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IAMCOMPLETE study**

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# Intraprocedural assessment of ablation margins using computed tomography co-registration in hepatocellular carcinoma treatment with percutaneous ablation: IAMCOMPLETE study



Pim Hendriks<sup>a,\*</sup>, Kiki M van Dijk<sup>a</sup>, Bas Boekestijn<sup>a</sup>, Alexander Broersen<sup>b</sup>,  
 Jacoba J van Duijn-de Vreugd<sup>a</sup>, Minneke J Coenraad<sup>c</sup>, Maarten E Tushuizen<sup>c</sup>,  
 Arian R van Erkel<sup>a</sup>, Rutger W van der Meer<sup>a</sup>, Catharina SP van Rijswijk<sup>a</sup>, Jouke Dijkstra<sup>b</sup>,  
 Lioe-Fee de Geus-Oei<sup>a,d,e</sup>, Mark C Burgmans<sup>a</sup>

<sup>a</sup> Department of Radiology, Leiden University Medical Center, 2333 ZA, Leiden, the Netherlands

<sup>b</sup> LKEB Laboratory of Clinical and Experimental Imaging, Department of Radiology, Leiden University Medical Center, 2333 ZA, Leiden, the Netherlands

<sup>c</sup> Department of Gastroenterology and Hepatology, Leiden University Medical Center, 2333 ZA Leiden, the Netherlands

<sup>d</sup> Biomedical Photonic Imaging Group, TechMed Centre, University of Twente, 7522 NB, Enschede, the Netherlands

<sup>e</sup> Department of Radiation Science & Technology, Delft University of Technology, 2628 CD, Delft, the Netherlands

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## ABSTRACT

**Purpose:** The primary objective of this study was to determine the feasibility of ablation margin quantification using a standardized scanning protocol during thermal ablation (TA) of hepatocellular carcinoma (HCC), and a rigid registration algorithm. Secondary objectives were to determine the inter- and intra-observer variability of tumor segmentation and quantification of the minimal ablation margin (MAM).

**Materials and methods:** Twenty patients who underwent thermal ablation for HCC were included. There were thirteen men and seven women with a mean age of  $67.1 \pm 10.8$  (standard deviation [SD]) years (age range: 49.1–81.1 years). All patients underwent contrast-enhanced computed tomography examination under general anesthesia directly before and after TA, with preoxygenated breath hold. Contrast-enhanced computed tomography examinations were analyzed by radiologists using rigid registration software. Registration was deemed feasible when accurate rigid co-registration could be obtained. Inter- and intra-observer rates of tumor segmentation and MAM quantification were calculated. MAM values were correlated with local tumor progression (LTP) after one year of follow-up.

**Results:** Co-registration of pre- and post-ablation images was feasible in 16 out of 20 patients (80%) and 26 out of 31 tumors (84%). Mean Dice similarity coefficient for inter- and intra-observer variability of tumor segmentation were 0.815 and 0.830, respectively. Mean MAM was  $0.63 \pm 3.589$  (SD) mm (range: -6.26–6.65 mm). LTP occurred in four out of 20 patients (20%). The mean MAM value for patients who developed LTP was -4.00 mm, as compared to 0.727 mm for patients who did not develop LTP.

**Conclusion:** Ablation margin quantification is feasible using a standardized contrast-enhanced computed tomography protocol. Interpretation of MAM was hampered by the occurrence of tissue shrinkage during TA. Further validation in a larger cohort should lead to meaningful cut-off values for technical success of TA.

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**Abbreviations:** BCLC, Barcelona Clinic Liver Cancer; CECT, Contrast-enhanced computed tomography; CT, Computed tomography; CTCAE, Common terminology criteria for adverse events; DoS, Degree of similarity; EASL, European Association for the Study of the Liver; eGFR, Estimated glomerular filtration rate; HCC, Hepatocellular carcinoma; ICC, Intraclass correlation coefficient; LI-RADS, Liver Imaging Reporting and Data System; LTP, Local tumor progression; MAM, Minimal ablation margin; MRI, Magnetic resonance imaging; MWA, Microwave ablation; RFA, Radiofrequency ablation; SD, Standard deviation; TA, Thermal ablation; vDSC, Volumetric Dice similarity coefficient

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\* Corresponding author.

E-mail address: [p.hendriks@lumc.nl](mailto:p.hendriks@lumc.nl) (P. Hendriks).

## 1. Introduction

Hepatocellular carcinoma (HCC) represents the sixth most common malignancy worldwide and the fourth most common cause of cancer-related mortality. It occurs predominantly in patients with chronic liver disease, in particular cirrhosis [1]. Thermal ablation (TA) is an effective treatment for small HCC lesions and considered the preferred modality for very early stage HCC according to the Barcelona Clinic for Liver Cancer (BCLC) guidelines [2]. For solitary lesions  $\leq 2$  cm, the effectiveness is equal to that of surgery but at a lower risk of

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complications [3–6]. For lesions > 2 cm surgical resection remains the preferred treatment as local tumor progression (LTP) is more prevalent after TA with relative risk ratios of up to 1.42 [7].

