

Title	Prominence of NUDT15 genetic variation associated with 6-mercaptopurine tolerance in a genome-wide association study of Japanese children with acute lymphoblastic leukaemia
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Citation	British journal of haematology, 199(2), 260-269 https://doi.org/10.1111/bjh.18405
Issue Date	2022-10-01
Doc URL	http://hdl.handle.net/2115/90588
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Туре	article (author version)
File Information	Br J Haematol_bjh.18405.pdf



1 Prominence of NUDT15 genetic variation associated with 6-mercaptopurine tolerance in a 2 genome-wide association study of Japanese children with acute lymphoblastic leukemia 3 4 Yoichi Tanaka¹, PhD; Kevin Y. Urayama^{2,3}, PhD, MPH; Makiko Mori⁴, MD; Yuki Arakawa⁴, MD; 5 Daisuke Hasegawa⁵, MD, PhD; Yasushi Noguchi⁶, MD; Masakatsu Yanagimachi^{7,8}, MD, PhD; Dai Keino^{7,9}, MD, PhD;Setsuo Ota¹⁰, MD, PhD; Koshi Akahane¹¹, MD, PhD; Takeshi Inukai¹¹, MD, PhD; 6 7 Mayumi Hangai³, MD; Takahisa Kawaguchi¹², PhD; Masatoshi Takagi¹³, MD, PhD; Katsuyoshi Koh⁴, 8 MD; Fumihiko Matsuda¹², PhD; Atsushi Manabe¹⁴, MD, PhD 9 10 ¹Division of Medical Safety Sciences, National Institute of Health Sciences, Kanagawa, Japan 11 ²Graduate School of Public Health, St. Luke's International University, Tokyo, Japan 12 ³Department of Social Medicine, National Center for Child Health and Development, Tokyo, Japan 13 ⁴Department of Hematology/Oncology, Saitama Children's Medical Center, Saitama, Japan 14 ⁵Department of Pediatrics, St. Luke's International Hospital, Tokyo, Japan 15 ⁶Department of Pediatrics, Japanese Red Cross Narita Hospital, Chiba, Japan 16 ⁷Department of Hematology/Oncology, Children's Cancer Center, Kanagawa Children's Medical 17 Center, Kanagawa, Japan 18 ⁸Department of Pediatrics, Yokohama City University Hospital, Kanagawa, Japan 19 ⁹Department of Pediatrics, St. Marianna University, Kanagawa, Japan 20 ¹⁰Department of Pediatrics, Teikyo University Chiba Medical Center, Chiba, Japan 21 ¹¹Department of Pediatrics, University of Yamanashi, Yamanashi, Japan ¹²Center for Genomic Medicine, Kyoto University, Kyoto, Japan 22 23 ¹³ Department of Pediatrics, Tokyo Medical and Dental University, Tokyo, Japan 24 ¹⁴Department of Pediatrics, Hokkaido University, Hokkaido, Japan 25 26 Correspondence: Yoichi Tanaka 27 Division of Medical Safety Sciences, National Institute of Health Sciences 28 3-25-26, Tonomachi, Kawasaki-ku, Kawasaki-shi, Kanagawa 210-9501 JAPAN 29 Tel: +81-44-270-6626; FAX: +81-44-270-6627 30 E-mail: tanakayoichi@nihs.go.jp 31 32 Running Title: Genetic risk factors for 6-MP tolerance in Japanese ALL 33 Keywords: 6-mercaptopurine, tolerance, genetic variant, genome-wide association study, Japanese 34 Counts: Abstract- 193 words; Text- 3,106 words; Tables- 3; Figure- 2; References- 37 35

