Restricting calcium currents is required for correct fiber type specification in skeletal muscle

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ABSTRACT

Skeletal muscle excitation-contraction (EC) coupling is independent of calcium influx. In fact, alternative splicing of the voltage-gated calcium channel CaV₁.1 actively suppresses calcium currents in mature muscle. Whether this is necessary for normal development and function of muscle is not known. However, splicing defects that cause aberrant expression of the calcium-conducting developmental CaV₁.1e splice variant correlate with muscle weakness in myotonic dystrophy. Here, we deleted CaV₁.1e (Cacna1s) exon 29 in mice. These mice displayed normal overall motor performance, although grip force and voluntary running were reduced. Continued expression of the developmental CaV₁.1e splice variant in adult mice caused increased calcium influx during EC coupling, altered calcium homeostasis, and spontaneous calcium sparklets in isolated muscle fibers. Contractile force was reduced and endurance enhanced. Key regulators of fiber type specification were dysregulated and the fiber type composition was shifted toward slower fibers. However, oxidative enzyme activity and mitochondrial content declined. These findings indicate that limiting calcium influx during skeletal muscle EC coupling is important for the secondary function of the calcium signal in the activity-dependent regulation of fiber type composition and to prevent muscle disease.

KEY WORDS: Voltage-gated calcium channel, Skeletal muscle excitation-contraction coupling, Muscle fiber type specification, Mouse

INTRODUCTION

Calcium is the principal second messenger regulating skeletal muscle contraction, growth and differentiation. In excitation-contraction (EC) coupling, cytoplasmic calcium levels are rapidly increased in response to action potentials and the magnitude of these calcium signals regulates the force of contraction. In skeletal muscle, voltage-gated calcium channels (CaV₁.1) and calcium release channels (type 1 ryanodine receptors, RyR1) are physically coupled to one another so that voltage-dependent activation of CaV₁.1 can directly activate opening of the RyR1. In mature muscles, calcium is released from the sarcoplasmic reticulum (SR) calcium stores, whereas calcium influx is dispensable for skeletal muscle EC coupling (Melzer et al., 1995). Actually, calcium currents through the major CaV₁.1a splice variant are small and activate slowly, only at strong membrane depolarizations.

Interestingly, during maturation of mammalian skeletal muscles activity-dependent calcium influx is actively suppressed by alternative splicing of CaV₁.1. In fetal muscles, exclusion of exon 29 produces a CaV₁.1e channel variant that conducts sizable L-type calcium currents and activates in parallel to SR calcium release at physiological voltages (Tuluc et al., 2009). However, after birth, the developmental CaV₁.1e splice variant is almost completely replaced by the adult, poorly conducting CaV₁.1a splice variant that includes exon 29 (Flucher and Tuluc, 2011; Tang et al., 2012). Why calcium influx is present in developing muscle but is then curtailed in mature skeletal muscles is not known. Conversely, it remains to be determined whether continued expression of the calcium-conducting CaV₁.1e splice variant alters contractile properties of mature skeletal muscles.

In addition to their primary role in EC coupling, activity-induced calcium signals in skeletal muscle are important for maintaining calcium homeostasis and for the regulation of muscle growth and differentiation. For example, calcium signals regulate the transcription of genes involved in the adaptive response to exercise (Bassel-Duby and Olson, 2006). Therefore, the tight control of calcium influx by alternative splicing of the CaV₁.1 channel is probably important for tuning muscle function to varying activity levels. Conversely, calcium influx through CaV₁.1e channels in mature muscles might be harmful. Abnormal expression of the embryonic, calcium-conducting CaV₁.1e splice variant in myotonic dystrophy type 1 (DM1) patients correlates with their degree of muscle weakness (Tang et al., 2012). Moreover, aberrant splicing of calcium channels and transporters in cultured myotubes from DM1 patients leads to altered intracellular calcium signaling (Santoro et al., 2014), and experimentally induced skipping of exon 29 aggravated the disease phenotype in muscles of a myotonia mouse model (Tang et al., 2012).

To identify the physiological importance of curtailing calcium influx through CaV₁.1 channels in adult skeletal muscle and to reveal a possible involvement of aberrant calcium signaling in DM1, we generated a genetic mouse model in which exon 29 has been permanently deleted. As expected, skeletal muscles of CaV₁.1ΔE₂⁹ knockout mice experienced increased calcium influx during EC coupling and at rest. In addition, their contractile properties were altered, calcium-activated downstream regulators were upregulated, and the fiber type composition was shifted towards slower fiber types. However, mitochondrial content and oxidative enzyme activity were reduced. Together, these findings...
indicate that chronically increased calcium influx through the developmental CaV1.1e isoform has little effect on EC coupling, but disturbs the normal regulation of muscle fiber type composition. Furthermore, the increased calcium influx causes mitochondrial damage and may thus contribute to muscle wasting in DM1. Conversely, these results suggest that, during normal development, limiting L-type calcium currents is important to enable the proper specification of fiber type composition and to protect the muscles from calcium-induced damage.

