Super-resolution ultrasound imaging method for microvasculature in vivo with a high temporal accuracy

Jaesok Yu1,2, Linda Lavery1 & Kang Kim1,2,3,4

Traditional ultrasound imaging techniques are limited in spatial resolution to visualize angiogenic vasa vasorum that is considered as an important marker for atherosclerotic plaque progression and vulnerability. The recently introduced super-resolution imaging technique based on microbubble center localization has shown potential to achieve unprecedented high spatial resolution beyond the acoustic diffraction limit. However, a major drawback of the current super-resolution imaging approach is low temporal resolution because it requires a large number of imaging frames. In this study, a new imaging sequence and signal processing approach for super-resolution ultrasound imaging are presented to improve temporal resolution by employing deconvolution and spatio-temporal-interframe-correlation based data acquisition. In vivo feasibility of the developed technology is demonstrated and evaluated in imaging vasa vasorum in the rabbit atherosclerosis model. The proposed method not only identifies a tiny vessel with a diameter of 41 μm, 5 times higher spatial resolution than the acoustic diffraction limit at 7.7 MHz, but also significantly improves temporal resolution that allows for imaging vessels over cardiac motion.

Acute coronary syndromes (ACS), a leading cause of morbidity and mortality in the US and Europe, is generally caused by the plaque rupture or erosion1–4. Extensive efforts to characterize and predict vulnerable plaques assessing a few known markers have been made for several decades1–7. It has been found that abnormally dense neovascularization of the vessel wall, often infiltrating into the plaque core, is associated with development of atherosclerotic plaque and progression of the disease4,5. These micro-vasculatures may lead to intraplaque hemorrhage, which typically accompanies unstable plaque4,5,6. Therefore, abnormal proliferation of adventitial vasa vasorum (VV) is an important clinical imaging target to assess vulnerability of the atherosclerotic plaques4–8,10. However, the lack of adequate noninvasive and high-resolution imaging technology to visualize VV is a big challenge. Micro-CT, optical coherent tomography (OCT), intravascular ultrasound (IVUS) angiography and contrast-enhanced ultrasound (CEU) imaging have been demonstrated to image VV in pre-clinical studies, but these technologies have suffered from hazardous radiation (micro-CT), poor imaging depth (OCT), invasive approach (IVUS) and insufficient spatial resolution (CEU)5,6,11–19.

CEU is one of the well-established imaging modalities to evaluate microvasculature in vivo animal study using gas-filled microbubbles that can provide high echogenic contrast6,13,17. In previous studies with CEU using a mid-frequency transcutaneous linear array transducer (Transmit at 7 MHz and receive at 15 MHz), they have shown the correlation of adventitial VV density based on ultrasonic image intensity with VV progression6,13,17. However, identifying individual VV, especially those near the lumen of the main vessel, was limited mainly due to low spatial resolution at mid-frequency6. Because of this limitation with spatial resolution, the accuracy of VV density measure therefore was limited. High spatial resolution is essential for separating individual tiny VV...
from the lumen of the main vessel to improve the accuracy of adventitial VV density as an indicator of VV progression. A high-frequency transcutaneous linear array transducer centered at 40 MHz has shown its spatial resolution as high as 67 μm in a tissue-mimicking phantom but with a limited imaging depth less than 14 mm. A high-frequency single element transducer ranging 30–150 MHz is preferably utilized in IVUS. However, conventional (30–50 MHz) and high frequency (90–150 MHz) IVUS are limited to the spatial resolution (>100 μm) and the imaging depth (<2 mm), respectively. Recently, multi-frequency IVUS (or super-harmonic IVUS) has been widely studied to maintain the advantage of both deep imaging depth (>2.5 mm) and improved spatial resolution (<50 μm) by incorporating relatively low frequency transmit (30–50 MHz) and high frequency receive (90–150 MHz). However, this technology was only evaluated in vitro or ex vivo so far, therefore in vivo study should be followed. In addition, IVUS is not ideal if considering eventual translation with full noninvasiveness. Therefore, transcutaneous imaging approach using a mid-frequency linear array transducer commonly used in the clinic would be sought after.

