Tracking the engraftment and regenerative capabilities of transplanted lung stem cells using fluorescent nanodiamonds

Tsai-Jung Wu1,2,3†, Yan-Kai Tseng4,5†, Wei-Wei Chang2,3, Chi-An Cheng4, Yung Ku04,5, Chin-Hsiang Chien1, Huan-Cheng Chang2,4,5* and John Yu1,2,3,6*

Lung stem/progenitor cells are potentially useful for regenerative therapy, for example in repairing damaged or lost lung tissue in patients. Several optical imaging methods and probes have been used to track how stem cells incorporate and regenerate themselves in vivo over time. However, these approaches are limited by photobleaching, toxicity and interference from background tissue autofluorescence. Here we show that fluorescent nanodiamonds, in combination with fluorescence-activated cell sorting, fluorescence lifetime imaging microscopy and immunostaining, can identify transplanted CD45CD54CD157 lung stem/progenitor cells in vivo, and track their engraftment and regenerative capacities with single-cell resolution. Fluorescent nanodiamond labelling did not eliminate the cells’ properties of self-renewal and differentiation into type I and type II pneumocytes. Time-gated fluorescence imaging of tissue sections of naphthalene-injured mice indicates that the fluorescent nanodiamond-labelled lung stem/progenitor cells preferentially reside at terminal bronchioles of the lungs for 7 days after intravenous transplantation.

To enable the aforementioned therapeutic applications of LSCs, it is crucial to determine their tissue-specific engraftment and regenerative capacity in animals. Achieving this goal requires the use of a highly sensitive imaging modality, together with a highly stable (both biologically and photophysically) imaging probe, to allow tracking of the transplanted cells in vivo over an extended period of time. Compared with magnetic resonance imaging (MRI), optical imaging offers the advantages of higher sensitivity, larger throughputs, and greater potential for clinical translation. However, existing fluorescence-based technologies such as dye labelling and in vitro transfection of cells with plasmids to express fluorescent proteins (for example, GFP) or non-fluorescent compounds (for example, luciferase) often suffer from the problems of photobleaching, interference from background tissue autofluorescence and/or low transfection efficiency for primary cells. Although quantum dots have been adopted and highly regarded as alternatives, their application is hampered by their potential toxicity and facile degradation in vitro as well as in vivo. Fluorescent nanodiamonds (FNDs), a new member of the nanocarbon family, have emerged recently as a novel nanotechnology imaging probe for biological applications. The nanoscale diamond material is chemically robust, biologically inert, and non-toxic at both cellular and whole-organism levels. A number of experiments have demonstrated that bare FND particles can be readily taken up by cells through clathrin-mediated endocytosis, but with an exceptionally low exocytosis activity. When excited by green-yellow light, the fluorophores (for example, negatively charged nitrogen–vacancy centres, NV−) built into the FND emit perfectly stable far-red emission (~600–800 nm) with a fluorescence lifetime (τ) greater than 15 ns, even in cells. This lifetime is substantially longer than that (τ ≈ 1–4 ns) of the endogenous and exogenous fluorophores commonly used in cell biology, making it possible to separate FND emission from the strong autofluorescence background of host tissue using fluorescence lifetime imaging microscopy (FLIM) and various nanosecond time-gating techniques. In this work, we show that primary cells such as LSCs can be spontaneously labelled with FNDs by endocytosis, and that the labelling does not eliminate cellular functions such as division, proliferation and differentiation. By combining FND–FLIM with fluorescence-activated cell sorting (FACS) and immunostaining analysis, we have been able to identify, unequivocally, transplanted cells in histological lung sections after intravenous injection.
the FND-labelled LSCs into mice for more than a week, with single-cell resolution. The tissue-specific engraftment capacity of these cells is further elucidated with lung injury models.

