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Introduction

Drug delivery for the treatment of central nervous system (CNS) diseases has been substantially improved since the advent of nanotechnology. Indeed, the CNS is a difficult target for therapies delivered by the systemic route due to the comprehensive vascular organization of a physical and chemical barrier (the blood brain barrier, BBB) and to the absence of a classical lymphatic.¹ This is the scenario faced in most neurodegenerative diseases, including progressive phases of multiple sclerosis (MS), a chronic immuno-mediated demyelinating disease of the CNS, in which the BBB is restored, or closed, while

Myelin-specific T cells carry and release magnetite PGLA–PEG COOH nanoparticles in the mouse central nervous system †

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Progress in nanotechnology has determined new strategies concerning drug delivery into the central nervous system for the treatment of degenerative and inflammatory diseases. To date, brain targeting through systemic drug administration, even in a nano-composition, is often unsuccessful. Therefore, we investigated the possibility of loading T lymphocytes with PGLA–PEG COOH magnetite nanoparticles (30 nm), which can be built up to easily bind drugs and monoclonal antibodies, and to exploit the ability of activated T cells to cross the blood–brain barrier and infiltrate the brain parenchyma. Iron oxide nanoparticles have been widely used in biomedical applications due to their theranostic properties and are therefore a well-established nanomaterial. The magnetite core is easily hybridized with polymeric compounds that may enhance the possibility of the nanoparticles entering cells with low phagocytic properties. Taking advantage of these material characteristics, after *in vitro* assessment of the viability and functionality of nano-loaded MOG_{35–55} specific T cells, we transferred cells containing the nanocargo into naïve mice affected by experimental autoimmune encephalomyelitis, an animal model of multiple sclerosis. By means of histological and immunohistological methods, we were able to identify the nano-loaded T cells in the central nervous system. Our data demonstrated that T cells containing nanomaterials hold the possibility of carrying and releasing nanoparticles in the brain.

compartmentalized inflammatory cell reactions are ongoing and triggering neurodegenerative processes.² Localized clearing of the immune reaction and cure are dependent on the efficiency of CNS parenchyma penetration by the administered drug; therefore intrathecal delivery of immunosuppressants is a promising new approach for this more malign phase of the disease.3 For instance, Komori et al. recently investigated the effects of intrathecally-administered rituximab, a monoclonal antibody directed against CD20 that had no effect on clinical outcomes in MS progressive forms if delivered by the peripheral route.⁴ A decade of experience demonstrates that CNS direct drug administration is a possibility for anti-spastic therapies: for example, in patients resistant to orally administered Baclofen, intrathecal delivery was shown to be a good strategy in order to obtain an improvement in MS spasticity.5 We think that in MS, aside from the design of new drugs, the development of technologies promoting brain drug uptake in order to achieve therapeutic levels is becoming more and more crucial, as direct intrathecal delivery minimizes side effects and limits invasive procedures.

In recent years the development of nanotechnologies has provided a means for crossing the BBB and targeting the inflamed MS brain; many laboratories have developed nanocarriers for drug delivery and nano-vaccines constitute an

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efficient strategy to promote immune tolerance.⁶ Furthermore, nanomaterials have the ability to combine diagnostic with therapeutic approaches:⁷ ultra-small magnetite nanoparticles (USPIO) have been used as contrast medium in order to enhance MRI identification of CNS newly formed lesions in MS by visualizing macrophage cell infiltrates and different patterns of lesions.^{8,9} Lesions in MS have been classified into 4 different patterns¹⁰ depending on the composition of cell infiltrate: demyelination, remyelination, oligodendrocyte loss and BBB damage. To date, there is a large ongoing effort in order to correlate histology data and neuroimaging data. We think that the future development of a MRI protocol able to spot NBR-loaded T cells in MS active lesions may lead to the identification of those lesions characterized by the presence of T cells.

In the present work, we demonstrated that nanoparticles internalized by immune cells might be a promising strategy in order to deliver therapeutics into the brain. More specifically, we utilized nanoparticles feasible for future conjugation with drugs and exploited the ability of T lymphocytes to be loaded, to carry, and to cross the BBB and release these nanoparticles into the CNS. Our experimental design is based on antigen-specific (MOG₃₅₋₅₅) mice T cells efficiently loaded with nanobeads (PGLA-PEG-COOH plus a magnetite core; NBR) and then transferred i.v into C57Bl/6 mice, naive or during the chronic phase of the animal model of MS -experimental autoimmune encephalomyelitis (EAE). Our research demonstrates that T cells loaded with NBR are able to migrate and effectively infiltrate the CNS tissue, where most importantly they were found in demyelinating lesions. Furthermore, we showed that in the CNS, the presence of utilized magnetite nanoparticles was identifiable by magnetic resonance imaging (MRI). Altogether, these findings provide exciting opportunities regarding the future of direct CNS MS lesions treatment.

