# Constitutively active SARM1 variants found in ALS patients induce neuropathy A. Joseph Bloom<sup>1\*</sup>, Xianrong Mao<sup>1</sup>, Amy Strickland<sup>1</sup>, Yo Sasaki<sup>1</sup>, Jeffrey Milbrandt<sup>1\*</sup>, Aaron DiAntonio<sup>2\*</sup>

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- 22
- 23 **Running title:** *SARM1* ALS patient variants

#### 1 Abstract

In response to injury, neurons activate a program of organized axon self-destruction initiated by 2 3 the NAD<sup>+</sup> hydrolase SARM1. In healthy neurons SARM1 is autoinhibited, but single amino acid 4 changes can abolish autoinhibition leading to constitutively-active SARM1 enzymes that 5 promote degeneration when expressed in cultured neurons. To investigate whether naturally-6 occurring human variants might similarly disrupt SARM1 autoinhibition and potentially 7 contribute to risk for neurodegenerative disease, we assayed the enzymatic activity of 29 rare 8 SARM1 alleles identified among 8,507 amyotrophic lateral sclerosis (ALS) patients. Ten 9 missense variants or small in-frame deletions exhibit constitutive NADase activity, including 10 more than half of those that are unique to the ALS patients or that occur in multiple patients. 11 Expression of these constitutively active ALS-associated SARM1 alleles in cultured dorsal root 12 ganglion (DRG) neurons is pro-degenerative and cytotoxic. Intrathecal injection of an AAV 13 expressing the common SARM1 reference allele is innocuous to mice, but a construct harboring SARM1<sup>V184G</sup>, the constitutively active variant found most frequently in the ALS patients, causes 14 15 axon loss, motor dysfunction, and sustained neuroinflammation. These results implicate rare 16 hypermorphic SARM1 alleles as candidate genetic risk factors for ALS and other 17 neurodegenerative conditions.

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#### 19 Keywords: ALS; SARM1; neurodegeneration; axon; NAD

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#### 21 Main Text

Trauma and disease in the nervous system activate an intrinsic axon self-destruction pathway,also known as Wallerian degeneration, which facilitates the orderly clearance of damaged axon

1 segments. This choice between maintaining or actively dismantling axons is primarily 2 determined by the action of SARM1, a TIR-containing NAD<sup>+</sup> hydrolase that cleaves NAD<sup>+</sup> to generate nicotinamide and cyclic ADPR (cADPR), a useful biomarker of SARM1 activity<sup>1</sup>. In 3 4 healthy neurons, SARM1 is maintained in an autoinhibited state, but injury- or disease-induced 5 depletion of the axon survival factor NMNAT2 activates SARM1 leading to a rapid loss of NAD<sup>+</sup>, metabolic catastrophe, and axon fragmentation<sup>2-4</sup>. SARM1 knockout mice are viable and 6 7 without apparent phenotypes under routine conditions, but are protected against 8 neurodegeneration in models of axotomy, traumatic brain injury, peripheral neuropathy, glaucoma, and retinal degenerative diseases<sup>5-11</sup>. Conversely, mutations that decrease NMNAT2 9 activity lead to polyneuropathy in both humans and model organisms<sup>12,13</sup>, suggesting that 10 11 aberrant SARM1 activation has a role in human disease. Furthermore, the recent observation that single point mutations in *SARM1* can disrupt enzyme autoinhibition  $^{14-18}$  led us to speculate that 12 13 naturally-occurring human variants might similarly dysregulate SARM1 and thereby increase 14 disease risk.

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16 To investigate whether SARM1 mutations that disrupt its autoinhibition are associated with 17 neurogenerative disorders, we sought to identify rare prodegenerative SARM1 missense variants 18 in human databases. ALS warrants particular attention because peripheral axon degeneration accompanies and may precede motoneuron death during ALS progression<sup>19,20</sup>. Here, we identify 19 20 over two dozen such polymorphisms found in ALS patients and interrogate the activities of the 21 encoded enzymes in cultured neurons. Provocatively, the majority of our strongest candidate 22 variants disrupt SARM1 regulation and confer constitutive activity in vitro. Furthermore, 23 expression of a constitutively-active SARM1 allele found in three unrelated patients causes an

1 ALS-like phenotype-motor dysfunction, axon loss and sustained neuroinflammation-when

2 expressed in the mouse spinal cord via intrathecal delivery.

