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## Introduction

Cell therapy is becoming a potent treatment approach for multiple human diseases. To support the development and accelerate the clinical translation of such 'living' systems,

# Pivotal role of the protein corona in the cell uptake of fluorinated nanoparticles with increased sensitivity for <sup>19</sup>F-MR imaging<sup>+</sup>

*In vivo* cell tracking by non-invasive imaging technologies is needed to accelerate the clinical translation of innovative cell-based therapies. In this regard, <sup>19</sup>F-MRI has recently gained increased attention for unbiased localization of labeled cells over time. To push forward the use of <sup>19</sup>F-MRI for cell tracking, the development of highly performant <sup>19</sup>F-probes is required. PLGA-based NPs containing PERFECTA, a multibranched superfluorinated molecule with an optimal MRI profile thanks to its 36 magnetically equivalent fluorine atoms, are promising <sup>19</sup>F-MRI probes. In this work we demonstrate the importance of the surface functionalization of these NPs in relation to their interaction with the biological environment, stressing the pivotal role of the formation of the protein corona (PC) in their cellular labelling efficacy. In particular, our studies showed that the formation of PC NPs strongly promotes the cellular internalization of these NPs in microglia cells. We advocate that the formation of PC NPs in the culture medium can be a key element to be used for the optimization of cell labelling with a considerable increase of the detection sensitivity by <sup>19</sup>F-MRI.

longitudinal cell imaging tools would be needed and they allow the monitoring of cell biodistribution, therapy outcomes and prevent off-target toxicity. Accordingly, the Food and Drug Administration (FDA) recommended the development and, subsequently, the use of non-invasive imaging techniques to track cells. Magnetic resonance imaging (MRI), when compared to other approaches, has the advantage of being safe, exploitable in longitudinal studies, and able to provide whole-body data also useful for clinical use.

Nanoparticles (NPs) containing conventional MRI contrast agents (*i.e.*, gadolinium or iron based paramagnetic agents) can be used to label cells and render them detectable by MRI. A large number of preclinical studies with superparamagnetic iron oxide (SPIO) NPs have shown promising results.<sup>1</sup> However, a rather poor signal specificity, a low cellular uptake and the fact that SPIO NP concentration is not linearly proportional to MRI signal are considered limiting factors, preventing a quantitative and accurate analysis.<sup>2,3</sup> The detection of SPIO NPs is unspecific since it is determined as signal loss in the tissue and could lead to misinterpretation and to rise of artefacts because these NPs strongly interfere with the proton signal (<sup>1</sup>H-MRI). Beyond the incompatibility of SPIO NPs detection with standard MRI analysis, these contrast agents could be confused with vessels, haemorrhage, necrosis or endogenous iron traces normally present in healthy or diseased tissues.

A number of studies recently highlighted the potential of fluorinated NPs as labelling agents for cell tracking by <sup>19</sup>F-

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MRI.<sup>3-8</sup> Indeed, as endogenous fluorine in tissues is not detected by MRI, the signal originating from fluorinated NP-loaded cells specifically refers to the infused product and is directly quantifiable by <sup>19</sup>F-MRI.<sup>3-7,9</sup> Of note, the <sup>19</sup>F-MRI signal is directly proportional to the amount of <sup>19</sup>F atoms, it is stable and does not interfere with the standard signal from <sup>1</sup>H. Indeed, <sup>19</sup>F-MRI can be combined with any <sup>1</sup>H-MRI protocols on the same scanner. Therefore, <sup>19</sup>F-MRI is a suitable strategy for unbiased localization and tracking of labelled cells over time. The low sensitivity of <sup>19</sup>F-MRI currently limits its potential applicability and translation to the clinic. While the use of high field MRI scanner and cryogenic radiofrequency probes could improve the sensitivity,<sup>10,11</sup> the customized design of better performing fluorinated tracers bears the potential to maximize imaging performance. In this respect, we previously reported that a superfluorinated molecule, named PERFECTA, showed ideal chemical and structural properties for <sup>19</sup>F-MRI for improving imaging sensitivity, thanks to its large number of <sup>19</sup>F atoms per molecule.<sup>12-15</sup> Importantly, the 36 <sup>19</sup>F atoms are symmetrically distributed in this molecule resulting in intense MRI signals at a specific and unique chemical shift.12 In addition to a high 19F payload, the physicochemical properties of the formulation, necessary to disperse these fluorinated molecules in physiological solutions as <sup>19</sup>F-NPs, could be tuned to obtain optimal internalization by cells without affecting their functionality.

Considering that NP internalization mainly occurs through endocytosis mechanisms,16-18 their uptake is strongly dependent on NP size, shape, charge and composition.<sup>19-22</sup> Moreover, proteins can be adsorbed onto the NP surface, and form a layer called protein corona (PC) impacting on the surface and colloidal properties of NPs with consequences on the cellular uptake mechanism. This biomolecular layer is characterized by two components: a "hard" (HC) and a "soft" corona, formed by layers more strongly and more loosely bound to the NP layers, respectively. Since the inner HC layer persists on the NP surface for times long-enough to ensure cellular interaction, it is considered more biologically relevant.23 Thus, the PC adsorbed on NPs creates a new bio-nano interface with direct consequences on cellular uptake mechanisms.24-26 For example, adsorption of opsonin on the NP surface can preferentially activate the phagocytic pathway.27

