CT of hepatocellular carcinoma in non-alcoholic fatty liver disease: imaging characteristics and inter-rater agreement

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Abstract

Aim: To determine the computed tomography (CT) features of non-alcoholic fatty liver disease (NAFLD) associated hepatocellular carcinoma (HCC).

Methods: In this institutional review board approved study, we reviewed 38 patients with NAFLD (68.4% male; mean age 63 years) with histology confirmed HCC and triphasic liver CT. CT images were independently reviewed by four readers blinded to clinical and pathology data. The reviewers assessed HCC for arterial phase hyper enhancement (APHE), portal venous phase washout (PVWO), delayed phase washout (DPWO), and enhancing capsule. Features of cirrhotic morphology and portal hypertension (PH) were also evaluated. The final CT features were determined by majority and a fifth reader reviewed cases lacking majority. Inter-rater agreement was determined by prevalence-adjusted kappa.

Results: Mean HCC size was 3.6 ± 2.8 cm (range, 1.1-16.0 cm). The HCCs showed APHE in 92.1%, PVWO in 55.3%, DPWO in 81.6%, and enhancing capsule in 44.7%. Cirrhotic morphology was present in 65.8% and PH in 63.2%. Inter-rater agreement was moderate to almost perfect for APHE (0.74-1.0), cirrhosis (0.79-0.89), and PH (0.79-0.95), weak to perfect for DPWO (0.47-0.95) and poor for PVWO (0.42).
Conclusion: NAFLD associated HCC demonstrate less frequent portal venous washout on CT which may affect their imaging diagnosis.

Keywords: Hepatocellular carcinoma, computed tomography, fatty liver, inter-rater agreement, non-alcoholic fatty liver disease

INTRODUCTION

Hepatocellular carcinoma (HCC) is the second most common cause of cancer related death worldwide, with increasing mortality rates in Europe, North America, South America and Africa[1,2]. Imaging plays a pivotal role in management of HCC and is an established method for diagnosis with radiological hallmarks on contrast enhanced multiphase computed tomography (CT) or magnetic resonance imaging (MRI). The imaging hallmark features include arterial phase hyper enhancement (APHE), portal venous phase washout (PVWO) and/or delayed phase washout (DPWO), and presence of enhancing capsule. Based on some or all of the three features, several guidelines have been developed for the non-invasive imaging diagnosis and standardization in reporting of observations suspicious for HCC such as European Association for the Study of the Liver (EASL), European Organization for Research and Treatment of Cancer, Organ Procurement and Transplantation Network (OPTN), American Association for the Study of Liver Diseases (AASLD) and Liver Imaging Reporting and Data System. It should be noted that EASL does not recognize capsule as a major imaging feature of HCC. However, these guidelines have only been validated in patients with most commonly recognized risk factors for development of HCC including, alcoholic cirrhosis and chronic viral hepatitis and not in non-alcoholic fatty liver disease (NAFLD)[3-4].

Although most HCCs (75%-90%) develop in cirrhosis resulting from chronic hepatitis B or C infections and alcoholic injury[9], an estimated 4%-22% of HCC occur in the setting of NAFLD[10-12]. NAFLD has now become the most common cause chronic liver disease in developed countries[13,14]. Given its increasing prevalence worldwide, NAFLD may become the most common chronic liver disease associated with HCC.

NAFLD is a spectrum of disease ranging simple non-alcoholic fatty liver (NAFL) to non-alcoholic steatohepatitis (NASH) that can progress to cirrhosis[15]. NAFL is considered to have minimal risk of progression to cirrhosis and liver failure, while NASH can progress to cirrhosis, liver failure and develop HCC. NASH is thought to be a common underlying cause of cryptogenic cirrhosis as the patients with cryptogenic cirrhosis are comprised mostly of patients with metabolic risk factors including obesity, metabolic syndrome and diabetes[15], but other etiologies such as burnt-out autoimmune hepatitis and occult alcoholism may also result in cryptogenic cirrhosis[16-18]. HCCs are known to occur in patients with NAFLD in the absence of cirrhosis[10,19-21] and these HCCs may not meet the imaging criteria based on the current guidelines including LIRADS[22].

