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To the Editor:

Fisher and Cronstein are to be commended for their recent metaanalysis on methylenetetrahydrofolate reductase (MTHFR) polymorphisms and methotrexate (MTX) toxicity in rheumatoid arthritis (RA)1. Their work exposes the lack of sound pharmacogenetic data on MTX in RA and underscores that there is a long way to go before MTX pharmacogenetics is ready to guide clinical decision-making in RA.

In the discussion, the authors note the paucity of data in the literature to determine whether the presence or absence of a single-nucleotide polymorphism (SNP) has an effect on time to adverse events from MTX. We have reported that the presence of ABCC2 IVS 23+56 T>C, an intronic SNP in one of the ABC transporter genes, correlated with time to MTX discontinuation and/or dose decrease due to toxicity in Caucasian patients with RA (p < 0.0001)2. Although this is not a MTHFR SNP, to our knowledge, this is the only study to date to demonstrate such a genotype to time to MTX discontinuation association.

The authors also point out that none of the studies included in the analysis provided details on the racial background of the participants and emphasize the importance of race in influencing pharmacogenetics in different populations. Indeed, the SNP frequencies in several of the key genes in MTX pharmacogenetics including MTHFR differ significantly by race3,4. Further, we have shown that genotype-toxicity associations vary by race. In our cohort of Caucasians and African Americans with RA, we found that the MTHFR 677 C>T SNP was associated with MTX toxicity in African Americans with RA but not in Caucasians, and the intronic ABCC2 SNP with time to MTX discontinuation in Caucasians as described above, but not in African Americans with RA2. Others have demonstrated the MTHFR 677 C>T SNP to be associated with MTX toxicity in Caucasians and African Americans with RA4. These results supplement those of Fisher and Cronstein’s metaanalysis, which showed the MTHFR 677 C>T SNP to be associated with MTX toxicity in studies that examined populations of Caucasians and Asians, but did not include African Americans. We concede that the sample sizes of these studies were small, a problem all too common with almost all the studies related to MTX pharmacogenetics in RA.

Fisher and Cronstein’s metaanalysis once again demonstrates the futility of small-scale pharmacogenetic studies in providing meaningful results that can be applied to individualize patient therapy in clinical practice. Collaborative, multicenter, adequately powered studies, stratified by race, are needed to clarify the muddled state that exists in MTX pharmacogenetics today.

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