Tetraphenylethylene-conjugated polycation covered iron oxide nanoparticles for magnetic resonance/optical dual-mode imaging

Li Yang¹, Shengxiang Fu¹, Li Liu¹, Zhongyuan Cai¹, Chunchao Xia², Bin Song², Qiyong Gong³,⁴, Zhiyun Lu⁵ and Hua Ai¹,², *

¹National Engineering Research Center for Biomaterials, Sichuan University, Chengdu 610065, China; ²Department of Radiology, West China Hospital, Sichuan University, Chengdu 610041, China; ³Huaxi MR Research Center (HMRRC), Department of Radiology, West China Hospital of Sichuan University, Chengdu, China; ⁴Psychoradiology Research Unit of Chinese Academy of Medical Sciences, Sichuan University, Chengdu, China and ⁵Key Laboratory of Green Chemistry and Technology (Ministry of Education), College of Chemistry, Sichuan University, Chengdu 610065, China

*Correspondence address. National Engineering Research Center for Biomaterials, No.29 Jiuyanqiao Wangjiang Road, Sichuan University, Chengdu, 610065, China. Tel: +86-28-8541-3991; Fax: +86-28-8541-3991; E-mail: huaai@scu.edu.cn

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Abstract

Magnetic resonance (MR)/optical dual-mode imaging with high sensitivity and high tissue resolution have attracted many attentions in biomedical applications. To avert aggregation-caused quenching of conventional fluorescence chromophores, an aggregation-induced emission molecule tetraphenylethylene (TPE)-conjugated amphiphilic polyethylenimine (PEI) covered superparamagnetic iron oxide (Alkyl-PEI-LAC-TPE/SPIO nanocomposites) was prepared as an MR/optical dual-mode probe. Alkyl-PEI-LAC-TPE/SPIO nanocomposites exhibited good fluorescence property and presented higher T² relaxivity (352 Fe mM⁻¹s⁻¹/C₀) than a commercial contrast agent Feridex (120 Fe mM⁻¹s⁻¹/C₀) at 1.5 T. The alkylation degree of Alkyl-PEI-LAC-TPE effects the restriction of intramolecular rotation process of TPE. Reducing alkane chain grafting ratio aggravated the stack of TPE, increasing the fluorescence lifetime of Alkyl-PEI-LAC-TPE/SPIO nanocomposites. Alkyl-PEI-LAC-TPE/SPIO nanocomposites can effectively labelled HeLa cells and resulted in high fluorescence intensity and excellent MR imaging sensitivity. As an MR/optical imaging probe, Alkyl-PEI-LAC-TPE/SPIO nanocomposites may be used in biomedical imaging for certain applications.

Keywords: magnetic resonance imaging; optical imaging; superparamagnetic iron oxide; aggregation-induced emission; dual-mode imaging

Introduction

Recently, aggregation-induced emission (AIE) [1, 2] has been widely studied and used in biomedical imaging applications, such as non-specific cell imaging [3], specific organelle imaging [4] and long-term cell tracking [5]. Different from conventional fluorescence chromophores’ aggregation-caused quenching, AIE molecules have significantly enhanced fluorescence in the aggregated state [1, 2, 6]. AIE nanoprobes with excellent photostability and easy cell internalization have attracted attentions in biomedical imaging, but weak tissue penetration and low spatial resolution limit their further applications [7, 8].

Multi-mode imaging modalities such as magnetic resonance (MR)/ optical [9, 10], MR/positron emission tomography (PET) [11, 12] and PET/computed tomography [13, 14] with complementary information about anatomical structure and function have been extensively explored and developed in biomedical applications. MR/optical imaging has its unique advantages. MRI is a non-invasive medical imaging technique with high spatial resolution to acquire subtle anatomical and functional information of tissues and organs. However, the sensitivity of MRI is relatively low than other imaging modalities such as PET or optical imaging. The combination of these MRI and optical
imaging modalities enables one to achieve both detailed tissue resolution and high sensitivity. Recently, the clinically used gadolinium (Gd)-based MR $T_1$ contrast agents have been reported to combine with AIE molecules as a dual-mode imaging probe which realized the liver-specific imaging [15], the visualization and quantification of changes in brain barrier functions [16]. However, Gd-based MR contrast agents have a strong correlation with nephrogenic systemic fibrosis in patients with renal insufficiency, which limits their applications [17, 18]. Due to their excellent biosafety, high sensitivity and easy modification, iron oxide nanoparticles (IONPs)-based MR $T_2$ contrast agents have been used for cancer diagnosis [19, 20], atherosclerosis diagnosis [21, 22], long-term cell tracking [23, 24] in clinical trials. In addition, previous studies showed that multiple superparamagnetic iron oxide (SPIO) nanocrystal containing micelles have much higher $T_2$ relaxivity than single SPIO nanocrystal suspensions at the same Fe concentration [25]. The controlled aggregation of SPIO nanocrystals can enhance relaxivity, similar to the effects that AIE molecules brought to us. So, the development of MR/optical imaging probes based on IONP and AIE molecules may find its unique applications.

