Morning plasma cortisol as a cardiovascular risk factor: findings from prospective cohort and
Mendelian randomization studies

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Abstract

Objective: The identification of new causal risk factors has the potential to improve cardiovascular disease (CVD) risk prediction and the development of new treatments to reduce CVD deaths. In the general population, we sought to determine whether cortisol is a causal risk factor for CVD and coronary heart disease (CHD).

Design and Methods: Three approaches were adopted to investigate the association between cortisol and CVD/CHD. First, we used multivariable regression in two prospective nested case-control studies (total 798 participants, 313 incident CVD/CHD with complete data). Second, a random-effects meta-analysis of these data and previously published prospective associations was performed (total 6680 controls, 696 incident CVD/CHD). Finally, one- and two-sample Mendelian randomization analyses were performed (122,737 CHD cases, 547,261 controls for two-sample analyses).

Results: In the two prospective nested case-control studies, logistic regression adjusting for sex, age, body mass index, smoking and time of sampling, demonstrated a positive association between morning plasma cortisol and incident CVD (OR 1.28 per 1 SD higher cortisol, 95% CI 1.06-1.54). In the meta-analysis of prospective studies the equivalent result was OR 1.18, 95% CI 1.06-1.31. Results from the two-sample Mendelian randomization were consistent with these positive associations: OR 1.06, 95% CI 0.98-1.15.

Conclusions: All three approaches demonstrated a positive association between morning plasma cortisol and incident CVD. Together these findings suggest that elevated morning cortisol is a causal risk factor for CVD. The current data suggest strategies targeted at lowering cortisol action should be evaluated for their effects on CVD.
Introduction

Despite lifestyle improvements and the successful targeting of cardiovascular disease (CVD) risk factors, such as hypercholesterolemia and hypertension, CVD is a leading cause of death in high income countries and increasingly so in low and middle income countries. The identification of new causal risk factors, such as elevated morning plasma cortisol, has the potential to improve CVD risk prediction and the development of new treatments to reduce CVD deaths.

Cortisol is a glucocorticoid produced by the adrenal glands, which is responsible for cardiovascular and metabolic adaptations during stress. Plasma cortisol has a circadian rhythm with a peak on waking and then declining throughout the day. Inappropriately sustained cortisol production in patients with tumours causing Cushing’s syndrome results in abdominal obesity, hyperglycemia, hypertension, and dyslipidemia and is associated with a four-fold increase in mortality rate, predominantly due to accelerated atherosclerosis and increased CVD. A similar higher rate of CVD has been documented in patients prescribed exogenous glucocorticoid therapy at supra-physiological doses.

Several population-based cross-sectional studies have reported morning plasma levels of cortisol to be positively associated with plasma glucose, blood pressure, and other cardiovascular risk factors. Associations have also been found between cortisol and markers of subclinical atherosclerosis including carotid plaques and coronary artery calcification. Evidence of an association between circulating cortisol levels and CVD events is, however, less conclusive, with studies to date mostly being cross-sectional. Three prospective cohort studies have reported a positive association between morning plasma cortisol and CVD, but effects were imprecisely estimated due to the small study sizes (number of incident CVD cases between 63 and 320). Larger prospective studies, and the use of genetic predictors of morning plasma cortisol as instrumental variables in Mendelian randomization analyses, are required to address whether cortisol is a causal risk factor for CVD.

A recent genome-wide association meta-analysis (GWAMA) study identified independent variants in one locus on chromosome 14 which together explained 0.5% of the variance in morning plasma
cortisol in European populations\textsuperscript{17}. These findings provide instrumental variables for a Mendelian randomization analysis of the effect of morning plasma cortisol on CVD\textsuperscript{16}.

