Detection of *Frederiksenia* sp. isolated from a cat with nephritis – Short communication

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**ABSTRACT**

In this paper we report the phenotypic and partial genetic characterisation of a novel bacterium strain isolated from a cat with severe nephritis. Multilocus sequence analysis was performed on the 16S rRNA and three housekeeping (*recN*, *rpoB*, *infB*) gene sequences obtained by PCR. In accordance with the results of phenotypic tests, the phylogenetic analyses confirmed the relatedness of the new strain (6036) to the family *Pasteurellaceae*. On the phylogenetic trees, strain 6036 appeared in a separate branch, closest to that of the type species (*Frederiksenia canicola*) of the genus *Frederiksenia*. These two bacteria shared 95.14 and 76.88% identity in their partial 16S rRNA and *recN* gene sequences, respectively. The *rpoB*- and *infB*-based phylogenetic analyses indicated that strain 6036 is most closely related to *Bibersteinia trehalosi* (with 90.58% identity) and *Haemophilus felis* ATCC 49733 (89.50% identity), respectively. The predicted genome identity values, based on the *recN* gene sequences, suggested that strain 6036 can be classified into the genus *Frederiksenia* as a novel species. A PCR method, specific to strain 6036, was developed to allow its rapid and accurate identification and differentiation from *F. canicola* and other species of *Pasteurellaceae*. The minimal inhibitory concentrations of 18 antimicrobial agents for strain 6036 were also determined.

**KEYWORDS**

*Frederiksenia*, *Pasteurellaceae*, phylogeny, antimicrobial susceptibility, MIC

The family *Pasteurellaceae* is a large bacterial family that comprises 30 genera with validly published names (Christensen and Bisgaard, 2018; Dickerman et al., 2020). Bacteria belonging to the family *Pasteurellaceae* are commonly isolated from various animal species and humans; most are considered opportunistic pathogens (Olsen et al., 2005). Several newly described genera have recently been added to the family, including the most recently classified *Caviibacterium* and *Conservatibacter*, which are associated mainly with guinea pigs (Adhikary et al., 2018), and *Glaesserella* established by the reclassification of *Haemophilus parasuis* (Dickerman et al., 2020). These are expected to be supplemented with further new genera and species in the future.

Members of the family *Pasteurellaceae* have frequently been isolated from the oral cavity and upper respiratory tract of cats and dogs (Christensen and Bisgaard, 2008), including *Haemophilus felis*, *Frederiksenia canicola* and species of *Pasteurella* sensu stricto (*Pasteurella multocida*, *Pasteurella dagmatis*, *Pasteurella canis*, *Pasteurella stomatis* and *Pasteurella oralis*) (Mutters et al., 1985; Inzana et al., 1992; Korczak et al., 2014). Recently, *P. dagmatis*-like organisms have also been identified (Sellyei et al., 2010) from the oral cavity of cats. These species often cause bite wound infections in humans (Mutters et al., 1985).

Acquiring knowledge of newly isolated species is essential for epidemiological studies and routine diagnostic clinical microbiology laboratories. The aim of this study was to characterise a phenotypically and phylogenetically distinct strain, isolated from a diseased cat, that differed from the other members of *Pasteurellaceae*. 
In September 2018, a 4-month-old female Maine Coon cat suddenly died without apparent clinical signs and was submitted for routine postmortem examination to the NFCSO Veterinary Diagnostic Directorate (Budapest, Hungary). Tissue samples were collected from brain, heart, lungs, liver, spleen, kidney and mesenteric lymph nodes, and after fixation in 10% formalin they were processed for histological and immunohistochemical (IHC) examinations. Besides Giemsa staining, tissue sections were examined with an anti-Mycobacterium bovis-based IHC (Szeredi et al., 2008). This method detects several bacteria, fungi and parasites, and it is far more sensitive than traditional histological staining procedures, detecting even low numbers of the above-listed pathogens in tissue samples. Furthermore, tissue sections were immunolabelled with an anti-Bibersteinia trehalosi reagent, which shows cross-reaction with Mannheimia haemolytica and P. multocida types A, B and D by IHC (Szeredi et al., 2010). Samples from kidney and spleen were cultured on Columbia agar (LAB M Ltd., Bury, UK) supplemented with 5% sheep blood under aerobic conditions at 37 °C for 24 h. Biochemical tests were performed for phenotypic characterisation according to the recommendation of Christensen et al. (2007).

