The consequences of a high-calorie diet background before calorie restriction on skeletal muscles in a mouse model

Martin Maldonado¹, Jianying Chen¹, Yang Lujun², Huiqin Duan¹, Mazhar Ali Raja¹, Ting Qu¹, Tianhua Huang², Jiang Gu¹, Ying Zhong¹&

¹Chengdu Jinxin Institute of Reproductive Medicine and Genetics, Chengdu Jinjiang Hospital for Maternal and Child Health Care, Chengdu 610066, China
²Translational Medical Center, Second Affiliated Hospital of Shantou University Medical College, Shantou 515041, Guangdong, P.R. China

Correspondence to: Ying Zhong; email: yzhong08@126.com
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ABSTRACT

The beneficial effects of calorie restriction (CR) are numerous. However, there is no scientific evidence about how a high-calorie diet (HCD) background influences the mechanisms underlying CR on skeletal muscles in an experimental mouse model.

Herein we present empirical evidence showing significant interactions between HCD (4 months) and CR (3 months).

Pectoralis major and quadriceps femoris vastus medialis, in the experimental and control groups, displayed metabolic and physiologic heterogeneity and remarkable plasticity, according to the dietary interventions. HCD-CR not only altered genetic activation patterns of satellite SC markers but also boosted the expression of myogenic regulatory factors and key activators of mitochondrial biogenesis, which in turn were also associated with metabolic fiber transition.

Our data prompt us to theorize that the effects of CR may vary according to the physiologic, metabolic, and genetic peculiarities of the skeletal muscle described here and that INTM/IM lipid infiltration and tissue-specific fuel-energy status (demand/supply) both hold dependent-interacting roles with other key anti-aging mechanisms triggered by CR.

Systematic integration of an HCD with CR appears to bring potential benefits for skeletal muscle function and energy metabolism. However, at this stage of our research, an optimal balance between the two dietary conditions, where anti-aging effects can be accomplished, is under intensive investigation in combination with other tissues and organs at different levels of organization within the organ system.

INTRODUCTION

The idea that reductions in calorie intake without malnutrition could slow aging, decrease age-related chronic diseases, and lengthen lifespan was postulated nearly a century ago by McCay and colleagues when they observed that rodents subjected to calorie restriction (CR) lived far longer than their ad libitum-fed counterparts [1].

CR rapidly alters the physiology and function of muscle stem cells (SC) in such a way that even short-term CR in rats that begins late in life (~18 months of age) has significant positive impacts in strengthening myogenic activity [2].

Muscle satellite cells in the adult skeletal muscles are characterized by the expression of the paired-domain transcription factors Pax7 [3] and Pax3 [4], the
myogenic regulatory factors Myf5 and MyoD [5, 6], and the cluster of differentiation protein CD34 [7], among others. Of these, PAX7 is the canonical biomarker for satellite cells because it is particularly expressed in all quiescent and proliferating satellite cells across multiple species, including primates (humans and monkeys) and mice [3].

One of the beneficial effects of CR on satellite cell frequency in murine models is associated with metabolic rearranging that facilitates oxidative over glycolytic metabolism and is linked to pivotal modulators of mitochondrial mass and function, such as the NAD-dependent protein deacetylase SIRT1 [2].

SIRT1, the mammalian ortholog of Sir2 in yeast, is a nutrient-sensing and longevity factor that mediates the effects of dietary restriction in diverse species. In rodents, the expression of SIRT1 increases in response to prolonged CR in different tissues such as adipose, liver, kidney, and brain tissue [8, 9]. SIRT1 cross-talk and integrate signals with the energy-nutrient sensors AMPK, PGC-1α, and the mammalian target of Rapamycin (mTOR) [10–12] ensuring management and regulation of many cellular and metabolic processes, autophagy, and cell homeostasis.

CR has been extensively linked with mitochondria proliferation, biogenesis, dynamics, morphology, and reductions in oxidative stress [8, 13–15]. Oxidative stress induces telomere dysfunction, damage [16], and shortening of telomere to a critical length that can trigger aging and reduce life spans in mice and humans by mechanisms that involve the induction of persistent DNA damage response at chromosome ends [17]. However, CR has been suggested to synergize with telomerase expression, resulting in significant lifespan extension [18].

Moreover, adiponectin, a protein hormone encoded by the gene ADIPOQ in humans and mice and produced in adipose tissue, has been linked with CR. Indeed, CR appears not only to upregulate plasma adiponectin levels [19] but also appears to be associated with distinct metabolic states, suggesting that adipose tissue signaling is a suitable target for interventions to delay aging [20].

Herein, we attempt to correlate the above effects of CR, in two anatomically distinct skeletal muscles, with a high-calorie diet (HCD) background. We present evidence for the first time of interactions between CR and HCD and discuss different genetic mechanisms and pathways that may interact to modulate physiologic and metabolic changes elicited by the dietary interventions.

RESULTS

Development and characterization of the experimental animal model

The steps for the generation of animal models can be seen in Figure 1 and Materials and Methods. Mice fed ad libitum HCD showed significant increases in body weight (Figure 2A). Interestingly, the animals subjected to HCD (HC) presented reduced daily volumes of ingested food (Figure 2B) compared to those fed the standard 3.1 kcal/g diet ad libitum (SD). Nevertheless, due to food composition, the calorie intake of the HC was significantly higher than that of its control counterpart (Figure 2C).

At the end of the CR period, the SD-CR and HC-CR showed no bodyweight differences between them (Figure 2D).

To determine total lipid content in PM and QF-VM, we used the Oil Red O technique (Figure 3A, 3B). Moreover, we analyzed the gene and protein expression of the adipose-type cytoplasmic fatty acid-binding protein (Fabp4, Figure 3C), used to predict inter- (INTM) and intra-myocellular (IM) fat infiltration [21], and Perilipin (Figure 3D), the most abundant phosphoprotein on adipocyte lipid droplets [22].

The combined data showed that the HC-CR contains higher amounts of lipid content compared to that in the SD-CR, even though, as mentioned before, the average body weight of the two experimental groups was almost identical at the end of the CR period.

Due to the small size of the animals, we did not distinguish INTM and IM adiposity; a limitation that has been already observed in other available studies of IM adipogenesis in mice [23].

To characterize the metabolic status of the experimental and control groups, we determined the oxidative and glycolytic muscle fiber properties of PM and QF-VM with the SDH technique (Figure 4A, 4B).

