1	Fine mapping of genes determining extrafusal fiber properties in murine soleus
2	muscle
3	Carroll AM ^{1,2} , Cheng R ³ , Collie-Duguid ESR ^{1,4} ,Meharg C ⁵ , Scholz ME ¹ , Fiering S ⁶ , Fields
4	JL ⁶ , Palmer AA ⁷ , Lionikas A ¹ .
5	
6	Affiliation:
7	¹ School of Medicine, Medical Sciences and Nutrition, University of Aberdeen, Foresterhill
8	Aberdeen, AB25 2ZD, UK
9	² The New Zealand Institute for Plant & Food Research Limited, Batchelar Road, Private Bag
10	11600, Palmerston North 4442, New Zealand
11	³ Research School of Biology, The Australian National University, 134 Linnaeus Way, Acton,
12	ACT 2601, Australia
13	⁴ Centre for Genome Enabled Biology and Medicine, University of Aberdeen, Old Aberdeen,
14	Aberdeen, AB24 3RY, UK
15	⁵ Institute for Global Food Security, Queen's University Belfast, 18-30 Malone Road, Belfast,
16	BT9 5BN, UK
17	⁶ Department of Microbiology/immunology, Geisel School of Medicine at Dartmouth,
18	Hanover, NH 03755, USA
19	⁷ Department of Psychiatry, University of California San Diego, 9500 Gilman Drive, Mail Code
20	0667, BRF2 3A24, La Jolla, CA 92093, USA
21	
22	
23	
24	
25	

27 Abstract

28 Introduction. Muscle fiber cross-sectional area (CSA) and proportion of different fiber types 29 are important determinants of muscle function and overall metabolism. Genetic variation 30 plays a substantial role in phenotypic variation of these traits, however, the underlying genes 31 remain poorly understood. 32 Aims. This study aimed to map quantitative trait loci (QTL) affecting differences in soleus 33 muscle fiber traits between the LG/J and SM/J mouse strains. 34 Methods. Fiber number, CSA, and proportion of oxidative type I fibers were assessed in the 35 soleus of 334 genotyped female and male mice of the F₃₄ generation of advanced intercross 36 lines (AIL) derived from the LG/J and SM/J strains. To increase the QTL detection power, 37 these data were combined with 94 soleus samples from the F_2 intercross of the same 38 strains. Transcriptome of the soleus muscle of LG/J and SM/J females was analysed using 39 microarray. 40 **Results.** Genome-wide association analysis mapped 4 QTL (genome-wide p<0.05) 41 affecting the properties of muscle fibers to Chromosome 2, 3, 4 and 11. A 1.5-LOD QTL 42 support interval ranged between 2.36 Mb and 4.67 Mb. Based on the genomic sequence 43 information, functional and transcriptome data, candidate genes were identified for each of 44 these QTL. 45 **Conclusion.** Combination of analyses in F_2 and F_{34} AlL populations with transcriptome and 46 genomic sequence data in the parental strains is an effective strategy for refining QTL and 47 nomination of the candidate genes. 48

49 Key words: skeletal muscle, muscle fiber types, genetic variation

51 Introduction

52 Skeletal muscle plays a broad range of biological functions including locomotion,

53 thermoregulation, respiration, postural support, protection of bones and viscera; as well as 54 serving as a source of amino acids in times of starvation or disease. Muscle tissue in 55 livestock also provides an essential source of dietary proteins. In humans, there is more 56 than a 2-fold difference in muscle mass between individuals of similar age and same sex (3, 57 33). This is the outcome of variability in the number of muscle fibers and their size (51). 58 These differences are of clinical relevance. Variability in muscle mass significantly impacts 59 energy expenditure (58), influencing preponderance to obesity. In addition, individuals with 60 lower muscle mass may be more vulnerable to impairment of these vital functions due to 61 aging and/or disease related muscle loss. It has recently been reported that there is a 62 positive association between muscle mass and longevity in older adults (66).

63

Human skeletal muscles are mainly comprised of a mixture of type I, IIA and IIX muscle
fibers (62). The number of fibers, their size and varying proportions of the fiber types affect
morphological and functional properties of the muscle (6). A larger diameter of the fibers
and higher number of fibers typically leads to augmented muscular strength and power (25,
28). The proportion of type I muscle fibers is a factor determining success in endurance
sporting events (15, 18) and overall metabolism in humans (24, 29, 44, 74). In livestock,
proportion of oxidative type I fibers is associated with meat quality (65).

71

In humans, genetic factors account for around half of the variation in strength (19, 24, 74)
and the upper limit heritability is even greater (over 0.9) for muscle mass (26). Heritability
estimates of proportion of type I fibers is also high, ranging between 0.4 and 0.9, indicating
that genetic factors play an important role in determining muscle fiber properties (37, 63).
Effects of genetic factors on muscle fibers have also been demonstrated in mouse (20, 22,

59), cattle (68), sheep (10, 38) and pig (71). However, the specific genes underlying these
effects remain largely elusive.

79

80 Attempts at mapping the polygenic architecture of muscle fiber properties in mouse (11), pig 81 (17, 43, 52, 55, 77), cattle (1) and carp (80) have been made. A number of QTL have been 82 identified in these studies. However, the resolution achieved in the F₂ population is not 83 adequate for reliable nomination of the candidate genes in the majority of the QTLs of 84 polygenic traits. The mouse soleus muscle (primarily consists of type I and IIA fiber types), 85 closely resembles the fiber type composition of human skeletal muscles (primarily comprised 86 of type I, IIA and IIX fiber types), and is therefore a particularly interesting experimental 87 model. In our previous study, we mapped soleus muscle fiber traits in an F_2 intercross 88 between the LG/J and SM/J laboratory mouse strains (11). These strains differ in a number 89 of muscular phenotypes, with the LG/J strain displaying a greater proportion of type I fibers, 90 and a greater cross-sectional area (CSA) of type I and IIA muscle fibers. We identified in 91 that study three significant QTLs contributing to the difference in the CSA of muscle fibers 92 between LG/J and SM/J strains (11). Regions of conserved synteny from the identified loci 93 were also implicated in fiber phenotypes in pig supporting the importance of these genomic 94 regions in determining muscle fiber properties. However, the exact genes underlying their 95 effects remain to be determined.

96

Integration of advanced study populations, high throughput gene expression technology and
increasing availability of knockout models aid identification of the causative genes.
Nomination of the genes underlying QTL effects can be facilitated by improving the mapping

100 resolution, and by utilising genomic sequence and transcriptome information. Advanced

101 intercross lines (AIL) have been proposed as a powerful population for mapping QTLs (16).

102 It has been demonstrated recently that a joint F₂ and AIL analysis can combine the

advantages of both mapping populations by increasing the power to detect QTLs and
achieving a higher mapping resolution of various traits in mice (13, 47). Additionally, testing
for differences in specific gene expression has led to several nominations of quantitative trait
genes (30, 35). For validation of such candidate genes, phenotypic effects of relevant
alleles can be examined in experimental populations where these alleles segregate albeit on
a different genetic background. In addition, available knockout models offers particularly
attractive option for validation experiments.

110 In the present study we aimed to fine-map QTL and nominate candidate genes affecting the

111 CSA and proportion of oxidative type I fibers in the soleus muscle in a combined analysis of

112 F_2 and F_{34} AlL mice, and by cross referencing QTL data with soleus transcriptome profiles in

the parental strains. Further filtering of the emerged candidates was carried out in an

114 independent AIL and a knockout model.

