Cystatin C Identifies Patients with Stable Chronic Heart Failure at Increased Risk for Adverse Cardiovascular Events

Dupont et al: Cystatin C in Stable Heart Failure

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Abstract

Background—Renal function is a strong predictor of adverse events in heart failure. Current renal function measures are imperfect and Cystatin C (CysC) is promoted as a better marker of glomerular filtration rate (GFR). This study compares the prognostic utility of CysC and derived GFR estimates with other measures of renal function in patients with chronic heart failure.

Methods and Results—We measured serum CysC levels in 823 heart failure patients undergoing coronary angiography with follow-up of major adverse cardiovascular events (MACE = death, myocardial infarction, stroke). Cystatin C levels strongly correlated with creatinine ($r = 0.73$), blood urea nitrogen ($r = 0.70$), and eGFR$_{MDRD}$ ($r = -0.62$) (all $p < 0.001$). However, the correlation was lower in eGFR $\geq 60$ml/min/1.73m$^2$. CysC-based measures significantly improved areas under the ROC curve for the prediction of MACE, especially in eGFR $\geq 60$ml/min/1.73m$^2$ ($p < 0.01$). Net reclassification improvement was 22.2% ($p < 0.001$) in this group. CysC remained an independent predictor of MACE ($p < 0.001$) after adjustment for traditional risk factors and BNP.

Conclusions—Cystatin C is an independent predictor of adverse events in chronic heart failure. It adds prognostic value to creatinine, particularly in patients with “preserved” renal function.

Key Words: heart failure, kidney, prognosis
Although risk prediction in individual patients with heart failure remains challenging, many factors have shown to be predictive of adverse long-term outcomes. Renal dysfunction and elevated natriuretic peptides are among the strongest predictors. Renal insufficiency has often been considered as a reduction in glomerular filtration rate (GFR), and is traditionally quantified based on serum creatinine (sCr) levels or equations that are derived from sCr measurements. However, the prognostic utility of these sCr-based estimates in the chronic heart failure population remains inferior to GFR, directly quantified by $^{125}$I-iodotamale clearance, suggesting that confounders exist. Nevertheless, cardiovascular risk is believed to be significantly elevated below an estimated GFR of 60 ml/min/1.73m$^2$ based on these equations.

In recent years, cystatin C (CysC) has emerged as a sensitive measure of renal function. Cystatin C is a cysteine protease inhibitor produced by nearly all nucleated human cells at a fairly constant rate (housekeeping gene) and is subsequently circulated in the bloodstream. Compared to creatinine, levels of CysC are less influenced by age, gender, or race. Because of its low molecular weight (13 kDa), it is freely filtered and neither secreted from nor reabsorbed back into the bloodstream (although it is metabolized by the proximal tubular cells). All these properties render CysC a potentially better estimate of GFR than sCr-based estimates. In the chronic kidney disease population, estimates of GFR have incorporated CysC as part of the new equations. CysC has also proven to be a better predictor of mortality and cardiovascular events than sCr-based estimates in different populations studied (elderly, coronary heart disease, acute and chronic heart failure).

Natriuretic peptides have found their way to everyday cardiology practice for diagnostic as well as prognostic purposes in patients with heart failure. Natriuretic peptides reflect...
myocardial stress, and have proven to predict outcome after coronary syndromes, after hospitalization for acute heart failure and in chronic heart failure. Despite BNP levels being somewhat dependent on renal function, BNP and CysC potentially constitute a unique biomarker combination for risk prediction, particularly in the setting of relatively preserved renal function.

The aims of this study were: 1) to examine the prognostic ability of CysC and derived equations, independent of BNP; 2) to examine its incremental value over traditional markers of renal function; and 3) to investigate risk stratification by the combination of BNP and CysC in a chronic stable heart failure population undergoing cardiac evaluation.

