

Detectability of Target Lesion During CT-Guided Tumor Ablations: Impact on Ablation Outcome

Nachweisbarkeit der Zielläsion bei CT-gesteuerten Tumorablationen: Auswirkungen auf das Ablationsergebnis

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Key words

microwave ablation, radiofrequency (RF) ablation, liver metastases, image guidance

received 18.04.2021

accepted 07.10.2021

published online 18.11.2021

Bibliography

Fortschr Röntgenstr 2022; 194: 515–520

DOI 10.1055/a-1669-9342

ISSN 1438-9029

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Georg Thieme Verlag KG, Rüdigerstraße 14, 70469 Stuttgart, Germany

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ABSTRACT

Purpose Small hepatic malignancies scheduled for CT-guided percutaneous ablation may have been identified in the hepatobiliary phase of liver MRI or in a specific phase of multi-phase CT but may be occult on unenhanced CT used to guide the ablation. We investigated whether the detectability of the target lesion would impact the efficacy of CT-guided hepatic tumor ablations.

Materials and Methods We included 69 patients with 99 malignant liver lesions (25 primary, 44 metastases) who underwent IRE (n = 35), RFA (n = 41), or MWA (n = 23) between 01/2015 and 06/2018. All procedures were performed under CT guidance. Lesions not detectable on CT (NDL) were targeted through identification of anatomical

landmarks on preinterventional contrast-enhanced CT or MRI. Rates of incomplete ablation, size of ablation zone, local tumor recurrence, intrahepatic progression-free survival (ihPFS), and adverse event rates were compared for detectable lesions (DL) vs. NDL.

Results 40 lesions were NDL, and 59 lesions were DL on unenhanced CT. The mean follow-up was 16.2 months (14.8 for DL and 18.2 for NDL). The mean diameter of NDL and DL was similar (12.9 mm vs. 14.9 mm). The mean ablation zone size was similar (37.1 mm vs. 38.8 mm). Incomplete ablation did not differ between NDL vs. DL (5.0% [2/40; 0.6–16.9%] vs. 3.4% [2/59; 0.4–11.7%]), nor did local tumor recurrence (15.4% [6/39; 5.7–30.5%] vs. 16.9% [10/59; 8.4–29.0%]), or median ihPFS (15.5 months vs. 14.3 months).

Conclusion Target lesion detectability on interventional CT does not have a significant impact on outcome after percutaneous liver ablation when anatomical landmarks are used to guide needle placement.

Key Points:

- Liver tumors can be successfully ablated even if they are not detectable on the navigational CT scan.
- Anatomical landmarks should be used and compared to preinterventional imaging.

Citation Format

- Barzakova E, Senthilvel N, Bruners P et al. Detectability of Target Lesion During CT-Guided Tumor Ablations: Impact on Ablation Outcome. Fortschr Röntgenstr 2022; 194: 515–520

ZUSAMMENFASSUNG

Ziel Intrahepatische maligne Läsionen, geplant für CT-gesteuerte perkutane Ablation, können MR-tomographisch oder CT-graphisch in einer bestimmten Kontrastmittelphase abgrenzbar sein, jedoch okkult in der nativen CT-Untersuchung, genutzt zur Planung der Ablation. Ziel dieser Studie ist zu untersuchen, ob die Abgrenzbarkeit der Zielläsion in der Planungsuntersuchung einen Einfluss auf das kurz- und mittelfristige Ablationsergebnis hat.

Material und Methoden Es wurden 69 Patienten mit insgesamt 99 malignen Leberläsionen (25 Primärtumoren, 44 Metastasen) eingeschlossen, die zwischen 01/2015 und 06/2018

irreversibler Elektroporation (n = 35), Radiofrequenzablation (n = 41) oder Mikrowellenablation (n = 23) unterzogen wurden. Alle Verfahren wurden unter CT-Führung durchgeführt. Läsionen, die in der Planungs-CT nicht nachweisbar waren (NDL), wurden durch die Korrelation zu anatomischen Landmarken in der kontrastverstärkten CT oder MRT-Voruntersuchung lokalisiert. Es wurden die Raten unvollständiger Ablation, lokalen Tumorrezidivs, die Größe der Ablationszone, das intrahepatische progresssfreie Überleben (ihPFS) und die Komplikationsraten für nachweisbare Läsionen (DL) gegenüber NDL verglichen.

