Growing resistance to antibacterial agents has increased the need for the development of new drugs to treat bacterial infections. Given increasing pressure on limited health budgets, it is important to study the cost-effectiveness of these drugs, as well as their safety and efficacy, to find out whether or not they provide value for money and should be reimbursed. In this article, we systematically reviewed 38 cost-effectiveness analyses of new antibacterial agents. Most studies showed the new antibacterial drugs were cost-effective compared to older generation drugs. Drug pricing is a complicated process, involving different stakeholders, and has a large influence on cost-effectiveness. Value-based pricing is a method to determine the price of a drug at which it can be cost-effective. It is currently unclear what the influence of value-based pricing will be on the prices of new antibacterial agents, but an important factor will be the definition of ‘value’, which as well as the impact of the drug on patient health might also include other factors such as wider social impact and the health impact of disease.

Key words: antibacterial drugs, antibiotics, cost-effectiveness, health economics, value-based pricing

Since the discovery of the first antibacterial agents in the 1930s, many new antibacterial drugs have been developed. This has had a large influence on health because of the decreased mortality and morbidity of bacterial infections (1,2). Between the 1970s and the 1990s, few new antibacterial drug classes were launched (3,4). Pharmaceutical companies were reluctant to invest in the development of antibacterial agents; the revenues of the new drugs were expected to be low, because of the short use of antibacterial drugs and the high competition with many cheap generic drugs (3,5). The lack of new antibacterial agents and the growing resistance to the available drugs has limited the treatment options for infection in recent years. However, since 2000, a series of new classes of antibacterial drugs such as oxazolidinones and lipopeptides have been launched (2–4). Increasingly, cost-effectiveness is an important factor affecting the reimbursement of new drugs, and it is important to investigate whether these drugs provide value for money. In this article, we will describe and explain some basic concepts of cost-effectiveness and review the available evidence on the cost-effectiveness of new antibacterial agents. Also, we will provide a brief historical overview of drug pricing (in the UK) and describe the role of value-based pricing in determining the price of new antibacterial agents.

What is Cost-effectiveness Analysis?

In a cost-effectiveness analysis (CEA), the total costs and effects of two or more treatment options are compared. When calculating costs, not only should the costs of the drugs be considered, including administration costs and costs of treating adverse drug reactions, but also other costs related to the treatment or disease, such as hospitalization costs. To study cost-effectiveness, both incremental (extra) costs and incremental effects are calculated for the new treatment option compared to the comparator option, which might be current best practice and which might be ‘do nothing’. Figure 1 depicts the possible results of a CEA. Compared with the comparator option, the new treatment may be more or less costly and more or less effective. When the new treatment leads to increased effects while decreasing costs (bottom right quadrant), it is the dominant treatment option, which means that it is more attractive than the comparator on economic grounds. When the new treatment leads to decreased effect while increasing costs (top left quadrant), it is dominated by the comparator, which means that the comparator is a more attractive option. When both costs and effects are increased (top right quadrant), the attractiveness of the new treatment depends on how much payers are prepared to pay for the extra effect. When the incremental costs per extra unit of effect are lower than the willingness-to-pay threshold (top right quadrant below the 45 degree line), the new treatment is cost-effective. Conversely, when the incremental costs per extra unit of effect are higher than the threshold (top right quadrant above the 45 degree line), the new treatment is not cost-effective. When costs and effects are higher with a new drug compared with the comparator, the incremental cost-effectiveness ratio (ICER) can be calculated as shown in the equation below.
Calculation of the ICER

\[ \text{ICER} = \frac{\text{Cost of new treatment} - \text{cost of comparator}}{\text{Effect of new treatment} - \text{effect of comparator}} \]

Effectiveness can be measured in many ways usually related to the treatment goal of the drug (e.g. cure rate). When effectiveness is measured in terms of cure rates, cost-effectiveness measured using the ICER is expressed as the incremental cost per extra case cured. This outcome measure is, however, very disease-specific, and therefore, it is difficult to compare with treatments in other diseases. A recommended measure is quality-adjusted life-years (QALYs), which account for both quality and quantity of life. There are various ways of measuring the quality of life, including direct valuation methods such as standard gamble, time trade-off and visual analogue scale (6) or questionnaires such as the EQ-5D (7). When effects are measured by QALYs, the ICER is the incremental cost per QALY gained, and as this is a generic health outcome measure, it can facilitate comparisons between treatments for different diseases. For this reason, some regulatory authorities or health insurance companies require evidence of a favourable incremental cost per QALY gained before they approve the use of a new drug. Another generic measure is life-years gained which is based on life expectancy only and does not take into account quality of life. This measure is easier to calculate, because it does not require information on health-related quality of life impacts, but it is not as comprehensive as QALYs.

