Characteristics and management of chronic destructive pneumonia

E W J CAMERON, P C APPELBAM, D PUDIFIN, W S HUTTON, S A CHATTERTON, AND J DUURSMA

From the Departments of Thoracic Surgery, Microbiology, and Medicine, University of Natal, Durban, South Africa

ABSTRACT In 12 years 627 patients presented to Wentworth Hospital, Natal with chronic destructive pneumonia (CDP). Common symptoms were haemoptysis, the production of foul-smelling sputum, and chest pain. The disease pursued a chronic course with acute exacerbations which may be lethal. The majority of patients were African men aged between 20 and 50 years who were free from other significant disease apart from dental infection. Radiographically and pathologically CDP had the characteristics of a necrotising pneumonia, and microbiological investigation showed mixed aerobic and anaerobic flora in the lower respiratory tract. Gram-positive aerobic cocci and Bacteroides species were the predominant organisms. In 120 patients treatment regimens were based on chloramphenicol, in 429 cephalosporins, and in 78 on combination therapy with cephalosporins, penicillin, and metronidazole. One hundred and seventy patients also required operative management in an attempt to control progress of the disease. The overall inpatient mortality rate from CDP was 7.8%. In the group of patients treated with combination therapy the mortality rate was 1.3%.

Le Roux1 introduced the term “chronic destructive pneumonia” (CDP) when describing a pulmonary lesion which occurs in Natal, South Africa. The disease has the characteristics of a bacterial infection and its pathological hallmark is necrosis of pulmonary tissue.

Between 1966 and 1977, 627 patients with CDP were referred to the Thoracic Surgical Unit, Wentworth Hospital from medical clinics where the natural history of the disease had been modified by antimicrobial therapy. The common reason for referral of the patients was failure of such therapy to control the disease. In this paper we summarise the characteristics and management of CDP in this group of patients, of whom 75% were African men aged between 21 and 60 years.

Clinical features

The cardinal symptoms of CDP are haemoptysis, cough productive of foul-smelling sputum, and chest pain. Fifteen per cent of patients had a history shorter than one month. Their symptoms were acute in onset and life-threatening in severity. Forty-seven per cent of patients had a history of more than one acute episode of pulmonary infection treated at a medical clinic. Some patients had experienced up to five recurrences of acute illness with intervening periods of quiescence and wellbeing. However, many of these patients eventually developed symptoms refractory to medical therapy. Thirty-two per cent of patients were referred from tuberculosis clinics where failure to isolate mycobacteria and lack of response to chemotherapy prompted review of the original diagnosis of pulmonary tuberculosis. Six per cent of patients were thought to have bronchial carcinoma. The symptomatology in the last two groups of patients was chronic and undramatic. Significant other disease, most commonly grand mal epilepsy and diabetes mellitus, coexisted with CDP in 6% of patients.

The family and social histories of the patients showed the disease to be endemic throughout urban and rural Natal. Among the men over 21 years of age (88% of all patients) spree drinking and cigarette smoking were prevalent habits.

On admission to Wentworth Hospital 18% of patients were in extremis; of the remainder half were febrile and ill and half were clinically...
undisturbed. On examination the pertinent findings included severe dental caries, gingivitis, or pyorrhoea in 60% and finger clubbing in 40% of patients. Almost all patients had clinical signs in their chests. Evidence of high alcohol consumption was seen in a number of patients who had patchy depigmentation of the lips, but other evidence of vitamin deficiency or malnutrition was rare.

**Radiography**

The radiographic appearance of CDP is variable. The lesions are single, circumscribed, and rarely lobar or segmental in shape. Chronic destructive pneumonia may present as a uniform opacity (figs 1, 2), or may show more diffuse shadowing, frequently becoming cavitated, usually with multiple loculi (figs 3, 4). The chest radiograph does not return to normal after CDP has become clinically quiescent. Residual dense linear opacities are evidence of a healed lesion.

Bronchography was carried out in 88% of patients. Bronchial distortion and displacement were delineated and the almost inevitable sequelae of CDP were bronchiecstasis and destruction of pulmonary tissue and bronchi by cavitation.

In this series CDP originated in the right upper lobe in 30% of patients, in the left upper lobe in 13%, and in the lower lobar apical segments in 12%. When the disease recurred it was at the site of the original lesion.

**Microbiology**

The microbiology of CDP was investigated in 50 consecutive cases. Specimens were collected by invasive methods—percutaneous lung puncture, transtracheal aspiration, and open lung biopsy—and were processed by aerobic and anaerobic techniques. The results are set out in table 1.