**Isolation and characterization of LSCs**

Single-cell suspensions of pulmonary cells were prepared from the proteolytic digestion of neonatal mouse lung tissues, and removal of red blood cells by lysis. Viable LSC candidates were isolated by FACS through staining with two specific cell surface markers (CD54 and CD157), propidium iodide and the CD45 antibody to exclude haematopoietic lineage cells. Flow cytometric analysis with CD31 and immunostaining with von Willebrand factor and (CD54 and CD157), propidium iodide and the CD45 antibody to FACS through staining with two specific cell surface markers (Supplementary Fig. S3, S4), the expression of these endothelial markers, indicating that they were not contaminated with endothelial cells (Supplementary Fig. S1). Only a few cells of the subpopulation expressed e6-integrin, CD24 and EpCAM (Supplementary Fig. S2). The expression pattern of these epithelial markers suggests that the presently isolated CD45^−CD54^−CD157^− cells markedly differ from the LSC cohorts reported in the literature (see Supplementary section, ‘Results and Discussion’ for details).

To fully characterize the sorted CD45^−CD54^−CD157^− cells, immunofluorescence analysis was first performed at the time of isolation. Through a careful assessment for the specificity of the detection using isotype control antibodies (Supplementary Figs S3, S4), we found that these cells expressed the undifferentiated embryonic stem cell markers (Oct-4 and Nanog), the epithelial cell marker (cytokeratin-7), and the club cell marker (CCSP), but not the differentiated type II and type I pneumocyte markers (SP-C and Aqp-5, respectively) (Fig. 1a). There were no changes in the expression of these markers after cell culture on collagen I-coated plates using DMEM medium supplemented with 10% fetal bovine serum (FBS), 1% insulin–transferrin–selenium (ITS) and 10 ng ml ^{−1} epithelial growth factor (EGF) for 10 days (Supplementary Fig. S5).

We next examined whether the cultured CD45^−CD54^−CD157^− cells had the capacity for cell differentiation. Cells were incubated in MCDB201 medium supplemented with 1% FBS, 1% ITS and 10 ng ml ^{−1} EGF for ~1–2 weeks. After 7 days of incubation, the cells became flattened and larger, and SP-C expression was detected in the cytoplasm close to the perinuclear region, which is indicative of cell differentiation into type II pneumocytes (Fig. 1b). Extension of the incubation to 14 days led to further flattening and enlargement of the cells (approximately fivefold greater than that of the original undifferentiated cells). These morphological changes were accompanied with a decrease in SP-C expression and an increase in Aqp-5 expression, an indication that the prospectively isolated CD45^−CD54^−CD157^− cells have the potential to differentiate into type II and then type I pneumocytes in a sequential manner.

**Labelling of LSCs with FNDs**

FNDs with dimensions of ~100 nm were produced by ion irradiation and thermal annealing of type I b diamond powders. To carry out FND labelling, the cultured CD45^−CD54^−CD157^− cells (identified hereafter as LSCs) were incubated with 100 nm FNDs in DMEM at 37 °C with 5% CO₂ for 4 h, after which the cells were washed thoroughly with phosphate-buffered saline (PBS) to remove free FNDs, and then collected by trypsin treatment. Flow cytometric analysis showed that the amount of 100 nm FNDs taken up by the LSCs reached a plateau at a concentration of 100 μg ml ^{−1} (Supplementary Fig. S6). This behaviour closely resembles that of many other cell types including HeLa cells, 3T3-L1 pre-adipocytes and 489-2.1 multipotent stromal cells. An extraordinary feature of the FND labelling is that in addition to far-red fluorescence, data collected in the side scatter (SSC) channel also provide useful information on FND uptake (Fig. 2a). This is attributable to the fact that diamond has the highest refractive index of all transparent minerals, and diamond nanocrystallites scatter light strongly in the visible region and thus produce strong SSC signals. By properly gating the bivariate plot (SSC versus Far-Red) in quadrants, two distinct separate populations of FND-labelled and unlabelled cells could be detected in these two channels (Fig. 2a). The double-positive cell population, denoted SSC^−Far-Red^+, was subsequently identified as FND-labelled cells without additional staining.

