**Left atrial structure and function are associated with cardiovascular outcomes independent of left ventricular measures: a UK Biobank CMR study**

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**Abstract**

**Aims:** We evaluated associations of left atrial (LA) structure and function with prevalent and incident cardiovascular disease (CVD), independent of left ventricular (LV) metrics, in 25,896 UK Biobank participants.

**Methods and results:** We estimated association of cardiovascular magnetic resonance (CMR) metrics [LAV (LA maximum volume), LAEF (LA ejection fraction), LVM:LVEDV (LV mass: LV end-diastolic volume ratio), GLS (global longitudinal strain), LVFGI (LV global function index)] with vascular risk factors (hypertension, diabetes, high cholesterol, smoking), prevalent and incident CVDs [atrial fibrillation (AF), stroke, ischaemic heart disease (IHD), myocardial infarction (MI)], all-cause mortality, and CVD mortality. We created uncorrelated CMR variables using orthogonal principal component analysis rotation. All five CMR metrics were simultaneously entered into multivariable regression models adjusted for sex, age, ethnicity, deprivation, education, body size, and physical activity. Lower LAEF was associated with diabetes, smoking, and all the prevalent and incident CVDs. Diabetes, smoking, and high cholesterol were associated with smaller LAV. Hypertension, IHD, AF (incident and prevalent), incident stroke, and CVD mortality were associated with larger LAV. LV and LA metrics were both independently informative in associations with prevalent disease, however LAEF showed the most consistent associations with incident CVDs. Lower LVGFI was associated with greater all-cause and CVD mortality. In secondary analyses, compared to LVGFI, LV ejection fraction showed similar but less consistent associations.

**Conclusion:** LA structure and function measures (LAEF, LAV) demonstrate significant associations with key prevalent and incident cardiovascular outcomes, independent of LV metrics. These measures have potential clinical utility for disease discrimination and outcome prediction.

**Keywords:** left atrium, left ventricle, cardiovascular magnetic resonance, vascular risk factors, atrial fibrillation, stroke, ischaemic heart disease, cardiovascular outcomes, mortality.

**Introduction**

The left atrium (LA) is highly sensitive to subtle left ventricular (LV) haemodynamic changes.1,2 Alterations in LA structure and function may precede detectable LV dysfunction and, as such, have potential utility for earlier and more accurate disease discrimination than LV metrics.1–3 In particular, LA size and function are altered in response to elevated LV filling pressures, an early feature of diastolic dysfunction and a key component of heart failure with preserved ejection fraction (HFpEF).1,3 Furthermore, clinically important arrhythmias, such as atrial fibrillation (AF), primarily result in atrial (rather than ventricular) remodelling. Thus, atrial metrics may provide better indicators for the presence and occurrence of these conditions and provide incremental predictive value for key related health outcomes, such as stroke.4

The association of echocardiography derived measures of LA structure and function with incident and prevalent cardiovascular diseases (CVDs) has been repeatedly demonstrated.5–9 However, whilst the incremental value of LA over LV metrics seems biologically plausible, formal demonstration of this requires further study. Furthermore, although echocardiography is a valuable first line modality in clinical settings, cardiovascular magnetic resonance (CMR) is the reference standard for cardiac chamber quantification providing highly reproducible metrics calculated with fewer geometric assumptions than in echocardiography. Existing CMR studies of the utility of LA metrics are mostly based on small select samples of clinical cohorts,10–12 with a paucity of data from larger population-based samples.

The UK Biobank is a very large population-based cohort study including detailed participant characterisation, linked longitudinally tracked health outcome data, and detailed standardised CMR. Thus, we evaluated, in 25,896 UK Biobank participants, clinical associations of LA structure and function independent of LV metrics. We estimated associations of CMR derived LA and LV metrics with vascular risk factors (VRFs), prevalent CVD, incident CVD, and mortality outcomes. We considered a wide range of demographic and clinical confounders and, critically, we assessed the independent value of LA metrics over measures of LV structure and function.

**Methods**

**Setting and study participants**

The UK Biobank includes over 500,000 participants from across the UK. Individuals aged 40-69 years-old were identified using National Health Service (NHS) registers and recruited between 2006-2010 through postal invitations.13 Baseline assessment comprised detailed characterisation of participant demographic, lifestyle, environmental, and medical factors, as well as a series of physical measures and blood sampling. Individuals who could not complete baseline assessment due to discomfort or ill health were not recruited. The UK Biobank protocol is publicly available.14 Linkages have been established with key routine health data including hospital episode statistics (HES) and death registers, with health outcomes documented according to standardised International Classification of Diseases (ICD) codes. This linked information is continually updated allowing reliable longitudinal tracking of incident events for all participants. Furthermore, the UK Biobank has produced adjudicated algorithmically defined incident health outcome data for key illnesses, such as myocardial infarction (MI) and stroke.15 The UK Biobank imaging study, launched in 2015, aims to scan a random 100,000 subset of the original participants and includes, amongst other things, detailed CMR imaging.16

**CMR image acquisition**

The UK Biobank imaging study is performed using standardised pre-defined operating procedures, equipment, and staff training. CMR imaging was with 1.5 Tesla scanners (MAGNETOM Aera, Syngo Platform VD13A, Siemens Healthcare, Erlangen, Germany), the acquisition protocol is published elsewhere.17 In brief, cardiac function assessment comprised three long axis cines and a complete short axis stack covering the left and right ventricles acquired at one slice per breath hold using balanced steady-state free precession sequences.

