Management of bone metastases in renal cell carcinoma: bone-targeted treatments, systemic therapies, and radiotherapy

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Abstract
Metastatic renal cell carcinoma (mRCC) presents with bone metastases in around 20%-30% of patients enrolled in the most recent first-line clinical trials. Emergence of several new agents in first line, in both monotherapy and combination, has significantly improved patient outcomes. However, the activity of such agents on bone metastases is unclear and management of these patients is complex, due to potential complications that can significantly impair quality of life. This review addresses mRCC diagnosis and monitoring and summarizes the current evidence on systemic therapy, ablative therapies such as stereotactic ablative radiotherapy and surgery, and supportive therapy with bone-targeting agents for these patients, with the goal of improving their outcomes.

Keywords: Bone metastases, renal cell carcinoma, bone targeted treatments, radiotherapy, systemic therapy

INTRODUCTION
Renal cell carcinoma (RCC) is a heterogeneous disease, with clear cell RCC (ccRCC) representing about
75% of all cases. ccRCC is characterized by frequent alterations in the Von Hippel-Lindau (VHL) gene, upregulation of VEGF, activation of the phosphoinositols-3-kinase-Akt-mTOR pathway, and increased migration ability and angiogenesis[1]. Among the non-clear cell RCC, papillary RCC is the most prevalent (10%-15%), and characterized by MET alterations (Type I) or a panoply of molecular alterations (Type II)[2].

Evidence suggests that metastasis in RCC is mostly driven by hematogenous dispersion of tumor-derived microvesicles, which carry CD105-positive RCC cancer stem cells and pro-angiogenic mRNAs[3]. In VHL-mutated ccRCC, HIF-regulated CUB-domain-containing protein and MUC1 are considered to be metastasis drivers, by increasing migration[4, 5]. As in other cancers, the chemokine receptor CXCR4 is also upregulated in ccRCC and associated with increased metastatic ability[6].

Bone can be affected by metastatic disease in virtually any cancer. One pivotal discovery in cancer research was breast cancer cell tropism to bone, which led to the “seed and soil hypothesis”[7]. Bone metastases (BM), also common among patients with RCC, are present in up to one third of patients with metastatic disease [Table 1][8].

BM can lead to significant morbidity in cancer patients, with progressive autonomy loss and impaired quality of life (QoL)[9]. In metastatic RCC (mRCC), BM are typically osteolytic, compromising bone integrity and eventually leading to skeletal-related events (SREs), including pathologic fractures, impending fracture requiring surgical intervention, bone pain requiring radiotherapy, spinal cord compression, and hypercalcemia[10]. In addition to these devastating complications, BM have a negative impact on the progression-free survival (PFS) and overall survival (OS) of mRCC patients treated with systemic therapies compared to patients without BM[11]. Moreover, development of BM at first-line treatment start resulted in significantly shorter survival compared with later points in time. Despite several studies reporting that BM are associated with higher risk of morbidity and shorter survival in mRCC patients, heterogeneity of study populations has been a critical drawback[12]. Median OS after diagnosis of RCC BM ranges from 12 to 28 months[13]. Suggested reasons for poor survival outcomes in BM patients include interactions between cancer cells and tumor microenvironment, particularly bone microenvironment, with subsequent bone destruction and rapid tumor growth[14]. A retrospective analysis of 94 patients with RCC and BM suggested five prognosis-predictive risk factors in this setting: sarcomatoid differentiation of the primary tumor (P = 0.001), spinal involvement (P = 0.003), extraosseous metastases (P = 0.021), increased alkaline phosphatase levels (> 1.5 times the upper limit of normal; P = 0.0003), and increased C-reactive protein levels (> 0.3 mg/dL, P = 0.018)[15].

