Managing treatment-related peripheral neuropathy in patients with multiple myeloma

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Abstract: Peripheral neuropathy is one of the most important complications of multiple myeloma treatment. Neurological damage can be observed at the onset of the disease, due to the effect of monoclonal protein or radicular compression, but more often is treatment related. Vinca alkaloids in the past era, and more recently, thalidomide and bortezomib are mainly responsible. Degeneration of dorsal root ganglion is common, prevalently related to angiogenesis inhibition and cytokine modulation in the case of thalidomide and inhibition of the ubiquitin proteasome system in the case of bortezomib. Sensory neuropathy and neuropathic pain are more common; motor neuropathy and autonomic damage are less frequently observed. Neurotoxicity often affects patient’s quality of life and requires dose modification or withdrawal of therapy, with a possible effect on the overall response. A prompt recognition of predisposing factors (such as diabetes mellitus, alcohol abuse, vitamin deficiencies, or viral infections) and appearance of signs and symptoms, through a periodic neurological assessment with appropriate scales, is extremely important. Effective management of treatment at the emergence of peripheral neuropathy can minimize the incidence and severity of this complication and preserve therapeutic efficacy. Dose adjustment could be necessary during treatment; moreover, gabapentin or pregabalin, tricyclic antidepressants, serotonin and norepinephrine reuptake inhibitors, carbamazepine, and opioid-type analgesics are suggested according to the pain severity. Some authors reported that patients who develop peripheral neuropathy during their multiple myeloma treatments presented a particular gene expression profile; therefore, future studies could be helpful for a better understanding of possible biological pathways underlying neurotoxicity.

Keywords: neurotoxicity, thalidomide-induced peripheral neuropathy, bortezomib-induced peripheral neuropathy

Introduction

Over the past decade, new treatment options, such as the proteasome inhibitor (PI) bortezomib and the immunomodulatory drugs (IMiDs) thalidomide and lenalidomide, have dramatically changed the outcome of multiple myeloma (MM) patients, improving response and long-term survival. These new drugs are now the milestones of MM treatment regimens, for either newly diagnosed or relapsed/refractory patients. However, these compounds are not free from side effects. Treatment-related peripheral neuropathy (PN), defined as damage, inflammation, or degeneration of the peripheral nerves, is an important complication observed in MM patients, which often leads to reduction or withdrawal of therapy, with an impact on efficacy and response to treatment and a significant effect on patient’s quality of life.
This review focuses on the clinical signs and symptoms, diagnosis, and risk factors of PN, particularly dealing with the incidence, mechanisms, and management of thalidomide-induced peripheral neuropathy (TiPN) and bortezomib-induced peripheral neuropathy (BiPN).

**Signs and symptoms**

Patients prevalently describe numbness/tingling in hands and feet, burning pain, altered sensitivity to touch and heat, muscle weakness, skin or nail changes, and/or lack of coordination. In fact, sensory neuropathy and neuropathic pain are more common; therefore, hyperesthesia, hypoesthesia, and paresthesia are very often referred, usually in a distal stocking-and-glove distribution over the hands and feet, frequently associated with altered heat and cold sensation. Additionally, sensory PN can lead to areflexia and loss of proprioception. Symptoms or signs of motor and/or autonomic nervous system damage can also emerge, even if less frequently. Motor symptoms prevalently occur in the case of a severe sensory PN causing muscle cramps, muscle atrophy, or loss of strength in distal muscles. Rarely, orthostatic hypotension, bradycardia, constipation, or impotence can occur as signs of autonomic damage.1-6

