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Coronary Microvascular Function and Cardiovascular Risk Factors in Women With Angina Pectoris and No Obstructive Coronary Artery Disease: The iPOWER Study

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**Background**—The majority of women with angina-like chest pain have no obstructive coronary artery disease when evaluated with coronary angiography. Coronary microvascular dysfunction is a possible explanation and associated with a poor prognosis. This study evaluated the prevalence of coronary microvascular dysfunction and the association with symptoms, cardiovascular risk factors, psychosocial factors, and results from diagnostic stress testing.

**Methods and Results**—After screening 3568 women, 963 women with angina-like chest pain and a diagnostic coronary angiogram without significant coronary artery stenosis (<50%) were consecutively included. Mean age (SD) was 62.1 (9.7). Assessment included demographic and clinical data, blood samples, questionnaires, and transthoracic echocardiography during rest and high-dose dipyridamole (0.84 mg/kg) with measurement of coronary flow velocity reserve (CFVR) by Doppler examination of the left anterior descending coronary artery. CFVR was successfully measured in 919 (95%) women. Median (IQR) CFVR was 2.33 (1.98–2.76), and 241 (26%) had markedly impaired CFVR (<2). In multivariable regression analysis, predictors of impaired CFVR were age (P<0.01), hypertension (P=0.02), current smoking (P<0.01), elevated heart rate (P<0.01), and low high-density lipoprotein cholesterol (P=0.02), but these variables explained only a little of the CFVR variation (r²=0.09). CFVR was not associated with chest pain characteristics or results from diagnostic stress testing.

**Conclusion**—Impaired CFVR was detected in a substantial proportion, which suggests that coronary microvascular dysfunction plays a role in the development of angina pectoris. CFVR was associated with few cardiovascular risk factors, suggesting that CFVR is an independent parameter in the risk evaluation of these women. Symptom characteristics and results from stress testing did not identify individuals with impaired CFVR. (J Am Heart Assoc. 2016;5:e003064 doi: 10.1161/JAHA.115.003064)

**Key Words:** angina pectoris • coronary artery disease • echocardiography • microvascular dysfunction • women

More than half of women with angina-like chest pain referred for clinical coronary angiography (CAG) have no obstructive coronary artery disease (CAD), and this is twice as often seen in women compared with men after the CAG. While previously considered a benign condition, recent studies have found the condition to be associated with persistent chest pain, repeated angiograms, reduced quality of life, and increased cardiovascular morbidity and mortality. A possible explanation for the discrepancy between symptoms and CAG findings could be ischemia caused by coronary microvascular dysfunction (CMD). Recent studies have convincingly demonstrated that CMD is a strong predictor of cardiovascular prognosis, and CMD is common in both men and women with no obstructive CAD.
Previous studies of CMD among subjects with angina with no obstructive CAD have been based on relatively small, selected populations and little is known about the burden of CMD. CMD can be assessed as reduced coronary flow velocity reserve (CFVR) invasively during the CAG,5 by positron emission tomography7,8 or by transthoracic Doppler echocardiography (TTDE) of the left anterior descending coronary artery (LAD) during dipyridamole or adenosine infusion. With perhaps as little as 5% of patients with angina needing revascularization, as found in the PROspective Multicenter Imaging Study for Evaluation of Chest Pain (PROMISE) trial,9 the need for assessment of CMD is expanding and it is important to also evaluate CMD outside of the few dedicated centers that routinely perform invasive assessment of CMD during the CAG. Although there is an increasing interest in CMD, several gaps in knowledge remain. Thus, little is known about the burden of CMD among women with angina and no obstructive CAD and the correlation with symptoms, results of stress tests, and risk factors, since this has never been systematically assessed in an unselected population. Also, knowledge regarding the prognostic implications of CMD and effective treatment targeting both symptoms and prognosis are needed.

The iPowers study (ImProve diagnOsis and treatment of Women with angina pEctoris and micRovessel disease) aims to investigate diagnostic possibilities and prognosis of impaired CFVR in women with angina-like chest pain and no obstructive CAD.10 We investigated CMD prevalence measured with TTDE-assessed CFVR and the association with cardiovascular risk factors in women who were consecutively sampled after diagnostic invasive CAG. Further, we examined whether CMD was associated with characteristics, severity, and frequency of angina-like chest pain and results of diagnostic stress testing.

