Effect of azelastine on sulphur dioxide induced impairment of ciliary motility in airway epithelium

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Abstract
Objective—The effect of azelastine on airway mucociliary transport function was studied by measuring ciliary motility of human bronchial epithelium in vitro with a photoelectric method.

Method—Bronchial epithelial cells were obtained by fibreoptic bronchoscopy, mounted in a Rose chamber, and perfused with Krebs-Henseleit solution. The preparations were placed on a microscope stage equipped with an illuminator, and the variations of light intensity caused by ciliary beating were detected by a photometer.

Results—The addition of azelastine to the perfusate increased ciliary beat frequency (CBF) in a dose dependent manner without ciliary discoordination. The mean (SE) maximal increase from the baseline value and the concentration required to produce a half maximal effect were 27±0.4(2.2)% and 9.2×10^-8 mol/l, respectively. Exposure of the cells to the perfusate containing 3 ppm sulphur dioxide rapidly decreased CBF by 59-2(5-0)%, and was accompanied by a reduction in intracellular cyclic AMP levels from 38±1(4-3) to 10±1(2-4) pmol/mg protein. This effect was prevented by pretreatment of cells with azelastine in a dose dependent manner.

Conclusions—Azelastine not only stimulates ciliary motility of airway epithelium and hence mucociliary transport function, but may also protect against sulphur dioxide induced ciliary dysfunction, probably by inhibiting intracellular cyclic AMP loss.

(Thorax 1993;48:542–546)

Removal of cellular debris, inhaled particles, and bacteria from the conducting airways of the respiratory tract are important factors in the defence mechanism of the lung. Movement of mucus containing adherent materials is directed posteriorly from the nasal cavity and upward from the lung, converging at the pharynx. This transport function is dependent on the beat frequency and coordination of cilia, mucus production, and the rheological properties of mucus. Several bronchial disorders including chronic bronchitis, asthma, and cystic fibrosis are associated with an impairment of mucociliary clearance, and exposure to various noxious agents such as cigarette smoke, ammonia, formaldehyde, and sulphur dioxide (SO2) can result in inhibition of ciliary function or ciliostasis.

Azelastine, 4-(p-chlorobenzyl)-2-(hexahydro-1-methyl-1H-azepin-4-yl)-1-(2H)-phthalazino hydrochloride, is an orally effective and long lasting antiallergic drug that prevents bronchospasm by inhibiting histamine release from mast cells and leukotriene synthesis in the lung. In addition, this drug possesses anti-inflammatory actions including suppression of superoxide generation by leukocytes, prostaglandin release from macrophages, and microvascular leakage. The purpose of the present study was (a) to characterise the SO2 induced inhibition of airway ciliary motility; (b) to determine whether azelastine can protect against airway mucociliary dysfunction; and (c) if so, to investigate its mechanism of action. We therefore studied ciliary motility of human airway epithelium in vitro by a photoelectric method.

Methods
Tissue Preparation
Fibreoptic bronchoscopy including transbronchial lung biopsy was carried out on 24 patients with a peripheral lung cancer. None had a history of respiratory tract infection for at least one month and any medication was stopped 14 days before the bronchoscopy. After topical anaesthesia of the upper airway with 4% lignocaine, a fibreoptic bronchoscope (Type 1T20; Olympus, Tokyo, Japan) was inserted transorally through an endotracheal tube. A standard bronchial brush (BC-9C; Olympus) was passed through the biopsy channel and epithelial cells were scraped at three different levels: the carina of the right upper lobe; the orifice of the middle lobe; and in a segment of the right lower lobe. Collection of samples was restricted to areas free of obvious mucosal invasion by carcinoma or acute inflammation, swelling, pus, or reddening. The cells were extracted from the brush by brisk agitation in a petri dish containing Krebs-Henseleit solution of the following composition (in mmol/l): NaCl, 118; KCl, 5-9; CaCl2, 2-5; MgSO4, 1-2; NaH2PO4, 1-2; NaHCO3, 25-5; glucose, 5-6. The cell
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Effect of sulphur dioxide on determination of CBF

