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Abstract

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Keywords

myostatin, SNP, growth, mortality, broiler chicken

Disciplines

Agriculture | Animal Sciences | Genetics and Genomics | Poultry or Avian Science

Comments

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Associations of myostatin gene polymorphisms with performance and mortality traits in broiler chickens

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myostatin / SNP / growth / mortality / broiler chicken

1. INTRODUCTION

Myostatin is a member of the transforming growth factor-beta (TGF-beta) superfamily and highly conserved in gene structure among vertebrate species [15, 16]. The myostatin gene is highly polymorphic. Nineteen single

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nucleotide polymorphisms (SNPs) and 20 haplotypes have been detected in 28 European bovine breeds, of which some were breed-specific [6]. Baron *et al.* [1] identified seven SNPs and one deletion in exon 2 of the myostatin gene in broiler and/or layer chicken lines. Gu *et al.* [12] identified three SNPs in the 5'-regulatory region and two SNPs in the 3'-regulatory region in the chicken, and these differed in allele frequencies between breeds.

Myostatin is mainly expressed in muscular tissues. Several studies have reported that myostatin negatively regulates skeletal muscle growth. Muscle weights of mice lacking the myostatin gene are about two to three times greater than those of wild type mice [18]. A loss-of-function mutation in the myostatin gene causes double muscling phenotypes in Belgian Blue and Piedmontese cattle [17]. Myostatin negatively regulates activation of satellite cells that promote postnatal muscle growth and repair in mice [16]. Overexpression of myostatin during infection with chronic infectious respiratory diseases has been shown to reduce skeletal muscle growth in pigs [7]. In humans, a loss-of-function mutation in the myostatin gene in children increases muscle bulk and strength [21]. The function of myostatin in negative regulation of skeletal muscle growth is modulated by additional genetic factors, as illustrated by the finding that myostatin protein levels are not significantly different between broilers and layers [11]. Other genes have been shown to affect muscularity in the presence of *Mstn*^{Cmpt-dl1Abc} (a 12-bp deletion in the myostatin gene) in mice [23]. Potts *et al.* [19] reported 19 genes to be differentially expressed between normal and double-muscled bovine embryos.

Myostatin gene polymorphisms have effects on multiple traits. Mutations in myostatin regulatory regions have been shown to be associated with abdominal fat weight, abdominal fat percentage, birth weight, breast muscle percentage and breast muscle weight in chickens [12]. Furthermore, Casas *et al.* [5] found that calves with two copies of the inactive allele of myostatin were more likely to die before weaning, were heavier at birth and leaner, and had more muscle mass than animals with zero or one copy; heterozygous calves had the heaviest bodyweight at weaning and the highest live weight; calves with zero copy were the highest in fat content.

The objectives of the current research were to identify genetic polymorphisms in the myostatin gene in elite commercial broiler chicken lines and to determine their associations with growth, mortality, yield and support traits in high- and low-health environments. Results of this study will increase our understanding of the many functions of the myostatin gene.

2. MATERIALS AND METHODS

2.1. Populations and traits

About 100 sires from each of three elite commercial broiler chicken lines (coded X, Y and Z) were used. Mortality and performance data were recorded on 17 to 437 progeny from each sire (Tab. I). A sire's progeny were raised in one high hygiene environment (HH) or in one of two low hygiene environments (LH: LH1 and LH2). Traits recorded (Tab. I) in all three environments were body weight at 7 and 40 days of age (BW7 and BW40) and mortality from hatching to 14 days (EMORT) and from 14 to 40 days of age (LMORT). Nine additional traits were recorded in the HH environment: ultrasound breast depth (US), percentage of breast (BR), feed conversion efficiency (FCR), eviscerated carcass weight (EV), three leg-defect related traits (measured on a subjective scale by trained technicians) [twisted legs or evident tibial dischondroplasia (Leg), X-ray-inspection-based sub-clinical or incipient development of tibial dischondroplasia (Lixi), and curly or crooked toes and/or bowed legs (Tobo)], and blood oxygen content measured by pulse oximeter (Oxi) and female's antibody titer to infectious bursal disease at 27 wks (IBD).

