Sarm1 deletion, but not WldS, confers lifelong rescue in a mouse model of severe axonopathy

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SUMMARY

Studies with the Wld<sup>S</sup> mutant mouse have shown that axon and synapse pathology in several models of neurodegenerative diseases are mechanistically related to injury-induced axon degeneration (Wallerian degeneration). Crucially, an absence of SARM1 delays Wallerian degeneration as robustly as Wld<sup>S</sup>, but their relative capacities to confer long-term protection against related, non-injury axonopathy and/or synaptopathy have not been directly compared. Whilst Sarm1 deletion or Wld<sup>S</sup> can each rescue perinatal lethality and widespread Wallerian-like axonopathy in young NMNAT2-deficient mice, we now report that an absence of SARM1 enables these mice to survive into old age with no overt phenotype whereas those rescued by Wld<sup>S</sup> invariably develop a progressive neuromuscular defect in their hindlimbs from around 3 months of age. We therefore propose Sarm1 deletion as a more reliable tool than Wld<sup>S</sup> for investigating Wallerian-like mechanisms in disease models and suggest that SARM1 blockade may have greater therapeutic potential than WLD<sup>S</sup>-related strategies.
INTRODUCTION

*Wld*<sup>S</sup>, a spontaneous mutant mouse allele encoding a fusion protein (*WLD*<sup>S</sup>) with nicotinamide mononucleotide adenylyltransferase (NMNAT) activity, robustly delays injury-induced axon and synapse degeneration (Wallerian degeneration) by locally substituting for loss of the endogenous NMNAT2 isoform (Mack et al., 2001; Gilley and Coleman, 2010; Cohen et al., 2012; Conforti et al., 2014). *Wld*<sup>S</sup> has been the tool of choice for investigating the molecular basis of axon pathology in animal models of neurodegenerative diseases and has revealed an involvement of Wallerian-like mechanisms in several cases (Conforti et al., 2014). Key steps in this process are thus potential targets for intervention in patients.

Sterile alpha and TIR motif-containing protein 1 (*SARM1*) acts downstream of NMNAT2 loss to promote axon degeneration (Osterloh et al., 2012; Gerdts et al., 2013; Gilley et al., 2015; Loreto et al., 2015; Walker et al., 2017). Depletion of SARM1 is, to date, the only other manipulation that can delay Wallerian degeneration and related axon degeneration in mice as robustly as exogenous expression of *WLD*<sup>S</sup> or other NMNAT variants (Conforti et al., 2014), but its effectiveness in maintaining long-term health of axons and synapses in mouse models of axonopathy and/or synaptopathy has not yet been directly compared to *Wld*<sup>S</sup>. Such a comparison is needed to ascertain the relative usefulness of *Sarm1* deletion in determining whether Wallerian-like mechanisms are involved in models of neurodegeneration and should also be informative in terms of therapeutic strategies for those disorders.

An absence of NMNAT2 in mice causes widespread axon truncation during embryogenesis and perinatal lethality (Gilley et al., 2013). Interestingly, early rescue by *Wld*<sup>S</sup> (dose-dependently) or by *Sarm1* deletion has shown that outgrowth of NMNAT2-deficient axons stalls due to a Wallerian-like degenerative mechanism (Gilley et al., 2013; Gilley et al., 2015). Reduced NMNAT2 levels have already been linked to tauopathy in mice and to decreased cognitive function in humans.
(Ljungberg et al., 2012; Ali et al., 2016), but the severity of the phenotype in mice lacking NMNAT2 suggests a complete lack of the protein is unlikely to directly model any neurodegenerative conditions. Nevertheless, these mice do represent a well-defined and robust system for comparing the longer-term protective effects of \textit{Wld}^{S} and \textit{Sarm1} deletion against a severe Wallerian-like axonopathy. Whilst the survival of NMNAT2-deficient mice homozygous either for \textit{Wld}^{S} or a \textit{Sarm1} knockout allele up to 3 months of age with no overt problems initially suggested similarly robust rescue in each case (Gilley et al., 2013; Gilley et al., 2015), we now report striking age-dependent differences between the two lines which are likely to have important experimental and therapeutic implications.
RESULTS

Locomotor defects and muscle atrophy in \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice, but not \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}^{-/-} mice

Mice homozygous for the \textit{Nmnat2}^{gtE} gene trap allele, lacking NMNAT2, that are additionally homozygous for \textit{Wld}^{S} or a \textit{Sarm1} knockout allele (\textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} or \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}^{-/-} mice respectively) are born at the expected frequencies and are outwardly indistinguishable from NMNAT2-expressing littermates up to three months of age (Gilley et al., 2013; Gilley et al., 2015) (Table S1). However, despite continued silencing of the trapped \textit{Nmnat2} alleles in each case (Fig. S1A), further aging has revealed clear differences between the lines: \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}^{-/-} mice remarkably survived for up to 2 years with no noticeable behavioral deficiency or phenotype, whereas \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice invariantly developed a conspicuous, progressive hindlimb defect from around 3-5 months of age.

