Cigarette smoke inhalation and the acute airway response

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ABSTRACT The acute airway response to smoking varying numbers (one to four) of identical cigarettes in rapid succession and smoking single cigarettes of differing tar/nicotine yields was assessed repeatedly in 13 healthy smokers. The airway response was variable, indicating airway narrowing consistently in only three subjects. There appeared no difference between forced expiratory volume in one second (FEV₁) and forced vital capacity (FVC) by changes in venous blood nicotine or percentage carboxyhaemoglobin. When five smokers inhaled smoke directly from a cigarette acute airway narrowing was consistently observed. A normal smoking pattern consisting of an initial drag of smoke into the mouth, followed by a pause before inhalation of smoke diluted with air, did not consistently cause airway narrowing although similar amounts of smoke as the direct drag were inhaled as assessed by changes in venous blood nicotine levels.

The manner of smoke inhalation affects the relative concentrations of the different constituents of smoke reaching the lungs and also appears to be the main determinant of the acute airway response to smoking, which was unrelated to the number of cigarettes smoked or the tar content of the smoke. This suggests that patterns of smoke inhalation may influence the pathogenesis of bronchial disease associated with smoking.

Despite a well-advertised association with carcinoma of the lung and chronic obstructive bronchitis, cigarette smoking remains a remarkably common habit, and this is even more surprising as the habit involves inhalation of smoke into the lungs. Cigarette smoke, when inhaled, behaves in a similar way to other irritant gases, vapours, and dusts, in causing acute dyspnoea, coughing, and acute airway narrowing. These responses probably represent vagally mediated protective reflexes as they promote proximal deposition of smoke particles and their subsequent ejection from the lungs by coughing and mucociliary clearance. However, when large numbers of smokers have been studied only a minority appear to develop any of these responses and only a few develop acute airway narrowing after smoking a single cigarette.

The question as to how the majority of smokers avoid an acute airway response to inhaled cigarette smoke has not been answered and is important as it suggests two conflicting theories concerning the relation between the acute response and the development of chronic obstructive bronchitis in a minority of smokers. It could be argued that the absence of an acute protective airway response enables airways to be chronically exposed to those constituents of cigarette smoke which promote chronic intrinsic airway disease. Alternatively, those individual smokers exhibiting an acute cigarette-induced response may represent a susceptible population who through repeated provocation proceed to develop chronic obstructive bronchitis.

Previous work on the subject has suffered from difficulties with measurement of the amount of smoke inhaled so that differences in inhaled dose may account for the previously reported variation in acute response. Furthermore, cigarettes have undergone a progressive reduction in tar and nicotine yield over the last few years which makes interpretation of earlier results difficult as most modern cigarettes yield a potentially less irritant smoke.

We have attempted to identify those factors which may be considered important in determining the acute airway response to cigarette smoke.
and by using changes in venous blood nicotine or carboxyhaemoglobin we have estimated the amount of smoke inhaled. By varying the number of identical cigarettes smoked in a rapid succession and varying the tar/nicotine yield of cigarettes we have attempted to identify whether quantity or type of cigarette smoke are also important determinants. Finally, the effects of differing patterns of smoke inhalation have been compared in order to decide whether the way in which smoke is inhaled, rather than quantity or type of smoke, influences the acute airway response.

Methods

Five separate experiments were undertaken. From a group of 13 asymptomatic male smokers, individuals were selected to take part in each (table 1). Each subject had a normal FEV1 and their mean age was 27 years.

<table>
<thead>
<tr>
<th>ONE TO FOUR CIGARETTES</th>
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In experiments 1 and 2 an assessment was made of the acute airway response to smoking differing numbers of a low to middle tar filtered cigarette (table 2) in rapid succession. Each of four subjects smoked on four separate days one to four cigarettes, the order being randomised by means of a 4×4 latin square experimental design. Airway response to smoking in experiment 1 was determined by forced spirometry, using a dry spirometer on-line to a minicomputer providing 12 variables from an analysis of maximal expiratory flow volume curves (MEFVC) (table 3). Using the same experimental design inspiratory airway resistance and associated lung volume measured in a constant volume whole body plethysmography were used to assess acute airway narrowing in experiment 2.