Experimental

Assembling nano-shells: NBR and NBR-Fluo

Magnetite (Fe₃O₄) nanoparticles (NP^{Fe}) were synthesized as previously reported (Patent: Baldi et al., Magnetic Nanoparticles functionalized with cathecol, production and use thereof, WO/ 2015 104 664).11 Briefly, 40 mL of NPFe suspended in diethylene glycol was sonicated for 1 hour in an ultrasound bath and then a solution of [N-(3,4-dihydroxyphenethyl)dodecanamide (DDA)] (1.09 g, 3.2 mmol) dissolved in 120 mL of ethanol was slowly added to the glycolic suspension. The system was again sonicated for 1 hour at room temperature. The solution was then diluted with 60 mL of ultrapure water, magnetically sedimented and dispersed in 90 mL of THF (Fe₃O₄-DDA). Then 440 milligrams of PGLA-b-PEG-COOH (43-3 kDa, 9.6 µmmol) was dissolved in 10 mL of THF and then added to 90 mL of Fe₃O₄-DDA suspension (440 mg of Fe_3O_4). Fluorescent nanoparticles (Fe₃O₄-DDA-Fluo) were achieved by adding a DMSO solution of Fluorescein 488-NHS (synthesized as reported within the ESI, Fig. S1 and methods†), properly derivatized with 4-aminobutylphosphonic acid, to the Fe₃O₄-DDA suspension. The suspension was left reacting overnight at room temperature. The formation of NBR was achieved by the nanoprecipitation

method: the organic phase was mixed with 900 mL of ultrapure water under vigorous stirring, maintaining a water : organic ratio of 10 : 1 with constant removal of the resulting solution. The mixture was then purified by washing with a phosphate-buffered solution and concentrated (Pellicon XL, cut-off 500 kDa) to a final volume of 150 mL (concentration: 0.3% (w/w) in Fe₃O₄); it was then filtered through a syringe filter (Millipore Sterivex, 0.22 μ m, polyethersulfone membrane). Fluorescent NBR (NBR-Fluo) were prepared using the same method by substituting Fe₃O₄-DDA with Fe₃O₄-DDA-Fluo.

Experimental animals

Female C57BL/6 mice (Envigo RMS Srl, Italy) were bred in the CeSaL animal house of the University of Florence, in Makrolon cages, with free access to food and water and kept at 23 °C with a 12 hours light/dark cycle. Every manipulation of the animals was executed according to the guidelines of the European community for animal care (DL 116/92, application of the European Communities Council Directive 86/609/EEC) and approved by the Committee for Animal Care (DGSAF, Italy, Authorization no. 514-2016-PR).

EAE induction

EAE was induced in 8 to 10 week-old female mice by s.c. injection of 200 mg myelin oligodendrocyte glycoprotein $(MOG)_{35-55}$ peptide (MEVGWYRSPFSRVVHLYRNGK, purity 85%; Espikem, Florence, Italy) in PBS, emulsified with an equal volume of complete Freund's adjuvant (CFA, Sigma) supplemented with 3 mg mL⁻¹ *Mycobacterium tuberculosis* H37Ra (Difco Laboratories, USA). *Bordetella pertussis* toxin (300 ng) was administered by i.p. injection at days 0 and 2 post immunization (Calbiochem, Germany). EAE-induced mice were scored daily for clinical signs of disease.

Generation of MOG₃₅₋₅₅ T cell lines (TCL)