Herein, an MR/optical imaging probe based on Alkyl-PEI-LAC-TPE/SPIO nanocomposites was developed (Scheme 1). Amphiphilic polymer Alkyl-PEI-LAC-TPE was designed for encapsulation of hydrophobic SPIO nanocrystals, the modification of lactobionic acid (LAC) on alkyl-polyethylenimine (PEI) was used to reduce the toxicity of PEI and the conjugation of tetraphenylethylene (TPE) endowed both hydrophobic and AIE features. Besides, different grafting degrees of alkane chain (C12) on polymer were used to understand the impact of amphiphilic structure on TPE’s fluorescence property. Then, SPIO nanocrystals were encapsulated inside the hydrophobic core of Alkyl-PEI-LAC-TPE micelles by self-assembly. The size distribution and morphology of Alkyl-PEI-LAC-TPE/SPIO nanocomposites were investigated, and the fluorescence properties and in vitro fluorescence imaging were evaluated, respectively. Additionally, $T_2$ relaxivity and in vitro MRI study of Alkyl-PEI-LAC-TPE/SPIO nanocomposites were tested under clinical MRI scanners.

Materials and methods

Materials

PEI (MW:1.8 kD), N-Hydroxysuccinimide (NHS, 98%), N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride (EDC, 98%) and 1-iodododecane (>98%) were purchased from Aladdin Industrial Corporation. Aldehyde functionalized tetraphenylethene (TPE-CHO) was purchased from AIEgen Biotech Co., Ltd. Lactobionic acid (LAC) was purchased from TCI (Shanghai) Chemical Industry Development Corporation. Iron (III) acetylacetonate (97%), 1,2-hexadecanediol (90%), benzyl ether (98%), oleic acid (70%) and oleylamine (70%) were purchased from Sigma-Aldrich Corporation. All other chemicals and solvents were of analytical grade and used without further purification.

**Scheme 1. The illustrated structure of Alkyl-PEI-LAC-TPE/SPIO nanocomposites**

**Synthesis of Alkyl-PEI-LAC-TPE polymer**

The synthesis route of Alkyl-PEI-LAC-TPE was displayed in Fig. S1 (online supplementary material). PEI-LAC was synthesized followed the reported protocol [26] with slight modification. Then, TPE-CHO was added for reaction with the amine of PEI-LAC. 1-Iodododecane was reacted with PEI-LAC-TPE in dimethyl sulfoxide (DMSO)/dimethylformamide (DMF) mixed solution following a previously reported method [25, 27] with slight modification.

PEI (4.3 g, 100 mmol) was dissolved in absolute DMSO (30 ml), LAC (3.58 g, 10 mmol), EDC (2.1 g, 11 mmol) and NHS (1.27 g, 11 mmol) were added and dissolved in anhydrous DMSO (40 ml) in a 100-ml round-bottom flask, and performed at 30°C for 4 h under stirring. The activated LAC was added dropwise to a stirring solution of PEI under nitrogen. The reaction was performed at 30°C for 24 h under stirring. The crude product (PEI-LAC) was dialyzed against deionized water (molecular mass cutoff, 1 kDa), collected and freeze-dried.

PEI-LAC (5.5 g, 70 mmol) and TPE-CHO (2.5 g, 7 mmol) were dissolved in anhydrous DMSO, respectively. TPE-CHO was added dropwise to a stirring solution of PEI-LAC, and the reaction was performed at 55°C for 18 h under stirring. The crude product was added dropwise to cold ethyl ether, and the target product PEI-LAC-TPE in the bottom was collected, dialyzed against deionized water and freeze-dried. 1-Iodododecane (0.24 g, 0.8 mmol) was added dropwise to PEI-LAC-TPE (0.92 g, 8 mmol) in DMSO/DMF mixed solution under nitrogen, and heated to 55°C for 6 h with stirring. After cooling to room temperature, the solution was incubated overnight. The crude product was dialyzed against gradient EtOH/H$_2$O solution. By controlling the ratio of 1-iodododecane and PEI-LAC-TPE, Alkyl-PEI-LAC-TPE with different alkane grafting degrees were synthesized. The crude product (Alkyl-PEI-LAC-TPE) was collected and freeze-dried. Alkyl-PEI-LAC-TPE with different grafting degrees of alkane chains were synthesized to investigate probes’ fluorescence properties.

