Multiple Myeloma: An Update

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Abstract

Multiple myeloma is a rare, largely incurable malignant disease of plasma cells. Patients usually present with hypercalcemia, renal insufficiency, anemia and/or lytic bony lesions along with a monoclonal protein in the serum and/or urine in addition to an increase in the number of clonal plasma cells in the bone marrow. Patients with myeloma live on an average for five to seven years, with their survival dependent on the presence or absence of different prognostic markers. Treatment of younger fit patients is with induction therapy consisting of steroids with one or more novel anti-myeloma agents followed by high dose melphalan and autologous stem cell transplantation, while older and less fit patients are treated with melphalan-based combination chemotherapy. Supportive care is of paramount importance and includes the use of bisphosphonates, prophylactic antibiotics, thrombosis prophylaxis and the use of hematopoietic growth factors along with the treatment of complications of disease and its therapy. As more progress is being made and deeper responses are being attained, the disease might turn into a potentially curable one in the near future.

Keywords: Myeloma; Multiple myeloma; Plasma cell myeloma; Plasma cell dyscrasia; Monoclonal gammopathy.

Introduction

Multiple myeloma (MM) is a malignant disorder characterized by uncontrolled proliferation of clonal plasma cells causing a wide variety of complications leading to organ dysfunction and eventually death. Since the report of the first case of myeloma by Solly in 1844 and the case of Thomas Alexander McBean whose urine was examined by Henry Bence Jones, there has been a steady increase in the number of reported cases of MM.1 Recently, there has been a dramatic improvement in our understanding of the pathogenesis of myeloma leading to major advances in the management of patients with MM. This review summarizes the latest advances in the diagnosis, staging, prognosis and management of MM incorporating results of major landmark trials and recent international consensus guidelines.

Epidemiology

MM accounts for 10% of hematological malignancies. It evolves from a pre-malignant condition called monoclonal gammopathy of undetermined significance (MGUS). This is found in 3% of those over the age of 50,2 and progresses to myeloma (or related disorders) at a rate of 1% per year.3 Some have an intermediate stage (smoldering or asymptomatic myeloma) which carries a transformation risk of 10% per year for the first 5 years, 3% per year for the next 5 years, and 1% per year for the subsequent 10 years (cumulative probability of 73% at 15 years).4 The median age at diagnosis is between 65 and 70 years. In Oman, there were 13 new cases of myeloma in 2009 (7 males and 6 females; all above the age of 50).5 Older age, immunosuppression and exposure to ionizing radiation, benzene, herbicides and insecticides have been reported as risk factors. The cause, however, remains unknown in most patients. The risk of developing MM is 4-folds higher in people with first degree relatives with myeloma and a small number of familial cases have been described.6

Pathogenesis of myeloma

Plasma cells arise from B-lymphocytes and normally produce immunoglobulins. Myeloma cells arise from post-germinal center plasma cells in lymph nodes and migrate back to the bone marrow (BM). All cases are preceded by MGUS.7 MM is believed to result from cytogenetic abnormalities that cause cells to undergo uncontrolled proliferation, escape apoptosis and evade the immune system. The initial cytogenetic abnormalities include translocations that lead to juxtaposition of oncogenes (like cyclin D1 on 11q13, cyclin D3 on 6q21, MMSE on 4p16.3, c-maf on 16q23 and maf-B on 20q11) to the immunoglobulin heavy chain locus on chromosome 14. As the disease progresses, the clone acquires additional cytogenetic abnormalities like del 17p13 (the p53 locus), Ras mutation, activation of NFκB (Nuclear Factor Kappa B; a transcription factor) and over-expression of BCL-2 (enables cells to escape apoptosis). These secondary changes lead to increased angiogenesis in the BM, change its microenvironment and promote further proliferation. Up to 90% of patients will have chromosomal abnormalities which strongly supports their central role in the pathogenesis (and prognosis) of myeloma.8

Pathophysiology of the clinical presentation

The malignant cells produce monoclonal immunoglobulins or light chains. The expanding clone replaces the normal BM leading...
to anemia and other cytopenias. The malignant cells also cause inhibition of osteoblasts and stimulation of osteoclasts leading to the appearance of lytic bone lesions. This is mediated through an increase in RANKL (receptor activator of NFκB ligand), which stimulates osteoclasts and a decrease in its inhibitor osteoprotegrin (OPG). MIP-1 alpha (macrophage inflammatory protein-1 alpha) and IL-6 also stimulate osteoclasts. Increased levels of DKK1 (dickkopf 1) inhibit osteoblasts. Inhibition of osteoblasts explains why alkaline phosphatase levels are not increased and why radionucleotide bone scans are not helpful in these patients. The increase in osteoclastic activity also explains the hypercalcemia seen in these patients.

