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Identity, developmental restriction and reactivity of extralaminar cells capping mammalian neuromuscular junctions

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Short Title: Capping Cells at Mammalian NMJ
Summary

Neuromuscular junctions are normally thought to comprise three major cell types: skeletal muscle fibres, motor neurone terminals and perisynaptic terminal Schwann cells. Here we studied a fourth population of junctional cells in mice and rats, revealed using a novel cytoskeletal antibody (2166). These cells lie outside the synaptic basal lamina but form caps over neuromuscular junctions during postnatal development. NMJ-capping cells also bound rPH, HM-24, CD34 antibodies and cholera toxin B-subunit. Bromodeoxyuridine incorporation indicated activation, proliferation and spread of NMJ-capping cells following denervation in adults, in advance of terminal Schwann cell sprouting. The NMJ-capping cell reaction coincided with expression of tenascin-C but was independent of this molecule since capping cells also dispersed after denervation in tenascin-C null mutant mice. NMJ-capping cells also dispersed after local paralysis with botulinum toxin and in atrophic muscles of transgenic R6/2 mice. We conclude that NMJ-capping cells ("kranocytes") represent a neglected, canonical cellular constituent of neuromuscular junctions where they may play a permissive role in synaptic regeneration.
Introduction

Neuromuscular junctions (NMJ) are interfaces between components of two main cell types: motor axon terminals and motor endplates of skeletal muscle fibres. In adult mammals and most other vertebrates, each motor endplate is normally innervated by a single motor axon terminal and in most muscle fibres this NMJ is constrained to less than 0.1% of the muscle fibre cell surface (Sanes & Lichtman, 1999; Beeson et al., 2006). The strength of synaptic transmission at each NMJ in a motor unit virtually guarantees that all the muscle fibres supplied by a motor neurone contract in response to every action potential conducted into its motor nerve terminals (Wood & Slater, 2001). NMJ’s are therefore crucial structures in the intercellular signalling processes that control movement and behaviour. Degeneration of motor nerve terminals occurs rapidly after motor axonal injury, axonal transport block or neuromuscular paralysis (Miledi & Slater, 1970; Brown et al., 1980; Hudson et al., 1984) and in early stages of motor neurone diseases such as amyotrophic lateral sclerosis (ALS; Fischer et al., 2004; Pun et al., 2006; David et al. 2007). Compensatory responses to such denervation include sprouting from intact neighbouring axons and regeneration of damaged axons (Brown & Ironton, 1978; Barry & Ribchester, 1996; Costanzo et al., 1999, 2000; Schaefer et al., 2005). Understanding the cellular organisation of NMJ’s and the mediators of their plasticity is therefore important: not only for the insights it provides into fundamental cell biology but also for identifying suitable cellular and molecular targets for effective treatment for ALS and other neuromuscular diseases.

Motor neurones and muscle fibres are not the only cell types present at an NMJ. In the endplate region, myelinating Schwann cells ensheath intramuscular axon collaterals up to the last heminode of Ranvier, ensuring high-fidelity conduction of action potentials (Court et al., 2004). The nerve terminal itself is capped by non-myelinating terminal Schwann cells (tSC; Robertson, 1956; Birks et al., 1960; Kang et al., 2003; Hayworth et al, 2006). These cells are contained in a continuation of the synaptic basal lamina and drape motor terminal branches and synaptic boutons, in register with other pre- and postsynaptic specialisations. Though not essential for the initial formation of neuromuscular synapses, tSC are required for maintenance of preterminal axon structure and function during development; for long-term maintenance of neuromuscular transmission in adult life; and they appear to play a path-finding role in compensatory sprouting responses following nerve injury and repair (Jahromi et al, 1992; Son & Thompson, 1995; Trachtenberg & Thompson, 1996; Castonguay and Robitaille, 2001; Reddy et al, 2003; Court et al., 2008). Thus, together, motor nerve terminals, muscle fibres and terminal Schwann Cells are
generally considered the three essential cellular elements that constitute the vertebrate neuromuscular junction (Sanes & Lichtman, 1999; Hughes et al., 2006).

Here we report strong association of a fourth cell type with NMJ’s. This arose from a serendipitous initial finding: whole-mount immunofluorescent staining of skeletal muscle with a novel polyclonal antibody (2166) produced vivid cytoskeletal labelling of a sub-set of cells capping the NMJ but lying outside the synaptic basal lamina. These NMJ-capping cells may correspond to “endoneurial cells” first identified in the neighbourhood of NMJ’s in pioneering ultrastructural studies of the NMJ by Robertson (1956), or to cells of similar appearance referred to by Weis et al. (1991) as “perisynaptic fibroblasts”. However, the data here suggest a much tighter association of NMJ-capping cells than that indicated by these previous studies. NMJ–capping cells become restricted to neuromuscular junctions during postnatal development but within 24 hours of denervation or paralysis in adults, they proliferate and spread throughout the perijunctional region, ahead of reactive sprouting of either terminal Schwann cells or regenerating motor axons. Taken together, these findings suggest that mammalian neuromuscular junctions comprise four cell types, not three, and that NMJ–capping cells should be considered as integral constituents of neuromuscular junctions.

Results

2166-antibody stains NMJ-capping cells

Antibody 2166 was raised in rabbits against an epitope of Tspan-2, an oligodendrocytespecific tetraspanin protein of 25kDa (Birling et al, 1999; Terada et al, 2002). The original intention was to use this antibody to map oligodendrocyte disposition and lineage in the brain. However, preliminary analysis using western blots suggested the antibody recognised a protein of 47kDa, which was not Tspan-2 (see Methods). Immunostained sections of brain indicated the antibody did not stain oligodendrocytes either. Since the potential utility of the 2166 antibody did not depend on knowledge of its primary antigen we proceeded to test it for potential binding to other neural tissues, including neuromuscular junctions, mainly anticipating the possibility of peripheral myelin or terminal Schwann cell staining.
Remarkably, 2166-immunostaining of whole-mount preparations of the mouse triangularis sterni (TS) muscle revealed a sub-population of cells, some of which were localised to neuromuscular junctions, where they formed caps over the motor nerve terminals. The 2166 antibody evidently stained the cytoskeleton in these capping cells (Fig. 1A-C; and see below) and frequently took the form of a lasso around the cell nucleus, appearing to extend into numerous filopodial or lamellipodial cellular processes. In most cases these extended over and beyond the boundaries of the motor endplate, as indicated by binding of rhodamine-conjugated α-Bungarotoxin (TRITC-α-BTX), a specific ligand for postsynaptic AChR. We subsequently observed 2166-positive cells capping neuromuscular junctions in several other skeletal muscles in mice and rats (diaphragm, soleus, EDL, FDB, and DL). Thus, we refer to the NMJ-localized 2166-positive cells throughout the Results section of this report as “NMJ-capping cells” (see Discussion, for justification of a proposed alternative name: “kranocytes”).

Although the disposition of NMJ-capping cells was unequivocal, several other forms of 2166-immunopositive cells were observed beyond the junctional region in confocal micrographs, including the optical sections above and below the NMJ’s. However, there was only enrichment of 2166-positive cells in the vicinity of intramuscular nerves and in contrast to the morphology of NMJ-capping cells, virtually all of the 2166-positive extrajunctional cells had a bipolar form, with extensions running along the longitudinal axis of the muscle fibres (Fig. 1D). 2166-positive cells showing a similar bipolar form were also associated with intramuscular capillaries (Fig. 1E).

