Microbial aetiology of healthcare associated pneumonia in Spain: a prospective, multicentre, case–control study

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ABSTRACT

Introduction Healthcare-associated pneumonia (HCAP) is actually considered a subgroup of hospital-acquired pneumonia due to the reported high risk of multidrug-resistant pathogens in the USA. Therefore, current American Thoracic Society/Infectious Diseases Society of America guidelines suggest a nosocomial antibiotic treatment for HCAP. Unfortunately, the scientific evidence supporting this is contradictory.

Methods We conducted a prospective multicentre case–control study in Spain, comparing clinical presentation, outcomes and microbial aetiology of HCAP and community-acquired pneumonia (CAP) patients matched by age (±10 years), gender and period of admission (±10 weeks).

Results 476 patients (238 cases, 238 controls) were recruited for 2 years from June 2008. HCAP cases showed significantly more comorbidities (including dysphagia), higher frequency of previous antibiotic use in the preceding month, higher pneumonia severity score and worse clinical status (Charlon and Barthel scores). While microbial aetiology did not differ between the two groups (HCAP and CAP: Streptococcus pneumoniae: 51% vs 55%; viruses: 22% vs 12%; Legionella: 4% vs 9%; Gram-negative bacilli: 5% vs 4%; Pseudomonas aeruginosa: 4% vs 1%), HCAP patients showed worse mortality rates (1-month: HCAP, 12%; CAP 5%; 1-year: HCAP, 24%; CAP, 9%), length of hospital stay (9 vs 7 days), 1-month treatment failure (5.5% vs 1.5%) and readmission rate (18% vs 11%) (p<0.05, each).

Conclusions Despite a similar clinical presentation, HCAP was more severe due to patients’ conditions (comorbidities) and showed worse clinical outcomes. Microbial aetiology of HCAP did not differ from CAP indicating that it is not related to increased mortality and in Spain most HCAP patients do not need nosocomial antibiotic coverage.

Key messages

What is the key question?
◆ This multicentre case–control study investigates the microbial aetiology of HCAP showing that MDR infections are infrequent in Europe and that a nosocomial antibiotic coverage is not justified in these patients.

What is the bottom line?
◆ The poor outcomes observed in HCAP are clearly a consequence of the patients’ poor clinical condition (comorbidities).

Why read on?
◆ There results possibly indicate the need for a closer medical follow-up rather than the use of broad-spectrum antibiotics.

INTRODUCTION

The current American Thoracic Society/Infectious Diseases Society of America (ATS/IDSA) guidelines for nosocomial pneumonia (2005) describe a new clinical entity, healthcare-associated pneumonia (HCAP), which includes a number of heterogeneous conditions possibly associated with a high risk of multidrug-resistant (MDR) infections and mortality.1

This new clinical entity (HCAP) indirectly reflects substantial progress in healthcare (day hospital, outpatient clinics, etc) and profound demographic (life expectancy) and sociocultural changes in the general population.2–4 Indeed, the amount of long-term care facility patients has rapidly increased in recent years, pneumonia being one of the most frequent causes of hospitalisation and mortality.2–4 Different studies have shown that the incidence, clinical presentation and mortality for pneumonia in HCAP patients are similar to hospital-acquired pneumonia.1–5–7 Similar outcomes have been described for patients fulfilling other HCAP criteria such as patients receiving haemodialysis, previous antibiotic therapy, or immunocompromised patients.5–11 Unfortunately, there is no general consensus about the microbial aetiology of HCAP and its influence on outcomes. Whereas in the USA and Asia a high prevalence of MDR pneumonia (ie, Gram-negative bacilli or
Staphylococcus aureus) has been described, in Europe the HCAP concept is progressively weakening due to limited evidence of ‘nosocomial pathogens’.12–22

Significant differences in local healthcare organisation and in population composition could be used to explain such microbiological disparities.

Finally, the two major concerns about HCAP are: (1) the excessive heterogeneity of HCAP definition (ie, immunosuppressed patients); and (2) the inadequacy of HCAP definition to predict the risk of MDR infections.23 24

For these reasons we performed a prospective case–control investigation on the microbial aetiology of HCAP in order to confirm the reliability of the HCAP concept in Europe and the role of its microbial aetiology in determining clinical outcomes.

MATERIALS AND METHODS

Study subjects

All consecutive immunocompetent adult patients hospitalised in the 12 collaborating Spanish hospitals with a suspicion of pneumonia were considered for recruitment between June 2008 and June 2010.

