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Multicolor spectral photon counting CT monitors and quantifies therapeutic cells and their encapsulating scaffold in a model of brain damage

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Abstract

Rationale & aim: Various types of cell therapies are currently under investigation for the treatment of ischemic stroke patients. To bridge the gap between cell administration and therapeutic outcome, there is a need for non-invasive monitoring of these innovative therapeutic approaches. Spectral photon counting computed tomography (SPCCT) is a new imaging modality that may be suitable for cell tracking. SPCCT is the next generation of clinical CT that allows the selective visualization and quantification of multiple contrast agents. The aims of this study are: (i) to demonstrate the feasibility of using SPCCT to longitudinally monitor and quantify therapeutic cells, i.e. bone marrow-derived M2-polarized macrophages transplanted in rats with brain damage; and (ii) to evaluate the potential of this approach to discriminate M2-polarized macrophages from their encapsulating scaffold.

Methods: Twenty one rats received an intralesional transplantation of bone marrow-derived M2-polarized macrophages. In the first set of experiments, cells were labeled with gold nanoparticles and tracked for up to two weeks post-injection in a monocolor study via gold K-edge imaging. In the second set of experiments, the same protocol was repeated for a bicolor study, in which the labeled cells are embedded in iodine nanoparticle-labeled scaffold. The amount of gold in the brain was longitudinally quantified using gold K-edge images reconstructed from SPCCT acquisition. Animals were sacrificed at different time points post-injection, and ICP-OES was used to validate the accuracy of gold quantification from SPCCT imaging.

Results: The feasibility of therapeutic cell tracking was successfully demonstrated in brain-damaged rats with SPCCT imaging. The imaging modality enabled cell monitoring for up to 2 weeks post-injection, in a specific and quantitative manner. Differentiation of labeled cells and their embedding scaffold was also feasible with SPCCT imaging, with a detection limit as low as 5,000 cells in a voxel of $250 \times 250 \times 250 \ \mu m$ in dimension *in vivo*.

Conclusion: Multicolor SPCCT is an innovative translational imaging tool that allows monitoring and quantification of therapeutic cells and their encapsulating scaffold transplanted in the damaged rat brain.

Key words: multicolor spectral photon-counting CT, regenerative medicine, cell therapy, cell tracking, neurology

Introduction

Stroke is the leading cause of long-term disability with no effective cure after the first hours of onset. Various types of cell therapies are currently

being evaluated in many clinical trials involving patients with ischemic stroke (1). A meta-analysis of preclinical studies found that the intracerebral administration is associated with the best treatment efficacy compared to other administration routes, as they can thus act locally by secreting neurotrophic factors for example (2-4). Recently, two independent clinical trials reported the safety of intracerebral administration of different types of therapeutic cells in stroke patients (5, 6). The infarct cavity allows the injection of a large volume of cells; however, it is a hostile environment for cell engraftment. To improve the integration of the grafted cells, these cells can be embedded within a bioengineered scaffold (7). To bridge the gap between the administration of regenerative therapy and the clinical trial outcome, the regulatory agencies that approve advanced-therapy medicinal products (ATMPs) have highlighted the need for non-invasive imaging tools (8). The challenge is to propose a solution that involves clinically relevant imaging techniques, in order to foster the clinical translation of these innovative theranostic approaches.

In human trials of cell tracking, nuclear medicine imaging, such as positron emission tomography (PET) and single photon emission computed tomography (SPECT), is mainly used (9). However, the relatively short half-lives of radiotracers prevent the long-term tracking of administrated cells. Gene reporter imaging with PET or SPECT may overcome this problem (10), although to date these technologies remain costly and not widely available. Magnetic resonance imaging (MRI) and X-ray computed tomography (CT) are the leading three-dimensional (3D) radiological technologies around the world. Coupled with contrast agents, both modalities are regarded as the imaging techniques that can potentially track administered cells up to several weeks. To render cells detectable, a contrast agent is incorporated into cells prior to transplantation. Superparamagnetic particles of iron oxide (SPIO) and gold nanoparticles (AuNPs) are examples of widely studied contrast agents for MRI and CT, respectively (11, 12). One limitation of both these approaches is the lack of specificity of the signal produced by the labeled cells, which makes it difficult to non-equivocally distinguish the signal of labeled sources hyperdensities cells from other of (microhemorrhages or calcifications for instance). A recent breakthrough in CT technology has been the introduction of dual-energy CT (DECT) into clinical CT scanners. DECT has several advantages compared to conventional CT, including better contrast-to-noise ratio and fewer artifacts (13). Importantly, after the administration of an iodinated contrast agent, DECT allows the reconstruction of non-contrast and contrast-enhanced images based on a single acquisition, eliminating the need for pre-scan. This leads to X-ray dose reduction without altering the

diagnostic value. The development of spectral photon-counting detectors rather than energy-integrating detectors has extended the DECT approach to a multispectral approach. This next generation spectral photon counting CT (SPCCT) is expected to further expand the boundaries of spectral CT in terms of noise reduction and tissue differentiation (14). It also improves the intrinsic spatial resolution owing to small pixel detector size (~250 µm) (15, 16). In addition, SPCCT allows selective visualization and quantification of multiple contrast agents by exploiting the K-edge discontinuity in X-ray absorption, a concept coined 'K-edge imaging' or 'multicolor CT' (17). The first reports of bicolor imaging with SPCCT have demonstrated the potential of this innovative approach (18-23), with many future applications yet to be developed.

