Intra-observer and inter-observer agreements for the measurement of dual-input whole tumor computed tomography perfusion in patients with lung cancer: Influences of the size and inner-air density of tumors

Qingle Wang1,2,3*, Zhiyong Zhang1,2,3*, Fei Shan2,3,4, Yuxin Shi12,3,4, Wei Xing5, Liangrong Shi6 & Xingwei Zhang1,2,3

1 Department of Radiology, Zhongshan Hospital, Fudan University, Shanghai, China
2 Shanghai Institute of Medical Imaging, Shanghai, China
3 Department of Medical Imaging, Shanghai Medical College, Fudan University, Shanghai, China
4 Department of Radiology, Shanghai Public Health Clinical Center, Fudan University, Shanghai, China
5 Department of Radiology, Third Affiliated Hospital of Suzhou University, Suzhou, China
6 Department of Oncology, Third Affiliated Hospital of Suzhou University, Suzhou, China

Keywords
CT perfusion; dual-blood supply; lung cancer; observer variation.

Correspondence
Fei Shan, Department of Radiology, Shanghai Public Health Clinical Center, Fudan University, No. 2901, Caolang Road, Jinshan, Shanghai 201508, China.
Tel: +86 189 3081 8986
Fax: +86 21 5724 8769
Email: shanfei@shphc.org.cn

Abstract
Background: This study was conducted to assess intra-observer and inter-observer agreements for the measurement of dual-input whole tumor computed tomography perfusion (DCTP) in patients with lung cancer.

Methods: A total of 88 patients who had undergone DCTP, which had proved a diagnosis of primary lung cancer, were divided into two groups: (i) nodules (diameter ≤3 cm) and masses (diameter >3 cm) by size, and (ii) tumors with and without air density. Pulmonary flow, bronchial flow, and pulmonary index were measured in each group. Intra-observer and inter-observer agreements for measurement were assessed using intraclass correlation coefficient, within-subject coefficient of variation, and Bland–Altman analysis.

Results: In all lung cancers, the reproducibility coefficient for intra-observer agreement (range 26.1–38.3%) was superior to inter-observer agreement (range 38.1–81.2%). Further analysis revealed lower agreements for nodules compared to masses. Additionally, inner-air density reduced both agreements for lung cancer.

Conclusion: The intra-observer agreement for measuring lung cancer DCTP was satisfied, while the inter-observer agreement was limited. The effects of tumoral size and inner-air density to agreements, especially between two observers, should be emphasized. In future, an automatic computer-aided segment of perfusion value of the tumor should be developed.

Introduction
Computed tomography perfusion (CTP) is an important functional imaging technique of pulmonary lesions for the quantitative evaluation of blood perfusion and the associated metabolic activity of tumors.1–3 CTP can be used in single or double input mode for the diagnosis of lung nodules or masses.4–8 Recently, with the advances in CT equipment and mathematical methods for analysis, lung cancer CTP with a first-pass maximum slope dual-input artery mode was introduced to routine clinical practice.9 Compared to the single input artery mode, the dual-input artery mode applied in lung CTP is based on the theory that lung cancers are supplied by both bronchial and pulmonary circulations.10,11 Using the dual-input artery mode, Yuan et al. first reported that the pulmonary index (PI) derived from dual-input whole tumor computed tomography perfusion (DCTP) analysis is a valuable...
biomarker for identifying malignancy in solitary pulmonary nodules (SPNs).\textsuperscript{12} Compared to magnetic resonance imaging (MRI) and fluorine 18 fluorodeoxyglucose (FDG) positron emission tomography (PET)/CT, CTP is more specific and accurate in the diagnosis of SPNs.\textsuperscript{13,14} Li et al. and Ohno et al. also demonstrated that DCTP analysis is useful in the prediction or early assessment of the treatment effect of chemoradiotherapy for non-small cell lung cancer (NSCLC).\textsuperscript{15,16}

Previous studies have proposed that DCTP analysis is valuable in clinical use to estimate the hemodynamics of lung cancer.\textsuperscript{9,12–17} Some of these studies involved inter-observer agreement analysis to measure advanced NSCLC or intra-observer and inter-observer agreement analysis to measure lung nodules.\textsuperscript{13–15} However, to our knowledge, tumor size-based observer agreement for measuring DCTP in lung cancer has not been reported in a major academic journal. Furthermore, the inner-air density of lung cancer (i.e. the bronchus sign and cavity), leading to an obvious difference of density in the tumoral area, might have an effect on observer agreement for the measurement of CTP and has also not been investigated in previous studies.\textsuperscript{18–23}

Therefore, the purpose of our study was to investigate the influence of tumor size and inner-air density on the agreement of measurements of the first-pass maximum slope DCTP analysis in lung cancer.

