Optical coherence tomography (OCT) is an evolving imaging modality that combines interferometry with low-coherence light to produce high-resolution tissue imaging. Cross-sectional in vivo images were obtained using an OCT device consisting of a Michelson interferometer, 1.3-µm broadband light source, and a handheld fiberoptic imaging probe. Image pixel resolution approached 10 µm. The mucosa of the oral cavity and oropharynx were examined in 41 patients during operative endoscopy. Optical coherence tomographic imaging was combined with endoscopic photography for gross and histologic image correlation. Optical coherence tomographic images of the oral cavity and oropharynx provided microanatomical information on the epithelium, basement membrane (BM), and supporting lamina propria (LP) of the mucosa. Normal microstructures identified in these tissues included an overlying keratin layer, papillae, ducts, glands, and blood vessels. Regions of pathologic features studied included mature scar, granulation tissue, mucous cysts, leukoplakia, and invasive cancer. Optical coherence tomographic imaging showed distinct zones of normal, altered, and ablated tissue microstructures for each pathologic process studied. Abnormal findings were directly compared with regions of normal tissue or conventional histopathologic features when tissue for analysis was available. This study provides a composite series of in vivo OCT images of the oral cavity and oropharynx in a variety of normal regions and pathologic states as well as outline future applications of OCT technology.
coherence light source with a Michelson interferometer to produce cross-sectional images of tissue structures. Conceptually, OCT is analogous to B-mode ultrasonography but uses broadband near-infrared light, rather than sound energy, to produce images with a resolution approaching that of light microscopy (10-15 µm).5,6 Optical coherence tomography images differences in the tissue optical properties, which includes the effects of both optical absorption and scattering. Tomographic systems are used in ophthalmology and dermatology, but in vivo study of OCT in the human oral cavity and oropharynx has been limited.7,8

In this study, we used a multipurpose OCT system for imaging the oral cavity and oropharynx during surgical endoscopy and in the clinical outpatient setting with the primary objectives of characterizing tissue microstructures and evaluating the efficacy of OCT to image the surface and submucosal tissues.

**METHODS**

**PATIENTS**

Optical coherence tomographic imaging was performed on patients undergoing oral examination in the outpatient clinic and during surgical endoscopy at the University of California Irvine Medical Center under protocol approved by the Human Subjects Institutional Review Board at the University of California Irvine, Orange. While the patient was under general anesthesia, OCT imaging was performed with a handheld probe positioned manually or with endoscopic guidance. Digital photographs of imaged areas were obtained and, when clinically indicated, correlative biopsy specimens were obtained.

**OCT SYSTEM AND IMAGING**

The components of our fiber-based OCT system are schematically illustrated in Figure 1. Near-infrared light from a broadband light source (central wavelength \( \lambda = 1310 \) nm, FWHM [full

---

**Figure 1.** Schematic of optical coherence tomographic (OCT) imaging system and oral cavity/oropharynx target. The OCT system parameters are as follows: \( \lambda = 1310 \) nm and full width half maximum = 80 nm; lateral resolution = 15 µm; axial resolution = 10 µm; frame rate: 1 to 2 Hz. A/D means analog to digital conversion; DSP indicates digital signal processing; and RSOD, Rapid Scanning Optical Delayline.

---

**Figure 2.** View of optical coherence tomographic handheld probe.

---

width at half maximum] \( \Delta \lambda = 80 \) nm, AFC BT 1310; JDS Uniphase, San Jose, Calif) enters a \( 2 \times 2 \) fiber coupler. In the reference arm, a rapid scanning optical delay line attains A-scan at 500 Hz without phase modulation. A phase modulator generates 500-kHz phase modulation for heterodyne detection. Signals backscattered from the sample arm are obtained by phase-resolved processing with the interference fringes. The lateral and axial resolution of the system is approximately 10 µm per pixel. The horizontal image window was set laterally from 2 to 10 mm in length while detailed images of tissue microstructure were recorded up to a depth of 1.6 mm, depending on the turbidity of the media.

To image the oral cavity and oropharynx in vivo, a custom flexible OCT probe was designed to accommodate specific anatomical structures (Figure 2). The probe consists of a 900-µm, single-mode, 1310-nm OCT imaging fiber distally mounted with a gradient refractive index lens and a 0.7-mm right-angle prism (Figure 3). The gradient refractive index lens is 1.00 mm in diameter, has a 0.23 pitch, and works as a focuser. Mounting of the prism and gradient refractive index lens is performed with an optics grade, low-viscosity, wicking UV glue.
Scanning is achieved by linearly translating the fiber optic along the long axis using a motorized stage (model 663.4pr; PI Line, Tustin, Calif). The fiber optic and optical elements are enclosed by a transparent plastic tube (1.95-mm inner diameter, 0.21-mm thickness, fluorinated ethylene propylene material), which is mechanically supported and protected by a secondary outer stainless steel cylinder. To help orient the user, colored markings were made along the sides of the optic fiber and opposite light exit along the fiber tip.

