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- 1 The effect of caloric restriction on the forelimb skeletal muscle fibers of
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25 Abstract

Skeletal muscle mass loss has a broad impact on body performance and 26 physical activity. Muscle wasting occurs due to genetic mutation as in muscular 27 dystrophy, age-related muscle loss (sarcopenia) as well as in chronic wasting 28 disorders as in cancer cachexia. Food restriction reduces muscle mass 29 underpinned by increased muscle protein break down. However the influence 30 of dietary restriction on the morphometry and phenotype of forelimb muscles in 31 a genetically modified myostatin null mice are not fully characterized. The effect 32 of a five week dietary limitation on five anatomically and structurally different 33 forelimb muscles was examined. C57/BL6 wild type (*Mstn*^{+/+}) and *myostatin* null 34 (*Mstn^{-/-}*) mice were either given a standard rodent normal daily diet ad libitum 35 (ND) or 60% food restriction (FR) for a 5 week period. M. triceps brachii Caput 36 laterale (T.lateral), M. triceps brachii Caput longum (T.long), M. triceps brachii 37 Caput mediale (T.medial), M. extensor carpi ulnaris (ECU) and M. flexor carpi 38 39 ulnaris (FCU) were dissected. weighted and processed for immunohistochemistry. Muscle mass, fibers cross sectional areas (CSA) and 40 myosin heavy chain types IIB, IIX, IIA and type I were analyzed. We provide 41 evidence that caloric restriction results in muscle specific weight reduction with 42 the fast myofibers being more prone to atrophy. We show that slow fibers are 43 less liable to dietary restriction induced muscle atrophy. The effect of dietary 44 restriction was more pronounced in *Mstn^{-/-}* muscles to implicate the oxidative 45 fibers compared to Mstn^{+/+}. Furthermore, peripherally located myofibers are 46 more susceptible to dietary induced reduction compared to deep fibers. We 47 additionally report that dietary restriction alters the glycolytic phenotype of the 48 *Mstn^{-/-}* into the oxidative form in a muscle dependent manner. In summary our 49 study shows that calorie restriction alters muscle fiber profile of forelimb 50 muscles of Myostatin null mice. 51

52 **Keywords:** Skeletal muscle, Myostatin, food restriction, myosin heavy chain

53

54 **1. Introduction**

Skeletal muscle the main protein reservoir in the body, is a highly adaptable 55 tissue that changes its physical as well as composition based on physiological 56 demands. Mechanical and nutritional stimuli cause an increase in muscle mass. 57 In contrast undernutrition, aging and diseases as cancer cachexia reduces the 58 muscle mass. Unbalanced nutrient intake with increased energy requirements 59 enhances the muscle catabolism resulting in fiber atrophy and muscle mass 60 loss (Koskelo et al., 1990). Muscle loss also occurring in progressive wasting 61 diseases e.g. cancer cachexia (Bruggeman et al., 2016) and age-related 62 sarcopenia leads to a decrease in muscle mass and strength (Thomas, 2007). 63 Additionally the role of physiological and metabolic stimuli e.g. physical activity, 64 disuse and immobilization, food restriction, drugs and diseases cause 65 molecular and cellular dysregulation which results in loss of muscle mass 66 (Carmeli and Reznick, 1994). It has been reported that ageing induces muscle 67 mass loss of 16%, 18%, 37% and 38% for soleus, extensor digitorum longus, 68 plantaris and gastrocnemius muscles respectively with greater fiber area loss 69 compared to the decrease in body mass (Brown and Hasser, 1996). Similarly it 70 was reported that dietary restriction causes a decrease in muscle fiber size but 71 not fiber number in chicken and rabbits (Tanaka et al., 1992 and Timson et al., 72 1983). In addition the glycolytic fibers were selectively decreased in the cross 73 sectional area lead to a proportional increase in the area of oxidative fibers in 74 bovine muscle (Greenwood et al., 2009). Muscle phenotype also changed 75 according to the dietary challenges, It has been reported that dietary deprivation 76 for 48 hours upregulates the expression of fast myosin heavy chain 2b mRNA 77 with no change in fiber type composition for EDL and soleus muscles in rat 78 (Mizunoya et al., 2013b) However 4 weeks fatty diet administration induced 79 reduction in the fast MHC2b and improved the oxidative metabolism in the EDL 80 of rat (Matsakas et al., 2013; Mizunoya et al., 2013a). Previously, it was shown 81 that fasting induces muscle proteolysis via glucocorticoid activation (Wing and 82 Goldberg, 1993) as well as causing an increase in the ATP-dependent 83 proteolysis and upregulation of ubiquitin conjugates and polyubiquitins in rat 84 skeletal muscle (Medina et al., 1991). 85

