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(54) Title: METHODS AND APPARATUS FOR ULTRASOUND IMAGING

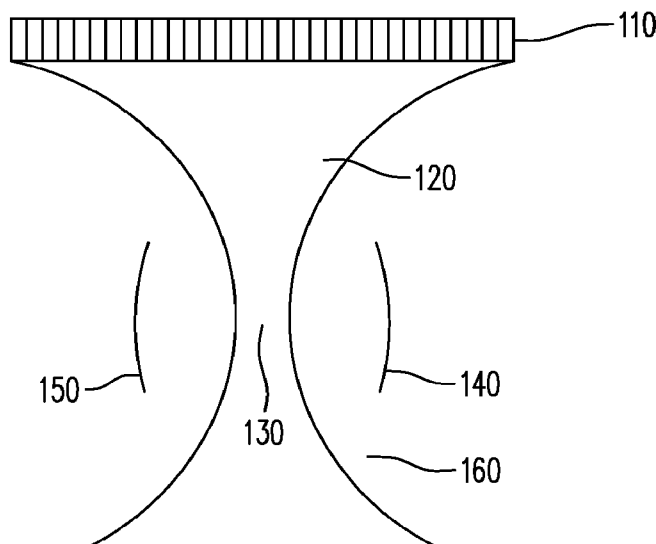


Figure 1

(57) Abstract: A first ultrasound pulse is applied to biological tissue to create shear waves in the biological tissue, a focused ultrasound pulse is transmitted into the biological tissue, one or more ultrasound signals is received from the biological tissue, and shear waves are detected in the biological tissue based on the received one or more ultrasound signals. At least one propagation property associated with the detected shear waves is determined, and the determined at least one propagation property is displayed.



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Methods and Apparatus for Ultrasound Imaging

Cross-reference to related applications

The present application claims priority to U.S. Provisional Patent Application
5 Serial No. 61/321,005, filed on April 5, 2010 and entitled “Method and Apparatus for
Ultrasound Imaging”, U.S. Provisional Patent Application Serial No. 61/321,341, filed on
April 6, 2010 and entitled “Method and Apparatus for Ultrasound Imaging” and U.S.
Provisional Patent Application Serial No. 61/350,585, filed on June 2, 2010 and entitled
“Method and Apparatus for Ultrasound Imaging”, the contents of which are incorporated
10 herein by reference for all purposes.

Background

Systems and methods described herein generally relate to the field of ultrasound
imaging. More specifically, embodiments described below relate to methods and systems
15 for measuring shear wave velocity in tissue.

Pathological conditions may result in soft tissue which is stiffer than would be
present under physiological conditions. Physicians therefore use palpation to locate stiff
tissue within a body and thereby identify pathological conditions. For example, breast
cancers are known to be generally harder than healthy breast tissue and may be detected
20 as a hard lump through palpation.

The propagation velocity of shear waves in tissue is related to the stiffness
(Young’s modulus or shear modulus) of tissue by the following equation,

$$E = 3\rho \cdot c^2 \quad (1)$$

where

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c is the propagation velocity of shear wave, E is Young's modulus, and ρ is the tissue density. Therefore, cancers or other pathological conditions may be detected in tissue by measuring the propagation velocity of shear waves passing through the tissue.

A shear wave may be created within tissue by applying a strong ultrasound pulse to the tissue. The ultrasound pulse may exhibit a high amplitude and a long duration (e.g., on the order of 100 microseconds). The ultrasound pulse generates an acoustic radiation force which pushes the tissue, thereby causing layers of tissue to slide along the direction of the ultrasound pulse. These sliding (shear) movements of tissue may be considered shear waves, which are of low frequencies (e.g., from 10 to 500 Hz) and may propagate in a direction perpendicular to the direction of the ultrasound pulse. The ultrasound pulse may propagate at a speed of 1540 m/s in tissue. However, the shear wave propagates much more slowly in tissue, approximately on the order of 1-10 m/s.

Since the tissue motion is generally in the axial direction (i.e., the ultrasound pulse direction) the shear waves may be detected using conventional ultrasound Doppler techniques. In this regard, the ultrasound Doppler technique is best suited to detect velocity in the axial direction. Alternately, shear waves may be detected by measuring a tissue displacement caused by the acoustic radiation force.

In order to accurately measure the propagation velocity of the shear wave, the shear wave needs to be tracked at a fast rate or a fast frame rate of several thousands frames per second. An image in a frame may consist of a few hundred ultrasound lines. A typical frame rate of regular ultrasound imaging is about 50 frames/s, which is too slow to track the shear wave propagation. Therefore, there exists a need to increase the frame rate while maintaining a good signal to noise ratio and good spatial resolution. Also, there exists a need to efficiently provide an indication of tissue stiffness.

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Summary

A method, medium and system may provide application of a first ultrasound pulse to biological tissue to create shear waves in the biological tissue, simultaneous transmission of a plurality of focused ultrasound pulses into the biological tissue, reception of a plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses, detection of the shear waves in the biological tissue based on the received one or more ultrasound signals, determination of a shear wave propagation velocity based on the detected shear waves, and display of the shear wave propagation velocity.

10

Brief Description Of The Drawings

Figure 1. A diagram of shear wave generation resulting from an acoustic radiation force.

Figure 2. A diagram of an ultrasound imaging system of some embodiments.

15 Figure 3. A diagram of a conventional ultrasound imaging system.

Figure 4. A diagram of multiple ultrasound transmitted/received beams.

Figure 5. A diagram of an ultrasound transmitted beam and multiple ultrasound received beams.

Figure 6. Color coding of shear wave propagation velocity squared.

20 Figure 7. Color coding of shear wave propagation velocity squared.

Figure 8. A diagram illustrating generation of shear waves by acoustic radiation forces and the propagation of shear waves.

Figure 9. A diagram illustrating sliding movements of shear waves.

Figure 10. A diagram illustrating the propagation of shear waves.

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Figure 11. A diagram illustrating the propagation of shear waves.

Figure 12. An example of a color-coded image of shear wave propagation velocity squared in tissue.

Figure 13. A diagram to illustrate tissue displacement caused by an acoustic
5 radiation force.

Figure 14. Scale of shear wave velocity squared c^2 by color coding bar composed of RGB representation.

Figure 15. A diagram to show the ultrasound coordinate system with respect to an
ultrasound transducer.

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Detailed Description

Embodiments will be described with reference to the accompanying drawing figures wherein like numbers represent like elements throughout. Before embodiments of the invention are explained in detail, it is to be understood that embodiments are not
15 limited in their application to the details of the examples set forth in the following description or illustrated in the figures. Other embodiments may be practiced or carried out in a variety of applications and in various ways. Also, it is to be understood that the phraseology and terminology used herein is for the purpose of description and should not be regarded as limiting. The use of “including,” “comprising,” or “having,” and
20 variations thereof herein is meant to encompass the items listed thereafter and equivalents thereof as well as additional items. The terms “mounted,” “connected,” and “coupled,” are used broadly and encompass both direct and indirect mounting, connecting, and coupling. Further, “connected,” and “coupled” are not restricted to physical or mechanical connections or couplings.

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Acoustic radiation force is created by a strong ultrasound pulse 120 as shown in Figure 1. The ultrasound pulse 120 exhibits a high amplitude as well as a long duration, (e.g., on the order of 100 microseconds). The ultrasound pulse 120 is transmitted from an ultrasound transducer array 110. The ultrasound pulse 120 is focused at a focal point 130 in biological tissue 160, resulting in an acoustic radiation force which pushes the tissue 160 at the focal point 130. The ultrasound pulse 120 may be transmitted multiple times and may be focused at a different focal point for each of multiple transmitted ultrasound pulses.

The tissue 160 is pushed mostly in the axial direction of the ultrasound pulse 120, creating shear waves 140, 150 which may propagate in the lateral direction or directions other than the axial direction (i.e., vertical direction). The propagation velocity of the shear waves 140, 150 depends on the stiffness (Young's modulus or the shear modulus) of the tissue 160. Greater tissue stiffness results in greater shear wave propagation velocity as shown in equation 1. Pathological conditions such as cancer may increase tissue stiffness thus these conditions may be diagnosed by determining the propagation velocity. For example, the shear wave propagation velocity may vary from 1 m/s to 10 m/s, depending on tissue conditions.

