Comprehensive Molecular Testing for Respiratory Pathogens in Community-Acquired Pneumonia

Naomi J. Gadsby, Clark D. Russell, Martin P. McHugh, Harriet Mark, Andrew Conway Morris, Ian F. Laurenson, Adam T. Hill, and Kate E. Templeton

1Medical Microbiology, Department of Laboratory Medicine, Royal Infirmary of Edinburgh, 2College of Medicine and Veterinary Medicine, University of Edinburgh, 3Department of Anaesthesia, University of Cambridge, and 4Respiratory Medicine, Royal Infirmary of Edinburgh, United Kingdom

(See the Editorial Commentaries by Musher on pages 824–5 and Jain and Pavia on pages 826–8.)

Background. The frequent lack of a microbiological diagnosis in community-acquired pneumonia (CAP) impairs pathogen-directed antimicrobial therapy. This study assessed the use of comprehensive multibacterial, multiviral molecular testing, including quantification, in adults hospitalized with CAP.

Methods. Clinical and laboratory data were collected for 323 adults with radiologically-confirmed CAP admitted to 2 UK tertiary care hospitals. Sputum (96%) or endotracheal aspirate (4%) specimens were cultured as per routine practice and also tested with fast multiplex real-time polymerase-chain reaction (PCR) assays for 26 respiratory bacteria and viruses. Bacterial loads were also calculated for 8 bacterial pathogens. Appropriate pathogen-directed therapy was retrospectively assessed using national guidelines adapted for local antimicrobial susceptibility patterns.

Results. Comprehensive molecular testing of single lower respiratory tract (LRT) specimens achieved pathogen detection in 87% of CAP patients compared with 39% with culture-based methods. Haemophilus influenzae and Streptococcus pneumoniae were the main agents detected, along with a wide variety of typical and atypical pathogens. Viruses were present in 30% of cases; 82% of these were codetections with bacteria. Most (85%) patients had received antimicrobials in the 72 hours before admission. Of these, 78% had a bacterial pathogen detected by PCR but only 32% were culture-positive (P < .0001). Molecular testing had the potential to enable de-escalation in number and/or spectrum of antimicrobials in 77% of patients.

Conclusions. Comprehensive molecular testing significantly improves pathogen detection in CAP, particularly in antimicrobial-exposed patients, and requires only a single LRT specimen. It also has the potential to enable early de-escalation from broad-spectrum empirical antimicrobials to pathogen-directed therapy.

Keywords. community-acquired pneumonia; bacterial load; viral; molecular testing; PCR.

Community-acquired pneumonia (CAP) is a common infectious disease with an estimated incidence of 2–11 cases per 1000 adults in the developed world and a mortality rate of 2%–14% [1–5]. Due to the range of pathogens responsible for CAP, in moderate or severe infection, broad-spectrum antimicrobial cover should be initiated before de-escalating to narrow-spectrum pathogen-directed agents once a microbiological diagnosis has been made [1, 5]. Unfortunately, de-escalation is uncommon in practice because current diagnostic methods may identify a pathogen in only 30%–40% of patients with CAP [4, 6, 7]. Recent studies have highlighted the need for more timely and sensitive microbiological diagnostic methods in CAP, particularly for bacteria and in the common scenario of antibiotic administration prior to sampling [4, 8, 9].

The development of multiplex real-time polymerase-chain reaction (PCR) assays currently enables a respiratory specimen to be rapidly screened for a wide range of viral and atypical bacterial pathogens in a small number of reactions [10–15]. However, a similar approach has not yet been adopted for typical bacterial pathogens. It is difficult to interpret PCR results for typical respiratory bacteria in nonsterile samples such as sputum due to the potential for contamination with the same organisms from oropharyngeal flora. However, accurate molecular quantification of bacterial loads may aid in distinguishing infection from contamination in a way that is analogous to the use of quantitative cultures.

Quantitative molecular assays for typical respiratory bacteria have been used previously in patients with pneumonia, but these assays lack a number of key targets, use targets that have been shown to lack specificity, or may be only semiquantitative [16–32]. However, our group recently developed a suitable assay that is capable of quantifying bacterial loads of 8 typical bacterial pathogens from lower respiratory tract (LRT) specimens [33]. This assay was extensively validated on clinical isolates, reference
strains, and clinical specimens such as sputa and bronchoalveolar lavage fluids, and all bacterial targets could be reliably quantified, even when present in mixtures of different concentrations [33].

