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RESEARCH ARTICLE



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A genome-wide association study identifies common variants influencing serum uric acid concentrations in a Chinese population

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Abstract

Background: Uric acid (UA) is a complex phenotype influenced by both genetic and environmental factors as well as their interactions. Current genome-wide association studies (GWASs) have identified a variety of genetic determinants of UA in Europeans; however, such studies in Asians, especially in Chinese populations remain limited.

Methods: A two-stage GWAS was performed to identify single nucleotide polymorphisms (SNPs) that were associated with serum uric acid (UA) in a Chinese population of 12,281 participants (GWAS discovery stage included 1452 participants from the Dongfeng-Tongji cohort (DFTJ-cohort) and 1999 participants from the Fangchenggang Area Male Health and Examination Survey (FAMHES). The validation stage included another independent 8830 individuals from the DFTJ-cohort). Affymetrix Genome-Wide Human SNP Array 6.0 chips and Illumina Omni-Express platform were used for genotyping for DFTJ-cohort and FAMHES, respectively. Gene-environment interactions on serum UA levels were further explored in 10,282 participants from the DFTJ-cohort.

Results: Briefly, we identified two previously reported UA loci of *SLC2A9* (rs1172228, combined $P = 8.98 \times 10^{-31}$) and *ABCG2* (rs2231142, combined $P = 3.34 \times 10^{-42}$). The two independent SNPs rs1172228 and rs2231142 explained 1.03% and 1.09% of the total variation of UA levels, respectively. Heterogeneity was observed across different populations. More importantly, both independent SNPs rs1172228 and rs2231142 were nominally significantly interacted with gender on serum UA levels (*P* for interaction = 4.0×10^{-2} and 2.0×10^{-2} , respectively). The minor allele (T) for rs1172228 in *SLC2A9* has greater influence in elevating serum UA levels in females compared to males and the minor allele (T) of rs2231142 in *ABCG2* had stronger effects on serum UA levels in males than that in females.

Conclusions: Two genetic loci (*SLC2A9* and *ABCG2*) were confirmed to be associated with serum UA concentration. These findings strongly support the evidence that *SLC2A9* and *ABCG2* function in UA metabolism across human populations. Furthermore, we observed these associations are modified by gender.

Keywords: Genome-wide association study, Serum uric acid, Ethnic differences, Gene-environment interaction

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Background

Uric acid (UA) is the primary end-product of purine metabolism in human beings. Most of the UA is derived from the metabolism of endogenous purine including cell turnover and synthesis. The UA excretion and reabsorption is mostly in kidney [1,2]. The UA concentration in human blood is more than fifty times higher than that in other mammals, because in most of the animals UA could be further catalyzed to allantoin by urate oxidase or uricase (the copper-binding enzyme) [1,3]; however, human beings lack uricase and have higher UA levels which could result in hyperuricemia and gouty arthritis [4]. UA could serve as an antioxidant by removing singlet oxygen and radicals [5]. However, elevated UA concentration can lead to a variety of disorders, including gout, hypertension, metabolic syndrome, diabetes mellitus, and cardiovascular disease [6-10].

It is indicated that the conventional factors including age, body mass index (BMI), alcohol consumption, and cigarette smoking could influence serum UA concentrations [11-16]. In addition, serum UA levels were also determined by genetic factors with heritability ranged from 25% to 63% [17-19]. Recent GWASs have identified multiple loci of ABCG2, SLC2A9, SLC17A1, SLC16A9, SLC22A11, GCKR, PDZK1, GCKR, RREB1, LRRC16A, WDR1, TRIM46, INHBB, SFMBT1, TMEM171, VEGFA, BAZ1B, PRKAG2, STC1, HNF4G, A1CF, ATXN2, UBE2Q2, IGF1R, NFAT5, MAF, HLF, ACVR1B-ACVRL1, and B3GNT4 associated with UA levels in European [20-26]. However, only two GWA studies have been conducted in Asians [26,27]. In addition, previous study indicated that the serum UA concentration was influenced by geneenvironmental interactions [22]. In order to investigate the genetic determines of serum UA in Asians, especially in Chinese, we conducted a GWA study in 3,451 individuals, and subsequently replicated the top SNPs in additional 8,830 healthy participants. In addition, we further examined whether the top SNPs interacted with gender, BMI, alcohol drinking, and cigarette smoking in determining serum UA levels respectively.