The only clusters of capping-cells were found around motor endplates. We counted the number of 2166-positive cells at neuromuscular junctions in adult TS muscles, counterstained with the nuclear marker DAPI (n=5 muscles, 100 neuromuscular junctions analysed). All neuromuscular junctions were capped by at least one 2166-positive cell. About 25% exhibited two, with fewer than 10% showing three (Fig. 1F). Outside the region of the NMJ’s, 2166-positive cells did not show any discernible clustering.

NMJ-capping cells show a distinct immunocytochemical profile

Immunocytochemical analysis (Table 1) clearly resolved NMJ-capping cells from terminal Schwann cells and suggested a distinct identity. First, double immunostaining with 2166 antibody and antibodies for either GFAP or nestin showed no co-localisation, demonstrating that
NMJ-capping cells are not terminal Schwann cells. We confirmed this by 2166 immunostaining in S100-eGFP mice (Zuo et al., 2004) in which all myelinating and terminal Schwann cells are endogenously fluorescent (Fig. 2A). The 2166 immunostaining pattern was completely different and extended beyond the boundaries of the Schwann cell plasma membrane. Immunostaining for M-cadherin, NCAM or desmin also did not label NMJ-capping cells, suggesting that they were not muscle satellite cells (Sanes et al., 1986; Fukada et al., 2004). Finally, absence of F4/80 immunostaining appears to rule out any identity with macrophages (Austyn & Gordon, 1981).

We next examined whether NMJ-capping cells might be a specialised form of fibroblast. The exact disposition of “perisynaptic fibroblasts” is unclear from previous reports, since they were depicted diagrammatically as near to, but not overlaying, neuromuscular junctions (Murray & Robbins, 1982; Gatchalian et al., 1989; Weis et al., 1991). No discernible immunofluorescence of NMJ-capping cells was detected using Thy-1 antibody, although axons, which also express Thy-1 antigen were immunolabeled (Reynolds & Woolf, 1992; Feng et al., 2000; Van der Putten et al., 2000). There were other Thy-1 positive cells near the NMJ but these did not include the 2166-positive NMJ-capping cells (Supplementary Fig. 1A). These positive controls confirmed the antibody’s specificity and affinity. Thus, 2166 cells may not correspond to the Thy-1 positive perisynaptic fibroblasts previously reported.

As a further test, we immunostained rat TS muscles using a mouse antibody against the rat isoform of prolyl-4-hydroxylase (rPH), an enzyme regulating the synthesis of collagen (Aiba et al., 1994; Smith et al., 1998). Positive staining for rPH was localised to 2166 positive cells above the NMJ although it did not overlap with 2166 immunostaining (Fig. 2B). Scrolling through individual optical slices (Z-series; see for example Supplementary Movies 1 and 2) showed rPH and 2166 staining were in the same cells. Thus, rPH is an enzyme evidently located in an intracellular compartment, possibly the endoplasmic reticulum (Nissi et al., 2001), near to but not overlapping the cytoskeletal components stained by 2166 antibody. Since 2166 antibody appeared to label specific cytoskeletal components, it is not surprising that it failed to colocalise with rPH. We did not examine whether rPH-positive fibroblasts in other tissues are also 2166-positive but it seemed clear that at least some cells with fibroblast-like characteristics also express the 2166 antigen.
The pattern of labelling of NMJ-capping cells suggests that the 2166 antigen is spatially quite restricted: for instance, there were substantial parts of the cells -mostly near the periphery -where there was no 2166 labelling. To test whether the 2166 staining colocalises with other components of the cytoskeleton, we co-immunostained NMJ-capping cells with actin, vimentin or tubulin. The 2166 immunostaining was more restricted, with no strong co-localisation with these other cytoskeletal markers (data not shown).

Next, while searching cytological literature for evidence of NMJ-capping cell disposition, we noted the strong resemblance of 2166 antibody staining of the NMJ-capping cells to staining for neuregulin (GGF II) at NMJ, reported by Trinidad et al (2000). These authors used the HM-24 antibody and attributed the staining pattern to terminal Schwann cells. However, in our hands, HM-24 positive staining was not observed in S100-eGFP positive terminal Schwann cells (Fig. 2B). It was not possible to double-stain NMJ-capping cells with 2166 and HM-24 antibodies because both were raised in rabbits but the HM-24 immunostaining pattern extended beyond the boundaries of the tSC’s. Interestingly, the pattern of HM-24 staining also appeared similar to the cytoskeletal labelling pattern of the 2166 antibody (see also Figure 4D3 in Trinidad et al., 2000), which is perhaps surprising for neuregulin distribution. It is puzzling that Trinidad et al did not report seeing HM-24 staining in cells lying outside the synaptic basal lamina (see below). However, the present data suggest that junctional HM-24 positive cells appear to be identical to the NMJ-capping cells that we identified using the 2166 antibody.

Next, we looked for possible cell-surface markers of NMJ-capping cells. A stem cell population located in interstitial spaces of skeletal muscle, outside the basal lamina, has been identified previously and isolated by immunostaining for the hematopoietic stem cell marker CD34 (Torrente et al, 2001;Tamaki et al, 2002). We found that the surfaces of NMJ-capping cells showed uniform, faint immunostaining for CD34 (Fig. 3A,B). These preparations revealed additional aspects to the disposition of NMJ-capping cells: they frequently covered and extended beyond the NMJ and contacted the muscle fibre outside the end-plate perimeter, sometimes extending processes between adjacent motor endplates (Fig. 3B). This analysis also confirmed that the 2166 antigen was contained within the cytoskeletal framework of CD34-positive NMJ-capping cells. Orthogonal projections of the NMJ-capping cell showed that they were separated from the muscle fibre surface by about 3 µm and therefore that they were located outside the
synaptic basal lamina (Fig. 3B, Supplementary Figure 2). We confirmed this in transverse muscle sections co-immunostained with anti-laminin and 2166 antibodies, together with TRITC-α-BTX staining of ACh receptors (Fig. 3C).

Finally, we found that fluorescent conjugates of cholera toxin B-subunit (CTB) highlighted the plasma membranes of NMJ-capping cells (Fig. 3D). We chose this reagent because it binds to GM1 gangliosides, which are absent from terminal Schwann cells (Halstead et al., 2005). Like CD34 staining, membrane staining with CTB extended beyond the cytoskeletal staining with 2166 and also revealed several instances of fine cellular processes extending between motor endplates (Supplementary Movies 3, 4, and 5). We did not attempt double-labelling with CD34 antibody and CTB but they otherwise appeared to have similar distributions.
Ultrastructural disposition of NMJ-capping cells

We noted that many electron micrographs of NMJ’s in textbooks (e.g. Peters et al., 1991) crop images of NMJ’s to exclude cells other than tSC’s, nerve terminals and muscle fibres. However, original published transmission electron micrographs of NMJ’s in frogs, lizards, chickens, mice, rats and humans frequently show slender cellular processes overlying the other principal cell types of the NMJ (Robertson, 1956; Desaki & Uehara, 1981; Connor & McMahan, 1987; Slater et al, 1992; Connor, 1997; see also Fig.4D). Thus, to extend the confocal microscopy data above we made electron micrographs from 300 ultrathin (75-100nm) serial-sections of one randomly-selected NMJ in a mouse TS muscle. We highlighted the profiles of the muscle fibre, motor nerve terminal, terminal Schwann cells and two putative NMJ-capping cells identified at this NMJ (Fig. 4A,B).