Our aim in the study was to assess the utility of a comprehensive molecular diagnostic approach encompassing 26 respiratory bacterial and viral pathogens and including bacterial quantification in patients with CAP.

METHODS

Patients and Study Criteria
Patients presented to 2 tertiary care hospitals in Edinburgh, United Kingdom, over an 18-month period between September 2012 and February 2014. Consecutive cases with clinical and radiological evidence of CAP were identified through daily retrospective electronic review of all respiratory samples received by our microbiology laboratory. Inclusion criteria for the study included the following: adult (aged ≥18 years); LRT specimen obtained within 48 hours of admission to the hospital; new radiographic infiltrate as determined by radiologist or consultant respiratory physician; and 3 of more of the following signs or symptoms: new or worsening cough, new or worsening expectoration of sputum, hemoptysis, new or worsening dyspnea, pleuritic chest pain, fever, headache, or abnormalities on chest auscultation or percussion. Exclusion criteria included the following: bronchiectasis, cystic fibrosis, or healthcare-associated pneumonia (hospitalized in an acute care hospital for 2 or more days within 90 days of the infection or resided in a nursing home or long-term care facility) [34]. Radiology reports were reviewed for all patients, and only patients with radiological changes consistent with pneumonia were included.

Where there was any uncertainty, the images were reviewed by an experienced respiratory physician (A. T. H.). Clinical data including comorbidities, outcomes, and antibiotic prescriptions were collected retrospectively from patient hospital records using a standard pro forma as part of a clinical audit of pneumonia management (approved by the Quality Improvement Team, Royal Infirmary of Edinburgh). Drug administration prior to hospitalization, including antibiotic prescription, was obtained from the electronic primary care prescription records on admission and later verified by a pharmacist.

Microbiological Culture and Molecular Testing
LRT specimens were cultured according to national standard protocols to detect common respiratory bacteria [35]. Isolates were identified using standard biochemical methods and/or matrix-assisted laser desorption/ionization time-of-flight (Bruker, Coventry, United Kingdom). Specimens with any named bacterial species identified by culture were considered to be culture positive. Following routine culture, LRT specimens were centrifuged, and mucopurulent material was stored at −80°C for up to 15 months. As part of the study protocol, specimens and associated clinical data were collected and anonymized before molecular testing in accordance with local ethical approval (South East Scotland Scottish Academic Health Sciences Collaboration Human Annotated BioResource reference no. 10/S1402/33). Thawed specimens were homogenized by vortexing with sterile glass beads, and a 200-µL aliquot was treated with lysozyme at 37°C followed by proteinase K at 56°C for 1 hour each. Total nucleic acid was then extracted using the automated nucliSENS easyMAG (BioMérieux, Basingstoke, United Kingdom) instrument with an off-board lysis protocol, including the addition of phocine herpes virus as an internal extraction control. Extracts were stored at 4°C for up to 5 days or longer at −80°C. Fast multiplex real-time PCR was performed using a combination of multiplex assays developed or adapted in-house for the following 26 pathogens: Streptococcus pneumoniae; Haemophilus influenzae; Moraxella catarrhalis; Staphylococcus aureus; Escherichia coli; Klebsiella pneumoniae; Pseudomonas aeruginosa; Acinetobacter baumannii; Mycoplasma pneumoniae; Chlamyaphila pneumoniae; Chlamyaphila psittaci; Legionella pneumophila; Legionella spp.; influenza A; influenza B; respiratory syncytial virus; parainfluenza virus types 1–3; adenovirus; human coronaviruses 229E, HKU1, NL63, and OC43; human metapneumovirus; and rhinovirus [10–15, 33]. As previously described [33], bacterial loads for S. pneumoniae, H. influenzae, M. catarrhalis, S. aureus, E. coli, K. pneumoniae, P. aeruginosa, and A. baumannii were expressed as colony forming units (CFUs) per milliliter of purulent material and calculated from standard curves generated by target gene plasmid dilution series on each PCR run. Quality control reactions (quantitative real-time PCR for GAPDH human gene and qualitative real-time PCR for gB phocine herpes virus internal control [IC]) were carried out on all specimens [33]. Quantitative GAPDH PCR was used as an indication of the cell content of each specimen because the GAPDH gene is present in human DNA. Negative (no template) controls were included on every extraction and PCR run to check purity of reagents. Runs were accepted if the negative controls were negative and the standard curves were linear; runs that failed quality control were repeated. Qualitative results were accepted if the internal control was positive or if at least 1 other target was positive. Quantitative results were accepted if the internal control quantification cycle (Cq) value fell within the range ±1 log Cq difference to negative extraction controls. Outside of this range, quantitative results were not accepted due to the potential for partial PCR inhibition leading to inaccuracy of measurement.