Methods

Study participants

In the discovery stage, we performed a GWAS of two studies in the Chinese Han population: the DFTJ-cohort consisted of 1,461 healthy individuals and the FAMHES included 2,012 Han healthy individuals aged 20 to 69 years old. The DFTJ cohort [28] and the FAMHES [29] were described in detail elsewhere. All of the participants included in the GWAS stage were recruited at health check-ups without chronic diseases such as cardiovascular disease and cancer. Briefly, the DFTJ-cohort initiated in 2008 is a long-term prospective, populationbased cohort study designed to determine the geneenvironmental interaction on several chronic diseases (obesity, diabetes mellitus, cardiovascular disease, etc.) and cancer in employees from Dongfeng Motor Corporation (DMC). The FAMHES was launched in 2009 in Fangchenggang city, Guangxi, southwest China in 2009 and enrolled 4,303 Chinese men with 17 to 88 years-old; this study was designed to examine the genetic, environmental, and their interactions on the development of age-related chronic diseases. The 8,830 healthy individuals included in the validation stage were selected from the DFTJ-cohort excluding the initial 1,461 subjects and had no diagnosed chronic diseases such as cardiovascular disease, cancer, and gout et al.

The detailed information about the GWAS population and the replication samples is shown in (Additional file 1: Table S1). All the participants provided written informed consent and the ethical committees in the Tongji Medical College and Guangxi Medical University approved this research project.

Measurement of serum UA levels and the covariates

A baseline physical examination was conducted and demographic information was collected via standard questionnaire. Overnight fasting venous blood specimens were obtained and serum UA levels were measured by the ARCHITECT Ci8200 automatic analyzer (ABBOTT Laboratories. Abbott Park, Illinois, U.S.A) using the Abbott Diagnostics reagents following the manufacturer's instructions in the DFTJ cohort [28] and using automatic analyzer (Dade Behring, USA) with original reagents in the FAMHES study. Weight and standing height were measured with light indoor clothing and in bare feet. Those who had smoked at least one cigarette per day for more than half a year either currently or formerly were defined as smokers; otherwise they were viewed as non-smokers. Alcohol drinking was divided into two categories: drinkers and non-drinkers. Those who had drunk at least once a week for more than half a year either currently or formerly were defined as drinkers; otherwise they were viewed as nondrinkers.

Sample genotyping and quality control

We performed the GWAS scan in 1,461 subjects from the DFTJ-cohort using Affymetrix Genome-Wide Human SNP Array 6.0 chips following the manufacture's protocol. Totally, we genotyped 906,703 SNPs among 1,461 subjects. After stringent QC filtering individuals with genotyping call rate < 95% were excluded (9 subjects) for further analysis. SNPs were excluded when 1) MAF < 0.01; 2) Hardy-Weinberg Equilibrium (HWE) test *P*-value < 0.0001; 3) SNPs call rate < 95%. Finally, 658,288 SNPs in 1,452 subjects with an overall call rate of 99.68% were used for further analysis. We used the Illumina Omni-Express platform to carry out the GWAS scan in FAMHES. There were 1,999 individuals (sample call rate >95%) included in the final statistical analysis. Based on quality control criteria, SNPs were excluded when P < 0.001 for the HWE test, MAF < 0.01, or genotype call rate < 95%. Finally, 709,211 SNPs were kept for further analysis.

In the validation stage, ten SNPs were selected based on the following criteria: 1) SNP with $P < 1.0 \times 10^{-5}$ for all GWAS samples; 2) when multiple SNPs showed a strong LD ($r^2 \ge 0.8$), SNPs previously reported in the literature were prior selected; 3) Clear genotyping clusters; 4) MAF ≥ 0.05 . We used the iPLEX system (Sequenom) and/or the TaqMan assay (Applied Biosystems) [30,31] to genotype the 10 SNPs. The primers and probes were available upon request.