Ergebnisse 40 Läsionen waren NDL, 59 Läsionen waren DL auf die native Planungs-CT. Die mittlere Beobachtungsdauer war 16,2 Monate (14,8 für DL und 18,2 für NDL). Die mittleren Durchmesser von NDL und DL waren ähnlich (12,9 mm vs. 14,9 mm). Die mittleren Ablationszonengrößen waren ähnlich (37,1 mm vs. 38,8 mm). Es zeigten sich keine wesent-

lichen Unterschiede zwischen NDL und DL bezüglich der Häufigkeit unvollständiger Ablationen (5,0 % [2/40; 0,6 %–16,9 %] gegenüber 3,4 % [2/59; 0,4 %–11,7 %]), des Auftretens eines lokalen Tumorrezidivs (15,4 % [6/39; 5,7 %–30,5 %] gegenüber 16,9 % [10/59; 8,4 %–29,0 %]) oder des medianen ihPFS (15,5 Monate gegenüber 14,3 Monate).

Schlussfolgerung Die Nachweisbarkeit der Zielläsion in dem periinterventionellen Planungs-CT hat keinen signifikanten Einfluss auf das Ergebnis nach perkutanen Leberablationen, wenn anatomische Landmarken zur Nadelplatzierung genutzt werden.

Kernaussagen:

- Lebertumoren können erfolgreich ablatiert werden, auch wenn sie in dem nativen Planungs-CT nicht nachweisbar sind.
- Anatomische Landmarken sollten verwendet und mit der präinterventionellen Bildgebung verglichen werden.

Introduction

Percutaneous ablation, e. g., microwave (MWA), radiofrequency ablation (RFA), or irreversible electroporation (IRE), is a well-established option for the treatment of selected hepatic tumors [1–6]. Successful ablation needs to cover the whole lesion including a sufficient safety margin of at least 5–10 mm beyond the tumor margin [7–9], while critical anatomic structures need to be spared. High-quality imaging is an important prerequisite to determine the ablation method of choice [10]. However, some tumors might be challenging to treat because they are associated with transient enhancement on multiphase CT, or are only visible on MRI, or even only on specific MRI sequences such as diffusion-weighted imaging, or hepatobiliary-phase imaging after liver-specific contrast.

On the other hand, image guidance during ablation is typically achieved by means of unenhanced CT and/or ultrasound. Different methods are used to improve the detectability of the target lesion during CT-guided ablation in order to overcome this shortcoming [11]. One approach to directly improve lesion visibility is to inject small amounts of contrast agent through an intra-arterial catheter, placed in the hepatic artery [12] – a method that increases the invasiveness of the procedure, requires access to DSA to place the catheter, and works only for lesions with a strong arterial blood supply. Indirect approaches use image co-registration of diagnostic CECT or MR imaging with the corresponding intraprocedural imaging [13–15]. However, co-registration of hepatic lesions can be error-prone.

Another, simple method is to use surrounding anatomical landmarks that can usually be defined on unenhanced CT, like portal or liver vein branches, or distance to liver capsule or other anatomical structures [16]. However, such procedures can be challenging since not all landmarks are visible on unenhanced CT. Image quality is often worse on intra-procedural control scans due to beam-hardening artifacts. Moreover, gross patient motion, patient breathing, and bowel motion, as well as deformation of

the liver secondary to the penetration of the probe can all contribute to difficulties when using this approach.

Accordingly, the aim of this study was to investigate whether the detectability of the target lesions in unenhanced CT would affect the outcome of liver ablation procedures done on the basis of unenhanced CT, using the anatomical landmark approach to identify target lesions.

Materials and Methods

This retrospective study was in accordance with the ethical standards of the institutional research committee and with the 1964 Helsinki declaration and its later amendments.

Patients

We included all patients who underwent CT-guided hepatic ablation procedures (RFA, MWA, or IRE) in our department between 06/2015 and 06/2018. The decision to perform percutaneous ablation was established in consensus in a multidisciplinary tumor board, attended by hepatobiliary surgeons, oncologists, radiotherapists, pathologists, radiologists, and interventional radiologists. Hepatic tumor ablation was recommended in patients, who were considered illegible for surgical resection and had no extrahepatic tumor burden.

