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Advanced heart sound analysis as a new prognostic marker in stable coronary artery disease

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Aims

Recent technological advances enable diagnosing of obstructive coronary artery disease (CAD) from heart sound analysis with a high negative predictive value. However, the prognostic impact of this approach remains unknown. To investigate the prognostic value of heart sound analysis as two scores, the Acoustic-score and the CAD-score, in patients with suspected CAD which is treated according to standard of care.

Methods and results

Consecutive patients with angina symptoms referred for coronary computed tomography angiography (CTA) were enrolled. The Acoustic-score was developed from eight acoustic CAD-related features. This score was combined with risk factors to generate the CAD-score. A cut-off score >20 was pre-specified for both scores to indicate disease. If coronary CTA raised suspicion of obstructive CAD, patients were referred to invasive angiography and revascularized when indicated. Of 1675 enrolled patients, 1464 (87.4%) were included in this substudy. The combined primary endpoint was all-cause mortality and myocardial infarction (n = 26). Follow-up was 3.1 (2.7–3.4) years. Of patients with primary endpoints, the Acoustic-score was >20 in 25 (96%); the CAD-score was >20 in 22 (85%). In an unadjusted Cox analysis of the primary endpoints, the hazard ratio for scores >20 under current standard clinical care was 12.6 (1.7–93.2) for the Acoustic-score and 5.4 (1.9–15.7) for the CAD-score. The CAD-score contained prognostic information even after adjusting for lipid-lowering therapy initiation, stenosis at CTA, and early revascularization.

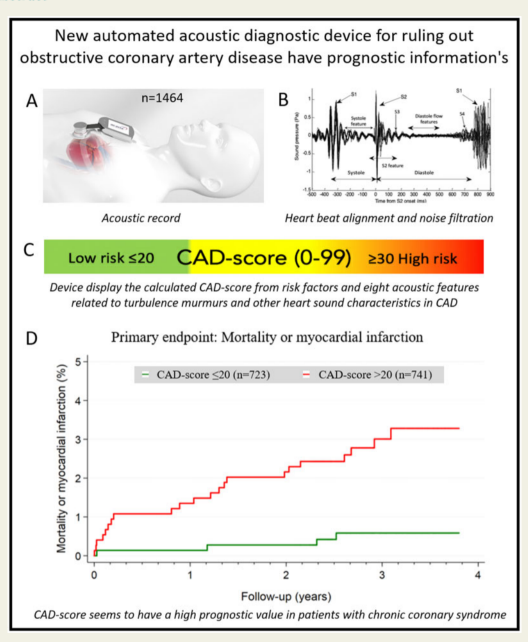
Conclusion

Heart sound analysis seems to carry prognostic information and may improve initial risk stratification of patients with suspected CAD.

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Graphical Abstract



Keywords

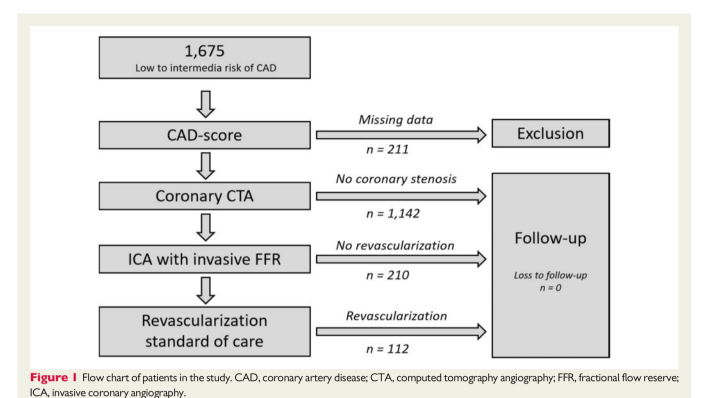
Prognosis • Coronary stenosis • Cardiovascular diagnostic technique • Heart sound • Acoustic cardiography

Introduction

Pre-test probability stratification of patients with suspected obstructive coronary artery disease (CAD) has been suboptimal for many years. Hence, most diagnostic tests to rule out CAD show normal conditions. $^{1-3}$

Acoustic devices measuring the heart sound originating from turbulent blood flow in the coronary circulation and altered myocardial relaxation have therefore been developed. These devices have been shown to be effective in ruling out CAD, and to outperform other methods by being faster, risk-free and more cost-effective.⁴

The diagnostic performance of two acoustic algorithms was recently tested in two large studies of patients presenting with symptoms suggestive of obstructive CAD.^{5–7} The studies demonstrated that these algorithms were positively correlated with the extent of CAD in terms of coronary artery calcium score, maximal diameter



stenosis, and number of vessels with disease. The studies demonstrated high sensitivities of 78–81%, moderate specificities of 35–53% but a high negative predictive value of 91–96%. It is therefore possible that these algorithms may serve as rule-out tests in patients with sus-

pected stable CAD.

