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Andrew F. Shorr  
*Washington Hospital Center*

Daniela E. Myers  
*Pfizer*

David B. Huang  
*Pfizer*

Brian H. Nathanson  
*OptiStatim*

Matthew F. Emons  
*Cerner Corporation*

*See next page for additional authors*

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Authors
Andrew F. Shorr, Daniela E. Myers, David B. Huang, Brian H. Nathanson, Matthew F. Emons, and Marin H. Kollef
A risk score for identifying methicillin-resistant \textit{Staphylococcus aureus} in patients presenting to the hospital with pneumonia

Andrew F Shorr\textsuperscript{1*}, Daniela E Myers\textsuperscript{2}, David B Huang\textsuperscript{3}, Brian H Nathanson\textsuperscript{4}, Matthew F Emons\textsuperscript{5} and Marin H Kollef\textsuperscript{6}

Abstract

\textbf{Background:} Methicillin-resistant \textit{Staphylococcus aureus} (MRSA) represents an important pathogen in healthcare-associated pneumonia (HCAP). The concept of HCAP, though, may not perform well as a screening test for MRSA and can lead to overuse of antibiotics. We developed a risk score to identify patients presenting to the hospital with pneumonia unlikely to have MRSA.

\textbf{Methods:} We identified patients admitted with pneumonia (Apr 2005 – Mar 2009) at 62 hospitals in the US. We only included patients with lab evidence of bacterial infection (e.g., positive respiratory secretions, blood, or pleural cultures or urinary antigen testing). We determined variables independently associated with the presence of MRSA based on logistic regression (two-thirds of cohort) and developed a risk prediction model based on these factors. We validated the model in the remaining population.

\textbf{Results:} The cohort included 5975 patients and MRSA was identified in 14%. The final risk score consisted of eight variables and a potential total score of 10. Points were assigned as follows: two for recent hospitalization or ICU admission; one each for age < 30 or > 79 years, prior IV antibiotic exposure, dementia, cerebrovascular disease, female with diabetes, or recent exposure to a nursing home/long term acute care facility/skilled nursing facility. This study shows how the prevalence of MRSA rose with increasing score after stratifying the scores into Low (0 to 1 points), Medium (2 to 5 points) and High (6 or more points) risk. When the score was 0 or 1, the prevalence of MRSA was < 10% while the prevalence of MRSA climbed to > 30% when the score was 6 or greater.

\textbf{Conclusions:} MRSA represents a cause of pneumonia presenting to the hospital. This simple risk score identifies patients at low risk for MRSA and in whom anti-MRSA therapy might be withheld.
risk factors for resistant pathogens in pneumonia presenting to the hospital suggest that grouping Gram-negative organisms such as Pseudomonas aeruginosa together with MRSA may be inappropriate [5-7]. These distinct pathogens possess different risk factors, or select risk factors for resistance may carry differential significance for the varying organisms. For example, theoretically, nursing home residence may be associated more with MRSA than P. aeruginosa [8]. Additionally, clinicians are able to either add or withhold anti-MRSA therapy without compromising the extent of the Gram-negative coverage. The decision to prescribe an anti-MRSA agent can be decoupled from the option of Gram-negative treatment. Several agents exist which possess only Gram-positive activity and are essentially employed only for their anti-MRSA properties. Thus, it seems prudent to better recognize the specific variables that are associated with an increased risk for MRSA. Understanding these patient characteristics would more precisely allow clinicians to segregate persons at risk for MRSA from those unlikely to be infected with MRSA. Furthermore, identifying those at low risk for MRSA would facilitate antibiotic stewardship by rationally limiting unnecessary anti-MRSA therapy which would help contain costs and reduce future antibiotic resistance [9]. Therefore, we conducted a retrospective analysis in order to develop a score to stratify patients presenting to the hospital with pneumonia by their risk for MRSA infection.

