# Extent to which array genotyping and imputation with large reference panels approximate deep whole-genome sequencing

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This manuscript presents—across multiple ancestries, genotype arrays, and imputation reference panels—the minor-allele frequency thresholds above which array genotyping and imputation accurately capture genetic variants genotyped with deep whole-genome sequencing.



# Extent to which array genotyping and imputation with large reference panels approximate deep whole-genome sequencing

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### Summary

Understanding the genetic basis of human diseases and traits is dependent on the identification and accurate genotyping of genetic variants. Deep whole-genome sequencing (WGS), the gold standard technology for SNP and indel identification and genotyping, remains very expensive for most large studies. Here, we quantify the extent to which array genotyping followed by genotype imputation can approximate WGS in studies of individuals of African, Hispanic/Latino, and European ancestry in the US and of Finnish ancestry in Finland (a population isolate). For each study, we performed genotype imputation by using the genetic variants present on the Illumina Core, OmniExpress, MEGA, and Omni 2.5M arrays with the 1000G, HRC, and TOPMed imputation reference panels. Using the Omni 2.5M array and the TOPMed panel,  $\geq$ 90% of bi-allelic single-nucleotide variants (SNVs) are well imputed ( $r^2 > 0.8$ ) down to minor-allele frequencies (MAFs) of 0.14% in African, 0.11% in Hispanic/Latino, 0.35% in European, and 0.85% in Finnish ancestries. There was little difference in TOPMed-based imputation quality among the arrays with >700k variants. Individual-level imputation quality varied widely between and within the three US studies. Imputation quality also varied across genomic regions, producing regions where even common (MAF > 5%) variants were consistently not well imputed across ancestries. The extent to which array genotyping and imputation can approximate WGS therefore depends on reference panel, genotype array, sample ancestry, and genomic location. Imputation quality by variant or genomic region can be queried with our new tool, RsqBrowser, now deployed on the Michigan Imputation Server.

### Introduction

Short-read deep whole-genome sequencing (WGS) accurately captures most single-nucleotide variants (SNVs) and short indels across the genome and minor-allele fre-quency (MAF) spectrum.<sup>[1](#page-12-0)</sup> Advances in sequencing technologies, and corresponding decreases in sequencing cost, have enabled ever larger human sequencing studies. $2-5$  Such studies have identified rare alleles that cause Mendelian diseases $6-8$  and contribute to risk of com-mon diseases<sup>[9](#page-12-3)</sup> and variation in quantitative traits.<sup>[2](#page-12-1),[4](#page-12-4)</sup> However, deep WGS remains prohibitively expensive and computationally intensive for large studies. $1,10$  $1,10$ 

In contrast to WGS, genotype arrays assay hundreds of thousands to millions of variants, representing only a small fraction of genetic variation but at a much lower cost. Variants that are not array genotyped can be statistically inferred by comparing sample haplotypes to an external reference panel of sequenced haplotypes via geno-type imputation.<sup>[11](#page-12-6)</sup> Most common (MAF  $>$  5%) variants are present in recent reference panels and can be imputed with high accuracy from genotype arrays.<sup>5[,12,](#page-12-8)[13](#page-12-9)</sup> However, lowfrequency  $(0.5\% < \text{MAF} \leq 5\%)$  and rare (MAF  $\leq 0.5\%$ ) variants appear less often or may be absent from the reference panel, making their imputation less accurate or impos-sible.<sup>[14](#page-12-10)</sup> Therefore, using inexpensive genotype arrays and imputation in place of costly deep WGS can result in lower coverage and less accurate genotyping of rare genetic variation.

Reference panel, genotype array, sample ancestry, and genomic location all influence imputation quality. $12-15$ Previous studies have evaluated imputation quality with

<span id="page-1-5"></span><span id="page-1-4"></span><span id="page-1-3"></span><span id="page-1-2"></span><span id="page-1-1"></span><span id="page-1-0"></span><sup>1</sup>Department of Biostatistics and Center for Statistical Genetics, School of Public Health, University of Michigan, Ann Arbor, MI, USA; <sup>2</sup>Institute of Genetic Epidemiology, Medical University of Innsbruck, Innsbruck, Austria; <sup>3</sup>Department of Medicine and Department of Neurosciences, Université de Montréal, Montreal, QC, Canada; <sup>4</sup>Research Centre, Montreal Heart Institute, Montreal, QC, Canada; <sup>5</sup>Department of Psychiatry, University of California San Diego, La Jolla, CA, USA; <sup>6</sup>Research Institute, Bloodworks, Seattle, WA, USA; <sup>7</sup>Department of Medicine, University of Washington, Seattle, WA, USA; <sup>8</sup>Department of Medicine, Icahn School of Medicine at Mount Sinai, New York, NY, USA; <sup>9</sup>Department of Genetics and Genomic Sciences, Icahn School of Medicine at Mount Sinai, New York, NY, USA; <sup>10</sup>Institute for Genomic Health, Icahn School of Medicine at Mount Sinai, New York, NY, USA; <sup>11</sup>Department of Medicine, University of Washington, Seattle, WA, USA; 12Institute of Clinical Medicine, Internal Medicine, University of Eastern Finland, Kuopio, Finland; 13Charles Bronfman Institute for Personalized Medicine, Icahn School of Medicine at Mount Sinai, New York, NY, USA; <sup>14</sup>Stanley Center for Psychiatric Research, Broad Institute of MIT and Harvard, Cambridge, MA, USA; <sup>15</sup>Department of Genetics, Harvard Medical School, Boston, MA, USA; <sup>16</sup>Departments of Psychiatry, Rutgers University, Robert Wood Johnson Medical School and New Jersey Medical School, New Brunswick, NJ, USA; <sup>17</sup>Institute for Biomedicine (Affiliated with the University of Lübeck), Eurac Research, Bolzano, Italy

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The study name, ancestry, mean sequencing depth, sample size (total and unrelated subset), and number of variants, including single-nucleotide variants (SNVs) and indels, for the four WGS datasets.

the multiethnic 1000 Genomes Phase 3  $(1000G),$ <sup>[12](#page-12-8)</sup> the predominantly European Haplotype Reference Consortium  $(HRC)<sup>13</sup>$  $(HRC)<sup>13</sup>$  $(HRC)<sup>13</sup>$  and two releases of the multiethnic Trans-Omics for Precision Medicine (TOPMed)<sup>[5,](#page-12-7)[16](#page-12-11)</sup> panels, finding that larger, more deeply sequenced panels support more accurate imputation. Closer ancestry matching between the sample and reference panel haplotypes also improves imputation quality, particularly for rare variants, as demonstrated by the use of population-specific reference panels in isolated $17-19$  and non-isolated European populations. $2,20,21$  $2,20,21$  $2,20,21$  Likewise, denser genotype arrays are associated with higher imputation quality,  $13,14,22$  $13,14,22$  $13,14,22$  although the effect of array size on imputation quality has not been studied with the TOPMed panels. Regional variability in imputation quality with the 1000G panel is associated with genomic features including repeats and GC content, $15$ but the degree to which imputation quality varies across the genome with the larger HRC or TOPMed panels is unknown. It is also unknown to what extent individual-level imputation quality varies within populations for any reference panel.

Here, we determine the extent to which genotyping with the Illumina Core, OmniExpress, MEGA, and Omni 2.5M arrays followed by imputation with the 1000G, HRC, and TOPMed reference panels can approximate deep WGS in studies with individuals of African, Hispanic/Latino, non-Finnish European, and Finnish ancestries. Depending on the MAF of variants relevant to the research question, study ancestry, and genomic location, we found that array genotyping and imputation can approximate WGS. Our findings, together with our new RsqBrowser tool for querying imputation quality, should help guide investigator decisions between these two technologies.

