Airway constrictor response to cotton bract extracts in the absence of endotoxin

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ABSTRACT Crude and purified aqueous extracts of cotton bracts shown to cause airway constriction in naive subjects were assayed for endotoxin content. Pulmonary function measured by flow changes on partial expiratory flow volume curves was used to assess airway responses to the bract extracts after their inhalation by a panel of volunteers. These responses are similar to the acute responses experienced by cotton textile workers. Crude aqueous extracts from various bracts harvested before and after senescence of the cotton plant displayed endotoxin concentrations ranging widely from 0·086 to 50 μg/ml. No correlation was found between these differences in endotoxin content and the severity of the airway constrictor response. Purifying the bract extract by a series of procedures that included precipitation of polymeric material by addition of methanol and chromatography on an anion exchange column of DEAE-Sephacel resulted in the elimination of almost all the endotoxin. The partially purified extract contained less than 1 ng/ml of endotoxin. The panel of volunteers responded to this purified bract extract, however, with a decrease in pulmonary function which was more than 60% of that seen with the crude extract of bracts. It is concluded that aqueous extracts of cotton bract contain an agent(s) other than endotoxin that causes acute airway constriction in people.

Cotton dust and cotton bract contain endotoxin, the amount of which varies widely depending, at least in part, on cotton quality, growing conditions, and growing location.² Pernis et al suggested that byssinosis, the specific lung disease of cotton textile workers, may be due to endotoxins produced by the Gram negative bacteria normally found on cotton bracts.² The bacterial content of the cotton bales being processed in one experiment³ and airborne microorganisms in another⁴ correlated with the prevalence of byssinotic symptoms. In the latter study, however, airborne endotoxin did not correlate with symptoms. In recent studies Diem et al found that decreases in pulmonary function over the work shift correlated with airborne dust concentration, not with bacterial or endotoxin concentrations.⁵ By contrast, Olenchok et al found in an experimental card room environment that FEV₁ decreases correlated with endotoxin content in the air significantly better than with airborne dust levels,⁶ and Rylander et al also found a dose response relation between airborne endotoxin and decreases in FEV₁, although the latter cautioned that the endotoxin levels may be only an indicator of some other agent present in the dust with which it maintains a close relation.⁷

Aqueous extracts of cotton bract induce an acute airway constriction in healthy subjects never before exposed to cotton dust.⁸ This constriction is similar in its changes of lung function parameters and its time course to the acute response experienced by textile workers.⁹¹⁰ The response is present with both the crude aqueous extract and partially purified extracts. The agent causing the airway constriction has not yet been identified but several of its characteristics are known, including solubility in water (a characteristic of endotoxin) and a molecular weight of less than 1000 daltons¹¹ (which does not characterise endotoxin). To determine the possible involvement of endotoxin in the acute airway constrictor reaction to cotton dust, the endotoxin content of the crude and purified cotton bract extracts which induce airway constriction in people was determined.

Materials and methods

PREPARATION OF COTTON BRACT EXTRACTS

The flow chart in fig 1 illustrates our procedures for...
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Dried cotton bracts
6 ml H₂O/g
2h, 23°C
16 000 x g, 10 min
0.45 μm Filter
Precipitate discarded
Standard crude extract
Freeze dry for storage (dry wt 60 mg/ml)
Resuspend, 700 mg/5 ml H₂O
Add methanol, 35 ml
16 000 x g, 5 min
Precipitate discarded
Rotovap, 42°C
Resuspend, 4 ml H₂O
7:1 MeOH Extract (concentrated 4 x’s)
DEAE-Sepharcell column (10 x 1.5 cm)
equilibrated and eluted with H₂O (120 ml)
DEAE Extract
Freeze dry for storage (dry wt 15 mg/ml)
Resuspend, 15 mg/ml
Add equal volume of diethyl ether
Shake vigorously 3 min
Aqueous layer
Ether layer
Bubble with N₂ to remove ether
DEAE ether extract
Freeze dry for storage (dry wt 15 mg/ml)

Fig 1 Procedure for preparing cotton bract extracts.

the preparation of the various bract extracts. Unless otherwise indicated, bracts were handpicked from frost killed plants just before harvest from cotton fields in the Lubbock, Texas area; they were stored at room temperature and pulversised in a Waring blender immediately before extraction. Extracts were freeze dried and stored at −4°C. For inhalation challenge by our subjects each extract was reconstituted with water or saline, as indicated, at a concentration equivalent to the standard crude extract. This ensured that for challenge purposes components were not concentrated as purification progressed. Pyrogen free water was used throughout.

