**PAPERS**

**Leucoplakia of the oral cavity, smoking and arylhydrocarbon-hydroxylase inducibility**

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**Summary**

In a consecutive series of 53 cases of oral leucoplakia, smoking history, oro-dental status and arylhydrocarbon-hydroxylase (AHH) inducibility were assessed. There was an increased association of leucoplakia with smoking, particularly in the male cases, but no remarkable findings as to dental status. Overall, there was a slightly higher frequency of cases with high AHH inducibility than expected from a normal control population. This difference was confined to the smokers, where there was a statistically highly significant over-representation of high AHH inducibility and under-representation of low AHH inducibility. The figures in the non-smoking patients were close to the expected.

**Introduction**

The familial clustering of certain malignancies, such as carcinoma of the breast and carcinomas of the large intestine, has been known for a number of years and genetic factors have been reported to be of aetiological significance (Penrose, Mackenzie and Karn, 1948; Woolf, 1958). Tokuhata (1964), demonstrated in lung cancer a synergistic interaction between familial and smoking factors. These epidemiological data became explicable in biological terms when Kellerman and associates presented their data on arylhydrocarbon-hydroxylase (AHH) inducibility (Kellerman, Luyten-Kellerman and Shaw, 1973 a; Kellerman, Shaw and Luyten-Kellerman, 1973b). They observed a statistically highly significant over-representation of high and intermediate AHH inducibility in cultured lymphocytes not previously exposed to hydrocarbons in the patients with pulmonary carcinomas. All these patients were heavy smokers. The association between high and intermediate AHH inducibility, exposure to polycyclic aromatic hydrocarbons (PAH) and the occurrence of pulmonary carcinomas was putatively explained by AHH-mediated activation of PAH to epoxides (Huberman et al., 1971, 1972; Huberman and Sachs, 1977; Buty, Thompson and Slaga, 1976) or epoxide metabolites (Booth and Sims, 1974; Daudel et al., 1975), initiating malignant cell transformation. In *vitro* experiments have given evidence of a preferential binding of the PAH epoxide products to the guanine sequences of DNA (Grunberger et al., 1974), and it has even been postulated that this DNA-alteration might represent a biochemical basis of carcinogenesis (Korsgaard, 1979).

Subsequent studies were inconsistent with or even contradictory to the original report. There were difficulties in confirming the postulated model of genetically determined AHH inducibility in man (Gurtoo, Bejoa and Minowada, 1975; Nebert and Atlas, 1977). More recently, however, several studies including twin studies have verified that AHH inducibility is under genetic control (Atlas, Vessel and Nebert, 1975; Paigen, Gurtoo and Minowada, 1977). In clinical material, the findings of Kellerman’s group have been confirmed by some reports (Guirgis et al., 1976; Korsgaard et al., 1977). A clustering of smokers with spinocellular carcinoma...
of the larynx (Trell et al., 1976), oral carcinoma (Trell et al., 1978, 1981) and laryngeal precancers (Trell et al., 1980), into the high and intermediate AHH induction groups has also been demonstrated. The authors now report the findings of AHH inducibility in relation to smoking habits and oro-dental status in oral leucoplakia.

Material and Methods

Study group

All cases with a clinical diagnosis of oral leucoplakia (n = 102) diagnosed at the Department of Oral Surgery and confirmed by histopathological examination at the Department of Pathology, Malmö General Hospital between the years 1964–1978, were invited to a smoking interview and AHH assessment at the Department of Preventive Medicine, Malmö General Hospital, in 1978–1979; 57 (56%) of the cases responded to the invitation; 4 were excluded because the histopathological examination revealed carcinoma in situ. The retrospective study thus finally comprised 53 cases. In one case, the AHH sample was lost. All cases showed typical microscopical changes with hyperplasia and dysplasia of the epithelial cells, occasional hyperkeratosis but only slight to moderate, unspecific inflammation. In 7 cases, more pronounced metaplastic changes, atypias and mitoses in the epithelial cells were noted. There were 29 men aged 22–74 years (mean 49) and 24 women aged 32–72 years (mean 52).

Control group

The control group consisted of 118 healthy employees of the University Hospital, Lund, and various research workers who voluntarily offered blood samples for assay of AHH inducibility. Forty-one were smokers, 19 with a daily consumption exceeding 20 cigarettes. None of the control group had malignant disease at the time of examination or had previously undergone treatment for malignant neoplasm.