The effect of hepatic steatosis in NAFLD on the imaging features of HCC has not yet been fully explored. For example, hepatic steatosis may decrease the liver attenuation on CT and as a result, washout observation may be absent or less conspicuous which could render the LIRADSv2017 imaging criteria not applicable[22-25]. HCCs can also occur in the absence of cirrhosis in patients with NAFLD as mentioned earlier. Therefore based on these premises, the purpose of our study was to determine the major imaging features of HCC on multiphase CT and the inter-observer agreement in patients with NAFLD.

METHODS

In this institutional review board (IRB) approved (ID: 15-004925), HIPPA-compliant retrospective study. Written informed consent for retrospective review of data was waived.
Patient selection

We reviewed our institutional pathology and imaging database between January 2006 and December 2016 with key words NAFLD, hepatic steatosis or steatohepatitis and HCC or hepatocellular carcinoma. This yielded a cohort of 400 patients. Among these 400 patients, only 38 patients met the AASLD criteria for NAFLD[15] and had a triple phasic CT (late arterial, portal venous and delayed phase) before histological confirmation of HCC. Patients with cryptogenic cirrhosis were excluded due to uncertainty of the underlying etiology. Among the final group of 38 patients, 24 had NASH cirrhosis (cirrhosis with current or past evidence of steatosis or steatohepatitis) and 14 patients had no cirrhosis- 7 NASH (hepatic steatosis ≥ 5% with inflammation ± fibrosis) and 7 NAFL (hepatic steatosis ≥ 5% without evidence of hepatocellular injury or fibrosis). A flowchart detailing the patient selection and subgroups is shown in Figure 1.

Patient age, sex, height, weight, body mass index (BMI), serum cholesterol, serum triglycerides, presence or absence of obesity, tumor histopathological and clinical management information were obtained from electronic medical records.

Cytological and pathological TNM staging was evaluated according to criteria of the 7th American Joint Committee on Cancer[26]. The diagnosis of NAFL, NASH was established at pathology. The time interval between CT and surgical pathology was 137 ± 387 days (range 3 to 1802 days). Twenty-four cases had surgical pathology within 6 months of CT. Histology of the HCC and background liver was evaluated by an experienced pathologist (TM) with expertise in NAFLD. HCCs were graded based on WHO classification and NAFLD was graded based on NAS score.

CT imaging review

All the CT images were independently reviewed on PACS Workstation (Centricity, GE Healthcare, Waukesha, WI) by four board-certified abdominal radiologists (SPS, ECE, AK, CAB) who were blinded to clinical and pathological findings other than the presence of HCC. Each reader recorded the imaging features of HCC, including size, location, APHE, PVWO, DPWO, and presence capsule. Readers also assessed for findings of cirrhosis-surface nodularity, caudate lobe hypertrophy, left lobe enlargement, widened fissures, widened gallbladder fossa, and portal hypertension (PH), splenomegaly, collaterals
and gastroesophageal varices. In patients with multiple HCCs, only the largest HCC with histological confirmation was assessed. The final imaging features of HCCs were determined by majority. A fifth reader blinded to clinical and pathological findings reviewed cases lacking majority. LIRADs criteria is only applicable in patients with cirrhosis or chronic viral hepatitis and therefore would not be applicable for patients with NAFLD without cirrhosis.

Statistical analysis
Continuous variables were expressed as mean ± standard deviation and categorical data as percentage. Inter-rater agreement was determined by prevalence-adjusted bias-adjusted Cohen's kappa[27]. Agreement between cirrhotic on CT and pathology was also determined by prevalence-adjusted Cohen's kappa[27]. Inter observer agreement was classified as none (0-0.2), minimal (0.21-0.39), weak (0.40-0.59), moderate (0.60-0.79) and strong (0.80-0.90) and almost perfect (> 0.90). Differences between non-cirrhotic and cirrhotic sub-groups were compared using an unpaired $t$-test with equal variance assumption for continuous data and Fischer's exact test for categorical data. Statistical significance was assumed for $P$ values of less than 0.05. Data were analyzed using JMP 11.0 (SAS, Cary, NC) and Prism 5.0 (GraphPad Software, Inc, La Jolla, CA).

