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EJECTION FRACTION AND MORTALITY: A NATIONWIDE REGISTER BASED COHORT STUDY OF 499,153 WOMEN AND MEN

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ABSTRACT

AIMS

We investigated the sex-based risk of mortality across the spectrum of left ventricular ejection fraction (LVEF) in a large cohort of patients in Australia.

METHODS AND RESULTS

Quantified levels of LVEF from 237,046 women (48.1%) and 256,109 men undergoing first-time, routine echocardiography (2000-2019) were linked to 119,232 deaths (median 5.6 years follow-up). Overall, 17.6% of men versus 8.3% women had an LVEF <50%. An LVEF <40% was associated with the highest crude cardiovascular-related and all-cause mortality at 5-years; ~20-30% and ~40-50%, respectively. Thereafter, actual cardiovascular-related and all-cause mortality at 5-years in both sexes steeply improved to a nadir LVEF of 65.0-69.9% (reference group). Below this LVEF level, the adjusted hazard ratio (HR) for cardiovascular-related mortality for a LVEF of 55.0-59.9% was 1.36 (95% CI 1.16, 1.59; $p < 0.001$) in women and 1.21 (95% CI 1.05, 1.39; $p = 0.008$) in men. In women, an LVEF of 60.0-64.9% was also associated with a HR 1.33 (95% CI 1.16, 1.52; $p < 0.001$) for cardiovascular-related mortality. These associations were most striking women and men aged <65 years and were replicated in those with suspected heart failure (32,403 cases aged 65.2 ± 16.1 years, 57.0% women). For pre-existing heart failure (33,738 cases aged 67.6 ± 16.9 years, 46.5% women), the specific threshold of increased mortality was at and below 50.0-54.9%.

CONCLUSIONS

Among patients investigated for suspected or established cardiovascular disease, we found clinically relevant sex-based differences in the distribution and mortality associated with an LVEF below 65.0-69.9%. Specifically, they suggest a greater risk of mortality at higher LVEF levels among women.

KEY WORDS: Left ventricular ejection fraction; mortality; cardiac function; sex-specific; outcomes

INTRODUCTION

Despite many attempts to identify a suitable alternative, left ventricular ejection fraction (LVEF), typically measured by transthoracic echocardiography, remains the most commonly applied measure of LV systolic function.¹ Expert guidelines recognise specific thresholds of LVEF to define LV dysfunction and increased risk of premature mortality with an LVEF of 52% and 54% measured by the Simpson's biplane method being routinely considered "normal" in men and women, respectively.² Therapies targeting symptomatic patients with heart failure with reduced (<40%) ejection fraction (HFrEF) are well-established.³ However, the definitive treatment of a broader range of patients with less impaired systolic function/more preserved LVEF (including many women^{4, 5}), remains elusive. Reports from the PARAGON-HF Study⁶ and TOPCAT Trial⁷ suggesting differential treatment responses based on sex, reflects the ongoing clinical conundrum on who might benefit from more proactive management and surveillance when presenting with an LVEF >45%.

Remarkably, with a few notable exceptions,⁸⁻¹⁰ there is a paucity of large-scale studies from routine clinical practice examining the relationship between quantified LVEF levels and mortality to address this issue. This critical gap in the literature is particularly evident when considering the predominance of men and a lack of sex-specific data.⁸⁻¹⁰ Previously, the National Echocardiography Database of Australia (NEDA¹¹) has identified clinically important thresholds of mortality risk in respect to pulmonary hypertension,¹² aortic stenosis¹³ and, most recently, diastolic dysfunction.¹⁴ Applying an expanded version of this unique resource, the primary aim of this study was to generate sex-specific data on the distribution of routinely observed LVEF levels and then examine their relationship to the risk of subsequent mortality. Given the overall heterogeneity of the cohort, a secondary aim (where possible) was to examine the pattern of mortality according to LVEF in specific patient groups.

METHODS

STUDY DESIGN

As previously reported,¹¹ NEDA is a very large, ongoing observational registry that captures individual echocardiographic data on a retrospective and prospective basis from participating centres Australia-wide (<https://www.neda.net.au/participating-sites/>). During the second iteration of data collection, 23 centres contributed to the registry. Australia's public-private, health care system provides universal coverage to the entire population. Complete provision of all echocardiographs from participating sites is standard practice and includes all parameters generated for each investigation/case. Consequently, NEDA represents a

reliable and robust barometer of the clinical caseload and outcomes of patients being investigated (predominantly via General Practitioner and Cardiology referral) and managed for suspected or established heart disease derived from Australia's ethnically diverse population (~25 million people). NEDA is registered with the Australian New Zealand Clinical Trials Registry (ACTRN12617001387314). Approval has been obtained from all relevant Human Research Ethics Committees and the study adheres to the Declaration of Helsinki. Original study data are only available to investigators from contributing NEDA centres. However, sharing of data outputs can be provided by the Corresponding Author if requested.

ECHOCARDIOGRAPHY REPORTS

All echocardiographic measurement and report data, including basic demographic profiling (biological sex and date of birth) and date of investigation, of participating centres were collected and remotely transferred into a central database via a "vendor-agnostic" automated data extraction process – the study period being 1/1/2000 to 21/5/2019. Although not always available, body mass index (BMI), blood pressure (BP) and heart rate data were also collected. All data are then transformed into standard NEDA format. Applying the *NEDA Study Protocol*, precise definitions for each echocardiography variable are created and duplicate measurements/investigations combined. A continuously updated *NEDA Data Dictionary* is maintained via a *Master NEDA Database*. Specialised text recognition software was applied to free text/clinical comments/conclusions to enhance the detection of specific patient groups; including those referred with suspected or a pre-existing diagnosis of HF.

For this study, all men and women aged ≥ 18 years with a quantified LVEF (with ranges and text descriptors not accepted) were considered eligible. Consistent with current guidelines,² a hierarchal preference for Simpson's biplane-derived LVEF (27.6% of cases) over 2-D Teichholz (55.4% of cases) and other quantification methods was applied. All primary data analyses were case-based and focused on the first-recorded echocardiogram.

ENDPOINTS

To derive the primary outcomes of cardiovascular-related and all-cause mortality, data-linkage was performed via Australia's National Death Index.¹⁵ This validated resource provides an accurate list of primary and secondary diagnoses linked to each death according to ICD-10 coding. Via an exhaustive probability matching process, reliable data on the survival status of all individuals, up to the study census date of

21/5/2019, were generated. Study follow-up comprised a median of 2,027 (interquartile range [IQR] 1,134, 3,191) days and a combined total of >3 million person-years follow-up. Consistent with previous NEDA reports,¹²⁻¹⁴ those deaths linked to a ICD-10 chapter code of I00-I99 (primary diagnosis) were considered a cardiovascular-related death.

STATISTICAL METHODS

NEDA data analyses and reports conform to STROBE guidelines.¹⁶ For descriptive purposes, LVEF groups were generated per 10-unit increments (see *Table 1*) and for all distribution and survival analyses in 5-unit increments. Standard methods for describing/comparing grouped data, including means (\pm standard deviation), median (IQR) and proportions (with 95% confidence intervals [CI]) were performed. Incidence rates (with 95% CI) of cardiovascular-related and all-cause mortality were calculated as events per 1,000 person-years follow-up. Due to the non-linearity of the relationship between LVEF and mortality, restricted cubic spline analyses were undertaken; a common LVEF of 65.0-69.9% being established as the reference group for all mortality comparisons. Actual 1- and 5-year mortality were calculable in 456,644 and 272,375 cases, respectively. The Kaplan Meier method followed by Cox-Proportional Hazard Models (entry method with proportional hazards confirmed by visual inspection) were used to derive adjusted hazard ratios (HR) for mortality. For the main analyses of mortality according to 5-unit increments in LVEF, adjustments were made for age, year of echocardiogram (3-year epochs), BMI, heart rate, right heart function (TR peak velocity), parameters of diastolic function (LAVI, LVDD, E' velocity and E wave velocity), LVH and valvular heart disease (see *Table 1*). The size of models being determined by those with complete profiling data. Applying the same methods (with adjustment for age, year of investigation, LVH and valvular heart disease), the equivalent pattern of mortality according to 5-unit increments in LVEF were specifically examined in the following groups on a sex-specific basis – **i)** Aged above and below 65 years (all cases), **ii)** Referred for the investigation of potential HF (32,403 cases) or with a pre-existing diagnosis of HF (33,738 cases), and **iii)** Those presenting with severe aortic stenosis (6924 cases) or LVH (105,858 cases). All analyses were performed with SPSS v26.0 and statistical significance accepted at a 2-sided p-value of <0.05.

