after incubation with anti-CD9, -CD63 and -CD81 antibodies, respectively. These values were 7%, 2% and 24% for EV/unconjugated HRP-A488. 27%, 24% and 20% of fluorescent particles were double positives for A488 and

respectively CD9, CD63 and CD81 in EVs~CLS-PEG₂₀₀₀-HRP-A488 sample in contrast to EV/unconjugated HRP-A488 showing respectively 1%, 1% and 6% of double positive EVs.

Finally, we used super-resolution microscopy (STORM) observations on EVs labelled with DiI lipophilic dye and anti-CD9 antibody (Figure 3C,D). This last technology enabled the observation of the colocalisation of HRP with EVs, both qualitatively and quantitatively. In contrast to controls, $14 \pm 1\%$ of EV~CLS-PEG₂₀₀₀-HRP-A488 particles were triple positive. Details in number of cluster types are provided in Figure S5.

We also tested a second EV product to investigate if this relatively low proportion of triple-positive EV was due to the EV origin and to control the versatility of this approach. These EVs were

produced from mMSC under 2D-culture conditions and purified by ultracentrifugation (characterisation is shown in Figure S6). $42 \pm 2\%$ of these EVs were triple positive after association with CLS-PEG₂₀₀₀-HRP-A488 (Figure S7). This could be due to differences in membrane composition and tetraspanin expression. Overall, these results highlight the versatility of this approach to associate a protein with different EVs and demonstrate a high engineering efficiency approximatively equal to 60%.

3.1.4 | Lipid-PEG-Cargo Association Does Not Modify EV Size and Surface Characteristics

Then, we investigated the potential impact of LA conjugation on the physico-chemical properties of EVs since proteins are large cargos that could disturb EV membranes. EV~CLS-PEG₂₀₀₀-HRP were analysed by NTA for size (Figure 4A) and Zeta Potential (ZP, Figure 4B). We did not observe significant changes neither in mode (110 ± 4 nm for EV~CLS-PEG₂₀₀₀-HRP and 112 ± 5 nm for control) nor in negative ZP (-20 ± 1 mV for EV~CLS-PEG₂₀₀₀-HRP and -19 ± 1 mV for control) after association. We compared EV/unconjugated HRP and EV~CLS-PEG₂₀₀₀-HRP using MACSPlex assay to evaluate the impact of the association on EVs antigens accessibility. Interestingly, EV~CLS-PEG₂₀₀₀-HRP and control exhibited similar profiles (Figure 4C). Thirty-three EV surface epitopes assessed by the test were detected in both types of EVs, highlighting the preservation of the accessibility of EV surface antigens using this association method. CryoTEM imaging revealed no difference between EV samples associated or not with CLS-PEG₂₀₀₀-HRP (Figure 4D).

3.1.5 | Lipid-PEG-Cargo Anchorage to EVs Demonstrates a Good Stability in the Presence of Serum

A common reported issue using LA conjugation to EVs is the redistribution of the cargo to plasma components as described in some studies with small cargos (Zheng et al. 2023, Tréton et al. 2023). For comparison purpose, we carried out a similar stability study on EV~Lipid-PEG₂₀₀₀-HRP with DSPE and CLS lipid. For the DSPE lipid (Figure 5A), $107 \pm 34\%$, $94 \pm 13\%$, $106 \pm 15\%$ and $65 \pm 3\%$ of the associated protein remained with EVs after 4°C ON in DPBS, a freeze/thaw (F/T) in DPBS, 37°C during 2 h in DPBS or DPBS supplemented with Foetal Bovine Serum (FBS, 1:50), respectively. For CLS lipid (Figure 5B), these values were $75 \pm 16\%$, $159 \pm 52\%$, $83 \pm 14\%$ and $71 \pm 1\%$, respectively. Overall, these results showed a good stability of the anchorage with EVs despite some partial redistribution of Lipid-PEG-HRP to FBS components that seems to be impacted by the lipid nature of LA.

3.2 | Control of EV/LA-Protein Association Process Enables Its Efficient and Functional Delivery Into Carcinoma Cell Lines

3.2.1 | EV Association Enables Concentration-, Time- and Cell-Dependent Intracellular Delivery of Active Cargo Protein Into PANC-1, A549 and SKBR3 Cells

After optimising LA-protein cargo association to EVs to obtain high asso, we aimed at demonstrating that this strategy was able to

turn EVs into effective protein delivery systems by characterising the intracellular delivery with an absolute amount of vectorised cargo. EV~Lipid-PEG-HRP samples were incubated with 1.10^5 PANC-1 cells (adherent epithelioid carcinoma cell line derived from human pancreas) per well. Cells were subsequently washed before lysis, and HRP enzymatic activity was measured in cell lysate. Importantly, we carried out all these experiments in the presence of FBS in order to verify the stability of the anchorage suggested by the previous stability experiment. The number of EVs per cell was variable between experiments depending on the EV/HRP association capacity but remained in the range of 500–2000 EVs/cell to still be in physiologically relevant conditions (Hagey et al. 2023).

