Variable Association of Reactive Intermediate Genes with Systemic Lupus Erythematosus in Populations with Different African Ancestry


ABSTRACT. Objective. Little is known about the genetic etiology of systemic lupus erythematosus (SLE) in individuals of African ancestry, despite its higher prevalence and greater disease severity. Overproduction of nitric oxide (NO) and reactive oxygen species are implicated in the pathogenesis and severity of SLE, making NO synthases and other reactive intermediate-related genes biological candidates for disease susceptibility. We analyzed variation in reactive intermediate genes for association with SLE in 2 populations with African ancestry.

Methods. A total of 244 single-nucleotide polymorphisms (SNP) from 53 regions were analyzed in non-Gullah African Americans (AA; 1432 cases and 1687 controls) and the genetically more homogeneous Gullah of the Sea Islands of South Carolina (133 cases and 112 controls). Single-marker, haplotype, and 2-locus interaction tests were computed for these populations.

Results. The glutathione reductase gene GSR (rs2253409; p = 0.0014, OR 1.26, 95% CI 1.09–1.44) was the most significant single SNP association in AA. In the Gullah, the NADH dehydrogenase NDUF4 (rs381575; p = 0.0065, OR 2.10, 95% CI 1.23–3.59) and NO synthase gene NOS1 (rs561712; p = 0.0072, OR 0.62, 95% CI 0.44–0.88) were most strongly associated with SLE. When both populations were analyzed together, GSR remained the most significant effect (rs2253409; p = 0.00072, OR 1.26, 95% CI 1.10–1.44). Haplotype and 2-locus interaction analyses also uncovered different loci in each population.

Conclusion. These results suggest distinct patterns of association with SLE in African-derived populations; specific loci may be more strongly associated within select population groups. (First Release May 1 2013; J Rheumatol 2013;40:842–9; doi:10.3899/jrheum.120989)

Key Indexing Terms:
SYSTEMIC LUPUS ERYTHEMATOSUS  AFRICAN AMERICANS  OXYGEN COMPOUNDS  GENETIC ASSOCIATION STUDIES  SINGLE-NUCLEOTIDE POLYMORPHISM
Systemic lupus erythematosus (SLE; MIM 152700) is a chronic, often systemic autoimmune disease characterized by the production of high titers of autoantibodies directed against native DNA and other cellular antigens. SLE disproportionately affects women and African Americans (AA), i.e., 0.009% of white men, 0.066% of white women, 0.038% of AA men, and 0.282% of AA women. A genetic contribution to SLE is unequivocal; recent genome-wide association studies in whites and Asians have identified nearly 40 validated susceptibility loci and implicated a broad array of biological pathways. Despite a higher prevalence, incidence, and disease severity, little is known about the genetic etiology of SLE in individuals of African ancestry. Recently, large candidate gene studies have uncovered associations of specific loci in AA.

Overproduction of nitric oxide (NO) and reactive oxygen intermediates is implicated in disease pathogenesis. Markers of systemic NO production and reactive oxygen species (ROS) are higher in patients with SLE than in controls; these markers correlate with disease activity, and early studies suggest that failure to suppress these markers associates with lack of clinical response to therapy for lupus nephritis. These combined observations make NO synthases and other reactive intermediate producing and scavenging genes biological candidates for disease susceptibility.

The Gullah are a unique population of African ancestry in the United States. Their ancestors were forcibly brought from the Sierra Leone and Ivory Coast area in West Africa, and were kept in the geographically isolated Sea Islands along the South Carolina and Georgia coasts. Until recent times, the estimated 100,000 to 300,000 Gullah remained relatively isolated. While continental AA average about 80% West African and 20% European ancestry, the white admixture in the Gullah is <3.5%, and the Gullah are the most homogeneous AA population described. Interestingly, there is the perception that SLE is rare in Africa, suggesting that comparative studies of related cohorts from the 2 continents may provide insight into the genetic etiology of SLE. A higher than predicted prevalence of SLE multiplex families and a high prevalence of seropositivity in SLE first-degree relatives was observed in the Gullah, suggesting a major genetic influence in this population. In parallel, studies report a higher prevalence of certain common complex traits in the Gullah when compared to other AA. Because of their genetic and environmental homogeneity, low European admixture, and increased prevalence and familial clustering of certain diseases, the Gullah are a unique population for deciphering the African heritability in these diseases. The power to detect associations may be higher in more genetically homogeneous populations, such as the Gullah.