Super-resolution US imaging technology has been recently introduced to overcome the limitation of inherent spatial resolution of US imaging defined by the acoustic diffraction limit. This approach utilizes two state-of-the-art technologies; tissue rejection and microbubbles localization technique. It is known that ultrafast plane wave imaging in general significantly improves the performance of the eigen decomposition based adaptive clutter filtering technique, outperforming in suppressing stationary signal that comes from the clutters, compared to conventional clutter filtering techniques used in typical Doppler imaging. Moreover, microbubble center localization technique allows to estimate each location of microbubbles in sub-pixel level precision. Each microbubble could be considered as a point source because its size (~3 μm) is much smaller than the spatial resolution of the imaging system that operates at mid-frequency ultrasound of around 5–10 MHz (100–200 μm). Therefore, received echo signal from individual microbubble would be presented as a point spread function (PSF) of the imaging system, and each microbubble is expected to be located at the centroid of the PSF. After summing up localized microbubbles in blood flow at different times over a large number of frames, a vascular network image in high spatial resolution beyond the acoustic diffraction limit can be formed. Errico et al. successfully reconstructed single static super-resolved image of rat brain vasculature network, identifying the micro-vasculature in diameter as small as 9 μm full-width at half-maximum (FWHM) with using a 20 MHz linear array transducer. However, the major drawback of this method requires a long data acquisition time of 150 seconds for a total of 75,000 frames at 500 frames per second. Requiring a long data acquisition time makes this imaging technology susceptible to motion artifacts and therefore hinders widespread of this technology for various applications.

Here, we present a new approach of super-resolution US imaging technology to achieve a high temporal resolution as well. A strategically designed approach in two steps is employed to drastically improve temporal resolution; (1) deconvolution localization technique to reduce data acquisition time, and (2) spatio-temporal-interframe-correlation (STIC) based data acquisition to compensate motion over reduced data acquisition time. First, applying deconvolution separates each center of microbubbles from densely grouped microbubbles while previous approach, 2D Gaussian fitting, typically requires discarding such frames in which clumped microbubbles cannot be separated. As such, deconvolution technique can significantly reduce data acquisition time. However, reduced data acquisition time using deconvolution only is not short enough to ignore fast physiologic motion such as cardiac-dependent motion. In addition to deconvolution technique, therefore, STIC data acquisition inspired from 3D fetal echo-cardiology imaging technique is adapted to further overcome fast physiologic motion. STIC acquisition allows it to synchronize collected images over multiple cardiac cycles. In vivo feasibility of the developed technique in identifying VV is demonstrated in rabbit atherosclerotic plaque model.

Results

Deconvolution based super-resolution ultrasound imaging. In Fig. 1, our developed imaging technique on a rabbit femoral artery is compared conventional imaging technologies. The white rectangle represents the balloon injured area that is expected to develop the plaque with VV, and white arrows in all images indicate a big branch near the femoral artery that is used as the landmark to match the image plane among different modalities. Cadence harmonic contrast-enhanced imaging technique with using microbubbles operated by a commercial ultrasound scanner (Accuson Sequoia 512, SIEMENS, Mountain view, CA) shown in Fig. 1(a) is a typical standard method in the clinic to evaluate microvasculature perfusion. However, this technique is limited in spatial resolution to identify and separate VV from the major vessel such as the femoral artery in this study. Conventional B-mode image shown in Fig. 1(b) only illustrates structural information of bifurcation area of the rabbit femoral artery and small vasculatures surrounding the femoral artery cannot be identified due to limited spatial resolution. To amplify signal from the blood, power Doppler based temporal maximum intensity persistence (MIP) imaging technique (4 frames = 2,000) implemented in Verasonics system is used as shown in Fig. 1(c). This method employs a tissue-rejection filter that suppresses stationary signal from tissue and extracts only signal from flowing microbubbles. However, its spatial resolution is still insufficient to clearly identify individual VV. The proposed super-resolution ultrasound imaging method implemented in Verasonics system shown in Fig. 1(d) delineates detail of microvasculature that is shown blurred in other imaging methods due to low spatial resolution that is mainly limited by diffraction limit of the operating frequency. To estimate maximum spatial resolution of our developed method, the detectable smallest vessel in the ROI is chosen among discernable vasculatures in the Fig. 2(a) (Selected region of Fig. 1(d)). FWHM on lateral axis of the selected vessel is estimated to be 41 micron that is 5-fold smaller than wavelength as shown in Fig. 2(b).

