

Distribution of the CD4 Alleles in *Sus scrofa* Demonstrates the Genetic Profiles of Western Breeds and Miniature Pigs

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3 **Distribution of the *CD4* alleles in *Sus scrofa* demonstrates the genetic**
4 **profiles of Western breeds and miniature pigs**

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32

33 Running title: Distribution of *CD4* alleles in *Sus scrofa*

34 **Abstract**

35 Widely used anti-pig CD4 monoclonal antibodies (mAbs) fail to recognize CD4 alleles
36 characteristic of miniature pig lines such as NIH miniature pigs and microminipigs. We
37 surveyed polymorphisms in the coding sequence of the porcine *CD4* gene among
38 Western and Oriental pig breeds and Japanese wild boars and investigated their
39 distribution. Of the 13 alleles that we identified among the 47 animals, two in group I
40 and three in group II were found exclusively in Western breed pigs. Group IV alleles,
41 which included mAb-nonbinding alleles, were found frequently in Oriental breed pigs,
42 suggesting that the mAb-nonbinding allele arose from the gene pool of Oriental pigs.
43 Group IV alleles were also found in Duroc and Large White pigs, suggesting genetic
44 inflow from Oriental pig breeds into Western breeds. Comparison of the *CD4* sequences
45 of species in Cetartiodactyla suggested that the group IV alleles in *Sus scrofa* occurred
46 before the divergence of this species from the other artiodactyls. The different antibody
47 specificities of the various CD4 alleles may facilitate the discrimination of T-cell
48 populations in transplantation studies using miniature pigs. The significance of the
49 preservation of CD4 polymorphisms to immune function in pigs warrants further
50 investigation.

51

52 Keywords: allele frequency; CD4; genetic polymorphisms; miniature pig; swine

53 INTRODUCTION

54

55 The CD4 molecule characterizes various T-cell populations, including helper
56 and regulatory T cells (1). The CD4 molecule interacts with MHC class II molecules,
57 and most CD4-positive T cells are MHC-II restricted.

58 Miniature pigs, generated by crossbreeding of pigs with a small body size, have
59 become popular research models, given the similarity of their organ size and immune-
60 related genes to those of humans and their ease of handling. NIH miniature pigs, a
61 representative line of miniature pigs, have a characteristic *CD4* allele (*CD4.2*) that fails
62 to react with widely used anti-pig CD4 monoclonal antibodies (mAbs). Compared with
63 the major *CD4* allele (*CD4.1*), *CD4.2* has nucleotide substitutions within exons 3 and 4,
64 thus preventing the molecule from binding to the mAbs (2).

65 Microminipigs (MMPs) are a unique miniature pig line whose characteristic
66 extremely small body size (<10 kg in mature animals) provides cost benefits in terms of
67 husbandry and materials during preclinical testing (3). The major *CD4* allele in MMPs
68 (*CD4.A*; LC064059.1) shows good agreement with the pig *CD4* reference sequence
69 (NM_001001908.1). In addition, MMPs carry an alternative allele (*CD4.B*;
70 LC064060.1) that contains multiple altered bases in exons 3 and 4 and that is highly
71 similar to *CD4.2* in NIH miniature pigs (4). T cells derived from MMPs homozygous
72 for *CD4.B* fail to bind to currently available anti-pig CD4 mAbs, indicating that the
73 CD4.B molecule has lost the epitope recognized by the mAbs, again similar to CD4.2
74 (5). To clarify the origin and functional relevance of the mAb-nonbinding CD4 alleles,
75 we investigated the distribution of the *CD4* alleles in populations of pigs and wild boars
76 and analyzed the origin of the mAb-nonbinding *CD4* allele.