However, no study has investigated whether heart sound carries any prognostic information. The acoustic features included in these acoustic algorithms have been linked to coronary stenosis degree, coronary flow volume and velocity, vascular stiffness, left ventricular diastolic, and systolic dysfunction—all markers which are predictive of a poor prognosis. ^{6,8–14}

The aim of this study was to investigate the prognostic value of pre-specified heart sound analysis (Acoustic-score) and an acoustic pre-test probability score (CAD-score) in patients with symptoms suggestive of CAD referred for coronary computed tomography angiography (CTA) and subsequently treated according to standard of care.

Materials and methods

Study design and population

The present predefined substudy used patients from the Danish study of the Non-Invasive Testing in Coronary Artery Disease (Dan-NICAD) trial 1. ^{15,16} Patients referred to coronary CTA due to symptoms suggestive of obstructive CAD were enrolled consecutively between September 2014 and March 2016. Patients were referred to coronary CTA after evaluation by a cardiologist in an outpatient cardiology unit. Decisions regarding referral to coronary CTA were based on patients' history, symptoms, risk profile, and echocardiographic findings according to national and

European Society of Cardiology guidelines. Exclusion criteria were: (i) age <40; (ii) previous known CAD; (iii) estimated glomerular filtration <40 mL/min; (iv) pregnancy; and (v) contra-indication for iodine-containing contrast medium, magnetic resonance imaging, or adenosine (severe asthma, advanced atrioventricular block, or critical aorta stenosis). All patients signed a written informed consent form. The Central Denmark Region Committees on Health Research Ethics, The Danish Medicines Agency, and The Danish Data Protection Agency approved the study, including the present 10-year follow-up substudy.

All included patients underwent a systematic interview to assess risk factors and symptoms and underwent coronary CTA. During their visit for coronary CTA, an examination with the acoustic microphone, CADScor®System (Acarix A/S, Denmark) was performed. Patients with suspicion of a coronary stenosis at coronary CTA all underwent invasive coronary angiography (ICA) with fractional flow reserve (ICA-FFR) measurement approximately 4 weeks after the coronary CTA. Patients without stenosis at coronary CTA did not undergo further testing. Follow-up data were obtained from electronic patient records and Danish national health registries (*Figure 1*).

The study design, imaging protocols, and analysis strategy have been published. 16

CAD-score

Heart sound recordings were obtained with the CADScor®System which is a small transportable electronic stethoscope device consisting of microphones and a micro-computer with a display. The microphone is mounted at the 4th intercostal space just to the left of the sternum with a dedicated adhesive patch during 5 min of rest. During the 3-min recording, the patient is asked to hold his/her breath for a period of 8 s, four times.

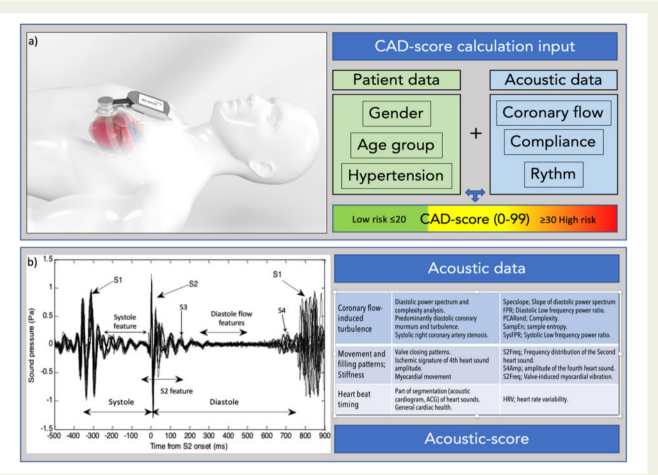


Figure 2 Depiction of the acoustic diagnostic method. (A) Coronary artery disease-score is calculated based on a 3-min recording session from the IC4-L region. Three patient risk factors are combined with acoustic derived data to generate a coronary artery disease-score. A score ≤20 is considered low risk and used to substantiate rule-out for coronary artery disease. (B) The acoustic data are derived from both the systolic and diastolic periods. The acoustic recording is segmented into discrete heart beats and aligned according to S2. Heart beats containing excess noise are filtered out, and the Acoustic-score is calculated. Several acoustic features having different, distinct coronary and myocardial origins are combined into the Acoustic-score.

Recordings were conducted in an undisturbed room. The CADScor®System analyses the quality of the recording and in case of poor quality, the device requests a second recording.

The CADScor®System algorithm Version 3 (V3) algorithm was utilized to calculate the Acoustic-score *per* se based on post-processing of audio recordings of the heart sound. In total, eight acoustic features related to turbulence murmurs and other heart sound characteristics in CAD were included in the Acoustic-score (*Figure 2*). The CAD-score V3 was generated by combining the Acoustic-score with the risk factors; gender, age, and hypertension (*Figure 2*).