Three sequential data-set are independently acquired using our super-resolution imaging sequence to verify repeatability of our proposed method. For three independent imaging sessions, bifurcation of femoral artery and the branched vessel indicated in white arrow in Fig. 1 are identified as landmark to maintain the same imaging plane. Figure 3(a) depicts temporal MIP vascular imaging using the first data set of 2,000 frames, and Fig. 3(b–d)
Figure 1. Comparison of several imaging modalities. (a) CadenceTR contrast-enhanced imaging with microbubbles acquired by commercial ultrasound scanner (Sequoia 512, Siemens), (b) conventional B-mode imaging, (c) temporal MIP vascular imaging, (d) proposed super-resolution imaging of ROI. Same raw data is used to reconstruct images (b–d). The white solid rectangle represents balloon-injured area that the plaque is expected to be developed. The white arrow indicates the same vessel branch that shows a correlation of images acquired by two different ultrasound scanners. White dashed rectangle represents selected ROI used in Figs 2 and 3.

Figure 2. The spatial resolution of the proposed imaging method. (a) The detectable smallest vessel is chosen (white solid line) in the ROI of Fig. 1(b–d) indicated by the white dashed rectangle, (b) Spatial profile of the selected vessel. FWHM is estimated by 41 μm (<λ/5).
represents reconstructed images from sequentially acquired the first, second and third data set (2,000 frames for each acquisition), respectively using our super-resolution imaging sequence. Note that Fig. 3(a,b) are reconstructed from same raw dataset. An obviously same major vessel is observed at the center in all three reconstructed images to assure the same imaging plane among three imaging sessions. Some microvasculature in the background shown slightly different can be attributed to tilted imaging plane within three different imaging sessions.

In vivo feasibility of the developed super-resolution technique in imaging vasa vasorum on a rabbit femoral artery. Conventional B-mode imaging and super resolution imaging for vascular network of both side femoral artery, injured (experimental side) and contralateral non-injured side (control side), are shown in Fig. 4. In B-mode images (Fig. 4(a,d)), adventitia area is represented in the yellow dotted line. Figure 4(b,c,e,f) show super-resolution images at diastole and systole on injured and the non-injured side, respectively. Supplementary Movies (1 and 2) for an entire cardiac cycle are provided. Each image at different stage of cardiac cycle is reconstructed from around 300 frames synchronized by STIC algorithm. The abundance of VV in adventitia indicated by white arrows can be clearly observed on the injured-side in the super-resolution ultrasound imaging. Normalized VV density on the injured-side is 0.027 ± 0.004, which is approximately 3 times greater than non-injured side with 0.010 ± 0.001. Furthermore, apparent uneven surface on the vessel wall on the injured side is observed most likely due to plaque formation on the lumen wall. The vessel wall is overall thickened (injured: 410 μm vs non-injured: 220 μm) and the lumen diameter is decreased (injured: 1.1 mm vs non-injured: 1.4 mm) in injured side. These observations are more clearly confirmed by closely looking at blood flowing through VV from adventitia into media shown in Supplementary Movies (1 and 2).

VV's are more populated and believed to be infiltrated into the medial area of injured area, while less populated VV are found only in adventitial and connective tissue area in non-injured area. Atherosclerotic plaque formation and corresponding VV development is evidenced by histology of femoral artery section shown in Fig. 5. The atherosclerotic lesions are characterized on injured side by neointimal proliferation, shown in Fig. 5(b) by hematoxylin and eosin (H&E) stain and abnormally enriched VV development on adventitia, shown in Fig. 5(d).
by immunofluorescence of von Willebrand Factor (vWF) stain. Pink solid line in Fig. 5(b) was drawn along endothelium in the tunica intima of the lumen. Total cholesterol level is measured 291 mg/dL that is approximately 3 times higher than standard range (<100 mg/dL) due to feeding high-fat diet.

