

# First detection of Mycoplasma wenyonii in France: Identification, evaluation of the clinical impact and development of a new specific detection assay

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#### 1 Title

2 First detection of *Mycoplasma wenyonii* in France: identification, evaluation of the clinical
3 impact and development of a new specific detection assay.

4

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20

#### 21 Abstract

- 22 Mycoplasma wenyonii, a hemoplasma infecting cattle, was never detected in France. In 2014,
- 23 evocative inclusions were observed in erythrocytes from cattle presenting milk drops, anemia,
- and edema in Brittany (France). A survey was then initiated to investigate the epidemiological
- situation and correlate mycoplasma detection with clinical signs. For this purpose, a new PCR

assay targeting *polC* gene was designed. Comparative results with published PCR assays 26 place this new one as more specific, allowing a one-step diagnosis without further sequencing. 27 A total of 181 cows were included in this study and 4.97% (n=9) were positive, resulting in 28 the first molecular identification of *M. wenyonii* in France. All positive animals presented 29 anemia, edema and milk drop. When selecting animals presenting evocative clinical signs, the 30 prevalence of *M. wenyonii* in Brittany was estimated to 25.6%. Further studies are needed to 31 evaluate the importance of the infection, the implication of arthropods and the existence of 32 asymptomatic carriers. 33

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35 Key words: *Mycoplasma wenyonii*, *Mycoplasma ovis*, hemoplasma, cattle, anemia, *polC*.

36

#### 37 **Introduction**:

38 After a period of confusion, Haemobartonella and Eperythrozoon species were reclassified as haemotropic mycoplasmas (hemoplasmas) based on molecular phylogenetic evidences 39 40 (Neimark et al., 2001; Neimark et al. 2002). Haemoplasmas species are commonly infecting erythrocytes of a wide range of animals. Two species are suspected to infect cattle: 41 Mycoplasma wenyonii, formerly known as Eperythrozoon wenyonii, and Candidatus 42 Mycoplasma haemobos. Data on Candidatus Mycoplasma haemobos are very scarce 43 (Niethammer et al., 2018) and most reports on cattle haemoplasmas only relate to *M. wenvonii* 44 (Tagawa et al., 2008 and 2013). 45

*M. wenyonii* is an acknowledged pathogenic species infecting cattle worldwide. Blood smears
reveal that this mycoplasma is found attached at the surface of red blood cells and/or free in
the plasma (Neimlark et al., 2001). Co-infections have been demonstrated with *A. marginale*, *A. phagocytophilum*, *Babesia* spp. and *Theileria* spp. (Hofmnn et al., 2004). In cattle, the
majority of *M. wenyonii* infections remains subclinical (Montes et al., 1994; Smith et al.,

51 1990), but signs such anemia, pyrexia, hind limb edema (hock), swollen teats, decreased milk 52 production and weight loss have been observed (Smith et al., 1990; Genova et al., 2011; 53 Collin et al., 2015). Chronical infection has also been reported in a Charolais bull in the USA 54 (Montes et al., 1994). The use of oxytetracycline is an effective therapeutics in most of the 55 cases. However, no antibiotic protocol has been proven to eliminate infection and infected 56 animals may remain carriers for life (Genova et al., 2011).

Modalities of dissemination are unknown but horizontal transmission by blood-sucking 57 arthropods has been basically accepted as the most common route for haemoplasma infection 58 (Hofmann-Lehmann et al., 2004). Abortion in association with M. wenyonii detection from 59 60 dam blood suggested that vertical transmission to the conceptus may be an alternative route (Sasaoka et al., 2015). Some studies have reported that dam-to-fetus transmission is possible, 61 but the importance of this route for infection is unknown too (Hornok et al., 2011; Fujihara et 62 63 al., 2011). Vertical transmission seems possible for Candidatus Mycoplasma haemobos too (Niethammer et al., 2018). Risk factors associated to infection have been poorly explored. 64 65 Housed cattle seem more susceptible to infection that grazing cattle and cattle between 1 and 3 years old appear to be the more susceptible to infection (Tagawa et al., 2012). 66

In cattle presenting clinical signs, the usual diagnostic method relies on blood smears stained 67 with Giemsa, but this technique shows low specificity and sensitivity (i.e. possible Howell-68 Jolli bodies artifact) (Smith et al., 1990). More recently, methods developed for diagnosis 69 included qualitative PCR (McAuliffe et al., 2006), real-time PCR (Willi et al., 2009; Meli et 70 al., 2010), direct PCR from whole blood sample without DNA extraction (Tagawa et al., 71 72 2012), and the loop-mediated isothermal amplification (LAMP) assay (Song et al., 2013). Yet, these methods have on main drawback in that they were unsatisfactory to discriminate easily 73 bovine hemoplasmas from other mycoplasmas of ruminant origin (i.e. M. ovis), which is an 74

important objective (Aktas and Ozubek, 2017). To overcome this issue, we developed ahighly specific PCR assay including a plasmid for positive control.