Enzyme assessment

Samples of venous blood (8–10 ml) were procured from each subject and kept overnight in heparinized tubes at +4°C. The blood was then diluted with Parker 199 medium and layered over a column of Lymphoprep®. After 40 min centrifugation at 400 g at room temperature, a distinct white ring on top of the Lymphoprep® column contained the lymphocytes (Böyum, 1968; Thorsby and Bratlie, 1970). These were washed in Parker 199, suspended in the same nutrient medium, and the lymphocyte yield was calculated by the standard Trypan blue dye exclusion test. After resuspension in Parker 199 complemented with fetal bovine serum, glutamine and antibiotics, the lymphocytes were stimulated with phytohaemagglutinin and pokeweed mitogen, initiating blast transformation, a modification of the method described by Busbee, Shaw and Cantrell (1972).
At the end of 72 hr in culture the lymphoblasts were tested for viability, divided into aliquots, and 1.5 mmol 3-methylcholanthrene in acetone was added to the test samples and allowed to take effect for an additional 24 hr. Enzyme activity was then assayed according to a modified procedure of Nebert and Gelboin (1968) and Busbee et al. (1972), and the degree of inducibility determined spectro-photofluorometrically as fold induction in the presence and absence of 3-methylcholanthrene.

Clinical assessment

At the time of the AHH analyses, all patients were asked to give a detailed account of the type, duration and consumption of tobacco up to the time of diagnosis of the oral leucoplakia.

All patient records were re-assessed with particular reference to oral hygiene, dental and denture status, location of oral leucoplakia and histological investigation of the leucoplakia lesions. Statistical significance was calculated according to the $\chi^2$ test with 2 degrees of freedom.

Results

Ninety per cent. of the male patients and 46% of the female were smokers. In general, the male smokers showed a higher tobacco consumption than the females.

There were no obvious associations between the histological appearance, size, shape or location of the lesions and age, sex or smoking habits of the patients.

Fourteen patients had dentures of some kind. These were functioning well and neither in the non-smokers nor in the smokers did mechanical irritation by the denture appear to be the cause of the leucoplaikias. The frequency of dentures in the material was not higher than expected in these age- and sex groups. In the control group the frequencies of high, intermediate and low AHH inducibility were 7-6, 41-5 and 50-9% respectively, which is in accordance with the data of Kellerman et al. (1973a) in a normal white U.S. population and in equilibrium with the Hardy-Weinberg law. There were no differences in the distribution in relation to smoking habits or sex. In the study group (Table 1) there were 9 cases with high AHH inducibility, 26 with intermediate and 17 with low inducibility. The frequencies expected from the control group were calculated as 5, 21 and 27 cases. When the material is divided into smokers and non-smokers, there is an obvious over-representation of individuals with high AHH inducibility among the smokers (8 against 3 expected cases) and also a slight under-representation of low values. These differences are significant ($P<0.005$) while the figures for the non-smokers are close to the expected.

Discussion

There is ample evidence that carcinoma of the oral cavity is one group of malignant diseases associated with heavy smoking (Graham, 1977; Pindborg, 1980). The same association with smoking is documented also in oral leucoplakia, which is regarded as the most frequent precancerous state of the mouth (Bánóczy, 1977; Pindborg, 1980). In the discussion of the aetiology of oral malignancy, ill-fitting prostheses, bad crowns and fillings also receive considerable attention (Thumfart et al., 1978). In a previous series of oral carcinoma (Trell et al., 1981), no remarkable abnormalities in the oro-dental and denture status were found, while smoking was present in all the male cases and 50% of the female cases. In the present consecutive series of oral leucoplakia, the oro-dental and denture findings were also unremarkable, smoking was recorded in 90% of the male cases and about 50% of the female cases. In a recent population investigation in Malmö, 48.6% of 50-year old males were found to be smokers (Janzon, et al., 1982).

The authors believe that these results add further support to the concept of AHH as an activator of proximate carcinogens, such as the PAH of tobacco smoke. However, it is hard to assess the spontaneous transformation rate of leucoplakia to carcinoma in the present material as this metabolic activation represents only one of several factors possibly involved in malignant transformation. Cessation of smoking might represent the most adequate prevention of malignant development in cases of oral leucoplakia where an association with smoking is apparent. The authors are continuously following their study group, but the present observation time is insufficient to draw definite conclusions on the further natural history in relation to factors like age, AHH inducibility, elimination or reduction of tobacco consumption and restorative oro-dental measures.

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References


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