In vitro culturing of porcine tracheal mucosa as an ideal model for investigating the influence of drugs on human respiratory mucosa

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Abstract It has been previously shown that fresh mucosa from different mammals could serve as raw material for in vitro culturing with the differentiation of cilia, which are the most important morphological structures for the function of the mucociliary system. Increasing legal restrictions on the removal of human tissue and changing surgical techniques have led to a lack of fresh human mucosa for culturing. Most of the animals that have been used as donors up to now are genetically not very close to human beings and must all be sacrificed for such studies. We, therefore, established a modified system of culturing mucosa cells from the trachea of pigs, which is available as a regular by-product after slaughtering. With respect to the possibility of developing “beating” cilia, it could be shown that the speed of cell proliferation until adhesion to the coated culture dishes, the formation of conjunctions of cell clusters and the proliferation of cilia were comparable for porcine and human mucosa. Moreover, it could be demonstrated that the porcine cilia beat frequency of 7.57 ± 1.39 Hz was comparable to the human mucosa cells beat frequency of 7.3 ± 1.4 Hz and that this beat frequency was absolutely constant over the investigation time of 360 min. In order to prove whether the reaction to different drugs is comparable between the porcine and human cilia, we initially tested benzalkonium chloride, which is known to be toxic for human cells, followed by naphazoline, which we found in previous studies on human mucosa to be non-toxic. The results clearly showed that the functional and morphological reactions of the porcine ciliated cells to these substances were similar to the reaction we found in the in vitro cultured human mucosa.

Keywords In vitro cultured ciliated cells · Porcine · Human · Naphazoline · Benzalkonium chloride

Introduction The mucociliary system of the airways plays a decisive role in the defence mechanism for cleaning the air of bacteria, allergens, dust and other foreign particulate matter. It, therefore, has an important protective function for the whole organism. Moreover, it regulates the temperature and humidity of the inhaled air, which are important parameters for the function of the lung.

Because of the importance of these functions, it is vital to possess an effective and reliable evaluation method for observing under constant conditions the reaction of the mucociliary system to the influence of different noxious substances as well as protecting or healing agents.

Until recently, our study group has evaluated these influences using in vitro cultured human nasal mucosa. Increasing legal restrictions on the removal of human tissue and changing surgical techniques have led to a lack of this material. In the past, experiments on the in vitro cultured mucociliary system have been accomplished in several...
mammals, however most of these are not suitable either because of similar legal problems, as in monkeys, or because they are herbivores and therefore, not genetically very close to human beings. Furthermore, to gain fresh mucosa in order to culture mucociliary tissue, it is inevitable that all these experimental animals must be sacrificed.

We, therefore, developed an alternative model by using porcine mucosa from the trachea, which has the advantages, that the pig as an omnivore has a similar biology in comparison to the human being and that the tracheal mucosa is available as a normal by-product after slaughtering.

In order to prove the suitability of the in vitro culturing of this porcine mucosa, we investigated the capability to develop cilia, the minimal time range of their constant beating frequency and the reaction of the cilia to different drugs compared to the reaction of human mucosa.

Materials and methods

Origin and isolation of porcine tracheal epithelial cells

Respiratory mucosa for cell culturing was harvested from fresh porcine trachea. The porcine–trachea was first cut into halves and cleaned by mechanical brushing in an acetylcysteine solution. The tissue was placed for less than 5 h into DMEM (Sigma) supplemented with antibiotic/antimycotic solution (penicillin 200 U/ml, streptomycin 200 μg/ml) and amphotericin B (0.5 μg/ml). To minimize the risk of contamination, the tissue was immersed in 10% iodine polyvidone (Betaisodona®) in Phosphate Buffered Saline (PBS) for 3 min and was then rinsed three times in PBS. After washing, the specimens were immersed in 0.01% protease from Streptomyces griseus (Sigma) in serum free culture medium for 3–4 h at 34°C. By gentle agitation with a scalpel, epithelial cells were released into the medium as single cells, cell clusters and larger mucosal sheaths. After filtration through a stainless steel sieve, the cell suspension was centrifuged at 150g for 10 min and resuspended in a serum-free culture medium [17, 18, 20].