Methods

Study overview and patients

We conducted a multi-centre, retrospective analysis of patients admitted to the hospital with pneumonia between April 2005 and March 2009. The data are from 62 hospitals in the United States (US) via the Cerner Health Facts database (see Additional file 1). We included adults with a primary discharge diagnosis of pneumonia or a primary diagnosis of sepsis with a secondary diagnosis of pneumonia and no other source of infection by International Classification of Diseases, 9th Revision, Clinical Modification (ICD-9-CM) codes (see Additional file 2). The codes were selected based on existing published studies [10-12] in combination with clinical judgment. We excluded those younger than 18 years of age and individuals with only nosocomial pneumonia (e.g., HAP, VAP). Nosocomial pneumonia was defined as occurring within 48 hours of hospitalization in accordance with the ATS/IDSA definitions [3]. We only included persons with pneumonia and laboratory confirmation of a bacterial aetiology. Presence of a bacterial aetiology was determined based on a review of electronic health records for culture results (blood, respiratory secretions, pleural fluid) and urinary antigens for either S. pneumoniae or Legionella spp. We only reviewed cultures obtained within 48 hours of admission. The database contains only de-identified data and is HIPAA compliant. Because of this and the nature of the present analysis (review of existing records), this study was classified as exempt from Institutional Review Board (IRB) review.

Endpoints and covariates

The isolation of MRSA served as our primary endpoint. We compared those with MRSA to those infected with other bacterial organisms. Patients with MRSA isolated along with another pathogen were counted as MRSA for the purposes of this study. Polymicrobial status (more than one isolate) was captured as a separate variable in the study. The determination of methicillin-resistance was made at each individual site in accordance with local laboratory practices. We collected information regarding patient demographics, co morbidities (based on ICD-9-CM codes), severity of illness, and source of admission. Need for intensive care unit (ICU) admission represented our primary measure of disease severity. We identified patients (based on ICD-9-CM codes) with coronary artery disease (CAD), congestive heart failure (CHF), diabetes mellitus (DM), chronic obstructive pulmonary disease (COPD), stroke, dementia, or chronic kidney disease necessitating haemodialysis (HD). We calculated Charlson Co morbidity scores to determine the extent of co morbidities [13]. We additionally categorized patients as to whether they met criteria for HCAP [3]. We defined HCAP as present if any one of the following criteria were met: nursing home, long term acute care facility, or skilled nursing facility (NH/LTC/SNF) exposure within the last 90 days, recent hospitalization in the last 90 days with length of stay ≥ 2 days, prior intravenous antibiotic therapy or wound care in the last 30 days prior to admission, chronic HD, or a history of immunosuppression. Chronic HD persons were identified as those who have undergone dialysis at any setting plus a diagnosis of chronic kidney disease (based on ICD-9-CM codes) within 30 days prior to the index admission or during index admission. Based on clinical expertise, we classified immunosuppressed persons as those who had: 1) recent or current treatment with systemic corticosteroids or other immunosuppressive/chemotherapy agents; 2) a diagnosis of human immunodeficiency disease (HIV), leukaemia/lymphoma, or metastatic cancer, or 3) had undergone a solid/liquid organ transplant. All patients not fulfilling the HCAP definition were categorized as CAP.

Statistics and risk score development

For continuous variables, MRSA and non-MRSA patients were compared with either the Student’s t-test or Wilcoxon 2-sample test, as appropriate. Categorical data were analyzed with either the chi-square test or the Fisher’s exact test. We defined a p-value of <0.05 to represent statistical significance.
For risk score development and validation, we employed a split sample approach. The entire population was randomly split into a development cohort (two-thirds of patients) and a validation cohort (remaining one-third of patients). We then utilized a bootstrapping based, stepwise logistic regression procedure on the development cohort to determine variables most strongly associated with MRSA [14]. We also assessed plausible interaction terms between various co morbidities as well as co morbidities and gender in a series of models. In each stepwise regression model, a p-value of 0.05 was required for entry into the model and a p-value of 0.1 to remain in the model. The stepwise algorithm was applied to 100 bootstrapped samples in the development set and each time the variable entered the model, this was recorded. Variables that consistently entered the model were kept for further analysis. The final 8 variables that comprised the risk score were those that entered the samples most often and had face validity for clinical relevance.

