Exploring epistatic relationships of NO biosynthesis pathway genes in susceptibility to CHD

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Aim: To assess the epistatic relationships of nitric oxide (NO) biosynthesis pathway genes in susceptibility to coronary heart disease (CHD).

Methods: A total of 2142 subjects enrolled in two case-control studies was genotyped for 7 single-nucleotide polymorphisms (SNP) within NO biosynthesis pathway genes using TaqMan assays. The association analyses were performed at both SNP and haplotype levels. Two-way SNP-SNP interactions and high-order interactions were tested using multiple unconditional logistic regression analyses and generalized multifactor dimensionality reduction (GMDR) analyses, respectively.

Results: Two alleles (rs1049255*C and rs841*A) were identified that were significantly associated with increased risk of CHD after adjusting for all confounders (OR=1.21, 95% CI: 1.06−1.39, combined \( P_{corr} =0.007 \) and OR=1.30, 95% CI 1.12−1.50, combined \( P_{corr} <0.001 \), respectively). Significant two-way SNP–SNP interactions were found between SNP rs2297518 and these two significant polymorphisms, affecting the risk of CHD (\( P<0.001 \) for both). No significant high-order interactions were identified.

Conclusion: The results suggested that two-way SNP–SNP interactions of polymorphisms within NO biosynthesis pathway genes contribute to CHD risk.

Keywords: coronary heart disease; nitric oxide; genetics; epistasis; interaction; single-nucleotide polymorphism; case control study
factor dimensionality reduction (GMDR) analysis\(^\text{[17]}\). We also systematically evaluated these approaches for their ability to predict those individuals who were affected by CHD and had any combination of two or more polymorphisms from all of the genotyped markers.

**Materials and methods**

**Study participants and CHD definition**

CHD patients (n=557) were consecutively recruited from Tongji Hospital and the Institute of Hypertension (Wuhan, China) between May 2004 and September 2008. The diagnostic criteria for CHD were at least one of the following: (1) the presence of a stenosis >50% in at least one of the major segments of the coronary arteries (right coronary artery, left circumflex, or left anterior descending arteries) on coronary angiography; (2) elevation of cardiac enzymes (troponin T, troponin I, creatine kinase-MB, aspartate aminotransferase, and glutamic pyruvic transaminase), typical ECG changes (Minnesota Code 1.1 or 1.2 in ECG) and clinical symptoms according to the World Health Organization (WHO) criteria; or (3) a documented history of coronary artery bypass graft or percutaneous coronary intervention. Subjects with congenital heart disease, cardiomyopathy, valvular disease, or renal or hepatic disease were excluded from the study. Fifty-five-seven ethnically and geographically matched controls were randomly selected either from healthy residents in the community (89.6%) or from inpatients (10.4%) with minor illnesses. All control subjects were free of cardiovascular disease and were subjected to the same exclusion criteria as the patients with CHD. All participants were asked for a detailed medical history and received a physical examination of cardiovascular systems, including evaluation of body mass index.

To confirm the credibility of the results obtained from our first study cohort described above, we obtained another CHD case-control study cohort that comprised 507 CHD patients and 502 unaffected controls recruited from the Tongji Hospital between September 2008 and February 2010 (Wuhan, China). The diagnostic criteria for CHD as well as the inclusion and exclusion criteria for qualified participants were identical to those used in our first CHD discovery sample.

All sensitive personal information was de-identified to protect patient privacy. The institutional review board of Tongji Hospital approved this study. Written informed consent was obtained from all participants. Experiments were conducted according to the principles expressed in the Declaration of Helsinki.

**Selection of candidate genes and polymorphisms**

With regard to the regulation of the NO biosynthesis pathway, we selected seven single-nucleotide polymorphisms (SNPs) based on previous evidence of potential functionality, validated allele frequency, and sequence-proven allelic variation: Leu608Ser (rs2297518) in nNOS\(^{[18, 19]}\), G-84A (rs41279104) in the promoter region of eNOS\(^{[20]}\), Glu298Asp (rs1799983) and T-786C (rs2002744) in the promoter region of eNOS\(^{[21, 22]}\), Tyr72His (rs4673) and C+640T (rs1049255) in the 3'-untranslated region (UTR) of the cytochrome b-245, alpha polypeptide gene (CYBA)\(^{[23-26]}\), which encodes the p22 phox subunit of the NADPH oxidase, and G+243A (rs841) in the 3'-UTR of the GTP cyclohydrolase 1 gene (GCH1)\(^{[27]}\).