**CMR image analysis**

CMR indices were derived using a fully automated quality-controlled image analysis pipeline previously developed and validated in a large subset of the UK Biobank.18,19 CMR metrics were available for the first 26, 891 UK Biobank CMR studies, of these both LA and LV data was available for 25,896 participants, which we include in the study (**Supplementary Figure 1**). We considered the following CMR measures: LAV (LA maximum volume), LAEF (LA ejection fraction, calculated as: LA maximum volume – LA minimum volume/ LA maximum volume), LVM: LVEDV (LV mass: LV end-diastolic volume ratio), GLS (global longitudinal strain). We considered LV global function index (LVGFI) as an additional measure of LV function. Previous reports have identified LVGFI as a strong predictor of heart failure and CVD events with incremental utility over LV ejection fraction.20,21 As per previous descriptions20,21, we defined LVGFI (%) as LV stroke volume/LV global volume\*100 where LV global volume was calculated as the sum of the LV mean cavity volume ((LV end-diastolic volume + LV end-systolic volume)/2) and myocardium volume (LV mass/density). Density of LV was specified as 1.05 g/mL. A higher LVGFI reflects better LV function. As LV ejection fraction (LVEF) is a more clinically established metric, we also considered associations with LVEF and compared its performance to LVGFI.

**Defining participant characteristics**

Sex, and ethnicity were taken as self-reported at baseline. Ethnicity was converted into a binary variable of White and BAME (Black Asian and minority ethnic) groups. Socioeconomic deprivation was recorded at the baseline UK Biobank assessment as the Townsend index, a measure of deprivation relative to national averages.22 Age was calculated at the time of imaging. Body mass index (BMI) was calculated from height and weight recorded at imaging. Educational level and smoking status were taken from self-report. Physical activity level was expressed as a continuous value of metabolic equivalent (MET) minutes/week, calculated by weighting different types of activity (walking, moderate, or vigorous) by its energy requirements using values derived from the International Physical Activity Questionnaire (IPAQ) study.23

**Ascertainment of vascular risk factors, cardiovascular disease, and mortality outcomes**

We considered the following VRFs: hypertension, diabetes, high cholesterol, and smoking, and the following CVDs (incident and prevalent): atrial fibrillation (AF), stroke, ischaemic heart disease (IHD), and MI. Mortality outcomes were ascertained from death register data. We considered all-cause and CVD mortality; the latter was defined as primary cause of death recorded as any CVD (ICD10 Chapter IX I00-I99). Incident CVDs and mortality outcomes were considered as those occurring after CMR imaging. The average follow-up time available for HES and mortality data was 4.2 ±1.2 (range: 2.5-6.9) years.

For ascertainment of prevalent VRFs and CVDs, we referred to baseline verbal interview, documentation of relevant HES codes, or record in UK Biobank algorithmically defined health outcomes (for MI and stroke). For diabetes and high cholesterol, we also referred to biochemistry data (glycosylated haemoglobin >48 mmol/mol and total cholesterol >7 mmol/L, respectively). The approach to ascertainment of VRFs and CVDs along with a full list of ICD codes used is presented in **Supplementary Table 1.**

**Statistical analysis**

Statistical analysis was performed using R version 4.0.324 and RStudio Version 1.3.1093.25 We included all UK Biobank participants with quality-controlled CMR data available.

We performed orthogonal principal component analysis (PCA) rotation of the five CMR metrics (LAVi, LAEF, LVM: LVEDV, GLS, LVGFI), creating uncorrelated CMR variables whilst retaining more than 90% of their individual variance, as described in previous work.26–28 Thus, we removed significant interdependencies between the CMR metrics and were able to include the rotated CMR variables together as exposures in the same model. For inclusion to the PCA, LAV and LAEF contained 0.03% missing values, and GLS contained 3.7% missing values that were imputed with the mean. For comparison, we created a separate set of PCA rotated CMR metrics replacing LVGFI with LVEF. The PCA loadings are presented in **Supplementary Table 2** and **Supplementary Table 3**. We estimated the independent association of the PCA rotated CMR metrics with VRFs (hypertension, diabetes, high cholesterol, smoking) and prevalent CVDs (AF, stroke, IHD, MI) in multivariable logistic regression models simultaneously modelling all five CMR metrics and adjusting for confounders (age, sex, ethnicity, deprivation, education, physical activity, BMI). We used Cox proportional hazards regression for incident CVDs and mortality outcomes (with covariate adjustment as before). In associations with incident CVDs, we excluded participants who had already had the same outcome prior to CMR.

