Macroglossia and less advanced dystrophic change in the tongue muscle of the Duchenne muscular dystrophy rat

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Abstract

Background: Duchenne muscular dystrophy (DMD) is an X-linked muscle disease caused by a complete lack of dystrophin, which stabilizes the plasma membrane of myofibers. The orofacial function is affected in an advanced stage of DMD and this often leads to an eating disorder such as dysphagia. Dysphagia is caused by multiple etiologies including decreased mastication and swallowing. Therefore, preventing the functional declines of mastication and swallowing in DMD is important to improve the patient's quality of life. In the present study, using a rat model of DMD we generated previously, we performed analyses on the masseter and tongue muscles, both are required for proper eating function.

Methods: Age-related changes of the masseter and tongue muscle of DMD rats were analyzed morphometrically, histologically, and immunohistochemically. Also, transcription of cellular senescent markers, and utrophin (Utrn), a functional analog of dystrophin, was examined.

Results: The masseter muscle of DMD rats showed progressive dystrophic changes as observed in their hindlimb muscle, accompanied by increased transcription of p16 and p19. On the other hand, the tongue of DMD rats showed macroglossia due to hypertrophy of myofibers with less dystrophic changes. Proliferative activity was preserved in the satellite cells from the tongue muscle but was perturbed severely in those from the masseter muscle. While Utrn transcription was increased in the masseter muscle of DMD rats compared to WT rats, probably due to a compensatory mechanism, its level in the tongue muscle was comparable between WT and DMD rats and was similar to that in the masseter muscle of DMD rats.

Conclusions: Muscular dystrophy is less advanced in the tongue muscle compared to the masseter muscle in the DMD rat.

Keywords: Duchenne muscular dystrophy, Masseter muscle, Tongue, Macroglossia, Rat

Background

Duchenne muscular dystrophy (DMD) is an X-linked muscle disease caused by out-of-frame mutations in the dystrophin gene that results in the complete loss of dystrophin protein [9]. Dystrophin is a structural protein that stabilizes the plasma membrane of the myofibers, and the lack of dystrophin protein causes myofibers to become fragile [9]. Skeletal muscle myofibers normally...
have a remarkable regenerative capacity [11]. Once
myofibers are damaged, the myogenic stem/progenitor
cells, satellite cells, are activated and proliferate, then
eventually fuse each other to form multinucleated myo-
tubes. The myotubes mature to constitute newly regen-
rated myofibers. DMD is characterized by repeated
cycles of degeneration and regeneration of myofibers,
which finally result in a loss of regenerative capacity due
to exhaustion of the satellite cell pool [21]. DMD patients
show progressive weakness and premature death due to a
loss of ambulation [9].

In an advanced stage of DMD, an orofacial function is
affected [3] and eating disorders such as dysphagia are
often a problem [37]. Dysphagia is caused by multiple
etiologies. Decreased mastication is one of the causes
dysphagia. Masticatory movement involves masseter,
temporalis, medial, and lateral pterygoid muscles [17].
Masticated foods are then swallowed using the tongue
muscle in the oral cavity. Swallowing is initiated by the
contraction of digastic and tongue muscles against the
palatine to move the bolus into the oropharynx, followed
by a complex coordination of pharyngeal and esophageal
muscles. The tongue muscles consist of the extrinsic mus-
cles, which move the tongue from the outside, and the
intrinsic muscles, which change the shape of the tongue.
The former consists of four types of tongue muscles: the
genioglossus, hyoglossus, styloglossus, and palatoglossus
muscles, and the latter consists of four types: the superior
longitudinal, inferior longitudinal, transverse, and verti-
cal muscles. Several reports demonstrated that dysphagia
in DMD patients is associated with dystrophic changes in
muscles related to mastication [36]. MacroGLOSSIA, which
is common in DMD patients [28, 37], is defined as an
enlarged tongue and is often associated with dysphagia
[5]. Therefore, preventing the deterioration of mastic-
ation and swallowing functions in DMD is important to
improve the patient’s quality of life.

We previously reported the generation of a rat model
of DMD (DMD rat) [22, 33]. DMD rats show a pro-
gressive muscle pathology in their hindlimb muscle
and diaphragm like human DMD [22, 33]. The patho-
logical changes seen in the muscle of DMD rats include
decreased strength, increased fibrosis, and fatty infil-
tration [22, 33]. We further demonstrated the occurrence of
premature cellular senescence in the muscle of DMD rats
and suggested that this could be one of the causes of sat-
ellite cell depletion, which leads to a loss of regenerative
capacity [33].

In the above our studies using DMD rats [22, 33], we
analyzed only the muscle of the hindlimbs and dia-
phragm. During breeding and maintenance of DMD rats,
we had experienced a decreased food intake in accord-
ance with the decrease in the body weight in their later
life, and this prompted us to examine whether the mus-
cles related to food intake are also affected. Thus, in this
study, we focused on the masseter and tongue muscles as
the muscles involved in mastication and swallowing and
investigated their development and histological changes,
as well as the dynamics of satellite cells.

