KURAに登録されているコンテンツの著作権は、執筆者、出版社（学協会）などが有します。
KURAに登録されているコンテンツの利用については、著作権法に規定されている私的使用や引用などの範囲内で行ってください。
著作権法に規定されている私的使用や引用などの範囲を超える利用を行う場合には、著作権者の許諾を得てください。ただし、著作権者から著作権等管理事業者（学術著作権協会、日本著作出版権管理システムなど）に権利委託されているコンテンツの利用手続については、各著作権等管理事業者に確認してください。

Title
Early prediction of histopathological tumor response to preoperative chemotherapy by Tc-99m MIBI imaging in bone and soft tissue sarcomas

Author(s)
Taki, Junichi; Inaki, Anri; Wakabayashi, Hiroshi; Sumiya, Hisashi; Tsuchiya, Hiroyuki; Zen, Yoh; Kinuya, Seigo

Citation
Clinical Nuclear Medicine, 35(3): 154-159

Issue Date
2010-03

Type
Journal Article

URL
http://hdl.handle.net/2297/23484
Early Prediction of Histopathological Tumor Response to Preoperative Chemotherapy by Tc-99m MIBI imaging in Bone and Soft Tissue Sarcomas

Abbreviated title: Prediction of chemotherapeutic effect by Tc-99m-MIBI

Original Manuscript

Junichi Taki, MD, PhD¹, Anri Inaki, MD², Hiroshi Wakabayashi, MD², Hisashi Sumiya, MD, PhD¹, Hiroyuki Tsuchiya, MD, PhD³, Yoh Zen, MD, PhD⁴, Seigo Kinuya, MD, PhD⁵

¹Department of Nuclear Medicine, Kanazawa University Hospital, ²Department of Biotracer Medicine, Kanazawa University Graduate School of Medical Sciences, Kanazawa, Japan; ³Department of Orthopedic Surgery, Kanazawa University Graduate School of Medical Sciences, Kanazawa, Japan; and ⁴Division of Pathology, Kanazawa University Hospital, Kanazawa, Japan

Correspondence and reprint requests:

Junichi Taki
Department of Nuclear Medicine, Kanazawa University Hospital,
13-1 Takara-machi, Kanazawa, 920-8640, Japan
E-mail; taki@med.kanazawa-u.ac.jp
Tel; +81-76-265-2333, Fax;+81-76-234-4257,

There are no financial disclosures from any authors.
Abstract.

**Purpose:** Tc-99m-MIBI accumulates in only viable cells. In patients with bone and soft tissue sarcomas, preoperative chemotherapy is essential and the early prediction of the tumor response to chemotherapy would be beneficial for the planning of treatment strategy. The purpose of this study was to assess whether the change of Tc-99m-MIBI image from pre-chemotherapy to the early to middle of the chemotherapy can predict final histopathological tumor response as accurately as the change of the image after completion of chemotherapy.

**Methods:** Seventy-three patients with bone and soft tissue sarcomas underwent Tc-99m-MIBI scintigraphy before chemotherapy and at least 2 times after the second or third or fifth chemotherapy. The change of the tracer uptake ($\Delta UR$) and perfusion ($\Delta PI$) from pre-chemotherapy to post-chemotherapy were compared with histologic response.

**Results:** The sensitivity, specificity, and accuracy for the prediction of effective chemotherapy in $\Delta PI$ were 88%, 83%, 85% after 2nd, 85%, 72%, 78% after 3rd, and 81%, 71%, 76% after 5th chemotherapy, and those in $\Delta UR$ were 88%, 83%, 85% after 2nd, 85%, 92%, 89% after 3rd, and 94%, 77%, 85% after 5th chemotherapy, respectively. The area under the receiver operator characteristic curve of the $\Delta PI$ after 2nd, 3rd, and 5th chemotherapy were similarly good (0.842, 0.858, 0.811, respectively) and those of $\Delta UR$ were similarly excellent (0.915, 0.936, 0.931, respectively).

