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(71) Applicants: (INSERM) INSTITUT NATIONAL DE LA SANTÉ ET DE LA RECHERCHE MÉDI-

CALE [FR/FR]; 101 rue Tolbiac, 75013 PARIS (FR). UNIVERSITÉ DE TOURS [FR/FR]; 60 rue du Plat d'Etain, BP 12050, 37020 TOURS CEDEX 1 (FR). CENTRE NATIONAL DE LA RECHERCHE SCIENTIFIQUE [FR/FR]; 3 rue Michel Ange, 75016 PARIS (FR). UNIVERSITE DE BORDEAUX [FR/FR]; 35, place Pey Berland, 33000 BORDEAUX (FR). INSTITUT POLYTECHNIQUE DE BORDEAUX (IPB) [FR/FR]; 1 rue du Docteur Schweitzer, 33400 TALENCE (FR).

(72) Inventors: DENIS DE SENNEVILLE, Baudouin; Institut de Mathématiques de Bordeaux, Université de Bordeaux, 351, cours de la Libération, 33405 TALENCE (FR). PERROTIN, Frank; Sce Gynécologie Obstétrique B, Centre Olympe de Gouges - CHU Bretonneau, 2 Bd Tonnellé, 37044 TOURS Cedex 01 (FR). ESCOFFRE, Jean-Michel; Inserm U930 Imagerie et Cerveau, Université F.Rabelais, 10 boulevard Tonnellé, 37032 TOURS (FR).

(54) Title: APPARATUS AND METHOD FOR CONTRAST-ENHANCED ULTRASOUND IMAGING

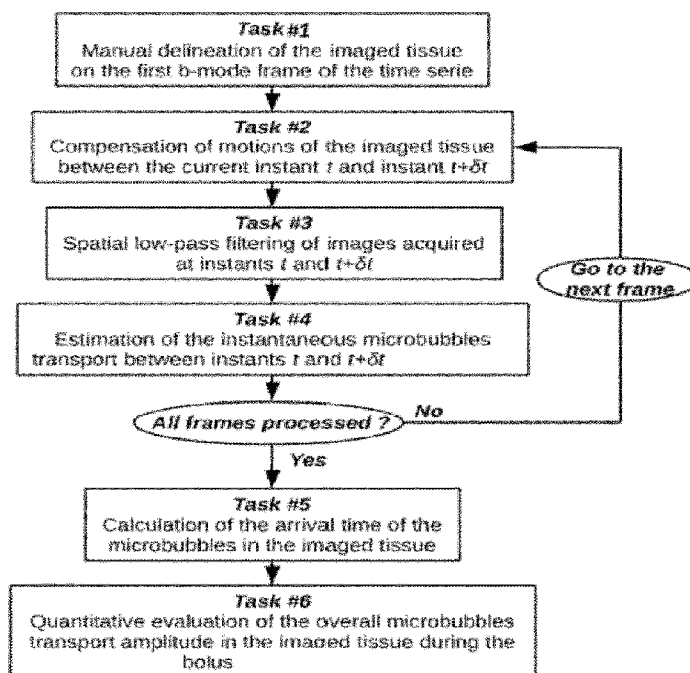


Fig. 1

(57) Abstract: The present invention relates to an apparatus and a method for contrast-enhanced ultrasound (CEUS). Known CEUS techniques remain unsatisfactory to quantify microcirculation and to provide accurate clinical diagnosis. The present invention introduces the use of a fluid dynamics model for the analysis of dynamic contrast-enhanced ultrasound (DCEUS). The efficiency of the invention was evaluated both in-vitro, for the quantitative estimation of microbubble flow rates, and in-vivo, for the classification of placental insufficiency (control vs. ligature) of pregnant rats. This experimental evaluation demonstrates that the invention outperforms the performance of known CEUS techniques. Examples of applications might include but are not limited to the quantitative analysis of the vascularisation of an organ such as the placenta or a tumour, the clinical diagnostic of obstetrical disorders, or the assessment of vascularisation in abnormal placental perfusion.



BOUAKAZ, Ayache; Inserm UMR U930, 10 boulevard
Tonnellé, 37032 TOURS (FR).

(74) **Agent: CABINET GERMAIN & MAUREAU**; 31-33 rue
de la Baume, 75008 PARIS (FR).

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Apparatus and method for contrast-enhanced ultrasound imaging

Field of the invention

The present invention relates to contrast-enhanced ultrasound, also referred to in the abbreviated form as "CEUS".

Background of the invention

5 Contrast-enhanced ultrasound is a non-invasive imaging technique that has been used extensively in blood perfusion imaging of various organs. This modality is based on the acoustic detection of gas-filled microbubble contrast agents used as intravascular flow tracers. These contrast agents significantly increase the ultrasound scattering of blood, thus allowing the visualization and the assessment of
10 microcirculation (i.e., blood velocity, blood volume fractions) commonly undetectable by Doppler ultrasound (DUS). The rheology of microbubbles in the microcirculation is similar to that of red blood cells, thus demonstrating the microbubbles do not interfere with hemodynamics and have a good safety profile in both cardiac and abdominal ultrasound applications.

15 To achieve this objective, computer tools are required for the spatio-temporal and quantitative assessment of organ perfusion. Practically, once the bolus injection of the ultrasound contrast agent is performed, the enhancement in the vascular compartment is screened using contrast-specific ultrasound-imaging sequences. Recent efforts are focused on the quantification of parameters related to this
20 enhancement and on the use of these latter as indicators for several pathological conditions. To this end, the average intensity within a region of interest (ROI) is analyzed as a function of time, in the form of a time-intensity curve (TIC). The wash-in and wash-out of the contrast agent within the ROI are subsequently quantified by calculating amplitude and temporal parameters, such as the peak enhancement (PE),
25 the wash-in rate (WiR), the time to peak (TTP), the area under the curve during wash-in (WiAUC).

One of the most recent applications in this field is quantitative assessment of placental perfusion. In contrast to other circulatory beds of the systemic circulation where blood traverses from arteries via capillary beds to veins, placental circulation
30 acts as an open system with the spiral arteries opening in a large lake of blood, namely the intervillous space, with little or no impedance to blood flow. That way, using dedicated small animal ultrasound scanner (Vevo®2100) and TIC-based post-processing software (Vevo®CQ), a recent study reported on real-time monitoring of microbubble perfusion in the uteroplacental unit, across the gestation of rats (C. J.

Arthuis, A. Novell, J. M. Escoffre, F. Patat, A. Bouakaz, and F. Perrotin, *“New insights into uteroplacental perfusion: quantitative analysis using doppler and contrast-enhanced ultrasound imaging”* *Placenta*, vol. 34, no. 5, pp. 424–31, 2013; hereinafter referred to as “Arthuis et al. 2013”). The enhancement of ultrasound contrast in the mesometrial triangle and in the maternal spiral arteries was visualized 1s after intravenous injection of contrast agents. These agents were then detected through the maternal vessel in the centre of the placenta to the chorionic plate. The perfusion of these agents slowly expanded in the placenta to the basal plate. During the gestation of rats, the blood flow increased from days 14 to 20 in the mesometrial triangle and the placenta. In other recent studies, the CEUS of uteroplacental perfusion in nonhuman primates were conducted using clinical scanners (e.g., Sequoia Systems) and custom-designed CEUS analysis program or image post-processing software were provided by the manufacturers (VueBox®, Bracco, Milan, Italy). Keator et al. reported that CEUS was able to detect the primary placental disc and underlying vessels two days earlier than DUS identified endometrial thickening (C. S. Keator, J. R. Lindner, J. T. Belcik, C. V. Bishop, and O. D. Slayden, *“Contrast-enhanced ultrasound reveals real-time spatial changes in vascular perfusion during early implantation in the macaque uterus”* *Fertil Steril*, vol. 95, no. 4, pp. 1316–21 e1–3, 2011). Using CEUS, spatial differences in vascular perfusion between the endometrium, myometrium and endometrial-myometrial zone were observed and quantified. Thus, myometrium displayed a higher blood flow than that detected endometrium and junctional zone.

However, using the TIC-based approach, the quantification is typically hampered by two main reasons: first, the exact relationship between the acquired ultrasound-signal and the local microbubbles concentration is hardly achievable. Second, the quantification intrinsically solely relies on temporal intensity variations, the explicit spatial transport of the contrast agent being left out.

Object of the invention

It is an object of the present invention to provide an apparatus and a method for contrast-enhanced ultrasound imaging without the drawbacks associated with the prior art.

Summary of the invention

In that respect, the present invention relates to an apparatus for contrast-enhanced ultrasound imaging, comprising:

- injection means for injecting a contrast agent within a region of interest,

- an ultrasound system for propagating ultrasound waves within said region of interest and detecting ultrasound waves backscattered within said region of interest,
- imaging means for producing a temporal succession of images according to said backscattered ultrasound waves, each image being representative of a spatial configuration of said contrast agent within said region of interest.

The contrast agent may preferably comprise gas microbubbles, preferably inert gas microbubbles.

In a non-limitative example, the region of interest may be located within an organ such as the placenta.

Said images are preferably B-mode images.

According to the invention, this apparatus comprises transport calculating means arranged to determine a spatial transport of the contrast agent from successive images.

According to an aspect of the invention, the transport calculating means may comprise numerical resolution means for numerical resolution of the following transport equation:

$$I_t + \vec{V} \cdot \vec{\nabla} I = 0$$

where I is a grey level intensity on said images, I_t is a partial temporal derivative of I between successive images, $\vec{\nabla}$ is a spatial gradient operator, $\vec{\nabla} I$ is the spatial gradient of I , and \vec{V} corresponds to said spatial transport between those successive images.

Practically, $\vec{\nabla} I = (I_x, I_y)$ for 2D images, and $\vec{\nabla} I = (I_x, I_y, I_z)$ for 3D images, $I_{x,y,z}$ being the spatial partial derivatives of I .

The spatial transport calculated with this transport equation remains identical whichever function is used to convert the grey levels of the images. It is therefore not necessary to convert the grey levels of the images.

In this transport equation, the transport is calculated taking simultaneously into account both spatial and temporal dimensions.

