Alternative methods for functional assessment of intermediate coronary lesions

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Abstract

Wire-based fractional flow reserve (FFR) is a diagnostic tool used to evaluate the ischemic burden of coronary lesions. Large-scale studies have shown that FFR-guided revascularization is associated with better clinical outcomes. However, wide adoption of this technology is limited due to the considerable cost, additional time needed for set-up and performance of the measurement as well as the invasiveness of the procedure which requires pressure wire placement across the lesion into the distal segment of the coronary artery. To overcome these limitations new, promising, and less-/non-invasive methods were developed. These methods are based on computational fluid dynamics analysis and three-dimensional lumen reconstruction. The aim of this paper is to review scientific evidence supporting the clinical safety and efficacy of these techniques, such as instantaneous wave-free ratio, quantitative flow ratio and FFR calculated from computed tomographic angiography. (Cardiol J 2020; 27, 6: 825–835)

Key words: coronary angiography, quantitative flow ratio, computational fluid dynamics, fractional flow reserve

Introduction

Coronary artery disease (CAD) is one of the main causes of morbidity and mortality in developed countries [1, 2]. Coronary angiography remains the gold standard for the diagnosis of CAD, however, its ability to differentiate ischemic from non-ischemic lesions is limited. In this respect, fractional flow reserve (FFR), which takes into consideration the functional severity of coronary stenosis, outperforms the traditional diagnostic approach, based solely on morphometric assessment [3, 4]. Unfortunately, the adoption of FFR in everyday clinical practice is slow and is utilized in only a minority of centers [5, 6]. Härle et al. [7] found that FFR was used in 3.2% of all diagnostic procedures performed in Germany. In Poland penetration rate of FFR was even lower and did not exceed 2% in 2014 [8]. The main limiting factors include: 1) considerable time need for set-up and conduction of the examination; 2) high cost of diagnostic probe and adenosine infusion; 3) invasiveness, as it requires insertion of a pressure wire across the lesion into the distal part of the vessel, which is associated with increased risk of serious complications, e.g. ventricular arrhythmias and coronary vessel dissection (occurring in 0.5% of procedures), and 4) patient-related contraindications (hypotension, asthma, second-degree atrioventricular blocks) [9–11]. To overcome these limitations, less invasive techniques based on computational fluid dynamics (CFD) and three-dimensional (3D) lumen reconstruction have been proposed [12–16].
**Pressure wire methods**

**Instantaneous wave-free ratio**

Instantaneous wave-free ratio (iFR) is one alternative method that does not require adenosine infusion (Fig. 1). Although vessel wiring is still necessary, iFR measurements are quicker to perform and are cheaper than FFR. The scientific basis came from findings by Sen et al. [17] who demonstrated that functional assessment of coronary lesions comparable to FFR is possible without drug induced hyperemia, during the so-called “wave-free period”. This period is seen in diastole and characterized by minimal and stable coronary resistance (similar to “hyperemic-like” conditions), which makes the trans-stenotic pressure gradient corresponding directly to flow and lesion severity [17].

The first published clinical study evaluating the correlation between iFR and FFR (ADenosine Vasodilator Independent Stenosis Evaluation [ADVISE]) demonstrated a close correlation between values obtained with these two methods ($r = 0.9; p < 0.001$) [17]. The possibility of iFR real-time measurement was proven by ADVISE in-practice study. The authors assessed 392 angiographically intermediate lesions and demonstrated that the best cutoff value of iFR corresponding to $FFR \leq 0.80$ was an iFR $\leq 0.90$ and resulted in classification agreement in 80% of cases, specificity of 79%, sensitivity of 81%, positive predictive value (PPV) of 71% and negative predictive value (NPV) of 87% [18]. What is more, it was shown that iFR correlates more closely than FFR with coronary flow velocity reserve, which suggests that iFR may be a more physiological parameter of disease severity [19].

