Research Paper

Day-Night Oscillation of Atrogin1 and Timing-Dependent Preventive Effect of Weight-Bearing on Muscle Atrophy

Shinya Aoyama,a,b, Shuichi Kojima,a Keisuke Sasaki,a, Ryosuke Ishikawa,a Mizuho Tanaka,a Takeru Shimoda,a Yuta Hattori,a Natsumi Aoki,a,b, Kengo Takahashi,a Rina Hirooka,a, Miku Takizawa,a Atsushi Haraguchi,a Shigenobu Shibata,a,b

Laboratory of Physiology and Pharmacology, School of Advanced Science and Engineering, Waseda University, Tokyo, Japan, Organization for University Research Initiatives, Waseda University, Tokyo, Japan

ABSTRACT

Background: Atrogin1, which is one of the key genes for the promotion of muscle atrophy, exhibits day-night variation. However, its mechanism and the role of its day-night variation are largely unknown in a muscle atrophic context.

Methods: The mice were induced a muscle atrophy by hindlimb-unloading (HU). To examine a role of circadian clock, Wild-type (WT) and Clock mutant mice were used. To test the effects of a neuronal effects, an unilateral ablation of sciatic nerve was performed in HU mice. To test a timing-dependent effects of weight-bearing, mice were released from HU for 4 h in a day at early or late active phase (W-EAP and W-LAP groups, respectively).

Findings: We found that the day-night oscillation of Atrogin1 expression was not observed in Clock mutant mice or in the sciatic denervated muscle. In addition, the therapeutic effects of weight-bearing were dependent on its timing with a better effect in the early active phase.

Interpretation: These findings suggest that the circadian clock controls the day-night oscillation of Atrogin1 expression and the therapeutic effects of weight-bearing are dependent on its timing.

1. Introduction

Various physiological events, including the sleep wake cycle, body temperature, and locomotor activity, exhibit circadian rhythms. These day-night fluctuations are driven by an internal circadian clock. A mammalian circadian clock is divided into two parts: the central clock in the suprachiasmatic nucleus (SCN) of the hypothalamus and peripheral clocks in peripheral tissues [1]. The central clock in the SCN has a role as a time keeper, and orchestrates and integrates the peripheral circadian clocks via neuronal and hormonal signals such as the sympathetic nervous system and glucocorticoid signaling [2,3]. Light is the major entraining factor for the SCN. Peripheral clocks are entrained not only by neuronal and hormonal signals via light-dependent regulation of the central clock in the SCN but also additionally by feeding and locomotor activity stimuli independent from the central clock in the SCN [4–6].

The molecular system of a mammalian circadian clock consists of a transcriptional and translational negative feedback loop of core clock genes, which include Brain and muscle ARNT-like 1 (Bmal1), Circadian locomotor output cycles kaput (CLOCK), Period1 (Per1), Period2 (Per2), Cryptochrome1 (Cry1), and Cryptochrome2 (Cry2) [7]. In brief, a heterodimer of BMAL1 and CLOCK acts as a transcriptional factor for Pers and Cry5 via binding to their E-box binding element. After their transcription and translation, PER1 and CRY1 inhibit the CLOCK and BMAL1-induced transcription of Pers and Cry5 and are degraded by ubiquitination systems.

Circadian transcriptomic studies have revealed that global rhythmic genes expression occurs in a tissue-specific manner, suggesting that the peripheral clocks generate the circadian fluctuation of tissue-specific physiological functions [8,9]. The circadian transcriptomes of mouse and human adult skeletal muscle have been reported [8,10–13], and the rhythmic gene expression is muscle-fiber type-specific [10]. In skeletal muscle, it is thought that the rhythm of muscular gene expression is regulated not only by an intrinsic muscle clock but additionally by feeding time and locomotor activity [10,14,15]. These rhythmic genes include several muscle-specific genes, such as Myogenic differentiation...
Physical inactivity and immobility induced muscle loss, and weight-bearing exercise is effective therapy for prevention of disused-muscle atrophy. An intermittent weight-bearing routine attenuates an increase of Atrogin1 and Murf1 expressions in the muscles of hindlimb unloaded (HU) mice [28,29]. However, a timing-dependent effect of weight-bearing has not yet been studied, although it has been suggested that muscle catabolic processes and muscle mass are controlled by a circadian clock [12,18].

In the present study, we examined the expression rhythm of Atrogin1 and Murf1 under the HU condition and the role of the circadian clock on their expression rhythm and muscle loss. Additionally, we investigate a timing-dependent preventive effect of intermittent weight-bearing on a HU-induced muscle atrophy.

