Analyze the Expression of Cytokeratin 5 on the Epithelial Cells of the Buccal Mucosa in Batik Workers

Juni Handajani¹, Dhinintya Hyta Narissi²

¹Department of Oral Biology, Faculty of Dentistry, Gadjah Mada University, Sekip Utara, Bulaksumur, Yogyakarta, Indonesia
²Faculty of Dentistry, Gadjah Mada University, Sekip Utara, Bulaksumur, Yogyakarta, Indonesia

Abstract: The industry of batik is often use synthetic dyes as azo. Azo dyes contain naphthol and diazonium salts that are toxic to the tissue when exposed via inhalation, swallowing or direct contact. Some preliminary studies mentioned the exposure on the azo dyes batik could be expected to cause abnormalities in the buccal mucosal epithelial cell nuclei marked increase in the frequency of micronucleus, karyolysis, and pyknosis. Objective: the purpose of this study was to analyze the expression of cytokeratin 5 on the epithelial cells of the buccal mucosa in batik workers Yogyakarta. Material and Methods: the study included 30 male subjects were divided into 2 groups: 15 subjects exposed to azo dyes and 15 subjects as control. Criteria for subject were batik workers exposed azo dyes in The Unit of Coloring at batik home industry Yogyakarta, while control is not exposed to azo dyes. Buccal mucosal epithelial cells were swabbed using cytobrush. Analysis of the expression of cytokeratin 5 in the buccal mucosal epithelial cells was used monoclonal antibody cytokeratin 5 (Biocare Medical, USA) and immunohistochemical method (ABC Staining Kit, ImmunoCruz, Santa Cruz Biotechnology, USA). Data were analyzed using independent t-test. Results: the results showed the expression cytokeratin 5 of epithelial cells buccal mucosa was significantly higher in batik workers group than the control (p <0.05). Conclusion: It is concluded that exposed azo could increase the expression of cytokeratin 5 in the epithelial cells of the buccal mucosa batik workers in Yogyakarta.

Keywords: cytokeratin 5, epithelial cell, buccal mucosa, azo dye, batik worker

1. Introduction

Oral cavity is port d’entry of toxic and non toxic substance to the body, so susceptible to pathological change. Oral mucosa is demarcating oral cavity consisting of two basic layers that are separated by a basement membrane. Both the base layers are stratified squamous epithelium on the outer and inner layer of connective tissue (lamina propria). Some parts known to have a third layer (submucosal) are found between the lamina propria adjacent to the bone (palate) or muscles (cheeks and lips). Submucosal tissue is composed of loose connective tissue containing nerves and blood vessels, also glands salivarius.

Function of oral mucosal protects mechanically against shear and tensile stress (compressive and shearing strength) providing defense against microorganisms, toxins, and some antigens contributing to the immunological defense both humoral and cell-mediated. Salivarius glands in the oral mucosa have function to secrete saliva which has several roles, there are lubrication and buffer activity and provide some antibodies.

Variation regional of the oral mucosa is associated with the degrees and types of pressure during mastication, speech and facial expressions. Oral mucosal structure varies according to the thickness of the epithelium, the degree of keratinization, connective tissue interface complexity to the epithelium, lamina propria and the composition of the existing or the absence of submucosal. Classification of oral mucosa is divided into masticatory, lining and specialized. Buccal mucosa is categorized lining, can be distended and bonded to the surrounding tissue structure through connective tissue which is rich in elastin. Oral mucosa is consisting 60% lining, 25% masticatory and the remaining 15% specialized.

Characteristics of lining epithelium mucosa are not keratinized and slightly under pressure. Differences between lining epithelial cells and keratinization epithelium are the surface layer of cells lining epithelium has little or no keratohyalin granules, filigran protein and loricin, but contains involucrin.

Cytokeratin (CK) is a protein that contains keratin intermediate filaments. The function of this protein is a component of the cytoskeleton and to contact the cell (desmosome and hemidesmosome) in epithelial tissues. Each product cytokeratin gene family is divided into neutral or basic type II cytokeratin (numbers 1-8) and the acidic type I cytokeratin (numbers 9-20). Cytokeratin usually found in pairs and type I have a shorter size. Specific distribution of cytokeratin 5 and 15 in epithelial are usually confined to the basal and parabasal layers although cytokeratin 14 may also be expressed by suprabasal keratinocytes. Cytokeratin 1 and 10 are found in the suprabasal layers of masticatory mucosa. Mucosal lining, mainly suprabasal keratinocytes stained for CK 4 and 13. In the epithelium lining the soft palate expressed CK 7, 8, and 18.

