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Analysis of MRI-derived spleen iron in the UK Biobank identifies genetic variation linked to iron homeostasis and hemolysis

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Summary

The spleen plays a key role in iron homeostasis. It is the largest filter of the blood and performs iron reuptake from old or damaged erythrocytes. Despite this role, spleen iron concentration has not been measured in a large, population-based cohort. In this study, we quantify spleen iron in 41,764 participants of the UK Biobank by using magnetic resonance imaging and provide a reference range for spleen iron in an unselected population. Through genome-wide association study, we identify associations between spleen iron and regulatory variation at two hereditary spherocytosis genes, ANKI and SPTA1. Spherocytosis-causing coding mutations in these genes are associated with lower reticulocyte volume and increased reticulocyte percentage, while these common alleles are associated with increased expression of ANKI and SPTA1 in blood and with larger reticulocyte volume and reduced reticulocyte percentage. As genetic modifiers, these common alleles may explain mild spherocytosis phenotypes that have been observed clinically. Our genetic study also identifies a signal that co-localizes with a splicing quantitative trait locus for MS4A7, and we show this gene is abundantly expressed in the spleen and in macrophages. The combination of deep learning and efficient image processing enables non-invasive measurement of spleen iron and, in turn, characterization of genetic factors related to the lytic phase of the erythrocyte life cycle and iron reuptake in the spleen.

Introduction

In normal human physiology, iron is recycled much faster than new dietary iron is absorbed, and this iron economy is regulated by the spleen.1–3 The spleen plays a critical role in removing senescent erythrocytes from the blood and does so via a population of splenic macrophages as well as by the action of the protein ferroportin, which transports iron back to the plasma.4 Given the large iron flux through the normal spleen due to erythrocyte recycling, measurement of spleen iron has the potential to reflect activity of iron salvage pathways.

Spleen iron also may reflect erythrocyte biology and dysfunction. Hereditary spherocytosis (HS [MIM: 270970 and 182900]) is a relatively common hemolytic anemia, occurring at a prevalence of 1:1,000–2,500 in European populations.5 Affected individuals have defects in membrane and cytoskeletal genes that contribute to erythrocyte membrane integrity and deformability, including SPTA1 (encoding the filamentous protein alpha-spectrin [MIM: 182860]) and ANKI (encoding a protein that tethers spectrin filaments to erythrocyte membranes [MIM: 612641]).5,7 The protein products of these genes interact in the formation of the mature erythrocyte cytoskeleton, a process accompanying cellular ferroportin during reticulocytosis.8 In HS, the resulting erythrocytes are spherically shaped and lose deformability and are ultimately trapped and ingested by macrophages within the red pulp of the spleen, resulting in an enlarged spleen and anemia despite brisk reticulocytosis. When measured directly with radiolabeled cells, erythrocyte turnover is dramatically accelerated in individuals with severe HS.9 While laboratory measures such as sphered cell and reticulocyte volume can diagnose HS,10 less is understood about the phenotypic and genetic heterogeneity of the disease.11–13

Although spleen iron has been investigated in specific disease groups,14–20 few studies were performed in unascertained cohorts, partly because of limitations in detecting and quantifying low levels of tissue iron in non-overloaded populations.21–23 The UK Biobank (UKBB) is a prospective study of half a million adults in the UK24 that has genetic and phenotypic information including magnetic resonance imaging (MRI).25 In this study, we applied computer vision techniques to quantify spleen iron non-invasively and at scale by repurposing the dedicated liver MRI acquisition. Spleen iron is only moderately correlated with other measures of iron stores in the body. Through genome-wide association study (GWAS), we characterized associations between spleen iron and genes involved in the human iron economy including common regulatory variation in the ferroportin gene, SLC40A1. Our GWAS identified a signal that co-localizes with a splicing quantitative trait locus for MS4A7, which we found to be abundantly expressed in the spleen and macrophages. Regulatory variation in the genes encoding alpha-spectrin (SPTA1) and ankyrin (ANKI) were also linked to spleen iron and...
increased mRNA expression of these genes and further-
more exhibited effects on reticulocyte and erythrocyte
parameters opposite to the effects observed in HS-affected
individuals. Non-invasive imaging coupled with genetic
analysis enabled us to develop a quantitative trait that illu-
minated aspects of iron homeostasis relevant to hemolysis
and iron recycling in the spleen.

