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A central principle in cardiovascular disease (CVD) management is that the first lifetime diagnosis signals the failure of primary prevention and the need to initiate secondary prevention of recurrent or related CVD events. The decades-long emphasis given to prevention of myocardial infarction (MI) and stroke is reflected in remarkable declines—≈33% over the past decade—in their incidence in developed countries. Incidence rates for chronic CVD presentations such as angina or heart failure, although less studied, do not appear to have similarly declined. Consequently, the spectrum of initial presentations of CVD in contemporary practice is likely to have changed in comparison with the latter part of the last century. Cohort studies that report only fatal end points (final presentations), may have less relevance to informing the success of primary prevention than those which investigate initial presentations. Within studies that incorporate nonfatal events, acute MI and stroke have been more commonly investigated.
than other chronic presentations.5–7 Large-scale contemporary studies that evaluate the first lifetime diagnosis in women and men across a wide range of acute and chronic CVDs including both fatal and nonfatal presentations can provide additional insight into the understanding of CVDs.

Fundamental unanswered questions about initial CVD presentation arise. First, what is the relative frequency of different CVDs as they affect women and men in contemporary practice? Second, is male sex an equally strong risk factor common to all CVDs, or does the association differ across a range of diseases?

The lack of large, contemporary, population-based cohorts with detailed clinical follow-up spanning hospital and ambulatory care has hindered the study of the initial presentation of a wide range of acute and chronic CVDs. It has been suggested that electronic health record (EHR) data might be meaningfully reused to create mega-cohorts for such research.9 We studied a contemporary, population-based cohort based on linked EHRs across primary, secondary, disease registry, and death records10–13 to address these 2 questions. We investigated a wide range of acquired symptomatic CVDs that are recognized to have differing pathogenic mechanisms.

Methods

Data Sources

Anonymized patients were selected from the Cardiovascular Research Using Linked Bespoke Studies and Electronic Records (CALIBER) program, described14 and validated10–13 elsewhere. Patients were linked across 4 clinical data sources: the Clinical Practice Research Database (CPRD), the Myocardial Ischemia National Audit Project registry, Hospital Episodes Statistics, and the national death registry from the Office for National Statistics. CPRD provides primary care data on anthropometric measurements, laboratory tests, medical history, clinical diagnoses, prescriptions, medical procedures, and health behaviors, coded using the Read clinical coding scheme. Patients registered in practices submitting linkable data to CPRD, covering ≈4% of the English population, have been found to be representative of the English population in terms of age, sex, and ethnicity.16,17 The Myocardial Ischemia National Audit Project is a national registry of patients admitted to the hospital with acute coronary syndromes. Hospital Episodes Statistics provides information on diagnoses and medical procedures related to all elective and emergency hospital admissions across all National Health Service hospitals in England.

Study Population

We studied 1937360 patients from 225 general practices across England registered between January 1997 and March 2010. We required that at study entry patients were aged ≥30 years, were free of diagnosed CVD, and had been followed up for at least 1 year. We used the entire medical history available on each patient to confirm they were free of diagnosed CVD. The look-back period ranged from 20 years to the minimum of 1 year, which previous research has indicated is a sufficient period to ensure accurate assessment of initial disease presentations.18 We used an open cohort design, so patients effectively entered the study when they met the inclusion criteria. Patients were censored on the earliest date from among: the date of first CVD presentation, date of death from other causes, date leaving the practice, or date of last practice data collection. (See Figure I in the online-only Data Supplement for study flow diagram.)

Risk Factors

The exposures of interest were sex and baseline age, analyzed as 10-year age groups between 30 and 80. A priori confounders were baseline age as a continuous variable (in analyses estimating associations with sex), smoking status, body mass index, systolic blood pressure, total and high-density lipoprotein cholesterol, diabetes mellitus, socioeconomic status (based on area deprivation measure), use of statins, use of blood pressure medication, and, in women only, use of oral contraceptives or hormone replacement therapy. The baseline value for these confounders was taken as the most recent measurement as recorded during consultations in primary care (CPRD) up to 1 year before study entry. (Detailed definitions are in online-only Data Supplement Methods I.)