**Methods**

**Patients and clinical manifestations**

This prospective study was conducted as a part of an institutional review board-approved clinical study of DCTP in patients with lung cancer. All patients were informed of the radiation exposure and potential harm and provided written informed consent before participating. From July 2012 to July 2014, 126 consecutive patients with suspected lung cancer were prospectively enrolled. The inclusion criteria were: (i) patients with suspected lung cancer; (ii) lesions sized $\geq 1.0$ cm in diameter; (iii) solid or part-solid nodules or mass; (iv) no history of any treatment, including chemotherapy, radiation therapy, or surgery for lung cancer; and (v) no history of hypersensitivity to iodinated contrast media. The exclusion criteria were: (i) contrast media injection failure, (ii) unfit Z axial coverage for dynamic CT scan, (iii) obvious image artifacts, (iv) follow up with proven metastatic tumors or benign lesions and (v) patients lost to follow up.

Thirty-eight patients were excluded, leaving a total of 88 patients with proven bronchogenic lung cancer. The mean age was 63.4 $\pm$ 8.5 years (standard deviation [SD]; range 29–80; median 63 years). There were 55 men and 33 women. A definite histopathological diagnosis was obtained in 83 subjects by tumor resection (51 patients), CT-guided biopsy (14 patients), and bronchofibroscopic biopsy (18 patients). The remaining patients were diagnosed by sputum examination (1 patient) or clinical treatment and follow-up (4 patients). Forty-eight patients had adenocarcinomas, 20 squamous cell carcinomas, seven small cell lung carcinomas, two sarcomatoid carcinomas, six NSCLC without definite pathological types, one poorly differentiated carcinoma, and four patients had malignant lung cancers clinically detected during follow-up.

**Dual-input whole tumor computed tomography perfusion (DCTP)**

Dual-input whole tumor computed tomography perfusion was performed using a 320-MDCT scanner (Aquilion ONE, Toshiba Medical Systems, Otawara, Japan). All patients were trained to hold their breath before examination. An 18-gauge cannula was placed in the antecubital vein. A plain CT scan of the entire thorax (120 kV, detector configuration 160 $\times$ 0.5 mm, rotation time 0.75 seconds, pitch 0.869, slice thickness 5.0 mm, reconstruction interval 5.0 mm) was taken to localize the tumor. A dynamic volume scan was then taken using a flexible Z axial range of 6–16 cm (median 10 cm) to cover the entire tumor and the trunk of the pulmonary artery and left atrium. The DCTP dynamic volume scan mode has 18 intermittent volumes comprising three parts with different temporal intervals (TI) were acquired using breath-hold behavior for a duration of approximately 31 seconds: a TI of two seconds for the first three scans from one to five seconds, a TI of 1.5 seconds for nine subsequent scans from 6.5 to 19 seconds, and a final TI of two seconds for the last six scans from 21 to 31 seconds.

**Figure 1** The time sequence display of the dynamic volume scan using the Aquilion ONE system. After bolus injection, 18 intermittent volumes comprising three parts with different temporal intervals (TI) were acquired using breath-hold behavior for a duration of approximately 31 seconds: a TI of two seconds for the first three scans from one to five seconds, a TI of 1.5 seconds for nine subsequent scans from 6.5 to 19 seconds, and a final TI of two seconds for the last six scans from 21 to 31 seconds.
different temporal intervals (TI = 1.5 or 2 seconds). The total scan time was 30.5 seconds (the time sequence display is shown in Fig 1; 1 mm thickness and 1 mm interval, 512 × 512 matrix, 320 mm field of view [FOV], reconstruction kernel FC 02, 100 mA, 0.5 seconds gantry rotation time). The adjusted parameters of the dynamic scan included the tuber voltage according to the patient’s body mass index (BMI). The standard for chest CTP was used: BMI ≤ 20, 80 kV in 14 patients; 20 < BMI < 30, 100 kV for 74 patients; and BMI ≥ 30, 120 kV did not apply to any of the patients.

