Bone metastases from differentiated thyroid carcinoma

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Abstract

The presence of distant metastases from differentiated thyroid carcinoma decreases the 10-year survival of patients by 50%. Bone metastases represent a frequent complication especially of follicular thyroid cancer and severely reduce the quality of life causing pain, fractures, and spinal cord compression. Diagnosis is established by correlating clinical suspicion with imaging. Imaging is essential to detect, localize, and assess the extension of the lesions and should be used in conjunction with clinical evidence. Bone metastases are typically associated with elevated markers of bone turnover, but these markers have not been evaluated in differentiated thyroid cancer. Skeletal and whole-body magnetic resonance imaging and fusion 2-deoxy-2-[18F]fluoro-D-glucose whole-body positron emission tomography/computed tomography (PET/CT) are the best anatomic and functional imaging techniques available in specialized centers. For well-differentiated lesions, iodine-PET scan combined with ¹₂⁴I-PET/CT is the newest imaging development and ¹³¹I is the first line of treatment. Bisphosphonates reduce the complications rate and pain, alone or in combination with radiiodine, radionuclides, or external beam radiotherapy and should be employed. Surgery and novel minimally invasive consolidation techniques demand an appropriate patient selection for best results on a multimodal approach. Basic research on interactions between tumor cells and bone microenvironment are identifying potential novel targets for future more effective therapeutic interventions for less differentiated tumors.

Endocrine-Related Cancer (2008) 15 37–49

Introduction

Patients with differentiated thyroid carcinoma (DTC) have a 10-year survival rate of 80–95%. However, when distant metastases are present, the overall 10-year survival rate is 40% (Hoie et al. 1988, Ruegemer et al. 1988, Mizukami et al. 1990, Samaan et al. 1992, Schlumberger 1998). One selected compendium of 13 studies found that among 1231 patients, 25% of metastases were to the bone, 49% to the lung, 15% to both lung and bone, and 10% to other soft tissues (Mazzaferr 1993). The methods used more than 15 years ago for diagnosis and treatment differ, however, from the present management. A more recent study shows that survival drops to 14% for patients older than 40 years with macronodular lung metastases or multiple bone metastases (Durante et al. 2006).

After the age of 40 years, 10% of patients with papillary thyroid carcinoma (PTC), 25% of patients with follicular thyroid carcinoma (FTC), and 35% of patients with Hurthle cell carcinoma (Lopez-Penabad et al. 2003) develop distant metastases. Bone metastases from DTC occur in 2–13% of patients (McCormack 1966, Marocci et al. 1989, Proye et al. 1992, Schlumberger et al. 1996, Fanchiang et al. 1998, Lin et al. 1999, Pittas et al. 2000, Bernier et al. 2001). The study of Durante et al. (2006) found 44% of bone metastases in 444 metastatic DTC patients. They are more frequent in FTC (7–28%) compared with PTC (1.4–7%).

When bone metastases are present, the overall survival at 10 years was reported to range from 13 to 21% (Beierwaltes et al. 1982, Schlumberger et al. 1996).
Physiopathology of bone metastases from DTC

Disseminated cells of various cancers are clonally different from primary tumors. Different genotypes might be selected by different microenvironments. Some studies have shown that bone involvement is more frequent in well-DTC compared with moderately or poorly DTC (Nagamine et al. 1985, Marcocci et al. 1989, Glinsky et al. 2005). These findings gave rise to terms such as benign metastasizing thyroid tumor, metastasizing adenoma, and malignant adenoma. However, the study of Marcocci did not include poorly DTC. Out of a total of 780 patients, only 30 (3.8%) had bone metastases. In two studies reporting a total of 190 follicular carcinoma patients with bone metastases, 148 were moderately differentiated (Massin et al. 1984, Schlumberger et al. 1986). In the series from Memorial Sloan-Kettering Cancer Center (MSKCC) on 79 DTC patients with bone metastases, two-thirds had either poorly differentiated or undifferentiated lesions at both the primary and metastatic sites (Tickoo et al. 2000). In one series of metastatic patients, multiple bone metastases correlated with poor survival (Durante et al. 2006).