**Diagnosis and evaluation**

Accurate evaluation of signs and symptoms is extremely important. The most widely accepted scale for a quantitative evaluation of PN is the National Cancer Institute – Common Toxicity Criteria score, Version 4.0 (Table 1).7 An alternative is the neurotoxicity subscale defined by the neuropathy-specific Functional Assessment of Cancer Therapy/Gynecologic Oncology Group, which is a questionnaire for patients undergoing systemic chemotherapy.8,9 The first assesses the extent of sensory and motor peripheral nerve damage caused by chemotherapy, while the neurotoxicity subscale evaluates sensory, motor, and hearing neuropathies and dysfunctions associated with neurotoxicity. Unfortunately, the degree of PN is very subjective and dependent on patients’ reporting, so a uniform and reliable interpretation of data is not so easy. Another evaluation method is the total neuropathy score (TNS), which is a complete assessment of symptoms, signs, ability aspects, and electrophysiology. The TNS can be reduced or purely clinical (Table 2). The TNS in the complete version includes the quantitative determination of the vibration perception threshold. The reduced TNS includes the electrophysiological examination of one sensory (sural) nerve and one motor (common peroneal) nerve. The purely clinical version is based on the evaluation of sensory, motor, and autonomic symptoms, pin and vibration sensibility, muscle strength, and deep tendon reflexes. However, electrophysiological examination is not always available and applicable.10

Recently, also the Treatment-Induced Neuropathy Assessment Scale has been evaluated.11 Regardless of the method or scale used, a regular and focused neurological examination (touch, pain, temperature, vibration, proprioception), distal muscle strength, ankle reflexes, and supine versus upright blood pressure evaluations are recommended. Nerve conduction studies (NCS) and needle electromyography may confirm the diagnosis of PN, identifying the neural structures involved, the presence of axonal degeneration or demyelination, and the severity of axonal damage. Particularly, thalidomide often causes axonal sensorimotor PN with reductions in sensory nerve action potentials (SNAPs) and distal compound motor nerve action potentials with denervation and reinnervation in distal muscles on needle electromyography. Bortezomib produces axonopathy less frequently and leads to a decrease in nerve conduction velocity only if demyelination is present. However, a discrepancy between neurophysiological assessment and clinical presentation is common because the damage to small fibers, which are prevalently involved, does not produce clear changes in the classic nerve conduction. Moreover, even if alterations in SNAPs are detected, recovery of SNAPs does not seem to correlate necessarily with clinical improvement. Recently, quantitative sensory testing has been introduced for evaluating sensation of small fibers. This method evaluates thermal (warm, cold) and pain (cold pain, heat pain) thresholds in the selected dermatomes. This test appears simple and fast, but it needs good cooperation between an examiner and a patient.12 Also, the evaluation of the quality of life in case of a chemotherapy-induced PN is fundamental. The European Organisation for Research and Treatment of Cancer developed a questionnaire for self-evaluation by patients (QLQ-CIPN20) that is currently used in international studies.13,14

**Risk factors**

Sometimes PN is caused by the disease itself and can be observed at the diagnosis, before starting treatment, in approximately one-quarter of patients. Therefore, symptoms are amplified by the therapy. Neurological symptoms related to plasma cell dyscrasias can be observed in not only MM but also amyloidosis, cryoglobulinemia, immunoglobulin M deposition, or POEMS syndrome. The possible etiologies are multiple: direct compression of nerves by plasmacytomas or bone lesions, amyloid deposition, immunoglobulin M antibodies against the myelin-associated glycoprotein,
or cytokine-mediated injury against a glycoconjugate component of nerves involved in the interactions between Schwann cells and axons. These factors can produce a small-fiber injury, segmental demyelination, or axonal degeneration. MM-associated PN is primarily sensory or sensorimotor, and symptoms are predominantly symmetric, including paresthesia, numbness, burning sensation, and weakness, often with mild intensity, rarely inactivating or life threatening.\(^7\)\(^6\)\(^-\)\(^10\)

Even if rare, it is important to investigate a neuronal damage that is typically demyelinating, peripheral, ascending, symmetrical, and affecting both sensation and motor function. This type of PN is frequently observed in POEMS syndrome, a rare entity, often misdiagnosed.\(^20\) Finally, Kosturakis et al\(^1\) recently demonstrated that patients with MM commonly present sensory and sensorimotor deficits at the onset as a result of disease-related decreases in peripheral innervation density.

In addition to the disease itself, comorbidities such as the presence of diabetes mellitus, alcohol abuse, vitamin deficiencies, or viral infections can increase PN; therefore, an accurate investigation and clinical history are mandatory at the diagnosis.

