Part IV. Human infections and antibiotic resistance

Principal authors: P Crowther-Gibson, N Govender, D A Lewis, C Bamford, A Brink

Co-authors: A von Gottberg, K Klugman, M du Plessis, A Fali, B Harris, K H Keddy, M Botha

Keywords: antibiotics; antibiotic ( antimicrobial) resistance; pneumonia; acute respiratory infection; enteric infections; sexually transmitted infections; hospital-acquired infections

South Africa has a high burden of infectious diseases, including a large portion that are of bacterial origin. This section reviews the national burden of disease and levels of antibiotic resistance in common bacterial infections in the human population. The consequences of resistance on clinical outcomes, through either treatment failures or the development of more virulent infections, are largely unknown. The full impact of antibiotic resistance on health in South Africa therefore remains to be assessed.

National burden of disease
South Africa faces a quadruple burden of disease, as a result of the HIV/AIDS epidemic, other infectious diseases, injuries, and non-communicable diseases. Tables I and II show the top five causes of death for all ages and for children under the age of 5 (information from the Revised Burden of Disease Estimates for South Africa 20001 and the 2010 South African Health Review2).

The largest single cause of death for all ages is HIV/AIDS, accounting for 26% of deaths.1 This is 5 times greater than the next largest single cause of death, ischaemic heart disease and stroke (7% each) followed by tuberculosis (TR) and interpersonal violence, each accounting for about 6%. While males have higher proportions of deaths owing to homicide/violence and TB than females, females have higher proportions of deaths due to HIV/AIDS, heart disease and stroke.

There is considerable uncertainty around estimates of child mortality in South Africa because of incomplete vital registration.2 Existing numbers suggest that HIV/AIDS is the leading cause of death (46%), followed by neonatal causes dominated by perterm complications, asphyxia, and infection. Diarrhoea, pneumonia and injuries together account for 17% of mortality.

HIV/AIDS and TB
The most pressing health concern in South Africa is the HIV/AIDS epidemic, with around 29% of the population infected with the virus (2009). In addition to a high incidence of chronic illness and violence-related deaths, South Africa has the largest number of people living with HIV/AIDS in the world (over 5 million), and 1 000 people are estimated to die as a result of AIDS daily.1 The Health Economics and HIV/AIDS Research Division predicts that HIV patients will soon account for around 60 - 70% of all hospital expenditures. HIV-related illnesses currently account for 50% of hospital admissions.

In absolute terms, South Africa has the fourth-largest TB population in the world (behind India, China and Indonesia) and bears 28% of the global burden of TB related to HIV. In 2007 data from the Global TB database, almost 1 000 per 100 000 of the population are infected with the disease annually. The emergence in South Africa of extremely drug-resistant tuberculosis (XDR-TB) that is considered virtually untreatable is of particular concern in a country with a high prevalence of HIV and a poor record of TB treatment.

Bacterial disease and antibiotic resistance
Our data are summarised from national surveillance efforts and site-specific case studies. The picture is incomplete because causes of illnesses and deaths are not well counted in South Africa, as is often the case in low-resource countries. Furthermore, separating bacterial from viral diseases requires a level of detail that, in most cases, does not exist. Nonetheless, the available information provides a basic idea of the current situation.

We present information on the burden of disease, current treatment options and antibiotic resistance for acute respiratory infections, diarrhoeal infections, sexually transmitted infections and nosocomial infections.

Acute respiratory and meningeal infections
As causes of severe respiratory tract, systemic and meningeal infections such as pneumonia, bacteraemia, and meningitis, the bacterial pathogens Streptococcus pneumoniae, Neisseria meningitidis and Haemophilus influenzae are major contributors to morbidity and mortality worldwide.

Antimicrobial chemotherapy has substantially decreased morbidity and mortality from these infectious diseases. However, their control is threatened by the global increase in antimicrobial resistance (AMR), including multidrug resistance. Resistant infections may adversely affect mortality, treatment costs, disease spread and duration of illness, increasing pressure on the choice of appropriate antibiotics.

<table>
<thead>
<tr>
<th>Age group</th>
<th>HIV/AIDS</th>
<th>Ischaemic heart disease</th>
<th>Stroke</th>
<th>Tuberculosis</th>
<th>Interpersonal violence and injuries</th>
</tr>
</thead>
<tbody>
<tr>
<td>All ages</td>
<td>25.5%</td>
<td>6.6%</td>
<td>6.5%</td>
<td>5.5%</td>
<td>5.3%</td>
</tr>
</tbody>
</table>

Source: Revised Burden of Disease Estimates for South Africa 20001

<table>
<thead>
<tr>
<th>Children under 5</th>
<th>HIV/AIDS</th>
<th>Age group</th>
<th>Diarrhoea</th>
<th>Pneumonia</th>
<th>Injuries</th>
</tr>
</thead>
<tbody>
<tr>
<td>46%</td>
<td>29%</td>
<td>9%</td>
<td>6%</td>
<td>2%</td>
<td></td>
</tr>
</tbody>
</table>

In developing countries such as South Africa and sub-Saharan Africa, where respiratory and meningeal diseases are more frequent because of the high burden of HIV infections, limited access to health care, costly antibiotics and low vaccination coverage, lack of access to antimicrobials and resistance to those available may increase the morbidity and mortality of vaccine-preventable diseases.

**Streptococcus pneumoniae**

*S. pneumoniae* is a leading cause of bacterial infection worldwide. The World Health Organization (WHO) estimates that 1.6 million people, including up to 1 million children aged <5 years, die annually of pneumococcal infection, with most deaths in developing countries.

**Treatment**

Successful management of pneumococcal disease involves use of early antimicrobial therapy. During the 1940s, clinical pneumococcal isolates exhibited complete susceptibility to antibiotics such as penicillin, the antibiotic of choice for the treatment of pneumococcal infections. However, in Australia, intermediate penicillin resistance was observed for the first time in 1967. In South Africa, fully penicillin-resistant *S. pneumoniae* strains were detected in 1977, and in 1978 the occurrence of multidrug-resistant and highly resistant strains was reported. Since then, the prevalence of *S. pneumoniae* antibiotic resistance has increased around the world, not only to penicillin but also to non-β-lactam drugs, such as the macrolides, tetracycline, chloramphenicol, the fluoroquinolones and co-trimoxazole. Resistance to non-β-lactam drugs is often associated with decreased susceptibility to penicillin, so the prevalence of multidrug-resistant strains is also increasing.

**Antibiotic resistance**

The increasing prevalence of pneumococcal resistance to single and multiple antimicrobials in South Africa demonstrates the need for new strategies to combat the problem, especially in terms of preventing increased mortality and treatment failures in penicillin-resistant pneumococcal meningitis. The clinical impact of antibiotic resistance has been reported as treatment failures for acute otitis media, and for pneumococcal meningitis. However, using the revised penicillin susceptibility breakpoints of 4 µg/ml for intermediate and ≥8 µg/ml for resistant strains, there is no evidence for a relationship between penicillin resistance and pneumococcal pneumonia treatment failures.

Surveillance data reveal that rates of resistance to penicillin and other antibiotics among *S. pneumoniae* vary by geographical location. Penicillin-resistant pneumococci have been reported with particularly high frequencies in South Africa since the mid-1970s and in other African countries since the 1980s.

A 1997 African multicountry study revealed that penicillin resistance levels among all isolates of *S. pneumoniae* ranged from 9% to 61%, and that an increase in resistance has been observed in four countries across Africa. Resistance levels in North Africa are generally high, as reported by Algeria (35% of all isolates), Egypt (49% of invasive isolates), and Tunisia (41% of all isolates), although in Morocco levels are much lower (9% of all isolates). Penicillin resistance in West Africa varies from 62% in Senegal to 31% in Ghana and 22% in Ivory Coast down to 7% of invasive isolates in The Gambia.

