Effect of bronchodilators on the cough response to inhaled citric acid in normal and asthmatic subjects

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ABSTRACT  Coughing was induced in seven normal and eight asthmatic subjects by giving successive inhalations of citric acid aerosols of progressively higher concentration (range 0.5–32%). A baseline cough response was obtained on each of four experimental days, and there was no significant difference between days in this respect. Then the subjects received by inhalation either a bronchodilator (salbutamol 5 mg or ipratropium 1 mg) or placebo, in a paired double blind crossover design. A second citric acid run followed and was used for paired drug-placebo comparisons. In the asthmatic subjects the cough response was diminished by both bronchodilators (p < 0.05), and the cough threshold was significantly higher after ipratropium but not salbutamol. In normal subjects no significant effects were found. Airways calibre was assessed, by an oscillatory technique that measures the resistance of the respiratory system (Siemens Siregnost FD 5), in four of the seven normal and all eight asthmatic subjects. The mean respiratory resistance was higher in asthmatic than in normal subjects, and fell significantly after both bronchodilators. In normal subjects smaller decreases in respiratory resistance occurred, significant only with salbutamol. The simplest hypothesis which explains the results relates change in cough response to altered neuroreceptor sensitivity associated with rapid changes in bronchial calibre.

Cough is frequently the only presenting symptom in patients with bronchial asthma. The cough reflex arises from rapidly adapting receptors located in the larynx, trachea, and major bronchi. Impulses travel in afferent fibres in the vagus nerve to the cough centre in the brainstem. The tracheobronchial cough receptors are stimulated by touch and inhaled irritants and sensitised by bronchoconstriction, and this could explain why bronchoconstriction is associated with cough. Cough, in patients with uncomplicated bronchial asthma, is relieved by conventional aerosol bronchodilators. We have investigated cough induced by inhalation of citric acid in a group of normal and asthmatic subjects, and have assessed the effect of a β2 stimulant (salbutamol) and an anticholinergic (ipratropium) on the cough response.

Methods

SUBJECTS

Eight normal and eight asthmatic subjects were selected. All were non-smokers and had not had a respiratory infection in the eight weeks preceding the study. One of the normal subjects developed a respiratory infection during the study. Results for the remaining seven are presented. The normal subjects had no history of atopy. The asthmatic subjects had all been inpatients with an acute attack of bronchial asthma during the two years before the study. They were stable and had been out of hospital for at least six months. All were being treated with inhaled salbutamol only, and all gave a history of atopy. Medication was omitted for 12 hours before the experiments.

ASSESSMENT OF COUGH RESPONSE TO INHALED CITRIC ACID

The subjects were asked to breathe through a mouthpiece and a Fleisch pneumotachograph. Nebulised citric acid solutions were either vented to atmosphere or directed to the mouthpiece via a side tube during a selected inspiration, according to the setting of an electrically operated valve. To record cough a mercury strain gauge was loosely attached round the subject’s neck so that movement of the thyroid cartilage could be detected. The strain gauge was balanced with a Wheatstone bridge circuit and its sensitivity adjusted so that carotid pulsation could be recorded.
Airway flow, coughing movements, and valve opening were recorded on a Medelec fibreoptic recorder.

Each subject was positioned so that he could see a stop clock marked in seconds and had been trained to inhale from residual volume (RV) to total lung capacity (TLC) in five seconds. After breathing normally for about 20 seconds the subjects exhaled to RV and then inhaled to TLC over a five second period. During this inspiration the valve was opened to deliver the aerosol of citric acid to the subject. After inhalation the subjects exhaled to functional residual capacity and then continued to breathe normally through the mouthpiece for a further 30 seconds, unless coughing occurred.

The citric acid was prepared as solutions at concentrations of 0.5%, 1%, 2%, 4%, 8%, 16%, and 32%, and delivered as an aerosol from a Wright's nebuliser driven at 10 l.min⁻¹. Each dose was given as a single inhalation (as described above); the starting concentration was 0.5%, and the dose was increased at four minute intervals to the maximum of 32%.