**Characterization of Alkyl-PEI-LAC-TPE polymer**

$^1$H NMR spectra of PEI, PEI-LAC, PEI-LAC-TPE and Alkyl-PEI-LAC-TPE were acquired on an NMR spectrometer (Bruker, 400 MHz). Element analysis of PEI, PEI-LAC and PEI-LAC-TPE were acquired on an element analyser (Teledyne Leeman Labs). Fourier-transform infrared spectroscopy (FTIR) spectra of PEI, PEI-LAC, PEI-LAC-TPE and Alkyl-PEI-LAC-TPE were recorded on a FTIR spectrometer (Thermo Fisher Scientific).

**Synthesis of SPIO nanocrystals**

SPIO nanocrystals were synthesized following a reported method from Sun et al. [28]. Briefly, Fe(acac)$_3$ (2 mmol), 1,2-hexadecanediol (2 mmol), oleic acid (2 mmol) and oleylamine (2 mmol) were added in benzyl ether (20 ml) with nitrogen. The mixture was heated to 200°C for 2 h, then the mixture was heated to reflux for 1 h. After cooling to room temperature, ethyl acetate was used to yield a
HeLa cells were cultured in DMEM medium containing 10% FBS at 37°C with 5% CO₂. HeLa cells were plated at a density of 5 × 10⁴ cell per well in a 24-well plate. After cell attachment, Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were diluted into DMEM medium with TPE concentration of 2.5 μg/ml for 12 h. Next, the medium was removed and the cells were washed with PBS twice. After the cells were fixed with 4% paraformaldehyde for 15 min, the fluorescence images were captured under a confocal laser scanning microscope (CLSM). The mean fluorescence intensities of HeLa cells incubated with Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were calculated by Image J.

**Preparation of Alkyl-PEI-LAC-TPE micelles and SPIO-loaded Alkyl-PEI-LAC-TPE (Alkyl-PEI-LAC-TPE/SPIO) nanocomposites**

SPIO nanocrystals in hexane were dried with nitrogen, then dispersed in chloroform. Alkyl-PEI-LAC-TPE was dissolved in DMSO, then added into chloroform under sonication. Then, the mixture of Alkyl-PEI-LAC-TPE with SPIO at a mass ratio of 3:1 was added into deionized water under sonication and kept shaking for 0.5 h. Chloroform was removed from by evaporation and dialyzed against deionized water to clear DMSO to obtain Alkyl-PEI-LAC-TPE/SPIO nanocomposites. The fabrication of Alkyl-PEI-LAC-TPE micelles was following the same protocol.

**Size distribution and morphology of SPIO nanocrystals, Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites**

The size distribution of SPIO, Alkyl-PEI-LAC-TPE and Alkyl-PEI-LAC-TPE/SPIO nanocomposites was performed at 25°C via dynamic light scattering (DLS) (Zetasizer Nano ZS, Malvern Instruments). The surface morphology of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were investigated by scanning electron microscopy (S-4800, Hitachi), for which 50 μl of the samples were dried on a piece of silicon. The size distribution and ultrastructure of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were studied by transmission electron microscopy (TEM) (Tecnai G2 F20 S-TWIN, FEI), for which 10 μl of the samples were dried on a copper grid.

**Fluorescence properties of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites**

The ultraviolet absorption (UV) spectrum of Alkyl-PEI-LAC-TPE micelles was obtained by ultraviolet–visible (UV–VIS) spectrophotometer (U-3900, Hitachi). The emission spectrum and fluorescence intensity at the maximum emission wavelength of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were investigated by scanning electron microscopy (S-4800, Hitachi), for which 50 μl of the samples were dried on a piece of silicon. Characteristic peaks of PEI-LAC were appeared in the 1H NMR spectrum (400 MHz, DMSO-d₆). The characteristic peaks at 3410 cm⁻¹ (-OH), 1650–1560 cm⁻¹ (-C–O) were attributed to LAC, indicating the successful modification with PEI-LAC. Characteristic peaks of PEI-LAC-TPE were displayed in Fig. S2 (online supplementary material).