Postmortem examination revealed severe icterus and slightly enlarged lungs, spleen and mesenteric lymph nodes presenting congestion. Kidneys were enlarged about 1.5× to the normal size, they were dark red in colour, and numerous greyish-white foci of 1 mm in diameter were dispersed in the cortex. The main histological lesion was observed in the kidneys presenting severe embolic supplicative nephritis. Large numbers of bacterial emboli occluded the glomerular capillaries. Numerous acute thrombi associated with large areas of necrosis and haemorrhages were also evident. Lesions indicative of bacterial septicaemia were detected in other organs as well: acute serous hepatitis accompanied by scattered foci of acute inflammation and necrosis, acute sepsis-purulent myocarditis with segmental or circumflex vascular necrosis, mild acute interstitial pneumonia, mild non-purulent encephahtitis, and lymphoid depletion in the spleen. Large numbers of bacteria were detected in the glomerular capillaries of the kidney with Giemsa staining and by an anti-M. bovis-based IHC method. Similar bacteria were identified in the foci of acute necrotic inflammation in the liver, and in the sinusoids of the spleen. These bacteria were not immunolabelled by an anti-B. trehalosi reagent (Fig. 1).

From the kidney and the spleen, Pasteurella-like bacterial colonies were grown in pure culture, which we designated strain 6036. Strain 6036 was non-haemolytic and after 24 h of aerobic incubation on Columbia agar plates the colonies were 0.5–1 mm in diameter with regular circular shape. They did not adhere to the agar surface. Their surface was smooth, shiny and greyish. The cells were Gram-negative, non-motile rods. V- or X-factor dependence was not observed and strain 6036 did not grow on MacConkey agar. Strain 6036 was oxidase, catalase and indole positive, but negative for ornithine decarboxylase and urea. Acid was produced from glucose, maltose and xylose, but not from trehalose, lactose, arabinose, dulcitol and sorbitol.

Fig. 1. A–C. Kidney, cat: A: Severe embolic (arrow) glomerulonephritis associated with acute haemorrhages and necrosis. Haematoxylin and eosin (HE), ×100; B: Several bacterial emboli (arrows) in the capillaries of glomeruli. Giemsa, ×100; C: Strong immunostaining of bacterial emboli. Anti-Mycobacterium bovis based IHC, ×100. Inset: Bacterial emboli are not immunostained with reagent specific for B. trehalosi, M. haemolytica and P. multocida A, B and D. IHC, ×200; D: Liver, cat. Acute serous hepatitis associated with foci of acute inflammation and necrosis (arrows). HE, ×200; Inset: Large amount of bacteria are immunostained in the area of inflammatory infiltration and in focal necrosis. Anti-M. bovis based IHC, ×200

Phylogenetic analysis was performed by sequencing segments of their 16S rRNA, rpoB (encoding the beta subunit of the RNA polymerase), recN (encoding the DNA repair protein) and infB (encoding translation initiation factor 2) genes as described previously (Korczak et al., 2014). Sequencing of PCR products was performed by Macrogen Europe (Amsterdam, The Netherlands). Nucleotide sequences were aligned and compared using Geneious Prime software (version 2019.2.1; http://www.geneious.com). The GenBank accession numbers for sequences obtained in this study are MN701139–MN701142. Nucleotide sequence data were analysed using MEGA7 software (Kumar et al., 2016). The evolutionary history was inferred using the neighbour-joining model and the dataset was subjected to bootstrap analysis of 1000 replicates. Multilocus sequence analysis (MLSA) was also performed (Korczak et al., 2014). Sequence identity data of the four genes were calculated using the Geneious Prime program and used in the formula described by Zeigler (2003):

\[ SI \text{ genome } = -1.30 + 2.25(S\text{rec}N), \]

where SI is sequence identity.

Phylogenetic trees based on the alignment of the 16S rRNA gene sequences (Fig. 2) and MLSA of the four concatenated genes confirmed the relatedness of strain 6036 to the Pasteurellaceae family and placed it on a species-like branch (Fig. 3). The 16S rRNA gene sequence comparison showed the highest sequence identity with the type strain of F. canicola (95.14%). Based on rpoB sequence identity, strain 6036 showed the highest identity with B. trehalosi (90.58%). InfB-based phylogeny showed the closest relatedness of
strain 6036 with *Haemophilus felis* ATCC 49733 (89.50%); the recN sequence identity was the highest with the type species of the *Frederiksenia* genus (76.88%) again. Lower genetic relatedness was detected when comparing the 16S rRNA (91.28–92.43%), rpoB (84.37–86.37%), infB (78.86–82.43%) and recN (59.55–61.11%) gene sequences with species representing the genus *Pasteurella*. Sequence data confirmed that strain 6036 is reliably identifiable using DNA sequence-based approaches.