The contrast medium (CM) dose (350 mg I/mL) was also adjusted according to patient weight: at least 35 mL for patients ≤70 kg; if the patient weighed >70 kg, then the dose would be equal to the weight multiplied by 0.5 mL/kg. The CM was injected using a dual-head power injector (REF XD 2060 Touch, Ulrich Medical, Ulm, Germany) at an injection rate of 8 mL/seconds, followed by a 30 mL saline flush at the same injection rate.

The recorded dose-length product and the estimated effective dose (measured with a conversion factor of 0.014 mSv/mGy × cm) for the dynamic DCTP scan were 395.49 ± 157.20 (SD) mGy × cm (median 376.7 mGy × cm) and 5.54 ± 2.20 (SD) mSv (median 5.27 mSv).

Quantitative DCTP analysis

Quantitative DCTP analysis was performed using perfusion software for the CT equipment (Body Perfusion, Toshiba Medical Systems, Otawara, Japan). Two radiologists with eight and three years experience in chest CTP imaging measured the perfusion parameters independently to evaluate inter-observer agreement. The second radiologist calculated the measurement twice a month to evaluate intra-observer agreement. Both radiologists were blinded to all clinical data and results.

The DCTP analysis was calculated using the maximum slope mode. The perfusion parameters contain the bronchial flow (BF), pulmonary flow (PF), and the perfusion index (PI, PI = PF/[PF + BF]). A global value representing the perfusion parameter for the entire tumor was calculated as the median of the perfusion value of all measured slices. The regions of interest (ROIs) were drawn free-hand around the solid part of the tumor, excluding the surrounding air, calcification, and nearby pulmonary vessels. The radiologist also placed circular 10 mm diameter ROIs in the bilateral lung parenchyma, excluding the large vessels, to calculate the signal-to-noise ratios (SNRs) of the first dynamic CT image. Subsequently, the SNRs in both the tumor and lung parenchyma were calculated.

Statistical analysis

The patient sample was divided into two groups depending on tumor size (nodules ≤3 cm and mass >3 cm) or inner density (with or without bronchus sign or cavity). Bland-Altman statistics were then used to assess the intra-observer and inter-observer agreements of DCTP measurements. The target percent of variation for reproducibility is <30%. The within-subject coefficient of variation (WCV) and single-measure intraclass correlation coefficient (ICC) were also calculated. All statistical analyses were performed using SPSS version 18.0 (SPSS Inc., Chicago, IL, USA) and MedCalc software version 12.2.1 (MedCalc Software, Ostend, Belgium). A two-sided P value of <0.05 was considered statistically significant.

### Table 1 Dual-input computed tomography perfusion parameters measured in patients with lung cancer

| Perfusion parameters | Masses (n = 37) | Nodules (n = 51) | Tumors without inner-air density (n = 68) | Tumors with inner-air density (n = 20) |
|----------------------|-----------------|-----------------|-----------------------------------------|--------------------------------------|
|                      | Pulmonary flow (mL/min/100 mL) | Bronchial flow (mL/min/100 mL) | Pulmonary index (%) | Pulmonary flow (mL/min/100 mL) | Bronchial flow (mL/min/100 mL) | Pulmonary index (%) |
| Observer 1, first    | 45.07 ± 27.39   | 83.73 ± 31.69   | 35.34 ± 12.20   | 51.10 ± 32.55   | 78.80 ± 36.26   | 40.47 ± 17.50    |
| Observer 2           | 49.34 ± 35.29   | 84.69 ± 33.78   | 35.58 ± 12.00   | 54.83 ± 35.07   | 79.61 ± 38.46   | 41.55 ± 17.92    |
| Observer 1, second   | 46.65 ± 30.49   | 82.89 ± 33.77   | 36.11 ± 12.79   | 52.11 ± 34.11   | 78.56 ± 38.30   | 41.17 ± 17.87    |
| Observer 1, second   | 43.90 ± 37.00   | 82.34 ± 33.87   | 36.11 ± 12.79   | 52.11 ± 34.11   | 78.56 ± 38.30   | 41.17 ± 17.87    |
| Observer 2           | 54.83 ± 35.07   | 79.61 ± 38.46   | 41.55 ± 17.92   | 52.11 ± 34.11   | 78.56 ± 38.30   | 41.17 ± 17.87    |
| Observer 1, second   | 61.70 ± 35.00   | 60.61 ± 34.94   | 52.16 ± 22.51   | 63.07 ± 33.60   | 56.81 ± 38.83   | 54.21 ± 22.30    |
| Observer 1, second   | 66.90 ± 38.40   | 57.45 ± 35.65   | 54.53 ± 22.08   | 66.90 ± 38.40   | 57.45 ± 35.65   | 54.53 ± 22.08    |
Results

