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Exploiting phenotype diversity in a local animal genetic resource: Identification of a single nucleotide polymorphism associated with the tail shape phenotype in the autochthonous Casertana pig breed

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9	Exploiting phenotype diversity in a local animal genetic resource: identification of a single
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11	Casertana pig breed
12	
13	Francesca Bertolini ^{1,2,*} , Giuseppina Schiavo ^{1,*} , Silvia Tinarelli ^{1,3} , Laura Santoro ⁴ , Valerio Joe
14	Utzeri ¹ , Stefania Dall'Olio ¹ , Leonardo Nanni Costa ¹ , Maurizio Gallo ³ and Luca Fontanesi ¹
15	
16	¹ Department of Agricultural and Food Sciences, Division of Animal Sciences, University of Bologna,
17	Viale Fanin 46, 40127 Bologna, Italy
18	² Department of Bio and Health Informatics, Technical University of Denmark, Kemitorvet, Building
19	208, Room 007, 2800 Kgs. Lyngby, Denmark
20	³ Associazione Nazionale Allevatori Suini, Via L. Spallanzani 4, 00161 Roma, Italy
21	⁴ ConSDABI - National Focal Point Italiano FAO, Contrada Piano Cappelle, 82100 Benevento, Italy
22	
23	Corresponding author: Luca Fontanesi, e-mail: luca.fontanesi@unibo.it
24	
25	* These authors contributed equally to this work.
26	

Running title: SNPs and tail shape phenotype in pigs

28 Highlights

- Casertana is an autochthonous pig genetic resource reared in Central-South of Italy.
- Tail shape phenotype variability in the breed was investigated in a GWAS.
- A single nucleotide polymorphism on porcine chromosome 12 was associated with this trait.
- This marker is close to the *SRY-box* 9 (*SOX9*) gene that is essential in skeletogenesis.

33 Abstract

34 Casertana is a local pig breed mainly raised in Central-South regions of Italy. Pigs of this breed 35 are considered the descendants of the ancient Neapolitan population that largely influenced the 36 constitution of the modern commercial pigs. The pigs of this breed are usually curly-tailed, like several other domestic pig populations. However, Casertana population shows some variability for 37 38 this trait, including animals having strait tail as observed in wild boars. In this study, we run, for the 39 first time, a genome wide association study (GWAS) comparing the curly-tailed (no. = 53) and strait-40 tailed (no. = 19) Casertana pigs to identify genomic regions associated with the tail shape phenotype 41 in Sus scrofa. All animals were genotyped with the Illumina PorcineSNP60 BeadChip v.2. GEMMA 42 software was used in the GWAS for which we were able to correct for stratification in the analysed cohort. A single nucleotide polymorphism (rs81439488), located on porcine chromosome 12, was 43 44 significantly associated with the investigated trait. This marker is close to the SRY-box 9 (SOX9) gene 45 that encodes for a transcription factor that is required during sequential steps of the chondrocyte 46 differentiation pathway, notochord maintenance and skeletogenesis. As the shape of the tail could be 47 important in relation to the problem of tail biting in pigs, the obtained results might open new perspectives for defining selection programs answering indirectly animal welfare issues. This work 48 49 demonstrated that autochthonous animal genetic resources might be used to disclose genetic factors 50 affecting peculiar traits by exploiting segregating phenotypes and genetic variability.

51

52 Keywords: Animal genetic resource; autochthonous breed; GWAS; morphological trait; SNP; *Sus* 53 *scrofa*.

54 **1. Introduction**

55 Conservation of animal genetic resources is mainly aimed to preserve genetic diversity and 56 associated inheritable phenotypes characterizing different populations that might be interesting for 57 current or future purposes, including potential use in breeding programs. These resources can be also 58 useful to understand biological mechanisms determining unique phenotypes derived by diversity in 59 selection pressures or as result of adaptation to environmental and production conditions (Leroy et 60 al., 2016).

