Serological and histological indices of hepatocellular carcinoma and tumor volume doubling time

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Received June 14, 2013; Accepted August 29, 2013

DOI: 10.3892/mco.2013.186

Abstract. Hepatocellular carcinoma (HCC) frequently develops in cirrhotic liver and is one of the most common malignancies worldwide. The tumor volume doubling time (TVDT) reflects the natural tumor growth rate and is an indicator of the biological malignant potential of a tumor. The present study aimed to elucidate the association between the serological and histological indices of HCC and TVDT. TVDT was analyzed for 53 HCCs by measuring the enlargement of the tumor diameter on dynamic computed tomography. Differences in TVDT were compared among histological grades of HCC differentiation. The α-fetoprotein (AFP) doubling time (T2AFP) for 44 HCCs with AFP levels >200 ng/ml was calculated and the differences in T2AFP were compared according to the histological grade of HCC differentiation and positivity for Lens culinaris agglutinin-reactive fraction of α-fetoprotein (AFP-L3). Among these 44 HCCs, the correlation between T2AFP and TVDT was analyzed for the 27 tumors for which TVDT could be calculated. The mean ± standard deviation (SD) TVDT in Edmondson grade 1 (Ed1), Ed2 and Ed3 HCC was 138.3±110.3, 94.9±91.5 and 32.2±20.8 days, respectively (P<0.01). The mean ± SD T2AFP in Ed2 and Ed3 HCC was 121.0±167.5 and 37.3±24.6 days, respectively (P<0.01). TVDT and T2AFP decreased with histological dedifferentiation of HCC. The mean ± SD T2AFP in the AFP-L3-positive and -negative groups was 63.2±101.2 and 191.9±209.9 days, respectively, with a statistically significant difference between the groups (P<0.01). A significant correlation was observed between T2AFP and TVDT (correlation coefficient, 0.70; P<0.01). A significant correlation was also observed among TVDT, serological indices and histological grades of HCC differentiation. A short T2AFP and/or AFP-L3-positivity were shown to reflect a poorly differentiated HCC histopathology and a higher malignant potential.

Introduction

Hepatocellular carcinoma (HCC) frequently develops in cirrhotic liver and is one of the most common malignancies worldwide. HCC typically develops in a multistep fashion from a low-grade dysplastic nodule to advanced HCC (1). During the course of carcinogenesis, a steady stepwise histological dedifferentiation of the neoplastic cells occurs in conjunction with increasing malignant potential, leading from a well-differentiated HCC to a moderately and finally to a poorly differentiated HCC, with a significantly increased tendency for vascular invasion and metastasis (2). The prognosis of patients worsening with histological dedifferentiation of HCC (3,4).

The tumor volume doubling time (TVDT) reflects the natural tumor growth rate and is an indicator of the biological malignant potential of a tumor. The assessment of TVDT at the time of diagnosis provides a useful parameter for clinical decision making regarding the optimal therapeutic strategy and appropriate follow-up intervals. However, the calculation of TVDT requires values from multiple time points, making this value impossible to determine in numerous cases where treatment is initiated soon after diagnosis. To calculate TVDT, an obvious increase in tumor size must be detected by comparing two subsequent imaging examinations.

The serum α-fetoprotein (AFP) level is one of the most commonly used tumor markers of HCC and is currently used for HCC surveillance and monitoring of recurrence after treatment. The serum levels of AFP and/or Lens culinaris agglutinin-reactive fraction of α-fetoprotein (AFP-L3) are reportedly significantly higher in poorly differentiated HCC compared to those in well- or moderately differentiated HCC (5-7). Although AFP may increase non-specifically during the process of highly active hepatitis, the AFP doubling time (T2AFP) may reflect the tumor growth rate in AFP-producing HCCs. However, the correlation among TVDT, histological grade of differentiation and T2AFP has not been fully elucidated. If T2AFP correlates strongly with TVDT, the malignant potential of AFP-producing HCCs, which is reflected by the natural tumor growth rate, would be easily estimated, even over short intervals.

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Key words: α-fetoprotein, α-fetoprotein doubling time, hepatocellular carcinoma, histological differentiation, tumor volume doubling time
The present study investigated the correlation among histological grade of HCC differentiation, TVDT and T2AFP, as well as between T2AFP and serum AFP-L3 levels, as an indicator of poor prognosis in HCC.

