Accurate sequential detection of primary tumor and metastatic lymphatics using a temperature-induced phase transition nanoparticulate system

Abstract: Primary tumor and tumor-associated metastatic lymphatics have emerged as new targets for anticancer therapy, given that these are difficult to treat using traditional chemotherapy. In this study, docetaxel-loaded Pluronic nanoparticles with Flamma™ (FPR-675, fluorescence molecular imaging dye; DTX/FPR-675 Pluronic NPs) were prepared using a temperature-induced phase transition for accurate detection of metastatic lymphatics. Significant accumulation was seen at the primary tumor and in metastatic lymph nodes within a short time. Particle size, maximum drug loading capacity, and drug encapsulation efficiency of the docetaxel-loaded Pluronic NPs were approximately 10.34±2.82 nm, 3.84 wt%, and 94±2.67 wt%, respectively. Lymphatic tracking after local and systemic delivery showed that DTX/FPR-675 Pluronic NPs were more potent in tumor-bearing mice than in normal mice, and excised mouse lymphatics showed stronger near-infrared fluorescence intensity on the tumor-bearing side than on the non-tumor-bearing side at 60 minutes post-injection. In vivo cytotoxicity and efficacy data for the NPs demonstrated that the systemically administered NPs caused little tissue damage and had minimal side effects in terms of slow renal excretion and prolonged circulation in tumor-bearing mice, and rapid renal excretion in non-tumor-bearing mice using an in vivo real-time near-infrared fluorescence imaging system. These results clearly indicate that docetaxel-loaded Pluronic NPs could provide a strategy to achieve effective cancer therapy by simultaneous delivery to primary tumors, tumor lymphatics, and tumor-associated metastatic lymphatics.

Keywords: metastatic lymphatics, primary tumor targeting, lymphatic tracking, temperature-induced phase transition, Pluronic nanoparticles

Introduction

Following on from the understanding that the first lymph nodes receive metastatic cancer cells via direct lymphatic drainage from the primary tumor, the sentinel lymph node (SLN) is recognized as a crucial element in cancer staging that limits the morbidity of treatment with excellent oncologic results.1–4 Moreover, as the clinical standard of care for cancer staging, tumor involvement of the SLN is identified using peritumoral injections of radiolabeled agents or imaging dyes (inorganic or organic), followed by SLN biopsy, which is an invasive surgical procedure.5–8 Therefore, there is a significant need to establish a noninvasive method for more accurate assessment of lymph node involvement with improved sensitivity and target specificity.

New macromolecular agents, including nanoparticles (NPs), have recently been used to image the lymphatics and SLN by magnetic resonance and optical techniques.9–11 Magnetic resonance imaging with ultrasmall superparamagnetic iron oxide NPs, which accumulate in the lymph nodes due to their prolonged circulation in the blood, has been used to reveal changes in the status of the SLN...
Based on cellular uptake behavior in various cell lines, an in vivo lymphatic drainage and retention of NPs in the lymph nodes was confirmed because the particle size range was optimal for the lymphatic uptake of injected NPs appears to be NP size, ie, NPs larger than 100 nm in diameter primarily remain at the injection site whereas those 10–80 nm in diameter are taken up well by the lymphatics.

In our previous reports, the formation of a nanocarrier with a core-shell structure was demonstrated through temperature-induced phase transitions in the melt state of a polyethylene glycol (PEG)/paclitaxel/Pluronic mixture. Liquid PEG (molecular weight 400) was used as a paclitaxel solubilizer, and the polymer that encapsulated the paclitaxel was composed of Pluronic F-68. For the preparation of docetaxel (DTX)-loaded Pluronic NPs, a soybean oil/Tween 80 mixture was used as the solubilizer instead of PEG in a temperature-induced phase transition. The Pluronic NPs delivering paclitaxel or DTX showed improved antitumor efficacy compared with a Cremophor® EL-based paclitaxel formulation and a Tween 80/ethanol-based DTX formulation (Taxotere®, Sanofi-aventis, Paris, France). Interestingly, a significant decrease in diameter was observed in the Pluronic NPs formed with soybean oil/Tween 80 as the DTX solubilizer. The diameter of the optimized NPs changed as a function of the soybean oil/Tween 80 ratio, and their size distribution ranged from 10 nm to 20 nm.

