Evidence robustly demonstrates that ischemia, rather than anatomy, is the optimal target for coronary revascularization. In the cardiac catheter laboratory, fractional flow reserve (FFR) and corresponding diastolic indices are regarded as the gold standard for physiological lesion assessment and ischemia detection (Table 1). Yet, despite a wealth of supporting data and indications in international guidelines, the use of FFR remains surprisingly low in the diagnostic assessment of coronary artery disease across the world.1,2 To address this, multiple groups have developed methods for computing FFR from invasive angiography, without the need for passing a pressure wire or inducing hyperemia, thus removing the main barriers to uptake. Angiography-derived FFR therefore has the potential to extend the benefits of physiological coronary lesion assessment to considerably more patients. Given the size of the interventional cardiology market, clinical and commercial motivation to deliver these tools as quickly as possible could hardly be greater. Several models are now approved as medical devices. Imminently, physicians and healthcare providers will have to decide whether to use these tools. But do they truly deliver physiology, and are they accurate enough? There are 3 particular areas of that deserve close scrutiny.

SIMPLIFICATION

Methods for computing angiography-derived FFR are software based. Three-dimensional arterial anatomy is reconstructed from paired 2-dimensional angiogram images. Mathematical equations that define hemodynamic laws are then applied to the reconstructed artery to predict the pressure dynamics along the artery, which are displayed as a color-mapped 3-dimensional artery. In an effort to rationalize these models to make them practical and expedient for clinical use, many groups have abandoned complex, numerical, computational fluid dynamics simulation in favor of analytical solutions based broadly upon the laws of Bernoulli and/or Poiseuille. These simpler physical laws characterize pressure losses attributable to convective acceleration and viscous friction, respectively. They are quick and simple to execute and perform well under steady (nonpulsatile), laminar flow conditions, in straight conduits. Coronary arteries, however, are not straight, and flow is pulsatile. Furthermore, these laws are unable to accurately characterize complex translesional pressure dynamics, particularly poststenosis pressure recovery, which is the basis of FFR. Some stenosis models make empiric assumptions or corrections for pressure loss and recovery. On average, these may perform adequately, but cannot represent the potentially complex flow patterns in a specific case. Moreover, they may be particularly vulnerable to inaccuracy in the context of serial lesions and diffuse disease in which 3-dimensional computational fluid dynamics computations more reliably characterize interstenosis hemodynamic interaction. The impact this has on accuracy, in all disease patterns, is yet to be fully determined.

Key Words: computational flow dynamics ■ computer-based model ■ coronary microvascular resistance ■ fractional flow reserve ■ imaging
ASSUMPTIONS
The discordance between angiographic severity and physiological (FFR) significance is well described and affects ≥30% of lesions. Discrepancies occur because, unlike angiography, FFR elegantly and automatically incorporates the combined and inter-related effects of coronary flow and microvascular resistance. It is therefore imperative that computational models of angiography-derived FFR include adequate physiological inputs or “tuning” to represent the maximum blood flow or minimum microvascular resistance; the latter dictates the former, which, in turn, dictates the pressure gradient and FFR. Hemodynamic equations are capable of accurately deriving a variety of physiological parameters, but only if other appropriate physiological inputs, such as flow or microvascular resistance, are included. A sensitivity analysis demonstrated that microvascular resistance was the dominant influence on angiography-derived FFR, above and beyond the severity or anatomy of epicardial disease. Hyperemic flow and minimal microvascular resistance are variable in health and disease and are hard to measure, even with invasive instrumentation. Noninvasive models of angiography-derived FFR therefore rely upon assumptions about these parameters, or predict them from surrogate markers such as arterial diameter. Again, empiric assumptions may be sufficient overall, for many cases, but will be inaccurate in nonaverage cases with discordant anatomy and physiology, that is, the very cases where FFR is superior to angiography. Therefore, unless models have an accurate method for achieving this, on a patient-specific basis, the “physiological” prediction becomes simply a function of stenosis geometry and they cannot be a genuine model of FFR at all (Figure). As an example, 1 study of angiographically derived FFR observed a significant reduction in diagnostic accuracy in patients with elevated microvascular resistance. Paradoxically, physiologically weak models will appear more feasible relative to angiographic appearance, and a potential danger is that user confidence may therefore be increased with poorer methods. FFR has enabled a great stride forward in terms of physiologically guided revascularization. It would be unfortunate if, in an attempt to increase physiological assessment, we were to take half a step back toward assessment based on epicardial arterial anatomy. Table 2 summarizes major trials of angiography-derived FFR.