End Points

Primary end points were defined as the first recorded diagnosis of the 12 most common symptomatic manifestations of CVD, irrespective of underlying disease mechanism, arising from pathology in the head, heart, abdomen, or legs. The first diagnosis could occur in primary care, secondary care, or at death. We studied the following CVDs: stable angina, unstable angina, nonfatal MI, unheralded coronary death (UCD), heart failure, a composite of cardiac arrest, ventricular arrhythmia, and sudden cardiac death (SCD), transient ischemic attack, ischemic stroke, subarachnoid hemorrhage (SAH), intracerebral hemorrhage, abdominal aortic aneurysm (AAA), peripheral arterial disease (PAD), composite CVD, and other deaths. In secondary analysis, we examined associations in a subset of nonfatal MIs that were classified into ST-segment–elevation MI and non–ST-segment–elevation MI. Coronary heart disease (CHD) and stroke that were not otherwise specified (NOS) were also studied. We classified as fatal events where a death record exists for the same calendar date. (Overview of codes and data sources used to define cardiovascular end points available in online-only Data Supplement Methods II.)

Statistical Analysis

Hazard ratios (HRs) were estimated for the disease-specific Cox proportional-hazards models with length of follow-up as the timescale, stratified by practice, with women as the reference category, and included interactions between age (linear and quadratic term) and sex. Where we estimated the HR for baseline age, we additionally stratified by sex, to allow the baseline hazard to vary. The proportional hazard assumption was tested using Schoenfeld residuals, with no significant effects found.

In the main analyses, we estimated the association of each end point with age groups, the age-adjusted association with sex across all subjects, and by age group in a model with sex interactions. Assuming mutual independence between initial presentations, we assessed heterogeneity in the reported associations based on r², an estimate of the between-group variance of the log hazard ratio, and a way of summarizing the variability in effect sizes across all the end points in a single statistic.19

In secondary analysis, we examined whether associations with sex change after adjusting for smoking status, body mass index, diabetes mellitus, systolic blood pressure, total cholesterol, high-density lipoprotein cholesterol, and social deprivation, or additionally for baseline use of blood pressure–lowering medications (diazides, β-blockers, angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, or calcium channel blockers), statins, oral contraceptives and hormone replacement therapy. Missing covariate data were handled by multiple imputation. (Methods used for multiple imputation are described in online-only Data Supplement Methods III.) In sensitivity analyses we studied associations between sex and CVDs (1) ignoring primary care diagnoses and (2) restricting end points to fatal events. In a post hoc analysis, we assessed the discrimination of age- and sex-adjusted models for each of the 12 end points by calculating the separate concordance index (C-index) for each.20

Approval was granted by the Independent Scientific Advisory Committee of the Medicines and Healthcare Products Regulatory Agency and the Myocardial Ischemia National Audit Project Academic Group. We registered the protocol at clinicaltrials.gov (NCT01164371).
Results
Baseline characteristics of the cohort are shown in the Table. The cohort was young at baseline, as would be expected from a population free from CVD, and 90% were white. Both systolic and diastolic blood pressure increased with age, as did the proportion on blood pressure–lowering medication, with more women than men treated at all ages. More men than women were current or ex-smokers, the proportion of current smokers declining at >60 years of age. Rates of statin prescription were low, but were higher in men than in women at all ages.

Initial CVD Presentations
Over a 6-year median follow-up (interquartile range, 2–10), 114 859 initial CVD presentations were observed (52.3% in men), among which nonfatal MI, UCD, and ischemic or NOS stroke together accounted for 32.5%. The proportion of events varied by sex and age group (Figure 1; Table I in the online-only Data Supplement). The most frequent initial CVD presentation for men was nonfatal MI, which accounted for 27.9% of events in the 30 to 39 age group and more than double the proportion in women in the same age group (11.2%). This proportion declined in men as age increased, becoming similar to that in women in the >80 age group. In contrast, stable angina and unstable angina accounted for similar proportions of initial presentations in both men and women and declined with age. Although evident in younger age groups, heart failure and ischemic stroke as an initial presentation started to increase in both sexes at age 60 to form the 2 most common initial presentations at age >80.

Associations With Age
The strength and shape of the association of CVDs with age varied from predominantly linear (in angina and nonfatal MI) to strongly quadratic (UCD, stroke, AAA), and from weak (SAH, unstable angina, and cardiac arrest/SCD) to very strong (heart failure and AAA. (See Figure II in the online-only Data Supplement.)