Dynamic image quality

The SNR for lung parenchyma of all patients was $30.29 \pm 9.07$ (SD, range 11.94–50.92, median 28.26), and the SNR for the tumors was $1.82 \pm 0.85$ (SD, range 0.57–4.34, median 1.60).

Intra-observer and inter-observer agreements for lung nodules and masses

There were 37 lung masses and 51 nodules. The mean mass diameter was $4.6 \pm 1.1$ cm (SD, median 4.5 cm), while that of the nodules was $2.3 \pm 0.5$ cm (SD, median 2.4 cm). The mean values of DCTP parameters measured in patients with lung nodules and masses are listed in Table 1. Observer agreements for measuring DCTP parameters in all patients are listed in Table 2. Bland–Altman plots for inter-observer agreements are shown in Figure 2. Observer agreements with nodules and masses, respectively, are listed in Table 3 and Bland–Altman plots are shown in Figure 3. The mean differences in intra-observer agreement for measuring DCTP parameters of lung masses and nodules ranged from $-1.30$ to $2.5\%$ and from $-2.14$ to $4.90\%$, and the WCVs ranged from 9.4 to 11.5% and 10.4 to 16.8%, respectively. Correspondingly, the mean difference in inter-observer agreement (except for the measurement of BF value of lung nodules), ranged from $-0.60$ to $-4.8\%$ and $-5.55$ to $-6.91\%$, and the WCVs from 12.1 to 14.4% and 14.1 to 19.8%, respectively.

Table 2 Observer agreement for measuring all lung cancers ($n = 88$)

| Perfusion parameters | Intra-observer agreement | Inter-observer agreement |
|----------------------|--------------------------|--------------------------|
|                      | ICC  | MD ± SD (%) | 95% limits (%) | RC (%) | WCV (%) | ICC  | MD ± SD (%) | 95% limits (%) | RC (%) | WCV (%) |
| Pulmonary flow       | 0.964 | $-2.90 \pm 18.0$ | $-38.2$ to $32.4$ | 35.3 | 12.8 | 0.942 | $-5.91 \pm 24.6$ | $-54.2$ to $42.4$ | 48.3 | 17.8 |
| Bronchial flow       | 0.971 | $3.84 \pm 19.5$ | $-34.5$ to $42.1$ | 38.3 | 14.0 | 0.930 | $8.50 \pm 41.4$ | $-72.7$ to $89.7$ | 81.2 | 29.7 |
| Pulmonary index      | 0.970 | $-2.78 \pm 13.3$ | $-28.9$ to $23.4$ | 26.1 | 9.6 | 0.945 | $-3.22 \pm 19.4$ | $-41.3$ to $34.8$ | 38.1 | 13.8 |

95% limits, 95% limits for mean difference; ICC, intraclass correlation coefficient for a single measure; MD, mean difference; RC, reproducibility coefficient; SD, standard deviation; WCV, within-subject coefficient of variation.

Figure 2 Intra-observer and inter-observer agreements using Bland–Altman plots for all lung cancers ($n = 88$) measured by two different observers. The solid line shows the mean value, where the mean = (value 1 + value 2)/2. The dashed lines denote the means ± (1.96 x standard deviation). Ordinate value = (value 1 − value 2)/mean x 100. Pulmonary flow (PF) 1, 2, 3 = PF measured by observer 1 the first time, observer 1 the second time, and observer 2, respectively. The same process applies to bronchial flow (BF) 1, 2, 3 and perfusion index (PI) 1, 2, 3. Graphs show Bland–Altman plots for (a) PF, (b) BF, and (c) PI values measured by observer 1 and (d) PF, (e) BF, and (f) PI values measured between observers 1 and 2.
Intra-observer and inter-observer agreements for lung cancers with or without inner-air density

