In Vivo Optical Coherence Tomography for the Diagnosis of Oral Malignancy

Petra Wilder-Smith, DDS, PhD,1* Woong-Gyu Jung, MSc,1 Matthew Brenner, MD,2 Kathryn Osann, PhD,2
Hamza Beydoun, BS,1 Diana Messadi, DDS, DMSc,3 and Zhongping Chen, PhD1
1Beckman Laser Institute, University of California, Irvine, California 92612
2University of California, Irvine, California 92612
3University of California, Los Angeles, California 90095

Background and Objective: Oral cancer results in 10,000 U.S. deaths annually. Improved highly sensitive diagnostics allowing early detection of oral cancer would benefit patient survival and quality of life. Objective was to investigate in vivo non-invasive optical coherence tomography (OCT) techniques for imaging and diagnosing neoplasia-related epithelial, sub-epithelial changes throughout carcinogenesis.

Study Design/Materials and Methods: In the standard hamster cheek pouch model for oral carcinogenesis (n = 36), in vivo OCT was used to image epithelial and sub-epithelial change. OCT- and histopathology-based diagnoses on a scale of 0 (healthy) to 6 (squamous cell carcinoma, SCC) were performed at all stages throughout carcinogenesis by two blinded investigators.

Results: Epithelial, sub-epithelial structures were clearly discernible using OCT. OCT diagnosis agreed with the histopathological gold standard in 80% of readings.

Conclusion: In vivo OCT demonstrates excellent potential as a diagnostic tool in the oral cavity. Lasers Surg. Med. 35:269–275, 2004. © 2004 Wiley-Liss, Inc.

Key words: carcinoma; imaging; leukoplakia; non-invasive diagnostics; oral diagnosis

INTRODUCTION

According to the American Cancer Society, 1,220,100 patients were diagnosed with cancer in the year 2000. In the same year, 552,200 persons were expected to succumb to cancer [1]. Despite significant advances in cancer treatment, early detection of cancer and its curable precursors remains the best way to ensure patient survival and quality of life. Oral cancer will claim approximately 10,000 lives in the U.S. this year [2,3]. Accounting for 96% of all oral cancers, squamous cell carcinoma (SCC) is usually preceded by dysplasia presenting as white epithelial lesions on the oral mucosa (leukoplakia). Leukoplakias develop in 1–4% of the population [2]. Malignant transformation, which is quite unpredictable, develops in 1–40% of leukoplakias over 5 years [2]. Dysplastic lesions in the form of erythroplakias carry a risk for malignant conversion of 90% [2]. Tumor detection is further complicated by a tendency towards field cancerization, leading to multicentric lesions [4]. Current techniques require surgical biopsy of lesions. Benign lesions are often biopsied, reducing patient motivation to agree to further diagnostic biopsies in the future. Conversely, many lesions are only detected by biopsy at an advanced stage, when treatment options and outcome are far from optimal. Of all oral cancer cases documented by the National Cancer Institute Surveillance, Epidemiology, and End Results Program, advanced lesions outnumbered localized lesions more than 2:1. Five-year survival rate is 75% for those with localized disease at diagnosis, but only 16% for those with cancer metastasis [2,3]. A modality for the direct, non-invasive early detection, diagnosis, and monitoring of oral dysplasia and malignancy and for the screening of high-risk populations is urgently required to identify treatment needs at early, more treatable stages of pathological development. Such multi-use clinical capabilities would be likely to produce a sharp drop in morbidity and mortality due to cancer, with substantial reductions in patient anxiety and suffering as well as treatment cost.

Optical Coherence Tomography (OCT)

OCT is a new high-resolution optical technique that permits minimally invasive imaging of near surface abnormalities in complex tissues. It has been compared to ultrasound scanning conceptually [5]. Both ultrasound and OCT provide real time structural imaging, but unlike ultrasound, which utilizes sound waves, OCT is based on low coherence interferometry, using broadband light to...
provide cross-sectional high resolution sub-surface tissue images [6,7]. The engineering principles behind OCT have been described previously [8–11]. Broadband laser light waves are emitted from a source and directed toward a beam splitter. One wave from the beam splitter is sent toward a reference mirror with known path length and the other toward the tissue sample. After the two beams reflect off the reference mirror and tissue sample surfaces at varying depths within the sample, respectively, the reflected light is directed back towards the beam splitter, where the waves are recombined and read with a photo detector (Fig. 1). The image is produced by analyzing interference of the recombined light waves. Cross-sectional images of tissues are constructed in real time, at near histologic resolution (approximately 10 μm with the current prototype technology). Previous studies using OCT have demonstrated that the imaging penetration depth suffices to evaluate macroscopic characteristics of epithelial and sub-epithelial structures with potential for near histopathological level resolution and close correlation with histologic appearance [12–14].