Renal insufficiency in MM is mostly caused by cast nephropathy (tubular damage and obstruction resulting from binding of light chains to Tamm-Horsfall mucoproteins). Other causes include amyloidosis, light chain or heavy chain deposition disease, plasma cell infiltration and interstitial nephritis. Other potential causes include: volume depletion, hypercalcemia, hyperviscosity, nephrotoxic drugs (e.g. NSAIDs), contrast nephropathy, infections, bisphosphonates and hyperuricemia.

The production of dysfunctional immunoglobulins, suppression of uninvolved normal globulins, abnormal lymphocyte function and leukopenia all lead to immunoparesis and recurrent infections. High levels of immunoglobulins can lead to hyperviscosity (typically when serum viscosity is >5 centipoises) causing headache, blurring of vision or even coma. Accumulation of free light chains in tissues can lead to amyloidosis. Fig. 1 illustrates the pathophysiology of the different clinical manifestations in MM.

Clinical features
Presenting Features
Patients generally present with bone pain (60%), fatigue (30%), weight loss (25%), paresthesia (5%), and recurrent infections (particularly pneumonia and urinary tract infections) with streptococcus pneumonia, Hemophilus influenza and Escherichia coli being the most common causative agents. Examination may reveal fever (0.7%), hepatomegaly (4%), splenomegaly (1%) and lymphadenopathy (1%). Pulmonary and pleural involvement are rare except in advanced disease. Cardiac involvement is seen with accompanying amyloidosis. Extra-medullary plasmacytomas are seen in 7% upfront and up to 20% at relapse. Cord compression (caused by an extra-medullary plasmacytoma or bony fragments from vertebral fractures) is rare (5%). Peripheral neuropathy is also rare at initial presentation and suggests amyloidosis or POEMS syndrome.

Laboratory Findings
Anemia (mostly normocytic normochromic) is seen in 75% of patients. Peripheral blood examination shows increased background staining and rouleaux formation. Circulating plasma cells and a leuko-erythrobalstic picture are occasionally seen. Because of increased immunoglobulins, the ESR is high: >20 mm/hour in 85% and >100 in one third. Elevated creatinine is seen in 50% and hypercalcemia in 25%. Urinalysis is typically negative (Bence-Jones proteins are not detected by dipstick) unless there is amyloidosis or light chain deposition disease leading to albuminuria. Also, 24-hour urine is required for quantification of monoclonal proteins (M-proteins) or light chains.

![Figure 1: Pathophysiology of MM. PCs: plasma cells, RANKL: receptor activator of nuclear factor Kappa B ligand; OPG: osteoprotegrin; MIP-1: macrophage inflammatory protein-1 alpha; IL: inter-leukine; Igs: immunoglobulins.](image-url)
An M-protein and/or light chain in the serum and/or urine is found in 97% of the patients. These usually present as a single peak in the gamma (or beta or alpha-2) region on densitometry or as a discrete band on agarose gel electrophoresis. Immunofixation (IF) is used to confirm their presence and determine their subtype. A serum free light chain (FLC) assay is useful in detecting patients with no monoclonal bands on electrophoresis and/or IF.\textsuperscript{12} Myeloma is classified into different types depending on their immunoglobulin heavy chain and light chain as follows: IgG (52%), IgA (21%), light chain (16%), Bi-clonal (2%), and IgM (0.5%), while IgD and IgE are rare.\textsuperscript{10} Patients with light chain myeloma are more prone to develop renal failure. About 3% of patients have non-secretory myeloma.