**Results and discussion**

**Synthesis and characterization of Alkyl-PEI-LAC-TPE polymer**

The 1H NMR spectrum of PEI, PEI-LAC, PEI-LAC-TPE and Alkyl-PEI-LAC-TPE was shown in Fig. 1. Low molecular weight branched PEI was firstly modified with LAC to improve its biocompatibility. Characteristic peaks of PEI-LAC were appeared in the 1H NMR spectrum (400 MHz, DMSO-d₆): δ (ppm) 4.33–2.96 (LAC), 2.81–2.09 (–CH₂–CH₃–NH–). TPE with good AIE effect was then conjugated with PEI-LAC. Characteristic peaks of PEI-LAC-TPE were appeared in the 1H NMR spectrum (400 MHz, DMSO-d₆): δ (ppm) 7.19–6.74 (TPE), 4.39–2.91 (LAC), 2.83–1.90 (–CH₂–CH₃–NH–). To explore the effect of alkane chains on the fluorescence properties of TPE, PEI-LAC-TPE was modified with different degrees of alkane chains by reacting with 1-iododecane. Characteristic peaks of Alkyl-PEI-LAC-TPE were appeared in the 1H NMR spectrum (400 MHz, DMSO-d₆): δ (ppm) 7.22–6.74 (TPE), 4.31–2.94 (LAC), 2.89–1.90 [-CH₂–CH₃–NH–, –OCH₃–(CH₃)–O–CH₃], 1.31–1.02 [-CH₂–(CH₃)–O–CH₃], 0.81 [-CH₂–(CH₃)–O–CH₃]. The 1H NMR spectrum of Alkyl-PEI-LAC-TPE with different grafting degrees of alkane chains was displayed in Fig. S2 (online supplementary material).

**FTIR was used to characterize PEI-LAC, PEI-LAC-TPE and Alkyl-PEI-LAC-TPE. As shown in Fig. 2, the characteristic peaks at 3290 and 1460 cm⁻¹ were attributed to the –NH and at 2940–2820 cm⁻¹ were attributed to the –CH₃ of PEI. The characteristic peaks at 3410 cm⁻¹ (–OH), 1630–1560 cm⁻¹ (–C=O) and 1060 cm⁻¹ (–C–O) were attributed to LAC, indicating the successful synthesis of PEI-LAC. For PEI-LAC-TPE, a peak of the stretching vibration of aromatic rings of TPE at 750–700 cm⁻¹ could be seen.
The -CH₂ peak of Alkyl-PEI-LAC-TPE at 2940–2820 cm⁻¹/C₀ was strengthened after modified with alkane chains. The LAC and TPE grafting ratios were calculated by elemental analysis according to the changes of ratio of carbon (C) and nitrogen (N) in PEI, PEI-LAC and PEI-LAC-TPE. The results showed that the LAC and TPE grafting ratios were 11% and 10%, respectively.

Besides, the alkane chain grafting ratio was calculated by the ¹H NMR of Alkyl-PEI-LAC-TPE according to the integral values at δ 7.22–6.74 and δ 0.81 ppm (Fig. 1 and Fig. S2, online supplementary material). Results showed that the alkane chain grafting ratios of Alkyl-PEI-LAC-TPE were 13%, 27% and 40%, respectively.

Size distribution and morphology of SPIO nanocrystals, Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites

Organic SPIO nanocrystal has a narrow size distribution with a mean diameter of 9 nm characterized by DLS in n-hexane (Fig. 3). Alkyl-PEI-LAC-TPE and Alkyl-PEI-LAC-TPE/SPIO nanocomposites with gradient alkane chains were prepared by self-assembly in deionized water. Alkyl-PEI-LAC-TPE micelles with gradient alkane chains have a similar diameter of 70 nm measured by DLS (Fig. 3), indicating that the different of alkane chain grafting ratios did not have significant impact on their size distribution. Alkyl-PEI-LAC-TPE/SPIO nanocomposites with gradient alkane chains have a similar diameter of 50 nm in DLS, smaller than that of Alkyl-PEI-LAC-TPE micelles. TEM results (Fig. 3) showed that Alkyl-PEI-LAC-TPE micelles were spherical-like structure, and organic SPIO nanocrystals were encapsulated in Alkyl-PEI-LAC-TPE micelles as isolated spherical nanocluster structure.

Fluorescence properties of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites

As fluorescence probes, the fluorescence properties of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites were studied, respectively. As shown in Fig. 4, the UV, excitation and emission spectrum of Alkyl-PEI-LAC-TPE micelles of different alkyl chains were the same in deionized water, with max excitation wavelength of 330 nm and max emission wavelength of 480 nm, indicating that Alkyl-PEI-LAC-TPE micelles with gradient alkyl chains had same fluorescence chromophore. Solid powders of gradient alkyl chain Alkyl-PEI-LAC-TPE appeared yellow at day-light and emitted a blue light excited by a UV-light (Fig. S3, online supplementary material).

