INTRODUCTION

Evolution of the mammalian motor system affords many morphologically and functionally different animals to thrive in diverse environmental conditions (Brownstone, 2020). Yet, the constantly changing environment—intrinsic and extrinsic—requires that an organism’s motor systems are capable of functional adaptations.

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The muscular system itself is responsive to chronic changes in functional demand. For example, aerobic exercise training improves endurance capacity through coordinated increases in muscle
activation patterns, which drives targeted, functional adaptation of skeletal muscle, altering metabolic signalling and expanding the vascular bed (Jensen et al., 2004). Additionally, chronic increases in loading drive muscle hypertrophy and increase force capacity (Mitchell et al., 2012). Moreover, constitutive fibre types, usually classified as slow oxidative (S-Type I), fast fatigue resistant (FR-Type IIA) and fast fatigable (FF-Type IIB/X), can undergo adaptive changes that alter muscle functional properties. For example, endurance training increases the proportion of Type I and IIA oxidative fibres resulting in improved fatigue resistance (Green et al., 1983).

Motoneurons innervating skeletal muscle also adapt to changing demands with electrophysiological adaptations matched to properties of the innervated muscle. This was elegantly shown by cross-reinnervation studies in which forced mismatches between muscle and motoneuron properties were induced by surgically re-routing motor axons from one muscle to another (e.g. medial gastrocnemius [MG] to soleus; Dum et al., 1985; Foehring et al., 1987). Following recovery, assessments of motoneuron properties at a chronic stage showed a shift towards those of the reinnervated muscle, suggesting mechanisms of plasticity exist within the motor unit to match muscle and motoneuron properties.

Physiological stimuli such as exercise also induce functional adaptations in motoneuron properties: endurance training in rats induces increases in the motoneuron medium after hyperpolarisation (mAHP) amplitude, consistent with a shift to a more fatigue-resistant phenotype (Gardiner et al., 2006). Thus, physiological adaptations in motor units should always be reflected centrally and peripherally.

Altered neuromuscular demand can also stimulate changes in organisation of synaptic inputs to motoneurons and in expression of post-synaptic membrane proteins, both of which reflect changes in neuromuscular activity (Arbat-Plana et al., 2017; Woodrow et al., 2013). Thus, adaptations in spinal cord physiology parallel changes in the muscular system. However, it is less clear what type of inputs are sensitive to chronic changes in activity, and the degree to which they might adapt. For example, gain-of-function adaptations in neuromodulatory inputs that amplify motor output may be a beneficial response to chronic increases in load. Equally, neuromodulatory systems may already be equipped to respond to chronic increases in load and thus do not need to adapt the pre- or post-synaptic molecular machinery.

Motoneurons receive neuromodulatory cholinergic synapses, termed C-boutons, because of their association with post-synaptic subsurface cisternae (SSC; Conradi, 1969) that regulate the mAHP (Miles et al., 2007). These somatic/proximal dendritic synapses can amplify motor output in a task-specific manner. The system is comprised of V_{O2} premotor interneurons (Zagoraiou et al., 2009), C-bouton synapses, and clusters of several different post-synaptic membrane proteins (Witts et al., 2014). Recent work has suggested mechanisms for how some of these components may contribute to amplification of motor output during high force output tasks such as swimming (Nascimento et al., 2020; Romer et al., 2019; Soulard et al., 2020). Activation of type 2 muscarinic acetylcholine receptors (m2AChR) on spinal motoneurons results in a reduction in the mAHP amplitude and an increase in excitability as measured by the frequency/current (f–I) relationship (Miles et al., 2007). The mAHP current is carried by small conductance, calcium-activated potassium channels SK2 & SK3, which are differentially expressed in fast and slow motoneurons, endowing them with their respective mAHP characteristics (Deardorff et al., 2013).

Also clustered opposite C-boutons are delayed rectifier, voltage-gated potassium channels, Kv2.1 (Muennich & Fyffe, 2004). The role of these channels is less well understood, but recent evidence suggests they may act as ‘molecular rheostats’, capable of maintaining firing during high synaptic drive or suppressing firing to protect motoneurons from excitotoxicity (Romer et al., 2019). It has been shown that m2AChRs modulate Kv2.1 channels by reducing action potential half-widths and increasing the inter-spike AHP, which aids recovery of Na^+ channels during high synaptic drive; thus, supporting high-frequency firing (Nascimento et al., 2020). Furthermore, as in various brain regions (Murakoshi et al., 1997; Park et al., 2006), the distribution of motoneuron Kv2.1 channels is plastic, suggesting they may play a role in neuromuscular adaptation in health and disease (Romer et al., 2014). Specifically, high activity states that increase intracellular calcium concentrations cause Kv2.1 channels to rapidly de-cluster, which lowers their activation threshold and increases conductance. In lower activity states, Kv2.1 channels coalesce into macro-clusters that form physical links with the SCC (Deardorff et al., 2021), and are thought to be non-conducting. However, the functional significance of Kv2.1 channel conducting and non-conducting roles in behaviour has yet to be determined.

