Pathogenic role of delta 2 tubulin in bortezomib-induced peripheral neuropathy

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The pathogenesis of chemotherapy-induced peripheral neuropathy (CIPN) is poorly understood. Here, we report that the CIPN-causing drug bortezomib (Bort) promotes delta 2 tubulin (D2) accumulation while affecting microtubule stability and dynamics in sensory neurons in vitro and in vivo and that the accumulation of D2 is predominant in unmyelinated fibers and a hallmark of bortezomib-induced peripheral neuropathy (BIPN) in humans. Furthermore, while D2 overexpression was sufficient to cause axonopathy and inhibit mitochondria motility, reduction of D2 levels alleviated both axonal degeneration and the loss of mitochondria motility induced by Bort. Together, our data demonstrate that Bort, a compound structurally unrelated to tubulin poisons, affects the tubulin cytoskeleton in sensory neurons in vitro, in vivo, and in human tissue, indicating that the pathogenic mechanisms of seemingly unrelated CIPN drugs may converge on tubulin damage. The results reveal a previously unrecognized pathogenic role for D2 in BIPN that may occur through altered regulation of mitochondria motility.

delta 2 tubulin | mitochondria | DRG | bortezomib | axonopathy

Chemotherapy-induced peripheral neuropathy (CIPN) is a “dying back” neuropathy that features a distal-to-proximal peripheral nerve degeneration that frequently occurs in cancer patients undergoing chemotherapy (1). Modification of the chemotherapy treatment of a patient is sometimes required to limit the severity of CIPN, preventing patients from receiving effective cancer treatment (2). The pathogenesis of CIPN is largely unknown, and this incomplete knowledge is a main reason for the absence of effective neuroprotection strategies while maintaining chemotherapy drug anticancer activities (2–5). Several classes of anticancer drugs with different antineoplastic mechanisms can induce CIPN (6–8). However, sensory impairment is the predominant adverse effect associated with each class, suggesting the existence of a common mechanism of pathogenesis (2).

Tubulin and microtubules (MTs) are well-established targets for multiple anticancer drugs that induce CIPN (9). The contribution of the MT changes to the onset of CIPN is not well understood but is strongly implicated as the determining factor. MT hyperstabilization plays a direct role in paclitaxel neurotoxicity (10), and a single nucleotide polymorphism in TUBB2a, a gene encoding a tubulin isoform, is associated with an enhanced risk of CIPN (11). Moreover, MT dynamics and stability are influenced by nicotinamide adenine dinucleotide (NAD+) levels through sirtuin modulation (12), suggesting a tubulin-mediated mechanism also for the NAD+-consuming activity of Sterile Alpha and TIR Motifs–containing protein 1 (SARM1) in driving axonal degeneration in CIPN models (13).

Tubulin, its posttranslational modifications (PTMs), MT dynamics, and stability play critical roles in neurons via the regulation of long-distance transport, MT severing, Ca2+ homeostasis, and mitochondrial energetics (14–20). Each of these functions provides a potential mechanism that could result in the axonal degeneration in CIPN (21). Furthermore, tubulin and MTs functionally interact with Transient Receptor Potential Cation Channel Subfamily V (TRPV) members 1 and 4, two nonselective cation channels that are implicated in both sporadic and familial cases of peripheral neuropathy (22–24). Importantly, anomalies in MT dynamics and tubulin PTMs can drive axon regeneration failure and neurodegeneration (25–32), and α-tubulin acetylation is a critical component of the mammalian mechanotransduction machinery and a pathological hallmark in the neuropathy induced by the drug vincristine and genetically inherited forms of peripheral neuropathy (33, 33–36).

In addition to taxanes and vincra alkaldoids, tubulin changes have been reported downstream of the reversible 26S proteasomal subunit inhibitor bortezomib (Bort), a widely employed drug with anti-tumor activity in hematological malignancies (37). Bortezomib-induced peripheral neuropathy (BIPN) is a painful

Significance
Chemotherapy-induced peripheral neuropathy (CIPN) is a debilitating “dying back” neuropathy featuring a distal-to-proximal peripheral nerve degeneration seen in cancer patients undergoing chemotherapy. The pathogenenic mechanisms of CIPN are largely unknown. We report that in sensory neurons, the CIPN-inducing drug bortezomib caused axonopathy and disrupted mitochondria motility by increasing delta 2 tubulin (D2), the only irreversible tubulin posttranslational modification and a marker of hyper-stable microtubules. These data provide a new paradigm for the risk associated with enhanced tubulin longevity in peripheral neuropathy and suggest that targeting the enzymes regulating this tubulin modification may provide therapies that prevent the axonal injury observed in bortezomib-induced peripheral neuropathy.

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The authors declare no competing interest.

