Value of intracellular bacteria detection in the diagnosis of ventilator associated pneumonia

A Torres, M El-Ebiary, N Fábregas, J González, J Puig de la Bellacasa, C Hernández, J Ramirez, R Rodríguez-Roisin

Abstract

**Background** – Markers of ventilator associated pneumonia are of interest for confirming the diagnosis and for guiding the initial management of this frequent complication of mechanical ventilation. The detection of intracellular organisms in the polymorphonuclear leucocytes (PMNLs) and/or macrophages of bronchoalveolar lavage (BAL) fluid has been suggested as a specific test for the early indication of an infectious pulmonary process.

**Methods** – The diagnostic value of detecting intracellular organisms in two types of BAL fluid – protected (P-BAL) and conventional (C-BAL) – in 25 patients who died in one unit was prospectively studied. Immediately after death both P-BAL and C-BAL were performed bilaterally. Through a minithoracotomy on both sides of the chest bilateral bronchoscopically guided open lung biopsy samples were obtained from the same area, and an average of eight open lung blind biopsy samples (not bronchoscopically guided) were taken from each lung for histological examination. BAL fluid was examined for quantitative cultures (threshold 10⁵ cfu/ml) and for the presence of intracellular organisms and extracellular organisms, and differential cell counts were also performed.

**Results** – Using the histopathology of the bronchoscopically guided open lung biopsies as the gold standard, detection of intracellular organisms in P-BAL (≥5%) and C-BAL (≥5%) fluids yielded 75% and 57% positive predictive values, and 83% negative predictive values, respectively. Prior treatment with antibiotics decreased the positive and negative predictive values of intracellular organism detection for both types of BAL fluid. The presence of intracellular organisms was correlated with the quantitative cultures of P-BAL and C-BAL samples. Quantitative cultures from P-BAL fluid were less sensitive (22% versus 45%) and more specific (100% versus 55%) than those from C-BAL samples. The percentage of extracellular organisms and the differential cell count in P-BAL and C-BAL samples could not discriminate between the presence or absence of pneumonia.

**Conclusions** – The presence of ≥5% intracellular organisms infecting PMNLs or macrophages in P-BAL or C-BAL fluids is a specific marker of ventilator associated pneumonia.

Keywords: intracellular organisms, protected bronchoalveolar lavage, ventilator associated pneumonia.
Intracellular bacteria detection in diagnosis of ventilator associated pneumonia

Table 1  Mean (SD) characteristics of the study population

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>56-7 (16)</td>
</tr>
<tr>
<td>Male (M)</td>
<td>16/9</td>
</tr>
<tr>
<td>APACHE II before death</td>
<td>27 (6)</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>36 (1)</td>
</tr>
<tr>
<td>Leucocyte count (× 10⁹/µl)</td>
<td>14:7 (8:4)</td>
</tr>
<tr>
<td>Pao₂/Fio₂</td>
<td>207 (126)</td>
</tr>
<tr>
<td>Mechanical ventilation period (days)</td>
<td>13 (13)</td>
</tr>
<tr>
<td>No. of antibiotics</td>
<td>4 (2)</td>
</tr>
<tr>
<td>Duration of antibiotic treatment (days)</td>
<td>9-5 (7-9)</td>
</tr>
<tr>
<td>Days without antibiotics (n=8)</td>
<td>3 (2)</td>
</tr>
</tbody>
</table>

Values were obtained immediately after death (±30 minutes) at the time of initiation of the study.

APACHE—Acute Physiology and Chronic Health Evaluation score; COPD—chronic obstructive pulmonary disease; Pao₂=arterial oxygen tension; Fio₂=fractional inspired oxygen.