### Material and methods

#### Genetic data resources

#### WGS data and processing

We used WGS data from the BioMe, $^{23}$  $^{23}$  $^{23}$  InPSYght, METSIM, $^{24,25}$  $^{24,25}$  $^{24,25}$  and MLOF<sup>26</sup> studies. Detailed descriptions of sample collection, sequencing, and data processing for BioMe and MLOF are provided by the TOPMed Informatics Research Center.<sup>5</sup> Corresponding information is available for the METSIM study.<sup>25</sup> The InPSYght study is a deep whole-genome sequencing US-based case-control study of individuals of admixed African-European or African genetic ancestry. Cases have either bipolar disorder or schizophrenia. The study is composed of samples from the Genomic Psychiatry Cohort (GPC), $27,28$  $27,28$  Consortium on the Genetics of Schizophrenia  $(COGS)<sub>29</sub>$  $(COGS)<sub>29</sub>$  $(COGS)<sub>29</sub>$  Bipolar Genome Study (BIGS)<sub> $30$ </sub> Lithium Treatment Moderate Dose Use Study (LiTMUS),<sup>[31](#page-13-7)</sup> and Systematic Treatment Enhancement Program for Bipolar Disorder (STEP-BD) studies,<sup>[32](#page-13-8)</sup> all obtained from the NIMH repository. Whole-genome sequencing of the samples (mean depth  $27 + -5.5$  X) was performed at the Broad Institute. We excluded individuals with sex mismatches ( $n = 20$ ), non-XX or XY sex karyotypes ( $n = 17$ ), duplicates (n = 366), >5% DNA contamination (n = 4), an excess of singletons ( $n = 39$ ), or <25% global African ancestry as determined by ADMIXTURE<sup>33</sup> analysis of array genotype data or for whom <98% of sites were at a sequencing depth of  $\geq$ 10 (n = 14). We complied with TOPMed data use agreements for the BioMe and MLOF studies. The InPSYght study was approved by the institutional review boards at each site. The METSIM study was approved by the ethics committee at the University of Eastern Finland and the institutional review board at the University of Michigan. All InPSYght and METSIM participants provided informed consent.

Participants from the BioMe biobank self-reported as Hispanic/ Latino and were recruited at the Mount Sinai Health System in New York City ( $n = 4,677$ ; [Table 1\)](#page-2-0). Participants in the MLOF study self-reported as non-Hispanic White and were recruited throughout the US ( $n = 2,987$ ). Participants in the METSIM study were recruited in Kuopio, Finland ( $n = 3,045$ ). On the basis of recruiting location and self-reported and genetic ancestry, we designated the population groups Hispanic/Latino (BioMe), African (InPSYght), Finnish (METSIM), and European (MLOF) ancestry for the purposes of this study.

In all studies, we removed participants inferred by KING<sup>[34](#page-13-10)</sup> to be related at a second degree or closer relationship to any other individual genotyped in TOPMed Freeze 9 ( $n = 157,675$ ), including all participants in these four studies and all individuals in the TOPMed imputation reference panel. This filtering yielded 3,141 participants in BioMe, 7,169 in InPSYght, 2,703 in METSIM, and 2,429 in MLOF. We then randomly downsampled to 2,429 individuals in each study [\(Figure S1](#page-11-0)).

WGS variant calling for all four studies was performed jointly with TOPMed Freeze 9 by the TOPMed Informatics Research Cen-ter (IRC) with the TOPMed Variant Calling/GotCloud pipeline.<sup>[5](#page-12-7),[35](#page-13-11)</sup>

We analyzed bi-allelic SNVs, multi-allelic SNVs, bi-allelic indels, and multi-allelic indels separately with n-allele variants recoded and analyzed as  $n-1$  bi-allelic variants at the same position.

#### Array genotyping in METSIM

METSIM participants were genotyped with the Illumina Human OmniExpress array. Variants with poor mapping of probes to GRCh37, call rate < 95%, or deviations from Hardy-Weinberg equilibrium (p  $< 10^{-6}$ ) were removed. $^{36}$  $^{36}$  $^{36}$ 

#### Genotype imputation

For each study, we subsetted WGS variants to those present on the Illumina Infinium Core (0.3M markers), Illumina Omni Express (0.7M), Infinium Omni 2.5M (2.4M), and Multi-Ethnic Genotyping (MEGA; 1.8M) arrays [\(Table S1](#page-11-0)). We refer to these WGS variant subsets as WGS-based arrays. For each study, we phased the selected variants with Eagle 2.4.1 and imputed genotypes by using Minimac4 on the Michigan Imputation Server (pipeline version 1.2.4)<sup>37</sup> with the (1) 1000 Genomes Phase 3 (n = 2,504), (2) Haplotype Reference Consortium  $(n = 32,470)$ , and (3) modified TOPMed ( $n = 88,804$ ) reference panels. MLOF and BioMe are included in the full, publicly available TOPMed R2 ( $n = 97,256$ ) panel. To avoid overlap of participants and the presence of close relatives in the reference panel, we removed 4,694 BioMe (4,668 Hispanic/Latino and 26 missing ethnicity) and 3,758 MLOF (2,977 non-Hispanic White and 781 missing race/ethnicity) individuals from the full TOPMed R2 panel to create our modified TOPMed panel.

## Evaluation of imputation quality

### Observed imputation  $r^2$

For each variant, we calculated the observed imputation  $r^2$  as the squared Pearson correlation coefficient between the imputed genotype dosages and the sequence-based genotypes. We assigned  $r^2 = 0$  for any variant present in the sequenced individuals but absent from the reference panels and so not imputed. We also assigned  $r^2 = 0$  for any variant with undefined correlation due to being monomorphic in the imputed data. For each variant category (bi-allelic SNVs, bi-allelic indels, multi-allelic SNVs, and multiallelic indels) and each WGS-based array, we calculated the proportion of variants that were well-imputed (observed imputation  $\rm r^2 > 0.8$ ) within study-specific MAF bins of size 0.00025 for MAF between 0.0002 and 0.002 and of size 0.001 for MAF > 0.002. We also calculated the mean  $r^2$  for each MAF bin. Each minor allele for multi-allelic variants was analyzed independently of the other minor alleles of the same variant so that multi-allelic variants had the same number of  $r^2$  measurements as minor alleles.

#### Genotype concordance

Separately for common, low-frequency, and rare bi-allelic SNVs, we calculated the heterozygous concordance rate between the imputed best-guess genotypes and sequenced-based genotypes as the proportion of heterozygous variants in WGS that were present in the reference panel that were also heterozygous in the imputed data by using bed-diff.<sup>38</sup> We excluded bi-allelic SNVs that were absent from the reference panels in these calculations.

#### Predicted variant consequences

In each study, we used  $VEP<sup>39</sup>$  to predict the functional consequences of bi-allelic SNVs. We partitioned variants into four classes based on the predicted impact on protein coding: high, moderate, low, and modifier. While variants in the high and moderate classes are likely to change protein behavior, variants in the low impact class are unlikely to do so. Modifier variants are mostly non-coding with no evidence of impacting protein coding.

#### Fine-scale ancestry estimation

For InPSYght, we estimated the proportion of African ancestry present in each individual by using  $RFMix^{40}$  $RFMix^{40}$  $RFMix^{40}$  with two reference groups representing African and European ancestry from 1000G. For BioMe, participants had previously been grouped by continental origin and into identity-by-descent (IBD) communities representing groups with shared recent genetic ancestry. $^{23}$  $^{23}$  $^{23}$  We labeled BioMe participants as from a Caribbean population if their continental origin was Caribbean or if they were members of the Puerto Rican or Dominican IBD communities. We labeled all other BioMe participants with non-missing continental origin as non-Caribbean. Participants not from Puerto Rican or Dominican IBD communities and with missing continental origin information were not included in comparisons between Caribbean and non-Caribbean populations.