RESULTS
Clinical and pathological characteristics
Mean age of subjects was 63 years (range 45-79). Patients were predominantly male ($n = 26$, 68.4%), diabetic ($n = 28$, 73.7%), and obese (BMI ≥ 30; $n = 25$, 65.8%). Mean ± SD total serum cholesterol and triglyceride values were 151.9 ± 39.7 (range: 36-229 mg/dL) and 130.6 ± 53.5 (range: 54-237 mg/dL), respectively. Most of the patients underwent liver transplant ($n = 21$, 55.3%) or hepatic resection or segmentectomy ($n = 12$, 31.6%). Of the 21 patients that underwent liver transplant, 16 patients received chemoembolization (14) or ablation (2) before transplantation, 4 patients received only ablation or chemoembolization, and 1 patient was lost to follow-up. Of 38 patients, 2 (5.3%) patients developed recurrent HCC. CT studies performed before any treatment was used for imaging analysis.

At pathology, HCCs were well differentiated in 14 patients (36.8%), well to moderately differentiated in 5 patients (13.2%), moderately differentiated in 16 patients (42.1%), moderate to poorly differentiated in 2 patients (5.3%) and poorly differentiated in 1 patient (2.6%). Hepatic steatosis was minimal (26.3%) in 10 patients, mild in 26 patients (68.4%), and moderate in 2 patients (5.3%). Steatohepatitis was present in 25 patients (65.8%) and cirrhosis in 24 patients (63.2%). The clinical and pathological characteristics are summarized in Table 1.

HCC and liver parenchyma imaging characteristics - majority features
Mean HCC size was 3.6 ± 2.8 cm (range: 1.1-16.0 cm). APHE was seen in 92.1%, PVWO in 55.3% DPWO in 81.6% and enhancing capsule in 44.7% [Figures 2 and 3]. Fat within the HCC was present in only one patient. Cirrhotic morphology was present in 25 patients (65.8%) and portal hypertension in 24 patients (63.2%). The imaging features are summarized in Table 2.

Non-cirrhotic NAFLD vs. cirrhotic NAFLD
Non-Cirrhotic NAFLD (NAFL in 7 and NASH in 7) and cirrhotic NAFLD were present in 14 (36.8%) and 24 (63.2%) patients respectively. Patients with non-cirrhotic NAFLD were older ($P = 0.03$), had larger mean HCC size ($P = 0.008$) and higher degree of hepatic steatosis ($P = 0.003$). PVWO feature was observed significantly more in the non-cirrhotic group as compared to the cirrhotic group (78.6% vs. 41.7%, $P = 0.04$). Portal hypertension features were more commonly seen in patients with cirrhotic NAFLD (91.7% vs. 14.3%, $P < 0.0001$). There was no significant difference between the two groups with respect to gender distribution ($P = 0.47$), BMI ($P = 0.14$), presence of diabetes ($P = 0.45$), cholesterol level ($P = 0.24$), triglyceride level
Inter-rater agreement was moderate to almost perfect for HCC APHE (0.74-1.0), none to moderate for PVWO (0-0.42), weak to almost perfect for DPWO (0.47-0.95) [Figure 4] and none to moderate for capsule (0.05-0.79). The inter-rater agreement was moderate to almost perfect for cirrhosis (0.79-0.95). Pathology and CT agreement for presence of cirrhosis was strong at 0.84. Inter-rater agreement for imaging features is summarized in Table 3.

DISCUSSION

In our study of NAFLD associated HCC, many HCCs did not demonstrate major imaging features particularly PVWO and enhancing capsule were absent in nearly half and DPWO was absent in nearly 20% of the patients. Cirrhotic morphology was present in 25 (65.8%) patients. A third (36.8%) of HCC occurring in non-cirrhotic livers would not be eligible for LIRADs classification.