We report hereafter the first clinical cases due to M. wenyonii (MW) observed in a dairy herd 77 Brittany, France. Suspected cases, based on visual examination of blood smears or imprecise 78 PCR assays have been presented as oral presentation in cattle medicine congresses, but this is 79 subsequently the first identification of MW in France. The clinical signs consisted in 80 hyperthermia (up to 40°C), anemia, and lower limbs and teat edema, and the disease affected 81 up to 40% of the cows. Blood smears and molecular evidence using protocols as described by 82 Tagawa et al. (2008) and Sasaoka et al. (2015), confirmed the diagnosis. Our second 83 objective was to evaluate the feasibility and performances of an original one step detection 84 based on gene *polC*. 85

86

#### 87 Materials and Methods

#### 88 DNA samples

89 DNA was extracted from 200 µl of EDTA-blood samples collected from 2 cattle with clinical signs and positive smear evocating infection by Mycoplasma wenyonii (sample 141145 and 90 141146, collected in France (Collin et al., 2015) using a QIAamp DNA Blood Mini Kit 91 (QIAGEN), eluted with 100  $\mu$ l of buffer AE and stored at  $-20^{\circ}$ C until use. DNA extract from 92 sample known as positive from previous investigation on *M. wenyonii* in Japan[16] was used 93 as positive control for PCR detection. For defining the specificity of the PCR assay, genomic 94 DNA from other mycoplasma species (Mycoplasma ovis, M. fermentans; M. hominis; M. 95 orale; Achleplasma laidlawi; M. capricolum subsp. capricolum; M. mycoides subsp. capri; 96 *M. leachi*; *M. putrefaciens*; *M. ovipneumoniae*; *M. bovigenitalium*; 97 *M. conjunctivae*; M. arginini; M. gallisepticum; M. canis; U. diversum; M. bovis; M. agalactiae) were used and 98 extracted from liquid culture as described elsewhere (Sambrook and Russel, 2001; Chen and 99

100 Kuo, 1993) except for *M. ovis* for which it was extracted from whole blood of a positive 101 animal. The *Mycoplasma* species have been chosen either for their renown existence in 102 ruminants (n=13) or for their existence in common domestic animals or humankind (n=5)

#### 103 PCR detection assays

Four set of primers were used for PCR assays (Table 1). Primers for 16S\_rRNA and primers for rnpB were designed according to Tagawa et al.2008 and Sasaoka et al. 2015 respectively. Based on the alignment *pol*C sequences available in databases for MW and MO, forward and reverse primers were designed to amplify a 500 bp product. Using the Primer blast software, these primers were predicted *in silico* to specifically amplify MW. The same strategy was used to designed MO specific primers.

PCR assays were performed using an Eppendorf Mastercycler ep-Gradient thermocycler in 25 110 µL reaction mixtures containing 0.4 mM of each primer, 1X PCR reaction buffer (with 111 112 MgSO4, New England Biolabs [NEB], Evry, France), 200 mM dNTPs and 2.5 U Taq DNA polymerase (NEB). Reaction mixtures were subjected to 2 min at 94°C, 30 cycles (except for 113 114 16S rRNA: 35) of 30s at 94 °C, 30s at 58°C (except for 16S rRNA: 62°C), 30s at 72 °C, and a 115 final elongation step of 5 min at 72°C. Annealing temperature was determined based on data obtained from a gradient of temperature. PCR products were analyzed by gel electrophoresis 116 in 2% agarose for 16S RNA (Tagawa et al. 2008) and *rnpB* (Sasaoka et al. 2015) assays or in 117 1% agarose for MW-polC and MO-polC assays (Molecular weight standard: Smart Ladder, 118 Eurogentec, Liège, Belgium). 119

