

**Clinical Pharmacogenetics Implementation Consortium (CPIC) guideline for thiopurine dosing based on *TPMT* and *NUDT15* genotypes: 2018 update**

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**CONFLICT OF INTEREST**

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## ABSTRACT

TPMT activity exhibits a monogenic co-dominant inheritance and catabolizes thiopurines. *TPMT* variant alleles are associated with low enzyme activity and pronounced pharmacologic effects of thiopurines. Loss-of-function alleles in the *NUDT15* gene are common in Asians and Hispanics and reduces the degradation of active thiopurine nucleotide metabolites, also predisposing to myelosuppression. We provide recommendations for adjusting starting doses of azathioprine, mercaptopurine, and thioguanine based on *TPMT* and *NUDT15* genotypes (updates on [www.cpicpgx.org](http://www.cpicpgx.org)) .

## INTRODUCTION

This document is an update to the Clinical Implementation Consortium (CPIC) Guidelines for Thiopurine Methyltransferase Genotype and Thiopurine guideline updated last in April 2013. The guideline text, evidence table and recommendations have been updated to reflect any new evidence. Specifically, this guideline adds a recommendation for *NUDT15* genotype with minor changes to the *TPMT* recommendation. Although most of the dosing recommendations have been generated from clinical studies in just a few diseases, we have extrapolated recommended doses to all conditions, given the pharmacokinetic nature of the genotype/phenotype associations. CPIC guidelines are published and periodically updated on [www.cpicpgx.org](http://www.cpicpgx.org). Detailed guidelines for use of phenotypic tests (e.g. *TPMT* activity and thiopurine metabolite levels), as well as analyses of cost effectiveness, are beyond the scope of this document.

## FOCUSED LITERATURE REVIEW

A systematic literature review focused on *TPMT* and *NUDT15* genotypes and thiopurine use was conducted (details in **Supplement**). Definitive reviews (1-4) were relied upon to summarize much of the earlier literature.

## DRUGS: THIOPURINES

### Background

Three thiopurines are used clinically: azathioprine (a prodrug for mercaptopurine), mercaptopurine, and thioguanine. Although all three medications share many of the same pharmacologic effects, mercaptopurine and azathioprine are generally used for non-malignant immunologic disorders, mercaptopurine for lymphoid malignancies, and thioguanine for myeloid leukemias. Because azathioprine is a prodrug for mercaptopurine, the two drugs can be considered to have identical interactions with TPMT and NUDT15. Recommendations for individuals with variants in one or both of these genes will be addressed in detail in the following sections.

## GENES: *TPMT* AND *NUDT15*

### Background

***TPMT***. TPMT activity is inherited as a monogenic, autosomal co-dominant trait (**Supplement, Figure S1**). Three *TPMT* single nucleotide polymorphisms (SNPs), which result in unstable proteins and enhanced TPMT protein degradation (2, 3), account for over 90% of low activity phenotypes and are the most common inactivating alleles, and so genotyping tests including these three variants have a high likelihood of being informative for TPMT phenotype (5, 6). Complementary phenotype laboratory tests can be helpful adjuncts to genotyping tests (**Supplement, Other Considerations**) (7).

TPMT catabolizes mercaptopurine to an inactive methylmercaptopurine base, leaving less parent drug available for eventual anabolism to active thioguanine nucleotides (TGNs, **Figure 1**). The secondary metabolite of mercaptopurine, thioinosine monophosphate (TIMP), is also a substrate for TPMT, and methylTIMP (and its further phosphorylated metabolites, methylmercaptopurine nucleotides or MeMPN) have pharmacologic activity (mostly immunosuppressive and hepatotoxic), inhibit *de novo* purine synthesis, and may contribute to some of the adverse effects of mercaptopurine, generally hepatotoxicity (2, 8, 9). Individuals who inherit two loss-of-function *TPMT* alleles (homozygous or

compound heterozygous *TPMT* deficient individuals) are at very high risk for life-threatening myelosuppression, due to very high TGNs, if given conventional doses of mercaptopurine (or azathioprine). Despite having higher TGNs than wild-type patients, only about 30-60% of *TPMT* heterozygotes cannot tolerate full doses of mercaptopurine or azathioprine (8, 10, 11). Good thiopurine tolerance in some heterozygotes may be because although they have higher TGNs than homozygous wild-type patients, they have lower concentrations (and thus fewer toxic effects) of the MeMPNs than do normal metabolizers, which may offset the toxic effects of having higher TGNs. Thus, there is less of a consensus over how to dose azathioprine and mercaptopurine in patients who are heterozygous for *TPMT* compared to those that are homozygous, although they are at a higher risk for toxicity compared to patients carrying two normal function alleles (12).

Although there is lower affinity between thioguanine and *TPMT* than between mercaptopurine and *TPMT*, *TPMT* significantly affects thioguanine pharmacokinetics and its cytotoxic effects (12-16). Thioguanine is directly metabolized by *TPMT* to inactive methylthioguanine base, leaving less drug available for anabolism by *HPRT* and other enzymes to active TGN metabolites. There is not an analogous secondary metabolite of thioguanine to undergo activation via *TPMT* (i.e. there are no methylTIMP or methylmercaptopurine nucleotides). As a result, patients receiving thioguanine are able to tolerate substantially higher TGN concentrations than do those receiving mercaptopurine or azathioprine (15). Within each *TPMT* phenotypic group, the initial recommended relative dosage decreases are similar for thioguanine, mercaptopurine, and azathioprine (**Table 2**).