First, EV~CLS-PEG₂₀₀₀-HRP were tested for cell internalisation at different HRP amounts (1, 5, 10 and 15 ng per well, Figure 6A). Native EVs without HRP did not trigger any peroxidase activity in PANC-1 cells lysate, validating the absence of detectable endogenous peroxidase activity. We observed a slight increase in HRP activity when cells were treated with free CLS-PEG₂₀₀₀-HRP control with up to 64 ± 9 pg recovered at 15 ng incubated. By contrast, we observed a significant increase in HRP activity when cells were incubated with EV~CLS-PEG₂₀₀₀-HRP: from 18 ± 2 pg at 1 ng/well to 310 ± 50 pg at 15 ng/well incubated (Figure 6A). Incubation time impact on HRP internalisation was assessed after 2, 4 and 24 h for an amount of HRP fixed at 10 ng/well (Figure 6B). Internalised HRP increased over time from 14 ± 2 pg after 2 h to 27 ± 4 pg after 24 h for controls and from 56 ± 5 pg after 2 h to 194 ± 27 pg after 24 h for EV~CLS-PEG₂₀₀₀-HRP. Then, we investigated the mechanism of this internalisation by inhibiting it at 4°C (Figure 6C). Only 4 ± 1 and 15 ± 4 pg of HRP were recovered at 4°C compared to 8 ± 2 and 97 ± 12 pg at 37°C for CLS-PEG₂₀₀₀-HRP control and EV~CLS-PEG₂₀₀₀-HRP, respectively, suggesting a temperature-dependent internalisation process as expected from endocytosis. Finally, we tested two other recipient cell lines in addition to PANC-1 cells, namely A549 and SKBR3 cells (adherent epithelioid carcinoma cell line derived from human lung and breast, respectively), to evaluate the versatility in protein delivery of this approach for other cell lines (Figure 6D). 358 ± 16 , 230 ± 21 and 119 ± 55 pg of HRP were recovered in cells lysates of A549, PANC-1 and SKBR3 cells, respectively, demonstrating the ability of EV~CLS-PEG₂₀₀₀-HRP to be internalised in different human adherent epithelioid carcinoma cell line in vitro. We also validated the versatility of this approach by using the second source of mMSC-EVs (previously described) to associate and deliver HRP to PANC-1 cells (Figure S8).

In order to control and validate the intracellular location of anchored cargo after delivery, A549 cell internalisation was observed using both epifluorescence microscopy (Figure 7A,E) and confocal microscopy (Figure 7F). Fluorescent spots were only visible in EV~CLS-PEG₂₀₀₀-HRP-A488 (Figure 7E) and not in controls. Due to the limited Alexa-Fluor labelling of the HRP (resulting from a limited number of free lysines after LA-conjugation), we analysed 3D-stack images using a random-forest algorithm to determine the number of fluorescent spots inside cells (Figure 7F). We considered this approach to be more objective than “naked-eyes” observation of the stack, especially due to the expected location of this lipidated HRP to the inner cell membrane once delivered. 52 ± 6 spots per 1.10^6 voxels were detected in EV~CLS-PEG₂₀₀₀-HRP-A488 in contrast to controls