The lymphatic system is not easily accessible using conventional intravenous injections or infusions of chemotherapeutic agents, and this limits the amount of drug that can access the lymphatic tissues, including a metastatic lymph node. It has been reported that the primary controlling factor for the lymphatic uptake of injected NPs appears to be NP size, ie, NPs larger than 100 nm in diameter primarily remain at the injection site whereas those 10–80 nm in diameter are taken up well by the lymphatics.

In this study, the suitability of DTX-loaded Pluronic nanoparticles with Flamma™ (FPR-675, a molecular imaging dye [DTX/FPR-675 Pluronic NPs]) as an efficient imaging modality and drug carrier for lymphatic cancer metastasis was confirmed because the particle size range was optimal for lymphatic drainage and retention of NPs in the lymph nodes. Based on cellular uptake behavior in various cell lines, an in vivo biodistribution study was conducted in order to confirm the increased tumor targetability and lymphatic tracking of the DTX/FPR-675 Pluronic NPs in tumor-bearing mice.

Materials and methods

Materials

Pluronic F-68 [poly(ethylene oxide)-poly (propylene oxide)-poly (ethylene oxide)] triblock copolymer (molecular weight 8,350; (EO)75(PO)25(EO)75) was obtained from BASF Company Ltd (Seoul, Republic of Korea). DTX (in anhydrous form) was purchased from Parling Pharma Tech Co Ltd (Shanghai, People’s Republic of China). Polyoxymethylene sorbitan monoooleate (Tween 80), soybean oil, and thiazoyl blue tetrazolium bromide (MTT) were purchased from Sigma (St Louis, MO, USA). FPR-675 dye was purchased from Bioacts (Incheon, Republic of Korea). Bovine aortic endothelial cells, murine macrophages (Raw264.7), and squamous cell carcinoma (SCC-7) cells were purchased from the American Type Culture Collection (Rockville, MD, USA). Roswell Park Memorial Institute 1640 medium, Dulbecco’s Modified Eagle’s Medium (low glucose), fetal bovine serum, and Dulbecco’s phosphate-buffered saline were purchased from WelGENE Inc (Daegu, Republic of Korea). All experiments with live animals were performed in compliance with the relevant laws and institutional guidelines of the Korea Institute of Science and Technology, and institutional committees approved the experiments. BALB/c nude mice (aged 5 weeks, 20–25 g, male) and athymic nude mice (aged 5.5 weeks, 20–25 g, male) were purchased from Orient Bio Inc (Gyeonggi-do, Republic of Korea). All other chemicals were analytical grade and used without further purification.

Methods

Preparation of the DTX/FPR-675 Pluronic NPs

The DTX/FPR-675 Pluronic NPs were prepared by temperature-induced phase transition. In order to form a drug-loaded phase at room temperature, 300 mg of liquid soybean oil/Tween 80, 1 mg of FPR-675 (loading amount 0.0033 wt%), and 20 mg of DTX (loading amount 6.25 wt%) were mixed, and that mixture was subsequently mixed with 200 mg of Pluronic F-68. As the temperature was increased to 60°C, the mixture melted into a liquid phase. Equilibrium was maintained at 60°C for 10 minutes with stirring, and the liquid mixture was then sufficiently cooled to 0°C for 10 minutes to induce the phase transition. For in vitro and in vivo NIRF imaging, the FPR-675 dye was mixed with a soybean oil/Tween 80/DTX mixture to observe lymphatic tracking in various organs
and tumor distributions of the DTX/FPR-675 Pluronic NPs in SCC-7 tumor-bearing mice.