Imputation

We performed ungenotyped SNPs imputation using MACH 1.0 software (see URLs) via LD information from the HapMap phase II database (CHB + JPT as reference set, 2007-08_rel22, released 2007-03-02) in the DFTJ-cohort GWAS. Genotyped SNPs in the FAMHES GWAS were inferred using the IMPUTE program [32], and the reference panels used for imputation in the FAMHES study were HapMap rel.24, build 36, CHB + JPT. The Imputed SNPs with high genotype information content (proper info > 0.5 for IMPUTE and Rsq > 0.3 for MACH) were retained for the further association analysis. Finally, 2,468,160 SNPs were used for the further analysis.

Statistical analysis

Analysis for serum UA was performed on natural logarithmic (ln)-transformed values because of skewed distributions. Genome-wide association tests were performed using the additive model by linear regression analysis with adjustment for age, gender, BMI, cigarette smoking, and alcohol drinking implement using PLINK1.06 [33]. The top two eigenvectors were also adjusted as covariates in the linear regression analysis. Population structure was evaluated using principal components analysis (PCA), as implemented by EIGENSTRAT software [34] and quantile-quantile (QQ) plot was generated by using R 2.11.1 (see URLs). Heterogeneity among the study populations was evaluated by the I^2 statistic [35]. The Manhattan plot of -log10 P, LD structures and haplotype block plots were generated by using Haploview (v4.1) [36]. The association studies with the imputation data were performed using the ProbABEL software [37]. The meta-analysis of the DFTJ-cohort GWAS data and the FAMHES GWAS data was performed using a fixedeffects meta-analysis with inverse variance weighted method using the metal software [38]. The regional association plots were drawing using SNAP software [39].

The power of the present study was calculated using the Quanto software package [40]. We used the mean of serum UA 292.5 μ mol/L, MAF values obtained from the combined genotype dataset, and assumed an additive genetic model with $\alpha = 0.05$ in two tail tests to calculate the statistical power.

A conditional analysis was done in linear regression model to examine the independence of the four associated SNPs in the combined data (partial r^2 indicated the proportion of the serum UA variation explained by each SNP). We did the conditional analysis via including the four significant SNPs in the linear regression model adjusting for age, gender, BMI, cigarette smoking, and alcohol drinking in 10,282 individuals from the DFTJ-cohort. The SNPs with *P* value < 0.05 remained in the multivariate model were considered to be associated with the serum UA levels independently.

The combined dataset including the DFTJ-cohort GWAS data and the validation stage data totaling 10,282 individuals used in gene-environment interaction analysis was tested by introducing the interaction terms (SNP × gender, SNP × BMI, SNP × alcohol drinking, and SNP × cigarette smoking) into the model, adjusting for the covariates including age, gender, BMI, alcohol drinking, and cigarette smoking. The *P* value less than 0.05 for the interaction term was considered statistical significant. Considering multiple interaction tests were conducted, we further did the multiple test [41] for the gene-environmental interaction analysis. All statistical analyses were performed with SPSS (version 15.0; SPSS, Chicago, IL), and SAS version 9.2 (SAS Institute, Cary, NC, USA).

Results

Genome-wide association of serum UA levels

The demographics of the participants are displayed in (Additional file 1: Table S1). The Q-Q plot revealed no inflation of type I error rate due to population stratification, with a genomic control inflation factor of 1.007 (Additional file 2: Figure S1). No heterogeneity was observed for the SNPs presented in the Table 1 between the DFTJ-cohort study and the FAMHES. As the Manhattan plot (Figure 1) and the regional association plots (Additional file 3: Figure S2) indicated, at the discovery stage the *ABCG2* was significantly associated with the serum UA at a genome-wide significance level.

Totally we further investigated ten SNPs in the validation stage and four common variants were associated with serum UA levels at a genome-wide significant level of 5×10^{-8} : *SLC2A9* (rs11722228, combined $P = 8.98 \times 10^{-31}$) and *ABCG2* (Combined $P = 3.34 \times 10^{-42}$ for rs2231142) (Table 1). Both loci of *ABCG2* and *SLC2A9* have been reported previously [20,22,27,42]. If SNPs are in the same locus, we chosen r² of SNPs less than 0.8 for further

