Characterization of Hepatic Focal Lesions by Diffusion Tensor Imaging and How Far it Can Predict Post-Treatment Response?

NEHAD M.S. FOUDA, M.D.; KARIM M. ABDELHAMID, M.SC. and NEHAL THARWAT, M.D.
The Department of Diagnostic Radiology, Faculty of Medicine, Mansoura University, Egypt

Abstract

Background: Liver diseases have been known to be a major health problem principally because of their world-wide distribution. Focal liver disease is a common diagnostic problem referred to radiologists for evaluation owing to its nonspecific clinical presentation and marked interobserver variation on clinical examination.

Aim of Study: The purpose of this study is to evaluate the role of diffusion tensor imaging in characterization of hepatic focal lesions and its value in post-treatment response.

Patients and Methods: This prospective study included 30 patients previously diagnosed to have hepatic focal lesions (HFLs) by their characteristic triphasic computed tomography (CT), dynamic magnetic resonance imaging (MRI) features and/or biopsy. The study included 43 lesions in 30 patients (16 benign lesions, 23 malignant lesions, and 4 treated malignant lesions). MRI with diffusion tensor imaging (DTI) was performed for all patients. Apparent diffusion coefficient (ADC) and fractional anisotropy (FA) values were evaluated for all lesions.

Results: There was a statistically significant difference in ADC values between the benign and malignant lesions ($p<0.001$) and between the treated malignant and untreated malignant lesions ($p=0.002$). There was a statistically significant difference in FA values between the benign and malignant lesions ($p<0.001$) and between the treated malignant and untreated malignant lesions ($p=0.004$). The best cut-off ADC value to differentiate between benign and malignant lesions respectively was $1.42 \times 10^{-3}$ mm$^2$/s with 95.7% sensitivity and 82.8% specificity. The best cut-off ADC value to differentiate between treated malignant and malignant lesions respectively was $1.65 \times 10^{-3}$ mm$^2$/s with 97.8% sensitivity and 95.7% specificity. The best cut-off FA value to differentiate between benign and malignant lesions respectively was 0.29 with 95% sensitivity and 70% specificity. The best cut-off FA value to differentiate between treated malignant and untreated malignant lesions respectively was 0.297 with 100% sensitivity and 69.2% specificity.

Conclusion: Diffusion tensor imaging is an evolving technique that can be used to characterize different hepatic focal lesions either benign or malignant with significant additive value to dynamic contrast enhanced MRI examination. It can also be used to monitor treatment response.

Key Words: Diffusion tensor imaging – Hepatic focal lesions – Benign – Malignant.

Introduction

LIVER diseases have been known to be a major health problem principally because of their world-wide distribution [1,2]. Focal liver disease is a common diagnostic problem referred to radiologists for evaluation owing to its nonspecific clinical presentation and marked interobserver variation on clinical examination [3].

Hepatic focal lesions (HFLs) are classified into benign and malignant lesions. Hemangiomas are the commonest benign tumor while hepatocellular carcinoma (HCC) it is the commonest primary malignant liver tumor. HCC is the fifth most common cancer in the world and the third most frequent cause of death amongst oncological patients [4].

Modern operative techniques and local therapies such as radiofrequency (RF) ablation are effective methods to treat primary hepatic malignancies or liver metastases. Therefore, accurate determination

Abbreviations:

HFLs : Hepatic focal lesions.
CT : Computed tomography.
MRI : Magnetic resonance imaging.
DTI : Diffusion tensor imaging.
ADC : Apparent diffusion coefficient.
FA : Fractional anisotropy.
HCC : Hepatocellular carcinoma.
RF : Radiofrequency.
USG : Ultrasonography.
DWI : Diffusion-weighted imaging.
TACE : Trans-arterial chemoembolization.
PPV : Positive predictive value.
NPV : Negative predictive value.
of liver lesion count, and nature of the lesion are important [5].

HFLs are diagnosed using ultrasonography (USG) and/or computed tomography (CT). Triphasic CT has traditionally been considered the optimal diagnostic modality for HFLs. However, many limitations have been reported concerning triphasic CT study such as renal impairment, radiation dose, and inability to confirm the specific tissue properties of focal lesions in some cases, leading to indeterminate diagnosis. Conventional and dynamic contrast-enhanced magnetic resonance imaging (MRI) are ideal tools for obtaining anatomical details but cannot provide functional details for HFLs [6-9].