We performed principal-component analysis (PCA) to obtain fine-scale ancestry information for all four studies. We used genotypes subset to those present on the Omni 2.5M array to project participants from each of the four studies onto the 938 reference samples from the Human Genome Diversity Project<sup>41</sup> by using the LASER server.<sup>[42](#page-13-18)</sup>

## Effect of regional genomic features on imputation quality

#### Genomic features datasets

We downloaded GC content over 5 bp intervals, the genomic positions of segmental duplications, the genomic positions of structural variants annotated with the Database of Genomic Variants, and the genomic positions of repeats identified with RepeatMasker from the UCSC Genome Browser database.<sup>[43](#page-13-19)</sup> Recombination rate was calculated with the HapMap GrCh38 genetic map<sup>44</sup> as centimorgans per megabase.

#### Relationships between genomic features and TOPMed imputation quality

In each study, we performed LD pruning to obtain a set of near-independent bi-allelic SNVs on chromosome 20, retaining variants with pairwise  $r^2$  < 0.2 within a sliding 50 kb window with a five variant step size with PLINK v2.0. $45-47$  For each retained variant, we defined five aggregate measures of genomic features over 10 kb windows centered at the variant: mean GC content, number of repeats, number of structural variants, presence of  $\geq 1$ segmental duplication, and mean recombination rate. We defined the linear distance of the variant from the nearest array-genotyped variant. For each of the six genomic features, we performed a logistic regression to test the association between dichotomous imputation quality (observed imputation  $r^2 > 0.8$  versus  $\leq 0.8$ ) and the feature, adjusting for variant MAF as a categorical variable with nine bins and breaks at 0, 0.0003, 0.0006, 0.0009, 0.001, 0.0032, 0.01, 0.032, 0.1, and 0.5. We also performed zero-one inflated beta regression to test the association between the continuous observed imputation  $r^2$  and each feature with the same MAF adjustment. Zero-one inflated beta regression models the association of the genomic features with the observed imputation  $r^2$ in the open interval  $0 < r^2 < 1$  (mean  $\mu$  and variance  $\sigma^2 \mu (1 - \mu)$ ) and the probabilities of observed imputation  $r^2 = 0$  (v) and observed imputation  $r^2 = 1$  ( $\tau$ ) in a piecewise manner.<sup>[48](#page-13-22)</sup> In both regression models, we centered and scaled continuous and count predictors for comparability.

### Effect of real versus WGS-based array genotypes on evaluation of imputation quality

To determine whether the WGS-based imputation results were consistent with the genotype array-based results, we imputed the real OmniExpress arrays with each of the three reference panels in METSIM. For each reference panel, we compared the observed imputation  $r^2$  and genotype concordance metrics for the real array-based genotype imputation to WGS-based array genotype imputation results (from above).

## Effect of variant caller on evaluation of imputation quality

Variants in TOPMed and the four study datasets were called with the TOPMed Variant Calling/GotCloud pipeline.<sup>[5](#page-12-7),[35](#page-13-11)</sup> To assess the impact of variant calling tool, we used an additional set of WGS variants called with Haplotype Caller from GATK version  $3.5<sup>49</sup>$  $3.5<sup>49</sup>$  $3.5<sup>49</sup>$ in the METSIM study.<sup>50</sup> Variants that deviated from Hardy-Weinberg equilibrium ( $p < 10^{-6}$ ), had >2% missingness, or had allelic imbalance <0.3 or >0.7 were excluded from this callset. Variants in regions of low complexity, centromeres, segmental duplications, or satellite regions were also excluded.<sup>50</sup> After filtering, 21.8M variants remained in the subset of 2,429 individuals used for imputation analysis in METSIM. We then created each of the four WGS-based arrays by using both METSIM callsets and evaluated imputation performance by comparing the imputed variants to the respective sequenced variants.

### Imputability tool for the Michigan Imputation Server

We developed RsqBrowser, a tool that allows researchers to query for the observed imputation  $r^2$  for variants or regions of interest. Users specify the genomic position in build GRCh38 and select the genotype array, imputation reference panel, and sample ancestry. RsqBrowser returns a table with the position and observed imputation  $r^2$  for all variants in the specified regions or genes. We have deployed this tool on the Michigan Imputation Server.

## Results

### Whole-genome sequencing studies of four ancestries

We used WGS data in four studies as gold standard genotypes. These four represent three major US populations, African American/African Diaspora (African), Hispanic/ Latino, and European American (European) ancestry, and a population isolate, Finnish ancestry from Finland [\(Ta](#page-2-0)[ble 1](#page-2-0)). We observed that our primary metric of imputation quality, the observed imputation  $r^2$ , was upwardly biased in small samples for low-frequency  $(0.5\% < \text{MAF} \leq 5\%)$ and rare (MAF  $\leq$  0.5%) variants ([Figure S1](#page-11-0)). To avoid any biases comparing across datasets of different sample sizes, we randomly downsampled the African, Hispanic/Latino, and Finnish ancestry datasets to 2,429 individuals to match the smaller European ancestry dataset. After all sample- and variant-level filtering, we included in our analysis 79.3M, 68.8M, 62.2M, and 22.0M variants in the African, Hispanic/Latino, European, and Finnish ancestry studies, respectively [\(Table 1](#page-2-0)). In each study, >91% of these variants were bi-allelic SNVs. The others were multi-allelic

SNVs (0.3%–1.6%), and bi-allelic (6.5%–6.6%) and multiallelic indels (0.005%–0.2%).

## Impact of reference panel on genotype imputation quality

For each WGS study participant, we subsetted WGS genotypes to those present on the Illumina Core (0.3M markers), OmniExpress (0.7M), Multi-Ethnic Genotyping (MEGA) (1.8M), and Omni 2.5M (2.4M) arrays. We then carried out genotype imputation on these genotype array subsets by using the 1000G and HRC imputation reference panels, as well as a modified TOPMed panel. Because the Hispanic/Latino and European WGS datasets were included in the TOPMed panel, we restricted the TOPMed panel to a subset of 88,804 reference samples that did not overlap our WGS datasets for all analyses. To measure the imputation quality of each sequenced variant in the study, we calculated the squared Pearson correlation between sequenced genotypes and imputed genotype dosages (observed imputation  $r^2$ ). We considered two aggregate measures of imputation quality: the mean observed imputation  $r^2$  and the proportion of well-imputed variants (observed imputation  $r^2 > 0.8$ ) by MAF bin. We selected a stringent threshold (>90% of variants well imputed) to define sets of variants for which array genotyping followed by imputation can approximate WGS. For variants satisfying this threshold, the two technologies are expected to provide very similar information.<sup>51</sup>

As expected, across all combinations of reference panels, ancestries, and MAF, the densest genotype array (Omni 2.5M) had both the highest mean observed imputation  $r^2$  and highest number and proportion of wellimputed variants [\(Figure 1](#page-5-0), [Figure S2,](#page-11-0) [Tables S2](#page-11-0) and [S3\)](#page-11-0). For the Omni 2.5M array and in all ancestries, TOPMedbased imputation approximated WGS for variants of lower MAF compared to the HRC or 1000G panels. TOPMED-based panel imputation approximated WGS at lower MAF thresholds in African and Hispanic/Latino ancestry (0.14% and 0.11%) than in European or Finnish ancestry (0.35% and 0.85%) ([Figures 1](#page-5-0)A and 1B, [Table S4\)](#page-11-0). The ordering of imputation quality by reference panel, genotyping array, and ancestry did not change when using mean observed imputation  $r^2$  as a metric of average imputation quality [\(Figure S2\)](#page-11-0). The ordering of results for both metrics also did not change by either the use of subsets of WGS genotypes instead of actual genotyping arrays [\(Figure S3](#page-11-0)) or the choice of variant caller ([Figure S4\)](#page-11-0).