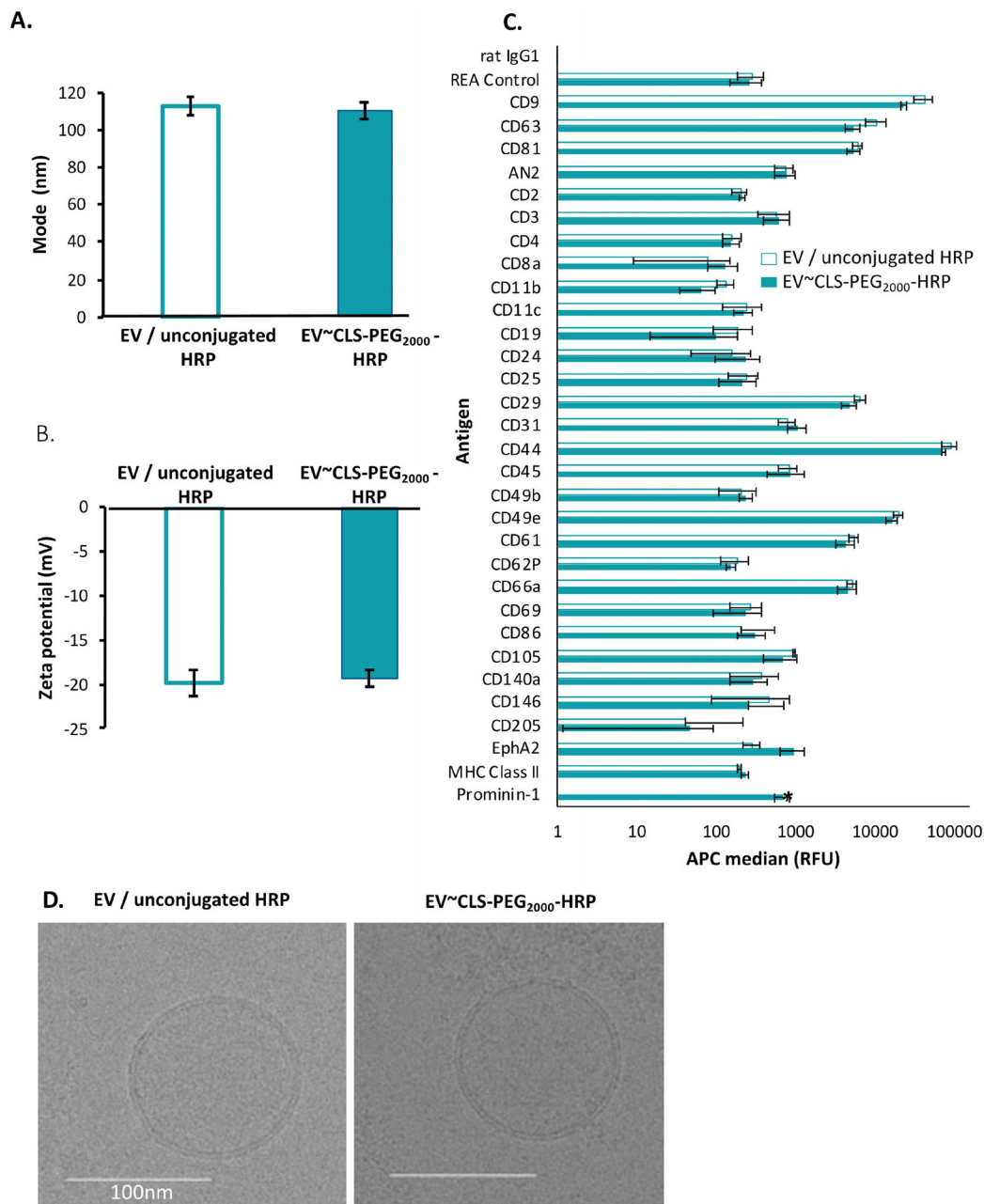


FIGURE 4 | Characteristics of mMSC-EV associated or not with Lipid-PEG-HRP. (A) Size (mode, NTA, NS300, $n = 5$) and (B) Zeta potential (NTA, Zetaview Twin, $n = 3$) after incubation of EVs with unconjugated HRP or CLS-PEG₂₀₀₀-HRP and SEC separation. (C) MACSplex Assay analysis as APC median after subtraction of buffer background signal. p values were calculated using a two-tailed Welch test and only Prominin-1 was significant ($*p < 0.05$). (D) CryoTEM images of EVs after incubation with unconjugated HRP or CLS-PEG₂₀₀₀-HRP. No SEC separation was processed on these samples ($n = 3$).

that presented <1 spot per 1.10^6 voxels. Taken together, these results confirmed quantitative cell lysis experiments.

3.2.2 | Lipid-PEG Linker MW Impacts Cell Internalisation of EVs~Lipid-PEG-HRP by Steric Hindrance of Native EV Surface Ligands

Finally, we evaluated the impacts of lipid and PEG moieties of Lipid-PEG anchors on the cell internalisation (Figure 8A). In contrast to controls, EV~CLS-PEG₂₀₀₀-HRP, EV~DSPE-PEG₁₀₀₀-

HRP, EV~DSPE-PEG₂₀₀₀-HRP and EV~DSPE-PEG₅₀₀₀-HRP were internalised at 124 ± 22 , 140 ± 8 , 87 ± 4 and 12 ± 5 pg respectively. Regarding these results, it seemed that the PEG linker MW strongly influence the cell internalisation efficiency. We hypothesised that this could be due to steric hindrance and that a control of the density of DSPE-PEG₅₀₀₀-HRP per EVs could restore their internalisation properties. To assess this assumption, we investigated the influence of a decreased density of cargos associated per EV on cell internalisation (Figure 8B,D). Interestingly, only EV~DSPE-PEG₅₀₀₀-HRP showed improvements in cell internalisation by 1.4 ± 0.1 and 2.1 ± 0.3 fold-change, thanks to

