The assessment of intermediate coronary lesions using intracoronary imaging

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Abstract: Intermediate coronary artery stenosis, defined as visual angiographic stenosis severity of between 30–70%, is present in up to one quarter of patients undergoing coronary angiography. Patients with this particular lesion subset represent a distinct clinical challenge, with operators often uncertain on the need for revascularization. Although international guidelines appropriately recommend physiological pressure-based assessment of these lesions utilizing either fractional flow reserve (FFR) or quantitative flow ratio (QFR), there are specific clinical scenarios and lesion subsets where the use of such indices may not be reliable. Intravascular imaging, mainly utilizing intravascular ultrasound (IVUS) and optical coherence tomography (OCT) represents an alternate and at times complementary diagnostic modality for the evaluation of intermediate coronary stenoses. Studies have attempted to validate these specific imaging measures with physiological markers of lesion-specific ischaemia with varied results. Intravascular imaging however also provides additional benefits that include portrayal of plaque morphology, guidance on stent implantation and sizing and may portend improved clinical outcomes. Looking forward, research in computational fluid dynamics now seeks to integrate both lesion-based physiology and anatomical assessment using intravascular imaging. This review will discuss the rationale and indications for the use of intravascular imaging assessment of intermediate lesions, while highlighting the current limitations and benefits to this approach.

Keywords: Optical coherence tomography (OCT); intravascular ultrasound (IVUS); coronary artery disease (CAD)

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Introduction

Coronary artery disease (CAD) remains a major cause of morbidity and mortality worldwide. Many patients with obstructive CAD will experience symptoms of myocardial ischaemia and a reliable assessment of stenosis severity is required to ensure clinicians make appropriate revascularisation decisions. Traditionally, invasive coronary angiography has been the gold standard investigation to assess severity and extent of CAD. However, angiography is limited as it is a two-dimensional representation of a more complex three-dimensional structure. Interpretation of images can also be hampered by vessel foreshortening or overlay, along with challenges in reliably assessing stenosis severity in highly eccentric lesions. The fallibility of stenosis severity on angiography has been documented in previous studies, which demonstrate an inter-observer variance in diameter stenosis of 15–45%. Thus, visual estimation
of stenosis severity lacks precision with up to 30% of angiographic assessments found to be erroneous (1). While some of this subjectivity can be addressed through use of quantitative coronary angiography (QCA) software, there remains issues with spatial and temporal resolution along with an inability to account for extraluminal abnormalities, marked vessel tortuosity and lesion asymmetry (2). Three dimensional QCA has gone some way in addressing these concerns allowing fusion of two or more angiographic views. However, even 3D QCA cannot overcome the limitation of angiographic resolution (3).

Given these inherent difficulties in accurate stenosis assessment on angiography, patients often fall into a category termed ‘intermediate’ stenosis severity. Intermediate stenosis is defined as an angiographic stenosis of between 30–70% and presents a distinct clinical challenge with uncertainty in regards to optimal assessment and management strategy (4). Although its prevalence in the general population is not well known, registry data suggests it may be present in up to 25% of patients undergoing coronary angiography (5).

Invasive physiological assessment is recognized as the optimal method for assessment of intermediate angiographic lesions. Myocardial fractional flow reserve (FFR) is defined as the ratio between maximum coronary blood flow in a stenotic artery compared with maximum blood flow in the theoretical absence of stenosis. This assessment is undertaken during the conditions of maximal hyperaemia, to ensure the effects of the coronary microcirculation are minimised (6). Pressure derived FFR is the most commonly utilised invasive tool and has been found to be superior to angiography when used to guide revascularization strategy (7,8). Recent data also now support the role of hyperaemic-free pressure derived indices, such as the instantaneous wave-free ratio (iFR) (9,10). Both physiological approaches are supported by current European Society of Cardiology (ESC) guidelines on myocardial revascularization suggesting that, when evidence of ischaemia is not readily available, FFR or iFR should be utilized to assess the haemodynamic significance of intermediate grade coronary stenosis (11).

More recently, there has also been the development of imaging based technology that provides lesion-specific physiological measurements without the need for guidewire-based technology. The most well investigated method, termed Quantitative Flow Ratio (QFR), has demonstrated good diagnostic accuracy when compared to invasive indices. QFR utilises an image-based virtual FFR index, computed by three-dimesional QCA combined with coronary flow velocity estimated by thrombolysis in myocardial infarction (TIMI) frame count (12). In a series of 265 patients assessing diagnostic performance of QFR, there was strong correlation with both FFR (r=0.863, P<0.001) and iFR (r=0.740, P<0.001) producing a diagnostic accuracy of 90.8% and 81.3% (P<0.001), respectively (13). Despite these initial encouraging results, the uptake of QFR remains limited due to time constraints of calculating results in real-time, along with cost issues relating to software and licensing.