Only cancer cells that exchange biological information with bone environments are able to establish bone metastases. Paget (1989) ‘seed and soil’ hypothesis is widely accepted (Liotta & Kohn 2001, Fidler 2002): circulating cancer cells (seeds) have a propensity to metastasize to organs with the micro-environment (soil) advantageous for their growth. The ability of cells to survive, multiply, and recruit a blood supply gives rise to metastases. Bone is a large repository for immobilized growth factors, including transforming growth factor, insulin-like growth factor-I and -II (IGF-I and -II), fibroblast growth factors, platelet-derived growth factors, bone morphogenetic proteins, and calcium. Released and activated during bone resorption these factors render the bone fertile for tumor growth (Roodman 2004).

More than 80% of bone metastases from all tumors including DTC are located in axial skeleton red marrow where blood flow is high (vertebrae, ribs, and hips). Tumor cell adhesive molecules bind the tumor cells to marrow stromal cells and bone matrix allowing them to grow and produce angiogenic and bone-resorbing factors.

One recent review concentrated on the physiopathology of bone metastases from thyroid cancers (Wexler & Sharretts 2007). Molecular biology studies try to explain the higher propensity of follicular and less differentiated cancers to metastasize to bone. One hypothesis was the difference in the expression of tumor suppressor genes, caveolin-1 and caveolin-2. Although the expression is up-regulated in FTC and down-regulated in PTC, the down-regulation was also found in the anaplastic thyroid cancer cells (Aldred et al. 2003, 2004). The latter group expresses in large amounts focal adhesion kinase (FAK). FAK affects adhesion, motility, and distant site tumor growth (Owens et al. 1996). Follicular thyroid cancer cells express less fibronectin resulting in higher cellular adhesion and migration (Chen et al. 2001).

Most DTC patients have predominantly osteolytic lesions, with secondary formation of bone in response to bone destruction. There is usually a concomitant increase in both technetium-labeled bisphosphonates uptake on bone scintigraphy (indicating bone formation) and N-telopeptide values (NTx, a marker of bone resorption). Bone sialoprotein (Bellahcene et al. 1998) and integrin avb3 (Pecheur et al. 2002) were found to be up-regulated in thyroid cancer cells and involved in tumor aggressiveness and tumor osteolysis.

Clinical evolution and symptoms

The most common manifestations of bone metastases from DTC: pain, fractures, and spinal cord compression are associated with lesions in the axial skeleton.

Pain often presents as the principal symptom of metastatic bone involvement (Coleman 2001) and progressively becomes more severe and resistant to commonly used non-opioid analgesics. Metastatic bone pain is associated with local chemical release of cytokines by tumor cells causing stimulation of intraosseous nerves, and pressure or mass effect of the tumor tissue within the bone (Selvaggi & Scagliotti 2005).
One retrospective study found that spinal cord compression occurs more frequently in DTC when compared with other bone-seeking cancers: 28% compared with prostate (10%) and breast cancers (8%) (Coleman 2006). The incidence of pathological fractures was 13%, and the incidence of both spinal cord compression and pathological fracture was 6% (Bernier et al. 2001). Early diagnosis, high-dose corticosteroid treatment, decompressive surgery, and spinal stabilization or radiotherapy are essential for neurological recovery within the first 24–48 h of presentation for bone metastases (Onimus et al. 1996, Alvarez et al. 2003). Back pain associated with an abnormality on a plain spinal radiograph in these cancer patients has a 60% incidence of epidural disease on magnetic resonance imaging (MRI).

**Diagnosis**

Diagnosis is established by correlating clinical suspicion with imaging. For cancers with high tropism to bone, elevated bone turnover markers could be the first sign of bone involvement but are not yet explored in thyroid cancer.