In this paper, we present data from the Acoustic-score version 3 without risk factor modification and the CAD-score version 3. Both an Acoustic-score and a CAD-score value >20 were pre-specified as abnormal.⁶

Coronary computed tomography angiography

All patients in the Dan-NICAD trial underwent coronary CTA scans on a 320-slice volume CT scanner (Aquillion One, Toshiba Medical Systems, Japan) according to clinical guidelines.

CT imaging analysis included an Agatston calcium score and evaluation of CAD including luminal diameter stenosis estimation in each segment of the coronary tree using an 18-segment model. Coronary lesions were evaluated blinded to patient history. Stenosis severity was classified in all segments with a reference vessel diameter >2 mm. Severe stenosis was defined as 50–100% diameter reduction ($\approx\!70\%$ to 100% area reduction). Segments with suspected severe stenosis and non-evaluable segments with a reference vessel diameter >2 mm were defined as having obstructive CAD by coronary CTA and referred to ICA.

Invasive coronary angiography

Patients with suspected obstructive CAD at the coronary CTA were referred to ICA. ICA-FFR was measured in lesions with a visually estimated 30–90% diameter stenosis located in vessels with a reference diameter >2.0 mm using a clinical ICA-FFR system (Aeris, St. Jude Medical, Minnetonka, MN, USA). Haemodynamically obstructive CAD was identified in a blinded core lab as: (i) ICA-FFR value <0.80, (ii) luminal diameter stenosis reduction >90%, or (iii) luminal diameter stenosis reduction ≥50% if ICA-FFR was indicated but not performed for technical, anatomical, or other reasons.

Revascularizations were performed according to standard clinical practice after the ICA. Decisions were made blinded for CAD-score and other diagnostic test results.

Follow-up

Clinical endpoint data were extracted from patient records and Danish national health databases. Using the Civil Personal Registration number assigned to each Danish citizen at birth, we obtained information regarding mortality, diagnosed diseases, in-hospital procedures, and medical treatment. The CAD-score remained blinded to clinicians during follow-up. End of follow-up was set to 30 June 2018.

Information on mortality, myocardial infarction, and revascularization with either percutaneous coronary intervention (PCI) or coronary artery bypass graft (CABG) was obtained from the Western Denmark Heart Registry, the Danish National Patient Registry, and/or Danish Civil Personal Register. Data on Medical treatment and compliance were extracted from reimbursed medical prescriptions at Danish pharmacies through the Danish National Health Service Prescription Database. Changes in lipid lowering therapy after primary cardiac evaluation were defined as continued, discontinued, initiated, or no therapy according to both pre- and post-test-issued prescriptions within a window of 180 days before and 120 days after the examination.

Statistical analysis

The primary study endpoint was defined as a composite endpoint of all-cause mortality and myocardial infarction blinded for the first 120 days after coronary CTA. Secondary endpoints were early and late revascularization with a cut-off of 120 days with death as competing risk, and any coronary events blinded for the first 120 days after coronary CTA (all-cause mortality, myocardial infarction, and late revascularization). Patients who were revascularized early in relation to the baseline cardiac diagnostic evaluation were not excluded from the follow-up.

The primary aim was to compare scores with a cut-off value of 20, however for the CAD-score, we also considered the three groups: CAD-scores \leq 20 (reference), CAD-scores 21–29, and CAD-scores \geq 30, as well as an increment of the CAD-score of 10 units.

Time-to-event analysis was performed with univariate and multivariate Cox regression of the cause-specific hazard ratios (HRs). Cumulative incidence functions for each endpoint were generated to illustrate the risk over time. For Cox multiple regression analysis, we pre-specified models which included an updated Diamond–Forrester score, coronary disease severity at coronary CTA with stratification for change in medication, and revascularization as part of the Dan-NICAD trial's cardiac evaluation. However, *post hoc* we chose to develop several models with the endpoint of any cardiac event due to the low number of events. Cox multiple regression analysis was pre-specified to be blinded for the period of the first 120 days from baseline to enable adjustment of change in medication and revascularization as part of the baseline cardiac evaluation. In the blinded period, two patients died. They were the only ones excluded in the Cox multiple regression analysis.

For all statistical analyses, 95% confidence intervals (Cls) were reported when appropriate. The statistical analyses were performed using STATA-15.

Results

Of 1675 patients enrolled in the Dan-NICAD trial, 1464 (87.4%) were included in this study. In total, 211 (12.6%) were excluded due to missing CAD-score V3 (characteristics specified in Supplementary material online, *Table S1*). Baseline characteristics are listed in *Table 1*.