Methods
Animals
We previously generated and characterized a strain of
DMD rat [22, 33]. In this strain of DMD rat, a deletion
of 329 bp around the splice site in intron2 leads to exon3
skipping, and an insertion of 1 bp in exon16 causes the
generation of a stop codon, resulting in a loss of dys-
throphin protein [22, 33]. Adult X^DmdX female rats were
mated with wild-type (WT) male rats to generate male
WT (XY) and DMD (X^DmdY) rats. They were maintained
under controlled environmental conditions, at 23 °C
with a light/dark cycle (lights on 0800–2000). Labora-
tory chow (Labo MR Standard, Nihon Nousan Co., Yoko-
hama, Japan) and water were given ad libitum. All animal
experiments performed in this study were in accordance
with the Guide for the Care and Use of Laboratory Ani-
mals of the University of Tokyo and were approved by
the Institutional Animal Care and Use Committee of the
University of Tokyo (P18-125).

Measurement of food intake
At 3 or 7 months old, the rats were transferred individu-
ally to a plastic cage and their amount of food intake was
measured for two consecutive days. The measured values
were divided by 2 and expressed as food intake per day.

Histological and immunohistochemical analyses
At indicated age, the rats were killed by inhalation of car-
bon dioxide gas. After removal of facial skin, the area cor-
responding to the masseter muscle was measured. The
masseter muscle was considered as an ellipse (Fig. 1B),
and its long and short diameters were measured with
calipers to calculate the area. Then, the lower half part of
the masseter muscle was removed. The tongue was cut
out at the root. Collected tissues were fixed in 4% para-
formaldehyde (PFA) in phosphate-buffered saline (PBS)
for paraffin-embedded sections or snap-frozen in liquid
nitrogen-cooled isopentane for cryosections.

The paraffin-embedded sections were used for hema-
toxylin–eosin (HE) or Masson’s trichrome staining. The
sections were observed and photographed using a micro-
scope (BX51, Olympus, Tokyo, Japan) equipped with a
digital camera (DP73, Olympus). The HE-stained tongue
sections were used for measurement of the width and
diagonal of the tongue (Fig. 1D). Masson’s trichrome-stained
sections were used for quantitative analysis of the fibrotic
area. Two fields were randomly selected in the section using a 4 × objective and the area occupied by fibrotic tissues stained blue and the total area of sections were calculated using ImageJ software (v1.47; National Institutes of Health, Bethesda, MD, USA).

The cryosections (7 μm) were used for immunohistochemistry. The sections were fixed with 4% PFA in PBS for 15 min, followed by blocking with 5% normal donkey serum (NDS) in PBS. The sections were incubated with primary antibodies overnight at 4 °C, then secondary antibodies for 1 h.

For identification of satellite cells, vascular endothelial cells, and mesenchymal progenitor cells, anti-Pax7 (1:200 with 5% NDS in PBS, mouse, clone P3U1; Developmental Studies Hybridoma Bank (DSHB), Iowa City, IA, USA), anti-CD31 (1:400 with 5% NDS in PBS, rabbit, NB100-2284, Novus Biological, Centennial, CO, USA), and anti-chondroitin sulfate proteoglycan 4 (CSPG4) (1:50 with 5% NDS in PBS, mouse, clone 5C12 [34]) were used as primary antibodies, respectively, followed by AlexaFluor-labeled donkey anti-mouse IgG (1:500 with 5% NDS in PBS, Jackson ImmunolResearch, West Grove, PA, USA).

For identification of necrotic myofibers, anti-laminin (1:400 with 5% NDS in PBS, rabbit, L9393 Sigma, St Louis, MO, USA) was used as the primary antibody, and AlexaFluor-labeled goat anti-rat IgG and donkey anti-rabbit IgG (1:500 with 2.5% NDS/2.5% normal goat serum (NGS) in PBS, Jackson ImmunolResearch, West Grove) were used as secondary antibodies.

For identification of newly formed regenerated myofibers, anti-embryonic myosin heavy chain (eMHC) (1:400 with 5% NDS in PBS, mouse, clone F1.652, DSHB) and anti-laminin were used as primary antibodies, and AlexaFluor-labeled donkey anti-mouse IgG and donkey anti-rabbit IgG (1:500 with 5% NDS in PBS, Jackson ImmunolResearch) were used as secondary antibodies. After the reaction with secondary antibodies, nuclei were counterstained with Hoechst 33258. The sections were photographed as described above. The myofibers were automatically identified by Cellpose Google Colab script (https://colab.research.google.com/drive/1958UQIH-XAYogKvbxnaUHALYvR73KLj2) [38], and their diameters (minimum Feret diameters) and numbers were measured using ImageJ software. The number of nuclei per myofiber was calculated by dividing the total number of nuclei by the number of myofibers.

The area occupied by CD31-positive cells and CSPG4-positive cells was calculated by ImageJ.

