Mitochondrial co-chaperone protein Tid1 is required for energy homeostasis during skeletal myogenesis

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Abstract

Background: Tid1 is a mitochondrial co-chaperone protein and its transcript is abundantly expressed in skeletal muscle tissues. However, the physiological function of Tid1 during skeletal myogenesis remains unclear.

Methods: In vitro induced differentiation assay of mouse myoblast C2C12 cells was applied to examine the physiological role of Tid1 during skeletal myogenesis. In addition, transgenic mice with muscle specific (HSA-Cre) Tid1 deletion were established and examined to determine the physiological function of Tid1 during skeletal muscle development in vivo.

Results: Expression of Tid1 protein was upregulated in the differentiated C2C12 cells, and the HSA-Tid1⁻/⁻ mice displayed muscular dystrophic phenotype. The expression of myosin heavy chain (MyHC), the protein served as the muscular development marker, was reduced in HSA-Tid1⁻/⁻ mice at postnatal day (P)5 and P8. The protein levels of ATP sensor (p-AMPK) and mitochondrial biogenesis protein (PGC-1α) were also significantly reduced in HSA-Tid1⁻/⁻ mice. Moreover, Tid1 deficiency induced apoptotic marker Caspase-3 in muscle tissues of HSA-Tid1⁻/⁻ mice. Consistent with the in vivo finding, we observed that downregulation of Tid1 not only reduced the ATP production but also abolished the differentiation ability of C2C12 cells by impairing the mitochondrial activity.

Conclusion: Together, our results suggest that Tid1 deficiency reduces ATP production and abolishes mitochondrial activity, resulting in energy imbalance and promoting apoptosis of muscle cells during myogenesis. It will be of importance to understand the function of Tid1 during human muscular dystrophy in the future.

Keywords: Skeletal muscle, Co-chaperone, ATP, AMPK and PGC-1α

Background

Muscles are soft tissues and function to produce force and motion in animals. Muscles are primarily responsible for posture and locomotion, as well as movement of internal organs such as the contraction of the heart and the movement of food through the digestive system [1]. Muscle tissues are derived from the mesodermal layer during embryogenesis in a process known as myogenesis [2]. Muscles are primarily powered by the oxidation of fats and carbohydrates, but anaerobic chemical reactions are also used, particularly by fast twitch fibers [3]. These catabolic reactions produce adenosine triphosphate (ATP) molecules which are used to power the contraction of muscle cells [3]. Dysfunction of muscles can cause muscle fatigue and severe diseases [4]. As reported by others, the AMPK signaling pathways are important to maintain the energy homeostasis of muscle tissues [5–7].

Accumulating studies suggest that mitochondria are associated with the regulation of the skeletal muscle physiology [8–15]. In addition, increase of mitochondrial biogenesis and mitochondrial DNA (mtDNA) were observed during myoblast differentiation [8, 9, 16]. Peroxisome proliferator-activated receptor-gamma coactivator 1 alpha (PGC-1α) is a transcription co-activator critically involved in energy metabolism [17, 18]. The PGC-1α is expressed in skeletal...
**Methods**

**Cell cultivation and induced differentiation of C2C12 cells in vitro**

A detailed protocol for cell cultivation and induced differentiation of C2C12 cells (ATCC® CRL-1772™) is as described previously [29]. In brief, C2C12 cells were expanded in growth medium of Dulbecco’s modified Eagle medium (DMEM), 10% fetal bovine serum (FBS), 1% L-glutamine, and 1% penicillin-streptomycin amphotericin (PSA) at 37 °C under 5% CO₂. When cultured cells reached 90% confluence, the growth medium was replaced with differentiation medium consisting of DMEM, 2% horse serum (#16050-130, Life technologies), and 1% PSA to stimulate myotube formation. The differentiation medium was replaced every 130, Life technologies), and 1% PSA to stimulate myotube

**Generation of Tid1 floxed mice and mouse genotyping**

The following procedures were performed according to the method described by Lo et al. to generate Tid1floxed mice [24]. In brief, the first loxP site was inserted upstream of the promoter region of Tid1 which is near to a transposon sequence, and the second loxP site was inserted into intron between exon 2 and 3 as indicated in Additional file 1. Deletion of the Tid1 gene in Tid1 floxed mice was achieved by crossing Tid1floxed mice with transgenic Cre mice. After the Cre-mediated recombination, the exon 1 and exon 2 of the Tid1 gene were deleted and the rest of the exon sequences lead to encoding an early stop codon. This truncated protein caused by frameshift mutation contains 18 amino acids whose amino acid sequence is not identical to that of the Tid1 protein. All genotyping of the Cre transgene and the Tid1-deficient mice was performed by polymerase chain reaction (PCR) using genomic DNA isolated from the tail tip. Genomic DNA collected from mice tails was prepared by sodium hydroxide lyzes [30]. The genotype screening was confirmed by PCR (the primer sets: IJ-FD-2.1 (5'TTAAGGCCAGTTTGCTCTAAAAC-3') and KL-RV-3 (5'-ACTTGACTAGCCCTTTCAGCATC-3')).