BM aspirate and biopsy show increased numbers of clonal plasma cells. Immunophenotyping using flow-cytometry and immunohistochemistry is important in proving clonality and can also provide some prognostic information. Myeloma cells are typically CD79a, VS38c, CD138, and CD38 positive. Occasionally, they are CD20 positive; seen in patients with t(11;14). Cytogenetic studies (karyotype and FISH) can reveal one or more of the genetic changes described earlier.

**Radiographic Studies\textsuperscript{13}**

On skeletal surveys, 80% of patients will have lytic lesions, diffuse osteopenia or fractures at diagnosis. These most commonly involve areas of active hematopoiesis (vertebral bodies, skull, thoracic cage, pelvis, and proximal humeri and femora). Osteosclerotic lesions are rare. Bone scans are not helpful. Contrast-enhanced CT scans should be avoided (renal toxicity). MRI can detect diffuse and focal BM lesions in 50% of those with negative skeletal surveys. MRI is also used in patients with suspected cord compression. The use of the gadolinium-based imaging should be avoided if GFR is <30 mL/min (risk of nephrogenic systemic fibrosis). PET scans correlate with areas of active lytic lesions but their use remains investigational. Fig. 2 demonstrates some of the features seen in MM.

**Diagnosis**

Myeloma is suspected in patients presenting with back pain, other bony pains, unexplained anemia, renal insufficiency and/or hypercalcemia. The diagnostic criteria for MM,\textsuperscript{14} put emphasis on the presence of otherwise un-explained end-organ damage (CRAB: hyper-Calcemia, Renal insufficiency, Anemia and Bony lesions) in patients with M-proteins/light chains and clonal plasma cells (Table 1). The differential diagnosis includes MGUS, smoldering myeloma, solitary plasmacytomas, plasma cell leukemia (PCL), AL amyloidosis (primary amyloidosis), POEMS syndrome (Polyneuropathy, Organomegaly, Endocrinopathies excluding diabetes and hypothyroidism, Monoclonal protein and Skin changes with no increase in BM clonal plasma cells and no end-organ damage), Waldenstrom’s macroglobulinemia (lymphoplasmacytic lymphoma with IgM monoclonal-protein) and metastatic malignancies (like kidney, breast and lung cancers). PCL is an advanced phase of myeloma with a plasma cell count of >2000/μL or 20% of the peripheral blood white cells.

![Figure 2: Some of the clinical findings seen in patients with multiple myeloma. A: Rouleaux formation on the peripheral blood film. B: Increased numbers of plasma cell on a BM aspirate. C: Lateral X-ray of the skull showing multiple lytic lesions; D, E, and F: Detection of monoclonal proteins on PEP and IF (D: normal pattern of SPEP; E: abnormal monoclonal peak in the gamma region on SPEP; F: Immunofixation identifying the M-protein as IgG kappa).](image-url)
Table 1: Diagnostic criteria of myeloma and related disorders.

<table>
<thead>
<tr>
<th>Disease entity</th>
<th>M-protein</th>
<th>Clonal PCs in BM</th>
<th>End-organ damage (CRAB*)</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>MGUS</td>
<td>&lt;30 g/L</td>
<td>&lt;10%</td>
<td>No</td>
<td>No other B-cell lymphoproliferative disorder</td>
</tr>
<tr>
<td>Smoldering Myeloma</td>
<td>&gt;30 g/L in serum or &gt;1 g/24hour in the urine</td>
<td>&gt;10%</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Symptomatic Myeloma</td>
<td>Any value</td>
<td>&gt;10%</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>Solitary Plasmacytoma</td>
<td>All of the following:</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Biopsy proven solitary lesion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>No clonal plasma cells in the bone marrow</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>An otherwise normal skeletal survey and MRI of the spine and pelvis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>No end organ damage</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*CRAB:Hyper-Calcemia serum calcium >2.875 mmol/L; Renal insufficiency serum creatinine >177 umol/L; Anemia hemoglobin <10g/dL or 2g/dL below normal; Lytic Bone lesions or severe osteopenia