#### 120 PCR sensitivity and design of replicative plasmid as positive controls

PCR products were purified (QIAquick PCR Purification Kit, QIAGEN, Courtaboeuf, France)
and cloned into pGEMT-easy plasmid vector (Promega, Charbonnières les bains, France).
Recombinant plasmid vectors were transformed in *Escherichia coli* strain DH5a
(Invitrogen/Life-Technologies, Courtaboeuf, France) grown in LB medium supplemented

with ampicillin, and purified with QIAprep Spin Miniprep kit (QIAGEN). DNA fragment 125 insertion was monitored by PCR with the appropriate specific primers, followed by 126 sequencing. Plasmid concentration was determined by Qubit Fluorometric Quantitation 127 (ThermoFischer, Villebon sur Yvette, France). PCR assays were performed with serial 128 dilution of the purified plasmid as templates to determine the minimal concentration for which 129 amplification is detected. Each experiment was repeated three times to assess repeatability 130 and we obtained every time the same results. For comparison, the PCR assay was run in 131 parallel with serial dilution of positive samples. 132

#### 133 **DNA sequencing**

Sequences of PCR product and plasmid were obtained by Sanger direct sequencing method
using specific primers at the sequencing facility of UMR 5165 (CNRS, UPS, CHU Purpan,
Toulouse, France). Sequence analysis was done with Chromas lite (available at
http://www.technelysium.com.au/chromas\_lite.html) and NCBI blast tools
(https://blast.ncbi.nlm.nih.gov/).

#### 139 Case detection study

140 A case detection study was implemented in 2016 with the help of veterinarians in French Brittany. Cases were included in the study on the basis of cattle presenting: anemia without 141 hemoglobinuria, edema of the limbs and/or the udder, transient milk drop, excluding 142 anaplasmosis and babesiosis, medical history was recorded. A total of 181 cows in 6 herds 143 were included in the study: 34 suspected clinical cases were collected and 147 samples from 144 animals without clinical signs were also collected at the same time and from the same region. 145 Animal blood samples were collected (5 mL, EDTA) for blood-smear primary detection using 146 Diff Ouick® coloration and, DNA extraction (OIAamp DNA Blood Mini Kit, Oiagen) for 147 PCR detection using MW-polC amplification. 148

149 Nucleotide sequence accession numbers.

150 Nucleotide sequences have been submitted to GenBank under accession numbers MH001426,

151 MH001427, MH001428 and MH001429 respectively.

- 152
- 153 **Results**

#### 154 First detection of hemoplasmas in French ruminants

155 Using the PCR assay developed by Tagawa et al. in 2008, the presence of MW was detected in 9 blood samples (i) collected in France. We selected only 2 after the sequencing process 156 (due to the 100% level of similarity between the 9) from two cows suspected to be infected 157 158 by MW, namely 141145 and 141146, (see Material and methods) and (ii) from an animal known as being infected by MW, as positive control. More specifically, with these samples, a 159 PCR product of the expected size, 193-bp, was detected. Regardless of the stringency of the 160 PCR conditions, the same PCR assay yielded a similar size product with DNA extracted from 161 M. bovis, another ruminant mycoplasma species known to infect ruminant and initially 162 included in our study as a negative control (Figure 1A). The 193-bp PCR product of 141145, 163 141146 and *M. bovis* were sequenced. Data showed that 141145 and 141146 sequences are 164 identical and closely matched sequences belonging to the 16s rRNA of MW and MO, with 0 165 to 2 SNPs, depending on the strains used for comparison (Figure 1B). For *M. bovis*, the 166 sequence was identical to the of *M. bovis* 16s rRNA. All together these data suggested that the 167 PCR developed by Tagawa et al. (2008), while detecting hemoplasmas, does not discriminate 168 169 between MO, MW and MB. When we tested the PCR assay developed by Sasaoka et al. (2015), which one targets the *rnpB* gene, *M. bovis* DNA gave a negative result while 141145 170 and 141146 yield the expected size product which sequences match specifically the MW. Yet, 171 172 MO was positive (Figure 1C).