Origin and isolation of human epithelial cells

Respiratory mucosa for cell culturing was harvested from partial resection of hyperplastic inferior turbinates in rhinosurgery. The patients had preoperatively given their informed consent. For isolating the cells, afterwards we applied the same procedure as for the porcine cells.

Culture medium for porcine and human epithelial cells

Both culture media were a mixture of DMEM and Ham’s F12 (Sigma), supplemented with insulin (1 μg/ml), transferrin (1 μg/ml), hydrocortisone (0.5 μg/ml), retinoic acid (10 ng/ml), L-glutamine (3.2 mM), 1.25% antibiotic/antimycotic solution (penicillin–streptomycin). For the porcine–trachea cells, it was necessary to add a gentamicin and kanamicin solution at a concentration of 0.1%, because this tissue was frequently severely contaminated with bacteria.

Submersion cell culture for porcine and human epithelial cells

Tissue culture dishes (EASY GRIP™, Falcon®) were used for incubation. The bottom of the culture dishes was coated with extra cellular matrix (ECM) molecules of the basal lamina. The following stock solutions of ECM-molecules were used for coating: 50 μl collagen IV [collagen IV (0.5 mg/ml) in 0.25% acetic acid], 10 μl laminin and 1 μl heparan sulphate proteoglycan (HSPG) were added to 1 ml PBS in each culture dish and incubated at 34°C for 24 h. The ECM-coating solution was removed and the dishes were washed and filled with the culture medium. The resuspended cells were seeded in submersion in high density on the tissue culture dishes and kept in an incubator (humidified atmosphere, 10% CO₂, 34°C). The culture medium was changed every 3 days, [21].

Measurement of ciliary beat frequency

Ciliary beat frequency of cultured cells was measured in a custom-made test facility. An inverted research microscope (DM IRB, Leica) with a heated stage (HT 200, Minitüb) was used. A colour video camera (CCD-Iris/RGB, Sony) relayed the microscope image to a monitor. The culture medium was controlled for temperature (Temperature-Controller HT 1.2, Vetec) and continuously changed (Masterflex digital console drive, Cole-Parmer Instruments) with a rate of 1 ml/min. Thus, different concentrations of the pharmaceutics solution could be completely replaced by the culture medium within 7 min. Temperature and pH-value were constantly controlled in the cell cultures. Each experiment was recorded on videotape. The CBF of single cells was measured according to the following log (Table 1).

CBF was sampled optically (custom made) from the monitor and computerized by FFT-Analyzer (CF-350 Z, Ono Sokki).

Table 1 Log for measurement of CBF with different pharmaceutics concentrations

| Concentration | Equilibration | Concentration 1 | Concentration 2 | Rinse out |
|---------------|---------------|-----------------|-----------------|----------|
| At time (min) | 0, 20, 40     | 60, 80          | 100, 120        | 140, 160 |

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Results

Constancy

An adherent monolayer culture of human and porcine ciliated cells (Fig. 1) was present after 4 days in vitro. The ciliary beat frequency of untreated cultured human nasal cells was measured over 180 min and appeared to be constant at least over this time range ($n = 7$; 29 single cells, two-sided $t$-test, $P < 0.05$). The ciliary beat frequency of porcine tracheal cells, which was controlled up to 360 min, also remained constant over this time period and additionally showed no significant difference in comparison to the CBF during the equilibration time ($n = 12$; 53 single cells, paired $t$-test, $P < 0.05$) (Fig. 2).