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axonally predominant and length-dependent peripheral neuropathy that affects ~40% of Bort-treated patients (3). Bort has been shown to increase MT polymer without disrupting MT ultrastructure and to affect MT-dependent axonal transport (37, 38). However, the in vivo and in vitro effects of Bort on tubulin PTMs and MT behavior as a potential BIPN pathogenic pathway have not been characterized. Furthermore, whether the perturbation of one or more tubulin PTMs may be sufficient and necessary to induce CIPN has not been determined.

Herein, we report that delta 2 tubulin (D2), an irreversible tubulin PTM residing on hyper-stable MTs, was increased in dorsal root ganglia (DRG) and sciatic nerves (SNs) from rats treated with Bort and in the sural nerve from a patient suffering from BIPN. D2 is a tubulin PTM that accumulates on very long-lived MTs and is promoted in vitro by MT stabilizers and drugs that cause CIPN, such as paclitaxel and vinca alkaloids (39, 40). The function of D2 remains unknown, although it is abundant in neurons as well as in long-lasting MT conformations in cilia. D2 is produced by the sequential action of a tubulin carboxypeptidase (VASH1/2) that cleaves the last residue of tyrosine from an α-tubulin subunit residing on stable MTs followed by the irreversible cleavage of a residue of glutamic acid by carboxypeptidase 1 (CCP1), a member of a family of tubulin de-glutamylases that is also expressed in neurons (41–44). Tubulin tyrosine ligase (TTL) is a rate-limiting enzyme that re-tyrosinates tubulin and so controls the level of de-tyrosinated tubulin, a reversible short-lived D2 precursor (45). Interestingly, constitutive suppression of TTL in mice resulted in perinatal death linked to a disorganization of CNS neuronal networks, underscoring a critical role for de-tyrosination and D2 in neuronal development (46, 47). A pathogenic role for D2 in neurodegeneration, however, has never been demonstrated. We found that Bort affects MT dynamics in vitro and in vivo and that D2 accumulation in sensory neurons may result from Bort-induced MT stabilization and increased tubulin heterodimer stability. Importantly, D2 was sufficient and necessary for both the axonopathy and disrupted mitochondria motility caused by Bort. These findings reveal a previously unrecognized role for the disruption of the tubulin tyrosination/de-tyrosination cycle in the onset of axonal injury that may occur through D2-dependent regulation of mitochondria transport.

**Results**

**Bortezomib Induces D2 in BIPN.** We measured tubulin PTM levels in the cell bodies of DRG and SNs isolated from rats that received acute and chronic administrations of Bort or untreated control (Fig. 1 and SI Appendix, Figs. S1–S3). In all cases, neuropathy was tested by mechanical and thermal nociception (SI Appendix, Fig. S1 A and B). Notably, however, only the chronic regimen appeared to significantly affect behavior and nerve conduction, and the changes in conduction velocity and evoked response amplitude were more severe in the caudal than in the digital nerve (SI Appendix, Fig. S1B). We quantified the fate of selected tubulin PTMs (de-tyrosinated, polyglutamylated, acetylated, and D2 tubulin) in DRG and SNs isolated from chronically treated rats and found that they were not significantly affected, with the exception of de-tyrosinated and D2 tubulins which appeared to be slightly, yet significantly, reduced in DRG (Fig. 1C and SI Appendix, Fig. S3). When we measured the fate of the same tubulin PTMs in tissues isolated from acutely treated rats prior to manifested neuronal injury or neuropathic behavior, however, we found that D2 was significantly increased in SNs and almost fivefold higher in the DRG cell bodies of Bort-treated rats than in controls (Fig. 1A and B and SI Appendix, Fig. S2). Furthermore, when classified by the two DRG neuron diameter markers NF200 and peripherin (48, 49), quantification of fluorescence intensity within the cell bodies revealed that D2 accumulation was higher within small-caliber DRG neurons labeled by peripherin (Fig. 1 D and E), phenocopying the cell specificity of the damage to unmyelinated fibers observed in BIPN patients (50–52).

We tested whether D2 accumulation was promoted by Bort in human BIPN by examining an available sample from a sural nerve biopsy of a cancer patient affected by the disorder. D2 levels were increased in tissue sections and whole tissue lysates of the sural nerve biopsy from the patient with BIPN but not in three control patients suffering from unrelated neuropathies, suggesting conservation and specificity of the pathways that induce D2 by Bort (Fig. 1 F–H).

We investigated whether D2 increase could be detected in vitro upon exposure of sensory neurons to pathogenic doses of Bort and analyzed how D2 levels correlated with axonal injury. To this end, adult mouse DRG neurons were grown in culture for 12 d, to allow for full recovery from axotomy and axonal extension. The cultures were exposed to Bort for up to 72 h prior to quantifying relative D2 levels and axonal degeneration in individual axons (Fig. 2 A–C). D2 was also detected by Western blot analysis of whole-cell (Fig. 2 D and E) and fractionated DRG lysates to evaluate total and partitioned D2 present in soluble and polymerized tubulin fractions (Fig. 2 F and G). We found that Bort promoted D2 accumulation and axonal injury in a time-dependent manner with a peak at 48 h. Noticeably, a significant increase in D2 levels already occurred at 12 h at a dose that did not induce a dose-dependent cellular death up to 48 h (SI Appendix, Fig. S4).

