Concordance between glucose-6-phosphate dehydrogenase (G6PD) genotype and phenotype and rasburicase use in patients with hematologic malignancies

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Abstract
Phenotypic rather than genotypic tests remain the gold standard for diagnosing glucose-6-phosphate dehydrogenase (G6PD) deficiency. However, with increasing use of genomic arrays and whole exome or genome sequencing, G6PD genetic data are increasingly available. We examined the utility of G6PD genetic data in patients with hematologic malignancies and the association of G6PD genotype and phenotype with rasburicase-induced methemoglobinemia. We analyzed G6PD activity for 990 patients. Genotype data were available from the Affymetrix DMET array (n = 379), whole exome sequencing (n = 374), and/or the Illumina exome array (n = 634) for 645 patients. Medical records of 341 patients with methemoglobin measures were assessed for the administration of rasburicase. We observed 5 non-synonymous SNPs, 4 of which were known to be associated with deficient G6PD activity (WHO Class I–III). Genotyping 367 males resulted in a positive predictive value of 81.8% (47.8–96.8%), and two males with a Class I-III allele having normal activity both received a red blood cell transfusion prior to the activity assay. However, genotyping males had only 39.1% (20.5–61.2%) sensitivity. Two of the 12 heterozygous females had deficient G6PD activity. Rasburicase-induced methemoglobinemia occurred in 6 patients, 5 of whom had at least one Class I-III allele, despite 2 of these having normal G6PD activity. We conclude that although an apparent nondeficient genotype does not necessarily imply a normal phenotype, a deficient genotype result indicates a deficient phenotype in those without transfusions, and may be a useful adjunct to phenotype to prevent adverse drug reactions.

Introduction
Glucose-6-phosphate-dehydrogenase (G6PD) deficiency is a well-recognized pharmacogenetic trait. [1] G6PD functions as the rate-limiting step of the hexose monophosphate shunt, which maintains the supply of NADPH available to replenish the cellular stores of reduced glutathione used for detoxifying reactive oxygen species. [1, 2] G6PD is the only source of NADPH in erythrocytes, leaving erythrocytes prone to oxidative stress from either endogenous or exogenous sources, including medications. Thus, erythrocytes are especially sensitive to G6PD deficiency, [1–4] and patients with G6PD deficiency are susceptible to hemolytic anemia and methemoglobinemia. [3–6] G6PD deficiency is estimated to affect almost 5% of the global population. [7]

The gold-standard method to assess clinical G6PD status is quantitating G6PD activity in whole red blood cells through spectrophotometry. [8, 9] Over 180 variant genetic alleles of G6PD have been reported, and uptake of genetic
tests rather than phenotypic tests of activity has been slow. This is partly because genetic tests are thought to be poorly predictive in females, for whom X-linked mosaicism in red cells results in variable expression of G6PD activity. [10–13] and because G6PD status is often important to ascertain with short turn-around-time [14] and few point-of-care G6PD genetic tests have yet been developed. [15] Moreover, most low-activity G6PD alleles are rare, and the optimal set of alleles for genotyping has not been established [16] and may differ by ancestry. [7] The World Health Organization (WHO) classifies variant alleles into five different classes based on relative enzyme activity compared to the wildtype enzyme and the clinical presentation of deficiency. [16] The Clinical Pharmacogenetics Implementation Consortium (CPIC®) makes pharmacogenetic recommendations based on the assumption that genotypes may be available pre-emptively, and has assigned four different likely G6PD phenotypes based on the presence of WHO class alleles: normal (Class IV), deficient (Classes II and III), deficient with chronic nonspherocytic hemolytic anemia (Class I), and variable (heterozygous females with one Class IV and one deficient (Class I–III) allele) (Supplemental Table S1). [6]

As array-based, whole exome and whole genome sequencing becomes more common, [6, 7, 17] clinicians will be faced with G6PD genetic information that is generated “incidentally.” These data will identify patients who carry class I–III low-function G6PD alleles, making it potentially possible to assign G6PD status to individuals based on genomic information alone. It is unknown how well genetic testing accurately predicts G6PD deficiency. We tested for this concordance of genotype with phenotype in an American pediatric population with hematological malignancies. We also tested how G6PD genotype and phenotype were associated with rasburicase-induced methemoglobinemia in pediatric patients receiving treatment for acute leukemia.