***NUDT15***. Through agnostic genome-wide association studies, variants in *NUDT15* have been identified that strongly influence thiopurine tolerance in patients with acute lymphoblastic leukemia (ALL) (17) and those with inflammatory bowel diseases (18). As a nucleoside diphosphatase, *NUDT15*, catalyzes the conversion of cytotoxic thioguanine triphosphate (TGTP) metabolites to the non-toxic thioguanine monophosphate (TGMP). Defects in *NUDT15*-mediated degradation of TGTP results in more TGTP available for incorporation into DNA (DNA-TG, the primary antileukemic

metabolite (19)), thus allowing for DNA damage and apoptosis. The SNP (rs 116855232; c.415C>T) causing p.R139C was the first *NUDT15* variant linked to thiopurine toxicity. It was shown that this amino acid change results in a nearly complete loss of enzymatic activity and protein stability *in vitro*. Patients carrying this allele showed excessive DNA-TG and severe myelosuppression (20). In children with ALL, patients homozygous for the p.R139C variant allele tolerated only 8% of the standard dose of mercaptopurine, whereas tolerated dose intensity was 63% and 83.5% for those heterozygous and wildtype for this SNP, respectively (17). While most clinical studies focused on mercaptopurine, *in vitro* experiments using laboratory models indicated similar influence of *NUDT15* on the cytotoxicity of azathioprine and thioguanine (20). Additional variant alleles have been identified with varying prevalence among differing ancestral groups and varying degrees of functional effects (***NUDT15* Allele Functionality Table and Frequency Table**). The variant p.R139C has been studied most extensively in patients receiving thiopurine therapy, thus, providing the strongest evidence for clinical implementation. Subsequent studies reported additional variants, most of which are rare, and their associations with clinical thiopurine toxicity do not rise to clinical actionability at this point, even though some showed decreased *NUDT15* activity in *in vitro*. For this reason, these variants (\*4 to \*9) are designated as unclear function but may be clarified as more data emerge.

Inherited TPMT deficiency is the primary genetic cause of thiopurine intolerance in Europeans and Africans, whereas risk alleles in *NUDT15* explain the majority of thiopurine-related myelosuppression in Asians and have been found in Hispanics.

### Genetic Test Interpretation

Genetic testing analyzes the DNA sequence at specific SNP locations in the *TPMT* and *NUDT15* genes (**Supplement**). Each named star (\*) allele is defined by the genotype at one or more specific loci (***TPMT* Allele Definition Table** (21, 22) and ***NUDT15* Allele Definition Table** (21, 23)) and is associated with a level of enzyme activity (***TPMT* Allele Functionality Table** (21, 22) and ***NUDT15* Allele Functionality Table** (21, 23)). **Table 1** summarizes the assignment of the likely *TPMT* and

NUDT15 phenotypes, based on the most common \* allele diplotypes, and these assignments are used to link genotypes with thiopurine prescribing recommendations. Of note, the phenotype of “possible intermediate metabolizer” has been introduced to the this guideline to describe an individual carrying one uncertain/unknown function allele PLUS one known no function allele, as this individual should be treated with “at least” the same precautions as would apply to an intermediate metabolizer. Although inactivating *TPMT* and *NUDT15* alleles have been identified in multiple populations (***TPMT* Frequency Table** (21, 22) and ***NUDT15* Frequency Table** (21, 23)), one of the limitations inherent in a commercial genotype-only test is that rare or previously undiscovered variants may not be included.

### **Available Genetic Test Options**

See **Supplementary material** and the Genetic Testing Registry (<https://www.ncbi.nlm.nih.gov/gtr/>) for more information on commercially available clinical testing options.

### **Incidental findings**

There are no diseases or phenotypic traits that have been linked to variation in *TPMT* or *NUDT15* in the absence of thiopurine treatment (2).

### **Linking genetic variability to variability in drug-related phenotypes**

There is substantial evidence linking *TPMT* and *NUDT15* genotype with phenotypic variability (see **Table S1**). Pre-emptive dose adjustments based on *TPMT* genotype have reduced thiopurine-induced adverse effects without compromising desired antitumor and immunosuppressive therapeutic effects in several clinical settings (**Table S1**). Similarly, retrospective studies strongly indicate that patients with loss-of-function *NUDT15* alleles are at excessive risk of thiopurine toxicity if the standard dose is administered. This body of evidence, rather than randomized clinical trials, provides the basis for most of the dosing recommendations in **Tables 2 and 3**.

## Therapeutic Recommendations

Thiopurines are used to treat malignant and non-malignant conditions, and thus the approach to dosing adjustments and the propensity to initiate therapy at higher vs. lower starting doses based on TPMT/NUDT15 status may differ depending on the clinical indication.. Thiopurines have a unique role in the treatment of many malignancies. The “normal” starting doses of thiopurines are generally “high” because they have been derived from trials which have been heavily weighted by the ~90% of the population who are wild-type for *TPMT* and *NUDT15* and receive maximal tolerable doses by the standards of anticancer treatment (hence, full doses should be given to those who are normal metabolizers for TPMT and NUDT15, **Tables 2 and 3**). Because the level of thiopurine tolerance is highly correlated with genetic ancestry (17), the “normal” starting doses can also vary by geographic regions and clinical practice.