## Less influence of genotype array size with TOPMed compared to 1000G- and HRC-based imputation

For all four ancestries and all three imputation reference panels, imputation quality increased with larger array size [\(Figure 1](#page-5-0)C). However, the difference in TOPMed imputation quality among the Omni 2.5M, MEGA, and OmniExpress arrays was minimal. For example, in the

<span id="page-5-0"></span>



(B) Comparison across the four studies using the Illumina Omni 2.5M array.

(C) Comparison across four Illumina genotyping arrays: Omni 2.5M, MEGA, Omni Express, and Core by ancestry (columns) and imputation reference panels (rows). In all plots, the x axes show minor-allele frequency (MAF) calculated separately by study. Sequenced biallelic SNVs not present in reference panels were assigned  $r^2 = 0$ . Bi-allelic SNVs were then aggregated by MAF bins of width 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.032, 0.1, 0.32, and 0.5.

African ancestry study, TOPMed imputation approximated WGS for variants with MAF  $\geq 0.14\%$  with the Omni 2.5M array,  $\geq 0.16\%$  with the MEGA array, and  $\geq$ 0.24% with the OmniExpress array [\(Table S4\)](#page-11-0). This threshold was higher with the smaller Core array  $(\geq 0.84\%)$ . In contrast, genotype array size had a larger effect on imputation quality with the HRC and 1000G

panels in African and Hispanic/Latino ancestry studies. For African ancestry, 1000G imputation approximated WGS at a much lower MAF with the Omni 2.5M array  $(\geq 2.5\%)$  compared to the OmniExpress array ( $\geq 14.0\%$ ). With the HRC panel, imputation with the OmniExpress array could not approximate WGS at any MAF in African ancestry.

#### Individual-level imputation accuracy varies with finerscale ancestry

Because imputation quality depends on the shared ancestry between reference panel and sample haplotypes, $11$  we hypothesized that imputation quality within the four WGS studies would vary with finer-scale ancestry. To measure individual-level imputation quality, we calculated concordance rates between heterozygous sequenced and imputed genotypes separately for study-specific rare, low-frequency, and common bi-allelic SNVs in each individual. As expected, concordance rates varied across individuals more for rare variants than for low-frequency and common variants (TOPMed: [Figures 2,](#page-7-0) [S5,](#page-11-0) and [S6](#page-11-0); all panels: [Table S5\)](#page-11-0). With the TOPMed panel, mean heterozygous concordance rates for rare variants were higher in individuals of African and Hispanic/Latino ancestry (0.93 in both) compared to individuals of European and Finnish ancestry (0.86 and 0.82) [\(Figure 2A](#page-7-0)). Concordance rates varied most within Hispanic/Latino individuals  $(10^{th} 90<sup>th</sup>$  percentile: 0.80–0.98).

We next stratified African and Hispanic/Latino study participants by finer-scale measures of ancestry. The African American population in the United States is primarily of African and European ancestries.<sup>[52](#page-13-26)</sup> We therefore estimated the proportion of African ancestry for each individual in the African ancestry study assuming two populations, which ranged from 0.26 to 1.00 (mean 0.82). For the 2,307 individuals with an estimated African ancestry < 0.95, individuals with higher proportions of African ancestry had higher genotype concordance rates with the TOPMed panel ([Figure 2B](#page-7-0)). For instance, concordance rates for those with an estimated proportion between 0.86 and 0.95 were higher (mean 0.93) than for those between 0.26 and 0.35 (mean 0.89). In contrast, concordance rates for the 122 individuals with estimated proportion of African ancestry  $> 0.95$  were lower (mean 0.91) than for individuals with smaller estimated proportions of African ancestry.

Hispanic/Latino populations in the United States are admixed with primarily European, Native American, and African ancestry, and individuals of Caribbean origin usually have more African ancestry. $53$  The concordance rates for individuals from Caribbean populations were higher (mean 0.96) compared to those from non-Caribbean (mean 0.79) populations with the TOPMed panel ([Figure 2C](#page-7-0)).

We also estimated finer-scale ancestry in all four studies with principal-component analysis (PCA), projecting the study individuals onto 938 reference samples from the Hu-man Genome Diversity Project (HGDP).<sup>[41](#page-13-17)</sup> The first two principal components (PCs) reflect clines of European (high PC2), African (high PC1 and low PC2), and Native American (low PC1 and low PC2) ancestries [\(Figure S7\)](#page-11-0). To see how TOPMed imputation quality varied with finescale ancestry, we divided individuals into concordance rate quintiles calculated jointly across all four studies. In all studies, individuals clustering closer to HGDP individuals of African ancestry were more likely to be in higher concordance rate quintiles, while those clustering closer to Native American or European populations were more likely to be in lower concordance rate quintiles for rare ([Figure 2D](#page-7-0)) and low-frequency variants [\(Figure S5](#page-11-0)D). As expected, there was little variability in common variant imputation quality ([Figure S6](#page-11-0)D).

Taken together, these results demonstrate that TOPMed imputation quality varies across individuals with finerscale ancestry. Among the populations studied here, population subsets with large proportions of African ancestry, including Hispanic/Latino ancestry individuals of Caribbean origin, were on average the most accurately imputed for rare variants. However, individuals with the greatest proportions of African ancestry in the African study were not the most accurately imputed. The heterozygote concordance rates from HRC and 1000G imputation also varied with finer-scale ancestry for rare variants [\(Table S5\)](#page-11-0).

### Imputation quality varies across the genome

Sequence quality and genotype array density are not uniform across the genome. Because these factors influence imputation, we sought to quantify the regional variability in imputation quality. We first visualized the observed imputation  $r^2$  for common variants (MAF  $>$  5%) across the chromosomes. Although the vast majority (>99.6%) of common variants are well imputed (observed imputation  $r^2 > 0.8$ ) in all four ancestries with the Omni 2.5M array and TOPMed reference panel ([Table S3](#page-11-0)), we identified clusters of common variants that were not well imputed at the same genomic positions across ancestries and genotype arrays [\(Figures 3](#page-8-0)A and [S8\)](#page-11-0). There are likewise regions with better-than-average imputation quality, including the HLA region on chromosome 6 [\(Figure S8](#page-11-0)), which is characterized by high LD and dense genotype array coverage. $22,54$  $22,54$ 

To assess regional variability in imputation quality, we calculated the lengths of runs of consecutively wellimputed variants separately for rare, low-frequency, and common bi-allelic SNVs across the genome ([Figure 3B](#page-8-0), [Tables S6](#page-11-0) and [S7\)](#page-11-0). We identified a large variability in the number of consecutively well-imputed common and lowfrequency variants with the TOPMed panel (e.g., interquartile range [IQR] in African ancestry is 41–750 common variants [10.4–253.2 kb] and 9–287 low-frequency variants [2.3–84.3 kb] with the Omni 2.5M array). As expected, the lengths of consecutive well-imputed rare variants were much shorter, having a maximum length of 34–45 variants depending on ancestry.

## Local genomic features explain little variability in imputation quality

Genomic features including high GC content and the presence of large duplications or repeats have been associated with regions of poor imputation quality in Europeans when using the  $1000G$  panel.<sup>[15](#page-12-16)</sup> To test the effects of genomic features on imputation quality with the TOPMed panel, we performed logistic regressions in each of our

<span id="page-7-0"></span>

Figure 2. Heterozygous genotype concordance rates for rare variants by ancestry with TOPMed panel imputation Heterozygous concordance rates were calculated between sequenced and TOPMed-imputed genotypes for rare (MAF < 0.5%, calculated separately in each study) bi-allelic SNVs with the Omni 2.5M array.

