No demonstrable effect of S-carboxymethylcysteine on clearance of secretions from the human lung

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Thomson, M. L., Pavia, D., Jones, C. J., and McQuiston, T. A. C. (1975). Thorax, 30, 669–673. No demonstrable effect of S-carboxymethylcysteine on clearance of secretions from the human lung. The mucolytic efficacy of S-carboxymethylcysteine has been assessed in a double-blind crossover trial in 16 patients with chronic obstructive bronchitis. No significant difference was found between drug and placebo after four or seven days' treatment in the rate of clearance of secretions from the lung. This was measured by external counting of previously inhaled polystyrene tracer particles tagged with technetium-99m (⁹⁹ᵐTc). Lateral scans across the right chest after inhaling the aerosol showed equal penetration of particles towards the periphery of the lung in drug and placebo runs; this indicated that the airways had not been cleared of mucus by the drug. There was no significant difference between drug and placebo runs in the number of coughs or the weight and radioactive content of sputum voided or raised at the end of the run by chest percussion and postural drainage. Ventilatory capacity was not significantly changed nor was there any subjective improvement in the patients as a result of taking the drug.

S-carboxymethylcysteine (L-3-[(carboxymethyl)thio]-alanine; SCMC) is a derivative of the amino acid cysteine in which the sulphhydryl (SH) group has been blocked by a carboxylic acid residue. Despite this it is said to possess with acetyl cysteine (Sheffner et al., 1964) and other compounds with free sulphhydryl groups the ability to split the disulphide bonds of the long chain glycoproteins of mucus and thereby reduce its viscosity; it may also reduce the density of the fibrillar structure of the mucus by replacing fucosucin by sialomucins (Havez et al., 1970). Recently introduced into the United Kingdom, SCMC has been promoted as an effective respiratory mucolytic in chronic obstructive bronchitis when administered orally.

Although few double-blind trials have been done, most authors have reported reduced sputum viscosity by Brookfield viscometer (Autran, Bowtin, and Kleibasur, 1970; Edwards, Steel, and Leszczynski, 1970) or increased pourability (Aylward, 1973) in chronic obstructive bronchitis during both relapse and remission after taking the drug (2-0–2.5 g/day). There is less agreement about the effect on sputum production (Edwards et al., 1970; Lemercier et al., 1970; Aylward, 1973; Aylward, 1974) or ventilatory capacity (Lemercier et al., 1970; Aylward, 1973; Aylward, 1974) but most authors have noted subjective and objective clinical improvement in a variable proportion of their subjects. Patients with all types of sputum responded to the drug but those with mucoid or mucopurulent sputum were more affected in one study (Autran et al., 1970) while in another a change from purulent to mucoid sputum during treatment was reported (Sadoul, 1970).

We report below the effect of S-carboxymethylcysteine on mucociliary clearance in chronic obstructive bronchitis in a double-blind crossover trial using a new objective method. The procedure and radioactive dose (maximum 80 µCi) have been approved by the Isotopes Panel of the UK Medical Research Council, the Hospital for Sick Children (Great Ormond Street), the ethical subcommittee of this School, and by the United Kingdom Atomic Energy Authority Research Establishment, Harwell for participation by their own staff (Booker et al., 1967).

**PATIENTS AND METHODS**

**PATIENTS** Sixteen male patients with chronic obstructive bronchitis (Medical Research Council, 1965) took part in the study; four had chronic bronchitis
with varying degrees of bronchospasm and eight had chronic bronchitis with radiological evidence of emphysema. Although the patients were not chosen specifically because of difficulty in raising phlegm, we have shown that patients similar to these are capable of responding to expectorant and mucolytic drugs (Thomson, Pavia, and McNicol, 1973; Thomson et al., 1974). Informed consent was obtained from each patient. All were in remission throughout the trial period. Their mean (± SD) age was 63.6 ± 8.9 years and height 1.71 ± 0.07 metres. There were nine smokers and seven ex-smokers who had smoked on average 15 270 pack-years. They had moderate to severe loss of ventilatory capacity as assessed by a dry bellows spirometer (Vitalograph): after a week without mucolytic drugs their mean (± SD) forced vital capacity (FVC), forced expiratory volume in 1 sec (FEV1), and FEV1/FVC were respectively 2.42 ± 0.75 l. (62.0 ± 17.3 %), 1.09 ± 0.38 l. (37.3 ± 12.7 %), and 46.2 ± 11.2 l. (68.2 ± 16.9 %). (The values in parentheses are percentages of predicted (Cotes, 1968).)

