Loss of skeletal muscle strength by ablation of the sarcoplasmic reticulum protein JP45

Osvaldo Delbono*, Jinyu Xia†, Susan Treves‡, Zhong-Min Wang*, Ramon Jimenez-Moreno*, Anthony M. Payne*, Maria Laura Messi*, Alexandre Brigiuet†, Florian Schaeer†, Miyuki Nishi®, Hiroshi Takeshima®, and Francesco Zorzato**

Departments of Anaesthesia and Research, Basel University Hospital, Hebelstrasse 20, 4031 Basel, Switzerland; *Departments of Physiology, Pharmacology, and Internal Medicine; Gerontology; and Geriatric Medicine, Wake Forest University School of Medicine, Winston-Salem, NC 27157; †Santhera Pharmaceuticals, CH-4410 Liestal, Switzerland; ‡Department of Biological Chemistry, Graduate School of Pharmaceutical Sciences, Kyoto University, Kyoto 606-8501, Japan; and ¶Department of Experimental and Diagnostic Medicine, General Pathology Section, University of Ferrara, Via Borsari 46, 44100 Ferrara, Italy

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Skeletal muscle constitutes ∼40% of the human body mass, and alterations in muscle mass and strength may result in physical disability. Therefore, the elucidation of the factors responsible for muscle force development is of paramount importance. Excitation–contraction coupling (ECC) is a process during which the skeletal muscle surface membrane is depolarized, causing a transient release of calcium from the sarcoplasmic reticulum that activates the contractile proteins. The ECC machinery is complex, and the functional role of many of its protein components remains elusive. This study demonstrates that deletion of the gene encoding the sarcoplasmic reticulum protein JP45 results in decreased muscle strength in young mice. Specifically, this loss of muscle strength in JP45 knockout mice is caused by decreased functional expression of the voltage-dependent Ca2+ channel Ca2,1, which is the molecule that couples membrane depolarization and calcium release from the sarcoplasmic reticulum. These results point to JP45 as one of the molecules involved in the development or maintenance of skeletal muscle strength.

Results and Discussion

Phenotype of the JP45 KO Mice. The absence of JP45 in the skeletal muscle of the mutant mice was confirmed by Western blot analysis of SR membranes isolated from JP45 KO and wild-type mice (Fig. 1D). The ablation of JP45 is not associated with major alterations in other SR proteins (Fig. 1E). Homozygous JP45 KO mice did not exhibit either a lethal phenotype or defects in embryonic or postnatal development. To look for a skeletal muscle phenotype, we analyzed the animals using a voluntary running wheel setup. The mice had free access to the wheel at any time of day. This experimental approach avoids potential problems linked to the effect of circadian rhythm on animal activity and/or animal compliance in performing nonvoluntary motor activity. We assessed spontaneous dark-phase (17.00–5.00 h) motor activity of 3-month-old wild-type and JP45 KO mice (Fig. 2). In the first week, the total running distance of the wild-type mice was approximately double that of the JP45 KO mice. Two weeks of training improved skeletal muscle performance in both groups. After 3 weeks of training, the total running distance increased 2- to 3-fold until it plateaued at ∼9–10 kilometers per night for both sets of mice, although the total running distance remained significantly lower in the JP45


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†To whom correspondence should be addressed. E-mail: zor@unife.it.

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KO mice (Mann–Whitney test, two-tailed, *P* < 0.05). We also found that the median cruise speed of the JP45 KO mice was significantly lower than that of the wild-type mice (Mann–Whitney test, two-tailed, *P* < 0.05).

