Axon death signalling in Wallerian degeneration among species and in disease

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1. Introduction

Neurons use their axons to communicate with remote cells. These axons can be extremely long, ranging from millimetres to centimetres to metres depending on the host, the type of neuron and the target cell [1,2]. In other words, axons can take up more than 99.9% of the neuronal volume. Axons also harbour a remarkably elaborate axonal cytoskeleton (axoskeleton) [3], which allows them to withstand stretch, compression, tension and torsion [4–8]. To ensure continued circuit function, the nervous system established soma-independent, local axonal-intrinsic and glial-extrinsic mechanisms to support lifelong axon survival [9]. If these survival mechanisms are impaired, axons will undergo axon degeneration [10,11]. Axonopathies are increasingly recognized as major contributors in neurological conditions, such as Alzheimer’s disease (AD), amyotrophic lateral sclerosis (ALS) and multiple sclerosis (MS) [12], Parkinson’s disease (PD) [13], traumatic brain injury (TBI) [14], and chemotherapy-induced peripheral neuropathy (CIPN) [15]. Axon degeneration occurs prior to neuronal loss in a broad range of injured and diseased nervous systems, thus targeting it by therapeutics serves as a promising opportunity to ameliorate neurological disorders [16,17]. Gaining insights into underlying mechanisms executing axon degeneration will help to...
define targets for the development of efficacious drugs for therapeutic intervention. However, how axon degeneration is mechanistically regulated and executed remains currently largely unknown.

To date, distinct morphological modes of axon degeneration have been observed and underlying molecular mechanisms described [10]. Among them are dying back axon degeneration, retraction, axosome shedding, focal axonal degeneration induced by growth factor withdrawal and pruning, to name a few. Axon degeneration can also be triggered through axonal injury (axotomy), which is probably one of the simplest models to study how axons execute their own destruction. Identified by and named after Augustus Waller, Wallerian degeneration (WD) is an umbrella term under which two distinct mechanisms occur [18] (figure 1): first, severed axons—separated from the soma—actively execute their own disassembly (axon death) within 1 day, through an evolutionary conserved axon death signalling cascade; and second, surrounding glial cells engage and clear the resulting debris within 3–5 days. Axon death and glial clearance are separate processes (figure 1): the attenuation of axon death results in severed axons which remain functionally and morphologically preserved for weeks to months, while defective glial clearance culminates in axonal debris which persists for a similar time in vivo [19,20].

Axon death signalling is activated not only when the axon is cut, crushed or stretched [21,22], it also seems to be a major contributor in different animal models of neurological conditions, e.g. where axones degenerate in the absence of injury [23]. Therefore, axon death signalling could provide insights into axon degeneration beyond a simple injury model or, as August Waller wrote [18], ‘It is impossible not to anticipate important results from the application of this inquiry to the different nerves of the animal system. But it is particularly with reference to nervous diseases that it will be most desirable to extend these researches’. Over recent years, significant advancements broadened our understanding of axon death signalling in WD. While these discoveries shed light onto distinct mediators of axon death, they also led to more unanswered questions. First, how complete is our understanding of axon death signalling? Second, what differences or commonalities have been observed across species? And third, how do these findings bring us closer to defining new pharmacological targets that could prevent axon degeneration in the diseased and injured nervous system?

In this review, we aim to provide answers to the above questions. First, we will comprehensively summarize our current understanding of axon death signalling across species. Second, we will highlight differences observed across model systems and discuss a potential core mechanism shared across species. And third, we will conclude with a brief overview of engaged axon death signalling in the diseased or injured nervous system.

2. Phases of Wallerian degeneration

Immediately after axotomy, the axon that has been separated from the soma goes through a lag phase where its overall morphology remains unchanged for 6–24 h depending on a number of factors, such as in vitro or in vivo models, and animal systems used (figure 2a). During this phase, the earliest feature observed immediately after injury is a rapid, short-term increase of axonal calcium (Ca^{2+}) levels in both the proximal and distal axon stump (figure 2b). This is mostly due to extracellular Ca^{2+} influx at the lesion site, and to a lesser extent from axon internal Ca^{2+} stores [24–27]. After the first, rapid short-term Ca^{2+} wave, nicotinamide adenine dinucleotide (NAD^+) and adenosine triphosphate (ATP) are rapidly depleted, thus impairing axonal energy homeostasis [28–30]. Mitochondria lose their membrane potential and begin to swell, thereby increasing the generation of reactive oxygen species (ROS). Ultimately, they release their internal Ca^{2+} stores, which culminates in a second, long-term Ca^{2+} wave [28,31–33]. At this point, the gross axonal morphology remains unchanged, despite the already initiated destabilization of microtubules [28,29].

Suddenly, the execution phase starts (figure 2): the axon begins to disassemble. Catastrophic granular fragmentation is observed at the molecular, ultrastructural and morphological level [28]. Microtubules start to disrupt, alongside the dismantling of the axoskeleton. The axon starts ‘beading’ or swelling, culminating in catastrophic axonal fragmentation [30,34,35]. During the execution phase, surrounding glial cells and specialized phagocytes not only clear the resulting axonal debris by activating multiple signalling pathways [19,36–39], but also actively enhance axonal fragmentation [40]. To date,
several key questions remain unresolved, such as what kind of axonal ‘eat me’ signals are presented to surrounding glia [41], and where within the execution phase glial cells engage to clear axonal debris.

Once the execution phase is over, e.g. when the injured axon is disassembled and the resulting debris cleared by surrounding glia, WD is also over. While WD occurs in both the central and peripheral nervous system (CNS and PNS, respectively), it is of particular importance in the PNS to pave the way for the proximal axon still attached to the neuronal soma, which ultimately activates its regenerative programme, to regrow and thereby to re-establish circuit function [33].

3. The discovery and implications of WldS

Axon death within WD was long thought to be a simple passive wasting away of severed axons [18]. In 1989, the serendipitous discovery of the ‘Wallerian degeneration slow’ (WldS) mouse challenged this idea. Severed WldS axons remained preserved for weeks rather than undergoing axon death within a day (figure 2) [42]. Subsequently, the molecular change in WldS mice was identified as a tandem triplication of two neighbouring genes: the N-terminal 70 amino acid fragment of the ubiquitination factor E4B (Ube4b) fused to full length nicotinamide mononucleotide adenylyltransferase 1 (Nmnat1) [43]. This fusion results in a translocation of nuclear Nmnat1 to the axon, where it exerts axon death attenuation (this will be discussed further below). The WldS neomorph provided the first evidence that axon death could be an active self-destruction programme to eliminate damaged axons; that is, because this programme is profoundly attenuated by the sole over-expression of WldS. Following the identification of WldS, several important discoveries were made, which are summarized below.

First, WldS harbours an evolutionary conserved function. While it was first found to delay axon death in mice [42,43], the mouse chimeric protein performs its function equally well in rats [44], fruit flies [19] and zebrafish [33,45]. WldS is therefore capable of delaying axon death across multiple species.

Second, WldS acts autonomously in neurons and is dosage dependent. The WldS mouse harbours a tandem triplication which results in over-expressed WldS [43]. The higher the levels, the better its capability to preserve severed axons [43].

Third, WldS acts locally within axons. While abundant in both nuclei and axons, WldS confers potent attenuation of axon death signalling specifically in axons [46]. Viral transduction of multiple modified versions of WldS into severed axons up to 4 h after axotomy (e.g. early within the lag phase) is sufficient to attenuate axon death signalling [47].

Fourth, WldS is specific for axon death. Its expression blocks axon death and seems to have largely no effect on developmental pruning [48], or other types of neuronal death such as apoptosis [49].

And fifth, WldS is beneficial in many models of neurodegenerative conditions. Since its discovery, the WldS mouse has been crossed into a vast number of different neurological models to assess its protective ability in the injured and diseased nervous system [23]. These findings support the idea that axon death is shared in injury and disease, and, more importantly, that axon degeneration is a major driver in different neurological conditions.