Two groups of patients were studied:

1. Patients with HCAP according to ATS/IDSA criteria.1 HCAP criteria for inclusion were slightly modified in order to focus the investigation on the risk of MDR pathogens in this population: the time of ‘previous hospitalisation’ was restricted to the last 2 months (instead of three); the following criteria were added: permanent nasogastric tube; gastroscope; permanent catheters; permanent urinary catheter.

2. Selected ‘control’ community-acquired pneumonia (CAP) patients. Matching criteria were: age (±10 years), gender, and the hospitalisation period (±10 weeks) in order to respect patients’ demographics and seasonal variations in microbial aetiology of pneumonia (flu, etc).

Exclusion criteria were: all causes of immunosuppression (HIV, neoplasia, etc, excluding steroids), no written consent for study recruitment.

Witnessed bronchoaspiration pneumonia (the inhalation of oropharyngeal or gastric contents into the lower respiratory tract) was not considered for the study because chemical pneumonitis could not be ruled out.

Data collection

Data were collected on age, gender, smoking and alcohol habits (80 g/day, corresponding to approximately 11 of wine, eight standard-sized beers, or one-half pint of hard liquor each day), previous influenza and pneumococcal vaccinations, chronic comorbidities (cardiovascular, hepatic, renal, rheumatological, dementia and neurological diseases, diabetes mellitus, neoplasia in remission, etc), modified Charlson index and Barthel scale (see tables E1 and E2 in supplementary data, available online only) to assess comorbidities and physical activity, respectively, chronic therapy (inhaled and systemic corticosteroids, gastroprotective drugs and statins), previous pneumonia, previous anti-biotic therapy in the last month and aspiration evidence (dysphagia scale, see table E3 in supplementary data, available online only).

Main clinical signs and symptoms (cough, expectoration, etc) were recorded at admission, as well as analytical data (leucocyte count, C-reactive protein, etc), pneumonia severity index (PSI) and CURB-65 scores (confusion, blood urea, respiratory rate, blood pressure, age >65 years), radiological data (multilobar infiltration, etc), antimicrobial treatment (with reference to Spanish CAP guidelines) and use of systemic corticosteroids.25–27

Pulmonary (pleural effusion, empyema, atelectasis, cavitary, respiratory distress, pneumothorax and surgical pleural drainage) and extrapulmonary complications of pneumonia (septic shock, acute renal failure, endocarditis, meningitis, cardiac arrhythmias, diarrhoea, myocardial infarction, cardiac arrest, pancreatitis) were also recorded, as well as 30-day, 90-day and 1-year mortality rate (and cause of death), hospital length of stay (LOS), clinical stability, intensive care unit (ICU) admission and ventilatory support (invasive or non-invasive mechanical ventilation, MV) requirement, treatment failure and 30-day and 1-year hospital readmissions.28

Patient follow-up

Surviving patients underwent a radiological and serological follow-up at 4–6 weeks. Serial phone calls were performed every 3 months up to 1 year after admission in order to assess eventual readmissions, new antibiotic treatments, pneumonia complications or fatality cases and their causes.

Microbiological investigations

Routine sampling to determine the microbial aetiology of pneumonia included sputum, urine for Streptococcus pneumoniae and Legionella pneumophila antigens, blood cultures, nasopharyngeal swabs for respiratory viruses and for methicillin-resistant S aureus (MRSA). Pleural puncture, tracheobronchial aspirates (TBAS) and bronchoalveolar lavage, when available, were collected for Gram and Ziehl–Nielsen stains and for cultures for bacterial, fungal and mycobacterial pathogens. We only considered valid sputum samples according to current microbiological standards.29 Positive swabs for MRSA were not considered indicative of MRSA-induced pneumonia if this potential aetiology was not confirmed by a positive culture of any other respiratory sample (sputum, TBAS, etc). Serology tests for atypical pathogens were performed at admission and after 4–6 weeks (see supplementary data, available online only).

In case of failure to respond, a new microbiological investigation was conducted for bacterial, fungal and mycobacterial pathogens.

Sample processing

Sample processing techniques have been described previously.30 The following respiratory viruses were studied through a reverse transcriptase PCR technique (CLART-Pneumovir, Genomica; see supplementary data, available online only): adenovirus; bocavirus; coronavirus; enterovirus (echovirus); influenza virus A (subtypes H3N2 human, H1N1 human, B, C and H1N1/2009); metapneumovirus (subtypes A and B); para-influenza virus 1, 2, 3, and 4 (subtypes A and B); rhinovirus; respiratory syncytial virus type A (VSR-A) and B (VSR-B).

Statistical analysis

A descriptive analysis of all variables was performed, analysing qualitative variables by means of absolute and relative frequencies, while quantitative variables have been represented by means, SD and CI if normally distributed, or through medians, minimal and maximal values and IQR if they did not follow a Gaussian distribution.