Volume-rendering of a three-dimensional reconstruction of all of these cellular components confirmed the shape and relationship of the NMJ-capping cells with the other cellular components of the NMJ, as observed using light microscopy (Fig. 4C; Supplementary movies 6 and 7). For instance, the EM reconstruction confirmed the impression from confocal microscopy, indicating that the cellular processes of NMJ-capping cells are quite laminar.

Confocal microscopy suggested that NMJ-capping cells themselves lack a basal lamina (see Fig 3C). However, micrographs like that shown in Fig. 4D indicate that capping cells are tightly ensheathed by a thin layer of extracellular matrix (See also Supplementary Figure 3). Previous studies suggest that perisynaptic fibroblasts are also rich in endoplasmic reticulum. This appears also to be the case, at least for the capping cell shown in Fig. 4D/Supplementary Figure 3.

In summary, whether NMJ-capping cells are identical to “perisynaptic fibroblasts” described previously by others remains open. The data reported above rather suggests that NMJ-capping cells represent a distinct sub-population of cells, defined by their shape, strong association with neuromuscular junctions, and a distinctive molecular profile as indicated by staining with a unique combination of five cytological markers: antibodies 2166, rPH, HM-24, CD34, and with CTB.
Development and Plasticity of NMJ-capping cells

The remarkable localisation of NMJ-capping cells, revealed here both by light microscopy of whole-mounts and by their reconstruction from electron micrographs, naturally led us to enquire about their possible functions. We focused initially on the patterns of 2166-immunostaining during early postnatal development and then we studied the effects of muscle atrophy, induced either by denervation, paralysis or disease. Finally, we investigated an association with tenascin-C, an early molecular marker of denervation, seeking insight into the possible mechanism of NMJ-capping cell plasticity.

2166-positive cells become restricted to NMJ postnatally

We immunostained TS muscles of postnatal mice aged 1-28 days (P1-P28). Neuromuscular junctions in these preparations were co-stained using neurofilament antibodies and fluorescent TRITC-α-BTX. At P1, 2166-positive cells were homogeneously distributed along the TS muscle with no clear pattern of localisation to NMJ’s (Fig. 5A). Between P5 and P10, 2166-positive cells became more restricted to endplate bands adjacent to intramuscular nerve branches but the degree of refinement between these times was not compelling (Fig. 5B-C). The pattern changed dramatically after P10 and by P28 2166-positive cells were highly restricted to the neuromuscular junctions, as shown in Figures 1-3.

2166-positive cells spread after denervation, paralysis or muscle atrophy

Next, we partially denervated TS muscles in adult mice, then immunostained them 1-6 days later with nestin antibody, a marker of reactive terminal Schwann cells (Hayworth et al, 2006; Kang et al., 2007). As expected, at innervated neuromuscular junctions nestin was expressed only in muscle fibres; terminal Schwann cells remained nestin-negative for at least 2 days after denervation (Fig. 5D). However, the images suggest that profusion of 2166-positive cells was evident in the denervated endplate regions of TS muscles within one day of intercostal nerve injury; and many of these cells appeared to have grown processes interconnecting NMJ’s (Fig. 5E, Supplementary Figures 1 & 4). This response was local to denervated endplates: innervated NMJ’s in the same partially denervated muscles were covered by NMJ-capping cells confined to the AChR cluster region, as in unoperated muscles. Some of the reactive, nestin-positive Schwann cell processes that formed 1-3 days later evidently became associated with the 2166-positive cell processes interconnecting motor endplates (Fig. 5F, Supplementary Fig. 5; 41
junctions observed with this association in 4 muscles). We were unable to apply immunocytochemistry to study the relationship between S100-positive Schwann cells and 2166-positive NMJ capping cells in mice, because both the 2166 antibody and commercial S100 antibody were raised in rabbits (hence unresolvable with anti-rabbit secondary antibodies). However, reactive Schwann cells in a partially denervated rat 4th deep lumbrical muscle stained using a mouse S100 antibody showed Schwann cell sprouts aligned with inter-endplate processes from 2166-positive NMJ-capping cells (Supplementary Fig. 1B, Supplementary Figure 4).

The above findings suggested two hypotheses: first, that NMJ-capping cells selectively proliferate after nerve injury and second, that these cells play an instructive role in sprouting by first activating terminal Schwann cells and then guiding their sprouts to denervated endplates. The data below support the first hypothesis but not the second, suggesting that the NMJ-capping cell response to denervation, paralysis or muscle atrophy probably occurs independently of the terminal Schwann cell reaction.

To test the first hypothesis we denervated TS muscles of seven adult mice and injected them with bromodeoxyuridine (BrdU) either 1 day (three mice) or 3 days (four mice) later. Incorporation of BrdU by NMJ-capping cells was assessed by immunofluorescence. Nuclei of NMJ-capping cells were distinguished from others (muscle and Schwann cell nuclei) by the characteristic 2166 staining forming lasso-like structures around the nuclei (see also Fig. 1, for example). We expected that the BrdU pulse would only mark cells within the first hour of injection, thus labelling only a small fraction of the 2166-positive cells that may have proliferated. We were therefore not surprised by the apparent absence of BrdU labelling at 1d; and by three days, only about 7% of all 2166-positive cells were BrdU positive. However, more than 65% of these cells were located within 40 µm of the neuromuscular junctions (Fig. 5G). These observations suggest that NMJ-capping cells were stimulated to spread then divide as an early consequence of endplate denervation. However, from these data we cannot exclude the possibility that the 2166-positive cells in the denervated endplate region may become supplemented by invasion of reactive cells from outside the endplate zone as well.
To resolve whether the spread and proliferation of NMJ-capping cells could constitute an instructive signal that triggers the Schwann cell reaction to denervation (Son et al, 1996), we took advantage of the slow response of Schwann cells and axons in the TS muscle to neuromuscular paralysis (unpublished observations), similar to that reported for “fasyn” types of muscle by Pun et al (2002). We injected botulinum toxin type A (BoNT/A) into the interstitial spaces between the TS and intercostal muscles. The animals were sacrificed 1-6 days later. Stimulation of the intercostal nerves supplying the isolated muscles gave no contractile responses, confirming the neuromuscular block, whereas stimulation of saline-injected control muscles produced vigorous muscle contractions.

The responses of NMJ-capping cells and tSC in BoNT/A-injected muscles were quite different from those in denervated muscle. After only one day of muscle paralysis the profusion of 2166-positive cells in the junctional region was not discernibly different from that induced by denervation. However, tSC remained nestin-negative and failed to show discernible evidence of sprouting for at least 6 days (Fig. 6A-B). The 2166-positive cells were clearly dispersed over and between neuromuscular junctions, as in denervated muscle (compare with Fig. 5). Since the Schwann cell and NMJ-capping cell reactions were dissociated in time by BoNT/A paralysis, we conclude that the role of NMJ-capping cells in adaptive responses to nerve injury may be to lay down a substrate that could be conducive to reactive Schwann cell outgrowth, but this does not necessarily trigger or instruct tSC’s to react. These cells evidently respond to cues arising from denervated or paralysed muscles independently of those that trigger earlier reaction of NMJ-capping cells.