Methods
Population
Participants were recruited from the database PATS (Patient Analysis & Tracking System; Dendrite Clinical Systems), which covers eastern Denmark with \( \approx \)3 million inhabitants. All women (18–80 years) referred between March 2012 and September 2014 for a clinically indicated diagnostic CAG due to angina-like chest pain and suspected obstructive CAD were screened according to previously described well-defined inclusion and exclusion criteria (Figure 1).10 We included both women referred for stable angina and women hospitalized suspected of unstable angina, since the latter may be first manifestation of stable angina. Women with elevated cardiac markers or ST-segment elevation were excluded (Figure 1). All women were included within 1 year of their CAG.

Figure 1. Inclusion and exclusion criteria in the iPowers study.

Basic Examination
Basic assessment included clinical and demographic data. Trained health professionals interviewed participants regarding cardiac symptoms with respect to location, character, duration, radiation, frequency, and provoking and alleviating factors. According to the classical classification of chest pain symptoms were classified as typical angina pectoris, atypical angina pectoris and non-cardiac chest pain.11,12 Questionnaires regarding chest pain symptoms included the Seattle Angina Questionnaire, which evaluates 5 dimensions of functional status,13 and the World Health Organization’s Rose’s Angina Questionnaire, which evaluates symptoms as definite angina or not, further subdividing those with definite angina as severe or nonsevere.14

We obtained information regarding cardiovascular risk factors (age, body mass index [BMI], diabetes, hypertension, hyperlipidemia, smoking, family history of cardiovascular disease, and menopausal status), comorbidity, previous hospital admissions, and previous diagnostic tests, which included noninvasive cardiac computed tomography–angiography (CTA), exercise electrocardiography (ECG), and single-photon emission computed tomography (SPECT) performed within 6 months before CAG from interviews and charts. ECG, blood pressure, and heart rate measures were obtained at rest, and abdominal circumference was mea-
sured. The ECG was analyzed for signs of ischemia (inverted T wave, pathological Q wave, ST-segment depression), bundle-branch block, and atrial fibrillation. Blood samples were analyzed for cholesterol levels (total, low-density lipoprotein [LDL], and high-density lipoprotein [HDL] cholesterol), triglycerides, creatinine level, and glycated hemoglobin (HbA1c). A spot urine sample was analyzed for microalbuminuria.

Echocardiographic Examination

All participants underwent standard transthoracic echocardiography and TTDE of the LAD during rest and high-dose dipyridamole stress (0.84 mg/kg) over 6 minutes to obtain coronary flow velocities (CFVs) at baseline and at maximal hyperemia. Different vasodilators can be used to induce hyperemia, but the majority of TTDE studies of CFVR have used dipyridamole. Adenosine and dipyridamole are regarded as equal to achieve peak coronary vasodilation and are used interchangeably in clinical practice. Echocardiographic examinations were performed by using the GE Healthcare Vivid E9 cardiovascular ultrasound system (GE Healthcare) with a 1.3- to 4.0-MHz transducer (GE Vivid 5S probe) for standard echocardiography and a 2.7- to 8-MHz transducer (GE Vivid 6S probe) for TTDE. Images were stored for offline analysis (GE EchoPac v.112). The same 4 experienced echocardiographers performed all examinations in the same settings. Before examination, participants were instructed to be abstinent from caffeine or food containing significant amount of methylexanthine (coffee, tea, chocolate, cola, and banana) for 24 hours. Medication containing dipyridamole was paused for 48 hours; long-lasting nitroglycerin, β-blockers, angiotensin-converting enzyme inhibitors, angiotensin type II antagonists, calcium antagonists, and diuretics were paused for 24 hours; and short-lasting nitroglycerin was paused for 1 hour before the examination. Participants were studied in the left lateral decubitus position. The octave was set at 3.1/6.2 MHz, frequency at 8 MHz for B-mode (2D), while a baseline color scale between 1.00–2.50 kHz (velocity range ±10–24 cm/s) was chosen according to low or high flow velocities respectively. Color gain was adjusted to provide optimal 2-dimensional imaging quality. LAD was visualized with color Doppler in an apical modified foreshortened 2- or 4-chamber view or in a modified short-axis view of the left ventricle. CFV was measured with pulsed-wave Doppler as a laminar flow toward the transducer. We aimed to align the ultrasound beam direction to the LAD flow by adjusting probe position during recording of 2-dimensional and pulse-wave images. In case of difficulty in visualization of the LAD, a microbubble contrast agent was used (SonoVue; Bracco Imaging). Acquisitions of CFV during dipyridamole infusion were obtained throughout the infusion or up to 3 minutes after the infusion had terminated until flow had reached peak velocity. Blood pressure and heart rate were measured every 3 minutes during the examination of CFVR. After the examination, intravenous theophylline (maximum dose 220 mg) was administered to relieve potential side effects of dipyridamole.