The photometer (NFX-II; Hamamatsu Photonics, Hamamatsu, Japan) with an inbuilt periplanetary eyepiece, a limiting aperture, and a lateral focusing telescope was attached to the head of the microscope. The Rose chamber containing epithelial cells was set on a microscope stage maintained at 37°C by an air stream stage incubator (EB52; Omron, Tokyo) and perfused with warmed Krebs-Henseleit solution aerated with 95% O₂ and 5% CO₂ at a constant pressure of 20 cm H₂O. Light from the illuminator passed through the preparation in varying intensities because of the beating action of the cilia; this was viewed directly at a magnification of × 400. Variations of light intensity were detected by the photometer and transduced to voltage impulses. These electrical signals were amplified, displayed on the oscilloscope screen (LBO-522; Leader, Tokyo), and recorded by a pen recorder (VP6213; Panasonic, Tokyo). Throughout the experiment, the same group of cilia in each preparation was studied—that is, the relative position of the diaphragm of the photometer and the selected epithelial border were kept constant. The choice of the ciliated area was randomised by using an eyepiece graticule and selecting the group of cilia which lay closest to the crosspiece of the graticule on the equatorial line. The longitudinal axis of the field of the photometer was viewed perpendicular to the cell border and the measurements of CBF were averaged from clumps of two or more cells with borders free of debris. The mean intracellular coefficient of variation of CBF, defined as the ratio of the standard deviation to the mean value as a percentage, was less than 5%. There were no significant differences in CBF among epithelial cells obtained from three different levels of the airway.

In addition to CBF measurements, ciliary coordination was assessed by the image of beating recorded on a video camera (Type 1860–01; Hamamatsu Photonics) with a 0.75 inch video cassette recorder (VO-5800; Sony, Tokyo) capable of freeze frame replay. Ciliary discoordination was defined as the loss of the metachronal wave on the free border of the cell clump.15

STUDY DESIGN

The preparation was allowed to stabilise for 30 minutes before the measurement of CBF. After determination of baseline CBF, azelastine (Eizai Company, Tokyo) in a concentration of 10⁻⁴ mol/l was added to the perfusate and CBF was continuously recorded for the next 20 minutes. For time controls, measurements were obtained when only Krebs-Henseleit solution was used for perfusion. To generate the CBF dose-response curves, azelastine was cumulatively added in half molar increments from 10⁻⁷ to 10⁻³ mol/l, and the response to a given concentration was allowed to reach a plateau before proceeding to the next higher concentration.

To test any protective action of azelastine on ciliary dysfunction, we examined the effect of pretreatment of the cells with azelastine on the SO₂ induced decrease in CBF. The preparation was incubated with 10⁻⁴ mol/l azelastine for 20 minutes. When the CBF response reached a plateau, Krebs-Henseleit solution with a mixture of 95% O₂, 5% CO₂ and 3 ppm SO₂ bubbling through it perfused the cells and azelastine mixture for 20 minutes. In this series of experiments, cells were preincubated with various concentrations of azelastine at 10⁻⁷ to 10⁻¹ mol/l and the responses of CBF to SO₂ were determined in order to assess a dose-response relationship.

MEASUREMENT OF CYCLIC AMP

After 20 minutes exposure of epithelial cells to azelastine alone, SO₂ alone, or SO₂ plus azelastine, preparations were quickly removed from the chambers and sonicated with a bath type sonicator (PT-101; Sino Company, Tokyo) in ice cold 10% trichloroacetic acid. After the extraction of trichloroacetic acid with ether, the residue was dissolved in acetate buffer. The cyclic AMP levels were measured in duplicate by [³H] cyclic AMP (New England Nuclear, Boston, Massachusetts, USA),16 corrected for ether extraction of 86% recovery, and normalised for protein content of the cells by the Lowry method17 with bovine serum albumin as a standard.

STATISTICAL ANALYSES

The effects of azelastine and SO₂ on CBF were compared with the appropriate time controls (Krebs-Henseleit solution alone) using Friedman’s two way analysis of variance. Comparisons between the effects of SO₂ with or without azelastine were made with the Kruskal-Wallis one way analysis of variance test. A different donor of ciliated cells was used in each experiment; n refers to the number of patients from which samples were obtained, and a p value of <0.05 was considered significant.

