Prolonged intracellular accumulation of light-inducible nanoparticles in leukemia cells allows their remote activation

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Leukaemia cells that are resistant to conventional therapies are thought to reside in protective niches. Here, we describe light-inducible polymeric retinoic acid (RA)-containing nanoparticles (NPs) with the capacity to accumulate in the cytoplasm of leukaemia cells for several days and release their RA payloads within a few minutes upon exposure to blue/UV light. Compared to NPs that are not activated by light exposure, these NPs more efficiently reduce the clonogenicity of bone marrow cancer cells from patients with acute myeloid leukaemia (AML) and induce the differentiation of RA-low sensitive leukaemia cells. Importantly, we show that leukaemia cells transfected with light-inducible NPs containing RA can engraft into bone marrow in vivo in the proximity of other leukaemic cells, differentiate upon exposure to blue light and release paracrine factors that modulate nearby cells. The NPs described here offer a promising strategy for controlling distant cell populations and remotely modulating leukaemic niches.
The differentiation of leukaemia cells is a therapeutic strategy often used in the clinic to eradicate blood cancers. The concentration of the agent used for inducing leukaemia cell differentiation and the spatio-temporal control of its application are important variables for the success of this therapeutic approach. Induction of leukaemia cell differentiation by RA is a therapeutic strategy that has been used with great success in the treatment of acute promyelocytic leukaemia (APL) and RA activates nuclear RA receptors (RARs) that induce cell growth arrest and differentiation. Despite its clear therapeutic efficacy, approximately 25% of patients receiving RA will develop serious complications, such as differentiation syndrome. Hence, there is a need for more effective formulations to deliver RA into leukaemia cells while preventing RA side effects. In addition, leukaemia cells resistant to conventional therapies reside in microenvironmental niches in the bone marrow that are difficult to access by therapeutic interventions. New strategies are required to address these problems.

Nanoparticles (NPs) that disassemble in response to light offer a promising approach for reducing the side effects of conventional therapies and increasing access of therapeutic agents to the target cells. Recently, light-inducible NPs have been reported to target solid tumours due to their specific accumulation in tumour vasculature after intravenous injection. However, such an approach is not applicable to leukaemia. The hypotheses of the present work are: (i) light-inducible NPs containing RA may be a more effective strategy for differentiating leukaemia cells because they release high and more effective concentrations of RA in a short period of time after NP disassembly, and (ii) light-inducible NPs containing RA accumulated in the cytoplasm of leukaemia cells may offer a unique opportunity to remotely differentiate these cells in leukaemic niches in the bone marrow, which in turn may interfere with the differentiation profile of leukaemia cells in a paracrine manner.

Here, we describe light-inducible polymeric NPs containing RA that effectively disassemble within cells after light activation. These NPs accumulate in the cytoplasm of leukaemia cells for more than 6 days. They are internalized primarily through a clathrin-mediated mechanism and at minor extent by macro-pinocytosis. They escape in few hours the endolysosomal compartment and accumulate in cell cytoplasm. We show that these NPs are more efficient and quicker at inducing transcription from the RARE-luciferase locus than RA in solution. We further show that these NPs can be activated to release RA in vivo in a highly controlled manner. Finally, we demonstrate that leukaemia cells transfected with these cells can home in the bone marrow in the same niche as other leukaemia cells, differentiate after blue laser activation and modulate the activity/phenotype of the resident leukaemia cells.

Results

Photo-disassembly and release properties of light-inducible NPs. To prepare light-inducible polymeric NPs, poly(ethyleneimine) (PEI) was initially derivatized with 4,5-dimethoxy-2-nitrobenzyl chloroformate (DMNC), a light-sensitive photochrome (Fig. 1a and Supplementary Fig. 1). PEI was selected as the initial NP block because it facilitates the cellular internalization of NPs and their subsequent escape from endosomes, while DMNC was selected because it responds rapidly to light and its degradation products are relatively non-cytotoxic. PEI–DMNC was then added to dextran sulfate (DS) to form NPs by electrostatic (PEI:DS) and hydrophobic (DMNC:DMNC) interactions. To stabilize the NP formulation, zinc sulfate was added, and NPs with an average diameter of 108.1 ± 9.9 nm and a zeta potential of 27.4 ± 1.6 mV were obtained.

To demonstrate that the NP formulation could be photo-disassembled, a suspension of NPs was exposed to UV light for up to 10 min. The number of NPs (as assessed by Kcps) decreased to below half of the initial number after 1 min of UV exposure, confirming NP disassembly (Fig. 1b). The disassembly was likely due to a change in the NP’s hydrophilic/hydrophobic balance after DMNC photo-cleavage. Importantly, a conventional blue laser (405 nm, 80 mW) (Supplementary Fig. 2B) or a blue confocal laser (405 nm, 30 mW) (Supplementary Fig. 2C) can replace the UV light to induce the photo-disassembly of NPs. The response of the NPs to a blue laser was mediated by DMNC coupled to PEI, as NPs without DMNC did not respond to the laser (Supplementary Fig. 2B). We further evaluated the feasibility of triggering intracellular NP disassembly. For that purpose, HUVECs were transfected with quantum dots 525-labelled NPs for 4 h (Supplementary Fig. 2D), and a small region of the cell housing NPs was excited by a blue laser (405 nm) under a confocal microscope. Under these conditions, NP fluorescence decreases compared to a reference region not excited with blue laser. This increase is due to NP disassembly and a resultant decrease in the quenching of the quantum dots immobilized in the NPs. Overall, our results show that we can disassemble NPs by light, either in vitro or within cells.

