Quantitative estimation of AgNORs in normal, dysplastic and malignant oral mucosa

Aman Chowdhrya, Revati Shailesh Deshmukhb, Deepika Shuklaa, Deepika Bablani, Shashwat Mishrac

Aims. Silver stainable nucleolar organizer regions (AgNORs) have received a great deal of attention recently as their frequency within the nuclei is significantly higher in malignant cells than in normal, reactive or benign neoplastic cells. The objective of this study was to carry out a quantitative assessment of large and small AgNORs in oral normal mucosa, precancerous lesions and infiltrating squamous cell carcinomas.

Methods. The study comprised 110 formalin-fixed, paraffin-embedded oral mucosal biopsies consisting of 30 oral dysplasia, 60 oral squamous cell carcinomas and 20 normal oral mucosa. AgNORs were counted in each nucleus, categorized as small, large and total number of AgNORs in each cell and their means were calculated.

Results. The mean value of small AgNORs, large AgNORs and total AgNORs increased gradually from normal mucosa to dysplastic lesions to squamous cell carcinomas. The study clearly indicates that in oral squamous cell carcinomas, AgNORs diminish in size as they increase in number. Further, AgNOR counts increase as the degree of malignant potential of the cell increases.

Conclusions. By combining both the enumeration of AgNORs and their size, good distinction can be made between normal, dysplastic and infiltrating squamous cell carcinomas. This could help in the early diagnosis and prognosis of dysplastic mucosal lesions and their malignant transformation.

Key words: silver stainable nucleolar organizer regions (AgNORs), oral dysplasia, squamous cell carcinomas

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INTRODUCTION

Millions of new cases of invasive cancer are diagnosed each year of which oral cancer has a considerably high incidence. This high incidence of oral cancer and its attendant morbidity and mortality has instigated workers to devise methods for its early diagnosis. Nucleolar organizer regions (NORs) are segments of chromosomes encrypted for ribosomal RNA (rRNA) which are present on specific loops of DNA (ref.1). NORs have received a great deal of attention recently because of the observations that their frequency within the nuclei is significantly higher in malignant cells than in normal, reactive or benign neoplastic cells2-3. Consequently, they are of diagnostic value for in characterization of the invasiveness in carcinomas4. They also play a role in the estimation of the cellular activity that is applied to a variety of neoplastic or hyperplastic lesions5-6. Since some of the NOR associated proteins are argyrophilic and can be demonstrated as black dots by a silver staining technique, such demonstrated structures are known as AgNORs (ref.7). The exact biochemical nature of these proteins is yet to be defined: they have been characterized as B 23, C 23 (ref.8) and RNA polymerase 1 (ref.9) and are attributed to acidic, non-histone constituents10.

According to studies, increased expression of NOR sites is expected in actively proliferating cells and they are visualized in those cells which are functionally active in the preceding interphase. Counting AgNORs in interphase nucleus may assist in diagnosing and grading malignant neoplastic lesions5-9. Moreover, malignant cells have larger AgNOR than benign cells10. The increased AgNOR size could be attributed to increased transcriptionally active ribosomal sites. Malignant cells are characterized by small size, larger number and scattered distribution of AgNORs (ref.11). AgNOR silver staining technique enables the distinction to be made between certain grades of malignancy and may enable prognostic assessment12. AgNOR technique, with its cost-effectiveness and easily reproducible results, can be very effective in resource-poor settings.

The endeavor of this study was to carry out a quantitative assessment of large and small AgNORs in specimens of oral normal mucosa, precancerous mucosal lesions and infiltrating squamous cell carcinomas previously diagnosed on H & E stain.
MATERIAL AND METHODS

30 samples with dysplastic lesions of the oral cavity and 60 samples of well differentiated carcinomas were selected from the archives of Department of Oral Pathology & Microbiology, Bharati Vidyapeeth Dental College & Hospital, Pune, for which histopathological diagnosis and reporting had been done previously. As controls, 20 normal oral mucosa biopsies were taken from twenty consenting, healthy volunteers without any tobacco habit/smoking. Samples were from different intra-oral sites i.e. buccal mucosa, alveolus, tongue, palate, retromolar area, vestibule and floor of mouth. 82 men and 28 women patients (age ranging from 21 to 75 yrs) participated in the study. Using hematoxylin-eosin (H&E) stained sections, the cases were divided into the following three groups.

Group 1: normal oral mucosa (n=20).

Group 2: mild, moderate and severe dysplasia, histopathologically graded by two oral pathologists evaluating independently (to avoid observer bias in the grading of dysplasia) based on established criteria by Pindborg and Smith 1997 (ref.13) (n=30).

