Executive summary

Improving the care of patients with community-acquired pneumonia (CAP) in South Africa is particularly important because of the high burden of disease and the need to improve standards of antibiotic prescribing in the face of rising antimicrobial resistance (AMR). The purpose of this document is to provide clinicians guidance as to the recommended management of patients with CAP. This is an update for clinicians, which takes into account important advances and controversies in the management of patients with CAP.

Diagnosing CAP

Primary care

The definitive clinical diagnosis of pneumonia requires the presence of compatible symptoms and signs for <2 weeks plus a new or worsening consolidation on chest X-ray (CXR). CXR may not be available in primary care settings in which case the diagnosis can be made on clinical grounds alone.

- CAP should be diagnosed in patients in primary care who present with a combination of well-established clinical features of CAP, including vital sign and examination abnormalities (A II).

Hospital level care

In contrast to primary care, CXRs are widely available and all patients presenting to hospital with suspected CAP require a CXR to confirm the diagnosis and exclude other potential causes for their illness. Otherwise the principles of CAP diagnosis are the same as in primary care.

- A CXR should be performed in all patients presenting to hospital with suspected CAP (A II).
- In the vast majority of cases a normal CXR excludes the diagnosis of CAP; however, empiric antibiotic therapy can be considered for severely ill hospitalised patients with suspected CAP and a negative CXR study. CAP is excluded if a repeat CXR at 24–48 hours is negative (A III).

Severity of illness scores

Assessment of the severity of CAP is important since it will
determine the appropriate site of care, the extent of the microbiological work-up and the choice of initial empiric antibiotic treatment.

- The CURB-65 score (CRB-65 for outpatients) is the recommended disease severity score for patients with CAP (A II).
- Severity scoring systems should not be the sole basis for making decisions regarding site of care. Disease severity score should always be interpreted in conjunction with a thorough clinical assessment of the patient (A II).

**Site of care decisions**

Site of care decisions, such as outpatient vs. inpatient care or general ward vs. intensive care unit, are important areas for improvement of CAP care. Decisions should be based on the clinical condition of the patients, on the disease severity scoring, on the social circumstances of the patients and on available resources.

- Patients with a CRB-65 score of 0 or a CURB-65 score of 0 or 1 are at low risk of death and may be considered for treatment at home (A II).
- Patients with a CRB-65 score of 1 or 2 or a CURB-65 score of 2 are at increased risk of death, and should be referred to hospital (A II).
- Patients with a CRB-65 score or CURB-65 score of 3 or more are at high risk of death and require urgent hospital admission and even consideration for possible admission to a high-care or intensive care unit (A II).

**Additional tests**

**Blood-based biomarkers**

Blood-based biomarkers may be used to aid the diagnosis of CAP and to assist in severity assessment.

- Routine measurement of CRP or PCT when the diagnosis is not in doubt is discouraged but may be used to measure response to therapy in the critically ill (A III).
- Measurement of CRP, particularly in primary care settings and when CXR is unavailable, may aid the diagnosis of CAP (A II).
- Measurement of CRP or PCT in emergency departments may be considered in patients with acute respiratory illness when the diagnosis of CAP is in doubt (B II).
- Urea should be measured in all hospitalised patients with CAP to assist in severity scoring (A I).

**Microbiological tests**

- Blood cultures (BCs) should be taken prior to antibiotic therapy in all patients with CAP with a CURB-65 score of ≥2 (A II).
- BCs should be considered in patients with lower CURB-65 scores, but who require hospitalisation for other reasons (B II).
- BCs should not be performed on patients with CAP who are being treated as outpatients (A II).
- A sputum sample or tracheal aspirate (collected at intubation) should be submitted for Gram stain and culture for all patients with CAP with a CURB-65 score of ≥2 (A II).
- Sputum samples can be considered in patients with CURB-65 scores of <2 who require hospitalisation for reasons such as comorbidities (B II).
- Sputum samples should not be submitted on patients with CAP who are being treated as outpatients (A II).
- The use of the pneumococcal UAT is not routinely recommended for patients with CAP (B II).
- The Legionella UAT should be considered, where available, for patients with severe CAP (B II).
- The use of rapid antigen tests for influenza is not recommended (B II).
- In patients with severe CAP during the influenza season (typically June to September) nasopharyngeal samples may be considered for detection of influenza (B II).
- The routine use of molecular tests to detect additional pathogens is not recommended (B II).
- Serology for ‘atypical’ pathogens should not be routinely performed (A II).

**Investigating for tuberculosis**

TB is a cause of CAP and clinical features are not reliable in distinguishing TB from other aetiologies. However, TB should be suspected in patients presenting with CAP who are co-infected with HIV, have a subacute history and in those who initially do not respond to antibiotics. Specific investigations for TB should be performed as indicated.

- In the following high risk patient groups presenting with CAP there should be a low threshold for investigation for pulmonary TB: HIV-infected, diabetics, admission to ICU, subacute illness or those not responding to empiric antimicrobial therapy (A II).
- A GeneXpert MTB/RIF™ (Cepheid, Sunnyvale, USA) assay performed on a single expectorated or induced sputum specimen is the preferred first line
diagnostic test for pulmonary TB. Alternatively, WHO endorsed rapid molecular tests, such as line probe assays, are recommended when they are more readily available (A II).

- TB culture should be performed in the following patients with a negative Gene Xpert MTB/RIF: non-resolving pneumonia or an ongoing suspicion of TB (A III).
- When sputum is unavailable Determine™ TB-LAM Ag (Alere, Waltham, MA, USA) testing should be performed in HIV-infected patients with CD4 counts <100 cells/µl or stage 3 or 4 disease who present with CAP (A I).

Investigating for pneumocystis pneumonia

PCP typically presents in immunocompromised patients as a subacute illness with constitutional symptoms and dry cough, and is characterised by bilateral infiltrates on CXR.

- The WHO clinical case definition should be used to clinically diagnose PCP (B III).
- Diagnostic testing of HIV-infected patients who fit the WHO case definition or in whom PCP is suspected on clinical grounds depends on local availability of tests and may include an immunofluorescent assay (IFA), direct fluorescent antibody test (DFAT) or PCR (B III).
- The preferred specimen for diagnostic tests for PCP is bronchoalveolar lavage fluid (BAL) although induced or expectorated sputum may be used when bronchoscopy is unavailable (B II).
- There is limited evidence to support for the use of beta-glucan to diagnose PCP in a South African setting. Its use is only recommended as part of a clinical registry or trial (A III).

Initial empiric therapy

Initial empiric therapy for CAP should be guided by the setting in which the patient is being treated, their age, use of antibiotics within the previous 90 days, the presence of comorbidities (cardiovascular disease, chronic respiratory disease, chronic renal failure, diabetes mellitus and HIV infection) and drug intolerance. Empiric therapy for PCP and influenza may be necessary when clinical and epidemiological criteria are met. It is rarely necessary to give empiric treatment for TB unless there is a miliary pattern on CXR.

Initial antibiotic therapy

- Patients treated at home who are <65 years old, without antibiotic exposure in the past 90 days or comorbidities should receive oral high dose amoxicillin (A II).
- Patients treated at home who are <65 years old, without antibiotic exposure in the past 90 days, or comorbidities, in the setting of low macrolide resistance, could receive an oral macrolide/azalide in the presence of severe beta-lactam allergy (A II).
- Patients treated at home who are ≥65 years old, have received antibiotics within the previous 90 days, or who have comorbidities, should receive oral amoxicillin-clavulanate or an oral second generation cephalosporin (A II).
- Patients whose admission to hospital is precipitated by advanced age, personal or family preference, inadequate home care or adverse social circumstances who have non-severe pneumonia, can be treated with oral antibiotics as described above (A II).
- Patients requiring admission to hospital who are <65 years old, without antibiotic exposure in the past 90 days, or comorbidities, should receive intravenous ampicillin or penicillin (if IVI ampicillin not available) (A II).
- Patients requiring admission to hospital who are ≥65 years old, have received antibiotics within the previous 90 days, or who have comorbidities, should receive intravenous amoxicillin-clavulanate, or cefuroxime or a third-generation cephalosporin (ceftriaxone or cefotaxime) (A II).
- Patients with severe pneumonia should receive amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin (ceftriaxone or cefotaxime) plus a macrolide antibiotic (A II).
- Respiratory fluoroquinolones (moxifloxacin or levofloxacin) are an alternative therapy but because of their activity against tuberculosis these agents should not be used as first line in CAP. They may be used in patients with severe beta-lactam allergy or as an alternative to beta-lactam/macrolide therapy but should be reserved for use in patients who have no alternative treatment options (A II).
- Antibiotics should be administered early, preferably within the emergency unit, to patients with confirmed CAP (A II).

Definitive therapy

When microbiological testing detects a causative organism, it may be possible to change from empiric to definitive therapy based on the drug susceptibility testing.
When a causative organism is identified by microbiological testing, antibiotics should be changed to the narrowest spectrum agent that effectively treats the organism (A II).

Ceftaroline is recommended as directed therapy based on the results of microbiological testing in cases of high level penicillin resistant (penicillin MIC ≥8 mg/L) S. pneumoniae or MRSA (A I).

Ertapenem is recommended as directed therapy based on the results of microbiological testing in cases of resistant Enterobacteriaceae such as ESBL-producing pathogens (A I).

When to add therapy for PCP and TB

Empiric therapy for PCP should be added when patients fulfill the WHO case definition and it should not be withheld based on negative immunohistochemical staining on sputum specimens (A II).

Empiric therapy for TB prior to initial testing is rarely required unless there is a miliary pattern on CXR or the patient is severely ill and TB is suspected (A III).

When to add empiric therapy for influenza

During the influenza season oseltamivir should be provided for any patient with severe pneumonia and can be stopped if PCR testing of nasopharyngeal aspirate is negative (A II).

During the influenza season oseltamivir should be provided for any patient with moderate CAP who is suspected of having influenza if they have a specific risk factor for severe disease and can be stopped if PCR testing of nasopharyngeal aspirate is negative (B II).

Adjunctive therapies

Given the significant burden of disease caused by CAP there have been many attempts to find adjunctive therapies to improve outcomes.

There is not enough evidence to recommend the routine use of statins for either prevention or treatment of CAP (A II).

The addition of a macrolide to standard beta-lactam therapy is associated with a better outcome in patients with severe CAP requiring ICU admission and while this may relate to the antimicrobial activity of macrolides, it may also be due to their anti-inflammatory, immunomodulatory effects (A II).

Use of systemic corticosteroids (e.g., methylprednisone 0.5 mg/kg/12 h or equivalent) should be considered in patients with severe CAP requiring ICU admission unless influenza or tuberculosis is likely, or there is a history of gastro-intestinal bleeding within the previous 3 months (A I).

Intravenous to oral switching (IVPOS) and duration of antibiotics

Prompt switching from intravenous to oral therapy is a cornerstone of antibiotic stewardship, as is the use of the minimum effective duration of therapy.

 Patients can switch from intravenous to oral antibiotics when they are haemodynamically stable, have a respiratory rate <25/min, temperature <37.8 °C and are able to take oral medication (A I).

For patients managed in the community, and for most patients admitted to hospital with low or moderate severity and uncomplicated pneumonia, 5–7 days of appropriate antibiotics is recommended (A II).

Treatment duration may be extended beyond 14 days for specific clinical scenarios such as Staphylococcus aureus bacteraemia (A II).

Patients with confirmed Legionella pneumonia should be treated with azithromycin for 7 days (A II).

