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Introduction

Regenerative medicine holds significant promise for leveraging the therapeutic potential of stem cells. Amniotic-derived cells, with their distinctive characteristics such as widespread availability, absence of ethical concerns, and immune privileges facilitating their use in allo- and xenotransplantation, exhibit broad clinical potential in both medical and veterinary fields, albeit only partially explored.

Moreover, the inherent stemness of these cells takes advantage of remarkable plasticity, evident in both *in situ* and *ex situ*

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Optimization of a nanoparticle uptake protocol applied to amniotic-derived cells: unlocking the therapeutic potential[†]

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Stem cell-based therapy implementation relies heavily on advancements in cell tracking. The present research has been designed to develop a gold nanorod (AuNR) labeling protocol applied to amniotic epithelial cells (AECs) leveraging the pro-regenerative properties of this placental stem cell source which is widely used for both human and veterinary biomedical regenerative applications, although not yet exploited with tracking technologies. Ovine AECs, in native or induced mesenchymal (mAECs) phenotypes via epithelial-mesenchymal transition (EMT), served as the model. Initially, various uptake methods validated on other sources of mesenchymal stromal cells (MSCs) were assessed on mAECs before optimization for AECs. Furthermore, the protocol was implemented by adopting the biological strategy of MitoCeption to improve endocytosis. The results indicate that the most efficient, affordable, and easy protocol leading to internalization of AuNRs in living mAECs recognized the combination of the one-step uptake condition (cell in suspension), centrifugation-mediated internalization method (G-force) and MitoCeption (mitochondrial isolated from mAECs). This protocol produced labeled vital mAECs within minutes, suitable for preclinical and clinical trials. The optimized protocol has the potential to yield feasible labeled amniotic-derived cells for biomedical purposes: up to 10 million starting from a single amniotic membrane. Similar and even higher efficiency was found when the protocol was applied to ovine and human AECs, thereby demonstrating the transferability of the method to cells of different phenotypes and species-specificity, hence validating its great potential for the development of improved biomedical applications in cell-based therapy and diagnostic imaging.

trans-differentiation processes.^{1,2} Similar to other stem cell sources, they demonstrate diverse paracrine regulatory functions, including immunomodulatory, antimicrobial, and antifibrotic activities.^{3–6} Unlike other stem cell sources, these biological attributes are constitutively expressed, defining unique functions of amniotic-derived cells, such as maintaining an immunotolerance barrier between the fetus and the mother.

Additionally, these abilities can be potentiated in response to the surrounding inputs, such as inflammatory mediators. Nevertheless, it has recently been demonstrated that there is an active conversation between them and neighboring cells which can also occur through mitochondrial transfer pathways.⁷ Based on these premises, amniotic-derived cells have been proposed as candidates for the treatment of severely damaged tissues that cannot be saved even by the most advanced surgical or pharmacological treatments.^{8,9}

Amniotic-derived cells combine the great pro-regenerative actions with the ability of a robust and enduring engraftment

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into damaged tissues.^{10–17} This capability enables their substantial contribution to the long-term healing, replacement, and restoration of tissues and organs.¹⁸ In addition, epithelial amniotic-derived cells (AECs) as mesenchymal stromal ones (AMSCs) exploit the migratory activity as a mechanism to reach damaged tissues.¹⁹ In AECs, the mobility is enhanced because of the activation of the EMT (EMT) process. This activation occurs in response to specific chemoattractant signals released by stromal or resident cells, triggering a recruitment response.^{20,21} This process, known as "homing", becomes crucial for the success of tissue regeneration and healing.

Cell mobility also plays a vital role in determining the appropriate injection strategy for transplanting stem cells.^{22,23} To develop cell therapy systems that can be readily translated into the clinical setting, a precise method for delivering cells into the body still needs to be established. Unfortunately, current protocols suffer from low delivery efficiency, as only a limited number of cells reach the target tissue and remain there after systemic administration.²⁴

Drawing inspiration from stem cell trafficking within the body under physiological and pathological conditions, recent advances have been made in inducing stem cell mobilization and directing patients' stem cells to sites of interest for treating a broad spectrum of diseases.²⁵ This perspective has paved the way for a new paradigm in the field of regenerative medicine, allowing for targeted and personalized management of stem cell-based therapy. Additionally, understanding the mechanisms of cell trafficking represents a challenging biological advancement for standardizing transplant protocols, aiming to maximize the homing abilities of stem cells into injured tissues and establish their long-lasting presence in damaged areas. Achieving these goals would significantly enhance the therapeutic potential of stem cell-based therapies.

A first fundamental prerequisite for achieving these targets is to have a reliable cell tracking system, capable of monitoring the diffusion of stem cells in the host bodies and defining the timing and conditions for the cells to get concentrated in the target zone without affecting their viability and/or their biological functions. In this context, labelling stem cells with nanomaterials working as contrast agents for imaging devices has recently attracted great attention in both the medical and veterinary fields.^{26,27} Different protocols have been developed so far for labelling mesenchymal stem cells (MSCs) to follow their homing and engraftment,²² monitoring cell survival and viability,²⁸ assessing tissue regeneration,²⁹ optimizing transplantation routes and studying biodistribution³⁰ and understanding their mechanisms of action.³¹ Importantly, in the veterinary field, tracking models of allogeneic mesenchymal cells have been developed in canines^{32,33} and equines.^{34,35}

To date, no approaches for tracking amnion-derived cells have been disclosed, despite their significant usage in preclinical and clinical models of stem cell-based treatments.^{1,2,20,36–39}

The current study aimed to optimize a 10×41 nm nanoparticle (NP) uptake protocol applied to ovine mesenchymal and epithelial amniotic-derived cells taking the advantage of the capability to manipulate their phenotype *in vitro*, thereby influencing their immunological and homing characteristics.

Experimental

Ethics statement

No ethical statement is necessary for the use of the ovinederived amniotic cells since they are obtained from waste tissue obtained from animals slaughtered for feed purposes.

Human amniotic epithelial cells derived from discarded human placentae from uncomplicated cesarean resections at 37–42 weeks of gestational age were obtained from Karolinska University Hospital (Stockholm, Sweden) with institutional review board approval and signed informed consent.

Cell isolation and culture

AECs were isolated from the AM of female sheep in midpregnancy (3 AM totally) according to Canciello *et al.*⁴⁰ AECs were maintained in MEM Eagle Alpha (ECM0850L, Euroclone) supplemented with 10% fetal bovine serum (FBS; #10270-106, Gibco), 100 IU mL⁻¹ penicillin–streptomycin (DE 17-602E, Lonza), 100 IU mL⁻¹ amphotericin B (ECM0009D, Euroclone) and 100 IU mL⁻¹ L-glutamine (ECB300D, Euroclone). AECs and mAECs were obtained, respectively, by culturing AECs for three passages in the presence and absence of 25 mM P₄ (4 pregnene-3,20-dione, P₄; P8783, Sigma-Aldrich) at 38.5 °C and 5% CO₂ in the incubator.