3 We identified a total of 29 SARM1 coding variants (missense and small in-frame deletions)

4 culled from three large publicly-accessible ALS consortia databases that include 8,507 cases in

total (Table 1) $^{21-23}$ . 5

6

7	Table 1 Rare SARM1 missense variants and in-frame deletions found in ALS patients
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Variant	rsID	occu		Number of occurrences in ALS patients	
		African	East Asian	European <sup>b</sup>	P2
Constitutively	/ active				
∆226-232	rs782325355	0	0	0.01	2 <sup>d</sup>
∆249-252		0	0	0	1 <sup>d</sup>
V184G	rs71373646	0.007	0.006	0	3 <sup>e</sup>
G206S	rs1555585199	0	0	0	2 <sup>e</sup>
L223P		0	0	0	1 <sup>d</sup>
R267W	rs11652384	0	0	0.001	1 <sup>e</sup>
V331E	rs1555585331	0	0	0	1 <sup>d</sup>
E340K	rs781854217	0	0	0.003	1 <sup>d</sup>
T385A		0	0	0	1 <sup>d</sup>
T502P	rs782421919	0	0	0.006	2 <sup>d,f</sup>
Not constituti	vely active				
A240E	rs1449836804	0	0	0.004	1 <sup>d</sup>
R244S		0	0	0	1 <sup>d</sup>
A250T	rs1555585243	0	0.06	0	1 <sup>d</sup>
A275V	rs376587698	0	0	0.006	1 <sup>d</sup>
R310H	rs369186722	0	0	0.01	1 <sup>d</sup>
A341V	rs373458416	0	0	0.01	2 <sup>d</sup>
R403P	rs782706244	0	0	0.0009	1 <sup>f</sup>
E431G	rs1555585662	0	0	0.001	1 <sup>d</sup>
R465T		0	0	0	1 <sup>d</sup>
R484C	rs1555585809	0	0	0.0009	1 <sup>d</sup>
A488E	rs782228906	0.004	0	0.02	2 <sup>d</sup>
V518L	rs782106973	0	0	0c	3 <sup>d</sup>
R569C	rs571724138	0	0	0.005	1 <sup>d</sup>
R570Q	rs539229444	0	0.008	0.005	1 <sup>e</sup>
D637Y	rs1451417529	0	0	0	1 <sup>d</sup>
A646S	rs782676389	0	0	0.0008	1 <sup>d</sup>
M672V	rs782774927	0	0	0.004	2 <sup>e,f</sup>
S684F	rs782256561	0.004	0	0.004	1 <sup>f</sup>
R702C	rs781850558	0	0.005	0.002	1 <sup>f</sup>

<sup>9</sup> 10 11 12 13 14 15 16

bnon-Finnish °V518L = 0.009% in gnomAD v3 non-Finnish Europeans

<sup>a</sup>gnomAD v2

dMinE<sup>32</sup>

eALS Variant Server<sup>34</sup>

fALS Knowledge Portal<sup>33</sup>

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1 Altogether, rare *SARM1* variants (i.e. with allele frequencies <0.01% in all gnomAD 2 populations<sup>24</sup>) occur in >0.9% of ALS cases but in only ~0.25% of controls<sup>21,22</sup>, and in only 3 ~0.3% of the general population<sup>24</sup>. For comparison, potentially pathogenic *TBK1* variants were 4 reported in 1.1% of ALS cases and 0.19% of controls<sup>25</sup>. This suggests that rare pathogenic 5 *SARM1* alleles might contribute to ALS risk.

6

7 To investigate whether these variants disrupt autoinhibition, we assayed the NAD<sup>+</sup> hydrolase 8 activities of the encoded enzymes. We prioritized the variants and first tested a) those identified 9 in multiple ALS patients but not in healthy controls and b) those unique to ALS patients (i.e. not 10 reported in any prior human study as of January 2020). These 15 SARM1 variants (Figure 1A, 11 Table 1) account for 51% (20/39) of rare SARM1 variants in ALS patients genotyped in the three 12 large ALS databases we investigated. To examine the properties of these mutants, we tested them 13 in  $Sarm1^{-/-}$  mouse dorsal root ganglion DRG neurons. We prepared lentiviruses for the 15 SARM1 mutant constructs, infected Sarm1<sup>-/-</sup> neurons, and assessed their NAD<sup>+</sup> hydrolase 14 15 activity. Eight of these variants were determined to be constitutively active, i.e. we found that the baseline level of NAD<sup>+</sup> was decreased in neurons expressing these mutant constructs and the 16 level of cADPR, a specific *in vitro* and *in vivo* SARM1 biomarker<sup>26</sup> was increased (Fig. 1). By 17 contrast, *SARM1*<sup>P332Q</sup>, the only variant common in any gnomAD population (1.1% in Europeans) 18 19 is not constitutively active (Fig 1).