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37 Summary

38 Inherited genetic variation is associated with 6-mercaptopurine (6-MP) dose reduction and 39 frequent 6-MP induced toxicities. However, tolerable dose for 6-MP is not completely predicted by 40 the known variation in NUDT15 and TPMT among Asian children with acute lymphoblastic leukemia 41 (ALL). We performed a genome-wide association study (GWAS) related to 6-MP dose among 42 Japanese children with ALL. This GWAS comprised 224 patients previously enrolled in Tokyo 43 Children's Cancer Study Group clinical studies with replication attempted in 55 patients. Genome-44 wide single nucleotide polymorphism (SNP) genotypes were evaluated for association with 6-MP 45 average dose during initial 168 days of maintenance therapy. Possible associations were observed 46 across 5 gene coding regions, among which only variants at 13q14.2 were genome-wide significant 47 and replicated (rs116855232, NUDT15, β =-10.99, P=3.7×10⁻¹³). Notable findings were observed for variants in AFF3 (rs75364948, P=2.05×10⁻⁶) and CHST11 (rs1148407, P=2.09×10⁻⁶), but were not 48 49 replicated possibly due to small numbers. A previously reported candidate SNP in MTHFR was associated with higher 6-MP average dose (rs1801133, P=0.045), and FOLH1 (rs12574928) was 50 51 associated in a candidate region evaluation ($P_{adjust}=0.013$). This study provides strong evidence that rs116855232 in NUDT15 is the prominent genetic factor associated with 6-MP tolerable dose in 52 53 Japanese. 54 55 56 57

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61 Introduction

A survival probability of 80-90% in children with acute lymphoblastic leukemia (ALL) has 62 63 been achieved due largely in part to advances in combination chemotherapy¹. 6-mercaptopurine (6-64 MP) is a main component for improving therapeutic outcomes, but tolerability is different in each 65 patient. Failure to minimize incidents of therapy interruption may affect prognostic outcomes for 66 childhood ALL patients². Response to 6-MP has been associated with variants in genes participating in the 6-MP metabolism pathway³. It is well known that genetic variation of the TPMT gene is 67 associated with 6-MP intolerance³. However, the TPMT variant was not shown to be associated with 68 69 thiopurine induced toxicities in Japanese⁴, and the frequencies of TPMT poor metabolizer was lower 70 in East Asians compared with other races. In 2014, the NUDT15 rs116855232 variant was reported to 71 influence 6-MP intolerance identified through a genome-wide association study (GWAS), and the 72 frequency of that variant is higher in Asians compared to populations of European and African 73 ancestries⁵. Subsequently, NUDT15 genotype was shown to affect 6-MP tolerability among Asians 74 undergoing childhood ALL therapy⁴.

75 To date, several candidate gene studies have shown that genetic variation in NUDT15 is associated with 6-MP dose reduction and frequent 6-MP-induced toxicities in childhood ALL^{4,6-13}. 76 77 NUDT15 enzyme dephosphorylates thio-guanosine triphosphate (GTP) and deoxy thio-GTP to thio-78 guanosine monophosphate (GMP) and deoxy thio-GMP, respectively. Lower NUDT15 enzyme leads 79 to increased thioguanosine incorporation ratio to 6-MP dose and 6-MP related toxicities. In some 80 instances, severe toxicities requiring significant 6-MP dose reduction have been observed in patients 81 who do not carry the known genetic risk factors for 6-MP tolerability. Thus, tolerable dose for 6-MP 82 is not completely predicted by the variation in NUDT15 and TPMT.

83 The aim of this study was to perform a GWAS to examine the role of inherited genetic variation
84 related to 6-MP dose, with objectives to both identify newly associated variation, as well as to

85 characterize the variants previously reported in diverse populations, among Japanese childhood ALL

86 patients. Moreover, we pursued a targeted examination of genetic variation across the candidate gene

87 regions to identify additional associations with 6-MP dose.