For the analysis of CFVR, diastolic peak flow velocities were measured at rest and at peak hyperemia (Figure 2). CFVR was calculated as the ratio between peak velocities during stress and during rest. Two experts, blinded to participant data, analyzed every CFVR examination independently. The first reading was used, except for estimates that differed by >0.2, in which case the 2 analyzers reanalyzed the CFVR examination and reached agreement. In our previous validation study with repeated TTDE CFVR examinations in 10 young, healthy subjects by the same observer, we found an intraclass correlation coefficient of 0.97 (95% CI 0.92–1.00) and coefficient of variation of 7% (95% CI 3–10%) for repeat examinations. In a subsample of 50 participants from the iPOWER study, CFVR readings for the 2 observers were highly reproducible.

Left ventricular ejection fraction (LVEF) was analyzed by a skilled echocardiographer as an automated biplane calculation (Auto-EF tool; GE EchoPac v.112).

Statistical Analyses

Continuous variables with a Gaussian distribution are expressed as mean±SD (standard deviation) values. Median±IQR (interquartile range) values are used for variables with a non-Gaussian distribution. Count in percent
values is used for categorical variables. Distribution was assessed graphically. Difference between participants and nonparticipants was tested by using 1-way ANOVA or \( \chi^2 \) test for continuous and categorical variables, respectively. Missing values in the Seattle Angina Questionnaire were imputed according to the validated scoring system.\(^{13}\)

Participants were divided into 3 groups according to CFVR, based on current guidelines for determination of CMD by using a cutoff point of 2.0\(^{17}\) and a previously used cutoff point of 2.5.\(^{18}\) Age-adjusted trend tests by multivariable adjusted logistic or linear regression analysis were used to evaluate the distribution of variables (cardiovascular risk factors, clinical assessment, laboratory tests, results from diagnostic stress testing, medical history including medication, and psychosocial factors) among the 3 CFVR groups. Dependent variables with skewed distribution (smoking duration and menopause duration) were logarithmically transformed to base 2. Symptom characteristics from questionnaires were evaluated by using age-adjusted trend tests or \( \chi^2 \) if parameters of interest were divided into 3 categories. Moreover, to explore whether each symptom variable was a predictor of reduced CFVR, age-adjusted linear regression analyses were performed with natural logarithmically transformed CFVR as outcome variable because of non-Gaussian distribution. Interaction analysis was performed to investigate possible differences in association between participants regarded as having stable or unstable angina at the time of the CAG.

To explore predictors of reduced CFVR, multivariable linear regression analyses were performed with natural logarithmically transformed CFVR. All potential explanatory variables with an a priori defined hypothesis (age, hypertension, smoking status, diabetes, BMI, cholesterol, postmenopausal status, systolic blood pressure, resting heart rate, nonobstructive atherosclerosis at CAG, HDL, non-high-density lipoprotein cholesterol (non-HDL), and triglycerides) were tested in a prioritized order as determinants of CFVR and discarded at a cutoff level of \( P \geq 0.10 \). Assumptions of linearity, variance homogeneity, and Gaussian distribution of residuals were assessed graphically. The logarithmically transformed parameter estimates in the regression equation were converted back to the original scale for interpretation as an expected percentage change in CFVR value for each parameter by using the equation: \( 1-(e^{(\text{parameterestimate})}) \times 100\% \). Interaction analysis was performed to investigate difference in association between participants with stable or unstable angina. A likelihood ratio test was used to test difference between the final model and a model including interactions.

