The co-existence of elevated high sensitivity C-reactive protein and homocysteine levels is associated with increased risk of metabolic syndrome: A 6-year follow-up study

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Abstract

Accumulating evidence has revealed that both high sensitivity C-reactive protein (hsCRP) and homocysteine (HCY) are associated with increased risk of metabolic syndrome (MetS) and cardiovascular disease. However, it is unclear whether the coexistence of these conditions accelerates the risk of metabolic syndrome (MetS). We hypothesized that the combination of high sensitivity C-reactive protein (hsCRP) and homocysteine (HCY) levels could exacerbate the development of MetS in a large prospective cohort study. We selected data from 3,170 individuals (1,614 men and 1,556 women) who participated in the Korean Genome and Epidemiology Study. Participants with high hsCRP and HCY levels were categorized into quartiles. MetS was defined based on the criteria of the modified National Cholesterol Education Program, Adult Treatment Panel III. The prevalence of MetS was higher in participants with concurrent high hsCRP and HCY compared to those with low hsCRP and HCY levels. The incidence of MetS at the 6-year follow-up was the highest in participants with concomitant high hsCRP and HCY levels, regardless of obesity. Even after adjusting for potential confounding factors including body mass index in a multivariate logistic regression model, subjects with elevated hsCRP and HCY levels had a 2.50-fold increased risk of developing MetS at the six-year follow-up compared to those who did not have high hsCRP and HCY level. MetS is more prevalent in the concurrent presence of elevated hsCRP and HCY levels. The combination of the two conditions may contribute to an increased risk of MetS, but these factors may not be synergistic.
Introduction

Metabolic syndrome (MetS) is characterized by a cluster of predisposing factors including central obesity, insulin resistance, dyslipidemia, and hypertension (HTN), which directly increase the risk of cardiovascular diseases (CVD). These relationships have been confirmed in several longitudinal studies [1]. These components of MetS are interconnected with various risk factors such as age, genetic variations related to inflammation, and various lifestyle factors [2–7]. Among these predisposing factors, visceral obesity is the most prevalent manifestation of MetS [8]. Considering the rapidly increasing prevalence of obesity, the substantial attention to MetS-related comorbidities is expected to increase [9].

C-reactive protein (hsCRP), which is known to be an important biomarker of CVD, is an acute-phase protein that is generated in the liver and is stimulated by pro-inflammatory cytokines including TNF-α and IL-6 [10]. Substantial evidence shows that obese individuals have higher hsCRP level than non-obese individuals [11, 12], indicating that the abdominal adipose tissue may be involved in the production and regulation of hsCRP level [13–15]. A number of studies have also demonstrated that high hsCRP level is associated with increased risk for CVD and MetS [16–18], suggesting that hsCRP could be an important marker for the prediction of cardio-metabolic risk [19, 20]. Homocysteine (HCY) is a non-essential amino acid that is biosynthesized from methionine. In the last few decades, it has been suggested that an elevated HCY level is pathophysiologically involved in the promotion of cardiovascular events and increased risk of MetS [21–23]. Although previous studies reported that elevated hsCRP and HCY levels are both associated with increased risk of MetS, to the best of our knowledge, no study has examined their combined effects on the risk of MetS in a large prospective cohort study. We hypothesized that the presence of elevated HCY and hsCRP levels would accelerate the risk of MetS in a cohort study.

Materials and methods

Subjects

Participants in the present study were part of a larger study, the Korean Genome and Epidemiology Study (KoGES), which is an ongoing, population-based cohort study that started in 2001 under the original title “The Korean Health and Genome Study.” Detailed information on the study design and aims of the KoGES have been previously reported [24]. In brief, the original study was designed to establish a representative adult cohort in the city of Ansan, Korea, and to identify the epidemiologic characteristics, frequency, and determinants of chronic diseases among Koreans. It began in June 2001 with 5,015 participants (2,521 men and 2,494 women; ages, 40–69 years). The participants were followed biennially, with a scheduled visit for a comprehensive health examination and on-site interviews at Korea University Ansan Hospital. Data from the 4th examination, conducted from March 2007 to February 2009 (for baseline assessment), and the 7th examination, conducted between March 2013 and February 2015 (for the 6-year follow-up assessment), were used in the present study. All participants signed an informed consent form, and the study protocol was approved by the Institutional Review Board of Korea University Ansan Hospital (Protocol ID: ED0624). All methods and experiments were performed in accordance with the relevant guidelines and regulations. Among 3,255 participants who were initially screened at baseline, 3,170 were included in the study. All participants underwent anthropometric measurements and a general health examination. Participants were excluded if they had any known systemic inflammatory disease or genetic abnormality and extreme outliers of hsCRP (n = 56, hsCRP > 10 mg/dL) and HCY (n = 29, HCY > 30 μmol/L) [25, 26]. High hsCRP was defined as >1.43 mg/dL for men and >1.20 mg/dL for women, and high HCY was defined as >14.71 μmol/L for men and >11.03 μmol/L for women, which correspond to the
75th percentiles of the whole study cohort. For the purpose of the present study, participants were divided into 4 groups based on the presence of high hsCRP and HCY level. The population was also sub-divided into tertiles of hsCRP and HCY levels for sub-group analysis.

**Anthropometric and biochemical measurements**

All subjects were surveyed for age, sex, marital status, smoking, drinking, and other habits. Blood pressure (BP) was measured in a standardized manner using a mercury sphygmomanometer. Body mass index (BMI) was calculated as weight in kilograms divided by height in meters squared. Waist circumference was measured at the midpoint between the lower rib margin and the iliac crest in a standing position. Blood was drawn for biochemical analysis after overnight fasting. Biochemical data including glucose, total cholesterol, triglycerides, HDL cholesterol, and hsCRP levels were determined using an auto-analyzer (ADVIA 1650 and 1800, Siemens, Tarrytown, NY) at Seoul Clinical Laboratories (Seoul, Korea). HCY level was measured using a chemiluminescence immunoassay assay.