2.2. DNA samples, identification of SNPs and genotyping

Blood samples from each sire were individually collected in tubes containing EDTA and immediately frozen at -20°C . Individual genomic DNA was extracted from a $15\ \mu\text{L}$ aliquot of blood with DNAzol reagent (Invitrogen, Carlsbad, California, USA) following the manufacturer's protocol. DNA samples were quantified by spectrophotometer and visualized in agarose gel. One hundred ng of genomic DNA was used as a PCR template.

Five pairs of gene-specific primers were designed based upon published myostatin gene sequence (GenBank accession number: AF346599). Primer sequences and their annealing temperature are in Table II and were used to amplify the coding region, part of the promoter and intron regions of myostatin. Three primer pairs, *MSTpr*, *MSTex1* and *MSTex3*, were used to identify and genotype SNPs. Primers *MSTex2-1* and *MSTex2-R* were used to identify the SNP that was genotyped using primers *MSTex2-2* and *MSTex2-R*. Independent amplifications were conducted for individual DNA samples from each line using a high fidelity Taq DNA polymerase (Invitrogen, Carlsbad, California, USA). PCR products were cloned using Topo Cloning Kit (Invitrogen, Carlsbad, California, USA) and 192 plasmids clones from each line and myostatin region were sequenced using the BigDye terminator kit on an

Table 1. Trait abbreviations, definitions and numbers of sires genotyped and progeny evaluated in three elite commercial broiler breeding lines (X, Y, and Z).

Progeny group	Trait	Abbreviation	Description	# sires per line			Mean # of progeny per sire in line		
				X	Y	Z	X	Y	Z
Low hygiene (LH1)		BW7-LH1	body weight at 7 days (g)	82	69	92	94	67	41
		BW40-LH1	body weight at 40 days (dag)	82	69	92	94	68	41
		EMORT-LH1	mortality prior to day 14 (%)	82	69	92	116	92	50
		LMORT-LH1	mortality after 14 days (%)	82	69	92	116	92	50
Low hygiene (LH2)		BW7-LH2	body weight at 7 days (g)	102	99	113	93	65	66
		BW40-LH2	body weight at 40 days (dag)	102	99	113	93	65	70
		EMORT-LH2	mortality prior to day 14 (%)	102	99	113	105	77	78
		LMORT-LH2	mortality after 14 days (%)	102	99	100	105	77	21
High hygiene (HH)		BW7-HH	body weight at 7 days (g)	101	100	113	379	309	210
		BW40-HH	body weight at 40 days (dag)	101	100	113	383	310	211
		EMORT-HH	mortality prior to day 14 (%)	101	100	113	437	335	248
		LMORT-HH	mortality after 14 days (%)	101	100	113	437	335	248
	US-HH	ultrasound breast depth at 5 wks (mm)		99			88		
	BR-HH	percentage of breast at 6 wks (%)	92	92	110	45	41	55	
	EV-HH	percentage of eviscerated yield at 6 wks (%)	92	92	110	35	40	56	
	FCR-HH	feed conversion ratio between 6 and 8 wks	92	94	110	28	29	61	
	IBD-HH	hen antibody titer to infectious bursal disease at 27 wks	70	0	74	19	0	17	
	Leg-HH	score for twisted legs or evident tibial dyschondroplasia (TD) at 5 wks	101	100	113	383	310	211	
	Lixi-HH	score for X-ray-inspection-based sub-clinical or incipient development of TD (Lixiscope) at 5 wks	93	96	102	31	31	26	
	Tobo-HH	score for curly or crooked toes and/or bowed legs at 5 wks	101	100	113	383	310	211	
	Oxi-HH	oximeter estimate of blood oxygen at 5 wks (%)	91	0	114	28	0	118	

ABI 3100 automated sequencer (Applied Biosystems Foster City, CA, USA). Phred/Phrap and Consed programs were used for sequence quality evaluation, contig assembling and SNP inspection, as previously described by Gordon *et al.* [8]. Clustal-W software [22] and Vector NTI (Invitrogen, Carlsbad, California, USA) were used for sequence alignments, translation and polymorphism identification.

