Mouse lymphatic endothelial cell targeted probes: anti-LYVE-1 antibody-based magnetic nanoparticles

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Purpose: To investigate the specific targeting property of lymphatic vessel endothelial hyaluronan receptor-1 binding polyethylene glycol-coated ultrasmall superparamagnetic iron oxide (LYVE-1-PEG-USPIO) nanoparticles to mouse lymphatic endothelial cells (MLECs).

Methods: A ligand specific target to lymphatic vessels was selected by immunohistochemical staining on the sections of a Lewis subcutaneous transplanted tumor. The z-average hydrodynamic diameter (HD), zeta potential, and the relaxivity of PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles were determined with a laser particle analyzer and magnetic resonance T2 spin echo sequence, respectively. Prussian blue staining and transmission electron microscopy (TEM) of nanoparticle labeled cells were performed to determine the nanoparticles’ binding form. Magnetic resonance imaging (MRI) was performed in vitro to evaluate the signal enhancement on the T2 spin echo sequence of the nanoparticle labeled cells. The iron content of the labeled cells after the Prussian blue staining and MRI scanning was determined by atomic absorption spectroscopy (AAS).

Results: The anti-LYVE-1 antibody was used as the specific ligand to synthesize the target probe to the MLECs. The mean z-average HDs of the LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles were 57.42 ± 0.31 nm and 47.91 ± 0.73 nm, respectively, and the mean zeta potentials of the LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles were 12.38 ± 4.87 mV and 2.57 ± 0.83 mV, respectively. The relaxivities of the LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles were 185.48 mM–1s–1 and 608.32 mM–1s–1. Cells binding nanoparticles were visualized as blue granules in the Prussian blue staining. The TEM results of the labeled cells showed the specific localization of nanoparticles. The AAS results of labeled cells after the Prussian blue staining and MRI scanning showed that the LYVE-1-PEG-USPIO nanoparticles had good binding selectivity for MLECs. MRI results indicated that the PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles could generate contrast on T2-weighted imaging, and the correlation between R2 and the iron content of the labeled cells was significantly positive.

Conclusion: This study demonstrated that LYVE-1-PEG-USPIO nanoparticles might potentially be used as an MRI contrast agent for targeting MLECs, and the magnetic properties of LYVE-1-PEG-USPIO nanoparticles were suitable for MRI.

Keywords: nanoparticles, lymphatic vessel endothelial hyaluronan receptor-1 (LYVE-1), ultrasmall superparamagnetic iron oxide (USPIO), mouse lymphatic endothelial cells (MLECs), magnetic resonance imaging (MRI)

Introduction

The lymphatic system is essential for the maintenance of tissue fluid homeostasis. With tumor growth, intratumoral pressure will gradually increase, mainly as the result of mechanical pressure generated from the proliferation of tumor cells and the perfusion...
pressure of intratumoral microvessels. Thus, intratumoral lymphatic vessels may become an important decompression pathway, enabling the outflow of interstitial fluid and tumor cells and their migration to other regions.\textsuperscript{11–13} Many studies have shown that there is a relationship between malignant tumor metastases and increased density of intratumoral lymphatic vessels.\textsuperscript{3–6} Most of these results come from the immunohistochemical staining of intratumoral lymphatic vessels in sections of tumor specimens, and the specific biomarkers used for lymphatic vessels are several proteins, which are expressed in lymphatic endothelial cells (LECs), such as the mucin-type transmembrane protein Podoplanin,\textsuperscript{7} lymphatic vessel endothelial hyaluronan receptor-1 (LYVE-1),\textsuperscript{8} homeobox transcription factor Prox-1,\textsuperscript{9} and vascular endothelial growth factor receptor-3 (VEGFR-3).\textsuperscript{10–12}

Although the in vitro method is the current gold standard for the study of intratumoral lymphatic vessels, the occurrence and development of lymphatic vessels cannot be dynamically observed. Meanwhile, theoretically, the contrast agents of computed tomography (CT) and magnetic resonance (MR) injected intravenously could enter the tumor stroma, and drain into the tumor lymphatic vessels, and finally flow back into the venous system, so that the intratumoral lymphatic vessels could be imaged. However, it is difficult to distinguish between the intratumoral blood vessels and intratumoral lymphatic vessels due to the simultaneous imaging of them. With the emergence and rapid development of molecular imaging, this could make it possible to image intratumoral lymphangiogenesis in vivo with specific targeted probes,\textsuperscript{13–15} and provide a direct method to evaluate the biological characteristics of intratumoral lymphatic vessels. However, this is on the condition that high sensitivity probes and high resolution imaging equipment are acquired.

Because of its high spatial resolution, noninvasiveness, and ability to provide anatomic and physiologic information simultaneously, MR imaging (MRI) is widely used in molecular imaging, such as MR in vitro cell imaging and MR in vivo imaging of small animals. Ultrasmall superparamagnetic iron oxide (USPIO) nanoparticles with higher sensitivity can improve the contrast enhancement effect by decreasing the $T_2^*$ and $T_2$*-weighted relaxation time of tissues.\textsuperscript{16} However, for most in vivo applications, nanoparticles need to overcome the nonspecific phagocytosis by mononuclear phagocyte system (MPS). One strategy to decrease nonspecific phagocytosis is to modify the nanoparticles with polyethylene glycol (PEG).\textsuperscript{17} PEG can also form steric barriers to the agglomeration that aid in particle dispersion. So, a specific antibody to lymphatic vessels should be chosen, which is then used to bind with polyethylene-coated USPIO (PEG-USPIO) nanoparticles.

