ABSTRACT: Molecular biological markers have been suggested to be of value in the diagnosis and prognostic evaluation of precancerous lesions. The purpose of the current study was to investigate the expression of p53 protein in normal oral mucosa, oral dysplastic lesions (DL), and oral squamous cell carcinoma (SCC), comparing patients with the habit of reverse smoking and conventional smokers. The patients were subjected to incisional biopsy and the soft tissue specimens were routinely fixed in 10% formalin and processed in the laboratory. Immunohistochemical technique was performed using the avidin-biotin-peroxidase protocol. The 102 patients included 42 reverse smokers and 60 conventional smokers. There were 79 cases of mild, 15 of moderate, and 3 of severe epithelial dysplasia. Five microinvasive oral SCC included one male and four females. The clinical characteristics of microinvasive tumors included patches, plaques or erosions; 100% showed positive nuclear staining for p53. It was found a significant association of p53 expression and exposure of reverse smoking among microinvasive oral SCC and oral DL patients in the population studied.

KEY WORDS: p53, immunohistochemistry, oral squamous cell carcinoma, reverse smoker.
Consumption of tobacco can cause genetic and molecular alterations in clinically distinct oral premalignant lesions (Schmidt et al., 2004). Reverse smoking is an unusual tobacco habit performed by the smoker placing the lighted end of a cigarette inside the mouth; the seal provided by the lips allows to the slow inhaling of the cigarette. This habit is practiced in various parts of the world, including The Netherlands, Southeast Asia, Indian subcontinent, Sri Lanka, Jamaica, Panama, Venezuela, Caribbean Islands and Colombia; occasionally immigrants to Western countries continue the habit of reverse smokers. In Colombia there are some very well defined regions with presence of this habit that seems to be transmitted through generations.

The clinical aspect of the oral mucosa in patients with the habit of reverse smoke varies when compared with conventional smokers. The most commonly affected areas are tongue and palate include both benign and malignant mucosal pathologies. Besides, reverse smoking is claimed to intensify the carcinogenic effect of heat and smoke resulting in increased risk of cancer development (Mercado-Ortiz et al., 1996).

To our knowledge no studies have examined the expression of p53 protein in oral dysplastic lesions (DL) and oral SCC, in patients with the habit of reverse smoking. Therefore, the aim of the present study was to investigate the expression of p53 protein in normal oral mucosa, oral DL, and oral SCC, using immunohistochemistry, comparing patients with the habit of reverse smoking and conventional smokers.

**MATERIAL AND METHOD**

This cross sectional study was performed during the first semester of 2009. An informative campaign was prepared through field advisors and community leaders of Hato Nuevo, San Francisco and Cayo de Palma villages, Sucre Department in Colombia, in order to notify the population about the investigation. Then, the researchers group visited the villages with the purpose of accomplish the epidemiologic gathering, to identify and select people with the habit of reverse smoking according to the criteria of inclusion (regular reverse smokers of any type of tobacco, during a period longer than 6 months, with a minimum dose of one cigarette per day). Conventional smokers also were included as control. This process was done with a home to home survey, with socio demographic variables, concerning attitudes and behaviors related with the use of tobacco.

All the 102 patients selected were subjected to incisional biopsy and the soft tissue specimens were routinely fixed in 10% formalin and processed in the laboratory (Pathology Department, School of Medicine of the University of Antioquia in Medellín, Colombia).

Dysplasia was classified as mild, moderate or severe following the WHO tumor classification (Gale et al., 2005). A representative block for each case was selected and multiple 4 μm sections were cut. The remaining 15 specimens included cases with normal mucosa. Normal controls were selected based on no history of alcohol or tobacco use.

Hematoxylin and eosin stained sections were prepared from the paraffin blocks and examined to determine the histologic tumor type, grade of tumor differentiation, depth of tumor invasion, and to assess the degree of dysplasia in the oral DL (Gale et al.).

Immunohistochemical technique was performed using the avidin-biotin-peroxidase protocol, according to Abrahao et al. (2011). Briefly, antigen retrieval was performed with Target antigen retrieval solution pH 9 (Dako A/S, CA, USA) in a water bath, followed by incubation with 6% hydrogen peroxide to quench endogenous peroxidase. The sections were then incubated in blocking solution (3% bovine serum albumin) for 1 hour at room temperature, followed by primary antibody incubation, previously diluted in blocking solution. Anti-p53 (clone DO-7, 1:200 dilution — DAKO A/S, CA, USA) antibody was incubated for 30 minutes at room temperature. Sections were exposed to the LSABTM system (DAKO A/S, CA, USA), developed in diaminobenzidine (Dako A/S, CA, USA) and counterstained in Mayer’s hematoxylin.

