Genome-wide association analysis of red blood cell traits in African Americans: the COGENT Network

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Laboratory red blood cell (RBC) measurements are clinically important, heritable and differ among ethnic groups. To identify genetic variants that contribute to RBC phenotypes in African Americans (AAs), we conducted a genome-wide association study in up to \sim 16500 AAs. The alpha-globin locus on chromosome 16pter [lead SNP rs13335629 in ITFG3 gene; P < 1E - 13 for hemoglobin (Hgb), RBC count, mean corpuscular volume (MCV), MCH and MCHC] and the G6PD locus on Xq28 [lead SNP rs1050828; P < 1E - 13 for Hgb, hematocrit (Hct), MCV, RBC count and red cell distribution width (RDW)] were each associated with multiple RBC traits. At the alpha-globin region, both the common African 3.7 kb deletion and common single nucleotide polymorphisms (SNPs) appear to contribute independently to RBC phenotypes among AAs. In the 2p21 region, we identified a novel variant of PRKCE distinctly associated with Hct in AAs. In a genome-wide admixture mapping scan, local European ancestry at the 6p22 region containing HFE and LRRC16A was associated with higher Hgb. LRRC16A has been previously associated with the platelet count and mean platelet volume in AAs, but not with Hgb. Finally, we extended to AAs the findings of association of erythrocyte traits with several loci previously reported in Europeans and/or Asians, including CD164 and HBS1L-MYB. In summary, this large-scale genome-wide analysis in AAs has extended the importance of several RBC-associated genetic loci to AAs and identified allelic heterogeneity and pleiotropy at several previously known genetic loci associated with blood cell traits in AAs.

INTRODUCTION

Laboratory red blood cell (RBC) measurements are important for the diagnosis and classification of various hematologic disorders. Some disorders of RBCs, such as sickle cell anemia and alpha thalassemia, are single-gene diseases with higher frequency among populations of African descent (1,2). Even among healthy individuals, African Americans (AAs) have lower hemoglobin (Hgb), hematocrit (Hct) and mean corpuscular volume (MCV) compared with other racial/ethnic groups across all ages (3-5).

Heritability studies suggest that RBC traits are under significant genetic influence. Genome-wide association studies (GWASs) of RBC indices have been reported among European and Japanese populations (6–8), but to our knowledge have not yet been reported for AA. In a gene-centric association study from the CARe consortium, the common African glucose-6-phosphate dehydrogenase (*G6PD*) A-variant on chromosome X and another variant of the α -globin (*HBA2-HBA1*) locus were associated with multiple RBC traits in AAs (9).

The genetic loci reported to date explain only a small fraction of heritability in RBC traits, highlighting the need for larger studies that include ethnic minorities and complementary analytic approaches (10). Thus, we performed a GWA meta-analysis of RBC traits among AA participants from cohorts of the Continental Origins and Genetic Epidemiology Network (COGENT). As AAs are an admixed population, the resulting genomic architecture can be leveraged to identify regions where either African or European ancestral alleles are associated with traits such as Hgb which differ significantly between European and African populations. Therefore, we performed admixture mapping for the association between available RBC traits (Hgb, Hct, MCHC) and local ancestry.

RESULTS

Descriptive analysis

Since not all RBC traits were available in every COGENT cohort, the numbers of individuals available for meta-analysis varied by each RBC trait (Supplementary Material, Table S1). Only Hgb (n = 16485) and Hct (n = 16496) were available in all cohorts. MCHC (n = 12152), MCV (n = 6438), RBC count (4818), MCH (n = 4066) and RDW (n = 3811) were available in subsets of participating cohorts. There were varying degrees of pairwise correlation between RBC traits (Supplementary Material, Table S2). Pearson's correlation coefficients were highest (>0.95) between Hgb and Hct, and between MCV and MCH and were lowest between the RDW and RBC count (0.03).

