Imaging Assessment of Visceral Pleural Surface Invasion by Lung Cancer: Comparison of CT and Contrast-Enhanced Radial T1-Weighted Gradient Echo 3-Tesla MRI

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Objective: To compare the diagnostic performance of contrast-enhanced radial T1-weighted gradient-echo 3-tesla (3T) magnetic resonance imaging (MRI) and computed tomography (CT) for the detection of visceral pleural surface invasion (VPSI). Visceral pleural invasion by non-small-cell lung cancer (NSCLC) can be classified into two types: PL1 (without VPSI), invasion of the elastic layer of the visceral pleura without reaching the visceral pleural surface, and PL2 (with VPSI), full invasion of the visceral pleura.

Materials and Methods: Thirty-three patients with pathologically confirmed VPSI by NSCLC were retrospectively reviewed. Multidetector CT and contrast-enhanced 3T MRI with a free-breathing radial three-dimensional fat-suppressed volumetric interpolated breath-hold examination (VIBE) pulse sequence were compared in terms of the length of contact, angle of mass margin, and arch distance-to-maximum tumor diameter ratio. Supplemental evaluation of the tumor-pleura interface (smooth versus irregular) could only be performed with MRI (not discernible on CT).

Results: At the tumor-pleura interface, radial VIBE MRI revealed a smooth margin in 20 of 21 patients without VPSI and an irregular margin in 10 of 12 patients with VPSI, yielding an accuracy, sensitivity, specificity, positive predictive value, negative predictive value, and F-score for VPSI detection of 91%, 83%, 95%, 91%, 91%, and 87%, respectively. The McNemar test and receiver operating characteristics curve analysis revealed no significant differences between the diagnostic accuracies of CT and MRI for evaluating the contact length, angle of mass margin, or arch distance-to-maximum tumor diameter ratio as predictors of VPSI.

Conclusion: The diagnostic performance of contrast-enhanced radial T1-weighted gradient-echo 3T MRI and CT were equal in terms of the contact length, angle of mass margin, and arch distance-to-maximum tumor diameter ratio. The advantage of MRI is its clear depiction of the tumor-pleura interface margin, facilitating VPSI detection.

Keywords: Magnetic resonance imaging; Lung cancer; Diagnostic imaging
of the TNM classification for lung cancer classifies both invasion of the elastic layer of the visceral pleura without reaching the visceral pleural surface (PL1) and full invasion of the visceral pleura (PL2) as stage T2 [2], suggesting that PL1 and PL2 tumors have similar prognosis and survival [3]. However, a study showed that the 5-year overall survival rate was significantly higher for patients with PL1 tumors (61.9%) than for those with PL2 tumors (39.2%) [4]. The 5-year survival rates of patients with and without VPSI were 57.9% and 83.0%, respectively, for N0–N2 disease and 74.3% and 88.5%, respectively, for N0 disease [1]. VPSI might therefore be considered an independent risk factor for poor prognosis, and its presence might be a potential indication for adjuvant chemotherapy [1].

Computed tomography (CT) is a noninvasive imaging modality for preoperative staging of lung cancers. However, it has limitations in the detection of more subtle cases of pleural invasion, as contiguity of the tumor with the pleural surface is not necessarily equivalent to its invasion [5]. A skirt-like 3-dimensional (3D) CT pattern of pleural morphology adjacent to the tumor [6], pleural tag with a soft tissue component [7], and type-5 border (convex border with a perpendicular or blunt angle) [8] have been shown to have predictive value for pleural invasion; however, these methods cannot be used to differentiate PL1 from PL2.

Magnetic resonance imaging (MRI) is potentially advantageous for VPSI detection because of its superior soft tissue contrast and tissue characterization properties, without ionizing radiation exposure. Conventional MRI can yield high-quality images of the thorax, particularly with breath-hold techniques; however, breath-holding can be challenging for some patients. Free-breathing, intravenous contrast-enhanced, radial, 3D ultrafast gradient-echo (volumetric interpolated breath-hold examination [VIBE]) T1-weighted imaging, hereinafter referred to as “radial VIBE,” has been proposed as an alternative to breath-hold post-contrast imaging. It enables patients to breathe freely during scanning, yielding excellent image quality and diagnostic performance owing to the enhanced lesion conspicuity, clarity of the tumor interface, and reduced respiratory motion artifacts compared to other free-breathing techniques [9,10].