RESULTS
Selective deletion of CaV1.1 exon 29 prevents the developmental switch from the CaV1.1e to the CaV1.1a isoform
In order to study the importance of the isoform switch from the calcium-conducting developmental CaV1.1e splice variant to the poorly conducting adult CaV1.1a splice variant we generated a mouse model with a constitutive knockout of exon 29 of the CaV1.1 (Cacna1s) gene (Fig. 1A). We reasoned that because the short transcript CaV1.1e is predominant during fetal development, CaV1.1ΔE29 mice would develop normally up to birth, but that the aberrant continuing expression of the high-conductance developmental calcium channel splice variant throughout postnatal development and adult life would reveal any influence of the extra calcium influx on EC coupling and/or other calcium-mediated signaling processes regulating muscle growth and differentiation. Furthermore, the CaV1.1ΔE29 mouse will expose whether aberrant splicing of CaV1.1 is itself sufficient to cause a disease phenotype reminiscent of DM1.

Heterozygous CaV1.1ΔE29 and homozygous CaV1.1ΔE29ΔE29 mice were viable, they developed normally, and home cage activity of CaV1.1ΔE29 mice was not significantly different from that of wild-type siblings (Fig. 1B, Fig. S1A). Expression of the two CaV1.1 transcripts at different developmental stages was analyzed in the predominantly slow oxidative soleus muscle, the predominantly fast glycolytic extensor digitorum longus (EDL) muscle, and the mixed diaphragm muscle. Quantitative RT-PCR demonstrated that wild-type fetal muscles express moderate levels of both splice variants, with a higher proportion of the splice variant lacking exon 29 (CaV1.1e) (Fig. 1C). After birth, wild-type muscles expressed a strong upregulation of the CaV1.1a transcript, whereas expression of the CaV1.1e transcript declined to less than 3% in 16-week-old mice. In muscles of ageing mice (15-18 months), total CaV1.1 transcript levels declined but the overall predominance of the CaV1.1a variant was maintained. As expected, in homozygous CaV1.1ΔE29ΔE29 mice the CaV1.1e transcript was found exclusively. At all developmental stages its expression levels resembled those of total CaV1.1 transcripts in wild-type mice. Western blot analysis confirmed normal expression levels of total CaV1.1 protein in soleus (Fig. 1D). However, in EDL muscle total CaV1.1 protein was reduced. This might, at least in part, reflect a reduced content of triad junctions due to the fiber type shift observed in CaV1.1ΔE29 mice (see below).

Aberrant expression of the developmental CaV1.1e isoform in mature muscles is not sufficient to cause severe myotonic dystrophy symptoms in mice
Because aberrant expression of CaV1.1e in adults has been linked to the DM1 phenotype in mouse and human (Santoro et al., 2014; Tang et al., 2012), we subjected CaV1.1ΔE29 mice at 2 and 8 months of age to a range of behavioral tests to examine different aspects of muscle performance (Fig. 2). A wire hang test was used to assess muscle strength. Endurance was tested by making the mice run on a treadmill at accelerating speed until exhaustion. Overall motor performance was examined with the Rotarod test. In none of these tests was a significant difference in the performance of wild-type, heterozygous CaV1.1ΔE29ΔE29 and homozygous CaV1.1ΔE29ΔE29ΔE29 mice observed. However, directly measuring grip strength revealed that the grip force of the front paws was significantly reduced in homozygous CaV1.1ΔE29ΔE29ΔE29 mice (Fig. 2D). Finally, voluntary running of the mice in a running wheel was recorded over the period of 7 days. Both the distance run and the duration the mice spent running per day were significantly reduced in homozygous CaV1.1ΔE29ΔE29ΔE29 mice compared with wild-type controls (Fig. 2E). These behavioral and functional analyses indicate that aberrant expression of CaV1.1e alters muscle performance without causing severe motor deficits as assessed in tests that revealed the disease phenotype in other DM mouse models (Gomes-Pereira et al., 2011). Also, histological staining of muscle sections did not reveal an increase in centrally located nuclei in CaV1.1ΔE29ΔE29ΔE29 muscles (Fig. S1B).