Methods

Patient Population and Study Design. We evaluated 823 stable patients with a clinical history of heart failure undergoing elective coronary angiography as part of the GeneBank study. GeneBank was a large prospective study, conducted at the Cleveland Clinic between 2001 and 2006 (NCT 00590200) that established a well-characterized clinical repository with clinical and longitudinal outcomes. Inclusion criteria were: Age ≥ 18 years and planned coronary angiography, There were no specific exclusion criteria. All participants gave written informed consent approved by the Cleveland Clinic Institutional Review Board. At the time of cardiac catheterization, blood samples were collected, processed, and stored at -80°F until analyses.

Renal function was expressed as sCr, urea levels (BUN), CysC, estimated GFR by either the four-variable MDRD equation (eGFRMDRD = 186 x sCr-1.154 x age-0.203 x [0.742 if female] x [1.21 if black]), the CKD-EPI equation for Cystatin C (eGFRCysC = 127.7 x CysC-1.17 x age-0.13 x [0.91 if female] x [1.06 if black]), or the eGFRsCr+CysC equation (eGFRsCr+CysC = 177.6 x sCr-0.65 x CysC-0.57 x age-0.20 x [0.82 if female] x [1.11 if black]). “Preserved” and “impaired” renal
function were defined as \( \text{eGFR}_{\text{MDRD}} \geq 60 \text{ ml/min/1.73m}^2 \) and \( \text{eGFR}_{\text{MDRD}} < 60 \text{ ml/min/1.73m}^2 \) respectively.

The endpoint for this study was a composite of major adverse cardiovascular events (MACE), including all-cause mortality, non-fatal myocardial infarction and non-fatal cerebrovascular accident (CVA). Non-fatal events were defined as MI or CVA in patients who survived at least 48 hours following the onset of symptoms. These clinical outcomes were prospectively ascertained over the ensuing 3 years for all subjects after enrollment, adjudicated and verified by source documentation.

**Cystatin C Assay.** Plasma CysC level was determined with a particle enhanced immunoturbidimetric immunoassay on the Architect ci8200 platform (Abbott Laboratories, Abbott Park, IL, USA). Briefly, latex particles are coated with anti-human CysC antibody and agglutinate with CysC present in the patient’s sample. The result is a change in absorbance which is proportional to the amount of CysC present in the sample. The analytical range spans 0.05mg/L to the highest calibration point. Intra- and inter-assay coefficients are 3.1 and 6% respectively. B-type natriuretic peptide, high-sensitivity C-reactive protein (hsCRP), sCr, BUN, fasting blood glucose and lipid profiles were all measured by clinically-approved assays on the same platform. The intra- and inter-assay coefficients were respectively 0.9% and 0.8% for BUN, 1.7% and 1% for sCr, 2.6% and 3.5% for BNP and 4% and 2.4% for hsCRP.

**Statistical Analyses.** Continuous variables were expressed as mean ± standard deviation if normally distributed or as median and inter-quartile range for non-normally distributed data. Categorical data were summarized as proportions and frequencies. Baseline characteristics between patients with preserved and decreased renal function were compared, using the student t-test, the Wilcoxon rank sum test or the chi-square test as appropriate. Spearman’s rank
correlation method was used as a nonparametric measure of association for correlations between CysC and the different measures of renal function (sCr, BUN, eGFR equations). Areas under the ROC curves (c-statistics) for the prediction of MACE were compared between different measures of renal function and net reclassification improvement was calculated. Univariate Cox proportional-hazards models were used to evaluate the association of the different measures of renal function along other variables, with the composite outcome. Measures of renal function were entered either as continuous variables (logarithmic transformed when non-normally distributed), either categorized in quartiles. Since sCr levels differ substantially between genders, sex-specific quartiles for sCr were used to equalize the distribution of men and women. Afterwards two types of multivariate Cox proportional-hazards models were constructed; a shorter model, using one measure of renal function in combination with BNP, and a longer model with a measure of renal function adjusted for all variables with a p-value < 0.05 in univariate analysis. The 3-year outcome was evaluated by Kaplan-Meier survival analyses, using the aforementioned quartiles. Differences between groups were evaluated with the log-rank test. Statistical significance was set at a 2-tailed probability level <0.05. All statistical analyses were performed using JMP Pro 9.0 (SAS Institute, Cary, NC). All the authors had full access to all of the data in the study, and take responsibility for the integrity of the data and the accuracy of the data analysis. All authors have read and agree to the manuscript as written.