61 Casertana pigs constitute a local breed mainly raised in Central-South regions of Italy. Pigs of 62 this breed are considered the descendants of the ancient Neapolitan pig population that largely 63 influenced the constitution of the modern commercial pig breeds through introgression of blood into British pig populations during the 19th century (Porter, 1993). Neapolitan pigs were, in turn, 64 influenced by Asian blood in the late 18th century (Porter, 1993). Casertana is enlisted among the 65 66 endangered animal genetic resources as the herd book of this breed accounts for about 100 boars and 67 sows currently registered (ANAS, 2016). Animals are mainly raised in extensive or semi-extensive 68 production systems with possible contacts and crossbreeding with European wild boars that could 69 have contributed, at least in part, to shape their morphological characteristics. Casertana pigs have a 70 black or grey coat colour, wrinkled skin, forward ears, and usually a typical hairless phenotype. The 71 pigs of this breed are usually curly-tailed, like several other domestic pig populations. However, 72 Casertana population shows some variability for this trait, including animals having strait and wavy 73 tail as in a few other pig breeds and in wild boars.

Domestication in mammals has been a complex and continuous process associated with a series of changes in the domesticated animals compared to the wild counterparts, derived by selective breeding of animals showing favourable production and reproduction performances, and increased docility that indirectly shaped the genome of domesticated populations (Wiener and Wilkinson, 2011; Larson and Burger, 2013; Carneiro et al., 2014; Wang et al., 2014; Wilkins et al., 2014). Several morphological features have been also directly or indirectly selected and, in most cases, fixed in

domesticated populations as result of the domestication process (Darwin, 1868). Coat colour is one of the most common phenotypic traits that has been modified as result of reduced selective pressure against colours with low fitness in the wild and of aesthetic preferences of the breeders, sometimes associated with higher production performances (Clutton-Brock, 1999). Among several other morphological characters, curliness of the tail and shape has been associated with domestication in mammals (Trut et al., 2009).

The tail is considered an extension of the spinal column usually composed of specifically shaped vertebrae. Spontaneous curly tail phenotypes in mice have been the matter of studies that investigated the role of embryonic development in this morphological anomaly (Copp et al., 1988; van Straaten and Copp, 2001; Ohnishi et al., 2017). Curly tail is also commonly observed in many dog breeds. Vaysse et al. (2011) compared the genome of dog breeds having curly tails with that of breeds with straight tails using single nucleotide polymorphisms (SNPs) chip data and identified a genomic region on chromosome 1 significantly associated with these alternative tail shapes.

93 In pigs, few studies have been reported on the genetic factors affecting tail shape. A putative 94 recessive genetic defect known as kinky tail (or flexed or screw tail), derived by fused caudal vertebrae associated in some cases with other defects, has been described in the mid of the last century 95 96 (Nordby, 1934; Donald, 1949; Brooksbank, 1958). It is not known if this defect could be, in some 97 way, related or not to the normal curling of the tail that is common in domestic pigs. This signature 98 of domestication, however, seems not fixed in all pig breeds (Porter, 1993) but no systematic study 99 has been conducted so far, probably because the difficulties in retrieving phenotype information due 100 to the usual practice of tail docking in most herds.

In this study, we took advantage from the variability of the shape of the tail that we recorded in the Casertana pig population and run a genome wide association study (GWAS) comparing the genome of curly-tailed and strait-tailed animals to identify genomic regions associated with the tail shape phenotype in *Sus scrofa*.

106 **2. Materials and methods**

107 **2.1. Animals**

A total of 101 Casertana pigs (of about 7 to 20 months old) from six different farms were 108 109 evaluated. Photographic records of each animal were obtained to capture information on the tail shape in standardized restraining conditions (including a direct evaluation of the personnel on this 110 111 phenotype during this phase for the biological sampling) for all animals and after release (Figure 1). 112 Pigs were classified as follows: 53 (25 males and 28 females) showed the curly tail phenotype; 19 113 (five males and 14 females) showed the strait-tail phenotype; 29 were not classified and excluded 114 from the study as tail docking, that was practised by the farmers as routine before weaning of the 115 piglets, prevented the recording of any tail phenotype.

