Priority Report

Real-time Monitoring of In Vivo Acute Necrotic Cancer Cell Death Induced by Near Infrared Photoimmunotherapy Using Fluorescence Lifetime Imaging

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Abstract

A new type of monoclonal antibody (mAb)-based, highly specific phototherapy (photoimmunotherapy; PIT) that uses a near infrared (NIR) phthalocyanine dye, IRDye700DX (IR700) conjugated with a mAb, has recently been described. NIR light exposure leads to immediate, target-selective necrotic cell death in vitro. Detecting immediate in vivo cell death is more difficult because it takes at least 3 days for the tumor to begin to shrink in size. In this study, fluorescence lifetime (FLT) was evaluated before and after PIT for monitoring the immediate cytotoxic effects of NIR mediated mAb-IR700 PIT. Anti-epidermal growth factor receptor (EGFR) panitumumab-IR700 was used for targeting EGFR-expressing A431 tumor cells. PIT with various doses of NIR light was conducted in cell pellets in vitro and in subcutaneously xenografted tumors in mice in vivo. FLT measurements were obtained before and 0, 6, 24, and 48 hours after PIT. In vitro, PIT at higher doses of NIR light immediately led to FLT shortening in A431 cells. In vivo PIT induced immediate shortening of FLT in treated tumors after a threshold NIR dose of 30 J/cm² or greater. In contrast, lower levels of NIR light (10 J/cm² or smaller) did not induce shortening of FLT. Prolongation of FLT in tissue surrounding the tumor site was noted 6 hours after PIT, likely reflecting phagocytosis by macrophages. In conclusion, FLT imaging can be used to monitor the acute cytotoxic effects of mAb-IR700-induced PIT even before morphological changes can be seen in the targeted tumors. Cancer Res; 72(18): 4622–8. © 2012 AACR.

Introduction

Most of the side effects of current cancer therapies are caused by damage to normal cells. Therefore, maximizing cytotoxicity to cancer cells while minimizing damage to normal cells is a highly desirable design criterion for new therapies. We recently developed a new type of monoclonal antibody (mAb)-based, highly specific phototherapy (photoimmunotherapy; PIT) that uses a near infrared (NIR) phthalocyanine dye, IRDye700DX (IR700) conjugated with mAb (1, 2). When the mAb-IR700 conjugate binds to the target cell and is exposed to sufficient, but still nonthermal, doses of NIR light, highly specific and immediate cell death occurs. The therapeutic effect is only observed if the mAb-IR700 binds to the target cell membrane and is exposed to NIR, otherwise the conjugate is harmless to nontargeted cells. Although targeted cell death can be observed immediately in vitro it is more challenging to see immediate changes in vivo because size changes take 3 to 4 days to become visible (3). Real-time monitoring of PIT effects could be important for ascertaining whether a PIT session has been effective and whether additional cycles of therapy are needed (1). This might include additional doses of light, higher intensity light or additional doses of the mAb-IR700 conjugate or all of these. Immediate feedback is especially important during surgical or interventional procedures under endoscopy. However, no clinically applicable imaging technology exists for assessing real-time effects of PIT (4, 5).

In addition to being a potent photosensitizer, IR700 is also a fluorophore. Its fluorescence can be used to direct NIR light, thus further reducing potential toxicity. IR700 has a relatively long fluorescence lifetime (FLT; refs. 6–12), and therefore, a change in IR700 FLT could be a good predictor of target cell death. In this study, we measured the FLT of mAb-IR700 after PIT with various doses of NIR light exposure and used FLT for real-time monitoring of the cytotoxic effects of PIT.

Materials and Methods

Reagents

Panitumumab, a fully humanized IgG2 mAb directed against the human epidermal growth factor receptor (EGFR), or HER1, was purchased from AMGEN Inc. A water soluble, silicon-phthalocyanine derivative, IRDye 700DX NHS ester (IR700; C21H22NO14Na2O2S3Si3, molecular weight of 1954.22) was purchased from LI-COR Bioscience. All other chemicals used were of reagent grade.

