

1 Short Communication

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3 **Genetic relatedness of *Haemophilus parasuis* among reference strains and Chinese**
4 **epidemic isolates**

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24 **Abstract**

25 *Haemophilus parasuis* is the causative agent of Glässer's disease and a commensal
26 coloniser of the porcine upper respiratory tract. Multiple complex factors, including the early
27 weaning of piglets and the management of high health status farms, make it a re-emerging
28 agent, responsible for a recent increase in the prevalence and severity of disease in pigs in
29 China. However, little genetic information is known about Chinese epidemic isolates. In this
30 study, a phylogenetic method for genotyping the *H. parasuis* population with available
31 Chinese epidemic isolates and reference strains from different origins is presented.
32 Phylogenetic analysis confirmed that there are at least two different genotypes in *H. parasuis*
33 population and a unique Chinese lineage with virulence results in the previous epidemics.

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35 *Keywords:* *Haemophilus parasuis*; Phylogenetic analysis; Genotype; OMP type; Chinese
36 lineage

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38 *Haemophilus parasuis* can colonise the porcine upper respiratory tract and also cause
39 severe disease (Oliveira and Pijoan, 2004; Olvera et al., 2007). It is an important bacterial
40 pathogen of pigs in China, being responsible for substantial morbidity, mortality and
41 economic losses (Cai et al., 2005; Li et al., 2009). However, little is known about the genetic
42 relationship between Chinese epidemic isolates, those from other countries and the reference
43 strains. Thus, we performed a phylogenetic analysis of eighteen orthologous genes of strains
44 of *H. parasuis* from different geographical locations and obtained from healthy and disease
45 animals.

46
47 Eighteen orthologous genes (*gapA*, *ompP1*, *hscA*, *luxS*, *murG*, *pyrH*, *hlyX*, *ompP2*, *hemN*,
48 *dapA*, *queF*, *lpdA*, *ksgA*, *sodA*, HAPS_1299 (conserved hypothetical protein), *smtA*, *dapB* and
49 *ompP5*) were chosen from orthologous groups according to different functional categories.
50 These were submitted for phylogenomic analysis, along with orthologous genes from 18
51 complete genomes and two draft genome sequences available for *Pasteurellaceae* (M. Yue et
52 al., unpublished data). MEGA 4.0 was used to create Neighbour-Joining trees with interior
53 branch test (2000 replicates; seed = 80650) (Tamura et al., 2007). The strains and their
54 characteristics and primers used to amplify PCR products for DNA sequencing are detailed in
55 Supplementary Tables S1 and S2, respectively.

56
57 Phylogenetically, the strains branched into two main lineages (Fig. 1), indicating that
58 there were two independent genotypes present in the *H. parasuis* population examined. These
59 accorded with results from analysis of *H. parasuis* population by a multilocus sequencing

60 typing (MLST) method used by Olvera et al. (2006b); the only difference was that most of
61 their strains in analysis originated from Europe. Genotype A contained many virulent strains,
62 including a specific Chinese lineage associated with virulence, which also included a cluster
63 (independent sample t test, $P < 0.001$) containing strains that could not be typed according to
64 the previous serological procedure (F599, F641, F603, F663, F685, F687 and F593) (Cai et al.,
65 2005).

66

67 Most of non-typeable strains came from central China (F641, F603, F585 and F593). The
68 majority of Chinese epidemic isolates were composed of serovar 4, serovar 5 and
69 non-typeable strains; the relatively high numbers of non-typeable isolates may reflect the use
70 of serovar 4 and serovar 5 vaccines within China. The reference virulent strains (29755, HS80,
71 HS1080, HS50, HS1079 and HS1081) and another two Chinese clinical isolates (F042 and
72 F093) formed other virulent lineages but there was lack of convergence. However, the
73 American Strain 29755 and HS1080 and Japanese HS80 had a closer genetic relationship with
74 Chinese virulent lineage. The majority of epidemic strains (14/17) in the Chinese lineage
75 (independent sample t test, $P < 0.001$) indicated that previous *H. parasuis* in China had a
76 genetic specificity different from other virulent lineages.

