

The Myostatin gene: an overview of mechanisms of action and its relevance to livestock animals

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1 Review: The *Myostatin* gene: an overview of mechanisms of action and its

2 relevance to livestock animals

- 3 D. Aiello¹, K. Patel² and E. Lasagna¹
- 4
- ⁵ ¹ Dipartimento di Scienze Agrarie, Alimentari e Ambientali, Università degli Studi di
- 6 Perugia, Borgo XX Giugno 74, 06121, Perugia, Italy
- ⁷ ² School of Biological Sciences, University of Reading, Berkshire, RG6 6UB, United
- 8 Kingdom
- 9
- 10 Corresponding author: Emiliano Lasagna. Fax: +39 075 5857122. Tel: +39 075
- 11 5857102. E-mail address: emiliano.lasagna@unipg.it
- 12
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15 Summary

Myostatin, also known as Growth Differentiation Factor 8, a member of the 16 17 Transforming Growth Factor-beta (TGF-β) super-family is a negative regulator of 18 muscle development. Myostatin acts at key points during pre- and post-natal life of 19 amniotes which ultimately determine the overall muscle mass of an animal. Mutations 20 have already demonstrated the impact of attenuating Myostatin activity on muscle 21 development. A number of large animals including cattle, sheep, dogs and humans 22 display the 'double muscled' phenotype due to mutations in the Myostatin gene. Here we firstly give an overview of the molecular pathways regulated by Myostatin that 23 24 control muscle development. Then we describe the natural mutations and their 25 associated phenotypes as well as the physiological influence of altering Myostatin 26 expression in livestock animals (cattle, sheep, goat, horse, pig, rabbit and chicken). Knowledge of null alleles and polymorphisms in the Myostatin gene are of great 27 interest in the animal breeding field and it could be utilized to improve meat 28 29 production in livestock animals.

30

31 **Keywords:** double muscling, single nucleotide polymorphisms, muscle hypertrophy,

32 muscle hyperplasia, meat production.

33

34 Introduction

35 Myostatin

36 Myostatin (MSTN), also known as Growth and Differentiation Factor 8 (GDF8), is one

of the major regulators of skeletal muscle development (Beyer *et al.*, 2013). The

38 MSTN gene (*MSTN*) is highly conserved among mammalian species and it acts in an

almost unique manner to reduce muscle size. MSTN-deficient animals display an
increase in skeletal muscle mass known as double-muscling (DBM). Mutations in *MSTN* have been described in numerous species including dog (Mosher *et al.,*2007), sheep (Kijas *et al.,* 2007), cattle (Grobet *et al.,* 1997), pig (Stinckens *et al.,*2008) as well as in one human (Schuelke *et al.,* 2004).

44

45 Myostatin signalling pathway and its control of skeletal muscle development

46 MSTN is expressed in many tissues (including the mammary gland) but most

47 prominently in skeletal muscle (Ji *et al.*, 1998). The *MSTN* has been highly conserved

48 throughout evolution and comprises 3 exons and 2 introns.

49 In all species reported in this review, *MSTN* exons code for a 375 amino acid latent

50 protein which undergoes significant post-translational modification in order to become

51 biologically active (Wolfman *et al.*, 2003). Firstly, the polypeptide undergoes

52 intracellular homodimerization through the formation of disulphide bonds. Thereafter

53 it is cleaved to form the N-terminal propeptide region and the C-terminal mature

region. The 12- KDa C-terminal mature fragment of MSTN initiates an intracellular

55 signalling cascade through its ability to bind and activate the Activin type II receptor

56 at the cell surface (ActRIIB and to a lesser extent ActRIIA). Subsequent

57 autophosphorylation of the ActRIIB leads to the recruitment and activation of low

58 affinity type I receptor for Activin ALK-4 or ALK-5. Activated type I receptor kinase

59 phosphorylates the transcription factors Smad2 and Smad3, allowing them to interact

60 with Smad4 (co-Smad) and translocate to the nucleus, to activate target gene

61 transcription. Importantly the activation of the MSTN receptor also inhibits Akt

62 (protein kinase B) activity, a major determinant in muscle protein synthesis and cell

63 proliferation. Enlargement of muscle fibre size, a process called fibre hypertrophy (or

simply hypertrophy) is in large part controlled by Akt activity (Trendelenburg et al., 64 65 2009). Myogenic differentiation is a highly orchestrated sequential program that ultimately generates mature skeletal muscle. Highly proliferative muscle precursors 66 67 which arise during embryogenesis differentiate into myoblasts. The commitment of the myogenic lineage is regulated by Muscle Regulatory Factors (MRFs) a collective 68 69 group of helix-loop-helix transcription factors; namely, MyoD, Myf5, Myogenin and MRF4 (Fig. 1). Additionally, exit from the cell cycle is a vital step during myoblast 70 71 differentiation (Bryson-Richardson & Currie, 2008).

MSTN regulates muscle development at key points during the process of pre-natal 72 73 muscle development: muscle precursor proliferation, myoblast proliferation and 74 differentiation. Studies by Amthor et al. (2002) have shown that ectopic expression (in 75 limb muscle) of MSTN down regulates Pax3; a key marker of proliferating muscle 76 precursors (Amthor et al., 2002). Additionally, MSTN upregulates p21 expression, 77 which ultimately inhibits proliferation of MyoD expressing myoblasts (Thomas et al., 78 2000). Of relevance to this review is the relationship between MyoD activity and the 79 expression of MSTN. MyoD is an important regulator of MSTN expression during myogenesis. This is demonstrated by a critical role of E-box motifs that were 80 identified in the MSTN promoter region; these motifs are known to be the binding 81 82 sites for basic helix-loop-helix transcription factors (MRFs) (Hu et al., 2013). 83 The interrelationship between MyoD and MSTN ensure that promiscuous differentiation mediated by an over-active MyoD induced cascade is checked by the 84 85 up-regulation of MSTN. Therefore MSTN serves to limit the size of both the myoblast precursor (Pax3⁺/MyoD⁺) and myoblast (Pax3⁻/MyoD⁺) pools. Down-regulating the 86 87 expression of MSTN would lead to an expansion of both populations (Amthor et al., 1999). 88

89 Examination of mouse development shows that muscle mass is determined by the 90 ability of myoblasts to form fibres, a process that occurs in two phases; primary and 91 secondary fibre formation. Matsakas et al. (2010) have shown an increase in the 92 myoblast pool, just before the fibre formation process in *Myostatin* null mouse (*Mstn*^{-/-} 93) embryos, which supports the development of extranumerary primary and secondary 94 myofibres. Any programme that promotes an increase in fibre formation is called fibre hyperplasia or simply, hyperplasia (Amthor *et al.*, 2002). Therefore the *Mstn*^{-/-} mouse 95 96 displays hyperplasia as a consequence of developing an increased number of mono-97 nucleated muscle cells (Matsakas et al., 2010).

Shortly before birth, muscle in *Mstn^{-/-}* mice not only contain extra muscle fibres, but 98 99 also each fibre has undergone a small, albeit significant, increase in size (18%). 100 However this is not enough to explain why the muscles in this species often weigh 2-101 3 times more than their normal counterpart (Omairi *et al.*, 2016). The resolution to 102 this issue comes by examining the size of each muscle fibre in adult mice. This 103 reveals that in the mouse, the increased muscle mass has arisen due to a 104 combination of a pre-natal increase in the number of fibres (hyperplasia) and a 105 precocious post-natal increase (43%) in the size of each fibre (hypertrophy) 106 (McPherron & Lee, 1997).