To evaluate the light-inducible polymeric NPs as a method for the controlled release of RA, a solution of RA with PEI–DMNC was assembled with DS in aqueous solution to form NPs (RA + NPs). Under these conditions, the carboxyl groups of RA tend to form electrostatic interactions with the amine groups of PEI. The resultant NP formulation contained approximately 150 μg of RA per mg of NP, an average diameter of 160 nm and a zeta potential of 22 mV. The release profile of the NP formulation containing [3H]RA was monitored by scintillation after NP exposure to UV light or a blue laser for up to 10 min (Fig. 1c). Up to 50 ng of RA was released per μg of NP after 10 min of activation. In the absence of light, up to 10 ng of RA was released per μg of NP after 8 days (Supplementary Fig. 3).

Internalization and intracellular trafficking of RA + NPs. Next, we asked whether leukaemia cells could uptake light-inducible NPs. According to our dynamic laser scattering studies, RA + NPs are positively charged when resuspended in different culture media and relatively stable for at least 24 h (Supplementary Fig. 4). RA + NPs did not exhibit significant cell uptake. As an equimolar solution of RA, these NPs had no substantial effect on leukaemia cell metabolism, as evaluated by ATP assay for concentrations up to 20 μg ml−1 (Supplementary Fig. 5a; in case of non-leukaemic cells no cytotoxicity was observed for RA + NPs up to 10 μg ml−1—Supplementary Fig. 6). Human bone marrow APL NB4 cells and human myelomonoblastic U937 cells (expressing a chimeric PLZF/RAR fusion protein) and showing impaired sensitivity to RA17,18 were exposed to NPs for 4 h, washed to remove excess NPs not taken up by the cells, either exposed or not to UV light for 10 min, and finally cultured for an additional 20 h. Because the activation of RA + NPs by UV or blue light can induce double-stranded DNA breaks, we evaluated the effect of this radiation on the phosphorylation of histone H2A (γH2AX), an early biomarker of DNA damage. Our results indicate that neither UV nor blue light exposure for 10 min had a significant impact on DNA damage (Supplementary Fig. 5B and C).

The intracellular accumulation of RA + NPs was monitored by flow cytometry and inductively coupled plasma mass spectrometry (ICP-MS) (Supplementary Fig. 7). Our results show that NP internalization peaked at 6 h (Supplementary Fig. 7A and B) and the intracellular concentration was dependent on the initial
**Figure 1 | NP photo-disassembly and cellular interaction.** (a) Schematic representation for the photo-disassembly of RA⁺ NPs. (b) Size, zeta potential and number of NPs (Kcps) of an aqueous suspension of light-activatable NPs (50 µg ml⁻¹) exposed to UV light (365 nm, 100 W) for up to 10 min. (c) Release of [3H]-RA from light-activatable NPs (10 µg ml⁻¹ in water) after exposure to UV or a blue laser (405 nm, 80 mW). (d) Dilution of TRITC-labelled RA⁺ NPs during THP-1 cell culture as monitored by flow cytometry. Percentages of positive cells were calculated using non-transfected cells as control. (e) THP-1 and NB4 cells were transfected with RA⁺ NPs in serum-free medium for 4 h. Left: the concentration of NPs in THP-1 and NB4 cells was monitored over 6 days by ICP-MS using Zn levels in NP. The amount of Zn in the cells after transfection (4 h) was defined as 100%. Right: cell doublings over 6 days as assessed by cell counting. (f) Internalization mechanism of RA⁺ NPs. Left: uptake of TRITC-labelled RA⁺ NPs in U937 cells after silencing key regulators of endocytosis with siRNAs. Right: schematic representation of NP internalization pathways. (g) Intracellular trafficking of FITC-labelled RA⁺ NPs. HUVEC cells were stained with LysoTracker DND-99 to monitor NPs trafficking through the endolysosomal compartment overtime. (h) Fluorescence of NPs in the cytoplasm measured by confocal microscopy. Scale bar represents 10 µm. In b,c,d,e,f results are expressed as mean ± s.e.m., n = 3. In g,h, results are expressed as mean ± s.e.m., n = 5. In f, statistical analyses were performed between control and the remain experimental groups using a one-way Anova followed by a Newman–Keuls post-test. *P < 0.05, ***P < 0.001, ****P < 0.0001.
NP loading (Supplementary Fig. 7C). No significant differences were observed in the internalization profiles between cells. Importantly, 60–100% of the leukaemia cells retained NPs in culture for at least 6 days, as confirmed by flow cytometry (Fig. 1d and Supplementary Fig. 8). This corresponded to the retention of 25–50% of the initial mass of the NPs during 8–10 cell doublings, as obtained by ICP-MS analyses (Fig. 1e). Altogether, NPs were efficiently internalized by human leukaemia cells and accumulated intracellularly for several days.

To identify the pathways of RA⁺ NP internalization, U937 cells were incubated in the presence of endocytosis chemical inhibitors at non-cytotoxic concentrations. Fluorescent-labelled RA⁺ NPs were then added and the internalization monitored by flow cytometry. Whenever possible, molecules that enter by a specific internalization pathway were used as positive controls to show the efficacy of our inhibitors (Supplementary Fig. 9A). Dynasore treatment (clathrin-mediated endocytosis (CME) inhibitor) reduced the uptake of RA⁺ NPs by 90% compared to control cells (Supplementary Fig. 9B). To further confirm the endocytosis mechanism involved in RA⁺ NP internalization, U937 cells were transfected with siRNAs to downregulate key components of different endocytosis pathways (Fig. 1f). We observed a ~60% and ~70% reduction in NP uptake upon downregulation of clathrin heavy chain (CLTC), and low-density lipoprotein receptor (LDLR), respectively, confirming a role for CME. The knockdown of macropinocytosis regulators Rac1 and CTBP1 led to a ~40% and ~50% decrease in RA⁺ NPs uptake, respectively, suggesting that macropinocytosis was also involved in RA⁺ NP internalization (Fig. 1f).

