

**MOLECULAR AND GENETIC CHARACTERIZATION OF *LEPTOSPIRA*
SPP. COLLECTION STRAINS FROM THE ST. PETERSBURG PASTEUR
INSTITUTE BASED ON 16S rRNA GENE SEQUENCING DATA**

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LEPTOSPIRA SPP. 16S rRNA ANALYSIS

АНАЛИЗ ЛЕПТОСПИР ПО 16S РРНК

МОЛЕКУЛЯРНО-ГЕНЕТИЧЕСКАЯ

КОЛЛЕКЦИОННЫХ ШТАММОВ *LEPTOSPIRA* SPP. САНКТ-
ПЕТЕРБУРГСКОГО ИНСТИТУТА ПАСТЕРА НА ОСНОВЕ ДАННЫХ
СЕКВЕНИРОВАНИЯ ГЕНА 16S РРНК

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ХАРАКТЕРИСТИКА

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Abstract.

Leptospirosis is a zoonotic disease found virtually worldwide. Microscopic Agglutination Test with live *leptospira* (MAT) is the reference method for the serological diagnosis of leptospirosis. MAT is based on assessing serum potential to agglutinate live reference serovar *Leptospira* maintained at a reference laboratory. At some laboratories having own collections of isolated and reference *Leptospira* strains applicable for serological diagnosis, those microorganisms are maintained for many years by repeated subculturing, that increases markedly a chance of strain cross-contamination. The lack of adequate quality control for reference strains may affect data of epidemiological studies. Control of *Leptospira spp.* reference strains purity and stability of their antigenic composition is very important for diagnosis of leptospirosis. The study objective was to compare the 16S rRNA gene nucleotide sequences of some *Leptospira* strains from the collection of the St. Petersburg Pasteur Institute to with relevant sequences uploaded to GenBank. In this study, 38 *Leptospira* strains were investigated. Nucleotide sequences of 36 strains were deposited in the international GenBank database, inconsistencies were revealed in two strains. The study found that the control *Leptospira* strains from the collection of the St. Petersburg Pasteur Institute had minimal dissimilarities from international control strains. The analysis of the resultant 16S rRNA sequences has shown the presence of point mutations, transitions, deletions and insertions, regardless of the strain species. The open *leptospira* pan-genome demonstrates high genomic variability in species due to the capability of *leptospira* for lateral gene transfer in order to adapt to changing environmental conditions. The massive acquisition and loss of genes give rise to an increased species diversity. The 16S rRNA gene is suitable for screening diagnostics; however, high level of the fragment similarity and close phylogenetic relationship between different species put bounds to its use in genotyping. The presence of point nucleotide mutations is most likely associated with the evolutionary mechanisms of *leptospira*, their ability to horizontal gene transfer and crossing-over, including ribosomal genes, but this assumption

necessitates additional research. For specimen genotyping it is necessary to select alternative genes with high specificity and sufficient level of nucleotide divergence. The study shows a need for genetic analysis of collection strains in order to control the purity of cultures.

Keywords: leptospirosis; leptospira; 16S rRNA; collection; zoonotic disease; cultivating.

Резюме.

Лептоспироз является распространенным практически по всему миру зооантропонозным заболеванием. Эталонным методом для серологической диагностики лептоспироза является реакция микроскопической агглютинации с живыми лептоспиралами (РМА). Этот метод основан на оценке способности сыворотки агглютинировать живые бактерии *Leptospira* эталонных сероваров, поддерживаемых в референс-лаборатории. В лабораториях, которые имеют коллекцию собственных изолированных и референсных штаммов *Leptospira*, используемых для серологической диагностики, эти микроорганизмы сохраняются в течение многих лет путем повторного пассирования, что значительно увеличивает шансы кросс-контаминации штаммов. Отсутствие адекватного контроля качества референсных штаммов, может отрицательно повлиять на эпидемиологические исследования. Контроль чистоты и постоянства антигенного состава референсных штаммов *Leptospira spp.* имеет большое значение для диагностики лептоспироза. Целью данного исследования было сравнение нуклеотидных последовательностей гена 16S, некоторых коллекционных штаммов лептоспир СПбИП им. Пастера, с последовательностями, загруженными в международную базу данных. В работе были изучены 38 штаммов лептоспир. Нуклеотидные последовательности 36 штаммов были депонированы в международную базу данных GenBank, в двух штаммах были обнаружены несоответствия. В результате исследования установлено, что контрольные штаммы *Leptospira* из коллекции Санкт-Петербургского института Пастера имеют минимальные отличия от международных контрольных штаммов. Анализ полученных последовательностей региона 16S рРНК, показал наличие точечных мутаций, транзиций, делеций и инсерций, независимо от видовой принадлежности штамма. Открытый пангеном лептоспир демонстрирует высокую геномную вариабельность у видов, что обусловлено способностью лептоспир к латеральному переносу генов, с целью приспособления к

изменяющимся условиям среды. Массовое приобретение и потеря генов ведут к увеличению разнообразия видов. Ген 16S рРНК подходит для скрининговой диагностики, однако высокое сходство данного фрагмента и тесное филогенетическое родство разных видов ограничивает его использование для генотипирования. Наличие точечных нуклеотидных мутаций вероятнее всего связано с эволюционными механизмами лептоспир, их способностью к горизонтальному переносу генов и кроссинговеру, в том числе и рибосомальных генов, однако это предположение обуславливает необходимость проведения дополнительных исследований. Для генотипирования образцов необходим подбор альтернативных генов, с высокой специфичностью и достаточным уровнем дивергенции нуклеотидов. Проведенное исследование показывает необходимость проведения генетического анализа коллекционных штаммов, с целью контроля чистоты культур.

Ключевые слова: лептоспироз; лептоспира; 16S рРНК; коллекции; зоонозные заболевания; культивирование.

1 Introduction.

Leptospirosis is a zoonotic disease occurring worldwide. The manifestations of its epidemic process are most typical for countries with humid, especially tropical and subtropical climates, as well as for the so-called developing countries, however, the disease cases occur as well in countries with temperate climates [21, 43]. Leptospirosis is endemic in tropical regions of Asia, Africa, Central and South America [45]. Numerous animal species including those synanthropic and farm livestock are considered as reservoir hosts of pathogenic *Leptospira*.

