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Authentication of "mono-breed" pork products: Identification of a coat colour gene marker in Cinta Senese pigs useful to this purpose

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1	Authentication of "mono-breed" pork products: identification of a coat colour gene marker in
2	Cinta Senese pigs useful to this purpose
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14	Running title: A DNA marker useful for pork product authentication
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16 Highlights

- Cinta Senese is a local Italian pig breed with a belted coat colour
- The value of its meat production chain should be protected from frauds
- We analyzed the *KIT* gene to identify a suitable DNA marker for this purpose
- A single nucleotide polymorphism (g.43597545C>T) was almost fixed in Cinta Senese
- Probability to correctly assign an unknown meat sample to Cinta Senese was ~1.00.

22 Abstract

The possibility to authenticate food products is crucial to defend local livestock production chains 23 from frauds. Cinta Senese is an autochthonous pig breed reared under extensive or semi-extensive 24 management systems, mainly in the Tuscany (Italy). A Protected Designation of Origin (PDO) 25 brand for Cinta Senese meat was recently obtained. The breed is characterised by a typical black 26 with a white-belted coat colour pattern. We analyzed a coat colour gene (KIT) to identify a DNA 27 28 marker that could be useful for Cinta Senese meat product authentication. An informative single 29 nucleotide polymorphism (SNP) was identified among different KIT gene haplotypes that were obtained from several pigs of different breeds. This SNP (g.43597545C>T; position on porcine 30 chromosome 8 in the Sscrofa10.2 genome assembly) was genotyped by PCR-restriction fragment 31 length polymorphism (RFLP) in 631 animals of 11 different pig breeds and one wild boar 32 population. Allele T was almost fixed in Cinta Senese (95.9%) and absent in many breeds and was 33 34 considered the tag SNP of the belted allele. Probability to correctly assign an unknown meat sample to Cinta Senese was 0.97-1.00. This DNA marker can be useful to distinguish meat of Cinta Senese 35 pigs from meat of non-belted pigs. Thus, it could be an important tool not only to defend Cinta 36 37 Senese pork chain from frauds but also to design breeding plans to eliminate non belted alleles from this pig population. 38

39

40 **Keywords:** SNP; coat colour gene; *KIT*; traceability; authenticity; pig breed

41 **1. Introduction**

42 The identification of the origin and the authentication of food products are important issues to defend livestock production chains from frauds that produce consumer distrust and undermine 43 commercial valorisation of many local and niche products (Montowska and Pospiech, 2012). 44 Among these products, an increasing interest during the last few years has been directed to the 45 development of "mono-breed" labelled lines of meat and dairy products (Fontanesi, 2009). The 46 marketing link between a breed and its products is positively considered by the consumers in terms 47 of perceived quality and contributes to improve profitability as the products are sold at a higher 48 price compared to undifferentiated ones. This link is mainly used to improve the economic incomes 49 50 derived by local and endangered breeds that are usually less productive. The market added value is important for a sustainable exploitation of rural economies and is the fundamental driver for the 51 conservation of endangered animal genetic resources (Fontanesi, 2009; Hoffmann, 2011). 52

53 Cinta Senese is an autochthonous pig breed that is reared under extensive or semi-extensive management systems, mainly in the Tuscany region (Italy). The breed is characterised by a typical 54 55 black with a white belted coat colour pattern. Its origin dates back to the XIII-XIV century when belted pigs were raised in the hills around Siena as demonstrated by a famous painting of Ambrogio 56 Lorenzetti in the Palazzo Comunale of Siena (a.D. 1340). The importance of this breed was 57 58 recognized with the early constitution of the breed national herdbook that worked from the 1936 to 1966 and then by a regional herdbook that was active since 1976. Just after the second world war 59 the breed was also used to produce grey or "tramacchiati" crossbred pigs by crossing with white 60 61 pigs that were fed with whey produced by the cheese factories of Pianura Padana in the North of Italy. This use was stopped by the transportation ban of the pigs due to an outbreak of diseases in 62 63 1968. Since then the number of animals of this breed dropped down, almost leading to the extinction of the breed. At the end of the eighties a few projects started the recovery of this breed 64 and in 1997 the national pig breeders herdbook preliminarily re-activated a section dedicated to 65 Cinta Senese to promote conservation programs that made it possible to constitute a definitive 66

herdbook section for this breed in 2001 (ANAS, 2015; Franci et al., 2007). These alternate periods 67 68 influenced the number of Cinta Senese heads: the number increased reaching about 160,000 in the fifties, then it decreased reaching the lowest number of 81 sows and 3 boars recorded in 1986 and 69 after conservation programs the number of pigs raised to about 5000 heads (Franci et al., 2007; 70 Raimondi, 1954). At present about 900 sows and 150 boars are registered in the National Herdbook 71 (ANAS, 2015). The current stabilized number is supported by the constitution of a Protected 72 73 Designation of Origin (PDO) brand for Cinta Senese meat in 2011 and the development of the Cinta Senese Consortium (Consorzio di Tutela della Cinta Senese). This consortium and the PDO 74 contributed to the visibility of Cinta Senese products and to the added value of the meat of this 75 76 breed that should be defended from potential frauds.

Coat colour is one of the most important traits that differentiate livestock breeds (Fontanesi,
2009). DNA markers associated with coat colours in different livestock species have been already
used to authenticate mono-breed dairy and meat products (D'Alessandro et al., 2007; Russo et al.,
2007; Fontanesi et al., 2010, 2011).