received a mean (SD) of 4 (2) drugs for a mean duration of 9-5 (7-9) days. These included: cefonicid, cefotaxime, ceftazidime, ciprofloxacin, fluconazol, vancomycin, clindamycin, netilmicin, trimethoprime/co-trimoxazole, imipenem, amikacin, and amphotericin B. Patients were considered not to have received antibiotics if they had not been given any for at least 48 hours before death. The administered antibiotics in these patients were: erythromycin, cefonicid, cefotaxime, ciprofloxacin, fluconazol, vancomycin, imipenem, amikacin, and amphotericin B. Causes of death included: cerebral death (12), multiple organ failure (9), and severe hypoxaemia (4). All patients except four had either localised (n=14) or diffuse (n=7) infiltrates on their chest radiographs. The clinical diagnoses of pulmonary infiltrates (n=21) on the day of death were: pneumonia (19), congestive heart failure (1), and both alveolar haemorrhage and pneumonia (1). The percentage of samples showing pneumonia was higher in lung samples from patients who presented with chest radiographic infiltrates (50% localised infiltrates and 62% diffuse infiltrates) than in those with normal chest radiographs (19%). The correlation between the percentage of positive lung biopsy samples for histological pneumonia in each patient and the time interval between chest radiographic infiltrate appearance and death was poor (r=0:2, p=NS). The main reasons for admission to the intensive care unit (ICU) were: acute respiratory failure (12), stroke (6), postoperative (3), cranial trauma (3), and myocardial infarction (1). Of the 12 cases of acute respiratory failure six were admitted with pneumonia, two with congestive heart failure, two with alveolar haemorrhage, and two with pulmonary neoplasia. Family members in each case gave informed written consent and permission was obtained from the ethical committee of our centre.

Study design
Immediately after death patients remained mechanically ventilated while breathing 100% O₂. Bilateral fibroptic bronchoscopies were performed using different bronchoscopes for each lung following standard techniques. The BAL specimens were retrieved in the order: protected bronchoalveolar lave (P-BAL) (Mill-Rose Laboratories Inc, Ohio, USA) followed by conventional BAL (C-BAL). Both types of BAL were performed by instilling an aliquot of 20 ml sterile saline which was discarded, followed by four aliquots each of 30 ml sterile saline (mean (SD) recovered BAL fluid 26 (15) ml and 45 (20) ml for P-BAL and C-BAL, respectively). Open lung bronschoscopically guided biopsy and lavage samples were taken from the same place – that is, from the area of maximal infiltrates on the chest radiograph or from lower lobes in patients without infiltrates. A total of 94 lavages (47 P-BAL and 47 C-BAL) were performed as three lung from three patients could not be studied. A minithoracotomy was performed on each side of the chest and a guided biopsy sample was taken from each lung using the light tip of the bronchoscope which was seen shining from within the lung in areas of maximal inflammation as seen on the chest radiograph. If there were no infiltrates on the radiograph the bronchoscope was placed in a lower lobe segment and biopsy samples were taken from that area. In addition, non-guided (blind) open lung biopsies were taken using strict aseptic techniques from each lung through each thora-cotomy incision. Three fragments were taken from both upper and lower lobes from each lung and from the middle lobe and lingular so that the overall number of biopsy samples per patient was 16 (two guided and 14 blind). The total number of biopsies was 375 (in three patients only one lung was sampled). The smallest size of both the bronchoscopically guided and the blind biopsy samples was 2 x 2 x 2 cm each. Each sample was divided into two pieces: one for histological examination and the other for microbiological processing. Pathological criteria for pneumonia included foci with accumulation of polymorphonuclear leucocytes (PMNs) in the capillaries and adjacent alveolar spaces corresponding to various degrees of evolution and extension.¹⁴

Microbiological processing and isolate identification
Serial dilutions (10⁻¹, 10⁻², 10⁻³) from each of the P-BAL and C-BAL samples were prepared in sterile normal saline. Biopsy samples were placed in a mortar with sterile sand and 3 ml sterile saline (Vidarroc, Barcelona, Spain) and homogenised. Serial dilutions (10⁻¹, 10⁻², 10⁻³) of each sample were prepared in sterile normal saline. One hundred µl of each dilution was inoculated into the following agar media: 5% sheep blood, chocolate, Center for Disease Control (CDC) blood, Wilkins-Chalgren, McConkey, blood charcoal yeast extract (BCYE-9), and Sabouraud-dextrose. All cultures were incubated at 37°C under aerobic and anaerobic conditions and in CO₂ enriched atmosphere. Cultures were evaluated for growth 24 hours and 48 hours later and discarded if negative after five days, except for CDC and Wilkins-Chalgren which were evaluated at seven days and for Sabouraud which was evaluated at four weeks. All micro-

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organisms isolated were identified by standard laboratory methods. Total cell counts of BAL fluid were performed using an aliquot of the original lavage fluid. Preparations were centrifuged with a Cytospin 2 cytocentrifuge (Shandon, Southern Products, UK) and two samples were stained with May-Grünwald Giemsa stain. Differential cell counts were made by examining at least 500 cells. In addition, 300 cells were examined at 1000× magnification using an oil immersion lens and the percentage of cells containing intracellular organisms per field was determined. A Gram stain of the cytocentrifuged material was made to evaluate the morphological characteristics of the bacteria.