Acute complications

A number of possible complications of CAP may occur and are recognised in patients who fail to respond to the first few days of empiric therapy or who deteriorate after an initial improvement.

Complicated para-pneumonic effusion and empyema

Repeat CXR should be performed for any patient failing to respond to the first few days of empiric therapy or who deteriorates after an initial improvement (A II).

If follow-up CXR demonstrates effusion or lung abscess, further imaging with CT or thoracic ultrasonography should be considered (B II).

Any significant amount of pleural fluid should prompt diagnostic pleurocentesis to exclude empyema (A II).

Fluid drainage by mean of an intercostal drain is necessary in all cases of complicated para-pneumonic effusion or empyema (A II).

Lung abscess

Patients diagnosed with lung abscess as a complication
of CAP should receive a prolonged course of antibiotics, usually 4–6 weeks, along with physiotherapy to effect postural drainage (B II).

**Cardiovascular events**
- Patients with CAP with well-recognised risk factors for a cardiovascular event or cases that fail to show adequate clinical recovery should be investigated for the possibility of a cardiovascular event (A III).

**Aspiration pneumonia**
The term ‘aspiration’ refers to the abnormal entry of a large inoculum of exogenous substances or endogenous secretions into the lower airways. This can cause pneumonitis or pneumonia, which has important clinical and microbiological differences from CAP.
- Acute aspiration events, particularly in the absence of systemic inflammation or impaired respiratory function, do not require antimicrobial therapy, even if associated with a new CXR infiltrate (A III).
- Antimicrobials should be considered for patients with aspiration pneumonitis and persistent or progressive signs and symptoms 48 hours after aspirating (B III).
- Aspiration pneumonia may be a more indolent process, usually occurring late after the aspiration event, and may be associated with suppurative complications. The diagnosis implies bacterial infection of the lung, and is therefore an indication for antimicrobial therapy (A II).
- Recommended empiric antibiotic therapy is amoxicillin-clavulanate; a cephalosporin plus clindamycin or metronidazole may be an acceptable alternative (B II).

**Vaccination for prevention of pneumonia**
Appropriate vaccination, typically against *Streptococcus pneumoniae* and influenza virus is a key pillar of antibiotic stewardship.
- All adults ≥50 years who are vaccine naïve should receive a single dose of PCV13 (A II).
- All adults ≥50 years who have received PPV23 should receive a single dose of PCV13 one year later (A II).
- All adults ≥65 years who are vaccine naïve should receive a single dose of PCV13 followed a year later by PPV23 (A II).
- All adults ≥65 years who have received PPV23 should receive a single dose of PCV13 at least one year later (A II).
- Younger adults (≥18 years) who are vaccine naïve with severe underlying comorbid or immunocompromising conditions including HIV infection should receive a single dose of PCV13 followed at least 2 months later by PPV23 (A II).
- Younger adults (≥18 years) who have previously received PPV23 and have severe underlying comorbid or immunocompromising conditions including HIV infection should receive a single dose of PCV13 one year later (A II).
- All women who are pregnant in the period of influenza vaccine availability (approximately March to June) should be offered influenza vaccination with IIV3 (All). Adults aged ≥65 years should be offered annual influenza vaccination with IIV3 (A I).
- All adults with specific chronic diseases (chronic pulmonary [including tuberculosis] and cardiac diseases, chronic renal diseases, diabetes mellitus and similar metabolic disorders, individuals who are immunosuppressed including HIV-infected individuals, and individuals who are morbidly obese (body mass index ≥40 kg/m²)) should be offered annual influenza vaccination with IIV3 (A II).
- All healthcare workers should be offered annual influenza vaccination with IIV3 (A II).

**Introduction**
CAP causes considerable morbidity and mortality throughout the world (1-3). Improving the care of patients with CAP in South Africa is particularly important because of the high burden of disease and the need to improve standards of antibiotic prescribing in the face of rising AMR (4).

There are a number of important international guidelines including those of the Infectious Diseases Society of America (IDSA)/American Thoracic Society (ATS) (5) and the British Thoracic Society (BTS) (6). However, South Africa represents a unique environment with a high prevalence of both HIV infection and AMR such that guidelines must be locally applicable. Groups interested in approaches to the management of CAP in South Africa include the South African Thoracic Society (SATS) and the Federation of Infectious Diseases Societies of Southern Africa (FIDSSA). This guideline, under the auspices of SATS and FIDSSA, represents an update of the SATS CAP Guideline published in 2007 (7).
Purpose and scope

The purpose of this document is to update clinicians regarding important advances and controversies in the management of patients with CAP. South Africa has a high prevalence of HIV infection, which is a risk factor for a number of pulmonary infections with overlapping presentations including tuberculosis and *Pneumocystis jirovecii* pneumonia (PCP). This guideline seeks to provide practical advice on the approach to all adult patients with acute community-acquired infection of the lung parenchyma. These guidelines do not apply to the much larger group of adults with non-pneumonic lower respiratory tract infections (LRTI), including acute bronchitis, acute exacerbations of chronic obstructive pulmonary disease or illnesses labelled as ‘chest infections’. Pneumonia in non-ambulatory residents of nursing homes and other long-term care facilities epidemiologically mirrors healthcare-associated pneumonia (HCAP) and should be treated according to institutional HCAP guidelines.

Methodology

A committee of specialists from the SATS and the FIDSSA was assembled with individuals and/or small groups being assigned to write a section of the guideline. A non-systematic literature search was undertaken to identify published evidence relevant to each section. Searches were conducted in PubMed and The Cochrane Library databases, as well as by checking reference lists of highly relevant papers and recent international CAP guidelines. Sections were collated and distributed to the entire group for comment. Disagreements were resolved by online discussion. Once all comments were addressed the committee agreed to the final draft of the guideline. Attempts have been made to align the guideline with the Department of Health Essential Medicines List, however it is aimed at both public and private sectors and therefore differs in some respects. The committee recognises that most patients with CAP are cared for by primary care, internal medicine, and emergency medicine physicians, and this guideline is therefore directed primarily at them.

Epidemiology

Influenza and pneumonia, along with tuberculosis were amongst the top five leading underlying natural causes of death in South African adults aged ≥15 years in 2013 (8). The incidence of LRTI amongst individuals aged ≥15 years in South Africa is approximately 400 per 100,000 populations with the peak incidence in individuals aged 25–64 years, likely driven by the high HIV prevalence in this age group (9). Underlying HIV infection is the most important risk condition for LRTI hospitalisation in South Africa, with an HIV prevalence of 74% among hospitalised patients aged ≥5 years (and >90% among those aged 25–44 years) documented at 4 sentinel surveillance sites in South Africa between 2009–2012 (9). HIV-infected individuals have substantially greater risk of being hospitalised with LRTI than HIV-uninfected individuals. Other important risk factors for hospitalised LRTI in adults include increasing age and underlying lung disease such as asthma or chronic obstructive airways disease. In South Africa from 2009–2012, the case fatality ratio (CFR) in adults with hospitalised LRTI was 7% with HIV-infected individuals experiencing a higher CFR (8%) than HIV-uninfected individuals (5%) (9). Other significant predictors of mortality in this setting were increasing age-group and receiving mechanical ventilation or supplemental oxygen.

Many organisms can cause pneumonia in adults. Mixed infections with multiple viral and/or bacterial infections are common. *Streptococcus pneumoniae* is the most common cause of CAP [approximately 27% of all adult CAP in the pre-pneumococcal conjugate vaccine (PCV) era] (10), although the incidence of invasive pneumococcal
pneumonia in South African adults has decreased as a result of indirect effect following the introduction of PCV into the routine infant immunisation programme in 2009 (11). Other bacterial causes of pneumonia include *Haemophilus influenzae*, *Staphylococcus aureus* and gram-negative bacilli (such as *Klebsiella pneumoniae*). Atypical bacterial causes of pneumonia are uncommon in South Africa (<2% of all adult CAP) but they have cyclical circulation with periodic increases in incidence and some (e.g., *Legionella* and *Mycoplasma* species) may cause outbreaks (12). *Bordetella pertussis* may present as CAP more atypically in older children and adults. In 2014, pertussis was identified in <5% of all patients with LRTI, however, rates of pertussis in all ages may be expected to increase following the shift from whole-cell to acellular pertussis vaccine in the routine infant immunisation schedule in 2009. Anaerobic bacteria may cause pneumonia particularly in patients at increased risk of aspiration. *Pseudomonas aeruginosa* is another uncommon bacterial cause of CAP and may be considered in individuals with severe pneumonia, especially those with underlying malignancy or cardiovascular disease and structural lung disease e.g., cystic fibrosis (13).

Influenza is the most common viral cause of CAP in adults, identified in 9% of patients. Other respiratory viruses such as respiratory syncytial virus, rhinovirus and adenovirus may also be identified (14). Co-infection is common with respiratory viruses and bacteria and some viruses may also be identified from healthy controls with no respiratory symptoms; therefore, identification of a virus should not preclude initiation of antibiotic treatment.

Importantly, between 18% to 40% of patients with CAP in South Africa may test positive for tuberculosis (15,16). The spectrum of aetiological agents in HIV-infected individuals may differ from HIV-uninfected individuals with *Streptococcus pneumoniae* and tuberculosis being more commonly identified in HIV-infected individuals (9). In addition, it is important to consider the diagnosis of PCP which has a prevalence of 22% among HIV-infected adults admitted with CAP in sub-Saharan Africa (17,18). *P. jirovecii* was identified on quantitative PCR from 18% of adult CAP patients in South Africa in 2014; however in some cases, identification on PCR may indicate colonisation (12). The risk of CAP in HIV-infected individuals, as well as the probability of identifying mixed or atypical infections, increases with decreasing CD4+T cell count (19).

With increasing globalisation and increasing contact at the animal-human interface, the risks of emerging respiratory viral infections such as avian influenza or Middle East Respiratory Syndrome Coronavirus (MERS-CoV) infections remain present (20). Clinicians should keep themselves updated regarding global trends in emerging viruses. Clusters of two or more individuals hospitalised with severe respiratory illness or respiratory deaths with no identified aetiology should be investigated.

**Diagnosing CAP**

**Primary care**

Pneumonia refers to an infection of the lower respiratory tract resulting in parenchymal lung inflammation and symptoms of an acute illness. The clinical diagnosis of pneumonia requires the presence of compatible symptoms and signs plus a new or worsening infiltrate on CXR (21). In this guideline patients with ‘suspected CAP’ will be defined as having supportive clinical features (described below) without CXR confirmation.

Although cough is the most common reason for presentation to outpatient care (22), only a minority of these patients will have CAP (6,23,24). Other LRTIs, including acute bronchitis, are most commonly caused by viruses (25,26), are associated with normal chest radiographs, and do not require antibiotic therapy (27,28). In a recent randomised controlled trial involving over 2,000 outpatients with LRTI in whom pneumonia was not suspected, the group assigned amoxicillin had no benefit in terms of duration or severity of symptoms. Side effects were significantly more common in those receiving antibiotics compared to placebo, translating into a net harmful effect (27). Similar findings have been observed in other clinical trials (29-31), emphasising the need to correctly identify those with CAP to limit overuse of antibiotics and to appropriately treat and refer those with more serious infections.