Estimating Impact on Antimicrobial Prescribing
The prescribed empirical therapy for each patient was compared with what antimicrobial(s) would have been appropriate for pathogen-directed therapy, based on the molecular identification result. Appropriate pathogen-directed therapy was determined using UK national guidelines adapted for local antimicrobial susceptibility patterns (Supplementary Table 1) [1].
Comprehensive Molecular Testing in CAP • CID 2016:62 (1 April) • 819

Statistical Analyses
Categorical variables were compared using Fisher exact test or \( \chi^2 \) test. Continuous data were assessed using the Shapiro–Wiik W test for nonnormality and analyzed using the Mann–Whitney U test or t test. The McNemar test was used to compare paired proportions. The Bonferroni method was used to correct for multiple comparisons during significance testing. To control for potential confounding factors that affect associations with the outcome of infection, we used logistic regression analysis.

For the analysis of the outcome of infection, a combined outcome measure of 30-day mortality and/or intensive care unit (ICU) admission was used because of the infrequent occurrence of either event in our cohort. Data were analyzed using StatsDirect software, version 2.8.0 (Altrincham, United Kingdom).

RESULTS

Patient Characteristics
A total of 323 hospitalized adult patients with CAP were included in the study cohort; 55% were male and the median age was 67 years (Table 1). Critical care admission (ICU or high-dependency unit) was required for 18.6% of patients, with 67 years (Table 1). Chronic obstructive pulmonary disease (75.4%) and/or intensive care unit admission was used because of the infrequent occurrence of either event in our cohort. Data were analyzed using StatsDirect software, version 2.8.0 (Altrincham, United Kingdom).

Pathogen Detection by Molecular and Conventional Testing of LRT Specimens
A total of 323 LRT specimens were received from 323 patients; 310 (96.0%) were sputa and 13 (4.0%) were endotracheal aspirates. Culture-based microbiological testing identified a bacterial pathogen in 127 (39.3%) patients. All specimens were positive by PCR for the human gene target GAPDH, with an average load of \( 1.24 \times 10^5 \pm 1.18 \times 10^5 \) gene copies per reaction; all specimens met quality control criteria for qualitative molecular testing.

Bacteria were detected by molecular testing in specimens from 262 (81.1%) patients (Table 2). When a cutoff of \( \geq 10^5 \) CFU/mL was applied for assays where bacterial load was quantified, bacteria were detected in specimens from 231 (71.5%) patients. Including respiratory virus results, overall pathogen detection by molecular testing rose to 86.7% (n = 280; Table 2).

H. influenzae and S. pneumoniae were the most frequently identified bacteria in patients with CAP, detected in 130 (40.2%) and 115 (35.6%) patients, respectively. Detection of more than 1 bacterial species occurred in 102 (31.6%) patients. H. influenzae or S. pneumoniae were present in 94 (92.2%) codetections. Viruses were detected in 98 (30.3%) patients (Table 2); rhinovirus was most commonly detected (12.7%, n = 41) followed by influenza A/B (7.1%, n = 23). However, viruses alone were found in only 18 (5.6%) of our patients, with 80 (81.6%) viruses cocodetected with bacteria (Table 2). The most common combination was rhinovirus with H. influenzae and/or S. pneumoniae in 30 (37.5%) cases.

PCR detected significantly more H. influenzae, S. pneumoniae, M. catarrhalis, S. aureus, E. coli, and K. pneumoniae than standard culture-based methods (Supplementary Table 2).