BM are associated with substantial morbidity, mortality, QoL deterioration, and economic burden. A recent propensity score matching analysis revealed that the presence of BM negatively affects OS outcomes in mRCC patients treated with first-line tyrosine kinase inhibitors (TKI), particularly in the International Metastatic Renal Cell Carcinoma Database (IMDC) intermediate-risk group[16]. Treatment toxicity and disease progression are expected to have a negative impact on QoL while increasing costs, due to the associated need for additional treatment, hospitalizations, and other management requirements. Therefore, correctly identifying patients with BM for preemptive treatment is of utmost importance for minimizing morbidity and improving patient outcomes and QoL. This review addresses RCC BM pathophysiology, as well as diagnosis and treatment in the mRCC setting.

**BONE METASTASES PATHOPHYSIOLOGY**

The metastatic cascade is an intricate and complex process that culminates with cancer cell seeding and colonization at a distant site of the primary tumor[17]. Bone marrow is a preferential site for seeding of
cancer cells expressing C-X-C Motif Chemokine Receptor 4 and 7 (CXCR4 and CXCR7), due to the stromal cell-derived factor 1, also known as C-X-C Motif Chemokine 12 (CXCL12), chemotactic effect\[\textsuperscript{23}\]. After seeding, interaction with bone microenvironment is crucial for colonization, and, as cancer cells adapt and grow, they ultimately become able to exploit and manipulate the stroma, in the so-called “vicious cycle of bone metastases” by Mundy and Guise\[\textsuperscript{24}\].

The balance between bone resorption and formation is well preserved in healthy bone, whereas it is disrupted and shifted towards bone resorption in BM. In physiological conditions, the triad of receptor activator of nuclear factor κB ligand (RANKL), RANK, and osteoprotegerin (OPG) assures homeostasis\[\textsuperscript{25}\]. RANK is a transmembrane receptor that is present on the surface of osteoclast precursor cells and can be activated by its ligand RANKL, produced by osteoblasts, osteocytes, and bone marrow stromal cells. RANK activation by its ligand regulates osteoclast activity, survival, and differentiation\[\textsuperscript{26}\]. The RANKL soluble decoy OPG, secreted by mature osteoblasts and stromal cells, can block RANK activation and prevent bone resorption. In pathological conditions, such as those of osteolytic metastases, tumor-derived osteoclastogenic factors - such as interleukin 1 (IL-1), IL-6, parathyroid hormone-related protein (PTHrP), prostaglandin E2, colony-stimulating factor 1, and tumor necrosis factor-α - elicit increased bone resorption\[\textsuperscript{27}\]. RANKL expression is upregulated by increased PTHrP and IL-11, leading to OPG downregulation and activated osteclastogenesis. Therefore, mitogenic factors are further released from bone matrix, further feeding the “vicious cycle” of tumor growth and bone resorption.

In mRCC with BM, the hepatocyte growth factor/c-Met pathway has an important role in disease pathophysiology, as recently reviewed\[\textsuperscript{28}\]. c-Met is abnormally expressed in different tumors, including RCC\[\textsuperscript{29-31}\]. In a retrospective analysis of nephrectomy specimens from 17 mRCC patients, high c-Met expression at primary sites was identified in 47% of nephrectomy and 86% of bone specimens\[\textsuperscript{32}\]. Pre-clinical data suggest that c-Met contributes to cancer stem cell (CSC) maintenance and CSCs-induced BM in mRCC, but its mechanism is still unknown.

**DIAGNOSIS**

As previously stated, BM early diagnosis and preemptive treatment are key for preventing SREs and reducing morbidity and QoL deterioration.

Conventional radiography performed in two projections is usually the first exam when patients have BM-suggestive symptoms, as it can be useful to retrieve information on size and location, as well as fracture risk, when performed by an experienced orthopedic surgeon. Additionally, this exam [Figure 1] enables classifying whether the tumor is osteolytic (bone-destroying), osteoblastic (bone-building), or mixed. Although quick and easy to obtain, as well as inexpensive, it needs to be further complemented with other methods, as it is not suitable for use as a screening test, due to its low sensitivity.
Figure 1. Right calcaneus osteolytic bone metastasis from mRCC (conventional radiography).

