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openheart Prevalence of early stages of heart failure in an elderly risk population: the Copenhagen Heart Failure Risk Study

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Preliminary data from this study has been presented at the moderated poster session at the ESC HF 2018 Congress, Vienna, Austria.

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ABSTRACT

Aims To describe the prevalence of heart failure (HF) stages in elderly outpatients with risk factors for HF but without known HF, and characterise the clinical, biochemical and echocardiographic findings in each stage.

Background Early stages of HF are underdiagnosed; nevertheless, they are associated with an increased risk of hospitalisation and increased mortality. The prevalence of HF stages in elderly high-risk patients is unknown.

Methods A total of 400 patients were consecutively included: ≥ 60 years old, ≥ 1 risk factor for HF (diabetes, chronic kidney disease, cardiovascular disease, atrial fibrillation, hypertension), and without known or suspected HF. HF stages were defined as the following: stage A (risk factor for HF, normal echocardiography), stage B (abnormal echocardiography, without symptoms of HF) and stage C (abnormal echocardiography, symptoms of HF, clinical signs/increased plasma aminoterminal pro-B-type natriuretic peptide [NT-proBNP] concentrations).

Results After thorough examination 44.25% of patients were categorised as HF stage A, 37.5% were HF stage B and 18.25% HF stage C. Those with higher stages of HF were older ($p < 0.001$) and more often had atrial fibrillation ($p = 0.006$). The median plasma NT-proBNP concentrations (pg/mL) were 132.5 for HF stage A, 275.5 for HF stage B and 400.0 for HF stage C ($p < 0.001$). Detectable plasma troponin-I was more frequent with abnormal echocardiography: HF stage A 9.7%, HF stage B 27.3% and HF stage C 30.1% ($p < 0.001$). HF stage C reported higher score on the Minnesota Living with Heart Failure Questionnaire ($p < 0.001$).

Conclusions In an elderly high-risk population without known or suspected HF, more than half were HF stage B or C. Higher stages of HF had increased plasma concentrations of NT-proBNP and troponin-I, besides a reduced quality of life. Focus on symptoms and signs of HF in this population are warranted.

INTRODUCTION

A large proportion of elderly people have early stages of heart failure (HF) based on the American College of Cardiology and the American Heart Association definition of HF stages.¹ Increasing age of the general population and increasing prevalence of risk

Key questions

What is already known about this subject?

- The heart failure (HF) staging definition introduced by the American College of Cardiology and the American Heart Association underlines the importance of recognition of asymptomatic patients with early stages of HF.
- Previous population-based studies have demonstrated a high prevalence of early stages of HF, HF stages A and B, with an increased risk of hospitalisation with HF and increased mortality.

What does this study add?

- In an elderly cohort living with risk factors for HF without known or suspected HF, more than half of the patients had an abnormal echocardiography and one-fifth had undiagnosed HF.
- The most common type of HF in this cohort was HF with preserved ejection fraction (HFpEF).
- Our findings suggest that increased focus on HF symptoms and thorough examination of high-risk patients can identify undiagnosed HF, in particular patients with HFpEF.
- This should be considered in specialised outpatient clinics treating the risk factors for HF and at the department of cardiology.

How might this impact on clinical practice?

- Our results underscore the need for increased focus on HF symptoms in high-risk populations to improve management of early stages of HF and improve patients' quality of life, besides allocating patients with HFpEF to high-quality randomised clinical trials.

conditions (eg, hypertension, diabetes) lead to an increased number of persons with a risk of HF, categorised as HF stage A.^{2,3} Overt HF, categorised as HF stage C or D, is associated with poor prognosis, but also preclinical stages of HF, HF stage B, has an increased risk of hospitalisation with congestion and an increased mortality risk.⁴⁻⁷ Early recognition and improved treatment of risk conditions may delay the progression of early stages of HF.⁸ Furthermore, promising research in

the treatment of HF with preserved ejection fraction (HFpEF) is ongoing, and recognition of these patients needs to be improved to secure an optimal treatment.

Large community studies have observed that a considerable number of patients have early stages of HF. HF stage B was observed in 24%–34% and HF stage C in 12%–13% of these populations, with increasing prevalence with increasing age.^{4–6} The prevalence of HF stages B and C may, therefore, be even higher if a high-risk population is investigated, for example, elderly patients living with risk factors for HF.

The aim of this study was to evaluate the prevalence and clinical characteristics of HF stages in elderly patients with a high risk of HF, but without known or suspected HF.

METHODS

Study population

The Copenhagen Heart Failure Risk Study is a prospective cohort study. Patients were included from the Department of Cardiology, the Clinic of Diabetes and the Clinic of Nephrology at the Herlev and Gentofte University Hospital, Copenhagen, Denmark, enrolled from December 2014 to June 2016. Patients were screened consecutively during admission at the Department of Cardiology (screening 5 days a week), included from the day of discharge until 6 months later. Patients from the Clinic of Diabetes were screened consecutively by a treating physician at a scheduled visit (screening 2 days a week), included within 6 months. Patients from the Clinic of Nephrology were screened consecutively from the list of scheduled visits (screening 2 days a week), included within 6 months.

Inclusion and exclusion criteria are presented in [Box 1](#). Data were collected during a single visit. Physical examination, medical history including current symptoms and medication, height, weight, non-invasive blood pressure, and 12-lead ECG were obtained; venous blood samples and spot-urinary sample were collected; patients fulfilled the Minnesota Living with Heart Failure Questionnaire (MLHFQ); and advanced echocardiography was performed. The Framingham probability of congestive HF within 4 years score was calculated. This is a risk prediction model that includes age, systolic blood pressure, heart rate, left ventricle (LV) hypertrophy on ECG, coronary heart disease, valve disease, diabetes and body mass index (if female).⁹

In total 3359 patients were screened and 400 patients were enrolled; inclusion flow is shown in [figure 1](#). The primary reason for patients not being eligible for this study was due to the presence of one or more exclusion criteria; especially patients from the Department of Cardiology often had advanced/ongoing cardiological diseases. No patients were excluded after inclusion, a full echocardiography was missing in one patient (n=1), and plasma concentrations of aminoterminal pro-B-type

Box 1 Inclusion and exclusion criteria for the Copenhagen Heart Failure Risk Study

Inclusion criteria.

- ▶ Age ≥60 years.
- ▶ Plus, one or more risk factor for heart failure (HF).
 - Stable ischaemic heart disease.
 - Peripheral arterial disease.
 - Hypertension.
 - History of atrial fibrillation/atrial flutter, diabetes.
 - Apoplexia cerebri.
 - Chronic kidney disease.

Exclusion criteria.