Some authors reported that the expression of genes involved in drug-induced apoptosis, mitochondrial dysfunction, and peripheral nervous system development, analyzed by gene expression profiling, seems to be significantly associated with the early onset of BiPN.\(^22\)\(^23\) In addition, single-nucleotide polymorphisms located in genes involved

| Table 1 Peripheral neuropathy according to NCI-CTC Version 4.0 |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Adverse event**               | **Grade**       | **1**           | **2**           | **3**           | **4**           |
| Peripheral motor neuropathy    | Asymptomatic, clinical or diagnostic observations only, intervention not indicated | Moderate symptoms, limiting instrumental ADL | Severe symptoms, limiting instrumental ADL | Life-threatening consequences, urgent intervention indicated | Death |
| Peripheral sensory neuropathy   | Asymptomatic, loss of deep tendon reflexes or paresthesia | Moderate symptoms, limiting instrumental ADL | Severe symptoms, limiting instrumental ADL | Life-threatening consequences, urgent intervention indicated | Death |
| Neuralgia                       | Mild pain       | Moderate pain, limiting instrumental ADL | Severe pain, limiting self-care ADL | –               | –               |

**Note:** Data from National Cancer Institute.\(^7\)

**Abbreviations:** NCI-CTC, National Cancer Institute – Common Toxicity Criteria; ADL, activities of daily living.

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<table>
<thead>
<tr>
<th>Table 2 Total neuropathy score</th>
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<tr>
<td><strong>Parameter</strong></td>
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<tr>
<td>Sensory symptoms</td>
</tr>
<tr>
<td>Motor symptoms</td>
</tr>
<tr>
<td>Number of autonomic symptoms</td>
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<tr>
<td>Pin sensibility</td>
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<tr>
<td>Vibration sensibility</td>
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<tr>
<td>Strength</td>
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<tr>
<td>Tendon reflex</td>
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<tr>
<td>Vibration sensibility (QST vibration)(^a)</td>
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<tr>
<td>Sural amplitude(^b)</td>
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<td>Peroneal amplitude(^b)</td>
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</table>

**Notes:** Score 0 indicates no PN; score 1/9 mild PN; score 10/19 moderate PN; and score > 20 severe PN. \(^a\)For the complete TNS. \(^b\)For the reduced TNS.

**Abbreviations:** PN, peripheral neuropathy; TNS, total neuropathy score; QST, quantitative sensory testing; ULN, upper limit of normal; LLN, lower limit of normal.
in cell death, DNA repair, and the development and function of the nervous system may also be associated with BiPN and TiPN, although the identified genes were distinct for each one.24

**Treatment-related PN**

Treatment-related PN depends on the drug used, the dosage, and the cumulative dose administered; it is usually symmetric and distal and dramatically affects the quality of life of patients, leading frequently to dose reduction, delay, or even discontinuation of a successful treatment. Vinca alkaloids and platinum compounds in the past era and more recently thalidomide and bortezomib mainly caused PN.

**Vinca alkaloids**

Vinca alkaloids inhibit the mitotic spindle and microtubule assembly, leading to distal axonopathy because axonal transport is microtubule dependent. Vincristine is the most neurotoxic, leading to PN in 30%–40% of treated patients. The severity is related to total cumulative dose; PN usually appears over the dose of 4 mg/m². Sensory signs appear first; distal motor weakness as well as neuropathic pain and autonomic dysfunction is not uncommon. Distal segments of the nerves are predominantly involved.25,26 Vincristine-induced PN may start with paresthesia in the fingers instead of toes but can also lead to ataxia, pain, and (distal) muscle weakness, resulting in foot drop. However, a significant proportion of patients will also develop autonomic symptoms, like urinary retention and erectile dysfunction. NCS may show a reduction in the amplitude of both sensory nerve and compound motor nerve action potentials. Signs and symptoms usually improve in months with dose reduction or discontinuation of the drug.27,28