**Conclusion:** In patients with bone and soft tissue sarcomas, the change of Tc-99m-MIBI image from pre-chemotherapy to early to middle of chemotherapy can predict the final histopathological tumor response to chemotherapy as accurately as the change of Tc-99m-MIBI image from pre-chemotherapy to the completion of the preoperative chemotherapy.
Key Words: bone and soft tissue sarcoma; Tc-99m-MIBI; preoperative chemotherapy; treatment monitoring; osteosarcoma; Ewing’s sarcoma.

Introduction

In patients with bone and soft tissue sarcomas, the combination of aggressive chemotherapy before surgical intervention and limb-sparing surgery permits excellent quality of life and dramatic improvement of the prognosis. The accurate evaluation of chemotherapeutic effects is essential for proper treatment strategy because a good response to preoperative chemotherapy is crucial to limb-sparing surgery. In addition, early prediction of the tumor response to chemotherapy during preoperative chemotherapy would be beneficial for the planning of treatment strategy. Furthermore, noninvasive and quantitative assessment of tumor response to chemotherapy may provide prognostic information that could be useful for subsequent patient’s management, because good pathological response to chemotherapy has been proved to be a predictor of good prognosis in osteosarcoma.

Recently, there has been growing interest in using Tc-99m-hexakis-2-methoxyisobutylisonitrile (MIBI), used as a myocardial perfusion imaging agent, for tumor imaging including bone and soft-tissue tumors. As a tumor imaging agent Tc-99m-MIBI has unique characteristics. Administration of high radionuclide activity permits radionuclide angiography, which provides information of tumor blood flow. After Tc-99m-MIBI accumulation to the tumor, the degree of washout of the tracer from the tumor reflects P-glycoprotein overexpression which implies multidrug resistance. In addition, several studies have revealed that the change of Tc-99m-MIBI uptake from pre-chemotherapy to post-chemotherapy could reflect the chemotherapeutic effect in
patients with bone sarcomas.\textsuperscript{8,12-14} However, there has been no report that investigated whether Tc-99m-MIBI imaging performed during preoperative chemotherapy as well as performed after the completion of the chemotherapy can equally well predict final tumor response to preoperative chemotherapy.

Therefore, the purpose of this study was to assess whether the combination of Tc-99m-MIBI scintigraphy performed before and in the early to middle of the preoperative chemotherapy can predict final histopathological tumor response to chemotherapy without deterioration of the diagnostic accuracy compared with the combination of the imaging performed before chemotherapy and after final cycle of chemotherapy in bone and soft tissue sarcomas.

**MATERIALS AND METHODS**

**Patients**

Seventy three patients (54 men and 19 women; age range, 7 to 74 years, average age, 28.8 ± 18 years) with various bone and soft-tissue sarcomas proven pathologically in specimens obtained by biopsy at pretreatment and operation after chemotherapy were recruited in this study, and all gave informed consent to participate in this study. There were 36 osteosarcomas (28 osteoblastic type, 5 chondroblastic type, 2 fibroblastic type, and 1 small cell osteosarcoma), 7 malignant fibrous histiocytomas (5 pleomorphic type, 1 spindle cell type, 1 myxoid type), 7 liposarcomas (6 myxoid liposarcoma and 1 pleomorphic liposarcoma), 6 Ewing sarcomas, 4 synovial sarcomas, 3 chondrosarcomas (2 mesenchymal chondrosarcoma and 1 extraskeletal myxoid chondrosarcoma), 3 rhabdomyosarcomas, 2 angiosarcomas, 2 leiomyosarcomas, 1 malignant schwannoma, 1 alveolar soft part sarcoma, and 1 malignant giant cell tumor. Sixty-three lesions were located in the extremities, 5 were located in pelvic area, 2 were located at shoulder, 2 were in ribs, 1 was at axilla.
Forty-six tumors were located in bone and 27 were in soft tissue.