**Biology**

In other types of cancer, elevated markers of bone turnover predict survival and early tumor response to treatment (Coleman et al. 2005). Specific tests include markers of bone formation (osteocalcin, bone-specific alkaline phosphatase (BSAP), and the cleaved amino and carboxy terminal peptides from procollagen: the N and C terminal propeptides of human procollagen type I) and especially bone resorption (the C terminal telopeptide and N terminal telopeptide NTx of collagen type I and deoxypyridinoline; Demers et al. 2003) and markers of osteoclastogenesis: the soluble receptor activator of nuclear factor-κB ligand (sRANK-L), its receptor (RANK), and its decoy receptor osteoprotegerin (OPG; Demers et al. 2003). These markers have not been evaluated in differentiated thyroid cancer.

**Imaging**

Imaging is essential to detect, localize, and assess the extension of the lesions and should be used in conjunction with clinical and biochemical evidence (Rosenthal 1997, Rybak & Rosenthal 2001). Imaging can also guide biopsies for histomorphological diagnosis.

**Anatomic imaging**

Plain radiographs can show bone destruction, sclerosis, and the absence of pedicles by tumor infiltration, but lesions may not appear on X-rays for several months with the limit of detection of lesions >1 cm. In the study of Schlumberger, among 115 thyroid cancer patients who had bone metastases only, bone radiographs documented a single bone metastasis in 33 patients, multiple bone metastases in 74 patients, and were normal in 8 patients (Durante et al. 2006).

Computed tomography (CT) can evaluate the extent of metastatic lesions and is particularly useful for sites that are difficult to evaluate, i.e., the spine and pelvis. For bone-seeking cancers, diagnostic sensitivity is 71–100%. CT describes foci before bone destruction has occurred as increased attenuation of the normally fatty bone marrow. Low soft tissue contrast and lack of information on broad areas of the spine represent the technique’s shortcomings.

Skeletal and whole-body MRI provides detailed images of both bone and bone marrow, so it is best employed when spinal cord compression and involvement by the tumor is suspected. Whole-body MRI yielded a sensitivity of 94% and diagnostic accuracy of 91% when screening for bone metastases in patients with different cancer types (Schmidt et al. 2006) with a cut-off size for the detection of malignant bone lesions of 2 mm.

No data are available on the sensitivity and specificity of CT and whole-body MRI in screening for bone metastases from DTC, but when bone involvement is suspected (aggressive tumor behavior, metastases at other sites, clinical data), whole-body MRI or (non)enhanced CT must be employed for risk assessment and treatment planning.

**Functional imaging**

The perimetastatic osteoblastic reaction (increased blood flow, new bone formation, and enhanced bone matrix turnover) accounts for abnormal radiopharmaceutical uptake on bone scan. 99mTc-labeled diphosphonates such as methylene diphosphonate (MDP) are most commonly used for localization and staging. Bone scan can also assess response to treatment of some bone-seeking cancers. When compared with anatomic imaging, the single-photon emission CT (SPECT) and SPECT/CT could make MRI better for vertebral body lesions and SPECT for the posterior elements (Ryan & Fogelman 1995, Kosuda et al. 1996).

However, bone scintigraphy relies on the detection of osteoblastic reaction. In thyroid cancer, bone metastases are primarily osteolytic, so the proportion of false-negative and false-positive results is high (Ito et al. 2007).
Tumor cell imaging by $^{131}$I or $^{123}$I whole-body scan (WBS) are both more specific and sensitive than $^{99m}$Tc-MDP but only for well-differentiated, NIS-positive thyroid tumors (Schirrmeister et al. 2001, de Geus-Oei et al. 2002). However, multiple site bone involvement usually correlates with aggressive, less differentiated tumor types.