Without the benefit of CXR the diagnosis of CAP in outpatient settings is challenging and relies on clinical finding (6) and, if available, point-of-care (POC) biomarkers (32). A suggested way of diagnosing CAP in primary care when CXR is unavailable using clinical features and CRP has recently been presented (33). The accuracy of clinical criteria for diagnosing CAP is poor; a systematic review has shown that even combinations of symptoms and signs rarely increase the probability of the diagnosis by more than 50% (34). However, the absence of vital sign abnormalities is a relatively reliable way to exclude the
diagnosis (35), and one study found that patients presenting with LRTI and normal vital signs and clinical examination have a low risk of pneumonia, and that this could be used to rule out the diagnosis in 95% of cases without the use of CXR (36).

Although there is a poor correlation between clinical findings and radiologically confirmed CAP, most physicians in primary care rely on history and examination to exclude or diagnose this disease (35). The following features are most commonly associated with CAP (6,29,37,38) and should be sought in all patients presenting with acute cough to primary care:

- Symptoms of an acute LRTI—cough, pleuritic chest pain, shortness of breath, sputum production (and the absence of rhinorrhoea or sore throat) (24).
- New focal chest signs on examination—localised bronchial breathing or crackles, dullness on percussion, decreased chest expansion.
- Systemic abnormalities—temperature ≥38 °C, subjective fever or chills, sweating, tachycardia.
- No other explanation for the illness—acute aspiration, carcinoma, pulmonary embolism, pulmonary oedema, acute exacerbation of COPD.

Pneumonia is a leading cause of morbidity and mortality in the elderly (39-41), who are at increased risk for this disease because of immunosenescence and concomitant illness (42). Atypical presentations are more common (43), leading to delays in diagnosis and treatment (44). Fever and cough are frequently absent in frail older adults, in whom CAP may manifest as new or worsening confusion, deterioration in functional status or decompensation of an underlying illness (44-47). In additional to these non-specific clinical findings, the elderly more commonly have normal CXRs (48), making the diagnosis especially challenging.

**Recommendations**

- CAP should be diagnosed in patients in primary care who present with a combination of well-established clinical features of CAP, including vital sign and examination abnormalities (A II).

**Hospital level care**

The principles of CAP diagnosis in patients presenting to hospital are the same as for outpatients. In contrast to primary care, X-rays are widely available and all patients presenting to hospital with suspected CAP require a CXR to confirm the diagnosis and exclude other potential causes for their illness (5).

The accuracy of CXR for the diagnosis of pneumonia is not known but it is considered the reference standard for ruling-out the diagnosis of CAP when normal. However, even a normal CXR is not sufficient to eliminate the diagnosis in the setting of a very high pre-test probability (34). CXR is not a good test for ruling-in CAP and alternative diagnoses should be considered for patients with abnormal CXRs in the absence of compatible clinical features of CAP. Therefore, as in all infectious diseases, the diagnosis of CAP in severely ill patients depends on the likelihood (or risk) of having the disease and the finding of supportive features using a combination of history, examination and chest radiography. In severely ill hospitalised patients with suspected pneumonia and a normal CXR, it is not unreasonable to start empiric antibiotic therapy (48), which may be stopped if appropriate cultures are negative and a repeat CXR at 24–48 hours remains clear of new infiltrates (5). This is supported by a large cohort study where 7% of patients admitted with suspected CAP and an initially normal CXR developed changes consistent with CAP on a repeat CXR (48). In most circumstances, however, the administration of antibiotics for suspected CAP without CXR confirmation has been associated with inaccurate diagnosis (49) and overuse of antibiotics (50).

**Recommendations**

- A CXR should be performed in all patients presenting to hospital with suspected CAP (A II).
- In the vast majority of cases a normal CXR excludes the diagnosis of CAP; however, empiric antibiotic therapy can be considered for severely ill hospitalised patients with suspected CAP and a negative CXR study. CAP is excluded if a repeat CXR at 24–48 hours is negative (A III).

**Severity of illness scores**

A number of assessment tools have been developed to assist clinicians in assessing the severity of CAP with each having its own advantages and limitations (51). However, assessment of the severity of infection is important since it will determine the appropriate site of care, the extent of the microbiological work-up and the choice of initial, empiric antibiotic treatment. One such scoring system recommended for use in patients admitted to hospital is the CURB-65 score, which was derived from the BTS rules.
The original international derivation and validation study of the CURB-65 was published in 2003 (52), and although it has also been validated in other settings, it has not been formally validated in the South African setting, as is the case with most of these scoring systems. The CURB-65 scoring system uses 5 components namely:

- Confusion.
- Urea >7 mmol/L.
- Respiratory rate ≥30 breaths/min.
- Low blood pressure (systolic <90 mmHg and/or diastolic ≤60 mmHg).
- Age ≥65 years.

A point is assigned for each of the parameters, if present in the patient. This scoring system has been recommended for use because of its simplicity, but also because its accuracy is similar to that of more complicated scoring systems, such as the pneumonia severity index (PSI).

A variation of the CURB-65 is the CRB-65, which does not require the measurement of the serum urea, thus making it particularly suitable for outpatient use, although it may be a little less accurate. The potential accuracy of the CRB-65 score in determining the need for hospitalisation of patients with CAP in an emergency department of a teaching hospital in Johannesburg, a setting of resource constraints and high HIV infection prevalence, has been evaluated. This was a prospective observational study in a single centre (53), which, although not an interventional study, documented that the CRB-65 accurately predicted time to clinical stability and risk of death in the patients with CAP. It appeared from the investigation that if the CRB-65 were to have been implemented in that setting it would have performed well in decision-making regarding the need for hospital admission of the patients. None of the scoring systems can replace clinical assessment and important additional factors, such as socioeconomic deprivation and comorbidity, must also influence the decision on hospitalisation and treatment.

**Recommendations**

- The CURB-65 score (CRB-65 for outpatients) is the recommended disease severity score for patients with CAP (A II).
- Severity scoring systems should not be the sole basis for making decisions regarding site of care. Disease severity score should always be interpreted in conjunction with a thorough clinical assessment of the patient (A II).

**Site of care decisions**

Patients may be referred from primary care to hospital for confirmation of the diagnosis with additional tests such as CXR or because admission is warranted. For patients seen in emergency departments with confirmed CAP a decision must be made regarding the need for admission. These decisions should be based on a combination of objective severity scoring, clinical assessment and factors such as advanced age, presence of significant co-morbid conditions such as HIV infection and COPD, and social circumstances. In general, patients seen in primary care with a CRB-65 score of 0 should be considered for treatment at home and those with scores of ≥1 should be considered for referral for admission to hospital.

The decision to admit patients seen in emergency departments with confirmed CAP should be based on the CURB-65 score supplemented with a subjective clinical assessment. It is preferable to manage patients with CAP outside of hospital (54), because of the increased costs (55,56) and complication rates associated with hospital admission. Patients with a CURB-65 score of 0 and 1 are thought to have a mild infection and are potentially suitable for management at home. Patients with scores of 2 are considered moderately ill and need to be observed in hospital, at least initially. Patients with scores of 3 or more are thought to be severely ill and therefore these cases, and especially those with a score of 4 or 5, need evaluation for possible admission to a high-care or even intensive care unit.

**Recommendations**

- Patients with a CRB-65 score of 0 or a CURB-65 score of 0 or 1 are at low risk of death and may be considered for treatment at home (B II).
- Patients with a CRB-65 score of 1 or 2 or a CURB-65 score of 2 are at increased risk of death, and should be referred to hospital (A II).
- Patients with a CRB-65 or CURB-65 score of 3 or more are at high risk of death and require urgent hospital admission and even consideration for admission to a high-care or intensive care unit (A II).

**Additional tests**

**Blood markers**

CRP can be useful when the diagnosis of CAP is in doubt (57), although primary care settings are less likely to have the test.
available. One exception is point-of-care CRP machines that are available in some primary healthcare facilities, particularly in the private sector. A systematic review and meta-analysis of CRP use in patients with respiratory tract infection presenting to primary care facilities showed a significant reduction in antibiotic use with a suggestion that this may lead to slightly more hospital admissions (32).

If CAP is confirmed by clinical presentation and CXR the value of CRP is very limited as it is raised in almost all cases (58). There are conflicting data regarding its use as a prognostic marker and it is a poor discriminator between different aetiologies of CAP and between TB and CAP. CRP is a sensitive marker of response to therapy with a failure to fall by 50% at day 3 or 4 being associated with increased 30-day mortality (59). In practice, however, clinical evaluation, including serial measurement of temperature, is likely to be sufficient to identify patients requiring further investigation and/or a change in antibiotic therapy.

The use of procalcitonin to guide initiation and discontinuation of antibiotics in patients with a range of respiratory tract infections across a range of settings has shown a significant reduction in the use of antibiotics with no increase in mortality or treatment failure (60). However, sub-group analysis of patients with CAP shows little difference in the proportion of patients initially prescribed antibiotics. It is therefore not recommended that procalcitonin be measured when the diagnosis of CAP is confirmed. Procalcitonin is expensive and routine measurement to guide duration of antibiotics is not recommended.

Full blood count (FBC) is of limited diagnostic value in CAP. While a raised white cell count (WCC) implicates a bacterial aetiology this is not excluded if the count is normal. FBC is of more value when the diagnosis is in doubt and alternative diagnoses are being sought. Routine measurement of urea in patients with CAP is important as it forms part of the CURB-65 severity score. Routine measurement of liver function is not recommended.

**Recommendations**

- Routine measurement of CRP or PCT when the diagnosis is not in doubt is discouraged but may be used to measure response to therapy in the critically ill (A III).
- Measurement of CRP, particularly in primary care settings and when CXR is unavailable, may aid the diagnosis of CAP (A II).
- Measurement of CRP or PCT in emergency departments may be considered in patients with acute respiratory illness when the diagnosis of CAP is in doubt (B II).
- Urea should be measured in all hospitalised patients with CAP to assist severity scoring (A I).

**Microbiological investigations**

Microbiological investigations are performed in order to identify the aetiological agent, and are not intended to confirm the diagnosis of CAP, which is diagnosed primarily clinically and radiologically. The common microbiological investigations that would be performed in the setting of CAP are BC, sputum culture, and detection of *Legionella* and pneumococcal antigens in urine. In addition, molecular tests are becoming more commonly employed, and serology may also be used. The benefit of identifying the aetiological agent is primarily to assist in directing antimicrobial therapy. This can result either in a change to broaden the spectrum of activity for an organism not adequately treated by the initial, empiric, regimen (which would directly benefit the patient), or to change to a narrower spectrum agent if a susceptible organism is identified (which is less likely to directly benefit the patient, but is in line with antibiotic stewardship principles, and will hopefully have a broader benefit on society). Other reasons to attempt to identify aetiological agents are to identify organisms of potential public health importance (e.g., influenza) and to better understand the epidemiology of CAP—which in turn influences clinical practice. However, these investigations are associated with increased health care costs. The decision of whether or not to perform microbiological investigations in patients with CAP is sometimes debatable. As a general rule, investigations would be recommended where there is a reasonable chance of the result influencing the management of the patient.

