Genome-wide meta-analysis identifies multiple novel loci associated with serum uric acid levels in Japanese individuals

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Gout is a common arthritis caused by elevated serum uric acid (SUA) levels. Here we investigated loci influencing SUA in a genome-wide meta-analysis with 121,745 Japanese subjects. We identified 8948 variants at 36 genomic loci (P<5×10⁻⁸) including eight novel loci. Of these, missense variants of SESN2 and PNPLA3 were predicted to be damaging to the function of these proteins; another five loci—TMEM18, TM4SF4, MXD3-LMAN2, PSORS1C1-PSORS1C2, and HNF4A—are related to cell metabolism, proliferation, or oxidative stress; and the remaining locus, LINC01578, is unknown. We also identified 132 correlated genes whose expression levels are associated with SUA-increasing alleles. These genes are enriched for the UniProt transport term, suggesting the importance of transport-related genes in SUA regulation. Furthermore, trans-ethnic meta-analysis across our own meta-analysis and the Global Urate Genetics Consortium has revealed 15 more novel loci associated with SUA. Our findings provide insight into the pathogenesis, treatment, and prevention of hyperuricemia/gout.
Serum uric acid (SUA) is reported to have an antioxidative effect\(^1\)–\(^2\), whereas elevated SUA, or hyperuricemia, results in crystal deposition and causes gout\(^3\). Gout is a common disease characterized by noninfectious acute arthritis. Both gout and hyperuricemia can result from an unhealthful lifestyle\(^4\)–\(^6\), but recent genetic studies, including genome-wide association studies (GWASs), have also revealed a genetic contribution to the development of these conditions, with this contribution being larger than that for other common diseases\(^7\)–\(^12\). Moreover, epidemiologic studies have revealed their relationship among other diseases such as cardiovascular diseases\(^13\),\(^14\), indicating the demicological studies have revealed their relationship among other diseases have a broader genetic basis than other common diseases\(^1\)–\(^6\),\(^13\)–\(^14\), indicating the importance of elucidation of the pathophysiology of these conditions. To date, several GWAS of SUA have been performed with Caucasian populations\(^15\)–\(^24\) as well as Asian populations including Japanese subjects\(^25\),\(^26\). Although there are genetic differences between Caucasian and Asian populations, they have many shared associated genes\(^2\),\(^27\) that exert major effects, such as \(ABCG2\), \(SLC2A9\), and \(SLC22A12\), all of which are well-known representative urate transporters in humans and which are important as therapeutic target molecules for gout and hyperuricemia. Therefore, identifying new loci may not only help elucidate the pathophysiology of these diseases, but may also reveal their target molecules, taking into account the fact that these diseases have a broader genetic basis than other common diseases as described above. Furthermore, the gene expression patterns to which the identified loci contribute should enable us to estimate effective pathways for drug delivery. In the present study, we have investigated the genetic loci that influence SUA with more than 120,000 Japanese individuals in a genome-wide meta-analysis and have compared our findings with those of previous GWASs\(^24\),\(^28\).

We identified 36 loci for SUA, including eight previously unreported loci, that suggest key cellular processes which contribute to elevated serum uric acid levels, followed by the identification of 15 more loci by trans-ethnic meta-analysis.

Results

Genome-wide meta-analysis. We performed a genome-wide meta-analysis based on three Japanese cohorts including those of the Japan Multi-institutional Collaborative Cohort (J-MICC) Study\(^29\)–\(^30\), the Kita-Nagoya Genomic Epidemiology (KING) Study\(^31\),\(^32\), and the BioBank Japan (BJJ) Project\(^33\),\(^34\). Detailed information regarding the baseline characteristics of the study subjects, genotyping arrays, and imputation is summarized in Supplementary Tables 1 and 2. We performed a genome-wide meta-analysis for SUA with data sets encompassing 121,745 Japanese subjects. Intercepts of linkage disequilibrium (LD) score regression and the genomic control lambda for each study are shown in Supplementary Table 2. The intercepts of LD score regression and the genomic control lambda for our meta-analysis were 1.043 and 1.165, respectively. Genomic control adjustment was not applied for genomic control at the level of each study because intercepts of LD score regression did not show inflation of test statistics. The quantile–quantile (Q–Q) plot for \(P\) values is shown in Supplementary Fig. 1. The results of the meta-analysis identified 8948 variants at 36 genetic loci with a \(P\) value of \(<5 \times 10^{-8}\) for SUA (Fig. 1). Among these 36 genetic loci, 8 were not previously reported, 10 were recently identified in a GWAS for SUA in Japanese performed by BJJ\(^28\), and 18 were previously identified by other GWASs for SUA\(^15\)–\(^17\),\(^19\)–\(^20\),\(^24\)–\(^26\). The eight novel loci were the following: \(rs74896528\) of \(SESN2\), \(rs10188118\) of \(LOC105373352\) - \(TMEM18\), \(rs6774054\) of \(TM4SF4\), \(rs11952102\) of \(MXD3\)–\(LMAN2\), \(rs16898823\) of \(PSORS1C1\)–\(PSORS1C2\), \(rs8024067\) of \(LINC01578\), \(rs6031598\) of \(PNPLA3\), and \(rs2281293\) of \(PNPLA3\).