The defect in \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice (male and female) first presented as a modest hindlimb gait abnormality during spontaneous locomotion, but this progressively deteriorated resulting in mice invariantly dragging their hindlimbs regularly during locomotion from around 6 months onwards as a result of worsening paraparesis (Movies 1-4). Consistent with this, locomotor ability of \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice in an accelerating Rotarod task deteriorated rapidly between 4 and 6 months (Fig. 1A). Movement became so limited by 10-12 months that it impaired free access to food and water so \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice were not aged further. In contrast, locomotor performance of \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}^{-/-} mice did not decline during the same period (Fig. 1A and Movies 5 and 6) and \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}^{-/-} mice still performed as well as \textit{Sarm1}^{-/-} controls up to at least 15 months (Fig. 1B).

Deteriorating locomotor function in \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice coincided with progressive and widespread wasting of hindlimb muscles (Fig. S1B). A specific
analysis of gastrocnemius muscle revealed evidence of some muscle fiber atrophy and slightly reduced muscle weight even at 10 weeks in \textit{Nmnat2}^{gtE/gtE}; \textit{Wld}\textsuperscript{S/S} mice, prior to the onset of overt locomotor dysfunction, but this had progressed to severe muscle fiber atrophy and loss of mass by 10 months (Fig. 1C and 1D). In contrast, no muscle fiber atrophy or loss of mass was seen in \textit{Nmnat2}^{gtE/gtE}; \textit{Sarm1}\textsuperscript{-/-} gastrocnemius up to 24 months (Fig. 1C-E).

Interestingly, although \textit{Nmnat2}^{gtE/gtE}; \textit{Wld}\textsuperscript{S/S} mice did not lose body weight between 10 weeks and 10-12 months, they did fail to gain weight as expected (Fig. S1C). This presumably reflected loss of muscle mass in the hindlimb being broadly matched by normal weight gain in the upper torso and forelimbs, which appeared largely unaffected. In contrast, developmental weight gain in \textit{Nmnat2}^{gtE/gtE}; \textit{Sarm1}\textsuperscript{-/-} mice was comparable to \textit{Sarm1}\textsuperscript{-/-} controls up to 24 months (Fig. S1D).

\textbf{Neuromuscular denervation in \textit{Nmnat2}^{gtE/gtE}; \textit{Wld}\textsuperscript{S/S} mice, but not \textit{Nmnat2}^{gtE/gtE}; \textit{Sarm1}\textsuperscript{-/-} mice}

Changes in neuromuscular junction (NMJ) innervation indicated that the muscle defect in \textit{Nmnat2}^{gtE/gtE}; \textit{Wld}\textsuperscript{S/S} mice is neurogenic. Motor endplate occupancy in \textit{Nmnat2}^{gtE/gtE}; \textit{Wld}\textsuperscript{S/S} gastrocnemius was found to be already moderately reduced at 10 weeks, but, consistent with the timing of gastrocnemius muscle fiber atrophy and weight loss in these mice, by 10 months denervation was extensive with only around 10 per cent of endplates showing normal innervation (Fig. 2A and 2B).

We also investigated motor endplate occupancy in a more distal hindlimb muscle, flexor digitorum brevis (FDB), at 10 months. Whilst muscle denervation was also evident in this muscle (Fig. 3A), it was actually less severe than in gastrocnemius at the same age. Interestingly, clear regional variation in the pattern of endplate occupancy was seen in FDB with discrete zones of normal innervation being found adjacent to zones of complete denervation (Fig. 3B and 3C). Isometric tension recordings indicated that this distinctive pattern of NMJ innervation reflects
discrete loss of entire motor units (Fig. 3D and 3E) with apparently normal function of remaining motor units (Fig. S2).

In contrast, and consistent with the lack of any locomotor problems, no significant endplate denervation and/or motor unit loss was evident in either gastrocnemius or FDB muscles from 10-month-old Nmnat2\textsuperscript{gtE/gtE};Sarm1\textsuperscript{−/−} mice (Fig. 2, Fig. 3A, 3D and 3E, and Fig. S2) and innervation remained comparable to Sarm1\textsuperscript{−/−} controls at 24 months, despite modest age-dependent denervation in both (Fig. 2A).