**Platinum compounds**

PN occurs in ~30% of patients treated with cisplatinum, and 20% of them have to stop treatment. The toxicity is related to the total cumulative dose (500–600 mg/m²) but not to the dose intensity of treatment and is prevalently directed to the dorsal root ganglion (DRG) by damage of mitochondrial DNA and crosslink of DNA, leading to its alteration and consequent apoptosis. Cisplatinum typically affects sensory neurons of large diameter. The patients almost invariably present with nonpainful paresthesias in hands and feet, with loss of sense of vibration, position, and movement and reduced myotatic reflexes. Symptoms may start or progress after the end of treatment, but a gradual improvement occurs in most patients. NCS demonstrate sensory axonal damage with reduced SNAPs, while conduction velocity and motor nerve conduction remain normal.29

**Thalidomide**

TiPN is mainly a sensory axonal polyneuropathy that can also occur at the end of treatment, which is cumulative, dose dependent, and often irreversible. As usual, the clinical manifestations include bilateral and symmetrical sensory disorders, more rarely motor disorders or dysautonomia. The patients initially report stinging sensations or numbness (distal paresthesia and hyperesthesia) affecting the toes or sometimes the fingers, which may extend proximally. Trembling is very common but rarely interferes with daily activities initially. With the increase in severity, the deep vibratory sensitivity and proprioception may also cause progressive ataxia, difficulty in walking, and trembling. In NCS, TiPN reveals most frequently a length-dependent axonal neuropathy, a sign of a dying-back degeneration of sensory and motor nerves. Large myelinated neurons are mainly affected, with preservation of small fibers and limited evidence for demyelination. Degeneration of DRG has also been reported.30–34

As a sign of sensorimotor axonal neuropathy, thalidomide usually produces a 50% decrease in SNAPs, concurrently with sensory changes.31,35–38 The mechanism is not clear, but probably linked to the angiogenesis inhibition and cytokine modulation.29 Another hypothesis is the capillary damage and secondary anoxemia in nerve fibers. Additionally, thalidomide could reduce neuronal cell survival by downregulation of tumor necrosis factor alpha (TNFα), inhibition of nuclear factor kappa beta, and subsequent acceleration of neuronal cell death, even if these data are debated.39 In fact, also lenalidomide shows a potent TNFα inhibition but does not cause important PN. Moreover, thalidomide has been proven to reduce neuropathic pain in refractory reflex sympathetic dystrophy, maybe through inhibition of cytokines, including TNFα. This anti-inflammatory effect might also explain why a relevant increase in PN is not often observed when thalidomide or lenalidomide are combined with bortezomib.40,41

The incidence of TiPN reported in the literature is extremely variable, from 23% to 70%, according to the study, the population analyzed (pretreated or naïve patients), and the length of exposure and dosage;42,43 however the majority of studies report PN in one-third or half of treated patients.44,45

The risk of neurotoxicity increases in a dose-dependent manner up to a cumulative dose of 20 g. Actually, thalidomide is administered at a maximum dose of 200 mg daily and for a limited treatment duration.46 Prolonged therapy >1 year is discouraged, not only for neurotoxicity but also for resistance;
PN after 1 year of treatment reaches 75% of cases, with one-third of patients with grade 3 toxicity.\textsuperscript{47,48} In the recent years, thalidomide has become widely incorporated in front-line myeloma treatment for young patients in preparation for autologous transplantation in combination with bortezomib and high-dose dexamethasone. Also, elderly and/or frail patients are often treated with thalidomide with low-dose melphalan without apparent increase in PN compared to low-dose melphalan alone.\textsuperscript{49} Nevertheless, motor PN can complicate thalidomide treatment considering that concomitant high-dose dexamethasone can worsen muscle weakness and cause mild or moderate tremor. In addition, constipation occurs rapidly and frequently in patients with autonomic nerve fiber injury, with \textasciitilde75% of patients (all grades of toxicity).