As already mentioned, Cinta Senese pigs are characterised by a typical belted coat colour that 81 82 can be the basis for the development of DNA markers useful for the authentication of Cinta Senese PDO products. The belted allele, in the past thought to be caused by a specific coat colour locus, is 83 84 one allele of the *Dominant white (I)* locus series that lists several alleles derived by complex mutations in the v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene homolog (KIT) gene 85 (whose product is involved in the migration of the melanoblast), namely copy number variations 86 (CNV) and a splice site mutation (Fontanesi et al., 2010; Johansson Moller et al., 1996; Marklund et 87 al., 1998; Pielberg et al., 2002). The combination of these mutations produces the classical white 88 coat colour phenotype (CNV and the splice site mutation) or the patch phenotype (CNV). The 89 molecular basis of the roan or I^d allele is not completely known even though it is due to variants 90 affecting the KIT gene (Fontanesi et al., 2010). The belted allele was suggested to be derived by an 91 uncharacterised regulatory mutation in the KIT gene, as it was not associated to any duplication of 92

the *KIT* gene described for other *I* alleles (Giuffra et al., 1999). Rubin et al. (2012) reported that the
belted allele could be due to duplication events in the promoter region. We recently characterised
different *KIT* gene haplotypes by Sanger sequencing in several cosmopolitan and local pig breeds
including a few Cinta Senese pigs and identified potential breed informative haplotypes (Fontanesi
et al., 2010). In this study we further analyzed the *KIT* gene and identified a DNA marker that, by
comparing 11 different pig breeds and one wild boar population, was useful to design a simple
genotyping test for the authentication of Cinta Senese meat.

100

101 **2. Materials and methods**

102 2.1. Animals and DNA extraction

DNA was extracted from blood samples, liver and muscle specimens and hair roots collected 103 from a total of 602 pigs of 11 different breeds (110 Cinta Senese; 105 Italian Large White; 52 104 105 Italian Landrace; 86 Italian Duroc; 32 Pietrain; 16 Hampshire; 50 Mora Romagnola; 47 Casertana; 50 Apulo Calabrese; 42 Nero Siciliano; and 12 Meishan) and one wild boar population (29 animals) 106 107 for a total of 631 animals. Samples were mainly obtained from previous projects (Fontanesi et al., 108 2010, 2014). Novel blood samples were collected during slaughtering in commercial abattoirs. DNA extraction was carried out using a standard phenol-chloroform protocol (Sambrook et al., 109 1989) or using the Wizard® Genomic DNA Purification kit (Promega Corporation, Madison, WI, 110 USA), following the manufacturer instructions. 111

112

113 *2.2. PCR analysis*

114 PCR was carried out using two primer pairs. The first primer pair (forward: 5'-

115 CCTCGCAGCAGGAGCAGT-3'; reverse: 5'-CTCAGGGCTGAGCATTCG-3') was used to

amplify a fragment of 388 bp encompassing a portion of intron 17, exon 18, intron 18 and a portion

of exon 19 of the porcine *KIT* gene that was used to re-sequence this gene region in 12 Cinta Senese

pigs to confirm previous sequencing data obtained by Fontanesi et al. (2010). The second primer

119 pair (forward: 5'-TGAACATTGCTGACTCCCCT-3'; reverse: 5'-

TGCATTTTACCTAAAGAGAAGAGC-3') was used to amplify a fragment in all 631 animals. 120 The amplicon of 157 bp was used for the PCR-RFLP analysis described below. The amplification 121 reactions were cycled in a 2720 Life Technologies thermal cycler (Life Technologies, Foster City, 122 CA, USA) with the following steps: 5 min at 95 °C; 35 amplification cycles of 30 sec at 95 °C, 30 123 sec at 56 °C, 30 sec at 72 °C; 10 min at 72 °C. The final reaction volume was of 20 µL and 124 included: 50-100 ng of template DNA, 1 U of Taq DNA polymerase (AmpliBioTherm Taq DNA 125 polymerase, Fisher Molecular Biology, Trevose, PA, USA; or EuroTaq DNA polymerase, 126 EuroClone Ltd., Paington, Devon, UK); 1X PCR buffer; 2.5 mM dNTPs; 10 pmol of each primer; 127 128 2.0 mM of MgCl₂.

129

130 2.3. Sequencing and haplotype analysis

131 Amplified fragments obtained using the PCR primers of the first pair reported above were preliminarily treated with 1 µl of ExoSAP-IT[®] (USB Corporation, Cleveland, Ohio, USA) for 15 132 min at 37°C. Treated amplicons were sequenced using the same PCR primers and the Big Dye v3.1 133 cycle sequencing kit (Life Technologies, Foster City, CA, USA). Sequencing reactions were 134 purified using EDTA 0.125 M, Ethanol 100% and Ethanol 70%, following a standard protocol. 135 Then, the purified products were loaded on an ABI3100 Avant sequencer (Life Technologies). 136 Obtained sequences were visually inspected and aligned with the help of the CodonCode Aligner 137 software (http://www.codoncode.com/aligner) using the reference sequence of the corresponding 138 pig KIT gene region (Fontanesi et al., 2010). The 28 KIT gene haplotypes previously reported by 139 Fontanesi et al. (2010) from several pig breeds were aligned and compared with sequences obtained 140 in the current study. These datasets (the previously reported *KIT* haplotypes and the additional 141 sequences obtained here) were used to identify the most informative single nucleotide 142 polymorphism (SNP) of the most frequent Cinta Senese haplotype. 143

