**CASE REPORT**

Rat bite fever caused by *Streptobacillus moniliformis* in a child: human infection and rat carriage diagnosed by PCR

J-M Andre, A M Freydiere, Y Benito, A Rousson, S Lansiaux, A Kodjo, C Mazzocchi, J-C Berthier, F Vandenesch, D Floret

A 7 year old boy presented with fever, rash on the extremities, and intense bilateral arthralgia in the knees, ankles, elbows, and wrists. Physical examination showed maculopapular morbilliform exanthema on the palms and soles, associated with several blisters 3–8 mm in diameter, containing a whitish fluid, on the face and elbows. None of the painful joints was mobilisable. The right knee and both ankles were swollen.

Laboratory investigations showed an intense inflammatory syndrome (C reactive protein, 300 mg/litre; erythrocyte sedimentation rate, 60 mm/hour), normal leucocyte count, and normal cerebrospinal fluid examination. Samples of cerebrospinal fluid, blood, and blister fluid were cultured.

Possible diagnoses included rat bite fever (RBF). One month before onset the patient had acquired two pet rats. His parents reported frequent physical contact between the boy and his pets, but they were not aware that he had been bitten.

Pending the laboratory results, empirical intravenous combination treatment was started with erythromycin (30 mg/kg/day) and amoxicillin (100 mg/kg/day).

Serological tests were negative for *Mycoplasma pneumoniae*, *Francisella tularensis*, *Leptospira* spp, and *Rickettsia* spp. Polymerase chain reaction (PCR) analyses were negative for *Rickettsia* spp and *Leptospira interrogans*.

Blood and cerebrospinal fluid cultures remained sterile. However, blister fluid samples yielded smooth, shiny, grey colonies 1–2 mm in diameter on horse blood agar plates incubated in aerobic conditions, and on Columbia blood agar incubated in anaerobic conditions, after 48 hours at 36°C. Microscopic examination showed pleomorphic Gram negative rods. Tests for oxidase, catalase, nitrate reductase, and urease were negative. Because growth was slow, despite serum enrichment, biochemical identification was not possible and antibiotic susceptibility testing was also unsuccessful.

The isolate was finally identified using a broad range PCR technique based on amplification and sequencing of bacterial 16S ribosomal DNA. The amplified DNA of the isolate showed 99% similarity with the Genbank *Streptobacillus moniliformis* sequence Z35305 (type strain ATCC16467).

After five days of antibiotic treatment, the patient’s general status improved. At this time he developed spectacular bilateral desquamation of the fingers and toes (fig 1). The echocardiogram was normal, arguing against atypical Kawasaki disease. Erythromycin and amoxicillin were discontinued after seven and 15 days, respectively, without symptom recurrence. The patient was seen 15 days after discharge, and his physical examination was normal. He admitted at this time that he used to eat his pets’ faeces.

One of the apparently healthy pet rats was euthanised. Necropsy showed no macroscopic lesions. Fragments of oropharyngeal and tracheal mucosa were cultured. Various commensal microorganisms (mainly *Pasteurella* sp.) were isolated, together with small Gram negative bacilli with similar morphology to those isolated from the child’s blisters. Despite multiple subcultures, pure colonies of the putative *Streptobacillus* spp could not be obtained. We used molecular tools to detect *S moniliformis*. DNA was extracted from mixed culture on an agar plate and from the rat’s tracheal biopsy. The broad range PCR technique could not be used because of the polymicrobial DNA extracts; therefore, a specific PCR targeting *S moniliformis* using primers described by Boot and colleagues was performed. The 296 bp DNA fragment was amplified and restricted using BfaI, as described previously.

This yielded supernumerary fragments but a *S moniliformis* signature (128, 92, and 76 bp fragments) was detected in the culture and biopsy DNA when compared with the isolated strain (fig 2).

**DISCUSSION**

RBF is a rare zoonosis caused by *S moniliformis*, a small fastidious pleomorphic Gram negative bacillus. It has been

**Abbreviations:** PCR, polymerase chain reaction; RBF, rat bite fever
reported that S moniliformis is present in the nasopharyngeal flora of 50–100% of healthy wild living and laboratory rats, but few recent data are available. Asymptomatic carriage appears to be frequent. RBF is classically transmitted by rat bites, but human cases caused by scratches or mouth to mouth contact have also been described; the mortality rate in untreated cases is about 10–13%. Ingestion of rat stools has not been reported previously.

Molecular methods in the pet of a human suffering from RBF biopsy. To our knowledge, the detection of S moniliformis polymorphism on DNA extracts from the cultures and rat colonies by culture from pet specimens, its presence was identified after culture of the child’s blister fluid and was detected in rat samples by molecular methods. This is the first report of such a detection by molecular methods in the pet of a human victim of rat bite fever that RBF is not a notifiable disease may explain the lack of data on the incidence of this zoonosis. More cases of RBF have been reported in the past decade, but it is difficult to know whether this is because of the increasing number of unusual pets or improved microbiological methods and vigilance in reporting. Molecular studies of S moniliformis carriage among pet rodents would be of interest.

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**Take home messages**
- We report the case of a child who owned pet rats and who developed an eruptive fever with blisters, polyarthritis, and spectacular desquamation of the hands.
- *Streptobacillus moniliformis* was identified after culture of the child’s blister fluid and was detected in rat samples by molecular methods.
- This is the first report of such a detection by molecular methods in the pet of a human victim of rat bite fever.

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