Myostatin is a TGF- β family member of secreted proteins that negatively 86 regulates skeletal muscle proliferation (Sharma et al., 2001). Myostatin protein 87 interference via knock out results in significant myofiber hyperplasia and 88 hypertrophy (McPherron et al., 1997). Furthermore, systemic administration of 89 Myostatin induced muscle atrophy in mice (Zimmers et al., 2002). Increased 90 myostatin expression was also associated with muscle wasting in cancer 91 cachexia, ageing, after HIV infection and in the course of chronic obstructive 92 pulmonary disease (COPD) (Costelli et al., 2008; Gonzalez-Cadavid et al., 93 94 1998; Plant et al., 2010; Yarasheski et al., 2002). Myostatin knock out mice showed a greater atrophy in response to unloading (McMahon et al., 2003). 95 Our previous study demonstrated a massive reduction in forelimb muscles 96 mass of aged myostatin null compared to wild type mice (Elashry et al., 2009). 97 These studies suggest that hypertrophic muscle has decreased adaptability 98 compared to WT. In this study we hypothesized that five weeks of 60% dietary 99 restriction would impact to a greater degree on the muscle of Mstn^{-/-} compared 100 to WT. We provide evidence that caloric restriction results in muscle specific 101 weight reduction with the fast myofibers being more prone to atrophy. We show 102 that slow fibers are less liable to dietary restriction induced muscle atrophy. 103 Furthermore, peripherally located myofibers are more susceptible to dietary 104 induced reduction. Additionally, we report that dietary restriction alters the 105 glycolytic phenotype of the myostatin null into more oxidative form in a muscle 106 dependent manner. 107

108 2. Materials and Methods

109 2. 1. Animals

Three months old male transgenic myostatin knockout (*Mstn^{-/-}*) and C57/BL6 110 wild type littermates (*Mstn*^{+/+}) mice were bred in the biological resource unit, 111 Reading University, UK. Mstn^{-/-} mice were generously provided from Dr Lee 112 (McPherron et al., 1997). The experiments were performed under a project 113 license from the United Kingdom Home Office in agreement with the Animals 114 (Scientific Procedures) Act 1986 (license number 7516). The University of 115 Reading Animal Care and Ethical Review Committee (AWERB) approved all 116 procedures. The mice were divided into 4 subgroups: a normal diet group (N=8) 117

 $Mstn^{+/+}$ and $Mstn^{-/-}$ kept with a normal diet with food and water provided *ad libitum* and a food restriction group (N=8) with an up to 60% reduction of the daily diet compared to normal mice for a 5 week period. Animals were humanely sacrificed via Schedule 1 killing.

122 2. 2. Tissue isolation and processing

Aged matched mice were sacrificed through schedule-1 killing following the 123 approval of the ethics committee. Five forelimb muscles were carefully chosen 124 to represent the arm and the forearm region *M. triceps brachii Caput longum*, 125 T.long; M. triceps brachii Caput laterale, T.lateral; M. triceps brachii Caput 126 mediale, T.medial; M. extensor carpi ulnaris, ECU; and M. flexor carpi ulnaris, 127 FCU. The muscles were dissected, snap frozen using isopentane pre-cooled 128 with liquid nitrogen and were embedded in tissue tech OCT (Sakura, VWR) 129 using dry ice cooled ethanol.10 µm transverse mid-belly cryosections for each 130 muscle were obtained on a poly-L-lysine coated slides (VWR) and left to dry at 131 room temperature for 1 hour and stored at -80C°. 132

133 2. 3. Immunohistochemical staining

Frozen muscle sections were washed in PBS for 10 minutes, permeabilized 134 using buffer solution containing 20 mM Hepes, 300 mM sucrose, 50 mM NaCl, 135 3 mM MgCl₂ and 0.5% Triton-X100 (pH7) at for 15 minutes. Non-specific 136 binding was blocked using wash buffer (5% fetal calf serum (v/v) and 0.05% 137 Triton X-100 (v/v) in PBS) for 30 minutes at room temperature before the 138 addition of antibodies. Myosin heavy chain (MHC) type I, IIA and IIB expressing 139 myofibers were identified by incubating the slides with A4.840 mouse IgM (1:1), 140 A.474 mouse IgG (1:4) and BFF3 mouse IgM (1:1) monoclonal primary 141 antibodies (DSHB) respectively as previously described (Matsakas et al., 142 2009). The primary antibodies were tested separately against the 143 corresponding MHC type on a muscle section and the following section with no 144 primary antibody was served as a negative control (Fig. 1g-I). Additionally, MHC 145 type IIA⁺ and IIB⁺ myofibers were stained together and MHC type I⁺ myofibers 146 were stained on consecutive section. All primary antibodies were pre-blocked 147 in washing buffer 30 minutes prior to use and incubated on samples overnight 148 at 4°C. Primary antibodies were highlighted via incubation with Alexa Fluor 633 149

goat anti-mouse IgM (Molecular Probes A21046, 1:200) for MHC I and MHC
IIB and Alexa Fluor 488 Goat-anti-mouse IgG (Molecular probes A11029,
1:200) for MHC IIA secondary antibodies in dark at room temperature for 45
minutes. Slides were mounted using a fluorescent mounting medium (Mowiol
4-88, Calbiochem) containing 2.5 μg/ml DAPI.

