1	First isolation of "Brachyspira hampsonii" from pigs in Europe						
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Abstract. Swine dysentery in Europe is classically attributed to *Brachyspira hyodysenteriae*. 21 However, other Brachyspira species have been increasingly associated with intestinal 22 disorders in pigs. This case report describes the first diagnosis of a "Brachyspira hampsonii" 23 infection in European pigs. In a routine quarantine monitoring protocol, two gilts were 24 presented for necropsy, in which soft watery non-haemorrhagic colonic content was found. 25 Microbial culture from the colonic content and from faecal samples revealed the presence of 26 strongly haemolytic, ring-phenomenon positive spirochetes indicative for Brachyspira 27 hyodysenteriae. A diagnostic commercial PCR could not confirm the presence of B. 28 hyodysenteriae. Phenotypic characterisation and PCRs targeting the 16S rRNA, 23S rRNA, 29 nox, hlyA and tlyA genes of different swine-related Brachyspira spp. were performed. 30 Phylogenetic analysis of sequences of the partial nox and 16S rRNA genes and multi locus 31 sequence typing demonstrated that the isolates in this case were "B. hampsonii" isolates. This 32 33 case report shows that the diagnosis of infections caused by new, emerging Brachyspira species is not self-evident and that the combination of microbial culture and PCR is 34 35 recommended, completed with more extensive genotyping if necessary.

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Case report: first isolation of "Brachyspira hampsonii" from pigs in Europe

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Infections with Brachyspira spp. in swine occur in most swine-rearing countries and 45 can result in substantial economic losses. Of all swine-related Brachyspira spp. infections 46 classical swine dysentery, caused by *Brachyspira hyodysenteriae*, results in the most severe 47 clinical symptoms (eg. mucohaemorrhagic diarrhea, weight loss, poor feed conversion). B. 48 hvodysenteriae was first recognized as the cause of swine dysentery in 1971 (Taylor and 49 Alexander 1971). At that time, the strong haemolysis of *B. hyodysenteriae* appeared indicative 50 for pathogenicity since other, weakly haemolytic Brachyspira (formerly Serpulina, Serpula 51 52 and *Treponema*) appeared to be commensal and were therefore named *Brachyspira innocens* (Kinyon and Harris 1979). Several reports of clinical disease caused by weakly hemolytic 53 Brachyspira indicated that not all weakly hemolytic Brachyspira spp. were non-pathogenic 54 55 for pigs (Taylor and others 1980, Neef and others 1994). Further research of these weakly haemolytic isolates including DNA-DNA hybridisation, resulted in the designation of three 56 57 more weakly haemolytic species namely B. intermedia, B. murdochii and B. pilosicoli (Trott and others 1996, Stanton and others 1997). 58

These weakly haemolytic species of *Brachyspira* diverge in the severity of clinical symptoms they cause. *B. pilosicoli* is pathogenic and causes spirochetal colitis in pigs, which is marked by non-haemorrhagic diarrhea and a poor feed conversion. For *B. intermedia* and *B.murdochii* the pathogenic potential is less clear-cut. Although both species have been isolated from clinical cases of diarrhea, the clinical symptoms are mild or absent in experimental infections and yet high numbers of spirochetes are necessary to cause an effect (Jensen and others 2004, Jensen and others 2010).

66 Recently, a new type of *Brachyspira* infection has been described. Outbreaks of 67 mucohaemorrhagic diarrhea, caused by strongly haemolytic *Brachyspira* strains inconsistent

with *B. hyodysenteriae*, were reported in the USA and Canada. Phylogenetic analysis of these
strains showed such a large genetic divergence between those isolates and all other *Brachyspira* spp. that these isolates likely represent a novel species, for which the name *"Brachyspira hampsonii"* has been proposed (Chander and others 2012). The current case
report describes, to the best of our knowledge, the first confirmed *"B. hampsonii"* infection in
pigs outside North-America.