Primary cells like LSCs divide slowly (~2 days for one division). Moreover, they are sensitive and fragile, and properties such as growth rate and differentiation capacity are likely to be affected by the nanoparticle-based labelling. To address this issue, we tracked
the proliferation of the FND-labelled LSCs continuously over two weeks by flow cytometry (Fig. 2b). Immediately after FND labelling, the mean fluorescence intensity of the LSCs was ~45-fold greater than that of the unlabelled control cells. A successive decrease in fluorescence intensity by half occurred approximately every 48 h due to cell division. LSCs labelled with FNDs at 100 fluorescence intensity by half occurred approximately every 48 h than that of the unlabelled control cells. A successive decrease in total population of viable pulmonary cells appeared as FND-labelled cells were not functionally engrafted. It is most likely that they were only initially trapped in the lung microvasculature and were eventually lost during the first week following transplantation.

To exclude the possibility that the sorted SSC−Far-Red+ cells were false positives, lung tissue sections were stained with haematoxylin and eosin (H&E) for morphological analysis, and then fluorescently imaged. However, when searching for FND-labelled LSCs in the tissue sections using confocal fluorescence microscopy, the images acquired using a continuous-wave 561 nm laser as the light source were overwhelmed by tissue autofluorescence and by the fluorescence derived from photoexcitation of the stain. To address the gating thresholds in the bivariate plots were carefully chosen by referring to the in vitro result (Fig. 2a) as well as the profiles of the saline controls (Supplementary Fig. S8) to ensure good reliability. With a false positive rate of less than 0.05%, as determined from the controls, the observed approximately tenfold decline in the SSC−Far-Red+ subpopulation was a reflection of the fact that most of the transplanted cells were not functionally engrafted. It is most likely that they were only initially trapped in the lung microvasculature and were eventually lost during the first week following transplantation.

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this issue, lung tissue sections were stained with the macrophage-specific antibody, F4/80, followed by haematoxylin counterstaining and fluorescence imaging. Overlapping of the bright-field and time-gated fluorescence images (Fig. 3c) showed no sign of FND colocalization with the F4/80-stained macrophages, suggesting that the observed FND-labelled LSCs were not phagocytosed after i.v. injection. Such identification could not have been made using organic dyes such as carboxyfluorescein succinimidyl ester (CFSE) because of the similarity in lifetime between CFSE and the background fluorescence (Supplementary Fig. S9).

**Engraftment of FND-labelled LSCs in lung injury models**

It is known that the regenerative capacity of LSCs is determined not only by their intrinsic developmental potential, but also by their interaction with other cell elements in their niches. This capacity could be substantially activated after tissue injury. To illustrate this effect, we tracked LSCs using mice pretreated with naphthalene, which selectively ablated club cells in the epithelium of terminal and respiratory bronchioles. Club cells (or Clara cells) are secretory cells that play a protective role in the bronchial tissue against damage. In this experiment, $5 \times 10^5$ FND-labelled LSCs were injected into the mice after lung injury for 2 days. Because LSCs express CCSP (Fig. 1a), the extent of the injury and the repair of the bronchiolar epithelium could be examined by immunostaining against CCSP (club cell secretory protein). On day 1, the bronchiolar epithelium in the lung-injured mice was sparsely surrounded by CCSP$^+$ cells in both the control and treatment groups (Fig. 4a), showing low degrees of lung repair. Although