145 2.4. Genotyping and data analyses

The genotyping protocol of the selected SNP (g.43597545C>T; position of the nucleotide 146 coordinate on porcine chromosome 8 in the Sscrofa10.2 genome assembly of the Sus scrofa 147 genome) of the Cinta Senese haplotype was based on PCR-RFLP. The amplified fragments of 157 148 bp obtained with the second primer pair reported above was digested with the restriction enzyme 149 DdeI. Briefly, restriction analysis was carried out overnight at 37 °C in a total of 13 µL of reaction 150 volume including 5 µL of PCR product, 3 U of DdeI (Fermentas, Vilnius, Lithuania) and 1X 151 reaction buffer. Produced DNA fragments were electrophoresed in 2.5% agarose gels running in 152 TBE 1X and visualized with 1X GelRed Nucleic Acid Gel Stain (Biotium Inc., Hayward, CA, 153 USA). 154

Hardy-Weinberg equilibrium of the genotyped SNP in the analysed populations was tested 155 using the HWE software program (Linkage Utility Programs, Rockefeller University, New York, 156 157 NY, USA). Pairwise allele and genotype frequency differences between Cinta Senese breed and all other populations were evaluated using absolute delta (δ) allele frequency differential. GENEPOP 158 software version 4.0.7 (Rousset, 2008) was used to calculate population pairwise F_{st} genetic 159 distance and G genic differentiation for each population pair (exact G test; Markov chain 160 parameters were: Dememorisation: 10000; Batches: 100; Iterations per batch: 5000). GenAlEx 6.5 161 162 software (Peakall and Smouse, 2012) was used to calculate population assignment of the pigs to the 12 populations using the leave one out option. Probability to incorrectly assign an unknown meat 163 164 sample to populations different from Cinta Senese or crossbred products of this breed versus all 165 other populations (error rate) was calculated using the frequency of occurrence of Cinta Senese pigs carrying allele C. Vice versa probability to correctly assign an unknown meat sample to Cinta 166 Senese (P_{CS}) was calculated using the following formula: 167

168 $P_{CS} = 1 - (f_{TT} + f_{CT})$ (1)

where f_{TT} and f_{CT} are the frequency of occurrence of pigs with genotype TT or CT in the other populations. 171

172 **3. Results**

173 *3.1. Identification and analysis of a breed-informative SNP*

Haplotype analyses of sequencing data reported by Fontanesi et al. (2010) who sequenced the 174 KIT gene in 35 pigs of different breeds (including 6 Cinta Senese pigs) showed that putative Cinta 175 Senese informative haplotypes could be identified (Figure 1). From these preliminary data, only two 176 177 haplotypes were observed in Cinta Senese pigs (Haplotype 9 and Haplotype 24). However Haplotype 24 was identified in just one animal of this breed (in heterozygous condition with 178 Haplotype 9). In addition, based on this information, it seemed that just a fragment of this gene, 179 180 including exon 18 and part of exon 19 (Figure 2) could capture most of Haplotype 9 sequence information due to the presence of a tag marker (g.43597545C>T SNP; a synonymous SNP on exon 181 18). Resequencing 24 other Cinta Senese haplotypes (from 12 unrelated Cinta Senese pigs), we 182 183 confirmed that obtained sequences were the same as already reported for the most frequent Cinta Senese haplotype. Allele T of the tag g.43597545C>T marker was present in Cinta Senese pigs 184 185 whereas allele C was present in all other haplotypes except in one rare haplotype observed in a Nero 186 Siciliano pig (as reported from the previous data ; Fontanesi et al., 2010; Figure 1). To further validate these results and to identify a marker that could be useful to authenticate Cinta Senese meat 187 188 products we set up a PCR-RFLP genotyping protocol to analyze a larger number of animals (a total of 631 pigs from different populations were genotyped). The digestion of the amplified product 189 with DdeI produced two fragments of 93 + 64 bp when the amplicon contained allele T whereas 190 191 when allele C was present the fragment of 157 bp remained undigested (Figure 3).

Allele and genotype frequencies at the g.43597545C>T SNP were obtained from 11 pig breeds including Cinta Senese and four other local Italian pig breeds (Mora Romagnola, Casertana, Apulo Calabrese and Nero Siciliano), three commercial heavy pig breeds (Italian Large White, Italian Landrace and Italian Duroc), two cosmopolitan breeds (Pietrain and Hampshire, the other belted breed included in this study), one Chinese breed (Meishan) and a European wild boar

population sampled in Italy (Table 1). None of the populations in which at least two alleles of the 197 g.43597545C>T SNP were detected were in Hardy-Weinberg disequilibrium. Allele T was the most 198 frequent in Cinta Senese pigs (95.9%). The same allele was the most frequent in the other belted 199 200 breed (Hampshire) included in our survey (89.9%). In all other populations, allele T was not identified (Italian Large White, Italian Landrace, Pietrain, Mora Romagnola, Apulo-Calabrese, 201 Meishan and European wild boars, in which only allele C was detected) or its frequency was $\leq 6\%$ 202 (Italian Duroc, Casertana and Nero Siciliano). Comparing the frequency of allele T in Cinta Senese 203 204 vs all other populations (Table 1), δ was ≥ 0.90 in all comparisons (δ was equal to 0.899 in Nero Siciliano) except against the Hampshire breed ($\delta = 0.084$). The assignment test indicated that 91.8% 205 of Cinta Senese pigs could be correctly assigned to their breed based on just the genotyped SNP 206 whereas for all other breeds this test assigned only 25% of Hampshire pigs to the correct breed and 207 for all other breeds assignment was 0%, mainly because the genotyped SNP was not very 208 209 informative or was fixed for allele C.