GWAS of RBC traits in COGENT AAs

The GWA results for each RBC trait are summarized by Manhattan (Fig. 1) and quantile–quantile (Supplementary Material, Fig. S1) plots. The meta-analysis inflation factors were all near unity (0.998–1.005), suggesting that confounders and other technical artifacts were well-controlled. In total, seven independent genomic loci met the experiment-wide significance threshold ($P < 1 \times 10^{-8}$) for one or more RBC traits (Table 1 and Supplementary Material, Table S3). Three loci (1p31.1, 13q31.2, 16p13.3centromeric) have not been previously associated with RBC traits, whereas four loci (2p21,

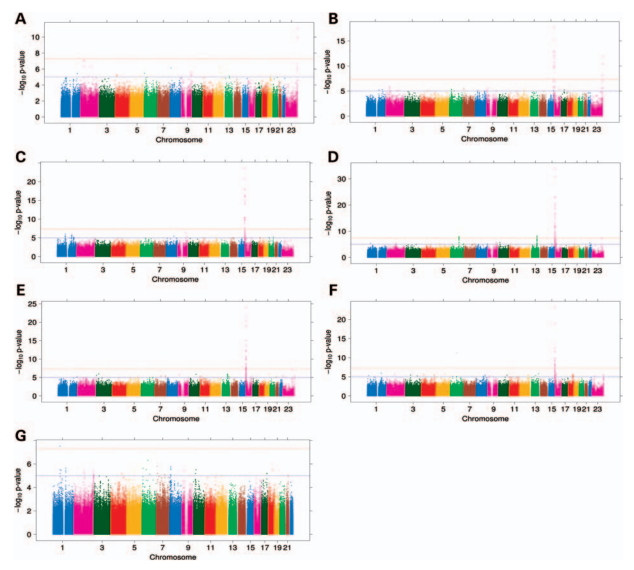


Figure 1. Manhattan plots of GWAS analysis for RBC traits (A) Hct; (B) Hgb; (C) MCHC; (D) MCH; (E) MCV; (F) RBC count and (G) RDW. The dashed horizontal red line indicates $P = 1 \times 10^{-8}$. The dashed horizontal blue line indicates $P = 5 \times 10^{-3}$.

6q21, 16p31.3telomeric, and Xq28) have been associated with at least one such trait in populations of European, Japanese or African descent.

Previously reported RBC loci

The two top Xq28 single nucleotide polymorphisms (SNPs) (rs762516, rs1050828) for Hgb, Hct, MCV and RBC count are located in the *G6PD* gene. rs1050828 encodes the G6PD amino acid substitution Val68Met that results in the *G6PD* A^- allele known to cause *G6PD* deficiency (MIM #305900). The G6PD A-variant has been previously associated with lower Hct, Hgb and RBC count, and with higher MCV in AAs (9). Here, we additionally report that the G6PD A^- allele is associated with the lower RDW. Given the extent of the association signal at Xq28, we repeated the Hgb and Hct association analyses in women from Women Health Initiative (WHI), the largest AA cohort (n = 8304) for Hgb and conditioning the lead SNPs, rs1050828. After adjusting for

rs1050828, the strength of association with Hgb for the remaining SNPs on Xq28 was greatly attenuated and no longer significant (data not shown).

The index SNP on 16p13, which encompasses the α -globin (*HBA2-HBA1*) locus, was rs13335629 within an intron of *ITFG3*. rs13335629 met the genome-wide significance threshold for association with lower Hgb, MCH, MCHC and MCV and also with a higher RBC count. The rs13335629 variant was also nominally associated with lower Hct ($\beta = -0.215 \pm 0.056$; P = 1.33E - 04) and higher RDW ($\beta = 0.0053 \pm 0.0021$; P = 0.01). Lo *et al.* previously reported a common rs1211375 variant within the 16p13 region associated with lower Hgb, MCH and MCV in AAs, and that these associations were not present in Caucasians (9). Our index SNP rs13335629 is in moderate linkage disequilibrium (LD) with rs1211375 ($r^2 = 0.33$ in HapMap YRI).

Three intronic variants of the protein kinase C (PKC)epsilon gene *PRKCE* on 2p21 were associated with lower