In East Africa, the prevalence in Ethiopia and Kenya has increased over the years to 29% of clinically significant isolates, and 48% of paediatric invasive isolates, respectively. In Malawi, a study of paediatric nasopharyngeal isolates revealed penicillin resistance of 21%. In Zambia, 14% of paediatric non-invasive isolates were penicillin-resistant in 1994 and in Mozambique, a prevalence of 14% of paediatric invasive isolates was reported. No systemic studies have defined the reasons for the diversity in resistance rates reported.

South Africa has been the primary site of pneumococcal penicillin resistance surveillance and research in Africa, and has had one of the highest reported rates in the world. Since the first reports of resistance in the 1970s, the prevalence of resistance in *S. pneumoniae* in South Africa has increased. Between 1979 and 1986, the prevalence of resistance to one or more antibiotics in pneumococcal cerebrospinal fluid and blood isolates increased from 4% to 14%. A continuation of the same survey, between 1991 and 1998, reported that antibiotic resistance increased from 19% to 25% among all age groups, and in children from 32% to 38%. Among all age groups during this period, penicillin resistance increased from 10% to 18%.

In 1992, 40% of isolates causing community-acquired meningitis or bacteraemia in children were penicillin-resistant. A 1999 Alexander Project study of pneumococcal isolates from the private sector in Johannesburg revealed that 79% were penicillin-resistant. Also in 1999, a study of nasopharyngeal isolates from private paediatric practices in Johannesburg found that antibiotic resistance was 69%, with 37% being multiply resistant. In 2003, a report on isolates from private clinical laboratories in South Africa showed that the rate of penicillin resistance among all age groups was 76%. Other studies in that year on invasive pneumococcal isolates from adults outside the private sector who were likely to have been exposed to less antibiotic prescribing revealed penicillin resistance levels of 13% and a study on non-invasive isolates from HIV-infected adults reported penicillin resistance at 15%. A study conducted in Gauteng in 2006 on adults with bacteraemic pneumonia showed that 33% of isolates were penicillin-resistant, when using the historical meningitis susceptibility breakpoints. Penicillin resistance in South Africa remains mainly intermediate in level, with a low prevalence of fully resistant isolates. Although, annually, resistance levels have increased overall, these are dependent on the site of specimen collection, age of the patients, and their location in the country.

**Macrolide resistance.** The increasing incidence of penicillin-resistant *S. pneumoniae* has been paralleled by an increase in resistance to other classes of antimicrobials, suggesting that penicillin resistance serves as a marker of resistance to other drugs. Almost 25% of *S. pneumoniae* isolates from South Africa show full erythromycin resistance, with over 90% of these also resistant to clindamycin. Additionally, 40 - 50% of penicillin-resistant isolates show cross-resistance to macrolides. In 2001, a national multicentre study of private clinical laboratories in South Africa revealed that a high prevalence (61%) of non-invasive isolates were macrolide (clarithromycin and azithromycin) resistant, whereas a national 2005 study showed that 14% of invasive pneumococcal isolates were resistant to macrolides.

**Co-trimoxazole resistance.** The Alexander Project in Johannesburg in 1996 - 1997 revealed that 15 - 20% of isolates were resistant to co-trimoxazole, while a 2001 study showed that co-trimoxazole resistance was as high as 72%. Other South African studies have found that co-trimoxazole resistance is associated with multidrug resistance in pneumococcal isolates from childhood carriers in hospitals and healthy children in the community. In a 1986 study, 43% of penicillin-resistant strains among childhood carriers in South Africa were also resistant to co-trimoxazole.

**Fluoroquinolone resistance.** A Canadian study showed that 1% of pneumococci have reduced susceptibility to fluoroquinolones. An increase in the frequency and degree of resistance to fluoroquinolones among pneumococci occurred particularly in penicillin-resistant *S. pneumoniae*.
pneumoniae, and in adults over 65 years.47 In South Africa, 2008 study data suggest that the use of fluoroquinolones to treat multidrug-resistant TB in children has led to the emergence of invasive pneumococcal disease (IPD) caused by levofloxacin-non-susceptible S. pneumoniae,48 although these strains remain rare outside those institutions.

**Multidrug resistance.** S. pneumoniae resistance to three or more different classes of antibiotics, defined as multidrug resistance, is a problem of increasing concern worldwide.48 The emergence of multidrug resistance was first reported in Soweto, South Africa, in 1977.21 Subsequently, multidrug resistance emerged globally.20-23,35-37 In South Africa in 2004, a third of pneumococcal isolates studied displayed multidrug resistance.49 Successful multidrug-resistant clones that are disseminated worldwide belong to only 10% of the 93 pneumococcal serotypes, including serotypes 3, 6A, 6B, 9N, 9V, 14, 19A, 19F and 23F.50

**Neisseria meningitidis**

*N. meningitidis* causing meningitis and other meningococcal diseases, such as meningococcaemia, is a major cause of morbidity and mortality in children worldwide, and of epidemics in Africa and Asia.

**Antibiotic resistance**

In Africa, limited data are available regarding antimicrobial non-susceptibility of *N. meningitidis*. Few studies have documented the existence of penicillin and other antimicrobial non-susceptibility in Africa other than South Africa. A study in Morocco reported an average rate of 4% for penicillin intermediate resistant invasive meningococcal isolates collected from 1992 to 2000.51 All isolates tested were susceptible to cefotaxime, chloramphenicol and rifampicin. Laboratory-based surveillance in Egypt from 1998 to 2004 reported high rates (86%) of resistance to co-trimoxazole but low rates of resistance to penicillin (1%) and ampicillin (5%).40% of isolates were intermediate resistant to either ampicillin (minimum inhibitory concentration (MIC) 0.25 - 1 μg/ml) or penicillin (MIC 0.12 - 0.25 μg/ml) and 34% were intermediate resistant to both penicillin and co-trimoxazole.44 One isolate, with intermediate resistance to penicillin, tested positive for β-lactamase production.

A serogroup A meningitis outbreak in northern Ghana in 1998 showed no evidence of resistance to any of the drugs tested, with the exception of sulphadiazine.55 No resistance to β-lactam agents or chloramphenicol was reported during surveillance of meningococcal meningitis in Cameroon during the 2007 and 2008 meningitis seasons. In Ethiopia, epidemic meningococcal isolates collected during 2002 - 2003 were compared with those from the 1988 - 1999 epidemic.56 All 40 isolates were fully susceptible to the antibiotics tested, except for sulphonamethoxazole (MIC >256 μg/ml). A study of the aetiology of bacterial meningitis in Nigeria was conducted between 1987 and 1992, and *N. meningitidis* was the most common pathogen isolated.57 Antimicrobial susceptibility testing by disc diffusion of meningococcal isolates collected from 1992 to 2000 showed no evidence of resistance to any of the drugs tested, with the exception of sulphadiazine.55 No resistance to β-lactam agents or chloramphenicol was reported during surveillance of meningococcal meningitis in South Africa was initiated during 1999. A study that genotypically characterised invasive meningococci collected from 2001 to 2005 reported a relatively low prevalence of penicillin non-susceptibility.59 During this period 6% of isolates were intermediately resistant to penicillin, with MICs ranging from 0.094 μg/ml to 0.25 μg/ml. No isolates tested were fully resistant or tested positive for β-lactamase production and all were susceptible to other drugs tested, with the exception of rifampin (0.3%). In 2009, South Africa reported its first case of fluoroquinolone-resistant *N. meningitidis*.60 MICs for ciprofloxacin and levofloxacin were 0.125 μg/ml and 0.25 μg/ml for ofloxacin. Resistance appeared to be mediated by a single amino acid substitution in the DNA gyrase enzyme. The isolate was susceptible to other drugs tested but was resistant to nalidixic acid (12 μg/ml). No subsequent cases of fluoroquinolone-resistant meningococci have been reported.