Individual sires were genotyped for five of the identified SNPs using PCR-RFLP methods. These five SNPs were chosen based on the feasibility and cost of genotyping using PCR-RFLP, partially in view of their potential future routine application in breeding programs. The PCR reaction was performed in 55 μL of reaction solution with 0.025 units/ μL Taq (Invitrogen Ltd., Paisley, UK), 1.5 mM Mg, 200 μM dNTPs, 0.5 μM primers, and 5 μL DNA sample (25 ng/ μL). Cycling conditions for PCR were 94 °C 2 min for 1 cycle, 94 °C 30 s, annealing temperature (Tab. II) 30 s and extension 72 °C 30 s for 40 cycles, and extra extension 72 °C for 2 min. Digestion of PCR products was performed in a 12 μL of reaction solution following manufacturer's directions (New England Biolabs, Ipswich, MA, USA) and other conditions (Tab. II). DNA fragments were electrophoretically separated on 3% Metaphor gel (Cambrex Bio Science Wokingham, Ltd., Berkshire, UK).

2.3. Statistical analysis

Associations of polymorphisms with traits were identified based on analysis of the mean adjusted performance of progeny of each sire in a given environment. These were derived from progeny performance records that were each adjusted for the systematic environmental effects of sex, age, hatch and mating group, and for mate effects (half the EBV of the dam) by subtracting solutions obtained from routine animal model genetic evaluation procedures that are used within these lines. Association analyses were conducted using weighted least squares of SAS[®] PROC GLM (SAS[®] 9.1), with weights equal to the number of progeny included in the adjusted mean to account for differences in variance of residuals. Only additive associations were evaluated because progeny means primarily reflect the additive effects of genes, being related to half the sire's breeding value. Association analyses were performed separately for single SNPs and for haplotypes of SNPs. In addition, an outlier analysis was performed. Twenty-one possible outlier data points were identified in the 14 traits by visual inspection of distributions of progeny means against numbers of progeny. Comparison of results from analyses with all data points with those after removal of the outliers showed that the identified outliers had minor

Table II. Primer sequences, annealing temperatures (T_m), single nucleotide polymorphisms (SNPs), and digestion conditions for restriction enzymes.

Primer name	Primer sequence	T _m (°C)	Restriction digestion condition	
			SNP	Enzyme °C Time (h)
<i>MSTpr</i>	Forward	60	A/G	<i>Bst</i> U1 60 8
	Reverse	60	A/G	<i>Hpa</i> I 37 8
<i>MSTex1</i>	Forward	60	C/G	<i>Hin</i> PI 37 8
	Reverse	60	C/T	<i>Bbs</i> I 37 12
<i>MSTex2*</i>	1 Forward	55	G/T	<i>Bsr</i> I 65 8
	2 Forward	55	G/T	<i>Bsr</i> I 65 8
<i>MSTex3</i>	R Forward	64		
	Reverse	64		

* For *MSTex2*, the primer pair *MSTex2-1* and *MSTex2-R* was used to identify SNPs, and the primer pair *MSTex2-2* and *MSTex2-R* was used to genotype SNP G/T with *Bsr* I.

effects on the SNP-trait association results. The results reported in this paper are, therefore, based on all data points in the original dataset.

In all analyses, residuals were assumed uncorrelated. Genetic relationships among sires were limited and were ignored in the analyses; the 102, 100, 114 sires that were analyzed for lines X, Y, and Z originated from a total of 51, 50, and 66 sires and 71, 75, and 93 dams. Thus, half- and full-sib relationships among the evaluated sires were small and ignoring them is not expected to bias results.

Analysis with single SNPs: Three allele substitution effect models were employed to determine associations between an SNP and a trait. Model 1 was for within-line data analysis and Models 2 and 3 were for analyses across lines.

Model 1: $y_i = \mu + b f_i + \varepsilon_i$ (fitted for each line separately).

Model 2: $y_{ij} = L_j + b_j f_{ji} + \varepsilon_{ji}$ (fitted across lines, with separate effects per line).