In particular, we have developed polylactic-co-glycolic acid (PLGA) NPs loaded with PERFECTA at high yields with different surface functionalizations. Specifically, PERFECTA@PLGA NPs (<sup>19</sup>F-PLGA NPs) were stabilized by three different biocompatible coating agents, namely poly-vinyl alcohol (PVA), sodium cholate (NaC) and chitosan (CS), forming NPs with a neutral, negative and slightly positive surface electrokinetic potential, respectively. As expected, these NPs were characterized by a specific bio-nano interface in the biological environment<sup>23-28</sup> with the formation, in some cases, of PC NPs with larger size and different surface properties (composition and electrokinetic potential) compared to the original NPs.28,29 The PC formation can be essential to enhance cell labelling performance as it was demonstrated with ferumoxytol NPs for stem cell labelling.30 In other cases, it has been shown that PC can strongly affect the relaxation times of different SPIO NPs depending on their

surface properties and PC composition.31,32 We tested these 19F-PLGA NPs as labelling agents on microglial cells (BV2), which are characterized by a high phagocytic capacity<sup>33</sup> similar to the resident immune cells of the central nervous system that play a fundamental role in immune defence.34 The in vivo tracking of myeloid cells with a natural capacity to phagocyte NPs such as macrophages, microglia, and dendritic cells (DCs) by <sup>19</sup>F-MRI helped monitor the progression of different inflammations,35,36 including the dynamics of cancer-associated immune response.37 In a first clinical trial it was used to trace immunotherapeutic DCs in colorectal adenocarcinoma patients.3 More recent studies exploited human peripheral blood mononuclear cells to validate clinical scanners and evaluate safety compliance.<sup>38</sup> In spite of technology improvements, sensitivity remains an issue when tracking small number of cells. Various NPs have been formulated to maximize the <sup>19</sup>F loading, to increase the interaction with cells and to facilitate their internalization to overcome the sensitivity limits.38,39 In the present work, we performed an optimization of superfluorinated PLGAbased nanoformulations modulating their surface properties to further improve cell labelling efficiency and imaging sensitivity. We compared the cell labelling efficiency of <sup>19</sup>F-PLGA NPs in relation to their surface features by a systematic <sup>19</sup>F-MRI study. The NPs with a higher tendency to form a PC in the culture medium resulted to be the most efficient labelling agents enabling the best <sup>19</sup>F-MRI response (Fig. 1). Moreover, studies on NP cellular uptake mechanisms, using pharmacological inhibitors for specific endocytosis pathways, showed that the highest NP uptake was associated with an enhancement of the phagocytic pathway indicating a significant role of the PC formation in this process.

### Results and discussion

#### Physico-chemical characterization of <sup>19</sup>F-PLGA NPs

<sup>19</sup>F–PLGA NPs were stabilized by sodium cholate (NaC), polyvinyl alcohol (PVA) or chitosan (CS), to obtain NPs with different surface properties.<sup>40</sup> While <sup>19</sup>F–PLGA–NaC and <sup>19</sup>F–PLGA–PVA NPs were formulated following an established procedure,<sup>15</sup> an initial study was dedicated to optimize the preparation protocol of positively charged NPs, which should exhibit a higher electrostatic interaction with the negatively charged cell membrane.<sup>20,41,42</sup> The optimization was carried out to obtain small and positively charged <sup>19</sup>F–PLGA–PVA–CS NPs with a homogeneous size distribution, a fine colloidal stability and high PERFECTA loading (ESI, Section 1.2 and Fig. S1†). Overall, using a CS/PLGA (w/w) ratio of 0.8 and PVA content of 2% (w/v) we obtained NPs with a positive electrokinetic potential (19 ± 8 mV) and the best colloidal properties among all the tested conditions to produce <sup>19</sup>F–PLGA–PVA–CS NPs (Fig. S1E and F†).

On the other hand, <sup>19</sup>F–PLGA NPs coated with either NaC or PVA were characterized as expected by negative and neutral electrokinetic potentials, respectively (Table 1). Dynamic Light Scattering (DLS) characterization in aqueous solution showed stable monodisperse NP size distributions for all the NPs ranging from 100 to 200 nm depending on the stabilizer (Fig. S2A–C†). Of note, <sup>19</sup>F–PLGA NPs stabilized with NaC

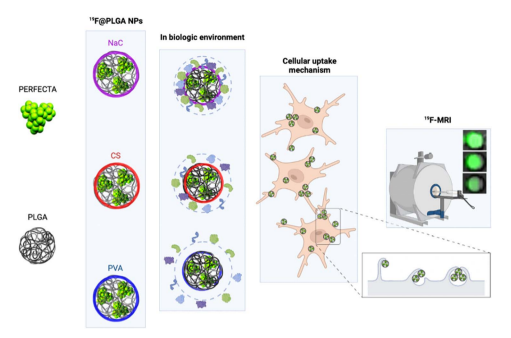


Fig. 1 Schematic representation of the components used to produce <sup>19</sup>F–PLGA NPs containing the superfluorinated probe (PERFECTA) and coated by different stabilizers. The obtained <sup>19</sup>F–PLGA NPs show a different behaviour in the biological environment with a diverse tendency to form the protein corona (PC). The PC modulates cell labelling efficiency through different internalization pathways and has a significant impact on <sup>19</sup>F-MRI sensitivity.

showed a particularly smaller hydrodynamic size, around 100 nm compared to the others. Interestingly, comparable results have been reported on similar NaC coated PLGA NPs loaded with a hydrophilic drug.<sup>43,44</sup> A different level of fluorine encapsulation was found through <sup>19</sup>F-NMR experiments with an external standard for the NPs, ranging from 2 to  $4 \times 10^{20}$  atoms per ml (Table 1).

Considering the concentrations of NPs measured by NTA (Table 1), the average amount of fluorine (PERFECTA) per single NP can be assumed to be considerably higher in PVA-CS and NaC-<sup>19</sup>F-PLGA NPs (almost  $27 \times 10^7$  atoms per NP) rather than in PVA formulation (around  $5.8 \times 10^7$  atoms per NP).