While some research has been reported in ophthalmologic, dermatologic, GI, gynecological, cardiac, and other OCT applications [11–29], there is limited information on intra-oral OCT, mainly on the topics of periodontal disease [30–33]. Broadband laser light waves are emitted from a source and directed toward a beam splitter. One wave from the beam splitter is sent toward a reference mirror with known path length and the other toward the tissue sample. After the two beams reflect off the reference mirror and tissue sample surfaces at varying depths within the sample, respectively, the reflected light is directed back towards the beam splitter, where the waves are recombined and read with a photo detector (Fig. 1). The image is produced by analyzing interference of the recombined light waves. Cross-sectional images of tissues are constructed in real time, at near histologic resolution (approximately 10 μm with the current prototype technology). Previous studies using OCT have demonstrated that the imaging penetration depth suffices to evaluate macroscopic characteristics of epithelial and sub-epithelial structures with potential for near histopathological level resolution and close correlation with histologic appearance [12–14].

The 10-μm resolution of OCT permits in vivo non-invasive imaging of the macroscopic characteristics of epithelial and sub-epithelial structures including: (1) depth and thickness, (2) peripheral margins, and (3) potential histopathological appearance. Thus OCT improves on existing clinical capabilities, particularly for identification of multifocal biopsy sites, for regular monitoring of lesions, and for screening high-risk populations. With tissue penetration depth of 1–3 mm, the imaging range of OCT diagnostics is suitable for the oral mucosa. The normal human oral mucosa is very thin, ranging from 0.2 to 1 mm. In hamsters, it is usually somewhat thicker (up to 2 mm), due to the marked surface keratotic layer.

Goal of these feasibility studies was to determine whether premalignant and malignant transformation can be detected and diagnosed in vivo using non-invasive OCT. Specifically, our aims were to (1) identify to what extent OCT can characterize epithelial and sub-epithelial change during carcinogenesis in the hamster cheek pouch model, and (2) compare these data with histopathological diagnosis and staging and determine diagnostic capability.

MATERIALS AND METHODS

Animal Model

The standard Golden Syrian Hamster (Mesocricetus auratus) cheek pouch model was used. By application of 0.5% DMBA (9,10 dimethyl-1,2-benzanthracene (Sigma, St. Louis, MO) in mineral oil three times per week, mild to severe dysplasia developed at 3–6 weeks, progressing to SCC at 10 weeks. Histological features in this model have been shown to correspond closely with premalignancy and malignancy in human oral mucosa [34]. The study used 36 female animals, 10–12 weeks old. One animal died at 2 days carcinogenesis, leaving a total of 35 animals in this study. To improve study logistics, the animals were randomly divided into three groups of 12 animals each. Only when all the animals from one group had been imaged and sacrificed was the next group was incorporated into the study. The animals were housed and treated in accordance with animal research committee guidelines at the University of California, Irvine (approval 97–1972). The median-lining wall of one cheek pouch of each hamster was treated with carcinogen; the other cheek pouch served as control. Previous studies have shown that this carcinogenesis process in one cheek pouch does not affect the other cheek pouch [35,36], and that therefore the untreated cheek pouch can be used as control.