(A) Distribution of concordance rates in each of the four studies. Boxplots correspond to  $25<sup>th</sup>$ ,  $50<sup>th</sup>$ , and  $75<sup>th</sup>$  percentiles.

(B) Distribution of concordance rates by bins of estimated proportion of African ancestry in the admixed African study.

(C) Distribution of concordance rates in Caribbean and non-Caribbean populations in the Hispanic/Latino study.

(D) Principal-component analysis (PCA) by genotype concordance quintile and ancestry. PCA was performed by projecting onto the Human Genome Diversity Project reference samples. Genotype concordance quintiles were calculated across all four studies and correspond to concordance rates of 0.37–0.82 (Q1), 0.82–0.86 (Q2), 0.86–0.93 (Q3), 0.93–0.95 (Q4), and 0.95–0.99 (Q5). Points are colored by ancestry.

four studies with the imputed quality status (observed imputation  $r^2 > 0.8$ ) as the dichotomous outcome for independent variants on chromosome 20. In separate models, we tested the associations of distance to the nearest genotyped variant and the following features aggregated over a 10 kb window centered at the variant: mean GC content, mean recombination rate, number of repeats, number of structural variants, and presence of  $\geq$ 1 segmental duplication, adjusting for bins of MAF. We found that higher recombination rate, lower GC content, greater distance to genotype array variants, more structural variants, and the presence of segmental duplications were all associated with lower imputation quality [\(Figure 4A](#page-9-0), [Table S8](#page-11-0)). The effect of nearby repeats was not consistent across ancestries or

<span id="page-8-0"></span>

Figure 3. Regional variability in TOPMed reference panel imputation quality

(A) Observed imputation  $r^2$  by genomic position (Mb) of common (MAF > 0.05) bi-allelic SNVs on chromosome 20. Sequenced bi-allelic SNVs not present in reference panels were assigned  $r^2 = 0$ . The horizontal line at  $r^2 = 0.8$  corresponds to the threshold used to determine well-imputed variants.

(B) Cumulative distribution of the number of consecutively well-imputed ( $r^2 > 0.8$ ) bi-allelic SNVs in each MAF category: common (MAF  $\geq$  0.05), low frequency (0.005  $\leq$  MAF  $<$  0.05), and rare (MAF  $<$  0.005), as calculated separately in each study. For common variants, European and Finnish curves appear to overlap and African and Hispanic/Latino curves appear to overlap.

repeat class, although nearby simple repeats were associated with worse imputation quality in all ancestries [\(Figure S9,](#page-11-0) [Table S9](#page-11-0)). However, none of the tested genomic features meaningfully impacted the proportion of variability in imputation quality beyond variant MAF [\(Figure 4](#page-9-0)B). Results were similar when modeling imputation quality as a continuous variable ([Figure S10](#page-11-0), [Table S10\)](#page-11-0) and were consistent across reference panels [\(Table S8](#page-11-0)).

### Impact of variant predicted function and type on imputation quality

Protein-coding variants are often of high clinical significance and easier to interpret compared to non-coding variants; they are also more likely to be rare and more difficult to impute. $5,55$  $5,55$  To determine the extent to which variants that impact protein coding are well imputed, we classified sequenced bi-allelic SNVs by predicted impact on protein coding. With the TOPMed panel and Omni 2.5M array, we found that 45.6%–64.1% of variants predicted to have high impact and 51.0%–66.8% of variants predicted to have moderate impact on protein coding were well imputed [\(Table S11\)](#page-11-0). We found no meaningful difference in imputation quality between the protein-coding classes when controlling for MAF [\(Figure S11](#page-11-0)).

All results presented above are for bi-allelic SNVs. Multiallelic SNVs and all indels have been shown to have lower imputation quality than bi-allelic SNVs with the 1000G panel, $12,15$  $12,15$  and indels are absent from the HRC panel.<sup>[13](#page-12-9)</sup> To quantify the effect of variant type on imputation quality, we calculated the proportion of well-imputed indels, multi-allelic SNVs, and multi-allelic indels by using the TOPMed panel and all four genotype arrays. We observed very similar MAF thresholds for which imputation could approximate WGS among bi-allelic SNVs, bi-allelic indels, and multi-allelic SNVs [\(Figure S12\)](#page-11-0). Multi-allelic indels were less well imputed. For example, in African ancestry, TOPMed imputation with the Omni 2.5M array approximated WGS at similar MAF thresholds for bi-allelic SNVs and indels and multi-allelic SNVs (0.14%, 0.24%, and 0.16% respectively) compared to 0.55% for multi-allelic indels [\(Table S12](#page-11-0)).

## **Discussion**

Here, we used deep WGS from studies of African, Hispanic/ Latino, European, and Finnish ancestries to quantify the extent to which array genotyping followed by genotype imputation can approximate WGS. We performed imputation by using genotypes present on the Illumina Core, OmniExpress, MEGA, and Omni 2.5M arrays with the 1000G, HRC, and TOPMed reference panels. We found

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Figure 4. Genomic features associated with TOPMed imputation quality of bi-allelic SNVs by ancestry (A) The odds ratios and corresponding unadjusted 95% confidence intervals from logistic regression models. Estimates are from separate

models testing the associations between characteristics of regional genomic features and whether or not a variant is well imputed (observed imputation  $r^2 > 0.8$ ), adjusting for variant MAF.

(B) The proportion of variance explained (Nagelkerke R<sup>2</sup>) for each logistic regression models with MAF only or with MAF and all six tested genomic features in one joint model.

that with the largest array (Omni 2.5M) and largest reference panel (TOPMed), array genotyping followed by imputation can approximate WGS at a population level for variants with MAF  $\geq 0.14\%$  in African ancestry,  $\geq 0.11\%$  in Hispanic/Latino ancestry,  $\geq$ 0.35% in European ancestry, and  $\geq$  0.84% in Finnish ancestry. Particularly for the African and Hispanic/Latino ancestry studies, TOPMed imputation approximated WGS at much lower MAF than HRC or 1000G imputation, which is consistent with previous analyses showing improvements in these populations even with a smaller version of the TOPMed panel.<sup>[16](#page-12-11)</sup> For analyses primarily investigating the genetic effects of common and low-frequency variants, such as single-variant genome-wide association studies (GWASs), in any of the four populations, array genotyping and imputation is sufficient to accurately capture genetic variants and given differences in cost allows for much larger sample sizes than WGS. Large proportions  $(\sim44\% - 60\%)$  of rare variants with MAF even lower than the reported thresholds were also well-imputed, highlighting the potential for well-powered rare-variant studies without WGS, although not all rare variants can be reliably imputed. Because we restricted the TOPMed panel to reference samples that did not overlap the WGS datasets, we expect that imputation quality with the full TOPMed R2 panel to be even better than reported here. In particular, we would expect higher imputation quality for Hispanic/Latino studies, as a large proportion of the Hispanic/Latino individuals in the TOPMed R2 panel were excluded here.

As previously reported with the 1000G and HRC panels, $12,13$  $12,13$  imputation quality with the TOPMed panel was higher when using larger arrays. However, the effect of genotype array choice on TOPMed imputation was much smaller than on HRC or 1000G imputation. The difference between the Omni 2.5M, MEGA, and OmniExpress arrays was minimal, suggesting that researchers imputing with the TOPMed panel in these populations may opt for the less expensive OmniExpress array with little loss of information. However, we did find lower imputation quality when using the smaller Core array  $(\sim]307k$  variants) and might expect even lower quality for arrays with fewer markers.