RESULTS

STUDY COHORT

Overall, 237,046 women and 256,109 men with a minimum of one quantified LVEF were studied ([Supplementary Figure S1](#)). Overall, mean age was 61 years and 48% were women. As shown in **Table 1**, there were distinctive trends in the distribution of women (increasing numbers) and average age of both sexes (decreasing age) as LVEF levels increased; with a reversal in age-related trends at the higher distribution of LVEF. Concurrent evidence of LVH (21.4% of all cases), valvular heart disease (12.3%) pulmonary hypertension (10.1%) were increasingly evident among those with moderate-to-severe LV systolic dysfunction. Overall, the sex-based distribution of LVEF was markedly different ([Supplementary Figure S2](#) and **Graphical Abstract**), with 8.3% versus 17.6% of women and men, respectively, with a LVEF <50%.

ALL-CAUSE & CARDIOVASCULAR-RELATED MORTALITY

INCIDENCE RATES

During 1.47 million and 1.55 million person-years follow-up, respectively, 52,705 women (22.2%) and 66,527 men (26.0%) died from all-causes (119,232 deaths overall). Of these, 16,827 (31.9%) and 20,762 (31.2%), respectively, were cardiovascular-related; with markedly higher rates of mortality in those with an LVEF <50% (peaking at ~70 deaths per 1,000 person-years) – see **Figure 1**. A similar pattern was observed when examining all-cause mortality ([Supplementary Figure S3](#)). In both sexes, cardiovascular-related mortality was lowest among those with a LVEF of 65.0-69.9%; an incident rate of ~8-10 deaths per 1,000 person-years being evident in the 60.0-75.0% LVEF range. Above these levels, mortality rates slightly increased, predominantly driven by non-cardiovascular deaths.

1- AND 5-YEAR ACTUAL MORTALITY

The overall pattern of 1- and 5-year actual cardiovascular-related and all-cause mortality according to LVEF levels was similar for women and men ([Supplementary Figure S4](#)). An LVEF <40% was associated with the poorest 1- and 5-year survival profiles; peaking at ~20-30% and ~40-50%, respectively, for cardiovascular-related and all-cause mortality at 5-years. There was a common nadir in cardiovascular-related (<5%) and all-cause mortality (~12-15%) at 5-years associated with a LVEF of 65.0-69.9% in women and men. Although a small absolute increase in 1- and 5-year mortality was evident at the upper distribution of LVEF levels, much of these excess mortality risks were attenuated on a fully adjusted basis – see [Supplementary Table S1](#).

ADJUSTED LONG-TERM MORTALITY

Figure 2 shows the fully adjusted risk of cardiovascular-related mortality above and below the reference LVEF level of 65.0-69.9% among the 56,715 women and 50,978 men with all available profiling data. Reflecting crude mortality rates, overall, the risks of cardiovascular-related mortality ($p < 0.001$ for all comparisons) were markedly elevated below a LVEF of 55.0%. Despite lower 5-year mortality rates (5.3% - 5.8%) overall, the adjusted risk of cardiovascular-related mortality associated with a LVEF of 55.0-59.9% was elevated in both women (HR 1.36, 95% CI 1.16, 1.59; $p < 0.001$) and men (HR 1.21, 95% CI 1.05, 1.39; $p = 0.008$). For women (HR 1.33, 95% CI 1.16, 1.52; $p < 0.001$) but not men (1.03, 95% CI 0.91, 1.18; $p = 0.620$) within the 60.0-64.9% LVEF group, this risk remained elevated (**Graphical Abstract**). Analyses of the contributory causes of death (any diagnostic position) according to LVEF levels showed that patterns of mortality largely reflect the competing risk posed by underlying cardio-renal-metabolic disease versus malignancy - [Supplementary Figure S5](#).

SUB-GROUP ANALYSES

Figure 3 shows the adjusted risk of cardiovascular-related mortality in those women and men aged above and below 65 years. In both sexes, the same pattern of mortality below a LVEF of 65.0-69.9% was evident in both groups. However, this pattern was more striking in the younger age group. Similarly, the threshold of significantly increased risk of mortality just below the identified reference group was 60.0-64.9% for women compared to 55.0-59.9% for men. **Figure 4** shows a similar pattern of mortality (but without a substantive level of mortality risk above the reference group) among the 18,456 women and 13,947 men referred for suspected HF. Alternatively, among the 15,710 women and 18,028 men with a pre-existing diagnosis of HF, the threshold of LVEF associated with a significantly increased risk of cardiovascular-related mortality was 50.0-54.9%. Similar patterns of mortality were specifically observed among severe AS and LVH cases; women in the latter group once again showing a higher threshold of mortality than men - [Supplementary Figure S6 and S7](#).

DISCUSSION

To our knowledge, this is the largest ever study of cardiovascular-related and all-cause mortality across the full spectrum of quantified LVEF observed in routine clinical practice. Moreover, unlike previous reports, outcomes are reported on a sex-specific basis. Specifically, routinely acquired echocardiographic data of almost 500,000 men and women were linked to 119,000 deaths during 3 million person-years follow-up. Within this large and heterogeneous cohort, women presented with a different pattern of LV systolic function compared to men. Although twice as many men presented with a LVEF <50%, below this threshold, cardiovascular-related mortality rates were similarly high in both sexes. In both women and men unadjusted mortality was lowest at a LVEF of 65.0-69.9%. In women but not men, an increased risk of cardiovascular-related mortality persisted to a LVEF threshold of 60.0-64.9%. In men, the equivalent threshold of increased mortality occurred at a lower LVEF (55.0-59.9%). This subtle but important sex-based difference persisted in nearly all sub-group analyses excepting those with a pre-existing diagnosis of HF. Overall, these data support recent efforts to determine if there are indeed important sex-based differences in therapeutic responses and outcomes in those with a LVEF >45% and evidence of HFpEF.^{6,7} More importantly, regardless of the specific reasons for our findings, they indicate that current applied thresholds for interpreting and acting upon routinely acquired LVEF levels may need to be revisited on a sex-specific basis.

Somewhat surprisingly, the distribution and prognostic implications of routinely observed LVEF (the most measured and utilised parameter for detecting and managing LV dysfunction) remains under-reported – especially in women. Until very recently, the most informative studies typically involved modestly sized cohorts with HF_rEF¹⁷ or clinical trials.¹⁸⁻²¹ As summarised in [Supplementary Table 2](#), recently published studies have somewhat addressed this evidence-gap.⁸⁻¹⁰ However, unlike NEDA, these studies largely rely on qualitative (visual) LVEF estimates rather than quantitative (such as Simpson's biplane method). Moreover, none provide specific data for women. Alternatively, these same studies do provide the more granular clinical profiling and outcome data that is not yet available to NEDA. Consistent with our main findings, in a large meta-analysis of mortality in 41,972 patients (35% women) with HF_rEF versus HF_pEF, the latter had much higher mortality rates overall. However, within this more select cohort, no discernible differences in mortality above a LVEF of 40% was evident.²¹ Alternatively, consistent with our report (particularly those relating to those with a pre-existing diagnosis of HF), a combined analysis of the PARADIGM-HF and PARAGON-HF cohorts (13,195 patients with HF_rEF to HF_pEF), demonstrated that the

