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Abstract. Due to their low cost, hand-held convenience, wide selection of bandwidths, and ultrasound imaging capability, linear ultrasonic transducer arrays have been widely studied for photoacoustic computed tomography (PACT). As linear-array PACT suffers from a limited view, full-view imaging requires either the transducer or the object to be rotated. So far, both the central frequencies and bandwidth of linear transducer arrays applied in full-view PACT are low, limiting the spatial resolutions of the reconstructed images. Here, we present a multiview high-frequency PACT imaging system implemented with a commercial 40-MHz central frequency linear transducer array. By rotating the object through multiple angles with respect to the linear transducer array, we acquired full-view photoacoustic pressure measurements. Further, to quantify the unipolar initial pressures and overcome the limitations of the single-view Hilbert transformation, we developed a multiview Hilbert transformation method. The in-plane spatial resolution of this full-view linear-array PACT was quantified to be isotropically 60 μm within a 10 × 10 mm² field of view. The system was demonstrated by imaging both a leaf skeleton and a zebrafish in vivo.

Keywords: photoacoustic computed tomography; high-frequency linear transducer array; Hilbert transformation.

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1 Introduction

Photoacoustic computed tomography (PACT) is an incarnation of photoacoustic tomography that provides optical contrast information in deep biological tissues. Unlike focused-scanningphotoacoustic microscopy (PAM) which forms images by combining A-lines, PACT forms images through reconstruction algorithms. To accelerate imaging speed, PACT is usually implemented with an ultrasonic transducer array employing 64–512 elements to detect photoacoustic (PA) signals in parallel. Currently, both curved and linear transducer arrays are widely used in PACT. Although curved transducer arrays, such as full-ring transducer arrays and partial-ring transducer arrays, have greater detection coverage, they are custom designed, costly, and offer fewer frequency choices. In contrast, linear transducer arrays have the advantages of low cost, wide bandwidth selection, hand-held operation, and ultrasonic imaging integration. Although linear-array PACT suffers from the limited-view problem, rotating either the transducer array or the imaging objects circularly enables full-view linear-array PACT at the expense of a longer imaging time.

Challenges still remain in full-view linear-array PACT. First, no full-view linear-array PACT has been reported with a central ultrasonic frequency higher than 24 MHz, which limits the spatial resolution. A higher central frequency would provide finer lateral resolution, and the associated wider bandwidth would result in higher axial resolution, allowing imaging of finer structures in biological tissues at the expense of penetration. Second, due to the band-pass frequency response of the transducer elements, the reconstructed images present bipolar (i.e., both positive and negative) pixel values unless the raw channel data are deconvolved with the system’s electrical impulse response (EIR) and non-negativity is enforced. Because PACT should ideally image the initial pressure or optical energy deposition, which is always non-negative, bipolar pixel values are artificial. It is also counterintuitive for physicians and biologists to interpret the images because both positive and negative peaks represent strong optical absorption. Moreover, bipolar pixel values are not conducive to quantification of physiological parameters, such as oxygen saturation (sO₂) and blood flow speed.

Thus far, three solutions have been proposed. The first solution is to simply threshold negative signals to zero. However, thresholding removes structures from the image and introduces image artifacts. The second solution is to deconvolve the raw channel data with the EIR. However, this method requires a high signal-to-noise ratio and accurate measurement of each element’s EIR. The third solution is to iteratively reconstruct an image with the non-negativity constraint. However, this method requires both accurate modeling of the whole imaging system and time-consuming computation. Thus, there is an urgent need for a simple and effective method to make bipolar pixel values unipolar.

Hilbert transformation along the acoustic axis recovers the envelope without knowledge of the EIR. Therefore, single-view Hilbert transformation has been widely applied in computing the A-lines of a PAM image and beamformed A-lines of a single-view linear-array PACT image, along the acoustic axis. However, single-view Hilbert transformation cannot be directly

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applied to full-view linear-array PACT because the full-view linear-array PACT has different acoustic axis directions. Hilbert transformation along a single direction will cause artifacts due to the mismatch between the Hilbert transformation direction and acoustic axis. Therefore, we propose a multiview Hilbert transformation method that solves this problem by coherently reconstructing bipolar images in each view, then applying Hilbert transformation along each view’s acoustic axis and taking the absolute value to recover the envelope, and finally adding unipolar single-view envelope images together to form a full-view unipolar image. This algorithm has never been previously reported.

