Detection of hydroxyapatite in calcified cardiovascular tissues

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\textbf{A B S T R A C T}

\textit{Objective:} The objective of this study is to develop a method for selective detection of the calcific (hydroxyapatite) component in human aortic smooth muscle cells in vitro and in calcified cardiovascular tissues ex vivo. This method uses a novel optical molecular imaging contrast dye, Cy-HABP-19, to target calcified cells and tissues.

\textit{Methods:} A peptide that mimics the binding affinity of osteocalcin was used to label hydroxyapatite in vitro and ex vivo. Morphological changes in vascular smooth muscle cells were evaluated at an early stage of the mineralization process induced by extrinsic stimuli, osteogenic factors and a magnetic suspension cell culture. Hydroxyapatite components were detected in monolayers of these cells in the presence of osteogenic factors and a magnetic suspension environment.

\textit{Results:} Atherosclerotic plaque contains multiple components including lipidic, fibrotic, thrombotic, and calcific materials. Using optical imaging and the Cy-HABP-19 molecular imaging probe, we demonstrated that hydroxyapatite components could be selectively distinguished from various calcium salts in human aortic smooth muscle cells in vitro and in calcified cardiovascular tissues, carotid endarterectomy samples and aortic valves, ex vivo.

\textit{Conclusion:} Hydroxyapatite deposits in cardiovascular tissues were selectively detected in the early stage of the calcification process using our Cy-HABP-19 probe. This new probe makes it possible to study the earliest events associated with vascular hydroxyapatite deposition at the cellular and molecular levels. This target-selective molecular imaging probe approach holds high potential for revealing early pathological changes, leading to progression, regression, or stabilization of cardiovascular diseases.

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1. Introduction

Cardiovascular calcification results in the deposition of insoluble calcium in tissues such as arteries, heart valves, and cardiac muscle [1,2]. A rising from a systemic mineral imbalance, this physiological change is commonly observed in patients with cardiovascular disease (CVD) [3,4]. Although the mechanism of vascular calcification is not fully understood, it has been suggested that vascular wall resident cells such as pericytes or vascular smooth muscle cells (VSMCs) may transdifferentiate and produce a mineralized matrix [5–7]. Accumulating evidence supports the hypothesis that most atherosclerosis-associated calcification results from highly regulated and organized active cellular processes [1,2,8–10]. It has been proposed that circulating osteoblast precursors or resident mesenchymal stem cells could produce ectopic calcification. However, it is more likely that calcifying vascular cells (CVCs), identified as a subpopulation of VSMCs, induce pathological calcification in response to environmental stimuli [7,11]. CVCs secrete several osteogenic-associated proteins such as osteopontin, osteonectin, osteocalcin (OCN), bone sialoprotein (BSP), bone morphogenetic proteins (BMPs), and matrix Gla protein (MGP) into the extracellular matrix [11,12]. Many of these proteins have a high binding affinity for calcium salts and accumulate in areas of vascular calcification, where they may play various roles including the regulation of calcium crystal nucleation and growth. In normal arterial walls, VSMCs constitutively express potent inhibitors of calcification, such as MGP, whose absence results in spontaneous calcification [1]. Understanding how these proteins regulate calcification is crucial for designing novel therapeutic approaches to treat calcification associated cardiovascular diseases.

The calcium deposits in cardiovascular tissues are comprised of various calcium salts, such as calcium carbonate (CC), calcium oxalate (CO), calcium phosphate (CP), calcium pyrophosphate (CPP), and hydroxyapatite (HA). However, little information has been reported about these different forms of calcium salts in cardiovascular diseases. Even the most advanced clinical imaging modalities, such
as optical coherence tomography, MRI, and ultrasonography have difficulty distinguishing them. Without a reliable analytical method, a clear correlation between various calcium salts and calcified vascular tissues cannot be obtained. Recently, we have developed an HA binding peptide (HABP-19), which can potentially be used as a tool to distinguish HA from other calcium salts in biological tissues. HABP-19 is a 19-residue peptide, derived from the bone-binding protein, OCN. Maturation of OCN requires posttranslational modifications, specifically vitamin K-dependent γ-carboxylation, resulting in the formation of three γ-carboxylated glutamic acid (γ-Glu) residues from three glutamic acid (Glu) residues [13,14]. The γ-carboxylated OCN is primarily associated with bone formation through its high binding affinity to HA, although a fraction is released into the circulation [15,16]. In contrast, the under-carboxylated circulating form of OCN has been implicated as a novel hormone and positive regulator of glucose homeostasis [17,18].