The epidemic state of leptospirosis varies significantly in European countries. In recent years, in non-endemic countries, there was an increase in imported leptospirosis due to the rise in popularity of ecotourism in regions with humid subtropical or tropical climates [1]. In Russia leptospirosis remains one of widespread zoonotic infections due to the existence of both natural and anthropurgic foci of leptospirosis in many administrative regions [2-6]. Some assessments come to more than 1 million severe cases of leptospirosis and about 60,000 fatal outcomes occur annually worldwide [16].

According to the serological classification, all currently recognized species of pathogenic *Leptospira* are categorized into more than 260 serovars [10]. Based on the phylogenetic classification, the genus *Leptospira* is divided into 64 species. In line with genome-wide analysis, the genus is divided into 2 clades: pathogens and saprophytes, which, in turn, are divided into 4 subclades (P1, P2, S1, S2). Subclade P1 and P2 involve 17 pathogenic and 21 intermediate species, respectively. Subclade S1 includes 22 saprophytic species, while subclade S2 includes 4 [37].

Leptospirosis in humans is mostly caused by *Leptospira interrogans*, *Leptospira borgpetersenii* or *Leptospira kirschneri* [9, 26, 35]. The pathogen can circulate in the environment for a long time thanks to sensitive or reservoir hosts that shed bacteria in their urine, contaminating soil and water. Humans usually get infected through direct contacts of their damaged skin or mucous membranes with the urine of infected wild or domestic animals, or through indirect contacts with

30 contaminated environment [8, 29, 36]. In humans, there is a variety of possible
31 clinical manifestations of the disease, ranging from subclinical infection to severe
32 progressive forms that can lead to death [19, 29, 39].

33 Laboratory confirmation of suspected cases is essential for the diagnosis of
34 leptospirosis, as there is significant polymorphism of its clinical manifestations.
35 Most cases are diagnosed through serological tests of patients' blood. Microscopic
36 Agglutination Test with live *Leptospira* (MAT) is the reference method for the
37 serological diagnosis of leptospirosis. MAT is based on the assessment of the serum
38 ability to agglutinate live *Leptospira* of reference serovars maintained at a reference
39 laboratory [22, 32, 42, 44]. At some laboratories that have their own collections of
40 isolated and reference *Leptospira* strains applicable for serological diagnosis, those
41 microorganisms are maintained for many years by repeated subculturing, that
42 increases significantly the chance of strain cross-contamination [13, 14, 19, 26].

43 Contamination of strains in *Leptospira* reference collections may entail severe
44 consequences. In an outbreak investigation, the pathogen serogroup is usually
45 identified with the help of MAT. The lack of adequate quality control of reference
46 strains may affect the results of epidemiological studies. Conventional serogroup
47 control of reference strains is carried out using monoclonal antibodies or control sera
48 samples [7, 28, 46].

49 Control of *Leptospira* spp. reference strains purity and stability of its antigenic
50 composition is very important for the diagnosis of leptospirosis. Sequencing of 16S
51 rRNA gene is a reliable method for molecular characterization of bacteria species,
52 and it is applicable to *Leptospira* spp. [15, 25, 31, 38]. The method was considered
53 to be an effective and simple tool for *Leptospira* species identification in the clinical
54 setting. Its important advantages are fast analysis, wide availability, and relatively
55 low cost. The study objective was to compare the 16S rRNA gene nucleotide
56 sequences of some collection *Leptospira* strains, including reference strains, with
57 sequences uploaded to GenBank.

58 2 Materials and Methods.

59 In this study, we investigated 38 *Leptospira* strains from the collection of the
60 Laboratory of Zoonotic Infections at the St. Petersburg Pasteur Institute.

61 The cultures were characterized by serological tests using MAT [18]. The
62 strains have undergone repeated subculturing at the laboratory for many decades.
63 Cultivation was carried out in a liquid nutrient medium based on distilled water
64 containing 10% phosphate buffer with the addition of 10% rabbit blood serum. The
65 strains were cultivated at 29° C. We used 7- to 14-days grown *Leptospira* cultures,
66 containing at least 10⁷ microbial cells/ml. Genomic DNA was extracted using a
67 QiaAmp DNA Mini kit (Qiagen, Germany) in compliance with the manufacturer's
68 instruction. For PCR, two pairs of primers flanking a 1423 bp fragment were used.
69 Primer sequence: 16S Out F 5'-AGAGTTGATCCTGGCTCAG- 3', 16S Out R 5'-
70 GGYTACCTTGTTACGACTT- 3' [28, 33]. A clinical sample was used for a
71 positive amplification control. The presence of *Leptospira* DNA in it was confirmed
72 by two commercial kits: AmpliSens® *Leptospira*-FRT (InterLabService Ltd.,
73 Russia), and BactoReal® Kit *Leptospira* spp. Multiplex (16S rDNA+LipL32)
74 (Ingenetix GmbH, Austria). ClearBand Nuclease Free Ultra-Pure Water (EcoTech
75 Biotechnology Turkey) was used for a negative control.

76 PCR amplification was carried out in 25 µl of the amplification mixture with
77 the addition of primers (15 pM each) 67 mM plus Tris HCl (pH 8.8), 16.6 mM
78 ammonium sulfate, 6.7 mM MgCl₂; 6.7 mM EDTA; 10 mM mercaptoethanol; 170
79 mg BCAA; 1.0 mM each dNTP; 1 unit Taq DNA polymerase (Fermentas).
80 Denaturation (5 min at 94°C,) was followed by 40 cycles of amplification: 30 s at
81 94°C, 30 s at 55°C, 1 min 20 s at 72°C, final elongation 7 min at 72°C.

82 The PCR products were separated on 2% agarose gel stained with ethidium
83 bromide, in comparison with the GeneRuler 1Kb molecular weight marker (Thermo
84 Scientific, USA). Electrophoresis was conducted for 40 minutes at 120V and
85 visualized by UV radiation.

86 The amplification products were purified according to the procedure as
87 follows: the mixture (including 2 µl of 3 M sodium acetate and 2 µl of 0.125 M

88 EDTA, and 1 μ l of glycogen) was added to 20 μ l of the amplification product and
89 incubated for 15 minutes at room temperature in the presence of chilled 96% ethanol.
90 Centrifugation was carried out for 15 minutes at 4 $^{\circ}$ C, 14,000 rpm/min. Then
91 supernatant was removed and the precipitate was washed twice with cold 70%
92 ethanol, repeating the cold centrifugation procedure. The washed precipitate was
93 dried in the air at room temperature. For the quality analysis, the purified precipitate
94 was dissolved in 30 μ l of TE buffer and visualized on an agarose gel. The purified
95 fragment of sufficient concentration was used to set up a sequence of reactions with
96 forward and reverse primers. For analysis, the reaction product was dissolved in
97 formamide and placed in an ABI Prism 3500 genetic analyzer (Applied Biosystems,
98 USA).