To elucidate the intracellular trafficking of our NPs, we used adherent cells (HUVECs) to allow for this characterization by confocal microscopy. The intracellular trafficking of our NPs was assessed first by performing a LysoTracker staining to see the general distribution of FITC-labelled RA⁺ NPs in the endolysosomal system. During the first few hours, there was an increase in the intracellular signal of FITC-labelled RA⁺ NPs and a clear drop in the intensity of LysoTracker, suggesting an increase in the pH of the endolysosomal vesicles by the presence of intracellular PEI (a strong base), as well as possible vesicle disruption, as the FITC-labelled NP signal was increased in the cytoplasm (Fig. 1g,h). At later time points (12 h) of incubation with FITC-labelled NPs, the intensity of LysoTracker reached control levels, suggesting that the endolysosomal system regained its normal characteristics. Our results further show that most of the NPs (~80% of NP fluorescence) escape endolysosomes within a few hours (Fig. 1h). At 6 h, most of the FITC-labelled NPs were distributed throughout the cytoplasm (~80% of the fluorescence) (Fig. 1h), and those that accumulated in the endolysosomal compartment were located in vesicles positive for Rab5 (early/late endosomes) and Rab7 (endosome/lysosomes) (Supplementary Fig. 10A and B). This is consistent with a rapid escape of the NPs from endosomes after uptake by the cell, likely due to their buffering capacity which gives rise to osmotic swelling and endosome rupture12.

Next, we asked whether RA⁺ NPs would be effluxed by leukaemia cells overtime. It is known that tumour cells express high levels of P-glycoprotein (P-gp), a membrane transporter that is responsible for the efflux of drugs20 and NPs21. Therefore, we used flow cytometry to study the effects of the P-gp antagonist, verapamil22, on the intracellular accumulation of RA⁺ NPs in Zn-induced U937-B412 cells expressing P-gp ( Supplementary Fig. 11A). Our results show that the intracellular accumulation of RA⁺ NPs in U937 slightly decreases from 4 to 12 h (Supplementary Fig. 11B), in contrast to the control, ultra-small paramagnetic iron oxide. The intracellular accumulation of ultra-small paramagnetic iron oxide was highly dependent on the inhibition of P-gp (Supplementary Fig. 11C). Overall, our results indicate that RA⁺ NP endocytosis takes place primarily through a clathrin-mediated mechanism and at minor extent by macropinocytosis. It is likely that both endocytic pathways are interconnected, as has been demonstrated recently for lipid NPs21,22. Our results further show that the internalization of RA⁺ NPs occurs quickly, and within the first 2 h, a significant percentage of NPs tend to escape the endolysosomal compartment, while the ones that do not escape accumulate in early/late endosomes.

In vitro differentiation of leukaemia cells with RA⁺ NPs. Previously, liposomes22 and block copolymer NPs23 have been used for the release of RA in myeloid leukaemia cell lines. However, no significant differences were observed between liposome/NP formulations and soluble RA in their capacity to induce leukaemia cell differentiation. Therefore, we evaluated the capacity of light-inducible RA⁺ NPs to prompt the differentiation of human CD34⁺ primary leukaemia cells isolated from the bone marrow aspirates of acute myeloid leukaemia (AML) patients, taken at diagnosis. We evaluated their clonogenic potentials in short-term (blast colony-forming units, CFUs) and long-term (long-term culture-initiating cell, LTC-IC) assays following treatment with RA or RA⁺ NPs (Fig. 2a). Our results indicate that light-inducible RA⁺ NPs were more effective than soluble RA at decreasing the number of CFUs (Fig. 2b). Our results further indicate that RA⁺ NPs activated by light are more effective in decreasing the number of CFUs (Fig. 2c) and LTC-ICs (Fig. 2d) than non-activated RA⁺ NPs. We extended the previous studies to different leukaemia cell lines, including U937, NB4 and THP-1 (Fig. 2e,g and Supplementary Fig. 12B). The differentiation process was monitored by the expression of myeloid maturation marker CD11b, a protein involved in the regulation of leukocyte adhesion and migration. For equivalent concentrations of RA, U937 (Fig. 2e) or NB4 (Fig. 2g) cells treated with RA⁺ NPs more readily differentiated into myeloid cells than cells treated with soluble RA, likely due to the higher efficiency of the NPs to deliver RA intracellularly (Fig. 2f) and Supplementary Fig. 12A). The differences in the intracellular accumulation of RA are likely explained by the differences in RA uptake mechanism. The RA is a hydrophobic small molecule with limited water solubility (63 mg l⁻¹ ref. (24)) and thus in vivo, it is transported by proteins called cellular retinoic acid-binding proteins (CRABP) both in extracellular and intracellular compartments25. The expression of CRABP is relatively low in undifferentiated leukaemic cells26 and thus the transport of soluble RA to inside of the cell might be limited. In contrast, the transport of RA⁺ NPs occurs by endocytosis which might facilitate the intracellular accumulation of RA.