**Definition of MetS**

Data were also collected on factors used to define components of MetS, which was defined based on the Third Report of the National Cholesterol Education Program Expert Panel III on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (NCEP-III) [27]. The components included abdominal obesity, low HDL-cholesterol, hypertension, hypertriglyceridemia, and high fasting glucose. High blood pressure was defined when the systolic blood pressure was 130 mmHg or higher, diastolic blood pressure was 85 mmHg or higher, or the patient was taking antihypertensive medication. Low HDL-cholesterol was defined as lower than 40 mg/dL for males and 50 mg/dL for females. Hypertriglyceridemia was defined as triglycerides of 150 mg/dL or greater. High fasting glucose was defined as a glucose level of at least 100 mg/dL or when a subject was receiving oral hypoglycemic agents or insulin therapy. Abdominal obesity was defined as a waist circumference wider than 90 cm in males and 85 cm in females [28]. The metabolic score was calculated by summing the number of positive diagnostic criteria on the NCEP-III. The incidence of MetS was defined as the percentage of participants who were newly diagnosed with MetS during the 6-year follow-up period.

**Statistical analysis**

Differences in the means for normal variables were examined by one-way ANOVA. The probabilistic distribution for non-normal variables was compared with the Kolmogorov-Smirnov test. Categorical variables were expressed as n (%) and analyzed with a chi-square test. Multivariate logistic regression was applied with adjustment for the factors identified to be significant risk factors for MetS in univariate analyses, which were age, sex, smoking status, alcohol use, baseline BMI, and change in BMI at the six-year follow-up (%BMI). The Bonferroni correction was applied for pairwise comparisons. Adjusted odds ratios were estimated with 95% confidence intervals (CIs), referenced to participants with normal hsCRP and HCY levels. All statistical analyses were performed using IBM SPSS version 23.0 (IBM Corp., Armonk, NY, USA). Statistical significance was identified at the 0.05 significance level.

**Results**

**General characteristics of study participants**

Originally, the KoGES was designed to establish a representative adult cohort and to identify the epidemiologic characteristics and determinants of chronic diseases among Koreans in
2001. Measurement of HCY level in the KoGES participants was included in the study protocol at the 4th examination. Thus, among 3,255 participants who were initially screened at baseline (4th examination), 3,170 were included in the study (Fig 1). Table 1 shows the demographic and biochemical characteristics of participants at baseline, classified into four groups according to the presence or absence of high hsCRP and HCY. Among the 3170 participants at the baseline examination, 435 dropped out over the 6-year follow-up period (non-response rate, 22.3%). There were no significant differences between the respondents and the non-respondents with regard to gender, smoking status, alcohol use, and BMI (p>0.05). However, the mean age was significantly higher in non-respondents than in respondents (p<0.05). Among participants with low hsCRP, the average HCY for those in the “low” and “high” groups at baseline was 10.2 and 15.5 μM, respectively; among the participants with high hsCRP, the average HCY for those in the “low” and “high” groups was 10.4 and 15.4 μM, respectively. Systolic and diastolic BP, glucose, triglycerides, total cholesterol, and HDL cholesterol levels were significantly different among the four groups (p<0.01). Moreover, the group with high hsCRP and HCY levels had the highest metabolic score.

Odd ratios for risk of MetS according to the concurrent presence of high hsCRP and Hcy levels

The prevalence and incidence of MetS among the four groups stratified by the presence of high hsCRP and HCY levels are shown in Tables 2 and 3. The prevalence of MetS was significantly different between the groups (hsCRP[−]/HCY[−] vs. hsCRP[−]/HCY[+] vs. hsCRP[+]/HCY[−] vs. hsCRP[+]/HCY[+], 17.9% vs. 27.8% vs. 35.0% vs. 47.2%, p<0.001). Moreover, there was also a significant difference in the incidence of MetS at the 6-year follow-up among
Table 1. General characteristics of participants according to the presence of high HCY and hsCRP levels.