Calculation of whole-genome relatedness by the presented recN sequence-based method is a reliable application for cross-comparison of *Pasteurellaceae* species and serves as an alternative to the classical hybridisation method (Zeigler, 2003, 2005; Kuhnert and Korczak, 2006; Christensen et al., 2012). Genome similarity values calculated for strain 6036 are listed in Table 1. Comparison of predicted genome similarity of strain 6036 and genera of *Pasteurellaceae* revealed that the similarity of *F. canicola* (strain HPA 21) and strain 6036 was 0.43, which is above the threshold of 0.4 for genus separation (Kuhnert and Korczak, 2006). However, this value is below the threshold of 0.85 for species separation, which indicates that strain 6036 can be classified to the *Frederiksenia* genus but represents a species different from *F. canicola*.

A PCR based on the recN gene sequence was developed for identification and discrimination from other species.
Primers 6036-F TTTGAATGGCTCAAACAGC and 6036-R CGGCAAGTTCTACACAACG, amplifying an 862-bp fragment, were used in the reaction. The reaction was complemented using primers PasrpoB-L and RpoB-R amplifying a 560-bp fragment of the rpoB gene (Korczak et al., 2004). Thus, by conducting a multiplex PCR we were able to exclude the possibility of a negative reaction due to PCR inhibition or lack of enough DNA. PCR was performed in a total volume of 25 µl containing 2.5 µl of 10× DreamTaq buffer, 0.5 mM MgCl₂, 200 µM dNTPs, 0.5 µM of each primer and 1 U of DreamTaq polymerase (Thermo Scientific, Waltham, USA). Reactions were run in a Bio-Rad C1000 Touch thermal cycler. DNA was amplified for 30 cycles using the following settings: denaturation at 95 °C for 30 s, annealing at 60 °C for 30 s and extension at 72 °C for 1 min. An initial denaturation step (95 °C, 10 min) and a final extension step (72 °C, 7 min) were also performed. Each reaction mixture was analysed using electrophoresis in a 1.5% agarose gel (SeaKem, Lonza, Basle, Switzerland) stained with GelRed (Biotium Inc., Hayward, USA) and visualised under UV light. Strain 6036 and species representing the family Pasteurellaceae were used to validate the PCR. The newly designed species-specific PCR clearly discriminated strain 6036 from other members of Pasteurellaceae. The PCR resulted in the expected amplification product with strain 6036 but did not produce any PCR product with other tested Pasteurellaceae species (Fig. 4).
Antimicrobial susceptibility was tested to 18 antimicrobial agents using minimal inhibitory concentration (MIC) test strips according to the manufacturer’s instructions (Liofilchem, Roseto, Italy; Table 2). *Escherichia coli* ATCC 25922 served as a quality control strain. MIC values were read after 24-h incubation at 37°C and were defined as the lowest concentration with complete inhibitory action according to the CLSI standards and as described in a previous study (CLSI, 2018a; CLSI, 2018b; Gutman et al., 2016). Very limited information is available on the antibiotic susceptibility of *F. canicola*, a novel isolate from Tasmanian devils (Gutman et al., 2016). *Frederiksenia* is a relatively recently described genus (Korczak et al., 2014), so there are no breakpoints for antibiotic resistance testing. It is

### Table 1. Calculated genome similarity values of strain 6036 to type species of Pasteurellaceae genera based on recN sequences

| Isolate 6036 | Isolate 6036 |
|--------------|--------------|
| Frederiksenia canicola HPA 21 | 0.43 |
| Vesperilibacter pulmonis DSM 27238 | 0.39 |
| Ursidibacter maritimus Pb43106 | 0.38 |
| Muribacter muris NCTC 12432 | 0.38 |
| Otardidibacter oris CCUG 59994 | 0.37 |
| Bibersteinia trehalosi NCTC 10370 | 0.36 |
| Glaesseriella parasuis CCUG 3712 | 0.35 |
| Nicoletella semolina CCUG 43639 | 0.35 |
| Actinobacillus lignieresii NCTC 4189 | 0.32 |
| Mannheimia haemolytica NCTC 9380 | 0.28 |
| Phocoenobacter uteri NCTC 12872 | 0.27 |
| Avibacterium gallinarum NCTC 11188 | 0.11 |
| Volucribacter psittaccicida CCUG 47536 | 0.10 |
| Conservatibacter flavesens CCUG 24852 | 0.10 |
| Haemophilus influenzae ATCC 33391 | 0.09 |
| Pasteurella stomatis CCUG 17979 | 0.07 |
| Aggregatibacter actinomycetemcomitans ATCC 33384 | 0.07 |