***TPMT recommendation.*** If starting doses are already high (e.g., 75 mg/m<sup>2</sup> of mercaptopurine), as is true in some ALL treatment regimens, lower than normal starting doses should be considered in TPMT intermediate metabolizers (11, 15, 24, 25) and markedly reduced doses (10-fold reduction) should be used in TPMT poor metabolizers (26) (**Table 2**). This approach has decreased the risk of acute toxicity without compromising relapse rate in ALL (27). Even at these markedly reduced dosages, erythrocyte TGN concentrations in TPMT poor metabolizers remain well above those tolerated and achieved by the majority of patients (who are TPMT normal metabolizers) (4, 26).

In some nonmalignant conditions, alternative agents may be chosen for TPMT intermediate or poor metabolizers rather than reduced doses of thiopurines; if thiopurines are used, full starting doses are recommended for TPMT normal metabolizers, reduced doses (30-80% of target dose) in TPMT intermediate metabolizers (28, 29), and substantially reduced doses (or use of an alternative agent) in TPMT poor metabolizers (**Table 2**) (4, 30).



Some of the clinical data upon which dosing recommendations are based (**Table 2**) rely on measures of TPMT phenotype rather than genotype; however, because *TPMT* genotype is strongly linked to TPMT phenotype (5-7, 31), these recommendations apply regardless of the method used to assess TPMT status.

***NUDT15 recommendation.*** Similar to *TPMT*, tolerated mercaptopurine dosage is also correlated with the number of non-functional alleles of the *NUDT15* gene (17, 18). In fact, the degree of thiopurine intolerance (e.g., for mercaptopurine) is also largely comparable between carriers of *TPMT* vs. *NUDT15* decreased function alleles (17), although there remains a paucity of multi-ethnic studies examining both *TPMT* and *NUDT15* variants. Therefore, our *NUDT15* recommendations parallel those for *TPMT*. For *NUDT15* normal metabolizers (*NUDT15*\*1/\*1), starting doses do not need to be altered. For *NUDT15* intermediate metabolizers (e.g., *NUDT15*\*1/\*3, **Table 2**), reduced starting doses should be considered to minimize toxicity, particularly if the starting doses is high (e.g., 75 mg/m<sup>2</sup>/day for mercaptopurine). For *NUDT15* poor metabolizers (e.g., \*3/\*3), substantially reduced doses (e.g., 10 mg/m<sup>2</sup>/day of mercaptopurine) or the use of an alternative agent should be used (**Table 2**) (20).

As for *TPMT*, there has been some variability in the tolerated thiopurine dosages within *NUDT15* intermediate metabolizers, with a minority of individuals who do not seem to require significant dose reduction (17, 20). Therefore, genotype-guided prescribing recommendations apply primarily to starting doses; subsequent dosing adjustments should be made based on close monitoring of clinical myelosuppression (or disease-specific guidelines). In contrast, a full dose of mercaptopurine poses a severe risk of prolonged hematopoietic toxicity in *NUDT15* poor metabolizers and preemptive dose reductions are strongly recommended (32, 33).

The NUDT15 poor metabolizer phenotype is observed at a frequency of about one in every 50 patients of East Asian descent, which is more common than the TPMT poor metabolizer phenotype in Europeans, and thus genotyping NUDT15 in the Asian populations may be of particular clinical importance. NUDT15 deficiency is also more prevalent in individuals of Hispanic ethnicity, particularly those with high levels of Native American genetic ancestry (17).

***TPMT and NUDT15 recommendation.*** Figure 2 outlines the recommended course of action if both *TPMT* and *NUDT15* genotypes are known. There have been reports of patients with intermediate metabolizer status for both TPMT and NUDT15 (i.e., compound intermediate metabolizers), and there was a trend for a lower thiopurine tolerance in these individuals compared to intermediate metabolizers for only TPMT or NUDT15. The two genes are independent: the likelihood of an individual being an intermediate metabolizer for both genes depends upon the population frequencies for variant alleles. For example, given estimates of no function alleles for *NUDT15* of 11% and of no function alleles for *TPMT* of 2%, the frequency of the compound intermediate phenotype is estimated at 0.2%. However, the evidence for a different starting dosage recommendation for the compound intermediate metabolizers remains limited.

#### **Recommendations for Incidental Findings**

Not applicable.

#### **Other considerations**

If test results are available for only one gene (*TPMT* or *NUDT15*, but not both), prescribing recommendations based on that gene's results may be implemented, with the caveat that the other gene's results are missing and may have important implications. The higher frequency of decreased function *NUDT15* variants among individuals of Asian and Hispanic backgrounds and of *TPMT* variants in those with European and African backgrounds should be considered. In addition, there

may be other reasons underlying poor tolerance to thiopurines that are not related to *TPMT* or to *NUDT15* genetic variation.

Complementary clinical laboratory tests are available to measure thiopurine metabolites in erythrocytes: TGNs (for mercaptopurine, azathioprine, and thioguanine) and MeMPNs (or MeTIMP) for those on mercaptopurine or azathioprine (see **Supplement** for details on associations with *TPMT*). Erythrocyte TGNs or MeMPNs are not related to *NUDT15* genotypes (34-36) because clinical assays do not distinguish among the mono-, di-, and tri-phosphate forms of active TGNs, but there is evidence that intermediate and poor metabolizers for *NUDT15* accumulate higher level of DNA-TG than normal metabolizers given the same mercaptopurine dosage (20). Thus, currently available erythrocyte therapeutic drug monitoring tests do not distinguish *NUDT15* metabolizer phenotypes.