Whether or not a new treatment is cost-effective depends on the decision-makers’ willingness to pay for a QALY, which is different across different settings. In the USA, a threshold of US$50 000–100 000 per QALY gained is commonly reported (8); in the UK, a range of £20 000–30 000 is used (9). Two important issues when considering cost-effectiveness of a new intervention are perspective and time horizon. The perspective is the viewpoint from which the analysis is performed (e.g. patient, hospital, healthcare system, society), and this influences the type of costs that should be collected (healthcare costs, patient-borne costs, costs borne by the rest of society, etc.). The time horizon should reflect the period over which the main differences between two interventions are expected. In many cases, this is a lifetime horizon. When the time horizon is more than 1 year, discounting of costs and effects is usually recommended, which has the effect of giving less weight to costs and benefits that occur in the future. The recommended discount rates for costs and effects vary by country, for example, in the UK the discount rates for both costs and effects should be 3.5% and in the Netherlands the discount rate for costs should be 4% and the discount rate for effects only 1.5% (10).

Every CEA has some level of uncertainty around the results. This can be because of uncertainty around the effectiveness estimate or around the costs. There may also be variability in costs and effects between patients, for example, because of differences in severity of the disease. This uncertainty should be assessed in a sensitivity analysis. In a one-way sensitivity analysis, the value of one parameter is varied over a plausible range, to show the effect of this parameter on the cost-effectiveness results. This can be performed for several or all parameters, varying only one at a time. It is the simplest form of sensitivity analysis and is useful to identify the parameters with the largest influence on the results. It is, however, not regarded as sufficient because it does not take into account any combined effect of parameters. In a two-way (or multiway) sensitivity analysis, two (or several) parameters are varied at the same time to assess their joint influence on the results.

Probabilistic sensitivity analysis is a technique that can be used to investigate joint parameter uncertainty. It can be used to generate cost-effectiveness acceptability curves, showing the chance that a drug would be cost-effective at a range of cost-effectiveness threshold values. This can be performed by carrying out a large number of simulations, drawing random samples from probability distributions for the ranges of key parameters. The number of simulations below the cost-effectiveness threshold represents the chance that the drug would be cost-effective given that threshold.

Other forms of sensitivity analysis include threshold analysis (identifying a threshold value for a parameter at which the new treatment would just be cost-effective) and scenario analysis (for example, showing a best-case scenario and a worst-case scenario).
Cost-effectiveness of New Antibacterial Agents

Since 2000, 22 new antibacterial agents have become available on the market (Box 1) (2). In this section, we will review the available evidence on the cost-effectiveness of these new drugs, in comparison to the older antibacterial agents. Using the NHS Economic Evaluation Database (NHS EED), we performed an initial search for CEAs of all 22 new drugs. Using the drugs in Box 1 as search terms, we identified 41 records. Of these, 12 were excluded, three studies were published in a non-English language (9 records. Of these, 12 were excluded, three studies were published in a non-English language (9–11), one did not concern any of the new drugs (12) and eight did not link costs and effects explicitly (13–20). Hence, an initial 29 studies were included for review (21–48, 58). We then undertook a more extensive search using Embase and PubMed using the same search terms including cost-effectiveness terms and found nine additional papers (49–57). Only studies reporting both costs and effects of the new drug compared to the comparator were included. Conference abstracts and non-English papers were excluded. From the papers included for review, we collected the following information: year of study, comparator, disease, outcome measure, whether sensitivity analysis was performed to assess uncertainty and which type of sensitivity analysis and final result (cost-effective or not). We used the PRISMA guidelines for systematic searching.

Several CEAs have been published on linezolid, which was launched in 2000. Other drugs that have been evaluated are daptomycin, telithromycin, ertapenem, gemifloxacin, doripenem, telavancin and fidaxomicin (see Table 1).