99 The resulting strain sequences were compared with those available from the
100 GenBank database.

101 For phylogenetic analysis, the MEGA11 software was used, the resulting
102 sequences were aligned using the ClustalW algorithm. The sequence coverage
103 length was 1144 bp. When constructing the tree, the sequences of 16S region of
104 *Borrelia* spp. from the international GenBank database, were chosen to be used for
105 the outgroup. Based on the alignment, a tree was built using the Neighbor-joining
106 method, bootstrap N = 1000.

107 3 Results.

108 Nucleotide sequences of the 16S rRNA gene of 38 strains were obtained. The
109 length of the sequences ranged from 1186 to 1423 bp. The sequences of 36 strains
110 were deposited in the international GenBank database. For some details, such as
111 serogroup of strain, date of isolation, origin and location see Table 1. The overall
112 sequence similarity is 99%, the number of polyform variants of the gene among the
113 obtained sequences ranges from 1 to 20 nt, which is consistent with the results of
114 previous studies [34].

115 Strain sequences were identified and validated using NCBI BLAST and by
116 alignment in MEGA 11 (Fig. 1).

117 The sequence of the *L. borgpetersenii* strain Perepelicin, obtained at our
118 laboratory, forms a cluster with foreign reference strains, although it differs by one
119 nucleotide.

120 Reference strains of *L. kirschneri* and *L. Interrogans* species, obtained by
121 foreign researchers, are clustered together, indicating their high similarity and
122 inapplicability of 16S rRNA gene to typing.

123 The results of comparison with nucleotide sequences of the 16S rRNA gene,
124 obtained by other researchers, were mostly concordant, although some discrepancies
125 were observed.

126 The analysis of the resultant sequences of 16S rRNA region has shown the
127 presence of point mutations, transitions, deletions and insertions, regardless of the
128 strain species.

129 Three points with a variable nucleotide were found in 10 *L. interrogans*
130 sequences obtained by us. In the sequences of GenBank strains thymine is more
131 common at position 38, however, there are some strains with cytosine. The reverse
132 is true for the strains here studied: only 8 of 36 strains contain thymine at this point,
133 while the rest of them contain cytosine (Fig. 2).

134 The sequence of the *L. kirschneri* strain is 99.0% similar to that of the *L.*
135 *interrogans* species, but 99.72% sequence identity was found by NCBI BLAST
136 analysis.

137 In two strains out of 38, inconsistencies were revealed. The sequences of the
138 collection strains of the species *L. borgpetersenii* and *L. noguchii* were analyzed and
139 proved to be identical to the strains belonging to *L. interrogans*.

140 4 Discussion.

141 Historically, *Leptospira* strains were classified according to their virulence,
142 all saprophytic strains were assigned to *L. biflexa*, and pathogenic strains to *L.*
143 *interrogans* [41]. Further subdivision of *Leptospira*, taking into account their
144 antigenic determinants, was based on serological techniques [28].

145 *Leptospira* classification was significantly expanded due to the introduction
146 of such genetic techniques as DNA-DNA hybridization, pulsed-field gel
147 electrophoresis (PFGE), and Multilocus sequence typing [11, 13, 20]. Serological
148 characteristics do not correlate with genovariants, one species of *Leptospira* may
149 contain several serogroups, while one serogroup may include strains of different
150 species. The open pangenome of *Leptospira* demonstrates high genomic variability
151 in species due to the capability of *Leptospira* for lateral gene transfer in order to
152 adapt to changing environmental conditions. The massive acquisition and loss of
153 genes give rise to an increase in species diversity [24, 36].

154 Sequencing of the 16S rRNA region is currently the most accessible method
155 for genotyping. This gene provides distinction between pathogenic, intermediate,
156 and saprophytic clades, but its discriminatory ability is too low to distinguish
157 between *Leptospira* genotypes. For example, it is impossible to differentiate *L.*
158 *interrogans*, *L. kirschnerii*, and *L. noguchii*, since they differ by one or two bases
159 only and belong to the same operational taxonomic unit (mOTU) that is shown by
160 the phylogenetic tree (Fig. 1) [23]. The sequence of the *L.kirschneri* strain obtained
161 in our study also has a minimal difference only from the type strains of
162 *L.interrogans*.

163 A large number of 16S rRNA sequences uploaded to GenBank are not full-
164 length (<1400 bp), therefore it is difficult to identify strains correctly. Long-term
165 subculturing of *Leptospira* in collections does not exclude erroneous identification
166 of sequences uploaded to public databases, it also can give rise to single mutations
167 [12].

168 When sequencing this gene, it was found that the control strains of *Leptospira*
169 from the collection of the St. Petersburg Pasteur Institute have minimal
170 dissimilarities from international control strains and are applicable in serological
171 diagnosis of leptospirosis.

172 In this study, two out of 38 strains did not match the sequences of similar
173 strains uploaded to GenBank, which may be due to its contamination or mislabeling.

174 Long-term storage of *Leptospira* by freezing is not available at all practical
175 laboratories owing to its high price, therefore collection strains of *Leptospira* for
176 many years undergo repeated subculturing, possibly resulting in strain mutations and
177 increasing the risk of cross-contamination or erroneous labeling of samples. This
178 justifies the need for genetic analysis of collections in order to circumvent the
179 possible risks associated with under- or overdiagnosis.

180 The 16S rRNA gene is suitable for screening diagnostics; however, high level
181 of the fragment similarity and close phylogenetic relationship between different
182 species put bounds to its use in genotyping.

183 The presence of point nucleotide mutations is most likely associated with the
184 evolutionary mechanisms of *Leptospira*, their ability to horizontal gene transfer and
185 crossingover, including ribosomal genes, but this assumption necessitates additional
186 research [34].

187 For genotyping of samples, it is necessary to select alternative genes with
188 high specificity and a sufficient level of nucleotide divergence.

189 Nowadays numerous genetic targets have been proposed for *Leptospira*
190 diagnosis and typing. The gene encoding the outer membrane lipoprotein of
191 pathogenic *Leptospira* *lipL32* is one of the most common alternatives to 16S rRNA,
192 however, its use is also limited by its discriminating abilities, and it is mostly applied
193 in screening diagnostics.

