INTRODUCTION

Oral cancer is considered to be the sixth most prevalent type of cancer worldwide. Approximately 90% of oral malignant neoplasms are squamous cell carcinomas (OSCC) [1]. Although OSCC may develop de novo, some of them originate from potentially malignant disorders (PMD). Epithelial dysplasia may be an early sign of potency for the future transformation into oral squamous cell carcinoma (OSCC). A routine biopsy and close observation are recommended for persistent white oral lesions. As frictional keratosis may mimic oral leukoplakia, the question arises: Is there a need for a biopsy of persistent white lesion of traumatic origin?

Materials and methods. Data from 643 oral tissue biopsies were retrospectively analyzed. A total of 176 (27.37%) results with provisional diagnosis of leukoplakia (36 cases), OLP (77 cases) and frictional keratosis (63 cases) were selected. Retrospective data collected included age, gender, smoking status, provisional and histopathological diagnosis. The data was analyzed to assess the prevalence of epithelial dysplasia and OSCC in terms of age, gender and smoking status.

Results. Five (2.84%) cases of OSSC were reported, all of them were graded as G1; four cases of OSCC were found in clinically defined leukoplakia lesions; one case of OSCC (1.3%) was found in OLP biopsy; epithelial dysplasia was reported in 5 lesions (2.84%) provisionally diagnosed as OLP (3 cases), and leukoplakia in 2 cases. No dysplasia or OSCC were found in the lesions diagnosed as frictional keratosis.

Conclusions. Epithelial dysplasia and OSCC may be found in leukoplakia or OLP lesions not initially suspected of any malignancy. In some cases, clinical features are not sufficient to diagnose a lesion without histopathology. Frictional keratosis is easily identified by clinicians, and may not require a biopsy in every case. Clinical and histopathological evaluation of the white lesions still needs improvement.

Key words
oral lichen planus, oral cancer, leukoplakia, oral epithelial dysplasia, frictional keratosis

Abstract
Introduction. Leukoplakia and oral lichen planus (OLP) are common diseases manifesting as white lesions that are considered potentially malignant disorders (PMD). Epithelial dysplasia may be an early sign of potency for the future transformation into oral squamous cell carcinoma (OSCC). A routine biopsy and close observation are recommended for persistent white oral lesions. As frictional keratosis may mimic oral leukoplakia, the question arises: Is there a need for a biopsy of persistent white lesion of traumatic origin?

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Conclusions. Epithelial dysplasia and OSCC may be found in leukoplakia or OLP lesions not initially suspected of any malignancy. In some cases, clinical features are not sufficient to diagnose a lesion without histopathology. Frictional keratosis is easily identified by clinicians, and may not require a biopsy in every case. Clinical and histopathological evaluation of the white lesions still needs improvement.

Key words
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white patch [12]. The most common sites of appearance are alveolar ridges, buccal and labial mucosa, but the tongue can also be affected. The prevalence of frictional keratosis is difficult to assess as it can be easily mistaken with leukoplakia. Some reports suggest that it may appear in about 5% of the population [2]. After elimination of the possible mechanical irritation, the lesion should disappear within a period of 4–8 weeks. There do not appear to be any reports of malignant transformation in frictional keratosis.

Clinically, leukoplakia and lichen planus, especially the plaque type, could mimic each other and cause diagnostic difficulties. Furthermore, a similar clinical appearance can be seen in cases of frictional keratosis [13]. As both leukoplakia and OLP are PMDs, clear criteria are needed to identify those lesions that might progress to OSCC. This study aims to assess the rate of epithelial dysplasia and oral squamous cell carcinoma found in routine histopathology of white lesions.

