

POSITION PAPER

Bisphosphonate guidelines for treatment and prevention of myeloma bone disease

Oi Lin Lee,¹ Noemi Horvath,^{1,2} Cindy Lee,^{2,3} Doug Joshua,^{2,4,5} Joy Ho,^{2,4,5} Jeff Szer ,^{2,6,7} Hang Quach,^{2,6,8} Andrew Spencer,^{2,6,9} Simon Harrison,^{2,6,10} Peter Mollee,^{2,11,12} Andrew W. Roberts,^{2,6,7,13} Dipti Talaulikar,^{2,14,15} Ross Brown,^{2,4} Bradley Augustson,^{2,16} Silvia Ling,^{2,17,18} Wilfrid Jaksic,^{2,3} John Gibson,^{2,4,5} Anna Kalff,^{2,9} Anna Johnston,^{2,19,20} Akash Kalro,^{2,21} Chris Ward,^{2,5,22} H. Miles Prince^{2,10,6} and Andrew Zannettino ^{2,23,24}

¹Department of Haematology, Royal Adelaide Hospital, ³Department of Haematology, Queen Elizabeth Hospital, ²³Adelaide Medical School, Faculty of Health and Medical Sciences, University of Adelaide, and ²⁴Cancer Theme, South Australian Health and Medical Research Institute, Adelaide, South Australia, ²Medical and Scientific Advisory Group, Myeloma Australia, ⁶Faculty of Medicine, Dentistry and Health Sciences, University of Melbourne, ⁷Department of Clinical Haematology and BMT, Royal Melbourne Hospital, ⁸Department of Haematology, St Vincent's Hospital, ⁹Department of Haematology, The Alfred Hospital, ¹⁰Department of Haematology, Peter MacCallum Cancer Centre, and ¹³Walter and Eliza Hall Institute of Medical Research, Melbourne, Victoria, ⁴Department of Haematology, Royal Prince Alfred Hospital, ⁵Department of Cancer and Haematology, Sydney Medical School, University of Sydney, ¹⁷Department of Haematology, Liverpool Hospital, ¹⁸South Western Sydney Clinical School, University of New South Wales, and ²²Department of Medicine, Royal North Shore Hospital, Sydney, New South Wales, ¹¹Department of Haematology, Princess Alexandra Hospital, and ¹²School of Medicine, University of Queensland, Brisbane, Queensland, ¹⁴Department of Haematology, Canberra Hospital, and ¹⁵College of Medicine, Biology and Environment, Australian National University, Canberra, Australian Capital Territory, ¹⁶Department of Haematology, Sir Charles Gairdner Hospital, Perth, Western Australia, ¹⁹Department of Haematology, Royal Hobart Hospital, and ²⁰Faculty of Health, University of Tasmania, Hobart, Tasmania and ²¹Department of Haematology, Royal Darwin Hospital, Darwin, Northern Territory, Australia

Key words

myeloma, bisphosphonate, skeletal-related event (SRE), osteolysis, osteoblast, osteoclast.

Correspondence

Andrew C. W. Zannettino, Adelaide Medical School, Faculty of Health and Medical Sciences, University of Adelaide, Frome Road, Adelaide, SA 5000, Australia.
 Email: andrew.zannettino@adelaide.edu.au

Received 7 February 2017; accepted 15 May 2017.

Abstract

Multiple myeloma (MM) is a haematological malignancy characterised by the clonal proliferation of plasma cells in the bone marrow. More than 80% of patients with MM display evidence of myeloma bone disease (MBD), characterised by the formation of osteolytic lesions throughout the axial and appendicular skeleton. MBD significantly increases the risk of skeletal-related events such as pathologic fracture, spinal cord compression and hypercalcaemia. MBD is the result of MM plasma cells-mediated activation of osteoclast activity and suppression of osteoblast activity. Bisphosphonates (BP), pyrophosphate analogues with high bone affinity, are the only pharmacological agents currently recommended for the treatment and prevention of MBD and remain the standard of care. Pamidronate and zoledronic acid are the most commonly used BP to treat MBD. Although generally safe, frequent high doses of BP are associated with adverse events such as renal toxicity and osteonecrosis of the jaw. As such, optimal duration and dosing of BP therapy is required in order to minimise BP-associated adverse events. The following guidelines provide currently available evidence for the adoption of a tailored approach when using BP for the management of MBD.