194 The housekeeping gene encoding the *secY* preprotein translocase is next in
195 frequency of use; the use of this gene for *Leptospira* genotyping seems to be the
196 most promising at the moment. Other proposed targets are: the gene encoding
197 flagellin, the class B polypeptide subunit of periplasmic flagella *flaB*, the gene for
198 the β-subunit of RNA polymerase *rpoB*, and the genes *gyrB*, *Adk*, *glmU*, *LigB*, etc.
199 [17, 27, 30, 40].

200 As a variety of targets is used by many researchers, the comparison of their
201 results is hardly possible. Therefore, the lack of standardization is currently the main
202 restriction on the molecular diagnosis of leptospirosis.

203 This study shows the need for genetic analysis of collection strains in order to
204 control the purity of cultures.

ТАБЛИЦЫ

Table 1. *Leptospira* strain sequences deposited in GenBank.

№ of strain sequence deposited in GenBank	Serogroup	Date of isolation	Origin	Location
OL703051	Canicola	1985	Human	Leningrad
OL703052	Icterohaemorrhagiae	1999	Human	Saint Petersburg
OL703053	Canicola	1962	Human	Kaliningrad
OL703054	Canicola	1969	Human	Kaliningrad
OL703055	Icterohaemorrhagiae,	1992	Human	Leningrad
OL703056	Icterohaemorrhagiae	1991	Human	Leningrad
OL703057	Icterohaemorrhagiae	1992	Human	Leningrad
OL703058	Canicola	2008	Human	Saint Petersburg
OL703059	Icterohaemorrhagiae	1968	Human	Leningrad
OL703060	Canicola	1968	Human	Kaliningrad
OL703061	Canicola	1985	Human	Leningrad
OL703062	Icterohaemorrhagiae	1990	Human	Leningrad
OL703063	Icterohaemorrhagiae	1990	Human	Leningrad
OL703064	Autumnalis	1925	Human	Japan
OL703065	Australis	1951	<i>Erinaceus europaeus</i>	Moscow
OL703068	Icterohaemorrhagiae	2018	Human	Saint Petersburg
OL703069	Icterohaemorrhagiae	1990	Human	Leningrad

OL825730	Icterohaemorrhagiae	2009	Human	Saint Petersburg
OL825731	Icterohaemorrhagiae	1980	<i>Rattus norvegicus</i>	Leningrad
OL825732	Icterohaemorrhagiae	1954	<i>Rattus norvegicus</i>	Leningrad
OL825733	Icterohaemorrhagiae	1961	<i>Rattus norvegicus</i>	Leningrad
OL825734	Icterohaemorrhagiae	1991	Human	Leningrad
OL825735	Icterohaemorrhagiae	1997	Human	Saint Petersburg
OL825736	Icterohaemorrhagiae	1990	Human	Leningrad
OL825737	Icterohaemorrhagiae	1991	Human	Leningrad
KR080516	Canicola	1931	<i>Canis lupus familiaris</i>	Holland
KY075908	Tarassovi	1938	Human	Moscow
KY075909	Pyrogenes	1923	Human	Indonesia
KY075910	Javanica	1938	<i>Rattus rattus</i>	Indonesia
KY075911	Australis	1937	Human	Australia
KY075912	Bataviae	1938	Human	Indonesia
KY075913	Autumnalis	1938	Human	Congo
KR107202	Sejroe	1944	<i>Apodemus flavicollis</i>	Denmark
KR091970	Icterohaemorrhagiae	1935	Human	Denmark
KR091971	Pomona	1936	Human	Australia
KR091973	Grippotyphosa	1929	Human	Moscow

РИСУНКИ

Figure 1. Phylogenetic tree built on the obtained *Leptospira* spp 16S rRNA sequences and sequences of reference strains taken from GenBank.