77

78 MATERIALS AND METHODS

79 Animals and Genomic DNA

80 We used 47 animals that had no obvious kinship with each other: 32 Western-
81 breed subjects (Berkshire, 8; Duroc, 8; Landrace, 8; Large White, 8), 8 pigs representing
82 Oriental breeds (Meishan, 4; Jinhua, 4), and 7 Japanese wild boars. Sperm, blood, and
83 tissue samples (tail ends, ear auricles, and kidney) were commercially purchased or
84 kindly provided by the Miyagi Livestock Experimental Station (Osaki, Miyagi, Japan);
85 the Chiba Prefecture Livestock Research Center (Yachimata, Chiba, Japan); the
86 Shizuoka Swine and Poultry Research Center (Kikugawa, Shizuoka, Japan); the
87 Tokushima Agriculture, Forestry, and Fisheries Technology Support Center (Kamiita,
88 Tokushima, Japan); and the Ibaraki Farm of the National Livestock Breeding Center
89 (Chikusei, Ibaraki, Japan). Samples for DNA extraction were obtained according to the
90 regulations regarding animal experiments at the Institute of Livestock and Grassland
91 Science, National Agriculture and Food Research Organization, and the Institute of
92 Japan Association for Techno-innovation in Agriculture, Forestry and Fisheries, and at
93 the institutes that provided the samples. Genomic DNA was obtained from sperm or
94 tissue samples by using phenol extraction followed by ethanol precipitation (6).

95

96 Sequencing

97 Primers were designed and used to PCR-amplify exons 2 through 10 of the
98 porcine *CD4* gene. For each reaction, 20 ng of genomic DNA was used as a template.
99 PCR amplification was performed by using AmpliTaq Gold polymerase (Thermo Fisher
100 Scientific, Waltham, MA, USA) in 35 cycles consisting of 94 °C for 30 s, 55 °C for 30
101 s, and 72 °C for 1 min after preincubation at 94 °C for 9 min. The amplified products
102 were kept at 4 °C until further use. Residual primers in the amplified products were
103 removed by treating with Exo-SAP IT solution (Affymetrix, Santa Clara, CA, USA)
104 according to the manufacturer's instructions. The products were sequenced with the

105 same primers used for the PCR amplification and BigDye Terminator v3.1 by using an
106 ABI 3730xl Genetic Analyzer (Thermo Fisher Scientific). Primers used in the PCR and
107 sequencing reactions are shown in Table 1. The sequences thus obtained were processed
108 as previously described (7). Polymorphisms were detected by using the PolyPhred
109 program (8). The pig *CD4* mRNA sequence (NM_001001908.1) was used as a
110 reference; it is 2715 bp long, comprises 11 exons, and contains a 1374-bp coding
111 sequence (CDS) corresponding to 457 amino acids. The locations of SNPs and the
112 exon–intron structure were determined by comparing the reference and the genomic
113 sequence of pig chromosome 5 (NC_010447.4). Phylogenetic analysis according to the
114 neighbor-joining method (9) was performed with ClustalX2 (10). Haplotypes of SNPs
115 within pig *CD4* genes were predicted by using PHASE with default parameters (11).

116

117 **RESULTS AND DISCUSSION**

118 We identified 40 SNPs, which were located in exons 3 through 9 within the CDS
119 of the reference pig *CD4* gene (Table 2). Of these, 14 SNPs were in exon 3, and 10 of
120 these SNPs were nonsynonymous. We then estimated 13 *CD4* alleles in the surveyed
121 animals according to computational reconstruction of the SNP haplotypes (Tables 3 and
122 4). The alleles with reconstructed haplotypes were derived from computational
123 estimations. Furthermore, some of the alleles (*CD4*02*, *CD4*05*, *CD4*08*, *CD4*09*,
124 and *CD4*13*) were not observed as homozygotes in the population thus examined. The
125 actual combination of SNPs in single alleles was difficult to confirm because of
126 chimerism between alleles during PCR amplification. However, because alleles that
127 were not found as homozygotes (except for *CD4*09* and *CD4*13*) were observed as
128 heterozygotes with other major alleles, we think that the estimated alleles are reliable.
129 All of the *CD4*09* and *CD4*13* alleles were found in combination with each other, thus
130 leading to uncertainty in their estimation; consequently the locations of these alleles in

131 the phylogenetic tree may be slightly inaccurate.