116 <u>Methods</u>

117 <u>Muscle Samples</u>

118 This study was carried out on soleus muscles dissected from females and males of the F₃₄ 119 advanced intercross lines (AIL) of the LG/J and SM/J inbred strains. Animals were 120 maintained as previously described (13) and sacrificed at 94 ± 4 days. All procedures were 121 approved by the Institutional Animal Care and Use Committee of the University of Chicago. 122 Soleus muscle samples from F_{34} AIL mice described in our previous study (47) were 123 subjected to histological analyses. The final sample size used in the present study was 334 124 F_{34} mice, 142 females and 192 males, after discarding samples of poor tissue quality. A set 125 of 94 F2 samples (38 females and 56 males) described in our previous study (11) was also 126 used in order to increase the QTL detection power. 127 In addition, we also analysed soleus muscle samples for two hypothesis driven studies 128 aimed at testing the effects of identified candidate genes on percentage of oxidative, type I 129 fibers. First, we examined solei samples from the Chd6 ATPase knockout (n=6), 130 heterozygous (n=4) and wild type (n=4) females. The generation of the Chd6 mutant mice 131 has been previously reported (40). Briefly, the genetic manipulation generated an allele with 132 the ATPase domain of Chd6 (exon 12) flanked by loxP sites so that the action of Cre 133 recombinase would delete this domain. The mice were mated to a germline Cre-expressing 134 strain (Jackson lab strain 003465) to delete both exon 12 and the neomycin resistance 135 marker used for the targeting. Subsequently breeding generated the Chd6 ATPase 136 knockout mice utilized in the present study. Second, solei of the advanced intercross mice 137 (generations F_9 - F_{12}), all homozygous carries of the C57BL/6J (n=22) or DBA/2J (n=23) 138 alleles at the region harbouring the *Alad* gene were selected from the tissue bank of our 139 previous study (9).

140 Phenotype assessment

The soleus muscles were frozen in isopentane cooled in liquid nitrogen. Transverse
sections from the belly of the muscle were cut at a thickness of 10 µm with a cryotome
(Leica CM1850UV) at -20°C. The muscle sections were subjected to ATPase staining (acid
pre-incubation, pH 4.47) to distinguish between fiber types (8). Microscopic images of
stained sections were taken at x5 and x20 magnification.

146 The following phenotypes were assessed: muscle fiber number (type I and IIA) and percent 147 of type I muscle fibers, cross-sectional area (CSA) of type I and type IIA fibers (Figure 1). 148 Muscle fiber traits were manually analysed using ImageJ software (NIH-version 1.43). 25 149 measurements of each fiber type were taken using the freehand selection tool at x20 150 magnification to obtain a value representing the mean CSA of type I or type IIA fibers for that 151 muscle. This was deemed as a representative sample by empirical testing as described 152 previously (11). Total number of type I and type IIA muscle fibers were counted using the 153 ImageJ cell counter plugin on x5 magnification images. As all fibers in mouse soleus pass 154 through the belly of the muscle (69), this method provides an accurate estimate of the 155 number of fibers constituting the muscle. Total number of type I fibers and total number of 156 type IIA fibers were counted, permitting derivation of percentage of type I fibers. Over the 157 course of the study ~200,000 muscle fibers were counted and ~6,700 fibers measured for 158 CSA.

159 <u>Statistical analyses</u>

The GraphPad Prism version 5.0 statistical package was used (GraphPad software, La Jolla,
CA). Data are presented as mean ± SD, unless otherwise stated. The CSA of type I and
type IIA fibers were analysed using a two-way (sex and fiber type) paired-measures (type I
and type IIA fibers) ANOVA.

164 Genotyping and QTL mapping

Mice were genotyped using a custom designed SNP array that included 4,610 polymorphic SNPs that were approximately evenly distributed across the genome, as described 167 previously (13). The genome-wide association analysis was performed in the combined 168 population of the F_{34} and recently published F_2 intercrosses (11) using the R package 169 QTLRel (12). This software accounted for the complex relationships (e.g., sibling, half-170 sibling, cousins) among the F₃₄ mice by using a mixed model, as previously described (12, 171 13). Due to the sex differences in muscle mass in these mice (47), and the discovery of sex 172 specific QTL in other studies (45, 46), we included sex as an additive and interacting 173 covariate. Threshold of significance was estimated by 1000 permutations (14). We defined 174 the support interval for each QTL as the 1.5-LOD drop off on either side of the peak marker. 175 This interval was expressed in physical map position (Mb) by using the nearest genotyped 176 SNP that flanked the support interval, based on the mouse genome build GRCm38.p3.

177 <u>Transcriptome analysis</u>

178 Soleus muscle tissues from 92-day old LG/J and SM/J females (n=3 of each strain) were 179 used. RNA was isolated using TRIzol (Invitrogen Life Technologies, Carlsbad, CA) followed 180 by purification and DNase digestion using RNeasy minikits (Qiagen, Venlo, Netherlands) 181 according to the manufacturer's instructions. Quantification of total RNA was performed on 182 a NanoDrop spectrophotometer (Thermo Scientific) and quality tested on an Agilent 183 Tapestation with R6K Screentapes (RIN ≥7.3). Generation of sense strand cDNA from purified total RNA (Ambion[®] WT expression kit, Ambion, Austin, Texas) followed by 184 185 fragmentation and labelling (GeneChip WT labelling kit, Affymetrix, Santa Clara, CA) were 186 performed according to the manufacturer's instructions. Hybridisation, washing, staining and 187 scanning of microarrays were carried out on Affymetrix Mouse Gene 2.0 ST microarrays 188 according to the manufacturer's standard protocols using a GeneChip Fluidics station 450 189 and GCS3000 scanner (Affymetrix®, Santa Clara, CA). Microarray data are available in the 190 ArrayExpress database (www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-191 5290.

193 Data pre-processing and quality control analysis was performed using Affymetrix®

194 Genechip® Expression Console v1.2. Probe cell intensity data on the Mouse Gene 2.0 ST

array (CEL files) were processed using the RMA16 algorithm (Affymetrix, Santa Clara, CA,

196 USA) which fits a robust linear model at probe level by employing background correction,

197 quantile normalisation of log2 transformed data and summarisation to probe level data (CHP

198 files, 41,345 probe sets).

199

200 Data was analysed for differentially expressed genes in Partek® Genomics Suite® version 201 6.6, build 6.15.0730 (Partek Inc., St Louis, MO) using a Mus musculus build mm10 202 annotation file for Mouse Gene 2.0 ST microarrays (MoGene-2 0-st-v1.na35.mm10). CEL 203 files (Expression Console v 1.2, Affymetrix, Santa Clara, CA) were imported to Partek 204 Genomics Suite v 6.6 and processed using RMA normalisation with background correction 205 of log2 transformed data and probe set summarisation by median polish. Differential 206 expression analysis between the LG/J and SM/J strains of all genes (n=41,345 transcript 207 clusters) was determined by 1-way ANOVA with Storey's FDR, and q-value ≤0.05 208 considered significant (n=819 genes differentially expressed \geq 1.2 fold; see **Supplementary** 209 Table 1). 210 To assess transcription of positional candidate genes in each strain, a hypothesis driven 211 analysis of differential gene expression was performed between the LG/J and SM/J strains 212 on all genes mapping to the support interval defined for each QTL in the GWAS described 213 above. Using Partek Genomics Suite v.6.6, a total of 159 genes that were represented on 214 the mouse Gene 2.0ST microarray, were identified in Mus musculus genome build 215 GRCm38, mm10 within mapping co-ordinates Chr2:158908559–162608559 (26 genes),

216 Chr3:33308451–35708451 (15 genes), Chr4:57605946–62913639 (77 genes) or Chr

11:27900000–31500000 (41 genes). 1-way ANOVA identified differentially expressed genes

between the LG/J and SM/J strains (P<0.05). Fold change was calculated using the

219 geometric mean of samples in each group.

