QUANTIFICATION OF ENDOTHELIAL $\alpha_v\beta_3$ EXPRESSION WITH HIGH-FREQUENCY ULTRASOUND AND TARGETED MICROBUBBLES: IN VITRO AND IN VIVO STUDIES

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Abstract—Angiogenesis is a critical feature of plaque development in atherosclerosis and might play a key role in both the initiation and later rupture of plaques. The precursory molecular or cellular pro-angiogenic events that initiate plaque growth and that ultimately contribute to plaque instability, however, cannot be detected directly with any current diagnostic modality. This study was designed to investigate the feasibility of ultrasound molecular imaging of endothelial $\alpha_v\beta_3$ expression in vitro and in vivo using $\alpha_v\beta_3$-targeted ultrasound contrast agents (UCAs). In the in vitro study, $\alpha_v\beta_3$ expression was confirmed by immunofluorescence in a murine endothelial cell line and detected using the targeted UCA and ultrasound imaging at 18-MHz transmit frequency. In the in vivo study, expression of endothelial $\alpha_v\beta_3$ integrin was measured using targeted UCA and high-frequency ultrasound in seven animals. Our results indicated that endothelial $\alpha_v\beta_3$ expression was significantly higher in the carotid arterial wall containing atherosclerotic lesions than in arterial segments without any lesions. We also found that the salivary gland can be used as an internal positive control for successful binding of targeted UCA to $\alpha_v\beta_3$ integrin. In conclusion, $\alpha_v\beta_3$-targeted UCA allows non-invasive assessment of the expression levels of $\alpha_v\beta_3$ on the vascular endothelium and may provide potential insights into early atherosclerotic plaque detection and treatment monitoring. (E-mail: v.daeichin@erasmusmc.nl or http://www.erasmusmc.nl/thoraxcenterbme/) © 2016 World Federation for Ultrasound in Medicine & Biology.

Key Words: Molecular imaging, Targeted microbubble, $\alpha_v\beta_3$ integrin, High-frequency ultrasound, Murine atherosclerosis, Endothelial cell, In vitro, In vivo.

INTRODUCTION

Ultrasound molecular imaging features high sensitivity, availability, rapid execution of imaging protocols and relatively low cost (Kaufmann and Lindner 2007; van Rooij et al. 2015), and its potential for imaging biological processes at the molecular level has been illustrated. The key element of this technique compared with regular diagnostic ultrasound imaging is the use of ultrasound contrast agents (UCAs) decorated with binding ligands such as antibodies and small peptides. These functionalized UCAs, so-called targeted UCAs (tUCAs) or targeted microbubbles, can bind to biomarkers involved in various disease processes. As UCAs are blood pool agents (Greis 2009), tUCAs can only bind to intravascular biomarkers on intravenous administration. Combined with dedicated ultrasound imaging sequences and the latest transducer technology, ultrasound molecular imaging allows quantitative assessment of molecular target expression with high sensitivity. The aforementioned features of this technique have opened new diagnostic applications, including the detection of atherosclerosis (Kaufmann 2009; Khanicheh et al. 2013a, 2013b; Liu et al. 2013; Shim et al. 2015), thrombosis (Alonso et al. 2007),...
neovascularization (Deshpande et al. 2011; Leong-Poi et al. 2003; Shelton et al. 2016; Stieger et al. 2008; Streeter et al. 2013), lymph nodes (Hauff et al. 2004) and inflammation (Betinger et al. 2012; Davidson et al. 2014; Lindner 2010; ten Kate et al. 2010). It also provides helpful insights into the genesis, progress and prevention of diseases (Dayton and Ferrara 2002; Khanicheh et al. 2013a; Klibanov 2006; Lanza et al. 1996; Lindner 2004; Liu et al. 2013; Palmowski et al. 2008; Shim et al. 2015; Streeter et al. 2013; van Rooij et al. 2015).