77

78 A cluster containing strains considered to be of lesser virulence (isolated from nasal
79 cavities) formed a unique lineage (independent sample t test, $P < 0.001$), which included
80 avirulent strains (HS1077, HS1073, HS81 and HS1072), mid-virulent strains (HS83, HS79
81 and HS1065), two confirmed virulent strains (HS82 and HS1076) and an unclarified strain

82 HS145. Interesting, the nasal isolates strains HS82 and HS1076 confirmed their virulence in
83 animal challenge, which contradicts the knowledge that healthy nasal isolates were less
84 virulent (Aragon et al., 2009). However, genetic separation between nasal and virulent strains
85 had been observed previously (Olvera et al., 2006a and b).

86

87 While genotype A was more robust; genotype B had only two strains (HS1075 and
88 FGXBB), which had a different origin but shared a virulent phenotype. These were also
89 characterised as genotype 2 from analysis of the *H. parasuis* population by the MLST method,
90 which were mostly comprised of clinical disease isolates (24/30) (Olvera et al., 2006b).
91 Analysis of further strains is required to determine the robustness of genotype B.

92

93 In addition, we analysed the population structure of the *H. parasuis* strains by comparison
94 of the amino acid sequences of the major outer membrane proteins (OMPs) P1, P2 and P5 in
95 an analogous manner to that described recently (Mullins et al., 2009). The derived
96 phylogenetic tree is shown in Fig. 2. There was a clear separation between OMP type A and B
97 lineages and also a unique Chinese OMP profile within the type A lineage, which also
98 contained genetically closer strains 29755, HS1080 and HS80 (Fig. 1). In a comparison
99 between the two polygenetic trees, the low virulent lineage in Fig. 1 was separated as the two
100 OMP lineages in Fig. 2 (line in green), indicating that strains of nasal origin had more
101 heterogeneity in OMP profiles than their genetic profile. Although immune selection may
102 contribute to a variable OMP profile, a relatively convergent Chinese OMP profile
103 demonstrated the Chinese lineage (independent sample *t* test, $P < 0.001$).

104

105 Taken together, the data indicate that there are at least two dramatically different
106 genotypes in the *H. parasuis* population. Phylogenetic analysis of both conserved orthologous
107 genes and OMPs indicate there is a unique Chinese lineage, which is associated with a
108 virulent phenotype. Although the first complete genome sequence of Chinese lineage SH0165
109 gave valuable genetic information, more work needs to be performed targeting which
110 virulence genes result in disease outcome (Yue et al., 2009).

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112 **Conflict of interest statement**

113 None of the authors of this paper has a financial or personal relationship with other
114 people or organisations that could inappropriately influence or bias the content of the paper.