132 One of the 13 alleles, *CD4*06* was identical to *CD4.A*, which was the major
133 allele in MMPs. These alleles were classified into four groups through phylogenetic
134 analysis (Figure 1A). The most frequently observed allele, *CD4*01*, accounted for
135 approximately two-thirds of the alleles in Western breeds. *CD4*01*, as well as the other
136 alleles in group I including *CD4.1* of NIH miniature pigs, had the identical sequence to
137 the reference *CD4* sequence in the extent of *CD4.1*; *CD4*01* was therefore considered
138 to be the representative allele in Western breeds (Figure 1B). Group II alleles were
139 common in Western breeds and shared several characteristic nucleotides in exons 3, 4,
140 5, and 8 with group I alleles; overall their similarity to group I alleles was higher than
141 that of other groups (Table 3 and Figure 1A). Alleles in groups I and II were found
142 exclusively in Western breed pigs in this study. Duroc pigs examined here did not have
143 any group II alleles. Although our small sample size might have accounted for this
144 finding, the genetic population size of the Duroc breed may be smaller than that for
145 other Western breeds, such as Landrace and Large White (12), and pigs with group II
146 alleles might have had no, or very limited, contribution to the establishment of the
147 Duroc breed. The representative alleles of Oriental breeds composed a particular clade
148 in group III, including *CD4.A/CD4*06*. *CD4* alleles derived from Japanese wild boars
149 were also included in group III (Table 4 and Figure 1).

150 Groups I through III of pig *CD4* alleles were highly similar to each other within
151 the region covering exon 3. However, the remaining *CD4* alleles, group IV, had many
152 polymorphic sites within the CDR2-like regions of exons 3 and 4 (13), compared with
153 those in the other groups (Table 3). Group IV included *CD4.2* and *CD4.B*, which are
154 regarded as characteristic alleles in miniature pigs, as well as alleles from Oriental
155 breeds: *CD4*12* in Jinhua pigs and *CD4*13* in Meishan pigs. Furthermore, *CD4*12*,
156 which comprised three-eighths of the alleles of Jinhua pigs, also occurred in Duroc and

157 Large White pigs, albeit at low frequency (Table 4). The deduced amino-acid sequence
158 *CD4*12* was completely identical to those of *CD4.2* and *CD4.B* in the region
159 represented in the registered *CD4.2* sequence (X65630.1). Taken together, our findings
160 indicate that the mAb-nonbinding pig *CD4* alleles that were first noted in miniature pig
161 lines occur widely throughout various pig populations, including representative Oriental
162 and Western breeds.

163 We revealed that the atypical *CD4* alleles (group IV) were not specific to
164 miniature pig lines but occurred relatively frequently in Oriental breeds and
165 occasionally in some Western breeds, thus showing good agreement with the genetic
166 influx from Oriental breeds into Western breeds, such as Duroc (14). Oriental-breed
167 pigs are frequently used as founders for miniature pig lines: for example, the generation
168 of MMPs included Vietnamese Pot-bellied pigs (3). The relatively high frequency of
169 group IV *CD4* alleles in Oriental breeds and the influence of these pigs in the
170 establishment of Western breeds and miniature pig lines suggest that the group IV
171 alleles originated in Oriental-breed pigs. However, none of the wild boars that we
172 analyzed contained any group IV *CD4* alleles, suggesting that the origin of the group IV
173 alleles might have been restricted to animals in a particular area of Asia. In addition,
174 cannot rule out the possibility that the group IV alleles in Duroc and Large White breeds
175 were derived from minor alleles in European wild boars. The existence and frequency of
176 group IV alleles in Japanese and European wild boars should be investigated to answer
177 this question.