The fluorescence intensity of gradient alkyl chain Alkyl-PEI-LAC-TPE micelles was measured at a series of TPE concentrations to evaluate the effect of alkane chain grafting ratios on the fluorescence properties of Alkyl-PEI-LAC-TPE. As shown in Fig. 5, the fluorescence intensity of Alkyl-PEI-LAC-TPE micelles with different alkyl chains was increased with the increasing of TPE concentration. For the same TPE concentration, the fluorescence intensity of Alkyl-PEI-LAC-TPE micelles was ranging from strong to weak with 13%, 27% and 40% alkyl chains (Fig. 5). In other words, the lower of alkane chain grafting ratio, the higher of fluorescence intensity at the same TPE concentration. It was possible that the different degrees of alkane chain grafting ratios of Alkyl-PEI-LAC-TPE had an effect on the fluorescence property.

To understand if the degree of alkane chain grafting ratios has any effect on the fluorescence property of Alkyl-PEI-LAC-TPE micelles, their fluorescence lifetime was measured. Fluorescence lifetime of fluorescence probes can be used to evaluate the impact of microenvironment (solvent polarity, viscosity, hydrophobicity, pH, etc.) around the fluorophores on their fluorescence property [29]. To obtain fluorescence lifetime, the fluorescence decay curves of gradient alkyl chain Alkyl-PEI-LAC-TPE micelles were measured by transient fluorescence spectrophotometer with the excitation wavelength of 370 nm and the emission wavelength of 480 nm, exhibiting biexponential decay in deionized water system (Fig. 6). The average
fluorescence lifetime of Alkyl-PEI-LAC-TPE micelles were 1.71, 1.29 and 1.05 ns according to the alkyl chain ratios of 13%, 27% and 40%, respectively (Table 1). The lower alkane chain grafting ratio, the longer the fluorescence lifetime, and the higher fluorescence intensity of Alkyl-PEI-LAC-TPE micelles.

As previously reported, restriction of intramolecular rotation (RIR) is the main mechanism for the AIE characteristic of TPE fluorophores [6, 30, 31]. The tightly stacked of TPE molecules in the hydrophobic core of Alkyl-PEI-LAC-TPE micelles resulted in the RIR process of TPE and that accounts for its AIE characters. The TPE and alkane chains together formed the hydrophobic core of Alkyl-PEI-LAC-TPE micelles. Therefore, the surrounding alkane chains near TPE molecules may affect the RIR process of TPE, reflecting in the change of fluorescence lifetime and intensity of Alkyl-PEI-LAC-TPE micelles. The lower of alkane chain grafting ratio, the more tightly the stacked of TPE, the higher degree of RIR process, the longer of fluorescence lifetime and the higher of fluorescence intensity of Alkyl-PEI-LAC-TPE micelles.

In addition, the fluorescence lifetime of Alkyl-PEI-LAC-TPE/SPIO nanocomposites have also been studied. As shown in Fig. S4 (online supplementary material), the fluorescence decay curves of Alkyl-PEI-LAC-TPE/SPIO nanocomposites exhibited biexponential decay, the same as Alkyl-PEI-LAC-TPE micelles. And the average fluorescence lifetime of Alkyl-PEI-LAC-TPE/SPIO nanocomposites decreased from 1.53, 1.20 to 1.00 ns as Alkyl-PEI-LAC-TPE grafted with 13%, 27% and 40% alkyl chains, respectively (Table S1,

![Figure 3](image-url)  
**Figure 3.** (a) DLS of SPIO nanocrystals in hexane, Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites in H2O. (b) TEM images of Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites with indicated alkyl chain gradients

![Figure 4](image-url)  
**Figure 4.** UV spectrum (a), excitation spectrum and emission spectrum (b) of gradient alkyl chain Alkyl-PEI-LAC-TPE micelles in H2O
The fluorescence lifetime of Alkyl-PEI-LAC-TPE/SPIO nanocomposites were slightly shorter than Alkyl-PEI-LAC-TPE micelles. The encapsulated SPIO nanocrystals did not have major impact on the fluorescence properties of Alkyl-PEI-LAC-TPE micelles.