Patients and methods

**Patients.** In order to compare TVDT by histological grade of HCC differentiation, based on our HCC database including patients admitted between April, 2001 and December, 2011, 53 patients with a histological diagnosis of HCC who met the following criteria were retrospectively identified: i) nodular type HCC; ii) availability of baseline and follow-up computed tomography (CT); iii) no treatment during the follow-up period; and iv) measurable enlargement of tumor diameter on dynamic CT during the waiting time prior to treatment.

To calculate T2AFP, based on our HCC database including patients admitted between April, 2001 and December, 2011, 44 HCC patients who met the following criteria were identified: i) serum AFP level measured twice over a 14-day interval; ii) histologically confirmed diagnosis of HCC; iii) baseline serum AFP level >200 ng/ml, to exclude the possibility of non-specific AFP elevation (8); iv) follow-up serum AFP level higher than the baseline serum AFP level; and v) baseline serum AFP-L3 level measured. Furthermore, among the 44 HCC patients in whom T2AFP could be calculated, data from 27 HCC patients for whom TVDT could be calculated were analyzed to evaluate the correlation between T2AFP and TVDT. This retrospective study was approved by the Ethics Committee of Wakayama Medical University. The need for patient consent was waived by the Ethics Committee.

**Histological diagnosis of HCC.** For the evaluation of the histological grade of HCC differentiation, tissue samples were obtained from surgically resected specimens or needle biopsies of the above-described HCC patients. Tumor needle biopsy was performed using an 18-gauge core biopsy needle (Bard Monopty™ no. 121820; Bard Biopsy Systems, Covington, GA, USA) under ultrasound guidance within 1 week prior to treatment. In patients with multiple HCC nodules, the largest nodule, which was presumed to affect prognosis, was assessed. Hematoxylin and eosin staining was performed on all the liver tumor specimens and histology was assessed by two pathologists who were blinded to the clinical course of the patients. Histological grading of HCC was performed according to the criteria described by Edmondson and Steiner (9). When heterogeneous histological grades of differentiation were observed in the obtained tissue sample, the highest grade was adopted. The fibrosis stage of the background liver was assessed using the METAVIR scoring system (10). In case of discrepancies between the evaluations of the two pathologists, a consensus was reached through discussion.

**Dynamic CT.** All contrast CT examinations were performed using multi-detector row CT scanners with at least 4 detectors (Aquilion; Toshiba Medical Systems, Tochigi, Japan, or LightSpeed VCT; GE Medical Systems, Milwaukee, WI, USA), with 5-mm sections. In addition to the plain images, arterial-phase images were obtained 40 sec after the start of bolus administration of 580 mgI/kg of non-ionic iodinated contrast medium for 30 sec via an automated power injector. From 2005 onward, the arterial phase was scanned using an automatic bolus-tracking program. Portal- and equilibrium-phase images were obtained at 70 and 180 sec after the start of bolus administration, respectively.

**Measurement of TVDT.** The maximum tumor diameter was measured in the equilibrium phase of dynamic CT using electronic calipers. The baseline and follow-up CTs of the same patient were evaluated by the same examiner, who was blinded to the clinical patient data. The TVDT was calculated using the following equation (11,12):

\[
TVDT = \left( \frac{T \log2}{3 \log(M1)-\log(M0)} \right)
\]

where M0 is the first-time maximum diameter of the HCC, M1 is the second-time maximum diameter and T is the observation period. The differences in TVDT were compared based on the histological grade of HCC differentiation.

**Measurement of T2AFP.** The T2AFP was calculated using the following equation:

\[
T2AFP = \left( \frac{T \log2}{[\log(\text{AFP1})-\log(\text{AFP0})]} \right)
\]

where AFP0 is the first-time AFP level, AFP1 is the second-time AFP level and T is the observation period.

The AFP-L3 fraction level was measured simultaneously using a liquid-phase binding assay, with a result of <10% being considered as negative and ≥10% as positive. The differences in T2AFP were compared among different histological grades and between AFP-L3-positive and -negative groups. Furthermore, the correlation between T2AFP and TVDT was analyzed.

**Statistical analysis.** The differences in tumor size and TVDT among histological grades of HCC differentiation were compared using the Kruskal-Wallis and Mann-Whitney U tests. The differences in T2AFP among histological grades of HCC differentiation were compared using the Mann-Whitney U test. The difference in T2AFP between AFP-L3-positive and -negative groups was compared using the Mann-Whitney U test. The Pearson correlation coefficient was used to evaluate the association between T2AFP and TVDT. For each statistical test, values of P<0.05 were considered to indicate a statistically significant difference. Data were analyzed using SPSS for Windows, version 20.0J (SPSS Japan Inc., Tokyo, Japan).