Benzalkonium chloride (BAC)

Human nasal cells

Increasing concentrations of BAC initially caused an opposing reaction of the CBF of the cultured human nasal cells (Fig. 3). A concentration of 0.001% BAC significantly increased the CBF to 112.73% ($\pm 17.26$, $P = 0.042$). But...
Comparison between human versus porcine cells

In comparison to the human nasal cell culture, the porcine tracheal cells showed only a slightly significant difference at the concentration of 0.001% \( (P = 0.0367) \). The other concentration of 0.001% \( (P = 0.675) \) as well as the rinsing out \( (P = 1) \) stimulated equal reactions.

Naphazoline

**Human nasal cells**

Increasing concentrations of naphazoline caused no significant stimulation or decrease of the CBF of the cultured human nasal mucosa cells (Fig. 4). Concentrations of 0.01% \( (108.75 \pm 15.84\%, \ P = 0.237) \), 0.1% \( (105.99 \pm 16.46\%, \ P = 0.366) \) and during rinsing out \( (104.09 \pm 30.41\%, \ P = 0.636) \) did not change the CBF relative to the equilibration.

**Porcine tracheal cells**

The porcine-tracheal cells again showed almost the same effect of naphazoline on the ciliary beat frequency (Fig. 4). The concentration of 0.01% naphazoline caused no significant variation from the equilibration \( (103.74 \pm 10.26\%, \ P = 0.319) \). Only the concentration of 0.1% showed a significant decrease of CBF \( (92.46 \pm 21.22\%, \ P = 0.002) \). However, the rinsing out with substrate-free medium showed that the decrease was statistically also reversible because of lower standard deviation \( (80.77 \pm 25.04\%, \ P = 0.234) \).

**Comparison between human versus porcine cells**

Compared to the human nasal cell culture, the porcine tracheal cells showed no significant difference in all concentrations of 0.01% \( (P = 0.385) \). The other concentration of 0.1% \( (P = 0.051) \) as well as the rinsing out \( (P = 0.083) \) also caused equal reactions.

**Discussion**

There are growing limitations of the removal of human tissue. Following legislation, it is permitted only with written consent of the patients, which increases administrative costs. Further, changing surgical techniques may lead to a lack of human tissue. In order to judge the effect of medicines on ciliary beat frequencies, a sufficient number of primary cultures must always be available.

In the past 30 years (since 1978), experiments on in vitro cultured mucociliary cells have been conducted on the following mammals: monkey [23], cow [30–35], sheep [8, 15, 24–27], rabbit [4, 13, 29, 36, 37], ferret [16, 19], hamster [14] and rat [5, 7, 12, 22, 28].

Cows, oines, rabbits and hamsters are all herbivore and are therefore not very close to human beings genetically. Closest to the humans are monkeys, but the use of them for
Influence of naphazoline on CBF (n=12)

Fig. 4 Influence of naphazoline on the ciliary beat frequency of cultured human nasal mucosa cells (blue columns, n = 12; 59 single cells) and of cultured porcine tracheal mucosa cells (red columns, n = 12; 60 single cells). There is no significant difference between human and pig (paired t-test, P < 0.05). The level of significance refers only to the comparison of CBF under the influence of naphazoline in relation to CBF during the equilibration period of each species. In human cells, no significance could be seen (paired t-test, P < 0.05). In porcine tracheal cells, there was only slight difference between CBF under the influence of 0.1% naphazoline (filled circle) in relation to CBF during the equilibration period (paired t-test, P < 0.05)

The capability of developing beating cilia

In comparison with our previous investigations of in vitro cultured human nasal mucosa cells, we found the following characteristics in the cultured porcine tracheal mucosa:

- The speed of cell proliferation until adhesion to the coated tissue culture dishes happened within approximately 48 h and was similar to human cells;
- The formation of cell conjugations began as irregular clusters and became increasingly confluent after a few days. This process was also comparable to the human mucosa;
- The proliferation of the cells was associated with a simultaneous growth of cilia as in the human cultures and showed the same appearance (Fig. 1). We did not observe a delayed proliferation of cilia;
- The survival time of its cells with a constant beating frequency is at least as long as the one of human mucosa (Fig. 2).