$T_2$ relaxivity of Alkyl-PEI-LAC-TPE/SPIO nanocomposites

$T_2$ relaxivities of Alkyl-PEI-LAC-TPE/SPIO nanocomposites with gradient alkyl chains were measured on a 1.5T clinical MRI scanner (Siemens). IONP is an effective MR $T_2$ contrast agent, which can significantly shorten the $T_2$ relaxation time of surrounding water protons [32, 33]. MR images became darker with the increased of iron concentration (Fig. 7b), and the $T_2$ relaxivity of the Alkyl-PEI-LAC-TPE (13%)/SPIO, Alkyl-PEI-LAC-TPE (27%)/SPIO and Alkyl-PEI-LAC-TPE (40%)/SPIO nanocomposites was 284.6, 337.2 and 351.8 Fe mM$^{-1}$s$^{-1}$, respectively. Results were similar to previous studies that nanocluster containing multiple SPIO nanocrystals performed higher $T_2$ relaxivities than those containing single SPIO nanocrystal [25]. What is more, the $T_2$ relaxivities of Alkyl-PEI-LAC-TPE/SPIO nanocomposites became higher with the increased alkane chain grafting ratio. One possible reason might be that there are more SPIO nanocrystals stacked in the hydrophobic core of Alkyl-PEI-LAC-TPE/SPIO nanocomposites with higher alkane chain grafting ratios. Moreover, the $T_2$ relaxivities

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**Table 1.** The average fluorescence lifetimes of gradient alkyl chain Alkyl-PEI-LAC-TPE micelles ($\lambda_{ex} = 370$ nm, $\lambda_{em} = 480$ nm)

| Grafting rate | Lifetime (s/nm) | Percentage (%) | Average lifetime (s/nm) |
|---------------|----------------|---------------|-----------------------|
| 13%           | $\tau_1$ 1.15  | 0.83          | 1.71                  | 1.04                  |
|               | $\tau_2$ 4.39  | 0.17          |                       |                      |
| 27%           | $\tau_1$ 0.92  | 0.85          | 1.29                  | 1.02                  |
|               | $\tau_2$ 3.38  | 0.15          |                       |                      |
| 40%           | $\tau_1$ 0.74  | 0.86          | 1.05                  | 1.02                  |
|               | $\tau_2$ 2.88  | 0.14          |                       |                      |
of all Alkyl-PEI-LAC-TPE/SPIO nanocomposites were higher than the commercial contrast agent Feridex (120 Fe mM$^{-1}$s$^{-1}$, 1.5T) [32]. This imaging property indicated that Alkyl-PEI-LAC-TPE/SPIO nanocomposites may be used as an effective $T_2$ contrast agent or cell labelling probe.

**In vitro** fluorescence imaging of labelled HeLa cells with Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites

Both Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites have shown good fluorescence properties, and their efficacies were tested with cell labelling. Alkyl-PEI-LAC-TPE/SPIO nanocomposites shown good biocompatibility to HeLa cells with TPE concentration at 2.5 mg/ml (Fig. S5, online supplementary material). CLSM was used to study the fluorescence imaging of labelled HeLa cells with Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites. As shown in Fig. 8, after 12 h incubation, Alkyl-PEI-LAC-TPE/SPIO nanocomposites were efficiently internalized by HeLa cells and mainly accumulated in the cytoplasm with high fluorescence intensity, the same as Alkyl-PEI-LAC-TPE micelles. Besides, similar fluorescence intensity in Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites labelled HeLa cells were detected (Fig. S6, online supplementary material), which further reveal the encapsulated SPIO nanocrystals did not have major impact on the fluorescence properties of Alkyl-PEI-LAC-TPE micelles.