Human AECs were isolated and collected as reported by Gramignoli et al.41 Briefly, the amnion membrane was mechanically separated from the chorion layer and washed with Ringer's solution (Baxter, Sweden) to remove blood cells. Subsequently, the amnion was immersed in TrypLE solution (Life Tech, Paisley, UK) and incubated for 30 min at 37 °C. The released cells were sedimented by centrifugation at 500 \times g for 5 minutes and re-suspended in a cryogenic preservation solution composed by University of Wisconsin solution (Lisieux, France) supplemented with 10% dimethylsulfoxide (DMSO; Sigma-Aldrich, MO, USA). Cells were transferred in controlled freezing containers and stored in the vapor phase in a liquid nitrogen storage tank until uptake experiments. Human AECs were maintained in DMEM Low Glucose (ECM0060, Euroclone) supplemented with 10% fetal bovine serum (FBS; #10270-106, Gibco), 100 IU mL^{-1} penicillin-streptomycin (DE 17-602E, Lonza), and 100 IU mL⁻¹ amphotericin B (ECM0009D, Euroclone). Human AECs were not passaged before uptake experiements. Cells were kept at 37 °C and 5% CO_2 in the incubator for all the experimental procedures.

AuNRs

Rhodamine (535/575 nm) labelled 10 × 41 nm gold nanorods (AuNRs; CF12-10-808-RHO-FB-DIH-50-1-CS-EP) with a surface plasmon resonance (SPR) 808 nm were obtained from NanopartzTM. AuNRs were managed according to the manifacturer's instructions.

Chemicals

The stock solutions of gold (Au) at 100 mg L^{-1} in 10% (v/v) HCl and rhenium (Re) at 1000 mg L^{-1} in 2% (v/v) HNO₃ were purchased, respectively, from CPAchem Ltd (Bogomilovo, Bulgaria) and Merck KGaA, (Darmstadt, Germany). The nitric and hydrochloric acid used for all dilutions were grade ultrapure, $\rm HNO_3 \geq 60\%$ and $\rm HCl \geq 30\%$ both supplied by Supelco (Merck KGaA, Darmstadt, Germany). All working solutions, standard, blanks and samples were prepared with type I (resistivity 18.2 M Ω cm) water, obtained from a ELGA LabWater PURELAB Option-Q water purification system (High Wycombe, United Kingdom). Certified reference material NPs SRM 8011 gold nanoparticles (nominal 10 nm diameter) was bought from NIST (Gaithersburg, USA). Argon and helium gases of 99.9995% purity were supplied by Sapio (Monza, Italy).

Immunofluorescence

Cells were grown on glass coverslips, treated as indicated in the cell culture section and then fixed with 4% paraformaldehyde (#11699404, VWR) for 10 min and permeabilized with 0.1% (v/v) Triton X-100 (T8787, Sigma-Aldrich) in $1 \times$ phosphate buffered saline (PBS; P3813, Sigma-Aldrich) for 10 min. Glass coverslips were then blocked in % (w/v) bovine serum albumin (BSA; A3059, Sigma-Aldrich) in $1 \times PBS$ for 1 h at room temperature (RT) and stained overnight at 4 °C with the following primary antibodies: anti-cytokeratin-8 (1:200) (ab2530, Abcam) and anti-vimentin (1:200) (M0725, Agilent DAKO). Cy3 (AP124C, Millipore) or Alexa Fluor 488 (ab150113, Abcam) conjugated anti-mouse secondary antibodies were diluted 1:200 in 1% (w/v) BSA/PBS and incubated on glass coverslips for 40 min at RT. Nuclear counterstaining was obtained with 4',6-diamidino-2-phenylindole (DAPI; D9542 Sigma-Aldrich) diluted 1:2000 in 1× PBS. Coverslips were mounted with fluoromount (F4680, Sigma-Aldrich), and cell samples were analyzed using an Axioskop 2 plus incident light fluorescence microscope (Zeiss) equipped with a CCD camera (Axiovision Cam 208color, Zeiss), with a resolution of 1300×1030 pixels, configured for fluorescence microscopy, and interfaced to a computer workstation, provided with an interactive and automatic image analyser (Zen 3.4 Blue edition Software, Zeiss). Digital images were acquired using standard filters setup for Cy3, Alexa Fluor 488 or DAPI.

Cytotoxicity assay

The CellTiter 96 aqueous one solution cell proliferation assay (MTS; #G3582; Promega) was used to determine the viability of mAECs cells treated with different concentrations of AuNRs. In detail, cells were seeded in 96-well plates at a density of 2 \times 10⁴ cells per cm² and were allowed to attach overnight in a CO2 incubator. Cells were then treated with 0, 1.1, 4.4, and 18 μ g mL⁻¹ of AuNRs for 12, 24 and 48 hours. The cells were then washed three times with PBS (P3813, Sigma-Aldrich) to remove the unbound AuNRs. The MTT reagent was added 2 h before the termination of the experiment, and the optical density was acquired at 490 nm. The NP concentration and timeframe were determined based on cell viability below 70%. For long term experiments ovine or human AECs labelled by following the optimized condition for uptake (24 hours onestep with 4.4 μ g mL⁻¹ AuNRs + G-force + MitoCeption) were checked for their viability over time (24, 48, 72 and 96 hours) by MTS following the manufacturer's instructions.

Internalization of AuNRs

1. One-step and two-step cultural strategies. Two distinctive cultural approaches were tested.

In the two-step method, mAECs adhering as a cell monolayer to the bottom of the support dish were exposed to AuNRs. Specifically, mAECs (2 \times 10⁴ cells per cm²) were plated, and after 24 hours, the growth medium was removed, and the cells were cultured with AuNRs dispersed in fresh medium.

The one-step method involved the addition of AuNRs to mAECs as a cell suspension, followed by cell seeding into a low-adherent culture dish (2×10^4 cells per cm²) and subsequent incubation using a cell shaker. A wide concentration range of AuNRs spanning from 0 µg mL⁻¹ to 18 µg mL⁻¹ and at different time points (12, 24 and 48 hours) were tested for evaluating AuNRs uptake performances gathered, respectively, with the one-step and the two-step methods.

2. Passive incubation with AuNRs. The passive incubation strategy was assessed for both one-step and two-step protocols. In the two-step, mAECs $(2 \times 10^4 \text{ cells per cm}^2)$ were plated, and after 24 hours, the growth medium was removed. Subsequently, cells were exposed to AuNRs dispersed in fresh medium at a concentration of 1.1, 4.4 and 18 µg mL⁻¹ for 12, 24 and 48 hours within the incubator. A control cell plate was prepared similarly, without the addition of AuNRs (0 µg mL⁻¹ AuNRs).