2-Deoxy-2-$[^{18}$F$]$fluoro-D-glucose whole-body positron emission tomography (FDG-PET) shows preferential tracer uptake in malignant cells with a high turnover rate by increased glucose metabolism. In thyroid cancer, PET imaging is useful in patients with metastatic poorly differentiated tumors, with high thyroglobulin (Tg) levels and negative $^{131}$I-WBS. A positive FDG scan has a strong negative prognostic value as it was associated, by itself, with an eightfold increased risk of death (Wang et al. 2000, Robbins et al. 2006).

In one study that compared different imaging techniques for the detection of distant metastases from thyroid cancer, whole-body $^{18}$F-FDG PET, $^{99m}$Tc-MIBI SPET, and post-therapeutic $^{131}$I-Na scintigraphy all demonstrated five of the six bone metastases (83.3%; Iwata et al. 2004).

Thyrotrophin (TSH) stimulates thyrocyte metabolism, glucose transport, and glycolysis. FDG is a glucose analog. Several studies showed that rhTSH stimulation improves the detection of occult thyroid metastases with FDG PET, compared with scans performed on TSH suppression (Chin et al. 2004).

The advantages of rhTSH-stimulated FDG PET and PET-CT fusion scanning in metastatic thyroid cancer are currently under evaluation (ClinicalTrials.gov Identifier: NCT00181168).

**Developments in tumor cell imaging for DTC**

Positron-emitting radioisotope $^{124}$I (half-life, 4.2 days) uses the sodium iodide symporter and is employed for three-dimensional imaging of the distribution of radioactive uptake within the benign and malignant thyroid disease since (the 1980s) 1985 (Frey et al. 1986). In DTC CT, $^{131}$I-WBS, $^{124}$I-PET, and combined $^{124}$I-PET/CT imaging detected 56, 83, 87, and 100% of lesions respectively (Freudenberg et al. 2004). $^{124}$I-PET detected more small bone metastases compared with $^{131}$I-WBS. $^{124}$I-PET is performed as soon as 24 h after the administration of the radiotracer and yields the same diagnostic accuracy as high-dose WBS does. These results warranted confirmation through a clinical trial that will start recruiting at the Stanford University (ClinicalTrials.gov Identifier: NCT00373711).

**Treatment**

Management of these patients must involve a multi-disciplinary approach consisting of medical treatment (analgesia and bisphosphonates), radiotherapy, surgery, and radioisotopes.

**Current treatments**

**Radioactive iodine**

All series on radioactive iodine therapy for bone metastases showed that bone metastases are generally resistant to commonly used activities of $^{131}$I and may require other approaches (Schlumberger et al. 1996). The low remission rate (29–35%) could be related to the large extent of disease at presentation (Marcocci et al. 1989, Pacini et al. 1994, Zettinig et al. 2002) and the dose delivery to bone metastases. Patient-specific, three-dimensional (3D) imaging-based dosimetry with $^{124}$I-PET is the subject of ongoing investigation for thyroid cancer (Sgouros et al. 2004).

A retrospective study evaluated therapeutic outcome, total administered radioiodine activities, and side effects (blood count alterations grades I–IV, WHO classification, and acute leukemia) in 107 patients with initial bone metastases (Petrich et al. 2001). Patients younger than 45 years were classified as group 1 (stage II, ‘low risk’, WHO classification) and those over 45 years as group 2 (stage IV, ‘high risk’). Total or partial remissions were more frequent in group 1 than group 2 (62.5% vs 49.5%) with lower delivered activities (18.89 ± 15.08 GBq vs 41.97 ± 31.25 GBq). Three out of four group 1 patients with three or less bone metastases had complete remission (11.1 GBq). Long-term and partial remission were obtained in a large proportion of patients, 24 and 27% of cases respectively, with low overall rate and degree of side effects. This study concluded that initial bone metastases in selected DTC patients up to 45 years and especially in those with less than three bone metastases can be treated with curative intent.