Like many other markers used in molecular pathology, none of the LEC-associated molecule markers is entirely specific for LECs. Due to the transient expression in the nucleus, prox-1 is difficult to use for clinical applications.\textsuperscript{18–20} Although VEGFR-3 expresses in LECs, it lacks lymphatic vessel specificity in cancer because it also expresses in blood vessel epithelium.\textsuperscript{21} LYVE-1 is specially expressed in LECs, and does not exist in blood vessel epithelium, except in normal liver blood sinusoids, spleen endothelium, high endothelial venules, and activated tissue macrophages.\textsuperscript{22–25} Podoplanin is a specific marker of the LECs and is not expressed in blood vessel epithelium,\textsuperscript{26} but it is expressed in the epithelium of the choroid plexus,\textsuperscript{27} kidney podocytes,\textsuperscript{28} and the lung is a major site of podoplanin expression.\textsuperscript{29}

In this study, first, we selected a relatively specific ligand between the anti-LYVE-1 antibody and anti-podoplanin antibody by immunohistochemical staining on the sections of a Lewis subcutaneous transplanted tumor, then used it to synthesize the molecular targeted probe. The physical characteristics of the probe were analyzed, including particle size, zeta potential, and relaxivity. Prussian blue staining, transmission electron microscopy (TEM), and atomic absorption spectrometry (AAS) of the labeled cells were performed to determine whether mouse LECs (MLECs) could be specifically and efficiently identified by this probe.

**Material and methods**

**Establish Lewis tumor model**

To establish the Lewis tumor mouse model, $1.0 \times 10^6$ Lewis tumor cells (purchased from AiBiological Research, Shanghai, People’s Republic of China) in 100 µL of Roswell Park Memorial Institute (RPMI)-1640 medium were inoculated subcutaneously in the right armpit region of ten 6-week-old male BALB/c mice (provided by the Laboratory Animal Center of our university, the First Hospital of China Medical University). Our study was approved by the institutional animal care committee of our university.

**Immunohistochemistry**

When the tumor volume grew to about 1 cm$^3$, the tumor-bearing mice were executed with overdose anesthesia. The tumors were removed from the right armpit of the mice, fixed with 4% paraformaldehyde for 24 hours, paraffin-embedded, and serial 5 µm thicknesses were cut. Immunostaining with the anti-LYVE-1 antibody and anti-podoplanin antibody were described in a previous report.\textsuperscript{30} Sections were deparaffinized,
rehydrated, and pretreated with 3% H₂O₂ for 20 minutes to block endogenous peroxidase activity. The following primary antibodies were used: hamster monoclonal anti-mouse podoplanin antibody (1:200, ab11936; Abcam, Cambridge, UK), rat monoclonal anti-mouse LYVE-1 antibody (1:100, MAB2125; R&D SYSTEMS Inc, Minneapolis, MN, USA), biotinylated anti-hamster immunoglobulin G (IgG) (1:100, ab6782; Abcam), and biotinylated anti-rat IgG (1:100, ZB2307; ZSGB-BIO, Beijing, People’s Republic of China). The slides were incubated with the anti-LYVE-1 antibody and anti-podoplanin antibody overnight at 4°C, then treated with biotinylated secondary antibodies followed by incubation with hors eradish peroxidase-conjugated streptavidin at room temperature for 30 minutes. Diaminobenzidine (DAB) was used as the chromogen. The slides were counterstained with hematoxylin.

LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles

The PEG-coated USPIO nanoparticles (PEG-USPIO) and PEG-USPIO coupled with rat monoclonal anti-mouse LYVE-1 antibody (MAB2125; R&D SYSTEMS Inc) (LYVE-1-PEG-USPIO) nanoparticles were provided by The Institute of Chemistry, Chinese Academy of Sciences (Beijing, People’s Republic of China). For the LYVE-1-PEG-USPIO nanoparticles, the feed ratio of the anti-LYVE-1 antibody to the PEG-USPIO nanoparticles was 6:1.

Z-average hydrodynamic diameter (HD) and zeta potential measurement

The z-average HD and zeta potential of two types of nanoparticles were measured with a laser particle analyzer (Nano ZS90; Malvern Instruments, Malvern, UK), the two types of nanoparticles were diluted with double distilled water (pH 6.0) and put into a sample chamber; the temperature was 25°C ± 0.05°C.

Relaxivity of LYVE-1-PEG-USPIO and PEG-USPIO

LYVE-1-PEG-USPIO and PEG-USPIO agarose gel suspensions (1% agarose gel) were prepared with iron concentrations of 0, 0.0017, 0.0034, 0.0068, 0.0136, and 0.0272 mM in 0.25 mL Eppendorf tubes. For the MR measurements, these tubes were embedded in a tank filled with 1% agarose gel to avoid interference produced by macroscopic external magnetic field B0 inhomogeneities. MRI was performed by using a 3.0 T MRI scanner (Signa HDxt imager; GE Healthcare Bio-Sciences Corp, Piscataway, NJ, USA) equipped with a small animal coil provided by GE. Imaging was achieved by using the T₂ spin echo sequence. The parameters were as follows: repetition time (TR) = 4000 ms; echo time (TE) = 10, 20, 40, 60, 80, and 100 ms; field of view (FOV) = 8 × 8 cm²; matrix size = 256 × 256; slice thickness = 1 mm; number of excitation (NEX) = 3. Source images analysis was made on a GE ADW4.4 workstation. A T₂ map program was used to calculate the sample T₂ relaxation times, and based on the T₂-relaxation time, the relaxation rate (R₂) was calculated as the inverse of the T₂-relaxation time in seconds (1/T₂). Each Eppendorf tube’s axial section was measured at three regions of interest (ROIs) (3 × 4 pixels) in at least three layers.