Associations With Sex
SAH was less common in men (HR men versus women, 0.69; 95% confidence interval [CI], 0.59–0.79); other CVDs were positively associated with male sex but with considerable heterogeneity ($\tau^2=0.196$; Figure 2). Specifically, the age-adjusted HR (all $P<0.001$) was <1.5 for transient ischemic attack, intracerebral hemorrhage, and unstable angina, 1.5 to 2.0 for stable angina, ischemic stroke, PAD, heart failure, and cardiac arrest/SCD, and 3.6 to 5.0 for AAA, MI, and UCD. The age-adjusted HR for men versus women was 4.14 (95% CI, 3.72–4.60) in ST-segment–elevation MI and 3.18 (95% CI, 2.86–3.52) in non–ST-segment–elevation MI. These associations changed little after adjustment for conventional CVD risk factors and baseline medications, with the exception of intracerebral hemorrhage, where the association reduced to null (Figure III in the online-only Data Supplement).

Table. Baseline Characteristics in Men and Women by 10-Year Age Groups

<table>
<thead>
<tr>
<th></th>
<th>30 to 39</th>
<th>40 to 49</th>
<th>50 to 59</th>
<th>60 to 69</th>
<th>70 to 79</th>
<th>&gt;80</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No. of patients</strong></td>
<td>418 755</td>
<td>385 486</td>
<td>211 109</td>
<td>194 172</td>
<td>159 919</td>
<td>110 478 52</td>
</tr>
<tr>
<td>Age, y</td>
<td>33.3 (3.2)</td>
<td>33.3 (3.2)</td>
<td>44.5 (2.9)</td>
<td>44.6 (2.9)</td>
<td>54.4 (2.9)</td>
<td>54.5 (2.9)</td>
</tr>
<tr>
<td>White</td>
<td>85.7</td>
<td>86.6</td>
<td>89.1</td>
<td>89.2</td>
<td>94.2</td>
<td>93.5</td>
</tr>
<tr>
<td>Most deprived quintile</td>
<td>22.8</td>
<td>21.9</td>
<td>19.7</td>
<td>18.6</td>
<td>17.5</td>
<td>16.6</td>
</tr>
<tr>
<td>Number of GP visits in previous year</td>
<td>3.6 (4.9)</td>
<td>7.1 (6.8)</td>
<td>4.1 (5.5)</td>
<td>6.2 (6.7)</td>
<td>4.4 (6.0)</td>
<td>6.4 (6.8)</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td>26.2 (4.5)</td>
<td>25.5 (5.7)</td>
<td>27.1 (4.6)</td>
<td>26.4 (5.9)</td>
<td>27.5 (4.6)</td>
<td>27.1 (5.7)</td>
</tr>
<tr>
<td>Systolic BP, mm Hg</td>
<td>126 (14)</td>
<td>117 (13)</td>
<td>130 (15)</td>
<td>124 (16)</td>
<td>136 (17)</td>
<td>133 (18)</td>
</tr>
<tr>
<td>Diastolic BP, mm Hg</td>
<td>78 (10)</td>
<td>74 (9)</td>
<td>81 (10)</td>
<td>77 (10)</td>
<td>83 (10)</td>
<td>81 (10)</td>
</tr>
<tr>
<td>Total cholesterol, mmol/L</td>
<td>5.3 (1.1)</td>
<td>4.9 (1.0)</td>
<td>5.5 (1.1)</td>
<td>5.3 (1.0)</td>
<td>5.4 (1.1)</td>
<td>5.7 (1.1)</td>
</tr>
<tr>
<td>HDL, mmol/L</td>
<td>1.2 (0.3)</td>
<td>1.5 (0.4)</td>
<td>1.2 (0.4)</td>
<td>1.5 (0.4)</td>
<td>1.3 (0.4)</td>
<td>1.6 (0.5)</td>
</tr>
<tr>
<td>Ex-smoker</td>
<td>13.2</td>
<td>14.5</td>
<td>15.2</td>
<td>12.7</td>
<td>21.2</td>
<td>14.8</td>
</tr>
<tr>
<td>Current smoker</td>
<td>27.5</td>
<td>21.3</td>
<td>25.5</td>
<td>19.4</td>
<td>22.1</td>
<td>17.3</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>0.9</td>
<td>0.8</td>
<td>2.1</td>
<td>1.4</td>
<td>3.9</td>
<td>2.6</td>
</tr>
<tr>
<td>Statins</td>
<td>0.4</td>
<td>0.2</td>
<td>2.1</td>
<td>1.0</td>
<td>5.0</td>
<td>3.5</td>
</tr>
<tr>
<td>BP-lowering medications</td>
<td>4.2</td>
<td>7.6</td>
<td>8.8</td>
<td>14.1</td>
<td>17.9</td>
<td>25.8</td>
</tr>
<tr>
<td>Oral contraceptives</td>
<td>. . .</td>
<td>35.0</td>
<td>. . .</td>
<td>10.4</td>
<td>. . .</td>
<td>1.2</td>
</tr>
<tr>
<td>HRT</td>
<td>.</td>
<td>0.7</td>
<td>9.1</td>
<td>30.0</td>
<td>15.1</td>
<td>3.3</td>
</tr>
</tbody>
</table>