***Implementation of this guideline.*** The guideline supplement contains resources that can be used within electronic health records (EHRs) to assist clinicians in applying genetic information to patient care for the purpose of drug therapy optimization (see *Resources to incorporate pharmacogenetics into an electronic health record with clinical decision support* sections of supplement).

## **POTENTIAL BENEFITS AND RISKS FOR THE PATIENT**

The benefits of pre-emptive *TPMT* testing are that doses that are customized based on *TPMT* status reduce the likelihood of acute myelosuppression without compromising disease control (4, 8, 24, 25). The risks would be that a proportion of *TPMT* intermediate metabolizers may spend a period of time at lower thiopurine doses than they can eventually tolerate, because only ~30-60% of *TPMT* heterozygous patients receiving conventional thiopurine doses experience severe myelosuppression (4, 8, 11). However, because steady state is reached in 2-4 weeks, any period of “under-dosing” should be short, and using this approach, at least in ALL and in inflammatory bowel disease, outcomes were not compromised (4, 8, 24, 25, 28).

Similar benefits are expected with pre-emptive *NUDT15* genotyping, especially for Asian patients, given that these variants have comparable effects as risk alleles in *TPMT*. At least in ALL, leukemia cells with loss-of-function *NUDT15* alleles are also more sensitive to mercaptopurine (20) and thus in theory *NUDT15* genotyped guided dosing would not compromise anti-leukemic efficacy of this drug.

A possible risk to the patient is an error in genotyping (4). Some *TPMT* and/or *NUDT15* variants may not be included in the genotype test used and patients with these variants may be assigned a "wild-type" (\*1) genotype by default. Thus, an assigned "wild-type" allele could potentially harbor a no or decreased function variant. Because genotypes are life-long test results, any such error could stay in the medical record for the life of the patient.

#### **CAVEATS: APPROPRIATE USE AND/OR POTENTIAL MISUSE OF GENETIC TESTS**

Most of the time, thiopurines are given orally daily for a period of at least several months. Genotype-based starting doses are just that—starting doses, and in most diseases, titration to the desired degree (or lack thereof) of myelosuppression is required. Thus, clinicians must continue to evaluate markers of disease progression and/or of myelosuppression to adjust thiopurine doses up or down from the genotype-directed starting doses. One caveat is that some serious long-term adverse effects (secondary tumors) have been associated with defective *TPMT* activity without necessarily causing serious acute myelosuppression; whether capping doses of thiopurines in those with a *TPMT* defect will decrease the risk of the late effect of secondary cancer is not known. Some adverse reactions to thiopurines, such as pancreatitis and hepatotoxicity, are not related to low *TPMT* activity.

The discovery and clinical implementation of *NUDT15* variants in thiopurine dosing is relatively recent and the exact impact of *NUDT15* genotype-guided dose adjustments on toxicity and efficacy are less clear compared to *TPMT*.

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## REFERENCES

- (1) Sandborn, W.J. Pharmacogenomics and IBD: TPMT and thiopurines. *Inflamm Bowel Dis* **10 Suppl 1**, S35-7 (2004).
- (2) Evans, W.E. Pharmacogenetics of thiopurine S-methyltransferase and thiopurine therapy. *Ther Drug Monit* **26**, 186-91 (2004).
- (3) Weinshilboum, R. Inheritance and drug response. *N Engl J Med* **348**, 529-37 (2003).
- (4) Ford, L.T. & Berg, J.D. Thiopurine S-methyltransferase (TPMT) assessment prior to starting thiopurine drug treatment; a pharmacogenomic test whose time has come. *J Clin Pathol* **63**, 288-95 (2010).
- (5) Schaeffeler, E. *et al.* Comprehensive analysis of thiopurine S-methyltransferase phenotype-genotype correlation in a large population of German-Caucasians and identification of novel TPMT variants. *Pharmacogenetics* **14**, 407-17 (2004).