155 2. 4. Imaging and quantifications

Muscle sections were examined and photographed using a Zeiss Axioscop2 156 fluorescence microscope loaded with a digital camera and connected to a 157 computer equipped with the axiovision computer software (Zeiss). The whole 158 muscle cross sections was reconstructed, total number, number of IIB, IIA and 159 type I fibers quantifications were performed by manual counting using the 160 Photoshop CS6 extended software. Type IIX fibers were counted via 161 subtraction of all other fibers from the total fiber number. Cross sectional area 162 (CSA) for each fiber type for all muscles were measured using the axiovision 163 software (Zeiss). 164

165 2. 5. Statistical analysis

The effect of genotype (Mstn^{+/+} vs. Mstn^{-/-}) and diet regime (ND vs. FR) on total 166 muscle fiber numbers, CSA and the percentage of type IIB, IIX, IIA and I fibers 167 were analyzed by two-way ANOVA for T.lateral, T.long, T.medial, ECU and 168 FCU muscles. The effect of genotype, diet regime and region of myofiber 169 (peripheral vs central) on CSA for each muscle were analyzed via three-way 170 ANOVA. Multiple comparisons and the interactions were evaluated using 171 Tukey's Post Hoc test. Chi-square test was performed to examine frequency 172 distribution of fiber types CSA following FR. Statistical analysis was conducted 173 by using Graph Pad prism 6 software. All values are presented as mean ± SEM, 174 *P* values less than 0.05 were considered significant. 175

176 **3. Results**

177 3. 1. Dietary restriction induces body weight and muscle mass specific
 178 reduction

In order to evaluate the effect of dietary restriction on skeletal muscle of Mstn^{+/+} 179 and *Mstn^{-/-}* mice for a five week period, average body weight measurements (g) 180 revealed that *Mstn^{-/-}ND* mice were significantly heavier compared to *Mstn^{+/+}ND* 181 mice (P<0.001). However both genotypes showed approximately 33% 182 reduction in the body mass following dietary restriction (Fig. 1a). Individual 183 muscle weight measurements showed that *Mstn^{-/-}ND* exhibited increases in 184 muscle mass of T.lateral (P<0.0001), T.long (P<0.0001), T.medial (P<0.05), 185 ECU (P<0.01) and FCU muscles (P<0.0001) compared to Mstn^{+/+}ND. To 186 evaluate whether dietary restriction impact on different muscles mass. Our data 187 showed that dietary restriction induced muscle specific weight loss e.g. T.lateral 188 showed large reduction in muscle mass for Mstn^{-/-}FR and Mstn^{+/+}FR (P<0.0001 189 and P<0.001) respectively compared to genotype-matched ND as well as 190 significant interaction (P<0.001) between diet and genotype (Fig.1b). Similarly, 191 T.long muscle exhibited more pronounced reduction in the Mstn⁻FR and 192 Mstn^{+/+}FR (P<0.0001 and P<0.001) respectively compared to ND matched 193 control also with significant interaction (P<0.0001) between the effect of diet 194 and the genotype (Fig. 1c). Furthermore, ECU muscle showed more weight 195 reduction following FR in *Mstn^{-/-}* than *Mstn^{+/+}* (P<0.001 and P<0.05) compared 196 to genotype matched control (Fig.1e). On the other hand, T.medial and FCU 197 muscles displayed a diet induced decreases in the muscle mass only in Mstn^{-/-} 198 mice (P<0.01 and 0.0001) compared to genotype-matched control (Fig.1d and 199 f). Representative immunofluorescent staining for T.long muscle of *Mstn*^{+/+}ND 200 show MHC types IIA, IIB and I expressing myofibers compared to the negative 201 control (Fig. 1g-I). 202

3. 2. Caloric restriction does not alter muscle fiber number

Based on previous experiments (Elashry et al., 2009) we examined whether a FR related muscle mass decrease was due to either loss in fiber number or loss of fiber size. Quantification of the average fibers number revealed that myostatin deletion increases muscle fiber number for T.lateral and T.long (P<0.0001), ECU and FCU (P<0.01) muscles on normal diet could be detected. In contrast, T.medial muscle displayed no fiber number change in *Mstn*^{-/-}. However dietary restriction for five weeks caused no change in the fibers number for all muscles of both genotypes (Fig. 2a, b, c, d and e). Representatives whole muscle reconstruction showed clearly the effect of *myostatin* deletion and dietary restriction on the cross section of the FCU muscle of $Mstn^{+/+}$ and $Mstn^{-/-}$ compared to normal diet control (Fig. 2f-i).