Given these advantages of an homogeneous population, we attempted to identify specific genetic variants in genes involved in reactive intermediate production and scavenging predisposing to SLE in the Gullah population and admixed AA.
MATERIALS AND METHODS

Patients and genotyping. The study population consisted of 1565 SLE cases and 1799 controls of African ancestry from the collaborative L Lupus Association Study 2 (LLAS2), including Gullah (133 cases and 112 controls) and non-Gullah AA (1432 cases and 1687 controls). All study participants provided written informed consent that was approved by institutional review boards at each institution. Cases met the 1997 American College of Rheumatology (ACR) criteria for SLE. Race was self-reported. Gullah ancestry was self-identified as AA Gullah from the Sea Islands region of South Carolina and Georgia, with all known grand-parents being of Gullah descent.

Genotyping was performed with the LLAS2 project, which involved multiple investigators and > 32,000 single-nucleotide polymorphisms (SNP). A total of 244 SNP from 53 regions harboring genes selected for their role in producing NO, ROS, or scavenging reactive oxygen and nitrogen species were analyzed (data available from author on request). SNP were chosen for their position and function in each gene, with priority given to those with potentially stronger phenotypic risks (data available from author on request). SNP were genotyped on a customized Illumina Infinium II platform.

Statistical analyses. Only SNP meeting the quality control criteria of < 10% overall missing data, no evidence of differential missingness between cases and controls (p > 0.05), consistency with Hardy-Weinberg equilibrium genotype frequency expectations (p < 0.01 controls, p < 0.0001 cases), and minor allele frequency (MAF) > 5% were included. Related and duplicate individuals were removed and sex inconsistencies and heterozygosity outliers excluded. Potential confounding substructure or admixture was controlled for using 306 ancestry informative markers to compute admixture proportions as implemented in the Admix program; principal component analysis using all SNP confirmed the results. After adjustment for population substructure, the inflation factor using all SNP was \( \lambda = 0.98 \) in the Gullah and \( \lambda = 1.19 \) in the AA. Principal component analysis plots of the AA and Gullah samples were conducted (data available from the author on request). Although the higher inflation factor would be expected given the selection of SNP in candidate genes, a genomic control-adjusted \( p \) value was also computed in AA. In contrast, the \( \lambda \) value close to 1 in the Gullah ensured that false-positive associations due to population stratification were excluded.

The computer program SNPGLA was used for the association analysis (Division of Public Health Sciences, Wake Forest School of Medicine; www.phs.wfubmc.edu/public/bios/gene/home.cfm). The additive genetic model is reported unless the lack-of-fit test for the additive model reached significance (p < 0.05). In that case, the minimum \( p \) value from the additive, recessive, and dominant genetic models is reported. Tests can be affected by low genotype counts; therefore, a minimum of 30 homozygotes and 10 heterozygotes for the minor allele were required to consider the recessive or additive models, respectively. Genetic models were defined relative to the minor allele, and reported results were adjusted for population substructure. In addition to the joint analysis of AA and Gullah samples, a weighted Z score metaanalysis was computed as implemented in METAL software (Center for Statistical Genetics, University of Michigan; www.sph.umich.edu/csg/abecasis/metal), with weights being the square root of the sample size for each dataset; thus, the metaanalysis incorporates direction, magnitude of association, and sample size. To uncover potential haplotype associations, a sliding window haplotype analysis of 3 to 8 SNP was performed in each region. A logistic regression model was employed, adjusting for population structure as implemented in PLINK software (Harvard University; www.mgh.harvard.edu/~purcell/plink).35 Haplotypes with frequency < 10% were excluded.

To test for 2-locus interactions among SNP, all reported results met the quality criteria defined above. Specifically, SAS software (SAS Institute Inc.) was used to compute a logistic regression model, with each SNP modeled under an additive genetic model and the interaction as the centered crossproduct of the SNP under the additive model. To reduce false-positive interactions due to low MAF, we rejected all pairs for which the expected number of individuals in the dataset was < 5 for minor allele homozygotes. In addition, all SNP pairs with a linkage disequilibrium (LD) measure of \( r^2 > 0.2 \) in YRI (a West African ancestry population) were excluded. Interactions were adjusted for population substructure. For chromosome X, only females were included.