116

117 **2.2. Genotyping**

118 Hairs (with roots) were collected from the investigated pigs. DNA extraction was carried out 119 using the Wizard® Genomic DNA Purification kit (Promega Corporation, Madison, WI, USA) 120 following the manufacturer's instructions. Genotyping of the extracted DNA was obtained with the 121 Illumina PorcineSNP60 BeadChip v.2 (Illumina, Inc., San Diego, CA, USA) that interrogated 61,565 SNPs. Single nucleotide polymorphisms were assigned to the Sscrofa11.1 genome version, as 122 123 previously described (Fontanesi et al., 2012). PLINK 1.9 software (Chang et al., 2015) was used to 124 filter SNPs and genotyping data using the following criteria already used in a similar study (Schiavo 125 et al., 2018): genotyping call rate >0.9, minor allele frequency >0.01 and Hardy-Weinberg 126 equilibrium *P*>0.001.

127

128 **2.3. Data analysis and genome wide association**

To evaluate distance relationships among the animals of the investigated cohort,
multidimensional scaling (MDS) was obtained with the PLINK 1.9 software (Chang et al., 2015).
Genome wide association study was carried out by applying the univariate mixed model of GEMMA

132 (Zhou and Stephens, 2012) that can accommodate the centered relatedness matrix calculated from 133 SNP genotypes to correct for population stratification in a case and control analysis. The model also 134 included the farm and the sex as fixed effects. To be able to identify associated markers in this 135 experiment that included a low number of animals (derived by the fact that the analysed pigs were 136 almost a complete representation of the whole population of the Casertana breed) and that used a SNP 137 chip that might originally have an ascertain bias (as local breeds were not used for the selection of 138 the informative SNPs), the significant threshold was defined at the $P_{nominal value} < 5.00 \text{E-}05$ level, 139 according to the Wellcome Trust Case Control Consortium (2007) and as also applied in several other 140 GWAS in livestock (e.g. Fontanesi et al., 2012; Sanchez et al., 2014). Genomic inflation factor (λ) 141 and quantile-quantile (Q-Q) plot were obtained with GenABEL (Aulchenko et al., 2007). Gene 142 annotation information was retrieved from the Sscrofall.1 genome version available at the Ensembl 143 database (http://www.ensembl.org/Sus_scrofa/Info/Index), release 91.

144

145 **3. Results and discussion**

146 A recent phenotypic characterization of the endangered Casertana pig population that we 147 carried out noted several morphological differences among distinct animals of this autochthonous 148 breed (data not shown). For example, in addition to the hairless or hypotrichotic condition (that is the 149 characteristic phenotype of the Casertana animals), we already described the presence of haired pigs 150 in this population and this morphological variability was used for a GWAS that we have recently 151 reported (Schiavo et al. 2018). Despite a limited number of animals was included in that study, we 152 were able to identify genomic regions associated with the hairless phenotype, demonstrating that local 153 animal genetic resources can be used to genetically describe phenotypic variability of simple traits 154 (Schiavo et al., 2018).

Another morphological trait that is not fixed in this breed is the shape of the tail (Figure 1). Of the animals for which we could record this phenotype, 26% (19 out of 72) showed a strait tail without any curls, similarly to the usual shape of wild boars. This shape was clearly different from the curly tails reported in the remaining investigated pigs (74%). There was no effect of the age and the sex did not exclusively explain the observed phenotype as both sexes were included in the two phenotype groups. In addition, we could exclude the possible effect of the behavioral change of tail posture on this phenotype (Zonderland et al., 2009). The recording system was based on standardized conditions and subsequent photographic records of the animals confirmed their assignment to one or to the other group of tail shape phenotype. The two groups were observed in animals from all six farms. Limited pedigree record prevented the possibility to evaluate any potential founder effect.

A total of 36,533 autosomal SNPs, mapped to a unique position in the Sscrofa11.1 genome version, was used for MDS. The obtained MDS plot showed some structures not well defined in the analysed pigs that however did not clearly separate the curly and strait tailed Casertana pigs (Figure 2). A stratified sample could be a critical point in GWAS in a very small population where, to some extent, all animals might be related. Figure S1 reports the genomic inflation factor (λ) and Q–Q plot that did not show any biased test statistic distribution, suggesting that the investigated cohort was corrected for a possible stratification effect.