We hypothesised that elevated morning plasma cortisol is a causative risk factor for incident CVD and tested this in multiple prospective studies and using Mendelian randomization.
Methods

New prospective cohort studies of plasma cortisol and cardiovascular disease

The Northern Sweden VIP, MONICA and MSP study

A prospective case-control study of fatal and non-fatal myocardial infarction (MI) and stroke was nested within the combined data from three studies conducted in the same population (Northern Sweden) using identical study protocols (the Västerbotten Intervention Program (VIP), the Northern Sweden MONItoring Of trends and Determinants in CArdiovascular Disease (MONICA) survey and the Mammary Screening Program (MSP); details are provided in supplementary material)19, 20. Incident cases were selected after removal of all prevalent cases, and two controls for every case were matched to cases by sex, age (median difference in age 0.33 years), study and date of baseline assessment (median difference in baseline assessment date 79 days). The study was approved by the Research Ethics Committee of Umeå University and informed consent was obtained from all participants.

Cortisol was measured by immunoassay on plasma samples obtained at the baseline assessment after at least 4 hours of fasting and stored at -80°C 21. Genotyping of three single nucleotide polymorphisms (SNPs) from the GWAMA which together contributed most to morning plasma cortisol (rs12589136, rs2749527, rs11621961) 17 was performed on the OpenArray platform.

British Women’s Heart and Health Study (BWHHS)

BWHHS is a prospective cohort study of 4,286 women who were randomly selected from 23 British towns between 1999 and 2001 and who were aged 60-79 years at baseline; details are provided in supplementary material 22. We used participants from a prospective case-control study of coronary heart disease (CHD) nested within this cohort, as previously described 23. Incident cases of CHD were selected, and two controls were randomly selected, within 5-year age groups of the cases, from women without CHD at the baseline assessment and who had been followed-up over the same time as the cases without a CHD event.
Serum cortisol levels were determined by radio-immunoassay (MP Biomedicals, UK; intra-assay CV 5.1-7.0%, inter-assay CV 6.0-7.9%) on blood samples obtained at the baseline examination after a minimum six-hour fast between 0800h and 1700h. The BWHHS has Local- and Multi-Centre Research Ethics Committee (LREC and MREC) approvals and all participants provided written informed consent.

Statistical Analysis

In the VIP, MONICA and MSP cohort, conditional logistic regression models, adjusting for matching variables (age and baseline assessment date, Sex and study did not vary between matched cases and controls), smoking, body mass index (BMI) and time of sampling (continuous variable as number of minutes since the earliest sampling time) were used to investigate the effect of a 1 SD increase in morning plasma cortisol on CVD. The controls in BWHHS were matched to cases within five-year age bands meaning control for the matching factor can be obtained, with no loss of validity and possible increase in precision, using (unconditional) logistic regression as there are no problems of sparse data. In BWHHS, logistic regression models, adjusting for the matching variable (age), smoking, BMI and time of sampling were used to investigate the effect of a 1 SD increase in morning plasma cortisol on CHD.

The main analysis used samples collected before 1100h, adjusted for time of sampling, as used for the GWAMA which identified genetic variants associated with morning plasma cortisol. The impact of missing data was investigated using multiple imputation by chained equations; details of this procedure and all additional analyses are in the supplementary material. Analyses were performed using Stata v.14. We report exact two-sided p-values throughout.

Meta-analysis of prospective studies

The multivariable regression meta-analysis of prospective studies combined data from the unpublished analyses of VIP, MONICA and MSP (fatal and non-fatal CVD), and BWHHS (fatal and non-fatal CHD) described above, as well as published data for morning plasma cortisol from the Caerphilly Study (fatal and non-fatal ischemic heart disease (IHD)) and the Vietnam Experience.
Study (fatal CVD). Details of these cohorts are available in the supplementary material. We were not able to include a third published prospective study because the necessary data were not available.

Statistical Analysis

A random-effects (DerSimonian-Laird) method was used to pool the maximally confounder adjusted (for each study) association of 1 SD higher morning plasma cortisol with CVD/CHD. A random-effect method was used as we were aware of clinical/study heterogeneity (i.e. different confounder adjustment, outcome definitions and genders/geographical locations), and statistical evidence of heterogeneity between studies was assessed with chi-squared goodness of fit and the $I^2$ statistic, together with its 95% confidence interval. Sensitivity analyses are described in the supplementary material.