Subjects and methods

Image analysis
We trained a convolutional neural network based on the U-net ar-
tecture26 for 3D organ segmentation from the neck-to-knee
Dixon MRI data by using 119 manual spleen annotations. The dice
score on an out-of-sample held-out test set was 0.922. For the
quantitative MRI data, we estimated the proton density fat
fraction (PDFF) and R2* from the single-slice liver multi-echo
data by using the PRESCO (phase regularized estimation using
smoothing and constrained optimization) algorithm.27 We con-
verted R2* into iron concentration (mg/g) by using the widely
used formula proposed by Wood et al.: iron concentration
= 0.202 + 0.0254 R2*.28,29 We extracted 2D masks from the 3D
spleen segmentations at their intersection with the liver acquisi-
tion (Figure 1A).30 We applied one-pixel erosion before computing
the median value within that mask. We excluded 2D masks that
had <1% of 3D volume or <20 voxels. Additional methods,
including image acquisition details, are described in the supple-
mental information.

Epidemiological modeling of spleen iron risk factors
Associations of spleen iron with age, genetic sex, and self-reported
ethnicity were performed in R v3.6.3. with the linear and logistic
models for spleen iron as a quantitative or binary trait, respecti-
ately, after adjusting for covariates including imaging center,
date, and time.

Phenome-wide association study
We generated a list of variables derived from raw data by using
PHESANT13 and removed procedural metrics (e.g., measurement
date), duplicates, and raw measures, resulting in 1,824 traits
(Table S2). We used PheWAS32 to combine ICD10 codes (Field
41270) into distinct phenotype codes or phecodes (Table S3). In
addition, we included 11 quantitative traits defined in our previ-
ous study.33 We performed linear (quantitative traits) or logistic
regression (binary traits) on spleen iron, adjusting for imaging cen-
ter, date, scan time, age, sex, BMI, height, and ethnicity.

GWAS
We performed a GWAS in n = 35,324 participants as described24
by using UKBB-imputed genotypes24 version 3 for our GWAS,
excluding single-nucleotide polymorphisms (SNPs) with minor
allele frequency < 1% or imputation quality (NFO score)
< 0.9. We included participants identified as White British by using
UK Biobank Field 22006, which is based on self-reported
ancestry and genetic ancestry based on principal-component
analysis.24 We excluded participants exhibiting sex chromosome
aneuploidy, participants with a discrepancy between genetic and
self-reported sex, heterozygosity and missingness outliers, and
genotype call rate outliers.24 9,911,384 SNPs passed quality
control (QC). We used BOLT-LMM35 v2.3.2 to conduct the genetic
association study. We included age at imaging, age2, sex, imaging
center, scan date and time, and genotyping batch as fixed-effect
covariates and genetic relatedness derived from genotyped SNPs
as a random effect to control for population structure and relat-
edness. We normalized the outcome variable by using inverse-
rank normalization. In the genetic association study, we found
no evidence for global inflation of test statistics (λgc = 1.035;
linkage disequilibrium [LD] score regression intercept 1.027 [SE
0.0072]).