Sentinel single-nucleotide polymorphisms (SNPs) with the lowest \(P\) values for SUA at each of the 36 loci are shown in Table 1. Association results of each study are shown in Supplementary Data 1. We determined the effect allele frequencies (EAFs) of these sentinel SNPs for each population in 1000 Genomes phase 3 (Supplementary Data 2). The EAFs indicated the sentinel SNPs with the lowest \(P\) values for SUA at each of the 36 loci are shown in Fig. 1. Eighteen loci shown in Supplementary Fig. 2. The results for these 27 SNPs, identified in our meta-analysis with the recent results by BJJ\(^28\) (Supplementary Data 3), with regional association plots for the eight loci newly identified in the present study are shown in Fig. 2. The BJJ data recently revealed that SNPs located at 27 loci showed genome-wide significant associations with SUA including 10 novel loci (Table 1)\(^28\). About the 27 reported SNPs, we compared the results in our meta-analysis with the recent results by BJJ\(^28\) (Supplementary Data 3), with regional association plots for the 10 loci also identified in the present study being shown in Supplementary Figure 2. The results for these 27 SNPs, identified in our meta-analysis, revealed a higher level of significance for the association with SUA in our meta-analysis than in the BJJ study.

A European GWAS for SUA was previously performed by the Global Urate Genetics Consortium (GUGC)\(^24\). We examined the publicly available data provided by the GUGC-based study for...
the sentinel SNPs or SNPs showing high LD ($r^2$ of ≥0.8 in JPT of 1000 Genomes phase 3) with the sentinel SNPs at the eight novel loci identified in the present study. Three of these loci, including 5q35.3, 2q13.12, and 22q13.31, were significantly associated with SUA in the GUGC-based GWAS, with the same direction of effect size as in our study (Supplementary Table 3). The 2p25.3 locus was nominally significantly associated with SUA. Although the 3q25.1 locus was not significantly associated with SUA in the GUGC-based study, it was nominally significantly associated with gout in the same study.

### Functional annotations for novel loci

We searched for SNPs at the newly identified loci associated with SUA that were associated with gene expression level or amino acid substitution of protein and that were in high LD ($r^2$ of ≥0.8 in JPT of 1000 Genomes phase 3) with sentinel SNPs and had a P value of $<1 \times 10^{-6}$ for SUA in our meta-analysis. We identified two nonsynonymous SNPs of $SESN2$ at the 1p35.3 locus and $PNPLA3$ at the 22q13.31 locus (Supplementary Table 4), and we found that six of the eight novel loci harbor variants with expression quantitative trait loci (eQTLs) for at least one tissue in the Genotype-Tissue Expression (GTEx) database. The two nonsynonymous SNPs, rs738409 (I148M) of $SESN2$, and rs74896528 (P87S) of $SESN2$, were predicted by SIFT, PolyPhen2 HVAR, and PolyPhen2 HDIV to be damaging or probably damaging.