Since denervation and paralysis had distinct temporal effects on terminal Schwann cells in TS muscles but both were sufficient to induce NMJ-capping cell proliferation and spread, we next asked whether inactivity was strictly necessary for this reaction or whether some other, intrinsic signal from muscle fibres was sufficient. To test this we examined NMJ-capping cells in TS muscles of R6/2 transgenic mice (normally used to model Huntington’s disease; Mangiarini et al, 1996). These mice develop profound, constitutive muscle atrophy and show several other characteristics of denervated muscle even though their muscle fibres are fully innervated and they show no physiological evidence of paralysis. In fact, there are very few discernible abnormalities in the morphology of either Schwann cells or motor nerve terminals at most
(>95%) of the NMJ’s in these mice (Ribchester et al 2004). Muscle atrophy in R6/2 mice is therefore likely to be either myogenic or to have some other non-neural explanation. Moreover, Schwann cells and axons retain their capacity to sprout when muscles in R6/2 mice are surgically denervated, as in normal mice (Ribchester et al 2004). We therefore used 2166 antibody to immunostain TS muscles dissected from two 15-week old R6/2 mice, towards the endstage of disease. As expected (Carter et al, 1999) both of these mice showed decreased body weight and showed significant muscle atrophy. If denervation or paralysis were necessary for the NMJ-capping cell reaction, we should have expected to see no differences in their distribution in R6/2 mouse muscles compared with wild-type mice, in spite of their muscle atrophy.

Surprisingly, the appearance and disposition of 2166-positive cells in R6/2 mice were similar to those observed following either denervation or paralysis of wild-type muscle (Fig. 6C,D). These observations support the interpretation that activation of NMJ-capping cells does not inevitably or invariably lead to activation of Schwann cells. The data also suggest that denervation and paralysis, though sufficient (see above), are not necessary for the induction of the NMJ-capping proliferation and spread. Thus, taken together, the NMJ-capping cell reaction that evidently occurred in all three circumstances – denervation, paralysis and atrophy (without paralysis) – suggests that factors triggering or triggered by atrophy of muscle fibres are responsible for their reactivity, rather than factors associated directly with reactive terminal Schwann cells.
What molecules cue the NMJ-capping cell reaction?

Previous data suggest that reactive perisynaptic fibroblasts express several adhesion molecules including tenascin, NCAM, fibronectin and heparan sulfate proteoglycan and that perisynaptic fibroblasts may be one source of tenascin (Covault & Sanes, 1985; Sanes et al, 1986; Gatchalian et al, 1989; Weis et al, 1991). These molecules have also been implicated in nerve sprouting and regeneration (Cifuentes-Diaz et al, 1998). Confirming this, we observed no positive immunostaining for tenascin-C in innervated neuromuscular synapses of wild-type mice but strong immunostaining for this molecule at neuromuscular junctions denervated for three days (Fig. 7A,B). After 6 days, tenascin-C expression had also spread from the neuromuscular perimeter and 2166-positive cells lay in register with these tenascin-C immunopositive regions (Fig. 7C).

Next, we examined the pattern of tenascin-C immunostaining following TS muscle denervation in tenascin-C null mutant mice. We carried out X-gal staining of denervated muscles in a group of heterozygous tenascin-C null-mutant mice in which the deleted sequence was substituted with a lac-Z reporter gene (Saga et al, 1992; Garci on et al, 2004). As expected, we found no positive lac-Z positive staining in innervated regions of TS muscles from these mice. However, 5 days after nerve section there was strong positive staining of the intramuscular axon and motor endplate region only: areas expected of Schwann cells or endoneurial or perineurial fibroblasts, and also consistent with tenascin-C expression by NMJ-capping cells (Fig. 7D,E). However, there was no compelling evidence that other 2166-positive cells expressed tenascin-C (compare Fig. 7C with 7D,E). Finally, we immunostained TS muscles with 2166 antibody in four homozygous tenascin-C null-mutant mice before and after intercostal nerve section. In unoperated muscles, NMJ-capping cellular localisation was indistinguishable from wild-type controls, suggesting that tenascin-C is not necessary for accumulation of these cells to NMJ’s during postnatal development (compare Fig. 7F with Fig. 1A-D and Fig. 5E,F). There was no discernible difference in the NMJ-capping reaction within the first 48 hours of denervation either (Fig. 7G).
We therefore conclude that expression of tenascin-C, although normally associated with the spread of NMJ-capping cells following denervation, is neither required for the initial localisation of NMJ-capping cells to endplates during development, nor for their proliferation and spread following nerve injury. Presumably, other molecules – or combinations of molecules – mediate the NMJ-capping cell reaction.

**Discussion**

The results suggest that mammalian NMJ’s normally comprise four cell types: these canonical constituents being muscle fibre, motor neurone, terminal Schwann cell and the NMJ-capping cell: a neglected cell-type that merits further investigation as a potential moderator of neuromuscular synaptic development, maintenance and plasticity. Evidence for this may be summarised as follows; then we discuss in more detail the identity of NMJ-capping cells; their possible significance of their immunocytochemical profile; and their development and plasticity.

First, serendipitously, we found that antibody 2166 immunostains a distinct population of cells capping NMJ’s, lying outside the synaptic basal lamina but co-localised with terminal Schwann cells, motor nerve terminals and muscle fibres at motor endplates. Confocal microscopy of whole mounts immunostained with the 2166 antibody and other markers were pivotal in demonstrating the consistent presence and association of NMJ-capping cells with motor endplates. It is perhaps important to note that estimates of the numbers of tSC’s at motor endplates should therefore take into account the evidence that most NMJ also localize 1-2 kranocytes (see data in Figure 1).

Second, NMJ-capping cells become restricted to the endplate region during postnatal development but react to either denervation or paralysis in adults by proliferation and spread from endplates. This occurs more rapidly than the reactive response of terminal Schwann cells. NMJ-capping cells also take the lead in forming bridges (or consolidating existing ones; see Figure 3) between neuromuscular junctions following muscle denervation or paralysis, ahead of terminal Schwann cells. This novel finding may be significant for understanding mechanisms of compensatory nerve sprouting and regeneration. NMJ-capping cells are also dispersed from NMJ in a disease model (R6/2 transgenic mice), where there is muscle atrophy but no paralysis or denervation. Since the structure and function of motor nerve terminals and terminal Schwann cells appear largely normal in these mice, a parsimonious explanation is that the signalling
mechanism that induces the proliferation and spread of NMJ-capping cells derives from
denervated, paralysed or atrophic muscle fibres rather than any other cell type at NMJ.

Third, while the above suggest that one of the functions of NMJ-capping cells may be in repair
of damaged neuromuscular connections, the data also suggest that the NMJ-capping cell reaction
does not require or depend on tenascin-C, one of the early molecular markers previously
associated with axonal sprouting and regeneration.