When labeled with a near-infrared (NIR) fluorochrome, the HABP-19 probe was able to display HA in the skeletons of living animals for 6 weeks without significant signal reduction [19]. We hypothesized that this probe could distinguish HA from other calcium deposits in calcified cardiovascular tissues. In the present study, we have utilized fluorescently labeled HABP-19 to directly visualize the early-stage osteogenesis-like activity of VSMCs in vitro. We have also visualized HA calcification in CVD tissues, carotid endarterectomy tissues, and resected aortic valves ex vivo. This visualization of HA-dependent calcification is otherwise unidentifiable by conventional imaging modalities or routine histological methods.

2. Methods

Fluorescein isothiocyanate (FITC) labeled HABP-19, Cy5.5 labeled Cy-HABP-19, and the corresponding FITC-labeled control dye (Sigma–Aldrich) was also included as a negative control dye.

2.1. Binding kinetics of various calcium salts and HABPs

The binding affinities of FITC, cHABP, and HABP-19 to HA was evaluated by incubating 1 ml of 10 µM peptide solution with 10 mg HA (Ca₃(PO₄)₂·OH; Sigma, St. Louis, MO) at room temperature (RT) for 3 h with constant agitation. The amount of peptide in solution bound to HA was determined indirectly by measuring unbound peptide in solution (490 nm) using a SpectraMax M2 Microplate (Molecular Devices, Sunnyvale, CA) and comparing these values to the FITC standard in PBS buffer (pH = 7.2) at 490 nm (λ = 67,000). To further characterize the binding affinity of HABP-19 to calcium carbonate (CaCO₃; Sigma), calcium oxalate (Ca(C₂O₄)₂·H₂O; Fisher Scientific, Waltham, MA), calcium phosphate (Ca₃(PO₄)₂; Fisher Scientific), or calcium pyrophosphate (Ca₃P₂O₇; Sigma), HABP-19 was incubated separately with each calcium salt for 3 h at RT. The amount of HABP-19 bound to each calcium salt was determined indirectly by measuring unbound HABP-19 in solution as aforementioned.

2.2. Cell culture and induction of mineralization

Human aortic vascular smooth muscle cells (VSMCs; 1.5 × 10⁴ per cm²) (Genlantis, San Diego, CA) were grown in 35 mm² petri dishes supplemented with 2 ml of smooth muscle cell growth medium (SMCGM; Genlantis) in triplicate. Mineralization was induced by culturing the cells in SMCGM as the control medium, or SMCGM supplemented with 50 µg/ml ascorbic acid (AA), 7.5 mM β-glycerophosphate (β-GP), and 10 mM 1,2-lysophosphatidylcholine (LPC; Type I, 99%, Sigma) as the stimulating medium. In order to provide in vitro magnetic stimuli to VSMCs, cells grown to 80% confluence in T-175 flask was transferred to 35 mm² petri dishes and treated with Nanoshuttle-PL™ (Ns), a polylysine based hydrogel containing gold and magnetite nanoparticles (n3D Biosciences, Houston, TX), for 8 h. After the cells had taken up the Ns, they were magnetically levitated using a magnetic drive (n3D Biosciences) and cultured for 4 days in control or stimulating media (in vitro magnetic suspension). The first day of culture in stimulating media for the induction of mineralization was defined as day 0. The culture medium was replaced with fresh medium every 2 days.

2.3. Quantification of mineralized layers by Alizarin Red S (ARS) staining

Quantification of mineralized tissue formation was performed as previously described [20,21]. In brief, the magnetic drive was removed from the top of each dish to allow the cell suspensions in the 3D cultures to settle and spread at the bottom for 4 h. The resultant cell monolayer was washed with PBS, fixed in formalin, and incubated with ARS (pH 4.2). To quantify ARS staining, the monolayer was incubated in 400 µl of 10% (v/v) acetic acid, scraped from the plate, and transferred to a 1.5-ml tube. The mixture was overlaid with mineral oil, heated to exactly 85 °C, centrifuged, and 300 µl of the supernatant was transferred to a new 1.5-ml tube. 200 µl of 10% (v/v) ammonium hydroxide was added to neutralize the acid. The absorbance of the aliquots was measured in triplicate at 405 nm in a 96-well plate.