We also present associations with individually entered raw CMR metrics. For associations with prevalent diseases, we used multivariable linear regression, considering individual raw CMR metrics as the model outcome and VRFs and prevalent CVDs as exposure variables. For incident outcomes, we used Cox proportional hazards regression with raw CMR metrics entered individually as exposure variables. We adjusted for confounders as before.In all models, LAV was log-transformed to remove skew. We corrected for multiple comparisons using a false discovery rate of 0.05 across exposure variables.

**Results**

**Population characteristics**

We studied 25,896 participants for whom CMR data was available (**Supplementary Figure 1)**. Thecohort had an average age of 62.9 (±7.5) years-old; 52% (n=13,488) were women (**Table 1**). The proportion of participants with hypertension, diabetes, high cholesterol, and smoking was 32.7%, 5.7%, 34.5%, and 3.7%, respectively. There was, overall, less socio-economic deprivation than UK national averages. The proportion of participants with prevalent AF, stroke, IHD, and MI at time of CMR was 1.5%, 1.9%, 6.0%, and 2.4%, respectively (**Table 1**).

LA size and function were comparable in men and women after adjustment for body size (**Table 1**). Compared to women, men had, on average, more concentric LV remodelling patterns (higher LVM: LVEDV) and poorer LV function by GLS, LVGFI, and LVEF (**Table 1**). We additionally examined CMR metrics in subsets of participants 1)without VRFs or CVD (healthy), 2)with VRFs, but without CVD, and 3)with CVD (**Figure 1, Supplementary Table 4**). There was a stepwise decline in LV (by LVGFI and GLS) and LA function (by LAEF) from the healthy subset to those with VRFs and to those with CVD (**Figure 1**). Average LVEF was higher in the participants with VRFs compared to the healthy subset, and lower than both subsets in those with CVD. Individuals with VRFs had smaller LAVi than healthy participants; those with CVD had the largest LAVi. The VRF and CVDs subset had higher LVM: LVEDV than the healthy cohort (**Figure 1, Supplementary Table 4**).

Over the average follow-up time of 4.2 ±1.2 years, we observed 180 (0.7%), 178 (0.7%), 530 (2.0%), and 197 (0.8%) incident cases of AF, stroke, IHD, and MI respectively. There were 331 deaths during the available follow-up period; of these, 58 were attributed to CVD. In total 880 (3.4%) participants had at least one incident event, of these 34% (n=297) were women (**Supplementary Table 5**). These participants had higher burden of VRFs than the whole cohort, with hypertension, diabetes, high cholesterol, and smoking documented in 50.5%, 9.9%, 46.7%, and 3.9% respectively (**Supplementary Table 5**).

**Association of CMR metrics with vascular risk factors**

In fully adjusted logistic regression models, including all the PCA rotated CMR metrics, we observed association of all the VRFs with poorer LA function (lower LAEF), with statistically significant relationships observed with diabetes and smoking (**Table 2**). Diabetes, high cholesterol, and smoking were associated with smaller LA sizes (lower LAV), whilst hypertension was associated with larger LA size (**Table 2**). There was significant association of all the VRFs with concentric LV remodelling patterns (higher LVM: LVEDV). Hypertension, diabetes, and smoking were associated with significantly poorer LV function by LVGFI and GLS (**Table 2**). In mutually adjusted models with LVEF instead of LVGFI, diabetes was associated with significantly lower LVEF; associations of LVEF with other VRFs were not statistically significant (**Supplementary Table 6**). There was a similar pattern of associations in models using raw CMR metrics entered individually (**Supplementary Table 7)**.

**Association of CMR metrics with prevalent cardiovascular disease**

In fully adjusted logistic regression models, including all the PCA rotated CMR metrics, all the prevalent CVDs were associated with significantly lower LAEF (**Table 2)**. AF and IHD were associated with significantly larger LA sizes (**Table 2**). As expected, these relationships appeared most dominant for AF (**Table 2**). AF, IHD, and MI were associated with more eccentric LV remodelling pattern (lower LVM: LVEDV). IHD, and MI were associated with poorer LV function by LVGFI (**Table 2**). The same pattern of associations was observed with LVEF in mutually adjusted models with LVEF instead of LVGFI (**Supplementary Table 6**). These relationships were broadly similar in models using individual raw CMR metrics; in these models, MI and AF were additionally associated with significantly poorer GLS, LVGFI, and LVEF, but these relationships were attenuated in the mutually adjusted models (**Supplementary Table 7**).

