Graded elevation of c-Jun in Schwann cells in vivo: gene dosage determines effects on development, re-myelination, tumorigenesis and hypomyelination.

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Abstract

Schwann cell c-Jun is implicated in adaptive and maladaptive functions in peripheral nerves. In injured nerves, this transcription factor promotes the repair Schwann cell phenotype and regeneration, and it promotes Schwann cell mediated neurotrophic support in models of peripheral neuropathies. However, c-Jun is associated with tumour formation in some systems, it potentially suppresses myelin genes, and has been implicated in demyelinating neuropathies. To clarify these issues, and determine how c-Jun levels determine its function, we have generated, c-Jun OE/+ and c-Jun OE/OE mice, with graded expression of c-Jun in Schwann cells, and examined these lines during development, in adulthood and after injury using RNA sequencing analysis, quantitative electron microscopic morphometry, Western blotting and functional tests. Schwann cells are remarkably tolerant of elevated c-Jun, since the nerves of c-Jun OE/+ mice, where c-Jun in elevated about six fold, are normal with the exception of modestly reduced myelin thickness. The stronger elevation of c-Jun in c-Jun OE/OE mice is, however, sufficient to induce significant hypomyelination pathology, implicating c-Jun as a potential player in demyelinating neuropathies. The tumour suppressor P19ARF is strongly activated in the nerves of these mice, and even in aged c-Jun OE/OE mice, there is no evidence of tumours, in agreement with the fact that tumours do not form in injured nerves, although they contain proliferating Schwann cells with strikingly elevated c-Jun. Furthermore, in crushed nerves of c-Jun OE/+ mice, where c-Jun levels are over-expressed sufficiently to accelerate axonal regeneration, myelination and function are restored after injury.
Significance statement

In injured and diseased nerves, the transcription factor c-Jun in Schwann cells is elevated, and variously implicated in controlling beneficial or adverse functions, including the trophic Schwann cell support for neurons, promotion of regeneration, tumorigenesis and suppression of myelination. To analyse the functions of c-Jun, we have used transgenic mice with graded elevation of Schwann cell c-Jun. We show that high c-Jun elevation is a potential pathogenic mechanism, since it inhibits myelination. On the other hand, we do not find a link between c-Jun elevation and tumorigenesis. Modest c-Jun elevation, which is beneficial for regeneration, is well tolerated during Schwann cell development and in the adult, and is compatible with restoration of myelination and nerve function after injury.
Schwann cell c-Jun has been implicated both in adaptive and maladaptive functions in peripheral nerves. On the one hand, in injured nerves, this transcription factor is a global amplifier of the repair Schwann cell phenotype and promotes regeneration, and in models of peripheral neuropathies Schwann cell c-Jun supports axonal survival, trophic factor expression and sensory-motor function (Arthur-Farraj et al., 2012, 2017; Hantke et al., 2014; Klein et al., 2014; Jessen and Mirsky 2016). Reduced c-Jun levels in Schwann cells are also implicated in failure of regeneration due to ageing and long-term denervation (Painter et al., 2014; Jessen and Mirsky 2016). The c-Jun pathway is therefore of interest for the development of a pharmacology for nerve repair. On the other hand, c-Jun is associated with tumour formation in some systems, and c-Jun potentially suppresses myelin genes (Eferl and Wagner 2003; Parkinson et al. 2008). Based on this, it has been characterized as a negative regulator of myelination, and implicated in demyelinating neuropathies (Jessen and Mirsky 2008).

In the present work we have generated and analysed mouse lines with graded expression of c-Jun in Schwann cells in order to clarify these issues, and determine how c-Jun levels determine its function.

c-Jun is present at low levels in Schwann cells of uninjured nerves, but is rapidly elevated 80-100-fold after nerve cut (De Felice and Hunt 1994; Shy et al., 1996; Parkinson et al., 2008; J. Gomez-Sanchez, K.R. Jessen, R. Mirsky unpublished). c-Jun elevation is also seen in human neuropathies (Hutton et al., 2011). Although c-Jun is implicated in the promotion of a number of tumours, in other situations c-Jun may have a role in the prevention of tumourigenesis by mechanisms that include activation of tumour suppressors such as P14ARF/p19ARF and Dmp1 (Eferl and Wagner 2003; Ameyar-Zazoua et al., 2005; Shaulian et al., 2010). P19ARF is elevated in Schwann cells after nerve transection, and the striking activation of Schwann cell c-Jun after injury is not associated with tumour formation (Gomez-Sanchez et al., 2013; Jessen et al., 2015b). Rather, the role of c-Jun is to take part in controlling the conversion of myelin and Remak cells to Schwann cell specialized to carry out injury-specific tasks and promote repair (Jessen et al., 2015a; Jessen and Mirsky 2016; Arthur-Farraj et al., 2017). This includes preventing the death of injured neurons and promoting axon growth by expression of trophic factors, guiding axons back to their targets by forming regeneration tracks (Bungner bands), and breakdown of myelin directly by autophagy and indirectly by cytokine expression to recruit macrophages. Inactivation of Schwann cell c-Jun results in defective repair Schwann cells and impaired regeneration (Arthur-Farraj et al., 2012; Jessen and Mirsky 2016).
The injury-induced extinction of myelin genes is also delayed without c-Jun, indicating that c-
Jun has a dual function, promoting the expression of the repair phenotype and the
suppression of the myelin phenotype. c-Jun suppression of myelin genes has only been
studied directly in culture, where c-Jun suppresses the Krox20- or cAMP-induced activation
of myelin genes, and enforced c-Jun inhibits myelination in co-cultures (Parkinson et al.,
2004; 2008). Negative transcriptional regulation of myelination has also been shown for
Notch1 and Sox2 in vivo and suggested for other factors including Pax-3, Id2 and Sox-2
based on cell culture experiments (Jessen and Mirsky 2008; Roberts et al., 2017).

The present results show that the function of c-Jun in Schwann cells depends on gene
dosage, and that Schwann cells are surprisingly tolerant of the moderately (~ 6-fold)
elevated c-Jun, seen in c-Jun OE/+ mice. In these mice, where over-expression of c-Jun is
sufficient to accelerate axonal regeneration (Wagstaff et al., 2017), myelination and function
are restored after nerve injury. Further, even high expression of c-Jun is not associated with
tumour formation in Schwann cells, although this is sufficient to cause hypo-myelination
neuropathy.

**Materials and methods**

Transgenic mice

Animal experiments conformed to UK Home Office guidelines under the supervision of UCL
Biological Services. To generate mice that overexpress c-Jun selectively in Schwann cells,
female R26c-Junstopf mice, generated in the laboratory of Klaus Rajewsky, which carry a
lox-P flanked STOP cassette in front of a CAG promoter driven c-Jun cDNA in the ROSA26
locus, were crossed with male P0Cre+/+ mice (Feltri et al., 1999). This generated P0-Cre+/+
;R26c-Junstopf/+ mice, which we refer to as c-Jun OE/+ mice. These male mice were back-
crossed with female R26c-Junstopf mice to generate P0-Cre+/+;R26c-Junstopf/+ mice,
referred to as c-Jun OE/OE mice. P0-Cre+/+ littermates were used as controls. Mice of either
sex were used in the experiments. The mice are on the C57BL/6 background.

Genotyping

DNA for genotyping was extracted from ear or tail samples using the HotSHot method
(Truett et al., 2000). Primers for genotyping the R26c-Junstopf transgene were 5'-
TGGCACAGCTTAAGCAGAAA-3' and 5'-GCAATATGGTGGAAAATAAC-3' (270bp ). The
primers for the Rosa26 wildtype locus were 5'GGAGTGTTGCAATACCTTTCTGGGAGTTC-
3' and 5'TGTCCCTCAATTTTACACCTGTTCAATTC-3' (217bp band). The primers for the P0-Cre transgene were 5'-GCTGGCCCAAATGTTGCTGG-3' and 5'CCACCACCTCTCCATTGCAC-3' (480bp band).

Nerve injury

The sciatic nerve was exposed and crushed (3x15 sec at three rotation angles) at the sciatic notch using angled forceps. The wound was closed using veterinary autoclips. The nerve distal to the crush was excised for analysis at various time points. Contralateral uninjured sciatic nerves were used as controls for western blotting, immunofluorescence or electron microscopy.