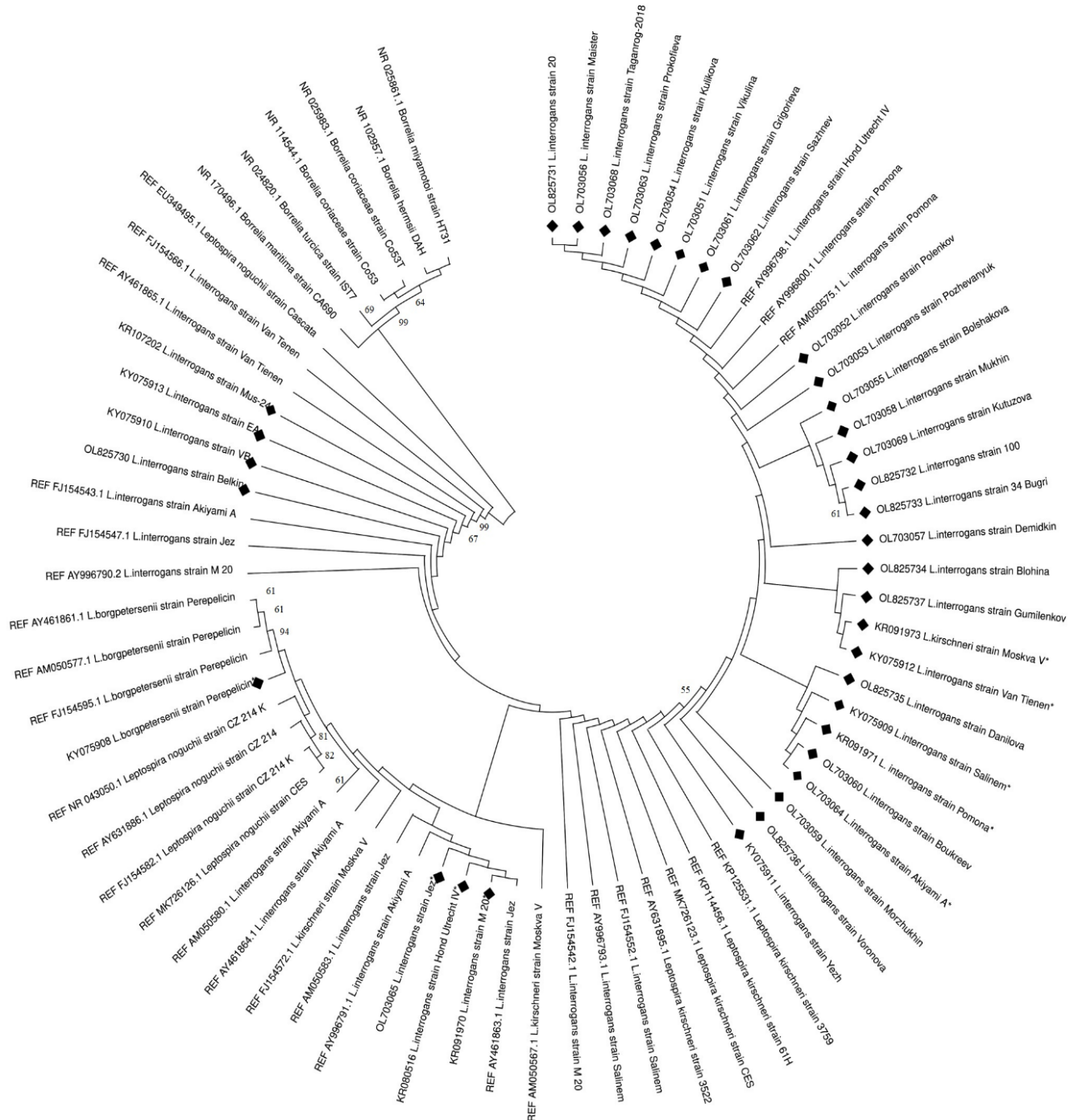
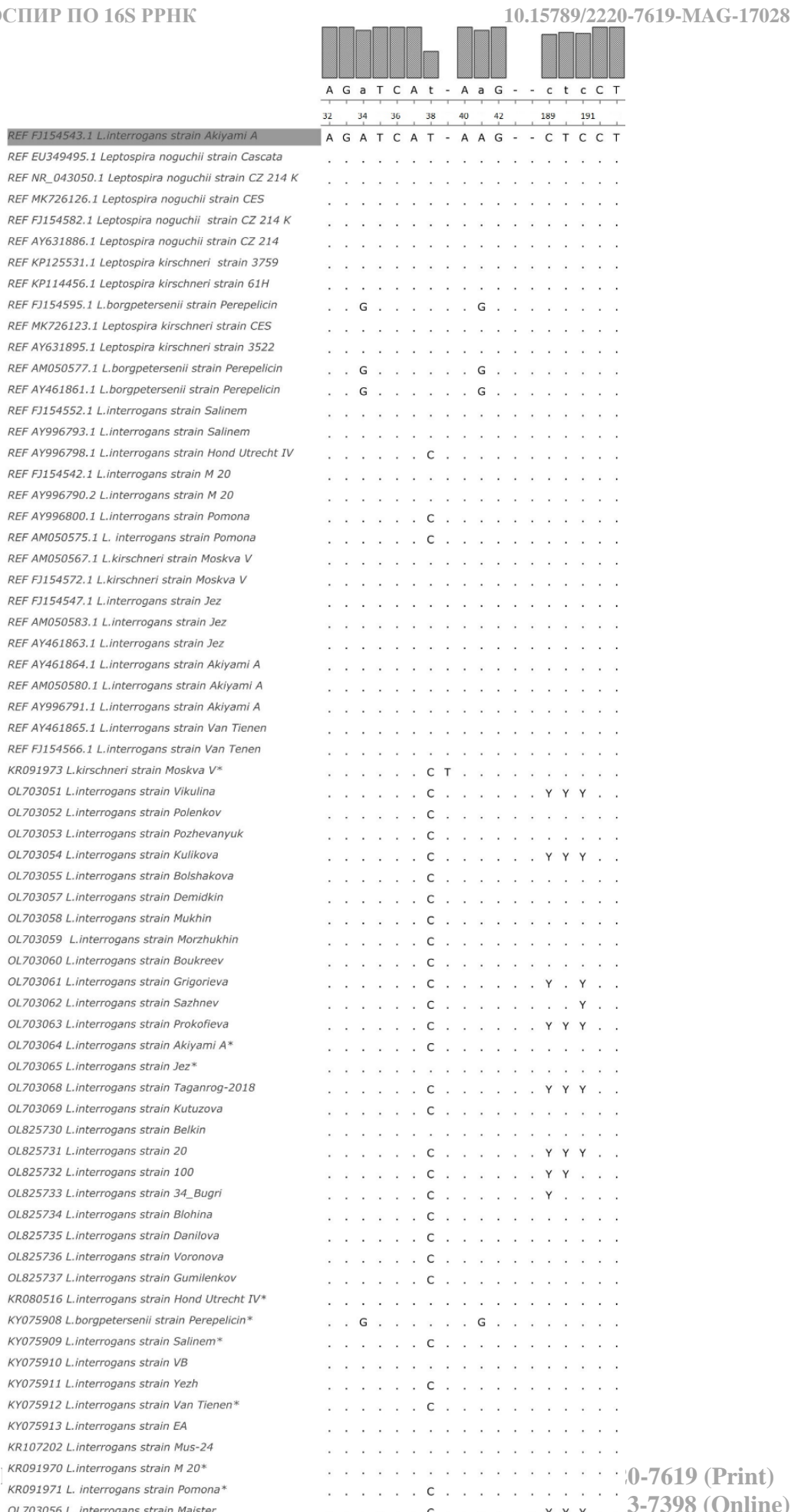


Figure 2. Multiple alignment of the obtained sequences with reference strains obtained from the international GenBank database.

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Notes: REF - sequences of reference strains from GenBank. *- sequences of reference strains obtained at our laboratory. The studied strains are described in Table 1;

The main differences are shown.

ТИТУЛЬНЫЙ ЛИСТ_МЕТАДАННЫЕ**Блок 1. Информация об авторе ответственном за переписку**

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Блок 3. Метаданные статьи

MOLECULAR AND GENETIC CHARACTERIZATION OF *LEPTOSPIRA* spp.
COLLECTION STRAINS FROM THE ST. PETERSBURG PASTEUR
INSTITUTE BASED ON 16S rRNA GENE SEQUENCING DATA

МОЛЕКУЛЯРНО-ГЕНЕТИЧЕСКАЯ

ХАРАКТЕРИСТИКА

КОЛЛЕКЦИОННЫХ ШТАММОВ *LEPTOSPIRA* spp. САНКТ-
ПЕТЕРБУРГСКОГО ИНСТИТУТА ПАСТЕРА НА ОСНОВЕ ДАННЫХ
СЕКВЕНИРОВАНИЯ ГЕНА 16S РРНК

Keywords: leptospirosis; leptospira; 16S rRNA; collection; zoonotic disease; cultivating.

Ключевые слова: лептоспироз; лептоспира; 16S рРНК; коллекции; зоонозные заболевания; культивирование.

Оригинальные статьи.