Several groups have studied plasticity of C-boutons in disease states. For example, Landoni et al. (2019) showed that C-bouton transmission initially compensates for progression of motor deficits during motoneuron loss in SOD1 amyotrophic lateral sclerosis (ALS) mice. Conversely, Konsolaki et al. (2020) have shown that C-bouton inactivation improves motor performance but not survival in SOD1 ALS mice. It is difficult, however, to separate mechanisms associated with disease or injury and chronic physiological overload of the neuromuscular system. Therefore, it is important to study how chronic changes in neuromuscular demand affect central components of the motor system, such as the motoneuron and its modulatory inputs.

Here, we asked whether such changes in muscular demand lead to corresponding adaptations in motoneurons and at C-bouton synapses. We used a model of chronic neuromuscular overload, as similar models have previously been shown to induce central and peripheral adaptations in motor units (Chalmers et al., 1991; Ianuzzo et al., 1976; Krutki et al., 2015; Rosenblatt & Parry, 1992). This involved extirpating the tibialis anterior (TA) muscle to increase loading of the remaining synergist extensor digitorum longus (EDL) muscle in adult rats for 21 days. We confirmed effectiveness of the overload stimulus by assessing changes in muscle physiology, showing a shift to a more fatigue-resistant phenotype. We then studied EDL motoneuron adaptations using retrograde tracers, and showed a corresponding reduction in cross-sectional area. Although we hypothesised that overload would induce adaptations to C-bouton organisation that correlated with adaptations seen in muscle physiology, there

**References**

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were no measurable differences in sizes and densities of both pre-
(C-boutons) and post-synaptic (K\(_{\text{v}2.1}\) & SK3) components, however,
there was a reduction in the proportion of SK3\(^{+}\) cells following over-
load. Our results show that in conjunction with a slower muscle
phenotype following overload, there is a corresponding decrease in
motoneuron size. We suggest that this central adaptation may com-
penstate for increased functional demands by reducing motoneuron
rheobase and increasing excitability. Furthermore, anatomical plas-
ticity of the neuromodulatory C-bouton complex is not necessary to
produce increased force output in this model of chronic functional
overload.

2 | METHODS AND MATERIALS

2.1 | Ethical approval

All surgical and experimental protocols were approved by the
University of Leeds Animal Welfare and Ethics Committee and con-
ducted in accordance with United Kingdom (UK) Animals (Scientific
Procedures) Act 1986 (ASPA). The investigators understand the
ethical principles under which the journal operates and confirm this
work complies with the journal animal ethics guidelines.

2.2 | Animals

Male Wistar rats (\(N = 14; 283 \pm 29\) g) were housed under a 12:12
light-dark cycle in a temperature-controlled 21°C environment, with
ad libitum access to food and water. Animals were randomly allo-
cated to either control of overload conditions.

2.3 | Animal surgical procedures

All animal surgeries were completed by competent Home Office-
approved PIL holders, under aseptic conditions. Surgical anaesthesia
was induced and maintained with isoflurane (5% and 2%, respec-
tively, in 100% \(\text{O}_2\). IsoFlo®; Zoetis UK Ltd).

2.4 | Muscle overload

An incision was made two thirds up the length of the right TA to-
wards the lateral side of the muscle. The covering fascia was cleared
exposing the TA muscle, enabling sectioning of the distal tendon
above the retinaculum and as close as possible to the proximal inser-
tion (Egginton et al., 2011). Upon releasing the tendons, the TA was
bluntly dissected from the lateral tibia surface and removed, taking
care not to damage the underlying EDL. Skin was closed with 5-0
Mersilk suture (Ethicon, Johnson & Johnson Medical Ltd). Animals
received subcutaneous analgesia (0.015 mg/kg, Vetagesic®; Ceva)
and antibiotic (2.5 mg/kg, Baytril®; Bayer) for 2 days post-surgery.