Introduction

Multiple myeloma (MM) is a clonal plasma cell (PC) malignancy characterised by osteolytic bone disease leading to devastating complications including debilitating pain, pathological fractures and spinal cord compression resulting in significant disability. Myeloma bone disease (MBD) is observed in more than 80% of patients during the course of their disease and severely affects their quality of life, increases morbidity and has a

significant economic impact.^{1,2} Moreover, MBD is also associated with a 30% increased risk of mortality.³

Pathophysiology of MBD

Bone health, under normal physiological conditions, is maintained by a dynamic balance between bone formation by osteoblasts and bone resorption by osteoclasts and occurs in response to physiological influences and mechanical forces. In MM, this tightly controlled process of bone formation and resorption is disrupted leading to increased osteoclast activity and decreased osteoblast activity.^{6,7} Several soluble MM PC-derived factors have

Funding: None.
 Conflict of interest: None.

been implicated in promoting bone destruction. Furthermore, factors released by bone resorption further promote MM cell growth perpetuating the vicious cycle of malignant cell expansion and bone destruction. Factors that influence osteoclast activation include the receptor activator of NF- κ B ligand (RANKL)/osteoprotegerin ratio,^{6,8–10} macrophage inhibitory protein-1 α (MIP-1 α),^{11,12} IL-6¹³ and tumour necrosis factor- α (TNF α).¹⁴ The factors that inhibit osteoblast activity include the inhibitors of the Wnt signalling pathway such as Dickkopf-1 (Dkk-1), soluble-frizzled receptor-like proteins (sFRP) and sclerostin.^{15–18}

Definition and diagnosis of MBD

MBD is traditionally diagnosed by plain radiograph-based skeletal surveys, which reveal the presence of osteolytic bone lesions or osteoporosis with compression fractures that are attributed to the underlying clonal PC disorder.^{19,20} However, a destructive bone lesion needs to be at least 1 cm and associated with a loss of at least 50% of the bone mineral content before it can be detected by plain radiograph.²¹ Whole body low dose CT (WBLD-CT), PET/CT and whole body MRI (WB-MRI) represent more sensitive imaging modalities for the detection of osteolytic lesions. However, if WB-MRI is not widely available, MRI of the spine and pelvis will detect approximately 90% of all osteolytic lesions.²²

Most existing guidelines still recommend skeletal survey by conventional radiography as the initial method for the detection of MBD. Other modalities such as WBLD-CT, MRI or PET/CT are indicated when there is a suspicion of bony disease even if conventional radiography is negative.^{20,23–25} A systematic review comparing conventional imaging with more modern techniques supports the use of WBLD-CT or MRI.²⁶ With the availability and use of more sensitive modalities, the 2014

International Myeloma Working Group (IMWG) guidelines recommend that osteoporosis or compression fracture alone without the presence of osteolytic lesion is insufficient to meet the criteria. Notably, bone densitometry or evidence of increased fluorodeoxyglucose uptake on PET without accompanying destructive bone lesions are also insufficient to meet the diagnostic criteria of MBD.²⁰ Recently, the IMWG recommended that patients with high-risk smouldering MM should be treated as symptomatic MM based on certain biomarkers of malignancy, including more than one focal lesion of at least 5 mm on MRI.²⁰ Focal lesions on MRI indicate bone marrow involvement and not actual bone destruction.²²

Recommendations (Table 1)

- 1 Skeletal survey by conventional radiology may be performed initially but WBLD-CT or PET/CT should be used to clarify ambiguous radiological findings or if suspicion of bony disease is high even with negative conventional radiological findings. (Grade A, Level I)
- 2 MBD is defined as one or more osteolytic lesions seen on conventional radiology, CT (including WBLD-CT) or CT/PET. On CT, lesions have to be ≥ 5 mm. Increased activity on PET scan without the accompanying destructive bone lesion is not sufficient. (Grade A, Level I)
- 3 In patients with smouldering MM, more than one focal lesion on MRI at least 5 mm is also an indication for treatment. (Grade A, Level I)
- 4 Osteoporosis or compression fracture alone without accompanying osteolytic bone lesion is also insufficient to meet the criteria. Neither are bone densitometry studies. (Grade A, Level II)
- 5 Bone (technetium-99) scintigraphy has no role in the diagnosis of MBD. (Grade A, Level I)

Table 1 National Health and Medical Research Council grades for recommendation and levels of evidence^{4,5}

Grades of recommendation	Levels of evidence
A Body of evidence can be trusted to guide practice	I Evidence obtained from a systematic review of all relevant randomised controlled trials
B Body of evidence can be trusted to guide practice in most situations	II Evidence obtained from at least one properly designed randomised controlled trial
C Body of evidence provides some support for recommendation(s) but care should be taken in its application	III-1 Evidence obtained from well-designed pseudo-randomised controlled trials III-2 Evidence obtained from comparative studies with concurrent controls and allocation not randomised, cohort studies, case-control studies, or interrupted time series with a control group III-3 Evidence obtained from comparative studies with historical control, two or more single arm studies, or interrupted time series without a parallel control group
D Body of evidence is weak and recommendation must be applied with caution	IV Evidence obtained from case series, either post-test or pretest/post-test

Bisphosphonates (BP)

