Significance of Expression of Cyclin D as an Early Indicator in Dysplastic Transformation of Oral Mucosa in Tobacco Users

OBJECTIVES: To observe the expression of Cyclin D in transition of normal oral mucosa to dysplastic lesions and to find out the possible association of immunostaining in normal oral mucosa and different grades of oral dysplasia.

METHODOLOGY: In this cross sectional analytical study, total of 120 diagnosed paraffin embedded blocks were included comprising of 60 samples of normal oral mucosa (Group 1) and 60 cases of various grades of oral epithelial dysplastic lesions (Group 2). Patient’s record files were reviewed for age, gender and tobacco habits. Immunohistochemistry was performed by using Cyclin D monoclonal antibodies on all the tissue samples. Staining with Cyclin D was observed in each of the cases to find out their possible association as early indicator of transition from normal mucosa to oral dysplasia.

RESULTS: In Group 1, 45/60 (75%) patients were negative for Cyclin D. In Group 2, 40/60 (66%) were negative for Cyclin D. We found non significant association for Cyclin D staining in transition of normal oral mucosa to low grade lesions. But significant association was found in Cyclin D positivity in transition from normal mucosa to high grade dysplastic lesions.

CONCLUSION: We found no association of Cyclin D as diagnostic marker between normal and early dysplastic lesions, but the expression for Cyclin D was shown to be increased with increasing irreversible grades of dysplasia ie: from normal oral mucosa to severe dysplasia.

KEY WORDS: Cyclin D, Normal oral mucosa, Oral lesions, High grade lesions, Low grade lesions.

INTRODUCTION

Most common malignancy found in oral cavity is oral squamous cell carcinoma (OSCC) forming about 3% of total head and neck neoplasm. They usually follow a premalignant progenitor termed as oral epithelial dysplastic lesions (OED). These include red and white lesions such as erythro-leukoplakias, oral submucus fibrosis, actinic cheilitis, chronic candidiasis and not the least oral lichen planus (OLP). It is documented that OED have malignant transformation rate of about 0-20%. Because of high prevalence of OSCC, these must be diagnosed before they progress to carcinoma in-situ or invasive carcinoma as suspected dysplastic lesions are usually associated with history of prolonged tobacco use, betel nut and areca nut chewing, alcohol intake, chronic irritation and oral infections. If OED associated with these chronic irritations are screened and diagnosed in reversible stage, it would definitely encourage the patient to change their lifestyle and eating habits so that oral lesion could be managed without progressing to irreversible OSCC. Suspected oral lesions are evaluated histologically for identification of dysplasia. According to pathologists, dysplasia is graded into three types based on the level of differentiation. Low grade
dysplasia rarely leads to cancer. Moderate grade dysplasia involves the epithelial layers up to the two third of its thickness and are considered at borderline. High grade dysplasia is a type of early cancer that has high risk of becoming malignant but has not yet spread and is isolated within the basement membrane. Although routine hematoxylin and eosin staining gives sufficient idea of dysplasia grading, but immunohistochemistry is being widely used to observe more accurate cellular details as well as intracellular changes. Immunohistochemistry (IHC) is helpful in categorizing the tumor type, its progress and it shows the therapeutic usefulness. Immunoreactivity pattern with nuclear or cytoplasmic components plays a key role in investigating the early location of cellular disturbance. Among different immunomarkers being widely used for OSCC, Cyclin D is focused by researchers which has proven its role in carcinogenesis. It acts as cell cycle regulator. Variety of cyclins are being used for diagnostic as well as prognostic marker. Among them Cyclin D is a key regulator of G1-S phase of cell cycle and its activation is controlled by Cyclin dependent kinase inhibitors (CDKI). Cyclin D has been shown to get amplified in B cell lymphomas and oral cavity cancers. According to literature there is strong evidence proving advanced OSCCs a result of disturbed amplification of Cyclin D which causes shortened G1-S phase and uncontrollable cell growth. However little is known for the overexpression of Cyclin D in oral dysplastic lesions and its role as anticancer drug by blocking cellular proliferation. Early diagnosis of oral dysplastic lesion is an important step in abstaining tobacco chewing habit and can improve patient survival. Therefore this study began with the aim to observe if there is any altered expression of Cyclin D in transition of normal oral mucosa to dysplastic oral lesions in tobacco users.