Atherosclerosis is characterized by accumulation of lipids, inflammatory cells and connective tissue within the arterial wall. A number of pathophysiologic mechanisms are thought to be involved in the progression of an atherosclerotic lesion into a vulnerable plaque (Naghavi et al. 2003). Intraplaque neovascularization and adventitial vasa vasorum have gained interest as a preceding or associated factor in the development, progression and instability of atherosclerotic plaques (Barger et al. 1984; Hellings et al. 2010; Langheinrich et al. 2007; Michel et al. 2011; Moreno et al. 2006; Sluimer et al. 2008). The presence of intraplaque neovascularization has been identified as an independent predictor of intraplaque hemorrhage and plaque rupture (Moreno et al. 2004; Purushothaman et al. 2003). Intraplaque neovascularization is therefore increasingly investigated as a marker for the non-invasive identification of vulnerable plaques. It has been reported that angiogenic activity in the atherosclerotic plaques coincides with \( \alpha_v \beta_3 \)-integrin by the adventitial vasa vasorum and intraplaque neovascularization (Brooks et al. 1994; Haubner 2006; Hoshiga et al. 1995; Hynes 1987). Therefore, \( \alpha_v \beta_3 \) integrin is an appropriate biomarker for targeted ultrasound imaging of intraplaque neovascularization and adventitial vasa vasorum, whereas adhesion of tUCAs to endothelial cells provides the signal enhancement. This molecular marker has been used for ultrasound molecular imaging of angiogenesis in tumors and hindlimb ischemia (Ellegala 2003; Leong-Poi et al. 2003). Although \( \alpha_v \beta_3 \) integrin expression in atherosclerotic plaques has been quantified \textit{ex vivo} using near-infrared optical imaging in mice (Tedesco et al. 2009) and \textit{in vivo} with magnetic resonance imaging in rabbits (Winter et al. 2003) and positron emission tomography in mice (Laitinen et al. 2009), to the best of our knowledge, no \textit{in vivo} ultrasound molecular imaging of adventitial vasa vasorum and/or intraplaque neovascularization with \( \alpha_v \beta_3 \) integrin has been reported.

Here, we provide \textit{in vitro} and \textit{in vivo} proof-of-concept for molecular imaging of \( \alpha_v \beta_3 \) in carotid artery lesions in apolipoprotein E-deficient (ApoE\textsuperscript{−/−}) mice with proven expansion of adventitial vasa vasorum (Rademakers et al. 2013). We hypothesized that molecular imaging of \( \alpha_v \beta_3 \)-integrin with tUCA could be used to quantify the adventitial vasa vasorum underlying atherosclerosis plaques. To test our hypothesis, attachment of \( \alpha_v \beta_3 \)-targeted MicroMarker UCA to endothelial SVEC cells was evaluated \textit{in vitro}. Attachment of UCA \textit{in vivo} and signal enhancement of the carotid plaques were assessed in the ApoE\textsuperscript{−/−} mouse model.

**METHODS**

A high-frequency pre-clinical ultrasound scanner, with a 256-element linear array transducer (Vevo 2100 with MS250 probe, VisualSonics, Toronto, ON, Canada) was used at 18 MHz. The transducer was operated at 10% transmit power, and a wide beam-width setting was chosen to have a low, more uniform transmit pressure over depth (Needles et al. 2010) (~400 kPa, mechanical index = 0.1). Side-by-side B-mode and non-linear contrast mode (amplitude modulation [Needles et al. 2010]) images were acquired at a frame rate of 20 frames/s and a dynamic range of 35. UCA was prepared from Target-Ready Vevo MicroMarker UCA (VisualSonics) according the manufacturer’s instructions. The UCA was targeted with biotinylated antibodies against \( \alpha_v \beta_3 \) integrin (Catalog No. 551380, Clone RMV-7, BD Biosciences, San Jose, CA, USA), an antibody previously successfully used to prepare \( \alpha_v \beta_3 \)-tUCA for ultrasound molecular imaging of tumor angiogenesis (Leong-Poi et al. 2003). An inactive form of biotinylated IgG(\( \kappa \)) (Catalog No. 553923, BD Biosciences) was used for making control UCA (cUCA).

