Adverse Pregnancy Outcomes and Maternal Chronic Diseases in the Future: a Cross-sectional Study Using KoGES-HEXA Data

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Research Article

Keywords: metabolic syndrome, preeclampsia, gestational diabetes mellitus, low birth weight, macrosomia, cardiovascular disease

Posted Date: December 13th, 2021

DOI: https://doi.org/10.21203/rs.3.rs-1062386/v1

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Abstract

**Purpose** Adverse pregnancy outcomes (APOs) are associated with an increased risk of chronic diseases, including cardiovascular disease (CVD) and metabolic syndrome (MS), in the future. We designed a large-scale cohort study to evaluate the influence of APOs (preeclampsia, gestational diabetes mellitus [GDM], stillbirth, macrosomia, and low birth weight) on the incidence of chronic diseases, body measurements, and serum biochemistry in the future and investigate whether combinations of APOs had additive effects on chronic diseases.

**Methods** We used health examinee data from the Korean Genome and Epidemiology Study (KoGES-HEXA) and extracted data of parous women (n = 30,174; mean age, 53.02 years) for the analysis. A logistic regression model was used to estimate adjusted odds ratios (aORs) and 95% confidence intervals.

**Results** The prevalence rate of APOs was 17.4% (5,264 women). Univariate analysis revealed that women with APOs were more frequently diagnosed with chronic diseases and had a family history of chronic diseases compared with women without APOs. In logistic regression analysis, composite APOs were associated with an increased risk of hypertension, diabetes mellitus, hyperlipidemia, angina pectoris, stroke, and MS (aOR: 1.093, 1.379, 1.269, 1.351, 1.414, and 1.104, respectively), after adjustment for family history and social behaviors. Preeclampsia and GDM were associated with an increased risk of some chronic diseases; however, the combination of preeclampsia and GDM did not have an additive effect on the risk.

**Conclusion** APOs moderately influenced the future development of maternal CVD and metabolic derangements, independent of family history and social behaviors.

Introduction

Chronic diseases, including cardiovascular disease (CVD) and its risk factors such as diabetes mellitus and metabolic syndrome (MS), are now recognized as the leading factors that threaten women's health worldwide. Prevalence rates of both coronary heart disease (CHD) and stroke have risen to similar levels as those of men of a similar age [1, 2]. CHD mortality rates have increased in American women within the age range of 35–54 years, contrary to a decreasing trend in the past four decades from 1980 to 2000 [1, 3, 4]. Additionally, in Korea, the CVD mortality rate is even 1.1 times higher in women than that in men [5]. These suggest that underlying factors leading to CVD may be different in women compared with men, and identification of pathophysiology unique to women has considerable potential to improve the long-term health status of women.

With an improved understanding of cardiovascular changes during normal pregnancy and the changes that are associated with adverse pregnancy outcomes (APOs), it is now recognized that obstetric events can contribute to gender-specific differences in cardiovascular risk and the development of CVD later in life [6]. APOs have also been linked to MS, a major risk factor for CVD [7, 8]. Guidelines for the prevention...
of CVD in women include a history of APOs, such as preeclampsia, gestational diabetes mellitus (GDM), preterm delivery, pregnancy-induced hypertension, and delivery of a low birth weight (LBW) infant as major CVD risk factors [1, 9–11]. The link between APOs and future chronic diseases mainly focuses on preeclampsia and GDM. Many epidemiologic studies and meta-analyses have shown that preeclampsia is related to an increased risk of developing or dying from overall CVD [12–14]. Moreover, preeclampsia is positively associated with cardiovascular risk factors such as hypertension, diabetes mellitus, and MS [12, 15–17]. GDM is also associated with a significant increase in the risk factors for CVD, including diabetes mellitus and hypertension [18–20]. However, other APOs, such as macrosomia, LBW, and stillbirth, have not been extensively studied. Furthermore, whether APOs cause various chronic diseases and affect serum biochemistry in later life and whether a combination of APOs confers higher risks of chronic diseases need to be investigated. Additionally, many studies have failed to adjust for factors such as social behaviors and family history of cardiometabolic status, thus limiting the understanding of CVD risk trajectory.

