Introduction

Atherosclerosis is the main cause of coronary artery disease (CAD). Atherosclerosis is developed and promoted by mainly the influx of LDL particles and the production of inflammatory cytokines in the arterial wall (1). These atherogenic mechanisms build non-obstacle atherosclerotic plaques in its early stage. Majority of plaques remain quiescent, whereas some plaques progress and others suddenly cause its rupture, leading to the acute coronary events (2). Therapeutic modulation of atherogenic targets is a key for the prevention of CAD. Numerous large-scale clinical trials have demonstrated anti-atherosclerotic effects of 3-hydroxy-3-methylglutaryl coenzyme A reductase inhibitors or statins for the primary and secondary prevention (3-6). Statin has the ability to reduce atherosclerotic cardiovascular events and attenuate disease progression. In addition, high-intensity statin has the potential to regress coronary atheroma in patients with CAD (5,7,8). Despite these favorable effects of a statin, many patients still continue to develop cardiovascular events (9,10). This ongoing disease risk presents a considerable challenge suggesting the need to identify additional key therapeutic targets, and develop more effective forms of risk prediction and preventive therapies.

Coronary angiography (CAG) has been widely used for the diagnosis and quantification of obstructive stenosis in the clinical settings. CAG has enabled to characterize culprit plaques associated with CAD, assess acute and chronic vascular response after revascularization strategies, and investigate the effect of medical therapies. While CAG is still a gold standard for the evaluation of coronary
artery stenosis, several limitations of CAG are recognized. CAG generates a 2-D silhouette of the arterial lumen and does not image the vessel wall, the site at which plaque accumulates. This disadvantage of CAG has stimulated the need to develop imaging modalities which enable to directly visualize plaques in the vessel wall.

Recent technological advances permit direct visualization of the full burden of coronary atherosclerosis. Intravascular imaging is increasingly becoming an important role in the evaluation of revascularization procedure but also in the assessment of drug efficacy. In addition, intravascular imaging has contributed to the identification of high-risk plaques that are likely to cause future cardiovascular events, so called “vulnerable plaques” (11,12). In this article, we summarize currently available intravascular imaging modalities to elucidate pathophysiology and characteristics of coronary atheroma in vivo.

**Grayscale intravascular ultrasound (IVUS)**

Grayscale IVUS uses miniaturized crystals which generate high-resolution, cross-sectional images of the vessel wall and lumen. Axial resolution is approximately 150 μm and the lateral resolution 300 μm. Grayscale IVUS allows robust quantitative measurements including lumen, vessel, and plaque area (13). Furthermore, Grayscale IVUS imaging enables to assess procedural outcomes after percutaneous coronary intervention (PCI). PCI with the guidance of grayscale IVUS has been demonstrated to reduce incidence of restenosis, stent thrombosis and cardiovascular events after PCI compared to angiographic guidance (14-17). Recently, IVUS has been increasingly used as the gold standard to evaluate natural history of coronary atherosclerosis under medical therapies (3,5,18-20).

**Plaque phenotypes on grayscale IVUS**

Based on echogenicity, plaques are classified in four categories by grayscale IVUS: soft plaque showing lesion echogenicity less than the surrounding adventitia, fibrous plaque with intermediate echogenicity between soft plaque and highly echogenic calcified plaques, calcified plaque having higher echogenicity than the adventitia with acoustic shadow, and mixed plaque (21). In addition to these grayscale IVUS-derived classifications of atherosclerotic plaques, distinct plaque phenotypes have been reported as follows.

**Attenuated plaque**

Attenuated plaque is defined as hypoechoic plaque with deep ultrasound attenuation without calcification or very dense fibrous plaque (Figure 1A). Histologically, attenuated plaques on IVUS correlate with a fibroatheroma containing large necrotic core or pathological intimal thickening (PIT) with a large lipid pool (22). Attenuated plaque is frequently observed at culprit lesion in patients with acute coronary syndrome (ACS) and is associated with deterioration of coronary flow and periprocedural myocardial infarction following PCI (23-25).