**Association of CMR metrics with incident cardiovascular disease**

In fully adjusted Cox regression models, with mutual inclusion of all the PCA rotated CMR metrics, poorer LA function (lower LAEF) was associated with significantly higher risk of incidence of all the CVDs considered, specifically AF, stroke, IHD, and MI (**Table 3, Graphical Abstract**). Larger LA size was associated with significantly higher risk of incident AF. More concentric LV remodelling patterns (higher LVM: LVEDV) were associated with significantly increased risk of incident stroke and incident IHD (**Table 3**). Lower LVGFI was associated with significantly higher risk of incident IHD (**Table 3**). In mutually adjusted models with LVEF instead of LVGFI, there was no significant association between LVEF and any of the incident CVDs (**Supplementary Table 8**). In equivalent Cox regression models with raw individually entered CMR metrics, the associations with LA metrics were largely unchanged (**Supplementary Table 9**). In these models, AF, stroke, and IHD were associated with significantly lower LVGFI, stroke and IHD were associated with significantly poorer GLS, and AF was associated with lower LVEF (**Supplementary Table 9**); these relationships (with exception of IHD and LVGFI) were attenuated in models mutually adjusting for all the CMR metrics (**Table 3)**.

**Association of CMR metrics with mortality outcomes**

In fully adjusted Cox regression models, including all the PCA rotated CMR metrics, larger LAVi was associated with significantly greater hazard of CVD mortality. Poorer GLS was associated with significantly higher risk of all-cause mortality. Lower LVGFI was associated with significantly higher risk of both all-cause and CVD mortality (**Table 3**). In mutually adjusted models with LVEF instead of LVGFI, LVEF was also associated with significantly lower risk of all-cause and CVD mortality, but with slightly smaller effect sizes than LVGFI (**Supplementary Table 8, Table 3**).

**Discussion**

**Summary of findings**

In this study of 25,896 UK Biobank participants, we demonstrate associations of CMR derived LA structure and function metrics with VRFs, prevalent CVDs, incident CVDs and mortality outcomes, independent of LV measures and a wide range of clinical confounders.

Lower LAEF emerged as a consistent and independent indicator of VRFs (diabetes, smoking) and prevalent and incident CVDs (AF, stroke, IHD, MI). Diabetes, high cholesterol, and smoking were associated with smaller LAV. Hypertension and IHD were associated with larger LAV, perhaps reflecting more advanced diastolic dysfunction in these conditions. Both prevalent and incident AF were associated with large LA sizes. More concentric LV remodelling patterns were associated with VRFs and incident CVDs, whilst prevalent CVDs were associated with more eccentric LV remodelling. These observations likely reflect differential dominance of LV pressure and volume overload in the transition from risk factor to disease, with volume overload becoming dominant after disease occurrence. LVGFI, GLS, and LVEF provided good indications of VRFs and prevalent CVDs, with LVGFI showing the most consistent results. LVGFI and LVEF were independent predictors of all-cause and CVD mortality, with larger effect sizes observed with LVGFI. Higher LAVi was independently associated with significantly higher CVD mortality.

Both the LV and LA metrics were independently informative in associations with risk factors and prevalent disease. In associations with incident outcomes many of the LV associations were attenuated, whilst LAEF associations with all incident CVDs remained robust independent of LV metrics and other confounders. Larger LAVi appeared a strong independent predictor for CVD mortality. These observations demonstrate the independent utility of LA structure and function metrics, particularly for prediction of incident outcomes, which likely reflects pre-clinical LA remodelling before establishment of LV alterations.

**Comparison with existing research**

We observed strong and significant associations of lower LAEF and larger LAV with both prevalent and incident AF. Consistently, Bertelsen et al.29 also demonstrate significant association of larger CMR derived LA volumes and poorer LA function with greater risk of AF detected on an implantable loop recorder in 203 participants with stroke risk factors but without pre-existing AF.29 These LA alterations likely reflect underlying atrial remodelling, which predisposes to (and can also occur as a result of) AF. In a study of 1,148 MESA (Multi-Ethnic Study of Atherosclerosis) participants, Heckbert et al.30 demonstrate association of lower total LAEF and larger LAV with greater burden of premature atrial contractions on ambulatory electrocardiographic monitoring; such arrhythmias may be precursors of AF and indicative of atrial fibrosis. Indeed, in a study of 111 patients without a prior history of atrial arrhythmia, Quail et al.11 demonstrate association of LA late gadolinium enhancement (a marker of atrial fibrosis) with incident atrial arrhythmias.

We observed, association of poorer LAEF with both prevalent and incident stroke independent of other CMR metrics. Larger LAV was associated with significantly greater risk of incident stroke in individual models, but not in models including other CMR metrics. In a study of 169 patients with AF referred for catheter ablation, Inoue et al.12 similarly demonstrate association of poorer LA function (LAEF) with prior stroke or transient ischaemic attack. Habibi et al.31 also report significant association of lower LAEF, but not LA size, with incident ischaemic stroke in 4,261 MESA participants. We additionally demonstrate significant associations of lower LAEF with incident IHD and incident MI, independent of other CMR metrics. In a study of 536 diabetic MESA participants without clinical CVD, Markman et al.32 also demonstrate association of poorer LA function with incident CVD (defined as composite of MI, resuscitated cardiac arrest, angina, stroke, heart failure, or AF). Our findings add to the literature by demonstrating specific independent association of larger LAVi and higher CVD mortality risk.