Since the shear wave may be characterized by tissue movement (or motion), the shear wave may be detected by the ultrasound Doppler technique (e.g., see US4573477, US4622977, US4641668, US4651742, US4651745, US4759375, US4766905, US4768515, US4771789, US4780837, US4799490, and US4961427). To detect this tissue movement (motion), the ultrasound pulse is transmitted multiple times to the tissue, and the ultrasound is scattered by scatterers in tissue and received by an ultrasound transducer as received ultrasound signals. The received ultrasound signals from the

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ultrasound array transducers are filtered, amplified, digitized, apotized, and beamformed (i.e. summed) after applying delays and/or phase-rotations for focusing and steering. The order of these processing steps may be interchanged. Received beamformed RF ultrasound signals undergo quadrature demodulation, resulting in complex, Doppler I-Q
5 signals. In a color Doppler technique, the ultrasound is transmitted at a pulse repetition frequency (PRF) and the velocity is detected as the shift in frequency (Doppler shift frequency) in the received ultrasound signal. The received ultrasound is mixed with in-phase (0 degrees) and quadrature (90 degrees) reference signals of the same frequency as the transmitted ultrasound frequency, resulting in complex I-Q Doppler signals.

10 Generally, the complex I-Q signal is used to derive the Doppler shift frequency because the Doppler shift frequency and the blood velocity have the following relationship

$$\Delta f = \frac{2f_t v \cos \theta}{c_s} \quad (2),$$

15

where Δf is the Doppler shift frequency, f_t is the transmitted frequency, v is the blood velocity, θ is the angle between the ultrasound beam direction and the velocity vector, and c_s is the speed of sound. The Doppler shift frequency is thus dependent on the angle between the velocity direction and the ultrasound beam direction
20 and is a measurement that an ultrasound color Doppler system may obtain.

In the case of color Doppler, the number of the sampled signals may be limited to several. Therefore, an auto-correlation technique is usually used to determine the phase

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differences between the I-Q signals and then to determine the Doppler shift frequency and the velocity as follows. The color Doppler's I-Q signals $z(m) = x(m) + jy(m)$ are used to calculate "auto-correlation" r as shown in the following equation, where $z(m)$ is the complex I-Q Doppler signal, $x(m)$ is the in-phase (real) signal, $y(m)$ is the quadrature phase (imaginary) signal, m indicates the signal number, j is the imaginary unit and $*$ indicates the complex conjugate.

$$r = \sum z(m) \cdot z^*(m-1) \quad (3)$$

The real ($Real(r)$) and imaginary ($Imag(r)$) parts of r are used to obtain the phase φ as shown in the following equation.

$$\varphi = \tan^{-1} \frac{Imag(r)}{Real(r)} \quad (4)$$

Since \tan^{-1} usually provides only -0.5π to 0.5π , the position of complex value r in the complex coordinate may be also used to derive φ in the range of $-\pi$ to π . The phase (i.e., color Doppler phase) φ is then related to the Doppler shift frequency as shown in the following equation.

$$\Delta f = \frac{\varphi_{PRF}}{2\pi} \quad (5)$$

Autocorrelation r between the received complex baseband ultrasound signals is thus obtained to detect tissue velocity or movement.

Tissue movement is detected at multiple lateral points in a field of tissue region by multiple ultrasound beams (for example, 540, 545, 550 in Figure 5) in order to monitor movement. This movement reflects action of the shear wave at those multiple lateral points (or multiple ultrasound beams). Consequently, the lateral propagation

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velocity of the shear wave may be determined from the detected tissue movement.

Alternately, the shear wave may be detected by measuring tissue displacement caused by acoustic radiation force which is in turn caused by a strong ultrasound pulse as shown in Figure 13. Tissue 1310 is positioned at a position 1320 before the acoustic radiation is applied and then is moved to a position 1330 after the acoustic radiation force was applied. To measure tissue displacement caused by the strong ultrasound pulse, ultrasound pulses are transmitted to tissue from an ultrasound transducer 1305 and then the ultrasound pulses are scattered from scatterers in tissue and returned to the transducer 1305 and received by the transducer 1305 as received ultrasound signals. The ultrasound pulses are focused at a depth in order to increase a signal-to-noise ratio of the resulting received ultrasound signals in comparison to unfocused ultrasound pulses. Using correlation of the received ultrasound signals from tissue the displacement 1340 (from the position 1320 to the position 1330) of the tissue 1310 due to the acoustic radiation force may be obtained and the tissue 1310 may be tracked thereafter. The ultrasound pulses may thereby track shear waves after shear waves are created by acoustic radiation force.

Ultrasound signals resulting from the first ultrasound pulse and received from the tissue 1310 before acoustic radiation force is applied are cross-correlated with received ultrasound signals resulting from the second ultrasound pulse after the acoustic radiation force is applied in order to find the best match between the received ultrasound signals. The best match may be found by finding a maximum correlation value to track the tissue and its displacement due to the acoustic radiation force. Therefore, when tissue displacement is observed or measured, a shear wave is detected. The displacement and tissue velocity may be related in that the displacement is a time integral $\int v_s dt$ of tissue

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velocity v_s . Therefore, the tissue displacement may be obtained by calculating the time integral of color Doppler velocity. Received ultrasound signals may be RF (Radio Frequency), IF (Intermediate Frequency) or baseband signals after demodulation. Alternately, the displacement may be further differentiated to obtain tissue strain, which
 5 may be then used to detect the shear wave propagation velocity.

Cross correlation $CC(t, \tau)$ of signals in the previous paragraphs may be mathematically expressed as follows,

$$CC(t, \tau) = \int_t^{t+W} S_1(t')S_2(t'-\tau)dt' \quad (6)$$

where $CC(t, \tau)$: cross correlation; $S_1(t')$: received signal from the first ultrasound
 10 transmission; $S_2(t'-\tau)$: received ultrasound signal from the second ultrasound transmission; W: window length; t: time, t': time; τ : time displacement. Time displacement value τ , which makes the maximum cross correlation (or the best match), determines the tissue displacement. Interpolation of signals using an interpolation function (e.g. cubic-spline) may be performed before cross correlation to increase spatial
 15 resolution.

The cross correlation may be replaced by the sum of absolute differences (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD), or the sum of absolute power differences (SPD) as follows.

$$SAD[l, k] = \sum_{n=0}^N |S_1[l+n] - S_2[l+n-k]| \quad (7)$$

$$20 \quad SSD[l, k] = \sum_{n=0}^N (S_1[l+n] - S_2[l+n-k])^2 \quad (8)$$

$$SCD[l, k] = \sum_{n=0}^N |S_1[l+n] - S_2[l+n-k]|^3 \quad (9)$$

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$$SPD[l, k] = \sum_{n=0}^N |S_1[l+n] - S_2[l+n-k]|^p \quad (10)$$

S_1 is the received ultrasound signal from the first ultrasound transmission before displacement, S_2 is the received ultrasound signal from the second ultrasound transmission after displacement. N : the number of signals in the signal window. k : window displacement by the number of signals and equivalent of τ . l : the position of the window. p is a real number. For SAD, SSD, SCD and SPD, the tissue displacement is determined based on the value of k that makes the minimum (or best match) of each of the SAD, SSD, SCD and SPD.

Figures 8 and 9 are used to illustrate shear wave generation and detection in detail.

10 A strong ultrasound pulse 820 is applied to tissue 860, 960 from an ultrasound transducer 810, 910 once or more times to increase the amplitude of shear waves which are caused by acoustic radiation forces resulting from the ultrasound pulse. Shear waves attenuate very quickly in tissue and thus a greater amplitude results in a greater propagation distance. One or multiple ultrasound pulses may be focused at one focal point or different

15 focal points. The ultrasound pulse creates acoustic radiation forces which push a layer of tissue, resulting in tissue movement 830, 910 mostly in the axial (vertical) direction as illustrated in Figure 9. The tissue layer movement 910 causes adjacent tissue layer movements 920, 925 mostly in the axial direction. The tissue layer movements 920, 925 then in turn cause next tissue layer movements 930, 935 which then cause adjacent tissue

20 layer movements 940, 945. This succession of tissue movements represents a propagation of shear waves 840, 850 in the lateral (horizontal) direction as shown in Figure 8. Since the tissue movements (or motions) caused by acoustic radiation forces are mostly in the axial direction, the motion may be detected by the color Doppler technique, which is

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sensitive to motions in the axial direction.

