Tobacco smoking alters the number of oral epithelial cells with apoptotic features

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Abstract: Tobacco smoking is a global problem associated with the occurrence of many systemic diseases and tumors. Oral cavity tumors are common tobacco-related cancers, and of all the anatomical structures that are exposed to the effects of smoking, the oral cavity remains the least-explored area. Changes that occur in the biology of oral epithelial keratinocytes under the influence of the components of tobacco smoke often go unnoticed, if they are asymptomatic. The proper functioning of the oral epithelium is determined by the proliferation and differentiation of the cells in keratinization — the process of programmed cell death, which extends through to the mechanisms of apoptosis. Due to incomplete knowledge of the impact of tobacco smoke on the biology of keratinocytes, an evaluation of the cell cycle was conducted and the apoptosis of oral epithelial keratinocytes was analyzed. The study involved 77 patients divided into four groups according to their intensity of smoking, ranging from 0 to 27 pack-years. There were no differences in the cell count between nonsmokers and smokers in the proper cell-cycle phases. The percentage of proliferating cells in the oral epithelium is about 11%. A reduction in the number of early-apoptotic cells (caspase positive/propidium iodide negative) and an increase in the number of late-apoptotic cells (caspase positive/annexin V positive/propidium iodide positive) were observed to occur with increasing pack-years. The present study demonstrates that smoking does not affect the oral keratinocyte cell cycle, but does modify the number of cells with early and late apoptotic features. An intensification of the impact of tobacco smoke components on the biology of the oral keratinocytes is clearly noticeable at approximately 6 pack-years. This indicates that the biology of the first organ exposed to tobacco smoke — the oral epithelium — is altered by tobacco smoking. (Folia Histochemica et Cytobiologica 2014, Vol. 52, No. 1, 60–68)

Key words: tobacco smoking; human oral keratinocytes; apoptosis; cell cycle; flow cytometry

Introduction

Tobacco smoking is a global problem of civilization, with the number of tobacco smokers estimated at about 1.3 billion [1]. Epidemiological studies show that one third of tobacco smokers die of cardiovascular diseases [2]. Statistically, the life expectancy of smokers is 15 years less than that of nonsmokers [3, 4]. Tobacco smoking not only increases the risk of systemic disease, but it also contributes to the risk of cancerogenesis [5, 6]. Long-term exposure to carcinogens in the air is a high-risk factor for the development of lung cancer [1, 3, 4] and oral cavity cancers [7]. Tobacco smoke contains over 4000 substances that affect the human body, of which a number (about 60) are known carcinogens [8, 9]. Tobacco smoke is a physically heterogeneous aerosol formed by the incomplete combustion of tobacco. During combustion, the particles are pyrolyzed (decomposed under the influence of temperature) and connect with each
other in the process of combustion synthesis. While smoking a cigarette, the smoker absorbs about 500 mg of smoke, of which about 65% is nitrogen and oxygen; the remaining 35% consists of biologically active substances [10].

Tobacco smoke reduces the effectiveness of the immunological system, and thus increases the risk of respiratory infections, which may also contribute to the development of malignancies [5, 11].

The effects of nicotine on epithelial cells of the respiratory tract [12–14] are well documented, but the oral cavity has not been so well explored. The results of a few studies comparing the morphology of the oral epithelium of smokers with that of nonsmokers pointed to an increase in its thickness and also to higher levels of cytokeratins in smokers’ cells [15, 16]. It is well known that epithelia exposed to adverse chemical or physical effects defend themselves by increasing cytotkeratin levels [17].

The exposure of cell lines cultured in vitro in the presence of nicotine extract (the dominant substance in tobacco smoke) leads to irreversible changes that may impair fundamental processes of the cells, such as proliferation and differentiation [18–21]. Yet it is difficult to conclude the effects of tobacco smoke in vivo, where the cells are in contact with tobacco tar at a different frequency and for a long time — measured in years, and not in hours as in culture conditions. It is also extremely difficult to determine the amount of nicotine to match the levels present during smoking [22]. The assumption that cells treated in vitro with a single dose of nicotine extract will react in the same way as the cells of smokers — which are exposed to tobacco smoke many times each day, often for years — is questionable.