Importantly, Wld\textsuperscript{S/S} and Sarm1\textsuperscript{−/−} mice both performed maximally in locomotor tests and neither showed any signs of neuromuscular denervation at the ages studied. Although a direct comparison with wild-type mice will be needed to establish if more subtle differences in motor function exist in either line, our data suggest that both have broadly normal neuromuscular function. The defect in Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S} mice thus appears to be specific to a declining inability of WLD\textsuperscript{S} to counter the lack of NMNAT2 in older mice rather than other intrinsic differences.

**No loss of myelinated tibial nerve axons in either Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S} or Nmnat2\textsuperscript{gtE/gtE};Sarm1\textsuperscript{−/−} mice**

Despite progressive denervation of motor endplates in hindlimb muscles from Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S} mice, no concurrent loss of myelinated axons was seen in the tibial nerve (Fig. 4A) and axons remained morphologically normal (Fig. 4B). A gross assessment revealed that most hindlimb muscles became extensively atrophied in these mice (Fig. S1B) so significantly reduced numbers of axons would have been expected, even in a mixed nerve such as this, if motor axon loss was the underlying cause. The age-dependent neuromuscular denervation in Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S} muscles thus appears to result from selective loss of the distal ends of motor axons and/or their terminals. Interestingly, this mirrors the age-dependent loss of protection of synapses at NMJs after axotomy in homozygous Wld\textsuperscript{S} mice, despite continued protection of the main body of the transected axon (Gillingwater et al., 2002).
Counts of myelinated axons in $Nmnat^{2EgE^{-}};Sarm1^{-/-}$ tibial nerves also remained comparable to $Sarm1^{-/-}$ controls even up to 24 months (Fig. 4A) with no age-related axon loss in either group up to this age. We also found no significant axon loss in a separate cohort of wild-type mice (on a related background) up to 24 months ($1514 \pm 22$ myelinated axons at 1.5 months compared to $1473 \pm 40$ at 24 months). This contrasts a previous study that reported significant loss of myelinated tibial nerve axons by 24 months in wild-type mice (Valdez et al., 2010), although this could simply reflect strain differences.
To date, \textit{Wld}^S has been the preferred tool for assessing the involvement of Wallerian-like axon and synapse degeneration in rodent models of neurodegeneration (Conforti et al., 2014). However, the relatively short-term preservation of neuromuscular innervation by \textit{Wld}^S in hindlimb muscles of \textit{Nmnat2}^{gtE/gtE} mice raises the possibility that this strategy might have greatly underestimated the involvement of Wallerian-like mechanisms in some models. Likely candidates are wabbler-lethal (\textit{Atp8a2}^{wl/wl}) and gracile axonal dystrophy (\textit{Uchl1}^{gad/gad}) mice where \textit{Wld}^S robustly protects (proximal) axons but does not rescue neuromuscular symptoms (Mi et al., 2005; Zhu et al., 2012). Human SOD1\textsuperscript{G37R}, SOD1\textsuperscript{G85R} and SOD1\textsuperscript{G93A} transgenic mouse models of amyotrophic lateral sclerosis (ALS) are also candidates, although \textit{Wld}^S largely fails to even protect axons in these models suggesting that unrelated degenerative mechanisms contribute substantially to symptoms (Vande Velde et al., 2004; Fischer et al., 2005). Crucially, because SARM1 deficiency confers longer-lasting preservation of NMJ innervation in \textit{Nmnat2}-deficient mice than \textit{Wld}^S, it could also confer a better outcome in these or related models.

We consider that prolonged preservation of \textit{Nmnat2}^{gtE/gtE};\textit{Sarm1}/\textit{Sarm1} motor axon terminals compared to those in \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} mice might reflect that local availability of WLD\textsubscript{S}, which is required for protection (Beirowski et al., 2009; Cohen et al., 2012), is likely to be subject to a variety of influences, whereas protection conferred by an absence of SARM1 will be invariant. Global expression of WLD\textsubscript{S} in homozygous \textit{Wld}^S mice does not diminish significantly with age up to 12 months (Gillingwater et al., 2002), but we propose that normal changes in physiology, from as young as 2 months of age, could alter the stability, delivery or activity of WLD\textsubscript{S} in \textit{Nmnat2}^{gtE/gtE};\textit{Wld}^{S/S} motor axon terminals, or otherwise alter the local environment, such that it can no longer effectively substitute for the lack of NMNAT2 to promote survival. Notably, more widespread NMJ denervation in gastrocnemius compared to
FDB in \textit{Nmnat2^{gtE/gtE}, Wld^{S/S}} mice suggests that changes specific to different muscle or motor unit types are more critical to the loss of WLD^{S}-mediated protection than those relating to axon length (FDB being more distal). These considerations will similarly apply to the age-dependent loss of protection of motor axon terminals after axotomy (and the resulting NMNAT2 loss) in homozygous \textit{Wld^{S}} mice (Gillingwater et al., 2002). Interestingly, annulospiral (sensory) nerve endings in the muscle remained protected in older mice in that context, suggesting a motor-specific defect (Oyebode et al., 2012). Whilst we have also seen qualitative preservation of annulospiral endings in \textit{Nmnat2^{gtE/gtE}, Wld^{S/S}} FDB at 10 months (not shown), a comprehensive analysis of sensory innervation will be required to determine whether sensory endings in general are better preserved than motor axon terminals.