The median Acoustic-score was 24 (14–29). The Acoustic-score correlated vaguely with age, Spearman's rho 0.11 (P<0.001), and was higher in males than in females, 27 (21–32) vs. 21 (16–25) (P<0.001). A total of 488 (33%) patients had an Acoustic-score \leq 20, indicating low risk.

The median CAD-score was 20 (10–31). The CAD-score was positively correlated with age, Spearman's rho 0.50 (P<0.001), and was higher in males than in females: 28 (15–31) vs. 15 (7–24) (P<0.001), respectively. A total of 723 (49%) patients had a CAD-score \leq 20, indicating low risk. The CAD-score was positively correlated with the Acoustic-score, Spearman's rho 0.54 (P<0.001). Furthermore, 2 \times 2 tables for the scores are available in Supplementary material online, *Table* S2).

Diagnostic imaging characteristics of the baseline cardiac evaluation including echo, coronary CTA, ICA-FFR measurements, and revascularization related to the baseline cardiac evaluation, which were part of the Dan-NICAD trial, are summarized in *Table 2*. The median inter-test interval between coronary CTA and ICA was 30 days (10th and 90th percentiles: 14 and 50 days).

Treatments initiated at baseline

Lipid-lowering therapy was initiated based on baseline cardiac evaluation in 103 (14.3%) patients with a CAD-score \leq 20 and in 184 (24.8%) with a CAD-score \geq 20 (Supplementary material online, Figure S1). However, at baseline, plasma cholesterol levels were also higher in patients initiating lipid lowering therapy than in others, $5.8 \pm 1.1 \text{ vs.} 5.3 \pm 1.0 \text{ mmol/L}$, P < 0.001.

In total, 140 patients were diagnosed with haemodynamically obstructive CAD at ICA based on the baseline cardiac evaluation; and 112 patients were subsequently early revascularized according to standard clinical practice, 62 with PCI and 50 with CABG (secondary endpoint, *Tables 3* and 4). CAD-scores were significantly higher in the 112 revascularized patients than in the 1352 patients without revascularization related to the primary cardiac evaluation, 31 (21–40) vs. 19 (10–30) (P < 0.001), respectively. A detailed description of clinical information and imaging characteristics of the 26 patients who were revascularized and had a CAD-score \leq 20 is presented in Supplementary material online, *Table* S3.

Prognostic value of heart sound analysis

The combined primary endpoint occurred in 26 patients; 16 patients died and 10 patients had a myocardial infarction within a median follow-up time of 3.1 (2.7–3.4) years. In total, nine patients had late revascularization (>120 days after inclusion) (Supplementary material online, *Table S4*).

The isolated Acoustic-score without inclusion of risk factors was >20 in 25 out of 26 (96%) patients with the primary endpoint, mortality and myocardial infarction (*Table 3*). The unadjusted HR was 12.6 (1.7–93.2), P < 0.05. The HR of the Acoustic-score did not increase for Acoustics-scores \geq 30 compared with scores in the 21–29 range. Acoustic-score >20 had an HR for early revascularization within 120 days of 2.3 (1.4–3.6) but did not predict late revascularization (*Figure 3*, Supplementary material online, *Figure S2*).

The CAD-score was >20 in 22 of 26 (85%) patients with the combined primary endpoint—in 12 of 16 (75%) of patients who died, and in all 10 patients who had myocardial infarction. In addition, CAD-

Table I Baseline characteristic of included patient demographics (N = 1464)

	Total	CAD-score ≤20	CAD-score >20
Number of patients	1464	723 (49.3%)	741 (50.7%)
Characteristic		,	, ,
Race, Caucasian	1454 (99.3%)	717 (99.2%)	737 (99.5%)
Sex, male	716 (48.9%)	257 (35.6%)	459 (62.9%)
Age (years)	57.1 ± 8.7	53.3 ± 8.0	60.8 ± 7.8
Body mass index (kg/m²)	26.6 ± 4.1	26.3 ± 4.1	27.0 ± 4.1
Abdominal circumference (cm)	93.0 ± 12.6	90.6 ± 12.3	95.3 ± 12.5
Blood pressure			
Systolic	138 ± 19	129 ± 14	148 ± 18
Diastolic	83 ± 11	79 ± 10	86 ± 11
Heart rate ^a	65 ± 11	65 ± 11	66 ± 11
Smoking			
Never	697 (47.6%)	343 (47.4%)	354 (47.8%)
Former	536 (36.6%)	245 (33.9%)	291 (39.3%)
Active	231 (15.8%)	135 (18.7%)	96 (13.0%)
Diabetes	79 (5.4%)	24 (3.3%)	55 (7.4%)
Symptoms			
Typical chest pain	407 (27.8%)	203 (28.1%)	204 (27.5%)
Atypical chest pain	493 (33.7%)	259 (35.8%)	234 (31.6%)
Non-specific	564 (38.5%)	261 (36.1%)	303 (40.9%)
Updated Diamond Forrester score	39% (20–54%)	28% (16–45%)	42% (29–59%)
Low risk (<15%)	210 (14.3%)	168 (23.2%)	42 (5.7%)
Moderate risk (≥15% to 85%)	1220 (83.3%)	555 (76.8%)	665 (89.7%)
High risk (≥85%)	34 (2.3%)	0 (0.0%)	34 (4.6%)
Medication			
Hypertension	491 (33.8%)	86 (12.1%)	405 (45.9%)
Mean number of antihypertensives	1.9 ± 1.0	1.9 ± 0.9	1.8 ± 1.0
Hypercholesterolaemia	398 (27.2%)	145 (20.0%)	253 (34.1%)
Biochemistry ^b			
Cholesterol (total, mmol/L)	5.4 ± 1.1	5.4 ± 1.1	5.3 ± 1.1
HbA1c (mmol/mol)	37.7 ± 7.2	37.1 ± 7.0	38.3 ± 7.2
Creatinine (mmol/L)	75.4 ± 14.6	72.4 ± 12.8	78.3 ± 15.6