The outer layer of the oral mucosa is a stratified squamous epithelium that is keratinized in areas subjected to masticatory forces (e.g., the gingiva and palate). The different layers of the oral epithelium represent a progressive maturation process. Cells from the superficial keratinized layer are constantly being shed and replaced from below [23]. The homeostasis of the oral epithelium is the result of a balance between the dying and proliferating cells forming the squamous multilayered epithelium [24]. The process of keratinization is thought to be a physiological cell death, extending partly to an increase in its thickness and also to higher levels of cytokeratins in smokers’ cells [15, 16]. It is well known that epithelium exposed to adverse chemical or physical effects defend themselves by increasing cytotkeratin levels [17].

The sequence of molecular events occurring during apoptosis depends on a number of factors, inducing apoptosis (inducer) and the cell’s sensitivity to this type of death. There are two classic means of inducing apoptosis: through the activation of the receptors localized in the plasma membrane (the death domain receptors, such as TNFR, Fas, TRAILR) or through changes in the mitochondrial membrane [33, 34]. The extrinsic pathway activation of the receptors leads to procaspase 8 activation, which activates the caspase 3 involved in the cell’s self-destruction phase [35, 36]. In the intrinsic (mitochondrial) pathway, apoptosis is induced by an internal factor inside a cell, such as DNA damage, increased levels of reactive oxygen species, oxidative stress, or abnormal electrolyte transport, as a result of which factors activating procaspase 9 which activates caspase 3 are released from the intermembrane space [34, 35]. Caspase 3 induces mechanisms leading to proteolysis of important proteins (e.g., poly (ADP-ribose) polymerase and cytoskeletal proteins) and the activation of endonucleases that cut DNA into oligonucleosomal fragments [33, 37]. Caspases are a family of cysteine proteases involved in both the initial and executive phases of apoptosis [33, 37].

The participation of caspases in the process of squamous keratinocyte differentiation is controversial [25, 26, 38–40]. Studies by Lippens et al. have shown that the activation of proapoptotic caspases is not required for the induction of epithelial keratinocyte differentiation [25]; on the other hand, Wu et al. report the participation of these proteases in the maturation of keratinocytes [26].

Proliferation, cornification, and apoptosis are fundamental processes that condition the proper functioning of the oral cavity epithelium. Due to the conflicting information about the effects of smoking on oral epithelial cells, studies of keratinocytes obtained directly from smokers could provide important data on the effects of smoke on the biology of the oral epithelium.
no markers have been set for any biological alterations produced in oral keratinocytes by tobacco smoke prior to the appearance of clinical changes. The aim of this investigation was thus to compare the cell cycle and basic parameters of apoptosis (such as caspase activation and the externalization of phosphatidylserine in the plasma membrane) of oral epithelial cells obtained from nonsmokers with those from smokers.

Material and methods

Patients. Fragments of oral mucosa were harvested during routine dental treatment (extraction of lower molars). All patients completed a detailed survey before entering the project. The form included questions on the patient’s current health status, medications, smoking (the number of cigarettes smoked per day and the number of years of addiction). Patients with systemic diseases, treated chronically with drugs, and who had consumed alcohol in the past 3 days were excluded from the project. All patients were examined clinically prior to obtaining the material. Any pathology of the oral mucosa, as well as symptoms of inflammation, led patients to be excluded from the project. All patients were given sufficient information regarding the study, and all individuals signed an informed consent form. The procedures were accepted by the Ethics Committee of the Medical University of Gdansk (NKEBN/72/2011).