DATA ANALYSIS

The diagnostic values of the two types of BAL fluid were calculated using 10^6 cfu/ml as a cutoff point to differentiate between colonisation and infection. Diagnostic values of quantitative cultures of both types of BAL samples were calculated excluding yeasts and non-pathogenic staphylococci (Staphylococcus epidermidis). The diagnostic values of intracellular organisms were calculated using different cutoff points (2%, 5%, and 10%) and analysed statistically using the SPSS/PC+ version 4-0.19 and the EPIInfo (version 5-01). Results are expressed as mean (SD). Quantitative variables were compared with the Mann-Whitney test and proportions with the χ² test (Fisher’s exact test when needed). Correlations were tested by linear regression analysis. Diagnostic values were calculated and interpreted according to standard methods.17,18 Data were analysed using histopathological criteria of pneumonia as the gold standard in two parallel ways: (1) open lung bronchoscopically guided lung biopsies (lung was considered to be positive for pneumonia if the open lung bronchoscopically guided biopsy sample showed histological evidence of pneumonia), and (2) blind open lung biopsies (a lung was considered positive for pneumonia if at least one of the blind open lung biopsy samples showed histopathological evidence of pneumonia). All p values are two tailed and the level of significance was set at 5%.

Results

INTRACELLULAR MICROORGANISMS, GRAM STAIN, AND CELL COUNTS

In P-BAL samples the mean percentage of intracellular organisms with colony counts of ≥10^6 cfu/ml was higher than those with cultures yielding <10^6 cfu/ml both in cases that had and had not received antibiotics (23% (range 0–80) versus 6% (range 0–90), p=NS, and 43% (range 0–90) versus 3% (range 0–12), p<0.05, respectively). The same was true for C-BAL (36% (range 0–80) versus 4% (0–90), p=0.002, and 49% (0–90) versus 14% (0–40), p<0.05, respectively).

Quantitative cultures of both P-BAL and C-BAL samples showed moderate correlations with the percentages of intracellular organisms found (r=0.7; p=0.0001; r=0.45; p=0.001, respectively), and these correlations persisted when results from patients who had and had not received antibiotics were compared (P-BAL: r=0.58 and r=0.67, p=0.001 each; C-BAL: r=0.4 and r=0.48, p=0.01 each, respectively).

Gold standard: bronchoscopically guided open lung biopsies

Cutoff points of 2%, 5%, and 10% were used to evaluate the sensitivity and specificity and positive and negative predictive values of intracellular organisms. Table 2 shows the different diagnostic parameter values for the detection of intracellular organisms in BAL fluid using the bronchoscopically guided pulmonary biopsy as a gold standard. For P-BAL in patients without prior antibiotic treatment, the threshold of 10% had the most acceptable diagnostic accuracy (sensitivity 50%, specificity 78%). However, in patients who had previously received antibiotics a threshold yielding an acceptable diagnostic value was not found. For C-BAL samples the best threshold was 5% when samples from patients who had not received antibiotics were analysed (sensitivity 75%, specificity 55%). Again, no acceptable ratio of sensitivity/specificity was found for the threshold of intracellular organisms giving an acceptable diagnostic value in patients who had been treated with antibiotics. There were no differences in the diagnostic value parameters of detection of intracellular organisms between P-BAL and C-BAL samples.
Gold standard: blind open lung biopsies
The best sensitivities for P-BAL samples were obtained using a threshold of 5% and 10% in both patients who had and had not received antibiotics (58% each). Calculated specificities were 100%. For C-BAL samples a cutoff point of 2% was the most accurate (sensitivity 67%, specificity 100%) in patients who had not been treated with antibiotics, while in those who had received antibiotics the specificity decreased to 67% (p<0.0001).

When the diagnostic value parameters obtained using bronchoscopically guided biopsies and blind biopsies as gold standards were compared, most sensitivities for P-BAL samples (both for patients with and without antibiotics and independently of the threshold chosen) were significantly higher and the specificities were invariably lower. Similar changes were seen for specificities in C-BAL specimens from patients who had not received antibiotics (tables 2 and 3).

