Efficient mucosal delivery of optical contrast agents using imidazole-modified chitosan

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1 Introduction

Early detection is critical for reducing cancer morbidity and mortality. Optical imaging and spectroscopy can be used to interrogate biochemical and architectural changes associated with early epithelial neoplasia to identify suspicious lesions. Molecular-specific optical contrast agents, capable of selectively labeling biomarkers up-regulated during neoplastic progression, have received much interest for their potential to improve early detection. When interrogated optically, these contrast agents can provide dynamic, real-time information about important molecular hallmarks of neoplasia. Studies of small-molecule optical probes with tissue-permeant properties, such as fluorescent sugar derivatives and nucleic acid dyes, have highlighted the promise of using optical imag-
ing to distinguish molecular changes in small populations of cells.

In recent years, tremendous progress has been made in developing sophisticated, molecularly targeted contrast agents, including antibodies, quantum dots, and various metal nanoparticles (e.g., gold nanoparticles, nanorods, nanoshells, etc.). It has also been demonstrated that targeted gold nanoparticles can provide high contrast between normal and precancerous epithelial tissues. However, their clinical applicability as a topically applied diagnostic agent for early detection of cancer is severely limited, due to our inability to efficiently deliver these agents across mucosal surfaces. Morphological changes associated with precancer generally begin in the basal layers of epithelium; thus, early-detection strategies require contrast agents to be delivered through several hundred microns of epithelial tissue. It is important that these agents be delivered efficiently and uniformly throughout the region of interest, ensuring that they reach and bind to their targets. In addition, unbound agents should also be easily “washed out” to reduce nonspecific signaling and false-positive detection. The penetration of molecules through mucosal tissue depends in part on their size and generally decreases exponentially with increasing hydrodynamic volume. Chemical modification and encapsulation strategies to improve tissue penetration have proven useful only for molecules up to 6 kDa in size.

Permeation enhancers (also called penetration or absorption enhancers) have been investigated to facilitate the delivery of larger molecules. These are typically applied substances that disrupt epithelial tissue by various mechanisms to increase paracellular (between cell) and/or transcellular (across cell) transport. Numerous permeation enhancers have demonstrated efficacy in skin, allowing transdermal penetration of drugs, macromolecules, and nano- and microparticles as large as 1 to 5 μm. However, studies utilizing similar formulations in the mucosal epithelium have enhanced the penetration only of small molecules, including insulin (6 kDa), calcitonin (3.5 kDa), and dextrans up to 70 kDa in weight. The surfactant Triton-X100 has recently demonstrated reversible mucosal permeation enhancement for efficient delivery of molecules up to 150 kDa in size. Nevertheless, the slow rate of tissue recovery associated with Triton-X100 limits the clinical feasibility of this approach, as morphological and histochemical analysis of tissues following treatment with the permeabilizing agent is not highly practical.

Chitosan and its analogs have been studied extensively for in vivo delivery of DNA and siRNA via mucosal delivery routes. Chitosan is a biocompatible, biodegradable, cationic polysaccharide comprised of repeating units of glucosamine and N-acetyl glucosamine. It has been shown to reversibly disrupt epithelial tight junctions to allow the paracellular transport of small molecules. Topical application of chitosan has been shown to enhance drug uptake in nasal, buccal, and intestinal epithelia, and its bioactivity is mediated by its cationic properties. However, a major drawback of chitosan for clinical use has been its limited solubility at physiological pH and above, which also leads to loss of charge. To improve these characteristics and enhance its permeation properties, we developed a molecular analog in which the primary amines are partially modified with 4-imidazole acetic acid monohydrochloride (henceforth referred to as chitosan-IAA).

In this paper, we describe the first use of chitosan-IAA to deliver tissue-impermeant optical contrast agents through the epithelium of freshly resected mucosal tissues. Topically applied, untargeted macromolecules and nanoparticles were used as probes to assess mucosal permeability following a 15-min pretreatment with chitosan-IAA. Contrast agent localization within the epithelium was interrogated using confocal microscopy to determine the route of transepithelial delivery. The dynamic recovery of barrier function after permeation treatment was assessed using untargeted probes of different sizes. Fluorescently labeled antibodies targeted to epidermal growth factor receptor (EGFR) were delivered into biopsies collected from patients with oral cancer and pretreated with chitosan-IAA to evaluate the clinical potential of this approach. The results of these studies demonstrate that chitosan-IAA reversibly enhances mucosal permeability in a rapid, reproducible manner sufficient to facilitate the transepithelial delivery of optical contrast agents up to 44 nm in diameter.

