

Table 4. Urinary excretion of capecitabine and its metabolites

| | Urinary excretion (% of dose) | | | |
|--------------|-------------------------------|---------------|--------|---------------|
| | Day 1 | | Day 14 | |
| | N | Mean ± SD | N | Mean ± SD |
| Capecitabine | 16 | 3.21 ± 2.04 | 19 | 3.42 ± 1.48 |
| 5'-DFCR | 16 | 8.39 ± 3.73 | 19 | 8.42 ± 3.44 |
| 5'-DFUR | 16 | 12.1 ± 4.34 | 19 | 14.6 ± 5.35 |
| 5-FU | 16 | 0.691 ± 0.835 | 19 | 0.782 ± 0.642 |
| FUPA | 16 | 2.78 ± 0.808 | 19 | 2.98 ± 1.05 |
| FBAL | 16 | 50.3 ± 9.66 | 19 | 49.5 ± 11.3 |
| Total | 16 | 77.5 ± 14.8 | 19 | 79.6 ± 16.9 |

5'-DFCR, 5'-deoxy-5-fluorocytidine; 5'-DFUR, 5'-deoxy-5-fluorouridine; FUPA, α -fluoro- β -ureidopropionic acid; FBAL, α -fluoro- β -alanine.

2 versus 13%) (12). Though pigmentation, which was not reported more than 5% in the phase III trials, was frequently observed in this study (38%), all events of pigmentation were grade 1 and did not lead to interruption or reduction. The rate of other adverse drug reactions in our study was almost identical to that reported in the phase III trials (12). With regard to severe abnormalities in laboratory parameters, AST elevation was more frequently observed in the present study (10 versus 1%), although the rate of hyperbilirubinemia was similar to phase III observations (12 versus 23%) (12). Dose reduction was executed more frequently than the phase III trials (53 versus 34%), but the rate of dose reduction to second level was almost similar (17 versus 12%). Median time to reduction to the first level was similar to phase III trials (2.6 months versus 2.5 months), and median time to reduction to the second level was longer in our study (5.3 months versus 3.6 months). From these results, the current 3-week regimen seems quite feasible for the treatment of MCRC in Japan.

The pharmacokinetic findings in the present study were basically similar to those reported in Caucasian patients (8,21). Pharmacokinetic analysis of plasma concentrations and urinary excretion showed rapid gastrointestinal absorption of capecitabine and efficient conversion to its metabolites. Peak concentrations of capecitabine and its metabolites, including 5-FU, were reached shortly after drug administration and declined exponentially with a half-life of approximately 1 h. Pharmacokinetic data obtained on days 1 and 14 showed no difference in pharmacokinetics over time and there was no clinically significant accumulation of capecitabine and its metabolites, except for 5-FU. The AUC of 5-FU on day 14 was 1.6 times higher than on day 1. A similar increase of 5-FU with multiple administration has been also reported in other clinical studies of capecitabine (7,8,21).

From these results, we conclude that the 3-week regimen of capecitabine is effective and well tolerated in Japanese patients with MCRC. Capecitabine has been reported to show good activity when combined with irinotecan (14,15)

and oxaliplatin (16,17). Further investigation of this 3-week schedule is warranted in Japan.

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References

- de Gramont A, Bosset JF, Milan C, Rougier P, Bouche O, Etienne PL, et al. Randomized trial comparing monthly low-dose leucovorin and fluorouracil bolus with bimonthly high-dose leucovorin and fluorouracil bolus plus continuous infusion for advanced colorectal cancer: a French intergroup study. *J Clin Oncol* 1997;15:808-15.
- Anon. Efficacy of intravenous continuous infusion of fluorouracil compared with bolus administration in advanced colorectal cancer. Meta-analysis Group In Cancer. *J Clin Oncol* 1998;16:301-8.
- Liu G, Franssen E, Fitch MI, Warner E. Patient preferences for oral versus intravenous palliative chemotherapy. *J Clin Oncol* 1997;15:110-5.
- Borner MM, Schoffski P, de Wit R, Caponigro F, Comella G, Sulkes A, et al. Patient preference and pharmacokinetics of oral modulated UFT versus intravenous fluorouracil and leucovorin: a randomised crossover trial in advanced colorectal cancer. *Eur J Cancer* 2002;38:349-58.
- Miwa M, Ura M, Nishida M, Sawada N, Ishikawa T, Mori K, et al. Design of a novel oral fluoropyrimidine carbamate, capecitabine, which generates 5-fluorouracil selectively in tumors by enzymes concentrated in human liver and cancer tissue. *Eur J Cancer* 1998;34:1274-81.
- Ishikawa T, Utoh M, Sawada N, Nishida M, Fukase Y, Sekiguchi F, et al. Tumor selective delivery of 5-fluorouracil by capecitabine, a new oral fluoropyrimidine carbamate, in human cancer xenografts. *Biochem Pharmacol* 1998;55:1091-7.
- Saeki T, Takashima S, Terashima M, Satoh A, Toi M, Osaki A, et al. A Japanese phase I study of continuous oral capecitabine in patients with malignant solid tumors. *Int J Clin Oncol* 2005;10:51-7.
- Mackean M, Planting A, Twelves C, Schellens J, Allman D, Osterwalder B, et al. Phase I and pharmacologic study of intermittent twice-daily oral therapy with Capecitabine in patients with advanced and/or metastatic cancer. *J Clin Oncol* 1998;16:2977-85.
- Kondo Y, Terashima M, Sato A, Taguchi T. A pilot phase II study of capecitabine in advanced or recurrent colorectal cancer. *Jpn J Clin Oncol* 2004;34:195-201.
- Sakamoto J, Kondo Y, Takemiya S, Sakamoto N, Nishisho I, on behalf of the clinical study group of capecitabine. A phase II Japanese study of a modified capecitabine regimen for advanced or metastatic colorectal cancer. *Anti-Cancer Drugs* 2004;15:137-43.
- Twelves C. Capecitabine as first-line treatment in colorectal cancer: pooled data from two large, phase III trials. *Eur J Cancer* 2002;38(Suppl. 2):15-20.
- Cassidy J, Twelves C, Van Cutsem E, Hoff P, Bajetta E, Boyer M, et al. Capecitabine Colorectal Cancer Study Group. First-line oral capecitabine therapy in metastatic colorectal cancer: a favorable safety profile compared with intravenous 5-fluorouracil/leucovorin. *Ann Oncol* 2002;13:566-75.
- Van Cutsem E, Hoff PM, Harper P, Bukowski RM, Cunningham D, Dufour P, et al. Oral capecitabine vs intravenous 5-fluorouracil and leucovorin: integrated efficacy data and novel analyses from two large, randomized, phase III trials. *Br J Cancer* 2004;90:1190-7.
- Tewes M, Schleucher N, Achterrath W, Wilke HJ, Frings S, Seeber S, et al. Capecitabine and irinotecan as first-line chemotherapy in patients with metastatic colorectal cancer: results of an extended phase I study. *Ann Oncol* 2003;14:1442-8.

15. Borner MM, Bernhard J, Dietrich D, Popescu R, Wernli M, Saletti D, et al. A randomized phase II trial of capecitabine and two different schedules of irinotecan in first-line treatment of metastatic colorectal cancer: efficacy, quality-of-life and toxicity. *Ann Oncol* 2005;16:282–8.
16. Diaz-Rubio E, Evans TR, Tabernero J, Cassidy J, Sastre J, Eatock M, et al. Capecitabine (Xeloda) in combination with oxaliplatin: a phase I, dose-escalation study in patients with advanced or metastatic solid tumors. *Ann Oncol* 2002;13:558–65.
17. Cassidy J, Tabernero J, Twelves C, Brunet R, Butts C, Conroy T, et al. XELOX (Capecitabine Plus Oxaliplatin): Active first-line therapy for patients with metastatic colorectal cancer. *J Clin Oncol* 2004;22:2084–91.
18. Fernando N, Yu D, Morse M, Blobe G, Odogwu L, Crews J, et al. A phase II study of oxaliplatin, capecitabine and bevacizumab in the treatment of metastatic colorectal cancer. *J Clin Oncol* 2005;23:6S(Abst. 3556).
19. Therasse P, Arbuck S, Eisenhauer E, Wanders J, Kaplan R, Rubinstein L, et al. New Guidelines to Evaluate the Response to Treatment in Solid Tumors. *J Natl Cancer Inst* 2000;92: 205–16.
20. National Cancer Institute—Common Toxicity Criteria (NCI-CTC Version 2.0, April 30, 1999).
21. Reigner B, Watanabe T, Schuller J, Lucraft H, Sasaki Y, Bridgewater J, et al. Pharmacokinetics of capecitabine (Xeloda) in Japanese and Caucasian patients with breast cancer. *Cancer Chemother Pharmacol* 2003;52:193–201.

APPENDIX

List of participating centers: NHO Shikoku Cancer Center, National Cancer Center Hospital, National Cancer Center Hospital East, Cancer Institute Hospital, Aichi Cancer Center, Saitama Cancer Center, Kobe University Graduate School of Medicine, Kanagawa Cancer Center, Osaka Medical College, Kinki University, NHO Osaka National Hospital.

Staging performance of carbon-11 choline positron emission tomography/computed tomography in patients with bone and soft tissue sarcoma: Comparison with conventional imaging

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The present study was conducted to compare the diagnostic accuracy between carbon-11 choline (¹¹C-choline) positron emission tomography (PET)/computed tomography (CT) and conventional imaging for the staging of bone and soft tissue sarcomas. Sixteen patients who underwent ¹¹C-choline PET/CT prior to treatment were evaluated retrospectively for staging accuracy. Conventional imaging methods consisted of ^{99m}Tc-hydroxymethylene diphosphonate bone scintigraphy, chest CT and magnetic resonance imaging of the primary site. The images were reviewed and a consensus was reached by two board-certified radiologists who were unaware of any clinical or radiological information using hard-copy films and multimodality computer platform. Tumor stage was confirmed by histological examination and/or by an obvious progression in number and/or size of the lesions on follow-up examinations. Reviewers examining both ¹¹C-choline PET/CT and conventional imaging classified T stage in all patients. Interpretation based on ¹¹C-choline PET/CT, the Node (N) stage was correctly diagnosed in all patients, whereas the accuracy of conventional imaging in N stage was 63%. Tumor Node Metastasis (TNM) stage was assessed correctly with ¹¹C-choline PET/CT in 15 of 16 patients (94%) and with conventional imaging in eight of 16 patients (50%). The overall TNM staging and N staging accuracy of ¹¹C-choline PET/CT were significantly higher than that of conventional imaging ($P < 0.05$). ¹¹C-choline PET/CT is more accurate than conventional imaging regarding clinical staging of patients with bone and soft tissue sarcomas. A whole body ¹¹C-choline PET/CT might be acceptable for imaging studies of tumor staging prior to treatment. (*Cancer Sci* 2006; 97: 1125–1128)

The general diagnostic tools for staging bone and soft tissue sarcomas are clinical examination, magnetic resonance imaging (MRI) and X-ray of the primary tumor site, chest X-ray or computed tomography (CT), and bone scintigraphy.⁽¹⁾

Positron emission tomography (PET) with [¹⁸F]-fluoro-2-deoxy-D-glucose (FDG) has been used in the evaluation of patients with bone and soft tissue sarcomas for grading and therapy monitoring.^(2–7) Most of these studies reveal that ¹⁸FDG-PET is superior in the assessment of grading and therapy monitoring compared with conventional imaging.

Recently, carbon-11 choline (¹¹C-choline) has been introduced as a new oncological positron-emitting radiopharmaceutical for evaluation of a variety of malignant tumors.^(8–11) Choline is an essential component of the cell membrane, and choline uptake may be via a choline-specific transporter protein.⁽¹²⁾ Choline kinase, which catalyzes the phosphorylation of choline, is upregulated in malignant cells. Some studies have demonstrated additional gains in diagnostic accuracy using ¹¹C-choline.⁽¹³⁾ ¹¹C-choline uptake is significantly higher in malignant tumors than in benign tumors and correlates well with the degree of ¹⁸FDG accumulation with

the lesion, while the high background activity owing to excretion via urinary tract interferes with evaluation on ¹⁸FDG-PET.^(14,15) However, the role of ¹¹C-choline PET scan in the staging of bone and soft tissue sarcomas has not been clarified. To fully elucidate the role of ¹¹C-choline PET, the comparison with ¹⁸FDG-PET and conventional imaging modalities are needed.

A new-modality PET/CT can improve the localization of tumors and accuracy of staging in patients because anatomic and molecular information can be coregistered precisely.⁽¹⁶⁾ The aim of the current study was to compare the diagnostic accuracy between ¹¹C-choline PET/CT and conventional imaging for the staging of bone and soft tissue sarcomas.