However, pressure wire assessment of intermediate lesions is not without its limitations and/or contraindications. Certain patient and lesion specific factors may decrease the efficacy of FFR and/or iFR and therefore physiological assessment may not be suitable in all patients. Patient specific characteristics may include; patient contraindication to adenosine (e.g., profound bradycardia), concurrent or recent ischaemia, significant left ventricular hypertrophy, severe aortic valve stenosis and situations where maximal hyperaemia (in the case of FFR) cannot be achieved due to significant microvascular impairment. In these scenarios, FFR may over- or underestimate lesion-specific ischaemia and may not be as reliable in guiding revascularization (14). Lesion specific features may also hamper physiological assessment including aorto-ostial, left main and tandem lesions along with segments of diffuse disease (15). These specific patient/lesion subsets may benefit from assessment with the use of intravascular imaging, chiefly intravascular ultrasound (IVUS) or optical coherence tomography (OCT). This review will therefore review the evidence and discuss the role of intravascular imaging in the assessment of intermediate coronary artery lesions.

**Basic principles of IVUS**

Intravascular ultrasound makes use of a miniaturized piezoelectric transducer mounted to a catheter tip to produce ultrasound signals, enabling quantitative assessment of the vessel wall and intimal atherosclerotic lesions. IVUS is now widely utilised as an adjunct to coronary angiography, allowing clinicians to better identify and characterize plaque morphology, measure reference luminal dimensions and optimize stent size, placement and deployment (16,17). Several manufacturers provide IVUS systems globally, enabling the majority of catheterization laboratories to have reliable access to the technology. The capabilities of IVUS continue to evolve, with improving resolution by higher acoustic frequency (up to 60 MHz for the opticross HD) and increasing pullback speeds; providing more accurate...
and rapid lesion assessment. Several post-processing mathematical methods have also been developed to improve characterization of plaque and tissue morphology above that of standard grey-scale IVUS imaging, including iMAP-IVUS, virtual-histology (VH-IVUS) and integrated-backscatter (IB-IVUS) (18,19). Further innovations have included integration of other intravascular imaging technologies such as infrared, fluorescent spectroscopy, or photoacoustic imaging with IVUS to better assess lesions for markers of vulnerability such as fibrous cap thickness and plaque composition (20). These multi-modal catheters are currently limited in their commercial availability, as they are yet to be proven to provide significant additional clinical benefit to patient management beyond grayscale IVUS alone.

**Clinical outcomes of IVUS-guided PCI**

Several large meta-analyses investigating the clinical benefits of IVUS-guided PCI with bare metal- (BMS) and drug eluting stents (DES) have been reported, with general consensus that IVUS improves patient outcomes (21,22). Focusing on DES, meta-analyses of RCTs and observational studies investigating IVUS-guided PCI have reported significant reductions in all-cause and cardiac death, target lesion revascularization (TLR), myocardial infarction (MI), and stent thrombosis (ST) (21,23,24); which has been confirmed at patient-level analysis (25). In complex lesions, IVUS reduced MACE and TLR with no effect on death or ST (26). These analyses however vary in their inclusion/exclusion criteria, with some limited to RCTs and others including observational studies, increasing the risk of confounding factors and treatment-selection bias. This was highlighted by Buccheri et al. who found that when their analysis was limited to only RCTs, IVUS-associated reduction in all-cause death was lost (23). Nevertheless, there is increasing consensus that IVUS-guided PCI improves clinical outcomes and now features in the American College of Cardiology (ACC) and ESC guidelines for guiding treatment of unprotected left-main coronary artery (LMCA) lesions (Class IIa, level B evidence), and is indicated for non-LMCA lesions of angiographically intermediate stenosis by the ACC guidelines (Class IIb, level B evidence) (27,28).

**IVUS assessment of LMCA disease**

LMCA disease is particularly challenging to accurately assess on angiography, as the arterial segment is often short, calcified and has disease that frequently involves the bifurcation. Unsurprisingly, it is known that there is high interobserver variability and low (41% to 59%) agreement when assessing angiographic lesion severity at this particular location (29,30). Given the critical importance of the LMCA to the coronary circulation and consequently its high prognostic value, accurate lesion assessment is crucial in guiding clinical management.