Group 3: histopathologically diagnosed well differentiated squamous cell carcinomas (n=60).

Two sets of sections were cut from each block at a thickness of 3-4µ. One set of sections was stained by Ehrlich’s Hematoxylin and Eosin for confirmation of the previous histopathologic diagnosis. All H&E-based diagnoses were independently verified by two experienced oral pathologists. Cases with ambiguity in the histopathological diagnosis were excluded. Following this, the other set of sections was stained for the Argyrophilic Nucleolar Organizer regions by an improved method of Ploton et al 1986 (ref.4) and the AgNOR count was established for each specimen. The histopathological diagnosis of each specimen was then compared with the characteristics evident on AgNOR staining.

AgNOR staining and counting:
AgNOR staining was performed according to the procedure established by the Committee on AgNOR quantification incorporating recommended safeguards to reduce variability and bias in staining and counting15. Each section was immersed in sodium citrate buffer (at pH 6.0) incubated in wet autoclave at 1208 °C for 20 min,

Fig. 1. Photomicrograph (a) showing Small AgNORs (100X; AgNOR stain), (b) showing Large AgNORs (100X; AgNOR stain).
then allowed to cool down to 378 °C. The slides were then immersed in a freshly prepared silver staining solution containing one part by volume of 2% gelatin in 1% formic acid, two parts of 25% aqueous silver nitrate solution at 378 °C in a thermostatically controlled environment for 11 min. The slides were then washed with double-distilled de-ionized water and the reaction was stopped. All stained sections were dehydrated in increasing grades of concentration of ethanol, and then clarified in xylene. All AgNOR quantification was performed by one of the authors.

Quantitative analysis: All stained sections were examined under oil immersion (100X) objective lens. In each of the sections, 100 individual cells were examined for methodical and systematic quantification of AgNORs, using a 100X oil immersion objective and 10X eye piece giving a total magnification of 1000X. Fields were selected at random and 100 cells were examined continuously. Each nucleus was examined for number of nucleoli. A nucleolus was defined as an AgNOR dot and the mean number of AgNOR was calculated.

In all sections, the argyrophilic, NORs were clearly evident as black “dots” or “blebs” of varying size, in the brown stained nucleus on a pale yellow background. The overall slide background was clear without any extraneous silver deposits. The black dots, which were visible in each cell, were evaluated depending upon the size into “large” NORs (>=2m) and “small” NORs (<2m). No counterstain was used to avoid obscuring the NOR dots. The AgNORs were counted as per the standardized guidelines recommended by Crocker et al\textsuperscript{12}. When AgNORs were seen in groups, each cluster was counted as one AgNOR. When AgNORs could be seen separately but large enough to occupy a substantial area, it was counted as one “large” NOR, and the smaller NORs which were distinctly visible, were counted as one “small” NOR.

AgNORs were counted in each nucleus and categorized as large NORs which appeared as blebs and small NORs which appeared as distinct small dots. Further, the total number of AgNORs in each cell was calculated by summing both counts and was initially recorded in a separate table. From these values the means of large, small and total AgNORs were obtained and tabulated.

Statistical analysis was performed using SPSS software version 14.0 (Statistical Package for Social Sciences). The results processed using unpaired students “t” tests. Values for $P$ less than 0.05 were regarded as significant.

RESULTS

The maximum number of squamous cell carcinomas was in males and the most common site being the buccal mucosa.

NORs were strictly located within the cell nucleus and were distinctly stained in black as dots and blebs that were

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<th>Table 1. Mean values of small, large and total AgNOR counts in Normal, Dysplasia and carcinoma patients.</th>
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N- number of patients, Min- minimum value, Max- maximum value, S.D.- Standard deviation
either round or elongated: the rest of the nucleus stained yellow-brown. The mean number of small NORs (Fig. 1a), large NORs (Fig. 1b) and total AgNORs, calculated for normal, dysplastic and squamous cell carcinoma cases is shown in Table 1.

In all categories of specimen, both small and large AgNORs were most common in the basal layers and became progressively less frequent in the superficial layers.

As observed throughout, there was an increase in number of small AgNORs in squamous cell carcinomas (mean 3.25) compared to that of dysplastic lesions (mean 2.05) and normal mucosa (mean 1.17) (Table 1, Fig. 2). The large AgNORs were increased in dysplastic lesions (1.71) and squamous cell carcinomas (1.94) as compared to normal mucosa (0.78) but the difference is not as remarkable as seen with the small AgNORs.