Overall, the SD and SD-CR in PM displayed extensive areas of oxidative fibers, while the animals fed HCD (HC) showed fewer clusters of the oxidative type (vs SD-CR) and denser areas of intermediate and glycolytic fibers. Although not statistically significant (when the metabolic fiber types were analyzed individually), the HC-CR-PM presented a mild transition from glycolytic to oxidative activity seen by the increase in oxidative and intermediate fibers and a decrease in the glycolytic type.
In addition, QF-VM in the different groups appeared to have denser areas of glycolytic fibers than those seen in PM, and the HC presented significantly higher amounts of glycolytic fibers than the rest of the groups analyzed. However, when these animals with an HCD background (HC) were subjected to CR, we observed a clear transition from glycolytic to the oxidative phenotype.

Serum glucose levels were measured after 8 h of fasting conditions (Figure 4C). The data showed that mice subjected to CR (SD-CR and HC-CR) presented

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**Figure 1. Generation of the experimental animals.** CD-1 mice were fed *ad libitum* standard food (First stage) until they reached 12 months of age. Animals were separated into a group fed *ad libitum* standard food (SD) and *ad libitum* HCD (HC), for 17 weeks (Second stage). SD and HC animals were subdivided into control and experimental groups (Third stage). The experimental groups (SD-CR and HC-CR) were subjected to calorie restriction (CR) for 13 weeks before sacrifice.

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**Figure 2. Food and calorie consumption during the development of the experimental animal model.** The body weights of the average of the animal groups before calorie restriction (CR) (A). Food ingestion of the average of the animal groups expressed in grams before CR (B). Kcal consumption of the average of the animal groups before CR (C). The animal weights expressed in grams during the CR period (D). The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
improved blood glucose levels (compared to *ad libitum* fed mice), eight weeks after the beginning of the dietary restriction.

We also analyzed the insulin-like growth factor 1 (IGF-1) and its receptor IGF-1R (Figure 4D) because of its wide range of physiological and cellular functions, one of which is the regulation of glucose metabolism. We chose tissue-specific expression rather than circulating levels mainly because the technique allows the comparison of the two skeletal muscles and because intrinsic secretion of muscle IGF-1, rather than circulating plasma IGF-1, is a key element for switching on anabolic pathways [24].

Our results showed that CR decreased mRNA and protein levels of IGF-1 and IGF-1R (IGF-1R-SD-CR-QF-VM, not statistically significant) in the animals previously fed standard food (SD-CR), in both PM and QF-VM. Although CR decreased IGF-1 protein levels in HC-CR-PM, we did not observe significant changes in IGF-1 translational expression in the HC-CR-QF-VM. This was also accompanied by a marked increase in IGF-1/IGF-1R mRNA and IGF-1R protein expression. Furthermore, the HC animals presented IGF-1 and IGF-1R mRNA and protein upregulations in both skeletal muscles.

**Expression of satellite SC markers in mice subjected to CR**

We analyzed the satellite cell biomarkers PAX3, PAX7, PAX9, CD34, MyoD, and MyF5 from the QF-VM and the PM tissues in the different groups by qPCR (Figure 5A, 5B). After 13 weeks of CR, we observed 2 different and independent phenomena: First, the gene expression patterns differed between the muscles analyzed: Although the SD-CR presented an overall tendency

![Figure 3. Lipid content determination in PM and QF-VM of the experimental animal model.](image-url)

Representative images of lipid content stained with the oil red O technique (A). Scale bar: 100 μm. SD (n=6), SD-CR (n=8), HCD (n=9), and HCD-CR (n=12). The total area stained with Oil red O (B). Fabp4 (C) and Perilipin (D) were used to predict inter-myocellular (INTM) and intra-myocellular (IM) fat infiltration. Fabp4 and Perilipin were analyzed by qPCR (mRNA) and expressed as %, the SD value was set to 0, and the compared samples were normalized to this level. Positive % values represent upregulation. Negative values represent downregulation. Protein expression of Fabp4 and Perilipin was obtained by Western Blot analysis and quantified with Image Lab 6.1 software. SD expression was set to 1 and the relative protein levels were normalized as a ratio of GAPDH expression. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
toward overexpression, only PAX7 (97%, ± 26) and PAX9 (89%, ± 20) were significantly upregulated in PM (compared to those in the SD), whereas in QF-VM all the satellite transcripts were significantly increased. Furthermore, the animals in the HC-CR-PM presented higher expression levels in PAX3 (33%, ± 7), PAX7 (176%, ± 27), CD34 (56%, ± 12), MyoD (220%, ± 24), and Myf5 (121%, ± 12), whereas in the HC-CR-QF-VM only PAX9 (21%, ± 4), MyoD (70%, ± 8) and Myf5 (62%, ± 13) were upregulated.

Second, the gene expression patterns differed between the animals exposed to CR with different alimentary backgrounds (SD-CR and HC-CR), as PAX9-PM was strongly upregulated (89%, ± 20) in the SD-CR (compared to those in the SD) while significantly downregulated to -29%, ± 10 in the HC-CR (compared to those in the SD). Other examples are PAX3-QF-VM (SD-CR upregulation to 52%, ± 14; HC-CR downregulation to -60%, ± 21) and Myf5-PM (SD-CR downregulation to -24%, ± 6; HC-CR upregulation to 121%, ± 12).

The gene expression of satellite and other markers, expressed as a percentage (%) or fold change values can be found in the Supplementary Table 4.

Mitochondrial DNA copy numbers and mitochondrial biogenesis-related activators in mice subjected to CR

The mitochondrial DNA copy numbers in the groups subjected to CR (the SD-CR and HC-CR) were increased in PM tissue by 28%, ± 3.5, and 66% ± 6.4 respectively (vs SD-PM), and in QF-VM tissue by 48.1 ± 4 and 51.2% ± 4.9 respectively (vs SD-QF-VM; Figure 6A). Our results appear to agree with scientific data supporting the idea that CR induces mitochondrial proliferation in rodents [16, 8].

