Managing UTI in primary care: should we be sending midstream urine samples?

The decision to prescribe antibiotics is one of the most common treatment decisions faced by frontline primary care clinicians daily, and urinary tract infection (UTI) is one of the most common bacterial infections encountered.1 Although antimicrobial treatment for UTI is accepted in clinical practice, antibiotic resistance in urinary bacteria is increasing,2 with rates cited in the literature of between 20% and 40% to trimethoprim and amoxicillin respectively.2 Bacteria are adept at side-stepping human intervention (for example, antibiotics and vaccines) and are developing resistance to antibiotics faster than the pharmaceutical industry is developing new ones: only two new classes of antibiotics have been developed in the last 30 years.3

These problems highlight the need for high quality evidence to help primary care clinicians optimise the diagnosis and management of UTI. Fortunately, a plethora of high quality, clinically valuable, primary care research papers have been published recently, including four in the Daily Mail4–6 and two in this month’s BJGP.7,8

One of the BMJ papers reported that a range of management strategies (immediate versus delayed empirical antibiotics versus antibiotics guided by the results of dipstick versus symptom score versus midstream urine culture) achieved similar symptomatic results, while delayed empirical and dipstick-guided prescribing reduced antibiotic consumption.2 The authors concluded that there is ‘no advantage in routinely sending midstream urine [MSU] samples for [culture]’. This makes for an interesting and contrasting background to one of the papers reported in this month’s BJGP,7 which provides evidence in support of sending MSUs for culture.

This study by Vellinga et al provides some original and novel evidence for a practice I suspect, is commonly used in primary care: that of looking for a previous urine culture to guide the treatment of a current, suspected UTI. Although it has made intuitive sense to avoid an antibiotic to which a patient’s urinary bacteria have previously been found to be resistant, until now there has been no evidence to support this practice. Furthermore, when it was used, clinicians must have wondered how long previous results remain relevant and if resistance to some antibiotics is more important than others.

The study by Vellinga and colleagues addresses both these questions. They found that a previous measure of resistance to co-amoxiclav, amoxicillin, ciprofloxacin, and trimethoprim remains predictive of further resistance at 3 and 12 months. The long duration of effects, although not as strong as at 3 months, may surprise many clinicians, but is not an isolated finding. A recently published systematic review9 reports a similar trend of effects in three studies10–12 that, when pooled, show that the relationship between antibiotics prescribed over a 12-month period and bacterial resistance cannot be explained by chance (odds ratio = 1.33, 95% CI = 1.15 to 1.53).

Happily, their findings do not contradict the Health Protection Agency’s antibiotic prescribing guidance14 that nitrofurantoin should be used as a first line treatment. In the study dataset, this was the antibiotic to which fewest bacteria were resistant at baseline, 3 and 12 months; and the antibiotic with the lowest positive and highest negative predictive values. In other words, previous resistance was least likely to predict future resistance, and previous susceptibility was most likely to predict future susceptibility.

I think there are two potential study limitations of Vellinga et al’s study which we should be aware of. First, the need to distinguish between a recurrent infection versus detecting the same infection twice. The authors’ chose a threshold of 2 weeks; that is, if a patient was found to have a repeat infection more than 2 weeks after the index infection, it was deemed a recurrence. This is a classic point of uncertainty for any study design, as there will be little, if any, evidence available to assist decision making and whatever is selected could be criticised as being too generous or lenient. However, given that one trial showed that 88% of women with UTI treated with nitrofurantoin had both improved and were microbiologically cured at 7 days,15 2 weeks does not seem unreasonable. Second, it would help us interpret the generalisability (otherwise known as the external validity or the extent to which the study patients are similar to our patients) if we know how Galway GPs in the Vellinga et al study use urine sampling. For example, is the local policy to send samples on all patients with suspected UTI, or only those with atypical symptoms, or patients deemed to be at high risk?

The second study16 in this issue of the BJGP makes an important contribution to improving our understanding of the relative contribution that symptoms, signs, and near-patient tests make to the diagnosis and prognosis of infections, or put more simply, developing diagnostic/prognostic algorithms. There are three stages in the development of such algorithms. First derivation, in which candidate symptoms, signs, or near-patient test results (the ‘predictors’) are identified as being statistically associated with the diagnosis (or prognosis) in question. Second validation, where predictors are checked in a different dataset to see if they remain associated, and to a similar degree. Finally, the use of the validated algorithm should be compared to usual care in a randomised controlled trial to see if patient outcomes are improved.

The Little et al study17 is an example of a second (validation) stage study, of which there are surprisingly few in primary care. It addresses the issue of accurately diagnosing UTI in women, which is vital if we are to target antibiotics at those who are most likely to benefit (and withhold them from those who are unlikely to). And the diagnosis is by no means clinically straightforward with only 50% to 66% of women with suspected UTI having a
microbiologically confirmed infection. The Little et al study is well designed and executed, and largely confirms the findings of its sister derivation study, showing that the study cannot assess the emergency clinic. However, it also means that the study cannot assess the diagnostic value (positive or negative prediction) of these signs.

In summary, the evidence reviewed suggests a role for urine dipstick in ruling in UTI and reducing antibiotic use. What is less clear is whether we should routinely send MSUs. If we stop, we could save money, but as Srirangam and Bondin describe, we could be losing in three ways. First, urine cultures provide valuable information regarding the changing patterns of antimicrobial resistance (a societal benefit). Second, as shown by Vellinga et al, a proportion of patients experience repeated UTIs, and previous results can help guide subsequent treatment (a patient benefit). Third, as discussed, between 34% and 50% of women presenting with urinary symptoms do not have a UTI, and we currently do not have any other sufficiently accurate diagnostic test to detect these patients. To these three I would add a fourth, namely clinical curiosity. Positive and negative predictive values only estimate group average probabilities of UTI, but the only way of establishing the diagnosis for the individual patient, with their unique history and set of circumstances, is a urine culture.

It is surely the case that ‘not all UTIs are the same’ and that there may be some patients in whom MSU is not cost-effective (from either a patient or societal perspective), but that in others (perhaps older patients, those with more severe symptoms, or those with recurrent UTI) there is a role for MSU. Further research is needed.

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REFERENCES

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