Trait	Chromo-some	Number of SNPs with $P < 5 \times 10^{-8}$	Top SNP in the region	Position Hg18	Candidate genes	Minor/ major allele	MAF	Effect size (SE)	P-value
HCT	Xq28	6	rs762516	153 417 857	G6PD, TKTL1, MECP2, MPP1	T/C	0.148	-0.452 (0.055)	2.17E - 16
HCT	2p21	3	rs13008603	46 209 352	PRKCE	A/C	0.163	-0.277(0.047)	4.09E - 09
HGB	16p13.3	11	rs13335629	250 381	ITFG3, LUCL7, NPRL3, POLR3K, RPL2B, MPG	A/G	0.120	-0.190 (0.019)	2.63E - 23
HGB	Xq28	9	rs762516	153 417 857	G6PD, TKTL1, MECP2, MPP1	T/C	0.146	-0.1614 (0.0186)	3.73E - 18
MCHC	16p13.3	38	rs13335629	250 381	ITFG3, LUCL7, NPRL3, POLR3K, RPL2B, MPG, NME4. DECR	A/G	0.117	-0.3298 (0.0227)	8.66E - 48
MCH	16p13.3	28	rs13339636	238 589	ITFG3, LUCL7, NPRL3, POLR3K, RPL2B, MPG, NME4, DECR	G/A	0.132	-0.6847 (0.0559)	1.87E - 34
MCH	13q31.2	4	rs9559892 ^a	88 166 665	_	A/C	0.253	-0.2589(0.0444)	5.46E - 09
MCH	16p13.3	1	rs7192051 ^a	4 482 118	HMOX2	G/T	0.360	-0.2396 (0.0411)	5.70E - 09
MCH	6q21	10	rs9386791	109 715 190	CD164	C/T	0.416	-0.2294(0.0401)	1.20E - 08
MCV	16p13.3	21	rs13335629	250 381	ITFG3, LUCL7, NPRL3, POLR3K, RPL2B, MPG, NME4, DECR, RHOT2, LMF1, WDR90	A/G	0.134	-0.648 (0.0669)	3.61E - 22
MCV	Xq28	6	rs762516	153 417 857	G6PD, FAM3A, F8, MPP1	T/C	0.137	1.5768 (0.2083)	3.76E - 14
MCV	16p13.3	1	rs7192051 ^a	4 482 118	HMOX2	G/T	0.363	-0.259(0.0475)	4.83E - 08
RBC	16p13.3	12	rs13335629	250 381	ITFG3, LUCL7	A/G	0.120	0.1699 (0.0169)	7.48E - 24
RBC	Xq28	9	rs1050828	153 417 411	G6PD, F8, MPP1, MECP2, CTAG2	T/C	0.108	-0.1424 (0.0159)	4.00E - 19
RDW	Xq28	1	rs1050828	153 417 411	G6PD	T/C	0.116	-0.0326 (0.0048)	1.70E - 11
RDW	1p31.1	1	rs10493739 ^a	83 698 745	_	T/C	0.334	0.0128 (0.0023)	3.02E - 08

Table 1. Results of genome-wide significant SNPs for RBC traits in COGENT AA

^aNovel loci.

Hct. The index SNP rs13008603 was also nominally associated with a lower RBC count ($\beta = -0.044 \pm 0.013$; P =4.69E-04), but not with other RBC traits (P > 0.05 for Hgb, MCV, MCH, MCHC, RDW). The three Hct-associated *PRKCE* variants are in strong LD (pairwise $r^2 > 0.7$). Another intronic variant of *PRKCE* (rs10495928) was previously associated with Hgb and Hct in Europeans (8) and with the RBC count in Japanese (6), but showed no evidence of association in AAs (P = 0.50 and 0.71 for Hct and Hgb, respectively). In European and African HapMap populations, there is no evidence of LD between rs10495928 and any of the three Hct-associated variants observed in COGENT AA. These results strongly suggest ethnicity-specific allelic heterogeneity for RBC traits at the *PRKCE* locus.

At 6q21, a haplotype comprised of 10 SNPs (lead SNP = rs9386791) was associated with a lower MCH, and nominally with a lower MCV (P = 1.09E - 05), Hgb (P = 0.007), Hct (P = 0.03), MCHC (P = 0.02), RBC count (P = 0.01). These variants are located ~50 kb upstream of *CD164*, which encodes a mucin-like molecule expressed by human CD34(+) hematopoietic progenitor cells that regulate erythropoiesis. Other variants of the *CD164 5'* flanking region have been associated with RBC, MCH and MCV in Japanese (rs11966072) (6) and with MCV in Europeans (rs9374080) (8). In HapMap CEU, rs9374080 is in LD with our AA index SNP rs9386791 ($r^2 = 0.87$).

