Delivery of optical contrast agents using Triton-X100, part 1: reversible permeabilization of live cells for intracellular labeling

Anne L. van de Ven
Rice University
Department of Bioengineering, MS 142
Keck Hall, Suite 116, 6100 Main Street
Houston, Texas 77005

Karen Adler-Storthz
University of Texas, Dental Branch
Department of Diagnostic Sciences
6516 M.D. Anderson Blvd., DBB, 4.133
Houston, Texas 77030

Rebecca Richards-Kortum
Rice University
Department of Bioengineering, MS 142
Keck Hall, Suite 116, 6100 Main Street
Houston, Texas 77005

Abstract. Effective delivery of optical contrast agents into live cells remains a significant challenge. We sought to determine whether Triton-X100, a detergent commonly used for membrane isolation and protein purification, could be used to effectively and reversibly permeabilize live cells for delivery of targeted optical contrast agents. Although Triton-X100 is widely recognized as a good cell permeabilization agent, no systematic study has evaluated the efficiency, reproducibility, and reversibility of Triton-X100–mediated permeabilization in live mammalian cells. We report a series of studies to characterize macromolecule delivery in cells following Triton-X100 treatment. Using this approach, we demonstrate that molecules ranging from 1 to 150 kDa in molecular weight can be reproducibly delivered into live cells by controlling the moles of Triton-X100 relative to the number of cells to be treated. When Triton-X100 is administered at or near the minimum effective concentration, cell permeabilization is generally reversed within 24 h, and treated cells continue to proliferate and show metabolic activity during the restoration of membrane integrity. We conclude that Triton-X100 is a promising permeabilization agent for efficient and reproducible delivery of optical contrast agents into live mammalian cells, © 2009 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.3090448]

Keywords: intracellular delivery; detergents; human telomerase reverse transcriptase.

Address all correspondence to: Rebecca Richards-Kortum, Rice University, Department of Bioengineering, MS 142, Keck Hall, Suite 116, 6100 Main Street, Houston, Texas 77005. Tel: 713-348-1823; Fax: 713-348-5877; Email: rnkortum@rice.edu.

1 Introduction

The identification of unique molecular markers of cancer and its precursors has stimulated the development of new cancer-screening tools. A technology facilitating the noninvasive identification of precancerous lesions at high risk for progression would have important clinical benefits. Molecular-specific optical contrast agents have garnered attention for their ability to selectively label cells expressing markers of interest. Optical probes targeting extracellular biomarkers have advanced to preclinical animal models and pilot clinical trials. The development of contrast agents targeted to intracellular biomarkers has proved to be more challenging. Multifunctional contrast agents designed to utilize cell uptake mechanisms have been reported; however, the engineering of highly specific, optical probes with the appropriate physiochemical properties for cell internalization is difficult and not always feasible. There is need for a strategy to facilitate the intracellular delivery of existing contrast agents into live cells.

Chemical detergents have traditionally been used as solubilizing agents for the study of biological membranes. At subsolubilizing concentrations, detergents can be used to perturb membrane structure and increase cell permeability. Chemical detergents have achieved widespread use as permeabilization agents for immunohistochemistry. In live cells, detergents have shown limited success as permeabilization agents due to the challenge of retaining cell viability following permeabilization. Of the detergents tested to date, Triton-X100 has shown the most promise as a cell permeabilization agent. Triton-X100 has been used to nondestructively release secondary metabolites in plant cells and to study in situ enzymatic assays in yeast. No systematic study has been performed in mammalian cells to determine how universally efficient, reproducible, and reversible this process is.

We sought to determine whether Triton-X100 can be used to effectively and reversibly permeabilize live mammalian cells for contrast agent delivery. In this paper, we describe the use of macromolecule uptake assays to evaluate the efficiency of cell permeabilization. We find that there is a minimum effective Triton-X100 concentration, in which cell permeabilization efficiency is balanced with cell loss after treatment. Cell recovery after treatment is evaluated using cell viability, population doubling, and membrane integrity assays. Our findings indicate that Triton-X100, when used at or near the

DOI: 10.1117/1.3090448

Paper 08269SSR received Aug. 4, 2008; revised manuscript received Nov. 11, 2008; accepted for publication Dec. 9, 2008; published online Mar. 10, 2009.
minimum effective concentration, can permeabilize a variety of live cells in a reproducible and reversible manner.