EXPERIMENTAL DESIGN Mucolytic and expectorant drugs were discontinued one week before the trial but other therapy was maintained throughout the trial period. In randomized double-blind order, four of the patients took 4 g per day in four doses of SCMC in syrup (Mucodyne) and four took the same quantity of syrup only, a placebo indistinguishable from the drug. The first trial run was done after seven days of these treatments. After a further seven days with placebo only, the drug/placebo routine was repeated in reverse order and a second identical trial run was done after seven days of treatment. The same procedure was followed throughout for the remaining eight patients save that treatment was cut from seven to four days by increasing the rest period from seven to 10 days. In addition, at the end of both trial runs, this latter group had chest percussion and postural drainage done by a qualified physiotherapist. Placebo was taken throughout all non-treatment periods and a last dose of drug or placebo was given in the morning before both trial runs. Since no significant differences were found between the four- and seven-day treatment groups, the findings from these were pooled before the final analysis of the results.

A fluorescent marker (fluorescein) (Editorial, Lancet, 1972), incorporated in both drug and placebo, was identified in the urine of all subjects on the days of the trial runs. Apart from the two baseline ventilatory capacity determinations all measurements were made during the day of the trial runs.

TRACER PARTICLES The method has been fully described (Thomson et al., 1974). The patients inhaled monodispersed 5(±0.7 SD) μm particles of polystyrene which had been tagged unencapsulated with 99mTc (Few, Short, and Thomson, 1970). The volume of aerosol inhaled at each breath was automatically cut off at 500 ml. The flow rate during inhalation (mean average 21.2 ± 1.7 SD 1/min) was (fortuitously) identical in drug and placebo runs. These factors, especially the first, should be strictly controlled since they affect the depth of penetration of the particles (Booker et al., 1967) and therefore their subsequent rate of clearance from the lung. After inhalation, whole lung gamma counts were made at half-hourly intervals to 6 hours by NaI(Tl) scintillation detectors: one detector was at the midpoint of the sternum and a second axially opposite in the seated subjects.

Initial depth of deposition of the particles was recorded by rectilinear gamma scanning (Dawson et al., 1971) across the right lung at one-inch intervals after inhaling the tracer particles.

RESULTS WHOLE LUNG CLEARANCE Figure 1 shows for drug and placebo runs the mean serial whole lung gamma counts at half-hourly intervals as percentages of the

![FIG. 1. Mean clearance curves from the whole lung counts for 16 patients in placebo and drug runs. Also shown are standard errors (vertical bars ± 1 SE), the paired t values, and probabilities at half-hourly intervals. (Right) the effect of physiotherapy in 8 patients.](http://thorax.bmj.com/)
drug and placebo runs were not significant. The mean losses during physiotherapy expressed as percentages of the six-hour values (19.9 and 19.4%) were almost the same in placebo and drug runs.

LUNG SCANS  Figure 2 shows the initial depth of penetration of particles across the right lung for drug and placebo runs. For each subject the counts for each 1 in (2.5 cm) traverse have been expressed as percentages of the total count for all traverses: the heights of the columns are the means of these percentages for 16 patients. A rough index of penetration of the particles is obtained from the ratio of the sum of columns 1 and 2 (near the sternum) to the sum of columns 4 and 5 (near the periphery). There was no significant difference between drug and placebo runs in this penetration index (t=0.52; P not significant) nor in the sums of the control (t=0.67) or peripheral (t=1.00) pairs of columns.

VENTILATORY FUNCTION  On average the FEV₁ fell by 10 and 30 ml and the FVC rose by 60 and 50 ml over the placebo and drug periods respectively. These differences were not statistically significant.

COUGH AND SPUTUM  The Table shows for the drug and placebo runs the mean number of coughs and sputum samples and the mean weight and radioactive content of sputum voided during the runs. The difference between means was not significant nor was there any consistent pattern in these variables indicating any preponderance in drug or placebo runs. During placebo runs sputa were classified as 10 mucoid, one mucopurulent, and five purulent: in the drug runs four were more purulent, three were less purulent, and nine were unchanged.

Physiotherapy on eight patients caused mean losses of sputum of 2.6 and 2.1 g with 7400 and 16300 gamma counts for placebo and drug runs respectively; these differences did not differ significantly (t=0.58 and 0.43).