Moreover, this sequence has been proven to be useful for evaluating the morphological features of lung cancer and for readily demonstrating the tumor-pleura interface [11]. Therefore, this study aimed to compare the diagnostic performance of contrast-enhanced, radial VIBE 3-tesla (3T) MRI, and CT for VPSI detection.

**MATERIALS AND METHODS**

This retrospective, bi-institutional study was conducted at Wonju Severance Christian Hospital and Samsung Medical Center. The study was approved by the Institutional Review Boards of both institutions and the local ethics committee (IRB No. CR319094). The requirement for written informed consent was waived.

**Patients**

Initially, 191 consecutive lung cancer patients who underwent both CT and 3T MRI between January 2016 and May 2019 were enrolled. Patients with poor quality CT or MRI scans were excluded (Fig. 1). Finally, a total of 33 patients met the following inclusion criteria: 1) nonmetastatic, primary NSCLC treated with surgical resection; 2) available pathology reports describing the extent of pleural invasion; 3) preoperative CT and 3T MRI scans with the same scan parameters available on a picture archiving and communication system (PACS); 4) available contrast-enhanced, radial VIBE 3-tesla (3T) MRI, and CT for VPSI detection.
free-breathing fat-saturated radial VIBE sequence on MRI; 5) suspected pleural invasion on CT; 6) CT and MRI performed within 3 months; and 7) no prior or current chemotherapy or radiation therapy. The sample including 23 patients from the Blinded Hospital and 10 from the Blinded Medical Center was comprised of 28 male and five female, aged 46–88 years (mean age, 68 ± 10 years). This study analyzed 17 adenocarcinomas, 13 squamous cell carcinomas, and three large cell carcinomas (Table 1).

### CT Scanning
A 64-channel multidetector CT scanner (Brilliance 64, Philips Medical System) was used at both institutions for breath-hold imaging in the supine position, with a 7–8 second breath-hold. The technical parameters of the CT scans were as follows: 0.625-mm detector collimation, 512 x 512 matrix, 340-mm field-of-view, 80–120-mAs tube current, 120-kV tube voltage, 2.5-mm slice thickness, and 0.5-seconds rotation time. For each patient, 350 mg/mL iohexol contrast material was administered intravenously at a rate of 2.5 cc/s [11].

### MRI Scanning
All MRI examinations were performed using a 3T system (MAGNETOM Skyra) with a 60-channel body coil. Patients were imaged in the supine position with their arms overhead to eliminate potential artifacts from the arms positioned on each side. The following pulse sequences were used in 23 patients: axial breath-hold T1- and T2-weighted turbo spin echo, axial T2-weighted half-Fourier acquisition single-shot turbo spin echo, contrast-enhanced fat-saturated T1-weighted, and contrast-enhanced free-breathing, fat-saturated radial VIBE. The pulse sequences common to all included patients were breath-hold TI-weighted and contrast-enhanced free-breathing fat-suppressed radial VIBE. Gadoteridol (0.1 mmol/kg; ProHance; Bracco Imaging) was injected at a rate of 1.5 cc/s. The radial VIBE imaging parameters with an isotropic resolution of 0.9 mm were as follows: repetition time, 3.36 ms; echo time, 1.66 ms; flip angle, 5°; field of view, 260 mm x 260 mm; and matrix size, 288 x 288 mm.

### Imaging Analysis
Two chest radiologists (readers 1 and 2, with 23 and 3 years of clinical experience, respectively), who were blinded to the patients’ clinical information and pathological results, evaluated the CT and MRI scans by consensus. To make the final decision, the radiologists were allowed to refer to the images several times and adjust the window and level settings if necessary. The observers were instructed to read all CT scans first, followed by MRI scans after 1 month to avoid interference bias between CT and MRI results in interpretation.