220 Candidate genes

221 Nomination of the candidate genes was based on the following three criteria. First, we 222 scrutinized polymorphisms in positional candidates between the LG/J and SM/J strains. The 223 emphasis was on the indels and SNPs that would affect the coding sequence and lead to 224 changes in amino acids. To assess whether amino acid substitution would influence the 225 function of a protein, evolutionary conservation at the site of substitution and properties of 226 substituted amino acids were considered using three different bioinformatics tools as 227 described by Nikolskiy and colleagues (56). Second, we examined expression of positional 228 candidates across a panel of over ninety mouse tissues and cell types available in BioGPS 229 GeneAtlas MOE430, gcrma dataset (79). This analysis permits a quantitative comparison of 230 transcript abundance of a gene between tissues. We considered that an abundant 231 expression in skeletal muscle lineage, i.e. muscle tissue and/or C2C12 myogenic cell line, 232 implies functional and/or structural relevance of a gene in this tissue. Third, we compared 233 gene expression levels in the soleus muscle between the two strains as described in the 234 previous section. Expression difference in this analysis might point at the strain-specific, 235 genotype-dependent mechanism underlying the phenotypic difference.

236

237 **Results**

238 Phenotypic analyses

CSA. Cross section analysis of soleus muscle fibers were done on mice of both sexes from the F_{34} cohort. For muscle fiber cross-sectional area, we observed a statistically significant sex by fiber type interaction (P<0.0001). In the female F_{34} mice there was no significant difference between type I and type IIA muscle fiber areas (913 ± 229 µm², n=140; and 952 ± 242 µm², n=140 respectively; P=0.2). However, there was a significant difference within the males, with the type I muscle fiber area being smaller than IIA fiber area (1084 ± 238 µm², n=187; and 1215 ± 294 μ m², n=187 respectively; P< 0.0001). Muscle fiber area was lower in females than males for type I CSA, (P < 0.0001) and type IIA CSA (P < 0.0001).

Percentage of type I fibers. The number of type I fibers as a percentage of total fibers varied substantially between individuals, ranging from 30% to 67% in females, and from 26% to 59% in males (**Figure 1**) and was greater in females than males ($46 \pm 8\%$, n=142; and 39 $\pm 6\%$, n=189; respectively; P < 0.0001).

Total fiber number. No difference was observed in the total soleus fiber number between females and males (646 ± 102 , n=120, and 667 ± 105 , n=177, respectively; P= 0.0979).

253

254 QTL analyses

Muscle fiber traits approximated the normal distribution in both the F_2 and F_{34} population

256 (**Supplementary Figure 1**). We identified significant QTL (at the 1% or 5% level of genome-

wide statistical significance) (39) for CSA of type I and type IIA fibers and the percentage of

type I fibers. We also identified chromosome-wide significant QTL for CSA of type I and type

259 IIA fibers, the percentage of type I fibers and total fiber number (**Table 1**). The size of the

support interval of these QTL ranged from 0.4-40.7 Mb, with a median of 4.6 Mb.

261 The QTL at the genome-wide level of significance for CSA of type I and type IIA fibers on

chromosome 3 was named *Mfq5*. The QTL at the genome-wide level of significance for the

263 percentage of type I fibers on chromosome 2 and 4 were named *Mfq4* and *Mfq6*,

respectively. The SM/J allele conferred a greater percentage of type I fibers at *Mfq4*, and a

greater CSA at *Mfq5*. The LG/J allele conferred a greater percentage of type I fibers at *Mfq6*

266 locus.

267 A significant QTL affecting CSA of type I and type IIA fibers was also detected on

chromosome 11 (Figure 2) within the same region as locus *Mfq3*, previously identified in the

 F_2 intercross of the same parental strains (11). The QTL exhibited male-specificity in both

type I and IIA fibers of the F_{34} mice (**Figure 3**). Because this QTL recapitulated properties of the *Mfq3* locus, which we also found to be male specific in the F_2 population, we concluded that the same locus has been refined in F_{34} and did not assign a new name for this QTL. Earlier reported *Mfq2* locus has been refined in a similar manner; a QTL on chromosome 6 affecting CSA of type I and type IIA fibers (at 1% chromosome specific threshold) was engulfed by the support interval of *Mfq2* and also replicated its increasing allele, LG/J, in both females and males (not shown).

277 <u>Gene expression analyses</u>

278 We hypothesized that each identified QTL harbours one or more genetic variants that drive 279 phenotypic differences by means of differential gene expression. Hypothesis driven analysis 280 of differential expression in soleus muscle was performed between LG/J and SM/J strains for 281 the genes in the most robust QTLs affecting fiber CSA or % Type I fibers (*Mfq3*, *Mfq4*, *Mfq5* 282 and Mfq6). The Mouse Gene 2.0 ST expression array contains 159 genes residing within 283 the support intervals of these QTLs (Supplementary Table 2). Twenty genes (Table 2) 284 showed evidence of differential expression (ANOVA, p≤0.05), 2 of which, Alad and Hdhd3, 285 were significant after correction for the multiple testing problem (Storey's FDR q \leq 0.05). 286 Compared to other tissues and cell types, expression of differentially expressed genes Mafb, 287 Acyp2 and Mtif2 (Table 2), is particularly enriched in skeletal muscle (BioGPS, Mouse 288 MOE430 gene expression data).

289 <u>Genomic analyses</u>

Positional candidates with non-synonymous polymorphisms provide a plausible genetic
cause for the phenotypic differences. Based on the genomic sequence of the LG/J and
SM/J strains (56), we identified 21 genes in the QTL regions with non-synonymous
polymorphisms predicted to affect protein function by at least one out of three algorithms
used in the analysis (Supplementary Table 3). Four of those genes (*Mfq3: Mtif2, Rtn4, Psme4; Mfq5: Dnajc19*) are prioritized further because of their preferential expression in

- 296 muscle lineage (differentiated muscle and/or C2C12 myoblasts) compared to other tissues
- and cell types. Among those, the *Mtif2* gene differs by 3 (rs26871496, rs26871494,
- rs29436813) and *Rtn4* by 9 (rs29473364, rs29469198, rs13463765, rs29465940,
- rs26857726, rs26857725, rs29474377, rs26857722, rs26857721) amino acids between the
- 300 two strains. At all SNPs the SM/J strains carries reference while the LG/J strain the
- 301 alternative allele.

302 Candidate gene analyses

303 The *Chd6* gene emerged as a differentially expressed positional candidate for the *Mfq4*

locus affecting percentage of type I fibers (**Table 2**). To test its effect we examined soleus

305 muscles of *Chd6* knockout, heterozygous and wild type littermates. This analysis however

revealed that the genotype of the animals did not have a significant effect (P=0.30) on the

307 percentage of type I fibers (**Figure 4**).

308 The Alad gene emerged as a candidate for another locus affecting proportion of type I fibers,

309 *Mfq6*. In the animals of an advanced intercross between the C57BL/6J and DBA/2J strains

- 310 (these strains carry one or three copies of *Alad*, respectively (3)), we examined if percentage
- 311 of type I fibers was genotype-dependent. The analysis revealed no difference in the

percentage of type I fibres between the carriers of the C57BL/6J and the DBA/2J alleles, 42

 \pm 7% and 42 \pm 8%, respectively.

314 Discussion

315 A previous study on muscle weight in LG/J and SM/J strains identified a two-fold difference 316 in soleus muscle size (47). We then explored the cellular and genetic mechanisms 317 contributing to this phenomenon, finding that the difference was largely due to the CSA of 318 muscle fibers and we mapped QTL affecting muscle fiber traits in an F₂ intercross between 319 the LG/J and SM/J strains (11). The present study, which utilizes the F₃₄ advanced 320 intercross, verified, refined and expanded our earlier findings. 321 A number of studies have previously reported the effects of Stat5a and Stat5b (36), Pgc-1a 322 (42), Ky (4), myostatin (54), leptin (61), calcineurin (76), Sod1 (5), alpha-actinin-3 (50), 323 dystrophin (7), Tbx15 (41) and IIB myosin heavy chains (2) genes on muscle fiber area in 324 knockout or mutant models. In addition, $Pgc-1\alpha$ (75), calcineurin (76), Foxo1 (34) and 325 myostatin (20) are reported to affect the proportion of muscle fiber types. However, the 326 genomic positions of these genes have not been linked to muscle fiber differences between 327 the LG/J and SM/J strains, implicating involvement of novel genes.