**Bortezomib**

BiPN generally starts distally and may progress proximally; it is related to dose, schedule, and mode of administration and is usually reversible. It typically occurs within the first treatment cycles with bortezomib, reaching a plateau around cycle 5, without apparent increase in subsequent cycles. The incidence achieves a plateau at cumulative doses of 30–45 mg/m\textsuperscript{2}. From the clinical point of view, BiPN is more a sensory rather than a motor PN (pain, paresthesia, burning sensation, dysesthesia, numbness, sensory loss) affecting the feet more than the hands. Reports of cold pain, positive sensory symptoms in a stocking-and-glove distribution, proprioception changes, impaired sharpness detection, and elevated thresholds for the detection of skin warming and heat pain are common; sometimes, patients may also present a suppression/reduction in their deep tendon reflexes.\textsuperscript{38,41,50–52} Motor impairment is rare, even if the pain and stinging of the extremities result in a reduction in activity and in distal weakness in the lower limbs. In terms of dysautonomic neurotoxicity, orthostatic hypotension has been reported in \textasciitilde10% of the patients. From the electrophysiological point of view, NCS predominantly reveal low amplitude of SNAPs, in keeping with a length-dependent, sensory, axonal polyneuropathy, with predominant involvement of small fibers.\textsuperscript{53,54}

BiPN is multifactorial and is prevalently related to the proteasome inhibition (Figure 1). The ubiquitin proteasome system (UPS) is the main proteolytic extralysosomal system in both cytoplasm and the nucleus, playing a key role in transcription, cell cycle regulation, proliferation, signaling, and apoptosis. Neoplastic cells, which have a high protein turnover, are extremely dependent on these regulatory pathways; hence, inhibition of the UPS reduces survival of tumor cells. Neurons are quiescent cells but suffer from proteasome inhibition because of their high metabolic activity; UPS impairment leads to protein aggregation due to failure to remove misfolded proteins. As support for this hypothesis, impairment of the proteasome has been related to some neurodegenerative diseases, such as Alzheimer, Parkinson,

![Diagram](Figure 1 Principal mechanisms of neuronal damage induced by bortezomib: ubiquitinated protein accumulated in the cytoplasm with production of aggresomes, endoplasmic reticulum stress, mitochondria dysfunction, alterations in calcium homeostasis, dysregulation of cytokines.)
Oral ixazomib in

Huntington’s chorea, and amyotrophic lateral sclerosis.\textsuperscript{55} The inhibition of UPS by bortezomib causes accumulation of ubiquitinated proteins in the cytoplasm with the creation of autophagic/lysosomal vesicles that induce endoplasmic reticulum stress, which leads to alterations in calcium homeostasis and mitochondria impairment.\textsuperscript{56,57} In addition, mitochondria dysfunctions are observed for mitotoxicity or damaged axonal transport due to cytoskeleton alterations that cause impairment of the mitochondria trafficking.\textsuperscript{58,59} DNA damage signaling alters nuclear structure and dysregulates protein synthesis\textsuperscript{53,60-67} Finally, induction of a proinflammatory response and secretion of cytokines can also lead to neurotoxicity, and the inhibition of the activation of nuclear factor-kappa beta can block the nerve growth factor-mediated neuronal survival.\textsuperscript{68} The most damaged cells during bortezomib treatment are usually neuronal cell bodies within the DRG and their axons extending in the extremities to a lesser extent. Mitochondrial and endoplasmic reticulum damages in both Schwann and satellite cells have also been observed (Figure 1).