Table 2: Staging systems and prognostic markers in patients with multiple myeloma (survival times are according to the ISS).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Durie-Salmon Staging System</th>
<th>International Staging System (ISS)</th>
<th>Survival*</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>All of:</td>
<td>Serum albumin &gt;35 g/L Serum β2-microglobulin &lt;3.5 mg/L</td>
<td>62m</td>
</tr>
<tr>
<td></td>
<td>Hemoglobin &gt;10 g/dL</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Normal serum calcium</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Low concentration of M-protein (IgG &lt;50 g/L, IgA &lt;30 g/L, Bence-Jones protein &lt;4 g/24hours)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>No bony lesions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>II</td>
<td>Neither stage I nor stage III</td>
<td></td>
<td>44m</td>
</tr>
<tr>
<td>III</td>
<td>Any of:</td>
<td>Serum β2-microglobulin &gt;5.5 mg/L</td>
<td>29m</td>
</tr>
<tr>
<td></td>
<td>Hemoglobin &lt;8.5 g/dL</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Serum calcium &gt;3 mmol/L</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>High concentrations of M-protein (IgG&gt;70 g/L, IgA&gt;50 g/L or Bence-Jones protein &gt;12g/24 hours)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Other prognostic markers:
Cyto genetic (by conventional karyotyping and by FISH):
Good/Average: Hyperdiploidy, t(11;14)
Bad/poor: Hypodiploidy, deletion 13q on conventional cytogenetics, t(4;14), t(14;16), chromosome 1 abnormalities, 17p deletion
Others: Advanced age, poor performance status, high LDH, high CRP, low platelets, M-protein (e.g., IgD having more renal failure and amyloidosis), circulating plasma cell, BM plasma cells >50%, plasmablastic morphology, high BM plasma cell labeling index (PCLI >3%), and serum FLC ratio (<0.03 or >32 having a significantly shorter survival).

For Durie-Salmon staging, each stage is sub-classified into A and B where: (A) normal renal function (serum creatinine <177 umol/L); (B) abnormal renal function (serum creatinine >177 umol/L)

Staging and Prognosis
Staging is done to determine the burden of disease and to predict survival. There are two main staging systems: the international staging system (ISS), and the Durie-Salmon staging system (Table 2). Cyto genetic abnormalities detected by conventional karyotyping or FISH are also important in the prognosis of myeloma. These and other prognostic factors are listed in Table 2.

Management
Guidelines are now available to guide the management of patients with plasma cell disorders in the current era with many available active agents. Historically however, treatment was based on the use of the combination of steroids and alkylating agents (typically melphalan and prednisone; MP). VAD (Vincristine, Adriamycin and Dexamethasone) later became a common initial therapy in younger patients. Most of the activity in this combination was from dexamethasone and dexamethasone alone became the regimen of choice. This group (younger patients) was later shown to have a survival advantage with autologous stem cell transplantation (ASCT) and this became the new standard of care, while MP remained the gold standard in older and less fit patients.

The discovery of the anti-angiogenic and immune-modulating properties of thalidomide (a compound withdrawn in the 1950-60s for its teratogenic effects) led to it being added to steroids with a significant improvement in response. Modifications to its structure led to the formation of a new less toxic immunomodulating drug (IMiD) called lenalidomide. At the same time, with the discovery...
of the ubiquitin-proteasome pathway that is responsible for the degradation of cellular proteins, a proteasome inhibitor (bortezomib) was developed. This targets the proteasome pathway leading to cellular apoptosis (mainly via inhibition of NF-kB). Remarkable responses were shown in the SUMMIT, CREST and APEX trials. These novel agents were tested in combination with steroids and/or other chemotherapeutic agents and seem to have transformed the prognosis of multiple myeloma with overall response rate of over 80%.32

In this era of novel agents, treatment varies depending on whether the patient is eligible for ASCT or not. Age >65, poor performance status (ECOG performance status of 3-4) and significant organ dysfunction (significant liver disease, renal disease with creatinine >220 umol/L unless on stable chronic dialysis and/or NYHA class III-IV) render patients unsuitable for such intensive therapy.