This study aims to: (i) demonstrate the feasibility of using SPCCT to longitudinally monitor and quantify AuNP-labeled therapeutic cells transplanted into brain-damaged rats; and (ii) evaluate the potential of this approach to discriminate the labeled cells from their encapsulating scaffold, both labeled with different types of contrast agents. Bone marrow-derived M2-polarized (repair) macrophages were used as the therapeutic cells (24, 25). The commercially-available scaffold PuraMatrix® was employed to encapsulate the cells (26, 27). Stereotaxic injection into rodent brains was chosen as a mean to model intracerebral administration in humans. A SPCCT prototype system (Philips Healthcare, Haifa, Israel) with small field of view (16.8 cm) was used for image acquisition (28). This system has five thresholds that could be adjusted in order to allow photon energy-based discrimination of any given element that has a K-edge in the range of 30 - 120 keV.

Materials and Methods

Animals

All experimental procedures involving animals and their care were carried out in accordance with the European regulation for animal use (APAFIS agreement number #4688). This study was approved by the local ethics committee of our institution (C2EA – 42, local ethic board). Adult male Sprague-Dawley rats (Janvier, France) that were 6 to 7 weeks-old at reception and with a body weight in the range of 250-300 g and C57B1/6 mice (Janvier, France) that were 5 weeks-old at reception and with a body weight in the range of 25-30 g were used. An acclimation period of at least 7 days was respected. Animals were housed in a temperature- and humidity-controlled environment (21 \pm 3 °C), on 12:12 h light-dark cycle, having free access to standard chow and tap water. Housing Pexiglas cages were bedded with wood dust and enriched with tunnels and wood sticks (rats) or domes (mice). The data are reported according to the ARRIVE guidelines (Animal Research: Reporting of *In vivo* Experiments).

Contrast agents

Gold nanoparticles (AuNPs)

11-Mercaptoundecanoic acid capped gold nanoparticles (11-MUDA AuNPs) were synthesized via a previously reported adaptation of the Turkevich method (29, 30). In brief, 85 mg of gold (III) chloride salt was dissolved in 500 ml of ultrapure water and brought to a boil while stirring. 25 mL of sodium citrate (38.8 mM) was added and the solution allowed to boil for additional 15 minutes before cooling to room temperature. A wine-red solution of gold nanoparticles resulted from this procedure. To cap the gold nanoparticles, 2.6 mg of 11-MUDA dissolved in 1 mL of ethanol was added, and the solution was stirred overnight. The resulting 11-MUDA AuNPs were purified by centrifuging them three times at 8.5 krcf and exchanging the supernatant with ultrapure water each time. AuNPs were then sterilized via syringe filtration (size: 0.45 µm) before further use. These nanoparticles had the following characteristics: peak absorbance of 524 nm, average hydrodynamic diameter of 22 nm with PDI of 0.2, core size of 11 ± 1 nm, and zeta potential of -44.4 mV.

lodinated nanoparticles (INPs)

Concentrated aqueous suspensions of INPs were prepared in two steps of emulsification and concentration as follows (31). The iodinated polymer TIB-PVAL was a 2,3,5-triodobenzoyl ester of poly(vinyl alcohol) containing 70 wt% of iodine. The emulsification of TIB-PVAL in water was performed by mixing 25 mL of 4 wt% TIB-PVAL in THF and 50 mL of deionized water. A block copolymer polycaprolactone-*block*-poly(ethylene glycol) was used as an emulsifier. After evaporation of THF under reduced pressure, the resulting aqueous suspension of INPs had a concentration of 15 mg iodine/mL. As the second step, the iodine concentration was increased up to 100 mg(I)/mL by means of centrifugation and redispersion of the pellet into a small volume of water. INPs had the following characteristics: peak absorbance of 237 nm, hydrodynamic diameter of 122 nm with PDI of 0.2, and core size of 100 ± 15 nm.

Cell culture and cell labeling

Bone marrow cells were harvested by flushing out tibias and femurs of donor animals. Total bone marrow cells were seeded on uncoated 6-well plates at 5×10⁵ cells/mL in RPMI 1640 with Glutamax (Fisher Scientific, Illkirch, France) supplemented with 10% fetal calf serum (SVF premium, Dutscher, France), 1% penicillin-streptomycin Fisher Scientific, Illkirch, France) and 25 ng/mL of mouse macrophage-colony stimulating factor (mM-CSF) (Miltenyi Biotec, Paris, France). The cell cultures were incubated at 37 °C in humidified 5% CO₂. Seven days after plating, non-adherent cells were harvested and adherent cells were washed twice in phosphate-buffered saline (PBS). The harvested bone marrow-derived macrophages were then stimulated with interleukin-4 (20 ng/mL) (Miltenyi Biotec, Paris, France) for 48 h to generate M2 alternatively-activated macrophages. Macrophage cell culture purity under these experimental conditions was repeatedly found to exceed 95% on flow cytometry analysis of CD11b+ cells (data not shown). At day 6, cells were incubated with AuNPs at 0.1 mg/mL for 18 h, based on a protocol that provided high AuNPs uptake while maintaining cell viability (29). Immediately after labeling, cells were detached by trypsination, washed once in PBS, counted and resuspended in Ca2+-Mg2+ sterile PBS for in vivo injection. AuNP internalization and cell morphology post-labeling were assessed by light microscopy. The viability of the AuNP labeled cells was examined using the LIVE/DEAD assay (Invitrogen, Carlsbad, CA, USA). The efficiency of labeling was determined based on inductively coupled plasma-optical emission spectrometry (ICP-OES) using three different bone-marrows and triplicates for each bone-marrow.