Muscle fiber number. The number of fibers is an important determinant of muscle size and functional properties. It is set during embryogenesis and the first post-natal week in mice (78). The number of muscle fibers in males (667 ± 105) and females (646 ± 102) of the F₃₄ population was comparable to that observed in the soleus of the F₂ population (645 ± 102 and 595 ± 107 , respectively), and within the range of the fiber count observed in solei of a variety of different strains of mice ~250-~900 fibers (32, 49, 57, 70, 72).

From these data it emerged that males and females are born with a similar number of fibers in soleus muscle, and that the sex difference in muscle weight (males have approximately 30% larger soleus than females) is due to the difference in fiber size. Comparison of the parental strains also revealed a similar number of fibers (11), despite the 2-fold difference in soleus weight (47), demonstrating that size rather than number of fibers determines variation in muscle weight between the LG/J and SM/J strains. Fiber area. The CSA of muscle fibers in the LG/J strain is 49% to 90% greater than the corresponding fibers in the SM/J strain, indicating that this variable accounts for a large portion of the muscle mass difference between the strains (47).

The area of type I (1084 ± 238 μ m² and 913 ± 229 μ m² for males and females, respectively) and type IIA (1215 ± 294 μ m² and 952 ± 242 μ m², respectively) of the F₃₄ mice was comparable to the corresponding fiber area of the F₂ mice of the same lineage (11) and it is within the range reported for the type I, between 920 μ m² and 1780 μ m² (32, 57, 70), and type IIA fiber area, between 700 μ m² and 1400 μ m² (32, 70), in various inbred mouse strains.

Percentage of type I fibers. The percentage of type I fibers in male $(39 \pm 6\%)$ and female (47 ± 8%) F₃₄ mice were also within the range of previous studies, which showed the percentage of type I fibers in the soleus muscle fluctuates between ~25 and ~66% (32, 57, 70).

In the F_{34} mice we replicated our observation in the F_2 population that the percentage of type I fibers was significantly greater in females than males. This sex difference is also observed in various human muscles where, in general, women have a higher percentage of type I muscle fibers than males (27, 53, 60, 64, 67). The phenomenon is likely to be explained, at least partly, by the effect of androgens; castration leads to a higher percentage of type I fibers in the soleus of male mice (73).

Validation and refinement of genetic architecture. In the present study, we validated and refined the genetic architecture of muscle fibers identified in an F_2 intercross between the same parental strains (11). In order to increase QTL detection power, we increased sample size by combining the F_{34} and F_2 data. The median mapping resolution of 4.6 Mb for muscle fiber QTLs was comparable with 3.7 Mb of muscle weight QTLs obtained in the same population albeit using ~1,600 fewer genetic markers than in the present analysis (47). A genome-wide significant QTL identified in the present study between 27.9 Mb and 31.4 Mb on chromosome 11 (**Table 1**) overlapped with a significant QTL, *Mfq3*, mapped in the F_2 population (11). In addition to the chromosomal location, the increasing allele of this locus (LG/J) and its male-specific effect (**Figure 3**) were also replicated in F_{34} , suggesting that the same gene(s) were involved in two different populations and permitting us to refine the *Mfq3* locus from 51.6 Mb to 3.57 Mb. The presence of two satellite QTL proximal of the refined *Mfq3* (**Table 1**) suggests that the QTL observed in the F_2 population (11) might have been an outcome of up to three linked loci.

The recently reported "mini-muscle" locus, mapped to 67.1–70.2 Mb on chromosome 11, affects muscle fiber area and proportion of fiber types (21-23). However, the mutation responsible for the "mini-muscle" phenotype maps to an intron of *Myh4* gene located at 67.2 Mb (31), between the support intervals of two adjacent QTLs affecting fiber type between the LG/J and SM/J strains (**Table 1**). Together, these data suggest that a number of genes residing on chromosome 11 might be involved in the regulation of muscle fiber phenotypes.

379 The QTL affecting the CSA of type I and type IIA fibers on chromosome 6, albeit at 1% 380 chromosome-wide threshold of significance (**Table 1**), overlapped with the *Mfq2* locus found 381 in the F_2 population, characterized by the same increasing allele, LG/J. Thus, the support 382 interval of Mfq2 could be considered to be 5.18 Mb rather than the previously reported 56.5 383 Mb. Importantly, the immediate proximity of the refined region (Chr 6: 110.8-116.0 Mb) to 384 the syntenic region (Chr 6:116.0-118.0 Mb) implicated in the QTL affecting the diameter of 385 pig IIA fibers (17) suggest that the same genes could be underlying the effects of these 386 QTLs in mice.

A QTL affecting percentage of type I fibers (at 10% chromosome-wide threshold) on chromosome 1 (67.6to 70.8 Mb) overlapped with *Mfq1* locus which influenced the CSA of type I and type IIA fiber area in the F₂ population (11). However, because the CSA and percentage of type I fibers are poorly correlated traits both in the F₃₄ (**Supplementary Table** **4**) and the F_2 mice (11), it is likely that different genes are underlying the *Mfq1* locus and the QTL identified in the F_{34} population. Further studies are required to clarify this observation.

393 Transcriptome analysis

394 In the present study, the expressed transcriptome in soleus muscle of the parental strains 395 was examined in order to facilitate nomination of the candidate genes within the refined QTL. 396 We hypothesized that if the phenotypic effect of the QTL was brought about by the allele 397 specific abundance of transcripts encoded by genes within the QTL, such genes would be 398 differentially expressed in the transcriptome between the parental strains. Comparison of 399 expression of the genes within the four most robust QTLs identified Alad and Hdhd3 genes 400 as potential candidates for the *Mfq6* locus, which affects the proportion of type I fibers. 401 Transcripts of both genes are more abundant in the LG/J compared to the SM/J strain. This 402 is consistent with our findings in the TA muscle of the same strains (48). Of these two 403 identified candidate genes, transcripts of Alad are ~20 times more abundant in the mouse 404 muscle than Hdhd3, regardless of strain. In addition, Alad may play a role during 405 myogenesis as its expression in C2C12 myogenic cells is 5-fold higher compared to 406 differentiated muscle (79).

407 Candidate genes.

408 The support intervals of four most robust QTLs harbor 159 genes (Supplementary Table 2). 409 These regions were scrutinized further for the genes fulfilling one of the following criteria: 410 presence of the functional variants (i.e. non-synonymous SNPs predicted to alter function of 411 encoded protein); abundance of transcript in muscle lineage, particularly in comparison to 412 other tissues and cell types; differential expression in the soleus of the two strains; and by 413 comparing genomic sequence between the LG/J and SM/J strains a list of 21 genes was 414 highlighted (Supplementary Table 3) with the strain-specific functional variants. Using 415 bioinformatics, 4 genes abundantly and/or preferentially expressed in skeletal muscle 416 compared to other types of tissues and cells were identified. Our own analysis of gene

417 expression in soleus muscle highlighted a set of 20 genes differentially expressed between 418 the two strains (Table 2). Intersection of all these lists permitted us to prioritise nine 419 candidate genes which appeared on more than one of these lists and/or for which 420 independent and accessible validation models were available (i.e. Chd6 and Alad). Because 421 neither the Chd6 (Figure 4) nor Alad genes were found to affect proportion of type I fibres in 422 the way predicted by the QTL analyses, the list of prioritised candidates was reduced to 7 423 genes annotated in **Supplementary Table 5**. Three out of four QTLs contain one (*Mfq6*) or 424 more candidate genes. All candidates are abundantly transcribed in muscle lineage with 425 Psme4, Acyp2 and Mafb showing the highest level of expression in skeletal muscle 426 compared to other tissues and cells. None of the seven candidates have been previously 427 implicated to affect properties of skeletal muscle fibres although some of them have been 428 implicated in cardiomyopathy or function as transcription factors (Supplementary Table 5). 429 Thus, genomic and gene expression analyses permitted focusing on a limited number of 430 positional candidates in the future validation studies for establishing the causative genes.