In the final risk score the standardized coefficients from the logistic regression represented the weight assigned to each predictor variable. Specifically, we fully standardized the coefficients of each variable in the final logistic regression model (i.e., they were standardized on the X and Y variables). Based on the standardized coefficients, we rounded up to get integer “point values” for the score with the smallest standardized coefficients getting a point value of 1 for ease of computation at the bedside. We then validated the risk score in the validation cohort, determining the prevalence of MRSA as a function of the score. We also examined traditional model performance characteristics (e.g., the area under the receiver operating characteristic curve [AUROC]).

**Results**
The final study population (both development and validation cohorts) included 5975 patients. MRSA occurred in 14.0% (n = 837) of patients. Other commonly recovered

| Demographics | MRSA (n = 837) | No MRSA (n = 5138) | p    |
|--------------|---------------|--------------------|------|
| Age, mean ± SD, years | 69.4 ± 17.1 | 67.1 ± 17.0 | <0.001 |
| Male, % | 53.4% | 55.7% | 0.435 |
| Race | | | 0.606 |
| Caucasian, % | 80.6% | 82.1% |
| African-American, % | 14.6% | 13.4% |
| Other/Unknown, % | 4.8% | 4.5% |

| Co morbid illnesses (current or prior to index admission unless noted) | MRSA (n = 837) | No MRSA (n = 5138) | p    |
|---------------------------------------------------------------|---------------|--------------------|------|
| Hypertension, % | 56.2% | 52.8% | 0.075 |
| Diabetes mellitus, % | 28.0% | 25.4% | 0.117 |
| Coronary artery disease, % | 29.2% | 27.6% | 0.358 |
| Congestive heart failure, % | 35.0% | 26.8% | <0.001 |
| Chronic obstructive pulmonary disease, % | 50.1% | 44.7% | 0.004 |
| Malignancy (includes lymphoma, myeloma, secondary neoplasm), % | 13.7% | 14.4% | 0.433 |
| Cerebrovascular disease (any) prior to index admission, % | 10.6% | 5.0% | <0.001 |
| Ischemic stroke prior to index admission, % | 5.0% | 3.2% | 0.007 |
| Dementia, % | 9.3% | 4.6% | <0.001 |
| Chronic haemodialysis, % | 21.1% | 15.6% | <0.001 |

| Healthcare-associated risk factors | MRSA (n = 837) | No MRSA (n = 5138) | p    |
|-----------------------------------|---------------|--------------------|------|
| Nursing home/skilled nursing facility/long term acute care exposure within last 90 days, % | 25.9% | 11.4% | <0.001 |
| Hospital-based wound care within 30 days prior to index admission, % | 5.38% | 1.32% | <0.001 |
| Recent hospitalization (for ≥ 2 days and within the last 90 days), % | 38.7% | 21.7% | <0.001 |
| Prior intravenous antibiotic therapy within the last 30 days, % | 5.4% | 1.3% | <0.001 |
| Immunosuppression, % | 23.4% | 16.2% | <0.001 |

| Severity of illness | MRSA (n = 837) | No MRSA (n = 5138) | p    |
|---------------------|---------------|--------------------|------|
| Intensive care unit admission (on or before index culture), % | 22.9% | 14.9% | <0.001 |
pathogens included: *S. pneumoniae* (17.5%), *P. aeruginosa* (13.8%), methicillin-sensitive *S. aureus* (13.7%), *Escherichia coli* (6.0%), and *H. influenzae* (5.6%). As Table 1 demonstrates, in the overall sample patients with MRSA were older and suffered from more co morbidities. Every chronic condition of interest, other than malignancy, occurred more often in patients with MRSA compared to those patients without MRSA. Reflecting this, the Charlson Co morbidity score [mean, standard deviation (SD)] was higher among patients with MRSA compared to no MRSA (2.6, 2.2) versus (2.2, 2.1), p < 0.001. A significant portion of MRSA versus no MRSA patients had an ICU admission on or before their index culture (22.9 versus 14.9%, p < 0.001).