These studies are extremely insightful when attempting to determine the cellular mechanism underpinning double muscling in large mammalian species harbouring a *MSTN* mutation (Elashry *et al.*, 2012). They predict that for an animal to develop fibre hyperplasia and a small degree of hypertrophy as a consequence of a *MSTN* mutation, the gene must normally be expressed and properly translated into a mature form during pre-natal development. However, in order to display significant fibre hypertrophy these conditions need to be satisfied during post-natal life. If the mouse

114 is taken as a guide, then changes in fibre number and small changes in fibre

diameter (less than 20%) can be explained by pre-natal action of MSTN. In cattle,

116 very low levels of MSTN are detected from day 15 to day 29 embryos, and increased

expression is detected from day 31 onwards (Kambadur *et al.*, 1997). The increase

of *MSTN* expression in the bovine embryos is thought to occur at a gestational stage

119 when primary myoblasts are starting to fuse and differentiate into myofibres.

120 Therefore the null mutation in the bovine *MSTN* lead to hyperplasia.

121

122 Double muscling phenotypes

123 The term hypertrophy has often been used to describe large mammalian species,

124 which display at the gross anatomical level, the enlargement of muscle.

125 Mechanistically this term has been used loosely, since in many cases enlargement of

126 muscle is solely through pre-natal muscle hyperplasia without any post-natal fibre

127 hypertrophy.

128 DBM in large animals has been reported in several species. Generally, muscle with a

129 large superficial area tends to be the most enlarged, while deeper muscles tend to be

reduced in size relative to normal muscle (Ouhayon & Beaumont, 1968). Large

131 commercially important DBM animals, especially cattle, have an excellent

132 conformation and an extremely high carcass yield, coinciding with a reduced internal

133 organ mass (Fiems, 2012).

134 However, these animals are more susceptible to respiratory disease, urolithiasis,

135 lameness, nutritional stress, heat and dystocia resulting in lower robustness (Holmes

et al., 1973). Also the reproductive performance can be influenced by hypertrophy:

i.e. in the South Devon breed, the gestation period for DBM calves is longer, resulting

138 in offspring with higher birth weights than the normal calves, also evidenced by the

139 higher instances of dystocia with high mortality rates if births are unassisted; the 140 findings highlighted therefore that the segregating alleles at the MSTN have 141 significant effects on calving ease in this breed (Wiener et al., 2002). 142 DBM cattle showed signs of fatiguing faster than normal cattle during forced exercise; relating to metabolic acidosis, because of a reduced blood circulation 143 144 leading to a deficiency in the transport of oxygen and a reduction of aerobic metabolic activity in the muscle (Holmes et al., 1973). DBM cattle have in fact an 145 146 increase in the proportion of fast twitch glycolytic fibres, resulting in a faster and more 147 glycolytic phenotype (Girgenrath et al., 2005).

148 Mutations in the MSTN are responsible for DBM in other large animals including one 149 case in humans. In the latter, Schuelke et al. (2004), observed that a G to A transition 150 at nucleotide gIVS1+5 caused extraordinary muscling in a young boy, especially in 151 the thighs and upper arms. No health problems were reported in the patient and the 152 testosterone and IGF-1 levels were normal. In dogs known as "bully" whippets, a 2-153 bp deletion was discovered in the third exon of the MSTN is associated with the DBM 154 phenotype. This deletion removes nucleotides 939 and 940 within exon three and leads to a premature stop codon at amino-acid 313 instead of the normal cysteine, 155 156 removing 63 amino acids from the predicted 375-aa protein (Mosher et al., 2007). A 157 gene targeting approach using the CRISPR/Cas9 system has been used to create 158 *MSTN* null Beagles; although mutant dogs displayed the DMB phenotype, very little 159 detail is available regarding their cellular phenotype (Zou *et al.*, 2015). Due to the 160 effects of MSTN on muscle mass, growth and other traits, the variations in *MSTN* 161 expression levels in skeletal muscles are of great interest in the animal breeding 162 field. Knowledge of null alleles and polymorphisms in the *MSTN* has been utilized to 163 improve the selection of beef cattle and sheep (Georges, 2010). The aim of this

section of the review is to describe known double-muscling in livestock animals thatharbour *MSTN* mutations.

166

167 Mutations in the Myostatin gene in cattle

168 Monogenic determination of muscular hypertrophy in Belgian Blue cattle was first 169 described in the 1980's (Hanset & Michaux, 1985; Grobet et al., 1997). Double 170 muscling was shown to be inherited as a single major autosomal locus which 171 nevertheless was affected by several modifier loci manifesting in incomplete penetrance. The causal loss of function mutation in Belgian Blue MSTN, located on 172 173 chromosome 2, was first reported by Grobet (1997) followed shortly thereafter by the study of McPherron and Lee who not only substantiated the finding of Grobet but 174 175 also reported a missense mutation in exon 3 in the Piedmontese breed MSTN 176 (McPherron & Lee, 1997). Approximately 20 different types of genetic variants 177 (deletions, insertions and nucleotide substitutions, also known as single nucleotide 178 polymorphisms - SNPs) have been identified in the bovine MSTN. Some of these 179 genetic variants give rise to muscular hypertrophy by inactivation of the gene (Grobet 180 et al., 1997). Mutated alleles and inactive MSTN have a significant association with growth speed and carcass favourite traits, so these polymorphisms could be used in 181 182 beef cattle in order to increase the quality and quantity of meat (Mirhoseini & Zare, 183 2012). In the view of quality meat production, this is an outstanding trait, since these 184 animals produce not just more, but leaner and more tender meat (Kobolák & Gócza, 185 2002). The carcass and meat quality traits are superior in these animals because of a 186 reduction in fat (decreased by 50%), muscle mass increase (by 20%) lower 187 proportions of bone and also less connective tissue, which contributes to tenderness 188 (McPherron & Lee, 1997; Vincenti et al., 2007). However, dystocia-related problems

are often observed in DBM cattle because hyperplasia occurs before birth, resulting
in larger calves (Deveaux *et al.*, 2001). Homozygous DBM animals manifest more
problems of dystocia than heterozygous. Therefore in order to generate homozygous
animals and at the same time keep costs down as well as reducing calve death
probability, it is worth considering mating heterozygous animals (Bellinge *et al.*,
2005).

A summary of the detected genetic variants in cattle is reported in Table 1.