**Haemophilus influenzae**

*H. influenzae* is an important cause of acute otitis media, sinusitis, chronic bronchitis, community-acquired pneumonia and meningitis.50 Before the introduction of *H. influenzae* type b (Hib) conjugate vaccines, globally Hib was estimated to be responsible for approximately 3 million serious illnesses and 386 000 deaths annually.50,51 95% of these cases and 98% of all deaths occurred in patients from developing countries, mainly in children <5 years.61 In sub-Saharan African children, Hib is responsible for 20% of all radiologically confirmed pneumonia cases and 40% of all meningitis cases.62,63

**Treatment**

Antimicrobial treatment is pivotal in the management of *H. influenzae* disease. Until the early 1970s, when *H. influenzae* resistance to ampicillin was first reported,44 ampicillin was the cornerstone of therapy.52,64 In sub-Saharan Africa, chloramphenicol and penicillin are the first-line antibiotics to treat meningitis and severe pneumonia, while mild pneumonia is treated with co-trimoxazole, ampicillin or amoxicillin.50-70 In South Africa, β-lactams such as penicillin, ampicillin or amoxicillin are still recommended as empirical first-line therapy for the treatment of respiratory tract infections in patients <65 years old and without co-morbid illness.70,71 Alternative agents recommended for treating patients >65 years old, or who have co-morbid illness, include amoxycillin-clavulanate or selected oral cephalosporins (cefuroxime axetil or cefpodoxime).71,72

**Antibiotic resistance**

The increasing prevalence of resistance among *H. influenzae* isolates to commonly used antibiotics is of concern. Resistance to penicillin is high, with prevalence rates of >45% reported in some settings.50,70,72,73 Resistance to ampicillin and other β-lactams is almost exclusively due to β-lactamase production. Isolates expressing this mechanism remain susceptible to β-lactamase-inhibitor combinations such as amoxicillin-clavulanic acid. A second non-β-lactamase-mediated resistance mechanism is conferred by mutations in the *fbpA* gene, encoding the transpeptidase region of penicillin-binding protein 3 (PBP3), which results in decreased affinities of the PBP3 for β-lactams.73 Such strains are termed β-lactamase-negative ampicillin-resistant (BLNAR). Worldwide, BLNAR strains continue to be isolated at very low frequencies.73-75 However, their prevalence has recently increased in countries such as Japan,76 Spain,77-79 and Korea.81

In Africa, data for *H. influenzae* AMR, especially regarding trends, are sparse.50,70,72,82-84 Increasing rates of chloramphenicol and co-trimoxazole resistance have been reported in Africa.70,72,84 In Cameroon, chloramphenicol resistance levels of up to 84% have been reported,44 while high prevalence of co-trimoxazole resistance have been reported in Mozambique (46%)82 and Kenya (66%).75
Beta-lactamase production is by far the most common mechanism of ampicillin resistance in South African isolates of *H. influenzae*. From 2003 to 2008, 2 177 cases of invasive *H. influenzae* were reported to the national laboratory-based surveillance system, of which 54% had viable isolates available for antimicrobial susceptibility testing. Of the viable isolates, 2% and 15% were found to be intermediate resistant and resistant to ampicillin, respectively. Of the 190 ampicillin non-susceptible isolates, 99% were β-lactamase producing and 1% were phenotypically β-lactamase-negative ampicillin resistant (BLNAR) and were characterised as low-level BLNAR (MIC 2 µg/ml). In addition, a β-lactamase-positive amoxicillin-clavulanate-resistant (BLPACR) strain was identified (MIC 8 µg/ml).

In the only previous report of South African BLNAR strains (ampicillin MIC 2 µg/ml), a BLNAR prevalence of 6% among isolates collected from various sources, including respiratory secretions and blood, was reported during a SENTRY worldwide surveillance programme in the Asia-Pacific region.

**Diarrhoeal infections**

**Non-typhoidal Salmonella**

Salmonellosis due to non-typhoidal *Salmonella enterica* spp. accounts for a large burden of disease worldwide. Illness is usually self-limiting and antimicrobial therapy is not required, but in cases of invasive disease antimicrobial therapy is important for a successful clinical outcome. Over the period 2003 - 2010, the Enteric Diseases Research Unit (EDRU) at the National Institute for Communicable Diseases (NICD) has documented 16 435 records of laboratory-confirmed cases of non-typhoidal *Salmonella enterica* isolates from human and non-human sources for South Africa. Isolates received from non-human sources (N=224) include samples of water, food and animal specimens processed at the EDRU for study purposes, or as a service by special request and not as part of their routine surveillance activities. These isolates were therefore not screened for antimicrobial susceptibility. Of the 16 211 human isolates, 13 702 were viable and were screened using antimicrobial agents.

**Treatment**

The treatment of choice for such infections are third-generation cephalosporins and fluoroquinolones, as resistance to ampicillin, chloramphenicol and co-trimoxazole has been present worldwide for many years. Failure to respond to treatment with the fluoroquinolones, as isolates have displayed decreased susceptibility to ciprofloxacin, has recently been reported. AMR to nalidixic acid has been used as a proxy to identify isolates that may not respond to treatment with ciprofloxacin.

**Antibiotic resistance**

Resistance to quinolones usually occurs as a result of alterations in the target enzymes (DNA gyrase and topoisomerase IV) and as a result of changes in drug entry and drug efflux. Resistance to quinolones can also be mediated by plasmids that carry genes coding for Qnr proteins, which protect the quinolone targets from inhibition. Plasmid-mediated quinolone resistance among South African strains of non-typhoidal *Salmonella* has been previously reported, as well as the detection of mutations in the DNA gyrase enzyme of clinical non-typhoidal *Salmonella*. In the period 2003 - 2010 there has been a decrease in the proportion of non-typhoidal *Salmonella* isolates showing resistance to ampicillin from 64% to 16%, chloramphenicol from 47% to 14%, ceftriaxone from 40% to 10%, and nalidixic acid from 38% in 2003 to 10% in 2010. Although the overall proportion of non-typhoidal *Salmonella* isolates showing resistance to nalidixic acid has decreased over time, when comparing non-typhoidal *Salmonella* isolates causing invasive disease with non-typhoidal *Salmonella* isolates causing non-invasive disease, isolates causing invasive disease account for the greater proportion of isolates showing resistance to nalidixic acid. There has been no increase in the proportion of non-typhoidal *Salmonella* isolates exhibiting resistance to ciprofloxacin. In 2004, the greatest proportion of non-typhoidal *Salmonella* isolates, just less than 2% (26/1 597), showed resistance to ciprofloxacin. Overall, just less than 1% of all non-typhoidal *Salmonella* isolates exhibited resistance to ciprofloxacin from 2003 to 2010. Over this same period the proportion of non-typhoidal *Salmonella* isolates exhibiting resistance to sulfamethoxazole has fluctuated from 40% of isolates in 2003, to a high of 78% of isolates for 2004 and 2005, to 48% of isolates in 2010, but overall there has been a general decrease in resistance to sulfamethoxazole since the highs of 2004/2005.