Next, we assessed whether the increase in D2 was limited to the MT fraction or extended to the soluble pool. We found that while the bulk marker βIII-tubulin (βIII) redistributed to the MT pellet in a time-dependent fashion, D2 accumulated with a similar kinetic in the soluble tubulin fraction. We hypothesized that this might be dependent on induction of MT stabilization and selective inhibition of soluble D2 turnover caused by loss of proteasome activity (Fig. 2 F and G). To examine this possibility directly, the consequences of Bort exposure on the stability of axonal MTs and levels of TTL and CCP1, two rate-limiting enzymes implicated in D2 generation, were measured in adult DRG neurons (12 days in vitro [DIV]) (Fig. 3 A–D and SI Appendix, Fig. S5 A and B). Endogenous levels of CCP1 and TTL were only slightly affected even at 24 h of exposure, a time point at which significant D2 accumulation and axonal injury was observed (SI Appendix, Fig. S5 A and B). When we tested whether Bort could acutely perturb MT behavior, however, we observed that it promoted axonal MT stability, as measured by higher residual levels of βIII that were resistant to mild MT depolymerization induced by nocodazole in neurons treated with Bort for 6 h (Fig. 3 A and B). MT stabilization was further consistent with a time-dependent increase in acetylated and de-tyrosinated tubulins, two tubulin PTMs associated with enhanced MT longevity (Fig. 3 C and D).

We investigated whether MT stabilization resulted from acute perturbation of MT dynamics by measuring the behavior of EGFP-EB3 comets that only label the plus ends of polymerizing MTs (Fig. 3 E and F and Movies S1 and S2). Bort led to time-dependent inhibition of MT dynamics in cultured DRG neurons starting at 1 h. We tested whether acute inhibition of MT dynamics also occurred in vivo by recording MT behavior in nociceptive neurons from adult zebrafish exposed to Bort by bath immersion and third-instar *Drosophila* larvae that had been fed on Bort-containing food (Fig. 4). An acute perturbation of MT behavior was observed in DRG neurons in adult zebrafish exposed to a dose of Bort that caused reversible degeneration of peripheral nerve endings at 96 h (Fig. 4A–C). Unlike the in vitro recordings, however, Bort did not seem to dampen MT dynamics in the nerve fibers of fish skin but rather only increased growth rates at 6 h (Fig. 4 B and C and SI Appendix, Fig. S6 and Movies S3 and S4). To test MT behavior using a different model system, we set up an analogous experiment in *Drosophila* larvae fed on...
Fig. 1. Bort induces D2 levels in vivo. (A) Representative D2 and βIII IF staining of DRG cell bodies and the SN isolated from control rats or rats acutely treated with Bort (i.v. 0.2 mg/kg; 24 h). (B and C) Relative tubulin PTM levels measured by quantitative IF in randomly selected DRG cell bodies (35–60) and SN (five sections per condition) from rats treated with acute (n = 3 to 5 per group) (B) or chronic (n = 4 per group) (C) (0.2 mg/kg, 3×/week for 8 wk) doses of Bort. (D) Representative peripherin (smaller unmyelinated C-fibers), NF200 (medium and large myelinated A-β fibers), and D2 IF staining of DRG cell bodies acutely treated with Bort. (E) Relative D2 tubulin values measured in DRG cell bodies (n = 35 to 60) positive to either peripherin or NF200 from rats (n = 6) acutely treated with Bort (i.v. 0.2 mg/kg; 24 h). (F) Representative D2 and βIII IF staining of sural nerve biopsy from a patient with BIPN. (G) Ratio analysis of D2/βIII levels measured by IF of fixed tissue (three to six sections per condition) from one sural nerve biopsy of a BIPN patient versus three sural nerve biopsies from control patients. (H) Immunoblot analyses of D2 levels in whole cell lysates from one sural nerve biopsy from a patient treated with Bort displayed with a control patient. Tyr, tyrosinated tubulin. GADPH, loading control. All data in B, C, E, and G are shown as medians plus interquartile range, and statistics were analyzed by Mann–Whitney U test. *P < 0.05, **P < 0.01. Scale bars in A, D, and F, 20 μm. Red writings are P values when compared to control; black writings are P values when compared between groups.
Bort-containing food for up to 6 h at a concentration of the drug that caused significant degeneration of the dendritic arbors when Bort was administered chronically (Fig. 4 D and E). This system was adopted as an alternative to administration by bath immersion that may have an impact on the drug intracellular concentration. Similarly to cultured adult DRG neurons and unlike adult DRG nociceptive neurons in zebrafish, we found that in Drosophila nociceptive neurons, Bort reduced catastrophe, rescue/nucleation frequencies and comet density at 3 h while affecting growth rates at 6 h (Fig. 4 F and G and Movies S5 and S6). Because either inhibition of MT dynamics or increase in MT polymerization alone might lead to MT stabilization, our results indicate that administration of the drug to either cultured DRG neurons or nociceptive neurons residing in live tissue stabilizes MTs through acute perturbation of MT dynamics. Together, these results demonstrate that in sensory neurons, Bort promotes D2 accumulation in vitro, in vivo, and in human tissue while acutely modifying MT behavior both in vitro and in vivo. They further indicate that in sensory neurons, D2 accumulation may result from Bort-dependent MT stabilization and increased tubulin heterodimer stability caused by proteasome inhibition.