For the antibody, positive and negative controls were used. Two independent observers reviewed the immunohistochemically-stained sections. In normal mucosa and dysplasia, p53, according to Cruz et al. (1998) and to Pirog et al. (2002) were classified as positive or negative, attributing a score taking into account both the relative number of stained nuclei and their localization in the upper 2/3 of the epithelium. In the carcinomas, evaluation was made in the areas of infiltration, where cell proliferation was more active. Breast tumor tissue which had previously been identified as p53 immunopositive was used as a positive control.
On admission the patients gave their consent to use the data obtained during the examination and treatment for further scientific projects. The study design was approved by the Ethics Committee on Human Research of the School of Dentistry of the University of Antioquia according to the Declaration of Helsinki.

Data collected were analyzed using the statistical package for social sciences (SPSS) for Windows version 19. Data was summarized in form of proportions and frequency tables for categorical variables. Means, median and standard deviation were used to summarize continuous variables. The Chi-Square test was used to compare the counts of categorical response between two independent variables. The statistical significance was fixed at 0.05.

RESULTS

This study was performed on resection specimens from five oral microinvasive SCC and 97 biopsies from oral DL. The 102 patients included 42 (41.2%) reverse smokers and 60 (58.8%) conventional smokers. Fifteen normal samples of oral mucosa were used as control. A total of 41 reverse smokers were women, while there were no gender differences in conventional smokers. Reverse smokers revealed a longer duration of smoking (32.7±18 years) than conventional smokers (30.2±15 years).

There were 79 cases of mild, 15 of moderate, and 3 of severe epithelial dysplasia. In all cases, consensus on the grading of dysplasia was achieved following review. Five microinvasive oral SCC included one male (conventional smoker) and four females (reverse smokers). In the present study, the clinical characteristics of microinvasive tumors included patches, plaques or erosions rather than ulcers or verrucous lesions.

p53 was confined to the basal layer in the normal oral epithelium (Fig. 1). In oral DL, p53 was present in the basal and suprabasal layers (Fig. 2). Of the five microinvasive oral SCC, 100% showed positive nuclear staining for p53 (Fig. 3).

In the 97 oral DL, p53 was immunoexpressed in 74% and 78% of the reverse and conventional smokers respectively.
The pattern of immunoexpression of p53 has been studied in a broad diversity of investigations (Abbas et al.; Kerdpon et al., 2001; Abrahao et al.). However, the current study has a particular interest because it investigates the expression of p53 in SCC and DL of reverse smokers which was not previously reported to date. In those countries where reverse smoking is practiced, a wide range of oral tissue changes have been observed (Mercado-Ortiz et al.; Alvarez-G—mez et al., 2008), including both benign and malignant oral mucosa pathologies. This different form of tobacco habit has been shown to be associated with oral cancer and precancerous lesions confirming our results (Alvarez-G—mez et al.). The evidence that the p53 pathway is very important in the biology of oral cancer has led to the application of such immunohistochemical analysis as a simple rapid and inexpensive method to potentially malignant lesions in an attempt to find a useful marker to predict progression to oral SCC (Montebugnoli et al., 2008). Moreover, the advantage of immunohistochemical staining is the direct demonstration of the spatial relationship of cells that have altered p53 protein expression, which is of particular importance in the study of clonal expansion of altered cell populations during multistage tumor genesis (Kushner et al., 1997).

In this study, p53 expression was confined to the basal cell layer of normal oral mucosa. In the dysplastic group, p53 was expressed in the basal and suprabasal layers, whereas the five microinvasive oral SCC showed positive nuclear staining for p53. Diverse investigations revealed comparable findings (Cruz et al.; Iamaroon et al., 2004). p53, when detected in oral mucosa without malignant potential was interpreted as physiological response of cells to genotoxic stress (Cruz et al.). In contrast, p53 expression in suprabasal cells was only detected in premalignant lesions, reflecting the presence of proliferating cells with DNA damage in more superficial layers of the epithelium (Iamaroon et al.).

To our knowledge this is the first investigation that founded p53 expression in microinvasive oral SCC. Our study revealed that in microinvasive carcinoma, the number of specimens positive for p53 was higher in comparison with the dysplastic lesions. In this respect, Warnakulasuriya & Johnson (1992) reported that positive included carcinomas without
deep invasion suggesting that p53 mutation may occur in the early stages of progression of a malignancy. Pentenero et al., (2011) found one cancer-related death observed in patients with oral microinvasive lesions. Additionally, they indicated that microinvasive carcinomas were diagnosed incidentally through a habitual clinical evaluation or during a follow-up of a premalignant lesion, rather than due to the presence of symptoms. Confirming our results, a previous investigation showed that microinvasive SCC had similar clinical aspects (erosions, patches and plaques) to our findings. As microinvasive lesions tend to present clinically as premalignant lesions, accurate clinical examination is essential if misdiagnosis of early lesions is to be avoided (Pentenero et al.).