**Methods**

**G6PD activity**

We retrospectively analyzed 990 pediatric patients with hematological malignancies (acute myeloid leukemia, acute lymphoblastic leukemia, chronic myeloid leukemia, or other hematological malignancies) enrolled in St. Jude Children’s Research Hospital from 1993–2013 on research protocols with institutional review board approval who had G6PD activity measured in their blood as part of clinical care. Four different quantitative spectrophotometric G6PD assays were used over time by St. Jude’s Department of Pathology to measure G6PD activity for 990 patients, with different ranges defining “normal activity” by date: before December 1996, 4.6–13.5 units/g Hb; between December 1996 and August 2002, 7–20.5 units/g Hb; between August 2002 and September 2004, 10.8–16.2 units/g Hb; and after September 2004, 6.3–18.5 units/g Hb. For patients with leukocyte counts > 100 x 10⁶ cells/μl, a buffy-coat-free sample was used to measure G6PD activity; however, this could not be confirmed for samples prior to 1996. Twelve patients were excluded from analysis as described in the results section for a total of 978 patients in the cohort (Fig. 1). Patients were considered deficient if their activity fell below the lower normal limit for the quantitative spectrophotometric assay.

**G6PD genotype**

DNA was extracted from peripheral blood after patients achieved clinical remission. 645 patients had their genotype assessed using one or more of the following platforms: Affymetrix Drug Metabolizing Enzymes and Transporters (DMET) array (Santa Clara, CA) (n = 379) [18, 19], Illumina HumanExome BeadChip (Exomechip) (San Diego, CA) (n = 634) [20], and whole exome sequencing (WES) (n = 374) [21] (Supplemental Fig. S1). The Affymetrix DMET array interrogated 6 nonsynonymous single nucleotide polymorphisms (SNPs) (A (p.N126D), Canton (p.R459L), Chatham (p.A335T), Mediterranean (p.S188F), Sao Borja (p.D113N), 1 with unknown function (p.V77M)). The Illumina Exomechip interrogated 8 nonsynonymous SNPs (A, 968 (p.L323P), Asahi (p.V68M), Mediterranean (p.S188F), Malaga (p.D181V), Sierra Leone (p.R104H), Seattle (p.D282H), Mira d’Aire (p.D350H), and one with unknown function (p.Q11H)) (Supplemental Fig. S2, Table S2). Nonsynonymous coding SNPs were classified according to the WHO classification, [16] and patients were assigned a phenotype based on the CPIC guidelines for rasburicase. [6]

**G6PD genotype and phenotype concordance**

We estimated the positive predictive value (PPV), negative predictive value (NPV), sensitivity, and specificity when using the patient’s genotype to predict G6PD phenotype, with 95% confidence intervals also estimated. Fisher’s exact test was used to compare the frequency of rasburicase-induced methemoglobinemia.

**Rasburicase and methemoglobinemia**

Of the 978 patients with evaluable G6PD activity, 341 had at least one methemoglobin level measured during therapy. Each medical record was assessed for administration of rasburicase, and we compared the dates of the methemoglobin level to
rasburicase administration. Methemoglobinemia was defined as methemoglobin > 3%, which was three times the standard deviation plus the mean for methemoglobin levels in our cohort. For patients with methemoglobinemia in the absence of rasburicase administration, we reviewed records to identify any other drugs known to cause methemoglobinemia, such as sulfamethoxazole/trimethoprim and nitric oxide.

Results

G6PD activity

Of 990 patients with G6PD activity measures, 8 were excluded because we could not confirm if the buffy coat free method was performed, and their leukocyte counts were > 100 × 10^3 cells/µl (Fig. 1). Four patients had conflicting activity measurements and were excluded from the analysis (Supplemental Material [Concordance of multiple G6PD activity assays] and Fig. 1).

Of 978 patients with evaluable G6PD activity, 51 had deficient G6PD activity (5.2%). Twenty-three (45%) of the deficient patients were black males, and there were more than twice as many deficient males as deficient females (Table 1). The mean (SD) age for patients with deficient G6PD activity was 8.5 ± 5.4 years, and the mean age for patients with normal G6PD activity was 8.0 ± 5.4 years.

