Methods for determining methicillin sensitivity of staphylococci showed that since the first resistant strains were reported, they were originally developed for S aureus; the same modifications are now being applied for CNS. However, CNS grow less luxuriantly than S aureus and the resistant subpopulation is smaller, making it harder to detect methicillin resistance. Where laboratories incorporate salt in their media for methicillin testing, a concentration of 5% is usually used, but this concentration is not well tolerated by staphylococci. Previous investigations showed that 2% salt provides an osmotically supportive medium and, in conjunction with Columbia agar, encourages enhanced growth and therefore more clearly demonstrates methicillin resistance. These results are also in general agreement with those of other studies and we recommend the use of Columbia agar incorporating 2% salt for the detection of methicillin resistance in CNS.

JRD SCOTT LM J MORGAN JVS PETHERT J W JONES Public Health Laboratory, Maegusor Park Hospital, Tauton, Somerset TA1 5DB JF RICHARDSON Staphylococcus Reference Laboratory, Central Public Health Laboratory, 61 Colindale Avenue, London NW9 5HT RHF BOLEY Ei Lilly and Co. Ltd, Kincleridge Road, Basingstoke, Hampshire RG21 2XJ


Correspondence

Effects of culture media on detection of methicillin resistance in coagulase-negative staphylococci

We read with interest the report by Milne et al on the effects of culture media on the detection of methicillin resistance in coagulase-negative staphylococci (CNS). We agree with their conclusions that the incorporation of salt in Columbia agar is the most reliable culture medium for the detection of methicillin resistance in CNS by disc testing.

The UK National External Assessment Scheme for Microbiology (NEQAS) has shown that generally most laboratories can recognise methicillin resistance in Staphylococcus aureus but have more problems with Staphylococcus epidermidis (Snell, personal communication). Some laboratories may seriously underestimating the percentage of isolates of CNS that are resistant to methicillin.

We examined 248 CNS for susceptibility to methicillin by disc testing using three different methodologies. Method 1 used Columbia agar with 2% salt added (CA + 2% salt) whilst method 2 used Mueller Hinton agar containing 2% salt (MH + 2% salt); in both methods incubation was at 35°C for 24 hours. Method 3 incorporated Diagnostic Sensitivity Test (DST) agar plus 1% lysed blood, incubated at 30°C for 21 hours. Strains showing equivocal results were further examined by plate MIC using the three media and respective incubation conditions.

Of the 248 strains tested, 36 were excluded because they were novobiocin resistant and two more were excluded because they were not identified on the ATB 32 Staph system (bioMerieux SA). Of the remaining 210 stains, the MIC results showed 107 (51%) to be resistant to methicillin. The results for the three methodologies are shown in the table.

The method using Columbia agar incorporating 2% salt incubated at 35°C detected all the resistant strains, but two sensitive strains were identified as resistant. This result is clearly superior to both of the other methods tested, although variations of methods based on DST incubated at 30°C appear to be the most common in use in the UK for determining methicillin sensitivity (Snell, personal communication).

Diagnosis of Helicobacter pylori in biopsy specimens

I read with interest the report by Veennendaal et al who showed that a 24 hour delay and storage of antral biopsy specimens in physiological saline solution did not alter the positive culture rate of Helicobacter pylori. However, I feel that the definition they have used for the diagnosis of H pylori infection may be misleading, if used without consideration. As they state, it is clear that culture is a 100% specific, though "probably not the most sensitive test" for the identification of H pylori infection. Their case with a positive serology and histology result, but with a negative culture seems to justify this view. Concerning their remaining two possible cases, my opinion is that microscopic identification of an H pylori-like organism in a haematoxylin and eosin stained biopsy specimen alone cannot be accepted as a diagnostic criterion for infection. Haematoxylin and eosin staining is not a generally accepted tool for demonstrating the micro-organisms and is insufficiently sensitive, though it is ideal for indicating antral gastritis. It is not suitable for detecting early colonisation of the mucosa by the bacterium. Stains such as Wright-Giemsa, Brown-Hopps or Warthin-Starry silver stain are more commonly accepted for this purpose because they are more sensitive.

In routine diagnostic work, however, it is advisable to use both of the two tests that are available (immunomembranous assay (ELISA), for IgG H pylori antibodies, used in the same population (143 positive patients). Other authors also confirmed the diagnostic value of histology and haematoxylin and eosin staining methods. However, experience with the test used is essential and here we agree with the correspondents.