Removal of the TA muscle increases the load burden on its syn-
ergist, the EDL. We, thus, use the term “overload” for this condition.
The overload period lasted 21 days.

2.5 | Motoneuron tracing

Retrograde fluorescent tracers were injected into the EDL 5 days
prior to terminal experiments. Tracers were injected into the medial
and lateral compartments of the EDL for separate assessment of mo-
toneurons innervating these compartments (data not included). One
\(\mu\)L of 1.5% 647 nm CTβi Alexa Fluor™ Conjugate (Invitrogen) was in-
jected into both medial and lateral compartments; 3 \(\mu\)L of 1.5% Fast
Blue (Polyscience, Inc.) was injected only into the medial compart-
ment. Skin was closed using 5-0 Mersilk suture (Ethicon, Johnson
& Johnson Medical Ltd). Animals received analgesic (0.015 mg/kg,
Vetagesic®; Ceva) and antibiotic (2.5 mg/kg, Baytril®; Bayer) subcuta-
neously for 2 days post-surgery.

2.6 | In situ muscle fatigability

Anaesthesia was induced with isoflurane (4% in 100% \(\text{O}_2\) and
maintained by constant infusion (30–35 mg kg\(^{-1}\) h\(^{-1}\)) of alf-
alone (Alfaxan: Jurox) via a catheter implanted into the external
jugular vein. A tracheotomy was performed to facilitate spontane-
ous breathing. Blood pressure and heart rate were monitored in
LabChart 8 (AD Instruments) via a carotid artery catheter connected
to a pressure transducer (AD Instruments).

EDL twitch force was quantified using a lever arm transducer sys-
tem (305B-LR; Aurora Scientific) and LabChart 8 (AD Instruments).
Unimpeded access to the EDL was enabled by dissection of sur-
rounding fascia, the distal tendon was then cut and attached to the
leaver arm of the force transducer. The peroneal nerve was exposed
and indirectly stimulated using bipolar stainless-steel electrodes
(Hudlicka et al., 1977), with muscle length and electrical current de-
livery optimised to generate maximal isometric twitch force. Fatigue
resistance of the EDL was determined using a protocol (10 Hz electri-
cal stimulation, 0.3 ms pulse width) to elicit a series of isometric con-
tractions over 3 min. A fatigue index (FI) was calculated as the ratio
of end-stimulation tension to peak tension (FI = end-stimulation ten-
sion/peak tension), using the mean of 5 consecutive twitches.

2.7 | Tissue preparation

Following successful in situ recordings, and remaining under anaes-
thesia, EDL were dissected and the muscle mid-belly was snap frozen
in liquid nitrogen cooled isopentane for muscle capillary analysis. All
frozen muscle tissue was stored at ~80°C until cryosectioning. Next,
animals were transcardially perfused with 0.1 M phosphate buffer
and fixed with 4% paraformaldehyde. Spinal columns were removed
immediately after perfusion and post-fixed in 4% PFA for 24 h. Spinal
cords were carefully dissected and cryoprotected in 30% sucrose at 4°C for 7 days. Next, the lumbar segments were isolated, frozen in OCT (Agar Scientific) and stored at −20°C.

2.8 | Muscle processing

EDL muscles were cryo-sectioned (−20°C, 12 µm), mounted on polylysine-coated slides (VWR International, Lutterworth, Leicestershire, UK) and stored at −20°C until staining. Fibre boundaries were labelled with anti-laminin antibodies (Sigma-Aldrich, L9393) to identify the basement membrane. Capillaries were labelled by *Griffonia simplicifolia* lectin I (Vector Laboratories, FL-1101, 1:250), an endothelial cell carbohydrate-binding protein. Photomicrographs were taken via a QImaging MicroPublisher 5.0 RTV camera (Teledyne QImaging) on a Nikon Eclipse E600 microscope (Nikon) at 20× magnification (field of view 440 × 330 µm) using a 2-second exposure time.