Extended-spectrum β-lactamase (ESBL)-producing non-typhoidal *Salmonella* isolates have been identified by the EDRU since 2003. In 2003, 28% (452/1 597) of all non-typhoidal *Salmonella* isolates were found to be ESBL producing. The proportion of all non-typhoidal *Salmonella* isolates found to be ESBL producing has decreased to 8% in 2010. ESBL production in non-typhoidal *Salmonella* in South Africa is usually associated with nosocomial isolates of non-typhoidal *Salmonella* and there is co-expression of quinolone and ESBL.

**Salmonella enterica serotype Typhi**

*S. Typhi* bacterium causes typhoid fever and is transmitted via food or water contaminated with human faeces. It is of clinical importance, as humans are the only recognised reservoir of *S. Typhi*. Typhoid fever is a major contributor of illness and death in humans, particularly in developing countries. In 2000 it was estimated that typhoid fever caused approximately 22 million illnesses and 220 000 deaths globally.

**Treatment**

Antibiotics are vital in the management of typhoid fever. Various fluoroquinolones such as ciprofloxacin have become the treatment of choice for infection with *S. Typhi*. However, as with the non-typhoidal *Salmonella*, increased resistance to the quinolone nalidixic acid and reduced susceptibility to the fluoroquinolone ciprofloxacin have been reported.

**Antibiotic resistance**

South Africa, with an estimated typhoid fever burden of disease of 100/100 000 of the population, has not been spared nalidixic-acid-resistant *S. Typhi*. Smith *et al.* reported on 27 nalidixic-acid-resistant isolates collected between 2003 and 2007 that exhibited mutations in both gyrA and topoisomerase IV genes and an active efflux of antibiotic as mechanisms of quinolone resistance. Keddy *et al.* subsequently reported on the first locally isolated strain of fluoroquinolone-resistant *S. Typhi*. The associated mechanism of resistance was the presence of a single amino-acid mutation in the gyrA gene along with a QnrS protein and active efflux of antibiotic. They concluded that the strain was possibly imported through contact with a traveller from the Asian sub-continent.

In the period 2003 - 2010, the EDRU received 706 viable *S. Typhi* isolates that have been screened using antimicrobial agents. Of these 706 viable *S. Typhi* isolates 595 caused invasive disease. The proportion of *S. Typhi* isolates resistant to the older antibiotic ampicillin has fluctuated over this period from 10% in 2003 to a high of 40% in 2006, and 10% at the end of 2010. The proportion of
S. Typhi isolates resistant to sulfamethoxazole remained consistently around 30%. In terms of chloramphenicol, the proportion of S. Typhi isolates identified by the EDRU as resistant has more than doubled from 5% in 2003 to 13% in 2010. The proportion of S. Typhi isolates causing invasive disease resistant to chloramphenicol for the year 2010 was 15%. In 2009, 20% (N=60) of all S. Typhi were resistant to the quinolone nalidixic acid. This proportion of quinolone-resistant S. Typhi isolates has been the highest identified through laboratory surveillance by the EDRU since 2003. In 2003, the proportion of quinolone-resistant S. Typhi was 10%, which decreased to 5% in 2006 and increased to 15% at the end of 2010. Over this same 8-year period, the proportion of ciprofloxacin-resistant S. Typhi was zero, except in 2009 when that proportion rose to 2% with the isolation of the fluoroquinolone-resistant S. Typhi mentioned earlier. Although there have been reports of ESBL-producing S. Typhi, none has been isolated in South Africa to date.13

**Shigella**

Shigellosis is caused by the enteric bacteria *Shigella* species. The disease is a worldwide problem, particularly in areas with poor access to clean water and sanitation, causing an estimated 600 000 deaths annually. As a result *Shigella* is a pathogen associated with water or food contamination as it can easily be spread by the faecal-oral route. The only reservoirs of significance, except for primate colonies, are humans. *Shigella dysenteriae* type 1 is probably the most important *Shigella* variant because it is epidemic-prone and the production of Shiga toxin by this variant of *Shigella* results in severe illness.95 *S. sonnei* has been associated with food- and water-borne outbreaks.

**Treatment**

*Shigella* isolates that are multidrug-resistant to ampicillin, trimethoprim, sulfamethoxazole and tetracycline have become prevalent. As a result, reliance on antibiotic treatment has shifted toward fluoroquinolones such as ciprofloxacin as first-line treatment. Although optimal treatment is to replace fluid and electrolytes, the use of antibiotics to shorten the duration and severity of disease and to decrease the period of pathogen excretion is important.97

**Antibiotic resistance**

From 2003 to 2010, the EDRU received 9 538 viable *Shigella* isolates. Of the 9 538 *Shigella* isolates only 337 caused invasive disease. Antimicrobial screening shows that the proportion of *Shigella* isolates resistant to older antibiotics over the 8-year period has been consistent: 50% for ampicillin, 50% for tetracycline, 50% for sulfamethoxazole and 40% for chloramphenicol. In terms of what has now become first-line treatment, consistently from 2003 to 2010 the proportion of *Shigella* isolates resistant to nalidixic acid has been 1% and for both ciprofloxacin and ceftriaxone the proportion of resistant *Shigella* isolates has been just below 1%. The proportion of *Shigella* isolates exhibiting ESBL production has also consistently been less than 1%. Despite the consistent low levels of resistance to both quinolones and fluoroquinolones, there is concern that the numbers may increase over time.

**Vibrio species**

*Vibrio* spp. are commonly found in aquatic environments and infection occurs as a result of poor access to clean water and sanitation. Of more than 30 species of *Vibrio*, 12 have been associated with illness in humans,89 of which the most important are *V. cholerae* subgroups O1 and O139, the causative agent of epidemic cholera.99 Although infection occurs with non-O1 *V. cholerae* the clinical manifestation is milder because this subgroup of *V. cholerae* lacks the cholera-toxin-producing gene. Pandemics of the devastating diarrhoeal disease caused by *V. cholerae* have been documented since 1817.98,99 Most epidemics occur in developing countries where it is endemic. The debilitating disease caused by *V. cholerae* is the result of an enterotoxin known as choleragen. *V. cholerae* O1 occurs in 3 serotypes (Ogawa, Inaba and Hikojima), and is further characterised into two biotypes – El Tor and classic.99

**Treatment**

Although antimicrobials are prescribed for the management of severe cases, to shorten the duration of illness and reduce the volume of rehydration solution required, *V. cholerae* strains are resistant to a number of antimicrobials including tetracycline, co-trimoxazole, trimethoprim and sulfamethoxazole. Knowledge of the AMR profile of local strains is important for the management of complicated cases, but adequate and timely rehydration therapy remains the gold-standard treatment for cholera.99

**Antibiotic resistance**

In 2008, an outbreak of cholera started in South Africa and continued into 2009. This was linked to cholera in Zimbabwe, with patients crossing the border to seek health care in South Africa. During 2009, the EDRU processed 570 *V. cholerae* O1 isolates associated with the outbreak. Further laboratory characterisation showed that 98% of the isolates were serotype Ogawa and 2% were serotype Inaba; all were biotype El Tor and 99.5% of the isolates were positive for the cholera toxin. The 2008/2009 outbreak isolates showed 100% resistance to co-trimoxazole, 48% resistance to chloramphenicol, 100% resistance to nalidixic acid, 3% resistance to tetracycline and 39% resistance to erythromycin. Although there was 100% resistance to nalidixic acid, none of the isolates associated with this outbreak was resistant to ciprofloxacin.99

In a second outbreak in 2008, reported from Shebagold Mine in the Ehlanzeni district of Mpumalanga, 31 isolates were submitted for analysis to the EDRU. All were biotype El Tor and displayed resistance to ampicillin, amoxycillin-clavulanate, sulfamethoxazole, trimethoprim, chloramphenicol, nalidixic acid, kanamycin, streptomycin and tetracycline, which was initially the antimicrobial agent of choice in the treatment of cholera in Africa. Although the isolates exhibited resistance to nalidixic acid they were susceptible to ciprofloxacin and imipenem. Further resistance to third-generation cephalosporins ceftriaxone and cefazidime was observed, indicative of ESBL activity.101

The EDRU routinely conducts antimicrobial screening on all *V. cholerae* O1 isolates and has data available from 2007. Since 2007, the EDRU has received 899 viable *V. cholerae* O1 isolates. In 2007, 13 of the 30 isolates received were resistant to sulfamethoxazole. The summary of these recent outbreaks is the most accurate description of the current situation of AMR among *V. cholerae* isolates in South Africa.