**In vitro**

Murine SV40 virus-modified endothelial cells (SVEC) were purchased from ATCC (LGC Standards, We- sel, Germany) and cultured in Dulbecco’s Modified Eagle’s Medium/Ham’s F-12 (Gibco, Life Technologies Europe, Bleiswijk, Netherlands) with penicillin–streptomycin solution (1% v/v solution containing 5,000 units penicillin [Gibco] and 5 mg streptomycin [Gibco] per milliliter) and 10% heat-inactivated fetal calf serum (Gibco) in a humidified atmosphere of 5% CO\textsubscript{2} at 37°C. Cells were cultured up to passage 5 in acoustically transparent OptiCell (NUNC, Wiesbaden, Germany) chambers on only one of the two membranes until 100% confluence. \( \alpha_v \beta_3 \) expression was determined by fluorescence immunostaining. The cells were fixed in 4% formaldehyde for 5 min and blocked in 5% goat serum in phosphate-buffered saline (PBS) solution for 1 h. After fixation, 1 × 1-cm pieces of the OptiCell membrane containing cells were cut out. These specimens were incubated with rat anti-mouse antibody against \( \alpha_v \beta_3 \) integrin (1:100; BD Biosciences) for 1.5 h. After thorough washing with PBS, the specimens were incubated with goat anti-rat antibody conjugated with AlexaFluor 488 (1:1,000; Invitrogen, Groningen, Netherlands) for 1 h. As a control, fixed cells were blocked and incubated with rat anti-mouse antibody against \( \alpha_v \beta_3 \) integrin only or goat anti-rat antibody conjugated with
AlexaFluor488 only. All procedures were carried out at room temperature. After incubation, the specimens were washed with PBS and embedded in Moviol (Sigma-Aldrich, Zwijndrecht, Netherlands) with Hoechst 33342 (Invitrogen; 5 μg/mL) to stain the cell nuclei. Fluorescence microscopy was performed with a fluorescence Olympus microscope equipped with an AxioCam MRc camera (Carl Zeiss, Sliedrecht, Netherlands) and 40× water-immersed objective (LUMPlanFL, NA 0.8, Olympus).

To assess the adhesion of tUCAs to SVEC cells, the cells were cultured as previously described, and 30 μL of αvβ3-tUCA or cUCA was added to the cells. The OptiCell was then turned cell side up, to let UCA adhere to the cells by flotation, and incubated at 37°C for 5 min. After incubation, the OptiCell was inverted (cell side down) and incubated for another 5 min at 37°C to allow UCA that did not adhere to the cells to float to the top membrane of the OptiCell chamber. For each specimen, 20 random fields of view were acquired for both UCA adhering to cells on the bottom and free-floating UCA on top of the fields of view were acquired for both UCA adhering to cells on the bottom and free-floating UCA on top of the OptiCell. The ratio of adhered UCA to total adhered cells on the bottom and free-floating UCA on top of the OptiCell was calculated and presented as the percentage of attached UCA.

To determine whether UCA adherent to αvβ3-integrin can be detected by high-frequency non-linear contrast imaging, cells were incubated with tUCA as described above, and then the membrane of the OptiCell that did not contain the cells was cut out. Cells were washed with 37°C fresh warm medium and placed in the tank filled with PBS, the temperature of which was maintained at 37°C. The 18-MHz probe was then positioned above the OptiCell. To prevent the overwhelming specular reflection from the OptiCell interface and to obscure the scattering signal from the sample under examination received by the US transducer, the OptiCell was placed tilted at a 20° angle with respect to the direction of the ultrasound beam; thus, specular reflected US signals from the OptiCell could not reach the US transducer (Klibanov 2006). First, a 2-D image was acquired, and the tUCA in this 2-D imaging plane were destroyed using a flash burst. In addition to the 2-D images, 3-D B-mode and non-linear contrast mode images were acquired by mounting the probe on a linear stepper motor (VisualSonics). The probe was translated over ~4.9 mm, while acquiring frames every 0.032 mm (beam overlap percentage ~50%, pixel size of 0.0262 mm in contrast mode). The series of images was then rendered in 3-D using the Vevo 2100 3-D Mode software (VisualSonics).