The most common causes of LTP are insufficient heat propagation resulting in remaining viable tumor tissue at the peripheral parts of the tumor, heat sink due to bordering intrahepatic blood vessels, and the presence of satellite nodules [8]. In order to minimize the risk of LTP after TA, it is generally recommended to achieve a minimal ablation margin (MAM) ≥ 5 mm encompassing the tumor [9,10]. After TA, contrast-enhanced computed tomography (CECT) examination is made to determine whether complete tumor necrosis with sufficient MAM has been achieved. The evaluation is usually based on a qualitative visual interpretation of side-to-side positioned diagnostic imaging and post-ablation CECT, assisted by two-dimensional in-plane measurements using anatomical landmarks.

In an attempt to identify patients at risk of developing LTP, software-assisted ablation margin assessment is considered as a promising tool [11–17]. Such software tool objectifies the MAM by using co-registration of diagnostic and post-ablation imaging. By segmenting the tumor on the diagnostic scan and the ablation zone on post-ablation CECT, a three-dimensional volumetric analysis can be performed to determine the MAM after coregistration of the images. Multiple retrospective studies found a correlation between software-assisted ablation margins and the occurrence of LTP [11–14].

Although software-assisted MAM determination is promising, no standardized or widely validated workflow has been described yet. Co-registration of preprocedural diagnostic images with a post-ablation CECT may result in registration errors as a result of differences in patient positioning, inhalation mode and imaging modality [11]. Moreover, tumor growth in the time interval between diagnostic imaging and TA may cause overestimation of the MAM. Lastly, little is known about the robustness of the co-registration process. Dissimilarities between different operators may occur while segmenting or registering medical images. We hypothesized that standardized acquisition of pre- and post-ablation images would contribute to the efficacy of software-assisted ablation margin quantification.

The primary objective of this study was to investigate the feasibility of ablation margin quantification using a standardized CECT protocol and a rigid registration algorithm. Secondary objectives were to investigate inter- and intra-observer variability, the time needed for CT-CT co-registration, and the correlation between ablation margins and local recurrence.

## 2. Materials and methods

The IAMCOMPLETE study was a prospective, single-arm, single center phase II feasibility study. The study protocol was approved by the institutional medical ethical board, and informed consent was obtained from all patients. The study was registered on ClinicalTrials.gov with number NCT04123340.

### 2.1. Patients

Twenty patients who underwent TA for HCC were recruited for this pilot study. Eligibility for TA and study participation of all patients were discussed in a multidisciplinary tumor board meeting. Inclusion criteria were: (i), (very) early stage HCC according to the BCLC staging system (solitary lesion ≤ 5 cm or a maximum of 3 lesions of ≤ 3 cm each); (ii), general eligibility for percutaneous TA; and (iii), age ≥ 18 years. HCC was diagnosed according to European Association for the Study of the Liver (EASL) guidelines based on histology or the Liver Imaging Reporting and Data System (LI-RADS) imaging characteristics on CT or magnetic resonance imaging (MRI) [18,19]. Exclusion criteria were: (i), estimated glomerular filtration rate (eGFR) < 30 mL/min/1.73 m<sup>2</sup>; (ii), morbid obesity (body mass index ≥ 40); (iii), any pulmonary condition that would

be a contraindication to prolonged apnea; (iv), Child Pugh C liver cirrhosis; (v), extrahepatic metastasis; and (vi), uncorrectable coagulopathy. Baseline patient and tumor characteristics were pseudonymized and stored in an encrypted database.

### 2.2. Thermal ablation procedure

TA was performed using either radiofrequency ablation (RFA) or microwave ablation (MWA). RFA was performed using Medtronic Cool-tip (single 3 or 4 cm needles, 3 × 3 cm or 3 × 4 cm needles) with up to six needle positions and MWA using Medtronic Emprint with thermosphere technology. Factors influencing the choice of TA technique and settings included tumor size, tumor location, bordering blood vessels and operator preference. Needle positioning was performed using ultrasound guidance, ultrasound-CT/MRI fusion (General Electric LOGIC E9, General Electric) or CT-guidance.

### 2.3. Scanning protocol

All TA procedures were performed under general anesthesia in a procedure room using an Aquilion One™ CT scanner (Canon Medical Systems). Prior to ablation, CECT was acquired using a patient weight dependent bolus of contrast material containing 350 mg of iodine per mL (Xenetix® 350, Guerbet). The full contrast protocol is reported in Table 1. Pre-oxygenated apnea was used while scanning by disconnection of the ventilation tube just before intravenous administration of the contrast agent. CT images were acquired using Sure Start™ bolus tracking at the descending aorta with a 10- and 50 s delay for arterial and venous phases, respectively.

Immediately after TA a second CECT examination was obtained using the same scanning protocol. This CECT examination was used by the performing interventional radiologist to determine whether technical success was achieved. The ablation was considered successful when the interventional radiologist estimated that a safety margin > 5 mm was obtained based on a qualitative interpretation of the pre- and post-TA imaging. The quantitative assessment was performed after the procedure and was not used to determine the technical success per-procedurally.