Synthesis of IR700-conjugated panitumumab

Panitumumab (1 mg, 6.8 nmol) was incubated with IR700 (66.8 μg, 34.2 nmol, 5 mmol/L in DMSO) in 0.1 mmol/L Na2...
HPO₄ (pH 8.6) at room temperature for 1 hour. Then the mixture was purified with a Sephadex G50 column (PD-10; GE Healthcare). The protein concentrations were determined with Coomassie Plus Protein Assay Kit (Pierce Biotechnology) by measuring light absorption at 595 nm (8453 Value System; Agilent Technologies). The concentration of IR700 was measured by absorption with spectroscopy to confirm the average number of fluorophore molecules conjugated to each panitumumab molecule. The number of IR700 per antibody was approximately 4 for the 1:4.5 reaction conditions. The addition of 0.4% SDS to the sample divided by conjugation is defined as the fluorescence intensity with SDS divided by fluorescence intensity without SDS. Panitumumab-IR700 conjugate (Pan-IR700) showed a quenching efficiency for a particular conjugation is defined as the fluorescence intensity with SDS divided by fluorescence intensity without SDS. Panitumumab-IR700 conjugate (Pan-IR700) showed a quenching efficiency of about 4.0 at pH 7.2. Pan-IR700 was kept at 4°C in the refrigerator as a stock solution.

Fluorescence lifetime measurements

All FLT experiments were conducted with the eXplore Optix-MX2 system (ART Advanced Research Technologies, Inc.; refs. 13, 14). A fixed pulsed laser diode was used as an excitation source at a wavelength of 670 nm. Region of interest (ROI) measurements with a spot size of 1.5 mm were selected at the image plane. The laser power was automatically chosen as the highest power that does not saturate the photon detector.

Lifetime analysis was conducted by using the ART OptiView (ART Advanced Research Technologies, Inc.). Lifetime values and lifetime mapping were calculated to fit fluorescence temporal point-spread functions (TPSFs) as single-exponential models with the Fit TPSF tool.

PIT for in vitro and in vivo models

PIT was conducted with a red light-emitting diode (LED) light at 680 to 700 nm wavelength (Tech-LED, Marubeni America Co.; ref. 2). Power densities were measured with an optical power meter (PM 100, Thorlabs).

Determination of FLT for Pan-IR700

Samples of Pan-IR700 at concentrations of 2.5, 5, 20, and 40 µg/mL were prepared by dilution with PBS. The fluorescence intensities and lifetimes of each sample were determined using the Optix MX2 system at room temperature within a 1.7 mL centrifuge tube.

To investigate the effect of PIT using Pan-IR700, the FLT of each sample at the concentration of 50 µg/mL was measured after irradiating the samples at a PIT dose of 0, 2, 4, 8, 15, and 30 J/cm².

Cell line

The HER1 positive cell line, A431 was used for HER1 targeting studies with panitumumab conjugates. The cell line was grown in RPMI 1640 (Life Technologies) containing 10% fetal bovine serum (Life Technologies), 0.03% t-glutamine, 100 units/mL penicillin, and 100 µg/mL streptomycin in 5% CO₂ at 37°C.

Cell pellet FLT studies

A431 cells were plated on 75 mm² cell culture flasks and incubated until confluent. Then Pan-IR700 conjugate was added to the media (1 µg/mL), and cells were incubated for 24 hours at 37°C. Upon completion of incubation, cells were removed from the flasks, and centrifuged to obtain pellets. The resulting cell pellets were washed three times with PBS and placed in 1.7 mL centrifuge tubes. The fluorescence intensities and lifetimes of each sample were then obtained.

To investigate the effect of cellular internalization with Pan-IR700 conjugates, A431 cells were plated on a 75 mm² flask and were incubated with Pan-IR700 for 1, 2, 4, 6, 15, and 24 hours. After removing the flasks and obtaining A431 cell pellets, FLT measurements of the A431 pellet were acquired.

After the A431 cell pellets were incubated overnight with Pan-IR700, cell pellets were irradiated at doses of 0, 2, 4, 8, 15, and 30 J/cm². After that, these pellets were gently washed with PBS × 1 and fluorescence intensity and lifetime images were obtained.

To detect the antigen specific localization of IR700 and to confirm the morphologic changes of A431 cells before and after PIT, fluorescence microscopy was conducted using Olympus BX61 microscope (Olympus America) equipped with the following filters: a 590 to 650 nm excitation filter, a 665 to 740 nm band pass emission filter. Transmitted light differential interference contrast images were also acquired. A431 cells were plated on a cover glass-bottomed culture well and incubated for 24 hours. Pan-IR700 was added to the medium (10 µg/mL), and the cells were incubated for either 6 or 24 hours. Once complete, the cells were washed once with PBS, and fluorescence microscopy was conducted before and after PIT.