2.4. Fluorescence microscopy

Each cell monolayer was washed with PBS and stained with 2 nmol of FITC, cHABP, or HABP-19 at RT for 60 min. The monolayer was then washed extensively with PBS to remove unincorporated probes. Fluorescence images were captured using an IX51 microscope (Olympus, Center Valley, PA) with excitation at 488 nm (FITC-tagged LP). FV 1000 Viewer software (Olympus) was used for image analysis.

2.5. Tissue acquisition and storage

Carotid endarterectomy and aortic valve specimens were acquired within 1 h after surgical resection and stored until use in 50% glycerol/PBS (4 °C) to preserve tissue morphology. The use of the specimens was approved by the institutional review board (IRB) of Baylor College of Medicine (Houston, TX).

2.6. Histology

Human aortic valves were fixed in 10% formalin, dehydrated in a graded series of ethanol washes, and embedded in glycolmethacrylate. After polymerization, thin sections (5 µm)
were prepared using the Exakt System modified sawing microtome technique [22]. Serial sections were stained with a Von Kossa reagent (American MasterTech, Lodi, CA) or with molecular imaging probes (HABPs).

2.7. Optical imaging acquisition

For ex vivo optical imaging, each specimen was incubated with 2 nmol of FITC, chABP, HABP-19, or Cy-HABP-19 for 1 h with constant agitation, thoroughly washed with PBS to remove unbound probe, resuspended into PBS solution, and images were acquired using the Maestro 2 optical imaging system (CRI, Woburn, MA). To validate the selective accumulation of imaging probes, a red (CY5.5) filter set (acquisition setting: 650–800 nm) steps and 10 ms exposure time) was used for FITC, chABP, and HABP-19. A red (CY5.5) filter set (acquisition setting: 680–950 nm steps and 10 ms exposure time) was utilized for Cy-HABP-19. The fluorescence images obtained were corrected to remove the auto-fluorescence background using the multi-excitation spectral analysis function (Maestro software v. 2.10).

2.8. Micro computed tomography (μCT) analysis

Micro CT was performed using a Siemens Inveon Preclinical Multimodel PET/SPECT/CT system (Malvern, PA) at medium resolution. Real-time images were reconstructed for correlation with the optical imaging system. An internal infrared video camera allows visual sample monitoring during scan acquisition. The scanner was operated in the 3D volume imaging acquisition mode. Specimens were laser aligned at the center of the field of view of the scanner for subsequent imaging. The μCT image was acquired in approximately 3 min, and concurrent image reconstruction was achieved using a COBRA (Siemens) Invenon Research Workplace software (Siemens) was used to view and adjust imaging.

2.9. Statistical analysis

In vitro and ex vivo data were analyzed for statistical significance by the Student’s t-test or one way ANOVA and Dunnett post hoc test, using the Statistical Package for the Social Sciences (SPSS) software, version 13 (SPSS, Chicago, IL). Means, standard deviations, and degrees of significance are shown on individual data graphs in the ‘Results’ section. A probability value (P) of <0.05 was considered statistically significant unless otherwise indicated.

3. Results

3.1. The HABP-19 probe specifically recognizes HA salt

The HA specificity of the prepared probes was determined by incubating HA with FITC, chABP, and HABP-19 solutions for 3 h. The HA binding affinity of each probe was quantified by measuring the absorbance of solution (490 nm) (Fig. 1). HABP-19, which has 6 Gla residues, exhibited a rather high HA binding affinity (86.2 ± 4.1%), whereas neither chABP (3.3 ± 3.2%) nor control FITC (4.2 ± 2.2%) showed appreciable HA binding (P < 0.005), suggesting a crucial role for the Gla residues within HABP-19 for binding to HA, as previously reported [19–21]. To confirm the HA binding specificity of HABP-19, a panel of calcium salts, CC, CO, CP, CPP and HA, were incubated in HABP-19 solutions. HABP-19 exhibited exceptional binding affinity for HA but not for the other calcium salts (P < 0.005). Our data indicate that HABP-19 has 25-fold higher binding affinity for HA than does chABP, and >6-fold higher specificity for HA than the other calcium salts tested (Fig. 1).