**Diarrhoeagenic Escherichia coli**

*E. coli* is commonly found in the normal flora of the colon and is used as an indicator of faecal contamination of water. Although a commensal organism, *E. coli* is an important human pathogen that has been associated with several gastro-intestinal syndromes. There are 6 major categories of diarrhoeagenic *E. coli*: enterotoxigenic (ETEC), enteroto-invasive (EIIEC), enteropathogenic (EPEC), enterohaemorrhagic (EHEC), diffusely adherent (DAEC) and entero-aggregative (EAggEC). The most clinically important is EHEC. The strain *E. coli* O157:H7 has been associated with outbreaks and clinical presentation of haemorrhagic diarrhoea, colitis and haemolytic...
uræmic syndrome.102, 103 E. coli O157:H7 produces two cytotoxins, one a verotoxin and the other a toxin identical to the Shiga toxin produced by Shigella dysenteriae type 1. These Shiga-toxin-producing E. coli are referred to as STEC. STECs are not limited to the E. coli O157:H7 serotype, as any of the non-O157:H7 serotypes may present as EHEC or STEC.

**Treatment**

Fluid replacement is recommended as treatment for gastro-enteritis caused by E. coli O157:H7 or non-O157:H7 STEC infection, as it believed (although evidence is lacking) that antimicrobial therapy is of no benefit and may increase the risk of haemolytic uræmic syndrome.103

**Antibiotic resistance**

As part of the EDRU’s surveillance activities, a screening multiplex polymerase chain reaction (M-PCR) analysis is conducted on all E. coli isolates submitted to the unit to categorise the isolate into one of the aforementioned diarrhoeagenic E. coli categories. This is done because antimicrobial screening is conducted only on isolates that are EHEC or STEC. Over the years 2003 - 2010, the EDRU received 3 109 viable E. coli isolates, of which 17 were found to be STEC and 21 to be EHEC by M-PCR. Antibiotic screening of these isolates shows that consistently less than 1% of all STEC or EHEC isolates are resistant to tetracycline, ampicillin, amoxicillin-clavulanate, co-trimoxazole, trimethoprim, sulfamethoxazole and chloramphenicol. The proportion of E. coli isolates showing ESBL activity for the same period was also consistently lower than 1%.

A recent study of clinical isolates of ESBL-producing E. coli isolates screened for ESBL enzymes found that 16 of the 22 isolates were resistant to ciprofloxacin as a result of the presence of aac (6’)-Ib-cr, a variant of an aminoglycoside modifying enzyme.104 Nothing from the EDRU surveillance data suggests that there may be E. coli resistant to the fluoroquinolones, as none was found to be resistant to ciprofloxacin, but these findings should be taken into consideration.

**Sexually transmitted infections**

Bacterial sexually transmitted infections (STIs) cause significant morbidity in South Africa and may rarely cause death, for example from ruptured ectopic pregnancy secondary to tubal damage from Neisseria gonorrhoeae and Chlamydia trachomatis or fetal death from congenital syphilis. They account for 87% of male urethritis syndrome (MUS) cases, 30% of vaginal discharge syndrome (VDS) cases and 10% of genital ulcer syndrome (GUS) cases. Importantly, both ulcerative and genital discharge syndromes are key co-factors for augmenting HIV infectiousness and susceptibility and increase transmission risk by 2 - 5 times in prospective studies.105 Patients with bacterial STIs may present with MUS, VDS, scrotal swelling syndrome (SSW, i.e. epididymo-orchitis), lower abdominal pain syndrome (LAP, i.e. pelvic inflammatory disease), GUS or buboes. As the syndromic management approach does not utilise laboratory testing, it is not possible to determine the total burden of bacterial STIs by individual STI pathogen. The bacterial burden also differs according to STI syndrome; recent aetiological surveillance data from South Africa showed that bacteria account for 87% of cases of MUS, 30% of cases of VDS and only 10% of GUS cases (Table III).

Between April 2004 and March 2005, 1 654 776 new STI episodes were treated in primary health care (PHC) clinics throughout South Africa. Incidence rates of new STI syndrome episodes, calculated per 1 000 population aged 15 - 49 years, demonstrated a national incidence rate of 63 per 1 000 population. The highest incidence rates were recorded in Limpopo (90 per 1 000), KwaZulu-Natal (87 per 1 000) and the Eastern Cape (73 per 1 000); the lowest incidence rate was recorded in the Western Cape (38 per 1 000). During the same time period, a total of 145 818 new STI syndrome episodes (46 222 in males, 99 596 in females, 8.8% of the national total) were reported among 126 656 patients in the sentinel survey, with a peak in the 20 - 24-year-old age group. In men with STIs, the most frequent syndromes were MUS and GUS, whereas for women they were VDS and LAP (Fig. 1). The relative prevalence and incidence of MUS, the most reliable indicator syndrome for ‘true’ STIs, seen at the sentinel sites during 2004 - 2005, is shown by province in Table IV.

**Neisseria gonorrhoeae**

At present in South Africa, AMR is solely an issue for N. gonorrhoeae infection. It is very important to have effective microbiological

| Table III. Bacteria causing the most prevalent STI syndromes in South Africa |
|-----------------------------|-----------------------------|-----------------------------|
| Category                    | Male urethritis syndrome (MUS) | Vaginal discharge syndrome (VDS) | Genital ulcer syndrome (GUS) |
| No. of enrolled cases       | 1 593 (100)                  | 1 462 (100)                  | 597 (100)                     |
| No. of bacterial cases      | 1 378 (87)                   | 423 (30)                     | 60 (10)                       |
| Bacterial aetiologies for MUS/VDS |                       |                                |                                |
| Neisseria gonorrhoeae       | 1 155 (73)                   | 180 (12)                     | NA                            |
| Chlamydia trachomatis       | 287 (18)                     | 203 (14)                     | NA                            |
| Mycoplasma genitalium       | 134 (8)                      | 144 (100)                    | NA                            |
| Bacterial aetiologies for GUS |                         |                                |                                |
| Treponema pallidum          | NA                           | NA                           | 44 (7)                        |
| Haemophilus ducreyi         | NA                           | NA                           | 5 (1)                         |
| Chlamydia trachomatis L1-L3 | NA                           | NA                           | 6 (1)                         |
| Klebsiella granulomatis     | NA                           | NA                           | -                             |

surveillance systems in place in South Africa and its neighbouring countries to facilitate early detection of such strains. There is mounting public health concern that gonorrhoea may become untreatable in years to come, which would have an extremely deleterious effect on HIV transmission in South Africa, where the prevalence of both diseases is high. Accordingly, efforts must be made locally to reduce the burden of gonorrhoea and for the international community to invest in the search for a new class of antimicrobial agents active against *N. gonorrhoeae*.