WldS provided the first evidence that, at the molecular level, axon death can be attenuated by over-expression of a single protein. Its discovery and characterization raised two key questions: first, how is WldS able to attenuate an axonal-intrinsic axon death signalling pathway actively executing the degeneration

Figure 2. Phases and events during Wallerian degeneration. (a) Morphological phases of Wallerian degeneration. After axotomy, the axon that is separated from the soma goes through a lag phase (red), where its morphology remains grossly intact. During the execution phase (brown), the axon undergoes catastrophic axon fragmentation (axon death). Surrounding glia engage and clear the resulting axonal debris. The attenuation of axon death signalling results in morphologically and functionally preserved axons and synapses. (b) Molecular and subcellular events during Wallerian degeneration. Axotomy, lag and execution phase (orange, red and brown, respectively). The precise onset and duration of each event depends on the model system used. Attenuated axon death signalling preserves the structure and function of axons and their synapses.
of severed axons; and second, what are the mediators of axon death signalling? The next section will summarize our current knowledge related to these two questions.

4. Axon death mediators

To date, axon death in WD is observed in several species such as mice, rats, flies and fish (figure 3). So far, each animal model has contributed to the discovery of important axon death mediators and also offered distinct insights into axon death signalling. Importantly, the modification of each mediator attenuates axon death comparable to the effects of WldS, e.g. in the range of days to weeks.

4.1. Ca\(^{2+}\)

As mentioned above, a local initial influx of Ca\(^{2+}\) right after axotomy from the site of injury is observed in both invertebrate and vertebrate models in vivo and in vitro [24,27,50,51]. The first, rapid short-term Ca\(^{2+}\) wave precedes the lag phase (figure 2). The disassembly of severed axons is initiated by high extra-axonal Ca\(^{2+}\) concentrations: if the extracellular environment is devoid of Ca\(^{2+}\) (by adding EGTA, or in a medium lacking Ca\(^{2+}\)), or when voltage-gated Ca\(^{2+}\) channels are inhibited, the fragmentation of severed axons is significantly delayed (figure 3a) [26,50,52–54].

The second, long-term Ca\(^{2+}\) wave is present solely in the separated axon. It appears prior to the onset of axonal fragmentation, and is believed to be a key instructive component for the execution phase [51,55]. During the second wave, Ca\(^{2+}\) is released from intra-axonal stores (e.g. mainly from mitochondria, and to a lesser extent from the endoplasmic reticulum), and the inhibition thereof can significantly delay the disassembly of the axon [25].

The expression of Wld\(^{5}\) in zebrafish and rodent neuronal cultures largely suppresses the second, but not the first, Ca\(^{2+}\) wave, suggesting that the second wave is the one responsible for triggering axon fragmentation [24,51]. In Drosophila larvae, Wld\(^{5}\) is also suppressing the first Ca\(^{2+}\) wave, reflecting minor differences between the experimental systems [27].

Taken together, the rise in intra-axonal Ca\(^{2+}\), both during the immediate first short-term as well as during the slower and delayed second long-term Ca\(^{2+}\) wave, is an important instructive signal to trigger axon death signalling. However, how transient high Ca\(^{2+}\) levels are linked to the initiation of axon death signalling remains completely unknown.

4.2. Nmnat2/dNmnat

Wld\(^{5}\) is an over-expressed version of Nmnat1 translocated from the nucleus to the axon [43,56]. Nmnat proteins are essential for ATP-dependent NAD\(^{+}\) synthesis from either...
nicotinamide mononucleotide (NMN) or nicotinic acid mononucleotide (NaMN) [57]. Mammals and zebrafish harbour three different Nmnat genes (Nmnat1–3) with different subcellular locations and kinetic properties [58]. In contrast, Drosophila relies on a single nmnat gene (dNmnat) [38], which is alternatively spliced to generate variants localized to the nucleus and the cytoplasm [59]. The discovery of WldS, and subsequently the extensive research on Nmnat proteins and their biosynthetic NAD\(^+\) activity, provided the following four major insights: levels and localization of Nmnat proteins as well as levels of NAD\(^+\) and its precursor NMN are all crucial for the execution of axon death signalling.

4.2.1. Nmnat2/dNmnat levels

Among the three mammalian Nmnat isoforms, Nmnat2 is the gatekeeper of axon death: it is a limiting, labile axon survival factor [60]. Synthesized in the soma and transported into the axon, Nmnat2 has to be constantly replenished in axons due to its rapid turnover (figure 3): Nmnat2 is the most labile member of its family with a half-life of less than 4 h, despite being the most highly expressed isoform in the brain [61]. Upon axotomy, Nmnat2 fails to be transported to the axon; thus, in severed axons, Nmnat2 levels drop rapidly, and axon death is initiated [60]. Vice versa, depletion of Nmnat2 in whole neurons is sufficient to induce axon degeneration in the absence of injury: axons undergo Wallerian-like degeneration—because they can be attenuated by Wld\(^6\)—while the soma remains unaffected [60,62]. Similarly, Nmnat2\(^{−/−}\) mice harbour perinatal lethality due to limited axon extensions, and their isolated neuronal cultures from both PNS and CNS contain neurite outgrowth consistently stalling at 1–2 mm [63]. Likewise, in Drosophila, RNAi-mediated knockdown, or a lack of dNmnat in mutants, leads to spontaneous axon degeneration prior to cell body degeneration [64,65]. In zebrafish, over-expression of either Wld\(^6\) or Nmnat2 results in a potent attenuation of axon death in vitro [33,45]. Whether knock-down of Nmnat2 also results in Wallerian-like degeneration remains to be determined. Taken together, levels of mammalian, fish and fly Nmnat2/dNmnat are crucial for axonal survival (figure 3). It is important to note that any engineered version of stabilized Nmnat protein is capable of attenuating axon death in a variety of different models, e.g. Wld\(^6\) as non-nuclear Nmnat1, Nmnat1 that fails to localize to nuclei (e.g. cytosolic and axonal Nmnat1, cyt-Nmnat1 and ax-Nmnat1, respectively), Nmnat2 able to persist longer in axons, Drosophila dNmnat over-expressed in cultured mouse dorsal root ganglia (DRG) or mouse Nmnat1 in Drosophila olfactory receptor neurons (ORNs) [45,61,62,66–68]. Therefore, above-threshold levels of Nmnat2 or modified versions thereof ensure the survival of the axon, while sub-threshold levels trigger axon death.

4.2.2. Nmnat2/dNmnat localization

Nmnat2 is predominantly found in cytoplasm and axoplasm, and is associated with membranes of Golgi-derived transport vesicles undergoing fast axonal transport [63]. Both the removal of the vesicle-association domain in Nmnat2 [62] or the modification of its residues required for palmitoylation [69] lead to increased Nmnat2 half-lives. Moreover, in mammals or flies, the over-expression of Nmnat3, which is predominantly found in mitochondria in mammals, potently attenuates axon death [67,70]. Therefore, the subcellular localization dictates turnover rates of Nmnat proteins. By targeting them to different axonal compartments relative to endogenously localized Nmnat2, their turnover is reduced, which results in increased Nmnat protein half-lives, and ultimately in attenuated axon death.

4.2.3. Nmnat substrate: nicotinamide mononucleotide (NMN)

The attenuation of axon death signalling requires NAD\(^+\) biosynthetic activity of Nmnat proteins [33,61,67,71,72]. Metabolic aspects of NAD\(^+\) in neurodegeneration and axon death signalling have recently been comprehensively reviewed in [57]. Importantly, genetic and pharmacological modifications of NAD\(^+\) metabolism result in different levels of attenuation [73]. Here, we will focus on manipulations that result in a robust long-term (days to weeks) or intermediate (2–3 days) attenuation of axon death signalling, respectively. However, we do note that there are manipulations which result in a short-term attenuation, e.g. a delay of axon death signalling in the range of a day. These observations remain subject to further analyses and will not be covered here.