Values are n (%) or mean \pm SD or median (IQR).

score was >20 in all patients who underwent late revascularization (>120 days after inclusion) (*Table 4*).

In an unadjusted Cox regression analysis of the combined primary endpoint, CAD-scores >20 had an HR of 5.4 (1.9–15.7), P < 0.01. The HR increased significantly with higher CAD-score; with CAD-scores ≤ 20 as reference, CAD-scores 21–29 and CAD-scores ≥ 30 had a HR of 3.0 (0.9–10.8), P = 0.09, and 7.7 (2.6–22.9), P < 0.001, respectively (Figure 4 and Supplementary material online, Figure \$2).

It was not possible to calculate the HR for late revascularization as there were no events in patients with CAD-scores \leq 20. However, the log-rank test for late revascularization was significant, P < 0.001 (Figure 4).

In Cox regression analysis of any cardiac events after 120 days (secondary endpoint: death, myocardial infarction, or revascularization >120 days), a CAD-score increment of 10 units had an HR of 1.8

(1.3-2.5) P < 0.01, which was not influenced by adjustment for sex and age. In contrast to the updated Diamond–Forrester score, the CAD-score remained significant after including the updated Diamond–Forrester score in the model. Finally, the CAD-score remained significantly associated with events after adjusting for (i) coronary stenosis at coronary CTA and (ii) lipid lowering therapy initiation and revascularization (*Table 5*).

In a stratified analysis of patients not revascularized as part of the baseline cardiac evaluation (n = 1352), a CAD-score >20 had an HR of 3.9 (1.1–13.9), P < 0.05, and a CAD-score increase of 10 units had an HR of 1.5 (1.0–2.3), P = 0.05, for events after 120 days (dead, myocardial infarction or revascularization >120 days). Similarly, in patients (n = 112) who were revascularized, all patients who died or had myocardial infarction or revascularization >120 days had a CAD-score >20, and a CAD-score increment of 10 units had an HR of 1.5 (0.8–2.6), P = 0.17, for events after 120 days.

^aMean heart rate at the time of CAD-score measurement was: 54 ± 7 b.p.m. and at the time of CTA: 56 ± 7 b.p.m.

^bData available in: cholesterol 94%, glucose fasting 8%, HbA1c 78%, and creatinine 99% of the cohort.

Table 2 Imaging study characteristics from the baseline cardiac evaluation which were protocolized in the Dan-NICAD trial

	Total	CAD-score ≤20	CAD-score >20
Number of patients	1464	723 (49.3%)	741 (50.7%)
Echo (n = 1438)			
Left ventricular ejection fraction	59.9% ± 3.4	60.0% ± 2.6	59.5% ± 4.0
Cardiac valve disease, any	69 (4.7%)	23 (3.2%)	46 (6.2%)
Coronary artery calcium score (n = 1459)			
Median	0 (0–81)	0 (0–13)	25 (0–192)
Coronary artery calcium score (CACS) groups			
None (CACS = 0)	747 (51.2%)	472 (65.7%)	275 (37.2%)
Low/moderate (CACS: 1–399)	564 (38.7%)	219 (30.5%)	345 (46.6%)
High (CACS ≥400)	148 (10.1%)	28 (3.9%)	120 (16.2%)
Cardiac computed tomography angiography ($n = 1457$)			
Coronary artery disease severity			
Non (stenosis 0% and CACS = 0)	696 (47.8%)	449 (62.5%)	247 (33.5%)
Mild (stenosis 0–30%)	306 (21.0%)	126 (17.5%)	180 (24.4%)
Moderate (stenosis 30–50%)	111 (7.6%)	51 (7.1%)	60 (8.1%)
Severe (stenosis 50–100%)	344 (23.6%)	93 (12.9%)	251 (34.0%)
Radiation dose per coronary CTA (mSv)	2.7 ± 1.7	2.4 ± 1.5	2.9 ± 1.8
Invasive coronary angiography with fractional flow reserve ($(n = 321)^a$		
Coronary artery disease severity			
Obstructive stenosis	140 (9.7%)	27 (3.8%)	113 (15.6%)
Number of coronary vessels with severe stenosis			
One-vessel disease	85 (5.9%)	16 (2.3%)	69 (9.5%)
Two-vessel disease	37 (2.6%)	8 (1.1%)	29 (4.0%)
Three-vessel disease or left main disease	18 (1.3%)	3 (0.4%)	15 (2.1%)