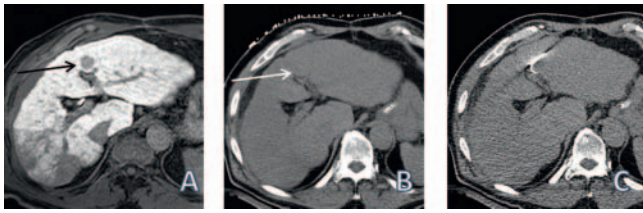
All patients had undergone either contrast-enhanced CT or MRI prior to ablation.

Patients, who had less than four weeks of follow-up, were excluded from the study.

Ablation

All procedures were performed by experienced interventional radiologists with more than 5 years of experience in abdominal interventions.

As per the standard protocol at our institution, all procedures were performed under general anesthesia. The individual



► **Fig. 1** Using anatomical landmarks for needle placement. **A** diagnostic, diffusion-weighted MR image. **B** unenhanced CT image with surface marker grid for planning needle placement (the known lesion is not visible, but the adjacent portal vein branches are well depictable). **C** RFA applicator in the right position.

► **Abb. 1** Nutzung anatomischer Landmarken für Nadelpositionierung.

approach and patient positioning during the ablation were determined by the interventionalist. For planning applicator placement, unenhanced CT was performed and compared to previous images using the anatomical landmark method (► **Fig. 1**). Lesions were defined as visible lesions (DLs) or non-visible lesions (NDLs) if the lesions could or could not be detected by an experienced interventional radiologist. The applicator was positioned using intermittent CT scans in expiratory breath hold.

If needed, overlapping ablation after repositioning of the electrodes was performed until the interventional radiologist felt that tumor ablation should be complete including a safety margin of at least 5–10 mm. The safety margin and the size of the ablation zone were planned according to the size and location of the lesion, independent of its detectability.

Before removing the electrode, a contrast-enhanced tri-phasic CT scan was obtained to verify treatment success and assess immediate complications. Track ablation was performed in all RFA and MWA procedures.

For RFA we used a monopolar system (RF 3000, Boston Scientific Corp, MA, USA) with umbrella-shaped applicators with a diameter of 2–4 cm and varying shaft lengths (LeVein, Boston Scientific Corp, MA, USA). Ablation is performed according to the vendor's ablation protocols.

MWA was performed using the Emprint system (Medtronic, MI, USA) with dedicated antennae with different shaft lengths. We always adhered to the protocols provided by the vendor.

We used the NanoKnife® system (AngioDynamics, Amsterdam, the Netherlands) for IRE with 2 to 5 probes with a tip exposure of 1.5 cm. IRE was performed with 70–90 pulses per probe pair, a pulse length of 90 µs, and a maximum voltage of 3000 V.

Evaluation of ablation and follow-up

The size of the ablation zone was measured in the immediate postinterventional contrast-enhanced CT examination, calculating the average diameter ((short + long axis)/2).

Adverse events were assessed according to the CIRSE classification system [17].

Routine follow-up imaging was performed 1 month after ablation and every 3 months thereafter and consisted of either contrast-enhanced multiphase CT or contrast-enhanced MRI. Technical efficacy was determined on the basis of the 1-month CT/MRI

examination according to the criteria defined by the international working group on image-guided tumor ablation [8].

Incomplete ablation (IA) was defined as the persistence of enhancing areas within the ablation zone or peripheral nodular enhancement within a 10-mm margin at the first follow-up scan four weeks after the procedure.

Local tumor progression (LTP) was defined as recurrent disease within a 10-mm margin during subsequent follow-up.

Intrahepatic progression-free survival (ihPFS) time was defined as the interval between ablation procedure and the first detection of local recurrence or new metastases elsewhere in the liver.

Statistical analysis

All statistical analyses were performed using SPSS software (version 24; IBM). IA and LTP rates as well as frequency of adverse events (AE) were compared between the groups using the Chi-square test. The lesion size and the size of the ablation zone were compared between the groups using the Mann-Whitney U-test. ihPFS was calculated for all patients using the Kaplan-Meier method and the difference among groups was determined by the log-rank test.

A value of $p < 0.05$ was considered statistically significant. All values are expressed as mean ± standard deviation (SD) or median and interquartile range (IQR).