A power analysis was computed with Quanto (University of Southern California; hydra.usc.edu/gxe) using a prevalence of 0.1% and \( \alpha = 0.01 \). LD between SNP was assessed with SNAP software using data from the 1000 Genomes Project in YRI (Broad Institute; www.broadinstitute.org/mpg/snap/index.php). SNP functionality was evaluated with the University of California at Santa Cruz (UCSC) genome browser (genome.ucsc.edu).

RESULTS

Genes with known association with production of or regulation of reactive oxygen and nitrogen intermediates were selected (data available from author on request). SNP mapping to 53 regions harboring reactive intermediate related genes were identified (data available from author on request). Association was assessed between these 244 SNP and SLE in the Gullah (133 cases and 112 controls) and AA populations (1432 cases and 1687 controls).

The most significant single-marker associations are shown in Table 1. To minimize potentially spurious associations, effects that were not supported by associations at neighboring SNP were excluded (Figure 1). In admixed AA, the most significant association was identified in the glutathione reductase (GSR) gene. In the Gullah, the most significant associations mapped to the NADH dehydrogenase (ubiquinone) FeS protein 4, 18 kDa (NADH-coenzyme Q reductase; NDUFS4), and NO synthase 1 (NOS1) genes. It should be noted that the variants reported here as associated with SLE met quality control thresholds in both AA and Gullah populations, but revealed association in only one of the populations (Figure 2).

The strongest signal in AA was observed at an intronic variant in a DNAseI hypersensitive cluster in the GSR gene (rs2253409; \( p = 1.43 \times 10^{-03} \), OR 1.26, 95% CI 1.09–1.44). Although modest, other associations with neighboring SNP corroborated the association observed at rs2253409, suggesting it is unlikely to be spurious (Figure 1). Samples had 74% power to detect this effect. Despite meeting quality control thresholds, rs2253409 was not associated in the Gullah (Figure 2).

In the Gullah, the most significant association was an intronic risk variant in the NDUFS4 gene (rs381575; \( p = 6.51 \times 10^{-03} \), OR 2.1, 95% CI 1.23–3.59). Interestingly, this SNP locates in the transcription factor binding site for the RE1-silencing transcription factor and GATA1 and GATA3 proteins. An intronic variant with a protective effect in NOS1 was also identified in the Gullah (rs561712; \( p = 7.18 \times 10^{-03} \), OR 0.62, 95% CI 0.44–0.88). Despite modest power (58% and 45% power to detect the effects reported for NDUFS1 and NOS1, respectively), it is noteworthy that both the NDUFS4 and NOS1 associations were corroborated by associations at neighboring SNP (Figure 1). These variants...
also met quality control thresholds in AA, but were associated only with SLE in the Gullah (Figure 2).

Combining all samples of African ancestry (AA and Gullah) in a joint analysis, the signals identified in the AA predominated, as expected given their larger sample size. Table 1 reveals that in the African population, among the top

Table 1. SNP with the most significant association with SLE.

<table>
<thead>
<tr>
<th>SNP</th>
<th>Chr</th>
<th>Pos (Mb)</th>
<th>Region</th>
<th>Risk</th>
<th>Allele</th>
<th>Case</th>
<th>Control</th>
<th>Best-P*</th>
<th>P-GC</th>
<th>OR (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>African (AA + Gullah)</td>
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<tr>
<td>rs2551715</td>
<td>8</td>
<td>30,666</td>
<td>GSR</td>
<td>A</td>
<td>0.30</td>
<td>0.27</td>
<td>4.57E-03d</td>
<td>6.43E-03</td>
<td>1.21</td>
<td>(1.06–1.38)</td>
</tr>
<tr>
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<td>8</td>
<td>30,667</td>
<td>GSR</td>
<td>G</td>
<td>0.33</td>
<td>0.30</td>
<td>7.21E-04d</td>
<td>1.16E-03</td>
<td>1.26</td>
<td>(1.10–1.44)</td>
</tr>
<tr>
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<tr>
<td>rs2253409</td>
<td>8</td>
<td>30,667</td>
<td>GSR</td>
<td>G</td>
<td>0.33</td>
<td>0.30</td>
<td>1.43E-03d</td>
<td>3.43E-03</td>
<td>1.26</td>
<td>(1.09–1.44)</td>
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<tr>
<td>Gullah</td>
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<tr>
<td>rs381575</td>
<td>5</td>
<td>52,949</td>
<td>NDUFS4</td>
<td>C</td>
<td>0.62</td>
<td>0.55</td>
<td>6.51E-03f</td>
<td>—</td>
<td>2.10</td>
<td>(1.23–3.59)</td>
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<tr>
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<td>NOS1</td>
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<td>0.38</td>
<td>0.51</td>
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<td>—</td>
<td>0.62</td>
<td>(0.44–0.88)</td>
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<td>NOS2A</td>
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<td>0.33</td>
<td>1.74E-03d</td>
<td>—</td>
<td>0.46</td>
<td>(0.28–0.75)</td>
</tr>
</tbody>
</table>