Aberrant expression of the developmental CaV1.1e isoform in mature muscles alters calcium signals during EC coupling and in resting muscle fibers
To clarify the cellular mechanisms underlying the altered muscle properties of CaV1.1ΔE29ΔE29 mice we analyzed calcium currents and cytoplasmic calcium signals directly in isolated flexor digitorum brevis (FDB) muscle fibers using several experimental paradigms. First, combined patch-clamp and cytoplasmic calcium recording was performed in FDB fibers loaded with the fluorescent calcium indicator Rhod-2. In line with the current properties of the CaV1.1e splice variant previously determined in reconstituted dysgenic myotubes (Tuluc et al., 2009), CaV1.1ΔE29ΔE29ΔE29 FDB fibers displayed sizable calcium currents starting at test potentials of −30 mV (Fig. 4A). Under the same conditions (1.8 mM extracellular calcium, 100 ms test pulses), control FDB fibers did
Fig. 1. Characterization of the \( \text{Ca}_{3.1}^{1.1\Delta E29} \) mouse. (A) Targeting strategy for generating the \( \text{Ca}_{3.1}^{1.1\Delta E29} \) exon 29 knockout allele. (B) Voluntary home cage activity at 2 and 8 months of age is similar in \( \text{Ca}_{3.1}^{1.1\Delta E29} \) mice compared with wild-type and \( \text{Ca}_{3.1}^{1.1\Delta E29} \) siblings (\( N=5 \)). (C) Expression levels of \( \text{Ca}_{3.1}^{1.1a} \) and \( \text{Ca}_{3.1}^{1.1e} \) mRNAs in wild-type, \( \text{Ca}_{3.1}^{1.1\Delta E29} \) and \( \text{Ca}_{3.1}^{1.1\Delta E29} \) mice were measured by quantitative RT-PCR (TaqMan) in soleus, EDL and diaphragm muscle at different developmental stages (\( N=3 \)). Numbers beneath show the fractional content of the two \( \text{Ca}_{3.1} \) transcripts. (D) Western blot analysis of total \( \text{Ca}_{3.1} \) protein (both splice variants) in soleus and EDL muscle of wild-type and \( \text{Ca}_{3.1}^{1.1\Delta E29} \) mice (\( N=3; \ *** P<0.001 \)). Mean±s.e.m. See also Fig. S1.
not display measurable currents at any test potential. In order to compare voltage sensitivity between the two genotypes, recordings in control fibers were repeated using 5 mM extracellular calcium and 500 ms test pulses. The current-voltage and current-conductance curves indicate that in \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) fibers half-maximal current activation is shifted by 38.5±3.1 mV in the hyperpolarizing direction (Fig. 4B). Accordingly, the simultaneously recorded calcium transients in \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) fibers showed a pronounced voltage-dependent component that peaked at \(-20\) mV and declined in parallel with the current density at positive potentials (Fig. 4C).

From the calcium transients the total calcium flux (influx and SR release) during the depolarizing pulses was calculated (Fig. 4A lower trace, Fig. S2). These calcium flux traces characteristically showed an early peak followed by a steady-state plateau phase (Szentesi et al., 1997). In control muscle, the voltage dependence of both the peak and plateau calcium fluxes displayed a monotonic increase, which could be fitted with a two-state Boltzmann function. This is consistent with a model of skeletal muscle EC coupling in which both parameters of the calcium flux (peak and plateau) represent a single process, i.e. calcium release from the SR. In \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) muscle fibers, the peak calcium flux behaved like that in controls, whereas the voltage dependence of the plateau flux was non-monotonic. At intermediate voltages the plateau flux was significantly larger than that of controls. Thus, the additional component in \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) muscles compared with controls corresponds to voltage-dependent calcium influx, which peaks at \(-20\) mV where most \( \text{CaV}1.1\text{e} \) channels are activated and then decreases in parallel with the declining driving force at positive membrane potentials. Together, these analyses clearly demonstrate that, during EC coupling, muscle fibers of \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) mice experience increased calcium signals due to a substantial component of calcium influx that is not observed in muscle fibers of wild-type mice.

In addition to its immediate role in EC coupling, this extra calcium influx in \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) mice might alter the calcium homeostasis in muscle cells. Analysis of cytoplasmic calcium concentrations in Fura-2-loaded isolated FDB fibers revealed no difference in resting calcium levels of \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) mice compared with wild-type controls (45.77±0.96 nM and 46.14±1.14 nM, respectively; \( P>0.5 \)). Next, we examined a possible contribution of L-type calcium currents to refilling of SR calcium stores. Both SR depletion and refilling resulted in a robust cytoplasmic calcium transient (Fig. 4D), the relative magnitude of which was expressed as the external to SR calcium transient ratio. In wild-type muscle fibers this ratio was not affected by the application of the L-type calcium channel blocker nisoldipine. However, in \( \text{CaV}1.1^{\Delta E29/\Delta E29} \) muscle fibers 5 µM nisoldipine dramatically reduced the ratio, indicating that in \( \text{CaV}1.1\text{e} \)-expressing muscle SR refilling is predominantly carried out by calcium influx through L-type channels (i.e. \( \text{CaV}1.1\text{e} \)).

Finally, the spontaneous occurrence of focal calcium transients – so called calcium sparklets – in resting muscle fibers was investigated. Because similar local calcium release events are exclusive to muscle cells expressing calcium-conducting \( \text{CaV}1.1\text{e} \) channels...
channels (e.g. CaV1.2 in cardiac myocytes), we hypothesized that aberrant expression of CaV1.1e might also bring about calcium sparklets in mature muscle fibers of CaV1.1ΔE29/ΔE29 mice. Indeed, isolated FDB fibers from CaV1.1ΔE29/ΔE29 mice loaded with Fluo-8 AM displayed spontaneous calcium sparklet-like behavior (Fig. 4E). The average amplitude of these calcium sparklets was 0.140±0.001 [ΔF/F]. The average amplitude, full width and half maxima of these calcium sparklets resembled those described by Rodriguez et al. (2014). In 5 mM extracellular calcium, these calcium sparklets occurred at a frequency of 0.724 ±0.211 10⁻³ s⁻¹ mm⁻¹. When the extracellular calcium concentration was reduced to 1.8 mM, or when L-type calcium
channels were blocked with 10 μM nisoldipine, the calcium sparklets were completely abolished, indicating their dependence on calcium influx through CaV1.1e. In wild-type control muscle fibers no such calcium release events were observed in normal or high extracellular calcium concentrations. Together, these findings demonstrate that calcium influx through the CaV1.1e splice variant not only alters calcium handling during EC coupling and during refilling of SR calcium stores, but also causes spontaneous calcium signals in resting intact muscle fibers.