Intriguingly, a model in which sustaining the effective potency of WLD^{S} locally is required for its protective effects leaves open the possibility that \textit{Sarm1} deletion may also be more effective than \textit{Wld^{S}} at rescuing symptoms in models of other types of neurodegenerative disease, not just those with early neuromuscular symptoms. If the disease-causing defect in a given model additionally reduces the activity or concentration of WLD^{S} within axons or synapses in some way, then its protective capacity might also be diminished. This could apply to disorders of axonal transport, protein synthesis or protein turnover, among others.

Crucially, our findings have important therapeutic implications for human disorders. Specifically, they suggest that strategies directed at blockade of SARM1 function have the potential to be more effective than WLD^{S}-related therapies in neuromuscular synaptopathies, and potentially also in a much broader group of neurodegenerative disorders. In addition, the remarkable survival and health of \textit{Nmnat2^{gtE/gtE}, Sarm1^{−/−}} mice into old age suggests that even long-term therapeutic interventions based on blocking SARM1 function might be both effective and well tolerated by patients.
This study also confirms SARM1 as a key regulator of degeneration caused by a NMNAT2 deficiency. Remarkably, Sarm1 deletion appears to block this process indefinitely without affecting long-term survival, even despite the predicted substantial reduction in NAD-synthesizing capacity (Gilley et al., 2015). SARM1 has recently been shown to possess NADase activity that promotes injury-induced axon/synapse degeneration and which can be inhibited by NMNAT activity (Gerdts et al., 2015; Sasaki et al., 2016; Essuman et al., 2017). Therefore, a model in which survival depends on NMNAT-dependent NAD production balancing NAD consumption, including any resulting from constitutive SARM1 NADase activity, is attractively simple. However, NAD consumption in uninjured Sarm1−/− axons has been shown to be comparable to wild-type consumption suggesting that SARM1 NADase activity under normal conditions is minimal (Sasaki et al., 2016). Instead, a situation in which a loss of NMNAT activity results in the up-regulation of SARM1 NADase activity via intermediate signals and/or interactions to trigger degeneration is more consistent with current findings (Sasaki et al., 2016; Di Stefano et al., 2017).

Finally, the similarity between the progressive phenotype in Nmnat2gtE/gtE; Wld<sup>S</sup>S mice and some mouse models of ALS is intriguing. Whilst there is, as yet, no established link between NMNAT2 and ALS, the SARM1 locus has been associated with sporadic ALS (Fogh et al., 2014) hinting at an involvement of Wallerian-like mechanisms. The neuromuscular defect in Nmnat2gtE/gtE; Wld<sup>S</sup>S mice could thus model some aspects of ALS disease pathogenesis. Even if the underlying mechanisms are unrelated, Nmnat2gtE/gtE; Wld<sup>S</sup>S mice would still be a useful tool for assessing reversibility of ALS-like symptoms as silencing of the Nmnat2<sup>gtE</sup> gene trap allele is conditional (Gilley et al., 2013).
EXPERIMENTAL PROCEDURES

Mouse breeding and maintenance.
Animal work was performed in accordance with the Animals (Scientific Procedures) Act, 1986, under Project License PPL 70/7620. Genotyping for the Nmnat2^{gtE}, Wld\(^S\) and Sarm1 knockout alleles was performed as described previously (Gilley et al., 2013; Gilley et al., 2015). Littermates were used where possible. The ages and genders of mice used in each individual experiment are described in Figure legends.

Reverse transcriptase-PCR (RT-PCR).
Semi-quantitative endpoint RT-PCR was used to confirm Nmnat2 gene silencing in the brains of Nmnat2^{gtE/gtE};Wld^{S/S} and Nmnat2^{gtE/gtE};Sarm1^{+/+} mice aged 10-12 months essentially as described previously (Gilley et al., 2013).