The yield of BC in the setting of CAP ranges from about 5–14%. Guidelines for appropriate collection of BCs have been published (61), and will not be reviewed here. The likelihood of a patient with CAP having a positive BC is increased with severity of illness. While a number of studies have described clinical parameters associated with increased chance of a positive BC, due to differences in study design and clinical parameters described, different studies have identified different parameters associated with bacteraemia. These include neutrophilia (62), low WCC, raised creatinine, hypoglycaemia, temperature >38 °C (63), elevated PSI score (64); tachycardia, tachypnoea and hypotension (65), and low oxygen saturation (66).
The use of sputum to identify aetiological agents is bedevilled by the need for an adequate sputum sample. When adequate samples are collected (as assessed by the presence of neutrophils and absence of epithelial cells), the Gram stain may be positive in up to 82% of patients with bacteremic pneumococcal pneumonia, 76% of patients with bacteremic *Staphylococcus aureus* pneumonia and 78% of patients with gram-negative bacillae (67). Thus, the major benefit of a Gram stain on a well-collected sputum sample is that it may allow for the broadening of antimicrobial cover for *S. aureus* or gram-negative bacilli. However, it must be borne in mind that in the majority of patients, the microscopy will be unhelpful, with no predominant micro-organism observed. Only approximately 25–40% of patients will have an adequate sample obtained (68,69) and the yield on culture of these adequate samples can be as high as 80%. However, when assessing the sputum culture yield in all patients with CAP, it can be as low as 15% overall, since a proportion of patients are unable to produce sputum, and a proportion of samples will be unsatisfactory. As with BCs, sputum samples should only be sent for culture if the result is likely to influence patient management.

The two commercially available urine antigen tests (UATs) are for *Legionella pneumophila* serogroup 1, and for *Streptococcus pneumoniae*. The *Legionella* antigen test has a high specificity (99%), but sensitivity is not well defined. The pneumococcal antigen test has a sensitivity of 60–80%, and specificity of about 90% (70,71).

The major issue with both of these tests is whether positive results will significantly alter therapy. In addition, for the pneumococcal antigen test, the question is what incremental yield is obtained over sputum and BC. A recent meta-analysis found that the UAT may identify an additional 11% of patients with pneumococcal pneumonia, and above those identified with culture (10). However, a study from Switzerland found that the addition of the UAT for *S. pneumoniae* did not influence antibiotic choices or clinical outcomes (72). Given that empiric regimens for CAP always include agents active against *S. pneumoniae* this is not surprising. A similar argument could be made regarding the use of the *Legionella* UAT, since patients with severe CAP will be treated with an agent active against this pathogen. However, there are no readily available alternative tests to detect *Legionella*, and the knowledge may have epidemiological importance. Thus, while further research is needed to guide practice regarding narrowing of spectrum when a UAT is positive, there may be some justification for testing patients with severe CAP for *Legionella* using a UAT.

Rapid antigen detection tests (RADT) are also available for influenza. Earlier versions of these assays had relatively poor sensitivity (50–70%), and although more recent versions have improved sensitivity there have been concerns that they are still not as sensitive as molecular assays (73). There is also little evidence to date on the clinical impact of using RADT in preference to molecular assays for influenza.

The addition of molecular tests to the diagnostic armamentarium has significantly increased the proportion of patients in whom an aetiological agent can be found—up to 80% (74,75). The majority of these additional agents have been viral, although molecular tests do offer the ability to detect some of the so-called ‘atypical’ pathogens. The challenge, however, remains in determining whether the presence of a positive molecular result proves causality, since the same viruses may be found in a proportion of asymptomatic patients. In addition, for many of the viral pathogens there is no specific treatment (apart from influenza).

The exception to the above discussion is influenza. Specific treatment is available, and the detection of influenza has epidemiologic and public health implications. Molecular assays also offer the most rapid and reliable method to detect influenza at present, although rapid antigen tests may prove superior and more cost effective in future.

The major disadvantage of serological investigation for pathogens such as *Legionella*, *Chlamydophila* and *Mycoplasma* spp., is the need for acute and convalescent sera to allow for appropriate interpretation of results. Thus, the results will not impact immediate patient management, and the routine use of serology is not recommended.

**Recommendations**

- BCs should be taken prior to antibiotic therapy in all patients with CAP with a CURB-65 score of ≥2 (A II).
- BCs should be considered in patients with lower CURB-65 scores, but who require hospitalisation for other reasons (B II).
- BCs should not be performed on patients with CAP who are being treated as outpatients (A II).
- A sputum sample or tracheal aspirate (collected at intubation) should be submitted for Gram stain and culture for all patients with CAP with a CURB-65 score of ≥2 (A II).
Sputum samples can be considered in patients with CURB-65 scores <2 who require hospitalisation for reasons such as comorbidities (B II).

Sputum should not be submitted on patients with CAP who are being treated as outpatients (A II).

The use of the pneumococcal UAT is not routinely recommended for patients with CAP (B II).

The Legionella UAT should be considered, where available, for patients with severe CAP (B III).

The use of rapid antigen tests for influenza is not recommended (B II).

In patients with severe CAP during the influenza season (typically June to September) nasopharyngeal samples may be considered for detection of influenza (B II).

The routine use of molecular tests to detect additional pathogens is not recommended (B II).

Serology for ‘atypical’ pathogens should not be routinely performed (A II).

**Investigating for tuberculosis**

South Africa faces a huge burden of TB; this is driven mainly by high rates of HIV infection (over 50% of incident TB cases are HIV-infected) and a large mining workforce (76,77). The National Department of Health (NDOH) guidelines recommend annual TB screening for all individuals in South Africa; this is supported by the WHO policy of intensified case finding for TB control in high prevalence regions. TB is a cause of CAP and the clinical features are not reliable in distinguishing TB from other aetiologies. However, TB should be suspected in patients presenting with CAP who are HIV-infected or who have diabetes mellitus (78,79), in any patient admitted to ICU (80), and in those with subacute illness or those not responding to empiric antimicrobial therapy.

Traditional diagnostic tests for TB have major drawbacks; automated liquid culture systems are the gold standard for the diagnosis of TB but are expensive and require prolonged incubation while sputum smear microscopy has a much lower diagnostic yield (81), particularly in HIV-infected patients, and does not provide drug susceptibility data. In response to these limitations, the NDOH recently rolled out GeneXpert MTB/RIF® (Xpert) (Cepheid, Sunnyvale, CA, USA) as a replacement for smear microscopy to diagnose pulmonary TB in South Africa. The assay has overall pooled sensitivities of 88% to detect culture-positive TB cases, and 67% after a negative smear microscopy result (82). A major advantage of the test is that it is able to rapidly detect patients with rifampicin resistance. Sputum induction provides an adequate sample and a bacteriological diagnosis more frequently than instruction by a health-care worker but is costlier and does not result in a higher proportion of same-day diagnoses (83). Other WHO endorsed and clinically validated rapid molecular TB tests are also in use in South Africa. Certain genotypic assays, such as line probe assays, may have diagnostic accuracy that is comparable to Xpert MTB/RIF® and have the added advantage of detecting both isoniazid and rifampicin resistance as well as susceptibility to second line TB drugs (84-86).

There is a growing evidence base for urine-based TB lipoarabinomannan (LAM) testing in HIV-infected patients, and a rapid point of care lateral flow assay is now available. The test performs particularly well for HIV-infected patients with CD4 counts <100 cells/µL, and studies have shown a sensitivity of ~40% for inpatients with confirmed TB (87,88), with an ability to detect the sickest patients with advanced immunosuppression (89,90). Appropriate training on LAM test performance and systems for quality control should be implemented in facilities where the test is in use.

**Recommendations**

- In the following high risk patient groups presenting with CAP there should be a low threshold for investigation for pulmonary TB: HIV-infected, diabetics, admission to ICU, subacute illness or those not responding to empiric antimicrobial therapy (A II).

- A GeneXpert MTB/RIF® (Cepheid, Sunnyvale, USA) assay performed on a single expectorated or induced sputum specimen is the preferred first line diagnostic test for pulmonary TB. Alternatively, WHO endorsed rapid molecular tests, such as line probe assays, are recommended when they are more readily available (A II).

- TB culture should be performed in the following patients with a negative GeneXpert MTB/RIF: non-resolving pneumonia or an ongoing suspicion of TB (A III).

- When sputum is unavailable Determine™ TB-LAM Ag (Alere, Waltham, MA, USA) testing should be performed in HIV-infected patients with CD4 counts < 100 cells/µL or stage 3 or 4 disease who present with CAP (A I).

**Investigating for pneumocystis pneumonia**

PCP typically presents as a subacute illness with constitutional symptoms and dry cough, and is characterised by bilateral infiltrates on CXR, normal chest auscultation and
desaturation on pulse oximetry after minimal exertion (91). It may progress to respiratory failure and ARDS, and carries an overall case fatality of 15% (18).

The gold standard diagnostic test for PCP is immunofluorescent staining (IFA) of <i>P. jirovecii</i> organisms on bronchoalveolar lavage samples. This test requires an invasive procedure and is not widely available in South Africa, and therefore is rarely used to diagnose PCP. Immunohistochemical stains (IFA and silver stains) are most commonly requested on expectorated and induced sputum samples, but the sensitivity of these tests is poor (560%) and they are inadequate to rule out the diagnosis of PCP (92). Sensitive PCR assays (93-97), including a commercial assay (98,99), have been developed and evaluated on a variety of respiratory specimens. Unfortunately, HIV-associated PCP has not been well represented in these evaluation studies. These assays are not available for routine use in the public sector in South Africa and whilst having good sensitivity they have low specificity and are not routinely recommended.

Plasma [1,3]-β-D-glucan (beta-glucan) (100) and lactate dehydrogenase (LDH) (101,102) have been used as supportive investigations but there is very limited evidence for their efficacy in South African settings. Therefore, the use of beta-glucan is only recommended as part of a clinical registry or trial. Because of these limitations, clinical assessment remains the most common method of diagnosis of PCP in SA, and should be based on the WHO case definition (Box 2) (103).

### Recommendations

- The WHO clinical case definition should be used to clinically diagnose PCP (B III).
- Diagnostic testing of HIV-infected patients who fit the WHO case definition or in whom PCP is suspected on clinical grounds, depends on local availability of tests and may include an IFA, DFAT or PCR (B III).
- The preferred specimen for diagnostic tests for PCP is BAL although induced or expectorated sputum may be used when bronchoscopy is unavailable (B II).
- There is limited evidence to support for the use of beta-glucan to diagnose PCP in a South African setting. Its use is only recommended as part of a clinical registry or trial (A III).

### Antibiotic therapy

#### Initial antibiotics

The choice of initial antibiotics for CAP in South Africa depends on the setting in which the patient is being treated, their age, use of antibiotics within the past 90 days and the presence of comorbidities (cardiovascular disease, chronic respiratory disease, chronic renal failure, diabetes mellitus and HIV infection) and drug intolerance. A recent systematic review found there were not enough trials to compare the effects of different antibiotics for pneumonia acquired and treated in the community (104) and guidance is therefore based on expert opinion (Tables 1, 2). Figure 1 is an algorithm for the management of community-acquired pneumonia in adults in South Africa.