Further analysis by HAADF-STEM confirmed the presence of spherical and colloidally stable NPs (Fig. S3A–C and G–I<sup>†</sup>) while semi-quantitative PERFECTA loading was verified by EDS chemical analysis (Fig. S3D–F<sup>†</sup>). Furthermore, the relaxation

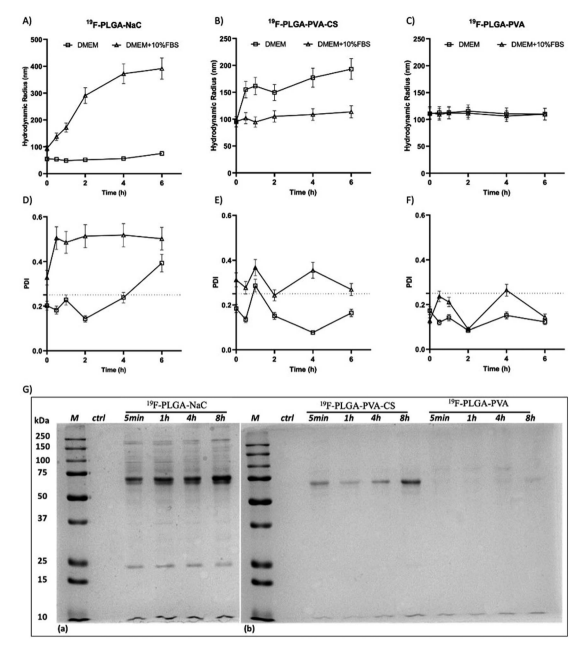
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times of the three <sup>19</sup>F–PLGA NPs were in the same range and suitable for <sup>19</sup>F-MRI, with relatively short  $T_1$  (260–350 ms, Table S1†) and long  $T_2$  (198–220 ms, Table S1†).

The colloidal stability of the NPs was also investigated under physiological conditions and in the presence of proteins. Specifically, size variations of the <sup>19</sup>F–PLGA NPs dispersed in cell culture medium (DMEM), with and without FBS, were evaluated by DLS over time up to 48 h (Fig. 2 and S2†). <sup>19</sup>F– PLGA–NaC NPs, when incubated with FBS, immediately showed after 30 min of incubation (Fig. 2A) an increase in size, while NP size remained stable in pure DMEM (Fig. 2A and D). This behavior was likely due to the NaC coating, which provides a good electrostatic stabilization in buffer solution, but it is known to promote strong interactions with serum proteins leading to the formation of aggregates.<sup>45</sup> Importantly, <sup>19</sup>F– PLGA–NaC NPs did not show sedimentation or precipitation

Surfactants	NP size <sup>b</sup>		Particle charge <sup>c</sup>	Fluorine content <sup>d</sup>		
	$D_{\rm H}$ (nm)	PDI	Zeta potential (mV)	Encapsulation (%)	Atoms per ml $(\times 10^{20})$	NPs per ml <sup><math>e</math></sup> (×10 <sup>11</sup> )
NaC PVA–CS PVA	$egin{array}{c} 104 \pm 10 \ 215 \pm 51 \ 206 \pm 9 \end{array}$	$\begin{array}{c} 0.20 \pm 0.03 \\ 0.16 \pm 0.06 \\ 0.12 \pm 0.03 \end{array}$	$-49 \pm 5 \\ +19 \pm 8 \\ -13 \pm 1$	$48 \pm 10 \\ 47 \pm 9 \\ 61 \pm 12$	$\begin{array}{c} 2.1 \pm 0.4 \\ 3.0 \pm 0.6 \\ 4.0 \pm 0.8 \end{array}$	$8.1 \pm 1.2 \ 11.2 \pm 6.7 \ 69.0 \pm 27.0$

<sup>*a*</sup> All reported values are the mean of six independent measurements performed on different NP batches. <sup>*b*</sup> Hydrodynamic diameter ( $D_{H}$ ) and polydispersity index (PDI) obtained by cumulant fitting of DLS measurements. <sup>*c*</sup> Particle charge determined by  $\zeta$ -potential measurement. <sup>*d*</sup> Fluorine content determined by <sup>19</sup>F-NMR. <sup>*e*</sup> NP concentration determined by Nanoparticle Tracking Analysis (NTA).



**Fig. 2** Hydrodynamic radius and PDI of all the NPs incubated in DMEM (square) or DMEM supplemented with 10% FBS (triangle) at 37 °C for time points from 5 min to 6 hours. Hydrodynamic radius and PDI values of PLGA–NaC (A–D), PLGA–PVA–CS (B–E) and PLGA–PVA (C–F). 1D-SDS–PAGE gel (G) of isolated HC PC from <sup>19</sup>F–PLGA–NaC NPs, <sup>19</sup>F–PLGA–PVA–CS and <sup>19</sup>F–PLGA–PVA NPs. "M" indicates the protein ladder reference, while "ctrl" is the control of the cell culture medium treated with the same procedure but without NPs.

effects during the overall experimental window. On the other hand, <sup>19</sup>F–PLGA–PVA NPs, stabilized by steric hindrance, showed the highest colloidal stability for both conditions, without effective interactions with serum proteins (Fig. 2C and F). For <sup>19</sup>F–PLGA–PVA–CS NPs, a slight increase of the hydrodynamic size was observed only in DMEM, indicating a stabilization effect of serum proteins (Fig. 2B and E).

Overall, we developed three PERFECTA loaded NPs with high yield of encapsulation, good colloidal stability and different patterns of interaction with the serum proteins as a function of their surface functionalization.

#### Protein corona characterization

In cell culture medium, <sup>19</sup>F–PLGA–NaC (Fig. 2A) showed a rapid increase of NP size, indicative of the formation of a protein corona (PC) due to the binding of medium-derived components. <sup>19</sup>F–PLGA–PVA–CS and <sup>19</sup>F–PLGA–PVA did not show such a relevant variation of size, while the increase of PDI values was related to the presence of the protein background.<sup>46,47</sup> The hard corona (HC) composition was then investigated upon NP incubation in cell culture medium supplemented with 10% FBS in the experimental window – from 5 min to 8 h – in accordance with cellular labelling conditions. SDS–PAGE gel analysis