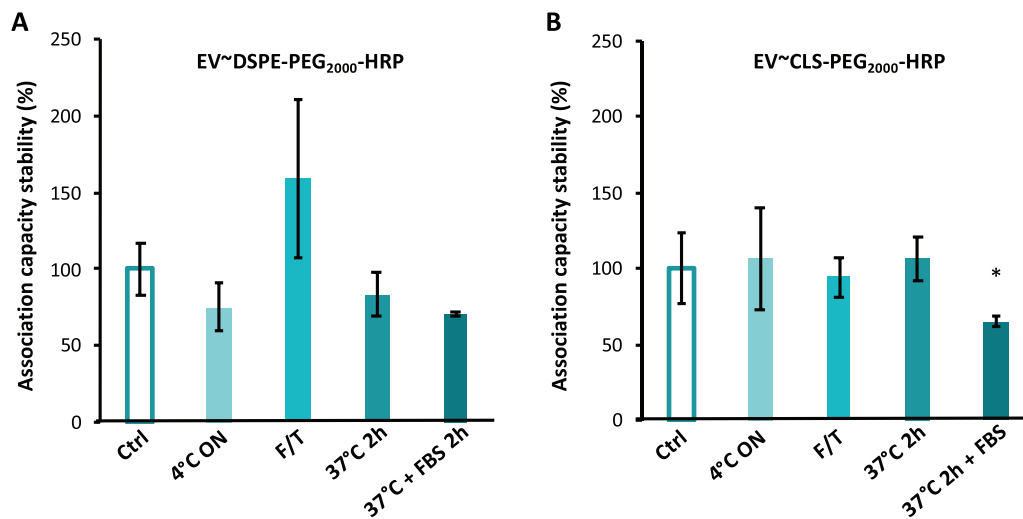


FIGURE 5 | Stability of the EV/Lipid-PEG₂₀₀₀-HRP association. Association capacity of (A) DSPE-PEG₂₀₀₀-HRP and (B) CLS-PEG₂₀₀₀-HRP in percentage of control samples. Samples were diluted at 1:11 in DPBS (or DPBS supplemented with 1:50 FBS if indicated) and incubated before SEC separation. 4°C ON samples were incubated at 4°C ON, F/T were frozen at -80°C before thawing and 37°C 2 h (+FBS) were incubated at 37°C during 2 h either with DPBS or DPBS + 1:50 FBS. $n = 3$ except for CLS-PEG₂₀₀₀-HRP 4°C ON ($n = 2$). p values were calculated using a two-tailed Welch test and samples were compared with Ctrl. $n.s > 0.05$, $* < 0.05$.

a decreased density from 337 to 171 and 84 HRP/EV. The lack of changes in EV~CLS-PEG₂₀₀₀-HRP and EV~DSPE-PEG₁₀₀₀-HRP strongly suggests that steric hindrance of PEG₅₀₀₀ was responsible for this observation, underlining the interest to control these parameters to maintain EV/target cells interactions.

4 | Discussion

4.1 | Lipid-Anchorage Is a Versatile Approach Compatible With Protein Cargos Whilst Maintaining Their Activity

The main aim of this study was to study fundamental parameters controlling EV/LA-protein association to provide a versatile way to use EVs as effective delivery systems for intracellular vectorisation of a functional protein. Indeed, no cellular internalisation of an active protein associated with EVs using an LA-based approach has been reported for now. We chose to use a click-chemistry reaction between maleimide and thiol to associate Lipid-PEG and cargo protein to take advantage of the high versatility of this approach, compatible with virtually all cargo proteins. Although no unconjugated cargo was loaded by simple incubation, up to 898 ± 247 cargos were associated per EV (Figure 2B). Assuming that HRP cargo and EV hydrodynamic radius are equal to 3 nm (Rennke and Venkatachalam 1979) and 50 nm, respectively, one can estimate an EV surface coverage of $\approx 90\%$ (using the *smoothed octagon packing onto a sphere* model [Reinhardt 1934, Fejes Tóth et al. 2023]). This calculation neglects EVs larger than 100 nm and the EV surface fraction already occupied by native membrane proteins but suggests that LA anchorage enables a very high association efficiency. Interestingly, this coverage density did not prevent EV/cell interaction using PEG <5000 g/mol, since decrease in density using medium PEG had no effect (Figure 8B,C).

By optimising the amount of cargo protein during incubation, we reached an association efficiency of 53% (after correction of the measured value which was underestimated due to the loss of EVs during SEC) (Figure 2C), which is among best protein loading efficiencies with EV reported into the literature (Rankin-Turner et al. 2021). We used three different single-particle analysis methods to characterise association heterogeneity, namely fluorescent NTA, nano-flow cytometry and super-resolution microscopy. Overall, up to 70% of particles were cargo+ (Figure 3A,B), among which half were tetraspanins+ (Figure 3B) according to nanoFCM measurements. This supports observations already done by other groups (Tréton et al. 2023, Chen et al. 2021) and highlights the importance of characterising the loading/association efficiency along with other metrics (i.e., association capacity and efficiency). However, it seems important to underline that our labelling strategy was experimentally limited because majority of HRP lysines was already used for the LA-conjugation, which left few free lysines for the labelling with Alexa-Fluor. This has likely resulted in the underestimation of HRP-positive EVs. Moreover, it has been demonstrated that antibody labelling of EVs could have a limited efficiency (Mitrut et al. 2024) and that all EVs do not express the three targeted tetraspanins (CD9, CD63 and CD81) (van Niel et al. 2018, Mizenko et al. 2021).