Younger fit patients i.e. transplant eligible
The aim of therapy in these patients is to induce a rapid and deep response, reverse organ damage and decrease the burden of disease. This is done by the use of combinations of high dose dexamethasone with thalidomide (TD; Thalidomide+Dexamethasone),23,24 bortezomib (VD; Velcade i.e. bortezomib+Dexamethasone),25 or lenalidomide (Rd; Revlimid i.e. lenalidomide+low dose dexamethasone),26 or with more than one novel agent. VTD (velcade with thalidomide and dexamethasone),27 CTD (cyclophosphamide, thalidomide and dexamethasone),28 and CyBorD (cyclophosphamide, bortezomib and dexamethasone),29 seem to produce the highest response rates. Alkylation agents like melphalan should be avoided as these are damaging to stem cells and will make it difficult to collect stem cell for transplant later.

After 2-4 cycles of induction therapy, responding patients undergo peripheral blood stem cell mobilization and collection using high dose cyclophosphamide followed by G-CSF.30,31 Stem cells are cryopreserved in 5% dimethylsulfoxide (DMSO) and then thawed at the time of transplant; done between 2-4 weeks after successful collection (except in cases where a delayed transplant strategy is followed, which is a valid option).32 Patients typically receive high dose melphalan 200 mg/m² (140 mg/m² in renal dysfunction). Stem cells are infused through a central line 24 hours later. Two large RCTs have shown increased survival by one year with ASCT in these patients.33,34 A large RCT also showed further improvement with tandem transplantation (a planned second transplant 3-6 months following the first one).35 The benefit was, however, limited to patients not achieving at least a very good partial response (VGPR) and a meta-analysis of 6 RCTs showed increased toxicity with no significant difference in overall survival (OS).36 A subset of patients might still benefit from tandem transplantation and some may benefit from a second transplant if they relapse later. Therefore, stem cells are collected for two transplants at the time of the initial collection.

Older and less fit patients i.e. transplant ineligible
The aim of therapy in this group is disease control with the least possible toxicity. MP has been the backbone of therapy for years producing 50% response rates. A meta-analysis of 27 RCTs showed similar survival rates between MP and various combinations of chemotherapy.37 The addition of novel agents to this combination has led to increased response rates, prolongation of progression free survival (PFS) and OS. The combination of thalidomide with MP (MPT) has been shown in several large RCTs to prolong PFS and in some OS, reaching a median survival of 40 months in some of these trials.38,39 A recent meta-analysis of 6 RCTs confirmed the improvement in PFS and OS.40 The combination of bortezomib (velcade) with MP (VMP) has also been shown in the VISTA trial to improve PFS and OS compared to MP with an OS of 70% at 3 years.41 More recently, lenalidomide (revlimid) has also been combined with MP (MPR) leading to significant improvement in survival.42 These regimens are also associated with improved quality of life.43,44

MPT, VMP and MPR have not been compared head-to-head in an RCT and the choice will depend on several factors like: renal function (bortezomib causes faster reversal of renal dysfunction while the dose of lenalidomide has to be adjusted in renal insufficiency), the presence and risk of peripheral neuropathy (more with thalidomide and bortezomib), the risk of thrombosis (more with thalidomide and lenalidomide), the presence of high risk cytogenetic abnormalities (bortezomib seems to overcome some of these) and some logistical considerations (thalidomide and lenalidomide are oral while bortezomib is given intravenously twice weekly although recent trials have shown similar efficacy with less toxicity with subcutaneous,45 and once weekly,46 dosing).

Maintenance therapy
With the excellent results obtained with novel agents, it seems intuitive to use something to maintain these responses for as long as possible. Thalidomide (alone or with prednisone) has been shown to improve PFS post-ACST in young patients in six studies (improving OS in three of them) and to prolong PFS in two trials in elderly patients. A meta-analysis has shown overall improvement in PFS and a marginally better OS.47 Bortezomib maintenance has also been shown to improve outcome but with no effect on OS.48 The best results with maintenance so far are with low dose lenalidomide. Two trials of lenalidomide maintenance after transplant,50,51 and one after MPR induction (the MM-015 trial),52 showed a significant improvement in PFS with an increase in OS in one of the transplant trials.53 It should be noted that achievement of an OS benefit with maintenance therapy is difficult with an increasing number of available effective salvage treatment at relapse and improvement in PFS might be a valid end-point, especially when there is improvement in quality of life. It should also be noted that maintenance therapy can be associated with some side effects related to the medications used as well as the reported increase in second primary malignancies (SPM) in these trials. The benefit therefore, has to be weighed against these risks.
Supportive care