173

#### 174 Development of a specific PCR for MW and MO

Overall, these results reflect the lack of a proper diagnostic assay that would allow 175 unambiguously the detection of MW in one single step. To fill this gap, we designed a PCR 176 assay targeting the *pol*C gene to amplify specifically a 500 bp product. This was further 177 confirmed *in vitro* using genomic DNA extracted from MW, and MB with a 55°C optimum 178 temperature (Figure 1D). The same strategy was used to designed MO specific primers. The 179 specificity of these PCR assays, further designated as MW-polC or MO-polC, was then tested 180 against 19 Mycoplasma species, of which 15 are known in ruminants (Table 2). Results 181 shown in Table 2 and illustrated in Figure 1 were compared to PCR assays previously 182 developed. All species but two, M. canis and U. diversum, were detected in our hands with the 183 assay targeting the 16S rRNA (Tagawa et al; 2008). The PCR assay developed by Sasaoka et 184 al. (2015) was more specific; yet amplification was obtained with MO. Overall, in our hands, 185 comparison of the different PCR assays currently available points towards MW-*polC* as being 186 187 the more specific for MW detection.

#### 188 Development of molecular positive controls

189 Since MW and MO are not cultivable, the use of a field sample as positive control which 190 availability is limited poses the question of the reproducibility and of the estimation of the sensitivity of the assay. To address these issues, we engineered a set of 4 replicative plasmids 191 carrying sequences targeted by the PCR assays and corresponding to: the MW- or MO- 530 192 193 bp or 545 bp *polC* sequence respectively, the 189 bp MW *rnpB* sequence and the MW 16s rRNA. Replicates of PCR assays using serial dilution of known plasmid concentration 194 indicates that MW-polC is less sensitive than other PCR assays, with that targeting the two 195 16s rRNA gene copies being as expected the most sensitive (Table 3 and Figure 2A). 196 Replicates of PCR assays using serial dilution of a field sample (Figure 2B) permit to confirm 197 198 these sensitivity differences and estimate the bacterial load of a clinically infected animal to

199  $10^7$  bacteria/ml of blood assessing that the sensitivity of MW-*polC* assay is sufficient enough 200 to detect clinical cases.

#### 201 Case detection study

Cases were recruited on basis of clinical signs evocating infection by *M. wenyonii* as detected by veterinarians in Brittany. Animals without clinical signs were also sampled at the same time. A total of 34 samples from adult dairy cattle presenting clinical signs were collected and 147 from animals without clinical signs. All were analyzed with MW-*polC* PCR assay. Of the 34 animals presenting signs evocating *M. wenyonii* infection, 9 were positive while the 147 samples from asymptomatic animals were all negative (Figure 3).

#### 208 Discussion

This study presents the molecular and clinical identification of a hemoplasma in cattle in 209 France. To the authors' knowledge, this is the first report of a molecular investigation of M. 210 211 wenyonii in this country. Hemoplasma organisms (named before 2001 Eperythrozoon) have been identified in a variety of mammals (including cattle) and the infection by these 212 213 organisms has been associated with extravascular hemolytic anemia, reproductive disorders, 214 milk drop syndrome, edema of dependent parts, growth retardation... of varying intensity: subclinical infection seems common (Messik, 2004). In Brittany (France), evocative 215 inclusions in bovine erythrocytes have been observed for the first time in 2014, associated 216 217 with clinical signs in adult cows (Collin et al., 2015).

Our study confirms the presence of *Mycoplasma wenyonii* in France. Surprisingly, in our survey 'Candidatus M. haemobos' was not detected unlike in other European countries (Ade et al., 2018; Niethammer et al. 2018), but this species is even less documented that MW. As expected, the number of infected animals by MW is much higher when selecting them on the basis of a clinical suspicion (9/34: 25.6 %) than when sampling them at random in the same area (9/181: 4.9 %). The selected significant clinical signs were: anemia without

hemoglobinuria, edema of the limbs and/or the udder, transient milk drop. Of course clinical
anaplasmosis and babesiosis were excluded by optical microscopy and PCR (data not shown).
No coinfection was observed in our study but is likely to occur from time to time since their
potential vectors share the same biotope (Hornok et al., 2011, Messik, 2004).

Additionally, the purpose of our study was to address the specificity of previously 228 published PCR assays, and to develop a specific molecular diagnosis a one-step assay, 229 without any subsequent amplification. We demonstrated that no other assay is as discriminant 230 as the one we developed: using MW-*polC* amplification (with the probes we developed) 231 allows detecting only Mycoplasma wenyonii, and no other mycoplasma of ruminant origin 232 (Figures 4 and 5). Moreover, the amplicon size is Mw-specific and quite different from M. 233 ovis amplicons, an important objective (Aktas and Ozubek, 2017) we fulfilled. At first, the 234 suitability of this assay in a real-time PCR was not the purpose of this study. 235

Sensitivity of the existing and new test has been evaluated. The detection threshold is as low as  $1.7 \ 10^6$  copies of the genome per mL total blood, but it is less sensitive than PCR assays targeting the 16S rRNA gene copies which are, as expected, the most sensitive. The future development of a real-time PCR will be the way to increase the sensitivity of the detection.