2. Materials and methods

2.1. Animals

Six-week-old male Kwl:ICR mice (body weight: 25.7–34.8 g) were obtained from Tokyo Laboratory Animals Science. C57BL/6-J-Clock<sup>Δ19</sup>-J (Clock mutant) mice were obtained from Jackson Laboratory (RRID: IMSR_JAX:002923) and backcrossed to ICR mice in our previous report [30]. Six- to ten-week-old male wild-type (WT; body weight: 27.6–42.7 g) and Clock mutant (Clock<sup>Δ19</sup>; body weight: 27.7–42.0 g) mice backcrossed to ICR mice were used. The animal facility is operated as a conventional room. Mice were kept in a room maintained at 22 ± 2°C, 60 ± 5% humidity on a 12-h light (08:00–20:00) -dark cycle. Zeitgeber time 0 (ZT0) was designated as lights-on time and ZT12 as lights-off time. The mice were provided with a standard diet (EF; Oriental Yeast) and water ad libitum. All mice were in group housing (4 mice per cage) during the acclimation period, and subsequently singly housed during experimental periods. After randomized grouping, experiments were conducted in a nonblinded condition. This study was approved by the Committee for Animal Experimentation at Waseda University (2017-A070) and animals were treated in accordance with the committee’s guidelines.

2.2. Hindlimb-unloading and weight-bearing

Hindlimb-unloading was performed as described previously [31]. A plastic band was fixed to the root and middle region of the mouse tail with surgical tape under isoflurane anesthesia. Three days after attachment of the plastic band, a rod was connected to the plastic band via a fishing swivel and the other end of the rod was attached to the top of the cage. Mice were able to move freely with their front legs and were allowed access to food and water without contact of each hindfoot to the floor. Intermittent weight-bearing was performed using a timing-regulated weight-bearing apparatus. Its design is shown in Supplementary Fig. 1. The intermittent weight-bearing was automatically regulated by a magnetic switch connected to a timer.

2.3. Ablation of the sciatic nerve

The unilateral sciatic nerve was excised as described previously [32]. Briefly, excision of the sciatic nerve was performed for the right leg, and sham surgery was performed for the left leg of each mouse during the same operation under isoflurane anesthesia (Wako Chemicals).

2.4. Measurement of locomotor activity

Locomotor activity of mice was monitored with an area sensor (F5B; Omron) and analyzed with ClockLab software (Actimetrics) as previously described [33]. Locomotor activity was continuously monitored during experimental periods.
2.5. Histological analysis

The gastrocnemius muscles were isolated for hematoxylin and eosin (H&E) staining. The muscles were frozen in a liquid nitrogen-cooled isopentane (Wako Chemicals). Subsequently, ten or sixteen-micrometer thick cross sections were cut from the middle portion of the muscle using a cryostat (Leica Microsystems). Sections were fixed in 10% formalin neutral buffer (Wako Chemicals) and then stained with hematoxylin solution (Wako Chemicals) followed by the eosin solution (Wako Chemicals). Cross-sectional area (CSA) of muscle fibers was determined using BZ-X analyzer software (Keyence). >500 fibers per mouse were analyzed from three areas in each muscle.

2.6. Measurement of serum corticosterone

Serum corticosterone levels were measured using a commercial kit (ASSAYPRO). Assays were performed according to the manufacturer's instructions.

2.7. Total RNA extraction and Real-time RT-PCR

Total RNA in skeletal muscle was extracted with TRIzol reagents (Thermo Fisher Scientific). Real-time reverse transcription PCR was performed using the One-Step SYBR RT-PCR Kit (Takara Bio Inc) with specific primer pairs (Supplementary Table 1) on a Piko Real PCR system (Thermo Fisher Scientific). The relative expression levels of target genes were normalized to those of TATA box binding protein (Tbp). Data were analyzed using the ΔΔCt method as described in our previously reports [33].

2.8. Protein extraction and western blotting

Frozen gastrocnemius muscles were ground into a powder using a frozen cell crusher (Cryo-Press, MICROTEC, Tokyo, Japan) and homogenized using TissueLyser II (Qiagen, Frederick, MD, USA) with RIPA buffer (50 mM Tris-HCl pH 7.6, 150 mM NaCl, 1 mM EDTA, 1% TritonX-100, 1% sodium deoxycholate, 0.1% SDS) in the presence of protease inhibitor cocktail (Roche Diagnostics, Indianapolis, IN, USA), and phosphatase inhibitor cocktail(Nacalai Tesque, Kyoto, Japan). After homogenizing, samples were rotated for 1.5 h at 4 °C, then centrifuged at 14,000 × g for 30 min at 4 °C. Protein concentrations were measured with a BCA protein assay kit (Thermo Fisher Scientific). SDS-PAGE was performed with 20 or 50 μg of protein and the proteins in the gel were transferred onto polyvinylidene difluoride (PVDF) membranes (GE Healthcare, Buckinghamshire, UK). The membranes were then incubated overnight at 4 °C with anti-p70S6K antibody (1:1000 dilution), anti-phospho-p70S6K antibody (1:1000 dilution), anti-phospho-S6 antibody (1:1000 dilution), anti-S6 antibody (1:1000 dilution), anti-phospho-Akt antibody (1:100 dilution), anti-Akt antibody (1:1000 dilution) or anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) antibody (1:250 dilution). Phospho-p70S6 kinase (Thr389) antibody (Cat#9205, RRID:AB_330944), p70S6 kinase antibody (Cat#9202, RRID:AB_331676), Phospho-S6 (Ser240, Ser244) antibody (Cat#2215, RRID:AB_331683), S6 antibody (Cat#2217, RRID:AB_331355), phospo-Akt (Ser473) antibody (Cat#9271, RRID:AB_329825), Akt-antibody (Cat#9272, RRID:AB_329827) and horseradish peroxidase (HRP)-conjugated anti-rabbit IgG antibody (Cat# 7074, RRID:AB_2099233) were purchased from Cell Signaling Technology (Danvers, MA, USA). Anti-GAPDH antibody (Cat# sc-20,357, RRID:AB_641107) and HRP-conjugated anti-goat IgG antibody (Cat# sc-2020, RRID:AB_631728) were purchased from Santa Cruz biotechnology (Santa Cruz, CA, USA). The bands of immunoreactive proteins were detected with an enhanced chemiluminescence kit (GE Healthcare, Buckinghamshire, UK) and quantified using a LAS-3000 system (GE healthcare, Buckinghamshire, UK).