The identity of NMJ-capping cells
The present data underscore the importance of a pioneering ultrastructural study by Robertson
(1956) who identified “endothelial” cells associated with the NMJ (see Text Fig 1 in Robertson,
1956). The consistency of this association and the potential role of these cells in maintenance of
the normal structure and function of the NMJ has been somewhat neglected since. However,
similar non-neural cells were identified in scanning electron micrographs by Desaki & Uehara
(1981), who described them as “flattened stellate cells of unknown nature”. Other investigators
recognised the association of fibroblast-like cells to NMJ, including Murray & Robbins (1982),
McMahan and colleagues (Connor & McMahan, 1987; Connor, 1997), Caroni & Schneider
(1994); and Sanes and colleagues, in their description of “perisynaptic fibroblasts” (Covault &
Sanes, 1985; Sanes et al, 1986; Connor & McMahan, 1987; Connor, 1997; Gatchalian et al, 1989;
Weis et al, 1991). Perhaps perisynaptic fibroblasts and NMJ-capping cells are the same (and
have identity with the endothelial cells of Robertson, 1956) but this remains unproven.
Nevertheless, the parallels are striking: like “perisynaptic fibroblasts” the capping cells
proliferate in response to denervation; they are fibroblast-like; and they appear to change their
gene expression in response to nerve injury. It is therefore rather surprising that reviewers have
been reluctant to depict these cell types as essential components of the NMJ (Sanes & Lichtman,
1999; Hughes et al, 2006).

Taking the present and previous observations together, we suggest that the NMJ-capping cells
ought now to be considered an integral cellular component of neuromuscular junctions and
therefore represented as such in diagrammatic depictions of the neuromuscular junction (e.g. Fig.
8A). At this time we suggest naming these capping cells descriptively, as neuromuscular
“kranocytes” (from Greek, κρανος, meaning helmet), which acknowledges their main
morphological characteristics. We use this nomenclature for the remainder of the present discussion.

**Immunocytochemical profile of kranocytes**

We screened kranocytes using a panel of antibodies, including those previously associated with different cell types in skeletal muscle. Of the antibodies tested, HM-24, prolyl-4-hydroxylase (rPH) and CD34 gave positive staining. Kranocytes also bound fluorescent cholera toxin B-fragment (CTB). The CD34 and CTB labels suggest a more complete covering of the NMJ by the capping cells than 2166 (or HM-24) staining. The explanation appears to be that 2166 stains the non-uniform distribution of a cytoskeletal protein whereas CD34 antibody and CTB label the plasma membranes of these cells.

Staining with HM-24 antibody revealed an intriguing contrast to a previous study of the localisation of the antigen to NMJ's. Trinidad et al (2000) reported HM-24 (ie neuregulin) immunoreactivity to be within terminal Schwann cells. However, our observations based on immunostaining in S100-GFP expressing transgenics (see Figure 2), suggest that kranocytes are HM-24 positive but tSC are not. The methods we used differ, principally in our use of triangularis sterni muscle whole-mounts rather than tissue sections. But since Trinidad et al (2000) did not consider or rule out the possibility that HM-24 antibody might have immunostained a fourth cell type, we suggest their conclusion may have been mistaken. The localization of neuregulin (or a neuregulin-like molecule) to kranocytes rather than tSC could have important implications for understanding the development and maintenance of neuromuscular junctions so it will be important to fully resolve this issue.

The enzyme rPH is involved in the synthesis of collagen (Pihlajaniemi et al, 1991) but cell types other than fibroblasts, including endothelial cells and myelin-forming Schwann cells, also express rPH (Singh et al, 1997; Vanderwinden et al 1999). Kranocytes were also immunopositive for CD34 antigen, previously identified as an endothelial or hematopoietic stem cell marker (Krause et al, 1994). CD34-positive/rPH-positive haematopoietic stem cells may give rise to fibroblasts (Lee et al, 2000; Zulli et al., 2005). It might therefore be interesting to investigate further the pluripotential (ie stem-cell) capacity of neuromuscular kranocytes, especially in light of a recent report that fibroblasts may be reprogrammed into a fully pluripotential state following ectopic expression (via retroviral vectors) of the transcription
factors Oct-4, Sox-2, c-Myc and Klf4 (Wernig et al., 2007; Takahashi et al., 2007).
The surface expression of the CTB receptor (a GM1 ganglioside) suggests another way that more
information might be obtained about the normal functions of these cells. For instance, ablation of
terminal Schwann cell at the neuromuscular junction can provide information about their short
and long term functions in sustaining motor nerve terminals (Reddy et al, 2003; Halstead et al,
2004, 2005). In preliminary experiments we found that antiganglioside antibody DG2 ablates
motor nerve terminals, terminal Schwann cell and NMJ-capping cells by complement-mediated
cell lysis (unpublished observations). If a more selective ablation of kranocytes becomes
possible, it will be interesting to examine the long-term effects of this on neuromuscular
development, maintenance and repair.

**Developmental and plasticity of kranocytes**

2166-positive cells become restricted to the neuromuscular contact zone during postnatal
development. Similar developmental restriction of non-neural cells to NMJ’s was reported in
frogs (Connor, 1997). Whether this is due to changes in the composition of the basal lamina at
the neuromuscular junction (Patton et al, 1997; Sanes et al, 1990; Cho et al, 1998) or other
phenomena at endplates, such as endogenous electric fields ((Betz et al, 1980b; Kinnamon et al,
1985) remains to be established. Abnormalities of synaptic structure have been demonstrated in
mutant mice that lack specific components of the synaptic basal lamina (Noakes et al, 1995;
VanSaun et al, 2003). It will be interesting to examine the disposition of kranocytes in mutant
mice lacking specific basal lamina components (or endogenous electric fields; etc.).
A role for kranocytes in neuromuscular plasticity is suggested by their reaction in adults to nerve injury or paralysis. This reaction occurred at least two days earlier than the reactive transformation of terminal Schwann cells. This early reaction does not detract from the pivotal role played by reactive tSc in leading regenerating axons to denervated or paralysed motor endplates (Son et al, 1996; Love & Thompson, 1998, 1999; Love et al., 2003; Kang et al, 2003). Rather, the data underscore the importance of tSC’s, by demonstrating that their transformation to a reactive state is not an inevitable consequence of early reaction and spread of kranocytes. Perhaps kranocytes consolidate the cellular bridges they appear to form between motor endplates and these are subsequently utilized by reactive tSc’s. Terminal Schwann cell bridges then support subsequent axonal sprouting and regeneration (Fig. 8B). It is noteworthy that electrical stimulation of denervated rat muscle does not prevent Schwann cells from becoming reactive, but it strongly inhibits the formation of Schwann cell bridges between endplates (Love et al, 2003). It would be interesting to know whether stimulation also prevents the reaction of neuromuscular kranocytes to denervation.

We also found evidence of dispersal of the population of 2166-positive cells in atrophic muscles of the R6/2 transgenic mouse model of Huntington’s disease. However, since motor nerve terminals, terminal Schwann cells and neuromuscular transmission appear largely normal in R6/2 mice (Ribchester et al, 2004), denervation, paralysis or muscle atrophy would appear to be sufficient to provoke the kranocyte cell reaction; but the reaction of these cells is evidently not sufficient to trigger the subsequent Schwann cell reaction in this disease model. It would be interesting to know the disposition and fate of kranocytes in other models of neuromuscular disease, particularly models of amyotrophic lateral sclerosis, where neuromuscular synaptic degeneration is among the earliest signs of disease (Fischer et al., 2004; Schaefer et al., 2005; Pun et al., 2006; Suzuki et al., 2007).
Conclusions
The focal distribution of a specific sub-population of antibody 2166-positive cells to neuromuscular junctions together with their progressive restriction to these areas during postnatal development and their persistence at synapses following muscle denervation, paralysis or atrophy, suggest important roles for kranocytes in the development, neurotrophic maintenance and repair of neuromuscular junctions. Several important questions remain: the possible gliotrophic function of kranocytes; the mechanism of their restriction to motor endplates, the functions of this developmental restriction; the significance of their proliferation and spread in atrophic muscle; and, finally, the identity of the 2166-antigen that was crucial in identifying these cells. Since we now have very good markers for kranocytes, there is a much greater potential than in the past for obtaining more information about the functions of these intriguing cells. Harnessing the molecular secretions of these plastic cells could perhaps also be utilised in targeting potential neuroprotective factors to neuromuscular junctions, potentially mitigating or preventing early signs of neuromuscular disease.