2 Materials and Methods

Here, we present a high-frequency full-view linear-array PACT imaging system, as shown in Fig. 1(a). The illumination source was a 532-nm wavelength laser beam from a second-harmonic generator, which was pumped by a Nd:YAG 1064-nm wavelength laser (Brilliant B, Quantel Inc.) at a repetition rate of 20 Hz. The laser beam was expanded by a circle-pattern engineered diffuser (EDC-5-A-2 s, RPC Photonics Inc.) to a diameter of 20 mm on the object’s surface. With a total energy of 35 mJ per laser pulse, the optical fluence on the object’s surface was estimated to be 11 mJ/cm², well below the American National Standards Institute safety limit (20 mJ/cm²). The excited PA waves were received by a linear transducer array (MS550D, Visualsonics Inc.) with 256 elements and an element-to-element pitch of 55 μm. Each element had a central frequency of 40 MHz and a bandwidth of 33 MHz. The MS550D transducer was connected to an ultrasound/photoacoustic (US/PA) dual modality imaging platform (Vevo LAZR, Visualsonics Inc.), where co-registered US/PA images were acquired and displayed. During experiments, the object to be imaged, either a zebrafish, leaf skeleton, or black microsphere, was fixed on a manual rotation stage (K6X, Thorlabs Inc.) and immersed in water. To get a two-dimensional (2D) full-view linear-array PACT image, samples were rotated, and images were acquired at multiple angles. After imaging, raw data acquired at all view angles were exported from the Vevo LAZR imaging platform to a computer.

A schematic diagram of the reconstruction process is shown in Fig. 1(b), where two coordinate systems are used: the global coordinates $\tilde{r} = (x, y)$ attached to the fixed linear transducer array, and the local coordinates $\tilde{r}_i = (x_i, y_i)$ attached to the sample to be rotated. Both coordinates share the same origin, but the local coordinates rotate with the sample, while the global coordinates are fixed. At rotation step $i$, the single-view PACT image is reconstructed under the global coordinates using the universal filtered backprojection (FBP) algorithm:

$$p_0^{(i)}(\tilde{r}) = \int \frac{d\Omega_0}{\Omega_0} \left[ 2p(\tilde{r}', t) - 2t \frac{\partial p(\tilde{r}', t)}{\partial t} \right]_{\tilde{r}'=\tilde{r}} \frac{d\Omega_0}{\Omega_0},$$

where $p(\tilde{r}', t)$ is the acoustic pressure measured by the transducer element located at $\tilde{r}'$ and time $t$. $p_0^{(i)}(\tilde{r})$ is the reconstructed initial PA pressure for the point at $\tilde{r}$. As previously mentioned, due to the limited view and finite bandwidth of the transducer, the reconstructed $p_0^{(i)}(\tilde{r})$ is bipolar. $c$ is the speed of sound and is assumed to be uniform. $\Omega_0$ is the solid angle subtended by the detection aperture of the transducer array with respect to the point $\tilde{r}$, and $d\Omega_0/\Omega_0$ represents the weighting factor of the contribution of the transducer element located at $\tilde{r}'$. Although the FBP is not a mathematically exact algorithm for the 2-D reconstruction, it can generate boundary-enhanced images that accurately reveal the internal structures of the object. In the future, a more accurate 2-D reconstruction algorithm may be applied.

With the assumption that the light illumination is rotationally symmetric, a full-view bipolar PACT image can be formed in the local coordinates by rotating the reconstructed images acquired at all angles from the global coordinates to the local coordinates, then averaging:

$$p_0(\tilde{r}_i) = \frac{1}{N} \sum_{i=1}^{N} R_{-a_i} [p_0^{(i)}(\tilde{r})].$$

Here, $\tilde{r}_i$ is the image point in the local coordinates, $p_0(\tilde{r}_i)$ is the reconstructed full-view bipolar PACT image, $N$ is the total number of rotation steps (views), $R$ is the rotation operator that transforms an image in the global coordinates $\tilde{r}$ to an image in the local coordinates $\tilde{r}_i$, and $a_i = (i - 1)2\pi/N$ is the rotation angle for the $i$th step.

To reconstruct the full-view unipolar image, we propose a four-step reconstruction procedure:

Step 1: At each rotation angle $a_i$, reconstruct the bipolar PACT image in the global coordinates $(x, y)$, using the universal FBP algorithm.