Adjunctive supportive care is very important in patients with MM and includes measures to treat complications of the disease and to prevent the morbidities associated with the disease and its treatment.53 These include:

- The use of bisphosphonates: Bisphosphonates (zolendronic acid or pamidronate) are used to prevent the progression of bony disease and to treat hypercalcemia.54 However, they have been associated with increased risks of renal toxicity and osteonecrosis of the jaw (ONJ). Bisphosphonates are given monthly for a maximum of two years and then stopped (or continued every 3 months in patients with ongoing active myeloma).55,56

- Thrombosis prophylaxis: Patients with MM are at higher risk of developing thromboembolism.57 Aspirin is the easiest to use and has similar efficacy when compared to other agents.58 However, thrombotic and bleeding risks should be assessed and the choice of prophylactic strategy individualized (patients with high risk for thrombosis should preferably be on LMWH or warfarin).

- Infection prophylaxis: Regular bacterial infection prophylaxis using daily trimethoprim-sulfamethoxazole (cotrimoxazole) or a fluoroquinolone (ciprofloxacin or levofloxacin) is controversial but is advisable in high-risk patients. Intravenous immunoglobulins should be considered in patients with recurrent life threatening infections. Prophylaxis against Pneumocystis carinii (PCP) is standard and herpes zoster prophylaxis (with acyclovir) is recommended, especially in patients getting bortezomib and those undergoing ASCT. Fungal prophylaxis is recommended in the latter group. Vaccination against viral influences, Streptococcus pneumonia and Haemophilus influenza is also recommended although the efficacy is not guaranteed.

Management of complications (CRAB and others)

- Hypercalcemia: Mild asymptomatic hypercalcemia (calcium <3 mmol/L) is managed by hydration and steroids while in moderate-severe cases, bisphosphonates (+/- calcitonin) are added.

- Renal insufficiency: Avoidance of nephrotoxic agents, dose-adjustments and treatment of hypercalcemia are important to prevent further damage.59 Plasmapheresis in patients with predominant cast nephropathy is controversial. When indicated, hemodialysis is used.

- Anemia: The use of erythropoietin or darbepoietin is recommended if hemoglobin is <10 g/dL. Hemoglobin should not be allowed to increase over 12 g/dL (risk of thrombosis).50 Patients with severe symptomatic anemia should also receive red cell transfusions.

- Bony lesions and bone pain: Patients should be assessed thoroughly for pain especially back pain, which may be a sign of impending cord compression. NSAIDs should be avoided (risk of renal damage). Patients with mild pain benefit from regular paracetamol while patients with moderate-severe pain will require tramadol or opioids. Focal lytic lesions causing severe pain not responding to therapy may respond to a single fraction 8-10Gy of RT. Pathologic fractures or impending fractures of long bones require stabilization with an intramedullary rod. Vertebral compression fractures may benefit from kyphoplasty or vertebroplasty.61

- Cord compression: This is treated as an emergency with RT plus high dose dexamethasone. Surgical decompression is only necessary if the neurologic deficit does not improve or in cases where the compression is due to a bony fragment or retropulsed bone.

- Peripheral neuropathy: This is present in a minority of patients at diagnosis but is a frequent complication of treatment and progressive disease. It is managed by gabapentin or pre-gabalin, treating the disease, dose-adjustment of the causative drug and treating any concomitant B12 deficiency.

- Hyperviscosity syndrome: Plasmapheresis should be done promptly to relieve symptoms if present regardless of the viscosity level and treatment for myeloma should be started immediately.

- Hyperuricemia and tumor lysis: Tumor lysis syndrome is uncommon in MM. Hydration and allopurinol (dose-adjusted in renal dysfunction) is used in patients with high disease burden.

Treatment of MGUS, Smoldering Myeloma and Plasmacytomas.