Model 3: $y_{ij} = L_j + b f_{ji} + \varepsilon_{ji}$ (fitted across lines with a common effect).

where, y_i and y_{ij} are the adjusted progeny mean of the i th sire of line j ; μ is a general mean; L_j is the effect of the j th line ($j = 1, 2, 3$); f_i and f_{ji} represent the number copies of a given allele of the SNP carried by the i th sire; b_j is the allele substitution effect for line j ; ε_i and ε_{ji} are the residual for sire i with variance $= \frac{1}{N_i} \sigma_e^2$ (N_i is the number of progeny for sire i). Significance level for addressing an association was set to be 0.031 corresponding to a 50% proportion of false positive associations (PFP).

Analysis with haplotypes: It is likely that the SNPs within the myostatin gene are in linkage disequilibrium because of their proximity on the chromosome, and the haplotypes formed by these SNPs may be associated with traits of interest. Thus, presence of linkage disequilibrium among myostatin SNPs was determined using a χ^2 test, amount of linkage disequilibrium was measured using r^2 [13], and haplotype frequencies were estimated by maximum likelihood using the software Arlequin version 2.000 [20]. For sires whose haplotypes could not be inferred with certainty, haplotype frequencies were used to assign haplotype probabilities. Models to analyze associations of haplotypes with traits were the same as described above but replacing SNP allele effects by $\sum_{h=1}^{n-1} b_{jh} f_{ijh}$, where f_{ijh} is the probability of haplotype h in sire i (or the sum of the probabilities that the first and second haplotype of the sire are h) and b_h is the substitution effect for haplotype h . In these models, the effect of the most frequent haplotype (GAGCT for each line) was set equal to zero, such that the models were of full rank. As a result, b_h represents the effect of substituting a copy of haplotype GAGCT by a copy of haplotype h . Significance tests for

haplotype effects on traits were performed in two steps: first, the overall haplotype effect for a trait was tested; if this was significant, then the contrast of each haplotype with the most frequent haplotype was tested.

3. RESULTS

3.1. SNPs and haplotypes

Thirteen SNPs were identified in the amplified regions of exons 1, 2 and 3 and introns 1 and 2 in myostatin gene (Tab. III). Of them, five exonic SNPs (*MST2100*, *MST2109*, *MST2244*, *MST2373* and *MST4842*) were genotyped on individual sires using the PCR-RFLP method to determine allele frequencies and associations with phenotypic traits. Allele frequencies of the five SNPs varied by line (Tab. IV). The *MST4842* SNP was fixed in lines Y and Z. All SNPs were in Hardy-Weinberg equilibrium ($P > 0.1$) but in linkage disequilibrium with each other ($P < 0.01$) (Tab. IV). The SNPs at positions 2100, 2109, and 2373 bp in the myostatin gene sequence (GenBank accession number: AF346599) were in complete disequilibrium in lines Y and Z, but an intervening SNP at 2244 bp was in incomplete disequilibrium ($r^2 < 0.84$) with the other SNPs. A total of eight haplotypes segregated in line X but only three of these segregated in lines Y and Z (Tab. IV), which was much fewer than the 32 haplotypes that were possible. Haplotypes GAGCT and GGCCG were prevalent in line X and haplotypes GAGCT and AGCTT were prevalent in lines Y and Z (Tab. IV). Haplotype GAGCT was the most frequent haplotype in all three lines.

3.2. Trait associations of single SNPs

Results for associations of individual myostatin SNPs with traits are in Figure 1. Results in Figure 1 were obtained from analysis with Model 1. Because *MST2373* was in complete disequilibrium with *MST2109* in lines Y and Z, results for the SNP are not shown. Compared with results from Model 1, analyses with Models 2 and 3 seldom provided extra information and are, therefore, not shown. For analyses with Model 2, significant interactions between SNPs and lines were detected when allele effects were significant in at least one line in Model 1. In these cases, the results for individual lines from Model 1 are reported. Analysis with Model 3 detected only one new association, for *MST2100* with Tobo.

Table III. Single nucleotide polymorphisms (SNPs) detected in three elite commercial broiler breeder lines.