![Figure 4. Metabolic characterization of PM and QF-VM in the experimental animal model.](image-url)

Representative images of SDH staining in muscle fibers of PM and QF-VM in the different groups (A). Scale bar: 100 μm. SD (n=6), SD-CR (n=8), HCD (n=9), and HCD-CR (n=12). The total area stained with Oil red O (B). Dark stained fibers were classified as oxidative (Ox); intermediate stained fibers were classified as oxidative-glycolytic (Ox-Gl); and very lightly or non-stained fibers were classified as glycolytic (Gl). Comparison of the blood glucose levels of mice during week 9 to 13 of the dietary restriction (C). IGF-1 and IGF-1R were analyzed by qPCR and Western blot analysis (D). mRNA was expressed as fold, the SD value was set to 0, and the compared samples were normalized to this level. Positive values represent upregulation. Negative values represent downregulation. Protein expression of IGF-1 and IGF-1R was obtained by Western blot analysis and quantified with Image Lab 6.1 software. SD expression was set to 1 and the relative protein levels were normalized as a ratio of GAPDH expression. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
Figure 5. Gene expression of satellite markers. cDNA from PM and QF-VM was obtained with a Takara RR047Q kit. qPCR was performed with 100 ng of target DNA to measure PAX3, PAX7, PAX9, CD34, MyoD, Myf5, and 18s rRNA (reference gene) expression (A). For each gene expressed as percentage (%), the SD value was set to 0, and the compared samples were normalized to this level. Positive values represent upregulation. Negative values represent downregulation. Each marker was analyzed with SYBR Green fluorescence detection, and the transcript levels were normalized to those of the endogenous control 18s rRNA. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD.

A hierarchical clustering illustration for the up/down-regulation of genes analyzed from RT-PCR array data. The right bar depicts the colors for upregulation and downregulation expressed as a % (B).
Then, we assayed the mRNA and protein expression of eNOS (Figure 6B, 6C), which has important roles in the regulation of mitochondrial function and induction of mitochondrial biogenesis [25], and PGC-1α (Figure 6D, 6E), a transcriptional co-activator, and also a master regulator of mitochondrial biogenesis [26, 27].

Interestingly, eNOS and PGC-1α in the HC-CR showed similar increased expression patterns (except for eNOS-QF-VM mRNA) compared to those found in the SD-CR, while the SD-CR-PM presented significant downregulations (vs SD).

We also measured the transcript levels of TOMM20, TOMM40, and TIMM9 (Figure 6F), which are responsible for encoding mitochondrial importing proteins, and NDUFS3 and ATP5C1 (Figure 6G), which encode electron transport chain associated enzyme components, because they reflect function and biogenesis rather than abundance [28].

Figure 6. Mitochondrial DNA copy numbers and the expression of mitochondrial biogenesis-related activators in mice subjected to CR. Mitochondrial DNA copy number (A) was analyzed with the Detroit-Mouse-mt-DNA analysis kit (Supplementary Material, 1). For each group, the value was expressed as a percentage (%); the SD value was set to 0, and the rest of the groups were normalized to this level for comparison. eNOS (B, C) and PGC-1α (D, E) were analyzed at the transcriptional and translational levels. TOMM20, TOMM40, TIMM9 (F), and NDUFS3 and ATP5C1 (G), were analyzed at the transcriptional level. For the qPCR assay, each primer was analyzed with SYBR Green fluorescence detection and the transcript levels, expressed as a %, were normalized to those of the endogenous control 18s rRNA. Protein expressions were obtained by Western Blot analysis and quantified with Image Lab 6.1 software. SD expression was set as 1 and the relative protein levels were normalized as a ratio of GAPDH expression. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
Similar to the previous patterns found in eNOS and PGC-1α, TOMM20, TOMM40, TIMM9, NDUFS3, and ATP5C1 were significantly upregulated in the HC-CR (except for TOMM20-QF-VM, and NDUFS3-QF-VM; vs SD-CR).

The SD-CR-QF-VM, unlike SD-CR-PM, presented overexpression of TOMM20, TIMM9, and ATP5C1 in (vs SD), while eNOS and PGC-1α remained unchanged (vs SD).

These findings suggest that although elevated amounts of mitochondria DNA content were found in both CR-treated groups, only the HC-CR and may present improved mitochondrial biogenesis and function.

**Telomerase activity was increased in animals subjected to CR**

The groups subjected to CR (SD-CR and HC-CR) expressed significantly higher telomerase activity in both PM and QF-VM tissue compared to the CGs (Figure 7), evidence that suggests a possible interaction between these two major anti-aging mechanisms. Moreover, the PM tissue appeared to have lower telomerase activity than that found in QF-VM.

**CR and the AMPK-SIRT1-mTOR network**

AMPKs and SIRT1, as well as mTOR (a central controller of cell growth and proliferation), are involved in the regulation of many cellular and metabolic processes. The cross-talk among these three enzymes occurs at several levels and the action of one may be modulated by the actions of the other(s) in some cases [13].

CR is well known to increase AMPKs and SIRT1 expression in different tissues and species [29, 9], so we analyzed the mRNA and protein levels of AMPK, T183-T172, and SIRT1 endogenous levels (Figure 8A–8E).

![Figure 7. Telomerase activity.](image-url) Telomerase activity was assayed with a TRAPEze RT Telomerase Detection Kit (Millipore) for fluorometric detection and real-time quantification in pectoralis major (PM) and quadriceps femoris vastus medialis (QF-VM). The telomerase values are arbitrary units relative to the TSR8 amplification, according to the assay kit protocol (Supplementary Material 2). The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
Although we presented mRNA and protein data of the different isoforms of AMPK, we focused on T183-T172 as AMPK becomes activated when phosphorylation takes place at T183 (AMPK α1, phosphorylated at threonine 183) and T172 (AMPK α2, phosphorylated threonine 172).

Our data showed higher AMPK expressions and AMPK activity (T183-T172) in the HC-CR than those found in the SD-CR, in PM (Figure 8A, 8B), and QF-VM (Figure 8C, 8D). Furthermore, SIRT1, which followed similar expression patterns in the two muscles analyzed, was slightly increased in the animals subjected to CR, but

Figure 8. CR and the AMPK-SIRT1-mTOR network. AMPKs were analyzed by qPCR (mRNA levels) and Western Blot (protein levels) in pectoralis major (PM) (A, B) and quadriceps femoris vastus medialis (QF-VM) (C, D). SIRT1 was analyzed in PM and QF-VM (E) with the Mouse NAD-Dependent Deacetylase Sirtuin-1 (SIRT1) ELISA Kit (CUSABIO) for quantitative determination, according to the assay kit protocol (Supplementary Material 3). Values are expressed in ng/ml. mTOR mRNA in PM and QF-VM (F). Immunoblot results and protein expression of mTOR (G) and mTOR-S2448 (H) in PM and QF-VM. For the qPCR assay, each primer was analyzed with SYBR Green fluorescence detection and the transcript levels, expressed as a %, were normalized to those of the endogenous control 18s rRNA. Protein expressions were obtained by Western Blot analysis and quantified with Image Lab 6.1 software. SD expression was set as 1 and the relative protein levels were normalized as a ratio of GAPDH expression. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
AMPK and SIRT1 are known to negatively regulate mTOR [30, 31], thus we checked mTOR (transcript, Figure 8F; protein, Figure 8G) expression, and the protein expression of mTOR phosphorylated at serine 2448 (mTOR-s2448; Figure 8H), as mTOR complex 1 (mTORC1–regulatory associated protein of mTOR [Raptor] positive, primarily phosphorylated at s2448) is a key regulator of skeletal muscle growth and cell proliferation through activation of mRNA translation/protein synthesis, and regulation of autophagy [32].