After each inhalation we counted the number of coughs induced and measured the time between the start of inhalation and the first cough (the latency). We divided the number of coughs by the latency to obtain a further assessment of the cough response, which we called the "cough index." The concentration of citric acid at which coughing first occurred was also recorded—the "cough threshold." A value of 32% was recorded as the cough threshold if coughing was not induced at all during a run.

ASSESSMENT OF AIRWAY CALibre
In the eight asthmatic subjects and four of the seven normal subjects we measured the resistance of the respiratory system (Ros) by an oscillatory technique (Siemens Siregnost SD 5) during normal tidal breathing. Ros varies with tidal breathing, being lowest at end inspiration. We drew a horizontal line of best fit, by eye, through the minimum values of five to eight breaths displayed on a pen recording.

PROTOCOLS
Each subject attended on four days, at the same time of day. First, a baseline series of citric acid inhalations was given. This was used only to check that baseline conditions were not significantly different in this respect on the four experimental days (see discussion).

Each subject then received inhaled salbutamol (5 mg in 3 ml normal saline) paired on another day with placebo (3 ml normal saline) or inhaled ipratropium (1 mg in 3 ml normal saline), again paired on another day with placebo; thus there were four experiments in all. The order of drugs and placebo was randomised in each pair, and the order of the salbutamol pair in respect to the ipratropium pair was also randomised. Neither subject nor investigator knew whether an active preparation or placebo was being given, although a few subjects commented on the taste of ipratropium. For each subject all four experiments were completed within 10 days at the most.

All drug and placebo solutions were nebulised by a raindrop nebuliser (Ideal, Bennett) driven by a Wright's pump at 12 l.min⁻¹.

Thirty minutes after inhalation of the drug or placebo aerosol the series of citric acid inhalations was repeated. The results from this second series were used to test for differences in cough response between drug and paired placebo days.

Airways calibre was assessed as Ros immediately before and after the baseline series of citric inhalations (that is, once to assess baseline conditions and a second time to assess any effect of citric acid), and a third time just before the second series of citric acid inhalations to test for any drug effect.

STATISTICS
In assessing changes in standard measurements (for example, Ros) we used the paired t test. When dealing with non-linear or scaled variables (for example, number of coughs, cough index) we used non-parametric methods (Wilcoxon's signed rank test, Friedman's test).

Results

BASELINE SERIES OF CITRIC ACID INHALATIONS
For both normal and asthmatic subjects we summed the total number of coughs and the total of cough indices for each run. There were no significant differences between the four experimental days (Friedman's test; table 1). We repeated the analysis for the number of coughs, looked separately at results for 8%, 16%, and 32% citric acid, and tested for differences between normal and asthmatic subjects. No significant differences were found at any citric acid concentration on any experimental day. Asthematics did not, in general, cough more than normal subjects in response to citric acid.

The baseline cough threshold did not vary by more than two dosage increments over the four days in any subject.

DRUG EFFECTS ON COUGH RESPONSE
Too few coughs occurred at citric acid concentrations of less than 8% to allow any statistical analysis. Figure 1 shows the mean number of coughs at 8%, 16%, and 32% for normal and asthmatic subjects during drug and placebo runs. A possibly significant
drug effect is seen at all three concentrations in asthmatics, and at 32% in normal subjects. To test this we used paired results (fig 2) and Wilcoxon's signed rank test. On a few occasions, even at the high citric acid dose concentrations, subjects did not cough either with placebo or with paired active drug. These results are omitted from figure 2. In normal subjects the coughs induced were not significantly different when placebo and paired active drug were compared (fig 2A). In asthmatic subjects the number of coughs were, with one exception, less after salbutamol or ipratropium (fig 2B; p < 0.05 in both cases) at both 16% and 32% citric acid doses.

We repeated the analysis using cough index instead of number of coughs, with identical statistical conclusions.

The cough threshold (fig 3) in asthmatic subjects tended to be higher for citric acid runs after active drug than after placebo. This was just significant for ipratropium (p < 0.05) but not so for salbutamol. There was no significant effect in normal subjects.