Table 1. Characteristics of Included Patients With Community-Acquired Pneumonia (n = 323)

| Demographics          | N (%) |
|-----------------------|-------|
| Male (%)              | 177 (54.8) |
| Age, median (interquartile range) years | 67, 51–78 |
| Age ≥65 y (%)         | 182 (56.3) |
| Age ≥75 y (%)         | 111 (34.4) |
| Comorbidity           |       |
| Chronic obstructive pulmonary disease | 128 (39.6) |
| Ischemic heart disease or heart failure | 91 (28.2) |
| Immunosuppressiona    | 43 (13.3) |
| Diabetes mellitus     | 36 (11.1) |
| Neoplastic disease    | 31 (9.6) |
| Cerebrovascular disease | 22 (6.8) |
| Chronic kidney disease | 22 (6.8) |
| Chronic liver disease | 6 (1.9)   |
| Severity index        |       |
|CURB-65 scoreb        |       |
| 0                     | 73 (23.1) |
| 1                     | 83 (26.3) |
| 2                     | 84 (26.6) |
| 3                     | 56 (17.7) |
| 4                     | 19 (6.0)  |
| 5                     | 1 (0.3)   |
|Pneumonia severity index classc|       |
| 1                     | 46 (12.1) |
| 2                     | 9 (4.2)   |
| 3                     | 18 (8.3)  |
| 4                     | 78 (35.9) |
| 5                     | 66 (30.4) |
| Admission C-reactive protein, mg/L (interquartile range)d | 149.9 (43–246.5) |
|Antimicrobial administration |       |
|Received antimicrobials in the 72 h prior to sputum sampling| 268 (84.8) |
|Outcome               |       |
|Intensive care unit admission | 40 (12.4) |
|Intubation and ventilation | 24 (7.4) |
|Vasopressor requirement | 19 (5.9)  |
|Total 30-day mortality | 20 (6.2)  |

Abbreviation: CURB-65, confusion of new onset, blood urea nitrogen >7 mmol/l, respiratory rate ≥30 breaths per minute, blood pressure <90 mmHg systolic or ≤60 mmHg diastolic, age ≥65 years.

a Immunosuppressive drugs, human immunodeficiency virus infection, inherited immunodeficiency syndromes, hematological malignancy.

b Information available for 216 patients.

c Information available for 217 patients.

d Information available for 295 patients.

Of 127 culture-positive specimens, 125 (98.4%) were also PCR positive for the same species of bacteria; the 2 discrepant specimens were H. influenzae culture positive but culture negative by PCR. Significantly, PCR was able to detect bacteria in 143 culture-negative specimens. Specimens from 27 (8.4%) patients grew bacteria that were not included in our PCR assays (Supplementary Table 3).

Bacterial Load
Bacterial load quantifications from 29 (9.1%) specimens were excluded from analysis due to the detection of partial inhibition.
by the IC assay. A higher bacterial load was detected by PCR in culture-positive specimens in comparison to culture-negative specimens. The mean combined bacterial load was $8.77 \times 10^8$ CFU/mL in culture-positive vs $7.03 \times 10^7$ CFU/mL in culture-negative specimens ($P < .0001$). Significant differences between culture-positive and culture-negative specimens were also observed when *H. influenzae* ($P < .0001$) and *S. pneumoniae* ($P < .0001$) were considered individually.

**Outcomes**

Neither culture-positivity nor PCR-positivity was associated with the outcome of infection ($P = .2$; odds ratio [OR] = 1.5; 95% confidence interval [CI], 0.9–2.9; and $P = .8$; OR = 0.9; 95% CI, 0.4–2.6, respectively). Polymicrobial detection was also not associated with the outcome of infection ($P = .4$; OR = 1.3; 95% CI, 0.7–2.5). The bacterial load of *S. pneumoniae* was associated with the outcome of infection on univariate analysis, with a higher load present in patients with 30-day mortality and/or ICU admission ($4.33 \times 10^8$ CFU/mL vs $1.29 \times 10^8$ CFU/mL; $P = .009$). However, the CURB-65 score (CURB-65; Confusion of new onset, blood Urea nitrogen $>7$ mmol/l, Respiratory rate $\geq 30$ breaths per minute, Blood pressure $<90$ mmHg systolic or $\leq 60$ mmHg diastolic, age $\geq 65$ years) was also associated with outcome on univariate analysis ($P = .01$). When logistic regression was used to correct for CURB-65 score in patients with *S. pneumoniae* infection, bacterial load was no longer significantly associated with outcome ($P = .09$; Supplementary Table 4).