**DNA isolation and genotyping**

Genomic DNA was isolated from whole blood collected in K\(_2\)-EDTA tubes using the QG-Mini80 workflow with a DB-S kit (FUJIFILM Corporation, Tokyo, Japan) according to the manufacturer’s instructions. DNA was quantified and diluted to a final concentration of 10 ng/µL.

All samples were genotyped using the Taqman™ 7900HT Sequence Detection System (Applied Biosystems, Foster City, CA, USA) according to the manufacturer’s instructions. Each assay was carried out using 10 ng DNA in a 5-µL reaction consisting of TaqMan universal PCR master mix (Applied Biosystems, Foster City, CA, USA), forward and reverse primers and FAM- and VIC-labeled probes designed by Applied Biosystems (ABI Assays-on-Demand (rs1799983, C_3219460_20; rs2297518, C_11889257_10; rs1049255, C_7516913_10; rs4673, C_2038_20) and Assays-on-Design; see Table 1). Allelic discrimination was measured automatically using the Sequence Detection Systems 2.1 software (auto caller confidence level 95%). A total of 10% of all genotypes was repeated in independent PCR reactions to check for consistency and to ensure intraplate and interplate genotype quality control. No genotyping discrepancies were detected between the repeated samples. DNA samples for cases and controls were run in the same batches.

**Statistical analysis**

Statistical analyses were performed with SPSS 13.0 (SPSS Inc, Illinois, Chicago) for Windows (Microsoft Corp, Redmond, Washington) and SNPassoc for the R statistical package\(^{[28]}\).

### Table 1. TaqMan primer and probe sequences.

| Primer (5’–3’) | Allele | Allelic probe |
|---------------|--------|----------------|
| rs41279104 Forward AGGCCGACGACTGG | G | VIC-CAGAGGCCGCTTCCA-NFQ |
| reverse CCCCCTGCCACAGCCTT | A | FAM-CAGAGGCCCTTCCA-NFQ |
| rs841 Forward TTGGTGGCAATATAAGGAACCTGCAA | G | VIC-CTTGTCAGTACTTAC-NFQ |
| reverse CAGGGCTCTGGTATCTG | A | FAM-CTTGTCAGTACTTAC-NFQ |
| rs2070744 Forward ACCAGGGATCAAGCTTTC | G | VIC-AGGGTCAGCGGAGCAG-NFQ |
| reverse GCAGGTCAGAGAGACTAG | A | FAM-AGGGTCAGCGGAGCAG-NFQ |
The level of linkage disequilibrium is indicated here as D’ (D-Prime). The presence of Hardy-Weinberg equilibrium (HWE) for each SNP was tested using Haplovie 4.0[29], which is based on the χ² test.

The normality of the distribution of quantitative variables was assessed using the one-sample Kolmogorov-Smirnov test, and transformations were applied for non-normal variables when necessary. All quantitative variables were generally described as means with standard deviations (SDs). For comparison of the baseline characteristics among different groups, one-way ANOVA tests were performed on quantitative variables, such as age, body mass index, high-density lipoprotein cholesterol (HDLC), and total cholesterol (TC). The χ² test was used for qualitative variables.

For each SNP, differences in allelic frequencies between cases and controls were determined by the χ² or Fisher’s exact test. Multiple unconditional logistic regression was used to estimate odds ratios (ORs) and 95% confidence intervals (CIs) under the additive model after adjusting for covariates such as gender, age, body mass index, hypertension, diabetes, hyperlipidemia, smoking status and different populations. Two-way SNP-SNP interactions between polymorphisms within the NO biosynthesis pathway genes for CHD were also determined by multiple unconditional logistic regression using allied genotypes after adjusting for significant confounders.

Haplotype frequencies for various SNP combinations were first estimated by haplo.stats[29] (version 1.2.1) in the R statistical package and then verified using Haplovie 4.0. Each of these programs applies the Expectation-Maximization (EM) algorithm when constructing the haplotypes. The haplo.stats program was used to compute global scores and haplotype-specific P values while allowing for adjusting covariates under the additive model using default settings. To minimize the false-positive results generated from multiple statistical testing in our aforementioned analyses, we adopted the Bonferroni correction method for multiple testing.