Sensitivity study was made possible thanks to the engineering of recombinant plasmids. These have several advantages over positive controls from the field infected animals whose reproducibility is poor and/or variable: they can be produce in large amount and offer reproducible and comparative data across labs.

Further studies are needed to evaluate the importance of MW infection. The systematic exploration of anemia in cattle could be the first step, as this symptom appears to be the more consistent in cattle of all ages. Studies on coinfections with *Theileria* spp., *Babesia* spp., *Anaplasma* spp. and *Bartonella* spp. are to be initiated too, because infectivity and disease

could be associated with coinfection, apart from other factors to determine. Correlations ofMW infection with productivity and growth might be evaluated subsequently.

As there is no clear consensus either on the potential vectors, or on transmission, arthropod

- trapping and investigation on asymptomatic or latent and chronic carriers are needed too.
- 252 The importance of vertical transmission is unknown too, and sampling calves before ingestion
- of colostral antibodies could help to specify the importance of this phenomenon.

254 The tools we developed can contribute to target these objectives.

255

#### 256 **Conflict of interest statement**

257 None of the authors of this paper has a financial or personal relationship with other people or

258 organizations that could inappropriately influence or bias the content of the paper

#### 259 Authorship

LX. Nouvel and MC. Hygonenq carried out DNA extraction and PCR design and analysis. E.

Martinelli and G. Catays carried out part of DNA extraction and PCR analysis. Ph. Lepage and E. Collin carried out animal sampling and case story collection. H.Inokuma provided positive controls and PCR protocols. C. Citti, LX. Nouvel and R. Maillard drafted the manuscript. LX. Nouvel, F. Schelcher, C. Citti and R. Maillard conceived and participated in the design of the study, which was coordinated by R. Maillard. All authors read and approved the final manuscript.

- 267
- 268 **Declarations of interest:**
- 269 None
- 270
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360	
361	Figure captions
362	Figure 1: M. wenyonii PCR assays
363 364 365 366 367 368 369	Princeps cases detection targeting 16S rRNA (A) or polC (C) genes, and assays specificity testing (B). T+: MW for 16S rRNA, <i>rnpB</i> and MW- <i>polC</i> ; MO for MO-polC; M: Smart ladder Eurogentec; 1: <i>Mycoplasma fermentans</i> ; 2: <i>Achleplasma laidlawi</i> ; 3: <i>M. capricolum</i> subsp. <i>capricolum</i> ; 4: <i>M. leachi</i> ; 5: <i>M. putrefaciens</i> ; 6: <i>M. ovipneumoniae</i> ; 7: <i>M. mycoides</i> subsp. <i>capri</i> ; 8: <i>M. bovigenitalium</i> ; 9: <i>M. conjunctivae</i> ; 10: <i>M. arginini</i> ; 11: <i>M. canis</i> ; 12: <i>U. diversum 310<sup>#</sup></i> ; 13: <i>U. diversum 246<sup>#£</sup></i> ; 14: <i>M. bovis</i> ; 15: <i>M. agalactiae</i> *; 16: <i>M. wenyonii*;</i> (*: DNA extract of blood from positive animal, <sup>#</sup> : culture after PK lysis, <sup>£</sup> : no culture growth)
370	Figure 1C: 16S and <i>polC</i> sequences from isolates 14-1145 and 14-1146
371	
372	Figure 2: Sensitivity testing of <i>M. wenyonii</i> PCR assays
373 374 375 376	(A) 16S rRNA, <i>rnpB</i> , MW- <i>polC and</i> MO- <i>polC</i> PCR with specific plasmid in serial dilution corresponding to 10 to 10 <sup>10</sup> gene copy as template (B) Quantification of gene copy in positive sample 14-1145: 16S rRNA, <i>rnpB</i> , MW- <i>polC</i> PCR with serial dilution of DNA extract as template.
377	Figure 3: Global survey results
378	Detection results in all animals or when selecting animal on clinical signs

