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Novel Urinary Peptidomic Classifier Predicts Incident Heart Failure

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Background—Detection of preclinical cardiac dysfunction and prognosis of left ventricular heart failure (HF) would allow targeted intervention, and appears to be the most promising approach in its management. Novel biomarker panels may support this approach and provide new insights into the pathophysiology.

Methods and Results—A retrospective comparison of urinary proteomic profiles generated by mass spectrometric analysis from 49 HF patients, 36 patients who progressed to HF within 2.6 ± 1.6 years, and 192 sex- and age-matched controls who did not progress to HF enabled identification of 96 potentially HF-specific peptide biomarkers. Based on these 96 peptides, the classifier called Heart Failure Predictor (HFP) was established by support vector machine modeling. The incremental prognostic value of HFP was subsequently evaluated in urine samples from 175 individuals with asymptomatic diastolic dysfunction from an independent population cohort. Within 4.8 years, 17 of these individuals progressed to overt HF. The area under receiver-operating characteristic curve was 0.70 (95% Cl, 0.56–0.82); *P*=0.0047 for HFP and 0.57 (0.42–0.72; *P*=0.62) for N-terminal pro b-type natriuretic peptide. Hazard ratios were 1.63 (Cl, 1.04–2.55; *P*=0.032) per 1-SD increment in HFP and 0.70 (Cl, 0.35–1.41; *P*=0.32) for a doubling of the logarithmically transformed N-terminal pro b-type natriuretic peptide.

Conclusions—HFP is a novel biomarker derived from the urinary proteome and might serve as a sensitive tool to improve risk stratification, patient management, and understanding of the pathophysiology of HF. (*J Am Heart Assoc.* 2017;6:e005432. DOI: 10.1161/JAHA.116.005432.)

Key Words: biomarker • heart failure • proteomics • risk stratification

eft ventricular (LV) heart failure (HF) is a clinical syndrome caused by adverse LV structural or functional alterations resulting in impaired ventricular filling and/or ejection and thus in the disability of the heart to pump a sufficient amount of blood to meet the metabolic needs of the body.^{1,2} HF represents an enormous public health and socioeconomic burden.^{1,2} Because of etiological diversity, it is difficult to clearly identify all contributing factors in a clinical setting and to depict the complex pathophysiology by single biomarkers. Difficulties thus arise in diagnosis, risk stratification, and management of HF patients.² A position

paper of the American Heart Association supported research into proteomics as applied to cardiovascular health and disease.³ In line with this recommendation, we developed multidimensional proteomic biomarkers characterizing distinct molecular manifestations of LV dysfunction that may provide additional diagnostic and prognostic value and identify new targets of treatment. We already identified specific peptide biomarker patterns helping in the diagnosis of coronary artery disease,⁴ and asymptomatic LV diastolic^{5,6} and systolic⁷ dysfunction. The present study aimed to extend the findings^{5–7} of these case–control^{5,7} and cross-

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Clinical Perspective

What Is New?

- We identified a novel multidimensional urinary biomarker consisting of 96 peptide fragments and named it heart failure predictor.
- Heart failure predictor predicts progression from asymptomatic left ventricular dysfunction to overt heart failure and is more accurate than a research-optimized N-terminal pro b-type natriuretic peptide assay.

What Are the Clinical Implications?

• Heart failure predictor might serve as a tool to improve risk stratification, patient management, and understanding the pathophysiology of heart failure.

sectional⁶ studies and to investigate whether the urinary proteomic signature might predict progression from asymptomatic LV dysfunction to overt symptomatic HF.

Methods

Study Participants

For discovery of the urinary biomarkers, we investigated 95 patients enrolled in the GS (Generation Scotland: Scottish Family Health Study),⁸ who had either HF at baseline (n=57) or progressed to HF during follow-up (n=38) and 192 sex- and age-matched healthy controls selected from the same cohort. For validation, we studied 175 patients with asymptomatic diastolic LV dysfunction at baseline enrolled in the FLEMENGHO (Flemish Study on Environment, Genes and Health Outcomes^{5,6}), of whom over a 4.8-year period 17 developed HF during follow-up. The diagnosis of HF was ascertained against the records held by general practitioners or hospitals in the catchment area of the FLEMENGHO study.9 Both GS (ethical approval registration number, 10/S1402/20) and FLEMEN-GHO (ML4804) complied with the Helsinki declaration for research in humans¹⁰ and received ethical approval. Permission for health record linkage in GS was obtained from the Privacy Advisory Committee of NHS National Services Scotland. The FLEMENGHO database is registered with the Belgian Privacy Commission (www.privacycommission.be). All participants gave informed written consent.