In our pathology department there is a long standing interest and experience in the detection of H pylori and its associated gastritis. When in doubt about the diagnosis, especially after treatment when organisms are infrequent or absent in the presence of chronic gastritis, additional staining techniques (Giemsa or Warthin-Starry) are occasionally necessary. This did not apply to our study. We therefore consider our "good standard" appropriate.

As pointed out in our article, we feel that culture is important for routine diagnostic

Dr Veennendaal and Lichendahl-Bernards correspondence

Although the results of our study as indicated by the correspondents were not influenced by the definition of Helicobacter pylori infection used, their comment addresses an important point.

There is no general agreement as to which tests should be used as the gold standard in the diagnosis of gastric H pylori infection. In a recent (unpublished) study we found the combination of histology (haematoxylin and eosin staining as well as a definition of H pylori infection superior with regard to sensitivity when compared with either test alone (121 patients culture positive, 125 patients positive by histology and 137 patients positive by histology, culture or both). These findings almost matched the results of a previous validated sensitive and specific enzyme linked immunomembranous assay (ELISA), for IgG H pylori antibodies, used in the same population (143 positive patients). Other authors also confirmed the diagnostic value of histology and haematoxylin and eosin staining methods. However, experience with the test used is essential and here we agree with the correspondents.

In our pathology department there is a long standing interest and experience in the detection of H pylori and its associated gastritis. When in doubt about the diagnosis, especially after treatment when organisms are infrequent or absent in the presence of chronic gastritis, additional staining techniques (Giemsa or Warthin-Starry) are occasionally necessary. This did not apply to our study. We therefore consider our "good standard" appropriate.

As pointed out in our article, we feel that culture is important for routine diagnostic
work, because it allows antimicrobial sus-
cceptibilities (metronidazole, clarithromycin) to be determined, and we would not replace it for a specific test that was not fully spec-
cific as suggested by the correspondents.


2 Peña AS, Endris HPH, Offerhaus CJA, Hoogendoorn-Verdegal A, et al. Value of serology (ELISA and Immunoblotting) for the diagnosis of campylobacter pylori infect-


Measurement of medical staff overload

Dr Bignardi is correct in his conclusion that it is difficult to measure medical workloads and requirements in microbiology departments. The current guidelines of the Royal College of Pathologists for consultant staffing suggest that for central laboratories in district general hospitals serving a popula-
tion of approximately 250 000 there should be at least two consultant medical microbiologists. A number of districts do not provide such staffing and cases need to be developed to persuade managers to provide appropriate levels. "Population served" is a crude measure of workload, even if referral patterns do not distort the picture. It is also clear that hospital bed numbers are not directly related to laboratory activity; indeed, historically bed numbers reducing bed numbers has resulted in an increase in laboratory tests from outpatients, day cases, and GPs. Numbers of specimens and the number and nature of tests can be more closely related to laboratory activity and can be made more sophisticated by such systems as WELCAN, but these are not a measure of medical input; neither are they a measure of the quality of a microbiology service. Particular problems in measuring consul-
tant microbiologist input are the contribu-
tions to core activities of the hospital(s) and clinical services—activities such as hospital infection control, policies for infection control, chemical disinfection—and the general pro-
vision of advice on the management of infected patients. The latter aspects depend to a large extent on the case mix profile of the units served: intensive care units, special care baby units and oncologywards make particularly heavy demands on medical microbiologists. Although these matters are generally clear in principle, the allocation of numbers to reflect the work load has proved to be very difficult. Some of the problems of consultant staffing levels have been discussed in a recent article in ACP News and the Microbiology Specialty Advisory Committee of the Royal College of Pathologists is currently examining this subject. It will not be easy to produce a uni-
versally acceptable measure, but the prob-
lems must be addressed in order to try to achieve a composite workload definition that reflects the range of input required of a consultant microbiologist.