The GMDR approach[17], which is an extension of multifactor dimensionality reduction (MDR)[31] for adjustments with covariates, was applied to identify multi-locus genetic interactions. It computed the maximum-likelihood estimates and the score values of all individuals under the null hypothesis. The cumulative score values were then calculated within each multifactor cell, which were each then labeled either as high-risk if the average score met or exceeded a pre-assigned threshold of 0 or as low-risk if the score was less than 0. We performed an exhaustive search for all possible combinations of one- to seven-locus models for all polymorphisms. The 10-fold cross validation (CV) consistency and the balanced prediction accuracy estimates were calculated for each combination of a pool of polymorphisms. The best model is the one with the highest prediction accuracy and maximal CV. The sensitivity, specificity, odds ratios, and sign tests (to determine P values) of the best model were also calculated using the GMDR software.

In CHD patients, population-attributable risks (PARs) were calculated by using the formula[32] PAR (%) = p(OR-1)/[p(OR-1)+1]×100%, where p is the proportion of individuals exposed to a risk gene (proportion or allele frequency of risk allele in CHD patients), and OR is the combined OR when cases and controls are compared in the risk model.

Power calculations to detect genetic associations were estimated using the QUANTO[33] program (Version 1.2.3). Assuming disease prevalence between 0.5% and 1%, our combined sample size can reach >80% power to detect a susceptibility locus with a genotypic relative risk >1.21 at the nominal Type I error rate of <0.05 for SNPs with minor allele frequencies >0.31 under the additive model. This indicates that our cohort sample size is sufficient to generate robust estimates in association analyses.

Results
Association between individual SNPs and the risk of CHD
The demographic details of two case-control studies are shown in Table 2. The observed frequencies of minor alleles and the genotype distributions of seven selected SNPs in cases and controls are summarized in Table 3. For all polymorphisms, the genotype distributions in cases and controls were under Hardy-Weinberg equilibrium (P>0.05). The Pallel Values for each SNP are shown in Table 3. Two SNPs (rs1049255 and rs841) were statistically significant associated with increased risk of CHD in the first study and were replicated in the second study. Combined analysis of the two studies, comprising a total of 1083 cases and 1059 control subjects, showed even stronger associations between CHD and these two SNPs (combined Pallel=0.001 for rs1049255, and combined Pallel<0.001 for rs841). Furthermore, these associations remained significant after the Bonferroni correction was applied (Pcorr=0.007 for rs1049255, and Pcorr<0.001 for rs841). We further conducted a genotypic association analysis assuming the additive genetic model to investigate how each of these SNPs confers a genetic risk for CHD. We found that same SNPs (rs1049255 and rs841) exerted effects on the disease trait in the additive models after adjusting for covariates such as gender, age, body mass index, hypertension, diabetes, hyperlipidemia, smoking status and different populations (OR=1.21, 95% CI 1.06–1.39, P=0.001, Pcorr=0.007 and OR=1.30, 95% CI 1.12–1.50, P<0.001, Pcorr<0.001, respectively; see Table 4). The overall PAR for rs1049255 and rs841 in the CHD patients were 11.52% (95%CI: 3.58%–19.47%) and 10.22% (95%CI: 4.36%–15.95%), respectively.

Haplotype analysis
We subsequently performed haplotype analysis using haplo.stats to study multiple SNPs within the eNOS and CYBA genes (Table 5). Consistent with the single-locus results (rs841 within CYBA), we only observed significant haplotype results in the CYBA gene (global P=0.01, Pcorr=0.04). To further evaluate the observed genetic effects independent of environmental factors, we conducted haplotype-based hypothesis tests using the software haplo.stats, which allowed for the adjustment of conventional risk factors. When the haplotype CG was chosen as the baseline, haplotype TG displayed a significantly increased risk for CHD (GT vs GC, OR=0.97, 95% CI 0.94–1.00,
Two-way SNP-SNP interactions

After adjusting for all covariates, significant interactions between rs2297518 and either rs1049255 or rs841 for risk of CHD were identified by the logistic regression analyses ($P<0.001$ for both). Individuals with the rs2297518 polymorphism (GA or AA) and the rs1049255 polymorphism (CT or TT) had a significantly lower risk of CHD ($OR=0.49$; 95% CI 0.38 to 0.67). In contrast, individuals with the rs2297518 polymorphism (GA or AA) and the rs841 polymorphism GG had a significantly higher risk of CHD ($OR=1.79$; 95% CI 1.33 to 2.36).