115

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119

120 **Appendix A. Supplementary material**

121 Supplementary Tables S1 and S2.

122

123 **References**

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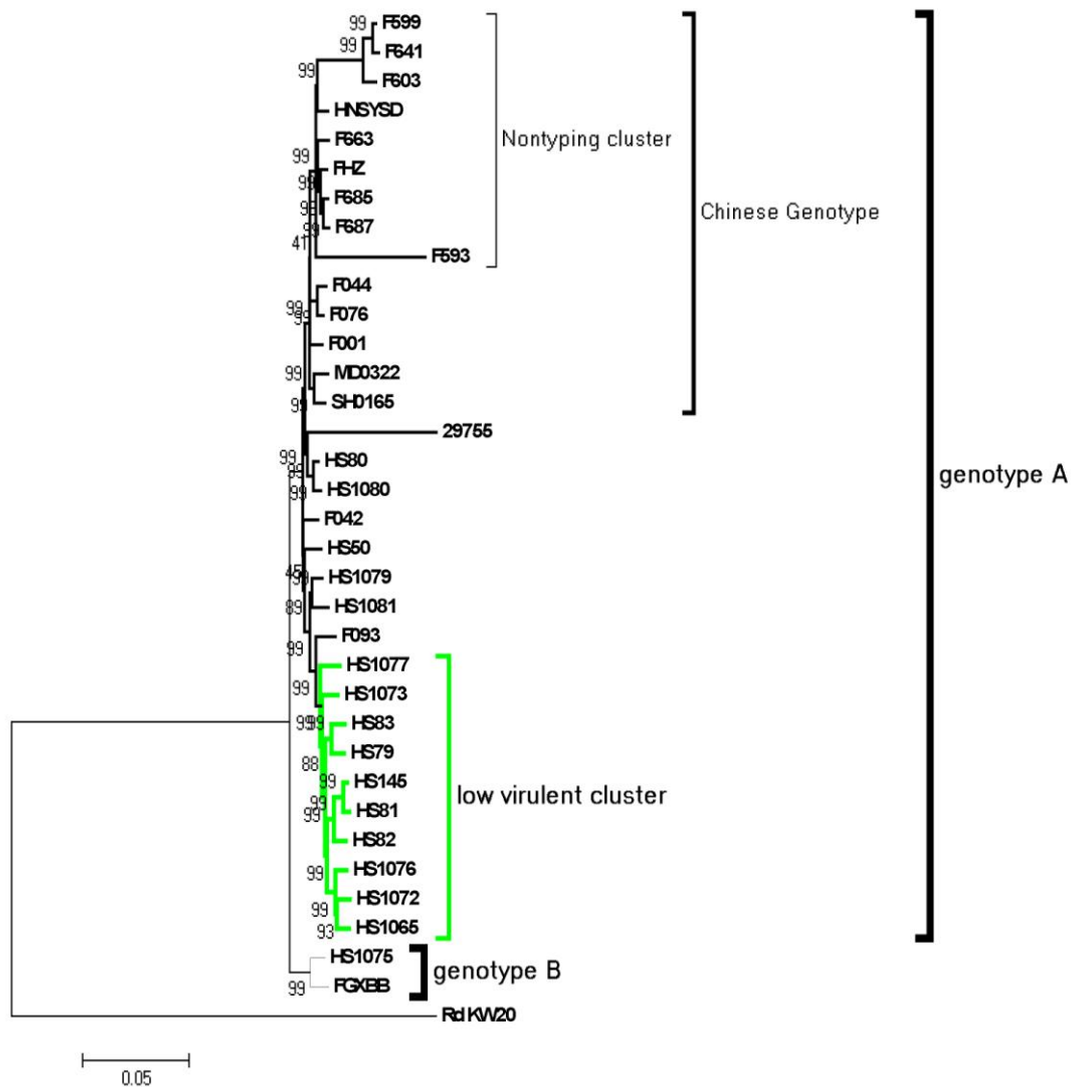
160 **Figure legends**