There were 20 lung cancers with inner-air density, including cavities (7 tumors) and CT bronchus sign (13 tumors), and 68 lung tumors without inner-air density. Figure 4 shows a colored map of the ROIs of the whole tumor, excluding the small cavity. The mean DCTP parameter values measured in patients with or without inner-air density are listed in Table 1. The observer agreements for measuring DCTP parameters of both groups are listed in Table 4. Bland–Altman plots for agreements between the observers are shown in Figure 5. A higher intra-observer agreement in lung cancers without inner-air density was represented as low reproducibility coefficients (RCs) of approximately 25%, narrow 95% limits, and low WCVs of approximately 10%. A lower intra-observer agreement in lung cancers with inner-air density was nearly 1.5–2 times that (PI = 35.4/22.7), compared to the values of lesions without bronchus sign or cavities. Except for the PI RC and WCV values, inter-observer agreement in the two groups was almost the same. However, the intra-observer

Table 3 Observer agreement for measuring lung cancer nodules (n = 51) and masses (n = 37)

| Perfusion parameters | Intra-observer agreement | Inter-observer agreement |
|----------------------|--------------------------|--------------------------|
|                      | ICC          | MD ± SD (%)  | 95% limits (%) | RC (%) | WCV (%) | ICC          | MD ± SD (%)  | 95% limits (%) | RC (%) | WCV (%) |
| Pulmonary flow       |              |              |                |        |         |              |              |                |        |         |
| Nodules              | 0.907        | −2.14 ± 23.8 | −48.9 to 44.6  | 46.7   | 16.8    | 0.937        | −6.91 ± 27.3 | −60.5 to 46.7 | 53.6   | 19.8    |
| Masses               | 0.966        | −1.3 ± 16.4  | −33.6 to 30.9  | 32.2   | 11.5    | 0.939        | −4.8 ± 20.1  | −44.2 to 34.7 | 39.5   | 14.4    |
| Bronchial flow       |              |              |                |        |         |              |              |                |        |         |
| Nodules              | 0.976        | 4.90 ± 22.4  | −39.1 to 48.9  | 44.0   | 16.1    | 0.942        | 16.5 ± 51.0  | −83.6 to 116.6 | 100.1 | 37.6    |
| Masses               | 0.952        | 2.5 ± 14.9   | −26.8 to 31.7  | 29.3   | 10.6    | 0.891        | −0.5 ± 17.3  | −34.5 to 33.4  | 33.9   | 12.1    |
| Pulmonary index      |              |              |                |        |         |              |              |                |        |         |
| Nodules              | 0.967        | −2.31 ± 14.7 | −31.2 to 26.5  | 28.8   | 10.4    | 0.952        | −5.55 ± 19.4 | −43.6 to 32.5  | 38.0   | 14.1    |
| Masses               | 0.910        | −2.2 ± 13.2  | −28.1 to 23.8  | 26.0   | 9.4     | 0.902        | −0.6 ± 18.8  | −37.4 to 36.2  | 36.8   | 13.1    |

95% limits, 95% limits for mean difference; ICC, intraclass correlation coefficient for a single measure; MD, mean difference; RC, reproducibility coefficient; WCV, within-subject coefficient of variation.

Intra-observer and inter-observer agreements for lung cancers with or without inner-air density

There were 20 lung cancers with inner-air density, including cavities (7 tumors) and CT bronchus sign (13 tumors), and 68 lung tumors without inner-air density. Figure 4 shows a colored map of the ROIs of the whole tumor, excluding the small cavity. The mean DCTP parameter values measured in patients with or without inner-air density are listed in Table 1. The observer agreements for measuring DCTP parameters of both groups are listed in Table 4. Bland–Altman plots for agreements between the observers are shown in Figure 5. A higher intra-observer agreement in lung cancers without inner-air density was represented as low reproducibility coefficients (RCs) of approximately 25%, narrow 95% limits, and low WCVs of approximately 10%. A lower intra-observer agreement in lung cancers with inner-air density was nearly 1.5–2 times that (PI = 35.4/22.7), compared to the values of lesions without bronchus sign or cavities. Except for the PI RC and WCV values, inter-observer agreement in the two groups was almost the same. However, the intra-observer

