**Rapid Communications**

**Louse-borne relapsing fever (**Borrelia recurrentis**) in an Eritrean refugee arriving in Switzerland, August 2015**

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We report an imported case of louse-borne relapsing fever in a young adult Eritrean refugee who presented with fever shortly after arriving in Switzerland. Analysis of blood smears revealed spirochetes identified as *Borrelia recurrentis* by 16S rRNA gene sequencing. We believe that louse-borne relapsing fever may be seen more frequently in Europe as a consequence of a recent increase in refugees from East Africa travelling to Europe under poor hygienic conditions in confined spaces.

Here we communicate a case of louse-borne relapsing fever (LBRF) in an Eritrean refugee after arrival in Switzerland in August 2015. *Borrelia recurrentis*, the causative agent of LBRF, was identified by 16S rRNA gene sequencing. In addition to diagnostic and therapeutic aspects, we discuss the epidemiology of this potentially re-emerging and serious disease in the context of a recent increase in refugees from East Africa travelling to Europe.

**Case description**

In August 2015, a refugee from Eritrea in their late 20s presented to our emergency department with fever for two days, nausea, headache, dysuria and bilateral flank pain. After leaving Eritrea about two months earlier with stopovers in Sudan (two weeks) and Libya (three weeks), the patient arrived in Italy ca 12 days before presenting to our hospital (Figure 1). On their way through Sudan, the patient had experienced a similar episode of dysuria and flank pain without fever that was not treated. Others travelling with them had febrile illnesses that were diagnosed as malaria. All of the patient’s clothes were exchanged once in Libya and a second time after arrival in Italy.

On presentation, the patient was afebrile and blood pressure and heart rate were within normal limits. Physical examination was significant for suprapubic and right flank tenderness, whereas skin examination was unremarkable. In particular, lice were not detected on the patient’s clothes. Routine blood tests demonstrated mild microcytic anaemia (haemoglobin 101 g/L; norm: 140–180), thrombocytopenia (108 x 10⁹/L; norm: 150–450) and elevated C-reactive protein (CRP: 108 mg/L; norm: < 10). Urine analysis revealed mild pyuria (266 /µL; norm 0–40) and abundant squamous epithelial cells (45/µL; norm 0–9). Blood smears demonstrated a large number of extracellular spirochetes (Figure 2). Blood cultures remained negative and only bifidobacteria grew in urine culture. Serological screening tests for human immunodeficiency virus (HIV), syphilis and Lyme disease were negative.

The patient was initially treated with doxycycline 100 mg twice per day over two days and ceftriaxone 2 g per day to cover *Borrelia*-associated relapsing fever and suspected pyelonephritis. We did not observe a Jarisch–Herxheimer reaction [1]. They remained afebrile throughout the admission, and their pain and dysuria settled immediately after initiation of antibiotic treatment. Ceftriaxone was stopped after five days of treatment and the patient was discharged for outpatient follow-up.

**Investigation of the disease-causing agent**

Subsequently, analyses were performed to further characterise the causative agent of the patients’ relapsing fever syndrome. The detection and identification of *B. recurrentis* was performed with broad-range bacterial PCR followed by partial sequencing of the 16S rRNA gene applied to two EDTA blood specimens drawn two hours (before treatment, microscopy-positive) and 18 hours (after the first dose of ceftriaxone, microscopy-negative) after presentation to our hospital [2]. PCR
was positive only on the first sample (100% identity with the *B. recurrentis* reference strain A1).

In a very recent LBRF patient, we additionally analysed the entire 16S rRNA gene, which yielded a 1,475 bp sequence (GenBank accession number: KT221542). Nucleotide sequence database analysis using the Basic Local Alignment Search Tool (BLAST, National Library of Medicine) showed 100% identity to the *B. recurrentis* reference strain A1 [3].

**Background**

LBRF once was a major worldwide endemic disease causing significant mortality in untreated patients (10–70%) [4-6]. The incubation period is usually short (four to eight days with a range of two to 15 days). In contrast to other *Borrelia*-associated relapsing fever syndromes, LBRF is considered a disease specific to the human host caused by *B. recurrentis* and transmitted by the human body louse *Pediculus humanus humanus* as the vector. In parallel with the markedly reduced incidence of body louse infestations after the 1940s, LBRF cases declined significantly worldwide with the exception of East African countries, in particular Ethiopia, where LBRF remains a common cause of hospital admission and death [7]. Interestingly, cases of LBRF have not recently been reported outside of Africa, including in refugees from affected countries, which may be indicative of a true decline in incidence or a lack of outbreaks in affected countries [8] rather than of difficulties in diagnosis, although microscopy of blood films has a low sensitivity and cannot differentiate between *Borrelia* species [9]. Hence, duration of treatment for *Borrelia*-associated relapsing fever syndromes, which differs according to the species, is usually determined based on geographical location and the respective predominant vector. Whereas single-dose procaine penicillin, tetracycline (500 mg) or doxycycline (200 mg) are effective treatment regimens for LBRF, treatment duration with doxycycline, penicillin G or ceftriaxone is much longer (10 to 14 days) in the case of TBRF.