For the one-step protocol, AuNRs at a concentration of 1.1, 4.4 μ g mL⁻¹ were added to a cell suspension (ovine mesenchymal and epithelial AECs and human epithelial AECs). Cells were then seeded into a low-adherent culture dish (2 × 10⁴ cells per cm²) and incubated using a cell shaker in the incubator for 12, 24 and 48 hours. A control cell plate was also prepared in a similar manner but without the addition of AuNRs (0 μ g mL⁻¹ AuNRs).

3. G-force-mediated internalization of AuNRs. The G-forcemediated internalization technique was implemented in the one-step cultural strategy, and its efficacy was evaluated using different AEC cell types (ovine mesenchymal and epithelial AECs and human epithelial AECs).

The protocol used refers to that described by O'Campo and colleagues⁴² with slight modifications. Briefly, AuNRs, at a concentration of 4.4 µg mL⁻¹, were dispersed in either 6 mL or 1 mL of fresh complete medium and applied to the cell pellet containing 2×10^5 cells for cytofluorimetric studies or 1.2×10^6 cells for ICP-MS analysis.

The samples were then subjected to centrifugal forces $(200 \times g)$ for 5 minutes and subsequently seeded in low-adherent culture dishes, achieving a cell density of 2×10^4 cells per cm². Following a 24 hour incubation at 38.5 °C or 37 °C, the samples were processed for gold quantification using ICP-MS and for the examination of cells that had internalized AuNRs through flow cytometry. A control cell plate was prepared in a similar manner, omitting the addition of AuNRs.

4. Mithocondria supplementation (MitoCeption). Freshly isolated mitochondria were obtained using the Qproteome Mitochondria Isolation Kit (#37612, Qiagen) following the manufacturer's instructions. Mitochondria to be used for Mito-Ception were obtained from a quantity of donor cells (mAECs)

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equal to that of the recipient cell population. Isolated mitochondria were suspended in fresh complete medium, combined with AuNRs at the tested dose and placed in contact with the recipient cells (ovine mesenchymal and epithelial AECs, and human epithelial AECs) prior to the application of G-force. Subsequently, the incubation by the one-step protocol was performed for 24 h.

30 minutes before the end of the AuNRs incubation protocol, mitochondria staining was carried out by adding MitoTracker[™] Green FM (M7514, Invitrogen) to the cells at a concentration of 20 nM. Subsequently, the cells were incubated at 38.5 °C or 37 °C for 30 minutes following manufacturer's instructions. The cells were then collected by centrifugation at 300 × g and extensively washed (3 times) with DPBS (#D8662; Sigma-Aldrich) before being used for further analyses.

Gold analytical determination with ICP-MS

Once uptake was completed, the cells were collected at the bottom of a tube by centrifugation at $300 \times g$. Three thorough washes with DPBS (#D8662; Sigma-Aldrich) were performed prior ICP-MS analysis to remove any remaining AuNRs possibly adhered to the cell surface.

At the beginning of the analyses, all equipment and glassware were flushed with 1–2% (v/v) HNO₃ and then rinsed with type I water. After vigorous mixing, a sample volume (50–100 μ l) was solubilized in an adequate quantity of a solution containing 20% (v/v) HNO₃ and 2% HCl (v/v) to obtain a sample dilution of 1:100 in HNO₃ 2% (v/v) and HCl 0.2% (v/v). The sample solutions obtained were analyzed using an inductively coupled plasma mass spectrometer ICP-MS Icap RQ (Thermo Fisher Scientific, Waltham, MA, USA) in KED (kinetic energy discrimination) mode. Instrumental parameters are shown in Table 1.

The isotopic masses acquired were ¹⁹⁷Au, ¹⁸⁷Re. Quantitative gold determination was performed using a calibration curve built with increasing addition standard concentrations (0.2, 0.4, 1, 2, 4, 10, 20, 50, 70, 100, 120 μ g L⁻¹) in 2% (v/v) HNO₃ and 0.2% (v/v) HCl. In all injected solutions (blank, standard and samples), the same amount of Re was added as an internal standard to compensate for the signal fluctuations typical of the ICPMS technique. In each analytical session, the absence of interferents and analytical accuracy were monitored. Therefore, a blank reagent was processed, and the gold concentration was lower than the LOQ value (equal to 0.10 mg L⁻¹) and the recovery of the certified reference material NPs SRM 8011 gold nanoparticles was acceptable in the range (98–103)%.

Table 1 ICP-MS instrumental parameters

Spray chamber Nebulizer Sample/skimmer cone Cool gas (L min⁻¹) Auxiliary gas (L min⁻¹) Plasma power (W) Nebulizer gas flow (L min⁻¹) KED mode: collision gas flow (helium) (L min⁻¹)

Quantification of rhodamine positive cells and MitoCeption by flow cytometry

For flow cytometry measurements, following exposure to AuNRs, cells underwent thorough washing with DPBS (#D8662; Sigma-Aldrich) and were subsequently suspended in a fresh complete medium. The flow cytometry analysis was conducted using a CytoFLEX SRT system (Beckman Coulter), where a total of 10 000 events were analyzed. AuNRs-rhodamine b was excited by a 561 nm yellow laser (Y585 channel), while mitotracker green dye was excited by a 488 nm blue laser (B525 channel). Fluorescence intensity was represented on a standard logarithmic scale. Data analysis and visualization were performed using CytExpert SRT software (version 1.2.10004, Beckman Coulter).

Representative flow cytometer plots were shown for each experimental condition. More in detail, the percentage of cells enclosing AuNRs or mitochondria were measured, respectively, by PE intensity (*x*-axis) or FITC (*x*-axis) and side scattering (SSC; *y*-axis) flow cytometry dot plot. First, the percentage of cells containing AuNRs or mitochondria was assigned by cells of higher PE or mitochondria intensity than the threshold intensity (black vertical line). The threshold value was determined by adding the robust SD to the median intensity of the untreated (0 µg mL⁻¹) cells.

Imaging AuNRs cell internalization with confocal laser scanner analysis

At the end of the incubation period, cells were gently washed three times with DPBS (#D8662; Sigma-Aldrich). Subsequently, the cells were seeded in confocal Petri dishes at a density of 2×10^4 cells per cm² and allowed the support to be adhered for approximately 3 hours before proceeding with confocal laser scanning analysis.