U937 (Fig. 2f) or NB4 (Fig. 2h) cells treated with light-activated RA⁺ NPs showed a higher capacity for myelocytic differentiation than cells treated with non-activated RA⁺ NPs. In the case of NB4 cells, RA released from light-inducible RA⁺ NPs promoted a higher percentage of cells expressing CD131higCD11b⁺ and CD131lowCD11b⁺ (Fig. 3a,b) than soluble RA. NB4 cells differentiated by light-inducible RA⁺ NPs have a different morphology from undifferentiated cells, showing polylobular nuclei, decreased nuclear:cytoplasmic ratios and decreased cytoplasm staining, as characterized by a May–Grunwald Giemsa staining (Fig. 3c). The efficiency of the RA⁺ NPs in delivering RA inside NB4 cells and inducing a RA-dependent signalling pathway was also confirmed by the use of an NB4–RARE reporter cell line. The RA-dependent induction of a RARE element driving the transcription of the firefly luciferase gene was used to evaluate the kinetics of RA-induction using RA⁺ NPs or RA in solution.
Figure 2 | Effect of RA + NPs on human leukaemia cells. (a) Schematic representation of the methodology used. (b) Differentiation of CD34+ AML cells isolated from the bone marrow aspirates of patients with AML cultured with light-activated RA + NPs or soluble RA. (c,d) Differentiation of CD34+ AML cells with RA + NPs (10 µg ml⁻¹) or blank (RA-NPs) NPs (10 µg ml⁻¹), exposed or not to UV light, by a CFU (c) or LTC-IC (d) assays. Results are expressed as a mean percentage of control plates containing only AML cells. (e,f) Myelocytic differentiation of human Zn-induced U937-B412 cells cultured with light-activated NPs or soluble RA at day 3 (e) or cultured with RA + NPs with or without light activation (f). (g.h) Myelocytic differentiation of human NB4 cells cultured with light-activated NPs or soluble RA at day 3 (g) or cultured with RA + NPs with or without light activation for 1 or 3 days (h). (i) Intracellular release of RA as evaluated by a RARE luciferase cell line. NB4-RARE cells were cultured with soluble RA (3 µg of RA per ml) for the entire duration of the experiment, or light-activatable RA + NPs (15 µg ml⁻¹, 0.6 µg of RA per ml). Cells were exposed to NPs for 1 h, washed with PBS and resuspended in cell media. Some samples were exposed to UV light for 5 min. The cells were then cultured for 24 h before luciferase luminescence reading. (j) [3H]-RA uptake by NB4 cells. NB4 cells were cultured with soluble [3H]-RA (0.3 and 3 µg ml⁻¹) for the entire duration of the experiment, or light-activatable [3H]-RA + NPs (1 and 10 µg ml⁻¹) for 4 h and then the cells were washed and cultured in cell medium for additional 20/68 h before scintillation counting. From b to j, results are expressed as mean ± s.e.m. (n = 3). In c,d,f,h, statistical analyses were performed by an unpaired t-test while in i were performed by one-way ANOVA followed by a Newman-Keuls post-test. *P<0.05, **P<0.01, ***P<0.001 and ****P<0.0001.
Our results showed that RA⁺ NPs were able to induce high levels of luciferase activity shortly after light activation (Fig. 2i). RA⁺ NPs were more efficient and quicker at inducing transcription from the RARE-luciferase locus than RA in solution.

Next, we monitored the effects of soluble RA and RA released from RA⁺ NPs in the transcriptional regulation of 96 genes involved in the differentiation program of NB4 and U937 cells. In the case of NB4 cells, CD52, CDKN1A (also known as the p21 gene) and TNFα are recognized as direct targets of RA. CD52 and CDKN1A are upregulated and TNFα is downregulated in response to RA. Our results showed that RA⁺ NPs have induced transcription from the RARE-luciferase locus more efficiently and more quickly than RA in solution. Other RA-responsive genes were also evaluated by qRT–PCR.

The number of mRNA transcripts for IL-8 and Ly6E genes was higher in cells treated with RA⁺ NPs than in those treated with soluble RA for 8–24 h and 24–48 h, respectively. In addition, the number of mRNA transcripts for CCR2 and CEBPβ genes was lower in cells treated with RA⁺ NPs than in those treated with soluble RA for 8–24 h and 8 h, respectively. Importantly, no significant variation in mRNA transcripts for RARα, RARβ and RARγ (Supplementary Figs 13 and 15) was found between all experimental groups. Therefore, our results suggest that the differentiation profile induced by RA⁺ NPs is similar to that observed for soluble RA; however, the magnitude of the effect in the initial stages (up to 24 h) is higher in cells treated with RA⁺ NPs than in those treated with soluble RA.

Recent studies report that lipid-based NPs are able to release siRNA cargo into the cytosol only during a limited window of time, when the NPs reside in a specific compartment that shares early and late endosomal characteristics. Therefore, we evaluated the impact of temporal activation of RA⁺ NPs on the
differentiation of NB4 and U937 cells. The effect of the intracellular release of RA was evaluated in terms of cell differentiation into the myeloid lineage (expression of CD11b at day 3) (Supplementary Fig. 16A). Cells that were incubated with RA⁺NPs for 4–8 h and subsequently activated by light showed the highest differentiation into the myeloid lineage. Interestingly, cells incubated with RA⁺NPs for 1 h showed an already high level of differentiation, indicating that NP uptake is rapid and their endosomal escape is efficient. Then, we asked whether cells exposed for the same time (4h) to NPs but activated at different times (up to 44 h) during the intracellular trafficking of RA⁺NPs would have differences in their differentiation pattern (Supplementary Fig. 16B). Our results show that even after 44 h following NP internalization, the NPs can be activated and the resultant delivery of RA in the cytosol can be relatively efficient. Similar results were obtained when the effect of temporal intracellular RA release was evaluated by the induction of the NB4-RARE reporter cell line (Supplementary Fig. 16B). Overall, these results show that the NPs can be accumulated in the intracellular environment and activated after a few days, without losing their capacity to promote cell differentiation. Importantly, a single activation procedure seems to be sufficient to activate RA release from the NPs because no significant increase in efficiency was observed when multiple rounds of light activation were performed (Supplementary Fig. 17).