The overall incidence of BiPN varies from 31% to 45% for all grades.\textsuperscript{52} In the first Phase II studies for pretreated relapsed/refractory MM patients, SUMMIT (Study of Uncontrolled Multiple Myeloma Managed with Proteasome Inhibition Therapy) and CREST (Clinical Response and Efficacy Study of Bortezomib in the Treatment of Relapsing Multiple Myeloma),\textsuperscript{69,70} and in the Phase III study APEX (Assessment of Proteasome Inhibition for Extending Remissions),\textsuperscript{71,72} BiPN of all grades occurred in 35%–37% of patients receiving bortezomib 1.3 mg/m\textsuperscript{2} and in 21% of patients receiving bortezomib 1.0 mg/m\textsuperscript{2}. BiPN of grade 3 was observed in 9%–13% of patients, and BiPN of grade 4 occurred in $\leq$1% of patients. Approximately 12% of patients needed dose adjustment and 5% of patients withdrew therapy for neurotoxic effects; resolution to baseline or improvement occurred in 64%–71% of patients with a median of 47 days or 110 days, respectively. In the Intergroupe Francophone du Myélome (IFM) trial for newly diagnosed young patients treated with bortezomib in the induction regimen followed by autologous stem cell transplantation, BiPN occurred in 47% of patients, with 16% showing grade 3 or 4.\textsuperscript{73} For the same population of patients, the HOVON-65/GMMG-HD4 trial used bortezomib as induction and also maintenance treatment; BiPN was observed in 37% of patients during induction, with 24% of grades 3–4.\textsuperscript{74} In elderly patients, bortezomib in combination with melphalan and prednisone leads to 44% of overall BiPN with 13% of grades 3–4;\textsuperscript{75} 74% of PN events had either resolved (56%) or decreased (18%) in a median of 2 months. Interestingly, when bortezomib was combined with thalidomide or lenalidomide, this did not further increase the rate of treatment-induced PN. In fact, PN of grades 3–4 occurred in 10% of newly diagnosed MM patients treated with bortezomib, thalidomide, and dexamethasone\textsuperscript{76} and in 14% of relapsed/refractory MM patients treated with bortezomib, lenalidomide, and dexamethasone.\textsuperscript{77} Considering untreated MM patients ineligible for transplantation, severe PN was observed in 11% of patients receiving velcade, melphalan, prednisone and thalidomide as induction followed by velcade plus thalidomide as maintenance treatment; on the other hand severe PN was reported in 5% of patients treated with velcade, melphalan and prednisone.\textsuperscript{78} The median time to onset of BiPN is $\sim$2.3 months. BiPN usually improves or completely resolves in most patients after a median of 3 months after discontinuation of treatment. However, there are reports in which the recovery from pain and other sensory neuropathic symptoms required up to 2 years after bortezomib discontinuation.\textsuperscript{79} Motor or autonomic PN rarely affects patients who receive bortezomib, but diarrhea or constipation and orthostasis can complicate therapy in 10%–15% of patients who receive bortezomib.

It is important to note that once BiPN has resolved, there is no increased risk for cumulative BiPN upon retreatment with bortezomib.\textsuperscript{80}

New generation PIs have been developed in an effort to overcome resistance to bortezomib and side effects. The incidence of PN with carfilzomib is $\sim$13%, including patients with baseline PN.\textsuperscript{66,81,82} Oral ixazomib in Phase II studies showed grades 3–4 PN in less than 6% of patients.\textsuperscript{83,84}

**Lenalidomide**

Lenalidomide is a second-generation IMiD with more potent anti-inflammatory and antiangiogenic activities than thalidomide, first approved in combination with dexamethasone for relapsed/refractory MM patients who had received at least one prior therapy. Recently, the US Food and Drug Administration extended the approval also as the first-line treatment. This drug downregulates proinflammatory cytokines, upregulates anti-inflammatory cytokines, and reduces cell surface adhesion molecules. In addition, it presents antiangiogenic and antiproliferative activities. Moreover, lenalidomide binds cereblon, a protein encoded by the \textit{CRBN} gene, which forms an E3 ubiquitin ligase complex involved in proteolysis of specific proteins. Particularly, lenalidomide potentiates the proteolysis of Ikaros family zinc finger proteins 1 and 3 (IKZF1 and IKZF3), important transcription factors for B-cell differentiation, and downregulation of interferon-regulatory factor 4 and c-Myc, inducing...
cytotoxicity of myeloma cells.\textsuperscript{35,36} Lenalidomide does not seem to cause substantial neurotoxicity. PN of grades 1–2 has been observed in 18%–24% cases, considering also that the majority of these patients had a prior history of PN.\textsuperscript{87–89} Rare cases of central neurotoxicity with IMiDs have been reported.\textsuperscript{90} New IMiD pomalidomide causes very few incidences of PN; <5%.\textsuperscript{91,92}