Materials and Methods

Patient. We retrospectively reviewed ¹¹C-choline PET/CT results from September 2005 to March 2006 for patients with bone and soft tissue sarcomas, who subsequently underwent surgical resection, chemotherapy and/or radiotherapy within 2 weeks. ¹¹C-choline PET/CT was performed for initial staging in 12 patients and for restaging of recurrent disease in four patients. The study population consisted of 13 men and three women with a mean age of 44 years (range, 13–75 years). The clinical records of all of the patients were available for review. This study was conducted in accordance with the amended Helsinki declaration and the protocol was approved by the Institutional Review Board (National Cancer Center, Research Center for Cancer Prevention and Screening). All of the patients provided their written informed consent to participate in the present study and to review their records and images.

Radiopharmaceuticals. Carbon-11 choline was synthesized with a commercial module essentially using the method described by Hara and Yuasa.⁽¹⁷⁾ ¹¹CO₂ was converted to ¹¹C-methyl iodide by LiAlH₄/HI reaction. ¹¹C-methyl iodide was trapped in dimethylaminoethanol. After a washing step with ethanol and water, ¹¹C-choline retained on a cation exchange resin was eluted with saline. Radiochemical purity of the solution was evaluated by liquid chromatography radiodetector. The organic solvents were analyzed by gas chromatography. Endotoxin was assayed by the lysosomal acid lipase method.

PET/CT. Scans were acquired with a PET/CT device (Aquiduo; Toshiba Medical Systems, Tokyo, Japan) that consisted of a PET scanner (ECAT HR+; CTI, Knoxville, TN, USA) and 16-section CT scanner (Aquilion V-detector; Toshiba Medical Systems) with a whole-body mode implemented as the standard software. Prior to the ¹¹C-choline PET/CT study, the patients fasted for at least

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Table 1. Summary of patients and confirmed staging

| Patient no. | Diagnosis | SUV | Size (mm) | Staging type | Location | TNM | Metastasis | Grade | Stage |
|-------------|--|-------|-----------|--------------|-----------------|----------|--------------------------------|-------|-------|
| 1 | Leiomyosarcoma | 4.63 | 110 | Initial | Retroperitoneum | T2bN0M1 | Soft tissue | High | IV |
| 2 | Rhabdomyosarcoma | 3.03 | 60 | Initial | Perineum | T2bN1M0 | Lymph node | High | IV |
| 3 | Pleomorphic malignant Fibrous histiocytoma | 15.05 | 133 | Initial | Chest wall | T2bN0M1 | Bone, pleura, lymph node | High | IV |
| 4 | Leiomyosarcoma | 4.10 | 80 | Initial | Retroperitoneum | T2bN0M,P | Lung | Low | IV |
| 5 | Osteosarcoma | 6.70 | 110 | Initial | Iliac bone | T2N0M1b | Bone, lung | High | IVB |
| 6 | Clear cell sarcoma | 13.03 | 80 | Initial | Chest wall | T2bN0M1 | Bone, lung, pleura, lymph node | High | IV |
| 7 | Myxoid liposarcoma | 2.15 | 50 | Initial | Leg | T1aN1M0 | Lymph node | Low | IVB |
| 8 | Osteosarcoma | 5.31 | 110 | Initial | Tibia | T2N1M0 | Lymph node | High | IV |
| 9 | Ewing sarcoma | 3.46 | 95 | Initial | Leg | T2bN0M0 | N/A | High | III |
| 10 | Ewing sarcoma | 9.86 | 102 | Initial | Shoulder | T2N0M0 | N/A | High | IIB |
| 11 | Ewing sarcoma | 6.14 | 16 | Initial | Spine | T1N0M0 | N/A | High | IA |
| 12 | Chondrosarcoma | 5.99 | 110 | Initial | Iliac bone | T2N0M1b | Bone | High | IVB |
| 13 | Leiomyosarcoma | 3.18 | 50 | Restaging | Thigh | T1bN1M1 | Bone, soft tissue, lymph node | High | IV |
| 14 | Osteosarcoma | 4.95 | 75 | Restaging | Jaw | T1N0M1a | Lung | High | IVA |
| 15 | Osteosarcoma | 3.60 | 50 | Restaging | Femur | T1N0M1b | Lung, bone | High | IVB |
| 16 | Alveolar soft part sarcoma | 3.60 | 25 | Restaging | Shoulder | T2N0M1 | Bone | High | IV |

N/A, not applicable; SUV, standardized uptake value; TNM, Tumor Node Metastasis.

6 h. CT was performed from the head to the mid-thigh according to a standardized protocol with the following setting: axial 3.0-mm collimation × 16 modes; 120 kVp; 100 mAs; and a 0.5-second tube rotation, pitch 11.0. Patients maintained normal shallow respiration during the three-dimensional acquisition of CT scans. No iodinated contrast material was administered. Emission scans from the base of the skull to the leg were obtained starting 5 min after the intravenous administration of 350–573 MBq of ¹¹C-choline. The acquisition time for PET was 2 min per table position. Images were reconstructed with attenuation-corrected ordered-subset expectation maximization with two iterations and eight subsets using emission scans and CT data.

Positron emission tomography, CT and coregistered PET/CT images were analyzed with dedicated software (e-soft; Siemens). The initial review of the attenuation-corrected PET images was performed using transaxial, coronal and sagittal planes. The images were reviewed and a consensus was reached by two board-certified radiologists who were unaware of any clinical or radiological information using a multimodality computer platform. ¹¹C-choline uptake was considered to be abnormal when it was substantially greater than the surrounding normal tissue. For ¹¹C-choline PET/CT, tumor sizes and T staging were determined by the CT part of PET/CT. ¹¹C-choline-avid lymph nodes or distant metastases on PET/CT were interpreted as positive for metastases regardless of size. Lymph nodes with abnormal uptake were deemed positive for metastases even when they were smaller than 10.0 mm in short axis nodal diameter. Lung nodules without abnormal uptake but highly suggestive of lung metastases on ¹¹C-choline PET/CT were considered to be positive for metastases. A pixel region of interest (ROI) was outlined within regions of increased ¹¹C-choline uptake and measured on each slice. For quantitative interpretations, standardized uptake value (SUV) was determined according to the standard formula, with activity in the ROI given in Bq per mL/injected dose in Bq per weight (kg). However, time decay correction for whole-body image acquisition was not conducted. A SUV of more than 2.5 was considered to characterize malignancy.

Conventional imaging. Conventional imaging methods, performed within 2 weeks of ¹¹C-choline PET/CT, either before or after, were ^{99m}Tc-hydroxymethylene diphosphonate (HMDP) bone scintigraphy, chest CT and MRI of the primary site. ^{99m}Tc-HMDP bone scintigraphy was performed 2 h after intravenous injection of 740 MBq of ^{99m}Tc-HMDP. Both anterior and posterior

whole-body planar images were obtained simultaneously with a dual-headed gamma camera (E.CAM; Siemens). Chest CT was performed using a multidetector scanner (Aquilion V-detector; Toshiba Medical Systems) with the following setting: axial 4.0-mm × 4 modes; 120 kVp, automated electric current; 0.5-second tube rotation; and pitch 5. Images were reconstructed with 10.0-mm slice thickness by means of a standard algorithm. MRI of the primary site was performed using a 1.5 Tesla system (Signa Horizon; GE Medical Systems, Milwaukee, WI, USA or Visart; MRI produced by Toshiba Medical Systems, Tokyo, Japan). Pulse sequences comprised T1-weighted spin echo (SE) images, T2-weighted fast spin echo (FSE) images, as well as post-contrast T1-weighted SE images with fat suppression after injection of contrast material. Pulse sequence parameters and slice orientation varied with the examined anatomic site. The images were reviewed and a consensus was reached by two board-certified radiologists who were unaware of any clinical or radiological information using hard-copy films and multimodality computer platform. The two readers for ¹¹C-choline PET/CT and those for conventional imaging were not the same persons.

Each tumor was staged according to the Tumor Node Metastasis (TNM) classification of the International Union Against Cancer for sarcoma of bone and the American Joint Committee staging protocol for sarcoma of the soft tissue.^(18,19) T, N and M stages were assigned for both PET/CT and conventional imaging. T staging was confirmed by pathological evaluation using specimens obtained from surgical resection of the primary tumors. N staging was confirmed by pathological examinations in two patients using specimens obtained from sampling of regional nodes. In terms of extraregional nodes in two patients, nodal staging was confirmed by an obvious progression in number and/or size of the lesions on follow-up examinations. The mean follow-up period was 172 days (range, 44–322 days).

Statistical analysis. All variables were assessed on a patient-by-patient basis. The McNemar test was used for paired comparisons between ¹¹C-choline PET/CT and conventional imaging. Statistical analysis was performed with the SPSS version 11 software program (SPSS, Chicago, IL, USA).

Results

There were eight bone sarcomas and eight soft tissue sarcomas (Table 1). The primary sites included shoulder (*n* = 2), chest wall

Table 2. Staging of bone and soft tissue sarcoma

| Variables | ¹¹ C-choline PET/CT | Conventional imaging | P-value |
|---------------|--------------------------------|----------------------|---------|
| Overall stage | | | 0.023 |
| Correct | 15 (94) | 8 (50) | |
| Understaged | 1 (6) | 8 (50) | |
| Overstaged | 0 | 0 | |
| N stage | | | 0.041 |
| Correct | 16 (100) | 10 (63) | |
| Understaged | 0 | 6 (38) | |
| Overstaged | 0 | 0 | |
| M stage | | | 0.617 |
| Correct | 15 (94) | 13 (81) | |
| Understaged | 1 (6) | 3 (19) | |
| Overstaged | 0 | 0 | |

Note: Data are presented as number (*n*). Numbers in parentheses are percentages. CT, computed tomography; PET, positron emission tomography.

(*n* = 2), retroperitoneum (*n* = 2), iliac bone (*n* = 2), leg (*n* = 2), thigh (*n* = 1), perineum (*n* = 1), tibia (*n* = 1), femur (*n* = 1), mandible (*n* = 1) and spine (*n* = 1). Pathological diagnoses were osteosarcoma (*n* = 4), Ewing sarcoma (*n* = 3), leiomyosarcoma (*n* = 3), clear cell sarcoma (*n* = 1), chondrosarcoma (*n* = 1), pleomorphic malignant fibrous histiocytoma (*n* = 1), myxoid liposarcoma (*n* = 1), rhabdomyosarcoma (*n* = 1), and alveolar soft part sarcoma (*n* = 1). Histological grade of tumors was grade 1 (*n* = 1), grade 2 (*n* = 1), grade 3 (*n* = 11) and grade 4 (*n* = 3).

All patients of initial staging had increased ¹¹C-choline uptake of the primary lesion (average maximal SUV ± SD: 5.92 ± 3.68 [range, 2.15–15.05]). Pathological T stages available in patients with initial staging are as follows: T1 (*n* = 1), T1a (*n* = 1), T1b (*n* = 1), T2 (*n* = 4) and T2b (*n* = 5). T stages in patients with restaging were T1 (*n* = 2), T1b (*n* = 1) and T2 (*n* = 1). Tumor size of patients for initial staging was 78.5 ± 34.0 mm (mean ± SD [range, 16.0–133.0 mm]). Both ¹¹C-choline PET/CT and conventional imaging classified the T stage correctly in all patients. Twelve (75%) of the 16 patients had N0 disease. Using ¹¹C-choline PET/CT, the N stage was correctly assigned in all patients, whereas the accuracy of conventional imaging in N stage was 63% (*P* = 0.041, Table 2). Understaging occurred in six patients (38%). Three of these patients (19%) had metastasis of inguinal lymph node whose largest diameter was less than 10.0 mm (Fig. 1). The incidence of distant metastases was high in our study population. Both ¹¹C-choline PET/CT and conventional imaging detected bone metastases in seven patients (44%), lung metastases in five (31%) and pleural dissemination in two (18%, Fig. 2). Using ¹¹C-choline PET/CT, the M stage was correctly assigned in 15 patients (94%), whereas the accuracy of conventional imaging in M stage was 81% (*P* = 0.617, Table 2).

The complete stages of all patients were stage IA (*n* = 1), stage IIB (*n* = 1), stage III (*n* = 1) and stage IV (*n* = 13). TNM stage was correctly assessed with ¹¹C-choline PET/CT in 15 of 16 patients (94%) and with conventional imaging in eight of 16 patients (50%, *P* = 0.023, Table 2). ¹¹C-choline PET/CT assigned an incorrect TNM stage in a patient. This patient was understaged due to small metastatic lung tumor which was not clearly visualized by CT part of ¹¹C-choline PET/CT. Eight patients were understaged by conventional imaging (50%). Of these, skip metastases of soft tissues were identified in two (25%) and small nodal metastases in six (75%). ¹¹C-choline PET/CT correctly determined TNM stage in seven patients (44%) in whom stage derived from conventional imaging was incorrect.