|                      | hsCRP[-] | HCY[+] | hsCRP[-] | HCY[+] | P value |
|----------------------|----------|--------|----------|--------|---------|
| Sample size, n (%)   | 1813 (57.2) | 563 (17.8) | 565 (17.8) | 229 (7.2) | -       |
| Age (years)          | 53.4±6.6 | 56.4±8.3 | 54.8±7.2* | 58.3±8.5* | <0.001  |
| BMI (kg/m2)          | 24.2±2.7 | 24.5±2.8 | 25.5±3.3# | 25.6±2.9* | <0.001  |
| ABMI at follow-up (kg/m2) | 0.14±1.21 | 0.03±1.28 | 0.20±1.15# | -0.09±1.29 | 0.07    |
| Men, n (%)           | 911 (50.2) | 298 (52.9) | 300 (53.1) | 105 (45.9) | 0.2     |
| Current smoker, n (%)| 267 (14.7) | 94 (16.7) | 107 (18.9) | 40 (17.5) | 0.11    |
| Current drinker, n (%)| 926 (51.1) | 288 (51.2) | 284 (50.3) | 106 (46.3) | 0.58    |
| Medication for hypertension, n (%) | 356 (19.6) | 152 (27.0) | 139 (24.6) | 72 (31.4) | <0.001  |
| Medication for diabetes, n (%) | 121 (6.7) | 47 (8.3) | 46 (8.1) | 29 (12.7) | <0.05   |
| Systolic blood pressure (mmHg) | 110.0±13.7 | 113.6±14.5$ | 112.4±14.1# | 116.3±14.3* | <0.001  |
| Diastolic blood pressure (mmHg) | 73.9±9.5 | 75.8±10.3$ | 75.7±9.9# | 76.3±9.7* | <0.001  |
| Fasting glucose (mg/dL) | 97.5±29.2 | 98.5±29.7 | 102.4±33.6# | 104.6±32.3* | <0.001  |
| Total cholesterol (mg/dL) | 198.8±34.3 | 202.3±34.3 | 203.4±36.2# | 206.7±35.9* | <0.001  |
| HDL cholesterol (mg/dL) | 46.0±10.8 | 45.5±11.2 | 42.8±10.0# | 41.2±8.7* | <0.001  |
| Triglycerides (mg/dL) | 130.6±78.8 | 143.1±95.25 | 157.9±102.1# | 176.6±99.3* | <0.001  |
| hCRP at baseline (mg/dL) | 0.54±0.32 | 0.59±0.34d | 2.86±1.8# | 2.83±1.57* | <0.001  |
| (log-transformed)     | (-0.36±0.31) | (-0.32±0.30) | (0.39±0.22) | (0.40±0.21) |        |
| hsCRP at follow-up (mg/dL) | 0.83±1.11 | 1.02±1.22 | 1.55±1.57# | 1.90±1.80* | <0.001  |
| (log-transformed)     | (-0.26±0.36) | (-0.16±0.37) | (0.02±0.38) | (0.11±0.38) |        |
| HCY at baseline (μmol/L) | 10.2±2.11 | 15.5±3.41# | 10.4±2.04 | 15.4±3.38* | <0.001  |
| (log-transformed)     | (1.00±0.09) | (1.18±0.09) | (1.01±0.09) | (1.18±0.09) |        |
| HCY at follow-up (μmol/L) | 12.3±3.14 | 16.0±5.2$ | 12.7±3.0 | 16.8±5.9* | <0.001  |
| (log-transformed)$^*$ | (1.07±0.09) | (1.18±0.12) | (1.09±0.09) | (1.20±0.12) |        |
| Vitamin intake, n (%) | 513 (28.3) | 98 (17.4) | 149 (26.4) | 35 (15.3) | <0.001  |
| Metabolic score       | 1.31±1.15 | 1.55±1.23# | 1.88±1.23# | 2.22±1.27* | <0.001  |
| Metabolic score at follow-up | 1.27±1.08 | 1.55±1.08 | 1.61±1.18# | 1.92±1.24* | <0.001  |

Abbreviation: BMI, body mass index; HDL, high-density lipoprotein; hsCRP, high sensitivity C-reactive protein; HCY, homocysteine.

Scale variables are summarized as mean±SD.

$^*$ Data from 2802 subjects at 6-year follow-up were included in the analysis.

$^P<0.05$, [HCY (-), hsCRP (+)] vs. [HCY (-), hsCRP (-)].

$^P<0.05$, [HCY (+), hsCRP (-)] vs. [HCY (-), hsCRP (+)].

$^P<0.05$, [HCY (+), hsCRP (+)] vs. [HCY (-), hsCRP (-)].

https://doi.org/10.1371/journal.pone.0206157.001

the four groups (hsCRP[-]/HCY[-] vs. hsCRP[-]/HCY[+] vs. hsCRP[+]/HCY[-] vs. hsCRP[+]/HCY[+], 14.1% vs. 20.2% vs. 20.3% vs. 33.0%, 22.3 vs. 26.3 vs. 33.8 vs. 58.6 per 1,000 person-year, $p<0.001$). In order to examine the odds ratios for the likelihood of MetS according to the presence of high hCRP and HCY levels, univariate and multiple logistic regression analyses were performed. The univariate analysis showed that the odds ratios for MetS in participants with either high hCRP or HCY level were 2.45 (95% CI, 1.99–3.03; $p<0.001$) and 1.77 (95% CI, 1.42–2.21; $p<0.001$) compared to participants with low hCRP and HCY levels, respectively. The multivariate analysis with adjustment for age, gender, smoking, alcohol status, vitamin supplement intake, and BMI showed that participants with the concurrent presence of high hCRP and HCY levels had a 2.70-fold increase (95% CI, 1.95–3.73; $p<0.01$) in the risk of MetS compared to those with low hCRP and HCY levels (Table 2). Moreover, Table 3 shows adjusted odds ratios for the estimated risk of developing MetS at the 6-year
follow-up among the four groups of participants who did not have MetS at baseline (n = 2383). After adjustment for age, gender, smoking, alcohol status, vitamin supplement intake, BMI at baseline, and ΔBMI at the 6-year follow-up in a regression model, the group with high hsCRP and HCY levels had a significantly higher risk of developing MetS upon follow-up (OR, 2.50, 95% CI, 1.50–4.16; p < 0.001) compared to the group with low hsCRP and HCY levels (Table 3).