For example, the color Doppler technique transmits and receives several ultrasound pulses, determines phase differences between the received ultrasound signals, and calculates a velocity of tissue or blood using the autocorrelation technique as previously discussed and known in the art. Variance and power of color Doppler signals may be also calculated in addition to the velocity. As in the conventional display of moving tissue or blood, one of these parameters may be used to display shear waves as shown in Figures 10, 11. It will be assumed that shear waves 1040 (1140), 1050 (1150) are determined in a color Doppler frame representing a certain time and shear waves 1060 (1160), 1070 (1170) are determined at a next moment or in a next frame. More image frames of shear waves may be obtained to track the shear waves and to create a movie of shear wave propagation. In alternate embodiments, tissue displacement due to acoustic radiation forces may be detected.

Figures 10 and 11 depict shear wave propagation at two points in time. Local shear wave propagation velocities, as illustrated by arrows 1080, 1090, may be derived by correlating two images of shear waves at two points in time. More image frames of shear waves may be used to track the propagation of shear waves in more image areas in order to present local shear wave propagation velocities or shear wave propagation velocity squared in a two-dimensional image as described below.

Correlation coefficient (CCV) between a first frame signal S^1 and the second frame signal S^2 may be obtained as speckle tracking as follows,

$$CCV(S^1, S^2) = \frac{\sum_{x=1}^m \sum_{z=1}^n (S^1_{x,z} - \bar{S}^1)(S^2_{x+X, z+Z} - \bar{S}^2)}{\sqrt{\sum_{x=1}^m \sum_{z=1}^n (S^1_{x,z} - \bar{S}^1)^2 \cdot \sum_{x=1}^m \sum_{z=1}^n (S^2_{x+X, z+Z} - \bar{S}^2)^2}} \quad (11)$$

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where $S^1_{x,z}$ is the ultrasound signal at x, z of the first frame, $S^2_{x+X,z+Z}$ is the ultrasound signal at $x+X, z+Z$ of the second frame, $\overline{S^1}$ is mean signal value in the window of the first frame signal, $\overline{S^2}$ is mean signal value in the window of the second frame signal. The coordinate system (x,y,z) is shown with respect to an ultrasound transducer 1510 in Figure 15. The elevational axis y is perpendicular to the paper of Figure 15 although it is shown slightly different for illustration purposes.

The displacement X, Z , that yields the maximum correlation coefficient determines the correct speckle tracking and the distance, and thus the velocity (i.e., the distance per time).

Similar to the 1D case, the correlation coefficient may be replaced by the sum of absolute differences (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD) and the sum of absolute power differences (SPD) as follows.

$$SAD(S^1, S^2, X, Z) = \sum_{x=1}^m \sum_{z=1}^n |S^1_{x,z} - S^2_{x+X,z+Z}| \quad (12)$$

$$SSD(S^1, S^2, X, Z) = \sum_{x=1}^m \sum_{z=1}^n (S^1_{x,z} - S^2_{x+X,z+Z})^2 \quad (13)$$

$$SCD(S^1, S^2, X, Z) = \sum_{x=1}^m \sum_{z=1}^n |S^1_{x,z} - S^2_{x+X,z+Z}|^3 \quad (14)$$

$$SPD(S^1, S^2, X, Z) = \sum_{x=1}^m \sum_{z=1}^n |S^1_{x,z} - S^2_{x+X,z+Z}|^p \quad (15)$$

p is a real number; m and n are integers. The 2D speckle tracking may be approximated by a 1D speckle tracking to obtain the shear wave propagation velocity and the shear wave propagation velocity squared. The mathematical expression will be similar to that used in the displacement measurement.

Alternately, a shear wave equation (16) may be used to derive the shear wave

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propagation velocity as follows,

$$\rho \frac{\partial^2 u_i}{\partial t^2} = \mu \left(\frac{\partial^2 u_i}{\partial x^2} + \frac{\partial^2 u_i}{\partial y^2} + \frac{\partial^2 u_i}{\partial z^2} \right) \quad (16)$$

where $i = x, y, z$, ρ is tissue density, μ is the shear modulus, u_i is the displacement vector, x is lateral coordinate, y is elevational coordinate and z is axial coordinate as shown in Figure 15. For incompressible materials, the Young's modulus E and the shear modulus μ have the following relationship.

$$E = 3\mu \quad (17)$$

Therefore, the shear wave propagation velocity squared may be obtained as a ratio of the shear modulus to the density as the following equation.

$$c^2 = \frac{\mu}{\rho} \quad (18)$$

One of the displacement components u_z in equation 16 may be determined by cross-correlation as previously discussed. By combining z component of equation 16 and equation 18, the shear wave propagation velocity squared and velocity are obtained as follows,

$$c^2 = \frac{\frac{\partial^2 u_z}{\partial t^2}}{\frac{\partial^2 u_z}{\partial x^2} + \frac{\partial^2 u_z}{\partial y^2} + \frac{\partial^2 u_z}{\partial z^2}} \quad (19)$$

and

$$c = \sqrt{\frac{\frac{\partial^2 u_z}{\partial t^2}}{\frac{\partial^2 u_z}{\partial x^2} + \frac{\partial^2 u_z}{\partial y^2} + \frac{\partial^2 u_z}{\partial z^2}}} \quad (20).$$

Therefore, the shear wave propagation velocity is obtained as the square root of the ratio

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between the temporal second-order derivative of the displacement and the spatial second-order derivatives of the displacement. Likewise, the shear wave propagation velocity squared is obtained as the ratio between the temporal second-order derivative of the displacement and the spatial second-order derivatives of the displacement. Since the spatial derivative of the displacement in elevational direction $\frac{\partial^2 u_z}{\partial y^2}$ may be considered negligible compared with the other spatial derivatives, the shear wave propagation velocity squared and velocity may be obtained from the other measurement values.

It is desirable to monitor and to track the shear wave frequently, meaning at a fast rate or frame rate. To speed up the frame rate, a wide, focused ultrasound pulse 520 may be transmitted and multiple ultrasound signals 540, 545, 550 may be simultaneously received as shown in Figure 5. The received ultrasound beams are used as described previously to detect shear waves and to derive shear wave propagation properties (i.e., velocity and velocity squared) therefrom. The focused transmit ultrasound beam 520 may be particularly suitable for maintaining a good signal-to-noise ratio of resulting received ultrasound beams during the detection of shear waves.

In some embodiments, multiple ultrasound beams (pulses) are simultaneously applied and transmitted to the tissue field and multiple ultrasound beams (pulses) per transmitted ultrasound pulse are received to increase the frame rate, as shown in Figure 4. In Figure 4, ultrasound pulses 420, 430 are simultaneously transmitted to biological tissue 480 from an ultrasound transducer array 410. For each transmitted ultrasound pulse 420, 430, multiple ultrasound receive signals 440, 445, 465, 460, 465, 470 are simultaneously received. The multiple ultrasound pulses may be transmitted simultaneously or at substantially simultaneous times. The multiple ultrasound pulses may be simultaneously

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transmitted. Or a second ultrasound pulse may be transmitted after a first ultrasound pulse is transmitted and before the first ultrasound pulse returns to the ultrasound transducer from a deepest depth of an ultrasound field. This transmission method increases the frame rate.

5 Figure 4 shows an example of two simultaneous transmitted ultrasound pulses but more than two transmitted ultrasound pulses may be also used. In some embodiments, coded ultrasound waveforms may be transmitted for better separation of simultaneous multiple ultrasound signals. For example, chirp codes, Barker codes, Golay codes or Hadamard codes may be used for better separation of ultrasound pulses. Again, the
10 received signals are analyzed using the methods previously described to determine tissue movement at multiple points, and shear wave propagation properties are derived therefrom.

 An image of a shear wave can be created based on the motion (or velocity) detected at multiple points in the imaging field. Subsequent transmit/receive sequences of
15 ultrasound may create multiple images of the shear wave at multiple points in time. Correlation between the images of the shear wave is then calculated to obtain the shear wave propagation velocity and velocity squared as previously discussed. Alternately, tissue displacement caused by acoustic radiation force is determined and the shear wave propagation velocity is calculated as the square root of the ratio between the temporal
20 second-order derivative of the displacement and the spatial second-order derivatives of the displacement. Likewise, the shear wave propagation velocity squared is calculated as the ratio between the temporal second-order derivative of the displacement and the spatial second-order derivatives of the displacement.