**In vitro** MRI study of the labelled HeLa cells with Alkyl-PEI-LAC-TPE/SPIO nanocomposites

As a dual-mode MR/fluorescence imaging probe, Alkyl-PEI-LAC-TPE/SPIO nanocomposites should have both MR and fluorescence imaging capabilities. In order to evaluate the probes’ MR visibility, HeLa cells were labelled by Alkyl-PEI-LAC-TPE/SPIO nanocomposites with different Fe concentration (5, 10, 15 and 20 µg/ml) for 12 h. Results (Fig. S7, online supplementary material) showed that Alkyl-PEI-LAC-TPE/SPIO nanocomposites displayed a dose-dependent labelling pattern with excellent labelling efficiency for HeLa cells. MRI study of the labelled HeLa cells with Alkyl-PEI-LAC-TPE/SPIO nanocomposites were evaluated on a 3.0 T clinical MRI scanner. HeLa cells were first labelled with Alkyl-PEI-LAC-TPE/SPIO nanocomposites at an Fe concentration of 5 µg/ml for 12 h. Then, the labelled HeLa cells were collected and homogenously dispersed in gelatine at graded cell numbers. MR images (Fig. 9b) showed that the labelled HeLa cells induced obvious reduction of $T_2$-weighted signal intensity, which was inversely to the number of cells. As shown in Fig. 9a, $T_2$ value decreased from 216 to 61 ms with the increasing number of Alkyl-PEI-LAC-TPE (13%)/SPIO labelled HeLa cells, which was similar to the results of Alkyl-PEI-LAC-TPE (27%)/SPIO and Alkyl-PEI-LAC-TPE (40%)/SPIO nanocomposites shown good biocompatibility to HeLa cells with TPE concentration at 2.5 µg/ml (Fig. S5, online supplementary material). CLSM was used to study the fluorescence imaging of labelled HeLa cells with Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites. As shown in Fig. 8, after 12 h incubation, Alkyl-PEI-LAC-TPE/SPIO nanocomposites were efficiently internalized by HeLa cells and mainly accumulated in the cytoplasm with high fluorescence intensity, the same as Alkyl-PEI-LAC-TPE micelles. Besides, similar fluorescence intensity in Alkyl-PEI-LAC-TPE micelles and Alkyl-PEI-LAC-TPE/SPIO nanocomposites labelled HeLa cells were detected (Fig. S6, online supplementary material), which further reveal the encapsulated SPIO nanocrystals did not have major impact on the fluorescence properties of Alkyl-PEI-LAC-TPE micelles.
nanocomposites labelled HeLa cells. These nanocomposites had demonstrated excellent MR imaging sensitivity in label HeLa cells and may find potential applications in cell imaging.

Conclusion

In conclusion, we have developed an MR/optical imaging probe by encapsulating multiple hydrophobic SPIO nanocrystals in AIE-based amphiphilic Alkyl-PEI-LAC-TPE polymer micelles. Alkyl-PEI-LAC-TPE/SPION nanocomposites are spherical with controllable size distribution. These nanocomposites exhibited good AIE properties with bright blue emission. Moreover, lower alkylation degree of Alkyl-PEI-LAC-TPE was helpful for longer fluorescence lifetime and aggregation-induced emission characteristic for liver specific imaging with long circulation lifetime. In conclusion, we have developed an MR/optical imaging probe by encapsulating multiple hydrophobic SPIO nanocrystals in AIE-based amphiphilic Alkyl-PEI-LAC-TPE polymer micelles. Alkyl-PEI-LAC-TPE/SPION nanocomposites are spherical with controllable size distribution. These nanocomposites exhibited good AIE properties with bright blue emission. Moreover, lower alkylation degree of Alkyl-PEI-LAC-TPE was helpful for longer fluorescence lifetime and aggregation-induced emission characteristic for liver specific imaging with long circulation lifetime.

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Conflict of interest statement. None declared.

Supplementary data

Supplementary data are available at REGBIO online.

References

1. Mei J, Leung NL, Kwok RT et al. Aggregation-induced emission: together we shine, united we soar! Chem Rev 2015;115:11718–940.

2. Hu R, Leung NL, Tang BZ. AIE macromolecules: syntheses, structures and functionalities. Chem Soc Rev 2014;43:4494–562.

3. Wang Z, Xu B, Zhang L et al. Folic acid-functionalized mesoporous silica nanospheres hybridized with AIE luminogens for targeted cancer cell imaging. Nanoscale 2013;5:2065–72.

4. Li S, Ling X, Lin Y et al. In situ generation of photoactivatable aggregation-induced emission probes for organelle-specific imaging. Chem Sci 2018;9:5730–5.

5. Liow SS, Dou Q, Kai D et al. Long-term real-time in vivo drug release monitoring with AIE thermogelling polymer. Small 2017;13:1603404.

6. Hong Y, Lam JW, Tang BZ. Aggregation-induced emission. Chem Soc Rev 2011;40:5361–88.

7. Wang LV, Yao J. A practical guide to photoacoustic tomography in the life sciences. Nat Methods 2016;13:627–38.

8. Rao J, Dragulescu-Andrasi A, Yao H. Fluorescence imaging in vivo: recent advances. Curr Opin Biotechnol 2007;18:17–25.

9. Yang KM, Cho HI, Choi HJ et al. Synthesis of water well-dispersed PEGylated iron oxide nanoparticles for MR/optical lymph node imaging. J Mater Chem B 2014;2:3535–64.

10. Song XR, Li SH, Dai J et al. Polyphenol-inspired facile construction of smart assemblies for ATP- and pH-responsive tumor MR/optical imaging and photothermal therapy. Small 2017;13:1603997.