Accelerating Rotarod task.
Locomotor performance was tested on an accelerating Rotarod (Ugo Basile, Model 7650, Varese, Italy). Mice were familiarized with the apparatus (2x 5 min runs at 10 rpm) one day before testing. At each test age mice performed 3x 5 min trials (3 to 30 rpm) separated by 30 min rests. Latency to fall (max 300 seconds) was recorded. Only involuntary falls were scored. Mice dismounting voluntarily were placed back onto the apparatus once but the run was excluded from the analysis if repeated. Best trial performance was used for statistical analyses.

Haematoxylin and eosin (H&E) staining.
Transverse cryosections of gastrocnemius muscles snap frozen in liquid nitrogen-chilled isopentane (8 µm thickness) or fixed in 4% paraformaldehyde (20 µm thickness) were stained with haematoxylin and eosin as previously described (Gilley et al., 2013). Images were captured using a MicroPublisher camera (QImaging) on
an Olympus BX50 microscope (20x objective). Staining of snap-frozen muscle sections was optimal for visualization of muscle structure without the artefactual muscle fiber separation seen on fixed sections.

**NMJ innervation.**

Innervation of neuromuscular junctions (NMJs) in gastrocnemius and flexor digitorum brevis (FDB) muscles was assessed by immunofluorescent staining. Staining was performed essentially as described previously (Krieger et al., 2013) on whole-mount muscles or longitudinal cryosections (60 µm thickness). Confocal z-stack series were acquired using Olympus FV1000 or Leica SPE scanning laser confocal microscopes (20x or 40x objectives). Multiple z-stack series were acquired for each muscle and z-projections generated for analysis. Endplate occupancy was determined by assessing the extent of overlap or direct abuttal of βIII-tubulin staining (axon terminal) with α-bungarotoxin staining (endplate). Endplates were scored as denervated when essentially none of the endplate (less than ~5%) was deemed occupied by the axon terminal, fully innervated with complete (greater than ~95%) occupancy, and partially innervated with intermediate occupancy (observer determined, scored blind). Original z-stack series were examined to exclude chance overlay of proximal axon segments and endplates in non-adjacent focal planes.

**Isometric muscle tension recordings.**

Force measurements for flexor digitorum brevis (FDB) muscle were made from FDB muscle-tibial nerve preparations as described previously (Beirowski et al., 2009), except that the proximal tendon was connected to a MLT0202 (0-25g) isometric force transducer (AD Instruments Ltd, Oxford, UK) and the tibial nerve was stimulated using 0.1-0.2 ms pulses of up to 10 V using a Digitimer DS2 isolated stimulator (Digitimer, Welwyn Garden City, UK) triggered via a Powerlab 26T interface. Tension
responses were digitized at 1kHz using Chart 7 or Scope 4 software (all ADInstruments Ltd.).

Counts of myelinated tibial nerve axons.
Transverse sections (20 µm) of fixed calf (from mid-way between knee and ankle) were stained with FluoroMyelin™ Red according to the manufacturer's instructions (Life Technologies). Images of tibial nerves were captured on an Olympus FV1000 point scanning confocal microscope imaging system (40x objective). Axon counts (inferred from numbers of myelin sheaths) were performed blind using the multi-point selection tool in ImageJ.

Statistical analysis.
Appropriate statistical testing of data was performed using Prism (GraphPad Software Inc., La Jolla, USA). Tests are described in Figure legends. A p value < 0.05 was considered significant.
AUTHOR CONTRIBUTIONS

Conceptualization, J.G. and M.P.C; Methodology, J.G. and R.R.R.; Investigation, J.G. and R.R.R.; Writing - Original Draft, J.G.; Writing - Review & Editing, J.G., R.R.R. and M.P.C; Visualization, J.G.; Funding Acquisition, M.P.C.; Supervision, M.P.C.

