

THE UNIVERSITY of EDINBURGH

### Edinburgh Research Explorer

## Computed tomography myocardial perfusion vs (15)O-water positron emission tomography and fractional flow reserve

#### Citation for published version:

Williams, MC, Mirsadraee, S, Dweck, MR, Weir, NW, Fletcher, A, Lucatelli, C, MacGillivray, T, Golay, SK, Cruden, NL, Henriksen, PA, Uren, N, McKillop, G, Lima, JAC, Reid, JH, van Beek, EJR, Patel, D & Newby, DE 2016, 'Computed tomography myocardial perfusion vs (15)O-water positron emission tomography and fractional flow reserve', *European Radiology*. https://doi.org/10.1007/s00330-016-4404-5, https://doi.org/10.1007/s00330-016-4404-5

#### **Digital Object Identifier (DOI):**

10.1007/s00330-016-4404-5 10.1007/s00330-016-4404-5

Link:

Link to publication record in Edinburgh Research Explorer

**Document Version:** Publisher's PDF, also known as Version of record

Published In: European Radiology

#### **Publisher Rights Statement:**

This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

#### **General rights**

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

#### Take down policy

The University of Édinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



#### CARDIAC



# Computed tomography myocardial perfusion vs <sup>15</sup>O-water positron emission tomography and fractional flow reserve

Michelle C. Williams<sup>1</sup> · Saeed Mirsadraee<sup>2</sup> · Marc R. Dweck<sup>1</sup> · Nicholas W. Weir<sup>2</sup> · Alison Fletcher<sup>2</sup> · Christophe Lucatelli<sup>2</sup> · Tom MacGillivray<sup>1,2</sup> · Saroj K. Golay<sup>1</sup> · Nicholas L. Cruden<sup>3</sup> · Peter A. Henriksen<sup>3</sup> · Neal Uren<sup>3</sup> · Graham McKillop<sup>4</sup> · João A. C. Lima<sup>5</sup> · John H. Reid<sup>2</sup> · Edwin J. R. van Beek<sup>1,2</sup> · Dilip Patel<sup>4</sup> · David E. Newby<sup>1,2</sup>

Received: 4 September 2015 / Revised: 5 May 2016 / Accepted: 12 May 2016 © The Author(s) 2016. This article is published with open access at Springerlink.com

#### Abstract

*Objectives* Computed tomography (CT) can perform comprehensive cardiac imaging. We compared CT coronary angiography (CTCA) and CT myocardial perfusion (CTP) with <sup>15</sup>O-water positron emission tomography (PET) and invasive coronary angiography (ICA) with fractional flow reserve (FFR). *Methods* 51 patients (63 (61–65) years, 80 % male) with known/suspected coronary artery disease (CAD) underwent 320-multidetector CTCA followed by "snapshot" adenosine stress CTP. Of these 22 underwent PET and 47 ICA/FFR. Obstructive CAD was defined as CTCA stenosis >50 % and CTP hypoperfusion, ICA stenosis >70 % or FFR <0.80.

*Results* PET hyperaemic myocardial blood flow (MBF) was lower in obstructive than non-obstructive territories defined by ICA/FFR (1.76 (1.32–2.20) vs 3.11 (2.44–3.79) mL/(g/min), P < 0.001) and CTCA/CTP (1.76 (1.32–2.20) vs 3.12 (2.44–

**Electronic supplementary material** The online version of this article (doi:10.1007/s00330-016-4404-5) contains supplementary material, which is available to authorized users.

Michelle C. Williams michelle.williams@ed.ac.uk

- <sup>1</sup> University of Edinburgh/British Heart Foundation Centre for Cardiovascular Science, Chancellor's Building, 49 Little France Crescent, Edinburgh, UK EH16 4SB
- <sup>2</sup> Clinical Research Imaging Centre, University of Edinburgh, Edinburgh, UK
- <sup>3</sup> Edinburgh Heart Centre, Royal Infirmary of Edinburgh, Edinburgh, UK
- <sup>4</sup> Department of Radiology, Royal Infirmary of Edinburgh, Edinburgh, UK
- <sup>5</sup> Departments of Medicine and Radiology, Johns Hopkins Hospital, Baltimore, MD, USA

3.79) mL/(g/min), P < 0.001). Baseline and hyperaemic CT attenuation density was lower in obstructive than non-obstructive territories (73 (71–76) vs 86 (84–88) HU, P < 0.001 and 101 (96–106) vs 111 (107–114) HU, P 0.001). PET hyperaemic MBF corrected for rate pressure product correlated with CT attenuation density (r = 0.579, P < 0.001). There was excellent per-patient sensitivity (96 %), specificity (85 %), negative predictive value (90 %) and positive predictive value (94 %) for CTCA/CTP vs ICA/FFR.

*Conclusion* CT myocardial attenuation density correlates with <sup>15</sup>O-water PET MBF. CTCA and CTP can accurately identify obstructive CAD.

Key Points

- CT myocardial perfusion can aid the assessment of suspected coronary artery disease.
- CT attenuation density from "snapshot" imaging is a marker of myocardial perfusion.
- CT myocardial attenuation density correlates with <sup>15</sup>O-water PET myocardial blood flow.
- *CT* attenuation density is lower in obstructive territories defined by invasive angiography.
- Diagnostic accuracy of CTCA+CTP is comparable to invasive angiography + fractional flow reserve.