**D2 Accumulation Is Sufficient and Necessary to Cause Axonopathy.** To test whether D2 accumulation induces axonopathy, lentiviral delivery of two independent TTL short hairpin RNA (shRNA) sequences was carried out in untreated adult DRG neurons starting at 5 DIV, to avoid any consequences of TTL loss on axon regeneration. Silencing of TTL was sufficient to increase both D2 levels and axonal fragmentation (Fig. 5 A–D). To rule out tubulin-independent functions of TTL in causing axonopathy, D2 was expressed in untreated neurons and its effects on axonal integrity were evaluated. Wild-type (WT) α-tubulin was also tested to control for undesired effects that expression of a tubulin subunit may have on cellular homeostasis.

Overexpression of recombinant D2 but not WT α-tubulin dramatically promoted axonal degeneration in naïve DRG neurons, suggesting a pathogenic role for accumulation of this irreversible tubulin PTM in the onset of BIPN (Fig. 5 E and F). To test this, we examined whether accumulation of D2 was necessary for Bort-induced axonopathy by administering Bort to adult DRG neurons that had been depleted of CCP1 expression. Lentiviral mediated depletion of CCP1 for either 7 or 10 d promoted substantial loss of D2 with no effects on polyglutamylated tubulin levels, a tubulin PTM also implicated in neurodegeneration and negatively regulated by the tubulin de-glutamylation activity of CCP1 (25, 28, 29) (SI Appendix, Fig. S7 A–C). We elected to use the shortest silencing strategy (7 d) in order to minimize any toxic consequences of sustained CCP1 silencing on axonal integrity. Under these experimental conditions, we found no effect of reducing CCP1 levels on axonal integrity of untreated neurons (Fig. 6 A and B). We observed, however, that a fivefold increase in fragmentation caused by Bort in control cells was only twofold when CCP1 was knocked down, consistent with a pathogenic role for D2 accumulation in...
Bort-induced axonal injury (Fig. 6A and B and SI Appendix, Fig. S7 G and H).

D2 Affects Mitochondria Motility. We explored whether the rescue observed in CCP1-depleted cells correlated with the ability of D2 accumulation to interfere with mitochondria motility, a hallmark of Bort-induced neurotoxicity in DRG neurons and a common feature of axonal degeneration (38). To this end, imaging of Mito-DrRed was conducted in cultured adult DRG neurons at 5 DIV, a time point at which mitochondria are motile and can be classified into subtypes according to mobility properties (Fig. 7).

We first investigated whether inhibition of motility by Bort was selective for mitochondria by imaging the dynamics of the lysosomal marker LAMP1-Cherry in cultured DRG neurons treated with Bort for 24 h (SI Appendix, Fig. S8 A and B and Movies S7 and S8). No perturbation in lysosomal motility was observed at Bort exposures that affect mitochondria motility, indicating a specific effect on mitochondria rather than on the overall MT-dependent transport machinery.

We then tested whether D2 depletion was essential for inhibition of mitochondria motility caused by Bort. We found that acute depletion of CCP1 to an extent that affected D2 levels without significant effects on polyglutamylated tubulin levels (SI Appendix, Fig. S7 D, E, F, I, and J) ameliorated loss of mitochondria motility in Bort-treated DRG neurons, with limited selectivity for dynamic pausing mitochondria, characterized by constrained bidirectional motility, compared to anterograde (AR) or retrograde (RR) running mitochondria pools (Fig. 7 A and B and Movies S9–S12).
To investigate whether excessive D2 inhibited mitochondria motility per se, the effect of D2 or WT α-tubulin overexpression was analyzed on mitochondria motility in untreated DRG neurons. While WT α-tubulin overexpression did not significantly interfere with any class of motile mitochondria, overexpression of D2 alone inhibited both dynamic pausing and fast-running mitochondria motility in both directions (Fig. 7 C and D and Movies S13–S15).