2 Materials and Methods

2.1 Cell Culture

The ability of Triton-X100 to permeabilize human cell lines was evaluated using carcinoma cells (1483, HeLa), SV-40 transformed cells (GM847), and primary cells (HDF, MCF-10A). The 1483 cell line was obtained from Dr. Reuben Lotan at the M.D. Anderson Cancer Center (Houston, Texas). HeLa and MCF-10A cells were purchased from American Type Culture Collection [(ATCC, Manassas, Virginia). GM847 cells were obtained from Coriell Cell Repositories (Camden, New Jersey). HDF cells were purchased from Lonza (Walkersville, Maryland). 1483 cells were cultured in Dulbecco’s Modified Eagle Medium: Nutrient Mix F-12™ medium supplemented with L-glutamine (Invitrogen, Carlsbad, California) and 10% fetal bovine serum [(FBS) Hyclone, Logan, Utah]. HeLa and GM847 cells were cultured in Minimum Essential Medium™ supplemented with L-glutamine, nonessential amino acids, sodium pyruvate, vitamins (Invitrogen) and 10% FBS. HDF cells were cultured in Fibroblast Growth Medium-2™ (Lonza). MCF-10A cells were cultured in Mammary Epithelium Growth Medium™ (Lonza).

2.2 Triton-X100 Dose Dependency of Cell Permeabilization

To determine the minimum effective Triton-X100 concentration required to permeabilize live cells, 1483, Hela, GM847, HDF, and MCF-10A cells were assayed for membrane integrity and cell viability following treatment with different concentrations of Triton-X100. Briefly, subconfluent monolayers of cells (plated 24 hours in advance at 5 x 10^4 cells/cm²) were washed once with cold phosphate buffered saline [(PBS) Sigma-Aldrich, St. Louis, Missouri] and treated with Triton-X100 (Sigma-Aldrich) for 10 min at 4 °C. Triton-X100 concentrations ranged from 0 to 5.5 pmol/cell, normalized relative to the cell count at the time of plating. After permeabilization, the cells were washed once in media and returned to the incubator. Cell viability was assessed 4 h following Triton-X100 treatment. Prewarmed 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT, Sigma-Aldrich) was added to the supernatant for 20 min at a concentration of 0.5 mg/ml. In viable cells, reduction of the MTT reagent led to the intracellular deposition of a blue formazan product visible via light microscopy. Thereafter, the cells were washed, labeled with 1 μM 3 kDa rhodamine-dextran (Invitrogen), and imaged using confocal microscopy (described below). Viable cells were identified by the presence of blue formazan deposits. Permeabilized cells were identified by presence of fluorescent dextrans inside the cytoplasmic and nuclear compartments. The minimum effective Triton-X100 concentration was defined as the concentration at which ≥95% of cells were permeabilized. Then, 300–500 cells were evaluated from representative fields of view for each treatment condition in five independent experiments. Differences between cell lines were assessed using a two-tailed, unpaired Student’s t-test, with p values of <0.05 considered statistically significant.

2.3 Permeabilization Detection Using Macromolecules

To determine whether Triton-X100 treated cells are permeable to macromolecules of different sizes, monolayers of treated cells were probed with 3, 40, and 150 kDa fluorescent macromolecules. Briefly, subconfluent monolayers of 1483 cells were permeabilized with 0 to 1.1 pmol/cell Triton-X100 for 10 min at 4 °C. Cells were washed once in media and covered with a 1:1:1 mixture of rhodamine-dextran (3 kDa, Invitrogen), fluorescein-dextran (40 kDa, Invitrogen), and AlexaFluor647 IgG (150 kDa, Invitrogen), each diluted to a concentration of 1 μM in PBS. Cells were incubated with this solution for 20 min at 4 °C and then imaged using fluorescence confocal microscopy at three different excitation wavelengths. The optical imaging plane was focused to yield a cross section of the cytoplasm and nucleus for each cell in the field of view. Using NIH ImageJ v1.34 software (http://rsbweb.nih.gov/ij/), the mean fluorescence intensity was determined for selected regions of interest. The mean fluorescence intensity of the cytoplasm and nucleus was compared to that of the extracellular solution on a cell-to-cell basis. Cells were defined as permeable if the mean cytoplasmic and/or nuclear fluorescence intensity was equal or greater than half of the mean extracellular fluorescence intensity. The threshold parameter was selected based on the observation that not all cells reach 100% of the extracellular mean fluorescence intensity. About 100–300 cells were evaluated from representative fields of view in three separate experiments.