**Results**

**Correlation between TVDT and histological grade of HCC differentiation.** To evaluate the grade of histological differentiation in each case, 22 nodules obtained from resected specimens and 31 nodules obtained by tumor biopsy were analyzed. The patient characteristics are summarized in Table 1. The patient sample comprised 35 men and 18 women with a mean ± SD age of 67.2±8.5 years (range, 38-80 years). The mean observation period was 45.7±25.4 days (range, 14-114 days). The mean TVDT was 91.4±93.6 days (range,
Histologically, 13 nodules were classified as Edmondson grade 1 (Ed1), 28 were Ed2 and 12 were Ed3.

The mean tumor diameter ± SD of all the HCCs, Ed1, Ed2 and Ed3 was 25.6±14.3, 20.0±8.0, 30.6±14.8 and 21.5±9.2 mm, respectively, exhibiting no statistically significant differences among different histological grades (P=0.066). No significant correlation was observed between TVDT and tumor diameter (correlation coefficient, -0.06; P=0.668).

The mean TVDT ± SD for Ed1, Ed2 and Ed3 was 138.3±110.3, 94.9±91.5 and 32.2±20.8 days, respectively. The TVDT was significantly longer for Ed1 and Ed2 compared to that for Ed3 (P<0.05), although no significant difference in TVDT was observed between Ed1 and Ed2 (P=0.11) (Fig. 1).

Correlation between T2AFP and histological grade of HCC differentiation.

In order to evaluate the grade of histological differentiation, 17 nodules obtained from resected specimens and 27 nodules obtained by tumor biopsy were analyzed.

The patient characteristics are summarized in Table II. The patient sample comprised 27 men and 17 women with a mean age ± SD of 67.5±8.2 years (range, 48-86 years). No Ed1 HCC exhibited serum baseline AFP level >200 ng/ml. A total of 29 nodules were Ed2 and 15 were Ed3. The mean ± SD T2AFP was 121.0±167.5 and 37.3±24.6 days for Ed2 and Ed3, respectively. There was a statistically significant difference in T2AFP between Ed2 and Ed3 (P<0.01) (Fig. 2).
Correlation between T2AFP and AFP-L3. A total of 34 patients were AFP-L3-positive and 10 were AFP-L3-negative. The mean ± SD T2AFP was 63.2±101.2 days for AFP-L3-positive patients and 191.9±209.9 days for AFP-L3-negative patients. The T2AFP was significantly shorter in the AFP-L3-positive group compared to that in the AFP-L3-negative group (P<0.01) (Fig. 3).

Correlation between T2AFP and TVDT. The T2AFP was calculated for 27 of the 44 cases and an analysis was performed. A strong and significant correlation was identified between T2AFP and TVDT (correlation coefficient, 0.70; P<0.01) (Fig. 4).

Discussion

Previous studies on the growth rate of HCC revealed a wide range of TVDTs, from 13.5 to 2,671 days (13-17). Ebara et al (18) suggested that the growth rate of HCC is not always consistent, even in the same patient, and may change during the course of HCC progression. Sheu et al (19) indicated that TVDT in HCC was positively correlated with initial tumor size. By contrast, other studies demonstrated no significant correlation between TVDT and tumor size (14,15,20,21). A previous study by Kubota et al (15) evaluated small HCCs (<20 mm) and demonstrated that the TVDT ranged widely, from 34.8 to 496.4 days. The present results indicated a similar range of the TVDTs, from 8.7 to 343.9 days. Furthermore, no significant correlation was observed between TVDT and tumor diameter. These findings strongly suggest that rapidly growing HCCs with high-grade malignant potential may be present even among small HCCs.

The wide range of TVDTs in small HCCs may be explained by differences in the histological grade of differentiation in each nodule. Although Sheu et al (19) observed that TVDT was not correlated with histological type or grade of malignancy, a significant correlation between TVDT and histological differentiation was demonstrated by some studies (18,21). Namely, slowly growing HCCs tend to be histologically well-differentiated, whereas rapidly growing HCCs tend to be moderately or poorly differentiated. In the present study, TVDT was significantly shorter for poorly differentiated compared to that for either well- or moderately differentiated HCCs and was the shortest among the three histological grades of HCC differentiation. Thus, the TVDT of each HCC appears to be significantly affected by the histological grade of differentiation.