Name	SNP	Position in the gene		Line in which detected
		Region	in AF346599 (bp)	
<i>MST2100</i>	G/A	Exon 1	2100	X, Y, Z
<i>MST2109</i>	G/A	Exon 1	2109	X, Y, Z
<i>MST2244</i>	G/C	Exon 1	2244	X, Y, Z
<i>MST2283</i>	A/G	Exon 1	2283	X, Z
<i>MST2346</i>	C/T	Exon 1	2346	X, Z
<i>MST2373</i>	C/T	Exon 1	2373	X, Y, Z
<i>MST2416</i>	A/G	Exon 1	2416	Z
<i>MST4405</i>	A/C, A/T	Intron 1	4405	X (A/T), Y and Z (A/C)
<i>MST4842</i>	T/G	Exon 2	4842	X
<i>MST4954</i>	A/G	Intron 2	4954	X, Y
<i>MST7434</i>	C/G	Exon 3	7219	X
<i>MST7435</i>	A/G	Exon 3	7220	X, Y, Z
<i>MST7436</i>	C/A	Exon 3	7221	X, Y, Z

MST2100: Allele 1 (nucleotide A) of *MST2100* had consistent negative effects on growth in the HH environment across ages and lines, although its effect was significant only on BW7 in line Z (Fig. 1A). Allele 1 also had a negative effect on IBD in lines X and Z and was significant ($P < 0.031$) in line Z. Allele 1 had a non-significant positive effect on Oxi in line Z but a significant ($P < 0.001$) negative effect on Oxi in line X.

MST2109: In lines Y and Z, the effects of allele 1 on the traits were similar in magnitude to those for *MST2100* because of the complete LD between these two SNPs, but the direction of effects was opposite (Fig. 1B). In line X, allele 1 had consistent positive effects on growth in the LH environments, with two of four traits having significant effects ($P < 0.031$). Allele 1 also had a negative effect on Leg ($P < 0.04$), a positive effect on Oxi ($P < 0.043$), and positive effects on LMORT in the LH environments (Fig. 1B).

MST2244: Allele 1 had consistent negative effects on growth in the LH environments in line X and Z and was significant for three of the four growth traits in line X ($P < 0.031$) (Fig. 1C). Allele 1 had a consistent negative effect on LMORT, with one allele effect being significant ($P < 0.01$) in the LH environments in line X (Fig. 1C).

MST2373: *MST2373* had the same associations with the traits as *MST2109* in lines Y and Z because of complete linkage disequilibrium (Tab. IV). In

Table IV. Allele and haplotype frequencies of single nucleotide polymorphisms in the myostatin gene, and amount of linkage disequilibrium among nucleotide polymorphic loci in three elite commercial broiler chicken lines.

Frequencies of alleles and haplotypes					
Locus or haplotype	Allele 1	haplotype	Line X	Line Y	Line Z
<i>MST2100</i> (A/G) ^a	A ^b		0.16	0.52	0.42
<i>MST2109</i> (A/G) ^a	A ^b		0.53	0.478	0.58
<i>MST2244</i> (C/G) ^a	C ^b		0.32	0.48	0.34
<i>MST2373</i> (C/T) ^a	C ^b		0.83	0.48	0.58
<i>MST4842</i> (G/T) ^a	G ^b		0.46	0	0
Haplotype ^c		AGCTT	0.02	0.48	0.34
		AGGCT	0.01	-	-
		AGGTG	0.03	-	-
		AGGTT	0.11	0.04	0.08
		GAGCG	0.12	-	-
		GAGCT	0.40	0.48	0.58
		GAGTT	0.01	-	-
		GGCCG	0.31	-	-
Linkage disequilibrium, measured by r^2					
Line	Locus ^d	<i>MST2100</i>	<i>MST2109</i>	<i>MST2244</i>	<i>MST2373</i>
X	<i>MST2109</i>	0.221 **			
	<i>MST2244</i>	0.045 **	0.537 **		
	<i>MST2373</i>	0.889 **	0.179 **	0.048 **	
	<i>MST4842</i>	0.055 **	0.242 **	0.463 **	0.060 **
Y	<i>MST2109</i>	1 **			
	<i>MST2244</i>	0.840 **	0.840 **		
	<i>MST2373</i>	1 **	1 **	0.840 **	
Z	<i>MST2109</i>	1 **			
	<i>MST2244</i>	0.7 **	0.7 **		
	<i>MST2373</i>	1 **	1 **	0.724 **	

^a SNP and its name. Sequence position is indicated by the number in the locus name.

^b Allele-defining nucleotide, designated allele 1 and whose frequency is shown.