Schwann cell culture

Schwann cell cultures were prepared from sciatic nerves of postnatal day 8-10 mouse pups essentially as in Morgan et al., (1991) and Arthur-Farraj et al., (2011). After enzymatic dissociation and centrifugation, the cell pellet was resuspended in defined medium (DM) (Meier et al., 1999), containing $10^6$ M insulin and 5% HS, and plated in drops on coverslips coated with Poly-L-lysine and laminin. Cells were incubated at 37°C/5%CO2 and allowed to adhere for 24 hr. After 24 hr, the medium was changed to DM/0.5% HS (controls), or DM with 10ng/ml NRG1 alone, or DM with NRG1 10ng/ml and dbcAMP 1mM for 48 hr before fixation and immunolabelling.

Antibodies

The following antibodies were used for Immunofluorescence: c-Jun (Cell Signalling, rabbit 1:800, 9165), Ki67 (Abcam, rabbit 1:100, ab15580), Krox20 (Covance, rabbit 1:100, PRB-236P), SOX-10 (R&D Systems, goat 1:100, AF2864), donkey anti-goat IgG (H+L) Alexa Fluor 488 conjugate (Molecular Probes, 1:1000, A11057), Cy3 donkey anti-rabbit IgG (H+L) (Jackson Immunoresearch, 1:500, 711-165-152), biotinylated anti-rabbit IgG (Amersham, 1:600, RPN1004), Cy3 Streptavidin (Jackson Immunoresearch, 1:500, 016-160-084).

The following antibodies were used for Western blot: GAPDH (Sigma, rabbit 1:5000, G9545), Calnexin (Enzo Life Sciences, rabbit 1:1000, ADI-SPA-860-D), c-Jun (Cell Signalling, rabbit 1:1000, 9165), Krox20 (Millipore, rabbit 1:500, ABE1374), Mpz (AvesLab, chick 1:2000, PZO), cyclin D1 (Santa Cruz, rabbit 1:200, sc-450), p19 Arf (5-c3-1) (Santa
Cruz, rat 1:100, sc-32748), anti-mouse IgG HRP-linked (Promega, 1:2000, W4028), anti-rabbit IgG, HRP-linked (Cell Signaling, 1:2000, 7074), anti-rat IgG, HRP-linked (Cell Signaling, 1:2000, 7077) anti-chicken IgY, HRP-linked (Promega, 1:2000, G1351).

Immunohistochemistry
Schwann cells were fixed in 4% paraformaldehyde/PBS for 15 min and then immunolabelled. Transverse sciatic nerve cryosections (10μm) were post fixed with 4% paraformaldehyde/PBS for 15 min, blocked in 0.2%Triton-X-100, 10%HS in PBS and subsequently incubated with primary antibodies in blocking solution overnight at 4ºC, followed by 2 hr in secondary antibodies and DAPI to identify cell nuclei (Thermofisher, 1:50000). For Ki67 staining biotin antibodies followed by Cy3 Streptavidin were used.

Western Blotting
For blotting, homogenates were obtained from injured and uninjured nerves as well as cultured nerve segments essentially as described previously (Gomez-Sanchez et al., 2015). Experiments were repeated at least three times with fresh samples and representative pictures are shown. Densitometric quantification was by Image Lab 4.1 (Bio-Rad Laboratories). Measurements were normalized to loading control GAPDH and/or calnexin.

RNA sequencing analysis
Library preparation: Total RNA was isolated using the RNeasy Lipid Tissue Mini Kit (Qiagen) with a DNase I step performed to eliminate traces of genomic DNA. The purified mRNA was fragmented, and primed with random hexamers. Strand-specific first strand cDNA was generated using Reverse Transcriptase in the presence of Actinomycin D. The second cDNA strand was synthesised using dUTP in place of dTTP, to mark the second strand. The resultant cDNA was then “A-tailed” at the 3’ end to prevent self-ligation and adapter dimerisation. Truncated adaptors, containing a T overhang were ligated to the A-Tailed cDNA. Successfully ligated cDNA molecules were then enriched with limited cycle PCR (10-14 cycles. The high fidelity polymerase used in the PCR is unable to extend through uracil. This means only the first strand is amplified for sequencing, thus making the library strand specific.
Sequencing: Libraries to be multiplexed in the same run were pooled in equimolar quantities. Samples were sequenced on the NextSeq 500 instrument (Illumina, San Diego, US) resulting in ~ 16M reads per sample.

Data Analysis: Run data were demultiplexed and converted to fastq files using Illumina’s bcl2fastq Conversion Software v2.18 on BaseSpace. Fastq files were aligned to a reference genome using STAR on the BaseSpace RNA-Seq alignment app v1.1.0. Reads per transcript were counted using HTSeq and differential expression was estimated using the BioConductor package DESeq2 (BaseSpace app v1.0.0).

The RNA Sequencing analysis was carried out by UCL Genomics, UCL Great Ormond Street Institute of Child Health.

Electron microscopy

Nerves were processed as previously described (Gomez-Sanchez et al., 2015). Transverse ultrathin sections from neonatal, P7 and P21 sciatic nerve or adult (P60) or aged (P300) sciatic nerve, or from injured distal stumps of adult sciatic nerves were taken 5 mm from the sciatic notch or cut site and mounted on film.

Photographs were taken using a Jeol 1010 electron microscope with Gatan camera and software. Images were analysed using NIH ImageJ. Photographs at were taken at 3000x to measure the number of myelinated axons, non-myelinated axons bigger than 1.5 μm and Schwann cell nuclei. The nerve area was measured from photographs taken at 200x magnification. Higher magnifications were used to show the Remak bundles or any abnormal morphology found in the nerve.

The percentage of extracellular matrix was analysed using ImageJ software by converting electron microscopy images at 5000x and 10000x into 8-bit images and tracing the presence of extracellular matrix.

Behavioural Tests

Experiments conformed to UK Home Office guidelines. Nine or more mice/ genotype were tested. Six week old mice were tested before surgery to ensure that there were no differences in normal responses between the genetic backgrounds. Sciatic function index (Inserra et al., 1998), toe spread reflex (Ma et al., 2011) and the toe pinch test, modified from Collier et al., (1961), were carried out as in Arthur-Farraj et al., (2012).
Statistical Analysis

Results are expressed as mean ± SEM. Statistical significance was estimated by one-way ANOVA with Tukey correction, two-way ANOVA with Bonferroni’s multiple comparison, Mann-Whitney U-test, or Student’s T-test. A P value < 0.05 was considered as statistically significant. Statistical analysis was performed using GraphPad software (version 6.0).

Results

Adult uninjured nerves of c-Jun OE/+ and c-Jun OE/OE mice have high levels of c-Jun protein in Schwann cell nuclei

Diagrammatic representation of how the c-Jun overexpressing mice were bred and produced is shown in Fig. 1A. The R26c-Junstopf mouse has a c-Jun cDNA insert in the Rosa26 WT locus with two flanking loxP sites either side of a STOP codon. These mice were bred with P0Cre+/− mice (Feltri et al., 1999). In the presence of Cre recombinase, the STOP codon is removed and c-Jun is overexpressed specifically in Schwann cells (Feltri et al., 1999). P0Cre+/−; R26c-Junstopf/+ control mice will be referred to as WT, while P0Cre+/−; R26c-Junstopf/+ will be referred to as c-Jun OE/+ mice, and P0Cre+/−; R26c-Junstopf/+ will be referred to as c-Jun OE/OE. Genotyping of WT, c-Jun OE/+ and c-Jun OE/OE mice is shown in Fig. 1B.

We examined c-Jun protein expression in adult (postnatal day (P) 60) uninjured sciatic nerves of c-Jun OE/+ and c-Jun OE/OE mice and compared this with that seen in WT mice. Double immunolabelling with c-Jun antibodies and Sox10 antibodies to specifically identify Schwann cell nuclei showed that Schwann cells in c-Jun OE/+ nerves expressed clearly elevated nuclear c-Jun levels compared to that seen in WT nerves, which showed barely detectable c-Jun using this staining protocol (see Materials and Methods). In c-Jun OE/OE nerves, nuclear c-Jun levels were further increased (Fig. 1C). No increase in c-Jun was seen in Sox10 negative nuclei, labelled with DAPI (not shown), indicating that c-Jun over expression in these mouse lines was Schwann cell specific in agreement with previous observations (Feltri et al., 1999).

Western blotting showed that c-Jun protein levels in uninjured adult sciatic nerves were elevated about six fold in c-Jun OE/+ mice and about 28 fold in c-Jun OE/OE mice, compared to WT (Fig. 1D). In c-Jun OE/+ mice, c-Jun mRNA levels were 4.5 fold higher than in WT nerves.
These data indicate that the axonal signals that normally suppress c-Jun during myelination in vivo fail to suppress c-Jun expression from the c-Jun OE transgene, as expected (Parkinson et al., 2008; Jessen and Mirsky 2008). We verified this by exposing purified Schwann cell cultures to signals that mimic axonal myelin signals in mice, namely the combined activation of cAMP and neuregulin pathways (Arthur-Farraj et al., 2011). In these experiments, a combination of 1 mM dbcAMP and 10nM neuregulin failed to suppress nuclear c-Jun expression in c-Jun OE/+ cells although down-regulation of c-Jun protein was seen in WT cells (Fig. 1E).