Diffusion-weighted imaging (DWI) relates to the diffusion properties of water molecules and reveals the histopathological tissue characteristics. Malignant tissues with high cellularity, constriction of extracellular spaces and high density of hydrophobic membranes have usually more restricted diffusion making apparent diffusion coefficient (ADC) values lower [10,11].

DWI provides limited information and does not provide information on tissue microstructure such as diffusion anisotropy. Diffusion tensor imaging (DTI) is MRI technique that reveals microstructural characteristics of biological tissue, which can detect the degree of diffusion in multiple directions by using additional gradients [12]. Compared to three gradient-directions applied to DWI, at least six or more gradient directions for every section in DTI are needed to calculate the diffusion tensor providing additional information on anisotropy diffusion and total diffusion orientations [13].

The literature is especially sparse regarding the usage of DTI in characterization of different HFLs. DTI can achieve more precise ADC calculation. In addition, fractional anisotropy (FA) values obtained by DTI are useful in evaluating the scalar properties of the diffusion of extracellular water molecules. DTI has been widely applied in the brain. Recently, application of DTI in the liver for diagnosis and staging of fibrosis and inflammation, and distinguishing cyst,malignancy, and hemangioma has shown reasonably good success [14-17].

DTI does not require any exogenous contrast agent and can be safely performed in patients presenting contraindications for gadolinium contrast agents (patients suffering from severe renal deficiency or nephrogenic systemic fibrosis). Early clinical applications of DTI demonstrated the sensitivity of this technique to hepatic lesions compared to standard MRI especially in metastases and lesions of atypical enhancing pattern. Also, DTI might be effectively employed to determine the biopsy target [18].

Recently, a multiparametric MRI approach including dynamic contrast-enhanced MRI and other functional imaging tools such as DWI, MR elastography, and MR spectroscopy, has drawn a lot of attention as it allows not only morphologic evaluation but also functional evaluation of various liver diseases. This can help in maximizing specificity and accuracy of cross-sectional imaging and avoid unnecessary biopsies, which may portend a postprocedural morbidity of 2.0% to 4.8% and mortality of 0.05% [19].

This study tried to introduce FA values (in addition to ADC) as a new biomarker that can be used in the diagnostic work-up of liver lesions.

The aim of this study is to evaluate the role of DTI in characterization of HFLs previously diagnosed with their characteristic triphasic CT, dynamic MRI features and/or biopsy and its value in post treatment response.

Patients and Methods

This prospective single-institution study was approved by our institutional review board and written informed consent from all patients was obtained. This study was conducted during the period from January 2018 to January 2020. Thirty patients (16 males and 14 females; mean age 51.8 years) who were referred from oncology center and tropical medicine unit with HFLs diagnosed by their characteristic triphasic CT, dynamic MRI features and/or biopsy were enrolled in this study.

There were 21 cases with single lesion, 5 cases with 2 lesions and 4 cases with 3 lesions with total of 43 lesions. Among the 43 lesions there were 16 benign lesions (37.2%), 23 malignant lesions (53.5%) and 4 completely treated malignant lesions (HCC) (9.3%). Regarding the type of the detected lesions, there were 12 haemangiomas (27.9%), 3 hydatid abscesses (7%), 1 simple cyst (2.3%), 12 HCC (27.9%), 7 metastases (16.3%), 2 cholangiocarcinomas (4.7%), 1 dysplastic nodule (2.3%), 1 epithelioid haemangioendothelioma (2.3%) and 4 HCC treated by trans-arterial chemoembolization (TACE) and showed complete response (9.3%).

All patients underwent MRI of the liver with DTI technique. We evaluated ADC and FA values of all lesions.
**Inclusion criteria:** Patients diagnosed to have HFLs by different imaging modalities and characterized by triphasic CT or dynamic MRI and/or biopsy (if needed) and agreed to participate in the study.

**The gold standard diagnostic criteria of HFLs:**

- **Haemangioma:** Hyperintensity on heavy T2-WI and the typical enhancement pattern seen in contrast-enhanced dynamic CT or MRI in the form of early peripheral nodular enhancement in arterial phase with centripetal progressive enhancement in portal and delayed phases [20].