88

89 Methods

90 Patients and sample collection

91 This GWAS comprised patients previously enrolled in a Tokyo Children's Cancer Study Group (TCCSG) clinical study including L89-12^{14,15}, L95-14^{15,16}, L99-15¹⁷ and L04-16¹⁸ from 23 clinical 92 93 centers collaborating on this genomic study. Patients were recruited at the time of routine outpatient 94 follow-up visit between 2013 and 2015, as previously described¹⁹. Briefly, patients were considered 95 eligible if they were aged 19 years or younger at the time of ALL diagnosis and self-identified as 96 Japanese. Saliva samples were collected at the time of follow-up visit after remission. DNA was 97 extracted using the Oragene prepIT DNA extraction kit (DNA Genotek, Ottawa, Canada), and stored at -80 °C. The study protocol was approved by the institutional review boards of all collaborating 98 99 institute and hospital involved in patient recruitment. Written informed consent was obtained from the 100 parents of each participant together with a written informed consent or assent by the child depend on 101 age.

102

103 Genotyping and quality control

Whole genome single nucleotide polymorphism (SNP) microarray genotyping was performed using the Illumina HumanCoreExome-12 v1.1 BeadChip (San Diego, CA) and has been described previously ¹⁹. Briefly, quality control (QC) procedures comprised excluding samples if the genotype call-rate was below 95% and samples that exhibited relatedness based on an identity-by-descent analysis. SNPs were excluded if genotype call-rate was less than 99%, genotype distribution deviated from that expected based on Hardy-Weinberg equilibrium ($P > 1 \times 10^{-6}$), or the minor allele frequency (MAF) was less than 0.01. Additionally, principal components (PC) analysis based on a subset of the post-QC genome-wide SNPs in low linkage disequilibrium (LD) was performed using EIGENSTRAT 2.0 software together with HapMap data from Japanese, and sample outliers were excluded based on a plot of the leading PCs.

114 Genome-wide SNP imputation was performed using ShapeIT2 and Minimac4 with reference 115 population from the 1000 Genomes Project Phase III Version 5. SNP imputation QC comprised 116 excluding poorly imputed SNPs defined by an R^2 of less than 0.5, resulting in a total of 6,236,137 117 SNPs available for analysis.

118

119 Clinical data

120 The clinical data and details of the administration of 6-MP doses during maintenance therapy were 121 available for 289 Japanese patients. In TCCSG protocols, maintenance therapy is initiated with 40 122 mg/m²/day of 6-MP and 25 mg/m²/week of oral methotrexate. These dosages were adjusted to 123 maintain the target leucocyte count at 1,500 - 3,000/mm³. We excluded patients who started at more 124 than 125% or less than 75% of protocol 6-MP dose, as therapeutic dose for 6-MP and MTX were 125 adjusted for drug induced toxicities before initiation of maintenance therapy. In total, we included 126 224 ALL patients (discovery cohort) who started maintenance therapy using the normal protocol 127 dose (30–50 mg/m²/day). The outcome variable for this GWAS was defined as 6-MP average dose 128 for initial 168 days of maintenance therapy.

129

130 Replication series

The replication series included 55 patients who started maintenance therapy by normal protocol
 recruited previously through TCCSG as part of a separate study⁴. DNA were extracted from peripheral

blood obtained at remission, and SNP genotyping was performed using Taqman real-time PCR assays(Applied Biosystems, Waltham, MA).

135

136 Statistical analysis

137 We performed genome-wide association analyses of SNPs in relation to 6-MP average dose for 138 initial 168 days of maintenance therapy among the discovery cohort using linear regression assuming 139 a log-additive genetic model of inheritance and adjusting for age at diagnosis. Association analysis 140 assuming dominant and recessive genetic inheritance models were also performed. Results showing a $P < 5 \times 10^{-8}$ were considered statistically significance at the genome-wide level, and a $P < 1 \times 10^{-5}$ 141 142 was considered as showing a suggestive association. In the replication series, association analysis was 143 conducted similarly using linear regression assuming a log-additive genetic model and adjusting for 144 age at diagnosis. We defined a Bonferroni corrected P < 0.05 as statistically significant in the 145 replication. Additionally, among the top SNPs, the difference of 6-MP average dose across the three 146 possible genotypes was evaluated using the Kruskal-Wallis test. Normality test of 6-MP distribution 147 was evaluated using the D'Agostino-Pearson test.