Lymphomonocytes were isolated from the spleen and by draining lymph nodes of immunized mice at 8 days postimmunization (dpi), by mechanical dissociation on nylon filters with 70 µm pores (BD Falcon), followed by density gradient centrifugation (Biocoll, Biochrom AG). Cells were seeded in 96-well U bottom plates (BD Falcon) at a density of 2 $\times~10^5$ per well with 10 $\mu g~mL^{-1}$ of MOG_{35-55} in complete medium (RPMI 1640 + 10% fetal bovine serum + 1% penicillinstreptomycin + 1% sodium pyruvate + 1% L-glutamine + 1% Hepes buffer, Sigma). A week later, cells were stimulated with 20 IU mL⁻¹ of IL-2 (InvivoGen). After another 7 days, the experiment proceeded with the proliferation assay, according to the "split-well" method: 100 µL of volume from the first plate was transferred to a second plate, divided into two volumes (50 µL per well). In one of the wells, we added 50 µL of MOG₃₅₋₅₅ and in the other one, 50 μ L of medium only. Then, 1 \times 10⁵ irradiated (6000 rad) spleen cells were placed in both. After 72 h of incubation at 37 °C, 5% CO₂, 1 μ Ci of ³H-thymidine (Perkin Elmer) was added into each well. Then, the cells were transferred to a glass fibre filter for the proliferation analysis with the Cell Harvester (TomtecMacIII). The proliferation activity was measured as counts per minute (cpm) with a Trimux Microbeta 1450 counter (Wallac). We considered the test positive when Δ cpm was >1000 cpm (Δ cpm = cpm of stimulated wells – cpm of not-stimulated wells) and the stimulation index (SI) > 2 (SI: cpm of stimulated wells/cpm of not-stimulated wells).

Loading of MOG₃₅₋₅₅ TCL with NBR nanoparticles

MOG₃₅₋₅₅ TCL were incubated with NBR or NBR-Fluo, (0.3%) at a density of 2×10^6 cells per mL in medium (RPMI 1640 + 10%) fetal bovine serum + 1% penicillin-streptomycin + 1% sodium pyruvate + 1% L-glutamine + 1% Hepes buffer; all from Sigma) at 37 °C for 2 h; where indicated, NBR-Fluo loading was carried out in parallel at 4 °C. Then, cells were washed 3 times and suspended in fresh medium for the next analysis. NBR-Fluo uptake by TCL was assessed by flow cytometry, evaluating the percentage of Fluorescein 488-positive cells and the related mean fluorescence intensity (MFI). MFI of samples loaded at 4 °C and 37 °C were compared by overlaying the corresponding FL-1 histograms; the number of events of all compared histograms were normalized based on the maximum peak value using the specific formatting histograms option of FCS Express6 software (DeNovo software, USA). For each sample, we acquired the same number of events. Dead cells, labelled with propidium iodide (2.5 μ g mL⁻¹, Molecular Probes), were excluded from the analysis.

Subcellular evaluation of NBR loaded TCL

The 1×10^6 NBR-loaded TCL were centrifuged at 1500 rpm for 5 min at RT, the supernatant was discarded and the cell pellet was fixed in Karnovsky solution; then it was embedded in epoxy resin. Samples were sectioned with an ultramicrotome in 50-60 nm slices. The sections were observed with TEM (Jeol, Japan). TEM analysis was performed on TCL immediately after NBR loading and upon 24 hours of stimulation in *vitro* with anti-CD3 ε antibodies 1 µg mL⁻¹ (clone 17A2, eBioscience). To investigate the NBR-loaded T cell shape, a further evaluation was performed on 2×10^6 TCL loaded with NBR-Fluo. Cells were seeded on positively charged glass slides (Bioptica), fixed with PFA 4% (Bioptica) for 15 min at RT, then washed with PBS and permeabilized with PBS/FBS 5%/Triton 0.3% solution for 30 min at RT in the dark. F-actin distribution was evaluated in TCL loaded with NBR-Fluo, or not, by labelling with AlexaFluor 633 phalloidin (Thermo Fisher Scientific) for 20 min at RT in the dark, and then the slides were washed three times in PBS and closed with glass coverslips using mounting medium containing DAPI (Prolong Gold with DAPI, Life Technologies). Images were acquired using a Leica DMIRE2 confocal microscope (Leica Microsystems, Germany) equipped with a He/Ne/Ar laser source, using a Leica Plan Apo x 63/1.40 NA oil immersion objective. A series of optical sections (1024 imes 1024 pixels each; pixel size 200 imes200 nm) were taken at intervals of 0.3 µm. Confocal images were analysed using ImageJ software (National Institutes of Health [NIH], Bethesda, MD).

Cell viability

NBR-loaded TCL were analysed for cell viability immediately after loading and upon *in vitro* stimulation for 24 and 72 hours with MOG_{35-55} 10 µg mL⁻¹ in complete medium. Cell viability was assessed by flow cytometry using the Annexin V Apoptosis Detection Kit FITC (eBioscence, U.S.A.) and the CyFlow Space cytometer (Sysmex-Partec, Germany).