In vivo
Animal experiments were approved by the regulatory authority of Maastricht University and performed in compliance with Dutch government regulation guidelines. Hypercholesterolemic male ApoE knockout mice (n = 7) on a C57/B16 background from an in-house breeding colony (originally from Iffa Credo, Lyon, France) were fed normal chow until the age of 50 wk. Mice were anesthetized with ketamine (0.1 g/kg) and xylazine (0.02 g/kg) by subcutaneous injection and pre-operative ibuprofen (subcutaneously). The animals were intubated and artificially ventilated using room air at the rate of 160 strokes/min and stroke volume of 250 μL (Mouse Microvent, Hugo Sachs, Germany). The left jugular vein was exposed, and a heat-stretched polyethylene-25 cannula was inserted (1.5 cm) and subcutaneously guided to the neck of the mouse. Here the catheter was fixed with sutures, extended, filled with heparinized saline (10 U/mL) and plugged. Animals were placed on the Vevo Rail System heating stage (VisualSonics), and ultrasound imaging was performed. For each animal, cUCA and tUCA (100 μL each) were injected with 20-min intervals between injections using a programmable syringe pump (11 Pico Plus Elite, Harvard Apparatus, Kent, UK) at the rate of 600 μL/min. The focus of the transducer was set at 7 mm, at the upper wall of the right common carotid artery. A field of view of the right common carotid artery and its bifurcation was chosen. The right salivary gland located above the right carotid was also in the imaging field. B-Mode gain was 20 dB, and contrast gain was 30 dB. DICOM (Digital Imaging and Communications in Medicine) images were exported to MATLAB (Version R2012 b, The MathWorks, Natick, MA, USA) for further off-line processing.

After injection of the UCA, about 10 min is required for UCA clearance from the blood (van Rooij et al. 2015). This waiting period also leaves more time for the tUCA to adhere to the αvβ3 binding sites. Next, a series of about 300 frames (~10 s) was acquired with high-intensity ultrasonic bursts (flash with standard internal destruction burst of 10-cycle square envelope at 18 MHz and 100% power for a total duration of exposure of 1 s) in the middle to disrupt the UCA in the imaging plane.

Subsets of the images (100 frames) captured 10 min after injecting the UCA were used for quantification of the bound UCA in the region of interest (ROI) around the carotid plaques, the healthy part of the carotid artery and in the salivary gland. The captured frames with a flash in the middle to destroy the UCA in the imaging plane were separated into two groups: pre-flash and post-flash groups (50 frames each from pre- and post-flash groups). The noise levels in each data set were obtained by drawing a ROI in the background of a contrast frame immediately after the flash pulse and calculating the mean intensity value (μ) and standard deviation (σ) in the ROI. Then, all image intensities below a noise threshold level of (μ + 4) × σ were set to zero (Daeichin et al. 2015a). Next, averaging was applied on all the frames of each pre- and post-flash group, resulting in a single mean image
for each of these groups. The averaging emphasizes the signal of stationary contrast spots. Each of the single pre- and post-flash mean images was then smoothed using a 2-D Gaussian filter of $3 \times 3$ pixels with sigma = 1 to remove speckles smaller than the characteristic bubble size $(0.0786 \times 0.0786 \text{ mm})$. The resulting post-flash mean images were subtracted from the pre-flash counterpart to cancel signals of non-stationary contrast spots, stationary artifacts and remaining tissue signal. In the subtracted mean images, mean intensity within each ROI was calculated and presented as molecular signals. Respiratory motion artifacts were absent and therefore not relevant.