148 The association of specific candidate SNPs reported previously, as well as the variants in coding 149 regions of those candidate genes were examined. Association results with a nominal p-value of less 150 than 0.05 was considered statistically significant for specific candidate SNPs. For regional 151 examination of association results across candidate genes, SNPs with a nominal p-value of less than 152 0.05 were considered noteworthy, and a condensed list of SNPs pruned on LD ($r^2 > 0.50$) within each 153 gene were adjusted for multiple testing based on 10,000 permutation of the data on the 6-MP dose 154 outcome in which a p-value below a family-wise type I error rate threshold of 0.05 was considered 155 statistically significant. Statistical analyses were performed using PLINK version 1.9, R software 156 (version 3.6.1) and Prism 9 (GraphPad Software, San Diego, CA).

157

158 Results

Genome-wide association analysis of 6-MP average dose during 168 days of maintenance therapy A total of 224 patients were included in the genome-wide association analysis. No patients needed to be excluded due to outlying genetic ancestry based on PC analysis evaluations. The characteristics of patients were shown in Table 1. Age at diagnosis was significantly correlated with 6-MP average dose ($\beta = -0.88$, $P = 6.96 \times 10^{-5}$). The median 6-MP average doses for the initial 168 days were 41.1 (interquartile range: IQR 25% – 75%, 32.8 – 48.6), and did not appear to deviate from a normal distribution (P > 0.05).

166 In the genome-wide analysis of the discovery cohort, linear regression of the 6-MP average 167 dose adjusted for age at diagnosis showed minimal evidence of genomic inflation ($\lambda = 1.01$) 168 (Supplementary Figure S1). The Manhattan plot of the results showed potential association of variants 169 representing 5 genetic coding regions ($P < 1 \times 10^{-5}$) with 6-MP average dose (Figure 1, Table 2 and 170 Supplementary Table S1). Genome-wide significant associations with 6-MP average dose were 171 observed for variants at the chromosome 13q14.2 region in which the leading SNP was rs116855232 located in NUDT15 SNP ($\beta = -11.45$, $P_{additive} = 8.5 \times 10^{-10}$, $P_{dominant} = 2.27 \times 10^{-9}$). Other variants 172 173 within this cluster of associated SNPs were in LD with this SNP (Supplementary Figure S2, $r^2 > 0.8$). 174 Suggestive associations were observed at chromosome 2q11.2 (rs75364948, AFF3), 2p21 175 (rs14452634, THADA), 12q23.3 (rs1148407, CHST11) and 16q23.2-3 (rs12934986 and rs10153053, 176 *CMIP*) (Table 2). In a sensitivity analysis, the results excluding outlier values of 6-MP (defined as the 177 values of more than the third IQR + $1.5 \times$ IQR or less than the first IQR + $1.5 \times$ IQR) showed a 178 persistent association with the chromosome 13q14.2 region, and slight attenuation of association for 179 the other loci (Supplementary Figure S3).

180

To identify associations in the absence of the effect of the well-known loci for 6-MP intolerance,

181 we excluded patients who were carriers of the *TPMT* rs1142345 and *NUDT15* rs116855232 variants. 182 In this cohort, 7 and 41 patients were carriers of the *TPMT* and *NUDT15* variants, respectively. Only 183 one patient carried both the *TPMT* and *NUDT15* variants. Among the remaining 178 patients, the 184 genome-wide analysis of 6-MP average dose showed potential associations with variants representing 185 13 genetic coding regions ($P < 1 \times 10^{-5}$) (Supplementary Table S2). The results of the suggestive 186 variants for *CHST11*, *CMIP* and *THADA* were similar to that of the full analysis.