- ▶ Age <60 years.
- ▶ No risk factor of HF.
- ▶ Previously diagnosed with or admitted for HF, known left ventricle ejection fraction ≤45%.
- ▶ Current admission with congestion or dyspnoea and suspected HF by the treating clinician.
- ▶ Advanced cardiac disease.
 - Moderate-severe valvular disease, third-degree atrioventricular block, pacemaker with >70% ventricular pace.
- ▶ Ongoing or recent cardiac disease or procedure.
 - Cardioversion within the previous 30 days, coronary angiography or percutaneous coronary intervention or coronary artery bypass operation within the previous 30 days, ongoing endocarditis or myocarditis, or pulmonary embolism.
- ▶ Dialysis/estimated glomerular filtration rate <15 mL/min/1.73 m².
- ▶ Moderate-severe chronic obstructive pulmonary disease (global initiative for chronic obstructive lung disease stadium (GOLD) ≥2C).
- ▶ Sepsis (systemic inflammatory response system criteria) within the previous 30 days or ongoing intravenous antibiotic.
- ▶ Estimated survival less than 1 year (eg, terminal cancer or due to frailty).

natriuretic peptide (NT-proBNP) was missing in one patient (n=1).

Echocardiography

Echocardiography was performed according to current guidelines.^{10–11} Examinations were performed on Vivid E9 (General Electric Vingmed Ultrasound, Horten, Norway) by the same trained physician. Analyses were made off-line in EchoPAC (V.201,70.1, General Electric Vingmed Ultrasound) by one physician, blinded to clinical and biochemical outcomes. Before examination patients were registered by name and social security number, height and weight were entered, and a three-lead ECG of good quality was continually registered. In patients with sinus rhythm, three loops were recorded, and for atrial fibrillation (AF) five loops were recorded. Images for strain should have a frame rate of 40–80 frames/s, while images for tissue Doppler >160 frames/s.

We predefined a study definition of an ‘abnormal echocardiography’ based on current recommendations and previous studies in the elderly patients (online supplementary table 1). Abnormal structure of the LV was defined as LV end-diastolic volume (LVEDV) indexed to body surface area (BSA) (female: >61 mL/m², male: >74

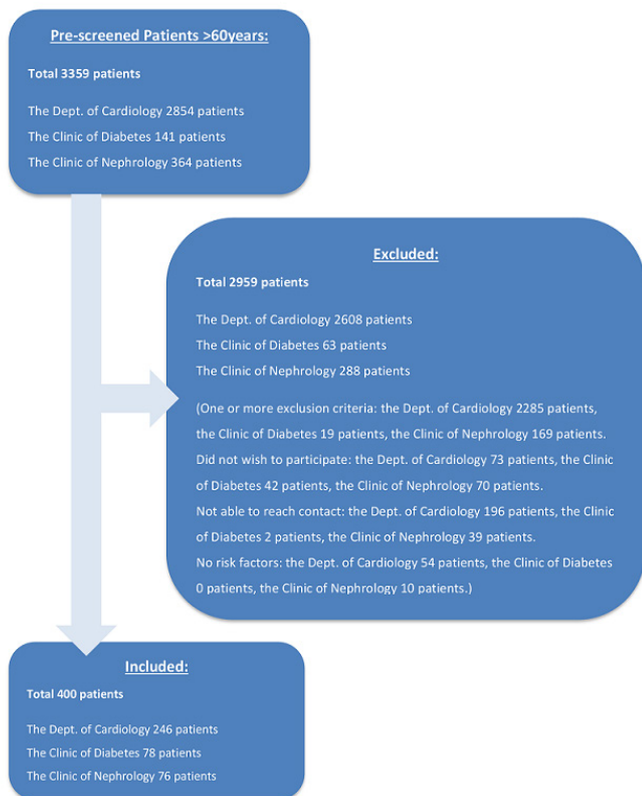


Figure 1 Inclusion flow of the Copenhagen Heart Failure Risk Study.

mL/m²)¹⁰ and LV mass indexed to BSA (female: >96 g/m², male: >116 g/m²).¹⁰ Abnormal systolic function was evaluated with left ventricle ejection fraction (LVEF) and global longitudinal strain (GLS). The limit for LVEF was set to detect mildly abnormal changes (female: ≤53%, male: ≤51%). The ARIC study has recently evaluated GLS in the HF staging system and confirmed its prognostic value for new-onset HF and mortality.⁴ However, validated reference limits do not exist, and therefore we used age-based and sex-based reference limits from the Framingham Heart Study¹² (female: ≥-15.3%, male: ≥-14.7%).

HF stages after examination

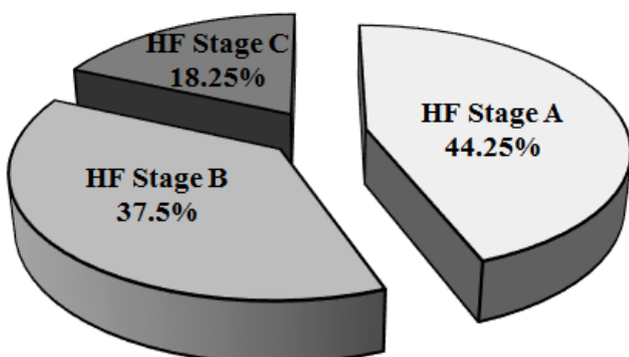


Figure 2 Prevalence of heart failure (HF) stages after clinical, biochemical and echocardiographic examination.

We included abnormal diastolic measures (tissue Doppler imaging and left atrial volume index), since these parameters have been associated to an increased risk of new-onset HF and mortality.^{4,5} The myocardial peak early velocity (e') is more affected by ageing, and we adjusted the study reference value according to normal values reported from the Atherosclerosis Risk in Communities Study (ARIC)⁴ (female: <4.1 cm/s, male <4.3 cm/s), while standard reference values were used for the more robust E/e' (lateral: >13, septal: >15) and for the left atrial volume indexed to BSA (≥ 35 mL/m²).¹¹ Significant valvular disease was defined as moderate-severe aortic stenosis (aortic valve area <1 cm²), moderate-severe aortic regurgitation (visual judgement) and severe mitral stenosis (peak gradient mean >12 mm Hg).

LVEF (%) was calculated by the Simpson biplane method. Regional Wall Motion Score Index was calculated using the 16-segment model. LVEDV (mL) was calculated from apical two-chamber and four-chamber views. LV mass index (g/m²) was calculated from linear dimensions in the parasternal view. Left atrial end-systolic volume (mL) was calculated from apical two-chamber and four-chamber views. Myocardial peak early velocity (e') (cm/s) and peak systolic velocity (s') (cm/s) were measured medially and septally at the mitral annulus with pulse wave tissue Doppler. Mitral inflow was measured with pulse wave Doppler, peak velocity of early filling (E) (cm/s) and atrial filling (A) (cm/s), and mitral valve deceleration time (ms). E/e' ratio was calculated septally and laterally. Tricuspid annular plane systolic excursion (mm) was measured with m-mode in the lateral tricuspid annulus. Longitudinal strain (%) was measured in three apical views, and GLS was calculated as the average of the three, an 18-segment model. The presence of valvular heart disease was evaluated: aortic sclerosis by visual judgement, aortic stenosis calculated (aortic velocity time integral [VTI], LV outflow tract VTI and diameter), aortic regurgitation by visual judgement of colour flow, mitral stenosis calculated (mitral valve VTI mean gradient), and mitral regurgitation by visual judgement of colour flow.