- (6) Yates, C.R. *et al.* Molecular diagnosis of thiopurine S-methyltransferase deficiency: genetic basis for azathioprine and mercaptopurine intolerance. *Ann Intern Med* **126**, 608-14 (1997).
- (7) Liu, C. *et al.* Genomewide Approach Validates Thiopurine Methyltransferase Activity Is a Monogenic Pharmacogenomic Trait. *Clin Pharmacol Ther* **101**, 373-81 (2017).
- (8) Relling, M.V. *et al.* Mercaptopurine therapy intolerance and heterozygosity at the thiopurine S-methyltransferase gene locus. *J Natl Cancer Inst* **91**, 2001-8 (1999).
- (9) Nygaard, U., Toft, N. & Schmiegelow, K. Methylated metabolites of 6-mercaptopurine are associated with hepatotoxicity. *Clin Pharmacol Ther* **75**, 274-81 (2004).
- (10) Evans, W.E. *et al.* Preponderance of thiopurine S-methyltransferase deficiency and heterozygosity among patients intolerant to mercaptopurine or azathioprine. *J Clin Oncol* **19**, 2293-301 (2001).
- (11) Stocco, G. *et al.* Genetic polymorphism of inosine triphosphate pyrophosphatase is a determinant of mercaptopurine metabolism and toxicity during treatment for acute lymphoblastic leukemia. *Clin Pharmacol Ther* **85**, 164-72 (2009).
- (12) Higgs, J.E., Payne, K., Roberts, C. & Newman, W.G. Are patients with intermediate TPMT activity at increased risk of myelosuppression when taking thiopurine medications? *Pharmacogenomics* **11**, 177-88.
- (13) Hartford, C. *et al.* Differential effects of targeted disruption of thiopurine methyltransferase on mercaptopurine and thioguanine pharmacodynamics. *Cancer Res* **67**, 4965-72 (2007).
- (14) Hosni-Ahmed, A., Barnes, J.D., Wan, J. & Jones, T.S. Thiopurine methyltransferase predicts the extent of cytotoxicity and DNA damage in astroglial cells after thioguanine exposure. *PLoS One* **6**, e29163 (2011).
- (15) Lennard, L. & Lilleyman, J.S. Individualizing therapy with 6-mercaptopurine and 6-thioguanine related to the thiopurine methyltransferase genetic polymorphism. *Ther Drug Monit* **18**, 328-34 (1996).
- (16) McBride, K.L., Gilchrist, G.S., Smithson, W.A., Weinshilboum, R.M. & Szumlanski, C.L. Severe 6-thioguanine-induced marrow aplasia in a child with acute lymphoblastic leukemia and inhibited thiopurine methyltransferase deficiency. *J Pediatr Hematol Oncol* **22**, 441-5 (2000).
- (17) Yang, J.J. *et al.* Inherited NUDT15 variant is a genetic determinant of mercaptopurine intolerance in children with acute lymphoblastic leukemia. *J Clin Oncol* **33**, 1235-42 (2015).
- (18) Yang, S.K. *et al.* A common missense variant in NUDT15 confers susceptibility to thiopurine-induced leukopenia. *Nat Genet* **46**, 1017-20 (2014).
- (19) Nielsen, S.N. *et al.* DNA-thioguanine nucleotide concentration and relapse-free survival during maintenance therapy of childhood acute lymphoblastic leukaemia (NOPHO ALL2008): a prospective substudy of a phase 3 trial. *Lancet Oncol* **18**, 515-24 (2017).
- (20) Moriyama, T. *et al.* NUDT15 polymorphisms alter thiopurine metabolism and hematopoietic toxicity. *Nat Genet* **48**, 367-73 (2016).
- (21) CPIC. *CPIC Guideline for Thiopurines and TPMT and NUDT15*.  
<<https://cpicpgx.org/guidelines/guideline-for-thiopurines-and-tpmt/>>.
- (22) PharmGKB. *Gene Reference Materials for TPMT*.  
<<https://www.pharmgkb.org/page/tpmtRefMaterials>>. Accessed January 1 2018.
- (23) PharmGKB. *Gene Reference Materials for NUDT15*.  
<<https://www.pharmgkb.org/page/nudt15RefMaterials>>.
- (24) Schmiegelow, K. *et al.* Thiopurine methyltransferase activity is related to the risk of relapse of childhood acute lymphoblastic leukemia: results from the NOPHO ALL-92 study. *Leukemia* **23**, 557-64 (2009).
- (25) Schmiegelow, K. *et al.* Long-term results of NOPHO ALL-92 and ALL-2000 studies of

- childhood acute lymphoblastic leukemia. *Leukemia* **24**, 345-54 (2010).
- (26) Evans, W.E., Horner, M., Chu, Y.Q., Kalwinsky, D. & Roberts, W.M. Altered mercaptopurine metabolism, toxic effects, and dosage requirement in a thiopurine methyltransferase-deficient child with acute lymphocytic leukemia. *J Pediatr* **119**, 985-9 (1991).
- (27) Relling, M.V., Pui, C.H., Cheng, C. & Evans, W.E. Thiopurine methyltransferase in acute lymphoblastic leukemia. *Blood* **107**, 843-4 (2006).
- (28) Meggitt, S.J., Gray, J.C. & Reynolds, N.J. Azathioprine dosed by thiopurine methyltransferase activity for moderate-to-severe atopic eczema: a double-blind, randomised controlled trial. *Lancet* **367**, 839-46 (2006).
- (29) Coenen, M.J. *et al.* Identification of Patients With Variants in TPMT and Dose Reduction Reduces Hematologic Events During Thiopurine Treatment of Inflammatory Bowel Disease. *Gastroenterology* **149**, 907-17 e7 (2015).
- (30) Sandborn, W.J. Rational dosing of azathioprine and 6-mercaptopurine. *Gut* **48**, 591-2 (2001).
- (31) Tamm, R. *et al.* Polymorphic variation in TPMT is the principal determinant of TPMT phenotype: A meta-analysis of three genome-wide association studies. *Clin Pharmacol Ther* **101**, 684-95 (2017).
- (32) Zhu, Y. *et al.* Combination of common and novel rare NUDT15 variants improves predictive sensitivity of thiopurine-induced leukopenia in children with acute lymphoblastic leukemia. *Haematologica*, (2018).
- (33) Ailing, Z., Jing, Y., Jingli, L., Yun, X. & Xiaojian, Z. Further evidence that a variant of the gene NUDT15 may be an important predictor of azathioprine-induced toxicity in Chinese subjects: a case report. *J Clin Pharm Ther* **41**, 572-4 (2016).
- (34) Moriyama, T. *et al.* The effects of inherited NUDT15 polymorphisms on thiopurine active metabolites in Japanese children with acute lymphoblastic leukemia. *Pharmacogenet Genomics* **27**, 236-9 (2017).
- (35) Lee, J.H. *et al.* Measurements of 6-thioguanine nucleotide levels with TPMT and NUDT15 genotyping in patients with Crohn's disease. *PLoS One* **12**, e0188925 (2017).
- (36) Asada, A. *et al.* NUDT15 R139C-related thiopurine leukocytopenia is mediated by 6-thioguanine nucleotide-independent mechanism in Japanese patients with inflammatory bowel disease. *J Gastroenterol* **51**, 22-9 (2016).
- (37) Anstey, A.V., Wakelin, S., Reynolds, N.J., British Association of Dermatologists Therapy, G. & Audit, S. Guidelines for prescribing azathioprine in dermatology. *Br J Dermatol* **151**, 1123-32 (2004).
- (38) Lichtenstein, G.R., Abreu, M.T., Cohen, R., Tremaine, W. & American Gastroenterological, A. American Gastroenterological Association Institute technical review on corticosteroids, immunomodulators, and infliximab in inflammatory bowel disease. *Gastroenterology* **130**, 940-87 (2006).
- (39) Krynetski, E.Y. & Evans, W.E. Pharmacogenetics of cancer therapy: getting personal. *Am J Hum Genet* **63**, 11-6 (1998).
- (40) Kaskas, B.A. *et al.* Safe treatment of thiopurine S-methyltransferase deficient Crohn's disease patients with azathioprine. *Gut* **52**, 140-2 (2003).
- (41) Zaza, G. *et al.* Thiopurine pathway. *Pharmacogenet Genomics* **20**, 573-4 (2010).