^c Haplotype frequency is the maximum likelihood estimate.

^d Locus sequence position is indicated by the number in the locus name.

** Represents that the linkage disequilibrium between two SNPs was very significant ($P < 0.01$).

line X, allele 1 had a positive effect on mortality in the HH environment, a positive effect on IBD (not significant), and a significant ($P < 0.001$) positive effect on Oxi (Fig. 1D).

MST4842: Associations of *MST4842* with traits were evaluated only in line X because of its fixation in lines Y and Z. In line X, allele 1 had consistent

negative, though partially significant ($P < 0.031$), effects on growth across ages in the LH environments (Fig. 1D). Allele 1 also had negative effects on LMORT in LH environments, though only significant ($P < 0.01$) in LH2.

3.3. Associations of haplotypes with traits

Few significant ($P < 0.1$) haplotype-trait associations were detected (Tab. V) and they generally matched results from the individual SNP analyses. In line X, haplotypes of the five SNPs had consistent associations ($P < 0.029$) with LMORT across environments and ages. Haplotype GAGCT had significantly lower LMORT than haplotype AGGTG in both LH and HH environments. Haplotype GAGCT also had significantly lower EV than AGCTT and GAGCG, but significantly higher EV than AGGTT. In line Y, there were no significant associations of haplotypes with the traits studied. In line Z, myostatin haplotypes had significant associations with BW7-LH1, BW7-HH, IBD-HH, and LMORT-LH2 (Tab. V). Of the three haplotypes in line Z, AGGTT had the highest BW7 in the LH1 environment but the lowest BW7 in the HH environment; GAGCT had the highest IBD-HH; and AGCTT had the lowest LMORT in the LH2 environments.

4. DISCUSSION

The high degree of polymorphism of the DNA sequence of the myostatin gene, which has been demonstrated in previous studies [1, 6, 12], was further displayed here in three elite commercial broiler chicken lines. Of the identified 13 SNPs, 11 were in exons and 2 were in introns. Four SNPs, *MST2283*, *MST2346*, *MST2416* and *MST7434*, resulted in disappearance/appearance of restriction sites that were expensive for PCR-RFLP genotyping. Four SNPs, *MST4405*, *MST4954*, *MST7435* and *MST7436*, did not result in disappearance/appearance of restriction sites. The remaining five of the 13 SNPs were genotyped using PCR-RFLP. The four genotyped SNPs, *MST2100*, *MST2109*, *MST2244* and *MST2373* in exon 1 were synonymous substitutions (GCG → GCA, CCG → CCA, CGC → CGG, GAC → GAT and ACG → ACT, respectively). Therefore, these four SNPs are genetic markers and likely not causative mutations for the traits with which they were found to be associated. The SNP *MST2373* was the same as that previously reported (as nt. 1204 C/T) by Baron *et al.* [1] in two chicken lines that were separately selected for bodyweight and egg production [1]. The SNP *MST4842* in exon 2 was a non-synonymous substitution (CTG → CGG), which resulted