Discussion
It has been shown that super-resolution ultrasound imaging technique has strong potential to extend the application of ultrasound imaging with unprecedented high spatial resolution24,25,32,33. We further developed this imaging technology to overcome the limitation of the temporal resolution, which makes it capable of imaging of fast event. Our proposed super-resolution imaging method successfully assesses VV density on the plaque in a rabbit femoral artery with enhanced temporal resolution as well as high spatial resolution. In our approach, deconvolution and STIC data acquisition were investigated to improve temporal resolution of US sub-diffraction imaging. It should be noted that deconvolution technique is known to be sensitive to noise as discussed in method section. Thresholding used in this study can successfully suppress noise, however it potentially also suppresses true signal when signal-to-noise ratio is relatively low for example in a fast plane-wave imaging. Therefore, more advanced adaptive noise reduction approach needs to be sought after to maximize signal-to-noise ratio while maintaining high temporal resolution in the future study. In addition, STIC data acquisition is generally used to collect and synchronize 3D data acquired from fast-moving fetal heart when using a relatively slow acquisition speed. It should be noted that there is an assumption in STIC acquisition; cardiac pulsation pattern over data acquisition period is stationary. For further translation of this technique, therefore a thorough performance analysis of STIC and an adaptive algorithm development if needed for non-stationary cardiac pulsation should be ensured. With reduced total number of frames thanks to deconvolution, we were able to generate ten super-resolution images using 3,000 acquired frames that can be collected within 6 seconds. Therefore, our approach has potential for free-hand scanning in the clinics because the acquisition time is significantly shorter than free-hand data acquisition time of 3D-fetal echocardiography when using STIC acquisition in the standard clinical protocol that takes around 4~5 minutes on average29,34.

Since this study is concentrated on the technical development and in vivo feasibility demonstration, it is not intended to investigate any correlation between VV development and plaque stage with enough statistics, although this well-established rabbit model has been studied for a long time for various diseases in association with atherosclerotic plaques. Further extended study with an increased number of rabbits and time points is planned in the following study. In addition, the histology is limited for providing anatomically matched cross sections as a gold standard. An established anatomical imaging modality with high spatial resolution is necessary.
as a gold standard to provide more direct comparison, for example, ex-vivo microCT that enables to provide volumetric microvasculature information\(^{35,36}\).

**Methods**

**Deconvolution-based super-resolution US imaging sequence.** Super-resolution imaging sequence was implemented into a fully-programmable ultrasound scanner (Vantage 128, Verasonics, Kirkland, WA, USA) equipped with a mid frequency hockey stick linear array transducer (CL15-7, ATL-Philips, Bothell, WA). Ultrasound plane waves of one and half cycle at frequency of 7.7 MHz were insonified to the target with high pulse repetition frequency (PRF) of 1,500 Hz with three different steering angles (-3°, 0°, 3°) for compounding, therefore the effective frame rate is 500 Hz. Signal processing algorithm was illustrated in Fig. 6. Acquired raw radio-frequency (RF) channel data were beamformed by delay-and-sum algorithm, and then downmixed to the baseband using a quadrature demodulator. The echo signal from microbubbles were extracted from compounded...
analytic baseband signal of 500 frames by using a spatio-temporal eigen-based decomposition clutter filter. Envelope of the echo signal from microbubbles was interpolated for an increased pixel resolution of 15 µm (lateral) × 10 µm (axial) by using the modified akima cubic hermite method and deconvolved with the measured system PSF to localize each microbubble. Richardson-Lucy (RL) deconvolution is a non-linear iterative deconvolution method that has been widely used for deblurring image with the presence of Poisson distributed noise in astronomy and biomedical applications. The iterative process is described by

Figure 7. Conceptual demonstration of sub-wavelength localization using deconvolution on synthetic data. (a) PSF of the imaging system. FWHM is assumed as 150 µm. (b) The locations of the two neighboring targets (Ground truth). Two targets are positioned 70 µm apart. (c) The synthetic signal received from two targets is shown in (b) using the imaging system with PSF shown in (a). This signal is modeled as a received signal in the imaging system whose has PSF shown in (a). Two targets cannot be separated in the image due to their distance is shorter than the spatial resolution of the imaging system. (d) Deconvolution results of received signal shown in (c) using the system PSF shown in (a). Two targets are distinctly identified. (e) The synthetic signal received from two targets when noise is added is shown in (b) using the same imaging system. (f) Deconvolution results of (e), where two targets are clearly identified with minimal interference due to noise.
where, \( * \) is the correlation operator, \( \otimes \) is the convolution operator, \( i^{(k)} \) is the estimated image after \( k \) iterations, \( h \) is the PSF of the imaging system and \( g \) is blurred image modeled by \( g = h \otimes i^{(0)} + \text{noise} \). The concept of this approach is illustrated in Fig. 7.

