Application of speCtraL computed tomogrAphy to impRove specIficity of cardiac compuTed tomogrAphyY (CLARITY study): rationale and design

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ABSTRACT

Introduction Anatomic stenosis evaluation on coronary CT angiography (CCTA) lacks specificity in indicating the functional significance of a stenosis. Recent developments in CT techniques (including dual-layer spectral detector CT [SDCT] and static stress CT perfusion [CTP]) and image analyses (including fractional flow reserve [FFR] derived from CCTA images [FFRct] and deep learning analysis [DL]) are potential strategies to increase the specificity of CCTA by combining both anatomical and functional information in one investigation. The aim of the current study is to assess the diagnostic performance of (combinations of) SDCT, CTP, FFRct, and DL for the identification of functionally significant coronary artery stenosis.

Methods and analysis Seventy-five patients aged 18 years and older with stable angina and known coronary artery disease and scheduled to undergo clinically indicated invasive FFR will be enrolled. All subjects will undergo the following SDCT scans: coronary calcium scoring, static stress CTP, rest CCTA and if indicated (history of myocardial infarction) a delayed enhancement scoring, static stress CTP, rest CCTA and if indicated invasive FFR will be enrolled. All subjects will undergo the following SDCT scans: coronary calcium scoring, static stress CTP, rest CCTA and if indicated (history of myocardial infarction) a delayed enhancement acquisition. Invasive FFR of ≤0.80, measured within 30 days after the SDCT scans, will be used as reference to identify a functionally significant stenosis. The primary study endpoint is the diagnostic performance of SDCT (including CTP) for the identification of functionally significant coronary artery stenosis. Secondary study endpoint is the diagnostic performance of SDCT, CTP, FFRct, and DL separately and combined for the identification of functionally significant coronary artery stenosis.

Ethics and dissemination Ethical approval was obtained. All subjects will provide written informed consent. Study findings will be disseminated through peer-reviewed conference presentations and journal publications.

Trial registration number NCT03139006; Pre-results.

INTRODUCTION

Anatomic evaluation of luminal stenosis on coronary CT angiography (CCTA) has high diagnostic performance to detect and rule out coronary artery disease.1–3 However, it lacks specificity in indicating the extent of myocardial ischaemia caused by a stenosis, especially in the presence of calcified lesions, due to volume artefacts caused by blooming.1–3 Invasively measured fractional flow reserve (FFR) is currently used to identify functionally significant coronary artery lesions and to guide revascularisation treatment.4 Results of the ISCHEMIA trial (NCT01471522) are awaited to shed light on the ongoing discussion concerning outcomes with revascularisation based only on ischaemia. Until these results are published, cardiac CT is the single modality able to simultaneously evaluate coronary artery anatomy and functional information in one investigation non-invasively. Recent developments in CT acquisition and image analysis techniques, such as dual-energy CT (DECT), static stress CT perfusion imaging (CTP), FFR derived from CCTA images (FFRct) and deep learning (DL) based image analysis, have shown the potential to improve the specificity of CCTA by combining both anatomical and functional information in one investigation.5–12 Dual-layer spectral detector CT (SDCT) is a DECT system with one X-ray tube and a dual-layer detector, which separately detects low and high-energy photons. In this way,
there is no time delay between the acquisitions of the different energy levels resulting in temporally coregistered dual-energy data. Also, the acquired images are spatiotemporally coregistered, making this DECT technique ideal for imaging a moving heart. In addition, by combining the information from both detector layers a conventional image can be reconstructed. Virtual monoenergetic images (40–200 keV) reconstructed from the SDCT may reduce blooming and beam-hardening artefacts and increase in-stent lumen visibility. A plaque-specific monoenergetic level can be applied to improve detection and quantification of coronary plaques. Together with the opportunity to assess myocardial iodine distribution and quantification in both rest and static stress images, SDCT may allow for improved anatomical and functional evaluation.

For FFR<sub>CT</sub>, the coronary lumen is segmented on CCTA images and assessed by algorithms based on fluid dynamics, lumped parameter models or machine learning. In this way, FFR<sub>CT</sub> is able to evaluate the functional significance of stenosis and has shown to improve diagnostic performance of CT. With SDCT, a plaque-specific monoenergetic level can be applied to improve lumen segmentation and the performance of FFR<sub>CT</sub>.