ACCURACY AND ERROR RANGE
Headline validation results report “diagnostic” accuracy. This quantifies how well a method predicts physiological significance or nonsignificance (FFR ≤0.80), relative to invasive FFR, expressed as sensitivity, specificity, positive, and negative predictive values, area under a receiver operating curve, and overall diagnostic accuracy. Diagnostic accuracy is a function of (1) the method’s accuracy and (2) the cases included in a particular study. The fewer cases close to the 0.80 threshold, the better the diagnostic accuracy will appear and vice versa. This is nicely illustrated in a study of FFR computed from computed tomography coronary angiography in which the diagnostic accuracy was 82% overall, but only 46% in cases in FFR were 0.70 to 0.80, which is precisely the range where most accuracy is required.

The best test of how accurately angiography-derived FFR agrees with invasive FFR is to plot the
### Table 1. Angiography-Based Coronary Physiological Assessment Techniques

| Index                                      | Abbreviation                | Calculated                                                                 | Equipment                                                      | Potential Benefits                                                                                           | Pitfalls/Limitations                                                                 |
|--------------------------------------------|-----------------------------|---------------------------------------------------------------------------|-----------------------------------------------------------------|------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|
| Fractional flow reserve                    | FFR                         | Whole cardiac cycle Pd/Pa at hyperemia                                    | Pressure wire                                                   | Predicts percentage improvement in flow with PCI. Good clinical outcomes data                               | Does not measure absolute flow and microvascular resistance                           |
| Instantaneous wave-free ratio/            | iFR/RFR                     | Pd/Pa during diastolic phase                                              | Pressure wire                                                   | Good clinical outcome data, relative to FFR                                                               | Does not measure absolute flow and microvascular resistance                           |
| resting full-cycle ratio                  |                             |                                                                           |                                                                  |                                                                                                           |                                                                                       |
| Index of myocardial resistance             | IMR                         | (Pd) · (thermodilution derived mean transit time)                         | Thermo- and pressure-sensitive wire                             | Microvascular resistance becoming of increasing interest (eg, PCI nonresponders, ANOCA, AMI, HFpEF)       | Thermodilution not widely used                                                       |
| Hyperemic microvascular resistance         | HMR                         | Pd/Doppler flow velocity                                                  | Doppler and pressure wire                                       | Microvascular resistance becoming of increasing interest (eg, PCI nonresponders, ANOCA, AMI, HFpEF)       | Doppler flow velocity challenging to measure. Doppler wires not widely used           |
| Hyperemic stenosis resistance              | HSR                         | (Pd-Pa)/Doppler flow velocity                                             | Doppler and pressure wire                                       | Objective, direct measure of the resistance of proximal disease                                         | Doppler flow velocity challenging to measure. Doppler wires not widely used. Surrogate index |
| Angiography-derived FFR                   | vFFR/QFR                    | Fluid dynamics equations informed by anatomy                              | Computational fluid dynamics software                            | Delivering clinical benefits of FFR without factors that limit the invasive technique                     | Relatively wide Bland–Altman limits of agreement compared with FFR. Requires excellent angiography. Less accurate in those with nonaverage microvascular resistance |
| CT-derived FFR                            | cFFR                        | Fluid dynamics equations informed by anatomy                              | Computational fluid dynamics software (offline)                  | Reduce the number of unnecessary invasive catheterizations                                              | Relatively wide Bland–Altman limits of agreement compared with FFR                   |
| Coronary flow reserve                      | CFR                         | (Hyperemic flow surrogate)/(baseline flow surrogate)                      | Doppler or thermosensitive wire                                 | A surrogate for flow and vasodilatory reserve. Flow more important than pressure, but hard to measure     | Prone to same limitations as those for Doppler wire or thermodilution. Variability in baseline measurement can impair interpretation |
| Absolute coronary flow                     | Qb                          | Infusion flow · (infusion temp/sensor temp) · 1.08                        | Thermosensitive wire, pressure wire, monorail infusion catheter  | Predicts absolute (not percentage) coronary flow changes and microvascular resistance                   | Additional time, expertise, and hardware                                             |
Table 2. Major Trials/Studies of Angiographically Derived FFR