Confidence interval (CI) refer to 95% intervals, and a 2-sided \( P \) value \(<0.05\) was considered significant. All analyses were performed by using STATA/IC 13.1 (StataCorp LP).

**Ethics**

This study was performed in accordance with the Helsinki Declaration and was approved by the Danish Regional Committee on Biomedical Research Ethics (H-3-2012-005). All participants have given written informed consent on oral and written information.

**Results**

**Study Population**

Of the 5288 women with angina undergoing CAG in eastern Denmark between March 2012 and September 2014, 2159 were eligible for the study, 963 were included, and 919 had successfully measured CFVR (Figure 3). Of the included participants, 72% were categorized as having stable angina and 28% as having unstable angina at the time of CAG. Median interval (IQR) between diagnostic clinical CAG and CFVR examination was 71 days (51–97 days). A microbubble contrast agent (SonoVue; Bracco Imaging) was used in 59 (6%) participants. Almost all participants experienced side effects during the CFVR examination (98%), and on a visual analog scale from 1 to 10, the mean (SD) severity of symptoms reported by the participants was 5.7 (2.6). Two participants had an inherent atrial fibrillation induced by dipyridamole, and one experienced a delayed universal urticarial reaction. A higher proportion of nonparticipants had hypertension, diabetes mellitus, or nonobstructive atherosclerosis at CAG and stable angina pectoris as CAG indication, and more were currently smoking compared with participants (Table 1). This was similar when including only participants referred with stable angina.

**Characteristics of Participants With CMD**

Median (IQR) CFVR was 2.33 (1.98–2.76) and did not differ between participants with stable angina and those with unstable angina (\( P=0.89 \)). A total of 241 (26%) participants had a CFVR \( \leq 2 \), 318 (35%) had a CFVR between 2 and 2.5, and 360 (39%) had a CFVR >2.5. Table 2 displays characteristics of the study population by CFVR level. Participants with lower CFVR were significantly older. After age adjustment, participants with impaired CFVR had significantly more history of hypertension, diabetes mellitus, current smoking, elevated heart rate and more nonobstructive atherosclerosis at CAG. Participants with low CFVR also had significantly lower HDL cholesterol levels (\( P=0.02 \)) whereas CFVR was not associated with other serum lipid measures. Low CFVR was associated with a higher use of acetylsalicylic acid (\( P=0.02 \)), which was partly explained by a higher proportion with nonobstructive CAD. CFVR was not associated with cardiovascular medica-
tion such as β-blockers, calcium antagonists, angiotensin-converting enzyme inhibitors, or statins (Table 2), and there was no relation between CFVR and comorbidities such as musculoskeletal, pulmonary, thyroid, gastrointestinal, or gynecologic disease. There was no significant association between CFVR and results from a resting 12-lead study ECG (signs of ischemia, bundle-branch block, or atrial fibrillation). However, only 25 (3%) participants had atrial fibrillation. LVEF was not associated with impaired CFVR, but participants with LVEF <45% were excluded.

Baseline CFV correlated with CFVR ($r = 0.42, P < 0.001$), but CFV was not associated with the same cardiovascular risk

