Blood culture bottle culture of pleural fluid in pleural infection

Sarah M Menzies,1 Najib M Rahman,1 John M Wrightson,1 Helen E Davies,1 Robert Shorten,2 Stephen H Gillespie,2 Christopher W H Davies,3 Nick A Maskell,4 Andrew A Jeffrey,5 Y C Gary Lee,1 Robert J O Davies1

ABSTRACT

Background Pleural infection is common, and has a >30% major morbidity and mortality—particularly when infection is caused by Gram-negative, Staphylococcus aureus or mixed aerobic pathogens. Standard pleural fluid culture is negative in ~40% of cases. Culturing pleural fluid in blood culture bottles may increase microbial yield, and is cheap and easy to perform.

Objectives To determine whether inoculating pleural fluid into blood culture bottles increases the culture positivity of pleural infection over standard laboratory culture, and to assess the optimum volume of inoculum to introduce.

Methods 62 patients with pleural infection were enrolled. Pairs of aerobic and anaerobic blood culture bottles were inoculated at the bedside with 2, 5 or 10 ml of pleural fluid, and two pleural fluid specimens were sent for standard culture. Pleural fluid from nine control patients was cultured to test for ‘false-positive’ results.

Results The addition of blood culture bottle culture to standard culture increased the proportion of patients with identifiable pathogens by 20.8% (20/53 (37.7%) to 31/53 (58.5%) (difference 20.8%, 95% CI difference 8.9% to 20.8%, p<0.001)). The second standard culture did not similarly improve the culture positivity (19/49 (38.8%) to 22/49 (44.9%) (difference 6.1%, 95% CI difference −2.5% to 6.1%, p=0.08)). The culture inoculum volume did not influence bacterial isolation frequency. The control fluids were culture negative.

Conclusions Blood culture bottle culture of infected pleural fluid increases microbial yield when used in addition to standard culture. This technique should be part of routine care.

INTRODUCTION

Pleural infection is a major cause of morbidity or death, and its incidence continues to rise in adults and children.1–4 Identification of the infecting bacteria by culture of pleural fluid is important for clinical care. This is often unsuccessful, as conventional Gram stain and culture are negative in ~40% of cases.2 These patients are treated with empirical antibiotics that cover the spectrum of likely pathogens, resulting in polypharmacy and its associated disadvantages. Anaerobic antibiotic treatment is frequently given empirically, as anaerobes are often implicated in this syndrome—although their pick-up rates by standard laboratory cultures are poor.5–7 More reliable culture methods for anaerobic bacteria could reduce the frequency with which this empirical treatment is given, if clinical confidence of the absence of anaerobic infection could be guaranteed.

Mortality is increased to ~40% in cases of Gram negative, Staphylococcus aureus or mixed aerobic infection,8–9 and reliable identification of these groups would allow the targeting of early aggressive therapy (such as video-assisted thoracoscopic surgery (VATS) surgical drainage) to selected patients.

Inoculating pleural fluid into blood culture bottles at the bedside can potentially improve the yield of pleural fluid culture.10–11 Previous studies of this approach are small, not restricted to patients with pleural infection and have not had a control intervention—making it unclear whether this benefit is real and clinically valuable. However, initial results suggest clinically higher bacterial isolate rates.12–16 In other clinical settings, inoculation of peritoneal dialysate,14 17–19 peritoneal fluid20–22 and synovial fluid14 23 24 into culture bottles has been shown to be clinically useful.

This study assesses whether inoculating possibly infected pleural fluid into aerobic and anaerobic blood culture bottles identifies more bacteria than simply repeating standard laboratory culture (the control intervention). It also assesses the optimum volume of pleural fluid inoculum to be used in this approach.

METHODS

This prospective study recruited 62 patients with suspected pleural infection and nine control patients from four centres in the UK—Oxford...
Patients with pleural infection

Sequential patients with potential pleural infection were identified at each centre and offered entry into the study if they fulfilled the inclusion criteria (box 1) which required written informed consent.