**Echolucent plaque**

Echolucent plaque is a lesion containing an intraplaque zone of absent or low echogenicity (Figure 1B) (26-28). Echolucent plaque is generally the result of high lipid content in a fibroatheroma, but relatively smaller size of necrotic core or lipid pool compared with attenuated plaque (22,29). This plaque phenotype has been shown as a signature of culprit lesions in patients with acute myocardial infarction (30).
Calcified nodule
Calcified nodule is a plaque phenotype associated with acute coronary events (31). Histologically, calcified nodule consists of fibrocalcific plaque with little or no underlying necrotic core, luminal surface that is disrupted by nodules of dense calcium, and overlying thrombus (32). Recent studies have validated features of calcified nodule on grayscale IVUS by comparing to pathological data in autopsy (33). Histopathological analyses have shown a distinct calcification with an irregular, protruding and convex luminal surface as grayscale-IVUS derived calcified nodule (Figure 1C). While this plaque feature is considered to associate with ACS, a sub-analysis of the PROSPECT study reported that patients with calcified nodule were less likely to develop major adverse cardiovascular events (34). Further investigation is warranted to elucidate clinical significance of calcified nodule on IVUS.

Spotty calcification
While extensive calcified lesions are generally considered to be clinically quiescent, scattered calcium pattern (spotty calcification) is associated with acute coronary events. Pathohistological study has shown the presence of small amount of calcium at high-risk lesions causing ACS or sudden cardiac death (35,36). Ehara et al. has demonstrated the presence of spotty calcification at culprit lesions in patients with acute coronary events in vivo (37). The definition of spotty calcification on IVUS is a lesion harboring only small calcium deposits with an arc of calcification of <90 (Figure 1D). Serial grayscale IVUS imaging has elucidated an accelerated plaque progression in patients with spotty calcification (38). As such, this calcium pattern on grayscale IVUS seems to be quite active and progressive form of disease.

Multiple layer appearance
Matsuo et al. has reported multiple layer appearance on grayscale IVUS as a high-risk feature of disease in heart transplant recipients (39). This phenotype exhibits multiple layers with distinct echogenicity (Figure 1E). In a serial grayscale IVUS study analyzing 132 heart transplant recipients, multiple layer appearance was associated with plaque progression. This might indicate the contribution of repeated episode of mural thrombosis to disease progression in heart transplant recipients.

Limitations of grayscale IVUS
Grayscale IVUS is an invasive procedure and is therefore currently limited to the assessment of patients who require CAG for a clinical indication. Conventional IVUS provides a suboptimal characterization of the composition of atherosclerotic plaque.

Virtual histology intravascular ultrasound (VH-IVUS)
While grayscale IVUS has the limited ability to analyze components within atherosclerotic lesions, VH-IVUS enables a detailed analysis of plaque composition. An advanced radiofrequency (RF) analysis of reflected ultrasound signals in a frequency domain analysis is used to visualize a reconstructed color-coded tissue map of plaque composition including fibrous, fibrofatty, necrotic core and dense calcium. The VH-IVUS algorithm has been validated against histology in autopsy specimens and in patients who underwent PCI. A high accuracy in detecting fibrous, fibrofatty, necrotic cores and calcium (autopsy: 79.7%, 81.2%, 85.5%, 92.8%, patients treated with PCI: 87.1%, 87.1%, 88.3%, 96.5%) has been already reported (40,41).

Plaque classifications by VH-IVUS
Currently, VH-IVUS classifies plaques into the following types.

Pathological intimal thickening (PIT)
PIT is considered as a progressive lesion in the early stages of atherosclerosis and represents a precursor lesion to fibroatheroma (31,42). It consists of mainly a mixture of fibrotic tissue and fibrofatty plaque with less than 10% each confluent necrotic core and dense calcium, respectively (Figure 2A) (44).

Thin-cap fibroatheroma (TCFA)
TCFA has been considered as a high-risk lesion causing acute coronary events (36,45). Histologically, TCFA is characterized by thin cap overlying a large necrotic core containing numerous cholesterol clefts (31). On VH-IVUS imaging, TCFA is characterized as a lesion having more than 10% confluent necrotic core with above 30 degree necrotic core abutting the lumen in at least three consecutive frames (Figure 2B) (44).

Thick-cap fibroatheroma (ThCFA)
ThCFA harbors a large lipid-necrotic core comprising large amount of extracellular lipid, cholesterol crystals, and necrotic
debris, surrounded by a thick fibrous cap (44). The definition of ThCFA is a lesion with more than 10% confluent necrotic core and a definable fibrous cap (Figure 2C).

