Hepatocellular Carcinoma Response to Local Regional Therapy; Correlations between Pre-Liver Transplants Imaging and Explant Pathology

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Keywords: HCC; Mrecist; Liver transplant; Explant; Pathology

Introduction

Hepatocellular carcinoma (HCC) is the fifth most common cancer in the world and ranks third in cancer related mortality [1]. Different modalities can be used for treatment according to Barcelona algorithm [2]. Locoregional therapy (LRT) which includes radiofrequency ablation (RFA) and Trans-arterial chemotherapy embolization (TACE) are used to treat patients through different stages of the disease according to BCLC algorithm [3]. At early stages of the tumor, RFA can be used with curative intent while TACE is considered a palliative treatment if patient deemed not a transplant candidate. In pre-transplant setting, these therapies frequently used as a bridge to transplant or for down staging the tumor to be within acceptable criteria for transplant. The ultimate goal of LRT to induce necrosis of the tumor. This can be achieved either by thermocoagulation (RFA) or by arterial occlusion and local administration of chemotherapy (TACE). The degree of tumor necrosis is a predictor of treatment success. Partial necrosis within the tumor leads to higher rate of tumor recurrence after liver transplantation [4]. Achieving complete necrosis is associated with lower rate of recurrence and improved survival post liver transplantation [5,6].

The radiological assessment of tumor response to Locoregional therapy has evolved to include the viability of tumor in comparison to the previously used criteria (RECIST and WHO criteria), which are based on the sum of the greatest dimensional measurement of target lesions [7]. Modified Response Evaluation Criteria in Solid Tumors (mRECIST) criteria has been introduced to overcome the limitation of measuring the tumour size only by taking into account the residual viable tumor [8,9]. The treatment response categorized into four groups according to the extent of necrosis: complete response, partial response, stable disease, or progressive disease [10].

The accuracy of measuring the tumor necrosis after LRT on dynamic imaging is under investigation. The main objective of this study, to compare the estimated viable HCC after LRT by CT imaging and before liver transplant to the histopathological assessment of viable HCC in the hepatic explant.

Abstract

Background/Aim: Modified Response Evaluation Criteria in Solid Tumors (mRECIST) were developed to assess the response to treatment in patients with HCC, based on measuring the amount of viable tumor using dynamic imaging (CT/MRI). Our aim to compare the estimated viable HCC after locoregional therapy (LRT) by CT imaging and prior to liver transplant to the histopathological assessment of viable HCC in the hepatic explant.

Methods: We prospectively evaluated 44 patients with HCC who underwent both LRT and liver transplantation at London health science center. Using mRECIST criteria, the response to LRT was assessed by two blinded radiologists and the percentage of necrosis was reported separately for the reference CT (rCT) done after the last LRT and prior to liver transplantation. The report obtained from each radiologist was combined then was compared to the findings of an expert pathologist reporting on viable tumour present and tumour necrosis in the hepatic explants. Both parties were blinded to prevent bias in the results.

Results: A total of forty-one transplant recipients fulfilled the inclusion criteria for the study. At time of listing 100% were within total volume criteria, 86% within UCSF, and 68% within Milan. The average time from the last reference CT scan to liver transplant was 57.7 days; the average time from last LRT to reference CT was 72.5 days. Thirty-four recipients (83%) had accurate assessment for necrosis (mRECIST) within 20% comparing rCT to explant (i.e. concordant). Nineteen, 19 (46%) of the 41 predicted 100% concordance. Only 7/41 (17%) had a poor correlation (>50%) between histology and reference CT images. Positive correlation was detected with the correlation coefficient is calculated as 0.5723. Our study demonstrated CT-pathologic correlation in predicting total volume with correlation coefficient is calculated as 0.014.

Conclusion: Dynamic CT is an accurate tool to evaluate the tumour response prior to liver transplantation and the likelihood of underestimating the tumour burden is low. With expert radiologists and pathologists, the correlation is acceptable and supports the ongoing use of frequent dynamic imaging to evaluate responses to LRT and determining transplant eligibility.
Materials and Methods

This is a single Centre, correlation analysis of a prospective study approved by the local Institutional Review Board (IRB). Between January 2010 to December 2015, we evaluated all patients at London Health Sciences Centre with known HCC who underwent LRT as bridging therapy prior to liver transplantation.

Inclusion criteria included any patients with proven diagnosis of HCC either, by histopathology or dynamic imaging (CT/MRI). All participants were eligible for liver transplant based on HCC tumor burden within Ontario criteria which corresponds to total volume of 115 mm or less and alpha fetoprotein less than 400 [11]. The patients were listed for liver transplant with MELD exceptional points of 22 and received LRT as bridging therapy to transplant.