Figure 3 reports the Manhattan plot obtained in this GWAS. One significant SNP (P=2.3E-05) was identified on porcine chromosome 12 (SSC12). This marker indicated as ALGA0064877 (rs81439488) is located at position 10,301,075 of this chromosome.

175 One of the closest annotated gene in this desert chromosome region is the SRY-box 9 (SOX9) 176 gene (positions 8,641,629-8,647,764, encoded by the -1 strand), that, according to its function, might 177 be the most plausible candidate gene, explaining the recorded phenotypic variability. It is well 178 established that the expression of this gene at the embryonal level marks the onset of cartilage 179 differentiation (Wright et al., 1995; Healy et al., 1996). SOX9 encodes for a transcription factor that 180 is required during sequential steps of the chondrocyte differentiation pathway, notochord 181 maintenance and skeletogenesis (Akiyama et al., 2002; Barrionuevo et al., 2006; Montero et al., 182 2017). Continued expression of Sox9 in differentiated chondrocytes is essential for subsequent 183 hypertrophy and sustains chondrocyte-specific survival mechanisms (Ikegami et al., 2011).

Heterozygous mutations within and around human *SOX9* cause campomelic dysplasia that is a malformation syndrome characterized by cartilage derived skeletal structure defects (Foster et al., 1994; Wagner et al., 1994). These mutations, most of which reduce the level of expression of this gene, are located upstream spanning a large region (from 50 kb to more than 1 Mb) in which regulatory elements are present (Wunderle et al., 1998; Bagheri-Fam et al., 2006). Close upstream mutations produce more severe defects whereas far upstream mutations cause mild defects (Pfeifer et al., 1999; Velagaleti et al., 2005; Leipoldt et al., 2007).

191 Based on these studies in other species it is tempting to suggest a possible regulatory mechanism 192 affecting SOX9 expression in porcine developing chondrocytes that would, in turn, produce a mild 193 cartilage/skeletal effect determining the shape of the tail. This hypothesis might be worth of further 194 investigation starting from a precise characterization of the structure and morphology of the pig tail 195 with different shapes for which, at present, there is no detailed investigation. Our phenotype records 196 were based only on an external morphological evaluation of the shape of the tail. Furthermore, 197 analysis of gene expression of SOX9 at different developmental stages should be also carried out to 198 evaluate the role of this gene in the phenotype observed in pigs.

199 The results we obtained might have broader impacts than those that would be limited to a simple 200 morphological characterization. The shape of the tail could be important in relation to the problem of 201 tail biting in pigs. Tail biting is a widespread behavioral vice with significant animal welfare 202 implications and economic losses in commercial pig farms (Bracke et al., 2004). A few studies have 203 established correlations between tail posture and tail biting incidence suggesting limited damages and 204 related welfare complications with behaviors of the pigs that tended to have a tail posture up that 205 those with tail posture down (Zonderland et al., 2009; Lahrmann et al., 2017). It would be interesting 206 to evaluate if pigs with genetically determined curly tails (as a possible adaptation derived by the 207 domestication process) are less affected by tail biting damages than pigs with strait tails.

208

209 **4.** Conclusions

210 This work demonstrated that autochthonous animal genetic resources, even constituted by very small populations, might be used to disclose genetic factors affecting peculiar traits by exploiting 211 212 segregating phenotypes and genetic variability. To our knowledge, this is the first study that reported 213 a frequency distribution of the tail shape phenotype in a pig population. Our results indicated that this 214 morphological trait is associated with a marker close to an important gene involved in embryonic 215 development, opening other hypothesis, worth of further investigations. It will be important to 216 validate the results we obtained in this GWAS in other breeds and populations, including a more 217 precise anatomical characterization of this trait, to further extend the impact of the results reported in 218 Casertana pigs. It would be however first needed to know if diversity for this morphological 219 characteristic is common in commercial pig populations as at present, there is not information on this 220 aspect, mainly due to the usual practice of tail docking that prevents the recording of this phenotype. 221 Considering the potential relationship between tail shape and tail biting damages (that, however, 222 remains to be formally demonstrated), it could be possible to envisage practical applications of the 223 identified marker in selection programs aimed to respond to animal welfare issues. Our study 224 represents one of the few examples of exploitation of animal genetic resources to recover information 225 that might have potential impacts in commercial populations.

