An original methodology for quantitative assessment of perfusion in small animal studies using contrast-enhanced ultrasound.

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To cite this version:


HAL Id: inserm-00191133
http://www.hal.inserm.fr/inserm-00191133
Submitted on 24 Nov 2007

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Abstract—The aim of this paper was to validate the combination of two original methods for assessing perfusion in small animal studies using Contrast-Enhanced Ultrasound. Respiratory motion was first removed using a frame selection method. This method, based on a Principal Component Analysis, enables the definition of two subsequences, corresponding to the end-of-inspiration plane and to the end-of-expiration plane. Attenuation caused by contrast agent microbubbles was then estimated. The developed method, based on the assumption that a linear relationship holds between local attenuation and backscatter, used the time-intensity variation in a highly echogenic distal area without contrast uptake to estimate local attenuation coefficients. Ten murine renal perfusion studies were investigated after a bolus injection of SonoVue. Replenishment kinetics were acquired too and were finally modeled with an exponential function. Preliminary results were promising. Indeed, the SonoVue concentration was more faithfully represented than on native images, which allowed a better assessment of perfusion parameters in the whole field of view.

I. INTRODUCTION

The development of ultrasound contrast agents (UCA) and contrast-specific imaging modalities has increased the interest in ultrasound for functional studies by improving the contrast resolution and suppressing signal from stationary tissues. In addition, UCA consist of microbubbles and thus are blood-pool agents, making Contrast-Enhanced Ultrasound (CEUS) an excellent tool for delineating perfused tissues. They allow functional assessment of their contrast enhancement, especially for tumor angiogenesis and response to new therapeutic regimes.

Several methods have been developed for quantifying microvascular perfusion using CEUS in vitro and in vivo [1], [2], [3]. These methods are based on the fact that microbubbles can be destroyed by ultrasound. During continuous infusion of a UCA, the kinetics of bubble destruction [2] or bubble replenishment after destruction [1], [3] can be used to describe and quantify organ perfusion. Quantification of contrast uptake kinetics starts with the design of user-defined regions-of-interest (ROI). The time-intensity variation for each ROI is then fitted with a model, e.g., an exponential function [1].

However perfusion coefficients estimated with these models could be impaired by several artefacts. The aim of this paper is to propose an original methodology for quantifying perfusion in small animal CEUS studies by investigating original methods to correct for these artefacts.

Artefacts may be due to motion that is present in medical images. As the same tissue should remain in the studied ROI for an accurate quantification, any motion has to be removed before. Motion artefacts may be related either to patient’s respiration or to probe displacement. Motion involves a degradation of estimates of perfusion indices and needs to be removed. Many methods for automatic motion compensation have been developed, using rigid or non-rigid registration algorithms. However these methods are based on the assumption that the motion is always contained in the acquisition plane, which is not necessarily true for in vivo acquisitions. Indeed, the motion is most of the time three-dimensional, whereas the acquisition is only bidimensional. New methods have thus to be considered and investigated.

In this paper, an original frame selection method that selects the frames acquired at the same respiratory phase is proposed, assuming that the investigated organ reaches almost the same position at each extreme phase of the respiratory cycle. This a posteriori respiratory gating is based on a Principal Component Analysis (PCA) and integrates a priori information about the respiratory frequency.

The other main limitation of accurate quantification is the shadowing effect resulting from an incorrect attenuation correction. Attenuation is typically compensated for by applying a time-gain compensation (TGC) curve. TGC is based on the assumption that attenuation is a function only of depth. This theory is insufficient, particularly when using UCA. Several methods for automatic attenuation compensation have been developed, either assuming sufficient homogeneity in the images [4], [5], [6] or making the less constraining assumption that local attenuation and backscatter are directly related [7]. Even if these methods provide useful attenuation correction for soft tissues, no method has yet been developed to assess microbubble attenuation in vivo. There is thus a crucial need to compensate for microbubble attenuation to quantify tissue perfusion accurately. A first approach for automatic microbubble attenuation in small animal studies, based on the time-intensity variation in a highly echogenic distal area without contrast uptake, has already been proposed [8]. In this paper, an improvement of this method which introduces a regularized estimation of attenuation coefficients is presented.
II. MATERIALS AND METHODS

A. Data acquisition and preprocessing

Ten mice were studied to validate the process. Mice were anaesthetized and placed on the right flank against a heating pad. Contrast ultrasound images were acquired at a low mechanical index of 0.09 after a bolus injection of 0.1 ml of Sonovue™, using an Acuson Sequoia 512 device (Siemens Medical Solutions). The linear-array transducer (15L8w, with a central transmit frequency of 7 MHz) was mechanically fixed above the mice allowing them to breathe freely. The penetration depth was adjusted to include both kidneys and the skin-table interface at the opposite side of the mouse in the field of view. A first bolus sequence was acquired during 10 seconds, followed by a 50 seconds delay to allow microbubble concentration to reach equilibrium. After destruction of microbubbles with destructive frames at maximum power during 5 seconds, an other sequence was acquired at 25 frames per seconds during 10 seconds. Dual images in Cadence™ Contrast Pulse Sequencing (CPS) [9] and fundamental modes were acquired. Both the frame selection and the attenuation estimation methods were applied to the acquisitions in fundamental mode, which were converted from gray levels into decibels using the grayscale colorbar displayed on the monitor.