The elevation of c-Jun specifically in Schwann cell nuclei in c-Jun OE/+ and c-Jun OE/OE mice allowed us to study in vivo the effects of a graded increase in c-Jun expression on Schwann cells in uninjured and injured nerves.

**Transcriptional profiling of uninjured nerves in WT, c-Jun OE/+ and c-Jun OE/OE mice**

To document changes in gene expression caused by c-Jun elevation in c-Jun OE/+ and OE/OE mice, we carried out RNA sequencing analysis on uninjured adult (P60) sciatic nerves. Heat map and Principal Component Analysis (PCA) confirmed that c-Jun overexpression was the dominant source of differential gene expression (Fig. 2A, B). In OE/+ nerves, which express about six fold WT levels of c-Jun protein, 67 genes were ≥2 fold up-regulated and 25 genes were ≥2 fold down-regulated compared to WT nerves. Among 13 genes we considered of particular interest, one gene was regulated ≥2 fold. This was *Shh* which was up-regulated (Fig. 2C). *c-Jun* was expressed at 153% of WT levels, and GDNF at 182% of WT levels, while the myelin protein genes *Mbp* and *Mpz* were expressed at about 65% and 75% of WT levels, respectively. Notably, the mRNA level for Krox20 (*Egr2*), a key myelin regulator, was essentially unchanged. The 15 most up- and down-regulated genes in c-Jun OE/+ are shown in Fig. 3A.

In OE/OE nerves, which express about 28 fold WT levels of c-Jun protein, 909 genes were ≥2 fold up-regulated and 1055 genes were down-regulated by ≥2 fold compared to WT nerves. Most of the 13 genes of particular interest changed expression by ≥2 fold in these mice (Fig. 2C). This included *c-Jun*, which was elevated four-to-five fold, *Gdnf*, which was elevated by about 56 fold, and *Shh* and *Olig1*, which were elevated 20 fold and 48 fold respectively. The myelin protein genes *Mbp* and *Mpz*, were reduced to 13-14 % of WT levels. The 15 most up- and down-regulated genes in c-Jun OE/OE nerves are shown in Fig. 3B.

A comparison of gene expression between OE/+ and OE/OE mice with respect to the 13 genes of interest and the most regulated genes is shown in Figs. 2C and 3C respectively.
The fact that in both c-Jun OE/+ and OE/OE mice, c-Jun protein was more strongly elevated, in terms of fold change from WT, than c-Jun mRNA, suggests that posttranscriptional controls are important in controlling c-Jun levels.

**Expression of myelin-related proteins in uninjured nerves of OE/+ and OE/OE mice**

We examined two key myelin related proteins, the pro-myelin transcription factor Krox20 (Egr2) and the myelin adhesion protein Pzero (Mpz) in uninjured sciatic nerves of c-Jun OE/- and OE/OE mice. In line with the mRNA data, Krox20 levels were essentially unaffected in c-Jun OE/+ mice, both in double label immunohistochemical experiments, which show Krox20 in Schwann cell nuclei, and in Western blotting experiments (Fig. 3D,E). Mpz levels in these mice were about 15% lower than those found in WT mice (Fig. 3F). In contrast, the c-Jun OE/OE mice expressed significantly less Krox20 protein in Schwann cell nuclei and Western blots (Fig. 3D,E), and much reduced levels of Mpz (Fig. 3F).

This indicates that in adult-Jun OE/+ nerves, the levels of key myelin-related proteins and their mRNA remain relatively mildly affected, in spite of about six fold elevation of Schwann cell c-Jun. This tolerance does, however, break down when c-Jun levels are elevated about 28 fold as seen in c-Jun OE/OE mice, in line with the capacity of c-Jun to negatively regulate myelin genes indicated in experiments in vitro and in mice with conditional inactivation of Schwann cell c-Jun (Parkinson et al., 2004; 2008; Arthur-Farraj et al., 2012).

**Structure of adult nerves is nearly normal in c-Jun OE/+ mice**

Although the levels of myelin proteins were normal in c-Jun OE/+ mice, it remained possible that the substantial c-Jun elevation affected myelination and nerve architecture. This was tested by a morphometric comparison of WT and c-Jun OE/+ nerves. The general appearance of WT and c-Jun OE/+ nerves of 60 day old mice was similar (Fig. 4A). The size of the cross-sectional profiles of the sciatic nerve and the number of Schwann cell nuclei were not significantly different between the two genotypes, and Ki67 labelling of Sox10 positive cells failed to show significant increase in Schwann cell proliferation (Fig 4B-D). WT and c-Jun OE/+ nerves had similar numbers of >1.5 μm axons per nerve profile, and the percentage of segregated (1:1), myelin-competent (>1.5μm diameter) axons that were myelinated, and the total number of myelinated axons were comparable (Fig 4E-G). Both WT and c-Jun OE/+ nerves contained similar very low numbers of myelin-competent (>1.5 μm diameter) axons in a 1:1 relationship that remained unmyelinated (Fig. 4H).

Measurements of g-ratios showed that myelin sheaths were slightly thinner in c-Jun OE/+
mice compared to WT (Fig. 4I). Remak bundles in c-Jun OE/+ mice appeared normal, and
the percentage of >1.5μm axons that were found within Remak bundles was similar and very
low in both genotypes (Fig. 4J).

We found that this similarity between WT and c-Jun over-expressing c-Jun OE/+ 60 day old
mice remained even in old (300 day) mice (Fig. 4K-S). As in young mice, observations of
general appearance and quantitative analysis failed to reveal significant differences between
the two genotypes, except for the difference in G-ratios, a difference that was also seen in
young mice (Fig. 4R). The only age-induced change related to Schwann cell numbers, which
were somewhat elevated in old mice, the difference between the genotypes reaching
statistical significance (Fig. 4M).

These observations show that although c-Jun OE/+ mice show about six fold elevation of c-
Jun protein that is localized to Schwann cell nuclei, they achieve essentially normal
Schwann cell and nerve architecture, with the exception of modestly reduced myelin
thickness.

Adult nerves of c-Jun OE/OE mice are hypomyelinated, show onion bulbs and hyperplasia
but do not form tumours

In contrast to c-Jun OE/+ mice, the higher (about 28 fold) c-Jun expression c-Jun OE/OE
mice resulted in obvious lack of myelin in 60 day old mice (Fig. 5A). Although the total
number >1.5μm axons was similar to WT (Fig. 5B), the percentage of segregated (1:1),
myelin-competent (>1.5 μm diameter) axons that were myelinated was reduced by about
40% (Fig. 5C), and there was a corresponding increase in the number of myelin-competent
axons that that had reached a 1:1 relationship but remained unmyelinated (Fig. 5E). The
myelin sheaths in c-Jun OE/OE mice were also thin compared to WT (Fig. 5F). While all of
this indicates impediment to myelination, a sorting defect was indicated by the fact that the
percentage of >1.5 μm axons that remained within Remak bundles was strikingly increased
to about 28% compared to <1% in WT (Fig. 5G). As a result of impaired myelination and
sorting, the number of myelinated axons in c-Jun OE/OE nerves was substantially lower than
in WT nerves (Fig. 5D). As seen in mouse models of CMT1A neuropathy and a number of
other mouse mutants with elevated, non-tumorigenic Schwann cell proliferation, the
organization of Remak bundles was somewhat altered (Robertson et al., 2002; Chen et al.,
2003; Ling et al., 2005; Verhamme et al., 2011). The cells sometimes showed increased
membranous structures and processes that were not in contact with axons, and they
contained fewer axons per transverse section of a bundle, suggesting the presence of a
larger number of Remak cells each taking care of fewer axons (Fig. 5A). Increased mast cell
Numbers are seen in several neuropathic conditions and mutant models and after mechanical nerve injury (Olson 1971; Ling et al. 2005; Ishii et al. 2016). We therefore counted mast cell numbers and found substantial elevation in c-Jun OE/OE nerves (Fig. 5H).