**Tissue handling**

At sacrifice, blood was drawn from the right ventricular apex. Subsequently the mice were perfusion fixed via the left cardiac ventricles with 20 mL sodium nitroprusside (0.1 mg/mL in PBS, Sigma-Aldrich, Seelze, Germany), followed by 1% paraformaldehyde (PFA). The right carotid arteries and salivary glands were excised, fixed in 1% PFA for 24 h and embedded in paraffin. The right carotid arteries and salivary glands were serially sectioned (4 μm) and stained at 20-μm intervals with hematoxylin and eosin (HE, Sigma-Aldrich) for microscopic examination. Serial sections were subjected to heat-induced epitope retrieval (DAKO target retrieval) and stained with rat anti-mouse MAC3 (1:400, clone M3/84, Becton & Dickinson) for 1 h at room temperature to visualize plaque macrophages. Slides were subsequently incubated with biotin-conjugated rabbit anti-rat secondary antibody (Jackson Immunoresearch), followed by horseradish peroxidase-conjugated rabbit anti-rat antibody (BrightVision, Immunologic) to amplify signal and visualized using diaminobenzidine.

**Statistical evaluation**

Statistical analysis for integrin expression and number of UCA microbubbles adherent to the SVEC cells was performed using an unpaired t-test. In vivo, quantified molecular signal based on video intensity of contrast-enhanced sonograms of processed images in seven animals was analyzed using a paired t-test and expressed as medians and interquartile ranges ($p < 0.01$ was considered to indicate significance).

**RESULTS**

**In vitro**

Immunofluorescence staining of α3β2-integrin expression on SVEC cell membranes (Fig. 1f) revealed that the ligand is expressed on the cell surface, especially in cell–cell border regions. No fluorescence was observed when only the α3β2-antibody or only the goat anti-rabbit antibody conjugated with AlexaFluor488 was used. These stainings also prove that the α3β2-tUCA used in this study can adhere to murine endothelial cells. To examine the binding affinity of UCA, α3β2-tUCA or cUCA was incubated with live SVEC cells. We observed that significantly more (3.7 times more, $p < 0.0001$) tUCA (Fig. 1b) than cUCA attached to the cells (Fig. 1a).

Figure 2a is a schematic of the imaging setup. Figure 2b illustrates the ultrasonic signals of SVEC cells to which UCA was bound. The contrast-to-tissue ratio was 30 dB. In Figure 2b, the gray-scale B-mode image overlies the contrast-mode image. The band in the middle of Figure 2b showing no green signal indicates the destruction of the tUCA with the flash burst.

In vivo

To examine further the application of α3β2-tUCA for imaging α3β2-integrin expression in vivo, the following locations were examined in ApoE−/− mice: atherosclerotic lesions at the bifurcation of the carotid artery, part of the wall of the common carotid artery containing no atherosclerotic lesions and the salivary gland. Figure 3a is a micrograph of an ApoE−/− mouse indicating the anatomic structure and presence of atherosclerotic lesions in the aortic arch (AA), brachiocephalic trunk (BT) and right carotid bifurcation (RCB). In Figure 3b and c are examples of cross-sectional histopathologic sections revealing the morphology of the carotid artery and the atherosclerosis lesions are presented. Large and advanced atherosclerosis lesions at and around the carotid bifurcation are visible in Figure 3b, and the absence of atherosclerotic lesions in the right common carotid (RCC) artery can be observed in Figure 3c.