Mouse model

All procedures were carried out in compliance with the Guide for the Care and Use of Laboratory Animal Resources (1996), National Research Council, and approved by the local Animal Care and Use Committee. A431 cells (1 × 10⁶ cells) were injected subcutaneously on both sides of the dorsum of female nude mice (National Cancer Institute Animal Production Facility). The experiments were conducted at 6 to 9 days after cell injection.

In vivo FLT imaging studies after PIT

Tumor-bearing mice were divided into 3 groups of 5 mice per group for the following irradiation doses of PIT: 10, 30, and 50 J/cm². As a control, 5 mice were prepared without PIT. One hundred micrograms of Pan-IR700 were injected intravenously via the tail vein into every mouse 24 hours before PIT. A431 tumors in the right-hand side of the dorsum were treated with PIT while the contralateral control tumors were shielded from light exposure with aluminum foil. After PIT, FLT images were obtained at the following time points: 0, 6, 24, and 48 hours. Zero hours acquisitions were conducted immediately after PIT. Maximum spot values of each ROI in the FLT images were calculated for tumors on both sides of the dorsum.
**Histologic analysis**

To evaluate serial histologic changes immediately (within 5 minutes) after PIT with various NIR light doses, microscopy was conducted (BX51, Olympus America). A431 tumors were harvested in 10% formalin immediately after 0, 10, 30, and 50 J/cm² of NIR light exposure. Serial 10-μm slice sections were fixed on a glass slide with H&E staining.

**Statistical analysis**

Statistical analyses were carried out using a statistics program (GraphPad Prism; GraphPad Software). Mann–Whitney’s U test was used to compare the lifetime value between those of treated tumors and untreated tumors. Student t test was used to compare with the lifetimes of treated tumors to no treatment control. P < 0.05 was considered to indicate a statistically significant difference.

**Results**

**FLT is independent from the Pan-IR700 concentration in the solution**

The FLTIs of various concentrations of Pan-IR700 were approximately the same, 3.56 ± 0.08 nanoseconds (ns); 3.62 (2.5 μg/mL), 3.58 (5 μg/mL), 3.44 (20 μg/mL), 3.60 ns (40 μg/mL), whereas the fluorescence intensities were decreased in proportion to the concentration (Fig. 1A and B).

**NIR light exposure alone does not affect the FLT of Pan-IR700**

Pan-IR700 (50 μg/mL) by itself was irradiated and FLT was measured. Both fluorescence intensity and lifetime did not change by irradiation of LED at the dose of 0, 2, 4, 8, 15, and 30 J/cm². The FLT was approximately 3.44 ± 0.06 ns.

**Internalization of Pan-IR700 prolonged the IR700 FLT**

FLT of A431 cells increased with the duration of the incubation with Pan-IR700. The FLTIs of A431 cell pellet at 1, 2, 4, 6, 15, and 24 hours of incubation were 2.98, 3.05, 3.13, 3.15, 3.36, and 3.41 ns, respectively. After 15 hours incubation, FLT of IR700 reached its peak and showed no further prolongation (Fig. 1D).

**Greater exposure of NIR light shortened the FLT of IR700 containing A431 cells**

PIT with greater NIR light doses induced greater shortening of FLT in A431 cell pellets incubated with Pan-IR700 for 24 hours before exposure to the NIR light (Fig. 1C). PIT shortened...
the FLT of A431 pellets down to 3.28, 3.09, 2.94, and 2.85 ns at doses of 0, 8, 15, and 30 J/cm², respectively.

**PIT induced typical necrotic cell death in A431 cells as well as rupture of lysosomes**

Under microscopy, Pan-IR700 was seen on the cell membrane and within endolysosomes at 24 hours after incubation. Following exposure to NIR light, immediate damage was induced in the cell membranes and lysosomes. Multifocal bleb formation was seen in the cellular membranes, characteristic of necrotic cell death induced by PIT (Fig. 2).

**Effective PIT induced immediate shortening of the FLT of IR700 in vivo**

The average FLT of A431 tumors 1 day after administration of 100 μg of Pan-IR700 in vivo was 3.27 ± 0.46 ns (n = 40). Significant shortening of FLT was induced immediately after PIT with NIR light doses of 30 and 50 J/cm² to experimental tumors (right dorsum, 30 J/cm²; down to 61.5% of 100 μg of Pan-IR700 in vivo Effective PIT induced immediate shortening of the FLT of IR700 in vivo). FLTs were immediately shortened to 69.1% of untreated tumors in the same mouse; P < 0.01, 50 J/cm²; down to 69.0% ± 10.9% of untreated tumors in the same mouse; P < 0.05).