The probe design was optimized during clinical evaluation to decrease bulk, improve acquisition speed, and optimize image quality. The current design was used for more than 95% of cases reported in this study. Image acquisition was monitored in real time on a 17-in LCD (liquid crystal display) monitor placed alongside the endoscopy video monitor. The entire OCT system is contained in a mobile endoscopy tower.

**OCT IMAGE ANALYSIS**

Images from multiple sites in the oral cavity and oropharynx were obtained. Acquired images were then sorted into the following 4 categories: normal healthy tissues, benign pathologic features (nonmalignant and precancerous lesions), invasive cancer, and transition zones at the borders of benign and malignant lesions. A breakdown of the images obtained at each anatomical site is tabulated in Table 1 and Table 2.

**HISTOLOGY**

In cases in which biopsy specimens were obtained, histopathologic features were compared with OCT images to assess epithelial thickness, basement membrane (BM) integrity, and characterization of tissue microstructures within the LP. The OCT images were then used to construct a database of normative in vivo tissue microanatomy and pathology.

**RESULTS**

Forty-one adult subjects ranging in age from 24 to 89 years participated in an OCT study of the oral cavity and oropharynx. Optical coherence tomographic images were collected from multiple sites during operative endoscopy in 34 subjects and in the outpatient clinic in an additional 7 subjects. When clinically indicated, histologic biopsy specimens were obtained and compared with OCT images and lesion photographs. In cases in which no biopsy was indicated, comparison of OCT images was made using standard histologic criteria. The distribution of images obtained is displayed in Table 1. Images were collected, and categorized, first separating normal anatomical regions into oral cavity and oropharynx. The images were then organized by diagnosis into categories of inflammation, cyst, acanthosis, hyperplasia, hyperkeratosis, parakeratosis, fibrosis, granulation tissue, metaplasia, dysplasia, ulceration, carcinoma in situ, verrucous carcinoma, and squamous cell carcinoma.
ical structures from pathologic lesions. Images from clinically evident pathologic lesions were subdivided into benign processes, premalignant lesions, and malignant lesions. Attention was also given to images demonstrating transitions from pathologic tissue changes and normal histopathology. Multiple benign and malignant pathologic lesions were identified, and are tabulated in Table 2.

The following tomographic series represents a survey of tissue imaging in which epithelium, BM, LP, and other regional microstructures were clearly identified in normal tissues using OCT (group 1). Each OCT image represents a specific region of interest with common and unique characteristics. Figure 4 is a classic OCT image of normal buccal mucosa with clear identification of the stratified squamous epithelium (SS) along the mucosal surface, the underlying LP, and the transition between these tissues along BM boundary. The darker appearance of the epithelium is directly related to its lower optical density and scattering properties, which, in turn, results in lower signal intensity. By contrast, the LP is a more optically dense tissue and appears brighter in comparison caused by a higher signal intensity. Taking note of these optical differences, OCT permits identification of seromucinous glands along the mucosal surface of the lower lip (Figure 5), filiform papillae along the dorsum of the tongue (Figure 6), and the elaborate glandular duct system within the floor of the mouth (Figure 7). Signal propagation or absorption allowed for further characterization of tissue microstructures. Duct and lymphatic systems are characterized by regions of low signal intensity (black regions), which likely represent clear fluid that does not scatter or absorb light. As a result, signal intensity is relatively undiminished after passing through these structures and is readily observed beyond their distal walls (Figure 8). In contrast, blood vessels produce an “optical shadow” with the absorption of light by hemoglobin and the result of the image loss beyond the point of the proximal vessel lumen (Figure 9). The normal tissue OCT studies (group 1) provides a set of reference images for comparative analysis with diseased tissues (groups 2-4).

In group 2, tissue microstructure was different in comparison with native tissues depending on the pathophysiology. For example, in Figure 10, OCT images of a mucous-filled vallecular cyst show 2 thin tissue layers and a region of hyperplastic epithelium enveloping a large region of low signal intensity (presumably turbid mucus); correlative endoscopic and histologic images are provided for comparison. In other accounts, signal intensity and uniformity of regional epithelium was markedly different compared with group 1. As seen in Figure 11, OCT imaging of leukoplakia revealed increased optical density of the epithelium, superficial signal “flash points” with secondary shadow formation, and a varying epithelial thickness (Figure 11). These findings represented acanthosis, hyperkeratosis, mild dysplasia, and chronic inflammation in correlative histologic analysis.