Representative B-mode and contrast mode images of the right carotid and salivary gland; analyzed ROIs for the plaque area and salivary gland; and a no-plaque ROI in the common carotid are provided in Figure 4. Similar to in vitro results, there was evident enhancement of the ultrasound signal from the α3β2-tUCA adhering to the vessel walls in the carotid bifurcation (Fig. 4c) and salivary gland (Fig. 4e) after applying our quantification method. Only a weak molecular signal was detected in the case of cUCA injections (Fig. 4d, f). Enhancement of the ultrasound signal in the no-plaque ROI (Fig. 4g, h) was significantly ($p < 0.01$) lower than that in the plaque ROI.
Quantitative analysis of the sonograms of the plaque ROI revealed that the difference between the molecular signals using $\alpha_v\beta_3$-tUCA (4.25 \pm 0.84 arbitrary units [a.u.]) and those using cUCA (1.08 \pm 0.39 a.u.) was statistically significant (Fig. 5). There was also a difference in signal intensity for the salivary gland (6.61 \pm 1.23 a.u. for tUCA vs. 2.63 \pm 0.60 a.u. for cUCA) (Fig. 5). However there was no statistically significant difference for $\alpha_v\beta_3$-tUCA and cUCA in the no-plaque ROI (1.16 \pm 0.24 a.u. for tUCA and 0.41 \pm 0.07 a.u. cUCA) (Fig. 5). Also, the molecular signal in the plaque ROI (4.25 \pm 0.84 a.u.) was significantly higher than that for the no-plaque ROI (1.16 \pm 0.24 a.u.) for the tUCA. As expected, this difference was not significant for the cUCA (1.08 \pm 0.39 a.u. vs. 0.41 \pm 0.07 a.u.).

To validate the results of ultrasonography, the carotid artery and salivary gland of mice were harvested and subsequently analyzed for the presence of blood vessels and macrophages by CD31 and MAC3 staining, respectively. Results indicated expanded adventitial vasa vasorum and numerous macrophages in the carotid artery plaques (Fig. 6a, c) and microvessels and macrophages in the salivary gland (Fig. 6b, d).

Also to validate the results of ultrasonography, the carotid artery and salivary gland of mice were harvested and subsequently analyzed for the presence of $\alpha_v\beta_3$, expression, blood vessels and macrophages by $\alpha_v\beta_3$, expression with targeted UCA.
CD31 and MAC3 staining, respectively. Results indicated the presence of \( \alpha_v \beta_3 \), expanded adventitial vasa vasorum, and numerous macrophages in the carotid artery plaques (Fig. 6a, c) and microvessels and macrophages in the salivary gland (Fig. 6b, d).

**DISCUSSION**

Our *in vitro* and *in vivo* results revealed successful binding of tUCA to \( \alpha_v \beta_3 \)-integrin, as well as successful detection of \( \alpha_v \beta_3 \)-tUCA corresponding to the intensity of \( \alpha_v \beta_3 \)-integrin expression on endothelial cells. Expression of \( \alpha_v \beta_3 \) was confirmed by immunocytochemistry *in vitro*. In *vivo*, immunohistochemical \( \alpha_v \beta_3 \) signals in plaques and the salivary gland are most likely derived from endothelial cells and macrophages, known to express \( \alpha_v \beta_3 \)-integrin (Antonov et al. 2004; Brooks et al. 1994; Hoshiga et al. 1995).

It is known that macrophages play an important role in the pathogenesis of atherosclerosis and strongly correlate with plaque vulnerability (Boyle 2005; Redgrave et al. 2006). The expression of \( \alpha_v \beta_3 \)-integrin on activated macrophages, which are recruited to the arterial wall, had been confirmed earlier (Antonov et al. 2004). However, our results are especially valuable because they can provide a strategy for direct and selective intraplaque neovascularization evaluation in terms of spatial extent and amount of endothelial \( \alpha_v \beta_3 \)-integrin in a non-invasive or minimally invasive manner. Such separation between \( \alpha_v \beta_3 \)-integrin expressed on macrophages and endothelial cells is possible with ultrasound molecular imaging because UCAs remain exclusively intravascular on intravenous administration, thereby minimizing non-specific signals from extravasated contrast material. This separation was not possible for the \( \alpha_v \beta_3 \) expression studies of atherosclerosis with near-infrared optical imaging (Tedesco et al. 2009), magnetic resonance imaging (Winter et al. 2003) and positron emission tomography (Laitinen et al. 2009) because the contrast agents used in these studies are small and can therefore extravasate. Although it can be argued that the micron-sized microbubbles in UCAs can extravasate from the leaky adventitial vasa vasorum, further investigation is required to confirm such a hypothesis.