Cell culture

MLECs (purchased from AiBiological Research) were the primary culture cell line (to keep the homology, MLECs were used). Colon 26 tumor cells (provided by the Kanazawa University School of Medicine, Kanazawa, Japan) were the cell line of mouse colon tumor that we used. MLECs were grown and maintained according to the vendor’s recommendation using Dulbecco’s modified eagle medium (DMEM; Life Technologies, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum (FBS; Life Technologies) and antibiotics (1% penicillin/streptomycin; Life Technologies). Colon 26 tumor cells were propagated in RPMI-1640 medium (Life Technologies) supplemented with 5% FBS and antibiotics. Cells were maintained under standard conditions of 37°C and 5% CO₂ and the passage numbers of all cells were between three and nine.

Immunofluorescence microscopy

In order to determine the expression of LYVE-1 receptor in MLECs and colon 26 cells, immunofluorescence was performed. MLECs and colon 26 tumor cells were grown to confluence on six-well plates (BD Biosciences, Franklin Lakes, NJ, USA), fixed with 4% paraformaldehyde for 20 minutes at room temperature followed by permeabilization with 0.3% TritonX-100 for 5 minutes. Cells were incubated with a blocking solution containing 5% bovine serum albumin-phosphate buffered saline with Tween for 1 hour at room temperature, and then treated with primary antibody (rat monoclonal anti-mouse LYVE-1 antibody, 1:20, MAB2125; R&D SYSTEMS Inc) and secondary antibody (goat anti-rat IgG/fluorescein isothiocyanate, 1:100, ZSGB-BIO). Nuclei were stained with DAPI (1023626001; Roche, South San Francisco, CA, USA). Specimens were examined via a Nikon TE2000-S inverted microscope (Nikon Corporation, Tokyo, Japan) equipped with a Spot-RT digital camera.
Prussian blue staining for labeled cells and corresponding AAS

To demonstrate the label efficiency of LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles, MLECs were plated at $1 \times 10^5$ cells/mL on each well of the six-well plates in 2 mL serum-free DMEM (Life Technologies), colon 26 cells were plated at $1 \times 10^5$ cells/mL on each well of the six-well plates in 2 mL serum-free RPMI-1640 medium (Life Technologies), cultured at 37°C with 5% CO$_2$ and 100% humid atmosphere for 16 hours, and then either LYVE-1-PEG-USPIO or PEG-USPIO nanoparticles were added at two different concentrations (25 and 100 µg Fe/mL) for 2 hours at 37°C with 5% CO$_2$ and 100% humid atmosphere, in 2 mL DMEM or RPMI-1640. Unlabeled MLECs and colon 26 cells were used as control groups. After 2 hours of incubation, the supernatant was removed, and the labeled and unlabeled MLECs and colon 26 cells were washed three times with phosphate-buffered solution (PBS) to remove nanoparticles that were free in solution or loosely adhered to the cell surfaces. Cells were fixed with 4% paraformaldehyde for 30 minutes and then washed three times with PBS. Prussian blue staining was performed by incubating fixed cells in a mixture of 10% potassium ferrocyanide and 10% hydrochloric acid (Shanghai Yuanye Biological Technology Co, Shanghai, People’s Republic of China) for 30 minutes, followed by washing with PBS, and counterstaining with eosin. Cells were photographed under a Nikon TE2000-S inverted microscope. Cells exhibiting blue particles were considered Prussian blue-positive. After being photographed, the labeled cells of each sample were harvested from the six-well plates separately by using a cell scraper and resuspended in 1 mL double distilled water for measuring the iron content of the labeled and unlabeled cells with AAS (Z2000, Hitachi Ltd, Tokyo, Japan). The experiment was performed three times.

**TEM**

MLECs and colon 26 cells were detached from the cell culture flasks using 0.05% trypsin (Life Technologies), and adjusted to a concentration of $1 \times 10^5$ cells/mL in growth media. Then 2 mL of the cell suspension was plated on each well of the six-well plates at 37°C with 5% CO$_2$ and 100% humid atmosphere for 16 hours. After incubating with two kinds of nanoparticles at 100 µg Fe/mL for 2 hours at 37°C with 5% CO$_2$ and 100% humid atmosphere in 2 mL DMEM or RPMI-1640, the MLECs and colon 26 cells were washed three times with PBS. The labeled MLECs and colon 26 cells were detached from the six-well plates using 0.05% trypsin, the same kind of cells within the six wells were collected in one group, and centrifuged at 1000 rpm for 10 minutes. After discarding the supernatants, cell pellets were fixed in 2.5% phosphate buffered glutaraldehyde for 4 hours at 4°C, and then washed three times in PBS. The samples were dehydrated by a graded series of ethanol, stained in 1% uranyl acetate and embedded in epon. Ultrathin sections (nm) were contrasted with 0.3% lead citrate and then examined and photographed at 80 kV by TEM (H-7650; Hitachi Ltd). In order to make sure of the morphology of the nanoparticles, the LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles suspension were diluted with PBS to a concentration of 10 µg Fe/mL, and was directly deposited onto a carbon-coated copper grid and air-dried at room temperature and examined and photographed by TEM.