Materials and Methods
Generation of 2166 antibody
The 2166 antibody was generated from a peptide (amino acid sequence CAIRNSRDVI) corresponding to the C-terminus of the Tspan-2 protein, coupled to keyhole limpet hemocyanin (KLH). This peptide was used to inoculate male New Zealand white rabbits of approximately 1.5 Kg in weight. After three injections of KLH-coupled peptide, serum was collected at appropriate time points and affinity purified. We made a preliminary attempt to identify the putative antigen by western blotting of a cytoskeletal protein fraction from cerebellum (data not shown). The antibody recognised a band of approximately 47 KDa. Analysis by mass spectrometry was consistent with an antigen of similar molecular weight to tubulin. Pre-immune serum was inactive and specific staining was blocked by preincubating the active serum with the peptide used to raise the antibody. However, we were unable to achieve immunoprecipitation of any specific protein with the 2166 antibody and it did not recognise purified tubulin on western blots.
A more stringent fractionation and/or proteomic analysis may therefore be necessary to identify the 2166 antigen.

Animals and surgery

All animals were maintained in a regulated, clean environment and treated in compliance with UK Home Office regulations. Adult female Sprague-Dawley rats were used in some experiments but most experiments were performed using the following mouse lines: C57Bl/6, S100-eGFP transgenic (accessed during a visit by one of us (FAC) to Prof P Caroni’s laboratory), R6/2 transgenic (Ribchester et al., 2004) and tenascin-C null mutant (TNC/TN-F18; Saga et al, 1992; donated by Prof C ffrench-Constant). Tenascin-C null mutant mice were genotyped as described by Garcion et al (2004). In some experiments, adult mice were anaesthetised intraperitoneally with 1mg/kg medetomidine (Domitor), 75mg/kg ketamine (Vetalar) and one or more of the intercostal nerves supplying the triangularis sterni (TS) muscle were crushed unilaterally. In other anaesthetised adult mice, intercostal muscles adjacent to the TS muscle were injected with 2.5 µl botulinum toxin type A (BoNT/A;10 ng/ml; Sigma) from a Hamilton syringe. Surgically-prepared animals were recovered for 1-6 days. All animals were sacrificed by cervical dislocation.

Immunocytochemistry and histochemistry

Whole-mount preparations were made by dissecting TS muscles in 0.1 M phosphate buffered saline (PBS, pH 7.4), fixed in 4% paraformaldehyde for 20 minutes, then incubated in either TRITC-or Alexafluor-647 conjugates of α-bungarotoxin (Molecular Probes; 5 µg/ml for 30 min). After washing in PBS and blocking/permeabilisation for one hour in 1% BSA, 0.4% Lysine, 0.5% Triton X-100 in PBS, primary antibodies (see below) were applied overnight at 4 °C. Muscles were washed and incubated with secondary antibodies for 3.5 hours at room temperature, washed and mounted in Vectashield (Vector Labs). Most of the observations reported in the Results were based on analysis of four whole-mount preparations per immunostain combination. For muscle section immunocytochemistry, dissected muscles were embedded in OCT (Tissue TEK) and frozen in isopentane precooled with liquid nitrogen. Cryostat sections were collected, fixed with 4% paraformaldehyde for 5 minutes, then blocked and permeabilised for 1 hour with 5% fish gelatine and 0.1% Triton X-100 in PBS. Primary antibodies (see below) were applied overnight at room temperature. Sections were washed and
incubated with secondary antibodies for 1 hour at room temperature, washed in PBS and mounted in Vectashield. Preparations were imaged using a BioRad Radiance 2000 confocal microscope on a Nikon Eclipse E600-FN platform. For staining with cholera toxin B (CTB), TS muscles from male Balb/c mice (6-8 weeks) were dissected, pinned on Sylgard-lined dishes and bathed in FITC-conjugated CTB (Sigma, 1:300) and a Texas-Red conjugate of α-bungarotoxin (Molecular Probes, Eugene), diluted 1:500 in oxygenated saline for 2 hours at 32 °C; followed by 30 min at 4°C, rinsed in buffer, fixed at room temperature for 15 minutes in 4% paraformaldehyde, rinsed in 0.1M glycine and mounted in Citifluor anti-fade reagent (Citifluor, Canterbury, UK). Images were obtained using a Zeiss Pascal confocal microscope. Voxx rendering software (Clendenon et al., 2002) was used for 3-D reconstruction.

Lac-Z expression in whole mounts of TS muscles from heterozygous tenascin-C null mutant mice was detected by X-gal staining solution, with a solution containing 2 mM MgCl₂, 5 mM K₃Fe(CN)₆, 5 mM 0.02% NP-40, and 1 mg/ml Xgal (5-bromo-4-chloro-3-indolyl-D-galactoside; Sigma). Image post-processing was carried out using Adobe Photoshop, and/or Image J software, in compliance with Journal of Cell Science image manipulation policy.

Sources and dilutions of species-specific antibodies

Antibodies were obtained from the following suppliers and used at indicated dilutions: rabbit 2166 (Peter Brophy, University of Edinburgh; 1:200); mouse anti-desmin (DAKO; 1:200); rat IgG2k F4/80 (Serotec; 1:50); mouse IgG1 anti-GFAP (Boehringer; 1:100); rat IgG anti-Laminin (a2 chain) (Alexis; 1:200); rabbit anti-M-Cadherin (Anton Wernig, University of Bonn; 1:50); rabbit anti-N-CAM (Elisabeth Bock, University of Copenhagen; 1:100); mouse IgG1 anti-Nestin (Susan Hockfield, Yale University; 1:200); mouse anti-neurofilament (165 kDa) (DHSB; 1:200); rabbit anti-S100 (DAKO; 1:200); mouse IgG1 anti-S100 (SA1259) (Affinity; 1:200); mouse anti-SV2 (SAPU; 1:200); mouse IgG1 anti-Thy 1.1 (Serotec; 1:100); rat IgG2b anti-CD34 (BD PharMingen; 1:50); mouse IgG1 anti-prolyl-4-hydroxylase (Acris Medicorp; 1:100); rat IgG anti-tenascin (Abcam; 1:200); rabbit HM-24 (Jonathan Cohen, Harvard Medical School; 1:200).