Cost-effectiveness was always assessed for one specific indication, such as pneumonia or skin infections caused by methicillin-resistant staphylococcus aureus (MRSA) or by other bacteria. Most of the published studies assessed the incremental cost per extra case cured with the new antibacterial drug. When both the costs and the cure rate of the new drug were higher than the comparator (top right quadrant in Figure 1) (21–23,36,56), it was difficult to assess whether the new drug was cost-effective or not, because there is no willingness-to-pay threshold for cost per extra case cured. In many studies, the new drugs were more effective, and total costs were lower, which means that the new treatment was the dominant treatment (bottom right quadrant in Figure 1). In this case, we can conclude that the new drug was cost-effective. In some studies, the authors found that the effectiveness of the new drug and the comparator were equal, and therefore, they performed a cost minimization study, only looking at the cost of the drug versus the comparator.

Nine studies assessed the incremental cost per QALY gained for linezolid, daptomycin, ertapenem, doripenem and fidaxomicin (32,33,43,45,48,49,51,52,58). In seven of these studies, the ICER was below the willingness-to-pay threshold. In one of these studies, the incremental cost per QALY gained of fidaxomicin compared to vancomycin was US$67 576 (45). The authors applied a willingness-to-pay threshold of US$100 000, so this drug was considered cost-effective for the treatment of Clostridium difficile infections.

Most studies showed that the new drugs were cost-effective, except in four cases. Both linezolid and daptomycin were dominated by trimethoprim/sulfamethoxazole for the treatment of MRSA infections (25), and telithromycin was dominated by moxifloxacin for the treatment of pneumonia (41). In two studies, on incremental cost per QALY gained, fidaxomicin was dominated by one or more of the comparators (51,58). All studies correctly considered not only the costs of the drug itself, but also other costs, such as hospitalization costs, which can affect the ICER.

Because uncertainty is present in every cost-effectiveness study, a good economic evaluation should perform a sensitivity analysis to examine the robustness of the model and assumptions. In some studies, no sensitivity analysis was performed, but most studies included at least some form of sensitivity analysis. Almost half of the studies included a probabilistic sensitivity analysis.

Some antibacterial agents can be used for various indications. The cost-effectiveness, however, has mostly been studied for one indication at a time. The cost-effectiveness of the drug can vary between different indications, because of different bacteria causing the infection or because of differences in location of the bacteria. It is therefore necessary to assess the cost-effectiveness of a new drug for a specific indication, even if the drug can be used for more than one indication. The cost-effectiveness

Box 1: New antibiotics launched since 2000 (2)

| Year | Drug                  |
|------|-----------------------|
| 2000 | Linezolid             |
| 2002 | Biapenem              |
| 2003 | Daptomycin            |
| 2004 | Gemifloxacin          |
| 2005 | Doripenem             |
| 2006 | Telithromycin         |
| 2007 | Ertapenem             |
| 2008 | Prulifloxacin         |
| 2009 | Pazufloxacin          |
| 2010 | Balofloxacin          |
| 2011 | Fidaxomicin           |
| 2012 | Bedaquiline           |
### Table 1: Reviewed cost-effectiveness studies on antibacterial agents