### Gene set enrichment analysis of SUA-associated loci

We searched for genes whose expression level was associated with SUA-associated SNPs in at least one tissue in the GTEx database. We found that 24 of the 36 loci identified in the present study harbor variants with eQTLs in at least one tissue in the GTEx database. We also identified 71 positively correlated genes whose expression level is increased by SUA-increasing alleles and 76 negatively correlated genes whose expression level is decreased by SUA-increasing alleles (Supplementary Data 5). Functional analysis of the sets of positively correlated genes and negatively correlated genes were performed with the Database for Annotation, Visualization, and Integrated Discovery (DAVID). For the positively correlated genes, the terms “Williams-Beuren...
Fig. 2 Regional association plots for the eight novel loci identified in the meta-analysis of SUA. The vertical axis represents $-\log_{10}(P\text{ value})$ for assessment of the association of each SNP with SUA. Panels a–h present plots for chromosome (chr) 1p35.3, 2p25.3, 3q25.1, 5q35.3, 6q21.33, 15q26.1, 20q13.12, or 22q13.31, respectively. Colors indicate LD ($r^2$) between each sentinel SNP and neighboring SNPs based on JPT of 1000 Genomes phase 3. SUA serum uric acid
syndrome”, “sodium”, “transport”, “sodium transport”, and “alternative splicing” were enriched (Supplementary Table 5). For the negatively correlated genes, the term “Williams–Beuren syndrome” was enriched.

Comparison between Japanese and European GWASs for SUA. SNPs located at 28 loci were recently found to show genome-wide significant associations with SUA based on data from individuals of European ancestry in the GUGC24. We examined the results obtained for these SNPs in our meta-analysis (Supplementary Data 6). Twenty-one of these 25 SNPs showed nominal or genome-wide significant associations with SUA in our meta-analysis, with the same direction of effect size in both studies.

We compared the SNP-based heritability ($h^2$) of SUA in our Japanese meta-analysis and the GUGC-based study24. The heritability estimates were calculated from summary statistics of 1,447,573 SNPs, which were assessed in both studies and have MAF $\geq$1% in both studies. The $h^2$ (standard error (SE)) estimates were 14.0 % (4.3%) for our Japanese study and 14.4% (3.9%) for the European study. Furthermore, we calculated the genetic correlation between Japanese and European studies employing the same data sets. The genetic correlation $\rho_{ge}$ (SE) was analyzed (0.591 (0.294), $P$ value = 0.164), and was not significantly less than 1.

Trans-ethnic meta-analysis with the use of GUGC-based study. We performed the trans-ethnic meta-analysis across our meta-analysis and the GUGC-based study to carry out fine-mapping analysis and identify further novel loci associated with SUA. We observed genome-wide significant ($\text{log}_{10}$ (Bayes’ factor) of $\geq$6) association signals at 59 loci (Fig. 3), of which 15 were novel. Shown in Supplementary Data 7 are sentinel SNPs with the highest $\text{log}_{10}$ (Bayes’ factor) for SUA at each of these 15 novel loci (rs302684 of TRABD2B-SKINT1L, rs2765545 of CCDC18, rs715 of CPS1, rs9942075 of TFD2, rs10471103 of INPP4B-LOC105377623, rs461660 of RAI14, rs2760181 of KIAA0319, rs6928482 of HLA-DQB1, rs10971419 of B4GALT1, rs2195525 of USP2, rs626277 of DACH1, rs2957742 of MYO9A, rs12451900 of ZBTB4, rs164009 of QRICH2, and rs1035941 of INSR).

Discussion

In a genome-wide meta-analysis performed with 121,745 Japanese subjects, we here identified eight novel loci significantly associated with SUA. Moreover, five of these loci were replicated in Caucasian populations.

Gout, which develops as a consequence of hyperuricemia, is a form of arthritis known from the time of ancient Egypt37, and modern Japanese are genetically known to be more susceptible to hyperuricemia and gout,10,38. To our knowledge, the present study is the largest genome-wide meta-analysis performed for SUA to date, and it thus provides important insight into the genetic background of hyperuricemia and gout.

Uric acid or urate is an end metabolite of purines such as adenosine derived from ATP and guanine derived from DNA. Urate is produced predominantly in the liver and is excreted by the kidneys and the intestine39,40. Genes for urate transporters and proteins associated with cell metabolism might therefore be expected to be associated with SUA. Indeed, urate transporter genes such as SLC22A12 (also known as URAT1), SLC2A9 (GLUT9), and ABCG2 (BCRP) have been markedly associated with SUA, hyperuricemia, and gout7–12.