risk of cardiovascular-related mortality extends well beyond currently accepted levels of normal LV systolic function.¹⁹ Recently, a report focussing on physician-derived LVEF examined the pattern of 46,258 deaths among 203,135 patients with 403,977 echocardiograms.⁸ Broadly consistent with our findings, this study demonstrated a nadir of all-cause mortality around an LVEF of 60-65% overall.⁸ A key finding of this study was an elevated risk of (all-cause mortality) associated with LVEF levels indicative of a hyper-dynamic LV. We observed a similar phenomenon, but found it was largely (but not exclusively) due to non-cardiovascular deaths and included relatively few cases. These specific findings are broadly consistent with the SCREEN-HF Study suggesting that higher LVEF levels are typically associated with advanced age, a smaller LV cavity and higher relative wall thickness; particularly in women.²² However, the contribution of non-cardiovascular mortality and a clear signal of increased mortality with higher LVEF levels in women aged <65 years requires further investigation. Regardless of the mechanisms, our findings reinforce those derived from more detailed analyses of LVEF derived from computed tomography,²³ and the EchoNoRMAL initiative,²⁴ in highlighting the need for more nuanced considerations of the prognostic significance of routinely observed levels of LVEF. In the common clinical setting of suspected, but not definitive diastolic dysfunction/HFpEF,¹⁴ our findings also support the need more definitive investigations (e.g. examining Global Longitudinal Strain²⁵) among those presenting with a LVEF of 50-60%. Early diastolic dysfunction, with normal filling pressures (and low E' velocities, E:A reversal and normal LAVi) may be seen across the spectrum of LVEF.²⁶ Progressively more abnormal diastolic function is associated with increased left ventricular filling pressures (and lower e' velocities, larger LAVi, and pseudonormalised or increased E:A ratio). Furthermore, in "normal" LVEF there is significant overlap between the Doppler diastolic indices of healthy individuals, normal ageing and diastolic dysfunction.

Despite the heterogeneity inherent to our study cohort, the clinical significance of our findings is reinforced by the outcomes of HFpEF trials. Firstly, in keeping with the CHARM Programme²⁷, TOPCAT⁷ and combined analyses of the PARADIGM-HF and PARAGON-HF Trials¹⁸, there is an evolving rationale to treat HF patients with an LVEF above 40% and below the "nadir" we identified at approximately 60%. Recent reports from a series of post-hoc analyses of the PARAGON-HF⁶ and TOPCAT⁷ trials, reinforced by a patient-level meta-analysis of mortality trials of neurohormonal modulating therapies,²⁸ appear to show that women derive treatment benefits at higher LVEF levels when compared to men. When combined with our "real-world" findings (with similar outcomes found in those presenting with suspected versus pre-existing HF), there is a

cogent rationale to systematically address the current gap in the evidence around sex-specific mechanisms of LV dysfunction and associated mortality, identifying optimal drug doses for women and applying sex-specific criteria for applying device-based therapies for HF.²⁹

LIMITATIONS

The NEDA cohort specifically reflects the broad characteristics and survival profile of those being investigated/managed for heart disease. As demonstrated by the specific pattern of mortality among women and men with a diagnosis of HF, as with our previous reports focussing on pulmonary hypertension¹² and aortic stenosis¹³, there is need to confirm our findings in specific patient populations. Excepting those with diagnoses derived from the National Death Registry of Australia, NEDA does not (yet) capture important clinical details on conditions such as coronary artery disease and other important determinants of health outcomes. We also do not have data on the ethnic profile of participants. NEDA also lacks clinical granularity in respect to individual patterns of treatment; our inclusion of the year of investigation at least reflecting broad changes in treatment over the study period. Unlike many previous reports, we specifically focussed on quantitative LVEF levels. However, reflecting real-world practice, many were derived from the non-recommended 2-D Teicholz method. To determine if the method of LVEF estimation confounded our findings, we undertook a sensitivity analysis ([Supplementary Figure S8](#)) that confirmed our overall findings. Study results were also highly consistent across all contributing NEDA centres. Finally, as highlighted by a recent expert consensus statement,³⁰ there are very little data on the clinical significance and prognostic implications of clinical variations in LVEF and this was not addressed in our current analyses. However, no substantive differences in the relationship between LVEF and mortality were noted when using the first or last recorded LVEF and a sensitivity analysis based on single versus multiple recorded LVEF levels also confirmed the consistency of our findings in this regard.

CONCLUSION

This analysis of a large cohort of patients routinely investigated with echocardiography, confirmed important sex-based differences in the distribution of LVEF and associated mortality (see **Graphical Abstract**). Within the range of LVEF associated with HFrEF, men were 2 to 3-fold more prevalent, but had broadly equivalent survival profiles to women. At the level of near equivalent sex-specific prevalence (a LVEF of 60.0-64.9%), women appeared to have a greater level of risk of cardiovascular-mortality compared to men. Overall, these

data reinforce the need for greater efforts to understand which women and men would benefit from more proactive clinical profiling and evidence-based treatments when presenting with a relatively preserved LVEF.

SUMMARY SENTENCE

During 3 million person-years follow-up of 237,046 women and 256,109 men subject to routine echocardiography, there were clinically important differences between men and women in the pattern of mortality according to the level of underlying left ventricular systolic dysfunction.

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REFERENCES

1. Celutkiene J, Spoletini I, Coats AJS, Chioncel O. Left ventricular function monitoring in heart failure. *Eur Heart J Suppl.* 2019;21:M17-M19.
2. Lang RM, Badano LP, Mor-Avi V, Afilalo J, Armstrong A, Ernande L, Flachskampf FA, Foster E, Goldstein SA, Kuznetsova T, Lancellotti P, Muraru D, Picard MH, Rietzschel ER, Rudski L, Spencer KT, Tsang W, Voigt JU. Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. *Eur Heart J Cardiovasc Imaging.* 2015;16:233-70.
3. Ponikowski P, Voors AA, Anker SD, Bueno H, Cleland JGF, Coats AJS, Falk V, Gonzalez-Juanatey JR, Harjola VP, Jankowska EA, Jessup M, Linde C, Nihoyannopoulos P, Parissis JT, Pieske B, Riley JP, Rosano GMC, Ruilope LM, Ruschitzka F, Rutten FH, van der Meer P. 2016 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure: The Task Force for the diagnosis and treatment of acute and chronic heart failure of the European Society of Cardiology (ESC) Developed with the special contribution of the Heart Failure Association (HFA) of the ESC. *Eur Heart J.* 2016;37:2129-2200.
4. Taylor CJ, Ordonez-Mena JM, Roalfe AK, Lay-Flurrie S, Jones NR, Marshall T, Hobbs FDR. Trends in survival after a diagnosis of heart failure in the United Kingdom 2000-2017: population based cohort study. *BMJ.* 2019;364:l223.
5. Stewart S, Ekman I, Ekman T, Oden A, Rosengren A. Population impact of heart failure and the most common forms of cancer: a study of 1 162 309 hospital cases in Sweden (1988 to 2004). *Circ Cardiovasc Qual Outcomes.* 2010;3:573-80.
6. McMurray JJV, Jackson AM, Lam CSP, Redfield MM, Anand IS, Ge J, Lefkowitz MP, Maggioni AP, Martinez F, Packer M, Pfeffer MA, Pieske B, Rizkala AR, Sabarwal SV, Shah AM, Shah SJ, Shi VC, van Veldhuisen DJ, Zannad F, Zile MR, Cikes M, Goncalvesova E, Katova T, Kosztin A, Lelonek M, Sweitzer N, Vardeny O, Claggett B, Jhund PS, Solomon SD. Effects of Sacubitril-Valsartan Versus Valsartan in Women Compared With Men With Heart Failure and Preserved Ejection Fraction: Insights From PARAGON-HF. *Circulation.* 2020;141:338-351.