Patients with MGUS and smoldering myeloma are followed every 3-6 months without treatment unless there is progression to MM.60 The use of bisphosphonates may decrease the incidence of bony lesions at progression. Plasmacytomas are treated with radiotherapy (≥45Gy to involved fields).65

Monitoring during and after therapy

Regular monitoring during and after treatment is essential. This consists of two parts; monitoring for disease response (according to the IMWG consensus guidelines),64 and assessment of treatment related toxicity. Patients should have a monthly detailed clinical assessment, as well as blood and urine tests. Once in a stable plateau phase, tests are repeated every three months. At the same time, they are assessed for the presence of side effects, which include:

- Thalidomide: sedation, constipation, skin rash, thrombosis and peripheral neuropathy.
- Lenalidomide: thrombosis and myelosuppression.
- Bortezomib: peripheral neuropathy, diarrhea and thrombocytopenia.

Of note, IMiDs are given in patients of childbearing age only if they use two contraceptive methods. Steroid related side effect, myelosuppression from melphalan and side effects of other drugs are also assessed.
Relapsed myeloma

Almost all patients with MM will relapse. Treatment at relapse depends on age, performance status and the presence of comorbidities at relapse. Patients eligible for ASCT who relapse >2 years after initial ASCT can benefit from another ASCT.59 Lenalidomide-dexamethasone is the regimen of choice given, it yielded impressive results in the MM-009 and MM-010 studies,66,67 unless the patient has progressed on or soon after a lenalidomide containing regimen. DVD (dexamethasone, velcade and doxil; a liposomal doxorubicin with less cardiotoxicity) based on the results of the DOXIL-MMY-3001 Trial,68 or other regimens (e.g. cyclophosphamide containing regimens) could also be used. In general, the same initial regimen can be repeated if the remission was long-lasting (>1 year) and a change to a combination containing another novel agent that the patient was not exposed to is preferable in patients with short remissions. There is no standard salvage regimen and therapy is tailored according to the pace of disease and fitness of the patient. Tolerance can be improved by dose-adjustments or use of newer strategies (e.g. weekly and/or subcutaneous bortezomib).

Role of allogeneic BMT

Historically, allogeneic transplantation was associated with high rates of complications including a prohibitive rate of treatment related mortality. With the use of reduced intensity conditioning (RIC) regimens, more patients have received this modality of treatment as upfront therapy, at relapse or as tandem post-ASCT. Unfortunately, the majority of patients might not be eligible because of advanced age, comorbidities and lack of available HLA-matched donors. In addition, long-term follow-up has shown continuing relapses post transplant and graft-versus-host disease in a significant proportion of patients. It remains an option though in very young fit patients especially those with high-risk features. However, until its safety and efficacy is established, it should only be used in the context of a clinical trial.69

Novel therapies and future directions

Several new agents have shown promising results in early phase clinical trials and are being explored as single agents or in combination with other agents in clinical trials like pomalidomide,70 (a new IMiD) and carfilzomib,71 (a second generation proteasome inhibitor). With better understanding of this disease and its pathogenesis, more novel agents are being discovered. With deeper responses to the currently available agents, it is not unrealistic to hope that one day in the near future, a cure will be the goal in the majority, if not all patients.

Conclusion

MM is a plasma cell disorder with an uncontrolled expansion of clonal plasma cells associated with end-organ damage (CRAB). It is preceded by MGUS and/or smoldering myeloma. The international staging system and cytogenetics are used to risk stratify patients. Major advances have occurred over the past few years, especially with the use of IMiDs and proteasome inhibitors. Younger patients can now enjoy a median survival of 7 years (with induction therapy, preferably bortezomib-based, followed by ASCT). Older patients also have longer survivals with MPT/MPV/MPR. Maintenance therapy seems to prolong survival further but more data on long-term survival (especially with regards to SPM) is needed. Supportive care is essential and includes the prevention and treatment of complications related to the disease and its therapy. The therapeutic paradigm for patients with myeloma seems to be shifting from the current dominant view of disease control,72 using sequential therapies to prolong PFS to that of intensive use of combinations of novel therapies followed by maintenance aiming at potential cure and prolongation of OS (while maintaining good quality of life).73

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References


Oman Medical Specialty Board