Computed tomography (CT) is another complementary imaging technique that can be used to diagnose and study BM, as it allows establishing the structure and volume of metastatic lesion, magnitude of cortical bone destruction, and presence of periostal reactions, as well as assess fracture risk[33]. It can be extremely sensitive for osteolytic and osteoblastic bone lesions involving cortical bone, but not for tumors restricted to the marrow space, being of limited use as a screening test[34].

Skeletal scintigraphy with labeled phosphonates enables bone metabolism visualization. The classical screening imaging method is Technetium-99m bone scintigraphy, which enables depicting the entire skeleton and identifying osteoblastic activity, by detecting hot spots in bone metabolism. Therefore, it is very sensitive for BM with marked reactive bone hypermetabolism but relatively insensitive for tumors with non-reactive osteolysis or isolated bone-marrow infiltration (such as mRCC)[35,36]. One additional issue is that bone matrix regeneration induced by BM treatment leads to metabolic activation, which can be read as increased activity in the bone scan (flare phenomenon).

Positron emission tomography combined with CT with 18F-fluorodeoxyglucose (PET-CT-FDG) and whole-body magnetic resonance imaging (MRI) have increased sensitivity for bone lesions compared to bone scintigraphy, with the latter excelling at renal BM detection [Figure 2]. MRI is based on cell proton content detection, as cancer cells comprise more water molecules than bone marrow or cortical bone ones[37]. Given its high soft tissue contrast and spatial resolution, MRI can detect metastases with 91% sensitivity and 95% specificity very early in time, before bone structure changes can be detected in CT scan[38]. PET-CT-FDG uses 18F-fluorodeoxyglucose marker, which enables identifying cells with increased glucose uptake, such as tumor cells. It can be combined with CT scan data, retrieving high-resolution images with metabolic information[39]. As BM are highly metabolic, they can be detected with high sensitivity and specificity. Bone lesions associated with RCC are typically osteolytic, so bone scintigraphy with Tc-99m has a limited sensitivity when compared, for instance, with FDG PET/CT. Wu et al.[40] compared FDG PET with bone scan in patients with RCC and suspected bone metastases confirmed by histopathology. PET-FDG had 100% diagnostic sensitivity and accuracy, while bone scintigraphy had 77.5% and 59.6%, respectively. Besides being two excellent diagnostic methods, PET-CT-FDG and CT scan are also better at assessing therapy response in bone lesions[41], however with the drawbacks of availability and price.
Figure 2. Osteolytic lesion of mRCC with soft tissue component and spinal cord compression on the 6th and 7th cervical vertebrae (MRI).

BONE METASTASES TREATMENT
Bone metastasis in mRCC can be targeted by several classes of agents or local ablative therapies, in both monotherapy and combination [Table 2].

Bone-targeted agents
According to radiographic density, BMs can be osteoblastic, osteolytic, or mixed. However, as osteoclastic activity increase is common to all BM types, targeting osteoclasts has been one of the main treatment approaches in this setting [42]. Bone-targeted agents (BTAs), such as bisphosphonates (BPs) and denosumab, are used to prevent SREs, and recent evidence suggests that BPs, such as zoledronic acid, can be adjunctively used with systemic therapy to decrease SREs in patients with mRCC and BM.

BPs have the particularity of high affinity for calcium ions and become attached to hydroxyapatite binding sites on the bone surface, especially when undergoing active resorption. BPs are internalized by active osteoclasts, inhibiting their function [43]. Pamidronate and zoledronic acid are European Medical Agency - and Food and Drug Administration - approved BPs for prevention of SREs in patients with solid tumors and BM.

Different types of BPs are currently available. First-generation (non-nitrogenous) BPs include clodronate, first studied in 1987 in a small trial with 34 breast cancer patients with osteolytic BM. The study showed that a 1.6 g/day dose of clodronate for 12 months reduced bone pain, development of new osteolytic foci, and BM extension and prevented severe hypercalcemia [44]. This was further validated some years later, in a larger, double-blind trial that showed a drastic SRE reduction [45].

