Developing a tool for the validation of quantitative DCE-MRI

Karin Bolabc, Joost C. Haeckabc, Lejla Alicab, Monique Bernsenb, Marion de Jongc, Wiro J. Niessenabd, Jifke F. Veenlandab

aBIGR – Biomedical Imaging Group Rotterdam, Departments of Radiology and Nuclear Medicine, Erasmus MC, Rotterdam, P.O. Box 2040, 3000 CA Rotterdam, The Netherlands; dFaculty of Applied Sciences, Delft University of Technology, Delft, The Netherlands

ABSTRACT

Dynamic Contrast-Enhanced Magnetic Resonance Imaging (DCE-MRI) is becoming an indispensable tool to non-invasively study tumor characteristics. However, many different DCE-analysis methods are currently being used. To compare and validate different methods, histology is the gold standard. For this purpose, exact co-localization between histology and MRI images is a prerequisite. In this study a methodology is developed to validate DCE-data with histology with an emphasis on correct registration of DCE-MR I and histological data. A pancreatic tumor was grown in a rat model. The tumor was dissected after MR imaging, embedded in paraffin, and cut into thin slices. These slices were stained with haematoxylin and eosin, digitized and stacked in a 3D volume. Next, the 3D histology was registered to ex-vivo SWI-weighted MR images, which in turn were registered to in-vivo SWI and DCE images to achieve correct co-localization. Semi-quantitative and quantitative parameters were calculated. Preliminary results suggest that both pharmacokinetic and heuristic DCE-parameters can discriminate between vital and non-vital tumor regions. The developed method offers the basis for an accurate spatial correlation between DCE-MRI derived parametric maps and histology, and facilitates the evaluation of different DCE-MRI analysis methods.

Keywords: DCE-MRI, histology, registration, DCE-analysis, quantitative, semi-quantitative, pharmacokinetic modeling, two-compartment model

1. INTRODUCTION

In oncology, Dynamic Contrast-Enhanced Magnetic Resonance Imaging (DCE-MRI), in which the signal intensity over time is measured following contrast injection, is frequently used as a tool to detect tumors or metastases, to monitor tumor growth and to assess the response to anti-tumor treatment. Although many different analysis methods have been used to quantify tumor characteristics from DCE-MRI data, a standard analysis method has not yet been adopted.

Both semi-quantitative and quantitative analysis methods can be employed to analyze the signal intensity time curves (SI-curves) in DCE-MRI. A semi-quantitative model derives parameters from the SI-curves such as the area under the curve (AUC), slope of wash-in and wash-out of the contrast agent and maximal enhancement. The semi-quantitative model is straightforward and easy to implement. However, derived parameters do not necessarily have a physical correlation with the underlying tumor characteristics and often are a mixture of several quantitative parameters/measures. Furthermore, it is difficult to compare measurements from scans acquired at different times or from different scanners, owing to the lack of an absolute intensity scale. Quantitative analysis methods are based on pharmacokinetic modeling, assuming that the contrast is divided into several compartments. Analysis using quantitative models is less straightforward as additional information is required, such as a T1-map and knowledge of the Arterial Input Function (AIF).

For the validation of DCE-MRI derived parameters, histology is still the gold-standard. The haematoxylin and eosin stain (H&E) is commonly used to discriminate viable from necrotic tumor tissue. However, not all studies show a correlation between DCE-parameters and histology. One of the factors influencing this correlation could be the poor co-localization of the histology with the MRI slices, which has not received much attention. Also changes in morphology (shrinkage) and orientation once specimens are dissected and placed in formalin fixative may play a role. In view of this, the goal of this study is to develop a novel methodology for validating DCE-MRI derived parameters, by achieving a more accurate co-localization between histology and DCE-MRI images. Hereto, a multi-step registration process involving an intermediate ex-vivo MRI scan will be used.
2. METHODS

2.1 Animal and tumor model

For this study a male Lewis rat (Harlan_CPB, Austerlitz, The Netherlands), weighing approximately 300 grams, was used with approval of the Committee for Animal Research of the Erasmus MC, Rotterdam, The Netherlands. Pancreatic tumor cells (CA20948 tumor) were inoculated subcutaneously in the right hind limb. The tumor was imaged when it reached a diameter of 1.5-2 cm. The animal was anesthetised before imaging using Isoflurane (Nicholas Pyramal Limited, London, UK). After completing the examinations the animal was sacrificed.