- **Hydatid abscesses:** Unilocular or multilocular cysts with thin or thick walls and calcifications, usually with daughter cysts. On MRI, hydatid cyst appears low on T1-WI and high on T2-WI. Intraluminal debris presence may alter the signal intensity. Both fibrous capsule and internal septa tend to be hypointense on T2-WI and show enhancement in post gadolinium phase [4,21].

- **Simple cyst:** Diagnosed according to their typical US, CT and MRI findings (thin smooth walls with no mural irregularity or nodularity or debris, hypodense non-enhancing on CT, non-enhancing on MRI with low T1 and high T2 signal intensity) accompanied with follow-up evaluations of at least 12 months [17,22].

- **HCC:** On multiphasic contrast CT or MRI, diagnosed by arterial phase hyperenhancement, subsequent washout appearance in portal and delayed phases and delayed enhancing pseudocapsule. Additionally, HCC have the propensity to invade vascular structures, most commonly the portal vein [23].

- **Metastases:** The diagnosis of metastases was proven by means of biopsy or follow-up imaging examinations including CT and MRI during routine controls of patients with known primary tumors [17].

- **Intrahepatic mass forming cholangiocarcinoma:** diagnosed on dynamic post-contrast scan by minor peripheral rim enhancement during both the arterial and portal venous phases, the central part frequently show gradual centripetal prolonged enhancement at delayed-phases with peripheral intrahepatic duct dilatation. Capsular retraction is highly suggestive of cholangiocarcinoma. Unlike HCC, cholangiocarcinoma only rarely forms a tumor thrombus. Also, cholangiocarcinomaappears hyperintense on T2-WI and central hypointensity in the tumor reflective of fibrosis may be seen [21,24].

- **Dysplastic nodule:** High-grade dysplastic nodules show iso to high signal on T1-WI and iso signal intensity on T2-WI. On dynamic post contrast study, high-grade nodules show early contrast enhancement in arterial phase and fade to isodensity or isointensity, but without washout on delayed phase (unlike HCC) [4].

- **Epithelioid haemangioendothelioma:** Typically seen as multiple lesions in a peripheral or subcapsular distribution, with a peripheral halo or target pattern of enhancement. Hepatic or portal veins or their branches may taper and terminate at or just within the edge of these lesions (lollipop sign) [28].

- **HCC treated by TACE:** The most important finding for a completely treated HCC is lack of internal enhancement. Treated masses usually demonstrate low signal intensity on T1 and T2-WI unless there is hemorrhagic or proteinaceous debris, in which case there is high signal intensity on T1-WI. In such instances, subtraction imaging help to identify subtle areas of arterial hyperenhancement indicative of viable tumor [26].

**Exclusion criteria:** General contra-indication for MRI scan (cardiac pacemaker, metallic implant), bad general condition and uncooperative patients who were unstable on machine table or cannot hold their breath.

**MR imaging technique:**

MRI was performed on high field system (1.5 Tesla) magnet units (Philips Ingenia) using a phased array coil to cover the whole liver. Patients were asked to avoid deep breathing during examination. Conventional MRI and DTI studies were performed. First; detection and localization of focal lesions were performed; second, the DTI with ADC and FA maps were performed.

**MR Protocol:**

- T1WI (TR=112 msec, TE=4.8msec, matrix 179x320, slice thickness 7-8mm, slice gap 1-2mm and FOV 300-400mm).
- T2WI (TR 1800 msec, TE=80msec, matrix 200x240, slice thickness 7-8mm, slice gap 1-2mm and FOV 300-400mm).
- T2 SPAIR (Spectral Attenuated Inversion Recovery) fatsuppression sequence: TR > 400msec, TE=80msec, matrix 204x384, slice thickness 7-8mm, slice gap 1-2mm and FOV300-400mm.

**Diffusion tensor study:**

An axial non-breathhold, single-shot gradient echo planar DTI sequence covering the whole liver was acquired using the following parameters: Acquisition time, 3:04 (min:sec), TE=94msec, TR > 2371 msec, matrix 256x160, slice thickness 8mm, slice gap 1mm, with different $b$-values ($b=0, 500,
800 & 1000s/mm²) with six diffusion directions applied.