Primary function of cytokeratin is to protect the epithelial cells of the pressure (stress) mechanical and non-mechanical resulting in cell death. The role of cytokeratin includes signaling when a cell responds to stress, apoptosis and other specific functions. Several human diseases are related to the alleged involvement of cytokeratin. Cytokeratin is increasingly widely used as a tumor marker for the purpose.
2. Material and Method

Subject and Study Design

The subject consisted of 30 men, divided into 2 groups, aged range 18-40 years old. They were 15 subjects exposed azo in The Unit of Coloring at batik home industry Yogyakarta and 15 as a control. Informed consent according to Helsinki II was obtained from each participant.

Intervention and Assessment

Approval ethical clearance from the Ethics Committee Faculty of Dentistry, Gadjah Mada University (Number: 00261/KKEP/FKG-UGM/EC/2015) on May 15, 2015. Subjects were asked to rinse prior to remove debris from the oral cavity. Cytobrush was moistened with 0.09% NaCl. Buccal epithelial cells were swabbed using cytobrush. Swab was done by turning the cytobrush in the direction of at least 360° on the right buccal mucosa. The same procedure was performed on the left buccal mucosa.

Immunohistochemical Staining

Samples were washed with PBS 3 times each for 5 minutes. The procedure was then performed using a blocking BSA 0.1% and 0.25% Triton for 20 minutes, then washed using with PBS 3 times each for 5 minutes. Incubation was carried out using a monoclonal antibody anti cytokeratin 5 (Biocare Medical, USA) for 24 hours at 4°C. Samples were washed using PBS 3 times each for 5 minutes.

Samples were stained using immunohistochemical method (ABC Staining Kit, Immunocruz, Santa Cruz Biotechnology, USA). Samples were incubated using a secondary antibody for 20 minutes in room temperature. The procedure followed by immersion in a substrate buffer for 20 minutes in room temperature.

Staining was used DAB around 20 minutes. Positive result was expressed brown colour on both nuclear and cytoplasmic cell. Samples were observed using a light microscope magnification of 200 times and a computer monitor with a magnification of 100 times. Each sample was collected at least 100 epithelial cells. Observation was done by counting positive cell of cytokeratin 5.

Statistical Analysis

The normal of the data and the homogeneity of variance were verified the Shapiro-Wilk and the Levene’s test respectively. The data of expression cytokeratin 5 epithelial buccal mucosa cells was then compared using independent t-test. In all the analysis, the level of significance was set at p<0.05
and was considered as significant. The calculations were handled with SPSS 12.0 software for Windows (SPSS Inc; Chicago, IL, USA).

3. Result

The expression of cytokeratin 5 was shown in cell nucleus and cytoplasm epithelial buccal mucosa cell (Figure 1). Positive expression of cytokeratin 5 was observed in both exposed to azo and control groups.

![Figure 1](image1.jpg)  
**Figure 1:** A. Expression of cytokeratin 5 was observed at the entire epithelial cell of buccal mucosa in exposed to azo group. B. Positive expression was shown brown in nucleus and cytoplasm (circle). There was vacuola in the cytoplasm cell.

![Figure 2](image2.jpg)  
**Figure 2:** Expression of cytokeratin 5 was in exposed to azo group (A), showed brown colour in both nucleus and cytoplasm, and vacuola in the cytoplasm (circle). The control group (B) observed brown colour only in the cytoplasm of the cell while the cell nuclei was blue.

Mean and standar deviation the number of expressed cytokeratin 5 in the epithelial buccal mucosa cells were shown in Table 1.

Table 1 showed mean the expression of cytokeratin 5 in the buccal mucosal epithelial cells of batik was higher than control. Normality data was calculated using the Shapiro-Wilk. The results of the normality data showed a group of exposed to azo in batik workers p = 0.964 and control p = 0.759. These results indicated the data were normally distributed (p > 0.05). Calculation of data homogeneity was used Levene's test showed p = 0.062 (p > 0.05) or data homogeneous. Furthermore, the data were tested using independent t-test p = 0.00 (p < 0.05). The result indicated the number of epithelial buccal mucosal cells that expressed cytokeratin 5 in batik workers were significantly different than the control.

