Predicting Response to Benzamide Riboside Chemotherapy in Hepatocellular Carcinoma Using Apparent Diffusion Coefficient of Water

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Abstract. Aim: To monitor the effects of the apoptotic agent benzamide riboside (BR) on tumor volume and water apparent diffusion coefficient (ADC) in rat hepatocellular carcinoma (HCC). Materials and Methods: Water ADC of the tumors and nearby liver tissue was measured using diffusion-weighted 1H MRI (DWI). The two groups of BR-treated animals, which differed in their sensitivity to the treatment, were identified as responsive (RBR) and non-responsive (NRBR). Results: Tumor growth in the RBR group was arrested and the mean tumor volume in this group was 1/6th and 1/16th compared to that of the NRBR group on days 7 and 14 after treatment, respectively. Water ADC of HCC was higher than in nearby normal liver tissue. Before BR treatment, the mean water ADC was significantly higher in the RBR group compared to the NRBR group. BR therapy did not change the water ADC value regardless of tumor sensitivity. Conclusion: Although the water ADC did not change after chemotherapy by BR, DWI has great potential for detecting and predicting response to chemotherapy in HCC.

Hepatocellular carcinoma (HCC) and liver metastases from colon and breast carcinomas are an increasing problem worldwide, representing the third largest cause of cancer-related death (1). It is becoming a great health problem not only in Asia and Africa, but also in Western countries mostly due to the major risk factors of HCC growth, such as cirrhosis and hepatitis, which encourage genetic changes that lead to neoplastic transformations (1). As many chemotherapy regimens have been developed for HCC treatment, accurate, non-invasive and quantitative monitoring of the early response to chemotherapy is critical as this data may indicate a need for alteration in therapy. An accurate monitoring scheme would also result in a reduction of invasive needle biopsies.

Chemotherapy efficiency has been estimated in a number of animal studies by using water apparent diffusion coefficient (ADC) measured by diffusion-weighted 1H MRI (DWI) (2-3). However, measurement of water ADC in abdominal tumors is challenging due to its sensitivity to respiratory, cardiovascular and/or gastrointestinal motions (4). To avoid these difficulties, most studies on the effect of chemotherapy on water ADC in rodent tumor models have used tumors placed subcutaneously (5-7). In patients, DWI of tumors and liver is a common technique because of the ability to breath-hold combined with the recent technology of single-shot echo-planar imaging with fat suppression (8-9). However, in the rat HCC model, water ADC in tumor and surrounding liver tissue has been evaluated only in a few studies (10-11). Thus, estimation of water ADC in orthotopic HCC before and after treatment is essential.

Benzamide riboside (1-β-D-ribofuranosylbenzene-3-carboxamide, BR) is novel antitumor drug (12). In fact, BR is a nucleoside produg that is metabolized intracellularly in malignant cells to an analog of NAD, benzamide adenine dinucleotide, that strongly and selectively inhibits inosine-5'-monophosphate dehydrogenase, which is the rate-limiting step of de novo guanylate synthesis. This action interferes with nucleic acid metabolism by halting DNA and RNA synthesis (12). BR induces apoptosis in K562, HL-60, and ovarian...
carcinoma cell lines by depleting GTP. The exact mechanism of this activity is under investigation, but may involve the observed property that benzamide adenine dinucleotide inhibits malate dehydrogenase, a mitochondrial enzyme involved in cellular respiration. Because BR is metabolized in normal hepatocytes by carboxylation to an inactive benzene carboxylic acid riboside, BR has selective toxicity in liver, tumors while preserving normal hepatocytes (12). In our study BR was administered through the hepatic artery. HCC tissue is highly vascular and the main blood transportation is switched from the portal vein to the hepatic artery (13). The main advantage of hepato-arterial intervention is the possibility of increasing drug concentration in certain liver tumor regions and decreasing systemic toxicity.

In this study, the effects of BR delivered through the hepatic artery on HCC tumor volume and water ADC in tumor and surrounding liver tissue were examined to answer the following questions: i) Are pretreatment values of water ADC useful for predicting response to therapy? ii) Is water ADC affected following therapy? iii) Are changes in HCC volume and water ADC correlated?