 In some embodiments, the propagation velocity of a detected shear wave (c) may

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be displayed. In some embodiments, the propagation velocity squared (c^2) of the detected shear wave may be displayed. Advantageously, the propagation velocity squared (c^2) may be more closely related than the propagation velocity (c) to the Young's modulus or the shear modulus as shown in equation 1. Therefore the propagation velocity squared (c^2) may provide an efficient proxy for the actual stiffness. In some embodiments, the propagation velocity squared (c^2) may be multiplied by three and then displayed. If tissue density is close to 1 g/cm^3 , this number (i.e., $3c^2$) may be close to the actual Young's modulus. In some embodiments, a product (bc^2) of any real number (b) and the propagation velocity squared (c^2) may be displayed. Determinations of actual stiffness are difficult and error-prone because the density of the tissue is unknown and must be estimated.

A color coding technique, a grayscale technique, or a graphical coding technique may be employed to present a shear wave propagation property (i.e., velocity c or velocity squared c^2) to a user. In some embodiments, a propagation velocity squared (c^2) of shear waves within tissue is displayed in a two-dimensional color image. Graphical-coding and/or two-dimensional images may also be used to represent the propagation velocity c or velocity squared c^2 in some embodiments.

A low value of shear wave propagation velocity squared c^2 may be coded using a red color while a high value of c^2 may be coded using a blue color. For example, Figure 6 illustrates a legend indicating that a red-colored tissue area includes shear waves associated with low c^2 values (e.g., $1 \text{ m}^2/\text{s}^2$) and that a blue-colored tissue area includes shear waves associated with high c^2 values (e.g., $100 \text{ m}^2/\text{s}^2$). Embodiments are not limited to color-based coding. Images of shear wave propagation properties within tissue may be coded using grayscale or any combination of graphical patterns (e.g., vertical lines,

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horizontal lines, cross-hatching, dot patterns of different densities, etc.) and colors.

After determining the propagation velocity squared (c^2), c^2 may be coded linearly with respect to the color wavelength as shown in Figure 6. For example, if c^2 within a tissue area is determined to be $50 \text{ m}^2/\text{s}^2$, the tissue area may be displayed using a yellow color 630.

Alternately, color-coding of the shear wave propagation velocity squared (c^2) may be defined as shown in Figure 7. Tissue areas associated with low values of the shear wave propagation velocity squared may be displayed as blue 710 while areas associated with high values of the velocity squared may be displayed as red 720. Different color-coding methods may be also used to represent the propagation velocity squared (c^2) or velocity c of shear waves. For example, color coding may be based on hue, brightness, and other color characteristics. The color-coded scale may represent different maximums and minimums of the shear wave propagation velocity squared or velocity than shown in Figure 6, 7. In this regard, the velocity squared maximum of $100 \text{ m}^2/\text{s}^2$ and velocity squared minimum of $1 \text{ m}^2/\text{s}^2$ in Figures 6 and 7 are only for the illustration purposes and do not limit the scope of the claims. Other values may represent the maximum or minimum values of the coding scale.

Color coding based on Red, Green and Blue (RGB) values may be used to represent the propagation velocity c or velocity squared (c^2) of shear waves as shown in Figure 14. In this example (Figure 14), the propagation velocity squared (c^2) of a shear wave within tissue is represented according to a color coding bar 1410 which is based on RGB values 1420, 1430 and 1440. The shear wave propagation velocity squared has 256 possible values in this example, as represented 256 colors in the color coding bar 1410. The smallest velocity squared $c^2(0)$ 1412 is represented by a color composed of a

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combination of R(0) 1422, G(0) 1432 and B(0) 1442. The middle velocity squared $c^2(127)$ 1415 is represented by a color composed of a combination of R(127) 1425, G(127) 1435 and B(127) 1445. The highest velocity squared $c^2(255)$ 1418 is represented by a color composed of a combination of R(255) 1428, G(255) 1438 and B(255) 1448. In this example, R(255) only indicates a Red color associated with the red index 255 and does not necessarily indicate a Red color value of 255, which is the brightest Red color. Likewise, G(255) indicates a Green color associated with the green index 255 and B(255) indicates a Blue color associated with the blue index 255.

Alternately, Red, Green, Blue and Yellow may be used to define a color coding bar. Alternately, a Hue-based color coding bar may be used.

Figure 12 represents an example of a color-coded image 1260 displaying a shear wave propagation velocity squared c^2 within human soft tissue (e.g. breast). A color coding scale 1250 is illustrated, in which a color code 1210 (i.e., representing a red color although displayed as white in this black/white document) represents a low shear wave propagation velocity squared value and a color code 1220 (i.e., representing a blue color although displayed as hatched in this black/white document) represents a higher shear wave propagation velocity squared value.

Based on the coding scale 1250, it can be seen that the color coded image 1260 includes an area 1280 of high propagation velocity squared c^2 . Since the shear wave propagation velocity squared c^2 is proportional to the Young's modulus, the tissue area corresponding to area 1280 is likely to be hard. Since a tumor is generally hard, image 1260 may indicate pathological conditions.

The color-coding method provides efficient distinction between an area including shear waves having a high propagation velocity squared value and other areas including

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shear waves having a low propagation velocity squared value. The color coding method therefore allows efficient identification of hard tissue areas within soft tissue areas. An image displaying shear wave propagation velocity or velocity squared may be combined (e.g., superimposed) with a regular image of ultrasound, e.g. B-mode image, or a combined B-mode image and color Doppler image and/or spectral Doppler image. Alternately, the shear wave propagation velocity squared or velocity may be displayed numerically. In some embodiments, the shear wave propagation velocity squared may be displayed in gray scale or based on other graphic coding methods such as using patterns rather than colors. For example, low values of shear wave propagation velocity or square of the shear wave propagation velocity may be displayed in black or dark gray while high values of shear wave propagation velocity or shear wave propagation velocity squared may be displayed in light gray or white using a grayscale coding method.

Figure 3 shows a diagram of a conventional ultrasound diagnostic imaging system with B-mode imaging, Doppler spectrum and color Doppler imaging. The system may include other imaging modes, e.g. elasticity imaging, 3D imaging, real-time 3D imaging, tissue Doppler imaging, tissue harmonic imaging, contrast imaging and others. An ultrasound signal is transmitted from an ultrasound probe 330 driven by a transmitter/transmit beamformer 310 through a transmit/receive switch 320. The probe 320 may consist of an array of ultrasound transducer elements which are separately driven by the transmitter/transmit beamformer 310 with different time-delays so that a transmit ultrasound beam is focused and steered. A receive beamformer 340 receives the received ultrasound signals from the probe 330 through the switch 320 and processes the signals 325. The receive beamformer 340 applies delays and/or phases to the signals and the resultant signals are summed for focusing and steering a received ultrasound beam. The

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receive beamformer 340 may also apply apodization, amplification and filtering.

The processed signal 345 is coupled to a Doppler spectrum processor 350, a color Doppler processor 360, and a B-mode image processor 370. The Doppler spectrum processor 350 includes a Doppler signal processor and a spectrum analyzer, and processes Doppler flow velocity signals and calculates and outputs a Doppler spectrum 355. The color Doppler processor 360 processes the received signal 345 and calculates and outputs velocity, power and variance signals 365. The B-mode image processor 370 processes the received signal 345 and calculates and outputs a B-mode image 375 or the amplitude of the signal by an amplitude detection.

The Doppler spectrum signals 355, color Doppler processor signals (velocity, power, and variance) 365 and B-mode processor signals 375 are coupled to a scan converter 380 that converts the signals to scan-converted signals. The output of scan converter 380 is coupled to a display monitor 390 for displaying ultrasound images.

Figure 2 shows a diagram of elements of an ultrasound imaging system including a shear wave processor 295 according to some embodiments. The ultrasound system in Figure 2 transmits strong ultrasound pulses to biological tissue to create acoustic radiation forces which push the biological tissue. Shear waves are created and propagate in the tissue after the biological tissue is pushed. The ultrasound system then transmits and receives ultrasound pulses to track the shear waves as the shear waves propagate in the biological tissue. Multiple received ultrasound beams may be simultaneously formed by the receive beamformer 240. Likewise, multiple transmitted ultrasound beams may be simultaneously formed by the transmitter/transmit beamformer 210. Received ultrasound signals from the receive beamformer 240 are processed to obtain tissue displacement, Doppler velocity, correlation, shear wave propagation velocity and/or shear wave

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propagation velocity squared as previously described. The shear wave processor 295 may perform the shear wave processing methods described previously. The shear wave processor 295 receives output 245 from the receive beamformer 240. Output 297 comprises shear wave velocity data or other shear wave properties. For example, the
5 shear wave processor 295 outputs the propagation velocity or the square of the propagation velocity of the shear wave to a scan converter 280 and a representation of the shear wave propagation velocity or the square of the shear wave propagation velocity is output to the display monitor along with the B-mode, color Doppler or spectral Doppler images.