### 2.4. Follow-up

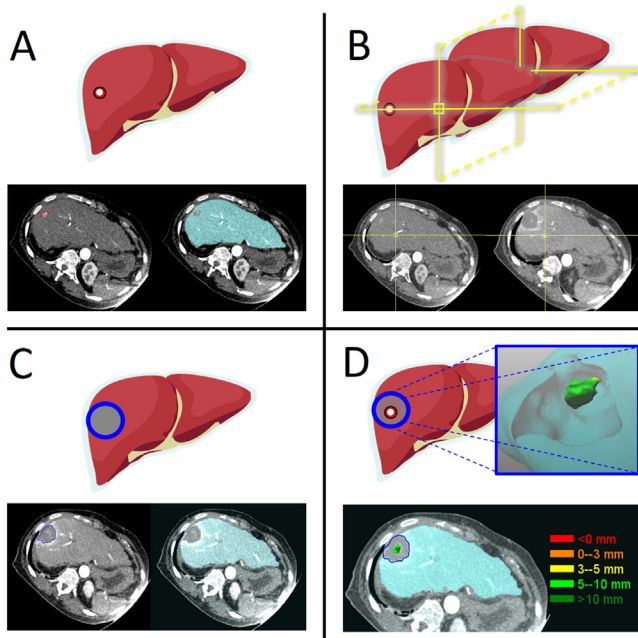
Study participants were followed for one year after TA. The follow-up schedule was similar to all regular TA patients and consisted of blood testing (including alpha-fetoprotein) at six weeks after the procedure, complemented with imaging using CECT or MRI every 3–4 months. Adverse events were graded according to the common terminology criteria for adverse events (CTCAE) 5.0 [20].

### 2.5. Software-assisted ablation margin quantification

Two teams performed the software-assisted ablation margin assessment, consisting of a researcher (PH, KMvD) and a radiologist with 8 or 20 years of experience in abdominal radiology (BB, MCB). In-house developed “deLIVERed” software was used, which uses an

**Table 1**  
Protocol of intravenous administration of iodinated contrast materials for computed tomography examination according to patient weight.

| Weight (kg) | Contrast agent |             | NaCl (0.9%) |             |
|-------------|----------------|-------------|-------------|-------------|
|             | Volume (mL)    | Flow (mL/s) | Volume (mL) | Flow (mL/s) |
| < 65        | 100            | 5.0         | 50          | 5.0         |
| 65–80       | 130            | 5.5         | 55          | 5.5         |
| 80–100      | 160            | 6.0         | 60          | 6.0         |
| > 100       | 190            | 7.0         | 70          | 7.0         |



**Fig. 1.** Software assisted ablation margin quantification. A, Step 1) semiautomatic segmentation of the liver and manual segmentation of the liver tumor. B, Step 2) semiautomatic gray-scale based rigid registration of pre- and post-ablation computed tomography images. C, Step 3) semiautomatic segmentation of the ablation zone. D, Step 4) Analysis of quantified ablation margins.

open-source Elastix-based registration algorithm, which is implemented in MeVisLab [21,22]. Axial 1-mm slices were used for all analyses.

A schematic overview of the software assisted workflow is presented in Fig. 1. In step 1) semiautomatic segmentation of the liver volume on the pre-ablation scan is performed to create a “liver mask” as region of interest for image registration. Manual tumor volume segmentation was performed by segmenting the tumor contour on every axial slice. Step 2) consisted of a semiautomatic rigid registration of pre- and post-TA CECT. This step was aided by a projection of the “liver mask” on the post-TA CECT. After initializing the starting position of the scans onto each other (*i.e.*, dragging the liver mask onto the liver in the post-TA CECT), the automatic rigid registration algorithm was used. This result could be refined with help of anatomical landmarks or manual adjustments by means of translation and rotation. Local optimized registration was strived for in the region of the tumor and TA zone. Step 3) was to semi-automatically define the “liver mask” on the post-TA CECT and to semi-automatically segment the ablation necrosis on the post-TA CECT. This was done by manual segmentation of the ablation zone on multiple slices and interpolation between these contours on intermediate slices. Step 4) was to transform the contours and registration into a three-dimensional model that was saved as a mesh structure in a Visualization Toolkit (VTK) file. This file was then quantitatively analyzed using Paraview 5.10.0 software. A lookup table was created in which the MAM (*i.e.*, the shortest distance between the tumor surface and ablation necrosis surface) was stored. The exported lookup table was used for further statistical analysis. All steps were repeated for every tumor. The segmented liver mask on post-TA CT ensured the MAM to always represent the shortest distance from pre-ablation tumor segmentation to post-TA viable liver tissue [13].

## 2.6. Image analysis

MAM quantification using the previously described workflow was performed for each tumor separately in both imaging phases. Moreover, all MAM quantifications were repeated by both teams to

determine intra-observer variability. The anatomical tumor side of the MAM was captured. A minimal time interval of two weeks between the two analyses was taken into account. The time needed to complete several steps of the workflow (*i.e.*, liver segmentation, tumor segmentation, co-registration and post-TA image analysis) was recorded. For inter-observer variability, the second analyses of both teams were used.