197 Double muscled cattle breeds

198 Belgian Blue

199 The breed in which this muscular hypertrophy and its effects have been analysed 200 most extensively is the Belgian Blue breed, which has been systematically selected 201 for double muscling to the point of fixation in many herds. Research by Grobet et al. 202 (1997) revealed an 11-bp deletion (nucleotides 821-831) in the open reading frame of 203 the Belgian Blue MSTN allele which results in the loss of 3 amino acids (275, 276, 204 and 277) and a frameshift after amino acid 274. The frameshift leads to a stop codon 205 after amino acid 287. Work by Wegner et al. (2000) showed that Semitendinosus from Belgian Blue was 1.6 times the weight of normal breeds solely due to an 206 207 increase in muscle fibre number. Indeed, muscle fibre size from the Belgian Blue was 208 actually smaller than other breeds (Wegner et al., 2000). Furthermore, these animals 209 have less collagen and connective tissue than the normal animals. The carcass fat 210 content in these animals is significantly lower than in normal cattle, especially 211 intramuscular fat (marbling) being influenced by the DBM phenotype with a strong 212 reduction of subcutaneous and internal fat tissues (Mirhoseini & Zare, 2012). The 213 results of many studies in fact have indicated that MSTN plays key roles in not only

214 myogenesis but also adipogenesis. *MSTN* deletion and inhibition in animals mainly 215 lead to increased muscle mass and reduced fat mass (Deng et al., 2017). 216 In beef cattle production, crossing with Belgian Blue cattle shows that although the 217 gene is recessive and monofactorial, its effect is apparent even in heterozygous 218 animals due to its partial dominance (Kobolák & Gócza, 2002). 219 The same mutation was also found in the Asturiana de los Valles (AV), a Spanish 220 beef cattle breed. MSTN polymorphisms in the AV breed have been described and 221 its diffusion into the breed has been continuous due to economic reasons (Grobet et

223

222

224 Piedmontese

al., 1997).

225 In Piedmontese cattle the double-muscled phenotype is an inherited condition 226 associated with a G to A mutation on nucleotide 938 (in exon 3) which translates to 227 C313Y in a highly conserved cysteine-knot structural motif region of the protein. This 228 is in the pre-helix loop, a region known to be important for ALK4/5 receptor 229 interaction (Cash et al., 2012). The mutation alters the function of MSTN, which disrupts a disulphide bridge that is essential for the correct conformation of the 230 231 protein (Kambadur et al., 1997). This breed has been systematically selected for 232 double muscling to the point of fixation in many herds (> 96% homozygous in the 233 Piedmonte region in Italy), but variability in muscle mass is still present (Miretti et al., 234 2013). Several studies support the notion that the double muscling phenotype, a 235 partially recessive trait, causes the relatively large effects on carcass conformation, 236 without a negative effect on calving, compared with animals with no copies of the 237 mutated allele (Casas *et al.*, 1998).

238

239 Marchigiana

240 The Marchigiana is one of the most important Italian beef cattle breeds and it is 241 renowned for its large body size, high weight daily gains and superior carcass 242 dressing percent. Marchigiana breed have a G to T transversion mutation at 243 nucleotide 874 in exon 3 (g.874G>T), translating to E291X in the MSTN. This point 244 mutation has a remarkable effect on the MSTN as it changes a codon for glutamic 245 acid into a stop codon (Marchitelli *et al.*, 2003). In Marchigiana, as in the other double 246 muscling breeds, the MSTN genotypes yield three different and distinct phenotypes. The homozygous G/G displays the normal phenotype whereas the T/T genotype 247 248 manifests as a double muscled body shape while maintaining its small frame, and is frequently associated with skeletal defects and serious survival problems due to 249 250 macroglossia and hypoplasia of the heart, lungs and other vital organs. The 251 heterozygous genotype (G/T) produces a well-muscled and large body structure and 252 excellent conformation without any of the above mentioned defects. Therefore, the 253 heterozygous animals are frequently selected as sires (Cappuccio et al., 1998). 254 Moreover heterozygous animals show a better meat quality than animals with a 255 normal genotype (Vincenti et al., 2007). Therefore they could be useful for breeders 256 to plan the matings to obtain a higher number of heterozygous animals. Obviously 257 this is possible only if the genotype at the MSTN locus of each animal is available. 258 Additionally two different SNPs have been found in the promoter region: g.-371T>A 259 and g.-805G>C, although Sarti et al. (2014) reported that these substitutions may not 260 be useful to be considered in the selection criteria, because there is no correlation 261 with productive traits or due to their homozygous genotype.

262

263 Other cattle breeds

264 An 11 bp deletion (nt821(del11)) resulting in a truncation of the bioactive C-terminal 265 domain of the protein has been found in Blonde d'Aguitaine, Limousine, and Parthenaise and Rubia Gallega breeds (Kambadur et al., 1997; Dunner et al., 2003). 266 267 A recent study (Bouyer et al., 2014) identified an unexpected mutation in the MSTN in Blonde D'Aquitaine cattle. The mutant allele is highly expressed leading to an 268 269 abnormal transcript consisting of a 41-bp inclusion between the exons 2 and 3, with a 270 premature termination codon predicted to translate into a protein lacking the entire 271 bioactive region.

An additional transversion mutation (g.433C>A) in Limousine breed has been

described that was shown to be functionally associated with the increased muscle

mass and carcass yield without any associated reproductive disadvantages (Sellick

275 *et al.*, 2007; Esmailizadeh *et al.*, 2008; Vankan *et al.*, 2010).

As in Piedmontese cattle, a G to A transition at nucleotide position 938 has been

reported in Gasconne (Kambadur *et al.*, 1997; Dunner *et al.*, 2003). An

insertion/deletion at position 419 replacing 7 bp with an unrelated stretch of 10 bp

was reported in Maine-Anjou cattle, resulting in a premature stop codon in the N-

terminal latency-associated peptide at amino-acid position 140 (nt419 (del7- ins10))

281 (McPherron & Lee, 1997). Additionally, a transversion (G to T) at nucleotide position

282 676, also causing a premature stop codon in the same N-terminal latency-associated

283 peptide at amino-acid position 226 (E226X) was identified in the same breed (Grobet

et al., 1997). Charolaise and Limousine have a C to T transition at nucleotide position

285 610 yielding a premature stop codon in the N-terminal latency associated peptide at

amino-acid positions 204 (Q204X) (Cappuccio et al., 1998). In addition to the genetic

variants found in *Bos taurus*, 14 polymorphisms (three in exon one, seven in exon

two, and four in exon three) have been reported in the coding part of the *MSTN* in

Nellore cattle (*Bos indicus*) genome. However, whether these polymorphisms are
functional mutations still remains to be elucidated (Grisolia *et al.*, 2009).

291

292 Double muscling in sheep

293 The *MSTN* is located at the end of the long arm (2g32.2 locus) on chromosome 2 in 294 the sheep (Ovis aries) (Bellinge et al., 2005). During the past decade a total of 77 *MSTN* SNPs have been reported in various sheep breeds such as Texel, Norwegian 295 296 Spælsau, commercial New Zealand sheep breeds and Latvian Darkhead (Kijas et al., 297 2007, Sjakste et al., 2011; Han et al., 2013), and the majority of these SNPs are 298 located in the non-coding regions of the gene. The exception is a 1-bp deletion 299 identified in nucleotide position 960 in the MSTN of Norwegian White Sheep and 300 c.101G/A in New Zealand Romney, c.120insA (Boman et al., 2009). Lastly in 2018, 301 Trukhachev et al., described for the first time eight variations in non-coding regions of 302 MSTN in the Stavropol Merino, a breed used for meat production in Russia. A 303 summary of the detected genetic variants in sheep is reported in Table 2. 304