In the group of 77 patients, there were 40 nonsmokers (Group 1, aged 27 ± 6.4 years, mean ± SD), 33 smokers (mean age 26 ± 4.7 years), and 4 heavy smokers with the highest average number of pack-years (mean age 51 ± 9.5 years). Smokers were divided into groups according to smoking intensity, expressed as pack-years. The pack-year (py) is the international unit for measuring the number of cigarettes a person has smoked over a long period, and is calculated as follows: Number of pack-years = (number of cigarettes smoked per day × number of years smoked)/20 (1 pack contains 20 cigarettes) [40]. The average number of pack-years for the group of 33 smokers was 3.0 ± 2.6. The smokers’ group was therefore divided into Group 2, with pack-years equal to or less than 3.0 (20 patients with average 1.4 py) and Group 3, with more than 3.0 pack-years (13 patients with average 5.6 py). Groups 2 and 3 were of similar ages (mean 25 ± 5.1 and 27 ± 4.2 years, respectively), as was Group 1 of nonsmokers. A Group 4 was also distinguished, with the highest average number of pack-years of 26.9. Due to their higher average age (4 patients of age 51 ± 9.5 years), these smokers were not included in the basic analysis with Groups 2 and 3. However, in the further evaluation of the results, Group 4 was used to highlight the influence of the smoking intensity on the oral epithelium.

Isolation of cells from the oral epithelium. Fragments of the gingival oral mucosa, about 0.5 cm² in size, were incubated in dispase (5 mg/mL, Gibco, Paisley, UK) for about 18 hours at 4°C to separate the epithelial sheets from the connective tissue. Single cells were obtained as the result of incubation of the oral epithelium in 0.25% trypsin-EDTA solution for 30 min at 37°C [41]. The viability of the obtained cells was over 80%, as determined by trypan blue staining, and 97% of cells were keratinocytes, as determined by staining cells with a mixture of antibodies against cytokeratins (anti human cytokeratin clone AE1/AE3, Dako, Glostrup, Denmark). The obtained keratinocytes originated from all layers of the oral epithelium, as was documented by microscopic analysis of the cells’ morphological heterogeneity (Figure 1).
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Cell cycle analysis. 1 × 10^5 ethanol-fixed keratinocytes were resuspended in 1 ml of staining solution (RNase A 200 μg/mL and propidium iodide (PI) 5 μg/mL in PBS). Then cells were incubated for 30 min at 37°C in the dark. 20,000 cells were analyzed on a FACS Calibur flow cytometer (Becton Dickinson Immunocytometry System, San Jose, CA, USA). The results were analyzed off-line using Cylogic software (CyFlo Ltd).

Cytofluorimetric estimation of number of cells with activated caspases. We applied the FLICA test (fluorochrome-labeled inhibitors of caspases) to estimate the number of cells with activated caspases [42]. The idea behind this method is based on the fact that fluorochrome-labeled inhibitors of caspases react covalently with the reactive enzymatic centers of the activated caspases. We used FITC-(fluorescein)-labeled pan-caspase inhibitor VAD-FMK (VAD, three amino acids peptide; FMK, fluoro-methyl ketone, which binds to cysteine of caspase), which detects all activated caspases in the cell, as the three-amino-acid target sequence binds to most caspases. In this method, simultaneous staining of cells with FITC-VAD-FMK and PI allows to distinguish the sequential stages of apoptosis which were described as: (i) early apoptosis: here the cells have a high intensity of green fluorescence, which indicates a high level of activated caspases (C+), but exclude PI, as the plasma membrane is not damaged (PI–); they were denoted C+/PI– (caspase positive/PI negative); (ii) late apoptosis: here the cells bind FITC-VAD-FMK to the activated caspases (C+) and are stained by PI, because the plasma membrane is damaged (PI+); denoted as C+/PI+ (caspase positive/PI positive); (iii) final stage of apoptosis: here the cells are stained by PI (PI+), but do not have active caspases (C–); denoted C–/PI+ (caspase negative/PI positive).

From each patient, 0.5 × 10^6 cells were incubated with 5 μM FITC-VAD-FMK (CaspACE™ FITC-VAD-FMK, Promega, Madison, USA) for 30 min at room temperature in the dark, before being washed and resuspended in PBS (phosphate-buffered saline) containing PI. 20,000 cells were analyzed on a flow cytometer.