Replication analysis
We replicated blood trait findings by using the non-European
meta-analyses from the Blood Cell Consortium Phase 2, which
investigated the genetic basis of 15 blood cell traits in cohorts of
diverse ancestries, including African ancestry (n = 15,171), East
Asian (n = 151,807), Hispanic/Latino (n = 9,368), and South
Asian participants (n = 8,180) in addition to European ancestry
(n = 563,946).36,37 To avoid overlap with UK Biobank Euro-
peans, we focused on summary statistics from the non-European
cohorts. We downloaded summary statistics for mean corpuscular
hemoglobin concentration (MCHC), mean corpuscular vol-
ume (MCV), and monocyte count from http://www.mhi-human
_genetics.org/en/resources/ on November 14, 2021, and harmo-
nized them by using dbSNP build 151 (GRCh37). We tested for
replication of MCHC and MCV at rs4737010[A] (ANK1) and
rs2479868[T] (SPTA1) and replication of monocyte count at
rs950802[A] (MS4A7).

Conditional analysis and fine-mapping
We performed conditional analysis by using GCTA,38 considering
variants within 500 kb of an index variant. We constructed a refer-
ence LD panel of 5,000 randomly selected, unrelated European
UKBB participants.24 We excluded the major histocompatibility
complex region because of the complexity of LD structure at this
locus (GRCh37:6: 28,477,797–33,448,354; see https://www.ncbi.
nlm.nih.gov/ucsc/human/regions/MHC). For each locus, we con-
sidered variants with locus-wide evidence of association
(pjoint < 10−6) to be conditionally independent. We followed an
iterative procedure to determine credible sets of causal variants
with 95% coverage.34

Colocalization studies
For gene expression studies, we used summary statistics from
GTex v8.39 For disease and quantitative trait studies, we used
UKBB summary statistics of phecodes,40 normalized quantita-
tive traits (http://www.nealelab.is/blog/2017/7/19/rapid-gwas-of-
thousands-of-phenotypes-for-337000-samples-in-the-uk-biobank).
We selected phenotypes with p < 5 × 10−8 within 500 kb of the in-
dex variant. We performed colocalization analysis by using coloc41
with default priors and considered variants within 500 kb of the in-
dex variant. We considered two genetic signals to have strong evi-
dence of colocalization if PP3 + PP4 ≥ 0.99 and PP4/PP3 ≥ 5.42

Heritability estimates
We estimated the heritability of each trait by using the restricted
maximum likelihood method,33 as implemented in BOLT-LMM.

Genetic correlation
We computed genetic correlation by using bivariate LD score
regression (LDSC).44 We computed the genetic correlation be-
tween spleen iron and 288 complex traits with a heritability of

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at least 5% from the Neale Lab [http://www.nealelab.is/uk-biobank/], plus organ iron measurements and volumes, and blood iron biomarkers. Following the recommendations of the developers, we (1) removed variants with imputation quality (INFO) < 0.9 because the INFO value is correlated with the LD score and could introduce bias, (2) excluded the major histocompatibility complex (MHC) region because of the complexity of LD structure at this locus (GRCh37:6: 28,477,797–33,448,354; see https://www.ncbi.nlm.nih.gov/grc/human/regions/MHC), and (3) restricted to HapMap3 SNPs. We used a Bonferroni-corrected p value of $1.7 \times 10^{-4}$ as the significance threshold to identify traits with a significant genetic correlation.

Exome sequence quality control

We performed quality control of $n = 200,643$ whole exomes from the UKBB. Raw genotype calls were filtered genotype-level quality metrics to identify quality outliers for a given site and remove poor-quality individual-level genotypes. Similarly to the functionally equivalent (FE) pipeline, we removed genotypes below a minimum read depth (for SNPs: 7 and for indels: 10) and genotypes below a minimum Phred-scaled genotype quality of 20. We removed genotypes where minor allele allelic balance < 0.15 for SNPs and 0.2 for indels. Supplementing FE filters with additional filters, we performed per-SNP QC, requiring the average genotype quality to be at least 30 and per-SNP depth of coverage to be at least 15, to filter out badly captured sites. Additionally, we removed variants with genotype missingness > 10% or that deviated meaningfully from Hardy-Weinberg equilibrium in a European ancestry cohort (HWE p < 1e–10). Of 17,981,897 total variants, 13,907,865 variants passed QC in the European exome cohort with MRI data ($n = 18,240$).