Student’s t test (parametric) and the Mann–Whitney (non-parametric) test were used to assess significant differences in quantitative variables. For qualitative variables, Pearson’s χ² test, or Fisher exact test for tables (2×2) and tests of likelihood ratios for tables (m×n) were used.31 Student’s t and Wilcoxon tests were used for associations of paired data in quantitative variables. For qualitative variables McNemar tests for tables
(2 × 2), or signs tests for tables (m × n) were used. A multivariate logistic regression analysis was performed to identify factors associated with short-term and long-term mortality. First, univariate relationships were analysed, in order to identify the possible factors associated with the event (death). Second, different models were developed with significant variables. Several sets of significant variables were introduced into the model as it was not possible to use them all at once, given the large sample size, and no confounder variables were found. After adjusting several models, one of the models with best adjustment was chosen. 95% CI for estimates and SPSS V.17.0 statistical pack were used.

RESULTS

Patients’ characteristics

We recruited 476 patients (238 HCAP cases and 238 CAP controls). The main HCAP patients fulfilled one HCAP criterion (n = 207, 87%) while 9.7% (n = 23) met two criteria, 3% (n = 8) three or more criteria.

The most frequent inclusion criterion was residency in long-term care facilities (n = 119, 50%), followed by recent hospitalisation (n = 100; 42%), domiciliary wound care (n = 22; 9.2%), haemodialysis (n = 11; 4.6%), permanent urinary tube (n = 10; 4.2%), gastrostomy (n = 9; 3.8%), nasogastric tube (n = 4; 1.7%), permanent catheter (n = 2; 0.8%) and close contact with MDR infection (n = 1; 0.4%).

Most patients were hospitalised in pulmonology departments (HCAP: 77.3%; CAP: 84.5%), while fewer patients were hospitalised in internal medicine departments (HCAP: 17.6%; CAP: 11.8%) or other services (HCAP: 8.9%; CAP: 6.3%).

The main patients’ characteristics are shown in table 1. The two groups did not differ in vaccination rates (flu and pneumococcal), smoking and alcohol history, chronic use of steroids (both inhaled and systemic) or in the percentage of previous pneumonia. By contrast, HCAP patients showed a more frequent use of pump inhibitors and of antibiotics in the last month before pneumonia (table 1). The most frequent antibiotics previously administered were: fluoroquinolones in both groups (HCAP: 29 (44%) vs CAP: 11 (41%); NS), β-lactams (HCAP: 20 (30.3%) vs CAP: 8 (29.6%); NS) and macrolides (HCAP: 1 (1.5%) vs CAP: 3 (11.1%); NS).

### Table 1 Patients’ main characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Cases (HCAP)</th>
<th>Controls (CAP)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Men, %</td>
<td>68</td>
<td>68</td>
<td>NS</td>
</tr>
<tr>
<td>Age, mean±SD (years)</td>
<td>78±10.8</td>
<td>77±10.1</td>
<td>NS</td>
</tr>
<tr>
<td>Current smokers, n (%)</td>
<td>21</td>
<td>8.8</td>
<td>NS</td>
</tr>
<tr>
<td>Ex-smokers, n (%)</td>
<td>102</td>
<td>42.9%</td>
<td>NS</td>
</tr>
<tr>
<td>Alcohol intake &gt;80 g/day, n (%)</td>
<td>3</td>
<td>1%</td>
<td>NS</td>
</tr>
<tr>
<td>Flu vaccine, n (%)</td>
<td>143</td>
<td>63%</td>
<td>NS</td>
</tr>
<tr>
<td>Pneumococcal vaccine, n (%)</td>
<td>41</td>
<td>18%</td>
<td>NS</td>
</tr>
<tr>
<td>Previous pneumonia, n (%)</td>
<td>73</td>
<td>32%</td>
<td>NS</td>
</tr>
<tr>
<td>Previous antibiotic therapy (last month)</td>
<td>78</td>
<td>33%</td>
<td>0.000</td>
</tr>
<tr>
<td>Inhaled corticosteroids, n (%)</td>
<td>51</td>
<td>21%</td>
<td>NS</td>
</tr>
<tr>
<td>Systemic steroids, n (%)</td>
<td>21</td>
<td>9%</td>
<td>NS</td>
</tr>
<tr>
<td>Statins</td>
<td>52</td>
<td>22%</td>
<td>NS</td>
</tr>
<tr>
<td>Proton-pump inhibitors, n (%)</td>
<td>124</td>
<td>52%</td>
<td>0.048</td>
</tr>
<tr>
<td>Anti-H2 drugs</td>
<td>25</td>
<td>17%</td>
<td>0.017</td>
</tr>
</tbody>
</table>

CAP, community-acquired pneumonia; HCAP, healthcare-associated pneumonia.