Patients received 5 cycles of preoperative chemotherapy at intervals of 2–3 weeks. In each chemotherapy cycle, cisplatin (120 mg / m²) was continuously infused through the intra-arterial catheter for 1 to 2 hours followed by 48 hours continuous infusion of doxorubicin (60 mg / m²) and 72 hours infusion of caffeine.¹⁵,¹⁶

**Study Protocol and Tc-99m-MIBI Scintigraphy**

The study protocol is shown in figure 1. All patients underwent imaging with Tc-99m-MIBI at least 3 times. Before chemotherapy all patients underwent the imaging with Tc-99m-MIBI (n=73), and after 1 week of the completion of the 2 (n=20) or 3 (n=69) or 5 cycles (n=66) of chemotherapy, same radionuclide imaging procedures were performed. After the completion of the 5 cycles of chemotherapy surgical operation was performed. Then the scintigraphic predictive power for the chemotherapeutic effect was assessed in comparison with the results of the postoperative histological examination.

Radionuclide angiography was performed after bolus injection of 600 - 740 MBq (16.2 – 20 mCi) of Tc-99m-MIBI with a gamma camera equipped with a low energy high-resolution parallel-hole collimator. Data were acquired every 2 sec for 2 min. Then, planar 2-min Tc-99m-MIBI image of the lesion was obtained at 15 min after radionuclide administration with 256 x 256 matrices.

**Image Analysis**

In the quantitative analysis of the pre-chemotherapeutic image, a manual region of interest (ROI) was set on the lesion and an automatically generated symmetrical ROI was set on the contralateral normal area as a background. When tumor Tc-99m-MIBI uptake is weak, the ROI of the tumor was delineated carefully to cover the whole lesion by referring to the computed tomography or magnetic resonance image. In the analysis of the images obtained after 2, 3 or 5 cycles of chemotherapy,
tumor ROI was newly set to cover the whole lesion as described above. Therefore, ROI size was changed depending on the size of the tumor. When the tumor uptake of radionuclide disappeared completely the same ROI delineated at prechemotherapy was applied to calculate the indices. Tc-99m-MIBI uptake ratio (UR) were calculated by dividing the count density of the lesion by that of the background ROI. For the prediction of the chemotherapeutic effect, the percent reduction of uptake ratio (ΔUR) was calculated as follows:

\[ \Delta UR (\%) = 100 \left( \frac{\text{prechemotherapy UR} - \text{postchemotherapy UR}}{\text{prechemotherapy UR}} \right) \]

The perfusion index (PI) was obtained by radionuclide angiography with Tc-99m-MIBI. Using the same ROI set to calculate the uptake ratio, the time-activity curve of each ROI was generated, and the PI was determined by dividing the peak count of the arterial phase of the lesion by that of the background ROI. When a peak count was not obtained, the time activity curve always showed a shoulder point, which was the flexion point between the rapid count increase due to the arterial phase and steady state or gradual count increase due to Tc-99m-MIBI accumulation to the lesion and normal tissue. Therefore, the count of the flexion point of the time activity curve was used to calculate the PI. Percent reduction of perfusion index (ΔPI) was also calculated to predict the response to chemotherapy by the following formula:

\[ \Delta PI (\%) = 100 \left( \frac{\text{prechemotherapy PI} - \text{postchemotherapy PI}}{\text{prechemotherapy PI}} \right) \]

**Assessment of the Chemotherapeutic effect by Histopathology**

After 5 cycle of chemotherapy surgical resection of the tumor was performed. Post-operative histological assessment was performed in all patients to evaluate the response of the tumor to chemotherapy. Histological grading of the effect of chemotherapy was determined based on the degree of viable cellularity and necrosis in the largest slice of the resected tumor. Grade IV (100% necrosis) and grade III (90%≤ to <100% necrosis) response were defined as good response to
chemotherapy (effective chemotherapy). Grade II (50%< to <90% necrosis) and Grade I (≤50% necrosis) responses were defined as poor response.