Indices for capillary-to-fibre ratio (C:F) and capillary density (CD) were derived from histological sections. These global indices describe gross changes in capillary supply, however, they lack descriptive power of local capillary distribution. The local capillary supply is a critical determinant of functional capacity and of significant importance in the functional overload model which presents with descriptive power of local capillary distribution. The local capillary supply, however, they lack descriptive power of local capillary distribution. The local capillary supply is a critical determinant of functional capacity and of significant importance in the functional overload model which presents with a significant angiogenic response and fibre hypertrophy (Kissane et al., 2020; Tickle et al., 2020). Therefore, to investigate the influence of concurrent expansion of the capillary bed and fibre hypertrophy on muscle function, we mathematically modelled skeletal muscle oxygen transport kinetics (Al-Shammari et al., 2019). Briefly, capillary distributions were digitally derived from histological sections and used to model as a point source of O₂ and estimations of tissue PO₂ were predicted using a number of model assumptions: oxygen demand (15.7 × 10⁻³ mol O₂ m⁻³ s⁻¹), myoglobin concentration (10.2 × 10⁻³ O₂ ml⁻¹), oxygen solubility (3.89 × 10⁻³ mol O₂ ml⁻¹ mmHg⁻¹), myoglobin diffusivity (1.73 × 10⁻⁷ cm² s⁻¹) and capillary radius (1.8–2.5 × 10⁻⁴ cm; Al-Shammari et al., 2019).

We performed histological assessments of the fibre-type distributions in both conditions, but tracer loading of the muscle reduced sample quality. These data were, therefore, excluded.

2.9 | Spinal cord immunohistochemistry

Spinal cord immunohistochemistry was performed as previously described (Smith et al., 2017). In brief, L3-L6 segments were sectioned at 50 µm on a cryostat (−20°C) and free-floating sections were collected and stored in PBS until staining. The sections were then washed in PBS (3 × 10 min) and incubated for 1 h in blocking solution (0.2% Triton X-100, PBS, NaCl and 10% normal donkey serum). The free-floating sections were then incubated for 48 h in primary antibodies diluted in blocking solution, washed and then incubated in secondary antibodies for 2 h, also in blocking solution. Primary antibodies: goat anti-vesicular acetylcholine transporter (anti-VACHT, Millipore Cat# AB100, RRID:AB_2630394, 1:1000), mouse anti-Kᵥ2.1 (UC Davis/NIH NeuroMab Facility Cat# 73–014, RRID:AB_10672253, 1:200) and rabbit anti-SK3 (Millipore Cat# AB5350-200UL, RRID:AB_91797, 1:200). Secondary antibodies at 1:200: Alexa Fluor® 555 donkey anti-mouse (Thermo Fisher Scientific Cat# A-31570, RRID:AB_2536180), Alexa Fluor® 488 donkey anti-goat (Jackson ImmunoResearch Labs Cat# 705-546-147, RRID:AB_2340430) and Alexa Fluor® 555 donkey anti-rabbit 555 nm (AB_2563181). Finally, sections were mounted on glass slides with Mowoil 4-88 (Carl Roth GmbH & Co. Kg).

2.10 | Confocal microscopy and quantitative analysis

Images were acquired with a Zeiss LSM 800 confocal microscope (Zeiss LSM 800 with Airyscan, RRID:SCR_015963), using a 40× oil immersion objective (1 AU aperture) and Zeiss ZEN Blue Edition software (ZEN Digital Imaging for Light Microscopy, RRID:SCR_013672). Motoneurons were identified by their location in the spinal cord ventral horn and presence of CTβ (467 nm) or FB (405 nm) fluorescence. Z-stacks of 30 µm at 0.40 µm intervals were acquired through the centre of each neuron, identified by the nucleus. Motoneurons that did not have a visible nucleus or had significant membrane disruption were excluded from due to poor reconstruction quality.

Researchers were blinded to the conditions for all data analyses. Three-dimensional (3D) reconstructions of each motoneuron were rendered from the confocal image z-stacks, utilising IMARIS Software (IMARIS, RRID:SCR_007370). In the 3D isometric view, solid surfaces of the motoneuron soma with dendrites, C-boutons and Kᵥ2.1 or SK3 were created via surface rendering and thresholding. CTβ or Fast Blue was used to model the motoneuron surface. A masking feature was then used to select Kᵥ2.1 or SK3 clusters contacting the motoneuron surface and/or proximal to the C-bouton. IMARIS was then used to generate volume and surface area data for each motoneuron, C-boutons, Kᵥ2.1 clusters and SK3 clusters. To determine the motoneuron cell size, cross-sectional area through the centre of the nucleus was calculated using Image J (ImageJ, RRID:SCR_003070).