The simulated echo signal (Fig. 7(c)) is represented by convolving the PSF of the imaging system (Fig. 7(a)) with two point-targets (Fig. 7(b)). Figure 7(c,e) shows the received signal (indicated by 'Overall') from two targets separated by 70 \( \mu \text{m} \), smaller than the spatial resolution of the system. With using deconvolution (\( \# \)s of iteration \( = 100 \)), however, this signal can be separated into two different targets as shown in Fig. 7(d,f). A potential problem with the RL deconvolution algorithm is noise amplification, which is known to be generic for all maximum likelihood techniques. Therefore, any false target can be expected when signal-to-noise ratio is not high enough due to noise-sensitive RL deconvolution. For example, in Fig. 7(e,f) deconvolving the received signal from two targets with significant Gaussian white noise resulted in imperfect target locations. This high sensitivity to noise can be alleviated with several available standard regulation approaches. In this study, thresholding at \(-10 \text{ dB}\) was applied to pre-deconvolved signal to suppress noise component.

Figure 8. Estimated cardiac pulsation by counting the numbers of flowing microbubbles (blue solid line). After applying low-pass-filtering, the frames with minimum value are chosen as reference frames for synchronization (red solid line).

Figure 9. Graphical diagram of STIC data acquisition. Sequentially acquired multiple datasets are synchronized to form a single cardiac cycle event based on the cardiac pulsation estimated by the numbers of bubbles.
Spatio-Temporal-Interframe-Correlation (STIC) based data re-alignment algorithm. STIC data re-alignment algorithm that synchronizes among collected cardiac cycles based on estimated cardiac pulsation is developed to monitor fast physiological event under limited imaging speed. To capture rapid physiological dynamics at a limited frame rate, sequentially acquired RF data over multiple cardiac cycles can be synchronized based on the cardiac cycle period estimated from the periodically changing signal intensity, which reflects the number of microbubbles. Rigid motion in frame-by-frame caused by respiratory or operator dependent motion was compensated by applying an offset to match acquired images. Offset was estimated by taking 2-dimensional cross-correlation between frames. If correlation coefficient between images is smaller than 0.9, the image can be excluded. The eigen-based spatio-temporal tissue rejection filter technically removes all stationary tissue information except for moving objects, such as microbubbles. At diastolic phase, microbubbles are less likely flow, therefore smaller number of moving microbubbles are detected. On the contrary, the number of detected microbubbles significantly increases as microbubbles are moving fast toward the systole. Low-pass filter is applied to the detected signal from flowing microbubbles to estimate the cardiac period. Figure 8 illustrates an example of the number of detected microbubbles as a function of time (blue solid line) over 2 seconds totaling 1,000 data points. The reference frame for synchronization (red solid line) was chosen at the minima after low pass filtering. The estimated period of 3 Hz by this approach has good agreement with the heart rate recorded by electrocardiogram of 290–310 beats per minute. Figure 9 shows overall graphical diagram of STIC acquisition method. First, we collected raw ultrasound RF data sequentially over multiple cardiac cycles. Acquired US frames are synchronously aligned based on cardiac period estimated by the numbers of microbubbles. Re-aligned images are integrated to form a super-resolution image of single cardiac cycle. In this study, we collected continuous 3,000 frames over 6 seconds, which is equivalent to approximately 15–30 cardiac cycles. Each cardiac cycle was divided by 10 sections and entire dataset was aligned based on section number. Aligned frames in each section were summed to reconstruct single super-resolution frame. Therefore, single cardiac cycle containing 10 reconstructed frames with super-resolution was generated and each super-resolution image contains around 300 acquired frames collected over multiple cardiac cycles.

Rabbit atherosclerotic plaque model. A New Zealand white rabbit (3.5 kg) was fed a high fat and cholesterol diet (cholesterol 1%, peanut oil 2.5%, and fat 10%) over 6 weeks to accelerate development of atherosclerosis6. Balloon injuries were induced to the superficial femoral artery on the right side of an anaesthetized rabbit (ketamine 150 mg IM, xylazine 8 mg IM and 2.5% inhaled isofluorane) by using a 2F Fogarty balloon catheter (Edwards Life Sciences, CA, USA) one-week after feeding. The balloon inflated at 2 atm was moved back and forth several times to apply injury to the vessel walls around bifurcation area under guidance of high-frequency ultrasound scanning (Vevo2100 equipped with a linear array transducer centered at 18.5 MHz, FUJIFILM Visualsonics, Toronto, On, Canada). Upon completion of imaging, blood of 5 mL was obtained to measure cholesterol concentration.