74 Two gilts, imported from the Czech Republic, were presented for necropsy in a routine 75 quarantine monitoring protocol. General macroscopic findings consisted of a low body weight and dilated large intestines in which soft watery non-haemorrhagic colonic content was 76 77 present. Histological examination of these large intestines was not performed. Microbial culture of the colonic content was performed on Tryptic Soy Agar (BD, Heidelberg, 78 Germany) supplemented with 5% sheep blood (IMP, Brussels, Belgium), 0.1% yeast extract 79 80 (Oxoid, Aalst, Belgium) and following antimicrobials: spectinomycin (200µg/ml), spiramycin (25 µg/ml), rifampin (12.5 µg/ml), colistin (6.25 µg/ml), and vancomycin (6.25µg/ml) 81 82 (Hommez and others 1998). The microbial cultures revealed strongly haemolytic, ring phenomenon-positive spirochetes, indicative for B. hyodysenteriae (Fellström and others 83 1995, Hommez and others 1998). Some of the pigs, housed in the same group as the two gilts 84 85 presented for necropsy, showed mild diarrhea. From the next batch of gilts from the same origin, additional faecal samples were taken in the quarantine. Strongly haemolytic 86 Brachyspira isolates, with ring phenomenon, were again found on microbial culture, whereas 87 commercial diagnostic PCR analysis (Adiavet Brachy, Paris, France) did not confirm the 88 89 presence of B. hyodysenteriae in these samples. All faecal samples were negative for Salmonella. 90

91 Phenotypic characterisation tests were performed on pure cultures which were 92 obtained by at least three subcultures on Tryptic Soy Agar (TSA) plates supplemented with

5% defibrinated sheep blood and 0,1% yeast extract (Jenkinson and Wingar, 1981). 93 Phenotypic characterisation was performed on 4-day old cultures and was based on beta 94 haemolysis, indole production, hippurate hydrolysis and the presence or absence of α -95 galactosidase, α-glucosidase and β-glucosidase (Fellström and others 1995). Indole 96 production was determined using a spot-indole test (Remel BactiDrop, Dartford, UK) and for 97 the other biochemical characteristics commercial discs were used according to the 98 manufacturer's instructions (Rosco Diatabs, Taastrup, Denmark). Type strains of B. 99 hyodysenteriae (ATCC 27164), B. pilosicoli (ATCC 51139) and B.innocens (ATCC 29796) 100 were included to provide positive controls for all the phenotypic characteristics that were 101 examined. 102

103 Several species-specific PCRs were performed, based on the following genes: *tlyA* 104 (Råsbäck and others 2006), 23S rRNA (Leser and others 1997) and *nox* (Phillips and others 105 2006) for *B. hyodysenteriae, nox* (Phillips and others 2010) and 23s rRNA (Leser and others 106 1997) for *B. intermedia*, 16S rRNA (Phillips and others 2006) for *B. pilosicoli* and *nox* 107 (Atyeo and others 1999) for *B.murdochii/B.innocens*. Additionally, PCR's were performed for 108 the haemolysis related genes *hlyA* and *hlyA*-ACP (Barth and others 2012).

Forward primer 5' TAGCYTGCGGTATYGCWCTTT 3' and reverse primer 5' 109 GCMTGWATAGCTTCRGCATGRT 3' were used to partially sequence the nox gene 110 (Weissenböck and others 2005). A product of 1014 base pairs was obtained. Forward primer 111 5' GTTTGATYCTGGCTCAGARCKAACG 3' 5' and reverse primer 112 CTTCCGGTACGGMTGCCTTGTTACG 3' were used to partially sequence the 16S rRNA 113 gene of which a 1044 base pair product was obtained (Johansson and others 2004). 114 Sequencing reactions were performed on purified PCR-product with the same primers as for 115 PCR. Nox and 16S rRNA sequences from other Brachyspira isolates were retrieved from 116

117 GenBank and compared with the sequences of the described field case isolate (D52) by118 BLAST analysis.

The sequences of the *nox* gene of the strain retreived in this case report (D52), of *B. hyodysenteriae, B. intermedia, B. murdochii* and *B. innocens* ATCC type strains and of 42 additional strains of several *Brachyspira* spp. retrieved from GenBank were aligned using ClustalW. Sequences of clade I strain 30599 and clade II strain 30446 of *B. "hampsonii"* were also included (Rubin and others, 2013b). Phylogenetic analysis was performed with an alignment sequence fragment of 540 bp and Kimura distance calculation and neighbourjoining method were used.

126 For multilocus sequence typing (MLST) primers and PCR conditions as described by Råsbäck and others in 2007(b) were used to analyse genes encoding alcohol dehydrogenase 127 (*adh*). esterase (est). glutamate dehydrogenase (gdh),glucose kinase 128 (glpK),129 phosphoglucomutase (pgm) and acetyl-coA acetyltransferase (thi). For each locus the sequence obtained from the D52 isolate was matched with the online MLST database 130 (www.pubmlst.org/brachyspira). 131

The phenotypic characteristics of isolate D52 corresponded to those of *B. "hampsonii*" as described by Chander and others (2012). The isolate was strongly beta haemolytic, indole negative, hippurate negative, negative for α -galactosidase and α -glucosidase, and positive for β -glucosidase. Although not exclusively, most isolates of clade I are positive for β glucosidase as compared to clade II, in which most isolates are negative for β -glucosidase.