Proliferation and cytokine production of MOG_{35-55} TCL loaded with NBR

Proliferation activity of TCL loaded with NBR was tested by ³Hthymidine incorporation test as previously described,¹² after *in vitro* stimulation with 10 μ g mL⁻¹ of MOG₃₅₋₅₅. Before adding ³H-thymidine, we collected 50 μ L of supernatant from each well for cytokine analysis. We measured IL-17, IFN γ , IL-4 and IL-10 production by Bioplex (Biorad), using the commercial kit Milliplex (MCYTOMAG-70K, Millipore) following the manufacturer's protocol.

Expression of activation markers

TCL, loaded or not with NBR, were stimulated with MOG_{35-55} (10 µg mL⁻¹) and analyzed after 12 and 24 hours in order to evaluate the expression of CD25 (activation) and CD62L (a key regulator of migration, early-modulated during cell activation). Cells were stained (20 minutes RT in the dark) with the following fluorescent anti-mouse monoclonal antibodies (all from eBioscience): anti-CD25 PE (1 : 100 diluted), anti-CD62L APC (1 : 600 diluted), anti-CD3 ϵ FITC (1 : 100 diluted). Then the cells were analysed with a CyFlow Space cytometer (Sysmex-Partec, Germany).

NBR release

NBR-Fluo-loaded MOG₃₅₋₅₅ TCL were incubated as previously described and then plated in 6-well plates coated, or not, with anti-CD3 ϵ antibodies (clone 17A2, eBioscience; 1 µg mL⁻¹) at a density of 600.000 cells/600 µL in complete culture medium. Cells were collected at baseline immediately after NBR-Fluo incubation (T0) and at different time points after loading: 2, 4, 6, 18, 24, 48 and 72 hours later (T2-T4-T6-T18-T24-T48-T72) upon in vitro culture with or without CD3 stimulation. Unstained cells were used as a negative control. The NBR-Fluo release was evaluated by flow cytometry (CyFlow Space cytometer, Sysmex-Partec, Germany) following the decrease of Fluorescein 488 mean fluorescence intensity (MFI) over time. The analysis was performed using FCS Express 6 software (De Novo Software, USA). In parallel, we determined the quantity of Fe_3O_4 in the cellular pellets by ICP-OES (Varian 710 ES). At the same time, we evaluated the percentage of NBR-Fluo-positive cells and the related MFI by fluorescence microscopy. Briefly, cells were seeded (10⁴ cells per spot) on positively charged glass slides, fixed and permeabilized as previously described. Slides were closed with aqueous mounting medium with DAPI and observed using a fluorescence microscope (Olympus BX600, Japan). MFI was quantified by ImageJ software, measuring the optical density of the green fluorescence conjugated to NBR in

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15 randomly selected fields chosen from 3 different experiments.

Passive transfer of NBR-loaded $\mathrm{MOG}_{\mathrm{35-55}}$ TCL in EAE and naïve mice

NBR loaded TCL were diluted in saline buffer at a density of 1×10^7 cells per mL and 100 µL of cell suspension was injected i.v. in C57BL/6 naïve mice (8 weeks old) and in EAE mice at 16 dpi, soon after the acute phase. We used 4 animals per group and we replicated the experiments three times. After 48 hours from passive transfer, mice were sacrificed and the brain and spinal cord were explanted in order to check the presence of NBRloaded T cells by histology/immunohistochemistry and post mortem MRI.

Histology and immunohistochemistry

Brain and spinal cord samples were fixed with 4% PFA, paraffinembedded and sectioned with a microtome in 5 µm slices. Slices were stained with Hematoxylin and Eosin (H&E), Luxol Fast Blue (LFB) and Perls, following standard protocols, or selected for immunofluorescence labelling. Briefly, slices were treated with citrate buffer (citric acid 10 mM, pH = 6; Sigma) in a thermostatic bath at 99 °C for 20 min and washed in hot water at 99 °C for antigen retrieval, permeabilized with PBS/FBS 5%/Triton 0.3% (Sigma) for 30 min at RT, labelled overnight at 4 °C with antimouse CD3ɛ antibodies (rabbit polyclonal, Abcam), and then diluted in permeabilization buffer 1:100. Then, slices were incubated for one hour in the dark at RT with the secondary antibodies AlexaFluor 488 anti-rabbit IgG (Life Technologies), and diluted 1:400 in permeabilization buffer. After 3 washes with PBS, the slices were closed with glass coverslips using mounting medium containing DAPI (Prolong Gold with DAPI, Life Technologies). Slides were observed by light or fluorescence microscopy (Leica Microsystems, Germany).