431 Conclusion

In conclusion, we have refined the genetic architecture affecting cross sectional area of
soleus muscle fibers and proportion of type I fibers in the LG/J and SM/J derived lineage.
Integrating QTL mapping, genomic and transcriptome data from homologous muscle
highlighted several candidate genes that may underpin muscle phenotypes critical to health
and disease and worthy of follow up analyses.

437

438

439

440

441 Grants

442 The study was supported by NIH R01AR056280 grant (AL), NIH R01DA024845 (AAP) and

443 Marie Curie International Reintegration Grant 249156 (AL).

444

445 Acknowledgements

The authors wish to acknowledge technical assistance from Diane Stewart, Jasmine C. Berry and Jake T. Brown. We acknowledge the Dartmouth Mouse Modelling Shared Resource of the Norris Cotton Cancer Center (support grant P30CA023108-27), for providing *Chd6* knockout mouse samples.

450

451 Author contributions

A.L. conceived and supervised the study, A.M.C. phenotyped muscle samples, A.A.P.
provided genotypes and oversaw the QTL analyses, R.C. designed the QTLRel software
used in the QTL analysis, A.M.C. carried out QTL mapping, M.E.S, C.M. and E.C.D. did
transcriptome analysis, S.N.F. and J.L.F. generated and provided *Chd6* knockout samples,
A.L. and A.M.C. wrote the manuscript with input from all co-authors.

457

459 Figure legends

Figure 1. Individual variability in proportion of oxidative fibers. Representative images
 of F₃₄ female soleus cross-sections following myosin ATPase staining (acid pre-incubation).
 Dark fibers type I, pale fibers type IIA.

463

Figure 2. Type I fiber cross-sectional area QTL on chromosome 11. Analyses were carried out in the F_2 intercross and in the combined F_2 and F_{34} populations. *X*-axis indicates the relative position in the linkage map in centimorgan (cM). The thresholds are at the level of 0.05 genome wise significance for the F_2 output (dotted line) and combined output (solid line).

469

470 Figure 3. Sex specificity of *Mfq3* locus on cross-sectional area (CSA) of soleus type I

- 471 and IIA fibers in the F_{34} intercross. Mean and SEM. Genotype at the peak marker: LG,
- homozygous for LG/J allele; H, heterozygous; SM, homozygous for SM/J allele.
- 473

474

475 Figure4. Percentage of type I fibers in the soleus muscle of 4 month old *Chd*6

476 knockout (KO), heterozygous (HET) and wild-type (WT) females. There is no difference

in percentage of type I muscle fibers in the soleus muscle between knockout, heterozygotes

and wild-type groups (P=0.3041). Each data point is from a single mouse, horizontal lines

479 represent group mean.

Chr	Thr**	Level***	Start Mb [†]	End Mb	Size Mb	Trait	Locus¥
1	С	0.1	67.6	70.7	3.1	% Type I	
1	С	0.1	193.9	194.3	0.4	% Type I & CSA2A	
2	С	0.1	92.4	104.8	12.4	% Type I	
2	С	0.05	139.6	145.6	6.0	% Type I	
2	G	0.01	158.8	162.5	3.7	% Type I	Mfq4 (SM)
3	G	0.05	33.6	40.0	6.4	CSA1 & CSA2A	Mfq5 (SM)
4	G	0.05	57.7	62.7	5.0	% Type I	Mfq6 (LG)
4	С	0.05	103.9	106.1	2.2	% Type I	
6	С	0.05	81.9	84.1	2.2	CSAIIA	
6	С	0.01	110.8	116.0	5.2	CSA1 & CSA2A	Mfq2* (LG)
7	С	0.05	138.4	140.0	1.6	% Type I	,
8	С	0.1	7.4	12.4	5.0	% Type I	
8	С	0.05	89.0	92.4	3.4	TOTAL	
8	С	0.01	121.9	128.6	6.7	TOTAL	
10	G	0.1	120.7	121.3	0.6	% Type I	
11	С	0.1	12.4	17.2	4.8	CSAIIA	
11	С	0.1	19.1	23.1	4.0	CSAIIA	
11	G	0.01	28.0	31.5	3.5	CSA1 & CSA2A	Mfq3* (LG)
11	С	0.1	62.5	64.2	1.7	% Type I	
11	С	0.1	70.6	76.2	5.6	% Type I	
12	С	0.1	27.6	29.3	1.7	CSA1	
13	С	0.01	5.3	9.9	4.6	% Type I	
13	С	0.05	71.5	74.0	2.5	CSAIIA	
14	С	0.05	93.6	102.3	8.7	CSAIIA	
15	С	0.1	12.1	20.3	8.2	TOTAL	
16	С	0.05	68.9	75.1	6.2	CSA1 & CSA2A	
Х	С	0.01	11.8	52.5	40.7	TOTAL	

481 **Table 1.** Characteristics of muscle fiber QTL in combined analyses of the F_2 and F_{34}

482 intercrosses derived from the LG/J and SM/J strains.

483 * refined previously identified QTL in the LG/J and SM/J F₂ intercross (47).

484 ** C – chromosome-wide threshold, G- genome-wide threshold

- 485 *** Level of significance
- 486 ¥ LG –increasing allele is LG/J, SM- increasing allele is SM/J
- 487 [†] Genomic positions based on GRCm38.p3.

Chr	QTL	Probe set ID	Gene	p-value*	Fold-Change**	Gene name***
2	Mfq4	17393868	Mafb	0.033	-1.77	v-maf musculoaponeurotic fibrosarcoma oncogene family, protein B (avian)
		17393910	Chd6	0.042724	-1.13	chromodomain helicase DNA binding protein 6
		17404652	Gm24780	0.032914	-1.98	Predicted gene Gm24780, predicted protein is B4HDV3.
3	Mfq5	17396801	Ttc14	0.033387	-1.14	tetratricopeptide repeat domain 14
		17396876		0.0219376	-1.80	There are no assigned mRNA sequences for this probe set. The probe set lies within IncRNA Sox2ot (Sox2 overlapping transcript, non-protein coding)
		17425606	Gm12526	0.046791	-1.17	predicted gene 12526
		17414380	Gm24277	0.00525644	-2.07	Gm24277 a known snRNA. The probeset also lies within an intronic region of RefSeq gene Pakap (PALM2-AKAP2), read through transcript on chromosome 4
		17425701	Mir3095	0.0298235	-1.78	Mir3095 (Entrez ID 100526502; EST ENSMUST00000175552).
		17426097	Мир3	0.00304	-1.49	major urinary protein 3
		17426126	Fkbp15	0.018038	-1.10	FK506 binding protein 15
4	Mfq6	17414545	Slc31a1	0.049799	1.12	solute carrier family 31, member 1
		17426166	Cdc26	0.021407	-1.29	cell division cycle 26
		17426198	Hdhd3	0.000869	1.86	haloacid dehalogenase-like hydrolase domain containing 3
		17426206	Alad	9.75E-05	1.91	aminolevulinate, delta-, dehydratase
		17414600	Rgs3	0.023096	1.08	regulator of G-protein signaling 3
		17248064	Mtif2	0.015517	-1.16	mitochondrial translational initiation factor 2
		17261285	LOC102637613	0.00432682	1.68	linc RNA [AK084560 (EST)/ Gm12092 (predicted gene)].
		17248127		0.00187422	-1.46	There are no assigned mRNA sequences for this transcript The probe set lies within an intron of <i>Sptbn1</i> .
11	Mfq3	17261393	Асур2	0.011777	1.26	acylphosphatase 2, muscle type
		17248196	Asb3	0.014221	-1.15	ankyrin repeat and SOCS box-containing 3

Table 2. Positional candidate genes differentially expressed between LG/J and SM/J soleus muscles.