Odds ratios for the risk of MetS according to the combination of hsCRP and HCY tertiles

Fig 2 presents the odds ratios for the estimated likelihood of MetS according to hsCRP and HCY tertiles. The multivariate analysis with adjustment for age, gender, smoking, alcohol status, vitamin supplement intake, and BMI showed that the odds ratios for MetS among the middle (0.42 mg/dl ≤ hsCRP < 1.01 mg/dl) and highest tertiles of hsCRP (hsCRP ≥ 1.01 mg/dl) with the highest tertile of HCY level (≥ 12.26 μmol/L) were 2.85 (95% CI, 1.82–4.47; p < 0.001) and 3.72 (95% CI, 2.40–5.76; p < 0.001), respectively, compared to those with the lowest hsCRP and HCY levels. In addition, the adjusted odds ratios for the risk of developing MetS at the 6-year follow-up among participants classified by similar cut-off points of hsCRP and HCY

### Table 2. Estimated odds ratios for the risk of MetS according to the combination of high hsCRP and HCY levels at baseline.

|                      | Estimated odds ratio (95% confidence interval) | P value |
|----------------------|-----------------------------------------------|---------|
|                      | hsCRP[-] HCY[-] hsCRP[+] HCY[+]               |         |
| Hypertriglyceridemia, n (%) | 501 (27.6) 192 (34.0) 233 (41.4) 117 (51.1) | <0.001  |
| Unadjusted           | Reference 1.36 (1.11–1.66)§ 1.84 (1.51–2.24)$ | <0.001  |
|                      | Adjusted, Model 1 1.33 (1.08–1.64)$           | <0.001  |
|                      | Adjusted, Model 2 Reference 1.23 (0.99–1.53)  | <0.001  |
| Low HDL cholesterol, n (%) | 886 (48.9) 292 (51.7) 352 (62.5) 154 (67.2) | <0.001  |
| Unadjusted           | Reference 1.13 (0.93–1.36) 1.73 (1.43–2.10)$ | <0.001  |
|                      | Adjusted, Model 1 Reference 1.10 (0.90–1.35) | <0.001  |
|                      | Adjusted, Model 2 Reference 1.05 (0.86–1.28)  | <0.001  |
| Hypertension, n (%)  | 513 (28.3) 262 (46.4) 212 (37.7) 127 (55.5) | <0.001  |
| Unadjusted           | Reference 2.21 (1.82–2.68)$ 1.52 (1.25–1.86)$ | <0.001  |
|                      | Adjusted, Model 1 Reference 1.90 (1.54–2.31)$ | <0.001  |
|                      | Adjusted, Model 2 Reference 1.80 (1.46–2.22)$ | <0.001  |
| High fasting glucose, n (%) | 422 (23.3) 138 (24.4) 188 (33.4) 90 (39.3) | <0.001  |
| Unadjusted           | Reference 1.070 (0.86–1.34) 1.64 (1.34–2.02)$ | <0.001  |
|                      | Adjusted, Model 1 Reference 0.93 (0.74–1.18) 1.58 (1.28–1.96)$ | <0.001  |
|                      | Adjusted, Model 2 Reference 0.83 (0.66–1.06) 1.20 (0.96–1.50) | <0.001  |
| MetS, n (%)          | 325 (17.9) 157 (27.8) 197 (35.0) 108 (47.2) | <0.001  |
| Unadjusted           | Reference 1.77 (1.42–2.21)$ 2.45 (1.99–3.03)$ 4.09 (3.07–5.44)$ | <0.001  |
|                      | Adjusted, Model 3 Reference 1.45 (1.13–1.86)$ 1.65 (1.30–2.09)$ 2.70 (1.95–3.73)$ | <0.001  |

Abbreviation: BMI, body mass index; HDL, high-density lipoprotein; hsCRP, high sensitivity C-reactive protein; HCY, homocysteine; MetS, metabolic syndrome.

Model 1: adjusted for age, sex, current smoking, current drinking, and vitamin intake.

Model 2: adjusted for Model 1 and waist circumference.

Model 3: adjusted for Model 1 and BMI.

* P < 0.05

§ P < 0.01

$ P < 0.001.

https://doi.org/10.1371/journal.pone.0206157.t002
Adjustments for age, gender, smoking, alcohol status, vitamin supplement intake, BMI at baseline, and ΔBMI at the 6-year follow-up showed that participants with the highest tertiles of hsCRP and HCY levels had a 2.15-fold increased risk of developing MetS (95% CI, 1.25–3.74; p < 0.01) compared to participants with the lowest hsCRP and Hcy levels. We investigated the interactions between hsCRP and HCY tertiles and their effects on the risk of MetS after adjusting for age, gender, smoking, alcohol status, vitamin supplement intake, BMI at baseline, and ΔBMI at the 6-year follow-up. No significant interactions were observed (Figs 2 and 3, p > 0.05).

**Discussion**

In a large population-based cohort study, we found that MetS was prevalent among participants with the concurrent presence of high hsCRP and HCY levels. The prevalence and incidence of MetS were highest in participants with elevated hsCRP and HCY levels (Tables 2 and 3), indicating that the combination of these conditions may exacerbate the risk for MetS. Even when adjusting for potential confounding factors in logistic regression models, participants with high hsCRP and HCY levels had a 2.70-fold risk of MetS at baseline and a 2.50-fold risk of MetS at the 6-year follow-up (Fig 3). Adjustments for age, gender, smoking, alcohol status, vitamin supplement intake, BMI at baseline, and ΔBMI at the 6-year follow-up showed that participants with the highest tertiles of hsCRP and HCY levels had a 2.15-fold increased risk of developing MetS (95% CI, 1.25–3.74; p < 0.01) compared to participants with the lowest hsCRP and Hcy levels. We investigated the interactions between hsCRP and HCY tertiles and their effects on the risk of MetS after adjusting for age, gender, smoking, alcohol status, vitamin supplement intake, BMI at baseline, and ΔBMI at the 6-year follow-up. No significant interactions were observed (Figs 2 and 3, p > 0.05).