20

Encouraged by these results, we assayed the activities of an additional 14 rare missense variants. These were considered poorer candidates because each is observed in only a single ALS patient and they are not unique to the patients as they are also found in the gnomAD database. Among

these, we identified two additional constitutively active variants (Table 1). In total, 40% (4/10) of
the SARM1 variants with constitutive NAD<sup>+</sup> hydrolase activity occur in multiple ALS patients.

4 Point mutants that disrupt SARM1's autoinhibitory interfaces result in dysregulation of SARM1 activity and promote the degeneration of cultured neurons $^{14-18}$ . Consistent with those findings, 5 6 lentiviral-mediated expression of all the constitutively active variants we tested (Table 1) alters the cell body morphology of cultured Sarm1<sup>-/-</sup> mouse DRG neurons consistent with cell death. 7 To quantify this pro-degenerative activity, two variant constructs,  $SARM1^{V184G}$  and  $SARM1^{A226-}$ 8 <sup>232</sup>, were studied further. The mutant enzymes were expressed in  $Sarm1^{-/-}$  DRGs neurons and 9 10 degeneration was measured by two methods. Fluorescently-labeled Annexin V, which binds to 11 phosphatidylserine, was used to determine whether the expression of either variant construct 12 significantly compromises axon health. Annexin V binding is a useful proxy for axon health as 13 neurites undergoing Wallerian degeneration expose phosphatidylserine on their extracellular surfaces similarly to apoptotic cells<sup>27</sup>. Neuronal death was quantified using an oxidoreductase 14 15 activity assay, a common measure of cell viability. Both assays demonstrated that both ALS-16 associated SARM1 variants produced a significant degenerative effect relative to the common 17 reference allele (Fig. 1).

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3

While these variants exhibit constitutive NAD<sup>+</sup> hydrolase activity, it is formally possible that they mediate their pro-degenerative effects via a distinct toxic mechanism. To investigate this alternate hypothesis, we generated constructs containing two mutations, either of the ALSassociated variants,  $SARM1^{V184G}$  or  $SARM1^{A226-232}$ , together with E642A, a point mutation in the TIR domain that disrupts the catalytic glutamate required for SARM1 NAD<sup>+</sup> hydrolase activity

and axon degeneration<sup>3</sup>. In both cases, introducing E642A abolishes enzymatic activity and the
detrimental effects of the constructs on cell body and axon health (Fig. 1). Hence, these ALS
patient-derived *SARM1* variants promote degeneration via loss of autoinhibition and resulting
constitutive NAD<sup>+</sup> hydrolase activity.

5

6 To test whether the rare SARM1 variants promote neurodegeneration in vivo, AAV viral vectors 7 were administered intrathecally to male and female six-week old wild-type mice, expressing either the common human allele of SARM1 (the reference allele) or SARM1<sup>V184G</sup>, the 8 9 constitutively active variant found most frequently in the ALS patient databases. In these 10 constructs, each SARM1 protein was fused to EGFP and expressed under the control of the 11 human synapsin promoter. The AAV viruses were produced with these constructs and a mixture of PHP.S and PHP.eB serotype capsids (both derived from AAV9<sup>28</sup>) in order to infect neurons in 12 13 the spinal cord and DRGs.

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Animals injected with AAV expressing the common *SARM1* allele had no discernible behavioral phenotypes for at least 12 weeks. By contrast, those injected with AAV-*SARM1*<sup>V184G</sup> exhibited motor impairment. Two of the mice rapidly progressed to full limb paralysis 3-4 days after injection. Other animals injected with *SARM1*<sup>V184G</sup> (7/9) displayed less dramatic motor deficits characterized by hindlimb clasping<sup>29</sup>, and significant muscle weakness, as measured by the inverted screen assay (Fig. 2). These deficits were detected within 3 weeks of injection and did not progress significantly through the 12-week observation period.

22

To characterize the neurodegeneration caused by  $SARM^{V184G}$  expression, the intrathecally 1 2 injected mice were examined for evidence of axon degeneration and neuron loss. We examined 3 the two mice that became rapidly paralyzed and the other mice with less severe disease as 4 separate cohorts because of the difference in phenotype. In the spinal cords of the paralyzed 5 mice, there was clear evidence of cell death around the ependymal canal as detected by TUNEL staining. Neuroinflammation was observed throughout the spinal cord of these mice as evidenced 6 7 by prevalent staining for CD68, a marker of activated macrophages. Neither of these phenotypes 8 were observed in animals injected with the common SARM1 allele construct (Additional File 1). 9 These mice did not display obvious myelin defects or vacuolization in the sural, sciatic or tibial 10 nerves at 3-4 days post-infection.