Ultrasound imaging protocol. At twelve weeks after surgery, ultrasound imaging were performed injecting a commercial microbubbles (Definity, Lantheus Medical Imaging, N. Billerica, MA). Mean diameter of microbubbles ranges 1.1–3.3 μm, and concentration of microbubbles is $1.2 \times 10^{10}$ bubbles/mL. A bolus of 0.2 mL microbubbles was intravenously administrated through an ear vein access catheter for each imaging session. An anaesthetized rabbit (same protocol of surgery) was prepared with hair-shaved legs (Fig. 10). US gel was applied for ultrasonically coupling. Super-resolution US imaging sequence was performed to the surgical side and thereafter contralateral side by using a fully programmable research ultrasound scanner (Vantage 128, Verasonics, Kirkland, WI) equipped with a hockey stick linear array transducer (CL15-7, ATL-Philips, Bothell,
A transducer holder was used to fix the transducer. Compounded plane wave images with 3 different angles of total 9,600 frames were acquired in each data acquisition. To verify repeatability, total three datasets were collected on the possibly same imaging plane. All signal processing was performed offline using MATLAB software (Mathworks, Natick, MA). Harmonic MIP imaging was sequentially conducted on the same location by using a commercial ultrasound imaging equipped with a linear array transducer (Acuson Sequoia 512 with 15L8, SIEMENS, Mountain view, CA). To quantify degree of vasa vasorum development, vessel density was calculated by

\[
Vasa\ Vasorum\ density = \frac{\sum\text{Microbubble signal intensity on adventitia area}}{\text{ adventitia area}}
\]

where, adventitia area was manually selected on the B-mode. The animal protocol for this study was approved by Institutional Animal Care and Use Committee (IACUC) of University of Pittsburgh. All experimental procedures of surgery and \textit{in vivo} imaging were performed in accordance with the all guidelines and regulations detailed in the approved protocol.

**Histology and immunofluorescence.** Femoral arteries were perfusion fixed using paraformaldehyde, and then carefully excised along with a block of the surrounding muscle to save neighbor vasculature in the connective tissues. Excised tissues were then carefully fixed in 4% paraformaldehyde, embedded in the paraffin-block and serially sectioned. Cross-sections of the paraffin block were stained with hematoxylin and eosin (H&E) and Von Willebrand Factor (vWF) for endothelium staining. In briefly, for antigen retrieval, treatment dewaxed slides were heated in sodium citrate buffer (pH 6.0, abcam, Cambridge, MA) at 95 °C for 30 min in blocking solution of goat serum and incubated with the anti-vWF antibody (Millipore, Burlington, MA) at 4 °C overnight. Incubated slides were washed with PBS 3 times for 3 mins. The slides were incubated with Cy3-conjugated IgG (Jackson ImmunoResearch Laboratories, West Grove, PA) and counterstained with DAPI. The stained slides were observed by using a fluorescent microscope (IX-81, Olympus, Center Valley, PA). Brightness (+40%) and contrast (+40%) of acquired fluorescence images were adjusted to improve visibility by using ImageJ software. The adventitia area was manually selected on the B-mode. The animal protocol for this study was approved by Institutional Animal Care and Use Committee (IACUC) of University of Pittsburgh. All experimental procedures of surgery and \textit{in vivo} imaging were performed in accordance with the all guidelines and regulations detailed in the approved protocol.