Tumor size (maximum dimension in any of the three planes [axial, sagittal, and coronal]) on CT and MRI scans, and tumor location were recorded. The angle of the mass margin was formed from the center of the tumor toward both ends of the pleural contact. The length of pleural contact was the length of the interface between the primary tumor and the pleura. It was drawn freehand and measured in the same image (Fig. 2B) [12,13]. The arch distance to maximum tumor diameter ratio was the length of the interface between the primary tumor and the neighboring structure (in this case, the pleura) (Fig. 2B) [14]. The imaging criteria utilized to determine the degree of

### Table 1. Clinicopathologic Features of Non-Small-Cell Lung Cancer Patients in This Study

| Clinicopathologic Features | Data          |
|----------------------------|---------------|
| No. of patients            | 33            |
| Age, years, mean ± SD (46–88) | 68.33 ± 9.96 |
| Sex, n (%)                 |               |
| Male                       | 28 (84.8)     |
| Female                     | 5 (15.2)      |
| Tumor size, mm, mean ± SD  | 53.4 ± 26.2   |
| Tumor pathology, n (%)     |               |
| Adenocarcinoma             | 17 (51.5)     |
| Squamous cell carcinoma    | 13 (39.4)     |
| Large cell carcinoma       | 3 (9.1)       |
| Tumor location, n (%)      |               |
| RUL                        | 11 (33.3)     |
| RML                        | 1 (3.0)       |
| RLL                        | 12 (36.4)     |
| LUL                        | 3 (9.1)       |
| LLL                        | 6 (18.2)      |
| T stage, n (%)             |               |
| T1                         | 1 (3.0)       |
| T2                         | 14 (42.4)     |
| T3                         | 12 (36.4)     |
| T4                         | 6 (18.2)      |
| Pleural invasion grade, n (%) |            |
| PL0                        | 11 (33.3)     |
| PL1                        | 10 (30.3)     |
| PL2                        | 3 (9.1)       |
| PL3                        | 9 (27.3)      |

LLL = left lower lobe, LUL = left upper lobe, RLL = right lower lobe, RML = right middle lobe, RUL = right upper lobe
suspicion for VPI were as follows (Fig. 2A): a high suspicion for VPI required a length of pleural contact > 5 cm or a > 180° angle between the margin of the mass and the pleural surface; moderate suspicion of VPI required a 3–5 cm contact length or a 90°–180° angle between the mass and the pleural surface; and mild suspicion of pleural invasion required a length of pleural contact < 3 cm or angle of mass margin < 90°.

Based on the smoothness or irregularity of the high signal intensity band interface between the tumor and the pleura, MRI scans were classified into two groups (Fig. 3): without VPSI, with smooth and clear margins, well-defined curvilinear pleural enhancement, and absence of protrusion of the interface between the tumor and the pleura (Fig. 3A); and with VPSI, with an irregular, undulating, or coarse margin or protrusion of the interface between the primary tumor and the pleura (Fig. 3B). The two groups were assessed in the axial, sagittal, and coronal planes. VPSI was determined if an irregular interface was observed in any plane. The lung cancer was deemed to be without VPSI only when there was a smooth tumor-pleura interface in all three planes.

**Pathological Analysis**

Pathological specimens were stained with the Verhoeff-Van Gieson stain to investigate the presence and extent of pleural invasion. A pathologist with 33 years of experience in analyses of lung specimens performed the analysis and staged the pleural invasion in each patient as follows: PL0, no pleural invasion; PL1, tumor invasion beyond the elastic layer of the visceral pleura but not the surface of the visceral pleura; PL2, tumor invasion of the surface of the visceral pleura; and PL3, tumor invasion of the parietal pleura or chest wall [15].

**Statistical Analysis**

Patient age and tumor size were assessed using the Wilcoxon rank-sum test. Sex, tumor pathology, and tumor location were assessed using Fisher's exact test between the two groups. Kappa statistics for categorical data were used to determine interobserver agreement. The Kappa result was interpreted as follows: < 0.20, slight; 0.21–0.40, fair; 0.41–0.60, moderate; 0.61–0.80, substantial; and
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> 0.81, excellent [16]. The chi-square or Fisher’s exact test for categorical data and Wilcoxon rank-sum test for continuous data were used to examine the significant differences between patients with and without VPSI. A \( p \) value < 0.05 was considered statistically significant. The diagnostic accuracy, sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), and F-score were calculated separately. The accuracies of MRI and CT for evaluating the length of contact and arch distance to maximum tumor diameter ratio were compared using the McNemar test. The receiver operating characteristic (ROC) curve analysis was used to assess the diagnostic values of CT and MRI for VPSI detection, and the area under the curve was calculated. All data analyses were performed using the SAS version 9.4 (SAS Institute Inc.).