*Mean (standard deviation) for continuous variables and % for categorical variables. Continuous variables are the most recently recorded value in the year before study entry. BP indicates blood pressure; GP, general practice; HDL, high-density lipoprotein cholesterol; and HRT, hormone replacement therapy.
Associations between sex and initial CVD presentation were differentially modified by age (Figure 3). The largest differences in HRs for men versus women were observed in the younger (coronary end points) and middle (ischemic stroke, PAD, AAA) age groups. Most dramatically, men <60 years old had an >4-fold higher risk of MI or UCD than similarly aged women. In all cases, associations with male sex diminished with age.

Sensitivity Analyses
The pattern and magnitude of associations with sex were similar in multiply-adjusted analyses to analyses adjusted for age alone (see Figure III in the online-only Data Supplement). Stable angina and PAD were the only initial presentations where the association with male sex differed when the EHRs used were restricted to secondary care and mortality (Figure IV in the online-only Data Supplement).

Discrimination of Age- and Sex-Adjusted Models for Different CVDs
Using disease-specific age and sex coefficients in risk prediction models resulted in markedly different discrimination performance (Figure V in the online-only Data Supplement), with C-indices ranging from very low for SAH (0.57; 95% CI, 0.55–0.59) to relatively high for AAA (0.86; 95% CI, 0.85–0.88) in comparison with a conventional composite CVD model with C-index of 0.73 (95% CI, 0.72–0.73).

Discussion
Objectives Addressed, Summary of Main Findings
By linking EHRs from multiple sources we curated a cohort of nearly 2 million patients with >100,000 nonfatal and fatal CVD end points of 12 different types. We found that the majority of CVD first presentations are not MI or ischemic stroke but rather heart failure, angina, transient ischemic attack, and PAD. In our contemporary population-based cohort, we find that 51.3% of men and 41.2% of women experienced some form of CVD during their lifetime, with heart failure and stroke (primarily ischemic and NOS) becoming more common as the initial presentations in both men and women in later life. The variable associations of sex and age with different CVDs have important consequences for risk prediction.

Importance of Studying First Manifestations of CVD
We compared the relative frequency of 12 of the most common CVDs affecting atherothrombotic processes in the coronary, cerebral, and peripheral circulations, aneurysms in the cerebral and peripheral circulations, and disorder of myocardial function and cardiac arrhythmia. This family of diseases is clinically relevant, because having one is strongly associated with the subsequent development of another and should initiate a range of secondary preventive interventions.
The size of cohort—nearly 2 million people—pos-
corresponds with a family physician,28 unlike other recent large
registered with a family physician,28 unlike other recent large
restricted to small cohorts, or reported in
restricted to small cohorts, or reported in

**Innovative Role of Large-Scale Health Record Linkages**

Through the use of linked EHRs, we were able to capture diseases first presenting in primary care and were not con-
fined to hospitalized cases. Our cohort is population based,
with >99% of the English population estimated to be reg-
istered with a family physician,28 unlike other recent large
cohorts, such as UK Biobank,29 and precision medicine initiatives 32 all place
the Research Program for Genes Environment and Health
men only.23–27

Despite the insights to be gained from considering the first
presentation among these diseases together, this first-life-
time-presentation approach has rarely been reported in the
literature22 and has tended to exclude major diseases such as
heart failure, been restricted to small cohorts, or reported in
men only.23–27

**Figure 2.** Hazard ratios of men in comparison with women for initial presentation of 12 different cardiovascular diseases among a population of 1.93 million adults. CHD indicates coronary heart
disease; CI, confidence interval; CVD, cardiovascular disease; HR, hazard ratio; NOS, not otherwise specified; and SCD, sudden cardiac death.

The vertical grey dotted line corresponds to the HR of the composite CVD endpoint. CHD NOS and Stroke NOS excluded from the main display because non-specific endpoints; their corresponding estimates are HR 2.03 (95% CI, 1.92-2.15; n=10,895)
and 1.37 (95% CI, 1.26-1.49; n=9,532).