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СПИСОК ЛИТЕРАТУРЫ

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1.	Киселева Е.Ю., Бренева Н.В., Лемешевская М.В., Бурданова Т.М. Завозной случай лептоспироза с летальным исходом из Вьетнама в Иркутскую область // Инфекционные болезни 2014 - . 12. - № 3. - С. 95 - 100	Kiseleva E.Yu., Breneva N.V., Lemeshevskaya M.V., Burdanova T.M. (2014). An imported case of leptospirosis with a lethal outcome from Vietnam to the Irkutsk region. Infectious Diseases (Russian journal) vol.12(3), pp.95-100	http://elib.fesmu.ru/elib/Article.aspx?id=305817
2.	Самсонова А.П., Петров Е.М., Савельева О.В., Иванова А.Е., Шарапова Н.Е. Анализ документированных результатов исследования сывороток крови больных, подозрительных на заболевание лептоспирозами, в реакции микроагглютинации // Инфекция и	Samsonova A.P., Petrov E.M., Savelyeva O.V., Ivanova A.E., Sharapova N.E. Analyzing the documented results by using microscopic agglutination test to examine sera from patients suspected of leptospirosis // Russian Journal of Infection and Immunity =	doi: 10.15789/2220-7619-ATD-1758

	иммунитет. 2022. Т. 12, № 5. С. 875–890. doi: 10.15789/2220-7619-ATD-1758	Infektsiya i immunitet, 2022, vol. 12, no. 5, pp. 875–890.	
3.	Соболева Г. Л., Ананьина Ю.В., Непоклонова И.В. Актуальные вопросы лептоспироза людей и животных /Российский ветеринарный журнал. — 2017. — № 8. — С. 13-17.	Soboleva G.L., Ananyina Y.V., Nepoklonova I.V., Actual Problems of Human and Animal Leptospirosis, Rossijskij veterinarnyj zhurnal (Russian veterinary journal), 2017, No. 8, pp. 13-17.	https://cyberleninka.ru/article/n/aktualnye-voprosy-leptospiroza-lyudey-i-zhivotnyh
4.	Стоянова Н. А., Токаревич Н.К., Волкова Г.В., Грачева Н. А., Кравченко С. С., Кузина Н. В., Лисеева Т. М., Мацьевская Е. А., Пьяных В. А., Снегирев В. И., Сосницкий В.И. Актуальные проблемы лептоспирозной инфекции в Северо-Западном федеральном округе // Эпидемиология и вакцинопрофилактика. 2003. №4 (11).	Stoianova N, Tokarevich N, Gracheva L, Volkova G, Gracheva N, Kravchenko S, Kuzina N, Liseeva T, Matsievskaya E, Snegirev V, Sosnitsky V. (2003). Actual problems of leptospirosis infection in the Northwestern Federal District. Epidemiology and vaccination (4 (11)), pp. 29-32.	URL: https://cyberleninka.ru/article/n/aktualnye-problemy-leptospiroznoy-infektsii-v-severo-zapadnom-federalnom-okruse (дата обращения: 27.09.2023).

5.	Стоянова Н А., Сергейко Л. М., Слепцова В. И. Иммунологический мониторинг и эпидемические особенности leptospiroza в Санкт-Петербурге // Микробиология эпидемиология и иммунобиология. 1996. - №6. - С. 120 - 122.	Stoianova N, Sergeiko L, Sleptsova V. (1996). Immunological monitoring and the epidemiological characteristics of leptospirosis in Saint Petersburg. Journal of Microbiology, Epidemiology and Immunobiology (6), pp.120-122.	-
6.	Токаревич Н.К., Стоянова Н.А. Эпидемиологические Аспекты Антропогенного Влияния На Эволюцию Leptospiroza // Инфекция и иммунитет. - 2011. - Т. 1. - №1. - С. 67-76.	Tokarevich N.K., Stoyanova N.A. (2011). Epidemiological aspects of anthropogenic influence to leptospirosis evolution // Russian Journal of Infection and Immunity Vol. 1. N. 1. pp. 67-76.	doi: 10.15789/2220-7619-2011-1-67-76
7.	Behera, S. K., Sabarinath, T., Ganesh, B., Mishra, P. K. K., Niloofa, R., Senthilkumar, K., Verma, M. R., Hota, A., Chandrasekar, S., Deneke, Y., Kumar, A., Nagarajan, M., Das, D., Khatua, S., Sahu, R., & Ali, S. A.	-	https://doi.org/10.3390/diagnostics12061455

	(2022). Diagnosis of Human Leptospirosis: Comparison of Microscopic Agglutination Test with Recombinant LigA/B Antigen-Based In-House IgM Dot ELISA Dipstick Test and Latex Agglutination Test Using Bayesian Latent Class Model and MAT as Gold Standard. <i>Diagnostics</i> (Basel, Switzerland), 12(6), 1455.		
8.	Bharti, A. R., Nally, J. E., Ricardi, J. N., Matthias, M. A., Diaz, M. M., Lovett, M. A., Levett, P. N., Gilman, R. H., Willig, M. R., Gotuzzo, E., Vinetz, J. M., & Peru-United States Leptospirosis Consortium (2003). Leptospirosis: a zoonotic disease of global importance. <i>The Lancet. Infectious diseases</i> , 3(12), 757–771.	-	https://doi.org/10.1016/s1473-3099(03)00830-2

9.	Boonsilp, S., Thaipadungpanit, J., Amornchai, P., Wuthiekanun, V., Bailey, M. S., Holden, M. T., Zhang, C., Jiang, X., Koizumi, N., Taylor, K., Galloway, R., Hoffmaster, A. R., Craig, S., Smythe, L. D., Hartskeerl, R. A., Day, N. P., Chantratita, N., Feil, E. J., Aanensen, D. M., Spratt, B. G., ... Peacock, S. J. (2013). A single multilocus sequence typing (MLST) scheme for seven pathogenic Leptospira species. <i>PLoS neglected tropical diseases</i> , 7(1), e1954.	-	https://doi.org/10.1371/journal.pntd.0001954
10.	Bourhy, P., Collet, L., Brisson, S., & Picardeau, M. (2014). <i>Leptospira mayottensis</i> sp. nov., a pathogenic species of the genus <i>Leptospira</i> isolated from humans. <i>International journal of systematic</i>	-	https://doi.org/10.1099/ijjs.0.066597-0

	and evolutionary microbiology, 64(Pt 12), 4061–4067.		
11.	Brenner, D. J., Kaufmann, A. F., Sulzer, K. R., Steigerwalt, A. G., Rogers, F. C., & Weyant, R. S. (1999). Further determination of DNA relatedness between serogroups and serovars in the family Leptospiraceae with a proposal for <i>Leptospira alexanderi</i> sp. nov. and four new <i>Leptospira</i> genomospecies. International journal of systematic bacteriology, 49 Pt 2, 839–858.	-	https://doi.org/10.1099/00207713-49-2-839
12.	Cerqueira, G. M., McBride, A. J., Queiroz, A., Pinto, L. S., Silva, E. F., Hartskeerl, R. A., Reis, M. G., Ko, A. I., & Dellagostin, O. A. (2010). Monitoring <i>Leptospira</i> strain collections: the need for quality control. The	-	https://doi.org/10.4269/ajtmh.010.09-0558