mTOR mRNA levels were significantly downregulated in PM (SD-CR -62% ± 16; HC -66.5 ± 13; HC-CR -103% ± 22) and in the QF-VM, only the HC-CR was downregulated (-34% ± 8). At the protein level, mTOR-HC-CR was significantly lowered compared to that in the SD-CR in PM (* < 0.05) and SD (PM, **<0.01; QF-VM, *<0.05).

The analysis of the mTORC1 complex showed activation in HC-PM and QF-VM (PM, **<0.01; QF-VM, **<0.01). On the other hand, the groups exposed to CR presented lower mTOR activity (PM-SD-CR, *<0.05; PM-HC-CR, **<0.01; QF-VM-HC-CR, *<0.05; vs SD). However, QF-VM-SD-CR displayed no statistical differences, vs SD.

**Adiponectin and adiponectin-receptors in mice subjected to CR**

The adipocyte-specific hormone and its receptors AdipoR1 and T-cadherin were examined in PM (Figure 9A, 9B) and QF-VM tissues (Figure 9C, 9D).

Adiponectin mRNA expression was drastically increased not only in the animals exposed to CR (except for the SD-CR-PM, -2.23- ± 0.3-fold) but also in the HC, in both muscle tissues. At the translational level, adiponectin also presented higher expression in the HC-CR compared to that in the SD-CR (Figure 9B, 9D), however, the differences were not significant in QF-VM.

Moreover, gene profiling of adiponectin in the HC-CR revealed positive correlations with the overexpression of the adiponectin receptors AdipoR1 (PM: r=0.77; QF-VM: r=0.66,) and T-cadherin (PM: r=0.79; QF-VM: r=0.72) in both tissues. Furthermore, the under-expression of adiponectin found in the SD-CR-PM was also strongly correlated with AdipoR1 (r=0.76) and T-cadherin (r=0.73) (Figure 9A).

**DISCUSSION**

In this study, we have examined the effects of an HCD background in two skeletal muscles from distant anatomical locations, in mice subjected to CR. In terms of the physiologic, metabolic, and phenotypic characteristics, the skeletal muscles in the experimental and control groups presented not only differences in the amounts of lipid infiltration but also exhibited discrepancies in terms of metabolic fiber properties, as well as metabolic fiber transition, according to the dietary interventions. PM enzymatic activity was predominantly oxidative in the SD whereas QF-VM presented more extensive areas of the glycolytic type. HCD induced fiber transition to the glycolytic phenotype in both muscles (HC groups), which was consistent with other reports showing inverse relationships between adiposity and the oxidative/type 1 muscle fiber [33–36]. Moreover, HC-CR increased the number of oxidative/intermediate fibers in PM and QF-VM. These results were associated with the protein and mRNA-overexpression of the transcriptional co-activator PGC-1α (PM-HC-CR, **<0.01; QF-VM-HC-CR, **<0.01) that has been reported to increase the proportion of red/oxidative type I fibers [37].

Numerous reasons such as nutrition, hormone levels, exercise, and other external environmental insults have been reported to affect muscle fiber type transition [38, 39], and the CR-mimetic Resveratrol has also been reported to induce myofiber type transition from glycolytic to oxidative in skeletal muscles [40–42].

Furthermore (except for HC-CR-QF-VM), CR decreased IGF-1 and IGF-1R protein expressions, consistent with other previous reports in rodents [43–45]. Nevertheless, IGF-1 translational levels in the HC-CR-QF-VM showed no differences than those found in the SD and displayed significantly higher expressions of IGF-1R. These results could be explained at least in part, by the effect of IGF-1 on mitochondrial mass via increased transcriptional activities of key factors involved in mitochondrial biogenesis like PGC-1α [46].

Prior studies in the rodent skeletal muscles [47, 2] suggested that short-term and lifelong CR increase and maintain the satellite cell number and function which normally declines with age, but whether short-term CR in combination with HCD in otherwise healthy aged animals reshapes the expression of specific markers of satellite SCs have not yet been addressed.

Due to the experimental design, in this work satellite SCs were not isolated, and the mRNA data represent the expression of the whole muscle tissue. Nevertheless, the two observed independent events already stated in the
results section (contrasting gene expression patterns differed between the muscles analyzed; different gene expression patterns between the animals exposed to CR with different alimentary backgrounds) may reflect the dynamics of the gene expression in the satellite SC pools. That being said, isolation and characterization of satellite SCs under the same experimental conditions would be necessary to confirm this hypothesis. Concerning the analysis of mitochondrial function, biogenesis, and number, CR promoted mitochondrial abundance by increasing the amounts of mtDNA [48] in both CR-treated groups in PM and QF-VM, regardless of the metabolic fiber type composition/activity, as the slow-twitch oxidative fibers (predominantly found in PM) contain many more mitochondria than the glycolytic type (predominantly found in QF-VM). The aerobic metabolism, which consumes O2, occurs in the mitochondria, thus the levels of oxidative enzymes in the different muscle fibers reflect the relative abundance of mitochondria, as determined by electron microscopy [49].

Figure 9. Expression of Adiponectin and Adiponectin-receptors. The mRNA levels of adiponectin, AdipoR1, and T-cadherin in pectoralis major (PM) (A). Immunoblot results and protein expression of adiponectin in PM (B). mRNA levels of adiponectin, AdipoR1, and T-cadherin in quadriceps femoris vastus medialis (QF-VM) (C). Immunoblot results and protein expression of adiponectin in QF-VM (D). Adiponectin, AdipoR1, T-cadherin, and 18s rRNA (reference gene) transcripts were analyzed by qPCR. Significant differences between groups are expressed as fold change values. For each gene expressed as a fold change, the SD value was set to 0, and the compared samples were normalized to this level. Positive values represent upregulation. Negative values represent downregulation. The fold changes are defined directly in terms of ratios. Each marker was analyzed with SYBR Green fluorescence detection, and the transcript levels were normalized to those of the endogenous control 18s rRNA. Protein expression of adiponectin in PM and QF-VM was obtained by Western Blot analysis and quantified with Image Lab 6.1 software. SD expression was set to 1 and the relative protein levels were normalized as a ratio of GAPDH expression. The data are the mean ± s.d. *P < 0.05; **P < 0.01 vs SD, unless otherwise specified.
Animals fed HCD previous to CR (HC-CR-PM and QF-VM) displayed increased expressions of the key regulators of mitochondrial biogenesis eNOS, PGC-1α, and AMPK (discussed in the AMPK-SIRT1-mTOR network).