Changes in the plasma membrane structure: externalization of phosphatidylserine. Externalization of phosphatidylserine (PS) is commonly used as a marker of apoptotic change in plasma membranes, which is a signal for macrophages to phagocytise the cell. The annexin V-binding test was used to estimate PS exposure on the plasma membrane surface (Annexin V-Fluos staining kit, Roche, Mannheim, Germany). Simultaneous staining with annexin V-FITC (detecting both apoptotic and necrotic cells with phosphatidylserine residues exposed on their surface) and PI (staining DNA in the cells with damaged plasma membrane, e.g., dead cells) allows discrimination between early and late apoptotic cells. Thus, the Annexin V test characterizes the sequential stages of apoptosis like the FLICA test: (i) early apoptotic cells bind annexin V but do not stain with PI (annexin positive/PI negative, An+/PI–); (ii) late apoptotic/necrotic cells bind annexin V and stain with PI (annexin positive/PI positive, An+/PI+). Cells were stained with Annexin V-FITC and PI according to manufacturer’s instructions and analyzed on a flow cytometer.

Microscopic analyses. Initial microscopic analysis of the oral epithelial cells in a bright field with Hoffman’s contrast was performed on a TS-100F light microscope (Nikon, Tokyo, Japan). The colocalization study of stained epithelial cells (FLICA, annexin V and PI) was performed using a Radiance 2100 confocal laser-scanning microscopy (CMLS) system (Bio-Rad, Hemel Hempstead, UK) mounted on Nikon Eclipse 600 microscope. The CMLS images were obtained with 40× and 60× oil-immersion objective lenses with N.A. of 1.3 and 1.4, respectively. The optimal iris diameter was used for each magnification. The CMLS images were analyzed using LaserSharp 2000 and LaserPix v. 2.0 software (Bio-Rad).

Statistical analysis. The group data are expressed as arithmetic means ± SD. Statistical analysis was performed using the ANOVA Kruskal-Wallis test and the Dunn test. Differences at p < 0.05 were considered significant. The tests were performed using Statistica data analysis software system, version 10 (StatSoft, Inc., 2011, www.statsoft.com).

Results

Analysis of cell cycle of oral epithelial cells

The analysis of the cell cycle (Table 1) in the oral epithelium of nonsmokers (Group 1) showed that the percentage of cells in the individual phases was 82.1% for G0/G1, 7.1% for S, and 4.3% for G2/M. For the Group 3 (smokers with 5.6 py), these values were 78.7%, 8.5%, and 3.4%, respectively. Thus, smoking about 6 pack-years did not significantly change the number of cells in the various cell cycle phases. There was, however, a tendency for the percentage of cells in the S phase to increase with pack-years of smoking (Table 1).

Morphologic assessment of apoptosis of oral keratinocytes with confocal microscopy

The FLICA-stained oral keratinocytes indicated that caspases were localized in the cytoplasm and that their activity was heterogeneous (Figure 2A). Caspase-positive cells occurred among cells of all sizes, and the largest cells from the outer epithelial layers also contained activated caspases (Figure 2B).
Annexin V-FITC staining showed that externalization of PS also affected cells of all sizes, (Figure 2C) among which are flattened cells with degenerating nuclei from the outer epithelial layers (Figure 2D).

Table 1. Cell cycle analysis of the oral epithelial cells from subjects with different smoking intensity (expressed as pack-years, py)

<table>
<thead>
<tr>
<th>Percentage of cells in the cell cycle phase</th>
<th>Group 1 (Control, 0 py)</th>
<th>Group 2 (1.4 ± 1 py)</th>
<th>Group 3 (5.6 ± 2.1 py)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G0/G1</td>
<td>82.1 ± 5.0</td>
<td>83.0 ± 4.0</td>
<td>78.7 ± 7.0</td>
</tr>
<tr>
<td>S</td>
<td>7.1 ± 2.2</td>
<td>7.6 ± 2.1</td>
<td>8.5 ± 2.9</td>
</tr>
<tr>
<td>G2/M</td>
<td>4.3 ± 1.7</td>
<td>3.8 ± 1.0</td>
<td>3.4 ± 1.5</td>
</tr>
</tbody>
</table>