## Figure Legend

**Figure 1:** Metabolism of azathioprine, thioguanine, and mercaptopurine (41). Permission has been given by PharmGKB and Stanford to use figure (<https://www.pharmgkb.org/pathway/PA2040>).

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**Figure 2.** Recommended Starting Doses of Thiopurines by TPMT and NUDT15 phenotype

<sup>a</sup> Whether a dose reduction is recommended from the starting dose depends on the level of the standard starting dose; for example, if the standard starting dose of mercaptopurine is 75 mg/m<sup>2</sup>/day or higher, then a lower starting dose may be considered in intermediate metabolizers and would be recommended in poor metabolizers, whereas if the starting dose is 50 mg/m<sup>2</sup>/day or lower, a reduced starting dose may not be necessary in intermediate metabolizers.

<sup>b</sup> See Table 2 for recommendation.

<sup>c</sup> For patients who are IM for both TPMT and NUDT15, further dose reduction might be needed compared to those who are only IM with respect to one gene (TPMT or NUDT15).

- Clinical Pharmacogenetics Implementation Consortium (CPIC) guideline for thiopurine dosing based on TPMT and NUDT15 genotypes: 2018 update
- CPIC Thiopurine Supplemental Tables



**Table 1. Assignment of likely TPMT and NUDT15 phenotypes based on genotypes**

Assignment of likely TPMT phenotypes based on genotypes		
Likely Phenotype	Genotypes	Examples of diplotypes
Normal metabolizer	an individual carrying normal function alleles	*1/*1
Intermediate metabolizer	an individual carrying one normal function allele PLUS one no function allele	*1/*2, *1/*3A, *1/*3B, *1/*3C, *1/*4
Possible Intermediate metabolizer	an individual carrying one uncertain/unknown function allele PLUS one no function allele	*2/*8, *3A/*7
Poor metabolizer	an individual carrying two no function alleles	*3A/*3A, *2/*3A, *3A/*3C, *3C/*4, *2/*3C, *3A/*4
Indeterminate	An individual carrying two uncertain/unknown function alleles OR one normal function allele plus one uncertain allele function allele	*6/*8  *1/*8
Assignment of likely NUDT15 phenotypes based on genotypes		

Normal metabolizer	an individual carrying two normal function alleles	*1/*1
Intermediate metabolizer	an individual carrying one normal function allele PLUS one no function allele	*1/*2, *1/*3
Possible Intermediate metabolizer	an individual carrying one uncertain function allele PLUS one no function allele	*2/*5, *3/*6
Poor metabolizer	an individual carrying two no function alleles	*2/*2, *2/*3, *3/*3
Indeterminate	one normal function allele PLUS one uncertain function allele OR two uncertain function alleles	*1/*4, *1/*5  *4/*5, *5/*6

<sup>†</sup> see TPMT and NUDT15 frequency table (21-23) for estimates of phenotype frequencies among different ethnic/geographic groups