**Muscles of CaV1.1e-expressing mice display an altered fiber type composition and oxidative metabolism**

In skeletal muscle, calcium signals also regulate activity-dependent control of muscle growth and fiber type specification. Moreover, the observed changes in contractile properties – decreased force, increased fatigue resistance and lower tetanic fusion frequency – are all reminiscent of the differences between fast and slow muscle types. Therefore, we hypothesized that the altered calcium signaling in CaV1.1ΔE29ΔE29 mice might affect contractile properties indirectly by a dysregulation of fiber type specification.

To analyze the fiber type composition of slow and fast muscles in wild-type and CaV1.1ΔE29ΔE29 mice, we immunostained sections of soleus and EDL muscle with antibodies against specific myosin heavy chain isoforms. The representative images in Fig. 5A,D demonstrate a substantial shift towards slower fiber types in both soleus and EDL muscles of CaV1.1ΔE29ΔE29 mice. Soleus muscles of CaV1.1ΔE29ΔE29 mice experienced a 48% increase in the fraction of type I fibers, mainly at the cost of type IIA and mixed fibers (Fig. 5A-C). In EDL muscle of CaV1.1ΔE29ΔE29 mice the fraction of type IIB fibers was reduced by 26%, whereas the fractions of IIA, IIX and mixed fibers increased 2- to 3-fold (Fig. 5D-F). Type I fibers were not detected in CaV1.1ΔE29ΔE29 EDL muscles. These findings indicate that the expression of CaV1.1e alters fiber type-specific activation patterns and function as master regulators of fiber type specification this might be reflected in the activity and/or expression levels of major calcium- and activity-regulated signaling proteins. In skeletal muscle, the calcium-dependent protein phosphatase calcineurin (protein phosphatase 2B) and the calmodulin-dependent protein kinase II (CaMKII) decode fiber type-specific activation patterns and function as master regulators of fast to slow fiber type changes (Chin et al., 1998; Wu et al., 2001; Chin, 2005). Using a colorimetric phosphatase assay we show that steady-state calcineurin activity is significantly increased in CaV1.1ΔE29ΔE29 mice compared with wild-type controls (Fig. 6B, Fig. S3A). The left shift of the intensity distribution diagrams was more pronounced in soleus, but it was still significant in EDL. It equally affected fibers with low and high SDH activity. Together, these results show that the initial increase of oxidative metabolism in young CaV1.1ΔE29ΔE29 mice is lost and even reversed at 6 months and older.

Because decreased SDH activity could arise from either a reduction in mitochondrial activity or content, we analyzed mitochondria in electron microscopy preparations. Consistent with the overall healthy state and normal motor performance of the CaV1.1ΔE29ΔE29 mice, electron microscopy did not reveal any defects in the myofibrils and EC coupling membranes (Fig. 6C). However, the mitochondria were distorted in CaV1.1ΔE29ΔE29 muscles. Morphometric analysis revealed that the mitochondrial content was significantly reduced in CaV1.1ΔE29ΔE29 mice to approximately half that in wild-type controls (Fig. 6D, Fig. S3B). This loss of intact mitochondria was paralleled by an increase in the fraction of damaged mitochondria up to 4.5-fold compared with wild-type controls, as well as a decrease in the average size of the healthy mitochondria. These findings explain the significantly reduced SDH activity observed in CaV1.1ΔE29ΔE29 compared with wild-type muscles of the same age.

**Aberrant expression of CaV1.1e changes key activity- and calcium-dependent regulators of fiber type specification and mitochondrial biogenesis**

If the altered calcium signals in CaV1.1e-expressing muscles impact the regulation of fiber type specification this might be reflected in the activity and/or expression levels of major calcium- and activity-regulated signaling proteins. In skeletal muscle, the calcium-dependent protein phosphatase calcineurin (protein phosphatase 2B) and the calmodulin-dependent protein kinase II (CaMKII) decode fiber type-specific activation patterns and function as master regulators of fast to slow fiber type changes (Chin et al., 1998; Wu et al., 2001; Chin, 2005). Using a colorimetric phosphatase assay we show that steady-state calcineurin activity is significantly increased in CaV1.1ΔE29ΔE29 soleus and EDL muscles (Fig. 7A). Western blot analysis using a phospho-specific antibody demonstrated that the activated forms of all three CaMKII isoforms were significantly increased in cytoplasmic fractions of soleus muscle (Fig. 7B), whereas in EDL muscles no changes in CaMKII activation were observed. A differential activation of CaMKII in slow versus fast muscles is consistent with its suggested role in differentially decoding slow and fast muscle calcium signals (Tavi and Westerblad, 2011).