Newly discovered RBC loci

Of the three novel loci associated with RBC traits, rs10493739 at 1p31.1 (associated with RDW) and rs9559892 at 13q31.2 (associated with MCH) are both located in regions devoid of known genes. *TTLL7* is the closest gene to rs10493739 (400 kb away) and encodes a tubulin polyglutamylase, which modifies beta-tubulin (11). There are no known genes within 500 kb on either side of rs9559892. The lead SNP at the third locus, rs7192051 is located within the second intron of the heme oxygenase-2 gene (*HMOX2*) and was associated with lower MCH and MCV. Heme oxygenase 2, the protein product of HMOX2, degrades heme and is important in erythropoiesis (12). Although *HMOX2* is located ~4 Mb centromic to the alpha-globin locus, it is not in LD with the previously identified 16p13 association signals (maximum $r^2 =$ 0.004 with rs13335629).

We attempted to validate two of our three novel RBC loci discovered in COGENT in two independent population-based samples: \sim 7700 AA youths ages 8–21 years from CHOP and 2010 AA adults from the Mount Sinai eMERGE study. There was no evidence of replication of rs9559892 with MCH, nor of rs7192051 with MCV or MCH (Supplementary Material, Table S4) in the validation sample. It was not possible to pursue replication of rs10493739 in CHOP and eMERGE because this SNP was not genotyped and it could not be imputed in the available replication samples. In over 20000 Europeans from the CHARGE consortium and 14 000 Japanese from RIKEN, there was no evidence of association of rs9559892 with MCH. Similarly, there was no evidence of association of rs7192051 with MCV or MCH in CHARGE Europeans (Supplementary Material, Table S4).

Admixture mapping analysis of Hgb, Hct and MCHC traits in WHI AA

As a complementary approach to identifying variants associated with RBC traits in AAs that occur at disparate frequencies in ancestral African versus European populations, we performed admixture mapping for Hgb, Hct and MCHC in WHI, the largest cohort comprising COGENT. For MCHC, there was one genome-wide significant association signal at the p-term of 16 containing the alpha-globin locus (Supplementary Material, Fig. S2). Local African ancestry in this region was associated with lower MCHC. The admixture association peak is at rs7203694 (P = 2.78e - 06) located within *RAB40C*, and the genome-wide significant region spans 0-0.78 mb (build 36). There were no genome-wide significant admixture associations for Hct (data not shown). For Hgb, a 2 mb region on chromosome 6p22.2-6p22.1 (25.2-27.1 mb, build 36) reached genome-wide significance, with increased European ancestry associated with higher Hgb levels (Fig. 2A). The Hgb admixture signal appears to be comprised of two peaks (Fig. 2B). Underlying the centromeric peak is HFE, the hemochromatosis protein-coding gene, which regulates iron absorption by modulating the interaction of the transferrin receptor with transferrin. Two known HFE mutations C282Y (rs1800562) and H63D (rs1799945) cause hereditary hemochromatosis, an autosomal recessive iron storage disorder (13). Among individuals of European descent, the frequencies of C282Y and H63D are 3.23 and 16.6%, respectively. Both the mutations are essentially absent in the HapMap YRI populations, and therefore the frequency in AAs is low and is the result of European admixture. C282Y was directly typed in WHI SHARe and other COGENT cohorts (total N = 15584); the genotype association test yielded an Hgb association P =0.0003 in WHI alone and 4.3×10^{-6} in COGENT overall (minor allele frequency = 0.015; $\beta = 0.239 \pm 0.052$). H63D was not directly typed; however, it is tagged by rs129128 $(r^2 = 1.0 \text{ in CEU})$, which was associated with Hgb levels in WHI (P = 0.008) but not when all COGENT cohorts were analyzed together (P = 0.07).

After adjusting for genotypes at C282Y and the H63D proxy rs129128 in WHI, the admixture *P* value for chromosome 6p22 was attenuated, but remained significant (from 3.28×10^{-7} to 1.12×10^{-4}), suggesting the existence of additional variants in this region that contribute to interpopulation differences in Hgb levels. Located within the telomeric peak of the admixture signal (Fig. 2B) are a number of additional variants that have Fst > 0.3, including several near *LRRC16A*, which has been associated with both serum transferrin levels in whites and the platelet count in AAs from COGENT (14). The most strongly associated *LRRC16A* variant rs9356970 is located ~25 kb upstream of the 5' flanking region (MAF = 0.09; $\beta = 0.118 \pm 0.028$; $P = 2.7 \times 10^{-5}$). According to the HapMap, the minor allele is present in 30% of European chromosomes, but only 2.5% of YRI