To remove the unloaded DTX, the NPs prepared by temperature-induced phase transition were mixed with distilled deionized water using a shaker under mild conditions at room temperature for 30 minutes. The dispersed NPs were then centrifuged at 3,000 rpm for 10 minutes followed by lyophilization or stored at 0°C for other experiments. To evaluate the physicochemical characteristics of the DTX/FPR-675 Pluronic NPs, a predetermined amount of distilled deionized water or phosphate-buffered saline (pH 7.4) was added to the dried sample and the NPs were freely dispersed in mild mixing conditions in each solution without specific mechanical assistance such as sonication.

**Particle size and morphology of the DTX/FPR-675 Pluronic NPs**

The average diameter, size distribution, and zeta potential of the NPs (1 mg/mL of NPs dispersed in phosphate-buffered saline) were measured by dynamic light scattering (Zetasizer Nano ZS90, Malvern Instruments Ltd, Malvern, UK) at 632.8 nm and 25°C. Transmission electron micrographs were also taken to observe the morphology of the DTX/FPR-675 Pluronic NPs. For the transmission electron microscopic measurements, the freeze-dried NPs were dispersed in distilled deionized water to obtain a solution of 0.1 wt%. In order to prepare a sample, each solution was dropped on a 300 mesh copper grid coated with carbon and then dried at 25°C in a vacuum oven for 24 hours. Subsequently, the sample was negatively stained with 2 wt% uranyl acetate solution and dried. The samples were examined using an H-7600 microscope (Hitachi, Tokyo, Japan) operated at 100 kV.

**Release behavior of DTX from the DTX/FPR-675 Pluronic NPs**

In order to measure the release profile of DTX from the DTX/FPR-675 Pluronic NPs, 20 mg of NPs were dispersed in 3 mL of phosphate-buffered saline and placed in a dialysis bag (Spectrum®, Rancho Dominequez, CA, USA; molecular weight cutoff 12,000–14,000), which was immersed in 15 mL of phosphate-buffered saline (pH 7.4) containing 0.1% (v/v) Tween 80 for sink conditions. The experimental setup was placed in a shaking water bath maintained at 37°C and was shaken horizontally at 150 rpm. At predetermined time intervals, 1.5 mL aliquots of the release medium (phosphate-buffered saline) were withdrawn and the total release medium was replaced with 15 mL of fresh phosphate-buffered saline to maintain sink conditions.

Quantification of the released DTX was determined using reverse-phase high performance liquid chromatography with a Capcell Pak-C18 column (4.6×250 mm, 5 μm) and an acetonitrile/water (55/45, v/v) mobile phase over 15 minutes at a flow rate of 1.0 mL per minute. The eluent was monitored by ultraviolet light absorption at 227 nm. As a positive control, 10 mg of DTX was dissolved in 0.27 g of Tween 80 (Sigma) and 0.2 g of ethanol (Tween 80/ethanol-based DTX formulation [Taxotere]) at room temperature. Prior to the release study, the free DTX stock solution was diluted with phosphate-buffered saline (pH 7.4) containing 0.1% (v/v) Tween 80 to a final concentration of about 250 μg DTX per mL.