We, therefore, designed a large-scale cohort study to (1) evaluate the influence of APOs (preeclampsia, GDM, stillbirth, macrosomia, LBW) on the incidence of chronic diseases, body measurements, and serum biochemistry in later life, (2) assess whether APOs were independent risk factors for chronic diseases, and (3) investigate the additive effects of combinations of APOs on chronic diseases.

**Methods**

**Study Population and design**

This cohort study relied on data from the Korean Genome and Epidemiology Study (KoGES). The KoGES is a national survey conducted by the Korean government (National Research Institute of Health, Centers for Disease Control and Prevention, and Ministry for Health and Welfare, Republic of Korea) to investigate the genetic and environmental etiology of common complex diseases in Koreans and causes of death. Details of the KoGES and the methods used were comprehensively explored [21]. Among the KoGES Consortium data, we used KoGES health examinee (HEXA) data consisting of data from participants older than 40 years who resided in urban areas. The HEXA cohort included participants from examination centers/medical institutions of 14 major cities across Korea who were recruited between 2004 and 2013. All participants voluntarily signed an informed consent form and attended follow-up from 2012 to 2016. This study was approved by the Ethics Committee of Korea University (2021GR0007).

**Participant Selection**

Among 113,945 women, women who lacked records of pregnancy outcome histories (n = 25,803), who lacked records of previous medical history, including family history (n = 57,808), and with a history of CVD before pregnancy (n = 139) were excluded. Finally, 30,174 participants were included in the study.

**Survey**
The HEXA consisted of two components: a health interview and a health examination. The health interview included questions regarding demographics, lifestyle status, medical history, and pregnancy outcomes. Each participant was asked regarding the previous history of pregnancy outcomes, including preeclampsia, GDM, stillbirth, macrosomia (birth weight of the neonate $\geq 4$ kg), and LBW (birth weight of the neonate $\leq 2.5$ kg), by trained interviewers. Each participant was also asked regarding the medical history of chronic diseases (hypertension, diabetes mellitus, hyperlipidemia, stroke, transient cerebral ischemia, angina pectoris, and Parkinson's disease) and family history of chronic diseases. Data for the following covariates were also obtained: age, age at delivery, lactation history, alcohol consumption, and smoking status. Smoking status was divided into two categories: current smoker and non-smoker. Drinking status was also classified into two groups: current drinker and non-drinker.

The health examination included anthropometric measurements and serum laboratory tests. All anthropometric measurements were obtained by trained and skilled examiners using a consistent and standardized methodology. Height was measured to the nearest 1 mm using a portable extensimeter (SECA 225; SECA, Hamburg, Germany). Weight was measured to the nearest 0.1 kg using a calibrated balance scale (GL-6000-20; CAS KOREA, Seoul, Korea). Waist circumference (WC) was measured at the narrowest point between the lower border of the rib cage and the iliac crest during minimal respiration. Hip circumference (HC) was measured at the widest part of the hip in the horizontal plane. Blood pressure (BP) was measured using a standard mercury sphygmomanometer.

All blood samples were obtained after the participants had fasted for a minimum of eight hours. The levels of fasting blood sugar (FBS), $\gamma$-glutamyl transferase, aspartate aminotransferase, alanine aminotransferase, albumin, blood urea nitrogen, creatinine, uric acid, total cholesterol, high-density lipoprotein cholesterol (HDL-C), triglycerides, and high-sensitivity C-reactive protein were measured using enzymatic methods.

**Covariates**

Composite APOs were defined if any of the APOs was present. MS was defined using the criteria proposed by the Joint Interim Statement of the International Diabetes Federation Task Force on Epidemiology and Prevention [22]. MS was defined as the presence of at least three of the five metabolic components described as follows: (i) a WC $\geq 80$ cm, according to the International Diabetes Federation criteria for Asian countries; (ii) FBS $\geq 100$ mg/dL or treatment for elevated glucose; (iii) serum fasting triglyceride $\geq 150$ mg/dL (1.7 mmol/L) or medication use; (iv) serum HDL-C $< 50$ mg/dL (1.3 mmol/L) or medication use; and (v) systolic BP $\geq 130$ mmHg, diastolic BP $\geq 85$ mmHg, or drug treatment for hypertension.