IVUS has been widely used in the assessment of intermediate LMCA lesions for many years. Indeed, IVUS-derived minimal lumen dimension is known to be a strong predictor of MACE (31). Numerous studies have sought to identify a standardized IVUS-derived measure that can be used to assist in guiding therapy, with the minimal lumen area (MLA) receiving the greatest interest (Figure 1). A wide range of MLA cut-off values have been proposed (between 4.5 and 7.5 mm²) to guide revascularisation versus medical management, based on the ability to correlate significantly with an FFR of <0.8 (32-36). The MLA cut-off of 6 mm² is often used as the standard to defer revascularisation, as this value was validated in the largest prospective multicentre (LITRO) study to date of 354 patients from 22 centres across Spain. Utilising IVUS, de la Torre Hernandez et al. found no significant difference in either cardiac-death (94.5% vs. 97.7%, P = 0.5) or event-free survival (80.6% vs. 87.3%, P = 0.3) between those who underwent revascularisation (MLA < 6 mm², n = 179) or in those is which it was deferred (MLA ≥6 mm², n = 186) after 2 years (34). Of the 179 deferred patients, just 8 required revascularisation and this was predominantly due to stable angina, with no significant clinical, angiographic or ultrasonographic differences from those not requiring revascularisation. This MLA cut-off is supported by a prior study that identified an MLA cut-off of 5.9 mm² correlated significantly with FFR <0.75 in 55 patients with moderate LMCA stenosis, with 93% sensitivity and 94% specificity (Table 1) (32). Opinion however does vary. The Mayo Clinic (USA) suggested a higher MLA cut-off value of 7.5 mm², derived from the mean ± 2 standard deviations of MLA from a cohort of 121 patients with angiographically normal/minimally diseased LMCA (33). Subsequent long-term follow-up of 214 patients with intermediate LMCA lesions at 3.3±2.0 years revealed no significant difference in MACE (TVR, MI, death) between those that underwent revascularization (n=83) or were deferred (n=131) using an MLA cut-off of 7.5 mm² alone (33).

Smaller MLA cut-off values have been suggested based on studies from South Korea, in which 4.5–4.8 mm²...
correlated significantly with FFR <0.8 in two studies, and an MLA of 4.1 mm$^2$ correlated more significantly with FFR<0.75 (Table 1) (35,36). Moreover, the same study found that among 40 lesions with MLA <6.0 mm$^2$, 38% had FFR ≥0.8, and 58% had FFR ≥0.75 (35). These studies suggest that ethnicity and possibly body habitus should be taken into consideration in determining a safe MLA cut-off value in clinical practice. However, when taken together, these studies highlight that an IVUS-defined MLA cut-off of >6 mm$^2$ appears to be a safe deferral threshold for revascularisation across Western and Asian populations.

IVUS guided LMCA PCI has been found to significantly reduce all-cause mortality by 40%, cardiac death by 53%, plus reduce the risk of stent thrombosis and TLR in a meta-analysis of 10 studies with 6,480 patients undergoing IVUS-versus angiographic-guided stent insertion alone (51). Sub-group analysis revealed this significance was maintained across different study types, ethnic populations, and duration of follow-up (both <3 and ≥3 years). A more recent meta-analysis of 7 studies including 4,592 patients also found IVUS guidance to significantly reduce all-cause death, cardiac death, stent thrombosis and MI, but had no significant benefit on TLR, likely due to the study being underpowered (52). It must be noted that only one

![Figure 1](image-url) Intravascular ultrasound assessment of an intermediate left main coronary artery stenosis. Eccentric and calcified distal left main coronary artery (LMCA) lesion (A). Representative cross-sectional intravascular ultrasound (IVUS) images of the proximal reference segment with highlighted minimal luminal area (MLA) (highlighted blue) (C) and target lesion MLA (blue) highlighting eccentricity and superficial calcification (D). The calculated MLA (shaded blue) was 5.1 mm$^2$, which is below the safe deferral threshold of 6 mm$^2$ (B). Panel B reproduced from Jasti et al. Circulation, 2004 (32).
A randomized controlled trial was included in each of these meta-analyses and, to our knowledge, just one further prospective RCT has been published since, in which IVUS-guidance significantly reduced MACE (driven by reduced cardiac death), with no significant influence on MI, TLR or ST in 336 patients (53). How IVUS-guidance reduced mortality while having no significant effect on MI, ST or TLR remains to be elucidated. In the aforementioned trial the authors suggest it is likely related to reduced procedural adverse events, whereby IVUS enables greater characterisation of the vessel and lesion, improving stent sizing, placement, expansion and apposition, resulting in greater mean stent diameter (53). This is supported by a patient-level pooled registry analysis where IVUS improved clinical outcomes post-PCI in LMCA lesions (54).