178 The CDR2-like region of *CD4*, which corresponds to the boundary of exons 3
179 and 4, protrudes to the outer side of the molecule (13). This region of the human *CD4*
180 molecule is required for binding to gp120 of the human immunodeficiency virus (15,
181 16), and its interaction with MHC class II molecules is debated (17, 18). Although
182 polymorphisms in the CDR2-like region might be hypothesized to influence T-cell

183 function, studies have failed to reveal any difference in immunologic traits between pigs
184 with typical and group IV *CD4* alleles (2, 5). The CDR2-like region of CD4 is
185 monomorphic in most species (13); however, among species of the order
186 Cetartiodactyla, the CD4 sequences corresponding to polymorphic sites in the CDR2-
187 like region of porcine CD4 showed high rates of polymorphism. Furthermore, four of
188 the amino acids of group IV CD4 at the 10 polymorphic sites between deduced amino
189 acid sequences of *CD4.2* and the pig *CD4* reference sequence are identical to the
190 residues in other Cetartiodactyla species, suggesting that the CD4.2/CD4.B allele arose
191 before the divergence of pigs from other artiodactyls (Figure 2).

192

193 **CONCLUSIONS**

194 Here, we have shown the distribution of *CD4* alleles—particularly those arising
195 from polymorphisms in CDR2-like regions—in several *Sus scrofa* populations. The
196 functional influence of these polymorphisms remains unclear, but the differences in
197 antibody specificity among these *CD4* alleles may facilitate the discrimination of T-cell
198 subpopulations in transplantation studies using miniature pigs. The significance of the
199 preservation of polymorphisms of porcine CD4 in terms of their interaction with MHC
200 molecules and other immune functions warrants further investigation.

201

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207

208 **Figure legends**

209

210 **Figure 1.** Phylogeny of pig *CD4* alleles. Sequences corresponding to (A) the complete
211 coding sequences and (B) the regions determined in a previous study (X65629.1 and
212 X65630.1; positions 99 to 397 in the coding sequence of NM_001001908.1) (13) were
213 analyzed by using the neighbor-joining method (9). Identical sequences were analyzed
214 as single sequences. The full-length coding sequence of pig *CD4* (NM_001001908.1),
215 as well as alleles in the MMP population and in NIH miniature pigs, was included in the
216 phylogenetic analysis. The coding sequence of cattle *CD4* (NM_001103225.1) was used
217 as an outgroup. Bootstrap trials were conducted 10,000 times, and the bootstrap values
218 are shown at the nodes. The lengths of branches with omissions are indicated in
219 parentheses.

220

221 **Figure 2.** Alignment of *CD4* sequences of Cetartiodactyla species corresponding to the
222 polymorphic region between the pig reference sequence and group IV *CD4* alleles. The
223 deduced amino-acid sequences of the porcine *CD4.2* allele (X65630.1) and the
224 corresponding regions of the pig reference sequence (NP_001001908.1) and of the *CD4*
225 proteins of representative Cetartiodactyla species (*Bos taurus*, NP_001096695.1; *Ovis*
226 *aries*, NP_001123374.1; *Capra hircus*, ACG76115.1; *Orcinus orca*, XP_004279107.1;
227 *Tursiops truncatus*, NP_001267583.1) were aligned. Triangles indicate polymorphic
228 sites between the pig reference sequence and *CD4.2*; fringed letters indicate amino acids
229 in non-pig species that are identical to those of the polymorphic sites in porcine *CD4.2*;
230 dots indicate residues identical to the pig reference sequence; and dashes indicate
231 deletions.

232

233

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- 286

Table 1 Primers sequences and procedures for PCR of the porcine *CD4* gene and preparation of sequencing templates.