In the present study, p53 was established in 74% and 78% of the oral dysplastic lesions in reverse and conventional smokers, respectively. This result confirms the previous reports which assumed that expression of p53 in the premalignant oral lesions may be an early episode in oral carcinogenesis (Ralhan et al., 2000). Consequently, the expression of p53 in the oral DL may facilitate in identifying patients who are at high risk for cancer development. This statement is supported by a follow-up study that found that no one of the p53 negative oral DL progressed to cancer, while three of the 26 p53 positive dysplasias developed cancer (Rich et al., 1999). Consistently, Cruz et al. observed that 86% of premalignant lesions that showed p53 expression above the basal layer developed into SCC.

In this sample, 82% of cases stained positive for p53. The high prevalence is comparable with those reported previously (Kerdpon et al., 1997; Ralhan et al.). Our finding is a result of the high prevalence of p53 expression associated with the habit of smoking tobacco, where reverse smokers increased significantly this high frequency. This is consistent with Field et al. (1994) who analyzed the smoking history of 71 patients by logistic regression and found that smoking correlates with p53 overexpression. Similarly, increased p53 positive cells were detected in patients with both betel quid chewing and smoking habits (Humayun & Prasad, 2011).

An important finding in our work is that reverse smokers were positively correlated with p53 expression and consequently with a higher number of oral DL and SCC. Some studies may explain this information. Quigley et al. (1965) showed that the maximum internal cigarette temperature in seven reverse smoking experiments was 760°C. The reverse smoking experiments lasted longer; in practice reverse smoking lasts as long as 18 min. Therefore, palatal temperatures rise appreciably during reverse smoking, indicating that substantial infrared inputs occur and transient tissue dehydration takes place. In line with this fact, White et al. (2001) found that tobacco heated to temperatures of 475°C or greater generated smoke aerosol that was detectably mutagenic. However, the relevance of these findings regarding cancer risk in humans is difficult to assess because of the lack of a direct correlation between mutagenicity and carcinogenicity (White et al.). Further investigations about the relationship between p53 and reverse smoking are needed to provide more understanding of habit-related oral carcinogenesis.

In this study, females were more affected with the premalignant and malignant lesions than males, because traditionally females are more likely to display reverse smoking. The fact that reverse smoking is predominantly a female habit is possibly a consequence of some of the sociologic roles played by women in cultures where this habit is observed (Mercado-Ortiz et al.; Alvarez-Gómez et al.).

The relationship of p53 expression and clinicopathological parameters has been reported with variable results. Most studies found no correlation of p53 expression with age, sex, site, disease stage and tumor differentiation (Rowley et al., 1997; Yao et al., 1999) while some reported a relation with tumor differentiation (Ng et al., 1999). In conclusion, the present study found a significant association of p53 expression and exposure of reverse smoking among microinvasive oral SCC and oral DL patients in the population studied. Additionally, the significant correlation between progression of oral epithelium from normal to neoplasia and increased expression of p53 suggests that it may be useful biomarker of malignant transformation in oral precancerous lesions and may serve as transitional point for cancer prevention programs.

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RESUMEN: Marcadores biológicos moleculares se han sugerido para ser de valor en el diagnóstico y evaluación pronóstica de lesiones precancerosas. El propósito del presente estudio fue investigar la expresión de la proteína p53 en mucosa oral normal, lesiones displásicas orales (LD), y en el carcinoma oral de células escamosas (COCO), comparando pacientes con el hábito de fumar convencional y fumar invertido. Los pacientes fueron sometidos a biopsia por incisión y los especímenes de tejidos blandos fueron fijados en formalina al 10% y se procesaron en el laboratorio. La técnica inmunohistoquímica se utilizó utilizando el protocolo de avidina-biotina-peroxidasa. De los 102 pacientes incluidos, 42 fueron fumadores invertidos y 60 fumadores convencionales. Hubo 79 casos de displasia epitelial leve, 15 moderada, y 3 severa. Cinco casos de COCC microinvasivo incluyeron un hombre y cuatro mujeres. Las características clínicas de los tumores microinvasivos incluyen parches, placas o erosiones; el 100% mostró tinción nuclear positiva para p53. Se encontró una asociación significativa de la expresión de p53 y la exposición de los fumadores invertidos entre pacientes con COCC microinvasivo y DL en la población estudiada.

PALABRAS CLAVE: p53, inmunohistoquímica, carcinoma oral de células escamosas, fumador invertido.

REFERENCES