11. Kaittanis C, Shaffer TM, Bolaender A et al. Multifunctional MRI/PET nanobeacons derived from the in situ self-assembly of translational polymers and clinical cargo through coalescent intermolecular forces. Nano Lett 2015;15:8032–43.

12. Lux J, White AG, Chan MN et al. Nanogels from metal-chelating cross-linkers as platforms applied to copper-64 PET imaging of tumors and metastases. Theranostics 2015;5:277–88.

13. Rahman WT, Wale DJ, Viglianti BL et al. The impact of infection and inflammation in oncologic (18)F-FDG PET/CT imaging. Biomed Pharmacother 2019;117:109168.

14. Beyer T, Townsend DW, Brun T et al. A combined PET/CT scanner for clinical oncology. J Nucl Med 2000;41:1369–79.

15. Chen Y, Li M, Hong Y et al. Dual-modal MRI contrast agent with aggregation-induced emission characteristic for liver specific imaging with long circulation lifetime. ACS Appl Mater Interfaces 2014;6:10783–91.

16. Feng G, Li JLY, Claser C et al. Dual modal ultra-bright nanodots with aggregation-induced emission and gadolinium-chelation for vascular integrity and leakage detection. Biomaterials 2018;152:77–85.

17. Weller A, Barber JL, Olsen OE. Gadolinium and nephrogenic systemic fibrosis: an update. Pediatr Nephrol 2014;29:1927–37.

18. Prince MR, Weinreb JC. Notice of withdrawal: MR imaging and gadolinium: reassessing the risk of nephrogenic systemic fibrosis in patients with severe renal disease. Radiology 2018;286:120–1.

19. Har-lon H, Shu-bo Z, Cai-jin L et al. Organic soluble superparamagnetic iron oxide nanoparticles coated with PEG-PEG-PEG-DDA and MRI imaging for cancer diagnosis. Acta Polym Sin 2014;194:201–201.

20. Xie L, Jin W, Chen H et al. Superparamagnetic iron oxide nanoparticles for cancer diagnosis and therapy. J Biomed Nanotechnol 2019;15:215–416.

21. Montiel Schneider MG, Lassalle VL. Magnetic iron oxide nanoparticles as novel and efficient tools for atherosclerosis diagnosis. Biomed Pharmacother 2017;93:1098–115.

22. Briley-Saebo KC, Cho YS, Shaw PX et al. Targeted iron oxide particles for in vivo magnetic resonance detection of atherosclerotic lesions with antibodies directed to oxidation-specific epitopes. J Am Coll Cardiol 2011;57:337–47.

23. Azevedo-Pereira RL, Rangel B, Tovar-Moll F et al. Superparamagnetic iron oxide nanoparticles as a tool to track mouse neural stem cells in vivo. Mol Biol Rep 2019;44:691–8.

24. Zheng B, Vazin T, Goodwill PW et al. Magnetic particle imaging tracks the long-term fate of in vivo neural cell implants with high image contrast. Sci Rep 2015;5:14055.

25. Wang Z, Liu G, Sun J et al. Self-assembly of magnetite nanocrystals with amphiphilic polyethyleneimine: structures and applications in magnetic resonance imaging. J Nanosci Nanotech 2009;9:378–85.
26. Lin WJ, Chen TD, Liu C-W. Synthesis and characterization of lactobionic acid grafted pegylated chitosan and nanoparticle complex application. *Polymer* 2009;50:4166–74.

27. Thomas M, Klíbanov AM. Enhancing polyethylenimine’s delivery of plasmid DNA into mammalian cells. *Proc Natl Acad Sci USA* 2002;99:14640–5.

28. Sun S, Zeng H, Robinson DB et al. Monodisperse MFe2O4 (M = Fe, Co, Mn) nanoparticles. *J Am Chem Soc* 2004;126:273–9.

29. Yang Z, Cao J, He Y et al. Macro-/micro-environment-sensitive chemosensing and biological imaging. *Chem Soc Rev* 2014;43:4563–601.

30. Wang H, Zhao E, Lam JWY et al. AIE luminogens: emission brightened by aggregation. *Mater Today* 2015;18:365–77.

31. Hong Y, Lam JW, Tang BZ. Aggregation-induced emission: phenomenon, mechanism and applications. *Chem Commun* 2009;4332–53.

32. Lee N, Yoo D, Ling D et al. Iron oxide based nanoparticles for multimodal imaging and magnetoresponsive therapy. *Chem Rev* 2015;115:10637–89.

33. Na HB, Song IC, Hyeon T. Inorganic nanoparticles for MRI contrast agents. *Adv Mater* 2009;21:2133–48.