Pairwise F_{st} measure based on the g.43597545C>T SNP indicated that all comparisons of Cinta Senese breed against all other breeds and populations were highly significant (P<0.0001) or in the case of Hampshire the comparison identified closeness with the Cinta Senese breed but was still significant (P<0.05) due to the higher frequency of allele C (0.125) in this cosmopolitan breed (Table 2). Genic differentiation for each population pair (exact G test) including Cinta Senese was highly significant for all other populations except against the Hampshire pigs confirming indirectly the results of the pairwise population matrix of mean genotypic genetic distance (Table 2).

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218 3.2. Usefulness of the KIT g.43597545C>T SNP to differentiate Cinta Senese meat

Based on the genotyping data produced for the g.43597545C>T SNP it is interesting to
evaluate if this *KIT* gene marker could be useful to differentiate Cinta Senese meat from meat of
other pig breeds and from wild boars. In this first analysis, we will exclude the other belted breed

(Hampshire) investigated in this study. The Hampshire breed is not present in Italy and it should not 222 223 be a problem for animals and meat coming from the same country in which Cinta Senese is raised. None of the analysed Cinta Senese pigs had genotype CC. That means that if genotyping 224 results obtained from meat of unknown origin produce genotype CC, origin from Cinta Senese can 225 be excluded with high confidence. If we assume that there is no selection against the CC genotype 226 in Cinta Senese pigs, the frequency of CC animals in this breed would be very low and equal to the 227 228 square of the frequency of allele C, according to the formula of Hardy Weinberg equilibrium (0.041 x 0.041 = 0.00168). This value would be the error rate that we might have when the genotyping test 229 produces CC. However, we should consider that animals of this genotype, based on results obtained 230 231 in this study and from our previous work (Fontanesi et al., 2010), might not have the classical belted coat colour pattern. This is because allele T can be considered the "belted allele". Therefore, 232 the real situation should consider a selection against the genotype CC in the Cinta Senese breed, as 233 234 animals may not have the characteristic belted trait that is used to register Cinta Senese pigs as animals of this breed. 235

If we consider the case in which the genotyping result of a meat of unknown origin is CT, the error rate in assigning this meat to a hybrid pig (obtained by crossing Cinta Senese animals with genotype TT with other animals with genotype CC) different from Cinta Senese is equal to the frequency of CT pigs in the Cinta Senese population (0.089).

If we suppose that our comparison would only include pigs of the investigated breeds 240 (excluding Hampshire) we should add information on the frequency of the CT genotype in all other 241 242 groups of pigs (Table 1). In this case we could also consider the frequency of this genotype to define the error rate in assigning meat of unknown origin with genotype TT to a hybrid pig 243 (obtained by crossing Cinta Senese animals with genotype TT with other animals with genotype 244 CT) different from Cinta Senese is equal to half the frequency of the CT genotypes in the other 245 populations, considering that crossing TT x CT, only half of the F1 animals would have genotype 246 TT. In this way, based on a total of 505 pigs deduced from Table 1 (all genotyped animals 247

excluding Cinta Senese and Hampshire) the number of CT non-Cinta Senese pigs was 15; therefore ((15/505)/2 = 0.015, is the error rate for this specific question.

Vice versa probability to correctly assign an unknown meat sample to Cinta Senese (P_{CS}) obtained from the data extracted from Table 1 could be $P_{CS} = 1 - 0.0317 = 0.9683$ that derives from the formula (1) of the Materials and methods section in which $f_{TT} = 1/505$ and $f_{CT} = 15/505$, where 1 is the only TT individual identified in the Duroc breed, 15 is the number of CT animals identified in Duroc (n. = 6), Casertana (n. = 4) and Nero Siciliano (n. = 5), and 505 is the total number of analysed animals, excluding Cinta Senese and Hampshire.

Following these procedures it is possible to easily calculate pairwise statistics for all breeds 256 257 against Cinta Senese (data not shown). Considering that potential frauds would not be originated by substituting meat coming from other local breeds (that are the only ones that have the T allele, 258 excluding the Duroc and the Hampshire breeds) but only using cheaper meat originated from 259 260 commercial populations or breeds that are usually of white belted coat colour and in which allele T is not present (or it could be present at a very low frequency, that was not possible to detect in our 261 survey), i) P_{CS} would be 1 and ii) the error rate in assigning to Cinta Senese a meat with genotype 262 TT in case there would be doubts from its possible hybrid origin would be equal to 0.00. 263

264

265 **4. Discussion**

Authentication of meat products obtained from DNA-based approaches is becoming more 266 precise and powerful considering the large amount of genomics data that is currently available and 267 that will be available in the future to extract useful information to answer a large number of new 268 questions and problems arising in this field (Bertolini et al., 2015). One of the most challenging 269 270 problems is the authentication of mono-breed labelled products usually obtained with local and less productive breeds (Fontanesi, 2009; Montowska and Pospiech, 2012). The problems arise by the 271 fact that it is difficult to identify breed-specific markers as animals of different breeds can 272 interbreed producing fertile hybrids and, for this reason, might share a large number of common 273

variants. Methods for the authentication of breed-specific products are key tools to defend the added
economic value of these products that is the strategy that can obtain a sustainable conservation of
local animal genetic resources, as part of an integrated development of their production chains
(Fontanesi, 2009).