Microbiological findings

Most patients underwent a broad microbiological investigation (see table E4 in supplementary data, available online only), with at least one positive microbiological result in 83 cases (34.9%) and 72 controls (30.3%) but two or three different isolates in 25 (10.3%) cases and 31 (13.1%) controls. Among the respiratory diseases, chronic obstructive pulmonary disease (COPD) was the most frequent in both groups, followed by bronchiectasis and asthma. As a consequence, patients with a high severity score (PSI classes IV–V 85% in HCAP vs 68% in CAP; p = 0.000) were more frequent in the HCAP group than in CAP indicating an increased mortality risk (see figure E1 in supplementary data, available online only).

The median number of comorbidities was clearly higher in HCAP due to the higher rate of neurological disorders in contrast with other diseases (respiratory, cardiac, diabetes, etc) that showed similar distribution between groups (table 2). Concordantly, in HCAP dysphagia was more frequent while body mass index decreased in comparison with CAP.

Among the respiratory diseases, chronic obstructive pulmonary disease (COPD) was the most frequent in both groups, followed by bronchiectasis and asthma. As a consequence, patients with a high severity score (PSI classes IV–V 85% in HCAP vs 68% in CAP; p = 0.000) were more frequent in the HCAP group than in CAP indicating an increased mortality risk (see figure E1 in supplementary data, available online only).

The HCAP subgroups of ‘nursing home’ (NH) and of ‘recent hospitalization’ (RH) showed a trend towards an increased incidence of respiratory diseases (COPD, NH: 67%; RH: 57%; HCAP: 30%) and of both inhaled (NH: 68%; RH: 67%; HCAP: 21%) and systemic steroids (NH: 32%; RH: 33%; HCAP: 9%).

### Table 2 Patients’ characteristics at presentation

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Cases (HCAP)</th>
<th>Controls (CAP)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMI, media±SD</td>
<td>25.6</td>
<td>4.3</td>
<td>26.8</td>
</tr>
<tr>
<td>Comorbidities ≥2, n (%)</td>
<td>158</td>
<td>75%</td>
<td>102</td>
</tr>
<tr>
<td>Respiratory disease, n (%)</td>
<td>118</td>
<td>50%</td>
<td>111</td>
</tr>
<tr>
<td>COPD</td>
<td>72</td>
<td>30%</td>
<td>61</td>
</tr>
<tr>
<td>Bronchiectasis</td>
<td>12</td>
<td>5%</td>
<td>6</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>57</td>
<td>27%</td>
<td>55</td>
</tr>
<tr>
<td>Cardiac failure</td>
<td>94</td>
<td>45%</td>
<td>99</td>
</tr>
<tr>
<td>Neurological diseases, n (%)</td>
<td>114</td>
<td>54%</td>
<td>46</td>
</tr>
<tr>
<td>Dysphagia (1–4), n (%)</td>
<td>90</td>
<td>39%</td>
<td>33</td>
</tr>
<tr>
<td>PSI, mean (SD)</td>
<td>123</td>
<td>133</td>
<td>104</td>
</tr>
<tr>
<td>PSI I–III, n (%)</td>
<td>35</td>
<td>15%</td>
<td>75</td>
</tr>
<tr>
<td>PSI IV–V, n (%)</td>
<td>198</td>
<td>85%</td>
<td>156</td>
</tr>
</tbody>
</table>

BMI, body mass index; CAP, community-acquired pneumonia; COPD, chronic obstructive pulmonary disease; HCAP, healthcare-associated pneumonia; PSI, pneumonia severity index.
microorganisms were found contemporarily in 11 cases (10.7%) and in 10 controls (10.9%) (p=0.200). 