Keywords Imaging · Perfusion · Ischaemia · Angiography · Angina

#### Abbreviations

BMI	Body mass index
CT	Computed tomography
CTCA	Computed tomography coronary angiography
CTP	Computed tomography myocardial perfusion
	imaging

ECG	Electrocardiogram
FFR	Fractional flow reserve
ICA	Invasive coronary angiography
MBF	Myocardial blood flow
MRI	Magnetic resonance imaging
PET	Positron emission tomography
SPECT	Single photon emission computed tomography

#### Introduction

Computed tomography coronary angiography (CTCA) can be performed at low radiation dose and has an excellent negative predictive value in the assessment of coronary artery disease (CAD) [1, 2]. However, areas of high density, such as calcification or stents, can reduce its accuracy. The concomitant assessment of myocardial perfusion could address this issue by providing additive functional information. Computed tomography myocardial perfusion (CTP) has the potential to improve the diagnostic accuracy of CTCA and can be performed during the same examination [3].

The selection of patients for coronary revascularization is based on the functional significance of stenoses. This is best characterised by fractional flow reserve (FFR) which is the current gold standard method of defining obstructive CAD and can lead to improved clinical outcomes when used to guide therapy [4]. However, FFR assessment requires invasive coronary angiography (ICA) and is a derived ratio of pressure rather than a direct measurement of blood flow. Other methods, such as magnetic resonance imaging (MRI) or single photon emission computed tomography (SPECT) can non-invasively assess myocardial perfusion. However, all investigations may be limited by spatial resolution, imaging artefacts, long imaging times and false negatives due to balanced ischaemia in three-vessel disease [5, 6].

Positron emission tomography (PET) using oxygen-15labeled water (<sup>15</sup>O-water) is the gold standard for the assessment of myocardial perfusion as it is a freely diffusible metabolically inert tracer, with a high extraction fraction that is independent of flow rate [7]. This tracer enables quantitative assessment of absolute myocardial blood flow, rather than relative myocardial blood flow identified by visual assessment in other techniques. <sup>15</sup>O-Water PET myocardial blood flow measurements have been validated with radiolabelled microspheres in animal models [8, 9] and compared with ICA, ICA/ FFR, SPECT, MRI and CTCA in human studies [10–14]. It has good intraobserver and interobserver variability as well as a low inter-scan variability for absolute myocardial blood flow [15, 16].

The diagnostic accuracy of CT myocardial perfusion imaging has been established in multi-centre studies as compared to SPECT [17] and SPECT+ICA [3]. In addition, there have been a number of single-centre studies comparing "snapshot" or "dynamic" CT myocardial perfusion imaging to a variety of gold standard assessments, including FFR [18, 19]. The acceptability of this non-invasive test to patients is also good [20]. However, snapshot CT myocardial perfusion imaging has not previously been assessed as compared to the physiological gold standard of <sup>15</sup>O-water PET myocardial blood flow. Furthermore, semi-quantitative assessment of CT myocardial perfusion imaging with measurements of attenuation density has not been assessed as compared to either clinical (FFR) or physiological (<sup>15</sup>O-water PET myocardial blood flow) gold standards.

This study assessed snapshot CT myocardial perfusion imaging (also known as single-shot, static and single-phase myocardial perfusion imaging), which obtains a small number of images at maximal contrast enhancement. In order to thoroughly assess this technique we compared it to both the physiological gold standard of absolute myocardial blood flow (measured with <sup>15</sup>O-water PET) and the clinical gold standard measure of coronary stenosis severity (FFR during ICA).

#### Methods

The study was conducted in accordance with the Declaration of Helsinki, with local research ethics committee approval and written informed consent of all participants.

#### **Study population**

Participants had known or suspected CAD and were due to undergo ICA for clinical indications. Exclusion criteria were renal failure (serum creatinine greater than 2.26 mg/dL or estimated glomerular filtration rate less than 30 mL/min), allergy to iodinated contrast, pregnancy or contraindication to adenosine. All patients who met the inclusion and exclusion criteria were invited to participate, and of these 19 % agreed. Cardiovascular risk was assessed using the Framingham 10year cardiovascular risk score.

#### Computed tomography

Participants underwent rest and adenosine stress computed tomography (CT) using a 320-multidetector scanner (Aquilion ONE, Toshiba Medical Systems, Japan). They were asked to refrain from caffeine for 12 h. Participants with a heart rate greater than 65/min were administered intravenous metoprolol (maximum 25 mg). Sublingual glyceryl trinitrate (300 µg tablet) was administered prior to, and removed immediately after, rest imaging.

After acquisition of scout images, patients underwent noncontrast wide volume CT using tube voltage 120 kV, tube current based on body mass index (BMI) and scan range from 2 cm below the carina to the base of the heart using volume sizes of 160, 140, 128, 120, 100 or 80 mm. Participants then underwent resting prospective wide volume CTCA. The detector coverage was based on non-contrast imaging and the acquisition window was based on heart rate (70-80 % of the RR interval for heart rates less than 60/min, 30-80 % of the RR interval for heart rates greater than 60/min or irregular heart rates). Tube voltage and current were based on BMI or scout image attenuation. A triphasic injection of iodinated intravenous contrast (iomeprol, 400 mg iodine/mL; Iomeron 400, Bracco, UK) was administered based on BMI (less than  $30 \text{ kg/m}^2$ , 50 mL;  $30-39 \text{ kg/m}^2$ , 60 mL; at least  $40 \text{ kg/m}^2$ , 70 mL). Bolus triggering started the scan when descending aorta enhancement reached 300 Hounsfield units (HU). Stress imaging was performed 10 min after rest imaging and after 4 min of adenosine (140  $\mu$ g/(kg/min)). The same tube voltage, tube current and contrast were used as for rest imaging. However, scan range was reduced to cover only the left ventricle, and tube current modulation obtained data over a single heartbeat. A further low dose CT (100 kV, targeted 75 % acquisition) was performed at 3 to 4 min to assess myocardial late enhancement.