Results

Table 1 illustrates the baseline characteristics of our study population, which included 35% of subjects with LVEF > 50%. In our study cohort, 208 subjects (25%) had impaired renal function as defined by eGFR_{MDRD} < 60ml/min/m^2. Subjects with impaired renal function were
likely older, female, with more co-morbid conditions, and higher median BNP levels (Table 1). Cystatin C levels demonstrated a non-normal distribution (Figure 1), with a median level of 1.11 mg/L (interquartile range of 0.92 - 1.41 mg/L). As expected, CysC had a strong correlation with other markers of renal function. However in the subgroup with preserved renal function, such correlations were less robust with non-CysC-based measures of renal function (Online-only Data Supplement Table). There were no significant differences in baseline CysC levels between those with impaired versus preserved LVEF.

During the 3-year follow-up period, the combined endpoint of MACE occurred in 201 (24%) of the patients, including 20% in the preserved renal function group (121 patients, 132 events: 80 deaths, 41 MI and 11 CVA) and 39% in the impaired renal function group (80 patients, 93 events: 62 deaths, 25 MI and 6 CVA). Areas under the ROC curves for Cys C and Cys C-based GFR estimations are significantly better than eGFR_{MDRD} both in the total cohort and in the subgroup of patients with eGFR_{MDRD} ≥ 60 ml/min/1.73m² but not in patients with eGFR_{MDRD} < 60 ml/min/1.73m² (Table 2). Nevertheless in the eGFR_{MDRD} ≥ 60 ml/min/1.73m² group, net reclassification improvement (compared to eGFR_{MDRD}) was 22.2% with Cys C, 22.8% with eGFR_{CysC} and 22.2% with eGFR_{sCr+CysC} (all P < 0.001). In univariate analysis, elevated levels of all renal measures were significantly associated with higher risks for the combined outcome in the overall cohort, both when entered as a continuous variable and when divided in quartiles (Table 3 and Figure 2). Age, body mass index, diabetes mellitus, coronary artery disease, hemoglobin, hsCRP, and BNP were also significantly associated with risk of future MACE in univariate analysis (data not shown). Increasing quartiles of CysC and decreasing quartiles for eGFR_{CysC} demonstrated progressively higher risk of future MACE (Figure 3). In particular, by adjusting for age, gender, and race in the eGFR_{CysC} equation, there was a better
separation of patients with higher eGFR_{CysC} values. However in an analysis with covariate information (BNP, age, body mass index, history of coronary artery disease, history of diabetes mellitus, hemoglobin, and hsCRP), the difference in C-statistics was not significant anymore.

Figure 2 also demonstrates that, when looking at eGFR_{MDRD}, the risk of MACE seems to increase only when eGFR_{MDRD} < 60 ml/min/1.73m^2, in agreement with classic teaching. But when patients are stratified by CysC or derived equation, the risk of MACE seems to increase earlier and looks more like a continuum. Fifty-two percent (212/409 subjects) in the two higher CysC quartiles and 67% (415/617 subjects) in the 3 lower eGFR_{CysC} quartiles had eGFR_{MDRD} ≥60 ml/min/1.73m^2 and were not deemed to be at higher risk of MACE based on their eGFR_{MDRD}. However CysC based measurements showed they were at risk. This is particularly apparent when comparing different measures of renal function in those with eGFR_{MDRD} ≥60ml/min/1.73m^2, indicating a progressive increased risk for future MACE with increasing quartiles of CysC and decreasing quartiles of eGFR_{CysC} (p= 0.002 and 0.001 respectively) but not other measures (Online-only Data Supplement Figure).