Materials and Methods

Tumor model. All animal studies were approved by the Indiana University Institutional Animal Care and Use Committee. N1S1 cells (American Type Tissue Culture Collection, Bethesda, MD, USA), a rat Novikoff hepatoma cell line, was maintained as an exponentially growing suspension in Iscove modified Dulbecco’s Medium (Sigma-Aldrich, St. Louis, MO, USA). To create the intrahepatic HCC, 1×10⁶ N1S1 cells were inoculated in the left lateral liver lobe of the male Sprague-Dawley rats (Harlan, Indianapolis, IN, USA) weighing 450-500 g. The tumors were allowed to grow for one week before the first MRI experiment and for two weeks before angiography. Tumor growth was measured from T₂⁺H images with b-value=0 s/mm². After the 28-day MRI experiment, livers containing tumors were excised for histology (Figure 1A). Doubling time of tumor growth was calculated using exponential curve of tumor volume on days 7, 14, 21, and 28 after cell injection.

Hepatic angiography and BR treatment. A 3-cm incision was made in the right groin to expose the femoral artery. A 5-mm incision was made longitudinally in the femoral artery and an Excel 14 microcatheter (Guidant, Indianapolis, IN, USA) were carefully inserted into the femoral artery, the tip was advanced under fluoroscopic guidance into the thoracic aorta and the catheter was able to track in the femoral artery, the tip was advanced under fluoroscopic guidance into the thoracic aorta and the catheter was able to track in the femoral artery, the tip was advanced under fluoroscopic guidance into the thoracic aorta and the catheter was able to track. The catheter and guide wire were then used to select the common hepatic artery, and a rotational angiogram was obtained with a KX0-100 digital subtraction Carm unit (Toshiba, Tokyo, Japan) with rotational digital subtraction at 15°/s rotation at 15 fps, 66 kV, 125 mA, and 43 ms exposures (Figure 1B). Total contrast agent injection during rotational digital subtraction angiography was 3 mL of Hypaquelemigleumene 60% (Amersham, Princeton, NJ, USA). BR (20 mg/kg) was then infused into 10 rats into the hepatic artery extending into the liver. The catheter was removed and the femoral artery was ligated(14).

MRI experiments. All in vivo MRI images were acquired with a Varian 9.4 Tesla, 31-cm horizontal bore system (Varian, Paolo Alto, CA, USA). HCC was examined weekly for 4 weeks after N1S1 cell inoculation. Animals were anesthetized with 1-1.5% isoflurane delivered in medical air at 1-1.5 l/min using a rat nose mask connected to a gas anesthesia machine (Vetland, Louisville, KY, USA). Rat respiration was monitored with a respiration module (SA Instruments, Stony Brook, NY, USA) using a sensitive transducer located under the animal’s abdominal area. The magnet was shimmed to less than 140 Hz line width at half height of the ¹H water signal.

¹H MRI. Water ADC of the tumors and nearby liver tissue were measured with a birdcage volume coil (ID=63 mm, length=190 mm) tuned to 400 MHz. The liver area was placed in the middle of the coil. Multi-slice DWIs of the tumor were collected using a spin-echo sequence consisting of two diffusion gradient pulses of Δ=6 ms duration separated by a Δ=11 ms period applied along all three axes. Interleaved DWI with four b-values (b=0, 256, 945 and 1679 s/mm²), 1100 ms repetition time (TR), 21 ms echo time (TE), 256x128 data points over a 64x64 field of view (FOV), 16 slices, 0.5 mm slice thickness, 1.5 mm slice gap were collected. Respiratory gating was used to minimize the motion effect on water ADC. In addition, the animal respiration rate was brought to a relatively stable level (≤40 breaths/min) to minimize variation in TR by slightly adjusting the isoflurane anesthesia. Water ADC was monitored only during the end-expiration time. Total data collection time for a set of DWI was 15 min. ¹H images and water ADC maps were reconstructed using the Image Browser software provided by Varian. The tumor volume and average water ADC were determined over a three-dimensional (3D) volume of interest for each temporal measurement.