3.2. Induction of calcification in human aortic smooth muscle cells

To study the early stages of the mineralization process in VSMCs, cells were cultured in control media (CM) or stimulating media (SM) supplemented with AA, β-GP, and LPC. AA and β-GP are known osteogenic factors that stimulate osteoblastic differentiation and mineralization [23–25]. LPC, a product of oxidized phosphatidylcholine hydrolysis, is a candidate compound thought to act on VSMCs and stimulate their transition into calcifying vascular cells, and it likely plays a significant role in the calcification of atherosclerotic plaques [7]. In a normal 2D culture, VSMCs cultured in SM grew into a dense monolayer in 3 days, whereas VSMCs grown in CM were smaller and less organized (Fig. 2A and Fig. 1 in the online-only Data Supplement). Cells treated with Nanoshuttle (Ns), without applying the magnetic field, exhibited elongated spindle-like cell shapes and formed nodules with pronounced dark ridges, reflecting the presence of iron nanoparticles (Fig. 2B). However, Ns treated and magnetically levitated VSMCs were capable of forming large cell clusters in the central region beneath the circular magnet drive at day 3 (Fig. 2C). Diffusely scattered spherical clusters of cells were also observed.

To validate the difference in mineralization, the treated cells were stained with ARS and quantified by absorbance spectroscopy (Fig. 2D and E). Cells cultured in 3D-Ns SM promoted the most mineral deposition. ARS measurements showed that the enhancement was 5-fold higher than it in normal 2D cultures (P = 0.0204) (Fig. 2E). However, the effect of CM and SM in all three study groups was not statistically significant. Hence the effects of extrinsic osteogenic factors didn’t influence the biological activity of VSMCs much in these early time points.

3.3. Identifying HA calcification in human aortic smooth muscle cells by HABP-19

Based on the above ARS mineralization results, 3D-Ns cultured VSMCs cells were selected for HABP-19 binding studies. Four days after stimulation, HA deposition was monitored after staining with...
HABP-19 and controls. As expected, a large fluorescence signal was seen in the HABP-19 stained cells, whereas minimal signal was observed in either the cHABP or FITC stained cells (Fig. 3). Other culture conditions (2D and 2D-Ns cultures) did not result in the deposition of a significant amount of HA (Figs. 2 and 3 in the online-only Data Supplement).

3.4. Detecting HA deposition in excised calcified human aortic valves

The potential detection of HA in biological tissues was then tested with human specimens. Calcified human aortic valves resected during valve replacement procedures were sliced into four pieces. The degree of calcification of these pieces was confirmed by μCT which is routinely used to visualize bones and calcified tissues [26]. The obtained 2D and 3D CT images indicate various degrees of calcification in these specimens (Fig. 4C and Movie I in the online-only Data Supplement). A characteristic pattern of mineralization was observed. Calcium depositions were mostly oriented toward the center cusp area of the aortic valves. Although the CT images revealed the area of calcification, they did not allow for differentiation of HA from other calcium components. The bright field image and fluorescence image of an aortic valve before incubation with probes are shown in Fig. 4A and B, respectively. The selective HA binding affinity of HABP-19 was then investigated by incubating the sliced specimens in PBS, FITC, cHABP, or HABP-19 for 1 h and removing the unbound probes (Fig. 4D–F). Fluorescence imaging showed a bright HABP-19-dependent signal. Minimal signal was observed with cHABP. Although a non-specific background signal was initially observed with FITC, it quickly decreased with time. Nevertheless, a strong HABP-19 signal was sustained throughout. Optical molecular images of calcified aortic valves treated with the HABP-19 probe exhibited intense HA accumulation at calcified regions, as indicated by an arrowhead (Fig. 4C and F).