Results and discussion

Production and characterization of magnetite nanobeads (NBR)

TEM imaging of nanoparticles is reported in Fig. 1A. Furthermore, we reported SEM-FEG-STEM imaging of the inorganic core and surface (Fig. 1B and C). The mean hydrodynamic diameter and mean polydispersity index for NBR was determined by dynamic light scattering, and found to be 44.1 (\pm 1.6) nm and 0.17 \pm 0.02, respectively, indicating a narrow size distribution. No major differences were detected for NBR-Fluo which showed an average hydrodynamic diameter of 41.0 (\pm 0.6) nm and PDI = 0.13 \pm 0.02. The ξ _potential of the two assemblies of particles was -41 mV and -38 mV, respectively, for NBR and NBR-Fluo, confirming their stability. NBR-Fluo show an emission band centered at 520 nm, typical of the fluorescent fluorophore (see ESI[†]).

Subcellular evaluation of MOG₃₅₋₅₅ TCL loaded with NBR

Upload of NBR was qualitatively determined by TEM. Furthermore, we analysed the ultrastructure of NBR-loaded T cells,



Fig. 1 Imaging of magnetite PGLA–PEG COOH nanoparticles. (A and B) TEM imaging of nanoparticles. (C) SEM-FEG-STEM imaging of the inorganic core and surface.

immediately after loading with nanoparticles (Fig. 2A(a and b)) and 24 hours later, upon in vitro stimulation with anti-CD3 antibody (Fig. 2A(c and d)). TEM micrographs show that the cells are healthy; the nanoparticles are distributed in the cytoplasm, mainly close to the cell membrane, and a few are spread within the nucleus; indeed, they are mostly concentrated in the cytoplasm, where they appear organized in the vesicle-like structures. During cell division, nanoparticles were in part transferred into the daughter cell (Fig. 2A(c and d)). To evaluate the possibility that NBR determines changes in T cell shape, we compared, using confocal microscopy, actin distribution (red) in control unloaded cells (Fig. 2B, first row panel) and cells loaded with NBR-Fluo (green) (Fig. 2B, second row panel). The figure shows that NBR does not affect cell shape, which remains circular, and there is a partial colocalization between the NBR-Fluo and actin (Fig. 2B, panel "merge"). Cell actin, compared to unloaded cells, showed a redistribution after NBR loading. In order to evaluate if loading of the NBR in T lymphocytes is a passive, or in part an active, event, we performed the NBR-Fluo loading protocol at 37 °C and 4 °C. As we have shown, at 4 °C, when metabolic activity is reduced or there is no activity, loading of NBR is reduced in terms of quantity (MFI is threefold diminished), but loading still occurs and leads to 100% of cells being positive for NBR-Fluo (Fig. 2C).

In vitro analysis of NBR-loaded MOG₃₅₋₅₅ TCL: viability, proliferation, cytokine production, and activation marker expression

First, we checked the viability of NBR-loaded TCL after 2 hours of incubation with nanoparticles and after *in vitro* stimulation with MOG₃₅₋₅₅ for 24 and 72 hours (Fig. 3A). We did not find any significant variation in apoptotic and necrotic cell percentage in NBR-loaded TCL (black columns) compared to the controls (white columns).

The proliferative response to MOG_{35-55} stimulation was comparable in NBR-loaded TCL and controls (Fig. 3B, left). When we measured cytokine production, we detected a significantly decreased level of IFN γ and IL17 (*p < 0.05; **p < 0.01) after MOG_{35-55} stimulation in NBR-loaded TCL compared to controls (Fig. 3B, right). Furthermore, we analysed the effect of



Fig. 2 Subcellular evaluation of NBR-loaded MOG₃₅₋₅₅ TCL. (A) TEM analysis of MOG₃₅₋₅₅ TCL after incubation with iron-oxide NBR nanoparticles (a and b) or 24 hours later (c and d). Micrograph "(d)" represents a higher magnification of the image "(c)" (red square), showing a cell under division. Slice thickness: 50-60 nm. (B) MOG₃₅₋₅₅ TCL loaded with NBR-Fluo analysed by confocal microscopy: NBR-Fluo (green) and Factin (red) (left panel). F-Actin distribution in control cells is reported in the right panel. One experiment is representative of three independent ones. Scale bar: $10 \ \mu$ m. (C) Flow cytometry evaluation of NBR-Fluo incorporation by MOG₃₅₋₅₅ TCL at 37 °C (red histogram) or 4 °C (black histogram); grey histogram represents the intrinsic auto-fluorescence of unstained control cells. The numbers of events in the overlay histograms were normalized based on the maximum peak value. One experiment is representative of three independent ones.

