Propionibacterium acnes as a Cause of Aggressive Aortic Valve Endocarditis and Importance of Tissue Grinding: Case Report and Review

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A case of prosthetic valve endocarditis with Propionibacterium acnes is described. The diagnosis was documented by histology and isolation of P. acnes from both blood and anulus tissue. Grinding of the tissue, which was first omitted to avoid contamination, was indispensable for cultivating the agent. The literature for P. acnes endocarditis is reviewed.

Propionibacterium acnes is a microorganism that usually has a low level of virulence. Nevertheless, it is associated with serious infections like brain abscesses (14), osteomyelitis (2), endophthalmitis after intraocular surgery and lens implantation (21, 23), subdural empyema (4, 11, 25), cerebral shunt infection (6), parotid and dental infections (8), and infective endocarditis (IE) (1, 5, 7, 10, 12, 13, 15–17). The strongest risk factor for serious infection is the presence of foreign bodies (1). As a member of the normal skin flora, P. acnes is mostly a contaminant in cultures of wound and blood specimens. In clinical settings it is often difficult to distinguish between contamination and infection. We describe here a case of a prosthetic valve endocarditis with P. acnes. Furthermore, the methods of processing tissue samples to increase sensitivity for growth of bacteria are discussed and the literature of IE caused by P. acnes is reviewed.

A 72-year-old man underwent aortic valve implantation with a St. Jude Medical prosthesis to replace a biologic aortic valve implanted in 1985 for degenerative aortic insufficiency. Three weeks postoperatively he developed fever, tiredness, headache, and right-side pleural effusion. Seven weeks after discharge, he again presented with chills and left-side hemiparesis. Transthoracic echocardiography showed free-floating masses on the St. Jude Medical valve. A first computed tomography scan of the skull was normal. A second computed tomography scan performed 2 weeks later showed subacute right-side supratentorial parietal and left-side infratentorial infarctions. On admission, his temperature was 36.8°C, his pulse was 76 beats per min, and his blood pressure was 125/75 mm Hg. Skin and lung examinations revealed no abnormalities. Cardiovascular examination showed a lateralized, elevated maximum impulse, crisp prosthetic valve opening and closing sounds, and a grade 4/6 systolic ejection murmur over the entire precordium. Neurologic examination showed signs of sensorimotorial left-side hemiparesis and urinary bladder retention. Vegetations and a paravalvular abscess in the region of the noncoronary sinus Valsalvae with a paravalvular aortic insufficiency were seen upon transesophageal echocardiography. Abnormal laboratory findings included a sedimentation rate of 35 mm/h, a C-reactive protein concentration of 60 mg/liter, and a hemoglobin level of 11.1 g/dl. The leukocyte count was 9.63 × 109/μl with no shift to the left but a few toxic granulocytes.

Three sets of blood samples for culture (one set consisting of an aerobic and an anaerobic bottle) were taken 1 day before the operation, and five sets were taken on the day of the operation. The St. Jude Medical aortic valve was replaced by an aortic homograft. The aortic anulus showed signs of prosthetic endocarditis, and an annular abscess, previously diagnosed by transesophageal echocardiography, was opened and debrided and tissue was sent for histologic and microbiologic investigations. Postoperatively, intravenous antibiotic therapy with amoxicillin-clavulanate (2.2 g every 8 h) and netilmicin (100 mg every 12 h) was started. At 7 and 8 days, respectively, after taking the blood cultures, three bottles (two aerobic, one anaerobic) of the first three sets and three bottles (two anaerobic, one aerobic) of the five sets of blood for culture taken 1 day later showed gram-positive rods that were later identified as P. acnes (see below). Fourteen days after the operation, the anaerobic cultures of the ground anulus tissue became positive with the same bacterium. The histologic staining of the anulus tissue showed large amounts of gram-positive rods which were morphologically consistent with Propionibacterium species (Fig. 1). After the MICs became known (penicillin, 0.016 mg/liter; amoxicillin-clavulanate, 0.032 mg/liter; cefoxitin, 0.016 mg/liter; chloramphenicol, 0.25 mg/liter; clindamycin, 0.047 mg/liter) the antibiotic therapy was changed to penicillin G (3 million IU every 4 h). This therapy was continued for another 6 weeks. The hemiparesis improved markedly. One year after implantation of the homograft and a double-chamber pacemaker because of bradycardia atrioventricular junctional heart rhythm the patient was found to be in good cardiac condition.