In vivo differentiation of leukaemia cells with RA⁺NPs. Next, we evaluated if RA⁺NPs can function in vivo. We investigated whether leukaemic cells loaded with RA⁺NPs could be activated in vivo after subcutaneous transplantation. We selected a Matrigel plug subcutaneous model because we know what is the attenuation of the blue laser through the murine skin (Supplementary Fig. 18A) and we could easily remove the cells from the Matrigel plug and evaluate their differentiation by flow cytometry. Initially we evaluated whether the in vivo environment could interfere with the differentiation program of leukaemic cells after ex vivo activation. Human NB4 cells were cultured with RA⁺NPs for 4 h, washed and activated ex vivo by exposure to a 405 nm blue laser (80 mW) for 5 min, embedded in Matrigel and then injected into a cylindrical poly(dimethylsiloxane) (PDMS) construct that was previously implanted subcutaneously in NOD/SCID recipients (Fig. 4a). The PDMS cylinder was used to restrict cell position within the implant site. After 5 days, human cells were isolated from the implants and CD11b expression measured by flow cytometry (Fig. 4b). Consistent with our in vitro data, CD11b expression was statistically higher in NB4 cells treated ex vivo with light-activated RA⁺NPs than in cells treated with RA⁺NPs without light activation (Fig. 4b). The experiment was repeated with in vivo activation. One day after implantation, the recipients were exposed to a 405 nm blue optical fibre for 5 min at the implant site (Fig. 4c). After 3 days, the recipients were killed, the human cells were isolated and CD11b expression was assessed. CD11b expression was higher in NB4 cells from mice that had been exposed to the blue laser, demonstrating that internalized RA⁺NPs can be activated to release RA in vivo in a highly controlled manner (Fig. 4c).

Homing and differentiation of leukaemia cells with RA⁺NPs. Leukaemia cells that resist therapy reside in microenvironmental niches in the bone marrow that are difficult to reach by conventional therapy. Therefore, we investigated whether leukaemic cells transfected with RA⁺NPs could engraft into the bone marrow and localize at nearby resident leukaemia cells. For this purpose, we transplanted leukaemia cells into immunodeficient CB17-Prkdcscid/J (NOD/SCID) mice treated with anti-CD122 to deplete innate immune cells (NS122). Green fluorescent protein (GFP)-labelled THP-1 cells, a human APL cell line (NB4 cells were unable to engraft), were injected into sub-lethally irradiated mice in the tail vein, followed by the injection of THP-1 cells transfected with RA⁺NPs after 24 h (Fig. 5a). After 1–2 days, animals were killed and the cells in the animal’s calvaria (cranium) were characterized by ex vivo imaging (Fig. 5b). Both cells (GFP-labelled and NP-labelled THP-1 cells) could engraft in the bone marrow of the animals and were detected after 1 or 2 days. Therefore, NP-labelled THP-1 cells (delivery cells) could enter into the same niche as the GFP-labelled cells (target cells) (Fig. 5c).

Next, we investigated whether we could differentiate the NP-labelled human THP-1 cells (delivery cells) at the bone marrow niche (more specifically at the calvaria), and whether they could modulate nearby cells (Fig. 6a). Blue laser attenuation studies in the calvaria bone using a photometer showed an attenuation of ≈75% (Supplementary Fig. 18B). Human THP-1 cells transfected with RA⁺NPs were injected intravenously into CB17-Prkdcscid/J (NOD/SCID) mice and 6 days after injection, the cranium was exposed to a blue laser for 5 min. Seventy-two hours after blue laser irradiation, the engraftment of human CD45⁺ cells in the long bones of non-irradiated and blue laser-irradiated mice was 47.7 ± 13.0 and 39.8 ± 3.5%, respectively (Fig. 6b). No statistical difference was found for the engraftment in both groups. Ex vivo staining of mice calvaria showed that human CD45⁺ cells in the non-irradiated mice did not express CD11b, while ~80% of the human CD45⁺ cells in blue-laser-irradiated mice expressed CD11b (Fig. 6c–e). In addition, cells surrounding CD45⁺CD11b⁺ cells in irradiated mice expressed CRABP, a RA protein transporter, while negligible levels of CRABP were found in non-irradiated mice (Fig. 6f). Therefore, THP-1 cells differentiated by a blue laser had the capacity to modulate the cells in the vicinity. This is likely due to RA secreted (including exosomes containing RA) by the THP-1 cells, as observed in vitro using the NB4 cell line (Supplementary Fig. 19). Overall, our results show that leukaemia cells transfected with RA⁺NPs could (i) engraft into the bone marrow and localize near to resident leukaemia cells, (ii) differentiate after blue laser activation and (iii) modulate the activity/phenotype of the resident leukaemia cells. It is possible that the cells that reach the bone marrow are the ones that are less affected by the passive diffusion of RA (that is, RA release without light activation) within the cells. Therefore, the cells that engraft are likely the ones that have less leaky RA⁺NPs and thus are undifferentiated leukaemic cells (CD11bnegative phenotype).