**Figure 4:** *polC* gel *Mycoplasma wenyonii* specific

380

- **Figure 5:** *polC* phylogenetic tree
- 382
- 383 Tables
- 384

## **Table 1: Primers and PCR conditions**

Targeted gene	Primers	Ta*	Expected amplicon size		References
			M. wenyonii	M. ovis	
16S_rRNA	16S-F2 (5'- GGCCCATATTCCTRCGGGAAG- 3') 16S-R2 (5'- ACRGGATTACTAGTGATTCCA- 3')	62°C	193 bp	176 and 193 bp	Tagawa <i>et</i> <i>al.</i> [2008]
rnpB	rnpB-F (5'- AGTCTGAGATGACTRTAGTG- 3') rnpB-R (5'- TRCTTGMGGGGGTTTGCCTCG- 3')	58°C	189 bp	187 bp	Sasaoka <i>et</i> <i>al.</i> [2015]
MW_polC	MW-PolCF1 (5'- GTAGTACCACCACCAGAGCAG- 3') MW-PolCR1 (5'- ACTACCTCGTGCCCAAGTG-3')	58°C	530 bp	-	This study
MO_polC	MO-PolCF1 (5'- GGACAGTGAGCGGAGATGG-3') MO-PolCR1 (5'- AAGAGTGGGGGCTTGAATGGG- 3')	58°C	-	545 bp	This study

<sup>385 \*:</sup> temperature of annealing

## Table 2: Specificity of the PCR assays targeting the detection of MW

Species and	Host	PCR assay			
reference of the strain or isolate		16S rRNA	rnpB	MW-polC	MO-polC
M. wenyonii	Ruminants	+	+	+	_*
Isolate 14-1146 this study					
M. ovis	Ruminants	+*	+	_*	+
isolate 10-373, this study					
<i>M. fermentans</i> type strain PG 12	Human	+	-	-	-
M. hominis	Human	+	-	-	-
type strain PG21					
M. orale	Human	+	-	_*	_*
type strain CH 19299					
A. laidlawii	Free	+	-	-	_*
type strain PG8					
<i>M. capricolum</i> subsp. <i>Capricolum</i> type strain California kid	Ruminants	+	-	-	-
<i>M. mycoides</i> subsp. <i>Capri</i> type strain Y goat	Ruminants	+	-	-	-
<i>M. leachii</i> type strain PG50	Ruminants	+	-	-	-
<i>M. putrefaciens</i> type strain KST	Ruminants	+	-	-	-

<i>M. ovipneumoniae</i> type strain Y98	Ruminants	+	-	-	-
<i>M. bovigenitalium</i> type strain PG 11	Ruminants	+	-	-	-
<i>M. conjunctivae</i> type strain HRC581	Ruminants	+	-	-	-
<i>M. arginini</i> type strain G230	Ruminants	+	-	-	-
<i>M. gallisepticum</i> type strain PG31	Poultry	+	+f	-	-
<i>M. canis</i> type strain PG14	Ruminants	-	-	-	-
<i>U. diversum</i> type strain A417	Ruminants	-	-	-	-
<i>M. bovis</i> type strain PG45	Ruminants	+	-	-	-
<i>M. agalactiae</i> type strain PG2	Ruminants	+	-	-	-

 +: amplicon at expected size; -: no amplification detected; -\*: weak signal at a different size than expected; +f: positive weak signal at the expected size

## 390 Table 3: Sensitivity of the PCR assays

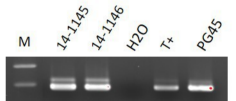
	Plasmid insert target	Detection threshold (genome/µL blood)
16S rRNA	MW	30
rnpB	MW	200
	МО	90
MW-polC	MW	1700
MO-polC	МО	240

391

MO: Mycoplasma ovis; MW: Mycoplasma wenyonii







>Seq1 [Mycoplasma wenyonii] M.wenyonii[14\_1145]partial16S

### 

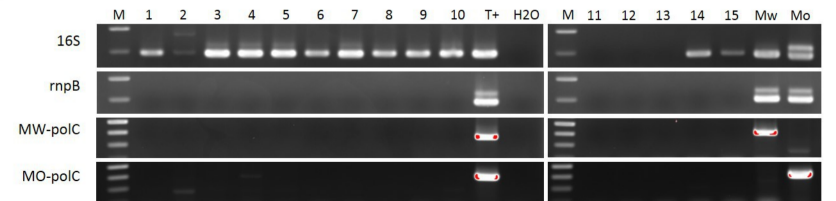
>Seq2 [Mycoplasma wenyonii] M.wenyonii[14\_1146]partial16S