Cell counts and extracellular microorganisms in the BAL fluid are shown in table 4. There were no significant differences between the percentages of PMNL, macrophages, lymphocytes, and extracellular microorganisms in samples from patients who had and had not received antibiotics in both types of BAL fluid samples. All samples contained less than 1% of squamous epithelial cells.

Quantitative cultures: P-BAL
Gold standard: bronchoscopically guided open lung biopsy
The sensitivity was 22%, specificity 76%, positive predictive value 54% and negative predictive value 43%. In the eight patients who were not treated with antibiotics the sensitivity of the P-BAL samples decreased significantly (from 22% to 0%; p<0.0001). No differences were shown in specificity.

Gold standard: blind open lung biopsies
The sensitivity was 22% and specificity 100% compared with the specificity of bronchoscopically guided biopsy samples; the positive predictive value was 100% and the negative predictive value 18%. In the eight patients who did not receive antibiotic treatment there were no differences in the sensitivity, specificity, and positive and negative predictive values.

Quantitative cultures: C-BAL
Gold standard: bronchoscopically guided open lung biopsy
A sensitivity of 35%, a specificity of 43%, and positive and negative predictive values of 43% and 35%, respectively, were obtained for C-BAL samples and no differences were seen between patients who did and did not receive antibiotic therapy.

Gold standard: blind open lung biopsies
The sensitivity was 45%, specificity 55%, and positive predictive and negative predictive values were 81% and 19%, respectively. In the eight patients who did not receive antibiotics the sensitivity of the C-BAL samples increased significantly (38% versus 58%; p<0.0001); there were no differences for specificity. Table 5 shows the different diagnostic values of both types of bronchoalveolar lavage using the two different types of gold standard (bronchoscopically guided and blind pulmonary biopsies). Overall, calculated sensitivities were significantly higher with C-BAL than with P-BAL samples (35% and 45% versus 22% and 22%; p<0.05 and p=0.0005, respectively). Conversely, specificities were higher with P-BAL than with C-BAL samples (76% and 100% versus 43% and 55%; p=0.00001 and p<0.0001, respectively).

When the diagnostic value parameters of P-BAL and C-BAL samples were considered using blind biopsy samples as the gold standard, higher specificity, higher positive predictive values, and lower negative predictive values were obtained in all cases than those obtained from bronchoscopically guided biopsies (table 5).
AGREEMENT BETWEEN BAL CULTURES
Sixty three microorganisms were isolated from P-BAL cultures, of which 50 (79%) were com-
comitantly isolated from C-BAL cultures. Conversely, 74 microorganisms were isolated from
C-BAL cultures of which 50 (67%) were also concomitantly isolated from P-BAL cultures.
Quantitative cultures of P-BAL correlated moderately with those of C-BAL (r = 0 5, p = 0 02).
Table 6 shows the number of species isolated in counts of $\geq 10^4$ cfu/ml from both P-BAL
and C-BAL fluids and the mean percentages of intracellular organisms in relation to
the presence or absence of histological pneumonia and as a function of the type of biopsy
sample obtained (bronchoscopically guided or blind pulmonary biopsy).

RELATIONSHIP BETWEEN BAL QUANTITATIVE CULTURES AND PNEUMONIA
The following potentially pathogenic micro-
organisms were isolated in counts of $\geq 10^4$ cfu/ml from P-BAL fluid cultures corresponding
with histological pneumonia in bronchosco-
icly guided biopsies: Pseudomonas aeruginosa (4), Staphylococcus aureus (1), Xan-
thomonas maltophilia (1), and Candida albicans
(1). The microorganisms isolated in counts of
$\geq 10^4$ cfu/ml from C-BAL cultures from lungs
with histological pneumonia were: Pseudomonas aeruginosa (3), Staphylococcus aureus (1), Xan-
thomonas maltophilia (1), Serratia marcescens (1),
Pseudomonas putida (1), Candida albicans (1),
Enterococcus faecalis (1), and Proteus mirabilis
(1).

HISTOLOGY AND MICROBIOLOGY OF LUNG
BIOPSIES
Twenty five of 47 (53%) pulmonary biopsy
specimens obtained showed histological signs
of pneumonia compared with 174 of 375 (46%)
of the blind pulmonary biopsy specimens.
Bronchiolitis was seen in 33 samples (9%) from
13 patients.