In 2017, two pivotal trials evaluating iFR in clinical practice were published. The Functional Lesion Assessment of Intermediate Stenosis to Guide Revascularization (DEFINE-FLAIR) trial consisting of almost 2500 patients with stable CAD, proved that iFR-guided is noninferior to FFR-guided coronary revascularization with respect to composite risk of death from any cause, nonfatal myocardial infarction (MI) or unplanned revascularization during 1-year follow-up. Additionally, study results showed that in iFR group median procedural time was significantly shorter (40.5 vs. 45 min; $p = 0.001$; iFR vs. FFR, respectively) and fewer patients had adverse procedural symptoms (3.1% vs. 30.8%; $p < 0.001$; iFR vs. FFR, respectively) [20]. The Instantaneous Wave-free Ratio versus Fractional Flow Reserve in Patients with Stable Angina Pectoris or Acute Coronary Syndrome (iFR-SWEDE-HEART) trial consisting of over 2000 patients with stable CAD or acute coronary syndrome (17.5% patients) showed similar results. The primary composite end-point (defined as composite of death from any cause, nonfatal MI or unplanned revascularization) occurred in 6.7% of the patients in the iFR group and in 6.1% of the patients in the FFR group in 1-year follow-up ($p = 0.007$ for noninferiority).
Authors, just as in the previously described study, reported that chest discomfort occurred less often during the iFR-guided procedure (3.0% vs. 68.3%; p < 0.001) [21]. Results of these two trials were reflected in European and in American guidelines, in which iFR was regarded as equivalent to FFR in hemodynamic assessment of intermediate-grade stenosis [22, 23].

**Alternative pressure wire methods**

Over the years other adenosine-free methods based on assessment of diastolic resting indices have been proposed. Recently published data proved a high correlation between iFR and resting distal coronary to aortic pressure (Pd/Pa). Both were associated with lesion anatomic and hemodynamic severity, showing excellent agreement between them [24, 25]. It seems that the adoption of Pd/Pa could be easier, in comparison to iFR, it was analyzable in a significantly higher number of cases [25]. Other diastolic resting indexes included resting Pd/Pa during the complete duration of diastole, in 25% to 75% of diastole, at midpoint of diastole (Fig. 2). All the above-mentioned parameters were proven to be identical to iFR, not only numerically, but also with respect to their agreement to FFR [26]. Though, they are all very promising, further studies are needed to evaluate their clinical value.

**Computational-based methods**

**Quantitative flow ratio**

In 2013 Morris et al. [13] published results from the VIRTUal FFR From Coronary Angiography (VIRTU-1) study, designed to demonstrate the feasibility of FFR computations based solely on two-dimensional (2D) coronary angiography images (virtual FFR [vFFR]). The study population consisted of 19 patients. Compared to FFR, vFFR had an accuracy of 97%, sensitivity of 86%, specificity of 100%, PPV of 100%, and NPV of 97%. Although, there was a strong correlation between vFFR and wire-based FFR (r = 0.84), the image analysis was labor- and time-consuming, requiring 24 h to process the above-mentioned data. Of note, authors used a “one-size fits all” approach, which assumed constant coronary vessel resistance. Such an assumption carries the risk of stenosis misclassification due to possible changes in downstream microcirculatory resistance [13].

Papafakis et al. [27] proposed virtual functional assessment index (vFAI) — a quick method of functional assessment of intermediate coronary lesions, which took only 15 min to analyze one vessel. This approach computes distal to proximal pressure ratio over the lesion based on 3D quantitative coronary angiography (QCA) reconstruction and steady-flow CFD. The method was compared to FFR in 120 patients showing accuracy of 88%, sensitivity of 90% and specificity of 86% for the optimal vFAI cut-off point (≤ 0.82). Additionally, the vFAI was superior to 3D QCA in predicting hemodynamic significance of coronary stenosis and demonstrated close correlation and good agreement with wire-based FFR values. The main limitation of vFAI, which is based solely on lesion geometry, is the fact that it does not take into account microvascular resistance and size of myocardial territory subtended by the vessel [27].

To overcome these limitations, the computed FFR (FFR_{QCA}) based on mean volumetric flow rate at hyperemia derived from 3D vessel invasive coronary angiography (ICA) reconstruction, Thrombolysis in Myocardial Infarction (TIMI) frame count and CFD utilization was proposed (Fig. 3). The analysis of 77 vessels provided an 88% overall accuracy of FFR_{QCA} for diagnosis of ischemia (defined as FFR ≤ 0.8). There was a strong correla-
tion between \( \text{FFR}_{\text{QCA}} \) and FFR values \((r = 0.81, p < 0.001)\) with a mean difference of \( \pm 0.06 \) \((p = 0.054)\) [12].

