Cigarette smoke inhalation patterns and bronchial reactivity

D R Taylor, W D Reid, P D Paré, J A Fleetham

From the Department of Medicine, University of British Columbia, Vancouver, Canada

ABSTRACT The manner in which a cigarette is smoked varies considerably between individuals and may be an important determinant of the altered bronchial reactivity observed in cigarette smokers. Twenty smokers were examined to determine the relationship between cigarette smoke inhalation patterns and bronchial reactivity. Inhalation patterns were measured non-invasively with a respiratory inductive plethysmograph and these were related to the provocative concentration of histamine that caused a 20% fall in FEV\(_1\) (PC\(_{20}\)) and to the cough threshold for inhaled citric acid. Histamine PC\(_{20}\) values were inversely correlated with depth and rate of inhalation. Cough threshold was inversely correlated with greater cigarette consumption and with depth of inhalation.

There is growing evidence that non-specific bronchial reactivity is greater in smokers than in non-smokers\(^1\) and that this effect is related to the numbers of cigarettes smoked.\(^2\) There are, however, several reports\(^3\)\(^-\)\(^7\) that have found no difference in bronchial reactivity between symptomless smokers and non-smokers. Cigarette smoke inhalation patterns vary considerably between individuals, with large variations in the depth and rate of inhalation and the duration of breath hold.\(^8\) These factors are likely to modify airway deposition of cigarette smoke particulate material and as such may be important determinants of cigarette smoke related lung disease.\(^9\)

We have measured inhalation patterns and bronchial reactivity in a group of chronic smokers with normal pulmonary function to examine further the relationship between cigarette smoking and bronchial reactivity to inhaled histamine.

Methods

SUBJECTS Twenty subjects (six men and 14 women) completed the study. Their ages ranged from 20 to 54 years and they smoked from 10 to 60 cigarettes a day, with a cumulative consumption ranging from four to 52 pack years. Eight subjects had a cough but none had any other symptoms suggesting atopy or asthma and none was taking any bronchodilator. All subjects had a forced expiratory volume in one second (FEV\(_1\)) and forced vital capacity (FVC) over 85% of the predicted value, on the basis of the best of three expirations into a spirometer (Collins Modular Function Analyser, Braintree, Massachusetts, USA). Subjects gave informed consent to the study, which was approved by the university ethics committee.

PROTOCOL The study was carried out during the morning on two separate days. On day 1 monitoring of smoke inhalation patterns was performed and followed by an assessment of the airway response to inhaled histamine or the cough response to inhaled citric acid. On day 2 the other bronchial challenge test was performed. The order of administration for the two challenge tests was randomised. The bronchoconstrictor response to histamine has been shown to be unaffected by cigarette smoking immediately before the test.\(^10\) On three additional consecutive days we restudied smoke inhalation patterns and cough threshold in five subjects to assess intrasubject variability.

MONITORING OF SMOKE INHALATION PATTERNS Respiratory inductive plethysmography (Respitrace, Non-invasive Monitoring Systems Inc, Ardsley, New York) was used to monitor the pattern of breathing and smoke inhalation.\(^11\) Calibration was carried out by the simultaneous equation method with subjects standing and semirecumbent. Validation against a known inspired volume from a Collins wet spirometer
was performed with subjects semirecumbent before and after observation of inhalation pattern. A change of more than 10% in volume calibration occurred in three subjects and their results were excluded from the analysis.

After calibration each subject remained semirecumbent in a quiet room. Tidal breathing was monitored for 10 minutes, after which subjects were instructed to smoke two of their usual brand of cigarettes with an intervening rest period of 10 minutes. They were permitted to read during the period of observation and every effort was made to minimise any disturbance. Subjects were not informed of our specific interest in their breathing pattern. Each cigarette inhalation was noted by continuous observation through a one way mirror.

Signals from the respiratory inductive plethysmograph were recorded on a Grass 78 polygraph recorder. Mean values for tidal volume (VT), inspiratory time (Ti), and mean inspiratory flow rate (VT/Ti) were calculated from 15 consecutive breaths during the final minute of tidal breathing in the initial rest period. During cigarette smoking, inhalation volume, (Vi), inhalation time (Ti), inhalation flow rate (Vi/Ti), and breath hold time were determined as mean values from all inhalations from both cigarettes apart from the first and last inhalation, which may be atypical. Inhalation volume (Vi) is expressed as a ratio of the vital capacity (Vi/VC) and mean inhalation flow rate (Vi/Ti) expressed as a ratio of the resting mean inspiratory flow rate (Vi/Ti:VT/Ti).