FIVE CIGARETTE TYPES

Experiments 3 and 4 each involved five subjects (table 1), smoking on separate days a single cigarette selected from one of five types (table 2) of differing tar/nicotine yields. One of these cigarettes contained 30% of a non-tobacco substitute (NSM) and one was a full strength unfiltered cigarette. The order by which each subject smoked each type of cigarette was randomised using a 5×5 latin square design. Airway response to smoking was measured in experiment 3 with MEFVC analysis and airway resistance was used in experiment 4.

| PATTERNS OF INHALATION |
In experiment 5 five subjects smoked a single cigarette of low to middle tar category (table 2) in three different ways on separate days (table 1): (1) drawing smoke into their mouths but not inhaling (mouth smoking); (2) drawing smoke directly from the cigarette into their lungs; (3) smoking normally. The airway response was determined with airway resistance measurements and the order of different smoking patterns was randomised. Breathing patterns during this study were also monitored with a single pair of magnetometer coils across the anteroposterior diameter

Table 1 The separate experiments (1–5) showing the type of study, the measurement used, and which subjects took part

<table>
<thead>
<tr>
<th>Method of assessment</th>
<th>MEFVC Airway resistance</th>
<th>MEFVC Airway resistance</th>
<th>MEFVC Airway resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experiment number</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Subject</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>1</td>
<td></td>
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<tr>
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<td>+</td>
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<td>+</td>
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</tr>
<tr>
<td>13</td>
<td>-</td>
<td>-</td>
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</tr>
</tbody>
</table>

Table 2 The different types of cigarette

<table>
<thead>
<tr>
<th>Tar category</th>
<th>Plain/filtered</th>
<th>Nicotine yield</th>
<th>Carbon monoxide yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low tar (30% NSM)</td>
<td>Filtered</td>
<td>0.70 mg/cig</td>
<td>11.0 mg/cig</td>
</tr>
<tr>
<td>Low tar</td>
<td>Filtered</td>
<td>1.01 mg/cig</td>
<td>12.1 mg/cig</td>
</tr>
<tr>
<td>Low to middle tar</td>
<td>Filtered</td>
<td>1.36 mg/cig</td>
<td>19.3 mg/cig</td>
</tr>
<tr>
<td>Middle to high tar</td>
<td>Plain</td>
<td>1.86 mg/cig</td>
<td>15.0 mg/cig</td>
</tr>
<tr>
<td>High tar</td>
<td>Plain</td>
<td>3.39 mg/cig</td>
<td>18.2 mg/cig</td>
</tr>
</tbody>
</table>

Table 3 Measurements from the maximal expiratory flow volume curves

<table>
<thead>
<tr>
<th>VITAL CAPACITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forced expired volume in one second</td>
</tr>
<tr>
<td>Forced expired ratio</td>
</tr>
<tr>
<td>Forced expired flow rate</td>
</tr>
<tr>
<td>Inspiratory flow at 75% vital capacity</td>
</tr>
<tr>
<td>Inspiratory flow at 60% vital capacity</td>
</tr>
<tr>
<td>Inspiratory flow at 50% vital capacity</td>
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<tr>
<td>Inspiratory flow at 25% vital capacity</td>
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<tr>
<td>Mean flow between 70-25% vital capacity</td>
</tr>
<tr>
<td>Mean flow between 60-20% vital capacity</td>
</tr>
<tr>
<td>Time to no flow</td>
</tr>
<tr>
<td>Time to peak flow</td>
</tr>
</tbody>
</table>
of the ribcage at the level of the xiphisternum, being calibrated for volume measurement before the experiment began. By observing the glow of the burning cone of the cigarette as each drag of smoke was taken, the duration and timing of each drag was recorded in relation to the breathing pattern on a multichannel recorder.

DOSE OF SMOKE INHALED
The amount of smoke inhaled while smoking, was measured in experiments 1, 2, and 5 by determining changes in level of venous blood nicotine from samples taken before and two minutes after completing the last cigarette. In experiments 3 and 4 change in venous carboxyhaemoglobin level was determined spectrophotometrically. We considered that this would provide a more reliable assessment of smoke inhalation, as carbon monoxide deliveries on the different cigarettes were more comparable than their nicotine yields (table 2).