**Image analysis:** ADC and FA maps were processed using secondary work station provided by the vendor (Phillips Advantage windows workstation with functional tool software). Analysis of ADC and FA values of HFLs was done by an expert radiologist (15 years’ experience in hepatic imaging, 8 years’ experience in DTI analysis) by applying regions of interest (ROIs) in each lesion on ADC and FA maps. In cystic lesions, ROIs were placed within the margins of the lesions.

**Statistical analysis of data:**

The collected data were coded, processed and analyzed using the SPSS (Statistical Package for Social Sciences) version 22 for Windows® (IBM SPSS Inc, Chicago, IL, USA). Data were tested for normal distribution using the Shapiro Walk test. Qualitative data were represented as frequencies and relative percentages. Quantitative data were expressed as median (minimum & maximum). Kruskal Wallis test was used to compare non parametric data between more than 2 groups with paired comparison by Mann Whitney U test. Receiver operator characteristic (ROC) curve was tested to calculate the diagnostic ability of quantitative variable in prediction of categorical outcome. Significance test results are quoted as two-tailed probabilities. For all the above-mentioned tests, the level of significance was tested, expressed as the probability of (p-value) and the results were considered significant if the p-value is ≤0.05 and highly significant if the p-value <0.001.

**Results**

This study included 30 patients with mean age of 51.8±12.7 years and age range between 23 and 73 years. Among the patients there were 16 males (53.3%) and 14 females (46.7%).

The mean ADC value of the different lesions in our study was 1.60±0.63x 10⁻³ mm²/s and the median value was 1.35x 10⁻³ mm²/s with range between 0.87 and 3.63x 10⁻³ mm²/s. The mean value of FA of the different lesions in our study was 0.33±0.14 and the median value was 0.33 with range between 0.08 and 0.74.

The median and minimum-maximum ADC and FA values of benign, malignant, and treated malignant (HCC) lesions are shown in Table (1). There was a statistically significant difference in ADC value between the benign and malignant lesions (p<0.001), between the treated HCC and untreated malignant lesions (p=0.002), but there was no statistically significant difference noted between the benign lesions and treated HCC (p=0.705). Also, there was a statistically significant difference in FA values between the benign and malignant lesions (p<0.001) and between treated HCC and untreated malignant lesions (p=0.004), but there was no statistically significant difference between the benign lesions and treated HCC (p=0.345) (Table 1) (Figs. 1,2).

<table>
<thead>
<tr>
<th>ADC</th>
<th>Benign (n=16)</th>
<th>Malignant (n=23)</th>
<th>Treated HCC (n=4)</th>
<th>Test of significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>2.152</td>
<td>1.212</td>
<td>2.024</td>
<td>KW $\chi^2=25.65$</td>
</tr>
<tr>
<td>Minimum-Maximum</td>
<td>1.21-3.63</td>
<td>0.87-1.46</td>
<td>1.86-2.17</td>
<td>$p&lt;0.001^*$</td>
</tr>
<tr>
<td>$p_1$</td>
<td>&lt;0.001*</td>
<td>0.705</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$p_2$</td>
<td>0.002*</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>FA</th>
<th>Benign (n=16)</th>
<th>Malignant (n=23)</th>
<th>Treated HCC (n=4)</th>
<th>Test of significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>0.211</td>
<td>0.381</td>
<td>0.2815</td>
<td>KW $\chi^2=16.34$</td>
</tr>
<tr>
<td>Minimum-Maximum</td>
<td>0.08-0.43</td>
<td>0.17-0.74</td>
<td>0.25-0.29</td>
<td>$p&lt;0.001^*$</td>
</tr>
<tr>
<td>$p_1$</td>
<td>&lt;0.001*</td>
<td>0.345</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$p_2$</td>
<td>0.004*</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

p: Probability. $p_1$: Statistical significant ($p<0.05$). $p_2$: Significance in relation to benign group. 

*: Statistically significant ($p<0.05$). **: Overall significance ($p<0.001$).
In our study, the best cut-off ADC value to differentiate between benign and malignant lesions respectively was $>1.42 \times 10^{-3}$ mm$^2$/s with 95.7% sensitivity, 82.8% specificity, 96.4% positive predictive value (PPV), 94.3% negative predictive value (NPV) and total accuracy of 88.6%. This value was considered statistically significant ($p=0.001$) (Fig. 3). The best cut-off ADC value to differentiate between treated HCC and untreated malignant lesions respectively was $>1.65 \times 10^{-3}$ mm$^2$/s with 97.8% sensitivity, 95.7% specificity, 96.4% PPV, 94.3% NPV and total accuracy of 95.2%. This value was considered statistically significant ($p<0.001$).