Exome variant annotation

We performed annotation by using VEPv100, LOFTEE, CADD, and ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/) with a custom pipeline to select variants meeting high-confidence loss-of-function criteria, filtered for rare variants (defined as cohort-specific minor allele frequency < 0.001). Of variants passing quality control, we subset 286,456 high-confidence, rare loss-of-function variants, 2,919,962 rare missense variants (CADD score $R > 20$), and 13,705 rare clinical pathogenic variants in 19,992 European ancestry samples for further analysis.

Rare variant association study

We performed rare variant burden and SKAT testing in SAIGE-GENE by using a mixed-effects model. A kinship matrix was built in SAIGE from a filtered set of 354,878 genotyped variants ($r^2 < 0.2$, minor allele frequency > 0.05, Hardy-Weinberg p value > 1e–10, excluding known regions of long-range LD). The linear mixed model regression equation was as follows:

$$y_i = \alpha + Xb + G\beta + b_i + \epsilon_i$$

In the model, $y_i$ is inverse-rank normalized spleen iron, and $X$ represents fixed-effect covariates: age at imaging visit, age$^2$, chromosomally determined sex expressed as a binary indicator variable, study center, standardized scan date, standardized scan time, and the first five principal components of European genetic ancestry. $G_i$ represents allele counts (0, 1, 2) for $q$ variants in each gene to test. We then performed SKAT and burden tests in SAIGE-GENE and reported the p value from SKAT-O. To inform directionality of effect, we reported the betas from the burden test. To avoid unstable results at low sample size, we calculated cumulative minor allele count and thresholded at $\geq 5$ minor alleles per gene, including singletons and doubletons.

Genetic identification of hereditary spherocytosis alleles

From the exome cohort filtered for European ancestry ($n = 167,246$), we annotated variants by clinical assertion as pathogenic according to ClinVar, downloaded from [https://ftp.ncbi.nlm.nih.gov/pub/clinvar/](https://ftp.ncbi.nlm.nih.gov/pub/clinvar/) on September 27, 2020, and called...
Characterization of spleen iron in a large, population-based cohort

We quantified spleen iron concentration (spleen iron hereafter) in 41,764 UKBB participants with both the 3D neck-to-knee and the quantitative liver single-slice MRI sequences available. We opportunistically measured spleen iron concentration by segmenting the spleen from the neck-to-knee image33 (Figure 1A) and subsequently extracting a 2D mask where the spleen volume intersects with the quantitative 2D liver slice (Figure 1B).30 The average spleen iron was 0.92 ± 0.32 mg/g, significantly lower than liver iron of 1.24 ± 0.29 mg/g (Table 1, Figure 1C; paired t test p < 2.2e−16). While there is no accepted normal range of spleen iron, 1.98 mg/g has been suggested as an upper cut-off, and 2.74 mg/g is reported to be pathological.51 Using 1.98 mg/g as the threshold, 1.04% (n = 435) of this cohort had elevated spleen iron, while 0.32% (n = 137) had spleen iron above the 2.74 mg/g threshold. 95% of the population fell into the range of 0.54 to 1.69 mg/g, and we propose this as a possible reference range in an unselected population (Table S2).

Spleen iron differed by age and sex. Men had higher spleen iron than women (men: 0.96 ± 0.34 mg/g; women: 0.87 ± 0.29 mg/g) (p = 6.3e−219; Tables S2 and S3). Increasing spleen iron was associated with age (0.0044 mg/g/year or 0.012 SD/year) (Figure 1D). In women, menopause was associated with 0.12 mg/g higher spleen iron [95% CI 0.08–0.16].