Calcium scoring images were reconstructed with filtered back projection (Quantum Denoising Software, QDS+; kernel FC02), and CTCA and CTP images were reconstructed with iterative reconstruction (Adaptive Iterative Dose Reduction 3D, Toshiba Medical Systems, Japan; FC05 or FC03 kernels [21] respectively). From the rest CTCA acquisition, the scanner console software identified the "best phase" and reconstructed this phase and those 10 ms either side. Additional reconstructions at any point in the rest acquisition could be produced as required. For the stress CTCA, images were reconstructed every 5 % of the acquisition which covered the entire RR interval of a single heartbeat. Additional reconstructions at any point in the stress acquisition could be produced as required.

#### Oxygen-15-labelled water positron emission tomography

Rest and stress <sup>15</sup>O-water PET-CT was preformed using a hybrid scanner (128-multidetector Biograph mCT, Siemens Medical Systems, Germany). Attenuation correction CT was performed before rest or before rest and stress imaging. An on-site cyclotron (PETtrace 8, GE Healthcare, UK) produced <sup>15</sup>O, and a radiowater generator (Hydex Oy, Finland) produced the <sup>15</sup>O-water bolus. Rest imaging was performed using a target of 500 MBq <sup>15</sup>O-water (15 s bolus, 2-min saline flush). Dynamic acquisition was performed over 5 min (14 × 5, 3 × 10, 3 × 20 and 4 × 30 s). After suitable decay (approximately 10 min), stress imaging was performed with a further 500 MBq <sup>15</sup>O-water. Dynamic acquisition was performed during adenosine infusion. Dynamic emission images were reconstructed using the UltraHD algorithm (Siemens Medical Systems, Germany) (zoom 2, matrix  $128 \times 128$ , voxels  $3.18 \times 3.18 \times 3$  mm).

## Invasive coronary angiography and fractional flow reserve

ICA was performed via radial artery as per standard practice. FFR was assessed for major epicardial vessels with stenosis greater than 50 %, where technically possible. FFR is the ratio between distal coronary pressure and aortic pressure at maximal hyperaemia [4]. Pharmacological stress was induced using adenosine (140  $\mu$ g/(kg/min)). Obstructive CAD was defined as luminal stenosis of at least 70 % on ICA or FFR less than 0.80.

#### **Image analysis**

Assessment of imaging was blinded to results of other modalities. CT was analysed using a post-processing workstation (Vitrea fX, Vital Images, USA). Two trained observers (MCW, DEN) assessed images separately, and differences were agreed by consensus. The single phase with the fewest artefacts was used to assess CT rest and stress imaging. CTP was assessed as per standard guidelines with adjustment of viewing parameters as required [22]. Visual analysis of CTCA was based on the 19-segment model [1] and CTP on the 17-segment model [23]. Segmental data were consolidated into three territories for per vessel perfusion analysis [23]. Obstructive CAD was defined as stenosis greater than 50 % with corresponding hypoperfusion on CTP.

Semi-quantitative assessment of CTP images was performed by assessing CT myocardial attenuation density at rest and during maximal hyperaemia. CT images were assessed in the short axis, and the apex was excluded. Automatically applied myocardial contours were edited manually as required. Attenuation density was measured in each myocardial segment for endocardium, mid-wall, epicardium and full thickness. Transmyocardial perfusion ratio was calculated as the ratio of segmental subendocardial attenuation divided by mean subepicardial attenuation of the short-axis slice [24].

PET was analysed using dedicated software (Carimas 2.4, Finland) using a single tissue compartment model with correction for perfusable tissue fraction and spillover [15]. Images were reoriented into short-axis and myocardial contours defined on digital subtraction images automatically, with manual adjustment. Myocardial blood flow was analysed for the whole left ventricle and segmental data was consolidated into three territories for per-vessel analysis [23]. Coronary vasodilator reserve was defined as the ratio of peak hyperaemic to resting myocardial blood flow [25]. Total coronary resistance was calculated as mean arterial pressure divided by myocardial blood flow at baseline and hyperaemia [25]. Myocardial blood flow and CT attenuation density were

corrected for rate pressure product, calculated from heart rate and blood pressure during imaging.

#### Statistical analysis

Statistical analysis was performed using SPSS (version 18 for Mac, IBM). Normally distributed quantitative variables are presented with mean and 95 % confidence interval. Non-normally distributed data are presented with median and interquartile range. Statistical significance was assessed using analysis of variance, Student t or Mann–Whitney U tests as appropriate. Sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and diagnostic accuracy were calculated on a per-vessel and per-patient basis. Receiver operator characteristic curves were constructed to assess diagnostic accuracy. Correlations were assessed using Pearson's correlations. Inter- and intraobserver variability was assessed using kappa statistics. A statistically significant difference was defined as a two-sided P value less than 0.05.

#### Results

#### Demographics

CT was performed in 51 patients (Table 1). ICA was planned for all patients but only completed in 47 because of either patient or clinician preference. PET was performed in 22 patients who agreed to additional imaging and had not undergone revascularization in the intervening time period. All imaging was therefore performed prior to any revascularization procedure.

According to the Framingham 10-year cardiovascular risk score, 23 (45 %) were high risk, 20 (39 %) intermediate risk and 8 (16 %) low risk. Thirteen patients had previously experienced an acute coronary syndrome. Mean coronary artery calcium score was 870 (580, 1161) Agatston units. ICA/FFR identified obstructive CAD in 27/47 (57 %) patients and 51/ 141 (36 %) of vessels. Participants who underwent PET had similar demographic details and risk factors compared to the other participants (Supplementary Table i).