Our study results are in agreement with previous reports in literature that HCC can occur even in the absence of steatohepatitis or fibrosis/cirrhosis\cite{15,19-21,24-26}. Also HCCs in non-cirrhotic liver were larger in agreement with existing literature that HCCs in non-cirrhotic liver usually present at a later stage and are

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Table 1. Clinical-pathological characteristics in patients with NAFLD-associated HCC (n = 38; main) by non-cirrhotic (n = 14) and cirrhotic (n = 24) liver

<table>
<thead>
<tr>
<th>Clinical and Pathological findings</th>
<th>Main Cohort (n = 38)</th>
<th>Non-cirrhotic (n = 14)</th>
<th>Cirrhotic (n = 24)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years) (mean ± SD)</td>
<td>63 ± 7.2</td>
<td>66.2 ± 6.3</td>
<td>60.9 ± 7.2</td>
<td>0.03</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>12 (31.6)</td>
<td>3 (21.4)</td>
<td>9 (37.5)</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>26 (68.4)</td>
<td>11 (78.6)</td>
<td>15 (62.5)</td>
<td></td>
</tr>
<tr>
<td>BMI (kg/m²) (mean ± SD)</td>
<td>32.2 ± 4.9</td>
<td>30.6 ± 4.6</td>
<td>33.1 ± 5.0</td>
<td>0.14</td>
</tr>
<tr>
<td>Diabetes (%)</td>
<td>28 (73.7)</td>
<td>9 (64.3)</td>
<td>19 (79.2)</td>
<td>0.45</td>
</tr>
<tr>
<td>Total Cholesterol (mean ± SD)</td>
<td>151.9 ± 39.7</td>
<td>171 ± 36.4</td>
<td>147.4 ± 39.9</td>
<td>0.24</td>
</tr>
<tr>
<td>Triglycerides (mean ± SD)</td>
<td>130.6 ± 53.5</td>
<td>129.2 ± 71.1</td>
<td>131 ± 50.7</td>
<td>0.95</td>
</tr>
<tr>
<td>HCC Pathology Source</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biopsy</td>
<td>5 (13.2)</td>
<td>3 (21.4)</td>
<td>2 (8.3)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Surgical Resection</td>
<td>12 (31.6)</td>
<td>10 (71.4)</td>
<td>2 (8.3)</td>
<td></td>
</tr>
<tr>
<td>Liver Explant at Transplant</td>
<td>21 (55.3)</td>
<td>1 (7.1)</td>
<td>20 (83.3)</td>
<td></td>
</tr>
<tr>
<td>Pathologic Features</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tumor Grade</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Well differentiated</td>
<td>14 (36.8)</td>
<td>4 (29)</td>
<td>10 (41.7)</td>
<td>0.31</td>
</tr>
<tr>
<td>Well-Moderately differentiated</td>
<td>5 (13.2)</td>
<td>1 (7.1)</td>
<td>4 (16.7)</td>
<td></td>
</tr>
<tr>
<td>Moderately differentiated</td>
<td>16 (42.1)</td>
<td>9 (64.3)</td>
<td>7 (29.2)</td>
<td></td>
</tr>
<tr>
<td>Moderate-Poorly differentiated</td>
<td>2 (5.3)</td>
<td>2 (8.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poorly differentiated</td>
<td>1 (2.6)</td>
<td>1 (4.2)</td>
<td></td>
<td></td>
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<tr>
<td>Hepatic Steatosis Grade</td>
<td></td>
<td></td>
<td></td>
<td>0.003</td>
</tr>
<tr>
<td>Minimal</td>
<td>10 (26.3)</td>
<td></td>
<td>10 (41.7)</td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>26 (68.4)</td>
<td>12 (85.7)</td>
<td>14 (58.3)</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>2 (5.3)</td>
<td>2 (14.3)</td>
<td></td>
<td></td>
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<tr>
<td>Hepatic Fibrosis Stage</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>9 (23.7)</td>
<td>9 (64.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>3 (7.9)</td>
<td>3 (21.4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2 (5.3)</td>
<td>2 (14.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>24 (63.2)</td>
<td></td>
<td>24 (100)</td>
<td></td>
</tr>
</tbody>
</table>