SUBJECTS
Healthy volunteer subjects (no respiratory symptoms, no history of asthma), aged 18–45, men, women, smokers, and non-smokers, were recruited from area universities. All were naive subjects never before exposed to cotton or other textile dust of any kind. For all procedures concerning volunteers informed consent was obtained according to the Yale University Human Investigating Committee guidelines.

ASSAY OF ACUTE AIRWAY CONSTRICTION
While breathing normally, each subject inhaled the aerosol of cotton bract extract for 10 minutes through a Dautrebande model D 30 aerosol generator operated at 20 psi air pressure. With these conditions the output of the nebuliser is constant, and the particle size delivered has been measured to be less than 1 μm in diameter. 13 The aerosol was inhaled through a mouthpiece from a J valve. The inspiratory port of the J valve was attached to a T piece, which in turn was connected to the aerosol generator and a 1 m long reservoir tube with internal diameter of 2.5 cm.

Constriction was assayed by comparing lung function values obtained from recordings of partial and maximum expiratory flow volume (PEFV, MEFV) curves 14 before and at 30 minute intervals for a two and a half hour to three hour period after the 10 minute inhalation of the aerosolised extract. These
Table 1  Endotoxin content and airway constrictor activity of cotton bract

<table>
<thead>
<tr>
<th>SCE from samples</th>
<th>Dry wt of SCE/ml</th>
<th>Endotoxin (µg/ml SCE)</th>
<th>MEF 40%(P) % decrease (Mean ± SEM)</th>
<th>Comparative MEF 40%(P) changes (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frost killed bracts 1979 harvest</td>
<td>60</td>
<td>36</td>
<td>36:0 ± 4:4 (n = 8)</td>
<td>100</td>
</tr>
<tr>
<td>Green bracts, air dried 1981 harvest (Texas)</td>
<td>41:2</td>
<td>0:25</td>
<td>12:8 ± 4:0 (n = 8)</td>
<td>35:6</td>
</tr>
<tr>
<td>Fresh green bracts, 1981 harvest</td>
<td>40:0</td>
<td>0:086</td>
<td>25:3 ± 5:1 (n = 8)</td>
<td>82:0</td>
</tr>
<tr>
<td>(New Orleans)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SCE = Standard crude extract.

were recorded with a pneumotachograph integrator device and an XY recorder; ordinate: expiratory flow rates, abscissa: expired volume. The subject first inspired to about 65% of the vital capacity (VC)—the exact level is not crucial and may vary from about 60% to 75% of VC—the subsequent forced expiration to residual volume yielded the PEFV curve. Without interruption, the subject next inspired maximally, and then again expired forcefully and maximally to residual volume, recording the MEFV curve. Forced vital capacity (FVC) and forced expiratory volume in one second (FEV₁) were measured from the MEFV curve. MEF 40%(P) is the instantaneous flow rate on the PEFV curve measured at the lung volumes corresponding to 40% of the forced vital capacity (maximum inspiration = 100% FVC). To compare responses to bract extract aerosol the MEF 40%(P) value was used where the 40% VC volume for each subject day was computed as an average from five control (before aerosol inhalation) MEFV curves of that day. This 40% volume was used throughout the test period.

Each subject was screened initially to determine responder status. For this, the crude aqueous extract (made from the frost killed bracts from the Lubbock, Texas, 1979 harvest) was used as the standard for screening all subjects. Responders to cotton bract were defined as subjects whose MEF 40%(P) decreased 20% or more after inhalation challenge with the standard extract. This value has been selected in previous studies because of its ability consistently to identify mild airway constrictor responses. Responders were selected to assay for constrictor activity in unknown extracts. Non-responders—that is subjects who showed a maximal decrease in MEF 40%(P) of 20% or less—were excluded.

Saline, administered in the same manner as the bract extract, was used as a control.

ENDOTOXIN DETERMINATION

The Limulus amoebocyte lysate test performed according to the Associates of Cape Cod, Inc (PO Box 224, Woods Hole, MA 02543) was used to determine endotoxin content of bract extracts. Serial dilutions of each bract extract were prepared with pyrogen free water. Equal aliquots (0·1 ml) of each dilution and lysate were mixed in tubes and incubated in a water bath at 37°C for one hour. The tubes were inverted and the last dilution to give a stable clot was compared with a similar clot formation in a dilution series of known amounts of commercial endotoxin (LPS E.cherichia coli serotype O 113, lot no 13, from Associates of Cape Cod, Inc). The water used to make the dilutions was used as a negative control.