The reason that the presence of comorbidities was considered an important reason for modifying and broadening antibiotic treatment in the current guideline relates to a number of studies that have been undertaken documenting that they are associated with distinct aetiological patterns in patients with CAP, frequently associated with a broader spectrum of pathogens (105-108). In the study by Ruiz and colleagues (105), comorbid pulmonary disease, hepatic and nervous system illnesses as well as current smoking and alcohol abuse were associated with distinct patterns of aetiology for each of those conditions. In that study, respiratory comorbidities were defined as treatment for asthma or COPD, or presence of interstitial lung disorders, hepatic comorbidities were defined as pre-existing viral or toxic hepatopathy and nervous system illnesses were defined as symptomatic acute or chronic vascular or nonvascular encephalopathy with or without dementia. For example, in that study patients with pulmonary disorders were at a greater risk of infections with gram-negative enteric bacilli and <i>Pseudomonas aeruginosa</i>. In the study by El-Solh et al. (106), activities of daily living index, and pulmonary (defined as treatment for COPD or interstitial lung disease), endocrine (defined as diabetes mellitus) and CNS (defined as symptomatic acute or chronic vascular or nonvascular encephalopathy) comorbidities were associated with similar...
Table 1 Empiric choice of antibiotics for CAP

<table>
<thead>
<tr>
<th>Setting</th>
<th>Route</th>
<th>&lt;65 years old, no antibiotics within 90 days and no comorbidities</th>
<th>≥65 years, antibiotics with 90 days or comorbidity*</th>
<th>Alternative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outpatient</td>
<td>PO</td>
<td>Amoxicillin</td>
<td>Amoxicillin-clavulanate or a second generation cephalosporin</td>
<td>Moxifloxacin or levofloxacin</td>
</tr>
<tr>
<td>Inpatients (non-severe)</td>
<td>PO/IV</td>
<td>Ampicillin</td>
<td>Amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin</td>
<td>Moxifloxacin or levofloxacin</td>
</tr>
<tr>
<td>Inpatients (severe/ICU)</td>
<td>IV</td>
<td>Amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin plus a macrolide/azalide</td>
<td>Amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin plus a macrolide/azalide</td>
<td>Moxifloxacin or levofloxacin plus amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin</td>
</tr>
</tbody>
</table>

* cardio vascular disease, chronic respiratory disease, chronic renal failure, diabetes mellitus, HIV infection. CAP, community-acquired pneumonia.

distinct aetiological patterns. The study by Cilloniz and co-workers (107) documented that pathogens such as *S. aureus, Enterobacteriaceae, Pseudomonas aeruginosa*, and *H. Influenzae* only occurred in CAP patients with one or more comorbidities, which included one or more of the following: chronic respiratory disease, diabetes mellitus, chronic cardiovascular disease, neurologic disease, chronic liver disease, and chronic renal disease, with these comorbidities not being further defined. In addition, multiple medical comorbid conditions have been documented to be a risk factor for penicillin-resistant and other drug-resistant pneumococcal infections (108).

Patients treated at home who are <65 years old, without antibiotic exposure in the past 90 days or comorbidities should receive oral high dose amoxicillin. Patients treated at home who are ≥65 years old, have received antibiotics within the previous 90 days or who have comorbidities should receive oral amoxicillin-clavulanate or an oral second generation cephalosporin. In both groups the alternative is an oral respiratory fluoroquinolone when there is severe beta-lactam allergy.

Patients admitted to hospital should be treated with intravenous ampicillin or penicillin in the first instance unless they are ≥65 years, have recent antibiotic exposure or co-morbidities when amoxicillin-clavulanate, cefuroxime or a third generation cephalosporin (ceftriaxone or cefotaxime) should be used. An alternative is a respiratory fluoroquinolone which is equally effective give orally or intravenously (109).

There is emerging evidence in patients with more severe pneumonia of either pneumococcal or non-pneumococcal aetiology, and including critically ill cases, that combination antibiotic therapy, most commonly the addition of a macrolide agent to standard beta-lactam therapy, may be associated with a better outcome than monotherapy (110,111). Recent systematic reviews and meta-analyses of critically ill patients with CAP, comparing macrolide-based therapies with other regimens clearly indicated that macrolide use was associated with a significant reduction in mortality compared with non-macrolide containing regimens, and the benefit became even more significant with the pooling of data from studies that provided adjusted risk estimates (112,113). This mortality benefit from the use of macrolide-based combination antibiotic regimens versus other antibiotic regimens in critically ill patients was also supported by studies of survival among intubated patients with CAP (114) and among CAP patients with severe sepsis, the latter even when evaluating patients with CAP infections due to macrolide-resistant pathogens (e.g., macrolide-resistant pneumococci and gram-negative pathogens) (115). Patients with severe CAP should therefore be treated with a combination of amoxicillin-clavulanate, cefuroxime or a third generation (ceftriaxone or cefotaxime) cephalosporin plus a macrolide/azalide antibiotic. An alternative regimen for severe CAP is a respiratory fluoroquinolone, which should be combined with another agent, most commonly a beta-lactam. In adults hospitalised with CAP, antibiotic therapy initiated within 4–8 hours of hospital arrival was associated with lower adjusted short-term mortality (113).

Additional considerations

In areas with epidemiologically high rates of penicillin-resistant *S. pneumoniae* (PRSP) pharmacokinetically enhanced amoxicillin-clavulanate (2,000 mg amoxicillin-125 mg clavulanate 12-hourly) may be the preferred agent.
Table 2  Dose and route of administration for antibiotics prescribed for CAP

<table>
<thead>
<tr>
<th>Antibiotics</th>
<th>Route</th>
<th>Dose</th>
<th>Time of administration</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Penicillins</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzylpenicillin (S. pneumoniae MIC ≤0.5 mg/L)</td>
<td>IV</td>
<td>2 MU</td>
<td>6 hourly</td>
</tr>
<tr>
<td>Benzylpenicillin (S. pneumoniae MIC ≤1 mg/L)</td>
<td>IV</td>
<td>4 MU</td>
<td>6 hourly</td>
</tr>
<tr>
<td>Benzylpenicillin (S. pneumoniae MIC ≤2 mg/L)</td>
<td>IV</td>
<td>4 MU</td>
<td>4 hourly</td>
</tr>
<tr>
<td>Amoxicillin</td>
<td>PO</td>
<td>1 g</td>
<td>8 hourly</td>
</tr>
<tr>
<td>Amoxicillin-clavulanate</td>
<td>PO</td>
<td>1 g</td>
<td>8 hourly</td>
</tr>
<tr>
<td></td>
<td>PO</td>
<td>2 g SR</td>
<td>12 hourly</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>1.2 g</td>
<td>8 hourly</td>
</tr>
<tr>
<td>Amoxicillin-clavulanate-plus amoxicillin</td>
<td>PO</td>
<td>375 mg/plus 500 mg</td>
<td>8 hourly/8 hourly</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>IVI</td>
<td>1–2 g</td>
<td>6 hourly</td>
</tr>
<tr>
<td><strong>Cephalosporins</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cefuroxime axetil</td>
<td>PO</td>
<td>750 mg</td>
<td>8 hourly</td>
</tr>
<tr>
<td>Cefuroxime</td>
<td>IVI</td>
<td>1.5 g</td>
<td>8 hourly</td>
</tr>
<tr>
<td>Cefpodoxime</td>
<td>PO</td>
<td>400 mg</td>
<td>12 hourly</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>IVI</td>
<td>1–2 g</td>
<td>24 hourly</td>
</tr>
<tr>
<td>Cefotaxime</td>
<td>IVI</td>
<td>1–2 g</td>
<td>8 hourly</td>
</tr>
<tr>
<td>Ceftaroline</td>
<td>IVI</td>
<td>600 mg</td>
<td>12 hourly</td>
</tr>
<tr>
<td><strong>Fluoroquinolones</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moxifloxacin</td>
<td>PO</td>
<td>400 mg</td>
<td>Daily</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>400 mg</td>
<td>daily</td>
</tr>
<tr>
<td>Levofloxacin</td>
<td>PO</td>
<td>750 mg/500 mg</td>
<td>Daily/12 hourly</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>750 mg/500 mg</td>
<td>Daily/12 hourly</td>
</tr>
<tr>
<td><strong>Macrolides/azalides</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Erythromycin</td>
<td>PO</td>
<td>500 mg</td>
<td>6 hourly</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>1 g</td>
<td>6 hourly</td>
</tr>
<tr>
<td>Clarithromycin</td>
<td>PO</td>
<td>500 mg</td>
<td>12 hourly</td>
</tr>
<tr>
<td></td>
<td>PO</td>
<td>1 g XL</td>
<td>12 hourly</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>500 mg</td>
<td>12 hourly</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>PO</td>
<td>500 mg</td>
<td>Daily</td>
</tr>
<tr>
<td></td>
<td>IVI</td>
<td>500 mg</td>
<td>Daily</td>
</tr>
<tr>
<td><strong>Tetracyclines</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxycycline</td>
<td>PO</td>
<td>200 mg/100 mg</td>
<td>Stat (200 mg) followed by 100 mg 12 hourly</td>
</tr>
<tr>
<td><strong>Carbapenems</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ertapenem</td>
<td>IVI/IM</td>
<td>1 g</td>
<td>Daily</td>
</tr>
</tbody>
</table>

CAP, community-acquired pneumonia.
Clinical studies with this formulation have demonstrated 97.1% bacteriological success against erythromycin-resistant (erythromycin MICs >1 mg/L) isolates and the formulation has also demonstrated high bacteriological efficacy against PRSP isolates non-susceptible to amoxicillin (amoxicillin MICs >4 mg/L), with 86.7% bacteriological success (116).

While significant macrolide resistance precludes the routine use of these agents as monotherapy for many patients with CAP, in areas of low macrolide resistance, and in young, otherwise healthy adults who have not recently received a course of antibiotics these agents may be used on their own, particularly in the setting of severe beta-lactam allergy. A macrolide/azalide/tetracycline may also be used alone or in combination with another agent in cases suspected or known to be infected with so-called ‘atypical’ pathogens.

**Recommendations**

- Patients treated at home who are <65 years old, without antibiotic exposure in the past 90 days, or comorbidities, should receive oral high dose amoxicillin (A II).
- Patients treated at home who are <65 years old, without antibiotic exposure in the past 90 days, or comorbidities,
in the setting of low macrolide resistance, could receive an oral macrolide/azalide in the presence of severe beta-lactam allergy (A II).

- Patients treated at home who are ≥65 years old, have received antibiotics within the previous 90 days, or who have comorbidities, should receive oral amoxicillin-clavulanate or an oral second generation cefalosporin (A II).

- Patients whose admission to hospital is precipitated by advanced age, personal or family preference, inadequate home care or adverse social circumstances who have non-severe pneumonia, can be treated with oral antibiotics as described above (A II).

- Patients requiring admission to hospital who are <65 years old, without antibiotic exposure in the past 90 days, or comorbidities, should receive intravenous ampicillin or penicillin (if IVI ampicillin not available) (A II).

- Patients requiring admission to hospital who are ≥65 years old, have received antibiotics within the previous 90 days, or who have comorbidities, should receive intravenous amoxicillin-clavulanate, cefuroxime or a third generation cephalosporin (ceftriaxone or cefotaxime) (A II).

- Patients with severe pneumonia should receive amoxicillin-clavulanate or cefuroxime or a third generation cephalosporin (ceftriaxone or cefotaxime) plus a macrolide antibiotic (A II).

- Respiratory fluoroquinolones (moxifloxacin or levofloxacin) are an alternative therapy but because of their activity against tuberculosis these agents should not be used as first line in CAP. They may be used in patients with severe beta-lactam allergy or as an alternative to beta-lactam/macrolide therapy but should be reserved for use in patients who have no alternative treatment options (A II).

- Antibiotics should be administered early, preferably within the emergency unit, to patients with confirmed CAP (A II).

**Definitive therapy**

Although microbiological confirmation of the aetiology of CAP will only be obtained in the minority of cases, it is important that when a causative organism is identified, antibiotics are changed to the narrowest spectrum agent that effectively treats the organism. When the causative organism is resistant to initial therapy, it is necessary to use a broader spectrum agent. Although standard agents may be used in patients with high-level resistance, two agents that should not be used as empiric therapy may be used.