P-value for non-cirrhotic group vs. cirrhotic group; continuous variables analyzed using unpaired t-test with equal variance assumption; categorical data analyzed using Fischer’s exact test. BMI: body mass index; NAFLD: non-alcoholic fatty liver disease; HCC: hepatocellular carcinoma

(P = 0.95), tumor size (P = 0.67), tumor grade (P = 0.31), APHE (P = 1.00), DPWO (P = 1.00) and enhancing capsule (P = 0.09).
larger in size at presentation\textsuperscript{[26]}. This may be due to lack of screening guidelines in non-cirrhotic patients with NAFLD\textsuperscript{[3,28]}. Larger tumor size seen in non-cirrhotic livers at presentation may make them ineligible for transplant based on Milan criteria for transplantation\textsuperscript{[35,36]}.

Imaging features of HCC on multiphase CT and MRI are based on sequential changes in the intra-lesional blood supply during hepatocarcinogenesis. Advanced HCCs receive their blood supply predominantly

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**Table 2. Imaging features of HCC and liver parenchyma at CT by majority consensus in patients with NAFLD-associated HCC (n = 38) by non-cirrhotic (n = 14) and cirrhotic (n = 24) liver morphology**

<table>
<thead>
<tr>
<th>HCC imaging features</th>
<th>Main cohort (n = 48)</th>
<th>Non-cirrhotic pathology (n = 14)</th>
<th>Cirrhotic pathology (n = 24)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tumor size (mean ± SD)</td>
<td>3.6 ± 2.8</td>
<td>5.1 ± 3.9</td>
<td>2.7 ± 1.3</td>
<td>0.008</td>
</tr>
<tr>
<td>APHE</td>
<td>35 (92.1)</td>
<td>13 (92.8)</td>
<td>22 (91.7)</td>
<td>1.00</td>
</tr>
<tr>
<td>PVW0</td>
<td>21 (55.3)</td>
<td>11 (78.6)</td>
<td>10 (41.7)</td>
<td>0.04</td>
</tr>
<tr>
<td>DPW0</td>
<td>31 (81.6)</td>
<td>12 (85.7)</td>
<td>19 (79.2)</td>
<td>1.00</td>
</tr>
<tr>
<td>Enhancing “Capsule”</td>
<td>17 (44.7)</td>
<td>9 (64.3)</td>
<td>8 (33.3)</td>
<td>0.09</td>
</tr>
<tr>
<td>Cirrhotic Liver Morphology</td>
<td>25 (65.8)</td>
<td>2 (14.3)</td>
<td>23 (95.8)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Portal hypertension</td>
<td>24 (63.2)</td>
<td>2 (14.3)</td>
<td>22 (91.7)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>LIRADS Score*</td>
<td></td>
<td></td>
<td></td>
<td>0.29</td>
</tr>
<tr>
<td>2</td>
<td>3 (7.9)</td>
<td>0</td>
<td>3 (12.5)</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>11 (28.9)</td>
<td>3 (21.4)</td>
<td>8 (33.3)</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>24 (63.2)</td>
<td>11 (78.6)</td>
<td>13 (54.2)</td>
<td></td>
</tr>
</tbody>
</table>

* LIRADS score applied to see if HCCs would meet the criteria, however if there is no cirrhosis, LIRADS criteria should not be applied as per guidelines. APHE: arterial phase hyperenhancement; PVW0: portal venous washout; DPW0: delayed phase washout

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**Figure 2.** A 70-year old male with histologically confirmed hepatocellular carcinoma in non-cirrhotic liver with mild steatosis and no steatohepatitis (non-alcoholic fatty liver). Patient was obese (BMI of 30.7), diabetic and dyslipidemic. On multiphase contrast enhanced CT (A-C), a heterogeneously enhancing mass (arrow) on arterial phase (A) with no washout on portal venous (B) and complete washout on delayed phase (C) images can be seen. In addition, a thin but incomplete capsule (arrowhead) can be seen on delayed phase image (C). No features of portal hypertension were seen on imaging.