In 27 of the 50 patients both aerobes and anaerobes were recovered, aerobes only in 13, anaerobes only in three, and isolates were sterile in seven. In six patients putrid specimens were obtained but there was a failure to recover anaerobes. If putrid odour is taken to be pathognomic of anaerobic infection cultural or clinical evidence of anaerobic infection was found in 36 patients.

Sputum specimens from each patient were examined concurrently for aerobic bacteria and fungi. The bacteria isolated from the sputum were similar to those found on invasive sampling in 11 of the 50 patients; in the others cultures of sputum were sterile or yielded species absent from the invasive samples.

All patients had at least six consecutive daily sputum samples examined by direct smear and concentration techniques and by culture for mycobacteria. Trap specimens were also taken at bronchoscopy with a MacRae mucus extractor. Mycobacteria were not isolated.
Table 1  Organisms recovered by invasive sampling

<table>
<thead>
<tr>
<th>Type of organism</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerobes</td>
<td></td>
</tr>
<tr>
<td>Streptococcus viridans</td>
<td>20</td>
</tr>
<tr>
<td>Streptococcus faecalis</td>
<td>11</td>
</tr>
<tr>
<td>Streptococcus haemolyticus</td>
<td>8</td>
</tr>
<tr>
<td>Streptococcus other subspecies</td>
<td>5</td>
</tr>
<tr>
<td>Staphylococcus epidermidis</td>
<td>10</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>6</td>
</tr>
<tr>
<td>Enterobacteria</td>
<td>13</td>
</tr>
<tr>
<td>Haemophilus species</td>
<td>9</td>
</tr>
<tr>
<td>Others</td>
<td>8</td>
</tr>
<tr>
<td>Anaerobes</td>
<td></td>
</tr>
<tr>
<td>Bacteroides melaninogenicus</td>
<td>11</td>
</tr>
<tr>
<td>Bacteroides other subspecies</td>
<td>19</td>
</tr>
<tr>
<td>Others</td>
<td>14</td>
</tr>
<tr>
<td>Fungi</td>
<td></td>
</tr>
<tr>
<td>Candida</td>
<td>3</td>
</tr>
</tbody>
</table>

operative specimens from 158 patients, and on the necropsy findings in five patients who died during medical treatment.

The pleurae covering the lesion were fused, thickened, and fibrotic yet highly vascularised (fig 5) by blood vessels derived from the parietes and the lung. Pleural effusion and secondary empyema thoracis have not been encountered.

Fig 3  Postero-anterior chest radiograph taken on the third admission for CDP of a 46-year-old Indian man known to be a heavy drinker. On examination he had pyorrhoea and finger clubbing. While on medical treatment he developed massive haemoptysis.

Fig 4  Typical postero-anterior chest radiograph of clinically acute (pathologically “active”) CDP of the right lung. Multiple cavities are seen, some containing fluid levels.

Fig 5  Vascular pleura from specimen shown in fig 6. Haematoxylin and eosin, original magnification x 25.

Pathology

The description of the gross and microscopic pathology of CDP is based on examination of the
Characteristics and management of chronic destructive pneumonia

The lung in CDP was airless and where the lesion had invaded a neighbouring lobe the confluent inflammatory mass obliterated all trace of the interlobar fissure (fig 6). The pulmonary and bronchial lesions in clinically acute and quiescent CDP are summarised in table 2. It is usual to find areas of “active” and “inactive” disease juxtaposed in CDP parenchyma with one or the other type of lesion dominant (figs 6, 7). As in bronchiectasis there is enlargement of the bronchial arteries supplying the affected lung, but the striking histopathological feature of the pulmonary blood vessels surviving in an active lesion is obliteration of the lumen.

Immunology

An assessment of immunological function was undertaken in a consecutive series of 20 African men aged from 20 to 50 years. The results were compared with those of a control group of healthy African men of similar age range. The tests used were total lymphocyte counts and T and B lymphocyte counts, assays of immunoglobulins G, A, and M, and of C3 by means of standard radial immunodiffusion technique, and lymphocyte response to stimulation by the mitogen PHA.

All patients had normal or somewhat increased numbers of circulating lymphocytes (mean 3517 cells/mm³, range 1512–5830 cells/mm³). Proportions of T, B, null, and “double marker” lymphocytes were normal. No instance of impaired ability to produce antibody was found, all patients having normal or raised levels of IgG, IgA, and IgM. C3 levels were normal. T lymphocyte formation was impaired in six of the 20 patients with tritiated thymidine uptakes between 20% and 40% of the normal control value.