According to another aspect of the invention, the apparatus may comprise resetting means arranged to compensate relative movements between said ultrasound system and said region of interest.

In an embodiment, said resetting means may be arranged to compensate relative movements between said ultrasound system and said region of interest using

a gradient driven descent algorithm maximizing an inter-correlation coefficient between successive images.

According to another aspect of the invention, the apparatus may comprise a spatial low-pass filter arranged to filter said images.

5 Said spatial low-pass filter may preferably be a spatial low-pass Butterworth filter.

Such a spatial low-pass filter allows mitigating the impact of local image structures on the spatial transport determined using said transport equation.

The invention also relates to a method for contrast-enhanced ultrasound
10 imaging, this method comprising:

- propagating ultrasound waves within a region of interest and detecting ultrasound waves backscattered within said region of interest,
- producing a temporal succession of images according to said backscattered ultrasound waves, each image being representative of a
15 spatial configuration of said contrast agent within said region of interest.

According to the invention, this method comprises determining a spatial transport of the contrast agent from successive images.

According to an aspect of the invention, the step of determining said spatial transport may comprise numerical solving of the following transport equation:

20
$$I_t + \vec{V} \cdot \vec{\nabla} I = 0$$

where I is a grey level intensity on said images, I_t is a partial temporal derivative of I between successive images, $\vec{\nabla}$ is a spatial gradient operator, $\vec{\nabla} I$ is the spatial gradient of I , and \vec{V} corresponds to said spatial transport between those images.

25 According to another aspect of the invention, the method may comprise compensating relative movements between said ultrasound system and said region of interest using a resetting means.

More specifically, the step of compensating said relative movements may comprise maximizing an inter-correlation coefficient between successive images using a gradient driven descent algorithm.

30 According to another aspect of the invention, the method comprises filtering said images using a spatial low-pass filter.

The invention also relates to an image processing tool which implements such a method.

The invention also relates to a device for contrast-enhanced ultrasound imaging, this device comprising such an image processing tool.

Brief description of the drawings

The above and others features, details and advantages of the present invention will become apparent from the following detailed description and the accompanying drawings in which:

- Figure 1 is a diagram representing different tasks of a method according to the invention;
- Figures 2a to 2d show results obtained from in-vitro experiments:
 - Figures 2a and 2b show typical example of DCEUS images of a silicone tubing undergoing a flow rate of 4.8 mL/min with bubble dilutions of 1:1000 (Fig. 2a) and 1:2000 (Fig. 2b) – for an easier visualization, only arrows associated to pixels with DCEUS intensity higher than 10 % of the maximal saturation value are displayed;
 - Figures 2c and 2d show DCEUS-based transports flow rate estimates, as a function of the pump flow rate, independently replicated ($N = 3$) for bubble dilutions of 1:1000 (Fig. 2c) and 1:2000 (Fig. 2d). The dashed line represents the linear regression line; its equation and the correlation coefficient are indicated in the text box in the bottom-right of each graph;
- Figures 3a to 3h provide example of microbubble transport estimates from the DCEUS of two rats using the proposed approach, obtained from in-vivo experiments. Figures 3a to 3d display results associated to a control rat and figures 3e to 3h display results associated to a ligature rat. Figures 3a and 3e display a B-mode image with superimposed the manually drawn mask (dashed line) underlying the placenta. Figures 3b and 3f, 3c and 3g, and 3d and 3h display DCEUS images acquired 10 s, 15 s and 20 s after the determined microbubble arrival in the placenta, respectively. The corresponding transport field estimates are superimposed. In these results, ΔT , δt , α and f_c were fixed to common typical values of 60 s, 3 s, 0.1 and $f_0 / 16$, respectively;
- Figures 4a and 4b show temporal evolution of the bolus velocity (averaged over the ROI encompassing the placenta, and weighted by the amount of grey level intensity in the DCEUS image) estimated for each control (Fig. 4a) and ligature (Fig. 4b) rat, obtained from in-vivo experiments;
- Figures 5a to 5h provide assessment of proposed and TIC-based approaches for the classification of the two rat populations (i.e. control and ligature):

- Figure 5a: boxplots of overall microbubbles velocities estimated over each rat populations using the proposed approach;
 - Figures 5b to 5e: distributions of PE, WiR, TTP and WiAUC estimated over the placenta for the two rat populations using the existing TIC-based approach. Median is shown by the central mark, the first and the third quartile are reported by the edges of the box, the whiskers extend to the most extreme time points which are not considered as outliers, and outliers are individually plotted in red;
 - Figure 5f: ROC curve obtained using γ ;
 - Figure 5g: ROC curves obtained using PE and WiR;
 - Figure 5h: ROC curves obtained using TTP and WiAUC.
- Figures 6a and 6b provide an analysis of the impact of the temporal window size (ΔT) and temporal derivative step (δT) on the overall results obtained in experiments. The bolus velocity was estimated for various values of ΔT and δT (α and f_c being fixed to typical values of 0.1 and $f_0 / 16$, respectively). The significance of the difference between bolus velocities obtained for the two rat populations (i.e. ligature and control) is reported using the p-value of an unpaired Mann-Whitney U test (Fig. 6a) and the AUROC (Fig. 6b);
- Figures 7a and 7b provide an analysis of the impact of the cut-of-frequency f_c of the low-pass spatial image filtering and of the regularization parameter α of the numerical scheme on the overall results:
- Figure 7a: bolus velocities estimated for various values of f_c (ΔT and δT and α and being fixed to typical values of 60 s, 3 s and 0.1, respectively);
 - Figure 7b: bolus velocities estimated for various values of α (ΔT , δT and f_c and being fixed to typical values of 60 s, 3 s and $f_0 / 16$, respectively). The significance of the difference between bolus velocities obtained for the two rat populations (i.e., ligature and control) is reported using the AUROC.
- Identical or similar elements are marked with identical reference signs in all of the figures.

Detailed description of the invention

The invention provides an apparatus and a method for contrast-enhanced ultrasound imaging.

- 35 The apparatus comprises injection means for injecting a contrast agent within a region of interest. For example, Definity® microbubbles (Lantheus Medical Imaging, Billerica, MA) can be injected within a tissue or an organ such as the placenta.

The apparatus also comprises an ultrasound system. This ultrasound system is arranged for propagating ultrasound waves within said region of interest. This ultrasound system is also arranged for detecting ultrasound waves that are backscattered within said region of interest. In this example, the ultrasound system comprises a MS-250 probe (21 MHz centre frequency, 13-24 MHz bandwidth, 75 μm axial and 165 μm lateral resolutions) and a Vevo®2100 ultrasound scanner (VisualSonics Inc., Toronto, Canada). The ultrasound system allows the method of the invention carrying out a step of propagating said ultrasound waves within said region of interest and a step of detecting ultrasound waves backscattered within said region of interest.

The apparatus also comprises imaging means for producing a temporal succession of images according to said backscattered ultrasound waves. Each image is representative of a spatial configuration of said contrast agent within said region of interest. Imaging means allows the method of the invention carrying out a step of producing said temporal succession of images according to said backscattered ultrasound waves. Said images are preferably B-mode images.

Basically, in this example, the method of the invention takes advantage of:

- simultaneous acquisition of dynamic contrast-enhanced ultrasound, also referred to in the abbreviated form as “DCEUS”, and
- B-mode images.

In particular, the instantaneous apparent microbubble bunches transport is estimated using a continuity equation of fluid dynamics applied on each acquired DCEUS image (see below). B-mode images are dedicated to the manual delineation of the imaged tissue and to the estimation of possible periodic, spontaneous or drift displacements of the tissue. The latter may be induced either by physiologic activity (breathing or peristaltic) or by motion of the ultrasound probe.

Figure 1 illustrates one way of carrying out some steps of the method of the invention, and describes more specifically six particular steps, or tasks, described below. Those steps or tasks are referred to as “task #1”, “task #2” ... “task #6”.

In task #1, said region of interest encompassing the imaged tissue is manually drawn on a B-mode image, and a binary mask (noted Γ) is constructed. Pixels of the image inside the mask have a value of one, and outside a value of zero.

The microbubble transport over the imaged tissue between the current processed DCEUS image (let t be the corresponding acquisition time) and the DCEUS image acquired at $t + \delta t$ (δt being a pre-defined interval of time) can then be estimated over the successive tasks #2 to #4.

In task #2, physiologic activity and motion of the ultrasound probe are compensated on the DCEUS images, since those relative movements may hamper the estimation of the apparent microbubble transport. In other words, task #2 is intended to compensate relative movements between said ultrasound system and said region
5 of interest.

Using the proposed microbubble transport estimation method, perturbations occur predominantly when the true transport field violates the physical model used for its estimation. Indeed, any temporal change in image intensity is likely to be attributed to microbubble transport. DCEUS image intensity changes can not only be
10 due to contrast change during bolus passage but also to the physiological activity.

In this example, the motion estimation process between t and $t + \delta t$ is a gradient driven descent algorithm maximizing an inter-correlation coefficient and applied on B-mode images, assuming a translational displacement restricted to the binary mask Γ encompassing the imaged tissue. The estimated spatial transformation
15 is subsequently employed to compensate motions of the imaged tissue on the DCEUS image acquired $t + \delta t$.

In task #3, a spatial low-pass filter is applied on each DCEUS image in order to mitigate the impact of local image structures on the employed continuity equation of fluid dynamics.

In this example, a spatial low-pass Butterworth filter (order 1) is applied on DCEUS images acquired at instants t and $t + \delta t$ in order to mitigate the impact of local image structures on the estimated microbubble transport, due to the differential nature of terms involved in the continuity equation employed afterwards (see below).
20

In task #4, the instantaneous apparent pixel wise microbubble transport is estimated. The step of the temporal derivative is referred to as δt . The transport between DCEUS images acquired at instants t and $t + \delta t$ is estimated.
25

In order to estimate the microbubble bunch transport (noted \vec{V}) occurring during the DCEUS between two close instants (i.e. t and $t + \delta t$), the following continuity equation can be employed in homogeneous environment: $I_t + \vec{\nabla} \cdot (I\vec{V}) = 0$,
30 where I denotes the grey level intensity on DCEUS images and I_t the partial temporal derivative of I . The left part of this equation is composed by a transient term (I_t) and a convection term ($\vec{\nabla} \cdot (I\vec{V})$), which stand for any temporal and spatial grey intensity variations, respectively. The estimated transport field \vec{V} thus accounts for spatio-temporal grey level intensity variations occurring during the dynamic imaging
35 sequence.