Results

Table 1 shows the endotoxin content of crude extracts prepared from four different bract samples. Two samples, the frost killed 1979 harvest and the brown bracts 1981 harvest, which were picked after the plant had dried and turned brown, contained 36 and 50 µg/ml endotoxin respectively.

A third sample prepared from bracts picked from the same field as the brown bracts 1981 harvest, but picked while the plants were still alive and green and

Table 2  Airway constrictor activity of purified cotton bract extracts

<table>
<thead>
<tr>
<th>Sample</th>
<th>No*</th>
<th>% Airway constrictor activity† (Mean ± SEM)</th>
<th>Dry wt (mg/ml)</th>
<th>Endotoxin (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCE</td>
<td>100</td>
<td></td>
<td>60</td>
<td>36000</td>
</tr>
<tr>
<td>7:1 MeOH extract</td>
<td>22</td>
<td>61 ± 5</td>
<td>24</td>
<td>1·3</td>
</tr>
<tr>
<td>DEAE extract</td>
<td>16</td>
<td>58 ± 8</td>
<td>15</td>
<td>0·3</td>
</tr>
<tr>
<td>Ether extracted DEAE extract</td>
<td>7</td>
<td>63 ± 10</td>
<td>15</td>
<td>0·2</td>
</tr>
</tbody>
</table>

SCE = Standard crude extract.

*NNumber of subjects tested with sample.

†dMEF 40%(P)purified extract × 100.
then air dried, contained about 200 times less endotoxin (only 0.25 µg/ml). Even less endotoxin was found in the fourth sample of bracts which was picked while still green and immediately freeze dried to prevent senescence from occurring. The extracts from these bracts contained only 0.086 µg/ml endotoxin.

The ability of these extracts to induce airway constriction in people was tested in a panel of eight volunteers who were known to be responders. The results are included in table 1. Responder status was determined by screening each volunteer with the standard crude extract of bracts prepared from the frost killed 1979 harvest. Compared with the 1979 harvest, the 1981 brown bracts were only about half as active (48.6%) even though they contained slightly more endotoxin. The green bracts from the 1981 harvest exhibited an airway constricting activity of 35.6%, only somewhat less than the brown bracts, whereas the endotoxin content was less by several orders of magnitude. The extract from the fresh green bracts, while containing the least amount of endotoxin, was more active in inducing airway constriction than either the air dried green or the brown bracts from the 1981 harvest. Its percentage activity was 82, close to that of the frost killed 1979 bract.

The crude aqueous extract from the 1979 harvest was partially purified in a series of sequential steps, the first of which was the addition of methanol. This resulted in the precipitation of a large amount of material including presumably proteins and other large molecules. As shown in table 2, the supernatant solution from this step, the 7:1 MeOH extract had a dry weight of 24 mg/ml reduced from the original 60 mg/ml. The airway constricting activity of this preparation was 61% that of the original extract, and the endotoxin concentration was at the nanogram level (1.3 ng/ml) compared with 36 µg/ml in the original crude extract. The precipitate from the 7:1 MeOH extract thus contained most of the endotoxin and accounted for only a 16% loss of activity.

The 7:1 MeOH extract was purified further by passing it through a DEAE anion exchange column. The fraction not bound by the column, designated the DEAE extract, showed additional decreases in dry weight material to 15 mg/ml and endotoxin content to 0.3 ng/ml. These changes resulted in no further de-

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**Table 3: Airway constrictor activity and endotoxin content of DEAE fractions of bract extracts**

<table>
<thead>
<tr>
<th>Sample</th>
<th>No*</th>
<th>ΔMEF 40% (P) † (Mean ± SEM) (%)</th>
<th>% Airway constrictor activity ‡</th>
<th>Endotoxin (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCE</td>
<td>5</td>
<td>-34 ± 6</td>
<td>100</td>
<td>36000</td>
</tr>
<tr>
<td>DEAE, Fr no 1</td>
<td>3</td>
<td>+4 ± 4</td>
<td>0</td>
<td>0.5</td>
</tr>
<tr>
<td>DEAE, Fr no 2</td>
<td>5</td>
<td>-24 ± 6</td>
<td>71</td>
<td>0.3</td>
</tr>
<tr>
<td>DEAE, Fr no 3</td>
<td>3</td>
<td>+3 ± 2</td>
<td>0</td>
<td>0.4</td>
</tr>
</tbody>
</table>

*SCE = Standard crude extract.

*Number of subjects tested with sample.

†Change in MEF 40% (P) after exposure to sample.