2.9. Statistical analysis

Data are represented as mean ± SE values. The GraphPad Prism version 7 (GraphPad Software) was used for the statistical analysis. P < .05 was considered as a statistically significant difference. To test whether data (sample size: n = 4) showed normal or non-normal distribution and equal or biased variation, we used Kolmogorov-Smirnov test and an F-test or Brown-Forsythe's test, respectively. If the data showed normal distribution and equal variation, statistical significance was determined by unpaired t-test or a one-way ANOVA with a Tukey test or two-way ANOVA with a Tukey test (if the interaction was significant) or Sidak test (if the interaction was not significant but the main effect was significant) for post-hoc analysis. If the data showed non-normal distribution or biased variation, statistical significance was determined by Mann-Whitney U test or Kruskal-Wallis test with a two-stage linear step-up procedure of the Benjamini, Krieger, and Yekutieli test for multiple comparisons. For the data of small sample size (n < 5), the statistical significance was determined using the Mann-Whitney U test or Kruskal-Wallis test with a two-stage linear step-up procedure of the Benjamini, Krieger, and Yekutieli test for multiple comparisons. Value of n in the figure legends represents the number of animals (The details were shown in Supplementary Table 2). The statistical details of experiments were shown in the figure legends and Supplementary Table 3. The JTK_cycle algorithm was used to detect 24-h rhythms and to analyze day-night variation and amplitude [34,35]. The results of the JTK_cycle algorithm were listed in Supplementary Table 4.

3. Results

3.1. Day-night fluctuations of catabolic processes in HU-induced atrophic muscles

Mice were placed into the HU model for 3 to 4 days to test whether the Atrogin1 and Murf1 showed day-night variation in a muscle atrophy conditioning (Fig. 1a). HU increased Atrogin1 expression significantly in the gastrocnemius muscle, and the gene exhibited day-night variation, which was lower at ZT9 and higher at ZT21 (Fig. 1b and Supplementary Table 4). Murf1 expression likewise exhibited day-night variation in a similar pattern as Atrogin1, though the level of Murf1 was not increased by HU (Fig. 1c and Supplementary Table 4). The clock gene expressions including Per1, Per2, Bmal1, Rev-erba, and Rorα showed circadian oscillation in the gastrocnemius muscle from CON and HU mice, and Per1 and Rorα expressions were increased by HU (Fig. 1d and Supplementary Table 3). The serum corticosterone level of HU group tended to be increased, and its circadian rhythm was maintained (Fig. 1f and Supplementary Table 3). The glucocorticoids receptor gene expression was not changed among all group and time points (Supplementary Fig. 2b). Phosphorylation of p70S6K, S6 and Akt, the key molecules for muscle anabolic processes, did not show day-night variation in both groups, and S6 and Akt phosphorylation were partially decreased by HU (Fig. 1k and Supplementary Tables 3 and 4). In the soleus muscle, Atrogin1 and Murf1 were increased by HU, however they were not changed over a day (Supplementary Fig. 3a and b). Per2 and Bmal1 showed circadian oscillation in the soleus muscle (Supplementary Fig. 3c and d and Supplementary Table 4). Furthermore, the peak time and periods of PER2::LUC luminescence in SCN slices were not changed among all groups (Supplementary Fig. 4).

3.2. Circadian clock drives HU-induced day-night expression of Atrogin1 and Murf1

To confirm the role of a circadian clock on the day-night expressions of Atrogin1 and Murf1, these expressions were assessed in the muscles of Clock−/− mice. In WT mice, Atrogin1 expression was increased by HU for 3 to 4 days and it showed the day-night fluctuation, while its fluctuation was not observed in Clock−/− mice (Fig. 2a and
Murf1 expression in ClockΔ19 mice was partially increased by HU and its time-dependent expression was not observed (Fig. 2b and Supplementary Table 4). We next evaluated the muscle weight in the hindlimb-unloaded WT and ClockΔ19 mice. HU for 2 weeks significantly decreased gastrocnemius muscle weight in the WT and ClockΔ19 mutant mice (Fig. 2c and d). The ratio of HU muscle weight to CON muscle in ClockΔ19 mice was significantly decreased compared with WT mice (Fig. 2e).