**In vitro cytotoxicity and cellular uptake of DTX/FPR-675 Pluronic NPs**

In order to evaluate the cytotoxicity and cellular behavior of the DTX/FPR-675 Pluronic NPs in various cell lines, bovine aortic endothelial cells (used as normal cells) were cultured in Dulbecco’s Modified Eagle’s Medium (low glucose; Gibco, Grand Island, NY, USA) containing 10% (v/v) fetal bovine serum (Gibco) and 1% (w/v) penicillin-streptomycin, and murine macrophages (Raw264.7) and squamous cell carcinoma (SCC-7) cells were cultured in Roswell Park Memorial Institute 1,640 medium containing 10% (v/v) fetal bovine serum and 1% (w/v) penicillin-streptomycin at 37°C in a humidified 5% CO₂ and 95% air atmosphere, respectively. The cells were seeded at a density of 5×10⁴ cells per well in 96-well flat-bottomed plates and allowed to adhere overnight. The three types of cells were washed twice with fresh phosphate-buffered saline and incubated for 24 hours with various concentrations of the DTX/FPR-675 Pluronic NPs. The cytotoxicity of the DTX/FPR-675 Pluronic NPs was evaluated using a MTT-based assay. After 24 hours of incubation, the cell culture medium was eliminated. The cells were then washed twice with phosphate-buffered saline to eliminate the remaining drug-loaded Pluronic NPs. Next, 25 μL of MTT solution (5 mg/mL in medium) was added to each well in the dark, and the cells were incubated for 2 hours at 37°C. The precipitates (formazan crystals) were then added and dissolved by adding 200 μL of dimethylsulfoxide. The absorbance was measured at 570 nm using a microplate reader (VersaMax™, Molecular Devices LLC, Sunnyvale, CA, USA). The percentages of viable cells were then determined by reduction of MTT relative to the negative control without the drug-loaded Pluronic NPs.

In order to compare the cellular uptake behavior of the DTX/FPR-675 Pluronic NPs, 1×10⁴ cells were seeded on a
dish with a cover slip and allowed to attach for 1 day. After cell attachment, the medium was replaced with 2 mL of serum with a culture medium containing the DTX/FPR-675 Pluronic NPs (5 mg of DTX) and then incubated for one hour. Next, the cells were washed twice with phosphate-buffered saline (pH 7.4) and fixed with a 4% paraformaldehyde solution. For nuclear staining, the cells were incubated with 4',6-diamidino-2-phenylindole (3 mmol) for 10 minutes at 25°C, followed by several washes in phosphate-buffered saline. The intracellular localization of the DTX/FPR-675 Pluronic NPs was observed using an IX81-ZDC focus shift compensating microscope (Olympus, Tokyo, Japan), where the excitation and emission wavelengths were 673 nm and 692 nm, respectively.

Lymphatic tracking and tissue distribution of the DTX/FPR-675 Pluronic NPs
The mice were anesthetized intraperitoneally using a solution containing 8 mg/mL ketamine (Ketalar®, Panpharma, Fougères, France) and 0.8 mg/mL xylazine (Rompun®, Bayer Pharma, Puteaux, France) at 0.015 mL/g of body weight. The lymph node was examined after intradermal injection into the left forepaw pad with the DTX/FPR-675 Pluronic NPs in an aqueous dispersion (10 μL, 1 mg/mL of DTX) using an NIRF image system. Images for lymphatic tracking were obtained at predetermined time points after NP injection using a 580 nm filtered charge coupled device camera under illumination at 460–490 nm, which was implemented in a high sensitivity imaging system (Olympus OV100) for data acquisition and analysis. In order to evaluate the distribution of NPs in lymphatic tissue, the lymph node (brachial) was obtained from normal mice and tumor-bearing mice and the obtained tissue was embedded in a Tissue Tek® OCT compound (Sakura Finetek, Torrance, CA, USA) and cryo-sectioned at 8 μm (CM1900 microtome, Leica). The fluorescent images were observed under a fluorescence microscope (Olympus BX51 and Zeiss Axioskop2 FS Plus).