* Best-P, OR, and CI reported under the following genetic models: a additive, d dominant, f recessive. Chr: chromosome; Pos: position; MAF: minor allele frequency; P-GC: best-P after a genomic control adjustment; SNP: single-nucleotide polymorphism; AA: African American; GSR: glutathione reductase; NDUF: nicotinamide adenin dinucleoacid hydrogen; NOS: nitric oxide synthase; SLE: systemic lupus erythematosus.

Figure 1. Regional plots of reactive intermediate gene loci in African, African American (AA), and Gullah populations. Genotyped SNP are plotted with their p-values [as $-\log_{10}(p$ values)] as a function of genomic position (Human Genome Build 18) within a region surrounding the most significant SNP (black diamond). Recombination rates from the 1000 Genomes AFR (11/2010) are plotted in gray to reflect the regional linkage disequilibrium (LD) structure. In each region the index SNP is represented by a large black diamond, and the shade of all other SNP (circles) indicates LD with the index SNP based on pairwise $r^2$ values from the 1000 Genomes AFR study. Known human genes in the University of California at Santa Cruz Genome Browser are at the bottom. GSR: glutathione reductase gene; NOS: nitric oxide synthase.
associations were the aforementioned variant in \textit{GSR} (rs2253409; \( p = 7.21 \times 10^{-04} \), OR 1.26, 95% CI 1.10–1.44) and another intronic variant in \textit{GSR} (rs2551715; \( p = 4.57 \times 10^{-03} \), OR 1.21, 95% CI 1.06–1.38) not in LD with the former (\( r^2 = 0.07 \) in YRI). Very similar results were obtained when a metaanalysis of the AA and Gullah results was computed (rs2253409, \( p = 6.85 \times 10^{-04} \); rs2551715, \( p = 4.49 \times 10^{-03} \)). The samples had 57% and 77% power to detect the effects reported for the first and second variants in \textit{GSR}, respectively. Figure 2 shows how the patterns of association for the reported genes vary among the AA, Gullah, and combined African populations.

Among the cases of SLE, 42% of the 1527 total AA and 46% of the 152 total Gullah cases show renal involvement. Keeping in mind the smaller sample sizes and reduced power to detect associations, testing for association of these SNP with lupus nephritis showed very modest associations: \textit{GSR} in African (rs2253409, \( p = 0.01 \)) and AA (rs2253409, \( p = 0.01 \)) and \textit{NOS1} in African (rs10850803, \( p = 0.01 \)) and AA (rs10850803, \( p = 0.007 \)).

Haplotype-association methods may have more power and accuracy than single markers to detect disease effects. As shown in Table 2, the most significant haplotype association detected in AA and combined African samples was a 3-SNP haplotype in an intronic \textit{NOS1} region (rs3741476, rs1875140, and rs1077490; \( p = 2.87 \times 10^{-04} \), frequency 0.21, OR 1.32 in AA; and \( p = 7.36 \times 10^{-04} \), frequency 0.21, OR 1.28 in all samples with African ancestry). A 4-SNP protective haplotype in an intronic glutathione synthetase (\textit{GSS}) region was also uncovered (rs6087651, rs2236270, rs17092180, and rs2273684; \( p = 2.65 \times 10^{-04} \), frequency 0.14, OR 0.73 in AA; and \( p = 1.71 \times 10^{-04} \), frequency 0.15, OR 0.73 in African). No significant haplotypes were identified in the Gullah population.