Together, these results reveal a pathogenic role for D2 in inducing axonal degeneration in DRG neurons. They further suggest that Bort-induced D2 accumulation caused by Bort exposure may contribute to the DRG damage by a selective inhibition of mitochondria motility.

**Discussion**

In this study, we adapted the BIPN model to test the hypothesis that seemingly unrelated CIPN-causing drugs share a pathogenic mechanism by converging on alterations of the tubulin cytoskeleton. In support of this hypothesis, we discovered that acute exposure to the proteasome inhibitor Bort induced accumulation of D2, a marker of hyperstable MTs and the only known irreversible tubulin PTM, in vitro and in vivo. Importantly, an increase in D2 could be measured in a rare human sural nerve biopsy from a cancer patient affected by BIPN, and D2 accumulation was sufficient and necessary for Bort induction of axonopathy and the inhibition of mitochondria motility in DRG neurons.

The innovation and potential of our findings is multifold as we demonstrate that 1) a nontaxane and nonvinca alkaloid compound unexpectedly leads to the neurotoxic accumulation of an irreversible tubulin PTM by affecting MT dynamics, stabilization, and tubulin turnover, providing a strong support for a model of pathogenesis in which CIPN drugs share a pathogenic mechanism converging on tubulin changes; 2) selective D2 accumulation within small unmyelinated fibers, which are preferentially affected in Bort-treated cancer patients (50–52), supports the notion that D2 detection could serve as an early biomarker of BIPN; 3) D2 levels may represent a general hallmark of premature MT and tubulin longevity associated with both acute and slow-developing neurodegeneration, as an increase in glutamylated D2 has recently been reported in the stratum lucidum of a kainate-induced epileptic seizure mouse model and in the hippocampi of postmortem patients with Alzheimer’s disease (53); and 4) D2 accumulation affects both axonal viability and mitochondria motility in DRG neurons, underscoring a previously unrecognized role for the tubulin tyrosination/de-tyrosination...
cycle in the control of axonal integrity that may occur through the regulation of mitochondria transport.

We found that in DRG neurons, D2 accumulation correlates with an acute increase in MT stability and a time-dependent buildup of soluble D2. The selective accumulation of soluble D2 is consistent with the proteasome inhibition activity of Bort and the irreversible nature of D2, the half-life of which, in contrast to other tubulin PTMs, is only dependent on degradation.

We explored the mechanism of D2 accumulation on MT polymer and found that Bort acutely affected MT behavior in vitro and in vivo with no substantial alteration of the levels of two rate limiting D2-regulating enzymes, even at time points when accumulation of D2 is already detectable. Unlike cultured DRG neurons and nociceptive neurons of *Drosophila* larvae in which Bort acutely inhibited dynamic MT behavior, Bort only caused induction of MT growth rates in DRG neurons residing in the skin of adult zebrafish. Either an inhibition of MT dynamics or an increase in MT polymerization alone might lead to MT stabilization, supporting the notion that Bort stabilizes MTs through acute perturbation of MT dynamics in vitro and in vivo. The discrepancy in MT behavior observed in the zebrafish may reflect differences in the intracellular concentration of the drug and/or a consequence of the shielding effect offered by the epithelial layer, which is also affected by bath administration of the drug. We further note that because of imaging limitations imposed by in vitro and in vivo recordings, MT dynamics were measured distally in the axons of the zebrafish and proximally in the axons of cultured DRG neurons and primary dendrites of nociceptive neurons from *Drosophila* larvae.

In the current study, D2 accumulation was sufficient and necessary to induce axonopathy and affected mitochondria motility in adult DRG neurons. Our results are consistent with published observations in which ablation of TTL correlates with severe defects in neuronal development and failed regeneration of axotomized axons, indicating a harmful role for increased D2 levels in neurite extension (27, 46). The findings that Bort led to neurotoxicity and accumulation of both polymeric and soluble D2, while expression of WT α-tubulin did not significantly affect axonal integrity or mitochondria motility despite leading to mild D2 accumulation on MTs, suggests that the increase in soluble D2, rather than polymeric, may predominantly contribute to toxicity.

