Immunoreactivity of the AAA+ chaperone ClpB from *Leptospira interrogans* with sera from *Leptospira*-infected animals

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**Abstract**

**Background:** *Leptospira interrogans* is a spirochaete responsible for leptospirosis in mammals. The molecular mechanisms of the *Leptospira* virulence remain mostly unknown. Recently, it has been demonstrated that *L. interrogans* ClpB (ClpB<sub>Li</sub>) is essential for bacterial survival under stressful conditions and also during infection. The aim of this study was to provide further insight into the role of ClpB in *L. interrogans* and answer the question whether ClpB<sub>Li</sub> as a potential virulence factor may be a target of the humoral immune response during leptosomal infections in mammals.

**Results:** ClpB<sub>Li</sub> consists of 860 amino acid residues with a predicted molecular mass of 96.3 kDa and shows multi-domain organization similar to that of the well-characterized ClpB from *Escherichia coli*. The amino acid sequence identity between ClpB<sub>Li</sub> and *E. coli* ClpB is 52%. The coding sequence of the clpB<sub>Li</sub> gene was cloned in *E. coli* BL21(DE3) strain. Immunoreactivity of the recombinant ClpB<sub>Li</sub> protein was assessed with the sera collected from *Leptospira*-infected animals and uninfected healthy controls. Western blotting and ELISA analysis demonstrated that ClpB<sub>Li</sub> activates the host immune system, as evidenced by an increased level of antibodies against ClpB<sub>Li</sub> in the sera from infected animals, as compared to the control group. Additionally, ClpB<sub>Li</sub> was found in kidney tissues of *Leptospira*-infected hamsters.

**Conclusions:** ClpB<sub>Li</sub> is both synthesized and immunogenic during the infectious process, further supporting its involvement in the pathogenicity of *Leptospira*. In addition, the immunological properties of ClpB<sub>Li</sub> point to its potential value as a diagnostic antigen for the detection of leptospirosis.

**Keywords:** ClpB, *Leptospira interrogans*, Leptospirosis, Molecular chaperone, Pathogen

**Background**

*Leptospira interrogans* belongs to pathogenic spirochaetes causing a serious disease in both humans and animals known as leptospirosis that is considered the most widespread zoonosis of worldwide importance [1]. The vectors of this pathogen are mostly wild rodents and domestic animals, which harbor the spirochetes in the proximal renal tubules of the kidneys and chronically excrete the leptospires with urine into the environment [2]. It is worth noting that leptospirosis is also a serious economic problem, because it causes abortions, stillbirths, infertility, failure to thrive, reduced milk production, and death in domestic animals such as cows, pigs, sheep, goats, horses and dogs [3–6]. In humans the disease varies from an asymptomatic flu-like illness to an acute life-threatening infection. Despite its severity and global importance, the molecular mechanisms of leptospiral pathogenesis remain largely unknown [1]. To date, only a few proteins have been identified as potential virulence factors in *Leptospira*. Among them, there is the chaperone ClpB, a member the Hsp100/Clp subfamily of the AAA+ ATPases that reactivates stress-aggregated proteins in cooperation with the DnaK system [7]. Recently, ClpB from *L. interrogans* (ClpB<sub>Li</sub>) has been shown to be essential for bacterial survival under stressful conditions (nutrient restriction, oxidative and heat stresses) and also for the pathogen’s virulence [8]. The involvement of ClpB in the response of *L. interrogans* to oxidative stress suggests that this chaperone may be...
one of key mediators of stress resistance, which is a prerequisite for Leptospira pathogenesis. The present study provides further insight into the role of ClpB\textsubscript{L1} during the infectious process. It is known that heat shock proteins (Hsps) play important roles during bacterial infections. They help pathogens to overcome stressful conditions to which they are exposed within the host cells, and represent major targets of the host’s immune system. Taking into account the fact that the chaperone ClpB from some pathogenic bacteria, Francisella tularensis and Mycoplasma pneumoniae, has been shown to be immunoreactive [9, 10], we decided to investigate an immunogenic potential of ClpB\textsubscript{L1}, which could point to this chaperone’s role in the pathogenicity of Leptospira and may translate into diagnostic applications.