In order to associate any spatio-temporal variations of pixel intensities to “transport”, without additional assumptions carried on the link between the microbubble dose concentration and grey level intensities, the contribution associated with the divergence of velocity field (i.e. $I\vec{\nabla}\cdot\vec{V}$) is left out. This simplifies the mass continuity equation in the latter equation to the following transport equation:

$$I_t + \vec{V}\cdot\vec{\nabla}I = 0$$

To sum up, the method of the invention comprises determining a spatial transport of the contrast agent from successive images. In this example, the step of determining said spatial transport comprises numerical solving of the latter transport equation.

The transport model of the latter transport equation is intrinsically under-determined, and thus leads to an ill-conditioned numerical scheme. The seek transport field \vec{V} can thus be estimated on a pixel-by-pixel basis through the following minimization process:

$$\arg \min_{\vec{V}} \int_{\Omega} |I_t + \vec{V}\cdot\vec{\nabla}I| + \alpha \left(\|\vec{\nabla}u\|_2^2 + \|\vec{\nabla}v\|_2^2 \right) d\vec{r}$$

where $\Omega \subseteq \mathfrak{R}^2$ is the image coordinates domain, (u, v) the estimated pixel wise transport vector components, and $\vec{r} \in \Omega$ the spatial location. The minimized functional accounts for the two following additive contributions:

- A data fidelity term (left part of the integral in the latter equation) which optimizes, through a L^1 norm, the transport model of said transport equation. Given that a L^1 penalizer is employed, transient variations act identically regardless the amount of grey level intensity.
 - A regularization term (right part of the integral in the latter equation) designed to provide a sufficient conditioning to the numerical scheme.
- The regularization term is given by $\|\vec{\nabla}u\|_2^2 = u_x^2 + u_y^2$ and $\|\vec{\nabla}v\|_2^2 = v_x^2 + v_y^2$ with u_x, u_y, v_x and v_y being the partial spatial derivatives of u and v , respectively. Physically, this regularization term assumes that the transport between neighbouring pixels is moderate.

α is a pre-defined weighting factor designed to link these two contributions.

In order to render the latter equation differentiable, $|s| = |I_t + \vec{V}\cdot\vec{\nabla}I|$ can be replaced by $\psi(s) = \sqrt{s^2 + \varepsilon^2}$ with $\varepsilon = 10^{-3}$.

Then, by applying the Euler-Lagrange equations on the latter equation on a pixel-by-pixel basis, one can derive the following system of equations for each $\vec{r} \in \Omega$:

10

$$\begin{cases} \frac{s}{\psi(s)} I_x - 2 \alpha \Delta u = 0 \\ \frac{s}{\psi(s)} I_y - 2 \alpha \Delta v = 0 \end{cases}$$

where $I_{x,y}$ denotes the partial spatial derivatives of I , and $\Delta(\cdot)$ the Laplacian operator. Neumann boundaries conditions can be employed.

From here, we have a set of $2 * |\Omega|$ non-linear equations with common
5 unknowns u and v .

$\Delta(\cdot)$ can be approximated in the discrete domain with $\Delta u = \bar{u} - u$, \bar{u} being the 3×3 local average of u . That way, two additional implicit linear contributions (along u and v , respectively) are obtained. This approximation allows linearizing the system as follows:

10
$$\begin{cases} A u + B v = C \\ D u + E v = F \end{cases}$$

with $A = I_x^2 + 2 \alpha \psi(s)$, $B = I_x I_y$, $C = 2 \alpha \psi(s) \bar{u} - I_x I_t$, $D = I_x I_y$, $E = I_y^2 + 2 \alpha \psi(s)$, and $F = 2 \alpha \psi(s) \bar{v} - I_y I_t$.

Solutions u and v can be found through a fixed-point scheme for which A, B, C, D, E and F are explicitly calculated, as follows:

15
$$\begin{cases} u^{n+1} = \bar{u}^n - I_x \frac{\bar{u}^n I_x + \bar{v}^n I_y + I_t}{I_x^2 + I_y^2 + 2\alpha\sqrt{(I_t + I_x u^n + I_y v^n)^2 + \varepsilon^2}} \\ v^{n+1} = \bar{v}^n - I_y \frac{\bar{u}^n I_x + \bar{v}^n I_y + I_t}{I_x^2 + I_y^2 + 2\alpha\sqrt{(I_t + I_x u^n + I_y v^n)^2 + \varepsilon^2}} \end{cases}$$

where $n + 1$ denotes a new iteration.

Note that the fixed-point scheme of this latter equation is designed to get the maximum benefit from implicit terms that can be expressed linearly along u and v , while leaving an explicit expression for \bar{u} , \bar{v} and $\sqrt{(I_t + I_x u^n + I_y v^n)^2 + \varepsilon^2}$. The
20 residual $\|\vec{v}^{n+1} - \vec{v}^n\|_2$ can be compared to a maximal allowed tolerance of 10^{-3} in order to ensure the convergence.

The differential nature of terms involved in said transport equation hampers the estimation of transport of high amplitude. Displacements that are larger than the size of one pixel cannot be estimated. In order to overcome this limitation, it can be
25 adopted a coarse-to-fine strategy as described in the following paper: I. Pratikakis, C. Barillot, P. Hellier, and E. Memin, "Robust multiscale deformable registration of 3d ultrasound images" International Journal of Image and Graphics, vol. 3, no. 4, pp. 547–565, 2003). According to this strategy, it can be made iteration of the registration

algorithm from a 16-fold down sampled image step by step to the original image resolution.

In addition, an iterative refinement of the transport estimates can be performed within each resolution. This implies running the algorithm several times at the same resolution, initializing the displacement field at the current run of the algorithm with the displacement field that resulted during the previous run. In this manner, the stability of the numerical scheme can be improved and at the same time a better quality of the estimates can be obtained. It is considered that the numerical scheme in the latter equation is converging when the average variation of the transport magnitude from one iteration to the next is smaller than 10^{-3} pixels.

In task #5, the instant time of the microbubble arrival is estimated, which for practical reasons is likely to vary from one experiment to the other.

For this purpose, each dynamically acquired image may be iteratively enumerated and the average pixel intensity over the binary mask Γ encompassing the imaged tissue can be calculated. Once, this value exceeded a typical pre-defined threshold of 1 % of the maximal intensity saturation value, the associated time instant is considered as the arrival time (referred to as t_0 throughout the present document).

In task #6, the quantitative microbubble transport amplitude during the bolus can be assessed.

At this stage, a set of pixel wise transport fields is obtained, each one being associated to each ultrasound image. The spatio-temporal averaged microbubble transport (noted γ), over the imaged tissue and during a time window (covering a duration ΔT , starting from the bolus arrival time t_0) can be then calculated as follows:

$$\gamma = \frac{\sum_{t=t_0}^{t_0+\Delta t} \sum_{\vec{r} \in \Gamma} I(\vec{r}, t) |\vec{V}(\vec{r}, t)|}{\sum_{t=t_0}^{t_0+\Delta t} \sum_{\vec{r} \in \Gamma} I(\vec{r}, t)}$$

Note that the module of each transport vector $\vec{V}(\vec{r}, t)$ is weighted by the amount of grey level intensity in the DCEUS image $I(\vec{r}, t)$. That way, similar γ values are obtained for scenarios with identical microbubble transports, but various pixel intensities and/or various amounts of pixels exhibiting microbubbles. $\vec{V}(\vec{r}, t)$ is converted in mm/s beforehand so as to express γ in a common metric unit.

The invention also relates to an image processing tool which implements such a method.

The invention also relates to a device for contrast-enhanced ultrasound imaging, this device comprising such an image processing tool.

The invention also relates to a computer program and/or an image post-processing software comprising instructions for carrying out the method of the invention.

5 Experimental evaluation

An experimental evaluation was performed using an Intel 2.5 GHz i7 workstation (8 cores) with 32 GB of RAM. The implementation was performed in C++ and parallelized through multi-threading.

Definity® microbubbles (Lantheus Medical Imaging, Billerica, MA) were used as
10 ultrasound contrast agent. Definity® microbubbles are a second-generation clinically approved contrast agent, composed of octafluoropropane gas encapsulated in a thin and flexible monolayer of PEGylated phospholipids. The mean diameter ranges from 1.1 to 3.3 μm . Definity® microbubbles were prepared according to the manufacturer's instructions. Briefly, a single vial of Definity® was warmed to room temperature and
15 then was activated using a Capmix® device (3M-ESPE, Cergy-Pontoise, France) for the full 45-second activation cycle.

A high frequency ultrasound scanner was used. More specifically, a MS-250 probe (21 MHz centre frequency, 13-24 MHz bandwidth, 75 μm axial and 165 μm lateral resolutions) connected to the Vevo®2100 ultrasound scanner (VisualSonics
20 Inc., Toronto, Canada) was used to acquire in-vitro and in-vivo images.

Concerning in-vitro experiments, a laboratory-made flow system was an open circuit consisting of 3.59 mm internal diameter flexible silicone tubing, through which a suspension of Definity® microbubbles circulated. The tubing flow system was submerged in a degassed water tank. One end was connected to reservoir filled with
25 degassed water, in which Definity® microbubbles were diluted in physiological serum. The diluted microbubbles solution was subsequently delivered by a peristaltic pump (MCP Process IP65, Cole-Parmer GmbH, Wertheim, Germany). The centre of the flow tube was positioned at a distance of 21 mm from the MS-250 probe. For each experimental condition, a video clip of 30 s was recorded at 20 frames per second
30 (pixel size= $0.027 \times 0.027 \text{ mm}^2$) using the Vevo®2100 scanner.