Data were then exported to Excel (Microsoft Excel, RRID:SCR_016137). Since all alpha-motoneurons contain C-boutons and Kᵥ2.1, cells with no C-bouton or Kᵥ2.1 surfaces were removed. C-bouton, SK3 and Kᵥ2.1 channel densities were normalised to the motoneuron surface area. For statistical analyses, the mean synaptic and channel density for each motoneuron was the observational unit (n) and the average density per animal was the experimental unit (N). Sample sizes were determined based on previous studies (Kissane et al., 2018). Shapiro–Wilks tests were performed to determine normality of the data, followed by either unpaired t-tests (normally distributed) or Mann-Whitney U tests (not normally distributed). Fisher’s exact test was used to compare the proportion of cells expressing SK3. Data are presented throughout as mean ± standard deviation. All analyses were performed using Python scripts (RRID: SCR_008394) in the Jupyter notebooks environment (RRID: SCR_013995).
2.11 | Data Availability Statement

The data that support the findings of this study and a digital analysis notebook are openly available in zenodo at https://zenodo.org/badge/latestdoi/336327574 reference number [RRID:SCR_002630].

3 | RESULTS

3.1 | Chronic functional overload induces functional shift to slower EDL phenotype

Previous studies of EDL muscle overload by removal of the TA synergist have shown an anatomic and physiologic shift to a slower phenotype (Rosenblatt & Parry, 1992; Rosenblatt & Parry, 1993). We were not able to reliably analyse fibre-type distribution in this study due to the loading of muscles with neuro-anatomical tracers. However, muscle weight was significantly greater in the overload condition, suggesting hypertrophy of EDL fibres (Control = 0.06 ± 0.01 g, N = 5 vs. Overload 0.09 ± 0.02 g, N = 7, p = 0.001).

3.2 | Chronic functional overload improves fatigability in EDL muscles

In order to determine if there were physiological adaptations in EDL muscles following overload, we assessed muscle twitch force and fatigability using an in vivo anaesthetised preparation. Measurement of maximal isometric force showed that overloaded muscle had reduced twitch (227 ± 55 g/g, N = 7 vs. 296 ± 36 g/g, p = 0.017, N = 5, Figure 1a,a1) and tetanic (Control = 1184 ± 189 g/g, N = 5 vs. Overload = 864 ± 155 g/g, N = 7, p = 0.009, Figure 1b,b1) force outputs compared to control. Correspondingly, the FI was increased in the overload condition (0.64 ± 0.04) compared to control (0.50 ± 0.08, p = 0.0003, Figure 1c) indicating a shift to slower, more FR fibres in the overload condition. Assessments of muscle capillary

![Figure 1](image-url)

**FIGURE 1** Effect of overload on EDL force production and fatigability. (a) Maximum twitch force in control and overload animals normalised to EDL weight (g/g). (a1) Representative normalised twitch traces from a control (Con, blue) and overload (OL, green) EDL. (b) Maximum tetanic force normalised to EDL weight (g/g). (b1) Representative peak normalised traces during tetanic stimulation in a control and overloaded EDL muscle. (c) Fatigue index ratio (end-stimulation force/peak force). (d) Capillary-to-fibre ratio. (e) Capillary density (CD) per mm² of muscle fibre. (f) Modelled partial pressure of O₂ (PO₂). (g, h) Representative 20× light microscope images of lectin stained (capillaries) muscle fibres in control (g) and overload conditions (h). (i, j) Representative images of modelled PO₂ spatial profile for control and overload conditions. Experimental units (animals, N) and statistical tests are as follows: (a–c), control N = 7, overload N = 7, unpaired t-test; (d–f), control N = 6, overload N = 5, unpaired t-test. *Represents a statistically significant difference (p < 0.05). EDL, extensor digitorum longus.
supply highlighted the potent angiogenic response to overload with a significant increase in capillary-to-fibre ratio (Control = 1.67 ± 0.10, N = 6; Overload = 2.15 ± 0.35, N = 5, p = 0.01, unpaired t-test, Figure 1d). However, due to hypertrophy induced by overload, the global CD (Control = 920 ± 122 mm^{-2}, N = 6; Overload = 761 ± 196, N = 5, p = 0.13, unpaired t-test, Figure 1e) and modelled spatial tissue PO_{2} remained unchanged (Control = 11.5 ± 1.8 mmHg N = 6; Overload = 9.1 ± 3.1 mmHg, N = 5, p = 0.12, unpaired t-test, Figure 1f–j). Thus, improved fatigue resistance is likely due to increased efficiency in aerobic metabolism.

Overall, our physiological data confirm a shift to a slower, more fatigue-resistant phenotype following overload.