The cells were fixed with 4% paraformaldehyde (#11699404, VWR) for 10 minutes and permeabilized with 0.1% (v/v) Triton X-100 (T8787, Sigma-Aldrich) in 1× phosphate buffered saline (PBS; P3813, Sigma-Aldrich) for 10 minutes. Following this, cell nuclei were stained with DAPI (D9542 Sigma-Aldrich) diluted 1:2000 in 1× PBS for 15 minutes.

In case of phalloidin staining, Alexa Fluor[™] 488 phalloidin (green; #A12379) was added at a concentration of 165 nM, incubated for 1 h at RT and washed extensively before confocal laser scanning microscopy (CLSM; Nikon A1R) analysis. CLSM was interfaced to a computer workstation, provided with an interactive and automatic image analyser (NIS-Element AR Analysis v4.40.00 64-bit). Digital images were acquired using a

> Cyclonic (at 2.7 °C) Micromist at 0.4 mL min⁻¹ Platinum 14 0.8 1550 (0.95-1.15) optimized daily (4.5-5.5) optimized daily

standard laser setup for Cy3 (for AuNRs), FITC 488 nm (for phalloidin) or DAPI (for nuclei).

Isolation of PBMCs

Following the collection of 16 milliliters of fresh peripheral blood at the slaughterhouse, ovine PBMCs were extracted using a density gradient centrifugation method. As directed by the manufacturer, a gradient was constructed using 12 mL of Ficoll-Paque PLUS (#GE17-1440-02, Cytiva). The cells were preserved using the above-discussed methods and kept in liquid nitrogen until they were required for immunological testing.

CM production from AECs

CM production was started once the AECs (either ovine or human) reached 24 hours of 4.4 µg mL⁻¹ AuNRs uptake with one-step + G-force + MitoCeption. The cells were cultured 4 hours in serum-free media (ovine AEC with MEM Eagle Alpha, human AECs with DMEM low glucose, both supplemented with 1% penicillin/streptomycin, 1% amphotericin B and 1% L-glutamine) before exposing them to the inflammatory lipopolysaccharide (LPS) stimulus (1 mg ml⁻¹, #L2637; Sigma-Aldrich) for 1 hour.⁴³⁻⁴⁵ The cells were then washed twice and cultured in serum-free medium for 24 hours before collecting the conditioned medium (CM). The CM was centrifuged at 300 × *g* for 10 minutes to remove any remaining cells or debris, and the supernatants were used for the PBMC proliferation assay.

PBMCs proliferation assay

The immunomodulatory effect of ovine and human AECderived CM was verified by the proliferation rate of PBMCs by using MTS. In detail, PBMCs were seeded in 96-well plates at a concentration of 2×105 cells, activated with 10 µg mL⁻¹ of phytohemagglutinin (PHA-L; #00-4977-93; invitrogen) and cultured for 48 h with or without CM. Then, PBMC proliferation was assessed by using the CellTiter 96 aqueous one solution cell proliferation assay (MTS; #G3582; Promega) following the manufacturer's instruction protocol. Data were normalized on PHAactivated PBMCs.

qPCR

The total RNA was extracted using a total RNA purification kit (# 17250 Norgen Biotek Corp.) following the manufacturer's instructions. 1 μ g of total RNA was retrotranscribed using oligodT primers (#BIO-38029, Bioline) and Tetro reverse transcriptase (#BIO-65050, Bioline), following the manufacturer's instructions. The qPCRs were carried out in triplicate using the SensiFAST SYBR Lo-ROX kit (#BIO-94050, Bioline) on a 7500 fast real-time PCR system (Life Technologies), according to the

manufacturer's instructions. The following PCR conditions were used for all the experiments: 95 °C for 10 min, followed by 40 cycles at 95 °C for 10 s and 60 °C for 30 s. Relative quantification was performed by using the $\Delta\Delta$ Ct method. GAPDH (glyceraldehyde 3-phosphate dehydrogenase) and YWHAZ (tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein zeta) were selected amongst the housekeeping genes for gene quantification. Gene expression data were the mean \pm SD, of the Δ Ct values related to the selected housekeeping. Specifically, one of the two housekeeping genes was reported. The expression profiles were similar with both reference genes. The primer sequences are reported in Table 2.

Statistical analysis

Data were generated from three independent experiments. Data were plotted and statistically analyzed using the software package GraphPad Prism version 5.0 for Windows. All samples were compared using a one-way ANOVA and Bonferroni *post hoc* test. Only significant differences (*p*-value < 0.05) among the samples are indicated in the charts.

Results and discussion

NP uptake optimization according to amniotic cell phenotypes

The major objective of the present study is to define whether the currently available protocols validated on mesenchymal stromal cells (MSCs) are effective in amniotic-derived cells. As a further novelty, a biological strategy (mitochondrial transfer, also known as MitoCeption) was tempted to enhance the uptake mechanisms.⁴²

Using this setting, the validation of the NP uptake protocol was first performed on ovine mesenchymal amniotic-derived cells (mAECs) resulting from the *in vitro* transition of epithelial ones (AECs).⁴⁴ Once the NP uptake protocol was verified and implemented on mAECs, the approach was then extended to the native pool of AECs of both ovine and human origin.

Amniotic-derived cells offer, indeed, the possibility to start from an epithelial phenotype and move towards the mesenchymal one exploiting the physiological pathway that controls epithelial-mesenchymal transition (EMT) in this cell type.

Indeed, amniotic cells with epithelial or mesenchymal phenotype coexist *in vivo* in the amniotic membrane (AM) where the EMT path is switched on in epithelial labor.⁴⁶ *In vitro*, AECs undergo spontaneous EMT after isolation from AM, acquiring a mesenchymal phenotype because of expansion in conventional media.^{47,48} This process can be prevented by progesterone (P_4) supplementation by mimicking in culture the hormonal asset of pregnancy as demonstrated recently by our group.⁴⁴

Table 2	Sequences of primers and conditions used in real-time qPCR
	· · ·

Gene	Forward sequence	Reverse sequence	Annealing temperature (°C)
GAPDH	5'-TCGGAGTGAACGGATTTGGC-3'	5'-CCGTTCTCTGCCTTGACTGT-3'	64.4
YWHAZ	5'-AGACGGAAGGTGCTGAGAAA-3'	5'-CGTTGGGGATCAAGAACTTT-3'	61.3
Vimentin	5'-GACCAGCTCACCAACGACA-3'	5'-CTCCTCCTGCAACTTCTCCC-3'	61.2
Cytokeratin-8	5'-CTCAAAGGCCAGGGCTTC-3'	5'-CTTGGCCTGGCATCCTTGA-3'	61.3

Crucially, our group proved, in addition, that the amniotic cell phenotype strongly affected the stem cell properties: in particular, to enhance stemness, *in vitro*⁴⁴ and *in vivo* plasticity^{16,49} and migratory properties.⁵⁰

Exploiting the effect of P_4 , for the current research, two different cell typologies (epithelial *vs.* mesenchymal) have been generated starting from AECs freshly isolated from 3 different fetuses. The phenotype characterization of both the epithelial and mesenchymal cell subpopulations was evaluated after three passages of expansion using specific protein and gene markers. Vimentin was employed as a late mesenchymal marker, while cytokeratin-8 was utilized for epithelial markers (Fig. S1, ESI†).