	American journal of tropical medicine and hygiene, 82(1), 83–87.		
13.	Chappel, R. J., Goris, M., Palmer, M. F., & Hartskeerl, R. A. (2004). Impact of proficiency testing on results of the microscopic agglutination test for diagnosis of leptospirosis. <i>Journal of clinical microbiology</i> , 42(12), 5484–5488.	-	https://doi.org/10.1128/JCM.42.12.5484-5488.2004
14.	Chen, H. W., Lukas, H., Becker, K., Weissenberger, G., Halsey, E. S., Guevara, C., Canal, E., Hall, E., Maves, R. C., Tilley, D. H., Kuo, L., Kochel, T. J., & Ching, W. M. (2018). An Improved Enzyme-Linked Immunoassay for the Detection of Leptospira-Specific Antibodies. <i>The American journal of tropical medicine and hygiene</i> , 99(2), 266–274.	-	https://doi.org/10.4269/ajtmh.17-0057

15.	Clarridge J. E., 3rd (2004). Impact of 16S rRNA gene sequence analysis for identification of bacteria on clinical microbiology and infectious diseases. <i>Clinical microbiology reviews</i> , 17(4), 840–862.	-	https://doi.org/10.1128/CMR.17.4.840-862.2004
16.	Costa, F., Hagan, J. E., Calcagno, J., Kane, M., Torgerson, P., Martinez-Silveira, M. S., Stein, C., Abela-Ridder, B., & Ko, A. I. (2015). Global Morbidity and Mortality of Leptospirosis: A Systematic Review. <i>PLoS neglected tropical diseases</i> , 9(9), e0003898.	-	https://doi.org/10.1371/journal.pntd.0003898
17.	Di Azevedo, M. I. N., & Lilienbaum, W. (2021). An overview on the molecular diagnosis of animal leptospirosis. <i>Letters in applied microbiology</i> , 72(5), 496–508.	-	https://doi.org/10.1111/lam.13442

18.	Faine, S., Adler, B., Bolin, C., & Perolat, P. (1999). "Leptospira" and leptospirosis. MediSci.	-	https://www.scirp.org/(S(i43dy n45teexjx455qlt3d2q))/reference/ReferencesPapers.aspx?ReferenceID=1391678
19.	Fortes-Gabriel, E., Guedes, M. S., Shetty, A., Gomes, C. K., Carreira, T., Vieira, M. L., Esteves, L., Mota-Vieira, L., & Gomes-Solecki, M. (2022). Enzyme immunoassays (EIA) for serodiagnosis of human leptospirosis: specific IgG3/IgG1 isotyping may further inform diagnosis of acute disease. PLoS neglected tropical diseases, 16(2), e0010241.	-	https://doi.org/10.1371/journal.pntd.0010241
20.	Fouts, D. E., Matthias, M. A., Adhikarla, H., Adler, B., Amorim-Santos, L., Berg, D. E., Bulach, D., Buschiazzo, A., Chang, Y. F., Galloway, R. L., Haake, D. A., Haft, D. H.,	-	https://doi.org/10.1371/journal.pntd.0004403

	Hartskeerl, R., Ko, A. I., Levett, P. N., Matsunaga, J., Mechaly, A. E., Monk, J. M., Nascimento, A. L., Nelson, K. E., ... Vinetz, J. M. (2016). What Makes a Bacterial Species Pathogenic?: Comparative Genomic Analysis of the Genus Leptospira. <i>PLoS neglected tropical diseases</i> , 10(2), e0004403.		
21.	Ghazaei C. (2018). Pathogenic Leptospira: Advances in understanding the molecular pathogenesis and virulence. <i>Open veterinary journal</i> , 8(1), 13–24.	-	https://doi.org/10.4314/ovj.v8i1 .4
22.	Guedes, I. B., de Souza, G. O., de Paula Castro, J. F., Cavalini, M. B., de Souza Filho, A. F., Maia, A. L. P., Dos Reis, E. A., Cortez, A., & Heinemann, M. B. (2021). Leptospira interrogans serogroup Pomona	-	https://doi.org/10.1007/s11250-021-02623-4

	strains isolated from river buffaloes. Tropical animal health and production, 53(2), 194.		
23.	Guernier, V., Allan, K. J., & Goarant, C. (2018). Advances and challenges in barcoding pathogenic and environmental Leptospira. <i>Parasitology</i> , 145(5), 595–607.	-	https://doi.org/10.1017/S0031182017001147
24.	Haake, D. A., Suchard, M. A., Kelley, M. M., Dundoo, M., Alt, D. P., & Zuerner, R. L. (2004). Molecular evolution and mosaicism of leptospiral outer membrane proteins involves horizontal DNA transfer. <i>Journal of bacteriology</i> , 186(9), 2818–2828.	-	https://doi.org/10.1128/JB.186.9.2818-2828.2004
25.	Hookey JV, Bryden J, Gatehouse L. (1993). The use of 16S rDNA sequence analysis to investigate the phylogeny of Leptospiraceae	-	doi: 10.1099/00221287-139-11-2585

	and related spirochaetes. J Gen Microbiol. 139(11):2585-2590.		
26.	Jayasundara, D., Senavirathna, I., Warnasekara, J., Gamage, C., Siribaddana, S., Kularatne, S. A. M., Matthias, M., Mariet, J. F., Picardeau, M., Agampodi, S., & M Vinetz, J. (2021). 12 Novel clonal groups of Leptospira infecting humans in multiple contrasting epidemiological contexts in Sri Lanka. PLoS neglected tropical diseases, 15(3), e0009272.	-	https://doi.org/10.1371/journal.pntd.0009272
27.	Khodaverdi Darian, E., Forghanifard, M. M., Moradi Bidhendi, S., Chang, Y. F., Yahaghi, E., Esmaelizad, M., Khaleghizadeh, M., & Khaki, P. (2013). Cloning and Sequence Analysis of LipL32, a Surface-Exposed Lipoprotein of	-	https://doi.org/10.5812/ircmj.8793