Local pig breeds represent an important resource that should be preserved and distinguished to generate economic values for niche markets that are based on their meat products. Cinta Senese products are probably one of the most valuable examples of niche pork production chain (Pugliese and Sirtori, 2012).

In this study, mining data obtained in genes affecting breed specific traits (in our case, coat 282 283 colour), we identified a DNA marker in the KIT gene (g.43597545C>T) that can be useful to distinguish meat from belted pigs (the characteristic trait differentiating Cinta Senese pigs from 284 many other pig breeds and commercial populations as well as from wild boars). Allele T was almost 285 286 fixed in this breed (0.959) and for this reason can be considered its breed-characteristic allele, associated with the belted phenotype. This allele was also the most frequent in Hampshire (0.875), 287 288 that is the other belted pig breed included in this study. However, at present Hampshire is not raised 289 in Italy (only a few animals of this breed might be present in Italy), therefore there is no risk of substitution of Cinta Senese meat with meat obtained from Hampshire pigs. A potential risk could 290 291 be derived by imported Hampshire meat but this is a remote possibility for the fact that pure Hampshire populations are maintained in breeding stocks for crossbreeding programs and are not 292 commonly used for large commercial productions of final slaughtered pigs. Anyway, in this case 293 294 the cost of the fraud would be high and not economically convenient.

Allele T was also observed in a few other breeds (Casertana, Nero Siciliano and Italian Duroc) in which it segregates at very low frequency. Only one TT animal, that apparently not had a belted phenotype, was observed in a non-belted breed. It could be possible that another very rare haplotype including allele T would be present in a few breeds or populations, as also suggested by Fontanesi et al. (2010). Moreover, the belted haplotype is also expected to segregate in non-belted

300 breeds as deduced by the results of crossbreeding programs that sometimes produce belted pigs. 301 However, the very low frequency of allele T in other pig populations is not a big problem for the authentication of Cinta Senese products. This is due to the fact that Casertana and Nero Siciliano 302 are local pig populations for which there is no convenience in using their meat to substitute Cinta 303 Senese pork products and usually pure Duroc pigs are not produced on large scale, as this breed is 304 305 constituted by a small nucleus used to produce sires useful for crossbreeding programs. Therefore 306 the frequency of the CT genotype in these breeds could not be considered to calculate the error rate and P_{CS} derived by this SNP for Cinta Senese products. On the other hand, the low frequency of the 307 alternative C allele in Cinta Senese does not substantially affect error rate and P_{CS} in this pig breed. 308 309 The presence of this allele in Cinta Senese (but also in Hampshire) can indirectly confirm what is 310 obtained from crossbreeding within the breed that sometimes produces piglets without the typical belted phenotype. 311

312 Other studies have investigated the possibility to authenticate mono-breed pork products or to distinguish meat of domesticated pig breeds from meat of wild boars using DNA markers. For 313 example, markers in genes affecting coat colour (MC1R; OCA2; and KIT) or other phenotypic traits, 314 like vertebral number (NR6A1), that are breed or population specific traits, have been already 315 proposed to this purpose (Carrión et al., 2003; Chung and Chung, 2010; Crovetti et al., 2007; 316 317 D'Alessandro et al., 2007; Fernández et al., 2004; Fontanesi et al., 2005, 2014; Okumura et al., 2000). Other approaches have used a large number of SNPs derived from the Illumina Porcine 318 SNP60 BeadChip array (that can genotype more than 60,000 SNPs) or identified by next generation 319 320 sequencing of breed specific DNA pools to identify a subset of 96 SNPs (Wilkinson et al., 2012) or 193 SNPs (Ramos et al., 2011) for breed genetic discrimination among several pig breeds using 321 322 different statistical approaches. Panels of microsatellites were also proposed for the same aim or to identify the level of admixture composition between different breeds (García et al., 2006; Oh et al., 323 2014). However the use of many markers for authentication of breed specific products has some 324 practical and cost-limiting aspects to be solved for routine applications. A few other attempts based 325