**Mechanical Properties of Intact Skeletal Muscle from Wild-Type and JP45 KO Mice.** The reduced speed and total running distance among the JP45 KO mice may be associated with decreased muscle strength resulting from alterations in their ECC machinery caused by the lack of JP45. To investigate this possibility, we studied the mechanical properties of intact extensor digitorum longus (EDL) and soleus muscles from wild-type and JP45 KO mice (Fig. 3A–D). Analysis of muscle contraction revealed significant differences in JP45 KO mice. Their EDL and soleus muscles exhibited decreased twitch and maximal tetanic absolute force compared with wild-type mice, whereas their contraction and relaxation kinetics did not differ significantly (Table 1). The reduced absolute force observed in their skeletal muscles cannot be attributed to loss of contractile protein content, because the wet weight of their EDL and soleus muscles did not significantly differ from those of wild-type mice. The myofibril area in EDL and soleus muscles also did not differ significantly between JP45 KO and wild-type mice [supporting information (SI) Fig. 6]. The muscle specific force (maximal tetanic force normalized to muscle cross-sectional area) of EDL and soleus muscles from JP45 KO mice was also significantly impaired. EDL maximal tetani recorded in JP45 KO (287 ± 38.6 mN·mm⁻², mean ± SD, *n* = 22) are ~25% lower than those recorded in wild-type mice (381 ± 46.7 mN·mm⁻², *n* = 13, *P* < 0.05). A similar decrease in maximal tetanic force generation was observed in solei from JP45 KO (244 ± 41.6, *n* = 22) compared with wild-type (301 ± 46.7, *n* = 13) mice (*P* < 0.05). Specific tetanic force decreased in single, intact flexor digitorum brevis (FDB) muscle fibers to an extent similar to that in EDL and soleus whole-muscle contraction, but the calcium sensitivity of the contractile proteins, tested in the same fiber after chemical skinning, did not differ significantly in JP45 KO compared with wild-type mice (SI Fig. 7). The decrease in force output is not due to fast-to-slow fiber transition because we observed no changes in the expression of myosin heavy chain (MHC) isoforms in EDL (E) and soleus (F) muscles. EDL from JP45 KO and wild-type mice expresses predominantly MHC IIb, whereas soleus expresses mainly MHC I and IIa. No gross alterations in the overall microscopic structure were observed in JP45 KO mice (data not shown).

![KO mice](image)

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**Mechanical Properties of Intact Skeletal Muscle from Wild-Type and JP45 KO Mice.** The reduced speed and total running distance among the JP45 KO mice may be associated with decreased muscle strength resulting from alterations in their ECC machinery caused by the lack of JP45. To investigate this possibility, we studied the mechanical properties of intact extensor digitorum longus (EDL) and soleus muscles from wild-type and JP45 KO mice (Fig. 3A–D). Analysis of muscle contraction revealed significant differences in JP45 KO mice. Their EDL and soleus muscles exhibited decreased twitch and maximal tetanic absolute force compared with wild-type mice, whereas their contraction and relaxation kinetics did not differ significantly (Table 1). The reduced absolute force observed in their skeletal muscles cannot be attributed to loss of contractile protein content, because the wet weight of their EDL and soleus muscles did not significantly differ from those of wild-type mice. The myofibril area in EDL and soleus muscles also did not differ significantly between JP45 KO and wild-type mice [supporting information (SI) Fig. 6]. The muscle specific force (maximal tetanic force normalized to muscle cross-sectional area) of EDL and soleus muscles from JP45 KO mice was also significantly impaired. EDL maximal tetani recorded in JP45 KO (287 ± 38.6 mN·mm⁻², mean ± SD, *n* = 22) are ~25% lower than those recorded in wild-type mice (381 ± 46.7 mN·mm⁻², *n* = 13, *P* < 0.05). A similar decrease in maximal tetanic force generation was observed in solei from JP45 KO (244 ± 41.6, *n* = 22) compared with wild-type (301 ± 46.7, *n* = 13) mice (*P* < 0.05). Specific tetanic force decreased in single, intact flexor digitorum brevis (FDB) muscle fibers to an extent similar to that in EDL and soleus whole-muscle contraction, but the calcium sensitivity of the contractile proteins, tested in the same fiber after chemical skinning, did not differ significantly in JP45 KO compared with wild-type mice (SI Fig. 7). The decrease in force output is not due to fast-to-slow fiber transition because we observed no changes in the expression of myosin heavy chain (MHC) isoforms in EDL (E) and soleus (F) muscles. EDL from JP45 KO and wild-type mice expresses predominantly MHC IIb, whereas soleus expresses mainly MHC I and IIa. No gross alterations in the overall microscopic structure were observed in JP45 KO mice (data not shown).