Values are n (%) or mean \pm SD or median (IQR).

CACS, Coronary artery calcium score.

^aICA were only indicated in patient with severe stenosis at coronary CTA.

Table 3 Tables of events during follow-up according to the Acoustic-score

	Total	Acoustic-score ≤20	Acoustic-score >20
Number of patients	1464	448 (33.3%)	667 (66.7%)
Primary endpoint			
Mortality and myocardial infarction	26 (1.8%)	1 (0.2%)	25 (2.6%)
Secondary endpoints			
Mortality, all-cause	16 (1.1%)	1 (0.2%)	15 (1.5%)
Myocardial infarction	10 (0.7%)	0 (0.0%)	10 (1.0%)
Procedure related/spontaneous	2/8	0/0	2/8
Early/late (cut-off 120 days)	7/3	0/0	7/3
Early revascularization (≤120 days)	115 (7.9%)	21 (4.3%)	94 (9.6%)
Part of the primary Dan-NICAD evaluation	112	21	91
Acute	3	0	3
Late revascularization (>120 days)	9 (0.6%)	3 (0.6%)	6 (0.6%)

Values are n (%).

In a 'worst case scenario' simulation, an event was assigned to all patients with early revascularization as part of the primary baseline cardiac evaluation and with a false negative CAD-score \leq 20 (n=26).

In this scenario, CAD-scores >20 had similarly 5-year events rates, for any mortality, myocardial infarction, late revascularization not part of the primary baseline evaluation, and early revascularization as

Tables of events during follow-up according to the CAD-score Table 4

	Total	CAD-score ≤20	CAD-score >20
Number of patients	1464	723 (49.3%)	741 (50.7%)
Primary endpoint			
Mortality and myocardial infarction	26 (1.8%)	4 (0.6%)	22 (3.0%)
Secondary endpoints			
Mortality, all-cause	16 (1.1%)	4 (0.6%)	12 (1.6%)
Myocardial infarction	10 (0.7%)	0 (0.0%)	10 (1.4%)
Procedure related/spontaneous	2/8	0/0	2/8
Early/late (cut-off 120 days)	7/3	0/0	7/3
Early revascularization (≤120 days)	115 (7.9%)	26 (3.6%)	89 (12.0%)
Part of the primary Dan-NICAD evaluation	112	26	86
Acute	3	0	3
Late revascularization (>120 days)	9 (0.6%)	0 (0.0%)	9 (1.2%)

Values are n (%).

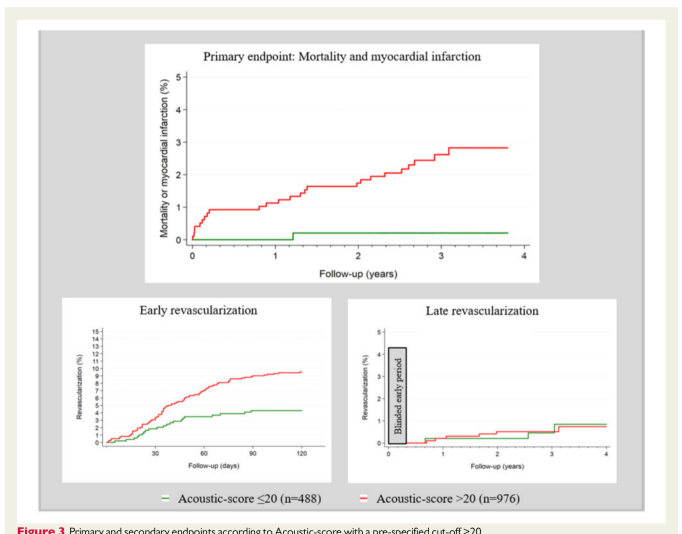


Figure 3 Primary and secondary endpoints according to Acoustic-score with a pre-specified cut-off >20.