**Fibrotic plaque**

VH-IVUS derived fibrotic plaque is mainly fibrous tissue without confluent necrotic core or calcium (Figure 2D) (44). Fibrotic plaque is different from PIT with regard to the presence of fibro-fatty tissue. Fibrotic plaque contains few fibro-fatty tissue (<15%), whereas PIT contains >15% fibro-fatty tissue.

**Fibrocalcific plaque**

Fibrocalcific plaque is a collagen-rich lesion with nearly all fibrotic tissue and dense calcium with less than 10% confluent necrotic core (Figure 2E) (44).

**Limitations of VH-IVUS**

Identification of intraluminal organizing thrombus is currently not possible by RF analysis. The acquisition of VH-IVUS images is gated at the R wave of the electrocardiography signal, which fails to allow to evaluate corresponding images in serial VH-IVUS imaging. Recent study raises questions about the accuracy of VH-IVUS to identify necrotic core. VH-IVUS-identified necrotic core was not correlated to histology in a swine atherosclerosis model (46). However, it is important to note that swine necrotic core is inherently different from human ones.

**iMAP-IVUS**

The iMAP-IVUS also acquires RF data continuously and then classifies plaques into four categories based on a spectra pattern; fibrous, fibrofatty, calcium, and necrotic core. Ex vivo validation study demonstrated accuracies for the detection of fibrous, lipidic, calcium and necrotic core as 95%, 98%, 98% and 97%, respectively (48). However, role and significance of iMAP imaging has not been fully investigated and established yet.

**Optical coherence tomography (OCT)**

OCT is a novel intravascular imaging modality using near-infrared light (1,300 nm). Compared to IVUS, OCT...
provides images of in vivo plaques with approximately 10 times higher resolution which is up to 10 μm in an axial resolution and to 20 μm in a lateral resolution. Since OCT has shallow penetration depth of imaging, this modality is superior to monitor plaque microstructures below the endothelial surface.

**Plaque composition on OCT imaging**

**Fibrous plaque**
A fibrous plaque exhibits high backscattering and a relatively homogeneous OCT signals (Figure 4A).

**Lipid plaques**
Lipid plaque is a signal-poor region with poorly delineated borders, a fast OCT signal drop-off, and little or no OCT signal backscattering (Figure 4B).

**Calcification plaques**
Calcification appears as a signal-poor or heterogeneous region with a sharply delineated border (Figure 4C).

**Plaque microstructures**

**Thin fibrous cap**
Fibrous cap on OCT is visualized as a signal-rich layer from the coronary artery lumen to the inner border of the underlying lipid-rich plaque. Thin fibrous cap is generally defined as fibrous cap <65 μm. A good correlation of the fibrous cap thickness between OCT image and histology has been reported (r=0.9, P<0.001) (49).

**Thin-cap fibroatheroma (TCFA)**
TCFA is an important feature of vulnerable plaque. While IVUS does not have enough abilities to visualize TCFA due to its limited spatial resolution, the high resolution of OCT enables to identify TCFA in vivo. OCT derived TCFA is characterized as the presence of thin fibrous cap (<65 μm) overlying a signal-poor lesion with diffuse border (lipid arc >90°) (Figure 4D).

**Macrophage**
Heavily infiltrated macrophage in thin fibrous cap is considered as a signature of vulnerable plaque (50).
visualizes macrophage accumulation as linear high-intensity signal on plaque surfaces with high attenuation (Figure 4E).

**Microchannel**
Microchannel is non-signal tubuloluminal structures without a connection to the vessel lumen and can usually be followed in multiple contiguous frames (Figure 4F). The presence of microchannel on OCT imaging has been shown to associate with a higher incidence of TCFA and future plaque progression (51,52).

**Cholesterol crystal**
Cholesterol crystal is defined as thin, linear region of high signal intensity within the lipid plaques (Figure 4G). The presence of cholesterol crystal is associated with enhanced plaque instability in patients with CAD (53).