**Surgery**

The main indications for surgery are persistent pain refractory to medical therapy, tumors with poor radioactive uptake, and spinal instability with or without neural compression. Various scoring systems guiding the surgical strategy are based on the primary tumor type, the presence of visceral metastases, and number of bone metastases (Tokuhashi et al. 1990, Enkaoua et al. 1997, Tomita et al. 2001).

Few studies have analyzed the impact of surgery on DTC bone metastases (Marcocci et al. 1989, Proye et al. 1992, Pittas et al. 2000). Reports have shown that
Radiopharmaceutical ablation (RFA) and ethanol injection are minimally invasive techniques that require a highly specialized team and appropriate patient selection. The role in differentiated thyroid cancer lesions is being investigated (Monchik et al. 2006, Wexler & Sharrett 2007).

**External radiotherapy**

DTC bone metastases without affinity for radioactive iodine may respond to external beam radiation therapy (EBRT; Simpson et al. 1988, Simpson 1990, Tsang et al. 1998). EBRT is a palliative treatment indicated only when pain, risk for fracture, and neurological complications of spinal cord compression are present.

Complete or partial pain relief is obtained in more than 80% of patients for at least 6 months in 50% of cases. Irrespective of the fractionation schedule, the number of subsequent spinal cord compression or pathological fractures is low. All prospective randomized trials for different bone-seeking cancers showed that single fraction regimens are at least as effective as fractionated regimens with increased convenience and cost control (Falkmer et al. 2003, Frassica 2003).

For DTC, the combination of external radiotherapy and radiiodine therapy has been reported to have an impact on cancer recurrence and pain relief (Tubiana et al. 1985). When employed after surgery, it results in reduced risk for further bone destruction (Fig. 1).

RFA prevents cytokine release from tumor cells and reduces tumor growth into the peristeum. When combined with surgery (Dupuy et al. 2001, Halpin et al. 2004, 2005), it may improve the outcome of bone-seeking cancers. Currently, there are no data on the association of RFA with radiiodine for bone metastases of DTC.

**Chemotherapy**

A variety of different chemotherapeutic regimens have been used for metastatic thyroid cancer (Shimaoka et al. 1985, Ahuja & Ernst 1987, Hoskin & Harmer 1987, De Besi et al. 1991, Casara et al. 1993), with limited effect and duration of response. The addition of TSH stimulation to a conventional regimen of chemotherapy could increase its therapeutic effect without increased morbidity (Santini et al. 2002). Consistent data on chemotherapy response of bone metastases from poorly DTC are not yet available.

**Bisphosphonates**

In addition to strictly palliative interventions, bisphosphonates represent a very effective therapeutic option for the prevention of skeletal complications of bone metastases.
metastases from different primary tumors. They bind preferentially to bone at the sites of active bone metabolism, are released from the bone matrix during bone resorption, and inhibit osteoclast activity, thereby reducing osteoclast-mediated bone resorption. There are several reports of sclerosis of lytic bone metastases on X-ray after bisphosphonates (Morton et al. 1988, Tanaka et al. 1996, Kokufu et al. 1998, 2000).

Nitrogen-containing bisphosphonates, such as zoledronic acid, pamidronate, and ibandronate, have a unique mechanism of action and greater clinical activity than first-generation bisphosphonates (Coleman 2004) with 50% reduction in the skeletal-related events. Still, these drug effects on tumor cells and reduction of tumor burden have to be further elucidated, as well as their potential as adjuvant therapy for cancer metastases.

**Anti-resorptive effect.** A recent study on patients with prostate cancer showed fewer skeletal-related events in the zoledronic acid arm compared with placebo (Canil & Tannock 2002). However, zoledronic acid failed to delay disease progression, lengthen survival, or improve quality of life. The same results were obtained in breast cancer.