In different multivariate Cox proportional hazard models (each containing one measure of renal function), all renal function measures (except urea) were associated with higher risk of future MACE when adjusted for BNP (model 1) and other risk factors (model 2) (Table 3).

Figure 4 depicts the additive value of CysC and BNP in overall risk prediction. Patients in the highest tertile of both variables had a 45% risk to experience death, myocardial infarction, or stroke after 3 years as compared to only 12% when both variables were in the lowest tertile (p<0.0001).
Discussion

There is a wealth of literature demonstrating the ability of CysC as a reliable and sensitive measure of renal dysfunction, and we and others have demonstrated the prognostic value of CysC in the setting of acute heart failure as well as in stable chronic heart failure. However, the majority of the studies have been limited to relatively small sample sizes that precluded adequately powered subgroup analyses with covariate adjustments to shed the light on how to best utilize this measure of renal function in the clinical setting. As the largest study looking at the clinical utility of CysC in stable chronic heart failure using contemporary statistical methodologies, we observed that elevated CysC may identify a large number of patients at increased cardiovascular risk with what was previously labeled as "preserved renal function" (eGFR_{MDRD} > 60ml/min/1.73m^2). Combining CysC with BNP provides excellent risk stratification, with an incidence of 12% of MACE at 3 years when both variables were in the lowest tertiles, versus an incidence of 45% of MACE at 3 years when both variables were in the highest tertiles. Taken together, our results imply that there is incremental prognostic value of CysC in patients with chronic stable heart failure, particularly in those with relatively preserved renal function based on creatinine-based estimates. This demonstrates that even in the setting of mild renal dysfunction there appears to be heightened risk for adverse cardiovascular outcomes.

Our findings corroborated prior studies that suggested CysC may improve overall risk prediction in heart failure. The limitations of sCr’s reciprocal relationship with GFR are well known, largely due to variable production (as a function of age, gender, muscle mass, and ethnicity), diet and tubular secretion. Creatinine-based equations try to correct for this by including determinants of muscle mass (age, weight, race, gender) in their formulas. The equations were developed in patients with chronic kidney disease where they are reasonably
accurate\textsuperscript{20}. It remains intriguing to see, in this and other studies\textsuperscript{21}, that CysC is consistently better in risk prediction whereas it is either equivalent or only slightly improves eGFR\textsubscript{MDRD} estimation\textsuperscript{7}. We see two potential explanations for this observation. First, CysC and derived equations are significantly better than sCr derived equations in estimating GFR in the non-chronic kidney disease population. This is very well possible because in these non-chronic kidney disease populations (elderly, heart failure, coronary patients), mean GFR is higher, patients are older and have more comorbidities. Current creatinine-based GFR equations tend to be less accurate in the higher GFR ranges as nicely demonstrated in the diabetic population and in the heart failure population\textsuperscript{4, 22, 23}. In addition, in the aforementioned settings, the dependence of creatinine on other factors than GFR (muscle mass, diet and general health condition) becomes more and more important. These influences are not all captured by correcting for age, gender and race. A second possibility is that CysC represents other factors aside renal impairment. There is indeed some mechanistic evidence to suggest that CysC, being an inhibitor of cysteine proteases, may be intricately involved in the pathogenesis of atherosclerosis\textsuperscript{24}. In addition it has been shown that the levels of CysC can be influenced by inflammation, glucocorticoid use, and thyroid function\textsuperscript{25}. All this may explain why the incremental prognostic value of CysC is mainly confined to those with relatively “preserved” renal function, whereas the role of measuring CysC in those with already impaired renal function as determined by sCr-based estimates is limited.

We have also applied for the first time the CysC-derived equation for estimating GFR in the heart failure population, and found eGFR\textsubscript{CysC} to be reliably predictive of long-term outcomes (and potentially superior to eGFR\textsubscript{sCr+CysC}). The observation that the eGFR\textsubscript{CysC} equation further improved risk prediction as compared to CysC was unexpected. In fact, the correlation between
CysC and eGFR\textsubscript{CysC} equation is very high (r = 0.98), and the equation only reclassified 14% of patients in a different quartile. However, in addition to a play of chance, it remains possible that adjusting for age, gender and race is important in this population. Interestingly, the addition of creatinine to the CysC-based equation (eGFR\textsubscript{SCr+CysC}) weakened the prognostic ability again. This suggests that, once GFR is estimated on the base of CysC, any potential improvement of GFR estimation by adding sCr is outweighed by its association with muscle mass or general health condition.