#### Radiation

Mean dose length product (DLP) for CT was 641.44 (558.50– 724.24) mGy cm. Using the 0.014 mSv/(mGy cm) conversion factor, we calculated the dose of rest CT as 2.73 mSv, stress CT 3.71 mSv and total CT protocol 8.98 mSv. Mean administered activity of <sup>15</sup>O-water was 474 (461–488) MBq for rest and 465 (448–482) MBq for stress imaging. Mean DLP of attenuation correction CT was 73 mGy cm. According to the conversion factors of 0.014 mSv/(mGy cm) and 1.1  $\mu$ Sv/MBq [26], this equates to a total effective dose of 2.05 mSv for PET.

Ν	51	
Age (years)	63	(61, 65)
Male/female	41/10	(80/20 %)
Weight (kg)	85.0	(80.0, 90.0)
BMI (kg/m <sup>2</sup> )	28.6	(27.3, 29.9)
Hypertension	36	(71)
Hypercholesterolaemia	48	(94)
Diabetes mellitus	5	(10)
Cerebrovascular disease	4	(8)
Peripheral vascular disease	2	(4)
Current smoker	9	(18)
Ex-smoker (>1 month)	23	(45)
Family history	24	(47)
Previous acute coronary syndrome	13	(25)
Previous revascularisation	13	(25)
Stent	12	(24)
Coronary artery bypass graft	1	(2)
Invasive coronary angiography		
Normal	18	(38)
One-vessel disease	14	(30)
Two-vessel disease	8	(17)
Three-vessel disease	7	(15)

Mean (95 % confidence interval) or number (%)

#### Assessment of myocardial perfusion

Tabla 1

Domographia

<sup>15</sup>O-Water PET demonstrated that the mean myocardial blood flow and myocardial blood flow corrected for rate pressure product for all segments was higher during hyperaemia (2.52 (2.40–2.64) and 2.65 (2.53–2.77) mL/(g/min) respectively) compared to baseline (0.80 (0.76–0.81) and 0.82 (0.79–0.85) mL/(g/min) respectively; P < 0.001 for both).

There were no differences in baseline <sup>15</sup>O-water PET myocardial blood flow between obstructive and non-obstructive lesions as determined by ICA/FFR (P = 0.74, Table 2, Fig. 1). However, during hyperaemia, <sup>15</sup>O-water PET myocardial blood flow was lower for obstructive lesions defined by ICA/FFR on both per-patient and per-vessel assessment (Table 2, Fig. 1).

There were no differences in baseline <sup>15</sup>O-water PET myocardial blood flow in territories supplied by arteries with obstructive or non-obstructive lesions as defined by CTCA/CTP on per-patient or per-vessel analysis (Table 2, Fig. 1). However, on per-segment analysis, baseline <sup>15</sup>O-water PET myocardial blood flow was lower in territories supplied by coronary arteries with obstructive lesions (0.65 versus 0.79 mL/(g/min), P < 0.001). Hyperaemic <sup>15</sup>O-water PET myocardial blood flow was lower in territories supplied by arteries with obstructive lesions defined by CTCA/CTP on per-patient, per-vessel and per-segment analysis (Table 2,

Table 2  $^{15}\mbox{O-Water PET}$  myocardial blood flow (mL/(g/min)) for obstructive and non-obstructive regions as defined by ICA/FFR and CTCA/CTP

		Non-obstructive		Obstructive		Р
ICA/FFR						
Per patient	Rest	0.88	(0.74, 1.03)	0.63	(0.35, 0.89)	0.074
	Stress	3.11	(2.44, 3.79)	1.76	(1.32, 2.20)	0.001
Per vessel	Rest	0.84	(0.77, 0.91)	0.78	(0.69, 0.88)	0.3
	Stress	2.92	(2.61, 3.23)	1.60	(1.28, 1.91)	< 0.001
CTCA/CTP						
Per patient	Rest	0.88	(0.74, 1.03)	0.63	(0.35, 0.90)	0.74
	Stress	3.12	(2.44, 3.79)	1.76	(1.32, 2.20)	0.001
Per vessel	Rest	0.86	(0.79, 0.94)	0.75	(0.67, 0.84)	0.45
	Stress	3.04	(2.70, 3.39)	1.78	(1.51, 2.05)	< 0.001
Per segment	Rest	0.79	(0.76, 0.82)	0.65	(0.61, 0.70)	< 0.001
	Stress	2.77	(2.63, 2.91)	2.09	(1.93, 2.24)	< 0.001

Mean (95 % confidence interval)

Fig. 1). In addition, vessel territories defined as normal by CTCA/CTP had a higher baseline and hyperaemic <sup>15</sup>O-water PET myocardial blood flow than those with non-obstructive or obstructive lesions, but this difference was not statistically significant (Fig. 2).

For per-vessel assessment, the optimal cut-off for <sup>15</sup>O-water PET myocardial blood flow for assessing the presence of obstructive stenosis as defined by either ICA/FFR or CTCA/CTP was 0.52 mL/(g/min) at baseline and 0.78 mL/ (g/min) during hyperaemia (Table 3, Supplementary Table ii). The area under the curve was higher for hyperaemic than baseline myocardial blood flow.

Mean CT myocardial attenuation density for all segments was 85 (83–87) HU at baseline and 111 (107–114) HU during hyperaemia (P < 0.01). The CT transmyocardial perfusion ratio (TPR) was higher at baseline than hyperaemia (1.68 (1.61–1.76) versus 1.17 (1.13–1.21), P < 0.001).

CT myocardial attenuation density was lower in territories supplied by arteries with obstructive as compared to nonobstructive lesions as defined by ICA/FFR (Table 4). <sup>15</sup>O-Water PET myocardial blood flow correlated with CT myocardial attenuation density (Table 5). The correlation was greater for the endocardium and when corrected for rate pressure product or left ventricular enhancement (Table 5). <sup>15</sup>O-Water PET myocardial blood flow corrected for rate pressure product correlated weakly with CT myocardial attenuation density at rest (r = 0.19, P < 0.001) but correlated well during hyperaemia (r = 0.579, P < 0.001, Fig. 1). CT transmyocardial perfusion ratio correlated only weakly with <sup>15</sup>O-water PET myocardial blood flow. However, <sup>15</sup>O-water PET defined coronary vasodilator reserve and total coronary resistance correlated more strongly with CT myocardial attenuation density.