on coat colour gene markers and multilocus microsatellite genotyping were also tested for 326 327 traceability of Cinta Senese products. In particular, Crovetti et al. (2007) analysed three MC1R polymorphisms, one of which can distinguish Duroc pigs from black pigs (Kijas et al., 1998), and 328 the duplication breakpoint test for the KIT gene that should give positive results only in white pigs 329 (Giuffra et al., 2002). However, these markers have some limits derived by the fact that Cinta 330 Senese breed is not fixed (or not almost fixed) for only one *MC1R* allele (Crovetti et al., 2007; 331 332 Fontanesi et al., 2005). In addition, as the test for the duplication breakpoint of the *KIT* gene is designed to have an amplified product in case there is a duplication of the KIT gene (usually in 333 white pigs) or absence of amplification in case there is no duplication (all other pigs with different 334 335 coat colours), it could be possible that the absence of amplification might be derived by PCR failure that would prevent the correct identification of the type of tested pigs. To avoid this problem, a 336 multiplex PCR should be designed including a control amplified fragment that should always be 337 338 produced in any types of pigs (D'Alessandro et al., 2007; Fontanesi et al., 2010). Anyway, the duplication breakpoint KIT gene test cannot give specific indications about the breed of the pigs. 339 340 Scali et al. (2012) proposed to use genotype information from 18 microsatellites to differentiate Cinta Senese pigs from Landrace, Large White, Large White x Landrace and Landrace x Cinta 341 Senese pigs. However, their approach was not statistically supported as just 3 or 4 pigs for the other 342 343 breeds or populations were included in the study that did not report the use of any standard sample to refer microsatellite allele size. In addition, more than one PCR and capillary electrophoresis 344 might be needed to obtain the multilocus microsatellite information, even if these details were not 345 346 reported in their work (Scali et al., 2012).

Our method based on the analysis of just one SNP (g.43597545C>T) that is highly informative for belted pigs can directly provide information from the type of pigs from which the meat is originated and is much more precise, cheaper and useful than the methods reported above based on *MC1R* and duplication breakpoint *KIT* gene analysis (Crovetti et al., 2007; Fontanesi et al., 2005) or microsatellite genotyping (Scali et al., 2012). 352

353 **5.** Conclusions

We have identified an SNP that is useful to distinguish meat of Cinta Senese pigs from meat 354 of other non-belted pigs. This marker can be easily genotyped by PCR-RFLP using basic 355 instruments commonly available in a molecular genetics laboratory. Therefore, it can be considered 356 as an important tool to defend Cinta Senese pork chain from frauds. In addition, as this marker 357 358 might capture the belted coat colour phenotype in pigs, it could be used to fix the belted phenotype in Cinta Senese population reducing the out-of-type animals in this breed obtained sometimes by 359 crossing Cinta Senese pigs. Moreover, it will be interesting to evaluate if this marker could be 360 361 associated with the belted phenotype in other local belted pigs that are present in Europe (i.e. Schwäbisch-Hällisches in Germany and Krškopoljski in Slovenia) and for which mono-breed 362 products have been already proposed or could be marketed as a possible way to improve economic 363 364 incomes for the farmers.

365

Conflict of interest statement 366

The authors declare that there are no conflicts of interest. 367

368

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References 377

- ANAS (2015). Registro Anagrafico. <u>http://www.anas.it/</u> (5 May 2015).
- Bertolini, F., Ghionda, M.C., D'Alessandro, E., Geraci, C., Chiofalo, V., Fontanesi, L., 2015. A next
 generation semiconductor based sequencing approach for the identification of meat species in
 DNA mixtures. *PLoS ONE 10*, e0121701.
- 382 Carrión, D., Day, A., Evans, G., Mitsuhashi, T., Archibald, A., Haley, C., Andersson, L., Plastow,
- G., 2003. The use of MC1R and KIT genotypes for breed characterisation. Archiv. Zootec.
 52, 237-244.
- Chung, E.-R., Chung, K.Y., 2010. Identification of Korean Native Pork using breed-specific DNA
 marker of KIT gene. Kor. J. Food Sci. Anim. Res. 30, 403-409.
- 387 Crovetti, A., Bozzi, R., Nardi, L., Franci, O., Fontanesi, L., Russo, V., 2007. Analysis of coat colour
- 388 genes for traceability of Cinta Senese products. Proceedings of 6th International Symposium
- on the Mediterranean Pig. October 11–13, 2007, Messina Capo d'Orlando (ME), Italy, pp.
 366-368.
- D'Alessandro, E., Fontanesi, L., Liotta, L., Davoli, R., Chiofalo, V., Russo, V., 2007. Analysis of
 the *MC1R* gene in the Nero Siciliano pig breed and usefulness of this locus for breed
 traceability. Vet. Res. Comm. 31 (Suppl. 1), 389-392.
- Fernández, A., Fabuel, E., Alves, E., Rodriguez, C., Silió, L., Óvilo, C., 2004. DNA tests based on
 coat colour genes for authentication of the raw material of meat products from Iberian pigs. J.
 Sci. Food Agric. 84, 1855-1860.
- Fontanesi, L., 2009. Genetic authentication and traceability of food products of animal origin: new
 developments and perspectives. Ital. J. Anim. Sci. 8 (Suppl. 2), 9-18.
- Fontanesi, L., Beretti, F., Dall'Olio, S., Portolano, B., Matassino, D., Russo, V., 2011. A
 melanocortin 1 receptor (*MC1R*) gene polymorphism is useful for authentication of Massese
 sheep dairy products. J. Dairy Res. 78, 122-128.
- 402 Fontanesi, L., Bozzi, R., Tazzoli, M., Crovetti, A., Davoli, R., Franci, O., Russo, V., 2005. Genetic
- 403 characterization of Cinta Senese pig breed: analysis of polymorphisms in four genes affecting