Definition of LV Dysfunction

In the GS study,⁸ HF was an ICD-coded admission to the hospital for symptomatic HF. In FLEMENGHO,^{5,6} the diagnosis of subclinical LV diastolic dysfunction relied on echocardiography. To ascertain the absence of symptoms, participants completed the London School of Hygiene questionnaires on cardiovascular and respiratory symptoms.¹¹ As described elsewhere,¹² guideline-driven echocardiographic criteria to stage patients with advanced diastolic LV dysfunction leave a large proportion of people unclassified in population studies. We therefore developed age-specific criteria in a healthy reference sample drawn from FLEMENGHO¹² and replicated these criteria in an independent European population study.¹³ Diastolic LV dysfunction included^{12,13}: (1) patients with an abnormally low age-specific transmitral E/A ratio indicative of impaired relaxation, but without evidence of increased LV filling pressures (E/e' \leq 8.5); (2) patients with a mildly-tomoderately elevated LV filling pressure (E/e' > 8.5) and an E/A ratio within the normal age-specific range; and (3) patients with an elevated E/e' ratio and an abnormally low age-specific E/A ratio (combined dysfunction).

Proteomic Urine Sample Analysis

Sample preparation and capillary electrophoresis–mass spectrometry analysis

For proteomic analysis, a 0.7-mL aliquot of stored urine was thawed immediately before use and diluted with 0.7 mL of 2 mol/L urea, and 10 mmol/L NH₄OH containing 0.02% sodium dodecyl sulphate. To remove higher molecular mass proteins, such as albumin and immunoglobulins, the sample was ultrafiltered using Centrisart ultracentrifugation filter devices (20 kDa MWCO; Sartorius, Göttingen, Germany) at 3000 relative centrifugal force units until 1.1 mL of filtrate was obtained. This filtrate was then applied onto a PD-10 desalting column (GE Healthcare, Uppsala, Sweden) equilibrated in 0.01% NH₄OH in HPLC-grade in H₂O (Carl Roth GmbH, Karlsruhe, Germany) to decrease matrix effects by removing urea, electrolytes, salts, and to enrich polypeptides. Finally, all samples were lyophilized, stored at 4°C, and suspended in HPLC-grade H₂O shortly before capillary electrophoresis-mass spectrometry (CE-MS) analyses.¹⁴

CE-MS analyses were performed using a P/ACE MDQ capillary electrophoresis system (Beckman Coulter, Fullerton, CA) on-line coupled to a micrOTOF MS (Bruker Daltonics, Bremen, Germany).^{14,15} The electrospray ionization device (Agilent Technologies, Palo Alto, CA) was grounded, and the ion spray interface potential was set between -4 and -4.5 kV. Data acquisition and MS acquisition methods were automatically controlled by the CE via contact-close-relays. Spectra were accumulated every 3 s, over a mass-to-charge ratio (m/z) ranging from 350 to 3000.

Quality control

Accuracy, precision, selectivity, sensitivity, reproducibility (Figure 1), and stability of the CE-MS have been previously



Figure 1. Proteome coverage of 6 CE-MS runs (A through F) of human urine standards. The molecular mass on a logarithmic scale (0.8-20 kDa on the y-axis) was plotted against the normalized CE migration time (15-45 minutes on the x-axis). Peak height and color represent average signal intensity. The human urine standard is a urine sample from a randomly selected healthy person that is used for quality control.¹⁶ CE-MS indicates capillary electrophoresis–mass spectrometry.

published.^{14,16} Quality control involves daily CE-MS runs of human urine standards.¹⁶ CE has a high reproducibility with at least 70% of peptide recovered compared with only 43% by liquid chromatography. To prevent variability because of carry-over effects from 1 to the next run, capillaries are reconditioned between runs (eg, 1 mol/L NaOH). Figure 1 shows proteome coverage of 6 CE-MS runs of human urine standards. The coefficient of variance estimated from over 600 urine samples collected once daily for over 3 years was 5.8%.¹⁷

Mass spectrometric data processing

Mass spectral peaks representing identical molecules at different charge states were deconvoluted into single masses, using MosaiquesVisu software.¹⁸ Only signals with z of more than 1 observed in a minimum of 3 consecutive spectra with a signal-to-noise ratio of at least 4 were considered. Reference

signals of 1770 urinary polypeptides were used for CE-time calibration by locally weighted regression. For normalization of analytical and urine dilution variances, signal intensities were normalized relative to 29 "housekeeping" peptides.^{19,20} The obtained peak lists characterize each polypeptide by its molecular mass, normalized CE migration time, and normalized signal intensity. All detected peptides were deposited, matched, and annotated in a Microsoft SQL database, allowing further statistical analysis.²¹ For clustering, peptides in different samples were considered identical, if mass deviation was <50 ppm. CE migration time was controlled to be below 0.35 minutes after calibration.