Definition of HF stages

HF stages were defined according to the American College of Cardiology/American Heart Association definition¹:

- ▶ HF stage A: patients with risk factors of HF, with normal echocardiography and without signs or symptoms of HF.
- ▶ HF stage B: patients with risk factors of HF, with an abnormal echocardiography but without past or current symptoms of HF.
- ▶ HF stage C: patients with an abnormal echocardiography and past or current symptoms of HF.

To ensure the HF diagnoses in patients categorised as HF stage C, we used the European Society of Cardiology definition of HF.¹⁴ Signs of HF may be absent, for which reason a biochemical confirmation of HF was included. Patients in HF stage C needed to fulfil three criteria:

(1) symptoms of HF (two of the following: dyspnoea or orthopnoea and oedema, or treatment with loop diuretics); (2) abnormal echocardiography according to study definition; and (3) clinical signs and/or abnormal plasma concentrations of NT-proBNP (sinus rhythm >125 ng/L,¹⁴ AF >524 pg/mL¹⁵).

Biochemistry

Venous blood samples were drawn on the day of examination. Routine blood samples (eg, haemoglobin, creatinine, highly sensitive C reactive peptide [hsCRP]) and urine-albumin were analysed successively. Blood samples for later tests were centrifuged at 4°, and plasma was stored at -80°C. Plasma concentrations of NT-proBNP (ng/L) were measured with the IMMULITE 2000 NT-proBNP, solid-phase, two-site chemiluminescent immunometric assay (Siemens Healthcare Diagnostics).¹⁶ Plasma concentrations of mid-regional proadrenomedullin (MR-proADM) (nmol/L) and copeptin (C-terminal provasopressin) (pmol/L) were measured on BRAHMS KRYPTOR with specified automated immunofluorescent assays.^{17,18} Plasma concentrations of troponin-I were measured with ADVIA Centaur TnI-Ultra assay (Siemens Healthcare Diagnostics).¹⁹ Concentrations of urinary albumin were measured in spot-urine. Microalbuminuria was defined as 20–199 mg/L and macroalbuminuria ≥200 mg/L.

Statistical methods

Patients were categorised in HF stages A, B and C. Dichotomous variables were presented as numbers and prevalences (%). Continuous variables were both normally and skewed distributed and are presented as medians with IQRs (25th and 75th percentiles). Baseline characteristics and echocardiographic parameters were compared across HF stages, with pairwise comparisons between groups. For continuous variables, we used Wilcoxon rank-sum test (Kruskal-Wallis test) and a trend test, and for dichotomous variables we used χ^2 test. In the pairwise comparison, we adjusted for multiple testing with the Bonferroni correction. Skewed distributed variables were log₂-transformed when appropriate to reduce the effects of skewness. Trend tests were performed using the Cochran-Armitage trend test for proportions and logistic regression models for continuous variables (dependent variable: HF stages). Multivariate linear and logistic regression models were performed to evaluate the associations between the HF stages and baseline characteristic with a p value <0.05 in the trend test. The regression models were adjusted for age, gender, AF, hypertension, ischaemic heart disease, stroke, diabetes, estimated glomerular filtration rate (eGFR) and department from where the patients were included. A two-sided p value <0.05 was considered significant, and in case of Bonferroni correction due to multiple testing a p value <0.05/3 (=0.017) was considered significant. Analyses were performed using the SAS Enterprise Guide V.7.11 statistical software.

Sample size

It was anticipated that the prevalence of patients with HF stage B or C would be 30%. With a power of 80% at a significance level of 5%, a total of 369 patients should be included. To account for dropouts and eventually missing data, 400 patients were included.

RESULTS

Prevalence of HF stages

Before examination by default, all patients were considered as HF stage A, based on the inclusion and exclusion criteria. After examination 177 (44.25%) remained as HF stage A, 150 (37.5%) were HF stage B and 73 (18.25%) were HF stage C, illustrated in figure 2. In HF stage A, 29.9% reported symptoms of HF, 36.7% had clinical signs of HF and 50.6% had elevated plasma concentrations of NT-proBNP. However, these patients did not fulfil all the criteria for an HF diagnosis, and their symptoms may therefore be considered non-cardiac. Only 28.25% in HF stage A had no symptoms of HF, no clinical signs and normal plasma concentrations of NT-proBNP. In HF stage B, all patients had an abnormal echocardiography, 38.0% had clinical signs of HF and 73.3% had elevated plasma concentrations of NT-proBNP, and 30.7% had both clinical signs of HF and elevated plasma concentrations of NT-proBNP. Two patients did not fit with the definition of HF stages; they had an abnormal echocardiography and symptoms of HF, but no clinical signs nor elevated plasma concentrations of NT-proBNP. They were categorised as HF stage B. Patients in HF stage C had an abnormal echocardiography and symptoms of HF; 78.1% had clinical signs of HF and 83.6% had elevated plasma concentrations of NT-proBNP, and 61.6% had both clinical signs of HF and elevated plasma concentrations of NT-proBNP.

Characterisation of HF stages

Baseline characteristics are presented in table 1. The median age was 72 years, ranging from 60 to 97, and patients in HF stage B and C were older than patients in HF stage A (p<0.001 for trend). Of the patients, 48.5% were female, with no significant difference between stages of HF.

Patients with HF stage C were more frequently admitted due to dyspnoea or palpitations. However, in all stages of HF, chest pain was the dominating reason for hospitalisation, and it did not differ between the different stages. The frequencies of chest pain and palpitations were almost identical to the frequencies of ischaemic heart disease and AF (table 1).

There was no difference in 'numbers of risk factors' (p=0.093 for trend); overall 119 patients had one risk factor, 184 had two risk factors, 79 had three risk factors and 18 had more than four risk factors. The only risk factor that differed in prevalence was a history of AF, with more patients in higher stages of HF (p=0.001 for trend). Patients with higher stages of HF had lower eGFR (p=0.003 for trend), haemoglobin (p=0.015 for trend)

Table 1 Clinical characteristics of patients in the Copenhagen Heart Failure Risk Study, according to HF stage