**Cell culture and immunocytochemistry**

Mononuclear cells were isolated from skeletal muscle as described previously [39]. In brief, the rats were killed by inhalation of carbon dioxide gas, and their tongue and masseter muscles were extirpated. As described above, the lower half part of the masseter muscle and tongue cut out at the root were used for cell isolation. They were minced with scissors and digested with 1.25 mg/mL protease (from Streptomyces griseus, type XIV; Sigma) at 37 °C for 1 h. Cells were separated from myofiber fragments through differential centrifugation and plated onto poly-L-lysine- and fibronectin-coated 48-well plates. Cells from each muscle piece were divided into two equal portions (one for immunocytochemistry of Pax7 and another for MyoD) and plated. Cells were cultured in Dulbecco’s modified Eagle medium (Gibco, Life Technologies, Palo Alto, CA, USA) containing 10% fetal bovine serum, 100 U/mL penicillin, 100 μg/mL streptomycin, and 50 μg/mL gentamicin for 4 days.

The cells were fixed with 4% PFA/PBS for 15 min and then blocked with 5% NGS/PBS containing 0.1% Triton X-100 for 10 min. The cells were incubated with anti-Pax7 (1:200 with 5% NGS/PBS) or anti-MyoD (1:200, with 5% NGS/PBS, mouse, clone 5.8A, Novocastra, Newcastle upon Tyne, UK) overnight at 4 °C. After washing, they were incubated with AlexaFluor-labeled secondary anti-rabbit IgG (1:500 with 2.5% NDS/PBS, mouse, clone P3U1; Developmental Studies Hybridoma Bank (DSHB), Iowa City, IA, USA), and anti-CD31 (1:400 with 5% NDS in PBS, rabbit, NB100-2284, Novus Biological, Centennial, CO, USA), and anti-laminin antibody. Data were expressed as percent positive myofibers or number of positive cells per area.

For quantification of myofiber size and the number of nuclei per myofiber, the sections were immunostained with anti-laminin, followed by labeling with AlexaFluor-labeled secondary antibody. The nuclei were counterstained with Hoechst 33258. The sections were photographed as described above. The myofibers were identified by double-staining with an anti-laminin antibody. Data were expressed as percent positive myofibers or number of positive cells per area.

For quantification of myofiber size and the number of positive cells per myofiber, sections were immunostained with anti-laminin, followed by labeling with AlexaFluor-labeled secondary antibody. The nuclei were counterstained with Hoechst 33258. The sections were photographed as described above. The myofibers were automatically identified by Cellpose Google Colab script, and their diameters (minimum Feret diameters) and numbers were measured using ImageJ software. The number of nuclei per myofiber was calculated by dividing the total number of nuclei by the number of myofibers.

The area occupied by CD31-positive cells and CSPG4-positive cells was calculated by ImageJ.

**Fig. 1** Food intake and morphometrical analyses of masseter and tongue muscles of WT and DMD rats. A The amount of food intake per day was measured at 3 and 7 months old. The data are expressed as mean ± SE. **, p < 0.01 by unpaired Student’s t-test. B The masseter muscle was considered as an ellipse (dotted white circle) and its long and short diameters were measured with calipers to calculate the area. C Changes in the area of the masseter muscle at 1, 3, 6, and 8 months old. The data are expressed as mean ± SE. **, p < 0.01 by unpaired Student’s t-test. D The tongue was cut at the root (dotted black line) (left) and the width (dotted black line) and area (dotted red circle) were measured. Scale bar = 1 cm. E Changes in the width and cross-sectional area of the tongue at 1, 3, and 8 months old. The data are expressed as mean ± SE. **, p < 0.01 by unpaired Student’s t-test.
Fig. 1 (See legend on previous page.)
antibody for 1 h. Nuclei were counterstained with Hoechst 33,258. The Pax7- and MyoD-positive cells were counted in randomly selected five fields at 20 × objective of a fluorescence microscope (BX50, Olympus). The total numbers of nuclei were also counted. The data were expressed as percent positive cells.

Quantitative real-time PCR
Total RNA was extracted from 100 cryosections (7 μm) with FastGene RNA Basic kit (Nippon Genetics Co., Ltd., Tokyo, Japan) and reverse transcribed to cDNA using Super Script II kit (Invitrogen). Quantitative real-time PCR was performed on a Light Cycler 2.0 (Roche Diagnostics, Roche, Basel, Switzerland) with the Thunderbird SYBR qPCR Mix (TOYOBO, Osaka, Japan). The following primer sets were used: p16: forward, 5′-TTC ACC AAA CGC CCC GAA CA-3′; reverse, 5′-CAG GAG AGC TGC CAC TTT GAC-3′; p19: forward, 5′-GTG TTG AGG CCA GAG AGG AT-3′; reverse, 5′-TTG CCC ATC ATC ATC ACC T-3′; p21: forward, 5′-GAC ATC TCA GGG CCG AAA-3′; reverse, 5′-GGC GCT TGG AGT GAT AGA AA-3′; p53: forward, 5′-AGA GAC GAG TGC CCA CCA-3′; reverse, 5′-AAC ATC TCG AAG CGC TCA C-3′; Utrn: forward, 5′-TAG AGC AAT AAC CTG CCA CAC GA-3′; reverse, 5′-ACG CTC TTC CTT CTC CAG AC-3′; MuRF1: forward, 5′-AGG ACT CCT GCC GAG TGA C-3′; reverse, 5′-TTG TGG CTC AGT TCC TTC TT-3′; Atrogin1: forward, 5′-GAA GAC CGG CTA CTG TGG AA-3′; Atrogin1: reverse, 5′-ATC AAT CGC TTG CGG ATC T-3′, and HPRT: forward, 5′-GAC CGG TTC TGT CAT GTC G-3′; reverse, 5′-ACC TGG TTC ATC ATC AAT CAC-3′. The expression of each gene was analyzed using the crossing-point method and expressed after normalization with that of HPRT.