2 Materials and Methods

2.1 Chemical Modification of Chitosan

Chitosan was chemically modified to improve its solubility and tissue permeation capacity. The conjugation of 4-imidazole acetic acid monohydrochloride (IAA) to the primary amines of chitosan for production of modified chitosan-IAA was performed as previously described. Briefly, 86% deacetylated chitosan of 130 kDa molecular weight (PCL 113, Novamatrix, Norway) and IAA (Acros Organics, Morris Plains, New Jersey) were dissolved in 0.1 M 2-(N-morpholino)ethanesulfonic acid (pH 6.5, Sigma-Aldrich, St. Louis, Missouri) to a concentration of 1.0% and 2.0% w/v, respectively. A theoretical modification of 50% was targeted for the reaction. IAA and chitosan were combined at a volumetric ratio of 1:5 while on ice. 20 M excess (in relation to IAA) of 1-ethyl-3-(3-dimethylaminopropyl carbodiimide hydrochloride (Pierce Biotechnology, Rockford, Illinois) was added and immediately vortexed to promote addition of IAA to the chitosan backbone. Reactions were allowed to continue overnight with end-over-end mixing. Solutions were dialyzed for 2 days against 5 mM hydrochloric acid (Acros Organics) thrice and then deionized water thrice using Snakeskin pleated dialysis tubing (10,000 MW cut-off; Pierce Biotechnology, Inc.). Following dialysis, the samples were lyophilized for 24 h. Degree of substitution was determined with via 1H NMR. Chitosan and chitosan-IAA were resuspended at a stock concentration of 1% w/v in 0.2 M sodium acetate (pH 4.5, Sigma Aldrich) immediately prior to dilution for tissue application.

2.2 Synthesis and Validation of Gold Nanoparticles

Gold nanoparticles were synthesized as untargeted tissue permeability probes. Gold spheres of 9.5 ± 0.35 nm and
20.7 ± 0.26 nm hydrodynamic diameter were synthesized by a citrate reduction of tetrachloroauric (III) acid (HAuCl₄, Sigma-Aldrich, St. Louis, Missouri) under reflux, as previously described. Nanoparticle size was confirmed by dynamic light scattering (DLS) with a ZetaPlus system (Brookhaven Instruments Corporation, Holtsville, New York). Each set of gold nanoparticles were coated with a fluorescent 5 kDa polyethylene glycol (PEG) to reduce their aggregation in media. Briefly, 0.1% w/v fluorescein-PEG-amine (Laysan Bio, Arab, Alabama) was reacted with 2 M excess of 2-iminothiolane (Pierce Biotechnology) for 2 h. Upon completion, the solution was filtered at room temperature with Amicon Ultra centrifugation filters (3000 MWCO; Millipore, Billerica, Massachusetts) at 3220 × g for up to 60 min. Gold nanoparticles, suspended at a concentration of 7 × 10¹⁰ particles/ml (determined by spectral absorbance analysis) were allowed to react with 10⁻⁴ M fluorescein-PEG-thiol at a volumetric ratio of 5:1 for 45 min on a shaker for each set of nanoparticles. PEG (Sigma-Aldrich) was subsequently added to a final concentration of 2% w/v. The particles were centrifuged at 1500 × g for 30 min and resuspended in deionized water. Gold coating of nanoparticles was confirmed by monitoring the ~494-nm absorbance peak (absorption maxima of fluorescein on our PEG) using a Synergy HT UV-Vis spectrophotometer (Biotek, Winooski, Vermont). The final hydrodynamic radius of the gold nanoparticles was measured to be 32.9 ± 1.27 nm and 43.9 ± 0.51 nm using DLS, demonstrating similar size increase following PEGylation as was previously described.