Table 1 shows the PCR results. Isolate D52 generated a positive result in the two species-specific PCRs for *B. intermedia* based on the 23S rRNA and *nox* gene respectively. Interestingly, the PCR targeting *tlyA*, presumed typical for *B. hyodysenteriae*, also generated a positive result. The PCRs for several haemolysis associated genes, *hlyA* and ACP(fabF-fabG), were positive as well.

The nox sequence of isolate D52 (GenBank accession nr KF202498) showed a 142 similarity of 100% over 547 basepairs with B. "hampsonii" isolate NSH-16, which is 143 described by Chander and others as the reference strain of B. "hampsonii" clade I (Chander 144 145 and others 2012). Besides, the nox sequence of isolate D52 showed a similarity of more than 99% over 874 basepairs with previously described isolates KC35 en EB106 (JX197410.1 and 146 JX197409.1) (Burrough and others 2012b). These isolates, originally described as strongly 147 haemolytic B. intermedia are recently refered to as "B. hampsonii clade I" (GenBank). With 148 149 the type strain B. hampsonii 30599 (clade I, NZ_AOMM01000255.1) as described by Rubin and others (2013b), the nox sequence of our isolate showed a similarity of 99% over 1014 bp 150 (difference of 2 nucleotides). The 16S rRNA sequence of isolate D52 (GenBank accession nr 151 KF586484) showed a sequence similarity of 99% over 1044 bp with "B. hampsonii" isolate 152 NSH-16. 153

Phylogenetic analysis of the *nox* sequence of isolate D52 and *nox* sequences of other Brachyspira spp. clearly place isolate D52 in the cluster of isolates comprising clade I of "*B*. *hampsonii*" (fig. 1)

As described for "*B. hampsonii*" in previous studies (Chander and others 2012) three of the seven loci for MLST could not be amplified. From the sequences of the 4 loci that could be amplified (*est, pgm, glp* and *thi*), none of them gave an exact match with known alleles in the MLST database. The *thi* sequence matched closest with allele 24 of "*Serpulina* sp. P280/1" (difference of 21 nucleotides) in accordance with the findings of Chander and others, 2012, for "*B. hampsonii*".

163 The results of the phenotypic characteristics, sequence comparisons, MLST and 164 phylogenetic analysis based on the *nox* sequence, identify the D52 isolate as "*B. hampsonii*" 165 clade I. To the best of our knowledge it is the first time that "*B. hampsonii*" isolates from

porcine origin are described in Europe, although the isolate *Serpulina* sp. P280/1 (Neef and
others 1994) in retrospect also may belong to "*B. hampsonii*".

The isolates of strain D52 obtained from the current field case, all contained the *hlyA*, 168 *tlyA* and ACP(*fabF*,*fabG*) genes. HlyA is the protein responsible for the strong haemolysis in 169 B. hyodysenteriae (Hsu and others 2001). In order to adequately perform its actions, the hlyA 170 gene has to be correctly placed between the accompanying fab-F and fab-G genes, coding for 171 an ACP-reductase and -synthetase (Zuerner and others 2004). Although the presence of hlyA 172 has been reported in some weakly haemolytic Brachysira spp. isolates, the fabF and fabG 173 genes were in those cases absent (Barth and others 2012), probably rendering the hlyA gene 174 functionally inactive. Another hemolysin, namely tlyA is consistently found in B. 175 hyodysenteriae. Although it has also been twice reported in weakly haemolytic species (Pati 176 and others 2010, Wanchanthuek and others 2010), these sequences show low sequence 177 178 similarity (82-83%) with tlyA of B. hyodysenteriae (Barth and others 2012). The presence of both these hemolysin encoding genes in the isolates in the current field case may be 179 responsible for the strong haemolysis displayed by these isolates. 180