*Streptococcus pneumoniae* was the most frequent isolated pathogen (63% vs 71%) followed by respiratory viruses (27% vs 15%), *Legionella pneumophila* (5% vs 11%) Gram-negative bacilli (including *Escherichia coli*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Proteus mirabilis*) (7% vs 6%), *Pseudomonas aeruginosa* (5% vs 1%), methicillin-sensitive *Staphylococcus aureus* (2% vs 3%) and atypical pathogens (including *Mycoplasma pneumoniae*, *Coxiella burnetii* and *Chlamydia pneumoniae*) (2% vs 4%). All isolated pathogens were similarly distributed between cases and controls but respiratory viruses showed a trend towards increased frequency among cases while *L. pneumophila* and other atypical pathogens showed a trend to be more frequent among controls. We did not observe any specific predominance among the different respiratory viruses identified in both groups (see table E5 in supplementary data, available online only). Among potential MDR pathogens *P. aeruginosa* showed only a mild non-significant trend towards a higher prevalence among cases with Gram-negative bacilli (table 4). The other Gram-negative isolates corresponded to five cases (one extended-spectrum betalactamase (ESBL) resistant *E. coli*, two non-MDR *E. coli*, one non-MDR *A. baumannii*, one non-MDR *P. mirabilis*) and four controls (one ESBL-resistant *E. coli*, three non-MDR *E. coli*).
A mixed infection was detected in 11% of the total population (n = 21), corresponding to 11 cases (11% of cases with at least one causative agent) and 10 controls (11% of controls) with a combination of two different micro-organisms (five cases and five controls with two different bacteria, two cases and four controls with a bacteria plus a virus, one case and one control with two viruses) and only three cases of three different pathogens (a bacteria plus two viruses in two cases, and three bacteria in one case) (see table E6 in supplementary data, available online only).

Most patients (n = 424; 92%) received an antibiotic coverage consistent with Spanish CAP guidelines, with a mild not significant trend towards greater adherence for CAP (antibiotic adequacy: HCAP, 89.6%; CAP, 94.3%; p = 0.085). Only 11% of the cases of pneumonia who did not receive recommended antibiotic therapy were caused by specific antibiotic allergies, while 5% of the cases were modified according to microbiological results.

### Clinical course and outcomes

The clinical outcomes of HCAP were clearly poorer than for CAP as both mean LOS (9 ± 8 days vs 7 ± 6 days, p = 0.008) and time to clinical stability (5.7 ± 5.4 days vs 4.8 ± 3.6 days, p = 0.030) were worse for HCAP, as were the rates of 30-day treatment failure (HCAP, 11 cases (5.5%) vs CAP, three cases (1.5%); p = 0.031) and of 30-day readmissions (HCAP, n = 37 (18%) vs CAP, n = 22 (11%); p = 0.030), the majority being for a respiratory cause (51% among HCAP, 55% among CAP). The rate of readmissions remained higher for HCAP than for CAP during the entire study follow-up period (3 months, 22% vs 14%, p = 0.054; 12 months, 43% vs 21%; p = 0.000), with a clear prevalence of respiratory causes (59–65%).

The mortality rate was also increased for HCAP patients at both 1-month (HCAP, n = 29 (12.2%) vs CAP, n = 11 (4.6%); p = 0.005) and 12-month follow-up (HCAP, n = 58 (24.4%) vs CAP, n = 22 (9.2%); p < 0.001).

Among patients who died during the study four cases (14%) and three controls (27%) were admitted to ICU, while six cases received MV (21%; three non-invasive MV; three invasive MV) and six controls (55%; four non-invasive MV; two invasive MV).

We analysed mortality rates across aetiological subgroups and we could only find higher 1-year mortality for S pneumoniae infection (HCAP, 10 (19.2%) vs CAP, one (2%); p = 0.008) and a slight non-significant trend towards increased 1-year mortality for respiratory viruses (HCAP six (27.3%) vs CAP two (18.2%), p = 0.68) among HCAP. Other aetiologies showed only very few fatality cases and did not show any statistical differences.

While LOS, readmission and 1-month mortality rates from NH and RH subgroups were similar to the whole HCAP population, treatment failure (12.9% vs 5.5%) and the 1-year mortality rate (27% vs 24%) showed an increasing trend in the RH group.

### Predictors of mortality

The multivariate analysis identified the following risk factors for 30-day mortality: a chronic cardiac disease (OR 9.95, 95% CI 2.84 to 34.94; p = <0.01), an increased LOS (OR 1.10) and a decreased autonomy at Barthel score (OR 0.97), a respiratory rate greater than 30 breaths per min (OR 1.16). Only the presence of an ‘alveolar’ pulmonary infiltrate at chest X-ray was a protective factor in comparison with a different pattern (interstitial or mixed pattern) (OR 0.16) (see table E7 in supplementary data, available online only).