Western blotting
Two hundred cryosections (7 μm) of the tongue muscle were homogenized using radioimmunoprecipitation assay (RIPA) buffer [10 mM NaH2PO4, 150 mM NaCl, 2 mM ethylenediaminetetraacetic acid (EDTA), 0.1% sodium deoxycholate, 1% Nonidet P-40, 10 μg/mL leupeptin, 5 μg/mL pepstatin, 1.84 g/L Na3VO4, 10 mg/mL p-nitrophenylphosphate (PNPP), 100 KIU/mL apronin, 20 μg/mL phenylmethanesulfonylfluoride (PMSF)], and protein concentration of the lysates was determined using Pierce BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, MA, USA). Lysates were then, diluted with RIPA buffer to the same concentrations of protein. The samples were subjected to sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) and western blotting as described in the previous study [1]. Antibodies used were as follows: anti-phospho-Akt antibody (Ser473, Cell Signaling Technology, Danvers, MA, USA, #9271), anti-Akt antibody (Cell Signaling Technology, #9272), anti-phospho-p70 S6K antibody (Thr389, Cell Signaling Technology, #9234), anti-p70 S6K antibody (Santa Cruz Biotechnology, C-18, sc-230), anti-phospho-S6 ribosomal protein antibody (Ser240/244, Cell Signaling Technology, #2215), anti-S6 ribosomal protein antibody (Cell Signaling Technology, 54D2, #2317), anti-4EBP1 antibody (Cell Signaling Technology, #9452), anti-α-tubulin antibody (Sigma Aldrich, T9026), anti-rabbit IgG, horseradish peroxidase (HRP)-linked whole antibody from donkey (GE healthcare, NA934V), and antier-mouse IgG, HRP-linked whole antibody from sheep (GE healthcare, NA931V). Band intensities of each blot were quantified using ImageJ.

Statistical analyses
Except for an analysis of myofiber diameters, graphed data are expressed as means±SE. Unpaired Student’s t-test (between two groups) and one-way analysis of variance (ANOVA) followed by a Tukey–Kramer test (for multiple group comparison) were used to evaluate statistical differences. For the experiment to assess the difference of myofiber diameters, the p-value was determined using the Wilcoxon rank sum test. p-values less than 0.05 were considered statistically significant.

Results
We previously reported that the body weight of DMD rats decreased progressively after the age of 6 months [33]. Suspecting that eating disorder occurs in DMD rats, we measured the amount of food intake at the ages of 3 months (before the onset of body weight decrease) and 7 months (after the onset of bodyweight decrease). The amount of food intake at 3 months old was comparable between WT and DMD rats (Fig. 1A). However, at 7 months old, significantly decreased food intake was observed in DMD rats, indicating the occurrence of an eating disorder (Fig. 1A).

The above results suggest that muscles involved in food intake are affected by the progress of the disease. We then performed morphometric analyses of muscles involved in food intake. We chose the masseter and tongue muscles because these two muscles are to be approached easily and distinguishable from other muscles. As shown in Fig. 1B, the area corresponding to the masseter muscle was considered as an ellipse, and its long and short diameters were measured with calipers to calculate the area. The area of the masseter muscle was measured from 1 to 8 months old. At 1 month old and after 6 months old, the area of the masseter muscle was significantly reduced in DMD rats compared to WT rats. Especially, the greater reduction was observed at 6 and 8 months old (Fig. 1C).
The appearance of the tongue at 8 months old is shown in Fig. 1D. The cross-section of the posterior part of the tongue was larger in DMD rats than in WT rats (Fig. 1D). Then, we measured the area and width of the upper half of the tongue sections. Quantitative analysis showed that the width of the tongue of DMD rats was significantly larger than that of WT rats after 3 months old (Fig. 1E). In addition, the area of the tongue was significantly larger in DMD rats at the age of 8 months than in WT rats (Fig. 1F). These results indicate that macroglossia is occurring in DMD rats.