In our study, the best cut-off FA value to differentiate between malignant and benign lesions respectively was $>0.29$ with 95% sensitivity, 70% specificity, 92% PPV, 82% NPV and total accuracy of 85%. This value was considered statistically significant ($p=0.001$) (Fig. 4). The best cut-off FA value to differentiate between treated malignant and untreated malignant lesions respectively was
<0.297 with 100% sensitivity, 69.2% specificity, 80% PPV, 95.7% NPV and total accuracy of 96.3%.

Yet, this value was considered statistically non-significant ($p=0.181$).

Fig. (2): Female patient 56 years old with cirrhotic liver secondary to chronic hepatitis C infection diagnosed to have HCC in segment VII on dynamic MRI displaying low T1-SI (A) and relatively high T2-SI (B) with heterogeneous enhancement in arterial phase (C) with wash-out in portal and delayed phases (D). DTI was performed, the lesion shows ADC value of $0.8 \times 10^{-3}$ mm$^2$/s and FA value of 0.39 suggestive of malignant nature (E, F) show ADC and FA maps respectively.

Fig. (3): ROC curve for ADC cut-off value in differentiation between benign and malignant lesions.

Fig. (4): ROC curve for cut-off FA value in differentiation between benign and malignant lesions.
Discussion

MRI is a preferred technique when further characterization of HFLs is needed. Lesion morphology, signal intensity, and contrast enhancement pattern are taken into consideration when characterizing masses with MRI. Yet, there can still be difficulties in the differentiation of benign and malignant lesions [27]. DW-MRI has proven a great and growing role in diagnosing hepatic pathology compared to other MRI techniques. Indeed, DW-MRI and quantitative ADC measurements is a fast and completely non-invasive technique that does not require contrast agent administration with possibility of better differentiation between benign and malignant lesions [28,29].

By using DTI, it is possible to obtain not only the ADC values but also the FA values that may also provide useful information regarding HFLs. The hypothesis is that the diffusion in malignant lesions should be more restricted and more anisotropic when compared to benign lesions resulting in outcomes as lower ADC values and higher FA values for malignant lesions [17].

In our study the most common malignant lesion was HCC (27.9%), the most common benign lesion was hemangioma (27.9%). This came in agreement with Abdel Kader et al., who declared that out of 312 lesions in his study, HCC was the most common malignant lesion (171/312), hemangioma was the commonest benign lesion (65/312) [30]. Also, nearly similar results were obtained in the study done by Debees et al., as they stated that among studied 27 malignant lesions; HCC came first (15/27) [31].

In this study, the median ADC value of benign lesions was 2.152 x 10^{-3} mm²/s, for malignant lesions 1.212 x 10^{-3} mm²/s and for treated HCC was 2.024 x 10^{-3} mm²/s. There was a statistically significant difference in ADC value between the benign and malignant lesions ($p < 0.001$), between the treated HCC and untreated malignant lesions ($p = 0.002$), but there was no statistically significant difference noted between the benign and treated HCC lesions ($p = 0.705$).

Our study came in agreement with Javadrashid et al., who found that the mean ADC value for benign lesions ($1.58 \pm 0.35 \times 10^{-3} \text{mm}^2/\text{s}$) was significantly higher than malignant lesions ($0.87 \pm 0.16 \times 10^{-3} \text{mm}^2/\text{s}$) with ($p = 0.001$) [32]. Also, Jain et al., stated that the mean ADC value for benign lesions was $1.678 \times 10^{-3} \text{mm}^2/\text{s}$, and for malignant lesions was $1.097 \times 10^{-3} \text{mm}^2/\text{s}$, with statistically significant difference in-between ($p < 0.001$) [33]. El-Refaei and colleagues found high level of mean ADC in simple cyst and hemangioma more than that of metastasis and HCC, with highly statistically significant difference ($p = 0.001$) in mean ADC between benign and malignant lesions [7]. This difference in ADC values or cut-off ADC values between mentioned studies may be due to the use of variable $b$-values as diffusion gradients.