> Seq3 [Mycoplasma wenyonii] M.wenyonii[14\_1145]polC

>Seq4 [Mycoplasma wenyonii] M.wenyonii[14-1146]polC

Figure 4 : 16S and *PolC* sequences of isolates 14-1145 et 14-1146

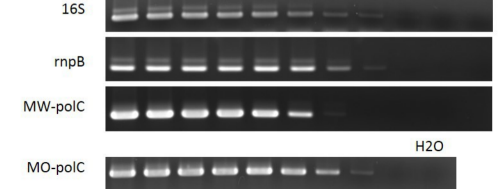
В

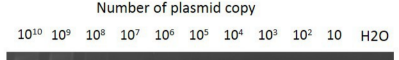








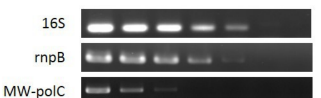






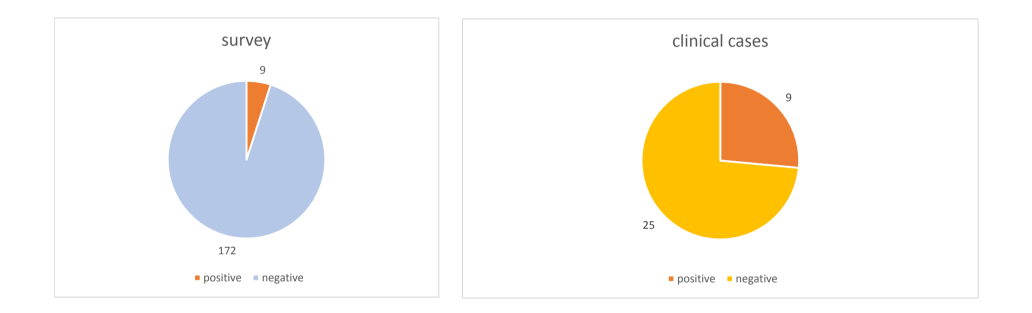


# Extract **14-1145**

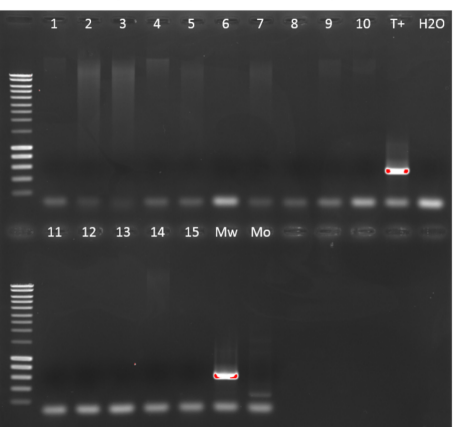


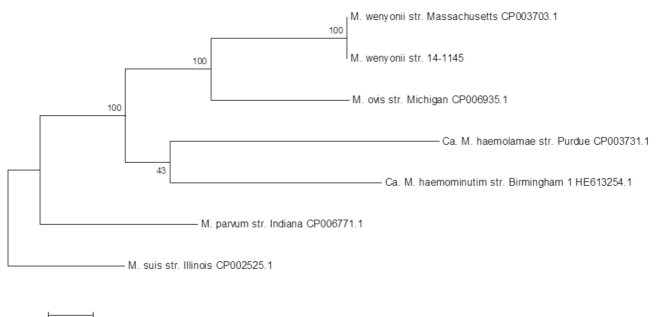


	survey		clinical cases
positive	9	positive	9
negative	172	negative	25



polC Mwen (2016-09-29 11hr16min)





#### 0.050

#### Phylogentic tree based on polC gene sequence

The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model [1]. The tree with the highest log likelihood (-22940.9572) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 7 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There were a total of 4202 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 [2].

1. Tamura K. and Nei M. (1993). Estimation of the number of nucleotide substitutions in the control region of mitochondrial DNA in humans and chimpanzees. Molecular Biology and Evolution 10:512-526.

2. Kumar S., Stecher G., and Tamura K. (2016). MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Molecular Biology and Evolution 33:1870-1874.

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