| Author           | Study                | Year | N=Arteries | Surrogate/Software/Company                                                                 | Mathematical Solution | Diagnostic Accuracy | 95% Limits of Agreement |
|------------------|----------------------|------|------------|------------------------------------------------------------------------------------------|------------------------|----------------------|-------------------------|
| Morris et al⁵    | VIRTU-1              | 2013 | 35         | vFFR from VIRTUheart (University of Sheffield)                                            | Transient 3D CFD      | 97%                 | FFR ±0.16               |
| Tu et al⁶        | FAVOUR Pilot         | 2016 | 84         | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | Empiric flow velocity (fQFR), TIMI frame counting-derived contrast velocity at baseline (cQFR) and under hyperemia (aQFR), Analytical equations based on laws of Bernoulli and Poiseuille | fQFR 80%              | cQFR 86%              | FFR ±0.14               |
|                  |                      |      |            |                                                                                           |                        | aQFR 87%              |                        | FFR ±0.13               |
| Komowski et al⁷  | FFR₂₃₀, FIM          | 2016 | 101        | FFR₂₃₀ (CathWorks, Israel)                                                                | Simple analytical equation, based on law of Poiseuille            | 94%                 | FFR ±0.10               |
| Trobs et al⁸     | FFR₂₃₀               | 2016 | 100        | FFR₂₃₀ from Syngo IZ3D and prototype software (Siemens Healthcare GmbH, Germany)           | CFD based on BP, anatomy, and literature estimates of microvascular resistance | 90%                 | FFR ±0.13               |
| Pellicano et al⁹ | FFR₂₃₀, validation   | 2017 | 203        | FFR₂₃₀ (CathWorks, Israel)                                                                | Simple analytical equation, based on law of Poiseuille            | 93%                 | FFR ±0.10               |
| Xu et al¹⁰       | FAVOUR II China      | 2017 | 328        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | 93%                 | FFR ±0.13               |
| Yazaki et al¹¹   | QFR in intermediate lesions | 2017 | 151        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | 88%                 | FFR ±0.10               |
| Westra et al¹²   | WIFI II              | 2018 | 240        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | 83%                 | FFR ±0.16               |
| Mejía-Rentería et al¹³ | QFR IMR study       | 2018 | 300        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | IMR <23 =88% IMR ≥23 =76% | FFR ±0.12               |
|                  |                      |      |            |                                                                                           |                        |                      | FFR ±0.15               |
| Westra et al¹⁴   | FAVOUR II EJ         | 2018 | 317        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | 87%                 | FFR ±0.12               |
| Fearon et al¹⁵   | FAST-FFR             | 2019 | 319        | FFR₂₃₀ (CathWorks, Israel)                                                                | Simple analytical equation, based on law of Poiseuille            | 92%                 | FFR ±0.13               |
| Omori et al¹⁶    | FFR₂₃₀, in multivessel disease | 2019 | 118        | FFR₂₃₀ (CathWorks, Israel)                                                                | Simple analytical equation, based on law of Poiseuille            | 92%                 | FFR ±0.14               |
| Stahli et al¹⁷   | All comer QFR        | 2019 | 516        | QFR from QAngio XA (Medis Medical Imaging Systems, NL)                                     | TIMI frame counting-derived contrast velocity at baseline (cQFR), Analytical equations based on laws of Bernoulli and Poiseuille | 93%                 | FFR ±0.07               |
| Masčedi et al¹⁸  | FAST-study           | 2019 | 100        | vFFR from 3D QCA software, CAAS workstation (PIE Medical Imaging, NL)                     | Simple analytical equation, based on laws of Bernoulli and Poiseuille | AUC=0.93              | FFR ±0.07               |
| Li et al¹⁹       | FLASH-FFR            | 2019 | 328        | caFFR from FlashAngio (Rainmed Ltd, China)                                                | CFD based on postangiography TIMI frame counting of flow velocity  | 96%                 | FFR ±0.10               |