Supplementary Table 4). Murf1 expression in ClockΔ19 mice was partially increased by HU and its time-dependent expression was not observed (Fig. 2b and Supplementary Table 4). We next evaluated the muscle weight in the hindlimb-unloaded WT and ClockΔ19 mice. HU for 2 weeks significantly decreased gastrocnemius muscle weight in the WT and ClockΔ19 mutant mice (Fig. 2c and d). The ratio of HU muscle weight to CON muscle in ClockΔ19 mice was significantly decreased compared with WT mice (Fig. 2e).
Considering that the expressions of Atrogin1 and Murf1 were regulated by nutritional and circadian mechanisms [36] and that feeding behaviors of mice showed a circadian manner, we next examined an effect of time-restricted feeding on the day-night expression of Atrogin1 and Murf1 to reveal a feeding cycle-dependent regulation of their rhythmic expressions. Mice were kept under the day-time (ZT2 to ZT10, DRF) or night-time (ZT14 to ZT22, NRF) restricted feeding schedule for 7 days before and 4 days after HU (Supplementary Fig. 5a). The day-night expressions of Atrogin1 and Murf1 were not changed between the DRF and NRF mice (Supplementary Fig. 5b and c), suggesting that the fluctuations of Atrogin1 and Murf1 expression could be independent of feeding time. In addition, the expression of clock genes except Rorα, which did not show circadian fluctuations under the HU condition (Supplementary Table 4), exhibited an antiphase oscillation in the gastrocnemius muscle of day- and night-time-restricted feeding groups (Supplementary Fig. 5d–h). Moreover, the amplitude of Bmal1 was decreased by NRF mice (Supplementary Table 4). This suggested that the day-night fluctuations of Atrogin1 and Murf1 were independent of the intrinsic muscle clock, and these fluctuations could be controlled by a time-dependent external signal such as neuronal and hormonal signals from the SCN.

Circadian oscillated genes in the skeletal muscle were regulated by the intrinsic muscle clock, locomotor activity, neuronal and hormonal stimuli, and feeding cycle [10,11,14]. We next determined the role of locomotor activity and neuronal signals on the day-night expression of Atrogin1 and Murf1 in the HU condition using the combination of HU model and catabolic stimulus (Supplementary Fig. 5a). The day-night expressions of Atrogin1 and Murf1 were significantly decreasing of Atrogin1 and Murf1 expression in the denervation group (Supplementary Fig. 5b and c), suggesting that the fluctuations of Atrogin1 and Murf1 expression could be independent of feeding time. In addition, the expression of clock genes except Rorα, which did not show circadian rhythm in the gastrocnemius muscle of HU mice (Fig. 3b), while it was observed in the sham-surgery contralateral muscle of HU mice (Fig. 3b). HU and denervation increased Atrogin1 and Murf1 independent of the intrinsic muscle clock, and these fluctuations could be controlled by a time-dependent external signal such as neuronal and hormonal signals from the SCN.

Muscle weight and their combination further increased it. HU and denervation increased muscle weight of HU mice (Fig. 3b). HU and denervation increased muscle weight of HU mice, while it was observed in the sham-surgery contralateral muscle of HU mice (Fig. 3b). HU and denervation increased Atrogin1 and Murf1 independent of the intrinsic muscle clock, and these fluctuations could be controlled by a time-dependent external signal such as neuronal and hormonal signals from the SCN.

To assess the effective timing of a weight-bearing on Atrogin1 and Murf1, mice were intermittently relieved from HU at a specific time within a day (Supplementary Fig. 1) and skeletal muscles were collected at 5 points within 3 to 4 days (Fig. 1a). Weight-bearing at the early active phase (W-EAP) significantly suppressed Atrogin1 and Murf1 expressions at 1st ZT21, when Atrogin1 was at a high point of its oscillation within a day in the HU mice, compared with weight-bearing at the late active phase (W-LAP) and HU mice (Fig. 1b). The Atrogin1 expression level was significantly increased by HU (Supplementary Table 3). The HU-induced up-regulation of Atrogin1 was attenuated by the intermittent weight-bearing, and this level in W-EAP mice was significantly lower than that in W-LAP mice (Supplementary Table 3). Chronic HU for 14 days significantly decreased the gastrocnemius muscle weight, and the W-EAP attenuated the HU-induced muscle atrophy and body weight loss (Fig. 4a–c). The gastrocnemius muscle weight of W-EAP mice was higher than that of W-LAP mice (Fig. 4a). Food intake was not different among all groups (Fig. 4d). The hourly locomotor activity level during active phase was decreased in HU, W-EAP and W-LAP mice (Fig. 4e), and W-EAP increase the locomotor activity during late active phase (no significant difference). The total activity levels were not changed significantly however seemed to decrease in HU, W-EAP and W-LAP mice (Fig. 4g). The activity level during the respective weight-bearing times were not different between the W-EAP and W-LAP mice (Fig. 4h).