MATERIALS AND METHODS

The records of patients who had been referred for diagnosis and treatment of oral pathologies in the period January 2013 – December 2019 were examined, from which cases with a provisional diagnosis of leukoplakia, OLP or frictional keratosis were selected. Leukoplakia was diagnosed by finding a symptomless white lesion, mainly in a smoking patient, the presence of which could not be explained by other means [7]. Oral lichen planus was diagnosed by finding papules that formed a network of white lines along which may also occur red areas of atrophy or erosions [11]. Frictional keratosis was diagnosed as white patches in areas prone to irritation, such as alveolar ridges, cheeks or lips, where an irritating factor was strongly suspected [12]. Retrospective data collected included age, gender, smoking status, provisional and histopathological diagnosis. Biopsy results that were not definitive and patients’ records with missing data were excluded from the study. Patients with a strong initial suspicion of OSCC were also excluded, and were referred to the Department of Oral and Maxillofacial Surgery to ensure a quick diagnostic process.

The data was analyzed using descriptive statistics to assess the prevalence of epithelial dysplasia and OSCC in terms of age, gender, and smoking status, by means of Statistica software.

RESULTS

Clinical and histopathological data related to 643 consecutive biopsies from the oral cavity were retrospectively analyzed. A total of 176 (27.37%) biopsy results with clinically determined white lesions that constituted 36 leukoplakia cases, 77 OLP cases and 63 frictional keratosis cases were selected. The study group included 125 (71.02%) females and 51 (28.98%) males. Mean age at presentation was 61.94 ± 14.84 years for males. The youngest patient was aged 18 years, and the oldest was 87 at the time of biopsy. Smoking habit was reported in 76 patients (43.18%).

Table 1 shows the histopathology of all cases. Among all histopathological diagnoses, 5 (2.84%) cases of OSCC were reported, all of them were graded as G1. OSCC was found on the lips (2 cases – 1 upper, 1 lower), floor of the mouth (2 cases) and hard palate (1 case). Four of five cases of OSCC were found in clinically defined leukoplakia lesions; thus, the prevalence of OSCC in leukoplakia was 11.11%. One case of OSCC (1.3%) was found in OLP biopsy. Three of four cases of OSCC in leukoplakia lesions were smokers, and the patient with a provisional diagnosis of OLP was a non-smoker. Clinical and histopathological diagnoses overlap and its relation to the cases of dysplasia/OSCC are presented in the Table 2.

Epithelial dysplasia was reported in 5 white lesions (2.84%), three of which were provisionally diagnosed as plaque type OLP (Tab. 3). However, the histopathological picture was lacking typical OLP features, and was subsequently considered as epithelial hyperplasia in leukoplakia. The other 2 cases were provisionally diagnosed as leukoplakia, which was not denied by histopathology (Tab. 4, Fig. 1). Four of five patients with dysplasia were smokers. Dysplastic lesions were located on the cheek (4 cases) and gingiva (1 case).

### Table 1. Histopathological diagnoses of all 176 white lesion biopsies

| Histopathological diagnosis | Total No. | Percentage of cases |
|----------------------------|-----------|---------------------|
| OLP                        | 76        | 43.18%              |
| Epithelial hyperplasia      | 6         | 3.41%               |
| Epithelial dysplasia        | 5         | 2.84%               |
| OSCC                       | 5         | 2.84%               |
| Frictional keratosis        | 84        | 47.73%              |

### Table 2. Distribution of white lesions, histopathological agreement, dysplasia, OSCC and smoking status (n=176)

| Provisional diagnosis | Total No. | H/P confirmation | Cases with dysplasia | Cases of OSCC | Smokers |
|-----------------------|-----------|------------------|----------------------|---------------|---------|
| Leukoplakia           | 36        | 8 (22.22%)       | 2 (5.56%)            | 4             | 32      | (88.89%) |
| OLP                   | 77        | 64 (83.12%)      | 3 (3.9%)             | 1 (1.3%)      | 19      | (24.66%) |
| Frictional keratosis  | 63        | 59 (93.65%)      | 0                    | 0             | 25      | (39.68%) |

### Table 3. Histopathological diagnoses of 77 biopsies provisionally diagnosed as OLP

| Histopathological diagnosis | Total No. | Percentage of cases |
|----------------------------|-----------|---------------------|
| OLP                        | 64        | 83.17%              |
| Epithelial dysplasia       | 3         | 3.9%                |
| OSCC                       | 1         | 1.3%                |
| Frictional keratosis       | 9         | 11.69%              |