305 Texel

306 Belgian Texel sheep muscle fibres show enlargement and therefore can be 307 considered to have fibre hypertrophy. Texels are utilized extensively as a terminal 308 crossbreed because of their exceptional conformation and potential to produce 309 higher-yielding carcasses with increased lean and decreased fat content (Leymaster 310 & Jenkins, 1993). Analysing the MSTN revealed no nucleotide differences in the 311 coding regions between DBM and normally muscled breeds (Kijas et al., 2007). This 312 suggests that genetic variation located outside the coding regions plays a more 313 important role in the regulation of muscle development in contrast to cattle, where

314 *MSTN* loss of function variants have been found within the three coding exons 315 (Grobet et al., 1997). Quantitative trait locus (QTL) analysis in Texel sheep 316 characterized a mutation (g.6723G>A) in the 3' UTR (Untranslated Region) of the 317 *MSTN* on chromosome 2 which has an effect on muscle mass. This creates a target 318 site for *miR1* and *miR206*; microRNAs (miRNAs) that are highly expressed in skeletal 319 muscle (Kijas et al., 2007). Other genetic variants have also been found including 320 c.*1232A, g+391G>T and another 18 SNPs: g.2449C>G; g.2379C>T; g.1405A>T; 321 g.1402G>A; g.1214C>T; g.1129C>T; g.41A>C; g.39T>C; g+474C>T; G+613T>C; g+616G>A; g+619T>C; g+622T>C; g+632G>T; g+696C>T; g+3135C>T; 322 323 g+4036A>C; g+4044C>T (Kijas *et al.*, 2007).

324

325 Norwegian sheep

326 The DBM phenotype in Norwegian white sheep was described to have extraordinary 327 over-development of the muscles, particularly on the hindguarters. Investigations 328 showed that these animals have not only extremely low levels of subcutaneous fat, 329 but also decreased internal fatty tissues. The DBM animals had lower bone mass compared with the wild type animal. Sequence analysis revealed a 1-bp deletion in 330 331 the MSTN at nucleotide position 960 in DBM individuals. The deletion of a G residue 332 (c.960delG) disrupted the reading frame from amino acid 320 onwards and produced 333 a premature stop codon at amino acid position 359 (compared to position 375 in the 334 wild type animals) (Boman & Vage, 2009).

The same *MSTN* 3'-UTR mutation (c.2360G>A) identified in Texel sheep was also
found in the Norwegian breed but with a less profound effect (Boman & Vage, 2009).

However a similar phenotype of increased muscle mass and fat was found in

338 Norwegian Spælsau sheep. The sequencing of the *MSTN* coding region revealed a

1-bp insertion at nucleotide position 120 (c.120insA) in DBM animals. The insertion of
an adenine residue disrupts the reading frame from amino acid position 40 onwards,
and generates a premature stop codon at amino-acid position 49 (Boman & Vage,
2009).

343

344 New Zealand

A comprehensive investigation of polymorphisms in MSTN in a diverse range of 345 346 sheep breeds (New Zealand Romney, Coopworth, Corriedale, Dorper, Perendale, Suffolk, Merino, Dorset Down, Poll Dorset, Texel and other NZ cross-bred sheep) 347 348 was performed using polymerase chain reaction-single strand conformational polymorphism (PCR-SSCP) analysis and DNA sequencing. A total of 28 nucleotide 349 350 substitutions were identified from nucleotide c.-1199 (in the promoter region) to 351 c.*1813 in the 3'UTR. Of these, three were located in the promoter region, three in 352 the 5'UTR, 11 in intron 1, five in intron 2 and five in the 3' UTR. Ten new substitutions 353 have been reported: c.-959C>T, c.-784A>G, c.373+563A>G, c.373+607A>G, c.374-354 654G>A, c.374-54T>C, c.748-54T>C, c.*83A>G, c.*455A>G and c.*709C>A (Han et 355 *al.*, 2013). 356 The other 18 substitutions had been reported previously. These include c.101G>A 357 which was already found in NZ Romney by Zhou et al. (2008) and also in Merino, 358 Corriedale and NZ cross-bred sheep (Clop et al., 2006; Kijas et al., 2007). In NZ 359 Romney a further two SNPs c.-2449G/C and c.-2379T/C were detected (Wang et al., 2016). The SNP c.*123A observed in NZ cross-bred sheep was also reported in 360

361 Texel (Kijas *et al.*, 2007), Charollais sheep from Britain (Hadjipavlou *et al.*, 2008),

362 White Suffolk, Poll Dorset and Lincoln breeds from Australia and showed significant

association with DBM phenotype as well as the other substitution c.373+18 T>G,
reported in Texel sheep (Clop *et al.,* 2006).

365

366 Other sheep breeds

Zel sheep, a meat breed in northern Iran, has a polymorphism in intron 2 as does the
Iranian Baluchi sheep (Dehnavi *et al.*, 2012). Three polymorphic sites in Indian sheep
have been identified in the 5'UTR, exon 1 and exon 2 regions. Both SNPs in the
exonic region were found to be non-synonymous. The genetic variants c.539T>G and
c.821T>A were in the exon 1 and exon 2, respectively (Pothuraju *et al.*, 2015). All
these genetics variants are not significantly associated with DBM phenotype.

373

374 Myostatin polymorphisms in goat

375 Several studies investigated the allelic variation in the goat MSTN. A 5 bp indel (1256 376 TTTTA/-) was identified in 5'UTR region in Boer, Matou, Haimen and Nubi goat 377 breeds, and a substitution (1388 T/A) in exon 1 region was detected only in Boer 378 (Zhang et al., 2012). Two novel single nucleotide polymorphisms were also identified 379 in Boer and Anhui white goat: g.197G>A, a substitution located in the 5'-UTR, and 380 345A>T in the exon 1 (Zhang et al., 2013). A thorough investigation was conducted 381 in 22 different goat breeds (Inner Mongolia Cashmere, Liaoning Cashmere, Taihang 382 Mountain, Chengde Polled, Jining Grey, Tibetan, Chengdu Brown, Jianchang Black, 383 Guizhou White, Guizhou Black, Longlin, Duan goat, Leizhou, Matou, Yichang White, 384 Shannan White, Nanjiang Brown, Angora, Toggenburg, Nubian, Saanen and Boer 385 goat) and a total of eight SNPs were detected (A1980G, G1981C, A1982G, G1984T, 386 A2121G, T2124C, G2174A and A2246G) (Li et al., 2006). Recently Nguluma et al. (2018) detected a polymorphic site T298C in the Boer goat population: the authors 387

concluded that the potential association of this polymorphism in *MSTN* with growth
performance could not be confirmed and that other genes for growth could be
responsible for the observed variation. A summary of the detected genetic variants in
goat is reported in Table 3.