11

The mice treated with  $SARM1^{V184G}$  that displayed a less severe behavioral response were 12 13 sacrificed twelve weeks post-injection. Pathological inspection of their spinal cords revealed no 14 evidence of ongoing apoptosis or elevated CD68 staining. Their peripheral nerves, however, 15 contain almost 10-fold more CD68-positive macrophages than those treated with the control SARM1 allele (Fig. 2). Macrophages also increase in size upon activation<sup>30</sup>, and the SARM1<sup>V184G</sup>-16 17 infected mice have a 1.6-fold greater CD68-stained area per cell than do control mice, yielding a 15.2-fold difference in total CD68-stained area. Hence, neuronal expression of SARM1<sup>V184G</sup> 18 19 triggers an elevated inflammatory response in peripheral nerves that persists for at least twelve weeks after treatment<sup>31</sup>. 20

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The average fiber densities in the peripheral nerves of the *SARM1*<sup>V184G</sup>-injected mice are also lower than those injected with AAV expressing the common human allele, demonstrating that

1 expression of the constitutively active SARM1 promotes axon loss. The number of axons were 2 counted in the sural, sciatic and tibial nerves. In both the sural and sciatic nerves, the density of 3 axons is significantly lower (p < 0.001), a trend that is evident in the tibial nerve, though it did not 4 reach statistical significance (Fig. 2). The average axon size and extent of myelination (g-ratio) 5 does not differ significantly between the variant and common SARM1 allele treated animals 6 (p>0.1, n=15). Myelin defects and vacuolization are not observed in these nerves, indicating a 7 lack of ongoing axon loss. The lack of axon defects at twelve weeks is consistent with the early, but stable, deficits in motor function observed in mice receiving the SARM1<sup>V184G</sup> virus (Fig. 2). 8 9 We interpret these data as evidence that a subset of neurons-those sufficiently susceptible to 10 SARM1-dependent degeneration and infected with virus—lost their axons before three weeks, 11 while others, including uninfected neurons, remained healthy and functional up to twelve weeks. 12 Inter-animal differences in motor dysfunction severity likely reflect variability in infection 13 efficiency.

14

15 In summary, we find that many rare SARM1 variants found in ALS patients also lack normal 16 autoinhibition, and that such an allele induces neurodegeneration and neuroinflammation when 17 expressed in the mouse nervous system. We therefore propose that hypermorphic SARM1 18 mutations are a candidate congenital risk factor for ALS. The mechanism by which constitutive 19 NAD<sup>+</sup> hydrolase activity would predispose to neurodegeneration appears straightforward as low 20 NAD<sup>+</sup> is a death sentence for energy-hungry neurons and is associated with both disease and aging-related functional defects<sup>32</sup>. We speculate that the contrast between virus-transfected mice 21 22 that rapidly display severe degenerative phenotypes, and human ALS patients who are typically 23 diagnosed only after several decades of life, likely reflects the difference in SARM1

1	expression-i.e. viral over-expression precipitates abrupt metabolic catastrophe in this model,
2	whereas chronic suboptimal $NAD^+$ levels lead to gradual motoneuron attrition in patients.
3	Fortunately, small molecule SARM1 inhibitors are already in development <sup>33</sup> , and we have shown
4	that a SARM1 dominant negative gene therapy can potently block the SARM1 programmed axon
5	degeneration pathway in mice <sup>34</sup> . Establishing that SARM1 inhibition is safe and effective in
6	carriers of pathogenic SARM1 variants could provide a vital stepping stone toward the use of
7	SARM1-directed therapeutics more generally for ALS and other diseases that involve axon
8	degeneration.
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12	Methods
13	Mice
14	Male and female WT and Sarm1 knockout C57/BL6 mice were housed and used under the
15	direction of institutional animal study guidelines at Washington University in St. Louis. The
16	inverted screen test of strength was performed as previously <sup>35</sup> . All protocols received
17	institutional IACUC approval.
18	
19	DRG culture
20	Mouse DRG culture was performed as previously described <sup>36</sup> . DRG were dissected from
21	embryonic day 13.5 Sarm1 knockout C57/BL6 mouse embryos and cells suspended in growth
22	medium at a concentration of $\sim 7 \times 10^6$ cells/ml in 96- well tissue culture plates (Corning) coated
23	with poly-d-Lysine (0.1 mg/ml; Sigma) and laminin (3 µg/ml; Invitrogen). For axotomy,

suspended neurons (2 μl) were placed as a drop in the center of each well and incubated at 37°C
with 5% CO<sub>2</sub> for 15 min, after which media was added to each well. Lentiviral particles
containing *SARM1* variants were generated as previously described<sup>36</sup>. Lentivirus was added (1–
10 × 10<sup>3</sup> pfu) after 1–2 days (DIV) and metabolites were extracted or assays were performed at
6–7 DIV. Cell death was quantified by assaying mitochondrial function (MTT assay), as
previously described<sup>37</sup>.