An increase in Schwann cell number, a feature of many neuropathies including CMT1A (Robertson et al., 2002; Lupski and Chance 2005), was also seen in c-Jun OE/OE nerves, the total number of Schwann cell nuclei per nerve profile being about six fold that in WT (Fig. 5I). Western blots of Cyclin D1 indicated ongoing proliferation among the cells of c-Jun OE/OE nerves (Fig. 5J). Proliferation of Schwann cells was indicated in double immunolabelling with Ki67 and Sox10 antibodies to detect dividing Schwann cells, since double labelled Schwann cells, although few, were about three times more common in c-Jun OE/OE nerves than in WT nerves (Fig. 5K). Observations in the electron microscope provided no evidence for the presence of a significant number of Schwann cells without contact with axons. The increased number of Schwann cell nuclei in nerve sections is likely due to non-myelinating cells in a 1:1 ratio with axons being shorter than myelin cells, cells with thin myelin sheaths being shorter than those with normal sheath thickness, and increased number of Remak cells.

The sciatic nerves of 60 day old c-Jun OE/OE mice were enlarged, showing total cross-sectional profiles that were about twice that in c-Jun OE/+ or WT nerves (Fig. 5L). Collagen containing extracellular space was also markedly increased in c-Jun OE/OE nerves, occupying 133,349 μm² (+/-19,891; n=3) (54% of nerve area) in 60 day c-Jun OE/OE nerves, but only 16,069 μm² (+/-2,834; n=5) (13% of nerve area) in WT nerve profiles in WT nerves (Fig. 4M). This amounts to an increase in extracellular space of 116,280 μm². Since the nerves of c-Jun OE/OE nerves are 121,486 μm² larger than WT nerves, more that 95% of the enlargement seen in c-Jun OE/OE nerves is due to increased collagen-containing extracellular space, with a likely contribution from increased number of Remak cells and cells other than Schwann cells. Increase in endoneurial connective tissue is seen in a number of neuropathies including CMT1A, and in the trembler and twitcher mouse mutants (Palumbo et al., 2002; Fledrich et al., 2012; Low 1977; Ling et al. 2005; Kagitani-Shimono 2008).

We examined the mutant nerves extensively for the presence tumours or cellular arrangements reminiscent of tumour formation, but failed to find any evidence in this direction. In line with this, the tumour suppressor p19ARF was strongly elevated in uninjured nerves of c-Jun OE/OE mice (Fig. 5N).

Examination of old (P300) mice showed that only three of the parameters studied above changed obviously with age. (Fig. 6A-J). This was the appearance of significant numbers of
onion bulbs (Fig. 6F,G), reduction in Schwann cell proliferation, which was no longer significantly elevated (Fig. 6J), and a reduction in the percentage of >1.5 μm axons that remained within Remak bundles, from about 28% at P60 (Fig. 5G) to less than 2% (Fig. 6E).

Thus, in nerves of c-Jun OE/OE mice, a large number of axons appear to gradually segregate from Remak bundles between P60 and P300. The proportion of these >1.5 μm axons that myelinate is similar to that of the >1.5 μm segregated axons in P60 nerves, or about 60% (Fig. 6B). No tumours were found in older mice (n=18).

**Developmental myelination is delayed in c-Jun OE/+ mice, but inhibited in c-Jun OE/OE mice**

Although adult nerves of c-Jun OE/- mice are essentially normal, we tested whether the c-Jun elevation in these nerves caused a delay in myelination during development. We also determined whether the lack of myelin in the adult c-Jun OE/OE nerves was due to delayed myelination in the adult or inhibition of myelination during development.

In developing nerves of c-Jun OE/+ mice, there was a trend towards c-Jun elevation at P1, but this was not significant, while at P7 Jun was elevated about six fold compared to WT nerves at the same age. This failed to suppress levels of the myelin proteins Mpz and Krox20 in Western blots (Fig. 7A,B). But nuclear Krox20 was reduced, judged by double labelling of nerve sections with Krox20 and Sox10 antibodies to identify Schwann cells (Fig. 7C).

In developing nerves of c-Jun OE/OE mice, c-Jun levels at P7 were about eight fold that found in WT mice at that age, and Mpz was suppressed in Western blots (Fig. 7A). Although Krox20 levels were not significantly reduced in Westerns, the number of Schwann cells that showed nuclear Krox20 was less 50% of that in WT nerves, as seen in double immunolabeling of nerve sections Fig. 7B,C).

Electron microscopy at P1, P7 and P21 showed that in c-Jun OE/+ mice, myelination was transiently delayed at P7, while in c-Jun OE/OE mice, myelination was severely inhibited (Fig. 7D).

In c-Jun OE/+ mice, nerve area and the percentage of >1.5 μm diameter axons that were found within Schwann cell families or Remak bundles, both of which were normal in the adult, were also normal during development at all three time points (Fig. 7E,K). However, a number of other parameters were abnormal at P7, although they were normal at P1 and P21, revealing a transient delay in myelination. This includes the number of Schwann cell nuclei, which was elevated (Fig. 7F), the percentage of segregated (1:1), myelin-competent (>1.5μm diameter) axons that were myelinated, which was reduced, the total number of myelinated axons, which was reduced (Fig. 7H), and the number of segregated (1:1) myelin-competent (>1.5 μm diameter) axons that were not myelinated,
which was elevated (Fig. 7I). In adult nerves of these mice, the myelin sheaths are slightly thinner than in WT, and this difference was already present at P7 and P21 (Fig. 7J).

In the developing c-Jun OE/OE nerves, nerve area was not significantly different from that seen in WT or c-Jun OE/+ mice (Fig. 7E). The large nerve area in P60 nerves of these mice therefore emerges in adulthood. At P7 and P21 these nerves contained about twice the number of Schwann cell nuclei seen in WT nerves, a smaller difference than that seen in the adult (Fig. 7F). This suggests ongoing, low-level Schwann cell proliferation in adult mutant nerves, supported by Ki67 labelling of Sox10 positive Schwann cells, although the differences between WT and c-Jun OE/OE nerves in Cyclin D1 levels and did not reach significance (Fig. 7L,M). In other respects, the differences between developing WT and mutant nerves at P7 and P21 already matched those seen in adult P60 nerves. This includes a reduced number of myelinated axons, an increased number of segregated myelin competent axons that remained unmyelinated, thinner myelin sheaths, and an increased number of >1.5 μm diameter axons that were seen within Schwann cell families or Remak bundles (Fig. 6G-K).

These experiments show that c-Jun negatively regulates developmental myelination in a dose-dependent manner. In the c-Jun OE/+ mouse about six fold overexpression of c-Jun results in a transient delay at P7, while the nerve has recovered at P21. On the other hand, in the developing nerves of c-Jun OE/OE mice where c-Jun levels are about 50% higher than in c-Jun OE/+ nerves, myelination is permanently inhibited and seen in only 30-40% of >1.5μm myelin competent axons, a figure comparable to that seen in the adult.

Re-myelination after nerve injury in c-Jun OE/+ mice

C-Jun is a key amplifier of the repair Schwann cell phenotype, which is generated in the distal stump of injured nerves. Therefore, elevation of c-Jun is a candidate approach for improving nerve repair under conditions where it falters, such as in older animals or due to long term Schwann cell denervation (Wagstaff et al., 2017). The observation that in c-Jun OE/+ mice, adult nerves with about six fold elevation of c-Jun achieve a relatively normal degree of nerve architecture and myelination during development, albeit with a delay, is encouraging for this approach, since it demonstrates that significant c-Jun elevation and myelination are compatible. However, after injury, re-myelination is slower and more easily disrupted than developmental myelination. The two processes are also partly controlled by distinct signals. We therefore tested the capacity of c-Jun OE/+ nerves to re-myelinate after nerve injury.

After sciatic nerve crush injury, c-Jun levels distal to the crush were elevated in WT mice and this elevation was enhanced in c-Jun OE/+ mice as expected (Fig. 8A). At one, seven and 14 days after injury, c-Jun levels in OE/+ nerves were two to three fold higher than those in crushed WT control nerves. This amounted to about 12 (at one day after crush) to about 30 (at...
seventeen and 14 days after crush) fold elevation of c-Jun in crushed c-Jun OE/+ nerves compared
to the levels found in uninjured control nerves. This was accompanied by somewhat lower
Krox20 levels (Fig. 8B). At 21 days after crush, c-Jun levels in WT nerves had declined
although they still remained significantly above those in uninjured nerves (data not shown).

When examined four days after nerve cut, nerves of c-Jun OE/+ mice showed accelerated
collapse/breakdown of myelin sheaths, and faster clearance of the myelin protein MBP, in
agreement with previous evidence that c-Jun promotes myelin clearance and myelin autophagy
(Arthur-Farraj et al 2012; Gomez-Sanchez et al. 2015) (Fig. 8C,D).