	All patients (n=400)	HF stage A (n=177)	HF stage B (n=150)	HF stage C (n=73)	P value for trend
Age, years	72 (67, 78)	69 (65, 73)*†	75 (69, 79)	74 (69, 82)	<0.001
Female sex, n (%)	194 (48.5)	78 (44.1)	74 (49.3)	42 (57.5)	0.053
BMI, kg/m ²	27.3 (24.5, 30.9)	27.1 (24.3, 30.7)	27.2 (24.5, 30.1)	28.3 (24.7, 32.8)	0.117
BMI ≥30, n (%)	119 (29.8)	52 (29.4)	39 (26.0)	28 (38.4)	0.309
Systolic blood pressure, mm Hg	137.0 (126.5, 149.0)	134.0 (126.0, 146.0)	140.0 (128.0, 150.0)	137 (125.0, 150.0)	0.312
Diastolic blood pressure, mm Hg	79.0 (72.0, 85.0)	81.0 (74.0, 85.0)†	78.0 (71.0, 86.0)	77.0 (69.0, 84.0)	0.0135
HR, beats/min	69.0 (60.0, 79.0)	69.0 (61.0, 78.0)	68.5 (60.0, 81.0)	69.0 (60.0, 79.0)	0.471
Left branch bundle block ECG, n (%)	13 (3.3)	1 (0.6)*†	8 (5.3)	4 (5.6)	0.015
Right branch bundle block ECG, n (%)	25 (6.3)	11 (6.2)	7 (4.7)	7 (9.7)	0.475
LV hypertrophy ECG, n (%)	7 (1.8)	2 (1.1)	4 (2.7)	1 (1.4)	0.675
NYHA class I, n (%)	235 (58.8)	117 (66.1)†	98 (65.3)‡	20 (27.4)	<0.001
NYHA class II, n (%)	145 (36.3)	55 (31.1)†	45 (30.0)‡	45 (61.6)	<0.001
NYHA class III, n (%)	20 (5.0)	5 (2.8)†	7 (4.7)	8 (11.0)	0.012
Smoking (former or active), n (%)	249 (62.3)	120 (67.8)*	81 (54.0)	48 (65.8)	0.316
Variables for HF staging					
Symptoms of heart failure, n (%)	128 (32.0)	53 (29.9)*†	2 (1.3)‡	73 (100)	
Clinical signs of heart failure, n (%)	179 (44.8)	65 (36.7)†	57 (38.0)‡	57 (78.1)	
Abnormal echocardiography, n (%)	223 (55.8)	0 (0.0)*†	150 (100)	73 (100)	
MLHFQ	12.0 (2.0, 28.0)	11.0 (2.0, 28.0)†	8.0 (1.0, 17.0)‡	25.0 (16.0, 42.0)	0.001
Framingham 4-year HF risk score	11.0 (8.0, 15.0)	9.0 (7.5, 14.0)*†	11.5 (9.0, 16.0)	12.0 (9.0, 16.0)	<0.001
Indication for hospitalisation					0.043
Chest pain, n (%)	115 (28.8)	55 (31.1)	38 (25.3)	22 (30.1)	0.646
Palpitations/Arrhythmia, n (%)	80 (20.0)	25 (14.1)	36 (24.0)	19 (26.0)	0.013
Syncope, n (%)	30 (7.5)	11 (6.2)	13 (8.7)	6 (8.2)	0.477
Dyspnoea, n (%)	12 (3.0)	4 (2.3)	2 (1.3)	6 (8.2)	0.044
Other, n (%)	9 (2.3)	4 (2.3)	5 (3.3)	0 (0)	0.545
Ambulatory visits—Clinic of Diabetes, n (%)	78 (19.5)	40 (22.6)	28 (18.7)	10 (13.7)	0.100
Ambulatory visits—Clinic of Nephrology, n (%)	76 (19.0)	38 (21.5)	28 (18.7)	10 (13.7)	0.160
Medical history					
Hypertension, n (%)	328 (82.0)	145 (81.9)	120 (80.0)	63 (86.3)	.568
Ischaemic heart disease, n (%)	98 (24.5)	37 (20.9)	41 (27.3)	20 (27.4)	0.187
Atrial fibrillation, n (%)	119 (29.8)	40 (22.6)†	48 (32.0)	31 (42.5)	0.001
Diabetes, n (%)	143 (35.8)	69 (39.0)	51 (34.0)	23 (31.5)	0.218
Chronic kidney disease, n (%)	63 (15.8)	30 (17.0)	23 (15.3)	10 (13.7)	0.506
Apoplexia cerebri, n (%)	48 (12.0)	19 (10.7)	23 (15.3)	6 (8.2)	0.915
Mild COPD or asthma, n (%)	34 (8.5)	18 (10.2)	6 (4.0)‡	10 (13.7)	0.840
Evaluation of risk factors					
Risk factors, n (%)	2.0 (1.0, 2.0)	2.0 (1.0, 2.0)	2.0 (1.0, 3.0)	2.0 (2.0, 2.0)	0.093
More than two risk factors, n (%)	97 (24.3)	39 (22.0)	41 (27.3)	17 (23.3)	0.615
Medication					
ACE inhibitor, n (%)	105 (26.3)	49 (27.7)	45 (30.0)‡	11 (15.1)	0.103
Angiotensin receptor antagonist, n (%)	139 (34.8)	59 (33.3)	51 (34.0)	29 (39.7)	0.388
Aldosterone antagonist, n (%)	5 (1.3)	0 (0.0)	3 (2.0)	2 (2.7)	0.047
Calcium antagonist, n (%)	125 (31.3)	48 (27.1)	50 (33.3)	27 (37.0)	0.097

Continued

Table 1 Continued

	All patients (n=400)	HF stage A (n=177)	HF stage B (n=150)	HF stage C (n=73)	P value for trend
Beta blocker, n (%)	187 (46.8)	71 (40.1)	75 (50.0)	41 (56.2)	0.012
Loop diuretics, n (%)	60 (15.0)	16 (9.0)*	15 (10.0)‡	29 (39.7)	<0.001
Thiazide, n (%)	125 (31.3)	62 (35.0)†	49 (32.7)	14 (19.2)	0.025
Statin, n (%)	247 (61.8)	115 (65.0)	90 (60.0)	42 (57.5)	0.226
Per oral antidiabetics, n (%)	111 (27.8)	57 (32.2)	35 (23.3)	19 (26.0)	0.172
Insulin, n (%)	65 (16.3)	34 (19.2)	22 (14.7)	9 (12.3)	0.141
Biochemistry					
Haemoglobin, g/L	138.6 (128.9, 146.6)	140.2 (132.1, 148.2)†	137.0 (125.7, 146.6)	133.7 (127.3, 145.0)	0.015
Thrombocytes, $\times 10^9/L$	232.2 (190.5, 280.0)	230.0 (193.0, 281.0)	228.5 (189.0, 270.0)	242.0 (184.0, 286.0)	0.960
Leucocytes, $\times 10^9/L$	7.2 (6.1, 8.4)	7.2 (6.1, 8.4)	7.0 (5.9, 8.1)‡	7.5 (6.6, 8.5)	0.092
hsCRP, mg/L	0 (0.0, 3.0)	0 (0.0, 3.0)	0 (0.0, 4.0)	0 (0.0, 4.0)	0.621
hsCRP ≥ 3 mg/L, %	130 (32.5)	48 (27.1)	55 (36.7)	27 (37.0)	0.067
Creatinine, $\mu\text{mol/L}$	82.0 (69.0, 106.0)	80.0 (66.0, 103.0)	82.0 (71.0, 109.0)	87.0 (68.0, 116.0)	0.213
eGFR, mL/min/1.73 m ²	70.0 (53.0, 86.0)	76.0 (57.5, 90.0)*†	65.5 (51.0, 84.0)	65.0 (47.5, 83.5)	0.003
Microalbuminuria, %	143 (37.9)	51 (31.1)	61 (43.3)	31 (43.1)	0.036
Macroalbuminuria, %	38 (10.1)	13 (7.9)*	18 (12.8)	7 (9.7)	0.457
HbA1c, %	5.8 (5.5, 6.6)	5.8 (5.6, 6.9)	5.8 (5.5, 6.5)	5.8 (5.6, 6.4)	0.186
Albumin, g/L	42.0 (40.0, 44.0)	43.0 (40.0, 45.0)*†	42.0 (40.0, 44.0)	41.0 (40.0, 44.0)	<0.001
Bilirubin, $\mu\text{mol/L}$	10.0 (8.0, 14.0)	10.0 (7.0, 14.0)	11.0 (8.0, 14.0)	10.0 (8.0, 14.5)	0.037
ALAT, U/L	29.0 (22.0, 39.0)	31.0 (24.0, 39.0)	27.0 (20.0, 40.0)	26.0 (20.0, 37.0)	0.524
TSH, $\times 10^{-3}$ IE/L	1.3 (0.8, 1.9)	1.3 (0.8, 1.8)	1.2 (0.8, 1.8)	1.6 (0.9, 2.4)	0.109
Cholesterol total, mmol/L	4.4 (3.7, 5.2)	4.5 (3.7, 5.3)	4.3 (3.7, 5.2)	4.3 (3.6, 5.1)	0.332
HDL, mmol/L	1.2 (1.0, 1.6)	1.3 (1.0, 1.5)	1.2 (1.0, 1.6)	1.3 (1.0, 1.6)	0.508
LDL, mmol/L	2.2 (1.6, 2.9)	2.2 (1.7, 2.9)	2.2 (1.7, 2.8)	2.2 (1.6, 2.7)	0.521
NT-proBNP, ng/L	203.0 (98.0, 601.0)	132.5 (70.0, 320.0)*†	275.5 (118.0, 746.0)	400.0 (179.0, 1210.0)	<0.001
Troponin-I, ng/L	0 (0.0, 0.0)	0 (0.0, 0.0)	0 (0.0, 12.0)	0 (0.0, 12.0)	0.490
Troponin-I ≥ 10 , ng/L, %	80 (20.1)	17 (9.7)	41 (27.3)	22 (30.1)	<0.001
MR-proADM, nmol/L	0.85 (0.66, 1.06)	0.75 (0.59, 0.95)*†	0.87 (0.69, 1.07)‡	1.00 (0.84, 1.17)	<0.001
Copeptin, pmol/L	7.2 (4.5, 14.3)	6.4 (3.8, 12.5)	7.6 (4.9, 14.5)	9.3 (5.2, 19.7)	0.013