Table 3 shows the most significant 2-locus interaction analysis results. Nearly all interactions were specific to one ethnic group or the other. Only an interaction between an \textit{NDUFS2} variant (rs4656993) with a \textit{minichromosome maintenance complex component 5} SNP (rs4645794) was observed in the combined African samples (\( p = 4.00 \times 10^{-04} \), OR 1.34, 95% CI 1.27–1.40) and AA (\( p = 9.74 \times 10^{-05} \), OR 1.40, 95% CI 1.32–1.48).

DISCUSSION

This is the first comprehensive analysis of reactive intermediate genes for their association with SLE in populations of African ancestry. Despite relatively modest power, strict quality criteria filters were applied to reduce the likelihood of false-positive associations. In the Gullah, the virtually perfect inflation factor ensures that false-positive associations due to population stratification can be conclusively excluded. Despite the small sample size of the Gullah being a limitation of our study, all the associations reported were corroborated by associations at SNP in LD with the top
The genes chosen were of interest because they all had the potential to affect reactive intermediate production or clearance/scavenging. ROS can oxidatively modify proteins to influence their activity. Transcription factors AP1 (c-Jun), nuclear factor-κB (NF-κB), hypoxia-inducible factor-1α, and p53 are all redox-regulated. For instance, H₂O₂ generated by endothelial cell (EC) NOX leads to NF-κB-mediated transcription of intercellular adhesion molecule and vascular cell adhesion molecule, both of which associate with lupus nephritis disease activity and atherosclerosis. Two inflammatory cytokines, interleukin 6 (IL-6) and monocyte chemoattractant proteins 1 (MCP-1; CCL2), important to the pathogenesis of lupus nephritis and atherosclerosis, have in common redox-regulated NF-κB response elements (UCSC Genome browser). Both MCP-1 and IL-6 secretion by activated EC is dependent on ROS production and can be inhibited by ROS scavengers.

The functional relevance of the described SNP in reactive intermediate genes is not known. However, reductions in the activity or expression of functionally protective genes could predispose to SLE, SLE disease activity, or target organ damage. SLE is associated with increased markers of oxidative stress, particularly among AA. The consequences of this increased oxidative stress may be increased antigenicity of self-antigens and pathogenic redox signaling. GSR catalyzes the reduction of glutathione disulfide to glutathione, an important antioxidant molecule. Reduced levels of GSH can lead to increased oxidant stress. Of significance to SLE, lower levels of reduced GSH were observed in T cells from patients with SLE, in association with mitochondrial hyperpolarization and ATP depletion, a process that can predispose cells to necrosis. Our study does not address whether the reduced levels of GSH observed in patients with SLE is due to increased production of ROS, reduced enzyme activity, or both.

The proteins encoded by NDUFS2 and NDUFS4 are subunits of the mitochondrial membrane respiratory chain NADH dehydrogenase (complex I). Reports are conflicting; however, defects in NDUFS2 can lead to increased oxidative intermediate production, while described mutations in NDUFS4 do not appear to lead to increased oxidative stress.
Low levels of NO produced by NOS3, also known as endothelial nitric oxide synthase (eNOS), are protective in vascular disease and inflammation. Expression of NOS3 is reduced in proliferative lupus nephritis, and the effects of low-level NO production are functionally reduced in the vasculature of patients with SLE. Similarly, NOS1 appears to prevent leukocyte adhesion in mice lacking NOS3. This finding is also seen in disease, because in both humans and mice with cirrhosis, NOS1 (normally expressed in vascular smooth muscle cells) is upregulated in eNOS (NOS3) deficiency, suggesting that NOS1 can compensate for reduced eNOS activity. Thus, NOS1 deficiency could exacerbate NOS3 dysfunction or deficiency, leading to inflammation and vascular dysfunction. There is evidence for this notion, because NOS1 is associated with endstage renal disease in AA.

We uncovered several novel associations of reactive intermediate-related genes with SLE in patients with African ancestry. We show that many of the loci associated with SLE differ in Gullah and AA, suggesting that specific loci may be more strongly associated in specific populations with African ancestry. This is not a surprising finding given the great genetic diversity present on the African continent. These results suggest that patterns of disease association for SLE may be distinct and specific loci may be more strongly SLE-associated in select African ancestry populations.

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