Nmnat proteins use either NMN or NaMN to generate NAD\(^+\) in an ATP-dependent manner. After injury, Nmnat protein levels drop, resulting in increased Nmnat substrates, e.g. NaMN and/or NMN, and decreased Nmnat products, e.g. NAD\(^+\). But which one is important for axon death signalling? While levels of NaMN seem to be negligible [73], levels of NMN are crucial: they temporally rise within 6 h in severed axons and seem to be an instructive signal [55,74]. Preventing this NMN rise results in long-term attenuation of axon death: either by the expression of Wld\(^6\) or modified Nmnat proteins that consume NMN to generate NAD\(^+\) or—interestingly—by the expression of a bacterial specific enzyme, NMN deamidase, which consumes NMN but does not generate NAD\(^+\), both in vitro [55,73,74] and in vivo in zebrafish and mice [75].

There is an alternative modification to keep NMN levels low: nicotinamide (Nam) is consumed by nicotinamide phosphoribosyltransferase (NAMPT) to generate NMN. The pharmacological inhibition of NAMPT by FK866, either right before or immediately after injury, prevents the rise of NMN, which results in short-term to intermediate attenuation of axon death signalling [73,74]. In summary, distinct pharmacological or genetic manipulations that lead to low levels of NMN also prevent axon death. Thus, the temporal rise of NMN after injury seems to be an instructive signal for axons to execute their own destruction.

4.2.4. Nmnat product: nicotinamide adenine dinucleotide (NAD\(^{+}\))

The involvement of NAD\(^+\) as a product of the biosynthetic activity of Nmnat proteins in axon death was observed right after the discovery of Wld\(^6\), before or immediately after axonal injury, the exogenous supply of high levels of NAD\(^+\) leads to long-range attenuation of axon death signalling [71,72]. Thus, axon death is triggered by NAD\(^+\) depletion in severed axons. It is important to note that Wld\(^6\) or engineered Nmnat proteins do not generate higher levels of NAD\(^+\), they solely prevent the depletion thereof by a—yet unknown—mechanism that inhibits an NAD\(^+\) consuming enzyme, which will be discussed further below [73,76].

Wld\(^6\) got its foot in the door of axon death signalling and paved the way for several key discoveries related to Nmnat proteins. Levels and localization of the axonal survival
factor Nmnat2/dNmnat are crucial for the survival or the degeneration of the axon. Its biosynthetic activity keeps levels of NMN as substrate low, and prevents the depletion of NAD⁺ as product (figure 3a).

Given the essential role of Nmnat proteins in axon death, an important question is how their turnover and half-life is regulated. What are the mechanisms regulating protein levels and therefore biosynthetic activity? Below, we will discuss the mechanisms that are crucial for Nmnat2 protein levels.

4.3. Atypical ubiquitin ligase complex

Following the cloning of Wld⁺ [43], the ubiquitin proteasome system (UPS) was also found to be involved in the early stages of axon death [29]. The subsequent discovery of Nmnat2 as a labile axon survival factor, which is subjected to rapid turnover in axons [60], led to an attractive hypothesis: could the UPS be responsible for the rapid turnover of Nmnat2 in axons? The first evidence for the UPS system to be involved in axon death was found in Drosophila. The E3 ubiquitin ligase Highwire (Hiw) regulates the turnover of dNmnat [20,77]. Likewise, the mammalian homologue PAM/Highwire/RPM-1 (Phr1) also fine-tunes levels of Nmnat2 [78]. Phr1 belongs to an evolutionarily conserved, atypical Skp/Cullin/F-box (SCF)-type E3 ubiquitin ligase complex consisting of S-phase kinase-associated protein 1A (Skp1A), Phr1 and F-box protein 45 (Fbxo45) [78,79]. This atypical SCF complex regulates—through polyubiquitination and the proteasome—levels of Nmnat2, by specifically targeting axoplasmic Nmnat2 for destruction [69,80]. The removal of either component slows down the turnover of Nmnat2, which results in attenuated axon death signalling (figure 3a).

4.4. MAPK signalling

The mitogen-activated protein kinase (MAPK) signalling pathway is also involved in axon death signalling. It is activated within 5 min after axonal injury and culminates in the phosphorylation of c-Jun N-terminal kinases (JNKs): loss-of-function analyses revealed that a partially redundant MAPK cascade is required to execute the degeneration of axons after injury (e.g. MKK4, MLK, DLK, MKK4/7, JNK1/3) [84]. The MAPK cascade limits levels of Nmnat2, by selectively degrading membrane-associated, palmitoylated Nmnat2 [69,82].

Interestingly, Nmnat2 levels are differentially regulated. Both MAPK signalling and the atypical ubiquitin ligase complex are important for the fine-tuning of Nmnat2 levels. While MAPK signalling targets membrane-associated Nmnat2, the atypical ubiquitin ligase complex selectively degrades axoplasmic Nmnat2 [69,85]. The pharmacological inhibition of both mechanisms results in a strong attenuation of axon death signalling [69]. This suggests that distinct axonal pools of Nmnat2 are differentially regulated (figure 3a).

The above discoveries revealed a central and conserved function for Nmnat2/dNmnat. Levels of the labile axonal survival factor are dictated by three branches (figure 3a): (i) continuous supply of Nmnat proteins by axonal transport, (ii) constant degradation of vesicle-bound Nmnat proteins by the MAPK pathway, and (iii) constant degradation of soluble Nmnat proteins by the atypical ubiquitin ligase complex.

In healthy uninjured axons, Nmnat protein levels are above threshold, ensuring low NMN and high NAD⁺ levels. Vice versa, in injured axons, the supply of Nmnat proteins by axonal transport is cut down, and, therefore, Nmnat protein degradation by the MAPK pathway and the atypical ubiquitin ligase complex takes over. This results in below-threshold levels of Nmnat proteins. Likewise, levels of NMN temporally rise, whereas NAD⁺ levels drop.

So far, the execution of axon death signalling is solely initiated by below-threshold levels of Nmnat proteins. Any modification that sustains levels of Nmnat proteins—e.g. gain of Nmnat stability or loss of Nmnat protein degradation—ultimately attenuates axon death.

At this point, no mediator in axon death signalling had been identified that actively contributes to signalling, e.g. where loss-of-function mutations result in attenuated axon death signalling regardless of Nmnat protein levels. Below, we will discuss such mediators.

4.5. Sarm1/dSarm

The first discovery of a mutation which attenuates axon death irrespective of Nmnat levels was made in Drosophila: through an unbiased forward-genetic screen for axon death defective mutants, several loss-of-function alleles of the gene ‘Drosophila sterile alpha and armadillo motif’ (dsarm) were isolated (figure 3). Mutations in dsarm block axon death for the lifespan of the fly [86], dsarm is therefore essential for injury-induced axon degeneration. Similarly, mutants or downregulation of the mammalian homologue Sarm1 harbour a potent attenuation of axon death signalling in vitro and in vivo [76,86–88]. Sarm1/dSarm is a toll-like receptor adaptor family member and mainly expressed in the nervous system [86], yet it also functions in glial cells [89], and in the immune system [90].