Interestingly, expression of their respective downstream transcriptional regulators, NFATC1 and HDAC4, in the cytoplasm and nuclei of CaV1.1ΔE29ΔE29 muscles was not altered (Fig. 7C,D, Fig. S4A,B). However, expression of peroxisome

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**Table 1. Contractile properties of soleus and EDL muscles of CaV1.1ΔE29ΔE29 and wild-type mice**

| Property                      | Soleus | EDL  |
|-------------------------------|--------|------|
|                               | (N=5)  | (N=15) |
| Twitch force (mN/mm²)          | 70.3±10.0 | 50.2±4.8 |
| Tetanic force (mN/mm²)         | 492.4±67.4 | 368.1±22.8 |
| Fatigue (100th tetanus) %      | 45.2±2.8 | 35.3±2.7 |
| Fatigue (150th tetanus) %      | 54.2±2.4 | 42.8±2.8 |
| Force-frequency (slope, k)     | 9.20±0.67 | 6.90±0.45 |
| Force-frequency (F₅₀, Hz)      | 19.74±0.85 | 11.97±0.44 |

**Wild type (N=5) vs Knockout (N=15)**

| Property | Wild type | Knockout | P-value |
|----------|-----------|----------|---------|
| Twitch | 78.0±7.3 | 54.2±4.1 | <0.001  |
| Tetanic | 447.6±27.3 | 289.1±19.3 | <0.0001 |
| Fatigue | 52.1±2.3 | 44.9±2.2 | <0.001  |
| Fatigue | 64.0±2.3 | 50.4±3.3 | <0.001  |
| Force-frequency (slope) | 17.34±0.67 | 12.90±0.54 | <0.001  |
| Force-frequency (F₅₀) | 34.75±1.61 | 27.09±1.02 | <0.0001 |

*Values represent means±s.e.m.

*Frequency of half-maximal force.
Proliferator-activated receptor γ co-activator 1α (PGC1α, Ppargc1a), a key downstream regulator of mitochondrial biogenesis and of oxidative metabolism in muscle (Handschin et al., 2007; Lin et al., 2002), was significantly increased in soleus muscle of CaV1.1ΔE29/ΔE29 mice compared with wild-type controls (Fig. 7E). In EDL muscle, PGC1α expression was not significantly altered. Expression of Six1, a key regulator of the fast fiber program (Grifone et al., 2004; Wu et al., 2013), was not affected.

To further examine potential effects on the expression patterns of the slow and fast program and on mitochondrial biogenesis, Affymetrix expression profiling was performed on mRNA preparations from soleus and EDL muscles of wild-type and CaV1.1ΔE29/ΔE29 mice. Although the analysis showed differential gene expression in soleus versus EDL muscles of both genotypes, comparison of wild-type versus CaV1.1ΔE29/ΔE29 soleus and EDL muscles did not reveal any significant differences (Fig. S4C). Also, the specific analysis of genes involved in mitochondrial fusion and fission revealed only small differences between slow and fast muscles but not between wild-type and CaV1.1ΔE29/ΔE29 mice. Thus, the increased calcium influx through CaV1.1e chronically hyperactivates calcineurin, CaMKII and PGC1α signaling and, over time, produces the observed changes in fiber type composition.

Fig. 4. Altered calcium signaling in isolated muscle fibers of CaV1.1ΔE29/ΔE29 mice. Enzymatically isolated FDB fibers from 3- to 4-month-old CaV1.1ΔE29/ΔE29 (red) and wild-type (black) mice were loaded with Rhod-2 (A–C) or Fluo-8 AM (D–G). (A) Representative voltage-clamp recording of calcium currents and parallel recording of cytoplasmic free calcium during a 100 ms depolarization to -30 mV. Bottom trace is the calculated total calcium flux. (B) Voltage dependence of peak current densities and conductance display the 38.5±3.1 mV shift of channel activation in CaV1.1ΔE29/ΔE29 fibers. Note that CaV1.1ΔE29/ΔE29 fibers were recorded in 1.8 mM extracellular calcium during 100 ms test pulses and control fibers in 5 mM calcium during 500 ms test pulses to experimentally normalize current densities. (C) Calcium transient amplitudes display a striking increase in CaV1.1ΔE29/ΔE29 muscle fibers at intermediate voltages, indicative of the calcium influx through CaV1.1e. (D) Representative calcium recordings of wild-type and CaV1.1ΔE29/ΔE29 fibers during an SR calcium release and reloading protocol. Note that the relative magnitude of the release and reloading transients is significantly decreased upon nisoldipine block of CaV1 channels in CaV1.1ΔE29/ΔE29 but not in wild-type muscle (**P<0.01). (E) Representative confocal images of a resting CaV1.1ΔE29/ΔE29 FDB fiber in 1.8 mM and 5 mM calcium. Spontaneous calcium release events occur only in 5 mM calcium and are blocked by addition of 10 µM nisoldipine. (F) Spatiotemporal properties in line-scan images identify the localized calcium signals as calcium sparklets. (G) The sparklets are sensitive to the extracellular calcium concentration and L-type channel block. n=32 images/N=4 animals (1.8 mM), n=183/N=7 (5 mM) and n=24/N=3 (5 mM+nisoldipine). Mean±s.e.m. See also Fig. S2.
without, however, a major induction of the slow muscle gene program at basal activity levels in adult mice.