2.4 Tracking the Time Course of Macromolecule Entry

Macromolecule entry was tracked in real time to evaluate macromolecule penetration as a function of time. Rhodamine-dextran (3 kDa), fluorescein-dextran (40, 70 kDa), and AlexaFluor647 IgG were individually diluted to a concentration of 1 μM in 0.05% Triton-X100. These solutions were topically coverslips. Based on the cell count at the time of plating, an appropriate volume was selected to yield a final concentration of 0.55 pmol/cell Triton-X100. This concentration was selected to facilitate both cytoplasmic and nuclear penetration of the larger macromolecules. The movement of macromolecules into the cells was monitored using time-lapse fluorescence confocal microscopy. Again, the optical imaging plane was focused to yield a cross section of the cytoplasm and nucleus for each cell in the field of view. Images were acquired every 15 s. The mean fluorescence intensities of the cytoplasm and nucleus were compared to that of the extracellular solution by selecting regions of interest within a time stack and normalizing relative to the mean fluorescence intensity of the extracellular solution in each image. No other background corrections were made. Ten cells were evaluated from a representative field of view in five separate experiments. All fluorescence quantitation was performed using ImageJ software. The mean fluorescence intensity of each cell compartment 1 h after macromolecule addition and the time required to achieve >98% fluorescence accumulation was determined from time versus intensity plots for each macromolecule.
2.5 Cell Metabolic Assay

Cells were assayed for metabolic activity 24 h following Triton-X100 treatment by monitoring the reduction of MTT as described in Ref.26. Subconfluent monolayers of 1483 cells were permeabilized with 0–5.5 pmol/cell Triton-X100 for 10 min at 4 or 37 °C, washed once, and then returned to a prewarmed complete medium for 24 h. Cellular metabolic activity was evaluated in triplicate in five separate experiments using the ATCC MTT assay. The absorbance measurement from each well was divided by the number of cells well remaining after treatment to determine the mean absorbance per cell.

2.6 Population-Doubling Assay

Population-doubling experiments were performed with subconfluent monolayers of 1483 and MCF-10A cells to determine the effect of Triton-X100 treatment on cell proliferation. Cells were plated at an appropriate concentration to allow log-linear growth for five days. Twenty-four hours after plating, cells were washed with PBS, treated with 0.55 pmol/cell Triton-X100 or PBS for 10 min at 4 or 37 °C, and then returned to prewarmed complete media. The number of cells remaining after treatment and the rate of population doubling was assessed in duplicate at four different time points in three separate experiments. Differences in rate of population doubling between each treatment group were assessed using a two-tailed, unpaired Student’s t-test, with p values of <0.05 being considered statistically significant.

2.7 Membrane Integrity Assay

To determine whether cells recover their membrane integrity after Triton-X100 treatment, cells were tested for their ability to exclude macromolecules 8–24 h after permeabilization. Subconfluent monolayers of 1483 cells cultured on glass coverslips were permeabilized for 10 min at 4 °C with 0.55 pmol/cell Triton-X100, washed, and returned to prewarmed complete media. At regular intervals after permeabilization, cells were treated with 0.5 mg/ml MTT for 20 min at 37 °C, washed, and covered with a 1:1:1 mixture of rhodamine-dextran, fluorescein-dextran, and AlexaFluor647 IgG (each diluted to 1 μM in PBS). Viable cells were identified by color deposition under bright-field imaging. Macromolecule exclusion was monitored using fluorescence confocal microscopy. Cells were considered to have an intact membrane if the mean fluorescence intensity of the cytoplasm and/or nucleus was <1% of the extracellular intensity. About 100–300 cells were evaluated from representative fields of view in three separate experiments.