Numerous investigators have adopted approaches in which ultrasound molecular images have been acquired through the use of UCAs targeted to many biomarkers related to atherosclerosis, such as CD81 (Yan et al. 2012), ICAM-1 (Demos et al. 1999), VCAM-1 (Hamilton et al. 2004; Khanicheh et al. 2013a, 2013b; Liu et al. 2013), P-selectin (Lindner et al. 2001; Liu et al. 2013), platelet glycoprotein Ibα (Khanicheh et al. 2013b; Liu et al. 2013; Shim et al. 2015) and von Willebrand factor (Shim et al. 2015). However, to the best of our knowledge, this is the first study reporting...
successful in vivo ultrasound molecular imaging of adventitial vasa vasorum in atherosclerotic plaques using UCA targeted to αvβ3-integrin in a mouse model of atherosclerosis. The advantage of using αvβ3-integrin as a target for tUCA over other atherosclerotic biomarkers, such as CD81, ICAM-1, VCAM-1, P-selectin, platelet glycoprotein Ibα and von Willebrand factor, is the ability to identify intraplaque neovascularization, which has been identified as an independent predictor of intraplaque hemorrhage and plaque rupture. The αvβ3-tUCA could therefore be used to assess the risk of rupture of atherosclerotic plaques. Use of αvβ3-tUCA to assess adventitial vasa vasorum will have advantages over use of non-targeted microbubbles that are currently already used in the clinic for adventitial vasa vasorum assessment (Staub et al. 2010; van den Oord et al. 2015). These advantages include (i) a better assessment of adventitial vasa vasorum and intraplaque neovascularization as the plaque can be imaged in three dimensions by, for example, mechanical stepping of the transducer; and (ii) assessment of the far-wall plaque which is now very limited for non-targeted microbubbles because of the pseudo-enhancement artifact resulting from free-floating microbubbles in the lumen (ten Kate et al. 2012).
In our study, we chose the carotid artery vessels as the target for examination for the following reasons: First, it is believed that the carotid artery is the most reproducible site for studies of lesions in mice (Ni et al. 2009; Rigatelli and Zanchetta 2005). Second, the carotid artery vessels are superficially located and hence readily accessible for ultrasound imaging systems. Third, the salivary gland, which is in the field of view on top of the carotid, is a suitable internal positive control for binding of tUCA in such a murine model. Finally, the common carotid distal of the bifurcation is usually free from lesions and, hence, an ideal intra-animal negative control because it is in the same field of view.

We observed \( \alpha_\beta_3 \)-tUCA attached to the middle part of the common carotid artery, a location where no atherosclerotic lesions were present. This may be explained by \( \alpha_\beta_3 \)-integrin expression in normal arteries, as reported previously (Hoshiga et al. 1995). However, the intensity of \( \alpha_\beta_3 \) expression in the healthy part of the common carotid artery was significantly lower than that in the plaque ROI. As expected, there was no statistical significant difference between the molecular signal in the plaque ROI and that in the no-plaque ROI for the cUCA, as well as for tUCA versus cUCA in the part of the vessel without plaques. Subsequent work to further improve the signal-to-baseline would add to the power of this technique.

The salivary gland is a large and very well vascularized organ, which is located above the right carotid in our imaging field of view. Microvessel density in the gland is high, hence so is the associated \( \alpha_\beta_3 \)-integrin expression. Therefore, it is an ideal target as an internal positive control for successful binding of \( \alpha_\beta_3 \)-tUCA to its targets. Our immunocytochemistry analysis of the salivary glands also confirmed the presence of macrophages and microvessels in this gland.