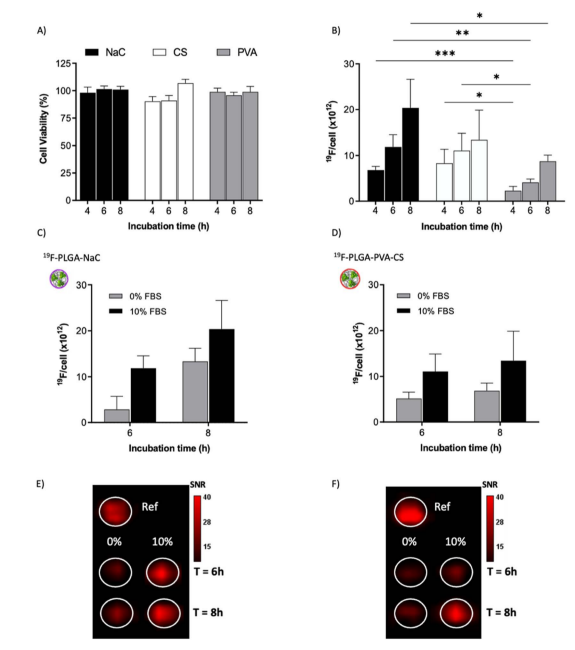
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indicated the formation of an extensive (HC) PC already within the first 5 min of incubation with <sup>19</sup>F–PLGA–NaC NPs (Fig. 2G). In the case of <sup>19</sup>F–PLGA–PVA–CS, few bands at 75 kDa becoming more intense with the incubation time were observed (Fig. 2G). In detail, PLGA–NaC NPs showed a rich PC profile with two more intense bands at 75 and 25 kDa. Of note, no PC formation was observed for PVA NPs at all incubation times, indicating the ability of PVA to prevent NP–protein interactions (Fig. 2G).

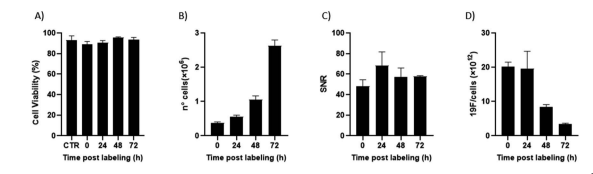
As expected, the PC composition was strongly dependent on the surface chemical composition of the NPs, increasing their complexity at the nano–bio interface.<sup>48–50</sup> Here, we demonstrate that <sup>19</sup>F–PLGA NPs coated with ionic stabilizers are more prone to form PC NPs than non-ionic analogues stabilized with PVA.

#### <sup>19</sup>F-PLGA NP cellular uptake studies

To test the cellular uptake and toxicity effects, microglial cells (BV2) were incubated with the <sup>19</sup>F-NP formulations for 4, 6 and 8 h, keeping constant the dose of fluorine content (*i.e.*, 1.54 mM PERFECTA corresponding to  $1 \times 10^{14}$  <sup>19</sup>F per cell). Under all



**Fig. 3** Labelling efficiency. (A) Cell viability of labeled BV2 cells (normalized to the cell survival of unlabeled cells, ctrl) evaluated by standard trypan blue staining after 4, 6 and 8 h of incubation with the different <sup>19</sup>F–PLGA-NPs. (B) <sup>19</sup>F cellular uptake as a function of incubation time measured by <sup>19</sup>F-MRI. (C and D) Labelling efficiency analyzed by <sup>19</sup>F-MRI in DMEM with or without FBS (0%; 10% FBS) for <sup>19</sup>F–PLGA–NaC (C) and <sup>19</sup>F–PLGA–PVA–CS (D) NPs. (E and F) <sup>19</sup>F-MRI phantoms of cells labeled with <sup>19</sup>F–PLGA–NaC (E) or <sup>19</sup>F–PLGA–PVA–CS (F) with or without FBS (0% or 10%, 250 000 cells per sample, ref. =  $5 \times 10^{19}$  fluorine atoms). All reported data are mean and standard error from 6 to 10 samples related to 3 to 5 independent experiments (n = 3 for 0% FBS).



**Fig. 4** Labelling stability during BV2 cell proliferation. Cell viability (A) and proliferation (B) of BV2 cells over time post labelling with <sup>19</sup>F–PLGA– NaC NPs. (C) Signal to noise ratio (SNR) related to the <sup>19</sup>F-signal from cell samples collected at each time post labelling. (D) <sup>19</sup>F content per cell decreased over time due to cell proliferation. All reported data are related to two independent experiments.

tested conditions, cell viability was similar to that of untreated cells (Fig. 3A), while <sup>19</sup>F labelling efficiency increased over time for all formulations. As expected, with the neutrally charged PLGA–PVA NPs, <sup>19</sup>F uptakes by cells were the lowest and limited over time (Fig. 3B), while an increasing trend and higher <sup>19</sup>F cellular uptake was observed for PLGA–NaC and PLGA–PVA–CS compared to PLGA–PVA NPs. Although the physical characteristics are almost similar, the main differences between <sup>19</sup>F–PLGA–PVA NPs and <sup>19</sup>F–PLGA–NaC or <sup>19</sup>F–PLGA–PVA–CS NPs are related both to the <sup>19</sup>F loading in single NP (lowest for <sup>19</sup>F–PLGA–PVA) and to the interaction with serum proteins, with formation of a rich PC for <sup>19</sup>F–PLGA–NaC and <sup>19</sup>F–PLGA–PVA–CS NPs.

Interestingly, the PC seems to enhance NP uptake by cells that consequently increase the <sup>19</sup>F-MRI signal and thus improve the detection sensitivity (Fig. 3C–F). Such a role of the PC in the cellular internalization process was supported by the lower uptake of NPs in the absence of serum, which resulted in a strong reduction of the <sup>19</sup>F-MRI signal. Thus, the improved efficiency to label cells shown by NaC and PVA–CS coated NPs seems not only due to higher <sup>19</sup>F concentrations in single NPs, but in particular due to the bio–nano interface formed once NPs were in contact with the biological environment. Importantly, NaC coated NPs also showed a significant increase in size upon incubation with serum proteins, likely affecting the cellular NP internalization pathways.