Electron microscopy

Isolated mouse TS muscle was fixed in 4% paraformaldehyde and 2.5% glutaraldehyde in ice-cold 0.1M phosphate buffer for 4 hours, then post-fixed in 1% osmium tetroxide for 45 minutes,
dehydrated in ethanol and propylene oxide and embedded in Durcupan resin. Three hundred ultrathin (75-100 nm) serial sections were cut from the endplate region, collected on formvar-coated grids (Agar Scientific, UK), stained with uranyl acetate and lead citrate then viewed in a Philips CM12 transmission electron microscope. Electron photomicrograph negatives were scanned at 1000 dpi. A few sections were lost during preparation (<20 out of 300), so these were replaced in the final image stack by a duplicate of a neighbouring serial section. Digitised micrographs were then transferred to a Unix workstation (Sun Microsystems) and a custom programme (Reconstruct) was used to generate 3D-voxel images. The programme “MAPaint” was used to reconstruct 3D volumes of the nerve terminal, skeletal muscle fibre, terminal Schwann cell and 2166 cells. Both Reconstruct and MAPaint were written and developed in the MRC Human Genetics Unit, Edinburgh [http://genex.hgu.mrc.ac.uk]. Different cell types were identified in each individual 2D image before manually delineating their borders. The 3D volumes were saved to independent files, calibrated and surface rendered using Imaris software (Bitplane; Zurich, Switzerland).

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ABBREVIATIONS (163 words)
2166 – Polyclonal antibody recognising kranocytes and other cell types AChR – Acetylcholine receptor BrdU – bromodeoxyuridine BoNT/A – botulinum toxin Type A C57Bl/6 – Standard laboratory mouse line CD34 – Stem cell marker CTB – Cholera Toxin B-subunit DAPI -4',6-diamidino-2-phenylindole DG2 – antiganglioside antibody DL – deep lumbrical muscles EDL – extensor digitorum longus muscle F4/80 – macrophage marker FDB – flexor digitorum brevis muscle GAP-43 – growth associated protein of molecular weight 43kDa GGFII – glial growth factor 2; neuregulin GFAP – glial fibrillary acidic protein GFP – Green Fluorescent Protein HM-24 – Neuregulin antisera ICC – Interstitial Cells of Cajal KLH -keyhole limpet hemocyanin Lac-Z – β-galactosidase NCAM – neural cell adhesion molecule NMJ -Neuromuscular Junctions ISC – Terminal Schwann Cells R6/2 – Huntington’s Disease model: transgenic mouse line rPH – rat prolyl-4-hydroxylase S100 – Ammonium-sulfate soluble Schwann cell antigen TRITC – tetramethylrhodamine isothiocyanate TRITC-α-BTX – TRITC conjugated with α-bungarotoxin TS – triangularis sterni muscle Tspan-2 – tetraspanin protein 2 WB – Western blot X-gal -β-galactosidase stain
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Immunostaining of mouse triangularis sterni muscles with 2166 antibody reveals capping cells at NMJ’s. A: low power confocal projection showing numerous 2166-positive interstitial cells and a prominent sub-population in the vicinity of NMJ’s. B: higher power confocal projection images of NMJ-capping cells (green), juxtaposed with motor endplates counterstained with TRITC-α-BTX (red). C: Triple-stained NMJ with axon stained with neurofilament antibodies (red), NMJ-capping cell (green) and motor endplate AChR stained with Alexafluor647-α-BTX (blue). D, E: other interstitial and endothelial cells with positive 2166 immunostaining were scattered at low density throughout the muscles. F: quantification of density of NMJ-capping cells at motor endplates. All endplates showed at least one and most were associated with only one NMJ-capping cell; no NMJ was associated with more than three of these cells. Scale bars: A, 100 µm; B, 20 µm (applies also to C and D).
Figure 2

NMJ-capping cells are not Schwann cells but express collagensynthesising enzyme and neuregulin. **A**: Confocal projection images of a triangularis sterni endplate in an S100-eGFP transgenic, showing endogenous fluorescence in terminal Schwann cells (green) overlying endplate ACh receptors (blue). The right panel of the pair shows superimposition of 2166 immunostaining (red). **B**: Immunostaining for the collagensynthesising enzyme prolyl-4hydroxylase (rPH) of an NMJ-capping cell in rat triangularis sterni muscle. **C**: Immunostaining of a NMJ-capping cell in mouse triangularis sterni muscle using HM-24 antibody, which recognizes GGFII/neuregulin. Scale bar: 50 µm.
Figure 3

NMJ-capping cells are CD34 antibody and CTB positive and reside outside the synaptic basal lamina. **A,B:** Confocal projections of an NMJ-capping cell in a mouse triangularis sterni muscle immunostained with 2166 (green) and CD34 (red) antibodies and with endplate AChR counterstained (blue). Lower panels in A show green and red channels separately; Panels 1 and 2 in B show orthogonal projections at the optical cuts indicated in the uppermost panel. **C:** Transverse section of a mouse TS muscle immunostained with 2166 antibody (green spots), laminin antibody (red) and endplate AChR (blue). The immunostained processes of the NMJ-capping cells clearly lie outside the basal lamina. **D:** Fluorescent conjugates of CTB also stain cells similar in form and location to NMJ-capping cells. Right panel shows the red channel indicating endplate AChR only. Scale bars: **A,** 50 µm (applies also to **C**); **D,** 50 µm.
Figure 4
NMJ-capping cells almost completely cover motor endplates.
Electron micrographs and 3D surface-rendered reconstructions of serially-sectioned electron micrographs showing the relationship between two putative NMJ-capping cells, motor nerve terminal, terminal Schwann cell and skeletal muscle fibre at a neuromuscular junction from mouse TS muscle. A, B. Sections 84 and 75 respectively from a 300-section series through a single NMJ. The soma and thin cytoplasmic processes of both NMJ-capping cells at this NMJ can be seen above the terminal Schwann cell and nerve terminal, lying outside the basal lamina. Each of the main cell types have been outlined on these electron micrographs (motor nerve terminal -green; muscle fibre -brown, terminal Schwann cell -yellow; 2166 cells -magenta and lilac). C: Surface rendering of reconstructed serial electron micrographs from the NMJ shown in A and B revealed two NMJ-capping cells (magenta and lilac) overlying a single terminal Schwann cell (yellow) and a motor nerve terminal (green) synapsing with a skeletal muscle fibre (brown). a: superior view of the nerve terminal and skeletal muscle fibre; b: nerve terminal only; c: nerve terminal and terminal Schwann cell; d: nerve terminal,
Schwann cell and two NMJ-capping cells; e: nerve terminal and two NMJ-capping cells; f: two NMJ-capping cells; g/h: the two NMJ-capping cells shown in isolation; i: orthogonal view of the reconstructed NMJ-capping cells. Animations of the volume-rendering of these two cells are presented in Supplementary Movies 6 & 7. The NMJ-capping cells cover approximately 60% of the nerve terminal at this NMJ. In contrast to the terminal Schwann cell, the veil-like processes of both NMJ-capping cells extend beyond the immediate boundaries of the endplate. D. Transmission electron micrograph through a neuromuscular junction in the rat diaphragm. The motor nerve terminal and terminal Schwann cell are covered by a slender process from a putative NMJ-capping cell (arrows; a high resolution image is reproduced in Supplementary Figure 3). Scale bar: 2.5 μm.
Figure 5