3.3. Timing-dependent effect of weight-bearing on HU-induced muscle atrophy

The effectiveness of a weight-bearing on Atrogin1 and Murf1, mice were intermittently relieved from HU at a specific time within a day (Supplementary Fig. 1) and skeletal muscles were collected at 5 points within 3 to 4 days (Fig. 1a). Weight-bearing at the early active phase (W-EAP) significantly suppressed Atrogin1 and Murf1 expressions at 1st ZT21, when Atrogin1 was at a high point of its oscillation within a day in the HU mice, compared with weight-bearing at the late active phase (W-LAP) and HU mice (Fig. 1b). The Atrogin1 expression level was significantly increased by HU (Supplementary Table 3). The HU-induced up-regulation of Atrogin1 was attenuated by the intermittent weight-bearing, and this level in W-EAP mice was significantly lower than that in W-LAP mice (Supplementary Table 3). Chronic HU for 14 days significantly decreased the gastrocnemius muscle weight, and the W-EAP attenuated the HU-induced muscle atrophy and body weight loss (Fig. 4a–c). The gastrocnemius muscle weight of W-EAP mice was higher than that of W-LAP mice (Fig. 4a). Food intake was not different among all groups (Fig. 4d). The hourly locomotor activity level during active phase was decreased in HU, W-EAP and W-LAP mice (Fig. 4e), and W-EAP increase the locomotor activity during late active phase (no significant difference). The total activity levels were not changed significantly however seemed to decrease in HU, W-EAP and W-LAP mice (Fig. 4g). The activity level during the respective weight-bearing times were not different between the W-EAP and W-LAP mice (Fig. 4h).

Fig. 2. Day-night variations of Atrogin1 and Murf1 expressions in gastrocnemius muscle and muscle weight of control (CON) and hindlimb-unloaded (HU) wild-type (WT) and Clock mutant (ClockΔflc) mice. The expression levels of (a) Atrogin1 and (b) Murf1 were examined by real-time RT-PCR. Data are expressed as the mean ± standard error (n = 4–5 at each time point) and normalized by the CON group of each genotype. *P < .05 by Kruskal-Wallis test with a two-stage linear step-up procedure of Benjamini, Krieger and Yekutieli test. Clock genes expression were shown in Supplemental Fig. 9. (c) Gastrocnemius muscle weight and (d) relative gastrocnemius muscle weight of CON (white) and 14 days HU (black) in WT and ClockΔflc mice. (e) Ratio of relative muscle weight in the HU group to that in the CON group of each genotype. Data are expressed as the mean ± standard error (n = 6). *P < .05, **P < .01, ***P < .001 by unpaired t-test or two-way analysis of variance (ANOVA) with a Sidak post hoc test.
In WT mice, HU-induced Atrogin1 up-regulation at ZT21 was not observed in the W-EAP mice while its increasing was remained in W-LAP mice (Fig. 5a left). On the other hand, weight-bearing at early and late active phase did not prevent HU-induced Atrogin1 expression in Clock-Δ19 mice (Fig. 5a right). Murf1 expression were remarkably increased by HU and the preventive timing-dependent effect of weight-bearing was not observed in Clock-Δ19 mice (Fig. 5b). In addition, the preventive effects of W-EAP in WT mice on the muscle atrophic phenotype, including the gastrocnemius muscle weight and muscle fiber CSA, were not observed in Clock-Δ19 mice (Fig. 5c−f). The pattern of locomotor activity in Clock-Δ19 mutant mice was changed by the timing of weight-bearing, and the locomotor activity of Clock-Δ19 mutant mice was increased during weight-bearing in each group. However, the total amount of locomotor activity was not changed (Supplementary Fig. 6).

4. Discussion

In this study, we showed that (i) Atrogin1 expression showed day-night variation in gastrocnemius muscles under the HU-induced atrophic condition, (ii) their day-night variations were regulated by a circadian clock, and (iii) the preventive effects of intermittent weight-bearing on the HU-induced muscle atrophy depended on its timing.

The rhythmic expression of Atrogin1 was observed in a normal skeletal muscle from microarray data reported previously [10,12]. The Atrogin1 expressions showed day-night variation under the HU condition, and their day-night variation was not observed in Clock-Δ19 mice. To elucidate the mechanism of rhythmic Atrogin1 expression, we first expected three factors to regulate their expression levels: feeding-cycle, an intrinsic muscle clock and a glucocorticoid effect. Atrogin1
expression is regulated by nutritional signaling via the Akt-FOXO pathway [36] or glucocorticoid receptor signaling [37]. Additionally, several E-box binding sites exist in the promoter region of Atrogin1 gene. These reports suggest that feeding time, glucocorticoid rhythm, or the intrinsic muscle clock drive the day-night expression of Atrogin1.