2.3 Synthesis and Validation of Fluorescent Macromolecules

Fluorescent macromolecules were prepared as targeted and untargeted tissue permeability probes. These contrast agents were selected based on size, shape, and charge to provide comparison to commonly known and utilized molecular contrast agents. 3 kDa rhodamine-dextran, 40 kDa fluorescein-dextran, and 150 kDa AlexaFluor 647-IgG were purchased from Invitrogen (Carlsbad, California) for use as untargeted probes. For targeted labeling studies, mouse antihuman antibodies specific for epidermal growth factor receptor (EGFR; clone 108; custom synthesized by the Baylor College of Medicine, Houston, Texas) were reacted with Alexa Fluor 464 carboxylic acid succinimidyl esters using commercially available labeling kits (Invitrogen). The purified conjugates were suspended in PBS at a concentration of 1.0 mg/ml. Dye-labeled isotype controls were synthesized at the same concentrations. Prior to tissue labeling, the bioactivity and specificity of the conjugates was confirmed using live EGFR-positive 1483 cells and EGFR-negative MDA-MB-435S cells as described in Ref. 54 in both the presence and absence of chitosan and chitosan-IAA.

2.4 Topical Permeation of Fresh Bladder Tissue

Guinea pig bladder mucosa was used as a model tissue to evaluate tissue permeability in the presence and absence of chitosan-based permeation enhancers because of the folding nature of the tissue. The guinea pig bladder allows for effective imaging of both epithelial and stromal layers of the tissue at once. All animals were cared for in accordance with institutional guidelines. The protocols were reviewed and approved by the IACUC at Rice University. This model has been previously validated for the study of topical permeation enhancement. Whole bladders were excised from 3- to 4-week-old female Hartley guinea pigs (Charles River Laboratories, Wilmington, Massachusetts) directly following animal sacrifice. The bladders were sectioned into 8 to 10 pieces of uniform surface area (3 × 3 mm) with a scalpel and washed once in PBS and then immediately used for permeation studies. Two topical permeation enhancers, chitosan and chitosan-IAA, diluted to 0.01% w/v in DMEM/F12 medium (Invitrogen), were evaluated for their ability to increase tissue permeability. Media alone was used as a negative control. 100 µL of permeation-enhancing solution was topically applied to the apical surface of each tissue biopsy for 15 min.

2.5 Assessment of Transepithelial Macromolecule Delivery

To determine whether chitosan-treated tissues are selectively permeable to macromolecules of specific sizes, fluorescent macromolecules ranging from 3 kDa to 150 kDa in size were topically applied to the tissue surfaces of bladder biopsies. Tissue penetration was then monitored optically. The apical surface of the epithelium was topically treated for 15 min with prewarmed (37 °C) 0.01% w/v chitosan or chitosan-IAA, washed once in cold media, and covered with a 1:1 mixture of 3 kDa rhodamine-dextran, 40 kDa fluorescein-dextran, and 150 kDa AlexaFluor 647-IgG, each diluted to a concentration of 1 µM in cold media. Tissues were immersed with this solution for 15 min at 4 °C and then imaged using confocal microscopy at three different excitation wavelengths (described in the following). Images were collected in 2- to 5-µm steps from the surface into the tissue. Following imaging, the tissues were washed three times in cold media (15 min total) and reimaged to assess the removal of unbound macromolecules. Experiments were repeated with chitosan-IAA treatment at 25 °C or 4 °C to determine the influence of temperature on tissue permeation. Each labeling condition was evaluated in six independent experiments.

2.6 Assessment of Transepithelial Nanoparticle Delivery

To determine whether chitosan-treated tissues permit the transepithelial delivery of nanoparticles, nanoparticles of different sizes were topically applied to the surface of permeabilized bladder biopsies. Nanoparticle delivery was then monitored optically. Briefly, the apical surface of the epithelium was topically treated for 15 min with prewarmed (37 °C) 0.01% w/v chitosan or chitosan-IAA diluted in DMEM/F12 medium, washed once in cold media, and covered with fluorescein-PEG-gold spheres of 33 or 44 nm diameter, each diluted to a concentration of 1 × 10⁶ nanoparticles/ml in cold media. Tissues were immersed with this solution for 15 min at 4 °C and then imaged using confocal microscopy. Images were collected in 2- to 5-µm steps from the surface into the tissue. Following imaging, the tissues were washed three times in cold media (15 min total) and reimaged to assess the removal of unbound nanoparticles. Each labeling condition was evaluated in six independent experiments.
2.7 Time-Course Analysis of Tissue Recovery
To monitor the recovery of bladder epithelial barrier function following the removal of chitosan-IAA, fresh bladder biopsies were treated with 0.01% w/v chitosan-IAA or media for 15 min at 37 °C, washed, and probed with nontargeted contrast agents at regular time intervals during tissue recovery. Briefly, chitosan-IAA and media-treated samples were washed three times with warm or cold media and then allowed to recover in media at 37 °C or 4 °C. At 0, 15, 30, 60, 90, or 120 min after chitosan-IAA removal, the samples were placed on ice and topically labeled with a 1:1 mixture of untargeted fluorescent macromolecules (3 kDa dextran, 40 kDa dextran, and IgG antibody) or 44-nm gold nanoparticles. Samples were allowed to incubate with the macromolecules or nanoparticles for 15 min prior to imaging via confocal microscopy. Images were collected 40 μm below the tissue surface to allow for optical sectioning of both the epithelium and underlying stroma. Tissue samples were assessed in duplicate using a minimum of three independent experiments.