Scaffold and scaffold labeling

Puramatrix (3-D Matrix, MA, USA) is a synthetic peptide that undergoes self-assembly into nanofiber hydrogels similar to the extracellular matrix upon introduction of monovalent cations in physiological conditions. PuraMatrix thus provides a suitable biological scaffold for cell transplantation, and it has been used for central nervous system regeneration (27, 32). For injection of AuNPs-labeled cells in INPs-labeled scaffold, the total volume of injectable solution (10 μ L) was composed of PuraMatrix (1/4 of 10 μ L), INPs solution (1/8 of 10 μ L) and AuNPs-labeled cells in PBS (remaining volume).

In vitro studies

The accuracy of quantification in SPCCT material imaging has been demonstrated previously by phantom imaging of iodine, gold and their mixture with other contrast agents (17). In this study, thirteen samples were prepared by suspending the gold or iodine nanoparticles in 1% agarose gel placed in Eppendorf tubes with a range of concentrations: 0, 10,

15, 20, 30, 40 and 60 mM for INPs and 0, 10, 15, 20, 30, and 40 mM for AuNPs (thus resulting in the same range of 0 – 8 mg/mL for each material). These phantoms were scanned at each imaging time point for calibration purpose in longitudinal *in vivo* studies (see below). In addition, to evaluate the performance of SPCCT quantification in our experimental setting, labeled cells pellets were prepared in the same conditions as for the *in vivo* administration, i.e. 10 μ L of PBS with decreasing quantity of cells: 1 x 10⁶, 0.5 x 10⁶, 0.25 x 10⁶, 0.125 x 10⁶ and no cells, placed at the bottom of Eppendorf tubes and secured with 1% agarose gel on top.

In vivo studies

Overall protocol

Figure 1 shows the experimental design of *in vivo* studies. In order to generate a lesion cavity that mimics stroke infarct at the chronic stage, 21 rats (experimental units) received an intracerebral injection of lipopolysaccharide (LPS) (33). To evaluate the exact lesion location for cell administration, rat brains were imaged 2 weeks later by T₂-weighted MRI. Subsequently, the rats received an intra-lesional injection of different therapeutic materials (e.g. cells \pm scaffold) as detailed thereafter. For this set of experiments, rats were randomly assigned to injection with or without scaffold. In the first set of experiments, cells were labeled with gold nanoparticles (AuNPs) and tracked up to two weeks post-injection using K-edge imaging of gold (monocolor study, n=13 LPS-treated rats). For the second set of experiments, the same protocol was repeated for a bicolor study, in which the labeled cells are embedded in iodine nanoparticle (INP)-labeled scaffold (bicolor study, n=8 LPS-treated rats). In both cases, the success of transplantation was determined

by performing μ CT immediately after intracerebral cell administration. The μ CT images are also shown as images of reference for accurate cell location. Sample size was determined in order to obtain enough data for this proof-of-concept study.

Animal models and cell transplantation

Animals were anesthetized by breathing 3-4% isoflurane (ISO-VET, Piramal Healthcare, Morpeth, UK) in air and maintained under 1.5% isoflurane in air using a face mask and mounted in a stereotaxic apparatus (D. Kopf Instruments). Isoflurane anesthesia was selected since it allows modulating anesthesia depth. Rectal temperature was kept at 37 ± 1 °C throughout the surgical procedures, using a feedback-regulated heating pad. To alleviate pain, buprenorphine was administered subcutaneously at the dose of 0.05 mg/kg prior to the surgery. Lubricant ophthalmic gel was applied on both eyes to prevent ocular dehydration, and the surgical site was shaved and disinfected with antiseptic solution (i.e. 10 % iodopovidone). To mimic stroke infarct, rats were stereotaxically injected with 50 µg of LPS from Escherichia coli (Sigma-Aldrich, Saint-Louis, USA) dissolved in 4 µL saline in the right hemisphere of the brain (0.5 mm anterior to bregma, 3.0 mm in lateral direction, 2 µL at 5.5 mm and 2 µL at 4.5 mm deep from the cortical surface). In addition, one mouse was submitted to ischemic stroke by permanent middle cerebral artery occlusion (pMCAO) according to a protocol published elsewhere (34). Briefly, the right distal middle cerebral artery was exposed by subtemporal craniotomy occluded and bv electrocoagulation. AuNP-labeled cells (in the range of 0.125 x 10⁶ to 0.5 x 10⁶) were intracerebrally transplanted in 10 µL vehicle (PBS or scaffold) inside the lesion, using stereotaxic coordinates based on

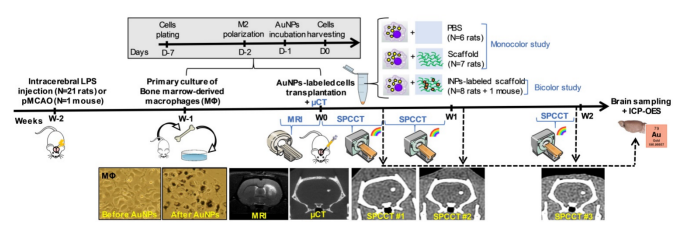


Figure 1. Experimental timeline for longitudinal multimodal imaging of brain-damaged rats transplanted with cell therapy. AuNPs: gold nanoparticles; ICP-OES: inductively coupled plasma-optical emission spectrometry; INPs: iodinated nanoparticles; LPS: lipopolysaccharide; µCT: micro-computed tomography; MRI: magnetic resonance imaging; pMCAO: permanent middle cerebral artery occlusion; SPCCT: spectral photon counting computed tomography.

pre-surgery MRI for each animal. At the end of the surgery, the wounds were cleaned with sterile saline solution, disinfected, and covered with lidocaine. Animals were allowed to completely wake up from anesthesia under a heating lamp and then housed individually for approximately 24 hours after the surgery. The surgical wound appearance was visually checked for correct healing.