**References**

1. Fowkes, F. G. \textit{et al.} Edinburgh Artery Study: prevalence of asymptomatic and symptomatic peripheral arterial disease in the general population. \textit{Int. J. Epidemiol.} \textbf{20}, 384–92 (1991).
2. Fuster, V. \textit{et al.} Atherothrombosis and High-Risk Plaque: Part II: Approaches by Noninvasive Computed Tomographic/Magnetic Resonance Imaging. \textit{J. Am. Coll. Cardiol.} \textbf{46}, 1209–1218 (2005).
3. Naghavi, M. \textit{et al.} From Vulnerable Plaque to Vulnerable Patient. \textit{Circulation} \textbf{108} (2003).
4. Moreno, P. R. \textit{et al.} Plaque Neovascularization Is Increased in Ruptured Atherosclerotic Lesions of Human Aorta. \textit{Circulation} \textbf{110} (2004).
5. Taruya, A. \textit{et al.} Vasa Vasorum Restructuring in Human Atherosclerotic Plaque Vulnerability. \textit{J. Am. Coll. Cardiol.} \textbf{65} (2015).
6. Moguilansky, D. \textit{et al.} Quantification of plaque neovascularization using contrast ultrasound: a histologic validation. \textit{Eur. Heart J.} 32, 646–653 (2011).
7. Koon, T.-G., Lerman, L. O. & Lerman, A. The Vasa Vasorum in Atherosclerosis. \textit{J. Am. Coll. Cardiol.} \textbf{65} (2015).
8. Portanova, A. \textit{et al.} Intracranial Vasa Vasorum: Insights and Implications for Imaging. \textit{Radiology} \textbf{267}, 667–679 (2013).
9. Kologode, F. D. \textit{et al.} Intraplaque Hemorrhage and Progression of Coronary Atheroma. \textit{N. Engl. J. Med.} \textbf{349}, 2316–2325 (2003).
10. Xu, J., Lu, X. & Shi, G.-P. Vasa vasorum in atherosclerosis and clinical significance. \textit{Int. J. Mol. Sci.} \textbf{16}, 11574–608 (2015).
11. Moritz, R. \textit{et al.} Quantification of vasa vasorum density in multiple slice computed tomographic coronary angiograms: role of computed tomographic image voxel size. \textit{J. Comput. Assist. Tomogr.} \textbf{34}, 273–81 (2010).
12. Arcidiacono, M. V. \textit{et al.} Left carotid adventitial vasa vasorum signal correlates directly with age and with left carotid intima-media thickness in individuals without atheromatous risk factors. \textit{Cardiovasc. Ultrasound} \textbf{13}, 20 (2015).
13. Staub, D. \textit{et al.} Contrast-Enhanced Ultrasound Imaging of the Vasa Vasorum. \textit{JACC Cardiovasc. Imaging} \textbf{3}, 761–771 (2010).
14. Moritz, R. \textit{et al.} IVUS detection of vasa vasorum blood flow distribution in coronary artery vessel wall. \textit{JACC Cardiovasc. Imaging} \textbf{5}, 935–40 (2012).
15. Cheng, J. M. \textit{et al.} \textit{In vivo} detection of high-risk coronary plaques by radiofrequency intravascular ultrasound and cardiovascular outcome: results of the Atheroremo-Ivus study. \textit{Eur. Heart J.} \textbf{35}, 639–647 (2014).
16. Kume, T. \textit{et al.} \textit{In vivo} assessment of vasa vasorum neovascularization using intravascular ultrasound: A comparison between acute coronary syndrome and stable angina pectoris. \textit{J. Cardiol.} \textbf{69}, 601–605 (2017).
17. You, X. \textit{et al.} Relationship between enhanced intensity of contrast enhanced ultrasound and microvessel density of aortic atherosclerotic plaque in rabbit model. \textit{PloS One} \textbf{9}, e92445 (2014).
18. Huang, C.-C., Chen, P.-Y., Peng, P.-H. & Lee, P.-Y. 40 MHz high-frequency ultrasound imaging. \textit{Med. Phys.} \textbf{44}, 2185–2195 (2017).
19. Ma, T. \textit{et al.} Multi-frequency intravascular ultrasound (IVUS) imaging. \textit{IEEE Trans. Ultrason. Ferroelectr. Freq. Control} \textbf{62}, 97–107 (2015).
20. Lee, J., Moon, J.-Y. & Chang, I. A 35 MHz/105 MHz Dual-Element Focused Transducer for Intravascular Ultrasound Tissue Imaging Using the Third Harmonic. Sensors \textbf{18}, 2290 (2018).
21. Wang, Z., Martin, K. H., Dayton, P. A. & Jiang, X. Real-time ultrasound angiography using superharmonic dual-frequency (2.25 MHz/30 MHz) cylindrical array: \textit{In vitro} study. \textit{Ultrasonsics} \textbf{82}, 298–303 (2018).