The consensus of a multidisciplinary team. In case of RFA, the procedure performed through percutaneous approach by using ultrasound guidance to insert 18-gauge, cooled-tip electrode with a 2 or 3 cm exposed tip which is attached to a radiofrequency generator. Radiofrequency energy delivered in pulses of 6-12 minutes and position changed in larger lesion to achieve complete ablation TACE involves identifying the blood vessels supplying the tumor during an angiogram and selectively delivering chemotherapeutic agent (doxorubicin 30 mg) mixed with Lipiodol followed by embolization of the vessel for complete occlusion of blood supply.

Baseline CT was done before each LRT as part of assessment. A reference CT (rCT) was done within 3 months after therapy to assess response to LRT. The radiological response was evaluated using the mRECIST criteria which are outlined in Table 1.

<table>
<thead>
<tr>
<th>Stable disease (SD)</th>
<th>Any cases that do not qualify for either partial response or progressive disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Progressive disease (PD)</td>
<td>An increase of at least 20% in the sum of the diameters of viable (enhancing) target lesions, taking as reference the smallest sum of the diameters of viable (enhancing) target lesions recorded since treatment started</td>
</tr>
<tr>
<td>Complete response (CR)</td>
<td>Disappearance of any intra tumoral arterial enhancement in all target lesions</td>
</tr>
<tr>
<td>Partial response (PD)</td>
<td>At least a 30% decrease in the sum of diameters of viable (enhancement in the arterial phase) target lesions, taking as reference the baseline sum of the diameters of target lesions.</td>
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</tbody>
</table>

Table 1: mRECIST, modified Response Evaluation Criteria in Solid Tumors.

Two blinded radiologists appraised the response to LRT and the percentage of necrosis was reported separately for each R CT done after each LRT and prior to liver transplantation. The results from the radiologists were compared to the findings of an expert pathologist reporting on viable tumour present and tumour necrosis in the hepatic explants. The pathologist determined the tumor viability by examining the largest diameter of the treated HCC. Both parties were blinded so prevent bias in the results.

To quantify the correlation between pre-liver transplantation radiological imaging to histopathological findings post-liver transplantation, we applied statistical analysis to four main factors under both radiology and pathology methods, which are percentage of Necrosis, number of tumors, tumor size and total tumor volume. In addition, two statistical methodologies are carried out to get consistent and reliable results. Even though of patients under observation have multiple tumors, detailed information is only accessible for the 1st tumor. Thus most of our statistical analysis is based upon the 1st tumor’s information.

First, a direct calculation of Pearson product-moment correlation coefficient is conducted for each of the four main factors. This statistic measures the linear dependence between two variables and lies between -1 and +1, with -1 indicating extremely negative correlation or moving in the opposite direction, and +1 indicating extremely positive correlation or moving in the same direction. Also, a corresponding plot of each main factor is provided to compare the movement of both results under both radiology and pathology visually.

Second, we assume there is a linear relationship between radiology and pathology results for each main factor and then analyzes how good this linear model is, or how strong the two variables of radiology and pathology results are linearly related. Through this regression analysis, we focus on the “Multiple R” in the output, which indicates the quality of the linear relationship. This Multiple R varies from 0 to 1 with larger number indicating stronger linear relationship. Also, a corresponding scatter plot with fitted linear trend line is provided for visual perception. Each dot on the plot represents a patient’s information with horizontal axis of radiology result and vertical axis of pathology results.

Results

A total of 44 transplant recipients fulfilled the inclusion criteria for the study. Three patients were excluded due to absence of a rCT after LRT and prior to liver transplant. The median age at diagnosis was 59.4 years.

There was male predominance in our patients (85%). The major indication for liver transplant in our group was liver cirrhosis secondary to hepatitis C (73%) (Table 2). Summarize the baseline characteristics of our cohort.
Mean Age at transplant: 59.4 years

Gender

<table>
<thead>
<tr>
<th>Gender</th>
<th>Male</th>
<th>Female</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>35/41 (85%)</td>
<td>6/41 (15%)</td>
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</table>

Diagnosis

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>HCV</th>
<th>NASH</th>
<th>ASH</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>30/41 (73%)</td>
<td>4/41 (9.75%)</td>
<td>3/41 (7.3%)</td>
<td>4/41 (9.75%)</td>
</tr>
</tbody>
</table>

AFP (Mean): 71 ng/mL

Transplant Criteria (Radiology)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Ontario</th>
<th>Milan</th>
<th>UCSF</th>
</tr>
</thead>
<tbody>
<tr>
<td>R value</td>
<td>41/41 (100%)</td>
<td>30/41 (68%)</td>
<td>37/41 (86%)</td>
</tr>
</tbody>
</table>

Transplant Criteria (Pathology)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Ontario</th>
<th>Milan</th>
<th>UCSF</th>
</tr>
</thead>
<tbody>
<tr>
<td>P value</td>
<td>41/41 (100%)</td>
<td>21/41 (50%)</td>
<td>33/41 (80%)</td>
</tr>
</tbody>
</table>

Local regional Therapy (LRT)