One- and two-sample Mendelian randomization analyses

Mendelian randomization analyses use genetic variants as a proxy for environmental exposures. As genetic variants are assigned at conception and tend not to be associated with other behavioural, social, or physiological factors, these analyses are less likely to be affected by reverse causation or confounding than conventional observational studies. The genetic instrument for morning plasma cortisol comprised of three SNPs (rs12589136, rs2749527, rs11621961) in low linkage disequilibrium ($r^2 < 0.3$), each associated independently with morning plasma cortisol in the CORtisol NETwork (CORNET) GWAMA ($n=12,597$). Biochemical investigation of these SNPs have also shown them to be associated with variation in total cortisol binding activity in plasma. The three SNPs explain approximately 0.5% variation in morning plasma cortisol levels and are located on chromosome 14 at a locus containing the SEPRINA6 gene. The SERPINA6 gene codes for corticosteroid-binding globulin (CBG) which binds to cortisol in the blood. To assess the potential of horizontal pleiotropy, where a genetic variant influences multiple outcomes through independent pathways, we investigated the association of these SNPs with confounders included in the multivariable prospective analysis in
the VIP, MONICA and MSP cohort, and with cardiovascular risk factors in publicly available GWAS consortium.

One-sample Mendelian randomization analysis in VIP, MONICA and MSP cohort

The causal estimate was derived using the two-stage method comprising a first-stage regression of the exposure on the SNP, and a second-stage regression of the outcome on the fitted values of the exposure from the first stage. Further details are available in the supplementary material. This provides the log OR for CVD per 1 SD increase in morning plasma cortisol due to each SNP. The effects of each SNP were combined in a random-effects meta-analysis.

Two-sample Mendelian randomization analysis

A two-sample Mendelian randomization approach, using publicly available data, was used to estimate the causal effect of morning plasma cortisol on CHD. We investigated the association between the genetic instrument for morning plasma cortisol and risk of CHD in a meta-analysis of 34,541 CHD cases and 261,984 controls of UK Biobank and 88,192 cases and 162,544 controls from CARDIoGRAMplusC4D. In the main analyses the causal effect of morning plasma cortisol on CHD was estimated for each SNP individually using the Wald ratio method and then combined in a random-effects meta-analysis. Additional analyses described in the supplementary material were performed to investigate the robustness of this causal estimate and any potential pleiotropic effects. The summary GWAS data for morning plasma cortisol and CHD can be downloaded from https://datashare.is.ed.ac.uk/handle/10283/2787 and https://data.mendeley.com/datasets/gbbsrpx6bs/1 respectively.
Results

New prospective cohort studies of plasma cortisol and cardiovascular disease

The Northern Sweden VIP, MONICA and MSP study

There were 905 cases and 1717 matched controls (93 cases with one control, 812 cases with two controls) included in the analysis. Of the 905 cases of CVD, there were 536 cases of MI and 369 cases of stroke, including fatal and non-fatal events. Median time to CVD event was 3 years 9 months.

Those with CHD were more likely to have hypertension, and higher BMI, total cholesterol, triglycerides, and fasting glucose, and to smoke tobacco (Table 1). Owing to the high proportion with missing data (Table S4), analyses were performed for all participants, those with complete data, and in a dataset with imputation of missing data. Time of blood sampling was not recorded in 52% of participants, who were more likely to be male, not fasted, and have lower BMI, leptin and fasting glucose levels (Table S5). The variables used in the imputation models are presented in Table S6 and distributions of imputed variables in Table S7.

There was no association between plasma cortisol and CVD incidence in the unadjusted model of all participants (Table 2). However, there was a positive association between morning plasma cortisol and CVD when including only participants with cortisol assessed between 0700h and 1100h (cases=297, controls=445), which was strengthened when adjusting for potential confounders (cases=268, controls=396) (Table 2). When the analysis was restricted to participants with complete data there was evidence of a positive association in all models (Table S1). The effect that each confounder has on the effect estimate can be seen in the one-at-a-time analysis with BMI having the largest effect in the observed (Table S2) and imputed (Table S3) data. The estimate obtained from the imputed data supported an association between higher morning plasma cortisol and CVD, but with an attenuated estimate in the adjusted model (OR 1.12, 95% CI 0.99 to 1.26) (Table S8).