7. Merrill M, Sweitzer NK, Lindenfeld J, Kao DP. Sex Differences in Outcomes and Responses to Spironolactone in Heart Failure With Preserved Ejection Fraction: A Secondary Analysis of TOPCAT Trial. *JACC Heart Fail.* 2019;7:228-238.
8. Wehner GJ, Jing L, Haggerty CM, Suever JD, Leader JB, Hartzel DN, Kirchner HL, Manus JNA, James N, Ayar Z, Gladding P, Good CW, Cleland JGF, Fornwalt BK. Routinely reported ejection fraction and mortality in clinical practice: where does the nadir of risk lie? *Eur Heart J.* 2020;41(12):1249-1257. .
9. Liang HY, Lo YC, Chiang HY, Chen MF, Kuo CC. Validation and Comparison of the 2003 and 2016 Diastolic Functional Assessments for Cardiovascular Mortality in a Large Single-Center Cohort. *J Am Soc Echocardiogr.* 2020;33:469-480.
10. Angaran P, Dorian P, Ha ACT, Thavendiranathan P, Tsang W, Leong-Poi H, Woo A, Dias B, Wang X, Austin PC, Lee DS. Association of Left Ventricular Ejection Fraction with Mortality and Hospitalizations. *J Am Soc Echocardiogr.* 2020;33:802-811 e6.
11. Strange G, Celermajer DS, Marwick T, Prior D, Ilton M, Codde J, Scalia GM, Stewart S, Bulsara M, Gabbay E, Playford D. The National Echocardiography Database Australia (NEDA): Rationale and methodology. *Am Heart J.* 2018;204:186-189.
12. Strange G, Stewart S, Celermajer DS, Prior D, Scalia GM, Marwick TH, Gabbay E, Ilton M, Joseph M, Codde J, Playford D. Threshold of Pulmonary Hypertension Associated With Increased Mortality. *J Am Coll Cardiol.* 2019;73:2660-2672.
13. Strange G, Stewart S, Celermajer D, Prior D, Scalia GM, Marwick T, Ilton M, Joseph M, Codde J, Playford D. Poor Long-Term Survival in Patients With Moderate Aortic Stenosis. *J Am Coll Cardiol.* 2019;74:1851-1863.
14. Playford DSG, Celermajer D, Evans G, Scalia GM, Stewart S, Prior D. Diastolic Dysfunction and Mortality in 436,360 men and women: The National Echo Database Australia (NEDA). *European heart Journal - Cardiovascular Imaging.* In press - doi:10.1093/ehjci/jeaa253.
15. Magliano D, Liew D, Pater H, Kirby A, Hunt D, Simes J, Sundararajan V, Tonkin A. Accuracy of the Australian National Death Index: comparison with adjudicated fatal outcomes among Australian participants in the Long-term Intervention with Pravastatin in Ischaemic Disease (LIPID) study. *Aust N Z J Public Health.* 2003;27:649-53.

16. Sharp MK, Tokalic R, Gomez G, Wager E, Altman DG, Hren D. A cross-sectional bibliometric study showed suboptimal journal endorsement rates of STROBE and its extensions. *J Clin Epidemiol.* 2019;107:42-50.
17. Solomon SD, McMurray JJV, Anand IS, Ge J, Lam CSP, Maggioni AP, Martinez F, Packer M, Pfeffer MA, Pieske B, Redfield MM, Rouleau JL, van Veldhuisen DJ, Zannad F, Zile MR, Desai AS, Claggett B, Jhund PS, Boytsov SA, Comin-Colet J, Cleland J, Dungen HD, Goncalvesova E, Katova T, Kerr Saraiva JF, Lelonek M, Merkely B, Senni M, Shah SJ, Zhou J, Rizkala AR, Gong J, Shi VC, Lefkowitz MP. Angiotensin-Nepriylsin Inhibition in Heart Failure with Preserved Ejection Fraction. *N Engl J Med.* 2019;381:1609-1620.
18. Solomon SD, Vaduganathan M, B LC, Packer M, Zile M, Swedberg K, Rouleau J, M AP, Desai A, L HL, Kober L, Anand I, Sweitzer N, Linssen G, Merkely B, Luis Arango J, Vinereanu D, Chen CH, Senni M, Sibulo A, Boytsov S, Shi V, Rizkala A, Lefkowitz M, McMurray JJV. Sacubitril/Valsartan Across the Spectrum of Ejection Fraction in Heart Failure. *Circulation.* 2020;141:352-361.
19. Solomon SD, Claggett B, Desai AS, Packer M, Zile M, Swedberg K, Rouleau JL, Shi VC, Starling RC, Kozan O, Dukat A, Lefkowitz MP, McMurray JJ. Influence of Ejection Fraction on Outcomes and Efficacy of Sacubitril/Valsartan (LCZ696) in Heart Failure with Reduced Ejection Fraction: The Prospective Comparison of ARNI with ACEI to Determine Impact on Global Mortality and Morbidity in Heart Failure (PARADIGM-HF) Trial. *Circ Heart Fail.* 2016;9:e002744.
20. Solomon SD, Claggett B, Lewis EF, Desai A, Anand I, Sweitzer NK, O'Meara E, Shah SJ, McKinlay S, Fleg JL, Sopko G, Pitt B, Pfeffer MA. Influence of ejection fraction on outcomes and efficacy of spironolactone in patients with heart failure with preserved ejection fraction. *Eur Heart J.* 2016;37:455-62.
21. Meta-analysis Global Group in Chronic Heart Failure. The survival of patients with heart failure with preserved or reduced left ventricular ejection fraction: an individual patient data meta-analysis. *Eur Heart J.* 2012;33:1750-7.
22. McGrady M, Reid CM, Shiel L, Wolfe R, Boffa U, Liew D, Campbell DJ, Prior D, Stewart S, Krum H. NT-proB natriuretic peptide, risk factors and asymptomatic left ventricular dysfunction: results of the SCReening Evaluation of the Evolution of New Heart Failure study (SCREEN-HF). *Int J Cardiol.* 2013;169:133-8.

23. Lin FY, Devereux RB, Roman MJ, Meng J, Jow VM, Jacobs A, Weinsaft JW, Shaw LJ, Berman DS, Callister TQ, Min JK. Cardiac chamber volumes, function, and mass as determined by 64-multidetector row computed tomography: mean values among healthy adults free of hypertension and obesity. *JACC Cardiovasc Imaging*. 2008;1:782-6.
24. Echocardiographic Normal Ranges Meta-Analysis of the Left Heart Collaborators. Ethnic-Specific Normative Reference Values for Echocardiographic LA and LV Size, LV Mass, and Systolic Function: The EchoNoRMAL Study. *JACC Cardiovasc Imaging*. 2015;8:656-65.
25. Cho GY, Marwick TH, Kim HS, Kim MK, Hong KS, Oh DJ. Global 2-dimensional strain as a new prognosticator in patients with heart failure. *J Am Coll Cardiol*. 2009;54:618-24.
26. Nagueh SF, Smiseth OA, Appleton CP, Byrd BF, 3rd, Dokainish H, Edvardsen T, Flachskampf FA, Gillebert TC, Klein AL, Lancellotti P, Marino P, Oh JK, Popescu BA, Waggoner AD. Recommendations for the Evaluation of Left Ventricular Diastolic Function by Echocardiography: An Update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging. *Journal Of The American Society Of Echocardiography: Official Publication Of The American Society Of Echocardiography*. 2016;29:277-314.
27. Pfeffer MA, Swedberg K, Granger CB, Held P, McMurray JJ, Michelson EL, Olofsson B, Ostergren J, Yusuf S, Pocock S. Effects of candesartan on mortality and morbidity in patients with chronic heart failure: the CHARM-Overall programme. *Lancet*. 2003;362:759-66.
28. Dewan P, Jackson A, Lam CSP, Pfeffer MA, Zannad F, Pitt B, Solomon SD, McMurray JJV. Interactions between left ventricular ejection fraction, sex and effect of neurohumoral modulators in heart failure. *Eur J Heart Fail*. 2020. 22. 10.1002/ejhf.1776.
29. Lam CSP, Arnott C, Beale AL, Chandramouli C, Hilfiker-Kleiner D, Kaye DM, Ky B, Santema BT, Sliwa K and Voors AA. Sex differences in heart failure. *Eur Heart J*. 2019;40:3859-3868c.
30. Wilcox JE, Fang JC, Margulies KB and Mann DL. Heart Failure With Recovered Left Ventricular Ejection Fraction: JACC Scientific Expert Panel. *J Am Coll Cardiol*. 2020;76:719-734.