Sarm1/dSarm contains three evolutionarily conserved protein domains: an Armadillo/HEAT (ARM) domain, a sterile alpha motif (SAM) (two in mammals and one in flies) and a Toll/interleukin-1 receptor homology (TIR) domain [86,87]. All three domains are essential for Sarm1 function in mice [76,82,87] and dsarm in Drosophila [65]. The ARM domain keeps dsarm/Sarm1 inactive, as previously reported in the C. elegans homologue TIR-1 [91], and the SAM domain is important for Sarm1 dimerization [76,81]. The TIR domain, rather than harbouring signalling activity, contains an enzymatic activity to consume NAD⁺ (NADase activity) [92]. Sarm1 activation by dimerization is necessary and sufficient locally within axons to execute axon degeneration in the absence of injury. Its activation triggers rapid depletion of NAD⁺ [73,76,92]. Importantly, the TIR domain NADase activity is evolutionarily conserved across flies, zebrafish and mice, where it cleaves NAD⁺ into nicotinamide (Nam) and ADP-ribose (ADPR) or cyclic ADPR (cADPR) with species-specific differences [92]. These findings support a model where, upon injury, Sarm1/dSarm is activated and actively depletes NAD⁺ levels in severed axons. Therefore, Sarm1/dSarm plays a central role in NAD⁺ depletion in severed axons after injury.

4.6. Axundead

Recently, another essential mediator of axon death signalling has been identified in Drosophila. Several axundead (axed) mutants were isolated by another unbiased forward-genetic screen [65]. Similar to highwire and dsarm mutants, mutations in axed attenuate axon death signalling for the lifespan of the...
fly (figure 3b) [20,65,86]. The axed gene consists of two isoforms (axed\textsuperscript{short} and axed\textsuperscript{long}), and both isoforms are capable of rescuing axed mutants. Axed proteins are predominantly found in axons and synapses, and its levels increase 4–6 h post axotomy and return to baseline 24 h after injury, suggesting either a transient change in localization and/or levels in response to axonal injury. Axed contains two evolutionary conserved domains: a BTB and a BACK domain, which suggests that Axed could dimerize and/or interact with cullin ubiquitin ligases [93]. There are four mammalian paralogues (BTBD1, BTBD2, BTBD3, BTBD6), and it remains to be determined which parologue(s) could have an active role in axon death in mammals.

4.7. Calpains and Calpastatin

Calpains have recently also been shown to be involved in axon death signalling (figure 3c) [81]. Calpains are Ca\textsuperscript{2+}-dependent non-lysosomal proteases, and they are involved in neuronal degeneration in traumatic brain injury, cerebral ischaemia and AD [94]. Among the 15 mammalian isoforms, Calpain-1 and Calpain-2 are ubiquitously expressed, predominantly in the brain, and are present in both neurons and glia [95]. Within minutes after experimental traumatic axonal injury (TAI), axonal Calpain activity is elevated [96]. Vice versa, mice lacking Calpain-1/2 show a significant delay in axon death compared with wild-type mice [81].

Calpastatin is an endogenous in vivo inhibitor of Calpains. After axonal injury, levels of Calpastatin drop within 10 h, which correlates with the morphological degeneration of axons [34]. In contrast, transgenic mice expressing human Calpastatin harbour robust attenuation of axon death in transected optic nerves in the CNS and sciatic nerves in the PNS [97].

Calpastatin-mediated regulation of Calpains is likely the most downstream cascade of axon death signalling. However, the mechanistic link between the above mediators of axon death signalling and Calpains in mammals, and whether Calpains are also involved in other species, remains to be determined.

4.8. Other mediators

So far, axon death mediators were discussed whose modification results in a robust attenuation. Yet also other modifications have been reported that are capable of delaying axon death, such as the mitochondrial permeability transition [31], recycling endosomes [98], autophagy [99,100], sodium and potassium currents [50], the ubiquitin ligase ZNRF-1 [101], microtubule destabilization mediated by CRMP2 [102,103] and the transcription factor Pebbled/RREB1 [104]. The precise interaction of these mediators with the signalling cascade remains to be determined, because of either tissue-specific phenotypes or the involvement of whole organelles.

5. Interspecies commonalities and differences

Wild\textsuperscript{S} attenuates axon death in a variety of models across evolution [19,32,42–44,48]. Each animal system offers unique insights into axon death signalling, thus also distinct observations have been made between them. How is axon death studied in different animal models, and what is common, or different, among them? Below, we will briefly discuss each animal model where axon death is extensively studied, followed by highlighting commonalities and differences in axon death signalling across species.

5.1. Animal models

Mice are frequently used for both in vitro and in vivo axon death assays: in vitro cultured neurons from superior cell ganglia (SCGs) and dorsal root ganglia (DRGs) from the PNS, and retinal ganglion cells (RGCs) from the CNS, are subjected to axon death assays. Moreover, pharmacogenetic manipulation of metabolism can also be applied in vitro. Broadly used in vivo assays are optic nerve injuries (CNS) and sciatic nerve lesions (PNS).

Rats facilitate stereotaxic injections because of their larger brains, and lesions of optic or facial nerves and nerve roots can also be readily performed. Moreover, nerves are longer than those in mice, putting them closer to human axons, although human axons can still be 10-fold longer than their rat counterparts [24,44]. Rats also provide more abundant sources of tissue for biochemical and proteomic studies.

Zebrafish provides a major advantage with its powerful in vivo live imaging, as the complete time course of axon death, with the resolution of single axons, can readily be observed. It is important to mention that zebrafish offers the unique ability to visualize temporal Ca\textsuperscript{2+} events in vivo.

Flies harbour the unique advantage of unbiased forward-genetic screens. Axon death is observed in vivo by antennal or maxillary palp ablation, wing injuries, and during larval development by nerve crush. These techniques were recently reviewed in [105]. Moreover, by the use of optogenetics, the functional preservation of axons and their synapses can readily be assessed by a simple grooming assay.

5.2. Commonalities and differences across species

Genetic analyses in the above animal systems provided important insights over recent years. These findings led to the definition of an axon death signalling pathway. It is tempting to combine all analyses to define a core signalling cascade across multiple species. Yet these analyses also revealed subtle differences that should not be neglected. Below, we will highlight common features as well as species-specific axon death signalling differences among mouse, fly and fish.

5.2.1. Conserved mediators

Ca\textsuperscript{2+} plays a crucial role in all model systems tested so far. Both Ca\textsuperscript{2+} entry into the axon after injury (first, rapid short-term Ca\textsuperscript{2+} wave) and the Ca\textsuperscript{2+} release from intracellular stores at the end of the lag phase (second, long-term Ca\textsuperscript{2+} wave) have been observed across multiple species. It is therefore likely that the first conserved mediator of axon death signalling is external Ca\textsuperscript{2+} influx into severed axons. Nmnat2/dNmnat in mice, fish and flies harbour an evolutionary conserved feature too. As a labile axon survival factor, levels of Nmnat proteins matter: high or robust levels potently protect severed axons from undergoing axon death, and low levels induce axon degeneration in all species tested [33,45,60,64]. Alongside Nmnat, its biosynthetic NAD\textsuperscript{+} activity is also conserved across species. It remains to be
determined whether levels of NMN and NAD⁺, or ratios thereof, define when axon death is initiated, not only in mice but also in flies and fish. Atypical ubiquitin ligase complex members, which regulate Nmnat protein levels, are conserved in mice and flies (Skp1a, Phr1, Fbxo45 in mouse, and Phr1 in fly). Interestingly, while the MAPK pathway plays an important role in mice, it seems to be negligible in flies [65]. Whether the fish homologues of the atypical ubiquitin ligase complex and the MAPK signalling cascade regulate levels of Nmnat2 in a similar way remains elusive.

The function of the Dsarm/Sarm1 homologues in flies and mice, and Sarm1 in fish [106], is also evolutionarily conserved. Loss-of-function mutants block axon death signalling, and the TIR domain of these species harbours the ability to pathologically degrade NAD⁺ to generate Nam and ADPR [92].

It is equally important to note that species-specific pathway analyses revealed some remarkable differences. This indicates that we are far from fully understanding axon death signalling mechanisms. Below, we will briefly discuss these differences.

5.2.2. Differences across species

Mice lacking Nmnat2 contain truncated axons during embryogenesis and die perinatally [63]. This lethality is partially rescued by over-expression of Wld⁺ [107], and fully rescued by Sarm1 mutants [88]. Therefore, Sarm1 executes axon death following below-threshold levels of Nmnat2. Sarm1 likely acts downstream of Nmnat2 (figure 3a). However, it could also act in parallel to it, but certainly not upstream of Nmnat2.