The interest of the present research towards the AECs, the most studied and used amniotic-derived cells to date,⁵¹ is related to their distinctive therapeutic properties and to the research expertise of our research team that widely characterized *in vitro* and *in vivo* them over the past years.^{1,10,16,17,49,52} Furthermore, our interest extends to mesenchymal amnioticderived cells, benefiting from the collective expertise of various stem cell research groups that occasionally utilize this placental source to isolate fetal mesenchymal stromal cells.

Effect of AuNRs on viability of mAECs

For the present research, gold nanorods (AuNRs) have been selected. Recently, there has been growing interest in a new approach that utilizes gold NPs as a contrast agent for computed tomography (CT) with potential applications in medicine protocols in both medicine²⁶ and veterinary.⁵³ This methodology is gaining attention due to its clinical applicability and low cost-effectiveness. Indeed, research indicates that gold NPs can effectively label various cell types, including stem cells and immune cells, without compromising their therapeutic effectiveness.⁵⁴ Nevertheless, *in vivo* experiments have successfully demonstrated noninvasive, quantitative, and longitudinal cell tracking with high sensitivity using this approach using animal models.^{55–60} ICP-MS approach with flow cytometry and confocal laser scanning analysis.

The initial experiments involved the preparation of AuNRs on mAECs to assess their impact on cell viability, employing a validated two-step protocol.²⁶ To employ AuNRs as cell-labeling agents, their cellular absorption and potential toxicity were assessed.

First, a dose-response curve with a range of Au-NRs (from 1.1 to 18 μ g mL⁻¹) encompassing the working dose indicated by the manufacturer (4.4 μ g mL⁻¹) was screened to assess the influence of AuNRs exposure on mAECs using the MTS assay. Three time points for AuNRs exposure were considered (12, 24, 48 h). As shown in Fig. 1, no cytotoxicity was seen after 12 h, 24 h, or 48 h of AuNRs incubation using the concentration range from 1.1 to 4.4 μ g mL⁻¹. In contrast, supplementation of 18 μ g mL⁻¹ dramatically reduced mAECs viability roughly 65% regardless of the incubation period. Based on these preliminary data, the experimental set up was then performed by incubating mAECs at both 1.1 and 4.4 μ g mL⁻¹.



Fig. 1 Viability of mAECs exposed to AuNRs using a two-step validated protocol. AuNRs were tested for cytotoxicity in mAECs at doses ranging from 0 to 18 µg mL⁻¹ and at various time points (12, 24, and 48 hours) with MTS. The significance vs. 0 µg ml⁻¹ was indicated with **** for p < 0.0001; vs. 1.1 µg mL⁻¹ was indicated with ### or ## for p < 0.001 and p < 0.01 respectively; vs. 4.4 µg mL⁻¹ was indicated with ^{\$\$} or ^{\$} for p < 0.01 and p < 0.01 and p < 0.05 respectively.

Impact of cellular adhesion on AuNRs uptake

To establish the link between AuNRs uptake protocol and in vitro cell conditions, a comparative analysis was conducted between one-step and two-step approaches. The two-step replicated a method extensively validated in the literature for NP uptake²⁶ where AuNRs are exposed to a cell monolayer where cells adhere to the bottom of the cultural support. On the other hand, the one-step method involves the addition of AuNRs to cells maintained in suspension, using low-adherent culture dish, and shaking. The one-step approach was regarded as promising due to its biological potential and practical advantages. The biological rationale behind employing this protocol is grounded by the evidence demonstrating as NP penetrability through the lipid bilayer is positively influenced by the extension of surface contact area.⁶¹ Nevertheless, it may provide the practical advantage of the immediate use of labeled cells without the need for further processing of the sample. The mAECs were exposed to different doses of AuNRs (1.1 and 4.4 μ g mL⁻¹) at three time points (12, 24 and 48 hours), utilizing both onestep and two-step approaches. Following each incubation with AuNRs, extensive washing was performed to ensure the removal of any AuNRs potentially attached to the cell surface. Fig. S2 (ESI⁺) illustrates a representative confocal laser scanner image of mAECs that, following the washing step, show that cells are free of AuNRs on the membrane surface while AuNRs are displayed internally. Subsequently, mAECs were lysed and analyzed for gold content using ICP-MS, a reference analytical technique, consolidated in the last decade for reliable NP-metal quantification.⁶² As depicted in Fig. 2, both systems reach their maximum uptake at 24 hours. Specifically, the one-step approach performs better, incorporating significantly higher quantities of AuNRs at all evaluated time points. When exposed to 1.1 μ g mL⁻¹ AuNRs, an increase of 11, 14, and 12-fold was observed at 12, 24 and 48 hours, respectively, compared to the two-step method. Similarly, exposure to 4.4 μ g mL⁻¹ AuNRs resulted in a 12, 19, and 19-fold increase at 12, 24 and 48 hours, respectively, compared to the twostep approach. Additionally, the highest incorporation is observed



Fig. 2 Cellular uptake dependency on the AuNR concetration, AuNR incubation time and cultural strategy. Quantification of AuNR uptake in 1×10^{6} mAECs cells by inducively coupled plasma mass spectrometry (ICP-MS) at different time points 12, 24 and 48 hours. One-step (blu line and symbol) and two-step (dark line and symbol) approaches were tested as cultural strategies for AuNR incubation. Two dose points of AuNRs were selected for testing dose dependency of uptake: 4.4 μ g mL⁻¹ (straight line) and 1.1 μ g mL⁻¹ (dashed line). The significance of data was reported as follows. One-step 4.4 μ g mL⁻¹ vs. two-step 4.4 μ g mL⁻¹ was indicated with **** for p < 0.0001; one-step 1.1 µg mL⁻¹ vs. two-step 1.1 µg mL⁻¹ was indicated with [#] or ^{####} for p < 0.05 and p < 0.0001; one-step 4.4 $\mu g~mL^{-1}$ vs. one-step 1.1 $\mu g~mL^{-1}$ was indicated with $^{\$}$ or $^{\$\$\$\$}$ for p~<0.05 and p < 0.0001. Two-step 4.4 µg mL⁻¹ vs. two-step 1.1 µg mL⁻¹ was indicated with § for p < 0.05. Statistically significant differences vs. time point 0 h were indicated with ^{aaaa} or ^{aa}, respectively, for p < 0.0001 and p < 0.01, vs. 12 h with ^{bbbb} (p < 0.0001) and vs. 24 h with ^{cccc} (p < 0.0001).

when cells are exposed to the highest dose of AuNRs ($4.4 \,\mu g \, mL^{-1}$), showing a peak uptake at 24 h followed by reaching a plateau at longer uptake times (Fig. 2). Consistently, literature evidence demonstrates a similar trend in nanorod uptake rates.^{63–66} The two-step culture strategy incorporates lower quantities of AuNRs, showing an increase at 24 h only when cells are exposed to a dose of AuNRs equal to $4.4 \,\mu g \, mL^{-1}$. Furthermore, extending the uptake times does not reach a plateau; instead, a decrease is observed (Fig. 2).