**Statistical analysis**

Continuous and categorical variables are expressed as mean $\pm$ standard deviation and percentages, respectively. Basic characteristics, body measurements, and serum biochemistry of the study population were compared between women with APOs and women without APOs, using t-test for continuous
variables and chi-square test for categorical variables. Multivariate logistic regression analysis was used to estimate the adjusted odds ratios (aORs) and 95% confidence intervals (CIs) for the development of chronic diseases. For multivariable analyses, we adjusted for maternal age, family history (hypertension, diabetes, angina, and stroke), present smoking, and present alcohol consumption. The significance level was defined as a p-value less than 0.05. Statistical analyses were performed using SAS for Windows (version 9.4; SAS Inc., Cary, NC, USA).

Results

APOs occurred in 5,264 women in the study population (prevalence rate, 17.4%), and the mean age of the participants was 53.02 years. The characteristics of the study participants are listed in Table 1. Women with APOs had lower age (51.70 ± 7.68 vs. 53.42 ± 8.08 years, p < 0.001) but had higher age at first delivery (25.81 ± 3.00 vs. 25.32 ± 3.01 years, p < 0.001) and last delivery (30.22 ± 3.84 vs. 29.75 ± 3.67 years, p < 0.001) than women without APOs. Compared with those without APOs, those with APOs were more frequently diagnosed with some chronic diseases such as diabetes mellitus and hypercholesterolemia and had a family history of chronic diseases, including hypertension, diabetes mellitus, angina pectoris, and stroke. They were also more likely to smoke and drink alcohol and less likely to breastfeed. Prevalence rates of MS and its components were not statistically different between the two groups, except in women with high FBS levels.
Table 1
The characteristics of the study population

| Baseline characteristics | Women without APOs (n = 24,910) | Women with APOs (n = 5,264) | p-value |
|--------------------------|----------------------------------|-----------------------------|---------|
| Age, y                   | 53.31 ± 8.27                     | 51.70 ± 7.56                | <0.001  |
| Age of first delivery, y | 25.09 ± 2.95                     | 25.65 ± 2.92                | <0.001  |
| Age of last delivery, y  | 29.88 ± 3.69                     | 30.21 ± 3.78                | <0.001  |

Chronic diseases

| Hypertension             | 4537 (18.21)                     | 942 (17.90)                 | 0.586   |
| Diabete                 | 1267 (5.09)                      | 314 (5.97)                  | 0.009   |
| Hyperlipidemia           | 1549 (6.22)                      | 381 (7.24)                  | 0.006   |
| Stroke                   | 275 (1.10)                       | 71 (1.35)                   | 0.130   |
| Transient cerebral ischemia | 41 (0.16)                      | 6 (0.11)                    | 0.398   |

Angina pectoris            | 463 (1.86)                      | 110 (2.09)                  | 0.265   |

Parkinson’s disease        | 20 (0.08)                       | 3 (0.06)                    | 0.659   |

Family history of chronic diseases

| Hypertension             | 6410 (25.73)                     | 1672 (31.76)                | <0.001  |
| Diabete                 | 3737 (15.00)                     | 1011 (19.21)                | <0.001  |
| Stroke                   | 3378 (13.56)                     | 848 (16.11)                 | <0.001  |

Angina pectoris            | 1558 (6.25)                      | 415 (7.88)                  | <0.001  |

Behavioral variables

| Current smoking          | 581 (2.33)                       | 148 (2.81)                  | 0.040   |
| Current alcohol drinking | 7381 (29.63)                     | 1652 (31.38)                | 0.012   |

Data are presented as mean ± standard error or n (%).

APOs adverse pregnancy outcomes; BP blood pressure; FBS fasting blood sugar; TG total triglyceride; HDL-C high-density lipoprotein cholesterol.