| Year  | Comparator                          | Disease                          | Outcome measure                  | Uncertainty analysis | Cost-effective? | Ref |
|-------|-------------------------------------|----------------------------------|----------------------------------|----------------------|-----------------|-----|
| 2001  | Flucloxacillin or vancomycin        | Cellulitis                        | Cost per extra cure/success      | 1-way                | Yes             | (55) |
| 2003  | Oxacillin or vancomycin             | Cellulitis                        | Cost per extra cure/success      | 1-way                | Yes             | (47) |
| 2004  | Vancomycin                          | Pneumonia                         | Cost per QALY gained             | 1-way, two-way, PSA  | Yes             | (43) |
| 2005  | Vancomycin                          | Pneumonia                         | Cost per QALY gained             | Scenario             | Yes             | (49) |
| 2005  | Teicoplanin                         | Gram-positive bacteraemia         | Cost per extra cure/success      | 1-way, PSA           | Yes             | (30) |
| 2006  | Vancomycin                          | MRSA pneumonia                    | Cost per extra cure/success      | No                   | Unsure          | (36) |
| 2007  | Teicoplanin                         | Gram-positive infections          | Cost per extra cure/success      | 1-way                | Yes             | (37) |
| 2007  | Vancomycin                          | MRSA surgical site infections     | Cost per extra cure/success      | 1-way, threshold     | Yes             | (40) |
| 2007  | Vancomycin                          | Skin infections                   | Cost per extra cure/success      | No                   | Yes             | (44) |
| 2009  | Vancomycin                          | MRSA skin infections              | Cost per extra cure/success      | 1-way, PSA           | Unsure          | (23) |
| 2009  | Vancomycin                          | MRSA pneumonia                    | Cost per life-year gained        | 1-way, 2-way         | Yes             | (27) |
| 2009  | Vancomycin                          | MRSA skin infections              | Cost per extra cure/success      | Scenario, 1-way, 2-way | Yes       | (28) |
| 2009  | Vancomycin                          | Skin infections                   | Cost per extra cure/success      | 1-way, PSA           | Yes             | (42) |
| 2009  | Vancomycin                          | MRSA pneumonia                    | Cost per extra cure/success      | Scenario             | Yes             | (54) |
| 2007  | Vancomycin                          | Skin infections                   | Cost per extra cure/success      | No                   | Yes             | (26) |
| 2009  | Vancomycin-gentamicin               | MRSA bacteremia/endocarditis      | Cost per extra cure/success      | Scenario             | Unsure          | (22) |
| 2011  | Vancomycin                          | MRSA skin infections              | Cost per extra cure/success      | Scenario, 1-way, PSA | Yes             | (24) |
| 2012  | TMP/SMK                             | MRSA infections                   | Cost savings                     | No                   | No              | (25) |
| 2014  | Vancomycin or β-lactam              | Enterococcal bacteraemia          | Cost per QALY gained             | 1-way, PSA           | Yes             | (52) |
| 2004  | Clarithromycin                      | Pneumonia                         | Cost savings                     | Subgroup analysis    | Yes             | (38) |
| 2004  | Clarithromycin                      | Pneumonia                         | Cost savings                     | No                   | Unsure          | (39) |
| 2004  | Clarithromycin                      | Pneumonia                         | Cost savings                     | No                   | Unsure          | (46) |
| 2008  | Several comparators*                | Pneumonia                         | Cost per extra cure/success      | 1-way                | No              | (41) |
| 2009  | Cefotetan                           | Prophylaxis before surgery        | Cost per failure avoided         | No                   | Yes             | (57) |
| 2009  | Piperacillin/tazobactam             | Intra-abdominal infections        | Cost per QALY gained             | Scenario, 1-way, PSA | Yes             | (32) |
| 2009  | Piperacillin/tazobactam             | Diabetic foot infections          | Cost per QALY gained             | Scenario, PSA        | Yes             | (33) |
| 2014  | Ceftriaxone                         | Pneumonia                         | Cost per extra cure/success      | 1-way, PSA           | Yes             | (50) |
| 2002  | Clarithromycin                      | Exacerbations of chronic bronchitis| Cost per extra cure/success      | Scenario, PSA        | Yes             | (31) |
| 2008  | Ceftriaxone/cefuroxime              | Pneumonia                         | Cost per extra cure/success      | No                   | Unsure          | (21) |
| 2010  | Imipenem                            | Pneumonia                         | Cost savings                     | 1-way                | Yes             | (34) |
| 2010  | Imipenem                            | Pneumonia                         | Cost savings                     | 1-way, PSA           | Yes             | (33) |
| 2010  | Imipenem-cilastatin                 | Pneumonia                         | Cost per QALY gained             | Scenario, 1-way, 2-way, PSA | Yes | (48) |
| 2008  | Vancomycin                          | Skin infections                   | Cost per extra cure/success      | 1-way, PSA           | Yes             | (35) |
of a specific drug might also change over time, when for example the resistance to the old drugs increases or when resistance to the new drug develops. The growing resistance to antibacterial agents highlights the urgent need for the development of new drugs to fight infections.

**Value-based Pricing**

Pricing of antibacterial drugs is an important topic to consider. For pharmaceutical companies, it is important to make profit on the sales of these drugs, because of the large investments needed to develop a new drug. But due to the limited budget, and the rising proportion of health spending that is accounted for by pharmaceuticals, good value for money is an important factor before reimbursement of a new drug. Drug prices are often an important driver of cost-effectiveness. Additionally, infections occur more frequently in developing countries with low insurance coverage where many patients cannot afford expensive antibacterial drugs. Pricing of the drugs is therefore a complicated process, involving different stakeholders.

Box 2 shows a historical overview of pharmaceutical pricing in the UK (59). In this section, we provide a brief summary of this, focusing in particular on value-based pricing. The timelines and examples are UK specific, but could easily be applied elsewhere. Many countries use cost-effectiveness analysis in a similar way to inform reimbursement decisions, although precise guidelines for undertaking such evaluations may vary by country.