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Figure 1. Progressive locomotor dysfunction and hindlimb muscle atrophy in \textit{Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S}} mice, but not \textit{Nmnat2\textsuperscript{gtE/gtE};Sarm1\textsuperscript{−/−}} mice. (A and B) Latency to fall in an accelerating Rotarod task for mice of the indicated genotypes and ages. Means ±SEM are plotted (maximum test duration 300 seconds) for \(n = 7/8\) male mice of each genotype in (A) (***>0.001 in two-way ANOVA with Dunnett's multiple comparisons), and \(n = 7\) female mice each genotype in (B) (n.s. not significant [\(p = 0.29\)] in \(t\) test). (C) Representative transverse sections of gastrocnemius muscle for mice of the selected genotypes and ages (as indicated) stained with haematoxylin and eosin. Modest fiber atrophy is seen in \textit{Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S}} gastrocnemius at 10 weeks and more severe atrophy with centrally-located nuclei, hypertrophic fibers (*) and pyknotic nuclear clumps (arrow) are all evident at 10 months. Images are representative of male and female mice at 10 weeks and 8-12 months but just female mice at 24 months. (D and E) Gastrocnemius muscle weights for mice of the indicated genotypes and ages. Individual animal values with means ±SEM are plotted for \(n = 3-5\) male mice per group in (D) (*\(p < 0.05\) and ***>0.001 in one-way ANOVA with Tukey's multiple comparisons, n.s. = not significant), and \(n = 4-5\) female mice per group in (E) (*\(p < 0.05\) in \(t\) test). See also Movies 1-6, Table S1 and Figure S1.

Figure 2. Progressive loss of NMJ innervation in gastrocnemius muscles of \textit{Nmnat2\textsuperscript{gtE/gtE};Wld\textsuperscript{S/S}} mice, but not \textit{Nmnat2\textsuperscript{gtE/gtE};Sarm1\textsuperscript{−/−}} mice. (A) Percentage of fully occupied (full), partially occupied (partial) and denervated endplates in gastrocnemius muscles from mice of the indicated genotypes and ages. Endplate occupancy was determined by assessing signal overlap between \(\alpha\)-bungarotoxin labeling of acetylcholine receptors in the motor endplate and \(\beta\text{III}-\text{tubulin}\) immunostaining of axon terminals (see Experimental Procedures). Total numbers of
NMJs analyzed in muscles from 3-5 mice per genotype (male and female) are listed at the base of each column. Means ±SEM of occupancy per animal are plotted (n.s. not significant \( p > 0.05 \) or ***\( p < 0.001 \) in one-way ANOVA with Tukey’s multiple comparisons of full innervation, selected comparisons). (B) Representative confocal z-series projections of fixed gastrocnemius (Gn) muscle preparations showing merged \( \alpha \)-bungarotoxin (red) and \( \beta \)III-tubulin (green) signals. Endplates in the images are marked as fully occupied (asterisks), partially occupied (arrows) or denervated (arrowheads).

Figure 3. Localized loss of NMJ innervation from distinct motor unit groups in flexor digitorum brevis (FDB) muscle of Nmnat2gtE/gtE; WldS/S mice. (A) Percentage of fully occupied (full), partially occupied (partial) and denervated endplates in FDB muscles of 10-month-old mice of the indicated genotypes (method as in Fig. 2A). Total numbers of NMJs analyzed in muscles from 5-9 mice per genotype (male and female) are listed at the base of each column. Means ±SEM of occupancy per animal are plotted (n.s. not significant \( p > 0.05 \) or **\( p < 0.01 \) in one-way ANOVA with Tukey’s multiple comparisons of full innervation, selected comparisons). (B) Percentage of full occupancy of endplates per imaged field revealed a zonal pattern of denervation in Nmnat2gtE/gtE; WldS/S FDB at 10 months with more widespread denervation in gastrocnemius (Gn). Fields with normal levels of full occupancy (80-100%) were relatively common in FDB (7/18) but absent in gastrocnemius (average of 16.9 and 11.6 endplates per field respectively). (C) Representative confocal z-series projections of fixed FDB muscle preparations showing merged \( \alpha \)-bungarotoxin (red) and \( \beta \)III-tubulin (green) signals highlighting regional variability in Nmnat2gtE/gtE; WldS/S endplate occupancy at 10 months. In addition to regions with widespread full occupancy (top panel) or denervation (middle panel), there were also areas containing degenerated endplates (bottom panel, not included in quantifications). (D) Isometric twitch tension recordings for representative
**Figure 4.** No loss of myelinated tibial nerve axons in either \( \text{Nmnat2}^{gtE/gtE;Wld^{S/S}} \) or \( \text{Nmnat2}^{gtE/gtE;Sarm1^{-/-}} \) mice. (A) Numbers of myelinated axons in tibial nerve (mid-calf level) from mice of the indicated genotypes and ages \( (Wld^{S/S} \text{ control groups include some } \text{Nmnat2}^{gtE/gtE;Wld^{S/S}} \text{ mice that were indistinguishable from } Wld^{S/S} \text{ mice}) \). Individual values (\( n = 3-8, \text{ as shown, male and female} \) and means ±SEM are plotted. No statistically significant differences were identified between groups (one-way ANOVA with Tukey’s multiple comparisons). (B) FluoroMyelin™ Red-stained tibial nerve cross-sections from 10-month-old \( Wld^{S/S} \) and \( \text{Nmnat2}^{gtE/gtE;Wld^{S/S}} \) mice (representative of \( n = 5 \) each genotype). No structural differences are evident even though \( \text{Nmnat2}^{gtE/gtE;Wld^{S/S}} \) mice have an advanced neuromuscular defect at this age.