Fig. 2. Morphological changes and mineralization of VSMCs. Cells were incubated in control media (CM), or in stimulating media (SM) supplemented with 50 μg/ml ascorbic acid (AA), 7.5 mM β-glycerophosphate (β-GP), and 10 nM L-α-lyso-phosphatidylcholine (LPC) for 4 days. Morphology of VSMCs in 2D (A), 2D-Ns (B), or 3D-Ns (C) SM cultures. Cells were washed with PBS and observed under light microscopy. Images of cells cultured in CM are shown in Supplemental Fig. 2. (D) Cells cultured in CM or SM were stained with Alizarin Red S (ARS) to detect the calcium mineral layer and images captured by a digital camera after the removal of unbound ARS dye by washing with water. (E) Mineral layers stained with ARS in CM or SM were quantified at 405 nm (results are mean ± SD, n = 4).

Fig. 3. Images of HA in VSMCs. HA calcification of VSMCs was induced by osteogenic factors and a 3D environment. For corroborative visualization of the extent of HA calcification, cells cultured in SM for 4 days were stained with FITC, cHABP, or HABP-19 and observed by fluorescent microscopy. Bright field (top) and fluorescence images (bottom) from mineral layers of VSMCs. HA deposits of VSMCs induced by AA, β-GP, LPC, and 3D-Ns were stained with control (A), FITC (B), cHABP (C) and HABP-19 (D) probes.
3.5. Histological analysis of calcified human aortic valves

Histological analysis of the calcified plaque was performed to determine HA distribution at the tissue level. It is known that calcium salts in paraffin-embedded tissues can be lost during the sectioning process. Thus the aortic valve tissue here was embedded in plastic to avoid this complication (Fig. 5). Consecutively sectioned aortic valves (Fig. 5A) were stained with Von Kossa reagent commonly used to visualize calcium salts in histological tissues (Fig. 5B); molecular imaging probes, including FITC dye, cHABP, and HABP-19 (Fig. 5C). Dark Von Kossa-dependent staining exhibited dense calcification in a band-like pattern (Fig. 5B).

However, staining of consecutive sections using HABP-19 revealed a more restricted pattern of HA deposition within the medial part (delineated by red lines in Fig. 5C). Aortic valves stained with FITC dye or cHABP did not exhibit detectable fluorescence. The HABP-19 and Von Kossa staining patterns were different in terms of their binding selectivity. The Von Kossa reagent stained all calcium salts, regardless of their phosphate content, while HABP-19 stain was selective for HA deposits.

3.6. Visualization of HA deposition in calcified cardiovascular specimens by the NIR HABP-19 probe

HA deposition in calcified cardiovascular tissues was further visualized using the NIR dye labeled Cy-HABP-19 imaging probe. This probe has increased sensitivity for real-time imaging and may
provide non-invasive optical detection of HA deposition in cardiovascular tissues [27]. Surgically excised human carotid atheroma specimens and aortic valves were incubated with Cy-HABP-19 for 1 h, and unbound probes were removed by washing with water. Images of calcified cardiovascular tissues were obtained and analyzed by an optical imaging system and μCT (Fig. 6). Digital images of carotid atheroma specimens (Fig. 6A) and aortic valves (Fig. 6B) clearly show an intense Cy-HABP-19-dependent signal within the calcified regions, primarily within tissue areas surrounding the bifurcation of carotid tissues and within the cusp area of aortic valves. μCT images showed large areas of calcium deposits (white) on the tissues (Fig. 6C and D), whereas NIR optical images facilitated distinct detection of HA on the calcified cardiovascular tissues (Fig. 6E and F). Background fluorescence signals induced by non-specific binding or auto-fluorescence in the aortic valves were not detectable in the NIR image; the tissue is therefore outlined with a dotted line (Fig. 6E). Although Cy-HABP-19 deposits matched the calcified regions seen on the CT images, there was a disparity between two imaging modalities with respect to selectivity. Cy-HABP-19 with NIR optical imaging was able to differentiate HA calcification from other components.

4. Discussion

There is considerable interest in furthering our understanding of the mechanisms of cardiovascular calcification and its implications in CVD [3,4]. Calcium salts of different forms are generated by locally produced stimuli; however, little is known regarding the correlation between the type and composition of the calcium salt, and tissue changes. In calcified vascular tissues, two distinct calcium compounds have been identified based on their chemical composition, CO (CaC$_2$O$_4$·2H$_2$O) and HA (Ca$_{10}$(PO$_4$)$_6$·6H$_2$O). It has been reported that calcium deposits seen in arteries and cartilage are mainly composed of HA, as seen in bone [1]. In this study, we describe a specific and sensitive tool to study HA, one of the main types of biological calcium deposits.

