Bone Markers in the Treatment of Cancer Related Bone Disease in Patients with Metastatic Breast Cancer

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Abstract

Bone metastases and treatment-induced osteoporosis are frequently the main causes of morbidity in patients with malignancies. Monitoring the level of bone markers (markers of bone metabolism) has been fairly well mapped in osteoporosis, where medical procedures can be modified according to the kinetics of marker levels before an answer can be evaluated by densitometry and before the onset of fractures. In bone metastases the role of these levels is not clear. The metabolic markers of bone resorption under review in this set were ß-Cross Laps (CTX) - a peptide that is a part of C-telopeptide, and N-terminal propeptide of collagen type 1 (P1NP).

Material and methods: We monitored a group of 52 female patients with metastatic breast cancer. The patients received appropriate systemic treatment based on the immunohistochemistry of the tumor; the treatment consisted of hormone therapy or chemotherapy, and included parenteral bisphosphonates (ibandronate and zoledronic acid alternately). Routine biochemical tests and blood count done prior to the initiation of therapy also included taking laboratory markers of bone metabolism CTX and P1NP and measuring bone mineral density (according to T-score). These bone markers were then checked in three, six, nine and twelve months, and matched with the progress of the disease. Total monitoring time was fifteen months.

Results: The patients in this set whose CTX value in the first sampling was less than 0.425 ran 8.5 times higher risk of death; the patients whose P1NP value reached more than 74 in the first collection ran 8.7 times higher risk of death. According to Cox proportional-hazards regression analysis for CTX, the significance level of p-value was 0.0452 and HR was 8.516 (95% CI 1.047 to 69.262), a difference which is not statistically significant. Regarding P1NP in Cox regression analysis, the significance level of p-value is 0.0433 and HR 8.673 (95% CI 1.067 to 70.520). Even this difference is therefore not statistically significant. When comparing the kinetics of marker levels, the difference is below statistical significance: p-value 0.6131 for P1NP and p-value 0.6357 for CTX.

Conclusion: The results of this study confirm a correlation between the starting levels of CTX and P1NP with the overall survival rate, which corresponds to the other results presented in literature.

Introduction

Bone metastases and treatment-induced osteoporosis are frequently the main causes of morbidity in patients with malignancies. The main indicator of low treatment efficacy of bone metastases is the incidence of skeleton-related events (SRE). Monitoring the levels of bone markers (biochemical markers of bone metabolism) has been fairly well mapped in osteoporosis, where medical procedures can be modified in accordance with the kinetics of marker levels before the answer can be evaluated by densitometry and before the development of a fracture [1,2]. In bone metastases, the role of these levels is not clear [3,4].

The first metabolic marker of bone resorption under review in this set was ß-Cross Laps (CTX) - a peptide that is a part of the C-telopeptide. It is localized on the beta chains of collagen type 1 and includes a position for possible pyridine coupling. It is easier to determine it from blood, something which is performed on an automated analyzer on a daily basis. It is recommended to determine CTX beta for monitoring the effectiveness of anti-resorptive therapy (e.g. by bisphosphonates, HRT) in osteoporosis or other bone disorders. In bone metastases, the optimal specificity seems to be (90%) at the value of 0.4 g/l. The sensitivity of this methodology is around 70% in bone metastases with breast cancer and 67% in prostate cancer.

What is also currently recommended for monitoring bone formation is N-terminal propeptide of collagen type 1 (P1NP), especially for osteoblastic metastases. P1NP is a highly specific marker showing the formation of osteoblastic (and also mixed) metastases; it shows little circadian variation, it can be detected even in serum and it has a relatively low intra-individual variation [5-8]. Should it be confirmed that patients with skeletal events show different kinetics in bone markers during treatment than patients without SRE, we would obtain another simple tool that would provide advance warning of an imminent risk of SRE, which is clearly a prognostic factor for shorter survival [9,10].

Material and Methods

From March 2009 to December 2011, we studied a group of 52 female patients with metastatic breast cancer (3 patients with only bone metastases, others had two or more organs affected, but in
The diagnosis of bone dissemination was determined both by means of bone scintigraphy, and confirmed by another imaging method - an X-ray or CT scan (3x NMR spine, 3x CT/PET). The patients received adequate systemic therapy based on the immunohistochemistry of the tumor, i.e., hormone therapy or chemotherapy, and also the parenteral bisphosphonate (alternately ibandronate and zoledronic acid). The routine biochemical tests and blood count done prior to the initiating therapy also included taking laboratory markers of bone metabolism CTX and P1NP and measuring the bone mineral density (according to the T-score). These bone markers were then checked in three, six, nine and twelve months, and matched with the progress of the disease. The total monitoring time was fifteen months. During each examination, the use of analgesics was recorded, as well as the occurrence of sudden SRE or other complications; before the end of monitoring, densitometry was done again. See Table 1 for the Monitoring Flow Sheet. The evaluation deals separately with the two groups (with or without SRE), and compares the kinetics of the levels of both markers during treatment as well as the occurrence of adverse effects and complications during the course of treatment, and the number of deaths.