In vivo biodistribution, tumor targetability, and lymphatic tracking of the DTX/FPR-675 Pluronic NPs in tumor-bearing mice
To observe the in vivo biodistribution, SCC-7 tumor cells were induced in male nude mice (aged 5.5 weeks, Orient Bio Inc) by a subcutaneous injection of 1.0×10⁶ cells suspended in cell culture (Roswell Park Memorial Institute 1,640 medium, 10% fetal bovine serum, 1% antibiotic agent). When the tumor volume reached approximately 150–200 mm³, the mice received an intravenous injection of the DTX/FPR-675 Pluronic NPs with 10 mg of DTX per kg. In order to observe the tissue distribution, tumor targeting capability, and lymphatic tracking, each animal was positioned on an IVIS imaging system (Xenogen Corporation, Alameda, CA, USA) equipped with a white-light excitation, charge coupled device camera-based fluorescence detection, and a filter set appropriate for near-infrared-emitting FPR-675 dye molecules (emission detected at 700 nm). The in vivo imaging system, consisting of a cooled charge-coupled device camera mounted on a light-tight specimen chamber (dark box), a camera controller, a camera cooling system, and a Windows computer system, was used for data acquisition and analysis. The mice were placed in the specimen chamber mounted with the charge coupled device camera cooled to −90°C, with the field of view set at 25 cm above the sample shelf. The photon emission transmitted from the mice was measured. The grayscale photographic images and bioluminescence color images were superimposed using the Living Image® version 2.11 software overlay (Xenogen) and IGOR® image analysis software (version 4.02A, WaveMetrics, Lake Oswego, OR, USA). A region of interest was manually selected over the signal intensity. The area of the region of interest was kept constant and the intensity was recorded as maximum (photons s⁻¹ cm⁻² sr⁻¹ [steradian]) within a region of interest. The mice, kept in an anesthetic state by inhalation of Gerolan® solution (enflurane as the active agent, Choong Wae Pharmaceuticals, Seoul, Republic of Korea), were automatically moved into the imaging chamber for scanning. The in vivo characteristics of the DTX/FPR-675 Pluronic NPs were confirmed by measuring the NIRF intensity in the SCC-7 tumor-bearing mice (n=3). To compare the distribution in various lymphatic tissues of each sample, the mice were euthanized at 60 minutes post-injection. The various lymph nodes (superficial cervical, brachial, and inguinal) were dissected from the mice, and their fluorescence intensities were determined using a 12-bit charge coupled device camera (Image Station 4000 MM; Kodak, New Haven, CT, USA) equipped with a special C-mount lens and an FPR-675 bandpass emission filter (680–720 nm; Omega Optical, Brattleboro, VT, USA). Identical illumination settings (eg, lamp voltage, filter, exposure time) were used in all animal imaging experiments.

Statistical analysis
The data are expressed as the mean ± standard deviation of at least three experiments. Data for the in vivo characteristics using an NIRF system were calculated using the region of interest function of the Analysis Workstation software package (ART Advanced Research Technologies Inc., Montreal,
Results
Preparation and characterization of DTX/FPR-675 Pluronic NPs
To demonstrate the targeting and tracking of the DTX/FPR-675 Pluronic NPs in the primary tumor and metastatic lymphatics, as described in Figure 1, various DTX/FPR-675 Pluronic NPs were prepared with different soybean oil/Tween 80 ratios as a solubilizer for DTX and FPR-675.

To optimize the composition of the DTX/FPR-675 Pluronic NPs, the soybean oil and Tween 80 mixtures were prepared with different weight ratios and characterized as shown in Figures 2A and 2B. Assessment confirmed stable dispersion of the DTX/FPR-675 Pluronic NPs with a soybean oil/Tween 80 ratio of 0.2 in the aqueous phase. The diagram illustrates the sequential detection of the primary tumor and metastatic lymphatics.

Figure 1: Schematic and expected behavior of DTX/FPR-675 Pluronic NPs.
Notes: (A) Schematic of DTX/FPR-675 Pluronic NPs according to temperature-induced phase transition and (B) expected behavior of NPs in blood vessels and lymphatic vessels.
Abbreviations: DTX, docetaxel; NPs, nanoparticles; FPR-675, Flamma™ fluorescence molecular imaging dye; NIR, near-infrared; PEO-PPO-PEO, poly(ethylene oxide)-poly(propylene oxide)-poly(ethylene oxide).
medium with a single distribution of 10.34±4.28 nm (Figure 2C). The maximum loading capacity and encapsulation efficiency of the NPs were 3.84±0.65 wt% and 94±2.67 wt%, respectively.