3. 3. Fast fibers are more prone to caloric restriction-induced muscle mass loss

Next we evaluated the impact of dietary restriction on individual muscle fiber 217 size via quantification of the CSA (μ m²) of IIB, IIX, IIA and type I fibers per 218 muscle. A common trend that dietary restriction reduced the CSA of glycolytic 219 type IIB fibers in $Mstn^{-/-}$ and $Mstn^{+/+}$ (P<0.0001) compared to controls was 220 observed (Fig. 3). Type IIB fibers CSA measurement of T.lateral muscle 221 displayed pronounced reduction (2074 \pm 7 to 1232 \pm 3 μ m²) in *Mstn^{-/-}* (P<0.0001) 222 compared to (1605±4 to 1246±3 μ m²) in *Mstn*^{+/+} (P<0.0001) following FR. 223 These results were supported with significant interaction (P<0.0001) indicating 224 the effect of FR on *Mstn^{-/-}* compared to *Mstn^{+/+}* (Fig. 3a). Furthermore, T.long 225 muscle showed similar reduction in type IIB fibers CSA (3596±52 to 1304±3 226 μ m²) and (3118±8 to 1085±3 μ m²) for *Mstn^{-/-}* and *Mstn^{+/+}* respectively 227 (P<0.0001) following dietary restriction and compared to genotype-matched 228 control (Fig. 3b). On the other hand, measurements of type IIB fibers CSA 229 showed more loss in the T.medial and ECU muscles of *Mstn^{-/-}* (P<0.0001) 230 compared to *Mstn*^{+/+} following FR. E.g. T.medial muscle of *Mstn*^{-/-}demonstrated 231 1647±42 to 1090.6±3 µm² CSA reduction (P<0.0001) compared to 1479.5±4 to 232 1257±3 μ m², (P<0.0001) reduction in *Mstn*^{+/+} provided with significant 233 interaction (P<0.0001) between the effect of diet and genotype (Fig. 3c and d). 234 Next, type IIX fibers CSA measurement for T.lateral, ECU and FCU muscles 235 displayed significant decreases in the CSA for both *Mstn^{-/-}* (P<0.05, 0.01 and 236 0.0001) and *Mstn*^{+/+} (P<0.01 and 0.0001) following dietary restriction compared 237 to control (Fig. 3a, d and e) however IIX fibers measurements showed more 238 loss in T.long and T.medial muscles in the $Mstn^{-1}$ (P<0.0001 and P<0.01) 239 compared to *Mstn*^{+/+} following FR (Fig. 3b and c). Additionally, we examined 240 whether the effect of dietary limitation involves the oxidative fibers, analysis of 241 type IIA fibers CSA demonstrated reduction in the T.long and T.medial muscles 242

(P<0.0001 and 0.05) respectively only in the $Mstn^{-/-}$ compared to $Mstn^{+/+}$ under 243 the FR regime (Fig. 3b and c). Surprisingly, IIA fibers CSA for ECU and FCU 244 muscles showed significant increase in *Mstn^{-/-}*FR (P<0.01 and 0.0001) 245 compared to *Mstn^{-/}*ND (Fig. 3d and e). Furthermore, we tested the oxidative 246 capacity of the muscles following dietary restriction by analyzing type I (slow 247 oxidative) fibers CSA, our results displayed large reduction in T.long muscle of 248 $Mstn^{-/-}$ (P<0.0001) compared to $Mstn^{+/+}$ (Fig. 3b). These data demonstrated that 249 glycolytic fibers were more liable to dietary restriction followed by IIX, IIA and I 250 oxidative fibers. Furthermore the impact of dietary restriction was more 251 pronounced in some of the *Mstn⁻⁻* muscles to implicate the oxidative fibers (IIA 252 and I) compared to Mstn^{+/+}. In order to confirm our data, T.long fibers CSA 253 measurements were categorized into small (0-1000 and 1000-2000 µm²), 254 medium (2000-4000 µm²) and large (4000-6000 µm²) for type IIB, small (0-1000 255 μ m²), medium (1000-2000 μ m²) and large fibers (2000-4000 μ m²) for IIX, IIA 256 and type I fibers. Frequency distribution assessment of type IIB, IIX, IIA and I 257 fibers of T.long muscle showed significant CSA shift from large size into the 258 small size population (P<0.0001) following dietary restriction compared to 259 genotype-matched control (Fig. 3f, g, h and i). 260