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#### 8 Automated quantification of axon degeneration

9 The axon degeneration index, the ratio of fragmented axon area over total axon area, was quantified as previously described<sup>36</sup>. To quantify Annexin V staining, the Alexa Fluor<sup>™</sup> 568 10 11 conjugate (ThermoFisher) was added to the cultured neurons at a 1:100 dilution four days after 12 viral infection. Bright field and fluorescent images were acquired one hour later using 13 Operetta. Unbiassed image analysis was performed using ImageJ as follows: total axon area was 14 measured from the binary bright field images after subtracting background. For Annexin 15 fluorescent intensity measurement, the fluorescent images were background subtracted and then 16 annexin positive area was defined using the particle analyzer. Data was reported as the total 17 fluorescent intensity of the annexin positive area divided by the axon area.

18

#### 19 DRG metabolite extraction and metabolite measurement

At DIV6, tissue culture plates were placed on ice and culture medium replaced with ice-cold
saline (0.9% NaCl in water, 500 μl per well). Saline was removed and replaced with 160 μl ice
cold 50% MeOH in water. Solution was transferred to tubes containing 50 μl chloroform on ice,
shaken vigorously, and centrifuged at 20,000*g* for 15 min at 4 °C. The clear aqueous phase

(140 μl) was transferred into microfuge tubes and lyophilized under vacuum. Lyophilized
 samples were reconstituted with 5 mM ammonium formate (15 μl), centrifuged (13,000 g, 10
 min, 4°C), and 10 μl of clear supernatant was analyzed. NAD<sup>+</sup> and cADPR were measured using
 LC-MS/MS as previously described<sup>26</sup>.

5

#### 6 AAV constructs and virus injections

AAV particles with a mixture of Php.s and Php.eB capsids<sup>28</sup>, containing a human *SARM1* gene construct fused to EGFP, under the control of the human synapsin promoter, were produced by the Viral Vector Core of the Hope Center for Neurological Disorders at Washington University in St. Louis. Viral particles were purified by iodixanol gradient ultracentrifugation and virus titers were measured by dot blot. Under light anesthesia with Avertin,  $6 \times 10^{11}$  viral genomes were injected intrathecally at L6/S1. Viral expression in mice 12-weeks post injection was confirmed by detecting EGFP expression via immunohistochemical analysis of sectioned DRGs.

#### 15 Immunohistochemistry, imaging and quantification

16 After perfusion with PBS followed by 4% PFA in PBS, tissues were fixed in 4% PFA in PBS for 17 1 h at room temperature and placed in 30% sucrose in PBS overnight at 4°C, then embedded in 18 OCT (Tissue-Tek), frozen on dry ice, and then stored at -80°C. Longitudinal sections of 6 µm or 19 cross-sections of 20 µm were obtained using a cryostat and slides were stored at -20°C. DRG 20 and nerve slides were post-fixed in cold acetone, then washed with PBS. Spinal cord slides were 21 simply washed three times in PBS. All slides were subsequently blocked with 4% BSA and 1% 22 Triton X-100 in PBS and incubated with rat anti-CD68 (1:500; Bio-Rad) and mouse-anti-GFP 23 conjugated to Alexa Fluor 488 (1:250; Thermo Fisher Scientific) overnight in the blocking