### Table 1. Background Characteristics on Included Participants and Nonparticipants

<table>
<thead>
<tr>
<th></th>
<th>Participants (n=963)</th>
<th>Nonparticipants (n=1196)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y, mean (SD)</td>
<td>62.1 (9.7)</td>
<td>62.7 (10.8)</td>
<td>0.19</td>
</tr>
<tr>
<td>Body mass index, kg/m², mean (SD)</td>
<td>27.3 (5.4)</td>
<td>27.4 (6.0)</td>
<td>0.71</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>487 (51)</td>
<td>598 (59)</td>
<td>0.001</td>
</tr>
<tr>
<td>Hyperlipidemia, n (%)</td>
<td>604 (63)</td>
<td>672 (66)</td>
<td>0.16</td>
</tr>
<tr>
<td>Diabetes mellitus, n (%)</td>
<td>127 (13)</td>
<td>177 (17)</td>
<td>0.02</td>
</tr>
<tr>
<td>Family history of CAD, n (%)</td>
<td>496 (53)</td>
<td>501 (51)</td>
<td>0.34</td>
</tr>
<tr>
<td>Smoking (current), n (%)</td>
<td>152 (16)</td>
<td>231 (22)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Stable angina pectoris, n (%)</td>
<td>693 (72)</td>
<td>918 (77)</td>
<td>0.01</td>
</tr>
<tr>
<td>Atherosclerosis at CAG, n (%)</td>
<td>335 (35)</td>
<td>513 (43)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Resident outside the capital region, n (%)</td>
<td>293 (30)</td>
<td>366 (31)</td>
<td>0.93</td>
</tr>
</tbody>
</table>

P value from 1-way ANOVA or $\chi^2$ test. CAD indicates coronary artery disease; CAG, coronary angiography.
factors as was CFVR. CFV was like CFVR associated with smoking ($P=0.004$) and heart rate ($P<0.001$). Cardiovascular risk factor associations with CFVR were similar for participants characterized as having stable ($n=693$) versus unstable angina ($P$ values for interaction $>0.05$).

### Determinants of CFVR

In multivariable regression analyses, CFVR remained associated with age, hypertension, smoking, resting heart rate, and HDL cholesterol in the final model (Table 3). However, the model explained only a minor part of the variation in CFVR ($r^2=0.09$). Adjusting for baseline CFV, which was strongly associated with CFVR, did not alter associations (results not shown). There was no significant interaction effect of stable versus unstable angina on the associations between cardiovascular risk factors and CFVR and no significant difference between our final model and a model including full interaction analysis ($P=0.54$).

When looking at smoking amount as a determinant of CFVR, adjusting only for age, CFVR decreased 4.6% (95% CI 2.0–7.2%) per 10 pack-year ([20 cigarettes/d]·10 y) for...
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**Table 3. Final Multivariable Regression Model in 885 Women**

<table>
<thead>
<tr>
<th></th>
<th>Expected Change of CFVR Value</th>
<th>95% CI</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (for 10 y of aging)</td>
<td>−6.2</td>
<td>−8.0 to −4.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Hypertension</td>
<td>−4.0</td>
<td>−7.2 to −0.8</td>
<td>0.016</td>
</tr>
<tr>
<td>Smoking</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Previous</td>
<td>−1.9</td>
<td>−5.2 to +1.6</td>
<td>0.29</td>
</tr>
<tr>
<td>Current</td>
<td>−8.6</td>
<td>−12.7 to −4.0</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Heart rate (for increase of 10 bpm)</td>
<td>−2.3</td>
<td>−3.8 to −0.8</td>
<td>0.002</td>
</tr>
<tr>
<td>High-density lipoprotein (per 1-mmol/L increase)</td>
<td>+4.1</td>
<td>+0.8 to +7.2</td>
<td>0.016</td>
</tr>
</tbody>
</table>

P value obtained by multivariable linear regression analyses with ln base logarithmic transformed coronary flow velocity reserve (CFVR) as outcome variable.

* Percent increase (indicated by +) or decrease (indicated by −) in percent per unit increase of independent variables.

Current smokers and 2.4% (95% CI 0.8–4.0%) per 10 pack-year ([20 cigarettes/d]·10 y) for previous smokers.

**Symptoms**

Of the participants, 471 (53%) had symptoms weekly and 306 (32%) had typical angina symptoms according to the classic characterization of chest pain.11,12 There was no association between CFVR level and symptom burden or symptom characteristics according to the classic classification of chest pain11,12 and Rose’s Angina Questionnaire. In addition, there was no association between CFVR level and angina frequency, angina stability, and treatment satisfaction evaluated by using the Seattle Angina Questionnaire, but participants with low CFVR had a significantly higher degree of physical limitation and a higher self-perception of disease as assessed by using the Seattle Angina Questionnaire (Figure 4). There was no association between impaired CFVR and whether angina pectoris occurred during rest, exertion, rest and exertion, or dipyridamole infusion. Further, we found no difference in number of hospital admissions or contacts with general practitioner (Table 4).