**AIRWAY RESPONSE TO HISTAMINE**

The airway response to inhaled histamine was determined by the technique of Juniper et al as modified by Lam et al. Doubling concentrations of histamine ranging from 0-03 to 16-0 mg/ml were administered until the maximum concentration had been given or a decrease in FEV1 of 20% or more had occurred. Each histamine solution was given for two minutes with a Bennett Twin nebuliser (output 0-23 ml min⁻¹) at a flow rate of 8 l min⁻¹. Subjects breathed normally wearing a face mask and nose clip. Forced expiratory manoeuvres were performed 30 and 90 seconds and three minutes after inhalation of a control solution of phosphate buffered saline, and at similar intervals after each concentration of histamine.

The concentration of histamine producing a 20% decrease in FEV1 (PC20) from the control value was determined by interpolation. In the seven subjects who did not show a 20% decrease in FEV1 after the maximum dose of histamine the PC20 was obtained from the line of best fit on the log dose-response curve extrapolated to infinity.

**COUGH THRESHOLD**

The cough threshold was determined by using a modification of the technique described by Bickerman and Barach and Empey et al. In a random single blind fashion each subject inhaled a control solution of normal saline followed by progressively increasing concentrations (the percentages being 0-5, 1, 2, 4, 6, 8, 12, 16, 24, 32, 48 and 64) of crystalline citric acid monohydrate (Fisher Scientific, New Jersey) dissolved in normal saline. Solutions were inhaled every five minutes from a Bird micronebuliser, the subject carrying out a slow inspiratory vital capacity manoeuvre over five seconds. The cough threshold was defined as the lowest concentration of citric acid that consistently elicited an involuntary cough during three separate inhalations.

**STATISTICAL ANALYSIS**

The relationship between variables was analysed by Spearman’s rank correlation method in view of the non-parametric distribution of the data.

**Results**

Smoking history, pulmonary function, cigarette smoke inhalation pattern, and histamine PC20 values are presented for each individual in the table. There was considerable intersubject variability in the mean depth of inhalation Vi/VC (range 7-6-34-5%), mean rate of inhalation Vi/Ti/VT/Ti (range 0-65-3-3), and mean duration of breath hold (range 0-4-3 s). There was no significant relationship between these three indices. There was also no significant relationship between cigarette smoke inhalation patterns and resting breathing patterns. In our subgroup of five subjects the mean (SD) coefficients of variation (SD/x 100) for these indices were 3-6 (1-8) for Vi/VC, 8-2 (3-2) for Vi/Ti/VT/Ti, and 52 (17) for breath hold time.

Histamine PC20 values ranged from 2-2 to >16 mg/ml. A cough threshold was obtained in all subjects, ranging from 2% to 64% citric acid. In our subgroup of five subjects the mean (SD) coefficient of variation for cough threshold was 12-6 (4-3).

Cough threshold values correlated positively with histamine PC20 values (rho(p)=0-38, p<0-05) and correlated negatively with cigarette consumption expressed as cigarettes/day (rho(p)=0-39, p<0-05) or pack/years (rho=0-49, p<0-05) (fig 1). There was no significant correlation between PC20 and cigarette consumption. When related to smoke inhalation patterns PC20 correlated negatively with depth of inhalation (Vi/VC)(rho=0-38, p<0-05) (fig 2) and rate of inhalation Vi/Ti/VT/Ti (rho=0-42, p<0-05) and cough threshold was also correlated negatively with depth of inhalation (Vi/VC) (rho=0-50, p<0-05) (fig 3).
Cigarette smoke inhalation patterns and bronchial reactivity

Smoking history, pulmonary function, smoke inhalation pattern, and bronchial reactivity to histamine in 20 normal subjects

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<th>Subject No</th>
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<th>Packs/year</th>
<th>FEV₁ (%)</th>
<th>Pred</th>
<th>Cough</th>
<th>V₁/Vc (%)</th>
<th>V₁/Ti (%)</th>
<th>V₁/Ti (1s⁻¹)</th>
<th>V₁/Ti (mg/ml)</th>
<th>Breath hold (mean, s)</th>
<th>PC₂₀ (mg/ml)</th>
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Mean ...

SEM ...

*) Derived by extrapolation.
†Geometric mean + SEM.
‡Geometric mean - SEM.

Vi—inhalation volume; VC—vital capacity; Vi/Ti—inhalation flow rate; Vi/Ti—inspiratory flow rate; PC₂₀—provocative concentration of histamine causing a 20% fall in FEV₁; + indicates presence and — absence of cough.

Fig 1 Cough threshold (% citric acid) compared with ranked cigarette consumption (pack years) in 20 subjects.