No changes were observed in the Atrogin1 expression level and its phase between NRF and DRF mice, suggesting that feeding timing did not involve in its day-night expression. However, the phase of Atrogin1 and Murf1 in NRF and DRF mice was shifted compared with that in the mice under the free feeding condition (Fig. 1 and Supplementary Fig. 5). Although timing of feeding is suggested not to be involved in its day-night expression, time-restricted feeding may partially affect the phase of Atrogin1 and Murf1, compared with the free feeding condition. It is likely to relate on the fasting periods because fasting regulates Atrogin1 and Murf1 expression [15]. In the present study, denervation increased Per2, Bmal1, and Rorα expression and abolished the rhythmicity of Rorα. The rhythmicity and phase of clock genes expression except for Rorα remained in the denervated muscles, suggesting that the involvement of these clock genes in day-night Atrogin1 expression could be small. In addition, considering that the ablation of Rorα rhythmic expression was observed in the sham muscle of HU mice, although the expression of Atrogin1 oscillated in this muscle, Rorα was less likely to regulate

In this study, serum corticosterone level in HU mice tended to be higher and exhibited a circadian rhythm (Fig. 1i), and its phase tended to be advanced by DRF compared with NRF (Supplementary Fig. 5i). It is well-known that the phase of corticosterone is affected by time-restricted feeding [38,39]. Considering that the DRF did not change the expression of Atrogin1 and Murf1 regardless of the phase shift of corticosterone by time-restricted feeding, this suggests that serum corticosterone rhythm does not affect their day-night expression under the HU condition.

It is thought that oscillation of global circadian genes expression in skeletal muscle is driven not only by a muscle clock but additionally by a neuronal external cue such as physical activity rhythm [10]. In the present study, denervated muscle did not show the normal day-night expression cycle of Atrogin1 under the HU condition, while their day-night variations were observed in the sham-surgery contralateral muscles from the same animals. Denervation influences clock gene expression [10,27]. In the present study, denervation increased Per2, Bmal1, and Rorα expression and abolished the rhythmicity of Rorα. The rhythmicity and phase of clock genes expression except for Rorα remained in the denervated muscles, suggesting that the involvement of these clock genes in day-night Atrogin1 expression could be small.

In addition, considering that the ablation of Rorα rhythmic expression was observed in the sham muscle of HU mice, although the expression of Atrogin1 oscillated in this muscle, Rorα was less likely to regulate
the day-night expression of Atrogin1. This result suggests that a circadian neuronal cue or locomotor activity controls the HU-induced Atrogin1 rhythmic expression independent of muscle clocks. Nakao et al. reported that Atrogin1 did not show a rhythmic expression in denervated muscles [27], in agreement with our results. The dissection of the sciatic nerve remarkably inhibits muscle activity, and Dyer et al. reported that physical activity regulated the rhythm of circadian gene expression via a muscle contraction-related Ca2+-dependent pathway in muscles [10], suggesting that circadian locomotor activity rhythms drove circadian genes expression. Circadian regulation of motor neuron activity could be important for the day-night variation of Atrogin1 expression. Additionally, the sciatic nerve includes not only a motor nerve but an autonomic nerve as well [40]. Further studies are warranted to determine the role of a clock via a motor or autonomic neuronal signal on the oscillation of Atrogin1 genes expression.

The day–night expression of Atrogin1 and MuRF1 was not observed in the soleus muscle (Supplementary Fig. 3). The preventive effect of W-EAP was likewise not observed in the soleus muscle (Supplementary Figs. 3 and 7). These results suggest that the effect of circadian rhythm on skeletal muscle function depends on the muscle fiber type. It is generally thought that skeletal muscle fibers are classified into two main types: slow-twitch (type1) muscle fibers and fast-twitch (type2A, type2X, and type2B) muscle fibers. Gastrocnemius muscles are predominantly made up of fast-type fibers, while soleus muscles mainly include slow-twitch fibers. It has been reported that the responses of gene expression such as that of Atrogin1 and MuRF1 and muscle atrophy were different between fast-type muscles and slow-type muscles [41,42]. Several rhythmic genes were differentially expressed between fast- and slow-type muscles [10]. Therefore, it is possible that the circadian regulation of Atrogin1 and timing-dependent effects of weight-bearing could depend on the muscle fiber types [42]. The circadian clock-controlled Atrogin1 and MuRF1 expression could be important for the maintenance of muscle mass in the HU condition. In our study, the ClockΔ19 mice showed higher response of Atrogin1 and MuRF1 to the HU condition and the HU-induced decreasing ratio of muscle weight to the control muscle was higher in ClockΔ19 mice. It suggests that ClockΔ19 mice are susceptible to the HU. Bmal1 knockout mice show extreme muscle loss with aging [18], though skeletal muscle-specific Bmal1 knockout mice did not show this phenotype [11], suggesting that a non-muscle clock could control muscle mass. Thus, in our study, the susceptibility to the HU in ClockΔ19 mice could be caused by a dysfunction of non-muscle clock. The effects of clock disturbance on HU-induced muscle loss were examined using constant light exposure model, considered the main model of central clock disruption. (Supplementary Fig. 8). The constant light exposure did not affect muscle loss, suggesting that central clock disruption is not involved in HU-
induced muscle atrophy. On the other hand, this might be due to the fluctuation of Atrogin1 expression was maintained under longer periods (~24 h) and did not disrupt its time variation because the constant light schedule in this experiment increased the periods of locomotor activity rhythm, however did not show the arrhythmic pattern. Thus, further studies are required using other clock disturbance models such as a constant high intensity light condition and a ultradian (T7) light cycle schedule.