Second-generation nitrogenous BPs were subsequently developed. Pamidronate at the dose of 90 mg intravenous monthly for 12 cycles was evaluated in a double-blind randomized trial in breast cancer
Table 2. Treatments for bone metastasis in mRCC

| Bone targeted agents      |   |
|---------------------------|---|
| Bisphosphonates           |   |
| Denosumab                 |   |

**Systemic therapy**
- TKI (sunitinib, cabozantinib)
- IO (nivolumab, pembrolizumab)

**Local treatments**
- Stereotactic ablative radiotherapy
- Surgery

Patients with BM under active systemic therapy. In this trial, significant SRE and pain reduction was observed\(^{46,47}\).

More recently, the third-generation nitrogenous BP zoledronic acid was compared with pamidronate, also in the breast cancer with BM setting, with no significant differences\(^{48}\). Zoledronic acid was further compared with ibandronate in the ZICE trial, a positive non-inferiority study that placed ibandronate as an alternative for patients for whom frequent hospital visits are not ideal\(^{46}\).

Data on BP efficacy in solid tumors other than breast and prostate are scarce. Zoledronic acid was evaluated in a placebo-controlled trial of 773 patients with other tumors and BM and shown to reduce SRE incidence and increase time to first SRE\(^{50}\).

In the mRCC setting, a phase II trial randomized 30 patients in a 1:1 ratio to receive everolimus alone or everolimus and zoledronic acid\(^{51}\). Despite everolimus no longer being the standard mRCC treatment, zoledronate may have an important role in delaying time to SREs. This was supported by a recent meta-analysis showing that zoledronic acid reduced SRE risk by 68% (HR = 0.32, 95%CI: 0.19-0.55) compared with placebo or no zoledronic acid\(^{52}\).

The BTA denosumab, a fully humanized anti-RANKL monoclonal antibody, emerged from research on key bone remodeling mediators\(^{53}\). Denosumab can be an alternative for patients with renal disfunction or other conditions, limiting the use of zoledronic acid. In a pivotal phase III trial, denosumab was non-inferior to zoledronic acid in delaying time to first SRE in various advanced tumors (HR = 0.84, 95%CI: 0.71-0.98)\(^{54}\). Additionally, a meta-analysis including 5723 patients from three trials found that denosumab significantly improved time to first SRE by a median of 8.21 months and reduced the risk of a first SRE by 17% compared with zoledronic acid (HR = 0.83, 95%CI: 0.76-0.90, \(P < 0.001\))\(^{55}\). Data regarding use of denosumab in urological cancers were also presented at ASCO 2013\(^{56}\), in a study including urological patients of the 20050244 trial and prostate cancer patients of the 2005103 trial. In total, 85 mRCC patients were randomized to zoledronic acid and 70 to denosumab in the 20050244 trial, which reported a trend in lowering the time to occurrence of first on-study SRE with denosumab (HR = 0.71, 95%CI: 0.43-1.17).

It should be noted that there is an added risk of osteonecrosis of the jaw (ONJ) with cumulative exposure to denosumab and BPs\(^{57}\). Therefore, physicians should consider a full dental examination in patients planned to start these agents, and any necessary oral health intervention should be done before starting therapy. Additionally, frequent clinical assessment of the oral mucosa should be a priority. Particular attention should be given when administering BTAs concomitantly with other therapies, as anti-angiogenic TKIs
further increase ONJ risk when combined with those agents. In a series of 41 patients treated with denosumab and an anti-angiogenic TKI, ONJ incidence was 17% (contrasting with pivotal denosumab trials)^58. Additionally, a multicenter retrospective study including 44 patients mostly under anti-angiogenic TKIs and BTAs reported an unexpectedly high number of ONJ cases, which contrasts with literature data and suggests that the frequency of ONJ in RCC patients may be largely underestimated^59.