DL is a machine learning-based approach that has been shown to outperform other image analysis approaches in a wide variety of medical image analysis tasks. One of these tasks includes segmentation and detection analysis of cardiac CT. The growing amount of data generated by new CT acquisition techniques (eg, SDCT and CTP) makes DL analysis a technique of high interest. A recent study showed that patients with functionally significant coronary artery stenosis can be identified using DL analysis on CCTA. By combining these new technologies (SDCT, CTP, FFR<sub>CT</sub> and DL), we hypothesise more accurate information will be available about the coronary anatomy, degree of stenosis, FFR<sub>CT</sub> and myocardial perfusion, thereby contributing to a higher specificity of CT for identification of functionally significant coronary artery stenosis. To test this hypothesis, we have designed the CLARITY study (Application of spectrAimed tomogrAphy to impRove specIficity of cardiac compuTed tomographY). This paper describes the methods, rationale and design of the study.

**METHODS AND ANALYSIS**

**Study design**

The CLARITY study is a prospective monocentre diagnostic study designed to assess the accuracy of SDCT, CTP, FFR<sub>CT</sub> and DL-based image analysis as a non-invasive way for the identification of functionally significant coronary artery stenosis, using invasive FFR as the standard of reference. Combinations of multiple new CT acquisition techniques (ie, SDCT and CTP) and image analysis techniques (ie, FFR<sub>CT</sub> and DL) will be investigated to evaluate the optimal diagnostic performance of SDCT for the identification of functionally significant coronary artery stenosis.

**Study objectives**

The primary study objective is to assess the diagnostic performance of SDCT (including CTP) for the identification of functionally significant coronary artery stenosis in patients with stable angina and known coronary artery disease, using invasive FFR as the standard of reference in a patient-based analysis. The secondary objective is to assess the diagnostic performance of SDCT, CTP, FFR<sub>CT</sub> and DL separately and combined for the identification of functionally significant coronary artery stenosis in patients with stable angina and known coronary artery disease, using invasive FFR as the standard of reference in a vessel-based and patient-based analysis. The tertiary objective is to investigate if and to what extent SDCT can increase the degree of blooming and beam-hardening artefacts of calcifications.

Patients will only be included in the study after informed consent is signed.

**Targeted population**

Patients aged 18 years and older with stable angina and known coronary artery disease and scheduled to undergo clinically indicated invasive FFR are eligible for inclusion. Further inclusion and exclusion criteria are described in table 1. Due to the high negative predictive value of CCTA, patients are usually referred to CCTA when the pretest likelihood for obstructive coronary artery disease is <50%. Because CCTA has a moderate specificity in indicating the functional significance of a stenosis, it is not (yet) recommended for patients with pretest likelihood higher than 50%. Because we are interested in increasing the specificity of CCTA, we have chosen to include patients with stable angina and known coronary artery disease.

**Patient and public involvement**

Neither patients nor the public were involved in the design of the study, development of the research question and deciding on the outcome measures. At inclusion, patients are asked if they would like to receive results of the study at the group level. After completion of the study, results will be disseminated through a report to the patients who wish to receive information about the study.

**Sample size calculation**

To allow for evaluation of sensitivity and specificity within our study population, the proportions of cases (FFR ≤0.80) and controls (FFR >0.8) in our sample should take the prevalence of disease into account. Internal research of patient records of the past 3 years showed that 50% of the patients who underwent invasive FFR had at least one positive FFR measurement (FFR ≤0.80). With a prevalence of 0.5, the proportion of cases to controls is ≤0.80. With a prevalence of 0.5, the proportion of cases to controls is 1, indicating that the same amount of cases and controls is required. The main goal of the CLARITY study is to increase the specificity of SDCT (including CTP) for the
identification of functionally significant coronary artery stenosis. In the targeted population, the sensitivity and specificity of CCTA are currently around 95% and 50%, respectively. Using SDCT (including CTP), we aim to increase the specificity from 50% to 85%, raising the lower limit of the 95% CI for specificity from a typical value of 35% to 60%, necessitates inclusion of at least 66 patients (33 cases and 33 controls). However, this should not come at a too high cost in sensitivity. Using SDCT (including CTP), we expect a sensitivity around 90% with a lower limit of the 95% CI of 65%, which necessitates inclusion of at least 62 patients (31 cases and 31 controls). Because we are interested in both the sensitivity and specificity, we take the largest number of patients of the two calculated sample sizes (n=66). To account for attrition, 75 patients will be included. Our internal research also showed that in most cases FFR is measured in more than one vessel, with an average of 1.6 vessels per patient. Thus, for evaluation on a vessel basis, the amount of measurements is expected to be around 120 individual measurements.