Sequencing of polypeptides

HF biomarkers were in silico assigned to the previously sequenced peptides from Human Urinary Proteome Database, version 2.0.²² Peptides from this database were sequenced,

as described elsewhere.^{23,24} Briefly, urinary peptides were fragmented using different tandem mass spectrometric techniques with a prior separation step with CE or HPLC. Fragmentation spectra were matched to the protein sequences from up-to-date public databases (IPI, NCBI Reference Sequence Database and Uniprot), using MS/MS search engines MASCOT (Matrix Sciences Ltd., London, UK) and OMSSA (National Center for Biotechnology Information, Bethesda, MD). In matching, we accounted for urinary proteins posttranslational modifications, such as hydroxylation of lysine and proline, and specific MS characteristics. Peptide sequences from liquid chromatography–MS analyses were verified by the comparison of experimental and theoretical CE migration time, which is dependent on the number of basic and neutral polar amino acids.

Identified HF-specific urinary peptides were combined into a single multidimensional classifier called Heart Failure Predictor (HFP), using the support vector machine-based MosaCluster software, version 1.7.0.²⁵ MosaCluster calculates classification scores based on the amplitudes of the selected biomarkers. Classification is performed by determining the Euclidian distance (defined as the support-vector machine classification score) of the vector to a maximal margin hyperplane. The parameters for derivation of the HFP classifier were 6.4 for C, 0.008192 for gamma, and 0.001 for epsilon. In sensitivity analyses, we forced sex and age into the computations of the classification scores.

Other Measurements

Hypertension was a blood pressure of at least 140 mm Hg systolic or 90 mm Hg diastolic or use of antihypertensive drugs. Venous blood samples were drawn after at least 6 hours of fasting for measurement of plasma glucose and serum total and high-density lipoprotein cholesterol and serum creatinine. We derived the estimated glomerular

 Table 1. Baseline Characteristics of Cases and Controls Nested in the GS

Characteristic	HF at Baseline	Incident HF	Healthy Controls		
Number	57	38	192		
Number of subjects, %					
Women	18 (31.6)	12 (31.6)	61 (31.3)		
Hypertension	46 (80.7)	26 (68.6)	157 (81.8)		
Diabetes mellitus	9 (15.8)	3 (7.9)	8 (4.2)		
Obesity	23 (41.1)	10 (26.3)	42 (22.0)		
Mean (SD) of characteristic					
Age, y	69±10*	64±11	64±9		
Body mass index, kg/m ²	29±5*	28±5	27±5		
Waist-to-hip ratio	0.94±0.08*	0.91±0.07	0.91±0.09		
Blood pressure, mm Hg					
Systolic pressure	136±23	143±21	141±17		
Diastolic pressure	78±13*	81±12*	82±10		
Heart rate, beats per minute	68±12	70±14	67±11		
Biochemical data					
Serum creatinine, µmol/L	94±28*	85±19*	80±13		
eGFR, mL/min per 1.73 m ²	74±22*	80±17*	85±15		
Total cholesterol, mmol/L	4.3±1.0*	4.9±1.4*	5.2±1.1		
HDL cholesterol, mmol/L	1.2±0.4*	1.3±0.4*	1.4±0.4		
Plasma glucose, mmol/L	5.8±2.2*	5.2±1.6*	5.1±1.3		
NT-proBNP, pg/mL	278 (93–774)*	132 (69–242)*	48 (32–72)		

Values are mean (\pm SD) or geometric mean (interquartile range). Hypertension was an office blood pressure of \geq 140 mm Hg systolic, \geq 90 mm Hg diastolic, or use of antihypertensive drugs. Diabetes mellitus was a self-reported diagnosis, a fasting glucose level of at least 7 mmol/L, or use of antidiabetic agents. Obesity was a body mass index of \geq 30 kg/m². For NT-proBNP, values are geometric mean (interquartile range). eGFR indicates estimated glomerular filtration rate; HDL, high-density lipoprotein; HF, heart failure; NT-proBNP, N-terminal pro-b-type natriuretic peptide.