- 404 performance and phenotypic traits. Proceedings of the International Workshop "The Role of
- 405 Biotechnology for the Characterisation and Conservation of Crop, Forestry, Animal and
- 406 Fishery Genetic Resources", Villa Gualino, Turin 5-7 March 2005. Book of Proceedings, pp.
- 407 175-176.
- Fontanesi, L., D'Alessandro, E., Scotti, E., Liotta, L., Crovetti, A., Chiofalo, V., Russo, V., 2010.
 Genetic heterogeneity and selection signature at the *KIT* gene in pigs showing different coat
 colours and patterns. Anim. Genet. 41, 478-492.
- 411 Fontanesi, L., Ribani, A., Scotti, E., Utzeri, V.J., Veličković, N., Dall'Olio, S., 2014. Differentiation
- of meat from European wild boars and domestic pigs using polymorphisms in the *MC1R* and *NR6A1* genes. Meat Sci. 98, 781-784.
- Fontanesi, L., Scotti, E., Russo, V., 2010. Analysis of SNPs in the *KIT* gene of cattle with different
 coat colour patterns and perspectives to use these markers for breed traceability and
 authentication of beef and dairy products. Ital. J. Anim. Sci. 9, e42.
- Franci, O., Gandini, G., Bozzi, R., 2007. Why and how to select a local porcine breed: the case of
 the Cinta Senese. *Options Méditerranéennes: Série A. Séminaires Méditerranéens* 76, 13-21.
- 419 García, D., Martínez, A., Dunner, S., Vega-Pla, J.L., Fernández, C., Delgado, J.V., Javier Cañón, J.,
- 420 2006. Estimation of the genetic admixture composition of Iberian dry-cured ham samples
- 421 using DNA multilocus genotypes. Meat Sci. 72, 560-566.
- 422 Giuffra, E., Evans, G., Törnsten, A., Wales, R., Day, A., Looft, H., Plastow, G., Andersson, L.,
 423 1999. The *Belt* mutation in pigs is an allele at the *Dominant white (I/KIT)* locus. Mamm.
 424 Genome 10, 1132-1136.
- 425 Giuffra, E., Törnsten, A., Marklund, S., Bongcam-Rudloff, E., Chardon, P., Kijas, J.M., Anderson,
- 426 S.I., Archibald, A.L., Andersson, L., 2002. A large duplication associated with dominant
- 427 white color in pigs originated by homologous recombination between LINE elements flanking
- 428 *KIT*. Mamm. Genome 13, 569-577.
- Hoffmann, I., 2011. Livestock biodiversity and sustainability. Liv. Sci. 139, 69-79.

- Johansson Moller, M., Chaudhary, R., Hellmén, E., Höyheim, B., Chowdhary, B., Andersson, L.,
 1996. Pigs with the dominant white coat color phenotype carry a duplication of the *KIT* gene
 encoding the mast/stem cell growth factor receptor. Mamm. Genome 7, 822-830.
- Kijas, J.M., Wales, R., Törnsten, A., Chardon, P., Moller, M., Andersson, L., 1998. Melanocortin
 receptor 1 (*MC1R*) mutations and coat color in pigs. Genetics 150, 1177-1185.
- 435 Marklund, S., Kijas, J., Rodriguez-Martinez, H., Rönnstrand, L., Funa, K., Moller, M., Lange, D.,
- Edfors-Lilja, I., Andersson, L., 1998. Molecular basis for the dominant white phenotype in the
 domestic pig. Genome Res. 8, 826-833.
- Maudet, C., Taberlet, P., 2002. Holstein's milk detection in cheeses inferred from melanocortin
 receptor 1 (MC1R) gene polymorphism. J. Dairy Sci. 85, 707-715.
- Montowska, M., Pospiech, E., 2012. Is authentication of regional and traditional food made of meat
 possible?. Crit. Rev. Food Sci. Nutr. 52, 475-487.
- 442 Oh, J.D., Song, K.D., Seo, J.H., Kim, D.K., Kim, S.H., Seo, K.S., Lim, H.T., Lee, J.B., Park, H.C.,
- 443 Ryu, Y.C., Kang, M.S., Cho, S., Kim, E.S., Choe, H.S., Kong, H.S., Lee, H.K., 2014. Genetic
- traceability of black pig meats using microsatellite markers. Asian-Austral. J. Anim. Sci. 27,
 926-931.
- 446 Okumura, N., Kobayashi, E., Suzuki, H., Morozumi, T., Hamashima, N., Mitsuhashi, T., 2000).
- 447 Breed specific mutations in *MC1R* and *KIT* genes in pigs. Anim. Sci. J. 8, 222-234.
- Peakall, R., Smouse, P.E., 2012. GenAlEx 6.5: genetic analysis in Excel. Population genetic
 software for teaching and research-an update. Bioinformatics 28, 2537-2539.
- Pielberg, G., Olsson, C., Syvänen, A.C., Andersson, L., 2002. Unexpectedly high allelic diversity at
 the *KIT* locus causing dominant white color in the domestic pig. Genetics 160, 305-311.
- 452 Pugliese, C., Sirtori, F., 2012. Quality of meat and meat products produced from southern European
- 453 pig breeds. Meat Sci. 90, 511-518.
- 454 Raimondi, R., 1954. Gli aspetti attuali della suinicoltura italiana. L'Italia Agricola 7, 1-16.