Monocolor study

Thirteen LPS-treated rats were intracerebrally administered with 0.5×10^6 AuNP-labeled cells encapsulated either in PBS (n=6) or in scaffold (n=7). These rats were imaged with SPCCT three times during the first 2 weeks of post-transplantation period - once in the first half of the first week (Day 0-3), once in the second half of the first week (Day 4-7), and once at the end of the second week (Day 13-14). Subgroups of rats were sacrificed at each time point to compare the gold content estimated with SPCCT analysis with the gold content measured by ICP-OES (n=2 at Day 0-3, n=7 at Day 4-7 and n=4 at Day 13-14).

Bicolor study

To evaluate the feasibility of bicolor imaging, two LPS-treated rats and one pMCAO mouse were intracerebrally administered with 0.5 x 10⁶ cells encapsulated in 10 μ L iodine-labeled scaffold and then imaged twice during the first week of post-administration (once in Day 1-2 and once in Day 4-6). Lastly, to evaluate the limits of detection for AuNP-labeled cells in the context of bicolor imaging, 6 rats received a range of cell quantities, i.e. 0.5 x 10⁶ (n=2), 0.25 x 10⁶ (n=2) or 0.125 x 10⁶ (n=2). The iodine quantity in the scaffold did not vary between groups. These rats were imaged twice during the first week (at Day 1-2 and Day 4-6) and once at the end of the second week (Day 12-14).

In vivo multimodal imaging

All imaging procedures were performed under 1-2% isoflurane anesthesia. System characteristics and acquisition parameters for MRI, µCT, CT and SPCCT are presented in Table 1. The SPCCT scanner is described in details in previous studies (17, 28). In brief, it is based on a clinical iCT platform (Philips Healthcare, Haifa, Israel), operating with a standard X-ray source and energy-resolving photon counting detectors made of cadmium-zinc telluride, with 5 energy thresholds that can be tuned according to the materials of interest. Tube voltages can be set to 80, 100 or 120 kVp and tube currents from 10 to 100 mA. The effective z-collimation is 2.5 mm in the isocenter, in-plane field of view is 168 mm, Z-coverage is 2 mm and minimum rotation time is 0.75 sec/rotation. We acquired images using both axial and helical acquisition modes at a tube current of 100 mA and a tube voltage of 120 kVp with the energy thresholds of 30, 53, 78, 83 and 98 keV. For reconstruction, the image chain used a two-step material decomposition approach that first generates material specific sinograms for each material of interest (e.g. gold, iodine, gadolinium, water), which is followed by reconstruction of each sinogram into specific material image, depending on the user's choice. Taken together, the specific images are generated by best fit of the measured count rates in the five energy bins available to calibration data from phantom imaging. A maximum-likelihood based material decomposition (35) based on literature data of the attenuations (36) of the target materials was applied to derive material sinograms. Further details are provided in previous studies (17, 22).

Table 1. System characteristics and acquisition parameters for each imagin	ing modality.
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	Computed tomography	(CT)	
	μCT	СТ	SPCCT
System	Siemens INVEON	GE Brightspeed	Philips prototype
Tension (kVp)	80	120	120
Current (mA)	500	150	100
In-plane resolution (µm)	56	310	250
Slice thickness (µm)	56	625	250
Energy thresholds (keV)	N/A	N/A	30, 53, 78, 83, 98
Acquisition time (sec)	792	3	17
	Magnetic resonance imaging (MRI)		
System	Bruker Avance II	Sequence	T2 - RARE factor 8
Field strength	7 T	TE/TR (ms/ms)	57.7/5000 ms
Gradients	440 mT/m	In-plane resolution (µm)	117
Software	ParaVision 5.1	Slice thickness (µm)	800
Coils diameters (mm)	15 / 25	Acquisition time (min)	4

μCT: micro-computed tomography; RARE: rapid acquisition with relaxation enhancement; SPCCT: spectral photon counting computed tomography; TE: echo time; TR: repetition time.

Image analysis

Image analysis was performed using ImageJ software (National Institute of Health, USA, imagej.nih.gox/ij/). The gold concentration in mg/mL was quantified by measuring the signal intensity on gold K-edge images in the regions of interests (ROI). In order to compare the quantitative performances of SPCCT scans with ICP-OES (which provides the gold mass measured in the sample in μ g), the gold content was estimated by multiplying the concentration by the ROI volume.

Inductively coupled plasma-optical emission spectrometry (ICP-OES)

Animals were sacrificed at the end of the experiment, and their brains were sampled for ICP-OES analysis of gold content, according to a protocol published elsewhere (37). In brief, the harvested brain samples were homogenized before being digested in 1 mL of nitric acid at 75 °C for 24 hours. The samples were further digested for another 3 hours at 60 °C after addition of 300 μ L of