Finally, we repeated the analysis, comparing the effect of active drug with the placebo, with which it was not originally paired. For ipratropium there was a significant fall in number of coughs and cough indices with 16% and 32% citric acid and a significant rise in threshold, as before (p < 0.05 in all cases). For salbutamol the cough index fell significantly with 16% and 32% citric acid, and the number of coughs with 16% also fell; but the number of coughs with 32% citric acid and the cough threshold were unchanged. Again, no significant effects were found in normal subjects.

**Table 1**  
Total coughs and cough indices (means with standard errors in parentheses) in all subjects on the four days of the experiment in the baseline citric acid runs

<table>
<thead>
<tr>
<th></th>
<th>Placebo</th>
<th>Salbutamol</th>
<th>Placebo</th>
<th>Ipratropium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal subjects (n = 7)</td>
<td></td>
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</tr>
<tr>
<td>Coughs</td>
<td>11.60 (3.8)</td>
<td>10.57 (2.6)</td>
<td>14.7 (7.6)</td>
<td>12.90 (5.3)</td>
</tr>
<tr>
<td>Cough indices</td>
<td>5.25 (2.6)</td>
<td>4.83 (2.0)</td>
<td>7.9 (2.0)</td>
<td>5.84 (2.3)</td>
</tr>
<tr>
<td>Asthmatic subjects (n = 8)</td>
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<td></td>
</tr>
<tr>
<td>Coughs</td>
<td>14.29 (3.3)</td>
<td>12.30 (4.3)</td>
<td>11.0 (4.2)</td>
<td>13.40 (4.0)</td>
</tr>
<tr>
<td>Cough indices</td>
<td>9.69 (3.6)</td>
<td>11.60 (5.3)</td>
<td>8.70 (4.4)</td>
<td>7.91 (3.1)</td>
</tr>
</tbody>
</table>

**Discussion**

Citrac acid induces cough in 90% of human subjects. The response is rapid and self limiting. The exact mechanism by which cough is stimulated is not known but the high osmolarity and the acidity of
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Fig 2  Individual results for number of coughs after active drugs salbutamol (S) and ipratropium (I), paired with placebo (P), at 16% and 32% citric acid concentrations.

Fig 3  Cough threshold in asthmatic subjects: the results obtained after administration of salbutamol and of ipratropium are connected to those for the results for the paired placebo.
Table 2  Airways impedance (Ros) (mbar.1⁻¹.s, means with standard errors in parentheses) at baseline (B), after baseline citric acid run (CA), and after drug (AD)

<table>
<thead>
<tr>
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<th>Placebo</th>
<th>Salbutamol</th>
<th>Ipratropium</th>
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<tbody>
<tr>
<td></td>
<td>Paired with salbutamol</td>
<td>Paired with ipratropium</td>
<td></td>
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<tr>
<td>Normal subjects</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n = 4)</td>
<td>B</td>
<td>1.86 (0.17)</td>
<td>2.16 (0.30)</td>
</tr>
<tr>
<td></td>
<td>CA</td>
<td>1.85 (0.12)</td>
<td>1.95 (0.22)</td>
</tr>
<tr>
<td></td>
<td>AD</td>
<td>2.14 (0.28)</td>
<td>1.30 (0.24)*</td>
</tr>
<tr>
<td>Asthmatic subjects</td>
<td>B</td>
<td>3.65 (0.80)</td>
<td>3.24 (0.62)</td>
</tr>
<tr>
<td>(n = 8)</td>
<td>CA</td>
<td>3.29 (0.87)</td>
<td>3.15 (0.46)</td>
</tr>
<tr>
<td></td>
<td>AD</td>
<td>2.23 (0.36)</td>
<td>1.92 (0.16)*</td>
</tr>
</tbody>
</table>

*p < 0.05 for the comparison between B and AD.

the nebulised solution may be partially responsible. In a study of this kind there are several technical problems which need consideration.

Since there is a diurnal variation in cough response, tests in an individual subject should be conducted at the same time of day. Since the response adapts over a short time period, comparisons for drug effect should be made on different days between drug and placebo, not immediately before and soon after drug administration.