*Indicates a difference (P≤0.05) between cases and controls.



Figure 2. Flow chart illustrating the peptides retained in the generation of HFP. HFP indicates Heart Failure Predictor.

filtration rate from serum creatinine by the Modification of Diet in Renal Disease formula.²⁶ Diabetes mellitus was a selfreported diagnosis, a fasting glucose level of at least 7 mmol/ L, or use of antidiabetic agents.²⁷ In the GS study,⁸ N-terminal proatrial natriuretic peptide (NT-proBNP) was measured using an automated ELISA assay (Roche Diagnostics, Basel, Switzerland) with an interassay coefficient of variation of <3%. The lower limit of detection was 5 pg of NT-proBNP per mL. In the FLEMENGHO study,^{12,13} NT-proBNP was measured in plasma by a competitive enzyme immunoassay for research use (Biomedica Gruppe, Vienna, Austria).²⁸ The interassay and intra-assay variations were lower than 15%. The lower detection limit was 5 pmol of NT-proBNP/L. The standard range provided by the manufacturer of the enzyme

immunoassay is from 0 to 1000 pmol/L (median, 208 pmol/L; 95% percentile, 300 pmol/L).

Statistical Analysis

We compared means and proportions characterizing the study participants at baseline by Student *t* test and Fisher exact test, respectively. We compared urinary peptide levels with a detectable signal in at least 30% of participants by the nonparametric Wilcoxon rank sum test. Unadjusted *P* values were calculated using the normal approximation of the Wilcoxon test statistic. In the GS study, we applied the Benjamini-Hochberg approach with the false discovery rate set at 5%.²⁹ We used Cox regression to compute hazard ratios