392

393 Myostatin polymorphisms in horse

394 Hosoyama et al. (2002) isolated and sequenced MSTN cDNA from a Thoroughbred 395 horse which was mapped to chromosome 18. Mutations in the equine MSTN have 396 been identified and are associated with racing phenotypes influencing racing 397 performance and muscle fibre proportions (Petersen et al., 2013). Dall'Olio et al. 398 (2010) sequenced in 16 horse breeds (Rapid Heavy Draft, Noric, Bardigiano, Haflinger, Lipizzan, Murgese, Tolfetano, Uruguayan Creole, Italian Saddle, 399 400 Maremmano, Quarter Horse, Salernitano, Andalusian, Ventasso, Italian trotter, 401 Thoroughbred horse) revealing seven SNPs: two transitions were located in the 402 promoter region at -646 (GQ183900: g.26T>C) and -156 (GQ183900: g.156T>C) bp 403 upstream from the start codon and are associated with breeds of different 404 morphological types. The g.26T>C SNP was polymorphic in 6/16 breeds with higher observed frequency of the g.26C allele. The g.156T>C polymorphism was detected 405 406 in 11/16 breeds and was identified in homozygous condition in a few Bardigiano, 407 Haflinger, Noric, Rapid Heavy Draft, and Uruguayan Creole horses (Dall'Olio et al., 408 2010). The other five SNPs were in intronic regions: four were localized in intron 1 409 and one in intron 2. Three of the SNPs of intron 1 (g.1634T>G, g.2115A>G, and 410 g.2327A>C) were also identified in Thoroughbred breeds (Petersen et al., 2013). One 411 polymorphism (g.2115A>G) has been associated with sprinting ability and racing 412 stamina in Thoroughbred horses. The association between *MSTN* and horse racing

413 performances was further evidenced by Binns et al. (2010) and Tozaki et al. (2010). 414 Subsequently 15 Chinese breeds were studied to select the best Chinese domestic 415 breed to evaluate the potential racing performances (Li et al., 2014). These studies 416 found six different SNPs in MSTN: two SNPs (g.26T>C and g.156T>C) in the promoter region, two (g.587A>G and g.598C>T) in the 5'-UTR region, and two 417 418 (g.1485C>T, g.2115A>G) in intron-1 of the equine MSTN, respectively. The SNPs 419 g.587A>G and g.598C>T were novel whereas the others had been previously 420 reported (Petersen et al., 2013).

Baron *et al.* (2012) described a genetic variant in exon 2 in some horse breeds. In
fact, they identified a substitution g.2279A>C in Arabians horses and a substitution
g.2478G>C in the Soraia breed horse.

424 Five polymorphisms (g.66495826T>C, g.66495696T>C, g.66493737T>C,

425 g.66495254C>T and g.66490010T>C) were recently observed (Stefaniuk *et al.*,

426 2016) in four Polish breeds (Arabians, Polish Konik, Hucul and Polish Heavy Draft).

427 The polymorphism g.66495254C>T (also known as g.598C>T), has been described

428 in Chinese horse breeds as well as in Polish Konik and Arabian horse breeds. The

429 g.66493737C>T polymorphism known to predict optimum distance in Thoroughbred

430 horses has been identified in four breeds in Egyptian bloodlines (Bower *et al.*, 2012)

431 which were introduced to Polish bloodstock through Egyptian stallions. The insertion

432 g.66495326_66495327Ins227 has been described for the first time in *MSTN* in

433 Thoroughbred horses. Recently, it has been found in the American Quarter Horse

434 (Petersen *et al.*, 2013), and in the Uruguayan Creole breeds (Dall'Olio *et al.*, 2014).

- 435 In the Quarter Horse breed, the Ins227 in *MSTN* is connected with changes to
- 436 Gluteus medius muscle fibre proportions. The higher Myosin Heavy Chain 2B fibre
- 437 type (fast contracting), is in line with pressure selection in Quarter Horse breed for

racing performance (Petersen *et al.*, 2013). A summary of the detected genetic
variants in horse is reported in Table 4.

440

441 Myostatin polymorphisms in pig

Jiang *et al.* (2002) reported three SNPs in porcine *MSTN* T>A, G>A and C>T, in the promoter, intron 1 and exon 3, respectively. Only one mutation (T to A) located in the region 383bp upstream of translation initiation site of porcine *MSTN* was associated with average daily gain in the growing period (from 60 to 100 kg of live weight) in Yorkshire pigs. Furthermore BW in pig with the heterozygous mutation (no AA was found) was increased (Jiang *et al.*, 2002).

Stinckens et al. (2008) compared the MSTN sequence of Belgian Piétrain, which 448 449 shows a heavily muscled phenotype with five other breeds (Piétrain, Landrace, Large 450 White, Meishan and Wild Boar). Fifteen polymorphic loci were found, three of which 451 were located in the promoter region (g.435G>A, g.447A>G, and g.879T>A), five in 452 intron 1 and seven in intron 2. The SNP g.879T>A only appears in Chinese Meishan 453 pigs whilst the polymorphism located at position 447 of the porcine MSTN promoter 454 had a very high allele frequency in the Piétrain pig breed. A g.447A>G mutation 455 which is associated with the expression of the porcine *MSTN* occurs at the putative 456 myocyte enhancer factor 3 (MEF3) binding site on the negative DNA strand. This 457 mutation disrupts a putative MEF3 binding site (Stinckens et al., 2008). 458 However, these results suggest that naturally occurring *MSTN* genetic variants 459 identified thus far in pigs do not have significant association with muscle phenotypes. 460 Nevertheless, a recent work, using an experimental approach has shown the role of 461 MSTN in the development of muscle in pigs. Qian et al. (2015) generated MSTN-

462 deficient Meishan pigs using zinc finger nucleases (ZFN) technology coupled with

463 somatic cell nucleus transfer. The resulting offspring show remarkable DBM 464 phenotype especially pronounced in the hindguarters. Muscle in the MSTN null pig increased mass by 50-100%. Incredibly the muscle fibre size in the null pigs was 465 466 smaller than the wild type. All the increase in mass could be attributed to fibre hyperplasia whereby some muscles from the null had twice the fibre number 467 468 compared to wild type. The animals displayed good overall health. As the technology 469 employed did not involve the introduction of any genetic material in to the genome 470 (e.g. selection markers), Qian et al. (2015) suggest that it is essentially the same as double muscle cattle which are used for human consumption. 471 472 A summary of the detected genetic variants in pigs is reported in Table 5.

473

474 Myostatin polymorphisms in rabbit

475 Fontanesi *et al.* (2011) investigated the variability of the effects of *MSTN*

476 polymorphisms on rabbit production traits. Four single SNPs have been identified by

477 comparative sequencing of 14 rabbits representing breeds or lines having different

478 conformation and muscle mass: one rare synonymous SNP in exon 1 (c.108C>T),

479 one synonymous SNP in exon 2 (c.713T>A), one SNP in the 3'-untranslated region

480 (c.*194A>G) and another SNP in intron 2 (c.747+34C>T) in Belgian hare, Burgundy

481 fawn, Checkered giant and Giant grey.

In commercial hybrids, Qiao *et al.* (2014) detected a SNP (T to C) in the 5' regulatory
region, but no mutation sites were detected in the exons. The correlation analysis
showed that the mutation was associated with increased liver and carcass weight.
These results suggest that the mutations in the upstream regulatory region of the *MSTN* are beneficial to the rabbit soma development, and the mutations can be used
as molecular markers for the selection of the meat quality in rabbits. Sternstein *et al.*

488 (2014) found polymorphisms in the *MSTN* in Giant Grey and NZ White breeds.

489 Comparative sequencing of these breeds revealed two SNPs located in the

490 regulatory region of the rabbit *MSTN* (c.-125T>C) and in intron 1 (c.373+234T>C).

