Detection of fastidious bacteria in cardiac valves in cases of blood culture negative endocarditis

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Abstract
The diagnosis of blood culture negative endocarditis is still a problem. Fastidious bacteria such as bartonella and coxiella are responsible for cases of blood culture negative endocarditis, the identification of which is mainly based on serological and DNA studies only available in specialised centres. Therefore, a routine technique is needed in surgical pathology laboratories to detect these bacteria in cardiac valve tissue sections. This report describes a staining technique, the Giménez stain, feasible and sensitive in detecting bartonella and coxiella in two cases of blood culture negative endocarditis. (J Clin Pathol 2001;54:238–240)

Keywords: endocarditis; bartonella; coxiella

The diagnosis of endocarditis remains a problem because about 10% of cases are diagnosed with negative blood cultures.1 There are at least two different reasons for the negativity of blood cultures. First, it is most frequently caused by previous antibiotic treatment. Second, it can be the result of bacterial species that are difficult to grow in the laboratory. Among these fastidious bacteria, some such as bartonella are responsible for re-emerging diseases in homeless people1 and can cause endocarditis.3 4 Coxiella burnetti is another bacterium implicated in blood culture negative endocarditis.5 6 The identification of these agents requires specific methods mainly based on serological studies, blood and valve bacterial cultures under special conditions, or the amplification and sequencing of bacterial DNA extracted from valve tissue.7 8 In some cases, immunofluorescence or immunohistochemistry on valve tissue sections has been found to be diagnostic.9 8 All these techniques are only available in specialised laboratories. Gram staining of valve tissue sections yields poor results. Given this background, we tested the feasibility and efficacy of Giménez staining done on valve tissue sections in a routine surgical pathology laboratory to detect bartonella and coxiella in cases of endocarditis with negative blood cultures.

Case reports

CASE 1
A 39 year old alcoholic homeless man was admitted into hospital because of fever of 38.5°C and polyarthralgia. He had an aortic diastolic murmur. Haematological laboratory values were: haemoglobin, 75 g/litre; white blood cell count, 13 000/mm³; and platelet count, 111 000/mm³. Because the fever persisted, transthoracic and transoesophageal echocardiographies were performed. They revealed aortic vegetations associated with aortic insufficiency. Four routine blood cultures were negative. The patient was treated with 8 g amoxicillin/day and 120 mg gentamicin/day. Despite the antibiotics, he remained febrile and the aortic insufficiency worsened. Surgical aortic valve replacement using a cryopreserved aortic valve homograft was performed two weeks after the beginning of treatment. The native aortic valve showed vegetations on the anterior right cusp, tearing and prolapse of the posterior cusp, and extensive loss of tissue in the anterior left cusp. The aortic valve bacterial culture was negative. An imprint of the valve tissue was performed. The serological studies showed IgG titres of 1/1600 against Bartonella henselae, 1/1600 against B henselae serotype Marseilles, and 1/3200 against B quintana. The serological studies against Afipia felis, brucella, coxiella, and chlamydia were negative. DNA extracted from the valve tissue was submitted for amplification of the intergenic spacer (IS) region as described previously.9 6 A specific signal was detected for B quintana.

CASE 2
In February 1999, a 44 year old Italian man was admitted for fever and surgical treatment of an aortic endocarditis of unknown aetiology. His remote medical history included rheumatic heart disease, and a splenectomy for congenital microspherocytic haemolytic anaemia. In 1996, he presented with fever and myalgia. He was treated with several short courses of doxycycline. In September 1998, dyspnea and worsened aortic insufficiency appeared despite a course of intravenous ceftriaxone followed by vancomycin and gentamicin. At this time, all routine blood cultures were negative. In January 1999, an aortic vegetation and a septal abscess were detected by echocardiography. He was then transferred for replacement of the aortic valve and treatment of the septal abscess. A cryopreserved aortic valve homograft was used. The aortic cusps were thickened and mildly calcified, and a small vegetation was adherent to the anterior cusp. Therefore, an imprint of the anterior cusp was performed. Serological tests for Legionella pneumophila, Chlamydia psittaci, B quintana, and B henselae were negative. The serological test for C burnetti was positive, with an IgG titre of 1/12800 and an IgA titre of 1/400 against phase I of C burnetti.10 The bacterial cultures of valve tissue performed on endothelial cell cultures and the polymerase chain reaction (PCR) with amplification of the IS 1111 gene on DNA extracted from valve tissue11 were both positive. The patient was treated with 200 mg...
doxycycline/day and 200 mg hydroxychloroquine sulphate/day. Two months later the patient was well.

Methods
In case 1, a small piece of formalin fixed aortic cusp measuring $5 \times 5 \times 3$ mm, moderately thickened and fibrous, without any gross vegetation, was available for pathology. In case 2, a 13 mm length fragment of valve tissue thickened by calcified fibrosis without gross vegetation, which was decalcified in RDO (Eurobio, Les Ulis, France) for six hours, was available. For this last case, a piece of aortic fibrous ring was also available. Paraform wax sections were cut at 2 µm and stained with haematoxylin and eosin, Gram, Giemsa, and Giménez stains. The valve imprints were also stained with Gram and Giménez stains. Giménez staining was performed as follows: (1) the dewaxed sections were incubated at room temperature for two minutes with a freshly filtered solution made from 2 ml of the stock solution of basic fuschin in 5 ml of buffer; (2) after rinsing well in tap water, they were incubated in malachite green for nine seconds; (3) after rinsing well in tap water, they were again incubated with the malachite green solution for nine seconds; (4) after rinsing well in tap water, the sections were air dried and mounted in Eukit (Labonord, Villeneuve d’Ascq, France).