To further gain insight into the age-related changes of the masseter and tongue muscles of DMD rats, histological analyses were performed. For the tongue muscles, the intrinsic and extrinsic muscles were examined separately. The masseter muscle of DMD rats showed myofiber degeneration and necrosis with infiltration of neutrophils and macrophages at 1 month old (Fig. 2B) and regenerative myofibers with enlarged centralized nuclei and basophilic cytoplasm at 3 months old (Fig. 2D). At 8 months old, in addition to inflammatory and degenerative changes, severe fibrosis and fatty infiltration were observed (Fig. 2F, H). No significant pathological changes were observed in the masseter muscle of WT rats at all ages (Fig. 2A, C, E, G).

In contrast, in the intrinsic tongue muscle of DMD rats, regenerative myofibers with enlarged centralized nuclei and basophilic cytoplasm were observed at 1 month old (Fig. 2J), and these regenerative changes were accompanied by mild fibrosis at 3 months old (Fig. 2L). At 8 months old, mild fibrosis, also confirmed by Masson’s trichrome stain (Fig. 2P), was persisted (Fig. 2N). The extrinsic tongue muscles generally showed similar findings as to the intrinsic tongue muscles, although regenerative changes were observed in some parts of the genioglossus muscles of DMD rats at 3 and 8 months.

![Fig. 2 Histological analyses of masseter and tongue muscles of WT and DMD rats. A-F HE- and G, H Masson’s trichrome-stained sections of the masseter muscle of WT and DMD rats. In the masseter muscle of DMD rats, myofiber degeneration and necrosis with infiltration of neutrophils and macrophages at 1 month old, regenerative myofibers with enlarged centralized nuclei and basophilic cytoplasm at 3 months old, and severe fibrosis, confirmed by Masson’s trichrome staining, at 8 months old were observed, respectively. No significant pathological changes were observed in the masseter muscle of WT rats at all ages. I-N HE- and O, P Masson’s trichrome-stained sections of the tongue muscle of WT and DMD rats. In the tongue muscle of DMD rats, regenerative myofibers with enlarged centralized nuclei and basophilic cytoplasm at 1 month old, the regenerative changes accompanied by mild fibrosis at 3 months old, and persistence of mild fibrosis, confirmed by Masson’s trichrome staining, at 8 months old were observed, respectively. No significant pathological changes were observed in the tongue muscle of WT rats at all ages.](image-url)
old (Supplementary Fig. 1). No significant pathological changes were observed in both the intrinsic and extrinsic tongue muscle of WT rats at all ages (Fig. 2I, K, M, O).

To further examine whether myofiber necrosis and muscle regeneration are ongoing after 6 months old when macroglossia is occurring in the tongue muscle of DMD rats, immunohistochemical analyses of endogenous IgG (myofiber necrosis) and embryonic MHC (muscle regeneration) were performed. The masseter muscle of DMD rats, though there is variation among individuals, showed marked myofiber necrosis (Fig. 3A, E) and muscle regeneration (Fig. 3C, F), while the intrinsic tongue muscle showed none of these (Fig. 3B, D–F). These results indicate that, although the masseter and tongue muscles of DMD rats undergo pathological changes that are not seen in WT rats, the changes seen in the tongue muscle are much milder than in the masseter muscle of DMD rats.

To investigate the factors that are responsible for macroglossia seen in DMD rats, we measured the myofiber diameters of the intrinsic tongue muscle of WT and DMD rats at 6 months old. The myofiber diameters of DMD rats were larger than those of WT rats (Fig. 3G, H; \( p < 0.01 \)) by the Wilcoxon rank sum test). The ratio of myofibers with central nuclei was significantly higher in DMD rats than in WT rats, indicating that muscle regeneration had occurred in the past (Fig. 3I). In addition, quantitative analysis of Masson’s trichrome-stained sections as shown in Fig. 2 reveals that the fibrotic area (myofiber necrosis) and embryonic MHC (muscle regeneration) were performed. The masseter muscle of DMD rats, though there is variation among individuals, showed marked myofiber necrosis (Fig. 3A, E) and muscle regeneration (Fig. 3C, F), while the intrinsic tongue muscle showed none of these (Fig. 3B, D–F). These results indicate that, although the masseter and tongue muscles of DMD rats undergo pathological changes that are not seen in WT rats, the changes seen in the tongue muscle are much milder than in the masseter muscle of DMD rats.

Hypertrophy of myofibers occurs by two mechanisms: increased protein synthesis through activation of the PI3K-Akt-mTOR pathway and the addition of new myonuclei by fusion of satellite cells [12]. To gain more insight into the mechanism of hypertrophy of myofibers observed in the tongue muscle of DMD rats, we investigated the amount and phosphorylation of molecules that act in the downstream of PI3K-Akt-mTOR pathway as well as the number of nuclei per myofiber in 6-month-old tongue muscle. No differences were observed between WT and DMD rats in the amount and phosphorylation of S6K, S6, and 4E-BP1 (Fig. 3K–R; Supplementary Fig. 3). Neither the expression nor the phosphorylation of Foxo1, which is thought to be involved in protein degradation, could be detected (data not shown). In addition, no significant differences were observed in the transcription of two skeletal muscle-specific ubiquitin ligase genes, Atrogin 1 and MuRF1, which are involved in protein degradation [2] (Fig. 3S). The phosphorylation of P65 and P38 was increased in WT and DMD rats (Fig. 3S).