As soon as Nmnat2 levels drop below threshold, Sarm1 unleashes its TIR domain to consume NAD⁺. Nmnat2, rather than maintaining NAD⁺ levels through its NAD⁺ biosynthetic activity, blocks NAD⁺ consumption of Sarm1, which seems central to axon death signalling [73,76]. It remains completely unknown how Nmnat2 inhibits the NADase activity of Sarm1.

As mentioned above, Nmnat2−/− mice contain truncated axons during embryogenesis and die perinatally [63]. Wld⁺ mice and Sarm1 mutants are not the only two candidates able to rescue both: surprisingly, the expression of the bacterial NMN deamidase does the same in a dosage-dependent manner [75]. This finding suggests that, besides the pathologically degraded NAD⁺ consumption of Sarm1, the temporal rise of NMN and lower axonal concentrations of NMN and higher concentrations of NAD⁺ [73,74,88] (figure 3a). Injury leads to below-threshold levels of the labile axonal survival factor Nmnat2, which in turn leads to a temporal rise of NLMN and lower levels of NAD⁺, thus reducing the ratio (NMN < NAD⁺). This ratio could already be sufficient to activate axon death. Importantly, below-threshold levels of Nmnat2 also discontinue Nmnat2-mediated inhibition of Sarm1, which discharges the NADase activity of Sarm1: the resulting Sarm1-mediated NAD⁺ consumption tips the ratio towards NMN (NMN ≫ NAD⁺). This ratio could therefore dictate whether axons should survive, or degenerate. Axon survival is favoured by NMN (NMN ≪ NAD⁺) preventing the rise of NMN, or the consumption of NAD⁺, both potently attenuate axon death. Vice versa, conditions altering this ratio will induce axon death: the forced depletion of NAD⁺ by a constitutively active NADase activity of Sarm1/Dsarm is sufficient to trigger axon (and cell body) death. If this were true, high NMN levels—significantly higher than NAD⁺—should also be capable of triggering axon death in vivo. This remains to be tested, but also harbours technical difficulties: Nmnat proteins efficiently synthesize NAD⁺ from NMN, thus a rise in NMN automatically results in a rise of NAD⁺ [58,73]. Interestingly, high levels of a cell-permeable analogue of NMN (which cannot be used as substrate for Nmnat-mediated NAD⁺ synthesis) are sufficient to activate Sarm1 in cultured neurons, thereby depleting NAD⁺, which results in non-apoptotic cell death. Whether this is also true in severed axons remains to be determined [108].

We are still far from understanding why and how axons execute their own disassembly. Our knowledge of axon death as an emerging signalling cascade is still in its infancy. Each axon death mediator revealed crucial mechanistic insights over recent years, and helped to define Sarm1 as a first target that can be translated to the clinic. In the section below, we will briefly discuss where targeting of Sarm1 is beneficial in mouse models of neurological conditions.

6. Axon death signalling in disease

Mutations in human Nmnat2 have been linked to fetal akinesia deformation sequence (FADS), and to childhood-onset polyneuropathy and erythromelalgia [109,110]. These discoveries provide the first direct molecular evidence that axon death in WD is involved in a human axonal disorder. They support
the idea that axonopathies are a major contributor to certain human neurodegenerative disorders [12–15].

WldS and its capacity to slow down injury-induced axon degeneration offered the first opportunity to attenuate axonopathies which occur in the absence of injury. Over recent years, distinct outcomes were observed in a broad range of disease models, including extended lifespan, age-dependent effects, improved performance, or a lack of amelioration [23]. The promising neuroprotective function of WldS also led to efforts developing drugs to stabilize Nmnat proteins [111]. However, while the efficacy of WldS or any other form of engineered Nmnat protein seems promising in certain models of neurological conditions, therapeutic potential may be limited due to a gain-of-function protein: its efficacy relies on dosage, which could be challenging during long-term therapeutic treatments [43]. Moreover, WldS harbours age-dependent effects [112], and while it is capable of slowing down axon death, it is not able to fully block it [107]. Therefore, alternative or complementary approaches would increase the chance of ameliorating axonopathies.

The manipulation of the NMN/NAD+ ratio could offer an attractive alternative. It could be achieved by pharmacological targeting of specific metabolic pathways, or by simple oral supplement of NAD+ or its precursors. Preventing either NAD+ consumption or a temporal rise of NMN could serve as attractive strategies [113,114]. However, without any specificity towards neuronal tissue, the manipulation of NAD+ metabolism also bears great risks [115].

The discovery of Sarm1/dSarm has offered a novel and unique therapeutic opportunity. In order to block axon death signalling, Sarm1 protein or its activity has to be decreased, rather than increased such as with Nmnat2. Like WldS, Sarm1 mutant mice offer a range of outcomes in mouse models of neurological conditions:

When axons are challenged by mechanical forces, Sarm1−/− mice harbour reduced neurological deficits and a better functional outcome in a model of TBI [116]. Sarm1−/− mice also contain significantly reduced axonal lesions in a model of TAI with impact acceleration of heads [117]. In addition, an in vivo gene therapy approach using adeno-associated virus to deliver a dominant-negative version of Sarm1 had similar effects in an injury-induced axon degeneration model to that in Sarm1−/− mice [118]. These findings strongly suggest that Sarm1 serves as a promising therapeutic target to ameliorate force-induced axonopathies.

In two models of CIPN (e.g. vincristine or paclitaxel), and in a model of metabolic-induced peripheral neuropathy, mice lacking Sarm1 prevent the distal degeneration of myelinated axons and electrophysiological abnormalities [119,120]. These findings fuel hope that axonopathies caused by chemotherapy or by diabetes—the most common causes of peripheral neuropathies [121,122]—could also be therapeutically targeted in patients.

However, while Sarm1 harbours the potential to ameliorate certain axonopathies, it is important to note that the lack of Sarm1 does not suppress motor neuron degeneration in a mouse model of ALS [123]. It remains to be determined whether Sarm1 is dispensable in other ALS mouse models.

So far, Sarm1−/− mutants have beneficial effects in certain models of neurological conditions. This raises hope that Sarm1 and other axon death mediators could serve as druggable targets to halt axon loss. Axon death signalling serves therefore as an attractive pathway to develop therapies against, with the ultimate goal to prevent and treat axon loss in a broad range of neurological diseases.

7. Conclusion

The discovery and characterization of axon death signalling, which is activated by injury, not only provided exciting insights into the underlying mechanism mediating axonal self-destruction after injury. It also revealed that the axon death pathway is hijacked in other challenging conditions for the nervous system where axonopathies occur in the absence of injury. Therefore, it is crucial to fill missing mechanistic gaps among already identified axon death mediators, and to identify other key mediators required for axon death signalling. This will help to expand our understanding of a signalling pathway that ultimately leads to the death of the axon. Overall, there is certainly a lot left to do, and axon death research will therefore continue to be a lively field in the future.

Data accessibility. This article has no additional data.

Competing interests. The authors declare that they have no competing interests.