Among participants referred for stable angina, 317 (47%) had previously undergone an exercise ECG and 101 (15%) had undergone SPECT. A positive stress test (exercise ECG or SPECT) could not identify participants with CMD assessed by using TTDE (Table 2).

**Discussion**

In this large study in which we systematically investigated the association between symptoms, cardiovascular risk factors, and noninvasively assessed CMD in women with angina-like chest pain and no obstructive CAD, we found that CFVR was impaired in a large proportion and was associated with age, hypertension, current smoking, elevated resting heart rate, and low HDL cholesterol. No convincing correlation between severity or characterization of chest pain and impaired CFVR was found, and a positive diagnostic stress test did not identify participants with CMD.

The burden of symptoms was similar to that of previous studies of obstructive CAD. Prevalence of typical angina in our population was identical to that of women with obstructive CAD from the CONFIRM registry (coronary CT angiography evaluation for clinical outcomes: an international multicenter registry).19 In our study, 50% of all participants had angina-like chest pain at least once a week. For comparison, in the clinical outcomes utilizing revascularization and aggressive drug evaluation (COURAGE) study of subjects with obstructive CAD, ≈40% had angina at least once a month at 3-year follow-up.20 Participants in the COURAGE study also scored higher on all 5 elements in the Seattle Angina Questionnaire compared with the participants in the current study, indicating better function.20 Another study found that participants with nonobstructive atherosclerosis at CAG have more persistent angina compared with participants with obstructive CAD evaluated by using the Seattle Angina Questionnaire.3 Together, the findings indicate an overall limited association between symptoms and angiographic CAD severity. We further found that degree of CFVR impairment was not associated with symptom characteristics and that the classic characterization of chest pain could not be used to identify CMD.11 The results may question whether CMD is the cause of symptoms in these women. However, we have only...
assessed 1 dimension of CMD. In this present study, we have not assessed the endothelium-dependent epicardial dysfunction or mechanisms pertaining to pain perception, epicardial disease, or noncardiac causes of chest pain such as gastric pain, musculoskeletal disorders, or pulmonary disease. This might as well explain the lack of association between angina pectoris and CFVR. Studies simultaneously targeting symptoms and CMD are needed to clarify this.

We found that 26% of the women had CFVR <2 and, thus, CMD according to current guidelines.17 This is in agreement with several other studies. In a previous study of TTDE-assessed CFVR in 394 participants (48% men) with angina pectoris and no angiographic stenosis, 22% of participants had CMD, and further, CMD was associated with a hazard ratio of 16 for death or nonfatal myocardial infarction.6 Another study in 65 women with angina and no obstructive CAD found a TTDE-measured CFVR <2 in 40%.21 Other studies assessing CMD invasively or by positron emission tomography with different cutoffs have indicated that 39% to 54% have CMD.4,5,22 In a recent large study with invasive assessment of CMD in 1439 men and women with chest pain and nonobstructive CAD included over a period of 19 years, 30% had abnormal CFVR in response to adenosine.23

In the present study, the prevalence of risk factors and the use of medication were similar to results from another large Danish study of 2253 women with angina pectoris and no obstructive CAD.1 Prevalence of risk factors was also comparable to the National Heart, Lung, and Blood Institute Women’s Ischemia Syndrome Evaluation (WISE) study of women with chest pain and no signs of obstructive CAD. However, our population was 8 years older and fewer were current smokers.24 Overall, the prevalence of cardiovascular risk factors was relatively high compared with that of a general Danish population of women at a similar age.25