Ceftaroline fosamil is a new, broad-spectrum cefalosporin prodrug that exhibits bactericidal activity against gram-positive pathogens, including methicillin-resistant Staphylococcus aureus (MRSA) and *S. pneumoniae*, as well as gram-negative pathogens. Integrated analysis of two registrational studies (Ceftaroline Community Acquired Pneumonia Trial versus Ceftriaxone in Hospitalized Patients) (FOCUS 1 and FOCUS 2), that compared ceftaroline with ceftriaxone in the treatment of adult patients requiring hospitalisation for CAP, demonstrated clinical cure rates for the ceftaroline group that were numerically higher than those for the ceftriaxone group and that this agent was well tolerated, with a safety profile similar to that of ceftriaxone (117). Ceftaroline should be reserved for use in patients with microbiologically confirmed CAP due to penicillin-resistant *S. pneumoniae* (penicillin MIC ≥8 mg/L) or MRSA.

Ertapenem, a group 1 carbapenem (limited activity against non-fermentative gram-negative bacilli) has excellent *in vitro* activity (i.e., ≥90% of isolates have an ertapenem MIC less than or equal to the susceptibility breakpoint) against bacteria that, in general, are associated with CAP, such as *S. pneumoniae*, methicillin-susceptible *Staphylococcus aureus* (MSSA) and many *Haemophilus* spp., *Enterobacteriaceae* including extended-spectrum β-lactamase (ESBL) producing strains and anaerobes (118). The efficacy and safety of ertapenem, for the treatment of CAP requiring parenteral therapy, were compared with those of ceftriaxone in two registration trials and demonstrated that ertapenem, 1 g once a day, was highly effective therapy for CAP in hospitalised adults with moderate-to-severe disease (118). Ertapenem should be reserved for use in patients with microbiologically-confirmed CAP due to resistant *Enterobacteriaceae*, such as ESBL-producing pathogens.

**Recommendations**

- When a causative organism is identified by microbiological testing, antibiotics should be changed to the narrowest spectrum agent that effectively treats the organism (A II).

- Ceftaroline is recommended as directed therapy based on the results of microbiological testing in cases of penicillin-resistant *S. pneumoniae* (penicillin MIC ≥8 mg/L) or MRSA (A I).

- Ertapenem is recommended as directed therapy based on the results of microbiological testing in cases of resistant *Enterobacteriaceae* such as ESBL-producing pathogens (A I).
When to add empiric therapy for PCP and TB

The WHO case definition of PCP in patients with HIV is shown in Box 2. Patients with HIV infection who fulfill these criteria or with a positive specific test for PCP should be initiated on therapy with co-trimoxazole (20 mg/kg TMP and 100 mg/kg SMX/day in divided doses) orally or intravenously and prednisone 40 mg twice daily for 5 days followed by 40 mg daily for 5 days and then 20 mg daily for 11 days.

Recommendations

- Empiric therapy for PCP should be added when patients fulfill the WHO case definition and it should not be withheld on the basis of negative immunohistochemical staining on sputum specimens (A II).
- Empiric therapy for TB prior to initial testing is rarely required unless there is a miliary pattern on CXR or the patient is severely ill and TB is suspected (A III).

When to add empiric therapy for influenza

The influenza season in South Africa typically starts in early June and runs until around September. Up to date information is available from the National Institute for Communicable Diseases website (www.NICD.ac.za). During this period influenza should be considered in any patient with severe pneumonia, particularly if there is a history of a preceding upper respiratory tract infection and/or diffuse bilateral infiltrates on CXR (119). If influenza is suspected on these grounds patients should be initiated on oseltamivir (75 mg twice daily) (120,121) and a nasopharyngeal aspirate tested for influenza by polymerase chain reaction. Specific risk factors for severe influenza are pregnancy, immune compromise (including diabetes mellitus and HIV infection), obesity and chronic lung, cardiac, neurological disease and age >65 years (122,123).

Recommendations

- During the influenza season oseltamivir should be provided for any patient with severe pneumonia and can be stopped if PCR testing of nasopharyngeal aspirate is negative (A II).
- During the influenza season oseltamivir should be provided for any patient with moderate CAP who is suspected of having influenza if they have a specific risk factor for severe disease and can be stopped if PCR testing of nasopharyngeal aspirate is negative (B II).

Adjunctive therapies

Since the mortality of patients with CAP, particularly those who need hospitalisation, and especially those in the intensive care unit remains high, even in the presence of effective antibiotic therapy, studies have been ongoing to find effective adjunctive therapies that could be used together with antibiotics to improve the outcome (124-126). Multiple agents have been recommended or tested but results have largely been very disappointing.

Statins

While prior statin use has been shown in systematic reviews and meta-analyses to be associated with a decreased risk and/or mortality of CAP (127-130), there is not enough evidence from randomised controlled trials to recommend their routine use to either prevent CAP or to improve its mortality. The only randomised, double-blind, placebo-controlled, intervention study investigating the impact of statin use on admission of patients with CAP to hospital was not associated with a reduction in cytokine levels, nor was it associated with a reduction in time to clinical stability among the patients (131).

Corticosteroids

Several recent studies and meta-analyses, measuring different end-points, have shown definite benefits of adjunctive corticosteroids in severe hospitalised patients with CAP (132-139). The data generated from these studies and systematic reviews have recently been extensively reviewed (140). One of the most comprehensive meta-analyses was that of Siemieniuk and colleagues (138), who in addition to undertaking an extensive extraction of the literature, analysed the data for all possible benefits and potential harms, using instruments to assess risk of bias in the individual studies, as well as publication bias, and the GRADE system to evaluate the quality/certainty of the evidence. The final assessment of findings was that corticosteroid use was associated with a lower mortality (significant only in the severe CAP group), reduction in need for mechanical ventilation, the occurrence of ARDS, time to clinical stability and length of hospital stay with the evidence being of moderate or higher quality (138). There are still unanswered questions regarding corticosteroid use, including which patients with CAP are most likely to benefit, which corticosteroids to use, at what dose and for how long. However, the data from the various studies and meta-analyses
suggest that those with severe CAP requiring ICU admission, those with the highest inflammatory indices (such as a CRP level above 150 mg/L) and those with shock requiring vasopressor support are most likely to benefit (140). The recommended regimen is methylprednisone 0.5 mg/kg/12 h or equivalent for 5 days (135). While patients with diabetes (141) and those receiving long-term steroids have been included in RCTs, there are limited data in patients with HIV infection.

**Recommendations**

- There is not enough evidence to recommend the routine use of statins in CAP (A I).
- The addition of a macrolide to standard beta-lactam therapy is associated with a better outcome in patients with severe CAP requiring ICU admission and while this may relate to the antimicrobial activity of macrolides, it may also be due to their anti-inflammatory, immunomodulatory effects (A II).
- Use of systemic corticosteroids (e.g., methylprednisone 0.5mg/kg/12 h or equivalent) should be considered in patients with severe CAP requiring ICU admission unless influenza or tuberculosis is likely, or there is a history of gastro-intestinal bleeding within the previous 3 months (A I).

**Severely ill patients with CAP**

Mortality is in the region of 12% for hospitalised CAP but >30% among those admitted to the ICU (142).

Obvious reasons for referral are the need for mechanical ventilation and the presence of septic shock. Otherwise patients with a CURB-65 of ≥3 should be evaluated for ICU admission. Clinical judgment, however, is important as elderly or immunocompromised patients may warrant ICU admission even with lower scores (143).

Organisms that cause severe CAP are similar to those that cause less severe disease; *S. pneumoniae*, *Legionella* spp., *S. aureus* and *K. pneumoniae* and viruses such as influenza (especially in unvaccinated patients, asthmatics, the obese, in immunocompromised cases and in pregnancy). However, other organisms are important to consider especially in the right clinical or geographical context: influenza H5N1 and H7N3, SARS and MERS-CoV, Hantavirus, *P. jirovecii*, enteric gram-negative bacilli (elderly, aspiration), MSSA or MRSA (influenza, travel, corticosteroids, diabetes), and *M. tuberculosis*.

Potentially useful interventions include those that are relevant to any patient with severe sepsis and mechanical ventilation should be utilised to restore adequate oxygenation without causing lung injury (144,145).

Antibiotics should be administered as soon as the diagnosis is made, preferably in the emergency room. These are similar to those described above except that intravenous macrolides should be administered to all severe pneumonias, particularly in the presence of septic shock, because of their immunomodulatory effect and because coincidentally they would cover *L. pneumophila* whether or not it is initially suspected. It does not appear, however, that empiric antibiotic coverage of other atypical pathogens such as *M. pneumoniae* and *C. pneumoniae* improves survival or clinical efficacy in hospitalised patients (146). In an influenza season oseltamivir should be initiated in any patient with severe pneumonia in whom influenza is suspected and stopped once it has been excluded by PCR. In patients with influenza pneumonia, the earlier that the neuraminidase inhibitor is started the better the outcome (147).

Severe pneumonia is itself a form of primary ARDS. However, spreading infiltrates may represent secondary ARDS due to capillary leak and not antibiotic failure. Where diagnostic uncertainty exists, particularly in immunocompromised patients with bilateral infiltrates and where pneumocystis PCR is unavailable, the use of biomarkers, including CRP, PCT, proBNP and BDG (plasma [1,3]-β-D-glucan (beta-glucan)) is extremely helpful in elucidating the cause and decreasing unnecessary antibiotic prescription. Biomarkers may also help to identify antibiotic failure. A declining CRP generally indicates that the antibiotics are appropriate despite worsening radiological features (148).

**IVPOS and duration of antibiotics**

Early IVPOS of antibiotics is a central pillar of antibiotic stewardship as it reduces costs and intravenous cannula infections and encourages reduced length of hospital stay (149). Several studies have shown that it is safe to switch patients with CAP from intravenous to oral therapy when they become clinically stable (150,151). Definitions of clinical stability vary and a typical definition is given in Box 3 (151).

The precise duration of antibiotic therapy for the management of microbiologically documented and non-documented CAP is not informed by robust evidence. The duration of therapy should be determined based on the clinical response of the patient and the causative agent. When fever defervesces rapidly and there is clinical
improvement it is safe to stop beta-lactam antibiotics after 5–7 days (6).

In patients who show a slow clinical improvement or who have a confirmed aetiological agent such as *Pseudomonas aeruginosa*, *S. aureus* or gram-negative enteric organisms, it may be necessary to continue antibiotics for longer.

**Recommendations**

- Patients can switch from intravenous to oral antibiotics when they are haemodynamically stable, have respiratory rate <25/min, temperature <37.8 °C and are able to take oral medication (A I).
- For community managed, and for most patients admitted to hospital with low or moderate severity and uncomplicated pneumonia, 5–7 days of appropriate antibiotics is recommended (A II).
- Treatment duration may be extended beyond 5–7 days for specific clinical scenarios such as *Staphylococcus aureus* bacteraemia (A II).
- Patients with confirmed *Legionella* pneumonia should be treated with azithromycin for 7 days (A II).

**Acute complications of CAP**

Most cases of pneumonia resolve completely with appropriate antibiotic treatment and supportive care. However, a number of important complications of CAP may occur that require specific management. These complications should be considered whenever a patient fails to respond adequately to therapy, although some patients present later with general ill health and ongoing constitutional symptoms. The diagnosis of complications of CAP is frequently delayed and clinicians should have a low threshold for investigations.