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References

1. Matsuda W, Furuta T, Nakamura KC, Hioki H, Fujiyama F, Anai R, Kaneko T. 2009 Single nigrostriatal dopaminergic neurons form widely spread and highly dense axonal arborizations in the neostriatum. J. Neurosci. 29, 444–453. (doi:10.1523/JNEUROSCI.4029-08.2009)

2. Wedel MJ. 2012 A monument of inefficiency: the presumed course of the recurrent laryngeal nerve in sauropod dinosaurs. Acta Palaeontol. Pol. 57, 251–256. (doi:10.4320/app.2011.0019)

3. Xu K, Zhong G, Zhuang X. 2013 Actin, spectrin, and associated proteins form a periodic cytoskeletal structure in axons. Science 339, 452–456. (doi:10.1126/science.1232251)

4. Smith DH, Wolf JA, Meaney DF. 2001 A new strategy to produce sustained growth of central nervous system axons: continuous mechanical tension. Tissue Eng. 7, 131–139. (doi:10.1089/107632701300062714)

5. Hammersland M, Jorgensen EM, Bastiani MJ. 2007 Axons break in animals lacking beta-spectrin. J. Cell Biol. 176, 269–275. (doi:10.1083/jcb.200611117)

6. Smith DH. 2009 Stretch growth of integrated axon tracts: extremes and exploitations. Prog. Neurobiol. 89, 231–239. (doi:10.1016/J.PNEUROBIO.2009.07.006)

7. Fournier AJ, Hogan JD, Rajbhandari L, Shrestha S, Venkatesan A, Ramesh KT. 2015 Changes in neurofilament and microtubule distribution following focal axon compression. PLoS ONE 10, e0131617. (doi:10.1371/journal.pone.0131617)

8. Krieg M, Stühmer J, Cueva JG, Fetter R, Spilker K, Cremers D, Shen K, Dunn AR, Goodman MB. 2017 Genetic defects in β-spectrin and tau sensitize C. elegans axons to movement-induced damage via torque-tension coupling. Elife 6, e20172. (doi:10.7554/eLife.20172)

9. Mariano V, Dominguez-Iturza N, Neukomm LJ, Bagni C. 2018 Maintenance mechanisms of circuit...
integrated axons. *Curr. Opin. Neurobiol.* **33**, 162–173. (doi:10.1016/j.conb.2018.08.007)

10. Neukomm LJ, Freeman MR. 2014 Diverse cellular and molecular modes of axon degeneration. *Trends Cell Biol.* **24**, 515–523. (doi:10.1016/j.tcb.2014.04.003)

11. Burgess RW, Crish SD. 2018 Editorial: Axonopathy in neurodegenerative disease. *Front. Neurosci.* **12**, 769. (doi:10.3389/fnins.2018.00769)

12. Vickers JC et al. 2009 Axonopathy and cytoskeletal disruption in degenerative diseases of the central nervous system. *Brain Res. Bull.* **80**, 217–223. (doi:10.1016/j.brainresbull.2009.08.004)

13. Burke RE, O'Malley K. 2013 Axon degeneration in Parkinson's disease. *Exp. Neurol.* **246**, 72–83. (doi:10.1016/j.expneurol.2012.01.011)

14. Hill CS, Coleman MP, Menon DK. 2016 Traumatic axonal injury: mechanisms and translational opportunities. *Trends Neurosci.* **39**, 311–324. (doi:10.1016/j.tins.2016.03.002)

15. Flatters SJL, Dougherty PM, Colvin LA. 2017 Clinical and preclinical perspectives on Chemotherapy-Induced Peripheral Neuropathy (CIPN). *a* narrative review. *Br. J. Anaesth.* **119**, 737–749. (doi:10.1093/bja/aex229)

16. Coleman MP, Perry VH. 2002 Axon pathology in Wallerian degeneration. *Neuron* **30**, 131–134. (doi:10.1016/S0896-6273(02)00245-X)

17. Park YJ, Jiang SY, Shin YK, Koh H, Suh DJ, Shinni T, Araki T, Park HT. 2013 Mitochondrial swelling and microtubule depolymerization are associated with energy depletion in axon degeneration. *Neuroscience* **238**, 258–269. (doi:10.1016/j.neuroscience.2013.02.033)

18. Waller A. 1850 Experiments on the section of the glossopharyngeal and hypoglossal nerves of the frog, and observations of the alterations produced thereby in the structure of their primitive fibres. *Phil. Trans. R. Soc.* **140**, 423–429.

19. MacDonald JM, Beach MG, Portiglia E, Sheehan AE, Watts RJ, Freeman MR. 2006 The Drosophila cell corpse engulfment receptor Draper mediates glial corpse engulfment but its exposure can be hindered regeneration in peripheral nerve. *J. Biol. Chem.* **281**, 36989–36999. (doi:10.1074/jbc.M609786200)

20. Bridge PM, Ball DJ, Mackinnon SE, Nakao Y, Brandt K, Hunter DA, Hertl C. 1994 Nerve crush injuries—a model for axonotmesis. *Exp. Neurol.* **127**, 284–290. (doi:10.1006/EXNR.1994.1104)

21. Maxwell WL, Bartlett E, Morgan H. 2015 Wallerian degeneration in the optic nerve stretch-injury model of traumatic brain injury: a stereological analysis. *J. Neurotrauma* **32**, 780–790. (doi:10.1089/neu.2014.3369)

22. Conforti L, Gilley J, Coleman MP. 2014 Wallerian degeneration: an emerging axon death pathway linking injury and disease. *Nat. Rev. Neurosci.* **15**, 394–409. (doi:10.1038/nrn3680)

23. Adalbert R, Moreale G, Paisz M, Conforti L, Walker SA, Roderick HL, Bootman MD, Slikos L, Coleman MP. 2012 Intra-axonal calcium changes after axotomy in wild-type and slow Wallerian degeneration axons. *Neuroscience* **225**, 44–54. (doi:10.1016/j.neuroscience.2012.08.056)

24. Villegas R, Martinez NW, Litto J, Phan P, Hernandez D, Twiss JL, Court FA. 2014 Calcium release from intra-axonal endoplasmic reticulum leads to axon degeneration through mitochondrial dysfunction. *J. Neurosci.* **34**, 7179–7189. (doi:10.1523/JNEUROSCI.4784-13.2014)

25. George B, Glass D, Griffin W. 1995 Axotomy-induced axonal degeneration influx through ion-specific channels. *J. Neurosci.* **15**, 6445–6452. (doi:10.1523/jneurosci.15.15.06-1945.1995)

26. Shacham-Silverman V, Sar Shalom H, Goldner R, Golan-Vaishenker Y, Gurwicz N, Gokhman I, Yaron A. 2018 Phosphatidylinositol4-sodium monomethylphosphonate is a marker for axonal debris engulfment but its exposure can be decoupled from degeneration. *Cell Death Dis.* **9**, 1116. (doi:10.1038/s41419-018-1155-x)

27. Lu-T-Y, MacDonald JM, Neukomm LJ, Sheehan AE, Bradshaw R, Logan MA, Freeman MR. 2017 Axon degeneration induces glial responses through Draper-TRAF4-JNK signalling. *Nat. Commun.* **8**, 14355. (doi:10.1038/ncomms14355)

28. Perez VH, Brown MC, Rosen H, Gordon S. 1987 The macrophage response to central and peripheral nerve injury. A possible role for macrophages in Wallerian degeneration. *J. Exp. Med.* **165**, 1218–1223. (doi:10.1084/jem.165.4.1218)

29. Purice MD, Speese SD, Logan MA. 2016 Delayed glial clearance of degenerating axons in aged Drosophila is due to reduced PI3 K/Draper activity. *Nat. Commun.* **7**, 12871. (doi:10.1038/ncomms12871)

30. Hoopfer ED, McLaughlin T, Watts RJ, Schuldiner O, Dey A. 2006 WldS-mediated axon protection. *Neuron* **49**, 367–378. (doi:10.1016/j.neuron.2006.05.023)

31. Barrientos SA, Martinez NW, Yoo S, Jara JS, Zamorano S, Hetz C, Twiss JL, Alvarez J, Court FA. 2011 Axonal degeneration is mediated by the corpse engulfment receptor Draper mediates glial corpse engulfment but its exposure can be hindered regeneration in peripheral nerve. *J. Neurosci.* **31**, 966–978. (doi:10.1523/JNEUROSCI.4065-10.2011)