Preventive effects of an intermittent weight-bearing on muscle loss have been known for a long time. An intermittent weight-bearing regime for 1 to 4 h per day ameliorates HU-induced gastrocnemius and soleus muscle loss (Fig. 4 and Supplementary Fig. 7). In addition, the present study shows that the timing of weight-bearing in a day is important to prevent from HU-induced muscle atrophy and that a circadian clock may be involved in its timing-dependent effect. The amount of physical activity is one of most important factors to maintain muscle mass [45]. In our study, W-EAP showed a greater preventive effect on gastrocnemius muscle atrophy compared with W-LAP, while the locomotor activity level during weight-bearing was not changed (Fig. 4). This suggests that the preventive effect of W-EAP is independent of the locomotor activity level. Although the amount of activity was not changed by the timing of weight-bearing, W-LAP has slightly increased locomotor activity during early inactive phase (Fig. 4). It is suggested that W-LAP may occur during sleep disturbance. Atrogin1 expression at ZT21 was suppressed by W-EAP. Considering that Atrogin1 expression was higher at ZT21 and lower at ZT9, this suggested that W-EAP suppressed a rise of Atrogin1 and Murf1 expression in the late active phase. While W-LAP partially suppressed an increase in Atrogin1 expression compared with HU in a day, it did not decrease completely at ZT21 compared with W-EAP. Atrogin1 is key molecules for progression of disused muscle atrophy and it is thought that the down-regulation of Atrogin1 expression attenuates muscle atrophy [23]. Therefore, the preventive effect of W-EAP on muscle atrophy could be caused by the down-regulation of Atrogin1 expression. Interestingly, the muscle of W-EAP mice showed the loss of the Atrogin1 oscillation. The amplitude of Rev-erba and Rora, tended to be decreased in W-EAP mice (Supplementary Table 4). Rev-erba KO mice exhibit muscle loss and the up-regulation of Atrogin1 and Murf1 [22]. Thus, the loss of Atrogin1 oscillation might be due to the down-regulation of Rev-erba rhythm by W-EAP.

An intermittent weight-bearing regimen additionally attenuates the down-regulation of mTOR signaling, which is one of the key anabolic processes [29]. In our study, while the phosphorylation of p70S6K was not down-regulation of mTOR signaling, which is one of the key anabolic processes responsible for the preventive effects of W-EAP.

In the present study, we showed that the muscle-specific catabolic factors Atrogin1 exhibited day-night oscillations in the HU condition. In addition, the preventive effects of weight-bearing are dependent on its timing via a circadian clock and W-EAP was more effective for the attenuation of muscle loss. This study provides the evidence to show the importance of timing for a preventive or therapeutic intervention of exercise and rehabilitation against muscle loss.

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Declaration of interests

The authors declare that no competing interests exist.

Author contributions

S.A. and S.S. planned experiments; S.A., S.K., K.S., R.I., M.T., T.S., N.A, K.T., R.H., A.H., and M.T. performed experiments and analyzed data; S.A., Y.H., K.S. and S.S. contributed to design the automated time-controlled intermittent weight-bearing device; S.A. and S.S wrote the manuscript.

Appendix A. Supplementary data

Supplementary data to this article can be found online at [https://doi.org/10.1016/j.ebiom.2018.10.057](https://doi.org/10.1016/j.ebiom.2018.10.057).