**Patient recruitment and evaluation**

On fulfilment of the eligibility criteria, patients are asked for participation in the study by their treating physician. After signing informed consent, each study participant will undergo a dedicated CT protocol on an SDCT system before undergoing the clinically indicated invasive FFR measurement. Patients are asked to refrain from coffee consumption from 24 hours prior to imaging. For each patient, a case report form will be filled in, including information from the SDCT scans, invasive FFR measurement(s) and baseline characteristics such as age, gender, medical history, cardiovascular risk factors and demographic information.

**SDCT protocol**

All CT acquisitions will be performed using a 64-slice SDCT scanner (IQ on Spectral CT, Philips Healthcare, Best, The Netherlands) with 64×0.625 mm collimation. All patients will undergo coronary calcium scoring (CCS), adenosine-induced static stress CTP scan and rest CCTA scan. In case of a history of myocardial infarction, an additional late phase acquisition will be performed for myocardial scar detection. An overview of the scan protocol is shown in figure 1. All acquisitions will be performed according to dedicated guidelines. As contrast agent, 300 mg I/mL Iopromide (Ultravist 300, Bayer Healthcare, Berlin, Germany) will be used. For all scans, both conventional and spectral-based images will be reconstructed, which allows accessing DECT options. All images will be analysed using a dedicated workstation (IntelliSpace Portal, Philips Healthcare, Best, The Netherlands).

**Coronary calcium scoring**

Acquisitions will be made using a prospectively ECG-triggered (78% R-R interval) protocol (‘Step & Shoot Cardiac’) at 120 kVp and 40 mAs. Images will be reconstructed with 3 mm slice thickness and 3 mm increment using iterative reconstruction. Coronary calcifications will be quantified with Agatston scores using commercially available software (HeartBeat CS, IntelliSpace Portal, Philips Healthcare, Best, The Netherlands).

**Static stress CTP**

On completion of CCS, stress will be induced using intravenous adenosine injection (140 mg/kg/min). After 3 min of adenosine injection, a test bolus with 10 mL iodinated contrast will be given and flushed with 20 mL saline at a flow rate of 4.5 mL/s (<80 kg) or 5 mL/s (≥80 kg). From this test bolus, a patient-specific optimal scan delay is obtained by placing a region of interest in the ascending aorta to determine the time to peak, after which an additional 4 s are added. Subsequently, the contrast agent will be administered using 71 mL at 4.5 mL/s (<80 kg) or 85 mL at 5 mL/s (≥80 kg) and flushed with 50 mL saline at the same flow rate. After the predefined patient-specific optimal scan delay, and still under peak adenosine stress, a retrospectively ECG-gated single CTP scan is acquired in helical mode. The scan is obtained using 120 kVp with cardiac dose modulation for the mAs (reference 270 mAs), with highest mAs at end-systolic phase (45% R-R interval) and reduced mAs during the other phases. Directly after the acquisition, the adenosine injection will be stopped. Images will be reconstructed in steps of

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
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<tbody>
<tr>
<td>Age ≥18 years</td>
<td>Subjects who because of age, general medical or psychiatric condition or physiologic status cannot give valid informed consent or tolerate the coronary CTA examination</td>
</tr>
<tr>
<td>Stable angina and known coronary artery disease</td>
<td>Subjects with (severe) renal insufficiency, indicated as glomerular filtration rate (GFR) &lt;60 mL/min</td>
</tr>
<tr>
<td>Scheduled to undergo clinically indicated invasive FFR</td>
<td>Subjects with unknown GFR or obtained ≥3 months before the planned scan</td>
</tr>
<tr>
<td>Willing and able to give informed consent</td>
<td>Contraindication or allergy to intravenous iodinated contrast agent</td>
</tr>
</tbody>
</table>

Willing and able to give informed consent

Scheduled to give informed consent

Age ≥18 years

Scheduled to undergo clinically indicated invasive FFR

Subjects who because of age, general medical or psychiatric condition or physiologic status cannot give valid informed consent or tolerate the coronary CTA examination

Stable angina and known coronary artery disease

Scheduled to undergo clinically indicated invasive FFR

Willing and able to give informed consent

Subjects with (severe) renal insufficiency, indicated as glomerular filtration rate (GFR) <60 mL/min