**Fig. 2.** Spontaneous activity of JP45 KO mice. Shown is dark-phase spontaneous activity in 12-week-old wild-type (squares) and JP45 KO (triangles) mice individually housed in cages equipped with a running wheel. Data points are expressed as mean ± SEM (*n* = 10–13 mice). *P* < 0.05 (Mann–Whitney two-tailed test).
Ablation of JP45 Decreases Cav1.1 Density in Sarcotubular Membranes. Because the decreased muscle strength observed in JP45 KO mice cannot be explained by loss of contractile proteins or changes in MHC isoforms, we thought that it might be linked to alterations of the ECC mechanism. We reasoned that the site of such alterations might be the supramolecular complex made up of Cav1.1 and RyR. Cav1.1 is the voltage sensor, which generates charge movement that is transmitted to the RyR to initiate the calcium release necessary to activate muscle contraction. Because JP45 forms a complex with Cav1.1 (30), we investigated whether the decreased muscle function caused by JP45 ablation was related to alterations in the expression of Cav1.1. Fig. 4 shows that, indeed, chronic JP45 depletion is accompanied by a decrease in the density of Cav1.1 in the sarcotubular membranes, as indicated by radioligand binding assays with the high-affinity Cav1.1 probe [3H] PN200-110 (Fig. 4A). Scatchard analysis of [3H] PN200-110 isothermal equilibrium binding to isolated sarcotubular membranes from JP45 KO and wild-type mice shows that the former have a significant decrease in PN200-110 $B_{\text{max}}$ with no alteration of its dissociation constant ($B_{\text{max}}$, pmol/mg of protein = 5.8 ± 0.40 vs. 7.9 ± 0.30, mean ± SD (n = 11 for JP45 KO and wild-type, respectively); $K_{d}$ = 1.3 ± 0.20 vs. 1.1 ± 0.10 nM, mean ± SD (n = 11 for JP45 KO and wild-type mice, respectively)]. The decrease of PN200-110 binding in the sarcotubular membranes
The decreased Ca_{1.1} membrane density in the skeletal muscle from JP45 KO mice causes modifications in the fiber’s electrophysiological properties: in fact, we found a reduction in the asymmetric capacitive current, which results in a 40% decrease in maximal Ca_{1.1} gating charge in intact FDB fibers (Fig. 4D) without significantly affecting charge movement half-activation potential (Table 2). These results together with previous studies on JP45 overexpression (30, 34) indicate that the stoichiometry of JP45–Ca_{1.1} subunit exhibited by mature skeletal muscle is crucial for maintaining the molecular and functional expression of the Ca_{1.1} subunit (31).

Because the Ca_{1.1} charge movement is the activating signal for RyR1-mediated calcium release from the SR, we performed experiments to evaluate whether depolarization-induced calcium release from the SR of JP45 KO mice is also affected. We found that the peak depolarization-induced calcium release is significantly decreased in JP45 KO fibers (Fig. 5A). The flux amplitudes J_r (μM·ms^{-1}) and J_s (μM·ms^{-1}) were 181 ± 13 and 115 ± 18 for fibers from wild-type mice and 108 ± 17 and 68 ± 7.3 for fibers from JP45 KO mice, respectively (P < 0.05). The decreased depolarization-induced calcium release is apparently not due to reduced RyR1 calcium-release channel density because Scatchard analysis of [H]ryanodine binding to heavy SR fractions revealed no changes (B_{max} values were 11.60 ± 2.0 and 13.51 ± 1.9 pmol/mg of protein for wild-type and JP45 KO mice, respectively) (Fig. 5B). The lack of significant differences in their resting calcium concentrations indicates that myoplasmic calcium-buffer capacity is preserved in JP45 KO mice (Fig. 5C). A nonsignificant difference in Ca ATPase activity in the light SR fraction supports this conclusion (SI Fig. 9).

We also investigated 4-chloro-m-cresol-induced force output in single, intact FDB fibers to finely assess maximal SR-releasable calcium (35). We found no differences in force generation between wild-type and JP45 KO muscle fibers induced by the RyR1 agonist 4-chloro-m-cresol (Fig. 5D). On the basis of 4-chloro-m-cresol-induced force experiments, we are confident that the reduction in voltage-dependent calcium release is not a result of SR calcium-store depletion but rather of altered Ca_{1.1} expression.

The most important and novel finding of this report is the demonstration that alterations to the molecular composition of the ECC machinery resulting from JP45 depletion are sufficient to induce a 20–30% decrease in skeletal muscle strength in young animals. In addition, this study elucidates the underlying molecular mechanism(s) responsible for the decay of muscle strength in JP45 KO mice. It unambiguously shows that the decrease in muscle strength is not due to changes in muscle fiber types or contractile protein properties but to alterations of the signal transduction machinery between Ca_{1.1} and RyR1 at the T tubule/SR junction. We believe that the crucial switch is the activating calcium signal from the SR, which is then delivered to the contractile proteins. Altering the components of this switch may affect global muscle function. This conclusion is supported by the data demonstrating that reduced membrane depolarization due to impaired functional expression of Ca_{1.1} reduces calcium release. The molecular basis of the reduction in Ca_{1.1} functional expression is likely linked to protein–protein interactions occurring within the EC-coupling macromolecular complex. Membrane targeting of Ca_{1.1} depends on amino acid sequences contained within the COOH terminal domain and in the I–II loop of the Ca_{1.1} (20). JP45 interacts with both Ca_{1.1} domains involved in membrane targeting (31), and the lack of these interactions may interfere with either proper Ca_{1.1} membrane targeting or the stabilization of the Ca_{1.1} complex in the
Differences are not statistically significant.