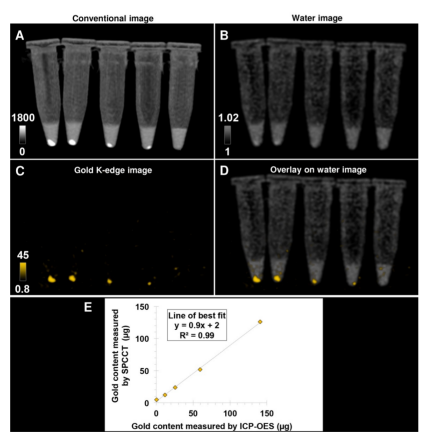


Figure 2. Pellets of AuNP-labeled macrophages at decreasing cell number imaged with a single SPCCT acquisition. (A) conventional image; (B) water image; (C) gold K-edge image; (D) overlay of gold K-edge image and water image; (E) Linear relationship between the gold content estimated by SPCCT on gold K-edge image and that measured by ICP-OES in each tube. From left to right: 1×10^6 , 0.5×10^6 , 0.125×10^6 and no cells. Color bars indicate Hounsfield units for conventional images and concentration in mg/mL for material images. CT: computed tomography; ICP-OES: inductively coupled plasma-optical emission spectrometry; SPCCT: spectral photon-counting computed tomography.

hydrochloric acid. 5 mL of deionized water was then added to 1 mL of the digested sample solution for ICP-OES analysis.

Statistics

Data are presented as mean ± standard deviation. Linear regression was used to correlate gold content estimated by SPCCT analysis with that measured by ICP-OES. The agreement between gold content estimated by SPCCT analysis and by ICP-OES was assessed with Bland-Altman analysis.

Results

For the purpose of illustration, Figure S1 shows the principle of bicolor imaging with SPCCT using a healthy rat that was intracerebrally injected with AuNP-labeled cells on one side and with gadolinium nanoparticles (GdNP)-labeled scaffold on the other side. The contrast agents could not be differentiated on conventional images, while gold K-edge images were able to detect AuNP-labeled cells in a non-equivocal manner. These GdNP were used in

another SPCCT project of our consortium; (38) however, we decided to use iodinated nanoparticles instead for this study because of the artifacts GdNP produce in MRI (Figure S1E).

In vitro studies

Bone-marrow derived M2polarized macrophages efficiently ingested AuNPs without any noticeable change in morphology compared to non-labeled cells in control group (Figure 1). AuNP uptake was confirmed by ICP-OES, with an average gold load per cell of 128 ± 34 pg. The cell viability was $86 \pm 1\%$ in the control group and 77 \pm 3% after labeling with AuNPs. In other words, the cell viability of AuNP-labeled cells was 89% the non-labeled cells. of Both conventional and gold K-edge images of cell pellets showed an observable increase in attenuation that was dependent on the number of cells in the pellets (Figure 2). The mean volume occupied by the cells in vitro was $2.8 \pm$ 1.9 µL. There was a strong linear relationship between the gold content measured on gold K-edge images by SPCCT analysis and that measured by ICP-OES in each cell pellet (R²=0.99). Gold content from K-edge images was slightly underestimated when

compared to the measurement from ICP-OES (slope of 0.9).

In vivo studies

Monocolor study

The conventional images reconstructed from SPCCT acquisition had significantly better resolution than the CT images acquired from a (conventional) clinical scanner (Figure S2). Gold K-edge images allowed a longitudinal monitoring of the brain distribution of AuNP-labeled macrophages in a non-equivocal manner (Figure 3).

One animal from the PBS group was inadvertently injected into the lateral ventricle and was therefore excluded from the quantitative analysis. This injection failure was clearly depicted by SPCCT despite cell dispersion (Figure S3). Thus 12/13 animals were included in the analysis (PBS: n=5, scaffold: n=7). Moreover, one rat did not get SPCCT data at Day 14 for technical reasons. Quantitative analysis of gold content from SPCCT images revealed a heterogeneity in the gold content that was present into the brain of each rat at the first scanning time point (i.e. Day 0-3 post-injection) (Figure 4A-B). In particular, one of the rats had a very low gold signal and cell volume (1.0 μ g in 0.5 μ L) (Figure 4B, dashed line), possibly due to suboptimal cell administration. The gold content remained relatively stable over time in both groups that had cells delivered in PBS (Figure 4A) and in scaffold (Figure 4B). The mean volume occupied by the cells *in vivo* was $2.7 \pm 1.0 \mu$ L. At the end of the experiment, the average gold content in the brain measured by ICP-OES was $48 \pm 18 \mu g$ (N=12). In comparison, SPCCT analysis provided an average

value of $44 \pm 23 \ \mu g$ in the last *in vivo* imaging time point for the same 12 animals. The animal with very low gold signal had only 7.4 μg of gold in its brain as measured by ICP-OES; thus confirming the injection problem that had been detected with SPCCT.

Figure 4C shows the Bland-Altman plot describing the agreement between the gold content estimated by SPCCT analysis and that measured by ICP-OES. In average, gold K-edge images underestimated gold content by 5 μ g (bias of 10%) with 95% limits of agreement of [-29; +19] µg. One animal with high gold value appeared to be an outlier (i.e. outside the limits of agreement). When excluding this animal from linear correlation analysis, the same relationship from the in vitro study was found between gold content from SPCCT analysis and from ICP-OES (Figure 4D, $R^2 = 0.80$, slope of 0.9).

Bicolor study

Conventional images did not allow the AuNP-labeled differentiation of cells from INP-labeled scaffold, while material images provided different gold and iodine locations, corresponding to the respective locations of cells and scaffold (Figure 5). As previously reported (28), there was some cross-talk between iodine and calcium, which resulted in appearance of the skull in the iodine image. As for monocolor imaging, bicolor imaging of gold and iodine allowed monitoring labeled cells and scaffold in the brain brain up to 2 weeks post-transplantation (Figure S4). Cells and scaffold monitoring was also feasible in the mouse brain with ischemic stroke (Figure S5).