Fig. 4. Multiplex PCR resulting in an 862-bp fragment exclusively with strain 6036, and a positive signal for all investigated strains amplified with primers PasrpoB-L and RpoB-R, respectively. Lane M: molecular weight standard; lane 1: strain 6036; lane 2: Frederiksenia canicola HPA21; lane 3: Pasteurella multocida CCUG 17976; lane 4: Pasteurella dagmatis CCUG 12397; lane 5: Pasteurella canis CCUG 12400; lane 6: Rodentibacter pneumotropicus NCTC 8141; lane 7: Mannheimia haemolytica NCTC 9380; lane 8: Bibersteinia trehalosi NCTC 10370; lane 9: negative control. 

### Table 2. Minimum inhibitory concentrations (MIC) of antimicrobial agents for isolate 6036

| Antimicrobial agent | MIC for isolate 6036 (μg/ml) |
|---------------------|-----------------------------|
| Penicillin G        | 0.023                       |
| Ampicillin          | 0.047                       |
| Streptomycin        | 96                          |
| Gentamicin          | 8                           |
| Tetracycline        | 0.38                        |
| Doxycycline         | 0.38                        |
| Erythromycin        | 0.75                        |
| Clindamycin         | 2                           |
| Enrofloxacin        | 0.006                       |
| Ciprofloxacin       | 0.012                       |
| Nalidixic acid      | 0.75                        |
| Chloramphenicol     | 0.75                        |
| Florfenicol         | 0.5                         |
| Cefazolin           | 0.19                        |
| Cefpodoxime         | 0.016                       |
| Colistin            | 0.5                         |
| Sulphamethoxazole   | 16                          |
| Trimethoprim – sulphamethoxazole | 0.19 |
Therefore not possible to determine unequivocally whether an isolate is susceptible or resistant to a given antibiotic, but it can be predicted from the exact MIC values. Low MIC values were determined for most of the antimicrobials tested (Table 2), including penicillin, ampicillin, tetracycline, doxycycline, erythromycin, enrofloxacin, ciprofloxacin, nalidixic acid, chloramphenicol, florfenicol, ceftazolin, cepodoxime, colistin, sulphamethoxazole and trimethoprim/sulphamethoxazole. Increased MIC values were obtained for streptomycin, sulphamethoxazole and trimethoprim/sulphamethoxazole. The MIC value for sulphamethoxazole (16 μg/mL) was not considered to be elevated because pathogens of veterinary importance can only be considered resistant to sulphonamides at an MIC equal to or greater than 512 μg/mL (CLSI, 2018a, 2018b). Aminoglycoside resistance occurs widely in Gram-negative bacteria, usually because of enzymatic inactivation of the drug (Kehrenberg et al., 2005; Giguère, 2013). Our results confirm the findings of Gutman et al. (2016), who found that resistance occurred intrinsically for aminoglycosides and lincosamides in bacterial species belonging to the family Pasteurellaceae. This highlights that avoiding the use of the aforementioned antibiotics can enable the application of appropriate therapeutic treatments and prevent the spread of antibiotic resistance.

The bacterium examined in the study was highly pathogenic for this kitten, causing severe glomerulonephritis and septicaemia. In the kidney and liver, the presence of bacteria was demonstrated by both Giemsa staining and an anti-M. bovis-based IHC method. The Pasteurella-like bacterium was antigenically clearly differentiated from B. trehalosi, M. haemolytica, P. multocida types A, B and D by the IHC test. Results of phenotypic and genetic analyses suggest that strain 6036 represents a novel species of the Frederiksenia genus, which is supported by individual and combined gene sequence analysis of the 16S rRNA gene and three housekeeping genes. Calculation of whole-genome relatedness with the recN sequence also confirmed this finding. This genus currently includes only the species F. canicola and, compared with strain 6036, we found differences in both genetic and phenotypic characteristics. The biochemical characteristics of strain 6036 show that xylose can be used for its phenotypic differentiation from F. canicola, which is a xylose-negative species (Korczak et al., 2014). We also designed a species-specific PCR that allows rapid and accurate identification and differentiation from F. canicola. In summary, new species may be added to the Frederiksenia genus in the future; in this report, we identified a possibly novel Frederiksenia-like organism by a combination of phenotypic and genetic analyses.

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