Previously, we developed fluorescence conjugated HA binding peptides (HABPs) inspired by the protein OCN; these peptides mimic natural OCN-HA binding properties by forming α-helices with HA in the skeleton of live animals [19]. Because bone mineralization and vascular calcification share a similar mechanism, our prior findings could be applied to detect HA calcification associated with cardiovascular diseases. We hypothesized that a HABP-19 probe could selectively detect HA calcification and may help predict potential risks for clinical complications in patients with CVD. The HA binding specificity of the HABP-19 probe was demonstrated using pure calcium salts, stimulated cell culture, and surgically excised specimens. Upon incubation with different types of salts, HABP-19 showed selective HA binding compared to chABP and FITC (negative control dye, Fig. 1). Substitution of the critical Gla residues with Glu amino acids in chABP resulted in a complete loss of its HA binding capability, indicating that the Gla residues within the HABP-19 probe are responsible for HA binding specificity [16,19–21,28]. The HA-binding selectivity was further tested by incubating HABP-19 with different forms of calcium salts. The fluorescent signal resulting from incubation of HABP-19 with HA salt was significantly higher than that resulting from all other salts, confirming the binding specificity of HABP-19 for HA.

It has been shown that VSMCs grown in the presence of osteogenic factors, such as AA, β-GP, and LPC, exhibit extensive matrix mineralization with minerals such as HA, calcifying collagen, extracellular matrix vesicles, and nodular calcification, similar to that observed in calcified atherosclerotic plaques and heart valves in vivo [29]. Thus, we evaluated whether HA deposition is present in the matrix monolayer during the early stage of VSMC mineralization in vitro. VSMCs were cultured with extrinsic stimuli, including osteogenic inducible factors, and in a 3D microenvironment. ARS analysis illustrated that mineral formation in VSMCs cultures was more dependent upon the 3D environment than on extrinsic factors (Fig. 2D). Following incubation with the fluorescent HABP-19 molecular imaging probe, fluorescence microscopy revealed that HA deposits were found on the mineral layer of both magnetically levitated 3D (Ns) VSMCs and VSMCs cultured in the presence of osteogenic factors under fluorescence microscopy. Neither 3D culture nor osteogenic factors alone promoted detectable HA calcification in vitro (Figs. 2 and 3 in the online-only Data Supplement). As expected, other control probes were unable to bind HA. Our data suggest that 3D stimuli were much more effective in inducing HA calcification of VSMCs than were extrinsic osteogenic factors, and importantly, that HABP-19 is capable of detecting newly formed HA.

![Image](https://example.com/image.png)

**Fig. 6.** Ex vivo images of cardiovascular tissues by NIR fluorescence. CEA tissues (A) and aortic valve (B) were incubated with Cy-HABP-19 for 1 h. Digital color images reveal extensive calcification (light cyan) in CEA specimens (A) and aortic valve (B), and μCT images. The white color represents calcified areas on the tissues. (E) and (F) NIR optical fluorescence images. These images are delineated as unmixed fluorescence-enhanced signal images (white) superimposed on the pseudo-background color image (gray). The control sample (marked by a dotted line and open arrowhead in panel C) did not exhibit any noticeable fluorescence in the optical imaging system.
Ectopic calcification has a detrimental effect on vascular function and is associated with increased mortality in cardiovascular diseases, type 2 diabetes, and chronic kidney disease [30]. The majority of ruptured plaques contain HA deposits [2,8–10]; therefore, HA deposition might be a good indicator of plaque vulnerability. Although early diagnosis is crucial for identifying vascular plaques induced by ectopic calcification before symptoms develop, there is no stable, selective, and sensitive indicator to detect plaques associated with HA, or to distinguish HA from other calcium salts without the possibility of negative feedback. Therefore, we examined the potential of HABP-19 to distinguish HA from calcium salts without the possibility of negative feedback. There-fore, we first surveyed by μCT. These μCT images confirmed that these patient tissues were highly calcified. In contrast to the large calcified areas identified by μCT, only small regions among the μCT highlighted areas was highlighted by HABP-19, presumably the HA rich regions. These differences suggest that not all calcium deposits on the aortic valves have the same composition. Previously, Ortlepp et al. have determined HA composition from the excised aortic valves [31]. Calcium contents were initially determined by atomic absorption analysis, and, according to the X-ray diffraction (XRD) measurements, the calcification in the valves was concluded to be HA deficient: CaO·(PO₄)₂(PO₃)₃(OH)₂·αH₂O, with 0 < α < 1, suggesting that HA was mixed with other calcified salts. Since the detail chemical information of calcium deposits in tissues is largely under-studied, this HABP-19 imaging probe might be useful in distinguishing HA rich depositions from other mineral components. Detail composition of the calcified atherosclerotic plaques could be analyzed by various SEM/XRD spectroscopies [32–34].