The treatment set

A group of 52 women at the age from 31 to 81 (the median was 63.5 years) with metastatic breast cancer and bone metastases. The monitoring period was from March 2009 to December 30, 2011. All patients were examined by scintigraphy, with a negative finding in 2 patients with clinical symptoms, where the finding was confirmed by CT examination of the site according to clinical symptoms.

Treatment by bisphosphonates

After the first examination, patients were alternately treated by zoledronic acid or ibandronate. Ibandronate was replaced by zoledronic acid only in five patients with reduced left ventricular ejection fraction and in the treatment of a light heart failure, due to the volume of hydration administered.

Antineoplastic treatment

In all cases there was first-line palliative treatment applied. It was administered according to the immunoprofile: the hormonal treatment was applied with a steroid receptor positivity of over 10% and negativity of HER 2 neu; in other cases, chemotherapy treatment was administered.

Hormonal therapy

It was administered in thirty patients using tamoxifen (5 patients) or the aromatic inhibitor (AI) in 19 patients, and selective destructor of steroid receptor (SERMD - fulvestrant) in one patient. In four patients, letrozole was replaced by anastrozole during the treatment, in order to achieve a better profile of side effects and better tolerability.

Antineoplastic chemotherapy

It was administered in 33 patients. In 4 patients there was apositive HER 2 neu receptor; in one, treatment started in taxane/ trastuzumab regimen, and in three patients there was the so-called triple-positive immunoprofile, so in the first line letrozole/herceptin was administered. 20 patients were triple-negative, of which one was treated by avastin+paclitaxel with a good result. Chemotherapy was administered according to the patient’s general condition, laboratory values, left ventricular ejection fraction, previous treatment, effect and tolerability.

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Table 1: Monitoring Flow Sheet.

Doxorubicin 60 mg/m² and cyclophosphamide 600 mg/m² interval 21 days, AT: doxorubicin 50 mg/m² and docetaxel 75 mg/m² 3x int 21 days, CBDA/GEM Carboplatin AUC 5 and gemcitabine 800 mg/m²/day 1-8x int 21 days, CMF cyclophosphamide 600 mg/m² and fluorouracil 600 mg/m² and methotrexate 40 mg/m² day 1 into 21 days. Pain management was carried out in accordance with a subjective evaluation of pain, with the help of the VAS visual scale, based on the WHO scales; however, the main measurable variable was the consumption of analgesics, converted to morphine units. Treatment was initiated mostly by non-steroidal analgesics (NSE) - mostly paracetamol+codeine or tramadol as tolerated, then continued with opiates in a combination of short-term ones for breakthrough pain, and long-term ones usually administered in the form of a patch. In total, opioid treatment was administered to 31 patients, i.e. 59.6%. Radiotherapy with analgesic intention was carried out immediately after the start of monitoring in six patients, and orthopedicsurgery was carried out in five patients (spinal stabilization twice, pathological fracture of the femoral neck four times).

The Results

The following conditions were considered to be SRE

Fractures, necessity of radiotherapy, transverse spinal cord lesions, hypercalcinia. Female patients were evaluated in the course of fifteen months during the study. In 8 patients it was necessary to reduce the dosage of bisphosphonates by 50% due to light progression (5x Zometa and 4x ibandronate), and the established therapy continued even with the progression. No statistically significant difference was found between the two bisphosphates during the time before SRE occurred. According to the chi-square log rank test, the significance level was 0.4007, and according to the Wilcoxon test the p-value was 0.4904. Clear prognostic factors for patients overall survival (OS) include, in addition to the stage of the disease, the immunohistochemical status and tumor grading, which explains the statistically significant difference between patients treated by chemotherapy or hormonal therapy. HR was 8.777 at 95% confidence limits (1.056-72.931). In patients without hormonal treatment, the risk of SRE is 8.8 times higher. The difference in OS is shown in Chart 1.

The age of patients is related to the metabolic state of the bones, to the occurrence of osteopenia, osteoporosis and pelvis fractures. In the monitored period of fifteen months, no differences in OS were found based on the age.