Following recruitment, patients were managed by the recruiting physicians according to local policy. Pleural fluid obtained at the time of diagnostic sampling was divided at the bedside as follows:
1. A 10 ml aliquot into a sterile container for standard Gram stain and culture at the recruiting hospital laboratory.
2. A 10 ml aliquot for repeat standard culture at a second laboratory (Royal Free Hospital, London, UK).
3. Pairs of aerobic and anaerobic blood culture bottles (BACTEC PLUS, Becton Dickinson, Sparks, Maryland, USA. Soybean-casein digest broth 2.75% w/v) were inoculated with either 2, 5 or 10 ml of pleural fluid for culture at the Royal Free Hospital.

Venous blood cultures were performed at each centre at the time of recruitment.

Sixty-two patients were enrolled in the study, 57 of which were included in the final analysis. Five patients were withdrawn—four had insufficient data and one was subsequently diagnosed with Mycobacterium tuberculosis, where this diagnostic strategy is inappropriate.

Control patients—patients without pleural infection

Control patients with pleural effusions secondary to heart failure or malignancy were offered entry into the study if they fulfilled the inclusion criteria (box 2). All of these patients were recruited at the Oxford Radcliffe Hospitals Trust during investigation and/or management of a pleural effusion of non-infective aetiology.

Pleural fluid obtained at sampling was divided by the same method as for patients with pleural infection, with the exception that only one pair of blood culture bottles was sent (each with 10 ml inoculum volume).

Sample handling and laboratory methods

Samples were transferred to a central analysis centre (Royal Free Hospital) by urgent courier. The standard sample was centrifuged at 3000 rpm for 10 min, the deposit being used for Gram staining and inoculation onto blood, MacConkey, chocolate and anaerobic media. Blood and MacConkey plates were incubated aerobically, and the chocolate plates in 5% CO2, all at 35°C for 18 h. The anaerobic plates (with metronidazole discs) were incubated anaerobically at 35°C for 48 h.

The blood culture bottles containing pleural fluid were incubated in the BacTec 960 (Becton Dickinson) at 35°C for up to 5 days. A Gram stain was performed on any positive bottles and subculture was performed onto appropriate solid media to identify the organism(s) and perform antibiotic susceptibility testing. Isolates of a coagulase-negative Staphylococcus were considered to be a contaminant and were not included in the data analysis.

Power calculation

Sixty patients were required to exclude the minimum clinically significant improvement in isolate rate of 25% (from 50% to 75%) with blood culture bottle culture (α=0.05, power 90%, the McNemar test).

Statistical analysis

Comparison of the microbiological sensitivity of tests was undertaken using the McNemar test for paired proportions (STATA version 9.2).

RESULTS

The baseline characteristics are given in table 1, and microbiological results from the two standard cultures and the blood culture bottle culture are given in table 2.

Patients with pleural infection

Primary analysis

In 53 patients, a full set of specimens allowed direct comparison between methods.

Addition of blood culture bottle culture to the standard laboratory culture increased the frequency of bacterial isolation from 20/53 (37.7%) cases with standard culture to 51/53 (58.5%) cases with the addition of blood culture bottle culture (difference 20.8%, 95% CI difference 8.9% to 20.8%, McNemar test=11.0, 1 degree of freedom (df), p<0.001).

Of the 31 cases with bacterial isolates combining these techniques, six had the same organisms identified by both, and four had different organisms identified. In nine cases only the standard culture was positive, and in 12 cases only the blood culture bottle culture was positive. Blood culture bottle culture also added microbiological value to standard culture in a further two (5.8%) cases, by identifying additional microbiologically relevant organisms when the standard culture was already positive (one Enterococcus spp., and one Streptococcus spp.—when standard culture was solely anaerobic).