Versatility of this approach has been controlled with another EV product. These EVs were also produced by mMSC but produced in 2D flask culture and isolated using ultracentrifugation (in contrast to the main commercial source of EVs used in this work, produced under turbulent flow from mMSC seeded onto microcarriers and isolated using SEC and tangential flow filtration). These EVs presented a different membrane composition in terms of surface antigens (Figure S6). We observed differences in association efficiency (Figure S7), capacity and cell delivery (Figure S8A,B, respectively) which could be due to differences in the corona density and/or composition along with the membrane lipid content. Such impact of the EV source of LA-conjugation has been reported by comparing EVs from different cells sources

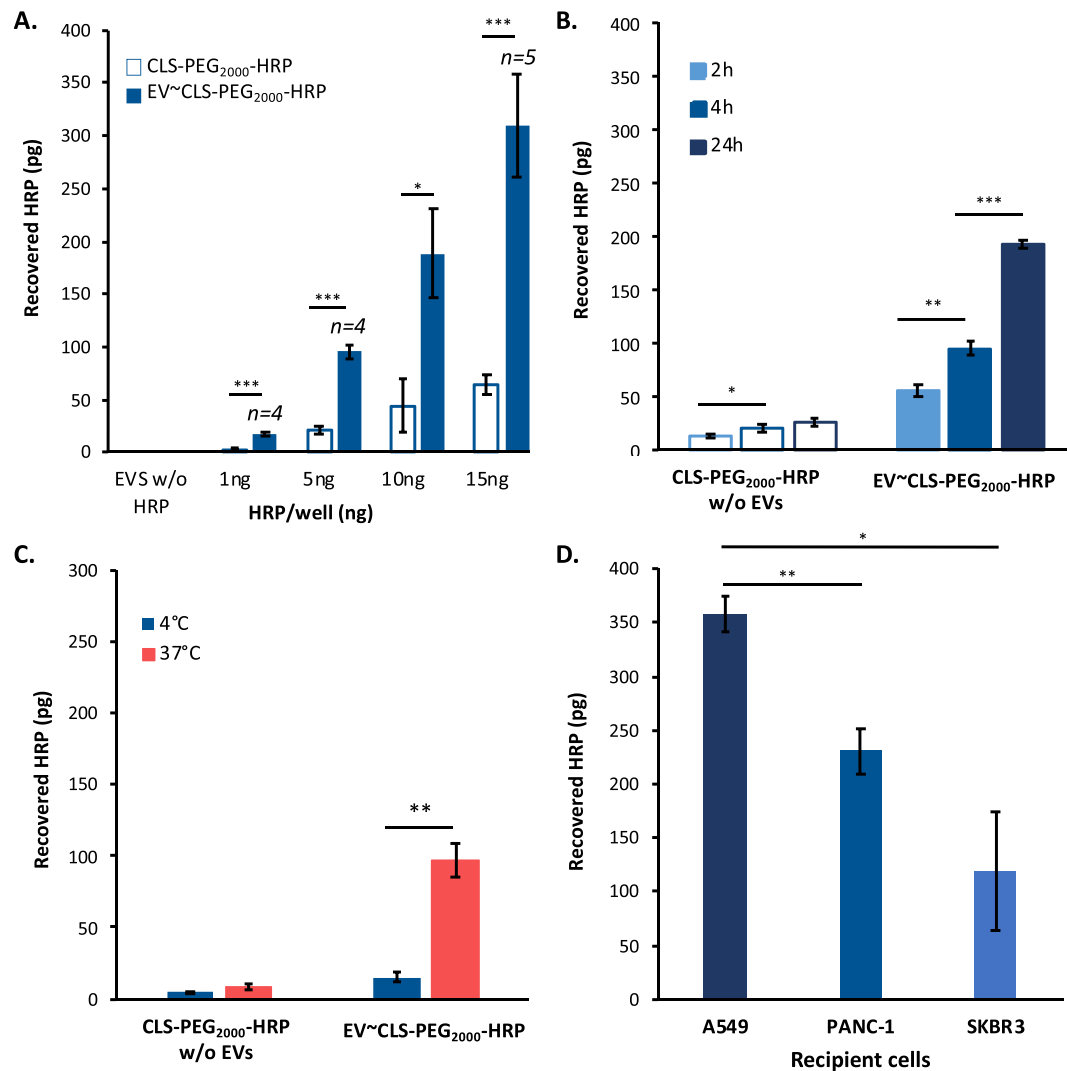


FIGURE 6 | Quantification of the absolute amount of internalised HRP depending on quantity, temperature and incubation time on cell internalisation of EV~CLS-PEG₂₀₀₀-HRP. Recovered CLS-PEG₂₀₀₀-HRP vectorised or not by EVs in PANC-1 lysate after incubation with 1.10^5 PANC-1 following these conditions: (A) 4 h, 37°C, 1, 5, 10, or 15 ng of HRP/well. (B) 2, 4, or 24 h of incubation, 37°C, 5 ng/well of HRP. (C) 2 h, 4°C or 37°C, 10 ng/well of HRP. (D) 4 h, 37°C, 10 ng with 1.10^5 A549, PANC-1, or SKBR3 cells. For each experiment, *p* values were calculated using a two-tailed Welch test. *n.s.* > 0.05, * < 0.05, ** < 0.01, *** < 0.001. *n* = 3 if unspecified.

(Zheng et al. 2023) but the impact of the production and purification process on LA-conjugation was not reported, to the best of our knowledge.