**Figure 3.** A 73-year old male with histologically confirmed hepatocellular carcinoma in non-cirrhotic liver with severe steatohepatitis (non-alcoholic steatohepatitis). Patient was obese (BMI of 31.1) and diabetic. On multiphase contrast enhanced CT (A-C), a heterogeneously enhancing mass (arrow) on arterial phase (A) showing no washout on portal venous (B) and delayed phase (C) images can be seen.
from the anomalous arteries (arterial blood supply)\[^{37,38}\]. This results in high arterial flow which manifests as APHE on dynamic multiphase imaging followed by washout (de-enhancement of a HCC, greater enhancement of the surrounding liver, or a combination of both factors) on portal venous and/or delayed phase imaging\[^{39-43}\]. The washout is attributed to diminished portal venous blood supply of the HCC, high tumoral cellularity with associated small extracellular volume, and expanded extracellular space of the surrounding fibrotic liver\[^{37,38,43,44}\].

Effect of NAFLD on imaging features on HCC is currently being explored. In our study APHE was present in most (92.1\%) of the cases. APHE is a major imaging criterion and has a good sensitivity for detection of HCC, ranging from 65\%-96\%\[^{39,43,45}\]. The lower sensitivity can be seen in early/well differentiated HCC with partial neovascularization. APHE is more sensitive than other enhancement features but lacks specificity for

**Table 3. Inter-observer agreement for HCC features and liver parenchyma morphology at MRI in patients with NAFLD-associated HCC (n = 38)**

<table>
<thead>
<tr>
<th>HCC imaging features</th>
<th>R1 vs. R2</th>
<th>R1 vs. R3</th>
<th>R1 vs. R4</th>
<th>R2 vs. R3</th>
<th>R2 vs. R4</th>
<th>R3 vs. R4</th>
</tr>
</thead>
<tbody>
<tr>
<td>APHE</td>
<td>0.74</td>
<td>0.79</td>
<td>0.79</td>
<td>0.74</td>
<td>0.84</td>
<td>1.0</td>
</tr>
<tr>
<td>PVWO</td>
<td>0.42</td>
<td>0.00</td>
<td>0.53</td>
<td>0.05</td>
<td>0.32</td>
<td>0.05</td>
</tr>
<tr>
<td>DPWO</td>
<td>0.95</td>
<td>0.84</td>
<td>0.74</td>
<td>0.79</td>
<td>0.68</td>
<td>0.47</td>
</tr>
<tr>
<td>Enhancing “Capsule”</td>
<td>0.47</td>
<td>0.05</td>
<td>0.42</td>
<td>0.37</td>
<td>0.79</td>
<td>0.37</td>
</tr>
<tr>
<td>Cirrhotic Liver Morphology</td>
<td>0.79</td>
<td>0.89</td>
<td>0.89</td>
<td>0.79</td>
<td>0.79</td>
<td>0.89</td>
</tr>
<tr>
<td>Portal hypertension</td>
<td>0.74</td>
<td>0.79</td>
<td>0.95</td>
<td>0.84</td>
<td>0.79</td>
<td>0.84</td>
</tr>
</tbody>
</table>

Data are presented as prevalence-adjusted bias-adjusted kappa. R1: reader 1; R2: reader 2; R3: reader 3; R4: reader 4; APHE: arterial phase hyperenhancement; PVWO: portal venous phase washout; DPWO: delayed phase washout
HCC. The specificity can be increased by combined assessment of APHE & washout (PVWO or DPWO). Sensitivity and specificity of combined APHE & washout for diagnosis of HCC ranges from 43% to 98%, and 81% to 100%, respectively.