**RESULTS**

The histopathological diagnoses of NSCLCs were as follows: 17 adenocarcinomas, 13 squamous cell carcinomas, and three large cell carcinomas. There were 11 tumors in the right upper lobe, one in the right middle lobe, 12 in the right lower lobe, three in the left upper lobe, and six in the left lower lobe (Table 1).

There were 21 patients without VPSI and 12 patients with VPSI, as determined pathologically. The mean and median tumor size of lung cancer were 49.7 ± 17.9 mm and 44 (33–55) mm, respectively, in the group without VPSI and 61.6 ± 24.2 mm and 55 (38.5–85) mm, respectively, in the group with VPSI. Although a larger tumor diameter was observed in patients with VPSI than in those without, tumor size was not significantly different between the two groups (\( p = 0.170 \)). There was no significant difference between the two groups in patient age, sex, tumor pathology, or tumor location (Table 2). Tables 3 and 4 show the \( p \) value, accuracy, sensitivity, specificity, PPV, NPV, and F-score for the length of contact, angle of mass margin, and arch distance-to-maximum on CT and MRI. The McNemar test showed no significant difference (\( p > 0.05 \)) in the accuracy of the contact length (MRI, 67%; CT, 61%), angle of mass margin > 90° (both MRI and CT, 61%), or arch distance to maximum tumor diameter ratio > 0.9 (CT, 67%; MRI, 61%) between CT and MRI (Table 5). Moreover, there was no significant difference in the area under the ROC curve for VPSI detection between CT and MRI, suggesting that the diagnostic performances of CT and MRI were similar (Fig. 4) regarding these characteristics.

Examination of the high-signal-intensity band interface between the tumor and the pleura on MRI revealed a smooth margin in 20 of 21 (95.2%) patients without VPSI and an irregular or coarse margin in 10 of 12 (83.3%) patients with VPSI. Similar signs in patients with and without VPSI were not appreciable on CT (Figs. 3, 5). The interface on MRI with the radial VIBE sequence significantly differed between patients with and without VPSI (\( p < 0.001 \)) (Table 3), and sensitivity, specificity, accuracy, PPV, NPV, and F-score were 83%, 95%, 91%, 91%, 91%, and 87%, respectively. The Kappa value of the comparison between

| Variables                                      | Without VPSI | With VPSI | \( P \)  |
|-----------------------------------------------|--------------|-----------|---------|
| Age, years, median (Q1–Q3)                    | 72 (64–77)   | 65 (54–73)| 0.060   |
| Sex, n (%)                                     |              |           | 0.133*  |
| Male                                          | 16 (76.2)    | 12 (100.0)|         |
| Female                                        | 5 (23.8)     | 0 (0.0)   |         |
| Tumor size (mm), median (Q1–Q3)               | 44 (33–55)   | 55 (38.5–85)| 0.170 |
| Tumor pathology, n (%)                        |              |           | 0.215*  |
| Adenocarcinoma                                | 12 (57.1)    | 5 (41.6)  |         |
| Squamous cell carcinoma                       | 8 (38.1)     | 5 (41.7)  |         |
| Large cell carcinoma                          | 1 (4.8)      | 2 (16.7)  |         |
| Location, n (%)                               |              |           | 0.106*  |
| RUL                                           | 6 (28.6)     | 5 (41.7)  |         |
| RML                                           | 1 (4.8)      | 0 (0.0)   |         |
| RLL                                           | 9 (42.9)     | 3 (25.0)  |         |
| LUL                                           | 0 (0.0)      | 3 (25.0)  |         |
| LLL                                           | 5 (23.8)     | 1 (8.3)   |         |

*Fisher’s exact test, †Includes PL0 and PL1, ‡Includes PL2 and PL3. LLL = left lower lobe, LUL = left upper lobe, RLL = right lower lobe, RML = right middle lobe, RUL = right upper lobe, VPSI = visceral pleural surface invasion.
the pathology and MRI was substantial at 0.800 for the radial VIBE sequence (Table 4).