We observed association of diabetes with smaller LAV and lower LAEF, along with more concentric LV remodelling and poorer LV function metrics. Two studies of small diabetic cohorts have also demonstrated lower LAEF in diabetics compared to controls, but have not demonstrated any significant difference in LA size.10,33 Similar to our observations, studies using the first release of the UK Biobank CMR data have demonstrated association of diabetes with smaller atrial volumes.34,35 Conversely, some echocardiography studies have demonstrated association of diabetes with larger LA sizes. For example, Armstrong et al.9 demonstrate association of larger LA diameter with prevalent diabetes in 2,903 CARDIA study participants. LA alterations evolve with disease progression, with LA dilatation reflecting persistently elevated LV filling pressures and advancement of diastolic (and systolic) LV dysfunction.3 Thus, the duration of exposure to and control of the diabetes, as well as the overall risk factor profile of participants likely influence associations with LA size. As the UK Biobank comprises a relatively healthy cohort, our observations reflect milder disease. Indeed, we observed significant association of larger LAV with pre-existing IHD, a condition associated with more advanced LV impairment. This is further supported by the observed association of larger LAV with greater CVD mortality risk. In our study, and in existing literature, LAEF appears as a reliable and consistent indicator of diabetes and other key morbidities.

We observed the association of more concentric LV remodelling patterns with VRFs and incident CVDs, whilst prevalent CVDs were associated with more eccentric LV remodelling patterns. These observations likely reflect predominance of pressure overload in the presence of VRFs and prior to disease occurrence, but dominance of volume overload after development of clinical cardiovascular disease. Our analysis also demonstrates consistent and significant association of lower LVGFI with VRFs, prevalent CVDs, incident IHD, and higher risk of all-cause and CVD mortality. In separate analyses comparing LVGFI to LVEF, the latter showed similar but less consistent associations. Our findings add strength to existing studies which have proposed the high utility of this LVGFI as a measure of LV function.20,21

**Clinical implications**

Inthis study of 25,896 UK Biobank participants, we describe independent clinical associations of CMR derived measures of LA structure and function (LAV, LAEF). These metrics, particularly LAEF, show robust associations with key cardiovascular outcomes independent of LV measures. Thus, there is potential utility for these metrics as components of clinical risk prediction algorithms. In the next stages towards development of such clinical models, there is need for evaluation of clinical relationships in other cohorts and settings. Any proposed clinical risk stratification models will require careful validation and evaluation of model performance prior to use in clinical practice.

**Strengths and limitations**

The highly detailed participant characterisation and standardised CMR data in the UK Biobank permitted evaluation of associations of CMR phenotypes with key VRFs and CVDs in a very large cohort, whilst considering a wide range of confounders. The linked reliably recorded health outcome data also permitted assessment of associations with incident CVDs. The duration of follow up was relatively short and the proportion of participants with incident events was small (n=880/ 25,896, 3.4%), and even fewer when considering subgroups of participants (**Supplementary Table 10**). Given that this limits our power to detect statistically significant associations with incident events, the observed significant relationships between LA metrics and incident CVDs are all the more notable. Identification of incident outcomes using HES is ideal for conditions such as acute MI and stroke, which almost always require hospitalisation. However, this approach is not optimal for endpoints that do not always require hospital admission, such as diastolic heart failure or mitral valve disease, which we were unable to consider in the analysis. There were few CVD mortality events, which means that analysis with this outcome is likely underpowered to appreciate the full picture of CMR associations (particularly small and moderate effect sizes). As events accrue in the UK Biobank, more adequately powered analyses may be conducted with possibility of evaluating associations with more granular disease-specific mortality outcomes. Finally, due to the observational nature of the study, we cannot exclude residual confounding or reverse causation, however, the primary aim of the present study is description of associations rather than causal inference.

**Conclusions**

LA structure and function measures (LAEF, LAV) demonstrate significant associations with key prevalent and incident cardiovascular outcomes, independent of LV metrics. These measures have potential clinical utility for disease discrimination and outcome prediction.

**Ethics statement**

This study complies with the Declaration of Helsinki; the work was covered by the ethical approval for UK Biobank studies from the NHS National Research Ethics Service on 17th June 2011 (Ref 11/NW/0382) and extended on 18 June 2021 (Ref 21/NW/0157) with written informed consent obtained from all participants.

**Data Availability Statement**

The data underlying this article were provided by the UK Biobank under access application 2964. UK Biobank will make the data available to bona fide researchers for all types of health-related research that is in the public interest, without preferential or exclusive access for any persons. All researchers will be subject to the same application process and approval criteria as specified by UK Biobank. For more details on the access procedure, see the UK Biobank website: http://www.ukbio bank.ac.uk/register-apply/.