In addition to bisphosphonates, radioisotopes have also been investigated in this setting. Radium-223 acts as a calcimimetic, and therefore it has a high affinity to skeletal lesions that typically have an increased bone turnover. Being an alpha-emitting agent, radium-223 can prevent the progression of bone metastasis by killing cancer cells and breaking the vicious cycle of bone metastasis, stabilizing the tumoral microenvironment. In a study by McKay et al.,^60 the addition of radium-223 to VEGF-targeted therapy in patients with mRCC and bone metastases was studied. This association produced a significant decrease in the serum levels of five bone turnover markers, with no limiting toxicity. These data suggest that radium-223 associated with VEGF-targeted therapy is safe and might be useful in treating mRCC patients, although we need clinical trial data to see if this approach reduces the rate and incidence of SREs in these patients^60.

The RadiCal trial is a phase II randomized protocol that is currently investigating this (SSE as a primary endpoint), by comparing cabozantinib with radium-223 in combination with cabozantinib (NCT04071223).

**Systemic therapy**

Systemic treatment options for patients with mRCC include TKIs targeting the vascular endothelial growth factor receptor, the mammalian target of rapamycin (mTOR), and other pathways^61. Additionally, monoclonal antibodies, such as the anti-VEGF bevacizumab, and immune checkpoint inhibitors, such as pembrolizumab, nivolumab, and ipilimumab, have yielded positive results, both alone and in combination^62,63.

Data regarding BM and systemic therapy is also scarce. Preclinical evidence suggests that sunitinib can inhibit osteoclastic activity^64. In a study by Shinya et al.,^64 the number of osteoclasts in mice treated with sunitinib was significantly lower than in the controls, suggesting sunitinib may inhibit osteoclast maturation. Additionally, a decline in serum and urine levels of amino-terminal telopeptide was seen in patients with mRCC during the first four weeks of sunitinib treatment, suggesting that it might have significant activity in patients with RCC and bone metastasis^65.

Although there are no prospective clinical data supporting this, two retrospective analyses found that sunitinib prolongs OS compared to historical controls^66 and decreases development and time to new bone lesions compared to sorafenib^67.

Cabozantinib is an approved treatment for mRCC, in both first and second line, and has been studied in the subgroup of BM patients, in the METEOR trial. In this study, the median survival of patients with BM was 20 months in the cabozantinib arm and 12 months in the everolimus arm in the second- or higher-line setting^66,69. In a preclinical study, it was shown that doses of cabozantinib below the threshold of cytotoxicity inhibit osteoclast differentiation and bone resorption activity. Additionally, cabozantinib downregulated the expression of osteoclast marker genes, and direct cell-to-cell contact between cabozantinib pre-treated osteoblasts and untreated osteoclasts confirmed the indirect anti-resorptive effect of cabozantinib^70. Moreover, recent data from a subgroup analysis of the Checkmate-9ER trial evaluating nivolumab plus cabozantinib confirm an OS and PFS benefit of the combination in BM patients^12. A favorable OS trend was also observed in a subgroup analysis of the Checkmate-214 trial^64, comparing nivolumab plus ipilimumab with sunitinib, and in the Checkmate-025 trial^71, comparing nivolumab with
everolimus in second or further lines of treatment.

There is an ongoing phase II trial investigating the combined use of pembrolizumab and denosumab in clear-cell mRCC that may also shed some light on this treatment option\[72].

**Stereotactic ablative radiation therapy**

RCC is refractory to conventional radiation therapy but responds to higher doses per fraction. In 2019, the National Comprehensive Cancer Network guidelines included stereotactic ablative radiotherapy (SBRT) in the treatment of recurrent and metastatic RCC\[73\]. Because tumor cell survival and proliferation directly rely on blood supply, SBRT has been shown to have a direct effect on tumor vasculature.