Target exon	Location (chromosome 5) ^a	Sequence (5'→3')	Amplicon (bp)
2	66,346,741–66,346,718	GTACCTGTGGGTGTCAGTTTAGAG	383
	66,346,359–66,346,382	CTTACCCAGCACCAGATATTTTC	
3	66,346,515–66,346,492	CTCAGACTCAAACCTGGGATGATTG	366
	66,346,150–66,346,173	GATCCCAGAGTTTACTAGGAGCTG	
4	66,333,536–66,333,512	AATGAGCAACTCAGATCAGAAGAGT	363
	66,333,174–66,333,195	CTTATCCATCTCTGGACGGTTG	
5	66,332,952–66,332,931	CTTCTCCTGGGGATAGTGCAT	358
	66,332,595–66,332,615	ACACTACAGCCACGAGCAGAG	
6	66,331,799–66,331,775	GCCTAGAGCTAGATGGGAATTTAAG	617
	66,331,183–66,331,204	GATTCCAGCCTCAGTTCAAACC	
7	66,329,886–66,329,862	CTTTAGAGCAGACAAGTGCTAGGAA	373
	66,329,514–66,329,537	ACCATACCCATAACCCACTGACTC	
8	66,328,935–66,328,912	AGCATAAGGATCAGACCCAAGTGT	400
	66,328,536–66,328,559	TAACTCTGTGGCTTCTTGTCTCTC	
9	66,328,539–66,328,516	GTTAATTCTGGGACAGATGGCTTC	238
	66,328,302–66,328,323	CTCTCTTACCCCTCCTCTTTG	
10	66,328,153–66,328,132	CCATCTCTGTGCAGGAAAAGTC	698
	66,327,456–66,327,476	AGCTGAGCTGCTTGGGTGATA	
10	66,327,556–66,327,537	ACTGACGGAGCCACAGACTC	668
	66,326,889–66,326,908	GGCTATCAACTTTCGCAGGA	

Table 2 SNPs in the porcine *CD4* gene.

Exon ^a	Pig genome		Coding sequence		Frequency of the reference allele in each breed (%) ^d							
					Western breeds				Oriental breeds			Overall (47)
	Position ^a	Allele ^b	Position ^c	Amino acid	LR (8)	LW (8)	DR (8)	BS (8)	MS (4)	JH (4)	WB (7)	
3	66,346,335	G A	126	Syn	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,306	C T	155	P→L	100.0	100.0	100.0	100.0	100.0	100.0	0.0	85.1
	66,346,300	A G	161	N→S	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,291	A G	170	N→S	93.7	100.0	75.0	100.0	25.0	37.5	100.0	83.0
	66,346,286	A G	175	Syn	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,279	C T	182	T→I	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,268	G A	193	G→S	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,266	C G	195	Syn	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,265	G A	196	G→S	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,260	T C	201	Syn	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
	66,346,259	G C	202	G→R	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6
66,346,255	G A	206	S→N	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6	
66,346,251	C G	210	F→L	100.0	93.8	93.8	100.0	75.0	62.5	100.0	92.6	
66,346,243	C A	218	T→K	50.0	56.3	93.8	43.8	75.0	62.5	0.0	53.2	
4	66,333,448	C T	231	Syn	100.0	100.0	100.0	100.0	50.0	100.0	100.0	95.7
	66,333,447	G C	232	E→K	100.0	100.0	100.0	100.0	75.0	100.0	42.9	89.4
	66,333,440	C G	239	T→S	50.0	56.3	93.8	43.8	100.0	62.5	100.0	70.2
5	66,332,898	A G	385	T→A	100.0	100.0	100.0	100.0	50.0	100.0	100.0	95.7
	66,332,728	T A	555	Syn	43.7	62.5	75.0	43.7	50.0	100.0	0.0	51.1
	66,332,683	T C	600	Syn	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0 ^e
6	66,331,624	C T	665	A→V	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0 ^e
	66,331,602	C T	687	Syn	100.0	100.0	100.0	68.8	100.0	100.0	100.0	94.7
	66,331,599	G A	690	Syn	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0 ^e
	66,331,420	C T	869	P→L	56.3	43.8	31.3	56.3	100.0	100.0	100.0	63.8
	66,331,411	C T	878	A→V	100.0	93.8	100.0	100.0	100.0	100.0	100.0	98.9
	66,331,410	G A	879	Syn	93.7	100.0	75.0	100.0	25.0	37.5	57.1	76.6
	66,331,380	T C	909	Syn	56.3	43.8	31.3	56.3	100.0	100.0	100.0	63.8
	66,331,341	G A	948	M→I	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0 ^e
7	66,329,815	C T	953	A→V	50.0	56.3	100.0	43.8	100.0	100.0	100.0	74.5
	66,329,814	G T	954	Syn	100.0	100.0	100.0	100.0	25.0	100.0	42.9	85.1
	66,329,781	T G	987	Syn	0.0	6.2	6.2	0.0	0.0	37.5	0.0	5.3
	66,329,724	G C	1044	M→I	56.3	43.8	31.3	56.3	100.0	100.0	100.0	63.8
	66,329,707	A G	1061	Q→R	6.3	6.3	31.3	0.0	25.0	100.0	42.9	24.5
	66,329,619	C T	1149	Syn	50.0	62.5	100.0	43.8	100.0	100.0	100.0	75.5
8	66,328,791	A G	1185	Syn	93.7	93.7	68.7	100.0	0.0	0.0	0.0	60.6
	66,328,790	G A	1186	E→K	93.7	93.7	68.7	100.0	0.0	0.0	0.0	60.6
	66,328,777	C T	1199	S→F	93.7	93.7	68.7	100.0	0.0	0.0	0.0	60.6
	66,328,726	C T	1250	A→V	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0 ^e
	66,328,713	C T	1263	Syn	93.7	93.7	68.7	100.0	0.0	0.0	0.0	60.6
9	66,328,348	G A	1335	Syn	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0 ^e