In a phenome-wide association study with over 3,200 quantitative traits and disease outcomes, spleen iron was correlated with erythrocyte parameters: reticulocyte percentage (beta = 0.091; p = 3.7e−66), reticulocyte count (beta = 0.087; p = 6.6e−64), and high light scatter reticulocyte count (beta = 0.089; p = 5.8e−64). Spleen iron was also associated with lifestyle factors, including consumption of lamb (beta = 0.150; p = 2.1e−44) and beef (beta = 0.143; p = 1.3e−43), and negatively associated with alcohol consumption (beta = −0.087; p = 1.7e−18). Spleen iron correlated with liver iron (beta = 0.409, p < 1e−300), brain iron content, specifically with T2* (inversely proportional to iron) in the caudate (beta = −0.063; p = 4.3e−16) and putamen (beta = −0.061, p = 4.1e−15).52 Spleen iron was associated with myeloid leukemia (beta = 0.386; p = 9.6e−10), chronic dermatitis (beta = 0.328; p = 4.1e−07), hypokalemia (beta = 0.286, p = 5.8e−6), and glaucoma (beta = 0.186, p = 1.0e−6) (Figure 2, Tables S4 and S5). Spleen iron was negatively correlated with iron-deficiency anemia, but this did not achieve Bonferroni significance (beta = −0.134, p = 0.002), although this diagnosis may not have been fully captured by medical billing codes (n = 580 affected individuals, n = 35,316 control individuals).

GWAS of spleen iron identifies DNA polymorphisms linked to global iron homeostasis

In a common variant GWAS of spleen iron, seven loci reached genome-wide significance (p < 5e−8; Figure 3, Table 2). Conditional analysis yielded no secondary signals. We estimated the narrow-sense heritability of spleen iron to be 16.7% (SE 1.64%). Spleen iron was moderately genetically correlated with ferritin (rF = 0.56) and MCHC (rF = 0.42), but the genetic correlations with other iron measures, including liver iron, were not significant (Tables S6 and S7).
We observed a signal on chromosome 2 at SLC40A1, which encodes the iron transporter ferroportin (lead variant: rs13008848[G], beta $\approx$0.057, $p \approx$2.9e$^{-10}$, Figure 4A). To test whether the spleen signal was shared across other body iron traits, we re-analyzed previous genetic studies of serum ferritin, serum iron, and liver iron$^{33,45}$ and observed evidence of co-localization at SLC40A1 (posterior probability $R \approx$0.99) (Table S8). Since the lead variant lies upstream of the SLC40A1 open reading frame, we tested for a shared effect between spleen iron and SLC40A1 expression. Across 54 human tissues of the GTEx Consortium,$^{53}$ we observed evidence for regional co-localization with a quantitative trait locus for SLC40A1 mRNA in many tissues, including whole blood (posterior probability $\geq 0.99$) (Figure 4D; Table S9). As expected, spleen iron was associated with SLC40A1, and furthermore, we showed that this locus most likely influences tissue iron levels through SLC40A1 mRNA abundance.

The GWAS of spleen iron identified other loci relevant to iron homeostasis, including a locus on chromosome 9 (Figure S5B). The lead SNP rs41276777[A] (beta $= 0.17$, $p = 2.1e^{-9}$) occurred in the 5’ untranslated region of PRPF4 and CDC26 (Figure S5A) and was associated with a regulatory locus affecting expression of both PRPF4 and CDC26 mRNA in whole blood and the spleen (Table S9), suggesting bi-directional regulation of gene expression. This region colocalized with signals for serum ferritin, serum iron, and other blood iron traits (Table S8),

Figure 2. Phenome-wide association between spleen iron and complex traits in the UK Biobank

(A–F) Traits are organized by category: (A) biological samples, (B) lifestyle and history, (C) physical measures, (D) online follow up, (E) medical history, and (F) disease diagnosis. Bonferroni significance threshold is shown by horizontal dashed line. The top three associations in each category are annotated. ML, myeloid leukemia; HLS reticulocyte, high light scatter reticulocyte.
suggested roles not only in spleen iron but body iron more broadly.