ID	Sequence	Protein Name	Accession Number	BAS	INC	Overlap
107929	DAaHKSEVAHRFKDLGEENFKALVL	Serum albumin	P02768	+28.1	+44.5	
2505	SpGEAGRpG	Collagen a-1(I) chain	P02452	-2.2	-1.3	2
2659	DDGEAGKpG	Collagen a-1(I) chain	P02452	-2.1	-1.2	2
5675	DGKTGPpGPA	Collagen a-1(I) chain	P02452	-2.1	-1.2	
14906	DGRpGPpGPpG	Collagen a-1(I) chain	P02452	-2.7	-1.4	2
16779	ApGDRGEpGPP	Collagen a-1(I) chain	P02452	-2.8	+1.3	
17694	ApGDRGEpGpP	Collagen a-1(I) chain	P02452	-2.8	-1.3	2
21365	PpGEAGKpGEQG	Collagen a-1(I) chain	P02452	+1.8	-1.0	2
23697	DDGEAGKpGRpG	Collagen α-1(I) chain	P02452	-2.1	-1.2	1
28561	SpGPDGKTGPpGPA	Collagen a-1(I) chain	P02452	-2.9	-1.5	1,2
30575	SpGSpGPDGKTGPp	Collagen a-1(I) chain	P02452	-3.3	-1.2	
32171	ApGDRGEpGPpGPA	Collagen a-1(I) chain	P02452	-1.5	-1.2	1,2
35339	ApGDRGEpGPpGPAG	Collagen a-1(I) chain	P02452	-1.4	-1.1	1,2
42594	VGPpGpPGPPGPPS	Collagen a-1(I) chain	P02452	-1.8	-1.2	
43442	VGPpGPpGPpGPPS	Collagen a-1(I) chain	P02452	-1.4	-1.1	
50638	PpGPpGKNGDDGEAGKP	Collagen α -1(I) chain	P02452	-2.1	+1.1	
51175	EGSpGRDGSpGAKGDRG	Collagen a-1(I) chain	P02452	-2.3	+1.1	
51875	VGPpGPpGPpGPPSAG	Collagen a-1(I) chain	P02452	+1.6	+1.1	
62504	TGPIGPpGPAGApGDKGESGP	Collagen a-1(I) chain	P02452	+2.1	+1.2	
63209	EGSpGRDGSpGAKGDRGET	Collagen a-1(I) chain	P02452	-2.0	-1.1	2
65257	SGEpGApGSKGDTGAKGEpGP	Collagen a-1(I) chain	P02452	+1.7	+3.2	2
72896	SGEpGApGSKGDTGAKGEpGPVG	Collagen a-1(I) chain	P02452	+1.2	+1.4	
75846	GPpGEAGKpGEQGVpGDLGApGP	Collagen a-1(I) chain	P02452	+1.2	+1.3	
77018	DGQPGAKGEpGDAGAKGDAGPPGp	Collagen a-1(I) chain	P02452	+1.2	+1.4	
78073	AEGSpGRDGSpGAKGDRGETGPA	Collagen a-1(I) chain	P02452	-1.1	-2.1	
81457	IGPpGPAGApGDKGESGPSGPAGPTG	Collagen a-1(I) chain	P02452	-1.1	-1.8	
82234	IGPpGPAGApGDkGESGPSGPAGPTG	Collagen a-1(I) chain	P02452	-1.2	-2.6	
87460	KGNSGEPGApGSKGDTGAKGEPGPVG	Collagen a-1(I) chain	P02452	+1.5	+1.9	
99808	LTGPIGPPGpAGApGDKGESGPSGPAGPTG	Collagen α -1(I) chain	P02452	-1.2	-1.2	
118163	LTGSpGSpGpDGKTGPPGPAGQDGRPGPpGppG	Collagen α -1(I) chain	P02452	-1.1	-1.5	
36769	DGPpGRDGQpGHKG	Collagen α -2(I) chain	P08123	-1.2	-2.1	
48093	GpAGPRGERGPpGESGA	Collagen α -2(I) chain	P08123	+1.2	+2.0	2
110240	LKGQpGApGVKGEpGApGENGTPGQTGARG	Collagen α -2(I) chain	P08123	+7.7	+2.2	
114086	TGEVGAVGPpGFAGEKGPSGEAGTAGPpGTpGP	Collagen α -2(I) chain	P08123	+1.3	+2.0	
18988	DGESGRpGRpG	Collagen α -1(III) chain	P02461	-1.2	-3.0	
28747	SpGERGETGPpGP	Collagen α -1(III) chain	P02461	+1.1	+1.4	1
30699	DGApGKNGERGGpG	Collagen α -1(III) chain	P02461	-1.0	-2.4	2
36784	DGVPGKDGPRGPTGP	Collagen α-1(III) chain	P02461	+1.0	+2.0	2
38798	GLpGTGGPpGENGKpG	Collagen α-1(III) chain	P02461	-1.1	-2.1	2
49295	ApGGKGDAGApGERGPpG	Collagen α -1(III) chain	P02461	+1.2	+1.8	
61304	GLpGTGGPpGENGKPGEPGp	Collagen α -1(III) chain	P02461	+1.5	+3.6	2
61945	GLpGTGGPpGENGKpGEPGp	Collagen α -1(III) chain	P02461	+1.6	+2.6	2

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Continued

Table 2. Continued

ID	Sequence	Protein Name	Accession Number	BAS	INC	Overlap
64887	GApGApGGKGDAGApGERGPpG	Collagen α -1(III) chain	P02461	+1.3	+2.4	
107460	KNGETGPQGPPGPTGPGGDKGDTGPpGpQG	Collagen α -1(III) chain	P02461	-1.2	-1.7	1
84484	pGFPGAQGEPGSQGEpGDpGLpGP	Collagen α -2(IV) chain	P08572	+1.6	+2.8	
30500	GApGLAGpAGpQGpS	Collagen α -1(VII) chain	Q02388–2	+1.7	+2.1	
86029	PpGppGPpGVPGSDGIDGDNGPPGK	Collagen α -2(IX) chain	H0Y409	+1.3	+2.0	
129940	DVGSYQEKVDVVLGPIQLQTPPRREEEPR	Deleted in malignant brain tumors 1 protein	Q9UGM3	+1.4	+2.0	
98089	DEAGSEADHEGTHSTKRGHAKSRP	Fibrinogen α chain	P02671	+1.6	+2.4	1
103912	DEAGSEADHEGTHSTKRGHAKSRPV	Fibrinogen α chain	P02671	+1.4	+1.9	
17968	DGGGSPKGDVDP	Sodium/potassium-transporting ATPase subunit $\boldsymbol{\gamma}$	P54710	-1.2	-1.4	2
13747	ATKTVGSDTF	Kininogen-1	P01042-2	+1.1	-2.8	
67263	GSGGSSYGSGGGSYGSGGGGGGGGGG	Keratin; type II cytoskeletal 1	P04264	-1.1	-1.6	
59368	FGASAGTGDLSDNHDIIS	Vesicular integral-membrane protein VIP36	Q12907	+1.7	+3.0	
73434	KDQGGYTmHQDQEGDTDAG	Microtubule–associated protein τ ; MAPT	P10636	-1.3	-1.7	1
87692	EDPQGDAAQKTDTSHHDQDHP	Short peptide from AAT	G3V387	-1.3	-2.1	
73015	ELTETGVEAAAASAISVARTL	Plasma protease C1 inhibitor	P05155	+1.9	+5.7	
111426	IPVKQADSGSSEEKQLYNKYPDAVAT	Osteopontin	P10451	+1.9	+1.5	
118694	IPVKQADSGSSEEKQLYNKYPDAVATW	Osteopontin	P10451	+2.9	+7.1	