Figure 3 Interobserver agreement using Bland–Altman plots for masses (n = 37) and nodules (n = 53) measured by two different observers. The solid line shows the mean value, where the mean = (value 1 + value 2)/2. The dashed lines denote the means ± (1.96 × standard deviation). Ordinate value = (value 1 − value 2)/mean) × 100. Pulmonary flow (PF) 1 = PF measured by observer 1, PF 3 = PF measured by observer 2. The same process applies to bronchial flow (BF) 1, BF 3, and perfusion index (PI) 1, PI 3. Graphs show Bland–Altman plots for (a) PF, (b) BF, and (c) PI values for lung masses and (d) PF, (e) BF and (f) PI values for lung nodules.
Colored maps of dual-input computed tomography perfusion in a 52-year-old man with peripheral adenocarcinoma in the right inferior lung. This nodule has a small cavity in the central portion. Air in the cavity may have resulted in miscalculation of the perfusion values. (a–c) show the maps color display of blood supply of the tumor for pulmonary flow (a), bronchial flow (b), and pulmonary flow (c). (d) shows computed tomography image without pseudo color processing.

Table 4 Observer agreement for measuring tumors with \((n = 20)\) or without inner-air density \((n = 68)\)

| Perfusion parameters | Intra-observer agreement | Inter-observer agreement |
|----------------------|--------------------------|--------------------------|
|                      | ICC          | MD ± SD | 95% limits (%) | RC | WCV (%) | ICC          | MD ± SD | 95% limits (%) | RC | WCV (%) |
| Pulmonary flow       |              |         |               |
| Tumors with air density | 0.930      | −7.91 ± 26.9 | −60.6 to 44.8 | 52.7 | 19.4 | 0.952      | −4.01 ± 24.6 | −52.1 to 44.1 | 48.1 | 17.2 |
| Tumors without air density | 0.976    | −1.43 ± 14.4 | −29.7 to 26.8 | 28.2 | 10.2 | 0.938      | −6.47 ± 24.8 | −55.1 to 42.2 | 48.6 | 18.0 |
| Bronchial flow       |              |         |               |
| Tumors with air density | 0.960      | 13.7 ± 31.0 | −47.1 to 74.5 | 60.8 | 23.5 | 0.916      | 20.4 ± 49.9 | −77.4 to 118.3 | 97.9 | 37.3 |
| Tumors without air density | 0.972     | 0.89 ± 13.5 | −25.6 to 27.4 | 24.5 | 9.50 | 0.931      | 4.94 ± 38.2 | −70.0 to 79.9 | 74.9 | 27.6 |
| Pulmonary index       |              |         |               |
| Tumors with air density | 0.967      | −6.35 ± 18.0 | −41.7 to 29.0 | 35.4 | 13.2 | 0.961      | −5.18 ± 19.2 | −42.8 to 32.4 | 37.6 | 13.7 |
| Tumors without air density | 0.968     | −1.73 ± 11.6 | −24.4 to 20.9 | 22.7 | 8.20 | 0.933      | −2.64 ± 19.6 | −41.0 to 35.8 | 38.4 | 13.9 |

95% limits, 95% limits for mean difference; ICC, intraclass correlation coefficient for a single measure; MD, mean difference; RC, reproducibility coefficient; SD, standard deviation; WCV, within-subject coefficient of variation.
agreement for both groups was decreased compared to the inter-observer agreement, with the exception of both agreements of PF and PI values approximated in lung cancers with air-density.

**Discussion**

Our results indicate that intra-observer agreement in lung cancer on the whole was adequate, while inter-observer agreement was more limited, especially for measurements of BF value in lung nodules and tumors with inner-air density. Lung cancer size and tumors with inner-air density might influence observer agreement of DCTP measurement.

This result was consistent with a previous finding of volume single-input CTP in lung cancer using motion correction. The more difficult delineation of smaller tumors might primarily reflect the use of a freehand contour drawing method. Thus, the application of automatic computer-aided delineation of the tumor and calculation of the volume perfusion value into future CT perfusion software might improve inter-observer and intra-observer agreements.

The agreements for measuring lung masses were better than those of the nodules. The lower intra-observer agreement for smaller lesions was observed with wider 95% limits and higher WCVs. Although inter-observer agreement was reduced in the nodule group, particularly the BF value, these results were acceptable for clinical use, and consistent with previous studies indicating the values for single or dual-input CTP in lung cancer. The volume scan mode, a measurement of the entire tumor and the introduction of motion correction techniques, might reflect relatively good agreement within observers.