187

188 Replication analysis

189 For the replication series, we selected three potentially associated SNPs that reside within genes 190 that have been previously implicated in 6-MP tolerability or progression of hematological malignancy 191 (Table 2). The association with NUDT15 rs116855232 strongly replicated in this series ($P = 5.43 \times$ 192 10⁻⁵); however, while the direction of association was consistent, rs75364948 (AFF3) and rs1148407 193 (CHST11) were not significantly associated with 6-MP average dose (Table 2). Combining the 194 estimates from the discovery and replication cohorts, a strong genome-wide significant association between rs116855232 (NUDT15) and 6-MP average dose was observed ($\beta = -10.99$, $P = 3.66 \times 10^{-13}$). 195 196 Variant rs116855232 in the NUDT15 gene represents the prominent association with 6-MP 197 average dose within our Japanese study. Among the combined discovery and replication cohorts, 51 198 patients were heterozygous for rs116855232 (C/T), 5 patients were homozygous for the variant (T/T). 199 The median of the 6-MP average dose for patients with CT and TT were 31.6 and 13.1 mg/m²/day, 200 respectively; these 6-MP doses were significantly lower compared to that of patients with the CC 201 genotype (41.9 mg/m²/day, $P = 3.74 \times 10^{-11}$, Figure 2A).

202

203 Previously reported candidate genes and 6-MP tolerable dose

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As a secondary aim, we selected candidate loci based on review of the literature and the

205 PharmGKB database (https://www.pharmgkb.org) and performed targeted examination of these candidate regions. SNPs rs1142345 (TPMT)⁵, rs3765534 (ABCC4)²⁰, rs1142345 (ITPA)^{21,22}, 206 rs4149056 (SLCO1B1)²³, rs1801133 (MTHFR)²⁴, rs61886492 (FOLH1)²⁵, rs12199316 (NHLRC1)²⁶ 207 and rs72846714 $(NT5C2)^{27}$ have been studied previously as candidate loci for 6-MP tolerable dose 208 209 and/or toxicities. Among these, MTHFR rs1801133 showed a significant association with 6-MP 210 average dose in our study ($\beta = 2.26$, P = 0.045). However, the other reported genetic variants were not 211 related with 6-MP dose in this analysis (Table 3). In addition, other variants in LD with specific 212 candidate SNPs were not associated. A range of genetic variation across these candidate genes were 213 targeted to examine whether other SNPs may be associated with 6-MP average dose in Japanese. SNP 214 associations showing nominal p-values of less than 0.05 were observed for MTHFR, FOLH1, 215 ABCC4, and ITPA (Table 3). Among these, rs12574928 in FOLH1 showed a statistically significant association with 6-MP average dose after adjusting for multiple testing (β = -8.02, P = 216 217 0.013).

218

219 Discussion

220 We observed strong evidence showing that genetic variation at the chromosome 13q14.2 region 221 is significantly associated with 6-MP average dose for initial 168 days of maintenance therapy. Among 222 the cluster of association signals, NUDT15 rs116855232 was the leading SNP, and the other variants 223 spanning the SUCLA2 and MED4 genes were in strong LD with this SNP. NUDT15 genetic variation 224 has been shown to effect thiopurine tolerance in our previous studies, as well as others from Asian 225 countries^{4,12,28} In this study, 19 of 279 patients experienced a reduction of average 6-MP dose during 226 the 168 days to a level less than 20 mg/m²/day. Five of 19 patients (26%) who needed a dose reduction 227 were carriers of the rs116855232 variant. Lower NUDT15 enzyme leads to increased thioguanosine 228 incorporation ratio to 6-MP dose and 6-MP related toxicities. Based on the characteristics of the 229 current study comprising an ethnically homogeneous group of Japanese children with ALL, NUDT15

230 was observed to have the strongest effect on 6-MP tolerance, and the associated variant rs116855232

231 is more frequent in Asians than other races/ethnicities.

The *TPMT* variant is a well-known genetic risk factor for 6-MP tolerance³. In this population, only 10 patients (1.8%) carried the *TPMT* rs1142345 variant, and their 6-MP average dose was 40.0 (19.1 – 57.4) mg/m²/day. The MAF of rs1142345 in this Japanese cohort was lower than populations of European and African ancestry (MAF; 4 – 6%), and there were no patients who were homozygous carriers. While not significant possibly due to limitations in statistical power, the effect size of this variant was also weaker compared to reports from other races/ethnicities.