**Discussion**

The current arrival of refugees from East Africa to Europe poses challenges for clinicians in European countries as they may be faced with tropical diseases including malaria [10] and diseases nowadays rarely diagnosed in Europe; for example, cases of cutaneous diphtheria have recently been diagnosed in refugees from East Africa at our institution (data not shown). In addition, some of these diseases have the potential to cause small outbreaks in refugee camps [11,12]. We present a case of LBRF imported into Europe in the context of the ongoing migration of refugees from Africa.

Taking into account the short incubation period and the patient’s travel dates with a possible episode while residing in Sudan, infection with LBRF probably occurred in Eritrea, Sudan or Libya. Apart from Ethiopia, LBRF has previously been reported from Eritrea and

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**Figure 1**

Travel route from Eritrea to Europe, louse-borne relapsing fever case, Switzerland, August 2015

**Figure 2**

Microscopic detection of spirochetes in blood, louse-borne relapsing fever case, Switzerland, August 2015

Panel A: Giemsa pH 7.2, stained thick film, 1,000-fold magnification.
Panel B: May-Grünwald Giemsa (MGG)-stained blood smear, 1,000-fold magnification.
were conducted. Comprehensive source tracing is difficult, in our opinion, as it is impossible to locate and investigate all refugees that have previously travelled with a case in crowded conditions. While the human body louse has been considered the principal vector of LBRF, *B. recurrentis* DNA was recently detected in head lice (*Pediculus humanus capitis*) [15], although it remains to be determined whether *B. recurrentis* can be transmitted via head lice from person to person. In any case, we consider screening arriving refugees for lice useful in order to prevent spreading of louse-borne diseases in refugee camps. Given that our case’s clothes had been exchanged twice before arrival in Switzerland and the fact that lice were not detected in their present clothes, neither formal contact tracing nor additional screening of refugees living in the same asylum seeker camp for lice or preventive delousing were conducted. Comprehensive source tracing is difficult, in our opinion, as it is impossible to locate and investigate all refugees that have previously travelled with a case in crowded conditions.

Patients with LBRF usually present with non-specific symptoms such as high fever, headache or pain in other parts of the body [16]. Hence, presentation of LBRF may resemble many other serious infections such as malaria, viral haemorrhagic fever, leptospirosis, typhus, TBRF, meningococcal meningitis or typhoid fever [17]. In addition, co-infection with malaria is common [16], although not detected in our patient. Most patients with LBRF are diagnosed using microscopy. This method lacks interspecies differentiation of *Borrelia* species causing relapsing fever syndromes, which is important for determining treatment duration (see below). Molecular diagnostic tests such as multiplex PCR or 16S rRNA gene analysis or are not universally available but can aid in the diagnosis of LBRF [18]. Our case demonstrates that the detection and identification of the aetiologic agent can be performed by 16S rRNA gene analysis directly from blood samples. Physicians should consider LBRF as differential diagnosis in refugees from East Africa presenting with fever of unknown origin, as mortality in untreated patients is high. Single-dose procaine penicillin, tetracycline (500 mg) or doxycycline (200 mg) are effective treatments for LBRF; the evidence for the superiority of tetracyclines is weak with regards to the risk of relapse and time to defervescence [1]. On the other hand, physicians should be aware of higher rates of the Jarisch–Herxheimer reaction with the use of tetracyclines [1].

**Conclusion**

Physicians in countries currently hosting Eritrean refugees need to consider LBRF in febrile migrants in addition to more common diseases like malaria.

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**Conflict of interest**

None declared.

**Authors’ contributions**

Wrote the manuscript: DG, GJC, AE, MO; performed laboratory investigations: DG, MM, AB, BS; revised manuscript: DG, GJC, CB, TB, MM, AN, AB, AE, MO; managed the patient: GJC, CB, TB, MO.

**References**