491 * ANOVA p-value for strain effect; ** Fold change uses SM/J as baseline (negative values indicate LG/J expression is down compared to SM/J,

492 positive values LG/J expression up compared to SM/J); bold indicates that gene is predominantly and/ or strongly expressed in skeletal muscle

tissue (79). *** For probe sets not designed against an annotated gene, genes at the genomic loci of the Affymetrix probeset were identified in

494 UCSC genome browser using mouse genome build GRCm38.

497

498 1. Abe T, Saburi J, Hasebe H, Nakagawa T, Kawamura T, Saito K, Nade T, Misumi S, Okumura 499 T, Kuchida K, Hayashi T, Nakane S, Mitsuhasi T, Nirasawa K, Sugimoto Y, and Kobayashi E. Bovine 500 quantitative trait loci analysis for growth, carcass, and meat quality traits in an F2 population from a 501 cross between Japanese Black and Limousin. Journal of animal science 86: 2821-2832, 2008. 502 Allen DL, Sartorius CA, Sycuro LK, and Leinwand LA. Different pathways regulate expression 2. 503 of the skeletal myosin heavy chain genes. The Journal of biological chemistry 276: 43524-43533, 504 2001. 505 3. Bishop TR, Miller MW, Wang A, and Dierks PM. Multiple copies of the ALA-D gene are 506 located at the Lv locus in Mus domesticus mice. Genomics 48: 221-231, 1998. Blanco G, Coulton GR, Biggin A, Grainge C, Moss J, Barrett M, Berquin A, Marechal G, 507 4. 508 Skynner M, van Mier P, Nikitopoulou A, Kraus M, Ponting CP, Mason RM, and Brown SD. The 509 kyphoscoliosis (ky) mouse is deficient in hypertrophic responses and is caused by a mutation in a 510 novel muscle-specific protein. *Human molecular genetics* 10: 9-16, 2001. 511 5. Bordet T, Schmalbruch H, Pettmann B, Hagege A, Castelnau-Ptakhine L, Kahn A, and Haase 512 **G**. Adenoviral cardiotrophin-1 gene transfer protects pmn mice from progressive motor 513 neuronopathy. The Journal of clinical investigation 104: 1077-1085, 1999. 514 6. Bottinelli R, and Reggiani C. Human skeletal muscle fibres: molecular and functional 515 diversity. Progress in biophysics and molecular biology 73: 195-262, 2000. 516 7. Briguet A, Courdier-Fruh I, Foster M, Meier T, and Magyar JP. Histological parameters for 517 the quantitative assessment of muscular dystrophy in the mdx-mouse. Neuromuscular disorders : 518 NMD 14: 675-682, 2004. 519 8. Brooke MH, and Kaiser KK. Muscle fiber types: how many and what kind? Archives of 520 neurology 23: 369-379, 1970. 521 Carbonetto P, Cheng R, Gyekis JP, Parker CC, Blizard DA, Palmer AA, and Lionikas A. 9. 522 Discovery and refinement of muscle weight QTLs in B6 × D2 advanced intercross mice. Physiological 523 Genomics 46: 571-582, 2014. 524 Carpenter CE, Rice OD, Cockett NE, and Snowder GD. Histology and composition of muscles 10. 525 from normal and callipyge lambs. Journal of animal science 74: 388-393, 1996. 526 Carroll AM, Palmer AA, and Lionikas A. QTL Analysis of Type I and Type IIA Fibers in Soleus 11. 527 Muscle in a Cross between LG/J and SM/J Mouse Strains. Frontiers in genetics 2: 99, 2011. 528 12. Cheng R, Abney M, Palmer AA, and Skol AD. QTLRel: an R package for genome-wide 529 association studies in which relatedness is a concern. BMC genetics 12: 66, 2011. Cheng R, Lim JE, Samocha KE, Sokoloff G, Abney M, Skol AD, and Palmer AA. Genome-wide 530 13. 531 association studies and the problem of relatedness among advanced intercross lines and other 532 highly recombinant populations. Genetics 185: 1033-1044, 2010. 533 14. Cheng R, and Palmer AA. A Simulation Study of Permutation, Bootstrap, and Gene Dropping 534 for Assessing Statistical Significance in the Case of Unequal Relatedness. *Genetics* 193: 1015-1018, 535 2013. 536 15. Costill DL, Daniels J, Evans W, Fink W, Krahenbuhl G, and Saltin B. Skeletal muscle enzymes 537 and fiber composition in male and female track athletes. Journal of applied physiology 40: 149-154, 538 1976. 539 Darvasi A, and Soller M. Advanced intercross lines, an experimental population for fine 16. 540 genetic mapping. Genetics 141: 1199-1207, 1995. 541 17. Estelle J, Gil F, Vazquez JM, Latorre R, Ramirez G, Barragan MC, Folch JM, Noguera JL, Toro 542 MA, and Perez-Enciso M. A quantitative trait locus genome scan for porcine muscle fiber traits 543 reveals overdominance and epistasis. Journal of animal science 86: 3290-3299, 2008.

544 18. Fink WJ, Costill DL, and Pollock ML. Submaximal and maximal working capacity of elite 545 distance runners. Part II. Muscle fiber composition and enzyme activities. Annals of the New York 546 Academy of Sciences 301: 323-327, 1977. 547 19. Frederiksen H, Gaist D, Petersen HC, Hjelmborg J, McGue M, Vaupel JW, and Christensen K. 548 Hand grip strength: a phenotype suitable for identifying genetic variants affecting mid- and late-life 549 physical functioning. Genetic epidemiology 23: 110-122, 2002. 550 20. Girgenrath S, Song K, and Whittemore LA. Loss of myostatin expression alters fiber-type 551 distribution and expression of myosin heavy chain isoforms in slow- and fast-type skeletal muscle. 552 Muscle & nerve 31: 34-40, 2005. 553 Guderley H, Houle-Leroy P, Diffee GM, Camp DM, and Garland T, Jr. Morphometry, 21. 554 ultrastructure, myosin isoforms, and metabolic capacities of the "mini muscles" favoured by 555 selection for high activity in house mice. Comparative biochemistry and physiology Part B, 556 Biochemistry & molecular biology 144: 271-282, 2006. 557 Guderley H, Joanisse DR, Mokas S, Bilodeau GM, and Garland T, Jr. Altered fibre types in 22. 558 gastrocnemius muscle of high wheel-running selected mice with mini-muscle phenotypes. 559 Comparative biochemistry and physiology Part B, Biochemistry & molecular biology 149: 490-500, 560 2008. 561 23. Hartmann J, Garland T, Jr., Hannon RM, Kelly SA, Munoz G, and Pomp D. Fine mapping of 562 "mini-muscle," a recessive mutation causing reduced hindlimb muscle mass in mice. The Journal of 563 heredity 99: 679-687, 2008. 564 24. Hernelahti M, Tikkanen HO, Karjalainen J, and Kujala UM. Muscle fiber-type distribution as 565 a predictor of blood pressure: a 19-year follow-up study. Hypertension 45: 1019-1023, 2005. 566 25. Hubal MJ, Gordish-Dressman H, Thompson PD, Price TB, Hoffman EP, Angelopoulos TJ, 567 Gordon PM, Moyna NM, Pescatello LS, Visich PS, Zoeller RF, Seip RL, and Clarkson PM. Variability in 568 muscle size and strength gain after unilateral resistance training. Medicine and science in sports and 569 exercise 37: 964-972, 2005. Huygens W, Thomis MA, Peeters MW, Vlietinck RF, and Beunen GP. Determinants and 570 26. 571 upper-limit heritabilities of skeletal muscle mass and strength. Canadian journal of applied 572 *physiology = Revue canadienne de physiologie appliquee* 29: 186-200, 2004. 573 Jaworowski A, Porter MM, Holmback AM, Downham D, and Lexell J. Enzyme activities in 27. 574 the tibialis anterior muscle of young moderately active men and women: relationship with body 575 composition, muscle cross-sectional area and fibre type composition. Acta physiologica Scandinavica

576 176: 215-225, 2002.