G6PD genotype

We observed variant genotypes at 5 nonsynonymous SNPs, 4 of which were part of Class II or III alleles in 645 patients...
with genotype data: A variant (p.N126D), Asahi variant (p.V68M), Viangchan variant (p.V291M), A-968 variant (p.L323P), and Pawnee variant (p.R439P) (Fig. 2, Supplemental Table S2). Two of the observed SNPs (Viangchan and Pawnee) were interrogated only by WES. Through WES, we found an additional 4 synonymous SNPs, 4 SNPs and 1 insertion/deletion (indel) in the 3’ untranslated region (UTR), 5 SNPs in the 5’ UTR, and 2 intronic SNPs (Supplemental Table S3). All variants were found in frequencies consistent with published allele frequencies (Supplemental Tables S2 and S3). All 645 patients were able to be assigned to a phenotype based on genotype (Fig. 1). Patients who had only the A variant were considered normal, as this is considered to be a Class IV allele.

Of 367 males, 11 were considered deficient based on genotype (hemizygous for a Class I–III allele). None of the 279 females were homozygous for a Class I–III allele, and thus none were considered deficient based on genotype. 12 females were heterozygous for a Class I–III allele and were assigned to have a “variable phenotype” (i.e., not assignable as normal or deficient) based on genotype.

### Table 1 Frequency of G6PD deficiency in pediatric patients with hematological malignancies by sex and race

|                  | Normal G6PD activity | Deficient G6PD activity |
|------------------|----------------------|-------------------------|
|                  | n    | %     | n    | %     |
| Male             |      |       |      |       |
| White            | 347  | 97.2% | 10   | 2.8%  |
| Black            | 131  | 85.1% | 23   | 14.9% |
| Other’           | 49   | 96.1% | 2    | 3.9%  |
| Total            | 527  | 93.8% | 35   | 6.2%  |
| Female           |      |       |      |       |
| White            | 262  | 98.1% | 5    | 1.9%  |
| Black            | 104  | 93.7% | 7    | 6.3%  |
| Other’           | 34   | 89.5% | 4    | 10.5% |
| Total            | 400  | 96.2% | 16   | 3.8%  |
| Both genders     | 927  | 94.8% | 51   | 5.2%  |

*Race self-reported; confirmed with genotype

‘Other includes Hispanics, Asians, and patients with mixed racial background

### G6PD genotype and phenotype concordance

Of the 11 males deficient according to genotype, 9 had deficient G6PD activity (PPV = 81.8% 47.8–96.8%), and of the 23 genotyped males with deficient G6PD activity, 9 had a Class I–III allele (sensitivity = 39.1% (20.5–61.2%)). Of the 356 males with a Class IV allele, 342 had normal G6PD activity (NPV = 96.1% (93.3–97.7%)). 342 out of 344 males with normal activity had a Class IV allele (specificity = 99.4% (97.7–99.9%)) (Fig. 3 and Supplemental Table S4). The two male patients with a Class I–III allele with normal activity had received a transfusion two days and 44 days prior to the G6PD activity assay respectively.

![Fig. 2 Five nonsynonymous SNPs in the G6PD gene were observed. Lines represent G6PD variants reported in the CPIC guidelines [6], and color indicates the WHO classification. SNPs interrogated in our cohort are listed by their codon number, amino acid substitution, and common name. The colored circles indicate which platforms include each SNP. The number in the circles indicates how many patients interrogated on that platform harbored that variant SNP. The gray line represents the whole exome sequencing coverage plotted against the left y-axis. The purple shading indicates the N-terminus domain, and the green indicates the C-terminus domain. We observed four Class I–III variants and one Class IV variant](image-url)
which may have artificially increased the apparent G6PD activity (Fig. 3).}

**Fig. 3** Genotyping G6PD has 81.8% positive predictive value and 39.1% sensitivity to predict G6PD phenotypic deficiency. The assigned phenotype based on genotype according to the CPIC guidelines is in parentheses in the flowchart. Nine of the 23 males with deficient activity were also deficient according to genotype, resulting in a sensitivity of 39.1% (20.5–61.2%). Nine of the 11 males with a Class I–III allele had deficient G6PD activity, resulting in a positive predictive value of 81.8% (47.8–96.8%). The two patients with a Class I–III allele with normal G6PD activity had received a transfusion (***) prior to the activity measure. The 95% confidence interval is listed in parentheses.

**Fig. 4** G6PD genotype in females does not correspond well with phenotype. Of 279 females, 268 had normal and 11 had deficient activity. No female was homozygous for a Class I–III allele, and 10/268 (3.7%) of those with normal activity and 2/11 (18%) of those with deficient activity carried one Class I–III allele. Of the 12 “variable” females (those heterozygous for a Class I–III allele), only 2 had deficient activity (16.7%). 258 out of 267 patients with only Class IV alleles had normal G6PD activity, resulting in a negative predictive value of 96.6% (93.5–98.3%).