Categorical values are presented in percentages, and continuous values are presented as median with 25th and 75th percentile.

*P<0.017 for HF stage A vs HF stage B.

†P<0.017 for HF stage A vs HF stage C.

‡P<0.017 for HF stage B vs HF stage C.

ALAT, alanine transaminase; BMI, body mass index; COPD, chronic obstructive pulmonary disease; eGFR, estimated glomerular filtration rate; HbA1c, haemoglobin A1c; HDL, high-density lipoprotein; HF, heart failure; HR, heart rate; hsCRP, highly sensitive C reactive peptide; LDL, low-density lipoprotein; LV, left ventricle; MLHFQ, Minnesota Living with Heart Failure Questionnaire; MR-proADM, mid-regional proadrenomedullin; NT-proBNP, aminoterminal pro-brain natriuretic peptide; NYHA, New York Heart Association; TSH, thyroid stimulating hormone.

and plasma albumin ($p<0.001$ for trend) (table 1). In the adjusted analyses the associations between HF stages and age, diastolic blood pressure, MLHFQ, haemoglobin, eGFR, prevalence of microalbuminuria and bilirubin remained significant (table 2).

Plasma concentrations of NT-proBNP increased with higher stages of HF ($p<0.001$), with a significant difference between each stage, illustrated in figure 3. Plasma concentrations of troponin-I were more frequently increased in patients with HF stages B and C ($p<0.001$ for trend). Plasma concentrations of copeptin and MR-proADM also

increased with higher stages of HF ($p<0.001$ for trend). MLHFQ score was higher in patients with HF stage C, reflecting a decreased quality of life ($p<0.001$ for trend). Framingham probability of congestive HF within 4 years score was higher in patients with HF stages B and C ($p<0.001$ for trend), illustrated in figure 3. In the adjusted analyses the associations between HF stages and increased plasma troponin-I concentrations ($p=0.010$), plasma NT-proBNP concentrations ($p<0.001$), plasma MR-proADM concentrations ($p<0.001$) and plasma copeptin concentrations ($p=0.009$) remained significant (table 2).

Table 2 Multivariate regression models according to HF stages

Multivariate linear regression models according to HF stages			
	Estimate	95% CI	P value
Age			
HF stage A	Ref		<0.001
HF stage B	3.55	2.137 to 4.956	<0.001
HF stage C	3.33	1.525 to 5.128	<0.001
Diastolic blood pressure			
HF stage A	Ref		<0.001
HF stage B	-0.09	-2.291 to 2.106	0.934
HF stage C	-1.87	-4.640 to 0.901	0.185
Minnesota Living with Heart Failure Questionnaire			
HF stage A	Ref		0.049
HF stage B	-6.11	-10.32 to 1.89	0.005
HF stage C	12.33	7.18 to 17.56	<0.001
Framingham 4-year risk of congestion			
HF stage A	Ref		0.118
HF stage B	0.36	-0.33 to 1.05	0.306
HF stage C	0.61	-0.26 to 1.48	0.17
Haemoglobin			
HF stage A	Ref		<0.001
HF stage B	-0.06	-0.26 to 0.14	0.528
HF stage C	-0.15	-0.41 to 0.10	0.232
eGFR			
HF stage A	Ref		<0.001
HF stage B	-1.36	-5.25 to 2.54	0.495
HF stage C	-2.77	-7.67 to 2.14	0.269
Bilirubin			
HF stage A	Ref		<0.001
HF stage B	1.04	-0.08 to 2.17	0.069
HF stage C	1.1	-0.32 to 2.52	0.128
Albumin			
HF stage A	Ref		<0.001
HF stage B	-0.65	-1.39 to 0.09	0.084
HF stage C	-0.84	-1.76 to 0.09	0.077
NT-proBNP			
HF stage A	Ref		<0.001
HF stage B	0.34	0.02 to 0.66	0.036
HF stage C	0.87	0.48 to 1.28	<0.001
MR-proADM			
HF stage A	Ref		<0.001
HF stage B	0.07	-0.03 to 0.17	0.179
HF stage C	0.29	0.16 to 0.42	<0.001

Continued

Table 2 Continued

Multivariate linear regression models according to HF stages			
	Estimate	95% CI	P value
Copeptin			
HF stage A	Ref		0.009
HF stage B	0.16	-0.15 to 0.35	0.422
HF stage C	0.16	0.11 to 0.74	0.008
Multivariate logistic regression models according to HF stages			
	OR	95% CI	P value
Left branch bundle block on ECG			
HF stage A vs HF stage B	6.44	0.68 to 60.56	0.104
HF stage A vs HF stage C	8.16	0.74 to 90.51	0.087
NYHA class			
HF stage A vs HF stage B	1.21	0.74 to 1.99	0.446
HF stage A vs HF stage C	0.29	0.16 to 0.52	<0.001
Atrial fibrillation			
HF stage A vs HF stage B	1.48	0.78 to 2.80	0.227
HF stage A vs HF stage C	2.39	1.12 to 5.10	0.024
Microalbuminuria			
HF stage A vs HF stage B	1.35	0.82 to 2.25	0.241
HF stage A vs HF stage C	1.3	0.69 to 2.44	0.412
Troponin-I ≥ 10 ng/L			
HF stage A vs HF stage B	2.35	1.21 to 4.54	0.011
HF stage A vs HF stage C	2.55	1.18 to 5.50	0.017

The response variables are specified below. The predictor variables are HF stages, age, gender, atrial fibrillation, hypertension, ischaemic heart disease, stroke, diabetes, kidney function (eGFR) and department from where the patients are included.