**MRI of labeled cells and corresponding AAS**

To demonstrate the MRI detection efficiency of LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles, the MLECs and colon 26 cells were incubated separately in 24-well plates (BD Biosciences) at an amount of $1 \times 10^5$ cells/mL each well for 16 hours at 37°C in a 5% CO$_2$ and 100% humid atmosphere, in 1 mL DMEM and RPMI-1640 respectively. Then, the growth media were discarded, serum-free DMEM or serum-free RPMI-1640 medium containing 6.25, 12.5, 25, 50, 100, or 200 µg Fe/mL of either LYVE-1-PEG-USPIO or PEG-USPIO nanoparticles was added into each well in 1 mL for 2 hours at 37°C in a 5% CO$_2$ and 100% humid atmosphere. Unlabeled MLECs and colon 26 cells were used as control groups. Labeled and unlabeled MLECs and colon 26 cells were washed three times with PBS, and detached using 0.05% trypsin. Labeled cells ($1 \times 10^5$ cells/tube) cultured within various concentrations and unlabeled cells were centrifuged at 800 rpm for 5 minutes, then the supernatant was discarded and the cells were resuspended with 0.25 mL 1% agarose gel in 0.25 mL Eppendorf tubes. In order to avoid interference produced by macroscopic external magnetic field B0 inhomogeneities, these tubes were embedded in a tank filled with 1% agarose gel. MRI was immediately performed by using a 3.0 T MRI scanner equipped with a small animal coil. Imaging was achieved by using the $T_2$ spin echo sequence. The parameters were as follows: TR = 4000 ms; TE = 10, 20, 40, 60, 80, and 100 ms; FOV = 8 × 8 cm$^2$; matrix size = 256 × 256; slice thickness = 1 mm; NEX = 3. Source images analysis was made on a GE ADW4.4 workstation. A $T_2$ map program was used to calculate the sample $T_2$.
relaxation times. The $R_2$-relaxation time in seconds ($1/T_2$). Each Eppendorf tube’s axial section was measured at three ROIs ($3 \times 4$ pixels) in at least three layers. When the MR test was finished, the iron content of labeled and unlabeled cells of each sample was measured with AAS.

**AAS quantification of labeled cells**
To determine the iron content of labeled and unlabeled cells, AAS measurement was performed. The experiment was performed three times. The average value of cell binding iron content was taken.

**Statistical analysis**
Statistical analysis was performed by using the SPSS 16.0 software (IBM Corporation, Armonk, NY, USA). All data were presented as mean ± standard deviation (SD). Statistical significance was assessed by an unpaired Student’s $t$-test for single comparisons or by one-way analysis of variance (ANOVA) for multiple comparisons. Pearson correlation analysis was run on comparisons of iron content and $R_2$. A $P$-value of less than 0.05 was considered statistically significant.

**Results**

**Ligand selected**
The immunohistochemical staining results showed that the anti-podoplanin antibody expressed not only in the lymphatic vessels, but also in a large number of other cells within the Lewis tumor. However, anti-LYVE-1 antibody was mainly expressed in the lymphatic vessels (Figure 1A and B). Therefore, we concluded that the anti-LYVE-1 antibody was more specific than the anti-podoplanin antibody in the identification of lymphatic vessels within the mouse Lewis subcutaneous transplanted tumor, so we selected anti-LYVE-1 antibody as the specific ligand.

**Figure 1** Immunohistochemical staining of Lewis tumor.
*Notes:* (A) LYVE-1-positive staining and (B) podoplanin-positive staining were seen in the Lewis tumor. Anti-podoplanin antibody could also stain a large number of other cells besides the lymphatic vessels (arrows), but anti-LYVE-1 antibody identified mainly only lymphatic vessels (arrows). Bar = 100 μm.

*Abbreviation:* LYVE-1, lymphatic vessel endothelial hyaluronan receptor-1.

**Nanoparticles characterization**
The results showed that the zeta potentials of PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles were $2.57 \pm 0.83$ mV and $12.38 \pm 4.87$ mV, respectively, and the z-average HDs were $47.91 \pm 0.73$ nm and $57.42 \pm 0.31$ nm, respectively. The $R_2$ was plotted on the y-axis versus iron concentration (x-axis). The relaxivity of the nanoparticles was obtained by the slope of the linear regression of $R_2$ with iron concentration. The relaxivity of the PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles was 608.32 mM$^{-1}$s$^{-1}$ and 185.48 mM$^{-1}$s$^{-1}$, respectively (Figure 2A and B).

**Immunofluorescence microscopy**
Immunofluorescence results showed that LYVE-1 receptors were expressed in MLECs, but not in colon 26 cells (Figure 3A–D).