238 Interestingly, 2 patients with TPMT rs1142345 requiring 6-MP average dose of less than 20 239 mg/m^2 , were carriers of the NUDT15 rs116855232 variant. Yang et al. reported that patients with 240 heterozygous genotypes for both TPMT rs1142345 and NUDT15 rs116855232 had 30-60% of 6-MP 241 standard dose during 6-months of maintenance therapy among the multi-ethnic patient cohort⁵. Choi 242 et al. reported NUDT15 rs116855232 was significantly associated with high 6-thioguanine nucleotide 243 levels in erythrocytes, 6-MP dose ratio, and frequency of leukopenia, but TPMT and other variants 244 were not associated among Korean children with ALL⁹. In our population, the average 6-MP dose for 245 heterozygous carriers of TPMT rs1142345 was similar to wild-type patients, but the average dose for 246 patients with both TPMT rs1142345 and NUDT15 rs116855232 was 50% of tolerable dose of the wild-247 type patients. Moreover, the standard dose of 6-MP on protocol for childhood ALL in Asian countries²⁹ 248 is set at 70% of the dose (40-60 mg/m²) applied to protocols in Europe and US (75 mg/m²). Although 249 there does appear to be an effect of TPMT rs1142345 on 6-MP tolerability in Asians in the presence of NUDT15 variation, these effects may be small due to low prevalence of the variant and a low 250 251 standard dose. Results suggest that heterozygous variants of both NUDT15 and TPMT may induce less 252 tolerability for 6-MP than NUDT15 heterozygous genotype alone.

253 Other genetic variants identified through candidate gene approaches have been reported to be 254 associated with thiopurine tolerable dose. In our study, *MTHFR* rs1801133 was associated with 6-MP 255 dose, but the direction of association is in contrast to other reports²⁴. *MTHFR* is involved in folate 256 metabolism and rs1801133 has been associated with hepatotoxicity. However, our study showed 257 rs1801133 to be associated with high 6-MP average dose. Other previously reported candidate genetic 258 variants were not associated with 6-MP average dose in our study.

259 In maintenance therapy, 6-MP dose and therapeutic duration is different by country, and allele 260 frequencies of candidate variants may be different. This circumstance makes validation of SNP 261 associations and consistency a challenge. The candidate missense SNP rs61886492 in FOLH1 could 262 not be evaluated in the current study due to its absence in the Japanese population. This variant was 263 previously reported to be associated with 6-MP mediated toxicity ²⁵. Interestingly, regional evaluation 264 of genetic variation of this gene in our study identified a significant association signal led by SNP rs12574928 located in an intronic region. This SNP has been described as an expression quantitative 265 266 trail locus (eQTL) for FOLH1 in multiple tissues as documented in the Genotype-Tissue Expression 267 (GTEx) Portal. Due to LD broadly spread across this relatively short gene, localizing a defined causal 268 region will require additional studies.

269 We showed that NUDT15 rs116855232 is the most prominently associated genetic loci for 6-270 MP tolerance in Japanese children with ALL. However, 6 of 20 patients who required significant 6-271 MP dose reduction to a level below 20 mg/m²/day did not carry the known risk variants for 6-MP 272 tolerance. Moreover, 17 patients needed to increase 6-MP dose to a level greater than $60 \text{ mg/m}^2/\text{day}$. 273 Five previously unreported variants located in four genes showed suggestive associations with 6-MP 274 tolerance, including loci residing in AFF3, CHST11, THADA and CMIP. However, these suggestive 275 variants are located in intronic regions, and their functional significance has not been reported. The 276 associated AFF3 rs75364948 and surrounding SNPs in strong LD showed to be potential eQTL for 277 AFF3 and KIAA1211L. AFF3 encodes a tissue-restricted nuclear transcriptional activator, and AFF3 upregulation was reported to be high in tamoxifen resistant breast cancer³⁰. While the function of 278 279 KIAA1211L is not clear, a previous study of molecular markers among childhood leukemia patients 280 showed this gene to be differentially expressed between patients with and without central nervous 281 system involvement³¹. CHST11 was reported previously in relation to the efficacy of methotrexate 282 treatment in rheumatoid arthritis ³². THADA and CMIP genetic variants were identified in GWAS as a susceptibility locus for type 2 diabetes mellitus^{33,34}, and other diseases^{35,36}. These suggestive findings 283 284 could not be replicated in our small patient series in the present study. Future activities will comprise 285 securing additional patient numbers for a statistically robust opportunity to confirm these findings in 286 further replication attempts.