Table 1 Genome-wide association analyses for serum uric acid levels

SNP	Chr	bp	Genes	Locus	Minor/major allele	GWAS(n = 3,451)				Validation(n = 8,830)				
						MAF	Effect size (s.e.m.)	<i>P</i> ₋ value ^a	MAF	Effect size (s.e.m.)	<i>P</i> ⁻ value ^a	P- value ^a		
rs11722228	4	9524839	SLC2A9	intron	T/C	0.31	0.028 (0.006)	3.68 × 10 ⁻⁶	0.31	0.043 (0.004)	4.16×10^{-26}	8.98 × 10 ⁻³¹		
rs4148152	4	89279933	ABCG2	intron	C/T	0.35	-0.029 (0.006)	1.22×10^{-7}	0.33	-0.029 (0.004)	4.49×10^{-13}	2.95 × 10 ⁻¹⁸		
rs3114018	4	89283605	ABCG2	intron	T/G	0.38	-0.034 (0.006)	2.75×10^{-9}	0.38	-0.029 (0.004)	1.02×10^{-12}	4.49×10^{-20}		
rs2231142	4	89271346	ABCG2	exon	T/G	0.29	0.046 (0.006)	1.19×10^{-14}	0.31	0.045 (0.004)	1.01×10^{-25}	3.34 × 10 ⁻⁴²		

The NCBI build 36 was used as the reference genome. Chr, chromosome. MAF indicates the minor allele frequency calculated using the data from all the subjects in the analysis. Effect size, represents the effect of a minor allele on the standardized trait (estimated coefficient of the term for the number of the minor alleles). ^aLinear regression analysis adjusted for age, sex, body mass index (BMI), cigarette smoking, and alcohol drinking.

investigating ($r^2 = 0.168$ for rs2231142 and rs4148152; $r^2 = 0.224$ for rs2231142 and rs3114018; $r^2 = 0.669$ for rs4148152 and rs3114018). (Additional file 3: Figure S2). However, we failed to replicate the top SNPs in the other loci of *SEC22B*, *CENTG2*, *TET2*, *VEGFC*, and *TNFRSF11B* in the validations stage (Additional file 4: Table S2).

The SNP rs11722228 in *SLC2A9* and rs2231142 in *ABCG2* were independently associated with serum UA levels and accounted for 1.03% and 1.09% of the serum UA variance respectively (Additional file 5: Table S3).

Ethnic differences in major genetic variants associated with serum UA levels

We compared the results in the present study with those in Japanese and Europeans. As Table 2 showes, the effect sizes of most of these loci showed consistent direction across the populations. *ABCG2* was associated with serum UA levels across different populations. We found significant association of serum UA with SNP rs4148152 in *ABCG2* in our study (combined $P = 2.95 \times 10^{-18}$). However, till now we did not find any reports of this SNP in European populations, which might be attributable to very low MAF (0.017) in Europeans. Furthermore, the SNP rs4148152 was in moderate LD with the SNP rs311408 in the present study $(r^2 = 0.669)$ but in very low LD in Europeans ($r^2 = 0.046$; HapMap CEU). SNP rs12356193 in SLC16A9, rs10480300 in PRKAG2 and rs653178 in ATXN2 were significantly associated with serum UA in European populations but were monoallelic in Asians. Similar findings were found for loci of SLC2A9 (rs16890979 and rs734553), SLC22A11 (rs17300741), and recently reported new loci of TRIM46 (rs11264341), VEGFA (rs729761), BAZ1B (rs1178977), STC1 (rs1778674), A1CF (rs10821905), UBE2Q2 (rs1394125), and HLF (rs7224610) [25] which were associated with serum UA in Europeans, however, these associations were not replicated in Asians (Japanese and Chinese in the present study). This might be due to the very lower MAF in Asians. However, the relative small sample size in the present study limited us to detect these associations with enough power. In addition, we failed to replicate the SNPs rs742132 in LRRC16A, rs780094 in GCKR, rs17632159 in TMEM171, rs17050272 in IINHBB, and rs7188445 in MAF which were reported in European [20,25], same as the findings in the Japanese population, albeit the MAFs of both of the SNPs are similar between Asians and