In the UK, since 1999, the National Institute for Health and Care Excellence (NICE) appraises the clinical effectiveness and cost-effectiveness of new drugs. Although this body has no direct influence on the price, they do give advice about whether or not drugs (and other healthcare and public health interventions) should be provided by the National Health Service (NHS). NICE accounts for several factors in its decision-making process, but when the incremental costs per QALY gained are higher than £30,000, NICE normally advises not to include this treatment in the NHS setting. As noted, a key driver of cost-effectiveness is drug price. The Prescription Pricing Regulation Scheme (PPRS), established in 1957 as the Voluntary Pricing Regulation Scheme, usually negotiates with pharmaceutical companies to have drugs with a reasonable price which are still profitable for the pharmaceutical industry. When NICE would consider a new drug not cost-effective at that price, pharmaceutical companies may be willing to negotiate a lower price in order to be able to sell the drug in the NHS setting. In 2002, the government established a risk-sharing scheme for new multiple sclerosis drugs. These drugs could be sold at a high price, but if the outcomes of a cohort study were lower than expected, the price was to be reduced. This way, the risk of the drug not being cost-effective was shared by the NHS and the drug company. Since 2009, it is possible for pharmaceutical companies to

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Table 1: continued

| Year | Comparator | Outcome measure | Disease | Disease | Comparator | Cost-effectiveness? | Cost per QALY gained | Cost per QALY gained | Cost per QALY gained | Cost per QALY gained | Cost per QALY avoided | Cost per QALY avoided | Cost per QALY avoided |
|------|------------|----------------|---------|---------|------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|
| 2013 | Fidaxomicin | Cost per QALY gained | Clostridium difficile infection | 1-way, two-way, PSA | Yes | 49 |
| 2013 | Vancomycin | Cost per QALY gained | Clostridium difficile infection | 1-way, two-way, PSA | Yes | 49 |
| 2014 | Vancomycin, metronidazole or FMT | Cost per QALY gained | Recurrent clostridium difficile infection | 1-way, two-way, PSA | Yes | 51 |
| 2014 | Vancomycin | Cost per QALY gained | Clostridium difficile infection | 1-way, two-way, PSA | Yes | 51 |

OPAT, Outpatient parenteral antimicrobial therapy; TMP/SMX, Trimethoprim/sulfamethoxazole; PSA, probabilistic sensitivity analysis; FMT, faecal microbiota transplant; QALY, quality-adjusted life-years.

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negotiate price reductions when the drug is considered not cost-effective by NICE and therefore not available on the NHS. A discount is provided, so that the costs of the new treatment are below the threshold set by NICE and the treatment can be available on the NHS (patient access schemes) (59). Also, NICE increased the willingness-to-pay threshold for drugs extending the life of patients with a short life expectancy (end of life criteria).

Instead of studying the cost-effectiveness of a drug with a certain price, NICE could also calculate the maximum price of the drug at which it would still be cost-effective. When the price of the drug would be no higher than this threshold value, it would represent 'value for money'. Given the fixed budget of the NHS, money spent on a new drug cannot be spent elsewhere. With a willingness-to-pay threshold of £20 000 per QALY gained, we assume that every £20 000 used for a new treatment displaces one QALY elsewhere in the NHS (60). The QALYs expected to be displaced by a new treatment can be calculated by dividing the total expected costs by the willingness-to-pay value. This forms the basis of value-based pricing. Figure 2 depicts the costs of a hypothetical new drug that increases health by 2 QALYs compared with the comparator at three different prices. At price 1, the total extra cost of this treatment to the NHS is £20 000, which leads to an ICER of £10 000 per QALY gained. So, at price 1, the drug is expected to improve health by two QALYs and displace 1 QALY (total costs of £20 000 divided by the willingness to pay of £20 000) elsewhere in the NHS. The net health benefit, defined as the difference between the total expected QALYs and the QALYs expected to be displaced elsewhere, would in this case be one QALY (2 expected QALYs minus 1 QALY displaced elsewhere). At price 2, the total extra costs of this treatment are £40 000, which leads to an ICER of £20 000 per QALY gained. Now the drug is still expected to improve health by 2 QALYs, but displaces 2 QALYs (£40 000) elsewhere. There is no net health benefit to the NHS. At price 3, the total extra costs of the treatment are £60 000, which leads to an ICER of £30 000 per QALY. In this case, 3 QALYs are displaced (£60 000), and the net health benefits to the NHS are minus 1 QALY (2 expected QALYs minus 3 QALYs forgone elsewhere). The aim of value-based pricing is to set the price at a level at which the net health benefit is positive or zero. A negative health benefit means that the drug does not provide value for money. Given that the definition of net health benefit is the difference between the total expected QALYs and the