Kadi F, Eriksson A, Holmner S, Butler-Browne GS, and Thornell LE. Cellular adaptation of the
 trapezius muscle in strength-trained athletes. *Histochemistry and cell biology* 111: 189-195, 1999.

579 29. Karjalainen J, Tikkanen H, Hernelahti M, and Kujala UM. Muscle fiber-type distribution
580 predicts weight gain and unfavorable left ventricular geometry: a 19 year follow-up study. *BMC*581 *cardiovascular disorders* 6: 2, 2006.

S82 30. Karp CL, Grupe A, Schadt E, Ewart SL, Keane-Moore M, Cuomo PJ, Kohl J, Wahl L,
 S83 Kuperman D, Germer S, Aud D, Peltz G, and Wills-Karp M. Identification of complement factor 5 as
 S84 a susceptibility locus for experimental allergic asthma. *Nature immunology* 1: 221-226, 2000.

585 31. Kelly SA, Bell TA, Selitsky SR, Buus RJ, Hua K, Weinstock GM, Garland T, Pardo-Manuel de
586 Villena F, and Pomp D. A Novel Intronic Single Nucleotide Polymorphism in the Myosin heavy
587 polypeptide 4 Gene Is Responsible for the Mini-Muscle Phenotype Characterized by Major Reduction
588 in Hind-Limb Muscle Mass in Mice. *Genetics* 195: 1385-1395, 2013.

589 32. Kilikevicius A, Venckunas T, Zelniene R, Carroll AM, Lionikaite S, Ratkevicius A, and
 590 Lionikas A. Divergent physiological characteristics and responses to endurance training among
 591 inbred mouse strains. Scandinavian journal of medicine & science in sports 23: 657-668, 2013.

592 33. Kim J, Wang Z, Heymsfield SB, Baumgartner RN, and Gallagher D. Total-body skeletal
 593 muscle mass: estimation by a new dual-energy X-ray absorptiometry method. *The American journal* 594 of clinical nutrition 76: 378-383, 2002.

595 34. Kitamura T, Kitamura YI, Funahashi Y, Shawber CJ, Castrillon DH, Kollipara R, DePinho RA, 596 Kitajewski J, and Accili D. A Foxo/Notch pathway controls myogenic differentiation and fiber type 597 specification. The Journal of clinical investigation 117: 2477-2485, 2007. 598 35. Klein RF, Allard J, Avnur Z, Nikolcheva T, Rotstein D, Carlos AS, Shea M, Waters RV, 599 Belknap JK, Peltz G, and Orwoll ES. Regulation of bone mass in mice by the lipoxygenase gene 600 Alox15. Science (New York, NY) 303: 229-232, 2004. 601 36. Klover P, Chen W, Zhu BM, and Hennighausen L. Skeletal muscle growth and fiber 602 composition in mice are regulated through the transcription factors STAT5a/b: linking growth 603 hormone to the androgen receptor. FASEB journal : official publication of the Federation of American 604 Societies for Experimental Biology 23: 3140-3148, 2009. 605 Komi PV, Viitasalo JH, Havu M, Thorstensson A, Sjodin B, and Karlsson J. Skeletal muscle 37. 606 fibres and muscle enzyme activities in monozygous and dizygous twins of both sexes. Acta 607 physiologica Scandinavica 100: 385-392, 1977. 608 Koohmaraie M, Shackelford SD, Wheeler TL, Lonergan SM, and Doumit ME. A muscle 38. 609 hypertrophy condition in lamb (callipyge): characterization of effects on muscle growth and meat 610 quality traits. Journal of animal science 73: 3596-3607, 1995. 611 39. Lander E, and Kruglyak L. Genetic dissection of complex traits: guidelines for interpreting 612 and reporting linkage results. Nature genetics 11: 241-247, 1995. 613 40. Lathrop MJ, Chakrabarti L, Eng J, Harker Rhodes C, Lutz T, Nieto A, Denny Liggitt H, Warner 614 S, Fields J, Stöger R, and Fiering S. Deletion of the Chd6 exon 12 affects motor coordination. 615 Mammalian Genome 21: 130-142, 2010. 616 41. Lee KY, Singh MK, Ussar S, Wetzel P, Hirshman MF, Goodyear LJ, Kispert A, and Kahn CR. 617 Tbx15 controls skeletal muscle fibre-type determination and muscle metabolism. Nature 618 communications 6: 8054, 2015. 619 42. Leick L, Hellsten Y, Fentz J, Lyngby SS, Wojtaszewski JF, Hidalgo J, and Pilegaard H. PGC-620 1alpha mediates exercise-induced skeletal muscle VEGF expression in mice. American journal of 621 physiology Endocrinology and metabolism 297: E92-103, 2009. 622 43. Li WB, Ren J, Zhu WC, Guo BL, Yang B, Liu LT, Ding NS, Ma JW, Li L, and Huang LS. Mapping 623 QTL for porcine muscle fibre traits in a White Duroc x Erhualian F(2) resource population. Journal of 624 animal breeding and genetics = Zeitschrift fur Tierzuchtung und Zuchtungsbiologie 126: 468-474, 625 2009. 626 44. Lillioja S, Young AA, Culter CL, Ivy JL, Abbott WG, Zawadzki JK, Yki-Jarvinen H, Christin L, 627 Secomb TW, and Bogardus C. Skeletal muscle capillary density and fiber type are possible 628 determinants of in vivo insulin resistance in man. The Journal of clinical investigation 80: 415-424, 629 1987. 630 45. Lionikas A, Blizard DA, Gerhard GS, Vandenbergh DJ, Stout JT, Vogler GP, McClearn GE, and 631 Larsson L. Genetic determinants of weight of fast- and slow-twitch skeletal muscle in 500-day-old 632 mice of the C57BL/6J and DBA/2J lineage. Physiological genomics 21: 184-192, 2005. 633 46. Lionikas A, Blizard DA, Vandenbergh DJ, Glover MG, Stout JT, Vogler GP, McClearn GE, and 634 Larsson L. Genetic architecture of fast- and slow-twitch skeletal muscle weight in 200-day-old mice 635 of the C57BL/6J and DBA/2J lineage. Physiological genomics 16: 141-152, 2003. 636 47. Lionikas A, Cheng R, Lim JE, Palmer AA, and Blizard DA. Fine-mapping of muscle weight QTL 637 in LG/J and SM/J intercrosses. *Physiological genomics* 42a: 33-38, 2010. 638 48. Lionikas A, Meharg C, Derry JM, Ratkevicius A, Carroll AM, Vandenbergh DJ, and Blizard 639 DA. Resolving candidate genes of mouse skeletal muscle QTL via RNA-Seq and expression network 640 analyses. BMC genomics 13: 592, 2012. 641 49. Luff AR, and Goldspink G. Total number of fibers in muscles of several strains of mice. 642 Journal of animal science 30: 891-893, 1970. 643 50. MacArthur DG, Seto JT, Chan S, Quinlan KG, Raftery JM, Turner N, Nicholson MD, Kee AJ, 644 Hardeman EC, Gunning PW, Cooney GJ, Head SI, Yang N, and North KN. An Actn3 knockout mouse