**Prussian blue staining and corresponding AAS**
Prussian blue staining was used to determine the existence of the nanoparticles in cells, and the cells binding nanoparticles were visualized as blue granules (Figure 4A–J). The number of blue granules increased with the increasing iron concentration, especially in the LYVE-1-PEG-USPIO labeled MLECs. To assess specific binding of LYVE-1-PEG-USPIO nanoparticles to MLECs, the iron content of the two cell lines was detected with AAS after 2 hours of incubation with LYVE-1-PEG-USPIO and PEG-USPIO. When the incubation concentrations of LYVE-1-PEG-USPIO and PEG-USPIO were 25 μg Fe/mL, we found the highest accumulation of LYVE-1-PEG-USPIO in the MLECs with the mean iron content of $48.4 \pm 8.5$ pg Fe/cell. The mean iron contents of LYVE-1-PEG-USPIO labeled colon 26 cells, PEG-USPIO labeled MLECs, and PEG-USPIO labeled colon 26 cells were $19.92 \pm 3.58$ pg Fe/cell, $17.56 \pm 5.76$ pg Fe/cell, and $15.61 \pm 7.23$ pg Fe/cell, respectively. When increasing the incubation concentration of LYVE-1-PEG-USPIO and PEG-USPIO to 100 μg Fe/mL, the mean iron content of LYVE-1-PEG-USPIO labeled MLECs was still the highest ($219.22 \pm 14.89$ pg Fe/cell), and approximate 4.6 times greater than the iron content of LYVE-1-PEG-USPIO labeled MLECs at the iron incubation concentration of 25 μg Fe/mL. Although the colon 26 cells showed an intense binding of LYVE-1-PEG-USPIO, they only reached a mean iron contents of $64.41 \pm 6.39$ pg Fe/cell. The mean iron contents of PEG-USPIO labeled MLECs and PEG-USPIO labeled colon 26 cells were $38.92 \pm 7.07$ pg Fe/cell and $35.03 \pm 9.89$ pg Fe/cell respectively (Figure 5).
TEM of labeled cells

TEM of the two cell lines after 2 hours of incubation with LYVE-1-PEG-USPIO (100 µg Fe/mL) and PEG-USPIO (100 µg Fe/mL) was used to study the subcellular localization of the two types of nanoparticles. Images of the LYVE-1-PEG-USPIO labeled MLECs revealed that the nanoparticles were mainly located in the cytoplasm endosomes, and a few nanoparticles were also attached to the cell membrane (Figure 6A). However, the PEG-USPIO nanoparticles were mainly attached to the MLEC membrane (Figure 6B). In the colon 26 cells, the LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles were mainly attached to the cell membrane (Figure 6C and D), but a few LYVE-1-PEG-USPIO nanoparticles were also found in the cytoplasm endosomes.

MRI and corresponding AAS in vitro

Tubes containing MLECs and colon 26 cells, which were incubated with serial concentrations of LYVE-1-PEG-USPIO and PEG-USPIO, were scanned with an MRI T2 spin echo sequence. The signal intensity of T2-weighted images decreased with the increasing iron concentration (Figure 7A). The MRI results showed a concentration-dependent gradient for the four kinds of labeled cells (Figure 7B). As the incubation concentrations of LYVE-1-PEG-USPIO and PEG-USPIO increased, the R2 of the labeled cells also increased. The R2 of tubes containing the same cell line incubated with LYVE-1-PEG-USPIO were higher than those incubated with PEG-USPIO at the same concentration. The R2 of tubes containing LYVE-1-PEG-USPIO labeled MLECs were higher than those containing LYVE-1-PEG-USPIO labeled colon 26 cells at the same concentration. The R2 of LYVE-1-PEG-USPIO labeled MLECs was significantly higher than other labeled cells at iron concentration of 25 to 200 µg Fe/mL (P < 0.05). There was no statistical difference in the R2 between the unlabeled MLECs and unlabeled colon 26 cell groups (P > 0.05). AAS results revealed a dose-dependent increase of nanoparticle labeled cells (Figure 7C). With increasing iron incubation concentration, the iron content of labeled cells increased. LYVE-1-PEG-USPIO labeled MLECs showed a significantly higher iron content than the other three kinds of labeled cells at the same incubation concentration. There was no statistical difference in the iron content between the unlabeled MLECs and unlabeled colon 26 cell groups (P > 0.05). The corresponding data were summarized in Table 1. Comparison of the R2 and iron content of labeled cells revealed significantly positive correlations. The Pearson correlation coefficient, r, for LYVE-1-PEG-USPIO labeled MLECs was r = 0.995 (P < 0.01), for PEG-USPIO labeled
MLECs was $r = 0.976$ ($P < 0.01$), for LYVE-1-PEG-USPIO labeled colon 26 cells was $r = 0.997$ ($P < 0.01$), and for PEG-USPIO labeled colon 26 cells was $r = 0.977$ ($P < 0.01$) (Figure 7D).

**Discussion**

In this study, an MRI probe was conducted with the anti-LYVE-1 monoclonal antibody, determined by the immunohistochemical staining of Lewis tumor specimens, as the specific targeting ligand and the PEG-coated USPIO nanoparticles as the signal ligand; the LYVE-1 receptor in LECs could be targeted and imaged by MRI. After evaluating the physical properties of this MRI probe, a series of cell experiments in vitro were performed to assess the specificity and effectiveness of the LYVE-1-PEG-USPIO nanoparticles.