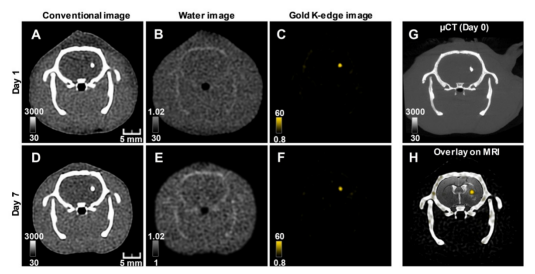


Figure 3. Longitudinal SPCCT imaging of a brain-damaged rat 1 day (A-C) and 7 days (D-F) post-transplantation of 0.5 x 10⁶ AuNPs-labeled macrophages. (A) & (D) conventional image; (B) & (E) water image; (C) & (F) gold K-edge image; (G) corresponding μ CT obtained on the day of injection; (H) overlay between baseline MRI (note the striatal lesion that appears as a hyperintense signal on T2-weighted imaging), conventional and gold K-edge images. Color bars indicate Hounsfield units for conventional images and concentration in mg/mL for material images. μ CT: micro-computed tomography; MRI: magnetic resonance imaging; SPCCT: spectral photon counting computed tomography.

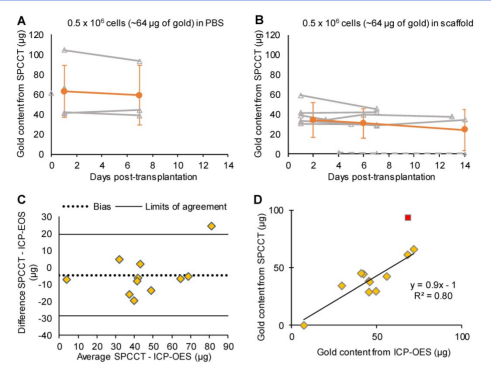


Figure 4. Monocolor study: quantitative analysis. (A) Graph plotting the average gold content estimated on gold K-edge images over time in the group of rats that received cells in PBS; (B) Graph plotting the average gold content estimated by SPCCT on gold K-edge images over time in the group of rats that received cells in scaffold; (C) Bland–Altman plot of gold content estimated by SPCCT on gold K-edge images and that measured by ICP-OES for each brain for the 2 groups; (D) Linear analysis of the same data; the red square represents an outlier that was excluded from linear regression. Grey lines represent individual animals and orange lines represent the mean ± standard deviation. Dotted line of graph B represents one animal that was excluded from analysis because of administration failure. ICP-OES: inductively coupled plasma-optical emission spectrometry; PBS: phosphate-buffered saline; SPCCT: spectral photon counting computed tomography.

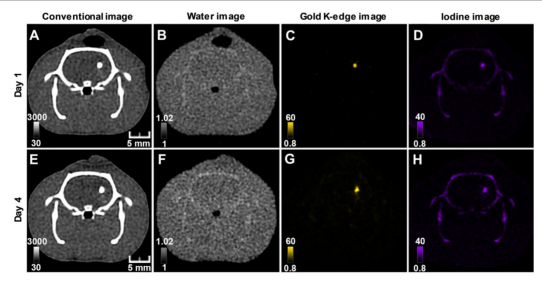


Figure 5. Longitudinal SPCCT imaging of a brain-damaged rat 1 day (A-D) and 4 days (F-I) post-transplantation of 0.5 x 10⁶AuNPs-labeled macrophages embedded in INP-labeled scaffold. (A) & (E) conventional image; (B) & (F) water image; (C) & (G) gold K-edge image; (D) & (H) iodine image. Color bars indicate Hounsfield units for conventional images and concentration in mg/mL for material images. SPCCT: spectral photon counting computed tomography; MRI: magnetic resonance imaging.

One animal died at Day 12 and was imaged *ex-vivo* for the last imaging point. All animals were included in quantitative analysis (n=9). The mean volume occupied by both cells and scaffold was $4.9 \pm 1.8 \mu$ L. In agreement with the monocolor study, the gold content that was present in the brain of each rat at the first scanning time point (i.e. Day 1-2 post-injection) was heterogeneous across animals and

remained stable afterwards (Figure 6). The mean volume occupied by cells on gold K-edge images was $1.4 \pm 0.3 \mu$ L. ICP-OES provided a value of $59 \pm 8 \mu g$ of gold in the brain of rats sacrificed at Day 4 of post-transplantation (injected with 0.5 x 10⁶ cells), compared to $72 \pm 4 \mu g$ estimated by SPCCT analysis at the same time point. The decrease in gold content estimated by SPCCT in each group paralleled the

decrease in injected cell quantity in each group (Figure 6). The limit of detection was 4902 ± 964 AuNP-labeled cells per voxel.

Discussion

In this study, we have shown that therapeutic cell tracking was feasible in brain-damaged rats with SPCCT. This technique enabled cell monitoring for up to 2 weeks post-injection, in a specific and quantitative manner, with a detection limit as low as 5,000 cells in a voxel of $250 \times 250 \times 250$ µm in vivo. Furthermore, we have provided the proof-of-concept that simultaneous imaging with discrimination of cells and their embedding scaffold was feasible with SPCCT, thus paving the way for monitoring ATMPs the long combinations in term with the next-generation multicolor clinical SPCCT scanners.