Table 2. Best-fitting parameters describing the voltage dependence of charge movement and intracellular Ca$^{2+}$ release in FDB fibers from wild-type and JP45 KO mice

<table>
<thead>
<tr>
<th>Mice</th>
<th>$Q_{\text{max}}$ μC/μF</th>
<th>$V_{1/2Q}$, mV</th>
<th>$K$</th>
<th>$\Delta F/F_{\text{max}}$</th>
<th>$V_{1/2F}$, mV</th>
<th>$K$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>30.5 ± 3.4</td>
<td>7.6 ± 0.92</td>
<td>14.2 ± 1.8</td>
<td>17.8 ± 2.6$^*$</td>
<td>6.4 ± 0.75</td>
<td>12.0 ± 1.5</td>
</tr>
<tr>
<td>JP45 KO</td>
<td>0.61 ± 0.08</td>
<td>12.9 ± 0.19</td>
<td>15.4 ± 1.8</td>
<td>0.39 ± 0.06$^*$</td>
<td>13.0 ± 0.15</td>
<td>16.3 ± 1.9</td>
</tr>
</tbody>
</table>

Values are mean ± SEM. $^*$, $P < 0.01$ (statistically significant differences between wild type and JP45 KO).

$^*_n$ = 15 (wild type) and 16 (JP45 KO) fibers.

$^*_{1n}$ = 17 (wild type) and 20 (JP45 KO) fibers.

T tubular membrane. In conclusion, this study provides new insights into some mechanism(s) responsible for diminished muscle strength.

**Experimental Procedures**

**Generation of JP45 KO Mice.** JP45 KO mice were generated as previously described (27). A mouse genomic library (from Stratagene) was screened with a cDNA probe to isolate genomic clones encompassing the 5’-end of the JP45 gene. A targeting vector was constructed by using a 5’-end JP45 genomic clone, Neo-resistance, and the diphtheria toxin gene DT-A as positive-selection and negative-selection cassettes, respectively (Fig. 1A).

**Ablation of JP45 reduced maximal voltage-dependent SR Ca$^{2+}$ release.** (A Upper) SR Ca$^{2+}$ release in FDB fibers from wild-type ($n = 17$) and JP45 KO ($n = 20$) mice. Data points (mean ± SEM) were fitted to a Boltzmann equation. (A Lower) OGB-SN transients in FDB fibers from wild-type and JP45 KO mice in response to various command pulses corresponding to the steepest part of the fluorescence/membrane voltage relationship. (B) [3H]Ryanodine binding to heavy SR fractions from wild-type and JP45 KO mice. Points represent the mean ± SD of five to eight determinations in two different R4 fraction preparations. (C) Fura-2 resting Ca$^{2+}$ concentration measurements in FDB fibers from wild-type ($n = 72$ fibers) and JP45 KO ($n = 71$) mice did not differ significantly. (D Upper) Typical contractures in response to 1 mM 4-CmC in FDB fibers from wild-type ($n = 8$) and JP45 KO ($n = 11$) mice. (D Lower) Differences are not statistically significant.

A Sall-linearized vector was used to transfect J1 mouse ES cells, and 140 ES clones carrying a homologous recombination were identified by Southern blot screening (Fig. 1B). JP45KO mice were backcrossed three times in a C57BL6 background and then intercrossed to obtain JP45KO mice. Genotyping was carried out by PCR using the following primers: JP45F2, 5’TAA AGA CAG AGA CCA CAT CCT CCC-3’. JP45R4, 5’-GAC AAG GGG TGT GGA TGA GGC-3’ (Fig. 1C).