226

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The authors declare that there is no conflict of interest regarding the publication of this paper.

237

238 Conflict of interest statement

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- 240

241 **References**

- Akiyama, H., Chaboissier, M.C., Martin, J.F., Schedl, A., de Crombrugghe, B. 2002. The
 transcription factor Sox9 has essential roles in successive steps of the chondrocyte
 differentiation pathway and is required for expression of Sox5 and Sox6. Genes Dev. 16, 2813–
 2828.
- 246 ANAS, 2016. Registro Anagrafico. http://www.anas.it/.
- Aulchenko, Y.S., Ripke, S., Isaacs, A., van Duijn C.M., 2007. GenABEL: an R library for genomewide association analysis. Bioinformatics 23, 1294–1296.
- Bagheri-Fam, S., Barrionuevo, F., Dohrmann, U., Günther, T., Schüle, R., Kemler, R., Mallo, M.,
 Kanzler, B., Scherer, G. 2006. Long-range upstream and downstream enhancers control distinct
 subsets of the complex spatiotemporal Sox9 expression pattern. Dev. Biol. 291, 382–397.
- Barrionuevo, F., Taketo, M. M., Scherer, G., Kispert, A. 2006. Sox9 is required for notochord
 maintenance in mice. Dev. Biol. 295, 128–140.
- Bracke, M.B.M., Hulsegge, B., Keeling, L., Blokhuis, H.J. 2004. Decision support system with
 semantic model to assess the risk of tail biting in pigs. 1. Modelling. Appl. Anim. Behav. Sci.
 87, 31–44.
- 257 Brooksbank, N.H. 1958. Congenital deformity of the tail in pigs. Br. Vet. J. 114, 50–55.
- 258 Carneiro, M., Rubin, C.J., Di Palma, F., Albert, F.W., Alföldi, J., Martinez Barrio, A., Pielberg, G.,
- 259 Rafati, N., Sayyab, S., Turner-Maier, J., Younis, S., Afonso, S., Aken, B., Alves, J.M., Barrell,
- 260 D., Bolet, G., Boucher, S., Burbano, H.A., Campos, R., Chang, J.L., Duranthon, V., Fontanesi,

261	L., Garreau, H., Heiman, D., Johnson, J., Mage, R.G., Peng, Z., Queney, G., Rogel-Gaillard,
262	C., Ruffier, M., Searle, S., Villafuerte, R., Xiong, A., Young, S., Forsberg-Nilsson, K., Good,
263	J.M., Lander, E.S., Ferrand, N., Lindblad-Toh, K., Andersson, L. 2014. Rabbit genome analysis
264	reveals a polygenic basis for phenotypic change during domestication. Science 345, 1074-
265	1079.
266	Chang, C.C., Chow, C.C., Tellier, L.C., Vattikuti, S., Purcell, S.M., Lee, J.J. 2015. Second-generation
267	PLINK: rising to the challenge of larger and richer datasets. Gigascience 4, 7.
268	Clutton-Brock, J. 1999. A Natural History of Domesticated Mammals. 2nd Edition, Cambridge, UK:
269	Cambridge University Press.
270	Copp, A. J., Brook, F. A., & Roberts, H. J. 1988. A cell-type-specific abnormality of cell proliferation
271	in mutant (curly tail) mouse embryos developing spinal neural tube defects. Development 104,
272	285–295.
273	Darwin, C. 1868. The Variation of Animals and Plants under Domestication. London, UK: John
274	Murray.
275	Donald, H. P., 1949. The inheritance of a tail abnormality associated with urogenital disorders in pigs.
276	J. Agric. Sci. 39, 164–173.
277	Fontanesi, L., Schiavo, G., Galimberti, G., Calò, D.G., Scotti, E., Martelli, P.L., Buttazzoni, L.,
278	Casadio, R., Russo, V. 2012. A genome wide association study for backfat thickness in Italian
279	Large White pigs highlights new regions affecting fat deposition including neuronal genes.
280	BMC Genomics 13, 583.
281	Foster, J.W., Dominguez-Steglich, M.A., Guioli, S., Kwok, C., Weller, P.A., Stevanović, M.,
282	Weissenbach, J., Mansour, S., Young, I.D., Goodfellow, P.N., et al. 1994. Campomelic
283	dysplasia and autosomal sex reversal caused by mutations in an SRY-related gene. Nature 372,
284	525-530.