The risk factors associated with 1-year mortality were: HCAP (OR 3.15), increasing age (+5 years, OR 1.29), prolonged ICU stay (+1 day, OR 1.26), increased respiratory rate (>30 per

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**Table 3 Clinical presentation of HCAP and CAP patients**

<table>
<thead>
<tr>
<th></th>
<th>Cases (HCAP)</th>
<th>Controls (CAP)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fever, n (%)</td>
<td>161 (68)</td>
<td>168 (71)</td>
<td>NS</td>
</tr>
<tr>
<td>Hypothermia, n (%)</td>
<td>16 (7)</td>
<td>4 (2)</td>
<td>0.005</td>
</tr>
<tr>
<td>Chills, n (%)</td>
<td>66 (28)</td>
<td>92 (39)</td>
<td>0.016</td>
</tr>
<tr>
<td>Cough, n (%)</td>
<td>161 (68)</td>
<td>176 (74)</td>
<td>NS</td>
</tr>
<tr>
<td>Purulent phlegm, n (%)</td>
<td>67 (31)</td>
<td>73 (31)</td>
<td>NS</td>
</tr>
<tr>
<td>Digestive symptoms, n (%)</td>
<td>38 (16)</td>
<td>25 (11)</td>
<td>NS</td>
</tr>
<tr>
<td>Dyspnoea, n (%)</td>
<td>168 (71)</td>
<td>155 (65)</td>
<td>NS</td>
</tr>
<tr>
<td>Altered mental status, n (%)</td>
<td>63 (27)</td>
<td>34 (14)</td>
<td>0.001</td>
</tr>
<tr>
<td>PaO2 &lt;65 mm Hg</td>
<td>146 (72)</td>
<td>122 (66)</td>
<td>NS</td>
</tr>
<tr>
<td>Bacteremia</td>
<td>24 (10.1)</td>
<td>19 (8.0)</td>
<td>NS</td>
</tr>
<tr>
<td>Multilobar infiltration, n (%)</td>
<td>57 (24)</td>
<td>47 (20)</td>
<td>NS</td>
</tr>
<tr>
<td>ICU admission, n (%)</td>
<td>10 (4)</td>
<td>11 (5)</td>
<td>NS</td>
</tr>
<tr>
<td>NIV, n (%)</td>
<td>10 (4)</td>
<td>12 (5)</td>
<td>NS</td>
</tr>
<tr>
<td>MV, n (%)</td>
<td>7 (3)</td>
<td>5 (2)</td>
<td>NS</td>
</tr>
<tr>
<td>Respiratory complications</td>
<td>37 (15.5)</td>
<td>38 (16.0)</td>
<td>NS</td>
</tr>
<tr>
<td>Distress</td>
<td>7 (2.9)</td>
<td>7 (3.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Empyema</td>
<td>2 (0.8)</td>
<td>4 (1.5)</td>
<td>NS</td>
</tr>
<tr>
<td>Pleural effusion, n (%)</td>
<td>26 (10.9)</td>
<td>26 (10.8)</td>
<td>NS</td>
</tr>
<tr>
<td>Cavitation</td>
<td>2 (0.8)</td>
<td>1 (0.5)</td>
<td>NS</td>
</tr>
<tr>
<td>Non-respiratory complications</td>
<td>92 (38.6)</td>
<td>62 (26)</td>
<td>0.005</td>
</tr>
<tr>
<td>Renal failure</td>
<td>33 (13.8)</td>
<td>20 (8.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Septic shock</td>
<td>11 (4.6)</td>
<td>6 (2.5)</td>
<td>NS</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>11 (4.6)</td>
<td>7 (2.9)</td>
<td>1.00</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>9 (3.8)</td>
<td>6 (2.5)</td>
<td>0.785</td>
</tr>
<tr>
<td>Others</td>
<td>28 (11.8)</td>
<td>23 (9.7)</td>
<td>0.72</td>
</tr>
</tbody>
</table>

CAP, community-acquired pneumonia; HCAP, healthcare-associated pneumonia; ICU, intensive care unit; MV, mechanical ventilation; NIV, non-invasive ventilation; PaO2, arterial oxygen tension.

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**Table 4 Microbiological aetiology of pneumonia among patients with known aetiology**