2.2 DCE-MRI Imaging protocol

A clinical 3 Tesla MRI scanner (Signa Excite, GE Healthcare, USA) was used for imaging. All images were acquired using the body coil. DCE images were obtained using TRICKS (Time Resolved Imaging of Contrast Kinetics), for achieving high temporal resolution. T1-weighted MR images were used for calibration of the DCE data and SWI-weighted MR image were used for registration purposes (see also figure 1 in section 2.4.1). In-vivo MRI settings: T1-weighted MRI: Tr/Te=20/4 ms, flip angles of 3, 5, 10, 15, 20, 35 and 50º and a XYZ resolution of 0.2x0.2x0.7; SWI-weighted MRI: Tr/Te=24.4/12.0 ms, flip angle of 10º and a XYZ resolution of 0.098x0.098x0.20; TRICKS DCE-MRI settings: Tr/Te=5.44/1.46 ms, flip angle of 10º, XYZ resolution of 0.2x0.2x0.7 and a temporal resolution of 4.13 ms. Ex-vivo MRI settings: SWI-weighted MRI: Tr/Te=23.5/8.9 ms, flip angle of 10º and a XYZ resolution of 0.098x0.098x0.40.

2.3 Histology

After in-vivo MR imaging the tumor was dissected and dyed in order to track its orientation with respect to the in-vivo situation. After dissection the tumor was placed in 200 cc 10% buffered formalin (Boom, The Netherlands) for fixation purposes. When the fixation process was completed, a thin section of the tumor was sliced off along the longest tumor axis in order to create a reference plane for ex-vivo MRI, thereby defining the plane for histological sectioning. The tumor was washed and embedded into agar to prevent tumor movement and to provide a water rich environment for ex-vivo MRI. After the ex-vivo MR imaging, the tumor was embedded in paraffin. Sections of 4 µm at approximately 80 µm intervals were cut using a microtome. These sections were fixed on a glass slide and stained with haematoxylin and eosin (H&E). The slides were digitized using the NanoZoomer (C9600. Hamamatsu, Japan) with a magnification of 20 and a resolution of 3.64 µm.

2.4 Data analysis

2.4.1 Registration DCE-MRI with histology

In order to match histology to DCE-MRI, the registration scheme of Alic et al.9 was used. In short this method consists of two steps: A) registration of 3D stacked histology to 3D ex-vivo SWI-weighted MRI, and B) registration of 3D ex-vivo SWI-weighted MRI to 3D in-vivo MRI. (figure 1). The 3D histology volume was created by stacking the digitized histology slices using a rigid registration scheme. Then the registration of histology to ex-vivo and ex-vivo to in-vivo was achieved using third order non-rigid B-splines with mutual information as similarity measure to be optimized. The separate transformations were subsequently concatenated in order to accomplish co-registration of 3D histology to in-vivo MRI. All registrations were performed using Elastix10 (http://elastix.isi.uu.nl/).
Figure 1: Different steps to achieve registration between 3D histology and DCE-MRI images: 1) in-vivo SWI to DCE-MRI registration, 2) ex-vivo SWI, 3) histology data, 4) stacking histology slices into a 3D volume, 5) registering histology to ex-vivo SWI, 6) registering ex-vivo SWI/histology to in-vivo SWI/DCE, 7) calculating DCE-parameter maps.