Statistics

Values are expressed as the mean ± SD. Differences in PIs and URs, and ∆PIs and ∆URs between good and poor response were analyzed by student t-test using the software JMP-5.0.1J for Macintosh computer. To appraise the discrimination abilities of the ∆PIs and ∆URs for the prediction of effective chemotherapy, receiver operator characteristic (ROC) curves were generated, and the areas under the curves (A_z) were calculated. The differences between the areas under the 2 ROC curves were examined using the 1-tailed paired t test.17,18 The optimal cutoff levels were identified as each index value minimizing the total number of false results.19 P < 0.05 was considered statistically significant. ROC analysis was performed using the software ROCKIT 0.9B (free software developed by Charles E. Metz, PhD for Macintosh computer).17,18

RESULTS

Post-operative histological examination of the resected tumors demonstrated good response in 33 patients and poor response in 40 patients. Good responders included 20 osteosarcomas, 3 malignant fibrous histiocytomas, 2 liposarcomas, 3 Ewing sarcomas, 1 synovial sarcomas, 1 rhabdomyosarcoma, 1 leiomyosarcoma, 1 malignant schwannoma, and 1 malignant giant cell tumor. Poor responders included 16 osteosarcomas, 4 malignant fibrous histiocytomas, 5 liposarcomas, 3 Ewing sarcomas, 3 synovial sarcomas, 3 chondrosarcomas, 2 rhabdomyosarcomas, 2 angiosarcomas, 1 leiomyosarcoma, and 1 alveolar soft part sarcoma.

All lesions showed significant tracer uptake. Accordingly, ROIs could be set without difficulty. In 2 patients after 3 cycles of chemotherapy and in 1 patient after 5 cycles of chemotherapy, PI of
Tc-99m-MIBI was not calculated because of poor bolus infusion of the radionuclide.

Before preoperative chemotherapy, there are no differences between the patients with good and poor response in PI (3.97 ± 2.51 and 3.38 ± 1.68, respectively, P=ns)) and UP (2.83 ± 0.93 and 2.74 ± 1.38, respectively, P=ns). After 2, 3, and 5 cycles of chemotherapy, in patients with good response, PI decreased to 1.81 ± 0.61 (P<0.01), 1.97 ± 1.25 (P<0.01), and 1.51 ± 0.76 (P<0.01), respectively, and UR also decreased to 1.56 ± 0.46 (P<0.01), 1.58 ± 0.51 (P<0.01), and 1.34 ± 0.46 (P<0.01), respectively. In patients with poor response, after 2, 3, and 5 cycles of chemotherapy, both PI (3.88 ± 2.17, 3.70 ± 2.77, 2.92 ± 2.48, respectively) and UR (2.64 ± 1.14, 2.61 ± 1.22, 2.19 ± 1.02, respectively) did not change significantly. Consequently, after 2, 3, and 5 cycles of chemotherapy, good responders demonstrated lower PI (P<0.05, P<0.01, P<0.01, respectively) and lower UR (P<0.05, P<0.01, P<0.01, respectively) than poor responders.

The ∆PI after 2 cycles of chemotherapy in patients with good and poor response were 36 ± 23% and -12 ± 42% (P < 0.01), respectively (Fig. 2A). The ∆PIs after 3 and 5 cycles of chemotherapy in patients with good and poor response were 45 ± 27% and -8.3 ± 50% (P < 0.0001), respectively (Fig. 2B), and 55 ± 22% and 19 ± 37% (P < 0.0001), respectively (Fig. 2C). The ∆UR obtained after 2 cycles of chemotherapy in patients with good and poor response were 34 ± 16% and -8 ± 32% (P=0.005), respectively (Fig. 3A), and those after 3 and 5 cycles of chemotherapy were 43 ± 14% and -0.7 ± 39% (P < 0.0001), respectively (Fig. 3B) and 52 ± 13% and 15 ± 27% (P < 0.0001), respectively (Fig. 3C).

The ROC curves for the predictability of effective chemotherapy of the ∆PI and ∆UR obtained after 2, 3, and 5 cycles of chemotherapy are shown in figure 4. The A of the ∆PI after 2, 3, and 5 cycles of chemotherapy were 0.842, 0.858, and 0.811, respectively and there were no significant differences among the values (P=0.13–0.49). The A of the ∆UR after 2, 3, and 5 cycles of
The accuracy of ∆PI after 2, 3, and 5 cycles after chemotherapy was 85%, 78%, 76%, respectively, and that of ∆UR after 2, 3, and 5 cycles after chemotherapy was 85%, 89%, 85%, respectively.