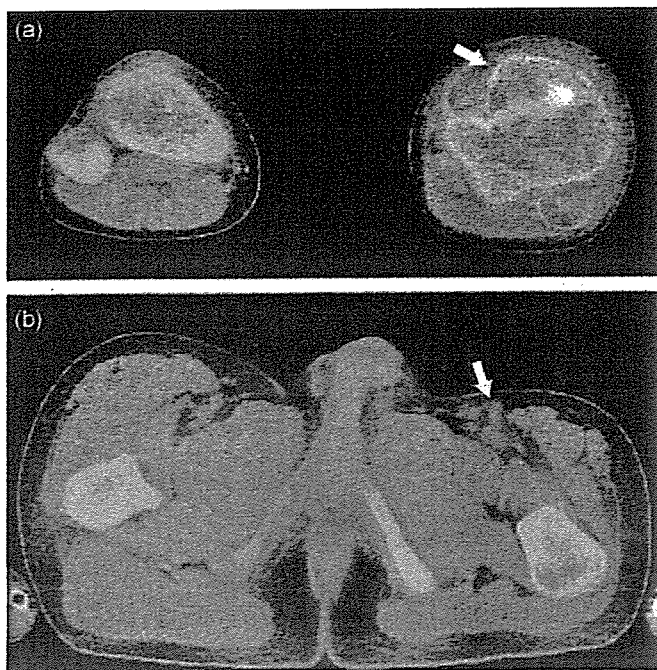


Fig. 1. A 13-year-old boy with osteosarcoma. (a) Transverse ¹¹C-choline positron emission tomography (PET)/computed tomography (CT) image revealed hypermetabolic focus in the proximal portion of the left tibia (arrow). PET/CT findings were verified at histopathological analysis. (b) Abnormal uptake of ¹¹C-choline was also noted in the left inguinal lymph node, which was interpreted as highly suspicious for malignancy (arrow). Subsequent resection revealed metastasis from osteosarcoma.

Discussion

The results of the present study show that ¹¹C-choline PET/CT improves the accuracy of staging in patients with bone and soft tissue sarcomas compared to conventional imaging. Specifically, ¹¹C-choline PET/CT has potentially significant implications for detecting nodal and distant metastases at overall staging. Reports about the efficacy of ¹¹C-choline in the localization and detection of bone and soft tissue sarcomas are still limited.⁽¹⁵⁾ To our knowledge, no study regarding ¹¹C-choline PET/CT for staging bone and soft tissue sarcomas was found. In our study, seven of the 16 patients had skip metastases of soft tissue or nodal metastases detected by ¹¹C-choline PET/CT that were not identified by routine clinical and conventional radiological evaluation.

The ability of PET to depict increased metabolism in malignancies has greatly improved the accuracy in detecting neoplasms.⁽⁴⁾ However, compared with conventional imaging studies, use of PET alone results in a lack of substantial detail.²⁰ The PET/CT device permits sequential acquisition of anatomic CT and functional PET images in a single scanning session. Morphological characterization of scintigraphic lesions by PET/CT resulted in a lower percentage of equivocal interpretations compared with that of conventional imaging. Tumor-detecting PET/CT technology is growing rapidly. However, there are only limited data available on staging of bone and soft tissue sarcomas with PET/CT.

Carbon-11 choline uptake was significantly higher in malignant soft tissue tumors and was due to the high utilization of cell membranes of these lesions. ¹¹C-choline uptake is observed physiologically in the liver, pancreas, kidney and duodenum. ¹¹C-choline is also secreted into phospholipid-rich pancreatic juice in a non-fasting state. A potential advantage of ¹¹C-choline PET/CT might be the assessment of tumors in the skull or retroperitoneum. Blood clearance of ¹¹C-choline is rapid and radioactive distribution

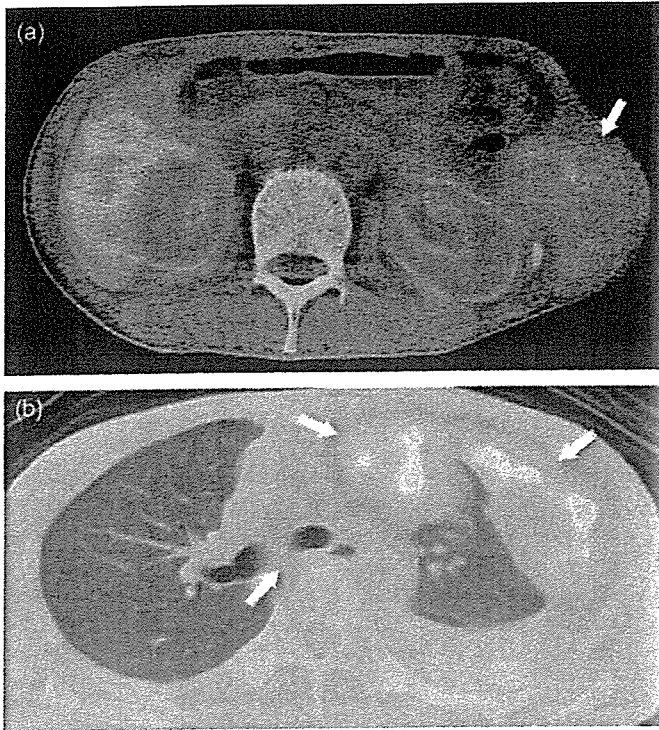


Fig. 2. A 34-year-old man with clear cell sarcoma. (a) Transverse ^{11}C -choline positron emission tomography (PET)/computed tomography (CT) image depicting abnormal uptake in the tumor arising from the left lateral chest wall (arrow). (b) PET/CT image also depicts pleural dissemination and mediastinal lymph node (arrows). Follow-up findings in this patient confirmed the diagnosis.

in tissues is constant in 5 min. The accumulation of ^{11}C -choline in the skull or retroperitoneum is hardly affected by background within the limits of short uptake time. In comparison to ^{18}F FDG, physiological background level in the urinary tract is low. This may be due to incomplete tubular reabsorption of the intact tracer, or enhanced excretion of labeled oxidative metabolites like betaine.⁽¹²⁾

References

- Reuther G, Mutschler W. Detection of local recurrent disease in musculoskeletal tumors: magnetic resonance imaging versus computed tomography. *Skeletal Radiol* 1990; **19**: 85–90.
- Nieweg OE, Pruim J, van Ginkel RJ *et al*. Fluorine-18-fluorodeoxyglucose PET imaging of soft-tissue sarcoma. *J Nucl Med* 1996; **37**: 257–61.
- Eary JF, Conrad EU, Bruckner JD, Folpe A, Hunt KJ, Mankoff DA, Howlett AT. Quantitative [^{18}F]fluorodeoxyglucose positron emission tomography in pretreatment and grading of sarcoma. *Clin Cancer Res* 1998; **4**: 1215–20.
- Franzius C, Sciuk J, Daldrup-Link HE *et al*. FDG-PET for detection of osseous metastases from malignant primary bone tumors: comparison with bone scintigraphy. *Eur J Nucl Med* 2000; **27**: 1305–11.
- Schwarzbach MHM, Dimitrakopoulou-Strauss A, Willeke F *et al*. Clinical value of [^{18}F]fluorodeoxyglucose positron emission tomography imaging in soft tissue sarcomas. *Ann Surg* 2000; **231**: 380–6.
- Ioannidis JP, Lau J. ^{18}F -FDG PET for the diagnosis of soft-tissue sarcoma: a meta-analysis. *J Nucl Med* 2003; **44**: 717–24.
- Tateishi U, Yamaguchi U, Seki K *et al*. Glut-1 expression and enhanced glucose metabolism are associated with tumor grade in bone and soft tissue sarcomas: a prospective evaluation by [^{18}F]fluorodeoxyglucose positron emission tomography. *Eur J Nucl Med Mol Imaging* 2006; **33**: 683–91.
- Hara T, Kosaka N, Shinoura N *et al*. PET imaging of brain tumor with [^3H]methyl- ^{11}C]-choline. *J Nucl Med* 1997; **38**: 842–7.
- Hara T, Kosaka N, Kishi H. PET imaging of prostate cancer using carbon-11-choline. *J Nucl Med* 1998; **39**: 990–5.
- Hara T, Inagaki K, Kosaka N *et al*. Sensitive detection of mediastinal lymph node metastasis of lung cancer with ^{11}C -choline PET. *J Nucl Med* 2000; **41**: 1507–13.
- Torizuka T, Kanno T, Futatsubashi M *et al*. Imaging of gynecologic tumors: comparison of ^{11}C -choline PET with ^{18}F -FDG PET. *J Nucl Med* 2003; **44**: 1051–6.
- Ishidate K. Choline/ethanolamine kinase from mammalian tissues. *Biochim Biophys Acta* 1997; **1348**: 70–8.
- Maeda T, Tateishi U, Komiyama M *et al*. Distant metastasis of prostate cancer: Early detection of recurrent tumor with dual-phase carbon-11 choline positron emission tomography/computed tomography in two cases. *Jpn J Clin Oncol* in press.
- Zhang H, Tian M, Oriuchi N *et al*. ^{11}C -choline PET for the detection of bone and soft tissue tumors in comparison with FDG PET. *Nucl Med Commun* 2003; **24**: 273–9.
- Tian M, Zhang H, Oriuchi N *et al*. Comparison of ^{11}C -choline PET and FDG PET for the differential diagnosis of malignant tumors. *Eur J Nucl Med Mol Imaging* 2004; **31**: 1064–72.
- Bar-Shalom R, Yefremov N, Guralnik L *et al*. Clinical performance of PET/CT in evaluation of cancer: additional value for diagnostic imaging and patient management. *J Nucl Med* 2003; **44**: 1200–9.
- Hara T, Yuasa M. Automated synthesis of [^{11}C]choline, a positron-emitting tracer for tumor imaging. *Appl Radiat Isot* 1999; **50**: 531–3.
- Green FL, Page DL, Fleming ID *et al*. *AJCC Cancer Staging Manual*, 6th edn. New York: Springer, 2002.
- Sobin LH, Wittekind C. *UICC TNM Classification of Malignant Tumours*, 6th edn. New York: Wiley, 2002.
- Franzius C, Daldrup-Link HE, Wagner-Bohn A *et al*. FDG-PET for detection of recurrences from malignant primary bone tumors: comparison with conventional imaging. *Ann Oncol* 2002; **13**: 157–60.
- Uchida T, Yamashita S. Molecular cloning, characterization, and expression in *Escherichia coli* of a cDNA encoding mammalian choline kinase. *J Biol Chem* 1992; **267**: 10 156–62.

Limited resolution of the present generation of ^{11}C -choline PET/CT and the partial volume effect result in failure to detect small lesions. In our study, one patient was understaged due to small metastatic lung tumor, which was not visualized clearly by the CT part of ^{11}C -choline PET/CT. Faint increase in tracer uptake and motion artifact caused by breathing contribute to false negative results. However, the advantage of ^{11}C -choline PET/CT is that the whole-body can be visualized in a single examination. In our study, 50% of patients were understaged by conventional imaging. The inaccuracy of conventional imaging in assessing skip metastases of soft tissues is due to the field of view.

We reported the accurate modality of ^{11}C -choline PET/CT as a non-invasive method for staging in patients with bone and soft tissue sarcomas compared to conventional imaging. Choline is an essential component of the cell membrane, and choline uptake may be via a choline-specific transporter protein. Choline kinase, which catalyzes the phosphorylation of choline, is upregulated in tumor cells.⁽¹²⁾ In some types of tumor cells, overexpression of choline-specific transporter protein and choline kinase were identified by *in situ* hybridization.⁽²¹⁾ ^{11}C -choline will be phosphorylated by choline kinase as a choline analog and retained in tumor cells.⁽²¹⁾ However, the precise pathway of metabolic trapping by tumor cells has not been elucidated, and further studies to clarify the mechanism of imaging by ^{11}C -choline are needed.

Our study has limitations. Most patients in this study had high-grade tumors (88%) and may differ from the patient population of previous studies. Our study was intended to examine the staging prior to treatment; therefore, patient population of high-grade tumors may explain the significant accuracy in overall staging compared to conventional imaging. A study with a larger patient population would clarify the influence of ^{11}C -choline PET/CT on staging. ^{11}C -choline is clearly a sensitive PET tracer for staging patients with bone and soft tissue sarcomas. The short half-life of ^{11}C -choline necessitates the availability of an on-site cyclotron, which causes practical restriction. More specific radiotracers will help overcome this limitation in the future.

In summary, the use of ^{11}C -choline PET/CT in patients with bone and soft tissue sarcomas increases the accuracy of overall staging and N staging compared to conventional staging. Our study suggests that whole-body ^{11}C -choline PET/CT should be the preferred modality for staging in patients with bone and soft tissue sarcomas.