Transient prolongation of IR700 FLT was found in and around PIT treated tumors 6 hours after PIT at NIR light doses of 30 and 50 J/cm² but continued to shorten at ≥24 hours after PIT. PIT with 10 J/cm² did not show this transiently prolonged FLT. IR700 FLT in untreated control tumors also slightly shortened at late time points (Fig. 3A).

Comparison with IR700 FLT between exposed and nonexposed tumors with NIR light of 30 and 50 J/cm² in the same mice showed significant differences within 5 minutes and, 24 and 48 hours after PIT (P < 0.05; Fig. 3B and C). The differences of IR700 FLT at 6 hours post-PIT were not statistically significant due to the diffuse temporal increase around exposed tumors. IR700 FLT of exposed and nonexposed tumors with NIR light of 10 J/cm² did not show significant difference at any time point (Fig. 3D).

FLT in PIT treated tumors with 50 and 30 J/cm² shortened significantly (P < 0.01) compared with no treatment controls (0 J/cm²). FLTs were immediately shortened to 69.1% ± 10.9% and 61.5% ± 5.1% by PIT with 50 and 30 J/cm², respectively. A431 tumors irradiated with only 10 J/cm² showed no significant shortening of FLT immediately after PIT. FLT shortened by only 7.7% at 48 hours after PIT compared with the untreated control (Fig. 4A). Interestingly, the FLT of nonirradiated tumors in PIT treated mice shortened slightly more than that in the untreated mice, but these changes were not significant, however, FLT became shorter with larger doses of NIR light to the treated tumors. These changes may be caused small amounts of light diffusing through the soft tissues from the "treated" side to the "untreated" side, thus explaining the dose-dependence of the effect.

**Histologic analysis**

Microscopy of treated tumors revealed various degrees of necrosis and microhemorrhage with clusters of healthy or damaged but potentially viable tumor cells after PIT. Necrotic damage was diffuse and intense and the amount of surviving tumor cells was reduced when 30 or 50 J/cm² of NIR light was administered. In contrast, when 10 J/cm² of NIR light was administered, necrotic cell damage was found in only limited areas with relatively large areas of viable cancer cells accounting for the majority of the tissue (Fig. 4C).

**Discussion**

Fluorescence microscopy studies showed Pan-IR700 gradually internalized into lysosomes in A431 cells at 37°C (Fig. 2). As Pan-IR700 internalized (Fig. 1D), IR700 FLT became longer as a function of incubation time. IR700 eventually accumulated in the lysosome. After exposure to a threshold intensity of NIR light, Pan-IR700 induced immediate outer cell membrane damage and damage to lysosomes, resulting in accumulation of IR700 within the cytoplasm and into the extracellular space. This damage was associated with a significant reduction in IR700 FLT. This implies that cellular internalization of the Pan-IR700 conjugate by itself prolongs IR700 FLT as it accumulates in the endolysosome. However, by damaging membrane structures, including the lysosomal membrane, PIT induces cell death and releases long FLT IR700, into the cytoplasm whereupon the FLT markedly shortens. Therefore, shortening FLT serves as an indicator of acute membrane damage induced by PIT.

Treatment with PIT with effective therapeutic light dose of NIR leads to shortened IR700 FLT in cancer cells in vitro and in tumors in vivo. The shortening of FLT was dependent on the dose of NIR light exposure in vitro (Fig. 1C). PIT with suboptimal doses of NIR light (10 J/cm²) did not show significant shortening of IR700 FLT in vivo. These differences could be
ascribed to the population of cancer cells, which received PIT effects. Our previous study showed that PIT with 50 J/cm² of NIR light exposure or more could eradicate A431 tumors. PIT with 30 J/cm² was not sufficient to totally eradicate tumors but caused tumor shrinkage and growth delay, indicating while not all cells were killed, most were severely and irreversibly damaged (2).

Shortened FLT of treated tumor in vivo was observed within 30 minutes of a single effective dose of NIR light and indicated a biologic effect several days before tumor size and shape changed. Although size of the lesion is considered a major indicator of cell death, it does not happen fast enough to determine if treatment has been effective. In the specific case of PIT, where light can be reapplied if necessary, a more immediate readout of cell death is needed. Size changes do not occur rapidly enough for monitoring cytotoxic effects. This is especially true of surgical or endoscopic procedures where it is preferable to complete treatments at one setting (2). FLT, because it is an immediate readout of the tumor’s condition, can assess the therapeutic effects of PIT to the cancer cells immediately after treatment and aids in deciding whether additional doses of NIR light exposure are necessary or not during the procedure (15, 16).