## 2.7. End points

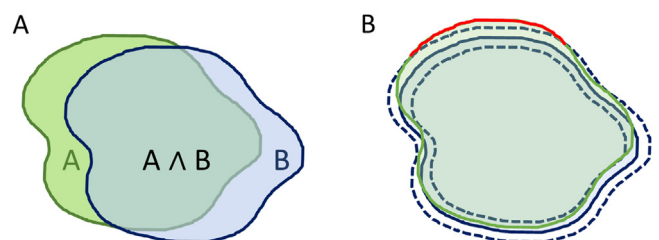
The primary endpoint of this study was met when CT-CT co-registration was feasible in  $\geq 80\%$  of all patients. This parameter was objectified using a 5-points scale. (Quality of registration: 1 = “poor”; 2 = “insufficient”; 3 = “moderate”; 4 = “good”; and 5 = “perfect”). In case of disagreement between the two teams, a consensus reading was performed. At a score of 4 or 5, the registration was considered feasible and well enough to pursue ablation margin quantification.

Secondary endpoints of this study were inter- and intra-observer agreement rates for tumor delineation and MAM measurement. The MAM was defined as the single shortest distance between the tumor volume and ablation volume. In this study the tumor volume segmented from the pre-ablation CECT and the ablation volume segmented from the post-ablation CECT were co-registered to obtain the MAM using three-dimensional, software-assisted analysis. The inter- and intra-observer agreement rates were expressed as volumetric Dice similarity coefficient (vDSC) and degree of similarity (DoS) within a two-pixel range (Fig. 2). The values from vDSC and DoS range from 0 to 1, meaning no spatial overlap and perfect agreement, respectively [23]. Values between 0.6 to 0.8 were considered to indicate substantial overlap and those between 0.8 and 1.0 to indicate almost perfect overlap, as they were derived from Kappa statistics [24]. For each lesion, a single MAM value was determined as the average value of the second measurements of both observers, on arterial phase images. For lesions that were invisible on arterial phase images, portal venous phase images were used.

All follow-up CT examinations of patients developing LTP or intra-hepatic metastases were double read by two radiologists with 8 and 20 years of experience in abdominal radiology (BB, MCB). In case of disagreement, consensus was reached on the anatomical orientation of LTP with respect to the tumor necrosis. This anatomical orientation was correlated to the anatomical orientation of the MAM.

## 2.8. Statistical analysis

Quantitative variables data were expressed as means  $\pm$  standard deviations (SD) and ranges. Categorical variables were expressed as numbers and proportions. Analyses were performed using Rstudio 1.4.1106. Inter- and intra-observer variability were analyzed using Bland Altman plots. Mean MAM-values of the two observers were



**Fig. 2.** Inter- and intra-observer variability outcome measures of tumor segmentation. A) shows the volumetric Dice similarity coefficient (vDSC), which is calculated by dividing the overlapping volume of two segmented volumes by the non-overlapping parts of these volumes. B) shows the degree of similarity (DoS) within 2 voxel distance between two segmentations. Dashed blue lines represent 2 voxel distance of the blue segmentation and the green segmentation exceeds this margin for approximately 20% at the top border (in red).



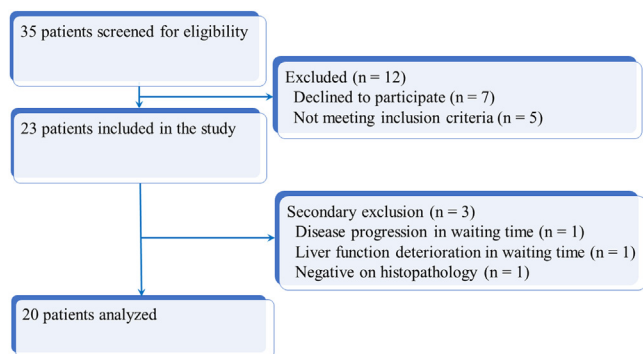


Fig. 3. Flowchart of the study population.

plotted in box-plots and expressed as medians with inter quartile ranges. The intraclass correlation coefficient (ICC) for inter-observer variability was determined [25].

### 3. Results

Informed consent was obtained from 23 patients, of whom 20 patients were treated according to the study protocol (Fig. 3). Reasons for preliminary exclusion were lack of histopathological confirmation of HCC after a combined biopsy and TA procedure for a LI-RADS-4 lesion (n = 1) [19], disease progression to Child-Pugh C liver cirrhosis (n = 1) or to intermediate HCC (n = 1) within a maximum of six weeks waiting time. Table 2 shows the baseline characteristics of the patients analyzed. A total of 20 patients with 31 HCC lesions were included with a mean lesion diameter of 18.8 ± 7.34 mm [SD] (range: 8–38 mm). Eight patients underwent prior HCC treatment before inclusion in this study.

Thirteen patients were treated with MWA and seven patients with RFA. TA was deemed successful with sufficient ablation margins in 30/31 ablations by discretion of the treating physician based on side-to-side reading of pre- and post-ablation CECT. In those patients, no additional ablation was performed. In one patient a CTCAE 5.0 grade 3 bleeding occurred that led to preliminary termination of the procedure. In this patient additional TA was performed in a second treatment. Other adverse events were CTCAE 5.0 grade 3: post-procedural pleural effusion (n = 1), grade 2: hematoma formation (n = 2), grade 2: postprocedural pain (n = 2), grade 2: postprocedural fever (n = 1), and grade 1: iatrogenic vascular thrombosis of a segmental vessel leading to segmental liver infarction (n = 1).