MRI coupled with cell labeling with SPIO has been extensively used to track cells in several clinical applications including ischemic stroke (11). One major limitation of SPIO-enhanced MRI, however, is that iron oxide labeling suppresses MR signal and thus hampers the evaluation of therapeutic cells' effect on tissue regeneration by other MRI sequences. Another limitation is that cell quantification is very difficult with SPIOs. This led us to seek an alternative imaging modality, with the pre-requisite to preserve MRI contrast and allow the quantitative monitoring of cells over time. One possibility is to use perfluorocarbon contrast agents to label the cells and to image them with ¹⁹F MRI (39, 40). CT represents another attractive translational option, as it is widely available and intrinsically quantitative, while some radiopaque elements such as iodine and gold do not affect MR signals. Tracking therapeutic cells with CT is an emerging field (reviewed in (12)). The main originality of spectral CT compared to other non-spectral X-ray-based approaches (µCT, clinical CT or synchrotron radiation CT (41)) is represented by its specificity with regard to the detection of gold or

other elements with K-edge energies in the appropriate range (\sim 40 – 100 keV) (42-44). Pioneering work for the specific imaging of macrophages with SPCCT was published 10 years ago in a mouse model of atherosclerosis (45). However, the technology was not yet mature and the image acquisition took several hours, which prevented *in vivo* applications, until the development of novel SPCCT prototypes such as the one used in the current study.

Using a specific imaging approach has many advantages. First, it simplifies both data acquisition and post-processing, which is a major asset for clinical translation at the diagnostic, organizational and financial levels. Second, it allows the exclusion of confounding signals from endogenous origin due, for instance, to recent hemorrhage (which is the most common complication of ischemic stroke), intracranial calcifications (which may be found in patients especially when they are aged (46)) or bone (i.e. the skull for cerebral imaging). Third, it allows differentiation of contrast agents of interest from other contrast agents. For example, iodine may be administered intravenously or intraarterially for diagnostic or interventional purposes and accumulate into the brain parenchyma as a result of blood brain barrier disruption. Alternatively, in the present study, we took advantage of this unique feature to track both cells and the cell-embedding scaffold. The capacity to discriminate two distinct contrast agents at the same location was limited to optical imaging techniques thus far. To the best of our knowledge, this is the first time that both therapeutic cells and scaffold may be tracked simultaneously in 3D and in a non-ambiguous manner using a medical imaging technique. This represents a technological leap and a great promise for evaluating ATMPs combination, for which fate elimination rate should and be evaluated independently. In this paper, our aim was to demonstrate that bicolor imaging with SPCCT can allow differentiation of therapeutic cells from their

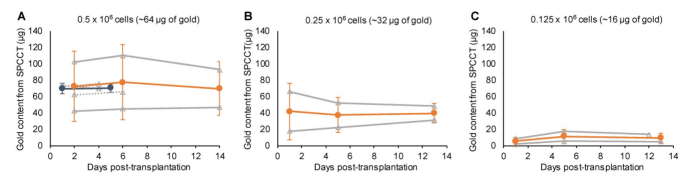


Figure 6. Bicolor study: quantitative analysis. Graph plotting the average gold content estimated on gold K-edge images over time in the group of rats that received (A) 0.5×10^6 ; (B) 0.25×10^6 ; and (C) 0.125×10^6 cells in INP-labeled scaffold. Grey lines represent individual animals (dotted lines: 1 week follow-up and plain lines: 2 weeks follow-up); blue, respectively orange, lines represent the mean ± standard deviation of animals with 1 week, respectively 2 weeks, follow-up. SPCCT: spectral photon counting computed tomography.

encapsulating scaffold, both labeled with distinct contrast agents. We have chosen to use gold for cell labeling and iodine for scaffold labeling in order to minimize the cross-talk that may arise from elements with closer K-edges, such as gold and gadolinium, based on previously published phantom and animal studies (17, 19, 20).

One key feature of using imaging in the context of restorative therapy is the possibility to monitor the success of cell transplantation. The variability in clinical outcome is still a clear obstacle for the adoption of cell therapies (47); however, it is crucial to determine if patients are non-responders due to a lack of treatment efficacy or to the failure of cell transplantation procedure and/or a suboptimal administration of cells. Another advantage of SPCCT is the fact that it is quantitative, thus allowing estimating the number of cells present in the sample. SPCCT provided an evaluation of gold amount that was in good agreement with that of ICP-OES taken as gold standard. This confirms and extends the quantitative performances previously obtained with different contrast agents using the same SPCCT prototype (17, 19, 38, 48). Our study demonstrates that there is an important variability in the number of transplanted cells from one animal to another, even in the highly controlled lab environment. One animal was excluded from the study because cells were injected in the lateral ventricle (which undergoes dilatation in the presence of focal cerebral injury), and one animal would have been excluded from the analysis because of the very low number of grafted cells. In the remaining animals, there was still a disparity at the baseline that could be accounted for in a therapeutic trial through a multivariate analysis. The use of SPCCT imaging coupled to cell labeling with AuNPs may thus allow a more rigorous inclusion of animals/patients and a more reliable statistical analysis of data in future preclinical and clinical trials of cell therapy. In case of intracerebral injection, using a technique that can specifically image AuNPs-labeled cells ascertains that other sources of hyperdensities, such as microhemorrhages caused by the injection procedure, are not mistakenly taken for therapeutic cells. These aspects taken together translate into a reduction of animals required for obtaining robust results in preclinical trials.

