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To cite this version:

Short communication

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Detection of *Francisella tularensis* subsp. *holarctica* in a European brown hare 
(*Lepus europaeus*) in Thuringia, Germany

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Abstract

The isolation of *Francisella tularensis* subsp. *holarctica* biovar II (strain 06T0001) from a European brown hare (*Lepus europaeus*) from Thuringia, Germany, is described for the first time. Identification of the microorganism was carried out by phenotypic characterisation, partial sequencing of the 16S rRNA gene and specific PCR using the primers TUL4-435/TUL4-863 and FtC1/FtC4. The epidemiology of tularemia in Germany is discussed and a risk assessment for humans is made.

Keywords: tularemia, *Francisella tularensis*, PCR, *Lepus europaeus*
1. Introduction

Tularemia is an infectious disease caused by the small, pleomorphic, heat-labile, Gram-negative, rod-shaped bacterium Francisella (F.) tularensis. The microorganism is a facultative intracellular pathogen affecting a wide range of animal species with more hosts than any other known zoonotic pathogen (Johannson et al., 2004). The main hosts are believed to be hares and rabbits (Leporidae), hamsters, water and field voles (Cricetidae), water and wood rats, lemmings and field mice (Muridae), squirrels (Sciuridae), and also aquatic rodents like beaver (Castoridae) and muskrats (Ondatrae). Other mammalian species susceptible for F. tularensis are monkeys (Anthropoidae), dogs and coyotes (Canidae), cats (Felidae), sheep (Ovidae) and cattle (Bovidae) but also predators like bears (Ursidae), and foxes (Vulpes). Infection of lagomorphs and rodents with F. tularensis often causes fatal disease. Usually, infected animals are found moribund or dead. Also several bird species as well as fish and amphibians are considered to be incidental hosts (Splettstoesser et al., 2005, Anda et al. 2001). Further vectors for transmission of F. tularensis are ticks, biting flies and mosquitoes (Blaškovič and Barak, 2005, Hubalek et al., 1996).

Tularemia is endemic in many regions of the Northern hemisphere worldwide. The species F. tularensis includes three subspecies (2 types). Type A strains, F. tularensis subsp. tularensis, have so far been found predominantly in North America, although type A strains have been isolated in the Danube region close to Bratislava (Guryčova, 1998). Type B strains (F. tularensis subsp. holarctica), F. tularensis subsp. mediasiatica and F. novicida are found in Europe, Northern Asia and Northern America (Ellis et al., 2002).

In Germany in 1983, 1990, 1991 and 1992 four cases of tularemia in hares or rabbits were notified from Lower-Saxony, Rhineland-Palatinate, North-Westfalia and
Baden-Württemberg, respectively (BMELV, 2006).

In November 2004 an outbreak of tularemia was reported among marmosets (Callithrix jaccus) in Central Lower Saxony, Germany (Splettstoesser et al., 2007). The detection of *F. tularensis* subsp. *holarctica* was a re-emergence of tularemia in Germany (BMELV, 2006).

In 2005 15 cases of tularemia were serologically confirmed in persons in Hesse who had contact with hares (Hofstetter et al., 2006, Splettstoesser et al., 2006). *F. tularensis* subsp. *holarctica* was also detected in several organ samples taken from these hares.

A variety of PCR methods has been established for the detection of *F. tularensis* DNA in both clinical and environmental specimens (Splettstoesser et al., 2005). For species identification PCR assays were used targeting the *tul4* gene (accession number M32059) which encodes a 17 kDa outer membrane protein (Sjöstedt et al., 1997). By Johansson et al. (2000) a PCR assay was evaluated to identify *F. tularensis* subsp. *holarctica* and distinguish it from all other subspecies (accession number AF240631).

Here we report a case of tularemia in a European brown hare. The causative agent *Francisella tularensis* was identified by phenotypic characterisation, Gram staining, agglutination test, DNA sequencing, PCR, and susceptibility to erythromycin.

2. Materials and Methods

Material
In February 2006, an European brown hare (Lepus europaeus) was shot and sent for microbiological investigation with the report of abnormal flight behaviour. It was an adult male weighing 3,600 g. No externally visible distinctive pathological-anatomical changes were observed. After opening of the carcass and examination of the organs the following significant morphological changes were found: swelling of body lymph nodes, moderate hyperplastic spleen tumour, swelling of the kidneys, diffuse reddening and petechiae of the tracheal mucosa and mucoid inflammation of the colon.