**Methods**

**Serum samples**

We studied archived serum samples from rabbits and cattle. Rabbit antisera (n = 8) against L. interrogans serovars: Icterohaemorrhagiae, Hardjo, and Canicola, and L. borgpetersenii serovars: Hardjo, Javanica, were prepared as described by [11]. Polyclonal rabbit antisera prepared against the L. interrogans ClpB (residues 158–334; anti-ClpB\textsubscript{L1:158–334} serum) [8] and provided by M. Picard was used as a positive control and the pre-immune serum was used as a negative control. Bovine sera were collected from cattle (n = 10) experimentally infected with L. borgpetersenii serovar Hardjo via conjunctival instillation of 1 x 10⁶ bacteria. Blood samples were collected 28 days after the challenge and in one case 210 days after the challenge (this serum was used as a positive control showing the highest OD in ELISA). Sera from uninfected cattle (n = 8) and also a fetal bovine serum were used as negative controls. To confirm the serological status of leptospiro infection, the sera were subjected to the microscopic agglutination test (MAT) [11, 12] and used at dilutions 1:100 for Western blotting or 1:200 for ELISA.

**Kidney homogenate preparation**

For detection of ClpB\textsubscript{L1} in kidney tissues from Leptospiro-infected hamsters, the kidneys were macerated with nine parts of a 1 % BSA diluent and inoculated into Tween80/40/ LH semi-solid medium. Cultures were incubated at 28–30 °C, for up to 10 weeks and examined weekly by dark-field microscopy to detect the growth of leptospires. The same macerated kidney tissues (20 μg sample of homogenate) were used for Western-blotting analysis. Total protein concentration in the homogenates was determined by the method of Bradford [13].

**Plasmid construction for protein overproduction**

L. interrogans clpB gene (2583 bp) was amplified from genomic DNA of L. interrogans by PCR using AccuTaq LA polymerase MIX (Sigma) with the following primers: CATATGAAATTAGATAAAGCTTT with the NdeI restriction site underlined, and AAGCTTT TAACTACAACAAGTCC with the HindIII restriction site underlined. DNA primers were synthesized by Genomed S.A. (Warsaw, Poland). First, the PCR product was cloned into pJET1.2 blunt vector (Fermentas), then digested with NdeI, HindIII, and ligated with the linearized pET NdeI-HindIII vector. The sequence of the resulting construct was confirmed by DNA sequencing (Genomed S.A.). Leptospira genomic DNA was extracted with a QIAamp DNA Mini Kit (Qiagen).

DNA plasmid preparation and transformation of E. coli cells were done according to [14].

**Purification of the recombinant ClpB**

L. interrogans ClpB protein was overproduced in E. coli BL21(DE3) strain (Novagen) and purified according to the procedure similar to that used to obtain ClpB from Ehrlichia chaffeensis [15]. Briefly, bacteria were grown at 37 °C to OD\textsubscript{600} = 0.6 and then induced with 0.5 mM IPTG for 2 h. Next, the cells were collected and suspended in 50 mM Tris–HCl (pH 7.4), 300 mM NaCl, 20 mM imidazole and 0.1 % Triton X-100, then disrupted by sonication in the presence of the protease inhibitor PMSF and centrifuged to collect the soluble extract. Next, polyethyleneimine (PEI) was added to precipitate nucleic acids. After centrifugation (20 000 g, 1 h), the supernatant was applied to a Ni-NTA column (Qiagen) and the bound protein was eluted with 50 mM Tris–HCl (pH 7.4), 300 mM NaCl, and 0.1 % Triton X-100 and 250 mM imidazole. Fractions containing a 6His-tagged ClpB\textsubscript{L1} (a calculated molecular mass of 98 488,59 Da) were identified with SDS-PAGE electrophoresis and Coomassie blue staining, then combined and further purified by gel filtration on Superdex 200 (Sigma) equilibrated with 50 mM Tris–HCl (pH 7.5), 10 % glycerol, 1 mM EDTA and 1 mM DTT. The pooled fractions containing ClpB\textsubscript{L1} were dialyzed against dialysis buffer (50 mM Tris–HCl pH 7.5, 1 mM EDTA, 1 mM DTT, 20 mM MgCl\textsubscript{2}, 200 mM KCl, 10 % glycerol) and stored at −70 °C. The N-terminal histidine tag was removed by proteolytic digestion using the Thrombin Cleavage Capture Kit (Novagen) according to the manufacturer’s protocol. The identity of the purified ClpB\textsubscript{L1} was confirmed by a liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis of tryptic peptides obtained after trypsin cleavage of the protein, performed at the MS LAB IBB PAN (Warsaw, Poland). The equipment used was sponsored in part by the Centre for Preclinical Research and Technology (CePT), a project co-sponsored by European Regional Development Fund and Innovative Economy, The National Cohesion Strategy of Poland.
SDS-PAGE and Western blotting analysis
To assess immune reactivity of ClpB<sub>L</sub>, SDS-PAGE electrophoresis was performed according to [16] using 10 % polyacrylamide gels and Western blotting was performed as described [17]. The blots were blocked with 0.1 % Tween 20 in Tris-buffered saline (TBS) for 1 h at room temperature and then incubated overnight at 4 °C with anti-ClpB<sub>L158--334</sub> serum (1: 2000 dilution) [8] or polyclonal