One of the main advantages of this method is short computation time, which did not exceed 10 min in total processing. Additionally, this method provides an evaluation of the entire coronary tree, whereas in wire-based FFR, only those lesions in which a pressure wire is inserted can be assessed [28].

Further confirmation of diagnostic accuracy of fast computational approaches came from prospective, observational, multicenter Functional Assessment by Various Flow Reconstruction (FAVOR) pilot study, in which 3 different quantitative flow ratio (QFR) computations were compared with standard wire-based FFR measurements. These included: 1) fixed-flow QFR (fQFR) that assumed a universal hyperemic flow velocity of 0.35 m/s; 2) contrast-flow QFR (cQFR) based on individual virtual flow derived from the frame count during contrast injection; 3) adenosine-flow (aQFR) based on individual virtual flow derived from the frame count during maximal adenosine-induced hyperemia. Authors confirmed good agreement between wire-based FFR and each QFR computation. The diagnostic accuracy was comparable for cQFR (86%) and aQFR (87%) and was significantly higher compared to fQFR (80%) indicating that the use of adenosine is not needed in this method [29]. Recently QFR received Conformité Européenne (CE) certificate, allows for wider adoption to everyday clinical practice.

In 2017, results from The FAVOR II China (Functional Diagnostic Accuracy of Quantitative Flow Ratio in Online Assessment of Coronary Stenosis) study were also published. They prospectively enrolled 308 consecutive patients at 5 centers in China. The primary endpoint was to assess if QFR would improve diagnostic accuracy of coronary angiography. Authors met the pre-specified performance goal for level of diagnostic accuracy of QFR in identifying hemodynamically significant stenosis. Additionally, they confirmed QFR to have 94.6% sensitivity, 91.7% specificity, 85.5% PPV, and 97.1% NPV and diagnostic accuracy of 92.4% in patient-level analysis, and 92.7% in vessel-level analysis [30].

Figure 3. Computation of quantitative flow ratio (QFR) from coronary angiography; A. Angiographic projections of the left anterior descending (LAD) artery at > 25° apart; B. Fractional flow reserve (FFR) measured during intravenous adenosine infusion was 0.73; C. Computed QFR value indicates ischemia (QFR = 0.73). Arrow indicates original location of pressure transducer.
A recently published study demonstrated retrospectively analyzed results of 306 intermediate lesions, which had been previously evaluated using FFR. In contradiction to previous studies, which utilized core-lab assessment, in this particular study used an on-site QFR calculation in all cases. It showed that the Pearson correlation was strong for QFR ($r = 0.85$). Additionally, optimal QFR decision value of 0.79 was identified, this corresponded to FFR = 0.80 (AUC = 0.94). After introduction of the cut-off value of $\leq 0.74$ and $> 0.83$, an excellent diagnostic performance of QFR was achieved, with sensitivity and specificity > 95%. Additionally, it was confirmed that the time for QFR analysis was relatively short and substantially decreased with the number of analyzed cases. The first 50 QFR analysis took an average of 5 min 59 s, whereas in the final 50 cases the mean time was 2 min 7 s [31].

Westra et al. [32] prospectively evaluated QFR in 240 lesions and correctly classified 83% of the lesions when an FFR cut off value of 0.8 was used. They also achieved a sensitivity of 77%, specificity of 86%, PPV of 87%, and NPV of 75%.

In 2018 the results from The FAVOR II Europe-Japan Study were published. In this international, multicenter trial 329 patients were enrolled. QFR values were calculated online in catheterization laboratories during the procedure. Sensitivity and specificity were $> 86\%$ for QFR, which was significantly higher than for 2D QCA (sensitivity 44.2%; $p < 0.001$ and specificity 76.5%; $p = 0.002$) [33].

The most recently published study demonstrated that QFR may also be utilized in acute coronary syndrome settings, particularly in guiding non-culprit lesion revascularization in patients presenting with ST-segment elevation MI [34]. Additionally, the QFR good inter-core laboratory reproducibility had already been proven [35].

Although QFR is a very promising method, there are some technical limitations that should be taken into account. At present, the degree of flow-limiting stenosis of the ostial left main and ostial right coronary artery lesions cannot be reliably measured. Supraventricular tachyarrhythmia leading to an altered filling pattern of coronary arteries remains an exclusion criterion for FFRQCA calculation. Additionally, patients with coronary artery bypass grafting supplying evaluated vessels or with collateral circulation have not been adequately studied [12]. Last, the timing of contrast injection may also affect the FFRQCA values.