Fig. 4 shows two examples of MAM quantification. Table 3 shows HCC lesion visibility on the different CT imaging phases and ablation margin quantification parameters. Out of 31 lesions, 18 lesions were

Table 2  
Characteristics of 20 patients with hepatocellular carcinomas treated with thermal ablation.

| Variable              | Value                   |
|-----------------------|-------------------------|
| Age (year)            | 67.1 ± 10.8 [49.1–81.1] |
| Sex                   |                         |
| Male                  | 13 (65%)                |
| Female                | 7 (35%)                 |
| Cirrhosis             |                         |
| Yes                   | 17 (85%)                |
| No                    | 3 (15%)                 |
| Etiology of cirrhosis |                         |
| Hepatitis B           | 1 (6%)                  |
| Hepatitis C           | 1 (6%)                  |
| Alcohol abuse         | 14 (82%)                |
| NASH                  | 1 (6%)                  |
| Child-Pugh score      |                         |
| A                     | 12 (88%)                |
| B                     | 5 (12%)                 |
| BCLC stage            |                         |
| Very early            | 8 (40%)                 |
| Early                 | 10 (50%)                |
| Intermediate          | 2 (10%)                 |
| Diagnostic imaging    |                         |
| CT                    | 9 (45%)                 |
| MRI                   | 11 (55%)                |
| Prior HCC treatment   |                         |
| None                  | 11 (60%)                |
| Surgical resection    | 1 (5%)                  |
| RFA                   | 4 (20%)                 |
| TACE                  | 4 (20%)                 |
| Number of HCC lesions |                         |
| 1                     | 13 (70%)                |
| 2                     | 5 (20%)                 |
| 3                     | 1 (5%)                  |
| 4                     | 0 (0%)                  |
| 5                     | 1 (5%)                  |
| Lesion size (mm)      | 18.8 ± 7.3 [8.0–38.0]   |
| Ablation technique    |                         |
| RFA                   | 7 (35%)                 |
| MWA                   | 13 (65%)                |

BCLC = Barcelona Clinic for Liver Cancer, CT = computed tomography; HCC = hepatocellular carcinoma; MWA = microwave ablation; MRI = magnetic resonance imaging; NASH = non-alcoholic steatohepatitis; RFA = radiofrequency ablation; TACE = trans-arterial chemoembolization. Quantitative data are reported as means ± standard deviations; numbers into brackets are ranges. Qualitative variables are expressed as raw numbers followed by percentages into parentheses.

visible on both CT phases and 11 lesions were visible on one CT phase only. Two solitary lesions in two different patients were not visible on both CT phases and therefore not suitable for analysis. These lesions were targeted by fusion of ultrasound images and MR images. The total time for tumor segmentation, scan registration and ablation zone segmentation was 19 min 31 s ± 6 min 39 s (SD) (range: 07 min

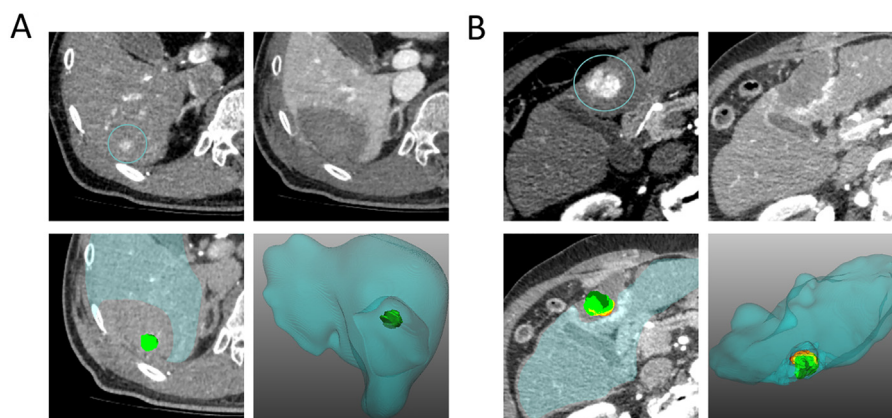


Fig. 4. Two examples of minimal ablation margin quantification of thermal ablation of hepatocellular carcinoma lesions, using in-house built software. A) Small lesion is segment 6 of the liver with an ablation zone > 5 mm. B) large lesion in segment 4 of the liver with 0 mm of minimal ablation margin.

**Table 3**  
Ablation margin quantification parameters.

| Variable                          | Results                       |
|-----------------------------------|-------------------------------|
| Visibility of HCC on CT phase     |                               |
| Arterial and portal venous phases | 18 (58%)                      |
| Portal venous phase only          | 6 (19%)                       |
| Arterial phase only               | 5 (16%)                       |
| Invisible                         | 2 (6%)                        |
| Duration for delineation (min:s)  |                               |
| Tumor                             | 04:28 ± 01:57 [01:13–11:00]   |
| Ablation zone                     | 04:06 ± 01:53 [01:16–11:29]   |
| Registration                      | 06:55 ± 03:38 [01:44–19:12]   |
| Total                             | 19:31 ± 06:39 [07:36–36:01]   |
| Minimal ablation margin (vDSC)    |                               |
| Arterial phase                    | 0.728 ± 3.349 [–5.834–6.203]  |
| Portal venous phase               | 0.554 ± 3.749 [–7.567–10.666] |
| Total                             | 0.626 ± 3.589 [–7.567–10.666] |

Quantitative data are reported as means ± standard deviations; numbers into brackets are ranges. Qualitative variables are expressed as raw numbers followed by percentages into parentheses.  
CT = computed tomography; HCC = hepatocellular carcinoma; vDSC = volumetric Dice similarity coefficient.