### Fig. 1
Proposed domain organization of ClpB from L. interrogans.

**a** The diagram shows structural domains of the protein: N-terminal domain (ND) with the double Clp_N motif, nucleotide binding domain 1 (NBD1), middle coiled-coil domain (MD) and nucleotide binding domain 2 (NBD2). Conserved ATPase motifs such as the Walker A (A), Walker B (B), sensor 1, sensor 2 (GAR) and the Arg fingers (R), coordinating ATP binding and hydrolysis are also indicated. Conserved residues of these motifs are marked in bold.

**b** Sequence alignment of ClpB from E. coli (B<sub>Ec</sub>) and L. interrogans (B<sub>L</sub>). Domain boundaries are indicated below the amino acid sequence. The conserved motifs are shown in red. Identical and similar amino acid residues are shaded in black and gray, respectively.
rabbit and bovine sera (1:100 dilution) against Leptospira strains. After primary antibody incubation, the blots were washed three times with TBS containing 0.05 % Tween 20 and incubated for 1 h at room temperature with the goat anti-rabbit IgG horseradish peroxidase (HRP) conjugate (Sigma) diluted 1: 3000 or the polyclonal rabbit anti-cow IgG/HRP conjugate (DakoCytomation) diluted 1: 1000. The blots were then washed three times as described above and were developed using 3,3’-diaminobenzidine (Sigma), and H2O2 as substrates.

ELISA procedure

ELISA (enzyme-linked immunosorbent assay), was performed to analyze the immune response in animals experimentally exposed to L. interrogans serovars. Costar 96 well ELISA plates were coated with 100 µl of 0.625 µg/ml of the recombinant ClpBli (a capture antigen) resuspended in phosphate-buffered saline (PBS) by incubation overnight at 4 °C. The plates were then washed five times with PBS buffer (PBS containing 0.05 % Tween 20) and non-specific binding sites were blocked by incubation with 100 µl of 0.1 % Tween 20 in PBS buffer for 1 h at room temperature. The wells were washed five times with PBS buffer. Control and duplicate animal serum samples were diluted 200-fold in PBS buffer and 50 µl of the diluted sera (in duplicate) were applied to each well and incubated at 37 °C for 1 h, followed by five rinses with PBS buffer. Next, secondary HRP-conjugated anti-rabbit (Abcam) (diluted 1:10 000) or anti-cow IgG (DakoCytomation) (diluted 1:2000) were added to each well and incubated for 1 h at 37 °C. The plates were then washed five times with PBS buffer and 3,3’5,5’-tetramethylbenzidine (TMB) (Sigma) was added to detect the antibodies. The reaction was stopped after 10 min by the addition of 50 µl of 1 M H2SO4. The absorbance at 450 nm was measured using PerkinElmer Multimode Plate Reader (Enspire). The assay was performed three times for each serum.

Data analysis

The statistical significance of differences between the ELISA results obtained for sera collected from uninfected and infected animals were determined using the Welch’s adjusted one-way ANOVA followed by the post-hoc Scheffe multiple comparison test. P < 0.05 was considered statistically significant. Results of data analysis are presented in the graphs as the median values. All statistical analyses were performed using STATISTICA PL program.