**In-stent neoatherosclerosis**
In-stent neoatherosclerosis is newly formed atherosclerotic form within the neointimal tissue of stented segments. Emerging evidence suggests chronic inflammation and/or incompetent endothelium as important contributors to this pathological change within stent (54,55). Recent studies demonstrated that neoatherosclerosis could be developed in late phase after both bare metal stent (BMS) and drug-eluting stent (DES) implantation (56,57). In addition, earlier and more frequent development of neoatherosclerosis after DES implantation has been observed in pathohistological and clinical studies (58,59). Currently, neoatherosclerosis is considered as one of important mechanisms associated with late stent thrombosis and restenosis (54,57,60,61). OCT has a great potential to evaluate in-stent neoatherosclerosis in vivo. In-stent neoatherosclerosis by OCT imaging has been defined as the combination of neointimal diffuse thickening with intimal lipid-laden (poor signal region with diffuse borders) and the presence of fibrous cap (57,61,62). In a recent study analyzing in-stent restenosis lesions after DES implantation by OCT, TCFA and neointimal rupture within stent were observed in 52% and 58% of study.

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**Figure 4** Images of plaque phenotype on OCT: (A) Fibrous plaque: fibrous plaque (asterisk) is imaged as a high backscattering and homogeneous region; (B) lipid-rich plaque: lipid-rich plaque (asterisk) is imaged as a signal-poor region with a poorly delineated borders; (C) fibrocalcific plaque: calcification (arrow head) is imaged by OCT from 12 to 3 o’clock, as well-delineated, signal-poor region; (D) fibrous cap: a fibrous cap (double arrow) is imaged as a signal-rich homogeneous region overlying a lipid core; (E) macrophage: macrophages (arrow heads) are imaged as signal-rich linear regions accompanied by high attenuation; (F) microchannel: microchannels (arrow heads) are imaged as signal poor voids observed in multiple contiguous frames; (G) cholesterol crystal: cholesterol crystal (arrows) is imaged by OCT as linear, high-backscattering structures within the plaque. OCT, optical coherence tomography.
Limitations of OCT

Limitations of OCT are poor tissue penetration and interference from blood. OCT has a penetration depth of 2 to 3 mm, which prohibits imaging beyond the internal elastic lamina. Due to this limited penetration, it is difficult to evaluate the entire amount of plaques. This imaging is not suitable for visualizing atherosclerotic plaques in large arteries. Infusion of contrast medium is required during OCT imaging because infrared light does not penetrate red blood cells. This feature makes it difficult to use this imaging in patients with kidney disease.

Near-infrared spectroscopy (NIRS)

Spectroscopy is the measurement of the electromagnetic spectrum through interaction of light with molecules. Based on this method, NIRS imaging within coronary artery allows for the chemical characterization of biological tissues and can be used to assess lipid and protein content in atherosclerotic plaques. In an experimental animal study and ex vivo validation studies, this modality was shown to accurately detect the lipid content within atherosclerotic plaques (63-65). The sensitivity and specificity to detect lipid component was 90% and 93%, respectively. Recently, catheter-based NIRS imaging system has been developed (InfraReDx, Burlington, Massachusetts, USA). This device incorporates the rotating core with optical fibers which emit and absorb near-infrared light. NIR spectrometer delivers the light into a sample and then evaluates the proportion of light which is returned over the range of optical wavelength.

Identification of high-risk plaques on NIRS

One recent study has investigated plaque features at culprit and non-culprit segments in patients with ST elevation myocardial infarction and autopsy segments without any lipid plaques (66). Culprit segments were more likely to exhibit a larger lipid core burden index (LCBI) compared to non-culprit segments (median LCBI; 523 vs. 90, P<0.001) and coronary autopsy lipid-free segments (median LCBI; 523 vs. 6, P<0.001). In addition, LCBI above 400 was an optimal cut-off to distinguish culprit segments in patients with ST elevation myocardial infarction from coronary specimens free of lipid component. Another study elucidated better evaluation of plaque characterization under the combination of NIRS and grayscale IVUS (29). In this study analyzing 131 lesions, attenuated (P<0.0001) and echolucent plaques (P=0.008) on grayscale IVUS were associated with a larger LCBI compared to calcified and non-calcified plaques. Another study demonstrated the utility of adding NIRS to conventional IVUS for the detection of fibroatheroma in the 116 coronary arteries from autopsy hearts (67). There were significant trends of increasing plaque burden and LCBI across more complex atheroma. Furthermore, adding NIRS derived-LCBI to the grayscale IVUS-derived plaque burden was significantly improved fibroatheroma detection accuracy. NIRS also has
the ability to predict periprocedural myocardial infarction following PCI (68). Goldstein et al. has reported that lesions with LCBI >500 were more likely to experience periprocedural myocardial infarction (P=0.0002) (69). These observations suggest the potential of NIRS imaging for the risk stratification of future coronary events and the management of revascularization procedure.