Radiologic Removal and Replacement of Port-Catheter Systems for Hepatic Arterial Infusion Chemotherapy

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OBJECTIVE. The purpose of our study was to retrospectively evaluate the safety and efficacy of radiologic removal and replacement of port-catheter systems.

MATERIALS AND METHODS. Between January 1999 and December 2004, 532 patients with unresectable advanced liver cancer underwent radiologic placement of port-catheter systems at our institution. Of these, 18 patients (nine men and nine women; age range, 32–83 years; mean age, 53.8 years) underwent removal of an implanted port-catheter system via the right femoral artery and radiographically guided replacement with a new system to allow continuous hepatic arterial infusion chemotherapy; we retrospectively reviewed these 18 cases. The reasons for removal of the previously implanted systems were as follows: catheter dislodgement ($n = 15$), catheter obstruction ($n = 1$), infection related to the implanted port ($n = 1$), and hemodynamic change ($n = 1$). Digital subtraction angiography and CT were performed, usually during injection of contrast medium through the implanted port-catheter system, within a few days after the replacement procedure and every 3 months thereafter.

RESULTS. We successfully performed radiologic removal and replacement of the port-catheter system while the patient was under local anesthesia in all 18 patients without complications requiring treatment. The cumulative patency rates of the hepatic artery after removal of the old port-catheter system and replacement with a new port-catheter system were 87.8% and 64.1% at 6 months and 1 year, respectively. Hepatic arterial infusion chemotherapy after replacement was performed 0–68 times (median, 19 times).

CONCLUSION. When an implanted port-catheter system can no longer be used but the patency of the hepatic artery is confirmed and continuous hepatic arterial infusion chemotherapy is required, removal and replacement of the port-catheter system are recommended.

Repeated hepatic arterial infusion chemotherapy using an implanted port-catheter system is reported to be effective for the treatment of patients with unresectable advanced liver malignancies [1–4]. Recent advancements in interventional radiologic techniques have led to nonsurgical placement of port-catheter systems being performed increasingly frequently, and the use of a side-hole catheter with tip fixation is recommended during this procedure to prevent catheter dislodgement and hepatic arterial occlusion [5–8]. Although continuous use of an implanted port-catheter system is the ideal scenario, this generally is not possible even with careful management because of complications such as occlusion, kinking, or dislodgement of the implanted catheter; hepatic artery occlusion; or infection of the implanted port-catheter system [6–12]. In such cases, if the hepatic artery is

patent, continued hepatic arterial infusion chemotherapy is possible after the original device has been replaced with a new system. The purpose of this study was to retrospectively evaluate the safety and efficacy of the radiologic removal and replacement of a port-catheter system.

Materials and Methods

This study is a retrospective one, and approval from the institutional review board of our hospital was obtained.

Patients

Between January 1999 and December 2004, 532 patients with unresectable advanced liver cancer underwent radiologically guided placement of port-catheter systems at our institution. Of those patients, 18 (nine men and nine women; age range, 32–83 years; mean age, 53.8 years) received a replacement system after the original device had been

Keywords: chemotherapy, implantable devices, liver cancer, port-catheter system

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removed to allow continuous hepatic arterial infusion chemotherapy. Seventeen patients had liver metastases that originated from colorectal cancer ($n = 9$), breast cancer ($n = 4$), gastric cancer ($n = 3$), or carcinoma of the papilla of Vater ($n = 1$), and the remaining patient had hepatocellular carcinoma.

The reasons for the removal of the previously implanted systems were as follows: catheter dislodgement ($n = 15$); catheter obstruction ($n = 1$); infection related to the implanted port ($n = 1$); and hemodynamic change ($n = 1$), such as hepatopetal flow of the common hepatic artery that was changed to hepatofugal flow as a result of altered flow in the gastroduodenal artery secondary to stenosis of the celiac artery. All 18 patients had only liver lesions that were well controlled by hepatic arterial infusion chemotherapy, so continuous hepatic arterial infusion chemotherapy was desired if the hepatic artery was patent. That information was obtained from the medical records.

First Placement of Port-Catheter Systems

The placement site of the port-catheter system was originally chosen according to the following method: All patients underwent angiography before catheter placement, which was performed using a 5-French angiographic catheter inserted from the right femoral artery to allow arterial mapping and to prevent extrahepatic influx of anticancer agents. The extrahepatic arteries branching from the hepatic artery, such as the right gastric artery, posterior superior pancreaticoduodenal artery, and superior duodenal artery, were embolized with microcoils (Tornado, Cook; or Trufill, Cordis) through a 2.9-French microcatheter (Jamiro, Kaneka; or Sniper, Clinical Supply) inserted coaxially [5, 13]. The left gastric artery and gastroduodenal artery were also embolized when the angiographic catheter tip was inserted into the splenic artery [5].

In patients with more than two hepatic arteries, these arteries were converted into a single arterial supply by microcoil embolization so that drugs could be distributed to the entire liver using a single indwelling catheter [5]. A 5-French angiographic catheter was then inserted from the left subclavian artery ($n = 14$) or the right femoral artery ($n = 4$) and was advanced to the common hepatic artery via the celiac artery.

Subsequently, using the catheter-exchange method, a 5-French indwelling catheter (Anthon P-U catheter, Toray; or W spiral catheter, Piolax) with ($n = 16$) or without ($n = 2$) a side hole was inserted. The tips of these catheters were tapered to 2.7-French and 20 cm in length; the catheters were inserted into the gastroduodenal artery ($n = 9$), the splenic artery ($n = 1$), the peripheral branch of the hepatic artery ($n = 2$), the right hepatic artery ($n = 3$), the common hepatic artery ($n = 2$), or the accessory

left gastric artery arising from the left hepatic artery ($n = 1$). In 12 of the 18 patients who had catheters inserted into the gastroduodenal artery ($n = 9$), splenic artery ($n = 1$), and others ($n = 2$), the artery around the tip of indwelling catheter was embolized using microcoils and a mixture (1:1.5) of *n*-butyl cyanoacrylate (Histoacryl, Braun) and iodized oil (Lipiodol Ultrafluide, Laboratoire Guerbet) through a microcatheter inserted coaxially via a 5-French angiographic catheter inserted from the femoral artery. The catheter tip was also fixed in these 12 patients.

In four of the remaining six patients in whom the catheter tip was not fixed, a W spiral catheter was used; the spiral-shaped tip of this catheter has the function of securing it. The side hole of the catheter was placed into the common hepatic artery or the celiac artery. Finally, the proximal end of the indwelling catheter was connected to a port implanted in a subcutaneous pocket created in the left chest wall or the right upper thigh.

Removal and Replacement of Port-Catheter Systems

Written informed consent was obtained from all the patients before these procedures. All the procedures were performed in an angiographic suite by interventional radiologists with the patient under local anesthesia. On the same day as the procedure or the day before the procedure, all patients underwent angiography using a 5-French angiographic catheter inserted from the right femoral artery to confirm patency of the hepatic arteries.

In the four patients in whom the catheter had previously been implanted from the right femoral artery, after opening the subcutaneous space housing the port, the indwelling catheter was directly withdrawn from the right femoral artery with the port.

In the 14 patients in whom the port-catheter system was previously implanted via the left subclavian artery, a 5-French hook-shaped catheter was first inserted from a right femoral artery through a 6-French sheath introducer and was then wrapped around the indwelling catheter. The hook-shaped catheter was then pulled to relocate the indwelling catheter tip to the aorta. After the hook-shaped catheter was withdrawn, a 5-French basket retriever was inserted via the right femoral artery through the sheath introducer to capture the distal tip of the indwelling catheter. After a small incision was made at the insertion site in the left chest wall, the implanted port was withdrawn, the proximal part of the indwelling catheter was cut, and the port was removed from the catheter. The indwelling catheter captured by the basket retriever was then withdrawn from the right femoral artery. Subsequently, replacement with a new port-catheter system was performed using the same methods described earlier.

The total time required for the procedure ranged from 107 to 225 minutes (mean, 155 minutes). Catheters were inserted from the left subclavian artery ($n = 15$), the right femoral artery ($n = 1$), and the left inferior epigastric artery ($n = 2$). In three of four patients in whom the first placement procedure was from the right femoral artery and had been performed at another institution, replacement was from the left subclavian artery. The catheters were advanced via the celiac artery ($n = 16$) or through the pancreaticoduodenal arcade via the superior mesenteric artery in cases of celiac artery stenosis ($n = 2$). Catheter tips were inserted into the gastroduodenal artery ($n = 2$), the splenic artery ($n = 3$), the peripheral branch of the hepatic artery ($n = 6$), the right hepatic artery ($n = 5$), the common hepatic artery ($n = 1$), and the middle hepatic artery ($n = 1$) (Table 1).

In one patient, because selecting a placement site for the catheter was difficult using the method mentioned earlier, placement was performed as follows: We first selected the celiac artery with a 5-French angiographic catheter (inserted via the femoral artery) and then inserted an indwelling catheter (Anthon P-U catheter, Toray) using the catheter-exchange method. A 2.9-French microcatheter (Sniper, Clinical Supply) was inserted coaxially into the right hepatic artery through the indwelling catheter, which was thereby relocated to the aorta. Finally, the proximal end of the microcatheter was connected directly to the implanted port using a connecting device. In six of 18 patients, the tip of the indwelling catheter was fixed using microcoils and a mixture of *n*-butyl cyanoacrylate and iodized oil. In eight of 12 patients in whom the catheter tip was not fixed, a W spiral catheter was used.

Using this system, hepatic arterial infusion chemotherapy was started a few days after the procedure, depending on the clinical circumstances. The details of hepatic arterial infusion chemotherapy and management of this system have been reported previously [7]. Digital subtraction angiography and CT were performed during injection of contrast medium through the implanted port-catheter system within a few days after the procedure and every 3 months thereafter to confirm that the catheter and hepatic artery were patent and that the entire liver was perfused adequately. These investigations were also performed whenever patients reported any symptoms that might be related to hepatic arterial infusion chemotherapy.

Evaluation

Outcome was evaluated in terms of the success rate for removal and replacement of the port-catheter systems, complications of the procedure, and number of sessions of hepatic arterial infusion chemotherapy after replacement with the new systems. The cumulative patency rate of the hepatic artery confirmed by digital subtraction angiography was calculated according to the Kaplan-Meier method.

Port-Catheter System for Hepatic Arterial Infusion

TABLE 1: Approach Artery and Location of Catheter Tip of Port-Catheter Systems

| Removal of Original Port-Catheter System | | | Placement of New Port-Catheter System | | | | | |
|--|--------------------------|-----------------|---------------------------------------|-------------------------------------|-----------------|------------------------|-------------------------------------|---|
| Approach Artery | Location of Catheter Tip | No. of Patients | Approach Artery | Location of Catheter Tip | No. of Patients | | | |
| Left subclavian artery | Gastroduodenal artery | 9 | Left subclavian artery | Gastroduodenal artery | 1 | | | |
| | | | | Splenic artery | 2 | | | |
| | | | | Right hepatic artery | 3 | | | |
| | | | Left inferior epigastric artery | Peripheral branch of hepatic artery | 1 | | | |
| | | | | Splenic artery | 1 | | | |
| | | | | Peripheral branch of hepatic artery | 1 | | | |
| | | | Right femoral artery | Common hepatic artery | 2 | Left subclavian artery | Right hepatic artery | 1 |
| | | | | | | Left subclavian artery | Right hepatic artery | 1 |
| | | | | | | Left subclavian artery | Peripheral branch of hepatic artery | 2 |
| Right femoral artery | Right hepatic artery | 2 | Left subclavian artery | Common hepatic artery | 1 | | | |
| | | | Left subclavian artery | Peripheral branch of hepatic artery | 1 | | | |
| | | | Left subclavian artery | Gastroduodenal artery | 1 | | | |
| Right femoral artery | Right hepatic artery | 2 | Left subclavian artery | Peripheral branch of hepatic artery | 1 | | | |
| | | | Right femoral artery | Middle hepatic artery | 1 | | | |

Results

Patients were followed up for a period ranging from 7 to 1,806 days (median, 373 days) after catheter replacement. We successfully performed removal and replacement of port-catheter systems in all 18 patients (Fig. 1). Although our patients experienced some minor complications requiring no treatment, such as hemorrhage, nausea, and pain, there were no other major complications, such as ischemia or infarction caused by extrahepatic arterial embolization and massive hematoma. Moreover, we have been able to perform hepatic arterial infusion chemotherapy continuously in our department in 16 of these patients.

The progress of one of the remaining two patients is not known because this patient only underwent removal and replacement of the port-catheter system at our institution and was not followed up by us. The other was a 33-year-old woman with multiple liver metastases from breast cancer. The tip of this patient's catheter was originally replaced into a peripheral branch of the hepatic artery; however, because this catheter became dislodged, it was removed 7 days after the original replacement procedure and was replaced with a new port-catheter system on the same day. In this procedure, the catheter tip was also inserted into another peripheral branch of the hepatic artery. In this patient, hepatic arterial infusion chemotherapy could be performed 58 times over 524 days until hepatic arterial occlusion occurred after the second replacement procedure (Fig. 2).