Results

Analysis of the amino-acid sequence of the molecular chaperone ClpB from L. interrogans

The clpBli gene encodes a protein of 860 amino acid residues with a predicted molecular mass of 96325.2 Da. Sequence alignment of ClpBli (Fig. 1) revealed that this protein shows a multi-domain organization similar to that of the well-characterized ClpB from Escherichia coli (ClpB Ec). Thus, ClpBli contains an N-terminal domain (ND1-145aa), two nucleotide binding domains (NBD1 161-342aa, NBD2 560-768aa) and a middle coiled-coil domain (MD 393-527aa) (Fig. 1). Both NBDs, involved in ATP binding and ATP hydrolysis, contain all characteristic and conserved sequence motifs of AAA+ ATPases (ATPases associated with a variety of cellular activities), i.e. Walker A (GXGXXGKT/S), Walker B (HyxDE) and sensor 1/2 motifs. Conserved arginine residues called Arg fingers are also present in both NBD domains. Sequence alignment of the ClpB sequences from bacteria L. interrogans and E. coli using the Clustal software revealed that the total sequence identity between them is only 52 %; 27.7 % within ND, 45.3 % within MD, 72 % within NBD1, and 65.7 % within NBD2. Therefore, the most highly conserved are the NBD domains and the main differences between L. interrogans and E. coli ClpB are in the N-terminal domain and the coiled-coil middle domain.

Expression of the clpBli gene in E. coli cells and purification of ClpBli

To examine whether ClpBli shows an immunogenic potential, which could point to its participation in the pathogenicity of Leptospira, we obtained a construct expressing clpBli (pET28clpBli) and then overproduced the recombinant ClpBli as a 6-histidine-tagged protein in E. coli B21(DE3) cells. As expected, the expression of pET28clpBli resulted in the ~100-kDa protein,

![Fig. 2](image-url)
Protein sequence coverage: 88%

1. MKLDKLTSLKNEAIYNAQSAEKLGNPEISEEHILKEVLSQPDGLVPLLI
2. SKILNLSPKFLSTENALGQRQKVGGNHTSAQDVGFSSAVSLKKAADEVRLK
3. ELKDEYLDTHILLGLRMSQTSSLKTEFLQLQLEYKHLKILTENRGKTV
4. IMDSPEGKTDALAKYAKNLNEALKQGKLDQVIRDEERIRTIQVLSRRT
5. KNNLVILGEPGVYKTAIVEGLANKIVQGEVPGEKINTLYTLDLGMIG
6. AKYRGGEFDRLLKALDEVKSQDGVILFIEHITVVGGATEGALASNM
7. LIPMLARGELRCCIGATTLYEQKYIEKDAELRRFQVVYLKESEPSTEVT
8. ILARCLGKRYELEAHGIRILDSALIAAATLSITYSDRFLPMKVADIDEAS
9. SKRRIEIDSMPEELDRANKRIQSKLTERAELKKQDTSIKERKRTL
10. SEQEQNFQTILKRWDELSKIRGKLQIKEIEYKINLEAEAEARKGIEIRV
11. ABIRYGLVDLQKEESANSELQKQESASRLLKEVESEEIANTVSRRTG
12. IPVSKMLQGERAKLMLMDVLKTKVGGQDHALRLVSEAVQRSRAGIADPN
13. RPIGTFFLGTPTQGKTETAKALAEFLPDDVNMATIDSYMEYAAHVAR
14. LIGAPPVYVYDEGQGLTEAAYRRPSYLLEDFIEKANPEVFNIFLQLD
15. EGRRTDGKVRDVDFKNTVILITSHIGSEILGSSEYTSEKENRLVEORLKK
16. HPIPEFNLKIDVULIFHSITDSVSHKIAQLEGLRQAKKSLNENGVSFTN
17. ELKDYVSKAGFPAEFYGRPLKQRLIQRVEGNALSRYIHDGRFTNGQNVFTD
18. YRVOGRKVVVV

Fig. 3 LC-MS/MS analysis of the purified ClpB$_{L_i}$. The amino acid sequence of ClpB$_{L_i}$ is shown with the peptides detected by LC-MS/MS indicated in red.