On the contrary, the accuracy of PI after 2, 3, and 5 cycles after chemotherapy was 70%, 69%, 70%, respectively, and that of UR after 2, 3, and 5 cycles after chemotherapy was 70%, 69%, 72%, respectively.

Representative cases are shown in Fig 5 (good responder) and Fig 6 (poor responder).

DISCUSSION

The present study demonstrated that the combination of the Tc-99m-MIBI scintigraphies performed before chemotherapy and in the early to middle of preoperative-chemotherapy (after the 2nd and 3rd cycle) as well as after final cycle of chemotherapy (5th cycle) can accurately predict the final response to chemotherapy which was proved by histopathology after 5 cycles of chemotherapy and operation. Based on the ROC analysis, both the performance of ∆PI and ∆UR for the prediction of the chemotherapeutic effect was similarly good after 2, 3 and 5 cycle of chemotherapy. Therefore accurate prediction of the chemotherapeutic effect can be possible during
early to middle of chemotherapy. However, when ΔPI and ΔUR was compared, predictive performance of ΔUR tended to be better than that of ΔPI after 2 and 3 cycle of therapy and significantly better after the final cycle of preoperative chemotherapy.

On the contrary the diagnostic accuracies for the prediction of chemotherapeutic effect of PI and UR itself after 2, 3, and 5 cycles of chemotherapy were 69% ~ 72%, and those were inferior to that of ΔPI and ΔUR (76% ~ 89%, Table 1).

Tc-99m-MIBI has been introduced primarily as a myocardial perfusion imaging agent to replace Tl-201. Tc-99m-MIBI has lipophilic cationic properties, accumulating largely in mitochondria by its negative trans-membrane potential. Moreover, its accumulation depends on the cell viability and metabolic status in myocardial cells. Like myocardial cells, human carcinoma cells accumulate Tc-99m-MIBI by depending on the mitochondria and plasma membrane potentials. In addition, radionuclide angiography with sufficient dose at the administration of the tracer can permit evaluation of the tumor blood flow. Accordingly, Tc-99m-MIBI scintigraphy provide integrated information of the tumor blood flow and tracer uptake which reflect tumor viability and metabolic status.

For the evaluation of the tumor response to preoperative chemotherapy, several modalities are performed such as CT, MRI, angiography, ultrasonography, and radionuclide imaging. Conventional angiography with contrast medium can observe tumor blood flow. Although, it can offer precise morphology of tumor vasculature, quantification of the tumor blood flow is somewhat difficult compared to radionuclide angiography. CT and MRI can precisely evaluate the tumor size. However, they cannot evaluate tumor viability and metabolic status so precisely even with contrast enhancement because fibrotic change, inflammatory change, or granulation tissue secondary to tumor necrosis after chemotherapy could be enhanced. Dynamic contrast-enhanced MRI provides
information about tissue perfusion, vascular permeability, micro-vessel density and may differentiate viable tumor tissue from reparative tissue. Several parameters by mathematical quantification of the dynamic data permit prediction of the chemotherapeutic response after the completion of the preoperative chemotherapy. However, early prediction of chemotherapeutic effect by dynamic MRI remains to be elucidated. The parameter calculated by dynamic MRI might mainly reflect tumor perfusion and might be similar to that obtained by radionuclide angiography with Tc-99m-labeled tracers. Color Doppler ultrasound also could predict chemotherapeutic response by assessing the change of tumor blood supply and intratumoral blood flow after chemotherapy. The method is relatively simple but further evaluation would be warranted including the issue of the reproducibility. Several FDG-PET studies has also demonstrated the efficacy in the prediction of the effect of chemotherapy at the end of preoperative chemotherapy, however, sensitive FDG accumulation in chemotherapy induced inflammation or therapy related fibrous tissue has impaired the diagnostic accuracy in some cases. Unfortunately, there has been no report in terms of the early prediction of preoperative chemotherapy using FDG PET in bone and soft tissue sarcomas.