<table>
<thead>
<tr>
<th></th>
<th>Cases (HCAP)</th>
<th>Controls (CAP)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streptococcus pneumoniae</td>
<td>52 (62.7)</td>
<td>51 (70.8)</td>
<td>0.282</td>
</tr>
<tr>
<td>Respiratory Viruses</td>
<td>22 (26.5)</td>
<td>11 (15.3)</td>
<td>0.116</td>
</tr>
<tr>
<td>Legionella pneumophila</td>
<td>4 (4.8)</td>
<td>8 (11.1)</td>
<td>0.227</td>
</tr>
<tr>
<td>Gram-negative bacilli*</td>
<td>5 (7.2)</td>
<td>4 (5.6)</td>
<td>1.000</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
<td>4 (4.8)</td>
<td>1 (1.4)</td>
<td>0.373</td>
</tr>
<tr>
<td>MSSA</td>
<td>2 (2.4)</td>
<td>2 (2.8)</td>
<td>1.000</td>
</tr>
<tr>
<td>Atypical pathogens</td>
<td>2 (2.4)</td>
<td>3 (4.2)</td>
<td>0.664</td>
</tr>
<tr>
<td>Haemophilus influenza</td>
<td>1 (1.2)</td>
<td>1 (1.4)</td>
<td>1.000</td>
</tr>
<tr>
<td>Moraxella catarrhalis</td>
<td>1 (1.2)</td>
<td>0 (0.0)</td>
<td>1.000</td>
</tr>
<tr>
<td>Total number</td>
<td>83 (72)</td>
<td>72</td>
<td></td>
</tr>
</tbody>
</table>

*Gram-negative bacilli include: Escherichia coli, Klebsiella pneumoniae, Acinetobacter baumannii, Proteus mirabilis. Only two of them were multidrug resistant (one case and one control of ESBL-resistant E. coli).

CAP, community-acquired pneumonia; ESBL, extended-spectrum beta-lactamase; HCAP, healthcare-associated pneumonia; Respiratory viruses include: respiratory syncytial virus, rhinovirus, adenovirus, influenza virus A and B, parainfluenza virus I–IV, metapneumovirus A and B. Atypical pathogens include: Mycoplasma pneumoniae, Chlamydia pneumonia, Coxiella burnetii.
min; OR 2.16). Only the presence of pleuritic pain was a protective factor (OR 0.40) (see table E8 in supplementary data, available online only).

We also adjusted the mortality rates for multivariate analysis for all relevant comorbidities (heart failure, neurological, diabetes, chronic respiratory disease, liver and renal diseases) to assess their role but no one of them individually achieved the statistical significance to enter the multivariate analysis. Nevertheless, the HCAP definition was associated with a twofold increased risk of mortality (OR 2.2, 95% CI 1.21 to 4.07; p=0.010). On the other hand, the Barthel index for autonomy was a protective factor for mortality (OR 0.99, 95% CI 0.98 to 0.99; p<0.01).

DISCUSSION
The main findings of this study are:
1. HCAP patients showed worse functional status than CAP patients despite similar age and clinical presentation.
2. Concordantly to initial PSI, HCAP clinical outcomes were clearly worse than for CAP; a low Barthel score was associated with increased 1-month mortality.
3. The microbiological investigation showed similar patterns in both groups indicating that microbial aetiology is not the leading cause of increased mortality from HCAP in Spain.

HCAP patients clearly differ from CAP patients despite similar age and gender due to the poorer functional status, as reflected by the number of comorbidities and level of autonomy for daily activities. In particular, HCAP patients showed a higher frequency of neurological disorders and dysphagia (although no anaerobic pathogens were found), particularly among NH patients. Moreover, a higher proportion of HCAP patients had recently received antibiotics before an episode of pneumonia and chronically use gastric acidity inhibitors, a recognised risk factor for pneumonia.32–34 NH and RH subgroups showed an increased frequency of COPD and, in parallel, of steroid use (both systemic and inhaled), a recognised risk factor for pneumonia.

The clinical presentation was very similar between the two study groups with the only exception being hypothermia, which was more frequent among HCAP and chills, which was more frequent among CAP indicating globally that a clinical distinction between the two groups is almost impossible at medical examination. Nevertheless, the PSI score was in general higher for HCAP patients, reflecting the relevant weight of comorbidities on initial pneumonia severity. Interestingly, the rate of septic shock of HCAP patients was slightly higher than the rate of ICU suggesting indirectly some treatment restriction for these patients. In general, a trend towards limited ICU admission and MV use was observed among HCAP patients, possibly reflecting a tendency towards a limited use of life-supporting measures or poorer patient conditions.

The main clinical outcomes clearly reflected the PSI estimation because HCAP showed increased mortality in both the short and long term, such as worse LOS, readmissions, etc.

The multivariate analyses showed that HCAP is more clearly associated with long-term mortality (1 year) rather than with short term (1 month), suggesting a predominant role of poor functional status on main outcomes. As a confirmation, the Barthel score for functional status is more strictly correlated with mortality (multivariate analysis) than comorbidities globally (Charlson) and individually, suggesting that a low Barthel score should indicate a more careful patient follow-up after discharge. Indeed, different authors have already described the prognostic relevance of a functional status assessment for mortality in CAP patients and, possibly, a simple assessment for mobility autonomy could help in follow-up decisions.35 36

Finally, the microbiological investigation demonstrated that despite the increased severity at admission and the higher use of antibiotics before the pneumonia episode, a known risk factor for MDR infection, HCAP patients showed an aetiologic pattern not far from the typical CAP microbiology.