Table 2. Clinical characteristics of two case-control studies.

| Characteristics | Controls | First study | CHD | Controls | Second study | CHD |
|-----------------|----------|------------|-----|----------|-------------|-----|
| $n$             | 557      | 576        | 502 | 507      |              |     |
| Age, y          | 62.3±9.3 | 61.0±9.8$^a$ | 62.6±8.7 | 60.3±10.6$^a$ |              |     |
| Men, %          | 62.1     | 63.9       | 52.2 | 76.5$^a$ |              |     |
| BMI, kg/m$^2$   | 23.7±3.2 | 24.4±3.1$^c$ | 23.6±3.8 | 24.6±3.3$^c$ |              |     |
| SBP, mmHg       | 131.3±20.8 | 145.9±24.2$^a$ | 123.6±17.7 | 134.4±21.6$^a$ |              |     |
| DBP, mmHg       | 78.0±11.1 | 86.2±13.3$^c$ | 77.9±10.1 | 80.1±13.7$^c$ |              |     |
| HDL-C, mmol/L   | 1.33±0.36 | 1.03±0.68$^a$ | 1.29±0.53 | 1.18±0.37$^a$ |              |     |
| TC, mmol/L      | 4.56±1.70 | 4.54±1.27     | 3.12±1.67 | 4.86±1.16$^a$ |              |     |
| Hypertension, % | 19.2     | 30.4$^c$    | 37.8 | 60.9$^c$ |              |     |
| Diabetes, %     | 3.2      | 18.7$^c$    | 7.8  | 16.6$^c$ |              |     |
| Hyperlipidemia, %| 21       | 35.1$^c$    | 30.1 | 40.2$^c$ |              |     |
| Smokers, %      | 37.3     | 46.5$^c$    | 26.3 | 57.4$^c$ |              |     |

$n$ indicates number of individuals; BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure; HDL-C, high-density lipoprotein cholesterol; TC, total cholesterol; Values are expressed as mean±SD unless otherwise noted; test for differences between cases and controls, $^aP<0.05, ^cP<0.01$.

Table 3. Distribution of genotypes and alleles in patients with CHD and controls.

| ID  | Gene   | SNP rs ID (M>m) | Function | Population | Control MM/MM/m | MAF     | CHD MM/MM/m | MAF     | $P_{\text{allele}}$ | Combined $P_{\text{allele}}$ | Bonferroni $P_{\text{allele}}$ |
|-----|--------|----------------|----------|------------|-----------------|---------|-------------|---------|-----------------|-----------------------------|------------------------------|
| 1   | iNOS   | rs2297518      | Leu608Ser | First      | 382/162/13      | 0.169   | 416/138/18  | 0.151   | 0.281           | 0.013                       | 0.001                        |
|     |        | (G>A)          |          | Second     | 346/144/12      | 0.167   | 394/98/15   | 0.126   | 0.009           |                            |                              |
| 2   | nNOS   | rs41279104     | promoter | First      | 366/167/24      | 0.193   | 414/140/22  | 0.160   | 0.041           | 0.143                       |                              |
|     |        | (G>A)          |          | Second     | 346/138/18      | 0.173   | 343/152/12  | 0.174   | 0.988           |                            |                              |
| 3   | eNOS   | rs1799983      | Glu298Asp| First      | 446/102/9       | 0.108   | 461/107/8   | 0.107   | 0.946           | 0.451                       |                              |
|     |        | (G>T)          |          | Second     | 398/99/5        | 0.109   | 419/81/7    | 0.094   | 0.268           |                            |                              |
| 4   | eNOS   | rs2020744      | promoter | First      | 451/96/10       | 0.104   | 457/113/6   | 0.109   | 0.735           | 0.881                       |                              |
|     |        | (T>C)          |          | Second     | 405/95/2        | 0.099   | 410/95/2    | 0.098   | 0.941           |                            |                              |
| 5   | CYBA   | rs1049255      | 3'-UTR   | First      | 188/261/108     | 0.428   | 229/264/83  | 0.373   | 0.008           | 0.001                       | 0.007                        |
|     |        | (C>T)          |          | Second     | 175/221/106     | 0.431   | 193/235/79  | 0.388   | 0.046           |                            |                              |
| 6   | CYBA   | rs4673         | Try72His  | First      | 471/85/1        | 0.078   | 492/83/1    | 0.074   | 0.698           | 0.538                       |                              |
|     |        | (G>A)          |          | Second     | 426/74/2        | 0.078   | 436/70/1    | 0.071   | 0.567           |                            |                              |
| 7   | GCH1   | rs841          | 3'-UTR   | First      | 267/234/56      | 0.311   | 236/253/87  | 0.373   | 0.003           | <0.001                      | <0.001                       |
|     |        | (G>A)          |          | Second     | 219/226/57      | 0.339   | 194/231/82  | 0.390   | 0.018           |                            |                              |