#### Diagnostic accuracy of perfusion imaging

When compared to ICA/FFR, the sensitivity, specificity, PPV and NPV of CTCA/CTP were 96, 85, 90 and 94 % on a perpatient basis and 88, 83, 75 and 93 % on a per-vessel basis (Supplementary Table iii). Inter- and intraobserver variabilities for CT were good (kappa value 0.772, P < 0.001 and 0.721, P < 0.001) and for PET were excellent (kappa value 0.817, P < 0.001 and 0.942, P < 0.001). Figure 3 shows example images. Late enhancement was identified on one CT in a patient with a previous myocardial infarction, and this was confirmed on subsequent MRI. This shows that the majority of perfusion defects identified represent reversible ischaemia rather than infarction.

#### Discussion

We have assessed combined CT coronary angiography and snapshot CT myocardial perfusion imaging and have undertaken simultaneous comparisons with the gold standard measure of absolute myocardial blood flow (<sup>15</sup>O-water PET) and the gold standard measure of coronary stenosis severity (FFR during invasive coronary angiography). We have demonstrated for the first time that CT attenuation density correlates with <sup>15</sup>O-water PET myocardial blood flow. In addition, CT perfusion identified clinically significant CAD as defined by FFR with excellent diagnostic accuracy. Thus we provide physiological and clinical evidence to support the use of CT myocardial perfusion imaging.

CTCA is an established technique for the non-invasive assessment of CAD. However, CTCA alone fails to identify a significant proportion of functionally significant lesions compared with ICA/FFR [27]. This is where perfusion imaging has an important discriminatory role and is additive to CTCA [28]. CORE-320 showed that the CTP and CTCA has a sensitivity, specificity, PPV and NPV for detecting greater than 50 % ICA stenosis with SPECT perfusion defects of 80, 74, 65 and 86 % on a per-patient basis [3]. However, although SPECT is widely used as the first-line perfusion imaging technique by many clinicians, it has limitations [29, 30]. Comparisons with SPECT may under-represent the true value of CTP. We therefore used ICA and FFR as the gold standard clinical reference comparator and confirmed the diagnostic accuracy of CTCA/CTP.

Our study supports the clinical use of CTP in the investigation of patients with suspected CAD. CTP can be used as an optional "add-on" investigation immediately after CTCA when heavily calcified vessels are identified or if the functional severity of stenosis is uncertain. Alternatively, it can be prospectively planned for patients with CAD being considered for coronary revascularisation. In addition to identifying major epicardial CAD, CTP can also identify microvascular dysfunction.



**Fig. 1** Assessment of myocardial blood flow by <sup>15</sup>O-water positron emission tomography as compared to computed tomography coronary angiography (CTCA) and myocardial perfusion (CTP) and invasive coronary angiography (ICA) with fractional flow reserve (FFR). Baseline and hyperaemic myocardial blood flow in obstructive and

non-obstructive vessels as defined by  $\mathbf{a}$ ,  $\mathbf{b}$  ICA/FFR and  $\mathbf{c}$ ,  $\mathbf{d}$  CTCA/CTP.  $\mathbf{e}$ ,  $\mathbf{f}$  Correlation between myocardial blood flow corrected for rate pressure product and CT contrast enhancement at rest and during hyperaemia respectively

In our study, some of the false positives were due to the presence of microvascular dysfunction, highlighted as "slow flow" during ICA with a normal FFR. FFR is primarily focused on identifying a pressure gradient across a functional stenosis but cannot assess microvascular resistance or flow directly [31]. Other reasons for false positive results may include motion artefact or beam-hardening artefact on CT. A high specificity is an important aim for CT myocardial perfusion imaging to



Fig. 2 Baseline (a) and hyperaemic (b) myocardial blood flow assessed by <sup>15</sup>O-water positron emission tomography in normal, non-obstructive and obstructive vessels as defined by computed tomography coronary angiography and myocardial perfusion

enable this to be used to guide referrals for invasive coronary angiography. CT can be used to minimize the number of normal invasive coronary angiographies performed and also provide additional information on anatomy in order to plan revascularization. It is hoped that in the future this could also be used to plan bypass surgery in relevant patients, without the requirement for invasive coronary angiography.

In our study, the optimal cut-off to identify hemodynamically significant stenoses was 0.78 mL/(g/min). Previous studies have reported optimal cut-off values between 1.86 and 2.5 mL/(g/min) [11, 13, 14, 32, 33]. However, our study did not include healthy volunteers and had a high proportion of patients with CAD. Myocardial blood flow is known to be lower in normal regions remote from those with an obstructive stenosis in patients after acute myocardial infarction, as compared to normal controls [25]. Reduced coronary flow velocity reserve has also been identified in women with chest pain without obstructive coronary artery disease on invasive coronary angiography, possibly highlighting microvascular dysfunction [34]. A variety of doses of <sup>15</sup>O-water have been used in previous studies, including lower doses than used in this study [35, 36]. Although current guidelines recommend a 700–1500 MBq <sup>15</sup>O-water bolus [37], improved detector sensitivity in the latest generation scanners means that lower doses are appropriate, but may lead to variations in the results. Together, these factors may help explain the lower thresholds identified in our study. The higher <sup>15</sup>O-water PET myocardial blood flow identified in normal as compared to non-obstructive coronary arteries suggests the potential impact of microvascular disease on myocardial blood flow in patients with nonobstructive coronary artery disease. This difference was not statistically significant in our study but this does warrant further investigation.