645 provides mechanistic insights into the association between alpha-actinin-3 deficiency and human 646 athletic performance. Human molecular genetics 17: 1076-1086, 2008. 647 MacDougall JD, Sale DG, Alway SE, and Sutton JR. Muscle fiber number in biceps brachii in 51. 648 bodybuilders and control subjects. Journal of applied physiology: respiratory, environmental and 649 exercise physiology 57: 1399-1403, 1984. 650 Malek M, Dekkers JC, Lee HK, Baas TJ, Prusa K, Huff-Lonergan E, and Rothschild MF. A 52. 651 molecular genome scan analysis to identify chromosomal regions influencing economic traits in the 652 pig. II. Meat and muscle composition. Mammalian genome : official journal of the International 653 Mammalian Genome Society 12: 637-645, 2001. 654 Mannion AF, Dumas GA, Cooper RG, Espinosa FJ, Faris MW, and Stevenson JM. Muscle 53. 655 fibre size and type distribution in thoracic and lumbar regions of erector spinae in healthy subjects 656 without low back pain: normal values and sex differences. Journal of anatomy 190 (Pt 4): 505-513, 657 1997. 658 54. McPherron AC, Lawler AM, and Lee SJ. Regulation of skeletal muscle mass in mice by a new 659 TGF-beta superfamily member. Nature 387: 83-90, 1997. 660 Nii M, Hayashi T, Mikawa S, Tani F, Niki A, Mori N, Uchida Y, Fujishima-Kanaya N, Komatsu 55. 661 M, and Awata T. Quantitative trait loci mapping for meat quality and muscle fiber traits in a 662 Japanese wild boar x Large White intercross. Journal of animal science 83: 308-315, 2005. 663 56. Nikolskiy I, Conrad DF, Chun S, Fay JC, Cheverud JM, and Lawson HA. Using whole-genome 664 sequences of the LG/J and SM/J inbred mouse strains to prioritize quantitative trait genes and 665 nucleotides. BMC genomics 16: 415, 2015. 666 57. Nimmo MA, Wilson RH, and Snow DH. The inheritance of skeletal muscle fibre composition 667 in mice. Comparative biochemistry and physiology A, Comparative physiology 81: 109-115, 1985. 668 58. Pourhassan M, Bosy-Westphal A, Schautz B, Braun W, Gluer CC, and Muller MJ. Impact of 669 body composition during weight change on resting energy expenditure and homeostasis model 670 assessment index in overweight nonsmoking adults. The American journal of clinical nutrition 99: 671 779-791, 2014. 672 59. Rehfeldt C, Ott G, Gerrard DE, Varga L, Schlote W, Williams JL, Renne U, and Bunger L. 673 Effects of the compact mutant myostatin allele Mstn (Cmpt-dl1Abc) introgressed into a high growth 674 mouse line on skeletal muscle cellularity. Journal of muscle research and cell motility 26: 103-112, 675 2005. 676 60. Roepstorff C, Donsmark M, Thiele M, Vistisen B, Stewart G, Vissing K, Schjerling P, Hardie 677 DG, Galbo H, and Kiens B. Sex differences in hormone-sensitive lipase expression, activity, and 678 phosphorylation in skeletal muscle at rest and during exercise. American journal of physiology 679 Endocrinology and metabolism 291: E1106-1114, 2006. 680 61. Sainz N, Rodriguez A, Catalan V, Becerril S, Ramirez B, Gomez-Ambrosi J, and Fruhbeck G. 681 Leptin administration favors muscle mass accretion by decreasing FoxO3a and increasing PGC-1alpha 682 in ob/ob mice. PloS one 4: e6808, 2009. 683 62. Schiaffino S. Fibre types in skeletal muscle: a personal account. Acta physiologica (Oxford, 684 England) 199: 451-463, 2010. 685 Simoneau JA, and Bouchard C. Genetic determinism of fiber type proportion in human 63. 686 skeletal muscle. FASEB journal : official publication of the Federation of American Societies for 687 Experimental Biology 9: 1091-1095, 1995. 688 64. Simoneau JA, and Bouchard C. Human variation in skeletal muscle fiber-type proportion and 689 enzyme activities. The American journal of physiology 257: E567-572, 1989. 690 65. Sosnicki A. Histopathological observation of stress myopathy in M. longissimus in the pig 691 and relationships with meat quality, fattening and slaughter traits. Journal of animal science 65: 584-692 596, 1987. 693 66. Srikanthan P, and Karlamangla AS. Muscle Mass Index As a Predictor of Longevity in Older

694 Adults. The American Journal of Medicine 127: 547-553, 2014.

695 67. Staron RS, Hagerman FC, Hikida RS, Murray TF, Hostler DP, Crill MT, Ragg KE, and Toma K. 696 Fiber type composition of the vastus lateralis muscle of young men and women. The journal of 697 histochemistry and cytochemistry : official journal of the Histochemistry Society 48: 623-629, 2000. 698 68. Stavaux D, Art T, McEntee K, Reznick M, and Lekeux P. Muscle fibre type and size, and 699 muscle capillary density in young double-muscled blue Belgian cattle. Zentralblatt fur 700 Veterinarmedizin Reihe A 41: 229-236, 1994. 701 Timson BF, Bowlin BK, Dudenhoeffer GA, and George JB. Fiber number, area, and 69. 702 composition of mouse soleus muscle following enlargement. Journal of applied physiology 703 (Bethesda, Md : 1985) 58: 619-624, 1985. 704 70. Totsuka Y, Nagao Y, Horii T, Yonekawa H, Imai H, Hatta H, Izaike Y, Tokunaga T, and Atomi 705 Y. Physical performance and soleus muscle fiber composition in wild-derived and laboratory inbred 706 mouse strains. Journal of applied physiology (Bethesda, Md : 1985) 95: 720-727, 2003. 707 71. Van den Maagdenberg K, Stinckens A, Lefaucheur L, Buys N, and De Smet S. The effect of 708 mutations in the insulin-like growth factor-II and ryanodine receptor-1 genes on biochemical and 709 histochemical muscle fibre characteristics in pigs. *Meat science* 79: 757-766, 2008. 710 72. van der Laarse WJ, Diegenbach PC, and Maslam S. Quantitative histochemistry of three 711 mouse hind-limb muscles: the relationship between calcium-stimulated myofibrillar ATPase and 712 succinate dehydrogenase activities. The Histochemical journal 16: 529-541, 1984. 713 73. Vaughan HS, Aziz U, Goldspink G, and Nowell NW. Sex and stock differences in the 714 histochemical myofibrillar adenosine triphosphatase reaction in the soleus muscle of the mouse. The 715 journal of histochemistry and cytochemistry : official journal of the Histochemistry Society 22: 155-716 159, 1974. 717 74. Wade AJ, Marbut MM, and Round JM. Muscle fibre type and aetiology of obesity. Lancet 718 335: 805-808, 1990. 719 75. Wang YX, Zhang CL, Yu RT, Cho HK, Nelson MC, Bayuga-Ocampo CR, Ham J, Kang H, and 720 **Evans RM**. Regulation of muscle fiber type and running endurance by PPARdelta. *PLoS biology* 2: 721 e294, 2004. 722 76. Wilkins BJ, Dai YS, Bueno OF, Parsons SA, Xu J, Plank DM, Jones F, Kimball TR, and 723 Molkentin JD. Calcineurin/NFAT coupling participates in pathological, but not physiological, cardiac 724 hypertrophy. Circulation research 94: 110-118, 2004. 725 Wimmers K, Fiedler I, Hardge T, Murani E, Schellander K, and Ponsuksili S. QTL for 77. 726 microstructural and biophysical muscle properties and body composition in pigs. BMC genetics 7: 15, 727 2006. 728 Wirtz P, Loermans HM, Peer PG, and Reintjes AG. Postnatal growth and differentiation of 78. 729 muscle fibres in the mouse. I. A histochemical and morphometrical investigation of normal muscle. 730 Journal of anatomy 137 (Pt 1): 109-126, 1983. 731 79. Wu C, Orozco C, Boyer J, Leglise M, Goodale J, Batalov S, Hodge CL, Haase J, Janes J, Huss 732 JW, 3rd, and Su AI. BioGPS: an extensible and customizable portal for querying and organizing gene 733 annotation resources. Genome biology 10: R130, 2009. 734 80. Zhang Y, Xu P, Lu C, Kuang Y, Zhang X, Cao D, Li C, Chang Y, Hou N, Li H, Wang S, and Sun X. 735 Genetic linkage mapping and analysis of muscle fiber-related QTLs in common carp (Cyprinus carpio 736 L.). Marine biotechnology (New York, NY) 13: 376-392, 2011.

- So L.). Wurne biotechnology (New F
- 737



Α