In a previous study with 12 osteosarcoma patients, reduction of Tc-99m-MIBI uptake after completion of chemotherapy roughly correlated to the histological response. Another studies with 28 bone sarcoma patients and 68 patients with bone and soft tissue sarcoma demonstrated good correlation between the reduction of Tc-99m-MIBI uptake after 3 or 4 cycles of chemotherapy and % necrosis of the tumor and good predictability of tumor response to chemotherapy. In the present study, in both bone and soft tissue sarcomas, reduction of tumor perfusion and Tc-99m-MIBI uptake in the early to middle of chemotherapy as well as after final cycle of chemotherapy can predict the final histological tumor response to chemotherapy. In the clinical
setting, a precise diagnostic method for the assessment of the chemotherapeutic effect at the earliest possible time after initiation of chemotherapy would be preferable. Our current study demonstrated that early imaging could permit prices estimation of chemotherapeutic effect after 2 or 3 cycle of preoperative chemotherapy as accurately as after final cycle of therapy, suggesting that the method may have an impact on modification of neoadjuvant chemotherapy protocol, on patient selection for the performance and timing of limb-sparing surgery, and on the selection of postoperative chemotherapy regimens.

MIBI has been proved to be the one of the substrates of P-glycoprotein, a product of the human multidrug resistance gene and multidrug resistance related protein.\textsuperscript{37,38} It has been proved that Tc-99m-MIBI once accumulated in tumor cells would be washed out from the cells by P-glycoprotein and the delayed Tc-99m-MIBI uptake was related to P-glycoprotein expression, while early uptake was not.\textsuperscript{10} Accordingly, early Tc-99m-MIBI uptake seems to reflect tumor viability more properly than delayed uptake. Therefore, as a marker of tumor viability, we analyzed only Tc-99m-MIBI uptake in early image obtained at 15 min after tracer injection.

For the assessment of tumor perfusion and Tc-99m-MIBI uptake, tumor ROI was set in each image before and after chemotherapy independently, i.e., new ROI was delineated after chemotherapy if tumor size changed to cover whole tumor more precisely instead of using the same ROI used before chemotherapy. This is because we assessed the histologic grading of chemotherapeutic effect based on the degree of cellularity and necrosis in the largest slice of the resected tumor as a gold standard of tumor response to chemotherapy. Accordingly ROI modification along with the change of the tumor size may reflect more precisely the tumor viability based on the histopathological examination.

Possible drawbacks of Tc-99m-MIBI include its high uptake by the liver and excretion to the
intestine and urinary system, which interfere with abdominal and pelvic evaluation. However, lesions of the extremities, where most bone and soft tissue sarcomas are found, are unaffected by these issues.

Since there was a great variety of tumor types included in this study, there was some dispersion in Tc-99m-MIBI tumor uptake. However, since the tumor Tc-99m-MIBI uptake reflects tumor viability and metabolic status, the degree of the change of tumor viability by chemotherapy might be reflected similarly to the change of Tc-99m-MIBI uptake among various tumor types.

CONCLUSION

The study demonstrates that the change in tumor perfusion and Tc-99m-MIBI uptake from pre-chemotherapy to early to middle of preoperative chemotherapy as well as to the final cycle of chemotherapy is highly predictive of histopathological tumor response to chemotherapy in bone and soft tissue sarcomas. This imaging technique, especially in AUR, might be a beneficial method in patient care and in elaborating treatment strategy during preoperative chemotherapy.
REFERENCES


FIGURE LEGENDS

FIGURE 1
Study protocol. Tc-99m-MIBI scintigraphy was performed before chemotherapy in all patients, and at least 2 scintigraphic studies were performed after 2, 3, or 5 cycle of chemotherapy. The change of perfusion index and uptake ratio from prechemotherapy to after each chemotherapy cycle were compared to the histopathological response to chemotherapy.