Subjects with unknown GFR or obtained ≥3 months before the planned scan

Contraindication or allergy to intravenous iodinated contrast agent

Subjects who participate in another study with radiation which is estimated to be in risk category III (ICRP 103)

Subjects who are pregnant

Subjects with contraindications to cardiac CT and/or intravenous contrast, intravenous adenosine, beta-blockers or nitroglycerine

CTA, CT angiography; FFR, fractional flow reserve; ICRP, international commission on radiological protection.
10% from 0% to 90% of the R-R interval with 0.9 mm slice thickness and 0.45 mm increment using iterative reconstruction. The myocardium will be evaluated using 5 mm multiplanar reformatted images in short axis and long axis view. Myocardial perfusion will be scored visually and quantitatively on conventional images, monoenergetic images and iodine density images.

Coronary CT angiography
After a pause of 5–15 min a resting CCTA will be performed. In accordance with the Society of Cardiovascular CT (SCCT) guidelines, intravenous beta-blocker will be administered in patients with a heart rate above 60 beats per minute. In all patients, nitroglycerin 0.4 mg is administered sublingually. Iodinated contrast will be administered using the same amount and flow rate as for the stress scan. Seven seconds after a threshold of 100 Hounsfield Units is reached in the descending aorta, a CCTA scan is acquired using a prospectively ECG-triggered (78% R-R interval) protocol (‘Step & Shoot Cardiac’) at 120 kVp and 100 mAs. Images will be reconstructed with 0.9 mm slice thickness and 0.45 mm increment using iterative reconstruction. In accordance with SCCT guidelines, the images (both conventional and monoenergetic) will be reported and assessed on image quality, plaque characteristics and degree of stenosis (categorised as 0%, 1%–24%, 25%–49%, 50%–69%, 70%–99% and 100%). For anatomical evaluation of stenosis, ≥50% degree of stenosis will be used to indicate a significant stenosis. In addition to standard assessment, myocardial perfusion will be evaluated and compared with the static stress CTP.

Late phase non-contrast scan
In case a patient has a history of myocardial infarction an additional late phase non-contrast scan is made 5–10 min after the rest CCTA scan. This scan is acquired using a prospectively ECG-triggered (78% R-R interval) protocol (‘Step & Shoot Cardiac’) at 120 kVp and 100 mAs. Images will be reconstructed with 0.9 mm slice thickness and 0.45 mm increment using iterative reconstruction. The myocardium will be evaluated for late enhancement, both visually and quantitatively, using 5 mm multiplanar reformatted images in short axis and long axis view of the conventional, monoenergetic and iodine density images.

Radiation dose estimation
The estimated radiation dose for the CT scans will be calculated by multiplying the dose length product with the conversion coefficient for the chest at 120 kVp (k=0.0145 mSv/(mGy/cm)). The estimated radiation doses for CCS scan, static stress CTP, CCTA and late phase scan are 0.9, 2.5–4.4, 2.5 and 2.2 mSv, respectively (table 2). Cumulative radiation dose for the total SDCT protocol is estimated to be ≤10 mSv.

ICA and invasive FFR
For the clinical purpose, ICA with FFR measurement(s) will be performed in each patient within 30 days after the SDCT scan. Invasive FFR will be measured, under maximal hyperaemia (intravenous adenosine injection (140 µg/kg/min)), by placing a pressure wire (PressureWire X Guidewire, St. Jude Medical, St. Paul, Minnesota, USA) distal to the stenosis. The FFR value will be calculated automatically by dividing the pressure measured distally from the stenosis using the pressure wire by the pressure proximally from the stenosis assessed with the guiding catheter. To allow for comparison with FFRCT, the location of the invasive FFR measurement will be recorded. An FFR value of ≤0.80 will be used as reference standard to indicate a functionally significant stenosis. The FFR value will be blinded for the observers who will perform
CT or FFR CT readings. And the outcome of the SDCT and image analyses will be blinded for the observer who performs the invasive FFR.

**FFR derived from CTA images**

FFR CT analysis will be performed on resting CCTA images using a research prototype on-site FFR CT method (FFR CT, IntelliSpace, Philips Healthcare). Coronary artery centre-line and lumen will be automatically segmented and manually corrected by an expert. Subsequently, FFR CT will be automatically calculated and simulated using these segmentations. FFR CT will be performed on both conventional and monoenenergetic images.