Another critical aspect requiring consideration is the determination of the percentage of cells containing AuNRs acquired at the end of the cell labeling protocol. Evaluating this factor is essential for confirming the efficacy of the proposed cell tracking technique. A rapid and effective method validated in literature for determining the percentage of cells in a specific subset is flow cytometry.²⁶ To date, this analytical technique has been extensively utilized to quantify NP uptake in mammalian cells since the introduction of NP to cells significantly increases side scatter, as documented in several studies.^{67–72} Importantly, its application for determining the percentage of NP-containing cells has gained significant interest in recent years⁷³ even if the sensitivity in measurements is considerably lower compared to more complex techniques such as ICP-MS-based technologies. 74

Of note, the applicative advantage of flow cytometry lies in its non-invasiveness. If required, the examined cells can be retrieved using a cell sorter and promptly employed for in vivo cell tracking purposes. A flow cytometric investigation was conducted to assess the percentage of mAECs containing AuNRs using both one and two-step. Notably, this analytical phase focused on the time window between 12 h and 24 h, guided by the results obtained from ICP-MS, which clearly indicated a plateau in AuNRs incorporation at 48 h. As shown in ESI[†] Fig. S3, a significant increase in the number of labelled cells was detected using the one-step protocol with 4.4 μ g mL⁻¹ AuNRs after 24 h of incubation. In more detail, mAECs cultured using the one-step protocol and exposed to 1.1 μ g mL⁻¹ and 4.4 μ g mL⁻¹ AuNRs for 24 h exhibited a respective 2.2-fold (2.1 \pm 0.7% vs. 0.95 \pm 0.2%) and 3-fold (5.9 \pm 0.4% vs. 2.1 \pm 0.3%) increase in positive cells compared to the percentage observed in the two-step approach. Consistent with the results obtained from ICP-MS concerning the internalized AuNRs, flow cytometric analysis exhibited the best performance in terms of cells labeled with AuNRs using the one-step approach at the optimized dose of 4.4 μ g mL⁻¹ after 24 hours of exposure. Remarkably, the analysis demonstrated a positive correlation between AuNRs data recorded with ICP-MS (amount of Au internalization) and flow cytometry (rhodamine positive cells) at 4.4 μ g mL⁻¹ after 24 h. This latter analysis definitively confirmed the one-step approach as the most efficient protocol for Au uptake and allowed the set-up of the cultural strategy before NP exposure (Fig. 3).

Even though the dynamics of uptake are closely dependent on a multitude of factors related to either NP features and cell type, biological context, and the function of the cells themselves,^{72,75} It is plausible to hypothesize that the one-step approach exhibits superior uptake capability as first it does not require preliminary steps, such as trypsinization, to prepare the batch of cells for NP labeling. These preliminary steps can be a source of stress and may negatively influence endocytosis dynamics.⁷⁶

By avoiding potentially harmful procedures such as trypsinization, the one-step approach ensures a more straightforward and less invasive process for NP labeling, leading to more reliable and accurate results. This methodological improvement is particularly advantageous for maintaining the integrity of the cells and the fidelity of the uptake process, thus making the one-step approach more favorable under specific experimental conditions here detailed. Experimentally, suspension cells can also be easier to work with for endocytosis studies because they do not need to be detached from a surface for analysis and/or further applications. This can reduce experimental variability and simplify the protocol potentially offering the practical advantage of having a labelled cell population ready-to-be used for tracking procedures. This avoids additional manipulation, which might potentially cause the loss of the same nanorods from the cells or/and cell viability.

The adoption of combined analytical approaches has yielded robust and highly confident data, providing a precise measure

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Fig. 3 Correlation between the average quantity of AuNRs internalized and the percentage of AuNRs positive cells after 24 hours of incubation with 4.4 μ g mL⁻¹ AuNRs using a one- or two-step cultural method. The average number of AuNRs was quantified by ICP-MS (1 \times 10⁶ mAECs), whereas the % of positive AuNRs was determined by flow cytometry. Statistic related to μ g of AuNRs internalized by cells was indicated with *** for p < 0.001. Statistic related to AuNRs positive cells percentage was indicated with ^{###} for p < 0.0001.

of uptake. Notably, at present, there is no primary method for measuring NP uptake, necessitating the use of orthogonal methods to achieve reliable results⁷⁷ for⁷⁸ silica⁷⁹ NPs. Furthermore, Böhme and colleagues utilized ICP-MS and flow cytometry, as in the present research, to accurately quantify the uptake of Al₂O₃ NP.⁸⁰

Cell uptake improvement by combining G-force with mitochondria supplementation

Overall, based on the previous findings, the one-step culture approach using 4.4 μ g mL⁻¹ AuNRs after 24 hours of incubation was chosen for further experiments aimed at identifying strategies to enhance cell uptake.

First, the addition of AuNRs to mAECs was performed under the influence of G-force, based on literature evidence attesting a positive influence of the centrifugal force (hereinafter referred as G-force) on the internalization efficiency of NP.42 This effect had been attributed to the forced NP deposition on cell plasma membrane. Subsequently, mAECs were incubated with 4.4 μ g mL⁻¹ AuNRs for 24 h using the one-step method. Data were then analyzed by comparing the AuNRs uptake performances using ICP-MS and flow cytometry (AuNRs internalized amount and % of AuNRs positive cells, respectively) with or without (referred as passive incubation) G-force. As depicted in Fig. 4, G-force induces a significant increase in both AuNRs amount and cell positivity to AuNRs. More in detail, the AU internalization in mAECs increased 25% (0.75 \pm 0.04 vs. 0.6 \pm 0.05 μ g/10⁶ pellet; Fig. 4A and C) whereas the percentage of positive cells passed from 5.9 to 7.73% (p < 0.05; Fig. 4B and C).

Then, an additional optimization step was tempted. To improve AuNRs uptake, the mechanism of endocytosis was modulated by introducing isolated mitochondria by exploiting a recognized mechanisms of cell-to-cell dialogue (*e.g.* passage/ exchange of organelles).