We showed that cholesterol (CLS) was 3.5-fold more efficient for association than DSPE with mMSC-derived EVs (Figure 2B). This greater affinity of CLS could be due to the cholesterol condensing effect (Alwarawrah et al. 2010) combined with its natural presence in high amounts in EV membranes (Skotland et al. 2020). Supporting this assumption, a recent study (Tréton et al. 2023) demonstrated the ability of the EV membrane to insert up to 20% exogenous cholesterol in addition to endogenous cholesterol instead of replacing it. Interestingly, this outperformance by the cholesterol anchor has also been observed for siRNA cargos (Haraszi et al. 2018, Biscans et al. 2018) but not by some other teams working with smaller chemical cargos (Zheng et al. 2023). These results suggest that the pres-

ence of a large protein cargo did not reduce CLS anchoring efficiency.

4.2 | Control of Lipid Anchorage Enables Intracellular Delivery of Cargo Protein and Is Affected by LA Structure

After a successful association study and optimisation, we investigated the intracellular delivery of cargo protein. Regarding these results, CLS-PEG₂₀₀₀ and a ≈ 1000 cargo/EV ratio were chosen as standard conditions. This high number of cargo associated per EVs obtained in this study enabled the use of low amount of EVs to treat cells, with 200–2000 EVs per cell, a dose compatible with cargo-specific cell response as recently established (Hagey et al. 2023). Even in the presence of serum, the association and vectorisation of protein cargo with EVs enabled up to 50-fold