Nevertheless, a few interesting findings were observed: the role of viral infections is relevant among HCAP and is probably also related to bacterial entrance;37 nonetheless, it is worth remembering that the PCR technique detects genetic material instead of live microbes; the rates of Gram-negative bacilli and P aeruginosa infections were similar between HCAP and CAP, discrediting the hypothesis of a ‘nosocomial microbial’ pattern for HCAP in Spain and justifying the general Spanish trend to follow CAP guidelines for antibiotic treatment. Indeed, the choice of modified criteria for HCAP definition for this study was taken to ensure a wider selection of potential MDR risky conditions (by reducing the time since previous hospitalisation to 2 months and by adding other potential sources of MDR infections currently and surprisingly not considered by guidelines). Nevertheless, few cases of MDR infections were detected among both HCAP and CAP groups, suggesting that specific risk factors for MDR infections should be systematically investigated in the future to improve the antibiotic coverage and possibly reduce the related excess mortality. Shorr et al24 described an association between potential MDR pathogens and specific risk factors (ICU, etc). Indeed, in our series the few HCAP and CAP cases of potential MDR pathogens made this correlation with MDR risk factors not evaluable.24 A lower frequency of L pneumophila was observed among HCAP possibly due to the limited mobility and reduced contagious risk of these patients.

Another important aspect of the study is the relatively high proportion of oral colonisation by MRSA, especially among HCAP (HCAP 16%, CAP 6%), that in association with the increased risk of broncho-aspiration, could create the alert of MRSA pneumonia in already compromised patients. Unfortunately, due to the bad clinical conditions of the main patients it was possible to perform only a minimal number of TBAS and bronchoalveolar lavage that would have considerably increased the rate of microbiological diagnosis and investigated the role of oral ‘colonisers’ in causing pneumonia through the aspiration of oropharyngeal secretions. Moreover, the percentage of microbiological findings is relatively low despite the investigators’ efforts, due to the limited ability of many patients to expectorate.

By contrast, the main strengths of the study are: this is the first prospective multicentre case–control study conducted in Europe on an HCAP population; a wide microbiological investigation was systematically performed in all centres using identical methods; we systematically recorded patients’ functional status and the long-term clinical outcomes (1 year).

We can therefore conclude that HCAP according to the ATS/IDSA definition, clearly corresponds to patients with poor functional status and with multiple comorbidities, who despite having a similar clinical presentation and similar microbial aetiology show worse clinical outcomes in both the short and long term. These findings perfectly match with other recent European studies (UK, Germany, Spain) and corroborate the idea that the usual CAP antibiotic coverage is still indicated in these patients in Europe.39 22 38 Moreover, patients with HCAP criteria possibly deserve more attention during hospitalisation...
regarding poor functional status and comorbidities that could easily decompensate as a consequence of a respiratory infection; a more prolonged HCAP follow-up after hospital discharge is needed due to the higher long-term mortality risk.

However, all our findings do not support the use of HCAP definition in Spain because the clinical approach to very elderly or multimorbid patients (follow-up) should not depend on this classification and the antibiotic choice can still follow CAP guidelines. Risk factors for potential MDR infections should always be investigated.

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**Contributors**

EP is the coordinator and main investigator of the study, directly involved in the project design, patient recruitment, data collection and analysis, and is the main author of the manuscript. The following authors were main investigators in the relative collaborating centres and directly involved in patient recruitment and data collection and reviewed and approved the final version of the manuscript: CC, JMVC, AC, IA, RZ, JA, LM, SB, FR, JR, IB and AD. MAM contributed to the project design, supervised the microbiological data of the study, and approved the final version of the study. RM contributed to the design of the project, and contributed and approved the final manuscript. AT led the study group, contributed to the design of the project, and contributed to and approved the final study; he is the guarantor of the entire study.

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**Competing interests**

None.

**Patient consent**

Obtained.

**Ethics approval**

Ethics approval was received from the ethics committees of all the local hospitals.

**Provenance and peer review**

Not commissioned; externally peer reviewed.

**REFERENCES**


Correction


The date reported under the row ‘Pleural effusion’ in table 3 of this published paper is incorrect. The correct data for that row is below:

<table>
<thead>
<tr>
<th></th>
<th>Cases (HCAP)</th>
<th>Control (CAP)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleural effusion, n (%)</td>
<td>26</td>
<td>10.9</td>
<td>NS</td>
</tr>
</tbody>
</table>

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