Importantly, recent evidence showed that internalization of exogenous mitochondria causes the activation of the actindependent endocytic pathway, demonstrated even by inhibiting endocytosis function.^{81–83} Starting from this premise and considering the recent successful attempts demonstrating a novel method to guide the transfer of mitochondria from cell donor to cell recipient, known as MitoCeption,⁸⁴ freshly isolated mitochondria from mAECs (donor cells) were incorporated into the one-step protocol for AuNRs uptake, utilizing G-force. Data reported in Fig. 4 showed that mitochondria supplementation determines a huge and significant increase in Au amount internalized in mAECs ($1.1 \pm 0.15 \ \mu$ g, about 2-fold; Fig. 4A and C) and, at the same time, it significantly increases the percentage of positive-rhodamine cells ($18.6 \pm 0.83\%$; more than 2.5-fold; Fig. 4B and C).

To demonstrate that mitochondrial transfer facilitated the enhanced internalization of AuNRs through MitoCeption, the organelles were pre-labelled with the fluorophore Mitotracker green. The MitoCeption was then quantified using flow cytometry.

As reported in Fig. 4B, mitochondria transfer occurred by involving $45.63 \pm 0.6\%$ of mAECs (Ctrl: 0 µg mL⁻¹ AuNRs) and $45.74 \pm 1\%$ of cells exposed to 4.4 µg mL⁻¹. Significantly, MitoCeption synergizes with the mechanism of NP micropinocytosis. In fact, nearly all cells tested positive for AuNRs (83.10 \pm 1%) exhibited incorporated mitochondria (Fig. S4, ESI†).

Confocal imaging was finally employed to further validate the enhancement in cell uptake achieved by combining G-force with MitoCeption. As depicted in Fig. 4D confocal laser scanning microscopy provided clear information consistent with the improved uptake data discussed earlier.

AuNRs uptake standardization applied to AECs

Finally, the protocol optimized for mAECs was extended to the native source of ovine AECs (see Fig. S1, ESI[†]) and then translated to human AECs. Human AECs have been extensively researched and utilized.²⁰ In contrast to ovine AECs, isolated from animals at mid-gestation, human AECs are term cells obtained from the amniotic membrane. Biologically, they have a distinct background; the ovine AECs used in this study are isolated at a stage where they actively support pregnancy, whereas the human one adopted for the present research are cells that have fulfilled their biological role. Nonetheless, they remain a significant source of fetal stem cells due to their broad pro-regenerative properties.

To this purpose, ovine AECs and human AECs were exposed to AuNRs uptake using the optimized protocol. More in detail, the previous source of isolated mitochondria collected from ovine-derived mAECs was supplemented to one-step cultured cells exposed to G force treatment.

ICP-MS analysis confirmed greater internalization in AECs compared to mAECs, regardless of species origin. More in detail, ovine AECs and human AECs internalized 1.5 (1.7 \pm 0.1 vs.

1.1 \pm 0.15 µg/10⁶ pellet) and 3.6 (4.0 \pm 0.4 vs. 1.1 \pm 0.15 µg/10⁶ pellet) times, respectively, more that ovine mAECs (Fig. 5A).

The result of improved uptake ability of the cells with the native epithelial phenotype was also confirmed by flow cytometry (Fig. 5B) showing a 1.3 ($25.0 \pm 0.5 vs. 18.6 \pm 0.83\%$) and 2.6 ($48.1 \pm 3.8 vs. 18.6 \pm 0.83\%$) fold increase in the percentage of rhodamine-positive ovine and human AECs, respectively. Confocal analysis corroborated these data (Fig. 5C). Notably, the human derived AECs showed a very high uptake efficiency that could be dependent of their species specificity (lipid





Fig. 4 Uptake effectiveness following passive incubation with AuNRs, Gforce mediated internalization of AuNRs and mitochondrial driven AuNRs uptake. (A) Panel A shows the quantification of AuNRs uptake in 1×10^6 AECs cells by ICP-MS following 24 h of incubation using a one-step approach. Average values of the AuNRs μ g/10⁶ pellet were reported. Statistics is reported below the table and summarized in panel C. (B) Panel B shown representative flow cytometer plots for each experimental condition. The table reports the average % values \pm SD related to the AuNRs positive mAECs detected by flow cytometry. Statistics is reported below the table and summarized in panel C. (C) Panel C shows the correlation between the average quantity of AuNRs internalized and the percentage of AuNRs positive cells after the centrifugation-mediated internalization method (G-force) and mitochondria supplementation. After 24 h of incubation, the average number of AuNRs was quantified by ICP-MS, whereas the % of positive AuNRs was determined by flow cytometry. Significance was reported as follows. Statistic related to µg of AuNRs internalized by cells was indicated with ** for p < 0.01. Statistic related to AuNRs positive cell percentage was indicated with # or ###, respectively, for p < 0.05 and p < 0.0001. (D) Panel D shows representative CLSM images at $40 \times$ magnification of mAECs before and after the addition of AuNRs at the selected concentration of 4.4 μ g mL⁻¹ with one-step cultural strategy for 24 h. Positive cells are shown in red due to the functionalization of the AuNRs with rhodamine dye, nuclei were stained with DAPI. Uptake improvement strategies (G-force and mitochondria supplementation) were shown together with the results of the passive incubation. Scale bars: 50 µm.

composition of plasma membrane, endocytosis/pynocitosis function, *etc.*) or from the different lifetime of the two amniotic stem cell sources (late human *vs.* middle stage ovine).

Considering that the objective of labeling these cells is their use in regenerative medicine protocols, it is essential to evaluate the long-term effects of nanoparticle incorporation, ensuring the preservation of cellular and biological functionalities.

To this end, the long-term effect of NP incorporation on cellular functionality was monitored by MTS assay in both ovine and human AECs labeled with the optimized labeling condition, with the observation endpoint set at 96 hours. AuNRs labeling remained stable over time, with an unchanged percentage of AuNR-positive cells detected by flow cytometry compared to the 24-hour time point (Fig. S5, ESI†). Notably, cell viability data indicate a perfect preservation of cellular functionality over time (Fig. 6A).

Also, biological properties of AECs bearing AuNRs were evaluated. Accordingly, our recent studies described how AECs exhibit significant immunomodulatory properties. These properties are expressed through the paracrine release of specific immunomodulatory molecules that regulate the inflammatory regeneration.50,85 microenvironment, promoting tissue Specifically, it has been observed that conditioned media (CM) from this cell source, whether from ovine or human AECs, play a crucial role in modulating immune cell activities by suppressing the proliferation of different immune cells, including peripheral blood mononuclear cells (PBMCs).^{50,85-87} This response was significantly heightened when the cells were exposed to inflammatory-like stimuli.85,86

The immunomodulatory influence of CM derived from LPS-stimulated ovine and human AECs was tested using the PBMCs activation test. As shown in Fig. 6B, CM inhibited the PHA-induced proliferation of PBMCs by approximately 70% (p < 0.0001 vs. PHA) and ~60% (p < 0.0001 vs. PHA) for ovine

and human AECs, respectively. These data confirm that the native attitudes of these cells are preserved upon AuNRs incorporation.