LMCA lesions are frequently associated with downstream

### Table 1 Optimal IVUS-defined minimal luminal area (MLA) values to best predict FFR

<table>
<thead>
<tr>
<th>Study</th>
<th>Patient/lesions</th>
<th>FFR</th>
<th>Best MLA cut-off (mm²)</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>NPV (%)</th>
<th>PPV (%)</th>
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<tbody>
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<td><strong>LMCA studies</strong></td>
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<tr>
<td>Park 2014 (36)</td>
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<td>0.80</td>
<td>4.5</td>
<td>78.7</td>
<td>80.4</td>
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<td>5.9</td>
<td>93.0</td>
<td>95.0</td>
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<td>–</td>
</tr>
<tr>
<td>Kang 2011 (35)</td>
<td>55/55</td>
<td>&lt;0.80</td>
<td>4.8</td>
<td>89.0</td>
<td>83.0</td>
<td>89.0</td>
<td>82.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&lt;0.75</td>
<td>4.1</td>
<td>95.0</td>
<td>83.0</td>
<td>97.0</td>
<td>75.0</td>
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<tr>
<td>Takagi 1999 (37)</td>
<td>42/51</td>
<td>0.75</td>
<td>3.00</td>
<td>83.0</td>
<td>92.3</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Lee 2010 (38)</td>
<td>94/94</td>
<td>0.75</td>
<td>2.00</td>
<td>87.9</td>
<td>78.9</td>
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<td>–</td>
</tr>
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<td>4.00</td>
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<td>56.0</td>
<td>96.0</td>
<td>46.0</td>
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<td>All 2.75</td>
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<td>65.0</td>
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<td></td>
<td></td>
<td></td>
<td>Prox-LAD 3.00</td>
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<td></td>
<td>Mid-LAD 2.75</td>
<td>73.0</td>
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<td>83.0</td>
<td>68.0</td>
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<td>3.09</td>
<td>69.2</td>
<td>79.5</td>
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<tr>
<td>Kang 2012 (42)</td>
<td>692/784</td>
<td>0.80</td>
<td>2.40</td>
<td>84.0</td>
<td>63.0</td>
<td>90.0</td>
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<td>206/206</td>
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<td>Prox-LAD 3.20</td>
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<td>66.7</td>
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<td></td>
<td></td>
<td></td>
<td>Mid-LAD 2.50</td>
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<td>87.7</td>
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<td>Han 2014 (44)</td>
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<td>61.0</td>
<td>63.0</td>
<td>73.0</td>
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<td>2.75</td>
<td>65.0</td>
<td>68.0</td>
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<tr>
<td></td>
<td>242/258</td>
<td>0.80</td>
<td>3.00</td>
<td>63.0</td>
<td>69.0</td>
<td>–</td>
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<tr>
<td>Waksman 2013 (45)</td>
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<td>0.80</td>
<td>3.07</td>
<td>64.0</td>
<td>65.0</td>
<td>83.0</td>
<td>40.0</td>
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<td>0.80</td>
<td>3.15</td>
<td>71.4</td>
<td>67.0</td>
<td>85.3</td>
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<td>2.70</td>
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<td>76.0</td>
<td>89.9</td>
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<td>56/61</td>
<td>0.80</td>
<td>2.36</td>
<td>67.0</td>
<td>65.0</td>
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<td>67.0</td>
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<td>Ben-Dor 2011(49)</td>
<td>84/92</td>
<td>0.80</td>
<td>3.20</td>
<td>69.2</td>
<td>68.3</td>
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<tr>
<td></td>
<td>0.75</td>
<td>2.80</td>
<td>79.7</td>
<td>80.3</td>
<td>–</td>
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</tr>
<tr>
<td>Brown 2017 (50)</td>
<td>89/92</td>
<td>0.80</td>
<td>2.65</td>
<td>58.3</td>
<td>67.6</td>
<td>–</td>
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NPV, negative predictive value; PPV, positive predictive value; MLA, minimal luminal area; FFR, fractional flow reserve; LMCA, left main coronary artery.
and/or multivessel disease (55), potentially confounding the ability of FFR to accurately assess the true functional significance of the LMCA stenoses. The effect of downstream disease was assessed by Fearon et al., who demonstrated that true FFR was significantly lower than that measured in the presence of a stenosis (0.81±0.08 vs. 0.83±0.08, P<0.001), even when the transducer was placed in a non-diseased vessel (56). Although the magnitude of this discrepancy is small and unlikely to affect clinical decision making in the vast majority of cases, the authors did note that the functional significance of LMCA disease cannot be accurately determined in cases where FFR in the non-diseased vessel is between 0.81 and 0.85 and in the presence of deep ischaemia due to severe downstream stenosis (FFR <0.45). These scenarios raise the possibility that intravascular imaging may be of use in assisting clinicians make revascularisation decisions during angiography (Figure 2).