2.8 Immunocytochemistry of Live Cells

Cells were labeled with nuclear-specific antibodies to validate the use of Triton-X100 for delivery of targeted contrast agents. Polyclonal rabbit antihuman PC563-hTERT antibody was purchased from EMD Biosciences (San Diego, California). Monoclonal mouse antihuman NCL-hTERT antibody was purchased from Novoceastra (Newcastle upon Tyne, United Kingdom). Goat antirabbit AlexaFluor 647 IgG, goat antimouse AlexaFluor 488 IgG, purified polyclonal rabbit IgG, and monoclonal mouse IgG2a were purchased from Invitrogen. Subconfluent monolayers of live 1483 cells (5 \times 10^4 cells/cm²) cultured on coverslips were labeled with antibodies as follows: 24 h after plating, the cells were washed with PBS, treated with 0.55 pmol/cell Triton-X100 for 10 min, blocked for 10 min with a saline buffer containing 1% BSA/2% goat serum, and then labeled for 1 h with PC563-hTERT (1:80 dilution) and NCL-hTERT (1:800 dilution) antibodies. Following three washes, cells were probed with a 1:500 dilution of AlexaFluor 647 and 488 IgG antibodies for 1 h, washed, and imaged live using confocal microscopy. As a control, live cells were labeled in parallel with primary IgG antibodies and secondary AlexaFluor antibodies. For comparison to established immunocytochemistry protocols, cells were fixed for 30 min with 10% formalin immediately before or after Triton-X100 treatment. Fixed cells were immunolabeled as described above. Cell labeling was evaluated in triplicate in three separate experiments. All cells were imaged in the same optical plane. All stock solutions were diluted in PBS; all labeling was performed at 4 °C.

2.9 Confocal Image Acquisition

All images were obtained using a Carl Zeiss LSM 510 confocal microscope (Thornwood, New York) equipped with 488 nm (30 mW), 543 nm (1 mW), and 633 nm (5 mW) lasers. Images were collected using photomultiplier tube detectors and Zeiss LSM 5 image examiner software. Samples were sequentially excited with each laser line, with power settings of 50, 100, and 100% respectively. Fluorescence emission was collected using 505–550, 565–615, and 650–710 nm bandpass filters, respectively. Images were acquired at 0.5 fps using a 63X oil objective with a pinhole of 2.56 Airy units. For macromolecule uptake and exclusion studies, the gain was held constant below the saturation level of the extracellular solution. For the immunocytochemistry studies, the gain was held constant at 535 (488-nm excitation) and 440 (633-nm excitation). Fluorescence recovery after photobleaching tests were performed using these imaging parameters to ensure that no photobleaching of the macromolecules occurred.

3 Results

3.1 Dose-Dependent Permeabilization of Live Cells

To determine the relationship between cell permeabilization and short-term viability, monolayers of 1483 squamous carcinoma cells were treated for 10 min with different concentrations of Triton-X100 and then assayed for MTT reduction and 3 kDa rhodamine-dextran uptake. The Triton-X100 concentration was normalized relative to the number of cells being treated. Figure 1 shows the dose-dependent response to Triton-X100 treatment, in terms of picomols of Triton-X100 added per cell. The solid triangles indicate the percentage of cells permeabilized; the hollow diamonds indicate the percentage of viable cells remaining after treatment. Cells treated with 0.18 pmol/cell or less showed little to no uptake of dextran following treatment. Cells treated with 0.28 pmol/cell or above showed dextran uptake in >95% of the cells. Cell loss was dependent on Triton-X100 dose, with the percentage of viable cells decreasing as the concentration of Triton-X100 was increased. After treatment with 0.28 pmol/cell Triton-
X100, 98 ± 3% of the cells were viable, compared to 99 ± 1% of PBS-treated controls.

Different cell lines were evaluated for their sensitivity to Triton-X100 treatment. Table 1 shows the minimum Triton-X100 concentration required to permeabilize ≥95% of each cell population for the delivery of 3 kDa molecules. No significant differences were observed between squamous carcinoma cells, SV-40 transformed cells, and primary cells. The minimum effective Triton-X100 concentration was 0.27 pmol/cell for all cell lines evaluated. Cell viability after treatment was >90% for all cell lines at the minimum effective Triton-X100 concentration.