**Effect of Antimicrobial Therapy Prior to Microbiological Sampling**

Of 316 patients with information available on antimicrobial administration prior to sampling, 268 (84.8%) had received antimicrobials either while in the hospital or in the community during the 72 hours before an LRT specimen was obtained. Prior antimicrobial administration was significantly associated with a patient having a culture-negative LRT specimen ($P < .0001$; OR = 9.1; 95% CI, 4.1–22.4). Of the 268 patients who had received prior antimicrobials, 77.6% (n = 208) had a bacterial pathogen detected by PCR, but only 32.1% (n = 86) were culture positive ($P < .0001$). The mean combined load of all detected bacteria per patient was significantly higher in patients who had not received antimicrobials prior to sampling ($5.40 \times 10^9$ CFU/mL) compared with those who had received antimicrobials ($2.67 \times 10^8$ CFU/mL; $P = .0001$).

**Estimation of Impact on Antimicrobial Prescribing**

Records of antimicrobial treatment during hospitalization were available for 99.1% (n = 320) of patients. Molecular testing had the potential to lead to de-escalation in number and/or spectrum of initial empirical antibiotic agents in 247 (77.2%) patients, escalation in number and/or spectrum of antibiotic agents in 19 (5.9%) patients, and no change in 54 (16.9%) patients (Table 3). The majority of the potential de-escalation events were related to switching of amoxicillin-clavulanate to narrower-spectrum agents such as amoxicillin and doxycycline in cases where *S. pneumoniae* or *H. influenzae* were detected by PCR and to the removal of clarithromycin in cases where atypical bacteria were not detected by PCR.

**DISCUSSION**

This study demonstrates that the use of a comprehensive multibacterial, multiviral molecular testing approach approximately doubles pathogen detection in patients with CAP from 39.3% to 86.7%, as well as provides valuable information about individual bacterial loads. Testing can be carried out within 1 working day to enable reporting of results in a clinically relevant time-frame, requires only a single LRT specimen, and is not
negatively impacted by antibiotic administration prior to sampling.

Only limited quantitative molecular bacterial testing has been carried out in a well-defined CAP setting [21,25,28,29,31]. As our study focused on testing sputum by molecular methods, our high level of pathogen detection for typical bacteria may not be directly comparable to results from other recent studies in hospitalized adult CAP. By combining several methodologies such as serology, culture, antigen detection, and PCR on multiple sample types, etiology was determined in 38%–76% of cases [4,7,21,31,36,38,39,43]. Atypical bacteria were uncommon in our cohort (<5% patients), but it was not an epidemic period for M. pneumoniae [42]. Viruses were detected in approximately one third of patients, which is in agreement with the 13%–56% range reported in previous studies; influenza was also detected at comparable levels (<10%) [4,7,21,31,36,38,39,43]. Although we were not able to test controls, viral detections by PCR from the upper respiratory tract have been found to be rare in asymptomatic adults compared with those with CAP in a recent study [43].

One limitation of our study was that it was restricted to CAP patients who could produce a sputum specimen. Also, we did not include blood culture or urinary antigen tests. Sputum is a pragmatic, although imperfect, choice of specimen type for the microbiological investigation of CAP. Microscopy for sputum quality is not routinely carried out locally; however, only mucopurulent material was tested, and the cellular content was quantified by human GAPDH gene PCR to rule out salivary specimens. However, sputum that is expectorated from the lower airways will always be at risk of oropharyngeal flora contamination. Despite the limitations, national CAP guidelines [1, 5] recommend sputum investigation for patients admitted to the hospital with moderate to severe CAP, both to aid microbiological diagnosis in order to guide therapy as well as for surveillance. Alternative lower respiratory specimen types such as bronchoalveolar lavage fluids may be expected to be less at risk of contamination, but these require a semi-invasive procedure that is certainly not routine in intubated patients.