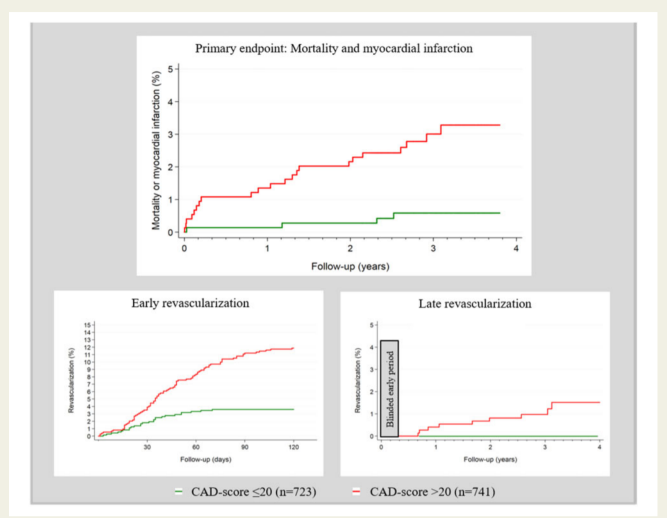


Figure 4 Primary and secondary endpoints according to coronary artery disease-score with a pre-specified cut-off >20.

part of the primary cardiac evaluation with CAD-score ≤20 (Supplementary material online, Figure S3).

Discussion

In this study, we present prognostic data showing that an Acoustic-score derived from heart sound has prognostic impact in patients with symptoms suggestive of CAD referred to cardiac CT. This Acoustic-score combined with risk factors, the CAD-score, demonstrated an HR >5 for the primary endpoint, a composite of myocardial infarction and all-cause mortality, at a pre-specified binary cut-point of CAD-score of 20. The prognostic value rose with higher CAD-score, HR >7.5 for CAD-score \ge 30 compared with \le 20. In addition, the CAD-core remained a prognostic predictor after adjusting for the updated Diamond–Forrester score and coronary stenosis at coronary CTA.

Acoustic detection of CAD

Acoustic detection of turbulent blood flow in coronary stenosis was first described in 1967 as a high-frequency diastolic murmur.¹⁷ Since

then, heart sound analysis has allowed differentiation between no, early, and advanced stages of CAD, as illustrated by diastolic frequency spectrum plots. Fig. Technological improvements in heart sound recording, segmentation, and analysis have enabled the development of an acoustic device for ruling out obstructive CAD.

Two large studies have tested the diagnostic accuracy of these acoustic rule-out devices in stable CAD. The CADence device was evaluated in the TURBULENCE study, a 21-sites multicentre study in the USA. The CADence device uses multiple recording positions and is handheld during the recording. Patients (n = 785) referred for nuclear stress testing due to chest pain symptoms and two or more CAD risk factors were included. Nuclear stress testing was followed by either coronary CTA or ICA as a reference standard with $\geq 70\%$ diameter stenosis indicating obstructive CAD. Disease prevalence was 15%, and the CADence device had a sensitivity of 78%, a specificity of 35%, and a positive and negative predictive value of 17% and 91%, respectively.

In the Dan-NICAD trial, the CADScor[®]System was tested in 1,675 patients with predominately intermediate risk of CAD who had been referred for coronary CTA.⁶ Patients were referred from five Danish hospitals, and coronary CTA was performed at two high-

Table 5 Cox regression analysis of any cardiac events after 120 days (dead, myocardial infarction or late revascularization >120 days)

	Univariate	Multivariate
Model A		
CAD-score per 10 unit	1.8 (1.3–2.5) <i>P</i> < 0.01	1.8 (1.1–2.8) <i>P</i> < 0.05
Sex, male	2.1 (0.9–5.0) <i>P</i> = 0.08	1.1 (0.4–3.0) <i>P</i> = 0.84
Age per 10 years	1.5 (0.9–2.3) <i>P</i> = 0.11	1.0 (0.6–1.7) <i>P</i> = 0.95
Model B		
CAD-score per 10 unit	1.8 (1.3–2.5) <i>P</i> < 0.01	1.7 (1.2–2.4) <i>P</i> < 0.01
Diamond–Forrester score 10 unit	1.2 (1.0–1.4) <i>P</i> < 0.05	1.1 (0.9–1.3) <i>P</i> = 0.52
Model C		
CAD-score per 10 unit	1.8 (1.3–2.5) <i>P</i> < 0.01	1.6 (1.1–2.4) <i>P</i> < 0.05
Diamond–Forrester score 10 unit	1.2 (1.0–1.4) <i>P</i> < 0.05	1.0 (0.8–1.2) <i>P</i> = 0.88
Severe stenosis at coronary CTA	6.1 (2.6–14.3) <i>P</i> < 0.001	4.2 (1.7–10.4) <i>P</i> < 0.01
Model D		
CAD-score per 10 unit	1.8 (1.3–2.5) <i>P</i> < 0.01	1.5 (1.1–2.1) <i>P</i> < 0.05
Lipid lowering therapy initiated	4.1 (1.9–9.2) <i>P</i> < 0.01	2.0 (0.8–4.9) <i>P</i> = 0.13
Revascularization at baseline	8.9 (3.9–20.0) <i>P</i> < 0.001	4.6 (1.8–11.6) <i>P</i> < 0.01