Mechanism of action

BP are pyrophosphate analogues that bind avidly to hydroxyapatite and are incorporated into areas of active bone remodelling.²⁷ All BP share the same core phosphate-carbon-phosphate backbone but their affinity for hydroxyapatite and their potency depend on the composition of the two side chains coupled to the central carbon atom of the nucleus.²⁸ The presence of a nitrogen or amino group, as in pamidronate and zoledronic acid, renders them 100–10 000 fold more potent than the non-nitrogen containing etidronate and clodronate.²⁹ BP are taken up by osteoclasts during bone resorption and this result in reduced osteoclast recruitment, maturation and activity and induction of apoptosis.^{30,31} In addition, recent studies suggest that BP can also stimulate osteoblastic bone formation *in vitro* and *in vivo*.^{32–34} To date, BP are the only pharmacological agents currently recommended for the treatment and prevention of MBD. While agents such as the anti-RANKL antibody Denosumab are used in patients with bone metastasis in breast and prostate cancer, Denosumab is still undergoing clinical trials in MM patients.^{35,36}

Evidence for using BP in MM

Oral clodronate has been shown to reduce the incidence of skeletal-related event (SRE) in patients compared to a placebo-control group.^{37,38} In patients with advanced disease and evidence of at least one osteolytic lesion, pamidronate was shown to reduce significantly SRE compared to placebo. Pamidronate-treated patients also experienced reduced bone pain.^{39,40} Zoledronic acid is at least as effective as pamidronate in reducing SRE, pain and delaying time to SRE in MM patients.^{41–43}

Comparison between BP

Currently, the two most commonly used BP in MBD are pamidronate and zoledronic acid. Zoledronic acid is the most potent BP and has demonstrated up to 180-fold potency compared to pamidronate.³⁵ A randomised, double-blind study comparing pamidronate and zoledronic acid in MM patients with lytic bone lesions and breast cancer patients with skeletal metastasis did not show any difference in terms of SRE in the MM cohort.⁴³ Although a more recent observational study suggested superiority of zoledronic acid over pamidronate in terms of both reduction of SRE and overall survival (OS), no long-term results from randomised controlled trials directly comparing these BP have been reported.⁴⁴

The Medical Research Council (MRC) of UK compared zoledronic acid with oral clodronate in symptomatic newly diagnosed MM patients. Not only did patients treated with zoledronic acid experience less SRE, they also showed increased OS and progression free survival (PFS) additional to that attributed to the effects of prevention of SRE.⁴⁵ Other BP have also been associated with improved survival; relapsed myeloma patients receiving pamidronate with second line therapy have slightly improved OS compared with the placebo-treated group.⁴⁰ These results support preclinical studies of anti-myeloma effects of BP.^{46–48} Furthermore, a Cochrane meta-analysis of 20 trials concluded that zoledronic acid improves OS when compared to placebo or etidronate but not compared to the other BP.⁴⁹

Adverse effects

Adverse effects of BP include inflammatory reactions at the site of injection, acute phase reactions like transient fever, myalgia and flu-like symptoms, hypocalcaemia, hypophosphataemia, renal impairment and osteonecrosis of the jaw (ONJ).^{50–53} Rarely, subtrochanteric and diaphyseal femoral fractures have also been reported.⁵⁴ Ocular side-effects including conjunctivitis, uveitis, episcleritis, scleritis and keratitis have also been associated with BP use. Symptoms become apparent within a few hours to days after commencement, requiring discontinuation.^{55–58} Acute phase reactions often occur after the first infusion and symptomatic treatment is normally sufficient.⁵⁹ Oral BP may also be associated with gastrointestinal side-effects like nausea, diarrhoea and abdominal pain.³⁸

Renal impairment

Intravenous BP are not metabolised but are eliminated exclusively by the kidneys.^{27,28} While acute and chronic renal impairment can occur, renal damage is dependent on drug levels in the blood. The risk is highest with high dosage and rapid infusion rates.⁴³ Renal injury may be multifactorial and may be due to glomerular, tubular or interstitial injury.^{60,61} Pamidronate is associated with acute kidney injury and nephrotic range proteinuria; this is attributed to a number of different mechanisms including collapsing focal segmental glomerulosclerosis.^{62,63} In contrast, zoledronic acid is more often associated with tubular toxicity resulting in acute tubular necrosis.⁶⁴

True incidence of BP-induced renal impairment is unknown, however an elevated baseline creatinine is a risk factor.⁶⁵ A study comparing zoledronic acid and pamidronate in patients with skeletal lesions in breast

cancer and MM found that the incidence of renal deterioration was similar in both drugs (10.7% in zoledronic acid vs 9.3% in pamidronate).⁴³ Notably, acute kidney injury from either drug may progress to renal failure requiring dialysis.⁶⁰

Renal impairment has also been rarely associated with oral clodronate especially when used simultaneously with non-steroidal anti-inflammatory drugs and as such, the manufacturers do not recommend its use in patients with severe renal impairment.⁶⁶

Osteonecrosis of the jaw

While occurring in only a minority of patients, ONJ is a potentially serious adverse effect of BP. ONJ commonly occurs following dental procedures and is characterised by exposed bone in the oral cavity with subsequent necrosis and bone death.⁶⁷

Pathogenesis of ONJ. The aetiology of ONJ remains unclear but may be due to a combination of infection, suppression of bone turnover and reduced vascularity of the bones of the maxilla and mandible. Dental infection is a well-established risk factor as infections are known to stimulate bone resorption.⁶⁸ Moreover, bacteria and neutrophils are often seen in affected tissue.^{69–71}