Elevated spleen iron colocalizes with a splicing quantitative trait locus for MS4A7 and monocyte traits

Our GWAS with spleen iron found a significant association at the MS4A7 locus (lead variant: rs950802[A], beta = 0.086, p = 7.7e−26; Table 2, Figure 5A). rs950802[A] causes a synonymous mutation at Leu57 in the third exon of MS4A7 and is also a variant in the first intron of MS4A14. This signal colocalized with a signal of serum ferritin (posterior probability ≥ 0.99). Using splicing quantitative trait data, we identified an alternative splicing event in MS4A7, which colocalized with spleen iron (Figure 5B). Based on an analysis of the open reading frame of MS4A7, this alternative splicing event was predicted to increase skipping of the second exon and therefore interrupt a conserved CD20-like transmembrane domain in MS4A7 (Figure 5C). We thus identified an association between MS4A7 and spleen iron and found a plausible molecular mechanism by which this variant disrupts MS4A7 function.

To explore the functional consequences of regulatory variation in MS4A7, we examined its association with hematological parameters in the UKBB and observed a significant association with monocyte count and percentage as well as platelet count and crit (Figure 5D). Using meta-analysis in non-European populations from the Blood Cell Consortium, a large study of blood cell traits in diverse cohorts, we replicated associations and directions of effect between this locus and monocyte count in an East Asian cohort (n = 151,807) at Bonferroni significance (p < 0.001) and observed the same direction of effect in African ancestry (n = 15,171) and Hispanic (n = 9,367) cohorts (Table S11). Further, we found enrichment of MS4A7 mRNA in monocytes compared to sixteen other hematopoietic cell types (Figure 5E) and also higher expression of MS4A7 mRNA in the spleen relative to other tissues (Figure 5F). A common allele at MS4A7 was thus associated with elevated spleen iron and monocyte traits. MS4A7 mRNA was enriched in the spleen and in monocytes, suggesting a role for this gene in iron recycling in the spleen.

GWAS of spleen iron identifies common alleles in SPTA1 and ANK1 linked to increased gene expression and erythrocyte function

In addition to signals linked to iron homeostasis, the GWAS of spleen iron revealed associations in SPTA1 and ANK1, encoding structural components of erythrocytes (Table 2, Figure S6). The lead variant rs2479868[A] (beta = −0.083, p = 2.5e−22) on chromosome 1 was located in the 3’ untranslated region of SPTA1, and the lead SNP rs4737010[A] (beta = 0.077, p = 1.0e−17) on chromosome 8 was the first intron of ANK1 (Figures 4B and 4C). We tested for a shared signal with mRNA expression levels of SPTA1 and ANK1 and observed co-localization with cis-regulatory variation for ANK1 in multiple tissues (Figures 4E and 4F).

Variation at each locus was associated with increased mRNA expression and mean reticulocyte volume and decreased MCHC, reticulocyte percentage, and spleen volume (Figure 6). The magnitude and directionality for the changes in red cell parameters associated with the lead SNPs at SPTA1 and ANK1 were similar, but a regression model testing for interaction between these two loci found that the effects are independent (beta = 0.016, p = 0.28). These signals did not co-localize with iron traits in other tissues such as serum or liver, suggesting specific effects to the spleen (Table S7).

Using non-European meta-analyses from the Blood Cell Consortium as independent cohorts, we replicated the associations and directions of effect between SPTA1 and ANK1 lead variants and both MCHC and mean corpuscular volume in an East Asian cohort at Bonferroni significance (p < 0.0042). In a cohort of African ancestry and in a cohort of Hispanic ancestry, we replicated directions of effect (Table S11).