2.8 Molecular-Specific Labeling of Human Oral Biopsies
To demonstrate the feasibility of delivering molecular-specific optical contrast agents targeted against biomarkers of neoplasia, human oral biopsies were co-treated with fluorescent EGFR antibodies and chitosan-IAA. Briefly, paired clinically normal and abnormal biopsies of the oral mucosa were obtained from five consenting patients at the University of Texas M.D. Anderson Cancer Center (MDACC). Patients gave written informed consent, and the clinical protocols were approved by the Institutional Review Boards at MDACC and Rice University. Biopsies were immediately placed and remained in chilled media until they arrived at Rice University. The biopsies were embedded vertically into 3% w/v ultrapure agarose (Invitrogen) to prevent the influx of permeation enhancers and contrast agents at the tissue margins, as previously described. The apical surface of the epithelium was left free of agarose to facilitate topical labeling. Based on the increased number of epithelial layers in human oral resections, tissues were topically treated with 1 μM antiEGFR-647 diluted in 0.01% w/v chitosan-IAA for 60 min at 37 °C, washed three times with cold media, and then sliced transversely into 200-μm-thick slices using a Krumdieck tissue slicer. Tissue slices were counterstained with 0.01% proflavine, a nucleic acid dye, and immediately imaged using confocal microscopy.

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 photo multiplier tube (PMT) detectors and Zeiss LSM 5 image examiner software. Samples were sequentially excited with each laser line with power settings held constant for each laser. Fluorescence emission was collected using 500- to 530-nm (40 kDa fluorescein-dextran), 565- to 615-nm (3 kDa rhodamine-dextran), and 650- to 710-nm (150 kDa Alexa647-IgG) bandpass filters. Tissue reflectance at 633 nm was collected using a 635-nm dichroic beamsplitter. Images were acquired at 0.5 frames per second using a 20× objective with a pinhole of 2.56 Airy units. For the untargeted macromolecule and nanoparticle permeation studies, the gain was held constant with excitation at 488, 543, and 633 nm. At the gains utilized, the fluorescence of solutions outside the tissue was oversaturated, but the fluorescence of solutions within the tissue was not. In the EGFR targeting studies, the gain was held constant for antiEGFR-647 imaging (633-nm excitation) and proflavine imaging (488-nm excitation).

2.10 Image Analysis
To quantify the fluorescence intensity of the stroma as a function of recovery time and temperature, representative confocal fluorescence images were analyzed using Image J v1.38 (NIH, Bethesda, Maryland). The mean fluorescence intensity per unit area was determined by selecting regions of interest (ROIs) within each image and then dividing the measured fluorescence intensity by the area of the ROI. The ROI margins were drawn just inside the stroma border as determined from reflectance overlay images. The entire stroma was included for each image. When images contained more than one region of discontinuous stroma, the mean intensity for the regions was averaged and treated as a single value. Background noise was determined at each time-point from samples pretreated with media in place of chitosan-IAA. The normalized mean fluorescence intensity of the stroma was calculated by subtracting the measured background and normalizing for the fluorescence intensity of the stroma at time of chitosan-IAA removal. The normalized mean fluorescence intensity of the stroma was assessed for each macromolecule using five representative images from three independent experiments (15 images total per macromolecule per time-point and temperature).

The mean reflectance intensity per unit area of the stroma in nanoparticle labeling studies was quantified in the same manner as described for the fluorescence studies, using five representative images from five independent experiments (25 images total per labeling condition).