**Mouse handling and muscular tissue collection**

The crossbreeding of homozygous Tid1floxed/Bos™ mice (control) with HSA–Cre transgenic mice of a C57BL/6 J background was attempted to give rise to muscle-specific Tid1-deficient mice (HSA-Tid1floxed). After birth, mice were weighed and recorded daily. They were then sacrificed at postnatal day (P)5 or P8, and the tissue samples were harvested for morphological, biochemical, and functional analyses.

**Histological and histochemical studies**

The collected tissues were dissected, fixed in a 20% formalin neutral buffer solution (Wako, Osaka, Japan) overnight, embedded in paraffin, and sectioned on a microtome using standard techniques. Hematoxylin and eosin (H&E) staining was performed as described previously [31]. For immunohistochemistry, tissues were incubated with the primary antibodies diluted in TBS-T solution, and then with secondary antibodies. The cross-sectional area was calculated by Image J [32].

**Behavioral experiments**

A Rotarod treadmill (Singa Technology Corporation) was used to analyze the motor coordination, balance, and exhaustion resistance. In brief, the mice were placed on a...
rotating rod running at different speeds and for different durations. The sensor on the bottom of the apparatus recorded the duration of mice falling down from the rotating rod. For data collection, mice were pre-trained for 3 days and trained four times per day on three consecutive days. For each training, the treadmill was held for 90 s with a steady speed of 12 rpm. Subsequently, there were three conditions for trials: test 1 (T1), 90-s trial at a steady speed of 33 rpm; test 2 (T2), 90-s trial at a steady speed of 40 rpm; and test 3 (T3), 600-s trial at a steady speed of 15 rpm.

Terminal dUTP nick-end labeling (TUNEL) assay
The collected tissues were dissected and embedded within OCT. Next, 7-μm thick frozen sections from mice tissues were sliced in a freezing cryostat at −20 °C. A fluorescein apoptosis detection kit (S7110; EMD Millipore Corporation) was used to measure the apoptotic cells according to the manufacturer’s protocol.

Mitochondrial morphology and membrane potential
The culture medium was removed from the dish and replaced with pre-warmed serum-free culture medium containing 100 nM Mito Tracker Red or 2 μg/mL JC-1. After incubation for 20 min at 37 °C, cells were immediately washed twice in cold phosphate-buffered saline (PBS) and analyzed using a Laser confocal microscope (Olympus FV1000).

ATP content assay
An ATP measurement kit for cells (K791-100; BIOVISION, USA) was used to measure the ATP concentration according to the manufacturer’s protocol. The ATP
concentration was measured using the chemiluminescence produced by the luciferase/luciferin reaction.

**Statistical analysis**
Data are expressed as means ± SD. Differences among groups were tested by analysis of variance (ANOVA). Comparisons between two groups were performed by the paired Student t test. P values less than 0.05 were considered significant.

**Results**
**Upregulation of Tid1 during induced differentiation of C2C12 myoblasts**
To evaluate the expression profile of Tid1 protein during myogenesis, we established and examined mouse myoblast C2C12 cells undergoing induced differentiation into myotubes (Fig. 1a). As shown in Fig. 1b, expression of myogenesis-related markers (myosin heavy chain (MyHC)) was upregulated, whereas PAX7 and MyoD were downregulated during the process of C2C12 differentiation. Of note, the protein level of Tid1 was also increased along with the induced differentiation (Fig. 1b). The decrease of the ATP sensor (p-AMPK) and increase of the mitochondrial biogenesis protein (PGC-1α) indicated that the mitochondrial activity and ATP production were both increased during C2C12 differentiation (Fig. 1b). The GEO Profiles database reveals that Tid1 expression of muscle is downregulated in the mitochondrial dysfunction mice in comparison with those from control mice (GEO accession: GDS4856 [33]) (Fig. 1c). Moreover, the search of the GEO Profiles database reveals that Tid1 expression of muscle is downregulated in muscular dystrophy model mice in comparison with those from control mice (GEO accession: GDS3371 [34] and GEO accession: GDS236 [35]) (Fig. 1d). These results suggest that Tid1 may play a role during myogenesis.
Phenotypic profile of postnatal transgenic and control mice on day 5

To extend the research on understanding the physiological function of Tid1 during muscular development, transgenic mice with muscle-specific Tid1 deficiency (HSA-Tid1f/f), for which cre recombinase-mediated deletion of the Tid1 gene is driven by actin promoter in muscle cells, were generated [36, 37] (Additional file 1). Interestingly, we found that the 5-day-old HSA-Tid1f/f mutant mice displayed less body weight compared to control mice (Fig. 2a and b). However, there was no significant size difference in the hind limb of HSA-Tid1f/f mice in comparison to that of the control mice (Fig. 2c). Next, the histology of the tibialis anterior (TA) muscle was analyzed; as shown in Fig. 2d, the size of muscle fibers of HSA-Tid1f/f mice were reduced compared to those of the control or HSA-Tid1f/+ mice.