**Fig. 5.** D2 accumulation is sufficient to drive Bort-induced axonopathy. (A) Schematic of the enzymes involved in D2 generation. (B) Immunoblot analyses of TTL, D2, and tyrosinated tubulin levels in adult DRG neurons (12 DIV) silenced of TTL expression by lentiviral shRNA delivery (5 DIV). (C) Representative IF images of neurofilament staining in DRG neurons (12 DIV) silenced of TTL expression for 7 d by lentiviral shRNA delivery. (D) Axonal degeneration in DRG neurons treated as in C. Data are pooled from 4 to 5 experiments (6 to 16 fields per condition for each experiment). (E) Immunoblot analyses of D2 and α-tubulin (DM1A) levels in adult DRG neurons (12 DIV) expressing D2-(pLVX-EGFP- D2) or WT α-tubulin (pLVX-EGFP-α-tub) (5 DIV). GAPDH, loading control. (F) Representative IF images of neurofilament staining in DRG neurons (12 DIV) transduced with lentiviruses expressing D2-(pLVX-EGFP-D2) or WT α-tubulin (pLVX-EGFP-α-tub) for 7 d. (G) Axonal degeneration in DRG neurons treated as in F. All data in D and G are shown as medians plus interquartile range from 4 to 6 experiments (4 to 17 fields per condition for each experiment). Statistical significance was analyzed by Mann–Whitney U test. GAPDH, loading control. shNC, shRNA noncoding control. *P* < 0.05, **P* < 0.01. Scale bars in C and F, 50 μm.
neurodegeneration or cortical development, however, has not been addressed. It is possible that compensation from CCP6 and/or regulation of the tyrosination/de-tyrosination cycle may vary among different types of neurons, and that loss of D2 is not as detrimental for neuronal viability as its accumulation. Indeed, we found that acute loss of D2 in cultured adult DRG neurons did not affect neither axonal integrity nor mitochondria motility. Nevertheless, our experiments of CCP1 depletion that did not result in MT hyperglutamylation while significantly inhibiting D2 formation strongly suggest an unexpected role for Bort-induced accumulation of D2 in driving axonal degeneration and defective mitochondria motility in DRG neurons.

At present, we do not have a mechanistic explanation for the selective loss of mitochondria motility caused by D2 accumulation (refer to SI Appendix, Fig. S9 for a schematic of our model). It may be that the irreversible accumulation of a soluble pool of D2 alters the composition of tubulin available for polymerization and in defects in tubulin/MT partitioning. Indeed, subtle changes in tubulin partitioning may impact mitochondria bioenergetics through tubulin interaction with VDACs with indirect consequences on motility (18, 19, 57, 58). On the other hand, D2 may interfere with the adaptor machinery that recruits mitochondria to MT-dependent motors or mitochondria-anchoring proteins, leading to inhibition of motility (59). For example, D2 may increase intracellular Ca$^{2+}$ levels through an unknown mechanism and induce Miro-dependent dissociation of MT-dependent motors. D2 accumulation may alternatively promote mitochondria localization of endogenous myosin V or syntaphilin, resulting in mitochondria arrest. Further work is necessary to provide insight into this regulation and the cross-talk between D2 and other tubulin PTMs, such as polyglutamylation and acetylation in the control of mitochondria energetics and dynamics. It will be equally crucial to investigate whether D2 accumulation affects responses to pro-survival and pro-death signals that could play a role in CIPN axon degeneration, including regulation of TRPV and Ca$^{2+}$ channels (12, 13, 21, 23, 60, 61).

Together, our data provide a strong rationale for pursuing the potential of targeting tubulin modifying enzymes of the tyrosination/de-tyrosination cycle in drug therapies aimed at preventing and/or rescuing axonal injury observed in BIPN.

**Materials and Methods**

**Reagents.** All chemicals were obtained from Sigma-Aldrich unless otherwise noted.

**Bortezomib Administration in Rats.** Bort (LC Laboratories, Woburn, MA) was formulated in 10% Tween 80, 10% ethanol, and 80% sterile saline. The study was divided into two experimental settings. Setting 1 involved acute treatment to test the modifications induced by a single intravenous (i.v.) administration of Bort at 0.20 mg/kg given as a single dose (and then killed after 24 h). Setting 2 involved chronic treatment to evaluate the neurotoxicity induced by Bort injected at 0.20 mg/kg i.v. three times per week for 8 wk as previously described (37). For each setting we analyzed untreated animals (the control, or Contr) and Bort-treated groups. The neurophysiological and behavioral tests were performed before the beginning of the treatment period (baseline value) and either 24 h after a single dose of Bort injection (setting 1) or at the end of 8 wk of Bort treatment (setting 2). All experiments were carried out between 9:00 AM and 4:00 PM.