‡ΔMEF 40% (P) purifled extract × 100.

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**Fig 2** Mean % fall in MEF 40% (P) after inhalation challenge versus endotoxin concentration (ng/ml) in various cotton bract extracts. Extracts compared were those listed in tables 1 and 2. In order of decreasing endotoxin content they are SCE of brown bracts (extreme right), SCE of frost killed 1979 bracts, SCE of green bracts, SCE of fresh green bracts, 7:1 MeOH extract, DEAE extract, and ether extracted DEAE extract (extreme left).
crease in airway constrictor activity (see table 2). Extracting the DEAE extract with diethyl ether caused little change in dry weight, endotoxin content, or constrictor activity but did remove lacinilene compounds.

The DEAE extract was fractionated into three fractions as it emerged from the column. Airway constrictor activity was associated exclusively with the second fraction despite the finding that all the fractions contained essentially the same low level (less than 1 ng/ml) of endotoxin. The results are presented in table 3. Results with the standard crude extract also are included for comparison. Subsequent assays performed in another laboratory confirmed these endotoxin values.¹⁷

To confirm that the purification scheme used for the bract extract does remove endotoxin, a solution of 180 ng/ml E coli endotoxin (from a commercial source) was carried through the scheme. With the addition of methanol no precipitation was observed. Nevertheless, after passage of the 180 ng/ml solution through the DEAE anion exchange column less than 0.2 ng/ml endotoxin remained in the solution. This confirms that endotoxin is consistently removed by the procedures and conditions used to purify our cotton bract extracts.

In order further to quantitate the relation between bract endotoxin content and airway constrictor response, the mean percentage fall in MEF 40%(P) (± 1 SD) was plotted (fig 2) against the endotoxin concentration of the various bract preparations used for each of the panels of subjects. As may be appreciated, over a several thousandfold variation in concentration, endotoxin did not influence the constrictor response.

Further, in table 4, the response between the bract preparations was compared in the same groups of individuals. The horizontal and vertical headings list the bract preparations and their endotoxin content. Numbers in the upper right of the diagram represent the average percentage difference in the MEF 40%(P) drop between the bract preparations (positive differences indicate that the preparations with the higher endotoxin content have a greater constrictor effect, negative differences indicate that the preparations with the lower endotoxin content have the greater constrictor effect). The second number in each comparison represents the number of paired comparisons.

The set of numbers in the lower left of the diagram indicates the t statistic for the paired comparison, the number of degrees of freedom (df), and the p value. Five of these comparisons were statistically significant. In these groups it is apparent that the standard crude extract is consistently more potent

Table 4 Differences in bronchoconstrictor responses (percentile in MEF 40%(P)) between bract preparations (Mean ± 1 SD)

<table>
<thead>
<tr>
<th>Sample*</th>
<th>Brown</th>
<th>SCE</th>
<th>Green</th>
<th>Fresh green</th>
<th>MEOH</th>
<th>DEAE</th>
<th>DEAE ether</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endotoxin (ng)</td>
<td>50 x 10³</td>
<td>36 x 10³</td>
<td>0.25 x 10³</td>
<td>0.086 x 10³</td>
<td>1.3</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>Brown</td>
<td>-19 ± 10.1</td>
<td>23 ± 18.4</td>
<td>9 ± 17.3</td>
<td>16 ± 13.5</td>
<td>19 ± 16.4</td>
<td>15 ± 9.4</td>
<td></td>
</tr>
<tr>
<td>50 x 10³</td>
<td>8</td>
<td>8</td>
<td>10</td>
<td>22</td>
<td>16</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>SCE</td>
<td>5-2</td>
<td>df = 7</td>
<td>t = 3-6</td>
<td>df = 7</td>
<td>p = 0.0045</td>
<td>1.7</td>
<td>df = 9</td>
</tr>
<tr>
<td>36 x 10³</td>
<td>23 ± 18.4</td>
<td>8</td>
<td>9 ± 17.3</td>
<td>10</td>
<td>16 ± 13.5</td>
<td>19 ± 16.4</td>
<td>15 ± 9.4</td>
</tr>
<tr>
<td>Green</td>
<td>0.25 x 10³</td>
<td>0.086 x 10³</td>
<td>1.3</td>
<td>0.3</td>
<td>0.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>t = 3-6</td>
<td>df = 7</td>
<td>p = 0.0015</td>
<td>df = 6</td>
<td>p = 0.0026</td>
<td>1-6</td>
<td>df = 7</td>
</tr>
<tr>
<td>Fresh green</td>
<td>0.086 x 10³</td>
<td>1.3</td>
<td>0.3</td>
<td>0.2</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SCE = Standard crude extract.
MEOH = 7-1 Methanol extract.
DEAE = DEAE extract.
DEAE ether = DEAE extract after ether extraction.
than the other preparations in terms of its constrictor effect. Clearly, however, no consistent pattern may be seen with regard to the endotoxin concentration.