Overall, after replacement of the port-catheter systems, hepatic arterial infusion chemotherapy was performed 0–68 times (median, 19 times). Chemotherapy was continued ($n = 4$) or was terminated because of death ($n = 5$), hepatic arterial occlusion ($n = 4$), catheter dislodgement ($n = 2$), or change to other treatments ($n = 2$). The cumulative patency rates of the hepatic artery at 6 months and 1 year after replacement with new systems were 87.8% and 64.1%, respectively.

Discussion

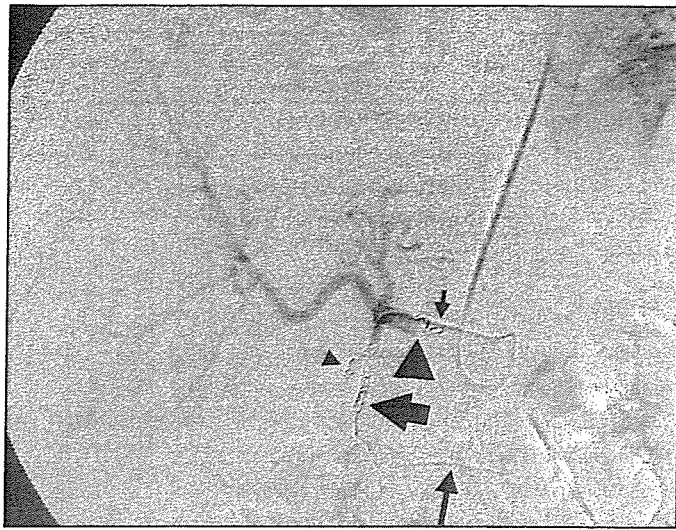
Repeated hepatic arterial infusion chemotherapy using an implanted port-catheter system is reported to be an effective therapy for patients with unresectable advanced liver malignancies and is used widely as a local approach [1–4]. When placing the catheter radiologically, which is less invasive than placing it using a surgical procedure, use of a side-hole catheter is recommended [5–8] and fixation of the catheter tip is recommended to prevent catheter dislodgement and hepatic arterial occlusion caused by mechanical stimulation resulting from movement of the unfixed catheter tip [5].

Hepatic arterial occlusion and catheter dislodgement are the most common complications that require hepatic arterial infusion chemotherapy to be stopped, with prevalences of 0–17% and 2.2–14.3%, respectively, recently reported during use of non-surgically inserted port-catheter systems [6–12]. However, placement of a side-hole

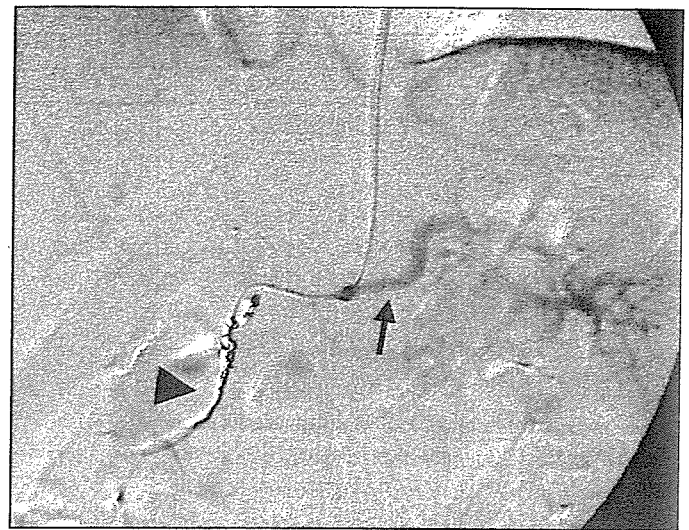
catheter with tip fixation is reported to be associated less frequently with hepatic arterial occlusion (5.4% [6]) or catheter dislodgement (2.2–2.8% [6, 7]). Accordingly, for the initial placement procedure, we usually insert the tip of a side-hole catheter into the deep portion of the gastroduodenal artery and fix it using microcoils and a mixture of *n*-butyl cyanoacrylate and iodized oil.

Although various complications such as catheter dislodgement can preclude the continued use of an implanted port-catheter system, we aim to continue to treat patients with hepatic arterial infusion chemotherapy if the hepatic artery is patent. A dislodged catheter causes flow into the extrahepatic arteries, resulting in reduced concentrations of drug in the liver. We overcome this complication by embolizing extrahepatic arteries, such as the left gastric artery, splenic artery, or dorsal pancreatic artery [5], if possible. However, in the present study, we removed implanted port-catheter systems and replaced them with new systems because catheter dislodgement was too great to be overcome by embolizing extrahepatic arteries.

We decided to remove the original implanted port-catheter system when replacing it with a new system to minimize the disturbance associated with replacement, the unnecessary stimulation of the artery, and the possibility of infection, and because this was generally the patient's request. Although we anticipated that removal of indwelling catheters with fixed tips would be difficult, it was possible to safely remove the catheter in all nine patients.



A



B



C

Fig. 1—43-year-old man with multiple liver metastases from rectal cancer. **A**, Arteriogram via port obtained after placement shows that all hepatic arteries are well visualized. Catheter tip was inserted into deep portion of gastroduodenal artery (*long thin arrow*), and side hole was placed in common hepatic artery (*large arrowhead*). To prevent extrahepatic influx of anticancer agents, gastroduodenal artery (*thick arrow*), right gastric artery (*small arrow*), and posterior superior pancreaticoduodenal artery (*small arrowhead*) were embolized with microcoils. Embolization of gastroduodenal artery was performed using mixture of *n*-butyl cyanoacrylate and iodized oil in addition to microcoils to fix catheter and occlude arteries. **B**, Arteriogram via port obtained 4 months after placement shows that splenic artery (*arrow*) is better visualized than hepatic arteries because of catheter dislodgement (*arrowhead*). **C**, Arteriogram via port obtained after replacement shows that all hepatic arteries are well visualized again. Catheter tip was inserted into peripheral branch of hepatic artery (*arrow*), and side hole was placed in common hepatic artery (*arrowhead*).

We removed implanted catheters via the right femoral artery in all patients. Particularly when the catheter is originally implanted from the left subclavian artery, removal should be performed via the femoral artery to prevent brain infarction due to release of thrombus around the indwelling catheter and subsequent vertebral arterial embolization [14]. We successfully removed implanted port-catheter systems in all patients without complications (such as brain infarction, hemorrhage, hematoma, infection, or pseudoaneurysm) requiring treatment and with the patient under local anesthesia.

We performed removal and replacement on the same day. After deciding the position of catheter tip insertion based on angiography performed before replacement, we inserted

new catheters from the left subclavian artery in many patients because this was the approach artery that had been used previously and patients had therefore previously experienced the procedure. Although the risk of complications such as hemorrhage, hematoma, and pseudoaneurysm is higher if the same route is used, we successfully performed removal and replacement of port-catheter systems using this approach without observing complications requiring treatment.

When an old system was replaced with a new system, the catheter tip was inserted into another artery because we had already embolized the gastroduodenal artery. Replacement of a side-hole catheter with its tip fixed and inserted into another artery was possible in only six of the 18 patients. In one of the remaining

12 patients in whom the tip was inserted into the peripheral branch of the hepatic artery, a second replacement procedure was required because the catheter became dislodged 7 days after the first replacement procedure.

In the present study, at 1 year after replacement, a 64.1% cumulative patency rate for the hepatic artery was achieved. This patency rate is lower than previously reported cumulative patency rates for first placement (81.4% [7] and 86.3% [8]). We think that this discrepancy results from nonfixation of the catheter tip, injury of the hepatic artery caused by prior hepatic arterial infusion chemotherapy, or both. Nonetheless, because we could perform hepatic arterial infusion chemotherapy a median of 19 times after port-catheter system replacement, it seems to be worth continuing

Port-Catheter System for Hepatic Arterial Infusion

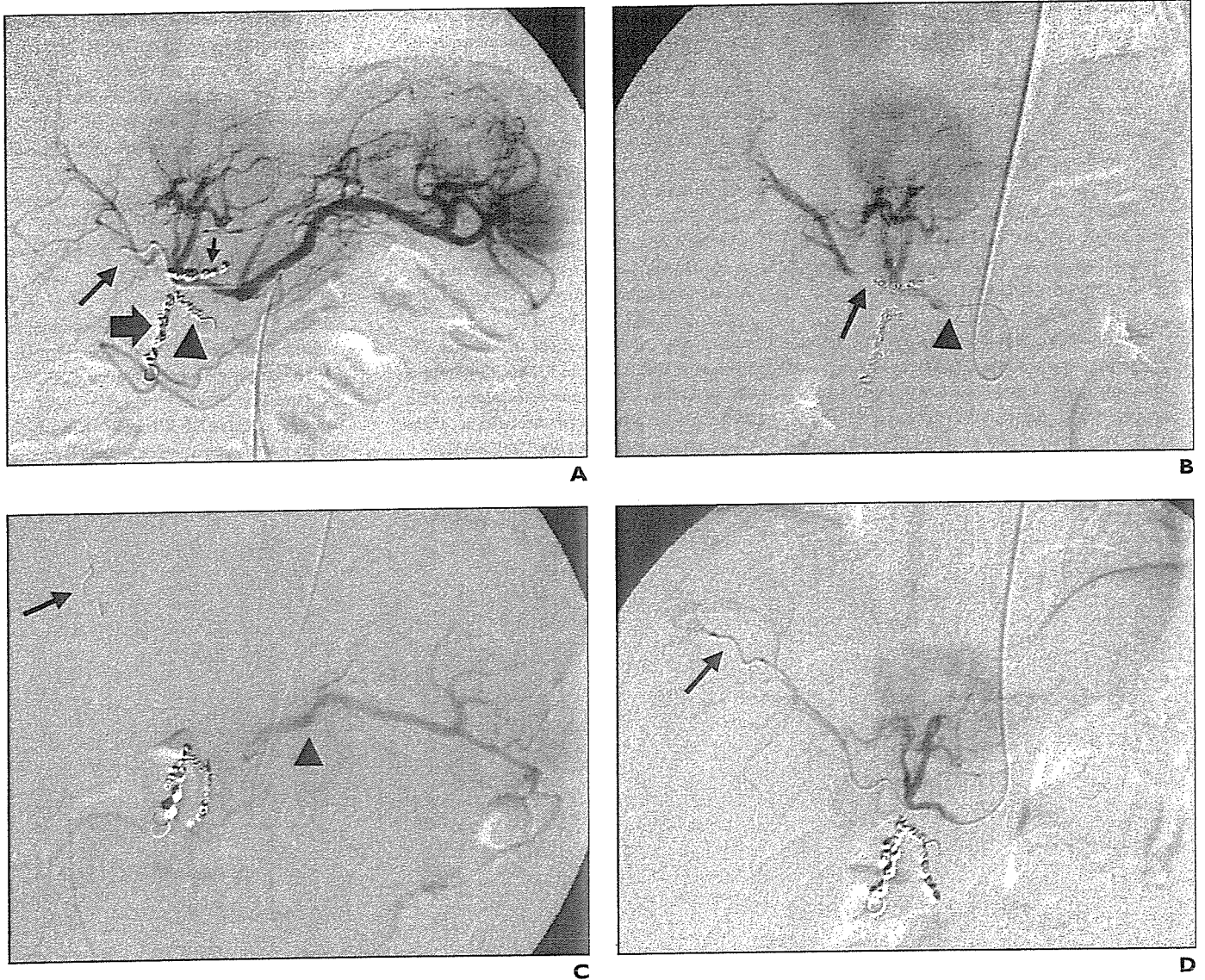


Fig. 2—33-year-old woman with multiple liver metastases from breast cancer.

A, Celiac arteriogram obtained after occlusion of implanted catheter (*long thin arrow*) shows that hepatic arteries are well visualized. Gastroduodenal artery (*thick arrow*), posterior superior pancreaticoduodenal artery (*arrowhead*), and right gastric artery (*small thin arrow*) were embolized with microcoils to prevent extrahepatic influx of anticancer agents.

B, Arteriogram via port obtained after replacement shows that all hepatic arteries can be visualized. Catheter tip was inserted into peripheral branch of hepatic artery and side hole was placed in common hepatic artery (*arrowhead*). Origin of right hepatic artery (*arrow*) was not visualized because of stenosis caused by tip of original catheter, but right hepatic artery is well visualized because of blood supply via left hepatic artery through intrahepatic arterial anastomoses.

C, Arteriogram via port obtained 1 week after replacement shows that splenic artery (*arrowhead*) is better visualized than hepatic arteries because of catheter dislodgement (*arrow*).

