Ethanol influence on gingival fibroblasts – a real-time in vitro study

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INTRODUCTION

Alcohol consumption is the world’s third largest risk factor for disease and disability, revealing the greatest risk in middle-income countries [1]. According to a WHO report (2011), over 70% of the urban and rural populations admit to alcohol consumption [2]. Lower socio-economic status and educational levels result in a greater risk of alcohol-related injury, disease and death. Alcohol is a common component of many medicines, as well as an ingredient in many oral hygiene home products. Mouthwashes containing alcohol are considered to inhibit wound healing in the oral cavity. Due to the fact that many different results are described for different concentrations of alcohol at different times, an attempt was made to visualise the direct impact of 7.2% and 22% alcohol on human gingival fibroblasts.

Materials and method. PANsystem 2000 was used for visualisation of the reaction of human gingival fibroblasts isolated from gingiva on ethanol in 2 different concentrations. PANsys 3000 is a multi-system fully-automated cell culture device used for in vitro culture and to study a variety of cell lines under conditions similar to in vivo. Observations were carried out for 48 hours since alcohol addition. Pictures were taken in a continuous process at 5 minute intervals and combined into a film.

Results. Both contamination of 7.2% and 22% ethyl alcohol negatively affected morphology and cell proliferation. Addition of ethanol at a concentration 7.2% enabled cells to regain their ability to divide and recover normal morphology after 10 hours; changes caused by 22% ethanol, however, were irreversible.

Conclusions. The obtained results suggest that daily usage of 7.2% alcohol contained in mouthwashes is non-toxic for gingival fibroblasts, and could be recommended after periodontal surgery.

Key words

alcohol, oral cavity, gingival fibroblasts

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to 10% alcohol for 48 h. Moreover, the increase in the number of cells was lower than in the control group, although the difference was not statistically significant. Isolated necrotic cells were observed [15].

The most effective mouthwashes contain chlorhexidine (CHX) or essential oils (EO). 0.1% CHX contains 7.2% ethanol, whereas EO contains 22% ethyl alcohol. The presence of ethanol in a mouth rinse containing 0.10% chlorhexidine has no deleterious effects on healing capacity; on the contrary, it helps stimulate wound healing. The combination of chlorhexidine plus ethanol is superior for healing. It is suggested that ethyl alcohol added to chlorhexidine decrease chlorhexidine cytotoxicity [16]. However in many cell-based experimental models, ethanol has been found to inhibit the effects of growth factors, including insulin and insulin-like growth factor [17, 18]. The growth-inhibitory effects of ethanol are often accompanied by increased apoptotic cell death [19, 20].

Due to the fact that many different results have been described for different concentrations of alcohol at different times, an attempt was made to visualise the direct impact of 7.2% and 22% ethanol on human gingival fibroblasts. For this purpose, a fully automated multi-system cell culture devised. This allowed observation of the impact of ethanol on human fibroblasts in conditions most similar to in vivo, and allow a longer time without intervals.

MATERIALS AND METHOD

PANsys 3000 (Systech GmbH, Augsburg, Germany) was used for the experiment. This is a multi-system fully automated cell culture device used for in vitro culture and for studying a variety of cell lines under conditions similar to in vivo. The system allows for the culturing of various cells and the use of various compositions of the media at the same time, using any culture conditions and a selected microscopic observation. All data associated with the experiments can be recorded and played back at a later time.

In order to perform the experiment, human fibroblast cells were isolated from gingival tissue obtained from 5 patients during the standard protocol of recession, and cultured in flasks with surface of 25 cm² in DMEM (Dulbecco’s Modified Eagle’s Medium) supplemented with 10% FBS (Foetal Bovine Serum) and 1% antibiotic-antimycotic solution (10,000 units penicillin, 10 mg streptomycin and 25 μg amphotericin B per ml) at 37 °C with 5% CO₂ concentration. The cell cultures with 70–80% confluence were washed with 3 ml of Hanks solution and trypsinized (0.25% trypsin, 0.02% EDTA). Cells were gathered by centrifugation (1,000 rpm, 10 min). 500 ml of the cell suspension after trypsinization was combined with 500 ml of culture medium, supplemented with 10% FBS and 1% antibiotic-antimycotic solution, and placed in a growth chamber device PANsys 3000 – a multi-system fully-automated cell culture device used for in vitro culture and for studying a variety of cell lines under conditions similar to in vivo. The cells were cultured for 24 h at 37 °C with 5% CO₂ concentration. After 24 h of culturing, 250 ml of ethyl alcohol (7.2% and 22%) was added to each chamber:

- Chamber 1 – 7.2% ethanol in DMEM solution;
- Chamber 2 – 22% ethanol in DMEM solution;
- Chamber 3 – control sample: full DMEM, no addition of ethanol.

All reagents were obtained from MERCK, Life Science Department, Poland.

Observations were carried out for 48 h after alcohol addition. Images were performed in a continuous process at 3 min intervals and combined into a movie. Images related to a specific region in the chamber – the same region (Region of Interest) and photographed for 72 h from the moment of placing cell solutions in the chambers – 24 h, and then after adding ethyl alcohol to the culture – 48 h.

All experiments were performed 3 times for each patient’s cells. All results were similar. Results from a representative experiment are shown below.