Rubin and others (2013a) could experimentally induce mucohaemorrhagic diarrhea in 181 swine when infected with a "B. hampsonii" strain 30446. The clinical signs were 182 indistinguishable from swine dysentery. It should, however, be noted that the strain 30446 183 clearly falls into the cluster II isolates of "B. hampsonii" whereas the strain from this case 184 falls into cluster I as shown in the phylogenetic tree in figure 1. Although experimentally "B. 185 hampsonii" strain 30599, which belongs to clade I, can induce severe clinical symptoms 186 (Harding and others, 2013), the symptoms in this case report were rather mild. This could be 187 due to difference in pathogenic potential between strains of clade I or be related to 188 Brachyspira colitis being a multifactorial disease. Environmental or nutritional factors may 189 alter the severity of clinical signs. 190

This case report and the recent case reports from Canada and USA (Burrough and 191 others 2012a, Rubin and others 2013a) indicate that new, emerging species of Brachyspira 192 can be important in swine-rearing countries. The results of the species-specific PCRs show 193 that diagnosis of infections caused by these emerging species can be confusing. When 194 diagnosis is solely based on microbial culture, all strongly haemolytic isolates will be reported 195 as B. hyodysenteriae, whereas they could belong to the provisionally named species "B. 196 hampsonii", "B. suanatina" (Chander and others 2012, Burrough and others 2012a, Råsbäck 197 198 and others 2007a, Rubin and others 2013a) or even other emerging Brachyspira species. On the other hand, when diagnosis is entirely based on PCR, strongly haemolytic isolates 199 inconsistent with B. hyodysenteriae could easily be missed. For now, the combination of 200 microbial culture and PCR, complemented with sequencing if necessary, is presumably the 201 most complete method for diagnosis of *Brachyspira* spp. infections. 202

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204 **Declaration of conflicting interests**

Sources of financial support have been acknowledged and the authors declare that they haveno competing interests.

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 rearrangements and sequence drift distinguish *B. pilosicoli* from *B. hyodysenteriae*. *Anaerobe* 10, 229–237.
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330 **Tables and figures**

- 331 Table 1: Primers used in PCRs and results for B.hyodysenteriae reference strain ATCC 27164, B. intermedia
- reference strain ATCC 51140 and the field case isolate D52.

Target Gene	Species-specificity	Primer name	Primer Sequence (5'-3')	Result D 52	Result ATCC 27164 <i>B.hyodysente</i> <i>riae</i>	Result ATCC 51140 <i>B. intermedia</i>
hlyA	Non- specific	hlyAFo	TCG ATG AAA TTA AAG ATG TTG TT	positive	positive	positive
		hlyARe	TTT TTC TTG ATC TTC TTG AGG A			
ACP(fabF- fabG)	Non-specific	ACPFo	AGG IGA AGT IAT AGC IGT TGA CG	positive	positive	positive
		ACPRe	GAA ACA CCA TTA AGI AIA TTA TCC CA			
23S	B. hyodysenteriae	Hyo23SFo	CGG TAA GTG ATG TAC TTG	negative	positive	negative
		Hyo23SRe	AGC CTC AAC CTT AAA GA			
nox	B. hyodysenteriae	HyonoxFo	ACT AAA GAT CCT GAT GTA TTT G	negative	positive	negative
		HyonoxRe	CTA ATA AAC GTC TGC TGC			
tlyA	B. hyodysenteriae	tlyAFo	GCA GAT CTA AAG CAC AGG AT	positive	positive	negative
		tlyARe	GCC TTT TGA AAC ATC ACC TC			
nox	B. intermedia	IntnoxFo	AGA GTT TGA AGA CAC TTA TGA C	positive	negative	positive
		IntnoxRe	ATA AAC ATC AGG ATC TTT GC			
238	B. intermedia	Int23SFo	CCG TTG AAG GTT TAC CGT G	positive	negative	positive
		Int23SRe	CGC CTG ACA ATG TCC GG			
16S	B. pilosicoli	Pilo16SFo	AGA GGA AAG TTT TTT CGC TTC	negative	negative	negative
		Pilo16SRe	GCA CCT ATG TTA AAC GTC CTT G			
nox	B. innocens/ B. murdochii	Innmurdnox Fo	CCT GAA AGT TTA AAA GCT G	negative	negative	negative
		Innmurdnox Re	CGA TGT ATT CTT CTT TTC C			

- 334 Figure 1: Phylogenetic tree based on the alignment (540bp) of the nox gene of Brachyspira spp. The
- alignment was created using CLUSTALw, distance calculation (Kimura) and neighbour joining using PHYLIP.
- Bootstrap values are indicated. Scale bar indicates 0,02 substitutions per site.