10 The shear wave processor 295 may comprise of general purpose central processing units (CPUs), digital signal processors (DSPs), field programmable Arrays (FPGAs), graphic processing units (GPUs) and/or discreet electronics devices.

Figure 2 represents a logical architecture according to some embodiments, and actual implementations may include more or different elements arranged in other manners.

15 Other topologies may be used in conjunction with other embodiments. Moreover, each element of the Figure 2 system may be implemented by any number of computing devices in communication with one another via any number of other public and/or private networks. Two or more of such computing devices may be located remote from one another and may communicate with one another via any known manner of network(s)
20 and/or a dedicated connection. The system may comprise any number of hardware and/or software elements suitable to provide the functions described herein as well as any other functions. For example, any computing device used in an implementation of the Figure 2

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system may include a processor to execute program code such that the computing device operates as described herein.

All systems and processes discussed herein may be embodied in program code stored on one or more non-transitory computer-readable media. Such media may include, 5 for example, a floppy disk, a CD-ROM, a DVD-ROM, a Blu-ray disk, a Flash drive, magnetic tape, and solid state Random Access Memory (RAM) or Read Only Memory (ROM) storage units. Embodiments are therefore not limited to any specific combination of hardware and software.

One or more embodiments have been described. Nevertheless, various 10 modifications will be apparent to those in the art.

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Claims

1. A method comprising:

applying a first ultrasound pulse to biological tissue to create shear waves in the biological tissue;

5 simultaneously transmitting a plurality of focused ultrasound pulses into the biological tissue;

receiving a plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses;

10 detecting the shear waves in the biological tissue based on the received one or more ultrasound signals;

determining a shear wave propagation velocity based on the detected shear waves; and

displaying the shear wave propagation velocity.

15 2. A method according to claim 1, wherein detecting the shear waves comprises calculating a correlation, the sum of absolute differences (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD) or the sum of absolute power differences (SPD) between the received ultrasound signals at one or multiple positions in time.

20

3. A method according to claim 1, wherein determining the at least one propagation property comprises calculating a correlation, the sum of absolute differences (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD) or the sum of absolute power differences (SPD) between the detected shear waves at one

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or multiple instances.

4. A method according to claim 1 wherein the plurality of focused ultrasound pulses comprise coded waveform signals.

5

5. A method according to claim 4 wherein the coded waveform signals comprise one or more of Chirp codes, Barker codes, Golay codes or Hadamard codes.

6. A method according to claim 1, wherein detecting the shear waves comprises determining a displacement of the biological tissue.

10

7. A method according to claim 1, wherein detecting the shear waves comprises determining a velocity of the biological tissue using a color Doppler technique.

15

8. A method according to claim 1, wherein the shear wave propagation velocity is calculated based on the square root of the ratio between a temporal second-order derivative of the displacement of the biological tissue and a spatial second-order derivative of the displacement of the biological tissue.

20

9. A method according to claim 6, wherein determining a displacement of the biological tissue comprises calculating a time integral of tissue color Doppler velocity.

10. A method according to claim 1, further comprising transmitting a second focused ultrasound pulse from a transducer after transmitting a first focused ultrasound

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pulse and before the first focused ultrasound pulse returns to the transducer from a deepest position in an ultrasound field.

11. A method according to claim 1, wherein applying the first ultrasound pulse
5 comprises applying a plurality of ultrasound pulses to the biological tissue to create shear waves in the biological tissue; and

wherein each of the plurality of ultrasound pulses is focused at a different focal point.

10 12. A method according to claim 1, wherein transmitting the plurality of focused ultrasound pulses comprises transmitting the plurality of focused ultrasound pulses into the biological tissue more than one time; and

wherein receiving the plurality of ultrasound signals from the biological tissue
comprises receiving the plurality of ultrasound signals from the biological tissue
15 generated in response to the plurality of focused ultrasound pulses at one or more instances.

13. A non-transitory medium storing processor-executable program code, the program code executable by a device to:

20 apply a first ultrasound pulse to biological tissue to create shear waves in the biological tissue;

simultaneously transmit a plurality of focused ultrasound pulses into the biological tissue;

25 receive a plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses;

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detect the shear waves in the biological tissue based on the received one or more ultrasound signals;

determine a shear wave propagation velocity based on the detected shear waves;
and

5 display the shear wave propagation velocity.

14. A medium according to claim 13, wherein detection of the shear waves comprises calculation of a correlation, the sum of absolute differences (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD) or the sum of
10 absolute power differences (SPD) between the received ultrasound signals at one or multiple positions in time.

15. A medium according to claim 13, wherein determination of the at least one propagation property comprises calculating a correlation, the sum of absolute differences
15 (SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD) or the sum of absolute power differences (SPD) between the detected shear waves at one or multiple instances.

16. A medium according to claim 13, wherein the plurality of focused ultrasound
20 pulses comprise coded waveform signals.

17. A medium according to claim 16, wherein the coded waveform signals comprise one or more of Chirp codes, Barker codes, Golay codes or Hadamard codes.

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18. A medium according to claim 13, wherein detection of the shear waves comprises determination of a displacement of the biological tissue.

19. A medium according to claim 13, wherein detection of the shear waves
5 comprises determination of a velocity of the biological tissue using a color Doppler technique.

20. A medium according to claim 13, wherein the shear wave propagation velocity is calculated based on the square root of the ratio between a temporal second-order
10 derivative of the displacement of the biological tissue and a spatial second-order derivative of the displacement of the biological tissue.

21. A medium according to claim 18, wherein determination of a displacement of the biological tissue comprises calculation of a time integral of tissue color Doppler
15 velocity.

22. A medium according to claim 13, further comprising transmission of a second focused ultrasound pulse from a transducer after transmission of a first focused ultrasound pulse and before the first focused ultrasound pulse returns to the transducer
20 from a deepest position in an ultrasound field.

23. A medium according to claim 13, wherein application of the first ultrasound pulse comprises application of a plurality of ultrasound pulses to the biological tissue to create shear waves in the biological tissue; and

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wherein each of the plurality of ultrasound pulses is focused at a different focal point.

24. A medium according to claim 13, wherein transmission of the plurality of focused ultrasound pulses comprises transmission of the plurality of focused ultrasound pulses into the biological tissue more than one time; and

wherein reception of the plurality of ultrasound signals from the biological tissue comprises reception of the plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses at one or more instances.

25. A system comprising:

a memory storing processor-executable program code; and

a processor to execute the processor-executable program code in order to cause the system to:

apply a first ultrasound pulse to biological tissue to create shear waves in the biological tissue;

simultaneously transmit a plurality of focused ultrasound pulses into the biological tissue;

receive a plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses;

detect the shear waves in the biological tissue based on the received one or more ultrasound signals;

determine a shear wave propagation velocity based on the detected shear waves;

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and

display the shear wave propagation velocity.

26. A system according to claim 25, wherein detection of the shear waves
5 comprises calculation of a correlation, the sum of absolute differences (SAD), the sum of
square differences (SSD), the sum of absolute cubic differences (SCD) or the sum of
absolute power differences (SPD) between the received ultrasound signals at one or
multiple positions in time.

10 27. A system according to claim 25, wherein determination of the at least one
propagation property comprises calculating a correlation, the sum of absolute differences
(SAD), the sum of square differences (SSD), the sum of absolute cubic differences (SCD)
or the sum of absolute power differences (SPD) between the detected shear waves at one
or multiple instances.

15

28. A system according to claim 25, wherein the plurality of focused ultrasound
pulses comprise coded waveform signals.

29. A system according to claim 28, wherein the coded waveform signals
20 comprise one or more of Chirp codes, Barker codes, Golay codes or Hadamard codes.

30. A system according to claim 25, wherein detection of the shear waves
comprises determination of a displacement of the biological tissue.

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31. A system according to claim 25, wherein detection of the shear waves comprises determination of a velocity of the biological tissue using a color Doppler technique.