One important observation from our study is the fact that one third of subjects in the subset with eGFR\textsubscript{MDRD} \geq 60 \text{ ml/min/1.73m}^2, (or up to one quarter of subjects in our entire study cohort) may demonstrate an elevated CysC level that is associated with heightened risk for future adverse cardiac events. This represents a relatively large population at risk. In this subgroup, CysC and CysC-derived equation were the only measures of renal function that stratified risk of adverse events. Taken together, the present data suggest that a decrease in GFR increases cardiovascular risk throughout its spectrum without a clear “normal” cut-off by sCr or sCr-based equations.

The current usage of CysC in clinical practice is limited despite its approval, clinical availability, and extensive nephrology and epidemiology literature to support its value. This is in part because no prior studies have relied on CysC as an inclusion criterion for treatment decisions, and limited therapeutic studies have provided analyses to test the differential impact of interventions in high versus low CysC subgroups. Nevertheless, the ability to identify a vulnerable population in a cohort of chronic stable heart failure is an opportunity to test therapeutic strategies that may potentially provide incremental benefit to existing therapies. Current clinical trials have utilized sCr-based cut-off values as exclusion criterion either due to
therapeutic contraindications or to remove confounding due to concomitant end-organ
dysfunction that may indirectly affect the demonstration of therapeutic efficacy. In contrast,
natriuretic peptide-based cut-off values, identifying a more at-risk sub-population, have been
increasingly utilized in clinical studies to facilitate the successful demonstration of the benefits
of drug therapy (such as eplerenone in mild heart failure\textsuperscript{26}). At the other end of the spectrum,
post-hoc subgroup analysis has implied that lower rather than higher natriuretic peptide levels
may demonstrate therapeutic benefits of statin therapy in ischemic cardiomyopathy\textsuperscript{27}. Based on
the inter-relationship between CysC and BNP, it is conceivable to identify those with elevated
natriuretic peptides and/or elevated CysC levels to test therapeutic strategies towards combined
or separate targets in the setting of mild heart failure.

**Study Limitations.** The biggest limitation of this study is that there was no direct “gold
standard” measurement of GFR (e.g., inulin or \textsuperscript{125}I-iothalamate clearance), yet recent reports of
relationships between iothalamate-derived GFR measures and CysC or eGFR\textsubscript{MDRD} have been
reported\textsuperscript{17}. Consequently, it remains speculative to state that the CysC-derived equation may
better reflect GFR in the heart failure population. Second, there was some degree of selection
bias as the population studied included patients undergoing elective cardiac catheterization for
symptom evaluation, and there were no prospective outcomes regarding heart failure
hospitalizations. Furthermore, the particle-enhanced turbidimetric immunoassay used to
determine CysC levels may have resulted in values that are 20\% to 30\% higher than in the
common particle-enhanced nephelometric immunoassay method\textsuperscript{28}. Similarly, the eGFR\textsubscript{CysC}
equation was derived with CysC values determined by the nephelometric and not turbidimetric
method, thus resulting in lower absolute eGFR\textsubscript{CysC} values. However, this should not alter the
correlation of CysC or eGFR\textsubscript{CysC} with other measures of renal function, nor its risk predictive
ability. Finally, we were not able to examine how CysC adds to other renal biomarkers (e.g., neutrophil gelatinase-associated lipocalin (NGAL)) that reflect complementary renal processes (e.g. tubular injury).

**Conclusion**

Cystatin C is a strong predictor of adverse events in stable chronic heart failure, independent of traditional risk factors and BNP. Cystatin C and its derived equation to estimate glomerular filtration rate add significant prognostic value to creatinine, predominantly in patients with relatively preserved renal function.