**DL analyses**

Conventional and SDCT data generated from the different acquisitions will be used as input for the analysis of cardiac anatomy and function using DL. This analysis will include segmentation of anatomical structures of interest (eg, myocardium, coronary arteries), detection and characterisation of atherosclerotic plaque and perfusion defects and subsequent identification of patients with functionally significant stenosis.

**Study endpoints and data analysis**

The primary study endpoint is the diagnostic performance of SDCT (including CTP) for the identification of functionally significant coronary artery stenosis defined as FFR ≤0.8. Anatomical evaluation of coronary stenosis will be performed on conventional CCTA images. These data will then be compared with data obtained from SDCT as described above. The secondary study endpoint is the diagnostic performance of SDCT, CTP, FFR CT and DL separately and combined for the identification of functionally significant coronary artery stenosis defined as FFR ≤0.8. Diagnostic performance of anatomical evaluation of coronary stenosis, FFR CT, CTP and DL will be evaluated separately using conventional CT reconstruction. Subsequently, the diagnostic performance of each technology including the use of DECT options provided by SDCT will be assessed separately. In this way, the results of each technique with and without the use of SDCT options can be evaluated and the added value of SDCT for each technique can be defined. Hereafter, FFR CT, CTP and DL (including SDCT options) will be combined with anatomical information. First, the degree of stenosis will be evaluated. If ≥25% degree of stenosis is present on CCTA (+SDCT options), further testing using either FFR CT (+SDCT options), CTP (+SDCT options) or DL (+SDCT options) will be used to indicate a functionally significant stenosis (figure 2). The DL analysis will be performed on data generated from the different CT acquisitions. For combinations of acquisitions, the DL algorithm will automatically define to which extent the information of each acquisition is weighted in the model.

**Table 2** Radiation dose estimation

<table>
<thead>
<tr>
<th>Type of scan</th>
<th>CTDIvol</th>
<th>DLP*</th>
<th>Estimated radiation dose†</th>
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<tbody>
<tr>
<td>Coronary calcium scoring</td>
<td>5</td>
<td>63</td>
<td>0.9 mSv</td>
</tr>
<tr>
<td>Static stress CTP scan</td>
<td>14–24‡</td>
<td>175–300‡</td>
<td>2.5–4.4 mSv‡</td>
</tr>
<tr>
<td>Coronary CTA</td>
<td>14</td>
<td>175</td>
<td>2.5 mSv</td>
</tr>
<tr>
<td>Late phase non-contrast§</td>
<td>12</td>
<td>150</td>
<td>2.2 mSv</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td>5.9–10.0 mSv</td>
</tr>
</tbody>
</table>

*Calculated by CTDIvolx12.5 cm. †ICRP103 is used to calculate estimated radiation dose whereby the conversion coefficient of 0.0145 is multiplied by the DLP.‡ Depending on weight. §Performed in case of a history of myocardial infarction. CTA, CT angiography; CTP, CT perfusion; CTDIvol, volume CT dose index; DLP, dose length product; ICRP, international commission on radiological protection.

**Figure 2** Flow chart of degree of stenosis analysis combined with FFR CT, CTP and DL (including dual-energy CT options provided by SDCT). First, degree of stenosis will be evaluated. If ≥25% degree of stenosis is present, further testing using either FFR CT (+SDCT options), CTP (+SDCT options) or DL (+SDCT options) will be used to indicate a functionally significant stenosis. CTA, coronary CT angiography; CTP, CT perfusion; DL, deep learning; FFR CT, fractional flow reserve derived from coronary CT angiography images; SDCT, dual-layer spectral detector CT.
For both the primary and secondary objective, the accuracy, sensitivity, specificity, negative predictive value and positive predictive value will be calculated with 95% CIs on a per-patient level (primary and secondary endpoint) and per-vessel level (secondary endpoint). Differences in diagnostic performance will be investigated using pairwise McNemar tests and area under the receiver-operating curve analyses. A p<0.05 will be considered statistically significant.