Management

The antibiotic regimens used in the management of CDP altered during the 12 years of this study. From 1966 to 1969 the regimen was based on chloramphenicol, and from 1969 to 1976 on cephalosporins. If indicated by isolation of particular sputum pathogens other antibiotics were also given. In order of frequency they were the penicillins, aminoglycosides, sulphonamides, and tetracyclines. Forty per cent of patients also had antituberculosis chemotherapy. Between 1976 and 1977 a combination of cephalosporin, penicillin, and metronidazole was standard therapy. The drug regimens are given in table 3.

Table 2 Pathology of CDP

<table>
<thead>
<tr>
<th>Classification</th>
<th>Tissue</th>
<th>Gross pathology</th>
<th>Histopathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active</td>
<td>Lung</td>
<td>Consolidated lung containing pus and slough-filled cavities</td>
<td>Exudative pneumonia with alveolar necrosis and cavitation limited by pyogenic membrane</td>
</tr>
<tr>
<td></td>
<td>Bronchi</td>
<td>Purulent bronchitis</td>
<td>Acute bronchitis with mucosal ulceration in the draining bronchi and bronchial destruction in the CDP lesion</td>
</tr>
<tr>
<td>Inactive</td>
<td>Lung</td>
<td>Fibrous replacement of lung tissue containing residual cavities lined by granulation tissue</td>
<td>Fibrosis of cavity walls and organising pneumonia</td>
</tr>
<tr>
<td></td>
<td>Bronchi</td>
<td>Bronchiectasis</td>
<td>Follicular bronchiectasis with peribronchial fibrosis and epithelial metaplasia</td>
</tr>
</tbody>
</table>
In general the drugs were given in high dose to inpatients who were severely ill (patients presenting with massive haemoptysis or uncontrolled pulmonary suppuration or both) or as cover therapy to patients submitted to operation. Low dose oral therapy was given to the other inpatients and to outpatients who remained on maintenance therapy until there was radiographic as well as clinical evidence of healing.

Chest physiotherapy was routine treatment for all hospital patients with CDP.

The decision to operate on patients with CDP was always taken with reluctance. The operative morbidity and mortality has been greater than in the management of other pulmonary lesions, and the selection of indications for operation in CDP has been restricted by fear of the postoperative sequelae. The indications for urgent operation were exsanguinating haemoptysis, uncontrolled pulmonary suppuration in patients who, despite intensive medical treatment both at the referring clinics and at Wentworth Hospital, continued to produce profuse purulent sputum and had fever, increasing anaemia, polymorphonuclear leucocytosis, and a rapidly enlarging pulmonary opacity on chest radiograph; and when the lesion might have been a bronchial carcinoma. Elective operations were confined to patients suffering a recurrence of symptoms with enlargement of the radiographic capacity at the site of previously recognised and medically treated CDP, and when there was symptomatic

Table 3 Drug regimens in CDP

<table>
<thead>
<tr>
<th>Clinical status</th>
<th>Duration of therapy</th>
<th>Drug regimen</th>
<th>Cephalosporin-based</th>
<th>Combination therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Chloramphenicol-based</td>
<td>Cephalosporin-based</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chloramphenicol</td>
<td>Cephalothin</td>
<td>Cephalothin</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1–4 g oral or im</td>
<td>8–12 g iv or cephradine</td>
<td>8 g iv, benzylpenicillin</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Chloramphenicol</td>
<td>4–8 g iv or cefoxitin 6 g iv</td>
<td>8 megaunits iv, metronidazole 600 mg oral</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1–2 g oral</td>
<td>Cefalexin</td>
<td>Cefalexin</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2 g oral or cephaloridine</td>
<td>2 g oral, phenoxymethylpenicillin</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2 g im</td>
<td>1–2 g oral, metronidazole 600 mg oral</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Phenoxymethylpenicillin 500 mg oral, metronidazole 400 mg oral</td>
</tr>
<tr>
<td>Inpatients</td>
<td>5–14 days</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe illness or submitted to operation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>7–14 days</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Outpatients</td>
<td>Up to 9 months</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

All dosages are given as daily totals for adults.
chronic sepsis in lower lobar lesions. The results of treatment are shown in Table 4.