### 3.2 Macromolecule Penetration Following Permeabilization

To determine whether cells are selectively permeable to macromolecules of different sizes, monolayers of Triton-X100-treated cells were topically labeled with a 1:1:1 mixture of 3 kDa rhodamine-dextran, 40 kDa fluorescein-dextran, and 150 kDa AlexaFluor647 IgG. Dextrans were selected for their neutral charge, and IgG was selected for its size and structural similarity to targeted antibodies. The fraction of cells permeable to each macromolecule is shown in Fig. 2. Figure 2(a) shows the dose-dependent permeabilization of the cytoplasm; Fig. 2(b) shows that of the nucleus. The 3 kDa dextran suc-

![Graph showing permeabilization of 1483 cell monolayers treated with different concentrations of Triton-X100.](image)

**Table 1** Sensitivity of different cell lines to Triton-X100 permeabilization.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Derivation</th>
<th>Min. Triton concentration to permeabilize ≥95% of cells (pmol/cell)</th>
<th>Cells remaining at min. concentration (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1483</td>
<td>Oral squamous cell carcinoma</td>
<td>0.27 ± 0.07</td>
<td>97 ± 4</td>
</tr>
<tr>
<td>Hela</td>
<td>Cervical squamous cell carcinoma</td>
<td>0.27 ± 0.06</td>
<td>92 ± 8</td>
</tr>
<tr>
<td>GM847</td>
<td>SV-40 transformed fibroblast</td>
<td>0.27 ± 0.06</td>
<td>93 ± 6</td>
</tr>
<tr>
<td>HDF</td>
<td>Primary fibroblast</td>
<td>0.26 ± 0.06</td>
<td>98 ± 3</td>
</tr>
<tr>
<td>MCF-10A</td>
<td>Primary epithelial cell</td>
<td>0.27 ± 0.04</td>
<td>94 ± 7</td>
</tr>
</tbody>
</table>

![Graph showing macromolecule penetration into the cytoplasm and nucleus as a function of Triton-X100 concentration.](image)
cessfully penetrated both cell compartments in over 95% of cells treated with 0.22-pmol/cell Triton-X100 and above. The penetration of larger molecules varied by cell compartment. At 0.22 pmol/cell Triton-X100, 83% of the cells that were permeable to 3 kDa dextran also contained 40 kDa dextran and 150 kDa IgG in the nucleus. Concentrations upwards of 1.1 pmol/cell were required to permeabilize the nucleus of /H33356 95% of cells. Thus, for Triton-X100 concentrations ranging from 0.18 to 0.55 pmol/cell, we observed that some nuclei were permeable to 3 kDa molecules but not 40 and 150 kDa molecules. Interestingly, there was no exclusion of 150 kDa molecules from nuclei that were permeable to 40 kDa molecules. The mean fraction of cells permeabilized was independently validated by treating cells with one macromolecule at a time (data not shown) to exclude any interacting effects.

Macromolecule penetration was tracked as a function of time using time-lapse confocal microscopy. Molecules of all sizes could be observed in the cytoplasm and nucleus of cells as early as 15 s after the administration of Triton-X100. Figure 3(a) shows the cumulative uptake of fluorescent IgG (normalized relative to the fluorescence intensity of the extracellular solution) following the administration of 0.55 pmol/cell Triton-X100. The solid triangles represent the mean fluorescence intensity of the cytoplasm and nucleus of cells treated with IgG in the absence of Triton-X100. After the addition Triton-X100, fluorescent IgG rapidly enters both the cytoplasm and nucleus of cells. At approximately 5 min, both cell compartments reached a mean fluorescence intensity equal to that of the extracellular space. The error bars represent one standard deviation.