In invasive malignant neoplasms (group 3), marked changes in the appearance of the OCT images were identified. Ablation of the tissue architecture and elimination of the BM were seen in all cases of histologically
proven malignancy. Figure 12 clearly demonstrates the characteristic loss of layering and BM integrity. Elimination of the BM in addition to the destruction of normative tissue microstructures produced images with poorly identifiable landmarks, boundaries, or microanatomical structures.

During the study, careful attention was made to image the clinical margins of both benign and malignant lesions (group 4). Disruption of tissue architecture was observed as pathologic tissues encroached into healthy regions. In Figure 13, the transition from normal epithelium and BM to invasive tumor is clearly demonstrated. Furthermore, tangential orientation of OCT imaging along the boundary of a lesion revealed multiple points of BM invasion offset by islands of intact microarchitecture (Figure 14).

The histologic images shown in Figures 10 through 14 illustrate the submucosal microanatomy for each of the related OCT Images. These histologic cross sections correlate with the OCT cross-sectional images, confirming the findings noted above.

The in vivo characterization of mucosal and submucosal tissues is of exceptional value in the treatment, diagnosis, and monitoring of both benign and malignant diseases within the upper aerodigestive tract. The tissues of the oral cavity and oropharynx are readily accessible with epithelial changes observed long before BM transgression in premalignant conditions. Unfortunately, considerable chal-
Challenges exist in distinguishing premalignant lesions from early tumor development on physical inspection and with current noninvasive modalities. Characterization of pathologic lesions is limited by the inability to look into the tissues. Visual imaging provides information about the surface of the lesion but very limited information about the BM and subsurface structures.

Within the oral cavity and oropharynx healthy epithelium varies from 75 to 550 µm in thickness while the underlying LP may extend up to 2000 µm.\textsuperscript{12,13} To reliably characterize mucosal pathologic processes an imaging modality must have a functional resolution approaching 10 µm.\textsuperscript{14} In comparison, the effective resolutions of magnetic resonance imaging (0.5-1 mm), computed tomography (1-5 mm), and ultrasonography (100-500 µm) are at least an order of magnitude larger than required to observe pathologic changes in the boundary between the epithelium and the LP.\textsuperscript{15-17} Other modalities, such as contact endoscopy, use in vivo staining with methylene blue, acetic acid, or indigo carmine to produce a resolution of 10 to 71 µm with the capability of observation up to 100 µm below the tissue surface.\textsuperscript{18} Only the most superficial layers of the epithelium can be visualized due to intrinsic optical limitations, and the images are obtained in an en face plane, an orientation that clinicians trained in reading cross-sectional histology are unaccustomed to. Furthermore, many disease processes of the oral mucosa result in immense epithelium thickening, thereby precluding any possibility of BM evaluation using existing technology.\textsuperscript{14,19} As a result of these limitations, tissue biopsy remains the diagnostic procedure of choice.

Optical coherence tomographic imaging can produce detailed cross-sectional images of the subsurface microstructure at an axial resolution of approximately 9 µm and a lateral resolution of 10 µm using safe low-power near infrared light in vivo and in real time. Optical coherence tomography has been used in ophthalmology,\textsuperscript{20-22} but its use in the head and neck has been limited, as imaging requires the patient to remain motionless and endoscopy is often needed to visualize the region of interest. Because of the combined requirement for the patients to remain motionless and the need for general anesthesia in most head and neck endoscopy, most studies to date have focused on the ex vivo imaging of analogous animal tissues.\textsuperscript{7,8,23-25} Ex vivo tissue has considerably different optical properties than in vivo due to the loss of blood, oxidation of hemoglobin, and tissue desiccation. In the head and neck, only a few limited in vivo imaging studies have been published that have focused on patients with cancer. Sergeev et al\textsuperscript{26} and Bouma and Tearney\textsuperscript{27} have reported the use of

![Figure 12](image1.png)

**Figure 12.** Invasive squamous cell carcinoma of maxilla illustrated using optical coherence tomography (A) and correlative histologic features (B). Note absence of basement membrane and tissue microstructures. Vertical bar indicates 500 µm.