**METHODOLOGY**

This was a cross sectional analytical study, which started after approval from institutional ethical review committee and was done in Department of Pathology Al-Tibri Medical College and Hospital, Karachi and Department of Histopathology Dow University of Health Sciences Karachi. This study started in March 2018 and was ended in November 2018. The samples were collected by nonprobability convenient sampling technique. Samples were grouped as 60 diagnosed samples of normal oral mucosa as group 1 and 60 diagnosed cases of different grades of OED lesions as group 2. All the grades of group 2 were further labelled as a, b and c according to increasing grade of dysplasia i.e. mild dysplasia as Group 2a (n=20), moderate dysplasia as Group 2b (n=21) and severe dysplasia as Group 2c (n=19).

Biopsy specimen from individuals who received radiotherapy or chemotherapy were excluded from the study. A total of 120 diagnosed paraffin embedded blocks were retrieved from histopathology laboratory of Dow University of Health Sciences Karachi. Institutional permission and approval was taken to use paraffin embedded histopathological blocks for research purpose. The study material was taken from biopsies received from patients who were treated between the year 2010 to 2017 and their medical records were studied for confirmation of tobacco usage. Histopathological grading was done by using H&E staining by experienced pathologist for reconfirmation for various grades of group 2.

For IHC 5microns thick sections were cut from each of the blocks. Manual's instructions were followed for IHC protocol. The validity of immunohistochemistry staining was provided by using positive controls which was B cell lymphoma for Cyclin D. Negative controls were taken by omitting primary antibody. The antibodies used was Anti Cyclin D (clone P2D11F11, RTU-CYCLIN-GM, dilution 1:100 Dakocytomotion).

Qualitative method which depended upon subjective perception and judgment was chosen for evaluation for immunoreactivity. Stained slides were observed using light microscope under x40 for evaluation of stained area and under x100 for evaluating intensity of staining. Random 10 fields were selected to visualize positive and negative stained cells. To avoid misleading interpretations, only homogenous staining was evaluated. For heterogeneous staining, dominant pattern of staining was chosen. Any light yellow to dark brown nuclear staining of at least 10% of focused area was considered as positive. For negative staining showing background stain only or barely visible intra nuclear brown staining in <10% area was considered.

SPSS IBM version 20 was used for data entry and analysis. Qualitative variables were calculated as frequencies and percentages. Chi-squared ($\chi^2$) test was used to compare qualitative variables. For quantitative variable like age mean and standard deviation were calculated. The p value ≤0.05 was considered as significant.

**RESULTS**

Demographical variables with frequencies and percentages are summarized in Table-1. All of the oral dysplastic cases were tobacco users of varying frequency. Most of the mild and moderate oral dysplastic cases belonged to regular tobacco users i.e. 34/41(83%) and rest of them were occasional tobacco users i.e. 7/41 (17%). However all of the severe dysplastic cases were regular tobacco users. We also found positive association of degree of dysplasia with tobacco history (p=0.047). Among group 1, we found...
majority of samples were negative for Cyclin D. Similarly in group 2, most of samples were negative for Cyclin D. These frequencies and percentages are shown in Table-2 and illustrated in Figure-1. According to Table-3, we further found non-significant association for Cyclin D staining in transition from normal oral mucosa to early dysplastic lesions. However significant association was seen for Cyclin D staining in transition of normal mucosa to high dysplastic lesions.