After image acquisition, microbubbles perfusion was quantitatively analyzed as follows. The DCEUS-based microbubble transport (γ) was computed for four regularly sampled flow rates delivered by the peristaltic pump i.e., 1.2, 2.4, 3.6 and 4.8 mL/min). All experiments were independently replicated ($N = 3$). A correlation
35 coefficient (R^2) and the γ -intercept of a linear regression were then calculated in order to assess quantitatively the relationship between the DCEUS-based microbubble

transport and the delivered flow rate. The complete evaluation process was repeated for two different microbubble concentrations: 1:1000 and 1:2000 (diluted in physiological serum).

Figures 2a to 2d report typical findings obtained using the employed in-vitro experimental platform. A temporal window size $\Delta T = 15$ s was employed here. In order to supply as much as possible the high inherent velocity of the microbubble in the silicone tubing, the transport was estimated between consecutive images. In other terms, since the imaging frame-rate was equal to 20 Hz, the temporal derivative step δt was set to a value of $1/20$ s. High instabilities of the numerical schemes were observed using the proposed implementation for $\alpha < 0.3$. A value of $\alpha = 0.3$ was therefore employed to generate the presented results. The cut-of-frequencies f_c of the spatial low-pass filtering of DCEUS images was set to the original image sampling frequency f_0 (i.e. no image filtering was applied) in order to mitigate the deterioration of the apparent local shapes of microbubble bunches. Figures 2a and 2b display typical examples of DCEUS-based microbubble transport fields (displayed in form of vector fields) estimated using the proposed methodology for microbubble dilutions of 1:1000 (Fig. 2a) and 1:2000 (Fig. 2b). Vectors of homogeneous amplitude and oriented along a common horizontal direction (accurately matching that of the delivered microbubble flow) are observable for each tested bubble dilution. As shown in Figures 2c and 2d, the proposed estimated DCEUS-based transport amplitude (γ) is a good quantitative indicator of the flow rate within the tube, for the two tested bubble concentrations ($R^2 > 0.96$, γ -intercept < 1 mm/s). It can be noticed that, for a bubble dilution 1:2000, the precision of γ diminished, especially for low delivered flow rates (i.e. 1.2 and 2.4 mL/min). The DCEUS-based microbubble estimation process was in this case impaired by few local image contrasts (see the reduced US contrast within the tube in Fig. 2b, as compared to Fig. 2a), which in turn hampered occasionally the conditioning of the employed numerical scheme.

Concerning in-vivo experiments, all procedures were performed in accordance with the ethical guidelines and were approved by the French Committee (n°19) for Animal Care and Ethics in Animal Experiments (APAFIS#3879-2016020117195710). Eleven pregnant Sprague-Dawley rats were purchased from Janvier Labs CERJ (Le Genest Saint-Isle, France). They were maintained at constant room temperature with 12h light cycle. The rats were 10-12 weeks old at the beginning of the experiments, weighing in average 377 ± 17 g. A ligature of the uterine pedicle was performed to induce an IUGR on the seventeenth days of gestation. Under gaseous anaesthesia (Aerrane®r, Baxter, Deerfield, IL), the pregnant rats were placed on a thermostatically controlled pad in order to maintain body temperature at about 37°C. Before the

surgery, an analgesia was achieved by intraperitoneal (i.p.) injection of buprenorphin (0.05 mg/kg; Buprecare®, Axience SAS, Pantin, France). After shaving of the abdomen, a surgeon performed a midline laparotomy incision with sterile precautions. The number of implantation sites was checked in each uterine horn. A 5-0 Ethilon nylon suture (Ethicon, Somerville, NJ) was placed around the uterine vessels near the lower end of one horn. The non-ligated horn served as a control. The abdominal incision was repaired in layers using standard surgical procedures. Five hours later, a single 5-mg/kg buprenorphine was intraperitoneally administered to manage postoperative pain.

10 Under gaseous anaesthesia (Aerrane®), the pregnant rats were placed on a thermostatically controlled pad and their abdomens were shaved before CEUS examination on the nineteenth day of gestation. A 24-gauge catheter was placed in a tail vein to inject Definity® contrast microbubbles. Ultrasound B-scans were used to image foetal-placental unit in cross-section. A bolus of 200 µL of contrast agent (0.5 mL/kg) was intravenously injected. Subsequently, a video clip of 150 s was recorded at 15 10 frames per second (pixel size = 0.035 × 0.035 mm²) to investigate the utero-placenta perfusion.

After image acquisition, utero-placental perfusion was quantitatively analyzed from the DICOM video data using the proposed methodology. The perfusion was also 20 analyzed using the existing TIC-based approach. For this purpose, we used the CEUS analysis software Vevo-CQ™, which is directly integrated in the ultrasound scanner. The four following parameters were extracted from the TIC: PE, WiR, TTP and WiAUC. The following paper provides a complete description of the above mentioned TIC-based parameters: F. Tranquart, L. Mercier, P. Frinking, E. Gaud, and M. Arditi, 25 *“Perfusion quantification in contrast-enhanced ultrasound (CEUS)–ready for research projects and routine clinical use”* *Ultraschall Med*, vol. 33 Suppl 1, pp. S31–8, 2012.. Note that, for both our approach and tested TIC-based methods, the DCEUS analysis was performed on an identical ROI encompassing the imaged tissue (see Task #1 in Fig. 1) in order to clarify the benefits on the final results.

30 For the two rat populations, all indicators (i.e. the overall microbubble transport amplitude γ as well as the four parameters extracted from the TIC) were expressed as medians and interquartile ranges, and were compared by using the unpaired Mann-Whitney U test. The results were considered significant when the p-value was lower than 0.025. In addition, considering each indicator as a classification 35 criterion, the performance of a binary classifier system was also assessed using a “receiver operating characteristic” (ROC) curve. The area under the ROC curve (AUROC) was subsequently computed: while a binary classifier acts like a completely

random guess for AUROC = 0.5, the best possible prediction method would yield to a value of 1.

Figures 3a to 3h report typical microbubble transport fields obtained from two dynamic contrast-enhanced experiments (one clip was selected in the control and one in the ligature population, respectively). For each rat, the mask encompassing the placenta, manually drawn in Task #1, is displayed in Figures 3a and 3e. The estimated microbubble transports are visually larger for the control population (Figs. 3b-d) as compared to the ligature population (Figs. 3f-h) (note that the estimated vector fields follow visually the dynamic contrast-enhancement in associated DCEUS images). This observation is confirmed in Figures 4a and 4b: the instantaneous transport of the apparent microbubble was larger for non-ligature rats, especially for images acquired during the first 10 seconds following the bolus arrival in the placenta. The spatio-temporally averaged microbubble transport amplitude (i.e., coefficient γ) was significantly higher (p-value = 1.1×10^{-3}) in the control group than in the ligature group (see Fig. 5a). The associated AUROC, displayed in Figure 5f, was equal to 0.9343. On the other hand, none of tested TIC-based parameters led to a significant difference between control and ligature populations (p-values obtained using PE, WIR, TTP and WiAUC were equal to 0.098, 0.348, 0.972 and 0.285, respectively), as shown in Figures 5b to 5e. The best achievable AUROC was obtained using PE (AUROC=0.72), as shown in Figures 5g and 5h.

Figures 6a and 6b and figures 7a and 7b analyze the sensitivity of the proposed methodology against the four following calibration parameters:

- The duration of the observation window ΔT , which determines the time interval after bolus arrival;
- The step of the temporal derivative δT , which determines the time interval for the estimation of the instantaneous microbubble transport;
- The cut-of-frequency f_c of the low-pass filter applied on the input DCEUS images. In the present document f_c is expressed as a fraction of the original image sampling frequency f_0 ;
- The regularization of the transport equation α , which infers the stability of the numerical scheme.

It can be observed in Figures 6a and 6b that γ (i.e., the overall microbubble transport amplitude) is a good classification criterion of the two rat populations for all tested temporal window size (ΔT) and temporal derivative step (δt) (p-value constantly $< 10^{-2}$, AUROC > 0.89), especially for low values of ΔT (i.e., < 20 s). Figure 7a shows that the performance of γ as a classification criterion for the two rat populations improved when reduced cut-of-frequencies f_c were employed in the spatial low-pass filtering of DCEUS images. This justifies the choice to set $f_c = f_0 / 16$ in

the previously shown results. In addition, it is important to report that high instabilities of the numerical schemes were observed for $\alpha < 0.1$. This justifies the choice to set a value of 0.1 as a default parameter for α in the scope of this experiment. Figure 7b shows that comparable classification performances were assessed for an α value fifteen times higher.

It appears from this experimental evaluation that the estimated transport is highly correlated with the microbubble velocity within our in-vitro system, for the delivered flow rates, whatever the tested microbubble concentration (see Figs. 2c and 2d). Moreover, the proposed classification criterion γ , estimated from DCEUS echography, was demonstrated to be a good binary classification criterion for ligature/non-ligature rat placentas. In particular, it outperformed TIC-based methods which provided best results using the parameter PE (AUROC = 0.72, as shown in Figs. 5g and 5h), but remained however well below our fluid dynamic approach (see Fig. 5f).

Conceptually, the proposed fluid dynamic approach analyses pixel wise microbubble velocity fields (both spatial and temporal derivative of the DCEUS image intensity are simultaneously involved in said transport equation). While existing compartment models or TIC-based approaches intrinsically solely rely on temporal intensity variations (the latter being averaged within a mask encompassing the organ of interest), the method of the invention performs a simultaneous spatio-temporal analysis. Consequently, this method provides an additional information for the kinetic analysis of DCEUS (see the quantitative estimates on the delivered flow rate in Figs. 2c and 2d, which are not accessible when using the TIC-based approach together with a continuous infusion of microbubbles). That being said, while a difficult conversion from grey level intensity to microbubble concentration is a common necessary prerequisite when using the existing TIC-based approach, this task is circumvented here.