Additionally, data on clinical outcomes i.e. patient quality of life and cost-effectiveness remains lacking. This gap may will hopefully be addressed by the upcoming FAVOR III trial, which is designed as a prospective, randomized, multicenter clinical outcome study. With a planned enrollment of approximately 2000 patients, it is powered to establish the role of this method in the diagnostic process of CAD patients.

**Computed tomographic angiography**

Computed tomographic angiography (CTA) of the coronary vessels was the first non-invasive diagnostic imaging method providing data for CFD analysis to derive FFR-equivalent measurements (Fig. 4). Koo et al. [14] analyzed 103 patients, who underwent coronary CTA, QCA and FFR measurement. They performed the computation of FFR from coronary CTA (FFRCT) using a powerful supercomputer to calculate the above-mentioned values. The proposed method utilized semi-automated segmentation of coronary arteries and approximation of the left ventricular mass. Despite the high computing power, a single analysis took approximately 5 h. The FFRCT had an accuracy of 84.3%, sensitivity of 87.9%, specificity of 82.2%, PPV of 73.9%, and NPV of 92.2% for the diagnosis of ischemia-inducing lesions on a per-vessel basis. Additionally, there was a good...
<table>
<thead>
<tr>
<th>Method</th>
<th>Author</th>
<th>Year</th>
<th>Sample size</th>
<th>Primary endpoint</th>
<th>Major findings</th>
<th>Cutoff values</th>
</tr>
</thead>
<tbody>
<tr>
<td>iFR</td>
<td>Sen et al. [17]</td>
<td>2012</td>
<td>157 stenoses</td>
<td>To determine time period when intracoronary resistance is minimal and stable and there is a possibility to assess stenosis severity without adenosine administration</td>
<td>During &quot;wave-free period&quot; the intracoronary resistance is minimal and stable. There is good correlation between iFR and FFR ($r = 0.9; p &lt; 0.001$)</td>
<td>Ischemia: FFR $&lt; 0.80$</td>
</tr>
<tr>
<td>iFR</td>
<td>Petraco et al. [18]</td>
<td>2014</td>
<td>392 lesions</td>
<td>To evaluate iFR and FFR diagnostic agreement in “real-world” conditions</td>
<td>The best cutoff to discriminate stenosis with FFR $\leq 0.80$ was an iFR $\leq 0.90$; sensitivity — 81%; specificity — 79%; PPV — 71%; NPV — 87%</td>
<td>Ischemia: FFR $&lt; 0.80$</td>
</tr>
<tr>
<td>iFR</td>
<td>Petraco et al. [19]</td>
<td>2014</td>
<td>182 patients 216 stenoses</td>
<td>To assess the diagnostic relationship between iFR, FFR coronary flow velocity reserve</td>
<td>iFR has stronger correlation with coronary flow velocity reserve than FFR</td>
<td>Ischemia: FFR $\leq 0.75$</td>
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<tr>
<td>iFR</td>
<td>Davies et al. [20]</td>
<td>2017</td>
<td>2492 patients</td>
<td>To determine the efficacy and safety of an iFR-guided vs. an FFR-guided strategy for coronary revascularization</td>
<td>iFR-guided revascularization was non-inferior to FFR-guided procedure. Less adverse procedural symptoms and clinical signs (including chest pain and dyspnea) in iFR group. Median procedural time is shorter in iFR group.</td>
<td>Ischemia: FFR $\leq 0.8$ iFR $\leq 0.89$</td>
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<tr>
<td>iFR</td>
<td>Götberg et al. [21]</td>
<td>2017</td>
<td>2037 patients</td>
<td>To determine if iFR is noninferior to FFR with respect to the rate of subsequent major adverse cardiac events</td>
<td>iFR-guided revascularization was non-inferior to FFR-guided procedure. More chest discomfort in FFR group.</td>
<td>Ischemia: FFR $\leq 0.8$ iFR $\leq 0.89$</td>
</tr>
<tr>
<td>vFFR</td>
<td>Morris et al. [13]</td>
<td>2013</td>
<td>19 patients</td>
<td>To compare vFFR and FFR values</td>
<td>Accuracy — 97%; sensitivity — 86%; specificity — 100%; PPV — 100%; NPV — 97%; strong correlation between vFFR and FFR</td>
<td>Ischemia: FFR $&lt; 0.80$ Anatomically obstructive CAD: ICA with stenosis &gt;50%</td>
</tr>
<tr>
<td>vFAI</td>
<td>Papafaklis et al. [27]</td>
<td>2014</td>
<td>120 patients 139 vessels</td>
<td>To test the diagnostic performance of the derived vFAI in a real-world patient population with intermediate lesions and compare with FFR values</td>
<td>Accuracy — 88%; sensitivity — 90%; specificity — 86% (for the optimal vFAI cut-off point ($\leq 0.82$) vFAI was superior to 3D QCA in predicting hemodynamic significance of coronary stenosis; close correlation and good agreement with wire-based FFR values.</td>
<td>Ischemia: FFR $\leq 0.80$</td>
</tr>
<tr>
<td>FFRQCA</td>
<td>Tu et al. [12]</td>
<td>2014</td>
<td>68 patients 77 vessels</td>
<td>To assess FFRQCA predictive value to diagnose ischemia</td>
<td>Overall accuracy — 88%; strong correlation between FFRQCA and FFR; total processing time of around 10 min.</td>
<td>Ischemia: FFR $\leq 0.80$</td>
</tr>
<tr>
<td>QFR</td>
<td>Tu et al. [29]</td>
<td>2016</td>
<td>73 patients 84 vessels</td>
<td>To assess diagnostic accuracy of fast computational approaches to derive FFR from diagnostic coronary angiography</td>
<td>High diagnostic accuracy: fQFR = 80%, cQFR = 86%, aQFR = 87%</td>
<td>Ischemia: FFR $\leq 0.80$</td>
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Table 1 (cont.). Summary of studies evaluating methods used to compute fractional flow reserve.