					Our study ^a (Chinese, discovery stage)				Japanese		European ancestry		
Gene	SNP	Chr	Position	Location	MAF (allele)	Effect size	P-value	MAF (allele)	Effect size	P-value	MAF (allele)	Effect size	P-value
PDZK1	rs12129861	1	144437046	5'-	0.20(A)	-0.011	0.074	0.087(A)	=	=	0.46(A)	-0.062 [20]	2.68 × 10 ⁻⁹
TRIM46	rs11264341	1	153418117	Intron	0.33(C)	0.017	4.49×10^{-3}	0.24(C)	0.074 [25]	9.5× 10 ⁻⁴	0.43(T)	-0.050 [25]	6.2×10^{-19}
INHBB	rs17050272	2	121022910	N/A	0.45(A)	0.008	0.174	0.49(A)	0.033 [25]	4.4× 10 ⁻²	0.43(A)	0.035 [25]	1.6×10^{-10}
GCKR	rs780094	2	27594741	Intron	0.44(T)	0.0133	0.014	0.43(T)	0.036 [27]	5.12×10^{-6}	0.42(T)	0.050 [20]	1.40×10^{-9}
LRP2	rs2544390	2	169913092	Intron	0.49(C)	-0.001	0.73	0.49(C)	-0.082 [27]	3.74×10^{-8}	0.37(T)	-	-
SFMBT1	rs6770152	3	53075254	N/A	0.36(T)	-0.010	0.071	0.46(T)	-0.016 [25]	0.31	0.42(G)	0.044 [25]	2.6×10^{-16}
SLC2A9	rs11722228	4	9524839	Intron	0.31(T)	0.028	3.68 × 10 ⁻⁶	0.45(T)	0.164 [27]	7.09 × 10 ⁻²⁴	0.50(T)	0.167 [20]	1.75 × 10 ⁻⁷⁵
SLC2A9	rs16890979	4	9531265	Exon	0.02(T)	-0.024	0.285	0.01(T)	-0.178 [27]	3.07×10^{-2}	0.29(T)	-0.340 [20]	3.55 × 10 ⁻¹⁸⁹
SLC2A9	rs734553	4	9532102	Intron	0.02(C)	-0.023	0.286	0.006(C)	-	-	0.23(C)	0.315 [20]	5.22 × 10 ⁻²⁰¹
ABCG2	rs3114018	4	89283605	Intron	0.38(T)	-0.034	2.75×10^{-9}	0.26(A)	-	-	0.48(G)	-0.057 [20]	2.93 × 10 ⁻¹²
ABCG2	rs2231142	4	89271347	Exon	0.29(T)	0.046	1.19×10 ⁻¹⁴	0.31(T)	0.121 [27]	1.62×10^{-13}	0.11(T)	0.173 [20]	3.10×10^{-26}
ABCG2	rs4148152	4	89279933	Intron	0.35(C)	-0.029	1.22×10^{-7}	0.19(C)	-	-	0.017(C)	-	-
ABCG2	rs4148155	4	89273691	Intron	0.29(A)	-0.046	9.23 × 10 ⁻¹⁵	0.30(G)	0.121 [27]	1.14 × 10 ⁻¹³	0.11(A)	-0.170 [20]	1.50×10^{-25}
TMEM171	rs17632159	5	72467238	N/A	0.31(C)	-0.009	0.129	0.28(C)	-0.045 [25]	0.011	0.31(C)	-0.039 [25]	3.5×10^{-11}