STATISTICAL ANALYSIS
In each experiment six measurements (either MEFVVC or airway resistance) were performed in rapid succession by every subject before each smoking session and repeated within two minutes of completing the test cigarettes. Factorial analysis of variance was used to assess the results in each experiment. By using the F ratios, the significance of the main factors could be determined—for example, variation caused by different subjects, order of each treatment, or effects of smoking were compared with the background or residual variation found from the variation within the groups of six replicated measurements. We were also able in a similar way to determine the significance of interaction between these factors; for example, to decide if all subjects reacted in the same way to smoking, the variation attributable to the interaction between the factors, subjects, and treatments being compared with the residual variation to provide an F ratio.

In all five experiments each variable used to assess the airway response was analysed separately. This amounted to 12 variables derived from the MEFVVC (table 3). In the body box studies, in addition to airway resistance (Raw) and the associated lung volume (Vtg), the variables airway conductance and specific airway conductance (sGaw) together with its logarithmic transformation were also calculated. Specific airway resistance (sRaw) and a covariance correction of airway resistance with lung volume as the covariate were determined as well. Each of these six variables was also analysed separately. Although many of these variables measured in either the MEFVVC and body box are not independent of one another, the risks of a falsely positive result are greatly enhanced by the large number of variables being examined. For example, in the MEFVVC studies where 12 variables are under study the chances of finding such a significant result by accident may well exceed 80%. To avoid this type of statistical error and to provide the equivalent of a p value of 0.05 for each variable, the overall p value for each experiment when testing the F ratios for significance was 0.0043 and 0.0073 for the MEFVVC studies and body box studies respectively.

When determining the size of change in each variable after smoking, the differences between mean values of each variable before and after smoking were contrasted with a 95% confidence interval derived from the standard error using a multiple comparison test of Dunn. Such a method was applied once more to reduce the risks of statistical error as many comparisons of means were being performed; for example in experiments 3 and 4, 25 pairs of means were compared.

![Figure 1](https://example.com/fig1.png)

**Fig 1** Upper graph shows the mean change in covariance corrected airway resistance for all subjects. Lower graph shows the changes in each individual related to cigarette consumption and the symbols represent each subject: subject 1 ■, subject 2 ○, subject 3 ●, subject 4 □.
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Results

Experiments 1 and 2
Airway resistance
There was no increase in airway resistance after smoking any of the number of cigarettes when the mean results for all four subjects were considered (fig 1). However, the subjects behaved differently after smoking, the interaction between subjects and treatments being significant for the variables Raw, log sGaw and sRaw (p values less than 0.0073). Viewing the changes in airways resistance after smoking against the 95% confidence interval it will be seen that subject 1 showed airway narrowing after one, two, three, and four cigarettes (fig 1), whereas no change in airways resistance occurred with the other subjects no matter how many cigarettes they had smoked.

Maximal expiratory flow volume curves
When using the MEFVC analysis to detect airway narrowing, no overall change occurred after smoking, taking all four subjects as a group (fig 2 for PEFR). Individual smokers again showed differing responses particularly for PEFR and FEV₁ (p values less than 0.0043) with only subject 6 showing evidence of reduction of PEFR and then only after two, three, and four cigarettes had been smoked in succession (fig 2).

Change in level of nicotine
Although there was a general tendency for more smoke to be inhaled with increasing numbers of cigarettes smoked (fig 3) subjects 1 and 6 did not appear to inhale more smoke than the other subjects, nor was there any apparent relationship between the change in venous blood nicotine and change in Raw or PEFR.

Experiments 3 and 4
Airway resistance
Only for the high tar cigarettes was there a mean increase in airway resistance for all five subjects (fig 4). However, most of this was contributed by one individual, subject 10, and he with subject 12 were the only ones to show evidence of acute airway narrowing after smoking (fig 4). For the remaining subjects there was no change in Raw after smoking. There was no relationship between the tar yields of the cigarettes and the increase in airway resistance in any of the subjects.

Maximal expiratory flow volume curves
The subjects reacted differently to smoking. There was a trend of a greater reduction in PEFR with higher tar cigarettes but for the variables PEFR and mean flow rates between 75-25% and 60-20%
of vital capacity there was significant interaction between treatments and subjects (p value less than 0.0043). This can be seen in fig 5 where subject 11 showed a reduction in PEFR on one occasion, subjects 8 and 12 on two occasions, and subject 10 reacted to all five cigarettes. Tar yield of the cigarettes did not appear to be related to the degree of reduction in PEFR except in subject 10.