On the other hand, mice fed standard food previous to CR (SD-CR) showed an overall reduction in the above-mentioned regulators of mitochondrial biogenesis (vs HC-CR), and part of these inconsistencies could be attributed to the processes involved in fiber transition to the oxidative phenotype, seen in the HC-CR-PM and -QF-VM (marked plasticity in the glycolytic/intermediate QF-VM); phenotypic changes that we did not observe in the animals treated with CR alone (SD-CR).

Moreover, differences between muscles, in the SD-CR, were also evident. The SD-CR-PM (vs SD and HC-CR) displayed reduced expressions of eNOS (mRNA, Figure 6B; protein levels, Figure 6C) and PGC-1α (mRNA, Figure 6D; protein levels, Figure 6E), TOMM20, TOMM40, TIMM9, NDUFS3, and ATP5C1 mRNA (Figure 6F, 6G). These peculiarities might be attributed at last in part, to the contrasting metabolic fiber activity (PM vs QF-VM), determined by the SDH technique (PM: Oxidative/intermediate/glycolytic; QF-VM: Glycolytic/intermediate/oxidative).

The CR-treated groups also presented mRNA upregulations of the myogenic regulatory factors (MRFs) MyoD and Myf5, except for the SC-CR-PM. Although little is known about Myf5, MyoD directly binds to numerous metabolic genes, including those associated with mitochondrial biogenesis, and regulates the oxidative metabolism cooperatively with alternative NF-kB [50].

Kraft, C. S. et al. (2006) have shown that increased signals of myogenesis (MyoD, Myf5) are associated with a metabolic transition from glycolysis to oxidative phosphorylation, with a consequent increase in mitochondrial biogenesis [51]. These assumptions are not only in line with our data in the HC-CR groups but remarkably with the MFRs downregulations (vs HC-CR) and reduced expression of markers of mitochondrial biogenesis observed in the SD-CR-PM, that follow the same notion.

Another report, also in agreement with our data on MRFs expressions and AMPK observed in the HC-CRs, postulate that AMPK-α1 is the predominant AMP isoform expressed in satellite cells, and AMPK-α1 deficiency in satellite cells impairs their activation and myogenic differentiation during muscle regeneration [52].

Then we analyzed the AMPK-SIRT1-mTOR network, which is implicated in the regulation of many cellular and metabolic processes, autophagy, and cell homeostasis.

Cellular stresses due to low glucose and nutrient deprivation cause a rise in the AMP/ATP ratio, thus AMPK becomes activated (T183-T172 results). AMPK activation suppresses anabolic pathways like gluconeogenesis, glycogen, fatty acid, triglyceride, cholesterol, and protein synthesis (mTOR-p70S6K-E2 pathway), and triggers catabolic pathways such as glycolysis, glucose uptake, and fatty acid oxidation to restore appropriate energy levels in the cell.

We observed in this work significantly increased transcript and protein expressions in the different AMPK subunits, and particularly in AMPK activity (T183-T172) in the HC-CR, in both PM and QF-VM (vs SD and SD-CR); results that were also associated with metabolic fiber transition towards the oxidative phenotype, and elevated mRNA/protein expressions of PGC-1α. These findings follow the notion that activation of AMPK leads to mitochondrial biogenesis by regulating PGC-1α which in turn improves the ATP synthesis capacity of the cell and promotes gene transcription in mitochondria, in skeletal muscles in response to chronic energy deprivation [53]. Moreover, Jager et al. [11] have also demonstrated that AMPK’s energy control function is exerted by activating mitochondrial biogenesis. This effect is realized by direct phosphorylation of PPAR-γ coactivator 1α (PGC-1α), the master regulator of this process, and while AMPK promotes the synthesis of new mitochondria, it also influences their turnover by activating their degradation through the autophagic process (mitophagy) [10].

Similar but not the same as our experimental design, Gutiérrez-Casado, et al. (2019) have shown that in skeletal muscle from CR mice, dietary fat influences mitochondrial mass and ultrastructure and may play a role in processes such as autophagy and mitophagy and mitochondrial dynamics during aging [54].

Although we did not analyze mitophagy, the above-reported data may suggest that our animals previously fed HCD before CR (HC-CR), due to increased AMPK activity, may have elevated mitophagy activity (vs SD and SD-CR), and further studies in the levels of autophagy and senescence would add valuable insights in the understanding of how HCD background affects CR.

An important question that remains to be acknowledged here is why animals fed standard food, subjected to CR
(SD-CR), presented lower AMPK activity (vs SD and HC-CR) in the two skeletal muscles?.

To answer this, we must take into consideration that the mechanisms of activation and the dynamics of AMPK are complex phenomena, not yet fully understood, and although it is recognized that phosphorylation at Threonine 172 (AMPK α2) - Threonine 183 (AMPK α1) is involved in AMPK activation, it may also be regulated by several other upstream kinase and pathways including CaMKK, Sirt1, LKB, CD36, ROS, allosteric regulation by AMP, AMP/ATP ratio, and the folliculin interacting protein 1 (Fnip1), among others [55–58]. Thus, we cannot rule out that distinct upstream regulators, stress signaling pathways, etc., are underpinning the decrease in AMPK activity in the SD-CR.

Interestingly, AMPK activity has been implicated in lipid accumulation in skeletal muscle cells through FTO-dependent demethylation of N6-methyladenosine [59]. Another report proposed that AMPK regulates muscle lipid storage, and reducing AMPK kinase activity decreases muscle perilipin-3 expression (which is consistent with our data) and intramyocellular triglyceride content [60].

Based upon the relation between AMPK with lipid accumulation, our data in the SD-CR (in line with the HC and HC-CR data) appear to be linked at least in part, with energy status (demand/supply) and contents of INTM/IM lipid infiltration.

Moreover, changes in the levels of various hormones have been revealed to activate or inhibit AMPK [61–64], suggesting that its regulation and physiological relevance are far more complex than what we have covered in this manuscript.