Discussion. In conclusion, we have developed a light-inducible polymeric NP for the spatio-temporal release of RA within leukaemia cells. We showed the efficiency of RA release both in vitro and in vivo. The efficiency is due to a combination of several factors, including (i) high concentration of intracellular RA, (ii) high endolysosomal escape (~80% of the NPs escape the endolysosomal compartment in the first 5 h), (iii) prolonged intracellular accumulation of the NPs (~6 days in leukaemia cells) and (iv) rapid disassembly of the NPs once activated by UV or blue light (within minutes). We further showed that cells transfected with light-inducible NPs can be activated after six days while maintaining the same inductive properties. This gives an opportunity to use cells for activation at specific sites in the human body and differentiate cells in their proximity. Although the present study uses a RA delivery system as a proof of concept, a similar approach could be implemented for other drugs relevant to therapeutic or regenerative medicine.
The present research reports a blue laser/UV-inducible NP. The platform was designed for initial proof of concept regarding the modulation of leukaemic cell niches in the bone marrow. Recent studies have explored UV/blue laser approaches to trigger in vivo the presentation of bioligands with spatial-temporal control to regulate cell adhesion, inflammation and vascularization of biomaterials and to detect protease activity at sites of disease. However, the in vivo applications of this light trigger are limited, as the optical properties of biological tissue cause attenuation and scattering of light rays, reducing their penetration depth. The recent advances in developing NPs activated by near-infrared light may provide a better option for enhancing light penetration in tissues. For example, upconversion nanocrystals that convert near infrared light into visible light (blue light) may be an interesting possibility for the photo-cleavage of linkers to which RA is chemically bound.

We have used leukaemic cells as a programmable delivery system that recognizes the leukaemic niches and modulate them after light activation. The aim of the study was to show the feasibility to remotely induce the differentiation of leukaemic cells in the bone marrow and not to demonstrate the exhaustion of cancer cells, as this will be difficult using the blue laser activation process. The concept here is different from previous studies that have used therapeutic NPs coated with aptamers to direct them to the bone marrow. Although the results achieved are interesting, the side effects of the systemic administration of the NPs containing a powerful anti-leukaemic agent remain to be determined. In the present study we have used cancer biology

**Figure 4 | In vivo differentiation of NB4 cells exposed to light-activatable RA⁺ NPs.** (a) Schematic representation of the in vivo experimental set up. Cells were treated with blank or RA⁺ NPs (both at 10 μg ml⁻¹) for 4 h, washed and then activated or not with a blue optical fibre (405 nm, 80 mW) for 5 min. Cells were then resuspended in a 1:1 (v/v) Matrigel solution and subcutaneously injected in a PDMS cylinder construct implanted in the dorsal region of mice. After 5 days, cells were removed from the construct and characterized by flow cytometry for CD11b expression. (b) Representative flow cytometry plots for mice recipient cells (left), human leukaemia NB4 cells (middle) and a mixture of mice recipient cells with human leukaemia NB4 cells (right) and percentage of CD11b⁺ cells in human leukaemia NB4 cells collected 5 days after subcutaneous injection. (c) Schematic representation of the in vivo experimental set up. Cells were treated with RA⁺ NPs (10 μg ml⁻¹) for 4 h, washed and then encapsulated in a 1:1 (v/v) Matrigel solution and subcutaneously injected in a PDMS cylinder construct implanted in the dorsal region of mice. After 24 h, some experimental groups were activated in vivo with a blue optical fibre for 5 min. Plot shows the percentage of CD11b⁺ cells in human leukaemia NB4 cells collected 3 days after the in vivo activation. In b, c, results are expressed as mean ± s.e.m. (n = 3–4). Statistical analyses were performed by one-way Anova followed by a Newman-Keuls post-test. **P<0.01, ***P<0.001, ****P<0.0001.
to fight cancer. We have used leukaemia cells loaded with a clinical-approved AML differentiating agent able to home to leukaemic niches in the bone marrow. Although not explored in the current study, normal hematopoietic stem cells may also be used as a nanoparticle carrier as they are able to home to leukaemic niches.32

Methods
Detailed methods are available in Supplementary Information.

Preparation and characterization of PEI conjugated with DMNC. DMNC (48.5 mg, Sigma) was slowly added to a solution of PEI in DMSO (2 ml containing 50 mg ml⁻¹ PEI, Sigma) containing triethylamine (24.5 µl, Sigma), and the reaction flask cooled to 0°C by immersion on ice. Then, the reaction was allowed to proceed for 24 h at 25°C with stirring. At the end, the PEI-DMNC conjugate was purified by dialysis (Spectra/Port 1 Regenerated Cellulose dialysis membrane, MWCO 6000-8000 Da, Spectrum) against DMSO overnight at room temperature. Reaction yields above 54% were obtained. For NMR characterization, PEI-DMNC conjugate was incubated in serum-free RPMI-1640 from 1 to 24 h with variable amounts of RA NPs. After incubations, NPs that were not internalized by the cells were washed (three times with PBS) and the cells were centrifuged; followed by the addition of an aqueous solution of nitric acid (1 ml, 69% v/v). The samples were analysed by ICP-MS for the concentration of Zn. The concentration of Zn was normalized per cell. The estimation of NPs was done based on controlled standard solutions.

Mechanism of NP uptake. U937 cells were cultured on 24-well plates (1 × 10⁵ cells per well) and incubated in serum-free RPMI-1640 from 1 to 24 h with variable amounts of RA NPs. After incubations, NPs that were not internalized by the cells were washed (three times with PBS) and the cells were centrifuged; followed by the addition of an aqueous solution of nitric acid (1 ml, 69% v/v). The samples were analysed by ICP-MS for the concentration of Zn. The concentration of Zn was normalized per cell. The estimation of NPs was done based on controlled standard solutions.