Values are means ± SD

Figure 2. Activated caspases and phosphatidylserine externalization in oral epithelial cells isolated from one nonsmoker studied by confocal laser-scanning microscopy. FLICA test (A, B) and Annexin V test (C, D) staining are shown. A. Activated caspases present in cells of all sizes; B. The arrow indicates cytoplasm with high FITC-VAD-FMK fluorescence; C. Phosphatidylserine externalization present in cells of all sizes; D. The arrow shows Annexin V-FITC fluorescence in the plasma membrane. Scale bar 10 μm

Analysis of cells with apoptotic death features

Activated caspases

The overall number of cells with activated caspases was similar in all examined groups, but the detailed
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Analysis of the apoptotic stages showed significant differences between nonsmokers and smokers (Table 2). Analysis of the oral keratinocytes of nonsmokers revealed that about 30% of all cells had activated caspases; in this group early apoptotic cells with intact plasma membrane (C+/PI–) dominated (Table 2). Smoking habit decreased the population of early apoptotic cells to 14.5% for Group 3 (5.6 py) with a reduction of 28% in comparison to nonsmokers (Table 2). Interestingly, the number of late apoptotic cells with damaged plasma membranes (C+/PI+) in the Group 4 (26.9 py) was significantly higher than in nonsmokers (Table 2, Figure 3). About 1% of oral keratinocytes of nonsmokers stained with PI, but lacked activated caspases (C–/PI+). This population increased for Group 3 to 2% (Table 2).

**Plasma membrane changes: phosphatidylserine externalization**

In the group of nonsmokers, about 29% of cells showed PS externalization, while those smoking 26.9 pack years had a significant increase, possessing over 40% of such cells. This change was mainly related to late apoptotic cells with damaged plasma membranes (An+/PI+, Table 2, Figure 3).

Summarizing the observed changes in caspase activation and plasma membrane, it was noted that, with increased pack-years of smoking, the number of cells with late apoptotic features (high activity of caspases, PS externalization, damaged cell membrane) increases, as illustrated in Figure 4.

**Effect of smoking on the content of cells with reduced DNA content**

The percentage of dying cells with reduced DNA levels (e.g., apoptotic bodies) located in the SubG0 part of the cell cycle increased with pack-years of smoking (Table 2). Among nonsmokers, these constituted 3% of the cells, while for the Group 3 smokers, the population of such cells almost doubled. In people smoking 26.9 pack-years the level of cells with reduced DNA amount was about 7% (Table 2).

**Discussion**

In the present study, we have demonstrated that tobacco smoking induced changes in the apoptosis of human oral keratinocytes, which may be signs of alterations in oral epithelium biology.

The limited information available in the literature concerning the proliferation potential of oral keratinocytes indicates that nicotine can increase or decrease the number of proliferating cells in vitro [19, 31]. Our results, based on cells obtained directly from smokers, showed no significant difference with cells from nonsmokers. Among nonsmokers, only 11% of the cells were in the S/G2/M phases — much lower than predicted by other authors who had examined oral keratinocytes [19, 28, 29] and epidermal keratinocytes [19, 43]. This higher percentage of cells in the S/G2/M phases is probably due to the fact that most of these observations were carried out on cultured cell lines, and not on cells obtained directly from patients.

### Table 2. Percentage of the oral epithelial cells with apoptotic features in subjects with different smoking intensity (expressed as pack-years, py)