The analysis of 49 cases with HF at baseline (BAS), 36 cases with incident HF (INC), and 192 controls identified 59 differentially excreted peptides that could be sequenced. The accession number is the identifier in the UniProtKB database (www.uniprot.org). BAS and INC are fold changes of amplitude comparing heart failure cases at baseline (BAS) and incident heart failure at follow-up (INC) to normal controls, respectively. The differential excretion was computed as (amplitude cases×frequency)/(amplitude control×frequency) or as (amplitude controls×frequency)/(amplitude cases×frequency) for upregulated (+) and downregulated (-) proteins in cases vs controls, respectively. Amplitude refers to the average mass spectrometric signal and frequency to the number of individuals with a detectable signal. Overlap refers to the peptide fragments also included in the previously published HF1 (1; reference 5) and HF patients with reduced ejection fraction (2; reference 7) classifiers. HF indicates heart failure.

and to determine clinical characteristics relevant for progression to overt HF. We identified covariables to be retained by a backward elimination with the *P* value set at 0.1. Variables with physiological relevance that were not retained by the stepdown procedure were combined in a propensity score³⁰ derived by regressing HFP on covariables, including sex, body mass index, mean arterial pressure, heart rate, LV mass index, treatment with inhibitors of the renin system, and use of β -blockers. To account for the small sample size in the replication sample (FLEMENGHO^{5,6}), we applied Firth regression.³¹ In FLEMENGHO participants,^{5,6} we evaluated the discriminatory performance of HFP by constructing the receiver operating characteristic curve and calculating the area under the receiver operating characteristic curve.

Results

Design of HFP in the GS Study

Cases were 57 patients with overt HF (ICD10 code, I50.1) present at baseline and 38 patients who over a 5-year followup period progressed to symptomatic HF requiring hospitalization (median time to event, 2.9 years). Among cases with incident HF, 19 (50%) had elevated levels of NT-proBNP (>125 pg/mL) at baseline. Controls were 192 sex- and agematched healthy individuals with normal NT-proBNP level (Table 1).

Impaired renal function is a potential confounder in urinary biomarker discovery.³² We therefore excluded the 10 patients with an estimated glomerular filtration rate of <45 mL/min per 1.73 m² (stage 3B according to the National Kidney Foundation Kidney Disease Outcomes Quality Initiative guideline [www.kidney.org]) from biomarker discovery, leaving 49 with overt HF at baseline and 36 with incident HF. The total number of detected peptide fragments was 5616, but only 1380 (24.6%) with a signal in at least 30% of study participants were analyzed. Of these 1380 peptides (Figure 2), 722 were not sequenced and 658 were sequenced. The sequenced peptides enabled identification of 157 parent proteins. Comparison of cases and controls with the false discovery rate set at 5% identified 96 potential peptide biomarkers, of which 59 were characterized by sequence and posttranslational modification (Table 2). The majority of the sequenced peptides originated from the extracellular matrix and were fragments of collagen I (n=33), III (n=10), as well as collagen IV, VII, and IX (each n=1). Other peptides originated

from α -1-antitrypsin, fibrinogen α (n=2), kininogen-1, microtubule-associated protein τ , osteopontin (n=2), plasma protease C1 inhibitor, and serum albumin (Table 2).