### Treatment

In South Africa, STIs have been treated using the syndromic management approach since the late 1990s. This approach is to manage symptomatic STIs and has the advantage of providing same-day treatment according to treatment flow charts, which can easily be adhered to by nursing staff at every PHC entry point across the country. Laboratory testing of STI patients is not required for case management, although the WHO recommended that periodic aetiological and AMR surveys are carried out in all countries using the approach. Lack of clinical samples has deskilled laboratory staff in terms of ability to culture and test gonococci for antimicrobial susceptibility. The syndromic approach generally works better for male-associated compared with female-associated STI syndromes. The poor specificity of syndromes such as VDS and LAP to predict the presence of STIs leads to overdiagnosis of STIs, unnecessary stigmatisation and potential relationship difficulties. Importantly, it results in substantial overprescribing of antimicrobial agents that may influence the development of AMR among sexually transmitted and non-sexually transmitted bacteria. Mathematical modelling has shown that syndromic management is the cheapest programmatic approach to the management of STIs, although there remains debate as to whether it is the most cost-effective.

Owing to the rapid emergence of quinolone-resistant gonococci in 2003, and their subsequent spread throughout the country, revised national guidelines were published in 2008. Gonorrhoea should now be treated with oral cefixime or intramuscular ceftriaxone. Gonococci exhibiting clinical resistance to oral cephalosporins have emerged in the Western Pacific region and have now spread to Europe. No such isolates have been found in Africa to date, but their emergence is likely in the near future. Other key changes include use of acyclovir in the GUS treatment algorithm and the replacement of erythromycin with amoxicillin for the treatment of presumptive chlamydial infection in pregnant women with VDS.

At least half of STI care episodes are estimated to be managed by the private sector, where the National Department of Health (NDoH) has less influence on prescribing practice. An interview-based study conducted among general practitioners (GPs) in Gauteng over a decade ago highlighted poor knowledge of STI syndromic

### Table IV. Male urethritis syndrome (MUS) indicators by province, primary health care

<table>
<thead>
<tr>
<th>Province</th>
<th>New episodes (N)</th>
<th>Relative prevalence of MUS (%)</th>
<th>Incidence rate per 1 000 population aged 15 - 49 (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eastern Cape</td>
<td>60 147</td>
<td>25.6</td>
<td>40.8 (39.8 - 41.8)</td>
</tr>
<tr>
<td>Free State</td>
<td>20 533</td>
<td>25.1</td>
<td>28.6 (28.2 - 29.0)</td>
</tr>
<tr>
<td>Gauteng</td>
<td>61 139</td>
<td>23.7</td>
<td>19.4 (18.9 - 19.9)</td>
</tr>
<tr>
<td>KwaZulu-Natal</td>
<td>121 972</td>
<td>26.7</td>
<td>50.2 (49.3 - 51.0)</td>
</tr>
<tr>
<td>Limpopo</td>
<td>59 409</td>
<td>24.6</td>
<td>50.1 (48.9 - 51.4)</td>
</tr>
<tr>
<td>Mpumalanga</td>
<td>40 227</td>
<td>39.5</td>
<td>47.9 (47.4 - 48.3)</td>
</tr>
<tr>
<td>North West</td>
<td>36 394</td>
<td>24.1</td>
<td>33.5 (32.4 - 34.5)</td>
</tr>
<tr>
<td>Northern Cape</td>
<td>7 364</td>
<td>32.7</td>
<td>33.7 (33.0 - 34.5)</td>
</tr>
<tr>
<td>Western Cape</td>
<td>32 062</td>
<td>30.1</td>
<td>23.5 (22.7 - 24.3)</td>
</tr>
<tr>
<td>National</td>
<td>439 247</td>
<td>26.5</td>
<td>35.2 (34.2 - 36.3)</td>
</tr>
</tbody>
</table>

Note: The denominator for the relative prevalence of MUS includes males and females.
Source: Report on the National Clinical Sentinel Surveillance of Sexually Transmitted Infections at Public Sector Primary Health Care Facilities (2005), prepared by the STI Reference Centre (NICD/NHLS) for the National Department of Health.
management, and less than half of prescriptions overall were judged to be effective. In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient antibiotic regimens. Prescribing correct drug treatment for STIs by GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid.

In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient to be effective. In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient to be effective. In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient to be effective. In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient to be effective. In addition, for most STI syndromes, uninsured patients were offered significantly cheaper and less convenient to be effective.

A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid. A study of GPs has been associated with male gender and recent graduation of the GP, as well as the patient having medical aid.

To make matters worse, at the time that the national STI guidelines were changed, the NDHiH had to purchase cefixime directly from Merck in Germany, and it was only made available at PHCs. This led to an inequality in the health care system, where cefixime was available to patients with presumptive gonorrhoea attending public clinics whereas similar patients attending tertiary-level hospital or GP facilities could only be treated with cefixime. Cefixime was finally made accessible to all practitioners for the treatment of gonorrhoea at the start of 2011.

Antibiotic resistance

The need for periodic aetiological and AMR surveillance, which is an integral part of syndromic management, has been largely ignored by most African countries. With the exception of South Africa, where good laboratory infrastructure and funding exist to support surveillance, Africa has minimal AMR data available for bacterial STI pathogens. Gonorrhoea is the only bacterial STI for which AMR surveys are currently undertaken in South Africa. Despite reports concerning AMR in chlamydial strains collected from patients failing treatment, it remains controversial whether documented stable homotypic drug resistance to antibiotics exists and AMR studies are not routinely performed for this STI pathogen anywhere in the world. Nevertheless, although a high prevalence of tetracycline resistance has been documented among Mycoplasma genitalium isolates, testing for this relatively new bacterial STI pathogen is performed in few specialist laboratories worldwide. Screening for resistance in Treponema pallidum remains a challenge because of inability to culture this organism in vitro. Although resistance of T. pallidum to penicillin has not been described to date, and a molecular assay for the macrolide resistance-associated A2058G mutation in 23S rRNA does exist, the STI Reference Centre has failed to detect this A2058G mutation in T. pallidum-positive DNA extracts from genital ulcer swabs recently collected in South Africa (D A Lewis and E E Müller, unpublished data). Chancroid is now a rare cause of GUS, and it is no longer feasible to culture isolates to determine AMR. Chancroid was the most frequent cause of GUS in the 1990s and surveys performed at that time reported that most strains were resistant to penicillin, co-trimoxazole and tetracyclines but susceptible to amoxicillin-clavulanate, macrolides, quinolones and extended-spectrum cephalosporins.

Gonococci isolated in South Africa remained fully susceptible to ciprofloxacin, the former first-line therapy used to treat gonorrhoea, until 2003 when researchers from the University of KwaZulu-Natal reported the abrupt emergence of quinolone-resistant N. gonorrhoeae (QRNG) among MUS patients attending an STI clinic in Durban. Subsequently, the NDHiH requested that the STI Reference Centre co-ordinate a gonococcal resistance survey in several South African cities, which included Cape Town, Durban, Johannesburg, Pietermaritzburg, Pretoria and Mthatha. The data revealed varying prevalence of QRNG, from 0% in Pretoria to 24% in Durban, although all isolates tested appeared susceptible to cephalosporins. Despite the widespread problem with QRNG, revised national guidelines were not published until 2008, at which point ciprofloxacin was replaced by either cefixime or ceftriaxone as first-line therapy for presumptive gonococcal infection. During this 4-year period, further rises in QRNG prevalence was reported from Durban (24% in 2004; 42% in 2005), Pretoria (0% in 2004, 7% in 2005), Cape Town (7% in 2004; 27% in 2007) and Johannesburg (11% in 2004; 32% in 2007). The STI Reference Centre has conducted additional surveys in Kimberley (2006), Bloemfontein (2008), East London (2010), Rustenburg (2011) and Polokwane (2011), and observed a QRNG prevalence of 53%, 16%, 41%, 15% and 40% respectively (D A Lewis, unpublished data).