**SUPPLEMENTAL MOVIE TITLES**

Movie 1. 9.5-month-old female \( \text{Nmnat2}^{gtE/gtE;Wld^{S/S}} \) mouse with advanced hindlimb defect. Related to Figure 1.
Movie 2. 8-month-old female $Nmnat^{2gtE/gtE}; Wld^{SS}$ mouse with moderate hindlimb defect. Related to Figure 1.

Movie 3. 8-month-old female $Nmnat^{2gtE/gtE}; Wld^{SS}$ mouse with advanced hindlimb defect. Related to Figure 1.

Movie 4. 8-month-old female $Wld^{SS}$ mouse (control). Related to Figure 1.

Movie 5. 8-month-old female $Nmnat^{2gtE/gtE}; Sarm1^{-/-}$ mouse. Related to Figure 1.

Movie 6. 8-month-old female $Sarm1^{-/-}$ mouse (control). Related to Figure 1.
Figure 2

A

```
innervation: full partial denervated

|     | 10 weeks | 10 months | 24 months |
|-----|----------|-----------|-----------|
| WldS/S  | 160      | 147       |           |
| Nmfnat2gtEgtE; WldS/S  | 238      | 167       |           |
| Nmfnat2gtEgtE; Sarm1+/+  | 151      | 138       |           |
| Nmfnat2gtEgtE; Sarm1+/-  | 150      | 174       |           |
```

Endplate occupancy (%)

B

```
Nmfnat2gtEgtE; WldS/S
10 months
mixed innervation
denervated

Nmfnat2gtEgtE; Sarm1+/+ 10 weeks
Nmfnat2gtEgtE; Sarm1+/- 10 months
```

50 μm
Table S1. Mendelian ratios of viable offspring from *Nmnat2^{+gtE};Wld^{S/S}* and *Nmnat2^{gtE/gtE};Sarm1^{-/-}* crosses (updated). Related to Figure 1.

| cross | viable offspring (>P10) | Genotype | Obs. | Exp. |
|-------|--------------------------|----------|------|------|
|       |                          | *Wld^{S/S}*           | 67   | 68.5 |
| *Nmnat2^{+gtE};Wld^{S/S}* x *Nmnat2^{gtE/gtE};Wld^{S/S}* | *Nmnat2^{+gtE};Wld^{S/S}* | 144   | 137  |
|       |                          | *Nmnat2^{gtE/gtE};Wld^{S/S}* | 63   | 68.5 |

$\chi^2$ test $p = 0.660$

| cross | viable offspring (>P10) | Genotype | Obs. | Exp. |
|-------|--------------------------|----------|------|------|
|       |                          | *Sarm1^{-/-}*       | 32   | 31.25 |
| *Nmnat2^{+gtE};Sarm1^{-/-}* x *Nmnat2^{+gtE};Sarm1^{-/-}* | *Nmnat2^{+gtE};Sarm1^{-/-}* | 61   | 62.5 |
|       |                          | *Nmnat2^{gtE/gtE};Sarm1^{-/-}* | 32   | 31.25 |