To reduce overfitting in the support vector machine modeling, for the generation of a novel multidimensional classifier for HF, we included all available 95 cases, irrespective of their renal function. The resulting classifier, HFP (threshold level -0.22), allowed correct discrimination of 57 patients with HF at baseline and 38 patients with incident HF versus 192 controls with 100% accuracy upon complete take-

Characteristic	Cases	Controls		
Number	17	158		
Number of subjects, %				
Women	11 (64.7)	90 (57.0)		
Hypertension	13 (76.5)	128 (81.0)		
Diabetes mellitus	3 (17.7)	13 (8.2)		
Obesity	4 (23.5)	52 (32.9)		
Mean (SD) of characteristic				
Age, y	72±6*	64±13		
Body mass index, kg/m ²	28±4	28±4		
Waist-to-hip ratio	0.89±0.08	0.90±0.08		
Blood pressure, mm Hg				
Systolic pressure	142±19	143±19		
Diastolic pressure	76±9*	82±10		
Heart rate, beats per minute	55±11*	63±11		
Biochemical data				
Serum creatinine, μ mol/L	85±19	87±21		
eGFR, mL/min per 1.73 m ²	71±16	72±14		
Total cholesterol, mmol/L	5.4±0.8	5.5±1.0		
HDL cholesterol, mmol/L	1.5±0.3	1.4±0.3		
Plasma glucose, mmol/L	5.6±1.9	5.2±1.1		
NT-proBNP, pmol/L	269 (251–432)	245 (166–389)		
Echocardiography data				
LVEF, %	68±9	70±9		
e' peak, cm/s	7.5±1.7	7.7±1.9		
E/e'	9.9±2.6	9.2±2.9		

 Table 3. Baseline Characteristics of Cases and Controls in the FLEMENGHO Study

Values are mean (\pm SD) or geometric mean (interquartile range). Hypertension was an office blood pressure of \geq 140 mm Hg systolic, \geq 90 mm Hg diastolic, or use of antihypertensive drugs. Diabetes mellitus was a self-reported diagnosis, a fasting glucose level of at least 7 mmol/L, or use of antidiabetic agents. Obesity was a body mass index of \geq 30 kg/m². For NT-proBNP, values are geometric mean (interquartile range). eGFR indicates estimated glomerular filtration rate; FLEMENGHO, Flemish Study on Environment, Genes and Health Outcomes; HDL, high-density lipoprotein; HF, heart failure; LVEF, left ventricular ejection fraction; NT-proBNP, N-terminal pro-b-type natriuretic peotide.

*Indicates a difference (P≤0.05) between cases and controls.

1-out cross-validation. HFP had 11 (11.5%) peptide fragments in common with the earlier published classifiers HF1 developed in hypertensive patients with asymptomatic LV diastolic dysfunction^{5,6} and shared 22 (22.9%) peptides with HFrEF103 derived in HF patients with reduced ejection fraction.⁷ Twenty-two of these common peptides have a known sequence (Table 2). Forcing sex and age into the computations of the classification scores did not affect the performance of HFP.

Replication of HFP in the FLEMENGHO Study

The prognostic utility of HFP was assessed in the independent FLEMENGHO cohort by applying the classifier onto the proteome profiles of 175 individuals with asymptomatic LV diastolic dysfunction but without previous coronary events (Table 3). Median follow-up was 4.7 years (interguartile range, 4.5–5.1; range, 1.1–8.4 years) in 158 patients who did not progress to HF and 5.0 years (interquartile range, 4.4-6.2; range, 3.4-7.2) in 17 patients who developed overt diastolic HF (P=0.20). In stepdown Cox regression, age was the only covariable retaining significance. With adjustment for age and the propensity score, the hazard ratio for HF associated with HFP in Firth regression was 1.64 (Cl, 1.05-2.53; P=0.029) for a 1-SD increment. The corresponding hazard ratio for logarithmically transformed NT-proBNP was 0.70 (Cl, 0.36-1.38; P=0.31). In similarly adjusted models including both biomarkers, the hazard ratios were 1.63 (Cl, 1.04-2.55; P=0.032) for HFP and 0.70 (Cl, 0.35-1.41; P=0.32) for NT-proBNP.

The area under the receiver operating characteristic curve for NT-proBNP was 0.57 (95% Cl, 0.42–0.72; P=0.62). For HFP, the area under the receiver operating characteristic curve was 0.70 (Cl, 0.56–0.82; P=0.0047; Figure 3). The performance of the classifier made up of only type I collagens fragments (n=33) generated an area under the receiver operating characteristic curve of 0.60 (Cl, 0.44–0.76; P=0.21).