3.3 | Chronic functional overload reduces EDL motoneuron soma cross-section area

Motoneuron properties are matched to the muscle fibre types they innervate. Motoneurons innervating Type 1 muscle fibres are smaller than those innervating Ila, which are smaller than those innervating IIb/IIx (Burke, 1967). In the overload model, we found a shift in the distribution of motoneuron sizes towards smaller motoneurons, leading to an overall reduction in EDL motoneuron size following removal of TA (Control = 174 ± 248, N = 6, n = 285 vs. Overload = 1372 ± 188, N = 5, n = 260, p = 0.021, unpaired t-test, Figure 2a–d1). This shift to smaller-sized motoneurons corresponds to the shift in muscle phenotype.

3.4 | Overload has no effect on C-bouton innervation of EDL motoneurons

C-bouton synapses are terminals of the V_{0}C interneuron circuit responsible for task-specific amplification of motor output (Miles et al., 2007; Zagoraiou et al., 2009). Because the overload condition removed the contribution of the synergist, TA, to locomotion and necessitated increased force output from the EDL, we asked whether an increase in C-bouton synapses (Figure 3a) occurs in order to meet the increased demands placed on EDL motoneurons. We thus assessed the effect of overload on the density and size of C-bouton inputs.
There was no significant effect of overload on C-bouton density (Control = 3.4 ± 0.8 per 100 μm², N = 6, n = 289 vs. Overload = 3.3 ± 0.8 per 100 μm², N = 5, n = 259, p = 0.61, Figure 3b,b1) or mean area (Control = 25.7 ± 3.7 μm², N = 6, n = 289 vs. Overload = 23.3 ± 6.9 μm², N = 5, n = 259, p = 0.48, Figure 3c,c1). That is, there were no discernible changes to the presynaptic component of C-bouton inputs to EDL motoneurons.

3.5 | \( K_{\text{v}}2.1 \) channel density and area are unaffected by overload

Although we saw no change in presynaptic C-bouton characteristics, changes in the post-synaptic protein complex could alter synapse function. \( K_{\text{v}}2.1 \) channels are thought to be important for facilitating high-frequency motoneuron firing and are recruited for C-bouton amplification of motor output (Nascimento et al., 2020). In addition to the large clusters found at the C-bouton, \( K_{\text{v}}2.1 \) channels are found in smaller clusters distributed throughout the membrane (Figure 4a-d1; Muennich & Fyffe, 2004). We therefore assessed the influence of overload on expression of both C-bouton-associated and -unassociated \( K_{\text{v}}2.1 \) channels by masking \( K_{\text{v}}2.1 \) signal to the presynaptic VAChT signal.

Chronic functional overload did not significantly alter the density of \( K_{\text{v}}2.1 \) clusters either associated (Control = 0.43 ± 0.17 per 100 μm², N = 6, n = 135 vs. Overload = 0.36 ± 0.12 per 100 μm², N = 5, n = 85, p = 0.41, t-test, Figure 4e,e1) or unassociated (Control = 4.2 ± 1.1 per 100 μm², N = 6, n = 135 vs. Overload = 2.9 ± 2.4 per 100 μm², N = 5, n = 85, p = 0.26, Figure 4f,f1) with C-boutons. Similarly, there was no difference in mean surface area between control (associated: 12.4 ± 3.5 per 100 μm², Figure 4g,g1; unassociated: 3.0 ± 0.4 per 100 μm², N = 6, n = 135, Figure 4h,h1) and overload groups (associated: 10.1 ± 5.4 per 100 μm², p = 0.41, Figure 4f,f1; unassociated: 3.0 ± 0.6 per 100 μm², N = 5, n = 85, p = 0.26). The lack of change in density and surface area of \( K_{\text{v}}2.1 \) for both associated and unassociated \( K_{\text{v}}2.1 \) channels was reflected in the unchanged percentage of \( K_{\text{v}}2.1 \) localised to the C-bouton following overload (Control = 16.8 ± 16.8 per 100 μm², N = 6, n = 140 vs. Overload = 24.5 ± 21.2 per 100 μm², N = 5, n = 79, p = 0.15, Figure 4i,i1).