**DISCUSSION**

This study showed that contrast-enhanced 3T MRI with the free-breathing fat-saturated radial VIBE sequence was superior to CT in depicting and distinguishing NSCLCs without VPSI from those with VPSI. VPSI is an independent factor for the poor prognosis of patients with NSCLCs and can influence the T stage, treatment, and prognosis [3]. Therefore, determining VPSI and the depth of pleural

**Table 3. CT and MRI Findings in Non-Small-Cell Lung Cancer Patients Without and with VPSI**

| Variables                        | Pathology                                                                 |
|----------------------------------|---------------------------------------------------------------------------|
|                                  | Without VPSI* (n = 21) | With VPSI† (n = 12) | P    |
| **Tumor size, mm, median (Q1–Q3)** |                       |                        |      |
| CT                               | 44 (32.1–54.9)        | 51 (41.7–82.7)        | 0.159|
| MRI                              | 45 (34.6–60.3)        | 54 (44.2–77.9)        | 0.165|
| **Length of contact, n (%)**     |                       |                        |      |
| CT (cm) ≤ 3 vs. > 3              | 10 (47.6)             | 2 (16.7)               | 0.133* |
| CT (cm) ≤ 5 vs. > 5              | 16 (76.2)             | 6 (50.0)               | 0.149* |
| MRI (cm) ≤ 3 vs. > 3             | 10 (47.62)            | 0 (0.0)                | 0.005* |
| MRI (cm) ≤ 5 vs. > 5             | 15 (71.4)             | 6 (50.0)               | 0.274 |
| **Angle of mass margin, n (%)**  |                       |                        |      |
| CT (°) ≤ 90 vs. > 90             | 8 (38.1)              | 0 (0.0)                | 0.030* |
| CT (°) ≤ 180 vs. > 180           | 13 (61.9)             | 12 (100.0)             | 0.538* |
| MRI (°) ≤ 90 vs. > 90            | 8 (38.1)              | 0 (0.0)                | 0.030* |
| MRI (°) ≤ 180 vs. > 180          | 13 (61.9)             | 12 (100.0)             | 0.538* |
| **Arch distance-to-maximum tumor diameter ratio, n (%)** | | | |
| CT                               | 13 (61.9)             | 3 (25)                 | 0.041 |
| MRI                              | 12 (57.1)             | 4 (33.3)               | 0.188 |
| Interface between the primary tumor and the pleura, n (%) | | | |
| Smooth and clear margin          | 20 (95.2)             | 2 (16.7)               | < 0.001* |
| Irregular or coarse margin       | 1 (4.8)               | 10 (83.3)              |      |

*Includes PL0 and PL1, †Includes PL2 and PL3. CT = computed tomography, MRI = magnetic resonance imaging, VPSI = visceral pleural surface invasion
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Table 4. Comparison of CT and MRI Findings for Prediction of Visceral Pleural Surface Invasion, with Pathology as the Reference Standard