### Table 4. Histopathological results of 36 biopsies provisionally diagnosed as leukoplakia

| Histopathological diagnosis | Total No. | Percentage of cases |
|----------------------------|-----------|---------------------|
| OLP                        | 8         | 22.22%              |
| Epithelial hyperplasia     | 6         | 16.67%              |
| Epithelial dysplasia       | 2         | 5.56%               |
| OSCC                       | 4         | 11.11%              |
| Frictional keratosis       | 16        | 44.44%              |
No case of OSCC or epithelial dysplasia was found in the biopsy of persistent frictional keratosis (Tab. 5).

Table 5. Histopathological diagnoses of 63 biopsies provisionally diagnosed as frictional keratosis

| Histopathological diagnosis | Total No. | Percentage of cases |
|-----------------------------|-----------|---------------------|
| OLP                         | 4         | 6.35%               |
| Frictional keratosis        | 59        | 93.65%              |

**DISCUSSION**

The study focused on determining OSCC and epithelial dysplasia rates in oral white lesions with no initial suspicion of malignancy. The lesions were subjected to routine biopsies to confirm or deny provisional diagnosis. When a potential causative factor was detected (smoking, trauma, candida infection), the lesion was observed for the 2–4 weeks needed to obtain a marked improvement or resolution. Cases of strong suspicion of OSCC were referred directly to the Department of Oral and Maxillofacial Surgery for further diagnostics and treatment. The most common OSCC location was the tongue, followed by the lip and floor of the mouth [14]. In this study, a similar frequency was reported for lip and floor of the mouth, while no case of tongue OSCC originating from a white lesion was reported. A recent study reported that different clinical types of leukoplakia were clinically misdiagnosed in 16.52% of cases, while malignancies were detected in 6.96% of cases when it was not initially suspected [15]. In the current study, the frequency of OSCC in leukoplakia was found to be even greater (11.11%), while the frequency of OSCC in the total number of white lesions subjected to histopathology was 2.84%. However, this frequency is comparatively low when compared to other study in which the dysplasia rate was 10.4%, and OSCC was found in 14.3% of all white lesions removed [16].

OLP is a PMD with approximately 1.2–4.9% rate of transformation. The known risk factors are smoking, alcohol consumption, HCV infection and male gender [17]. In the current study, one case of OSCC (1.3%) was found in OLP biopsy. This is in accordance with a recent meta-analysis reporting that malignant transformation of OLP was determined in 1.14% of cases [18].

Since OSCC may originate from PMD, such as leukoplakia or OLP, these lesions are routinely subjected to biopsy and histopathology. Some cases of leukoplakia and plaque type OLP are often confusing to clinicians. For leukoplakia, there are no specific clinical features to differentiate from other white lesions, and histopathological diagnosis is not always decisive. Banoczy et al. attempted to establish the typical histological changes present in leukoplakia lesions. These include abnormalities in keratinization (para-, hyper-, dyskeratosis) (Fig. 2), although neither are indices of premalignancy, changes in epithelial thickness (atrophy) and inappropriate epithelial maturation (dysplasia). In connective tissue, inflammation and hyaline and elacine degeneration were most commonly found [19]. However, some pathologists might deny a diagnosis of leukoplakia in the absence of epithelial dysplasia, which is in disagreement with some recommendations from the dental literature [7]. Moreover, the histopathological picture sometimes cannot be decisive, partially because dysplastic changes are not always visible in the specimen obtained as an incisional biopsy from a large leukoplakic lesion.

The diagnosis a plaque type OLP should be based on the biopsy and histopathology, as microscopic features are usually specific (Fig. 3). The histopathological aspect of lichen planus is described as follows: ‘In the area of clinically visible papules, the epithelium is thickened, with acanthosis and hyperparakeratosis and liquefactive degeneration of the basal layer. Round or ovoid ’colloid bodies’ appear mainly in the spinous layer of epithelium and in the lamina propria. Also a well-defined inflammatory zone in the connective tissue is present’ [3]. However, some samples may not show typical features and final diagnosis should be made by evaluating both clinical and histopathological aspects [9].