### Drugs modification and intervention

Prompt recognition of signs and symptoms and early intervention are extremely important. Particularly, TiPN symptoms can often aggravate and become irreversible. Since the occurrence of TiPN is not predictable and a preventive treatment still does not exist, it is important to minimize other potential risk factors for PN such as vitamin B12 deficiency or diabetes. Once treatment is initiated, it has been suggested that thalidomide has to be limited to 200 mg/d to minimize TiPN and should be dose reduced or discontinued in patients with grade 2 or 3 TiPN, respectively. After reduction, thalidomide can be restarted with a 50% dose reduction upon resolution to grade 1 if the risk–benefit ratio is favorable (Table 3). In some cases, also patients with grade 1 TiPN can benefit from a thalidomide dose reduction by 50%. During maintenance treatment, the thalidomide dose could be reduced to 50 mg/d as soon as the patient has achieved a plateau response; prolonged use >12 months is not recommended.\textsuperscript{41,93} For elderly patients (>75 years), a daily thalidomide dose of 100 mg can be considered, with dose reduction to 50 mg/d in the case of PN.\textsuperscript{94,95}

In the case of BiPN, lower doses of bortezomib, weekly administration, or different schedules (4-week cycles instead of 3-week cycles) may be used.\textsuperscript{86} In fact, bortezomib should be progressively reduced from 1.3 mg/m\textsuperscript{2} to 1.0 mg/m\textsuperscript{2} up to 0.7 mg/m\textsuperscript{2} in the case of severe or persistent damage (Table 4). Moreover, once-weekly instead of twice-weekly application can be performed. Recently, a multiagent study has shown that, in addition to dose reduction, weekly dosing may prevent the progression of PN and reduce severity without affecting efficacy.\textsuperscript{97} The Gruppo Italiano Malattie EMatologiche dell’Adulto (GIMEMA) showed a significantly lower incidence of PN with weekly versus biweekly bortezomib: an incidence of grades 3–4 PN of 8% in the once-weekly group and 28% in the twice-weekly group was observed; 5% of patients in the once-weekly group and 15% in the twice-weekly group discontinued therapy because of PN. The overlapping results in terms of overall survival and progression-free survival between the two arms were reported.\textsuperscript{98} Finally, subcutaneous (SC) rather than intravenous (IV) administration is actually recommended. In a recent trial, 222 relapsed MM patients were randomized to SC or IV administration of bortezomib. The overall response rates after four cycles were identical in both arms, and no significant difference was observed in the time to overall survival and progression-free survival. BiPN grades 1–4 (38% vs 53%), grade 2 (24% vs 41%), and grade 3 (6% vs 16%) were significantly lower with SC versus IV administration of bortezomib.\textsuperscript{99} Also other groups reported less neurotoxicity with SC administration.\textsuperscript{100–103}

As supportive care, vitamin B, antioxidants such as vitamin E, alpha-lipoic acid, glutathione, and supplements with glutamine or acetyl-L-carnitine can be used. However, administration of pyridoxine (vitamin B6) can cause additional sensory neuropathy in patients with impaired renal function and in association with a protein-deficient diet; vitamin C may interfere with bortezomib metabolism and may also abrogate bortezomib-mediated inhibition of proteasome activity. Therefore, careful attention is recommended, especially during the days of bortezomib administration. The drugs to alleviate BiPN include gabapentin or pregabalin; tricyclic antidepressants such as amitriptyline, nortriptyline, and imipramine; serotonin and norepinephrine reuptake inhibitors; carbamazepine; and opioid-type analgesics. The local application of lidocaine or menthol-containing analgesic cream can temporarily alleviate bortezomib-induced pain.

Some authors propose first to use pregabalin (150–600 mg/d) for at least 3 months or gabapentin (300–2,400 mg/d).