Our quantification method ensures that only the signal from bound UCA is enhanced and considered as molecular signal. Shortly after injection of the UCA, the intensity of contrast in the ROI increased rapidly and reached a maximum, and this was followed by a slow decay in intensity for both tUCA and cUCA. However, 10 min after the injection, because of the binding of the tUCA to the \( \alpha_\beta_3 \)-integrin, the level of intensity is expected to be higher for tUCA than for cUCA. Ideally, 10 min after injection of cUCA, image intensity in the ROI should fall back to the baseline level (similar to the level before UCA injection), and there should be no difference in image intensity before and after the flash pulse. On the other hand, there will be a difference in the intensity levels for tUCA before and after the flash because of the destruction of bound tUCA, which is a measure of the molecular signals in the ROI. Even 10 min after injection, there will be recirculating unbound UCA for both tUCA and cUCA injections. Therefore, the actual difference in the intensity of the pre- and post-flash frames depends on the amount of both bound and unbound UCA in the ROI. The increase in intensity after the flash is also an indication of the presence of recirculating UCA. The processing steps, such as averaging pre- and post-flash frames and subtracting the result frames from each other, ensures
that tissue artifacts are minimized and only the stationary contrast spots are enhanced. We had previously found, in both in vitro and in vivo comparison studies, that the amplitude-modulation pulse sequence used to extract the non-linear response of the UCA at the fundamental frequency is a suitable non-linear contrast mode for imaging targets such as mouse carotid artery with the imaging platform used in this study (Daeichin et al. 2015b).

There are several limitations to this study. First, in the preparation of the targeted UCA, biotin–avidin bridging was used to non-covalently and very stably couple the $\alpha_b\beta_3$ antibody to the lipid microbubble coating. However, this strategy may not be translated to humans, as avidin is immunogenic in humans (Petronzelli et al. 2010). On the other hand, this avidin–biotin coupling can be avoided by covalently linking the antibody to the lipid microbubble coating. A covalent linkage of cyclic RGD, a human compatible ligand for $\alpha_v\beta_3$, has already been reported for tUCA (Anderson et al. 2011). In addition, these components are commercially available. Second, it cannot be assumed that the murine expression of $\alpha_v\beta_3$ is fully identical to clinical disease-related $\alpha_v\beta_3$ expression. In the clinical setting, more complex factors such as disease-related inducers, individual differences and intricate signal pathways are often involved in the regulation of $\alpha_v\beta_3$ expression. Although there remains a long road to translation of this technique from the animal model to the clinical situation, it would be desirable to relate the level of video-intensity signal from the tUCA to the stage of the disease and to predict the risk of future clinical cardiovascular events. Also, precise spatial correlation between ultrasonography and immunohistology is not always possible because of the difficulty involved in spatially matching the ultrasound plane with histologic sections and the fact that 2-D ultrasonography is formed from signals received over a beam elevation of almost a millimeter. Accuracy of $\alpha_v\beta_3$ quantification using the tUCA may be limited by some degree of non-specific binding because of the expression of $\alpha_v\beta_3$ by the vascular endothelium. In addition, more complex factors, such as acoustic radiation force, UCA concentration, blood flow velocity and UCA affinity, are involved and need to be further investigated.

CONCLUSIONS

This article describes the successful preparation of $\alpha_v\beta_3$-targeted microbubbles and the potential of target-specific UCA to quantitatively detect the expression of $\alpha_v\beta_3$ integrin in vitro and in vivo. Our initial experience in molecular ultrasonography using $\alpha_v\beta_3$-tUCA has indicated that it may enable in vitro and in vivo molecular imaging of $\alpha_v\beta_3$ expression on actively sprouting vascular endothelium. This imaging modality may provide reference values for relative expression of $\alpha_v\beta_3$ and information likely to be very useful to address the detection, prognosis and vulnerable potential of atherosclerosis or susceptibility to anti-atherosclerosis drugs.

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