HF, heart failure; MR-proADM, mid-regional proadrenomedullin; NT-proBNP, aminoterminal pro-brain natriuretic peptide; NYHA, New York Heart Association; Ref, reference; eGFR, estimated glomerular filtration rate.

Echocardiographic parameters are presented in table 3. An 'abnormal echocardiography' was part of the HF stages B and C definition, and this reflects in the results with no major differences between the two. The only echocardiographic parameters with a difference between HF stages B and C were the proportion of abnormal LV mass index ($p < 0.001$ for trend) and the proportion of abnormal e' septal ($p < 0.0001$ for trend). The median LVEF was 62.64%, slightly lower in higher stages of HF ($p < 0.0001$). In total 12 patients (3%) had an LVEF $\leq 40\%$, of these 8 had no symptoms and were therefore HF stage B, while 4 reported symptoms of HF and were therefore HF stage C.

Despite having excluded patients with known moderate-severe valvular heart disease, two patients were encountered with moderate-severe aortic stenosis (aortic

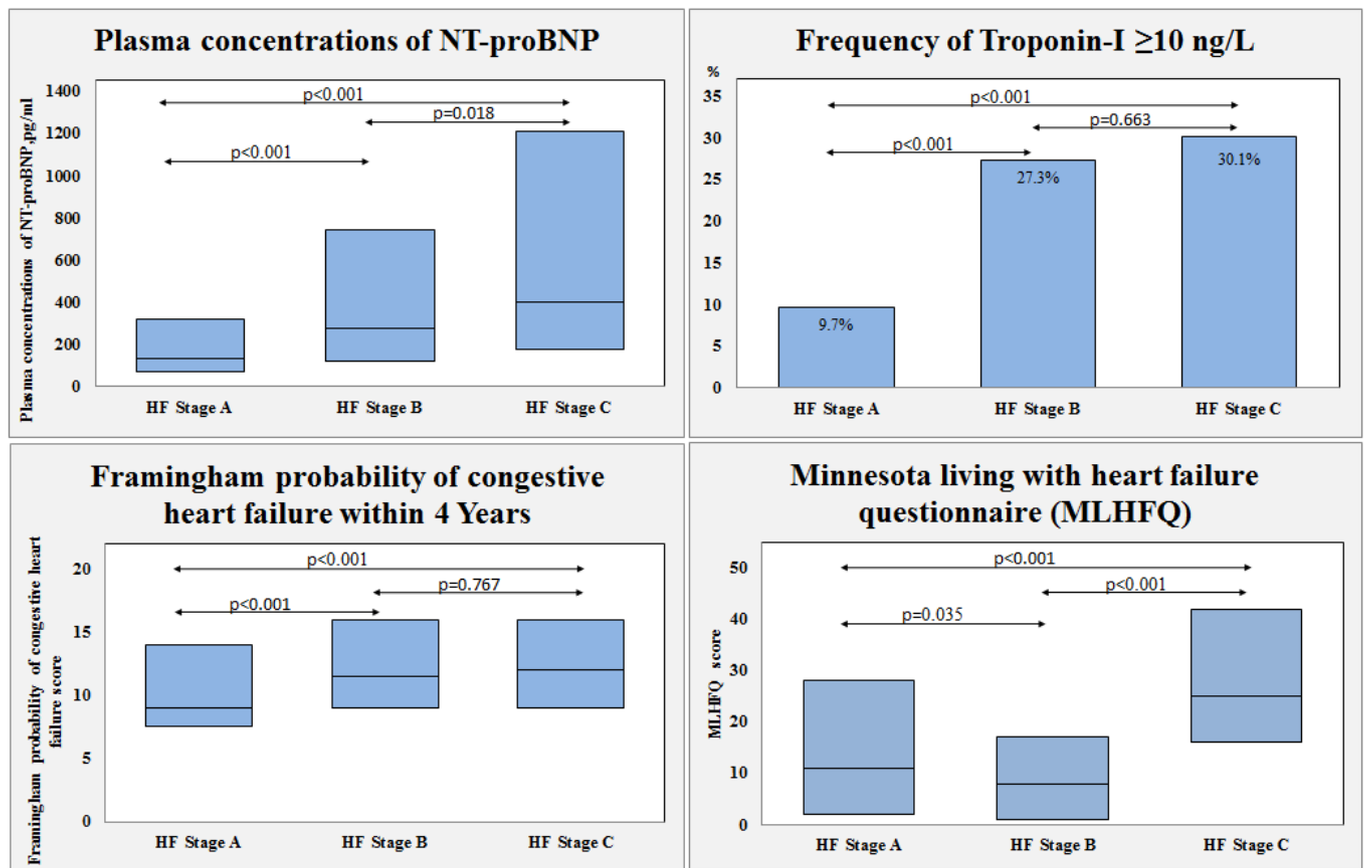


Figure 3 Cardiac biomarkers, quality of life and Framingham HF risk score according to HF stages. Plasma concentrations of NT-proBNP presented as median with 25th and 75th percentiles and patients with elevated troponin-I (≥ 10 ng/L) as proportion (%) according to HF stages. Minnesota Living with Heart Failure Questionnaire and Framingham probability of congestive HF within 4 years score presented as median with 25th and 75th percentiles. HF, heart failure; NT-proBNP, aminoterminal pro-brain natriuretic peptide.

valve area < 1.0 cm²) and one with moderate-severe aortic regurgitation. No patients with severe mitral regurgitation were encountered.

DISCUSSION

The main findings of this study were that HF stages B and C are frequent in an elderly high-risk population without known or suspected HF, and especially HFpEF is underdiagnosed. Clinical characteristics were similar in HF stages B and C; however, plasma concentrations of NT-proBNP increased with higher stages of HF, whereas troponin-I was more frequently detectable in patients with an abnormal echocardiography. Finally, higher stages of HF had decreased quality of life.