¹H diffusion data were fit to the Stejskal-Tanner equation (15). The average ADCs in intrahepatic tumor and surrounding liver were calculated using a 3×10⁻³ mm²/s threshold to eliminate abnormal ADC values that were clearly higher than the diffusion coefficient of free water (16). In general, the threshold process decreased the ADC values by 25-30% eliminating a signal artifact related to the motion effects. Water ADC in HCC was compared to water ADC in surrounding liver tissue, which also exhibited motion effects.

Histology. The intrahepatic tumors with the surrounding liver tissue were excised from the animal body, fixed in 10% formalin solution (Fisher Scientific, Pittsburgh, PA, USA), and then embedded in paraffin. The histological sections of the tumors were cut along the same plane as the MR images. Tissue sections were obtained at 5 μm thickness and stained with hematoxylin and eosin (H&E) to identify the viable, necrotic, and inflammation tumor regions (Figure 1C).

Data analysis and statistics. All statistical data are presented as the mean±standard error of mean (SEM) and represent the range across a cohort of animals (n=12 (control group), 5 (responsive (RB)), and 5 non-responsive to BR (NRBR) groups). Analysis of the data was performed by ANOVA with post-hoc comparisons among the experimental groups and time points using the least significant difference test (Statistica/v. 5.1 program). A p-value ≤0.05 was used to define statistical significance.
Results

Tumor volume. The two groups of BR-treated animals, which differed in their sensitivity to the treatment as was estimated from tumor volume changes, were identified as RBR and NRBR. Representative T2 ¹H MRI from control, NRBR, and RBR groups are presented in Figure 2. In the RBR group, 7 days post BR treatment (20 mg/kg), the tumor ceased to grow, and almost disappeared on 14 days post treatment. In the NRBR group, the tumor continued to grow after BR treatment with a rate similar to that of the control group. The mean tumor volumes are shown in Figure 3A as calculated using T2 weighted ¹H MRI. The mean tumor volumes for all three groups were similar before treatment on day 7 (0.13±0.05 cm³ for control group (A), 0.03±0.02 cm³ for NRBR group (B), and 0.04±0.01 cm³ for RBR group (C)). On day 14 the tumors grew to 0.85±0.19 cm³ (control, $p \leq 0.05$ vs. day 7), 0.40±0.16 cm³ (NRBR, $p \leq 0.05$ vs. day 7), and 0.20±0.11 cm³ (RBR, $p \leq 0.15$ vs. day 7). In the NRBR group, the tumors continued to grow during the next 1-and 2-week post-treatment to 1.51±0.71 cm³ ($p \leq 0.05$ vs. day 7) and 3.00±1.60 cm³ ($p \leq 0.07$ vs. day 7), respectively. Tumor growth in the RBR group was arrested: 0.24±0.12 cm³ and 0.19±0.09 cm³ in tumor volume on days 7 and 14 post-treatment, respectively. Furthermore, the mean tumor volume in the RBR group was 1/6th and 1/16th compared to the NRBR group ($p \leq 0.05$) on days 7 and 14 after treatment, respectively. The doubling time of the tumor growth in control and NRBR groups was 3.2 and 3.9 days, respectively. In the control group, the tumors continued to grow during the third and fourth weeks of experiments to 2.58±0.52 cm³ ($p \leq 0.05$ vs. day 7) and 5.33±1.2 cm³ ($p \leq 0.05$ vs. day 7), respectively.
Water ADC. Figure 4 shows selected DWI transaxial sections and the ADC map of the rat liver region. HCCs (28 days after cell inoculation) are marked by solid arrows. The image of the liver containing HCC was slightly blurred with $b$-values of 945 or 1678 s/mm² even though they were collected with respiratory gating, however, the ADC map shows only a moderate motion effect in the liver area. The mean water ADC values in HCC before BR treatment (on day 7 post cell inoculation) are shown in Figure 3B and time track changes in the mean of water ADC values in HCC and nearby normal liver are shown in Figure 5. On day 7 after cell inoculation, the mean water ADC was (in $10^{-3}\text{mm}^2/\text{s}$) 1.23±0.08 (control group), 1.13±0.06 (NR BR group), and 1.74±0.14 (R BR group) (Figure 3B). Seven days later, the mean water ADC was not significantly changed (in $10^{-3}\text{mm}^2/\text{s}$): 1.26±0.06 (control group), 1.26±0.07 (NR BR group), and 1.51±0.11 (R BR group) (Figure 5). Thus, before BR treatment, the mean water ADC was significantly ($p<0.05$) higher by 1.54 (day 7, $p<0.05$) and 1.2 times (day 14, $p<0.05$) in the R BR group compared to NR BR group.