**Male Sex as a Risk Factor for Different CVDs**

We demonstrate that male sex does not have a common under-
lying association on the incidence of different CVDs. Rather,
the strength of this association is highly variable, ranging from
protective for SAH; minor for transient ischemic attack, intra-
cerebral hemorrhage, and unstable angina; moderate for stable angina, ischemic stroke, PAD, heart failure, and cardiac arrest/SCD; and strong for AAA, MI, and UCD. Additionally, we found that these associations change with age, with sex differences in proportion of initial presentation of MI and coronary death reducing with age, and with heart failure and stroke (ischemic and NOS) emerging as the most common initial presentations in both sexes. These findings suggest that stratifying patients into low-, intermediate-, and high-risk groups based on their total and disease-specific risks, accompanied by the establishment of new cost-effective treatment thresholds, could improve risk management, particularly for diseases such as heart failure and

We expect the pattern of the age and sex associations we
found with the CVD outcomes to apply to the broader UK
population and other European populations free from symp-
tomatic CVD. Our patients were drawn from >200 practices representative of the English population. Indeed, a recent arti-
cle investigating similar questions in a smaller investigator-led
Dutch cohort found broadly similar associations, albeit with
fewer end points.22 Different cohorts, especially those with
more people from differing ethnic groups or differing baseline
risk profiles, may well present different associations.

**Validity of Risk Factor and Disease Measurements in EHRs**

Although a principal strength of this study is the ability to
resolve a wide range of CVDs in a large-scale cohort, the prin-
cipal limitation is the possibility of errors in the individual
EHR data sources.34,35 However, evidence for the validity of our
risk factor and disease end points comes from several sources.
First, in this population, using identical phenotypic definitions
for these same 12 diseases, we have replicated anticipated risk factor – disease associations with systolic and diastolic blood
pressure,11 type 2 diabetes mellitus,15 smoking,10 and socio-
economic deprivation.12 These findings support the prospec-
tive prognostic validity of both the risk factor and the disease
measurements. Second, a recent systematic review of studies
validating diagnoses in CPRD found a median positive predic-
tive value of 88% across a wide range of diagnoses,7 whereas
a separate systematic review found the accuracy of discharge
coding in Hospital Episodes Statistics to be 83%.35 Third, the
associations we found when considering events from all data
sources (Figure V in the online-only Data Supplement) were
consistent with those when excluding nonfatal cases or those
from primary care. The doctors and coders responsible, and
the information on which these diagnoses are based, differ for
each data source (primary care, hospital, and death); it was
reassuring that the associations were broadly similar. Finally,
we33 and others36 have demonstrated the validity of using
linked data for end point follow-up.
stroke that affect high proportions of women but are undermanaged based on current clinical risk assessment.39

Clinical Implications and Risk Prediction
Current risk algorithms in common use focus on CHD40 and CVD,41 as does the new American College of Cardiology/American Heart Association Guideline on the Assessment of Cardiovascular Risk,42 yet we show that chronic disease, such as heart failure and PAD, account for a substantial proportion of initial CVD presentations in contemporary practice. These diseases are associated with marked increased risk of subsequent events and death, yet have been excluded from many risk prediction algorithms. Given the recent decline in the incidence of acute events of MI and stroke, our findings raise the question of whether risk algorithms should take account of the current burden of CVDs and, in efforts to personalize cardiovascular risk, whether there is a need for risk algorithms tailored to account for specific diseases. For clinical use the latter would only have a role if decisions on prevention strategies were altered by using a more specific than a more generic risk prediction tool. Our post hoc analysis of the discrimination performance of risk prediction models using disease-specific age and sex coefficients supports the importance of having more tailored risk algorithms.

A more nuanced application of age and sex in the clinical setting that takes account of their heterogeneous associations with different CVDs is provided by the following example: A 69-year-old woman with untreated hypertension has a 20% 10-year general risk of CVD, fulfilling guideline criteria for primary prevention. With heart failure her most likely initial CVD presentation within that 10-year time frame (see Figure 1), a tailored blood pressure–lowering regime that excludes calcium antagonists would optimize CVD prevention because these drugs are relatively less effective at reducing risk of heart failure.43 At earlier ages, where CHD is the more common initial presentation, the choice of blood pressure–lowering medication is likely to make little difference to outcomes. This is just 1 example of the way in which understanding of the heterogeneity of risks associated with specific end points could lead to more personalized risk modification.