Table 4. Classification of Chest Pain Variables According to CFVR Level

<table>
<thead>
<tr>
<th>CFVR≤2.0 (n=241)</th>
<th>2&lt;CFVR≤2.5 (n=318)</th>
<th>CFVR&gt;2.5 (n=360)</th>
<th>P Value*</th>
<th>P Value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typical AP</td>
<td>71 (30)</td>
<td>102 (32)</td>
<td>122 (34)</td>
<td>0.17</td>
</tr>
<tr>
<td>Atypical AP</td>
<td>129 (54)</td>
<td>150 (47)</td>
<td>156 (43)</td>
<td></td>
</tr>
<tr>
<td>Noncardiac chest pain</td>
<td>41 (17)</td>
<td>66 (21)</td>
<td>82 (23)</td>
<td></td>
</tr>
<tr>
<td>Rose’s angina classification, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe definite AP</td>
<td>45 (19)</td>
<td>56 (18)</td>
<td>61 (17)</td>
<td>0.91</td>
</tr>
<tr>
<td>Nonsevere definite AP</td>
<td>56 (24)</td>
<td>73 (24)</td>
<td>94 (27)</td>
<td></td>
</tr>
<tr>
<td>Nondefinite AP</td>
<td>130 (56)</td>
<td>174 (57)</td>
<td>195 (56)</td>
<td></td>
</tr>
<tr>
<td>Physical limitation</td>
<td>71 (22)</td>
<td>73 (24)</td>
<td>77 (22)</td>
<td>0.02</td>
</tr>
<tr>
<td>Angina stability</td>
<td>63 (29)</td>
<td>62 (29)</td>
<td>65 (28)</td>
<td>0.17</td>
</tr>
<tr>
<td>Angina frequency</td>
<td>76 (23)</td>
<td>74 (24)</td>
<td>77 (22)</td>
<td>0.33</td>
</tr>
<tr>
<td>Treatment satisfaction</td>
<td>67 (25)</td>
<td>65 (24)</td>
<td>67 (24)</td>
<td>0.42</td>
</tr>
<tr>
<td>Perception/quality of life</td>
<td>48 (29)</td>
<td>50 (26)</td>
<td>51 (25)</td>
<td>0.04</td>
</tr>
<tr>
<td>Chest pain at exertion</td>
<td>41 (19)</td>
<td>50 (18)</td>
<td>51 (17)</td>
<td>0.88</td>
</tr>
<tr>
<td>Chest pain at rest</td>
<td>67 (32)</td>
<td>92 (33)</td>
<td>106 (36)</td>
<td></td>
</tr>
<tr>
<td>Chest pain at exertion and rest</td>
<td>104 (49)</td>
<td>137 (49)</td>
<td>139 (47)</td>
<td></td>
</tr>
<tr>
<td>Chest pain during dipyridamole infusion</td>
<td>70 (31)</td>
<td>112 (36)</td>
<td>119 (35)</td>
<td>0.91</td>
</tr>
<tr>
<td>Reproduced symptoms during dipyridamole infusion</td>
<td>68 (30)</td>
<td>96 (32)</td>
<td>95 (28)</td>
<td>0.22</td>
</tr>
<tr>
<td>Weekly chest discomfort</td>
<td>120 (53)</td>
<td>164 (56)</td>
<td>173 (51)</td>
<td>0.30</td>
</tr>
<tr>
<td>Previous hospitalization as a result of AP</td>
<td>1 (1)</td>
<td>1 (2)</td>
<td>1 (1)</td>
<td>0.60</td>
</tr>
<tr>
<td>Previous contacts with general practitioner as a result of AP</td>
<td>3 (4)</td>
<td>3 (4)</td>
<td>3 (3)</td>
<td>0.33</td>
</tr>
</tbody>
</table>

*P value from age-adjusted trend test (logistic or regression analyses) or chi-square test when symptom parameters of interest are divided into 3 categories.
†P value from age-adjusted linear regression analysis with natural logarithmically transformed coronary flow velocity reserve (CFVR) as outcome.
Reduced CFVR was associated with some, but not all, traditional cardiovascular risk factors. This was comparable to other studies that have found age, hypertension, diabetes mellitus, smoking status, resting heart rate, and HDL cholesterol to be determinants of CFVR, including in multiple adjusted models. One study in obese men and women with no obstructive CAD found impaired CFVR to be related to a high BMI. We did not find relations between CFVR and BMI, menopause or lack of hormonal therapy and CFVR in this large cohort and one reason may be that none of the mentioned studies adjusted for age, which is highly correlated to CFVR. However, the explanatory effect of the variation of CFVR in our final model was low, indicating that although associations are present, traditional cardiovascular risk factors account for little of the variation in CFVR, which is in accordance with previously published results from the National Heart, Lung, and Blood Institute WISE study and another recent large study.