VEGFA	rs729761	6	43912549	N/A	0.14(T)	-0.013	0.092	0.07(T)	0.002 [25]	0.97	0.30(T)	-0.047 [25]	8.0×10^{-16}
LRRC16A	rs742132	6	25715550	Intron	0.25(A)	-0.001	0.8705	0.31(A)	0.008 [27]	0.609	0.30(A)	0.054 [20]	8.50×10^{-9}
SLC17A1	rs1183201	6	25931423	Intron	0.18(A)	-0.009	0.2	0.16(A)	-0.063 [27]	1.73×10^{-3}	0.48(A)	-0.062 [20]	3.04×10^{-14}
SLC17A3	rs1165205	6	25978521	Intron	0.19(T)	-0.010	0.139	0.16(T)	-0.07 [27]	5.04×10^{-4}	0.48(T)	0.060 [20]	1.60×10^{-13}
PRKAG2	rs10480300 b	7	151036938	Intron	0.00(T)	-	-	0.00(T)	-	-	0.28(T)	0.035 [25]	4.1×10^{-9}
BAZ1B	rs1178977	7	72494985	Intron	0.08(G)	0.011	0.291	0.11(G)	-0.058 [25]	0.025	0.19(G)	-0.047 [25]	1.2×10^{-12}
STC1	rs17786744	8	23832951	N/A	0.35(G)	0.0001	0.989	0.25(G)	0.018 [25]	0.32	0.42(G)	0.029 [25]	1.4×10^{-8}
HNF4G	rs2941484	8	76641323	3utr	0.32(T)	0.024	1.7× 10 ⁻⁵	0.43(T)	0.050 [25]	1.8× 10 ⁻³	0.44(T)	0.044 [25]	4.4×10^{-17}
A1CF	rs10821905	10	52316099	5'near gene	0.04(A)	0.008	0.584	0.05(A)	0.075 [25]	0.042	0.18(A)	0.057 [25]	7.4×10^{-17}
SLC16A9	rs12356193 ^b	10	61083359	Intron	0.00(C)	-	-	0.00(C)	-	-	0.18(C)	0.080 [20]	1.07 × 10 ⁻⁸
SLC22A11	rs17300741	11	64088038	Intron	0.07(A)	0.018	0.087	0.03(A)	0.063 [27]	0.197	0.49(A)	0.060 [20]	6.68 × 10 ⁻¹⁴
SLC22A12	rs505802	11	64113648	Intron	0.23(T)	-0.008	0.196	0.18(T)	-0.231 [27]	1.00×10^{-31}	0.30(T)	-0.060 [20]	2.04 × 10 ⁻⁹
SLC22A12	rs506338	11	64197496	Intron	0.23(C)	-0.005	0.502	0.17(C)	-0.229 [27]	2.34 × 10 ⁻³¹	0.29(T)	-	-
ATXN2	rs653178 b	12	110492139	Intron	0.00(C)	-	-	0.00(C)	-	-	0.49(C)	0.035 [25]	7.2 × 10 ⁻¹²
UBE2Q2	rs1394125	15	73946038	Intron	0.008(A)	0.014	0.188	0.08(A)	0.021 [25]	0.48	0.34(A)	0.043 [25]	2.5×10^{-13}
IGF1R	rs6598541	15	97088658	Intron	0.41(A)	0.015	9.85× 10 ⁻³	0.50(A)	0.033 [25]	0.038	0.36(A)	0.043 [25]	4.8×10^{-15}