Common alleles in SPTA1 and ANK1 show effects on erythrocytes opposite to effects observed with rare deleterious alleles and in hereditary spherocytosis

Given the known associations of both SPTA1 and ANK1 gene defects with hereditary spherocytosis (HS), we analyzed rare variation (minor allele frequency < 0.001) predicted to cause loss of function. Starting with 167,246 exomes of European ancestry, we conducted rare variant association studies for
In this study, we combined deep-learning algorithms and efficient image processing to quantify spleen iron in 41,764 participants of the UKBB. This enabled us to estimate a reference range for spleen iron in a large, unselected population. Spleen iron was higher in men, and our analysis of pre- and post-menopausal women is consistent with prior reports of iron stores in other tissues. Spleen iron increased with age, is associated with red meat intake, and is inversely associated with alcohol consumption, extending prior observations in the liver. Further, spleen iron was only modestly associated with measures of iron in other tissues. Unlike other iron-rich organs such as the liver, we discovered that spleen iron was associated with indicators of reticulocytosis and reticulocyte turnover.

Our GWAS of spleen iron identifies regulatory loci in SPTA1 and ANK1, which when combined with other evidence, suggests a model of low splenic turnover due to relatively large, long-lived erythrocytes. These alleles are associated with decreased spleen iron and increased mRNA expression of their respective cis gene, as well as larger reticulocyte volume and reduced measures of reticulocytosis (Figure 6). Since the spleen is the major route of erythrocyte clearance and iron salvage, lower levels of reticulocytosis would be expected to result in lower spleen iron at steady state. Neither variant is associated with anemia, suggesting that they are not pathogenic. This reduced turnover model also predicts that iron levels in other organs not involved in erythrocyte clearance (such as the pancreas) would be unaffected by these two loci, consistent with the data.

As genetic modifiers, these alleles in SPTA1 and ANK1, which segregate frequently across global populations, may explain the variable penetrance and expressivity observed in HS. In a recent study of affected individuals with identified HS mutations, 64% involved SPTA1 or ANK1, and the investigators observed multiple HS families with broad phenotypic variability, including a compelling example of dizygotic twins sharing the same pathogenic ANK1 mutation presenting with mild disease in one affected individual and severe disease requiring splenectomy in the other. The variation could not be explained, leading to speculation that yet-unknown genetic factors may be contributing. We were able to identify the hallmarks of HS through rare loss-of-function variation in SPTA1, even within a cohort not ascertained for hemolytic anemias. Our analysis suggested that common genetic variants may modify the effects of rare deleterious alleles. The expression-increasing variants identified here may help to explain the heterogeneity that has been observed in HS patients.

In addition to genes specific to red cell turnover in the spleen, our genetic study of spleen iron also pointed to regulators of the human body’s iron economy. We characterized a spleen iron signal at M54A7, which belongs to the CD20

### Table 2. Fine-mapped lead SNPs from GWAS of spleen iron

| Lead SNP | Locus | Lead SNP consequence | Effect allele | Other allele | Beta | p | Minor allele frequency |
|----------|-------|----------------------|--------------|-------------|------|---|-----------------------|
| rs950802 | M54A7/M54A14 | synonymous variant | A | G | 0.086 | 7.7e-26 | 0.307 |
| rs2479868 | SPTA1 | intron variant | T | C | -0.083 | 2.5e-22 | 0.266 |
| rs4737010 | ANK1 | intron variant | A | G | -0.077 | 1.0e-17 | 0.228 |
| rs13008848 | SLC40A1 | 5’ UTR variant | G | C | -0.057 | 2.9e-10 | 0.226 |
| rs41276777 | PRPF4 | 5’ UTR variant | A | G | 0.175 | 2.1e-09 | 0.019 |
| rs115697725 | KLHL29 | intron variant | G | C | 0.059 | 4.9e-08 | 0.146 |