Fig. 4 Immune reactivity of the recombinant ClpB$_{L_i}$ with rabbit sera. a The purified ClpB$_{L_i}$ protein (250 ng) was resolved by SDS-PAGE and analyzed by Western blotting using: the antisera against ClpB$_{L_i}$ (a positive control), pre-immune control serum (a negative control), or polyclonal rabbit antisera raised against: L. interrogans and L. borgpetersenii serovars as indicated in the figure. The positions of protein size markers (M) (in kDa), PageRuler prestained Protein Ladder (Thermo Scientific), are shown on the left. The arrow indicates the position of ClpB$_{L_i}$. (b) ELISA analysis of the recombinant ClpB$_{L_i}$ protein as a capture antigen using all the above rabbit sera. The data were analyzed using Welch adjusted one-way ANOVA. Symbols: (▫), the median value; (box), 25 %–75 % range around the median value; (whiskers), min-max range. (****) denotes $P < 0.001$; ns, not statistically significant.
corresponding to ClpB_{Li} that was soluble in *E. coli* cells. The protein was purified from the soluble fraction using two separation techniques: immobilized metal affinity chromatography (IMAC) and gel filtration chromatography (Fig. 2a). The identity of ClpB_{Li} was confirmed with an LC-MS/MS analysis (Fig. 3). The obtained peptide map covered 88 % of the amino acid sequence of ClpB_{Li}. In addition, LC-MS/MS data indicated that the purified ClpB_{Li} was not contaminated with ClpB from the *E. coli* host strain. The purified ClpB_{Li} was subsequently digested with thrombin to remove the N-terminal 6His-tag (Fig. 2b). The post-cleavage N-terminal sequence of the recombinant ClpB_{Li} protein contains three additional amino acid residues, namely GlySerHis, and in such form the protein was further characterized by Western blotting analysis and ELISA assay.

### Immunogenic capacity of ClpB_{Li}

The immune reactivity of ClpB_{Li} with serologically positive sera from rabbits and cattle experimentally infected with two pathogenic *Leptospira* species (*L. interrogans* and *L. borgpetersenii*) was tested by Western blotting (Figs. 4a and 5a) and ELISA assay (Figs. 4b and 5b) and compared to the sera from uninfected healthy controls. We found that all the tested sera prepared from *Leptospira*-infected animals, but not from the uninfected controls,

![Fig. 5](image-url)

**Fig. 5** Immune reactivity of the recombinant ClpB_{Li} with bovine sera. a The purified ClpB_{Li} protein (250 ng) was resolved by SDS-PAGE and analyzed by Western blotting using the antiserum against ClpB_{Li}, 158–334 (a positive control; control +), polyclonal bovine antisera raised against: *L. borgpetersenii* serovar Hardjo (*Leptospira*-infected cattle), and sera collected from uninfected cattle (healthy group; negative control). The positions of protein size markers (M) (in kDa), PageRuler prestained Protein (Thermo Scientific), are shown on the left. The arrow indicates the position of ClpB_{Li} (~100 kDa). b ELISA analysis of the recombinant ClpB_{Li} protein as a capture antigen using the above bovine sera. Fetal bovine serum was also used. The data were analyzed using Welch adjusted one-way ANOVA. Symbols: (∗∗∗), the median value; (box), 25 %–75 % range around the median value; (whiskers), min-max range. (∗∗∗) denotes *P* < 0.001
Detection of ClpB<sub>Li</sub> in *Leptospira*-infected animals

Additionally, we detected ClpB<sub>Li</sub> (96-kDa protein) in the infected hamster kidney tissue (Fig. 6), from which leptospires were isolated using standard culture method. No reactivity of the 96-kDa protein with anti-ClpB<sub>Li</sub> 1158–334 serum was observed in the kidney homogenate obtained from an uninfected hamster (Fig. 6, lane 4). The result indicates that ClpB<sub>Li</sub> is produced during an experimental infection of animals.

In summary, our data indicate that the molecular chaperone ClpB<sub>Li</sub> is immunogenic and detectable in animals infected with pathogenic *Leptospira* spp.