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Fig. 3 (See legend on previous page.)
muscles at 9 months old. p16 and p19 were markedly upregulated in the masseter muscle of DMD rats compared to WT rats (Fig. 4A, B). On the other hand, both p16 and p19 were slightly upregulated in the tongue muscle of DMD rats but the degrees were significantly lower than those in the masseter muscle (Fig. 4A, B). p21 transcription was significantly upregulated only in the masseter muscle of DMD rats (Fig. 4C), and there was no difference in the p53 transcription in both the muscles between WT and DMD rats (Fig. 4D). There was no significant difference in the number of Pax7-positive satellite cells in the sections, although there was a large variation between individuals (Fig. 4E). On the other hand, when cultured in vitro for 4 days, the number of Pax7- or MyoD-positive satellite cells derived from the masseter muscle of DMD rats was significantly lower than that of WT rats (Fig. 4F, G, Supplementary Fig. 4A and B), while the number of satellite cells derived from the tongue muscle of DMD rats was only slightly lower than that of WT rats (Fig. 4F, G, Supplementary Fig. 4A and B). These results indicate that cellular senescence, possibly through p16- and p19-dependent pathway, occurs extensively in the masseter muscle of DMD rats as was seen in their hindlimb muscles [33], while it is significantly less severe in the tongue muscle.

The above results suggest that the dystrophic changes are less advanced in the tongue muscle of DMD rats. Utrophin, known as dystrophin-related protein, had been shown to compensate for the function of dystrophin [7, 13, 20, 26]. In a mouse model of DMD, mdx mouse, the skeletal muscle phenotype is milder than that seen in DMD patients [20], and this had been once thought to be due to an upregulation of utrophin [20]. Thus, we examined the Utrn transcription in the masseter and tongue muscles of WT and DMD rats at 9 months old. In the masseter muscle, Utrn transcription was significantly upregulated by the absence of dystrophin (Fig. 4H). Interestingly, in the tongue muscle, Utrn transcription was also upregulated but its basal expression in the presence of dystrophin (WT rats) was comparable to that in the masseter muscle of DMD rats (Fig. 4H). This result suggests that the tongue muscle is intrinsically expressing a higher level of Utrn transcription even in the presence of dystrophin, and this might be related to the resistance of

Fig. 4 Quantitative analyses of gene expression and satellite cells in muscles of WT and DMD rats. A p16, B p19, C p21, and D p53 transcriptions in the masseter and tongue muscle at 9 months old. The data are expressed as mean ± SE. * and **, p < 0.05 and p < 0.01 by unpaired Student’s t-test, respectively (WT, n = 5; DMD, n = 6). E Number of Pax7-positive satellite cells in the sections at 9 months old. The data are expressed as mean ± SE. There was no significant difference among all groups by Tukey–Kramer test (WT, n = 5; DMD, n = 6). F, G Relative number of Pax7- and MyoD-positive satellite cells from 9-month-old rats after 4-day culture in vitro. The data are expressed as mean ± SE. * and **, p < 0.05 and p < 0.01 by unpaired Student’s t-test, respectively (WT, n = 3; DMD, n = 3). H Utrn transcription in the masseter and tongue muscle at 9 months old. The data are expressed as mean ± SE. *, p < 0.05 by unpaired Student’s t-test (WT, n = 5; DMD, n = 6)
this tissue against degenerative changes in the absence of dystrophin.

Discussion

In the present study, we demonstrated that the tongue of DMD rats shows macroglossia and degenerative changes were observed transiently only in the early period of life. On the other hand, chronic degenerative and necrotic changes were seen in the masseter muscle of DMD rats.

Cellular senescence is caused by oxidative stress due to chronic inflammation [6]. We previously reported progressive myofiber degeneration and necrosis, along with cellular senescence with increased transcription of p16 and p19, in the tibialis anterior (TA) and diaphragm of DMD rats [33]. p16 transcript was mainly present in satellite cells and mesenchymal progenitor cells, and their proliferative activity was severely suppressed in vitro. The phenotype of the masseter muscle of DMD rats seen in this study is similar to our previous reports; the masseter muscle showed progressive myofiber degeneration and necrosis, and accordingly, p16 and p19 transcription was upregulated. In contrast, the transcription of p16 and p19 in the tongue muscle of DMD rats was much lower than those in the masseter muscle, and the proliferative activity of the satellite cells was relatively well-preserved in vitro. This is in agreement with the histological observation that myofiber necrosis and inflammation were not observed in the tongue muscle of DMD rats, suggesting the absence of oxidative stress that could induce cellular senescence.

