ABSTRACT

Material and methods: The study was of prospective-retrospective character. It was carried out at the AKH in Vienna (Austria), where 100 patients with focal liver lesions were included in the study. All patients underwent the routine MR sequences on appliances 1.5 and 3T (Siemens, Germany): T1, T2, HASTE, VIBE, and a DWI with three b values (b 50, b 300, b 600 s / mm²) and ADC map with ROI (regions of interest). The numerical value of ADC map was calculated, where n = 100 liver lesions, by two independent radiologists. Results: On the basis of matching the PH finding statistically we get DWI accuracy of 96.8% for the assessment of liver lesions. The average numerical value of ADC in benign hepatic lesions (FNH, Hemangiomas) in our study amounted to 1.88 (1.326 to 2.48) x10⁻³ mm²/s, while the value of malignant liver lesions (HCC, CCC, CRCLM) were significantly lower and amounted to 1.15 (1.024 to 1.343) x10⁻³ mm²/s (Figure 2). Differences between the mean ADC of benign and malignant lesions showed a statistically significant difference with p <0.0005. In our research, we get cut-off for the ADC value of 1.341 x10⁻³ mm²/s, which proved to be the optimal parameter for differentiation between benign and malignant lesions. Conclusion: Measuring ADC values with DWI as an additional MRI tool can help in oncological practice by distinguishing normal liver parenchyma from focal lesions, and in differentiating benign from malignant liver lesions, particularly in cases where administration of contrast is not possible.

Key words: diffusion weighted imaging, apparent diffusion coefficient, focal liver lesions.

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Excel (version 11th Microsoft Corporation, Redmond, WA, USA) were used.

3. RESULTS

Gender structure of the respondents in our study was balanced: 52% of men develop compared to 48% of women, with no statistically significant difference ($\chi^2 = 0.182$, $p = 0.670$). The youngest patient was 19 and the oldest 82 years.

Kendall’s tau-b coefficient of concordance showed a statistically significant correlation ($p<0.0005$) between MRI DWI findings of focal liver lesions and PH findings of focal liver lesions. PH findings confirmed MRI DWI diagnosis by: hemangioma in 11/12 (92%), by FHN (focal nodular hyperplasia) in 6/10 (60%), in patients with metastatic colorectal cancers (CRC) in 1/1 (100%), the cholangiocellular carcinoma (CCC) in 12/12 (100%) and hepatocellular carcinoma (HCC) in 20/22 (91%) patients (Figure 1). On the basis of matching the PH findings statistically we got DWI accuracy of 96.8% for the assessment of liver lesions. Single-shot T2-weighted fast SE and DWI MRI ($b$ 50, $b$ 300 $b$ 600 s / mm$^2$) were compared by two independent radiologists (A and B) in the assessment of liver lesions in which the Kappa coefficient (Measure of Agreement Kappa = 0.864) showed a statistically significant approval ($p<0.0005$). But comparing a single-shot T2-weighted fast SE and DWI by one radiologist (A) in the assessment of liver lesions Kendall’s tau-b coefficient of agreement (matching) did not show statistically significant matching ($b = 0.119$, $p = 0.437$). This is manifested in favor of detection by DWI, which detected more liver lesions, of total $n = 100$ liver lesions $n = 18$ were not detected on T2 w, which were confirmed by PH. From a total of $n = 18$ missed hepatic lesions that were not detected in T2 sequences, DWI detected $n = 17$ (94.4%). From a total of $n = 82$ detected focal liver lesions on T2w number, $n = 81$ (96.8%) was detected on DWI. The largest number of focal lesions $8/18$ (44%) that were not detected on the T2 sequences, the PH verified as HCC, 4/18 (22%) as FHN, 3/18 (17%) as meta CRC, 2/18 (11%) hemangioma and 1/18 (5%) CCC. (Figure 1)

Figure 1. Frequency of changes that were detected on DVI, but not single-shot T2-weighted fast SE

By measuring the ADC value of each lesion individually we got different results between different types of lesions. The average numerical value of ADC in benign hepatic lesions (FHN, Hemangiomas) in our study amounted to 1.88 (1.326 to 2.48) x10$^{-3}$ mm$^2$/s, while the value of malignant liver lesions (HCC, CCC, CRCLM) were significantly lower and amounted to 1.15 (1.024 to 1.343) x10$^{-3}$ mm$^2$/s (Figure 2)

Figure 2. The difference of ADC average values of benign and malignant lesions of the liver. Average values of ADC for benign lesions amounted to 1.88 (1.326 to 2.48)x10$^{-3}$ mm$^2$/s. Average values of ADC for malignant lesions amounted to 1.15 (1.024 to 1.343)x10$^{-3}$ mm$^2$/s

Figure 3. Metastatic lesion of CRC: a) ADC:1,1 x10$^{-3}$ mm$^2$/s; b) MR DWI black blood-b value=50sec/mm2; c) ADC : FHN in the left lobe of the liver 1,3 x 10$^{-3}$ mm$^2$/s

2). Differences between the mean ADC of benign and malignant lesions showed a statistically significant difference with $p<0.0005$. In our research, we get cut-off for the ADC value of 1,341x10$^{-3}$mm$^2$/s, which proved to be the optimal parameter for differentiation between benign and malignant lesions.