Isolation and cultivation of Francisella tularensis

Bacteria originating from brain, lung, liver, spleen, kidney, sections of small intestine, colon and bile were cultivated on blood agar plates by means of loop inoculation and incubated under aerobic conditions at 37°C for 3-4 days. In parallel, growth of Francisella bacteria was tested under aerobic and microaerophilic conditions on chocolate agar with 2% cysteine and cystine heart-agar (Oxoid, Wesel, Germany) with 9% sheep blood. Growth of F. tularensis was confirmed by the slide agglutination assay with a specific antiserum (BgVV, Berlin, Germany) and standard Gram-staining. The susceptibility of bacteria to erythromycin was examined (Etest™, Erythromycin discs [15 µg], Oxoid).

Differential diagnostic investigations

Other differential diagnostic assays applied were hemadsorption test for detection of European brown hare syndrome virus (EBHS-V), ELISA antigen assay for Chlamydia/Chlamydophila detection, direct immunofluorescence test for rabies
detection, general bacteriological investigation, and parasitological investigation of lung, small intestine and colon sections by means of flotation, sedimentation and migration techniques.

PCR detection and partial sequencing of 16S rRNA gene

A bacterial colony was resuspended in 100 µl phosphate-buffered saline and boiled for 10 minutes. The DNA was extracted using the High Pure PCR Template Preparation Kit™ (Roche Diagnostics, Mannheim, Germany) according to the recommendations of the manufacturer. The DNA was eluted in 200 µl elution buffer and 5 µl were applied in each PCR assay.

The PCR was carried out as described by Johansson et al. (2000). Briefly, the reaction mixture consisted of 5 µl 10x PCR buffer with 1.5 mmol MgCl₂ (Genaxxon, Stafflangen, Germany), 2 µl of dNTP mix (2.5 mM of each dNTP, Carl Roth GmbH, Karlsruhe, Germany), 1 µl of each of forward and reverse primers (10 pmol/µl, JenaBioscience, Jena, Germany), 0.2 µl of Taq DNA polymerase (5 U/µl, Genaxxon), 5 µl of DNA extract and deionised water to a final volume of 50 µl. The primers used are shown in table 1. After denaturation at 94 °C for 60 s, 35 cycles of amplification were performed according to the following protocol: denaturation at 94 °C for 30 s, primer annealing at 60 °C for 60 s, and primer extension at 72 °C for 30 s. After the final extension step at 72 °C for 60 s, each reaction mixture was subjected to electrophoresis in a 2% agarose gel (Eurogentech, Cologne, Germany). After ethidium bromide staining, the PCR products were visualized by UV light and documented using BioImage system GeneGenius (Syngene, Synoptics Ltd., UK).

Amplicons obtained by using primer pair 16S UNI-L/R (Kuhnert et al., 1996) and DNA of isolate 06T0001 were cut out of agarose gel plug. DNA was purified using
QIAquick Gel Extraction Kit™ (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. The extract was subjected to cycle sequencing with BigDye™ Terminator Cycle Sequencing Ready Reaction Kit (Applied Biosystems, Darmstadt, Germany). Amplification primers 16S UNI-L and 16S UNI-R were used as sequencing primers, too. Nucleotide sequences were determined on an ABI PRISM 310 Genetic Analyzer (Applied Biosystems). Resulting sequences were compared with those from public database entries (http://www.ncbi.nlm.nih.gov/blast/).

3. Results

The pathological findings in the sick hare were concordant with the clinical picture of tularemia in this species. *F. tularensis* could be isolated from different organs (lung, liver, spleen, kidney, small intestine section and bile) of this animal. After incubation for 72 to 96 h at 37°C, numerous glossy colonies were seen on chocolate agar plates. The characteristic glossy grey or dark green staining of the agar in the colony area was distinct on cysteine heart agar. The bacteria were Gram-negative, very small, had a coccoid rather than a bacillary shape, and showed resistance to erythromycin. Using the primer pair 16S UNI-L/16S UNI-R a 1,200 bp fragment of the 16S rRNA gene was amplified. Partial sequencing of this fragment and BLAST search resulted in the identification of the bacterium as a member of the species *F. tularensis*. This result was in concordance with species-specific PCR results using primers TUL4-435/TUL4-863 (Sjöstedt et al., 1997).
For distinguishing the *F. tularensis* subsp. *holarctica* from other *F. tularensis* subspecies, the primer set FtC1/FtC4 (Johansson et al., 2000) was applied. The amplicon obtained with DNA of isolate 06T0001 was only 300 bp in length suggesting that this isolate was a member of the subspecies *holarctica*. But the amplicons for *F. tularensis* subsp. *tularensis*, subsp. *mediasiatica*, *F. novicida* and *F. philomiragia* were 330 bp in size.

Other differential diagnostic investigations were applied including the hemadsorption test for detection of European brown hare syndrome virus (EBHS-V) and, *Chlamydia/Chlamydophila* capture ELISA, direct immunofluorescence test for rabies detection were negative. Parasitological investigation of the lungs, small intestine and colon resulted in detection of *Trichuris* and other nematode species, *Dicrocoelium* flatworms and *Eimeria* spp.