491 A summary of the detected genetic variants in rabbit is reported in Table 6.

492

493 Myostatin polymorphisms in poultry

494 In chickens MSTN maps to 7p11 (Sazanov et al., 1999), and like that of mammals is 495 composed of three exons (373 bp, 374 bp and 1567 bp, respectively) and two 496 introns. Gu et al. (2003) showed poultry MSTN not only regulates skeletal muscle 497 development, but also participates in the fat metabolism and disposition. This 498 research team identified seven SNPs: five were in the 5'-regulatory region (G167A, 499 T177C, G304A, A322G, and C334T) and two were in the 3'-regulatory region of 500 different chicken lines. These last two SNPs in the 3'-regulatory region of the MSTN 501 are A to T (7263) and A to G (6935). Ye et al. (2007) studied the association of MSTN 502 polymorphism with mortality rate, growth, feed conversion efficiency, ultrasound 503 breast depth, breast percentage, eviscerated carcass weight, leg defects, blood 504 oxygen level, and hen antibody titer to the infectious bursal disease virus in three commercial broiler chicken lines. The MSTN had pleiotropic effects on broiler 505 506 performance. This conclusion was reached by the discovery of fourteen SNPs: seven 507 genetic variants in exon 1 (G2100A, G2109A, G2244C, A2283G, C2346T, C2373T, 508 A2416G), one in exon 2 (T4842G), three in exon 3 (C7434G, A7435G, C7436A), and three in intron 1 and 2 (A4405C, A4405T and A4954G). 509

510 As the main function of MSTN is the regulation of skeletal muscle growth, Ye *et al.*

511 (2007) deemed that the non-synonymous SNP T4842G is associated with an amino

512 acid change in the *MSTN* and it could be responsible for variability in body weight.

513 The Bian chicken breed raised for dual purposes, is an important Chinese breed and 514 has a 234G>A in exon 1 of the MSTN (Zhang et al., 2012). Other Chinese chicken breeds (Jinghai, Youxi, and Arbor Acre) have shown four new mutations (A326G, 515 516 C334G, C1346T, G1375A) that were located in the 5'-regulatory region (Zhang et al., 517 2012). Further studies on the growth traits show that the SNPs in chicken *MSTN* may 518 affect the abdominal fat weight and percentage, breast muscle weigh and 519 percentage, birth weight, and adult weight (Zhang et al. 2012). Zhiliang et al. (2004) 520 identified three SNPs in the 5' regulatory region and two SNPs in the 3' regulatory region, and these differed in allele frequencies between breeds. They found that in 521 522 an F2 generation from a cross of broiler and silky chickens, homozygous genotypes AA and BB at a locus in the 5' regulatory region have a higher abdominal fat weight 523 524 and abdominal fat percentage than AB genotype (Zhiliang et al., 2004). The 525 upstream promoter region of *MSTN* was analysed in Wenshang Luhua chicken DNA. 526 Thirteen E-boxes were identified upstream of MSTN and the polymorphisms of E-527 boxes were explored for the first time (Hu et al., 2013). 528 Other interesting studies were carried out on ducks to investigate the association of 529 polymorphisms in MSTN with slaughter traits, breast muscle weight, breast muscle 530 percentage, leg muscle weight and leg muscle percentage. Analysis of the 5' 531 regulatory region of the MSTN showed that polymorphisms (753G>A, 658G>T and 532 235G>C) were associated with the breast muscle percentage and abdominal fat rate 533 (Lu et al., 2011). Furthermore Xu et al. (2013) studied polymorphisms in Pekin duck, and identified three significant variations. The first is a transition T to C in the ORF 534 535 (position 129) and revealed an association with breast muscle thickness. The second 536 SNP was located at 708 bp for the T/C mutation in the ORF and last 952T<C had a 537 significant association with the "Fossilia Ossis Mastodi, or dragon bone" length. In

538 Gaoyou ducks, a transition G>A at 2701bp in exon 3 of the MSTN is correlated with 539 the abdominal fat rate (Liu et al., 2012). In Sansui duck, six SNPs were identified in 540 the first and the third exons (g.106G>A, g.120A>G, g.159G>A, g.5368G>A, 541 g.5389A>C and g.5410G>A) with four loci seemingly associated to leg muscle 542 weight, leg muscle percentage and dressing percentage (Zhao et al., 2016). 543 A summary of the detected genetic variants in poultry is reported in Table 7. 544 545 Myostatin and future implications 546 According to some investigators, MSTN mutations are the main cause of

547 hypertrophy, with a lesser roles played by other gene mutations (Kobolák & Gócza,

548 2002). Inactivation of *MSTN* has therefore been proposed to be a strategy for

549 improving muscle growth of food animals and treating human diseases associated

550 with muscle weakness and dystrophy (Chen & Lee, 2016).

Research, especially on mice, has highlighted the potential of manipulating MSTN signalling in order to promote muscle growth. In null mutants of this species, some muscles are approximately three times their normal weight. Impressive as they are, muscle enlargement in large mammals carrying a null mutation in the same gene, to our knowledge, do not approach this level of muscle growth. Therefore it is important to ascertain the molecular basis underpinning these different responses with a view of translating these findings into increased meat production.

558 One picture that emerges through this review is that mutations that compromise

559 MSTN function have a consequence during development and give rise to

560 supernumerary muscle fibres (hyperplasia). However, one of the clear differences

561 between mice and large animals (cattle and pigs) is the post-natal phenotype. Mice

show considerable fibre hypertrophy whereas in both cattle and pigs display no

563 increase in fibre size. These findings need to be used as a benchmark for future work 564 on doubling muscle in large animals. First and foremost is the need to understand the basis of muscle growth in large mammals. Here it is very important to use the correct 565 566 terms to describe the phenotype of animals, as often this can lead to misinterpretations regarding mechanism. Often DBM animals are referred to as being 567 568 'Hypertrophic'. However this could infer fibre enlargement. As we have discussed, 569 especially in the case of cattle and pig, there is no fibre enlargement. We suggest 570 that accurate mechanistic descriptors are used when they have been precisely established and without this proof a more generic term needs to be applied. We 571 572 suggest the use of the four following terms: 1) Muscle enlargement through hyperplasia; 2) Muscle enlargement through hypertrophy; 3) Muscle enlargement 573 574 through hyperplasia and hypertrophy; 4) Muscle enlargement through unknown 575 cellular mechanisms.

576 Research is required to understand the mechanisms that underpin the role of MSTN 577 in post-natal muscle development in mammals, to answer the question as to why in 578 the absence of MSTN, fibres from mice undergo enlargement, whereas those from 579 large mammals do not. For a number of years the naturally occurring mutants in 580 cattle were our only reference model for large animals lacking MSTN. The lack of 581 fibre hypertrophy was usually explained by the presence of a secondary (to date 582 unidentified) modifying mutation that interfered with the post-natal effect but sparred 583 the pre-natal phenotype. However the work by Qian et al. (2015) in the pig which 584 targets only the MSTN undermines the modifying gene idea. Therefore loss of 585 function mutation in both small and large animals leads to hyperplasia. However it is 586 only in mice that the mutation has an effect on muscle fibre size where it presents as 587 hypertrophy.