1	buffer. Slides were then washed, incubated in secondary antibodies (Jackson ImmunoResearch
2	Laboratories) and mouse anti-GFP conjugated to AlexaFlour 488 (1:250, Thermo Fisher
3	Scientific), washed, and mounted in Vectashield with DAPI. Slides were imaged using a DMI
4	4000B confocal microscope (Leica Microsystems) with a $20\times$ oil objective and DFC 7000-T
5	camera (Leica Microsystems). For quantification, at least four images were measured per animal.
6	CD68-positive cells were counted by a researcher blinded to the images' treatment group. The
7	total CD68-stained area and nerve area in each image was quantified with the particle analyzer in
8	ImageJ using a uniform threshold.
9	
10	TUNEL apoptosis detection
11	TUNEL was performed as previously described <sup>38</sup> . Slides prepared for immunohistochemistry
12	were thawed then postfixed with 4% PFA for 10 min at room temperature, washed thoroughly
13	with PBS, incubated with 10 $\mu$ g/ml proteinase K for 15 min at 37°C, then washed with PBS. A
14	positive control slide was incubated in DNase I (1 U/ml) for 1 h at RT, then washed with PBS.
15	Slides were then pretreated with TdT buffer (25 mm Tris-HCl, 200 mm sodium cacodylate, 0.25
16	mg/ml BSA, 1 mm cobalt chloride, Roche Diagnostics) at RT for 10 min. To perform end-
17	labeling, TdT buffer was combined with terminal deoxynucleotidyl transferase (Roche
18	Diagnostics, 400 U/slide) and Biotin-16-dUTP (Roche Diagnostics, 4 $\mu$ m) and added to slides
19	for 1 h at 37°C. Slides were thoroughly washed with PBS, then blocked for 30 min with 5%
20	normal goat serum in PBS with 0.3% Triton-X, then incubated with Alexa-Fluor-conjugated
21	streptavidin (Jackson ImmunoResearch Laboratories) for 30 min at 37°C. Slides were washed
22	and mounted in Vectashield with DAPI.
23	

1	Toluidine blue staining and axon quantification
2	Sural, sciatic and tibial nerves were fixed in 3% glutaraldehyde in 0.1 M PBS, processed and
3	imaged as previously described <sup>8</sup> . Micrographs were stitched using Leica software and axons
4	were counted using ImageJ. To determine axon size distribution and G ratios of the sciatic nerve,
5	four nonoverlapping areas per cross section were imaged with a $100 \times$ oil objective of a Zeiss
6	Axioskop and photographed with a Hitachi camera. Photographs were analyzed using a
7	previously described binary imaging analysis method <sup>39</sup> .
8	
9	Statistical analysis
10	Two-tailed significance is reported throughout. All statistics were calculated using the R
11	software package. All data is available upon request.
12	
13	
14	Abbreviations
15	AAV: Adeno-associated virus
16	ALS: Amyotrophic Lateral Sclerosis
17	<b>ARM:</b> HEAT/Armadillo motif
18	cADPR: Cyclic adenosine diphosphate ribose
19	<b>CD68:</b> Cluster of Differentiation 68
20	DRG: Dorsal root ganglion
21	EGFP: Enhanced green fluorescent protein
22	MLS: Mitochondrial localization signal
23	MTT: 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide

- 1 NAD: Nicotinamide adenine dinucleotide
- 2 **NMNAT2:** Nicotinamide mononucleotide adenylyltransferase 2
- 3 **TIR:** Toll/Interleukin receptor
- 4 **TUNEL:** Terminal deoxynucleotidyl transferase dUTP nick end labeling
- 5 **SAM:** Sterile alpha motif
- 6 **SARM1:** Sterile alpha and TIR motif containing 1
- 7

#### 8 **Declarations**

#### 9 Ethical Approval and Consent to participate

- 10 All studies were approved by the Washington University Institutional Animal Care and Use
- 11 Committee.
- 12 **Consent for publication**
- 13 Not Applicable

#### 14 Availability of supporting data

15 All data relevant to this study are contained within the article.

#### 16 **Competing interests**

- 17 A.D. and J.M. are co-founders, scientific advisory board members and shareholders of Disarm
- 18 Therapeutics, a wholly owned subsidiary of Eli Lilly. A.J.B. and Y.S. are consultants to Disarm
- 19 Therapeutics. The authors have no other competing conflicts or financial interests.

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- 23 Authors' contributions

1	A.J.E	B, J.M and A.D designed the study. X.M performed in vitro experiments. A.S		
2	perfo	rmed in vivo experiments. Y.S assisted with mass spec analysis and method		
3	devel	opment. A.J.B. analyzed the data. A.J.B., J.M. and A.D. drafted and edited the		
4	figur	es and manuscript. All authors read and approved the final manuscript.		
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10	Auth	ors' information		
11	Not A	Applicable		
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14				
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#### 1 Figure Legends

2

#### 3 Figure 1. Dysregulated SARM1 variants found in ALS patients promote

4 **neurodegeneration.** (A) Schematic representation of the domain structure of SARM1.