NBR uptake on the cell surface expression of the activation markers CD25 and CD62L, both normally present after stimulation. CD25 is the IL2 receptor and CD62L is a lectin involved in cell migratory ability; they were evaluated upon MOG_{35-55} stimulation for 12 or 24 hours (Fig. 3C). The flow cytometry analysis indicated that activation of NBR T cells induces the expected up-regulation of CD25 and retains CD62L, and therefore T cells have the phenotype of normal active cells able to patrol the CNS parenchyma.

In vitro analysis of MOG₃₅₋₅₅ TCL NBR uptake and release

Before transfer of NBR-loaded TCL *in vivo*, we investigated for *in vitro* NBR release, by means of MFI determination in NBR-Fluoloaded cells and chemical determination of cell pellet Fe_3O_4 content. MFI value was measured (Fig. 4A) by flow cytometry immediately after NBR-Fluo cell incubation (T0) and after 2, 4, 6, 18, 24 and 72 hours; the green histograms in Fig. 4A show a reduction of MFI value from baseline to 72 hours of incubation compared to the control (unstained cells, white histogram). The evaluation of spontaneous or stimulationinduced release of nanoparticles by NBR-loaded TCL through measuring the Fe_3O_4 content gave the same result (Fig. 4B). After loading 2×10^6 antigen-specific T cells with NBR, we examined the cellular pellets, comparing them with the amount of Fe₃O₄ contained in an equal number of cells immediately after NBR loading. As shown in Fig. 3C, the Fe₃O₄ level decreased by half in the first 24 hours of culture, both in the stimulated cells and in the non-stimulated ones and continued to decrease slightly between 24 and 48 hours of culture, then remained stable between 48 and 72 hours. On the other hand, fluorescence microscopy (Fig. 4C) of the NBR-Fluo (green), and T cells nuclei (DAPI, blue), shows that NBR-Fluo are maintained in almost 60% of the antigen-specific T cells from baseline to 72 hours of incubation. This determination was performed by observation of 5 areas per slide (3 slides for each time point), containing 25 cells per area for a total of three independent experiments.

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Fig. 3 Viability, proliferation, cytokine production and early activation markers of NBR-loaded TCL. (A) Analysis of the viability of NBR-loaded MOG_{35-55} TCL compared to not-loaded cells (CTR) at different time points and experimental conditions: immediately after NBR loading (upper panel) and 24 and 72 hours later, upon *in vitro* stimulation with MOG_{35-55} peptide (middle panel) or cultured with medium alone (lower panel). Mean value \pm SEM of three different experiments. PI = propidium iodide. (B) Left panel: antigen specific proliferative response of NBR loaded MOG_{35-55} TCL determined by ³H-thymidine incorporation assay and expressed as Δ cpm (mean cpm observed in MOG_{35-55} stimulated cells – mean cpm in not stimulated cells; cpm = counts per minute). Mean value \pm SEM of three experiments. Right panel: cytokine profile of NBR-loaded MOG_{35-55} TCL compared to not-loaded ones (CTR); IFN γ , IL17 and IL10 production was evaluated by Milliplex assays in supernatants of antigen-stimulated cells. Mean value \pm SEM of three experiments. (C) Flow cytometry analysis of early activation markers expression: CD25 (left) and CD62L (right) in MOG_{35-55} TCL loaded with NBR and not loaded (CTR) after 12 hours and 24 hours of *in vitro* culture with MOG_{35-55} peptide and in the absence of stimuli (–). Column graphs report the percentage of CD3 positive cells expressing CD25 and CD62L, respectively. Mean values \pm SEM of three independent experiments.

Passive transfer of NBR loaded TCL into naïve mice

After assessing viability, proliferative ability and phenotype of MOG_{35-55} TCL containing NBR *in vitro*, we studied *in vivo* ability of NBR-loaded TCL to migrate in CNS after i.v. cell transfer in naive animals, where BBB is intact. As reported in the methods section, mice were sacrificed 48 hours after the passive transfer. Fig. 5 illustrates Perls staining and reveals the presence of cells containing magnetite (blue cells) in the spinal cord (panel A) and brain sections (panels B and C). Focusing on brain tissue, we observed the presence of magnetite-positive cells spread in the hippocampus region (B) and within the cerebellum in the proximity of the fourth ventricle (C).