Examination of crushed c-Jun OE/+ nerves by electron microscopy showed a significant
delay in re-myelination at 2 weeks after nerve crush (Fig. 8E). At this time point, about 35% of
myelin-competent (>1,5μm) axons were myelinated in OE/+ nerves, while over 95% were
myelinated in WT nerves (Fig. 8F). At two weeks after crush, the number of myelinated axons
was also reduced in OE/+ nerves (Fig. 8G) and the number of segregated, myelin-competent
axons without myelin was elevated (Fig. 8H). Significant recovery was seen four weeks after
the crush when about 75% of myelin-competent axons were myelinated in OE/+ nerves compared
to 98% in WT nerves and by 10 weeks, essentially all myelin-competent axons were myelinated
in both genotypes, a situation similar to that in uninjured nerves of these mice (Fig. 8E-H).

Myelin sheaths in adult c-Jun OE/+ mice are thinner than in WT (previous section), and this
difference was also seen in re-myelinated nerves (Fig. 8I). Tumours were not seen, and
regenerating WT and c-Jun OE/+ nerves did not differ in size (Fig. 8J) or other aspects of
general nerve architecture (Fig. 8E).

Importantly, the c-Jun OE/+ mice achieved full functional recovery after nerve crush. In the toe
pinch test, which is primarily a sensory test, time to full recovery was comparable in WT and c-
Jun OE/+ mice, while time to initial response (group average) was about 2 days longer in the
mutants (Fig. 9A,B). In the toe spread reflex, primarily a test of motor recovery, c-Jun OE/+ mice showed a transient delay in recovery on days 14 and 15 only (Fig. 9C), possibly caused by
delay in myelination at this time point (Fig. 8F,G). In the sciatic functional index (SFI) a sensory-
motor test, c-Jun OE/+ mice showed a non-significant trend towards a transient delay during the
second and third week after injury (Fig. 9D,E).

Discussion

We have generated c-Jun OE/+ and c-Jun OE/OE mice with enforced expression of c-Jun
in Schwann cell nuclei to study the effects of a graded increase in c-Jun expression on
Schwann cell development and on re-myelination after injury. This has shown, first, that
during development and in adult nerves, Schwann cells are remarkably tolerant of elevated
c-Jun levels. Although developing and adult nerves of c-Jun OE/+ mice show about six fold increase in c-Jun relative to WT nerves at the same age, myelination is only transiently affected at P7. By P21, myelination appears normal, and in the adult, Schwann cells and nerve architecture is similar to that in WT nerves, with the exception of modestly reduced myelin thickness, which is unlikely to have significant consequences for sensory-motor control. Second, although re-myelination after injury, which generally is more easily perturbed than in development, is delayed in c-Jun OE/+ mice, re-myelination shows strong recovery at four weeks, and 10 weeks after injury essentially all myelin competent axons are myelinated. As in uninjured nerves, the myelin sheaths of regenerated OE/+ nerves remain thinner than those in regenerated WT nerves. The sensory and motor tests used here show only a slight delay followed by complete functional recovery in c-Jun OE/+ mice. Therefore, the c-Jun elevation in c-Jun OE/+ mice is compatible with essentially normal restoration of myelin and nerve function to that found before injury. This is important, because we find that the c-Jun elevation in c-Jun OE/+ mice is sufficient to accelerate regeneration under conditions where it is compromised by ageing or long-term denervation (Wagstaff et al, 2017; LJ Wagstaff, J Gomez-Sanchez, R Mirsky and KR Jessen unpublished). Third, the higher over-expression achieved in c-Jun OE/OE mice confirms the potential of c-Jun to negatively regulate myelination, as previously seen in vitro. Myelination is strongly impaired during development, and this persists in adult nerves, which show hypomyelinating pathology, enlarged connective tissue and immature onion bulbs. Fourth, even in nerves of aged c-Jun OE/OE mice, there is no evidence of tumour formation, and these nerves show strong activation of the tumour suppressor P19ARF. The absence of tumorigenic effect of enforced c-Jun expression in Schwann cells is in agreement with the fact that mechanical nerve damage is not associated with tumour formation, although injured WT nerves contain proliferating cells with high c-Jun levels.

c-Jun is involved directly or indirectly in the control of about 180 of the approximately 4000 genes that change significantly after nerve injury. This allows c-Jun to take part in the regulation of a spectrum of properties of denervated repair Schwann cells, including morphology, autophagy-mediated myelin breakdown, and the expression of trophic factors linked to regeneration, including GDNF, artemin, BDNF, NGF and LIF (Arthur-Farraj et al., 2012; Fontana et al., 2012; Jessen et al 2015a; Gomez-Sanchez, 2015, Jessen and Mirsky, 2016). Of these GDNF, artemin and LIF have been shown to be direct targets of c-Jun. Additional evidence for direct regulation of injury-induced genes by c-Jun comes from a study of enhancer activation in Schwann cells. This showed c-Jun binding sites associated with injury-activated enhancers of genes elevated after nerve injury, including Shh, Olig1 and Runx2 (Hung et al., 2015).
The gene targets and function of AP-1 transcription factors, a family to which c-Jun belongs, are regulated by dimerization partners and ancillary proteins (Chinenov and Kerrpola 2001; Eferl and Wagner, 2003). Little is known about these components in Schwann cells.

The RNA sequencing analysis showed that in uninjured nerves of c-Jun OE/+ mice, 95 genes were expressed at levels that differed ≥2 fold from those in WT nerves. Sixty seven of these genes were up-regulated in response to increased c-Jun levels including Shh. In c-Jun OE/OE nerves, 1964 genes were changed ≥2 fold, and 909 of these were up-regulated, among them GDNF, Shh, Olig1, Id2, Sox2 and Runx2. The myelin genes Mpz, Mbp and Pmp22 were all strongly down-regulated in c-Jun OE/OE nerves. Examining injured nerves, we previously, identified 172 genes that were expressed at different levels in seven day cut nerves of mice in which c-Jun was genetically inactivated compared to unjured WT nerves. Of these 172 genes, 106 genes were up-regulated by higher c-Jun levels, namely expressed more highly in cut WT nerves than in cut c-Jun knockout nerves. A comparison of the 15 genes most up-regulated by c-Jun in uninjured c-Jun OE/+ and OE/OE nerves in the present work, with the 15 genes that are most up-regulated by c-Jun in seven day cut nerves reveals only two common genes, Shh and GDNF. This limited similarity indicates that the group of genes directly or indirectly regulated by c-Jun in Schwann cells that have adopted the repair phenotype after injury is significantly different from the set of genes, which responds to c-Jun in cells that ensheath axons, many of which retain myelin differentiation.

We find that the substantial c-Jun elevation in c-Jun OE/OE mice is sufficient to cause severe hypo-myelination. This is undoubtedly related to the ability of c-Jun to suppress myelin genes. Although the causal relationship between the increased c-Jun levels seen in human CMT1A, and demyelination has not been analysed (Hutton et al., 2011), it seems clear that sustained dys-regulation of c-Jun resulting in high expression in uninjured nerves is a potential hazard. c-Jun is therefore a candidate for a factor that could cause or promote pathological demyelination.

c-Jun OE/OE nerves and nerves affected by demyelinating neuropathies, in particular CMT1A, show many similarities, most obviously hypo-myelination. In c-Jun OE /OE mice, this involves substantially thinner myelin sheaths and about a 40% reduction in myelination among axons that have segregated and are myelin-competent. Axonal sorting is also adversely affected in younger (P60) c-Jun OE/OE mice, since in these mice, the percentage of unsorted >1.5 μm axons that remain in Remak bundles is 28%, compared to <1% in WT. In common with human CMT1A nerves, and nerves of the C22 and My41 mouse models of CMT1A, c-Jun OE/OE nerves also contain increased Schwann cell numbers (Robertson et al., 2002; Lupski and Chance 2005). However, neither mouse models of CMT1A nor the c-
Jun OE/OE mice show significant numbers of Schwann cells that are without axonal contact. Increase in endoneurial connective tissue, which is substantial in c-Jun OE/OE mice, is also a feature of human CMT1A nerves and of the CMT1A rat (Palumbo et al., 2002; Lupski and Chance 2005; Fledrich et al., 2012). Abnormalities of Remak fibres, including the formation of membranous structures that do not contact axons, which are seen in c-Jun OE/OE mice, are also described in the My41, C22 and C3 mouse models of CMT1A (Robertson et al., 2002; Verhamme et al., 2011). Lastly onion bulbs, which are prominent in human CMT1A nerves and seen in rodent CMT1A models (Lupski and Chance 2005; Fledrich et al., 2012), are also present in nerves of aged c-Jun OE/OE mice.