5 Constitutively-active variants are indicated above in red. Bold variants were prioritized because

6 they were identified in multiple ALS patients or were unique to ALS patients.  $\Delta$  indicates an in-

7 frame deletion. MLS, mitochondrial localization signal; ARM, HEAT/Armadillo motif; SAM,

8 sterile alpha motif; TIR, Toll/interleukin-1 receptor homology domain. (**B**) cADPR and NAD<sup>+</sup>

9 levels from cultured Sarm1<sup>-/-</sup> DRG neurons infected with variant human SARM1 constructs,

10 performed in triplicate, relative to the reference SARM1 allele. (C) Neuron death as measured by

11 the MTT assay and (**D**) axon degeneration as measured by Annexin V staining in Sarm1<sup>-/-</sup> DRG

12 neurons infected with lentivirus expressing SARM1 variant constructs as well as double mutant

13 constructs including E642A, a point mutation that disrupts SARM1 NAD<sup>+</sup> hydrolase activity,

14 relative to the common SARM1 reference allele, all performed in triplicate. (E) Representative

15 bright-field and Annexin V-stained images of axons from Sarm1<sup>-/-</sup> DRG cultures infected with

variant and *SARM1* reference allele constructs. \*p<0.05; \*\*p<0.005; \*\*\*p<0.0005 difference

17 from reference allele, two-tailed t-test.

18

#### 19 Figure 2. Motor dysfunction, neuroinflammation and axon loss in mice injected

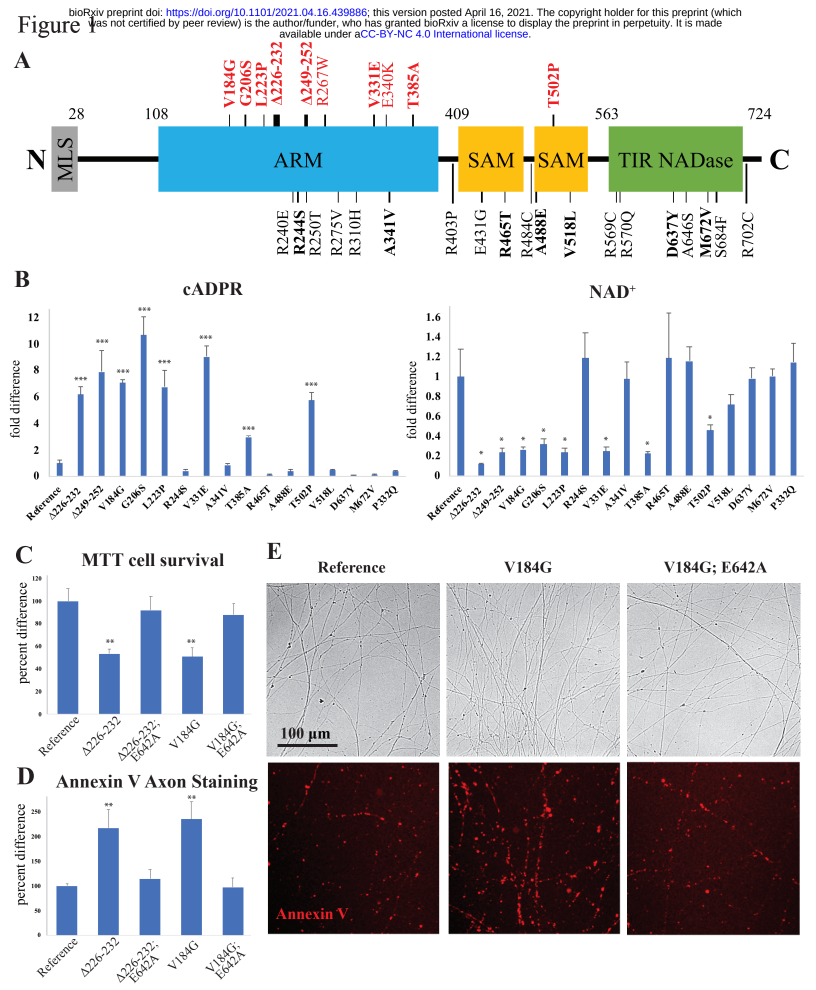
20 intrathecally with a SARM1<sup>V184G</sup> AAV construct. (A) Average time suspended from an

21 inverted screen (maximum 120 seconds, performed in triplicate) for C57/BL6 mice injected with

22 a human SARM1 reference allele (n=8) or SARM1<sup>V184G</sup> (n=7) AAV compared to uninjected

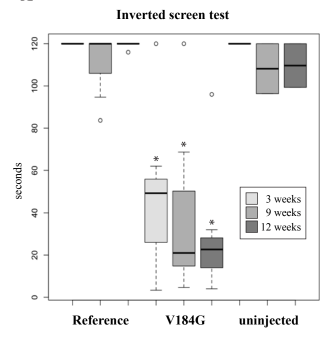
controls (n=3) 3, 9 and 12 weeks post-injection. \*p<0.005 difference from both the reference