3. *4. Dietary restriction causes peripheral myofiber atrophy*

Our results show that dietary restriction primarily target fast MHCIIB fibers. To 262 address whether the anatomical location of the fiber had an effect on the 263 myofibers CSA following caloric restriction, we analyzed the CSA of the 264 peripheral (PF) and central (CT) fibers within the mid-belly of the corresponding 265 muscle. Our results revealed that in T.lateral, T.long and ECU muscles but not 266 T.medial muscle showed a common observation that type IIB myofibers CSA 267 were larger in the PF (P>0.0001) compared to CT region for Mstn^{-/-} and Mstn^{+/+} 268 on normal diet (Fig. 4a, c, f and i). Furthermore, FCU muscles demonstrated 269 larger CSA of PF IIB compared to CT fibers (P<0.0001) only in the Mstn^{-/-} on a 270 normal diet (Fig. 4I). Furthermore, FR induced significant reduction in the CSA 271 of PF IIB fibers (P<0.0001) compared to CT fibers e.g. T.lateral muscle showed 272 (850±87 vs. 1247±67 µm²) and (609±1 vs. 1348±2 µm²) for Mstn^{-/-} and Mstn^{+/+} 273 respectively also significant interaction was detected (P<0.0001) indicating the 274

interference of diet and genotype (Fig. 4a). Next we analyzed type IIX fibers, 275 larger IIX fibers CSA of both regions in T.lateral muscle of *Mstn*^{+/+} (P<0.0001) 276 and T.long muscle of *Mstn^{-/-}* (P<0.0001) on normal diet were observed. Also, 277 PF IIX fibers of *Mstn^{-/-}*FR showed more reduction in the CSA in T.lateral and 278 T.long muscles (P<0.05 and P<0.0001) compared to *Mstn⁻*ND respectively 279 (Fig. 4b and d). However, reduction in the PF IIX fibers of both genotypes were 280 noticed in the T.medial (P<0.01), ECU and FCU muscles (P<0.0001) following 281 FR (Fig. 4g, j and m). Moreover, CT type IIX fibers displayed reduction in T.long 282 and T.medial muscles of $Mstn^{-}$ (P<0.0001) as well as T.lateral muscle of 283 Mstn^{+/+} (P<0.0001) following FR. Additionally, PF type IIX fiber CSA 284 demonstrated further reduction in T.medial (P<0.001 and P<0.0001) and ECU 285 muscle (P<0.0001) of both genotypes compared to CT fibers (Fig. 4 g and j) 286 following FR. On the other hand, more reduction in the PF IIX fibers was 287 detected in T.lateral and FCU muscles of only *Mstn^{-/-}* (P<0.0001) compared to 288 CT fibers following FR (Fig. 4b and m), also significant interaction (P<0.01) 289 indicating the effect of diet on *Mstn^{-/-}*. In contrast, analysis of type IIA fibers 290 showed some variations, PF IIA fibers of T.medial muscle displayed more 291 reduction in the CSA (P<0.01) compared to CT fibers only in *Mstn*^{+/+} (Fig. 4h) 292 however PF fibers of ECU muscle showed more CSA reduction compared to 293 CT fibers (P<0.0001 and P<0.001) for *Mstn^{-/-}* and *Mstn^{+/+}* respectively following 294 dietary restriction (Fig. 4k). However, FCU muscle displayed larger CSA 295 reduction in PF IIA fibers only in the *Mstn^{-/-}* compared to CT fibers after dietary 296 restriction (Fig. 4n). Type I fibers was only localized in the central part of the 297 muscle therefore were excluded from this comparison. 298

3. 5. Caloric restriction induces a slow phenotype conversion in a muscle specific manner

Our previous study demonstrated that *myostatin* deletion induced various degrees of slow to fast fiber type shift in the forelimb muscles (Elashry et al., 2009). Therefore, we examined whether dietary restriction alters the muscle phenotype of the *Mstn*^{-/-} compared to *Mstn*^{+/+}. Quantification of the average percentage of each fiber type population per muscle demonstrated percentage increases of type IIB fibers in the T.lateral, T.long, T.medial, ECU and FCU