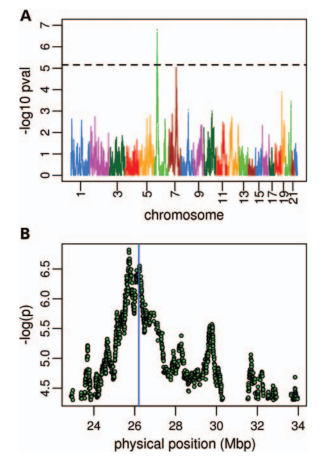


Figure 2. Admixture scan of Hgb concentration. (**A**) shows a genome-wide plot of $-\log(P$ -values) for local-ancestry association with Hgb. The dashed horizontal line indicates the experiment-wide admixture scan significance threshold of $P < 7 \times 10^{-6}$. (**B**) indicates a zoom-in of the genome-wide significant region on chromosome 6, where there appears to be a broad, bimodal admixture peak. The region corresponding to the *HFE* gene is shown in blue.

chromosomes. In a regression model simultaneously adjusting for *LRRC16A* rs9356970 in addition to *HFE* C282Y and rs129128, the association signal for local African ancestry at 6p22 was further attenuated, but remained nominally associated with lower Hgb ($\beta = -0.058 \pm 0.029$; P = 0.045). Together, these results suggest that several European-derived alleles in the 6p22 region, including those of *HFE* and *LRRC16A*, may contribute to higher Hgb levels observed in populations of European descent compared with AAs.

CNV analysis and assessment of allelic heterogeneity at 16p13 alpha-globin region

Given the genetic complexity of the alpha-globin locus on chromosome 16p13, including the presence of a common 3.7 kb alpha-thalassemia deletion in AAs (15), and the extent and magnitude of the observed GWAS signal for RBC traits at 16p13, we assessed structural variation at the 16p13 alpha-globin locus using data from 1000 genomes. First, we confirmed the presence of a common deletion $(-\alpha^{3.7})$ among African Americans (AAs) and West Africans that removes one alpha-globin gene copy (*HBA2*)

(Supplementary Material, Fig. S3). Using pooled sequence data from 16 samples (9 YRI, 5 LWK, 1 ASW, 1 CLM) that appear to be homozygous for $-\alpha^{3.7}$ deletion, we further localized the breakpoints, which appear to be bounded by ~300 bp of nearly identical sequence located within the 5' flanking regions of *HBA1* and *HBA2* (Supplementary Material, Fig. S4). Second, we identified a rare deletion spanning *HBM* through *HBQ1*, in three Han Chinese individuals, and several other possible (but uncertain) rare copy number variations (CNVs) including one duplication (Supplementary Material, Fig. S5) and a very rare deletion that deletes a known regulatory element MCS-R1 (16) (Supplementary Material, Fig. S6).

Among all typed and imputed SNPs in the WHI dataset, the strongest correlation with alpha37 in 63 YRI samples from 1000 Genomes was observed in the region of ITFG3. This includes rs13335629 (r-squared = 0.6), which is also the top Hgb- and MCHC-associated SNP in COGENT AA. We repeated the association analyses in WHI (n = 8304) for Hgb and MCHC conditioning on rs13335629. The top SNP for Hgb in the conditional analysis was POLR3K rs798693 (P = 8.7E - 06). For MCHC, rs2541612 in NPRL3 remained genome-wide significantly associated with lower MCHC (P = 1.14E - 09). When both *ITFG3* rs13335629 and *NPRL3* rs2541612 were included as covariates in the conditional analysis, the SNP most strongly associated with lower MCHC was *LUC7L* rs1211375 (P = 1.50E - 06). Taken together, the 1000 Genomes CNV analyses and the results of conditional regression analyses for Hgb and MCHC suggest that while some of the red cell GWAS association signal may be due to the common African alpha37 deletion, there appears to be independent signals coming from other structural variants and/or SNP(s) in the region.