Since there were no females homozygous for a Class I–III allele, a positive predictive value, sensitivity, and specificity could not be estimated for females. Of the 12 patients assigned as having a variable phenotype based on genotype, 2 (16.7%) had deficient G6PD activity. Genotyping females had a negative predictive value of 96.6% (93.5–98.3%): 258 out of 267 female patients with only Class IV alleles had normal G6PD activity (Fig. 4 and Supplemental Table S5).

We expected false negative results to occur because of the possible existence of Class I–III alleles that were not interrogated. All 14 males who had deficient activity but did not have a class I–III allele were interrogated by whole exome sequencing, which failed to detect any nonsynonymous SNPs in 12, and detected only the A (p.N126D) variant (a WHO class IV allele) in the other 2 males.
Fig. 5 Sixty patients with G6PD activity also had at least one methemoglobin measurement and received rasburicase at some point in therapy. Ten of these patients had deficient G6PD activity, and 50 had normal activity. The patients were then classified according to genotype. Eighteen patients had a methemoglobin level at the time of rasburicase administration: 3 patients with deficient activity and 15 patients with normal activity. The three patients with deficient activity all had a Class I-III allele and developed methemoglobinemia (>3%) after rasburicase administration. Of the 15 patients with normal activity who had evaluable methemoglobin after rasburicase, 3 developed methemoglobinemia: one male patient hemizygous for a Class I-III allele (this is the patient (**) who had received a transfusion prior to the G6PD activity assay); one was a female patient heterozygous for a Class I-III allele; and one was a female who did not have a Class I-III allele. Thus, of the 18 patients who had methemoglobin measurements at the time of rasburicase administration, 6 patients developed methemoglobinemia after rasburicase, and 5 of these 6 had a Class I-III allele

(Supplemental Table S4). We also found false negatives among females. Of the nine deficient females without a Class I-III allele, all were interrogated by WES, which did not detect any nonsynonymous variants in eight females, and detected the A variant alone in one female (Supplemental Table S5).

**Rasburicase and methemoglobinemia**

Of the 341 patients with at least one methemoglobin measurement in their medical records (196 males, 145 females), 320 had normal G6PD activity and 21 had deficient G6PD activity (Fig. 1). Sixty of the 341 patients received rasburicase (10 patients with deficient G6PD activity and 50 patients with normal G6PD activity), and 18 patients had methemoglobin measurements at the time of rasburicase administration (3 patients with deficient activity and 15 patients with normal activity) (Fig. 5) (Supplemental Fig. S3 for males and Supplemental Fig. S4 for females). All three of the patients with deficient G6PD activity who had an evaluable methemoglobin measurement after rasburicase administration developed rasburicase-induced methemoglobinemia, and each had a Class I-III allele. Three of the fifteen patients with normal G6PD activity and with documented methemoglobin measurements at the time of rasburicase administration developed methemoglobinemia after rasburicase. Two of these patients had at least one Class I-III allele: a male hemizygous for a Class I-III allele who had received a transfusion 44 days prior to the activity assay and a female heterozygous for a Class I-III allele. The third patient was a black female who did not have a Class I-III allele by Exomechip array. The other 12 patients with normal G6PD activity who had only Class IV alleles or did not have any genotype information available did not develop methemoglobinemia after rasburicase administration (9 males and 3 females). In summary, 6 patients developed rasburicase-induced methemoglobinemia, and 5 of these 6 had a Class I-III allele (Fig. 5). Among patients with normal G6PD activity, methemoglobinemia after rasburicase administration was more common (2 of 2) in those with a Class I-III allele than in those without a Class I-III allele (1 of 13) ($p = 0.029$).

We then examined all 341 patients with G6PD activity and at least one methemoglobin measurement, regardless of whether they ever received rasburicase (Fig. 6 and Supplemental Fig. S5 for males and Supplemental Fig. S6 for females). Three patients with deficient G6PD activity (all of whom also had Class I–III alleles) developed
341 patients with G6PD Activity and Methb levels

21 patients with deficient G6PD activity
- 3 patients with high Methb
- 3 patients received rasburicase the day before elevated Methb
- 1/3 had a Class I-III allele

24 patients with high Methb
- 1 female with a Class I-III allele
- 1 male with a Class I-III allele and Transfusion
- 3 patients received rasburicase the day of elevated Methb measurement