On days 14, 21, and 28 after cell inoculation, the water ADC in control HCC remained relatively stable in the range of 1.11-1.26×10^{-3} mm²/s. BR treatment did not change the water ADC values in both NR BR and R BR groups. However,
the differences between these two groups were not significant after BR therapy ($p<0.15$ and $p<0.31$ on days 7 and 14 post-treatment, respectively) mostly probably due to higher variability at these data points.

In all three groups, the mean water ADC in the normal liver tissue was almost identical and unchanged during the one-month experiment. The ranges of the mean water ADC in the liver were between (in $10^{-3}$ mm$^2$/s) 0.74-0.85 (control group), 0.76-0.85 (NR BR group), and 0.78-0.81 (R BR group) (Figure 5). Before BR treatment, the mean water ADC in HCC of all three groups was significantly ($p<0.05$) higher compared to the adjacent normal liver. For example, 7 days after cell inoculation, the ratio of water ADC in the tumor to that in the liver was 1.49±0.13 (control group), 1.48±0.06 (NR BR group), and 2.2±0.08 (R BR group). After BR treatment, the mean water ADC in HCC continued to be significantly ($p<0.05$) higher compared to the adjacent normal liver in control and R BR groups but not in the NR BR group ($p<0.13$). HCC-to-liver ADC ratio did not change significantly with tumor growth in control and NR BR groups or with arrest of tumor growth in the R BR group.

Discussion

In this work, the changes in tumor volume and water ADC in the rat HCC before and after interventional treatment of the tumor with BR were studied. The measurement of water ADC in the rat intrahepatic HCC was challenging due to respiratory, cardiac, bowel, etc. motions. To minimize these problems, we decreased the respiratory rate to 35–40 breaths/min and used gating to monitor water ADC only during the end-expiration time when diaphragmatic motion is reduced. In addition, the calculation of water ADC was done by: i) using thresh-holding to eliminate ADC values that were clearly higher than free water diffusion (>3 mm$^2$/s), ii) avoiding the undoubtedly bright spots on ADC maps when the region of interest was drawn, and iii) comparing water ADC in HCC and surrounding normal liver tissue, which was also undergoing similar motion effects. These approaches helped to achieve relatively steady ADC values for each experimental data point.

In HCC, the mean water ADC was significantly higher than in surrounding normal liver tissue by ~1.5 times (control and NR BR groups) and by ~2 times (R BR group) on all experimental days. BR did not change this ratio significantly. These data correlate with other publications showing higher water ADC in HCC compared to healthy liver tissue (3, 17-18). The most reasonable explanation of this effect is less differentiation of tumor cells and a net increase in the tumor relative extracellular space (ECS). Histology of HCC and liver after the last MRI experiment showed that HCC contains areas of inflammation and necrosis with increased ECS (Figure 1C) in which water ADC could be high. It has also been shown that in experimental mammary tumors, the high and low ADC values also correlate with high and low necrosis, respectively (5).

Intrahepatic infusion of BR was a semi-effective treatment of intrahepatic HCC in rats. Thus, the two groups of BR-treated animals, which differed in their sensitivity to the treatment, were identified as R BR and NR BR. Tumor growth in the R BR group was arrested while in control and NR BR groups the tumor continued to grow. We have previously shown that BR induces apoptosis in rabbit and rat liver tumors with minimal apoptosis in normal liver (19-20). For example, the mean tumor apoptosis rates were 71% with 10 mg/kg BR and only 1% with saline solution treatment. BR-mediated...
induction of tumor apoptosis makes BR an excellent candidate for chemotherapeutic application (12, 21).