Of note, <sup>19</sup>F-uptake was more variable at longer incubation times probably in relation to the active proliferation of BV2 cells. To further investigate this aspect, signal stability of BV2 cells was also monitored by <sup>19</sup>F-MRI from 24 to 72 h post labelling with <sup>19</sup>F–PLGA–NaC NPs. While cells rapidly proliferated as expected maintaining a good viability (Fig. 4A and B), <sup>19</sup>F signal remained stable over time although the collected cells were washed before imaging (Fig. 4C). Internalized NPs should be redistributed among dividing cells, as shown by the dilution of <sup>19</sup>F signal per cell (Fig. 4D). Cell culture supernatants failed to reveal significant <sup>19</sup>F signal (<1% of the total uptaken <sup>19</sup>F content, Fig. S4†) supporting redistribution of the internalized PERFECTA among dividing cells rather than release.

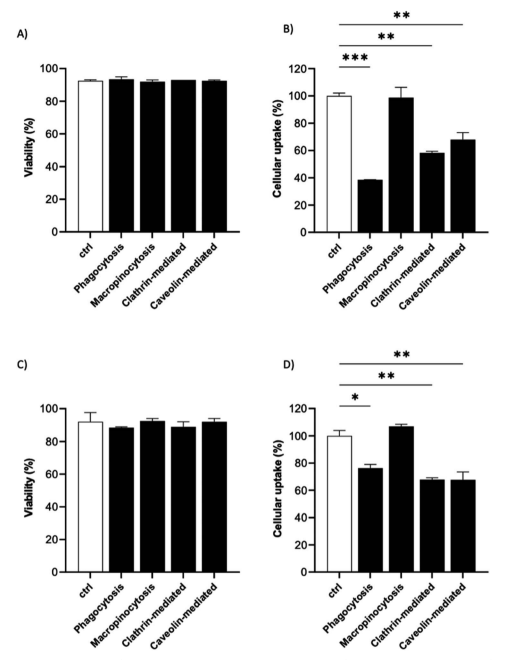
#### Cellular internalization mechanism studies

It is known that the active endocytic mechanism is a common process for NP uptake by cells,<sup>51</sup> but different pathways could be

involved as a function of NP size and surface properties. Considering the size range of the <sup>19</sup>F-PLGA-NPs (100-200 nm), phagocytosis and micropinocytosis pathways should be predominant.51,52 Using specific pharmacological inhibitors, we selectively evaluated the four main internalization pathways of phagocytic cells:51-54 phagocytosis, macropinocytosis, clathrinmediated micropinocytosis and caveolin-mediated micropinocytosis (details are reported in Table S2<sup>†</sup>). Optimal doses were chosen based on significant inhibition with a limited effect on cell viability (Fig. S5<sup>†</sup>). Inhibition assays were carried out on the most efficient formulations for cell labelling, namely <sup>19</sup>F-PLGA NPs formed with NaC or PVA-CS (Fig. 3B), and at short incubation time (4 h) to limit cell stress due to inhibitors which might alter the cell uptake mechanism. We used flow cytometry analysis to simultaneously assess cell viability (Fig. 5A and C) and NP uptake (Fig. 5B and D, reported singularly in Fig. S6<sup>†</sup>). To this end, fluorescent <sup>19</sup>F–PLGA NPs were prepared by including a small amount of poly(lactide-co-glycolide)rhodamine B in the formulation. While no effects were detected with the macropinocytosis inhibitor, those inhibiting phagocytosis and clathrin/caveolin-mediated uptakes significantly limited NP internalization (Fig. 5B and D). In the case of <sup>19</sup>F-PLGA-PVA-CS NPs, clathrin/caveolin mediated processes were equally involved with an inhibition of about 32% compared to untreated cells, and almost 25% for the phagocytic pathway. For <sup>19</sup>F-PLGA-NaC, phagocytosis was the predominant activated pathway, with a decrease of 61% in NP uptake followed by clathrin- and caveolin-mediated with 42 and 32% reduction, respectively. These differences in the uptake mechanisms could be expected considering that the size and composition of the protein corona were dependent on the pristine electrokinetic potential of the NPs.

In fact, with increased size and rich PC, it was expected that phagocytosis would have been the main mechanism in the case of <sup>19</sup>F–PLGA–NaC NPs.

Our results are in line with previous reports highlighting that perfluorocarbon loaded polymeric NPs with more hydrophilic domains are mainly internalized by clathrin-mediated endocytosis, while more hydrophobic NPs rely on different mechanisms.<sup>53</sup> Considering the overall results obtained, we suggest that these differences are mainly related to the interactions of the NPs with the biological media (formation of PC-NPs). Paper



**Fig. 5** Preferential endocytic pathways of <sup>19</sup>F–PLGA-NPs. The cell viability and uptake of <sup>19</sup>F–PLGA–NaC (A and B) and <sup>19</sup>F–PLGA–PVA–CS (C and D) NPs in BV2 cells were evaluated in the presence of pharmacological inhibitors of specific endocytic route and compared to untreated labeled cells (ctrl). Data were obtained by flow cytometry in independent experiments and expressed as follows: \*p < 0.05; \*\*p < 0.01; \*\*\*p < 0.001 (one way ANOVA test compared to the ctrl).

# Conclusions

Overall, our data indicate that a careful design of <sup>19</sup>F-nanoprobes together with a comprehensive analysis of their properties within biological fluids is needed to exploit <sup>19</sup>F-MRI as a valuable imaging method for cell tracking. Here we reported the development of biocompatible <sup>19</sup>F–PLGA-NPs containing PERFECTA and stabilized with different coating agents in order to modulate their surface properties and interaction with the biological environment and promote their cellular uptake. All the NPs were characterized by high <sup>19</sup>F loading and excellent <sup>19</sup>F-imaging

properties, but they showed distinctive affinities to serum proteins with formation of PC NPs of diverse size and composition as a function of their surface properties. In this sense, we demonstrated that the PC composition was pivotal to promote NP uptake by phagocytic cells as microglia and maximize <sup>19</sup>F-MRI response. Surface charged NPs, NaC and CS stabilized, showed a more significant interaction with serum proteins with consequently higher cell internalization and <sup>19</sup>F-MRI signals. The bio–nano interface should then be considered an essential element in the design of imaging nanoprobes for cell labelling, especially in the case of non-phagocytic cells. For example, in the