To quantify the fluorescence intensity of EGFR labeling in transverse images in human oral biopsies, representative confocal fluorescence images were analyzed. In normal samples, ROIs were bounded by the apical and basal surfaces of the epithelium, thereby including any labeling throughout the epithelium. In neoplastic samples, ROIs were drawn to include all squamous cells from the surface of the tissue to the lower margin of EGFR labeling. Regions of fibroblasts, as identified morphologically by proflavine counterstaining, were excluded from the analyses. The mean fluorescence intensity per unit area was assessed using 15 images per biopsy, consisting of three representative images from five separate tissue slices. No background corrections were made because the measured background was <1% of the measured mean EGFR labeling intensity for all images evaluated. Differences in mean labeling intensity between normal and neoplastic samples were assessed on a patient-to-patient basis using a two-tailed, unpaired Student’s t-test, with p-values of ≤0.01 being considered statistically significant. The relative contrast ratio was determined by dividing the mean fluorescence intensity of each cancer biopsy by that of the contralateral normal.
3 Results

3.1 Synthesis and Characterization of Imidazole-Modified Chitosan

A schematic for the synthesis of chitosan-IAA is shown in Fig. 1. We successfully introduced imidazole acetic acid to the primary amines of chitosan via a carbodiimide-mediated reaction. Following synthesis, $^1$H NMR characterization of the purified polymer provided on average a degree of substitution of about 3.0% of the primary amines modified for each chitosan molecule. Chitosan-IAA polysaccharide prepared through this type of reaction has been previously shown to provide enhanced polymer solubility and buffering capacity.

3.2 Transepithelial Macromolecule Delivery through Fresh Bladder Mucosa

Figure 2 shows representative confocal fluorescence images of bladder tissue treated topically with 0.01% w/v chitosan-IAA, chitosan, or media for 15 min at 37 °C followed by topical application of a 1:1:1 mixture of 3 kDa rhodamine-dextran, 40 kDa fluorescein-dextran, and 150 kDa Alexa647-IgG. The tissue reflectance images are shown on the left, and the corresponding fluorescence images are shown to the right. The yellow/white lines indicate the boundary between the stroma and the epithelium. Due to the three-dimensional (3-D) folding of the resected bladder, it was possible to image the epithelium in cross section using confocal microscopy at a depth of 40 μm. In the reflectance images, the epithelium was distinguished from the stroma by its darker appearance. Both chitosan and chitosan-IAA were found to facilitate the transepithelial delivery of macromolecules. Macromolecules of all three sizes accumulated in the stroma of permeation-enhanced tissues, causing the stroma to appear brighter than the epithelium in the fluorescence images. Chitosan-IAA treatment facilitated more intense stromal accumulation than nonmodified chitosan for all sizes of macromolecules evaluated. The macromolecules could be removed with several brief washes in media (data not shown), demonstrating that macromolecule accumulation is reversible. Experiments were independently repeated by labeling tissues with one macromolecule at a time (data not shown) to exclude macromolecule interactions. In media-treated controls, the macromolecule penetration was limited to the superficial epithelium. No significant macromolecule penetration was observed following chitosan-IAA or chitosan treatment at 25 °C or 4 °C (data not shown).

3.3 Transepithelial Nanoparticle Delivery through Fresh Bladder Mucosa

Figure 3 shows representative confocal fluorescence images of bladder tissue treated topically with 0.01% w/v chitosan-IAA, chitosan, or media for 15 min 37 °C and then probed with fluorescein-PEG-gold spheres of 44 or 33 nm diameter. Both chitosan and chitosan-IAA were found to facilitate the transepithelial delivery of nanoparticles. Following nanoparticle application, fluorescence was primarily observed in the stroma of permeable tissues. Stromal labeling was generally less uniform than observed with macromolecules. Pretreatment with chitosan-IAA resulted in more intense stromal accumulation than nonmodified chitosan. This difference was more obvious following application of larger nanoparticles. The stromal reflectance was visibly enhanced in samples demonstrating nanoparticle uptake. Tissues exposed to nanoparticles showed a stromal reflectance enhancement of 1.5 to 1.7 (44 nm gold) and 1.7 to 2.1 (33 nm gold) compared to tissues treated with media in place of chitosan following quantifica-
tion of reflectance at a fixed gain and regions of the stroma. Minimal to no nanoparticle accumulation was observed in media-treated controls.