Suppression of bone remodelling by BP may play an important role in the pathogenesis of ONJ. This is supported by increased risks with higher potency BP like zoledronic acid in comparison with pamidronate and alendronic acid.^{69,72,73} ONJ has also been described with other anti-resorption drugs like the anti-RANKL antibody, denosumab.⁷⁴ The predisposition of the jaw to osteonecrosis has been attributed to the increased rate of remodelling in the jaw due to biomedical load resulting in microtrauma and heightened bone turnover.⁷⁵

Furthermore, osteonecrosis is classically associated with an interruption of the blood supply⁷⁶ and BP are known to have anti-angiogenic properties.^{77,78} ONJ has also been described in cancer patients treated with other anti-angiogenic agents such as bevacizumab.⁷⁹

Risk factors. Risk factors for ONJ include the potency, dosage and duration of exposure to BP.^{52,67,72,73,80} The MRC Myeloma IX study found that the risk of ONJ with the use of zoledronic acid was 3.7% after a median follow up of 23.7 months versus 0.5% with clodronate.⁸¹ In a single centre study, the median time to development of ONJ with oral BP, pamidronate and zoledronic acid was 54, 34 and 16 months respectively.⁸² Notably, the incidence increases with more prolonged exposure. Patients exposed to zoledronic acid had an incidence of

ONJ of 0.5, 1 and 1.3% at 1, 2 and 3 years respectively.⁸³ In contrast, the risk of ONJ for osteoporotic patients treated with yearly zoledronic acid was very low reflecting the low cumulative dosage of BP used.⁵⁰

Concomitant oral disease and dental procedures especially dental extractions, represent additional risk factors.^{51,67,69,72} Badros *et al.* estimated a 9 times greater risks of ONJ after a dental extraction while Durie *et al.* found that underlying dental problems such as infection or dental extraction was found in 81% of MM patients who developed ONJ.^{67,73}

Other risk factors include older age, concomitant corticosteroid use, smoking, diabetes mellitus and cyclophosphamide therapy.^{67,72,84,85} Genetic factors are also thought to contribute with single nucleotide polymorphisms found within region of the genes associated with bone turnover and collagen formation.^{86,87} Furthermore, certain metabolic bone diseases may influence the predisposition to development of ONJ with one study showing that polymorphism in the farnesyl pyrophosphate synthase gene, which encodes the protein directly inhibited by BP, resulted in a positive correlation between carrier status and ONJ.⁸⁷

Subtrochanteric and other atypical femoral fractures

An association between long term BP use and the development of atypical femoral fractures in particular subtrochanteric fractures and fractures at the femoral shaft has recently been reported.⁵⁴ Of all the femoral fractures, typically 87% occur at the proximal femur with only 3% occurring at the subtrochanteric region and 5% at the femoral shaft.⁸⁸ The pathogenesis is not completely understood but may be related to long term suppression of bone remodelling leading to accumulation of microdamage.^{89,90} Although it may occur in patients who have not been exposed to BP, 93.9% of cases of atypical femoral fractures have a history of long-term BP use mostly for osteoporosis but a minority for malignancy.⁹¹ The majority of patients report prodromal symptoms such as groin or thigh pain before diagnosis, hence clinicians should be aware and recognise the signs of atypical femoral fractures.⁹¹

Guidelines for the use of BP

Which patients to start on BP?

Most international guidelines recommend starting BP on all symptomatic MM patients requiring chemotherapy including patients with no visible bone lesions on conventional radiology (Table 2).