Bases different from those in the reference sequence (NM_001001908.1) were regarded as SNPs. BS, Berkshire; DR, Duroc; JH, Jinhua; LR, Landrace; LW, Large White; MS, Meishan; Syn, synonymous; WB, Japanese wild boar

^a Positions of SNPs on the pig genome are indicated relative to p-ter of chromosome 5 (NC_010447.4).

^b The first allele in each pair is identical to the reference sequence (NM_001001908.1).

^c Exons and positions of SNPs in the coding sequence are shown relative to the nucleotide sequence NM_001001908.1.

^d The number of animals for each breed is indicated in parentheses.

^e Monomorphic in the animals used in this study.

Table 3. Alleles of pig *CD4* genes inferred by SNP haplotypes in the coding sequence.

Group	Allele	Position in the coding sequence of the reference pig <i>CD4</i> mRNA (NM_001001908.1)																																										
		Exon 3										Exon 4			Exon 5			Exon 6				Exon 7				Exon 8			Exon 9															
		126	155	161	170	175	182	193	195	196	201	202	206	210	218	231	232	239	385	555	600	665	687	690	869	878	879	909	948	953	954	987	1044	1061	1149	1185	1186	1199	1250	1263	1335			
I	Pig RefSeq <i>CD4</i> (NM_001001908.1)	G	C	A	A	A	C	G	C	G	T	G	G	C	C	C	G	C	A	T	T	C	C	G	C	C	G	T	G	C	G	T	G	A	C	A	G	C	C	C	G			
	<i>CD4.1</i> (X65629.1)
	<i>CD4*01</i>	C	T	.	A	T	.	.	C	.	.	.	G	C	G	T	.	.	.	A			
	<i>CD4*02</i>	C	T	.	A	T	.	.	C	.	T	.	G	C	G	T	.	.	.	A			
II	<i>CD4*03</i>	A	.	.	G	.	A	C	T	.	A	T	.	G	.	G	T	.	.	.	T	.	.	A			
	<i>CD4*04</i>	A	.	.	G	.	A	C	T	T	A	T	.	G	.	G	T	.	.	.	T	.	.	A			
	<i>CD4*05</i>	A	.	.	G	.	A	C	T	.	A	.	T	.	.	.	T	.	G	.	G	T	.	.	.	T	.	.	A			
III	<i>CD4.A</i> (LC064059.1) / <i>CD4*06</i>	.	.	.	G	T	.	.	G	.	C	T	.	A	.	.	A	.	.	T	G	.	G	.	G	A	T	T	T	.	A				
	<i>CD4*07</i>	.	.	.	G	C	T	.	A	.	.	A	.	.	.	G	.	.	.	G	A	T	T	T	.	A				
	<i>CD4*08</i>	.	.	.	G	A	C	T	.	A	.	.	A	.	.	.	G	.	.	.	G	A	T	T	T	.	A				
	<i>CD4*09</i>	A	A	C	T	.	A	.	.	A	.	.	.	G	.	.	.	G	A	T	T	T	.	A				
	<i>CD4*10</i>	.	T	A	A	C	T	.	A	.	.	A	.	.	.	G	.	.	.	G	A	T	T	T	.	A				
	<i>CD4*11</i>	.	T	A	.	A	.	.	A	C	T	.	A	T	G	.	G	.	G	A	T	T	T	.	A				
IV	<i>CD4.2</i> (X65630.1)	A	.	G	.	G	T	A	G	A	C	C	A	G	A	.	.	G		
	<i>CD4.B</i> (LC064060.1)	A	.	G	.	G	T	A	G	A	C	C	A	G	A	.	.	G	.	.	C	T	.	A	A	G	A	T	T	T	A		
	<i>CD4*12</i>	A	.	G	.	G	T	A	G	A	C	C	A	G	A	.	.	G	.	.	C	T	.	A	G	A	T	T	T	.	A		
	<i>CD4*13</i>	A	.	G	G	G	T	A	G	A	C	C	A	G	.	.	A	.	.	A	C	T	.	A	T	G	.	G	.	G	A	T	T	T	.	A			