<table>
<thead>
<tr>
<th>Method</th>
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</tr>
</thead>
<tbody>
<tr>
<td>QFR</td>
<td>Xu et al. [30]</td>
<td>2017</td>
<td>308 patients, 332 vessels</td>
<td>To assess the diagnostic performance of QFR for the diagnosis of hemodynamically significant coronary stenosis</td>
<td>Diagnostic accuracy: patient-level — 92.4%, vessel-level — 92.7%; sensitivity — 94.6%; specificity — 91.7%; PPV — 85.5%; NPV — 97.1%</td>
<td>Ischemia: FFR ≤ 0.80</td>
</tr>
<tr>
<td>QFR</td>
<td>Westra et al. [32]</td>
<td>2018</td>
<td>172 patients, 240 lesions</td>
<td>To evaluate the feasibility and diagnostic performance of QFR in unselected consecutive patients</td>
<td>QFR correctly classified 83% of the lesions; area under the receiver operating characteristic curve — 0.86; sensitivity — 77%; specificity — 86%; PPV — 87%; NPV — 75%</td>
<td>Ischemia: FFR ≤ 0.80</td>
</tr>
<tr>
<td>QFR</td>
<td>Westra et al. [33]</td>
<td>2018</td>
<td>272 patients</td>
<td>To evaluate the value of online QFR during routine ICA for procedural feasibility, diagnostic performance, and agreement with FFR</td>
<td>QFR analysis is superior to angiographic assessment; sensitivity — 86.5%; specificity — 86.9%; median time to QFR — 5 min</td>
<td>Ischemia: FFR ≤ 0.80</td>
</tr>
<tr>
<td>QFR</td>
<td>Kołtowski et al. [31]</td>
<td>2018</td>
<td>268 patients, 306 lesions</td>
<td>To assess diagnostic accuracy of QFR</td>
<td>Strong Pearson correlation for iQFR (r = 0.85), fQFR (r = 0.73), vQFR (r = 0.78) and iQFR (r = 0.70)</td>
<td>Sensitivity and specificity &gt; 95% for iQFR ≤ 0.74 (n = 89, 29%) and &gt; 0.83 (n = 116, 38%), respectively</td>
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<tr>
<td>FFRCT</td>
<td>Koo et al. [14]</td>
<td>2011</td>
<td>103 patients, 159 vessels</td>
<td>Diagnostic performance of FFR_{CT} and CCTA stenosis, compared with invasive FFR (as the reference standard)</td>
<td>Accuracy — 84.3%; sensitivity — 87.9%; specificity — 82.2%; PPV — 73.9%; NPV — 92.2% (for diagnosis of ischemia-inducing lesions on per-vessel basis); good correlation between FFR_{CT} and FFR values</td>
<td>Ischemia: FFR_{CT} and FFR ≤ 0.80 Anatomically obstructive CAD: CCTA with stenosis ≥ 50%</td>
</tr>
<tr>
<td>FFRCT</td>
<td>Min et al. [15]</td>
<td>2012</td>
<td>252 patients</td>
<td>Whether FFR_{CT} plus CT could improve the per-patient diagnostic accuracy such that the lower boundary of the 1-sided 95% confidence interval of this estimate exceeded 70%</td>
<td>Accuracy — 73%; sensitivity — 90%; specificity — 54%; PPV — 67%; NPV — 84% (for diagnosis of ischemia-inducing lesions on per-patient basis); high NPV; high sensitivity; authors did not achieve the prespecified level of per-patient diagnostic accuracy</td>
<td>Ischemia: FFR_{CT} and FFR ≤ 0.80 Anatomically obstructive CAD: CT and ICA with stenosis ≥ 50%</td>
</tr>
<tr>
<td>FFRCT</td>
<td>Nørgaard et al. [16]</td>
<td>2014</td>
<td>254 patients</td>
<td>Per-patient diagnostic performance as assessed by the area under the receiver-operating characteristic curve of FFR_{CT} ≤ 0.80 vs. CCTA (stenosis &gt; 50%) for the diagnosis of hemodynamically significant stenosis (FFR ≤ 0.80) in patients with CCTA stenosis of 30% to 90%</td>
<td>Accuracy — 86%; sensitivity — 84%; specificity — 86%; PPV — 61%; NPV — 95% (for diagnosis of ischemia-inducing lesions on per-vessel basis) FFR_{CT} under 0.8 correlated well with FFR values under 0.8</td>
<td>Ischemia: FFR_{CT} and FFR ≤ 0.80 Anatomically obstructive CAD: CT and ICA with stenosis &gt; 50%</td>
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### Summary of studies evaluating methods used to compute fractional flow reserve.