In one small study, DTC patients who had been administered pamidronate (90 mg, as a 2-hour i.v. infusion monthly for 12 consecutive cycles) showed a significant decrease in bone pain, improved performance status, and improved quality of life. However, no significant decrease in analgesic consumption was recorded. Partial radiographic response of bone lesions was observed in two out of ten patients. Side effects were mild and transient (Vitale et al. 2001).

**Vehicles for radioisotopes.** Bisphosphonates-triggered radioisotopes for the palliation of metastatic bone have been proposed for patients with painful disseminated skeletal metastases in whom external beam radiotherapy cannot be administered (Finlay et al. 2005).

Commercially available bone-seeking radiopharmaceuticals that target the perimetastatic osteoblastic reaction include: $^{153}$Sm-EDTMP, $^{186}$Re-HEDP, or $^{89}$Sr chloride (Pandit-Taskar et al. 2004). The $\beta$-emitting radionuclides deliver energy over a range of several millimeters. The newest developments are radionuclides with high-linear energy transfer (LET) $\alpha$-particles like $^{223}$Ra (Nilsson et al. 2005) enhance radiobiological potency with increased energy delivery, providing an anti-tumor effect in addition to palliation.

All isotopes seem to be effective in terms of pain control and response rates are between 40 and 95% (Lam et al. 2007) for breast and prostate cancers. Thrombocytopenia and neutropenia, which are common to all isotopes, are generally mild and reversible side effects. Dosimetric studies are underway to reduce side effects and provide response predictions. Treatment efficacy is improved with combination therapy. In 126 prostate cancer patients with bone metastases that all received external beam radiotherapy in combination with either

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**Figure 1** The vicious circle at the bone remodeling unit. Tumor cells produce factors that increase the formation of osteoclasts through the RANK-RANKL (receptor activator of nuclear factor-κB ligand) system: IL-6 (interleukin-6), prostaglandin E2 (PGE2), tumor necrosis factor (TNF), macrophage colony-stimulating factor (M-CSF), and parathyroid hormone-related peptide (PTH-rP). During bone resorption, osteoclasts release transforming growth factor-β (TGF-β), insulin-like growth factors (IGFs), fibroblast growth factors (FGFs), platelet-derived growth factor (PDGF), and bone morphogenetic proteins (BMPs), which increase the production of parathyroid hormone-related peptide by tumor cells, as well as growth factors that increase tumor growth. This represents the vicious circle that increases bone destruction and tumor growth. OPG, osteoprotegerin. Adapted from (Clamp et al. 2004, Roodman 2004).
89Sr or placebo; at 3 months, 59% of patients in the active group were free of new painful metastases, suggesting an 89Sr anti-tumor effect (Porter et al. 1993). The combination of zoledronic acid with 89Sr improves pain reduction (Storto et al. 2006), leading to the speculation that strontium could have increased availability due to greater bone remodeling associated with bisphosphonates. Finally, several studies evaluated the radio-nuclides/chemotherapy combined regimen for prostate cancer (Tu et al. 2001). These compounds have yet to be investigated for bone metastases from thyroid cancer.

**Anti-tumor effect.** Preliminary studies suggest that bisphosphonates can exhibit anti-tumor activity both in vitro and in vivo (Clezardin et al. 2003, 2005, Green 2003). Nitrogen-containing bisphosphonates inhibit the mevalonate pathway that induces apoptosis in osteoclasts and tumor cells alike. They can impair tumor cell adhesion, invasion (Woodward et al. 2005) and proliferation, reinforce the effects of cytotoxic agents, and exhibit anti-angiogenic effects. In thyroid cancer cell lines, clodronate inhibited cell growth of endocytic macrophages, osteoclasts, and several cancer cells in a dose-dependent manner. It transiently increased cytosolic Ca2+ on slow-growing SW579 thyroid cancer cells but not on fast-growing ARO cells. This implies that clodronate-mediated cell growth inhibition in slow-growing thyroid cancer cells might correlate with a Ca2+ signaling pathway (Yang et al. 2004).