5 32. A system according to claim 25, wherein the shear wave propagation velocity is calculated based on the square root of the ratio between a temporal second-order derivative of the displacement of the biological tissue and a spatial second-order derivative of the displacement of the biological tissue.

10 33. A system according to claim 30, wherein determination of a displacement of the biological tissue comprises calculation of a time integral of tissue color Doppler velocity.

15 34. A system according to claim 25, the processor further to execute the processor-executable program code in order to cause the system to:

transmit a second focused ultrasound pulse from a transducer after transmission of a first focused ultrasound pulse and before the first focused ultrasound pulse returns to the transducer from a deepest position in an ultrasound field.

20 35. A system according to claim 25, wherein application of the first ultrasound pulse comprises application of a plurality of ultrasound pulses to the biological tissue to create shear waves in the biological tissue; and

wherein each of the plurality of ultrasound pulses is focused at a different focal

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point.

36. A system according to claim 25, wherein transmission of the plurality of focused ultrasound pulses comprises transmission of the plurality of focused ultrasound pulses into the biological tissue more than one time; and

wherein reception of the plurality of ultrasound signals from the biological tissue comprises reception of the plurality of ultrasound signals from the biological tissue generated in response to the plurality of focused ultrasound pulses at one or more instances.

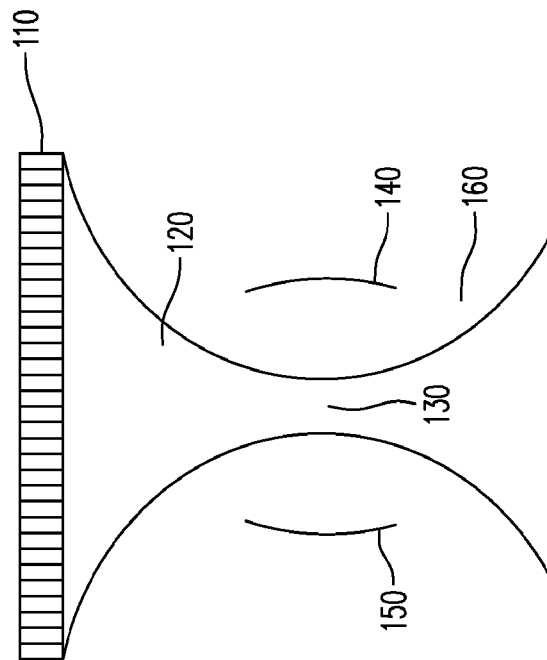


Figure 1

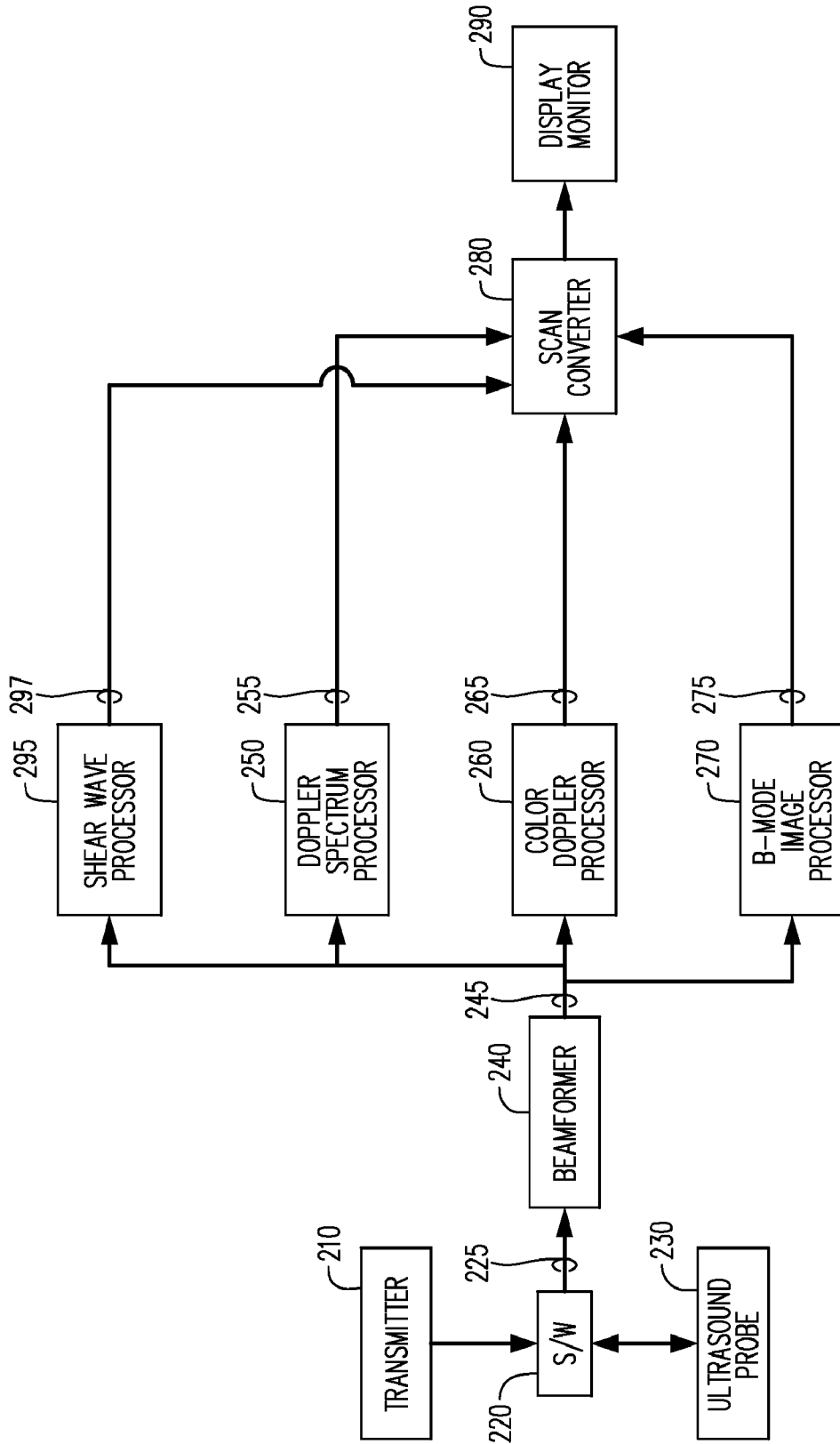


Figure 2

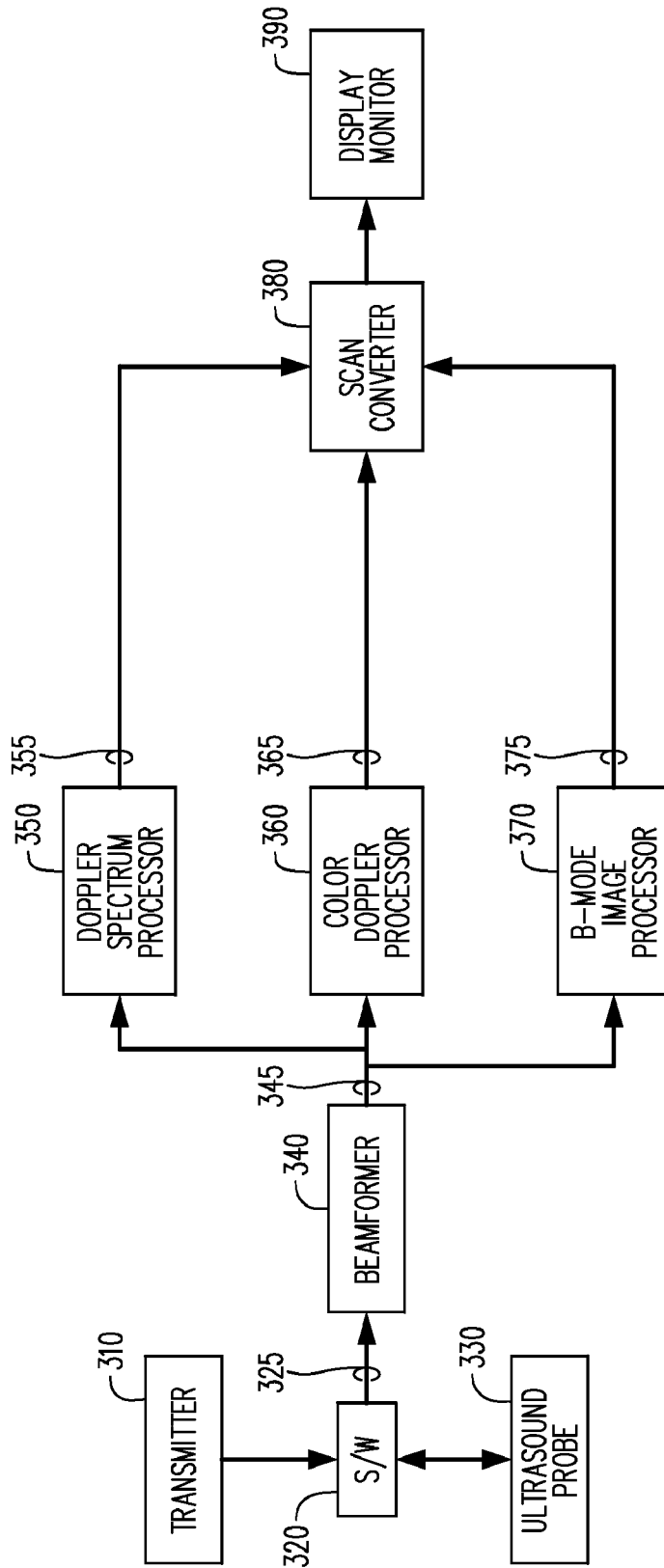


Figure 3
(PRIOR ART)

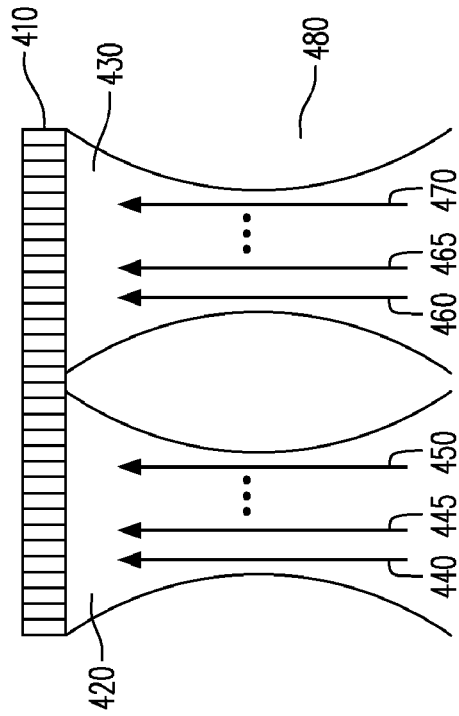


Figure 4

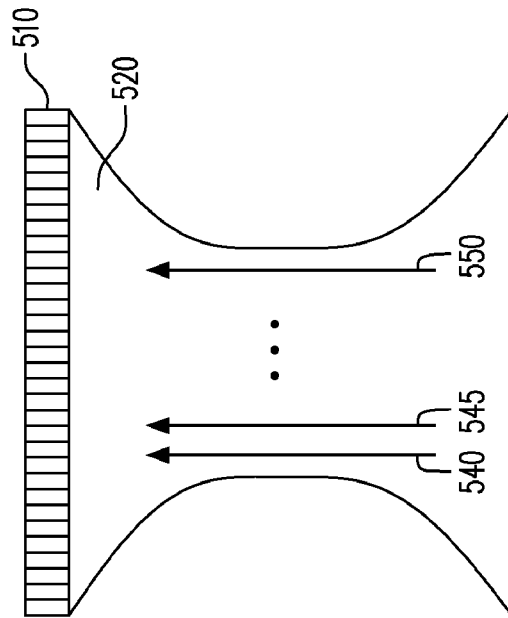


Figure 5

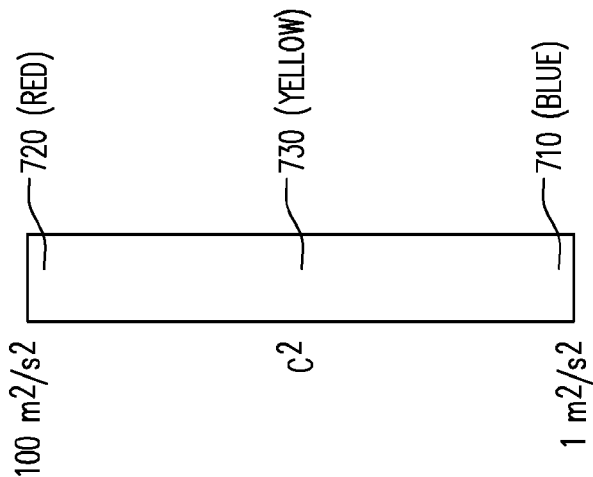


Figure 6

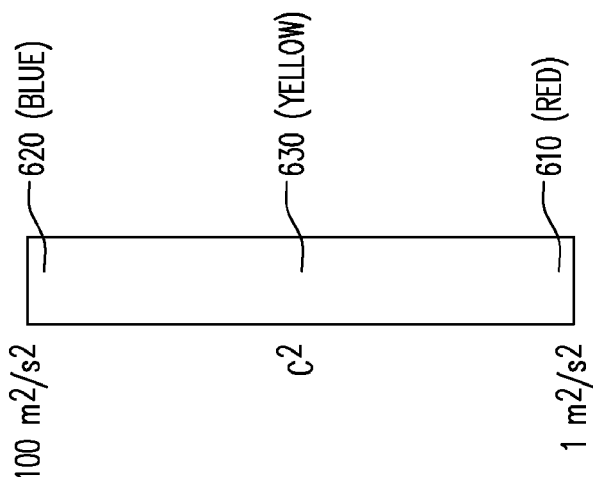


Figure 7

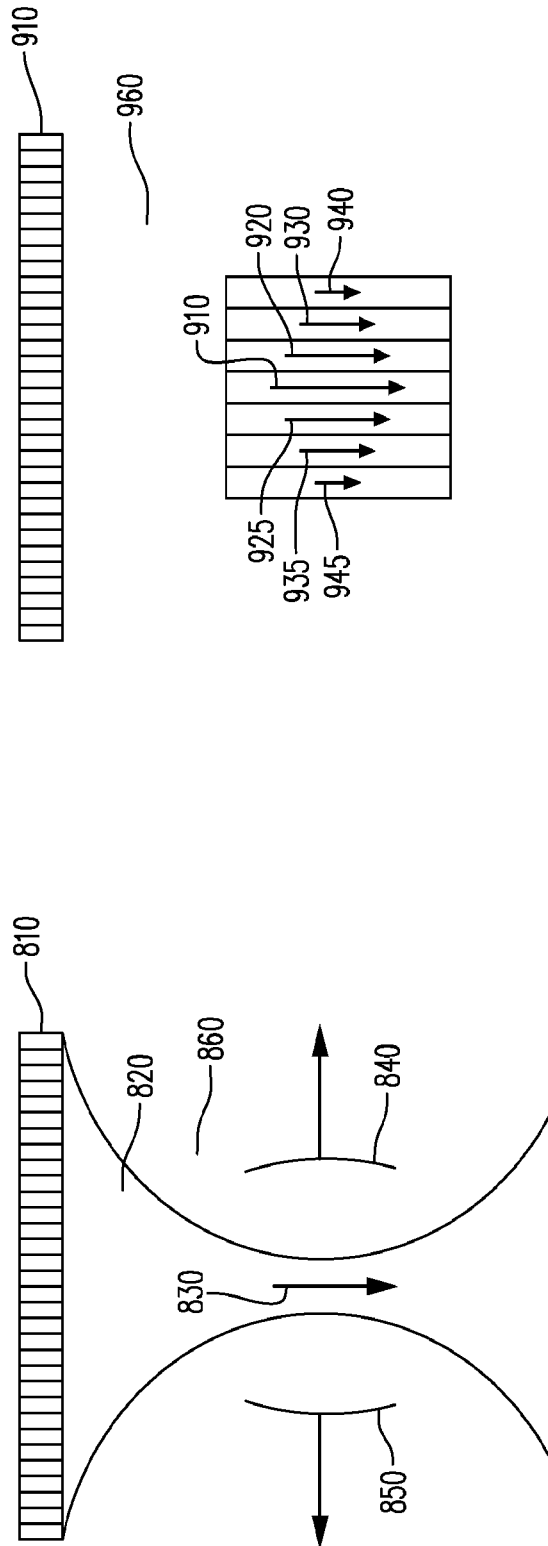


Figure 9

Figure 8

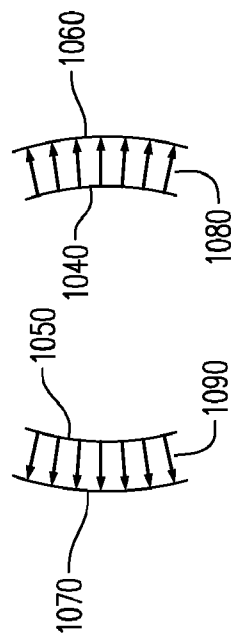
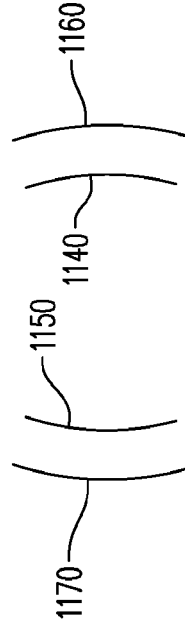