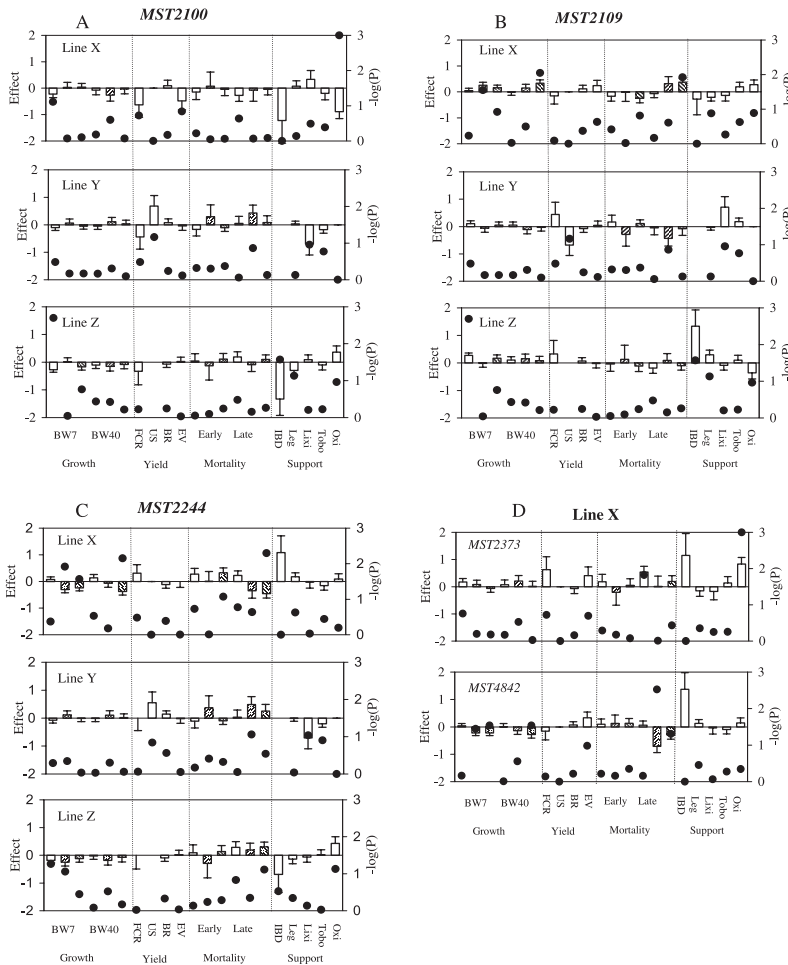


Figure 1. A, B, and C: Estimates of the effect of allele 1 relative to allele 2 on traits for myostatin SNPs *MST2100*, *MST2109* and *MST2244* in three elite commercial broiler lines (X, Y and Z). D: Estimates of effect of allele 1 over allele 2 on traits for SNPs *MST2373* and *MST4842* in line X. The traits on the x-axis are growth traits including body weight at 7 (BW7) and 40 days (BW40); yield traits including feed conversion ratio (FCR), ultra sound (US), breast yield (BR), and eviscerated yield (EV); mortality traits including early and late mortality; and support traits including oximeter (Oxi), score for curly or crooked toes and/or bowed legs at 5 wks (Tobo), score for X-ray-inspection-based sub-clinical or incipient development of TD at 5 wks (Lixi), score for twisted legs or evident tibial dyschondroplasia at 5 wks (Leg) and hen antibody titer to infectious bursal disease at 27 wks (IBD). In the Figure, bar = estimate of effect and its standard error, bullet points = significance levels ($-\text{Log}(P\text{-value})$); the horizontal line (x-axis) = a P -value of 0.031, which corresponds to a proportion of false positives of 0.5.

Table V. Estimates (\pm standard error) of significant ($P^b < 0.1$) associations of myostatin haplotypes with traits in three elite commercial broiler chicken lines.

Line	Trait	Haplotype ^a								P^b
		AGCTT	AGGCT	AGGTG	AGGTT	GAGCG	GAGTT	GGCCG		
X	BR-HH	0.76 \pm 0.60	-	0.97 \pm 0.39*	-0.34 \pm 0.30	0.32 \pm 0.23	0.00 \pm 0.44	-0.14 \pm 0.14	0.094	
	EV-HH	2.27 \pm 1.14*	-	0.39 \pm 0.71	-0.91 \pm 0.45*	1.08 \pm 0.35*	0.28 \pm 0.76	0.04 \pm 0.22	0.009	
	LMORT-LH1	1.39 \pm 1.47	-	1.79 \pm 0.86*	0.53 \pm 0.63	1.00 \pm 0.39*	0.10 \pm 0.69	0.40 \pm 0.30	0.027	
	LMORT-LH2	0.69 \pm 0.58	-0.52 \pm 1.13	1.34 \pm 0.51*	-0.04 \pm 0.25	-0.20 \pm 0.24	1.67 \pm 0.71*	0.41 \pm 0.17*	0.004	
	LMORT-HH	0.00 \pm 0.79	-1.08 \pm 1.56	1.64 \pm 0.64*	0.20 \pm 0.35	0.09 \pm 0.31	1.71 \pm 0.61*	-0.20 \pm 0.22	0.029	
Z	Oxi-HH	-0.64 \pm 0.69	0.38 \pm 1.25	-1.51 \pm 0.80	-1.13 \pm 0.37	-0.08 \pm 0.33	-0.50 \pm 0.42	-0.10 \pm 0.22	0.057	
	BW7-LH1	-0.13 \pm 0.14	-	-	0.59 \pm 0.23*	-	-	-	0.009	
	BW7-HH	-0.24 \pm 0.09*	-	-	-0.42 \pm 0.16*	-	-	-	0.005	
	IBD-HH	-0.98 \pm 0.65	-	-	-2.43 \pm 1.02*	-	-	-	0.037	
	Leg-HH	0.22 \pm 0.18	-	-	0.64 \pm 0.30*	-	-	-	0.083	
LMORT-LH2	-0.25 \pm 0.17	-	-	0.54 \pm 0.31	-	-	-	0.045		

