**Reliability of imputed genotype data for family-based analyses**

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**Introduction**

Family-based analyses use random genetic variation within families to remove confounding from population stratification and to separate out direct genetic effects (effects of variants in an individual on that individual) from indirect genetic effects (effects of variants in an individual on another individual mediated through the environment). The underlying principle relies on relationships between relatives’ genotypes due to random segregations of genetic material during meiosis1–5. Family-based genome-wide association studies (FGWASs) — which use parental genotypes as control variables — and sib-GWAS6 — which use sibling genotypes as control variables — have been proposed as solutions to the confounding issues known to affect standard GWAS designs and downstream applications including estimation of heritability and genetic correlation, Mendelian randomization, and inferences of natural selection.

While family-based analyses have favorable theoretical properties, issues arising from imperfections in real-world genotype data have not been explored. Most GWAS studies use data derived from genotyping arrays — which measure genotypes at an incomplete set of variants — followed by imputation from a panel of reference haplotypes with more complete genome sequences. The imputation methods work by finding related sequences in the reference panel and using the reference haplotypes to fill in the genotypes at the positions not directly observed on the genotyping array, but they do not take into account relationships between individuals in the target sample, including the sibling and parent-offspring relations used in family-based analyses. The output of imputation methods typically consists of a probability distribution over genotypes, from which the most likely genotype (‘hard-call’) or the expected genotype (‘dosage’) is used for downstream analyses. Imputation methods also provide a quality metric (INFO score or R2) that estimates the fraction of genotype variation the imputation has recovered and is thus bounded in [0,1].

**Results**

**Correlations between relatives’ imputed genotypes**

We sought to examine whether imputed data preserves the relationships between relatives’ genotypes implies by Mendelian Laws and from which the theoretical properties of family-based analyses derive. We analyzed correlations between relatives’ imputed genotypes (UKBv3 imputation) for 19,290 sibling pairs and 5,324 parent-offspring pairs from the UK Biobank White British subsample. We found that correlations between first-degree relatives’ imputed genotypes decreased below the theoretical expectation with imputation quality (measured by INFO score with values between 0 and 1): for imputed SNPs (MAF>1%) with INFO scores between 0.30–0.31, the mean correlation between siblings was 0.437 (S.E.=0.001) for genotype dosages and 0.376 (S.E.=0.001) for hard-calls, lower than the expected correlation of 0.5. Even at INFO scores of 0.96–0.97, the correlation remained below 0.5 (P=1.6x10⁻⁴) for hard-call genotypes. For parent-offspring pairs, SNPs (MAF>1%) with INFO scores between 0.30-0.31, the mean correlation between parent-offspring pairs was 0.434 (S.E.=0.001) for genotype dosages and 0.337 (S.E.=0.002) for hard-calls lower than the expected correlation of 0.5. Like sibling pairs even at INFO scores of 0.96-0.97 the correlation remained below 0.5 (P=2.59x10⁻2) (Figure 1).

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**Figure 1. Correlations between relatives’ genotypes as a function of imputation quality (INFO score).** We show the mean correlation from 1000 SNPs in each INFO score bin along with its 95% confidence interval for both imputed dosages (expected genotype given genotype probabilities) and imputed hard-call genotypes (most likely genotype). Results from 5,324 parent-offspring pairs and 19,290 sibling pairs from the white British subsample of the UK Biobank.

**Assessing bias due to imputation in sib-GWAS**

We now show how the bias in sib-GWAS induced by imputation can be assessed by comparing sibling imputed genotypes and genotypes from whole genome sequence data. For a sibling pair from family , the model is:

We assume that sibling genotypes are mean-normalized. The sib-difference regression can proceed by least-squares regression of onto . This will give an unbiased estimate of . We can rewrite the equation in terms of the regressions onto and (which are uncorrelated with each other):

for some uncorrelated with and . We can then ask what we would obtain if we instead performed regression onto the difference in imputed sibling genotypes:

We can thus assess the bias that comes from using imputed sibling genotypes by performing two regressions:

1. Slope of regression of on , which gives bias coming from the sib-difference component;
2. Slope of regression of on , which gives bias coming from the sib-sum component, i.e. the component that the sib-difference should be orthogonal with as it captures parental effects/stratification.

This can be done by obtaining the sibling genotypes from whole genome sequence (WGS) data (which we assume to be true) and forming and and regressing these values onto the difference in sib imputed genotypes, .

From these slopes, we could express the expected result of sib-difference regression as a linear combination of and .

**Comparison of sibling imputed and WGS genotypes.**

We regressed the difference between siblings’ genotypes, derived from whole-genome sequencing (WGS) data in the UKB, onto the difference from imputed data:

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Theoretically, we would expect an intercept of zero and a slope of one if the imputed genotypes are the same as the WGS genotypes.

We analyzed 68 high-quality imputed SNPs (mean info score of 0.966) and 46 low-quality imputed SNPs (mean info score of 0.312). For high-quality SNPs, the average slope estimate was 0.957 (S.E. = 0.0015), whereas for low-quality SNPs the average slope was 0.062 (S.E. = 0.0142), suggesting a much weaker relationship between imputed and WGS genotypes than would expected given the INFO score.

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**Figure 2.** **The distribution of intercepts and slopes from the regression of  onto  for two groups of SNPs**: high-quality and low-quality imputed SNPs. The analysis includes 68 high-quality SNPs and 46 low-quality SNPs, with models fitted for 19,052 White British sibling pairs from the UK Biobank.

**INFO score is an unreliable metric of imputation quality in UK Biobank**

The above results showed much poorer concordance between imputed and WGS genotypes than we expected based on INFO score. Thus, we performed a simpler analysis that looked at the regression R2 between imputed and WGS genotypes as a function of SNP INFO score using 19,052 White British individuals, one from each sibling pair.

Figure 3 presents the relationship between the INFO score and the R2 between imputed and WGS data. The results indicate that the INFO score is highly unreliable — rather than matching the R2, as expected, the actual R2 values are often much lower. For high-quality SNPs, some have R2 values as low as 0.5, while for low-quality SNPs (with INFO scores around 0.3), the imputed and WGS genotypes are mostly uncorrelated.

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**Figure 3. INFO score vs R2 between imputed and WGS data in UK Biobank.**

**Discussion**

These results raise concerns about the sib-GWAS meta-analysis by Howe et al.6 that used hard-call genotypes for all SNPs with MAF>1% and INFO score greater than 0.3, implying many low-quality imputed genotypes were used. To investigate the impact that low-quality imputed genotypes have on sib-GWAS and FGWAS, we are going to perform more analyses comparing the imputed genotypes in UK Biobank to the whole-genome sequencing data. For the next step we are going to extend the analysis to a greater number of SNPs across different INFO score values. We will then perform sib-GWAS and FGWAS using WGS data and compare the results from downstream analyses to those from imputed data.

Our results argue that stringent quality control should be applied for family-based analyses using imputed genotype data to ensure that such analyses have the theoretical properties that they promise. Furthermore, imputation methods that account for pedigree relations between close relatives should be developed to enable better family-based analyses of data derived from genotyping arrays.

Beyond implications for family-based analyses, our results raise more general questions about the reliability of imputed data, currently the vast majority of data that has been analysed in GWAS. An important question is whether the unreliability of the imputed data in the UK Biobank applies to other imputed datasets. Even if it does not, since the UK Biobank imputed genotype data7 is probably the most used genetic data in the world, the issues with this particular imputed dataset deserve attention.

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