22. Ma, J. \textit{et al.} Design factors of intravascular dual frequency transducers for super-harmonic contrast imaging and acoustic angioffgraphy. \textit{Phys. Med. Biol.} \textbf{60}, 3441–3457 (2015).
23. Siepmann, M., Schmitz, G., Bzyl, I., Palomowski, M. & Kiessling, F. Imaging tumor vascularity by tracing single microbubbles. In 2011 \textit{IEEE International Ultrasonics Symposium} 1906–1909, \url{https://doi.org/10.1109/ULTSYM.2011.0476} (IEEE, 2011).
24. Errico, C. \textit{et al.} Ultrafast ultrasound localization microscopy for deep super-resolution vascular imaging. \textit{Nature} \textbf{527}, 499–502 (2015).
25. Christensen-Jeffries, K., Browning, R. J., Tang, M. X., Dunsby, C. & Eckersley, R. J. \textit{In vivo} acoustic super-resolution and super-resolved velocity mapping using microbubbles. \textit{IEEE Trans. Med. Imaging} \textbf{34}, 433–440 (2015).
26. Demeen, C. \textit{et al.} Spatiotemporal Clutter Filtering of Ultrafast Ultrasound Data Highly Increases Doppler and \textit{f}Ultrasound Sensitivity. \textit{IEEE Trans. Med. Imaging} \textbf{34}, 2271–2285 (2015).
27. Rost, M. J., Bates, M. & Zhuang, X. Sub-diffraction-limit imaging by stochastic optical reconstruction microscopy (STORM). \textit{Nat. Methods} \textbf{3}, 793–796 (2006).
28. Betzig, E. et al. Imaging Intracellular Fluorescent Proteins at Nanometer Resolution. Science (80-.), 313, 1642–1645 (2006).
29. Viñals, F., Poblete, P. & Giuliano, A. Spatio-temporal image correlation (STIC): a new tool for the prenatal screening of congenital heart defects. Ultrasound Obstet. Gynecol. 22, 388–394 (2003).
30. Rissanen, T. T. et al. High-Resolution Ultrasound Perfusion Imaging of Therapeutic Angiogenesis. JACC Cardiovasc. Imaging 1, 83–91 (2008).
31. Lee, S. C. et al. Temporal Characterization of the Functional Density of the Vasa Vasorum by Contrast-Enhanced Ultrasonography Maximum Intensity Projection Imaging. JACC Cardiovasc. Imaging 3, 1265–1272 (2010).
32. Bar-Zion, A., Tremblay-Darveau, C., Solomon, O., Adam, D. & Eldar, Y. C. Fast Vascular Ultrasound Imaging with Enhanced Spatial Resolution and Background Rejection. IEEE Trans. Med. Imaging 36, 169–180 (2017).
33. Cox, B. & Beard, P. Imaging techniques: Super-resolution ultrasound. Nature 527, 451–452 (2015).
34. Zhao, L. et al. Feasibility Study on Prenatal Cardiac Screening Using Four-Dimensional Ultrasound with Spatiotemporal Image Correlation: A Multicenter Study. Plos One 11, e0157477 (2016).
35. Gössl, M., Malyar, N. M., Rosol, M., Beighley, P. E. & Ritman, E. L. Impact of coronary vasa vasorum functional structure on coronary vessel wall perfusion distribution. Am. J. Physiol. - Hear. Circ. Physiol. 285, H2019–H2026 (2003).
36. Yoo, Y. M. et al. Adaptive clutter filtering for ultrasound color flow imaging. Ultrasound Med. Biol. 29, 1311–20 (2003).
37. Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH Image to ImageJ: 25 years of image analysis. Nat. Methods 9, 671–675 (2012).

Acknowledgements
Small animal imaging system (Vevo2100) was supported by the NIH grant 1S10RR027383-01. Authors thank to Dr. You-Jin Choi, Dr. Hyunjung An and Dr. Jianhui Zhu for great helps and discussions in preparation and analysis of histology samples.

Author Contributions
Jaesok Yu conceived the idea, developed algorithm, designed and performed experiments, processed and analyzed the data, drafted and finalized the manuscript. Linda Lavery performed animal experiments and discussed the results. Kang Kim conceptualized and designed all experiments, supervised the project, analyzed the results, and outlined and edited the manuscript. All authors proof read the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-32235-2.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.