The in vitro release profiles for the DTX/FPR-675 Pluronic NPs were measured at pH 7.4 as shown in Figure 2D. The DTX/FPR-675 Pluronic NPs showed a sustained release pattern typical of that observed for drug release from a homogeneous polymer matrix, and differed from that found with free DTX. Therefore, DTX/FPR-675 Pluronic NPs of 10.34±4.28 nm were used throughout the experiments.

In vitro cytotoxicity and cellular uptake of DTX/FPR-675 Pluronic NPs

To compare the cytotoxicity and cellular uptake behavior of the DTX/FPR-675 Pluronic NPs, three cell types (normal [bovine aortic endothelial] cells, macrophage [Raw264.7] cells, and tumor [SCC-7] cells) were evaluated by MTT assay and fluorescent images (Figure 3).

Growth inhibition was found in all cells and the DTX/FPR-675 Pluronic NPs showed drug concentration-dependent cytotoxicity in the cell culture system. The DTX/FPR-675 Pluronic NPs had a more efficient cytotoxic effect on the SCC-7 tumor cells than on the bovine aortic endothelial cells and Raw264.7 cells.

To observe the cellular uptake behavior, the DTX/FPR-675 Pluronic NPs were incubated for 60 minutes and compared by the fluorescence intensity of each cell (Figure 3B). When their subcellular localization was observed under a fluorescent microscope, the fluorescence from the DTX/FPR-675 Pluronic NPs was not in the nuclear compartment of the SCC-7 tumor cells. All cells treated with the DTX/FPR-675 Pluronic NPs exhibited a red fluorescence signal in the cytoplasm, and the intensity was particularly high in the SCC-7 tumor cells. In contrast, the bovine aortic endothelial cells and Raw264.7 cells showed very low fluorescence intensity, indicating the possibility of low unintended uptake of...
the DTX/FPR-675 Pluronic NPs into normal cells and macrophages.

**Lymphatic tracking and tissue distribution of DTX/FPR-675 Pluronic NPs in normal and tumor-bearing mice**

To compare the lymphatic accumulation and tissue distribution in two types of mice (non-tumor-bearing and tumor-bearing), the lymph nodes were identified immediately after intradermal injection of DTX/FPR-675 Pluronic NPs into the forepaw pad of mice using a real-time NIRF imaging system. After injection, the DTX/FPR-675 Pluronic NPs in the tumor-bearing mice were rapidly transported from the injection site to the lymphatics. The lymph nodes and lymphatic vessels were visualized within one minute and then were continually visualized for a minimum of 60 minutes (early diagnosis time), and then we gradually decreased the fluorescence intensity at the lymph node. However, in the normal mice, there was no transport of the DTX/FPR-675 Pluronic NPs to the lymphatics and the flow status was observed at the initial time due to injected pressure on the mouse forepaw pad (Figure 4A). Therefore, accurate sites for the lymphatics and lymphatic vessels were more intensely recognized in tumor-bearing mice. Likewise, quantitative analyses of the fluorescence intensity at the brachial node were performed for an extended time period, and higher fluorescence intensities were observed in tumor-bearing mice than in normal non-tumor-bearing mice (Figure 4B). Further, fluorescence images from a frozen section of lymphatic tissue were observed to verify enhanced accumulation of DTX/FPR-675 Pluronic NPs in a tumor-bearing mouse. The DTX/FPR-675 Pluronic NPs accumulated in both the external and internal regions of the lymph node (Figure 4C).