This advantage of the proposed methodology is illustrated on the presented in-vitro experiment (see Figs. 2a-d), for which quantitative indicators of the flow rate could be accurately estimated, whatever tested microbubble concentration. Practically, microbubbles generate a hyper-intense signal, which saturates grey level intensities. The histogram of the DCEUS images within the imaged organ consistently depicts two main peaks: while high values correspond to pixels where microbubbles are present, low values are associated to the others. "Binary" DCEUS images are thus obtained which allows the estimation of a wave front displacement using the proposed methodology, regardless microbubble concentration considerations.

The proposed methodology is sensitive to the four above-mentioned calibration parameters. Figures 6a and 6b and figures 7a and 7b demonstrate that a compromise has to be found for each parameter, as follows. First, while the duration of the observation window (ΔT) must be high enough to screen a significant microbubble bolus (around 10 seconds were found to be sufficient), this value must be also reduced in order to diminish potential long-term motion bias (e.g., motion of the ultrasound probe, foetus motion...). Second, while the step of the temporal derivative (δt) must be high enough to observe a significant microbubble transport between instants t and $t + \delta t$ (several seconds provided good results in the employed experimental setup), this value must also be reduced in order to diminish the potential occurrence of physiological motions between the same instants. Third, while the cut-of-frequency of a low-pass filter applied on the input images (f_c) has to be low enough to filter regionalized ultrasound artefacts which may be attributed to "false transport" ($f_c = f_0 / 16$ provided best results on the presented in-vivo data), this value must also be high enough to prevent removing accurate microbubble information. Finally, the regularization parameter (α) in the optimized functional must be high enough to avoid ill conditioning of the numerical scheme (≥ 0.1 with the proposed implementation for the proposed in-vivo protocol), it must also be low enough in order to allow estimating local transports. Figures 6a and 6b and figures 7a and 7b demonstrate that the four calibration parameters δt , Δt , α and f_c may be fixed permanently for a specific DCEUS imaging protocol. Consequently, a great advantage of the invention is that an intervention of the user is only required for the manual determination of the ROI encompassing the imaged tissue.

Using the proposed approach, it is shown that a moderate imaging session duration is mandatory (< 20 seconds after the bolus arrival in the placenta, as shown in Figs. 6a and 6b). This is a great advantage since it diminishes the potential occurrence of long-term bias, including motion of ultrasound probe, foetus motion, etc...

It must be also underlined that the proposed implementation opens good perspectives toward a real-time diagnostic: the employed variational cost function renders itself compatible with a fast linear numerical scheme, while providing a dense pixel wise transport field. Calculations are consequently low time consuming: on the test platform used for this experimental evaluation, less than 100 ms were mandatory for the complete processing of one single frame. Taking into account that the employed imaging frame-rate was equal to 10 Hz for the in-vivo experiments, the processing of one image could be achieved within the interval of time available between two successive acquisitions. An embedded implementation on an ultrasound

scanner system for an immediate diagnostic is thus conceivable using available hardware.

Examples of applications

The invention comprises numerous applications, such as for example the
5 clinical diagnostic of obstetrical disorders, the quantitative analysis of the vascularisation of an organ such as the placenta or a tumour, the assessment of vascularisation in abnormal placental perfusion or the assessment of therapeutic efficacy.

10 Non-limitative examples of applications are described in the following documents:

- Arthuis et al. 2013.
- C. Bailey, T.A.G.M. Huisman, R.M. de Jong, and M. Hwang, "*Contrast-Enhanced Ultrasound and Elastography Imaging of the Neonatal Brain: A Review*" American Society of Neuroimaging, 2017.
- 15 - K.S. Mehta, J.J. Lee, A.A. Taha, E. Avgerinos, and R.A. Chaer, "*Vascular applications of contrast-enhanced ultrasound imaging*", Society for Vascular Surgery, vol. 66, no. 1, pp. 266-74, 2017.
- A.F.L. Schinkel, M. Kaspar, and D. Staub, "*Contrast-enhanced ultrasound: clinical applications in patients with atherosclerosis*", Int J Cardiovasc Imaging, vol. 32, pp. 35-48, 2016.
- 20 - Y. Hu, J. Zhu, Y. Jiang, and B. Hu, "*Ultrasound Microbubble Contrast Agents: Application to Therapy for Peripheral Vascular Disease*", vol. 26, no. 4, pp. 425-34, 2009.
- M. Kaspar, S. Partovi, M. Aschwanden, S. Imfeld, T. Baldi, H. Uthoff, and
25 D. Staub, "*Assessment of microcirculation by contrast-enhanced ultrasound: a new approach in vascular medicine*", Swiss Medical Weekly, 145:w14047, 2015.
- Y. Fei, and W. Li, "*Effectiveness of contrast-enhanced ultrasound for the diagnosis of acute pancreatitis: A systematic review and meta-analysis*", Digestive and Liver Disease, vol. 49, pp. 623-29, 2017.
- 30 - V. Rafailidis, A. Charitanti, T. Tegos, E. Destanis, and I. Chrysogonidis, "*Contrast-enhanced ultrasound of the carotid system: a review of the current literature*", J Ultrasound, vol. 20, pp. 97-109, 2017.
- B. Braden, A. Ignee, M. Hocke, R.M. Palmer, and C. Dietrich, "*Diagnostic value and clinical utility of contrast enhanced ultrasound in intestinal
35 diseases*", Digestive and Liver Disease, vol. 42, pp. 667-74, 2010.

Of course, the invention is not limited to the precise embodiments described hereinabove and various adaptations may be effected without departing from the

scope of the invention as defined in the appended claims. For example, it is to be understood that the present invention contemplates that, to the extent possible, one or more features of any embodiment can be combined with one or more features of any other embodiments.

CLAIMS

1. Apparatus for contrast-enhanced ultrasound imaging, comprising:
- injection means for injecting a contrast agent within a region of interest,
 - an ultrasound system for propagating ultrasound waves within said region of interest and detecting ultrasound waves backscattered within said region of interest,
 - imaging means for producing a temporal succession of images according to said backscattered ultrasound waves, each image being representative of a spatial configuration of said contrast agent within said region of interest,

characterised in that it comprises transport calculating means arranged to determine a spatial transport of the contrast agent from successive images.

15

2. Apparatus according to claim 1, in which the transport calculating means comprises numerical resolution means for numerical resolution of the following transport equation:

$$I_t + \vec{V} \cdot \vec{\nabla} I = 0$$

20 where I is a grey level intensity on said images, I_t is a partial temporal derivative of I between successive images, $\vec{\nabla}$ is a spatial gradient operator, $\vec{\nabla} I$ is the spatial gradient of I , and \vec{V} corresponds to said spatial transport between those successive images.

3. Apparatus according to claim 1 or 2, comprising resetting means arranged to compensate relative movements between said ultrasound system and said region of interest.

4. Apparatus according to claim 3, in which said resetting means is arranged to compensate relative movements between said ultrasound system and said region of interest using a gradient driven descent algorithm maximizing an inter-correlation coefficient between successive images.

5. Apparatus according to anyone of the preceding claims, comprising a spatial low-pass filter arranged to filter said images.

35

6. Method for contrast-enhanced ultrasound imaging, comprising:
- propagating ultrasound waves within a region of interest and detecting ultrasound waves backscattered within said region of interest,

- producing a temporal succession of images according to said backscattered ultrasound waves, each image being representative of a spatial configuration of said contrast agent within said region of interest, characterised in that it comprises determining a spatial transport of the contrast agent from successive images.

7. Method according to claim 6, in which determining said spatial transport comprises numerical solving of the following transport equation:

$$I_t + \vec{V} \cdot \vec{\nabla} I = 0$$

10 where I is a grey level intensity on said images, I_t is a partial temporal derivative of I between successive images, $\vec{\nabla}$ is a spatial gradient operator, $\vec{\nabla} I$ is the spatial gradient of I , and \vec{V} corresponds to said spatial transport between those images.

8. Method according to claim 6 or 7, comprising compensating relative movements between said ultrasound system and said region of interest using a resetting means.

9. Method according to claim 8, in which compensating said relative movements comprises maximizing an inter-correlation coefficient between successive images using a gradient driven descent algorithm.

10. Method according to anyone of claims 6 to 9, comprising filtering said images using a spatial low-pass filter.

25 11. Image processing tool, characterised in that it implements a method according to anyone of claims 6 to 10.

12. Device for contrast-enhanced ultrasound imaging, characterised in that it comprises an image processing tool according to claim 11.