Clinical characteristics of HF stages

Elderly patients with one or more risk factor for HF often present with symptoms compatible with HF. After a thorough examination, we observed that more than half of the patients had structural or functional abnormalities of the LV evaluated by echocardiography and almost one-fifth of the patients had undiagnosed HF. Previous studies have examined the distribution of HF stages in the community, but to our knowledge this is the first study to describe the

prevalence of HF stages in elderly patients living with risk factors for HF, under relevant medical care. The findings of this study are in fine line with the prevalence of HF stages found in large community studies.⁴⁻⁶ However, by focusing on high-risk patients, we increased the observed prevalence of HF stages B and C. In the present study the patients in HF stage C were more likely to be older and have a history of AF, which was also the case for the patients in HF stage B. Despite symptoms of HF, no clinical characteristics clearly distinguished HF stages. Other studies have found an increased prevalence of higher age, male gender, obesity, atrial fibrillation and diabetes with higher stages of HF.^{4,20} Thus, the lack of differences in clinical characteristics between HF stages underscores the importance of focusing on symptoms in daily clinical practice in elderly patients with a risk factor for HF. Patients in HF stages A and B are considered patients at risk of overt HF. Previous studies have demonstrated an increased mortality risk and risk of admission for HF in patients in HF stage B.⁴⁻⁶ Further, it should be noticed that patients in HF stage B may have increased plasma concentrations of NT-proBNP and eventually signs of HF, but due to lack of symptoms these patients do not fulfil the criteria for overt HF (HF stage C).

Table 3 Echocardiographic parameters of patients in Copenhagen Heart Failure Risk Study, according to HF stage

	All patients (n=400)	HF stage A (n=177)	HF stage B (n=150)	HF stage C (n=73)	P value for trend
Left ventricle structure and dimension					
IVSd, cm	0.94 (0.83, 1.06)	0.90 (0.81, 1.02)*†	0.97 (0.87, 1.07)	1.00 (0.88, 1.12)	<0.001
LVIDD, cm	4.75 (4.28, 5.21)	4.69 (4.22, 5.05)	4.80 (4.40, 5.31)	4.87 (4.39, 5.36)	0.071
LVEDVi, mL/m ²	39.22 (32.81, 48.39)	39.87 (32.60, 46.05)	40.42 (33.84, 51.68)	38.24 (31.58, 48.16)	0.480
LVEDVi abnormal†, n (%)	5 (1.3)	0 (0.0)	3 (2.1)	2 (2.9)	
LVmass, g/m ²	76.07 (64.53, 90.84)	71.34 (62.71, 82.70)*†	80.29 (68.69, 95.41)	83.95 (66.82, 106.75)	<0.001
LVmass abnormal†, n (%)	40 (10.3)	0 (0.0)*†	21 (14.4)	19 (26.4)	
Abnormal structure or dimension, n (%)	42 (10.5)	0 (0.0)*†	23 (15.3)	19 (26.4)	
Left ventricle systolic function					
LVEF biplane, %	62.62 (55.15, 68.01)	64.49 (58.60, 68.35)*	59.27 (51.98, 65.58)	61.20 (51.60, 69.10)	<0.001
LVEF biplane abnormal†, n(%)	59 (14.8)	0 (0.0)*†	39 (26.0)	20 (27.4)	
LVEF ≥50%, n (%)	360 (90.0)	177 (100.0)*†	124 (82.7)	59 (80.6)	<0.001
LVEF 40%–49%, n (%)	28 (7.0)	0 (0.0)*†	18 (12.0)	10 (13.7)	<0.001
LVEF <40%, n (%)	12 (3.0)	0 (0.0)*†	8 (5.3)	4 (5.5)	0.005
WMSI	1.00 (1.00, 1.00)	1.0 (1.0, 1.0)*	1.0 (1.0, 1.13)	1.0 (1.0, 1.0)	0.002
WMSI abnormal†, n (%)	28 (7.2)	0 (0.0)*†	22 (15.2)	6 (8.3)	
s' septal, cm/s	7.00 (6.00, 8.00)	7.29 (6.47, 8.71)*†	6.85 (5.77, 7.75)	6.39 (5.72, 7.15)	<0.001
s' lateral, cm/s	7.92 (6.69, 9.13)	8.44 (7.06, 9.40)*†	7.74 (6.49, 9.05)	7.22 (6.00, 8.66)	<0.001
GLS, %	-21.00 (-23.03, -18.10)	-21.28 (-23.03, -19.03)*	-20.23 (-22.80, -17.00)	-20.67 (-24.10, -17.87)	0.010
GLS abnormal†, n (%)	27 (7.0)	0 (0.0)*†	19 (13.1)	8 (11.3)	
Abnormal systolic function, n (%)	68 (17.0)	0 (0.0)*†	47 (31.3)	21 (29.2)	
Left ventricle diastolic function					
e' septal, cm/s	6.22 (5.15, 7.54)	6.81 (5.77, 7.98)*†	5.84 (4.66, 6.98)	5.90 (4.86, 7.33)	<0.001
e' septal abnormal†, n (%)	35 (8.8)	0 (0.0)*†	29 (19.6)	6 (8.2)	
E/e' septal	11.51 (9.37, 14.50)	10.51 (8.69, 12.07)*†	13.09 (10.13, 17.26)	13.17 (11.06, 16.19)	<0.001
E/e' septal abnormal†, n (%)	73 (20.6)	0 (0.0)*†	51 (39.5)	22 (41.5)	
E/e' lateral	8.60 (6.81, 11.18)	8.22 (6.59, 9.66)	9.62 (7.12, 12.88)	10.45 (7.32, 13.05)	<0.001
E/e' lateral abnormal†, n (%)	44 (12.4)	0 (0.0)*†	30 (23.3)	14 (26.4)	
E/A	0.87 (0.73, 1.02)	0.87 (0.75, 1.00)	0.87 (0.70, 1.08)	0.83 (0.73, 1.07)	0.003
MV deceleration time, ms	262.08 (223.48, 308.94)	265.03 (225.90, 308.94)	258.36 (219.85, 305.04)	260.69 (224.30, 329.28)	0.857
LAVol index, mL/m ²	29.66 (24.57, 37.41)	26.36 (22.47, 29.87)*†	35.48 (27.80, 43.64)	37.98 (29.56, 45.77)	<0.001
LAVol index abnormal†, n (%)	122 (30.7)	0 (0.0)*†	76 (51.4)	46 (63.0)	

Continued

Table 3 Continued

	All patients (n=400)	HF stage A (n=177)	HF stage B (n=150)	HF stage C (n=73)	P value for trend
Abnormal diastolic function, n (%)	150 (37.5)	0 (0.0)*†	105 (70.0)	45 (62.5)	
Right ventricle function					
TAPSE, mm	22.10 (19.52, 25.17)	22.48 (20.43, 25.17)†	22.10 (19.00, 25.55)	20.75 (18.33, 23.71)	0.006
TR velocity, m/s	2.45 (2.24, 2.62)	2.34 (2.16, 2.52)*†	2.51 (2.30, 2.65)	2.54 (2.29, 2.77)	0.008

Categorical values are presented in percentages, and continuous values are presented as median with 25th and 75th percentiles.

‡ Abnormal echocardiographic parameter according to the study definition.

*P<0.017 for HF stage A vs HF stage B.

†P<0.017 for HF stage A vs HF stage C.

‡ P<0.017 for HF stage B vs HF stage C.

§P<0.017 for HF stage B vs HF stage C.

¶

E, peak velocity of early mitral inflow; GLS, global longitudinal strain; HF, heart failure; IVSd, interventricular septal end diastole; LAVol, left atrial volume; LVEDVi, left ventricle end-diastolic volume indexed; LVEF, left ventricle ejection fraction; LVIDD, left ventricle internal diameter diastole; LVmass, left ventricle mass index; MV, mitral valve; TAPSE, tricuspid annular plane systolic excursion; TR, tricuspid regurgitation; WMSI, Wall Motion Score Index; e', myocardial peak early velocity; e'', myocardial peak early velocity; s', myocardial peak systolic velocity.