**DISCUSSION**

The developmental switch of CaV1.1 splice variants is important for the correct specification of skeletal muscle fiber types

Several lines of evidence at the molecular, functional and behavioral level demonstrate that the continued expression of the developmental CaV1.1e splice variant in mature muscles of CaV1.1\(^{ΔE29/ΔE29}\) mice alters the fiber type composition in the slow direction: (1) in soleus muscle the fraction of type I fibers increases at the expense of type IIA fibers, and in EDL muscles type IIA, IIX and mixed fibers increase at the expense of type IIB fibers; (2) in isolated soleus and EDL muscles maximal twitch and tetanic forces are reduced, fatigue resistance is increased, and tetanic fusion of contractions occurs at lower frequencies; (3) grip force and the duration of voluntary wheel running are reduced; (4) calcineurin and CaMKII activity and expression of PGC1α are upregulated.

However, in mature mice the increased slow fiber content and the increased expression of PGC1α were no longer accompanied by the expected increase in oxidative enzyme activity, most likely because the increased calcium influx in CaV1.1\(^{ΔE29/ΔE29}\) muscles also caused a severe loss of mitochondria.

The fiber type composition of skeletal muscles is primarily a genetically determined adaptation of muscle properties to their specific physiological functions. Furthermore, it is subject to continuous dynamic adaptation to altered demand. The changes in fiber type composition and contractile properties observed in the CaV1.1\(^{ΔE29/ΔE29}\) mice are reminiscent of adaptations occurring in response to endurance training. The magnitude of the observed shift in fiber type composition is within the range observed in mice subjected to stringent endurance training protocols (Allen et al., 2001; Krüger et al., 2013). However, neither monitoring home cage activity nor the behavioral tests revealed increased spontaneous activity. If anything, CaV1.1\(^{ΔE29/ΔE29}\) mice spent less time voluntarily running than wild-type controls. Therefore, the observed changes in fiber type composition cannot be explained.
Fig. 6. Mitochondrial content and enzyme activity are reduced in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} muscles. (A,B) SDH activity was analyzed in sections of wild-type (black) and Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} (red) mice. Staining intensity was measured in each fiber profile and plotted in intensity distribution diagrams. (A) In 7-week-old mice, SDH activity is markedly increased in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} soleus and EDL muscles, as seen by the right-shifted intensity distribution histogram. (B) In 6-month-old mice, SDH activity is significantly reduced in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} soleus muscle and in EDL muscles. In EDL also the full width at half maximum (FWHM) was reduced. N=3. (C) Electron micrographs of 4- and 5-month-old wild-type and Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} soleus and EDL muscles. Whereas myofibrils and EC coupling membranes appear normal, dilated and lysed mitochondria are found in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} muscles. (D) Morphometric analysis demonstrates significantly decreased fractional content (percentage area occupied by intact mitochondria) in both Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} muscle types. Mitochondrial size is decreased in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} muscles and the fraction of damaged mitochondria increased in Ca\textsubscript{v}1.1\textsuperscript{ΔE29/ΔE29} soleus muscles. N=30-40 images of two biological replicates per condition. *P<0.05, **P<0.01, ***P<0.001. Mean±s.e.m. See also Fig. S3. Scale bars: 100 µm in A,B; 0.5 µm in C.
as a normal adaptive response to altered activity, but represent a dysregulation of fiber type composition owing to the absence of postnatal inclusion of CaV1.1 exon 29 in CaV1.1ΔE29/ΔE29 mice. Since CaV1.1 is almost exclusively expressed in skeletal muscle and the functional effects (e.g. reduced force, increased fatigue resistance) were observed in isolated muscles, muscle-intrinsic mechanisms are likely to be responsible for the altered muscle fiber type composition in CaV1.1ΔE29/ΔE29 mice.

Expression of CaV1.1.1e alters skeletal muscle calcium signaling and activates the pathways for the slow twitch fiber program

Calcium is the principal second messenger regulating adaptive changes of muscle properties in response to training or experimentally altered innervation patterns (Bassel-Duby and Olson, 2006; Chin et al., 1998; Liu et al., 2005). As the primary defects in the CaV1.1ΔE29/ΔE29 mice are altered gating and conduction properties of the skeletal muscle L-type calcium channel expressed in adult mice, a role of increased calcium influx in determining the fiber type composition is very likely. In cultured dysgenic myotubes reconstituted with either of two CaV1.1 splice variants, we previously reported that exclusion of exon 29 caused a 30 mV left-shifted voltage dependence of current activation and an 8-fold increase in current density (Tuluc et al., 2009). Here, we demonstrate that also in isolated muscle fibers of CaV1.1ΔE29/ΔE29 mice, sole expression of the CaV1.1e splice variant causes an equally large left shift of voltage sensitivity and a substantially increased calcium influx during EC coupling.