Cross-ethnic transferability of previously reported RBC to COGENT AA

We assessed whether 72 SNPs previously associated with RBC traits in European or Japanese populations are associated with RBC traits in COGENT AA (Supplementary Material, Table S5). Using the conservative Bonferroni multiple comparison corrected significance threshold (P < 0.0001), we validated four associations. In addition to the association of *HFE* rs1800562 with Hgb, these include *ITFG3* rs1122794 (previously associated with MCH in Europeans) with higher MCHC ($P = 1.5 \times 10^{-8}$), MCH ($P = 7.2 \times 10^{-6}$) and MCV ($P = 7.1 \times 10^{-5}$) in AAs; *ITFG3* rs7189020 (previously associated with MCV in Europeans) with higher MCH ($P = 1.0 \times 10^{-5}$) and MCV ($P = 1.5 \times 10^{-5}$) in AAs; and *HBS1L-MYB* rs7775698 (previously associated with HCT, MCH, MCHC, MCV and RBC count in Japanese) with a lower RBC count ($P = 3.3 \times 10^{-5}$) in AAs.

RBC-associated genetic variants and anemia in AA women

To identify genetic variants associated with anemia, defined dichotomously as Hgb < 12 g/dl, we performed a GWA scan in 8304 AA women 50–79 years old from WHI. Two loci, Xq28 and 16p13, met the threshold of genome-wide significance. The G6PD rs1050828 A-variant was associated with a 1.49-fold (95% CI: 1.33–1.67) increased risk of anemia

 $(P = 3.3 \times 10^{-12})$. The index SNP at 16p13 (rs1088638) is located ~ 20 kb 3' to POLR3K, and was associated with a 1.42-fold (95% CI: 1.26-1.60) increased risk of anemia $(P = 1.2 \times 10^{-8})$. We also constructed a composite RBC genetic risk score (GRS) by summing genotyped or imputed allele dosage at the 15 SNPs associated with at least one RBC trait in AA through the GWA scan, admixture mapping scan, conditional analyses or cross-ethnic transferability analyses described above. The GRS ranged from 5 to 19, with a median of 12. When modeled as a quantitative trait, the GRS was strongly associated with anemia (P = 3.5×10^{-18}), explaining 1.4% of the anemia phenotypic variance, or 2.2% of the variance in Hgb concentration. When WHI participants were grouped into four GRS categories, those in the highest GRS category had a 1.95-fold increased risk of anemia (95% CI: 1.56-2.42) compared with those in the lowest GRS category ($P = 3.6 \times 10^{-9}$).

DISCUSSION

In this first reported GWAS meta-analysis of RBC traits in AAs, we report genome-wide associations for four loci (*G6PD* on Xq28, alpha-globin locus on 16pter, *PRKCE* on 2p21 and *CD164* on 6q21). We also validated the association in AAs of variants in genes such as *HFE* and *HBS1L-MYB*, which have previously been associated with RBC traits in other ethnicities. At the alpha-globin locus, there appears to be allelic heterogeneity (particularly for MCHC), with both copy number variants and SNPs having apparent independent effects. At *PRKCE*, the variants associated with lower Hct in our AA sample appear to be distinct from another set of *PRKCE* variants that have been associated with Hgb, Hct and RBC count in Europeans and Japanese.

Hemizygous males and in some instances female carriers of the X-linked G6PD A- allele are predisposed to acute episodes of drug- or infection-induced hemolytic anemia. Under basal conditions, however, the G6PD A-allele is not generally thought to be associated with RBC abnormalities, and hemizygous G6PD A-individuals have been reported to have normal baseline red cell survival in the absence of oxidant stress (17). Nonetheless, the association of low RDW with G6PD deficiency may be due to low grade hemolysis resulting in an increase in the MCV with rightward shift of the overall distribution of RBC volume without change in the shape of the distribution (18). The G6PD A-variant is in LD with other nearby genetic variants that plausibly could influence Hgb or RBC morphology. TKTL1 encodes a transketolase enzyme that links the pentose phosphate pathway with anaerobic glycolysis, which constitutes the two major metabolic pathways for glucose utilization in human erythrocytes. MPP1 encodes the red cell membrane protein p55, a scaffolding protein that anchors the actin cytoskeleton to the plasma membrane by forming a ternary complex with protein 4.1R and glycophorin C (19).