It is often assumed that a positive stress test in the presence of no obstructive CAD is indicative of microvascular dysfunction. One study of 68 women found that significantly more women with low CFVR had a positive clinical stress test. This could not be corroborated in our study: among the 47% of participants who had a stress test performed, a positive stress test was not predictive of CMD. A recent large study included 1439 subjects with angina and invasive assessment of CMD also found no association between results of stress testing and CMD. The cause of the positive stress tests is unclear; however, it is plausible that CMD can cause ischemia without positive stress testing since these are mostly based on demonstration of regional rather than diffuse ischemia. Other cardiovascular indices might explain more of the variation in CFVR than traditional cardiovascular risk factors. Studies that include factors of vessel stiffness such as brachial–ankle pulse-wave velocity, augmentation index, and aortic pulse-wave velocity achieve a greater explanatory effect of the CFVR variation with $r^2$ values ranging from 0.36 to 0.52.

A full invasive assessment including both endothelial and nonendothelial mechanisms is regarded as the “gold standard” for assessing CMD. However, given that only a minority of patients with angina will ultimately need revascularization; noninvasive methods for assessment of CMD deserve wider application. Recent evidence has shown that CMD assessed by positron emission tomography among patients with no visual evidence of CAD on rest/stress positron emission tomography–myocardial perfusion imaging was a powerful predictor of major cardiovascular events with a hazard ratio of 0.8 per 10% increase in CFVR. Thus, noninvasive assessment of CMD may prove an important means of risk-stratifying this group. The present study demonstrates that this is feasible in the vast majority of unselected subjects and that the results are similar to those found by using invasive assessment. In addition, TTDE assessment of CMD is easily available given appropriate training and can be repeatedly assessed with no concern of radiation.

Strengths and Limitations
A main strength of this study is the systematic inclusion of participants who represent women with clinically assessed angina pectoris and no obstructive CAD in a region covering almost 3.0 million inhabitants. All women referred to CAG for angina were systematically screened and invited. Participants represent both the capital and rural areas. Nonparticipants had a greater burden of some cardiovascular risk factors and more nonobstructive atherosclerosis at CAG compared with participants, and this may have led us to underestimate the prevalence of CMD. The most common reason not to participate in the study was exhaustion and lack of energy, and this could also explain why nonparticipants had more cardiovascular risk factors. However, the internal validity is not likely to be affected by nonparticipants. Prevalence of CMD was comparable to previous studies, but more information on whether CFVR differs between clinically determined symptomatic and atypically symptomatic women or asymptomatic women not referred could have been addressed by including a matched control group of asymptomatic women. Future studies will address this.

We have assessed 1 dimension of CMD; the adenosine-induced reduced CFVR, which is mainly caused by dysfunction of the endothelium-independent vasodilation of the microcirculation. We have not assessed the endothelium-dependent epicardial dysfunction or microvascular spasm, which can be detected by using invasive acetylcholine provocation test. The latter might have been the main mechanism responsible for chest pain in a subgroup of participants included in this study, in particular those with unstable angina pectoris. Therefore, the burden of CMD might be underestimated in this study.

Conclusion
In the iPOWER study, CMD was systematically assessed by TTDE with high feasibility. One-third of the women had impaired CFVR and this was not associated with symptom characteristics or severity, or with results from diagnostic stress testing. CFVR was associated with only a few traditional cardiovascular risk factors. Further follow-up will determine whether assessment of CMD is useful to risk

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stratify the large population of women with angina and no obstructive CAD and to monitor effects of intervention.

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Disclosures

None.

References


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