**Figure 3 | FND-labelled LSCs in uninjured mice.**

a. Flow cytometric analysis of total lung cells collected from uninjured mice receiving an i.v. injection of FND-labelled LSCs for 1, 4 and 7 days ($n = 3$ per group). b. Representative FLIM, TGF and bright-field H&E staining images of the same lung tissue sections from mice in a. The merged H&E and TGF images show that the FND-labelled cells (denoted by black arrows) are primarily located in the subepithelium of bronchiolar airways. Scale bar, 50 μm. c. Co-localization examination of FND-labelled LSCs and macrophages in a typical lung tissue section immunostained with macrophage-specific antigen F4/80 and haematoxylin counterstain. The enlarged view in the last panel shows that the transplanted FND-labelled cells (red, denoted by red arrows) and alveolar macrophages (brown, denoted by black arrows) are located at different positions. Scale bar, 10 μm. Experiments were repeated in triplicate.
they also exhibited different degrees of clustering in the subepithelium of bronchiolar airways in uninjured mice (Fig. 3b). In contrast, the localization of the cells in the lung tissue section of a naphthalene-injured mouse on day 7, showing engraftment of the transplanted FND-labelled LSCs (denoted by white and black arrows) to terminal bronchioles in cluster form. Scale bars, 50 μm. Experiments were repeated in triplicate.

Figure 4 | FND-labelled LSCs in lung-injured mice. a, b. Immunohistochemical analysis of lung tissue sections (a) and flow cytometric analysis of total lung cells (b) collected from naphthalene-injured mice receiving an i.v. injection of saline (control) or FND-labelled LSCs for 1 and 7 days (n = 3 per group). The tissue sections in a were stained with CCSP for club cells (brown). c. Representative FLIM, TGF and bright-field H&E staining images of the same lung tissue sections, showing the location of FND-labelled LSCs (denoted by white and black arrows) in terminal bronchioles of the lungs. d. FLIM and H&E/TGF images of the lung tissue section of a naphthalene-injured mouse on day 7, showing engraftment of the transplanted FND-labelled LSCs (denoted by white and black arrows) to terminal bronchioles in cluster form. Scale bars, 50 μm. Experiments were repeated in triplicate.

In the lung-injured mice, although most of the transplanted FND-labelled LSCs were monodispersed or in singlet form (Fig. 4c), they also exhibited different degrees of clustering in the terminal bronchioles (Fig. 4d). It is therefore speculated that the pattern of the distribution of these FND-labelled cells may reflect different degrees of injury in the lung tissues. For instance, the clustered cells may represent regions that have sustained more injury from the naphthalene. A close examination for the co-localization of the FND-labelled cells with the CCSP (Fig. 5a) and pan-cytokeratin (Fig. 5b) epithelial markers by high-power fluorescence imaging with single-cell resolution confirmed that the transplanted LSCs were not entrapped in the microvascular network of the lung, but rather had been functionally engrafted and integrated in the bronchiolar epithelium.

Our results have shown that the lung epithelia of the injured mice were restored more rapidly after transplantation of the FND-labelled LSCs than with saline control (Fig. 4a). To provide a semi-quantitative estimate, we calculated the percentage (P_h) of the transplanted LSCs engrafted to the lung based on the total number (N_h) of viable (PI^-) cells isolated from the mouse lung tissue, the percentage (P_v) of the FND-labelled cells identified in FACS analysis, and the number (N_v) of the cells actually transplanted, as P_h = N_h P_v / N_v. With N_v ≈ 7 × 10^6, P_v = 1.64% and N_v ≈ 5 × 10^5, the P_h for normal mice was estimated to be ~23% on day 1. This percentage, however, markedly declined to 1.7% on day 7 (Fig. 3a). In contrast, the P_h for the lung injury model exhibited a much smaller decrease, with P_h ≈ 13% on day 1 and P_h ≈ 11% on day 7 (Fig. 4b). The
distinct contrast between these two results provides compelling evidence that the lodgement of the transplanted FND-labelled cells in the injury models represents a ‘pro-active’ tissue-specific engraftment, instead of non-specific (or passive) entrapment.