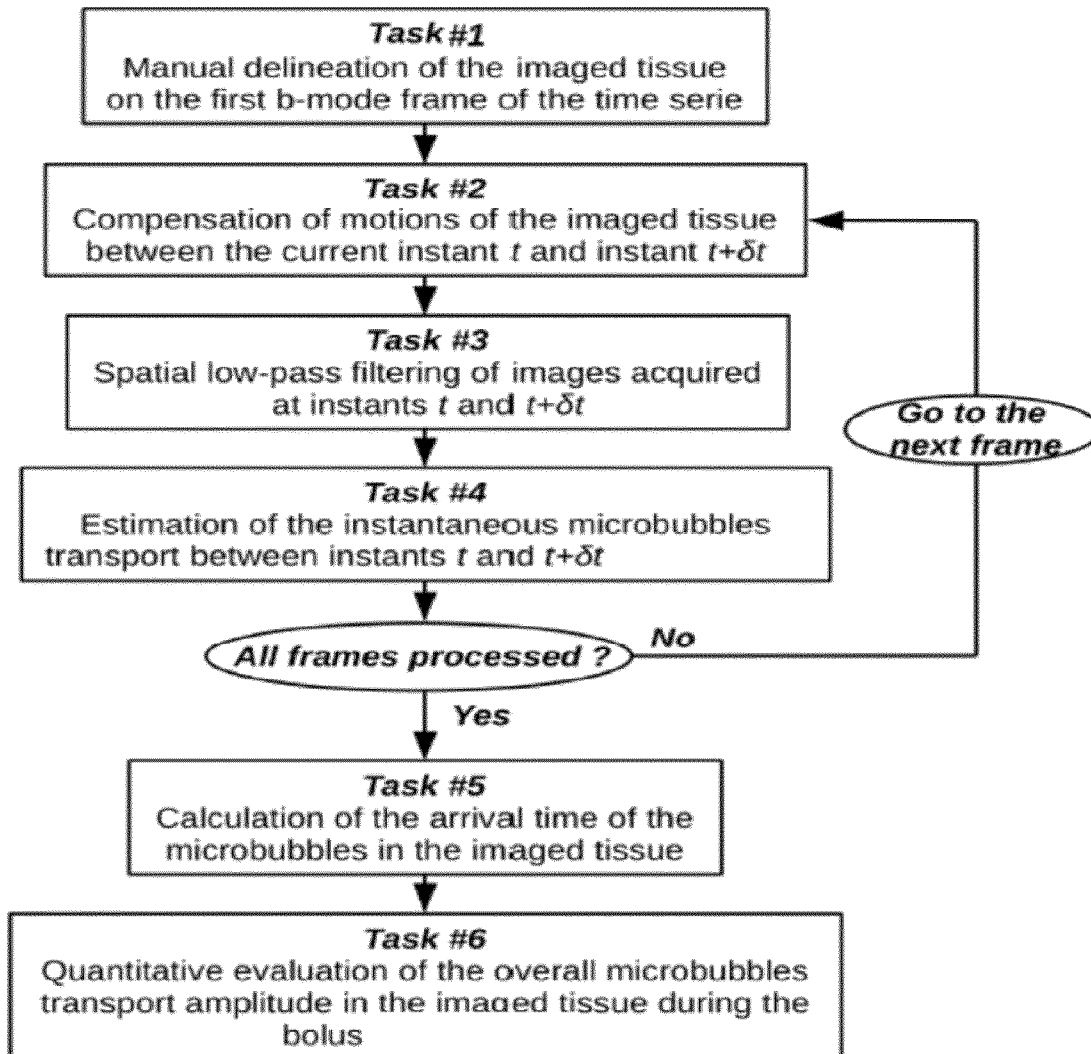
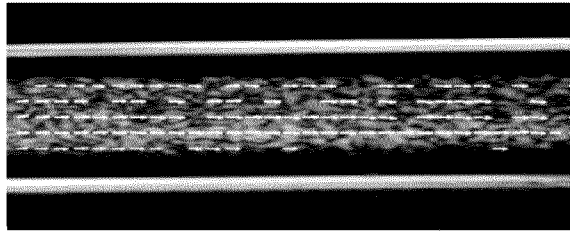
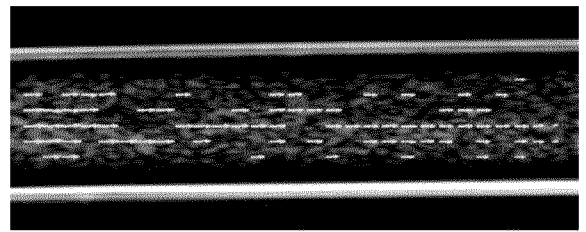


Fig. 1

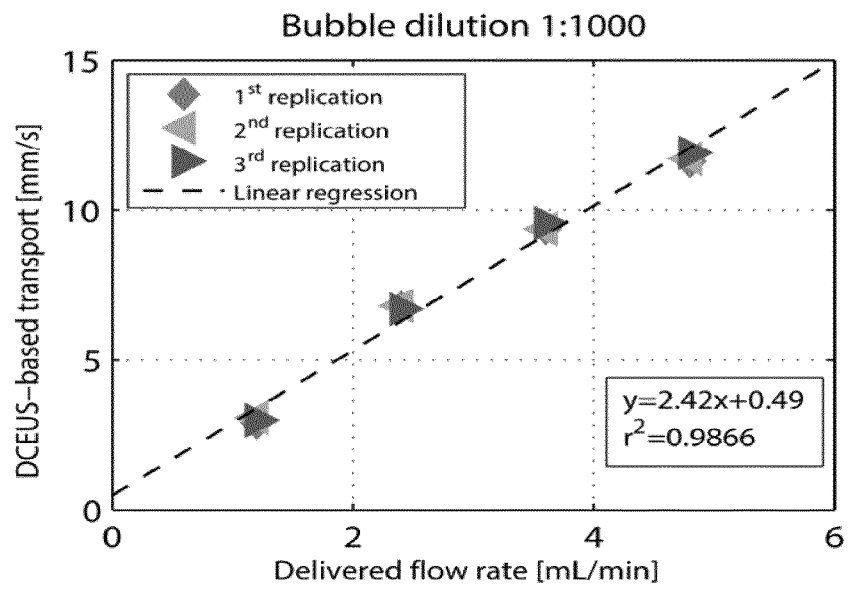
Fig. 2



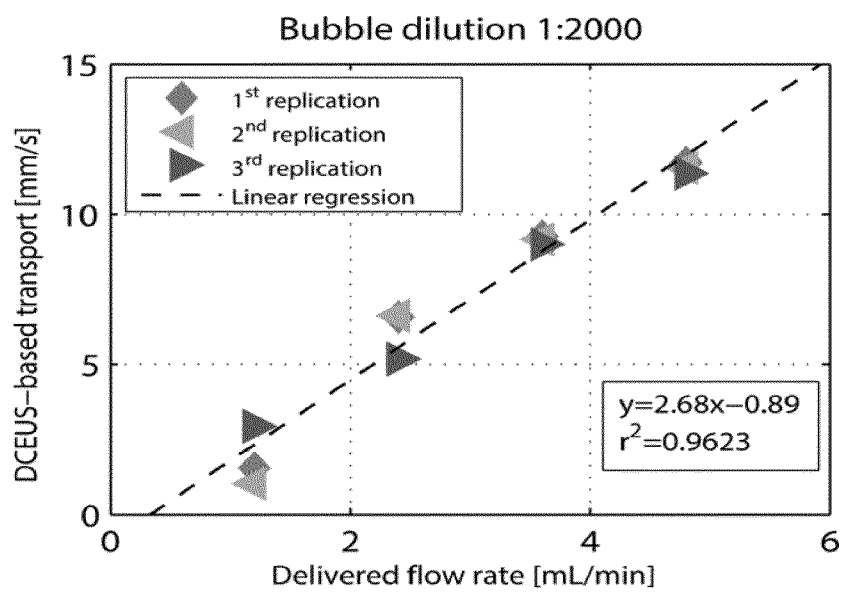
(a)



(b)



(c)



(d)

Fig. 3

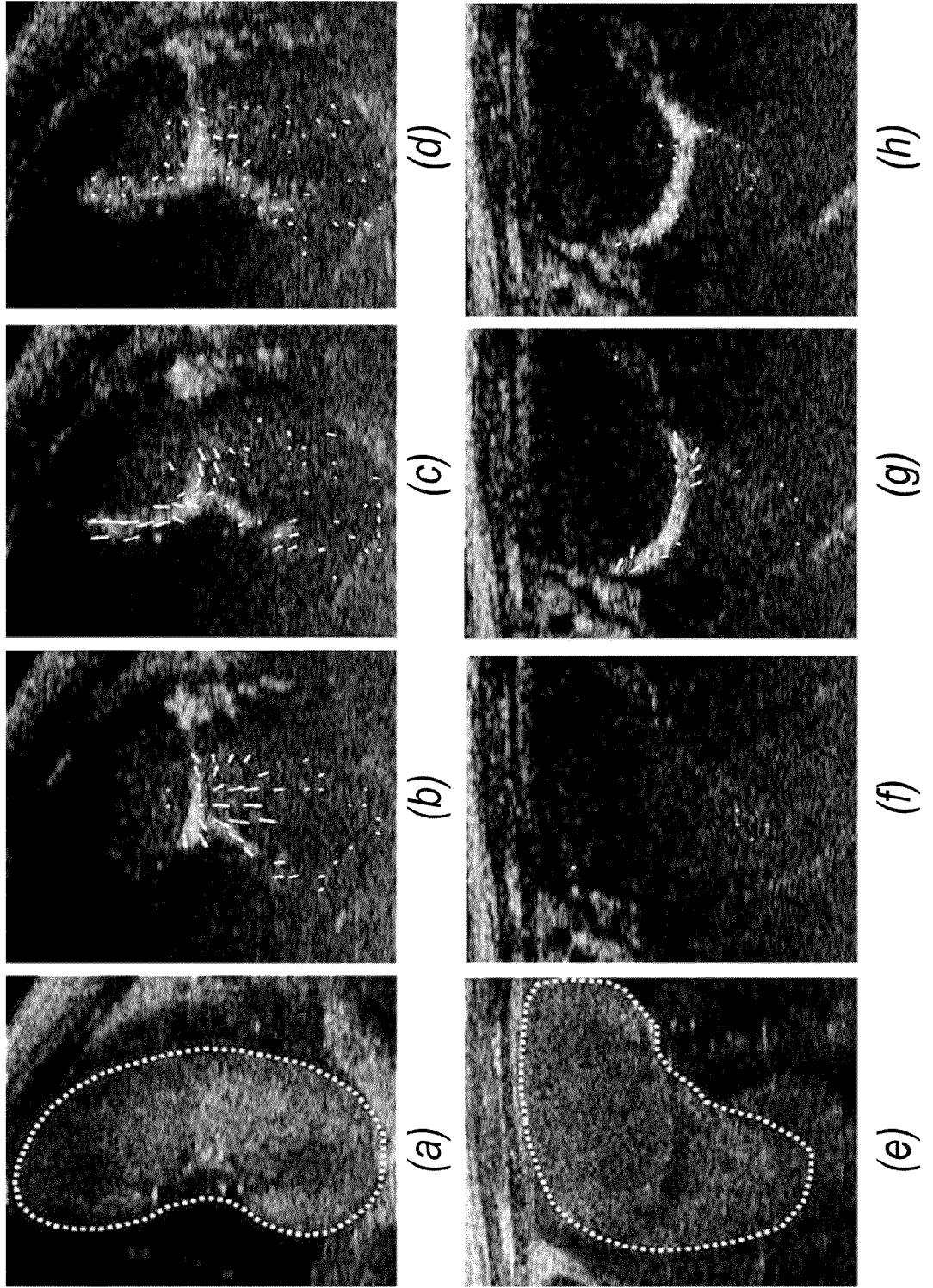
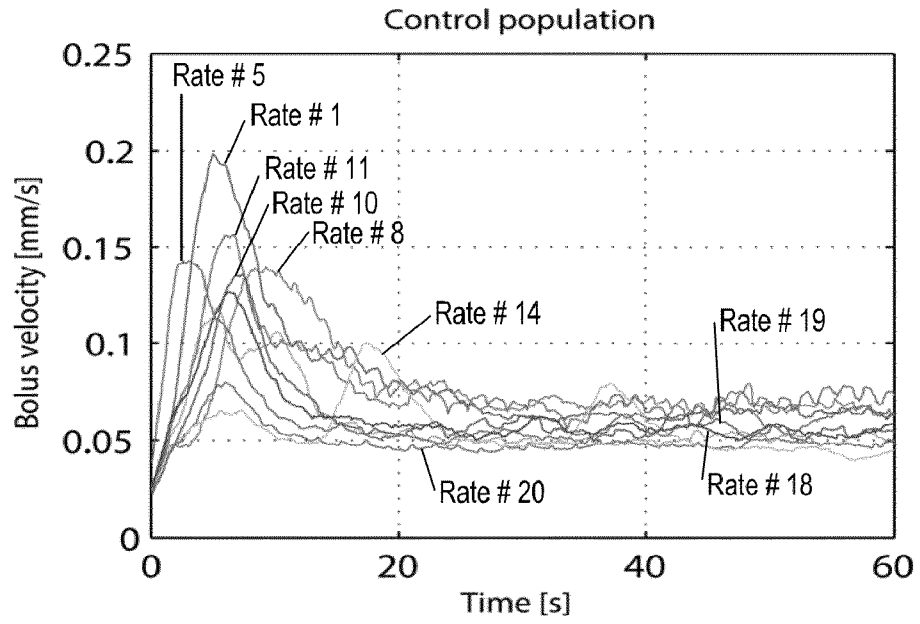
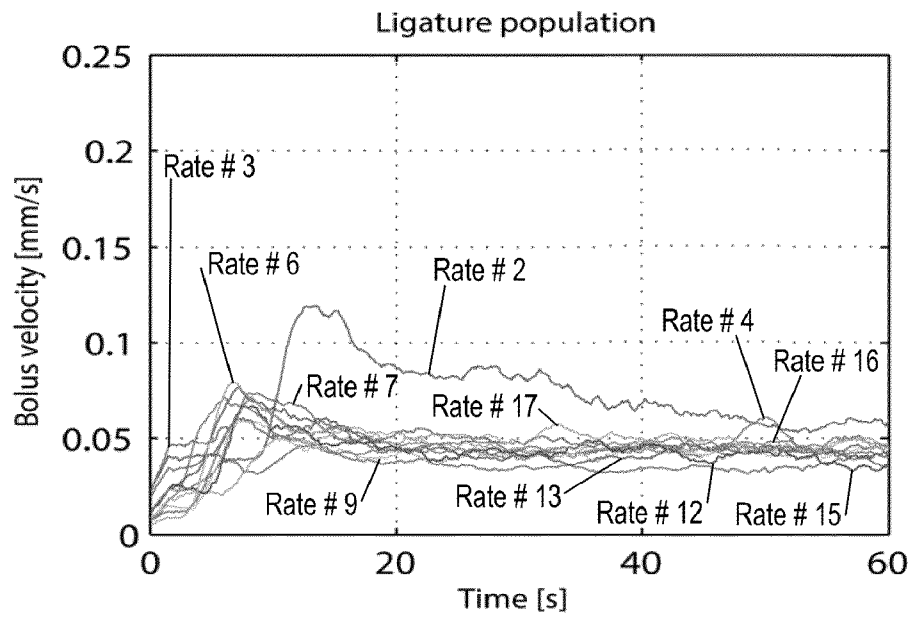