3.6 | SK3 channels are expressed in most EDL motoneurons but are unaltered by chronic functional overload

In addition to \( K_{\text{v}}2.1 \) channels, small conductance potassium channels (SK) are also clustered on the motoneuron post-synaptic membrane opposing C-boutons. SK2 and SK3 channels are responsible for the calcium-dependent potassium currents underlying the mAHP and, therefore, regulate motoneuron firing frequencies. Previous work has suggested that SK2 channels are expressed in all motoneurons, whereas in the specific pools studied (mainly tibial motoneurons), SK3 channels are selectively expressed in

![Figure 3](image-url)
slow motoneurons (Deardorff et al., 2013; Dukkipati et al., 2018). Although the EDL muscle is mainly comprised of fast fatiguable (FF) units, there are also fast FR and a small proportion of type I slow (S) units (Kissane et al., 2018). We reasoned that, given the shift to a more fatigue-resistant phenotype following chronic overload (Figure 1), there would be a corresponding upregulation of SK3 expression in motoneurons.

Interestingly, we found that out of all motoneurons assessed (regardless of condition) 227 of 285 (79%) expressed SK3 channel clusters (Figure 5a–e). In control animals, 130 of 141 cells expressed SK3...
(92%), while in the overload condition 97 of 144 motoneurons (67%, p = 9.5e-06) had SK3 expression. The lower proportion of SK3 in the overload condition may indicate that SK3 is in fact downregulated following chronic overload. However, in the neurons that expressed SK3, we found no difference between control and overload groups in either SK3 density (Control = 0.4 ± 0.04 per 100 µm², N = 6, n = 130 vs. Overload = 0.5 ± 0.3 per 100 µm², N = 5, n = 97, p = 0.65, Figure 5f) or area (Control = 7.4 ± 2.7 µm², N = 6, n = 130; vs. Overload = 7.0 ± 2.0 µm², N = 5, n = 97, p = 0.73, Figure 5g). Thus, while fewer cells were SK3 positive in the overload condition, there was no difference in SK3 expression between the conditions for cells that were positive.

In summary, we found that SK3 channels are not reliable binary markers for slow motoneurons in EDL and that there was a decrease in the proportion of SK3⁺ cells in the overload condition despite the shift to smaller cell sizes and decreased fatigability. Overall, though, we detected no significant effect of overload on presynaptic C-bouton terminals, or expression of either of the post-synaptic proteins assessed (SK3 and Kᵥ2.1). This suggests that the C-bouton complex is resistant to the increased functional demands of the innervated muscle.

4 | DISCUSSION

The results presented here demonstrate that chronic overload of the rat EDL muscle induces significant adaptations to muscle fibre capillary innervation and contractile properties. Specifically, 21 days functional overload resulted in an increase in EDL capillary-to-fibre ratio and fatigue resistance, paralleled by a decrease in twitch and tetanic force production. We reasoned that this shift to a slower phenotype in the overloaded EDL muscle may be reflected anatomically in their motoneuron characteristics. This was confirmed by a decrease in motoneuron soma cross-sectional area. We also hypothesised that a key spinal neuromodulatory input, C-boutons, would change so as to compensate for the increased neuromuscular demand during overload. However, while we did find a decrease in the proportion of SK3⁺ neurons, we were unable to detect significant adaptations in size or density of key components of the C-bouton complex (C-bouton synapse, SK3 and Kᵥ2.1 channels). Importantly, we also show that SK3, a suggested molecular marker for slow motoneurons (Dukkipati et al., 2018), was expressed in most EDL motoneurons, despite these being mainly fast type (Kissane et al., 2018).
4.1 | Increased EDL fatigue resistance following overload is paralleled by a reduction in motoneuron soma size

The overload-induced shift to a more aerobic phenotype in EDL muscle was reflected in the central portion of the motor unit, by a shift to smaller sizes for motoneurons in this condition. Motoneuron size is inversely related to input resistance and positively correlates with rheobase (Henneman, 1957), meaning changes in motoneuron size may lead to altered recruitment thresholds across the motor pool. Previous electrophysiological assessments of overloaded rat MG motoneurons demonstrated significant reductions in rheobase, associated with increased input resistance and a leftward shift in the frequency current (f-I) relationship in FF motoneurons, indicating that less input was required to activate these motoneurons (Krutki et al., 2015). Taken together, these changes in motoneuron properties would be matched to peripheral changes and mean that the probability of recruitment and firing rate is increased for a given synaptic drive in the overload compared to the control condition.