3.4 Route of Contrast Agent Delivery Following Chitosan-IAA Treatment

The route of contrast agent delivery through the epithelium was optically interrogated in fresh bladder tissues pretreated with chitosan-IAA. Confocal fluorescence images were collected in 2-μm steps from the surface of the tissue following the topical application of 3 kDa rhodamine-dextran or 44 nm fluorescein-PEG-gold. Representative videos are available online as Video 1 (3 kDa dextran) and Video 2 (44 nm gold). The videos start at 2 μm below the surface and advance at a rate of 2 μm/s. The stills shown for Video 1 and Video 2 represent frames collected at 10 μm below the tissue surface. 3 kDa rhodamine-dextran labeling, shown in Video 1, was characterized by ringlike fluorescence surrounding each cell in the field of view. The labeling appeared extracellular, suggesting a paracellular route of dextran delivery. These rings became progressively smaller in diameter with increasing depth in the epithelium, correlating well with known bladder epithelium morphology. In contrast, the delivery of larger paracellularly and transcellular routes. Fluorescein-PEG-gold delivered in a paracellular manner appeared as ringlike labeling that became progressively smaller with increasing depth. Fluorescence associated with nanoparticles was also observed in the cytoplasm of cells across the full thickness of the epithelium, suggestive of transcellular transport. Individual nuclei appeared as a black spot within each cell, suggesting that the transcellular movement of nanoparticles was limited to the cytoplasmic compartment. In both samples, the transition from epite-
Fluorescence intensity as a function of time and temperature. Tissues pretreated with chitosan-IAA were washed and allowed to recover at 37 °C (left) or 4 °C (right). The recovery of barrier function was probed at regular time intervals using fluorescent macromolecules and nanoparticles. Tissues held at 37 °C rapidly recovered barrier function, while tissues held at 4 °C displayed continued permeability as evidenced by stromal fluorescence. The ability of the recovering epithelium to block contrast agents was size dependent, with larger molecules excluded more rapidly than smaller molecules.

Epithelial Recovery Following Chitosan-IAA Treatment

Figure 5 shows tissue permeability as a function of time and temperature following the removal of chitosan-IAA. Tissue recovery at 37 °C is illustrated on the left, and tissue recovery at 4 °C is illustrated on the right. Samples were pretreated with 0.01% chitosan-IAA for 15 min and washed in warm or cold media, and then macromolecules or nanoparticles were applied topically at different time-points. The mean fluorescence intensity of the stroma was measured at each time-point with larger molecules excluded more rapidly than smaller molecules.

Epithelial permeability for all sizes for up to 2h was characterized by a relative increase in fluorescence. Dextran and nanoparticle accumulation in the stroma was generally uniform. Few morphological features were defined in stroma, other than blood vessels of horizontal and vertical orientation (see Video 1). These vessels were characterized by low-contrast agent accumulation, appearing as dark stripes and circles/ovals, respectively. Similar routes of transport for these two contrast agent sizes were seen following pretreatment of tissues with nonmodified chitosan (data not shown).

Delivery of Targeted Contrast Agents for Cancer Detection

To demonstrate the clinical potential of chitosan-IAA to aid in delivery of targeted optical contrast agents for the early detection of cancer biomarkers, paired human oral biopsies were topicaly labeled with fluorescent EGFR-specific antibodies diluted in 0.01% w/v chitosan-IAA. We have previously demonstrated that antibodies targeted to EGFR lack the ability to penetrate normal oral mucosa and squamous carcinomas.

EGFR labeling in normal epithelium was limited to the basal layer of epithelial cells, appearing as a faint stripe above the epithelium. EGFR labeling in squamous cell carcinoma tissue was characterized by bright, ring-like extracellular labeling extending downward from the tissue surface. The scale bar represents 100 μm.