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challenging cases of labelling T-cells and stem cells, the modulation of the PC can be of major importance since NP internalization is limited due to the restricted intracellular space and poor phagocytic activity. In fact, while the formation of a PC is often avoided for *in vivo* administration of drug delivery agents (to avoid sequestration of the NPs by the RES and immune response activation), we strongly advocate its exploitation for *ex vivo* cell labelling, as it might be key for increasing <sup>19</sup>F sensitivity and consequently *in vivo* cell tracing by <sup>19</sup>F-MRI tracking. In this sense, the labelling procedure of therapeutic cells for clinical applications should be performed with human serum, with the possibility to use patient-derived serum. This approach would allow the formation of a "personalized protein corona", as it has been recently proposed for the use of different nanomedicines with diagnostic and therapeutic purposes.<sup>55–57</sup>

### Experimental

#### Materials

PERFECTA was synthesized as previously described.<sup>58</sup> Poly(lactic-*co*-glycolic acid) (PLGA; Resomer 502H, 50:50, 7–17 kDa), polyvinyl alcohol (PVA; 30–70 kDa, 87–90% hydrolyzed), sodium cholate (NaC), and ethyl acetate (EtOAc, purity  $\geq$  99.5%) were supplied by Sigma-Aldrich (Germany). Water ultrapure Type-I Milli-Q water (mQw) (18.2 m $\Omega$  cm<sup>-1</sup>) was obtained using a Simplicity® Water Purification System (MERCK, Germany). Chitosan (CS, 100–300 kDa; 85% deacetylated) was purchased from Fisher Scientific.

For cellular labelling, the immortalized murine microglial cell line (BV2) was kindly provided by Dr R. Furlan (Clinical Neuroimmunology Unit, IRCCS Ospedale San Raffaele). Cell culture medium DMEM (Lonza Bioscience, Italy) was supplemented with 10% fetal bovine serum (FBS, Lonza Bioscience, Italy), 100 mg ml<sup>-1</sup> of streptomycin, 100 U ml<sup>-1</sup> of penicillin and 2 mM of Lglutamine (Gibco-Invitrogen). Trypsin (Invitrogen) allowed the dissociation of adherent cells from culture dishes and their collection. Trypan blue exclusion assay (0.4% solution, Sigma-Aldrich, Germany) was used to evaluate cell viability. Paraformaldehyde (PFA, 4% v/v in water, Sigma-Aldrich, Germany) was used to fix cells before MRI analysis. For the inhibition of cellular uptake, phenylarsine oxide (PAO), filipin III, amiloride hydrochloride, chlorpromazine hydrochloride, Bodipy-LacCER and transferrin-Alexa Fluor 488 were supplied by Invitrogen and Thermo Fisher, while dextran-rhodamine B (44 kDa or 155 kDa) was provided by Sigma-Aldrich.

For flow cytometry (FCM) analysis a fluorescent version of the polymer was used. In detail, PLGA-rhodamine B (50:50, 10-30 kDa; Sigma-Aldrich) was used in the NP formulation at 2% w/ w with respect to unlabeled PLGA. Finally, the impact of NPs on cell viability was evaluated by 7-aminoactinomycin D dye as a live-dead marker (7-AAD; Invitrogen).

#### Preparation of <sup>19</sup>F-PLGA NPs

<sup>19</sup>F–PLGA NPs were formulated using an optimized protocol previously reported.<sup>15</sup> Briefly, PERFECTA was firstly dissolved in 1.5 ml of EtOAc and then heated at 65 °C. PLGA was dissolved in the same organic solvent (1.5 ml) and heated at 65 °C. The two solutions were combined and the resulting organic solution was rapidly mixed and added dropwise to the aqueous solution, containing either 0.6% w/v NaC or 2% w/v PVA, depending on the formulation type. For the NPs coated with CS, PVA (2% w/v) was added to an aqueous solution where CS was previously dissolved (CS/PLGA w/w = 0.8, see the ESI, Section 1.2†). The solution was then sonicated using a tip sonicator at 60 W for 25 s. Samples were left stirring for 3 h and the excess of organic solvent was then removed by rotary evaporation. NPs were collected and purified by centrifugation (10 733*g* × 14 min for <sup>19</sup>F-PLGA-PVA-CS NPs and 10 733*g* × 40 min for both <sup>19</sup>F-PLGA-PVA and <sup>19</sup>F-PLGA-NaC NPs). The final product was then resuspended in mQw.

#### Characterization of <sup>19</sup>F-PLGA nanoparticles

NP colloidal stability and morphology were evaluated by multiangle Dynamic Light Scattering (DLS, ALV/CGS-3 system, ALV-Laser Vertriebsgesellschaft.m.b.H., Germany). Hydrodynamic diameter, size distribution and polydispersity (PDI) were determined by a cumulant fitting of the auto-correlation function obtained through DLS measurements. For these analyses, the stock solution was diluted 1:100 (v/v) in water and measured at 25 °C. The colloidal stability of the NPs in a biological environment was assessed diluting each formulation 1:6 (v/v) in water, DMEM and DMEM + 10% FBS. Samples were incubated at 37 °C and measured at *t*<sub>0</sub>, 30 min, 1, 2, 4, 6, 24 and 48 h at a dilution 1 : 25 (v/v) in water through DLS measurements. The concentration of NPs was estimated using the Nanoparticle Tracking Analysis (NTA, Nanosight NS3000, Malvern Panalytical, Malvern, England), the measurements were done at a dilution of 1:5000 for <sup>19</sup>F-PLGA-NaC and 1:10000 for <sup>19</sup>F-PLGA-PVA and <sup>19</sup>F-PLGA-PVA-CS (v/v) in water. The electrokinetic potential was obtained through ζ-potential measurements (Zetasizer Nano ZS, Malvern Panalytical, Malvern, England) at a dilution of 1:5 (v/v) in water. Encapsulation efficiency of PERFECTA was determined from the quantification of <sup>19</sup>F atoms through <sup>19</sup>F-NMR experiments (Bruker AV400, Biospec; Bruker Biospin, Germany) with an external reference (TFA, trifluoroacetic acid). <sup>19</sup>F-MRI experiments were also performed to characterize and compare the MRI signal properties of each <sup>19</sup>F-PLGA-NP.