| Variables                                | Accuracy (%) | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) | F-Score | Kappa |
|------------------------------------------|--------------|----------------|-----------------|---------|---------|---------|-------|
| **Length of contact**                    |              |                |                 |         |         |         |       |
| **CT (cm)**                              |              |                |                 |         |         |         |       |
| > 3                                      | 61           | 83             | 48              | 48      | 83      | 61      | 0.2667 |
| > 5                                      | 67           | 50             | 76              | 55      | 73      | 52      | 0.2667 |
| **MRI (cm)**                             |              |                |                 |         |         |         |       |
| > 3                                      | 67           | 100            | 48              | 52      | 100     | 69      | 0.3980 |
| > 5                                      | 64           | 50             | 71              | 50      | 71      | 50      | 0.2143 |
| **Angle of mass margin**                 |              |                |                 |         |         |         |       |
| **CT (°)**                               |              |                |                 |         |         |         |       |
| > 90                                     | 61           | 100            | 38              | 48      | 100     | 65      | 0.3092 |
| > 180                                    | 67           | 17             | 95              | 67      | 67      | 27      | 0.1418 |
| **MRI (°)**                              |              |                |                 |         |         |         |       |
| > 90                                     | 61           | 100            | 38              | 48      | 100     | 65      | 0.3092 |
| > 180                                    | 67           | 17             | 95              | 67      | 67      | 27      | 0.1418 |
| **Arch distance-to-maximum tumor diameter ratio** | | | | | | | |
| **CT (> 0.9)**                           | 67           | 75             | 62              | 53      | 81      | 62      | 0.3388 |
| **MRI (> 0.9)**                          | 61           | 67             | 57              | 47      | 75      | 55      | 0.2186 |
| **Interface**                            |              |                |                 |         |         |         |       |
| **MRI**                                  | 91           | 83             | 95              | 91      | 91      | 87      | 0.8000 |

CT = computed tomography, MRI = magnetic resonance imaging, NPV = negative predictive value, PPV = positive predictive value.

Table 5. Agreement Between Radial Volumetric Interpolated Breath-Hold Examination Magnetic Resonance Imaging and Computed Tomography When the Same Methods Were Employed

| Variables                                | Kappa Coefficient | McNemar Test (P) |
|------------------------------------------|-------------------|------------------|
| **Length of contact**                    | 0.7284            | 0.317            |
| **Length of contact**                    | 0.9333            | 0.317            |
| **Angle of mass margin**                 | 1.0000            | -                |
| **Angle of mass margin**                 | 1.0000            | -                |
| **Arch distance-to-maximum tumor diameter ratio** | 0.8787            | > 0.999          |

CT = computed tomography, MRI = magnetic resonance imaging.

Fig. 4. Summarized receiver operating characteristic curves demonstrating the area under the curve of 75% (95% CI: 58–92%), 72% (95% CI: 54–90%), 81% (95% CI: 66–96%) for CT and 79% (95% CI: 63–94%), 70% (95% CI: 51–88%), 81% (95% CI: 65–97%) for magnetic resonance imaging in the diagnosis of visceral pleural surface invasion on length of contact (A), angle of mass margin (B) and arch distance to maximum tumor diameter ratio (C), respectively. CI = confidence interval, CT = computed tomography, VIBE = volumetric interpolated breath-hold examination.
invasion is important [6]. To predict pleural invasion, several radiological tools have been used in recent years. Chest radiography, which is the most common modality for the initial investigation of lung disease, is unsuitable for the assessment of pleural invasion [17]. Although CT is widely used for staging lung cancers, its ability to predict pleural invasion is limited [2]. Dynamic free-breathing steady-state free precession MRI has been used to assess the movement of a tumor abutting the chest wall during breathing with 88.5% accuracy [18]. The whole-lesion histogram analysis of the apparent diffusion coefficient could assist in the assessment of pleural invasion [19]. However, to the best of our knowledge, no previous studies have distinguished PL1 from PL2 pleural invasion. Our results showed that the interface between the tumor and the pleura is a useful MRI marker for the determination of VPSI with 91% accuracy. CT and MRI were equivalent in terms of the other parameters examined. There were no significant differences in the areas under the ROC curve between CT and MRI for the contact length, angle of mass margin, or arch distance-to-maximum tumor diameter ratio.

Elastic fiber staining is an important and widely accepted pathological technique for the detection and assessment of pleural invasion by lung cancer [15]. However, a previous study has highlighted the difficulties and even the impracticality of clearly classifying the depth of tumor invasion on histological slices when relying on identification of small and sometimes barely visible structures, such as the lamina elastica interna [20]. Elastic stains may be helpful for identifying the visceral pleural surface in cases of adhered visceral and parietal pleurae [16]. Although the application of elastic stains is simple and inexpensive, interpretation of the histologic results can be challenging [21]. This study showed MRI to be a potential