Protocol

During carcinogenesis over 12 weeks, in vivo (i) clinical evaluation and photography, (ii) high resolution OCT/ODT were performed at weekly intervals. During in vivo measurements, the cheek pouch was continuously irrigated with isotonic saline at room temperature to avoid dehydration. The animals were kept warm using a heating pad and a thermal sock. Use of these measures eliminated animal mortality almost completely during the sometimes lengthy imaging sessions. Only one unplanned animal death occurred throughout the duration of these investigations. Early imaging sessions could last up to 10 minutes; once our techniques were established imaging sessions averaged <3 minutes. The anesthetized hamster’s everted cheek pouch was attached to the microscope stage using a specially designed and fabricated ring-shaped clamp fastened rigidly to the stage surface (Fig. 2). The clamping device was marked on its rim at 1-mm intervals to allow the use of localization coordinates for designating areas of specific interest and for achieving repeated, atraumatic scans with different modalities in exactly the same location. Accurate re-localization of the clamping device at weekly intervals was ensured by marking several coordinates of the device outline on the hamster cheek pouch using an animal micro tattooing device (Ketchum lab animal micro tattooer, Ottawa, CDN) at least 7 days prior to the commencement of the study. This time frame was selected as preliminary studies had demonstrated the absence of any tattooing-related tissue changes after a minimum of 5-day post-tattooing. Whilst recognizing the value of tattooing in very close proximity to the area of interest to facilitate the co-localization of corresponding OCT images and
Histological Evaluation

Histological evaluation of each stained section was quantified by two blinded, pre-standardized scorers (one oral pathologist, one dentist) according to the criteria established by Macdonald [34], whereby each characteristic listed below was assessed. Although a well-defined histological grading system for oral epithelial dysplasia has not been developed yet, most oral pathologists grade on the scale of mild-moderate-severe dysplasia depending on the range and severity of individual features and the proportion of epithelium thickness affected. The following numerical grading system was used for each slide: 0—healthy, 1—hyperkeratosis, 2—mild dysplasia, 3—moderate dysplasia, 4—severe dysplasia, 5—carcinoma-in-situ, and 6—SCC. The criteria for Oral Epithelial Dysplasia were as follows: drop-shaped rete ridges, irregular epithelial stratification, individual cell keratinization, basal cell hyperplasia, loss of intercellular adherence, loss of polarity, hyperchromatic nuclei, increased nucleo-cytoplasmic ratio, anisocytosis, pleomorphic cells and nuclei, abnormal mitotic figures, and increased mitotic activity. Each site was assessed for each of these characteristics at a level of either none (0), slight (1), or marked (2).

OCT Evaluation

The two pre-trained scorers classified each image diagnostically in a blind manner on a scale of 0 (normal) to 6 (SCC). This scale was designed to parallel the scale used for histopathological evaluation [34]. OCT diagnostic scores were based on changes in keratinization, epithelial thickening, epithelial proliferation and invasion, broadening of rete pegs, irregular epithelial stratification, and basal hyperplasia. Epithelial invasion was defined as loss of visible basement membrane. Each site was assessed for each of the above characteristics at a level of either none (0), slight (1), or marked (2). The score for each site depending on the range and severity of individual features and the proportion of epithelial thickness affected.

Data consisted of descriptive images. Scorers were pre-trained using a standard set of 50 OCT and 50 matching histopathological images. Initial training was repeated until at least 98% of images were identified correctly, then 50 new sets of OCT and histopathology images were identified by each scorer with at least 90% accuracy. At this stage, scorers were deemed “pre-standardized” and ready to participate in these studies. Each scorer evaluated all data in one session, which took place once all data accrual was complete. A second re-evaluation of all images by the same scorers in one session, 3 months later, was used to evaluate intra-observer variability.

Statistical Evaluation

To test for agreement between the two scorers, and between the same scorer at the first and second evaluation of each sample, a kappa statistic was used. A consensus diagnostic score for any specific image was created by taking the average of the two ratings rounded to the nearest whole integer. The histopathology score was used as the gold standard. Because histopathology was classified into seven different categories, agreement between OCT and histopathology was first described by correlations and percent agreement, then by sensitivity and specificity,
which are more appropriate for dichotomous discrete variables. The diagnostic sensitivity and specificity of OCT were defined using two approaches: (a) investigating the ability of OCT to differentiate between healthy (0–1) versus pathological (2–6) lesions, and (b) investigating the ability of OCT to differentiate between malignant (5–6) versus non-malignant (0–4) lesions. These values were calculated considering data from each scorer separately (n = 70) and also using the consensus score (n = 35).

RESULTS

The in vivo OCT technology used in these investigations was able to image multiple epithelial and sub-epithelial layers throughout carcinogenesis in the hamster cheek pouch model (Figs. 3–5). Light penetration and image resolution were consistently better in the non-malignant tissues.