<table>
<thead>
<tr>
<th>Apoptotic features</th>
<th>Group 1 (Control, 0 py)</th>
<th>Group 2 (1.4 ± 1 py)</th>
<th>Group 3 (5.6 ± 2.1 py)</th>
<th>Group 4 (26.9 ± 15.7 py)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Activated caspases</td>
<td></td>
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<tr>
<td>Early apoptotic C+/PI–</td>
<td>20.1 ± 6.2</td>
<td>16.6 ± 7.5</td>
<td>14.5 ± 5.2d</td>
<td>13.2 ± 4.6</td>
</tr>
<tr>
<td>Late apoptotic C+/PI+</td>
<td>9.2 ± 4.6</td>
<td>9.6 ± 4.0</td>
<td>10.6 ± 4.2</td>
<td>16.9 ± 5.0a</td>
</tr>
<tr>
<td>Cells at the final apoptosis C–/PI+</td>
<td>0.7 ± 0.8</td>
<td>1.7 ± 2.4</td>
<td>2.2 ± 2.3b</td>
<td>1.0 ± 0.4</td>
</tr>
<tr>
<td>2. Phosphatidylserine externalization:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early apoptotic An+/PI–</td>
<td>9.5 ± 5.2</td>
<td>10.0 ± 5.6</td>
<td>9.3 ± 3.1</td>
<td>9.5 ± 2.5</td>
</tr>
<tr>
<td>Late apoptotic An+/PI+</td>
<td>19.3 ± 7.3</td>
<td>21.8 ± 8.1</td>
<td>23.6 ± 7.9</td>
<td>32.5 ± 7.7a</td>
</tr>
<tr>
<td>3. Hypodiploid DNA content</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>(cells in the sub G0 region)</td>
<td>3.0 ± 2.5</td>
<td>2.4 ± 1.2</td>
<td>5.6 ± 4.9</td>
<td>7.5 ± 7.1</td>
</tr>
</tbody>
</table>

Classification of cells into the early, late and final apoptotic stages was based on criteria described in Materials and Methods. Values are means ± SD. *and ** significantly different from the corresponding values of the control group (nonsmokers), p < 0.05 and p < 0.01, respectively (Kruskal-Wallis test)
Our finding that there was no significant difference between nonsmokers and smokers with 6 pack-years of smoking in the percentages of cells in individual cell cycle phases are difficult to discuss, because there are no similar flow cytometric analyses in the available literature based on material obtained directly from smokers. It is worth noticing that the smokers examined in our study showed no macroscopic changes in the oral cavity that might be the result of disturbances in epithelial cell proliferation.

Keratinization is thought to be a process of physiological cell death which occurs as the result of the process of cell differentiation [26, 27]. The majority of publications which evaluated the influence of to-

Figure 3. Flow cytometric analysis of oral epithelial cells with active caspases (FLICA test) and externalized phosphatidylserine (Annexin V test) from examined groups of subjects with different smoking intensity expressed as pack-years (py). Density plots of a representative subject from each group. The plot is divided into identical quadrants containing: lower right: C+/PI−, An+/PI−, early apoptotic cells; upper right C+/PI+, An+/PI+, late apoptotic cells; upper left C−/PI+, cells in the final stage of apoptosis (C, caspases; An, annexin; PI, propidium iodide)

Figure 4. Percentage of late apoptotic cells with activated caspases (C+), phosphatidylserine externalization (An+), and discontinuous plasma membrane (PI+) in examined groups with different smoking intensity expressed as pack-years. *significantly different from control, p < 0.05 (Dunn test)
bacco smoke on the intracellular processes of oral keratinocytes, examined only the general features of apoptosis — such as caspase activation, and changes in mitochondria, plasma membrane, and chromatin. The participation of caspases in the process of differentiation of squamous keratinocytes is controversial [25, 26, 38, 39]. The studies of Lippens et al. have shown that the activation of proapoptotic caspases is not required for the induction of differentiation of epithelial keratinocytes [25], while Wu et al. have reported on the participation of these proteases in the maturation of keratinocytes [26]. Our results clearly document that this group of enzymes is presented in the oral keratinocytes of all epithelial layer cells, but that the amount of activated caspase differs. These observations need further studies to estimate the types of caspase that appear in oral keratinocytes, as there is uncertain need further studies to estimate the types of caspase amounts of activated caspase differs. These observations that this group of enzymes is presented in the oral mucosa. However, ultrastructural studies of the oral mucosa cytoarchitecture of heavy smokers revealed reductions in desmoglein 3 and keratin 10 expressions [30].

In summary, the present study demonstrates that smoking at the rate of 6 pack-years did not affect cell cycle of human oral keratinocytes, but altered the number of cells with apoptotic features with increasing pack-years of smoking. Our study is the first observation in humans, however, the investigations on a group of smokers with longer-lasting habit could provide more data about the interference of smoking with the biology of oral keratinocytes.

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