32. Bernardi P, Krauskopf A, Basso E, Petrocelli V, Blalch-Dyson E, Di Lisa F, Forte MA. 2006 The mitochondrial permeability transition pore. *J. Neurosci.* **26**, 83–86. (doi:10.1523/JNEUROSCI.0248-05.2006)

33. Martin SM, O'Brien GS, Portera-Cailliau C, Sagasti A. 2010 Wallerian degeneration of zebrafish trigeminal axons in the skin is required for regeneration and developmental pruning. *Development* **137**, 3895–3894. (doi:10.1242/dev.053611)

34. Yang J, Weiner RM, Kallop D, Olsen G, Wu Z, Renier N, Uyru K, Tessier-Lavigne M. 2013 Regulation of axon degeneration after injury and in development by the endogenous calpain inhibitor calpastatin. *Neuron* **80**, 1175–1189. (doi:10.1016/j.neuron.2013.08.014)

35. Villegas R, Martin SM, O'Donnell KC, Carrillo SA, Sagasti A, Allende ML. 2012 Dynamics of degeneration and regeneration in developing zebrafish peripheral axons reveals a requirement for extrinsic cell types. *Neural Dev.* **7**, 19. (doi:10.1186/1749-8104-7-19)
from naturally occurring developmental pruning. Neuron 50, 883–895. (doi:10.1016/j.neuron.2006.05.013)

49. Finn JT, Weil M, Ancher F, Simon R, Shiringavan A, Ruff MC. 2000 Evidence that Wallerian degeneration and localized axon degeneration induced by local neurotrophin deprivation do not involve caspases. J. Neurosci. 20, 1333–1341. (doi:10.1523/JNEUROSCI.04-01-2000.0333)

50. Mishra B, Carson R, Hume RJ, Collins CA. 2013 Sodium and potassium currents influence Wallerian degeneration of injured Drosophila axons. J. Neurosci. 33, 18 728–18 739. (doi:10.1523/JNEUROSCI.0077-13.2013)

51. Vargas ME, Yamagishi Y, Tessier-Lavigne M, Sagasti A. 2015 Live imaging of calcium dynamics during axon degeneration reveals two functionally distinct phases of calcium influx. J. Neurosci. 35, 15 026–15 038. (doi:10.1523/JNEUROSCI.2484-15.2015)

52. Schlaepfer WW. 1974 Calcium-induced degeneration of axoplasm in isolated segments of rat peripheral nerve. Brain Res. 69, 203–215. (doi:10.1016/0006-8993(74)90002-X)

53. Kerschesteiner M, Schwab ME, Lichtman JW, Misgeld T. 2005 In vivo imaging of axonal degeneration and remyelination in the injured spinal cord. Nat. Med. 11, 572–577. (doi:10.1038/nm1229)

54. Knørrle J et al. 2010 Mechanisms of acute axonal degeneration in the optic nerve in vivo. Proc Natl Acad Sci. USA 107, 6064–6069. (doi:10.1073/pnas.0907941107)

55. Loretto A, Di Stefano M, Gering M, Conforti L. 2015 Wallerian degeneration is executed by an NMN-SARM1-dependent late Ca2+ influx but only modestly influenced by mitochondria. Cell Rep. 13, 2539–2552. (doi:10.1016/j.celrep.2015.11.032)

56. Conforti L, Tarton A, Mack TG, Mi W, Buckmaster EA, Wagner D, Perry VH, Coleman MP. 2000 A U2af2/DAG10e chimeric protein and overexpression of Rbp9 in the slow Wallerian degeneration (Wdk) mouse. Proc. Natl Acad. Sci. USA 97, 11 377–11 382. (doi:10.1073/pnas.97.21.11377)

57. Sasai Y. 2018 Metabolic aspects of neuronal degeneration: from a NAD+ point of view. Neurosci. Res. 52, 442–447. (doi:10.1016/j.neures.2018.07.001)

58. Berger F, Lau C, Dahlmann M, Ziegler M. 2005 Subcellular compartmentation and differential catalytic properties of the three human nicotinamide mononucleotide adenyltransferase isoforms. J. Biol. Chem. 280, 36 334–36 341. (doi:10.1074/jbc.M508660200)

59. Ruan K, Zhu Y, Li C, Brazil JM, Zhai RG. 2015 Alternative splicing of Drosophila Nmnat functions as a switch to enhance neuroprotection under stress. Nat. Commun. 6, 10 057. (doi:10.1038/ncomms10057)

60. Gilley J, Coleman MP. 2010 Endogenous Nmnat2 is an essential survival factor for maintenance of healthy axons. PLoS Biol. 8, e1000300. (doi:10.1371/journal.pbio.1000300)

61. Yan T, Feng Y, Zheng J, Ge X, Zhang Y, Wu D, Zhao J, Qiao 2010 Nmnat2 delays axon degeneration in superior cervical ganglia dependent on its NAD synthesis activity. Neurochem. Int. 56, 101–106. (doi:10.1016/j.neuint.2009.09.007)

62. Milde S, Gilley J, Coleman MP. 2013 Subcellular localization determines the stability and axon protective capacity of axon survival factor Nmnat2. PLoS Biol. 11, e1001539. (doi:10.1371/journal.pbio.1001539)

63. Gilley J, Adalbert R, Yu G, Coleman MP. 2013 Rescue of peripheral and CNS axon defects in mice lacking Nmnat2. J. Neurosci. 33, 13 410–13 424. (doi:10.1523/JNEUROSCI.1534-13.2013)

64. Fang Y, Soares L, Teng X, Geary M, Bonini NM. 2012 A novel drosophila model of nerve injury reveals an essential role of Nmn1 in maintaining axonal integrity. Curr. Biol. 22, 590–595. (doi:10.1016/j.cub.2012.01.063)

65. Neuromm Li et al. 2017 Axon death pathways converge on Axundead to promote functional and axonal structure disassembly. Neuron 95, 78–91.e5. (doi:10.1016/j.neuron.2017.06.031)

66. Sasai Y, Vohra BPS, Baloh RH, Milbrandt J. 2009 Transgenic mice expressing the Nmnat1 protein manifest robust delay in axonal degeneration in vivo. J. Neurosci. 29, 6526–6534. (doi:10.1523/JNEUROSCI.1429-09.2009)

67. Avery MA, Sheehan AE, Kerr KS, Wang J, Freeman MR. 2009 WldS requires Nmnat1 enzymatic activity and NIF–VP3 interactions to suppress Wallerian degeneration. J. Cell Biol. 184, 501–513. (doi:10.1083/jcb.200808042)

68. Babetto E, Beirovski B, Janeckova L, Brown R, Gilley J, Thomson D, Ribchester RR, Coleman MP. 2010 Targeting Nmnat1 to axons and synapses transforms its neuroprotective potency in vivo. J. Neurosci. 30, 13 291–13 304. (doi:10.1523/JNEUROSCI.1189-10.2010)

69. Summers DW, Milbrandt J, DiAntonio A. 2013 The Phr1 ubiquitin ligase promotes axonal degeneration by tuning levels of Nmnat2 protein. PLoS Biol. 10, 1–18. (doi:10.1371/journal.pbio.1001440)

70. Desbois M, Crawley O, Evans PR, Baker ST, Masuho I, Yasuda R, Gill B. 2018 PAM forms an atypical SCF ligand complex that promotes Wallerian degeneration through regulation of axonal Nmnat2. Cell Rep. 17, 774–782. (doi:10.1016/j.celrep.2016.09.043)

71. Walker LJ, Summers DW, Sasai Y, Brace E, Milbrandt J, DiAntonio A. 2017 MAPK signaling promotes axonal degeneration by speeding the turnover of the axonal maintenance factor Nmnat2. Elife 6, e22540. (doi:10.7554/eLife.22540)

72. Miller BR, Press C, Daniels RW, Sasai Y, Milbrandt J, DiAntonio A. 2009 A dual leucine kinase-dependent axon self-destruction program promotes Wallerian degeneration. Nat. Neurosci. 12, 387–389. (doi:10.1038/nn.2290)