Fig. 5. A 61-year-old male with right lower lobe lung adenocarcinoma and no visceral pleural surface invasion (PL0, as determined pathologically).
A. Contrast-enhanced axial CT shows an ambiguous, blurred interface between the tumor and the pleura (arrow) highlighting the difficulty of evaluation of pleural invasion by CT. B-D. (B) Post-contrast free-breathing radial volumetric interpolated fat-saturated image shows a smooth and clear margin, with well-defined curvilinear pleural enhancement (arrows), and absence of protrusion at the interface between the tumor and the pleura in the axial, (C) coronal, and (D) sagittal planes. CT = computed tomography
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non-invasive, non-radiative adjunct for this assessment. In this study, the free-breathing radial VIBE sequence depicted a high-signal-intensity band at the interface between the tumor and the pleura. The smoothness or irregularity of this interface distinguished tumors without and with VPSI with substantial accuracy (Figs. 3, 5, 6). This high-signal-intensity band has been shown to correspond pathologically to edematous, thickened, adhered pleura.

Fig. 6. A 75-year-old male with left upper lobe lung adenocarcinoma and pathologically determined visceral pleural surface invasion (PL2, as determined pathologically).
A. Axial contrast-enhanced CT does not clearly show the interface between tumor and pleura (arrows), highlighting the difficulty in evaluation of pleural invasion by CT. B. Post-contrast radial volumetric interpolated breath-hold examination fat-saturated image shows an irregular undulating margin (arrows) of the tumor interface with the pleura, closely abutting the intercostal muscle, and irregular thickening of the enhanced pleura adjacent to the tumor. CT = computed tomography.

Fig. 7. A 64-year-old male with right upper lobe adenocarcinoma and no visceral pleural surface invasion (PL1, as determined pathologically).
A-C. In the non-contrast breath-hold T1-weighted (A), breath-hold T2-weighted (B), and axial fat-saturated T1-weighted breath-hold (C) images, the tumor exhibits a broad interface with the pleura (arrows) but no specific features for pleural invasion. D, E. In axial T2-weighted half-Fourier acquisition single-shot turbo spin-echo (D) and axial contrast-enhanced T1-weighted fat-suppressed images (E), the interface between the tumor and the pleura is blurred (arrows); however, there is no obvious morphologic sign enabling distinction between visceral pleural surface invasion and the lack thereof. F. On contrast-enhanced free-breathing radial volumetric interpolated breath-hold examination imaging, there is a smooth interface of the tumor with the pleura and well-defined, smooth, curvilinear pleural enhancement along the tumor (arrow).
with no parietal pleural involvement of the tumor [22].

Difficulties in distinguishing tumor invasion from non-malignant, inflammatory adhesions to the pleura and chest wall have been encountered with other diagnostic approaches also, including expiratory dynamic CT [13,23], ultrasound [24], pneumothorax CT [25], and dynamic cine MRI [18,26]. Therefore, this limitation is considered a common problem. This type of inflammatory adhesion was found on pathological examination in the two patients with false-positive MRI findings. MRI with non-contrast T1- and T2-weighted and post-contrast pulse sequences utilized to evaluate these two patients showed an indistinct margin between the tumor and the pleura, leading to the false-positive VPSI classification (Fig. 7).

Several other limitations should be mentioned. First, our study was retrospective in design. The diagnostic performances of other MRI pulse sequences could not be evaluated because only two pulse sequences were commonly shared among all MRI examinations. The assessment of the ability of other MRI pulse sequences to identify VPSI would be of value. Second, despite the enrollment of patients from two medical institutions, the number of patients was small, with three patients having PL2 and nine patients having PL3. This was probably because pathological proof was available only for patients who underwent an operation for a potentially resectable lung cancer. Therefore, patient selection bias was inevitable in this study. Further prospective multi-institutional studies with a larger sample size are required.

In conclusion, the diagnostic performance of contrast-enhanced radial T1-weighted gradient-echo 3T MRI and CT was equal in terms of the contact length, angle of mass margin, and arch distance-to-maximum tumor diameter ratio. The advantage of MRI is that it can clearly show the tumor-pleura interface margin, facilitating VPSI detection. Further studies with larger sample sizes are required to verify the clinical utility of MRI for prognostic use in NSCLC patients.

Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

Author Contributions

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