**Sources of Funding**

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**Disclosures**

Drs. Dupont and Wu had no relationships to disclose. Dr. Hazen reports being listed as co-inventor on pending and issued patents held by the Cleveland Clinic relating to cardiovascular
diagnostics. Dr. Hazen reports having been paid as a consultant or speaker for the following companies: Abbott Diagnostics, BG Medicine Inc., Cleveland Heart Lab, Esperion, Lilly, Liposcience Inc., Merck & Co., Inc., and Pfizer Inc. Dr. Hazen reports receiving research funds from Abbott, Cleveland Heart Lab, Liposcience Inc., and Pfizer Inc. Dr. Hazen reports having the right to receive royalty payments for inventions or discoveries related to cardiovascular diagnostics or therapeutics from the companies shown below: Abbott Laboratories, Inc., Cleveland Heart Lab., Esperion, Frantz Biomarkers, LLC, Liposcience Inc., and Siemens. Dr. Tang reports having received research grant support from Abbott Laboratories, Inc.

References


performance of the ckd epidemiology collaboration (ckd-epi) and the modification of diet in renal disease (mdrd) study equations for estimating gfr levels above 60 ml/min/1.73 m2. Am J Kidney Dis. 2010;56:486-495.


Table 1. Baseline Characteristics.

<table>
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<tr>
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<th>Total Cohort (n=823)</th>
<th>eGFR$_{MDRD}$ $\geq$60ml/min/1.73m$^2$ (n=615)</th>
<th>eGFR$_{MDRD}$ &lt;60ml/min/1.73m$^2$ (n=208)</th>
<th>P-value$^a$</th>
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<td>Demographics:</td>
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<tr>
<td>Age (Years)</td>
<td>66 ± 11</td>
<td>65 ± 11</td>
<td>71 ± 9</td>
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<td>Male Gender (%)</td>
<td>60</td>
<td>64</td>
<td>48</td>
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<td>BMI (kg/m$^2$)</td>
<td>29.5 ± 6.7</td>
<td>29.7 ± 6.7</td>
<td>28.9 ± 6.6</td>
<td>0.12</td>
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<td>Co-morbidities:</td>
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<td>Diabetes mellitus (%)</td>
<td>29</td>
<td>23</td>
<td>45</td>
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<tr>
<td>Hypertension (%)</td>
<td>76</td>
<td>72</td>
<td>86</td>
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<tr>
<td>Hyperlipidemia (%)</td>
<td>80</td>
<td>80</td>
<td>80</td>
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<tr>
<td>Current smoker (%)</td>
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<td>13</td>
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<td>CAD (%)</td>
<td>76</td>
<td>74</td>
<td>84</td>
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<td>LVEF (%)</td>
<td>35 (25-55)</td>
<td>35 (25-50)</td>
<td>40 (25-55)</td>
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<td>Laboratory Data:</td>
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<tr>
<td>Urea (mg/dl)</td>
<td>21 (16-27)</td>
<td>19 (15-23)</td>
<td>35 (27-44)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>Creatinine (mg/dl)</td>
<td>0.94 (0.8-1.17)</td>
<td>0.87 (0.76-1.15)</td>
<td>1.41 (1.22-1.81)</td>
<td>&lt; 0.0001</td>
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<td>Cystatin C (mg/dl)</td>
<td>1.11 (0.92-1.41)</td>
<td>1.01 (0.88-1.18)</td>
<td>1.69 (1.42-2.13)</td>
<td>&lt; 0.0001</td>
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<td>eGFR$_{MDRD}$ (ml/min/1.73m$^2$)</td>
<td>77 (60-93)</td>
<td>85 (72-97)</td>
<td>44 (36-53)</td>
<td>&lt; 0.0001</td>
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<td>eGFR$_{CysC}$ (ml/min/1.73m$^2$)</td>
<td>63 (47-80)</td>
<td>71 (58-84)</td>
<td>37 (30-46)</td>
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<td>eGFR$_{Scr+CysC}$ (ml/min/1.73m$^2$)</td>
<td>70 (53-86)</td>
<td>77 (66-90)</td>
<td>40 (32-48)</td>
<td>&lt; 0.0001</td>
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<td>BNP (pg/ml)</td>
<td>300 (118-675)</td>
<td>259 (106-540)</td>
<td>557 (220-1108)</td>
<td>&lt; 0.0001</td>
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<td>Hemoglobin (gr/dl)</td>
<td>13.2 ± 1.8</td>
<td>13.5 ± 1.7</td>
<td>12.4 ± 1.8</td>
<td>&lt; 0.0001</td>
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<td>CRP (mg/dl)</td>
<td>3.8 (1.6-9)</td>
<td>3.2 (1.4-7.8)</td>
<td>5.5 (2.6-14.1)</td>
<td>&lt; 0.0001</td>
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<tr>
<td>ACE-I or ARB (%)</td>
<td>68</td>
<td>68</td>
<td>66</td>
<td>0.49</td>
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<td>Beta-blockers (%)</td>
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<td>Diuretic (%)</td>
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<tr>
<td>Aspirin (%)</td>
<td>62</td>
<td>62</td>
<td>63</td>
<td>0.80</td>
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Statin (%) | 59 | 60 | 57 | 0.36