#### Cell labelling assays

Cellular uptake of <sup>19</sup>F–PLGA NPs were investigated using the immortalized murine microglial cell line (BV2). Cells were seeded at a density of  $3.3 \times 10^5$  cells per ml in a 6-well plate and incubated with 1.54 mM of PERFECTA (corresponding to  $1 \times 10^{14}$  <sup>19</sup>F per cell) at 37 °C and 5% CO<sub>2</sub> for 4, 6 and 8 h. Cells were then washed 3–5 times with supplemented culture medium in the well until complete removal of free NPs as assessed by light field microscopy. Labeled cells were then collected and washed in abundant culture medium (2 × 300*g* for 10 min). In addition, for determining the potential contribution due to the presence of free NPs that persist after repeated washing steps, cells incubated with NPs for few seconds ( $t_0$ ) were also included. As blank control, cells were treated with an equal volume of mQw

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and underwent the same experimental procedures. After labelling, cell viability was evaluated through trypan blue exclusion test. BV2 cells were washed in PBS and subsequently fixed with 4% PFA for 20 min and stored at 4 °C until MRI acquisitions.

For FBS-free labelling tests, BV2 were treated as previously described and the culture medium was replaced with FBS-free DMEM before incubation with NPs.

#### Labelling stability

To test the labelling stability upon incubation with <sup>19</sup>F–PLGA NPs (6 h with 1 × 10<sup>14</sup> <sup>19</sup>F per cell), BV2 cells were washed to remove the not internalized NPs and then were kept in culture. Cells were then collected at  $t_0$ , 24, 48 and 72 h. For each time point, cells were washed in abundant culture medium (2 × 300g for 10 min), and PBS (300g × 10 min), counted and fixed with 4% PFA. The collected cells were analyzed by <sup>19</sup>F-MRI. The cell culture supernatant was recovered and the <sup>19</sup>F content was evaluated by <sup>19</sup>F NMR, except for  $t_0$ .

#### Cell labelling inhibition assays

To study the endocytic pathways, phenylarsine oxide (PAO), amiloride chloride, chlorpromazine hydrochloride (CPZ) and filipin III were used to assess phagocytosis, macropinocytosis, and caveolin-mediated and clathrin-mediated pathways, respectively (see Table S2†). Each inhibitor was initially tested to determine the optimal dose using selective commonly used substrates, namely dextran–rhodamine B (44 kDa or 155 kDa) for phagocytosis and macropinocytosis, Bodipy–LacCER for caveolin-mediated and transferrin-Alexa Fluor 488 for clathrinmediated pathways. The inhibitory effect was assessed by flow cytometry as a decreasing fold in the mean fluorescence intensity (MFI) value compared to the uninhibited condition.

BV2 cells were seeded at a density of  $3.3 \times 10^5$  cells per ml in a 6-well plate and pre-incubated for 30 minutes with 0.0075, 400, 7.5 and 14  $\mu$ M of PAO, amiloride hydrochloride, filipin III and chlorpromazine hydrochloride, respectively. Fluorescent <sup>19</sup>F–PLGA NPs were added and incubated using a PERFECTA concentration equal to 0.84 mM for 4 h. Cells were then washed until the complete removal of free NPs, counted and analyzed by flow cytometry.

#### <sup>19</sup>F-magnetic resonance imaging

The relaxometric properties ( $T_1$  and  $T_2$ ) of <sup>19</sup>F–PLGA NPs were assessed by MRI. In detail, <sup>19</sup>F-MRI analysis was performed on all formulations at four different concentrations ranging from 6 to  $12 \times 10^{18}$  <sup>19</sup>F atoms in 100 µl of mQw. Each sample was loaded in 0.2 ml Eppendorf and placed into a phantom containing 2% agar gel.

The quantification of <sup>19</sup>F content in labeled cells was performed on pellets of cells, collected by centrifugation and placed in 0.2 ml tubes. The total amount of <sup>19</sup>F atoms was estimated using standards of PERFECTA formulations as reference  $(4-10 \times 10^{18} {}^{19}\text{F})$ .

All experiments were performed on a 7T scanner (Biospec; Bruker-Biospin) using a dual-transmit receive  $^{19}\text{F}/^{1}\text{H}$  volume coil (35  $\times$  59 mm) equipped with a horizontal bore magnet (ultra-

shielded). <sup>19</sup>F-MR images were acquired at PERFECTA specific resonance frequency (-73.4 ppm). For all MRI acquisitions, a 3D turbo-spin echo sequence was used with the same field of view  $(45 \times 30 \times 24 \text{ mm})$  for <sup>1</sup>H-MRI  $(T_{\rm R}/T_{\rm E} = 250/15 \text{ ms}, \text{matrix} = 128$  $\times$  64  $\times$  8, 6 averages) and for <sup>19</sup>F-MRI ( $T_{\rm R}/T_{\rm E}$  = 1500/40 ms, 64  $\times$  $32 \times 8$ , 32 averages). The relaxation times of <sup>19</sup>F–PLGA NPs were assessed using a RARE sequence with multiple echo times  $(10T_{\rm E})$ from 16 to 304 ms) and multiple repetition times ( $6T_{\rm B}$  from 350 to 1500 ms). For each sample,  $T_1$  and  $T_2$  were extrapolated from the fitting of the signal intensity as a function of  $T_{\rm E}$  or  $T_{\rm R}$ , respectively, using the scanner software tools (ISA tools Bruker Paravision 6). For each sample, the <sup>19</sup>F signal to noise ratio (SNR) was calculated by measuring the signal in each sample using the image analysis tool of the scanner (Paravision 6.0, Bruker-Biospin) and divided by the standard deviation of the noise (image area outside of the sample).