Results

There was no significant difference in baseline CBF between the cells treated with azelastine and the Krebs-Henseleit solution time controls. The CBF did not change significantly during a 20 minute observation period in the control experiment, whereas addition of azelastine (10⁻⁴ mol/l) rapidly increased the mean (SE) CBF from 17.2 ± 0.7 Hz to a peak value of 21.8 ± 1.2 Hz within five minutes, being significantly faster than the control (p<0.05; n = 11), and remained faster thereafter (fig 1). The stimulatory effect of
Azelastine on the cilia was dose dependent with the maximal increase from the baseline CBF and the concentration required to produce a half maximal effect being 27.0 (4.2)% (p < 0.05, n = 9) and 9.2 × 10^-6 mol/l, respectively (fig 2). This was accompanied by corresponding increases in intracellular cyclic AMP levels (fig 3) where the threshold concentration of azelastine (3 × 10^-6 mol/l) was identical to that for ciliary stimulation.

The addition of 3 ppm SO2 to the perfusate caused a rapid reduction of ciliary motility, with the CBF values significantly slower than the control experiments at 2.5–20 minutes after the initiation of SO2 exposure (p < 0.05, n = 10), an effect that was significantly inhibited by pretreatment of cells with 10^-6 mol/l azelastine (fig 4). The maximal decrease in CBF from the baseline value induced by SO2 was 59.2 (5.0)%, and azelastine protected against this inhibition in a dose dependent fashion. The decrease in CBF was 24.4 (3.9)%, 17.6 (2.2)%, 9.1 (2.8)%, and 6.6 (1.0)%, for concentrations of 3 × 10^-7, 10^-6, 3 × 10^-5, and 10^-4 mol/l azelastine, respectively (fig 5). Ciliary discoordination was not observed among adjacent cilia on the same cells or several bordering cells in any of the experiments.

Exposure of epithelial cells to SO2 decreased intracellular cyclic AMP levels from 38.2 (4.3) to 10.1 (2.4) pmol/mg protein (p < 0.001, n = 12). Azelastine pretreatment prevented this effect in a dose dependent manner so that the reduction of cyclic AMP levels was completely reversed at concentrations of 3 × 10^-6 and 10^-5 mol/l (fig 6).

Azelastine (10^-4 mol/l) was added to the perfusate at time 0 (arrow). Bar lines indicate 1 SE (n = 9 for each group). *p < 0.05 v Krebs-Henseleit controls.

Figure 1 Effect of azelastine on ciliary beat frequency (CBF) of human bronchial epithelium. Azelastine (10^-4 mol/l) was added to the perfusate at time 0 (arrow). Bar lines indicate 1 SE (n = 11 for each group), *p < 0.05 v Krebs-Henseleit controls.

Figure 2 Concentration dependent effect of azelastine on ciliary beat frequency (CBF) of human bronchial epithelium. Azelastine was cumulatively added to the perfusate in half molar increments. Data are expressed as percentage of the baseline CBF obtained before addition of azelastine. Bar lines indicate 1 SE (n = 9 for each group). *p < 0.05 v Krebs-Henseleit controls.

Figure 3 Effect of azelastine on intracellular cyclic AMP levels of human bronchial epithelium. The cells were incubated with various concentrations of azelastine for 20 minutes and the cyclic AMP concentrations were determined by radioimmunoassay. Bar lines indicate 1 SE (n = 9 for each bar). *p < 0.05, **p < 0.01 v Krebs-Henseleit controls.

Figure 4 Effect of SO2 on ciliary beat frequency (CBF) of human bronchial epithelium. SO2 (3 ppm) was added to the perfusate at time 0 (arrow) in the cells pretreated with and without azelastine (10^-4 mol/l). Preparations perfused with Krebs-Henseleit solution alone served as control experiments. Bar lines indicate 1 SE (n = 10 for each group). *p < 0.05 v controls; †significantly different from values for SO2 alone.