Interestingly, after an initial shortening of the FLT, it briefly became longer at about 6 hours after PIT. By 24 hours after PIT the FLT was reduced again (Fig. 3). Since we have observed prolongation of IR700 FLT as it is being internalized we hypothesize that after cell membrane disruption caused by PIT, the IR700 leaks into the extracellular space where it is internalized by macrophages mobilized to respond to the release of cytokines associated with cell necrosis. This is supported by histologic findings at 6 hours post-PIT that show inflammatory infiltrates composed of macrophages, which are entering the space formerly occupied by viable tumor (1). This transient prolongation of IR700 FLT may therefore be a sign of effective cell damage followed by initiating tissue repair possibly mediated by the chemokine release or the toll-like receptor system induced by the fragmented DNA and lipid bilayer (17–19).

Fluorescent proteins (FP) are a potential alternative for monitoring tumor growth in vivo (20–23). Fluorescence imaging using FPs is better suited for longitudinal monitoring of the effects of photo-therapy (24, 25). Acutely, FPs retain their signal regardless of the viability of the cells and even in necrotic cells may be taken up by macrophages. Thus, even though FLT requires postprocessing of the fluorescence signal and uses relatively expensive equipment, it is better suited for detecting...
acute changes than FPs (26, 27). Fluorescence imaging with FPs has been used for longitudinal monitoring of the therapeutic effects of PIT (2). However, PIT-induced acute cell death can only be detected with optical methods such as FLT while longer term changes can be measured with FPs. It should be noted, however, that FLT is potentially clinically translatable while FPs, which require cell transfection are unlikely to be used clinically.

This data suggests that the FLT of Pan-IR700 is a robust measurement that does not depend on the concentration of Pan-IR700 or light exposure in solution. For example, the in vitro Pan-IR700 solution did not change its FLT at varying concentrations or after NIR light exposure with various doses. Therefore, only the surrounding chemical microenvironment seems to affect the IR700 FLT. While IR700 is normally fluorescent and reflects tumor burden, after catabolism in the lysosome and photo-bleaching, fluorescence may be reduced, thus leading to ambiguity regarding tissue viability. However, those photo-chemical and biochemical changes do not affect FLT. Therefore, shortening of FLT is a better biomarker than IR700 fluorescence intensity.

In conclusion, FLT is a potential method of assessing in near real-time, the cytotoxic effects of PIT using a mAb-IR700 conjugate, during surgical or endoscopic procedures. FLT is prolonged during endolysosomal internalization but rapidly shortened after cell damage. FLT again is prolonged for a brief period about 6 hours after PIT due to internalization by migrating macrophages. After that there is a steady reduction in FLT. Thus, FLT imaging allows the assessment of the effect of PIT before morphologic changes become evident.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

**Authors’ Contributions**

Conception and design: P.L. Choyke, H. Kobayashi

Development of methodology: T. Nakajima, H. Kobayashi

Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): T. Nakajima, K. Sano, M. Mitsunaga, H. Kobayashi

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**Figure 4.** A, FLT in PIT treated tumors with 50 and 30 J/cm² shortened significantly (P < 0.01) compared with no treatment control (0 J/cm², control). FLT's were immediately shortened to 69.1 ± 10.9% and 61.5 ± 5.1% by PIT with 50 and 30 J/cm², respectively. A431 tumors irradiated with only 10 J/cm² showed no significant shortens of FLT that were seen immediately. FLT shortened by only 7.7% at 48 hours after PIT compared with no treatment control. B, FLT of nonirradiated tumors in PIT treated mice shortened slightly more than that in the untreated mice over time, but these changes were not significant. They may be caused by a small amount of light diffusing through tissue from the irradiated side, even though the surface of the tumor was covered. Student t test was used for the statistical analysis. C, histologic specimens of A431 tumors, which were treated with PIT at 0, 10, 30, and 50 J/cm², are shown. All specimens were stained with hematoxylin and eosin. Microscopic evaluation of treated tumors revealed various degrees of necrosis and microhemorrhage with clusters of healthy or damaged tumor cells after PIT. Necrotic damage was diffuse and intense and fewer surviving tumor cells were seen when 30 and 50 J/cm² of NIR light was administered. In contrast, when only 10 J/cm² of NIR light was administered, necrotic cell damage was found in only limited areas within the tumor while substantial amounts of healthy cancer foci remained. Scale, 50 μm.
References


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