The histological changes outlined here for c-Jun OE/OE and CMT1A nerves, are generally not specific to these conditions, but are also observed to a varying degree in a number of other non-tumour-associated human nerve pathologies, mutant mouse nerves or in injured nerves. (Low 1977; Haney et al., 1999; Chen et al., 2003; Ling et al. 2005; Kagitani-Shimono 2008; Ishii et al., 2016). The relative paucity of disease-specific structural changes in pathological nerves, and the sloppy relationship between molecular and histological phenotype, makes it hard to interpret a particular histology in terms of a causal sequence.

Although the availability of binding partners or other ancillary proteins are important regulators of c-Jun function, the levels of c-Jun protein are likely to be a key factor in determining whether c-Jun has beneficial or adverse effects on nerve biology. Previous work shows that already at low or moderate levels, which are compatible with myelination, c-Jun appears to promote neuron-supportive signalling from Schwann cells to neurons, including the activation of trophic factors, such as GDNF, while higher levels are required to suppress myelin genes. Thus, in the C3 mouse model of CMT1A, c-Jun is elevated, but this is not high enough to disrupt myelin, although it enhances axonal survival and sensory motor performance (Hantke et al., 2014). Similarly, in CMT1X mice, c-Jun is elevated and increases GDNF expression, but does not disrupt myelination (Klein et al., 2014).

In sum, the present results show that although moderate c-Jun increase is well tolerated during Schwann cell development and re-myelination after injury, strong elevation of c-Jun in uninjured nerves suffices to induce significant hypo-myelination pathology, implicating c-Jun in demyelinating neuropathies. On the other hand, we do not find a link between c-Jun elevation and tumorigenesis in line with the fact that tumours do not form after nerve injury, although c-Jun is strikingly elevated as Schwann cells lose myelin differentiation and proliferate. We also find that after crush injury of OE/+ nerves, myelination and nerve function can be restored in the face of c-Jun levels that are high enough to promote axonal
regeneration in mice in which regeneration has been compromised by long term denervation or advanced age.

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Figure legends

Figure 1. Graded over-expression of c-Jun Schwann cell nuclei of c-Jun OE/+ and c-Jun OE/OE mice.

(A) Genomic structure of the c-Jun floxed allele in the Rosa26 locus. Excision of the stop codon is effected by crossing Rosa26c-Junff/+ mice with P0cre expressing mice to generate c-Jun OE/+ and c-Jun OE/OE mice over-expressing c-Jun specifically in Schwann cells.

(B) PCR analysis showing the presence of c-Jun OE, Rosa26 WT and P0cre bands from DNA samples extracted from tails of WT, c-Jun OE/+ and c-Jun OE/OE mice.

(C) Representative immunofluorescence images from WT, OE/+ and OE/OE sciatic nerve cryosections showing Sox10 and c-Jun positive nuclei. Note graded increase in c-Jun in c-JunOE/+ and c-Jun OE/OE mice. Scale bar; 50μm.

(D) Western blot of sciatic nerve protein extracts from P60 WT, c-Jun OE/+ and c-Jun OE/OE mice showing increasing c-Jun levels. The graph quantifies c-Jun expression in WT (n=7), c-Jun OE/+ (n=6) and c-Jun OE/OE (n=6) mice. The quantifications are normalized to the levels in uninjured WT nerves, which are set as 1. Note that the difference in c-Jun expression between c-Jun OE/+ and c-Jun OE/OE nerves is also significant. One-way ANOVA with Tukey comparison; *p<0.05, ****p<0.0001.

(E) Representative immunofluorescence images from purified Schwann cell cultures from WT and c-Jun OE/+ mice. The cells were exposed to neuregulin (nrg) alone, or neuregulin plus cAMP analogue (dbcAMP), a combination that mimics axonal myelination signals. Note that neuregulin plus dbcAMP suppresses c-Jun in WT, but not in c-Jun OE/+, cells. Sox10 was used as a Schwann cell marker to show levels of c-Jun specifically in Schwann cells.

Figure 2. Gene expression in c-Jun OE/+ and c-Jun OE/OE mice.

(A) Heat map of the 400 most regulated genes in uninjured nerves of WT and c-Jun OE/OE mice.

(B) PCA map of gene regulation in WT and c-Jun OE/OE nerves

(C) Expression of 13 genes of interest in the sciatic nerve of OE/+ and OE/OE mice. The table shows how c-Jun elevation affects the expression of a sub-set of repair cell markers, myelin proteins and transcription factors in the mouse lines indicated. Note that in OE/+ nerves, only Shh is regulated ≥2 fold. In OE/OE nerves, repair cell markers are up-
regulated, and myelin genes are down-regulated, although two important myelin regulators, Krox20 and Sox10 are not strongly affected. WT (n=3), OE/+ (n=4) and OE/OE (n=4).

Figure 3. The 15 most up- and down-regulated genes in the sciatic nerve of OE/+ and OE/OE mice.

(A and B) The 15 most strongly elevated genes (the upper panels) and the 15 most suppressed genes (the lower panels) in response to c-Jun elevation in the mouse lines indicated.

(C) The 15 most strongly regulated gene in OE/OE nerves compared to expression in OE/+ nerves.

(D) Representative immunofluorescence images from WT, c-Jun OE/+ and c-Jun OE/OE sciatic nerve cryosections showing Krox20 in Sox10 positive Schwann cell nuclei. Note similar Krox20 expression in WT and c-Jun OE/+ nerves, but much reduced levels in c-Jun OE/OE nerves. Scale bar: 50μm.

(E) Western blot of sciatic nerve protein extracts from P60 mice showing similar levels of Krox20 in WT and c-Jun OE/+ nerves, but lower levels in c-Jun OE/OE nerves. The graph quantifies Krox20 expression in WT (n=5), c-Jun OE/+ (n=4) and c-Jun OE/OE (n=5) mice. The quantifications are normalized to the levels in uninjured WT nerves, which are set as 1. One-way ANOVA with Tukey comparison; *p<0.05, **p<0.01.

(F) Western blot of sciatic nerve protein extracts from P60 mice. Note that Mpz expression is 15% lower than WT in c-Jun OE/+ nerves, but strongly suppressed in c-Jun OE/OE nerves. The graph quantifies Mpz expression in WT (n=5), c-Jun OE/+ (n=4) and c-Jun OE/OE (n=5) mice. The quantifications are normalized to the levels in uninjured WT nerves, which are set as 1. One-way ANOVA with Tukey comparison; ****p<0.0001.

Figure 4. Electron Microscopic structure of adult nerves in WT and OE/+ mice.

(A) Electron micrographs showing similar overall appearance of nerves from P60 WT and OE/+ mice. Scale bar: 5μm.

(B) The total area of P60 WT and OE/+ mouse nerves is not significantly different. Mann-Whitney U test; p=0.5317 (n=5).

(C) The number of Schwann cell nuclei per sciatic nerve profiles not significantly different between P60 WT and OE/+ nerves. Mann-Whitney U test; p=0.0952 (n=5).
(D) Counts of Ki67 positive/Sox10 positive nuclei indicate that the difference in Schwann cell proliferation between WT and c-Jun OE/+ mice is not significantly different. Mann-Whitney U test; p=0.2000 (n=3).

(E) The total number of axons larger than 1.5μm in diameter is similar in P60 WT and OE/+ nerves. Mann-Whitney U test; p=0.9444 (n=5).

(F) The percentage of axons in a 1:1 relationship and greater than 1.5μm in diameter that are myelinated is similar in P60 WT and OE/+ nerves. Mann-Whitney U test; p>0.9999 (n=5).

(G) Per nerve profile, the number of myelinated axons is similar in P60 WT and OE/+ nerves. Mann-Whitney U test; p=0.8016 (n=5).

(H) Per nerve profile, the number of axons in a 1:1 relationship and greater than 1.5μm in diameter but not myelinated is not significantly different between P60 WT and OE/+ nerves. Mann-Whitney U test; p>0.999 (n=5).

(I) Myelin thickness measured by g-ratios is thinner in P60 OE/+ nerves compared to WT. The whiskers extend from the 5th to the 95th percentiles. Mann-Whitney U test; p=0.0079 (n=5).

(J) The percentage of axons greater than 1.5μm in diameter that remain unmyelinated and within Remak bundles is very low and similar in P60 WT and OE/+ nerves. Mann-Whitney U test; p=0.1508 (n=5).

(K) Electron micrographs show that the overall structure of adult P300 nerves in WT and OE/+ mice is similar. Scale bar: 5μm.