When the adjusted analysis was stratified by outcome (MI or stroke) the association between morning plasma cortisol and MI appeared to be stronger (OR 1.85, 95% CI 1.28 to 2.68) than with stroke (OR
1.08, 95%CI 0.84 to 1.39); p=0.01 for difference between the two estimates. There was no strong
statistical evidence to suggest the association differed by sex (p = 0.36).

*British Women’s Heart and Health Study (BWHHS)*

There were 169 cases (111 non-fatal and 58 fatal) and 363 controls included in the analysis of the
BWHHS nested case-control study. Those with CHD had higher triglyceride levels and systolic blood
pressure, and lower HDL cholesterol levels; they were more likely to have hypertension and to have
ever smoked (Table 1). The pattern of associations was similar to that in the VIP, MONICA and MSP
studies, with little evidence of an association in unadjusted analyses of the whole study population
and a positive association emerging as analyses were restricted to those with a sample taken between
0700h and 1100h and after adjustment for confounders (Table 2). Estimates in BWHHS were less
precise than in VIP, MONICA and MSP with 95% confidence intervals including the null value.

**Meta-analysis of prospective studies**

By combining data from the previously published analyses of the Caerphilly study and Vietnam
Experience study with data from the VIP, MONICA and MSP cohort and BWHHS cohort, a total of
7376 individuals (696 CVD events (fatal and non-fatal), 6680 controls) were included. The estimate
from the Vietnam Experience study was from an unadjusted regression model as that was all that was
available. The other estimates were from models adjusting for potential confounders (Table S9). A 1
SD higher cortisol was associated with 18% higher odds of experiencing a CVD event (OR 1.18, 95%
CI 1.06 to 1.31) (Figure 1). There was no strong evidence of heterogeneity ($\chi^2 = 3.14, p = 0.37, I^2 = 5\%$ (95% CI 0% to 86%), and the positive association remained in a leave-one-out analysis (Table
S10). Though power for these analyses is low, we found no strong evidence of association between
study effect size estimate and mean participant age (p=0.64) or percentage male (p=0.93).

Visual inspection of a funnel plot of the effect size (lnOR) against the standard error of the effect size
(lnOR) suggested asymmetry (Figure S1), indicating that some small studies that were consistent with
a small or null association between cortisol and risk of CVD may be missing. Egger’s test did not
provide evidence of small study bias (p=0.32), although this test has low statistical power given the low number of studies included.

One- and two-sample Mendelian randomization analyses

In the VIP, MONICA and MSP cohort, there was no strong evidence of an association between the SNPs comprising the instrumental variable with any of the matching variables (age, sex, survey date, cohort) or potential confounders (BMI, sampling time, smoking) either individually or when combined in a fixed-effects or random-effects meta-analysis (Table S11). In publicly available GWAS consortia the instrumental variable was positively associated with type 2 diabetes and negatively associated with being overweight, LDL cholesterol and cigarettes smoked per day (Table S12). The Mendelian randomization analysis in the VIP, MONICA and MSP cohort suggested a positive effect of morning plasma cortisol on CVD risk (OR 1.33, 95% CI 0.26 to 6.73), but this had a wide 95% confidence interval given the small sample size.