**Biodistribution, tumor targetability, and lymphatic tracking of DTX/FPR-675 Pluronic NPs in tumor-bearing mice**

Because the FPR-675 was physically entrapped in the NPs, the difference in biodistribution was observed between the DTX/FPR-675 Pluronic NPs and the free FPR-675.29,30 Figure 5A demonstrates that the DTX/FPR-675 Pluronic NPs have stronger fluorescence intensity of a yellow and red color than the free FPR-675 in the superficial cervical node and brachial node until 60 minutes post-injection. In addition, the quantification data for each lymph node showed a significant difference and an approximately four-fold higher intensity than the free FPR-675, as shown in Figure 5B and 5C. Similar results were observed with the excised lymphatic tissues as presented in Figure 5D. Interestingly, the fluorescence intensity from the tumor-bearing side was more significantly different than that from the non-tumor-bearing side.

The whole body distribution behavior of the DTX/FPR-675 Pluronic NPs in vivo was monitored using real-time NIRF imaging for up to 9 hours post-injection (Figure 6A). After intravenous injection, the DTX/FPR-675 Pluronic NPs showed strong fluorescence signals at the lymphatics (superficial cervical node and brachial node) from 3 hours post-injection. The fluorescence at the tumor site gradually increased after one hour post-injection, and the highest NIRF intensity in the tumor was observed at 7 hours post-injection. Figure 6A confirms that the experimental results presented in Figure 5 were maintained for extended periods of time.

The renal clearance of DTX/FPR-675 Pluronic NPs is presented in Figure 6B.31 Strong fluorescence intensity was observed in the bladder of the normal mouse at one hour post-injection and this indicates that the DTX/FPR-675 Pluronic

---

**Figure 3** Cytotoxicity and in vitro cellular uptake of the NPs in various cells.

**Notes:** (A) In vitro cytotoxicity at various concentrations of the DTX/FPR-675 Pluronic NPs from 0.1 g/ml to 1,000 g/ml, and (B) cellular behavior of the DTX/FPR-675 Pluronic NPs using normal cells (BAECs), macrophages (Raw264.7), and tumor (SCC-7) cells. Data are presented as the mean ± standard deviation (n=6).

**Abbreviations:** BAECs, bovine aortic endothelial cells; DTX, docetaxel; NPs, nanoparticles; FPR-675, Flamma™ fluorescence molecular imaging dye; TRITC, tetramethylrhodamine; DAPI, 4',6-diamidino-2-phenylindole.
NPs in normal mice were rapidly excreted from the body through the kidneys and bladder. However, minimal clearance of the DTX/FPR-675 Pluronic NPs was observed in the bladders of tumor-bearing mice.

**Discussion**

It is generally accepted that nanoparticulate systems with diameters of less than 200 nm can avoid uptake by the reticuloendothelial system, circulate in the bloodstream for a prolonged period, and consequently accumulate in the tumor region. Furthermore, if their size is less than 20–30 nm, they can be utilized for lymphatic delivery. Based on previous studies, DTX/FPR-675 Pluronic NPs with a diameter of approximately 10.34±4.28 nm were prepared by temperature-induced phase transitions. Given their size, DTX/FPR-675 Pluronic NPs can be utilized as vehicles for lymphatic delivery of molecular imaging probes and chemotherapeutic agents. As shown in Figure 3, the DTX/FPR-675 Pluronic NPs were successfully delivered to the cytoplasm of SCC-7 tumor cells, which is the target site of the drug; however, minimal uptake was observed in other cells (normal [bovine aortic endothelial] cells and macrophages [Raw264.7]). It has been reported that the major factors affecting efficient tumor targeting of NPs have a close relationship with greater uptake of NPs into tumor cells than other cell types. Based on this previous report, the minimal cellular uptake of the NPs in normal cells can be explained. If the DTX/FPR-675 Pluronic NPs are delivered to the tumor site, they can be taken up by cancer cells efficiently and lymphatic delivery can be expected, as described in Figure 1. As presented in Figure 4, conclusive evidence for delivery of DTX/FPR-675 Pluronic NPs to the lymph nodes in tumor-bearing mice strongly supports the potential application of DTX/FPR-675 Pluronic NPs as a multifunctional nanocarrier for molecular imaging probes and chemotherapeutic drugs.