B. Methods

1) Frame selection: The first step of the proposed frame selection method was the estimation of the respiratory cycle by applying a PCA to the image sequences. This estimation was based on the assumption that not only one but several components held information related to the respiratory motion \( R(t) \), so that the latter was found as a linear combination of the five main components provided by PCA, referred to as \( C_0, ..., C_4 \) (Fig. 1(a)):

\[
R(t) = C_0(t) + \lambda_1 C_1(t) + \lambda_2 C_2(t) + \lambda_3 C_3(t) + \lambda_4 C_4(t).
\]

As values of respiratory frequencies are always in a fixed frequential window between \( f_{\text{min}} \) and \( f_{\text{max}} \) (in practice we chose \( f_{\text{min}} = 1 \text{ Hz} \) and \( f_{\text{max}} = 5 \text{ Hz} \) for an anaesthetized mouse), the respiratory motion was estimated by maximization of a frequential criteria \( C_r \) defined by:

\[
C_r = \max_\lambda \left( \int_{f_{\text{min}}}^{f_{\text{max}}} \text{FFT}(R) \right),
\]

where \( \lambda = (\lambda_1, ..., \lambda_4) \) and \( \text{FFT}(X) \) is the fast Fourier transform of vector \( X \). \( C_r \) thus corresponds to the energy of \( R \) in the predefined frequential window.

After the estimation of the respiratory motion, the local minima and maxima of respiration were extracted following the method described in [10]. Frames corresponding to each extreme phase of the respiratory cycle were selected by applying a threshold of 5%, i.e. for each respiratory period, values lower than 5% of the local minima and values upper than 95% of the local maxima were retained for generating the minima subsequence \( S_{\text{min}} \) and the maxima subsequence \( S_{\text{max}} \) respectively. These subsequences corresponded to the
end-of inspiration plane and the end-of-expiration plane, respectively (Fig. 2).

2) Microbubble attenuation estimation: A propagation model of the ultrasound pulse specific to CEUS studies was proposed [8]. This model considered that the time-intensity variation in a highly echogenic distal area without contrast uptake (i.e., the skin-table interface at the opposite side of the mouse), thereafter referred to as $R_{\text{dist}}$, provided the cumulative attenuation from the transducer to $R_{\text{dist}}$. At each time (t), the variation of intensity of each pixel of an A-line (N pixels per line) was expressed as:

$$P_k(t) = \beta_k(t) - 2 \sum_{i=1}^{k-1} \alpha_i(t), \quad 1 \leq k \leq N - 1,$$

where $\alpha_k(t)$ is the attenuation caused by microbubbles and $\beta_k(t)$ is a function of the backscatter.

Pixels beyond $R_{\text{dist}}$ were not considered, so that the last pixel of each A-line (i.e., the $N^{th}$) belonged to $R_{\text{dist}}$. As there was no contrast uptake in $R_{\text{dist}}$, the backscatter coefficient associated with the last pixel was equal to zero: $\beta_N(t) = 0$. Therefore,

$$P_N(t) = -2 \sum_{i=1}^{N-1} \alpha_i(t).$$

So for each image of the sequence, sound propagation along each A-line of N pixels was modeled by N equations (i.e., (3) and (4)) for $2(N - 1)$ unknowns (i.e., $\alpha_1(t), ..., \alpha_{N-1}(t), \beta_1(t), ..., \beta_{N-1}(t)$), which could not be solved without additional information.

This equation system was previously analytically solved by assuming the ratio $\alpha_k(t) = \beta_k(t)/\alpha_k(t)$ constant on each A-line [8]. This paper proposed an improvement of that method. A smooth constraint on $a_k(t)$ was introduced, allowing regularized attenuation coefficients to be estimated. At each time (t), $a_k(t)$ was constrained to be homogeneous on the field of view. For each A-line, $\alpha_k(t)$ and $\beta_k(t)$ coefficients were obtained by minimization of an error depending on the propagation equations (3) and (4) and on the additional constraint of homogeneity on $a_k(t)$, which was initialized by applying the analytical method [8]. The process was then iterated. For each image, $a_k(t)$ was calculated for each pixel and its mean value was used as initial value for the next iteration. The process stopped when relative variation of $a_k(t)$ was lower than 5%.