Box 1 Clinical features of patients with pleural infection

- Clinical presentation compatible with pleural infection
- Gram stain or culture positive
- pH <7.2 by blood gas machine analysis
- Excess pleural fluid available after standard sampling

Box 2 Clinical features of patients without pleural infection

A. Heart failure patients
- Clinical presentation compatible with heart failure and not infection
- Pleural effusion present requiring drainage
- Transudative pleural effusion (protein <30 g/dl, or pleural fluid:plasma lactate dehydrogenase (LDH) ratio <2/3), with a pH >7.2
- Excess pleural fluid available after standard sampling

B. Patients with malignant effusions
- Proven malignant effusion requiring drainage
- Pleural fluid biochemistry supporting a non-infected effusion (pH >7.3, glucose >2.2 mmol/l and LDH <1000 IU/l)
- No clinical features of infection or raised temperature >37.5°C and/or venous blood C-reactive protein >20 mg/l
- Excess pleural fluid available after standard sampling


Radcliffe Hospitals Trust, Northampton General Hospital (Northampton), Southmead Hospital (Bristol) and the Royal Berkshire Hospital (Reading).
Repeating the standard culture in a second laboratory (control intervention)

Standard culture results from both the recruiting hospital and the second laboratory were available in 49 cases. Addition of the information provided by the second laboratory culture produced a clinically insignificant increase in bacterial yield (6.1%). This tended towards, but did not achieve, statistical significance—19/49 (38.8%) culture-positive cases on standard culture at the recruiting hospital, increased to 22/49 (44.9%) with the addition of a repeat culture-positive cases on standard culture at the recruiting hospital and the Royal Free laboratory standard culture.

Recent pleural procedure performed that may have introduced infection, no. of patients (%)

Hospital admission in the month prior to enrolment, no. of patients (%)

Pleural fluid characteristics

Visible purulent, n (%) 23 (51) 0 (0)

pH (SD) 6.9 (0.3) 7.6 (0.1)

Glucose (mmol/l), median (IQR) 1.0 (0.1–2.1) 6.0 (5.1–6.4)

Protein (g/l), median (IQR) 42 (34–49) 30 (25–34)

Lactate dehydrogenase* (IU/l), median (IQR) 3192 (1006–8864) 65 (49–125)

Venous blood results

White cell count (×10⁹/l), median (IQR) 13.6 (9.1–19.0) 6.3 (5.5–8.9)

*Serum upper limit of normal:200 IU/l.

Adding blood culture bottle culture to the combined results of both standard cultures

The frequency of pathogens identified by both standard laboratory cultures combined was 22/49 (44.9%) cases, which increased to 30/49 (61.2%) cases when blood culture bottle culture was added (difference 16.3%, 95% CI difference 4.3% to 16.3%, McNemar test=8.0, 1 df, p=0.005).

Blood culture bottle culture still provided extra microbiological value when standard cultures were already positive in the two (4.1%) additional cases described earlier. Figure 1 shows the additional sensitivity of the two standard cultures and blood culture bottle culture for each of the infecting bacterial classes. The pathogen classes match those shown to have prognostic significance.8 9

Venous blood cultures

Venous blood cultures were taken in 51/57 pleural infection cases, and were positive in only 6/51 (11.8%). Of the six positive cultures, three did not add any clinical value as the organism was also isolated from pleural fluid. In two cases they were the only positive culture for the patient (Streptococcus pneumoniae and Enterobacter aerogenes), and one cultured S pneumoniae—when only S aureus was isolated from the pleural fluid cultures (from the blood culture bottle culture).

Pleural fluid inoculum volume for blood culture bottle culture

It was possible to compare the 2, 5 and 10 ml inoculum volume in 11 patients with bacterial isolates from the aerobic culture bottles, and in 20 patients with bacterial isolates from the anaerobic culture bottles. There was no evidence of increased sensitivity with any of the inoculum volumes (table 3).

Control patients without pleural infection

In the control patients, both sets of standard cultures and all blood culture bottle cultures were negative, with the exception of one standard culture sample that isolated Streptococcus viridans. Neither follow-up clinical information nor pleural fluid parameters from sampling supported the diagnosis of pleural infection.

DISCUSSION

This is the first adequately powered prospective controlled study of culturing infected pleural fluid in blood culture bottles. Our results suggest that inoculation of pleural fluid into blood culture bottles increases diagnostic yield and the number of multiorganism infections detected. These data support the use of blood culture bottles in routine care for patients with suspected pleural infection.