Bronchoscopically guided biopsies yielded
positive cultures in 36 of 47 (76%) specimens.
In those samples with histological evidence of
pneumonia (n = 25), cultures were $\geq 10^4$ cfu/g
in seven biopsies, $\leq 10^3$ cfu/g in 13 biopsies,
and negative in five. Blind biopsies yielded
positive bacterial cultures on 279 of 375 (74%)
occasions. In those samples with histological
evidence of pneumonia (n = 174) cultures were
above the threshold of $10^4$ cfu/g in 71 cases
(41%), $10^3$ cfu/g in 67 (38%), and negative in
36 (21%). Culture of bronchoscopically guided
biopsies of P-BAL and C-BAL samples were
similar in 30 of 47 (64%) studies. Biopsy cul-
tures coincided with P-BAL cultures in 34 of
47 (72%) cases and with C-BAL in 36 (76%).
In seven studies (15%) there was no agreement
in culture results with either technique.

We found moderate correlations between
quantitative cultures of coincident micro-
organisms isolated from bronchoscopically
guided lung biopsies and quantitative cultures
of P-BAL (r = 0 55, p<0 005) or C-BAL
samples (r = 0 57, p<0 005).

Discussion
The main finding of this study is that the
presence of intracellular bacteria in the PMNLs
or macrophages of the centrifuged fluid from
P-BAL (≥5%) or C-BAL (≥2%) samples is
a reasonable diagnostic marker of ventilator
associated pneumonia. However, these results
are influenced by prior antibiotic treatment
(lower sensitivity and specificity).9,20

The diagnosis of ventilator associated pneu-
monia still remains elusive. An important issue
is that the results of quantitative culture are
delayed for at least three days, resulting in
empirical antibiotic treatment in the interim.
Given the fact that clinical features of pneu-
monia in mechanically ventilated patients are
neither specific nor sensitive, rapid markers of
pulmonary infection might be of great value to
the clinician to help prescribe an antibiotic
regimen. Several markers have been reported
in the literature. The direct examination for
the presence of elastin fibres may help in the
diagnosis of necrotising pneumonias.8,21
Other markers in respiratory secretions such as
lactate dehydrogenase and endotoxin are being
investigated.6,7

The percentage of intracellular organisms in
the PMNLs and macrophages of the recovered
BAL fluid deserves particular attention. Initial
studies from the same group5,22 found that the
use of this technique on C-BAL samples has a
reasonable sensitivity and 100% specificity for
the diagnosis of ventilator associated pneu-
monia. Yet, there was discrepancy in the threshold
for differentiating the presence of pneumonia.
More investigations using P-BAL
samples23 were very sensitive and specific when
choosing 2% intracellular organisms as a cutoff
point.

Table 6 Numbers of lungs with and without pneumonia (n = 47), numbers of isolates of bacterial species from BAL
fluids in significant counts ($\geq 10^4$ cfu/ml), and percentages of intracellular microorganisms

<table>
<thead>
<tr>
<th>Pneumonia*</th>
<th>No. of species ($\geq 10^4$ cfu/ml)</th>
<th>Mean (range) intracellular organisms (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P-BAL</td>
<td>C-BAL</td>
</tr>
<tr>
<td>Bronchoscopically guided open lung biopsies:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>25</td>
<td>6</td>
</tr>
<tr>
<td>No</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Lungs (at least one positive blind open lung biopsy per lung):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>40</td>
<td>13</td>
</tr>
<tr>
<td>No</td>
<td>7</td>
<td>3</td>
</tr>
</tbody>
</table>

* Presence or absence of histological pneumonia (bronchoscopically guided lung biopsy).
Other investigators have confirmed the value of the detection of intracellular organisms and have suggested 5% of PMNLs containing intracellular organisms as the ideal threshold to differentiate between the presence and absence of pneumonia. However, recent publications could not find an adequate cutoff point and the effect of antibiotic treatment was not specifically explored in these studies. Moreover, the gold standards used were limited to the presence of clinical signs of pneumonia or to the bacteriological cultures of protected specimen brush or BAL fluid only. When the histological results of blind necropsy open lung biopsy specimens were used as a gold standard we found that the presence of ≥5% of PMNLs with intracellular organisms in P-BAL samples or ≥2% in C-BAL samples was a sensitive (approximately 60%) and very specific (100%) tool for detecting the existence of pneumonia.