![Figure 13](image2.png)

**Figure 13.** Multimodal analysis revealing transition zone of cancer and adjacent tissue along anterior mandible margin using optical coherence tomography (A); endoscopic view (B), and correlative histologic features (C). CA indicates cancer (squamous cell carcinoma); LP, lamina propria; SS, stratified squamous epithelium; and vertical bar, 500 µm.
OCT in the aerodigestive tract. These findings were limited by the challenges in the depth of imaging, resolution, and speed of image acquisition inherent in OCT imaging. By using a different light source and a faster and more robust piezoelectric translation stage, we have been able to improve resolution from 15 to 20 µm to 10 µm, increase maximum image depth by almost 50%, vary the horizontal image width from 2 mm to up to 10 mm, and increase the frame capture rate from 1.5 to 2 seconds for a 2-mm scan to 1 second for a 6-mm scan. The improved resolution and depth of imaging allows more rapid and precise characterization of the tissue structures, which we have used to perform extensive in vivo imaging of normal, benign, and malignant tissues in the oral cavity and oropharynx.

This study confirms the feasibility of using OCT to identify differences in mucosal and submucosal tissue structures of the oral cavity and oropharynx. Epithelium, LP, and BM boundaries were clearly identified in all normal tissues imaged. Each tissue microstructure, including papillae, glands, ducts, and blood vessels, revealed unique optical characteristics that allowed for image correlation with known histologic features. Optical coherence tomographic imaging of pathologic conditions such as leukoplakia revealed benign changes in epithelial density and tissue thickness without alteration in the BM. In contrast, OCT images of early cancer revealed ablation of normal tissue microanatomy in addition to BM compromise. Further imaging of lesion borders revealed transition zones between cancerous and noncancerous tissues. Optical coherence tomographic images of these clinically suspicious lesions were correlated with histologic specimens and intraoperative digital photographs. Of particular note were the considerable histologic distortions created by tissue biopsy as well as fixation of specimens when compared with OCT images of the same region. The in vivo OCT imaging of living tissues may represent a truer biologic picture than any image produced by other high-resolution modalities.

The ability to perform in vivo tissue imaging with a noninvasive, high-resolution modality could significantly alter the detection and management of early stage disease. With the capability to perform a tomographic study, one could easily monitor disease, document progression, and target tissue biopsy specimens to regions of greater histologic yield. Additionally, OCT is an evolving imaging modality where both improvements in source coherence bandwidth and electromechanical design will lead to higher-resolution images and faster frame rates. The application of this high-resolution imaging modality provides valuable information on the physiology of benign mucosal disorders and on the development, transformation, and progression of cancer. Our ongoing OCT investigations are focused on new applications in children, microsurgery, and office-based instruments, in addition to continuing to expand our adult clinical series.

Submitted for Publication: May 3, 2005; final revision received March 12, 2006; accepted April 22, 2006.

Correspondence: Brian J.-F. Wong, MD, PhD, Zhongping Chen, PhD, Beckman Laser Institute and Medical Clinic, University of California, Irvine, 1002 Health Sciences Rd, East Irvine, CA 92617 (bjwong@uci.edu or z2chen@uci.edu).

Author Contributions: Drs Ridgway, Jackson, Crumley, and Wong had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Ridgway, Jackson, Guo, Armstrong, Crumley, Chen, and Wong. Acquisition of data: Ridgway, Guo, Mahmood, Su, Armstrong, Shibuya, Crumley, Chen, and Wong. Analysis and interpretation of data: Ridgway, Armstrong, Crumley, Gu, and Wong. Drafting of the manuscript: Ridgway, Su, Armstrong, and Wong. Critical revision of the manuscript for important intellectual content: Ridgway, Jackson, Guo, Mahmood, Armstrong, Shibuya, Crumley, Gu, Chen, and Wong. Statistical analysis: Ridgway. Obtained funding: Wong. Administrative, technical, and ma-

Financial Disclosure: None reported.

Funding/Support: This study was supported by grants DC 006026, CA 91717, EB 00293, RR 01192, and M01-RR00827-28 from the National Institutes of Health, grant 32456 from the Flight Attendant Medical Research Institute, grant 12RT-0113 from the State of California Tobacco Related Disease Research Program, and grant F49620-00-1-0371 from the Air Force Office of Scientific Research, and the Arnold and Mabel Beckman Foundation.

Previous Presentation: This study was presented in part at the annual meeting of the American Head and Neck Society; May 15, 2005; Boca Raton, Fla.

Acknowledgment: We thank master machinist Rudolph Limberg and Steve Knisley for assistance with construction of the instrument and Aya Yamamoto, RN, and Cleve Barclay, ST, for their help and patience in the operating room.

REFERENCES