Components: abdominal obesity was defined as a waist circumference ≥80 cm. Elevated BP was defined as systolic/diastolic pressure ≥130/85 mm Hg or drug treatment for hypertension. Elevated FBS was defined as FBS level ≥100 mg/dL or treatment for elevated glucose. High TG were defined as TG ≥150 mg/dL or medication use. Low HDL-C was defined as an HDL-C level <50 mg/dL or medication use.
|                                | Women without APOs (n = 24,910) | Women with APOs (n = 5,264) | p-value |
|--------------------------------|---------------------------------|-----------------------------|---------|
| Lactation history              |                                 |                             |         |
| Lactation                      | 22715 (91.19)                   | 4651 (88.35)                | <0.001  |
| Sibling number of lactations   | 2.55 ± 1.07                     | 2.35 ± 0.91                 | <0.001  |
| Duration of lactation, m       | 31.40 ± 28.65                   | 26.52 ± 22.38               | <0.001  |
| Metabolic syndrome and its components<sup>a</sup> |                                 |                             |         |
| Metabolic syndrome             | 5018 (20.14)                    | 1023 (19.43)                | 0.242   |
| Abdominal obesity alone        | 6710 (26.94)                    | 1455 (27.64)                | 0.297   |
| Elevated BP alone              | 9986 (40.09)                    | 2064 (39.21)                | 0.237   |
| Elevated FBS alone             | 5752 (23.09)                    | 1314 (24.96)                | 0.004   |
| High TG alone                  | 5432 (21.81)                    | 1110 (21.09)                | 0.250   |
| Low HDL-C alone                | 7452 (29.92)                    | 1519 (28.86)                | 0.127   |

Data are presented as mean ± standard error or n (%).

APOs adverse pregnancy outcomes; BP blood pressure; FBS fasting blood sugar; TG total triglyceride; HDL-C high-density lipoprotein cholesterol.

<sup>a</sup>Components: abdominal obesity was defined as a waist circumference ≥80 cm. Elevated BP was defined as systolic/diastolic pressure ≥130/85 mm Hg or drug treatment for hypertension. Elevated FBS was defined as FBS level ≥100 mg/dL or treatment for elevated glucose. High TG were defined as TG ≥150 mg/dL or medication use. Low HDL-C was defined as an HDL-C level <50 mg/dL or medication use.

Body measurements and laboratory data were also compared between the two groups (Table 2). Women with APOs had greater height, weight, WC, and HC than women without APOs. Mean values of FBS and serum albumin levels were statistically different between the two groups.
| Table 2                                                                 |
|------------------------------------------------------------------------|
| **The body measurements and laboratory data of the study population**  |

|                                | **Women without APOs** | **Women with APOs** | **p-value** |
|--------------------------------|------------------------|---------------------|-------------|
| (n = 24,910)                   | (n = 5,264)            |                     |             |
| **Body measurements**          |                        |                     |             |
| Height, cm                    | 155.65 ± 5.28          | 156.53 ± 5.23       | <0.001      |
| Weight, Kg                    | 57.82 ± 7.36           | 59.17 ± 7.88        | <0.001      |
| WC, cm                        | 79.73 ± 8.03           | 80.05 ± 8.20        | 0.009       |
| HC, cm                        | 94.37 ± 5.54           | 94.95 ± 5.74        | <0.001      |
| PR                            | 67.82 ± 9.46           | 67.49 ± 9.61        | 0.032       |
| Systolic BP, mmHg             | 121.93 ± 16.42         | 121.41 ± 16.12      | 0.035       |
| Diastolic BP, mmHg            | 75.86 ± 10.09          | 75.79 ± 10.17       | 0.630       |
| **Laboratory findings**       |                        |                     |             |
| FBS, mg/dL                    | 93.21 ± 19.65          | 94.62 ± 24.61       | <0.001      |
| GGT                           | 21.56 ± 22.67          | 21.59 ± 21.36       | 0.917       |
| AST                           | 23.11 ± 14.12          | 22.90 ± 13.60       | 0.320       |
| ALT                           | 20.16 ± 17.50          | 20.31 ± 24.04       | 0.666       |
| Albumin                       | 4.63 ± 0.28            | 4.64 ± 0.28         | 0.010       |
| BUN                           | 13.66 ± 3.78           | 13.56 ± 3.77        | 0.058       |
| Creatinine                    | 0.77 ± 0.18            | 0.77 ± 0.14         | 0.297       |
| Uric acid                     | 4.25 ± 0.97            | 4.28 ± 0.96         | 0.063       |
| TC, mg/dL                     | 199.56 ± 35.51         | 199.13 ± 35.60      | 0.428       |
| HDL-C, mg/dL                  | 56.92 ± 12.74          | 57.14 ± 12.68       | 0.259       |
| TG, mg/dL                     | 115.60 ± 76.05         | 113.51 ± 75.66      | 0.070       |
| hsCRP                         | 0.13 ± 0.37            | 0.14 ± 0.33         | 0.784       |