Fig. 3 Cumulative macromolecule penetration as a function of time following the administration of 0.55 pmol/cell Triton-X100. (a) The time course of AlexaFluo647 IgG accumulation in the cytoplasm and nucleus of live 1483 cells, averaged over five trials. Cells are permeabilized rapidly, with both the cytoplasm (○) and nucleus (□) achieving a mean fluorescence intensity equal to that of the extracellular space at ~5 min. Cells treated with IgG in the absence of Triton-X100 showed no significant uptake of fluorescence. All samples were normalized relative to the extracellular fluorescence. Error bars were removed for clarity; the error ranged from ±0.01 to 0.10 AU for one standard deviation. (b) The mean fluorescence intensity of each cell compartment relative to that of the extracellular space 1 h after Triton-X100 addition. (c) The time required for >98% macromolecule accumulation in the cytoplasm (■) and nucleus (□) of Triton-X100 treated cells. The time required for macromolecule uptake appears to be dependent on macromolecule size. No significant differences are observed between the cell compartments. The error bars represent one standard deviation.
Intensity of cell cytoplasm and nucleus was measured relative to that of the extracellular space 1 h after Triton-X100 addition. The total penetration varied by molecule size as well as the target cell compartment. The 3 kDa dextran uniformly penetrated both the cytoplasm and nucleus of treated cells. The 40 and 70 kDa dextrans penetrated to a lesser degree and preferentially localized to the cytoplasm. The 150 kDa IgG preferentially accumulated in both cell compartments and raised the mean fluorescence intensity of both cell compartments above that of the extracellular space.

The time required for macromolecule accumulation was measured using time-lapse confocal microscopy. Figure 3 shows the time required to achieve >98% macromolecule accumulation after Triton-X100 addition, as determined from the mean intracellular fluorescence accumulation over time. The time required for macromolecule uptake appeared dependent on macromolecule size, with the larger molecules requiring more time to accumulate inside treated cells. The macromolecule accumulation time ranged from 4.0 ± 0.8 to 9.1 ± 1.5 min for 3 and 40 kDa molecules, and from 11.8 ± 1.3 to 16.0 ± 1.3 min for the 70 and 150 kDa molecules. No significant differences were observed between cell compartments for any given macromolecule. These trends were independent of macromolecule concentration (data not shown).

3.3 Cell Recovery after Treatment
Monolayers of 1483 cells were assayed for metabolic activity 24 h after treatment with different concentrations of Triton-X100. Figure 5 shows the mean absorbance measured at 570 nm following MTT reduction by viable cells. Absorbance measurements of the total cell population at each Triton-X100 concentration yielded a dose-dependent decrease in absorbance. When the population data was adjusted for the number of cells remaining after 24 h after treatment, it was found that cells treated with 1.1 pmol/cell Triton-X100 or less retained full metabolic activity. Treatment with higher concentrations of Triton-X100 resulted in cell death and/or loss of metabolic activity, leading to a reduction in the mean absorbance per cell. No significant differences were observed in the metabolic activity of cells treated with 0.55 pmol/cell Triton-X100 at...
The proliferative capacity of 1483 cells following Triton-X100 treatment was determined using a population doubling assay. Figure 5(b) shows the cumulative cell number for three days following treatment with 0.55 pmol/cell Triton-X100 or PBS. Approximately 87 ± 1% of the adherent cells remained immediately after Triton-X100 treatment. Both treatment groups continued to proliferate with time. We observed a slight decrease in the rate of population doubling for each treatment group; however, the differences observed were not statistically significant (0.037 ± 0.012 cells/h for Triton-X100 treatment versus 0.041 ± 0.008 cells/h for PBS treatment). Similarly, no significant differences in population doubling were observed between Triton-X100 and PBS-treated primary MCF-10A cells or 1483 cells treated with Triton-X100 prewarmed to 37 °C (data not shown).

To determine whether cells recover their membrane integrity after Triton-X100 treatment, cell monolayers were tested for their ability to exclude macromolecules 8–24 h after treatment. Figure 5(c) shows the cumulative recovery of membrane integrity in metabolically active cells. The percentage of cells with intact membranes increases progressively with time. Membrane recovery was mostly restored at 24 h. (d) Confocal images of cells probed with MTT and fluorescent macromolecules 24 h after treatment with 0.55 pmol/cell Triton-X100. Of the cells that show metabolic activity, <10% of cells exhibit permeability to 3 kDa rhodamine-dextran (orange), 40 kDa fluorescein-dextran (green), or 150 kDa AlexaFluor 647 IgG (red). The arrows indicate cells that are permeable to macromolecules. The error bars represent one standard deviation.