TABLE 1. BASELINE PROFILE

LVEF Groups	ALL (N=493,155)	<30% (n=14,758)	30 – 39% (n=17,686)	40 – 49% (n=32,290)	50 – 59% (n=101,354)	60 – 69% (n=203,369)	≥70% (n=123,696)
Demographic Profile							
Age, years	60.8±17.7	66.4±15.5	68.0±14.9	65.5±16.3	59.8±18.4	58.7±18.0	62.2±16.9
Female (%)	237,046 (48.1%)	3826 (25.9%)	5304 (30.0%)	10,484 (32.5%)	41,054 (40.5%)	103,999 (51.5%)	72,379 (58.5%)
Anthropometrics							
BMI, m/kg ²	28.1±6.3	27.7±6.5	28.1±6.4	28.2±6.3	28.2±6.4	27.9±6.2	28.3±6.4
Vital Signs							
Heart rate, bpm	72.1±15.5	85.1±21.8	79.0±19.5	75.8±18.4	71.4±15.2	69.6±13.9	74.0±15.0
Systolic BP, mmHg	135±22	120±23	126±22	131±23	131±22	133±22	140±22
Diastolic BP, mmHg	77±11	72±13	73±13	75±13	75±12	76±12	80±10
Right Heart Function							
eRVSP, mmHg	32.6±10.8	38.9±12.3	37.1±12.3	34.7±11.6	31.6±10.7	31.0±9.9	33.7±10.8
TR peak velocity, m/s	2.6±0.5	2.9±0.5	2.8±0.5	2.7±0.5	2.5±0.5	2.5±0.5	2.6±0.5
Left Heart Dimensions & Function							
LVDD, cm	4.8±0.7	5.9±1.0	5.4±0.9	5.1±0.8	4.8±0.6	4.6±0.6	4.6±0.6
LVSD, cm	3.1±0.8	5.1±1.0	4.4±0.8	3.8±0.7	3.3±0.6	2.9±0.5	2.8±0.5
LVEF, %	61.7±12.8	22.0±5.5	34.8±13.0	44.9±3.0	55.6±2.8	64.3±2.9	75.5±4.9
Mitral E' velocity, cm/s	8.3±3.0	5.3±2.2	5.8±2.2	6.5±2.4	8.0±2.9	8.7±2.9	8.6±3.1
Mitral E wave velocity, cm/s	80.91±26.4	87.9±30.4	84.9±32.3	81.1±31.2	78.1±27.3	80.0±24.6	83.4±25.5
Mitral A wave velocity, cm/s	71.8±27.8	64.3±30.2	72.1±28.6	73.8±27.7	69.1±26.8	68.9±26.6	79.1±28.9
Mitral E:A ratio	1.2±0.7	1.7±1.5	1.3±1.0	1.2±0.8	1.2±0.7	1.2±0.6	1.1±0.6
SVi, ml/m ²	40.1±12.8	28.1±14.0	34.2±10.8	37.3±12.2	40.1±10.6	42.1±12.5	43.9±13.6
LVH (%)	105,858 (38.1%)	5331 (81.5%)	5789 (74.1%)	8798 (60.0%)	19,357 (35.4%)	33,619 (27.4%)	32,964 (46.2%)
LAVi, ml/m ²	40.5±27.3	61.7±38.3	55.2±35.6	47.8±30.9	34.6±19.6	32.5±18.6	52.1±33.8
Evidence of Left Heart Disease							
Pulmonary HT (%)	49,118 (18.4%)	4045 (41.6%)	3692 (35.0%)	4683 (26.4%)	8569 (16.3%)	14,387 (13.3%)	13,732 (20.0%)
Septal E:E' ratio >12.0 (%)	53,282 (10.8%)	2874 (19.6%)	3010 (17.0%)	4353 (14.0%)	9605 (9.5%)	18,138 (8.9%)	15,120 (12.2%)
LAVi >34 ml/m ² (%)	75,334 (15.3%)	2640 (17.9%)	2884 (16.3%)	4670 (14.5%)	11,541 (11.4%)	22,360 (11.0%)	31,239 (25.3%)
Valvular heart disease (%)	60,382 (15.7%)	5194 (40.5%)	4922 (32.7%)	6622 (25.0%)	12,402 (15.4%)	18,328 (11.5%)	12,914 (14.2%)
Outcomes							
Follow-up, months	89.0±49.4	95.8±49.4	94.3±49.0	89.4±47.8	84.2±46.9	86.3±46.1	95.8±48.3
All-cause deaths (%)	119,232 (24.2%)	7735 (52.4%)	8426 (47.6%)	12,114 (37.5%)	23,987 (23.7%)	37,851 (18.6%)	29,119 (23.5%)
CVD cause of death (%)/ % of all deaths	37,499 (7.6%) / 31.5%	4069 (21.5%) / 52.6%	3802 (21.5%) / 45.1%	4656 (14.4%) / 38.4%	7131 (7.0%) / 29.7%	9899 (4.9%) / 26.2%	7942 (6.4%) / 27.3

Legend: Body mass index (BMI) recorded in 343,868 cases, Heart rate in beats/min (bpm) – 208,138 cases, Systolic & Diastolic Blood Pressure (BP) – 60,522 and 60,265 cases, estimated right ventricular systolic Pressure (eRVSP) – 267,496 cases, tricuspid regurgitation (TR) peak velocity – 277,247 cases, left ventricular diastolic/systolic diameter (LVDD/LVSD) – 398,925 and 371,285 cases, mitral annular E' velocity – 227,650 cases, mitral E wave velocity – 406,978 cases, mitral A wave velocity – 374,757 cases, mitral E:A ratio – 379,054 cases , stroke volume index (SVi) – 128,569 cases, left ventricular hypertrophy (LVH) status quantified in 277,752 cases (applying a modified American Society of Echocardiography criteria), left atrial volume index (LAVi) – 179,722 cases, and presence of valvular heart disease determined in 385,483 cases according to the following criteria – a) presence of - moderate or greater aortic stenosis (mean aortic pressure gradient $>20\text{mmHg}$ or $\text{AVA} < 1.2\text{cm}^2$); b) moderate or greater aortic regurgitation (physician reported); c) moderate or greater mitral regurgitation (physician reported) and/or; d) mild or greater mitral stenosis (mean transmitral gradient $>5\text{mHg}$).

FIGURE LEGENDS

FIGURE 1. INCIDENT RATE OF CARDIOVASCULAR-RELATED MORTALITY

Legend: The rate of cardiovascular-related mortality per 1,000 person-years, is presented separately for women (top graph) and men (bottom) according to 5-unit increments in LVEF. The total number of deaths contributing to the rate of mortality in each group (red numerals) are provided above the horizontal axis.

FIGURE 2. ADJUSTED CARDIOVASCULAR-RELATED MORTALITY

Legend: The box inserts show the adjusted HR (95% CI) of those co-variables included in the fully adjusted models that were significantly associated with mortality. Plots show the adjusted HR (plus or minus 95% CI) for cardiovascular-related mortality for each 5-unit LVEF group relative to the reference group.

FIGURE 3. AGE-SPECIFIC CARDIOVASCULAR-RELATED MORTALITY

Legend: Data for those men and women aged <65 years (top graphs) and 65 years and above (bottom graphs) are presented separately. The box inserts show the adjusted HR (95% CI) for all co-variables included in the adjusted models. Plots show the adjusted HR (plus or minus 95% CI) for cardiovascular-related mortality for each 5-unit LVEF group relative to the reference group – the same overall pattern of adjusted risk were replicated when fully adjusted models (with fewer cases) were applied. *** $p < 0.001$, ** $p < 0.01$. * $p < 0.05$.

FIGURE 4. CARDIOVASCULAR-RELATED MORTALITY – SUSPECTED HF CASES

Legend: The box inserts show the adjusted HR (95% CI) for all co-variables included in the adjusted models. Plots show the adjusted HR (plus or minus 95% CI) for cardiovascular-related mortality for each 5-unit LVEF group relative to the reference group – the same overall pattern of adjusted risk were replicated when fully adjusted models (with fewer cases) were applied. *** $p < 0.001$, ** $p < 0.01$. * $p < 0.05$.

FIGURE 5. CARDIOVASCULAR-RELATED MORTALITY – PRE-EXISTING HF CASES

Legend: The box inserts show the adjusted HR (95% CI) for all co-variables included in the adjusted models. Plots show the adjusted HR (plus or minus 95% CI) for cardiovascular-related mortality for each 5-unit LVEF group relative to the reference group – the same overall pattern of adjusted risk were replicated when fully adjusted models (with fewer cases) were applied. * $p < 0.05$.