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**Table 3. Estimated odds ratio for the risk of MetS according to the combination of high hsCRP and HCY levels at 6-year follow-up.**

| No. of subjects, n (%) | HCY[-] | HCY[+] | HCY[-] | HCY[+] | P value |
|-----------------------|--------|--------|--------|--------|---------|
| Hypertension, n (%)   | 1228 (63.0) | 317 (16.3) | 306 (15.7) | 97 (5.0) | -       |
| Unadjusted Reference  | 1.26 (0.94–1.68) | 1.51 (1.14–2.00) | 1.96 (1.26–3.05) | <0.001 |
| Adjusted, Model 1     | 1.32 (0.98–1.77) | 1.50 (1.13–2.00) | 2.20 (1.40–3.47) | <0.001 |
| Adjusted, Model 2     | 1.29 (0.96–1.74) | 1.39 (1.04–1.86) | 1.98 (1.25–3.13) | <0.001 |
| Low HDL cholesterol, n (%) | 503 (41.0) | 144 (45.4) | 137 (44.8) | 52 (53.6) | 0.05    |
| Unadjusted Reference  | 1.20 (0.94–1.54) | 1.17 (0.91–1.50) | 1.67 (1.10–2.52) | <0.05   |
| Adjusted, Model 1     | 1.21 (0.93–1.56) | 1.16 (0.90–1.50) | 1.64 (1.07–2.51) | <0.05   |
| Adjusted, Model 2     | 1.17 (0.90–1.52) | 1.04 (0.80–1.35) | 1.42 (0.92–2.19) | 0.19    |
| Hypertension, n (%)   | 381 (31.0) | 142 (44.8) | 118 (38.6) | 47 (48.5) | <0.001  |
| Unadjusted Reference  | 1.80 (1.40–2.32) | 1.40 (1.08–1.81) | 2.09 (1.38–3.17) | <0.001  |
| Adjusted, Model 1     | 1.54 (1.18–2.00) | 1.33 (1.02–1.73) | 1.63 (1.06–2.52) | <0.01   |
| Adjusted, Model 2     | 1.50 (1.15–1.95) | 1.20 (0.91–1.57) | 1.43 (0.92–2.22) | 0.02    |
| High fasting glucose, n (%) | 198 (16.1) | 60 (18.9) | 65 (21.2) | 23 (23.7) | <0.05   |
| Unadjusted Reference  | 1.21 (0.88–1.67) | 1.40 (1.03–1.92) | 1.62 (0.99–2.64) | <0.01   |
| Adjusted, Model 1     | 1.10 (0.79–1.52) | 1.33 (0.97–1.82) | 1.43 (0.86–2.37) | <0.05   |
| Adjusted, Model 2     | 1.05 (0.75–1.46) | 1.17 (0.85–1.62) | 1.20 (0.72–2.00) | 0.27    |
| Metabolic syndrome, n (%) | 173 (14.1) | 64 (20.2) | 62 (20.3) | 32 (33.0) | <0.001  |
| Unadjusted Reference  | 1.54 (1.12–2.12) | 1.55 (1.12–2.14) | 3.00 (1.91–4.72) | <0.001  |
| Adjusted, Model 3     | 1.24 (0.87–1.77) | 1.20 (0.84–1.72) | 2.50 (1.50–4.16) | <0.001  |

Abbreviation: BMI, body mass index; HDL, high-density lipoprotein; hsCRP, high sensitivity C-reactive protein; HCY, homocysteine; MetS, metabolic syndrome.

Model 1: adjusted for age, sex, current smoking, current drinking, and vitamin intake.

Model 2: adjusted for Model 1 and waist circumference.

Model 3: adjusted for Model 1, BMI at baseline, and BMI change at follow-up (ΔBMI).

*P < 0.05

§P < 0.01

*P < 0.001.

https://doi.org/10.1371/journal.pone.0206157.t003
increased risk for developing MetS at the 6-year follow-up compared to those with low hsCRP and HCY levels. Moreover, in a model that investigated the risk of MetS and hsCRP and HCY tertiles, participants with the highest hsCRP (hsCRP ≥ 1.01 mg/dl) and HCY tertiles (≥12.26 μmol/L) had a 3.72-fold and a 2.15-fold increased risk of MetS at the 6-year follow-up compared to those with the lowest tertile of hsCRP and Hcy levels after adjustment for potential confounding factors. However, no significant interactions between the tertiles of hsCRP and Hcy levels were examined, suggesting that the two factors may not synergistically affect the increased risk of MetS. However, to the best of our knowledge, this is the first study to investigate the combinatory effect of elevated hsCRP and HCY on the development of MetS using data from a large prospective cohort study. Therefore, the magnitude and significance of the modifying effect of high hsCRP and HCY levels in individuals on the risk of MetS should be confirmed in other cohort studies.

Fig 2. Estimated odds ratios for risk of MetS according to hsCRP and HCY tertiles. The odds ratio was estimated after adjusting for age, sex, smoking and alcohol status, vitamin intake, and BMI (n = 3170). P-value for interaction = 0.45, *P < 0.05, **P < 0.01.