Our study has reported that there was statistically significant difference in the median ADC value between the treated malignant (HCC) and untreated malignant lesions ($p = 0.002$), but there was no statistically significant difference noted between the benign and treated malignant lesions ($p = 0.705$).

Increased ADC values in tumours after treatment generally show positive correlation with tumour response. The increase of ADC values as a response to treatment occurs earlier than size change in focal hepatic tumours. Correlation between ADC values and tumour response was mainly studied with colorectal metastases treated with chemotherapy and HCCs treated with chemoembolization therapy [34].

This came in agreement with Abduljaleel et al., who showed that ablation zones can be differentiated from surrounding liver parenchyma visually in the DWI and by means of ADC maps in all patients. They reported that the mean ADC of the well ablated lesion was $1.4 \times 10^{-3} \text{mm}^2/\text{s}$ and of the residual lesion $0.8 \times 10^{-3} \text{mm}^2/\text{s}$ with significant statistical difference between the residual viable tumor and the well ablated lesions [35].

This also came in accordance with Tantawy and Mohamed who found that ADC values were significantly higher in lesions that responded to TACE or radio frequency ablation (RFA) than in non-responding lesions. The mean ADC of the lesions before treatment was $1.27 \pm 0.25 \times 10^{-3} \text{mm}^2/\text{s}$, and increased after treatment in responding lesions to reach $1.57 \pm 0.22 \times 10^{-3} \text{mm}^2/\text{s}$ with a statistically significant difference ($p = 0.002$) [34].

This also agreed with Lo et al., who found that an ADC change of $\geq 25\%$ within 6 months post-stereotactic ablative radiotherapy (SABR) was an independent predictor of sustained HCC tumor control. The ADC values pre- and post-SABR were $1.43 \pm 0.28 \times 10^{-3} \text{mm}^2/\text{s}$ and $1.72 \pm 0.34 \times 10^{-3} \text{mm}^2/\text{s}$ respectively ($p < 0.001$) [36]. This indicated that the ADC values could be established as a monitoring tool to assess the response of malignant lesions for treatment.
In this study, the best cut-off ADC value to differentiate between benign and malignant lesions respectively was >1.42 x 10^{-3} mm^2/s with 95.7% sensitivity, 82.8% specificity, 96.4% PPV, 94.3% NPV and total accuracy of 88.6%.

This came in agreement with Madhu et al., who found that the best ADC cut-off value to differentiate between benign and malignant HFLs was 1.431 x 10^{-3} mm^2/swith sensitivity of 87.5%, specificity of 71%, PPV of 79.5% and NPV of 81.5% [37]. Also, Jain et al., defined an ADC value of 1.26 x 10^{-3} mm^2/s to be the best available cut-off value for differentiating benign and malignant lesions, achieving sensitivity and specificity of 92% and 80%, respectively [33].

Within the same context, Hasan et al., reported that by using ADC cut-off of 1.6 x 10^{-3} mm^2/s led to the highest accuracy for the differentiation of malignant and benign liver lesions (86%) with a sensitivity of 100% and specificity of 68% for malignant lesions. Its strength was in its 100% NPV where ADC values above 1.6 x 10^{-3} mm^2/s exclude the malignant lesions [38].

However, ADC fails to define the characteristics of diffusion in anisotropic environments. Anisotropic properties of tissues can be evaluated using DTI, which allows the analysis of diffusion along multiple directions by employing additional gradients. By using DTI, in addition to ADC values, FA values can be calculated. FA values show the fraction of anisotropic diffusion to total diffusion. Higher FA indicated higher tumor cell density and higher malignant potential [15,17].

In this study, the median FA value of the benign lesions was 0.211, for malignant lesions 0.381 and for treated malignant lesions (HCC) was 0.2815. There was a statistically significant difference in FA values between the benign and malignant lesions (p<0.001), and between treated malignant and untreated malignant lesions (p=0.004), but there was no statistically significant difference between the benign and treated malignant lesions (p=0.345).