This diagnostic value of IVUS in LMCA lesions is recognised in ESC guidelines, recommending IVUS should be considered to assess the severity of unprotected left main lesions (Class IIa Level B) and considered to optimize treatment of unprotected left main lesions (Class IIa Level B). However, a MLA cut-off value is not yet detailed in guidelines, likely due to the discrepancies across reports and lack of large trial data (28).

**IVUS assessment of non-LMCA lesions**

The current gold standard to assess significance of non-LMCA intermediate lesions remains physiological

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**Figure 2** Hybrid use of intracoronary imaging and physiology to guide coronary revascularisation. Coronary angiography demonstrating intermediate distal left main coronary artery (LMCA) stenosis (A) with downstream severe disease in the mid left anterior descending (LAD) artery (B). Both fractional flow reserve (C) and resting full-cycle ratio (F) assessment confirm ischaemia within the LAD territory. Optical coherence tomography imaging demonstrates the minimal luminal area within the LMCA is 7.95 mm² (D), highlighting that targeted revascularisation of the proximal to mid LAD should be considered and precluding need for intervention of the LMCA.
assessment (57). However, given the relative ease and safety of IVUS, several studies have sought to identify IVUS-derived criteria (namely MLA) that can identify significant lesions with high diagnostic accuracy. To date, studies have generally reported poor diagnostic accuracy (66–74%) of IVUS-derived MLA in correlating with FFR ≤0.8 in non-LMCA intermediate lesions (50). One promising study has reported an 83% sensitivity and 92.3% specificity to identify a FFR ≤0.75 (MLA cut-off 3 mm²) (37); however, subsequent studies have generally reported poorer diagnostic capability (Table 1) (42,46,58), and in one study of small vessels (<2.8 mm wide and <20 mm long) no correlation was found (59). Recent meta-analyses of non-LMCA studies report mean MLA cut-off of ~2.60 mm² with sensitivity and specificity of 75-79% and 65-66% respectively, to identify a FFR ≤0.8 (60,61). Further diagnostic parameters are outlined in Table 1, in which IVUS-derived MLA is associated with (generally) far greater negative predictive values than positive predictive values, indicating that IVUS-derived MLA can provide clinical benefit in identifying stenoses acceptable for deferral, but should not be used in isolation to validate revascularisation of non-LMCA vessels.

The diagnostic accuracy of IVUS is further driven by vessel location and anatomical variants. For example, a significant correlation between MLA and FFR <0.8 was identified in major epicardial vessels >3.5 mm, but not side branches <2.25 mm in 77 patients with intermediate ostial lesions (62). Table 1 lists the range of MLA cut-off values (2–4 mm²) reported in non-LMCA intermediate lesions including results from Koo et al., that revealed the need for different cut-off MLA values based on lesion location, and suggests that IVUS is unsuitable in vessels with high anatomical variation (40). Altering MLA cut-off values based on vessel diameter has been supported in one RCT in which 400 pairs of propensity score matched patients underwent PCI with a MLA of <4 mm² in vessels >3 mm, and <3.5 mm² in vessels 2.5–3 mm, or FFR ≤0.75 (63). MACE-free survival was equivalent between FFR and MLA-guided groups up to 2 years follow-up, with significantly more interventions performed in the IVUS compared with FFR group (48.8% vs. 28% P<0.001). Similarly, Nam et al. found IVUS-guidance (MLA cut-off <4 mm²) resulted in a greater number of interventions compared to FFR-guidance (≤0.8) in 167 patients with intermediate coronary lesions, with no significant difference in MACE at 1 year. They also calculated that reducing the MLA cut-off to <3 mm² equalized the number of interventions between groups, concluding that both FFR and IVUS were suitable for use in guiding PCI in patients with intermediate lesions (64). Increased stent use in IVUS-guidance is often reported, with Park et al. also finding IVUS-guided therapy is associated with a greater number (as well as larger and longer stents used) with no improvement in patient outcomes, but rather a significant increase in target lesion failure predominantly due to periprocedural MI in a non-randomized post-hoc analysis of 463 matched patients from the EXCELLENT trial (65).