Blood (5 ml of blood per bottle) was cultured with the Septi-Chek (BCB) system (Roche Diagnostics, Basel, Switzerland). The aerobic bottle was vented, and a BCB slide was added, inverted, and observed daily. The anaerobic bottle was incubated anaerobically and subcultured only onto anaerobically incubated brucella sheep blood agar, kanamycin-vancocin mycin laked blood agar, and phenylethyl alcohol blood agar (18) (all media from Becton Dickinson [BBL], Basel, Switzerland) when growth could be observed macroscopically or when
the corresponding aerobic blood culture bottle was positive for growth.

The valves and tissues were transported in sterile containers without any medium. Unfortunately, transport was delayed 36 h by a transportation problem. Cultures of tissue samples were first done in brain heart infusion broth (BHI) incubated aerobically at 37°C without growing to avoid contamination. Seven days after the operation, when histologic and blood culture results became positive but cultures of tissues and valves showed no growth, one-half of the tissue incubated in BHI was ground with a Ten Broeck tissue grinder and was cultured aerobically at 37°C with 5% CO2 on Columbia sheep blood agar as well as on Columbia chocolate agar. Anaerobic cultures were set up on brucella sheep blood agar. Eventually, at 7 days after grinding and 14 days postoperatively, *P. acnes* was isolated from the ground tissue; the unground fraction remained negative.

Identification was done by using the RapID ANA II System (Innovative Diagnostic Systems, Atlanta, Ga.) in combination with gas-liquid chromatography of volatile and nonvolatile fatty acids from prereduced, anaerobically sterilized chopped meat broth containing carbohydrates (9). All isolates produced predominantly propionic acid and showed the same enzymatic pattern (RapID ANA code 004 655, resulting in *P. acnes* with an identification level of >99.9% and a biofrequency of 1 of 37). MICs were determined by the E-test procedure (AB Biodisk, Solna, Sweden) on brucella blood agar, which has been shown to have a good correlation with the methods of the National Committee for Clinical Laboratory Standards (24).

*P. acnes* is a rare cause of serious infections, and it is difficult to prove that it is the causative agent of serious infections in general and of IE in particular (1). Of the 10 cases of *P. acnes* infection described in Table 1, only one case was undoubtedly caused by *P. acnes* (13). For three cases there were few details (1, 15). The remaining seven cases of IE seemed to be possibly or probably caused by *P. acnes* (5, 7, 10, 12, 16, 17).

In our patient two blood cultures each were found to be positive for *P. acnes* 7 and 8 days postoperatively. The tissue sample in BHI showed no growth at those times. The anulus tissue sample was ground only then. Seven days later the anaerobic cultures revealed growth of *P. acnes*, and the organism was biochemically identical to those from the blood cultures.

Recently, another patient underwent surgery for a mycotic aneurysm of the ascending aorta. Blood cultures showed *Salmonella enteritidis*, as did the crushed second half of the aortic tissue sample, in contrast to the compact first fragment, which did not. These two cases point out that grinding of tissue samples is necessary.

In our laboratory, during 1 year, we found *Propionibacterium* species in 138 blood culture sets from 129 patients (0.8% of all blood culture sets processed, or 5.1% of all positive blood cultures within 1 year). Contrary to the results of other investigators (1), two blood cultures positive for *Propionibacterium* species, found in 6 of 129 patients, were not found to prove serious infections. The present case report with four positive blood culture sets showed the only clinically significant infection caused by *P. acnes* during 1 year.

*Propionibacterium* spp. grew more often in anaerobic bottles (*n* = 95), but nevertheless, they were isolated 66 times in aerobic bottles (*P* = 0.004; chi-square test) and 25 times in both anaerobic and aerobic bottles. This finding supports the
fact that Propionibacterium spp. can no longer be considered strict anaerobes (3). In the literature, even in large series of blood culture analyses (19, 20, 22), there are no available comparable data concerning the frequency of growth of Propionibacterium spp. in anaerobic or aerobic bottles.

The mean time of detecting growth of Propionibacterium species in blood cultures was 6.4 days in anaerobic bottles and 6.1 days in aerobic bottles (range, 2 to 15 days). In our case, blood cultures needed 7 and 8 days, respectively, to become positive, in contrast to the tissue sample, which needed 14 days. This points out the importance of incubating blood cultures as well as tissue samples 3 weeks or more in cases in which Propionibacterium infection is suspected. Therefore, communication between clinicians and the laboratory is very important.

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