588 Clues to resolving this issue come from recent work in monkeys which shows that 589 MSTN and Activin act synergistically to inhibit fibre hypertrophy during adult life 590 (Latres et al., 2017). Based on these findings we suggest that muscle fibres of both 591 cows and pigs are sensitive to Myostatin/Activin signalling, in a similar manner to 592 monkeys. But the issue that still needs to be resolved is why do fibres in adult cows 593 and pigs fail to enlarge in the absence of MSTN. The most parsimonious explanation 594 is that there is a partial redundancy relationship between MSTN and Activin; in the 595 absence of MSTN, the expression levels of Activin become elevated to such a 596 degree that in cows and pigs the latter can completely cover the loss of the former. 597 Examples of gene expression compensation by related molecules, similar to our 598 proposal are abound in mammalian biology (Barbaric et al., 2007). One of the best 599 examples comes through the investigations of MRFs where genetic inactivation of 600 MyoD results in an up-regulation of the related gene-Myf5 (Rudnicki *et al.*, 1992). 601 The hypothesis outlined above has a number of important implications. Our assertion 602 of why the relationship between MSTN and Activin in cows and pigs is only partial 603 and not complete, come from the fact that loss of MSTN has some phenotypic 604 consequence (hyperplasia). Therefore compensation through an up-regulation of 605 Activin expression cannot have occurred during pre-natal life. The second implication 606 is that if there is a redundancy mechanism in mice, which must be very muted since 607 these animals develop a profound phenotype both during pre-natal and adult life. Our suggestions can be validated by quantifying the levels of MSTN and Activin at 608 609 different developmental stages in both large and small animals, an avenue now 610 possible following the development of specific ELISA for MSTN and Activin (Latres et 611 *al.*, 2017).

612 For the meat industry and for the human health sector who focus on muscle growth, 613 the hypothesis outlined here advocates a strategy of dual MSTN and Activin 614 antagonism to promote the growth of the tissue. This could be achieved through the 615 use of a combination of molecules that specifically antagonise the activity of MSTN 616 and Activin (antibodies or protein specific propeptides) or a single protein which acts 617 at a signalling convergence point (at the receptor level through the deployment of a 618 ligand trap or blocking antibody (Omairi et al., 2016, Lach-Trifilieff et al., 2014). 619 Moreover for beef production it will be very interesting to better understand the role of 620 MSTN in adipogenesis; Deng et al. (2017) in fact reported that muscle and adipose 621 tissue develop from the same mesenchymal stem cells, and researchers have found 622 that MSTN is expressed in fat tissues and plays a key role in adipogenesis. 623 Finally *MSTN* is a prime target for transgenic approaches aimed at enhancing meat 624 production in livestock (Georges, 2010). Possible strategies for this outcome include 625 the generation of MSTN knock-out animals. Also more elaborate transgenic 626 approaches, such as targeting post-natal or sex specific inhibition of MSTN need to 627 be considered. Wang et al. (2017), reported the successful application of the CRISPR/Cas9 system to engineer the goat genome through micro-injection of Cas9 628 629 mRNA and sqRNAs targeting *MSTN* in goat embryos. They demonstrate the utility of 630 this approach by disrupting MSTN, resulting in enhanced body weight and larger 631 muscle fiber size in Cas9-mediated gene modified goats. MSTN activity can also be 632 modified using non-genetic approaches using for example blocking antibodies or 633 ligand traps.

634

635 **Conclusions**

636 One picture that emerges through this review is that mutations that compromise

MSTN function have a consequence during development and give rise to
supernumerary muscle fibres (hyperplasia). However, one of the clear differences
between mice and large animals (cattle and pigs) is the post-natal phenotype. First
and foremost there is the need to understand the basis of muscle growth in large
mammals.

This review landscapes the genetics of DBM in mammalian species and chicken and demonstrates the huge number of genetic variants present in animals of commercial interest. It also highlights areas where greater research is required in order for progress to be made concerning the role of MSTN in the regulation of muscle development in economically important animals. Knowledge of null alleles and

647 polymorphisms in *MSTN* are of great interest in the animal breeding field and could

648 be utilized to improve the selection for meat production in livestock animals.

649

650 **Conflict of interest**

The authors have no conflict of interest to declare.

652

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656

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964 LEGENDS TO FIGURES

Figure 1 Myostatin action during myoblast proliferation and differentiation (modified 965 966 from Langley et al., 2002). Retinoblastoma protein (Rb), in a low phosphorylated 967 state, inhibits cell division. Rb activity is attenuated due to hyper-phosphorylation by 968 the kinase action of CKD2. However the activity of CDK2 is inhibited by p21 which is 969 induced by the action of MSTN. MSTN also activates Smad2/3 signalling which 970 inhibits the expression of *MyoD* which is needed for normal myoblast differentiation. 971 In the absence of MSTN, the activity of CDK2 is not inhibited which allows it to 972 inactivate Rb resulting in increased proliferation of myoblasts. At the same time the expression of MyoD is no longer inhibited by Smad2/3 signalling pathways allowing it 973 974 to promote differentiation of the extranumerary myoblasts.



975

Table 1 Polymorphisms on *Myostatin* gene in cattle.

Breed	Polymor on M	rphisms ISTN	Reference
	position	mutation	
Asturiana de los Valles	nt821	DEL11	981 Grobet <i>et al.</i> , 1997 ₈₂
Belgian Blue	nt821	DEL11	McPherron & Lee, 1987
Blonde d'Aquitaine	nt821 nt3811	DEL11 T>G	Kambadur <i>et al.</i> , 1997 Bouyer <i>et al.</i> , 2014
Charolaise	nt610	C>T	Kambadur <i>et al.</i> , 19987
Gasconne	nt938	G>A	Kambadur <i>et al.</i> , 1997 Dunner <i>et al.</i> , 2003
Limousine	nt821 nt610 g.433	DEL11 C>T C>A	Kambadur <i>et al.</i> , 1997 Cappuccio <i>et al.</i> , 1998 Sellick <i>et al.</i> , 2007 ₉₄
Maine-Anjou	nt419 nt676	del-7-ins10 G>T	McPherron & Lee, 1997 Grobet <i>et al.</i> , 1997 ₉₆
Marchigiana	g.874	G>T	Cappuccio <i>et al.</i> , 1998
Nellore	nt76 nt111 nt267 nt374 nt414 nt420 nt433 nt445 nt527 nt641 nt694 nt840 nt951 nt1083	A>T G>T A>G DEL16 C>T T>G A>T A>T T>A G>A G>A G>A A>G T>G C>T	999 Grisolia <i>et al</i> ., 2009
Parthenoise	nt821	DEL11	Kambadur <i>et al.</i> , 1997
Piedmontese	nt938	G>A	Kambadur <i>et al</i> ., 1997
Rubia Gallega	nt821	DEL11	Kambadur <i>et al.</i> , 1997

Brood	Polymorphisms on <i>MSTN</i>		Poforonoo
Breed	position	mutation	Reference
Texel	g.6723 g+391 g.2449 g.2379 g.1405 g.1402 g.1214 g.1129 g.41 g.39 g+474 G+613 g+616 g+619 g+622 g+632 g+632 g+696 g+3135 g+4036 g+4044	G>A G>T C>G C>T A>T G>A C>T C>T C>T T>C G>A T>C G>T C>T C>T C>T C>T C>T T>C C>T C>T T>C C>T T>C C>T T>C T>C T>C T>C T>C T>C T>C T>T C>T T>C T>T C>T T>C T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>C T>T C>T T>T C>T T>T C>T T>T C>T T>T C>T T>T C>T T>T	Kijas <i>et al.</i> , 2007
Norwegian White	c.960	DEL1	Wang <i>et al.</i> , 2016
Sheep	c.2360	G>A	
New Zealand Romney	c.101 c959 c784 c.373+18 c.373+563 c.373+607 c.374-654 c.374-54 c.748-54 c.*83 c.*455 c.*709 c.*123A c2449 c2379	G>A C>T A>G A>G G>A T>C T>C A>G C>A INSA T>G G>C T>C	Wang <i>et al.</i> , 2016 Kijas <i>et al.</i> , 2007
Charollais	c.*123A		Kijas <i>et al.</i> , 2007
White Suffolk	c.*123A		Kijas <i>et al.</i> , 2007
Poll Dorset	c.*123A		Kijas <i>et al</i> ., 2007
Lincoln	c.*123A		Kijas <i>et al.</i> , 2007

Table 2 Polymorphisms on *Myostatin* gene in sheep.