**Discussion**

There are published descriptions of infective pulmonary lesions which clinically and pathologically resemble CDP. The nomenclature of the lesions is various since they are sometimes associated with a demonstrable or inferred aetiology and sometimes with one or other part of the pneumonia-bronchiectasis-lung abscess-lung organisation complex. The best documented group of these CDP-like lesions is that in which anaerobes are incriminated as pathogens. There is also a more heterogeneous group where the aetiology is uncertain.

A recent editorial in the *Lancet* listed in historical order a series of papers, all Western in origin, associating anaerobes with pleuropulmonary disease. The gap in publication between 1946 and 1966 reflects an almost complete disappearance of CDP-like pneumonia from the West. "Deep lung rot"—a colloquialism of the 1930s and 1940s—had become a rarity whose passing was marked by the papers of Sellors et al and Logan and Nicholson. Since 1966 the role of anaerobes in necrotising pneumonia, bronchiectasis, and lung abscess has been elucidated, and present microbiological evidence suggests that CDP belongs to this group of pulmonary disease.

If mixed aerobic and anaerobic infection is the cause of CDP as the microbiological investigation indicates, then aspiration from the mouth is the probable initiating event in the largely male population of patients where dental disease and alcohol abuse are rife. The frequent radiographic localisation of CDP lesions in the upper lobes and apical segments of the lower lobes also suggests aspiration.

Other aetiological influences such as malnutrition and vitamin deficiency are not evident and the immunological study described above has failed to demonstrate immunological deficiency (the impaired PHA response noted in six of the 20 patients is likely to be the result of the severe infection rather than its cause).

Finegold has listed the effective antibiotics in anaerobic infection as penicillin, cephalosporins, tetracyclines, chloramphenicol, and clindamycin. Therefore the CDP patients on the chloramphenicol and cephalosporin based regimens were not given inappropriate therapy for a mixed aerobic-anaerobic infection. However, the pathology of CDP, in particular the presence of large volumes of slough in "active" lesions and of fibrosis in "inactive", complicates treatment since such damaged lung is inaccessible to antibiotics and must lack the normal immune mechanisms. The cephalosporin, penicillin, and metronidazole combination was, therefore, introduced to counter the microbiological and pathological characteristics of CDP. The combination is bactericidal and has the theoretical advantages of in vitro effectiveness against the range of pathogens, tissue penetration, dose flexibility, and relative absence of side effects. Favourable reports have been recently published on the use of antibiotic combination therapy in mixed aerobic-anaerobic infections and on the empirical use of combination therapy when such infection is suspected on clinical grounds.

It is recognised that death from pneumonia may occur despite treatment with effective antibiotics, and the mortality rates for certain types of pneumonia are high—25% in necrotising pneumonia, a disease which closely resembles CDP. Although surgical intervention is important in the general management of anaerobic infection, anaerobic pulmonary disease is an exception. Contemporary opinion holds that antimicrobial therapy, physiotherapy with postural drainage, and therapeutic bronchoscopy are adequate management. Surgical drainage is rarely if ever indicated and pulmonary resection is not considered. Nevertheless 25% of our patients have required pulmonary resection and 2% cavitary drainage because of failure of medical management to control progress of the disease and threat to life. Recognition of the importance of the anaerobic component of CDP and the consequent adjustment of antimicrobial therapy has not decreased the necessity for surgical intervention (table 4). Indeed the highest
resection rate of 33% occurs in patients treated with the cephalosporin, penicillin, and metronidazole combination. It has been shown that combination therapy lowers the inpatient mortality rate, but lung destruction is not prevented and the cycle of quiescence and recurrence of CDP is not broken in many of the medically treated patients. Should the trend to lower operative morbidity and mortality continue (table 4), it might be correct to widen the indications for surgery to include all those of Reed and Allbritten, since resection is the only means of ridding the patient of damaged lung, inaccessible to antibiotics and a permanent danger to health and life.

Bacterial pneumonia is a major public health problem in tropical developing countries, and based on the South African experience CDP is likely to be a significant facet of this problem. However there are no reports available from these regions which are probably the world reservoirs of the disease and where it may masquerade as pulmonary tuberculosis. Prevention of CDP awaits attainment of Western standards of living and health, particularly dental, care.

This study was supported in part by grants from the Group Chairman's Fund of Anglo-American Corporation, and the South African Medical Research Council.

References

Characteristics and management of chronic destructive pneumonia.

E W Cameron, P C Appelbaum, D Pudifin, W S Hutton, S A Chatterton and J Duursma

Thorax 1980 35: 340-346
doi: 10.1136/thx.35.5.340

Updated information and services can be found at:
http://thorax.bmj.com/content/35/5/340

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/