Monitoring the Effect of Doxorubicin Liposomal in VX2 Tumor Using Magnetic Resonance Imaging

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Abstract—Cancer is still one of the serious diseases threatening the lives of human beings. How to have an early diagnosis and effective treatment for tumors is a very important issue. The animal carcinoma model can provide a simulation tool for the studies of pathogenesis, biological characteristics, and therapeutic effects. Recently, drug delivery systems have been rapidly developed to effectively improve the therapeutic effects. Liposome plays an increasingly important role in clinical diagnosis and therapy for delivering a pharmaceutic or contrast agent to the targeted sites. Liposome can be absorbed and excreted by the human body, and is well known that no harm to the human body. This study aimed to compare the therapeutic effects between encapsulated (doxorubicin liposomal, Lipodox) and un-encapsulated (doxorubicin, Dox) anti-tumor drugs using magnetic resonance imaging (MRI). Twenty-four New Zealand rabbits implanted with VX2 carcinoma at left thighs were classified into three groups: control group (untreated), Dox-treated group, and LipoDox-treated group, 8 rabbits for each group. In each group, MRI scans were performed three days after tumor implantation. A 1.5T GE Sigma HDx1 whole body MRI scanner with a high resolution knee coil was used in this study. After a 3-plane localizer scan was performed, three-dimensional (3D) fast spin echo (FSE) T2-weighted Images (T2WI) was used for tumor volumetric quantification. Afterwards, two-dimensional (2D) spoiled gradient recalled echo (SPGR) dynamic contrast-enhanced (DCE) MRI was used for tumor perfusion evaluation. DCE-MRI was designed to acquire four baseline images, followed by contrast agent Gd-DOTA injection through the ear vein of rabbit. A series of 32 images were acquired to observe the signals change over time in the tumor and muscle. The MRI scanning was scheduled on a weekly basis for a period of four weeks to observe the tumor progression longitudinally. The Dox and LipoDox treatments were prescribed 3 times in the first week immediately after the first MRI scan; i.e. 3 days after VX2 tumor implantation. ImageJ was used to quantitate tumor volume and time course signal enhancement on DCE images. The changes of tumor size were monitored. The MRI scanning was performed immediately after the tumor implantation and every 3 days thereafter. ImageJ was used to measure the tumor volume and to analyze the time course signal change. The Dox and LipoDox treatments were prescribed 3 times in the first week immediately after the first MRI scan. The signal intensity of LipoDox-treated group was significantly lower than that of the other two groups, which implies that targeted therapeutic drug remained in the tumor tissue. This study provides a radiation-free and non-invasive MRI method for therapeutic monitoring of targeted liposome on an animal tumor model.

Keywords—Doxorubicin, dynamic contrast-enhanced MRI, lipodox, magnetic resonance imaging, VX2 tumor model.

I. INTRODUCTION

Drug targeting to specific tumors with minimum toxicity and superior therapy efficacy is one of the important goals for cancer treatment pharmaceutics [1]. Liposome is one of the hottest carriers in drug delivery system for years. It can be absorbed and discharged by the human body, and is well known that no harm to the human body clinically. Clinical-Grade Vasculature-Targeted Liposomal Doxorubicin has been shown that it can be enhanced antitumor efficacy [2]. It has been shown that Liposomal doxorubicin has a lower cardiologic toxicity than doxorubicin [3]. Lipodox has been demonstrated that it can improve radiotherapy response in hypoxic prostate cancer xenografts [4]. Recently, thermosensitive liposomal drug delivery system becomes a hot topic in the area of drug delivery systems [5], [6]. Importantly, the effective inhibition of in vivo tumor growth was proved using Lipodox. Hence, we proposed to investigate the treatment efficacy of free Dox and Lipodox for muscle VX2 tumor in an animal model. The aim of this proposal was to compare the differences in treatment efficacy between encapsulated and un-encapsulated anti-tumor drugs. The discrepancy among untreated, encapsulated and un-encapsulated drug treatment was investigated.

II. MATERIALS AND METHODS

This study intended to set up a New Zealand White (NZW) Rabbit VX2 tumor model to longitudinally observe the development of the tumor after drug therapy using MRI techniques.

A. Animal Model

VX2 carcinoma tumor was implanted at left thighs of NZW rabbits, while right thighs were referred as the normal tissue. Three days after tumor implantation, magnetic resonance imaging scan was performed.