200 patients with normal Methb
- 7 patients received another medication (nitric oxide (5), dapsone (1), lepirudin (1))
- 4 patients received another medication (phenazopyridine (1), dapsone (2), lidocaine (1))

311 patients with normal G6PD Activity
- 11 patients had only Class IV alleles
- 10 patients did not receive rasburicase at the time of the Methb measurement
- 12 patients received rasburicase at the time of the Methb
- 42 patients received rasburicase

Fig. 6 341 patients had at least one methemoglobin measurement in their medical record, 21 of whom had deficient G6PD activity. 27 patients developed methemoglobinemia at some point during therapy (12 males and 15 females). All 3 patients with deficient G6PD activity developed elevated methemoglobinemia (>3%) after rasburicase administration, and all had a class I-III allele. Of the 24 patients with normal G6PD activity who developed methemoglobinemia at any time during therapy, 3 experienced methemoglobinemia after rasburicase administration. The one male was hemizygous for a Class I-III allele (he had received a transfusion prior to the activity assay**). One female was heterozygous for a Class I-III allele, and one female did not have a Class I-III allele (but was interrogated by the Exomechip array only). Of the other 21 patients with normal G6PD activity who experienced methemoglobinemia, 10 patients received another medication associated with methemoglobinemia.

In 367 males, genetic testing yielded a positive predictive value of 81.8% (47.8–96.8%), a negative predictive value of 96.1% (93.3–97.7%), sensitivity of 39.1% (20.5–61.2%), and specificity of 99.4% (98.1–99.9%). Genetic testing in 279 females yielded a negative predictive value of 96.6% (93.5–98.3%). No female in our patient population was homozygous for a Class I–III allele, and 2 of 12 (16.7%) heterozygous females had deficient activity. Rasburicase-induced methemoglobinemia occurred in 6 patients, 5 of whom had a Class I–III allele.

Despite extensive characterization of the G6PD gene, phenotypic enzyme activity is currently the gold standard for determining G6PD deficiency. However, various factors can influence activity assay results. Mature erythrocytes have approximately 50 times less G6PD activity than reticulocytes in the same individual; thus, G6PD activity measurements taken in the setting of reticulocytosis can be artificially high in G6PD-deficient patients [32–34]. Although leukocytes usually contribute to a very small fraction of the measured G6PD activity, they could account for a substantial fraction in the setting of hyperleukocytosis (>100 × 10^9 cells/µL) and anemia, [35, 36] and for these cases, it is recommended to use a Buffy coat free method. Red blood cell transfusions can also influence a G6PD activity measure. The lifespan of transfused red blood cells is approximately 60 days [37], and an activity measurement within this time frame could be reflective of the activity of the red blood cells from the donor as well as from the

**Discussion**

The prevalence of G6PD deficiency in 978 pediatric patients with hematologic malignancies was 5.2%, which is consistent with the prevalence in the general population. [7]
patient. [6] In addition to patient factors, temperature and sample handling can also affect the results of activity assays. [9] A genetic test could avoid some of the artifacts inherent in measuring G6PD activity in blood.

Even if genetic variants are adequately interrogated, there are inherent limitations to genetic test for assigning G6PD status. It is difficult to predict activity in heterozygous females due to X-linked mosaicism, in which one copy of the gene is randomly inactivated in cells. [11–13] Most heterozygotes have more normal G6PD cells than deficient G6PD cells, suggesting a cell selection bias. [38] Age also plays a role in the distribution of normal and deficient red blood cells: younger heterozygotes have more G6PD normal cells, whereas elderly patients have more G6PD deficient cells. [39, 40] In our cohort, 2 of 12 (16.7%) heterozygous females had deficient G6PD activity, which is consistent with reports in the literature. [10, 41]

False positive results in males (deficient genotype with normal activity) were not expected. The only two male patients in our cohort who had normal activity despite having a Class I–III allele had received red blood cell transfusions 44 days and 2 days prior to the activity assay, and thus activity may have reflected that of the red blood cells from the donor. One of these two patients developed methemoglobinemia after rasburicase administration, which also further suggests this patient was actually G6PD deficient. Hence our potential true positive predictive value for males could be 100%, given that both discordant patients may have had artificially elevated G6PD activity.