36 s–36 min 01 s) (Table 3). vDSC rates of inter- and intra-observer variability for tumor segmentation were 0.815 ± 0.069 (SD) (range: 0.660–0.930) and 0.830 ± 0.073 (SD) (range: 0.512–0.930), respectively (Table 4). The mean MAM was 0.63 ± 3.59 (SD) mm (range: –7.57–10.67 mm). For individual lesions that were visible on both CT phases, a mean volumetric difference of 31.8% between both CT phases was found. However, there was no uniformity in whether delineation in arterial or portal-venous phase resulted in a larger tumor volume.

Two lesions in two patients were invisible on both arterial and portal venous phases, which made them infeasible for ablation margin quantification. In another three lesions in two patients, the registration quality was insufficient (with scores of 3/5, 3/5 and 2/5) to accurately quantify the ablation margins, resulting in a total feasibility of 26/31 (84%) lesions in 16/20 (80%) patients. For the feasible lesions, Bland Altman analysis revealed a mean absolute error between the two observers of 2.15 ± 2.12 mm [SD] (range: 0.39–10.45 mm) for ablation margin quantification, with an ICC of 0.794 (95% confidence interval: 0.594–0.895) (Fig. 5).

Four patients developed LTP in one of the ablated tumors. This included the single patient in whom the thermal ablation was technically unsuccessful but terminated due to a bleeding. In one patient, the tumor was invisible on ultrasound and pre-ablation CECT. This lesion was targeted using fusion navigation of ultrasound and diagnostic MRI. The mean MAM of the other two lesions that developed LTP was –4.00 mm, compared to an overall mean MAM of 0.626 ± 3.589 (SD) mm (range: –6.26–6.65 mm) (Fig. 6). Noteworthy, three patients who developed LTP were treated for recurrent

HCC as part of this study. Moreover, three lesions were peripherally located in the liver dome. Six patients developed new HCC lesions elsewhere in the liver. None of the patients with a MAM > 0 mm developed LTP. In the group of patients with a MAM < 0 mm, 2/9 (22%) patients developed LTP.

#### 4. Discussion

Quantitative ablation margin analysis using co-registration imaging is a promising method to assess technical success of liver tumor TA and has the potential to be superior to visual qualitative assessment [11–14]. Our study demonstrates that a standardized imaging protocol with intraprocedural pre- and post-TA CT under general anesthesia with preoxygenated breath hold allows accurate image co-registration. We found a feasibility for accurate co-registration of pre- and post-TA CECT examinations in 80% of patients and 83.9% of all lesions treated, using the standardized scanning protocol.

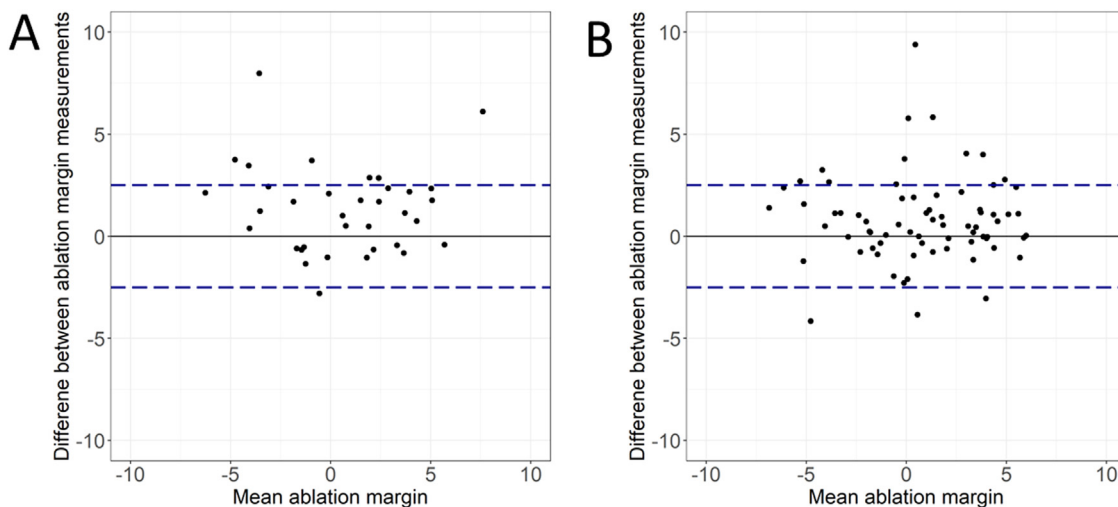
In previous retrospective studies on ablation margin quantification, exclusion rates up to 40% were reported, when co-registration of diagnostic CECT or MRI with post-TA CECT was used [11,14,26,27]. Reasons for inaccurate co-registration in these studies included differences in the position and shape of the liver as a result of differences in patient positioning and breathing mode, and motion artifacts. Feasibility rates similar to ours were found in a study that used intraprocedural pre- and post-ablation CECT, after selection of well demarcated HCC lesions [12]. Standardization of pre- and post-TA CECT protocols thus seem to contribute to the feasibility of ablation margin quantification.