Nucleotides that differ from the reference pig *CD4* mRNA sequence (NM_001001908.1) are shown; dots indicate nucleotides identical to the reference. Nonsynonymous substitutions leading to changes in amino acids (see Table 2) are bolded. Alleles detected in NIH miniature pigs (*CD4.1* and *CD4.2*) (13) and MMPs (*CD4.A* and *CD4.B*) (5) are included also; the determined sequences of the alleles in NIH miniature pigs correspond to positions 99 through 397 of the coding sequence of pig *CD4* (NM_001001908.1).

Table 4. Pig *CD4* alleles in each population.

Group	Allele	Western breeds (<i>n</i> = 8 each)				Oriental breeds (<i>n</i> = 4 each)		Japanese wild boar (<i>n</i> = 7)	Total alleles
		Berkshire	Duroc	Landrace	Large White	Meishan	Jinhua		
I	<i>CD4*01</i>	7	11	7	8				33
	<i>CD4*02</i>				1				1
II	<i>CD4*03</i>	4		8	5				17
	<i>CD4*04</i>	5							5
	<i>CD4*05</i>				1				1
III	<i>CD4*06</i>					4			4
	<i>CD4*07</i>						5		5
	<i>CD4*08</i>		4	1					5
	<i>CD4*09</i>					2			2
	<i>CD4*10</i>							6	6
	<i>CD4*11</i>							8	8
IV	<i>CD4*12</i>		1		1		3		5
	<i>CD4*13</i>					2			2
Total alleles		16	16	16	16	8	8	14	94