The addition of blood culture bottle culture to standard culture increased the number of culture-positive cases in pleural infection by 20.8% (from 37.7% to 58.5%). This is consistent with a previous study, in which blood culture bottle culture

Table 1  Baseline characteristics of pleural infection and control patients

<table>
<thead>
<tr>
<th>Variable</th>
<th>Patients with pleural infection (n = 57)</th>
<th>Patients without pleural infection (controls) (n = 9)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex, n (%)</td>
<td>Male 38 (67)</td>
<td>8 (89)</td>
</tr>
<tr>
<td>Age, years (SD)</td>
<td>60 (18)</td>
<td>83 (10)</td>
</tr>
<tr>
<td>Duration of pleural infection symptoms before enrolment (days), median (IQR)</td>
<td>19 (10–30)</td>
<td>NA</td>
</tr>
<tr>
<td>No. of antibiotic doses before enrolment, median (IQR)</td>
<td>15 (6–35)</td>
<td>NA</td>
</tr>
<tr>
<td>Recent pleural procedure performed that may have introduced infection, no. of patients (%)</td>
<td>12 (21)</td>
<td>5 (56)</td>
</tr>
<tr>
<td>Hospital admission in the month prior to enrolment, no. of patients (%)</td>
<td>7 (13)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

Table 2  Pleural fluid culture results for each culture method

<table>
<thead>
<tr>
<th>Home laboratory standard culture</th>
<th>Royal Free laboratory standard culture</th>
<th>BCB culture (combined aerobic and anaerobic)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infection patients, n (%)</td>
<td>Controls, n (%)</td>
<td>Infection patients, n (%)</td>
</tr>
<tr>
<td>No pathogen</td>
<td>33 (62)</td>
<td>8 (89)</td>
</tr>
<tr>
<td>Streptococcus pneumoniae</td>
<td>5 (9)</td>
<td>0</td>
</tr>
<tr>
<td>Streptococcus anginosus group*</td>
<td>4 (8)</td>
<td>0</td>
</tr>
<tr>
<td>Other streptococci</td>
<td>0</td>
<td>1 (11)</td>
</tr>
<tr>
<td>Anaerobic: aerobic</td>
<td>5 (9)</td>
<td>0</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mixed aerobic</td>
<td>1 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Gram-negative bacteria</td>
<td>5 (9)</td>
<td>0</td>
</tr>
</tbody>
</table>

*Previously termed ‘Streptococcus milleri group’. BCB, blood culture bottle.
increased culture positivity from 44% to 64%. This study adds to the former by being adequately powered and having a control intervention. Together, these two studies show that blood culture bottle culture is a valuable adjunct to standard laboratory plated culture within the context of pleural infection, and should be part of standard care.

It is interesting that this finding is very similar to the results in other ‘non-blood’ uses of blood culture bottle culture. In bacterial peritonitis, blood culture bottle culture increases the bacterial isolation rate by 29–49%,19–21 from a positivity rate on standard culture of 42–54%. In synovial fluid culture, blood culture bottle culture increases the bacterial isolation rate by up to 20% over the 10–20% culture positivity with standard culture.14 23 24

In addition to increasing the organism identification rate over standard culture at the recruiting hospital, blood culture bottle culture identified additional clinically important co-infecting bacteria in 2/53 (3.8%) cases. In the first case, standard culture isolated methicillin-resistant S aureus (MRSA), Pseudomonas aeruginosa and anaerobes, whereas bottle culture identified an Enterococcus spp.—which was of clinical value given the resistance patterns these organisms exhibit. In the second, standard culture grew only anaerobes, and bottle culture identified an aerobic growth of an organism from the Streptococcus anginosus group.

S pneumoniae isolation rates were higher from standard culture at the recruiting hospital compared with standard culture and blood culture bottle culture at the second laboratory (9.4, 1.9 and 1.8%, respectively). This may be due to the known tendency for delays in sample processing leading to impaired isolation of this fastidious organism.23 26 However, the frequency of isolation of non-pneumococcal streptococci also differed between standard and blood culture bottle culture, with the former identifying more organisms in the S anginosus group and the latter identifying more ‘other streptococci’ (namely other α-haemolytic streptococci). It is unclear why different bacteria culture preferentially in different media, and further studies are required to address this observation. This finding emphasises the importance of employing both standard and blood culture bottle culture strategies in parallel.