To evaluate the $T_2$ enhancing capabilities of PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles, agarose gel solutions (1% agarose gel) of the PEG-USPIO and LYVE-1-PEG-USPIO nanoparticles at different iron concentrations were imaged by an atomic absorption spectroscopy (AAS). The results showed that colon 26 cells with LYVE-1-PEG-USPIO had a higher iron content than PEG-USPIO at both 25 and 100 µg Fe/mL concentrations. The specific binding of the antibody to the receptor was confirmed by the increased iron content in the MLEC control groups compared to the colon 26 cells control groups.

**Abbreviations:** AAS, atomic absorption spectroscopy; LYVE-1-PEG-USPIO, lymphatic vessel endothelial hyaluronan receptor-1 binding polyethylene glycol-coated ultrasmall superparamagnetic iron oxide; MLECs, mouse lymphatic endothelial cells; PEG-USPIO, polyethylene glycol-coated ultrasmall superparamagnetic iron oxide.
MLECs, LYVE-1-PEG-USPIO labeled colon 26 cells, and of labeled cells and control group cells (MLECs was significantly higher than the other three kinds Fe/mL), the iron content of the LYVE-1-PEG-USPIO labeled kinds of cells also increased. At a low concentration (25 increasing, the amount of nanoparticles bound to the two of LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles Prussian blue staining, showed that with the concentration results of the same samples, which had been examined with AAS. AAS of cells had different binding capacities to the two types of nanoparticles, and this was further verified by AAS. AAS could generate contrast on T₁-weighted imaging, which had been reported that particles with radii 50 nm exhibited significantly greater uptake compared with particles >50 nm. Theoretically, without considering the effect of LYVE-1 receptor-mediated endocytosis and the nanoparticles’ surface charge, the number of cells binding PEG-USPIO nanoparticles would be much greater than the number of cells binding LYVE-1-PEG-USPIO nanoparticles at the same incubation time and at the same iron incubation concentration. However, for the two cell lines, the higher cell binding nanoparticle was LYVE-1-PEG-USPIO, not PEG-USPIO; this indicated that the size of the nanoparticles was not the major factor for the binding capacity of cells to the two types of nanoparticles.

PEG-USPIO labeled colon 26 cells. When increasing the incubation concentration of LYVE-1-PEG-USPIO and PEG-USPIO to 100 μg Fe/mL, the iron content of the LYVE-1-PEG-USPIO labeled MLECs which were harvested after the Prussian blue staining, increased most significantly, approximately 4.6 times greater than the iron content of the LYVE-1-PEG-USPIO labeled MLECs at the iron incubation concentration of 25 μg Fe/mL. Although the iron content of the other three kinds of labeled cells also increased when increasing iron incubation concentration, they were far below that of the LYVE-1-PEG-USPIO labeled MLECs.

MRI detection of the labeled cells showed the same effect as the results of the Prussian blue staining. The signal intensity of the LYVE-1-PEG-USPIO labeled MLECs decayed most rapidly when the iron concentration increased, and AAS results of the same samples, which had been examined by MRI also showed the highest iron content of LYVE-1-PEG-USPIO labeled MLECs (Figure 7). The iron content of the LYVE-1-PEG-USPIO labeled MLECs was approximately 2.3 times greater than the iron content of the LYVE-1-PEG-USPIO labeled colon 26 cells at the iron incubation concentration of 25 μg Fe/mL, and approximately 4.4 times greater at the iron incubation concentration of 200 μg Fe/mL.

As we know, the binding capacity of cells to nanoparticles might be attributed to receptor-mediated endocytosis, nanoparticles’ surface charge, and size. To analyze the different effects caused by the three factors, we chose colon 26 cells, which lack LYVE-1 receptor as confirmed by the immunofluorescence study, as the negative control under the uniform conditions of temperature, incubation concentration, and incubation time.

The mean hydrodynamic size of the LYVE-1-PEG-USPIO nanoparticles was 57.42 ± 0.31 nm, which was bigger than that of the PEG-USPIO nanoparticles (47.91 ± 0.73 nm). It had been reported that particles with radii <50 nm exhibited significantly greater uptake compared with particles >50 nm. Theoretically, without considering the effect of LYVE-1 receptor-mediated endocytosis and the nanoparticles’ surface charge, the number of cells binding PEG-USPIO nanoparticles would be much greater than the number of cells binding LYVE-1-PEG-USPIO nanoparticles at the same incubation time and at the same iron incubation concentration. However, for the two cell lines, the higher cell binding nanoparticle was LYVE-1-PEG-USPIO, not PEG-USPIO; this indicated that the size of the nanoparticles was not the major factor for the binding capacity of cells to the two types of nanoparticles.