**Table 2. Recommended Dosing of Thiopurines by TPMT phenotype**

	Mercaptopurine			Azathioprine		Thioguanine		
<b>Phenotype</b>	<i>Implications for mercaptopurine and azathioprine phenotypic measures</i>	<i>Dosing recommendations for mercaptopurine</i>	<i>Classification of recommendations</i>	<i>Dosing recommendations for azathioprine</i>	<i>Classification of recommendations</i>	<i>Implications for thioguanine phenotypic measures</i>	<i>Dosing recommendations for thioguanine</i>	<i>Classification of recommendations</i>
<b>TPMT Normal metabolizer</b>	Lower concentrations of TGN metabolites, higher MeTIMP, this is the 'normal'	Start with normal starting dose <sup>a</sup> (e.g. 75 mg/m <sup>2</sup> /day or 1.5 mg/kg/day) and adjust doses of mercaptopurine (and of any other	Strong	Start with normal starting dose <sup>a</sup> (e.g. 2-3 mg/kg/day) and adjust doses of azathioprine based on disease-specific guidelines. Allow	Strong	Lower concentrations of TGN metabolites, but note that TGN after thioguanine	Start with normal starting dose <sup>a</sup> (e.g. 40-60 mg/m <sup>2</sup> /day) and adjust doses of thioguanine and of other myelosuppressive	Strong

	pattern	myelosuppressive therapy) without		2 weeks to reach steady-state after each dose adjustment (4, 30, 37).		are 5-10X higher than TGN after mercaptopurine or azathioprine	therapy without any special emphasis on thioguanine. Allow 2 weeks to reach steady-state after each dose adjustment (4, 16).	
	Normal risk of thiopurine-related leukopenia, neutropenia, myelosuppression	any special emphasis on mercaptopurine compared to other agents. Allow at least 2 weeks to reach steady-state after each dose adjustment (4, 27, 30).				Normal risk of thiopurine-related leukopenia, neutropenia, myelosuppression		
<b>TPMT Intermediate</b>	Moderate to high	Start with reduced starting doses	Strong	Start with reduced starting doses	Strong	Moderate to high	Start with reduced doses (50% to 80%	Moderate

<b>metabolizer</b> <b>OR</b> <b>TPMT Possible</b> <b>intermediate</b> <b>metabolizer</b>	concentrations of TGN metabolites; low concentrations of MeTIMP Increased risk of thiopurine- related leukopenia, neutropenia, myelosuppress ion	(30%-80% of normal dose) if normal starting dose <sup>a</sup> is $\geq 75$ mg/m <sup>2</sup> /day or $\geq 1.5$ mg/kg/day (e.g. start at 25-60 mg/m <sup>2</sup> /day or 0.45- 1.2 mg/kg/day) and adjust doses of mercaptopurine based on degree of myelosuppression and disease- specific guidelines. Allow 2-4 weeks to		(30%-80% of normal dose) if normal starting dose <sup>a</sup> is 2-3 mg/kg/day, (e.g. 0.6 – 2.4 mg/kg/day), and adjust doses of azathioprine based on degree of myelosuppression and disease-specific guidelines. Allow 2-4 weeks to reach steady-state after each dose		concentrations of TGN metabolites; but note that TGN after thioguanine are 5-10X higher than TGN after mercaptopurin e or azathioprine Increased risk of thiopurine- related	of normal dose) if normal starting dose <sup>a</sup> is $\geq 40$ -60 mg/m <sup>2</sup> /day (e.g. 20-48 mg/m <sup>2</sup> /day) and adjust doses of thioguanine based on degree of myelosuppression and disease-specific guidelines. Allow 2- 4 weeks to reach steady-state after each dose adjustment. If myelosuppression	
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		reach steady-state after each dose adjustment. If myelosuppression occurs, and depending on other therapy, emphasis should be on reducing mercaptopurine over other agents (4, 11, 15, 24, 25, 27, 30, 38, 39). If normal starting dose is already <75mg/m <sup>2</sup> /day or		adjustment (4, 30, 37, 38).		leukopenia, neutropenia, myelosuppression	occurs, and depending on other therapy, emphasis should be on reducing thioguanine over other agents (4, 16).	
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		< 1.5mg/kg/day, dose reduction may not be recommended.						
<b>TPMT Poor metabolizer</b>	Extremely high concentrations of TGN metabolites; fatal toxicity possible without dose decrease; no MeTIMP metabolites	For malignancy, start with drastically reduced doses (reduce daily dose <sup>a</sup> by 10-fold and reduce frequency to thrice weekly instead of daily, e.g. 10 mg/m <sup>2</sup> /day given just 3 days/week) and adjust doses of	Strong	For non-malignant conditions, consider alternative non- thiopurine immunosuppressan t therapy.  For malignancy, start with drastically reduced doses (reduce daily	Strong	Extremely high concentrations of TGN metabolites; fatal toxicity possible without dose decrease  Greatly increased risk	Start with drastically reduced doses (16) (reduce daily dose <sup>a</sup> by 10-fold and dose thrice weekly instead of daily) and adjust doses of thioguanine based on degree of myelosuppression and disease-specific guidelines. Allow 4- 6 weeks to reach	Strong

	Greatly increased risk of thiopurine-related leukopenia, neutropenia, myelosuppression	mercaptopurine based on degree of myelosuppression and disease-specific guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment. If myelosuppression occurs, emphasis should be on reducing mercaptopurine over other agents. For non-malignant		dose <sup>a</sup> by 10-fold and dose thrice weekly instead of daily) and adjust doses of azathioprine based on degree of myelosuppression and disease-specific guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment (28, 30, 37, 38, 40).		of thiopurine-related leukopenia, neutropenia, myelosuppression	steady-state after each dose adjustment. If myelosuppression occurs, emphasis should be on reducing thioguanine over other agents. For non-malignant conditions, consider alternative non-thiopurine immunosuppressant therapy (4).	
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		conditions, consider alternative non-thiopurine immunosuppressant therapy (4, 26, 30, 38).						
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\*Normal starting doses vary by race/ethnicity and treatment regimens. If standard dose is below normal recommended dose, dose reduction might not be recommended for intermediate metabolizers.