Results

Patients and tumors

A total of 69 patients with 99 malignant hepatic liver lesions were included in the study. Patient characteristics are summarized in ► **Table 1**. The median number of treated tumors per patient was 1 (range 1–3). 59 of all treated lesions were DLs in the planning unenhanced CT examination and 40 were NDLs.

The mean lesion size showed no statistically significant difference between lesions that were visible and those that were not visible, with 14.9 ± 0.9 mm for DLs and 12.9 ± 0.8 mm for NDLs ($p = 0.28$).

The type of ablation procedure used is summarized in ► **Table 2**.

Technical success

The size of the ablation zone was not significantly different for both groups: 38.8 ± 1.3 mm for DLs and 37.1 ± 1.4 mm for NDLs ($p = 0.41$).

Incomplete ablation was observed in 2 out of 59 cases (3.4%; 0.4–11.7%) in the DL group and in 2 out of 40 cases (5.0%; 0.6–16.9%) in the NDL group, which was also not significantly different ($p = 0.69$).

Adverse events

AEs were observed in 6 of 69 patients (9%).

The following AEs were observed: One patient developed pneumothorax, two patients had perihepatic hematomas, two patients had arterioportal fistulae, and one patient was found to have segmental liver infarction due to procedure-related portal vein thrombosis. In two patients, an additional intervention in

► **Table 1** Patients characteristics.► **Tab. 1** Patientencharakteristika.

		N = 69
Age (y)	66.6 (± 14.8)	
Gender	Male	52 (75 %)
	Female	17 (25 %)
Tumor type	Hepatocellular cancer	20 (29 %)
	Cholangiocarcinoma	9 (13 %)
	Colorectal metastases	33 (48 %)
	Metastases of melanoma	2 (3 %)
	Metastases of ovarian cancer	2 (3 %)
	Metastases of gastric carcinoma	1 (1 %)
	Metastases of pancreas carcinoma	1 (1 %)
	Metastases of bronchial carcinoma	1 (1 %)
	Metastases of breast cancer	1 (1 %)
	Liver cirrhosis	Yes
No		55 (80 %)
Number of treated lesions per patient	n = 1	47 (68 %)
	n = 2	15 (22 %)
	n = 3	6 (9 %)
	n = 4	1 (1 %)

form of coil embolization (one of arterial bleeding and one of a symptomatic arterioportal fistula) was required (grade 3).

All other complications were categorized as grade 1. There was no procedure-related permanent morbidity or death. All complications were observed in the DL group in 6 out of 59 cases (10%), with a statistically significant difference ($p = 0.037$) compared to 0 out of 40 cases (0%) in the NDL group.

Follow-up, LTP, and ihPFS

The mean follow-up was 16.2 ± 11.7 months: 14.8 ± 1.5 months for patients after ablation of DLs and 18.2 ± 1.8 months for patients after ablation of NDLs.

One patient was lost to follow-up after 4 weeks.

During the follow-up-time, local tumor progression was observed in 10 out of 59 cases (16.9%) in the DL group and 6 out of 39 cases (15.4%) in the NDL group, which was not significantly different ($p = 0.84$).

Using Kaplan-Meier analysis, the mean estimated ihPFS time was 15.5 ± 2.2 months for DLs compared to 14.3 ± 2.4 months for NDLs ($p = 0.84$). These results are visualized in ► **Fig. 2**.

Discussion

This study investigated whether the detectability of target lesions during CT-guided hepatic ablations would influence the technical and oncological success or adverse event rates of ablation proce-

► **Table 2** Ablation characteristics.► **Tab. 2** Charakteristika der Ablationen.

		N = 99	
Detectability of tumor in unenhanced CT	Detectable (DL)	59 (59 %)	
	Non-detectable (NDL)	40 (40 %)	
Ablation type	MWA	DL n = 18	23 (23 %)
		NDL n = 5	
	RFA	DL n = 24	41 (41 %)
		NDL n = 17	
	IRE	DL n = 17	35 (35 %)
		NDL n = 28	