(L) The area of transverse profiles of P300 WT (n=4) and OE/+ (n=5) nerves is not statistically different. Mann-Whitney U test; p=0.2857.

(M) The number of Schwann cell nuclei per sciatic nerve profile is somewhat higher in P300 OE/+ (n=5) nerves compared to WT (n=4). Mann-Whitney U test; p=0.0317.

(N) The total number of axons larger than 1.5μm in diameter is similar in P300 WT (n=4) and c-Jun OE/+ (n=5) nerves. Mann-Whitney U test; p=0.1905.

(O) The percentage of axons in a 1:1 relationship and greater than 1.5 μm in diameter that are myelinated is similar in P300 WT (n=4) and OE/+ (n=5) nerves. Mann-Whitney U test; p=0.4444.

(P) The numbers of myelinated axons per nerve profile is similar in P300 WT (n=4) and OE/+ (n=5) nerves. Mann-Whitney U test; p=0.1905.
Figure 5. High c-Jun levels in c-Jun OE/OE nerves result in hypo-myelination.

(A) Electron micrographs showing lack of myelin and increased connective tissue spaces in P60 c-Jun OE/OE nerves compared to WT.

(B) The total number of axons larger than 1.5μm in diameter is not significantly different in P60 WT (n=5) and c-Jun OE/OE (n=3) nerves. Mann-Whitney U test; p=0.0714.

(C) The percentage of axons in a 1:1 relationship and greater than 1.5μm in diameter that are myelinated is lower in c-Jun OE/OE mice than in WT (n=5) and OE/OE (n=3) nerves. Mann-Whitney U test; p=0.0179.

(D) The number of myelinated axons per nerve profile is substantially reduced in c-Jun OE/OE (n=3) nerves compared to WT (n=5). Mann-Whitney U test; p=0.0357.

(E) Per nerve profile, the number of axons in a 1:1 relationship and greater than 1.5μm in diameter but not myelinated is much higher in OE/OE (n=3) nerves than in WT (n=5). Mann-Whitney U test; p=0.0179.

(F) Myelin, measured as g-ratios, is thinner in OE/OE (n=3) mice compared to WT (n=5). The whiskers extend from the 5th to the 95th percentiles. Mann-Whitney U test; p=0.0357.

(G) The percentage of unmyelinated axons greater than 1.5μm in diameter that remain in Remak bundles is higher in OE/OE (n=3) nerves than in WT (n=5) nerves. Mann-Whitney U test; p=0.0357.

(H) Nerves in c-Jun OE/OE mice (n=3) contain more mast cells that nerves in WT mice (n=5). Mann-Whitney U test; p=0.0179.

(I) OE/OE (n=3) nerves show more Schwann cell nuclei per nerve profile than WT (n=5) nerves. Mann-Whitney U test; p=0.0357.
Figure 6. Nerves of aged c-Jun OE/OE mice

(A) The nerves of P300 c-Jun OE/OE mice and WT mice contain comparable numbers of axons. Mann-Whitney U test; p=0.1143 (n=4).

(B) In P300 c-Jun OE/OE mice the percentage of axons in a 1:1 relationship and greater than 1.5μm in diameter that are myelinated is lower than in WT mice. Mann-Whitney U test; p=0.0286 (n=4).

(C) In P300 c-Jun OE/OE mice, the number of myelinated axons per nerve profile is reduced compared to WT. Mann-Whitney U test; p=0.0286 (n=4).

(D) Per nerve profile, P300 c-Jun OE/OE mice have a much larger number of unmyelinated axons that are greater than 1.5μm in diameter and in a 1:1 relationship and, compared to WT mice Mann-Whitney U test; p=0.0286 (n=4).

(E) The percentage of unmyelinated axons greater than 1.5μm in diameter that are found within Remak bundles is higher in OE/OE nerves than in WT nerves. Mann-Whitney U test; p=0.0286 (n=4).
(F) Electron micrographs from nerves of P300 c-Jun OE/OE mice, showing examples of onion bulbs. The central axon, which is sometimes myelinated (upper panels), is surrounded by relatively few layers of flattened Schwann cells, suggesting an early stage of bulb formation. Scale bar = 1 μm.

(G) The number of onion bulbs in P300 OE/OE nerves is much higher than in WT nerves. Mann-Whitney U test; p=0.0286 (n=4).

(H) P300 OE/OE nerves contain a higher number of mast cells than WT nerves. Mann-Whitney U test; p=0.0286 (n=4).

(I) Nerves in P300 c-Jun OE/OE mice show more Schwann cell nuclei per nerve profile than nerves in WT mice. Mann-Whitney U test; p=0.0286 (n=4).

(J) The rate of Schwann cell proliferation is not significantly higher in P300 OE/OE nerves than in WT nerves, judged by counts of Ki67 positive/Sox10 positive nuclei. Mann-Whitney U test; p=0.1000 (n=3).

Figure 7. Developmental over-expression of c-Jun delays myelination in c-Jun OE/+ mice, but inhibits myelination in c-Jun OE/OE mice.

(A) Western blot of nerve extracts from P1 and P7 sciatic nerves. The results are quantified in the graphs. Data from P1 nerves are normalized to levels in P1 WT nerve, which are set as 1, while data from P7 nerves are normalized to levels in P7 WT nerve, which are set as 1. Note that by P7, c-Jun is elevated in both OE/+ and OE/OE nerves, while Mpz is reduced in OE/OE nerves only. P1 WT (n=3), OE/+ (n=3); P7 WT (n=4); OE/+ (n=4), OE/OE (n=3). Statistical analysis for P1 is Student’s T-test; p=0.0608 for c-Jun, p=0.0174 for Mpz.

Statistical analysis for P7 is One-way ANOVA with Tukey comparison; *p<0.05, **p<0.01, ****p<0.0001.

(B) Western blot of nerve extracts from P7 WT, OE/+ and OE/OE nerves. The results are quantified in the graph. Krox20 levels are similar in all genotypes. One-way ANOVA with Tukey comparison; p=0.2053 (n=3).

(C) The percentage of Krox20/Sox10 positive Schwann cells in sections from WT (n=8), OE/+ (n=6) and OE/OE (n=3) sciatic nerves at P7. Note a graded decrease in Krox20 positive cells as levels of c-Jun increase. One-way ANOVA with Tukey comparison; *p<0.05, ***p<0.001.
(D) Representative electron micrographs from P1, P7 and P21 nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Note hypomyelination in OE/OE nerve at P7 and P21, and transient hypomyelination in OE/+ nerves at P7. Scale bar= 5μm.

(E) The nerve areas are similar in all three genotypes at all developmental stages. One-way ANOVA with Tukey comparison; P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5), p=0.1978; P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3), p=0.2261 and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4), p=0.084.

(F) The number of Schwann cell nuclei per sciatic nerve profile at P1, P7 and P21 in nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Note the transient difference between WT and OE/+ nerves at p7, while OE/OE nerves have more Schwann cells at p7 and p21. P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5); P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). One-way ANOVA with Tukey comparison. **p<0.01, ***p<0.001 and ****p<0.0001.

(G) The percentage of axons in a 1:1 relationship and greater than 1.5μm in diameter that are myelinated at P1, P7 and P21 in nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Note reduced myelination in OE/OE mice at P7 and 21, and transient reduction in OE/+ mice at P7. P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5); P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). One-way ANOVA with Tukey comparison; **p<0.01 and ***p<0.001.

(H) The number of myelinated axons per nerve profile at P1, P7 and P21 in nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Note the substantial reduction in myelinated axons in OE/OE nerves at P7 and P21, and transient decrease in OE/+ mice at P7. P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5); P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). One-way ANOVA with Tukey comparison; *p<0.05, **p<0.01, ***p<0.001 and ****p<0.0001.

(I) The number of axons in a 1:1 relationship and greater than 1.5μm in diameter that remain unmyelinated at P1, P7 and P21 in nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Note the increase in unmyelinated axons in OE/OE nerves at P7 and P21, but at p7 only in OE/+ mice. P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5); P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). One-way ANOVA with Tukey comparison; **p<0.01, ***p<0.001 and ****p<0.0001.

(J) In both OE/+ and OE/OE nerves, the reduction in myelin thickness, measured as g-ratios, which is seen in the adults is already present at P7 and p21. P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). The whiskers
extend from the 5th to the 95th percentiles. One-way ANOVA with Tukey comparison, *p<0.05, **p<0.01, and ***p<0.001.