D, Arteriogram via port obtained after second removal and replacement shows that hepatic arteries are well visualized. Catheter tip (*arrow*) was inserted into another peripheral branch of hepatic artery.

hepatic arterial infusion chemotherapy when this therapy is needed in situations such as absence of extrahepatic lesions or when liver metastases are thought to be the prognosis-limiting factor.

In conclusion, although the retrospective design of this study meant that many limitations exist, it is noteworthy that we could

safely remove and replace port-catheter systems so that hepatic arterial infusion chemotherapy could continue. Attempting these procedures appears worthwhile if continuing treatment using an implanted port-catheter system is not possible, the hepatic artery is confirmed patent, and continuous hepatic arterial infusion chemotherapy is required.

References

1. Allen-Merish TG, Earlam S, Fordy C, Abrams K, Houghton J. Quality of life and survival with continuous hepatic-artery floxuridine infusion for colorectal liver metastases. *Lancet* 1994; 344:1255–1260
2. Link KH, Sunelaitis E, Kommann M, et al. Regional chemotherapy of nonresectable colorectal liver me-

- tastases with mitoxantrone, 5-fluorouracil, folinic acid, and mitomycin C may prolong survival. *Cancer* 2001; 92:2746–2753
3. Arai Y, Inaba Y, Takeuchi Y, Ariyoshi Y. Intermittent hepatic arterial infusion of high-dose 5FU on a weekly schedule for liver metastases from colorectal cancer. *Cancer Chemother Pharmacol* 1997; 40:526–530
 4. Kumada T, Arai Y, Itoh K, et al. Phase II study of combined administration of 5-fluorouracil, epirubicin and mitomycin-C by hepatic artery infusion in patients with liver metastases of gastric cancer. *Oncology* 1999; 57:216–223
 5. Arai Y, Inaba Y, Takeuchi Y. Interventional techniques for hepatic arterial infusion chemotherapy. In: Castaneda-Zuniga WR, ed. *Interventional radiology*, 3rd ed. Baltimore, MD: Williams & Wilkins, 1997:192–205
 6. Yamagami T, Iida S, Kato T, et al. Using *n*-butyl cyanoacrylate and the fixed-catheter-tip technique in percutaneous implantation of a port-catheter system in patients undergoing repeated hepatic arterial chemotherapy. *AJR* 2002; 179:1611–1617
 7. Tanaka T, Arai Y, Inaba Y, et al. Radiologic placement of side-hole catheter with tip fixation for hepatic arterial infusion chemotherapy. *J Vasc Interv Radiol* 2003; 14:63–68
 8. Seki H, Kimura M, Yoshimura N, Yamamoto S, Ozaki T, Sakai K. Hepatic arterial infusion chemotherapy using percutaneous catheter placement with an implantable port: assessment of factors affecting patency of the hepatic artery. *Clin Radiol* 1999; 54:221–227
 9. Wacker FK, Boese-Landgraf J, Wagner A, Albrecht D, Wolf KJ, Fobbe F. Minimally invasive catheter implantation for regional chemotherapy of the liver: a new percutaneous transsubclavian approach. *Cardiovasc Intervent Radiol* 1997; 20:128–132
 10. Herrmann KA, Wagershauser T, Sittek H, Reiser MF. Liver intraarterial chemotherapy: use of the femoral artery for percutaneous implantation of catheter-port systems. *Radiology* 2000; 215:294–299
 11. Habbe TG, McCowan TC, Goertzen TC, Leveen RF, Culp WC, Tempero MA. Complications and technical limitations of hepatic arterial infusion catheter placement for chemotherapy. *J Vasc Interv Radiol* 1998; 9:233–239
 12. Jung HY, Shim HJ, Kwak BK, et al. Percutaneously implantable catheter-port system for chemotherapeutic infusion through the hepatic artery. *AJR* 1999; 172:641–644
 13. Inaba Y, Arai Y, Matsueda K, Takeuchi Y, Aramaki T. Right gastric artery embolization to prevent acute gastric mucosal lesions in patients undergoing repeat hepatic arterial infusion chemotherapy. *J Vasc Interv Radiol* 2001; 12:957–963
 14. Yamagami T, Kato T, Iida S, Tanaka O, Nishimura T. Withdraw of implanted port-catheter for hepatic arterial infusion chemotherapy with fixed catheter tip technique. *J Vasc Interv Radiol* 2003; 14:639–642



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CT-guided needle biopsy of lung lesions: A survey of severe complication based on 9783 biopsies in Japan

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Abstract

Purpose: The aim of our study was to update the rate of severe complications following CT-guided needle biopsy in Japan via a mailed survey.

Materials and methods: Postal questionnaires regarding CT-guided needle biopsy were sent out to multiple hospitals in Japan. The questions regarded: the total number and duration of CT-guided lung biopsies performed at each hospital, and the complication rates and numbers of pneumothorax, hemothorax, air embolism, tumor seeding, tension pneumothorax and other rare complications. Each severe complication was followed with additional questions.

Results: Data from 9783 biopsies was collected from 124 centers. Pneumothorax was the most common complication, and occurred in 2412 (35%) of 6881 cases. A total of 39 (35%) hospitals reported 74 (0.75%) cases with severe complications. There were six cases (0.061%) with air embolism, six cases (0.061%) with tumor seeding at the site of the biopsy route, 10 cases (0.10%) with tension pneumothorax, six cases (0.061%) with severe pulmonary hemorrhage or hemothorax, nine cases (0.092%) with hemothorax, and 27 cases (0.26%) with others, including heart arrest, shock, and respiratory arrest. From a total of 62 patients with severe complications, 54 patients (0.55%) recovered without sequela, however one patient (0.01%) recovered with hemiplegia due to cerebral infarction, and the remaining seven patients (0.07%) died.

Conclusions: This is the first national study documenting severe complications with respect to CT-guided needle biopsy in Japan. The complication rate in Japan is comparable to internationally published figures. We believe this data will improve both clinicians as well as patients understanding of the risk versus benefit of CT-guided needle biopsy, resulting better decisions.

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Keywords: CT-guided needle biopsy; Complication; Lung nodule

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1. Introduction

Transthoracic needle biopsy is a common procedure used mainly to elucidate the nature of pulmonary nodules [1,2]. CT has rapidly become the guidance modality of choice for performing transthoracic needle biopsy due to technical advances in CT and its better detection of pulmonary lesions, which sometimes cannot be identified on chest radiograph [3].

CT-guided needle biopsy is generally regarded as a safe procedure, although pneumothorax and other rare complications can sometimes occur [4]. There have been occasional reports of deaths due to severe complications, such as, air embolism following lung biopsy [5]. Fortunately, these complications are generally very rare; previously published data shows wide variations in complication rates, making them difficult to generalize [5–8].

The aim of our study was to update the rate of severe complications following CT-guided needle biopsy in Japan via a mailed survey.

2. Materials and methods

Postal questionnaires regarding CT-guided needle biopsy were sent out to named radiologists at 101 university hospitals and cancer centers in Japan in August 2001. The radiologists at these hospitals were asked to pass duplications of the questions to other associate hospitals. The questions required information regarding: the total number and duration of CT-guided lung biopsies performed at each hospital, and the complication rates, numbers of pneumothorax, hemothorax, air embolism, tumor seeding, tension pneumothorax, severe pulmonary hemorrhage or hemoptysis which was treated with drugs for hemostasis and other rare complications, and mortalities and morbidities after that.

We defined a case as having a severe complication when one of the following criteria was met: (1) the duration of hospital stay was prolonged due to the biopsy, (2) a special technique or treatment was required to treat the complication, (3) a special procedure was required for resuscitation, and (4) shock or pre-shock developed. Each severe complication was followed with additional questions, including diagnosis of the complication, the position of the pulmonary lesion, the distance of the pulmonary lesion from the peripheral pleura, whether the lesion was located near the hilum or large pulmonary vessel, whether there was any reasonable factor causing the complication such as cough during biopsy, biopsy technique (CT-fluoroscopy or Co-axial method), the number of biopsies for each case, type and size of the needle, and presence of significant sequela from the complication.

Furthermore, the questionnaire included the following enquiries: whether emergency medication was prepared for resuscitation in the operating room, whether the patient was treated by the intravenous route and monitors, such as automatic sphygmomanometer, pulse oximetry, and electrocar-

diography. Finally, availability of access to other departments in case of emergency was questioned. Postal replies of questionnaire had been received for a year, and these answers were analyzed.

3. Results

A total of 9783 biopsy data were collected from 124 centers. The average number of biopsies performed per center was 79 cases, and that per center per year was 21 cases. The number of institutions in which hyperbaric oxygen recompression can be performed was 41 of 114 (37%) hospitals. Patients were kept on peripheral intravenous drip infusion in 86 of 92 (93%) hospitals, automatic sphygmomanometer in 38 of 92 (41%) hospitals, pulse oximetry in 32 of 92 (35%) hospitals, and electrocardiography in 8 of 92 (9%) hospitals.

Pneumothorax was the most common complication, and occurred in 2412 (35%) of 6881 cases. The number of centers that reported severe complications was 39 (35%) of 114 centers. The total number of overall severe complications was 74 (0.75%) cases. Of these, details of the complications in 64 cases are described in Table 1. There were six cases (0.061%) with air embolism, six cases (0.061%) with tumor seeding at the site of the biopsy route, 10 cases (0.10%) with tension pneumothorax, six cases (0.061%) with severe pulmonary hemorrhage or hemoptysis, 10 cases (0.10%) with hemothorax, and 26 cases (0.26%) with others. The others included 14 cases of pneumothorax requiring temporal drainage of the pneumothorax or chest tube insertion, three cases of heart arrest, and so on. There was no report of coughing during needle placement into the thorax in any of the cases with air embolism. Two of six pulmonary lesions were complicated with air emboli located near the large pulmonary vessel, and one lesion contained a cavity (Table 2). Tumor seeding occurred in two cases following CT-guided biopsy performed

Table 1
Summary of 64 cases of severe complications

| Severe complications | No. |
|--|-----|
| Pneumothorax requiring drainage of air | 14 |
| Tension pneumothorax | 10 |
| Hemothorax | 10 |
| Air embolism | 6 |
| Tumor seeding | 6 |
| Pulmonary hemorrhage of hemoptysis | 6 |
| Heart arrest | 3 |
| Respiratory arrest | 1 |
| Shock | 1 |
| Cyanosis | 1 |
| Cardiac tamponade | 1 |
| Pneumomediastinum | 1 |
| Mediastinal hematoma | 1 |
| Loss of consciousness | 1 |
| Severe pain of biopsied site | 1 |
| disseminated intravascular coagulation (DIC) | 1 |
| Total | 64 |

Table 2
Summary of cases of air embolism

| No. | Age | Sex | Size (mm) | Location (lobe) | Distance from pleura (mm) | Large vessel near the nodule | Cavity | CT-fluoroscopy | Co-axial method | No. of biopsy | Technique of biopsy | Size of the needle | Sequela |
|-----|-----|-----|-----------|-----------------|---------------------------|------------------------------|--------|-----------------|-----------------|---------------|---------------------|--------------------|--------------------|
| 1 | 72 | F | 20 | Left lower | 40 | Yes | No | Yes | No | 2 | Core biopsy | 18G | Death |
| 2 | 59 | M | 10 | Left lower | 20 | No | No | NA ^a | Yes | 1 | Core biopsy | 18G | Totally improved |
| 3 | 57 | F | 7 | Right middle | 25 | No | No | Yes | No | 1 | Core biopsy | 18G | Totally improved |
| 4 | 74 | M | 20 | Right upper | 25 | Yes | No | Yes | No | 2 | Core biopsy | 20G | Partially improved |
| 5 | 57 | M | 12 | Right lower | 3 | No | No | No | Yes | 1 | Core biopsy | 20G | Totally improved |
| 6 | 75 | M | 25 | Right lower | 18 | No | Yes | No | No | 1 | Core biopsy | 18G | Totally improved |

^a NA, information was not available.

by the Co-axial method (Table 3). In one of these two cases, the tip of the outer cannula was placed within the chest wall, so that seeding obviously occurred by direct contact of the inner needle with the biopsy route.

From a total of 62 cases with severe complications, 54 cases (0.55%) were recovered without sequela, and one case (0.01%) recovered but with hemiplegia due to cerebral infarction. Unfortunately, four (0.04%) of the remaining seven cases died just after the CT-guided biopsy procedure; these consisted of one case of air embolism, one case of DIC, and two cases of heart arrest. Three cases (0.03%) of the remaining seven cases died several years later due to tumor seeding. Four cases complicated with air embolism, three of which were treated with hyperbaric oxygen recompression, were recovered without sequela out of a total of six cases. In 23 (50%) of 46 centers, an emergency team was able to attend when a severe complication occurred.