Among the eight novel loci identified in our study, TMEM18, TM4SF4, MXD3, and HNF4A are related to cell metabolism or proliferation. TMEM18 is a highly conserved gene related to obesity and plays a role in the central control of appetite and body weight regulation41–43. TM4SF4 is associated with gallstone disease and has been implicated in both liver regeneration and pancreas development44,45. Both MXD3 and HNF4A encode transcription factors. MXD3 forms a heterodimer with the cofactor MAX and is thought to promote uncontrolled cell proliferation and tumorigenesis46,47. HNF4A is associated with nonalcoholic steatohepatitis48 and plays a role in hepatic glucose-neogenesis and lipid metabolism49. In addition, HNF4A controls gene expression in pancreatic islets, with HNF4A mutations having been associated with maturity-onset diabetes of the young type 1 and hyperinsulinemic hypoglycemia50. Furthermore, three of the eight novel loci identified in the present study harbor genes related to oxidative stress and inflammation: SESN2, PSORS1C1, and PNPLA3. SESN2 encodes a highly conserved stress-inducible metabolic protein that protects cells from stressors such as hypoxia, starvation, DNA damage, and oxidative stress51,52. PSORS1C1 and PSORS1C2 encode psoriasis susceptibility 1 candidates 1 and 2, respectively. PSORS1C1 is implicated in synovial inflammation and bone destruction in rheumatoid arthritis53, which, like gout, is a common type of arthritis. Its expression is inhibited in synovial fibroblasts affected by rheumatoid arthritis, which results in a reduction in interleukin-17, osteoclastogenic factor, and interleukin-1 levels as well as attenuation of cell
proliferation. PNPLA3 encodes a membrane protein located at the surface of hepatocyte lipid droplets. A GWAS of nonalcoholic fatty liver disease identified PNPLA3 as a major genetic determinant of fatty liver and hepatic fat content. PNPLA3 is also associated with inflammation, fibrosis, and the development of hepatocellular carcinoma. Thus, novel loci associated with SUA were also related to oxidative stress and inflammation. Given that uric acid has an antioxidative effect, loci related to oxidative stress or inflammation might also be expected to be associated with SUA. However, further molecular functional analyses are required to confirm these associations. The functional relation of the last of the eight novel loci identified in the present study, LINC01578, to SUA is unknown. Indeed, LINC01578 encodes a long intergenic non-protein-coding RNA of unknown function. It is also possible that a gene located near LINC01578 is actually responsible for the observed association with SUA.

Previous candidate analyses and GWASs of clinically defined gout identified nonsynonymous variants of gout susceptibility genes such as ABCG2 (rs72552713, Q126X, rs2231142, Q141K) and GCKR (rs1260326, A446P). SLC22A12 (URAT1) and SLC2A9 (GLUT9) are also genetic loci that influence SUA and encode urate transporters that mediate physiological urate reabsorption in the kidney. We previously showed that dysfunctional nonsynonymous variants of SLC22A12 and SLC2A9 are responsible for renal hypouricemia type 1 and type 2, respectively. The present study also identified missense SNPs at two loci, rs738409 (I148M) of PNPLA3 and rs74896528 (P87S) of SESN2, that are predicted to impair the function of the encoded proteins (Supplementary Table 4). The rs738409 (I148M) polymorphism of PNPLA3 is in LD with rs2281293, which showed the most significant association with SUA at this locus in our genome-wide meta-analysis. The rs2281293 SNP of PNPLA3 is also an eQTL for this gene (Supplementary Data 4). On the other hand, rs74896528 of SESN2 has not been identified as an eQTL (Supplementary Data 4), and its SNP was not reported in the previous study based on GUGC data because of its low frequency in Caucasian populations (Supplementary Data 2). These results suggest that this missense (P87S) variant of SESN2 (rs74896528) is a novel locus that is associated with SUA specifically in Japanese or Asian populations.

The 28 loci identified in the European population of the GUGC study, SNPs at 21 loci showed a nominal or genome-wide significant association with SUA in our meta-analysis (Supplementary Data 6), again with the same direction of effect size. The SNP-based heritability for Japanese was 14.0%, and was similar to the 14.6% seen in Europeans. The genetic correlation between Japanese and Europeans was not significantly <1. These results suggest the possibility that most genetic causal variants of SUA are shared across ancestries.