**DNA and RNA Manipulation.** Nucleic acid manipulation was carried out as previously described (27, 36).

**Microsomal Preparation and Biochemical Analysis.** The isolation of SR membrane fractions (37), SDS/PAGE, and Western blot analysis were carried out as described by Anderson et al. (30). [3H]PN200-110 and [3H]Ryanodine binding was carried out according to Anderson et al. (38). Curve fitting was performed by using Prism4 software (GraphPad). MHC band separation was performed as described by Talmadge and Roy (39), and densitometry was analyzed with the NIH ImageJ 1.36 software.

**In Vivo and in Vitro Muscle Strength Assessment.** Animals were individually housed in cages equipped with a running wheel carrying a magnet. Wheel revolutions were registered by a reed sensor connected to an I-7033D Digital-Input module (Spectra), and the revolution counters were read by a standard laptop computer via an I-7520 RS-485-to-RS-232 interface converter (Spectra). Digitized signals were processed by the “mouse running” software developed at Santhera Pharmaceuticals. To test force in *vitro*, EDL and soleus muscles were dissected and mounted into a muscle testing setup (Heidelberg Scientific Instruments). Muscle force was digitized at 4 kHz by using an AD Instruments converter and stimulated with 15-V pulses for 0.5 ms. EDL tetanus was recorded in response to 400-ms pulses at 10–120 Hz, whereas, for soleus, 1,100-ms pulses at 10–150 Hz were applied. Specific force was normalized to the muscle cross-sectional area [CSA = wet weight (mg)/length (mm)] × 1.06 (density mg/mm$^3$)] (40).

**Cell Electrophysiology and Optical Recordings.** FDB fibers from wild-type and JP45 KO were enzymatically dissociated, plated, and recorded as previously described (41). For whole-cell patch clamp, the composition of the pipette solution was 140 mM Cs-aspartate, 5 mM Mg-aspartate, 10 mM CsEGTA [ethylene glycol-bis(2-aminoethyl ether)-N,N,N’,N’-tetraacetic acid], and 10 mM Hepes [N-(2-hydroxyethyl)piperazine-N’-(2-ethanesulfonic acid)]. The pH was adjusted to 7.4 with CsOH. The external solution contained 145 mM tetraethylammonium hydroxide (TEA)-Cl, 10 mM CaCl$_2$, 10 mM Hepes, and 0.001 mM tetrodotoxin. Solution pH was adjusted to 7.4 with TEA-OH.

For charge-movement recording, calcium current was blocked by adding 0.5 Cd$^{2+}$ plus 0.3 La$^{3+}$ to the external solution (8, 32). We recorded the charge movement corresponding to gating of the Ca$_{1.1}$ channel. To this end, we used a protocol consisting of a 2-s prepulse to $-30$ mV and a subsequent 5-s repolarization.
to a pedestal potential of −50 mV followed by a 25-ms depolarization from −50 mV to 50 mV with 10-mV intervals (8, 31). Intramembrane charge movements were calculated as the integral of the current in response to depolarizing pulses and are expressed per membrane capacitance (coulombs per farad). For analysis of the relationship between charge movement and membrane voltage, data points were fitted to a Boltzmann equation of the form $Q_{on} = Q_{max}[1 + \exp(V_{1/2} - V_m)/K]$, where $Q_{max}$ is the maximum charge, $V_m$ is the membrane potential, $V_{1/2}$ is the charge movement half-activation potential, and $K$ is the steepness of the curve. The best-fitting parameters are included in Table 1. For intracellular Ca$^{2+}$ recording, the fiber-holding potential was −80 mV, and the duration of the command pulses was 40 ms to better display the steady component of the SR release function. The pipette solution contained 20 mM Cs$_2$EGTA and 500 μM Oregon Green Bapta (OGB)-5N (Invitrogen). OGB-5N transients were recorded by using a laser scanning confocal microscope (Radiance 2100; Bio-Rad/Zeiss) in linescan mode. Images were converted into fluorescence intensity. SR Ca$^{2+}$ release flux was calculated by using a single-compartment model as described (42). The software for these calculations was kindly provided by J. Vergara (University of California, Los Angeles). Resting Ca$^{2+}$ concentration was measured by using fura-2AM as the Ca$^{2+}$-probe in enzymatically dissociated FDB fibers. Ratios of the fluorescence recorded at 380- and 340-nm excitation wavelengths were converted into Ca$^{2+}$ concentration following published procedures (43). Force recordings in single, intact FDB muscle fibers were as previously described (44, 45).

**Statistical Analysis.** Statistical analysis was performed by using Student’s unpaired t test and the Mann–Whitney U test when values were not normally distributed. $P < 0.05$ was considered significant.

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