Table 2 Ethnic differences in major genetic variants associated with serum uric acid levels

NFAT5	rs7193778	16	68121391	N/A	0.05(C)	0.004	0.769	0.10(C)	0.053 [25]	0.048	0.14(C)	0.046 [25]	8.2×10^{-10}
MAF	rs7188445	16	78292488	N/A	0.27(A)	-0.003	0.592	0.31(A)	-0.06 [25]	4.5× 10 ⁻⁴	0.33(A)	-0.032 [25]	1.6×10^{-9}
HLF	rs7224610	17	50719787	Intron	0.13(C)	0.004	0.579	0.18(C)	0.004 [25]	0.83	0.42(C)	0.042 [25]	5.4×10^{-17}

The NCBI build 36 was used as the reference genome. ^aSample size, 3451, the P values in Han Chinese population were from meta-analysis of two GWASs (DFTJ-cohort and FAMHES) in discovery stage; –, information is not available. ^bThese SNP is monoallelic in HapMap CHB and HapMap JPT.

Α

600

400



P_{interaction}=0.040

Europeans, suggesting that there were genetic discrepancy on serum UA levels among different ethnic groups. The moderate effect size of these loci on serum UA levels and the limited sample size in the present study might be another potential explanation.

Interaction of SNPs with gender, BMI, cigarette smoking, and alcohol drinking

We further explored the interactions between the two independently-associated SNPs (rs11722228 in SLC2A9 and rs2231142 in ABCG2) and gender, BMI, cigarette smoking, and alcohol drinking on serum UA levels in 10,282 individuals from the DFTJ-cohort. Both SNPs rs11722228 and rs2231142 were nominally interacted with gender on serum UA levels (P = 0.04 and P = 0.02,

respectively; Additional file 6: Table S4 and Figure 2). The minor allele (T) for rs11722228 in SLC2A9 has greater influence in elevating serum UA levels in females compared to males (beta = 0.051 in females vs. beta = 0.035 in males), similar with the previously reported results [20]. In contrast, for SNP rs2231142 in ABCG2, the minor allele (T) had stronger effects on serum UA levels in males than those in females (beta = 0.037 in females vs. beta = 0.057 in males), consistent with the findings in the European populations and African Americans [20,22]. However, the gene-gender interaction altered to null after the multiple test based on the FDR approach. Besides, the rs11722228 accounted for 1.33% and 0.98% and rs2231142 accounted for 0.93% and 2.25% of the total variance of the serum UA levels for females and males, respectively.

In addition, the SNP rs11722228 was also interacted with alcohol drinking and cigarette smoking (P for interaction = 0.016 and 0.035, respectively; Additional file 6: Table S4). Considering that the SNP rs11722228 had different effect size on serum UA levels in males and females and in the present study 95.1% smokers and 86.7% drinkers were males, we restricted our analysis in males. However, we failed to detect the interactions between rs11722228 and cigarette smoking and alcohol drinking on serum UA levels anymore (data not showed), suggesting that the SNP-smoking and the SNP-drinking interaction was driven by the gender differences in the serum UA concentrations.

Discussion

CC CT TT

In this two-stage GWAS, we replicated two previously reported loci (ABCG2 and SLC2A9) associated with serum UA levels and found significant gene-gender interactions in Chinese Han population. In addition, ethnic differences were observed between Asian and European populations.

SLC2A9 is located in chromosome 4p16-15.3 and encodes glucose transporter 9 (GLUT9) which can reabsorb UA in renal tubules [4]. Several studies have reported the association between SLC2A9 and serum UA levels [20,27,43,44]. Importantly, SLC2A9 is a transporter for both fructose and urate [45]. Fructose intake could facilitate UA formation in liver via increasing purine breakdown. In addition, animal evidences indicated a causal relationship between fructose intake, serum UA, and metabolic syndrome [11,46,47].

ABCG2 is an UA exporter that mediates urate excretion in the kidney. Multiple evidences indicated that the common variants in ABCG2 could reduce the transport function and result in the hyperuricemia and gout [44,48,49]. In the present study, the missense SNP rs2231142 in ABCG2 showed the strongest association with serum UA level. This missense SNP could result in a glutamine-tolysine amino acid substitution and the glutamine residue is highly conserved across species and the LD pattern differs in Chinese population and European population.

Comparisons of the SNPs of the association studies for serum UA in different populations are of great interest. The present study replicated two previously reported loci of SLC2A9 and ABCG2 associated with serum UA levels. SNP rs11722228 in SLC2A9 explained 1.03% variation compared to 1.33% in Japanese population [27] and rs2231142 in ABCG2 accounted for 1.09% of the variation of serum UA levels in our study compared to 1.20% in white individuals and 0.30% in African Americans [22]. Previous studies reported that variants in SLC2A9 show the strongest effect on serum uric acid levels compared to ABCG2 [20,22]. However, in the present study we observed that the loci of ABCG2 and SLC2A9 show comparable amounts of explained variance (1.09% and 1.03% respectively). Table 2 showed that there was a little difference in MAFs in both variants rs2231142 located in ABCG2 and rs11722228 located in SLC2A9 (0.29 and 0.31) in Chinese which were different from that observed in Europeans (0.11 and 0.50 respectively).