A similar phenotype to the tongue muscle of DMD rats had been observed in the hindlimb muscle of mdx mice [10, 19, 35]. In the tongue muscle of DMD rats, regeneration of myofibers was transiently observed at 1 month old, and at later periods of their life, myofibers with central nuclei persisted without any degenerative changes. In the skeletal muscle of mdx mice, necrotic regeneration is observed only up to 1 to 2 months of age, and regenerated myofibers with central nuclei remain for a long time [10, 19, 35]. At present, it is not clear why myofiber necrosis occurs only transiently in mdx mice, but it has been suggested that one reason is that compensatory increases in utrophin, a functional homolog of dystrophin, may replace the function of dystrophin [20]. In the present study, the Utrn transcript was higher in the tongue muscle of WT rats than in their masseter muscle. Therefore, one of the possibilities is that some mechanism may be responsible for the increased transcription of Utrn in the tongue muscle of the rat, and myofiber necrosis may not occur even under dystrophin deficiency. However, this study only showed differences in transcription levels of utrophin by quantitative RT-PCR, and immunohistochemical analysis to determine the localization, as well as quantitative analysis of protein levels by western blotting, is needed to draw a precise conclusion.

The degree of skeletal muscle regeneration after injury varies depending on the body part; it has been reported that the diameter of myofibers and the degree of fibrosis in masticatory muscles of 100-day-old mdx mice vary according to the muscles [31]. In addition, Yoshioka et al. reported that when injury and subsequent regeneration of skeletal muscle were induced by cardiotoxin injection in WT and mdx mice, there were differences in the rate of regeneration and the degree of increase in muscle weight after regeneration [40]. In particular, the masseter muscle had a longer time to recover its muscle weight after injury than the TA muscle, and no hypertrophy of regenerated myofibers occurred [40]. They suggested that these differences were due to a distinct nature of the satellite cells present in each skeletal muscle. Regarding this, the origin of the satellite cells varies among different muscles [27]. The tongue muscle including satellite cells is thought to be composed of mixed mesodermal origins [23, 27] but detailed convincing lineage tracing experiments of these muscles are not available. On the other hand, the satellite cells of the masseter muscle are derived from the first and second pharyngeal arches of the cranial paraxial mesoderm with contributions from the splanchnic mesoderm [27]. At present, it is unclear how the different origins of these satellite cells ultimately affect the nature of the skeletal muscle from which they arise. However, in the multiple types of myopathies, including DMD, it has been suggested that differences in the origin of satellite cells may contribute to the distinct pathological susceptibility among different muscles [27]. Thus, the phenotypic difference between the masseter and tongue muscles observed in DMD rats may be due to the difference in the nature of the satellite cells. However, we cannot exclude the possibility that the differences in dystrophic changes in each muscle are simply due to differences in the mechanical load on each muscle. In this regard, studies that verify differences in muscle loading in the rat are required.

The tongue muscle has a unique structural feature among skeletal muscles in that it is covered by epithelium, and its development is indirectly influenced by epithelium [14, 18]. During development, the epithelium-derived sonic hedgehog (shh) acts on cranial neural crest cells (CNCC), which constitute the mesenchyme [14]. When the hedgehog signal in CNCC is defective, migration of myoblasts from the occipital somite is perturbed, resulting in dysplasia of the tongue. Although the involvement of the epithelium in the adult tongue has not been reported, it is possible that factors derived from the epithelium act directly or indirectly via mesenchyme on the tongue muscle, which may be protective.
against degenerative changes of tongue myofibers caused by dystrophin deficiency. This point awaits further investigation.

Macroglossia has been reported to occur not only in human DMD but also in canine X-linked muscular dystrophy (CXMD) [30]. In human DMD, it is known that macroglossia interferes with tongue function [36, 37], but no detailed histological study has been performed. On the other hand, it has been suggested that hypertrophy of myofibers is the cause of macroglossia in CXMD [30]. This is consistent with the results of the present analysis in DMD rats. The mechanisms of skeletal muscle hypertrophy include increased protein synthesis, decreased degradation, and increased number of myonuclei supplied to myofibers by satellite cells [12]. It has been reported that some regenerated myofibers in mdx mice and golden retriever muscular dystrophy (GRMD) dogs show hypertrophy [4, 8, 15, 16, 24, 29], and one of the mechanisms is the vigorous activation of the Akt-mTOR pathway during the period before the occurrence of myofiber necrosis, which results in hypertrophy of myofibers leading to a subsequent reduced severity of pathology [25]. Since the mechanism of myofiber hypertrophy in the tongue has not been analyzed in depth in any of the species reported [28, 30, 37], we attempted to clarify the mechanism of myofiber hypertrophy in the tongue muscle of the rat DMD in the present study. As a result, the signaling molecules involved in the regulation of protein synthesis and degradation were not significantly altered compared to WT rats, at least at the age of 6 months examined in this study. However, this does not necessarily exclude the possibility that the Akt-mTOR pathway is involved in the hypertrophy of tongue myofibers in DMD rats, and it still remains possible that the transient activation of this pathway at a younger age (before 6 months old) had led to the hypertrophy of muscle fibers. On the other hand, although not statistically significant, the number of nuclei per myofiber showed an increasing trend, suggesting that a supply of new myonuclei by satellite cells may have occurred. This is supported by the present result that myogenicity of satellite cells derived from the tongue muscle of DMD rats was relatively preserved in vitro, even at the age of 9 months, unlike those from the masseter muscle.