<table>
<thead>
<tr>
<th>Therapy</th>
<th>TACE</th>
<th>RFA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>31/41 (75.5%)</td>
<td>10/41 (24.5%)</td>
</tr>
</tbody>
</table>

No. of tumors

<table>
<thead>
<tr>
<th>Source</th>
<th>Radiology</th>
<th>Pathology</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2.1 (1-6)</td>
<td>1.9 (1-4)</td>
</tr>
</tbody>
</table>

### Table 2: Summarize the baseline characteristics of our cohort.

- UCSF: University of California, San Francisco
- TACE: Trans-arterial chemotherapy embolization
- RFA: Radiofrequency ablation
- HCV: Hepatitis C vaccine
- NASH: Nonalcoholic steatohepatitis
- ASH: Hepatitis A
- AFP: Acute flaccid paralysis

At time of listing, all of our patients were within Ontario criteria. Majority of patients were within UCSF (86%) and Milan (68%). The HCC was single lesion was in 50% and only one lobe was affected in 70%. Most of our patients were treated with TACE (75%).

13 patients (31%) had more than one LRT (12 had >1 TACE and two had >1 RFA). No patients progressed beyond transplant criteria. The average time frame from the last reference CT scan to liver transplant was 57.7 days; the average time from last LRT to reference CT was 72.2 days.

Using mRECIST criteria for radiological evaluation of Tumor necrosis, 29 (70%) and 8 (19.5%) patients achieved complete response (CR) and partial response (PR), respectively. The histopathological assessment of liver explants revealed 24 (58.5%) and 14 (34%) patients with complete and partial response, respectively. Evaluation of tumor response summarized in Table 3.

### Table 3: Evaluation of tumor response according to mRECIST criteria.

<table>
<thead>
<tr>
<th></th>
<th>Complete response</th>
<th>Partial response</th>
<th>Stable disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiological response</td>
<td>29 (70%)</td>
<td>8 (19.5%)</td>
<td>3 (7%)</td>
</tr>
<tr>
<td>Pathological Response</td>
<td>24 (58.5%)</td>
<td>14 (34%)</td>
<td>3 (7%)</td>
</tr>
</tbody>
</table>

### Figure 1: Correlation of necrosis target lesions.

### Figure 2: Correlation of number of tumors.
Dynamic CT was accurate in assessing the number of target lesions compared to Pathology (Correlation Coefficient 0.6648) (Figure 2). The size of 1st tumor was only obtained in 35 patients. From these patients, the size measurement on dynamic CT corresponds well with the histopathological findings (Correlation Coefficient 0.5616) (Figure 3). The predicted total tumor volume from dynamic CT showed weak association with the predicted value from histopathology (Correlation Coefficient 0.2379) (Figure 4).

Discussion

EASL-AASLD guidelines adopted the modified RECIST (mRECIST) criteria which takes into account the viability of tumor on dynamic imaging (CT/MRI) to accurately predict response to locoregional treatment. mRECIST criteria provided significantly better prognostic value compared to other methods of assessment. A study by Gillmore et al. comparing assessment of response to TACE by conventional RECIST criteria to mRECIST criteria showed significant difference in response between the two criteria. There was a survival benefit when deemed a responder by the mRECIST criteria [12]. These results confirmed by another study by Shim et al. Conventional RECIST criteria underestimates the response to LRT [13]. Forner et al. demonstrated the use of conventional RECIST criteria missed all the Complete responder and underestimated the partial responders [14].

The use of Modified RECIST criteria in assessing response to molecular therapy for HCC lead to more accurate evaluation of response. In open label study for Brivanib for HCC, Response to treatment according to mRECIST criteria confers higher disease control and subsequently, increased overall survival [15]. In retrospective analysis for Sorafenib therapy, the prognosis of patient categorized as responder was better in the mRECIST group [16].

From our results, we can conclude that through the percentage of necrosis, number of tumors and tumor size, we derived a relative high correlation coefficient and hence a correlated relationship between the radiology and pathology results is implied. We also notice that the correlation coefficient of the total volume is not high enough. The reason for this is mainly due to the lack of the correlation for the 2nd or 3rd tumors sizes, while the total volume variable is considering all the tumors.

The limitation of this study include the extent of necrosis can be affected by interval between CT and liver transplant. In addition, the estimation of pathological necrosis was conducted by experienced pathologist but still represent a subjective measure.

Our results support the notion of dynamic CT as useful tool to estimate viable tumor after locoregional therapy and subsequently response. Similar study of 178 patients, sensitivity and specificity of CT in detecting complete necrosis were 87.5 and 68.9%, respectively [17]. Riaz et al., explored radiologic–pathologic correlation of HCC treated with internal radiation using yttrium-90 microspheres. The finding of the study showed Imaging including dynamic CT demonstrated a high predictive value for necrosis with complete response had a 100% PPV and specificity [18].

In conclusion, among expert radiologists and pathologists, the correlation is acceptable and supports the ongoing use of frequent dynamic imaging to evaluate responses to LRT and determining transplant eligibility.

References