Moreover, we showed that lung cancers without inner-air density (the cavity or the bronchus sign) had better agreement than those with inner-air density, particularly regarding observer agreement. This finding indicated that the partial volume effect produced between the parts of soft tissue, and the air in the tumor played an important role in the reproducibility of lung cancer CTP, even when the function of motion correction was used to reduce misregistered dynamic images.

We used personalized protocols of dynamic scan and CM injection based on the BMI and weight of the patients to control X-ray radiation and CM doses and to balance the image quality. To our knowledge, this study is the first published in a major academic journal. Following a previous study, we also designed a scan sequence comprising a TI of 1.5 seconds in the temporal window, covering the climax of pulmonary and systemic circulation, and a TI of two seconds before and after the window, to limit exposure time and maintain the accuracy of the perfusion calculation. The estimated radiation dose for the dynamic scan...
Observer agreements for perfusion CT

Q. Wang et al.

was 5.54 mSv. Considering the sensitivity to image noise, the maximum-slope CTP images should have an acceptable SNR (a balance must be made between temporal resolution and the SNR).\(^2\)\(^5\) To some extent, the choice should be determined by the analysis method used. Thus, time-attenuation data from protocols that adopt a higher mAs value but a lower image frequency are appropriate for compartmental analysis.\(^2\)\(^4\)\(^5\) The tumor SNR results illustrate this balance. For the broad use of CTP, detailed development, including dynamic protocol, hardware, and image algorithm are required to reduce the radiation dose and improve image quality.

This study has several limitations. First, the sample size was not predetermined in dimension with a power analysis. Second, other factors that might influence observer agreement analysis, such as the location, the duration of data acquisition, and image quality were not examined. Third, one of the two observers had limited experience in DCTP image analysis, and thus did not partake in intraobserver agreement. This choice might decrease the significance of agreements.

In conclusion, the intra-observer agreement for DCTP in lung cancer was satisfied, but the inter-observer agreement was limited. Although DCTP was helpful for the diagnosis of lung cancer, a professional radiologist experienced in DCTP is required. Overall, observer agreement of large-sized tumors with or without cavities or bronchus sign in solid parts was better than for small-sized tumors with air density. In the future, automated segment analysis based on the density, morphology, and perfusion value of lung lesions should be developed to increase the reproducibility between observers.

Acknowledgments

The National Natural Science Foundation of China (Grant No. 81301223) supported this study. We thank Professor Riwa Kishimoto for assistance with statistical work.

Disclosure

No authors report any conflict of interest.

References

1 Nishino M, Hatabu H, Johnson BE, McLoud TC. State of the art: Response assessment in lung cancer in the era of genomic medicine. Radiology 2014; 271: 6–27.

2 Miles KA, Lee TY, Goh V et al. Current status and guidelines for the assessment of tumour vascular support with dynamic contrast-enhanced computed tomography. Eur Radiol 2012; 22: 1430–41.

3 Kim H, Park CM, Goo J, Wildberger JE, Kauczor HU. Quantitative computed tomography imaging biomarkers in the diagnosis and management of lung cancer. Invest Radiol 2015; 50: 571–83.

4 Shan F, Zhang Z, Xing W et al. Differentiation between malignant and benign solitary pulmonary nodules: Use of volume first-pass perfusion and combined with routine computed tomography. Eur J Radiol 2012; 81: 3598–605.

5 Tacelli N, Remy-Jardin M, Copin MC et al. Assessment of non-small cell lung cancer perfusion: Pathologic–CT correlation in 15 patients. Radiology 2010; 257: 863–71.

6 Friaoli F, Anzidei M, Zaccagna F et al. Whole-tumor perfusion CT in patients with advanced lung adenocarcinoma treated with conventional and antiangiogenetic chemotherapy: Initial experience. Radiology 2011; 259: 574–82.

7 Tacelli N, Santangelo T, Scherpereel A et al. Perfusion CT allows prediction of therapy response in non-small cell lung cancer treated with conventional and anti-angiogenic chemotherapy. Eur Radiol 2013; 23: 2127–36.