Cell culture. HUVECs (Lonza) were cultured in EGM-2 medium (Lonza) in a CO₂ incubator at 37°C, 5% CO₂ in a humidified atmosphere, with media changes performed every other day. Cells were passaged every 2–3 days and used for experiments between passages 4 and 6. Human bone marrow APL NB4 cells, kindly provided by Dr Arthur Zelent (Institute of Cancer Research, Royal Cancer Hospital), were cultured in RPMI-1640 ( Gibco) supplemented with 10% fetal bovine serum (FBS) (Gibco) and 100 U ml⁻¹ penicillin (Lonza). Human myelomonoblastic cell lines U937-MT and U937-B412, kindly provided by Dr Estelle Duprez (Centre de Recherche en Cancérologie de Marseille, France), were maintained at exponential growth in RPMI-1640 medium supplemented with 10% FBS and 100 U ml⁻¹ of penicillin. For PLZF/RARA induction cells were stimulated with 0.1 mM ZnSO₄ for at least 24 h. Human acute monocytic leukaemia cell line THP-1 (DSMZ no. ACC16) was cultured in RPMI-1640 media supplemented with HEPE (10 mM), sodium pyruvate (1.0 mM) and 2-mercaptoethanol (0.05 mM).

Intracellular trafficking analyses of NPs. HUVECs were stained with FITC-conjugated TRITC labelled NPs for 4 h and then incubated in serum-free RPMI-1640 from 1 to 24 h with variable amounts of RA NPs. After incubations, NPs that were not internalized by the cells were washed (three times with PBS) and the cells were centrifuged; followed by the addition of an aqueous solution of nitric acid (1 ml, 69% v/v). The samples were analysed by ICP-MS for the concentration of Zn. The concentration of Zn was normalized per cell. The estimation of NPs was done based on controlled standard solutions.

Figure 5 | Leukaemia cells transfected with RA NPs can home at leukaemia niches. (a) Schematic representation of the protocol. Human GFP-labelled THP-1 cells were used as the target cells and human THP-1 cells without GFP label were employed as delivery cells. On day 0, the target cells were injected into the NOD/SCID mice by intravenous injection through the tail vein. On day 1, the delivery cells loaded with CRANUMS collected and examined with a confocal microscope. (b) Both target and delivery cells were observed in mice cranium at 24 and 48 h. Bar scale means 100 µm. (c) Delivery cells close (<100 µm) or distant (>100 µm) to target cells were quantified. According to our results, the delivery cells can enter into the same niche as the target cells. Results are expressed as mean ± s.e.m. (n = 3).
NP dilution during cell proliferation. NP dilution with cell growth was monitored over 6 days by ICP-MS by the quantification of intracellular levels of Zn. NB4 and THP-1 cells (0.5 × 10^6 cells per ml) were plated in six-well plates and incubated in serum-free RPMI-1640 with 20 μg ml⁻¹ of RA⁺ NPs. After 4 h incubation, NPs that were not internalized by the cells were washed three times with PBS and the cells were left to grow at 0.2 × 10^6 cells per ml in complete medium for additional 48 h.

**Figure 6 | In vivo activation of leukaemic THP-1 differentiation program by light at the bone marrow.** (a) Schematic representation for the activation of human THP-1 cells transfected with RA⁺ NPs in the bone marrow niche leading to the modulation of leukaemia cells. (b) Percentage of human CD45⁺ cells in the long bones as evaluated by flow cytometry. NOD/SCID mice were injected intravenously with THP-1 cells transfected with RA⁺ NPs through the tail vein. After 6 days, animals (n = 6) were either irradiated by a blue laser (5 min, 405 nm, 80 mW) or not in the calvaria (n = 6). Three days after irradiation, mice were killed and the long bones were collected. Cells in the long bones were characterized by flow cytometry for the expression of human CD45 epitope. The engraftment of THP-1 cells was slightly higher in the non-treated than light-activated animals, but not statistically different. Percentages of positive cells were calculated based in the isotype controls and are shown in each scatter plot. Results are mean ± s.e.m. (n = 3). (c–e) Differentiation of human CD45⁺ cells into CD11b⁺ cells, as evaluated by immunofluorescence, in the calvaria of animals irradiated or not with a blue laser. B, bone; BM, bone marrow. In d, results are mean ± s.e.m. (n = 3). ***P < 0.001. (f) Expression of CRABP2 in cells of calvaria of animals irradiated or not with a blue laser. Scale in c,e,f means 200, 50 and 100 μm.
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**Subcutaneous implantation in vivo.** The animal work has been conducted according to relevant national and international guidelines and approved by the Bioethics Committee of University of Salamanca. On the day before injecting the cells, PDMS cylindrical constructs (40 mm × 10 mm; exterior × interior = 1.5 cm) were implanted subcutaneously on NOD/SCID mice (Jackson Laboratory) maintained under appropriate conditions with anaesthesia. For the experiment in the day of the experiment, NB4 were suspended in serum-free medium with (i) no NPs, (ii) with empty NPs (10 μg ml⁻¹) or RA⁺ NPs (10 μg ml⁻¹) for 4 h. At the end, cells were washed by centrifugation (1,300 r.p.m., 5 min), and the ones treated with RA⁺ NPs were either adopted or not with a blue laser (405 nm, 80 mW) for 5 min. Next, NB4 cells (5 × 10⁶ cells per PDMS construct) were injected subcutaneously in the centre of the PDMS construct embedded in Matrigel (200 μl, BD Biosciences). Five days after injection of the cells, animals were killed by cervical dislocation and cells within the cylindrical construct were collected and characterized by flow cytometry. For the in vivo activation studies in the day of the experiment, NB4 cells were suspended in serum-free medium with (i) no NPs, (ii) with RA⁺ NPs (10 μg ml⁻¹) for 4 h. At the end, cells were washed by centrifugation (1,300 r.p.m., 5 min), and 5 × 10⁴ NB4 cells per PDMS construct were injected subcutaneously in the centre of the PDMS construct embedded in Matrigel (200 μl, BD Biosciences). One day after injection, experimental groups were either activated or not with a blue optical fibre (405 nm, 80 mW). Three days after injection of the cells, animals were killed by cervical dislocation and cells within the cylindrical construct were collected and characterized by flow cytometry.