More dystrophic phenotype in postnatal HSA-Tid1f/f mice on postnatal day 8

Phenotypically, the HSA-Tid1f/f mice displayed a significant reduction in body weight compared to the control mice on day P8 (Fig. 3a and b), and the size of the hind limb of HSA-Tid1f/f mice was also reduced in comparison to those of either control or HSA-Tid1f/+ mice (Fig. 3c). Histologically, the muscle biopsy displayed patchy lymphocytic infiltration in HSA-Tid1f/f mice. The presence of infiltrating lymphocytes indicated the induced inflammation in the muscular tissues of mutant mice (Fig. 3d). In addition, we observed that the HSA-Tid1f/f mice died between days P8 and P10. Because the mice died between P8 and P10, examination of the muscular physiology of adult mice with Tid1 deficiency was prevented. Nevertheless, we evaluated the muscular physiology of adult mice carrying Tid1...
heterogeneity (HSA-Tid1<sup>f/+</sup>). We assessed the motor coordination of HSA-Tid1<sup>f/+</sup> transgenic mice in comparison to that of the control mice. As shown in Fig. 3e and f, we found that the HSA-Tid1<sup>f/+</sup> transgenic mice displayed reduced running time and running distance. These results suggest that deletion of the Tid1 gene causes dysfunction of muscle tissue of transgenic mice. In addition, Tid1 haploinsufficiency could abolish the physiological muscular function.

**Imbalanced energy and induced apoptosis of muscle from HSA-Tid1<sup>f/f</sup> mice**

In order to further investigate the molecular mechanisms involved in muscular homeostasis at different time points, we harvested the protein extracts of muscle tissue from control and mutant mice at P5 and P8. Immunoblotting analyses showed that the expression levels of Tid1 and MyHC proteins were both reduced in HSA-Tid1<sup>f/f</sup> mice in comparison with those in control mice on P5 and P8 (Fig. 4a and b). In addition, we found that the expression of active AMPK (phosphorylated AMPK) protein within the muscle tissues from HSA-Tid1<sup>f/f</sup> mice was increased on P5 but reduced on P8 compared to control mice (Fig. 4a and b). Moreover, expression of mitochondrial biogenesis protein (PGC-1α) was also decreased in HSA-Tid1<sup>f/f</sup> mice (Fig. 4a and b). Of note, we observed that the expression of apoptosis indicator (cleavage Caspase-3) was increased on P5 and P8 in HSA-Tid1<sup>f/f</sup> mice (Fig. 4a and b). Meanwhile, the Tid1 deficiency induced the number of apoptotic cells within the muscle tissues from HSA-Tid1<sup>f/f</sup> compared to control mice (Fig. 4c). The above findings suggest that Tid1 deficiency dysregulates energy balance and induces muscular apoptosis during muscular homeostasis.

![Fig. 4](image-url) Imbalanced energy and induced apoptosis of muscle from HSA-Tid1<sup>f/f</sup> mice. Crude proteins extracted from the whole posterior limb of mutant and control mice on day 5 (P5) (a) and on day 8 (P8) (b) were collected, respectively. Expression of proteins of interest was detected by immunoblot analyses. c The apoptotic cells were detected by TdT-mediated dUTP nick end labeling assay (TUNEL). ***P < 0.001.
Downregulation of Tid1 abolishes the induced differentiation ability of C2C12 by reducing ATP production and impairing mitochondrial activity

To determine whether Tid1 deletion gives rise to abnormal myoblast differentiation in vitro, we manipulated the expression of Tid1 in the myoblast cell line (C2C12) undergoing induced differentiation. Knockdown of Tid1 in C2C12 cells by small hairpin RNA interference (shRNAi) reduced the myotube formation from morphological observation (Fig. 5a). In addition, we observed that C2C12 cells with Tid1 knockdown displayed both reduced mitochondrial mass (Fig. 5b and Additional file 2A) and abolished mitochondrial membrane potential (Fig. 5c and Additional file 2B) before induced differentiation in comparison with control C2C12 cells undergoing regular differentiation, but not on day 3. The intracellular ATP concentration of C2C12 cells with Tid1 knockdown was also decreased before induced differentiation but not on day 3 (Fig. 5d and Additional file 2C). Meanwhile, we found that expression of both total and active AMPK was increased in C2C12 cells with shRNAi-Tid1 before induced differentiation, and the upregulation of AMPK was reduced on induced differentiation day 3 (Fig. 6a and b). In agreement with previous results from HSA-Tid1f/f mice in vivo (see Fig. 4a and b), the expression levels of PGC-1α and MyHC protein in C2C12 cells with Tid1 knockdown by shRNAi were decreased on day 3 (Fig. 6b). We also observed that the expression level of Caspase-3 protein was increased before induced differentiation but not on day 3 within the C2C12 cell undergoing induced differentiation plus Tid1 knockdown (Fig. 6a and b). These findings suggest that Tid1 deficiency reduces the differentiation ability of myoblast through reducing ATP production and impairing mitochondrial activity.