Nearly 40% of those presenting to the hospital with pneumonia met criteria for HCAP, and the prevalence of MRSA was substantially higher in HCAP. Those with HCAP were 1.7 times (95% Confidence Interval (CI): 1.5-2.0) more likely to be infected with MRSA than persons classified as CAP (Figure 1).

The final logistic regression model consisted of eight variables and included demographics, prior healthcare exposure, disease acuity, and select co morbid conditions. Table 2 summarizes the risk score for MRSA pneumonia and the designation of points. Variables more strongly linked with MRSA (based on relative risks) included a recent inpatient hospitalization for ≥ 2 days within the last 90 days and need for ICU admission (see Additional file 3). The HCAP variable was an independent predictor of MRSA but not as strong a predictor as some of its components; therefore, it ultimately was not included in the final model. The total possible score ranged from 0 to 10. Figure 2 shows how the prevalence of MRSA rose with increasing score in both the development and validation cohorts after stratifying the scores into Low (0 to 1), Medium (2 to 5) and High (≥ 6) risk. When the score was ≤ 1, the prevalence of MRSA was < 10% while the prevalence of MRSA climbed to > 30% when the score was ≥ 6. Of note, the Low Risk subgroup (score of ≤ 1) represents 57.7% of the overall sample (e.g. both development and validation cohorts). In addition, in the overall sample, 100% of the patients in the High risk group have HCAP versus 18.17% in the Low risk group, p < 0.001.

As a screening test, a score of ≤ 1 versus greater than 1 had a sensitivity and specificity of 59.1% and 60.0% respectively. The positive predictive value was low at 19.2% but the negative predictive value equaled 90.1%. The pre-test probability for MRSA was approximately 14% (e.g., the prevalence of MRSA in the overall cohort). With a ≤ 1 classification threshold, patients with a score ≤ 1 would have a post-test (posterior) probability of MRSA of 10% (a reduction in risk of approximately one-third) and the post-test probability of MRSA would be 19.4% for those with scores > 1. The AUROCs of this model in the development and validation cohorts were 0.66 (95% CI: 0.63, 0.68) and 0.64 (95% CI: 0.60, 0.67), respectively.

**Discussion**

This large retrospective analysis of microbiologically-confirmed culture positive pneumonia patients presenting to the hospital reveals that MRSA constitutes an important pathogen in this setting. Similarly, HCAP accounts for a large proportion of all pneumonia patients admitted to the hospital and is nearly as common as CAP. Even though patients with HCAP face an increased risk for MRSA, the prevalence of MRSA in HCAP was less than 20%. A risk scoring tool based on factors independently associated with the recovery of MRSA segregates patients with moderate accuracy. A low total score describes a cohort of individuals unlikely to have MRSA and in whom anti-MRSA therapy can likely be withheld.

Our data add to the growing evidence elucidating the frequency of HCAP relative to CAP. In the initial study examining the epidemiology of HCAP, HCAP accounted for approximately one-third of all pneumonias presenting to the emergency department (ED) [15]. Other reports have documented that HCAP represents between 30% and 60% of pneumonias initially evaluated in the ED [5-7], and this trend is also observed in Asia and Europe [9,16]. Our study confirms that MRSA is a common pathogen in patients presenting to the hospital with their pulmonary infections. Classically thought only to be significant in nosocomial infections, it is now evident that physicians treating those who come to the hospital with pneumonia must consider MRSA when selecting antibiotic regimens.