In this study, 6-MP average dose for initial 168 days of maintenance therapy was negatively correlated with age at diagnosis. Therefore, the association of 6-MP and genetic variant was adjusted by age. Although there have been no reports on the relationship between 6-MP tolerable dose and age, some reports have shown that age was negatively correlated with TPMT activity³⁷.

291 There are some limitations of this study to acknowledge. Due the retrospective nature of this 292 study, we decided to exclude patients who started at more than 125% or less than 75% of protocol 6-293 MP dose. These patients were not suitable for evaluation for 6-MP response to standard therapy 294 in maintenance therapy because therapeutic dose for 6-MP and MTX were adjusted for drug 295 induced toxicities before maintenance therapy. Therefore, it is possible that our study cohort 296 underrepresented patients who deviated from the normal 6-MP tolerance. Despite this limitation, our 297 study is meaningful in describing the responsible genetic factors of 6-MP tolerance among patients 298 starting with standard protocol dose.

299

300 Conclusion

301	We provide strong evidence that NUDT15 rs116855232 is the prominent genetic factor
302	associated with 6-MP tolerable dose, and it currently represents the most meaningful clinically
303	predictive marker in Japanese. These findings indicated that NUDT15 genotyping should be
304	considered prior to initiation of 6-MP treatment. Prospects for the identification of additional loci of
305	6-MP tolerance are high after further validation of suggestive loci observed in this Japanese study.
306	
307	Acknowledgements
308	This work was support by funding from St. Luke's Life Science Institute (Tokyo, Japan), the Japan
309	Society for the Promotion of Science KAKENHI grant numbers (15K18932, 26253041), the
310	Children's Cancer Association of Japan, and the Japan Leukemia Research Fund.
311	
312	Conflict of interest
313	There are no conflicts of interest to declare.
314	
315	Authorship contributions
316	Y.T., K.Y.U., M.T. and A.T designed the experiments. Y.T., K.Y.U. and M.H analyzed data. M.M.,
317	Y.A., D.H., Y.N., M.Y., D.K., S.O., K.A., T.I., M.T. K.K., and A.M. collected the clinical data. T.K.
318	and F.M. contributed to perform genotyping. Y.T. and K.Y.U. wrote the paper. All authors approved
319	the manuscript and the interpretation of the data.
320	
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436 Figure legends

437 Figure 1. Manhattan plot of results of the GWAS using linear regression adjusting for age at diagnosis.

- 439 Figure 2. Average 6-MP dose for 168 days from initial maintenance therapy in the combined Discovery
- 440 and replication cohort (N = 279). Difference between genotypes were analyzed by the Kruskal-Wallis
- 441 test. (A) *NUDT15* rs116855232; (B) *AFF3* rs75364948; (C) *CHST11* rs1148407.

Table 1. Demographic and clinical characteristics of patients

	Discovery	Replication
Patient number	224	55
Gender		
Female	99 (44.2%)	24 (43.6%)
Male	125 (55.8%)	31 (56.4%)
Age at diagnosis (years)		
0-1	14 (6.3%)	3 (5.5%)
2-4	104 (46.4%)	19 (34.5%)
5-10	79 (35.3%)	26 (47.3%)
11-19	27 (12.0%)	7 (12.7%)
Median (range)	4.7 (0.9 – 15.7)	5.0 (1 – 17)
6-mercaptopurine initial dose (mg/m ² , median)	39.9 (30.2 - 50.0)	40.1 (30.8 - 49.5)
6-mercaptopurine average dose for 168 days (mg/m ² , median)	41.1 (13.3 – 78.5)	37.2 (5.5 - 60.0)
methotrexate dose (mg/m ² /week, median)	24.5 (0-31.5)	24.7 (0 - 30)