As secondary endpoints, we investigated inter- and intra-observer variability of segmentation of the tumor. Reproducibility of segmentation is a prerequisite for accurate quantitative ablation margin analysis. We found high inter- and intra-observer vDSC rates for tumor segmentation. Although the vDSC is an objective measure of volumetric overlap, it is less sensitive in larger volumes [28]. The DoS within a 2-voxel distance contributes to an objective measure for inter- and intra-observer agreement, as it is volume independent. With DoS values of 0.781 ± 0.140 [SD] (range: 0.536–0.978) and 0.828 ± 0.120 [SD] (range: 0.526–0.982) at a voxel size of 0.71 × 0.71 × 1 mm<sup>3</sup>, high inter- and intra-observer agreement rates were found. In a recent systematic review on ablation margin quantification by Minier et al., only a limited number of studies evaluated inter- and intra-operator characteristics of segmentation or MAM quantification [29]. Most studies used Kappa statistics, and therefore categorized the outcome measure to < 0 mm, 0–5 mm and > 5 mm. Values ranged from 0.68 to >0.95. Kim et al. reported a Kappa of 0.71 for inter-observer agreement of HCC delineation, resulting in a few recommendations for optimization, which were partly implemented in our study (small slice thickness < 3 mm, respiratory motion restriction, multiphase CT, and consensus meeting for overcoming discrepancies) [30]. Using MRI examinations with larger voxel sizes,

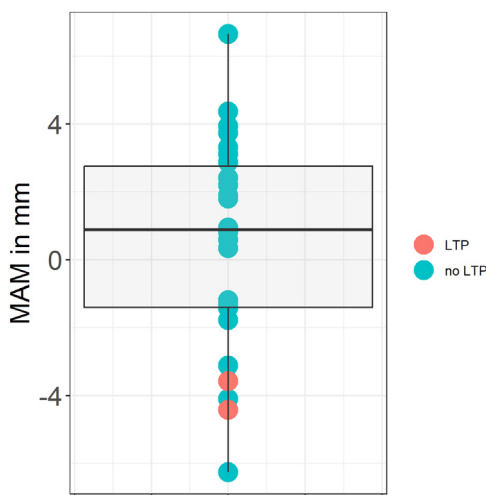
**Table 4**  
Results of inter- and intra observer variability in hepatocellular carcinoma delineation on the different imaging phases.

| Variable  | Results                     |                             |
|---|-----------------------------|-----------------------------|
|   | vDSC                        | DoS                         |
| Inter-observer variability in tumor delineation |                             |                             |
| Arterial phase                                  | 0.816 ± 0.085 [0.608–0.910] | 0.782 ± 0.161 [0.454–0.997] |
| Portal venous phase                             | 0.811 ± 0.087 [0.580–0.930] | 0.780 ± 0.163 [0.379–0.959] |
| Total   | 0.815 ± 0.069 [0.660–0.930] | 0.781 ± 0.140 [0.536–0.978] |
| Intra-observer variability in tumor delineation |                             |                             |
| Arterial phase                                  | 0.823 ± 0.061 [0.665–0.920] | 0.840 ± 0.088 [0.653–0.979] |
| Portal venous phase                             | 0.837 ± 0.085 [0.512–0.930] | 0.815 ± 0.143 [0.526–0.982] |
| Total   | 0.830 ± 0.073 [0.512–0.930] | 0.828 ± 0.120 [0.526–0.982] |

Quantitative data are reported as means ± standard deviations; numbers into brackets are ranges. vDSC= volumetric Dice similarity coefficient. DoS = degree of similarity (chances of two delineations to overlap within a 2 voxel distance).



**Fig. 5.** Bland-Altman plots for A) inter-observer variability and B) intra-observer variability of ablation margin quantification. X-axis shows the mean of the two measurements of quantified ablation margins and the Y-axis shows the difference between the two measurements. The blue dashed lines depict the mean inter-observer difference of 2.5 mm. The mean absolute error between two observers was  $2.15 \pm 2.12$  mm [SD] (range: 0.39–10.45 mm).



**Fig. 6.** Boxplot of minimal ablation margins (MAM) for ablated lesions that developed local tumor progression (LTP) ( $n = 2$ ) vs those that did not develop LTP ( $n = 26$ ).

Hocquelet et al. found DSC values  $> 0.9$  for each lesion in ablation margin quantification, [31]. In our study, we found a mean difference in tumor volume of 31% between segmentations of the same tumor in arterial phase and portal venous phases. This is an important factor to take into account when interpreting the result of quantitative margin analysis. CT level and window settings may influence the volumetric measurements when segmenting a tumor [32]. Other factors that may influence tumor segmentation are the timing of image acquisition, contrast volume and contrast load. Optimized scan timing, standardized CT level and window settings, target segmentation training, and consensus reading may contribute to the robustness of the workflow [33].