Table 2 Summary of current international BP guidelines for myeloma

Clinical scenario	ASCO	BCSH	EMN	ESMO	IMWG	Mayo clinic	NCCN
Patient selection	<ul style="list-style-type: none"> • Lytic lesions on plain X-rays • Compression fractures of spine from osteopenia • Osteopenia on plain Xray or BMD 	<ul style="list-style-type: none"> • Symptomatic patients requiring treatment whether or not bone lesions evident 	<ul style="list-style-type: none"> • Patients requiring chemotherapy • Severe osteopenia/osteoporosis • Osteolytic lesions/pathological fractures 	<ul style="list-style-type: none"> • Stage III (Salmon-Durie) • Relapsed patients receiving chemotherapy 	<ul style="list-style-type: none"> • Patients receiving anti-myeloma therapy • Patients with osteoporosis/osteopenia resulting from myeloma 	<ul style="list-style-type: none"> • Lytic lesion on plain X-ray • Osteoporosis/osteopenia on BMD 	<ul style="list-style-type: none"> • Patients receiving primary myeloma treatment
Choice of BP	<ul style="list-style-type: none"> • PAM over ZA 	<ul style="list-style-type: none"> • ZA over PAM 	<ul style="list-style-type: none"> • No preference PAM, ZA or clodronate 	<ul style="list-style-type: none"> • NA 	<ul style="list-style-type: none"> • ZA (first choice) PAM (second) then clodronate 	<ul style="list-style-type: none"> • PAM over ZA 	<ul style="list-style-type: none"> • No preference PAM or ZA
Duration	<ul style="list-style-type: none"> • 2 years in patients with responsive or SD. Further use at the discretion of treating physician • Resume on relapsed with SRE 	<ul style="list-style-type: none"> • Reasonable to consider stopping when patient has achieved CR or VGPR and no active bone disease • Reinstated at the time of relapse 	<ul style="list-style-type: none"> • Continue for 2 years • Administration beyond 2 years is not recommended • Alternative is to continue at a reduced dose or decreased frequency 	<ul style="list-style-type: none"> • Long term 	<ul style="list-style-type: none"> • Patients not in CR/VGPR, ongoing treatment • Patients in CR/VGPR, administer for at least 1 year and up to 2 years and then at physician's discretion 	<ul style="list-style-type: none"> • Continue for 2 years • If in remission and stable – stop if requires active treatment, frequency decreased to 3 monthly 	<ul style="list-style-type: none"> • NA
PAM infusion time	<ul style="list-style-type: none"> • 90 mg over ≥ 2 h • If CrCl < 30, 90 mg over 4–6 h 	<ul style="list-style-type: none"> • In severe renal impairment, 30 mg over 2–4 h in consultation with a renal physician 	<ul style="list-style-type: none"> • Resume on relapse • 90 mg 2–4 h 	<ul style="list-style-type: none"> • NA 	<ul style="list-style-type: none"> • NA 	<ul style="list-style-type: none"> • 90 mg over ≥ 2 h • NA 	<ul style="list-style-type: none"> • NA
Monitoring	<ul style="list-style-type: none"> • Serum Cr before each infusion • If renal deterioration without apparent cause, withhold until Cr returns to 10% of baseline 	<ul style="list-style-type: none"> • Renal function should be carefully monitored and doses reduced in line with the manufacturers' guidance 	<ul style="list-style-type: none"> • Monitor renal function • Patients with renal impairment should have CrCl, electrolytes and albuminuria monitored 	<ul style="list-style-type: none"> • NA 	<ul style="list-style-type: none"> • Monitor CrCl before each infusion • Monitor electrolytes and urinary albumin • Discontinue in patients renal problems until Cr returns to 10% baseline 	<ul style="list-style-type: none"> • NA 	<ul style="list-style-type: none"> • Monitor for renal dysfunction

Table 2 Continued

Clinical scenario	ASCO	BCSH	EMIN	ESMO	IMWG	Mayo clinic	NCCN
	<ul style="list-style-type: none"> • Monitor serum calcium, electrolytes, phosphate, magnesium • Monitor albuminuria 3–6 months by dipstick and 24-h collection if positive • No recommendation using bone markers 		<ul style="list-style-type: none"> • The use of bone markers is not recommended in SRE prediction or in optimising bisphosphonate 				

ASCO, American Society of Clinical Oncology; BCSH, British Committee for Standards in Haematology; BMD, bone mineral density; BP, bisphosphonate; Cr, creatinine; CrCl, creatinine clearance; CR, complete remission; EMIN, European Myeloma Network; ESMO, European Society for Medical Oncology; Hb, haemoglobin; IMWG, International Myeloma Working Group; NA, not applicable; NCCN, National Comprehensive Cancer Network; PAM, pamidronate; SD, stable disease; SRE, skeletal related events; VGPR, very good partial remission; ZA, zoledronic acid.

In smouldering myeloma, commencing BP has not resulted in delayed progression to symptomatic disease including development of MBD or in PFS.^{92,93}

Recommendations (Table 1)

- 1 In the light of the more recent guidelines and the fact that BP may confer a survival advantage over placebo, BP should be started on all symptomatic MM patients requiring treatment regardless of the evidence of MBD. (Grade A, Level II)
- 2 At present, there is insufficient evidence to recommend routine BP use in patients with smouldering myeloma, monoclonal gammopathy of undetermined significance or patients with isolated plasmacytoma. (Grade A, Level I)

Choice of BP

In Australia, oral clodronate and intravenous pamidronate and zoledronic acid are reimbursed by the PBS for the treatment and prevention of MBD. Based on the MRC Myeloma IX trial that showed superior OS and PFS of patients on zoledronic acid over clodronate, both the British Committee for Standards in Haematology (BCSH) and IMWG have recommended zoledronic acid.⁴⁵ In contrast, American Society of Clinical Oncology (ASCO) and the Mayo clinic favour the use of pamidronate because of the lower risk of developing ONJ and similar efficacy against SRE to zoledronic acid^{52,73,82} (see Table 2).