FIGURE – GRAPHICAL ABSTRACT

Legend: The observed associated risk of between LVEF (on a continuous/unit-level basis) and probability of all-cause mortality is presented as smoothed spline curves (age-adjusted) for women and men separately. Shaded areas represent the 95% CI. Box inserts show the fully adjusted risk (HR plus 95% CI) of cardiovascular (CV)-related mortality with a LVEF 65.0-69.9% the reference group.

Figure 1

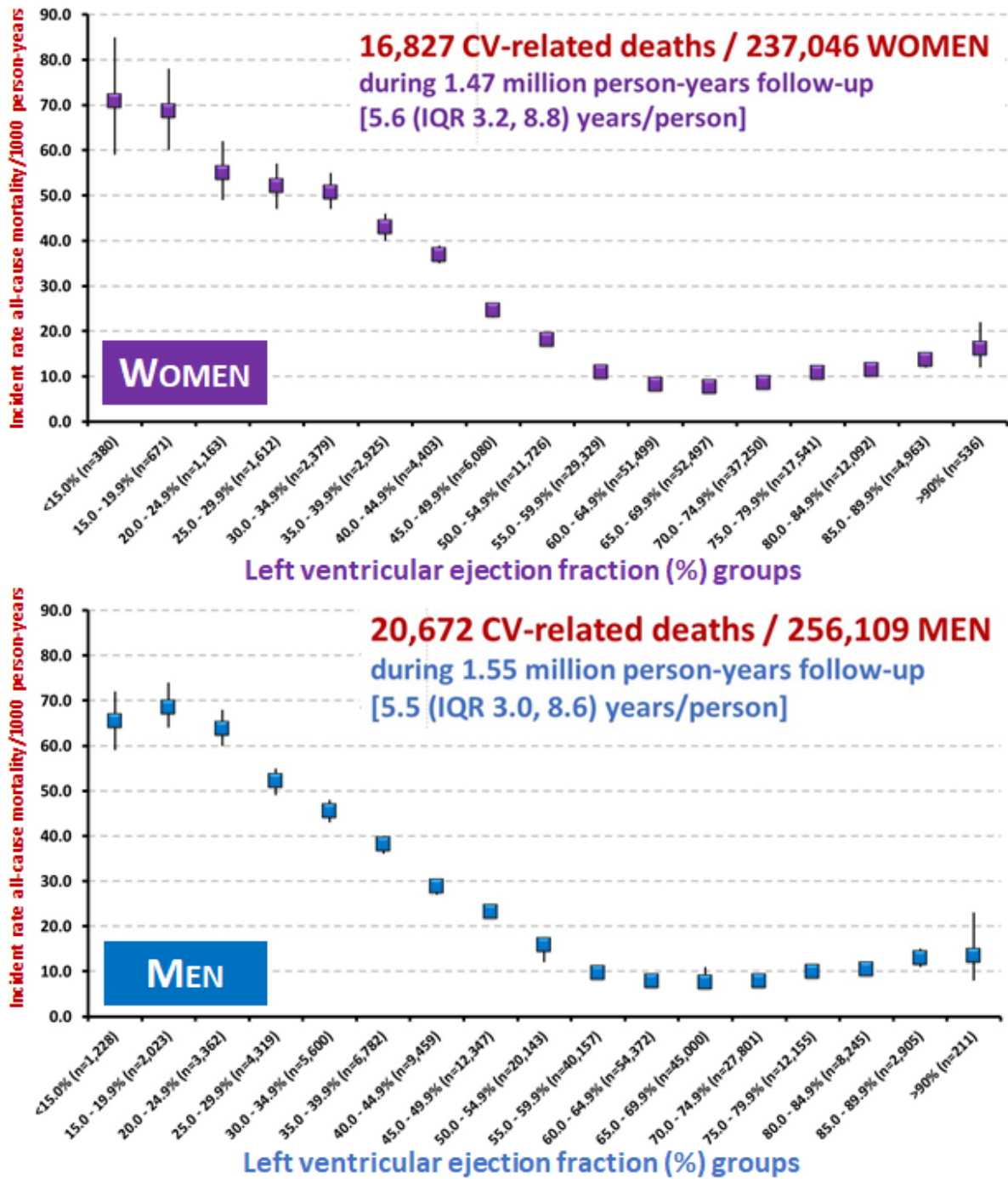


Figure 2

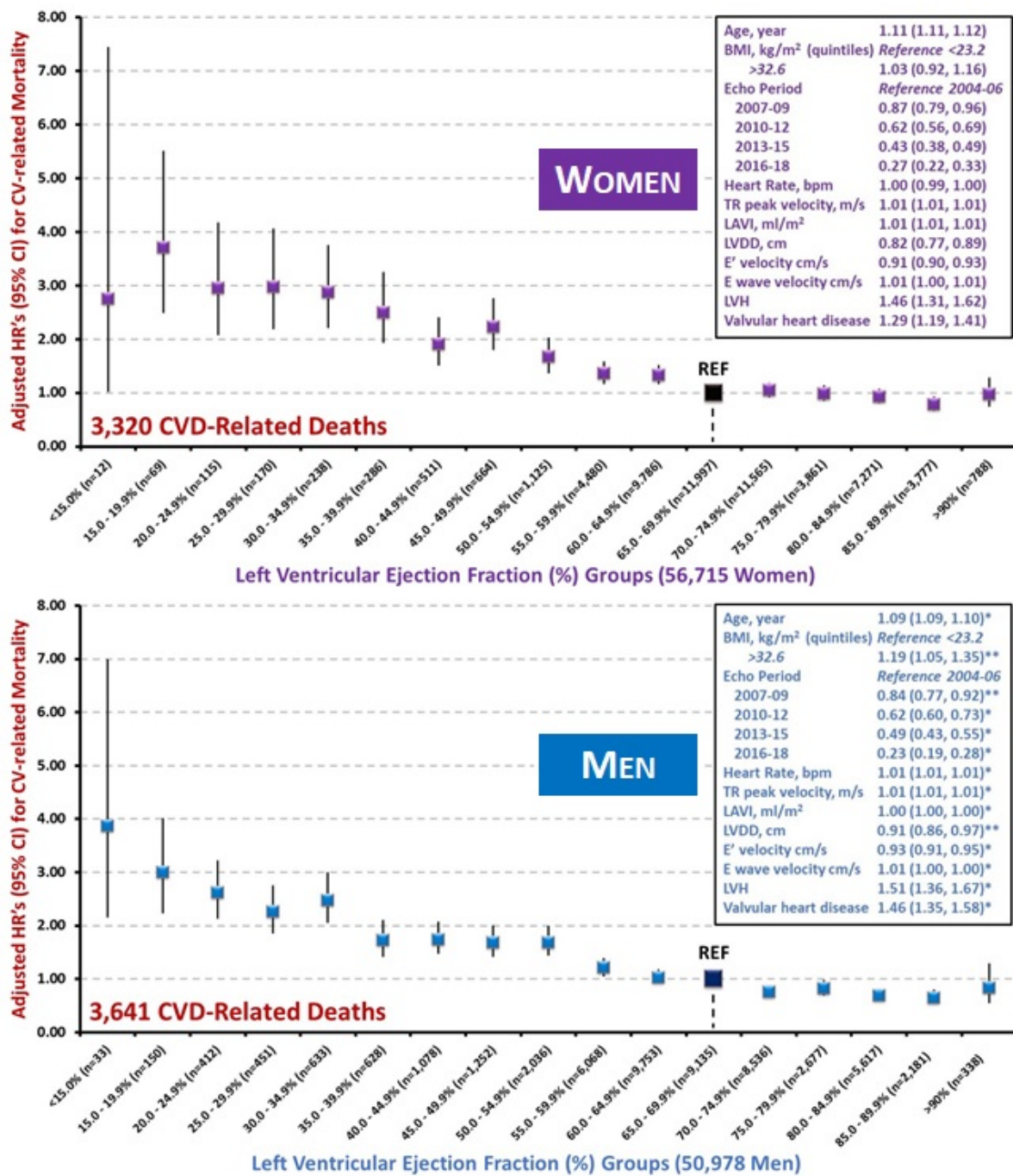


Figure 3

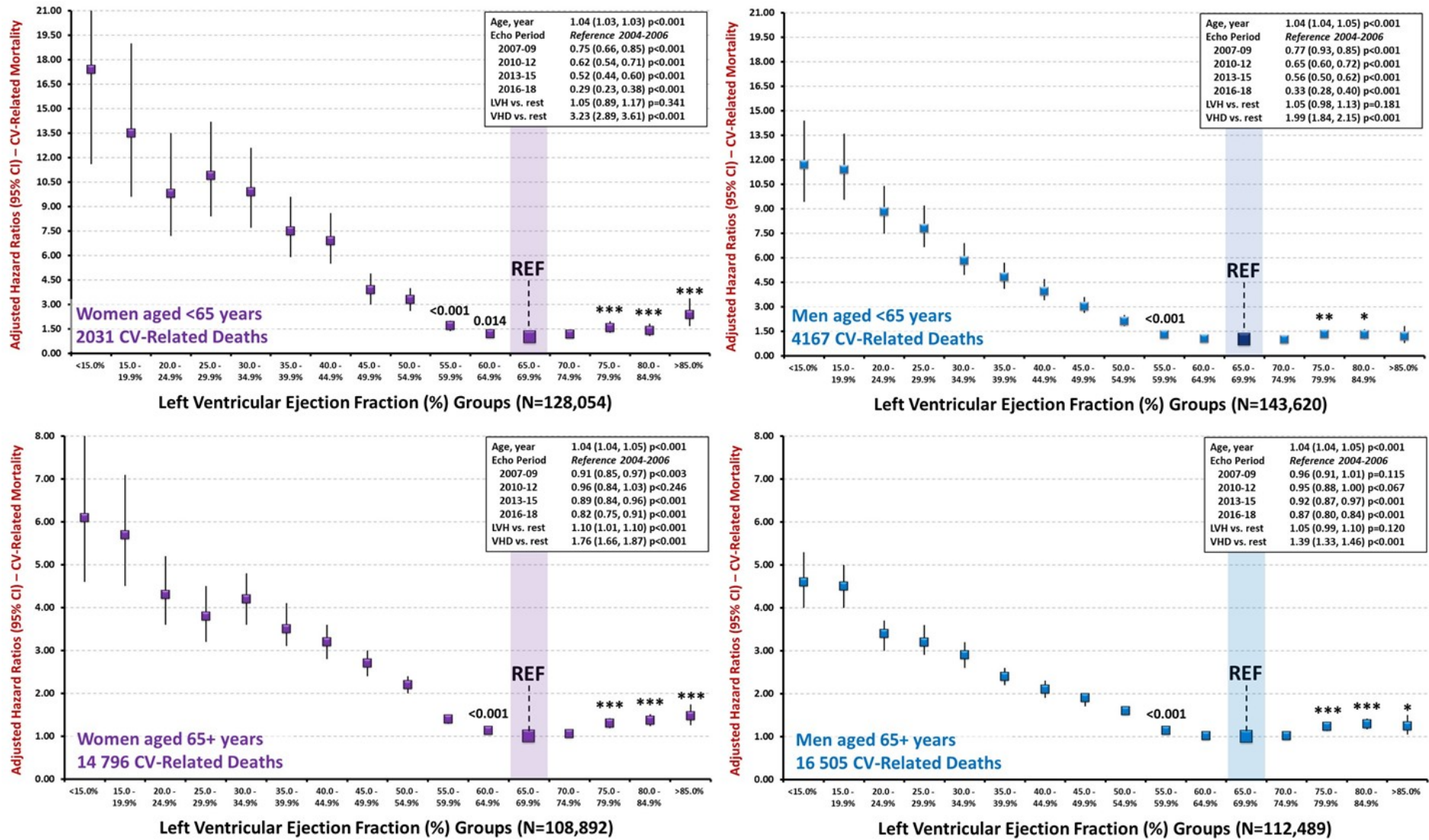


Figure 4

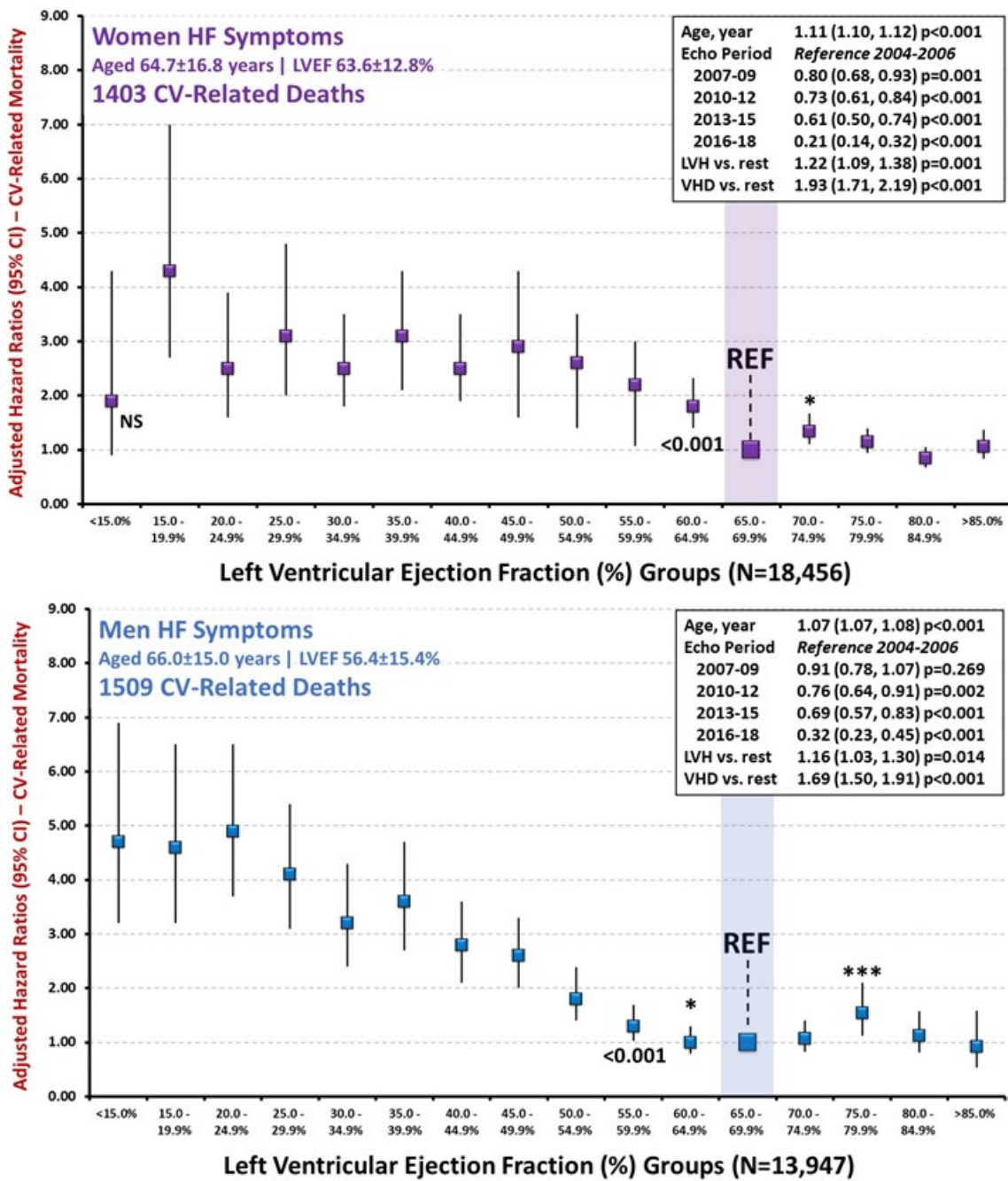


Figure 5

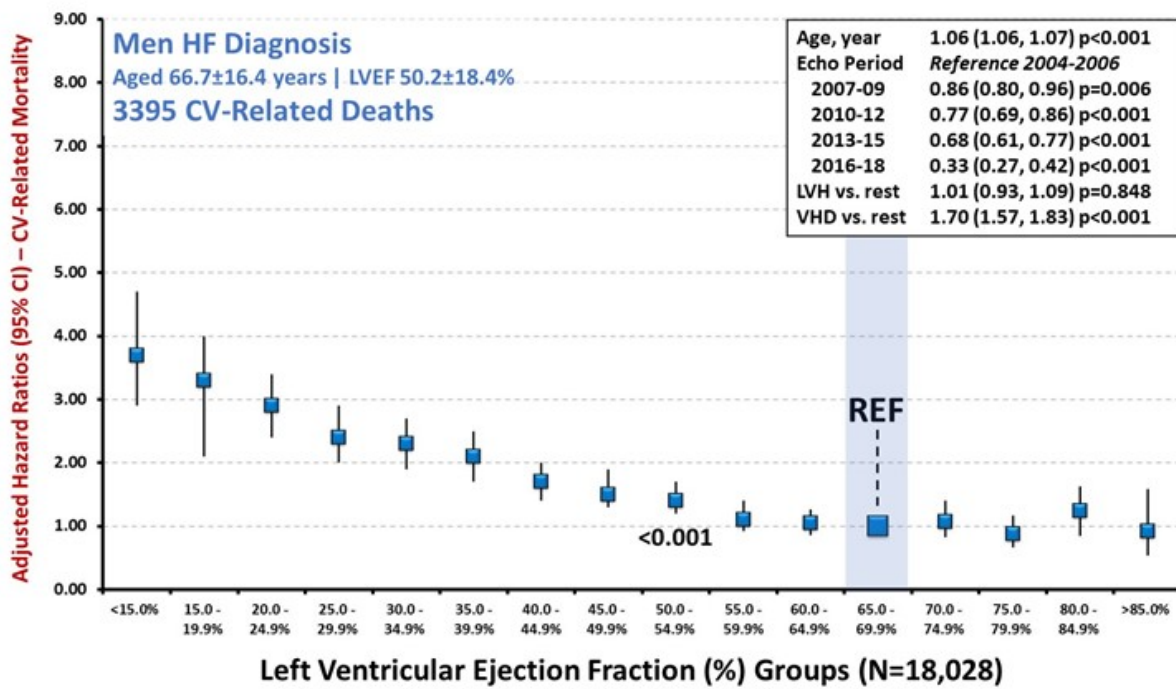
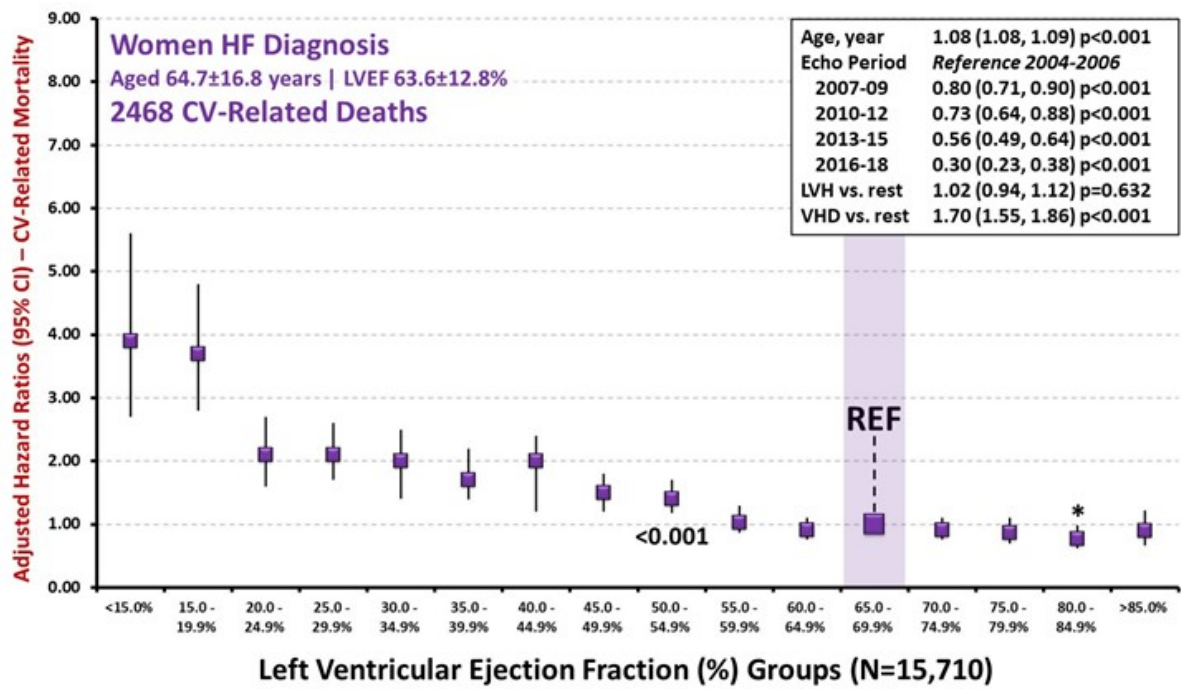
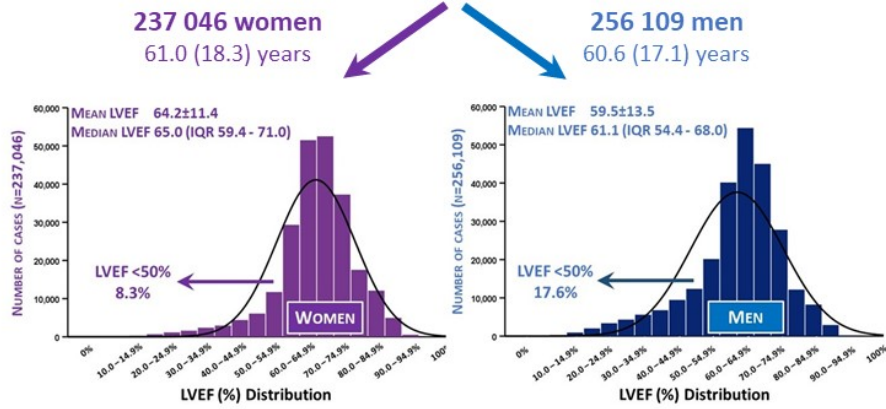


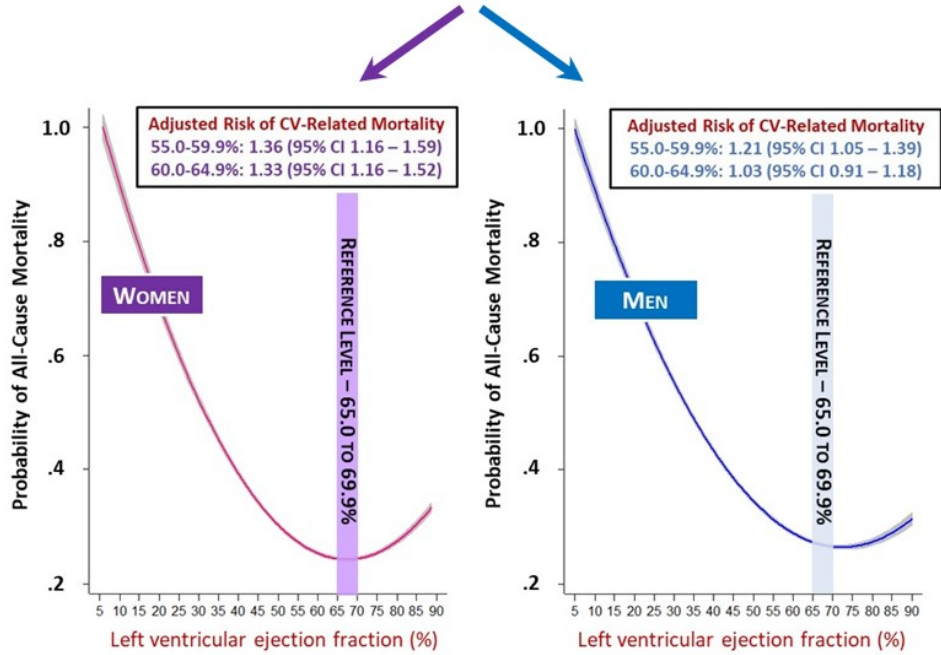
Figure – Graphical Abstract

Critical lack of large-scale, real world data on the sex-specific distribution & prognostic impact of LVEF

National Echocardiography Database of Australia
 493 155 cases investigated for suspected or established cardiovascular disease with echocardiography linked to mortality outcomes



Sex-specific differences in the distribution of LVEF



Sex-specific differences in CV-related mortality at equivalent LVEF