WGS is also used for clinical purposes, including diag-nosis, screening, and identifying therapeutic targets.<sup>[6](#page-12-2)</sup> Variants predicted to alter protein function are often of high clinical significance.<sup>[55](#page-14-2)</sup> In the populations studied here, we found that only 50.7%–66.7% of bi-allelic SNVs with moderate or high predicted impact on protein coding were well imputed, as might be expected given the generally low MAF of these variants. To quantify individual imputation quality in contrast to population-level imputation quality, we calculated the heterozygous concordance rates between sequenced bi-allelic SNVs and imputed best-guess genotypes. For all three reference panels, we found that the concordance rates for rare and low-frequency variants varied widely among individuals in the African, Hispanic/Latino, and European ancestry studies and were associated with finer-scale ancestry. Because of this variability and the large proportion of rare variants that are not accurately imputed with the available imputation reference panels, we believe that WGS cannot currently be reliably approximated in clinical settings with array genotyping and imputation with the reference panels studied here. Recent work in cystic fibrosis-affected individuals has shown that disease-specific WGS panels can provide additional information for imputation of diseasecausing variants and may play a role in future clinical use. $56$ 

Despite large numbers of individuals with African and Hispanic/Latino ancestry in the TOPMed reference panel, more than half of the individuals in the panel are estimated to be of >50% European ancestry. Still, we found that TOPMed imputation quality was highest for the African and Hispanic/Latino ancestry studies and for individuals with large proportions of African ancestry among the populations studied here. Higher imputation quality of rare variants in African populations compared to European populations was previously reported with the 1000G panel.<sup>[12](#page-12-8)</sup> A first possible explanation for the high relative imputation quality in African populations is that there are proportionally more rare variants in non-African populations that have undergone recent bottlenecks and subsequent population growth, as is true in the three US populations studied here [\(Figure S13\)](#page-11-0). In these populations, it can be more difficult to identify the haplotype background of the rare variation.<sup>[12](#page-12-8),[57](#page-14-4)</sup> A second possible explanation is that individuals with large proportions of African ancestry in these studies match more closely by chance with a subset of TOPMed haplotypes than do the individuals with large proportions of Native American or European ancestry. However, relatively higher imputation quality with the TOPMed panel in samples of African and Hispanic/Latino ancestry compared to European ancestry was previously reported in a separate set of samples.<sup>[5](#page-12-7)</sup> Third, admixture could impact the accuracy of haplotype phasing of the sample or reference haplotypes. Taken together, these results emphasize the importance of ancestrally diverse reference panels such as TOPMed and suggest that reference panel composition is not the only factor explaining ancestry differences in imputation quality.

While nearly all common and low-frequency variants are well imputed with the TOPMed panel in the populations studied, there was substantial variability by genomic region in imputation across the MAF spectrum. Some regions, such as the densely genotyped HLA locus, had higher imputation quality than what would be expected on the basis of variant MAF alone. We found that lower recombination rate and higher GC content around a variant were associated with higher imputation quality in all four studies but that none of these features except MAF explained a substantial proportion of variability in imputation quality. For all tested features except for GC content and repeats, the direction of effect was consistent with previous work examining regions of poor imputation quality via the 1000G panel in Europeans.[15](#page-12-16) The inconsistency in GC content direction

could be explained by our focus on well-imputed variants instead of poorly imputed regions and/or by continued improvements in sequencing quality. Given the difficulty of predicting hard-to-impute regions/variants, we developed RsqBrowser, a tool that allows researchers to query empirical imputation quality for specific variants or genomic regions of interest by ancestry, which is available on theMichigan Imputation Server.

The results presented here are limited by the use of high quality but imperfect WGS as a gold standard. We did not consider any variants that were imputed from the reference panels but not detected in the WGS. Comparisons of imputation quality by reference panel or genotyping array did not change when using real array data compared to WGS-based arrays or when using a different variant caller, although data were only available to carry out these sensitivity analyses in the study of Finnish ancestry. We also note that the results presented here cannot necessarily be extended to other populations or population isolates, particularly those such as East and South Asian populations, that are not represented or represented in smaller numbers in the TOPMed panel. Furthermore, we only used WGS from one study for each population that we analyzed. For some ancestries, particularly population isolates, other population-specific reference panels may perform better than the three commonly used imputation panels analyzed here.

While array genotyping and imputation cannot fully replace deep WGS, we found that it can approximate WGS for variants down to specific MAF thresholds depending on genotype array and reference panel choices as well as sample ancestry. Researchers' decision to invest in one technology over another will depend on these criteria, genomic location, and the MAF of variants relevant to their research questions.

## Consortia

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### Data and code availability

Genotypes can be accessed through dbGaP for the BioMe (dbGaP: phs001644) and MLOF (dbGaP: phs001515) studies. METSIM and InPSYght genotype data are not currently available because of privacy restrictions. The variant-level  $r^2$  statistics generated during this study are available to download from RsqBrowser [\(https://](https://imputationserver.sph.umich.edu/rsq-browser/downloads) [imputationserver.sph.umich.edu/rsq-browser/downloads\)](https://imputationserver.sph.umich.edu/rsq-browser/downloads). Analysis code and scripts are available at [https://github.com/schanks/](https://github.com/schanks/imputation_rsqbrowser) [imputation\\_rsqbrowser.](https://github.com/schanks/imputation_rsqbrowser)

### Supplemental information

Supplemental information can be found online at [https://doi.org/](https://doi.org/10.1016/j.ajhg.2022.07.012) [10.1016/j.ajhg.2022.07.012](https://doi.org/10.1016/j.ajhg.2022.07.012).

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Full acknowledgments are provided in the [supplemental](#page-11-0) [information.](#page-11-0)

### Declaration of interests

E.E.K. has received speaker honorarium from Regeneron, 23&Me, and Illumina and sits on the advisory board for Encompass Bio and Galateo Bio.

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### Web resources

RsqBrowser, [https://imputationserver.sph.umich.edu/rsq](https://imputationserver.sph.umich.edu/rsq-browser)[browser](https://imputationserver.sph.umich.edu/rsq-browser)

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## Supplemental information

## Extent to which array genotyping and

## imputation with large reference panels

## approximate deep whole-genome sequencing

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## Supplemental Figures



**Figure S1. Effect of sample size on imputation quality metrics.** Random subsets of individuals were taken from each of the WGS studies as the total sample size of unrelated individuals allowed (up to 7,000 for African, 3,000 for Hispanic/Latino, and 2,000 for European and Finnish). Imputation was performed with the Omni 2.5M array and the TOPMed imputation reference panel. A. The proportion of sequenced biallelic SNVs that are well-imputed ( $r^2$ >0.8) by sample size. B. The mean r2 by sample size. In both plots, the x-axes show minor allele frequency (MAF) calculated separately by study based on the 2,429 samples used in the main analyses. Sequenced biallelic SNVs not present in reference panels were assigned  $r^2$ =0. Biallelic SNVs were then aggregated by MAF bins of width 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.032, 0.1, 0.32, and 0.5.



**Figure S2. Mean observed imputation r2 of biallelic SNVs by reference panel, study ancestry, and genotyping array.** The mean observed imputation r<sup>2</sup> with the TOPMed, HRC, and 1000G imputation reference panels. A. Comparison across the reference panels using the Illumina Omni 2.5M array. B. Comparison across the four studies using the Illumina Omni 2.5M array. C. Comparison across four Illumina genotyping arrays: Omni 2.5M, MEGA, Omni Express, and Core by ancestry (columns) and imputation reference panels (rows). In all plots, the x-axes show minor allele frequency (MAF) calculated separately by study. Sequenced biallelic SNVs not present in reference panels were assigned  $r^2$ =0. Biallelic SNVs were then aggregated by MAF bins of width 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.032, 0.1, 0.32, and 0.5.