We also provide further evidence of the need to protect women against CVD with the same vigor as for men. The
current strategy of evaluating and treating short-term risk of total CVD has the consequence that almost all men aged >70 should be on treatment, irrespective of their CVD risk factors. However, a wider group of people with high risk of specific CVDs could be targeted and treated earlier by increasing the sensitivity (by extending the time horizon to lifetime, as suggested by the Joint British Societies latest recommendations) and specificity (by using more specific diagnoses) of risk predictions. Given that the majority of initial CVD presentations in our cohort were nonfatal (84% in men and 80% in women), such opportunities for earlier intervention via refinement of prediction tools should not be missed.

Furthermore, our findings have potentially important consequences for the accuracy of models used to predict CVD risk in clinical practice. We found large differences in the associations of different CVDs with age (from very weak with SAH, heart failure, stroke, and AAA) and male sex (from negative with SAH to very strong with AAA, nonfatal MI, and UCD). So far, most efforts to improve the prediction of CVD have focused on refining current models with new predictors. Although there are several models for specific CVDs (eg, heart failure, stroke), current guidelines recommend assessment of total CVD risk to simplify clinical decision making. Here we show that this one-size-fits-all approach reduces the ability to discriminate between individuals with high and low risk of specific CVDs.

**Implications for Research**

Our findings suggest that future research on the primary prevention of CVD should take account of current patterns of disease presentation and redress the imbalance of previous literature that has focused extensively on heart attack and stroke. Our findings have implications for the design and interpretation of observational studies, randomized trials, and meta-analyses investigating the primary prevention of CVDs. Because the fundamental risk factors of age and sex have such heterogeneous associations with different CVDs, and most studies are only sufficiently powered to examine CVD aggregates, it is important to account for the relative proportion of each disease in the composite end point in meta-analysis. Despite an extensive literature on the underlying biological and behavioral pathways by which sex may influence aggregates of CVD and CHD, there is a lack of mechanism studies that investigate why sex has such heterogeneous associations on different CVDs.

**Limitations**

Our study has important limitations. First, we were not able to resolve some disease subtypes, eg, systolic versus diastolic heart failure or ruptured versus nonruptured cases of AAA. We did find that the association of MI with male sex was more marked for ST-segment–elevation MI than non–ST-segment–elevation MI, suggesting an even greater degree of heterogeneity may be unmasked by investigating more specific diagnoses. Second, we did not evaluate common CVDs that are commonly asymptomatic such as atrial fibrillation. Third, EHRs contain limited covariates for explaining the heterogeneity in sex differences that we report. Fourth, there were 2 less well-specified diagnoses (CHD NOS and stroke NOS) which we were unable to resolve further, but which we included to ensure all potential initial presentations were taken into account. Stroke NOS is likely to be largely ischemic stroke, based on proportion of strokes that are ischemic and the behavior of this end point in modeling, indicating that we may have overestimated the association of ischemic stroke with male sex. We believe CHD NOS is a mixture of stable and unstable angina given the associations in this article and others, but are unable to substantiate this.

**Conclusion**

In an era of modern primary prevention, CVDs commonly first present with heart failure, transient ischemic attack, stable angina, and PAD—diseases that have seldom been the focus of primary prevention studies. Predicting CVD risk should take account a wide range of CVDs, and the different association each has with age and sex, as well.

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**Disclosures**

None.

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**CLINICAL PERSPECTIVES**

The first lifetime presentation of cardiovascular disease in men and women in the 21st century is not currently well understood, with contemporary studies of sufficient size and clinical resolution to distinguish the most common cardiovascular diseases (CVDs) lacking. Traditional cohort studies have, to date, commonly focused on incident heart attack and stroke, but it is well recognized that both have been rapidly declining in incidence. Understanding how CVDs first present is important for developing primary prevention strategies that protect against specific phenotypes and against the wider cascade of other CVDs that often follow. Electronic health records based on usual clinical practice in unselected, contemporary populations provide an important opportunity to assess how CVD first presents in women and men across a wide range of 12 different diseases affecting the head, heart, abdominal, and peripheral circulations. In a study of 1.9 million adults, 114,859 people experienced an incident cardiovascular diagnosis, the majority (66%) of which were neither myocardial infarction nor ischemic stroke. Sex has differing associations with different CVDs, with implications for risk prediction and management strategies. Chronic disease, such as heart failure and peripheral arterial disease, account for a substantial proportion of initial lifetime CVD presentations, yet are been excluded from many risk prediction algorithms. Given the recent decline in the incidence of acute events, our findings emphasize the relevance of risk algorithms that take account of the current burden of CVDs.

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