4 °C when compared to those treated at 37 °C (data not shown).

The proliferative capacity of 1483 cells following Triton-X100 treatment was determined using a population doubling assay. Figure 5(b) shows the cumulative cell number for three days following treatment with 0.55 pmol/cell Triton-X100 or PBS. Approximately 87 ± 1% of the adherent cells remained immediately after Triton-X100 treatment. Both treatment groups continued to proliferate with time. We observed a slight decrease in the rate of population doubling for each treatment group; however, the differences observed were not statistically significant (0.037 ± 0.012 cells/h for Triton-X100 treatment versus 0.041 ± 0.008 cells/h for PBS treatment). Similarly, no significant differences in population doubling were observed between Triton-X100 and PBS-treated primary MCF-10A cells or 1483 cells treated with Triton-X100 prewarmed to 37 °C (data not shown).

To determine whether cells recover their membrane integrity after Triton-X100 treatment, cell monolayers were tested for their ability to exclude macromolecules 8–24 h after treatment. Figure 5(c) shows the cumulative recovery of membrane integrity in metabolically active cells, as measured by MTT reduction and macromolecule uptake. The percentage of metabolically active cells exhibiting intact cell membranes increased progressively with time. At the 24-h time point, ~10% were still permeable to macromolecules. Fig. 5(d) shows representative confocal microscopy images of cells assayed for membrane integrity 24 h after Triton-X100 treatment. Metabolically active cells were identified by color deposition using light microscopy. Cells that remained permeable after 24 h (indicated by arrows) admitted entry of 3 kDa (orange), 40 kDa (green), and 150 kDa (red) macromolecules.

3.4 Delivery of Targeted Optical Contrast Agents

Live and fixed cells were labeled with antibodies before or after treatment with 0.55 pmol/cell Triton-X100. Figure 6 shows representative confocal fluorescence images of 1483 cells probed for two different nuclear targets. The PC563-hTERT labeling is shown in red, and the NCL-hTERT labeling is shown in green. In cells fixed with 10% formalin, both targets were available for immunolabeling. Pretreatment with Triton-X100 before fixation enhanced the availability of the PC563 target. Treatment of live cells with Triton-X100 allowed both targets to be successfully labeled and imaged without fixation. Positive dual labeling was observed in all live permeabilized cells and varied in intensity less than ±5% from cell to cell (data not shown). A slight decrease in NCL-hTERT labeling intensity was observed when comparing live to fixed cells, likely due to repeated washing of the non-cross-linked proteins. No antibody cross-reactivity or nonspecific labeling was observed.

4 Discussion

Our results demonstrate that Triton-X100 can be used at subsolubilizing concentrations to permeabilize the membrane and nucleus of live cells. Both targeted and untargeted optical
contrast agents can be reproducibly delivered into live cells by using the appropriate concentration of Triton-X100. In membrane-bound protein solubilization studies, it is generally recognized that the Triton-X100 concentration should be adjusted relative to the amount of protein to be extracted similarly, we found it useful to normalize the Triton-X100 concentration for live cell permeabilization relative to the number of cells to be treated. Using this approach, we found that the Triton-X100 concentration could be readily optimized and scaled for treating different numbers of cells.

The cell permeabilization efficiency was evaluated as a function of Triton-X100 concentration. Studies of fluorescent dextran uptake suggest that cell permeabilization has a threshold, since as little as a 1.5-fold change in Triton-X100 concentration increases the percentage of permeabilized cells by 88 ± 4%. The percentage of viable cells remaining after treatment is directly related to Triton-X100 concentration, with increasing doses of Triton-X100 leading to decreased cell viability. These results demonstrate the importance of determining the minimum effective Triton-X100 concentration for a given application. By using the lowest Triton-X100 concentration that provides sufficient permeabilization efficiency, one can minimize cell loss following treatment. The cell recovery studies suggest that permeabilization can be a reversible process when the appropriate concentration of Triton-X100 is used. The results of the MTT assay further highlight the importance of optimizing Triton-X100 addition, because concentrations of >1.1 pmol/cell lead to a decrease in the mean metabolic activity per cell. Our findings show that minimizing the use of Triton-X100 will reduce both immediate and long-term cell loss. A 10-min treatment with 0.55 pmol/cell Triton-X100, a concentration that facilitates intranuclear labeling with antibodies, allows cells to retain both metabolic and proliferative capacities. Surprisingly, the immediate restoration of membrane integrity does not appear to be a prerequisite for cell function. No lag in the rate of proliferation is observed in the first 24 h after treatment, despite that fact that only 90 ± 8% of metabolically active cells recover full membrane integrity in this time.