IVUS-derived MLA has also been shown to correlate significantly with physiological indices other than FFR, including iFR, diastolic pressure ratio (dPR), and resting full-cycle ratio (RFR) (66). Despite the significant correlation, the diagnostic accuracy remains low (60.5–62.4%; comparable with FFR 61.7%), with the best cut-off MLA being 3.4 mm² across all indices (66). Matsushita et al. also recently reported IVUS-derived MLA to have just 67% sensitivity and 69% specificity to predict significant stenosis (iFR ≤0.89) requiring a cut-off MLA <1.8 mm²; equivalent to the FFR <0.8 (cut-off MLA 2.0 mm²) (67). Together, these data suggest that IVUS provides limited diagnostic accuracy in predicting functionally significant stenosis identified by FFR and iFR.

Reasons for the relatively poor diagnostic accuracy of IVUS in identifying functionally significant CAD is due to several factors including ethnicity, FFR cut-off, vessel size, and high anatomical variations. It is well established that MLA cut-off differs between Asian and Western populations, with comparably lower MLA cut-off values reported in Asian studies of 2.75 vs. 3.0 mm² (44) and 2.68 vs. 3.03 mm² (61), respectively. The reasons suggested for this may be associated with a lower body mass index and myocardial mass in Asian patients, but also potentially different lesion characteristics between ethnicity’s (44,61). The accuracy of IVUS-derived MLA can be improved when the target FFR is reduced to <0.75 (49,60,61). However, most studies adopt an FFR of 0.8 to provide greater clinical relevance and minimise the number of untreated lesions that may result in ischemia, and as such validating an MLA cut-off value for a FFR <0.75 may be inappropriate despite the improved accuracy (61). Alternatively, accuracy may be improved by combining other IVUS-derived parameters with the MLA, including stenotic area, minimal luminal diameter (MLD), and lesion length; increasing the sensitivity up to 100% in one study (37-39). Furthermore, computational “machine learning algorithms” are now being developed which are significantly improving the
diagnostic accuracy of IVUS to identify lesions with an FFR \( \leq 0.8 \) up to 86\%, with the aim to reduce the need for pressure wire assessment (68).

**Basic principles of OCT**

Optical coherence tomography is an intravascular imaging technique which utilises light waves and time-delay information in a modality analogous to ultrasound providing a real-time portrayal of intraluminal and transmural coronary anatomy (69). By analysing the interference pattern of light waves emitted by the catheter using a tool known as the interferometer, the system is able to generate a ‘depth-profile’ of the luminal walls allowing detailed assessment. Since the light waves used have a much higher frequency than sound waves, the spatial resolution (i.e., the number of pixels used to construct a single image) offered by OCT is greater than that of IVUS, albeit with poorer tissue penetration (70).

First demonstrated *in vivo* in ophthalmological applications, OCT has found use in the characterisation of intracoronary lesions during percutaneous intervention (71). OCT systems begin by scanning the light beam across a cross-sectional sample to capture an ‘A-scan,’ which is a 1-dimensional depth profile. As the catheter automatically and rapidly rotates within the portion of coronary artery being assessed, this process is repeated to generate a 2-dimensional ‘B-scan’ (72). The pullback system then begins, imaging the vessel in this same manner along a pre-determined length of its lumen. The series of B-scans are then compiled into a 3-dimensional, volumetric image known as a ‘volume scan’ (73). As infrared light cannot penetrate blood, OCT systems require blood to be flushed from the arterial lumen prior to analysis as the presence of erythrocytes causes excess scatter, significantly decreasing imaging quality (71,73). Methods for removing blood from the imaging field are divided into occlusive, involving gentle balloon inflation proximal to the imaged segment, or non-occlusive. Non-occlusive methods are most common with the automated intracoronary administration of crystalloids, iso-osmolar solution, or more commonly contrast (74).

The use of OCT provides several technical and clinical benefits over IVUS. Firstly, OCT systems are capable of achieving faster rotational speeds and hence quicker data acquisition than IVUS, improving workflow and minimising procedural time (75). OCT also provides significantly higher resolution (10–15 versus 100 \( \mu \)m) when compared with IVUS (76). The improvement in resolution allows the operator to more readily identify dissections, ulceration, thrombus, characterise the size of the coronary vessel and detect stent malapposition (75–77). Furthermore, the features of the backscattered light captured by the catheter on their return can help differentiate atherosclerotic plaques into fibrous or fibrocalcific, and identify presence of a necrotic core, seen as signal-poor regions with poorly-delineated borders and little-to-no backscatter (78). Newer generation OCT catheters also have increased deliverability, being able to be manipulated through tortuous and small calibre coronary vessels.