This came in agreement with Li et al., who found that the mean FA value of HCC lesions (0.42 ±0.11) was significantly higher than that of normal liver parenchyma (0.32 ±0.10) (p=0.004) [15]. This also came in accordance with Erturk et al., as they showed that the mean FA values of cysts, hemangiomas, and metastases were 0.2 ±0.05, 0.37 ±0.1 and 0.46±0.1, respectively. The differences in FA values of cysts and metastases and of cysts and hemangiomas were statistically significant (p<0.01). On the other hand, the difference between metastases and hemangiomas was not significant (p=0.88) [17].

In this study, the best cut-off FA value to differentiate between malignant and benign lesions respectively was >0.29 with 95% sensitivity, 70% specificity, 92% PPV, 82% NPV and total accuracy of 85%. This value was considered statistically significant (p=0.001). Similar results were obtained by Erturk et al. who reported that in distinguishing metastases (malignant lesions) from cysts and hemangiomas (benign lesions) using FA value of 0.31 as the cut-off value, the sensitivity was 56.2% and the specificity was 80% [17].

As regard usage of FA values to monitor the response of malignant lesions for treatment, there was lack of researches assessing FA changes in patients with malignant HFLs after treatment. However, Abdel Razek et al., in a study about breast cancer found higher FA values in tumor recurrence than post-operative changes in patients with breast conserving surgery with statistically significant difference (p=0.003) that matched with our results [39]. On the other hand, D’Arco et al., in a study concerning pediatric brain tumour response stated that anisotropy within the matrix of tumour tissue is initially low making the evaluation or follow-up of diffusion anisotropy within tumours after treatment of little benefit [40].

The limitation of this study was the small number of patients and treated malignant lesions included in the study.

Conclusion:

MRI with DTI is an effective non-invasive tool with significant additive value to dynamic contrast enhanced MRI examination in detection and characterization of different HFLs due to its ease of acquisition and ability to obtain functional information in absence of intravenous contrast, especially in patients with abnormal renal function. It also may help in characterization of lesions with non-specific enhancing pattern. It can be used as a monitoring tool for assessment of treatment response in malignant hepatic lesions. Further studies should be performed including larger number of patients from more than one center.

References


توصيف الآفات البؤرية الكبدية باستخدام خاصية الانتشار الموتر
والإي م 의해 يمكنها التنبؤ بالاستجابة بعد العلاج؟

الهدف من الدراسة: تقييم دور الرين المغناطيسي بخارية الانتشار الموتر في توصيف الآفات البؤرية الكبدية وتقييم مدى استجابتها للعلاج.

المراقب: وفرت البحث: شملت هذه الدراسة المرتبطة 30 مريضاً تم تشخيصهم سابقاً بوجود آفات بؤرية كبدية من خلال التصوير بالأشعة المقطعية ثلاثية المراحل أو الرين المغناطيسي الدينيمائي أو الفرعية. وشملت الدراسة 33 مريضاً (16 مريضاً (3 أفق حميدة، و 2 أفق خبيثة، و 4 أفق خبيثة عالية). تم إجراء التصوير بالزين المغناطيسي مع خاصية الانتشار الموتر لجميع المرضى وتقييم معدل الانتشار الظاهري بالإضافة إلى معدل تفاوت الانتشار لجميع الآفات.

نتائج البحث: كان هناك فرق معقد بخصوصي في قيمة معدل الانتشار الظاهري بين الآفات الحميدة والخبيثة وبين الآفات الخبيثة والخبيثة المعالجة. كان هناك فرق معقد بخصوصي في قيمة معدل تفاوت الانتشار الظاهري بين الآفات الحميدة والخبيثة بنسبة 82.8% وخصوصية 95.7%. كانت أفضل قيمة فاصلة لقيم معدل تفاوت الانتشار الظاهري عند عالمات إجمالية وخصوبة 95% وخصوصية 100%. كانت أفضل قيمة فاصلة لقيم معدل تفاوت الانتشار الظاهري عند عالمات إجمالية وخصوبة 97.9% وخصوصية 100%.

الاستنتاج: تصور الرين المغناطيسي بخارية الانتشار الموتر هو تقنية متطورة يمكن استخدامها لتوصيف الآفات البؤرية الكبدية المختلفة إما حميدة أو خبيثة ويعتبر ذات قيمة مضافة كبيرة إلى جانب الرين المغناطيسي الدينيمائي ويمكن أيضاً أن يستخدم لرصد مدى الاستجابة للعلاج.