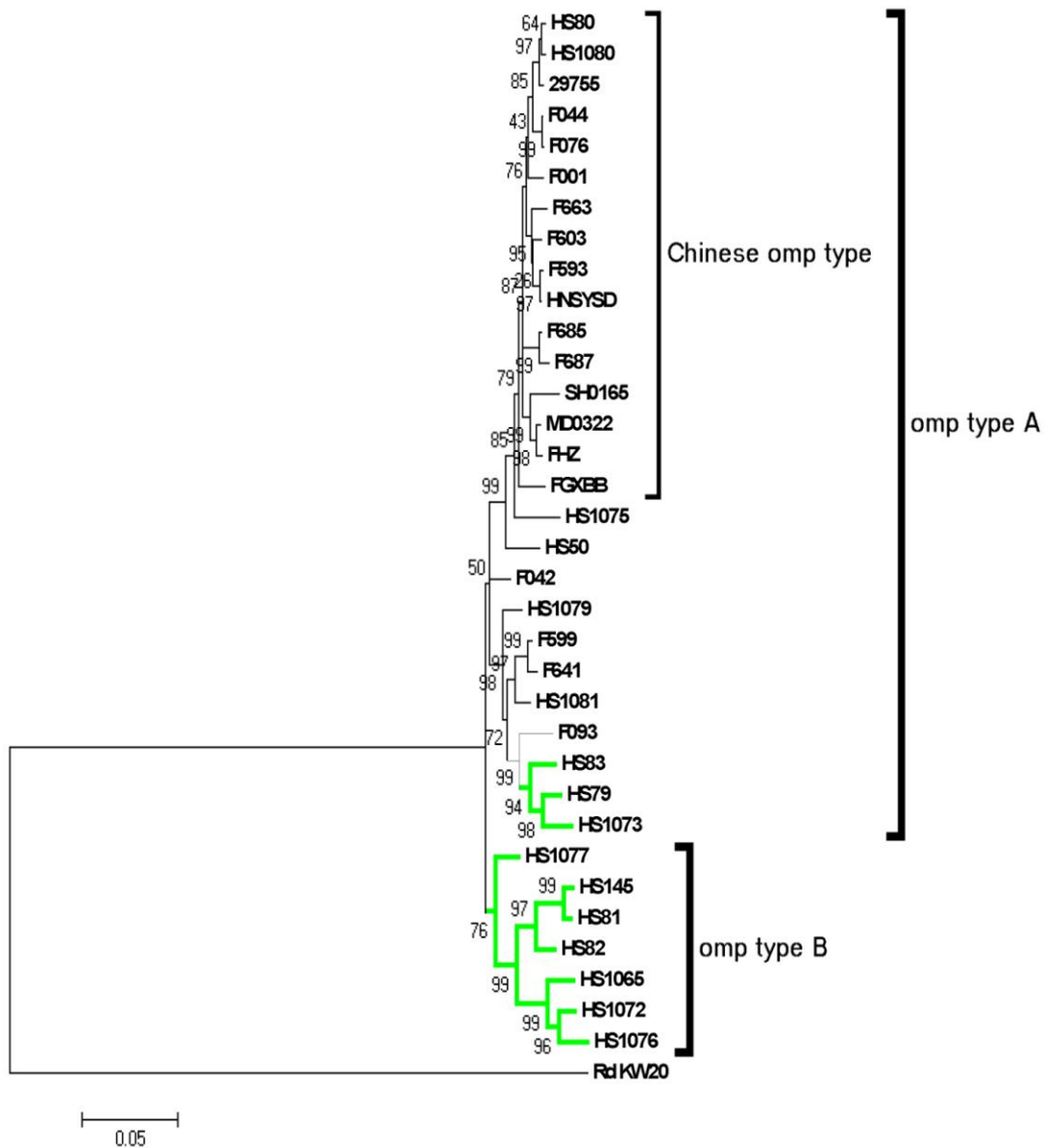
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162 Fig. 1. Neighbour-joining (NJ) tree with interior branch test of 2,000 replicates based on
163 concatenated nucleotide sequences of 18 conserved genes. Thirty-four *H. parasuis* strains
164 could be divided into two genotypes (genotypes A and B). The abundant genotype A
165 contained the Chinese genotype, including the non-typeable cluster and a low virulence
166 cluster (in green). Genotype B had only two strains. *H. influenzae* strain KW20 was used as
167 the reference for the construction of the phylogenetic tree. The NJ tree was produced using
168 MEGA 4.0. The scale bar is in unit of nucleotide substitutions per site. Bootstrap values (%)
169 are indicated at the branch nodes of the phylogenetic trees.

170

171 Fig. 2. Neighbour-joining (NJ) tree with interior branch test of 2,000 replicates based on
172 concatenated amino acid sequences of the P1, P2 and P5 proteins. Thirty-four *H. parasuis*
173 strains could be divided into two outer membrane protein (OMP) types or profiles (OMP
174 types A and B). OMP type A contained the Chinese OMP type. Green lines indicate nasal
175 isolates. *H. influenzae* strain KW20 was used as the reference for the construction of the
176 phylogenetic tree. The NJ tree was produced by MEGA 4.0. The scale bar is in units of amino
177 acid substitutions per site. Bootstrap values (%) are indicated at the branch nodes of the
178 phylogenetic trees.





1 **Supplementary Table S1**2 Strains and background information of *Haemophilus parasuis* used in the study.