Indian sheep	c.539	T>G T⊳A	Pothuraju <i>et al.</i> , 2015
	c 373+396	T>C	
	c.374-362	A>T	
	c.374-16	DELT	
Stavropol Merino	c.747+185	C>A	Trukhachev et al., 2018
	c.748-194	C>A	
	c.782_783	INST	
	c.940	G>T	
	c.*310	G>T	

1003	Table 3	Polymor	phisms on	Mvostatin de	ne in goat
1005		i orynnor		my oolaan go	no in goui

Table 3 Polymorphisms on <i>Myostatin</i> gene in goat.					
Breed	Polymorphisms on <i>MSTN</i>		Reference		
	position	mutation			
Anhui white	g.197 nt345	G>A A>T			
Boer	nt1256 g.197 nt1388 nt345 nt298	TTTA/- G>A T>A A>T T>C	Zhang <i>et al.</i> , 2013 Nguluma <i>et al.</i> 2018		
Haimen	nt1256	TTTA/-			
Motou	nt1256	TTTA/-			
Nubi	nt1256	TTTA/-			

Table 4 Polymorphisms on *Myostatin* gene in horse.

Duesd	Polymorphis	Poforonoo	
Breed	position	mutation	Reference
American Quarter	g 66495326 66495327	INS227	Petersen <i>et al.</i> 2013
Horse	9.00+00020_00+00027		
	g.26		
	g.150 g.1634	T>G	
Andalusian	a.2024	G>A	Dall'Olio <i>et al.</i> , 2010
	a2115	A>G	
	g.2327	A>C	
	g.4230	T>A	
	g.2279	A>C	Deren et el 2012
Arabians horses	g.66495696	T>C	Stofopiuk at al. 2016
	g.66495254	C>T	Stefanliuk et al., 2010
Bardigiano	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Haflinger	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
	g.26	T>C	
Hucul	g.66495696	T>C	Stefaniuk <i>et al.</i> , 2014
nacai	g.66493737	T>C	Stefaniuk <i>et al.</i> , 2016
	g.66490010	T>C	
Italian Saddle	g.26	T>C	Dall'Olio <i>et al.</i> . 2010
	g.156		
Italian trotter	g.26		Dall'Olio <i>et al.</i> , 2010
	g.66495254		
Dolioh Konik	g.66493696		Stefaniuk <i>et al.</i> , 2014
	g.00493737 g.66495254		Stefaniuk <i>et al.</i> , 2016
	g.66490010		
Lipizzan	g.00100010	T>C	Dall'Olio et al. 2010
Maremmano	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Murgese	a.156	T>C	Dall'Olio <i>et al.</i> , 2010
	g.26	T>C	
NOTIC	g.156	T>C	Dall Olio <i>et al.</i> , 2010
	g.26	T>C	
	g.66495254	C>T	Stofaniuk at al 2014
Polish Heavy Draft	g.66495696	T>C	Stefaniuk <i>et al.</i> 2014
	g.66493737	T>C	
	g.66490010	T>C	
Rapid Heavy Draft	g.26	T>C	Dall'Olio <i>et al.</i> . 2010
	g.156		
Salernitano	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Soraia	g.2478	G>C	Baron <i>et al.</i> , 2012
	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Thoroughbred	g.1634	T>G	Petersen <i>et al.</i> , 2013
horse	g.2115	A>G	Petersen <i>et al.</i> , 2013
	g.2327	A>C	Petersen <i>et al.</i> , 2013

Tolfetano	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Uruguayan Creole	g.156	T>C	Dall'Olio <i>et al.</i> , 2010
Ventasso	g.26	T>C	Dall'Olio <i>et al.</i> , 2010

Polymorphisms on MSTN on mutation Breed Reference position g.435 G>A **Belgian Pietrain** g.447 Stinckens et al., 2008 A>G g.879 T>A **Chinese Meishan** T>A Qian *et al.*, 2015 g.879 T>A nt383 exon 3 Yorkshire pig Jiang *et al.*, 2002 (position no G>A specified) C>T

1008 **Table 5** Polymorphisms on *Myostatin* gene in pig.

Table 6 Polymorphisms on *Myostatin* gene in rabbit.

Breed	Polymorphisms on <i>MSTN</i>		Reference
	position	mutation	
	c.108	C>T	
Belgian hare	c.713	T>A	Fontanesi <i>et al</i> 2013
	c.*194	A>G	i ontanesi et al., 2013
	c.747+34	C>T	
	c.108	C>T	
Burgundy fawn	c.713	T>A	Fontanesi et al. 2013
	c.*194	A>G	
	c.747+34	C>T	
	c.108	C>T	
Checkered giant	c.713	T>A	Fontanesi et al. 2013
	c.*194	A>G	i ontariesi et al., 2013
	c.747+34	C>T	
Commercial			
breeds (not	nt476	T>C	Qiao <i>et al.</i> , 2014
specified)			
	c.108	C>T	
Giant grov	c.713	T>A	Fontanesi <i>et al.</i> 2013
Giant grey	c.*194	A>G	
	c.747+34	C>T	
Giant Grev	c125	T>C	Sternstein et al 2014
Giant Grey	c.373+234	T>C	
New Zealand	c125	T>C	Stornstoin at al 2014
White	c.373+234	T>C	

1014	Table 7 Pol	ymorphisms on	Myostatin	gene in poultry
1014		ymorpmomo on	wyoolain	gone in pound

Breed	Polymorphisms Breed on MSTN		Beference
	position	mutation	
Arbor Acre	nt167 nt177 nt304 nt322 nt326 nt334 nt334 nt1346 nt1375 nt6935	G>A T>C G>A A>G A>G C>T C>G C>T G>A A>G	Gu <i>et al.</i> , 2003 Zhang <i>et al.</i> , 2012
	nt7263	A>T	
Bian chicken	nt234	G>A	Zhang <i>et al.</i> , 2012
Gaoyou ducks	nt2701	G>A	Liu <i>et al.</i> , 2012
Jinghai	nt326 nt334 nt1346 nt1375	A>G C>G C>T G>A	Zhang <i>et al</i> ., 2012
Pekin duck	nt129 nt708 nt952	T>C T>C T>C	Xu <i>et al</i> ., 2013
Sansui duck	g.106 g.120 g.159 g.5368 g.5389 g.5410	G>A A>G G>A G>A A>C G>A	Zhao <i>et al.</i> , 2016
Youxi	nt326 nt334 nt1346 nt1375	A>G C>G C>T G>A	Zhang <i>et al.</i> , 2012