The present study also identified 132 correlated genes whose expression levels are associated with SUA-increasing alleles (Supplementary Data 5). UniProt term enrichment analysis showed that these correlated genes are enriched in genes related to “transport” (Supplementary Table 5). A novel locus, rs6031598 of HNF4A, is correlated with the expression level of HNF4A. Of note, a noncoding genetic variant, rs1967017 of PDZK1, which encodes a scaffold protein for urate transporters, has been shown to be functionally linked to HNF4-dependent PDZK1 expression.

For SNP rs9394948 of ZNF318, ABC10 (MRP7), an ABC transporter gene, was a positively correlated gene, and SLC22A7 (OAT2), an SLC transporter gene, was a negatively correlated gene (Supplementary Data 5). SLC22A7 encodes organic anion transporter 2 (OAT2), which mediates urate transport and is expressed in kidney and liver. Furthermore, for SNP rs11952102 of MXD3, RAB24, and PRELID1 were positively correlated genes, and MXD3 was a negatively correlated gene. RAB24 is localized to the endoplasmic reticulum and is thought to participate in autophagosome maturation. RAB24 may influence SUA via autophagy, because there is a report on relationship between SUA and autophagy which is promoted by NLRP3 and results in phagocytosis of urate crystals by human osteoblasts. PRELID1 encodes PRELI, which forms a complex with TRAILP1 and mediates intramitochondrial transport of phosphatidic acid. It is possible that PRELI may function as a urate transporter that directly affects SUA or that it indirectly influences SUA via its function as a phosphatidic acid transporter.

In trans-ethnic meta-analysis across our own meta-analysis and the GUGC study, we have here identified 15 more novel loci significantly associated with SUA. Out of these, rs2760181 of KIAA0319 at 6p22.3 showed different direction of regression coefficients between Japanese and European studies, but showed genome-wide significant association (log_{10} Bayes’ factor >6). Future studies will therefore be necessary to validate our findings in independent cohorts.

The present genome-wide meta-analysis of SUA in Japan identified eight novel loci. Furthermore, trans-ethnic meta-analysis of SUA in the present study revealed 15 more novel loci associated with SUA. The present study also demonstrated that SUA is regulated by multiple “transport”-related genes, that is, not only urate transporter genes but also non-transporter genes such as PDZK1 and HNF4A. Our findings thus provide important insight into SUA regulation and the pathogenesis of hyperuricemia and gout, and they provide a potential basis for the development of new treatments for these diseases.

Methods

Study subjects and genotyping. We performed a genome-wide meta-analysis based on three Japanese cohorts including those of the J-MICC Study, KING Study, and BBJ Project. An overview of the characteristics of the study populations is provided in Supplementary Table 1. Information regarding study-specific genotyping, imputation, and analysis tools is provided in Supplementary Table 2. Data and sample collection for the cohorts participating in the present study were approved by the respective research ethics committees. All participants provided written informed consent.

Details of cohorts. The Japan Multi-institutional Collaborative Cohort (J-MICC) Study was launched in 2005. Through March 2014, 92,642 Japanese participants aged 35 to 69 years had provided blood samples and lifestyle data based on a questionnaire after having given their informed consent. The present study included 14,539 J-MICC Study participants randomly selected from the 12 targeted areas. The BioBank Japan (BBJ) Project (http://biobankjp.org/english/index.html) is an ongoing community-based prospective observational study of the genetic basis of cardiovascular disease and its risk factors. It recruited 3975 Japanese subjects aged 50–80 years who underwent community-based annual health checkups between May 2005 and December 2007. A total of 2095 of the KING Study samples was included in the present study. SUA was measured with the uricase-peroxidase method or the uricase–3,5-dimethoxy-4-fluorone (F-DAO5) method in 10,794 of the 14,091 participants. Individuals receiving treatment for hyperuricemia or gout were excluded. Finally, 10,621 participants remained for the association analysis (Supplementary Table 1). This study was approved by the ethics committees of Nagoya University Graduate School of Medicine (approval no. 939-14), Aichi Cancer Center, and all other participating institutions. All research procedures were conducted according to the Ethical Guidelines for Human Genome and Genetic Sequencing Research in Japan and the Declaration of Helsinki.