3

KRG serovar^a	Date of receipt^b	Strain	Country of origin^c	Diagnosis/isolation site	Experimental Infection	Phenotype virulence^d	Reference
1	12/86	HS82	Japan	Healthy/nose	Glässer's disease	virulent	Amano et al., 1994
1	2/90	HS145	Australia	ND	ND	ND	Cai et al., 2005
2	12/86	HS83	Japan	Healthy/nose	Healthy	mid-virulent	Nielsen et al., 1993
3	12/86	HS81	Japan	Healthy/nose	Healthy	avirulent	Nielsen et al., 1993
4	12/86	HS79	Japan	Healthy/nose	Subclinical	mid-virulent	Amano et al., 1994
5	12/86	HS80	Japan	Septicaemia/meningitis	Glässer's disease	virulent	Amano et al., 1996
6	9/96	HS1072	Switzerland	Healthy/nose	Healthy	avirulent	Nielsen et al., 1993
7	9/96	HS1073	Switzerland	Healthy/nose	Healthy	avirulent	Nielsen et al., 1993
8	9/96	HS1065	Sweden	ND	Subclinical	mid-virulent	Kielstein et al., 1992
9	11/85	HS50	Sweden	ND	Healthy	avirulent	Kielstein et al., 1992
10	9/96	HS1076	Germany	Healthy/nose	Glässer's disease	virulent	Kielstein et al., 1992
11	9/96	HS1077	Germany	Pneumonia/trachea	Healthy	avirulent	Kielstein et al., 1992
12	9/96	HS1075	Germany	Polyserositis/lung	Glässer's disease	virulent	Kielstein et al., 1992
13	9/96	HS1079	USA	ND/lung	Glässer's disease	virulent	Kielstein et al., 1992
14	9/96	HS1080	USA	ND/joint	Glässer's disease	virulent	Kielstein et al., 1992
15	9/96	HS1081	USA	Pneumonia/lung	Polyserositis	virulent	Kielstein et al., 1992
5	ND	29755	USA	Pneumonia/lung	Glässer's disease	virulent	Nielsen et al., 1993
5	4/01	SH0165	Hubei, China	Polyserositis/lung	Glässer's disease	virulent	Cai et al., 2005
4	8/01	MD0322	Hebei, China	Polyserositis/lung	Glässer's disease	virulent	Cai et al., 2005
6	/03	F001	Hubei, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
10	/04	F042	Anhui, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
7	/04	F044	Henan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
5	/04	F076	Fujian, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
2	/04	F093	Hainan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/03	F593	Hubei, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005

NT	/04	F599	Shandong, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/03	F603	Henan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/04	F641	Henan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/04	F663	Shanghai, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/04	F685	Hunan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
NT	/02	F687	Hainan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
5	/04	FHZ	Zhejiang, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
4	/04	HNSYSD	Hunan, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005
5	/04	FGXBB	Guangxi, China	ND/lung	Glässer's disease	virulent	Cai et al., 2005

4

5 ^a There are 15 serovars according to immunodiffusion using heat-stable antigen extracts and NT was defined as the non-typable strains of *H. parasuis* by serotyping method
6 described by Cai. ND indicated the unknown information.

7 ^b Date expressed in month/year format.

8 ^c All the 17 Chinese field isolates were collected in 17 distinct farms from 11 provinces representing the Central China (SH0165, MD0322, F001, F044, F593, F603, F641),
9 East China (F042, F076, F599, F663, FHZ), South China (F093, F685, F687, HNSYSD, FGXBB), without disease correlation with available data.

10 ^d Strains were categorised as the virulent, mid-virulent and avirulent phenotype of *H. parasuis* according to the accompanying reference about experimental infection model.
11 All the 17 Chinese field isolates were provided by the study by Dr. Cai and the detail information about experimental infection with each strains as follows: 9-10 week old
12 weaning piglets (seronegative pigs, 5 for each strain as a group and another 5 for the negative control) were chosen for intraperitoneally challenging with 7×10^9 CFU, all
13 the infection groups, except for the negative control, showed clinical syndrome as Glässer's disease within a week, and 3~5 piglets in each group died in the following
14 weeks (Cai et al., unpublished data).