Data are presented as mean ± standard error or n (%).

WC waist circumference; HC hip circumference; PR pulse rate; BP blood pressure; FBS fasting blood sugar; GGT γ-glutamyl transferase; AST aspartate aminotransferase; ALT alanine aminotransferase; BUN blood urea nitrogen; TC total cholesterol; HDL-C high-density lipoprotein cholesterol; TG total triglyceride; hsCRP high sensitivity C-reactive protein.
Table 3 presents the results of the multiple regression analysis from the model used to investigate the relationship between previous APOs and chronic diseases. After adjusting for confounding factors, women with composite APOs showed higher rates of hypertension (aOR, 1.093; 95% CI, 1.004–1.189), diabetes mellitus (aOR, 1.379; 95% CI, 1.207–1.575), hyperlipidemia (aOR, 1.269; 95% CI, 1.126–1.429), stroke (aOR, 1.414; 95% CI, 1.083–1.847), and angina pectoris (aOR, 1.351; 95% CI, 1.090–1.675) than those without APOs. Transient cerebral ischemia was not associated with composite APOs. Women with preeclampsia only had a higher risk of hypertension, hyperlipidemia, and angina pectoris. Women with GDM only had a higher risk of hypertension, diabetes mellitus, and hyperlipidemia. The risk estimate of GDM for future diabetes mellitus was the highest (aOR, 9.069; 95% CI, 5.938–13.851). In women with a combination of preeclampsia and GDM, the association was significant only with diabetes mellitus (aOR, 3.287; 95% CI, 1.473–7.338), and the association was not additive. Maternal risk of chronic diseases was not associated with stillbirth or LBW among the APOs. Macrosomia was associated with a higher risk of diabetes mellitus and hyperlipidemia, a trend that was similar to that of GDM.
Table 3
Logistic regression analysis for APOs for predicting chronic disease