Interestingly, the calcium influx on top of the calcium released from the SR was not reflected in a parallel increase in contractile force

Fig. 7. Altered expression of regulators of fiber type specification and mitochondrial biogenesis in slow and fast muscles of CaV1.1ΔE29/ΔE29 mice.

(A) Basal enzymatic activity of calcineurin is significantly increased in CaV1.1ΔE29/ΔE29 soleus and EDL muscles (red) compared with wild type (black). (B) Western blot analysis shows a significant increase of activated CaMKII β, γ and δ isoforms (phosphorylated at Thr286) in soleus but not EDL muscle of CaV1.1ΔE29/ΔE29 (C,D) Cytoplasmic and nuclear localization of NFATc1 and HDAC4 does not differ between the genotypes (age of mice 6-8 months). Controls for the cytoplasm and nuclear fraction are GAPDH and histone H3, respectively. N=3. (E) Quantitative RT-PCR analysis demonstrates that expression of PGC1a (Ppargc1a) mRNA is significantly increased in CaV1.1ΔE29/ΔE29 soleus muscles, whereas the decline of PGC1a in EDL and of Str1 expression in both muscles is not significant. N=3. *P<0.05, ***P<0.001. Mean±s.e.m. (F) Model of signaling pathways causing fiber type shift and mitochondrial damage in CaV1.1ΔE29/ΔE29 muscles. During EC coupling, increased calcium influx through CaV1.1.1e activates calcineurin (CN), CaMKII and PGC1a, the primary regulators of fiber type specification and mitochondrial biogenesis. The shift in fiber type composition in the slow direction offsets the direct effects of increased calcium signals on EC coupling. Parallel upregulation of oxidative metabolism by PGC1a is counteracted by mitochondrial damage caused by calcium overload due to increased activity-dependent and spontaneous calcium influx and altered calcium homeostasis. See also Fig. S4.
expression of PGC1α, were constitutively upregulated in skeletal muscles of CaV1.1 ΔE29/ΔE29 mice, and that activation of CaMKII and PGC1α was specific to slow muscles. In accordance with this, altered CaMKII signaling was also observed in the calcium permeation CaV1.1 mutant (Lee et al., 2015).

Thus, in CaV1.1 ΔE29/ΔE29 mice the continuous expression of CaV1.1e causes increased calcium influx during EC coupling, in homeostatic calcium regulation, and at rest. Which of these calcium influx events contribute to the activation of the slow muscle pathway remains to be determined. Because calcineurin and CaMKII signaling is highly sensitive to the calcium signaling patterns in response to slow fiber type-specific activity, we favor a role of altered calcium signal during EC coupling. In any case, if aberrant activation of these signaling pathways causes an increase in slow fibers in CaV1.1 ΔE29/ΔE29 mice, during normal development the alternative splicing event causing the shift from a calcium-conducting to a non-conducting CaV1.1 variant might be an important prerequisite for the proper regulation of fiber type composition at basal activity levels as well as in response to exercise.

**The contribution of aberrant CaV1.1 splicing to myotonic dystrophy**

Splicing defects of important muscle proteins, including the CLCN1 chloride channel, insulin receptor, SERCA1 (ATP2A1) and CaV1.1, lead to DM1 (Thornton, 2014). The myotonia is likely to be caused by a hyperexcitability of muscles due to the loss of CLCN1 function (Lueck et al., 2007). Because aberrant expression of the CaV1.1e splice variant correlates with the degree of muscle weakness in DM1 patients, and forced missplicing of CaV1.1 exon 29 caused centrally localized nuclei in a myotonia mouse model, it has been suggested that increased calcium influx through the developmental CaV1.1e splice variant may contribute to the myopathy (Santoro et al., 2014; Tang et al., 2012). This hypothesis is in line with the known role of increased calcium influx via various entry pathways in muscular dystrophy (Allen et al., 2010; Whitehead et al., 2006). Here, we examined whether aberrant missplicing of CaV1.1 exon 29 by itself is sufficient to cause DM-like symptoms. At the organismal level this was not the case. Homozygous CaV1.1 ΔE29/ΔE29 mice, which exclusively express the developmental CaV1.1e splice variant, did not show severe muscle weakness, and their muscle sections did not reveal centrally located nuclei, which is a histopathological hallmark of dystrophic muscle. Although the contractile force of isolated muscles was reduced, this was accompanied by increased fatigue resistance and might therefore be the consequence of the fiber type shift rather than a symptom of DM1.