The answers to a questionnaire about dyspnoea, the quantity and stickiness of phlegm, and the ease with which it was raised were equivocal.

DISCUSSION  Although SCMC reduces sputum viscosity moderately in vitro (Hirsch, Viernes, and Kory, 1970) we have been unable to demonstrate that SCMC affects the fast phase rate of clearance of secretions from the lung in patients with chronic obstructive bronchitis.

The topographical distribution of inhaled tracer particles as obtained by rectilinear scanning was necessary for validating the whole lung clearance findings because the rate of clearance of particles varies inversely with the depth in the lung at which the particles are deposited (Thomson and Pavia, 1974). Peripheral mucus and the entrained particles have a longer transfer path to the larynx and, more important in dogs (Asmundsson and Kilburn, 1970) and almost certainly in humans, the inherent rate of transfer is much slower in the finer, than in the larger airways (Hilding, 1963). Moreover coughing is much less effective in expelling mucus from the small airways (<2 mm diameter) because adequate linear gas velocities are not achieved in the expanded peripheral airway bed (Leith, 1968). The close similarity in particle penetration shown here by the scans in drug and placebo runs indicates that the race of the particles to the larynx was fair.

The failure of the tracer particles to penetrate deeper into the lung after four or seven days of treatment is retrospective evidence that the drug did

<p>| TABLE |
| --- | --- | --- | --- | --- |
| Mean (± SE) number of coughs and sputum samples and the sputum weight and radioactivity for 16 patients for drug and placebo trial runs together with the paired t values and probabilities |</p>
<table>
<thead>
<tr>
<th>No. of Coughs</th>
<th>No. of Sputum Samples</th>
<th>Sputum Weight (g)</th>
<th>Sputum Radioactive Content (counts) x 10³</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>D</td>
<td>P</td>
<td>D</td>
</tr>
<tr>
<td>Mean</td>
<td>53.7</td>
<td>59.7</td>
<td>5.1</td>
</tr>
<tr>
<td>SE</td>
<td>13.2</td>
<td>26.0</td>
<td>1.0</td>
</tr>
<tr>
<td>t</td>
<td>1.46</td>
<td>0.08</td>
<td>0.93</td>
</tr>
<tr>
<td>Probability</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
</tbody>
</table>

P = Placebo; D = Drug.
not clear the airways before or between the trial runs. In a previous trial (Thomson et al., 1974) increased particle penetration was found after treatment with another mucolytic.

**Sputum Viscosity** In the study reported here there is no evidence that those patients with mucoid sputum showed more mucolysis than those with purulent sputum. Purulent sputum which is mainly composed of DNA fibres would not be expected to respond to double-bond-splitting agents such as SCMC.

It has long been known that there is a minimum mucus viscosity below which the cilia beat ineffectually, for example, in pandemic influenza (Hilding, 1943). Mucolytic agents might defeat their aim by reducing viscosity below this minimum (Carson, Goldhamer, and Carpenter, 1966): the dose used here (4 g/day) was higher than that recommended (2–0–2.5 g/day). Had the ciliary mechanism failed for this reason in the present trial, however, more low viscosity sputum would have been produced by physiotherapy in the drug than in the placebo runs, but this was not observed.

Sputum was collected here only over the six hours of the trial runs. During this time, however, the subjects were continuously under observation and every effort was made to ensure that all sputum raised was collected. Charman and Reid (1972) found no difference between early morning sputum and that obtained later in the day by physiotherapy, and it is unlikely that 24-hour collections would have shown major differences between drug and placebo volumes. Although increase in sputum production is commonly associated with decrease in viscosity, it is unlikely *per se* to result in better lung function (Lopez-Vidriero et al., 1973).

**Effect of Physiotherapy** Chest percussion and postural drainage caused losses of activity from the lung which were greater on the average than those observed during the final three hours of the trial run. It would seem that during that three-hour period retention or ‘pooling’ of secretions must have occurred in the large airways in some subjects. If this is true, coughing must have been suppressed or ineffective, and in addition mucociliary transport must have been inadequate to overcome gravity. Presumably the postural change enabled gravity to be overcome or it stimulated coughing by altering the distribution of mucus. There was no evidence that this retention in the large airways was affected by the SCMC.

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**REFERENCES**


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