There is substantial public health concern about the global spread of gonococci with decreased susceptibility to oral cephalosporins which have resulted in gonorrhoea treatment failures in several countries, including Japan, China, Australia, Norway and the UK. Japan, China and Australia therefore now use intramuscular ceftriaxone to treat gonorrhoea. To date there has been no confirmed case of clinical failure with oral cephalosporins in Africa, but such strains will undoubtedly emerge over time, either through importation or de novo. All gonococci tested in South African surveys carried out by the STI Reference Centre (STIRC) over the past 5 years have remained fully susceptible to both cefixime and ceftriaxone (D A Lewis, unpublished data).

In terms of other antimicrobials, studies from Gauteng have confirmed that tetracyclines and penicillin should not be used to treat gonorrhoea in South Africa because of a high prevalence of plasmid-mediated tetracycline resistance (36 - 74%) and a lower, but still unacceptably high, prevalence of penicillinase-producing gonococci (16 - 26%). Gonococci isolated in Johannesburg in 2008 demonstrate no resistance as yet to azithromycin, spectinomycin and gentamicin (D A Lewis, unpublished data).

Where bacterial STI pathogens are resistant to treatment, patients may be at increased risk of pathogen-associated complications, such as epididymo-orchitis or pelvic inflammatory disease in the case of antimicrobial-resistant N. gonorrhoeae. From the public health viewpoint, such patients also remain infectious to others for longer and this may increase transmission of the pathogen within the community. STIs are also important co-factors in HIV transmission, and HIV viral loads are increased in cervicovaginal, seminal and ulcer-derived secretions in the presence of other STIs. In the case of gonorrhoea, for example, studies from Malawi demonstrated that urethritis can elevate the seminal HIV viral load approximately 8 times and, even with effective anti-gonococcal treatment, it may take over 3 weeks for the seminal viral loads to decline to levels seen in HIV-infected dermatology patients (controls). The risk of HIV transmission may be much greater in HIV-infected individuals with antimicrobial-resistant gonorrhoea, particularly in a country like South Africa where there are an estimated 5.3 million HIV-infected adults aged 15 years and older. Relevant to this argument, the STI Reference Centre demonstrated that the detection of QRNG in men with MUS in Cape Town and Johannesburg was significantly associated with co-infection with HIV.

Finally, treating patients with resistant STIs will require use of more expensive antimicrobial agents and also, when gonococcal resistance to oral cephalosporins emerges in South Africa, increased use of injectable antimicrobials such as ceftriaxone, spectinomycin or gentamicin. The widespread use of intramuscular antimicrobial agents to treat index STI patients and their partner(s) may have a deleterious public health effect by reducing patient and sexual partner access because of fears concerning injections. Widespread
use of intramuscularly administered antimicrobials also heightens the risk of needle-stick injuries for staff working with STI patients, who are at high risk of being HIV infected.

**Hospital-acquired infections**

**Public sector**

According to the 2009 National Health Laboratory Service (NHLS) public sector susceptibility data (Table V), *K. pneumoniae* remains

<table>
<thead>
<tr>
<th>Laboratories</th>
<th>GSH</th>
<th>TBH</th>
<th>GP</th>
<th>UNI*</th>
<th>DGM</th>
<th>SBAH</th>
<th>CMJAH</th>
<th>CHBH</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Klebsiella pneumoniae from blood cultures</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N = \text{total of isolates})</td>
<td>325</td>
<td>190</td>
<td>113</td>
<td>89</td>
<td>112</td>
<td>440</td>
<td>258</td>
<td>388</td>
</tr>
<tr>
<td>Gentamicin (% susceptible)</td>
<td>32</td>
<td>42</td>
<td>41</td>
<td>49</td>
<td>63</td>
<td>48</td>
<td>51</td>
<td>39</td>
</tr>
<tr>
<td>Amikacin (% susceptible)</td>
<td>70</td>
<td>87</td>
<td>76</td>
<td>90</td>
<td>98</td>
<td>64</td>
<td>63</td>
<td>59</td>
</tr>
<tr>
<td>Ciprofloxacin (% susceptible)</td>
<td>54</td>
<td>60</td>
<td>67</td>
<td>61</td>
<td>80</td>
<td>59</td>
<td>66</td>
<td>72</td>
</tr>
<tr>
<td>ESBL (% susceptible)</td>
<td>71</td>
<td>64</td>
<td>56</td>
<td>53</td>
<td>46</td>
<td>60</td>
<td>50</td>
<td>62</td>
</tr>
<tr>
<td>Ertapenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>96</td>
<td>99</td>
<td>100</td>
<td>100</td>
<td>98</td>
</tr>
<tr>
<td>Imipenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Meropenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td><strong>Escherichia coli from blood cultures</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N = \text{total of isolates})</td>
<td>281</td>
<td>131</td>
<td>135</td>
<td>40</td>
<td>62</td>
<td>193</td>
<td>219</td>
<td>417</td>
</tr>
<tr>
<td>Ciprofloxacin (% susceptible)</td>
<td>80</td>
<td>83</td>
<td>93</td>
<td>70</td>
<td>81</td>
<td>92</td>
<td>83</td>
<td>78</td>
</tr>
<tr>
<td>Gentamicin (% susceptible)</td>
<td>83</td>
<td>82</td>
<td>84</td>
<td>90</td>
<td>84</td>
<td>91</td>
<td>78</td>
<td>76</td>
</tr>
<tr>
<td>Amikacin (% susceptible)</td>
<td>88</td>
<td>96</td>
<td>94</td>
<td>98</td>
<td>95</td>
<td>94</td>
<td>69</td>
<td>78</td>
</tr>
<tr>
<td>ESBL (% susceptible)</td>
<td>10</td>
<td>11</td>
<td>10</td>
<td>13</td>
<td>16</td>
<td>6</td>
<td>8</td>
<td>48</td>
</tr>
<tr>
<td>Ertapenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Imipenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Meropenem (% susceptible)</td>
<td>100</td>
<td>100</td>
<td>-</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td><strong>Pseudomonas aeruginosa from blood cultures</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N = \text{total of isolates})</td>
<td>94</td>
<td>44</td>
<td>15</td>
<td>14</td>
<td>30</td>
<td>134</td>
<td>93</td>
<td>152</td>
</tr>
<tr>
<td>Gentamicin (% susceptible)</td>
<td>66</td>
<td>61</td>
<td>80</td>
<td>64</td>
<td>93</td>
<td>48</td>
<td>84</td>
<td>72</td>
</tr>
<tr>
<td>Cefipime (% susceptible)</td>
<td>51</td>
<td>64</td>
<td>80</td>
<td>71</td>
<td>90</td>
<td>52</td>
<td>81</td>
<td>79</td>
</tr>
<tr>
<td>Pip-taz (% susceptible)</td>
<td>40</td>
<td>43</td>
<td>40</td>
<td>79</td>
<td>97</td>
<td>60</td>
<td>90</td>
<td>74</td>
</tr>
<tr>
<td>Ciprofloxacin (% susceptible)</td>
<td>57</td>
<td>68</td>
<td>73</td>
<td>79</td>
<td>100</td>
<td>52</td>
<td>82</td>
<td>84</td>
</tr>
<tr>
<td>Ceftazidime (% susceptible)</td>
<td>66</td>
<td>82</td>
<td>93</td>
<td>86</td>
<td>100</td>
<td>57</td>
<td>85</td>
<td>79</td>
</tr>
<tr>
<td>Imipenem (% susceptible)</td>
<td>65</td>
<td>52</td>
<td>13</td>
<td>79</td>
<td>100</td>
<td>48</td>
<td>77</td>
<td>74</td>
</tr>
<tr>
<td>Meropenem (% susceptible)</td>
<td>66</td>
<td>70</td>
<td>13</td>
<td>86</td>
<td>97</td>
<td>52</td>
<td>78</td>
<td>75</td>
</tr>
<tr>
<td><strong>Acinetobacter from blood cultures</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N = \text{total of isolates})</td>
<td>241</td>
<td>175</td>
<td>21</td>
<td>22</td>
<td>38</td>
<td>173</td>
<td>98</td>
<td>323</td>
</tr>
<tr>
<td>Pip-taz (% susceptible)</td>
<td>20</td>
<td>8</td>
<td>38</td>
<td>9</td>
<td>89</td>
<td>20</td>
<td>41</td>
<td>14</td>
</tr>
<tr>
<td>Ciprofloxacin (% susceptible)</td>
<td>57</td>
<td>30</td>
<td>71</td>
<td>14</td>
<td>18</td>
<td>26</td>
<td>40</td>
<td>37</td>
</tr>
<tr>
<td>Ceftazidime (% susceptible)</td>
<td>57</td>
<td>43</td>
<td>67</td>
<td>0</td>
<td>24</td>
<td>27</td>
<td>42</td>
<td>50</td>
</tr>
<tr>
<td>Imipenem (% susceptible)</td>
<td>26</td>
<td>9</td>
<td>43</td>
<td>18</td>
<td>92</td>
<td>32</td>
<td>42</td>
<td>21</td>
</tr>
<tr>
<td>Meropenem (% susceptible)</td>
<td>25</td>
<td>9</td>
<td>43</td>
<td>14</td>
<td>79</td>
<td>32</td>
<td>32</td>
<td>27</td>
</tr>
<tr>
<td><strong>Staphylococcus aureus from blood cultures</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(N = \text{total of isolates})</td>
<td>250</td>
<td>175</td>
<td>121</td>
<td>41</td>
<td>94</td>
<td>476</td>
<td>228</td>
<td>411</td>
</tr>
<tr>
<td>Cloxacillin (% susceptible)</td>
<td>65</td>
<td>69</td>
<td>74</td>
<td>71</td>
<td>16</td>
<td>63</td>
<td>57</td>
<td>76</td>
</tr>
<tr>
<td>Erythromycin (% susceptible)</td>
<td>69</td>
<td>70</td>
<td>83</td>
<td>66</td>
<td>11</td>
<td>56</td>
<td>56</td>
<td>75</td>
</tr>
<tr>
<td>Clindamycin (% susceptible)</td>
<td>70</td>
<td>70</td>
<td>85</td>
<td>68</td>
<td>28</td>
<td>58</td>
<td>65</td>
<td>74</td>
</tr>
</tbody>
</table>