The results for diagnosis of each image by two pre-trained investigators are depicted in Table 1. OCT and histopathology were classified into one of seven categories (0 = normal tissue to 6 = SCC). Agreement within scorers, between scorers and between modalities, was assessed using kappa statistics. Intra-observer agreement for the two modalities (histopathology and OCT) at the two scoring timepoints was excellent. Using the kappa statistic for each of the two observers separately and for the observers combined agreement was greater than or equal to 90% for each modality (Table 2).

Fig. 3. Healthy cheek pouch H&E stained image (A, 20×; B, 40×) and in vivo OCT (C) of same healthy cheek pouch tissues. Thin normal epithelial and thick sub-epithelial layers are visible. 1—keratinized surface layer, 2—flat stratified squamous epithelium, 3—submucosa: dense fibrous connective tissue, 4—longitudinal striated muscle fiber, and 5—basement membrane.

Fig. 4. Dysplastic cheek pouch H&E (A, 20×; B, 40×): Epithelial thickening, increase in hyperchromatism, pleomorphism of individual cells, loss of polarity in basal cell layer. In vivo OCT (C), overall epithelial thickening, broader rete pegs, loss of polarity in basal cell layer. 1—keratinized surface layer, 2—flat stratified squamous epithelium, 3—submucosa: Dense fibrous connective tissue, 4—longitudinal striated muscle fiber, and 5—basement membrane.

Fig. 5. Malignant cheek pouch. H&E stained image (A, 20×; B, 40×) and in vivo OCT (C) of same cheek pouch with squamous cell carcinoma. In H&E, epithelial pearls are invading the connective tissue. Individual cancer cells exhibit pleomorphism, increased nuclear:cytoplasmic ratio, and hyperchromatism. In OCT, notice loss of normal epithelial stratification and basement membrane. Image definition from sites with SCC was consistently much poorer than from other sites, due to reduced penetration by light into malignant tissues.
Inter-observer agreement was assessed for both histopathology and OCT. For histopathology, there was perfect agreement between the two scores for any sample. For OCT, there was never a discrepancy of more than 1 point between scorers. The kappa statistic for OCT was 0.603 (SE = 0.091), indicating moderate agreement between observers for scoring.

Overall agreement between readings for OCT and histopathology are illustrated in Table 3. Because histopathology was classified into seven different categories, agreement between OCT and histopathology was first described by percent agreement and the kappa statistic, then by sensitivity and specificity, which are more appropriate for dichotomous discrete variables.

OCT diagnosis agreed with the histopathology for 56 of 70 (80%) readings (kappa = 0.767, SE = 0.056). Using OCT, the histopathology was underestimated by one category in six cases and overestimated in eight cases. In no instance was the difference between histopathology and OCT more than one level. Diagnostic sensitivity of OCT for differentiating between healthy (0–1) versus pathological (2–6) lesions was 0.98 (SE = 0.020) and specificity was 0.95 (SE = 0.049) if each score is considered separately (n = 70). Using this technique, each sample is counted twice, as each sample was evaluated separately by each of the two scorers. Using the consensus score (n = 35) from both investigators (in case of non-consensus, the higher of the two scores was used), sensitivity for OCT was 100% and specificity was 90%.

Diagnostic sensitivity and specificity for differentiating between malignant (5–6) versus non-malignant (0–4) lesions was 100% and specificity was 96% (SE = 0.028). Using the consensus score from both investigators (in case

### Table 3. Frequencies: OCT (Rows) by Histopathology (Columns)

|      | 0 | 1 | 2 | 3 | 4 | 5 | 6 | Total |
|------|---|---|---|---|---|---|---|-------|
| +    | 11| 9 | 2 | 1 | 0 | 0 | 0 | 10    |
| +    | 9 | 1 | 7 | 1 | 0 | 0 | 0 | 11    |
| +    | 9 | 0 | 1 | 0 | 0 | 0 | 0 | 9     |
| +    | 9 | 0 | 1 | 0 | 0 | 0 | 0 | 9     |
| +    | 11| 5 | 2 | 0 | 0 | 0 | 0 | 10    |
| +    | 11| 6 | 0 | 0 | 0 | 0 | 0 | 10    |

| Total | 10 | 10 | 10 | 10 | 10 | 10 | 70  |
of non-consensus, the higher of the two scores was used), sensitivity for OCT was 100% and specificity was 96%.