Table 3. Recommended Dosing of Thiopurines by NUDT15 phenotype

		Mercaptopurine		Azathioprine		Thioguanine	
Phenotype	Implications for thiopurine phenotypic measures	Dosing recommendations for mercaptopurine	Classification of recommendations	Dosing recommendations for azathioprine	Classification of recommendations	Dosing recommendations for thioguanine	Classification of recommendations
<b>NUDT15 Normal metabolizer</b>	Normal risk of thiopurine-related leukopenia, neutropenia, myelosuppression	Start with normal starting dose <sup>a</sup> (e.g., 75mg/m <sup>2</sup> /day or 1.5mg/kg/day) and adjust doses of mercaptopurine (and of any other myelosuppressive	Strong	Start with normal starting dose <sup>a</sup> (e.g., 2-3 mg/kg/day) and adjust doses of azathioprine based on disease-specific guidelines.	Strong	Start with normal starting dose <sup>a</sup> (40-60 mg/day). Adjust doses of thioguanine and of other myelosuppressive therapy without any special	Strong

		therapy) without any special emphasis on mercaptopurine compared to other agents. Allow at least 2 weeks to reach steady-state after each dose adjustment (4, 27, 30).		Allow 2 weeks to reach steady-state after each dose adjustment (4, 30, 37).		emphasis on thioguanine. Allow 2 weeks to reach steady-state after each dose adjustment (4, 16).	
<b>NUDT15 Intermediate metabolizer OR</b>	Increased risk of thiopurine-related leukopenia,	Start with reduced starting doses (30%-80% of normal dose) if	Strong	Start with reduced starting doses (30%-80% of normal dose)	Strong	Start with reduced doses (50% to 80% of normal dose) if normal	Moderate

<b>Possible</b>	neutropenia,	normal starting		if normal		starting dose <sup>a</sup> is	
<b>NUDT15</b>	myelosuppression	dose <sup>a</sup> is $\geq 75$		starting dose <sup>a</sup> is		$\geq 40-60$	
<b>Intermediate</b>	n	mg/m <sup>2</sup> /day or $\geq$		2-3 mg/kg/day,		mg/m <sup>2</sup> /day (e.g.	
<b>metabolizer</b>		1.5 mg/kg/day		(e.g. 0.6 – 2.4		20-48 mg/m <sup>2</sup> /day)	
		(e.g. start at 25-60		mg/kg/day), and		and adjust doses	
		mg/m <sup>2</sup> /day or		adjust doses of		of thioguanine	
		0.45-1.2		azathioprine		based on degree	
		mg/kg/day) and		based on degree		of	
		adjust doses of		of		myelosuppression	
		mercaptopurine		myelosuppression		and disease-	
		based on degree		n and disease-		specific	
		of		specific		guidelines. Allow	
		myelosuppression		guidelines.		2-4 weeks to	
		and disease-		Allow 2-4 weeks		reach steady-state	
		specific		to reach steady-		after each dose	

		<p>guidelines. Allow 2-4 weeks to reach steady-state after each dose adjustment. If myelosuppression occurs, and depending on other therapy, emphasis should be on reducing mercaptopurine over other agents (4, 11, 15, 24, 25, 27, 30, 38, 39).</p>		<p>state after each dose adjustment (4, 30, 37, 38).</p>		<p>adjustment. If myelosuppression occurs, and depending on other therapy, emphasis should be on reducing thioguanine over other agents (4, 16).</p>	
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		If normal starting dose is already <75mg/m <sup>2</sup> /day or < 1.5mg/kg/day, dose reduction may not be recommended.					
<b>NUDT15 Poor metabolizer</b>	Greatly increased risk of thiopurine-related leukopenia, neutropenia, myelosuppression	For malignancy, initiate dose at 10 mg/m <sup>2</sup> /day and adjust dose based on myelosuppression and disease-specific		For non-malignant conditions, consider alternative non-thiopurine immunosuppressant therapy.	Strong	Reduce doses to 25% of normal dose <sup>a</sup> and adjust doses of thioguanine based on degree of myelosuppression and disease-	Strong

		<p>guidelines. Allow 4-6 weeks to reach steady state after each dose adjustment. If myelosuppression occurs, emphasis should be on reducing mercaptopurine over other agents.</p> <p>For non-malignant conditions,</p>		<p>For malignant conditions, start with drastically reduced normal daily doses<sup>a</sup> (reduce daily dose by 10-fold) and adjust doses of azathioprine based on degree of myelosuppression and disease-specific</p>		<p>specific guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment. In setting of myelosuppression, emphasis should be on reducing thioguanine over other agents. For non-malignant conditions, consider</p>	
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		consider alternative non- thiopurine immunosuppressant therapy (4, 26, 30, 38).		guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment (28, 30, 37, 38, 40).		alternative non- thiopurine immunosuppressant therapy (4).	
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\*Normal starting doses vary by race/ethnicity and treatment regimens. If standard dose is below normal recommended dose, dose reduction might not be recommended for intermediate metabolizers.