Step 2: For each bipolar image from step 1, take Hilbert transformation along the depth direction (+z) and calculate the amplitude by taking the absolute value.

Step 3: Rotate the processed image from step 2 by an angle of $-a_i$, transforming it from the global coordinates $(x, y)$ to the local coordinates $(x_i, y_i)$.

Step 4: Repeat steps 1–3 for all rotation angles, and average pixel by pixel over all the images.

Fig. 1 (a) Setup of the high-frequency full-view linear-array PACT system. (b) Schematic of the scan process. PA: photoacoustic. The local coordinates are aligned with the global coordinates when the object is imaged at the first view angle (i.e., $a_i = 0$).
Mathematically, this process can be formulated as

\[ p_n^+ (\mathbf{r}_l) = \frac{1}{N} \sum_{i=1}^{N} R_{-\alpha_i} \{ A [ H [ p_0^+ (\mathbf{r}_l)] ] \}, \]

(3)

where \( p_n^+ (\mathbf{r}_l) \) is the final unipolar image, \( A \) is the absolute value operator and \( H \) is the Hilbert transformation operator. Note that Eq. (3) represents a nonlinear process because of the amplitude extraction, while the conventional FBP algorithm is linear.

3 Results

We imaged a dehydrated leaf skeleton to demonstrate the full-view linear-array PACT. The leaf skeleton was sealed in a 3% agar gel. Figure 2(a) shows the setup of the single-view linear-array PACT, and Figs. 2(b) and 2(c) show a single-view bipolar image and a unipolar image, respectively. Due to the limited view of single-view linear-array PACT, only features with surface normals aligned to the acoustic axis can be reconstructed. Figure 2(d) shows the setup for acquiring six views with an angular step size of 30 deg, and Figs. 2(e) and 2(f) show the corresponding bipolar image and unipolar image of the leaf skeleton, respectively. Compared with the single-view images, multiview images clearly recovered more features. Figure 2(g) shows the full-view PACT setup with 12 views and an angular step size of 30 deg. Figures 2(h) and 2(i) show a full-view bipolar image and unipolar image of the leaf skeleton, respectively. Compared with the six-view images, full-view images revealed even more features. An animation (Video 1) shows the scan setups and corresponding bipolar images and unipolar images of the leaf skeleton at different views. From the movie, we can see that more features of the leaf are gradually revealed as the number of views increases.

Comparing the full-view bipolar [Fig. 2(h)] and unipolar [Fig. 2(i)] images of the leaf skeleton, we can see that the two images have identical features. This identity confirms that the multiview Hilbert transformation followed by amplitude extraction keeps all the features of the original bipolar image. Figure 2(j) compares the profiles of Figs. 2(h) and 2(i) along the dashed lines. The contrast-to-noise ratios (CNRs) of selected features, marked with numbers 1–8 in Fig. 2(j), are presented in Fig. 2(k). Clearly, the full-view unipolar image has a better CNR than the bipolar image, mainly because, after the multiview Hilbert transformation, the noise level has been lowered more than the signals.

The in-plane spatial resolutions of the multiview PACT imaging system were also quantified by imaging 10-μm-diameter black-dyed microspheres (Polysciences Inc.) mixed in an agar gel. Figure 3(a) shows a single-view unipolar PACT image of the microsphere phantom, and Fig. 3(b) shows an enlarged image of a single microsphere, as enclosed by a dashed box in Fig. 3(a). Figure 3(c) shows the PA amplitude distribution along the dashed arrow \( y_1 \) in Fig. 3(b), and the Gaussian fitting result shows that the full width at half maximum (FWHM) is 89 μm, which is the lateral resolution of the single-view PACT system. With the same method, the axial resolution of the single-view PACT along the \( x_1 \) direction was quantified to be 52 μm. Figure 3(d) shows a full-view unipolar PACT image of the microspheres with 18 views and a 20-deg angular step size. Figure 3(e) is an enlarged image of the same microsphere shown in Fig. 3(b). It is obvious that the full-view microsphere image is isotropic in the \( x_1y_1 \) imaging plane. Figure 3(f) shows the PA amplitude distribution along the dashed arrow \( y_2 \) in Fig. 3(e), and the Gaussian fitting result shows that the FWHM value is 60 μm, less than the single-view FWHM of 89 μm along the direction in Fig. 3(b). The isotropic resolution of the full-view unipolar PACT is slightly worse than the axial resolution of the single-view PACT due to the blurring effect of the incoherent sum in the unipolar image reconstruction process.