| APOs                  | Hypertension aOR (95% CI) | Diabetes mellitus aOR (95% CI) | Hyperlipidemia aOR (95% CI) | Transient ischemia aOR (95% CI) | Stroke aOR (95% CI) | Angina pectoris aOR (95% CI) |
|-----------------------|---------------------------|-------------------------------|-----------------------------|--------------------------------|-------------------|-----------------------------|
| Composite APOs        | 1.093 (1.004, 1.189)      | 1.379 (1.207, 1.575)          | 1.269 (1.126, 1.429)        | 0.685 (0.289, 1.622)            | 1.414 (1.083, 1.847) | 1.351 (1.090, 1.675)        |
| Preeclampsia          | 1.394 (1.215, 1.599)      | 0.988 (0.775, 1.260)          | 1.194 (0.976, 1.461)        | 1.324 (0.406, 4.318)            | 1.313 (0.842, 2.048) | 1.429 (1.012, 2.019)        |
| GDM                   | 1.417 (0.974, 2.061)      | 6.640 (4.549, 9.693)          | 1.665 (1.024, 2.707)        | -                              | 1.092 (0.266, 4.477) | 0.671 (0.164, 2.739)        |
| Stillbirth            | 0.944 (0.680, 1.310)      | 0.953 (0.563, 1.613)          | 0.725 (0.426, 1.234)        | -                              | 1.597 (0.700, 3.645) | 0.753 (0.306, 1.851)        |
| Macrosomia            | 0.881 (0.780, 0.995)      | 1.573 (1.325, 1.867)          | 1.273 (1.083, 1.496)        | 0.250 (0.034, 1.817)            | 1.434 (1.000, 2.058) | 1.292 (0.959, 1.741)        |
| LBW                   | 1.093 (0.938, 1.273)      | 0.884 (0.667, 1.171)          | 1.200 (0.966, 1.491)        | 1.405 (0.431, 4.582)            | 1.383 (0.862, 2.218) | 1.276 (0.863, 1.886)        |
| Preeclampsia only     | 1.409 (1.227, 1.618)      | 1.111 (0.869, 1.422)          | 1.290 (1.055, 1.578)        | 1.303 (0.403, 4.214)            | 1.446 (0.933, 2.244) | 1.466 (1.036, 2.074)        |
| GDM only              | 1.593 (1.011, 2.511)      | 9.069 (5.938, 13.851)         | 2.496 (1.484, 4.198)        | 1.918 (0.469, 7.854)            | 0.566 (0.078, 4.085) |                             |
| Preeclampsia + GDM    | 1.579 (0.834, 2.991)      | 3.287 (1.473, 7.338)          | 0.559 (0.136, 2.298)        | 1.263 (0.173, 9.224)            |                  |                             |

Adjusted by age, family history (hypertension, diabetes, angina, stroke), present smoking, and present alcohol.

*aOR* adjusted odds ratio; *CI* confidence interval; *APOs* adverse pregnancy outcome; *GDM* gestational diabetes; *LBW* low birth weight.

The unadjusted basic model is described in supplementary Table S1.

We analyzed the relationship between APOs and future MS and its components (Table 4). Adjusted analysis showed that composite APOs were associated with an increased risk of MS (aOR, 1.104; 95% CI,
1.020–1.194), elevated WC (aOR, 1.172; 95% CI, 1.094–1.255), hypertension (aOR, 1.074; 95% CI, 1.006–1.147), and elevated FBS level (aOR, 1.193; 95% CI, 1.111–1.280). Low HDL and high triglyceride levels were not associated with composite APOs. The effect of preeclampsia only on MS was not statistically significant; however, the effects of GDM only and the combination of preeclampsia and GDM on MS were statistically significant (aOR, 1.908 and 1.866, respectively). The combination of APOs reduced the risk of MS rather than increasing it.
Table 4
Logistic regression analysis for APOs for predicting metabolic syndrome and its components

| Metabolic syndrome | Abdominal obesity | Elevated BP | Elevated FBS | Low HCL-C | High TG |
|--------------------|-------------------|-------------|--------------|-----------|---------|
| Composite APOs     | 1.104 (1.020, 1.194) | 1.172 (1.094, 1.255) | 1.074 (1.006, 1.147) | 1.193 (1.111, 1.280) | 1.009 (0.944, 1.078) | 1.038 (0.964, 1.118) |
| Preeclampsia       | 1.115 (0.974, 1.276) | 1.185 (1.051, 1.335) | 1.349 (1.204, 1.510) | 1.040 (0.916, 1.179) | 0.939 (0.835, 1.057) | 1.001 (0.879, 1.141) |
| GDM                | 1.814 (1.309, 2.515) | 1.300 (0.959, 1.763) | 1.372 (1.034, 1.821) | 2.512 (1.907, 3.310) | 1.210 (0.908, 1.613) | 1.281 (0.929, 1.766) |
| Stillbirth         | 1.015 (0.747, 1.379) | 1.025 (0.774, 1.356) | 1.133 (0.862, 1.488) | 1.029 (0.770, 1.376) | 0.969 (0.735, 1.276) | 1.111 (0.830, 1.487) |
| Macrosomia         | 1.098 (0.986, 1.223) | 1.350 (1.231, 1.479) | 0.888 (0.811, 0.972) | 1.297 (1.180, 1.426) | 1.042 (0.952, 1.141) | 1.001 (0.903, 1.109) |
| LBW                | 0.956 (0.824, 1.108) | 0.824 (0.721, 0.941) | 1.060 (0.942, 1.193) | 0.926 (0.809, 1.059) | 0.973 (0.862, 1.099) | 1.090 (0.954, 1.245) |
| Preeclampsia only  | 1.124 (0.980, 1.289) | 1.176 (1.042, 1.328) | 1.367 (1.219, 1.532) | 1.076 (0.947, 1.223) | 0.955 (0.847, 1.076) | 1.002 (0.878, 1.144) |
| GDM only           | 1.908 (1.286, 2.831) | 1.209 (0.827, 1.765) | 1.511 (1.075, 2.125) | 3.001 (2.167, 4.157) | 1.396 (0.997, 1.954) | 1.207 (0.813, 1.791) |
| Preeclampsia + GDM | 1.866 (1.064, 3.270) | 1.817 (1.104, 2.991) | 1.532 (0.940, 2.494) | 1.848 (1.121, 3.048) | 0.810 (0.469, 1.397) | 1.468 (0.857, 2.516) |