Fig. 4



(a)



(b)

Fig. 5

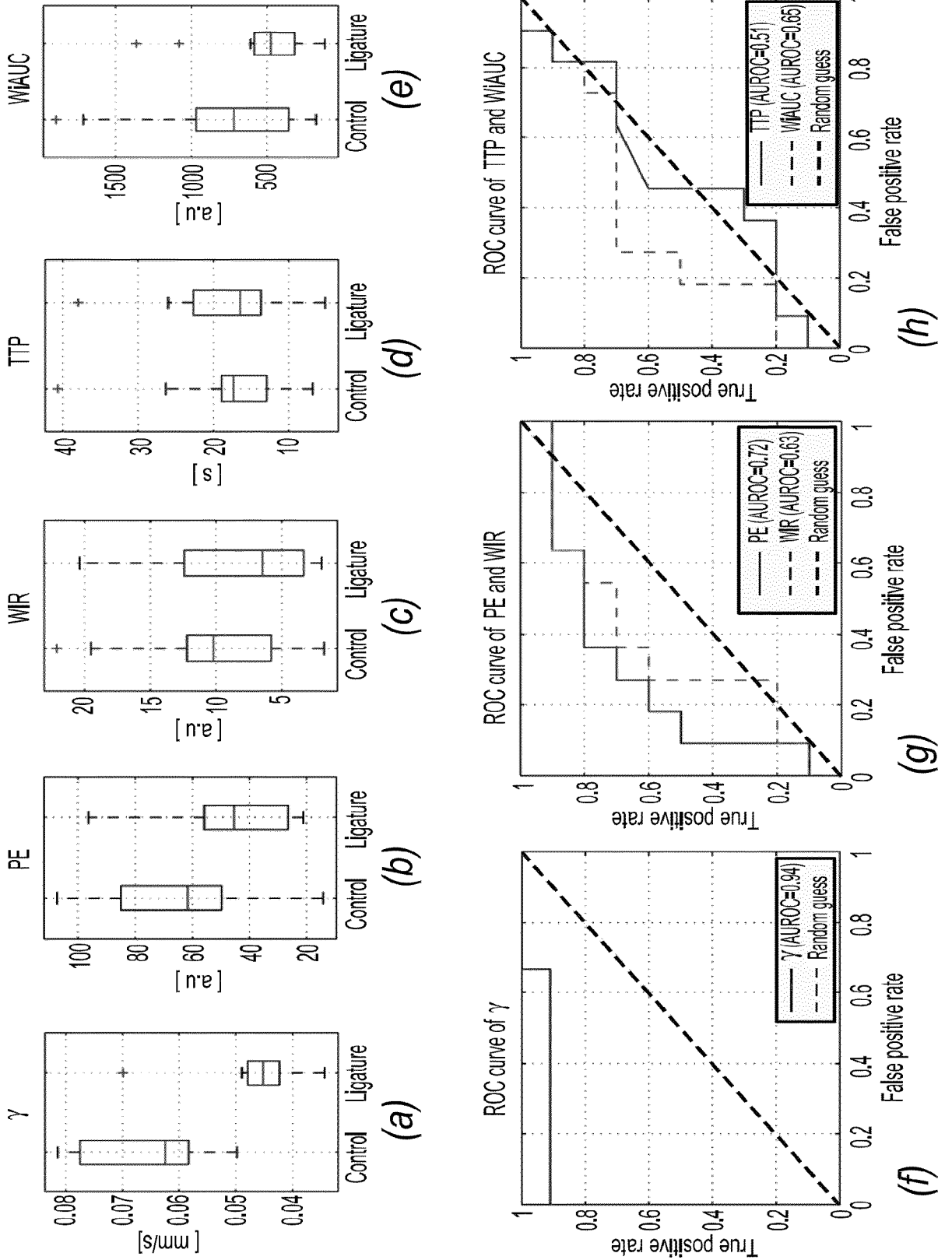


Fig. 6

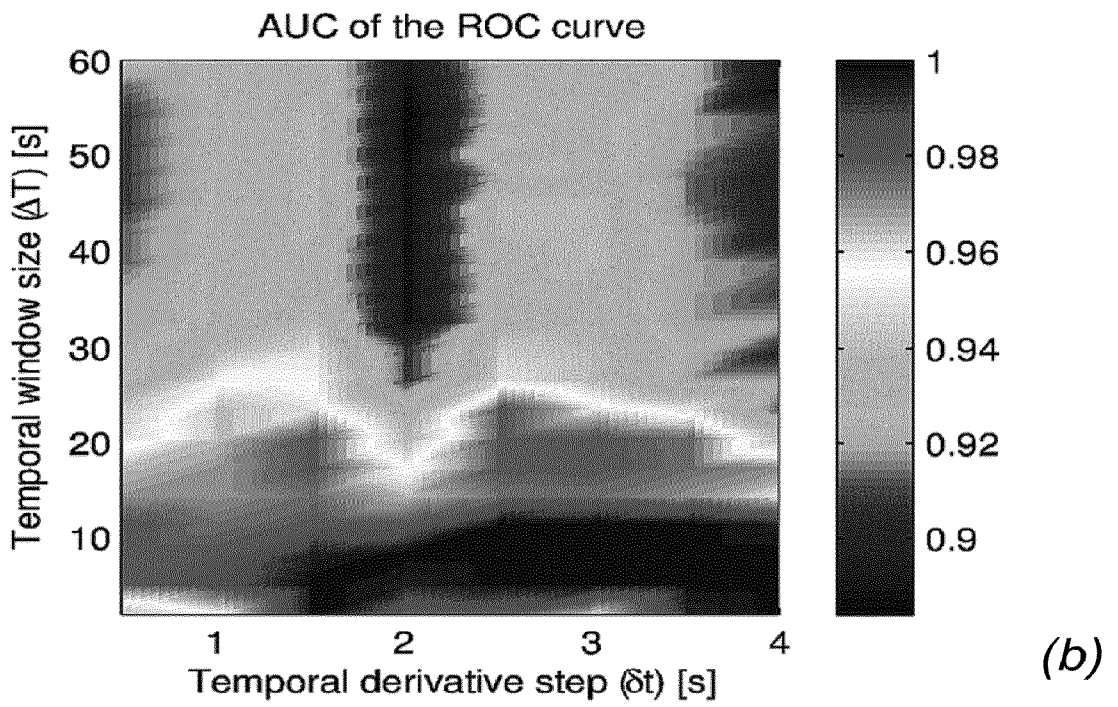
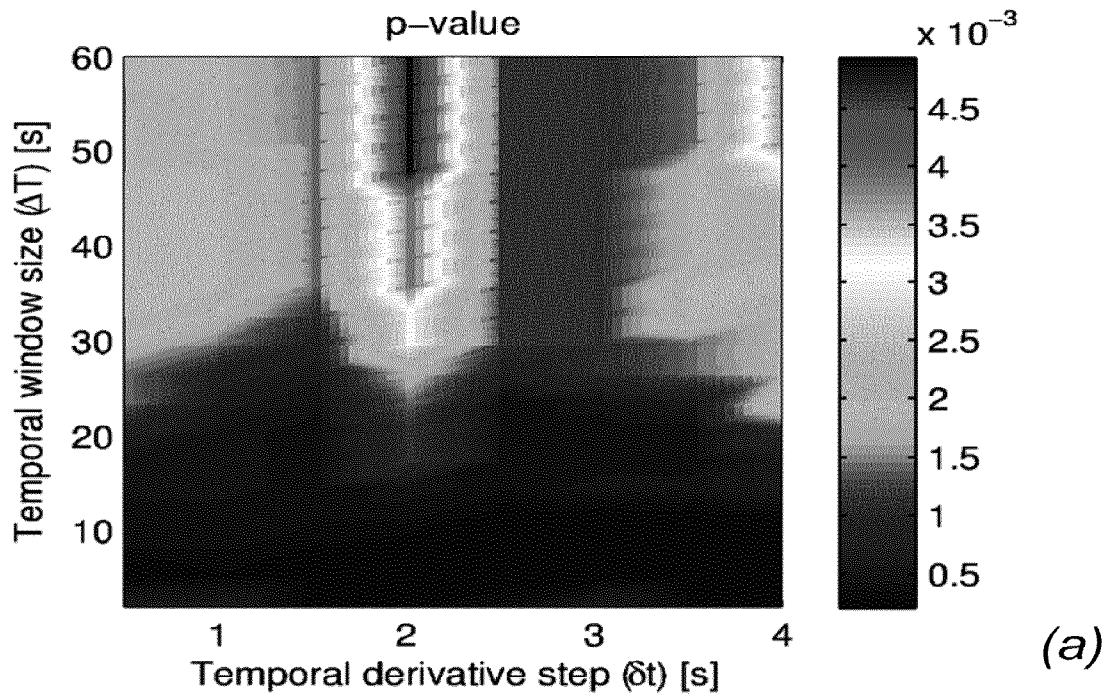
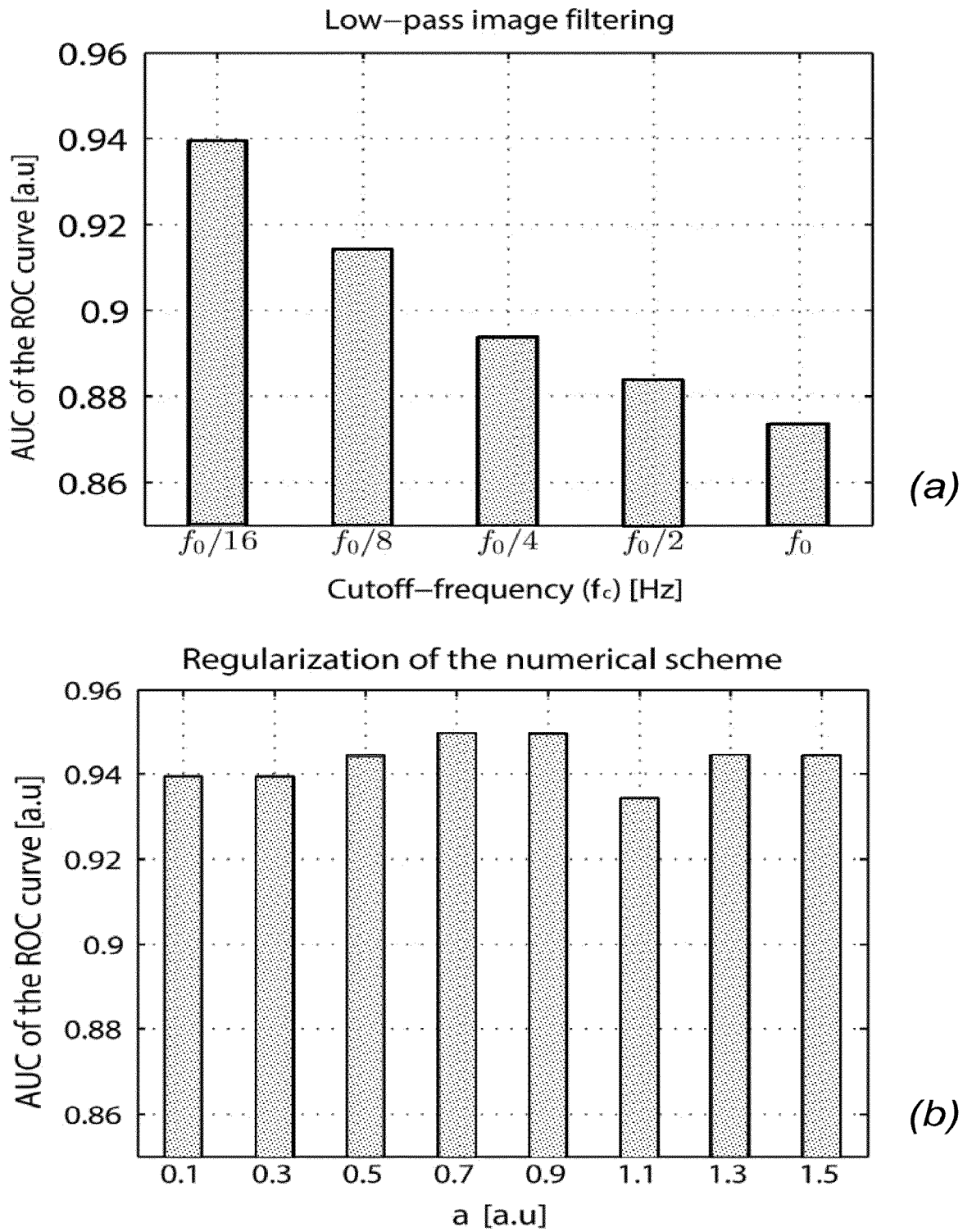


Fig. 7



INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2018/070383

A. CLASSIFICATION OF SUBJECT MATTER
 INV. A61B8/08
 ADD. A61B6/00 G01R33/56 G01S7/52 G06T5/50 G06T7/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
 Minimum documentation searched (classification system followed by classification symbols)
 A61B G01R G01S G06T

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
 EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X A	US 2008/228080 A1 (ARDITI MARCEL [CH] ET AL) 18 September 2008 (2008-09-18) paragraph [0012] paragraph [0025] paragraph [0053] paragraph [0052] paragraph [0055] paragraph [0063] paragraph [0104] paragraph [0099] figures 1,2b abstract ----- -/--	1,2,6,7, 11,12 4,9

Further documents are listed in the continuation of Box C.

See patent family annex.

- * Special categories of cited documents :
- "A" document defining the general state of the art which is not considered to be of particular relevance
 - "E" earlier application or patent but published on or after the international filing date
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 - "O" document referring to an oral disclosure, use, exhibition or other means
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 - "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
 - "&" document member of the same patent family

Date of the actual completion of the international search 5 September 2018	Date of mailing of the international search report 17/09/2018