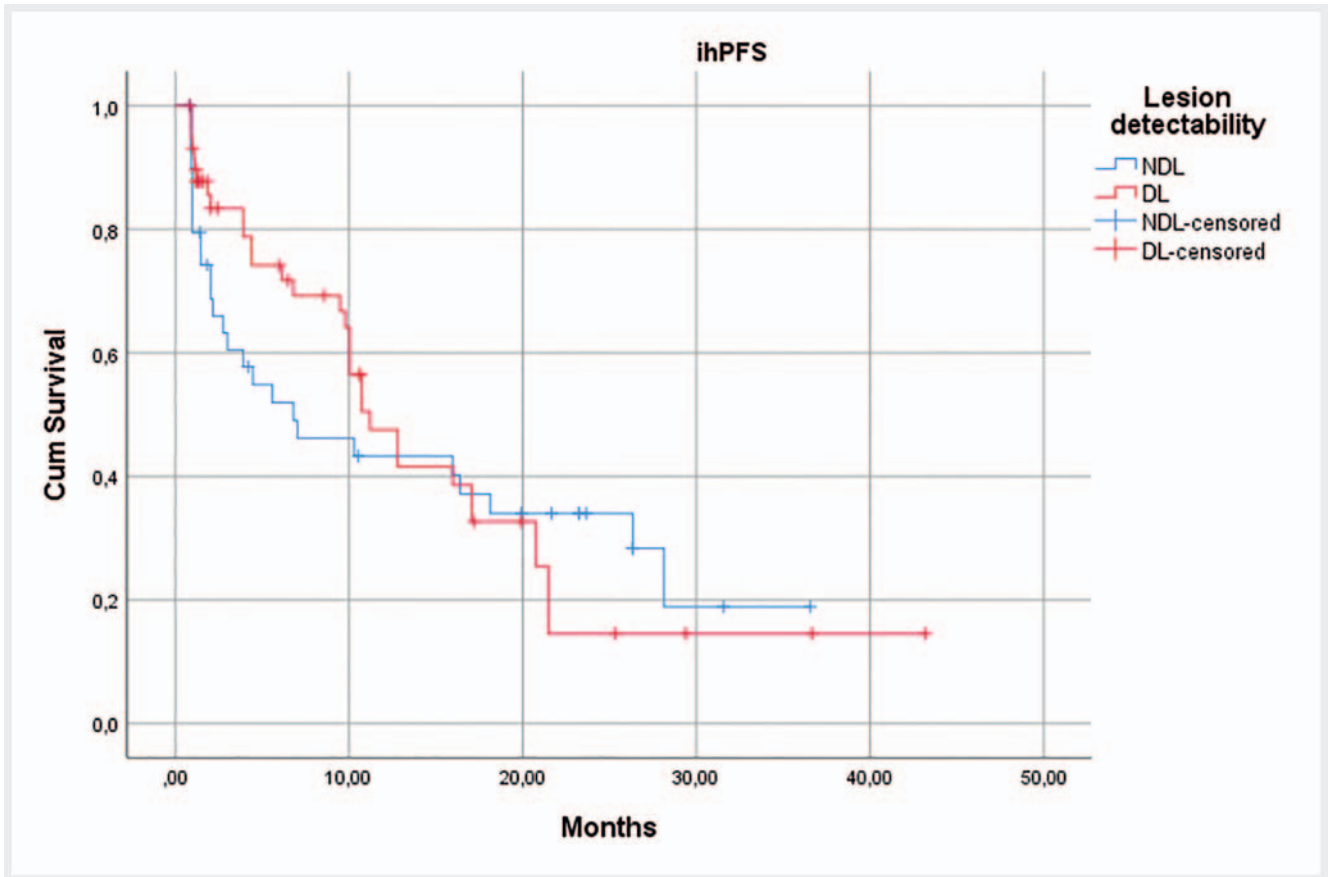
dures. Our results demonstrated that there was no significant difference comparing the groups with detectable and non-detectable lesions in terms of the completeness of ablation, rates of local recurrence, or intrahepatic recurrence-free survival. We expected that in patients with non-detectable lesions (NDL), the ablation zones would be larger than in patients with detectable lesions (DL) – but this was also not the case. Last, adverse events were even significantly lower in the NDL group, suggesting that they are not connected with the detectability of the target lesion.

Accordingly, when using the “anatomical landmark approach” to ablate non-detectable lesions, the outcome was similar compared with that of patients with detectable lesions. Applying the “anatomical landmark approach” it should be considered that the position of the target lesion in relation to a landmark might be distorted due to patient positioning (e. g., supine vs. prone position) or breathing movements. Therefore, we perform every needle placement and every control CT scan in breath hold under general anesthesia and we attempt to perform ablation in the same patient position as in the preoperative imaging.