**Homing of leukaemia cells to leukaemic niches in vivo.** NOD.CB17-Prkdcscid/J (NOD/SCID) female mice, 6–8-week-old, received 1.5 Gy of total body irradiation from a 137Cs source and were treated with mouse anti-CD122 monoclonal antibody (200 μg) by intraperitoneal injection. Human GFP-labelled THP-1 cells were employed as the target cells and human THP-1 cells without GFP label were employed as the delivery cells. On day 0, the target cells (10,000,000 cells per mouse) were injected into the NOD/SCID mice by intravenous injection through the tail vein. On day 1, the delivery cells (10,000,000 cells per mouse; cells were suspended in 200 μl of PBS) were injected into the NOD/SCID mice by intravenous injection through the tail vein. The loading of the cells was performed as follows: Cells were incubated for 4 h with TRITC-labelled RA⁺ NPs (20 μg ml⁻¹) in RPMI medium, washed with PBS to remove loosely bound NPs, resuspended in RPMI medium with 10% FBS for 18 h and then injected in the animals that had received the target cells. After 24 h (n = 3) or 48 h (n = 3) of the delivery cell injection, the mice were killed, the craniums collected and examined with a confocal microscope.

**Bone marrow modulation in vivo.** NOD.CB17-Prkdcscid/J (NOD/SCID) female mice (n = 12) aged 6–8 weeks were employed in this experiment. Before cell injection, mice received 1.5 Gy of total body irradiation from a 137Cs source and were also treated with 200 μg of mouse anti-CD122 monoclonal antibody (200 μg) by intraperitoneal injection. Human THP-1 cells were incubated with 20 μg ml⁻¹ of RA⁺ NPs in RPMI medium for 4 h, followed by extensive wash with PBS to remove non-internalized NPs. The cells were then resuspended in RPMI medium with 10% FBS and left in the culture incubator overnight. On the following day, the cells loaded with RA⁺ NPs were collected and 1 × 10⁶ cells per mouse in 200 μl PBS were injected into the NOD/SCID mice intravenously, through the tail vein. All the mice were thus divided into two groups: the first group was blue laser irradiated (n = 6) and the other group was not irradiated (n = 6). In the blue laser irradiated group, each mouse was anaesthetized with 3.5% chloral hydrate (405 nm, 80 mW) for 5 min. The mice in the non-irradiated group received the same treatment without blue laser irradiation. The bone was divided into two parts: the parietal and the occipital bone. The long bones including the femurs, tibias and pelvis were crushed in PBS containing 1% bovine serum albumin (BSA, sigma) and 2 mM EDTA (Invitrogen). The samples were then filtered with 70 μm cell strainers and treated with ACK solution to lyse red blood cells. The resulting cells were stained with PE-conjugated anti-human CD45 antibody (eBioscience). The results were measured by flow cytometry and the data were analysed with FlowJo software. The mouse craniums were collected for the ex vivo staining examination followed by the traditional protocol. Briefly, the mouse craniums were cut into four pieces: two parietal and one occipital part bone. Only parietal bones were used in this experiment. Bone pieces were fixed in 4% paraformaldehyde for 30 min, washed twice with PBS, blocked with 2% BSA-0.01%
Triton X-100 in PBS (BSA buffer) for 1 h, and incubated with primary antibodies (diluted 1:100 in BSA buffer) overnight at 4 °C. After washing with BSA buffer for 2 h, bones were incubated with secondary antibodies (diluted 1:200 in BSA buffer) for 2 h and then washed with BSA buffer. Nuclei were stained using DAPI. The bone pieces were finally examined with a confocal microscope (Nikon Eclipse Ti) under FITC/TRITC/DIC signal channel. Images were analysed in ImageJ.

Analyse Particles was used after thresholding to quantitatively measure signals and measure the total area occupied by these particles.

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Acknowledgements

We acknowledge the use of the Laboratório de Bio-imagem de Alta Resolução of the Faculty of Medicine of University of Coimbra. The authors would like to thank the financial support of FEDER through the programme COMPETE and by Portuguese funds through FCT (PTDC/CTM-NAN/120552/2010 and UID/NEU/04539/2013 to R.P.), FCT (SFRH/BPD/62419/2009 to C.B.; SFRH/BD/99642/2012 to E.O.; POCTI-01-0145-FEDER-016390 (CANCER STEM to L.F.), IC (ERC project n° 307384, ‘Nanotrigger’ to L.F.), MINECO (SAF2012-32810, SAF2015-64420-R and Red de Excelencia Con- solider OncBioSAF2014-57791-REDIC to I.S.-G.) and Bloodwise and CRUK programme grants to T.E. and D.H. contributed with animal studies and analysed data; A.M.-L., M.B.G.C. and D.H. contributed with animal studies; S.P. carried out cell characterization analyses; R.G. and T.E. contributed for the cell studies; R.P.d.N. and L.F. contributed conceptually, made measurements, analysed data and wrote the manuscript. All the authors have read and approved the manuscript.

Author contributions

C.B. contributed conceptually to the design of the nanoparticle system and together with E.O. carried out cell and animal studies, analysed data and wrote the manuscript; Y.C. and D.H. contributed with animal studies and analysed data; A.M.-L., M.B.G.C. and D.H. contributed for animal studies; S.P. carried out cell characterization analyses; R.G. and T.E. contributed conceptually, made measurements, analysed data and wrote the manuscript. All the authors have read and approved the manuscript.

ARTICLE NATURE COMMUNICATIONS | DOI: 10.1038/ncomms15204