**Figure S3. Imputation quality of biallelic SNVs by reference panel using WGSbased and real Illumina OmniExpress arrays.** A. The proportion of sequenced biallelic SNVs imputed from real array data (red line) or from WGS-based array (blue line) in the Finnish study that are well-imputed ( $r^2$ >0.8) by imputation reference panel. B. The mean observed imputation  $r^2$  for the same variants. In all plots, the x-axes show minor allele frequency (MAF) calculated separately by study. Variants were aggregated by MAF bins of size 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.316, 0.1, and 0.5. The lines appear entirely overlapping for the HRC and 1000G reference panels.



Figure S4. Proportion of well-imputed (r<sup>2</sup>>0.8) biallelic SNVs by reference panel,

**genotyping array, and variant caller in Finnish study.** The proportion of sequenced biallelic SNVs called with the GotCloud pipeline (red line) or GATK pipeline (blue line) in the Finnish study that are well-imputed  $(r^2>0.8)$  by reference panel (rows) and genotyping array (columns). In all plots, the x-axes show minor allele frequency (MAF) calculated separately by study. Variants were aggregated by MAF bins of size 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.316, 0.1, and 0.5.



**Figure S5. Heterozygous genotype concordance rates for low-frequency variants by ancestry with TOPMed panel imputation.** Heterozygous concordance rates were calculated between sequenced and TOPMed imputed genotypes for low-frequency (0.5%<MAF<5%, calculated separately in each study) biallelic SNVs with the Omni2.5M array. A. Distribution of concordance rates in each of the four studies. Boxplots correspond to 25th, 50th, and 75th percentiles. B. Distribution of concordance rates by bins of estimated proportion of African ancestry in the admixed African study. C. Distribution of concordance rates in Caribbean and non-Caribbean populations in the Hispanic/Latino study. The inset figures in panels A-C show the same distributions with a restricted y-axis. D. Principal component analysis (PCA) by genotype concordance quintile and ancestry. PCA was performed by projecting onto the Human Genome Diversity Project reference samples. Genotype concordance quintiles were calculated across all four studies and correspond to concordance rates of 0.903-0.964 (Q1), 0.964-0.971 (Q2), 0.971-0.973 (Q3), 0.973-0.974 (Q4), and 0.974-0.974 (Q5). Points are colored by ancestry.



**Figure S6. Heterozygous genotype concordance rates for common variants by ancestry with TOPMed panel imputation.** Heterozygous concordance rates were calculated between sequenced and TOPMed imputed genotypes for common (MAF>5%, calculated separately in each study) biallelic SNVs with the Omni2.5M array. A. Distribution of concordance rates in each of the four studies. Boxplots correspond to 25th, 50th, and 75th percentiles. B. Distribution of concordance rates by bins of estimated proportion of African ancestry in the admixed African study. C. Distribution of concordance rates in Caribbean and non-Caribbean populations in the Hispanic/Latino study. The inset figures in panels A-C show the same distributions with a restricted y-axis. D. Principal component analysis (PCA) by genotype concordance quintile and ancestry. PCA was performed by projecting onto the Human Genome Diversity Project reference samples. Genotype concordance quintiles were calculated across all four studies and correspond to concordance rates of 0.974-0.995 (Q1), 0.995-0.996 (Q2), 0.996-0.996 (Q3), 0.996-0.997 (Q4), and 0.997-0.997 (Q5). Points are colored by ancestry.



**Figure S7. Principal component analysis of WGS samples.** PC1 and PC2 for the four WGS studies and Human Genome Diversity Project (HGDP) reference samples from Africa (n=129), Europe (n=156), and Native America (n=63). PCA was performed by projecting onto all HGDP reference samples (n=938).





























## **Figure S8. Regional variability in imputation quality of common variants with the TOPMed reference panel by genotyping array and ancestry across all**

**chromosomes.** Observed imputation r<sup>2</sup> by genomic position (Mb) for common (MAF>0.05) biallelic SNVs across all chromosomes by genotyping array (columns) and ancestry (rows). Variants above the horizontal black lines are well-imputed (observed imputation  $r^2 > 0.08$ ).



**Figure S9. Repeat classes associated with TOPMed imputation quality of biallelic SNVs by ancestry.** The odds ratios and corresponding 95% confidence intervals from logistic regression models. Estimates are from separate models testing the associations between each repeat class and whether or not a variant is well-imputed (observed imputation r2>0.8) adjusting for variant MAF. Repeat classes as defined by RepeatMasker include DNA repeat elements (DNA), long interspersed repeated elements (LINE), low complexity repeats (LowComplex), long terminal repeat elements including retrotransposons (LTR), rolling circle repeats (RC), RNA repeats (RNA), satellite repeats, microsatellites (Simple), short interspersed repeat elements including ALUs (SINE), and repeats of unknown class.



**Figure S10. Genomic features associated with TOPMed imputation quality of biallelic SNVs by ancestry.** The odds ratios and corresponding 95% confidence intervals from zero-one inflated beta regression models testing the association of genomic features with the observed imputation  $r^2$  in the open interval 0<r<sup>2</sup><1 (mean  $\mu$ and variance-related parameter  $\sigma$ ) and the probabilities of observed imputation r<sup>2</sup>=0 (v) or  $r^2$ =1 ( $\tau$ ). Estimates are from separate models testing the associations between characteristics of regional genomic features and imputation quality (observed imputation r 2) adjusting for variant MAF.



Figure S11. Proportion of well-imputed (r<sup>2</sup>>0.8) biallelic SNVs by predicted **functional impact and ancestry**. The predicted functional impact of all sequenced biallelic SNVs was determined with VEP. The x-axes show minor allele frequency (MAF) calculated separately by study. Biallelic SNVs were then aggregated by MAF bins of width 0.00025 MAF for MAF between 0.0002 and 0.002 and of size 0.001 MAF for MAF > 0.002; those plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.001, 0.0032, 0.01, 0.032, 0.1, 0.32, and 0.5.



## Figure S12. Proportion of well-imputed (r<sup>2</sup>>0.8) variants by variant type,

**genotyping array, and ancestry with the TOPMed panel.** The proportion of sequenced variants that are well-imputed by genotyping array (rows) and ancestry (columns). X-axes show minor allele frequency (MAF) calculated separately in each study. Sequenced variants not present in reference panels were assigned  $r^2=0$ . Variants were then aggregated by MAF bins of width 0.00025 MAF for MAF between 0.0002 and 0.002, bins of width 0.001 MAF for MAF between 0.002 and 0.4, and one bin of width 0.1 MAF for MAF between 0.4 and 0.5. MAF bins plotted here correspond to singletons, doubletons, and tripletons in each study, as well as those with mean MAF closest to the values 0.01, 0.0032, 0.01, 0.316, 0.1, and 0.5.





## Supplemental Tables



**Table S1. Whole genome sequencing (WGS)-based genotype arrays.** The numbers of variants included on the Illumina arrays and the actual number of WGS variants in each study used to create the WGS-based arrays.



**Table S2. Number of well-imputed biallelic single nucleotide variants (SNVs) in each whole genome sequencing (WGS) study by reference panel, genotype array, ancestry, and minor allele frequency (MAF) category.**



**Table S3. Proportion of biallelic single nucleotide variants (SNVs) in each whole genome sequencing (WGS) study that are well-imputed (r 2 >0.8) by reference panel, genotype array, ancestry, and minor allele frequency (MAF) category.**



**Table S4. Minor allele frequency (MAF) threshold above which array genotyping and imputation can approximate whole genome sequencing (WGS) for biallelic single nucleotide variants (SNVs) by reference panel, genotype array, and ancestry**. Threshold is the smallest MAF for which >90% of biallelic SNVs are well-imputed (observed imputation  $r^2 > 0.8$ ).