Evaluation of the group after termination of treatment after twelve months

17.3% of patients died, which means that 82.7% of patients were still alive, out of which 46.2% had no fracture. Patients with the occurrence of SRE accounted for 53.5%, of which 7.1% of patients had a transverse
spinal lesion (2 patients). In both cases it was the primo-manifestation of the disease which was already present at the beginning of the study. Furthermore, a fracture was observed in seven patients (twice in the lower arm of the pubic bone, once in the femur, once in the forearm and three times in a vertebra without dislocation); in all cases, radiotherapy was administered. Only radiotherapy was administered against pain in six other patients (without a fracture). Orthopedic surgery was performed twice during the treatment and six times upon admission. During the treatment, death occurred in 17.3% of patients, of which one was diagnosed (post mortem) with double malignancy (a malignant melanoma of the right retina with dissemination to CNS); 3.6% of patients were diagnosed with ischemic stroke (despite early diagnosis without thrombolysis due to the disseminated malignancy) and 10.7% of patients died of the progression of the disease. Pulmonary embolism was observed in 25% of cases.

In this group, 25% of patients had to have their doses of analgesics increased by 1-2 morphine units per day. The number of patients with SRE reached 25%; in 21% of cases a palliative radiotherapy regimen was implemented (10 x 3 Gy); 2 patients were diagnosed with transverse spinal cord lesion (7.1%). Samples drawn from both groups of patients (with and without fractures) were compared at each stage. The first samples were drawn before the therapy was initiated, and then the others at each stage. The development of marker level kinetics was also compared between the two groups. According to the Cox analysis, the significance level of p-value was 0.0452 and HR was 8.516 (95% CI 1.047 to 69.262). Therefore, the recorded difference is not statistically significant, although the graphic representation shows a difference. The comparison of the first CTX sampling with overall survival was found to be significant. The patients whose CTX value in the first sampling was less than 0.425 ran an 8.5 times higher risk of death. Both groups of P1NP were evaluated after the first sampling, prior to the initiation of therapy. According to the Cox analysis, the significance level of the p-value was 0.0433 and HR was 8.673 (95% CI 1.067 to 70.520). Therefore, the recorded difference was not statistically significant, although the graphic representation shows a difference. However, it is lower than CTX. The comparison of the first P1NP samples with overall survival was found to be significant. The patients whose P1NP value in the first sampling was higher than 74 run 8.7 times higher risk of death. However, the difference (the p-value is 0.6131) does not reach statistical significance. None of the monitored parameters can be sufficiently discriminating for clinical practice to divide patients with SRE from those without it. With the minimum required specificity of 80%, the sensitivity reached was insufficient (Charts 2-5).

Discussion

This study confirms that hormone treatment is a significant predictor for the occurrence of SRE. Patients without hormonal therapy have 8.8 times higher risk of SRE, which probably results partly from the nature of cancer and also indicates the negative impact of chemotherapy-induced bone loss, while the role of age has not proved significant. The correlation of densitometrically-established osteoporosis and osteopenia according to the T-Score and bone markers was not statistically significant for the occurrence of SRE, which would correspond to an effective treatment of osteoporosis (both postmenopausal and treatment-induced) by parenteral bisphosphonates in effective doses. Therefore, the occurrence of SRE can be attributed to the bone cancer and not to osteoporosis. According to the data from literature, SRA usually occurs in 36–49% of patients treated for MBC with bone metastases during the first year of treatment; it usually takes 12 months before the first SRE occurs. In this study, there were 39.02% of SRE within 12 months. The patients whose
CTX value in the first sampling was less than 0.425 ran an 8.5 times higher risk of death; the patients whose P1NP value reached more than 74 in the first collection ran an 8.7 times higher risk of death. There is no clear explanation for this result; however, there has already been described a correlation between the higher values of P1NP in early breast cancer and the development of bone metastases and worse chances of survival [11], and in the TUGAMO study even a correlation between levels of serum CTX and P1NP in bone metastases of renal carcinoma [12].
Conclusion

The study failed to prove that testing fluctuating levels of bone markers CTX and P1NP is sufficiently specific and sensitive for predicting the occurrence of SRE in the first fifteen months of therapy. The patients whose CTX value in the first sampling was less than 0.425 ran an 8.5 times higher risk of death, and the patients whose P1NP value reached more than 74 in the first collection ran an 8.7 times higher risk of death. Therefore the value of the first sample was a negative predictive factor for survival. It cannot be confirmed that the levels of markers of bone resorption and remodeling actually copied the effect of treatment on the bone and that they were predictors of the risk of SRE, in correlation with the kinetics of the levels. The issue of the clinical significance of bone markers and selecting the most suitable ones is still under clinical testing and it cannot yet be recommended for current clinical practice when monitoring tumor bone disease.

References