Discussion

*Leptospira* like many other pathogenic bacteria are exposed to a significant stress during infection of host cells, frequently resulting in protein misfolding and aggregation. Despite being exposed to stressful conditions, pathogens survive, overcome host defense mechanisms, and cause the disease symptoms. The specific mechanisms of the host invasion by leptospires are not well defined. In particular, the molecular basis for virulence remains unknown, due to the lack of genetic tools for the manipulation of *Leptospira*. The fact that ClpB is usually up-regulated in pathogenic microorganisms [8, 15] suggests that the disaggregase activity of ClpB may be essential for their virulence. Moreover, the involvement of ClpB in the response of *L. interrogans* to oxidative stress [8] suggests that this chaperone may be one of key mediators of stress resistance, which is a prerequisite for *Leptospira* pathogenesis. The chaperone ClpB may function either as a true virulence factor directly involved in causing the disease or a virulence-associated protein that can be essential for colonization of the host. Virulence gene products are often immunogenic and responsible for acquired immunity that protects against disease [18]. At this point it should be also noted that molecular chaperones despite their cytosolic localization are strongly immunogenic in a number of bacterial infections [19]. It has been reported that some chaperones (e.g. GroEL) may be associated with the outer membrane of the pathogenic bacteria or exported from the bacterial cell after heat shock [19]. Therefore, exposure of bacterial Hsps to the host's immune system is possible during infection. Indeed, ClpB from some pathogens (i.e. *Mycoplasma pneumoniae, Francisella tularensis*) is an immunoreactive protein [9, 10]. The total sequence identity between ClpB proteins from these pathogens is only ~40%. It is likely that ClpB as an important mediator of resistance to oxidative stress could be also a potential target for the host immune response during leptospiral infections in mammals. Therefore, we decided to investigate an immunogenic potential of this chaperone in *Leptospira*. The use of the *E. coli* expression system allowed us to produce the recombinant ClpB<sub>Li</sub> protein and to assess its immune reactivity with sera collected from *Leptospira*-infected animals and the uninfected healthy controls. Our results show that ClpB is immunogenic during leptospiral infections because it was recognized by sera collected from experimentally infected animals (see Figs. 4 and 5). Thus, among the antibodies raised against leptospiral proteins, there were specific antibodies against ClpB<sub>Li</sub>. This is the first study where ClpB from pathogenic *Leptospira* species was evaluated for its ability to elicit immune responses in animals. Moreover, our results suggest that ClpB<sub>Li</sub> could be considered as a potential antigen candidate for a diagnostic test. We postulate that the presence of species-specific domains (e.g. ND or MD, see Fig. 1) in the antigen could minimize a cross-reactivity of antibodies with ClpB from different bacteria. Further prospective studies are needed to assess the ClpB<sub>Li</sub>’s predictive value in leptospirosis diagnostics. In addition, the presence of ClpB<sub>Li</sub> in the infected hamster kidney tissues (see Fig. 6) demonstrates that the chaperone is produced by pathogen during infection of the host further confirming the involvement of ClpB in the pathogenicity of *Leptospira*.

Conclusions

Identification of *Leptospira* virulence factors and understanding their properties is crucial for uncovering the diseases mechanisms. This study underlines the potential
importance of the chaperone ClpB in leptospiral infections. We believe that our data provide new information, which may lead to a better understanding of the role of ClpB and possibly other stress-response factors in the life cycle of the pathogenic bacterium L. interrogans. It is worth noting that since ClpB does not exist in animal cells, it might become a promising target for novel therapies against pathogenic Leptospira species. Further studies are needed to determine the biological role of ClpB during leptospiral infection in mammals and its diagnostic or even immunoprotective potential. The recombinant ClpB12 produced in this work will help in further biochemical characterization of this chaperone and the analysis of its function in the pathogen.

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Availability of supporting data
The DNA sequence of L. interrogans clpB gene was retrieved from the EMBL-EBI (accession number AAC18188.1) website (http://www.ebi.ac.uk/ena/data/view/AC18188). The protein sequences of L. interrogans and E. coli ClpB were retrieved from UniProtKB (accession number Q72QU2 and P63284 (CLPB_ECOLI)/http://www.uniprot.org/uniprot/P63284).

Authors’ contributions
JK, ZA, DW performed the experiments. SKM designed the experiments, analyzed the data and drafted the manuscript. MZ assisted in data analyses and the preparation of manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
The sera and kidney tissues used in this work were originally collected during another study (project license number PPL2608, date of approval 15 October 2008). All operators involved in the study, protocols, and premises were licensed under the Animals (Scientific Procedures) Act (1986) (ASPA).

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