**DISCUSSION**

These studies demonstrate that in vivo OCT can achieve high-resolution images of epithelial and sub-epithelial change, identifying epithelial thickening, as well as changes in stratification and structure.

Inter-observer variability with regard to diagnosis was acceptable, with discrepancies between the evaluators never exceeding 1 point. Scorer consistency was excellent, with repeat evaluations after 3 months showing a kappa equal to or exceeding 90%. Sensitivity was excellent for the diagnosis of SCC, and somewhat reduced for differentiating between different types of dysplasia. As the resolution capabilities of OCT technology are rapidly increasing, their ability to differentiate between different stages of dysplasia should also improve.

Thus, these feasibility studies confirm the usefulness of high-resolution, in vivo OCT imaging as a promising tool for clinical and research needs related to oral premalignancy and malignancy. Further studies are underway to engineer new approaches to the clinical diagnosis and management of oral pathologies, and to a better understanding of the processes predicting and paralleling premalignant and malignant change. In future, these capabilities will permit investigation, in patients, into the effects of chemoprevention and chemotherapy on the parameters described above, providing a better understanding of pathological mechanisms, predictors of malignant change in dysplasia, risk of tumor recurrence, and predictors of tumor response to therapy.

**REFERENCES**

1. American Cancer Society. Cancer facts and figures. In: American Cancer Society Report, 2000. 4 p.
2. Regezi J, Sciubba J, editors. Oral pathology. Philadelphia: W.B. Saunders Co.; 1993. pp 77–90.
3. California Department of Health Services. Cancer Surveillance Section Annual Report, March 1999.
4. Slaughter DP. Field cancerization in oral stratified squamous epithelium. Cancer 1953;6:963–968.
5. Izatt JA, Kobayashi K, Sivak MV, Barton JK, Welch AJ. Optical coherence tomography for biodiagnosis. Opt Photon News 1997;8:41–47.
6. Ding Z. High-resolution optical coherence tomography over a large depth range with an axicon lens. Opt Lett 2002;27:4.
7. Huang D, Swanson EA, Lin CP, Schuman JS, Stinson WG, Chang W. Optical coherence tomography. Science 1991;254(5035):1178–1191.
8. Swanson EA, Izatt JA, Hee MR, Huang D, Lin CP, Schuman JS. In vivo retinal imaging by optical coherence tomography. Opt Lett 1993;18(21):1486–1488.
9. Fujimoto JG, Hee MR, Izatt JA, Boppart SA, Swanson EA, Lin CP, et al. Biomedical imaging using optical coherence tomography. Proc SPIE 1999;3749:402.
10. Bouma B, Tearney GJ, Boppart SA, Hee MR, Brezinski ME, Fujimoto JG. High-resolution optical coherence tomographic imaging using a mode-locked Ti:Al2O3 laser source. Opt Lett 1995;20(13):1486–1488.
11. Boppart SA. Optical coherence tomography: Technology and applications for neuroimaging. Psychophysiology 2003;40(4):529–541.
33. Matheny ES, Hanna N, Mina-Araghi R, Jung WG, Chen Z, Wilder-Smith P, Brenner M. Optical coherence tomography of malignant hamster cheek pouches. J Invest Med 2003; 51(1):S78.

34. MacDonald DG. Comparison of epithelial dysplasia in hamster cheek pouch carcinogenesis and human oral mucosa. J Oral Pathol 1981;10:186–191.

35. Wilder-Smith P, Liaw LH, Krasieva TB, Messadi D. Laser-induced fluorescence for detection and diagnosis of oral malignancy. J Dent Res 1999;78:820.

36. Wilder-Smith P, Liaw L-H, Krasieva TB, Nguy L, Yoon Y, Messadi D. Topical ALA-induced fluorescence in oral dysplasia and malignancy. Lasers Surg Med 1999; 69:39.

37. Tearney GJ, Bouma BE, Fujimoto FG. High-speed phase- and group-delay scanning with a grating-based phase control delay line. Opt Lett 1997;22:1811–1813.

38. Rollins AM, Kulkarnis MD, Yazdanfar S, Ung-arunyawee R, Izatt JA. In vivo video rate optical coherence tomography. Opt Express 1998;3:219–229.