SIRT1, which values (ng/ml) were slightly increased, but not statistically significant, in the CR-treated groups could be rationalized as prolonged exposures to CR in rodents have shown to have stronger effects on SIRT1 in several different tissues [8]. Nevertheless, we emphasize here that the regulatory properties of SIRT1 may, nonetheless, be remarkably more intricate, as SIRT1 activity relies also on intracellular levels of nicotinamide [65], post-translational modifications [66], and/or interactions with several other protein mediators [67, 68].

mTORC activity, assayed in this work by the expression of mTOR-S2448, was reduced in the CR-treated groups (SD-CR-QF-VM was not statistically significant; vs SD); results that were expected as it is well documented that CR negatively regulates the mammalian target of rapamycin. However, considering that the mTOR pathways respond to various signals, including the nutrients (glucose and branched-chain amino acids), energy (ATP/AMP ratio), hypoxia, and growth factors (IGF-1/AKT pathway), etc., and that our data on T183-T172 (AMPK activity) showed mRNA and protein downregulations in the animals feed standard food previous to CR (SD-CR), another question mark related to which upstream mechanisms, other than AMPK activation, are implicated in mTOR inhibition, warrants deeper investigation.

It is now widely recognized that the adipose tissue is an essential and highly active metabolic-endocrine organ that secretes peptide hormones including leptin, cytokines (especially TNFα), adipin and acylation-stimulating protein (ASP), angiotensinogen, plasminogen activator inhibitor-1 (PAI-1), adiponectin, resistin, etc., as well as steroids hormones. [69–71].

In this work we choose to investigate adiponectin, not only because of its numerous anti-aging properties but because the characterization of the skeletal muscle tissues after HCD-CR demonstrated contrasting amounts of lipid infiltration in the groups.

Previous reports have inversely correlated adiponectin circulating levels with obesity [72, 73], and positively associated it with CR [74, 75]. Curiously, Miller et al. (2017), although they found a significant inverse relationship between HMW adiponectin and fat mass in mice of 10 or 20 months of age, no changes at 30 months of age were evidenced, indicating that in lean animals, the levels of HMW adiponectin, but not total adiponectin, are related to adiposity and that this relationship is sensitive to age. [20]. Moreover, Amany A Saleh, et al. (2020) found that adiponectin serum levels increase mainly due to secretion by the central adipose tissue but it does not reflect the tissue-specific expression of adiponectin and receptors, and this may vary in the different organs [76].

Our data showed that CR did not increase tissue-specific adiponectin protein expressions in the animals previously fed standard food (SD-CR-PM and -QF-VM), but it significantly upregulated adiponectin in the animals previously fed HCD (HC-CR-PM and -QF-VM). Because adiponectin is mainly produced by the adipocytes, we would expect lower protein expression of tissue-specific adiponectin in the SD-CR groups (SD-CR displayed markedly reduced lipid content). However, no changes were observed, compared to those found in the SD. This might be attributed to the fact that mechanistically, adiponectin, as a protective response against metabolic insults, can also be produced by the myocytes, in vitro and in vivo [77–80].
Surprisingly, the HC-PM displayed elevated translational expression of adiponectin (vs SD). Of note, HC-PM showed elevated amounts of fat infiltration and marked plasticity (fiber transition to the glycolytic phenotype), which was consistent with a report demonstrating that intramyocellular adiponectin levels are primarily associated with type IIA/D fibers containing elevated intramyocellular lipid accumulation [78].

It is evident that the magnitude of the subject embraced in this work goes far beyond the scope of the collected data. Thus, we would like to acknowledge this matter as one of the limitations that we have encountered so far. Here we merely present evidence of how HCD alters genetic patterns in the skeletal muscles of animals subjected to CR with the same BMI and how HCD in combination with CR affects specific key anti-aging mechanisms boosted by CR alone. That being said, further mechanistic studies on this topic are urgently required.

Also, it is important to mention that protein post-transcriptionally mRNA regulation or post-translational modifications (methylation, acetylation), differential protein degradation (proteasome-mediated or autophagy-mediated protein breakdown), pre-existing lysine acyl modifications, including succinylation, malonylation, glutarylation, crotonylation, beta-hydroxyisobutyrylation, 3-hydroxy 3-methylglutaryl, and fatty acylation, etc. [81, 82], influence protein structure and function, thus the transcriptional and translational expressions, in some of the genes analyzed in this manuscript, would not be always in line with each other.

Another aspect of our work that needs to be considered is that the data reflects the result of specific experimental conditions like animal species, age, tissue-specific expression, and characteristics of the dietary interventions, among others, so whether the results could be or could be not replicated, relies on the particularities of the experimental setup.

**CONCLUSIONS**

Herein we present empirical evidence showing significant interactions between HCD and CR. The skeletal muscles analyzed, in the experimental and control groups, displayed metabolic and physiologic heterogeneity and remarkable plasticity, according to the dietary interventions.

HCD in combination with CR not only altered genetic activation patterns of satellite SC markers but also boosted the expression of MRFs and key regulators of mitochondrial biogenesis, which in turn were also associated with metabolic fiber transition.

Our data prompt us to theorize that the effects of CR may vary according to the physiologic, metabolic, and genetic peculiarities of the skeletal muscle described here and that INTM/IM lipid infiltration and tissue-specific fuel-energy status (demand/supply) both hold dependent-interacting roles with other key anti-aging mechanisms triggered by CR.

Finally, in this manuscript, we expose an unexplored area from which controversy may arise regarding the effects of CR on skeletal muscles in higher organisms that have been subject to caloric overload.

Systematic integration of an HCD with CR appears to bring potential benefits for skeletal muscle function and energy metabolism. However, at this stage of our research, an optimal balance between the two dietary conditions, where anti-aging effects can be accomplished, is under intensive investigation in combination with other tissues with distinct metabolic demands, and organ structures at different levels of organization within the whole-body system.

**MATERIALS AND METHODS**

**Food design**

The animal food was developed by Beijing Keao Xieli Feed Co., Ltd (Beijing Chaoyang district, Yangshan road, number 4). The standard food (3.1 kcal/100 g) composition can be seen in the Supplementary Table 1. A hypercaloric food rich in fatty acids and carbohydrates (HCD; 5.5 kcal/100 g) was designed to induce obesity in the animal model (Supplementary Table 2).