Limitations of NIRS imaging

NIRS is not able to evaluate the depth of lipid core and the volume of lipid quantitatively. In addition, current NIRS system does not have capabilities to evaluate other features of vulnerable plaque such as thin fibrous cap, macrophage infiltration. Further advances may permit detection of other plaque features implicated in vulnerability.

Coronary angioscopy

Intracoronary coronary angioscopy has the capability of direct visualization of the surface of the plaque through fiber optics in vivo with high resolution (<150 μm).

Visualized plaque features by coronary angioscopy

The advantage of coronary angioscopy is to assess plaque color and surface characteristics. Plaque color can be graded as white, light yellow, yellow or intensive yellow by coronary angioscopy (Figure 6). White plaques on coronary angioscopy correlate with mainly fibrous plaque, whereas yellow plaques have been shown to harbor lipid rich tissue or necrotic core. The intensity of yellow color is associated with its instability, and high yellow intensity of plaques has been reported to exhibit thin, fibrous caps overlying a lipid core (70-72). Morphology of plaques is also evaluable by coronary angioscopy. Stable plaque is lesion with smooth surface, whereas complex plaque exhibits an irregular surface. Ruptured plaque, eroded plaque, intimal flap, fissure and ulceration are another complex lesions identified by coronary angioscopy (73). This modality has the ability to clearly visualize thrombus, which is a coalescent red or white mass, or both, adhering to the intima or protruding into the inner lumen (74). On histological validation, while the sensitivity of grayscale IVUS for the detection of thrombus was only 57%, its sensitivity was 100% under angioscopy imaging (73).

Limitations of angioscopy

There are some limitations of coronary angioscopy. Firstly, it is difficult to conduct quantitative analysis of plaques with regard to color and volume. Second, coronary angioscopy images only surface of the coronary lumen. It requires displacement of blood within coronary arteries by injecting saline continuously.

Potential links between imaging features and the corresponding cardiovascular risk

One of expected role of intracoronary imaging is the prediction of future cardiovascular events. Current available studies regarding the prediction of future cardiovascular events are summarized as follows and in the Table 1.

Grayscale IVUS

There is limited evidence showing the association of
echolucent plaques with subsequent ACS events. Yamagishi et al. has demonstrated that echolucent plaques within non-culprit segments were more likely to develop ACS during follow-up period (75). Another study analyzing IVUS data of 4,137 patients with CAD has reported that baseline plaque burden and its progression rate were associates with major adverse cardiovascular events (76). Clinical outcome in patients having grayscale IVUS derived plaque phenotypes requires further investigation.

**Virtual histology intravascular ultrasound (VH-IVUS)**

The Providing Regional Observations to Study Predictors of Events in Coronary Tree (PROSPECT) trial, used angiography, three-vessel grayscale and VH-IVUS to evaluate the natural history of atherosclerosis in a 697 patients with ACS. This study has demonstrated that minimal luminal area of <4 mm², a plaque burden of >70%, and VH-IVUS derived TCFA were independent determinants of future adverse cardiac events in non-culprit lesions (77). Another study also elucidated TCFA on VH-IVUS was associated with cardiovascular events in 170 patients with stable angina or ACS requiring PCI (78).

**Optical coherence tomography (OCT)**

Currently, there is no study demonstrating that high-risk plaques on OCT predict future cardiovascular events. One of reasons is the lack of large-scale study to elucidate predictive ability of OCT imaging in cardiovascular events. Specific OCT-derived plaque features associated with clinical outcomes might not be fully evaluated. There are still needs to validate the significance of plaque microstructures on OCT imaging.

**Near-infrared spectroscopy (NIRS)**

Oemrawsingh et al. has recently reported that LCBI predicts future cardiovascular events in patients with CAD. In this study, LCBI above 43.0 within non-culprit arteries had a 4-fold risk of major adverse cardiovascular events in 203 patients during 1-year follow-up (79).