This study assessed snapshot CTP where a small number of images are acquired just after the peak of contrast enhancement, rather than dynamic CTP that acquires multiple images during the wash-in and wash-out of contrast. Myocardial blood flow calculated using dynamic CTP correlates with <sup>15</sup>O-water PET myocardial blood flow [38]. However,

 Table 3
 Optimal cut-off value for hyperaemic myocardial blood flow (MBF) on per-vessel assessment to identify obstructive stenosis as defined by ICA/FFR or CTCA/CTP

		Cut-off value (mL/(g/min))	Area under the curve		Sensitivity (%)	Specificity (%)
			(95 % CI)	P value		
ICA/FFR	Hyperaemic MBF	0.78	0.897 (0.815, 0.979)	< 0.001	100	92
	CFR	1.02	0.880 (0.773, 0.986)	< 0.001	100	100
CTCA/CTP	Hyperaemic MBF	0.78	0.856 (0.761, 0.951)	< 0.001	100	95
	CFR	1.02	0.880 (0.773, 0.986)	0.02	100	94

 
 Table 4
 Myocardial contrast enhancement for obstructive and nonobstructive segments as defined by ICA and FFR

	Non-o	bstructive	Obstructive		Р		
Myocardial contrast enhancement (HU)							
Rest	86	(84, 88)	73	(71, 76)	< 0.001		
Stress	111	(107, 114)	101	(96, 106)	0.001		
Endocardiu	um contra	st enhancement	(HU)				
Rest	95	(93, 97)	84	(80, 87)	< 0.001		
Stress	115	(111, 118)	102	(98, 107)	< 0.001		
CT contrast corrected for left ventricle enhancement (HU)							
Rest	0.17	(0.17, 0.18)	0.15	(0.14, 0.16)	< 0.001		
Stress	0.37	(0.36, 0.39)	0.30	(0.28, 0.32)	< 0.001		
CT corrected for rate pressure product (HU)							
Rest	90	(86, 93)	81	(76, 86)	0.004		
Stress	124	(114, 134)	118	(108, 127)	0.330		
Transmyocardial perfusion ratio							
Rest	1.62	(1.54, 1.71)	1.91	(1.69, 2.13)	< 0.001		
Stress	1.23	(1.19, 1.28)	0.98	(0.93, 1.04)	< 0.001		

Mean (95 % confidence interval)

dynamic CTP has a higher radiation dose than snapshot CTP. Our study shows that semi-quantitative assessment of snapshot CTP using CT attenuation density also correlates with <sup>15</sup>O-water PET myocardial blood flow. Previous studies have suggested that rest images alone are sufficient to identify obstructive CAD [39]. However, our data do not support this as rest imaging alone was not sufficient to discriminate between obstructive and non-obstructive lesions.

The use of medications such as glyceryl trinitrate or betablockers prior to CTP could alter the presence or severity of perfusion defects [40]. However, previous studies have shown no difference in hyperaemic myocardial blood flow measured with <sup>13</sup>N-ammonia in patients taking beta-blockers [41]. In

Table 5Correlation betweenoxygen-15 PET myocardial bloodflow and computed tomographymyocardial perfusion imaging atrest and during adenosine stress:for PET parameters as comparedto CT myocardial attenuationdensity; and for CT parameters ascompared to oxygen-15 PETmyocardial blood flow correctedfor rate pressure product

addition the diagnostic accuracy of CTCA/CTP identified in our study was similar to previous studies where medications such as beta-blockers and nitrates were withheld. There is also significant heterogeneity in resting myocardial blood flow between individuals receiving these medications [10, 16, 33]. The number of patients included in this study was small and the overall prevalence of CAD was relatively high. Selection bias may also have been introduced as only 19 % consented to participate in this study. This means that assessment of the additive value of CTP over CTCA was difficult to assess. In addition FFR assessment was only performed in major epicardial vessels with stenosis greater than 50 % where technically possible, which may have led to misclassification. Indeed, it is not always possible in a clinical setting to perform FFR because of vessel size or tortuosity. This is an inherent limitation of this "gold standard", but nevertheless FFR is widely used clinically to guide patient management. Despite having high inter- and intraobserver variability [15] and good agreement between software packages [35], <sup>15</sup>O-water PET analysis has limitations including potential problems with bolus delivery, PET resolution, patient motion and suboptimal hyperaemia.

In conclusion, this study assessed CTP in comparison to <sup>15</sup>O-water PET myocardial blood flow and functional severity of coronary stenosis assessed by fractional flow reserve. We have shown that CTCA/CTP provides a robust physiological and clinical assessment of patients with suspected CAD, with excellent correlation with myocardial blood flow and comparable diagnostic accuracy to current invasive gold standard approaches of ICA and FFR. This is the first study to assess the semi-quantitative assessment of CT myocardial perfusion attenuation density as compared to <sup>15</sup>O-water PET. We believe that CTCA combined with CTP can now be performed in the clinical assessment of patients with suspected CAD.