Healy, C., Uwanogho, D., Sharpe, P.T. 1996. Expression of the chicken Sox9 gene marks the onset
of cartilage differentiation. Ann. N. Y. Acad. Sci. 785, 261–262.

- Ikegami, D., Akiyama, H., Suzuki, A., Nakamura, T., Nakano, T., Yoshikawa, H., Tsumaki, N. 2011.
 Sox9 sustains chondrocyte survival and hypertrophy in part through Pik3ca-Akt pathways.
 Development 138, 1507–1519.
- Lahrmann, H. P., Hansen, C. F., D'Eath, R., Busch, M. E., & Forkman, B. (2017). Tail posture
 predicts tail biting outbreaks at pen level in weaner pigs. Appl. Anim. Behav. Sci. doi:
 10.1016/j.applanim.2017.12.006.
- Larson, G., Burger, J. 2013. A population genetics view of animal domestication. Trends Genet. 29,
 197–205.
- Leipoldt, M., Erdel, M., Bien-Willner, G.A., Smyk, M., Theurl, M., Yatsenko, S.A., Lupski, J.R.,
- Lane, A.H., Shanske, A.L., Stankiewicz, P., Scherer, G. 2007. Two novel translocation breakpoints upstream of SOX9 define borders of the proximal and distal breakpoint cluster region in campomelic dysplasia. Clin. Genet. 71, 67–75.
- Leroy, G., Besbes, B., Boettcher, P., Hoffmann, I., Capitan, A., Baumung, R. 2016. Rare phenotypes
 in domestic animals: unique resources for multiple applications. Anim. Genet. 47, 141–153.
- 301 Montero, J.A., Lorda-Diez, C.I., Francisco-Morcillo, J., Chimal-Monroy, J., Garcia-Porrero, J.A.,
- Hurle, J.M. 2017. Sox9 expression in Amniotes: Species-specific differences in the formation
 of digits. Front. Cell Dev. Biol. 5, 23.
- 304 Nordby J.E. 1934. Kinky tail in swine. J. Hered. 25, 171–174.
- Ohnishi, T., Miura, I., Ohba, H., Shimamoto, C., Iwayama, Y., Wakana, S., Yoshikawa, T. 2017. A
 spontaneous and novel Pax3 mutant mouse that models Waardenburg syndrome and neural tube
 defects. Gene 607, 16–22.
- 308 Pfeifer, D., Kist, R., Dewar, K., Devon, K., Lander, E.S., Birren, B., Korniszewski, L., Back, E.,
- 309 Scherer, G. 1999. Campomelic dysplasia translocation breakpoints are scattered over 1 Mb
- 310 proximal to SOX9: evidence for an extended control region. Am. J. Hum. Genet. 65, 111–124.
- 311 Porter, V. 1993. Pigs: A Handbooks to the Breeds of the World. Cornell University Press,