Originally, the notion of HCAP was developed to help with this decision making process [3]. The specific components of the HCAP definition were derived from expert opinion and were initially adopted from a single centre study in bacteraemia [3,17]. HCAP was meant to serve as a tool to help clinicians stratify individuals as to the chance that a resistant pathogen, like MRSA, was a practical issue. Our finding that there are many patients

![Figure 1 Prevalence of methicillin-resistant Staphylococcus aureus in community-acquired and healthcare-associated pneumonia.](http://www.biomedcentral.com/1471-2334/13/268)
with HCAP who are not infected with MRSA suggests that HCAP alone cannot serve as a precise tool for risk stratification. Our findings further confirm the observations of earlier researchers. For example, Schrieber and colleagues found that HCAP performed poorly as a screening test for MRSA in persons presenting to the hospital with severe pneumonia necessitating mechanical ventilation [6]. Shorr et al. similarly found that HCAP misclassified many patients both as to their risk for resistant pathogens in general, and for MRSA, specifically [5]. Thus, it appears that broad use of the HCAP criteria for classification can lead to overuse of broad spectrum antibiotics which can result in unnecessary cost and antimicrobial resistance.

The proposed risk score is novel and improves on the status quo, as it provides a means for reliably classifying quickly and easily a large cohort of patients at low risk for MRSA. Since anti-MRSA therapies can often be added or withheld without necessarily adjusting the selected Gram-negative coverage, having an effective way to limit the prescription of anti-MRSA treatments becomes crucial. The situation with respect to anti-MRSA treatment options is even more acute given their costs and potential side effects.

Prior reports have proposed alternate risk scoring schemes for determining the probability of resistant organisms in those coming to the hospital with pneumonia [5-7]. However, most of these studies have pooled resistant Gram negative organisms (e.g., *P. aeruginosa*) with MRSA [5-7]. Our effort is therefore unique in that we have examined MRSA specifically. Our results are also novel in that we identify several factors associated with MRSA that have not previously been linked to pneumonia with this pathogen. For example, extremes of age have not been previously described as related to MRSA in pneumonia. Likewise the description of a nexus between certain comorbid diseases, such as cerebrovascular disease (CVD) and dementia, with MRSA has not been noted before in evaluations of pneumonia in the ED. The mechanism of these associations with MRSA is unclear. These specific factors might actually represent surrogates for other factors that we could neither address nor measure in the dataset. Additionally, it appears that not all factors contribute equally to the risk for MRSA as a cause of pneumonia on hospital presentation. Intense exposure to the healthcare system (e.g., recent inpatient stay) seems disproportionately important as does severity of illness. Moreover, different risk scores utilized thus far in pneumonia (e.g., CURB-65) focus solely on severity of illness while our risk score specifically deals with issues of aetiology.

Our study has several strengths. It is based on a large sample from multiple hospitals across the US. This provides

| Variable                                      | Points |
|-----------------------------------------------|--------|
| Age                                           |        |
| Age < 30 years or > 79 years                  | 1      |
| Prior healthcare exposure                     |        |
| Recent hospitalization (for ≥ 2 days and within the last 90 days) | 2      |
| Nursing home/skilled nursing facility/long term acute care exposure within last 90 days | 1      |
| Prior IV antibiotic therapy within the last 30 days | 1      |
| Severity of illness                           |        |
| Intensive care unit admission (on or before index culture) | 2      |
| Co morbid illness                             |        |
| Cerebrovascular disease (any), prior to admission | 1      |
| Dementia                                      | 1      |
| Female with diabetes mellitus                 | 1      |

Possible total point score: 10

![Figure 2](attachment:figure2.png)
Conclusions
MRSA represents a cause of pneumonia presenting to the hospital. A simple risk score for identifying MRSA in patients presenting to the hospital with pneumonia identifies those at low risk for MRSA and may facilitate efforts to better direct antibiotic prescribing.

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Author details
1Pulmonary and Critical Care Medicine, Washington Hospital Center, Washington DC, USA. 2Pfizer Inc, Outcomes Research, Collegeville, USA. 3Formerly of Pfizer Inc, US Medical Affairs, Collegeville, USA. 4OptiStatim, LLC, Longmeadow, USA. 5Cerner Corporation, Research Division, Culver City, USA. 6Pulmonary and Critical Care Division, Washington University School of Medicine, St. Louis, USA.

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