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**Conflicts of interest**

SEP provides consultancy to and owns stock of Cardiovascular Imaging Inc, Calgary, Alberta, Canada.

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**Figure legends**

**Figure 1 Title:** CMR metric means, and 95% confidence interval of the mean stratified by disease status

**Figure 1 Legend:** Within the “Healthy”, “VRFs”, and “CVD” subsets, we include participants without prevalent CVD or VRFs, with VRFs but without prevalent CVDs, and with prevalent CVDs, respectively. CMR: cardiovascular magnetic resonance, CVD: cardiovascular disease; LVGFI: left ventricular global function index, GLS: global longitudinal strain, i: indicates indexation to body surface area, LAEF: left atrial ejection fraction, LAV: maximum left atrial volume, LVEDV: left ventricular end-diastolic volume, LVEF: left ventricular ejection fraction, left ventricular mass, MI: myocardial infarction.

**Table 1. Participant characteristics**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Whole sample** | **Men** | **Women** |
|  | **(n= 25,896)** | **(n= 12,408)** | **(n= 13,488)** |
| Age at imaging (years) | 62.9 (±7.5) | 63.6 (±7.6) | 62.2 (±7.4) |
| Townsend deprivation index | -2.7 [-3.9, -0.7] | -2.7 [-4.0, -0.7] | -2.6 [-3.9, -0.7] |
| Education |  |  |  |
| Left school ≤14yrs without qualifications | 67 (0.3%) | 38 (0.3%) | 29 (0.2%) |
| Left school ≥ 15yrs without qualifications | 1,844 (7.1%) | 855 (6.9%) | 989 (7.3%) |
| Secondary school qualification | 3,485 (13.5%) | 1,313 (10.6%) | 2,172 (16.1%) |
| A levels / AS levels or equivalent | 1,465 (5.7%) | 653 (5.3%) | 812 (6.0%) |
| Other professional qualification | 7,258 (28.0%) | 3,666 (29.5%) | 3,592 (26.6%) |
| Higher education (e.g., university) degree | 11,511 (44.5%) | 5,755 (46.4%) | 5,756 (42.7%) |
| Missing | 266 (1.0%) | 128 (1.0%) | 138 (1.0%) |
| BMI (kg/m2) | 25.9 [23.5, 28.9] | 26.5 [24.3, 29.1] | 25.3 [22.8, 28.6] |
| Physical activity (summed MET-minutes/week) | 1899 [896, 3573] | 1971 [958, 3666] | 1838 [834, 3514] |
| Smoker (current) | 960 (3.7%) | 540 (4.4%) | 420 (3.1%) |
| Hypertension | 8,471 (32.7%) | 4,914 (39.6%) | 3,557 (26.4%) |
| High cholesterol | 8,947 (34.5%) | 5,117 (41.2%) | 3,830 (28.4%) |
| Diabetes | 1,485 (5.7%) | 924 (7.4%) | 561 (4.2%) |
| Prevalent cardiovascular disease |  |  |  |
| Atrial fibrillation  | 386 (1.5%) | 273 (2.2%) | 113 (0.8%) |
| Stroke  | 503 (1.9%) | 320 (2.6%) | 183 (1.4%) |
| IHD  | 1,560 (6.0%) | 1,092 (8.8%) | 468 (3.5%) |
| MI  | 633 (2.4%) | 502 (4.0%) | 131 (1.0%) |
| Incident CVD and mortality outcomes  |  |  |  |
| Atrial fibrillation   | 180 (0.7%) | 127 (1.0%) | 53 (0.4%) |
| Stroke   | 178 (0.7%) | 114 (0.9%) | 64 (0.5%) |
| IHD   | 530 (2.0%) | 347 (2.8%) | 183 (1.4%) |
| MI   | 197 (0.8%) | 140 (1.1%) | 57 (0.4%) |
| All-cause mortality | 331 (1.3%) | 220 (1.8%) | 111 (0.8%) |
| CVD mortality | 58 (0.2%) | 44 (0.4%) | 14 (0.1%) |
| Any of AF, Stroke, IHD, MI or CVD death | 880 (3.4%) | 583 (4.7%) | 297 (2.2%) |
| CMR Metrics |  |  |  |
| LAV (ml) | 70.0 [57.0, 85.3] | 75.9 [61.1, 92.6] | 65.9 [54.5, 78.7] |
| LAVi (ml/m2) | 38.0 [31.5, 45.4] | 38.0 [30.9, 46.0] | 38.1 [31.9, 45.0] |
| LAEF (%) | 61.3 (±9.1) | 60.6 (±9.6) | 61.9 (±8.5) |
| LVM: LVEDV (g/ml) | 0.57 [0.52, 0.63] | 0.60 [0.56, 0.66] | 0.54 [0.50, 0.59] |
| LVSVi (ml/m2) | 47.1 (±8.4) | 48.8 (±9.1) | 45.6 (±7.4) |
| LVEF (%) | 59.6 (±6.1) | 57.8 (±6.1) | 61.1 (±5.5) |
| LVGFI (%) | 47.7 (±6.8) | 44.8 (±6.2) | 50.4 (±6.2) |
| GLS (%) | -18.5 (±2.7) | -17.8 (±2.6) | -19.1 (±2.7) |