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Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer De la Hera, Germán
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INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2018/070383

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 2017/196539 A1 (GU XIAOLIN [CN] ET AL) 13 July 2017 (2017-07-13) paragraph [0041] - paragraph [0042] paragraph [0040] paragraph [0043] paragraph [0070] - paragraph [0071] paragraph [0048] paragraph [0051] - paragraph [0052] figures 1,4,5 paragraph [0054] - paragraph [0058] paragraph [0064] - paragraph [0067] -----	1,3-6, 8-12
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Information on patent family members

International application No PCT/EP2018/070383

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专利名称(译)	具有对比度增强的超声成像的装置和方法		
公开(公告)号	EP3658038A1	公开(公告)日	2020-06-03
申请号	EP2018750370	申请日	2018-07-27
[标]申请(专利权)人(译)	法国国家健康医学研究院 UNIV之旅 法国国家科学研究中心 波尔多大学 INSTITUT POLYTECHN波尔多平局		
申请(专利权)人(译)	CENTRE法国国家科学研究 Université电波尔多 工业聚合物 (IPB)		
当前申请(专利权)人(译)	CENTRE法国国家科学研究 Université电波尔多 工业聚合物 (IPB)		
[标]发明人	DENIS DE SENNEVILLE BAUDOUIN PERROTIN FRANCK ESCOFFRE JEAN MICHEL BOUAKAZ AYACHE		
发明人	DENIS DE SENNEVILLE, BAUDOUIN PERROTIN, FRANCK ESCOFFRE, JEAN-MICHEL BOUAKAZ, AYACHE		
IPC分类号	A61B8/08 A61B6/00 G01R33/56 G01S7/52 G06T5/50 G06T7/00		
CPC分类号	A61B8/481 A61B8/5276 G01S7/52039 G06T7/0016 G06T2207/10016 G06T2207/10132 G06T2207/30104		
优先权	2017183673 2017-07-28 EP		
外部链接	Espacenet		

摘要(译)

本发明涉及用于对比度增强超声 (CEUS) 的设备和方法。 已知的 CEUS 技术仍不能令人满意地量化微循环并提供准确的临床诊断。 本发明介绍了使用流体动力学模型来分析动态对比增强超声 (DCEUS)。 本发明的效率在体外 (用于定量估计微气泡流速) 和在体内 (用于体内对怀孕大鼠的胎盘功能不全分类 (对照 vs. 结扎)) 进行评估。 该实验评估表明, 本发明优于已知的 CEUS 技术的性能。 应用的示例可能包括但不限于器官 (如胎盘或肿瘤) 血管化的定量分析, 产科疾病的临床诊断或异常胎盘灌注中血管化的评估。