The presence of epithelial dysplasia is commonly used to evaluate the malignant potential of PMDs. It has been suggested that DNA ploidy analysis or p53 expression may be a more precise method of predicting malignancies, as cases of DNA aneuploidy and mutations in p53 show a higher risk of malignant transformation [4, 20]. Moreover, Nagler et al. found that biomarker levels in saliva, such as CA125, tissue polipeptide antigen and Cyfra 21–1, were significantly elevated in oral cancer patients compared to healthy controls [21].

Although oral leukoplakia and frictional keratosis often show similar clinical features, frictional keratosis is not believed to be a PMD. Histological features include hyperparakeratosis and peeling surface with fissures and clefts (Fig. 4). In most cases, lesions are colonized with
bacteria and show no signs of inflammation or epithelial dysplasia [22]. In the current study, no case of OSCC or epithelial dysplasia was found in the biopsy of persistent frictional keratosis, confirming that trauma-induced white lesions have no potential for malignant transformation. The biopsies were taken only when the lesion was not markedly improved or completely healed within the period of 2–4 weeks following elimination of the causative factor. This finding raises the question of unnecessary biopsies in such cases, and clearly demonstrates the need for determining a non-invasive method for white lesion diagnostics. However, Chi et al. reported that some of the biopsied samples of alveolar ridge keratosis (ARK) showed dysplasia [23]. Dysplasia was found only in the patients with confirmed tobacco smoking
and alcohol consumption; therefore, it was suggested that leukoplakia can mimic ARK, and biopsy of any persistent white lesions is recommended, especially, when other risk factors are present [23].

White lesions are commonly found in the oral cavity and can cause diagnostic difficulties. According to epidemiological studies performed on 17,000 Americans, white lesions were found in 27.9% of individuals [24]. Oral lesions, in spite of similar appearance, may be the result of various diseases, ranging from mechanical trauma, through precancerous lesions to squamous cell carcinoma [25]. Despite having typical and benign clinical features, a biopsy is advised of an oral white lesion when a patient presents himself for the first time. Familial history, trauma, drug intake, habits and patient’s complaints should all be included in the provisional diagnosis process [26, 27]. If the lesion is asymptomatic, clinician is often the first person to diagnose the disease. If the symptoms are present, it is important to collect relevant information about the condition: the date of first appearance, localization, duration, previous treatment and patient’s habits [26]. White lesions for which a possible causative factor has been identified, e.g. smokers’ lesion or frictional keratosis could be observed for 4–8 weeks after removal of the suggested cause to achieve spontaneous resolution. However, even such a period is already a long one in the case of a squamous cell carcinoma, a carcinoma in situ or severe epithelial dysplasia. Therefore, in the case of non-homogenous, symptomatic lesions, a biopsy is strongly recommended before elimination of possible causative factors and observation [7].

CONCLUSIONS

Clinical evaluation of white lesions still needs improvement and the histopathological criteria of the diagnosis require further standardization, related not only to dysplasia. In some lesions, oral cancer develops even without dysplasia in earlier biopsies; for this reason, a regular clinical and histopathological evaluation allows finding a potential malignancy at an early stage [28]. Further research of samples that showed features of malignancy might help in developing new diagnostic methods that should be precise and rapid. Molecular biomarkers may be valuable in the evaluation of leukoplakia for both diagnostic and prognostic aspects. So far, no method employing biological markers is available, but in future, the monitoring of many genes simultaneously might allow the prediction of the risk of malignant transformation of precancerous lesions. These methods might be more reliable than those depending on the monitoring of epithelial dysplasia [28]. A non-invasive tool that would help to reduce the number of unnecessary biopsies would greatly improve the cost-effectiveness, as well as patient compliance with screening programmes for high-risk patients. Moreover, such a tool might enable cost-effective screening of the general population. So far, every case of atypical white lesions of unknown origin or those that do not respond to treatment, should be biopsied to ensure early detection of potential malignancy.

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