A merit of the FND labelling technique is that it allows quantitative assessment of the distribution of transplanted LSCs in tissue due to the excellent chemical stability and photostability of the nanomaterial. We conducted this analysis by acquiring images of H&E staining using a bright-field section scanner. From these images, the cells showing FND labelling were counted visually in randomly selected regions. To reduce potential biases in the quantitative assessment, measurements were performed for more than 30 visual fields per section for six randomly selected sections of each sample, by independent workers. The results show that the percentages of FND-labelled LSCs nested in the terminal bronchioles and alveoli of the lung-injured mice are 59 ± 5% and 41 ± 5%, respectively, compared with 24 ± 3% and 76 ± 3% for the uninjured mice (Fig. 5c). The marked difference in the histomorphometric distribution of the FND-labelled LSC population between these two animal models serves as additional evidence that the transplanted LSCs reach lung tissues and further engraft to the terminal bronchioles (black) or alveoli (white) of the lungs of uninjured and injured mice on day 7 after i.v. injection. Results are averages of more than 30 visual fields per section, with error bars representing standard deviations of the measurements (n = 6 per group).

Methods

Cell isolation and immunofluorescence staining. Viable LSCs were isolated from neonatal CD-1 (ICR) mice by proteolytic digestion of lung tissues and lysis of red blood cells, followed by sorting with FACS after staining with propidium iodide, CD34, CD137 and CD45 antibodies. Cells were then cultured through several passages and seeded at a density of 2 × 10⁵ cells ml⁻¹ in a collagen I-coated plate using DMEM supplemented with 10% FBS, 1% ITS and 10 ng ml⁻¹ EGF. To perform immunofluorescence staining, cells in primary culture were first fixed in 4% paraformaldehyde/ PBS at room temperature, permeabilized, and blocked with 0.5% Triton X-100 in 3% bovine serum albumin (BSA)/ PBS. Cells were then incubated at 4 °C with the primary antibodies against antigens including SP-C, Aqp-5, cytokeratin-7 (all from Millipore), CCSP and Oct-4 (both from Santa Cruz). After overnight incubation, the cells were washed and incubated with respective FITC- or Cy3-labelled secondary antibodies (Jackson Immunoresearch), and then counterstained with 4,6-diamidino-2-phenylindole (DAPI). Photomicrographs of the cells were immediately taken after staining with a Zeiss fluorescence microscope.

Figure 5 | Identification of transplanted FND-labelled LSCs in bronchiolar epithelia and their histomorphometric distribution. a, b Club cell markers (CCSP, a) and epithelial markers (pan-cytokeratin, b) examined by FLIM and immunostaining of lung tissue sections prepared from naphthalene-injured mice on day 7 after i.v. injection of FND-labelled LSCs. The enlarged, merged immunostaining and TGF images in the red dashed boxes show that the FND-labelled LSCs (red) co-localize with CCSP⁺ (brown) and pan-cytokeratin⁺ (brown) cells and reside at the bronchiolar epithelial region of the lung. Scale bars, 10 μm. c, Histomorphometric distribution measured by visually counting the numbers of FND-labelled LSCs nested in terminal bronchioles (black) or alveoli (white) of the lungs of uninjured and injured mice on day 7 after i.v. injection. Results are averages of more than 30 visual fields per section, with error bars representing standard deviations of the measurements (n = 6 per group).

Conclusions

We have developed a novel nanodiamond-based platform that enables long-term imaging and tracking of transplanted stem/progenitor cells in vivo. The platform consists of FND labelling, FLIM and FACS technologies, with many unique features including simple labelling, high throughput and unequivocal detection of fluorescence signals that are not confounded by tissue autofluorescence and/or fluorescence derived from exogenous stains. Moreover, the FND labelling does not eliminate the cells’ in vitro differentiation and in vivo migration. Using the combined techniques, we have been able to follow the fate of FND-labelled LSCs in vivo after transplantation, precisely locate their position in tissue with single-cell resolution, and quantify their engraftment capacity over a week for the first time. The platform is particularly suitable for the study of rare stem cells that are difficult to transfect to express fluorescent proteins or non-fluorescent compounds. The method is general and also has broad applicability to other stem cell systems (such as haematopoietic stem/progenitor cells⁴³,⁴⁴).