Abbreviations:
BMI, Body Mass Index; CAD, Coronary Artery Disease; LVEF, left ventricular ejection fraction; eGFR, estimated glomerular filtration rate; BNP, brain natriuretic peptide; CRP, C-Reactive Protein; ACE-I, angiotensin converting enzyme inhibitor; ARB, angiotensin receptor blocker.

§ P-value for difference between eGFR\textsubscript{MDRD} \geq 60\text{ml/min/1.73m}^2 and eGFR\textsubscript{MDRD} < 60\text{ml/min/1.73m}^2.
Table 2. AUC-ROC for different measures of renal function as predictors of MACE.

<table>
<thead>
<tr>
<th></th>
<th>AUC-ROC</th>
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<td>Total Cohort</td>
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<td>eGFR&lt;60ml/min/1.73m²</td>
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<td></td>
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<td>n=615</td>
<td>n=208</td>
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<td>eGFR&lt;sub&gt;MDRD&lt;/sub&gt;</td>
<td>0.615</td>
<td>0.529</td>
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<td>Cys C</td>
<td>0.648</td>
<td>0.600</td>
<td>0.586</td>
</tr>
<tr>
<td></td>
<td>(p=0.07)</td>
<td>(p=0.006)</td>
<td>(p=0.18)</td>
</tr>
<tr>
<td>eGFR&lt;sub&gt;CysC&lt;/sub&gt;</td>
<td>0.652</td>
<td>0.603</td>
<td>0.589</td>
</tr>
<tr>
<td></td>
<td>(p=0.01)</td>
<td>(p=0.001)</td>
<td>(p=0.18)</td>
</tr>
<tr>
<td>eGFR&lt;sub&gt;Cr+CysC&lt;/sub&gt;</td>
<td>0.641</td>
<td>0.581</td>
<td>0.573</td>
</tr>
<tr>
<td></td>
<td>(p=0.02)</td>
<td>(p=0.02)</td>
<td>(p=0.25)</td>
</tr>
</tbody>
</table>

P-values are for comparison with eGFR<sub>MDRD</sub>.

MACE = Death, myocardial infarction or stroke
Table 3. Uni- and multivariate Cox proportional hazard models for different renal measures as predictors of MACE.