**Complicated para-pneumonic effusion and empyema**

Para-pneumonic effusions occur in at least 40% of bacterial CAP, and are usually small (152). They are characterised by exudative chemistries and an influx of neutrophils into the pleural space. Most effusions are uncomplicated and resolve with treatment of the pneumonia. However, if bacteria invade the pleural space, a complicated para-pneumonic effusion or empyema results. Anaerobic utilisation of glucose by the neutrophils results in pleural fluid acidosis, and lysis of neutrophils increases the LDH concentration in the pleural fluid to values often more than 1,000 IU/L. Empyema develops when there is evidence of bacterial infection of the pleural liquid, as indicated by the presence of pus in the pleural space and/or the presence of bacterial organisms on Gram stain. A positive culture is not required for diagnosis, since there are several reasons why bacteria may not be cultured from an empyema, including the presence of anaerobic organisms in the pleural space that may be difficult to culture. The characteristic putrid odour of the pleural fluid is classically considered diagnostic of anaerobic infection.

In a patient with non-resolution of CAP, the demonstration of any significant amount of pleural fluid on CXR should prompt diagnostic pleurocentesis. The fluid is considered an exudate if any of the following are found and a transudate if all are absent (Light's criteria) (153):

- Ratio of pleural fluid to serum protein >0.5.
- Ratio of pleural fluid to serum LDH >0.6.
- Pleural fluid LDH > two thirds of the upper limits of normal serum value.

In a more recent systematic review, pleural fluid cholesterol greater than 55 mg/dL and pleural LDH greater than 200 U/L each had better positive and negative likelihood ratios for distinguishing exudates from transudates than did Light's criteria (154).

A low pleural fluid pH level is more predictive of complicated effusions (that require drainage) than is a low pleural fluid glucose level. In such cases, a pleural fluid pH of <7.2 indicates the need for urgent drainage of the
effusion, if pH is 7.2–7.3 it is reasonable to observe the patient while a pleural fluid pH of >7.3 suggests that the effusion may be managed with systemic antibiotics alone (155,156). Fluid drainage by mean of an intercostal drain is necessary in all cases of complicated para-pneumonic effusion or empyema. Current international guidelines strongly recommend the routine use of ultrasonography for all pleural fluid drainage procedures.

**Recommendations**

- Repeat CXR should be performed for any patient failing to respond to the first few days of empiric therapy or who deteriorates after an initial improvement (A II).
- If follow-up CXR demonstrates effusion or lung abscess, further imaging with CT or thoracic ultrasonography should be considered (B II).
- Any significant amount of pleural fluid should prompt diagnostic pleurocentesis to exclude empyema (A II).
- Fluid drainage by mean of an intercostal drain is necessary in all cases of complicated para-pneumonic effusion or empyema (A II).

**Lung abscess**

Lung abscess is defined as necrosis of the pulmonary parenchyma with cavitation. Acute lung abscess, associated with a short duration of symptoms and a rapidly-evolving chest radiograph, is a rare complication of CAP. Risk factors include immunosuppression, inappropriate antibiotic selection, and infection with *S. aureus* and *K. pneumoniae*. Acute lung abscess should be differentiated from a chronic abscess (presenting with indolent symptoms that evolve over a period of weeks or months) seen most commonly in the debilitated or alcoholic patient, and following aspiration in a patient with a reduced level of consciousness or impaired swallowing mechanisms. Infection with anaerobic bacteria, *S. aureus*, gram-negative enteric bacilli or *S. anginosus* group (previously known as *S. milleri*) in the presence of poor dental hygiene should be considered in that instance.

A lung abscess is typically diagnosed when a chest radiograph reveals a pulmonary infiltrate with a cavity; an air-fluid level is frequently present. Better anatomic definition can be achieved with a CT scan, which can distinguish a cavitating lung lesion from a pleural collection, also a complication of CAP. It may also demonstrate previously unrecognised underlying conditions such as an aspirated foreign body, a pulmonary neoplasm, or bronchial stenosis. Most patients respond to appropriate antibiotics guided by the microbiology of the precipitating episode of CAP. If patients fail to respond to antibiotics it may be necessary to insert a pigtail drain. Antimicrobial therapy with amoxicillin-clavulanate is appropriate if a chronic abscess is suspected. In this instance, sputum culture is unreliable as it is contaminated by oral flora. A prolonged course of antibiotics (up to 6 weeks) may be required, although there is no evidence on the optimum duration of antimicrobial therapy. Monitoring response with serial CRP measurements may be useful in guiding the length of therapy. Percutaneous drainage of the abscess (guided by either ultrasound or CT) can be performed diagnostically and therapeutically in non-responders, and surgery is rarely required (less than 5% of cases in most series).

**Recommendations**

- Patients diagnosed with lung abscess as a complication of CAP should receive a prolonged course of antibiotics, usually 4–6 weeks, along with physiotherapy to effect postural drainage (B II).

**Cardiovascular events**

There is an emerging awareness of the possible occurrence of cardiovascular events in patients with CAP, with one of the earliest studies of cardiac changes in CAP having been undertaken in South Africa (157). With regard to the cardiac events these may include acute myocardial infarction (AMI), new or worsening cardiac failure and acute or worsening arrhythmia, occurring either alone or in combination, and have been documented in all-cause CAP, as well as in pneumococcal CAP specifically (158-163). While these cardiovascular events are more common in the elderly and in those patients with underlying cardiac and other risk factors, there is also evidence that these events may also occur in younger patients without a history of clinical cardiac disease or obvious additional risk factors (162). With regard to the pathogenesis, it is being increasingly recognised that platelet activation may play a central role in CAP-associated AMI, raising the possibility that anti-platelet agents, such as aspirin, among many other agents, may be beneficial in preventing these events, as has been documented in at least one study in the elderly (164-166). Furthermore, the pathogenesis of these cardiac events in pneumococcal CAP is increasingly being understood (167,168). It is recommended that any patients with CAP
who have well-recognised risk factors for a cardiovascular event, or patients who are complicated by clinical failure, should be investigated for the possibility of a cardiovascular event. Importantly, such events, when they occur in patients with CAP, are associated with a poorer prognosis acutely, as well as a poorer prognosis and associated increased risk of cardiovascular events on long-term follow-up (169-171). This topic has been reviewed in more detail elsewhere (172).

**Recommendations**

- Patients with CAP with well-recognised risk factors for a cardiovascular event or cases that fail to show adequate clinical recovery should be investigated for the possibility of a cardiovascular event (A III).

**Long-term sequelae of lung damage**

CAP occasionally leads to bronchiectasis, particularly if recurrent or associated with an underlying anatomic or immune defect. Bronchiectasis is defined as abnormal dilation and distortion of the bronchial tree, and is characterised by chronic sputum production with persistent airflow limitation. Significant infection, particularly in childhood, can cause structural damage that impairs mucociliary clearance and facilitates chronic bacterial infection. Over time, persistent infection may then result in bronchiectasis. Aetiological agents that have been implicated in the original precipitating bronchial wall injury include *Mycoplasma pneumoniae, Bordetella pertussis*, adenovirus and measles. Tuberculosis remains an important cause of bronchiectasis in patients from high-burden communities and in those with HIV infection.

**Aspiration pneumonia**

**Definitions**

The term ‘aspiration’ refers to the abnormal entry of a large inoculum of exogenous substances or endogenous secretions into the lower airways. This may result in lung inflammation with associated clinical consequences or resolve spontaneously without any therapy. The clinical presentation depends on the type and volume of aspirated material, the frequency of aspiration and the host immune response. The major syndromes related to aspiration include (173) airway obstruction from inhalation of particulate matter (174), aspiration pneumonitis, a chemical injury caused by aspiration of sterile liquid gastric contents (or other noxious fluids), and (175) aspiration pneumonia, an infectious process caused by the inhalation of oropharyngeal secretions colonised by microorganisms (173-176). While most cases of CAP are caused by microaspiration of relatively virulent bacteria residing in the upper respiratory tract (177), aspiration pneumonia refers to a syndrome of bacterial pneumonia in susceptible patients with defective lower airway clearance mechanisms who aspirate a large inoculum of normally nonvirulent pathogens (178,179).

Although some features of aspiration pneumonitis and aspiration pneumonia overlap, they represent distinct clinical entities in terms of pathophysiological mechanisms, clinical manifestations and treatment. So-called ‘bland’ aspiration, as occurs after haematemesis or aspiration of enteral feeds, may lead to CXR infiltrates but does not result in an inflammatory response in the lung and usually resolves spontaneously without antimicrobial therapy (178,180,181). There is often a failure by clinicians to distinguish aspiration pneumonitis or bland aspiration from aspiration pneumonia and a tendency to consider all aspiration syndromes to be infectious, resulting in overuse of antimicrobials (173).

**Epidemiology and risk factors**

Accurate estimation of the prevalence of community-acquired aspiration pneumonia (CAAP) is limited by the lack of a standardised case definition, and because most studies do not distinguish aspiration pneumonitis from pneumonia (179). Observational studies from developed countries have found that up to 15% of CAP episodes may result from aspiration (47,182-186). Stroke patients who aspirate have a 7-fold higher risk of developing pneumonia (187), which complicates 10% of acute strokes and is associated with a significantly increased risk of death (188). Patients with CAAP are more likely to be admitted to intensive care units (189,190) and require mechanical ventilation than those with non-aspiration CAP, and have a significantly increased in-hospital mortality and length of stay (182,191).

The primary predisposing mechanisms for aspiration include dysphagia and altered mental status (192), resulting in compromised glottic closure and cough reflexes (175,176). Dysphagia is regarded as the most important risk factor for aspiration pneumonia; it is most commonly due to neurological and oesophageal disorders, but also complicates COPD (193) and the use of antipsychotic medications (194). Alcohol abuse and seizures are strongly associated with anaerobic aspiration pneumonia because of reduced levels...
of consciousness, poor oral hygiene, immune dysfunction and delayed presentations (179). Conditions such as dental caries, periodontal disease and gingivitis increase the risk of oropharyngeal colonisation with pathogenic organisms and a higher overall bacterial load, and are associated with a higher risk of aspiration pneumonia (173,176,195). The elderly therefore represent a higher risk group (186) because of the more frequent neglect of oral hygiene (196) and higher rates of neurological disease (173). Nasogastric- or gastrostomy-tube feeding are independent risk factors for aspiration (197).

Microbiology

A dominant role for anaerobic organisms in aspiration pneumonia was suggested by early studies using animal models and invasive diagnostic procedures such as transthoracic and transtracheal needle aspiration (198). The most common isolates included Bacteroides melaninogenicus and other Bacteroides species, Fusobacterium nucleatum, and Peptostreptococcus spp. (175,199,200), many of which produce beta-lactamases. The relevance of these early studies assessing the bacteriology of aspiration pneumonia has been questioned, with concerns about the sterility of sampling techniques employed (201) and the inclusion of patients with established complications such as lung abscess and empyema (173). More recent studies using protected specimen brushes to sample the lower respiratory tract of patients with severe aspiration pneumonia isolated bacterial pathogens in a minority of cases. In these studies S. pneumoniae, S. aureus, H. influenzae, and Enterobacteriaceae predominated, and no pathogenic anaerobic organisms were isolated (202,203). In a group of institutionalised elders with severe aspiration pneumonia who underwent bronchoscopic sampling, gram-negative enteric bacilli were the predominant organisms isolated (49%), followed by anaerobic bacteria (16%), and S. aureus (12%) (204). This shift in microbiological profiles may reflect a true decline in anaerobic infection due to improved social conditions and access to health care, but the overlap with organisms found in health-care associated pneumonia suggests an increased occurrence of aspiration in these settings.