Discussion

We identified urinary peptide biomarkers with prognostic value for the progression from asymptomatic LV dysfunction to overt HF. The ensuing multidimensional classifier HFP surpassed a research-optimized NT-proBNP assay in the prediction of progression to symptomatic HF. NT-ProBNP, an inactive fragment of the cleaved pro-BNP molecule, is the guideline-endorsed state-of-the-art clinical marker to confirm HF diagnosis.³³

From a mechanistic point of view, HFP extensively depicts specific excretory molecular phenotypic alterations associated with progressive LV dysfunction. Fragments of fibrillar



Figure 3. Receiver operating characteristic curve for the HFP score factors and NT-proBNP values of the comparison between patients with preclinical left ventricular diastolic dysfunction who did and did not progress to overt heart failure in the FLEMENGHO cohort. FLEMENGHO indicates the Flemish Study on Environment, Genes and Health Outcomes; HFP, Heart Failure Predictor; NT-proBNP, N-terminal pro b-type natriuretic peptide.

type I and III collagen, important components of the myocardial extracellular matrix,³⁴ predominantly make up the proteomic urinary signature associated with HF. These observations are in line with altered collagen synthesis,³⁵ chemical/enzymatic cross-linking,³⁶ and/or turnover by different proteases^{35,37} as the mechanisms underlying the perturbed LV mechanics and geometry^{38,39} and progression to HF.⁴⁰ Renal dysfunction secondary to HF³² might also have contributed to the urinary peptide excretion pattern as captured by HFP.

In addition to collagens, the biomarker pattern included peptide fragments from α -1-antitrypsin and osteopontin, which showed an elevated differential excretion in cases (Table 2). Levels of α -1-antitrypsin progressively increase across the New York Heart Association classes of HF and correlate with B-type natriuretic peptide.41 This might be a compensatory mechanism for the loss of anti-protease activity as a consequence of oxidative stress. The presence of a kininogen-1 fragment in HFP indicated that alterations in kininogen-1 and therefore kinins, its cleavage products, may also be relevant for the diagnosis and prediction of HF. Kinins, such as bradykinin (kallidin-I) and lysyl-bradykinin (kallidin-II), are potent vasoactive and inflammatory peptides acting through the formation of nitric oxide radicals and prostacyclin.⁴² Inhibition of kinin degradation by angiotensin-converting enzyme inhibitors increases LV ejection fraction and decreases the LV end-diastolic volume,⁴³ thereby underscoring the relevance of kinins in the pathophysiology of HF. Moreover, in line with the observed increased HF-related excretion of osteopontin and altered excretion of collagen type I and III, López and coworkers demonstrated that elevated expression of osteopontin in HF patients correlated with collagen cross-linking lysyl oxidase and insoluble collagen.⁴⁴ Combined with NT-proBNP, osteopontin improves the diagnosis of acute heart failure and refines risk stratification.⁴⁵

Conclusions

HFP is a novel biomarker derived from the urinary proteome and might serve as a tool to improve risk stratification, patient management, and understanding of the pathophysiology of HF. While the prognostic utility of HFP has been validated in a fully independent cohort, our study must be interpreted within the context of its limitations. First, the number of patients progressing to overt HF in the validation cohort was relatively small (9.7%), thus limiting the statistical assessment of the prognostic utility. However, we applied Firth regression as a bias-corrected approach to conventional Cox regression. Second, we derived HFP from peptides with a detectable signal in at least 30% of study participants. Incomplete data might be perceived as a weakness. However, ignoring biomarkers with missing values wastes potentially important information, explaining why in proteomic studies missing values of 50% or more are commonly accepted without consensus about the threshold to be applied. Moreover, the 30% threshold in this article is in keeping with our previously published peer-reviewed research. Third, the fact that not all identified polypeptides were sequenced impedes to some extent the insight into the pathophysiological mechanisms underlying HF. Finally, our current study cannot prove the origin of the urinary collagen fragments. However, we are now running proteomics on biopsies taken from explanted (diseased) and implanted (healthy) hearts during cardiac transplantation surgery in an attempt to prove that the urinary and tissue proteomic signatures are similar (http://erc.europa.e u/projects-and-results/erc-funded-projects/prophet). In spite of these limitations, our study underscores the diagnostic and prognostic power of a multidimensional biomarker approach. Further studies are necessary to reach the high level of evidence sufficient to establish HFP as a clinically valuable test.

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Disclosures

Koeck and Nkuipou-Kenfack are employees of Mosaiques-Diagnostics GmbH. Mischak is the CEO of Mosaiques-Diagnostics GmbH. None of the other authors declared a conflict of interest.

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Novel Urinary Peptidomic Classifier Predicts Incident Heart Failure

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