*Data for Universitas are incomplete for certain organisms.

NHLS = National Health Laboratory Service; ESBL = extended-spectrum β-lactamase; GSH = Groote Schuur Hospital; TBH = Tygerberg Hospital; GP = Green Point NHLS Laboratory, Cape Town; UNI = Universitas Hospital, Bloemfontein; DGM = Dr George Mukhari Hospital, Pretoria; SBAH = Steve Biko Academic Hospital, Pretoria; CMJAH = Charlotte Maxeke Johannesburg Academic Hospital; CHBH = Chris Hani Baragwanath Hospital; Pip-taz = piperacillin-tazobactam.
a highly resistant nosocomial pathogen, with more than 50% of all strains producing ESBLs. These isolates were frequently multiresistant, with only 32 - 63% susceptible to gentamicin and 54 - 80% susceptible to ciprofloxacin.

*E. coli* strains exhibited less resistance than *K. pneumoniae*, with 76 - 91% susceptible to gentamicin, 78 - 92% susceptible to ciprofloxacin and only 6 - 16% producing ESBLs. The very high rate of ESBL production (48%) at Chris Hani Baragwanath Hospital (CHBH) remains unexplained.

Patterns of resistance among *P. aeruginosa* isolates vary between laboratories. Cefazidime remains the most active agent.

Carcapenem resistance among *Acinetobacter* spp. is common in the 5 hospitals with major intensive care units, with only 20 - 40% of isolates being susceptible to carbapenems. Approximately 60% of *S. aureus* isolates from blood are sensitive to cloxacinol.

### Private sector

For several reasons, including selective pressure from overuse of antibiotics and failure of hospital infection control practices, the incidence of colonisation and infection, particularly with resistant Gram-negative bacteria, in South African private institutions appears to be increasing. In addition, the worldwide emergence and spread of carbapenem-resistant *K. pneumoniae* and *E. coli* and reports of hospital outbreaks owing to such strains is cause for local concern. Increased use of carbapenems in the private sector in South Africa is driven by an increase in cephalosporin and fluoroquinolone resistance among ESBL-producing Enterobacteriaceae. Although extensive published data regarding antibiotic susceptibility of community-acquired respiratory tract pathogens especially *S. pneumoniae* are available, including those of invasive isolates, few data have been published for Gram-negative pathogens such as *A. baumannii* or *P. aeruginosa* or for Gram-positive pathogens, particularly *S. aureus*.

The SENTRY international antimicrobial surveillance programme documented the prevalence of ESBL production in Enterobacter cloacae among hospitalised patients in several Johannesburg private hospitals as 20% (N=1154) and that of oxacillin resistance in blood isolates of *E. coli* (N=471) was 84%, and 20% were resistant to fluoroquinolones (Table VI). Cephalosporin resistance among isolates of *K. pneumoniae* (N=636) was high; 52% were resistant to cefuroxime. The most active agents in *Enterobacter* spp. (N=242) were imipenem/meropenem, ertapenem, ciprofloxacin and levofloxacin, with 100%, 94%, 88% and 87% susceptibility, respectively. Carbapenem resistance in invasive isolates of *P. aeruginosa* (N=382) varied between 45% and 42% for imipenem and meropenem and in *A. baumannii* (N=190) between 33% and 32%, respectively. The overall incidence of methicillin resistance among *S. aureus* isolates was 36% (N=629). The prevalence of ESBL production among all-source isolates of *K. pneumoniae* (N=7514), *Enterobacter* spp. (N=4 031) and *E. coli* (N=28 412) was 26%, 12% and 5%, respectively.

### Table VI. Incidence (%) of ESBL production (number of isolates) in selected strains of Enterobacteriaceae in private practice in South Africa (all sources), January - June 2006

<table>
<thead>
<tr>
<th>City</th>
<th><em>K. pneumoniae</em></th>
<th>Enterobacter spp.</th>
<th><em>E. coli</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>26 (7 514)</td>
<td>12 (4 031)</td>
<td>5 (28 412)</td>
</tr>
<tr>
<td>Johannesburg</td>
<td>42 (3 010)</td>
<td>11 (1 486)</td>
<td>4 (12 600)</td>
</tr>
<tr>
<td>Pretoria</td>
<td>27 (2 244)</td>
<td>10 (1 061)</td>
<td>3 (7 406)</td>
</tr>
<tr>
<td>Durban</td>
<td>8 (1 359)</td>
<td>5 (1 093)</td>
<td>4 (5 637)</td>
</tr>
<tr>
<td>Cape Town</td>
<td>40 (805)</td>
<td>27 (328)</td>
<td>4 (1 380)</td>
</tr>
<tr>
<td>Bloemfontein</td>
<td>15 (96)</td>
<td>6 (63)</td>
<td>12 (3 189)</td>
</tr>
</tbody>
</table>

ESBL = extended-spectrum β-lactamase.

### References