					Discovery	(N = 224)	Replication $(N = 55)$		Combined ($N = 279$)	
Chr	Position	Gene	SNP	MAF in	β	Р	β	Р	β	Р
	(GRCh37)			discovery	(95% CI)		(95% CI)		(95% CI)	
13	48619855	NUDT15	rs116855232	0.094	-11.45	8.46×10^{-10}	-9.13	5.43×10^{-5}	-10.99	3.66 × 10 ⁻¹³
					(-14.95, -7.95)		(-13.29, -4.96)		(-13.82, -8.16)	
2	100399866	AFF3	rs75364948	0.203	6.49	2.05×10^{-6}	0.84	0.766	5.83	3.12×10^{-6}
					(3.88, 9.10)		(-4.76, 6.44)		(3.42, 8.24)	
12	105058593	CHST11	rs1148407	0.138	8.08	2.09×10^{-6}	2.22	0.421	6.52	2.03×10^{-5}
					(4.83, 11.3)		(-3.28, 7.74)		(3.56, 9.49)	
2	43501170	THADA	rs144526347	0.015	21.37	2.28×10^{-6}				
					(12.74, 30.00)					
16	81553070	CMIP	rs12934986	0.277	5.63	6.43×10^{-6}				
					(3.24, 8.01)					
16	81659083	CMIP	rs10153053	0.234	-5.91	5.79×10^{-6}				
					(-8.40, -3.14)					

Table 2. Results of the genome-wide association analysis of 6-mercaptopurine dose and replication (age adjusted)

Abbreviations; Chr, chromosome; SNP, single nucleotide polymorphism; MAF, minor allele frequency.

							Discovery (N=			
Chr	Position	Gene	SNP	Function	Alleles	MAF	β	Р	Р	Reference
	(GRCh37)						(95% CI)	(nominal)	(adjusted)	
Previ	ous reported S	NPs								
1	11856378	MTHFR	rs1801133	Missense	G/A	0.375	2.26 (0.06, 4.46)	0.0454		24
6	18123502	NHLRC1	rs12199316	Upstream	C/G	0.394	-0.69 (-3.06, 1.69)	0.553		26
6	18130918	TPMT	rs1142345	Missense	T/C	0.0156	-2.38 (-11.45, 6.70)	0.608		2
10	104878454	NT5C2	rs72846714	Intron	G/A	0.024	3.75 (-6.00, 13.5)	0.452		27
11	49186274	FOLH1	rs61886492	Missense	G/A					25
12	21331549	SLCO1B1	rs4149056	Missense	T/C	0.147	-2.45 (-5.47, 0.57)	0.113		23
13	95815415	ABCC4	rs3765534	Missense	C/T	0.143	-1.66 (-4.88, 1.55)	0.310		20
20	3193842	ITPA	rs1127354	Missense	C/A	0.152	1.01 (-2.05, 4.07)	0.520		21, 22
Gene	tic variants aci	ross previous	ly reported candi	date gene regi	ons (nominal	P-value<0.05	5)			
11	49213504	FOLH1	rs12574928	Intron	C/T	0.056	-8.02 (-12.9, -3.21)	1.59×10^{-3}	0.013	25
13	95707834	ABCC4	rs9561773	Intron	C/T	0.228	3.69 (1.02, 6.37)	7.35×10^{-3}	0.577	20
13	95953517	ABCC4	rs11568681	Missense	G/T	0.018	10.91 (2.54, 19.29)	0.011	0.707	20
20	3200778	ITPA	rs959815466	Intron	C/T	0.013	-11.55 (-21.2, -1.90)	0.019	0.126	21,22

Table 3. Results of the relationship between previously reported candidate genes and average 6-mercaptopurine dose for 168 days (age adjusted)

Abbreviations; Chr, chromosome; SNP, single nucleotide polymorphism; MAF, minor allele frequency.