In our study, a mean MAM of  $0.626 \pm 3.589$  [SD] mm (range:  $-6.26$ – $6.65$  mm) was found, with 36.5% of all MAM quantifications being  $< 0$  mm. This was considerably lower than the intended ablation margin of at least 5 mm. Despite the limited ablation margin, the LTP rates were in concordance with that of an earlier study in a similar population [34]. Other studies on quantitative ablation margin analyses also found margins that were considerably lower than 5 mm [12–14]. Although a MAM of  $> 5$  mm is associated with good

clinical outcomes, this may often not be achieved in practice. Previous studies reported a MAM  $> 5$  mm in only 2.7% [12] and 37.5% of lesions where they used software-assisted ablation margin quantification [14]. In one study, the overestimation of qualitative interpretation of TA was shown, as the number of patients requiring additional TA increased from 12/150 to 35/150 after introducing software-assisted quantification of MAM with a threshold of 2 mm [35]. In our study, LTP occurred in only 2/9 patients with a quantified MAM  $< 0$  mm. There may be several explanations for this. Experimental studies have demonstrated that considerable tissue shrinkage may occur during TA [36,37]. As a result of the contraction of the ablated tissue, the volume of the ablation zone may be smaller than the tumor volume as determined on the pre-procedural scan, even in a patient in whom the tumor was completely ablated with sufficient margins. Tissue shrinkage may vary considerably depending on individual patient and tumor characteristics, as well as the TA equipment and settings. Also, it occurs non-uniform over time and asymmetrical [38]. Moreover, the software may incorrectly calculate a negative MAM as a result of errors in image co-registration or segmentation of the tumor and/or ablation zone. In fact, our study demonstrates that segmentation is subject to inter- and intra-observer variability, and may be discordant between arterial and portal venous phases. Future research should focus on methods to further reduce such variability to increase reliability and reproducibility of ablation margin quantification, preferably with fast, reliable and fully automated co-registration and segmentation. It is important to stress that a negative MAM does not necessarily mean that a tumor was incompletely ablated as it is only investigated as a predictive factor. On the other hand, none of the patients in our study with a positive MAM-value developed LTP, and a positive MAM value should of course be aimed for.

In this study, a rigid registration algorithm was used for performing quantitative margin assessment. Previous research showed that non-rigid deformations may be present in the liver, especially caused by breathing motion differences [29,39]. Non-rigid registration allows for transformation and scaling of images rather than just translation and rotation to reach an optimal registration [40]. Since the process of TA actually causes local deformation of the tissue, non-rigid algorithms should be restricted in order not to overcompensate for local deformations [41]. Using the standardized intraprocedural scanning protocol with ventilation tube disconnection, the main reason for large deformation (breathing motion and positioning artifacts) were prevented and fast rigid registration with high accuracy was feasible.

Limitations of this feasibility study were the limited number of included patients, the retrospective use of ablation margin quantification software and the limited number of events, which brings limitations to statistical analysis on the MAM results. Despite these limitations we have been able to prospectively demonstrate a feasible pipeline for robust ablation margin quantification.

In conclusion, ablation margin quantification after TA of HCC can be made by implementing a standardized scanning protocol consisting of pre- and post-ablation CECT under breath hold by disconnection of the ventilation tube under general anesthesia. High inter- and intra-observer agreement rates were found for lesion delineation. LTP seemed to be associated with low MAM values (*i.e.*, < 0 mm), but this should further be studied in larger cohort trials.

## Human rights

The authors declare that the work described has been performed in accordance with the Declaration of Helsinki of the World Medical Association revised in 2013 for experiments involving humans.

## Informed consent and patient details

The authors declare that they obtained a written informed consent from the patients included in the article. The authors declare that this report does not contain any personal information that could lead to the identification of the patients.

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## Author contributions

All authors attest that they meet the current International Committee of Medical Journal Editors (ICMJE) criteria for Authorship.

## Declaration of Competing Interest

The authors have no conflicts of interest related to this work to declare.

## CRediT authorship contribution statement

**Pim Hendriks:** Conceptualization, Methodology, Validation, Formal analysis, Investigation, Data curation, Writing – original draft, Visualization. **Kiki M van Dijk:** Validation, Investigation, Writing – original draft. **Bas Boekestijn:** Formal analysis, Investigation, Writing – review & editing. **Alexander Broersen:** Methodology, Software, Investigation, Data curation, Writing – review & editing, Supervision. **Jacoba J van Duijn-de Vreugd:** Resources, Data curation. **Minneke J Coenraad:** Resources, Writing – review & editing. **Maarten E Tushuizen:** Resources, Writing – review & editing. **Arian R van Erkel:** Conceptualization, Resources, Writing – review & editing. **Rutger W van der Meer:** Conceptualization, Resources, Writing – review & editing. **Catharina SP van Rijswijk:** Conceptualization, Resources, Writing – review & editing. **Jouke Dijkstra:** Conceptualization, Software, Writing – review & editing, Supervision. **Lioe-Fee de Geus-Oei:** Methodology, Writing – review & editing, Supervision. **Mark C Burgmans:** Conceptualization, Methodology, Validation, Investigation, Resources, Data curation, Writing – review & editing, Visualization, Supervision.

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