The Kita-Nagoya Genomic Epidemiology (KING) Study (ClinicalTrials.gov identifier NCT02626291) is an ongoing community-based prospective observational study of the genetic basis of cardiovascular disease and its risk factors. It recruited 3975 Japanese subjects aged 50–80 years who underwent community-based annual health checkups between May 2005 and December 2007. A total of 2095 of the KING Study samples was included in the present study. SUA was measured with the uricase-peroxidase method or the uricase–3,5-dimethoxy-4-fluorone (F-DAO5) method in 10,794 of the 14,091 participants. Individuals receiving treatment for hyperuricemia or gout were excluded. The study was performed according to the guidelines of the Declaration of Helsinki; the study protocol was approved by the ethics committees of Aichi Gakuin University, Ichi Medical University, Nagoya University, and Kyushu University; and all participants provided written informed consent.

The BioBank Japan (BBJ) Project (http://biobankjp.org/english/index.html) was initiated in 2003 at the Institute of Medical Science, The University of Tokyo, and it
has constructed a large-scale, multi-institutional, hospital-based biobank. The BBJ recruited more than 200,000 Japanese patients with any of 47 target diseases between fiscal years 2003 and 2007. Patients were recruited from 66 hospitals of 12 medical institutes throughout Japan (Osaka Medical Center for Cancer and Cardiovascular Diseases, Cancer Institute Hospital of Japanese Foundation for Cancer Research, Juntendo University, Tokyo Metropolitan Geriatric Hospital, Nippon Medical School, Nihon University School of Medical Science, Tokushima University, Shiga University of Medical Science, Fukui University, National Hospital Organization Osaka National Hospital, and Iizuka Hospital). All patients were diagnosed with one or more of the 47 target diseases by physicians at the cooperating hospitals. Clinical information and biochemical measurements was collected through interviews and reviews of medical records with the use of a standard questionnaire. The present study included 109,029 individuals aged between 18 and 85 years with valid SUA measurements as described elsewhere. Subjects receiving urate-lowering therapy (allopurinol, febuxostat, probenecid, or benz bromarone) or with renal insufficiency (estimated glomerular filtration rate of < 15 ml min⁻¹ 1.73 m⁻²) were excluded. We obtained written informed consent from all participants, and this study was approved by the ethics committees of RIKEN Center for Integrative Medical Sciences and the Institute of Medical Science, The University of Tokyo.

Association analysis for SNPs and SUA. Individuals taking urate-lowering drugs were excluded from the present study. SUA was adjusted for age, sex, the top 10 principal components, and study-specific covariates in a linear regression model. We then standardized the resulting residuals. The association of the z-score of the residuals with SNP allele dose was tested by linear regression analysis. The effect sizes and standard errors estimated in linear regression analysis were used in the subsequent meta-analysis.

Quality control after genotype imputation. After genotype imputation, quality control was applied to each study. SNPs with an imputation quality of r² ≥ 0.3 or a minor allele frequency of < 0.005 were excluded. SNPs that passed quality control in both the J-MICC Study and BBJ cohorts were subjected to meta-analysis. To identify studies with inflated GWAS significance, which can result from population stratification, we computed the genomic control lambda (λ) and the intercept of LD score regression. We calculated the genomic control lambda in R. A study showing a score > 1.1 for both measures was regarded as inflated. Inflation was not detected in any study included in the present meta-analysis, and so genomic control adjustment was not applied.

Meta-analysis. The meta-analysis was performed with a total of 121,745 Japanese subjects from the three cohorts (Supplementary Table 1). The association results for each SNP across the studies were combined with METAL software by the fixed-effects inverse-variance-weighted method. Heterogeneity of effect sizes was assessed with the P² index. The meta-analysis included 5,864,938 SNPs and the results from at least both the J-MICC Study and BBJ Project. The genome-wide significance level was set to a P value < 5 × 10⁻⁸.

Replication study for novel loci with the GUGC-based study. To employ a replication study and compare our meta-analysis with publicly available results from Europeans conducted by the GUGC, we downloaded the summary statistics from their website. The EAP of the HapMap project phase 2 CEU samples for each SNP was converted to the summary statistics of the GUGC because these results of the GUGC study did not include EAfs. We excluded variants with MAF < 0.01. P values for the GUGC study were corrected for genomic control (λ = 1.12 for SUA and 1.03 for gout) and genomic inflation did not occur in the GUGC study because the intercepts of the LD score regression, on the raw P-values, were 1.01 for SUA and 1.09 for gout. We therefore calculated the raw P values from the corrected P values, and used the raw P values as a replication study for novel loci in our meta-analysis. For the replication of five novel loci, the significance level was determined by dividing 0.05 by the number of loci for Bonferroni correction (α = 0.05/5 = 0.01).