**Change in level of carboxyhaemoglobin**

The subjects who had shown acute airway narrowing did not appear to have inhaled more cigarette smoke as measured by change in carboxyhaemoglobin than the non-responders nor was there any relationship between the degree of airway narrowing and amount of smoke inhaled.

**EXPERIMENT 5**

**Patterns of smoke inhalation**

The “normal” pattern of smoking in each of the five subjects appeared to consist of two stages. The first stage was a “mouth” phase when smoke...
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Table 4  Mean volumes (with standard errors) of inhaled smoke obtained from the measurement of anterior chest wall for the differing patterns of smoking. The change in venous blood nicotine is also recorded together with number of drags of smoke.

<table>
<thead>
<tr>
<th>Subject</th>
<th>Pattern of smoking</th>
<th>Normal (ml)</th>
<th>Direct inhalation (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouth</td>
<td></td>
<td>Number of puffs</td>
<td>Change in nicotine</td>
</tr>
<tr>
<td>9</td>
<td></td>
<td>11</td>
<td>-1.5</td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>5</td>
<td>-0.1</td>
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<tr>
<td>11</td>
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<td>19</td>
<td>-0.3</td>
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<tr>
<td>12</td>
<td></td>
<td>7</td>
<td>-1.0</td>
</tr>
<tr>
<td>13</td>
<td></td>
<td>14</td>
<td>-1.1</td>
</tr>
</tbody>
</table>

was drawn into the mouth without any evidence of inhalation, the anterior chest wall remaining stationary. The second stage began with a pause of variable duration often associated with removing the cigarette from the mouth which was followed by inhalation of smoke (fig 6). By contrast on being directed only to mouth smoke, although a drag of smoke was taken from the cigarette, no inhalation of smoke took place (fig 6). Only three subjects were able to inhale smoke directly from the cigarette into the lungs to provide an assessment of direct smoke inhalation (fig 6). Subjects 10 and 11 failed to do so because of intolerance of the smoke, although both were able to perform "mouth" and normal smoking patterns.

Volumes of "smoke" inhaled and change in nicotine

There was little relationship between the volume of smoke inhaled, numbers of puffs of smoke and changes in venous blood nicotine (table 4). Both direct inhalation of smoke and normal smoking led to comparable volumes of smoke plus air being drawn into the lungs, suggesting that dilution of the smoke with air in both patterns of smoking was comparable. Where no smoke was inhaled (mouth smoking and subject 10), no real change in nicotine level was observed (table 4).

Airway resistance

Subject 10 once again showed an increase in airway resistance after normal smoking and was the only subject to do so in this experiment. In subjects 9, 12, and 13 who failed to respond to normal smoking patterns, a direct inhalation of smoke from the cigarette caused marked increase in airway resistance (fig 7). The increase in airway resistance appeared related to the volume of smoke plus air that was inhaled but was not related to changes in venous blood nicotine levels.

Discussion

While allowing our subjects to smoke in their usual way, we set out to vary the amount of smoke each could potentially inhale by asking them to smoke in rapid succession increasing numbers of identical cigarettes. We also varied the tar/nicotine yield of smoke inhaled by asking them to smoke different types of cigarettes. The acute changes in airway function, caused by smoking, determined by either MEFVC analysis or airway resistance measurements were small.
and appeared only to occur in certain individuals. Indeed, only three subjects (1, 6, and 10) consistently showed responses to smoking. No clear relationship could be found between the amount or type of smoke inhaled and size of the occasional acute airway response. Furthermore, the amount of smoke inhaled largely appeared independent of the number and type of cigarettes smoked. These observations suggest that smokers vary in the way they inhale cigarette smoke and that factors such as number and type of cigarette have only a minor influence upon their usual smoking response. Despite this, certain subjects appeared regularly to show evidence of acute narrowing after smoking which suggests an individual susceptibility.