Oral bisphosphonates can be considered in patients who are unable to attend hospital for infusions. However, dosing recommendations have to be followed meticulously in order for it to be effective. For example, it must be taken in the morning on an empty stomach with a glass of plain water and patients should refrain from eating, drinking or taking other drugs for at least 1 h, otherwise absorption of the BP may be affected.⁶⁶

Recommendations (Table 1)

- 1 Intravenous BP are more effective than oral agents. Either pamidronate or zoledronic acid are acceptable choices for most patients. (Grade A, Level 1)
- 2 The risk of ONJ is higher with zoledronic acid. In patients with increased risk of developing ONJ, pamidronate may be preferred. (Grade B, Level II)
- 3 Oral clodronate is a reasonable option in patients who are unable to attend hospitals for infusion; however, dosing recommendations have to be followed meticulously. (Grade D)

Dosing, frequency and monitoring

Doses are recommended as in Table 3 with adjustments made for renal function. If renal function deteriorates without any other apparent causes, ASCO and IMWG guidelines recommend that BP should be withheld until it returns to within 10% of the baseline (Table 2).

Pamidronate and zoledronic acid are not recommended below CrCl <30 mL/min and clodronate is contraindicated if CrCl <10 mL/min⁹⁴ (Table 3). However, either slowed infusion rate or reduced dose for pamidronate has been proposed by ASCO and BCSH (Table 2). In a randomised trial of either 30 or 90 mg pamidronate, 30 mg is as efficacious in terms of quality of life and time to first SRE. Notably, there was also a trend towards lower risks of renal toxicity and ONJ in the 30 mg group.⁹⁶

Recommendations (Table 1)

- 1 Pamidronate and zoledronic acid are administered every 3–4 weeks. Oral clodronate to be administered daily in one or divided doses. (Grade A)
- 2 Renal function should be measured prior to each infusion (Grade A). For unexplained renal deterioration, BP should be withheld until renal function returns to within 10% of the baseline. (Grade A, Level II)
- 3 In patients with renal impairment, the dose of zoledronic acid should be adjusted as per manufacturer's

Table 3 Bisphosphonate dosing in renal insufficiency. (Adapted from MIMS online⁹⁴ and Terpos 2015⁹⁵)

Creatinine clearance (mL/min)	Recommended dose for clodronate (daily)
>80	1600 mg
50–80	1600 mg (no dose reduction)
30–50	1200 mg
10–30	800 mg
<10 or on dialysis	Not recommended
Creatinine clearance (mL/min)	Recommended dose for zoledronic acid (3–4 weekly)
>60	4 mg over 15 min
50–60	3.5 mg over 15–30 min
40–49	3.3 mg over 15–30 min
30–39	3 mg over 15–30 min
<30	Not recommended
Creatinine clearance (mL/min)	Recommended infusion time for pamidronate 90 mg (3–4 weekly)
>60	2–4 h
30–60	Reduce dose or infuse over 4–6 h
<30	Not recommended unless life-threatening hypercalcaemia

recommendation (Grade C). It may also be reasonable to decrease infusion rate to 30 min. (Grade C, expert opinion). No similar dose reduction recommendation exists for pamidronate and again, it may be reasonable to either administer it over a longer duration or reduce the dose to 30 or 60 mg. (Grade C, Level IV)

- 4 Neither pamidronate nor zoledronic acid is recommended in patients with severe renal impairment however in the case of life-threatening hypercalcaemia or significant MBD, pamidronate 30 mg over 2–4 h may be used. (Grade C, Level IV)
- 5 Serum calcium, phosphate and magnesium should be measured regularly. Patients may need calcium and vitamin D supplementation (Grade A, Level I). Calcium should be used cautiously in patients with renal impairment and should not be taken concurrently with oral BP.

Duration

There are no data to indicate optimal duration of therapy. In the MRC Myeloma IX trial, long-term follow up of patients up to 4 years on 4-weekly zoledronic acid or daily oral clodronate demonstrated low incidence of adverse events including ONJ and acute renal failure.⁹⁷ Similarly, in the Z-MARK study, patients who had already received 1–2 years of prior BP therapy received either zoledronic acid 4 mg at 4- or 12-weekly intervals based on the level of bone resorption marker, urinary N-telopeptide of type 1 collagen (uNTX). The rate of SRE, as well as adverse events, was low in this study and the authors concluded that the 12-weekly dosing schedule is safe and effective for up to 4 years.⁹⁸

There may be a role in using risk stratification of SRE to adjust scheduling of BP therapy⁹⁹ (Table 4, Fig. 1).

Table 4 Risk stratification for development of further SRE. (Adapted from Dickinson *et al.*⁹⁹)

Risk for development of SRE	
Low	<ul style="list-style-type: none"> • CR or VGPR • <4 prior bone lesions and no osteoporosis
Intermediate	<ul style="list-style-type: none"> • SD • >4 prior bone lesions or osteoporosis
High	<ul style="list-style-type: none"> • No SRE within 4 months • Risk of hypercalcaemia • Progressive disease

CR, complete remission; SD, stable disease; SRE, skeletal related events; VGPR, very good partial remission.