- Ramos, A.M., Megens, H.J., Crooijmans, R.P., Schook, L.B., Groenen, M.A., 2011. Identification
 of high utility SNPs for population assignment and traceability purposes in the pig using highthroughput sequencing. Anim. Genet. 42, 613-620.
- Rousset, F., 2008. Genepop'007: a complete reimplementation of the Genepop software for
 Windows and Linux. Mol. Ecol. Res. 8, 103-106.
- 460 Rubin, C.J., Megens, H.J., Martinez Barrio, A., Maqbool, K., Sayyab, S., Schwochow, D., Wang,
- 461 C., Carlborg, Ö., Jern, P., Jørgensen, C.B., Archibald, A.L., Fredholm, M., Groenen, M.A.,
 462 Andersson, L., 2012. Strong signatures of selection in the domestic pig genome. Proc. Natl.
 463 Acad. Sci. USA 109, 19529-19536.
- 464 Russo, V., Fontanesi, L., Scotti, E., Tazzoli, M., Dall'Olio, S., & Davoli, R., 2007. Analysis of
- melanocortin 1 receptor (*MC1R*) gene polymorphisms in some cattle breeds: their usefulness
 and application for breed traceability and authentication of Parmigiano Reggiano cheese. Ital.
 J. Anim. Sci. 6, 257-272.
- 468 Sambrook, J., Fritsch, E.F., Maniatis, T., 1989. Molecular cloning: a laboratory manual. New York:
 469 Cold Spring Harbor Laboratory Press.
- 470 Scali, M., Vignani, R., Bigliazzi, J., Paolucci, E., Bernini, A., Spiga, O., Niccolai, N., Cresti, M.,
- 471 2012. Genetic differentiation between Cinta Senese and commercial pig breeds using
 472 microsatellite. Electronic J. Biotechnol. 15, 1-11.
- 473 Wilkinson, S., Archibald, A.L., Haley, C.S., Megens, H.J., Crooijmans, R.P., Groenen, M.A.,
- 474 Wiener, P., Ogden, R., 2012. Development of a genetic tool for product regulation in the
- diverse British pig breed market. BMC Genomics 13, 580.

477 **Figure captions**

478

Figure 1. *KIT* gene haplotypes identified in different pig populations (D = Italian Duroc; CS = Cinta Senese; G = Gray pigs; H = Hampshire; LW = Italian Large White; M = Meishan; NS = Nero Siciliano; P = Pietrain; WB = Wild boar) as defined in Fontanesi et al. (2010) with indicated the allele of the tag marker (g.43597545C>T). Haplotypes identified in Cinta Senese pigs are evidenced.

484

Figure 2. Resequenced *KIT* gene region including the tag marker g.43597545C>T (within squared
brackets) and encompassing part of intron 17, exon 18, intron 18 and part of exon 19. Exon regions
are indicated in bold and evidenced. Primer regions are underlined.

488

489 Figure 3. PCR-RFLP patterns obtained genotyping the g.43597545C>T SNP (M = molecular DNA
490 ladder; the genotypes are indicated above each gel line).

492 Table 1. Several statistics and data obtained for the g.43597545C>T single nucleotide
493 polymorphism of the *KIT* gene in different pig breeds and in wild boars.

494

Breed/population	No. of	Genotypes (no. of pigs)		Allele frequencies		HWE ¹	δ^2	
	pigs	CC	СТ	TT	С	Т		
Cinta Senese	110	0	9	101	0.041	0.959	0.535	-
Italian Large White	105	105	0	0	1.000	0.000	-	0.959
Italian Landrace	52	52	0	0	1.000	0.000	-	0.959
Italian Duroc	86	79	6	1	0.953	0.047	0.145	0.912
Pietrain	32	32	0	0	1.000	0.000	-	0.959
Hampshire	16	0	4	12	0.125	0.875	0.449	0.084
Mora Romagnola	50	50	0	0	1.000	0.000	-	0.959
Casertana	47	43	4	0	0.957	0.043	0.673	0.916
Apulo-Calabrese	50	50	0	0	1.000	0.000	-	0.959
Nero Siciliano	42	37	5	0	0.940	0.060	0.574	0.899
Meishan	12	12	0	0	1.000	0.000	-	0.959
European wild boars	29	29	0	0	1.000	0.000	-	0.959

495

496 ¹ Hardy Weinberg Equilibrium (P value).

498 breed and populations.

^{497 &}lt;sup>2</sup> Absolute delta (δ) allele frequency differential of the T allele between Cinta Senese and all other

Table 2. Pairwise population statistics comparing Cinta Senese data versus all other investigated
 breeds including F_{st}, genic differentiation (exact G test) and genotypic genetic distance

Breeds	F _{st} (P value)	P value of the G	Genotypic
		test	distance
Italian Large White	0.9580 (<0.0001)	< 0.0001	3.755
Italian Landrace	0.9449 (<0.0001)	< 0.0001	3.755
Italian Duroc	0.9089 (<0.0001)	< 0.0001	3.514
Pietrain	0.9376 (<0.0001)	< 0.0001	3.755
Hampshire	0.0532 (0.018)	0.2315	0.291
Mora Romagnola	0.9443 (<0.0001)	< 0.0001	3.755
Casertana	0.9131 (<0.0001)	< 0.0001	3.513
Apulo-Calabrese	0.9443 (<0.0001)	< 0.0001	3.755
Nero Siciliano	0.9016 (<0.0001)	< 0.0001	3.417
Meishan	0.9279 (<0.0001)	< 0.0001	3.755
European wild boars	0.9363 (<0.0001)	< 0.0001	3.755