STOCK SOLUTIONS
Stock solutions of basic fuschin were made up as follows. Solution 1: 10 g of base fuschin (RAL, Rieux, France) in 100 ml of 95% ethanol; solution 2: 11.25 g of phenol (Sigma, St Louis, Missouri, USA) in 250 ml of distilled water at $37^\circ C$; mix solutions 1 and 2 in 650 ml of distilled water. The 0.8% (wt/vol) solution of malachite green was made up as follows: 2 g of malachite green oxalate (Merck, Darmstadt, Germany) in 250 ml of distilled water. Buffer: 3.5 ml of 0.2 M monosodium phosphate (Sigma) plus 15.5 ml of 0.2 M disodium phosphate (Sigma) in 19 ml of distilled water.

CONTROLS
Thirteen cases of streptococcus or staphylococcus endocarditis with valve tissue sections showing full blown inflammation and many Gram positive cocci were selected from pathology files. They were cases of rapidly progressive valve incompetence necessitating surgical valve replacement after a short antibiotic treatment. An additional case of Staphylococcus epidermidis aortic endocarditis treated with antibiotics for four weeks was also available: it showed mild chronic inflammation and Gram stain negativity. In all these cases, valve tissue sections were stained with the Giménez stain as described above.

Results
In case 1, the valve tissue showed replacement fibrosis associated with a focus of inflammatory cells and necrosis. Inflammation consisted of mononuclear cells mixed with rare polymorphonuclear cells. In the superficial aspect of the valve, there was a thin fibrin vegetation without inflammation. Gram and Giemsa stains were negative. Giménez stain showed many intracellular short bacilli in the inflammatory focus in the valve. The bacilli strongly stained pink/red in a blue/green background (fig 1A). The valve imprint showed the same bacilli.

In case 2, the valve tissue showed fibrosis and large inflammatory foci with necrosis, mononuclear cells, and many polymorphonuclear cells. A superficial thin, leucocyte rich fibrin vegetation was observed. Gram and Giemsa stains were negative. Giménez stain showed many intracellular thin bacilli in the inflammatory focus in the valve (fig 1B). The valve imprint showed the same bacilli with Giménez stain, whereas the Giemsa and the Gram stains were negative.
Of the 13 control cases of Gram stain positive streptococcus or staphylococcus acute endocarditis, three cases were also positive with the Giménez stain showing pink/red cocci; staining was much less obvious than with the Gram stain, and 10 cases were negative. In the case of Gram stain negative S. epidermidis aortic endocarditis after prolonged antibiotic treatment, Giménez stain showed a few positive residual intracellular bacteria in the inflammatory foci (fig 1C).

Discussion

In both cases, endocarditis was confirmed by histological examination of the valve showing inflammation. However, standard diagnostic tests such as blood and valve cultures in routine bacteriological media, Giemsa, and Gram stains of valve tissue sections failed to demonstrate the aetiology. Finally, endocarditis was related to B. quintana or C. burnetii by serological studies and confirmed by amplification of DNA extracted from the valves in a specialised laboratory.

Given the increasing incidence of infections with fastidious Gram negative bacteria, mainly B. quintana, but also C. henselae,16,17 causing endocarditis and the fact that endocarditis can result from infection with C. psittaci18 and C. burnetii,5-7 it has become a major challenge for the pathologist to detect bacteria in valve tissue sections to help in diagnoses where usual microbiological techniques fail and where more sophisticated techniques, only available in specialised laboratories, are required.

In specialised centres where specific antibodies are available, immunofluorescence or immunohistochemistry can be performed in valve tissue sections to detect bartonella and C. burnetii.3 Gram and Giemsa stains are not always sensitive for bartonella15 and C. burnetii.17 The Warthin-Starry stain is difficult to perform and to analyse because of frequent heavy background staining, although it has been used to detect bartonella in tissue sections.15,18 Therefore, a feasible and sensitive staining technique for the detection of fastidious bacteria is needed in pathology laboratories dealing with the surgical pathology of cardiac valves. The Giménez stain has been performed in microbiology laboratories to detect such bacteria on pus and effusion drainage liquid19 smears, and on lymph node and valve imprints. We show here that it can be used successfully on formalin fixed, paraffin wax embedded tissue sections, even after mild decalcification, as long as the sections are thin (2–3 μm). The Giménez stain yields clear cut staining of bartonella and coxiella, which appear pink/red, whereas the underlying tissue is blue/green. Furthermore, it sometimes stains usual bacteria, both Gram positive and Gram negative, after prolonged antibiotic treatment. In the latter situation, Giménez stain positivity does not assume that the bacteria are still alive. Thus, we suggest the systematic use of the Giménez stain for the pathological examination of valves in cases of endocarditis.20

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