Discussion

The physiological function of chaperone and co-chaperone in muscle cell development remains elusive [38, 39]. In order to further understand the role of Tid1,
a mitochondrial co-chaperone, on mediating myogenesis, we determined the expression of Tid1 during the induced differentiation of myoblasts (C2C12 cells) in vitro. During the differentiation of C2C12 cells, we found that the protein levels of Tid1, PGC-1α, and MyHC, as well as the mitochondrial activity, were increased during the process of induced myoblast differentiation (Fig. 1b), consistent with previous studies [8–10, 16]. These results suggest that upregulation of Tid1 may play a role during the myoblast differentiation. Indeed, Linnola et al. have reported that downregulation of Tid1 may impair the function of the neuromuscular junction [40, 41]. Of note, the HSA-Tid1f/f mice displayed muscular dystrophic phenotype at P8 but not P5 (Fig. 3a–c). In addition, the apoptosis signal cleavage Caspase-3 was enhanced in skeletal muscles of HSA-Tid1f/f mice in comparison with the control mice (Fig. 4a–c). It has been shown that the apoptotic process occurs in muscle fibers of muscular dystrophy phenotype [42–44]. Moreover, we found that AMPK activity was increased at P5 but decreased at P8, and expression of PGC-1α protein was decreased at P8 (Fig. 4a and b). These results suggest that Tid1 deficiency impairs the mitochondrial activity, resulting in insufficient ATP production and cell apoptosis. Next, downregulation of Tid1 by RNA interference reduced the myotube formation and expression of MyHC in the induced C2C12 cells (Figs. 5a and 6b). We also assessed markers of mitochondrial mass/volume (Mito Tracker staining) and ΔΨ (JC-1 staining) by fluorescence microscopy. The Tid1 knockdown impaired the mitochondrial mass and membrane potential (Fig. 5b and c) and reduced the ATP amount compared to those before induced differentiation (Fig. 5d). Meanwhile, the mitochondrial biogenesis marker PGC-1α was reduced in Tid1-knockdown C2C12 cells at day 3 after induced differentiation (Fig. 6b). These in vitro results phenotypically correlate with the HSA-Tid1f/f mice in vivo. However, we did not find the induction of C-caspase 3 by downregulation of Tid1 at day 3 (Fig. 6b). It has been reported that Caspase 3 is required for muscle differentiation [45]. We speculate that the absence of C-caspase-3 changes at day 3 was caused by the induction of Caspase 3 activity of controls during the skeletal muscle differentiation. The discrepancy between the expression of C-caspase 3 under the distinct downregulation of Tid1 and Caspase 3 during the in vitro induced differentiation may be caused by the distinct physiological effect after the gene-specific knockdown. For example, downregulation of Caspase 3 may not cause the mitochondrial abnormality or ATP defect. However, in our previous studies, Tid1 knockout attenuates the mitochondrial mass/volume and induces cell apoptosis in mouse embryonic fibroblasts and pre-matured T cells [24, 25]. Together, these results suggest that downregulation of Tid1 would initially decrease ATP production along with increased cell death, followed by reduced mitochondrial biogenesis (reduction of PGC-1α). These in vitro results imply that energy insufficiency of muscle cells mediated by Tid1 gene deletion (HAS-Tid1f/f) is likely the reason for apoptotic cell death during the myogenesis and muscular dystrophy phenotype.

**Conclusion**

Our results demonstrate that Tid1 plays a pivotal role in myogenesis. Deletion of the Tid1 gene causes dysfunction of muscle tissues of transgenic mice. Downregulation of Tid1 abolished the ability of C2C12 to differentiate via reduced ATP production, impaired mitochondrial activity, and induced apoptotic cell death.

In summary, Tid1 deficiency reduces ATP production, impairs the mitochondrial activity of muscle cells during the myogenesis, and, consequently, causes muscle cell apoptosis. However, how Tid1 deficiency reduces ATP production and inhibits PGC-1α activity remains unclear. Thus, we will further investigate the molecular mechanisms mediated by Tid1 to regulate ATP production and PGC-1α activity during myogenesis. In addition, our muscle-specific Tid1 deletion mice will provide a useful tool to study the physiological function of Tid1 in normal muscular homeostasis.
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