References

[1] Bass J, Takahashi JS. Circadian integration of metabolism and energetics. Science 2010;330(6009):1340–54.
[2] Balsalobre A, Brown SA, Marucci L, et al. Resetting of circadian time in peripheral tissues by glucocorticoid signaling. Science 2000;289(5488):2344–7.
[3] Albrecht U. Timing to perfection: the biology of central and peripheral circadian clocks. Neuron 2012;74(2):246–60.
[4] Schibler U, Ripperger J, Brown SA. Peripheral circadian oscillators in mammals: time and food. J Biol Rhythms 2003;18(3):250–60.
[5] Shibata S. Neural regulation of the hepatic circadian rhythm. Anat Rec A Discov Mol Cell Evol Biol 2004;280(1):101–9.
[6] Tahara Y, Shibata S. Chronobiology and nutrition. Neuroscience 2013;253:78–88.
[7] Takahashi JS. Molecular components of the circadian clock in mammals. Diabetes Obes Metab 2015;17(Supp1):6–11.
[8] Miller BH, McDearmon EL, Panda S, et al. Circadian and CLOCK-controlled regulation of the mouse transcriptome and cell proliferation. Proc Natl Acad Sci U S A 2007;104(9):3342–7.
[9] Zhang R, Lahens NF, Ballance HI, et al. A circadian gene expression atlas in mammals: implications for biology and medicine. Proc Natl Acad Sci U S A 2014;111(45):16219–24.
[10] Dyar KA, Ciciliot S, Taglialozzi GM, et al. The calcineurin-NFAT pathway controls activity-dependent circadian gene expression in slow skeletal muscle. Mol Metab 2015;4(11):R23–33.
[11] Dyar KA, Ciciliot S, Wright LE, et al. Muscle insulin sensitivity and glucose metabolism are controlled by the intrinsic muscle clock. Mol Metab 2014;3(1):29–41.
[12] McCarthy JJ, Andrews JL, McDearmon EL, et al. Identification of the circadian transcriptome in adult mouse skeletal muscle. Physiol Genomics 2007;31(1):86–95.
[13] van Moorsel D, Hansen J, Havekes B, et al. Demonstration of a day-night rhythm in human skeletal muscle oxidative capacity. Mol Metab 2016;5(8):635–45.
[14] Anyama S, Shibata S. The role of circadian rhythms in muscular and osteous physiology and their regulation by nutrition and exercise. Front Neurosci 2017;11:63.
[15] Shavlakadze T, Anwari T, Soffe Z, et al. Impact of fasting on the rhythmic expression of myogenic and metabolic factors in skeletal muscle of adult mice. Am J Physiol Cell Physiol 2013;305(1):C26–35.
[16] Andrews JL, Zhang X, McCarthy JJ, et al. CLOCK and BMAL1 regulate Myod and are necessary for maintenance of skeletal muscle phenotype and function. Proc Natl Acad Sci U S A 2010;107(44):19090–5.
[17] Harfman BD, Schroder EA, Kachman MT, et al. Muscle-specific loss of Bmal1 leads to disrupted tissue glucose metabolism and systemic glucose homeostasis. Skelet Muscle 2016;6:12.
[18] Kondratov RV, Kondratova AA, Gorbacheva VY, et al. Early aging and age-related pathologies in mice deficient in Bmal1, the core component of the circadian clock. Genes Dev 2006;20(14):1868–73.
[19] Liu J, Zhou B, Yan M, et al. CLOCK and BMAL1 regulate muscle insulin sensitivity via SIRT1 in male mice. Endocrinology 2016;157(6):2259–69.
Schröder EA, Harfmann BD, Zhang X, et al. Intrinsic muscle clock is necessary for musculoskeletal health. J Physiol 2015;593(24):5387–404.

Woldt E, Sebti Y, Solt LA, et al. Rev-erα modulates skeletal muscle oxidative capacity by regulating mitochondrial biogenesis and autophagy. Nat Med 2013;19(8):1039–46.

Mayeuf-Louchart A, Thorel Q, Delhaye S, et al. Rev-erα regulates atrophy-related genes to control skeletal muscle mass. Sci Rep 2017;7(1):14383.

Bodine SC, Baehr LM. Skeletal muscle atrophy and the E3 ubiquitin ligases MuRF1 and MAFbx/atrogen-1. Am J Physiol Endocrinol Metab 2014;307(6):E469–84.

Mitch WE, Goldberg AL. Mechanisms of muscle wasting. The role of the ubiquitin-proteasome pathway. N Engl J Med 1996;335(25):1897–905.

Bodine SC, Latres E, Baumhueter S, et al. Identification of ubiquitin ligases required for skeletal muscle atrophy. Science 2001;294(5547):1704–8.

Gomes MD, Lecker SH, Jagoe RT, et al. Atrogin-1, a muscle-specific F-box protein highly expressed during muscle atrophy. Proc Natl Acad Sci U S A 2001;98(25):14440–5.

Nakao R, Yamamoto S, Horikawa K, et al. Atypical expression of circadian clock genes in denervated mouse skeletal muscle. Chronobiol Int 2015;32(4):486–96.

D'Aunno DS, Robinson RR, Smith GS, et al. Intermittent acceleration as a countermeasure to soleus muscle atrophy. J Appl Physiol (1985) 1987;63(1):138–44.

Aoyama S, Jia H, Nakazawa K, et al. Dietary genistein prevents denervation-induced muscle atrophy in male rodents via effects on estrogen receptor-alpha. J Nutr 2016;146(6):1147–54.

Yoshida D, Aoki N, Tanaka M, et al. The circadian clock controls fluctuations of colonic cell proliferation during the light/dark cycle via feeding behavior in mice. Chronobiol Int 2015;32(8):1145–55.