	Baseline	Hyperaemia
PET parameters vs CT myocardial attenuation density		
PET total coronary resistance	0.278, <0.001	0.274, <0.001
PET myocardial blood flow	-0.227, <0.001	0.230, <0.001
Change in PET myocardial blood flow	0.307, <0.001	0.278, <0.001
PET coronary vasodilator reserve	0.411, <0.001	0.460, <0.001
PET myocardial blood flow corrected for rate pressure product	-0.235, <0.001	0.553, <0.001
CT parameter vs oxygen-15 PET myocardial blood flow		
CT attenuation density	-0.235, <0.001	0.553, <0.001
Endocardium CT attenuation density	0.328, <0.001	0.601, <0.001
CT attenuation density corrected for left ventricular enhancement	-0.136, 0.01	0.269, <0.001
CT attenuation density corrected for rate pressure product	0.19, <0.001	0.579, <0.001
Transmyocardial perfusion ratio	0.172, 0.0002	0.214, <0.001

r value, P value



Fig. 3 Example images from CT, PET and ICA. This 74-year-old nonsmoker with hypertension, hypercholesterolaemia and atypical chest pain underwent multimodality assessment of his cardiac anatomy and physiology. a Short-axis basal, mid and apical images of the left ventricle at rest and during hyperaemia along with three-dimensional representations of the transmyocardial perfusion ratio and coronary anatomy. Hypoattenutation is seen in right coronary artery (RCA) and left anterior descending (LAD) artery during hyperaemia. There is also mild hypoattenuation in the RCA territory on rest imaging. **b** Corresponding <sup>15</sup>O-water PET images which identified the same perfusion abnormalities in terms of the absolute myocardial blood flow (mL/(g/min)). **c** Corresponding images from ICA which identified an occluded LAD and severe stenosis of the RCA with FFR less than 0.80

**Acknowledgments** MCW, DEN and MRD are funded by the British Heart Foundation (FS/11/014, CH/09/002). This research was funded by the British Heart Foundation and Chest Heart and Stroke Scotland (R12/ A143). DEN is the recipient of a Wellcome Trust Senior Investigator Award (WT103782AIA). The Centre for Cardiovascular Science is the recipient of a British Heart Foundation Centre of Research Excellence Award (RE/08/001). The Clinical Research Imaging Centre and Wellcome Trust Clinical Research Facility (Edinburgh) are supported by National Health Service Research Scotland (NRS).

The scientific guarantor of this publication is Professor D E Newby.

The authors of this manuscript declare relationships with the following companies: MCW, DEN and EJHvB have spoken at meetings sponsored by Toshiba Medical Systems. JACL receives grant support only from Toshiba Medical Systems. No complex statistical methods were necessary for this paper. Institutional review board approval was obtained. Written informed consent was obtained from all subjects (patients) in this study. No study subjects or cohorts have been previously reported. Methodology: prospective, observational, performed at one institution.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http:// creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

#### References

- Miller JM, Rochitte CE, Dewey M et al (2008) Diagnostic performance of coronary angiography by 64-row CT. N Engl J Med 359: 2324–2336
- Menke J, Kowalski J (2015) Diagnostic accuracy and utility of coronary CT angiography with consideration of unevaluable results: a systematic review and multivariate Bayesian randomeffects meta-analysis with intention to diagnose. Eur Radiol 26: 451–458
- Rochitte CE, George RT, Chen MY et al (2014) Computed tomography angiography and perfusion to assess coronary artery stenosis causing perfusion defects by single photon emission computed tomography: the CORE320 study. Eur Heart J 35:1120–1130
- Tonino PAL, Fearon WF, De Bruyne B et al (2010) Angiographic versus functional severity of coronary artery stenoses in the FAME study. J Am Coll Cardiol 55:2816–2821
- Okada DR, Ghoshhajra BB, Blankstein R et al (2009) Direct comparison of rest and adenosine stress myocardial perfusion CT with rest and stress SPECT. J Nucl Cardiol 17:27–37
- Blankstein R, Shturman LD, Rogers IS et al (2009) Adenosineinduced stress myocardial perfusion imaging using dual-source cardiac computed tomography. J Am Coll Cardiol 54:1072–1084
- Bergmann SR, Fox KA, Rand AL et al (1984) Quantification of regional myocardial blood flow in vivo with H215O. Circulation 70:724–733
- Bol A, Melin JA, Vanoverschelde JL et al (1993) Direct comparison of [13N]ammonia and [15O]water estimates of perfusion with quantification of regional myocardial blood flow by microspheres. Circulation 87:512–525
- 9. Huang SC, Schwaiger M, Carson RE et al (1985) Quantitative measurement of myocardial blood flow with oxygen-15 water and positron computed tomography: an assessment of potential and problems. J Nucl Med 26:616–625