The overall rate of local tumor recurrence observed in this study is in line with previously published data, ranging from 6% to 33% [18–22]. ► **Fig. 3** shows an example of an incomplete ablation in one of our patients, most likely due to the adjacency of the liver vein and thus the heat sink effect. Several studies investigated risk factors associated with partial ablation and local recurrence [23–26]. To our knowledge, there is no previous study investigating the detectability of the target lesion as a potential risk factor.

One study that compared the accuracy of computer-assisted versus cognitive registration for locating liver tumors had shown that non-rigid registration yielded better localization accuracy than cognitive registration performed by the interventional radiologist [27]. However, no clinical outcome was investigated in this study. In other words, the clinical benefit of image registration in liver ablation is largely unknown.

Limiting factors of the presented study include the use of unenhanced CT scans as a criterion for lesion visibility and the lack of additional ultrasound guidance. We generally use unenhanced CT for ablation planning but use contrast agent after



► **Fig. 2** Kaplan-Meier curve for intrahepatic progression-free survival.

► **Abb. 2** Kaplan-Meier-Kurve für das intrahepatische progresssfreie Überleben.



► **Fig. 3** Example of an incomplete ablation. **A** Pre-procedural T2-weighted MRI-Image of a colorectal metastasis adjacent to the middle liver vein (arrows). **B** Post-interventional control-CT-scan in hepatic portal phase one day after MWA of the metastasis. **C** T2-weighted MRI-Image four weeks after MWA with detectable residual tumor adjacent to the middle liver vein, in the sense of incomplete ablation, probably due to the heat sink effect.

► **Abb. 3** Beispiel einer unvollständigen Ablation. **A** Preinterventionelles, T2-gewichtetes MRT-Bild einer kolorektalen Metastase angrenzend an die mittlere Lebervene (Pfeile). **B** Postinterventionelle Kontroll-CT-Untersuchung in venöser Kontrastmittelfase ein Tag nach Mikrowellenablation der Metastase. **C** T2-gewichtetes MRT-Bild vier Wochen nach Ablation mit abgrenzbarem Resttumor angrenzend an die mittlere Lebervene, im Sinne einer unvollständigen Ablation, am ehesten aufgrund von heat sink Effekt.

the presumed completion of the procedure in order to evaluate the ablation result. We perform contrast-enhanced imaging only in cases requiring visualization of not only the target lesion but

also of critical anatomic structures e. g., the liver hilum – but usually after placement of the ablation probe near the target lesion. Often several rounds of contrast-enhanced CT imaging are required, though the reasonably applied amount of IV contrast is limited. Therefore, we tend to dispense with an initial contrast-enhanced CT examination for planning purposes in order to safely be able to perform a contrast-enhanced CT examination in the case of acute complications. Also, anatomical structures, as well as the target lesion, are visible only for a short time after application of contrast agent. Nevertheless, their exact position might change during the procedure as they can be displaced by the puncture needle.

Ultrasound is often used as a stand-alone or assisting technique for tumor ablation. In our institution we perform tumor ablation solely with CT guidance. The addition of US could help depict more NDLs and could be used as a real-time tool for accurate needle placement.

In view of our results, however, there seems to be little need for additional ultrasound for targeting purposes even in patients with NDLs.

Another often used technique is prior lesion marking by intraarterial lipiodol injection. In our experience marking was insufficient in many cases of liver metastases due to poor vascularization. Furthermore, the combination of intraarterial and CT-

guided interventions is often logistically challenging and places an additional strain on the patient with a further risk of complications. Nevertheless, our experience is based on very few cases and since there are some good results in the published literature [28], this technique should be considered in the appropriate cases.

We included a variety of liver malignancies, including primary and secondary liver tumors, which typically present different features on CT and MRI imaging, such as arterial hyperenhancement and venous washout. Also, the presence or absence of liver cirrhosis generates a different image in diagnostic and interventional imaging. However, the different tumor types and the cases of liver cirrhosis were evenly distributed in the DL and NDL groups, so that we are confident that this would not confound our results.

In conclusion, this study demonstrates that successful ablation of liver tumors that are not detectable on unenhanced CT is possible and safe when using the anatomical landmark method to guide needle placement.

Conflict of Interest

The authors declare that they have no conflict of interest.

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