The limitation of this study is that it is still unclear whether the macroglossia in DMD rats influences tongue function, and it is unknown whether the decreased food intake seen in DMD rats is attributed to macroglossia. Furthermore, it is also possible that the decreased food intake seen in the DMD rats is caused by general fatigue such as respiratory problems or cardiomyopathy. In fact, the DMD rats show extensive fibrosis in their heart [22, 33] and similar distributions and progression of heart involvement to those of patients with DMD [32]. Regarding these points, it is necessary to evaluate the tongue function experimentally and to elucidate the exact cause of decreased food intake in DMD rats in the future.

Conclusions
We demonstrated that, unlike the masseter and hindlimb muscles, degenerative/necrotic changes of myofibers, decreased satellite cell proliferative activity, and increased p16 and p19 expressions were seldomly observed in the tongue muscle of DMD rats. Furthermore, the tongue muscle of the rat showed higher Ultrn transcription than the masseter muscle, suggesting that this compensatory Ultrn transcription might protect the tongue myofibers from degenerative/necrotic changes under the lack of dystrophin. Unraveling the mechanism involved in less advanced dystrophic changes in the tongue muscle in DMD rats would benefit to establish an effective treatment for DMD.

Abbreviations
ANOVA: Analysis of variance; CNCC: Cranial neural crest cells; CSPG4: Chondroitin sulfate proteoglycan 4; CXMD: Canine X-linked muscular dystrophy; DMD: Duchenne muscular dystrophy; DMD rat: A rat model of DMD; DSHB: Developmental Studies Hybridoma Bank; EDTA: Ethylenediaminetetraacetic acid; eMHC: Embryonic myosin heavy chain; HE: Hematoxylin–eosin; HPRT: Hypoxanthine-guanine phosphoribosyltransferase; HRP: Horseradish peroxidase; NICHD: National Institute of Child Health and Human Development; NDS: Normal donkey serum; NGS: Normal goat serum; PBS: Phosphate-buffered saline; PFA: Paraformaldehyde; RIPA: Radioimmunoprecipitation assay; SDS-PAGE: Sodium dodecyl sulfate–polyacrylamide gel electrophoresis; Shh: Sonic hedgehog; TA: Tibialis anterior; WT: Wild-type; X: Male WT; X(dy): Male DMD.

Supplementary Information
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Additional file 1: Supplementary Fig. 1. HE-stained sections of the extrinsic tongue muscle of DMD rats. At 3- and 8-month-old, regenerative myofibers with centralized nuclei and basophilic cytoplasm were observed in some genioglossus muscles.

Additional file 2: Supplementary Fig. 2. Immunohistochemical analyses of vascular endothelial (CD31) and mesenchymal progenitor (CSPG4) cells. Representative immunohistochemistry of CD31 (red), and CSPG4 (red) and laminin (green) in the tongue muscle of WT and DMD rats at 6-month-old. Nuclei were stained with Hoechst 33342 (blue). Graphed data are quantitative analysis of CD31- and CSPG4-positive areas. The data are expressed as mean±SE (CD31, WT (n=5) and DMD (n=6); CSPG4, WT (n=3) and DMD (n=3)).

Additional file 3: Supplementary Fig. 3. Western blot of molecules involved downstream of Akt-mTOR pathway in the tongue muscle at 6-month-old. Two bands (doublet) were detected by the anti-4E-BP1 antibody. The upper (p-4E-BP1 (high)) and lower (p-4E-BP1 (low)) bands correspond to highly phosphorylated 4E-BP1 and non- or lower-phosphorylated 4E-BP1, respectively. α-tubulin was used as an internal control. p-Akt (S473), Akt phosphorylated at Ser473; p-S6K (T389), S6 kinase phosphorylated at Thr389; p-S6 (S240/244), S6 ribosomal protein phosphorylated at Ser240 and Ser244; n, the number of samples.

Additional file 4: Supplementary Fig. 4. Representative immunocytochemistry of satellite cells. The cultured cells were immunostained with...
anti-Pax7 (A) and anti-MyoD (B) on day 4 of culture. Nuclei were stained with Hoechst 33258. Arrowheads (white) indicate positive cells.

Authors’ contributions
KY conducted the experiments, analyzed the data, prepared the figures, and wrote the manuscript; YT, MI, RO, and HN conducted the experiments; SK prepared tissue sections and conducted the experiments; JC, FH, SIT, TM, and KU edited the manuscript. The authors read and approved the final manuscript.

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Declarations

Ethics approval and consent to participate
All animal experiments performed in this study were in accordance with the Guide for the Care and Use of Laboratory Animals of the University of Tokyo and were approved by the Institutional Animal Care and Use Committee of the University of Tokyo (P18-125). All studies were conducted in compliance with the Helsinki Declaration.

Competing interests
The authors declare that they have no competing interests.

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