DR DAVIES BI DURDEN
Microbiology Specialist Advisory Committee,
Royal College of Pathologists,
2 Carlton House Terrace,
London SW1Y 5AF


3 Workload figures: whose norms are they anyway? ACP News 1993:11–12.

Dr Bignardi comments:

I welcome the interest by the Microbiology Specialist Advisory Committee of the Royal College of Pathologists: eliciting such inter-
est was the main purpose of my report. In my opinion the current guidelines by the Royal College of Pathologists for consultant staffing in microbiology are so impractical that they cannot be implemented by the College itself. This is demonstrated by the fact that, during the period of my study, four job descriptions for single-handled con-
sultants were approved by the College despite the fact that the respective popula-
tions exceeded 250 000 (the College recom-
ends two consultants for departments serving a population of approximately 250 000). According to my analysis, the case for a second full-time consultant microbiologist was very strong in two of the three four hospitals.

One would hope that if a formula based on the weighted number of beds and speci-
mens (and perhaps on other factors) was sanctioned and policed by the College, at the least the worst cases of understaffing could be eliminated. Since writing my report I have noticed some important trends: the overall number of both consul-
tants and junior doctors in microbiology seems to be decreasing, pathologists and pathol-
yogy departments have been asked to take sub-
stantial cuts in their budget over the next years, and the NHS Management Executive has commissioned a strategic review of pathology services which may subsequently throw the door open to more pathology privatisations.

Given the current political climate, I think it most important that we try to iden-
tify and quantify the minimum medical staff requirement for a good quality service in microbiology.

Necrotising granulomas of the uterine corpus

We read with interest the report by Drs Akosa and Boret of necrotising granulomas of the uterine corpus following Nd YAG laser ablation of the endometrium, and noted their reference to the original report of peritoneal granulomas following laser ablation.1

We subsequently reported the histo-
logic findings from four hysterectomy specimens obtained for various indications following Nd YAG laser ablation.3 Our findings were essentially the same as those of Akosa and Boret, and we were able to demonstrate by energy dispersive x-ray analysis that the black foreign material within the necrotising granulomas consisted largely of aluminium oxide compatible with the known composition of the sapphire laser probe.

We also provided evidence to support the hypothesis that recurrent bleeding following laser ablation is due to inspissation of functional endometrium from the tubal ostia and isthmus,4 and we described how Akosa and Boret made no comment on the histological appearances of the endo-
metrium away from the obvious laser damage.

Finally, Akosa and Boret refer to the technique as endometrial resection which is in our view not correct, as the use of the Nd YAG laser is a technique for endometrial ablation.

JHF SMITH A KENNEDY
F SHARP
Northern General Hospital,
NHS Trust, Horsey Road,
Sheffield S7 7AU

PC REID
Lumon and Doncastro Hospital

W THURRELL
University College Hospital,
London

Dr Akosa and Boret comment:

We are grateful to Dr Smith et al for their prompt comment on our short report. This was basically intended to increase awareness among histopathologists of what has become a diagnostic quandary in the absence of adequate clinical information and in view of the increasing use of minimal invasive surgical techniques.

We noted in our report that the abnor-
malities in the endometrium were either complete or focal, the latter the cause of subsequent bleeding. The residual endo-
metrium, although not stated in our report, was not confined only to the cornu as in the case referred to in the paper by Baggish and Baltoyannis. If one assumes that in every case of endometrial ablation the entire endometrium is destroyed, the hypothesis of inspissation may be acceptable; in our experi-
ence this is not always the case.

Endometrial resection using laser and endometrial ablation have been and are used interchangeably. Our opening sen-
tence is now under discussion, as "Transcervical resection of the endo-
metrium is a hysteroscopic method of endo-
metrial ablation": this is self-explanatory.

Our literature search was confined to 1990 onwards, which explains why the papers by Baggish and Baltoyannis and Lomano were not cited. As for the paper by Reid et al, we can only assume that at the time of our search it had not been indexed.

We have now read all these papers and they


2 Baggish GE, Reid PC, Kennedy A, Smith JHF. Necrotising granulomas of the peri-

3 Reid PC, Thurrell W, Smith JHF, Kennedy A, Sharp P. Nd YAG laser endometrial ablation: histological aspects of uterine heal-

4 Lomano JM. Photocoagulation of the endo-

5 Baggs MS. Baltoyannis, Short report. This pa-
Diagnosis of Helicobacter pylori in biopsy specimens.

G Cserni

doi: 10.1136/jcp.47.4.380-b

Updated information and services can be found at:
http://jcp.bmj.com/content/47/4/380.2.citation

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/