muscles Mstn^{-/-}ND (P<0.0001) compared to Mstn^{+/+}ND. However FR altered 307 the fiber type composition for both genotypes (P<0.001) e.g. T.lateral muscle 308 demonstrated a reduction in the type IIB fibers (P<0.0001 and P<0.05) and 309 increases of type IIX fibers (P<0.0001 and P<0.05) for Mstn^{-/-}FR and Mstn^{+/+}FR 310 respectively compared to genotype matched control (Fig. 5a and b). Also FR 311 induced an increase in the type IIA fibers of *Mstn^{-/-}*FR compared to ND, however 312 *Mstn*^{+/+}FR showed a reduction in type IIA fibers compared to ND. These data 313 were confirmed by significant interaction between both genotypes (P<0.01, Fig. 314 5c). Furthermore, T.long muscle showed a reduction in type IIB fibers 315 (P<0.0001) with an increase in type IIX fibers (P<0.001 and P<0.0001) for Mstn⁻ 316 ^{/-}FR and *Mstn*^{+/+}FR respectively compared to genotype-matched ND (Fig. 5d) 317 and e). Although there were no significant changes, *Mstn^{/-}*FR demonstrated 318 marked increases in type IIA fibers compared to *Mstn^{-/-}ND*. However *Mstn^{+/+}FR* 319 showed significant increase in type IIA fibers (P<0.01) compared to Mstn^{+/+}ND 320 (Fig. 5f) as well as FR resulted in marked increase in type I fibers in both 321 genotypes (P<0.01) compared to ND control (Fig. 5g). On the other hand, 322 T.medial muscle analysis revealed fiber type population of 33%, 34%, 26% and 323 7% for IIB, IIX, IIA and I fibers respectively in the *Mstn*^{+/+}ND and showed almost 324 no fiber type shift following dietary restriction in the *Mstn^{-/-}* however *Mstn^{+/+}* FR 325 displayed fiber shift into type I fibers (IIB→IIX→IIA→I) following the same 326 treatment (P<0.05) compared to ND. These data was supported by significant 327 interaction (P<0.05) between diet and genotype effect (Fig. 5h, i and j and k). 328 On the other hand, ECU muscle displayed a different fiber shift pattern with 329 dietary restriction whereas IIB to IIX fibers transformation in the Mstn^{-/-} 330 (P<0.0001 and P<0.01) respectively (Fig. 5I and m) and exhibited type IIA to 331 type I fiber shift in *Mstn*^{+/+} (P<0.001) compared to ND (Fig. 5n and o). Similarly, 332 FCU muscle FR induced type IIB to IIX fiber shift for *Mstn^{-/-}* (P<0.0001) 333 compared to ND (Fig. 5p and q) however demonstrated fibers shift from IIB 334 directly into IIA fibers but not type IIX fibers in $Mstn^{+/+}$ (P<0.0001 and P<0.05) 335 *Mstn*^{+/+}ND (Fig. 5r compared to and s). Representative double 336 immunofluorescent images of T.long muscle for *Mstn*^{+/+} and *Mstn*^{-/-} showed the 337 transformation of the glycolytic fibers type IIB and IIX into the oxidative type IIA 338 fibers following dietary restriction. A negative control without addition of primary 339

antibodies was performed on a consecutive muscle section (Fig. 5t, u, v, w andx).

342 4. Discussion

Dietary intake plays a critical role to maintain the physiological composition of 343 the skeletal muscle. Previous studies revealed that myostatin deletion results 344 345 in an increase in muscle mass due to an increase in fiber number and fiber size (McPherron et al., 1997). However, the impact of dietary restriction on 346 anatomically and structurally different forelimb muscles of the myostatin null are 347 not fully characterized. Our results suggest that the impact of dietary restriction 348 was greater in the *Mstn^{-/-}* than in wild-type muscle. In agreement with our 349 results, previous work showed that dietary limitation caused more muscle loss 350 in the hind limb muscles of the *Mstn^{-/-}* compared to *Mstn^{+/+}* suggesting higher 351 catabolic activities (Matsakas et al., 2013). These results suggest that the 352 hypertrophic muscle of *Mstn^{-/-}* is more prone to protein breakdown than that of 353 wild-types. In support of this notion is the finding that *Mstn^{-/-}* mice exhibited 354 more muscle mass loss during hind limb unloading (McMahon et al., 2003). Our 355 results indicate that muscle size (enlarged) and composition (fast) might play a 356 role in the degree of muscle loss following dietary limitation. This in line with 357 previous reports indicating that fast twitch IIB fibers of rodents and monkeys 358 are more sensitive to starvation compared to slow twitch fiber (Goldspink and 359 Ward, 1979; Li and Goldberg, 1976; Matsakas et al., 2013; McKiernan et al., 360 2012). However we show that there is also a regional specificity of MHC types 361 362 in terms of atrophy.

We propose that the dietary restriction induced muscle loss is regulated via 363 364 inhibition of IGF-1 signaling which results in upregulation of atrogin-1 and MuRF-1 expression. A similar study showed that undernutrition for 28 days 365 causes reduced growth rate via regulation of IGF-1 and myostatin expressions 366 (Jeanplong et al., 2015). We shows that the oxidative fibers are resistant to 367 dietary restriction. In agreement with our results, it has been shown that the 368 atrophy resistance of the oxidative fibers was due to a protection afforded by 369 peroxisome proliferator-activated receptor gamma coactivator-1 370 alpha (PGC1a) (Wang and Pessin, 2013). Unexpectedly, dietary restriction causes 371

372 CSA increases in type IIA fibers for ECU and FCU of *Mstn^{-/-}* which might be an
 373 adaptive mechanism to compensate the decrease in glycolytic fibers number.