#### Protein corona characterization

The protein binding on the NP surface was investigated through the incubation of NPs in DMEM supplemented by 10% FBS (Biowest; South America) for different times from 5 minutes to 8 hours at 37 °C and 95% of humidity. NPs were diluted 1 : 6 v/v in order to mimic the experimental conditions of a cell labelling experiment. After incubation, NP-protein complexes were isolated by sedimentation through a sucrose cushion (CARLO ERBA Reagents; Italy) adapting a protocol that was already investigated.59 In detail, NP-protein samples were loaded over the sucrose cushion (0.7 M). The mixture was then centrifuged through the cushion in order to isolate NP-protein complexes from unbound proteins (10733  $\times$  g, 4 °C for 30 minutes). Isolation was followed by three washes in mQw water and resuspensions in 50 µl of mQw. The last supernatants and protein-NP pellets were analyzed by DLS (at 90°) to check the successful isolation and purification.

#### **SDS-PAGE electrophoresis**

A qualitative determination of hard-corona (HC) patterns was investigated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis. All reagents were purchased from Bio-Rad, Sigma-Aldrich. In detail, 5 µl of SDS-PAGE loading buffer (Bio-Rad, 65.8 mM Tris-HCl, pH 6.8, 2.1% SDS, 26.3% (w/v) glycerol, 0.01% bromophenol blue), supplemented with  $\beta$ -mercaptoethanol (1:20) was added to 15  $\mu$ l of each sample and kept at 99 °C for 5 minutes for digestion. Digested HC samples and control (ctrl) were loaded in the wells of SDS-PAGE gels. Ctrl consisted of a sample of FBS that underwent the same purification protocol. The gel was composed of a 4% polyacrylamide stacking gel (125 mM Tris-HCl, pH 6.8, 0.1%, m/v, SDS) and a 12% resolving polyacrylamide gel (in 375 mM Tris-HCl, pH 8.8, 0.1%, m/v, SDS buffer). A Tris-glycine buffer at pH 8.3 (with 0.1% SDS, m/v) was used as the cathode, whereas a Tris buffer at pH 8.8 represented the anode. Electrophoresis was performed in three steps with increasing voltage: the first step was set at 50 V for 20 min, second step at 100 V for 40 min and third step at 150 V until the dye front reached the bottom of the gel. Staining and distaining

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were performed with Colloidal Coomassie Blue (Serva) and 7% (v/v) acetic acid in water, respectively. Finally, the SDS-PAGE gels were scanned with a VersaDoc imaging system (Bio-Rad).

#### High-angle annular dark field scanning transmission electron microscopy (HAADF-STEM) supplemented by energydispersive X-ray (EDS) analysis

Scanning transmission imaging was performed using a STEM-HAADF and EDS in an FEI aberration corrected scanning transmission electron microscopy (Cs-corrected STEM) TITAN electron microscope operating at 300 kV. Samples for electron microscopy-analysis were prepared by drop casting 5  $\mu$ l concentrated particle solution onto a carbon coated TEM-grid and allowed to dry at room temperature overnight. Nanoparticle size distribution was determined with ImageJ software.

#### NMR experiments

For each formulation, <sup>19</sup>F-loading was quantified through <sup>19</sup>F-NMR experiments with an external standard, more precisely a sealed capillary containing 100 µl of 0.35 M trifluoroacetic acid (TFA) in D<sub>2</sub>O. All NMR spectra acquisitions were performed on a Bruker Avance III HD 400 spectrometer (Bruker BioSpin, Rheinstetten, Germany) using a Bruker Automatic Sample Changer (SampleXpress). <sup>19</sup>F-NMR spectra were recorded in the range of -65 to -85 ppm compared to TFA (-75 ppm). The NMR probe was maintained at 300 K for the entire duration of the experiment. A delay time between repetitions of 14.5 s was adopted to ensure full relaxation and 256 scans were collected. All <sup>19</sup>F-NMR spectra were automatically phased and baseline corrected for accurate quantitative analysis through MestReNova software. Each set of NPs was analyzed by <sup>19</sup>F-NMR for fluorine quantification. In detail, <sup>19</sup>F-NMR spectra were performed on NP stocks using TFA reference as previously described. From the ratio of integrals of the peaks, the fluorine content was calculated.

#### Flow cytometry analysis

<sup>19</sup>F–PLGA NPs were formulated with 2% of PLGA–rhodamine B to make NPs detectable by flow cytometry. For inhibition tests, cells were stained with the 7-AAD dye to assess their viability. The inhibition of NP uptake was evaluated as decreasing fold of the mean fluorescence intensity (MFI) compared to the uninhibited conditions.

All flow cytometry analyses were performed using a FACS Canto II (BD Biosciences) with the following laser configuration: 488 nm laser and 585/42 filter (detecting rhodamine B), 488 nm laser and 530/30 filter (detecting FITC and Alexa Fluor 488) and 488 nm laser and LP670 filter (detecting 7-AAD). Post-processing analyses were performed using the FlowJo software (BD Biosciences).

#### Statistical analysis

All measurements reported in the present work are given as the average ( $\pm$ standard deviation) of at least n = 3 replicates for each condition. Cellular experiments were run at least twice, and each assay contained two replicates (n = 4 measures/ condition). Data were plotted in Prism Software (GraphPad

Software 9) and statistical significance was calculated by oneway ANOVA followed by Dunnett *post hoc* test. Statistical significance was considered for p < 0.05.

### Author contributions

L. C. and F. B. B. designed and supervised the research work. L. G., C. C., L. C. and F. B. B. designed the experimental work. L. G., C. C., M. S. A., V. S. and P. M. performed the experimental work. G. R. supervised flow cytometry acquisitions and helped to analyse the obtained results. L. G., C. C., L. C. and F. B. B. wrote the manuscript with the support of all other authors. All authors discussed the results and contributed to the final manuscript.

# Conflicts of interest

There are no conflicts to declare.

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