We observed decreased SDH activity and severe mitochondrial damage in muscles of mature and aged CaV1.1 ΔE29/ΔE29 mice. Similar mitochondrial damage has been described in other mouse muscle disease models with aberrant calcium signaling (Irwin et al., 2003). Calcium overload leads to mitochondrial damage and ultimately produces the symptomatic central cores in the diseased muscles (Boncompagni et al., 2009; Brooks et al., 2004). Similarly, the CaV1.1e-mediated spontaneous calcium sparks, or the increased calcium influx after store depletion, might overburden the mitochondrial calcium handling capacity and cause their loss in CaV1.1 ΔE29/ΔE29 mice. In fact, both calcium sparks and store-operated calcium currents have previously been implicated in the pathophysiology of muscular dystrophy (Goonasekera et al., 2014; Weisleder and Ma, 2006). Thus, the mitochondrial damage observed in CaV1.1 ΔE29/ΔE29 mice might precede the appearance of histopathological and clinical features of DM1.
We conclude that, by itself, missplicing of exon 29 in Ca\textsubscript{v}1.1 is not sufficient to reproduce the full spectrum of DM1 symptoms in mice. As the disease phenotype in dystrophy and myotonia mouse models is often less severe than in human disease, this does not preclude the possibility that missplicing of Ca\textsubscript{v}1.1 makes a notable contribution to the disease in DM1 patients. Furthermore, it is likely that in combination with the splicing defects in other muscle genes involved in calcium handling (Sercal1) and excitability (Clen1), the mitochondrial damage observed in the Ca\textsubscript{v}1.1\textit{ΔE29/ΔE29} mice might be aggravated, and thus contribute to the myopathy (Tang et al., 2012). If so, the use of clinically approved L-type calcium channel blockers to target Ca\textsubscript{v}1.1c currents might be a viable strategy to alleviate the symptoms of DM1 (Benedetti et al., 2015).

In conclusion, the effects of Ca\textsubscript{v}1.1 missplicing in the Ca\textsubscript{v}1.1\textit{ΔE29/ΔE29} mice revealed a novel role of increased L-type calcium currents in the dysregulation of muscle fiber type composition. With regard to normal muscle physiology these findings suggest that, during development, skeletal muscles actively suppress L-type calcium currents by alternative splicing of Ca\textsubscript{v}1.1, so as to prevent unrestrained calcium influx during EC coupling from interfering with the second calcium signaling function in regulating muscle fiber type composition. Furthermore, aberrant expression of the developmental Ca\textsubscript{v}1.1e splice variant in mature muscles causes mitochondrial damage, which might contribute to the pathology of DM1.

MATERIALS AND METHODS

Mice and animal care

Ca\textsubscript{v}1.1\textit{ΔE29} mice were generated in a C57BL/6N background at Taconic Artemis (Cologne, Germany). All experimental protocols conformed to guidelines of the European Community (86/609/EEC) and were approved by the Austrian Ministry of Science (BMWF-66.011/0069-II/10b/2010) and by the Institutional Animal Care Committee of the University of Debrecen (22/2011/DE MAB). Genotyping is described in the supplementary Materials and Methods.

Quantitative RT-PCR

RNA was isolated from muscles of wild-type and Ca\textsubscript{v}1.1\textit{ΔE29} mice using the RNeasy Fibrous Tissue Mini Kit (Qiagen). After reverse transcription (SuperScript II reverse transcriptase, Invitrogen) the absolute number of transcripts was assessed by quantitative TaqMan PCR (50 cycles). For western blotting. Calcineurin activity was determined using a Calcineurin Phosphatase Activity Colorimetric Assay (Abcam) according to the manufacturer’s instructions. For antibodies and experimental details, see the supplementary Materials and Methods.

Western blot and calcineurin assay

Cytoplasmic and nuclear protein fractions were prepared from 6-month-old wild-type and Ca\textsubscript{v}1.1\textit{ΔE29/ΔE29} mice and separated on 6-10% bis-Tris gels for western blotting. Calcineurin activity was determined using a Calceinurin Phosphatase Activity Colorimetric Assay (Abcam) according to the manufacturer’s instructions. For antibodies and experimental details, see the supplementary Materials and Methods.

Electron microscopy

Tissue processing for transmission electron microscopy of EDL and soleus muscles, and the analysis of mitochondria, were performed as detailed in the supplementary Materials and Methods.

Statistical analysis

A two-way ANOVA with Bonferroni post-hoc test was used for home cage activity and fiber type analysis. One-way ANOVA was used for the other behavioral tests. Student’s t-test and Mann–Whitney U-test were used to calculate the statistical significance for fiber type and contractile properties analysis. N represents the number of animals. Data are presented as mean± s.e.m. The statistical analysis was performed with GraphPad Prism. For further details see the supplementary Materials and Methods.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

B.E.F., N.S. and L.C. designed research and wrote the manuscript. N.S., B.D., A.B., P.T., P.S., M.S., J.R., M.W.H., C.S. and G.J.O. performed experiments and/or analyzed data.

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Data availability

The raw and preprocessed microarray data are available at Gene Expression Omnibus under accession number GSE67803.

Supplementary information

Supplementary information available online at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.129676/-/DC1

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