Assessment of the magnitude of the response is difficult, since it seems very unlikely that a response of, say, three coughs, represents a receptor response of three times the magnitude of the neural output producing one cough. We have therefore used non-parametric statistics, and our results refer to directional change only—that is, we may say that subjects coughed significantly more, or less; but we cannot say how much more or less. This, however, is sufficient to detect the presence or absence of a drug effect, which is the purpose of this study.

There is considerable variability in the response (fig 2, table 1). We approached the problem by careful use of separate experiments pairing placebo with each drug, by testing separately for comparability of baseline conditions on each day, and by a double blind design. (Proper double blind trials cannot be done with inhaled ipratropium because of its distinctive taste.) Despite this variability, results for asthmatic subjects were clearcut (fig 2). The possibility of a type 2 statistical error certainly exists so far as negative conclusions are concerned. For example, although we did not show a significant effect in normal subjects, we may have missed one owing to high variability and small numbers.

So far as possible, constant volumes of inhaled citric acid aerosol were given by a standardised inhalation routine. Some subjects coughed before receiving the full dose. We tried to compensate for this by weighting the response according to the shorter latency. The "cough index" thus obtained gave similar results and identical conclusions.

We assessed airways resistance with a forced oscillation technique. This requires normal breathing through a mouthpiece and a reference impedance of fixed dimensions. An oscillatory flow of air (0.7 ml at 10 Hz) is superimposed on the resired air at the mouthpiece. Measurements of respiratory resistance thus obtained (Ros) correlate well with the results of conventional methods of measuring airways resistance using a body plethysmograph. Ros varies slightly during the respiratory cycle, and is also affected by involuntary movements of the upper airway structures (for example, glottic closure), and we have therefore used the lower values of Ros recorded during the respiratory cycle, which are less affected by artefact.

Cough arises from stimulation of rapidly adapting airway receptors, which are also stimulated and sensitised by bronchoconstriction. The higher airway tone in asthmatics might imply that the receptors have a higher resting discharge rate than in normal subjects and thus the stimulus required to induce cough would be less. Alternatively, the receptors might be more sensitive to the stimulus of an inhaled irritant. We were surprised that before treatment with bronchodilators there was no clear difference between the sensitivity of normal and asthmatic subjects to citric acid. Recent work does suggest that, in the absence of upper respiratory tract infection, the cough response is not necessarily related to the degree of bronchoconstriction. None of our asthmatic subjects had severe airways obstruction, but Ros was higher than in the normal subjects. The cough response appears to be not purely related to the degree of bronchoconstriction, when this is not rapidly changing. This is consistent with the clinical observation that cough in asthmatic patients is not related to the degree of airways obstruction.

Although the initial cough response to citric acid was similar we showed that the normal and asthmatic subjects reacted differently to citric acid after they had received bronchodilators. The bronchodilators had no significant effect in the normal
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subjects but they reduced cough number and cough index in the asthmatic patients, in whom ipratropium also increased the cough threshold.

As ipratropium and salbutamol both diminished the cough response, and both reduce airway tone by different mechanisms, the relatively rapid change in airway tone is likely to have contributed to our results. The change in Ros was as expected, greater in the asthmatic patients than in the normal subjects. A change in tone could reset irritant receptors and alter the threshold of airways smooth muscle receptors, which are more slowly adapting than irritant receptors. Discharge from smooth muscle receptors is known to alter with an increase in bronchial tone and we assume that a reverse effect would also occur.

The simplest explanation for our results is therefore that the cough response depends less on the static level of airways resistance than on sudden changes in its value, and that pulmonary neuroreceptors adapt to any constant level of bronchoconstriction but are then potentiated or inhibited by sudden increases or decreases in airway calibre. In asthmatic subjects a relatively large change in airway calibre after bronchodilators produced a demonstrable effect on cough response. In normal subjects a smaller change in calibre is to be expected (and was found), either too small to change the cough response or producing a change too small to be detected by our methods.

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