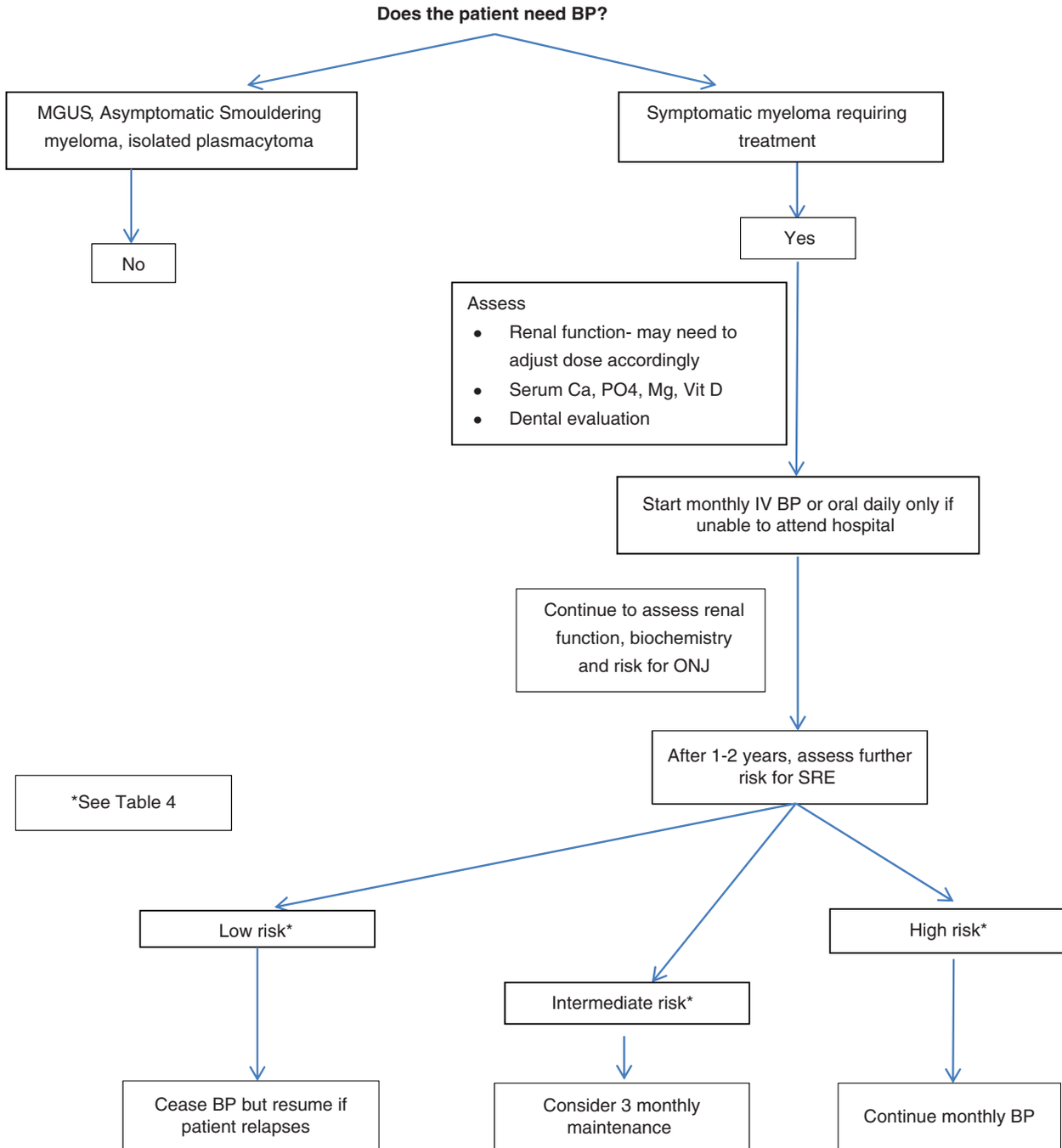


Figure 1 Proposed algorithm for initiation and continuation of bisphosphonate (BP) therapy. MGUS, monoclonal gammopathy of undetermined significance; ONJ, osteonecrosis of the jaw; SKE, skeletal-related event.

Recommendations (Table 1)

1 For patients who have achieved complete remission (CR) or very good partial remission (VGPR), monthly BP should be continued for up to 2 years. (Grade D). It can then either be stopped if risk for development of

further SRE is low or the frequency decreased to 3-monthly intervals if risk is intermediate (Grade D, Level III-3). Monthly administration should be resumed at relapsed. (Grade C, Level II)

2 For patients who do not achieve CR/VGPR but show evidence of stable disease (intermediate risk), consider

decreasing frequency to 3-monthly intervals after 2 years of therapy. (Grade D, Level III-3)

- 3 Patients with active or progressive disease (high risk), BP should continue on a monthly basis. (Grade B)

How to prevent ONJ?

Prior to initiation of BP therapy, except in cases where patients require immediate therapy, all the major guidelines recommend that patients should undergo a comprehensive dental examination and any existing dental problems be addressed.