8 Lind J, Meijerink MR, Dingemans AM et al. Dynamic contrast-enhanced CT in patients treated with sorafenib and erlotinib for non-small cell lung cancer: A new method of monitoring treatment? Eur Radiol 2010; 20: 2890–8.

9 Yuan X, Zhang J, Ao G, Quan C, Tian Y, Li H. Lung cancer perfusion: Can we measure pulmonary and bronchial circulation simultaneously? Eur Radiol 2012; 22: 1665–71.

10 Milne EN. Circulation of primary and metastatic pulmonary neoplasms. A postmortem microarteriographic study. Am J Roentgenol Radium Ther Nucl Med 1967; 100: 603–19.

11 Savai R, Wolf JC, Greschus S et al. Analysis of tumor vessel supply in Lewis lung carcinoma in mice by fluorescent microsphere distribution and imaging with micro- and flat-panel computed tomography. Am J Pathol 2005; 167: 937–46.

12 Yuan X, Zhang J, Quan C et al. Differentiation of malignant and benign pulmonary nodules with first-pass dual-input perfusion CT. Eur Radiol 2013; 23: 2469–74.

13 Ohno Y, Nishio M, Koyama H et al. Comparison of quantitatively analyzed dynamic area-detector CT using various mathematic methods with FDG PET/CT in management of solitary pulmonary nodules. AJR Am J Roentgenol 2013; 200: W593–602.

14 Ohno Y, Nishio M, Koyama H et al. Solitary pulmonary nodules: Comparison of dynamic first-pass contrast-enhanced perfusion area-detector CT, dynamic first-pass contrast-enhanced MR imaging, and FDG PET/CT. Radiology 2015; 274: 563–75.

15 Li XS, Fan HX, Fang H, Huang H, Song YL, Zhou CW. Value of whole-tumor dual-input perfusion CT in predicting the effect of multiarterial infusion chemotherapy on advanced non-small cell lung cancer. AJR Am J Roentgenol 2014; 203: W497–505.

16 Ohno Y, Koyama H, Fujisawa Y et al. Dynamic contrast-enhanced perfusion area detector CT for non-small cell lung
cancer patients: Influence of mathematical models on early prediction capabilities for treatment response and recurrence after chemoradiotherapy. *Eur J Radiol* 2016; **85**:176–86.

17 Nguyen-Kim TD, Frauenfelder T, Strobel K, Veit-Haibach P, Huellner MW. Assessment of bronchial and pulmonary blood supply in non-small cell lung cancer subtypes using computed tomography perfusion. *Invest Radiol* 2015; **50**:179–86.

18 Valentin J, International Commission on Radiation Protection. Managing patient dose in multi-detector computed tomography (MDCT). ICRP publication 102. *Ann ICRP* 2007; **37**:1–79, iii.

19 Ng QS, Goh V, Fichte H et al. Lung cancer perfusion at multi-detector row CT: Reproducibility of whole tumor quantitative measurements. *Radiology* 2006; **239**:547–53.

20 Nakayama Y, Awai K, Funama Y et al. Abdominal CT with low tube voltage: Preliminary observations about radiation dose, contrast enhancement, image quality, and noise. *Radiology* 2005; **237**:945–51.

21 Qiang JW, Zhou KR, Lu G et al. The relationship between solitary pulmonary nodules and bronchi: Multi-slice CT-pathological correlation. *Clin Radiol* 2004; **59**:1121–7.

22 Sauter AW, Merkle A, Schulze M et al. Intraobserver and interobserver agreement of volume perfusion CT (VPCT) measurements in patients with lung lesions. *Eur J Radiol* 2012; **81**:2853–9.

23 Lee SM, Lee HJ, Kim JI et al. Adaptive 4D volume perfusion CT of lung cancer: Effects of computerized motion correction and the range of volume coverage on measurement reproducibility. *AJR Am J Roentgenol* 2013; **200**:W603–9.

24 Shan F, Xing W, Qiu J, Zhang Z, Yang S. First-pass CT perfusion in small peripheral lung cancers: Effect of the temporal interval between scan acquisitions on the radiation dose and quantitative vascular parameters. *Acad Radiol* 2013; **20**:972–9.

25 Miles KA. Perfusion CT for the assessment of tumour vascularity: Which protocol? *Br J Radiol* 2003; **76** Spec No 1: S36–42.