Figure 11

Figure 10

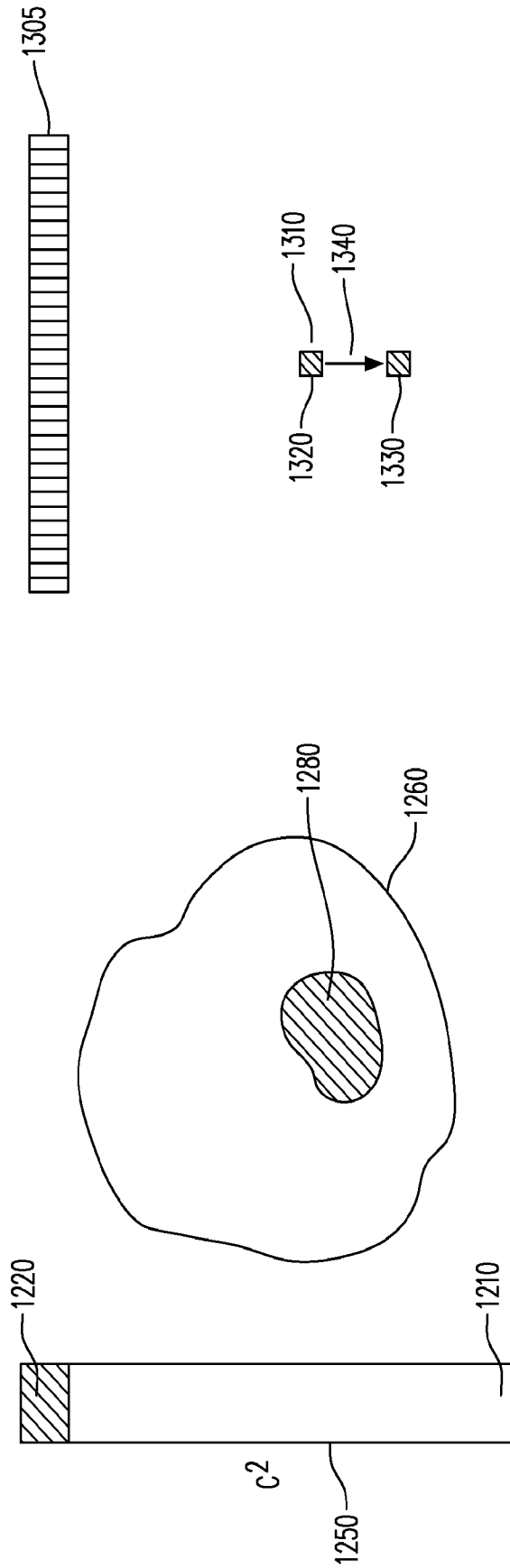


Figure 13

Figure 12

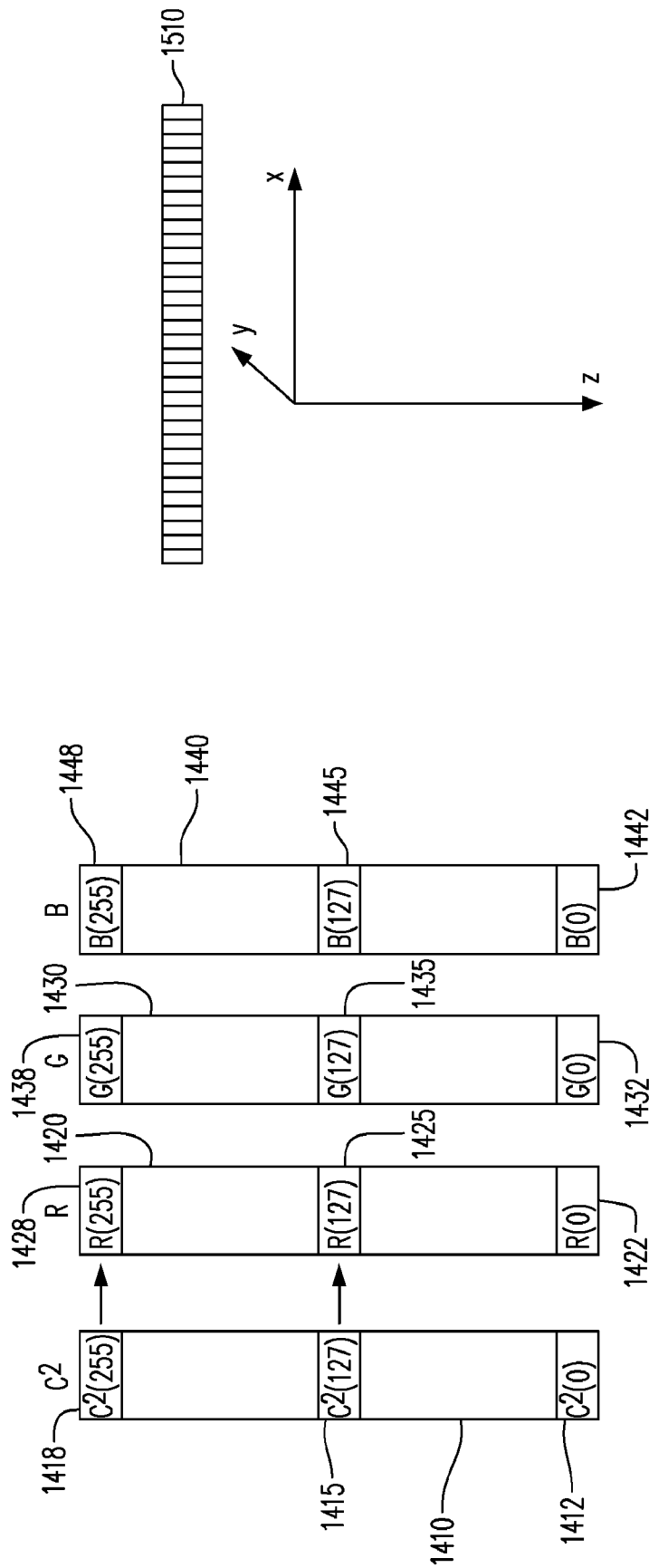


Figure 15

Figure 14

专利名称(译)	用于超声成像的方法和设备		
公开(公告)号	EP2555685A2	公开(公告)日	2013-02-13
申请号	EP2011766381	申请日	2011-03-22
[标]申请(专利权)人(译)	日立阿洛卡医疗株式会社		
申请(专利权)人(译)	日立ALOKA MEDICAL. , LTD.		
当前申请(专利权)人(译)	HITACHI , LTD.		
[标]发明人	TAMURA TADASHI		
发明人	TAMURA, TADASHI		
IPC分类号	A61B8/14 A61B8/08 G01S15/89 G06F19/00		
CPC分类号	A61B8/485 A61B8/08 A61B8/463 A61B8/488 G01S7/52036 G01S7/52042 G01S7/52071 G01S7/52095 G01S15/8979		
优先权	61/321341 2010-04-06 US 13/030831 2011-02-18 US 61/350585 2010-06-02 US 61/321005 2010-04-05 US		
其他公开文献	EP2555685A4		
外部链接	Espacenet		

摘要(译)

将第一超声脉冲施加到生物组织以在生物组织中产生剪切波，将聚焦的超声脉冲发射到生物组织中，从生物组织接收一个或多个超声信号，并且在生物组织中检测剪切波。基于所接收的一个或多个超声信号。确定与检测到的剪切波相关联的至少一个传播特性，并且显示所确定的至少一个传播特性。