**Table 1 footnote.** Counts variables are presented as number (percentage), continuous variables as mean (standard deviation) or median [interquartile range] based on skew. BMI: body mass index, CMR: cardiovascular magnetic resonance, CVD: cardiovascular disease, GLS: global longitudinal strain, i: indicates indexation to body surface area, IHD: ischaemic heart disease, LAEF: left atrial ejection fraction, LAV: maximum left atrial volume, LVEDV: left ventricular end-diastolic volume, LVEF: left ventricular ejection fraction, LVM: left ventricular mass, LVGFI: left ventricular global function index, MET: metabolic equivalent, MI: myocardial infarction

**Table 2. Associations of mutually adjusted CMR metrics with vascular risk factors and prevalent cardiovascular disease in multivariable logistic regression models with full confounder adjustment**

|  |  |  |
| --- | --- | --- |
|  | **Vascular risk factors** | **Prevalent cardiovascular disease** |
| **CMR Metric** | **Hypertension** | **Diabetes** | **High cholesterol** | **Smoking (current)** | **AF** | **Stroke** | **IHD** | **MI** |
| LAVi (ml/m2) | 1.24\* | 0.87\* | 0.96\* | 0.88\* | 1.30\* | 0.96 | 1.14\* | 1.09 |
|  | [1.21, 1.28] | [0.83, 0.92] | [0.94, 0.99] | [0.83, 0.94] | [1.18, 1.44] | [0.88, 1.05] | [1.08, 1.20] | [1.01, 1.18] |
|  | 7.59$×$10-46 | 1.31$×$10-6 | 0.0129 | 1.65$×$10-4 | 4.15$×$10-7 | 0.3661 | 2.73$×$10-6 | 0.0377 |
| LAEF (%) | 0.99 | 0.94\* | 0.99 | 0.93\* | 0.40\* | 0.88\* | 0.82\* | 0.82\* |
|  | [0.96, 1.02] | [0.89, 0.98] | [0.96, 1.02] | [0.87, 0.99] | [0.36, 0.43] | [0.82, 0.96] | [0.78, 0.86] | [0.76, 0.88] |
|  | 0.6064 | 0.0110 | 0.3950 | 0.0266 | 6.15$×$10-91 | 0.0027 | 1.36$×$10-14 | 3.82$×$10-8 |
| LVM: LVEDV | 1.43\* | 1.20\* | 1.10\* | 1.29\* | 0.80\* | 1.04 | 0.85\* | 0.75\* |
|  | [1.38, 1.48] | [1.14, 1.27] | [1.07, 1.14] | [1.21, 1.38] | [0.71, 0.90] | [0.95, 1.14] | [0.81, 0.91] | [0.68, 0.82] |
|  | 2.40$×$10-99 | 1.10$×$10-11 | 6.28$×$10-9 | 1.05$×$10-13 | 2.16$×$10-4 | 0.4062 | 1.19$×$10-7 | 1.52$×$10-10 |
| LVGFI (%) | 0.93\* | 0.87\* | 1.00 | 0.88\* | 1.23\* | 0.95 | 0.88\* | 0.71\* |
|  | [0.90, 0.96] | [0.82, 0.92] | [0.96, 1.03] | [0.82, 0.94] | [1.11, 1.37] | [0.87, 1.05] | [0.84, 0.94] | [0.65, 0.77] |
|  | 1.07$×$10-5 | 1.89$×$10-6 | 0.7519 | 2.43$×$10-4 | 8.50$×$10-5 | 0.3310 | 2.63$×$10-5 | 7.96$×$10-16 |
| GLS (%) | 1.03\* | 1.15\* | 0.97 | 1.12\* | 1.11 | 1.04 | 1.01 | 1.07 |
|  | [1.00, 1.07] | [1.09, 1.21] | [0.94, 1.00] | [1.05, 1.20] | [1.01, 1.22] | [0.96, 1.14] | [0.95, 1.06] | [0.99, 1.16] |
|  | 0.0251 | 9.34$×$10-7 | 0.0382 | 9.47$×$10-4 | 0.0387 | 0.3420 | 0.8414 | 0.0836 |