$\chi^2$ test $p = 0.965$
Figure S1. Muscle atrophy and an absence of weight gain in Nmnat2<sup>gtE/gtE;Wld<sup>S/S</sup></sup> mice but not Nmnat2<sup>gtE/gtE;Sarm1<sup>-/-</sup></sup> mice. Related to Figure 1. (A) RT-PCR analysis of Nmnat2 mRNA levels in brains of 10 to 12-month-old mice of the indicated genotypes. Actb acts as a sample reference. Images are representative of n = 3 mice per genotype. Cycle numbers were adjusted for Nmnat2 detection such that band intensity is close to saturation for the Wld<sup>S/S</sup> and Sarm1<sup>-/-</sup> control samples to emphasize the relative absence of Nmnat2 mRNA in the Nmnat2<sup>gtE/gtE;Wld<sup>S/S</sup></sup> and Nmnat2<sup>gtE/gtE;Sarm1<sup>-/-</sup></sup> samples. (B) Widespread hindlimb muscle wasting in Nmnat2<sup>gtE/gtE;Wld<sup>S/S</sup></sup> mice relative to matched Wld<sup>S/S</sup> controls. Representative 6-month-old female mice are shown but the phenotype invariantly develops from 3-5 months onwards in both male and female Nmnat2<sup>gtE/gtE;Wld<sup>S/S</sup></sup> mice. (C and D) Body weights of male and female Wld<sup>S/S</sup> and Nmnat2<sup>gtE/gtE;Wld<sup>S/S</sup></sup> mice (C) and Sarm1<sup>-/-</sup> and Nmnat2<sup>gtE/gtE;Sarm1<sup>-/-</sup></sup> mice (D) at the indicated ages. Individual values with means ±SEM are plotted (n.s. not significant [p > 0.05] or ***p <0.001 in two-way ANOVA with Bonferroni’s multiple comparisons, separate tests for males and females).
Figure S2. Motor unit numbers are reduced in 10 month \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} FDB muscles but remaining units function normally. Related to Figure 2. (A) Representative maximum tetanic tension response (50 Hz stimulation for 1 s) of FDB muscle preparations from 10-month-old male \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} and \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Sarm1}\textsuperscript{-/-} mice. (B) Maximum tetanic force of FDB muscles from 10-month-old male mice of the genotypes listed. Data are mean ±SEM (n = 6 muscles from three mice, except for n = 4 muscles from two \textit{Sarm1}\textsuperscript{-/-} mice). \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} force is noticeably reduced compared to the other genotypes due to reduced numbers of motor units (Fig. 3E), although substantial intra-genotype variation means that group sizes are not sufficient for statistical significance (one-way ANOVA, p > 0.05). (C) Correlation between (reduced) numbers of motor units in \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} FDB muscle explants and maximum tetanic force. Spearman correlation \(r = 0.986\) (\(p = 0.0028\)). This suggests that, although different numbers of motor units remain in individual 10-month-old \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} FDB muscles, those that are present are similar in size. (D) Representative fatigue profiles of FDB muscle explants of the four genotypes studied (as indicated) elicited by 50 Hz tetanic stimulation for 3 s, at intervals of 5 s (gaps between each tetanus were clipped). (E) Fatigue resistance of FDB muscle explants for 10-month-old mice of the genotypes listed. Individual values with means ±SEM are plotted (same groups sizes as in B). The index of fatigue resistance is the stimulus number that produced a tetanic force equal to 50% of the initial force. Fatigueability of \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} and \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Sarm1}\textsuperscript{-/-} FDB muscles is not significantly different from matched \textit{Wld}\textsuperscript{SS} or \textit{Sarm1}\textsuperscript{-/-} controls (one-way ANOVA, \(p > 0.05\)). However, grouping muscles based on \textit{Wld}\textsuperscript{SS} or \textit{Sarm1}\textsuperscript{-/-} status (i.e. excluding \textit{Nmnat2} status) suggests genetic background does influence fatigueability (\(t\) test for \textit{Wld}\textsuperscript{SS} and \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Wld}\textsuperscript{SS} versus \textit{Sarm1}\textsuperscript{-/-} and \textit{Nmnat2}\textsuperscript{gtE/gtE};\textit{Sarm1}\textsuperscript{-/-}, \(p = 0.0126\)).
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Table S1. Mendelian ratios of viable offspring from Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> and Nmnat2<sup>Mε2E</sup>; Sarm1<sup>−/−</sup> crosses (updated). Related to Figure 1.
Birth ratios to show that Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> and Nmnat2<sup>Mε2E</sup>; Sarm1<sup>−/−</sup> are born at the expected frequencies (updated in relation to information in the cited publications).

Figure S1. Muscle atrophy and an absence of weight gain in Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> mice but not Nmnat2<sup>Mε2E</sup>; Sarm1<sup>−/−</sup> mice. Related to Figure 1.
Additional data relating to the muscle atrophy in Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> mice.

Figure S2. Motor unit numbers are reduced in 10 month Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> FDB muscles but remaining units function normally. Related to Figure 2.
Data showing that motor units in Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> FDB muscles, although fewer in number, are of broadly equivalent size and have a fatigueability profile comparable to controls.

Movies 1-6. Movies of representative 8- to 9.5-month-old female Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> (1-3), Wld<sup>SS</sup> (4), Nmnat2<sup>Mε2E</sup>; Sarm1<sup>−/−</sup> (5), and Sarm1<sup>−/−</sup> (6) and mice. Related to Figure 1.
Movies highlighting the impaired hindlimb function in Nmnat2<sup>Mε2E</sup>; Wld<sup>SS</sup> mice but not Nmnat2<sup>Mε2E</sup>; Sarm1<sup>−/−</sup> mice or controls.