Using publicly available data, two-sample Mendelian randomization analysis provided evidence that each SD increase in morning plasma cortisol increased CHD: OR 1.06, 95% CI: 0.98 to 1.15 (Figure 2). There was no strong statistical evidence of heterogeneity between the instrumental variable effect estimate for each SNP (Q statistic = 0.88, p=0.64). Estimates obtained from the additional Mendelian randomization analyses, including multivariable Mendelian randomisation, lacked precision but consistent with the positive association obtained in the main analyses (Table S13 and S14).
Previous prospective studies have reported positive associations between morning plasma cortisol and later CVD risk but a lack of precision of the effect estimate cast doubt on the importance of cortisol\textsuperscript{13,15}. Moreover, these reports focused on associations with CVD of the ratio of cortisol to testosterone\textsuperscript{13}, estradiol\textsuperscript{14}, and dehydroepiandrosterone sulfate\textsuperscript{15} which are of questionable biological relevance. Here, we focused on morning plasma cortisol and were able to obtain precise estimates of its association with incident CVD/CHD by undertaking analyses in two new (previously unpublished) prospective nested case-control studies and pooling those results with existing published prospective studies in a random-effects meta-analysis. Following adjustment for key confounders (age, sex, BMI, smoking) where possible we found that a 1 SD higher morning plasma cortisol was associated with 18% higher odds of incident CVD/CHD. Furthermore, evidence from the Mendelian randomization analyses was consistent with these multivariable results, supporting a positive causal effect of morning cortisol levels on CVD/CHD. Given the key sources of biases in these two approaches are different – residual confounding in the multivariable regression analyses and horizontal pleiotropy, where a genetic variant influences multiple outcomes through independent pathways, in the Mendelian randomization analyses – the consistent finding with each strengthens the conclusion that there is a causal effect\textsuperscript{31}. Thus, our findings support the development of interventions that effectively lower cortisol as a means of reducing risk of CVD\textsuperscript{32}.

Plasma cortisol has a circadian rhythm with a peak on waking. In earlier studies, we found that several metabolic cardiovascular risk factors, such as blood glucose, and blood pressure, were more closely associated with morning plasma cortisol than with other measurements such as 24-hour urinary cortisol metabolites\textsuperscript{33}, and that these associations are strengthened by adjusting for the time of sampling, at least for samples obtained in the morning\textsuperscript{34}. We therefore used morning samples, adjusted for time of sampling, in our previous GWAMA which identified genetic variants associated with morning plasma cortisol\textsuperscript{17}. Here, in multivariable regression analyses in both of the two independent studies a positive association of cortisol with CVD/CHD was only observed when we excluded participants with samples outside of the 0700 to 1100hrs time range. This may be because...
afternoon cortisol values have less inter-individual variability, or because morning cortisol better reflects the reactivity of the hypothalamic-pituitary-adrenal axis or contributes more to the tissue actions of cortisol.

There is also some consistency of findings between cortisol measured in other sample types. The cumulative exposure to cortisol over many weeks or months has been assessed by measuring cortisol in hair, which has also been reported recently to be associated with self-reported history of CVD in a retrospective cohort study, with a comparable magnitude of effect as traditional risk factors (OR 2.7 for highest cortisol quartile compared to lowest quartile, compared with hypertension (OR 2.5), abdominal obesity (OR 2.2), and diabetes mellitus (OR 3.1)) \(^{35,36}\). Flatter slopes in decline of saliva cortisol across the day in the prospective Whitehall II cohort study among 4047 civil servants were associated with higher risk of cardiovascular deaths (hazard ratio for 1 SD reduction in slope steepness = 1.87; 95% CI 1.32 to 2.64) \(^{37}\). Also, higher twenty-four-hour urinary cortisol levels, as measured in 861 participants who were part of the prospective InCHIANTI study, were associated with higher risk of dying of CVD (highest tertile compared with lowest tertile, hazard ratio = 5.00; 95% CI 2.02 to 12.37) \(^{38}\). Of course, we recognise the potential for publication bias in this area so adding genetic evidence to our findings was deemed important.