4. Discussion

Recently, many small pulmonary lesions, which cannot be detected on chest radiograph, have been easily visualized by CT examination in daily clinical work. These lesions are usually followed with CT, or in some cases these are biopsies using CT-guided technique. CT-guided needle biopsy is a widely accepted technique and is one of the principal methods for evaluating a pulmonary lesion [9]. Although it is not rare to have minor complications due to CT-guided needle biopsy, such as, a small amount of pneumothorax and pulmonary hemorrhage, these complications improve without any treatment [5]. On the other hand, it is well known that potentially life-threatening complications such as air embolism and tumor seeding can occur. Fortunately, the frequency of these complications is considered very rare [5]. However, the number of published reports has shown that the incidence of air embolism has been increasing over the last several years. Only seven cases with air embolism were documented in the 20 years before 1995 [10–16], whereas six cases have already been published in the last 10 years [17–22].

This is the first national research study demonstrating the incidence rate of severe complications with respect to CT-guided needle biopsy based on a large number of biopsy cases using a multi-center survey.

The most common complication of transthoracic percutaneous needle biopsy is pneumothorax, with a frequency rate of 0–61%, whereas the incidence of pneumothorax requiring chest tube drainage ranges from 1.6% to 17% [23]. In the present study, the rate of pneumothorax was 35.1%, which is considered comparable to the previous studies.

Sinner's review of the literature determined that there were two cases suspected of air embolism in 2726 patients [5]. He estimated that the relative risk of air embolism per patient was about 0.07%. In the present study of 9783 biopsies, air embolism occurred in six patients, resulting in an incidence

Table 3
Summary of cases of tumor seeding

| No. | Age | Sex | Size (mm) | Location | Distance from pleura (mm) | Co-axial method | No. of biopsy | Technique of biopsy | Size of the needle |
|-----|-----|-----|-----------|-------------|---------------------------|-----------------|---------------|---------------------|--------------------|
| 1 | 72 | M | 30 | Right upper | 0 | No | 1 | Core biopsy | 18G |
| 2 | 73 | M | 30 | Left lower | 30 | Yes | 3 | Core biopsy | 18G |
| 3 | 71 | M | 10 | Right upper | 20 | No | 2 | Aspiration biopsy | 22G |
| 4 | 30 | F | 28 | Left upper | 76 | No | 2 | Core biopsy | 18G |
| 5 | 69 | M | 15 | Right lower | 0 | No | 2 | Core biopsy | 21G |
| 6 | 77 | M | 12 | Right upper | 30 | Yes | 2 | Core biopsy | 20G |

rate of 0.06%, which also shows no major difference from the previously reported complication rate. However, in the present study, there were several cases of severe complications including cardiac and respiratory arrest, and shock, which can be secondary to air embolism, although it is very difficult to confirm air embolism in the coronary artery in cases of myocardial infarction when the patient has not been scanned at the level of the heart. It is speculated that concurrent cough during the procedure has a high possibility of an air embolism misplacing the biopsy needle into the large vessel adjacent to the pulmonary lesion. Among the total of six cases with air emboli in the present study, two cases demonstrated biopsied pulmonary lesions located close to the large vessels, however the remaining four cases have no close relation to the large vessels. There were no reports of coughing during the procedure in any of the cases complicated by air embolism. Air embolism even occurred in a case in which the nodule was very near the pleura (case no. 5). In our study, all cases with air emboli had undergone CT-guided biopsy using a core biopsy needle of 18–20 gauge, which is greater in diameter than the usually used fine aspiration needles. Having said that, in the previous reviews, most cases with air emboli were biopsied by fine aspiration needles, and there are two prior reports of air embolism following CT-guided lung needle marking using thin needles without recent biopsy [24–26].

Tumor seeding into the needle tract seems to be a rare possibility in several case reports [27–34]. There were six cases (0.06%) of tumor seeding in our study, which is a relatively high frequency compared to previous studies [5,35]. The true incidence of tumor seeding along the needle may be underestimated as not all cases can be diagnosed, and many patients die before these metastases become clinically apparent. Tumor seeding appears to depend on the size of the needle, therefore large-bore needles carry a relatively greater risk of tumor seeding, however tumor seeding following a fine needle aspiration was reported in one case of our study. It is thought that CT-guided biopsy performed using the Co-axial method has less frequency of tumor seeding as the outer cannula minimizes direct contact of the tumor cells with the biopsy route. Surprisingly, tumor seeding occurred in two cases using the Co-axial method. We speculate that the outer cannula was not appropriately placed.

Unfortunately, there were seven patients (0.07%) who died in our study due to complications in the CT-guided needle biopsy. Greene [6] estimated the mortality rate associated with fine needle aspiration to be 0.02%, how-

ever Richardson et al. [8] reported eight deaths (0.15%) in their study due to complications in CT-guided needle biopsy. Most of the deaths in the present study were attributed to fatal air embolism. Three cases of air embolism that were treated with hyperbaric oxygen recompression were recovered without sequela, which may suggest hyperbaric oxygen recompression therapy is effective for treatment of air embolism, and for reducing the mortality rate.

Our study has several limitations, including selection bias, the long period of the study, multi-center analysis with a large variety of techniques and CT scanners, and the possibility of missing or misdiagnosing significant complications such as the number of air emboli and tumor seeding. Moreover, our study is a retrospective questionnaire-based analysis rather than a prospective survey.

In conclusion, this is the first nation-wide study documenting severe complications with respect to CT-guided needle biopsy in Japan. The complication rate in Japan is comparable to internationally published figures. We believe this data will improve both clinicians as well as patients understanding of the risk versus benefit of CT-guided needle biopsy, resulting better decisions.

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References

- [1] Sinner WN. Pulmonary neoplasms diagnosed with transthoracic needle biopsy. *Cancer* 1979;43:1533–40.
- [2] Klein JS, Zarka MA. Transthoracic needle biopsy. *J Thorac Imag* 1997;12:232–49.
- [3] Hirose T, Mori K, Machida S, et al. Computed tomographic fluoroscopy-guided transthoracic needle biopsy for diagnosis of pulmonary nodules. *Jpn J Clin Oncol* 2000;30:259–62.
- [4] Berquist TH, Bailey PB, Cortese DA, et al. Transthoracic needle biopsy: accuracy and complication in relation to location and type of lesion. *Mayo Clin Proc* 1980;55:475–81.
- [5] Sinner WN. Complications of percutaneous transthoracic needle aspiration biopsy. *Acta Radiol Diag* 1976;17:813–28.

- [6] Greene RE. Transthoracic needle aspiration biopsy. In: Athanasoulis CA, Pfister RC, Greene RE, Robertson GH, editors. *Interventional radiology*. Philadelphia: Sanders; 1982. p. 587–634.
- [7] Klein JS, Zarka MA. Transthoracic needle biopsy. *Radiol Clin North Am* 2000;38:235–66.
- [8] Richardson CM, Pointon KS, Manhire AR, et al. Percutaneous lung biopsies: a survey of UK practice based on 5444 biopsies. *Br J Radiol* 2002;75:731–5.
- [9] Belfiore G, Filippo SD, Guida C, et al. CT-guided needle biopsy of lesions. *Nucle Med Biol* 1994;21:713–9.
- [10] Wescott JL. Air embolism complicating percutaneous needle biopsy of the lung. *Chest* 1973;63. pp. 108–108.
- [11] Aberle DR, Gamsu G, Golden JA. Fatal systemic arterial air embolism following lung needle aspiration. *Radiology* 1987;165:351–3.
- [12] Cianci P, Posin JP, Shimshak RR, et al. Air embolism complicating percutaneous thin needle biopsy of lung. *Chest* 1987;92:749–50.
- [13] Tolly TL, Feldmeier JE, Czarnecki D. Air embolism complicating percutaneous lung biopsy. *AJR Am J Roentgenol* 1988;150:555–6.
- [14] Baker BK, Awwad EE. Computed tomography of fatal cerebral air embolism following percutaneous aspiration biopsy of the lung. *JCAT* 1988;12:1082–3.
- [15] Worth ER, Burton RJ, Landreneau RJ, Eggers GWN, et al. Left atrial air embolism during intraoperative needle biopsy of a deep pulmonary lesion. *Anesthesiology* 1990;73:342–5.
- [16] Wong RS, Ketai L, Temes RT, Follis FM, et al. Air embolus complicating transthoracic percutaneous needle biopsy. *Ann Thorac Surg* 1995;59:1010–1.
- [17] Khatri S. Cerebral artery gas embolism (CAGE) following fine needle aspiration biopsy of the lung. *Aust NZ J Med* 1997;27. pp. 27–27.
- [18] Regge D, Gallo T, Galli J, et al. Systemic arterial air embolism and tension pneumothorax: two complications of transthoracic percutaneous thin-needle biopsy in the same patient. *Eur Radiol* 1997;7:173–5.
- [19] Kodama F, Ogawa T, Hashimoto M, et al. Fatal air embolism as a complication of CT-guided needle biopsy of the lung. *JCAT* 1999;23:949–51.
- [20] Shetty PG, Fatterpekar GM, Manohar S, et al. Fat cerebral air embolism as a complication of transbronchoscopic lung biopsy: a case report. *Aust Radiol* 2001;45:215–7.
- [21] Arnold BW, Zwiebel WJ. Percutaneous transthoracic needle biopsy complicated by air embolism. *AJR Am J Roentgenol* 2002;178:1400–2.
- [22] Mokhlesi B, Ansaarie I, Bazen B, et al. Coronary artery air embolism complicating a CT-guided transthoracic needle biopsy of the lung. *Chest* 2002;121:993–6.
- [23] Laurent F, Montaudon M, Latrabe V, et al. Percutaneous biopsy in lung cancer. *Eur J Radiol* 2003;45:60–8.
- [24] Ohi S, Ito Y, Keiya H, et al. Air embolism following computed tomography-guided lung needle marking; report of a case. *Kyobu-Geka* 2004;57:421–3.
- [25] Kamiyoshihara M, Sakata K, Ishikawa S, et al. Cerebral arterial air embolism following CT-guided lung needle marking; report of a case. *J Cardiovasc Surg* 2001;42:699–700.
- [26] Sakiyama S, Kondo K, Matsuoka H, et al. Fatal air embolism during computed tomography-guided pulmonary marking with a hook-type maker. *J Thorac Cardiovasc Surg* 2003;126:1207–9.
- [27] Muller NL, Bergin CJ, Miller RR, et al. Seeding of malignant cells into the needle track after lung and pleural biopsy. *J Can Assoc Radiol* 1986;37:192–4.
- [28] Redwood N, Beggs D, Morgan WE. Dissemination of tumor cells from fine needle biopsy. *Thorax* 1989;44:826–7.
- [29] Berger RL, Dargan EL, Huang BL, et al. Dissemination of cancer cells by needle biopsy of the lung. *J Thor Cardiovasc Surg* 1972;63:430–2.
- [30] Freise G, Larios R, Takeno Y, et al. Cell dissemination and implantation of neoplasms through biopsy and excision of malignant tumors. *Dis Chest* 1967;52:485–9.
- [31] Christensen ES. Iatrogenic dissemination of tumor cells. Dissemination of tumour cells along the needle track after percutaneous, transthoracic lung biopsy. *Danish Med Bull* 1978;25:82–7.
- [32] Ferrucci JT, Wittenberg J, Margolies MN, et al. Malignant seeding of the tract after thin-needle aspiration biopsy. *Radiology* 1979;130:345–6.
- [33] Yoshikawa T, Yoshida J, Nishimura M, et al. Lung cancer implantation in the chest wall following percutaneous fine needle aspiration biopsy. *Jpn J Clin Oncol* 2000;30:450–2.
- [34] Kara M, Alver G, Sak SD, Kavukcu S. Implantation metastasis caused by fine needle aspiration biopsy following curative resection of stage IB non-small cell lung cancer. *Eur J Cardiothor Surg* 2001;20:868–70.
- [35] Ayar D, Golla B, Lee JY, Nath H. Needle-track metastasis after transthoracic needle biopsy. *J Thorac Imag* 1998;13:2–6.