<table>
<thead>
<tr>
<th>Log-transformed</th>
<th>UNADJUSTED</th>
<th>MODELL 1§</th>
<th>MODELL 2¶</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hazard Ratio*</td>
<td>P-value</td>
<td>Hazard Ratio*</td>
<td>P-value</td>
</tr>
<tr>
<td>Urea</td>
<td>1.40 (1.22-1.59)</td>
<td>&lt;0.0001</td>
<td>1.25 (1.09-1.42)</td>
</tr>
<tr>
<td>Creatinine</td>
<td>1.34 (1.21-1.48)</td>
<td>&lt;0.0001</td>
<td>1.22 (1.09-1.35)</td>
</tr>
<tr>
<td>eGFRMDRD</td>
<td>0.72 (0.65-0.80)</td>
<td>&lt;0.0001</td>
<td>0.80 (0.72-0.90)</td>
</tr>
<tr>
<td>Cystatin C</td>
<td>1.45 (1.30-1.60)</td>
<td>&lt;0.0001</td>
<td>1.29 (1.15-1.44)</td>
</tr>
<tr>
<td>eGFR_{CysC}</td>
<td>0.69 (0.62-0.77)</td>
<td>&lt;0.0001</td>
<td>0.77 (0.68-0.87)</td>
</tr>
<tr>
<td>eGFR_{sCr+CysC}</td>
<td>0.70 (0.63-0.78)</td>
<td>&lt;0.0001</td>
<td>0.78 (0.70-0.88)</td>
</tr>
</tbody>
</table>

Abbreviations as in Table 1. P-value for log(BNP) <0.001 in every model.

§ Model 1 is adjusted for log-transformed BNP

¶ Model 2 is further adjusted for age, BMI, history of coronary artery disease, history of diabetes mellitus, hemoglobin, and hsCRP.

*Hazard ratios are per 1-standard deviation (Urea 12.7, Creatinine 0.89, eGFRMDRD 26.2, Cystatin C 0.73, eGFR_{CysC} 23.7, eGFR_{sCr+CysC} 24.8)
Figure Legends

Figure 1. **Histogram of serum Cystatin C Concentrations**. Patients with eGFR_{MDRD} < 60ml/min/1.73m^2 are projected in darker shade.

Figure 2. **Forrest Plots (unadjusted) For Quartiles of Different Measures of Renal Function**. Hazards ratio compared to reference for quartiles of urea, cystatin, creatinine (all in mg/dL) as well as estimated glomerular filtration rate equations (all in ml/min/1.73m^2).

Figure 3. **Kaplan-Meier curves for Death/MI/Stroke according to Quartiles of Cystatin C (A) and eGFR_{CysC} (B).**

Figure 4. **Risk Stratification in Patients with Heart Failure According to Tertiles of Cystatin C and BNP**. The risk of the combined outcome (death, myocardial infarction, stroke) increases from 12.3% in patients with both biomarkers in the lowest tertile to 44.8% in patients with both biomarkers in the highest tertile (p<0.0001).
Cystatin C Identifies Patients with Stable Chronic Heart Failure at Increased Risk for Adverse Cardiovascular Events
Matthias Dupont, Yuping Wu, Stanley L. Hazen and W. H. Wilson Tang

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Supplemental Table.

Spearman Correlation coefficients between Cystatin C and other measures of GFR in patients with “preserved” and “impaired” renal function.

<table>
<thead>
<tr>
<th></th>
<th>CORRELATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total Cohort n=823</td>
</tr>
<tr>
<td>Urea</td>
<td>0.70</td>
</tr>
<tr>
<td>Creatinine</td>
<td>0.73</td>
</tr>
<tr>
<td>eGFR&lt;MDRD</td>
<td>-0.79</td>
</tr>
<tr>
<td>eGFR&lt;CysC</td>
<td>-0.98</td>
</tr>
<tr>
<td>eGFR&lt;sCr+CysC</td>
<td>-0.93</td>
</tr>
</tbody>
</table>
Supplemental Figure.

Incidences of Death, Myocardial Infarction, or Stroke according to quartiles of Different Renal Function Measures in the Preserved Renal Function Subgroup (eGFR_{MDRD} \geq 60 \text{ ml/min/1.73m}^2).

1^{st} quartile encompasses lowest values of creatinine, urea, cystatin C, eGFR_{MDRD} and eGFR_{Cys C}. P-values for trend according to Cochran-Armitage.