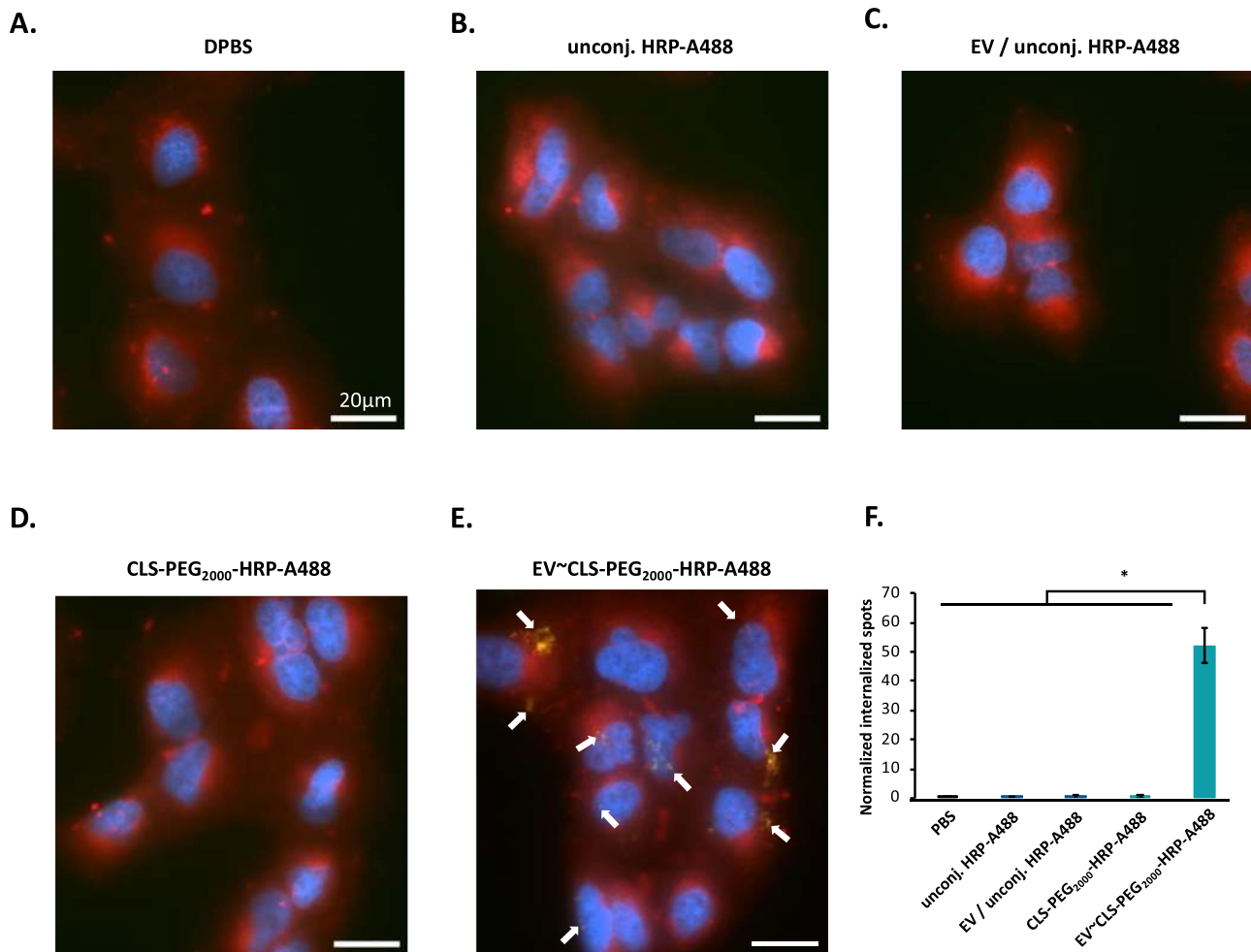


FIGURE 7 | Epifluorescence and confocal microscopy observation and quantification of HRP-A488 cell internalisation. A549 cells were incubated 4 h with HRP-A488, conjugated with CLS-PEG₂₀₀₀ or not and associated with EVs or not. Panel images have been acquired using full-field epifluorescence microscopy. Blue, Red and green fluorescence are DAPI, Cell Mask lipophilic dye and Alexa-fluor 488 from labelled HRP, respectively. (A) A549 cells control incubated with DPBS. (B and C) A549 cells incubated with unconjugated HRP-A488 alone and mixed with EVs, respectively. (D and E) A549 cells incubated with CLS-PEG₂₀₀₀-HRP-A488 alone and associated with EVs, respectively. Identical HRP mass was deposited between EV conditions and their respective controls. Identical number of EVs was deposited between EV conditions. Bar scale is 20 μm. White arrows highlight some HRP-A488 spots. (F) Quantitative analysis on confocal 3D image stacks obtained using spinning-disk confocal microscopy. Number of fluorescent spots in cells was normalised per total number of voxels corresponding to intracellular space. $n = 2$ images except for EV/unconj. HRP-A488 ($n = 3$). For each experiment, p values were calculated using a two-tailed Welch test. $n.s > 0.05$, $* < 0.05$.

more intracellular delivery in three different carcinoma-derived cell lines (PANC-1, A549 and SKBR3 cells). These results suggest that proteins could be more stable cargos for lipid anchorage than small chemicals (Zheng et al. 2023, Tréton et al. 2023). Some differences in protein delivery were observed among cell types, confirming the already described phenomenon that EV internalisation can differ depending on cell type (Ilahibaks et al. 2023). Interestingly, this internalisation was concentration-, time- and temperature-dependent, allowing for a fine-tuning of the amount that can be delivered. Unfortunately, it is difficult to compare these masses of internalised cargo with literature since such absolute measurements are barely ever reported. Finally, we highlighted the impact of three PEG linker MW on cell internalisation. Indeed, when the cargo protein was associated with EVs with a high MW PEG linker (5000 g/mol), the cell internalisation efficiency was significantly decreased. This effect was mitigated by decreasing the number and, therefore density

of DSPE-PEG₅₀₀₀-HRP per EV. Such impact of the LA linker on EV/cargo association and cellular uptake was never reported and should probably be considered in future studies.

5 | Conclusion

Although EVs are promising biomolecule vectors, their membrane is as impermeable to proteins as the cellular one. Thus, reaching acceptable protein loading capacity and intracellular delivery remains an important challenge. Altogether, we demonstrated that the exogenous association of active proteins to EVs using LA is possible and can lead to efficient intracellular delivery systems, even in the presence of serum. By thoroughly characterising this association and numerous governing factors (Lipid nature, PEG linker, temperature, EV origin, etc.), it was possible to control the number of HRP associated with EV surface,