Aside from genes involved in Hgb synthesis or metabolism, other genetic loci such as *CD164* and *PRKCE* may be associated with RBC traits through effects on erythropoiesis. CD164 (endolyn) is an adhesive receptor present on early hematopoietic progenitors and maturing erythroid cells that

regulates the adhesion of CD34+ cells to bone marrow stroma and affects migration and proliferation of hematopoietic stem cells and progenitor cells (20,21). The upstream region harboring the RBC trait-associated variants contains an erythroleukemia cell line (K562)-specific cluster of histone modifications and ENCODE transcription factor ChIP-seq binding sites including those for GATA-2 and c-Jun. *PRKCE* encodes an isoform of PKC, PKC epsilon, which is expressed in hematopoietic progenitor cells in a lineage- and stage-specific manner and appears to influence erythroid and megakaryocytic progenitor proliferation and differentiation by modulating the response of hematopoietic precursors to a tumor necrosis factor-related apoptosis-inducing ligand (22–24).

Though the finding was not validated in independent AA samples, one of our novel genome-wide significant associations in our discovery cohorts was the association of MCH and MCV with *HMOX2*, which encodes heme oxygenase-2, a constitutively expressed enzyme with a major role in heme catabolism. Heme induces expression of globin genes in erythrocyte progenitor cells and thus plays an important role in erythropoiesis (12,25,26). The lead SNP in this region, rs7192051, is within 5 kb of predicted *HMOX2* regulatory elements such as transcription factor binding sites, DNase sites and histone modification sites (27). Therefore, further study of this variant in larger, independent samples of AA may be warranted.

Our findings have potential clinical implications. Although previous studies have explored the role of common genetic variation in the regulation of these RBC phenotypes in populations of European and Asian descent (6,7,10), no systematic genetic association studies of these traits have been reported in African-ancestry populations. This is particularly important, as there are marked differences in these RBC indices among ethnic groups, and anemia is more prevalent in populations of African descent (28). While it appears that some of the phenotypic variations for RBC and other hematologic traits are controlled by genetic variation shared across ethnic groups (29), other RBC loci are relatively unique to Africans. Rare variants, which are not well captured by GWASs, and undetected common variants of more modest effect may account for additional genetic variance. Discovery and validation of these and additional genetic variants associated with RBC traits in other ethnic populations are likely to uncover new mechanisms and pathways that affect hematopoiesis and RBC turnover, offering insights that may inform further research into red cell biology. Indeed, recent reports have shown that genetic loci uncovered through an unbiased genome-wide study in human populations, together with follow-up functional studies incorporating gene expression, bioinformatic analyses and insights from mouse models and gene knockdown experiments, can greatly contribute to our understanding of the biological mechanisms underlying RBC production (30,31).

MATERIALS AND METHODS

Primary subjects and data collection

We performed GWA analysis of RBC traits in over 16 000 AAs from seven population-based cohorts that comprise the

Continental Origins and Genetic Epidemiology Network (COGENT). The characteristics of each cohort were described in previous publications (14,29). Fasting blood samples were drawn and analyzed for RBC traits at designated clinical laboratories using an automated electronic cell counter. These counters directly measure Hgb concentration (in grams per deciliter), RBC count (in millions per microliter) and MCV, the average size of the RBC in femtoliters. Electronic cell counters calculate MCH, MCHC, Hct and RDW. Hct is the percentage of blood by volume that is occupied by RBC and is calculated by multiplying the RBC count in millions/microliter by the MCV in femtoliters. MCH is the average amount of Hgb inside an RBC expressed in picograms and is calculated by dividing the Hgb concentration by the RBC count in millions per microliter, then multiplying by 10. The MCHC is the average concentration of Hgb in RBCs and is calculated by dividing Hgb in grams per deciliter by Hct. The RDW is a measure of the variance in RBC size and is calculated by dividing the standard deviation of RBC volume by the MCV and multiplying by 100.

All participants self-reported their race/ethnicity. Additional clinical information was collected by self-report and clinical examination. Participants provided written informed consent as approved by local Human Subjects Committees. Study participants who were pregnant or had a diagnosis of cancer or AIDS at the time of blood count were excluded.

Replication subjects and data collection

For validation of novel, genome-wide significant associations identified in the COGENT discovery sample, we performed association analyses in two independent population-based samples: \sim 7700 AA youths ages 8–21 years from Children's Hospital of Philadelphia (CHOP) and 2010 AA adults from the Mount Sinai electronic Medical Records and Genomics (eMERGE) study. We also attempted to replicate novel loci in two other ethnic populations: 14 088 Japanese from RIKEN and up to 30 000 European Americans from CHARGE. Details of each validation cohort are provided under Supplementary Material.