https://doi.org/10.1371/journal.pone.0206157.g002
Considering the ever-increasing body of evidence regarding MetS, chronic low-grade inflammation, accumulation of reactive oxygen species, and activation of the immune system may have an important role in the pathogenesis of obesity-related metabolic disorders [10, 29]. Previous evidence from clinical and epidemiologic studies suggested that elevated hsCRP and HCY levels share many common pathophysiological pathways inflicting damage on the cardiovascular system [30]. However, it is unclear whether the coexistence of elevated hsCRP and HCY levels accelerates the risk of MetS. HsCRP is a robust biomarker of underlying systemic inflammation and appears to be an important marker in both cardiovascular disease and metabolic syndrome [31, 32]. While the pathophysiologic mechanisms by which systemic inflammation leads to increased risk of MetS are not clearly evident, both adipokines produced in hypertrophic adipocytes [10] and production of pro-inflammatory cytokines such as TNF-α,
IL-6, and monocyte chemoattractant protein-1 released from adipose tissue of increased infiltration by macrophages may play an important role in the pathogenesis of MetS [33, 34]. In addition, data from clinical and epidemiologic studies also shows that elevated hsCRP level in healthy individuals is associated with increased risk of MetS [35] as well as future development of insulin resistance and diabetes [36] in healthy individuals. HCY is a thiol-containing intermediate amino-acid in the methionine to cysteine pathway. Over the past several decades, a number of studies reported that elevated HCY was related to endothelial dysfunction, arterial intimal-media wall thickening, and a pro-thrombotic state, providing a pathophysiologic explanation for the increased risk of CVD morbidity and mortality [21]. In animal studies, insulin affected the activity of enzymes involved in HCY turnover, and hyperhomocysteinemia has been suggested as a possible additional component of MetS [37]. Even though several population-based studies showed a significant association between MetS and plasma HCY level [38], results from these studies were limited due to cross-sectional design, small sample size, and inadequate control of confounders [39]. Therefore, a potential causal relationship between HCY level and MetS has not been clearly established. Recent evidence from a 4.8-year longitudinal study in a Chinese population also reported no significant association between baseline HCY and the incidence of MetS [40]. We found similar results; the highest prevalence and incidence of MetS at the 6-year follow-up were among participants with high hsCRP and HCY levels. However, the incidence rate of MetS among individuals with elevated hsCRP and HCY levels relative to those without them might be influenced by the high rate of non-respondents among subjects with high hsCRP and HCY levels at the 6-year follow-up, although this was not statistically different (non-respondents, n = 435; hsCRP[–]/HCY[–] vs. hsCRP[–]/HCY[+] vs. hsCRP[+]/HCY[–] vs. hsCRP[+]/HCY[+], 17.5% vs. 21.9% vs. 16.8% vs. 19.8%, p = 0.175). Moreover, elevated hsCRP level and elevated HCY level alone were significantly associated with increased risk of MetS at baseline; however, these were not predictive of the incidence of MetS at 6-year follow-up when compared to those with low hsCRP and HCY levels. Only the group with elevated hsCRP and HCY was significantly associated with increased risk of MetS, suggesting that the combination may exacerbate the risk for MetS. Therefore, the magnitude and significance of the modifying effects of elevated hsCRP and HCY on the development of MetS should be confirmed in other cohort studies.

The current study has several strengths that lend confidence to the findings. First, this study is the first to show that the co-existence of high hsCRP and HCY is correlated with development of MetS in a large general population. Second, this was a longitudinal study, which allowed us to establish a causal relationship of hsCRP and HCY levels with MetS. Since the characteristics of the study population including biochemical data, metabolic scores, BMI, hsCRP, HCY level, and blood pressure were not significantly different between participants and non-participants at baseline (p>0.05), our results are likely representative of the general population.

Although our study adapted a prospective design with follow-up measurements and used a large general population sample, several limitations should be acknowledged. First, we used hsCRP as a marker to define the inflammatory status of the participants, but the use of only hsCRP may not sufficiently reflect the inflammatory status of participants. The combined effects of other inflammatory cytokines and HCY on the development of MetS should be examined in further studies. Second, the group with high hsCRP and HCY levels has a relatively small sample size compared to the other groups. Third, mild to moderate elevation of HCY level in the general population may be partially influenced by participant nutritional habits and exercise status; these factors were not considered in the present study. Our data showed that participants with regular intake of any type of vitamin supplement had significantly lower levels of HCY compared to those who did not take vitamin supplements. Even though the
effect of vitamin supplementation on reducing the risk of type 2 diabetes and CVD is still controversial [41, 42], several lines of research suggest that vitamin supplementation with folate, B6, and B12 has the potential to lower HCY level and thus cardio-metabolic risk [43]. Finally, genetic variation or epigenetic modifications that might play an important role in the elevation of hsCRP and HCY level were not examined in the present study. Accordingly, future longitudinal studies that focus on interactions between hsCRP- and HCY-related genetic variations and the development of MetS should be conducted.

In conclusion, MetS is more prevalent in the concurrent presence of elevated hsCRP and HCY levels. Taken together, combined elevation of hsCRP and HCY levels were associated with increased risk of MetS, but these factors may not be synergistic. Additional longitudinal study is needed to further address the significance and magnitude of the coexistence of these conditions on the development of MetS in the context of reducing CVD risk.