Owing to the SNPs of rs12356193 in SLC16A9, rs10480300 in PRKAG2, and rs653178 in ATXN2 are monoallelic in Asians, we failed to replicate the SNPs of rs12356193, rs10480300 and rs653178 those were identified in Europeans. In addition, we found notable differences in MAFs for rs16890979 in SLC2A9, rs17300741 in SLC22A11, rs10821906 in AICF, rs1394125 in UBE2Q2, rs7193778 in NFAT5 and rs7224610 in HLF (0.02, 0.07, 0.04, 0.008, 0.05, 0.13 respectively for Chinese population and 0.29, 0.49, 0.18, 0.34, 0.14, 0.42 respectively for European populations) Besides, both rs16890979 ($r^2 =$ 0.005) and rs734553 ($r^2 = 0.005$) in the gene *SLC2A9* were in very low LD with the SNP rs11722228 in Asians compared to that in European populations $(r^2 = 0.202 \text{ for})$ rs16890979; $r^2 = 0.193$ for rs734553). The very low MAF and the difference in the LD structure might partly explain the discrepancy of the associations between the Europeans and Asians. In addition, relative small sample size in the present study and the moderate effect size of the loci on serum UA levels also limited us to have enough power to detect these associations.

Serum UA levels are lower in females than that in males. More importantly, gene-gender interactions were observed for the two independent SNPs of rs11722228 and rs2231142. The minor allele (T) for rs11722228 has greater influences in elevating serum UA levels in females compared to males, consistent with the previous study [23]. For rs2231142, the TT genotype was related to higher UA as compared with GG and GT genotype in both males and females. This might be due to that the rs2231142 T allele was related to the reduced ability to excrete UA [50]. In addition, the minor allele T allele of

rs2231142 has greater effects on the serum UA levels in males than those in females (P for interaction = 0.02), in consistent with the findings from the Europeans [22]. This gender difference might be due to the specific physiological characteristics in females. It is also reported that the estrogens might increase the renal clearance of serum UA [51], however, the mechanism underlying the genegender interaction remains to be further elucidated.

To our best of knowledge, this is the first GWA study on serum UA levels in Chinese population. Because of the relative small sample size in the discovery stage, we might have limited power to detect the associations of the SNPs with small effect size and/or low MAF. However, in the present study we identified two reported loci (SLC2A9 and ABCG2) associated with serum UA levels, suggesting that our study was capable of identifying significant loci associated with serum UA levels. In addition, the combined data in our study has more than 90% statistic power to detect the interaction between SNP rs11722228 and gender; 70.4% statistical power to detect the interaction between SNP rs2231142 and gender on serum UA levels. Furthermore, our study confirmed the gene-gender interaction on serum UA levels and observed that the ABCG2 and SLC2A9 functioned differently in males and females across different populations [20].

Conclusions

Our study replicated two loci of *SLC2A9* and *ABCG2* associated with serum UA levels in a Chinese Han population. Heterogeneity was observed among different ethnic populations. In addition, we found both the two loci interacted with gender on serum UA levels. These two loci had different effects on serum UA levels in males and females. Further studies are needed to validate our findings and investigate the underlying mechanisms of the gene-gender interaction.

Additional files

Additional file 1: Table S1. Characteristics of the subjects participated in this study.

Additional file 2: Figure S1. Q-Q plots for QTL analyses.

Additional file 3: Figure S2. Associations of SNPs in Table 2 with serum uric acid concentrations.

Additional file 4: Table S2. SNPs failed to be validated in the validation stage in the present study.

Additional file 5: Table S3. Covariates and SNPs in relation to serum uric acid levels in 10,282 individuals.

Additional file 6: Table S4. Interaction between SNPs and gender, BMI, alcohol drinking and cigarette smoking.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

Projects conception: WTC, MZN, HMA, FBH. Study design: WTC, HMA, YBY, LDX, WC, YHD, ZXM, YJ, GH. Sample processing and database establishment: YBY, YXB, HYF, ZL, DQF, HSL, ZJ, MXW, LMJ, LDF, QGK, GL, HD, ZX, FYY, and WT. Genotyping: HMA, YBY, And GLX. Imputation: HMA, YBY, GLX. Data analysis: YBY, DXY, and LJ. Interpretation of the data: HMA, MZN, YBY. Drafted the manuscript: YBY and MZN. Revised manuscript: WTC, HMA, MZN. All authors have read and approved of the final version of this manuscript.

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URLs

R statistical environment, http://cran.r-project.org/; PLINK, http://pngu.mgh.harvard.edu/~purcell/plink/; MACH 1.0, http://www.sph.umich.edu/csg/abecasis/mach/; International HapMap Project, http://www.hapmap.org/index.html.

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