**Primary Antibodies Used for Western Blot or Immunofluorescence Analysis.**

- De-tyrosinated tubulin (rabbit, 1:200 immunofluorescence (IF); 1:2,000 Western blot (WB), Millipore), tyrosinated tubulin (rat, 1:400 IF, 1:2,000 WB, Millipore), acetylated tubulin (mouse, 1:300 IF, 1:2,000 WB, Sigma Aldrich), D2 (rabbit, 1:400 IF, 1:1,000 WB, Millipore), polyglutamyalted tubulin (mouse G5T335, 1:200 IF or rabbit polyE, 1:2,000 WB, AdipoGen), jill tubulin (mouse and rabbit, 1:500 IF, 12,000 WB; Abcam), α-tubulin (mouse DM1A, 12,000 WB, Sigma Aldrich), CCP1 (rabbit, 1:1,000 WB, Proteintech), T7L (rat, 1:1,000 WB, Proteintech), GADPH (mouse and rabbit, 1:5,000 WB, Thermo Fisher and Sigma-Aldrich), peripherin (mouse, 1:200 IF, Abcam), and neurofilament (NF-200 chicken on tissue and mouse on DRG neurons, 1:200 IF, 

Partial CCP1 depletion, to levels that rescue axonal degeneration and cause no detectable changes in polyglutamylated tubulin, antagonized the loss of mitochondria motility induced by Bort. Conversely, D2 overexpression that caused axonal degeneration also promoted inhibition of mitochondria motility. Our findings are consistent with reports of a primary role of loss of mitochondria homeostasis in the disruption of axonal integrity and in the onset of CIPN (54). Interestingly, a link between neurodegeneration and MT hyperglutamylation by complete CCP1 loss of function has been recently reported in cerebellar atrophy observed in CCP1 knockout (KO) mouse models and infants with a defect of the gene encoding CCP1, presumably through regulation of mitochondria motility and fusion (28, 29, 55). Ablation of CCP1 expression has also been linked to the selective loss of mitochondria motility, including regulation of TRPV and Ca$^{2+}$ channels (12, 13, 21, 23, 60, 61).

Together, our data provide a strong rationale for pursuing the potential of targeting tubulin modifying enzymes of the tyrosination/de-tyrosination cycle in drug therapies aimed at preventing and/or rescuing axonal injury observed in BIPN.
Fig. 7. D2 accumulation is sufficient and necessary to drive defects in mitochondria motility caused by Bort. (A and C) Representative kymographs of mitochondria motility in DRG neurons infected with shCCP1 and shNC lentivirus (A) or pLVX-EGFP-D2 and WT-α-tubulin lentivirus (C) prior to infection with Mito-DSRed lentivirus and Bort treatment (A). Videos, 10 s/frame for 30 min. (B and D) Quantification of mitochondria dynamic state density in neurons treated as in A and C and expressed as a fraction of total mitochondria density. Dynamic Pause (DP); Stationary (ST); Anterograde Running (AR); Retrograde Running (RR). Data shown in B and D are from 9 to 19 (B) or 8 to 12 neurites (D) pooled from three independent experiments. All data in B and D are shown as medians plus interquartile range, and statistics were analyzed by Mann–Whitney U test. **P < 0.01, ***P < 0.001, and ****P < 0.0001.

DNA or shRNA. Production of lentiviral particles was conducted using the second-generation packaging system as previously described (63). In brief, HEK293T cells were co-transfected with lentiviral plasmid DNA or shRNA and the packaging vectors pLP1, pLP2, and pLP-VSV-G (Thermo Fisher) using the calcium phosphate transfection method. At 24, 36, and 48 h after transfection, the virus-containing supernatant was collected and the lentiviral particles concentrations (800-fold) by ultracentrifugation (100,000 × g at 4 °C for 2 h) prior to aliquoting and storage at −80 °C.

Plasmid DNA and shRNA. Lentiviral constructs to overexpress WT and D2 tubulins were generated by subcloning of WT and D2 tubulins from WT and D2 pEGFP-tubulin vectors (Clontech) into pLVX lentivector. Briefly, pEGFP-D2 tubulin was generated by a site-directed mutagenesis kit (Agilent) and confirmed by standard Sanger sequencing (Genewiz). Lentiviral WT and D2 tubulins were then generated by digesting pEGFP-WT or D2 tubulin with restriction enzymes, Afe1 and Not1 sequential digestions. Two TTL shRNA sequences were purchased from Sigma-Aldrich in pLKO.1 vectors: shTTL1 (TRCN0000019151; sequence: CCG GCT CAA AGA ACT ATG GGA AAT ACT CGA GTA TTT CCC ATT TTT TG) and shTTL2 (TRCN0000191515; sequence: CCG GCA TTC AGA AA GAG TAC TCA ACT CGA GTA TTT CCC ATA GTT CTT TGA TTG TT). The pLKO.1 vector with noncoding (NC) sequence was used as control. The CCP1 shRNA was purchased from Sigma-Aldrich in pLKO.1 vector (TRCN0000347379; sequence: GTG CCG GAG TGA AAC AGC TTA TTA TCT CCA GAG AAT CGG TTA TCT TTT).