**Angioscopy**

It has been reported that the presence of yellow plaques was associated with future ACS events (71). The number of yellow plaques has also been shown to predict future ACS events. Evaluation of angioscopic findings in

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**Table 1 Features of intravascular imaging modalities**

<table>
<thead>
<tr>
<th>Features</th>
<th>Grayscale IVUS</th>
<th>VH-IVUS</th>
<th>OCT</th>
<th>NIRS</th>
<th>Angioscopy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axial resolution</td>
<td>150 μm</td>
<td>200 μm</td>
<td>10-20 μm</td>
<td>N/A</td>
<td>&lt;150 μm</td>
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<tr>
<td>Studies regarding the prediction of cardiovascular events</td>
<td>Echolucent plaque is at risk for ACS (75)</td>
<td>Minimal lumen area of &lt;4 mm², plaque burden &gt;70% and VH-derived TCFA predicts ACS (77)</td>
<td>No available evidence exists</td>
<td>Lipid core burden index (LCBI) &gt;43.0 is at risk of MACE (79)</td>
<td>Yellow plaques predict ACS (70)</td>
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<td></td>
<td>Higher plaque burden is associated with MACE (76)</td>
<td>VH-derived TCFA predicts MACE (78)</td>
<td></td>
<td>Number of yellow plaques predicts ACS (80)</td>
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<td></td>
<td>In-stent yellow plaque is at risk of MACE (81)</td>
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<tr>
<td>Advantages</td>
<td>Precise tomographic measurement of lumen area, plaque size and distribution</td>
<td>Ability to differentiate plaque composition</td>
<td>Excellent axial resolution</td>
<td>Ability to assess fibrous cap thickness and macrophage filtration</td>
<td>High sensitivity and specificity in the detection of lipid rich plaque</td>
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<td>Direct visualization of plaque color and morphology of coronary surface</td>
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<tr>
<td>Disadvantages</td>
<td>Limited axial resolution, Inaccuracy in the assessment of plaque composition</td>
<td>Limited axial resolution, Inability in the assessment of plaque composition behind calcium</td>
<td>Low tissue penetration</td>
<td>Need of contrast medium flush for image acquisition</td>
<td>Lack of prospective data</td>
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<td>Need of continuous injection of saline during image acquisition</td>
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IVUS, intravascular ultrasound; VH-IVUS, virtual histology intravascular ultrasound; OCT, optical coherence tomography; NIRS, near infrared spectroscopy; ACS, acute coronary syndrome; MACE, major cardiovascular events.
552 patients has demonstrated number of yellow color plaque as the independent risk factors of ACS events (80). In another study, in-stent yellow plaque at 1-year after stent implantation was associated with higher incidence of major cardiovascular events (81).

Summary and future perspective

The present review describes that (I) various invasive imaging modalities allow to evaluate both plaque burden and composition in vivo; (II) plaque imaging provides a unique opportunity to identify vulnerable plaques, elucidate atherosclerotic mechanisms contributing to coronary events and the effect of anti-atherosclerotic medical therapies on coronary atheroma. Although these modalities are considerably promoting our knowledge of coronary atherosclerosis, there remains some limitations regarding axial resolution, penetration and differentiation of plaque composition. Several future directions can be considered to overcome these limitations. Firstly, Improvement in the spatial resolution of conventional modalities may enable more accurate analysis of atherosclerotic plaque. For example, micro OCT has been recently developed and tested in cadaver coronary arteries. As the resolution of this modality is around 10 μm, cellular and subcellular features associated with atherogenesis are clearly imaged (82). Secondly, fusion of different imaging modalities may cover inherent limitation of each modality (83). Combined system of NIRS and IVUS is one of examples enabling to analyze both vessel structure and plaque composition simultaneously. Given that IVUS has limited ability in evaluating plaque composition and NIRS does not provide measurers of vessel wall, combination of NIRS and IVUS helps clinicians to assess both morphological and compositional characteristics of plaques. As such, on-going innovation of technology will be expected to develop novel imaging devices which contain further sophisticated ability to image plaques. Additionally, we have to understand and investigate how these modalities can be used more effectively to manage patients with CAD.

Acknowledgements

None.

Footnote

Conflicts of Interest: Y Kataoka is supported by Cerenis Therapeutics. Other authors have no conflicts of interest to declare.

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