Nevertheless, the concentrations required for these effects in vitro and in animal models were usually much higher than those used in clinical practice. In humans, there is no evidence from any of the clinical trials of bisphosphonates suggesting a beneficial effect on soft tissue metastasis.

In light of these data and comparing the risk of skeletal events in thyroid cancer with other cancers, where bisphosphonates are conventionally employed, bisphosphonates treatment should be employed in bone metastases from DTC. We are using in our practice the same protocols as for breast or prostate cancer, but definitive studies in this area are warranted. Selection of patients for other techniques (surgical procedures and radiotherapy) is made on the basis of a multidisciplinary approach.

**Developments and perspectives**

**Bone resorption inhibitors: targeting osteoclastogenesis**

New approaches to prevent and treat bone metastases are being developed (Wittrant et al. 2004). Examples include molecules that directly counter the receptor activator of RANKL, PTH-rP inhibitors (the major mediator of osteolytic disease), and anti-Dkk1.

Targeting the OPG/RANK/RANKL axis may offer a novel therapeutic approach to malignant osteolytic pathologies (Wittrant et al. 2004, Croucher et al. 2005). Clinical trials with AMG 162 (denosumab, a human monoclonal antibody to RANKL) are currently underway in Europe for patients with myeloma, breast, and prostate cancer skeletal metastases (Body et al. 2003). In the United States, several Phase II/III Clinical Trials with denosumab in patients with solid organ tumors or multiple myeloma with radiographic evidence of one or more bone metastases currently receiving oral or i.v. bisphosphonates are underway.

There are no such studies available yet (or for other targets of common osteotropic cancers: TGF-β, PTH-rP/Gli2, BMP-7) specifically for thyroid cancer bone metastases.

**Anti-angiogenic factors**

In radioactive iodine unresponsive thyroid cancer, several therapeutic approaches are being investigated. Tyrosine kinase receptors control angiogenesis and tumor growth. Tyrosine kinase receptors are targeted through: anti-VEGF – bevacizumab (Avastin), anti-Epidermal growth factor receptor (EGFR; Iressa), anti-Her2/neu (Herceptin) antibodies, and through small molecules (ZD6474 and high-affinity PPARgamma agonist RS-5444) inhibiting VEGFR, EGFR, TIE-2, and RET oncoproteins. Thalidomide blocks the angiogenesis cascade and combrestatins have pro-apoptotic effects. Other molecules target Ras, Raf (ISIS 5132), MEK, TNF-related apoptosis-inducing ligand (TRAIL), Akt/mTOR and HSP90 (geldanamycin), histone deacetylase (FR901228, depsipeptide), and COX2 (celecoxib).

FTC is the most angiogenic-dependent tumor of the thyroid gland. VEGF mRNA and protein levels are associated with mitogenic activity and tumor growth in FTC cell lines. Inhibition of VEGF production or VEGFR phosphorylation has been shown to reduce the growth of FTC xenografts (Soh et al. 2000, Ye et al. 2002, Schoenberger et al. 2004). EGF and its receptor (EGFR) are also over-expressed in thyroid carcinomas (Aasland et al. 1990, van der Laan et al. 1995) and co-expression of EGF and EGFR is associated with FTC bone metastases (Gorgoulis et al. 1992).

Therefore, blockade of both EGFR and VEGFR kinase activities and their downstream targets (Akt and mitogen-activated protein kinase) can offer an attractive approach for the treatment of FTC bone.
metastases. One study is underway in France using sunitinib malate (Pfizer): Sutent (THYSU, ClinicalTrials.gov Identifier: NCT00510640).

AEE788 (Novartis Pharma) inhibits both the EGFR and VEGFR receptors (Traxler et al. 2004). AEE788 alone or in combination with paclitaxel can significantly reduce tumor size and subsequent bone destruction by FTC in nude mice by direct anti-tumor and anti-angiogenic effects. A phase I study is currently recruiting adult patients with stage IV solid tumors (ClinicalTrials.gov Identifier: NCT00118456).