**Table S5. Mean heterozygous concordance rates by reference panel, genotype array, ancestry, and MAF category.** Summary statistics are further broken down for the African ancestry study by estimated proportion of African ancestry (0.25-0.5, 0.5-0.75, 0.75-1.00) and for the Hispanic/Latino ancestry study by Caribbean (C) and non-Caribbean (NC) origin.



**Table S6. 25th, 50th, and 75th percentiles of the number of consecutive well-imputed**  (observed imputation r<sup>2</sup>>0.8) biallelic single nucleotide variants (SNVs) by reference **panel, genotype array, ancestry, and minor allele frequency (MAF) category.**

	Array	<b>MAF</b>	Length in kb of consecutively well-imputed (r2>0.8) biallelic SNVs											
			African			Hispanic/ Latino			European			Finnish		
			25 <sup>th</sup>	50 <sup>th</sup>	75 <sup>th</sup>	25 <sup>th</sup>	50 <sup>th</sup>	75 <sup>th</sup>	25 <sup>th</sup>	50 <sup>th</sup>	75 <sup>th</sup>	25 <sup>th</sup>	50 <sup>th</sup>	75 <sup>th</sup>
<b>TOPMed</b>	Omni 2.5M	Common	10.4	84.8	253.2	15.7	109.6	315.3	11.6	80.1	256.5	12.5	87.2	267.4
		Low frequency	2.3	23.9	84.3	5.2	20.7	60.1	1.6	7.8	29.8	2.3	8.0	19.2
		Rare	0.0	0.0	0.1	0.0	0.0	0.2	0.0	0.0	0.1	0.0	0.0	0.2
	MEGA	Common	5.1	73.3	241.6	11.4	101.3	302.2	5.3	55.7	210.5	6.3	64.9	224.4
		Low frequency	1.0	12.4	59.8	4.0	17.3	53.5	1.1	5.7	21.4	1.9	7.1	17.4
		Rare	0.0	0.0	0.1	0.0	0.0	0.2	0.0	0.0	0.1	0.0	0.0	0.2
	Express Omni	Common	0.7	31.3	162.8	4.6	70.4	245.3	2.5	23.9	113.6	3.2	31.1	144.6
		Low frequency	0.3	3.8	27.9	1.7	9.2	29.7	0.5	3.4	11.2	1.0	4.6	12.0
		Rare	0.0	0.0	0.1	0.0	0.0	0.1	0.0	0.0	0.1	0.0	0.0	0.2
	Core	Common	0.0	0.6	5.9	0.3	6.2	73.1	0.3	4.1	21.2	0.4	5.0	28.1
		Low frequency	0.0	0.7	3.4	0.3	2.1	8.9	0.0	1.2	4.5	0.1	1.9	6.0
		Rare	0.0	0.0	0.1	0.0	0.0	0.1	0.0	0.0	0.1	0.0	0.0	0.1
HRC	Omni 2.5M	Common	0.2	2.3	8.5	0.6	4.0	12.3	0.8	5.0	14.9	0.8	5.3	16.2
		Low frequency	0.0	0.3	1.0	0.0	0.6	1.9	0.1	1.7	5.2	1.4	5.6	13.8
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1
	MEGA	Common	0.0	0.9	4.3	0.3	2.9	10.1	0.7	4.7	14.2	0.8	5.3	16.2
		Low frequency	0.0	0.1	0.7	0.0	0.3	1.2	0.0	1.4	4.4	1.3	5.5	13.5
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1
	Express Omni	Common	0.0	0.7	3.4	0.2	2.2	8.2	0.7	4.2	12.4	0.8	5.3	16.1
		Low frequency	0.0	0.0	0.5	0.0	0.2	1.0	0.0	0.8	2.9	1.1	5.1	12.7
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1
	Core	Common	0.0	0.2	1.1	0.0	0.8	3.6	0.3	2.4	8.1	0.8	5.2	15.9
		Low frequency	0.0	0.0	0.2	0.0	0.0	0.5	0.0	0.3	1.7	0.8	4.0	10.6
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
1000G	Omni 2.5M	Common	0.3	4.0	21.2	1.2	9.7	33.4	1.9	10.6	31.6	2.1	12.9	40.0
		Low frequency	0.0	0.6	2.0	0.1	0.9	2.6	0.0	0.7	2.5	0.0	1.2	3.7
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
	MEGA	Common	0.0	1.0	6.0	0.4	4.2	20.4	0.9	6.7	23.2	1.5	9.9	33.1
		Low frequency	0.0	0.3	1.1	0.0	0.5	1.7	0.0	0.5	2.2	0.0	1.1	3.4
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
	Express Omni	Common	0.0	0.7	4.0	0.3	3.0	14.3	0.9	5.4	16.5	1.3	8.1	25.0
		Low frequency	0.0	0.2	0.9	0.0	0.3	1.3	0.0	0.2	$1.5$	0.0	0.8	2.7
		Rare	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
	Core	Common	0.0	0.2	1.3	0.0	0.8	4.2	0.1	1.9	7.6	0.4	3.4	12.5
		Low frequency	0.0	0.0	0.4	0.0	0.0	0.6	0.0	0.0	$1.0\,$	0.0	0.3	1.9
		Rare	0.0	$0.0\,$	0.0	0.0	$0.0\,$	0.0	$0.0\,$	0.0	$0.0\,$	0.0	0.0	0.0

**Table S7. 25th, 50th, and 75th percentiles of the length in kilobases (kb) of consecutively well-imputed (observed imputationr 2 >0.8) variants by reference panel, genotype array, ancestry, and minor allele frequency (MAF) category.**

**Table S8. Associations of genomic features with dichotomous TOPMed imputation quality status from logistic regressions by genotype array and ancestry.** MAF refers to the model with 9 MAF categories as predictors. ALL refers to the model with 9 MAF categories and all 6 genomic features as predictors. All other models refer to a model with 9 MAF categories and the named genomic feature as predictors. The features include mean GC content (GC), number of repeats (REP), number of structural variants (SV), presence of segmental duplications (SEG), mean recombination rate (RECOMB), and distance to nearest genotyped marker (DIST).

**Table S9. Associations of repeats with dichotomous TOPMed imputation quality status from logistic regressions by genotype array and ancestry.** All model names refer to the model with 9 MAF categories and membership in the named repeat class as predictors. Repeat classes as defined by RepeatMasker include DNA repeat elements (DNA), long interspersed repeated elements (LINE), low complexity repeats (LowComplex), long terminal repeat elements including retrotransposons (LTR), rolling circle repeats (RC), RNA repeats (RNA), satellite repeats, microsatellites (Simple), short interspersed repeat elements including ALUs (SINE), and repeats of unknown class.

**Table S10. Associations of genomic features with continuous TOPMed imputation quality status from zero-one inflated beta regressions by genotype array and ancestry.** All models used the same set of predictors for each of the four parameters (see methods). MAF refers to the model with 9 MAF categories as predictors. ALL refers to the model with 9 MAF categories and all 6 genomic features as predictors. All other models refer to a model with 9 MAF categories and the named genomic feature as predictors. The features include mean GC content (GC), number of repeats (REP), number of structural variants (SV), presence of segmental duplications (SEG), mean recombination rate (RECOMB), and distance to nearest genotyped marker (DIST).



**Table S11. Proportion of biallelic single nucleotide variants (SNVs) in each whole genome sequencing (WGS) study that are well-imputed (r 2 >0.8) by reference panel, genotype array, ancestry, and predicted impact on protein coding.** Predicted impact was estimated with VEP.



**Table S12. Minor allele frequency (MAF) threshold above which array genotyping and imputation can approximate whole genome sequencing (WGS) with the TOPMed panel by genotype array, ancestry, and variant type**. Threshold is the smallest MAF for which >90% of variants are well-imputed (observed imputation  $r^2$ >0.8).

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