With isotropic high resolution in the imaging plane, full-view linear-array PACT can resolve fine structures in biological tissues. The zebrafish is a major model organism for biomedical research, but low-frequency PACT cannot resolve its tiny features. Here, we use the high-frequency full-view PACT system to perform \( in vivo \) imaging of a zebrafish embedded in a low gelling temperature agarose (A9414, Sigma). All animal work was performed in compliance with Washington University’s
institutional animal protocols. During the scan process, the zebrafish was alive and its heart was beating. Figure 4(a) shows a single-view bipolar PACT image of the zebrafish. Only the trunk vessel (TV) and part of the eye can be identified. Figure 4(b) is the corresponding unipolar image of Fig. 4(a) obtained by Hilbert transformation and amplitude extraction along the acoustic axis. It shows a clearer boundary of the zebrafish but still suffers from the limited-view problem. Figure 4(c) is a full-view bipolar PACT image of the zebrafish with 18 views and a 20-deg angular step size. The eye now appears round, and

Fig. 3 Characterization of the full-view linear-array PACT system. (a) Single-view unipolar image of 10 μm microspheres, (b) enlarged image of the single microsphere in the dashed box of (a), (c) photoacoustic amplitude distribution along the dashed line in (b), (d) full-view unipolar image of 10 μm microspheres, (e) enlarged image of a single microsphere in the dashed box of (d), (f) photoacoustic amplitude distribution along the dashed line in (e).

Fig. 4 In vivo images of a zebrafish. (a) Bipolar single-view PACT image, (b) unipolar single-view PACT image, (c) bipolar full-view PACT image with an enlarged image at the lower left corner representing the dashed area, (d) unipolar full-view PACT image with an enlarged image at the lower left corner representing the dashed area, (e) unipolar full-view ultrasound image, (f) co-registered US/PA image of the zebrafish obtained by merging (d) and (e). Note that the spine and the trunk vessel (TV) can be clearly differentiated in (f). CF, caudal fin; DF, dorsal fin; SB, swim bladders; US, ultrasound; VF, ventral fin.
we can begin to discern the ventral fin and dorsal fin in addition to the TV. Figure 4(d) is a corresponding full-view unipolar PACT image, showing clearer features, such as boundaries, tails, and tiny bones close to the TV. Although Fig. 4(d) looks better than Fig. 4(c), they actually show the same details, as proved by comparing the features of the enlarged sections from the same area. Figure 4(e) shows a full-view unipolar ultrasound image of the zebrafish achieved by adding the beamformed enveloped ultrasound image at each rotation angle. Since the ultrasound image resolves mechanical contrasts, we can clearly see the swim bladders and spine. Figure 4(f) is a fused image of Figs. 4(d) and 4(e), and it shows both image contrasts (mechanical and optical contrasts). We can clearly identify the TV and spine in Fig. 4(f).

4 Conclusions

In summary, we present a high-frequency full-view linear-array PACT imaging system paired with a multiview Hilbert transformation method. The system has an isotropic spatial resolution of 60 μm within the imaging plane. Such a high resolution allows us to clearly identify different vessels and organs in a live zebrafish. Compared with the conventional FBP method, the multiview reconstruction and Hilbert transformation method render much clearer envelope images of the zebrafish. If the unipolar image recovered by applying the single-view Hilbert transformation along the acoustic axis and taking the absolute value correctly represents optical absorption, the multiview Hilbert transformation images, which are achieved by linearly adding unipolar images from different views, should also correctly represent the optical absorption. The maximum imaging depth of the single-view linear-array PACT system was ~10 mm in this study. By rotating the sample 360 deg, we are able to image samples with diameters up to ~20 mm. Currently, the imaging speed is limited by the slow manual rotation scan. Therefore, the imaging speed of full-view linear-array PACT is lower than that of circular-array PACT. However, linear-array PACT provides a low cost solution with more frequency choices. Moreover, its imaging speed can be further improved by using a faster laser system and a motorized rotation scan setup.

By applying side illumination plus elevational scanning, we will be able to acquire three-dimensional (3-D) whole-body images of small animals. Compared with whole-body PA macroscopy,16 the full-view linear-array PACT has higher in-plane image quality due to the full-view detection, but the 3-D whole-body image quality is worse because of the relatively poor elevational resolution of the linear transducer array.

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References


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