- 312 Sanchez M.P., Tribout T., Iannuccelli N., Bouffaud, M., Servin, B., Tenghe, A., Dehais, P., Muller,
- 313 N., Del Schneider, M.P., Mercat, M.J., Rogel-Gaillard, C., Milan, D., Bidanel, J.P., Gilbert, H.
- 314 2014. A genome-wide association study of production traits in a commercial population of
- Large White pigs: evidence of haplotypes affecting meat quality. Genet. Sel. Evol. 46, 12.
- 316 Schiavo, G., Bertolini, F., Utzeri, V.J., Ribani, R., Geraci, C., Santoro, L., Óvilo, C., Fernández, A.I.,
- Gallo, M., Fontanesi, L. 2018. Taking advantage from phenotype variability in a local animal
 genetic resource: identification of genomic regions associated with the hairless phenotype in
 Casertana pigs. Anim. Genet., in press.
- 320 Trut, L., Oskina, I., Kharlamova, A. 2009. Animal evolution during domestication: the domesticated
 321 fox as a model. Bioessays 31, 349–360.
- van Straaten, H. W., Copp, A. J. 2001. Curly tail: a 50-year history of the mouse spina bifida model.
 Anat. Embryol. 203, 225–238.
- 324 Vaysse, A., Ratnakumar, A., Derrien, T., Axelsson, E., Rosengren Pielberg, G., Sigurdsson, S., Fall,
- 325 T., Seppälä, E.H., Hansen, M.S., Lawley, C.T., Karlsson, E.K.; LUPA Consortium, Bannasch,
- 326 D., Vilà, C., Lohi, H., Galibert, F., Fredholm, M., Häggström, J., Hedhammar, A., André, C.,
- 327 Lindblad-Toh, K., Hitte, C., Webster, M.T. 2011. Identification of genomic regions associated
- with phenotypic variation between dog breeds using selection mapping. PLoS Genet. 7,e1002316.
- Velagaleti, G.V., Bien-Willner, G.A., Northup, J.K., Lockhart, L.H., Hawkins, J.C., Jalal, S.M.,
 Withers, M., Lupski, J.R., Stankiewicz, P. 2005. Position effects due to chromosome
 breakpoints that map approximately 900 Kb upstream and approximately 1.3 Mb downstream
 of SOX9 in two patients with campomelic dysplasia. Am. J. Hum. Genet. 76, 652–662.
- 334 Wagner, T., Wirth, J., Meyer, J., Zabel, B., Held, M., Zimmer, J., Pasantes, J., Bricarelli, F.D., Keutel,
- 335 J., Hustert, E., Wolf, U., Tommerup, N., Schempp, W., Scherer, G. 1994. Autosomal sex
- reversal and campomelic dysplasia are caused by mutations in and around the SRY-related gene
- 337 SOX9. Cell 79, 1111–1120.

- Wang, G.D., Xie, H.B., Peng, M.S., Irwin, D., Zhang, Y.P. 2014. Domestication genomics: evidence
 from animals. Annu. Rev. Anim. Biosci. 2, 65–84.
- Wiener, P., Wilkinson, S. 2011. Deciphering the genetic basis of animal domestication. Proc. Biol.
 Sci. 278, 3161–3170.
- Wilkins, A. S., Wrangham, R. W., Fitch, W. T. 2014. The "domestication syndrome" in mammals: a
 unified explanation based on neural crest cell behavior and genetics. Genetics 197, 795–808.
- Wright, E., Hargrave, M.R., Christiansen, J., Cooper, L., Kun, J., Evans, T., Gangadharan, U.,
 Greenfield, A., Koopman, P. 1995. The Sry-related gene Sox9 is expressed during
 chondrogenesis in mouse embryos. Nat. Genet. 9, 15–20.
- Wunderle, V.M., Critcher, R., Hastie, N., Goodfellow, P.N., Schedl, A. 1988. Deletion of long-range
 regulatory elements upstream of SOX9 causes campomelic dysplasia. Proc. Natl. Acad. Sci
 USA 95, 10649–10654.
- Zhou, X., Stephens, M. 2012. Genome-wide efficient mixed-model analysis for association studies.
 Nat. Genet. 44, 821–824.
- 352 Zonderland, J.J., van Riel, J.W., Bracke, M.B.M., Kemp, B., den Hartog, L.A., Spoolder, H.A.M.,
- 2009. Tail posture predicts tail damage among weaned piglets. Appl. Anim. Behav. Sci. 121,
 165–170.

Figure 1. Tail shape of Casertana pigs: a) curly tail; b) strait tail.







Figure 2. Multidimensional scaling (MDS) with represented the pigs (dots) included in this study
divided in the two groups of tail shape.



C1

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Figure S1. Quantile–quantile (Q–Q) plot obtained from the genome wide association analysis.



Lambda = 0.91