I IVRはいま：最近の動向

6. CTガイド下IVR

頭頸部悪性腫瘍による消化管狭窄症例に対するCTガイド下経皮的胃瘻造設術を中心に

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荒平 聡子/関口 隆三/直井 国治/中屋 良宏 国立がんセンター東病院放射線部
前田 哲雄/佐竹 光夫 国立がんセンター中央病院放射線診断部

CTガイド下IVRは、IVR CTアンギオシステムの導入¹⁾と、高速連続スキャン技術、高速画像再構成技術、連続画像表示技術等の発達でリアルタイムに画像が得られるCT透視の臨床応用によって、従来のX線透視や超音波では誘導・施行が困難であった病変や領域でひろく行われてきている。当施設でも、各種領域で臨床応用しているが、本稿では、頭頸部悪性腫瘍による消化管狭窄症例に対するCTガイド下経皮的胃瘻造設術（CT fluoroscopic Percutaneous Gastrostomy：CT-PG）について概説する。

CT-PGの有用性

頭頸部悪性腫瘍による消化管閉塞症例では、安全な栄養ルートを確認することは重要な問題である。栄養ルートの確保の方法としての中心静脈栄養法は、敗血症などの問題があり、長期管理に適さず、経鼻胃管がまず用いられている。しかしながら、頭頸部悪性腫瘍による消化管閉塞症例で経鼻胃管は留置不可能なことが多く、また、長期留置には合併症も少なくない。胃瘻あるいは胃十二指腸瘻の造設は、中心静脈栄養法や経鼻胃管による流動食注入に比べ、コスト、安全性、患者の日常生活の快適性に優れているとされ²⁾、当施設では胃瘻造設術を選択している。胃瘻造設術の方法としては、まず経皮的内視鏡的造設術（Percutaneous Endoscopic Gastrostomy：PEG）が選択されるが、その多くが内視鏡通過困難な頭頸部悪性腫瘍による消化管閉塞症例ではPEGの適応とならず、侵襲が大きな開腹的造設術を選択せざるを得なかった。1998年から2001年までに当施設で取り扱った頭頸部悪性腫瘍症例は、1358例であり、そのうち44例の消化管閉塞症例に対し、IVR CTアンギオシステム（東芝社製）を用いCT-PGを施行している。これら症例の経験をもとに、CT-PGの手技手順と留意点および成績について述べる。

CT-PGの手技手順と留意点

CT-PGの手技手順のシェーマを図1に、使用器具を図2に示す。

1) 穿刺部確認

穿刺部近くの皮膚に、CTで可視可能なマーカーを1cm間隔で水平に置き、単純CTを撮影する。胃内腔の状況、胃の形状と周囲臓器（特に大腸）との関係を十分に把握し、穿刺部位を確認する。少量でも飲水可能な症例では、発泡散の内服を併用する。

留意点：拡張した大腸（または小腸）が胃の全面に位置しルートがとれない症例は、CT-PGの適応とならない場合がある。局所麻酔下で徒手的に大腸が回避可能となるか、あるいは空気注入で分離される場合もあると思われるが、腸管損傷は重篤となる危険性が高く³⁾、十分な注意が必要とされる。経肝的なアプローチしかとれない場合、経皮経肝的胃瘻造設術の報告はあるが⁴⁾、われわれは、空気注入し胃が拡張していくと胃壁が肝から離れていく症例を経験しており、胃拡張が得られた時点で肝を経由しないルートを再度検索するようにしている。

2) 直接穿刺、空気注入

胃拡張（図3）

ブスコパン1A（禁忌例にはグルカゴン1A）、ソセゴン1Aを筋注し、局所麻

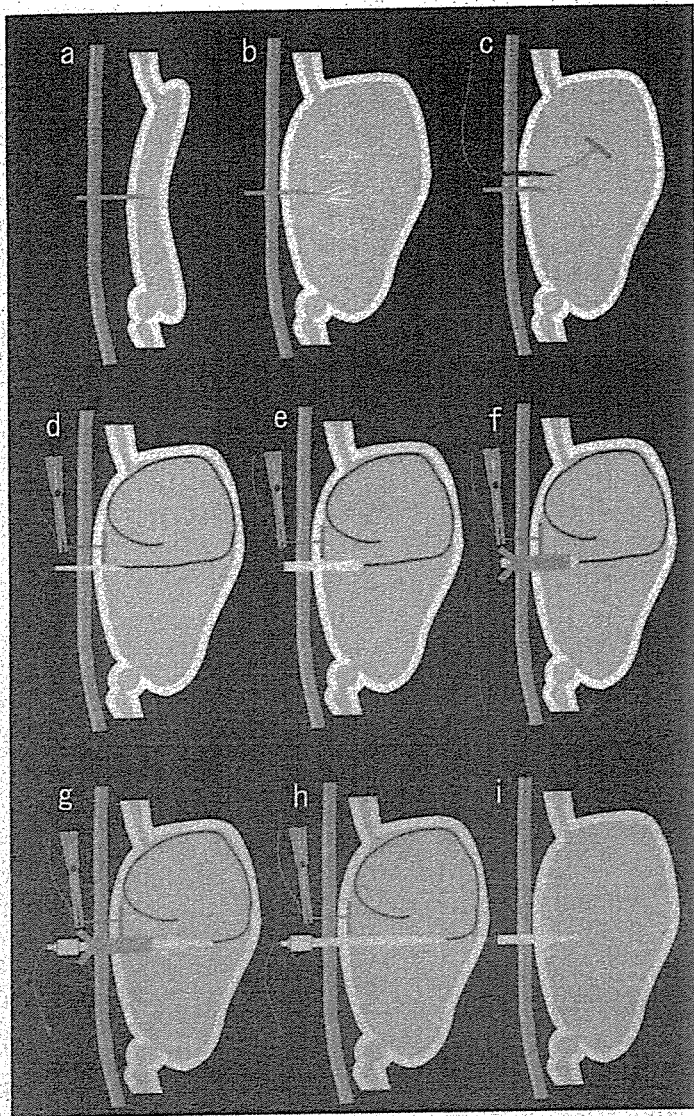


図1 手技手順のシエマ

- a : 直接穿刺
- b : 空気注入と胃拡張
- c : アンカー挿入
- d : アンカーによる胃壁固定下にガイドワイヤ挿入
- e : ルート拡張
- f : 16Fピールアウェイシース挿入
- g : 胃瘻チューブ挿入
- h : 16Fピールアウェイシース除去
- i : パラシュートを開き固定

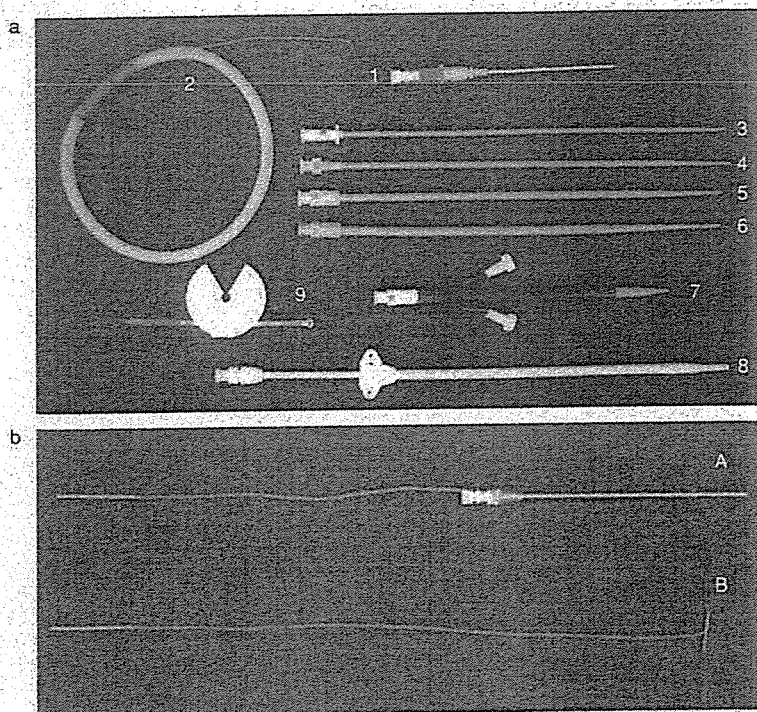


図2 CT-PG時の使用器具

a : FRICTION-LOCK™ MALECOT RUSSELL MODIFIED GASTROSTOMY SET (COOK社製)

- 1 穿刺針
- 2 ガイドワイヤ
- 3 8F ダイレータ
- 4 10F ダイレータ
- 5 12F ダイレータ
- 6 14F ダイレータ
- 7 16Fピールアウェイシース
- 8 胃瘻チューブ
- 9 固定板

b : COPE GASTROINTESTINAL SUTURE ANCHOR SET (COOK社製)

Aは18G針内に装着されたアンカー、Bは追加用アンカー。

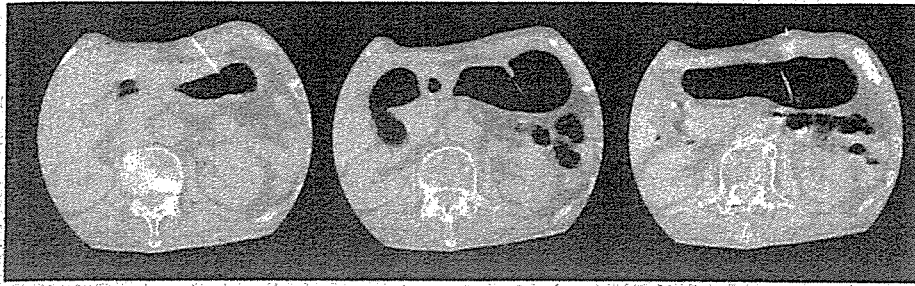


図3 直接穿刺による胃拡張



図4 アンカー挿入と胃壁固定

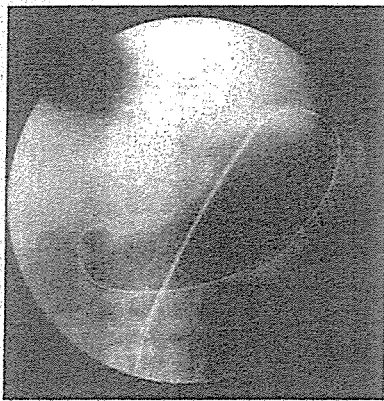


図5 ルート拡張 (8F ダイレータ)

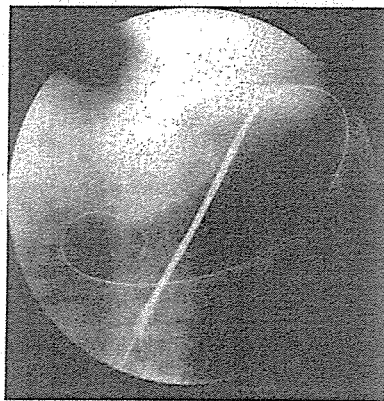


図6 ルート拡張 (16F ダイレータ)

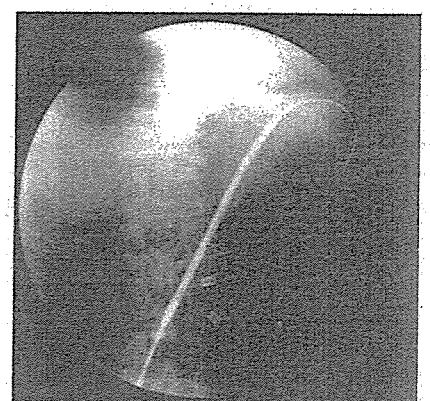


図7 16Fピールアウェイシース挿入

酔後穿刺する。拡張のない胃への初回穿刺針として、18G針では胃壁を押すだけで貫通するには太すぎるため、少なくとも21G以下の細針を用いている。あらかじめ、3方活栓付き延長チューブを装着し、CT透視下に穿針する。X線透視下に空気を約500ml注入し、十分胃内腔を拡張させる。

留意点：胃壁内注入、腹腔内注入となる場合があり、空気のひろがりに異常がある場合は、随時CT撮影にて確認する。CT透視を用いれば経時的判断が容易であるが、延長チューブを長くするなど、放射線防護に心がける。

① 胃壁外注入の場合、少量の時点であれば再施行が可能であるが、状況を判断し、後日、再施行を予定するこ

とも必要である。

② 炭酸ガスを用いれば、腹腔内あるいは血管内への誤注入時に、より安全と思われる⁹⁾。

3) アンカー挿入と胃壁固定 (図4)

十分拡張した時点で胃瘻チューブ挿入部位を想定し、その近傍から局所麻酔を追加し穿刺・挿入する。CT透視下に行うが、確実に胃壁固定ができていることを確認する。

以降の手技は、アンカーを引き上げ、胃壁固定させながらセルジンガー法に準じてX線透視下に行う⁹⁾。

4) 18Gサーフ口針穿刺

十分に局所麻酔を行い、約1.5cmほどの皮膚切開、剥離をする。穿刺は、胃壁固定しながら胃噴門側に向かうように

穿刺する。

留意点：空気追加注入あるいは少量の造影剤注入で、確実に胃内腔にあることを確認する。

5) ガイドワイヤ挿入

ワイヤの走行を確認しながら行い、十分余裕を持って留置する。

6) ルート拡張 (図5, 6)

8Fから順に手の抵抗、胃の形状、ガイドワイヤの形状を確かめながら16Fまで拡張する。

留意点：胃の形状に変化がある場合や、ガイドワイヤが屈曲する場合は、胃壁を貫通していないことが推測される。このような場合は、小径のものから拡張チューブを回しながら、胃壁を切り通す感覚で再施行する。