DISCUSSION

Various studies have been done on oral epithelial dysplasia with the objective of knowing diagnosis and prognosis. In our study 80% oral dysplastic samples belonged to males with M:F 4:1. Similar results were shown in the study of Juneja S in 2015 which showed 53% of their dysplastic
samples belonged to males. However, in contrast Jaber M in 2010, showed that oral dysplasia was common in both male and females equally with M:F 1:1. This might be due to larger sample size and westernized environment where their study was conducted. In our study most of the oral dysplastic cases were distributed in middle aged patients with mean age 45±5 years. Similar results were shown in study done Ho PS in 2009 which showed predominance of oral dysplasia in middle aged patients. This similarity might be due to similar sample size with similar gender distribution in their study. Gopinath D et al in 2016 showed evidence of oral dysplasia in individuals with mean age 51±17 years in males and 56±12 years in females. A study done in 2014 by Starzynska A et al found dysplastic lesions were most commonly found in patients with age range of 51-60 years. In this study strong history of chewing tobacco was seen in 88% in oral dysplastic cases with 12% cases from occasional tobacco users but nonsignificant association was shown in tobacco use and grade of oral dysplasia (p=0.136). These results were in concordance with the findings of Shetty P in 2017 showing evidence of dysplasia in individuals with low to moderate tobacco consumption. Similar findings were reported by Mohiuddin S in 2015 showing positive association of dysplasia with smokeless tobacco and areca nut. The present study showed 75% cases with negative scores for Cyclin D and 25% samples showing mild staining in normal oral mucosa. It has been previously shown in studies done by Bogoz B in 2012, Mishra R in 2009 that Cyclin D is totally negatively expressed in normal oral mucosa. However they used smaller sample size and all normal oral mucosal samples were strictly recruited from tobacco users. In contrast to our results, study done by Ramakrishna A in 2013 showed 40% of normal oral mucosal samples with moderate staining but the sample size was smaller than the present study. Our study showed 67% oral dysplastic lesions negative for Cyclin D which was in similarity with study done by Lam KY and colleagues in 2000 proved faint expression of Cyclin D is only limited to few high grade oral lesions and all of the OSCC cases. He further commented that this finding was because of shorter half life of Cyclin D during G1-S phase of cell cycle. This finding was further reinforced by the evidences provided by Mishra R in 2009 showing no immunoreactivity with Cyclin D in normal and potentially malignant lesions. Our study showed non-significant association of Cyclin D expression in various grades of oral dysplasia which was reinforced by the results of Turrati E in 2005. According to him Cyclin D was totally negative in low grade dysplasia and was positive in only 8% of high grade dysplasia. In present study Cyclin D expression was reported in 33% of dysplastic lesions and significant association (p=0.08) was observed in transition of normal mucosa to severe dysplasia which coincided with previous study done by Ramasubramanian A et al in 2013 showing Cyclin D positivity for dysplastic lesions and its strong expression was favored for irreversible grades of dysplastic lesions (p=0.001). Our study revealed 90% cases of mild dysplasia with negative expression of Cyclin D, moderate dysplastic cases showed 66% cases with negative expression for Cyclin D and 33% cases showed mild expression. Similarly severe dysplastic lesions showed negative expression for Cyclin D in 42% cases as well as positive expression was observed in 58% cases. In 2017 Patel SB and colleagues investigated all of the mild dysplasia cases with mild expression of Cyclin D. Moderate dysplastic cases showed 66% of positivity. Similarly severe dysplasia showed positive expression in 92% cases. These dissimilarities might be due to smaller sample sizes used in their studies.

CONCLUSION

Oral squamous cell carcinoma treatment in its late presentation, end up with high rates of mortalities and diminished chances of survival. It is not surprising that early oral epithelial dysplastic lesions are curable. Therefore it is a real need to imply methods for early detection of oral dysplastic lesions especially in those with risk factors like tobacco use. Cyclin D showed non-significant association in transition from normal mucosa to early dysplastic cases and hence it was not found as reliable diagnostic marker in early oral dysplastic lesions but strong and significant amplification of Cyclin D was seen in transition from normal oral mucosa to severe dysplasia which showed its usefulness as diagnostic tool in high grade dysplastic lesions.

CONFLICT OF INTEREST

None declared

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