	Pathogenic Leptospira Spp. Iranian Red Crescent medical journal, 15(11), e8793.		
28.	Kmety E, Dikken H. (1993). Classification of the species <i>Leptospira interrogans</i> and history of its serovars. Groningen, The Netherlands: University Press Groningen;	-	-
29.	Ko, A. I., Goarant, C., & Picardeau, M. (2009). <i>Leptospira</i> : the dawn of the molecular genetics era for an emerging zoonotic pathogen. <i>Nature reviews. Microbiology</i> , 7(10), 736–747.	-	https://doi.org/10.1038/nrmicro2208
30.	Lam, J. Y., Low, G. K., & Chee, H. Y. (2020). Diagnostic accuracy of genetic markers and nucleic acid techniques for the detection of <i>Leptospira</i> in clinical samples: A meta-analysis. <i>PLoS neglected tropical diseases</i> , 14(2), e0008074.	-	https://doi.org/10.1371/journal.pntd.0008074

31.	Landolt, N. Y., Chiani, Y. T., Pujato, N., Jacob, P., Schmeling, M. F., García Effron, G., & Vanasco, N. B. (2022). Utility evaluation of two molecular methods for Leptospira spp. typing in human serum samples. <i>Heliyon</i> , 9(2), e12564.	-	https://doi.org/10.1016/j.heliyon.2022.e12564
32.	Levett PN. (2001). Leptospirosis. <i>Clin Microbiol</i> 14(2):296-326.	-	doi:10.1128/CMR.14.2.296-326.2001
33.	Masuzawa, T., Sakakibara, K., Saito, M., Hidaka, Y., Villanueva, S. Y. A. M., Yanagihara, Y., & Yoshida, S. I. (2018). Characterization of Leptospira species isolated from soil collected in Japan. <i>Microbiology and immunology</i> , 62(1), 55–59.	-	https://doi.org/10.1111/1348-0421.12551
34.	Morey, R. E., Galloway, R. L., Bragg, S. L., Steigerwalt, A. G., Mayer, L. W., & Levett,	-	https://doi.org/10.1128/JCM.00670-06

	P. N. (2006). Species-specific identification of Leptospiraceae by 16S rRNA gene sequencing. <i>Journal of clinical microbiology</i> , 44(10), 3510–3516.		
35.	Philip N, Bahtiar Affendy N, Ramli SNA, Arif M, Raja P, Nagandran E, et al. (2020) <i>Leptospira interrogans</i> and <i>Leptospira kirschneri</i> are the dominant <i>Leptospira</i> species causing human leptospirosis in Central Malaysia. <i>PLoS Negl Trop Dis</i> 14(3): e0008197.	-	https://doi.org/10.1371/journal.pntd.0008197
36.	Picardeau M. (2017). Virulence of the zoonotic agent of leptospirosis: still terra incognita. <i>Nature reviews. Microbiology</i> , 15(5), 297–307.	-	https://doi.org/10.1038/nrmicro.2017.5

37.	Picardeau, Mathieu. (2020). Leptospira and Leptospirosis. 10.1007/978-1-0716-0459-5_24.	-	https://pubmed.ncbi.nlm.nih.gov/v/32632877/
38.	Postic D, Riquelme-Sertour N, Merien F, Perolat P, Baranton G. (2000). Interest of partial 16S rDNA gene sequences to resolve heterogeneities between Leptospira collections: application to <i>L. meyeri</i> . <i>Res Microbiol.</i> 151(5):333-341.	-	doi:10.1016/s0923-2508(00)00156-x
39.	Rajapakse S. (2022). Leptospirosis: clinical aspects. <i>Clinical medicine (London, England)</i> , 22(1), 14–17.	-	https://doi.org/10.7861/clinmed.2021-0784
40.	Slack, A. T., Symonds, M. L., Dohnt, M. F., & Smythe, L. D. (2006). Identification of pathogenic Leptospira species by conventional or real-time PCR and	-	https://doi.org/10.1186/1471-2180-6-95

	sequencing of the DNA gyrase subunit B encoding gene. BMC microbiology, 6, 95.		
41.	Stimson AM. (1907). Note on an Organism Found in Yellow-Fever Tissue. Public Health Reports (1896-1970) Vol. 22(18):541.	-	DOI: 10.2307/4559008
42.	Strutzberg-Minder, K., Ullerich, A., Dohmann, K., Boehmer, J., & Goris, M. (2022). Comparison of Two Leptospira Type Strains of Serovar Grippotyphosa in Microscopic Agglutination Test (MAT) Diagnostics for the Detection of Infections with Leptospires in Horses, Dogs and Pigs. Veterinary sciences, 9(9), 464.	-	https://doi.org/10.3390/vetsci9090464
43.	Sugunan, A. P., Vijayachari, P., Sharma, S., Roy, S., Manickam, P., Natarajaseenivasan,	-	https://pubmed.ncbi.nlm.nih.gov/19700804/

	K., Gupte, M. D., & Sehgal, S. C. (2009). Risk factors associated with leptospirosis during an outbreak in Middle Andaman, India. <i>The Indian journal of medical research</i> , 130(1), 67–73.		
44.	Sykes, J. E., Gamage, C. D., Haake, D. A., & Nally, J. E. (2022). Understanding leptospirosis: application of state-of-the-art molecular typing tools with a One Health lens. <i>American journal of veterinary research</i> , 83(10), ajvr.22.06.0104.	-	https://doi.org/10.2460/ajvr.22.06.0104
45.	Torgerson, P. R., Hagan, J. E., Costa, F., Calcagno, J., Kane, M., Martinez-Silveira, M. S., Goris, M. G., Stein, C., Ko, A. I., & Abela-Ridder, B. (2015). Global Burden of Leptospirosis: Estimated in Terms of Disability Adjusted Life Years. <i>PLoS</i>	-	https://doi.org/10.1371/journal.pntd.0004122

	neglected tropical diseases, 9(10), e0004122.		
46.	Zhang, C., Yang, H., Li, X., Cao, Z., Zhou, H., Zeng, L., Xu, J., Xu, Y., Chang, Y. F., Guo, X., Zhu, Y., & Jiang, X. (2015). Molecular Typing of Pathogenic Leptospira Serogroup Icterohaemorrhagiae Strains Circulating in China during the Past 50 Years. PLoS neglected tropical diseases, 9(5), e0003762.	-	https://doi.org/10.1371/journal.pntd.0003762