For patients who must undergo major oral surgical procedures, there is little evidence to support withholding BP as it can stay in the bone for many years. However, the International Task Force on ONJ recommends interrupting BP therapy until soft tissue healing has occurred.¹⁰⁰ The Mayo Clinic has recommended withholding BP for at least 1 month before the procedure but the IMWG has recommended discontinuation for 90 days before and after invasive dental procedures including extractions, dental implants and surgery to the jaw.^{101,102} Prophylactic antibiotics may also be beneficial for the prevention of ONJ in patients who require invasive dental procedures.^{103,104}

Recommendations (all Grade C, Level IV, except indicated) (Table 1)

- 1 Prior to initiation of BP therapy, patients should undergo a comprehensive dental examination, address any pre-existing dental problems and optimise periodontal health.
- 2 Patients should be educated about the importance of dental hygiene and early recognition of symptoms. They should have regular dental check-up at least every 12 months.
- 3 New dental problems should be managed conservatively and dental extractions and other surgical procedures should be avoided unless absolutely necessary.
- 4 Major invasive dental procedures should be performed by an experienced oral surgeon.
- 5 Prophylactic antibiotics may be beneficial in patients undergoing invasive dental procedures.
- 6 For patients who require invasive dental procedures, BP should be withheld until soft tissue healing has occurred.
- 7 For patients who require invasive dental procedures, in the absence of data, it would seem reasonable to withhold BP 1–3 months prior to the procedure taking into account the estimated risks and benefits for the individual patient. (Grade D)

Management of established ONJ

ONJ is defined as the presence of exposed bone in the maxillofacial region that does not heal within 8 weeks after identification by a healthcare professional and should be managed by an experienced oral surgeon.^{100,101} BP should be discontinued until healing occurs.¹⁰² If BP are to be resumed, Methrotra *et al.* recommends using pamidronate instead of zoledronic acid and administering it over longer intervals, as the incidence of development of ONJ is lower with the former agent.⁸² Treatment is aimed at reducing pain, controlling soft tissue and bone infections and minimising progression of bone necrosis. The majority of patients can be managed conservatively including maintenance of optimal dental hygiene with chlorhexidine mouth washes, limited debridement and antibiotics which may result in healing in 30–60% of cases.^{105,106} In more established disease, surgical excision of necrotic bone may be necessary.^{75,100}

Recommendations (Table 1)

- 1 Treatment of ONJ should be conservative in most cases. (Grade C, Level IV)
- 2 BP should be discontinued in patients who develop ONJ ideally until soft tissue healing occurs. However, this decision should be made on a case-by-case basis depending on the risk–benefit ratio especially in patients with active MM. It may be reasonable to resume therapy when there is an improvement in bone status. (Grade C, Level IV)
- 3 For patients who developed ONJ while on zoledronic acid, it may be reasonable to change to pamidronate. (Grade D, Level II)

Utility of bone resorption markers to guide BP therapy

Bone turnover markers measure collagen breakdown products and other molecules released from osteoblasts and osteoclasts during the process of bone resorption and formation. Bone resorption markers have been used as tools to evaluate the extent of bone disease, predict risk of SRE and also response to therapy.¹⁰⁷

BP results in a rapid decrease in bone resorption markers^{43,108,109} and there has been interest in using bone turnover markers to decide when to cease BP therapy or decide on the interval between doses. A small retrospective study found as expected serum C-terminal telopeptide of Type 1 collagen (CTX) was significantly suppressed with BP. Notably, patients with increasing levels of CTX (although still within the reference range)

whilst on BP was predictive of progression of bone disease.¹¹⁰ However, the FLEX study did not support these findings.¹¹¹ In this study, post-menopausal women who had received 4–5 years of alendronate were further randomised to receive 5 years of alendronate or placebo. uNTX and bone specific alkaline phosphatase (b-ALP) were performed at baseline, 1 and 3 years and the authors concluded that the bone markers did not predict fracture risks. Similarly, another study which looked at MM patients receiving either monthly or 3-monthly zoledronic acid based on levels of uNTX found that it is not predictive of SRE.⁹⁸

Serum CTX has also been proposed by some oral surgeons to assess risk and guide dental treatment in patients taking BP.¹¹² They proposed that a certain ‘cut-off value’ of CTX would identify patients who were at higher risk of developing ONJ. However, patients taking BP will nearly always have low levels of CTX which can persist for many months following the discontinuation of BP therapy.^{98,110} Moreover, the majority of these patients will not develop ONJ.¹¹³

Currently, the international guidelines do not recommend the use of biochemical markers of bone metabolism to monitor the use and optimization of BP nor in SRE risk prediction.

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Recommendations (all Grade B) (Table 1)

- Although there are studies that show that bone resorption markers are predictive of the risk of SRE, there is currently no recommendation to use them routinely.
- Currently, there is no role for the use of bone resorption markers in monitoring the use of BP therapy.
- There are insufficient data to support the use of bone resorption markers to determine the risk of ONJ in patients requiring invasive dental procedures.

Conclusion

MBD is common and a devastating complication of MM and contributes to increased morbidity and mortality of patients. These guidelines are based on the most recent data available and aim to clarify the role of BP in treatment and prevention of this condition. Although the use of BP in the management of MBD has demonstrated benefit, there are also long-term consequences that need to be recognised and managed appropriately. The proposed algorithm for the use of BP in MM is shown in Figure 1.

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