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References
1. Gami AS, Witt BJ, Howard DE, Erwin PJ, Gami LA, Somers VK, et al. Metabolic syndrome and risk of incident cardiovascular events and death: a systematic review and meta-analysis of longitudinal studies. J Am Coll Cardiol. 2007; 49(4):403–14. https://doi.org/10.1016/j.jacc.2006.09.032 PMID: 17258085
2. Collaboration IRGCERF, Sarwar N, Butterworth AS, Freitag DF, Gregson J, Willeit P, et al. Interleukin-6 receptor pathways in coronary heart disease: a collaborative meta-analysis of 82 studies. Lancet. 2012; 379(9822):1205–13. https://doi.org/10.1016/S0140-6736(11)61931-4 PMID: 22421339
3. Ferreira I, Twisk JW, van Mechelen W, Kemper HC, Stehouwer CD. Development of fatness, fitness, and lifestyle from adolescence to the age of 36 years: determinants of the metabolic syndrome in young adults: the amsterdam growth and health longitudinal study. Arch Intern Med. 2005; 165(1):42–8. https://doi.org/10.1001/archinte.165.1.42 PMID: 15642873
4. Kassi E, Pervanidou P, Kaltsas G, Chrousos G. Metabolic syndrome: definitions and controversies. BMC Med. 2011; 9:48. https://doi.org/10.1186/1741-7015-9-48 PMID: 21542944
5. Devaraj S, Hemarajata P, Versalovic J. The human gut microbiome and body metabolism: implications for obesity and diabetes. Clin Chem. 2013; 59(4):617–28. https://doi.org/10.1373/clinchem.2012.187617 PMID: 23401296
6. Zampetaki A, Mayr M. MicroRNAs in vascular and metabolic disease. Circ Res. 2012; 110(3):508–22. https://doi.org/10.1161/CIRCRESAHA.111.247445 PMID: 22302757
7. Jean-Louis G, Zizi F, Brown D, Ogudegbe G, Borer J, McFarlane S. Obstructive sleep apnea and cardiovascular disease: evidence and underlying mechanisms. Minerva Pneumol. 2009; 48(4):277–93. PMID: 21645544
8. Despres JP, Lemieux I, Bergeron J, Pibarot P, Mathieu P, Larose E, et al. Abdominal obesity and the metabolic syndrome: contribution to global cardiometabolic risk. Arterioscler Thromb Vasc Biol. 2008; 28(6):1039–49. https://doi.org/10.1161/ATVBAHA.107.159228 PMID: 18356555
9. Bray GA, Bellanger T. Epidemiology, trends, and morbidities of obesity and the metabolic syndrome. Endocrine. 2006; 29(1):109–17. https://doi.org/10.1385/ENDO:29:1:109 PMID: 16622298
10. Yudkin JS, Kumari M, Humphries SE, Mohamed-Ali V. Inflammation, obesity, stress and coronary heart disease: is interleukin-6 the link? Atherosclerosis. 2000; 148(2):209–14. PMID: 10657556

11. Choi J, Joseph L, Pilote L. Obesity and C-reactive protein in various populations: a systematic review and meta-analysis. Obes Rev. 2013; 14(3):232–44. https://doi.org/10.1111/obr.12003 PMID: 23171381

12. Brooks GC, Blisha MJ, Blumenthal RS. Relation of C-reactive protein to abdominal adiposity. Am J Cardiol. 2010; 106(1):56–61. https://doi.org/10.1016/j.amjcard.2010.02.017 PMID: 20690648

13. Nakamura H, Ito H, Egami Y, Kaji Y, Maruyama T, Koike G, et al. Waist circumference is the main determinant of elevated C-reactive protein in metabolic syndrome. Diabetes Res Clin Pract. 2008; 79(2):330–6. https://doi.org/10.1016/j.diabres.2007.09.004 PMID: 17964679

14. Visser M, Bouter LM, McQuillan GM, Wener MH, Harris TB. Elevated C-reactive protein levels in overweight and obese adults. JAMA. 1999; 282(22):2131–5. PMID: 10591334

15. Park HS, Park JY, Yu R. Relationship of obesity and visceral adiposity with serum concentrations of CRP, TNF-alpha and IL-6. Diabetes Res Clin Pract. 2005; 69(1):29–35. https://doi.org/10.1016/j.diabres.2004.11.007 PMID: 15955385

16. Freeman DJ, Norrie J, Castlake MJ, Gaw A, Ford I, Lowe GD, et al. C-reactive protein is an independent predictor of risk for the development of diabetes in the West of Scotland Coronary Prevention Study. Diabetes. 2002; 51(5):1596–600. PMID: 11978661

17. Devaraj S, Singh U, Jialal I. Human C-reactive protein and the metabolic syndrome. Curr Opin Lipidol. 2009; 20(3):182–9. https://doi.org/10.1097/MOL.0b013e32832ac03e PMID: 19369869

18. Francisco G, Hernandez C, Simo R. Serum markers of vascular inflammation in dyslipidaemia. Clin Chim Acta. 2006; 369(1):1–16. https://doi.org/10.1016/j.cca.2005.12.027 PMID: 16469304

19. Ridker PM, Wilson PW, Grundy SM. Should C-reactive protein be added to metabolic syndrome and to assessment of global cardiovascular risk? Circulation. 2004; 109(23):2818–25. https://doi.org/10.1161/01.CIR.0000132467.45278.59 PMID: 15197153

20. Rifai N. High-sensitivity C-reactive protein: a useful marker for cardiovascular disease risk prediction and the metabolic syndrome. Clin Chem. 2005; 51(3):504–5. https://doi.org/10.1373/clinchem.2004.044990 PMID: 15738514

21. Mangoni AA, Jackson SH. Homocysteine and cardiovascular disease: current evidence and future prospects. Am J Med. 2002; 112(7):556–65. PMID: 12015248

22. Bjorck J, Hellgren M, Rastam L, Lindblad U. Associations between serum insulin and homocysteine in a Swedish population-a potential link between the metabolic syndrome and hyperhomocysteinemia: the Skaraborg project. Metabolism. 2006; 55(8):1007–13. https://doi.org/10.1016/j.metabol.2006.03.010 PMID: 16839834

23. Sreckovic B, Sreckovic VD, Soldatovic I, Colak E, Sumarac-Dumanovic M, Janeski H, et al. Homocysteine is a marker for metabolic syndrome and atherosclerosis. Diabetes Metab Syndr. 2017; 11(3):179–82. https://doi.org/10.1016/j.dsx.2016.08.026 PMID: 27600468

24. Kim Y, Han BG. Cohort Profile: The Korean Genome and Epidemiology Study (KoGES) Consortium. Int J Epidemiol. 2017; 46(4):1350. https://doi.org/10.1093/ije/dyx105 PMID: 28937572