Clinical features and diagnosis

The early clinical features of aspiration pneumonia are difficult to distinguish from other causes of CAP, particularly because the aspiration event is usually not witnessed (175). The diagnosis is usually made in patients presenting with a pneumonic process with a predisposition for aspiration (i.e., difficulty swallowing or a reduced level of consciousness), plus involvement of dependent pulmonary segments (posterior segments of the upper lobes and the apical segments of the lower lobes when aspiration occurs in the recumbent position, or the basal segments of the lower lobes in an upright position), particularly in the right lower lobe (202,203). There are no clinical or biochemical findings that reliably distinguish anaerobic aspiration pneumonia from CAP (176,205), but the following may suggest an anaerobic cause of pneumonia (173-176,206):

- The production of foul-smelling sputum, suggesting infection with anaerobic organisms.
- The development of lung abscesses, necrotising pneumonia or empyema; the onset of these complications may be indolent, occurring around 2 weeks after the aspiration event.

In contrast, aspiration pneumonitis is a hyperacute illness that usually occurs after a witnessed aspiration event with rapid-onset dyspnoea (within 2 hours of aspiration), bronchospasm, bilateral patchy CXR infiltrates (including non-dependent areas), and frothy sputum. Hypoxia is a prominent feature, and patients may progress to develop acute respiratory distress syndrome. This may be accompanied by a systemic inflammatory response with fever, leukocytosis and tachycardia, despite the absence of infection (175,207). Based on animal models, more than 120 mL of gastric contents need to be aspirated to induce chemical pneumonitis in an average-sized adult (179), and so a witnessed large aspiration event supports this diagnosis.

Antimicrobial therapy

The frequent finding of anaerobic infections in CAAP in microbiological studies performed in the 1970s led to a change in recommendations for empiric antibiotic therapy for aspiration pneumonia, with a shift away from penicillin to the use of agents with specific anaerobic coverage such as clindamycin, metronidazole and beta-lactam-beta-lactamase inhibitor combinations. Small RCTs conducted in the 1980s and early 1990s comparing penicillin to clindamycin for patients with lung abscess and confirmed anaerobic pneumonia showed much better cure rates with the use of clindamycin (208,209).

Recommendations

- Acute aspiration events, particularly in the absence of
Vaccination to prevent CAP

Implementation of the South African National Guidelines for vaccination against influenza and pneumococcal infections may both assist in preventing CAP (see www.pulmonology.co.za).

Two pneumococcal vaccines are registered for use in adults in South Africa, the 23-valent polysaccharide vaccine (PPV23), and the 13-valent pneumococcal conjugate vaccine (PCV13).

The indications for PPV23 reported in the South African pneumococcal vaccination guideline are similar to that reported in the most recent Advisory Committee on Immunization Practices (ACIP) recommendations (210). In brief, PPV23 is recommended for use in adults aged 19–64 years with underlying comorbid conditions, who smoke cigarettes, have functional or anatomical asplenia and in those with immunocompromising conditions such as HIV infection, haematological malignancies and transplant patients. It is also recommended for use in persons 65 years of age or older.

In 2014, the PCV13 was registered for use in adults in South Africa, as in other parts of the world, based on initial immunogenicity studies. The initial registered indication in South Africa was the use of a single dose of PCV13 in adults ≥50 years, as had been registered elsewhere (211). However, at the end of 2015, PCV13 received registration in South Africa for use in adults aged 18 years and older as a single dose (212). Specific groups at high risk for pneumococcal infection were mentioned in the registration including cases with sickle cell disease and HIV infection that were recommended to receive at least one dose of PCV13 whether or not they had received one or more doses of PPV23 previously. A specific regimen was recommended for patients with haematopoietic stem cell transplants.

Elsewhere in the world, such as in the USA, PCV13 has also been registered for use in adults ≥19 years with underlying comorbid and immunocompromising conditions (213), and in adults ≥65 years (214). In those with high-risk factors for pneumococcal disease (e.g., CSF leak, cochlear transplant and functional or anatomical asplenia) and those with immunocompromising conditions, as well as those aged ≥65 years of age, the recommendation is for the use of PCV13 in sequence with PPV23. The recommendation is that PCV13 should always preferably be given first. In individuals who have not previously had PPV23 vaccination the PCV13 vaccine should be given first, followed a minimum of 2 months later with the PPV23 vaccine in the case of adults ≥19 years of age with the high-risk underlying comorbid and immunocompromising conditions, and 12 months later in individuals ≥65 years. However, in any of these cases, if the individual has already had a vaccination with PPV23, the PCV13 vaccination should be given a minimum of 1 year after the PPV23 vaccination.

Two recent clinical studies attest to the clinical efficacy of the PCVs in adults in different settings. The first study was that of the older 7-valent pneumococcal conjugate vaccine (PCV7) in predominantly HIV-infected adults and adolescents (aged >15 years of age) in Malawi, who had recently recovered from an episode of invasive pneumococcal disease (215). This was a randomised, double-blind, placebo controlled trial and in the active arm of the study, two doses of PCV7 were given to the patients 4 weeks apart. The efficacy of the vaccine for the primary endpoint, which was prevention of a further episode of vaccine serotype (or serotype 6A) pneumococcal infection, was 74% (95% CI, 30–90%). The second was a study in The Netherlands among adults >65 years which evaluated the efficacy of PCV13 in preventing first episode of vaccine type strains of community-acquired pneumococcal pneumonia (primary endpoint), non-bacteremic and noninvasive pneumococcal CAP and invasive pneumococcal disease (secondary endpoints) (216). Patients in the active arm were given one dose of PCV13. Vaccine efficacy for the primary endpoint was 45.6% (95.2% CI, 21.8% to 62.5%), and for the secondary endpoints 45.0% (95.2% CI, 14.2% to 65.3%) and 75% (95.2% CI, 41.4% to 90.8%), respectively. Efficacy persisted throughout the duration of the study of almost 4 years.

The trivalent inactivated influenza vaccine is the only licensed influenza vaccine available in South Africa (121).
Because of the changing nature of influenza viruses, each year the strains included in the vaccine for the following influenza season are updated. For this reason, as well as limited duration of effectiveness, influenza vaccination is recommended to be repeated each year. Influenza vaccine usually becomes available in South Africa from March and should be given sufficiently early to provide protection for the winter season. Influenza vaccine effectiveness depends on characteristics of those being vaccinated (age and health), whether there is a good match between the circulating viruses and the viruses contained in the vaccine, and on influenza types and subtypes. In general, influenza vaccines work best among children ≥2 years and healthy adults. Older people (≥65 years), children <2 years and severely immunocompromised individuals often have poorer immune responses to trivalent inactivated influenza vaccine (IIV) compared with healthy adults. However, even for these people the influenza vaccine still provides some protection. Other products, e.g., high-dose influenza vaccine and adjuvant vaccines, have been shown to be more effective in certain groups (217) but these vaccines are not available in South Africa. In the elderly, influenza vaccination has been shown to reduce the incidence of severe disease including bronchopneumonia, hospital admissions and mortality. Trivalent IIV has been shown to provide protection in HIV-infected adults without severe immunosuppression (218). Vaccination of healthcare workers may decrease the risk of spreading influenza to their patients. Vaccinating individuals at risk of severe influenza may provide direct protection for these individuals. In addition, vaccinating individuals in close contact with people at risk for severe influenza may provide indirect protection through preventing transmission to high-risk individuals. Groups who should receive influenza vaccine are shown in Box 4.

Recommendations

- All adults ≥50 years who are vaccine naïve should receive a single dose of PCV13 (A II).
- All adults ≥50 who have received PPV23 should receive a single dose of PCV13 one year later (A II).
- All adults ≥65 years who are vaccine naïve should receive a single dose of PCV13 followed a year later by PPV23 (A II).
- All adults ≥65 years who have received PPV23 should receive a single dose of PCV13 at least one year later (A II).
- Younger adults (≥18 years) who are vaccine naïve with severe underlying comorbid or immunocompromising conditions including HIV infection should receive a single dose of PCV13 followed at least 2 months later by PPV23 (A II).
- Younger adults (≥18 years) who have previously received PPV23 and have severe underlying comorbid or immunocompromising conditions including HIV infection should receive a single dose of PCV13 one year later (A II).
- All women who are pregnant in the period of influenza

<table>
<thead>
<tr>
<th>Box 4 Persons who should receive annual influenza vaccine</th>
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<tr>
<td>Pregnant women irrespective of stage of pregnancy, or postpartum</td>
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<tr>
<td>HIV-infected adults</td>
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<tr>
<td>Healthcare workers</td>
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<tr>
<td>Persons (adults or children) who are at high risk for influenza and its complications because of underlying medical conditions and who are receiving regular medical care for conditions such as chronic pulmonary (including tuberculosis) and cardiac diseases, chronic renal diseases, diabetes mellitus and similar metabolic disorders, individuals who are immunosuppressed, and individuals who are morbidly obese (body mass index ≥40 kg/m²)</td>
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<tr>
<td>Residents of old-age homes and chronic care and rehabilitation institutions</td>
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<tr>
<td>Persons aged ≥65 years</td>
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<td>Children aged 6–59 months (efficacy of trivalent IIV is low in this group)</td>
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<tr>
<td>Persons aged 6 months to ≤18 years on long-term aspirin therapy</td>
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<tr>
<td>Adults and children who are family contacts of individuals at high risk of severe influenza</td>
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<tr>
<td>Any persons wishing to minimise the risk of influenza acquisition, especially in industrial settings, where large-scale absenteeism could cause significant economic losses</td>
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vaccine availability (approximately March to June) should be offered influenza vaccination with IIV3 (A I).

- Adults aged ≥65 years should be offered annual influenza vaccination with IIV3 (A I).

- All adults with specific chronic diseases: chronic pulmonary [including tuberculosis] and cardiac diseases, chronic renal diseases, diabetes mellitus and similar metabolic disorders, individuals who are immunosuppressed including HIV-infected individuals, and individuals who are morbidly obese (body mass index ≥40 kg/m²) should be offered annual influenza vaccination with IIV3 (A II).

- All healthcare workers should be offered annual influenza vaccination with IIV3 (A II).

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Footnote

Conflicts of Interest: AC Whitelaw has received honoraria to the Institute from MSD, Novartis, AstraZeneca, and Takeda. A Brink has received honoraria from MSD, Cipla and Pfizer. C Feldman has acted on the advisory board and/or speakers bureau of companies manufacturing and/or marketing macrolide antibiotics (Abbott, Aspen GSK, Pfizer, Sandoz) and ceftaroline (AstraZeneca/Pfizer) and companies manufacturing and/or marketing the pneumococcal conjugate vaccine (Pfizer). C Cohen has received received grant funds to the Institute from Sanofi and travel paid for by Sanofi and Parexel. GA Richards has received honoraria from Aspen, Cipla, MSD, Pfizer, AstraZeneca, Sanofi, BMS and Fresenius Kabi. K Dheda has received grants, free products/kits from Hain Life Sciences, Alere Diagnostics, FIND Diagnostics, Pfizer, Novartis, Cipla, GSK, AstraZeneca and Sanofi-Aventis. The other authors have no conflicts of interest to declare.

Disclaimer: The recommendations are based on currently available scientific evidence together with the consensus opinion of the authors. Adherence to these recommendations is voluntary and does not account for individual variation among patients; the recommendations are not intended to supplant physician judgement with respect to particular patients or special clinical situations. In addition, the recommendations do not indicate an exclusive diagnostic work-up or course of treatment or serve as a standard of medical care.

References


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