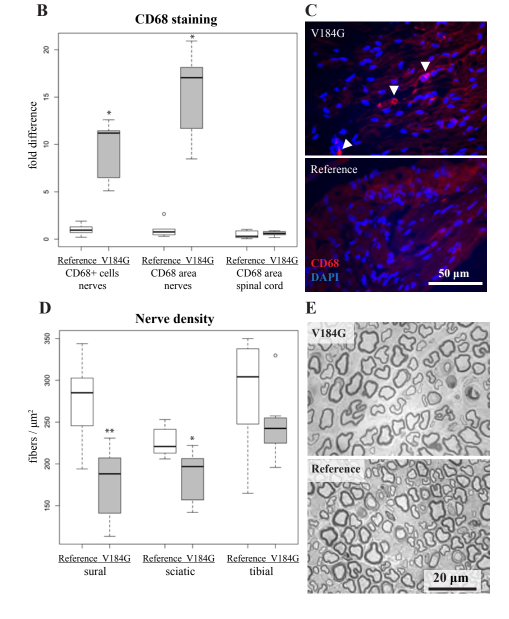
1	allele and uninjected controls, 2-tailed t-test. (B) The normalized average number of cells stained
2	by the macrophage marker anti-CD68 in nerve, and the average percent area of total anti-CD68
3	staining in nerve and in spinal cord sections, from C57/BL6 mice injected with a SARM1 <sup>V184G</sup>
4	AAV construct (3 images per mouse, n=7 mice) relative to those injected with a human SARM1
5	reference allele construct (n=8 mice) 12 weeks post-injection; $*p<10^{-4}$ , 2-tailed t-test. (C)
6	Representative images of nerve stained with DAPI and anti-CD68 from mice 12 weeks after
7	injection with a $SARM1^{V184G}$ or reference allele construct. ( <b>D</b> ) Average fibers per cross-sectional
8	$\mu m^2$ in sural, sciatic and tibial nerves from mice 12 weeks after injection with a SARM1 <sup>V184G</sup> (3)
9	images per mouse, n=7 mice) or reference allele construct (n=8 mice); *p<0.05; **p<0.001, 2-
10	tailed t-test. (E) Representative images of toluidine blue stained sural nerve sections.
11	
12	Additional file 1. Rapid cell death and neuroinflammation in mice injected intrathecally
13	with a SARM1 <sup>V184G</sup> AAV construct.
14	(A) Representative images of spinal cord sections, with closeup of ependymal canal, stained with
15	DAPI and the apoptosis marker TUNEL from mice 2 days after injection with a SARM1 <sup>V184G</sup> or
16	
	SARM1 human reference allele construct. (B) Representative images of spinal cord and (C)
17	<i>SARM1</i> human reference allele construct. ( <b>B</b> ) Representative images of spinal cord and ( <b>C</b> ) adjacent nerve sections stained with DAPI and the macrophage marker anti-CD68 from mice 2
17 18	



v

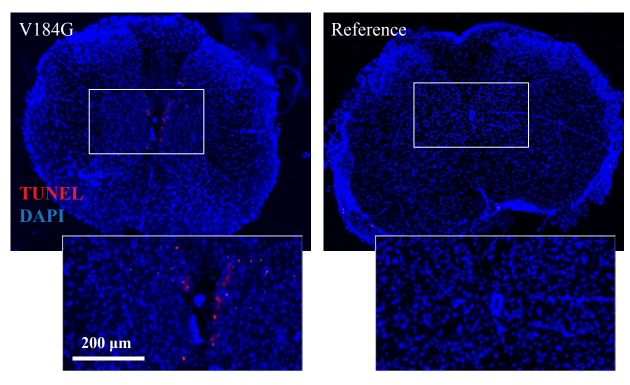
#### A



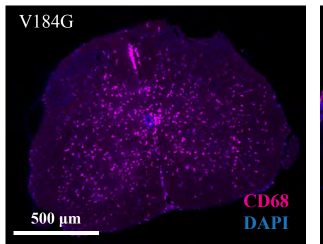


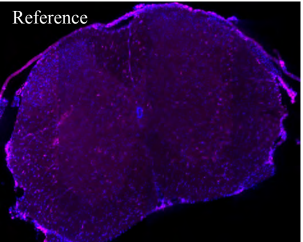
## Additional File 1

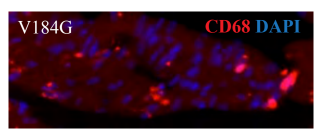
### A



B







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