Genotyping and quality-control

Genomic DNA was extracted from peripheral blood leukocytes and genotyping was performed on the Affymetrix 6.0 array or Illumina Omni or 1 M platforms within each cohort using methods described previously (14,29). DNA samples with a genome-wide genotyping success rate of <90% or sex discordance were excluded, as were genetic ancestry outliers (identified by cluster analysis using principal components analysis or multi-dimensional scaling). SNPs with a genotyping success rate of <95% or MAF <1%, monomorphic SNPs and SNPs that map to several genomic locations were removed from the analyses. Participants and SNPs passing basic quality control thresholds were imputed to >2.2 million autosomal SNPs based on HapMap2 haplotype data using a 1:1 mixture of Europeans (CEU) and Africans (YRI) as the reference panel. Details of the genotype imputation procedure have been described previously (14,29). Prior to discovery metaanalyses, SNPs were excluded if imputation quality metrics

(equivalent to the squared correlation between proximal imputed and genotyped SNPs) were <0.30.

Data analyses

For all cohorts, GWA analysis was performed on the raw, untransformed RBC trait using linear regression adjusted for covariates, implemented in either PLINK v1.07 or MACH2QTL v1.08. In GeneSTAR, the family structure was accounted for in the association tests using linear mixed-effects models implemented in R (32). For the 22 autosomes, analysis was performed using genotyped and imputed SNPs. For the X chromosome, only genotyped SNPs were analyzed due to the technical limitations of imputing X-linked SNPs. All analyses were performed under an additive genetic model using allelic dosage (genotyped or imputed) at each SNP, adjusted for age, age-squared, sex and clinic site (if applicable), 4-10 principal components.

For each phenotype, meta-analysis was conducted using inverse-variance weighted fixed-effects models to combine β coefficients and standard errors from study-level regression results for each SNP, to derive pooled estimates. Study-level results were corrected for genomic inflation factors (λ) by multiplying the standard errors (SEs) of the regression coefficients by the square-root of the study-specific λ . Meta-analyses were implemented in the METAL software. Between-study heterogeneity of results was assessed by using Cochran's Q statistic and the I^2 inconsistency metric. A threshold of $\alpha = 1 \times 10^{-8}$ was used to declare genomewide statistical significance. This statistical threshold accounts for the greater nucleotide diversity and lower LD in African descent populations combined with testing of multiple, correlated RBC traits (31,33). We carried out replication testing of 'suggestive' SNPs selected on the basis of a more liberal significance threshold in our primary AA discovery GWAS $(P < 5 \times 10^{-8}).$

To assess the potential existence of multiple, independent variants influencing a trait at the same locus (allelic heterogeneity), regression analyses were repeated in the largest sample (WHI, n = 8095), conditional on the most strongly associated (index) SNP in that region.

We also assessed the transferability to AAs of SNPs previously associated with RBC traits in populations of European or Japanese ancestry by assessing association with RBC traits in the COGENT discovery meta-analyses. For validation, we considered consistency of direction of effect, and assessed statistical significance using a simple Bonferroni adjustment for the total number of SNPs assessed, using a two-sided hypothesis test.

Local ancestry estimation and admixture mapping in WHI

For each AA individual in the WHI sample, locus-specific ancestry was estimated using an extension of the model described by Tang *et al.* (34). We used phased haplotype data from HapMap3 CEU and YRI individuals as reference panels. An admixture mapping analysis was performed in WHI to test for association between Hgb levels and ancestry at each genomic location (local ancestry), while adjusting for the first 10 principal components, regions of recruitment,

clinical trial, age and age-squared. The critical value for genome-wide significance level of admixture mapping is substantially lower than the genotype test due to the extensive correlation in local ancestry between adjacent markers that result from the recent admixture in AAs. We therefore adopted an empirically determined genome-wide significance threshold of $P < 7.1 \times 10^{-6}$, which corresponds to a Bonferroni correction of ~7000 independent tests (35).

Copy number variation (CNV) analysis using 1000 genomes data

We used the 1000 Genomes sequencing data to investigate CNVs at the chromosome 16 p31 alpha-globin locus, studying 946 African-ancestry samples at roughly $4 \times$ sequencing coverage. As a result of noise in depth-based genotyping at this locus (due to low-pass sequencing, high %GC and potential overlapping variants), some of our analyses were confined to the 76 YRI samples, which have higher sequence coverage in 1000 Genomes data and more complete genotyping (call rate 84 % at 95 % CI).

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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