Apoptosis and necrosis have to increase water ADC in tumor due to increase in ECS (22). However, this was not the issue in our case. After BR infusion, the water ADC remained unchanged in both $R_{BR}$ and $N_{BR}$ groups. In our experiments, it seems that BR induced apoptotic and necrotic changes leading to increase in ECS and water ADC are counterbalanced by the ADC reducing factors. Histological analysis has shown that HCC contained coagulative type of necrosis characterized by well-packed nucleus-less cells and macromolecular contamination that can restrict water motion and thereby increase the tortuosity of this space. Monte Carlo simulation of diffusion in the ECS have shown that the extracellular volume fraction $\alpha$ and tortuosity factor $\lambda$ are interrelated by Archie’s law $\lambda = \alpha^{-1}$, where $\lambda$ is the tortuosity exponent (23). The increase of tortuosity factor has been correlated experimentally with a dynamic decrease of the ADC in ischemic brain (24). Another factor that can counterbalance a possible increase in ADC could be changes in water diffusion in intracellular space (ICS). Usually the value of ADC in ICS is underestimated beside the fact that the cells occupy 82-85% of the tissue space in the liver (25). The experimental measurements of ADC in ECS and ICS have shown that ADC values in those compartments are more identical than different (16, 26). This is in contrast to the general belief that water ADC is lower in the ICS compared to ECS.

The perfusion factor that can contribute to water ADC value could be the issue in our case (27). However, the effect of perfusion is diminished when wide ranges of $b$-values are used as it was done in this study ($b=0$, 256, 945 and 1679 s/mm²). Nevertheless, this assumption needs further experimental estimation using additional $b$-values in the range 0-100 s/mm². Thoeny et al. (28) propose using the difference between ADCs obtained with low (0-100 s/mm²) and high $b$-values (500-1000 s/mm²) as a perfusion component of the tissue ADC.

It is unclear how the initial ADC values can predict tumor chemosensitivity that could be clinically relevant. Lemaire et al. (5) showed that 5-fluorouracil-treated experimental mammary tumors were distinguishable in terms of water ADC up to day 5 whereas the tumor volume, over the same period of time, did not distinguish the sensitive and non-sensitive groups. Contrary to these data, Seierstad et al. (29) did not find any correlations between pretreatment ADC values and changes in colon adenocarcinoma HT29 xenograft volumes after chemoradiation, whereas early changes in mean ADC quantitatively correlated with treatment outcome. We have found that the initial levels of water ADC in the $R_{BR}$ group were higher compared to the $N_{BR}$ group. These data suggest that a higher initial ADC level (1.5-1.7×10^{-3} mm²/s) could be a promising sign for effective BR treatment, and in contrast, tumors with a lower initial ADC value (1.1-1.3×10^{-3} mm²/s) are most likely to be resistant to BR treatment. The tumors with high initial values of water ADC were more vulnerable to BR therapy, suggesting that the cells were in weaker physiological condition. The weaker physiological condition of the cells may be related to the decreased cell membrane permeability. It has been previously shown that the more sensitive to chemotherapy with carmustine subcutaneous 9L glioma in rat had higher water ADC and $^{23}$Na signal intensity (30). Like water ADC, single quantum $^{23}$Na signal intensity mostly reflects changes in ECS. However, it was shown that ECS did not change in carmustine-treated subcutaneous 9L glioma (31). The authors found that the increase in $^{23}$Na MRI resulted from an increase in intracellular-treated subcutaneous 9L glioma (31). Thus, $^{23}$Na MRI in conjunction with DWI could provide synergistic information of post-treatment changes in liver tumors, allowing prediction and early assessment of treatment success.

In conclusion, the data presented here show that implantation of N1S1 cells in the rat liver can be used as a HCC model for pre-clinical study of transarterial therapy with BR. Water ADC of N1S1-inoculated HCC is higher than in nearby normal liver tissue. Before BR treatment the mean water ADC is higher in the $R_{BR}$ group compared to the $N_{BR}$ group. Intrahepatic infusion of BR is a semi-effective transarterial treatment of HCC in rats. BR therapy did not change the water ADC value regardless to tumor sensitivity. Accurate non-invasive and quantitative monitoring of the water ADC before and following chemotherapy of HCC is critical as this data may help to select an appropriate therapy or indicate a need for alteration in therapy.

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