The tertiary endpoint is an evaluation of blooming and beam-hardening artefacts of calcifications using SDCT compared with conventional CCTA. To determine the optimal virtual monoenergetic level at which calcium blooming and beam hardening is minimised and anatomy of the vessel is best visualised, degree of stenosis will be quantitatively scored at monoenergetic levels in steps of 10 (40–200 keV) and on conventional CCTA and compared with degree of stenosis at quantitative coronary angiography (QCA). The monoenergetic image with the least difference in the degree of stenosis compared with QCA is considered best. To determine the difference in blooming and beam-hardening artefacts of calcifications, volumes of circumscribed calcifications will be measured with dedicated volume analysis software (HeartBeat CS, IntelliSpace, Philips Healthcare). The percentage of volume measured on conventional CT minus the volume measured on the monoenergetic level with the least blooming (defined by the quantitative comparison with QCA) is considered as the degree of difference of blooming of calcifications. A paired t-test (normally distributed data) or Wilcoxon signed-rank test (non-normally distributed data) is used to assess the significance of the degree of decrease in blooming. Categorical variables will be expressed as counts and percentages. The Shapiro-Wilk test will be used to identify normally distributed data. Continuous variables will be expressed as mean±SD for normally distributed data and as median with IQR for non-parametric data.

**DISCUSSION**

The CLARITY study aims to evaluate the diagnostic performance of (combinations of) new CT acquisition techniques (SDCT and CTP) and image analysis techniques (FFR<sub>CT</sub> and DL) for the identification of functionally significant coronary artery stenosis. By analysing both anatomical and functional information, we hypothesise that the diagnostic performance of CT can be substantially improved. Multiple studies have demonstrated that assessment with FFR<sub>CT</sub> and image analysis exploiting DL individually lead to improved diagnostic performance. By combining these techniques with the assessment on monoenergetic and iodine density images, SDCT may allow for improved anatomical and functional evaluation. Each CT acquisition and image analysis technique has its own strengths and weaknesses. FFR<sub>CT</sub> can be performed on rest CCTA and thus does not require an additional scan. However, it can be challenging or even impossible in cases with high-density calcified plaque, a stent in the vessel of interest or in the presence of motion or misalignment artefacts. With the use of SDCT, some of these challenges may be overcome by reduction of blooming and beam-hardening artefacts. Stress CTP on SDCT is less or not influenced by these factors and may potentially be more representative of perfusion defects, as it is acquired during stress. On the other hand, stress CTP requires an additional scan and administration of adenosine, which increases radiation dose, risk and discomfort, which may be omitted if diagnostic performance is not superior to other techniques. DL-based image analysis may be able to gain insights into optimal combinations of information obtained from the different acquisitions. However, the number of patients in the current study is relatively low for DL-based analysis, which may need to be addressed by designing new approaches for sparsely sampled data. We expect that the highest diagnostic value can be found in a combination of diagnostic tests. The CLARITY study will be the first to investigate the combinations of SDCT, CTP, FFR<sub>CT</sub> and DL-based image analysis for the identification of functionally significant stenosis. Because the study is monocentric and only patients scheduled to undergo clinically indicated invasive angiography are included, a potential limitation is selection bias. Therefore, the results of the current study will be applicable to a relatively specific population with stable angina and known coronary artery disease. Future studies will be needed to translate the findings of the CLARITY study to a patient cohort with low to intermediate pretest likelihood for obstructive coronary artery disease, clinically more frequently encountered on CT.

**CONCLUSION**

The results of the CLARITY study will clarify to what extent SDCT, including CTP and image analysis techniques can improve the specificity for the identification of functionally significant coronary artery stenosis. This could potentially help with clinical decision-making and thereby decrease the number of patients unnecessary referred for invasive FFR.

**Status of study**

Patient enrolment started on 29 March 2017 and is anticipated to be completed before 1 August 2020. Until this moment, 21 patients are included in the study.

**Ethics and dissemination**

All subjects will provide written informed consent. Study findings will be disseminated through peer-reviewed conference presentations and journal publications.

**Contributors**

Design of the study: RWvH, II, PAdJ, MJW, MV and TL. Principal investigator: TL. Recruitment of patients: RWvH, MJMC, GEHL, MV and TL. Drafted study protocol and manuscript critically for important intellectual content: RWvH. Data collection, analysis and interpretation of the data: all authors. Revised study protocol and manuscript critically for important intellectual content: all authors. All authors have approved the final version of the manuscript.
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Disclaimer The authors are solely responsible for the design and conduct of the study, collection and analysis of the data, writing of the manuscript and its final contents.

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Patient consent for publication Not required.

Ethics approval The study protocol was reviewed and approved by the institutional review board of the University Medical Centre Utrecht (NL55917.041.16). In addition, the radiation safety committee approved this study.

Provenance and peer review Not commissioned; externally peer reviewed.

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