Echocardiographic characteristics of HF stages

Echocardiographic parameters were used to identify patients in both HF stages B and C. In this study comparable prevalence of structural and functional abnormalities of the myocardium was found in HF stages B and C, and only LV mass index and e' septal differed between HF stages B and C. Other studies have observed a more pronounced difference in echocardiographic variables, increasingly so with higher stages of HF.^{4 20}

An important finding of this study was the substantial number of patients with HF stage C (n=73), undiagnosed HF. Among patients with HF stage C, 80.6% had HFpEF (LVEF ≥50%), 13.7% had HF with mid-range LVEF (LVEF 41%–50%) and 5.5% had HF with reduced LVEF (LVEF ≤40%). Compared with other studies the echocardiographic findings in this study confirm the increased risk of HF these patients present, with E/e' septal >15 in one-fifth of the patients²¹ and left atrial size ≥35 mL/m² in almost one-third of the patients.²² Finally, a total of 88 patients (n=69 with sinus rhythm and n=19 with AF) had fulfilled the echocardiographic and NT-proBNP criteria for inclusion in the Paragon trial. However, the number was reduced to 27 when including the criteria of New York Heart Association (NYHA) class and treatment with a diuretic.²³ Currently there is no established treatment for HFpEF; instead, focus on the treatment of comorbidities, for example, hypertension and atrial fibrillation, is recommended. In patients with type 2 diabetes, treatment with a sodium glucose inhibitor should be considered.²⁴ Based on post-hoc analyses from the Irbesartan in Heart Failure With Preserved Systolic Function (I-PRESERVE) trial²⁵ and the Treatment of Preserved Cardiac Function Heart Failure with an Aldosterone Antagonist (TOPCAT) trial,²⁶ it may be speculated that HF stage C patients, with relatively low plasma concentrations of NT-proBNP, could benefit from treatment with an aldosterone antagonist.²⁷

Cardiac biomarkers and HF stages

Plasma concentrations of NT-proBNP increased with higher stages of HF. These observations are supported by similar findings in community-based studies of HF stages.^{4 6 7} Elevated plasma concentrations of NT-proBNP are expected in HF stage C, but the elevated concentrations in HF stage B suggest that already in this early stage of HF, in asymptomatic patients, the heart is subject to an increased wall stress. This study is the first to describe that plasma concentrations of copeptin increase with increasing stages of HF. Copeptin is a surrogate marker for vasopressin and reflects neurohumoural activation. It has prognostic value in both acute and chronic HF, and in elderly patients with symptoms of HF.²⁸ We also observed increased plasma concentrations of MR-proADM with higher stages of HF, which also reflects increased neurohumoural activation in higher stages of HF. This finding has previously been observed in stable outpatients with HF.²⁹

In this present study, an increased proportion of patients in both HF stages B and C had detectable

troponin-I, while other studies have found increasing plasma concentrations of troponin with each stage of HF.^{4 6} We found that only one-fifth of the patients had troponin-I above the detection limit, and from our observations elevated troponin-I is more coherent with an abnormal echocardiogram than with HF stages. Troponin-I has been associated with increased mortality risk.³⁰

In contrast, higher stages of HF were not associated with increased plasma concentrations of hsCRP. Coronary microvascular inflammation has been suggested as a driver in the development of HFpEF.³¹ Lack of systemic inflammation or the need for a more sensitive biomarker reflecting inflammation³² may be the reason.

Quality of life (MLHFQ) and Framingham risk score of HF

Symptoms of HF were part of the definition of HF stage C, and higher stages of HF more often presented with a higher NYHA class. Patients in HF stage C also presented with decreased quality of life compared with HF stages A and B, respectively; this aspect has not been described in other studies. The quality of life decreases with severity of overt HF,³³ but it is noteworthy that also patients with undiagnosed HF stage C have a reduced quality of life.

In the present study Framingham 4-year risk score of congestive HF was increased in both HF stages B and C. However, the risk score was low, and the median probability for hospitalisation for congestion within 4 years was only increased by 3% for both men and women. Other studies have reported increased mortality rate with higher stages of HF, most pronounced for HF stages C and D.^{4 5 20} These numbers are hardly comparable since this study has no follow-up data; however, such a difference might occur due to different inclusion criteria.

Methodological considerations and perspectives

Some methodological considerations of this study should be addressed. Patients in this study were selected from three highly specialised sections at the hospital, according to predefined inclusion and exclusion criteria, with the purpose to evaluate whether HF was underdiagnosed in a high-risk population, with no current suspicion of HF by the treating clinician. Therefore, the findings in this cohort do not reflect the general population. Instead, as we intended, the study reflects the risk of HF in an elderly high-risk population.

The definition of abnormal echocardiographic parameters and the definition of HF in the present study need to be discussed. It may be argued that the definition of an abnormal echocardiography is too wide and that this has resulted in an overestimation of the prevalence of HF stages B and C. However, the definition was predefined before data analysis and inspired by current guidelines and previous studies. The ARIC study examined the effect of including diastolic measures in the definition of structural heart disease, and concluded that diastolic measures should be included⁴; similar results have been reported from Olmsted County.⁵ Further, the definition of overt HF (HF stage C) can also be discussed. We used

the European Society of Cardiology's definition and required that symptoms of both dyspnoea or orthopnoea and oedema or use of loop diuretics were present. It was the intention to be conservative and avoid a type I error ('false positive cases'). In contrast, it may be argued that we have made a type II error ('false negative cases') because 30% of the patients in HF stage B were evaluated to be NYHA class II, but did not report symptoms themselves. A bicycle test with invasive measurement of filling pressures might have revealed reduced exercise capacity and increased filling pressures, thereby classifying these patients as HF stage C.³⁴ Functional class was evaluated by a single investigator, and a stress test, for example, 6 min walking test, was not conducted in cases of doubt. Misclassification of functional status may therefore be present, but it should be noted that NT-proBNP increased according to HF stages. Dyspnoea may also be explained by ageing, obesity, physical fitness, chronic obstructive pulmonary disease and AF. Finally, we did not evaluate the prevalence of HF in different subgroups due to the sample size, and our data therefore elucidate the prevalence of early stages of HF in an elderly high-risk population with different risk factors for HF.

CONCLUSION

HF stages B and C are frequent in an elderly high-risk population without known or suspected HF, and especially HFpEF is underdiagnosed. Clinical characteristics are similar in early stages of HF, but the plasma concentrations of NT-proBNP are increasing with higher stages of HF, and troponin-I is more frequently detectable if the echocardiography is abnormal. Finally, higher stages of HF are associated with a decreased quality of life even in patients with undiagnosed HF.

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Competing interests None declared.

Patient consent for publication Parental/guardian consent obtained.

Ethics approval The study was approved by the Committee on Health Research Ethics for the Capital Region of Denmark (H-3-2014-016) and conducted according to the Declaration of Helsinki.

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