Passive transfer of NBR-loaded TCL into EAE mice

We then investigated whether NBR-loaded TCL were able to migrate into injured CNS tissue in EAE mice and if there was any colocalization with demyelinated lesions. The spinal cord designated to the histopathological analysis underwent a MRI investigation, and results (ESI, Fig. 2†) were in agreement with Perls staining. In Fig. 6 we show the histological analysis in the thoraco-lumbar region of the spinal cord; within this area we observed infiltrates (H&E) and demyelination (LFB), together with the presence of magnetite-positive cells (yellow arrows, Perls), identified in consecutive slices also as T cells by anti-CD3 immunofluorescence staining (white arrows).

Therapeutic delivery to the CNS during MS is challenging; direct lateral ventricular injections have been one option, although highly invasive and not feasible for treatments requiring frequent drug administration.⁴ Injections into cerebrospinal fluid are also a possibility, but diffusion in the brain parenchyma remains restricted.13 The direct use of nanomaterials able to physically interact with epithelial cells and shown to cross barriers, for example the BBB, could be a strategy in order to project new nano-drug formulations.14-16 Nonetheless, the absence of specificity of this interaction leads to a possible reduced efficacy in treatment of CNS diseases. Here, we decided to target CNS with autoreactive T cells, loaded with magnetite nanoparticles encapsulated in a PGLA-PEG-COOH copolymer (NBR) planned to easily cross the plasma membrane and to bind proteins, as antibodies. NBR dimensions make them suitable to cross membranes and internalize into cells following different pathways,17 becoming a cargo that may be transported and delivered in the brain. Activated T



Fig. 4 Evaluation of nanoparticles release from NBR-loaded MOG_{35-55} TCL. (A) Green histograms show the mean fluorescence intensity (MFI) evaluated by flow cytometry of MOG_{35-55} T cells incubated with NBR-Fluo (Fluorescein 488) at different time points after loading: immediately after NBR loading (T0), and 2, 4, 6, 18, 24, 48 and 72 hours later (T2-T4-T6-T18-T24-T48-T72). Unstained cells were used as a negative control (white histogram). (B) Iron-oxide content expressed in µg of Fe₃O₄ measured by ICP-OES in MOG_{35-55} TCL 24, 48 and 72 hours after NBR loading, upon *in vitro* culture in the presence (black curve) or absence of CD3 stimulation (grey curve). Mean values \pm SEM of 3 experiments. (C) Upper panel: pictures of the same NBR-Fluo-loaded T cells analysed in panel A, taken using a fluorescence microscope at 4 different time points: T0-T6-T24-T72. Green: NBR-Fluo; blue: DAPI. Scale bar: 20 µm. One experiment is representative of 3 independent ones. Lower panel: quantification of the green fluorescence intensity in the NBR-Fluo-loaded T cells shown in the pictures. Column graph reports the average value \pm SEM of green fluorescence optical density measured by ImageJ in 15 randomly selected fields chosen from 3 different experiments (**p* = 0.017; ***p* = 0.0045, Mann-Whitney *t* test).

lymphocytes cross the BBB and patrol brain parenchyma and, more important, depending on their antigen specificity, they will persist in the CNS with a peak at day 5 post-injection.¹⁸ The primary goal of this study was to investigate if NBR loaded CD4+ T lymphocytes, autoreactive for myelin antigens, were vital, functional and able to reach the CNS with an intact BBB and during the chronic phase of EAE.

We described how NBR are able to enter antigen-specific T lymphocytes without altering their shape and migratory properties, but partially modifying their cytokine profile towards a less inflammatory pattern. Electron microscopy imaging suggests that NBR locate in vesicle-like structures, and confocal microscopy shows that NBR are in close contact with actin, which resulted in a different distribution in loaded cells compared to controls. In previous work, we have demonstrated that actin cytoskeleton modifications regulate proinflammatory effector functions in myeloid dendritic cells;^{19,20} here, we may speculate that a similar mechanism, driven by actin organization after NBR loading, is acting on T cells and is partially responsible of IL17 and IFN γ down-regulation. In contrast with other reports concerning bare magnetite nanoparticles (5–30 nm),²¹ NBR in part enters the nucleus, suggesting that this PGLA-covered nanomaterial is able to passively diffuse through all the membranes. Our data on cell loading at 4 °C indicates