4. DISCUSSION

Diffusion figures (average image between images obtained with three diffusion gradient coefficient) is displayed for each b value together with ADC folder. Image with high b value determines whether areas of limited diffusion, which appear as high intensity. Visual estimation of DWI images, including those with high b values (greater than 500sec / mm$^2$), can help in distinction between solid and cystic lesions. Both benign and malignant changes can maintain a high signal intensity with increasing b values, which makes characterization only on the basis of diffusion images difficult. Visual assessment of only diffusion image can lead to false positives because most of benign lesions will appear bright on T2 weighted images and partial volume effect of other structures, which occurs in the cellular benign lesions such as focal nodal hyperplasia-FNH, adenoma, abscess. False negative results can occur in the metastatic mucus–producing tumor, which can mimic cystic lesions, necrotic primary or secondary tumor processes, and image artifacts that may obscure the lesion (4). Problems with DWI may arise when benign lesions have a re-}

mimic cystic lesions, necrotic primary or secondary tumor processes, and image artifacts that may obscure the lesion (4). Problems with DWI may arise when benign lesions have a restriction of diffusion, cystic lesions can imitate necrotic malignant lesions.

In addition to the visual intensity of the signal being assessed (qualitatively) with ADC maps it is necessary to measure the coefficient of diffusion which expresses an average numerical value for each voxel (quantitative -x10$^{-3}$ mm$^2$/s),
setting the ROI on the place of pathological changes. Diffusion coefficient is connected with mobility of water molecules, which reflects the characteristics of the tissue. Cysts and hemangiomas have the highest ADC values due to the relatively unrestricted diffusion of water molecules within their content, while HCC, metastases and FNH show the lowest value, primarily because of its high cellularity (Figure 3). Therefore there is no overlapping between the ADC values of cysts and solid lesions. However, more important clinical problem is distinguishing metastases from hemangiomas, as hemangiomas may exhibit atypical contrast retention, similarly to the hypervascular metastasis or may be hyperintense and therefore exhibit decreased signal intensity on T2 sequences. Necrotic metastases may show marked hyperintensity on T2 signal imitating hemangiomas (Figure 4). ADC values of metastases and hemangiomas are significantly different with regard to their properties, but to some extent overlappings are possible. Also some overlap between FNH and adenomas is possible. ADC/DWI MRI of liver lesions has been researched significantly with focus on comparing DWI with different MRI techniques (standard breath-hold T2-weighted, breathhold, respiratory triggered, MnDPDP MR imaging, T2-weighted turbo spinecho sequences, etc.) (5-12). Regardless of the possible overlap between malignant and benign hepatic lesions and the non-uniformity in the cut-off value for normal liver parenchyma, according to the literature, ADC value of liver metastases is in the range of 0.94-2.85x 10^-3 mm^2/s and the normal liver parenchyma 0.69-2.28x 10^-3 mm^2/s. Each course has its own scan parameters (breath-hold, respiratory triggered, and navigator echo techniques) and different ADC values, and therefore different cut-off (13). Nevertheless, DWI/ADC is of great benefit because the measurement of the coefficient of diffusion—ADC values may represent a valorization factor in monitoring the oncological therapy (14).

In our study, we demonstrated that the ADC map is reliable in distinguishing benign (hemangioma, FNH) from malignant lesions (HCC, CCC, CRC metastases). Based on the obtained average ADC value of benign hepatic lesions of 1.88 (1.326 to 2.48) x 10^-3 mm^2/s and malignant liver lesions of 1.15 (1.024 to 1.343) x 10^-3 mm^2/s statistical testing showed a statistically significant difference (p <0.0005). The obtained cut-off ADC value between benign and malignant lesions is 1.341 x 10^-3 mm^2/s and DWI accuracy in the overall differentiation of liver lesions of 96.8%. Taouli and Koh on the work of review report the results of various studies in which the value of ADC cut-off ranged from 1.47 x 10 to 1.63 x 10^-3 mm^2/s, which can be used for optimal differentiation of benign from malignant lesions (15). Cut-off ADC value that we get from 1.341 x 10^-3 mm^2/s is slightly lower than the average of the above study, but higher than the one of Cieszanowski et al which was 1.25 x 10^-3 mm^2/s (16). Filipe et al. used the cut off value of 1.43x 10^-3 mm^2/s when differentiating benign from malignant lesions and have concluded that the ADC value of malignant lesions is significantly lower compared to benign lesions (17). Testa et al. obtained the results that showed statistically significant difference between benign and malignant lesions with the cut-off value of 1.2x10^-3 mm^2/s, and the accuracy of 71% (18). There are several possible reasons that explain these differences, including the use of different hardware, the lack of standardized protocols for image acquisition (using different b values), different methods for calculating ADC and different population of patients. The growing use and importance of DWI will certainly with future development contribute to uniformity of parameters for image acquisition.

5. CONCLUSION

Measuring ADC values with DWI as an additional MR imaging tool can help in oncological practice by distinguishing normal liver parenchyma from focal lesions, and in differentiating benign from malignant liver lesions, particularly in cases where administration of contrast is not possible.

• Declaration of interests: The authors declare no conflict of interests.

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