### Statistical Significance of AgNORs

In comparison to normal mucosa, a highly significant increase in small, large and total AgNORs was seen in dysplastic lesions and squamous cell carcinomas ($P<0.05$). Squamous cell carcinomas also showed a significantly higher frequency of AgNOR than dysplasia. Comparing large AgNORs in dysplastic lesions and squamous cell carcinomas, a nonsignificant increase in the mean values was noticed in squamous cell carcinomas. When mean values of total AgNORs were tallied between dysplastic lesions and squamous cell carcinomas, a significantly higher frequency of AgNOR in squamous cell carcinomas was found ($P<0.05$) (Table 1).

Of the different sites of occurrence, except for the palate, the mean frequency of large AgNORs in squamous cell carcinomas was statistically higher than in normal mucosa ($P<0.05$). However, the mean values for small and the total AgNOR counts showed a highly significant increase irrespective of its site of occurrence ($P<0.05$).

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**Fig. 2.** Photomicrograph of (a) Normal Mucosa showing AgNOR staining (100X; AgNOR stain), (b) Dysplastic Mucosa showing AgNOR staining (100X; AgNOR stain), (c) Carcinomatous Mucosa showing AgNOR staining (100X; AgNOR stain).
DISCUSSION

The nucleolar organizer regions (NORs) have been recognized for over a decade as areas of rDNA, which transcribe to rRNA. The most commonly used method for staining the NORs in vogue today is the AgNOR staining method, which identifies NOR associated proteins (NORAPs) whose function in unclear. The AgNOR technique helped cytogeneticists with a simple tool for the study of these structures in tumor pathology. Quantitative estimation of interphase AgNORs is useful for evaluation of cell kinetics. Protein synthesis is faster in rapidly dividing cells. Accumulation of interphase AgNORs during the mitotic cycle is correlated with the increased demand for ribosomal biogenesis. Accordingly, an increase in the nucleolar structures (AgNORs), where rRNA synthesis takes place, is expected in rapidly diving cells. The AgNOR parameter is thus a convenient marker for studying the rate of proliferation of cells in tissues being studied histologically.

Hence, in the present study this technique was followed for its convenience and feasibility. We studied a range of oral epithelial lesions which are commonly encountered in the natural history of oral carcinoma in order to examine whether these lesions share common or dissimilar features in relation to AgNOR patterns and numbers. The present study also evaluated the significance of the frequency of small, large and total AgNORs. To the best of our knowledge, ours is the first study assessing the size of AgNORs and using it as criteria for distinguishing normal mucosa, oral dysplasia and oral squamous cell carcinoma.

In the present study, the frequency of small, large and total AgNORs increased with malignant progression from normal mucosa to squamous cell carcinomas of the oral cavity ((P<0.05; Table 1), in agreement with earlier report. This clearly indicates that the total AgNOR counts increase with the changes occurring in the normal epithelium towards a dysplastic change and its malignant transformation. Interestingly, increase in AgNORs counts is associated with a concomitant increase in biochemically assayed DNA in the affected cell. In human cells, the degree of argyrophilia of individual NORs on chromosomes is constant and inheritable. The size of silver stained NORs is also related to the degree of transcriptional activity of the rDNA concerned. Thus larger AgNOR sites on chromosomes are more numerous in transcriptionally active cells than those of smaller size. But this may not be true in interphase nuclei in malignant cells.

The small AgNORs were more frequent in cancerous mucosa at all sites, whereas, the large AgNORs showed no significant increase in carcinoma of the palate. Although, this could be artifactual due to small sample size, quantification of small AgNORs might have a greater discriminative value for distinguishing dysplasia and carcinoma.

Even though the reason for the greater number of AgNORs in the nuclei of malignant lesions is uncertain, it may represent an escalation in ploidy, increased gene amplification or a rise in chromosomal segregation with more cells being in the S phase of their cycle. In agreement with Derenzini et al., we also found an inverse correlation between malignant change and the size of AgNORs. Although AgNORs became more numerous with malignant progression from dysplastic lesions to frank carcinoma, their sizes became smaller. This may well represent increased transcriptional activity resulting in more rRNA gene copies with increasing proliferative activity.

CONCLUSION

Combining enumeration of AgNORs and size, we were able to reliably distinguish between normal, dysplastic and infiltrating squamous cell carcinomas. This could help in the early diagnosis of dysplastic mucosal lesions and their malignant transformation. It may also be applicable in the prognosis of prophylactically treated premalignant lesions and in treatment planning.

CONFLICT OF INTEREST STATEMENT

Author’s conflict of interest disclosure: The authors stated that there are no conflicts of interest regarding the publication of this article.

REFERENCES