The mean zeta potential of the two types of nanoparticles measured with a laser particle analyzer was 2.57 ± 0.83 mV
for PEG-USPIO and 12.38 ± 4.87 mV for LYVE-1-PEG-USPIO. Studies had shown that the cell phagocytosis rate of surface-positively charged nanoparticles was faster than that of neutral or surface-negatively charged nanoparticles, because nanoparticles with positive surface charges could more easily adhere to the surface-negatively charged cell membrane by electrostatic interactions. The AAS results showed that the iron content of LYVE-1-PEG-USPIO labeled colon 26 cells was approximately 1.3 times greater than that of PEG-USPIO labeled colon 26 cells at the iron incubation concentration of 25 μg Fe/mL, and the differences gradually increased as the iron incubation concentration increased. The iron content of LYVE-1-PEG-USPIO labeled colon 26 cells was approximately 2.1 times greater than that of PEG-USPIO labeled colon 26 cells at the incubation concentration of 200 μg Fe/mL. Thus, we concluded that the different binding capacities of colon 26 cells to LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles were mainly caused by the different surface charges of the two types of nanoparticles. However, for the binding capacities of MLECs to the two types of nanoparticles, the iron content of LYVE-1-PEG-USPIO labeled MLECs was approximately 2.6 times greater than that of PEG-USPIO labeled MLECs at the iron incubation concentration of 25 μg Fe/mL, and increased to 8.5 times greater at the iron incubation concentration of 200 μg Fe/mL. Furthermore, the iron content of LYVE-1-PEG-USPIO labeled MLECs was approximately 2.3 times greater than the iron content of the LYVE-1-PEG-USPIO labeled colon 26 cells at the iron incubation concentration of 25 μg Fe/mL, and approximately 4.4 times greater at the iron incubation concentration of 200 μg Fe/mL. These data indicated that the MLEC binding capacity to LYVE-1-PEG-USPIO nanoparticles was considerably higher than that of the colon 26 cells. This should be attributed to not only the different surface charges of the two types of nanoparticles, but also...
Table 1  Iron content and R₂ of MLECs and colon 26 cells after incubation with LYVE-1-PEG-USPIO and PEG-USPIO at various concentrations for 2 hours

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<thead>
<tr>
<th>Incubation concentration (µg Fe/mL)</th>
<th>Nanoparticle labeled cells</th>
<th>AAS iron content (pg Fe/cell)</th>
<th>MR imaging R₂ (s⁻¹)</th>
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</thead>
<tbody>
<tr>
<td>0</td>
<td>MLEC + LYVE-1-PEG-USPIO</td>
<td>0.93 ± 0.23</td>
<td>2.12 ± 0.21</td>
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<tr>
<td></td>
<td>MLEC + PEG-USPIO</td>
<td>0.95 ± 0.32</td>
<td>2.32 ± 0.19</td>
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<td>Colon 26 + LYVE-1-PEG-USPIO</td>
<td>0.87 ± 0.39</td>
<td>2.24 ± 0.23</td>
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<td>Colon 26 + PEG-USPIO</td>
<td>0.88 ± 0.50</td>
<td>2.15 ± 0.25</td>
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<td>6.25</td>
<td>MLEC + LYVE-1-PEG-USPIO</td>
<td>9.32 ± 1.36</td>
<td>2.92 ± 0.23</td>
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<tr>
<td></td>
<td>MLEC + PEG-USPIO</td>
<td>8.16 ± 1.12</td>
<td>2.63 ± 0.21</td>
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<tr>
<td></td>
<td>Colon 26 + LYVE-1-PEG-USPIO</td>
<td>7.64 ± 1.21</td>
<td>2.67 ± 0.34</td>
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<tr>
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<td>Colon 26 + PEG-USPIO</td>
<td>7.44 ± 1.15</td>
<td>2.43 ± 0.28</td>
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<td>12.5</td>
<td>MLEC + LYVE-1-PEG-USPIO</td>
<td>17.52 ± 2.12</td>
<td>3.90 ± 0.57</td>
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<td>MLEC + PEG-USPIO</td>
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<td>2.71 ± 0.46</td>
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<td>Colon 26 + LYVE-1-PEG-USPIO</td>
<td>10.23 ± 1.42</td>
<td>3.03 ± 0.59</td>
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<td>Colon 26 + PEG-USPIO</td>
<td>8.64 ± 1.28</td>
<td>2.54 ± 0.35</td>
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<td>MLEC + LYVE-1-PEG-USPIO</td>
<td>46.04 ± 3.12</td>
<td>8.52 ± 0.34</td>
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<td>MLEC + PEG-USPIO</td>
<td>18.16 ± 2.04</td>
<td>3.71 ± 0.24</td>
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<td>Colon 26 + LYVE-1-PEG-USPIO</td>
<td>19.84 ± 2.16</td>
<td>4.85 ± 0.29</td>
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<td>2.97 ± 0.38</td>
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<td>7.26 ± 0.89</td>
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<td>12.36 ± 1.21</td>
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<tr>
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<td>Colon 26 + PEG-USPIO</td>
<td>31.64 ± 2.92</td>
<td>6.32 ± 0.98</td>
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<tr>
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<td>62.50 ± 3.06</td>
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<tr>
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<td>MLEC + PEG-USPIO</td>
<td>53.61 ± 4.32</td>
<td>12.11 ± 2.42</td>
</tr>
<tr>
<td></td>
<td>Colon 26 + LYVE-1-PEG-USPIO</td>
<td>101.92 ± 5.40</td>
<td>22.33 ± 3.38</td>
</tr>
<tr>
<td></td>
<td>Colon 26 + PEG-USPIO</td>
<td>47.96 ± 3.44</td>
<td>10.35 ± 1.56</td>
</tr>
</tbody>
</table>

Note: Results expressed as mean ± standard deviations (n = 3).