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<tr>
<td>FFR&lt;sub&gt;CT&lt;/sub&gt;</td>
<td>Douglas et al. [36]</td>
<td>2015</td>
<td>584 patients</td>
<td>Percentage of those with planned ICA in whom no significant obstructive CAD was found at ICA within 90 days</td>
<td>FFR &lt; 0.80</td>
<td>No major adverse cardiac events over 90-day follow-up period in any patient in whom ICA was cancelled based on negative results of CTA/FFR&lt;sub&gt;CT&lt;/sub&gt;</td>
</tr>
<tr>
<td>FFRCT</td>
<td>Nørgaard et al. [37]</td>
<td>2018</td>
<td>677 patients</td>
<td>To assess real-life clinical outcomes of introduction of strategy including CTA and selective FFRCT in patients with stable CAD</td>
<td>FFR&lt;sub&gt;CT&lt;/sub&gt; ≤ 0.80</td>
<td>FFRCT is an effective tool to differentiate patients with intermediate-grade coronary lesions who may need further invasive testing</td>
</tr>
<tr>
<td>FFR&lt;sub&gt;CT&lt;/sub&gt;</td>
<td>Min et al. [15]</td>
<td>2018</td>
<td>252 patients</td>
<td>To assess real-life clinical outcomes of introduction of strategy including CTA and selective FFRCT in patients with stable CAD</td>
<td>FFR&lt;sub&gt;CT&lt;/sub&gt; &lt; 0.80</td>
<td>Authors reported FFR&lt;sub&gt;CT&lt;/sub&gt;’s accuracy of 73%, sensitivity of 90%, specificity of 54%, PPV of 67%, and NPV of 84% for diagnosis of ischemia-inducing lesions on a per-patient basis. The study did not achieve its pre-specified level of per-patient diagnostic accuracy, however, it showed that adding FFR&lt;sub&gt;CT&lt;/sub&gt; analysis to plain CTA assessment improved diagnostic accuracy. Authors emphasized that FFR&lt;sub&gt;CT&lt;/sub&gt; had high negative predictive value and high sensitivity, indicating that coronary angiogram is not needed when FFR&lt;sub&gt;CT&lt;/sub&gt;’s results are normal, despite significant stenosis in CTA [15].</td>
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A refined version of FFR<sub>CT</sub> calculation was evaluated by Nørgaard et al. [16] who studied 254 patients with coronary CTA, QCA and FFR measurements. The new approach was significantly quicker with a mean time to results of less than 4 h (depending on CT scan quality and CAD burden). On a per-vessel basis, authors found diagnostic accuracy of 86%, sensitivity of 84%, specificity of 86%, PPV of 61%, and NPV of 95% for FFR<sub>CT</sub> under 0.8, which correlated well with FFR values under 0.8. They concluded that FFR<sub>CT</sub> has high diagnostic performance compared with standard FFR measurements [16].