In general bacterial investigations *Mannheimia haemolytica* was detected in lung tissue and *Clostridium perfringens* type A in the colon.

### 4. Discussion

Tularemia is widespread in Europe, its presence is reported from all countries except UK, Iceland, and Portugal (Tärnvik et al., 2004). In most countries, only sporadic human infections are reported every year, whereas in others, e.g. Sweden or Finland, outbreaks comprising hundreds of cases are recorded (Tärnvik et al., 2004). The first case of tularemia in Germany was reported 50 years ago (BMELV 2006). In October 2004 an outbreak of this disease in a semi-free living group of marmosets in Lower Saxony was detected (Splettstoesser et al., 2007), and in December 2005 infections in humans were reported from Hesse (Hofstetter et al. 2006, Splettstoesser...
et al., 2006). Here, a case of tularemia in a hare in Thuringia is described. All these cases were found within a region known as “Central Germany”. The confirmation of autochthonous infections in the geographical centre of Germany will stimulate the historical debate on the origin of tularemia in Central Europe and on its natural reservoirs, routes of transmission and the ecological niches of the bacteria. In this geographical area characteristic conditions are predominant for typical natural foci of tularemia (Pikula et al., 2003): alluvial forests, < 200 m above sea level, 8°-10°C mean annual temperature, 450-700 mm mean annual precipitation, and 2,000-2,200 h mean annual sunshine duration.

The most important natural occurring reservoir of *F. tularensis* may be amoeba or protozoa in surface water and ticks (e.g. *Dermacentor reticulatus* and *Ixodes* species) as well as fleas (Guryčova et al., 2001). Therefore the tenacity of the infectious bacteria was found in dust, damp hay, carcasses, water etc. The bacterium can survive in the environment and in vectors for a relevant period of time. The assumption that rodent and tick populations of non-endemic areas are very susceptible to a newly introduced, highly contagious and virulent agent may explain the focal emergence of tularemia in hitherto unaffected areas like Thuringia.

In the reported case a hare shot by a hunter was sent to a local authority for investigation because of its abnormal flight behaviour. It showed significant morphological changes of organs, like swelling of body lymph nodes, moderate spleen tumour, swelling of kidneys and diffuse reddening and petechiae of tracheal mucosa. These lesions are typical for tularemia. The investigation of the bacterial colonies revealed results, which were concordant with *Francisella* bacteria. Using DNA sequencing of 16S rRNA gene the occurrence of *F. tularensis* was confirmed. Consequently, we used the PCR assays as the latest diagnostic method for detection of *F. tularensis* (Splettstoesser et al. 2005). With PCR using primer pair FtC1/FtC4 for
subspecies identification of the investigated bacteria the agent could be identified as

*F. tularensis* subsp. *holarctica*. Erythromycin resistance was observed in the isolate
and was therefore assigned to biovar II (Tomaso et al., 2005). This finding indicates
that the possible geographic origin of the isolate is Eastern Europe or that the
Western border of this well-known region of endemicity is located in the German
mountain range.

This case of tularemia in a hare is the first report on the presence of this zoonosis in
Thuringia, Germany. The increased epizootic activity of *Francisella tularensis* in the
endemic region of Central Europe shows that tularemia has to be considered an
emerging disease for animals and humans too. The influence of Greenhouse effect
and the obviously changing climate in Central Europe might have an crucial impact
on the future spread of the disease and the stable establishing of permanent
endemic areas. Thus a higher risk for wildlife, pet animals and humans has to be
expected. Therefore, the aim of further investigations will be the characterisation of
the epidemiology of tularemia in whole Europe.

**Acknowledgements**

We thank the staff of the collaborating laboratories for their support. We wish to thank
Prof. Ian N. Clarke, Southampton, for reviewing the English language of the
manuscript.

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Table 1: Sequences of primers

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>Fragment size</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>16S UNI-L</td>
<td>AGA GTT TGA TCA TGG CTC AG</td>
<td>1,200 bp</td>
<td>Kuhnert 1996</td>
</tr>
<tr>
<td>16S UNI-R</td>
<td>GTG TGA CGG GCG GTG TGT AC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TUL4-435</td>
<td>GCT GTA TCA TCA TTT AAT AAA CTG CTG</td>
<td>400 bp</td>
<td>Sjöstedt 1997</td>
</tr>
<tr>
<td>TUL4-863</td>
<td>TTG GGA AGC TTG TAT CAT GGC ACT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FtC1</td>
<td>TCC GGT TGG ATA GGT GTT GGA TT</td>
<td>300 bp/330 bp</td>
<td>Johansson 2000</td>
</tr>
<tr>
<td>FtC4</td>
<td>GCG CGG ATA ATT TAA ATT TCT CAT A</td>
<td></td>
<td></td>
</tr>
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