For assessment of washout, either portal venous or delayed phase imaging can be used. In this study, PVWO was absent in 17 (44.7%) of the HCCs. In contrast, DPWO was absent in only 7 (18.4%) HCCs. In addition, the interobserver agreement was none to moderate for PVWO and weak to almost perfect for DPWO. These results support the added value of delayed phase imaging in assessment of washout. Triple phase CT is the standard protocol for HCC in chronic liver disease, however some centers may not include the delayed phase to reduce the radiation dose. However, this may lead to loss of important diagnostic information (washout characteristics). In a prior study with NAFLD associated HCC, PVWO was absent in 30% cases but DPWO was not reported. Ehman et al. reported lack of washout (portal or delayed) in only 18% (15% on CT and 21% on MRI) of the 184 pathologically proven HCCs with cirrhosis resulting from several different etiologies. Washout in NAFLD associated HCC may not be demonstrated well probably due to hepatic steatosis that can lead to liver hypo-attenuation on post-contrast enhanced images, thereby resulting in the appearance of persistent HCC hyper-attenuation or iso-attenuation during the portal venous and delayed phase imaging. Interestingly however in our study PVWO was more commonly seen in subgroup with non-cirrhotic NAFLD, which had higher proportion of cases with mild moderate steatosis in comparison to cirrhotic NAFLD subgroup. These findings suggest that degree of hepatic steatosis may have more determining effect than presence of fibrosis for washout appearance. Future studies with larger number of patients are required to confirm our findings. The findings should be confirmed in studies with larger population of NAFLD associated HCCs.

Capsule appearance is a highly specific but not very sensitive feature of HCC. Capsule is thought to be caused by expansile growth of HCC causing perilesional compression of liver tissue, which appears as enhancing rim around the HCC in portal venous or delayed phase. Capsule is supplied by the portal venous system leading to this delayed enhancement. Interestingly, capsule is a recognized major feature of HCC based on the LIRADS, OPTN guidelines but not on the AASLD and EASL guidelines. In our study, capsule appearance was seen in only 17 (44.7%) of the NAFLD-associated-HCCs but still within the reported range of 27 to 64% in studies including all chronic liver diseases.

Inter-rater agreement was poor for PVWO, variable for DPWO and capsule suggesting the difficulty in interpretation of washout and detection of capsule in NAFLD associated HCCs. We had four experienced abdominal radiologists working in tertiary level institute with large exposure to HCCs on a daily basis. Even with this level of expertise, the interobserver agreement for PVWO was poor. We think that this is due to the variable hepatic steatosis in background liver that affects interpretation of the washout particularly PVWO.

The study has some limitations. Despite searching a large patient database containing around 2,500 patients (pathological data for HCC), our final study cohort was relatively small. Due to retrospective nature of the study, the imaging (triphasic CT) technique including scanner, sequence protocol(s) and contrast agent(s) was not uniform over the study period and this may have introduced variability in the phase of image acquisition. Some cases of NAFLD associated HCCs may have been excluded that either lacked pathological assessment or were interpreted as cryptogenic cirrhosis. The readers were aware of the diagnosis of HCC in the lesions which may have introduced bias for imaging features. However this bias of prior knowledge did not inflate interobserver agreement for the imaging features. Degree of hepatic steatosis may have changed from time between histological analysis and radiological assessment, due to systemic interventions or natural progression of NAFLD. This was unavoidable as several patients received locoregional treatment before undergoing surgery or liver transplantation. These treatments may also have...
contributed to the changes in the liver parenchyma. However we minimized this variation as most of the studies had histological evaluation within 6 months of CT study.

In conclusion, NAFLD associated HCC may not show portal venous phase washout on CT and may impact the imaging diagnosis of HCC. Our study should be confirmed in studies with larger population of NAFLD associated HCCs. There may be a need for modification of criteria for multiphase CT based diagnosis of NAFLD associated HCCs particularly in the non-cirrhotic patients.

DECLARATIONS

Authors’ contributions
Made substantial contributions to conception and design of the study and performed data analysis and interpretation: Garg I, Thompson SM, Mounajjed T, Ehman EC, Venkatesh SK
Performed data acquisition, as well as provided administrative, technical, and material support: Garg I, Thompson SM, Sheedy SP, Mounajjed T, Khandelwal A, Ehman EC, Bookwalter CA, Venkatesh SK

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All authors declared that there are no conflicts of interest.

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Consent for publication
Not applicable.

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