The constancy of the beat frequency over a sufficient time

From our human experiments, we knew that a maximum investigation period of 180 min is sufficient for analyzing the effects of naphazoline and benzalkonium chloride. However, to make sure that the cilia of the porcine mucosa hold up the constancy of beat frequency over a time sufficient for testing the influence of various pharmacological substances in this species, we primarily measured the CBF of the porcine tracheal cells up to 360 min. It could be demonstrated, that the CBF was absolutely constant during the whole test period. Therewith we could show that the survival time of the porcine mucosa cells is by far sufficient for those experimental studies.

Moreover, in our previous studies, the human nasal mucosa showed a constant beating frequency of $7.3 \pm 1.4$ Hz over the chosen investigation time of 180 min, whereas the CBF of the porcine mucosa was $7.57 \pm 1.39$ Hz, thus demonstrating no statistical difference between both tissues. These findings also confirm that the porcine tracheal mucosa provides optimal prerequisites for investigating the influence of harmful or healing substances on the human mucosa.

The comparability of the reaction of the cilia to different drugs

The main goal of our study was to answer the question whether the reaction of porcine and human cilia to different
drugs is comparable. Therefore, we initially tested benzalkonium chloride, which we found in previous studies on human mucosa to be highly toxic on the cilia and their beat frequency. The functional and morphological reactions of the porcine ciliated cells to this substance were exactly the same as the reaction observed in the in vitro cultured human mucosa.

From the findings in our experiments with the human mucosa, we demonstrated, on the other hand, that naphazoline as a pure substance has no toxic influence on the survival of the cilia of human mucosa and their beat frequency with different concentrations. The average CBF was 105% compared with the mean equilibration frequency.

When testing the effect of naphazoline on the porcine mucosa we measured a CBF of 103.7% using a concentration of 0.01%, which was statistically equal to the CBF of the human mucosa.

With a concentration of 0.1%, the porcine CBF decreased to 92.5%. This value was significantly different to the equilibration period, but it was not significantly different to the human mucosal cells.

After rinsing out the substance, there was a rapid recovery of ciliary function. Whereas the first measurement after 20 min of rinsing out still showed the same CBF as at the end of exposure to naphazoline 0.1%, the CBF had already recovered to 96.8% (±19.6%) in the second measurement after 40 min.

Although there is no statistical difference, these initial data suggest that the porcine mucosal cells might have a slightly higher sensitivity than the human mucosa cells.

In conclusion, it became evident that the porcine tracheal mucosa can serve as an ideal tissue for the investigation of pharmacological effects on the sensitive physiology of mucosal cilia.

Our future investigations will also include hurtful substances causing neoplastic alteration or allergic reactions, for which this model also seems to be suitable.

In order to achieve comparable results to the earlier investigations with the human mucosa cells, in this study the DMEM culture medium has still been used. Due to our present studies, the airway epithelial cell growth medium (PromoCell) leads to a higher differentiation of cilia and therefore, will be used in our future experiments.

Summary

The presented data indicate that the in vitro culturing of porcine tracheal mucosa is well suited for investigation of the influence of different noxious substances as well as of protecting or healing agents under constant conditions. In the initial trial with the toxic substance benzalkonium chloride and the non-toxic z-symptomimetic drug naphazoline it could be shown that the reactions of the porcine tracheal mucosa are fairly equal to the reaction of in vitro cultured human mucosa. These results need to be confirmed by further research. Using porcine tracheal mucosa has the significant advantage that living animals do not need to be sacrificed because porcine tracheas are available as a normal by-product after slaughtering. We therefore believe that in vitro cultured mucosa from porcine–trachea can serve as an ideal material for researching the influence of substances affecting or healing human mucosa.

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