We further conducted histological analyses to identify calcium salts and their distribution in the sectioned tissues by comparing Von Kossa staining with our molecular imaging probe. Although routine histological methods are useful for visualizing calcification of cardiovascular tissues, the sectioning process itself often results in a loss of calcium crystals. Thus, a plastic embedding process was introduced in the present study to minimize calcium salt loss during sectioning. Calcification staining results showed that the Von Kossa assay stained entire mineralized layers and lacked adequate specificity. Conversely, fluorescence images using our HABP-19 probe clearly indicated a restricted stained area, mainly corresponding to HA deposits that were not selectively detected by Von Kossa staining (Fig. 5).

For in vivo HA imaging, NIR fluorescence (NIRF) molecular imaging offers several advantages, including relatively deep photon penetration into tissue and reduced auto-fluorescence [27,35]. NIR conjugated Cy-HABP-19 optical images exhibited distinct HA deposition at the surface of calcified cardiovascular tissues compared to μCT images or fluorescence imaging acquired with visible light (Fig. 4 vs. Fig. 6). Therefore, Cy-HABP-19 could potentially be used as a unique molecular imaging probe offering previously unavailable information on HA deposition, and detecting early stage cardiovascular disease. The distinct HA selectivity of this probe may be especially useful in studies of older individuals, in which HA deposition may have occurred, but the calcification being imaged is the result of deposition of non-HA calcium salts [27]. A bisphosphonate-based imaging contrast agent has previously been developed to image osteogenic activity in atherosclerotic plaques in mice models [27,36,37]. Our results indicate that HABP-19 could be applied to detect HA deposition in human tissues and, potentially, within patients, thereby offering an new diagnostic tool to assess clinical risk and monitor the efficacy of various interventions [37,38]. However, the optical technology is not suitable for whole body imaging. Other imaging reporters, such as PET, SPECT, or MRI tracers, could be tethered to Cy-HABP-19 for whole-body multi-functional imaging.

Although the contribution of atherosclerotic calcification to plaque rupture is undefined, previous studies have revealed that inflammation precedes calcification in early-stage atherosclerosis, suggesting that macrophages promote the proinflammatory environment and send specific signals to vascular wall cells to initiate osteogenic differentiation [37,39]. Subsequently, both inflammation and calcification processes developed in parallel and within close proximity. The microcalcification stage may cause plaque rupture and microfractures that may result in the acute clinical events [40]. Once equilibrium in the artery shifts toward calcification, HA deposition could quickly advance and drive disease progression [37]. Virmani et al. have demonstrated that a specific type of lesion with a calcified nodule is prone to rupture; these lesions consist of a fibrous cap disrupted by dense calcific nodules [41–43]. They hypothesize that these plaques may rupture owing to physical stress exerted by the nodules. On the other hand, other evidence indicates that calcified plaque may be more stable and, thus, less prone to rupture than noncalcified lesions [44]. In summary, the contribution of calcification to plaque vulnerability remains controversial and further investigation is required. The developed HABP-19 could represent a useful tool for further investigation of the mechanisms of vascular calcification, in particular HA formation.

Conflict of interests

None.

Acknowledgments

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Appendix A. Supplementary material

Supplementary data related to this article can be found online at http://dx.doi.org/10.1016/j.atherosclerosis.2012.07.023.

References