Adjusted by age, family history (hypertension, diabetes, angina, stroke), present smoking, and present alcohol.

BP blood pressure; FBS fasting blood sugar; HDL-C high-density lipoprotein cholesterol; TG triglyceride; aOR adjusted odds ratio; CI confidence interval; APO adverse pregnancy outcome; GDM gestational diabetes; LBW low birth weight.

The unadjusted basic model is described in supplementary Table S2.

Discussion
In this large cohort study, an association was found between APOs and subsequent risk of CVD and other cardiovascular risk factors, including MS, after adjusting for family history of chronic diseases and social behaviors. The risk of CVD and cardiovascular risk factors did not increase further when preeclampsia occurred in combination with GDM.

Similar to previous studies, our study showed that women with APOs were associated with a higher risk of CVD as well as cardiovascular risk factors, including MS. Moreover, women with APOs had a higher prevalence of CVD in the family, which is known as an important risk factor for developing CVD [23–25]. Although family history plays an important role in the development of CVD, a few large cohort studies have adjusted for family history to assess the correlation between APOs and the occurrence of CVD in later life. The increased risk of CVD attributable to family history can be caused by shared genetic, environmental, and behavioral factors [23]. The association between APOs and CVD remaining after adjustment for family history may also be intervened, in part, by maternal environmental and behavioral factors, which could be modifiable. In this study, women with APOs tended to be more obese and had bad habits for cardiometabolic health, especially smoking, alcohol consumption, and less frequent breastfeeding. A previous study also reported that people with a family history of CVD had less favorable risk factors, including physical inactivity and hypercholesterolemia [23]. In this study, we observed a modest but independent association between APOs and the occurrence of CVD in the future after adjusting for family history and social behaviors. Therefore, pregnancy could be a new opportunity to inform women at risk in advance of danger signals and to prevent the occurrence of CVD through lifestyle modifications.

Among the various APOs, preeclampsia and GDM are the most studied to clarify the relationship with chronic diseases. During pregnancy, GDM increases the risk of preeclampsia [26]. However, the combined effect of preeclampsia and GDM on maternal chronic diseases after delivery has not been extensively studied. Only two studies have investigated the combined effects of GDM and preeclampsia on chronic diseases. One study in Canada reported that having both APOs was associated with more than an additive effect on diabetes mellitus, hypertension, and CVD when compared with having neither GDM nor preeclampsia after six months of delivery [27]. Furthermore, a recent study with a small population that was conducted in Turkey also proved that patients with a combination of GDM and preeclampsia had a higher prevalence of CHD than patients with only GDM [28]. Unlike these previous studies, the present study failed to prove the additive effect of the combination of GDM and preeclampsia on chronic diseases. This may be due to differences in the study population and the duration of the follow-up period. One of these previous studies [27] had a limitation in that it was comprised of women with various ethnocultural backgrounds, and the other study [28] had the limitation of a small sample size. Moreover, these studies had a shorter follow-up period of CVD evaluation after delivery than that in the present study. To the best of our knowledge, the present study is the only study that has examined the long-term effect of the combination of preeclampsia and GDM on chronic diseases in a large sample.