M: major allele; m: minor allele; MAF: minor allele frequency; $P_{\text{allele}}$: significance of minor allele frequency differences determined by $\chi^2$ tests, CHD vs control; Combined Pallele from two case-control populations were combined using Mantel-Haenszel test.

Table 4. Association of CYBA and GCH1 variants with CHD.

| Gene | SNP     | Risk allele | Other allele | Adjusted OR (95% CI per risk allele) | Adjusted OR | $P$ value |
|------|---------|-------------|--------------|-------------------------------------|-------------|-----------|
| CYBA | rs1049255 | C           | T            | 1.21 (1.06–1.39)         | 0.006       |
|      | rs841   | A           | G            | 1.30 (1.12–1.50)         | <0.001      |

Significance of adjusted OR computed with multivariate unconditional logistic regression analysis, adjusting for age, body mass index, hypertension, diabetes, hyperlipidemia, smoking status and different populations.
2.40). All of the above results remained significant after correcting for multiple testing (Table 6).

**Table 5. Assessment of association between haplotypes with CHD.**

| Haplo-type | Frequency of Haplotype<sup>a</sup> | Control, n (%) | CHD, n (%) | Crude<sup>b</sup> ORs (95% CI) | Adjusted<sup>c</sup> ORs (95% CI) | Adjusted<sup>d</sup> P value<sup>e</sup> |
|------------|-----------------------------------|----------------|------------|-------------------------------|----------------------------------|---------------------------------------------|
| eNOS rs20207444-rs1799983 |                     |               |            |                               |                                  |                                            |
| GT        | 0.792                            | 0.800         | 0.616      | Baseline                      |                                  |                                            |
| GC        | 0.099                            | 0.099         | 0.883      | 1.00 (0.95–1.14)              | 0.903                            |                                            |
| TT        | 0.106                            | 0.097         | 0.385      | 0.98 (0.94–1.16)              | 0.471                            |                                            |
| Over all χ²=1.26 Global P=0.739  |                     |               |            |                               |                                  |                                            |
| CYBA rs4673-rs1049255 |                     |               |            |                               |                                  |                                            |
| GC        | 0.560                            | 0.612         | 0.001      | Baseline                      |                                  |                                            |
| GT        | 0.361                            | 0.315         | 0.002      | 0.97 (0.94–1.00)              | 0.012                            |                                            |
| AT        | 0.068                            | 0.064         | 0.531      | 0.96 (0.91–1.02)              | 0.167                            |                                            |
| Overall χ²=11.33 Global P=0.010  |                     |               |            |                               |                                  |                                            |

<sup>a</sup> Haplotype frequencies were inferred using the EM algorithm within the haplo.stats R package. Haplotypes are not listed if all the estimated frequencies are <0.02 in controls, patients with CHD;
<sup>b</sup> Crude P value based on haplotype-specific score tests;
<sup>c</sup> Odds ratio (95% confidence interval) were adjusted for gender, age, body mass index, hypertension, diabetes, hyperlipidemia smoking status and different populations.

### High-order interactions

To detect high-order SNP-SNP interactions, GMDR analyses were performed. The results are presented in Table 7. No significant high-order interactions were detected. However, the combination of rs2297518 and rs1049255 was the strongest among all two-factor models, indicating that some potential interaction exists between rs2297518 and rs1049255, which is consistent with our previous results.