Ebara et al (18) and Okazaki et al (20) identified TVDT as one of the factors affecting survival in HCC patients. Furthermore, Cucchetti et al (21) reported that TVDT predicts the recurrence of single HCCs after surgical resection. Those results demonstrated that HCCs with vascular invasion, poor or undifferentiation on histological examination and serum AFP levels >200 ng/ml exhibited faster tumor growth, unrelated to tumor size. Thus, the TVDT of HCC reflects the histological grade of differentiation or the grade of malignancy in the process of cancer progression and the assessment of TVDT is considered crucial for deciding the treatment of HCC patients, particularly those with small HCCs treatable by percutaneous local ablation or surgical resection.

However, in order to calculate TVDT, an obvious increase in tumor size must be evident from a comparison of two consecutive imaging examinations. If a close association exists between TVDT and a tumor marker doubling time, that particular tumor marker doubling time may serve as a surrogate for TVDT. Sheu et al (19) reported that, although no significant correlation was observed between TVDT and initial serum AFP levels, TVDT was closely correlated with T2AFP. Our results were consistent with the previous findings. Furthermore, T2AFP was correlated with histological grade of differentiation and the level of AFP-L3, which is another indicator of the grade of malignancy in HCC. Masuda et al (22) reported that the doubling times of preoperative serum AFP or des-γ carboxyprothrombin may be useful predictors of recurrence and prognosis. Those results together with ours, strongly indicate that T2AFP represents a valuable serological index reflecting the malignant potential of HCC. Since the assessment of T2AFP is easier, simpler and faster compared to that of TVDT, T2AFP appears to offer a more useful tool for evaluating tumor growth rates compared to TVDT.

Certain limitations must be considered when interpreting the results of the present study. First, this study on HCC growth rate was small and retrospective. A prospective study would be preferable; however, such a study is not ethically feasible, since
treatment would have to be delayed from the day of diagnosis until confirmation of tumor enlargement. For this reason, the majority of previous studies involved retrospective investigations of a relatively small number of subjects. Second, the histological grade of differentiation was assessed using not only resected specimens, but also tumor biopsy specimens. Since the histological differentiation of HCC is often heterogeneous, the assessment of the histological grade of differentiation may not reflect the least differentiated histology in the tumor. However, the present study strongly demonstrated that TVDT was significantly correlated with histological grade of differentiation, consistent with previous studies (18,21). Third, since the tumor size of massive or diffuse-type HCC cannot be limited to nodular-type HCC. Thus, the present results cannot be generalized, although the possibility that T2AFP reflects tumor growth rates even in massive or diffuse-type HCC requires further investigation. Fourth, T2AFP cannot be used to assess the tumor growth rate when HCC is not detected on imaging. In the present study, T2AFP was analyzed only in HCC patients with baseline serum AFP levels of >200 ng/ml that remained higher thereafter, to allow the exclusion of non-specific AFP elevations (8). However, patients with AFP >200 ng/ml without imaging evidence of HCC may be a frequent finding. Furthermore, the assessment of tumor growth rate using T2AFP is not possible for non-AFP-producing HCCs. Furthermore, tumor growth rates cannot be assessed when AFP-producing activity is low and T2AFP is long. However, since the HCCs for which T2AFP could be assessed in the present study were all moderately or poorly differentiated, T2AFP may be considered useful for identifying HCCs with a high-grade malignant potential. Finally, the interval used for T2AFP measurements in the present analysis varied greatly (range, 14-113 days). It was previously reported that the growth rate of HCC is not always consistent and the increases in the serum AFP levels are also not always linear (18). In addition, dedifferentiation of HCC is often observed during growth. Thus, the optimal interval between two AFP measurements has not been determined.

In conclusion, a significant correlation was observed among serological indices, histological grade of differentiation and TVDT, but not tumor size. The TVDT and T2AFP were shorter with advancing histological grade of differentiation, or grade of malignancy. The T2AFP was shorter in the AFP-L3-positive compared to that in the AFP-L3-negative group. A strong correlation was observed between TVDT and T2AFP; therefore, when deciding the treatment for HCC patients, attention should be paid to a short T2AFP and/or AFP-L3 positivity, since poorly differentiated HCC is more likely to exhibit microscopic vascular invasion or intrahepatic metastases.

Acknowledgements

The authors would like to thank Ms. Kazu Konishi for her excellent secretarial assistance. This study was supported in part by a grant-in-aid from Takeda Pharmaceutical Company Limited.

References