Genome-wide significant associations (p < 5e-8) are shown by locus and lead SNP after fine-mapping. A seventh association at the major histocompatibility locus (MHC) could not be fine-mapped.
family of membrane proteins, which are expressed within the hematopoietic lineage and largely uncharacterized.62,63 Here, we linked a splicing quantitative trait locus in MS4A7 to spleen iron and macrophage abundance and even found that this gene was enriched in the spleen and in macrophages. It is possible that excess circulating monocytes can be recruited to provide an expanded reservoir in the spleen, contributing to the body’s iron economy in addition to splenic macrophages.9,64 Our findings regarding MS4A7 potentially illuminate additional details of splenic erythrocyte clearance mechanisms. For example, the specific functional roles played by macrophages and monocytes in the splenic red pulp remain incompletely understood.9,64

This study has limitations. First, as the only study to quantify spleen iron in a large cohort, no replication cohort is available, though we are able to replicate all our findings on relevant blood cell traits in independent cohorts.66,67 Second, while we found limited evidence in the UK Biobank that spleen iron varies by ethnicity (Figures S3 and S4), additional imaging studies are warranted to quantify spleen iron across populations. Third, experimental studies in the hematopoietic lineage of a model system will be needed to test the functional consequences of the regulatory variation observed here. Fourth, while we genetically identified HS in the UKBB and showed that the common alleles in SPTA1 and ANK1 act as genetic modifiers of erythrocyte parameters, additional clinical validation is needed to substantiate these findings in an HS cohort.

In summary, we have quantified spleen iron by repurposing liver MRI acquisitions, maximizing use of the data at no extra cost to scanning or participant time. Our findings suggest that steady-state levels of spleen iron are sensitive to alterations in erythrocyte structure affecting cell turnover as well as alterations in iron transport by macrophages. We identified common regulatory variation in HS genes showing effects on erythrocytes that are opposite of the effects observed in HS-affected individuals. Imaging-derived spleen iron is thus a quantitative biomarker reflecting hemolysis and iron reuptake by the spleen, is tractable for genetic analysis, and has potential to contribute to future clinical characterization of hemolytic anemias.

Data and code availability
The derived datasets generated during this study are available from the UK Biobank (https://www.ukbiobank.ac.uk). Summary
Figure 5. Characterization of the MS4A7 locus

(A) The MS4A7 locus on chromosome 11 is associated with an increase in spleen iron. Purple triangle displays the lead SNP rs950802[G].

(B) This locus co-localizes with splicing quantitative trait locus of MS4A7 in whole blood (posterior probability \(R \geq 0.99\)).

(C) This splicing quantitative trait locus is associated with exon skipping in the MS4A7 locus. Corresponding betas and p value are shown. The MS4A7 gene model is displayed with the conserved CD20-like domain shown in green.

(D) Genetic associations with a panel of blood cell traits show associations with monocyte count, monocyte percentage, and mean platelet volume and negative associations with platelet count and platelet crit.

(E) Expression of MS4A7 was enriched in monocytes, including non-classical, intermediate, and classical forms.

(F) Expression of MS4A7 in human tissues displayed enrichment for lymphoid tissue, notably spleen and lymph nodes. DC, dendritic cell; MCHC, mean corpuscular hemoglobin concentration.
statistics are available from the GWAS catalog (https://www.ebi.ac.uk/gwas) at accession number GCST90101831. The code generated during this study is publicly available at https://github.com/calico/ukbb-mri-sseg and www.github.com/recoh/pipeline.

Supplemental information
Supplemental information can be found online at https://doi.org/10.1016/j.ajhg.2022.04.013.

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Author contributions
N.B., M.C., E.P.S., J.D.B., and E.L.T. designed the study. N.B., B.W., and Y.L. implemented the image processing methods. M.C. performed the data processing. E.P.S. and M.C. performed genetic analysis. E.L.T., E.P.S., R.L.C., M.C., and N.B. drafted the manuscript. All authors edited, read, and approved the manuscript.

Declaration of interests
E.P.S., R.L.C., and M.C. are employees of Calico Life Sciences LLC. Y.L. is a former employee of Calico Life Sciences LLC.

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