**Generation of the animal model and dietary interventions**

Female CD-1 (ICR) mice were maintained in a specific pathogen-free animal facility in individual and ventilated cages and were housed at 23°C under a 12-hour dark/light cycle. Water and food were given ad libitum prior to the dietary interventions. This study was approved by the Institutional Review Board of the Chengdu Jinjiang Maternity and Child Health Hospital and all methods were performed under the relevant guidelines and regulations.

For the generation of the animal models, a group (Figure 1) initially composed by ~ 12-month-old mice (n=80) fed ad libitum 3.1 kcal/100 g standard food (First stage) was divided (second stage) into a group fed...
ad libitum standard food (n=30) and another group of animals switched to a 5.5 kcal/100 g ad libitum HCD for a period of 17 weeks (n=50). Then, mice previously fed standard food that continued with the same diet (n=10) became the standard diet control group (SD). Animals previously fed standard food, subjected to CR for a period of 13 weeks with a diet consisting of a 60% of the ad libitum 3.1 kcal/100 g calorie intake (n=19), became the standard diet-calorie restriction group (SD-CR); Animals fed HCD diet for a period of 17 weeks that continued with the same nourishment (n=14) became the high-calorie control group (HC); Animals fed HCD for a period of 17 weeks and then switched to CR (3.1 kcal/100 g food equivalent to 60% of the 5.5 kcal/100 g ad libitum calorie intake) for a period of 13 weeks (n=33) became the high calorie-calorie restriction group (HC-CR). The daily amount of food of the CR-restricted groups was adjusted according to decreasing body weight.

Mice were monitored daily by laboratory members and by animal health technicians. Before the experimental endpoint, the mice experienced minimal pain and stress during routine handling, bodyweight determination, and blood collection from the tail vein for measurement of blood glucose levels.

No ill or deceased mice were observed before the experimental endpoint (when the mice were 19 to 20 months old). Animals were euthanized, under the same metabolic conditions, by the cervical dislocation technique.

**Tissue collection**

Muscle tissues were collected from the quadriceps femoris vastus medialis (QF-VM) of the right lower limb and the right pectoralis major (PM) of the chest. The samples were flash-frozen in liquid nitrogen and stored until the execution of the experiments.

**Oil red O staining assay for frozen sections**

PM and QF-VM frozen tissues from the SD (n=6), SD-CR (n=8), HCD (n=9), and HCD-CR (n=12) were cut into 8μm thick sections with a cryostat (Leica CM1860 UV cryomicrotome, Leica Biosystems) and mounted on glass slides, then air-dried for 30-60min at room temperature and fixed in ice-cold 10% formalin for 5-10 minutes. The step was repeated twice and the slides were immediately rinsed with distilled water. After drying for several minutes, the slides were placed in absolute propylene glycol for 2-5 minutes and stained in pre-warmed oil red O solution for 8-10 minutes. Samples were treated with 85% propylene glycol solution for 2-5 minutes and rinsed with 2 changes of distilled water. Slides were finally stained in Mayer’s hematoxylin for 30 minutes, placed in distilled water, and then mounted with an aqueous mounting medium for microscope visualization. Area distribution of frozen section tissues was calculated with the ImageJ software.

**Succinic dehydrogenase assay for the determination of metabolic muscle fiber type**

Metabolic fiber type (oxidative and glycolytic) in PM and QF-VM from the SD (n=6), SD-CR (n=8), HCD (n=9), and HCD-CR (n=12) was determined with the succinate dehydrogenase (SDH) staining solution (Tetra-salt method; Batch # G2000) kit (Beijing Solarbio Science and Technology Co., Ltd.). Six or more transverse sections (12 mm thick) were obtained with a cryostat (Leica CM1860 UV cryomicrotome, Leica Biosystems). Briefly, the flash-frozen sections were immersed in NBT incubation solution for 20 min at 37 C. Samples were rinsed with distilled water, and coverslips were mounted onto glass slides with a natural balsam for microscopic visualization. Myofibers were processed with a digital slide scanner (MoticEasyScan, Motic Microscopy) linked to a Motic digital slide assistant image capture software. Areas were randomly selected for the determination of metabolic fiber type.

**Quantitative real-time polymerase chain reaction (qPCR) assay**

Tissues preserved in liquid nitrogen were homogenized, and the RNA was extracted with Takara RNAiso PLUS Total RNA Extraction Reagent (Takara Bio, Inc.). DNA was extracted with a TIANamp Genomic DNA Kit (TIANGEN). The total nucleic acid concentration and optical density (OD) were assayed by UV spectrophotometry. cDNA was obtained with a Takara kit (RR047Q). qPCR was performed with 100 ng of target DNA; the expression of the pair box gene (PAX) 3, PAX7, PAX9, cluster of differentiation 34 (CD34), myoblast determination protein 1 (MyoD), myogenic factor 5 (Myf5), adipose-type cytoplasmic fatty acid binding protein (Fabp4), perilipin, Insulin growth factor 1 (IGF1), IGF-1 receptor (IGF-1R), nitric oxide synthase (eNOS), peroxisome proliferator-activated receptor-gamma coactivator-1alpha (PGC-1α), mitochondrial import receptor subunit TOM20 homolog (TOMM20), translocase of outer mitochondrial membrane 40 homolog (TOMM40), mitochondrial import inner membrane translocase subunit Tim9 (TIMM9), NADH dehydrogenase [ubiquinone] iron-sulfur protein 3, mitochondrial (NDUFS3), ATP synthase, H+ transporting, mitochondrial F1 complex,
gamma polypeptide 1 (ATP5C1), AMP-activated protein kinase (AMPK) Alpha 1-2, Beta 1-2, Gamma 1-2, AMPK phosphorylated at Threonine 183 and 172 (T183-T172), mammalian target of rapamycin (mTOR), and 18s ribosomal RNA (18s rRNA) or GAPDH (the two reference genes) was evaluated through relative quantification with SYBR Premix Ex Taq™ (Takara RR420Q) using an ABI7500 instrument (Applied Biosystems, Foster City, CA). The samples were run in triplicate and the list of primers can be seen in Supplementary Table 3.