RESULTS

As a result of carrying out 48 h of cell culture in DMEM solution containing alcohol at a concentration of 7.2 and 22%, changes were observed in morphology and cell division in both cell lines.

Immediately after the addition of alcohol at the concentration of 7.2%, changes in cell proliferation and cell morphology were observed, in comparison to the control (Fig. 1) (http://marzena.softweb.pl/control.avi; password wyga1302). Decrease in growth rate and the number of cell divisions was observed, resulting in a constant number of cells in the observed region. The cells were also morphologically changed: size was significantly reduced and shape was changed (from spindle to irregular) (Fig. 2). 10 hours after alcohol addition, the cells returned to their normal, regular shape and started again the regular dividing processes (http://marzena.softweb.pl/ethanol7_2.avi; password wyga1302).

Immediately after the addition of alcohol at the concentration of 22%, significant changes in cell proliferation and cell morphology were observed, in comparison to the control. The cells were also morphologically changed: size significantly reduced and shape changed (from spindle to irregular). After addition of the alcohol to the culture...
medium, the cells were incubated in a hypertonic solution, cells lost water so that cytoplasm became very thick and cell volume was significantly decreased. There was also complete inhibition of cell division. After 6 hours from the addition of alcohol, cells completely lost their ability to divide and took a spherical shape. There were no visible movements of the cytoplasm. A very visible effect of the addition of alcohol was also the immediate loss of cell projections (Fig. 3; 23 h and 2 minutes), which in turn made it impossible to connect to the fibroblast colony and thereby further divisions (http://marzena.softweb.pl/ethanol22.avi; password wyga1302).

Ethanol causes the off-state of normal operations and weakening of the cell. After the resignation of poisoning, the cell regains its previous efficiency (which is noticeable in the case of 7.2% ethyl alcohol addition), but if poisoning lasts longer, the function of the cell is permanently weakened or irreversibly disappears.

DISCUSSION

The observation let us to conclude that both the contamination with 7.2% and 22% ethyl alcohol negatively affects morphology and cell proliferation. The addition of ethanol at a concentration 7.2% enables cells to regain their ability to divide and recover normal morphology after 10 h, when changes caused by 22% ethanol are irreversible.

Alcohol consumption may be considered as a risk indicator for periodontitis [21]. There are several potentially important local mechanisms by which alcohol can interfere with fibroblasts. One of local mechanism by which alcohol can interfere with fibroblasts in the wound-healing process in maxillofacial injuries is the inhibition of fibroblast proliferation and ECM synthesis at the wound site [22]. Alcohol significantly reduced cell viability and increased reactive oxygen species in oral fibroblasts [6]. In animal models, alcohol consumption increased gingival oxidative damage and the production of TNF-α in periodontal ligament fibroblasts [23]. Decreasing cell viability in response to increasing alcohol concentration was dose-dependent. Alcohol can decrease host defence in oral cavity tissue by altering cytokine production and lymphocyte T function [11]. In another in vitro study, alcohol negatively influenced fibroblast growth factor-mediated aortic smooth muscle cell proliferation by reducing phosphorylation of downstream kinases, and disrupting the cell cycle regulation [24]. Fibroblast cells isolated from trauma patients caused impairments in re-epithelialisation, angiogenesis, and inflammation in wounds following acute alcohol exposure [25]. It has been suggested that the cytotoxicity of alcohol depends on concentration and how long the mouthwash was retained in the mouth [26].

Alcohol on its own causes damage to the oral mucosa and includes epithelial atrophy and decrease in basal cell size atrophy with associated hyper-regeneration [27]. However, a cytological study performed by Bagan et al. [28] revealed no changes in epithelial superficial, intermediate, parabasal, or basal cells in samples collected from patients who had regularly been using mouthwash containing 26% alcohol for 6 months.

In the presented study, a very similar method for in vivo conditions was used for cell behaviour observation. The reaction of fibroblasts after 7.2% alcohol stimulation was probably connected with the difference of concentration between the external environment and cell cytoplasm. The hydrophilic nature of alcohol renders its easy distribution to every tissue containing water. It is also possible that the difference obtained in the current study is due to only a single dose of alcohol. Ethanol has easily detectable promitogenic effects only when added to the cells only once, without sealing the wells [29]. It has been shown that ethanol can bind, although weakly, to certain proteins in various membranes which become saturable for it [30,31]. This could probably explain the non-toxic or even stimulatory effect...
of ethanol when combined with other active ingredients in medicines [32].

The present study shows that the applied system is convenient for monitoring the reaction of cells to different agents. However there are some limitations to the study. First of all, trypsinization influences the cells vitality. The visualization of fibroblasts reaction on ethanol, although performed in conditions most similar to in vivo, is still far from a real in vivo situation. In a real in vivo situation it is difficult to exclude the influence of ethanol metabolite products on cells during ethyl alcohol consumption.

CONCLUSION

The real-time in vitro study is a convenient model to observe and record cells in reaction to various agents, such as ethanol. This is the first time that the reversible effect of 7.2% alcohol on gingival fibroblasts viability has been visualized. These results suggest that the daily use of mouthwashes containing 7.2% alcohol is non-toxic for gingival fibroblasts, and can be recommended after periodontal surgery.

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REFERENCES