Fig. 5 Detection of magnetite-positive cells within the CNS of naïve mice 48 hours after passive transfer of MOG_{35-55} TCL loaded with NBR. (A) Consecutive spinal cord sections (5 μ m thickness) stained with Perls (blue: iron-positive cells; red: nuclei; pink: cytoplasm) and immunolabeled with anti-CD3 fluorescent antibodies (green = CD3+ T cells) and DAPI (blue = nuclei). Red rectangle maps a region where we found correspondence between Perls-positive cells and CD3 immunolabeling. (B and C) Perls staining revealed the presence of magnetite-positive cells (arrows) in the hippocampus region (B) and in the cerebellum, in the proximity of the fourth ventricle (C). Sections are observed at different magnifications as indicated (2.5×; 5×; 20×; 40×). Data are representative of three independent experiments.

that the mechanism of NBR cell entry is mostly passive, although an active component, an atypical macropinocytosis reported on HeLa cells loaded with SPION,²² might be present.

Our *in vitro* experiments demonstrated that nano-loaded cells start to gradually release the cargo during the first 24 hours. The apparent discrepancy between data from flow cytometry/chemical measures of NBR and fluorescence microscopy suggests that these cells are spontaneously releasing a part of the nano-cargo, but not all of it. Probably the flow cytometry and chemical analysis methods that we used for NBR content determination were not sufficiently sensitive, beneath a certain threshold of fluorescence, for a proper quantification of NBR-positive cells by flow cytometry. Notwithstanding the MFI data, fluorescence microscopy shows around 60% of cells still carrying NBR, even if to a lower extent, 72 hours after loading. Proliferation does not increase the spontaneous release of the nanomaterial, and we may

hypothesize that during the proliferative response the magnetic particles are diluted in daughter cells, as suggested by an equal amount of NBR-positive cells in stimulated and not-stimulated protocols. These kinetics makes the system suitable for transport and delivery of therapies (e.g. anti CD20) designed to be effective in the periphery and in the CNS. In this perspective, we demonstrated that cells loaded with NBR cross the BBB in naive animals, where barrier permeability and integrity are normal, and reach the brain in 48 hours. Therefore, in naive mice, we confirmed that NBR-loaded cells were able to reach the brain parenchyma. Of note, the number of injected cells was far below the usual threshold that induces BBB damage.23 On the other hand, thanks to their antigen specificity and activation state, they may stay and interact with neural cells.24 To further challenge this experimental delivery system, we injected NBRloaded T cells during the chronic phase of EAE, the animal model of multiple sclerosis. In these experiments, we suggest



Fig. 6 Histological analysis of EAE spinal cord after the passive transfer of NBR-loaded TCL. Histological analysis at the thoracic-lumbar region of spinal cord. Consecutive spinal cord slices (5 μm thickness) were stained with Hematoxylin/Eosin (HE) to reveal infiltrating immune cells (grey arrows), Luxol Fast Blue (LFB) to highlight demyelinating lesions (red arrows), Perls to detect the presence of magnetite-positive cells (yellow arrows) and anti-CD3 fluorescence immune-labelling to identify the presence of T cells (green, white arrows). Scale bars: 20 μm.

the possibility of tracing NBR cells, due to their ability to generate detectable signals in MRI, as previously described.²⁵ Furthermore, the presence of NBR/cells in the CNS was confirmed by histology and immunohistochemistry; we found transferred cells containing NBR and NBR alone, probably released by the cells, also in demyelinated inflammatory areas 48 hours after injection. In fact, by this time antigen-specific T lymphocytes directed to a CNS antigen should be between the CNS perivascular space and the parenchyma, in an intact BBB scenario, or directly in the parenchyma when BBB permeability is altered.²⁶

Conclusions

To the best of our knowledge, the use of T cells as carriers of nanoparticles has been previously assessed in cancer therapy, where nanoparticles were reversibly attached to activated CD8+ T cell surfaces^{27,28} and in autoimmunity, where poly(ethylene glycol)-functionalized hydrophilic carbon clusters specifically target T lymphocytes in the spleen *in vivo* and act as an anti-oxidant.²⁹ Therefore, this is the first step towards the development of nano-loaded self-antigen-specific T lymphocytes as a therapy delivery system in the context of brain autoimmune disease.

Conflicts of interest

Cericol Research Center Colorobbia, Italy, provided nanoparticles, named NBR, used in this work.

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