Analysis of the muscles phenotypes revealed a general conversion toward slow 374 oxidative phenotype following dietary restriction. However, we observed only 375 the degree and the mechanism of fiber shift was dependent on the original fiber 376 type population e.g. T.long and T.lateral muscles displayed IIB-IIX-IIA in the 377 $Mstn^{-/-}$ however only showed a IIB \rightarrow IIX fiber type conversion in $Mstn^{+/+}$. 378 Moreover, ECU and FCU muscles showed a IIB \rightarrow IIX fiber shift in *Mstn^{-/-}* 379 however demonstrated a IIA \rightarrow I and IIB \rightarrow IIA shift respectively in *Mstn*^{+/+}. These 380 data indicate that myostatin deletion and dietary limitation interactions alter 381 muscle adaptation and plasticity. Muscle oxidative conversion was previously 382 reported with ageing, although there was trophic activity in type II fibers, 383 additionally there was a relative increase in type I fibers (Kallman et al., 1990). 384 Furthermore, a recent study in muscular dystrophy showed that muscle atrophy 385 was accompanied with a myofiber oxidative shift perhaps due to an increase in 386 the expression of PGC1 α (Rocchi et al., 2016). On the other hand, soleus 387 muscle immobilization, hind limb unloading and spinal cord transection induced 388 muscle atrophy plus type I into type II fibers transformation see review by (Cho 389 et al., 2016). However in our model the physical performance and muscle 390 function were preserved compared to previous models which resulted in 391 improved phenotype particularly in *Mstn^{-/-}*. It could be explain that muscle 392 adaptability is dependent on the extend of the external stimuli therefore the high 393 demand for protein supplement in the *Mstn^{-/-}* induced fiber atrophy and fiber 394 type transformation. This is far greater compared to *Mstn*^{+/+} such a change in 395 muscle metabolism energy saving mechanisms are required. 396

³⁹⁷ Our data discussed the impact of caloric restriction in combination with ³⁹⁸ *myostatin* deletion on five muscles from different anatomical locations. We ³⁹⁹ provide evidence that 60% of dietary limitation results in body weight and ⁴⁰⁰ muscle specific mass reduction with fast myofibers more prone to atrophy ⁴⁰¹ compared to slow myofibers. The effect of dietary restriction was more ⁴⁰² pronounced in some of the *Mstn^{-/-}* muscles to implicate the oxidative fibers ⁴⁰³ compared to *Mstn^{+/+}*mice. Furthermore, peripheral myofibers are more

susceptible to dietary induced reduction compared to deep fibers. We provide
evidence that dietary restriction alters the fast phenotype of the *Mstn^{-/-}* into the
slower form in a muscle dependent manner. These results allow the conclusion
that caloric reduction alters the muscle fiber composition and the oxidative
pattern to compensate muscle mass loss which might impact on muscle
function particularly in the hypermuscular *myostatin* null animals.

410 **Competing interest**

All the authors have declared no conflict of interest regarding the publication ofthis paper.

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529 Figure legends

530 Figure 1 Dietary restriction induces body weight and muscle mass 531 specific reduction.

a Average body weight (g) for normal diet (ND) and food restricted (FR) Mstn+/+ 532 and Mstn^{-/-} mice (N=4 for all groups). Significant increase in the Mstn^{-/-} 533 (P<0.001) compared to the $Mstn^{+/+}$ on normal diet. FR decreases the body 534 weight for *Mstn*^{+/+} and *Mstn*^{-/-} compared to genotype-matched ND mice. **b-f** 535 Average muscle weight (g) for b T.lateral, c T.long, d T.medial, e ECU, and f 536 FCU muscles. Specific muscle mass increases in *Mstn^{-/-}* ND compared to 537 *Mstn*^{+/+} ND. FR induces more muscle mass reduction in T.lateral and T.long 538 muscles of $Mstn^{-/-}$ (diet and genotype interaction P<001 and 0.0001) 539 respectively compared to genotype matched control. Mstn^{-/-} display more 540 muscle loss in T.medial, ECU and FCU muscles compared to Mstn+/+ following 541 FR. g-I Immunofluorescent images for T.long muscle of Mstn+/+ND against 542 A.474, BFF3 and A.4840 specific MHC antibodies show **g** type IIA⁺ (green), **h** 543 type IIB+ (red) and i type I+ fibers (green) compared to j, k and I the 544 corresponding negative control (NC) respectively. DAPI was used to visualize 545 the myonuclei. All values displayed as mean \pm SEM. * = P < 0.05, ** = P < 0.01, 546 *** = P< 0.001 and ****=P<0.0001). Scale bar in **g-l** =20μm. 547

548 Figure 2 Caloric restriction does not alter muscle fiber number.

a-e Average total myofiber number for a T.lateral, b T.long, c T.medial, d ECU, 549 and e FCU muscles of ND and FR Mstn+/+ and Mstn-/- mice. The muscles of 550 *Mstn^{-/-}* show increases in the myofiber number compared to *Mstn^{+/+}*. T.medial 551 muscle display no fiber number change in both genotypes. FR reveals no fiber 552 number loss compared to ND mice. f-i Reconstructive immunolabelled images 553 for FCU muscle of *Mstn*^{+/+} and *Mstn*^{-/-} ND and FR against MHCIIB (red), MHCIIA 554 (green), MHCIIX (unstained). Myostatin deletion increases the total CSA of the 555 muscle compare to wild type ($\mathbf{f} vs \mathbf{g}$). FR reduces the CSA of *Mstn*^{+/+} ($\mathbf{f} vs \mathbf{h}$) 556 and $Mstn^{-/2}$ (**q** vs. **i**). All values displayed as mean ± SEM. *=P< 0.05, **= P< 557 0.01 and ****=P<0.0001). 558