The use of SBRT in metastatic bone lesions has also been investigated. Investigators from the Memorial Sloan Kettering Cancer Center compared the use of single-fraction (18-24 Gy) or hypofractioned (3-5 fractions to 20-30 Gy), by assessing tumor control rates in 105 patients with extracranial mRCC. Local control rates were best when using high single (24 Gy) compared with low single (< 24 Gy) dose or hypofractionated regimens, with three-year local PFS of 88%, 21%, and 17%, respectively\[74\]. Jhaveri et al.\[75\] investigated pain relief in RCC BM patients treated with SBRT and reported quicker and more durable pain control in those treated with BED of > 85 Gy compared with < 85 Gy, with 78% and 32% of patients, respectively, showing symptom control at 10 weeks. A study from the M.D. Anderson Center reported one-year spine tumor PFS of 82.1% with SBRT, with approximately half of all patients free of symptoms at one year\[76\].

With additional targeted therapies, the role of systemic therapy in RCC continues to improve, increasing patients’ survival and QoL. SBRT will play a valuable role in providing not only high local control rates but also prolonged symptomatic control from months to years.

**Surgery**

In mRCC, surgical metastasectomy (SM) is mostly performed for patients with pulmonary lesions, which represent the most frequent site for RCC metastases\[77\]. A systematic review of eight studies assessed the role of local therapy in mRCC and reported a longer OS and cancer-specific survival (CSS) in patients undergoing complete metastasectomy than in those undergoing either incomplete or no metastasectomy [40.8 months, interquartile range (IQR): 31.6-48.0 vs. 14.8 months, IQR: 13.3-21.0, respectively]. Patients with pancreatic, liver, and lung lesions derived the greatest advantage from complete metastasectomy when compared to incomplete or no metastasectomy\[77\].

Generally, BM are more resistant to systemic therapy and radiotherapy when compared to other metastases sites, and, additionally, they are highly destructive. The role of metastasectomy in mRCC with BM was recently reviewed in a retrospective study with 114 patients. Single BM was seen in 68 (59.6%) patients, whereas 46 (40.4%) had multiple (2-5) BM. BM were located in axial bones (spine, skull, and ribs) in 47 (41.2%) patients, in the appendicular skeleton (extremities, pelvis, clavicle, and scapula) in 20 (17.5%), in multiple sites in 33 (28.9%), and in unknown exact location in 14 (12.3%)\[78\]. BM excisional surgery, particularly in the spine, is a procedure which is technically very demanding, even for specialized orthopedic and spine surgeons. Bone metastases from mRCC are destructive and hypervascular [Table 3], and, in the majority of cases, reconstruction is required after tumor resection to support the operated lesion against the load. A case series of 36 consecutive patients concluded that, for selected patients, complete spinal resection of spinal metastases can potentially prolong survival, with a median CSS of 130 months. In this series, the presence of liver metastases was a potential drawback to this strategy, being associated with
Table 3. Typical characteristics of BM in renal cell carcinoma

| Clinical and pathologic features of renal cell carcinoma BM |
|----------------------------------------------------------|
| · Mostly osteolytic[15] |
| · High c-MET expression[29-31] |
| · Radioresistant (require high biologically effective dose (BED))[76] |
| · Highly vascularized and destructive[80] |

short-term survival after spinal metastasectomy[79].

In many cases, treatment goals for patients with BM do not include complete SM, but instead palliative surgery combined with non-surgical treatment to preserve or restore neurological function and improve pain control[80]. Spinal metastases can represent a particularly difficult case with regards to symptoms and treatment, as neurological pain is often added to nociceptive pain, and neurological impairment can be observed. Due to close contact with the spinal cord, spinal BM treatment is challenging and requires a multidisciplinary approach[81].

CONCLUSION

BM and SREs increase economic burden and decrease QoL, ultimately increasing morbidity throughout the metastatic disease course. Additionally, the overall prognosis of these patient is poor, compared with patients without BM. Advances in systemic and ablative therapy have improved these patients’ outcomes in recent years. BM in patients with RCC may have a special sensitivity to drugs targeting c-Met, and administration of BTAs can further improve time to SREs and minimize QoL deterioration. Combining and sequencing all these resources can be complex, making multidisciplinary intervention of utmost importance to maximize patient outcomes and treatment.

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Contributed significantly and similarly to the conceptualization, writing and review of the article: Mansinho A, Nejo P, Leitão T, Casimiro S, Costa L

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