![Figure 2: Net health benefit of a hypothetical new drug at three different prices (60).](image)
QALYs expected to be displaced elsewhere (total expected costs divided by the willingness-to-pay value), one could also say that the aim of value-based pricing is to set the price of a new drug to a value at which the ICER of that new drug would be at or below the willingness-to-pay threshold. In other words, the price can be varied up to the point at which the incremental cost per QALY is equal to the threshold.

Value-based pricing was first recommended by the Office of Fair Trading. A key issue with value-based pricing is how ‘value’ is measured. One option would be to express value as therapeutic value, measured by QALYs. When a new drug is very effective (it produces a large increase in QALYs), the drug can be more expensive than when it is moderately effective (small increase in QALYs). However, it has been suggested that not only QALYs should be taken into account when assessing value, and many other factors could also be included (see Box 3) (61,62). Based on recent consultation documents produced by NICE, two additional components of value that are being given careful consideration are the burden of disease and the wider social impact – the impact of disease on people’s ability to be part of society. Whichever values are included in value-based pricing, an important consideration is how these ought to be weighted to account for the fact that different drugs may have different effects on each factor.

There is little evidence from other countries about the prospects of value-based pricing, because no country currently performs value-based pricing as proposed in the UK. However, in some countries, the insurance coverage of a new drug is based on the cost-effectiveness of the drug. In Sweden, insurance coverage depends on approval from their health technology assessment body (TLV). For this decision, QALYs as well as production loss are taken into account.

**Future Prospects**

It is unclear yet what the influence of value-based pricing will be on the prices of new antibacterial agents. If we look at the different components NICE is planning to consider in their evaluation, three factors will be of importance: health gain (QALYs), burden of disease and wider social impact.

### Health gain

Many of the studies reviewed in this article showed that the new antibacterial agents were more effective than the comparator, mostly because of a higher cure rate. Because these drugs can be life-saving, a large ‘value’ can be expected for which a high price can be paid. However, there is currently little evidence about the incremental costs per QALY gained. More evidence is needed using the effect of new antibacterial agents on QALYs to set a value-based price for these drugs.

### Burden of disease

Because bacterial infections occur frequently, the total burden of these infections is large. However, as we have seen in this review, each indication (type of infection) needs to be considered separately. For some indications, the burden of disease might be lower than others. In general, the burden of one specific bacterial infection is probably low, also because of the short duration of the illness. Chronic conditions have a more significant effect on burden of disease than (treatable) acute conditions.

### Wider social impact

National Institute for Health and Care Excellence defines the wider social impact as the loss in capacity of a person with the disease to engage with society (e.g. working or taking care of someone), compared with their capacity without the disease. If infections occur among working people, the impact on productivity might be high. However, when the duration of the infection is usually short, the impact will not be as high as for example with chronic diseases.

**Conclusions**

Since 2000, several new antibacterial agents have been developed to treat (resistant) infections such as pneumonia.

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**Box 3: Possible factors to take into account when assessing value (61,62)**

| Patient or disease-related factors | Healthcare process-related factors | Factors outside patient + NHS |
|-----------------------------------|-----------------------------------|-------------------------------|
| Severity of disease               | Treatment time + location         | Ability to resume working     |
| Near the end of life              | Waiting times                     | Increased productivity        |
| Size of population                | Less unpleasant treatment         | Benefit to carers             |
| No other treatment options        | Degree of risk of the treatment   | Cost savings to other services|
| Socially disadvantaged patients   |                                   | Cost savings to patients/carers|
| Children                          |                                   | Quality of evidence           |
| Reduction in fear (e.g. of death) |                                   | Innovation                    |
| Unmet need                        |                                   |                               |

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and skin infections. To keep healthcare spending within reasonable limits, it is important that new drugs are only prescribed when these drugs provide value for money. Most new antibacterial agents are cost-effective alternatives to the old drugs, either because these are more effective and decrease healthcare costs or the increased costs are below the willingness-to-pay threshold. Value-based pricing is a method that could be used to determine a price for new antibacterial agents at which these drugs provide value for money.

Conflict of interest

None declared.

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Notes

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