Listed in chronological order. Invasive FFR (threshold ≤0.80) was comparator in each study. 3D indicates 3-dimensional; aQFR, adenosine QFR; AUC, area under the curve; BP, blood pressure; caFFR, coronary angiography-derived fractional flow reserve; CFD, computational fluid dynamics; cQFR, contrast QFR; EJ, Europe and Japan; FFR, fractional flow reserve; FFR₂₃₀, FFR derived from coronary angiography; FIM, first in man; fQFR, fixed QFR; IMR, index of microcirculatory resistance; QFR, quantitative flow ratio; TIMI, thrombolysis in myocardial infarction; and vFFR, virtual fractional flow reserve.
differences between predicted and observed FFR values against the mean (ie, a Bland–Altman plot). From this, the mean difference (delta), which quantifies any bias in the angiography-derived method, and the 95% limits of agreement, are calculated. The limits of agreement (±1.96 SDs) comprise 95% of observed differences and are akin to the 95% CI of a computed, angiography-derived FFR result or an error range (Figure). The wider the limits of agreement, the larger the method’s error and vice versa. Unlike diagnostic accuracy, the limits of agreement are only a function of how accurate a method is. A recent meta-analysis of 13 studies of angiography-derived FFR demonstrated impressive diagnostic accuracy (sensitivity, 89%; specificity, 90%), but more-sobering agreement, with limits of agreement of FFR ±0.14.20 This is remarkably similar to FFR computed from computed tomography in the NXT trial (limits of agreement FFR ±0.15).21 FFR computed from computed tomography, however, is a noninvasive screening tool, best used to reduce unnecessary invasive catheterization. Arguably, the accuracy “bar” should be set far higher for a test in the catheter laboratory, where results directly influence coronary anatomy, to provide personalized management and improved clinical outcomes. However, deriving physiology from anatomy is challenging and requires assumptions. Model simplification and physiological assumptions, based on extrapolated or averaged data, are likely to work in the majority of patients. However, much of FFR’s success lies in its ability to identify those cases where nonstandard microvascular resistance and/or flow result in discordant physiology and anatomy. It is therefore important that models of angiography-derived FFR retain the same patient-specific physiology that separates traditional FFR from angiography, or at least that they highlight which cases require more-reliable assessment. Operators must understand how accuracy and error are defined in all patient groups. Stringent validation is required to prove that models are accurate and physiologically sound, in the hands of those who will be using them. If this can be achieved, clinicians have the potential to achieve what could be a new level of patient-specific medicine.

CONCLUSIONS

Angiography-derived FFR has the potential to change clinical practice for the considerable benefit of patients by providing routine physiological data, together with coronary anatomy, to provide personalized management and improved clinical outcomes. However, deriving physiology from anatomy is challenging and requires assumptions. Model simplification and physiological assumptions, based on extrapolated or averaged data, are likely to work in the majority of patients. However, much of FFR’s success lies in its ability to identify those cases where nonstandard microvascular resistance and/or flow result in discordant physiology and anatomy. It is therefore important that models of angiography-derived FFR retain the same patient-specific physiology that separates traditional FFR from angiography, or at least that they highlight which cases require more-reliable assessment. Operators must understand how accuracy and error are defined in all patient groups. Stringent validation is required to prove that models are accurate and physiologically sound, in the hands of those who will be using them. If this can be achieved, clinicians have the potential to achieve what could be a new level of patient-specific medicine.

ARTICLE INFORMATION

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