2.4.2 DCE-MRI quantification
The semi-quantitative parameters (i) maximum enhancement (Smax), (ii) area under the curve (AUC), (iii) wash-in and (iv) wash-out were calculated from the SI-curve. Next this curve was converted into a contrast-enhancement curve using equation (1):

\[ C(t) = \frac{1}{\alpha} \left( \frac{1}{T_1(t)} - \frac{1}{T_1(0)} \right) \]  

with \( C(t) \) (mM) representing the concentration of the contrast agent at time \( t \), \( \alpha \) the relaxivity (mM/s) of the contrast agent (Gadovist; \( \alpha = 3.6 \)), \( T_1(t) \) the T1-value of the tissue at time \( t \) after injection of the contrast agent and \( T_1(0) \) the
T1-value of the tissue before contrast injection. The latter was calculated using a linearized T1-mapping method\textsuperscript{11}. Quantitative parameters were calculated by fitting a two-compartment model (originally proposed by Brix et al.\textsuperscript{12}, figure 2) to the contrast-enhancement curve.

![Diagram of two-compartment model](image)

This two-compartment model was solved by using a Laplace transform\textsuperscript{13}, where the bolus of the arterial input function (AIF) is assumed to be an instantaneous injection of contrast agent and thus modeled as a delta function. The derived equation becomes:

$$ C(t) = \frac{A}{a + b} \left( \exp^{-bt} - \exp^{-at} \right) $$

with $k_{12}$ a measure of influx of contrast agent into the tumor (also known as $k_{trans}$) and $k_{21}$ the out flux of contrast agent ($k_{op}$). Parameters $a$ and $b$ represent a combination of $k_{12}$, $k_{21}$ and $K_{out}$. $V_c$ and $V_p$ are the volumes of the central-blood plasma compartment and peripheral EES respectively, and $t$ is the time passed since injection.

2.4.3 Validating DCE-MRI with histology

In order to validate the different DCE parameters, vital and non-vital regions were selected by drawing regions of interest (ROI’s) on histology slices in the 3D volume. These regions were then transformed to the parametric maps, using the transformation parameters resulting from the registrations.

3. RESULTS

An example of derived DCE-MRI parametric maps is shown in figure 3. As can be seen from the r-square values in the figure (upper right), the two-compartment model used here does not always result in a good fit. Vital (black) and non-vital (gray) regions of interest were drawn in the registered histology slice. For these regions mean and standard deviation were calculated in the subsequent DCE parameter maps (table 1). Some of the standard deviations are considerable, however, a two-tailed t-test performed between the vital and non-vital regions per parameter showed that all difference were significant (p-value $<< 0.05$). Scatter plots were made to investigate whether the DCE parameters allow for segmentation between vital and non-vital regions (figure 4). It seems that segmentation is possible using a combination of two parameters.
Figure 3: An example of DCE-MRI parametric maps, with at the top left the registered histology slice. The black circle denotes a vital region of interest and the gray circle denotes a non-vital region.
Figure 4; Scatter plots for the different DCE parameters of pixel-values in the vital and non-vital regions of interest.
Table 1; Mean and standard deviation (sd) of vital and non-vital regions of interest for different DCE parameters

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>NON-VITAL ROI</th>
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4. CONCLUSION

A framework has been developed to accurately register DCE-MRI data to histology in order to validate MRI derived parametric maps. Non-rigid registration is necessary for exact co-localization of DCE-MRI data with histology. However, artifacts due to cutting (tearing and shattering) and alterations in morphology (shrinkage) hamper correct co-localization. Therefore, the DCE-MRI validation has to be limited to regions where registration was successful.

Preliminary results for the tumor we studied show significant different mean values between vital and non-vital regions for parameter maps calculated with the two-compartment model as well as with semi-quantitative measures. In addition, parameters b, wash-in and AUC90 can discriminate between the different tumor regions. For this tumor, the combination of two well chosen DCE-MRI parameters will enable segmentation into vital and non-vital regions. In the future this study will be expanded using more data and including multiple quantitative DCE-analysis methods, different AIF estimation procedures and calibration methods, and different histology stains. Also a method for segmenting vital and non-vital tumor tissue will be developed.

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REFERENCES