3.5 Epithelial Recovery Following Chitosan-IAA Treatment

Figure 5 shows representative confocal images of antiEGFR-647 labeling in fresh human oral biopsies co-treated with chitosan-IAA, collected at the same gain. Biopsies were topicaly treated with antiEGFR-647 diluted in 0.01% w/v chitosan-IAA for 1 h at 37 °C, washed three times, sliced transversely, and imaged. In normal tissues (top), EGFR labeling is limited to the basal layer of epithelial cells, appearing as a faint region of labeling at the base of the epithelium. Nonspecific labeling is observed at the apical surface of keratinized tissues such as the retromolar trigone tissue shown below. EGFR labeling in squamous cell carcinoma (bottom) is characterized by bright, ring-like extracellular labeling extending downward from the tissue surface. The scale bar represents 100 μm.

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Figure 6 shows representative transverse sections of normal (top) and cancerous (bottom) gingiva tissue labeled for EGFR and counterstained with the nucleic acid dye proflavine. Labeling with antiEGFR-647 is displayed in red and proflavine in green. Differential EGFR labeling was observed between normal and cancer samples from all five patients evaluated. EGFR labeling in normal epithelium was limited to cells in the basal layer. This labeling was observed in all normal samples evaluated, and was generally less intense and more diffuse than labeling in cancer samples. Nonspecific antibody labeling was observed in keratinized tissues, appearing as a bright stripe above the epithelium. EGFR labeling in squamous cell carcinoma tissue was characterized by intense, ring-like labeling extending downward from the tissue surface. This labeling appeared highly uniform, labeling all squamous cells within the treatment zone. Regions of fibroblasts, which were identified by the presence of small, widely spaced nuclei using proflavine staining, did not show significant EGFR labeling. With 1 h of chitosan-IAA treatment, antibody labeling reached a depth of approximately 150 to 250 μm in cancer tissue and ≤400 μm in normal epithelium.

Figure 6 shows the mean fluorescence intensity of paired oral biopsies collected from five patients. The biopsy pairs were labeled and imaged as described earlier. The mean fluorescence intensity of EGFR labeling in normal biopsies was generally low and showed little variation from patient to pa-
The mechanism by which chitosan and its analogs increase tissue permeability remains poorly understood. Experiments using human intestinal Caco-2 cell monolayers have suggested that the mechanism of action, which appears to be mediated by positive charges on the chitosan, includes interactions with tight junction proteins, the redistribution of F-actin, and a destabilization of the cell membrane. Our studies support the observation that this is an active, energy-dependent process, since no transepithelial contrast agent delivery was observed following permeation treatment at 25 °C or 4 °C. Similarly, the recovery of barrier function appears to be an active process, requiring incubation at 37 °C after washing of the tissue. Permeation of bladder mucosa is rapid, with a 15-min treatment sufficient to facilitate contrast agent transport through several layers of epithelial cells.

The temperature-dependent nature of chitosan-IAA treatment provides a unique opportunity for the study of this process, since the gain and loss of tissue permeability can be halted simply by placing the tissue on ice. Using this approach, we were able to measure size-dependent differences in agent exclusion. Tissues chilled at various time-points during the recovery process preferentially excluded larger contrast agents more rapidly than smaller contrast agents. The ability of the treated tissues to block transepithelial agent delivery increased progressively. Interestingly, in cell monolayers, chitosan concentrations of ≥0.01% have been shown to irreversibly block tight junction recovery. In contrast, three-dimensional (3-D) tissues washed after treatment with chitosan-IAA show evidence of recovery of the epithelial barrier function. This washing step is important for the rapid reversal of epithelial permeability; however, we have yet to determine whether washing physically removes chitosan-IAA or introduces new divalent cations to compete with chitosan-IAA binding sites.

The rapid reversibility of chitosan-IAA mediated permeation supports the clinical potential of this approach. There are concerns that topical permeation enhancers, which by nature interfere with the barrier function of the epithelium, could potentially permit the entry of toxins and/or allergens. Based on our results, we would expect the penetration of these agents to be blocked rapidly. Compared to the topical permeation enhancer Triton-X100 studied in the same tissue model, which has also shown promise for the mucosal delivery of macromolecular optical contrast agents, tissue recovery following chitosan-IAA treatment is much more rapid. Optimization of the treatment concentration, duration, and removal may be used to further improve the rate of tissue recovery.