Fig 2 Ratio of cigarette smoke inhalation volume to vital capacity (Vi/Vc) compared with the ranked values for the concentration of histamine producing a 20% decrease in FEV₁ (PC₂₀) in 20 subjects.
cigarette smoking
The association between cigarette smoking and chronic airflow obstruction is well established. Nevertheless, many people smoke for prolonged periods without developing any airflow obstruction. It has been customary in epidemiological studies of cigarette smoking to quantify smoking history with a single variable, the total number of cigarettes smoked. This variable, however, does not take into account other important factors that may determine the relative concentration of smoke constituents reaching the lung, such as the degree of inhalation. There are some data on the frequency and possible significance of inhalation of cigarette smoke. Sixty per cent of moderate smokers have been found to inhale, and they develop more airflow obstruction than smokers who do not inhale. There is also evidence that smokers who inhale have significantly higher carboxyhaemoglobin and plasma nicotine concentrations. These data, however, have been obtained by subjective assessment of the degree of inhalation, which is notoriously unreliable.

An objective study of smoking patterns has been difficult because most measuring devices have required some form of mouthpiece, which in itself may affect both the ability to smoke and the pattern of smoking. The respiratory inductive plethysmograph enables accurate monitoring of respiration without an airway and this has been used to quantify smoking patterns.

In this study we have confirmed that in a group of smokers with no airflow obstruction there are large differences between subjects in inhalation rate and volume and in duration of breath hold after inhalation. These different inhalation patterns might influence the concentration and deposition of cigarette smoke constituents within the lung. Aerosol deposition tends to increase with depth of inhalation, whereas more rapid inhalation may reduce lung deposition owing to proximal impaction. Furthermore, a breath hold after inhalation of cigarette smoke would favour gravity settlement of a small proportion of particles that would otherwise remain airborne.

Although some of the data are conflicting, there is increasing evidence that chronic smoking is associated with bronchial hyperreactivity. Gerrard and co-workers have reported greater bronchial reactivity in smokers with symptoms than in age matched non-smokers, and Malo et al have reported similar results in symptomless smokers. Buczko and colleagues also found greater non-specific airway responsiveness in smokers than in non-smokers and showed a relationship to the amount smoked. In a large prospective study Taylor and coworkers showed that, in men with a baseline FEV1 over 80% predicted, bronchial reactivity was significantly greater among smokers and slightly greater among ex-smokers than among non-smokers, and that the increased bronchial reactivity in the smokers was associated with an accelerated decline of FEV1. Several studies have found no difference in histamine response between smokers and non-smokers. These investigations, however, examined young symptomless smokers and as such may have selected the less responsive subjects. Our results are consistent with the evidence that chronic cigarette smoking is associated with increased bronchial reactivity and show that cigarette smoke inhalation patterns are significant determinants of this increased bronchial reactivity. Both the bronchoconstrictor response to histamine and the cough response were enhanced in the subjects who inhaled most deeply. Furthermore, subjects who inhaled faster also had increased bronchoconstrictor responses. Our results do not support an alternative hypothesis—namely, that increased bronchial reactivity in smokers might cause them to modify their inhalation pattern in an attempt to minimise the irritant effects of the cigarette smoke.

Wanner and co-workers have recently demonstrated that the variability in airway responsiveness to histamine in normal smokers is related to differences in the dose deposited in the airways. As the dose of histamine deposited is related in part to breathing pattern, an alternative explanation for the relationship between cigarette smoke inhalation patterns and increased bronchial reactivity is that subjects who inhale cigarette smoke most deeply also inhale larger quantities of the provocative agent. This explanation is not valid for the cough threshold data, as this was
performed with a slow vital capacity manoeuvre. The histamine bronchial challenge was performed over two minutes of tidal breathing and we do not have measurements of tidal volume at this time. We do know that before measurement of smoke inhalation patterns there was no significant relationship between smoke inhalation pattern and resting breathing pattern. We think therefore that the relationship between smoke inhalation patterns and bronchial reactivity is unlikely to be due to a greater depth of inhalation with histamine in subjects with large inhaled volumes during cigarette smoking.

Cough is a frequent manifestation of bronchial hyperreactivity, yet there are few published reports on the variability of cough threshold in health and disease. As such, measurement of cough threshold represents a valuable additional method of assessing non-specific bronchial reactivity. This study confirms the large intersubject differences in cough threshold observed in previous reports and shows that cough threshold is related to cigarette consumption and depth of inhalation. The greater the cigarette consumption and the deeper the smoke inhalation, the lower the cough threshold. Cough and wheeze are major symptoms of increased bronchial reactivity; yet some patients with asthma present primarily with cough, whereas others have bronchoconstriction and no cough. The present study shows that cough threshold is related to bronchial reactivity to histamine in chronic smokers. Previous reports suggest that cough and bronchoconstriction are due to quite different mechanisms. Inhaled bupivacaine blocks cough induced by citric acid but not bronchoconstriction induced by histamine in non-asthmatic subjects, and inhaled lignocaine blocks cough but not the bronchoconstrictor response to inhaled distilled water in subjects with asthma. It has also been shown that absence of a permeant anion in iso-osmolar aerosols causes cough but not bronchoconstriction.

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