B. Imaging Protocol

1.5T GE Sigma HDxt whole body magnetic resonance scanner and 8-channel high resolution knee coils was used in this study. NZW rabbit was anesthetized with an intramuscularly injected ketamine (10 mg/kg) and xylazine (5 mg/kg). The MRI scanning was performed immediately after the tumor implantation and every 3 days thereafter. The rabbits were placed in a supine position in the MRI scanner to minimize movement artifacts. The images were acquired using a T1-weighted spoiled gradient recalled echo (SPGR) sequence with a high resolution knee coil. A series of 32 images were acquired to observe the signals change over time in the tumor and muscle. The MRI scanning was scheduled on a weekly basis for a period of four weeks to observe the tumor progression longitudinally. The Dox and LipoDox treatments were prescribed 3 times in the first week immediately after the first MRI scan; i.e. 3 days after VX2 tumor implantation. ImageJ was used to quantitate tumor volume and time course signal enhancement on DCE images. The changes of tumor size were monitored. The MRI scanning was performed immediately after the tumor implantation and every 3 days thereafter. ImageJ was used to measure the tumor volume and to analyze the time course signal change. The Dox and LipoDox treatments were prescribed 3 times in the first week immediately after the first MRI scan. The signal intensity of LipoDox-treated group was significantly lower than that of the other two groups, which implies that targeted therapeutic drug remained in the tumor tissue. This study provides a radiation-free and non-invasive MRI method for therapeutic monitoring of targeted liposome on an animal tumor model.
injection using Zoltil 50 and Rompun 2% mix solution before
MRI scanning. The VX2 tumor site was landmarked at the
center of the knee coil. The scanning protocol was listed as
follows: (1) 2D fast spoiled gradient recalled echo (FSPGR)
pulse sequence was scanned to obtain multi-slice localizer
images for subsequent scanning, (2) T2-weighted (T2W) spin
echo (SE) pulse sequence was performed to obtain axial images,
(3) 3D fast spin echo (FSE) pulse sequence was performed to
achieve 3-D axial images on tumor sites, and finally (4) 2D
spoiled gradient recalled echo (SPGR) was used to assess the
signal changes over time before and after Gd-DOTA contrast
agent injection. Dynamic contrast enhanced (DCE) MRI
scanning was arranged to obtain 4 reference images first. After
Gd-DOTA was bolus injected into the ear vein of a NZW
rabbit, 32 consecutive images were acquired to evaluate the
signal intensity changes and enhancement pattern over time.

Twenty four NZW rabbits were classified into three groups
with 8 rabbits in each group: the control group (without
treatment), Dox treatment group (free doxorubicin) and
Lipodox (doxorubicin liposomal) treatment group. After
magnetic resonance imaging scanning protocol was performed
to obtain VX2 tumor images, Dox or Lipodox was injected via
the ear vein of NZW Rabbit for chemotherapy. Totally, three
times chemotherapy and four times MRI scans were performed
on a weekly basis for longitudinal investigation.

3D FSE images were used for quantitating tumor volume
changes over time using ImageJ software. The region of
interest (ROI) was segmented and the ROI values were
measured on VX2 tumor in DCE MRI images. Then, the
dynamic signal intensity changes of tumor over time were
obtained. Furthermore, the treatment efficacy between
Lipodox, and Dox treatment was compared.

Fig. 1 The time course tumor DCE MR images (Pre-treatment) show
contrast enhanced signal intensity changes with time after Gd-DOTA
injection.

The scanning parameters for FSE were TR = 2000 ms, TE =
72.36 ms, FA = 90°, FOV = 16 × 16 cm², BW = 25 kHz, matrix
= 256 × 128, NEX = 1, number of slice = 8 and slice thickness =
5 mm. And the scanning parameters for DCE-MRI were TR =
80 ms, TE = 1.32 ms, FA = 60°, FOV = 16 × 16 cm², BW =
31.25 kHz, matrix = 256 × 256, NEX = 1, number of slice = 8
and slice thickness = 5 mm.

Fig. 2 The time course tumor DCE MR images of control group

Fig. 3 The time course tumor DCE MR images of Dox treatment group

Fig. 4 The time course tumor DCE MR images of Lipodox treatment group

III. RESULTS

The time series tumor DCE-MR images are shown in Fig. 1.
It was found that the signal intensity increased with time, i.e.
contrast enhancement. The tumor rim was clearly shown. Fig. 2
shows the time course tumor DCE-MR images of control group
with larger and larger tumor size with time. Figs. 3 and 4 show
the time course tumor DCE-MR images of Dox treatment and
Lipodox treatment groups, respectively. Figs. 5-8 show the
time varying intensity curve of tumor on DCE images at
subsequent weeks, respectively. It is found that MR signal
The intensity of tumor for Lipodox treatment group was the lowest for all weeks. However, the MR signal intensity of Dox treatment group was lower than untreated group at the first week, and higher than untreated group at the 2nd and the 3rd week, and became lower again at the 4th week. The time varying tumor volume measured from FSE images is listed in Table I.

**TABLE I**

<table>
<thead>
<tr>
<th>Week</th>
<th>1st (cm³)</th>
<th>2nd (cm³)</th>
<th>3rd (cm³)</th>
<th>4th (cm³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated</td>
<td>10.2</td>
<td>28.2</td>
<td>58.3</td>
<td>80.8</td>
</tr>
<tr>
<td>DOX</td>
<td>31.8</td>
<td>32.9</td>
<td>55.9</td>
<td>36.2</td>
</tr>
<tr>
<td>LIPODOX</td>
<td>15.4</td>
<td>55.4</td>
<td>48.4</td>
<td>34.2</td>
</tr>
</tbody>
</table>

**Fig. 4** The time course tumor DCE MR images of Lipo-Dox treatment group

**Fig. 5** The time varying intensity of tumor on DCE images at first week (Pre-treatment)

**Fig. 6** The time varying intensity of tumor on DCE images at second week

**Fig. 7** The time varying intensity of tumor on DCE images at third week

**Fig. 8** The time varying intensity of tumor on DCE images at fourth week
IV. DISCUSSION AND CONCLUSIONS

The results found that both encapsulated treatment group and unencapsulated treatment group can effectively inhibit the growth of tumors. DCE MRI found the signal intensity in the encapsulated treatment group was significantly lower than the other two groups which implied that the encapsulated anti-cancer drugs targetedly remained in tumor tissue. T2W MRI found the tumor volume of the encapsulated treatment group was significantly lower than the other two groups. It was an evidence to show encapsulated treatment group is superior to unencapsulated treatment group. In conclusion, the treatment efficacy of Lipodox is superior to Dox treatment due to its targeted characteristics.

REFERENCES