The factors that determine the route of contrast agent penetration need to be further clarified. Chitosan solutions have been shown to increase both the paracellular and transcellular permeability of Caco-2 cell monolayers in a reversible, dose-dependent manner. We observed contrast agent uptake in tissues via both these routes. Contrast agents up to 150 kDa in size appeared to follow primarily a paracellular route, while nanoparticles followed both paracellular and transcellular routes. The observed paracellular transport likely resulted from an opening of the tight junctions. The observed transcellular transport was likely a consequence of cell membrane destabilization. Both of these mechanisms have been described in Caco-2 monolayers following chitosan treatment. All contrast agents evaluated here were essen-
tially neutral in charge (data not shown), suggesting that the shift toward a transcellular route may be determined by probe size. The formulation of chitosan into nanoparticle-drug complexes has been demonstrated to enhance the internalization of chitosan in cells and tissues, presenting a second potential approach for driving transcellular delivery. At this time, it is not yet clear whether using chitosan and its analogs as a nanoparticle or nanoparticle coating will be more efficient than chitosan free in solution. From the point of contrast agent delivery, the ability to selectively control the delivery route would be useful for distinguishing between true intracellular labeling and endocytic uptake.

Controlled, uniform delivery of molecular-specific contrast agents across the full thickness of the epithelium remains an important goal for design of early detection strategies. One-time topical application of 0.01% chitosan-IAA facilitates the permeation enhancement in a variety of tissues, including the transitional epithelium of the bladder, squamous epithelium of the oral cavity, and squamous cell carcinoma. Delivery of fluorescent EGFR-specific antibodies was observed across the full thickness of the epithelium (150 to 400 μm) in all normal oral biopsies. Differential labeling was observed between normal and cancer specimens, suggesting that the specificity of the antibody and the structure of the target are conserved in normal and cancer specimens, suggesting that the specificity of chitosan-IAA treatment will be important for understanding and controlling the 3-D spread of tissue permeation. These margins will need to be distinguished from the labeling margins in order to determine how to further improve the depth of labeling in cancer tissue. The selection of an appropriate animal model will be critical for optimizing the efficacy of chitosan-IAA treatment at doses that can be scaled for clinical use.

The successful translation of chitosan-IAA for in vivo work will require characterizing the safety, toxicity, biological activity, and reversibility of chitosan-IAA mediated permeation at clinically relevant doses. Potential in vivo application would include topical application of chitosan-IAA and contrast agents via gauze pad or similar devices or use of a spray catheter for endoscopically accessible tissues (e.g., gastrointestinal tract and cervix). Chitosan is currently FDA approved as a generally regarded as safe, or GRAS, compound for use as a hemostatic bandage. Drug delivery studies utilizing chitosan derivatives in animal models have demonstrated minimal cytotoxic effects.

Similar studies will be needed to monitor the safety of chitosan-IAA both locally and systemically at regular time intervals following topical treatment. Given the size of the macromolecules and nanoparticles that have been shown to penetrate the tissue following chitosan or chitosan-IAA treatment, study of the potential entry of small viruses should also be analyzed as a potential drawback for in vivo use of the permeation enhancers. Carcinogen-treated or orthotopic transplant models will be useful for establishing the sensitivity and specificity of contrast agents delivered with clinically relevant chitosan-IAA doses. Furthermore, studies for understanding the actual mechanism by which chitosan and chitosan-IAA enhance macromolecular transport are needed. Additional studies are also needed to determine the limitations of chitosan-IAA as a function of concentration, pH, and treatment duration.

5 Conclusions
In conclusion, this paper demonstrates that chitosan-IAA is a promising topical permeation enhancer for the mucosal delivery of tissue-impermeant optical contrast agents. Compared to other mucosal permeation enhancers, chitosan and chitosan-IAA can facilitate delivery of a broader size range of molecules. No chemical modification of optical contrast agents is needed. Nanoparticles as large as 44 nm can be delivered following a one-time permeation treatment. Chitosan-IAA, with its improved solubility, appears to be the more effective permeation enhancer, particularly for larger contrast agents. Importantly, permeation of the epithelium can be reversed by washing, leading to a progressive reduction in macromolecule penetration. Recognition that this is an active process provides new avenues for elucidating the mechanism of chitosan and its analogs, since the gain and loss of permeability can be halted simply by placing the treated tissue on ice. The co-administration of chitosan-IAA with molecular-specific contrast agents facilitates differential labeling between normal and cancerous tissues, supporting the clinical potential of this approach. Further work is needed to establish the safety of chitosan-IAA for in vivo use.

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