^a The effect of a haplotype is expressed in units of additive genetic standard deviation.^b P -value for test of total haplotype effect.* Haplotype is significantly different from haplotype GAGCT at the $P < 0.05$ level.

in a change of an amino acid (Leu → Arg). Baron *et al.* [1] did not identify *MST4842* but found six other non-synonymous SNPs in exon 2. The five SNPs found in the current study formed eight haplotypes in the broiler chickens; line X had five line-specific haplotypes, of which haplotype GGCCG was most frequent.

Each of the five identified myostatin SNPs was significantly ($P < 0.031$) or consistently associated with bodyweight at day 7 and/or day 40 in at least one broiler line. Associations of the non-synonymous SNP *MST4842* with bodyweight in line X may be due to the amino-acid change in myostatin, although functional studies will be needed to ascertain this. Associations of myostatin polymorphisms with growth have been frequently reported in humans and cattle [2–5, 9, 10, 14, 15, 17, 21]. These imply that, because the main function of myostatin is regulation of skeletal muscle growth, changes in its protein sequence can modify growth function. The detected associations of the synonymous SNPs with growth may be attributed to their linkage disequilibrium with other polymorphisms that exist in regulatory regions or exons of myostatin or to QTL beyond the myostatin gene. Gu *et al.* [12] reported a polymorphic site in the 5′ regulatory region of myostatin, which was associated with hatch weight in F₂ chickens from a broiler by Silky cross [12]. This or other regulatory-region polymorphisms could be in linkage disequilibrium with some of the synonymous SNPs detected in this study, which would explain their significant associations. Identification and further analysis of such polymorphisms were, however, beyond the scope of the present study.

The current study on myostatin SNP associations in chickens measured more extensive and diverse traits than have been reported in most previous studies. In addition to associations with growth, myostatin polymorphisms were also found to be associated with mortality, yield and support traits. This pleiotropic effect of the myostatin gene has also been reported by others in chicken and other species. Gu *et al.* [12] found homozygous genotypes AA and BB at a locus in the 5′ regulatory region to be associated with higher abdominal fat weight and abdominal fat percentage than AB in the F₂ chickens from a cross of broiler and Silky chickens. Casas *et al.* [5] showed that calves with two copies of the inactive allele of myostatin were more likely to die before weaning, that calves with one copy of the inactive allele had leaner and more muscled carcasses than animals without inactive alleles, and that calves with zero copy of the inactive allele were highest in fat content.

Polymorphisms for myostatin have been used in marker-assisted selection in beef cattle but not in chickens. The current results suggest that one non-synonymous SNP (*MST484*) and four synonymous SNPs may have pleiotropic

associations with a set of traits and show genotype by environment interactions. This represents the first full evaluation of pleiotropic effects of myostatin mutations on different traits, but these results require confirmation in larger data sets and other lines. In addition, the causative reasons for the associations are not clear, even for the non-synonymous SNP *MST484*.

Pleiotropic effects and presence of genotype by environment interactions of myostatin can complicate genetic improvement but knowledge about them will allow breeding programs to be structured to minimize their impact or maximize their benefits. Presence of pleiotropic effects will require identification of alleles or haplotypes that have a net beneficial effect across traits. Presence of genotype by environment interactions of myostatin implies that the network that the gene is involved in is sensitive to environmental changes. Knowledge of such genotype by environment interactions is important for genetic selection programs, to ensure that the allele(s) selected for has(have) beneficial effects in the target environment.

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