FIGURE 2
Percent reduction of perfusion index (PI) obtained from radionuclide angiography with Tc-99m-MIBI after 2 cycle (A), 3 cycle (B), and 5 cycle (C) of chemotherapy in patients with good response and poor response. Horizontal dashed line indicates threshold line derived from receiver operator characteristic curve analysis that minimized the total number of false results.

FIGURE 3
Percent reduction of Tc-99m-MIBI uptake ratio (UR) after 2 cycle (A), 3 cycle (B), and 5 cycle (C) of chemotherapy in patients with good response and poor response. Horizontal dashed line indicates threshold line derived from receiver operator characteristic curve analysis that minimized the total number of false results.

FIGURE 4
Receiver operator characteristic (ROC) curve analysis for the prediction of tumor response to chemotherapy using ΔPI and ΔUR. The areas under the ROC curve (A$_z$) of the percent change of
perfusion index (ΔPI) at after 2 cycle (0.842), 3 cycle (0.858), and 5 cycle (0.811) of chemotherapy were similar (A). The A of the percent change of Tc-99m-MIBI uptake ratio (ΔUR) at after 2 cycle (0.915), 3 cycle (0.936), and 5 cycle (0.931) of chemotherapy were also similar (B).

FIGURE 5
A 50-year-old female with pleomorphic malignant fibrous histiocytoma of the left shoulder. Before chemotherapy, radionuclide angiography with Tc-99m-MIBI demonstrated marked increase in tumor blood flow. Anterior chest image obtained 15min after the tracer injection showed intense accumulation of Tc-99m-MIBI in the left shoulder. After 2 cycles of chemotherapy, tumor blood flow and Tc-99m-MIBI uptake decreased dramatically. After 3 and 5 cycles of chemotherapy, no significant increase in tumor blood flow and Tc-99m-MIBI uptake was observed. All these findings implied a good response to chemotherapy. Wide resection of tumor was performed after 5 cycles of chemotherapy. Pathological study of the resected tumor specimen revealed no viable tumor cells.

FIGURE 6
A 16-year-old male with osteosarcoma of the right distal femur. Before chemotherapy, radionuclide angiography with Tc-99m-MIBI demonstrated marked increase in tumor blood flow and intense Tc-99m-MIBI uptake. After 2, 3, and 5 cycles of chemotherapy, tumor blood flow and tumor uptake of Tc-99m-MIBI were still increased. All these findings implied a poor response to chemotherapy. After 5 cycles of chemotherapy, resection of the tumor was performed and the pathological study of the resected tumor specimen revealed ≤ 50% tumor necrosis.
FIGURE 1

Tc-99m-MIBI imaging

1st chemotherapy

2nd chemotherapy

3rd chemotherapy

4th chemotherapy

5th chemotherapy

Operation and histopathological examination
FIGURE 3

A

B

C

% Reduction of UR after 2 cycle

% Reduction of UR after 3 cycle

% Reduction of UR after 5 cycle

P < 0.005

P < 0.001

P < 0.001

Good Response

Poor Response

Good Response

Poor Response

Good Response

Poor Response
**FIGURE 5**

<table>
<thead>
<tr>
<th>Radionuclide angiography with Tc-99m-MIBI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before Chemo</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tc-99m-MIBI 15min image</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before Chemo</td>
</tr>
</tbody>
</table>
FIGURE 6

<table>
<thead>
<tr>
<th>Radionuclide angiography with Tc-99m-MIBI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before Chemo</td>
</tr>
<tr>
<td>After 2 cycle of Chemo</td>
</tr>
<tr>
<td>After 3 cycle of Chemo</td>
</tr>
<tr>
<td>After 5 cycle of Chemo</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tc-99m-MIBI 15min image</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before Chemo</td>
</tr>
<tr>
<td>After 2 cycle of Chemo</td>
</tr>
<tr>
<td>After 3 cycle of Chemo</td>
</tr>
<tr>
<td>After 5 cycle of Chemo</td>
</tr>
</tbody>
</table>