4. Discussion

Observation of the effect of genotoxic, cytotoxic, an indication of exposure to chemicals, and toxic response can be performed using epithelial buccal cell exfoliation technique. Cytokeratin is very useful for tumor markers in oncology. Expression of cytokeratin in this study showed that the pattern did not the normal pattern. This pattern can be observed in Table 1, which showed increasing the expression of cytokeratin 5 in the epithelial buccal mucosal cells in the group exposed to azo batik workers. According to Garant [5] that cytokeratin 5 was not expressed on the superficial epithelial cells of the buccal mucosa. Buccal mucosa as lining mucosa was not keratinized. Expression of cytokeratin 5 and 14 observed in all types of basal cell keratinization and non-keratinization and the expression would be decreased in suprabasal layer. This study also showed the visible expression of cytokeratin 5 in the control group in small number (Table 1). The condition was probably due to post-transcriptional regulation of mitosis process. The number of expression of cytokeratin 5 were elevated in the batik workers group suspected pathological changes in circumstances on the buccal mucosa.

Increasing the number of epithelial cells expressed cytokeratin 5 in the buccal mucosa batik worker was significantly higher than the control may due to exposure to the azo material that always inhaled while working. Subjects had worked in the Unit of Coloring batik for at least five years so that the possibility of exposure time appropriate materials azo long time since they worked. Azo dyes entered to the the body through inhalation, then the dye was metabolized, resulting in abnormalities in the cells of epithelial the buccal mucosa. The use of azo dyes in batik that exceed normal limits suspected as the cause of an increase in the expression of cytokeratin 5 in batik workers. The use of azo dyes with a safe threshold value according to Decree No. LH 51 / MENLH / 10/1995 was in the range 200-400 mg / L, with the prohibition of the use permit by Permenkes No. 722 / Menkes / Per / IX / 2008 due to carcinogenic element. Results of the interviews indicated the use of azo dyes was about 10-15 g / L, or about 20 times higher than the maximum safe limit recommended by the Ministry of Environment.

The abnormality in the cells exposed to azo in batik worker also showed for vacuola in the cell cytoplasm (Figure 1 and
Positive expression of cytokeratin 5 showed brown colour in both the nucleus and cytoplasm of the buccal mucosa cell (Figure 2A) in exposed to azo batik workers group. In the control group only observed positive expression in the cell cytoplasm (Figure 2B), while in the cell nucleus was blue colour. The condition may cause the damage to the DNA in the basal cells that occurs when there was a change of epithelial cell differentiation patterns of differentiation. Cytokeratin expression was associated with epithelial cell differentiation stage buccal mucosa [17]. This study also supported previous results that exposed to azo dyes in batik workers in Yogyakarta was significantly effect on the increase in the frequency of micronuclei, karyolysis, and pyknosis the buccal mucosal epithelial cells. Increased frequency of micronuclei, pyknosis cell nucleus and karyolysis indicated damage of DNA nucleus [12, 13, 14].

Another possibility of changes in the pattern of differentiation in epithelial buccal mucosal cells in batik workers was a system of self protection to workers who were still very minimal. This could be seen when all workers in Unit of Colouring batik did not use personal protective equipment so that they could be vulnerable to get the risk exposure of azo dyes. Therefore, we concluded that exposure to azo colour can enhance the expression of cytokeratin 5 in the epithelial cells of the buccal mucosa batik workers in Yogyakarta.

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Compliance with Ethical Standards

Conflict of Interest: Author Juni Handajani declares that she has no conflict of interest. Author Dhinintya Hyta Narissi declares that she has no conflict of interest.

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Ethical approval: This article contains studies with human participants performed by the authors. The procedure in this study was approved from the Ethics Committee Faculty of Dentistry, Gadjah Mada University (Number: 00261/KKEP/FKG-UGM/EC/2015) on May 15, 2015.

Informed consent: All participants have agreed to participate in this study by signing the formal consent.

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**Author Profile**

**Juni Handajani** was graduated as a dentist, Master of Dental Science, and Doctoral degree from Faculty of Dentistry, Gadjah Mada University, Indonesia. In March 23, 2011, she received PhD degree in Dental Science from Niigata University Japan. She is as a lecturer and researcher at Department of Oral Biology, Faculty of Dentistry Gadjah Mada University since 1998 until now.