The multicenter Prospective Longitudinal Trial of FFR<sub>CT</sub>: Outcome and Resource Impacts (PLATFORM) trial evaluated FFR<sub>CT</sub> guided revascularization looking at clinical outcomes, cost/resource utilization and quality of life. Overall 584 patients with new onset of chest pain were included. Patients were randomized to standard evaluation (usual care arm) and CTA/FFR<sub>CT</sub> testing. In the usual care arm, significantly more patients who underwent coronary angiography had no obstructive CAD when compared to FFR<sub>CT</sub> care arm (73.3% vs. 12.4%; p < 0.0001). This observation was further confirmed in a propensity score matching analysis of 148 pairs (72% vs. 12%; p < 0.0001). Most importantly there were no major adverse cardiac events over the 90-day follow-up correlation between FFR<sub>CT</sub> and FFR values, with a slight underestimation by FFR<sub>CT</sub> (0.022 ± 0.116; p = 0.016). Authors concluded that the addition of FFR<sub>CT</sub> to standard coronary CTA measurements might enhance diagnostic accuracy and this method’s utility [14].
defined as total coronary vessel lumen volume relative to left ventricular mass, has a statistically significant influence on FFR CT’s accuracy and specificity.

Conclusions

Functional assessment of coronary arteries remains a gold standard in the diagnosis of patients with intermediate coronary artery stenosis. In current clinical practice, the adoption of traditional wire-based FFR technology is slow and limited by clinical safety and economic constraints. QFR and FFRCT are the new, less-/non-invasive computational methods that have recently emerged as promising diagnostic tools. The currently available body of evidence, though limited, provides solid grounds in recognizing these technologies as strong candidates to reduce the number of wire-based FFR examinations (Table 1). Advantages and disadvantages of above-mentioned diagnostic tools are summed up in Table 2.

It is thought herein, that all these methods will find their place in the management of patients with CAD. It seems that QFR and FFRCT should be perceived as more complementary, rather than competitive modalities. While FFRCT may lead to better identification of patients who would not benefit from ICA investigation, QFR may be used on-line to assess the hemodynamic significance of a lesion during ICA and eliminate risks associated with wiring of a coronary artery. It is essential to utilize cut-off values in which QFR has excellent agreement with FFR measurements (“grey-zone” concept). If results of the upcoming clinical validation will be positive, one may foresee a change in the current diagnostic algorithm by incorporating

<table>
<thead>
<tr>
<th>Comparator</th>
<th>FFR</th>
<th>iFR</th>
<th>QFR</th>
<th>FFRCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invasiveness:</td>
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<td>Contrast</td>
<td>+</td>
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<td>Invasive coronary angiography</td>
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<td>Pressure wire</td>
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<tr>
<td>Adenosine</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Data acquisition and processing time</td>
<td>8–10 min*</td>
<td>5–7 min*</td>
<td>3–5 min*</td>
<td>4–6 h*</td>
</tr>
<tr>
<td>Online/offline processing</td>
<td>Online</td>
<td>Online</td>
<td>Online</td>
<td>Offline</td>
</tr>
<tr>
<td>Costs</td>
<td>+++</td>
<td>+++</td>
<td>+</td>
<td>++++</td>
</tr>
</tbody>
</table>

*Excluding standard invasive coronary angiography time and standard computed tomography angiography time; FFR — fractional flow reserve; FFRCT — FFR calculated from computed tomography; iFR — instantaneous wave-free ratio; QFR — quantitative flow ratio.
alternative methods for functional assessment of intermediate coronary lesions (Fig. 5).

**Conflict of interest:** None declared

**References**