The question as to how most habitual smokers avoid the irritant effects of cigarette smoke may be answered by our comparison of the effects of different patterns of smoking upon the acute airway response. It would appear that when a direct inhalation of smoke from the cigarette into the chest is achieved, there is consistent evidence of airway narrowing. A response was not consistently seen in either mouth or normal smoking and did not appear to be related to the volume of smoke inhaled when normal and direct smoking were contrasted. This suggests that the usual pattern of smoking, which we found to be similar to that observed by Rawbone et al. and consisted of an initial “drag” of smoke into the mouth followed after a variable pause by a subsequent inhalation of smoke into the lungs, could minimize the irritant qualities of the tobacco smoke. The way of inhaling smoke, in particular the interval that smoke remains in the mouth, has an important role in determining the relative concentrations of the constituents reaching the lungs. Buccal absorption of water soluble compounds together with precipitation of particulate matter (tar) leads to a relative increase of volatile insoluble compounds in the smoke entering the lungs during normal smoking. Direct inhalation of cigarette smoke prevents these adjustments occurring, enabling the full concentration of soluble gases such as sulphur dioxide and acrolein together with tar to reach the airways and so perhaps stimulate laryngeal and bronchial irritant receptors involved in the acute airway response. It is noteworthy in this respect that subject 10 who repeatedly responded to normal smoking was unable to inhale smoke directly and failed to directly draw any smoke into his chest, whereas smoking normally he was able to inhale smoke and tolerate the airway response to it.

The findings that smokers differ in their response to smoking cigarettes and that in only a small proportion was acute airway narrowing observed despite varying the number and type of cigarettes smoked, would appear at first sight discordant with recent work showing responses in most subjects. However, they are consistent with results of other studies where much larger numbers of smokers have been studied and where a similar degree of care has been taken to assess the causes, other than smoking, of variation in the measurements of airway function. The randomised and balanced nature of the design of our studies, in which each subject underwent in random order each type of treatment (number of cigarettes smoked, type of cigarette smoked, and type of smoking pattern) enabled the significance of interactions between those main factors influencing variation of measurements to be tested using analysis of variance. The results showed that some smokers consistently behaved differently from the group findings.

Internal consistency of the results was also achieved in that the same intersubject variation in response was seen with two different measurements of airway function, MEFVC and airway resistance. Furthermore, the objectivity of these measurements was maintained as the observers were unaware of the type of treatment each subject underwent. The assessment of “dose” of smoke inhaled into the lungs by following changes in venous blood nicotine levels in the studies using identical cigarettes is based on previous work by Russell and Haines. Our results confirm reports that mouth smoking—that is simply indrawing smoke into the mouth without a subsequent inhalation—fails to alter blood nicotine levels. This occurs because acid smoke from flue-cured tobacco in English cigarettes prevents buccal absorption of nicotine. It was not possible to use the same assessment when testing the smoking of different types of cigarette as they varied in their nicotine yield. For this reason, changes in venous carboxyhaemoglobin were used to assess the amount of smoke inhaled, as carbon monoxide was absorbed predominantly by the lungs and not buccal mucosa. This was based on the assumption that CO deliveries of different cigarettes were roughly comparable, as CO delivery depends on air permeability of the paper which affects both smoke dilution and tobacco burning. The failure to show a relationship between the amount of smoke inhaled and airway response supports our notion that composition of the inhaled smoke is the most important determinant of airway response rather than the amount or type of smoke. 
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inhaled. The similarity of the volumes of smoke inhaled during normal and direct smoke inhalation, which produced a markedly different response, also provides further support for this view.

We therefore conclude that there is a large intersubject variation in bronchial response to smoking which cannot be accounted for solely by the amount or type of smoke inhaled into the mouth.

Most habitual smokers appear able to adopt a pattern of smoking which avoids or minimises the irritant nature of cigarette smoke. Despite such manoeuvres, certain individuals still develop an acute airway response after smoking which may reflect an enhanced bronchial reactivity to cigarette smoke, and they may pursue a more rapid age-related decline in FEV₁.

The pattern of smoke inhalation is important in determining the relative concentration of the constituents of cigarette smoke which reach the lungs. This is the first demonstration that the pattern of smoke inhalation is also important in the physiological response to smoking and certainly more important than the number of cigarettes smoked, or the tar content of the smoke. This observation raises the possibility that the pattern of smoking may also play a major contributory role in the development of bronchial disease associated with smoking. Future studies of the pathogenesis of smoke-related diseases should perhaps include some assessment of the manner of smoke inhalation.

We wish to thank the Research Department of the Imperial Group Bristol for their support and assistance with this study.

References