Research regarding the long-term health status of women who deliver infants with LBW or macrosomia or have a history of stillbirth is scarce. We observed that macrosomia was associated with an increased risk
of some chronic diseases. This observation may be related to the strong association between macrosomia and GDM, as both groups showed similar results in this study. A previous nationwide Swedish study also reported similar results that the risk of CVD in women giving birth to infants with macrosomia was attenuated after adjusting for cardiometabolic risk factors [29]. Previous studies have shown various results in women who deliver infants with LBW and have a history of stillbirth. In some cohort studies, women who delivered infants with LBW were found to be approximately twice as likely to have CVD in the future [29, 30]. In a recent meta-analysis, associations for composite CVD were two-fold for stillbirth, but no association was noted for LBW or small-for-gestational age [31]. Another meta-analysis reported that women with a history of miscarriage and/or stillbirth were more likely to develop CHD, but not stroke, compared with women without a similar history [32]. These various results could be due to the intervening by genetic and racial factors. A study on the association between stillbirth and all-cause mortality according to nationality and race reported that only in women of North African origin, the adjusted hazard ratio for all-cause mortality after stillbirth had increased significantly, whereas other groups showed no statistically significant differences [33]. Another reason could be that previous studies used different definitions for “a small fetus” and “stillbirth” and had different endpoints of the study, including prevalence and mortality. Further large-scale studies on the relationship between APOs and CVD according to racial and national differences are needed.

Our study has some limitations. The greatest limitation of this study is its cross-sectional design, which does not allow the assessment of the causal pathway underlying the observed relationships. Furthermore, it is possible that there was misclassification and under-reporting of diagnosis because diagnoses of diseases were self-reported by the participants. Additionally, selection bias might have occurred. The subjects were recruited from health examination centers and hospitals located in urban areas of Korea, and only those willing to participate were enrolled; therefore, they might not be entirely representative of the Korean population. This type of screening bias has been observed in many prospective cohort studies. Finally, there was insufficient information on factors that could influence the occurrence of chronic diseases, such as the period from childbirth to the study period, changes in diet or socioeconomic environment from childbirth to the study period, and hormone therapy. No information was available regarding the specific states of pregnancy, including the severity of pregnancy complications, gestational weight gain, and postpartum weight retention. Therefore, although we could adjust for many confounders in the present study, the potential for residual confounding by environmental factors remained. Despite these limitations, our study has several strengths. To the best of our knowledge, this was the only study that had examined the long-term effect of the combination of preeclampsia and GDM on various types of chronic diseases. Moreover, this study had a large sample size, and we were able to adjust for several potential confounders, including family history and social behaviors, and assess their independent and additive effects on the risk of CVD later in life.

In conclusion, APOs moderately influenced the future development of maternal CVD and metabolic derangements, independent of family history and social behaviors. Moreover, the risk of CVD and cardiovascular risk factors did not increase further when preeclampsia occurred in combination with
GDM. Further prospective studies are needed to clarify the relationship between APOs and maternal CVD in the future.

**Declarations**

**Acknowledgments**

The authors thank all the participants of this study. In addition, we would like to thank Editage (www.editage.cn) for English language editing.

**Funding**

Not applicable

**Conflict of interest**

The authors declare that they have no conflict of interest.

**Availability of data and material**

The data in this article are obtained from public databases that are open and transparent.

**Author’s contributions**

GJC: Data collection, data analysis, manuscript writing/editing. JK: Data collection, data analysis. JYK: Data collection, data analysis. SWH: Protocol development, statistical analysis. SBL: Protocol development, statistical analysis. MJO: Data collection, data analysis. SJK: Data collection, data analysis. JES: Data analysis, manuscript writing/editing. All authors reviewed and approved the final manuscript.

**Ethical approval**

All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards. The study was approved by the Ethics Committee of Korea University (2021GR0007).

**Consent to participate**

All participants voluntarily sign an informed consent form.

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