Background Percutaneous coronary intervention (PCI) guided by fractional flow reserve (FFR) measurement is superior to visual angiographic assessment alone. We have developed a workflow that takes a single rotational angiogram (RoCA), reconstructs the 3-dimensional arterial tree and applies computational fluid dynamics (CFD) to calculate the FFR without the need to induce hyperaemia or perform invasive pressure measurements.

Methods 20 patients, scheduled for elective PCI underwent RoCA. The FFR was measured with a Combowire (Volcano), under resting and hyperaemic conditions. Physiologically significant lesions were stented and the measurements repeated. The arterial anatomy was reconstructed on a Philips 3DCA workstation. Generic boundary conditions for CFD were derived from the measured data. The calculated (“virtual”) and measured FFR values were then compared.

Results There were 11 right coronary artery (RCA) cases (6 stented) and 12 left coronary artery (LCA) cases (5 stented). The anatomy was reconstructed, and the FFR computed in each case (pre- and post-stenting). The CFD model accurately predicted which lesions were physiologically significant (FFR <0.8) and which were not (FFR >0.8) in all cases. The virtual FFR values deviated from the measured by ±6% (SD=6%) for both RCA and LCA cases.

Conclusion We have developed a novel, user-friendly workflow, which has the potential to predict FFR without the need for invasive measurements or inducing hyperaemic conditions. Our model identified lesions requiring intervention in all cases. Further work will optimise and refine the model by better characterising the downstream generic boundary conditions. We aim to improve the accuracy of the optimised model with more complex patients and lesions.
Intravascular imaging modalities are used clinically

**Objectives** To determine the diagnostic accuracy of whole heart 3D myocardial perfusion CMR against invasively determined FFR.

**Methods** 55 patients referred for angiography underwent rest and adenosine stress 3D myocardial perfusion CMR at 3 Tesla (3D turbo gradient echo, flip angle 15, TR 2.0 ms/TE 1.0 ms, 12 slices of 5 mm thickness, in-plane resolution 2.3×2.3 mm², 10-fold k-space and time k-t broad linear speed up technique acceleration with k-t principal component analysis). Perfusion was scored visually as on a coronary territory basis on a score from 0 to 3. Ischaemic burden was calculated by quantitative segmentation of the volume of hypoenhancement. The FFR was measured in vessels with >50% severity stenosis. Fractional flow reserve <0.75 was considered haemodynamically significant.

**Results** Two patients were excluded (one due to claustrophobia, the other had poor image quality). From the remaining 55 patients and 159 coronary vessels, 64 underwent pressure wire assessment and 39 had an FFR <0.75. Sensitivity, specificity and diagnostic accuracy of CMR analysis per patient were 90%, 91% and 91%, respectively for the detection of significant coronary artery disease. By coronary territory the values were 79%, 92% and 85%.

**Conclusion** 3D CMR stress perfusion can detect functionally significant coronary artery disease with excellent sensitivity, specificity and predictive values when compared with FFR. 3D CMR perfusion imaging may offer an alternative method of detecting ischaemia for the purpose of guiding revascularisation and risk stratification.

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**AN EX-VIVO "WHOLE HUMAN HEART MODEL" FOR THE DEVELOPMENT OF INTRAVASCULAR IMAGING**

**Abstract 022 Figure 1** 3D CMR perfusion of a patient with a proximal LAD lesion with positive fractional flow reserve (FFR = 0.61). Note the subendocardial perfusion defect from the base towards the apex.

**Abstract 023 Figure 1**

**Background** Intravascular imaging modalities are used clinically to investigate ambiguous angiographic coronary lesions, guide and optimise stent deployment, and assess stent-related complications. Both intravascular ultrasound (IVUS) and optical coherence tomography (OCT) facilitate characterisation of plaque components, although a lack of adequate spatial resolution and depth of penetration, respectively, limit their clinical application. Adaptation of the existing technologies and novel techniques are in development but require further validation. We have developed an ex-vivo whole heart cadaveric model that facilitates multi-modality imaging and accurate comparison with histology.

**Methods** We have developed a model for fluoroscopic and invasive assessment of coronary arteries within "whole heart" cadaveric specimens. Hearts are provided by the West of England heart valve bio-bank, following harvesting of valves for homograft production. The coronary ostium is dissected and mobilised to allow cannulation with a modified 6F coronary guiding catheter (Abstract 023 figure 1A), secured with sutures. The cadaveric specimen is held within a purpose-built perspex container, with adaptors on both sides of the container's lid allowing connection of the guide catheter internally, and a Y-connector and pressure/injector manifold externally, see Abstract 023 figure 1 panel B. Cadaveric specimens undergo angiography (see Abstract 023 figure 1C,D), placement of a 0.014" guide wire and imaging catheter manipulation with the artery held at physiological temperature and pressure. In collaboration with the Department of Bioengineering, Erasmus MC, Rotterdam, we are using this model for assessment and validation of optical attenuation analysis as a tool to accurately delineate areas of macrophage infiltration, a major marker of plaque vulnerability. Optical attenuation governs the signal drop-off associated with tissue penetration. It is derived by fitting the following functional relation to the OCT data, \( I(z) = I_0 \exp(-\mu(z)) \), where \( \mu(z) \) is the local optical attenuation, the parameter of interest. The local signal intensity, \( I_0 \), is also a free parameter in the fit, but is currently not analysed. The data are fitted in windows of 200 μm length, after appropriate processing to reduce speckle noise.

**Results** Comparison of OCT, IVUS, and VH-IVUS against histology confirm the challenges in characterising plaque (Abstract 023 figure 2A–D; arrows indicate calcium). Analysis of optical attenuation appears to correlate with areas of macrophage infiltration (arrows in Abstract 023 figure 2E–H).

**Conclusions** Our ex-vivo whole heart cadaveric model facilitates accurate comparison of imaging modalities against histology. Developments in the imaging technologies are necessary to facilitate plaque characterisation as a clinical application. Optical attenuation may offer additional information regarding the macrophage content and “vulnerability” of plaque, validation work using our cadaveric model is ongoing.

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