73. Shin JE et al. 2012 SCG10 is a JNK target in the axonal degeneration pathway. Proc. Natl Acad. Sci. USA 109, E3696–E3705. (doi:10.1073/pnas.1216041109)

74. Milde S, Fox AN, Freeman MR, Coleman MP. 2013 Deletions within its subcellular targeting domain enhance the axon protective capacity of Nmnat2 in vivo. Sci. Rep. 3, 1–9. (doi:10.1038/srep02567)

75. Osterhof JM et al. 2012 eSarm/Shepr1 is required for activation of an injury-induced axon death pathway. Science 337, 481–484. (doi:10.1126/science.1223899)

76. Gerdis J, Summers DW, Sasai Y, DiAntonio A, Milbrandt J. 2013 Sarm1-mediated axon degeneration requires both SARM and TIR interactions. J. Neurosci. 33, 13 569–13 580. (doi:10.1523/JNEUROSCI.1197-13.2013)

77. Gilley J, Orsomando G, Nascimento-Ferreira I, Coleman MP. 2015 Absence of SARM1 rescues development and survival of Nmnat2-deficient axons. Cell Rep. 10, 1975–1982. (doi:10.1016/j.celrep.2015.02.060)
degeneration, J. Cell Biol. 216, 477–493. (doi:10.1083/jcb.201606020)

101. Araki T, Wakatsuki S. 2018 Regulation of neuronal/axonal degeneration by ZNRF1 ubiquitin ligase. Neurosci. Res. 139, 21–25. (doi:10.1016/j.neures.2017.08.008)

102. Kinoshita Y, Kondo S, Takahashi K, Nagai J, Wakatsuki S, Araki T, Goshima Y, Otshuma T. 2019 Genetic inhibition of CRMP2 phosphorylation delays Wallerian degeneration after optic nerve injury. Biochem. Biophys. Res. Commun. 514, 1037–1039. (doi:10.1016/j.bbrc.2019.05.060)

103. Zhang J-N et al. 2016 Calpain-mediated cleavage of collapsin response mediator protein-2 drives acute axonal degeneration. Sci. Rep. 6, 37050. (doi:10.1038/srep37050)

104. Farley JE, Burdett TC, Barria R, Neukomm LJ, Kenna KP, Landers JE, Freeman MR. 2018 Transcription factor Pebbled/RRB1 regulates injury-induced axon degeneration. Proc. Natl Acad. Sci. USA 115, 1358–1363. (doi:10.1073/pnas.1715371115)

105. Rooney TM, Freeman MR. 2014 Drosophila models of neuronal injury. ILAR J. 54, 291–295. (doi:10.1093/ilar/iru007)

106. Tian W, Asgharsarharghi A, Valera G, Czopka T, Henninger N, Bouley J, Sikoglu HT, López-Schier H. 2018 Systemic loss of Sarm1 is glioprotective after neurotrauma. bioRxiv. (doi:10.1101/493163)

107. Gilley J, Ribchester RR, Coleman MP. 2017 Sarm1 deletion, but not WldS, confers lifelong rescue in a mouse model of severe axonopathy. Cell Rep. 21, 10–16. (doi:10.1016/j.celrep.2017.09.027)

108. Zhao ZY et al. 2019 A cell-permeant mimetic of NMN activates SARM1 to produce cyclic ADP-ribose and induce non-apoptotic cell death. Science 15, 452–466. (doi:10.1126/science.aab2768)

109. Lukacs M et al. 2019 Severe biallelic loss-of-function mutations in nicotinamide mononucleotide dehydrogenase 2 (NMNAT2) in two fetuses with fetal akinesia deformation sequence. Exp. Neurol. 320, 112961. (doi:10.1016/j.expneurol.2019.112961)

110. Huppke P et al. 2019 Homozygous NMNAT2 mutation in sisters with polyneuropathy and erythemalagia. Exp. Neurol. 320, 112958. (doi:10.1016/j.expneurol.2019.112958)

111. Ali YO, Bradley G, Li H-C. 2017 Screening with a NMNAT2-MSD platform identifies small molecules that modulate NMNAT2 levels in cortical neurons. Sci. Rep. 7, 43846. (doi:10.1038/srep43846)

112. Ferri A, Sanes JR, Coleman MP, Cunningham JM, Kato AC. 2003 Inhibiting axon degeneration and synapse loss attenuates apoptosis and disease progression in a mouse model of motoneuron disease. Curr. Biol. 13, 669–673. (doi:10.1016/S0960-9822(03)00206-9)

113. Kaneko S, Wang J, Kaneko M, Yiu G, Hurrell JM, Chitnis T, Khouy SJ, He Z. 2006 Protecting axonal degeneration by increasing nicotinamide adenine dinucleotide levels in experimental autoimmune encephalomyelitis models. J. Neurosci. 26, 9794–9804. (doi:10.1523/JNEUROSCI.2116-06.2006)

114. Meyer zu Horste G, Miesbach TA, Muller JI, Feldrich R, Stassart RM, Kieseier BC, Coleman MP, Sereda MW. 2011 The WldS transgene reduces axon loss in a Charcot-Marie-Tooth disease 1A rat model and nicotinamide delays post-traumatic axonal degeneration. Neurobiol. Dis. 42, 1–8. (doi:10.1016/j.nbd.2010.12.006)

115. Yaku K, Okabe K, Hikosaka K, Nakagawa T. 2018 NAD metabolism in cancer therapeutics. Front. Oncol. 8, 622. (doi:10.3389/fonc.2018.00622)

116. Henninger N, Bouley J, Sikoglu EM, An J, Moore CM, King JA, Bowser R, Freeman MR, Brew BH. 2016 Attenuated traumatic axonal injury and improved functional outcome after traumatic brain injury in mice lacking Sarm1. Brain 139, 1094–1105. (doi:10.1093/brain/aww001)

117. Ziegas NK, Klotzlos VE. 2018 Primary traumatic axonopathy in mice subjected to impact acceleration: a reappraisal of pathology and mechanisms with high-resolution anatomical methods. J. Neurosci. 38, 4031–4047. (doi:10.1523/JNEUROSCI.2343-17.2018)

118. Geisler S, Huang SX, Strickland A, Doan RA, Summers DW, Mao X, Park J, DiAntonio A, Milbrandt J. 2019 Gene therapy targeting SARM1 blocks pathologic axonal degeneration in mice. J. Exp. Med. 216, 294–303. (doi:10.1084/jem.20181040)

119. Geisler S, Doan RA, Strickland A, Huang X, Milbrandt J, DiAntonio A. 2016 Prevention of vincristine-induced peripheral neuropathy by genetic deletion of SARM1 in mice. Brain 139, 3092–3106. (doi:10.1093/brain/awv251)

120. Turkew E, Falconer D, Reed N, Höke A. 2017 Deletion of Sarm1 gene is neuroprotective in two models of peripheral neuropathy. J. Peripheral Nerv. Syst. 22, 162–171. (doi:10.1111/jps.12219)

121. Grisold W, Cavaletti G, Windenbak AJ. 2012 Peripheral neuropathies from chemotherapeutics and targeted agents: diagnosis, treatment, and prevention. Neuro. Oncol. 14, iv45–iv54. (doi:10.1093/neucom/nos203)

122. Jaster-Swiftly K, Smith AG. 2016 Updates in diabetics peripheral neuropathy. F1000Res. 5, 738. (doi:10.12688/f1000research.7981.1)

123. Peters OM, Lewis EA, Osterloh JM, Weiss A, Salameh JS, Mettlerie J, Brown RH, Freeman MR. 2018 Loss of Sarm1 does not suppress motor neuron degeneration in the SOD1G93A mouse model of amyotrophic lateral sclerosis. Hum. Mol. Genet. 27, 3761–3771. (doi:10.1093/hmg/ddy260)