Because FPR-675 is physically entrapped in the NPs, release of FPR-675 is expected during systemic circulation of the NPs. The whole body distribution of the DTX/FPR-675 Pluronic NPs showed more enhanced accumulation at the lymph nodes (superficial cervical and brachial) than that of free FPR-675 (Figure 5A–C). This significant difference in biodistribution behavior between DTX/FPR-675 Pluronic NPs and free FPR-675 confirmed that the fluorescence from DTX/FPR-675 Pluronic NPs represented the tracking of NPs. Figure 5D indicates similar behavior of DTX/FPR-675 Pluronic NPs in the excised lymphatic tissue. In addition, a clear difference in the fluorescence intensity was observed between the tumor-bearing side and non-tumor-bearing side in the tumor-bearing mice, indicating accurate tracking of metastatic lymphatics using DTX/FPR-675 Pluronic NPs. These results suggest that a biodistribution study using DTX/FPR-675 Pluronic NPs could provide information on the lymphatic tracking of NPs in a tumor environment.

The whole body distribution and renal clearance behavior of the DTX/FPR-675 Pluronic NPs were examined for extended periods, as shown in Figure 6. Figure 6A confirms that the experimental results presented in Figure 5 were maintained for extended periods of time. In the absence of abnormal (ie, tumor) tissue for enhanced permeation and retention, rapid renal clearance was confirmed in the normal
Figure 5 In vivo real-time and excised lymph node near-infrared fluorescence imaging of tumor-bearing mice injected with DTX/FPR-675 Pluronic NPs for lymph node imaging. Notes: In vivo whole body near-infrared fluorescence images of mice (n=4) 60 minutes post-injection of (A) free FPR-675 (upper) and DTX/FPR-675 Pluronic NPs (lower). Comparison of near-infrared fluorescence intensity in (B) an excised superficial cervical (right) and a brachial (left) node and (C) near-infrared fluorescence images using DTX/FPR-675 Pluronic NPs. (D) Representative ex vivo fluorescence images of dissected lymph nodes (superficial cervical, brachial, and inguinal). 

Abbreviations: DTX, docetaxel; NPs, nanoparticles; FPR-675, Flamma™ fluorescence molecular imaging dye; IV, intravenous.
mice through observation of strong fluorescence intensity in the bladder one hour post-injection (Figure 6B). These results also indicate prolonged circulation of DTX/FPR-675 Pluronic NPs in the blood, leading to enhanced tumor accumulation at the primary tumor and metastatic lymphatics.13–35

Conclusion

DTX/FPR-675 Pluronic NPs were prepared using self-assembled Pluronic NPs with soybean oil/Tween 80 mixtures, and their use for accurate detection of primary tumors and metastatic lymphatics was demonstrated. At the optimum composition of soybean oil/Tween 80 mixtures with DTX and FPR-675, the DTX/FPR-675 Pluronic NPs were prepared with a single distribution of diameter (approximately 10 nm) and a clear difference in lymphatic accumulation was seen between normal and tumor-bearing mice. The preliminary results of this study demonstrated that DTX/FPR-675 Pluronic NPs may provide a strategy for improving cancer therapy through simultaneous delivery of molecular imaging probes and chemotherapeutic agents to the primary tumor, tumor lymphatics, and tumor-associated metastatic lymphatics.

Acknowledgment

This work was supported by National Research Foundation of Korea grants (20110027932, 2012028831, and 2013063969) funded by the Korean government.

Disclosure

The authors report no conflicts of interest in this work.

References

Sequential detection of primary tumor and metastatic lymphatics