NMJ-capping cells become restricted to the endplate zone postnatally but spread following denervation in adults. A-C: 2166 immunostaining (green) is diffuse at birth but becomes progressively concentrated near NMJ’s over the following 1-2 weeks. D-F: Denervation in adults triggers nestin reactivity and sprouting in terminal Schwann cells within 3 days but no discernible sprouting or nestin immunoreactivity is seen 1 day after axotomy. In contrast, extensive spreading of 2166 immunostaining occurs within 1 day of axotomy (See also Supplementary Figure 4). At 3 days, some of the Schwann cell sprouts were associated with 2166-positive links between motor endplates (arrowhead; see also Supplementary Figure 5). G: Immunostaining following injection of a BrdU pulse shows BrdU positive staining (red) of nuclei (blue) in both junctional and non-junctional 2166-positive cells (green), suggesting that NMJ-capping cells both divide and spread from endplates following axotomy. Arrowheads indicate BrdU positive nuclei. Scale bar: A-C, 100 µm; D-G, 50 µm (applies also to D, E and F).
Figure 6

NMJ-capping cells also spread in paralysed and atrophic muscles. **A,B:** Profusion of 2166-immunoreactivity (green) is seen within one day of injection of a locally-paralysing dose of botulinum toxin type-A but terminal Schwann-cell reactivity and sprouting, assessed by nestin immunostaining (red), is not observed until at least 6 days of paralysis in TS muscles. **C,D:** Similar profusion and spread of NMJ-capping cells is observed in constitutively atrophic muscles of 15-week old R6/2 transgenic mice. Littermate controls show normal, endplate-localised staining and distribution of NMJ-capping cells. Scale bar: 50 µm
Figure 7

Reactive NMJ-capping cells are associated with tenascin-C expression but do not require tenascin-C in order to localize, react or spread from NMJ’s.

A-C: Tenascin-C immunoreactivity (red) spread from the endplate region from 36 days after axotomy (left panels) and these tenascin-C rich regions coincide with the distribution of NMJ-capping cells (green, right panels; AChR counterstained blue). D,E: Selective X-Gal staining of intramuscular nerve and motor endplates in heterozygous tenascin-C null mutant mice with lac-Z substitution, 5 days after axotomy. The area shown in E corresponds to the area bounded by the dotted outline in D. Unoperated muscles were X-Gal negative (not shown). F,G: NMJ-capping cells are also localized to NMJ in homozygous tenascin-C null mutant mice and these cells spread from endplates 3 days after axotomy, as in wild-type mice. Scale bars: F, 50 µm (applies also to A, B, C and E); G, 50 µm
Figure 8

NMJ-capping cells are integral, plastic cellular components of the NMJ.

A: Cartoon depicting the relationship between four main cell types proposed to constitute the mammalian NMJ: muscle fibre, motor nerve terminal, terminal (perisynaptic) Schwann Cell and NMJ-capping cells, named here as “kranocytes” (see Discussion).

B: Stages in kranocyte reactivity to denervation, paralysis or muscle atrophy that could indicate a permissive function for these cells in terminal Schwann cell and axonal sprouting. Kranocyte sprouting is the first reaction, followed by terminal Schwann cell sprouting, then axonal sprouting.
LEGENDS TO SUPPLEMENTARY FIGURES AND MOVIES

Supplementary Figure 1
NMJ-capping cells are Thy-1 negative but co-localise after denervation with activated terminal Schwann cells. Panels showing combined (left) and separated (middle and left) colour channels of preparations immunostained with 2166 antibody and either Thy 1.1 (A) or S100 (B). A. Co-immunostaining of mouse triangularis sterni muscle with 2166 antibody (green) and Thy-1.1 antibody (red) show no co-localisation. However, Thy-1 immunopositivity was observed in some of the other perijunctional fibroblasts and intramuscular capillaries. Acetylcholine receptors were counterstained with fluorescent bungarotoxin (pseudocoloured blue). B. Co-immunostaining of rat 4th deep lumbrical muscle with 2166 antibody (green) and S100 antibody (red) 6 days after section of the tibial nerve. Sprouts between muscle fibres made by NMJ-capping cells and terminal Schwann cells are clearly coincident and associated, but the antibodies are evidently not colocalised in the same cells.

Supplementary Figure 2

Supplementary to Fig 3B/C, 2166 immunostaining of transverse muscle sections shows thin NMJ-capping cell processes lying outside the synaptic basal lamina, as indicated by co-staining for laminin (red) and Alexa647-α-BTX (blue). Calibration 20 µm

Supplementary Figure 3

High resolution image of the NMJ shown in Figure 4D. Note the thin basal lamina around most of the putative NMJ-capping cell and its processes above the NMJ, and the presence of rough ER in the cytoplasm. Key: MF – muscle fibre; NT-nerve terminal; tSC – terminal Schwann Cell; K-Kránocyte; BL – basal lamina. Calibration 2 µm.
Supplementary Figure 4
Co-staining of TS muscles in unoperated muscles (A,B) and 1 day after axotomy (C,D) with 2166 antibody (green), DAPI (blue; nuclei) and TRITC-α-BTX (AChR). These images show evidence of a rapid increase in the number of 2166-positive cells per NMJ following denervation. Calibration 30 µm (A,B); 20 µm (C,D).

Supplementary Figure 5
Co-staining of a 3-day denervated TS muscle with 2166 antibody (green), nestin (red) and Alexa647-α-BTX (blue). Note the association of a cell processes from an NMJ capping cell and a nestin-positive tSC sprout. This figure shows separation of the RGB channels for the same image as Figure 5E. Calibration 40 µm.

Supplementary Movie 1
Z-series of triple-stained NMJ shown in flat projection in Fig 2B rPH (red) /2166 (green)/AChR (blue)

Supplementary Movie 2
Animated rotating projection of z-series for triple-stained NMJ shown in Supplementary Movie 1 and in flat projection in Fig 2B. rPH (red) /2166 (green)/AChR (blue)

Supplementary Movie 3
Z-series of endplate AChR stained with TRITC-α-BTX, shown in projection and with volume rendering in Fig 3D

Supplementary Movie 4
Z-series of NMJ-capping cells stained with CTB, shown in projection and with volume rendering in Fig 3D
**Supplementary Movie 5**
Animated rotating projection of Z-series of NMJ-capping cells stained with CTB, shown in Supplementary Movie 4 and in projection with volume rendering in Fig 3D.

**Supplementary Movie 6**
Volume rendered animation of one of the reconstructed NMJ-capping cells shown in Fig 4A-C.

**Supplementary Movie 7**
Volume rendered animation of the other reconstructed NMJ-capping cell shown in Fig 4A-C.

| Antibody/Marker | Cell Type or Antigen       | 2166 Cells |
|-----------------|---------------------------|------------|
| S100            | Schwann Cells             | -          |
| GFAP            | Reactive Schwann Cells    | -          |
| Nestin          | Reactive Schwann Cells    | -          |
| M-Cadherin      | Satellite Cells           | -          |
| NCAM            | Satellite Cells           | -          |
| Desmin          | Satellite Cells           | -          |
| Thy-1           | Fibroblasts/Neurons       | -          |
| F4/80           | Macrophages               | -          |
| CD34            | Stem Cells                | +          |
| rPH             | Fibroblasts               | +          |
| HM-24           | GGFII-related molecules   | +          |
| CTB             | GM1 Ganglioside           | +          |

**Table 1**
Panel of cellular and molecular markers used to screen for NMJ-capping cell epitopes.
Positive staining was obtained with CD34, rPH, HM-24 and CTB. Other markers of Schwann cells, macrophages, fibroblasts and other satellite cells did not stain NMJ-capping cells.