Mendelian randomization analysis is less likely to be biased by confounding or reverse causation, and suggests that the elevated morning plasma cortisol that is associated with CVD/CHD is unlikely to be attributed to, for example, a response of the hypothalamic-pituitary-adrenal axis to chronic inflammation of subclinical cardiovascular disease. There was no strong evidence of an association between the instrumental variable used in the Mendelian randomization analyses and the potential confounders included in the multivariable regression analyses. In addition, estimates from the multivariable Mendelian randomization analyses were consistent with the positive association obtained in the main analyses. The three SNPs comprising the instrumental variable were not in LD \((r^2 < 0.1)\) with a SNP (rs112635299) associated with CHD in the same locus \(^{29}\). This suggests it is unlikely that horizontal pleiotropy is biasing the estimate between morning plasma cortisol and CHD observed in our two-sample Mendelian randomization analysis. Further, consistent estimates were
obtained using the Wald ratio, inverse variance weighted, likelihood method and weighted median approaches. The genetic variants comprising the instrument in the Mendelian randomization analyses explain 0.5% variance in morning plasma cortisol. This may cause weak instrument bias which in the two-sample analyses will bias the estimate towards the null.

Greater clarity in understanding the causal pathway between elevated cortisol and CHD will be achieved by: firstly, identifying additional variants associated with cortisol to strengthen the instrumental variable used in Mendelian randomization and enable use of additional methods more adept at handling pleiotropy; and secondly, validating the instrumental variable for plasma cortisol in non-European populations justifying the analysis of larger outcome datasets of mixed ancestry leading to greater precision of causal estimates.

Cortisol acts through intracellular glucocorticoid and mineralocorticoid receptors to modulate fuel metabolism, cardiovascular homeostasis and inflammation. As such there are multiple pathways through which cortisol may be increasing CVD risk. The effects of cortisol on CVD may be mediated by high blood pressure and glucose levels. If this is the case, interventions that target the source of the problem, cortisol, may prevent use of multiple, subsequent interventions (i.e. blood pressure lowering and glucose lowering medications). However, there is also evidence that cortisol is exerting a direct effect on CVD risk, independent of established CVD risk factors. In this case, current treatments will not be effective in reducing CVD risk in individuals with high cortisol levels.

Our stratified analysis suggested there may be a stronger association of morning plasma cortisol with CHD than stroke; although this was not conclusive it is consistent with the higher risk of CHD, rather than stroke, observed in patients exposed to exogenous glucocorticoids. The underlying mechanism for this difference is unknown.

The strengths of our studies include the meta-analysis of multiple cohorts, reducing sampling bias and increasing statistical power, and the careful case classification and use of matched controls in the VIP, MONICA and MSP cohort. A further strength is the consistency between different approaches, with
the positive prospective association supported by findings from the Mendelian randomization analyses

Limitations include the wide range of sampling times and large number of missing data for sampling time in the VIP, MONICA and MSP cohort, which reduced precision and potentially introduced bias when estimating the association between cortisol and CVD. To address these limitations, we performed multivariable imputation to replace missing sampling times with values imputed from the observed data. When the estimate from the imputed dataset, rather than the observed data, was used to contribute to the meta-analysis then the positive association remained (OR 1.12, 95% CI 1.03 to 1.22).

Given the known importance of sampling time for plasma cortisol it was not unexpected that there was no association between cortisol and CVD in the unadjusted analysis. The other cardiovascular risk factors that were measured and adjusted for in the studies included in the meta-analysis was not comprehensive so that the adjustments may have been incomplete. The Mendelian randomization within the VIP, MONICA and MSP cohort, which aimed to establish three-way associations between genotype, plasma cortisol and CVD, was under-powered.

Improving our understanding of the underlying mechanisms of CVD, as well as identifying biomarkers, is of public health importance as it can lead to the development of novel pharmaceutical treatments and guide targeted interventions for those most at risk. The current data suggest that strategies targeted at lowering cortisol action may reduce risk of CVD. Reducing cortisol action safely is challenging, given that the anti-inflammatory properties of cortisol are essential to survival during, for example, severe infection. However, approaches which target tissue-specific cortisol action, such as inhibition of the cortisol regenerating enzyme 11βHSD1, have shown modest improvements in glycaemic control, lipid profile and blood pressure in phase II clinical trials for type 2 diabetes and arguably should be evaluated for their effects on CVD.
Declaration of interest