**Table 7. Multilocus interaction model by GMDR method.**

| No of Loci | Best model | Prediction accuracy | Cross-validation consistency | Sign test (P) |
|------------|-------------|---------------------|-----------------------------|--------------|
| 1          | 5           | 52.13               | 8/10                        | 0.055        |
| 2          | 1, 5        | 51.80               | 6/10                        | 0.172        |
| 3          | 1, 4, 5     | 52.40               | 7/10                        | 0.623        |
| 4          | 1, 2, 5, 7  | 52.82               | 10/10                       | 0.172        |
| 5          | 1, 2, 4, 5, 7 | 51.31             | 8/10                        | 0.172        |
| 6          | 1, 2, 3, 4, 5, 7 | 51.18         | 10/10                       | 0.623        |
| 7          | 1, 2, 3, 4, 5, 6, 7 | 50.39       | 10/10                       | 0.828        |

1=rs2297518, 2=rs41279104, 3=rs1799983, 4=rs2020744, 5=rs1049255, 6=rs4673, 7=rs841.

One possible explanation for the single-locus analysis is that polymorphisms from these genes are all linked to NO production, which might be associated with the principal pathogenesis process of atherosclerosis. The C+640T polymorphism in the 3'-UTR of CYBA and the G+243A polymorphism (rs841) in the 3'-UTR of GCH1 were independently associated with an elevated risk of CHD in a Chinese Han population. Two-way SNP-SNP interaction analyses indicated that the iNOS Leu608Ser polymorphism (rs2297518) has an interaction with the two SNPs mentioned above for risk of CHD. We did not detect any high-order interactions between these SNPs.

### Discussion

In the present study, we examined the relationship between seven genetic polymorphisms within NO biosynthesis pathway genes and the risk of CHD. Single-locus analysis revealed that the C+640T polymorphism (rs1049255) in the 3'-UTR of CYBA and the G+243A polymorphism (rs841) in the 3'-UTR of GCH1 were independently associated with an elevated risk of CHD in a Chinese Han population. Two-way SNP-SNP interaction analyses indicated that the iNOS Leu608Ser polymorphism (rs2297518) has an interaction with the two SNPs mentioned above for risk of CHD. We did not detect any high-order interactions between these SNPs.

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sion has been localized to vascular smooth muscle cells and mononuclear leukocytes in early and advanced atherosclerotic lesions, which may contribute to lesion formation by increasing oxidative stress in the vessel wall[34]. Given that all of these loss-of-function polymorphisms are functionally involved in the development of atherosclerosis, they may confer risk for CHD though biological interactions with each other. Further in vitro studies are required to identify the molecular and cellular mechanisms that underlie the precise effects of these variants on endothelial function and their interaction with orthodox cardiovascular risk factors.

To control for potential false-positive results, we took several factors into consideration and carefully designed our study. First, we recruited only ethnically and geographically matched subjects from Chinese Han cohorts. Given the homogenous study population, we expect population substructure to be minimal. Second, all selected candidate SNPs have substantial functional effects, which are likely involved in the development of CHD. Third, we used the conservative Bonferroni correction to control for false-positive findings due to multiple testing. Finally, successful replication of the association signals in two independent cohorts as well as in the combined sample warrant the plausibility of our study.

Several limitations of our study should be acknowledged. First, only a limited number of genes and SNPs from the NO biosynthesis pathway were selected. The incomplete gene and SNP coverage likely does not represent the entire pathway and therefore may not fully describe the contributions of these genes. However, our report provides insight for future studies, which will focus on the elucidation of the mechanisms of these interactions. Second, population stratification may exist in this study and thus result in a spurious association between a marker and disease. Finally, although we observed an increasing statistical power in most cases when we performed analyses using the combined sample, we noticed that we failed to detect the associations of some SNPs (eg, rs4673) with CHD due to their low frequencies of minor alleles, which results in inadequate statistical power. Further validations from larger, independent populations as well as perspective studies are necessary to confirm our results.

In summary, we explored the epistatic relationships of NO biosynthesis pathway genes and CHD susceptibility in two independent case-control cohorts from a Chinese Han population. We have revealed possible interactions between SNPs and a risk for CHD. These results support the hypothesis that common polymorphisms within NO biosynthesis pathway genes modify CHD risk.

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**Author contribution**

Yuan-chao TU, Hu DING, Yu-jun XU, and Lan ZHANG performed the research; Xiao-jing WANG and Cong-xin HUANG contributed new analytical tools and reagents; Yuan-chao TU and Hu DING analyzed the data; Yuan-chao TU and Dao-wen WANG wrote the paper.

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