There are however limitations. Given the light source used by OCT approaches the infrared spectrum with a wavelength average of 1,350 nm, there is an inherent limitation to the depth of field of OCT imaging. This particularly becomes problematic during imaging of the LMCA, where the vessel dimensions can regularly exceed 4mm. Furthermore, it can be difficult to create a blood-free field in the LMCA due to the high coronary flow volumes in this anatomical location. This can interfere with image acquisition and limits assessment of luminal dimensions and plaque morphology. Finally, OCT cannot be used in the assessment of aorto-ostial lesions of the right or left coronary arteries as it is not possible to adequately clear blood from these locations and thus these lesions lend themselves more towards IVUS assessment (79).

**OCT assessment of intermediate lesions**

Similar to IVUS, OCT has also found relevance in the quantification of LMCA lesion severity (*Figure 3*). Although initial data correlating MLA and FFR was obtained and validated using IVUS (80), these criterion are increasingly being extrapolated to OCT (*Figure 4*). Experimental phantom models have shown that IVUS may overestimate luminal area by up to 10\%, while OCT measurements are highly accurate to ‘true’ luminal dimensions. Thus, *in vivo* OCT typically presents smaller cross-sectional luminal areas than IVUS and operators need be aware of this discrepancy when interpreting images. Based on these observations, LMCA lesions with MLA \( >5.4 \text{ mm}^2 \) on OCT are generally considered a safe cut-off for deferral of revascularisation (81).

There is increasing data supporting OCT-derived MLA as a superior predictor of functional ischaemia, when compared with IVUS in non-LMCA. In a study of 186 patients assessing 203 *de novo* lesions diagnostic efficacy of MLA obtained by OCT and IVUS to predict an FFR of \( <0.75 \) was assessed. Receiver operator characteristic
curve analysis showed that OCT-MLA had a statistically significantly better diagnostic efficacy than IVUS-MLA in identifying ischaemia (FFR \(<0.75\)) with the best MLA cut-off values of 1.39 mm\(^2\) (AUC: 0.732, 95% CI: 0.660–0.804) and 2.57 mm\(^2\) (AUC: 0.615, 95% CI: 0.53–0.696) for OCT and IVUS respectively. Interestingly, there was no significant difference between OCT- and IVUS-MLA for the prediction of FFR \(\leq 0.80\) (P=0.13) (82). Although this study suggests that OCT may have superior efficacy to IVUS in detecting functional ischaemia, it must be cautioned that intravascular imaging is currently not interchangeable with FFR in clinical decision making but rather these results suggest that OCT may be preferable to IVUS where there may be doubt, and may reduce the incidence of unnecessary coronary intervention.

Previous meta-analyses support these findings. A metaanalysis of 6,919 patients with 7537 non-LMCA lesions found OCT- and IVUS-derived MLA had similar sensitivity in predicting haemodynamically significant lesions (0.732 vs. 0.747, P=0.519). OCT-derived MLA however had higher specificity (76.3% vs. 66.5%, P<0.001) and diagnostic accuracy in detecting flow limiting lesions than IVUS respectively (AUC 0.810 vs. 0.754, P=0.045) (83). Of the seven trials included in the analysis specifically utilising OCT, 340 patients and 438 lesions were assessed. These and other similar studies are presented in Table 2. In the largest trial included in the analysis, Reith et al. found that amongst their 100 patients with 142 lesions, receiver operator characteristic analysis demonstrated a moderate diagnostic efficacy of OCT-derived MLA with a best cut-off values of 1.64 mm\(^2\) (AUC: 0.836, 95% CI: 0.772–0.901) for the prediction of FFR \(\leq 0.80\) (90). These combined results suggest that OCT, with its superior imaging resolution, is the preferred intravascular imaging modality for the
Figure 4 Optical coherence tomography and fractional flow reserve. Intermediate lesion of the distal left main coronary artery (A), which extends into the proximal left anterior descending (LAD). Initial optical coherence tomography images (B) demonstrate a minimal luminal area of 2.64 mm², with evidence of a small intimal dissection (at 11 o’clock). Fractional flow reserve (FFR) assessment of the vessel (C) demonstrates that the lesion is highly haemodynamically significant (FFR 0.67).

detection of haemodynamically significant stenosis in non-LMCA lesions.