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### VIP, MONICA and MSP

<table>
<thead>
<tr>
<th></th>
<th>Cases of incident CHD (n=905)</th>
<th>Controls (n=1717)</th>
<th>p</th>
<th>Cases of incident CHD (n=169)</th>
<th>Controls (n=327)</th>
<th>p</th>
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<tbody>
<tr>
<td>Cortisol, nmol/l (SD)</td>
<td>524 (193)</td>
<td>527 (183)</td>
<td>0.85</td>
<td>526 (222)</td>
<td>530 (207)</td>
<td>0.58</td>
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<td>Male, n (%)</td>
<td>655 (72)</td>
<td>1236 (72)</td>
<td>-</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>-</td>
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<tr>
<td>Age, years (SD)</td>
<td>54.5 (7.8)</td>
<td>54.3 (7.9)</td>
<td>0.11</td>
<td>70.4 (5.5)</td>
<td>70.1 (5.3)</td>
<td>0.47</td>
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<tr>
<td>Body Mass Index, kg/m² (SD)</td>
<td>27.0 (3.9)</td>
<td>25.9 (3.6)</td>
<td>&lt;0.0001</td>
<td>28.1 (4.9)</td>
<td>27.4 (4.5)</td>
<td>0.14</td>
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<td>Fasting glucose, mmol/l (SD)</td>
<td>5.63 (1.70)</td>
<td>5.36 (1.12)</td>
<td>&lt;0.0001</td>
<td>6.29 (2.17)</td>
<td>6.20 (1.99)</td>
<td>0.65</td>
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<td>Total Cholesterol, mmol/l (SD)</td>
<td>6.54 (1.30)</td>
<td>6.17 (1.25)</td>
<td>&lt;0.0001</td>
<td>6.76 (1.41)</td>
<td>6.63 (1.13)</td>
<td>0.27</td>
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<td>HDL Cholesterol, mmol/l (SD)</td>
<td>1.25 (0.98)</td>
<td>1.37 (0.92)</td>
<td>0.18</td>
<td>1.53 (0.42)</td>
<td>1.68 (0.47)</td>
<td>0.0003</td>
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<td>Triglycerides, mmol/l (SD)</td>
<td>1.89 (1.27)</td>
<td>1.54 (0.87)</td>
<td>&lt;0.0001</td>
<td>2.10 (1.02)</td>
<td>1.87 (1.00)</td>
<td>0.005</td>
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<td>Systolic blood pressure, mmHg (SD)</td>
<td>142 (19)</td>
<td>135 (17)</td>
<td>&lt;0.0001</td>
<td>153 (22)</td>
<td>148 (27)</td>
<td>0.05</td>
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<tr>
<td>Diastolic blood pressure, mmHg (SD)</td>
<td>89 (10)</td>
<td>85 (9)</td>
<td>&lt;0.0001</td>
<td>81 (11)</td>
<td>80 (12)</td>
<td>0.52</td>
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<td>Hypertensive, n (%)</td>
<td>574 (72)</td>
<td>752 (50)</td>
<td>&lt;0.0001</td>
<td>136 (80)</td>
<td>232 (71)</td>
<td>0.02</td>
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<td>Ever smoked, n (%)</td>
<td>282 (33)</td>
<td>337 (20)</td>
<td>&lt;0.0001</td>
<td>90 (53)</td>
<td>138 (42)</td>
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</tbody>
</table>