FND production and labelling. FNDs were produced by radiation damage of type Ib diamond powders (Micron+ MDA M0.10, Element Six) using a 40 keV He⁺ ion beam, followed by thermal annealing at 800 °C, air oxidation at 450 °C and purification in concentrated H₂SO₄–HNO₃ (3:1, vol/vol) solution at 100 °C.
Confocal fluorescence microscopy and FLIM. Confocal imaging was carried out using a 6-P5 inverted microscope (Leica) equipped with three solid-state lasers operating separately at 561, 488 and 405 nm for the excitation of FND, SP-C and DAPI, respectively. Fluorescence was collected through an oil-immersion objective (x100, NA 1.4) and detected with either a photomultiplier tube (PMT) for SP-C and DAPI or an avalanche photodiode (APD) for FND. To conduct FLIM, an external frequency-doubled picosecond Nd:YAG laser (IC-532-30, High Q Laser), operating at 332 nm with a pulse width of 7 ps and a repetition rate of 50 MHz, was coupled through a single-mode optical fibre into the Leica microscope featuring a FLIM system (Supplementary Fig. S1). FND fluorescence was collected using a x 63 oil-immersion objective and detected with an APD at a wavelength of 650–850 nm through a bandpass filter (OS196646-HQ750/210m, Chroma). A time-correlated single-photon counting system (Pico-Harp 3200, Pico Quant) was used to measure the fluorescence lifetimes, from which time-gated images were obtained with the SynPhoTime software.

In vivo tracking. Mouse LSCs were labelled with bare FNDs for 4 h at 100 μg ml⁻¹. After washing away excess FNDs, 5 × 10⁶ cells in 100 μl PBS were i.v. injected into adult mice (four weeks old) through tail veins. Mice were maintained until being killed on days 1, 4 and 7 (n = 3 per group), after which organs were collected for analysis with FACS and FLIM. All assays were performed for at least three independent experiments. For fluorescence imaging, tissues were first fixed in 10% formalin overnight and then embedded in paraffin and sectioned. Sections (5 μm) were deparaffinized in xylene, rehydrated in a graded ethanol series, and incubated in blocking buffer. Sections were incubated overnight with antibodies against antigens including F4/80 (eBioscience), CCSP and pan-cytokeratin (both from Santa Cruz) and then analysed immunohistochemically with a peroxidase detection kit (Vector Laboratories) using diaminobenzene as the substrate according to the manufacturer’s instructions. All sections were counterstained with Mayer’s haematoxylin. Images of lung tissues after H&E staining or immunostaining were acquired using a high-field scanner (Aperio Digital Pathology System) equipped with a x 40 objective. In separate experiments with lung injury models (n = 3 per group), naphthalene dissolved in corn oil was injected intraperitoneally at 275 mg kg⁻¹ body weight. For fluorescence imaging, tissues were first fixed in 10% formalin, washed by PBS to remove free FNDs and then analysed by using a FACSAria flow cytometer (BD Biosciences) equipped with a 594 nm laser and a 670 nm emission filter in the far-red channel. To conduct differentiation studies, injured FND-labelled LSCs were incubated in MCDB201 medium supplemented with 1% FBS, 1% ITS and 10 ng ml⁻¹ EGF. Cells were then fixed, permeabilized, and stained with primary antibodies against SP-C.

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Author contributions
T.J.W. and Y.K.T. conceived the study, conducted the experiments, interpreted the results, and wrote the manuscript. T.J.W. performed the tissue-culture work, cell sorting and tracking experiments, and data analysis. Y.K.T. and Y.K. assembled the FLIM systems and produced the image acquisition scripts. W.W.C. helped with the isolation of LSCs and carried out data analysis. C.A.C. prepared FNDs and performed the cell tracking experiments. C.H.C., H.C.C. and J.Y. supervised and planned the project, analysed the data, and wrote the manuscript. All authors designed the experiments and edited the manuscript.

Additional information
Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to H.C.C. and J.Y.

Competing financial interests
The authors declare no competing financial interests.