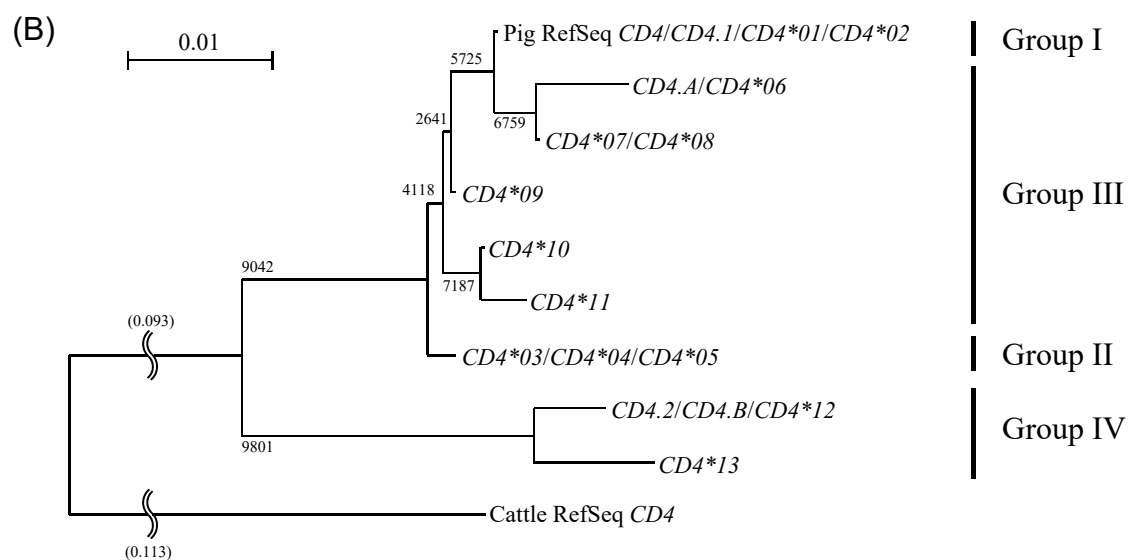
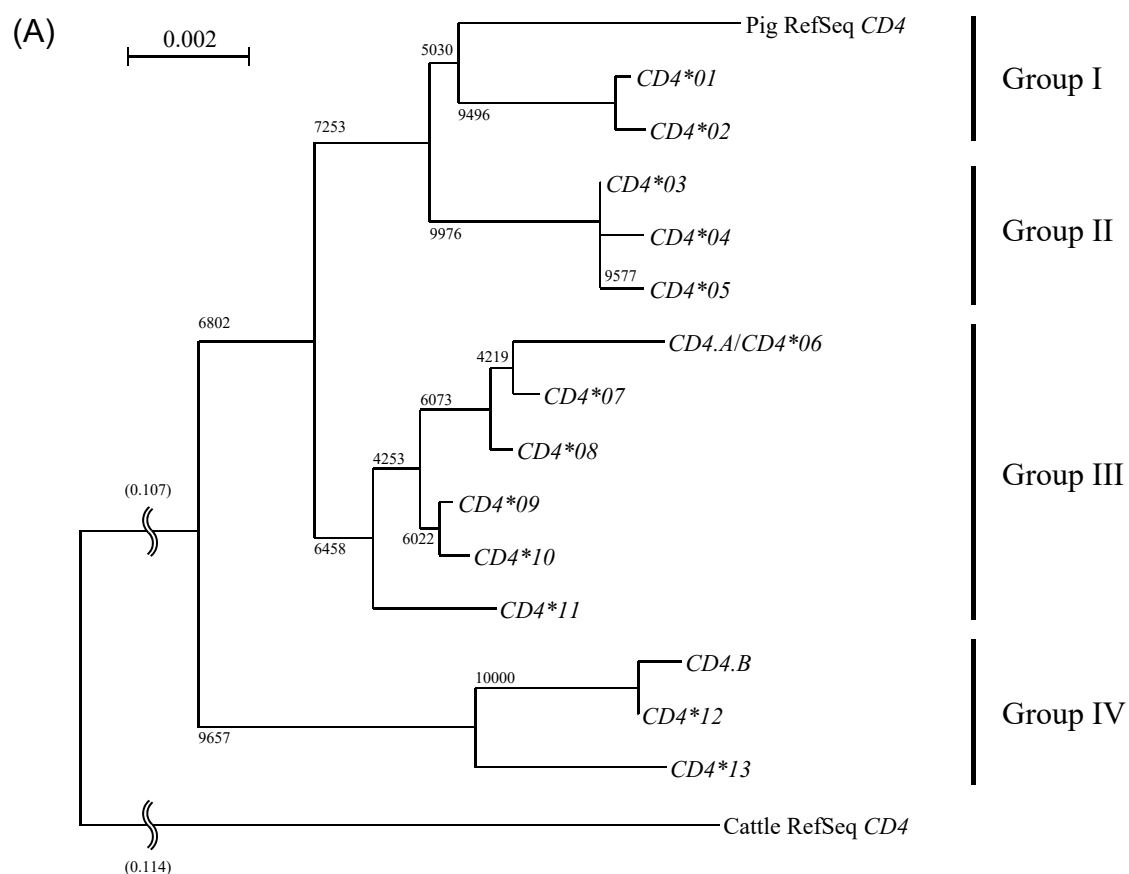


Figure 1