### Table 3 Algorithm for the management of TiPN

<table>
<thead>
<tr>
<th>PN grade</th>
<th>TiPN management</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>Thalidomide dose reduction by 50%</td>
</tr>
<tr>
<td>2</td>
<td>Thalidomide discontinuation until resolution or improvement to grade 1</td>
</tr>
<tr>
<td>3–4</td>
<td>Restart with a 50% dose reduction</td>
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<tr>
<td></td>
<td>Definitive discontinuation</td>
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</table>

**Abbreviations:** TiPN, thalidomide-induced peripheral neuropathy; PN, peripheral neuropathy.

### Table 4 Algorithm for the management of BiPN

<table>
<thead>
<tr>
<th>PN grade</th>
<th>BiPN management</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>Twice-weekly administration: dose reduction to level –1 or change to weekly administration</td>
</tr>
<tr>
<td></td>
<td>Weekly administration: dose reduction to level –1</td>
</tr>
<tr>
<td>1 with pain or 2</td>
<td>Temporary discontinuation of bortezomib or further dose reduction to level –2</td>
</tr>
<tr>
<td>3–4</td>
<td>Definitive discontinuation</td>
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</tbody>
</table>

**Note:** Dose reduction of bortezomib: standard dose, 1.3 mg/m\textsuperscript{2}; reduction to level –1, 1 mg/m\textsuperscript{2}; and reduction to level –2, 0.7 mg/m\textsuperscript{2}.

**Abbreviations:** BiPN, bortezomib-induced peripheral neuropathy; PN, peripheral neuropathy.
In case of failure, duloxetine (30–60 mg/d) is considered as a valid second-line choice. Tramadol can reduce chronic pain.\(^\text{18}\)

However, the recommendations from the British Committee for Standards in Haematology in MM are

- oral tramadol for mild–moderate pain;
- oral oxycodone or alternatively morphine for chronic moderate–severe pain; and
- SC opioid therapy (oxycodone or morphine injection) for acute severe pain.\(^\text{104,105}\)

Moreover, patients should be advised to:

- wear loose-fitting shoes, roomy socks, and padded slippers;
- soak feet in icy water and massage the feet for temporary pain relief;
- keep feet uncovered in bed because pressing on the toes can add to the problem; and
- walk to help circulation in the feet, even if too much walking or standing can make the problem worse.

Patients suffering from severe PN and having difficulty in the execution of daily activities can benefit from physical exercise and physiotherapy. Adequate fiber and fluid intake, stool softeners, and laxatives are recommended for the prevention of treatment-induced constipation (and also to avoid side effects of painkillers). Patients with autonomic dysfunctions should rise cautiously, avoid demanding physical tasks, and drive vehicles or operate machinery with caution; hydration, adjustment of antihypertensive medications, increased salt intake, and eventually low-dose mineralocorticoids can be helpful for orthostasis.\(^\text{28,29,38,41,105}\)

More recently, menthol, a topical transient receptor potential melastatin 8 receptor activator, seems to be efficient for BiPN.\(^\text{106}\)

**Conclusion**

Although the introduction of new drugs in the standard of care in MM patients leads to a significant improvement in their outcome, neurotoxicity is an important side effect, particularly related to the use of bortezomib and thalidomide causing BiPN and TiPN, respectively. Patients, who sometimes present PN at baseline, related to the disease, are often forced to reduce or stop treatment because of the development of important neurotoxicity. Even if they are able to complete the planned treatment, they can suffer from signs and symptoms that significantly affect their quality of life.

A close monitoring of predisposing factors at baseline and regular neurological assessment with scales for screening and grading of PN, such as the National Cancer Institute – Common Toxicity Criteria or the TNS, are extremely important. At the appearance of signs and symptoms, modification of dosage and/or frequency of administration are recommended, together with concomitant treatments to relieve pain. Second-generation PIs and IMiDs, such as carfilzomib and pomalidomide, revealed good efficacy and lower neurotoxicity, representing a promising option for the treatment of MM patients.

**Author contributions**

SG, LC, and MTP acquired, analyzed, and interpreted the data and drafted the article. SG and MTP revised it and approved the final version. The questions related to the accuracy or integrity of any part of the work were appropriately investigated and resolved by SG, LC, and MTP. All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