CAD-score and updated Diamond–Forrester scores are analysed as a continuous variable per 10 units. Values are hazard ratios (CI 95%).

volume centres. Patients with a coronary stenosis at coronary CTA underwent ICA with FFR, and haemodynamically obstructive CAD was defined as ICA-FFR <0.80. In Denmark, coronary CTA is the first-line test for ruling out CAD in *de novo* patients, which results in a low disease prevalence of 10% in the Dan-NICAD trial compared with 15% in TURBULENCE. The CADScor®System had sensitivity of 81%, a specificity of 53%, and a positive and negative predictive value of 16% and 96%, respectively; hence, similar sensitivities and positive predictive values but higher specificities and negative predictive values than the TURBULENCE study.

Prognostic value of acoustic analysis of heart sound

The present study is the first study to evaluate the prognostic value of an acoustic analysis of heart sounds. An isolated Acoustic-score of eight heart sound features related to CAD predicted 25 out of 26 myocardial infarctions or deaths. Combining the Acoustic-score with risk factors to obtain the CAD-score increased the proportion of patients with a score ≤20, and thus a low probability of CAD, from 33% to nearly 50%. Nonetheless, the CAD-score remains highly predictive for both deaths, myocardial infarctions, and late revascularization.

The prognostic value of the CAD-score remained significant after adjustment for coronary stenosis, which is a very strong prognostic predictor in stable CAD. This is due to specific acoustic features related to, e.g. high-risk plaque characteristic. However, further research is needed regarding the origin of these acoustic features and their correlation with specific pathologies.

The Dan-NICAD trial was designed to evaluate the diagnostic accuracy of the CAD-score with a reference of haemodynamic coronary stenosis at ICA.⁶ Decisions regarding medical treatment and revascularization were based on standard clinical care; thus, no

decision was taken based on the CAD-score. In patients with a CAD-score ≤20, lipid lowering therapy was initiated in 103 (14.3%) and 26 (3.6%) were revascularized. These patients had a therapy that might have altered their prognosis and affected the outcome of the present study. We performed multiple regression and stratified analyses to adjust for this confounder, and the conclusion on the prognosis was unaffected.

Finally, we performed a 'worst case scenario' presuming the situation that all early revascularized patients with CAD-scores \leq 20 had an event of death, myocardial infarction, or late revascularization—as mimic the most severe consequence if we had not offered early revascularization to these patients due to rule-out based on CAD-score. In this scenario, a CAD-score \geq 20 was no longer a predictor of prognosis. Contradict, 723 patients with a CAD-score \leq 20 would not have need coronary CTA for ruling out obstructive CAD. Hence, a reduction in contrast agent and iodinated radiation of \geq 1700 mSv (723 patients with an averaged radiation dose of 2.4 mSv per coronary CTA) could be achieved in this study despite we used newer generations of CT scanners.

Limitations

This cohort comprised almost exclusively Caucasians with low to intermedia pre-test probability of CAD and none had previously documented CAD. Patients were referred to coronary CTA as the first-line diagnostic test according to Danish standard of care. This approach might have introduced referral bias base on pre-test probability and due to the limitations of coronary CTA, e.g. irregular heart rate and severe obesity.

The development of CAD-score V3 required splitting the cohort into a consecutive training (n = 593) and a blinded validation (n = 1082) cohort, which might have over-fitted the results in the

total cohort analysis. However, diagnostic accuracy in the validation vs. training cohort indicated no over-fitting.⁶

The CAD-score does not distinguish between diastolic murmurs origination. Hence, diastolic valve disease may decrease CAD-score specificity as it increases the false positive rate but sensitivity remains high.¹⁹

Finally, this study is limited by the low event rates reflecting an overall favourable prognosis of patients with stable CAD. Hence, the results of this study are promising but should be interpreted with caution and final conclusions regarding the safety of acoustic rule-out strategies should await results of randomized controlled strategy trials.

Conclusion

The Acoustic-score derived from heart sound seems to give prognostic information about mortality, myocardial infarction, and late revascularization in patients with symptoms suggestive of CAD which is treated according to standard of care. Diagnostic algorithms of this Acoustic-score combined with risk factors developed for ruling out CAD, the CAD-score, risk-stratified patients even further. These findings are useful as new automated acoustic diagnostic devices have emerged for ruling out obstructive CAD.

Supplementary material

Supplementary material is available at European Heart Journal is available at online.

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