There are, however, some findings and limitations of our study that need to be emphasised. Firstly, the high specificity of intracellular organism detection found in our study is conditioned by the high prevalence of histological pneumonia (in blind open lung biopsy specimens) found in our deceased critically ill patients. Secondly, the diagnostic value parameters of intracellular organisms, particularly the sensitivity, were strongly influenced by prior antibiotic treatment. This has been reported by others who showed a reduction in the predictive accuracy of intracellular organisms in detecting ventilator associated pneumonia in patients receiving or having recently received antibiotics. This finding probably does not apply to all patients and depends upon the intrinsic characteristics of each antibiotic (intracellular penetration, accumulation, and disposition). Thirdly, the diagnostic value of intracellular organism detection was generally independent of the type of BAL fluid used. In C-BAL samples a lower threshold of intracellular organisms (≥2%) was enough to differentiate between the presence and absence of pneumonia in patients not receiving antibiotics. However, our results could have been biased as, due to design constraints, P-BAL sampling was always performed before C-BAL sampling in each lung of our cadaveric model without evaluating an order effect. This was because the main aim of our study was to explore the results of P-BAL. Fourthly, the diagnostic value of intracellular organisms was altered by the type of gold standard used – bronchoscopically guided open lung biopsy specimens or multiple blind open lung specimens. The diagnostic value of intracellular organisms for both types of BAL samples were more specific when the analysis was taken into consideration lung-by-lung and in the presence of histological pneumonia if at least one of the multiple blind open lung biopsy specimens was compatible with it. What may occur in ventilator associated pneumonia is that, once foci of pneumonia develop, infected secretions may be redistributed to other bronchopulmonary segments, possibly enhanced by the volume and flow patterns generated by the ventilator. This is why positive microbiological results (intracellular organisms or quantitative cultures) could be shown when anatomical areas, distant from the histological foci of the pneumonia, were explored and also explains why blind techniques, as other investigators have pointed out, may have similar results to bronchoscopic techniques. The presence of bronchiolitis could also partly explain the false positive results, although the incidence of bronchiolitis in our biopsies was small (9%). The correlation between quantitative cultures of BAL fluid and the percentage of intracellular organisms (r = 0.45) indicates that the detection of intracellular organisms is only partly dependent (20%) on the lung bacterial burden. A further limitation is that, as an experimental study, the detection of intracellular organisms is restricted to BAL fluid from patients with ventilator associated pneumonia. Whether this technique may be extended to other respiratory samples such as endotracheal aspirates has not been investigated, although it seems that the most adequate sample for processing is the BAL fluid. Finally, a potential pitfall of this study could be the inclusion of gold standard biopsy samples with histological pneumonia and negative cultures. This only occurred in 20% of lung samples containing histological evidence of pneumonia, although our results did not alter when these samples were excluded from the analysis.

Overall, quantitative cultures of C-BAL yielded both a poor sensitivity and specificity, irrespective of the gold standard (bronchoscopically guided or multiple blind samples) used. This reinforces previously published work by our group using a single necropsy bronchoscopically guided open lung biopsy specimen as a gold standard. Quantitative cultures of P-BAL were less sensitive and much more specific than those obtained with C-BAL, confirming the low value of bronchoscopically guided open lung biopsy samples for the diagnosis of ventilator associated pneumonia.

Meduri et al. found that P-BAL cultures had a high sensitivity and specificity for diagnosing ventilator associated pneumonia. However, our results showed a much lower sensitivity. This can be explained by differences in the gold standard used (histological results in our study), the type of study population (immediately deceased patients in our case), prior antibiotic treatment (only 48 hours without antibiotic treatment in those cases considered not previously treated), exclusion of non-pathogenic microorganisms, and the level of contamination with P-BAL fluid.

Meduri et al. have shown that Gram stains of P-BAL fluids were positive in all the cases of pneumonia, thus allowing for early and accurate diagnosis of lower respiratory tract...
infections before the results of cultures were available. Other investigators \(^1\), \(^2\), \(^3\) have reported that the percentage of neutrophils in the lavage fluid is greater in cases of pneumonia, yet no differences in the percentage of macrophages or lymphocytes were shown.\(^4\) Differences among studies have to be explained by differences in the underlying diseases of the study populations.

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