The other interest of using SPCCT imaging is in the possibility to depict the 3D cell distribution within the whole brain and to monitor their fate over a long period of time. This is essential to assess cell engraftment and to evaluate whether an early disappearance of cells might explain a lack of therapeutic efficacy. In our study, we observed very little changes in cell deployment after their injection in the brain, whether they were administered in buffer solution or embedded in a scaffold. The amount of gold that was estimated using K-edge gold imaging with SPCCT did not vary over time for a given individual. This might suggest that the number of transplanted cells remained relatively constant and that cells did not proliferate (as expected in case of macrophages). Of course, one limitation of the approach is that it does not allow determining whether cells are still labeled with AuNPs or, if this is the case, whether they are still viable and functional. However, this is true for any form of cell labeling with NPs (i.e. comparable approaches such as SPIO-enhanced or ¹⁹F MRI) (40, 49). Indeed, AuNPs may be released from cells and taken up by nearby endogenous macrophages. We were not able to evaluate this hypothesis in our study because we used repair macrophages as therapeutic cells. We have chosen these cells both because they represent promising candidates to treat ischemic stroke patients (24, 25) and because they are easy to label with NPs due to their phagocytic capacities (50, 51). Future studies should aim at discriminating injected cells from endogenous ones by making use of cells with bioluminescence/fluorescence or by revealing specific epitopes for instance for cells of human origin. We did not see any elimination of AuNPs during the course of the study, which demonstrates the feasibility of long-term tracking but may also bring up a safety concern. It would be interesting to conduct a longitudinal imaging study over longer periods of time to define the time course (and pathways) of gold elimination from the brain.

One limitation of X-ray based techniques for cellular imaging is their intrinsic low sensitivity. Nevertheless, it is interesting to note that the detection limit of SPCCT imaging was of the same order of magnitude as that of ¹⁹F MRI (i.e. 10³-10⁴ cells per voxel). This initial study used conventional reconstruction by filtered back-projection therefore there is scope for improvement of sensitivity, for example by implementation of iterative reconstruction (52, 53). In addition, AuNP load per cell may be further optimized (54). In our study, the detection limits have been basically determined for informative purposes and should be investigated more thoroughly as a next step. However, the main difficulty is the need for a gold standard technique that would allow precise evaluation of the gold amount in a given voxel. To address this issue, most brains of the current study have been further imaged with in vivo K-edge imaging and ex vivo phase-contrast imaging using X-rays from Synchrotron radiations (data analysis is on-going and will be published elsewhere). Another limitation of SPCCT for cell tracking may be patient exposure to

ionizing radiations. Radiotoxicity is not an issue for longitudinal follow-up of rodents with μ CT (55, 56). In the same way, SPCCT may allow safe, multiple exams of the same patient since the scans are performed at a much lower dose when compared to conventional CT for an equivalent image quality and also because of the elimination for the need of prescan (14, 48, 57-60).

The SPCCT prototype that we have used for this study displayed all the characteristics of a clinical CT scanner, but a reduced field of view (28). Due to improved image quality compared to conventional CT, we were able to image rats and mice brains with sufficient details to track therapeutic cells. SPCCT clinical scanners with full field of view are now available (61), including in our own institution. Therefore, there is hope that the SPCCT approach that we propose may help investigations of cell therapy and ATMP combinations in future clinical trials. Some hurdles, however, need to be overcome in the preclinical field beforehand. The main questions arise from the fate of contrast agents in the body (e.g. potential accumulation in the brain) and the possible interactions between the contrast agents and the therapeutic effects. It would be too long to debate these issues here, but two points might be interesting to consider with regard to the use of gold as the cell label. First, it has been shown recently that macrophages were able to scavenge AuNPs and eliminate them through exosomes (62). Second, previous studies have shown that macrophage function was not affected by gold loading (29, 63). Of course, these issues should be addressed in well-designed in vitro and in vivo experiments. We believe that our results represent a major milestone from a methodological point of view and should facilitate better characterization and optimization of cell therapies at the preclinical level. In the present study, rats were not submitted to stroke because our main purpose was to evaluate a novel imaging approach; thus a 'morphological' and 'radiological' model of brain lesion that did not induce mortality. One of the perspectives of our work is to evaluate the effects of the intracerebral administration of therapeutic cells aimed at modulating neuroinflammation (such as M2-polarized macrophages or human mesenchymal stem cells) in a rat model of ischemic stroke by coupling multimodal imaging (SPCCT for monitoring cells and scaffold and advanced multiparametric MRI methods for evaluating tissue repair) with neurobehavior and immunohistochemistry analyses. Because of its translational nature, this protocol may help bridging the gap between preclinical and clinical trials, and thus foster the

future approval of efficient cell therapies and ATMP combinations for stroke patients.

Conclusions

Multicolor CT is an innovative translational imaging tool that allows monitoring and quantifying therapeutic cells and their encapsulating scaffold transplanted into the damaged rat brain.

Abbreviations

11-MUDA: 11-mercaptoundecanoic acid; 3D: three-dimensional; ATMPs: advanced-therapy medicinal products; AuNPs: gold nanoparticles; CT: tomography; ICP-OES: computed inductively coupled plasma-optical emission spectrometry; INPs: iodinated nanoparticles; LPS: lipopolysaccharide; µCT: micro-computed tomography; mM-CSF: mouse macrophage-colony stimulating factor; MRI: magnetic resonance imaging; PBS: phosphate-buffered saline; PDI: polydispersity index; pMCAO: permanent middle cerebral artery occlusion; RPMI: Roswell Park Memorial Institute; SPCCT: spectral photon counting computed tomography; SPIO: superparamagnetic particles of iron oxide; THF: tetrahydrofuran; TIB-PVAL: 2,3,5-triodobenzoyl ester of poly(vinyl alcohol); UV: ultraviolet.

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Supplementary Material

Supplementary figures. http://www.ntno.org/v04p0129s1.pdf

Competing Interests

One of the authors (DPC) is supported by a research grant from Philips Healthcare, which is the manufacturer of the prototype SPCCT system used in this work.

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