**Table 2 footnote.** Results are odds ratios, 95% confidence intervals, and p-values. Models are logistic regression models with disease of interest entered as the response (outcome) variable. For the vascular risk factor models, covariates include mutually entered PCA rotated CMR metrics (LAV, LAEF, LVM/LVEDV, GLS, LVGLFI), age, sex, ethnicity, deprivation, education, body mass index, physical activity, and all the VRFs (except the one set as the model outcome). For the prevalent cardiovascular disease models covariates include mutually entered PCA rotated CMR metrics (LAV, LAEF, LVM/LVEDV, GLS, LVGLFI), age, sex, ethnicity, deprivation, education, body mass index, physical activity, hypertension, high cholesterol, diabetes, and smoking. \*indicates statistically significant p-values with a false discovery rate of 0.05, giving an approximate threshold of 0.025 for this table. AF: atrial fibrillation, CMR: cardiovascular magnetic resonance, LVGFI: left ventricular global function index, GLS: global longitudinal strain, i: indicates indexation to body surface area, IHD: ischaemic heart disease, LAEF: left atrial ejection fraction, LAV: maximum left atrial volume, LVEDV: left ventricular end-diastolic volume, LVM: left ventricular mass, MI: myocardial infarction, PCA: principal component analysis.

**Table 3. Associations of mutually adjusted CMR metrics with incident cardiovascular disease and mortality outcomes in Cox proportional hazard models with full confounder adjustment**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **CMR Metric** | **AF** | **Stroke** | **IHD** | **MI** | **All-cause mortality** | **CVD mortality** |
| LAVi (ml/m2) | 1.47\* | 1.13 | 1.10 | 1.06 | 1.11 | 1.34\* |
|  | [1.28, 1.70] | [0.98, 1.31] | [1.01, 1.19] | [0.93, 1.21] | [1.00, 1.23] | [1.05, 1.71] |
|  | 8.46x10-8 | 0.0807 | 0.0302 | 0.4002 | 0.0407 | 0.0185 |
| LAEF (%) | 0.64\* | 0.83\* | 0.88\* | 0.87\* | 0.96 | 0.85 |
|  | [0.56, 0.73] | [0.73, 0.95] | [0.81, 0.95] | [0.76, 0.99] | [0.87, 1.06] | [0.69, 1.04] |
|  | 2.50$×$10-11 | 0.0060 | 9.95$×$10-4 | 0.0294 | 0.4029 | 0.1119 |
| LVM: LVEDV | 1.06 | 1.22\* | 1.27\* | 1.14 | 1.09 | 1.05 |
|  | [0.92, 1.23] | [1.06, 1.40] | [1.17, 1.37] | [0.99, 1.30] | [0.98, 1.22] | [0.82, 1.35] |
|  | 0.4036 | 0.0065 | 1.27x10-8 | 0.0732 | 0.0959 | 0.7039 |
| LVGFI (%) | 0.92 | 0.89 | 0.88\* | 0.95 | 0.85\* | 0.61\* |
|  | [0.79, 1.06] | [0.76, 1.03] | [0.80, 0.96] | [0.82, 1.11] | [0.76, 0.95] | [0.48, 0.78] |
|  | 0.2521 | 0.1266 | 0.0063 | 0.5372 | 0.0050 | 5.95$×$10-5 |
| GLS (%) | 0.97 | 1.10 | 1.08 | 1.05 | 1.14\* | 1.10 |
|  | [0.85, 1.12] | [0.95, 1.27] | [0.99, 1.17] | [0.91, 1.20] | [1.02, 1.27] | [0.87, 1.38] |
|  | 0.7171 | 0.1880 | 0.0888 | 0.5254 | 0.0170 | 0.4314 |

**Table 3 footnote.** Results are hazard ratios, 95% confidence intervals, and p-values. Covariates are: LAV, LAEF, LVM/LVEDV, GLS, GLFI, age, sex, ethnicity, deprivation, education, body mass index, hypertension, high cholesterol, diabetes, physical activity, smoking. The CMR variables are principal component analysis rotated variables. \*indicates statistically significant p-values with a false discovery rate of 0.05, giving an approximate threshold of 0.028 for this table. AF: atrial fibrillation, CMR: cardiovascular magnetic resonance, CVD: cardiovascular disease; LVGFI: left ventricular global function index, GLS: global longitudinal strain, i: indicates indexation to body surface area, IHD: ischaemic heart disease, LAEF: left atrial ejection fraction, LAV: maximum left atrial volume, LVEDV: left ventricular end-diastolic volume, LVM: left ventricular mass, MI: myocardial infarction.

**Figure 1. CMR metric means, and 95% confidence interval of the mean stratified by disease status**



**Figure 1.** Within the “Healthy”, “VRFs”, and “CVD” subsets, we include participants without prevalent CVD or VRFs, with VRFs but without prevalent CVDs, and with prevalent CVDs, respectively. CMR: cardiovascular magnetic resonance, CVD: cardiovascular disease; LVGFI: left ventricular global function index, GLS: global longitudinal strain, i: indicates indexation to body surface area, LAEF: left atrial ejection fraction, LAV: maximum left atrial volume, LVEDV: left ventricular end-diastolic volume, LVEF: left ventricular ejection fraction, LVM: left ventricular mass.