Figure 5 Concentration dependent effect of azelastine on the SO2 induced decrease in ciliary beat frequency (CBF) of human bronchial epithelium. The cells were pretreated with various concentrations of azelastine for 20 minutes and exposed to Krebs-Henseleit solution containing SO2. Data are expressed as percentage of the baseline CBF obtained before SO2 exposure. Bar lines indicate 1 SE (n = 9 for each bar). *p < 0.05, **p < 0.01, ***p < 0.001 v SO2 alone.
Discussion

We have shown that azelastine stimulates ciliary motility of human bronchial epithelium and protects against inhibition of cilia by SO₂, presumably by the accumulation of intracellular cyclic AMP.

It has been shown that ciliary motility in epithelial cells is regulated by various endogenous receptors including cyclic AMP, 
\[ \text{Ca}^{2+}, \] and protein kinase C. In the present study azelastine increased CBF of airway epithelium in a dose dependent fashion, the concentration required to produce a half maximal effect (9.2 × 10⁻⁶ mol/l) being similar to that previously found to inhibit histamine release in mast cells (5 × 10⁻⁶ mol/l) and to inhibit platelet activating factor release in macrophages (1.6 × 10⁻⁵ mol/l). Azelastine also raised intracellular cyclic AMP levels, as shown by Akagi et al., which suggests that azelastine causes cyclic AMP to accumulate in mast cells. Increased availability of cyclic AMP may activate glycoprotein kinase C. The increase in cyclic AMP production is mediated by SO₂, which reverses the azelastine induced reduction of CBF and cyclic AMP levels. We therefore speculate that this reversal may be produced by the inhibition of cyclic AMP loss.

In conclusion, the present studies indicate that the antiallergic drug azelastine not only enhances the beating action of human bronchial cilia but also protects against the impairment of ciliary motility, and suggest that this drug could be of value in patients with mucociliary dysfunction.

We thank Yoshimi Sugimura and Masaaki Shino for their technical assistance.

This study was supported in part by a grant for scientific research No. 63770524 from the Ministry of Education, Science and Culture of Japan.

Adventitia

The lung has no parenchyma

In case this proposition seems too iconoclastic, I will restate it as follows: within the lung there is no tissue that can usefully be identified and named “parenchyma.” I have urged for many years that the use of this word in relation to the lung is inappropriate and should be avoided. Avoidance is not difficult. George Simom managed to write an excellent textbook on lung radiology (Principles of chest x-ray diagnosis, 3rd edn. Butterworth, London, 1971) without once using the phrase “lung parenchyma”. Fifteen years ago I confirmed in a formal study (Br J Dis Chest 1978;72:1–12) that the meanings attached to this phrase by physicians and radiologists on both sides of the Atlantic are so varied that it is virtually meaningless. In many instances the word “parenchyma” can be omitted without altering the apparently intended meaning; in others it is necessary to ascertain the sense in which this word is being used. This sense is usually discordant with accepted usages of “parenchyma” in relation to other organs.

How did this verbal confusion arise? In relation to secreting glands or the liver, “parenchyma” refers to tissues composed of cells having functions specific to the organ concerned, as opposed to supporting and vascular tissues; this usage is unequivocal and, I believe, generally accepted. In the lung I suppose the cells that might be considered to be most organ specific are the type II pneumocytes, but it would be absurd to call them the lung parenchyma! The function of the lung depends not upon any one type of cell, but upon the structural and functional interaction of elements of its vasculature and airways to bring about orderly gas exchange in the alveoli. In general it seems that what most people mean, when they speak of the lung parenchyma, is its peripheral gas exchanging part. There is a fairly well defined demarcation at the ends of the terminal bronchioles between the part in which movement of gas is by diffusion both within airspaces and across alveolar membranes, and airways in which it is by ventilatory to and fro movement. It is now agreed that the part of the lung associated with a terminal bronchiol should be called an acinus; thus I think that what is usually meant by the “lung parenchyma” is the pulmonary acini. If you hear or read these words, consider first whether the word “parenchyma” contributes anything; it can often be eliminated without loss of meaning. It may be evident from context that the intended reference is to the pulmonary acini, as opposed to the airways. If you do not think that anyone using these words has this meaning in mind, ask what he or she really means. I recommend that this weasel word be eliminated from discourse concerning the lung.

G SCADDING
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Thorax 1993 48: 542-546
doi: 10.1136/thx.48.5.542