(K) The percentage of unmyelinated axons that are greater than 1.5μm in diameter in Schwann cell families or Remak bundles at P1, P7 and P21 in nerves of WT, c-Jun OE/+ and c-Jun OE/OE mice. Abnormally high numbers are seen in OE/OE nerves only. P1 WT (n=5), OE/+ (n=4) and OE/OE (n=5); P7 WT (n=5), OE/+ (n=5) and OE/OE (n=3); and P21 WT (n=5), OE/+ (n=4) and OE/OE (n=4). One-way ANOVA with Tukey comparison, *p<0.05, **p<0.01, and ***p<0.001.

(L) Western blot of nerve extracts from P7 WT (n=3), OE/+ (n=3) and OE/OE (n=3) sciatic nerves showing Cyclin D1, an indicator of cell proliferation. Quantification of the data is normalized to levels in P7 WT nerve, which are set as 1. Cyclin D1 levels are similar in all mouse lines. One-way ANOVA with Tukey comparison; p=0.3871.

(M) Counts of Ki67 positive/Sox10 positive nuclei in P7 in WT (n=6), OE/+ (n=6) and OE/OE (n=3) nerves. OE/OE nerves show increased Schwann cell proliferation. One-way ANOVA with Tukey comparisons, *p=0.0121. In Figs. E-K, p values are calculated relative to WT at the same age.

Figure 8. Re-myelination of OE/+ nerves is delayed

(A) Western blot of c-Jun in nerve extracts from the distal stump of adult WT (n=4) and OE/+ (n=4) nerves 1 day, 7 days and 2 weeks after crush. The graph shows quantification of the results, normalized to levels in uninjured WT nerves, which are set as 1. Note significant elevation of c-Jun at all time points. Mann-Whitney U test: 1 day p=0.0006 (n=4); 7 days p=0.0002 (n=4); and 2 weeks p=0.0022(n=4).

(B) Western blot of nerve extracts from the distal stump of adult WT and OE/+ nerves 2 weeks after crush. The results are quantified in the graph, normalized to levels in 2 week crushed WT nerve, which are set as 1. Krox20 levels are reduced in OE/+ nerves. Mann-Whitney U test, p=0.0286, (n=4).

(C) Representative electron micrographs from the distal stump 4 days after sciatic nerve cut in WT and c-Jun OE/+ mice, illustrating collapsed myelin sheaths. The graph shows that fewer intact myelin sheaths per nerve profile remain in OE/+ nerves than in WT; Mann-Whitney U test; p=0.0286 (n=4).

(D) Transected c-Jun OE/+ nerves clear myelin protein faster than WT nerves. The graph shows the reduction in MBP 4 days after transection expressed as a percentage of MBP in
uninjured nerve. WT and c-Jun OE/+ nerves have cleared close to 40% and 60% of their MBP content, respectively. The data are obtained from quantitation of Western blots. WT (n=4) and OE/+ (n=8). Mann-Whitney U test; p=0.0070.

(E) Representative electron micrographs from the distal stump of WT and OE/+ nerves 2, 4 and 10 weeks after crush. In OE/+ nerves, the number of myelinated axons, which is reduced at 2 and 4 weeks, has recovered at 10 weeks. Scale bar: 5μm.

(F) The percentage of axons greater than 1.5μm in diameter and in a 1:1 ratio that are myelinated in the distal stump of WT and c-Jun OE/+ mice 2, 4 and 10 weeks after nerve crush. Note that myelination in OE/+ nerves, which is reduced at 2 weeks, has recovered substantially by 4 weeks and is normal at 10 weeks. Mann-Whitney U test: 2 weeks p=0.0286, n=4; 4 weeks P=0.0079, n=5; and 10 weeks p=0.0571 (n=4).

(G) The number of myelinated axons per nerve profile of the distal stump of WT and c-Jun OE/+ mice 2, 4 and 10 weeks after nerve crush. In c-Jun OE/+ mice, few myelinated axons are present at 2 weeks, but normal numbers are seen at 10 weeks. Mann-Whitney U test: 2 weeks p=0.0286 (n=4); 4 weeks p=0.0079 (n=5); and 10 weeks p=0.0571 (n=4).

(H) The number of unmyelinated axons greater than 1.5μm in diameter and in a 1:1 relationship that have not myelinated in the distal stump of WT and OE/+ nerves 2, 4 and 10 weeks after crush. Two and 4 week crushed OE/+ nerves contain elevated numbers of unmyelinated axons, but their number has fallen to normal levels at 10 weeks. Mann-Whitney U test: 2 weeks p=0.0286 (n=4); 4 weeks P=0.0079 (n=5); and 10 weeks p=0.1143 (n=4).

(I) Myelin thickness, measured as g-ratios, is reduced in the distal stump of OE/+ nerves at all time points after crush. The whiskers extend from the 5th to the 95th percentiles. Mann-Whitney U test: 2 weeks p=0.0286 (n=4); 4 weeks p=0.0079 (n=5); and 10 weeks p=0.0286 (n=4).

(J) The area of transverse sections through the distal stump 2, 4 and 10 weeks after crush is similar in WT and OE/+ nerves. Mann-Whitney U test: 2 weeks p=0.6571 (n=4); 4 weeks p=0.3095 (n=5); and 10 weeks p=0.9004 (n=4).

In Figs. A and F-J, p values are calculated relative to WT at same time after injury.

Figure 9. Functional recovery is slightly delayed in c-Jun OE/+ mice.
(A) Toe pinch assay, showing the percentage of mice that show a response to a pinch of toes 3, 4 and 5 at different times after sciatic nerve crush in WT and c-Jun OE/+ mice. In c-Jun OE/+ mice, all toes show a trend towards a delayed response.

(B) The average time in days after crush at which the first toe pinch response is seen in toe 3, toe 4 and toe 5. WT (n=10) and OE/+ (n=9). Mann-Whitney U test; p values are calculated relative to WT for each toe. Toe 3 p=0.0056, Toe 4 p=0.0043, Toe 5 p=0.0404.

(C) The toe spread reflex in WT (n=10) and OE/+ (n=9) mice following sciatic nerve crush. The reflex response is delayed at day 12, 14 and 15 in c-Jun OE/+ mice. Two-way ANOVA with Bonferroni comparison, p=0.0171.

(D) Representative digital footprints from WT and c-Jun OE/+ mice taken at 0, 7, 18, 21, 28 and 70 days after sciatic nerve crush, used in sciatic functional index (SFI) analysis.

(E) SFI results from WT (n=11) and c-Jun OE/+ (n=8) mice at different times after sciatic nerve crush. There is no significant difference between WT and c-Jun OE/+ mice. Two-way ANOVA with Bonferroni comparison, p=0.5545.
### C

| Repair Schwann cell markers | Gene                          | OE/+ vs WT Fold Change | OE/+ vs WT Log2(FC) | OE/OE vs WT Fold Change | OE/OE vs WT Log2(FC) | OE/OE vs OE/+ Fold Change | OE/OE vs OE/+ Log2(FC) |
|-----------------------------|-------------------------------|------------------------|---------------------|-------------------------|-----------------------|--------------------------|-------------------------|
|                             |                               |                        |                     |                         |                       |                          |                         |
|                             | c-Jun transcription factor AP-1-like | 1.53                   | 0.61                |                         |                       |                          |                         |
|                             | sIhh sonic hedgehog           | 5.21                   | 2.38                |                         |                       |                          |                         |
|                             | Gdnf glial cell line derived neurotrophic factor | 1.82                   | 0.86                |                         |                       |                          |                         |
|                             | Rabf brain derived neurotrophic factor | 0.77                   | -0.37               |                         |                       |                          |                         |
|                             | Olig1 oligodendrocyte transduction factor 1 | 1.24                   | 0.31                |                         |                       |                          |                         |
| Myelin proteins             | Mps myelin protein zero       | 0.65                   | -0.62               |                         |                       |                          |                         |
|                             | Mbp myelin basic protein      | 0.75                   | -0.41               |                         |                       |                          |                         |
|                             | Ptnp22 peripheral myelin protein 22 | 0.95                   | -0.07               |                         |                       |                          |                         |
| Transcriptional Factors     | Krox20 early growth response 2 | 0.96                   | -0.06               |                         |                       |                          |                         |
|                             | Sry2 inhibitor of DNA binding 2 | 2.11                   | 1.08                |                         |                       |                          |                         |
|                             | Sry2 (sex determining region Y)-box 2 | 1.48                   | 0.51                |                         |                       |                          |                         |
|                             | Sry2 (sex determining region Y)-box 10 | 0.58                   | -0.78               |                         |                       |                          |                         |
|                             | Runx2 runt related transcription factor 2 | 1.56                   | 0.64                |                         |                       |                          |                         |