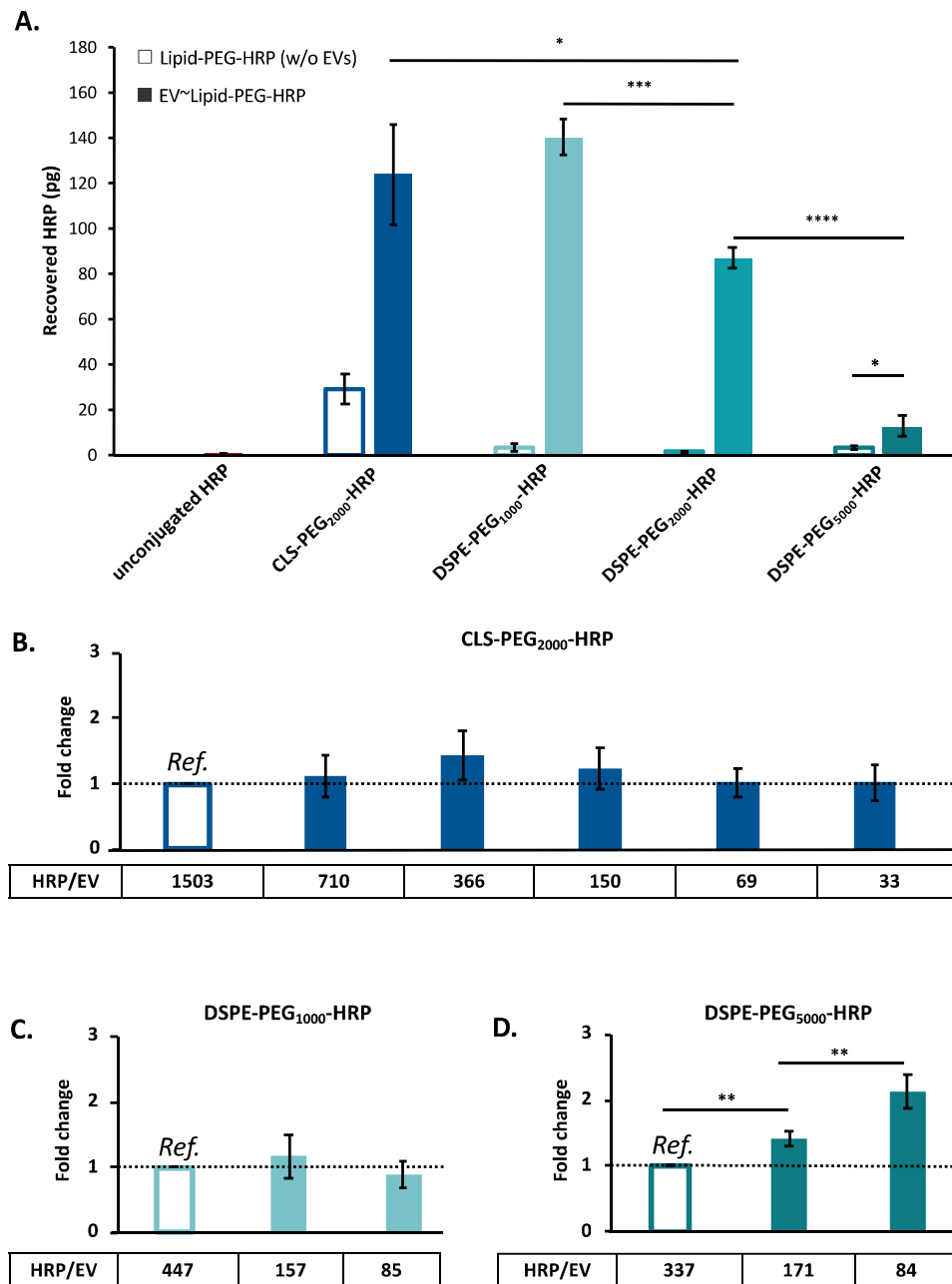


FIGURE 8 | Impact of lipid anchor nature, PEG linker MW and HRP/EV ratio on cell internalisation. (A) Recovered Lipid-PEG-HRP in PANC-1 lysate after incubation of 10 μ g of Lipid-PEG-HRP associated or not to EVs, for 4 h at 37°C with 1.10^5 PANC-1 cells ($n = 4$). (B, C and D) Fold change in recovered Lipid-PEG-HRP in PANC-1 lysate after incubation of 1500 EV~Lipid-PEG-HRP per cell for 4 h at 37°C with 1.10^5 PANC-1 cells. Decreasing HRP/EV ratio have been tested with CLS-PEG₂₀₀₀-HRP, DSPE-PEG₁₀₀₀-HRP and DSPE-PEG₅₀₀₀-HRP for (B), (C) and (D), respectively. The proportion of Lipid-PEG-HRP recovered in lysate was normalised with the highest ratio of the experiment (indicated in the figure as Ref.). Significances are compared to 1 (null hypothesis = no changes whatever the HRP/EV ratio) ($n = 3$). For each experiment, p values were calculated using a two-tailed Welch test. $n.s$ > 0.05, * < 0.05, ** < 0.01, *** < 0.001, **** < 0.0001.

allowing for efficient and tunable intracellular delivery. These results prove the particular interest to thoroughly characterise EV-association processes with the aim to preserve their natural properties for cell interactions. To improve this process, other molecules, such as lipids and polymers, could be investigated. In particular, the effect of the PEG linker has been by far less studied than lipid impact, whatever the type of cargo, despite being crucial, as shown in this study. Thus, this study highlights the value of controlled and rational engineering of EVs to make them

efficient biomolecule delivery systems, opening up promising prospects for various protein replacement or inhibition therapies.

Author Contributions

Antonin Marquant: Conceptualization, data curation, formal analysis, investigation, methodology, validation, visualization, writing—original draft, writing—review and editing. **Jade Berthelot:** conceptualization,

investigation, methodology, validation, visualization. **Claudia Bich:** conceptualization, investigation, methodology, validation, visualization. **Zeineb Ibn Elfekih:** investigation, methodology, validation, visualization. **Laurianne Simon:** conceptualization, investigation. **Baptiste Robin:** conceptualization, investigation. **Joël Chopineau:** conceptualization; resources. **David Tianpei Wang:** data curation, formal analysis, investigation, methodology, validation, visualization. **Samuel Jay Emerson:** data curation, formal analysis, investigation. **Aijun Wang:** conceptualization, resources. **Clément Benedetti:** software, visualization. **Simon Langlois:** formal analysis, software. **Laurence Guglielmi:** conceptualization, methodology, resources. **Pierre Martineau:** conceptualization, methodology, resources. **Anne Aubert-Pouëssel:** conceptualization, supervision, visualization, writing—original draft, writing—review and editing. **Marie Morille:** conceptualization, funding acquisition, project administration, resources, supervision, validation, visualization, writing—original draft, writing—review and editing.

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Conflicts of Interest

The authors declare no conflicts of interest.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Supporting Information

Additional supporting information can be found online in the Supporting Information section.