Overall, the present results proved the efficiency of the combo protocol involving physical (G-force) and biological (MitoCeption) strategies in improving AuNRs incorporation.



Fig. 5 Standardization of AuNR uptake in AECs. (A) Average number of AuNRs internalized into 1×10^6 AECs of ovine and human origin quantified by ICP-MS. Statistic related to μ g of AuNRs internalized by cells was indicated with * for p < 0.05 and with **** for p < 0.0001. (B) Left panel: Representative flow cytometer plots showing the % \pm SD of positive AuNRs AECs of ovine and human origin. Right panel: The graph reports the quantification of the average % values related to the AuNRs positive ovine and human AECs detected by flow cytometry. Statistic related to AuNRs positive cells percentage was indicated with * for p < 0.05 and with **** for p < 0.0001. (D) Panel D shows representative CLSM images at 40× magnification of ovine and human AECs before and after the addition of AuNRs at the selected concentration of 4.4 μ g mL⁻¹ with one-step cultural strategy + G-force + mitochondria for 24 h. Positive cells are shown in red due to the functionalization of the AuNRs with rhodamine dye, nuclei (blu) were stained with DAPI. Scale bars: 50 μ m.



24

48

500 72

D 96

Fig. 6 Long-term effect of AuNR incorporation using the optimized uptake conditions. (A) Cell viability of both ovine and human AECs was checked with MTS over time (24, 48, 72 and 96 hours) upon incorporation of 4.4 µg mL⁻¹ AuNRs by using the optimized condition (one-step + G-force + mitochondria). Statistical analysis indicated no significant differences between groups, denoted as ns in the graph. (B) Ovine and human AECs were labelled with the optimized condition (24 hours one-step + G-force + mitochondria). CM derived from labelled ovine od human AECs stimulated with LPS were collected and used for 48 hours treatment of PBMCs. Statistic between groups was indicated with ** for p < 0.0001.

Furthermore, the optimized uptake protocol can be effectively utilized with amniotic-derived cells, enabling the generation of a batch of labeled cells ranging from 2×10^5 to 5×10^5 , depending on the AECs phenotype and the species of origin used for AnNRs incorporation, starting from an initial concentration of 1×10^6 .

The availability of this quantity of cells does not impose a restriction on AECs, as each amniotic membrane can yield 100 to 500 million viable cells in the ovine placenta,⁴⁰ and 80 to 300 million viable cells in a full-term human placenta.⁴¹

Our research group has optimized the recovery protocol of ovine AECs from the AM at middle gestational stage (fetus of 25-30 mm in size), allowing over 20 millions of cells to be to obtained from 10 cm^2 of AM.⁴⁰ This practical aspect is quite relevant in order to document the feasibility of the proposed protocol in generating a realistic amount of autologous immune-privileged labelled cells ready to be used for

Paper biomedical applications in medicine and veterinary. The protocol formulation seems to exceed the intracellular AuNR concentration necessary to be visualized with CT as reported

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concentration necessary to be visualized with CT as reported as demonstrated by Kim and colleagues, who achieved CT detection with at least 2×10^4 cells⁶⁰ labeled with Au using a passive incubation protocol.

Of relevance, the present results were obtained by requiring the use of a small initial AuNR concentrations (4.4 μ g mL⁻¹) if compared with other tracking protocols that started from initial Au NP doses ranging from μ g mL⁻¹ up to mg mL⁻¹.^{56,60,64,65,88,89}

Finally, the optimized protocol proposed in this research appears to significantly enhance AU internalization efficiency, even at the single-cell level. Premising that our group did not undertake single-cell quantification, as previously documented by Wang and colleagues,⁶³ our results indirectly affirmed an internalized Au amount of 4.0 µg in a population of 0.48×10^6 labeled cells (equivalent to 48.1% out of 1×10^6 cells in human AECs). This yielded a single-cell incorporation of approximately 8.3 pg per cell, originating from a 4.4 µg mL⁻¹ of AuNRs solution. The amount of incorporated gold exceeds the yield levels reported in the literature concerning AuNRs, particularly considering that the nanoparticle exposure concentration in this study was maintained at lower levels than those documented in previous studies.^{64,65}

This high yield data coupled with a low exposure concentration is highly relevant from a biological standpoint, as it is preferable for avoiding changes in cellular functionality and cytotoxicity.⁹⁰ This becomes particularly critical, as in our case, for regenerative medicine applications, necessitating the development of systems that not only track cells but also require them to function at the site of repair.

Conclusions

In this study, an optimized protocol for the uptake of gold NPs (AuNRs) by amniotic-derived cells, leveraging cell suspension (onestep), G-force and MitoCeption, was developed. This new uptake method significantly improved AuNRs incorporation in ovine amniotic-derived cells independently of cell phenotype (epithelial and mesenchymal) and in human epithelial cells, enhancing their application for regenerative medicine and biomedical imaging.

The ability to greatly internalize NPs without compromising cell vitality or therapeutic properties opens new avenues in both human and veterinary medicine, particularly in non-invasive tracking of cell therapies and enhancing diagnostic precision with AuNRs as CT contrast agents. Future research will focus on standardizing this labeling protocol for clinical use, by assessing long-term effects in preclinical models to ensure safety and stem cell retention of their native regenerative potential. By establishing a new cell labeling protocol for cellular tracking, this study also unveils key insights into the mechanisms of cellular communication setting the stage for advanced studies to unravel the mechanisms behind NPs uptake and mitochondrial transfer, potentially revolutionizing personalized stem cell-based strategies of repair.

Author contributions

Alessia Peserico: validation, formal analysis, investigation, data curation, writing – original draft. Angelo Canciello: investigation, resources, visualization. Giuseppe Prencipe: investigation, resources, visualization. Roberto Gramignoli: resources. Valeria Melai: investigation. Giampiero Scortichini: resources. Mirella Bellocci: investigation. Giulia Capacchietti: investigation. Maura Turriani: resources. Chiara Di Pancrazio: resources. Paolo Beradinelli: supervision. Valentina Russo: supervision. Mauro Mattioli: conceptualization, methodology, supervision. Barbara Barboni: conceptualization, methodology, supervision, writing – review & editing, project administration, funding acquisition.

Data availability

Any additional information is available from the corresponding author upon request.

Conflicts of interest

There are no conflicts to declare.

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