More recent data has also suggested that OCT may be of equal, or even superior, to FFR in guiding PCI decisions in intermediate lesions. The FORZA trial was a direct comparison between OCT and FFR in a 1:1 randomisation pattern. If either FFR or OCT identified a lesion as significant, based on current criteria (FFR ≤0.80 or OCT area stenosis ≥75% or 50–75% with MLA <2.5 mm²), the patient underwent PCI with optimisation using the same technique. Using pre-defined endpoints of MACE and residual angina, FFR was found to be a more cost-effective strategy with more patients being medically managed (62.5% vs. 44.8%; P<0.001) however an OCT-guided PCI strategy saw a reduction in all-cause mortality, non-fatal MI, TVR, kidney injury, and significant angina at 13-month follow-up (92). Although the results of this trial are unlikely to alter the current guidelines on physiological assessment of intermediate lesions, it does raise the possibility that an intravascular image-guided approach may yield very similar outcomes and should not be considered inappropriate.

FFR is not without its limitation and as discussed, LMCA lesions may be associated with downstream disease limiting accurate assessment with FFR alone. There is no data utilizing OCT in this clinical scenario however as seen in Figure 2, OCT in combination with FFR assessment
may aid operators in accurate assessment of lesions which require intervention.

**Future directions**

Moving forward, the future of OCT and IVUS assessment may lie in coupling with computational fluid dynamics (CFD) simulations of coronary flow and pressure. Use of CFD technology has potential for clinicians to gather simultaneous anatomical and functional assessment of individual coronary lesions. Bezerra *et al.* (93) have applied these concepts to reconstructed 3D IVUS images, showing a moderate-to-high correlation with invasive pressure wire FFR ($r=0.79$; $P<0.001$). The overall accuracy, sensitivity, specificity, positive predictive value and negative predictive value of this approach was 91%, 89%, 92%, 80% and 96%, respectively to identify FFR <0.80. Similar results have also been shown with OCT, and may be preferable given its higher resolution and more robust estimates of luminal dimensions. In a study of 92 patients with intermediate stenosis undergoing both OCT and FFR assessment, a CFD algorithm was utilised to calculate computational FFR (FFROCT). With a functionally significant stenosis defined as an FFR of <0.80, FFROCT resulted in accuracy, sensitivity and specificity of 88%, 69% and 96% respectively. There remains some hurdles though before this technology is more widely adopted. Firstly, further validation work is required in more diverse lesion and patient cohorts. Secondly, computational time for CFD is still relatively high and this precludes use in a high paced cath lab environment (94). Nevertheless, there is clear potential for the future and there are several industry led collaborations attempting to deliver this technology to clinicians soon.

**Conclusions**

Optimal management of intermediate CAD on angiography remains a significant clinical challenge. Although invasive physiological assessment with FFR or iFR remains current gold standard, a number of clinical scenarios may push clinicians towards assessment of lesion severity using intravascular imaging.

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**Footnote**

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| Table 2 Optimal OCT-defined minimal luminal area (MLA) values to best predict FFR |
|-----------------|----------|-----------------|------------------------------|-----------------|-----------------|-----------------|-----------------|
| Study           | Patient/lesions | FFR  | Best MLA cut-off (mm$^2$) | Sensitivity (%) | Specificity (%) | NPV (%) | PPV (%) |
| Gonzalo 2012 (48) | 56/61 | 0.80 | 1.95 | 80.0 | 83.0 | 80.0 | 83.0 |
| Usui 2018 (84) | 186/203 | 0.75 | 1.39 | 72.5 | 66.4 | 54.3 | 81.4 |
| Shiono 2012 (85) | 59/62 | 0.75 | 1.91 | 93.5 | 77.4 | 92.3 | 80.6 |
| Belkacemi 2013 (86) | 27/27 | 0.80 | 2.54 | 71.0 | 84.0 | – | – |
| Pyxaras 2013 (87) | 36/55 | 0.80 | 2.88 | 73.0 | 71.0 | – | – |
| Pawlowski 2013 (88) | 48/71 | 0.80 | 2.05 | 75.0 | 90.0 | 92.6 | 70.6 |
| Zafar 2014 (89) | 30/41 | 0.80 | 1.62 | 70.0 | 97.0 | 91.0 | 89.0 |
| Reith 2015 (90) | 100/142 | 0.80 | 1.64 | 78.8 | 75.8 | 73.4 | 80.8 |
| Burzotta 2018 (91) | 40/40 | 0.80 | 2.50 | 64.0 | 93.0 | – | – |

NPV, negative predictive value; PPV, positive predictive value; MLA, minimal luminal area; FFR, fractional flow reserve; LMCA, left main coronary artery.
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