While a positive deficient genotype result should indicate a deficient phenotype, a negative deficient genotype (i.e., no detectable Class I–III alleles) does not necessarily indicate normal activity, because any genetic platform may fail to detect some important variants. [6] False negatives (G6PD deficient activity but without a Class I-III allele) were expected, especially in patients without whole exome sequencing. In our cohort, the sensitivity in males was only 39.1% (20.5–61.2%). There is the possibility that silent or noncoding variants may contribute to G6PD deficiency (Supplemental Material (Silent and noncoding variants in the G6PD gene)); however, we were unable to confirm that noncoding variants accounted for any of our cases of G6PD deficiency in the absence of Class I–III alleles (data not shown).

This is one of few large-scale G6PD genotype-phenotype studies done in an American population and the first in pediatric patients undergoing treatment for hematological malignancies. A study of neonates in Pennsylvania which genotyped 5 variants (A, Asahi, Mediterranean, Kaiping, and Canton) in over 4000 patients had similar genotype/phenotype concordance rates as our study. [17] Most other studies have focused on specific populations and on the alleles most common to that population. In these studies, even if whole exome sequencing was performed, 6–28% of deficient patients did not have a missense variant detected. [42–45] Thus, it seems that factors other than G6PD coding variants may cause G6PD deficiency.

G6PD activity was measured without regard to transfusion history, a potential limitation of our retrospective study of clinical phenotypes. G6PD activity measurements were made based on clinical considerations (e.g., the need to consider rasburicase for tumor lysis syndrome) and so may not have been optimally timed relative to hematologic status. If a patient has normal G6PD activity and receives a transfusion from a G6PD-deficient donor, the transfusion should not be enough for the patient to present as deficient. However, a red blood cell transfusion from a donor with normal G6PD activity could artificially increase the G6PD activity of a G6PD deficient patient. Therefore, the activity assay may have missed some G6PD deficient patients, as we believe it did with the two males hemizygous for a Class I–III allele who had normal G6PD activity after a red blood cell transfusion. These were the only patients we expected to be G6PD deficient according to genotype who had normal G6PD activity. A related limitation of this study is that data were limited to what was in the medical record, which may not have accurately reflected all red blood cell transfusions given outside of St. Jude. Methemoglobin levels were also limited to those patients who had their methemoglobin levels monitored for a clinical reason, which was not always clear from their medical record.

Rasburicase is a recombinant urate oxidase enzyme approved for the prevention and treatment of hyperuricemia of tumor lysis syndrome. As rasburicase breaks down uric acid to allantoin, it also produces hydrogen peroxide, an oxidizing agent, as a byproduct. [1, 46, 47] Rasburicase-induced methemoglobinemia is a potentially serious adverse effect that has been documented in several patients with G6PD deficiency. [48, 49] and rasburicase is thus contraindicated in patients with G6PD deficiency. A few patients in our cohort received rasburicase and were later determined to be G6PD deficient. We found that rasburicase-associated methemoglobinemia occurred in 6 patients, 5 of whom had at least one Class I–III allele. Importantly, performing only a phenotypic assay would have missed two of these patients: one hemizygous male who had received a blood transfusion prior to the activity assay and one heterozygous female with normal activity at the time of rasburicase administration. This indicates that having a fraction of deficient erythrocytes, even with a normal G6PD activity measure, could potentially be enough to predispose to methemoglobinemia after exposure to a strong oxidative trigger, like rasburicase. Similarly, in a retrospective analysis of a primaquine study, females heterozygous for the Mahidol variant, a class III allele, were more likely to require treatment for hemolytic anemia than wildtype females, even if the G6PD activity test was normal.
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In conclusion, while G6PD genotyping has its limitations, it can provide valuable information, and may become more widely available as whole exome and whole genome sequencing increases. For drugs such as rasburicase, the turnaround time for a G6PD genetic test is currently too long, and an activity test will still be required for obtaining a patient’s G6PD status quickly. However, genetic tests are rapidly evolving to include point-of-care tests, and G6PD genotype can be useful for confirmation of G6PD status when phenotypic tests may be compromised. When a Class I–III allele is observed in male patients or two Class I–III alleles in females, medications known to cause hemolytic anemia or methemoglobinemia in G6PD deficiency, such as rasburicase, should be avoided. For heterozygous female patients, caution must be used with strong oxidative drugs, even if the G6PD activity measure is normal, due to the potential for hemolytic anemia or methemoglobinemia. Genotyping has the advantage that results are not altered by the hematologic status of the patient, such as high white blood cell count, transfusion history, or anemic states. More work needs to be done to ensure greater sensitivity of DNA-based tests by increasing the alleles interrogated by commercially available genetic tests and to understand the mechanism of G6PD deficiency in the absence of a Class I–III allele.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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