Abbreviations: AAS, atomic absorption spectrometry; LYVE-1-PEG-USPIO, lymphatic vessel endothelial hyaluronan receptor-1 binding polyethylene glycol-coated ultrasmall superparamagnetic iron oxide; MLECs, mouse lymphatic endothelial cells; MR, magnetic resonance; PEG-USPIO, polyethylene glycol-coated ultrasmall superparamagnetic iron oxide.

receptor-mediated endocytosis, and the latter should be considered as the major factor.

In addition, the diameters of the two cell lines used in this study were different; MLEC diameters were bigger than colon 26 cell diameters, so the surface area of a single MLEC was bigger than that of a single colon 26 cell. In the condition of the same number of the two kinds of cells, the total surface area of the MLECs was much bigger than that of the colon 26 cells. The results of the AAS analysis showed that there was no statistical difference between the iron content of the PEG-USPIO labeled MLECs and PEG-USPIO labeled colon 26 cells (P > 0.05).

TEM results indicated that the localizations of LYVE-1-PEG-USPIO and PEG-USPIO nanoparticles in MLECs and colon 26 cells were different at the iron incubation concentration of 100 µg Fe/mL for 2 hours. For colon 26 cells, the LYVE-1-PEG-USPIO nanoparticles were mainly attached to the cell membrane, and a few were located in colon 26 cell cytoplasmic endosomes. The PEG-USPIO nanoparticles were almost all attached to the membranes of MLECs and colon 26 cells (Figure 6). Researchers found that the uptake of USPIO by cells was dose and time dependent.38,39 Sun et al reported that only a few cells showed the uptake of USPIO (0.3 µmol Fe/mL) after incubation for 20 minutes, and the uptake of USPIO gradually increased during the next 12 hours.38 In our study, TEM of labeled cells was performed at the iron incubation concentration of 100 µg Fe/mL for 2 hours. The PEG-USPIO nanoparticles were almost all attached to the cell membrane; we thought the PEG with which the USPIO nanoparticles were modified could reduce the nonspecific phagocytosis, and another reason might be the insufficient incubation time. We thought that with
prolonged incubation time and increased iron incubation concentration, the iron content of the labeled cells could have certain changes, which need further study. However, the LYVE-1-PEG-USPIO nanoparticles were mainly located in the MLECs cytoplasmic endosomes, with a few attached to the cell membranes. This should be attributed to receptor-mediated endocytosis. Researchers pointed out that, there were at least three types of endocytosis for nanoparticles: clathrin-mediated endocytosis, caveolae-mediated endocytosis, and clathrin-and caveolae-independent endocytosis. Prevo et al showed that LYVE-1 could mediate the endocytosis of hyaluronic acid by means of receptor-mediated endocytosis, and they inferred that LYVE-1 mediated the uptake of hyaluronic acid via caveolae-mediated endocytosis. Therefore, it could be concluded that the LYVE-1-PEG-USPIO nanoparticles located in the MLEC cytoplasmic endosomes were mainly caused by receptor-mediated endocytosis.

We also found that the correlation of the $R^2$ and the iron content of labeled cells as measured by AAS was significantly positive ($P < 0.05$). There was no significant difference among the $R^2$ of the four kinds of labeled cells when the iron incubation concentration of nanoparticles was lower than 25 µg Fe/mL ($P > 0.05$). However, when the iron incubation concentration was equal to or greater than 25 µg Fe/mL, the differences were significant ($P < 0.05$), and the differences were more apparent with the increasing of the iron incubation concentration. Therefore, the iron concentration ($\geq 25$ µg Fe/mL) of the nanoparticles could be used in an in vivo test.

Our study had several limitations. First, due to the different LEC sources and species, the expression of LYVE-1 receptors in LECs might be different. Our study could not demonstrate that the anti-LYVE-1 antibody-based probe could be applied to other species, especially to human LECs, so the feasibility of the application of the probe in other sources and species of LECs needs further research. Second, Nakao et al have reported that not all LECs express LYVE-1 and the expression of LYVE-1 would be downregulated with cells or species aging. Although the cultured MLECs were controlled in three to nine generations in our study, as the MLEC generation increases, there might be some changes in the expression of LYVE-1. Our in vitro study demonstrated that the LYVE-1-PEG-USPIO nanoparticles could effectively target to MLEC; however, we did not perform an in-depth study about the influence of different cultured MLEC generations on the expression of LYVE-1. Third, it has been indicated that, until now, none of the LEC-associated molecular markers is entirely specific for lymphatic endothelium; LYVE-1 is also expressed in liver blood sinusoids, spleen endothelium, and activated tissue macrophages. When the LYVE-1-PEG-USPIO is applied in vivo, it is hard to avoid being captured by these cells, which may disturb the probe to the real targets. How to solve these problems requires further research.

**Conclusion**

This study indicated that the anti-mouse LYVE-1 antibody-based probe had good binding affinity to LYVE-1 expressed in MLECs. The MRI results indicated that the LYVE-1-PEG-USPIO nanoparticles could generate contrast on T2-weighted imaging, and the correlation between $R^2$ and the iron content of labeled cells was significantly positive. The LYVE-1-PEG-USPIO nanoparticle was a promising agent for MRI of MLECs in vitro. Whether or not this probe could be used in the complex environment of the living organism needs further research.

**Acknowledgments**

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