**Western blot analysis**

The tissues were homogenated with 5 volumes of Radioimmunoprecipitation Assay (RIPA) buffer (Solarbio® life sciences), and the supernatants were fractionated by SDS-PAGE. The proteins were quantified with BCA protein quantification kit (Solarbio® life sciences), transferred to PVDF fiber membranes (Bio-RAD), and blocked with 5 % blocking solution for Western blot (Roche). The membranes were then exposed to Anti-Fabp4 antibody [EPR3579] ab92501(Abcam; dilution: 1/2000); Anti-Perilipin-1 [EPR3753(2)] ab172907 (Abcam; dilution: 1/2000); IGF1 Rabbit mAb #A0830 (Abclonal; dilution: 1/500); Anti-IGF1 Receptor antibody [EPR19322] ab182408 (Abcam; dilution: 1/1000); Anti-eNOS antibody [EPR19296] ab199956 (Abcam; dilution: 1/1000); PGC1 alpha Rabbit pab, #A1971 (ABclonal; dilution: 1/1000); Anti-AMPK alpha 1 + AMPK alpha 2 antibody [EPR19549] ab207442 (Abcam; dilution: 1/1000); Anti-AMPK beta 1 antibody [Y367] ab109268 (Abcam; dilution: 1/5000); Recombinant Anti-AMPK gamma 1 antibody [y308] ab32508 (Abcam; dilution: 1/5000); Recombinant Anti-AMPK alpha 1 (phospho T183) + AMPK alpha 2 (phospho T172) antibody [EPR5683] ab133448 (Abcam; dilution: 1/5000); Anti-mTOR antibody [EPR390(N)] ab134903 (Abcam; dilution: 1/5000); Anti-mTOR (phosphor s2448) antibody [EPR426(2)] ab109268 (Abcam; dilution: 1/5000); Anti-Adiponectin antibody [EPR17019] ab181281 (Abcam; dilution: 1/1000); Anti-GAPDH antibody [EPR 16891] – Loading Control ab181602 (Abcam; dilution: 1/5000). Immunodetection was performed using Goat Anti-Rabbit IgG H&L (HRP) ab6721 (Abcam; dilution: 1/3000) secondary antibody, and an enhanced chemiluminescence device (Bio-Rad gel imager).

**Mitochondrial DNA copy number assay**

DNA of PM and QF-VM tissues from the SD (n=9), SD-CR (n=16), HCD (n=14), and HCD-CR (n=30) was isolated with the TIANamp Genomic DNA Kit (TIANGEN) and the mitochondrial DNA was analyzed with a Mouse Mitochondrial DNA Copy Number Assay Kit (Detroit R&D, Inc.) through comparison of mitochondrial (mt) and nuclear (n) DNA measured by qPCR.

The mitochondrial DNA copy number was calculated as follows:

\[ \Delta C_{t1} = C_{t} \text{(control mitochondrial DNA)} - C_{t} \text{(control nuclear DNA)} \]

\[ \Delta C_{t2} = C_{t} \text{(experimental group mitochondrial DNA)} - C_{t} \text{(experimental group nuclear DNA)} \]

The Mouse Mitochondrial DNA Copy Number Assay Kit protocol is annexed as Supplementary Material 1.

**Telomerase detection assay**

Tissue homogenates were analyzed with a TRAPEze RT Telomerase Detection Kit (Millipore) for fluorometric detection and real-time quantification of telomerase activity.

Briefly, 50 to 100 mg of PM and QF-VM frozen tissues from the SD (n=10), SD-CR (n=19), HC (n=14), and HC-CR (n=33) were homogenized and resuspended in 200 μl of CHAPS Lysis Buffer. The samples were incubated in ice for 30 minutes and then centrifuged at 12,000 x g for 20 minutes at 4C. One hundred sixty μl of the supernatant was transferred into a fresh tube and used to determine protein concentration. The remaining extract was used to perform the RT telomerase assay according to the kit protocol (Supplementary Material 2).

**NAD-dependent deacetylase sirtuin-1 assay**

One hundred mg of PM and QF-VM tissue homogenates from the SD (n=10), SD-CR (n=19), HC (n=14), and HC-CR (n=33) were analyzed with a Mouse NAD-Dependent Deacetylase Sirtuin-1 (SIRT1) ELISA Kit (CUSABIO) for the quantitative determination of SIRT1, according to its manual instructions (Supplementary Material 3).

**Statistical analysis**

The data were analyzed with a two-tailed unpaired t-test with Welch’s correction for comparisons of two groups or a two-way ANOVA with a Bonferroni posthoc analysis to determine interactions between multiple groups. Correlations were tested by Pearson correlation analysis and the data were processed using GraphPad Prism 8.3.0. Western blot analysis and protein densitometry were conducted with Image Lab 6.1. A P value < 0.05 was considered to indicate statistical significance.
Abbreviations

CR: Calorie restriction; HC: High calorie control group; HC-CR: High calorie-calorie restriction group; HCD: High calorie diet; IM: Intra-myocellular; INTM: Inter-myocellular; HMW: High molecular weight; MRFs: Myogenic regulatory factors; PM: Pectoralis major; QF-VM: Quadriceps femoris vastus medialis; SC: Stem cells; SD: Standard diet control group; SD-CR: Standard diet-calorie restriction group.

AUTHOR CONTRIBUTIONS

Martin Maldonado: Conceptualization, Experiments, Data analysis, Investigation, Methodology, Visualization, Writing - review and editing. Jianying Chen: Formal analysis, Investigation, Visualization, experiments. Lujun Yang: Conceptualization. Huiqin Duan: Visualization, Mazhar Ali Raja: Visualization. Ting Qu: Visualization. Tianhua Huang: Supervision. Jiang Gu: Supervision. Ying Zhong: Resources. All authors have given approval to the final version of the manuscript.

CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

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Editorial Note

*This corresponding author has a verified history of publications using the personal email address for correspondence.

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PMID: 28458782

PMID: 27924077
SUPPLEMENTARY MATERIALS

Supplementary Tables

Please browse Full Text version to see the data of Supplementary Tables 2, 4.

**Supplementary Table 1. Standard food composition.**

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<th>Number</th>
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<td>Coarse fiber</td>
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Supplied by the manufacturer company: Beijing Keao Xieli Feed Co., Ltd. (Beijing Chaoyang district, Yangshan road, number 4).

*The standard food provides 3.1 kcal/kg of energy.

**Supplementary Table 2. High-Calorie food composition.**
## Supplementary Table 3. List of primers.

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## Supplementary Table 4. Satellite markers and other genes expression.
**Supplementary Materials**

Please browse Full Text version to see the data of Supplementary Materials 1–3.

**Supplementary Material 1.** Mouse mitochondrial DNA copy number assay kit.

**Supplementary Material 2.** TRAPEze® Kit RT telomerase detection kit

**Supplementary Material 3.** Mouse NAD-Dependent Deacetylase Sirtuin-1 (Sirt1) ELISA Kit.