This study used repeated standard laboratory culture as a control intervention to test whether the improved yield of organisms using blood culture bottles was simply due to repeating the culture process. We have shown that blood culture bottle culture is superior to simple repetition of culture, with repeat standard culture increasing the rate of organism identification by 6.1% (from 38.8% to 44.9% of cases), and blood culture bottle culture increasing this further to 61.2% (65.3% if the additional clinically important bacterial isolates are considered).

The results from control patients without clinical evidence of pleural infection showed no false-positive cases identified from blood culture bottle culture of pleural fluid. This provides reassurance that the proposed diagnostic approach is robust.

This study demonstrated no evidence of differential rates of bacterial identification between different volumes of pleural fluid inoculum. This may suggest that even small residual volumes of possibly infected pleural fluid should be subjected to blood culture bottle culture.

There are several potential limitations to this study. Firstly, there was extra transportation time involved in transferring samples to the second laboratory, which could have altered the yield of bacteria. In addition, as we showed that different bacteria may have preferential growth in different media, and the spectrum of bacterial infection differs worldwide, this study should be repeated in other regions to ascertain the exact magnitude of benefit with blood culture bottle culture. A larger study may also be needed to address whether bottle culture is particularly beneficial in subgroups of clinical settings, for example taking into consideration antibiotic treatment prior to sampling, the size of the effusion or the likely bacterial load (degree of sepsis).

**Table 3** Comparison of the different pleural fluid inoculum volumes in culture-positive patients

<table>
<thead>
<tr>
<th>Volume of inoculum</th>
<th>Aerobic blood culture bottles n = 11, number positive (%)</th>
<th>Anaerobic blood culture bottles n = 20, number positive (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 ml</td>
<td>8 (73)</td>
<td>18 (90)</td>
</tr>
<tr>
<td>5 ml</td>
<td>8 (73)</td>
<td>16 (80)</td>
</tr>
<tr>
<td>10 ml</td>
<td>8 (73)</td>
<td>15 (75)</td>
</tr>
</tbody>
</table>

Some organisms were cultured in two or three different volume bottles, hence totals are greater than the number of cases.

**CONCLUSIONS**

Inoculating pleural fluid into blood culture bottles at the bedside increases the rate of bacterial pathogen identification in pleural infection, when used in addition to standard laboratory culture. This increased yield appears to be specific to the use of blood culture bottles, is not a result of repeating the culture process and is clinically and statistically significant.

These results suggest that bedside pleural fluid inoculation into blood culture bottles should enter routine care in those with potential pleural infection.
Respiratory viruses are not associated with acute graft rejection during the acute phase of infection

Although several studies support the association between respiratory viruses and chronic lung rejection, the relationship between viral infection and acute rejection has not been established.

In this prospective cohort study, bronchoalveolar lavage (BAL) fluid and biopsy specimens from 343 bronchoscopic procedures performed in 77 patients were analysed over 27 months from November 2003 through March 2006. Real-time reverse-transcription polymerase chain reaction assays for detection of RNA respiratory viruses were performed on the BAL fluid specimens. The severity of acute rejection was graded as mild (A0, A1) to severe (A2–A4). The overall rate of viral infection in grade A0/A1 acute rejection cases was 25.4%, compared with 12.6% in grade A2 or higher rejection. Thus, patients with lower acute rejection grade were twice more likely to be positive for viral infection than those with higher rejection grades. The authors also found that in patients with simultaneous acute rejection and lower respiratory tract viral infection, the FEV1 recovery rate was significantly lower than in patients who had acute rejection without a simultaneous viral infection.

This study provides evidence that respiratory viruses per se do not promote acute graft rejection, at least during the acute phase of infection, but that they do worsen graft function recovery when simultaneously present with acute rejection.


M Elftise

Correspondence to Dr Maisun Elftise, ENT Department, University Hospital Coventry and Warwickshire, Clifford Bridge Road, Coventry, CV2 2DX, UK; maisun.elftise@uhcw.nhs.uk

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