MT Fractionation Assays. DRG neurons were treated with vehicle (dimethyl sulfoxide [DMSO]) or 100 nM Bort (LC Laboratories, Woburn, MA) up to 48 h at 37 °C. At the end of the incubation time, cells were gently washed with warm PHEM 1× buffer (60 mM Pipes, 25 mM Hepes, 10 mM ethylene glycol-bis[β-aminoethyl ether]-N,N,N′,N′-tetraacetic acid, and 4 mM MgCl2; pH 6.9) once before extraction with 80 μL PHEM 1× buffer, supplemented with...
0.05% Triton X-100, protease inhibitor mixture, and 10 μM taxol. After 5 min at 37 °C, the supernatant containing the soluble tubulin fraction was quickly collected in 20 μl 5x Laemmli buffer and 100 μl 1x Laemmli buffer was added to each tube to collect the MT pellet (32).

MT Stability Assays. DRG neurons (12 DIV) were treated with vehicle (DMSO) or 100 nM Bort up to 6 h before addition of 1 μM nocodazole for 45 min. At the end of the incubation time, DRG neurons were gently washed with warm PHEM 1x buffer once before extraction with PHEM 2x buffer, supplemented with 0.05% Triton X-100, protease inhibitor mixture, and 10 μM taxol. After 5 min at 37 °C, a matching volume of 2x fixative buffer (8% PFA and 0.2% glutaraldehyde in 1x PHEM) was added dropwise to the coverslips, and cells were incubated for another 30 min at 37 °C. DRG neurons were finally washed with PBS 1x and processed for IF labeling. All images were analyzed using ImageJ software by measuring mean jill intensity on major proximal neurites (within 100 μm from the cell body) (32).

MT Dynamics Assays in Cultured DRG Neurons. DRG neurons (6 DIV) grown on MatTek dishes were infected with EB3-EGFP lentivirus for 5 d and live-cell imaging of EB3 comets was performed in complete HBSS media (HBSS, 30 mM glucose, 1 mM MgSO₄, 4 mM NaHCO₃, and 2.5 mM Hepes; pH 7.4) using an epifluorescence microscope (Nikon Ti) with a temperature-controlled (37 °C) CO₂ incubator (2 frame for 5 min) at 0, 1, 2, 5, 10, and 15 min after addition of 100 μM taxol. After better visualization and reduce background, the mean projection of each time-lapse was subtrated from each frame. Kymographs were generated by drawing a region of 100 μm from the cell body using ImageJ or Metamorph software. Parameters describing MT dynamics were defined as follows: rescue/nucleation frequency is the number of rescue or nucleation events per micrometer squared and per min; frequency is the number of full track total duration of growth; comet density is the number of comets per micrometer squared per minute; growth length is the comet movement length in micrometers; comet lifetime is the duration of growth in seconds; and growth rate is the growth length/comet lifetime (32, 64).

MT Dynamics Assays in Zebrafish Nerve Endings. Adult zebrafish (Danio rerio) were raised in an Aquaneering fish habitat set to pH 7.5, 400- to 500-μs illumination, and 28 °C with 10% water exchanges daily. Fish used in the experiment were in a Narce mutant background and doubly heterozygous for the transgenic insertions P2 × 3.LexA.LexAop.3.BFP-GFP and P2 × 3.LexA.Lex-Aop.mCherry that we used as a cytosolic filler to visualize nerve endings. The mCherry-labeled strain was a generous gift from Alvaro Sagasti of the University of California, Los Angeles, and the EB3-GFP fish were generated in the Rolls laboratory using the gateway system and tool kit as previously published by Feng et al. (65). After the first imaging session, fish were housed in standard mating cages, one fish per cage with no divider, and their water exchanged every 24 h with fresh system water with 0.01% DMSO and 1.3 μM Bort or 0.01% DMSO for vehicle-treated control fish. Fish were mounted in 2% low-melt agarose and anesthetized with 0.012% tricaine but were not further warmed. Mitochondria and lysosomes were defined as stationary (ST) if fixed in an oscillatory movement or short oscillatory movements in either direction (approximately <5 μm), or AR and RR running if moving for short or longer distances in either direction (approximately >5 μm). The number of different movement states per movie was scored and quantified on kymographs. The density of each dynamic state was reported as a percentage of total density to normalize for fluctuations in total numbers.

Statistics. Data distributions that did not pass the Shapiro–Wilk normality test are shown as medians plus interquartile range, and statistical significance was analyzed by nonparametric unpaired two-tailed Mann–Whitney U test or Kruskal–Wallis with Dunn’s multiple comparisons test. Data distributions that passed the normality test are shown as means ± SEM, and statistical significance was analyzed by parametric one-way ANOVA for multiple comparisons (GraphPad Prism 8.0.2 software). For analyses of MT growth rates, length of growth, and comet lifetimes, when comparing groups, we assumed that each EB comet measure is an independent observation. All experiments were repeated at least three times unless stated otherwise. Detailed description of n values is included in each figure legend, and individual P values for each comparison are listed in SI Appendix, Table S1.

Data Availability. All study data are included in the article and supporting information.

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