A key strength of our study was the availability of quantitative bacterial load outputs in addition to qualitative detection results. Quantification of bacterial DNA load may be important in distinguishing infection from oropharyngeal contamination in sputum. Most molecular work done to date has focused on S. pneumoniae, and a cutoff of $10^4$–$10^5$ gene copies/mL is typically described as a significance threshold [20,21,29,44]. A cutoff of $10^5$ CFU/mL is also generally agreed upon for sputum culture as determined through years of experience. Based on these data, we applied a significance threshold of $\geq 10^5$ CFU/mL for all bacterial loads, and this did not significantly decrease overall pathogen detection. However, it remains unclear whether a single molecular cutoff is relevant for all bacterial species and whether cutoffs for sputum culture are relevant to sputum PCR. Quantitative molecular tools for many bacterial species are new [32,33], and there is a clear need for further exploration of their role. Determination of thresholds for significant and nonsignificant detection are key targets for future work.

We found a higher mean bacterial load by PCR in culture-positive specimens in comparison to culture-negative specimens. However, the culture-negative group was more frequently antibiotic exposed, and antibiotic exposure was also associated with lower bacterial loads. As PCR is able to detect dead as well as viable bacteria, it is not clear for how long bacterial loads might be detectable after initiation of appropriate antibiotics.
in patients with CAP and if monitoring of bacterial load by PCR would be useful in these patients. With the quantitative molecular tools now available for the relevant bacteria, these issues can also be further investigated.

A recent review has noted that interventions to reduce excessive and increase effective antibiotic prescribing in hospitalized patients can have a positive impact on antimicrobial resistance, hospital-acquired infections, and clinical outcomes [45]. However, studies in the pneumonia setting have been small and either focused on atypical bacteria and viruses [46] or nonquantitative molecular detection of bacteria [30], demonstrating 11% and 67% antibiotic modification, respectively. The results of comprehensive molecular testing in our study were not available to the treating physicians due to the anonymized study protocol. However, we estimated that de-escalation and/or reduction in the number of agents might have occurred in three quarters of our CAP patients. This was based on significantly improved bacterial detection capability, high rates of broad-spectrum empirical antibiotic usage in our cohort, and knowledge of local antimicrobial resistance patterns.

Clearly, a large number of additional factors influence antimicrobial selection (eg, severity of illness, concurrent infection at sites other than the lower respiratory tract, drug allergy, antimicrobial susceptibility testing, inflammatory markers), and we do not suggest that sputum PCR testing alone will be sufficient. However, the lack of positive microbiological identification is a significant issue, and it is highly likely that enhancing the detection of pathogens and reporting of bacterial loads would have a major impact on the clinical decision-making process. Therefore, our study illustrates the feasibility of providing the physician with significantly more information on which to base treatment decisions than is currently available and suggests that comprehensive PCR testing including bacterial load quantitation should be one of the inputs to future prospective studies in this area.

Supplementary Data
Supplementary materials are available at http://cid.oxfordjournals.org. Consisting of data provided by the author to benefit the reader, the posted materials are not copyrighted and are the sole responsibility of the author, so questions or comments should be addressed to the author.

Notes
Acknowledgments. The authors thank the staff of the Royal Infirmary of Edinburgh Medical Microbiology Laboratory for their help with data and specimen collection.

Financial support. This work was supported by the Chief Scientist Office (grant number ETM/250).

Potential conflicts of interest. All authors: No reported conflicts. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