3) Reperfusion kinetics modeling: After microbubble attenuation has been estimated, quantification of contrast uptake kinetics after microbubble destruction can be investigated, even in deep areas. Two ROIs were defined in the cortex of each kidney to compare their replenishment kinetics (Fig. 3). Time intensity curve (TIC) were extracted from fundamental and CPS images and from attenuation images and then fitted by a mono-exponential model [1]:

$$\text{TIC}(t) = BI + A(1 - e^{-\beta t}).$$

$BI$ is the baseline intensity. $A$ and $\beta$ indicate the amplitude of enhancement and the velocity parameter, respectively. They correspond to three functional parameters typically estimated in destruction-replenishment studies.

### III. RESULTS

A. Frame selection

Frame selection was performed on the 10 studies on both bolus and replenishment sequences. Almost half of the frames (mean value of 48.6%, standard deviation of 7.4%) were selected on average. Typical mouse respiratory frequency values were estimated, with a mean value of 2.68 Hz and a standard deviation of 0.36 Hz.

For all the studies, no more respiratory motion could be detected on $S_{\text{min}}$ and $S_{\text{max}}$ by visual analysis. A PCA was performed on the 40 subsequences (2 subsequences per original sequence) to confirm this result. The five main components estimated for one bolus study after frame selection are shown in Fig. 1(b). While respiratory information can be easily observed on the 5 components before frame selection (Fig. 1(a)), especially on $C_2$ and $C_4$, components after frame selection are respiratory motion free. Components $C_0^*$, $C_1^*$ and $C_2^*$ seem to fit to $C_0$, $C_1$ and $C_2$. They are related with physiological kinetics. On the other hand, a component equivalent to $C_2$, that mainly contains respiratory motion information, cannot be found after frame selection.

B. Attenuation estimation

Microbubble attenuation images were estimated for each of the 20 sequences (Fig. 3). While shadowing artefacts can
Fig. 4. Example of replenishment curves in the proximal kidney (left) and in the distal kidney (right) extracted from attenuation images. The same ROIs as those on Fig. 3 were used.

be easily noticed in native images, attenuation coefficients are homogeneous within each cortex and equivalent in both cortices, which agrees with a homogeneous concentration of UCA microbubbles in these areas.

In both fundamental and CPS images, time-intensity curves in the distal kidney revealed shadowing artefacts, while equivalent attenuation coefficients are found in both cortices (Fig. 3(b)). Similar results were obtained with the replenishment kinetics (Fig. 4).

C. Reperfusion kinetics modeling

An exponential modeling was performed on the reperfusion kinetics extracted from the attenuation images (Fig. 4). For this study, values estimated for the exponential model were $A = 0.028, \beta = 0.4$ for the proximal kidney (Fig. 4(a)) and $A = 0.03, \beta = 0.61$ for the distal kidney (Fig. 4(b)). In both cases, baseline intensity was equal to zero.

IV. DISCUSSION AND CONCLUSION

The assessment of replenishment kinetics following destruction of UCA microbubbles can allow the assessment of tissue perfusion parameters. However, a preliminary step consists of removing any artefacts such as respiratory motion and attenuation caused by microbubbles.

In this study, new tools for perfusion assessment in small animal studies have been presented. The developed frame selection method allowed the selection of images acquired at the same time of the respiratory cycle by generating two subsequences for each initial acquisition. Because of the relatively high respiratory frequency of mice, time for reaching the same extreme position two consecutive times is very short in front of replenishment speed and duration. Consequently, differentiation of the different contrast enhancement phases was not altered by the lower temporal resolution of $S_{\text{min}}$ and $S_{\text{max}}$.

UCA microbubble attenuation was then estimated. The method made use of the time-intensity variation in a highly echogenic distal area without contrast uptake, which provided cumulative attenuation in the perfused area. Results were promising. Compared with the initial method [8], attenuation estimation was more accurate and homogeneous in the cortices. Shadowing artefacts were highly reduced, allowing a more faithful representation of UCA concentration than both fundamental and CPS images. The assumption of proportionality between attenuation and backscatter coefficients appeared to be reasonable in a first approximation.

Making this relationship more flexible is under investigation. The pressure-dependence of microbubble attenuation will be further investigated too. However, this dependence should not be very substantial in the experimental conditions of this study. Indeed, Tang et Eckersley [11] found that attenuation of Sonovue rapidly decreased when the frequency was away from resonance, which was the case here.

After respiratory motion and microbubble attenuation artefacts have been removed, quantitative assessment of perfusion could be investigated, even in distal areas. Results showed that while native images only allowed quantitative study in areas close to the probe, replenishment curves extracted from attenuation images could be modeled to estimate functional parameters in the whole field of view. Next step would be the study of the correlation between these parameters and perfusion parameters, such as blow flow.

The complete processing described in this paper has been implemented in a user friendly software which could be very useful for the perfusion assessment of different organs in small animal studies.

REFERENCES