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1 **Supplementary Table S2**
 2 Primers used in this study.
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Homologous gene and locus ^a	Primers	Sequence	Expected Length (bp) ^b
<i>gapA</i>	gapA-F	ATGGCAATTA AAAATTGGTATTAAC	1020
HAPS_0001	gapA-R	TTAGCCTTTGTAGTTGTAAACGTG	
<i>hscA</i>	hscA-F	ATGTCATTACTTCAAATTGCAGAA	1677
HAPS_0059	hscA-R	TTAATTCCTCACCCGAAAGTAGT	
<i>luxS</i>	luxS-F	ATGCCTTTACTAGATAGCTTTAAAG	510
HAPS_0063	luxS-R	CTATGGATTTAGCAATTTCTCATCT	
<i>murG</i>	MurG-F	ATGACAAAAAAATTATTGGTAATGG	1056
HAPS_0119	MurG-R	TTACAAACTATTTTCCACAATCACT	
<i>pyrH</i>	pyrH-F	ATGAGCAATCCTATTTATAAACGTA	714
HAPS_0136	pyrH-R	TTAAGCAATCGTCGTACCTTC	
<i>hlyX</i>	hlyX-F	ATGAAAATTGTATCTGATTTTAAAGC	771
HAPS_0167	hlyX-R	TTATAAGTTTGGATTGCAATGGG	
<i>hemN</i>	hemN-F	ATGCAGCCCCCTTAAGC	1152
HAPS_0238	hemN-R	CTATTCCTTTAAAAAACCTTCCAAC	
<i>dapA</i>	dapA-F	ATGTCAAAACCTCTTTTTTCAGG	888
HAPS_0278	dapA-R	TTAGATTAATTGTGCTTTTTGTAATG	
<i>queF</i>	queF-F	ATGAATTACAATAATGAATGCTTTTC	840
HAPS_0318	queF-R	TTATTGTCTCACCATTCTTAAATTTT	
<i>lpdA</i>	lpdA-F	ATGAGCCAAGAAATTA AACACAA	1425
HAPS_0490	lpdA-R	TTATCTTTTCTTCGCTTTTGGAT	
<i>ksgA</i>	ksgA-F	ATGAGTTCAAATTCAAAAAACATTT	861
HAPS_0737	ksgA-R	TTATTCATCAAACAAGACTAGTTCTTT	
<i>sodA</i>	sodA-F	ATGGCATAACATTACCTGAGTTAG	621
HAPS_0815	sodA-R	TTATGCTTGGGATTCAAACCGT	
HAPS_1299	HAPS1299-F	ATGATTTACAGTATGACCGCTTTC	864
HAPS_1299	HAPS1299-R	CTACTCCAAATCTGAATTTGCTC	
<i>smtA</i>	smtA-F	ATGGGAACAAAGAATTGTTTCCTA	789
HAPS_1395	smtA-R	CTAATGAGATAGATATATATCAAACCGA	
<i>dapB</i>	dapB-F	ATGACATTA AAAAATTGGCGTTGT	813
HAPS_2274	dapB-R	TTATAAATTATTTAAATCCAACACATCG	
<i>ompP1</i>	ompP1-F	ATGAATAAATTTACTAAAACAGCACTT	1290
HAPS_0037	ompP1-R	TTAGAATTTGTAGTTTACGTTTAAGC	
<i>ompP2</i>	ompP2-F	ATGAAAAAACACTAGTAGCATTAGC	1092
HAPS_0164	ompP2-R	TTACCATAATACACGTAAACCAACA	
<i>ompP5</i>	ompP5-F	ATGAAAAAATCTTTAATTGCATTAGC	1116
HAPS_2298	ompP5-R	TTACATAGAACTTCTTTTGAACCTT	

4 ^a Gene name was assigned according to the *H. parasuis* genome SH0165. The homologous gene locus number is given below each
 5 gene name.

6 ^b Expected length of the PCR amplification product according to the homologous gene in SH0165 genome.