Functional annotations. For prioritization of associated SNPs at the novel loci, we adopted a series of bioinformatics approaches to collate functional annotation. We first used ANNOVAR71 to obtain an aggregate set of functional annotations, including gene location and impact of amino acid substitution based on the prediction tools SIFT and PolyPhen-2—for the sentinel SNPs and SNPs in high-LD (r² of > 0.8 in JPT of 1000 Genomes phase 3) with the sentinel SNPs and with a P value of < 1 × 10⁻⁸ for SUA. We also examined these sentinel and high-LD SNPs for identification of eQTLs in 14 tissues considered relevant to SUA regulation using the GTEx v7 database. The significant criteria for eQTL were based on the GTEx project21 variants with a nominal P value below the gene-level threshold were regarded as significant. The gene level threshold was determined by the permutation test in the GTEx project. UniProt term enrichment analysis for the sets of positively correlated genes and negatively correlated genes was performed with DAVID and with the threshold of a false discovery rate of < 0.05 as calculated by the Benjamini–Hochberg adjustment method.

SNP-based heritability in Japanese and European samples. We estimated the SNP-based heritability of SUA for our Japanese meta-analysis and GUGC-based study with the use of LD score regression.24 As explained in our replication study section, the EAF of the HapMap project phase 2 CEU samples for each SNP was added to the summary statistics of the GUGC because the results of the GUGC study did not include EAfs. The heritability estimates were calculated from the summary statistics of 1,447,573 SNPs, which were assessed in both studies and have MAF ≥ 1% in both studies and were not pan-Asian SNPs. The P values for the GUGC study were corrected for genomic control (λ = 1.12).24 Genomic inflation did not occur in GUGC because the intercept of LD score regression based on the raw P values was 1.01. Thus, we used raw P values calculated from corrected P values. Furthermore, we calculated the genetic correlation between Japanese and Europeans using the same data sets. The genetic correlation was calculated with the use of Popcorn.22

Trans-ethnic meta-analysis with the use of GUGC-based study. For our trans-ethnic meta-analysis across our meta-analysis and the GUGC-based study, we used MANTRA v1.2 software23, which has been developed for trans-ethnic meta-analysis allowing heterogeneity in allele effects. The trans-ethnic meta-analysis was calculated from the summary statistics of 1,986,983 SNPs, which were assessed in both studies and have MAF ≥ 1% in both. In our meta-analysis, the effect sizes were calculated from a linear regression analysis in which the z-score of residual values of SUA values after adjustment for covariates was used as a dependent variable. In the GUGC project, the effect sizes were calculated from the linear regression analysis in which the SUA value was used as a dependent variable. The scale of effect size for these studies was therefore different. Thus, before the MANTRA analysis, the effect sizes and standard errors of the GUGC study were divided by the standard deviation of SUA in the GUGC study (= 1.4 mg/dl) to approximate the scale of effect sizes. A prioritized model of the relationship between the studies was estimated by employing a dmatcal script in the software using the allele frequency of the analyzed SNPs. We regarded log₁₀ Bayes’ factor > 6 as a significant threshold in line with the previous simulation study.

Data availability
The summary statistics of our genome-wide meta-analysis based on three Japanese cohorts is available at the National Bioscience Database Center (Research ID: hum0167.v1.meta.v1).

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Author contributions

M. Nakatochi, A.H., Y.O. and H. Matsuo conceived and designed the study. A.N., Y. Kamatani, M. Kubo, Y. Kamatani, and N.S. contributed to research design. M. Nakatochi, S.I., K.Y., N.K., T. Takezaki, M. Kanai, Y. Kamatani, and Y.O. performed statistical analysis. A.N., Y. Kawamura, S. Shimizu, K.Y., M. Kawaguchi, M. Nakajima, M.T., and H. Matsuo analyzed data. H. Matsuo organized this collaborative study. H.N., T. Nakamura, N.K., K. Matsuda, Y.M., T.M., N.S., M.Y., and K.W. provided intellectual input and assisted with preparation of the manuscript. M. Nakatochi, M. Kanai, A.H., Y. Kawamura, Y.O. and H. Matsuo wrote the manuscript.

Additional information

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