References
1. Lim WS, Baudouin SV, George RC, et al. BTS guidelines for the management of community-acquired pneumonia in adults: update 2009. Thorax 2009; 64:iuii–55.
2. Jokinen C, Heisskanen I, Juvonen H, et al. Incidence of community-acquired pneumonia in the population of four municipalities in eastern Finland. Am J Epidemiol 1993; 137:977–88.
3. Myint PK, Kwock CS, Majumdar SR, et al. The International Community-Acquired Pneumonia (CAP) Collaboration Cohort (ICCC) study: rationale, design and description of study cohorts and patients. BMJ Open 2012; 2:e0001030.
4. Jain S, Self WH, Wunderink RG, et al. Community-acquired pneumonia requiring hospitalization among U.S. adults. N Engl J Med 2015; 373:415–22.
5. Mandell LA, Wunderink RG, Anzueto A, et al. Infectious Diseases Society of America/American Thoracic Society consensus guidelines on the management of community-acquired pneumonia in adults. Clin Infect Dis 2007; 44:278–327.
6. Chalmers JD, Taylor JK, Singanayagam A, et al. Epidemiology, antibiotic therapy, and clinical outcomes in health care associated pneumonia: a UK cohort study. Clin Infect Dis 2011; 53:107–13.
7. Musher DM, Roig IL, Cazes G, Stager CE, Logan N, Safar H. Can an etiologic agent be identified in adults who are hospitalized for community-acquired pneumonia: results of a one-year study. J Infect 2013; 67:11–8.
8. Jain S, Williams DJ, Arnold SR, et al. Community-acquired pneumonia requiring hospitalization among U.S. children. N Engl J Med 2015; 372:835–45.
9. Musher DM, Thorner AR. Community-acquired pneumonia. N Engl J Med 2014; 371:1619–28.
10. Templeton KE, Scheltina SA, Graffelman AW, et al. Comparison and evaluation of real-time PCR, real-time nucleic acid sequence-based amplification, conventional PCR, and serology for diagnosis of Mycoplasma pneumoniae. J Clin Microbiol 2003; 41:4366–71.
11. Templeton KE, Scheltina SA, Sillekens P, et al. Development and clinical evaluation of an internally controlled, single-tube multiplex real-time PCR assay for detection of Legionella pneumophila and other Legionella species. J Clin Microbiol 2004; 42:1564–9.
12. Scheltina SA, Templeton KE, Beersma MF, Claas EC. Diagnosis of human metapneumovirus and rhinovirus in patients with respiratory tract infections by an internally controlled multiplex real-time RNA PCR. J Clin Virol 2005; 33:306–11.
13. Gaunt ER, Hardie A, Claas EC, Simmonds P, Templeton KE. Epidemiology and clinical presentations of the four human coronaviruses 229E, HKU1, NL63, and OC43 detected over 3 years using a novel multiplex real-time PCR method. J Clin Microbiol 2010; 48:2940–7.
14. Yang S, Lin S, Khalil A, et al. Quantitative PCR assay using sputum samples for rapid diagnosis of pneumococcal pneumonia in adult emergency department patients. J Clin Microbiol 2005; 43:3221–6.
15. Aplhalter P, Stouer B, Barouwisch W, Nehr M, Kramer L, Burgmann H. Community-acquired bacteria frequently detected by means of quantitative polymerase chain reaction in nosocomial early-onset ventilator-associated pneumonia. Crit Care Med 2005; 33:1492–9.
16. Kais M, Spindler C, Kalin M, Ortvquist A, Giske CG. Quantitative detection of Streptococcus pneumoniae, Haemophilus influenzae, and Moraxella catarrhalis in lower respiratory tract samples by real-time PCR. Diagn Microbiol Infect Dis 2006; 55:169–78.
17. Bayram A, Kocoglu E, Balci I, Filiz A, Eksi F. Real-time polymerase chain reaction assay for detection of Streptococcus pneumoniae in sputum samples from patients with community-acquired pneumonia. J Microbiol Immunol Infect 2010; 43:452–7.
18. Abdeldaim GM, Stralin K, Olcen P, Blomberg J, Herrmann B. Toward a quantitative DNA-based definition of pneumococcal pneumonia: a comparison of Streptococcus pneumoniae target genes, with special reference to the Spn9802 fragment. Diagn Microbiol Infect Dis 2008; 60:143–50.
19. Johannson N, Kalin M, Trevylyang-Lindell A, Giske CG, Hedlund J. Etiology of community-acquired pneumonia: increased microbiological yield with new diagnostic methods. Clin Infect Dis 2010; 50:202–9.
20. Rios-Licea MM, Bosques FJ, Arroliga AC, Galindo-Galindo JO, Garza Gonzalez E. Quadruple real-time quantitative PCR assay for the detection of pathogens related to late-onset ventilator-associated pneumonia: a preliminary report. J Microbiol Methods 2010; 81:232–4.
21. Ost DE, Pech D, Fadel A, Wettimuny S, Ginocchio C, Wang XP. Mini-bronchoalveolar lavage quantitative polymerase chain reaction for diagnosis of meticillin-resistant Staphylococcus aureus pneumonia. Crit Care Med 2010; 38:1536–41.
22. Feizabadi MM, Majnnooni A, Nomanpour B, et al. Direct detection of Pseudomonas aeruginosa from patients with healthcare associated pneumonia by real time PCR. Infect Genet Evol 2010; 10:1247–51.
23. Werno AM, Anderson TP, Murdoch DR. Association between pneumococcal load and disease severity in adults with pneumonia. J Med Microbiol 2012; 61:1129–35.