- Uren NG, De Bruyne B, Wijns W et al (1994) Relation between myocardial blood flow and the severity of coronary-artery stenosis. N Engl J Med 330:1782–1788
- Karamitsos TD, Leccisotti L, Arnold JR et al (2010) Relationship between regional myocardial oxygenation and perfusion in patients with coronary artery disease: insights from cardiovascular magnetic resonance and positron emission tomography. Circ Cardiovasc Imaging 3:32–40
- Schaefer WM, Nowak B, Kaiser H-J et al (2003) Comparison of microsphere-equivalent blood flow (15O-water PET) and relative perfusion (99mTc-tetrofosmin SPECT) in myocardium showing metabolism-perfusion mismatch. J Nucl Med 44:33–39
- Kajander S, Joutsiniemi E, Saraste M et al (2010) Cardiac positron emission tomography/computed tomography imaging accurately detects anatomically and functionally significant coronary artery disease. Circulation 122:603–613
- Danad I, Raijmakers PG, Appelman YE et al (2013) Hybrid imaging using quantitative H215O PET and CT-based coronary angiography for the detection of coronary artery disease. J Nucl Med 54: 55–63
- Nesterov SV, Han C, Mäki M et al (2009) Myocardial perfusion quantitation with 15O-labelled water PET: high reproducibility of the new cardiac analysis software (Carimas). Eur J Nucl Med Mol Imaging 36:1594–1602
- Iida H, Yokoyama I, Agostini D et al (2000) Quantitative assessment of regional myocardial blood flow using oxygen-15-labelled water and positron emission tomography: a multicentre evaluation in Japan. Eur J Nucl Med 27:192–201
- Cury RC, Kitt TM, Feaheny K et al (2015) A randomized, multicenter, multivendor study of myocardial perfusion imaging with regadenoson CT perfusion vs single photon emission CT. J Cardiovasc Comput Tomogr 9:103–112, e1–2
- Greif M, von Ziegler F, Bamberg F et al (2013) CT stress perfusion imaging for detection of haemodynamically relevant coronary stenosis as defined by FFR. Heart 99:1004–1011
- Choo KS, Hwangbo L, Kim JH et al (2013) Adenosine-stress lowdose single-scan CT myocardial perfusion imaging using a 128slice dual-source CT: a comparison with fractional flow reserve. Acta Radiol 54:389–395
- Feger S, Rief M, Zimmermann E et al (2015) Patient satisfaction with coronary CT angiography, myocardial CT perfusion, myocardial perfusion MRI, SPECT myocardial perfusion imaging and conventional coronary angiography. Eur Radiol 25:2115–2124
- Kitagawa K, George RT, Arbab-Zadeh A et al (2010) Characterization and correction of beam-hardening artifacts during dynamic volume CT assessment of myocardial perfusion. Radiology 256:111–118
- Mehra VC, Valdiviezo C, Arbab-Zadeh A et al (2011) A stepwise approach to the visual interpretation of CT-based myocardial perfusion. J Cardiovasc Comput Tomogr 5:357–369
- 23. Cerqueira MD, Weissman NJ, Dilsizian V et al (2002) Standardized myocardial segmentation and nomenclature for tomographic imaging of the heart. A statement for healthcare professionals from the Cardiac Imaging Committee of the Council on Clinical Cardiology of the American Heart Association 18:539–42
- 24. George RT, Arbab-Zadeh A, Miller JM et al (2009) Adenosine stress 64- and 256-row detector computed tomography angiography and perfusion imaging: a pilot study evaluating the transmural extent of perfusion abnormalities to predict atherosclerosis causing myocardial ischemia. Circ Cardiovasc Imaging 2:174–182
- Uren NG (1994) Reduced coronary vasodilator function in infarcted and normal myocardium after myocardial infarction. N Engl J Med 331:222–227
- 26. ICRP (2008) Radiation dose to patients from radiopharmaceuticals. ICRP Publication 106. Ann ICRP 38:1

- Meijboom WB, van Mieghem CAG, van Pelt N et al (2008) Comprehensive assessment of coronary artery stenoses. J Am Coll Cardiol 52:636–643
- Bettencourt N, Chiribiri A, Schuster A et al (2013) Direct comparison of cardiac magnetic resonance and multidetector computed tomography stress-rest perfusion imaging for detection of coronary artery disease. J Am Coll Cardiol 61:1099–1107
- Rocha-Filho JA, Blankstein R, Shturman LD et al (2010) Incremental value of adenosine-induced stress myocardial perfusion imaging with dual-source CT at cardiac CT angiography. Radiology 254:410–419
- Bettencourt N, Rocha J, Ferreira N et al (2011) Incremental value of an integrated adenosine stress-rest MDCT perfusion protocol for detection of obstructive coronary artery disease. J Cardiovasc Comput Tomogr 5:392–405
- Plein S, Motwani M (2013) Fractional flow reserve as the reference standard for myocardial perfusion studies: fool's gold? Eur Heart J Cardiovasc Imaging 14:1211–1213
- Kajander SA, Joutsiniemi E, Saraste M et al (2011) Clinical value of absolute quantification of myocardial perfusion with (15)O-water in coronary artery disease. Circ Cardiovasc Imaging 4:678–684
- Chareonthaitawee P, Kaufmann PA, Rimoldi O et al (2001) Heterogeneity of resting and hyperemic myocardial blood flow in healthy humans. Cardiovasc Res 50:151–161
- 34. Reis SE, Holubkov R, Conrad Smith AJ et al (2001) Coronary microvascular dysfunction is highly prevalent in women with chest

pain in the absence of coronary artery disease: results from the NHLBI WISE study. Am Heart J 141:735-741

- 35. Harms HJ, Nesterov SV, Nesterov SV et al (2014) Comparison of clinical non-commercial tools for automated quantification of myocardial blood flow using oxygen-15-labelled water PET/CT. Eur Heart J Cardiovasc Imaging 15:431–441
- 36. Lubberink M, Harms HJ, Halbmeijer R et al (2010) Low-dose quantitative myocardial blood flow imaging using 15O-water and PET without attenuation correction. J Nucl Med 51:575–580
- Hesse B, Tägil K, Cuocolo A et al (2005) EANM/ESC procedural guidelines for myocardial perfusion imaging in nuclear cardiology. Eur J Nucl Med Mol Imaging 32:855–897
- Kikuchi Y, Oyama-Manabe N, Naya M et al (2014) Quantification of myocardial blood flow using dynamic 320-row multi-detector CT as compared with <sup>15</sup>O-H<sub>2</sub>O PET. Eur Radiol 24:1547–1556
- 39. Osawa K, Miyoshi T, Koyama Y et al (2014) Additional diagnostic value of first-pass myocardial perfusion imaging without stress when combined with 64-row detector coronary CT angiography in patients with coronary artery disease. Heart 100:1008–1015
- Zoghbi GJ, Dorfman TA, Iskandrian AE (2008) The effects of medications on myocardial perfusion. J Am Coll Cardiol 52:401– 416
- 41. Hajjiri MM, Leavitt MB, Zheng H et al (2009) Comparison of positron emission tomography measurement of adenosine-stimulated absolute myocardial blood flow versus relative myocardial tracer content for physiological assessment of coronary artery stenosis severity and location. JACC Cardiovasc Imaging 2:751–758