4.2 | C-bouton complex is largely unaltered by chronic functional overload

The C-bouton is a neuromodulatory synapse responsible for task-specific amplification of motor output, and is anatomically characterised by a dense aggregation of proteins at the opposing post-synaptic membrane (Conradi, 1969; Deardorff et al., 2014). C-bouton modulation increases the f-I slope of motoneurons through activation of m2AChRs and subsequent reductions in the amplitude and duration of the mAHP, mediated by SK channels (Miles et al., 2007). C-bouton SK2 channels are expressed in almost all motoneurons regardless of type, whereas previous work in extensor motoneurons shows that SK3 channels are preferentially found on S type motoneurons and thus are likely responsible for larger mAHP conductances (Deardorff et al., 2013; Dukkipati et al., 2018). Surprisingly, we found that most EDL motoneurons expressed large SK3 clusters to some degree. Thus, in the rat EDL motor pool, it would seem that SK3 expression cannot be reliably used as a binary molecular marker to identify slow motoneurons.

There was a decrease in the proportion of SK3⁺ motoneurons in the overload condition. Given prior evidence that SK3 channels are associated with longer AHPs and slower phenotypes (Deardorff et al., 2013), the “loss” of SK3 expression could be considered to indicate a shift to a faster phenotype—a finding in contrast to our other findings. However, considering that we did not detect a decrease in SK3 expression (cluster size or density) in the SK3-positive cells, it is possible that the motoneurons that presumably ‘lost’ SK3 clusters had lower expression levels before overload. Taken together, it is not clear whether the reduction in the number of SK3⁺ neurons had significant functional implications.

We found no effect of overload on the size or density of C-boutons, Kv2.1 channels or SK3 channels on EDL motoneurons, suggesting that both pre- and post-synaptic components of the synaptic complex are largely unaffected by a stimulus which induced changes in muscle fibres and motoneuron size. This was unexpected, especially for SK3, as the motoneuron mAHP has been shown to be increased following chronic overload (Krutki et al., 2015). It is possible that there were changes in other SK isoforms, but given the propensity of SK3 channels to be differentially expressed in motoneuron types, we focused on them.

There are several potential reasons why we did not detect adaptions in C-bouton complexes. Firstly, certain motoneuron properties, such as C-bouton synapses, might be somewhat resistant to plasticity (Chalmers et al., 1991). Secondy, C-bouton synapses may already be organised to meet the increased demand following overload, and so even if they were more active, there may be no need for anatomical adaptation. Thirdly, C-bouton innervation may be sensitive to certain types of stimuli, but overload is either not appropriate or sufficient to stimulate plasticity. In this vein, Kv2.1 organisation on the motoneuronal membrane seems to be affected mainly by extremely high or pathological levels of activity (Romer et al., 2019; Romer et al., 2014), whereas our overload model induces a physiological load increase. Moreover, it is possible that Kv2.1 clustering was modulated during the early phases of overload, when the stimulus is greatest, and returned to ‘normal’ once motor adaptation was complete. Furthermore, we now know that Kv2.1 channels make physical links with the SSC via vesicle associated membrane protein (VAMP)-associated proteins (VAPs; Deardorff et al., 2021; Johnson et al., 2018; Kirmiz et al., 2018). Although the physiological relevance of this structural role is unknown, evidence from brain neurons suggests that the Kv2.1-VAP interaction maintains tight plasma membrane–SSC junctions (Johnson et al., 2018). Because many of the proteins located at the C-bouton are Ca²⁺ modulated, and the region of the SSC functions as a local Ca²⁺ microdomain, C-bouton activity could affect associated protein function (SK, Kv2.1 etc) by modulating Ca²⁺ flux in the microdomain (Deardorff et al., 2021). Thus, dispersal of Kv2.1 would be counterproductive in response to overload, as it would theoretically limit C-bouton function. However, it is important to note that the molecular mechanisms downstream of C-bouton activation of m2AChRs have yet to be described.

In conclusion, our results show that a shift to a slower phenotype in EDL muscles is accompanied by a reduction in EDL motoneuron size, which perhaps allows EDL motor units to be recruited with less synaptic drive. The C-bouton complex, a key neuromodulatory synapse, is, however, anatomically unaffected by overload, suggesting that adaptation of these synapses was not necessary. Whether C-bouton function adapts in other environmental conditions remains to be seen.

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AUTHOR CONTRIBUTIONS
R.W.P.K performed muscle experiments and analyses, spinal cord tissue preparation, interpreted results, wrote and edited the manuscript. A.G performed spinal cord experiments, acquired confocal images, analysed images using IMARIS, interpreted results, wrote and edited the manuscript. P.G.T performed muscle experiments and edited the manuscript. S.C. & S.E provided technical advice and edited the manuscript. R.M.B helped design the study, interpreted the data, wrote and edited the manuscript. C.C.S conceptualised and designed the study, supervised experiments, analysed data, wrote and edited the manuscript.

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