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Table 1. Characteristics of cases and controls in the combined VIP, MONICA and MSP cohort and BWHHS cohort. SD, Standard deviation; HDL, High Density Lipoprotein; p, p value. Means and standard deviations are presented for continuous variables. The number and percentage of individuals are presented for binary variables. Hypertensive was generated as blood pressure at examination >= 160/90 mmHg and/or taking antihypertensive medication. The p value from VIP, MONICA and MSP is from a conditional logistic regression model accounting for the matched nature of the data.
<table>
<thead>
<tr>
<th></th>
<th>VIP, MONICA and MSP</th>
<th>BWHHS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cases</td>
<td>Controls</td>
</tr>
<tr>
<td>Unadjusted</td>
<td>905</td>
<td>1717</td>
</tr>
<tr>
<td>Adjusted</td>
<td>323</td>
<td>492</td>
</tr>
<tr>
<td>Unadjusted and 0700h-1100h</td>
<td>297</td>
<td>445</td>
</tr>
<tr>
<td>Adjusted and 0700h-1100h</td>
<td>268</td>
<td>396</td>
</tr>
</tbody>
</table>
Table 2. Associations of plasma cortisol with CVD in the combined VIP, MONICA and MSP cohort and BWHHS cohort. OR, odds ratio; CI, confidence intervals; p, p value. All models included the matching variables of age; for VIP, MONICA and MSP, models also included the matching variables of sex, study, and baseline assessment date. Adjusted models also included BMI, smoking and time of sampling in the regression models. Analyses were repeated restricted to individuals with blood sampling time between 0700h and 1100h.
New analyses
VIP, MONICA and MSP (CVD - fatal and non-fatal)
BWHHS (CHD - fatal and non-fatal)
Subtotal (I-squared = 0.0%, p = 0.554)

Previously published analyses
Caerphilly (IHD - fatal and non-fatal)
Vietnam (CVD - fatal)
Subtotal (I-squared = 37.5%, p = 0.206)

Overall (I-squared = 4.6%, p = 0.370)

NOTE: Weights are from random effects analysis
Figure 1. Meta-analysis of prospective multivariable regression association of morning plasma cortisol with cardiovascular disease. Meta-analysis provides evidence that a 1 SD increase of morning plasma cortisol is associated with 18% higher risk of later cardiovascular disease (Odds ratio (OR) 1.18 95% CI 1.06 to 1.31, p=0.002). The VIP, MONICA and MSP study consisted of 268 cases and 396 controls; the BWHHS consisted of 45 cases and 89 controls; the Caerphilly study consisted of 320 cases and 2,003 controls; the Vietnam study consisted of 63 cases and 4,192 controls. The number of controls in the Caerphilly and Vietnam Experience studies was assumed to be all individuals who did not experience a CVD event or CVD death. An assumption was made that the standard deviation of cortisol in the Vietnam Experience study was the same as the VIP, MONICA and MSP study; this allowed inclusion of the Vietnam Experience study. Bars represent individual study 95% confidence intervals, with a central block proportional to study size. The summary diamond represents the pooled effect size estimate and 95% CI. Weights are from random-effects analysis. a regression model included age, survey date, cohort, smoking, BMI and sampling time. b regression model included age, smoking, BMI and sampling time. c regression model including age, smoking status, sampling time, adult social class, alcohol consumption, height, FEV₁/height², fibrinogen (log transformed), white cell count (log transformed). d estimate from unadjusted regression model.
### Study Results

<table>
<thead>
<tr>
<th>Study</th>
<th>OR (95% CI)</th>
<th>% Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>UK Biobank and CARDioGRAMplusC4D</td>
<td></td>
<td></td>
</tr>
<tr>
<td>rs11621961</td>
<td>1.12 (0.96, 1.31)</td>
<td>26.94</td>
</tr>
<tr>
<td>rs12589136</td>
<td>1.02 (0.90, 1.16)</td>
<td>39.37</td>
</tr>
<tr>
<td>rs2749527</td>
<td>1.07 (0.94, 1.23)</td>
<td>33.70</td>
</tr>
<tr>
<td>Overall (I-squared = 0.0%, p = 0.643)</td>
<td>1.06 (0.98, 1.15)</td>
<td>100.00</td>
</tr>
</tbody>
</table>

**NOTE:** Weights are from random effects analysis.
Figure 2. Effect estimate of genetically elevated morning plasma cortisol on risk of CHD. The forest plot shows odds ratios and 95% confidence intervals for CHD per standard deviation increase in morning plasma cortisol. The overall genetic effect was estimated using random-effects meta-analysis of Wald ratio estimates. The genetic effect was estimated using data from the meta-analysis of 34,541 CHD cases and 261,984 controls from UK Biobank and 88,192 cases and 162,544 controls from CARDIoGRAMplusC4D 29.