**Gene therapy**

Strategies of gene therapy in oncology are: oncopgene silencing, tumor suppressor replacement, prodrug therapy, immunotherapy, genetic immunization, antisense therapy, ribozyme therapy, antibody neutralization, chemoprotection, and radiiodide therapy.

Several agents are being evaluated for iodine-negative metastases from thyroid cancer, aiming mainly at tumor re-differentiation and increased expression of hNIS. These agents include lithium, retinoids (isotretinoin Accutane, bexarotene Tagretin), and thiazolidinediones (Zarnegar et al. 2002, Park et al. 2005). hNIS gene transfer into hNIS-defective thyroid cancer could improve the response to radiiodine of iodine-negative bone metastases (Haberkorn et al. 2004).

**Assessment of therapeutic response**

Tg and 131I-WBS have been established as important markers for follow-up in DTC. These markers are not specific for metastatic bone disease in DTC. There are very little data on DTC bone metastases with respect to techniques usually employed for other cancer types due to the reliability of 131I-WBS. However, they are valuable in poorly differentiated or undifferentiated iodine-negative thyroid cancers.

Many studies’ aim is to identify biological or imaging markers able to select patients with bone metastases, who are likely to respond to bisphosphonates and anti-neoplastic therapy and to indicate whether these interventions are effective (Clamp et al. 2004). Studies that use bone-specific biomarkers to tailor treatment with bisphosphonate or anti-neoplastic therapy have been developed like the multicenter bisphosphonate therapy directed by bone resorption markers (BISMARK) trial recruiting patients with breast cancer and associated metastatic bone disease. The trial is powered to prove non-inferiority between standard and marker-directed administration of zoledronate for the prevention of skeletal-related events.

On X-ray, positive response of the skeletal lesions may be visible as lesion sclerosis progressing from the periphery toward its center.

On bone scan, healing lesions usually demonstrate decreased radioisotope tracer uptake. The ‘flare’ phenomenon with increased uptake in the early phases of therapy can also be seen due to isotope uptake, by healing bone, and it has been suggested that an increase in the number of lesions is a more reliable marker of disease progression than an increase in the intensity of uptake.

CT density change can evaluate re-mineralization of osteolytic lesions with palliative radiotherapy (Reinbold et al. 1989, Chow et al. 2004). For less differentiated cancers, 18F-FDG-PET allows for earlier assessment of response compared with bone scan (Stafford et al. 2002). In the case of thyroid cancer, one clinical trial will try to evaluate overall response (complete and partial responses) rate based on conventional imaging methods (CT scan, RECIST) and tumor markers as well as early (7 days) and late (3 months) response rates based on functional imaging 18F-FDG-PET and to compare response rates by CT scan versus FDG-PET in patients treated by sunitinib malate (ClinicalTrials.gov Identifier: NCT00519896).

**Conclusion**

The presence of bone metastases alters the prognosis of patients with DTC. Radioiodine is the first-line treatment for NIS-positive lesions, but significant results are obtained only in select number of patients. In bone-seeking cancers, bisphosphonates are known to increase the time to skeletal complications and relieve pain especially when employed in combination with radioiodine, radionuclides, or external beam radiotherapy. The optimal timing and duration of bisphosphonate therapy and their potential in the prevention of bone metastases is under investigation in clinical trials for solid tumors. Selection of patients is crucial to optimize the multimodal approach of surgery and novel minimally invasive techniques. Apart from tumor and bone-targeted treatment, pain management, and psychological support are essential in clinical decision making. Therefore, a multidisciplinary approach provides an improved quality of life. Basic research on interactions between tumor cells and bone microenvironment are identifying potential novel targets for future more effective therapeutic interventions for poorly DTC.
Acknowledgements

The authors declare that there is no conflict of interest that would prejudice the impartiality of this scientific work.

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