All cell lines evaluated here exhibited similar sensitivities to Triton-X100 treatment. The minimum effective concentration required for 3 kDa dextran uptake did not significantly vary between cell lines. All cells showed a dose-dependent response, with the percentage of viable cells remaining after treatment decreasing as the concentration of Triton-X100 was increased. No significant differences were observed in cell viability between cell lines. When the rate of population doubling was evaluated in both squamous cell carcinoma and primary epithelial cell lines, Triton-X100 treatment was found to have no significant effect on either population. On the basis of these results, we believe that Triton-X100 treatment may useful in a range of mammalian cells.
In thinking about potential applications of Triton-X100 mediated permeabilization, it is important to recognize the limitations of this approach. Triton-X100 treatment, even at the low concentrations used here, may have deleterious effects on cells. We observe that antibody labeling of permeabilized cells is specific and reproducible, suggesting that unbound antibodies are removed during cell washing steps. Triton-X100 has been reported to release intracellular proteins and modulate the activity of intracellular enzymes and reorganize membrane domains. These effects, however, do not appear to affect the ability of cells to recover membrane integrity, MTT reduction capacity, or proliferative activity. Triton-X100 cannot be used to target specific cell populations, because all topically treated cells are permeabilized. Thus, Triton-X100 mediated permeabilization will be most useful for applications requiring delivery of contrast agents to large populations of live cells.

Triton-X100 permeabilization holds several advantages over existing intracellular contrast agent delivery techniques. Unlike microinjection, which is only useful for studying small numbers of cells, Triton-X100 can simultaneously deliver both targeted and untargeted contrast agents to any number of cells. Cell loading is more uniform than can be achieved with electroporation; thus, no secondary purification step is required to isolate populations of uniformly loaded cells. Triton-X100 can be used as a pretreatment or coadministered with the macromolecules to be delivered. Contrast agent delivery is not dependent on natural cell uptake mechanisms, allowing cells to be labeled quickly and reproducibly. We believe that Triton-X100, when combined with molecular specific contrast agents, will provide a useful tool for in situ optical studies.

5 Conclusions
In summary, Triton-X100 can be used at subsolubilizing concentrations to permeabilize the cytoplasm and nucleus in a variety of mammalian cells. Targeted and untargeted molecules ranging from 1 to 150 kDa in weight can be reproducibly delivered into adherent cells by controlling the concentration of Triton-X100 relative to the number of cells to be treated. Permeabilization of the cell cytoplasm appears to be a threshold event, whereas nuclear permeabilization is dependent on both macromolecule size and Triton-X100 concentration. When Triton-X100 as used at or near the minimum effective concentration, cell permeabilization is generally reversed in 24 h. Treated cells continue to proliferate and show metabolic activity during the restoration of membrane integrity. Increasing Triton-X100 beyond the minimum effective concentration reduces both short- and long-term cell survival. Different cell lines exhibit similar sensitivities to Triton-X100 treatment, suggesting that Triton-X100 mediated permeabilization will be widely useful. We conclude that Triton-X100 is a promising permeabilization agent for efficient, reproducible, and scalable delivery of optical contrast agents to live mammalian cells.

Acknowledgments
We thank Vivian Mack for assistance with cell culture. This work was supported in part by an NIH BRP Grant (No. CA103830) and a training fellowship from the Keck Center Nanobiology Training Program of the Gulf Coast Consortia (NIH Grant No. 5 T90 DK071054-03).

References