Figure 3 Fast fibers are more prone to caloric restriction-induced muscle mass loss.

a-e Average CSA measurements (µm²) for type IIB, IIX, IIA and type I fibers of 561 a T.lateral, b T.long, c T.medial, d ECU and e FCU muscles of Mstn^{+/+} and 562 Mstn^{-/-}ND and FR mice. FR causes significant reduction in the CSA of type IIB 563 fibers in T.lateral, T.medial and ECU muscles of Mstn^{-/-} compared to Mstn^{+/+}. 564 FR induces significant type IIX reduction in T.medial muscle of only Mstn^{-/-}, 565 T.lateral muscle of only *Mstn*^{+/+} and in T.long, ECU and FCU muscles for both 566 genotypes compared to ND. FR causes reduction in CSA of IIA fibers of T.long 567 and T.medial muscles of *Mstn^{-/-}* however increases the CSA of ECU and FCU 568 muscles. FR reveals no change in type IIA fibers for all Mstn+/+ muscles. FR 569 induces only significant reduction in type I fibers of T.long muscle but not 570 significant (NS) in T.medial muscle. f-i T.long muscle frequency distribution of 571 fiber CSA measurements (µm²) for **f** type IIB, **q** type IIX, **h** type IIA and **i** type I 572 fibers for of *Mstn*^{+/+} and *Mstn*^{-/-} ND and FR mice. FR induces fibers shift into 573 small size population (arrow marked) compared to ND for both genotypes. All 574 values displayed as mean± SEM. *=P< 0.05, ** = P< 0.01 and ****=P<0.0001. 575

576 Figure 4 Dietary restriction causes peripheral myofiber atrophy.

a-n Average CSA (μ m²) measurements for peripheral (PF) and central fibers 577 (CT) type IIB, IIX and IIA fibers for the muscles T.lateral **a**, **b**; T.long **c**, **d**, **e**; 578 T.medial f, g, h; ECU i, j, k and FCU I, m, n of ND and FR Mstn^{+/+} and Mstn^{-/-} 579 mice. For all muscle except T.medial, PF IIB fibers show CSA increases in Mstn⁻ 580 ^{/-} and *Mstn*^{+/+} ND compared to CT fibers. PF IIB fibers of T.lateral, ECU and 581 FCU muscles display smaller CSA compared to CT fibers following dietary 582 restriction (a, c, f, i and I). FR reduces the CSA of PF IIX fibers compared to 583 CT fibers for *Mstn^{-/-}* and *Mstn^{+/+}* (**b**, **d**, **g**, **j** and **m**). FR show larger CSA in the 584 CT IIA fibers of ECU and FCU muscles of *Mstn^{-/-}* (k and n). All values displayed 585 as mean± SEM. *=P< 0.05, **= P< 0.01, ***= P< 0.001 and ****=P<0.0001. 586

587 Figure 5 Caloric restriction induces a slow phenotype conversion in a 588 muscle-specific manner.

Average percentages of type IIB, IIX, IIA, and I fibers for T.lateral a, b, c; T.long d, e, f, g; T.medial h, i, j, k; ECU I, m, n, o; and FCU muscles p, q, r, s of ND and FR Mstn^{+/+} and Mstn^{-/-} mice. Mstn^{-/-} ND show increases in type IIB and IIX fibers compared to *Mstn*^{+/+}ND for all muscles. FR induces an increase in type IIX and type IIA fibers of T.lateral muscle in *Mstn^{-/-}* and only type IIX increases in *Mstn*^{+/+}. T.long muscle exhibits type IIB to IIA fiber shift and IIB to IIX fiber shift for *Mstn*^{+/+} and *Mstn*^{-/-} following FR respectively. FR causes an increase in type I fibers for T.medial muscle of *Mstn*^{+/+}. ECU muscles show IIB to IIX fiber type shift for *Mstn^{-/-}* compared to IIA to I fibers for *Mstn^{+/+}* following FR. FCU muscle displays IIB to IIA fiber shift for *Mstn*^{+/+} following FR. t-x Representative double immunofluorescent images of mid-belly section for T.long muscle in ND and FR Mstn^{+/+} and Mstn^{-/-} mice. FR induces myofiber transformation from glycolytic type IIB fibers (red) into oxidative type IIA. Type IIX fibers remained unstained (black). A consecutive muscle section with only secondary antibodies and DAPI was used as a negative control (NC). Scale bar =100 µm. All values displayed as mean± SEM.



Figure 1 Dietary restriction induces body weight and muscle mass
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623 Figure 2 Caloric restriction does not alter muscle fiber number.



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