Discussion
Naive subjects never before exposed to cotton dust of any type respond to inhalation of crude aqueous extracts of cotton dust or bract with an acute airway constriction that has a delay of 90–120 minutes before reaching maximum. No fever was found associated with this response even though the standard crude extract contains about 36 mg/ml of endotoxin. Airborne respirable dust in cotton textile mills contains Gram negative bacteria as well as endotoxin, whereas the crude extracts used in this study have been sterilized by passage through a 0.45 micron filter before being administered to subjects. The question arises, is the endotoxin concentration in mill dust or bract extracts high enough to cause (or contribute to) the acute airway constriction observed? Two groups have studied and reported on the response of subjects to inhalation of pure endotoxin. In the earlier study 40 µg or 80 µg of purified *E coli* endotoxin were inhaled. Two subjects who inhaled 40 µg had no response and of the eight subjects who inhaled 80 µg, two responded with FEV₁ decreases of 17–20% which were maximal at 30 minutes, the other six did not experience significant changes in FEV₁. In a recent study Rylander's laboratory challenged 15 subjects with 4–20 µg of pure endotoxin in solution. Even at the higher concentration which was inhaled by four of the 15 subjects, they found no significant decreases in FEV₁. It appears from these studies that only a limited number of healthy individuals are sensitive to endotoxin at concentrations as high as 80 µg of pure endotoxin.

Airborne respirable dust in cotton textile mills contains 0.02–3.16 µg/m³ endotoxin. If it is assumed a worker breathes about 15 m³ during an eight hour shift he will breathe in 0.3–47 µg endotoxin over the workday. In volunteers, where one assumes tidal breathing at a rate of 12 breaths per minute at 800 ml/breath for the 10 minute inhalation test period, it is estimated that each subject inhales 0.3 ml of bract extracts. This is equivalent to inhaling 11 µg of endotoxin when the standard crude extract of bracts 1979 harvest is tested and only 0.09 ng for the purified bract extracts (DEAE extracts). These endotoxin concentrations are far below those reported to cause airway constriction in people, and thus it is not surprising that no correlation was seen between endotoxin concentrations and bronchoconstrictor effects. As in these challenge studies with the bract extracts the pure endotoxin exposures were administered over a short period in contrast to the exposure of mill workers, who have a prolonged exposure and who are exposed at the same time to Gram negative bacteria as well.

Crude aqueous extracts prepared from different samples of cotton bracts showed a wide range of endotoxin content from 0.086–50 µg/ml. No correlation was found between the airway constrictor activity of these crude extract samples and their endotoxin content—again not surprising in view of the findings of Cavagna et al. and Haglind et al.

Purified extracts, which are essentially endotoxin free (<1 ng/ml endotoxin), retain the ability to induce airway constriction in volunteers. The less than 1 ng/ml concentration of endotoxin remaining in purified extracts is without effect. DEAE fractions collected before and after the active fraction (fraction 2) contain similar low levels of endotoxin and yet are without airway constricting activity. After the three sequential purification steps, about 63% of the constricting activity remains. Thus aqueous extracts of cotton bract contain an agent(s) other than endotoxin which causes acute airway constriction in people.

Since cotton mill dust contains a high proportion of bract material, the bract constrictor agent(s) found in aqueous extracts is undoubtedly present in mill dust as well. Attempts to compare the relative dose of bract aqueous extract and respirable mill dust are rough approximations at best. Estimates of the proportion of bract in mill dust vary (32–52%) and no data are available to know the proportion of the airway constricting agent(s) extracted into the initial aqueous extract, the standard crude extract. The volume of extract inhaled by volunteers is about 0.3 ml prepared from 50 mg of bract. The amount of respirable mill dust inhaled by workers is 5–57 mg, assuming dust levels of 0.46–3.80 mg/m³ and an inhalation of 15 m³ during an eight hour workshift. It has not been shown that the quantities of endotoxin present in textile mill dust are sufficient alone to account for the acute airway constriction observed in mill workers. Possibly, however, the bract constrictor agent and sublevels of endotoxin are present together, as they may be in mill dust, a synergistic action exists.

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References