25. Markanday A. Acute Phase Reactants in Infections: Evidence-Based Review and a Guide for Clinicians. Open Forum Infect Dis. 2015; 2(3):ofv098. https://doi.org/10.1093/ofid/ofv098 PMID: 26258155

26. Lentz SR, Haynes WG. Homocysteine: is it a clinically important cardiovascular risk factor? Cleve Clin J Med. 2004; 71(9):729–34. PMID: 15478704

27. Expert Panel on Detection E, Treatment of High Blood Cholesterol in A. Executive Summary of The Third Report of The National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, And Treatment of High Blood Cholesterol In Adults (Adult Treatment Panel III). JAMA. 2001; 285(19):2486–97. PMID: 11369702

28. Kim JA, Choi CJ, Yum KS. Cut-off values of visceral fat area and waist circumference: diagnostic criteria for abdominal obesity in a Korean population. J Korean Med Sci. 2006; 21(6):1048–53. https://doi.org/10.3346/jkms.2006.21.6.1048 PMID: 17179685

29. Yudkin JS, Stehouwer CD, Emeis JJ, Coppack SW. C-reactive protein in healthy subjects: associations with obesity, insulin resistance, and endothelial dysfunction; a potential role for cytokines originating from adipose tissue? Arterioscler Thromb Vasc Biol. 1999; 19(4):972–8. PMID: 10195925

30. Nishida K, Otsu K. Inflammation and metabolic cardiomyopathy. Cardiovasc Res. 2017; 113(4):389–98. https://doi.org/10.1093/crv/cvx012 PMID: 28395010

31. Timpson NJ, Lawlor DA, Harbord RM, Gaunt TR, Day IN, Palmer LJ, et al. C-reactive protein and its role in metabolic syndrome: mendelian randomisation study. Lancet. 2005; 366(9501):1954–9. https://doi.org/10.1016/S0140-6736(05)67788-0 PMID: 16325697
32. Dehghan A, Dupuis J, Barbalic M, Bis JC, Eiriksdottir G, Lu C, et al. Meta-analysis of genome-wide association studies in >80,000 subjects identifies multiple loci for C-reactive protein levels. Circulation. 2011; 123(7):731–8. https://doi.org/10.1161/CIRCULATIONAHA.110.948570 PMID: 21300955

33. Maury E, Brichard SM. Adipokine dysregulation, adipose tissue inflammation and metabolic syndrome. Mol Cell Endocrinol. 2010; 314(1):1–16. https://doi.org/10.1016/j.mce.2009.07.031 PMID: 19682539

34. Murakami S. The physiological and pathophysiological roles of taurine in adipose tissue in relation to obesity. Life Sci. 2017; 186:80–6. https://doi.org/10.1016/j.lfs.2017.08.008 PMID: 28801262

35. Al Rifai M, Silverman MG, Nasir K, Budoff MJ, Blankstein R, Szkiel M, et al. The association of nonalcoholic fatty liver disease, obesity, and metabolic syndrome, with systemic inflammation and subclinical atherosclerosis: the Multi-Ethnic Study of Atherosclerosis (MESA). Atherosclerosis. 2015; 239(2):629–33. https://doi.org/10.1016/j.atherosclerosis.2015.02.011 PMID: 25683387

36. Ridker PM. A Test in Context: High-Sensitivity C-Reactive Protein. J Am Coll Cardiol. 2016; 67(6):712–23. https://doi.org/10.1016/j.jacc.2015.11.037 PMID: 26868696

37. Oron-Herman M, Rosenthal T, Sela BA. Hyperhomocysteinemia as a component of syndrome X. Metabolism. 2003; 52(11):1491–5. PMID: 14624412

38. Kumar A, Palfrey HA, Pathak R, Kadowitz PJ, Gettys TW, Murthy SN. The metabolism and significance of homocysteine in nutrition and health. Nutr Metab (Lond). 2017; 14:78. https://doi.org/10.1186/s12986-017-0233-z PMID: 29299040

39. Nabipour I, Ebrahimi A, Jafari SM, Vahdat K, Movahed A, et al. The metabolic syndrome is not associated with homocysteinemia: the Persian Gulf Healthy Heart Study. J Endocrinol Invest. 2009; 32(5):406–10. https://doi.org/10.1007/BF03346476 PMID: 19794288

40. Wang X, Ye P, Cao R, Yang X, Xiao W, Zhang Y, et al. The association of homocysteine with metabolic syndrome in a community-dwelling population: homocysteine might be concomitant with metabolic syndrome. PLoS One. 2014; 9(11):e113148. https://doi.org/10.1371/journal.pone.0113148 PMID: 25401978

41. Song Y, Cook NR, Albert CM, Van Denburgh M, Manson JE. Effect of homocysteine-lowering treatment with folate Acid and B vitamins on risk of type 2 diabetes in women: a randomized, controlled trial. Diabetes. 2009; 58(8):1921–8. https://doi.org/10.2337/db09-0087 PMID: 19491213

42. Hannibal L, Blom HJ. Homocysteine and disease: Causal associations or epiphenomenons? Mol Aspects Med. 2017; 53:36–42. https://doi.org/10.1016/j.mam.2016.11.003 PMID: 27876556

43. Alexander DD, Weed DL, Chang ET, Miller PE, Mohamed MA, Elkayam L. A systematic review of multivitamin-multimineral use and cardiovascular disease and cancer incidence and total mortality. J Am Coll Nutr. 2013; 32(5):339–54. https://doi.org/10.1080/07315724.2013.839909 PMID: 24219377