

# Efficient Estimation of Realized Kinship from Single Nucleotide Polymorphism Genotypes

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**ABSTRACT** Realized kinship is a key statistic in analyses of genetic data involving relatedness of individuals or structure of populations. There are several estimators of kinship that make use of dense SNP genotypes. We introduce a class of estimators, of which some existing estimators are special cases. Within this class, we derive properties of the estimators and determine an optimal estimator. Additionally, we introduce an alternative marker weighting that takes allelic associations [linkage disequilibrium (LD)] into account, and apply this weighting to several estimators. In a simulation study, we show that improved estimators are obtained (1) by optimal weighting of markers, (2) by taking physical contiguity of genome into account, and (3) by weighting on the basis of LD.

**KEYWORDS** realized kinship; genomic relationship matrix (GRM); local identity by descent (IBD); linkage disequilibrium; locus weighting

**G**ENES inherited from the same ancestral copy by related individuals are said to be identical by descent (IBD). At the locus level, a pair of individuals share 0–4 genes that are IBD. At the genome level, the kinship coefficient is often used to summarize the average amount of IBD sharing across all loci. The kinship coefficient is important in genetic data analyses that either use or adjust for pairwise relatedness. The *pedigree kinship*,  $\Psi$ , the probability that genes segregating from each individual at a randomly chosen locus are IBD, is a deterministic function of the pedigree relationship. However, the *realized kinship*,  $\Phi$ , the actual proportion of IBD genome between two individuals varies widely about its expectation ( $\Psi$ ) as a consequence of Mendelian sampling (Hill and Weir 2011). Additionally, in samples ascertained on the basis of trait information or samples affected by artificial selection, there are biases in the levels of relatedness of individuals (Liu *et al.* 2003; Purcell *et al.* 2007). Thus, even when pedigree information is available, estimates of realized kinship are often preferred to pedigree kinship (Visscher *et al.* 2006; VanRaden 2007, 2008; Hayes *et al.* 2009).

The availability of dense SNP genotypes makes it possible to estimate realized kinship accurately without pedigree in-

formation. Such estimates are particularly useful in studies that involve population samples (Choi *et al.* 2009; Yang *et al.* 2010; Day-Williams *et al.* 2011). Improving the precision of realized-kinship estimators has value for human genetics applications involving gene mapping or genotypic disease risk prediction, and also for animal and plant breeding, where incremental improvements in predictive accuracy are of economic value in trait optimization.

The matrix of inferred pairwise realized kinship is a measure of genomic similarity and a kernel in the sense of Gianola and van Kaam (2008). It is therefore appropriate for phenotype prediction or whole genome prediction (de los Campos *et al.* 2010). The use of this matrix for phenotype or breeding-value prediction is the genomic best linear unbiased prediction methodology (for example, Bernardo 2008). In plant and animal breeding applications, pedigree information has been combined with inferred kinships (Crossa *et al.* 2010). In the mixed model context, the matrix is used for variance component estimation (Yang *et al.* 2010), or as a proxy for polygenic effects in the presence of major gene effects (Kang *et al.* 2010).

One group of existing estimators of realized kinship makes use only of the population allele frequencies at each SNP marker, and not of additional information such as the ordering of markers along the chromosome. Estimators in this group require only dense SNP genotypes and reliable sources of marker allele frequencies as input. The PLINK (Purcell *et al.* 2007) method-of-moments estimator ( $\hat{\Phi}_P$ ) estimates realized kinship from the  $k$  coefficients; the proportion of genome at

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which two noninbred individuals share 0, 1, or 2 IBD genes. Choi *et al.* (2009) adopted a maximum likelihood estimator (MLE) ( $\hat{\Phi}_M$ ) that estimates the  $k$  coefficients using an EM algorithm. The classic genomic relationship matrix (GRM) estimator ( $\hat{\Phi}_G$ ) estimates kinship through the empirical correlation of genotypes; see, for example Hayes *et al.* (2009). An alternative version of the GRM estimator ( $\hat{\Phi}_R$ ) is more robust to presence of rare alleles (VanRaden 2008). Day-Williams *et al.* (2011) proposed a method-of-moments estimator ( $\hat{\Phi}_D$ ), which estimates kinship by exploring the relationship between identity by state (IBS) and IBD.

Other estimators make use of various sources of additional information to improve accuracy of kinship estimation. A number of methods have been developed to estimate IBD sharing at the locus level; see for example Moltke *et al.* (2011) and methods cited in Brown *et al.* (2012). These location-specific IBD estimates in turn provide estimates of kinship; note that location-specific kinships are constrained to the values 0, 1/4, 1/2, and 1 (Day-Williams *et al.* 2011). Here, we consider estimators from two of such local IBD methods that have not been previously used to estimate genome-wide realized kinship. The local method of Day-Williams *et al.* (2011) with resulting kinship estimator  $\hat{\Phi}_L$ , (to be distinguished from  $\hat{\Phi}_D$ ) requires information on the ordering of markers along the chromosomes. It predicts the amount of sharing at each marker by first estimating a neighborhood kinship for each marker and then applying a constrained smoothing algorithm on each chromosome. The hidden Markov model (HMM) proposed by Brown *et al.* (2012), with resulting kinship estimator  $\hat{\Phi}_H$ , estimates probabilities of IBD states between two or more individuals at each marker location. A genetic map is needed to compute transition probabilities along the Markov chain.

Finally, we consider efficient estimation of realized kinship in the presence of linkage disequilibrium (LD). An increase in the density of SNP panels, by itself, will not improve precision without limit in the presence of LD, as additional SNPs are not independent sources of information. Speed *et al.* (2012) developed LDAK ( $\hat{\Phi}_K$ ), a weighted version of the GRM that takes LD into account. By analogy with methods introduced in the population structure context by Patterson *et al.* (2006) and Zou *et al.* (2010), LDAK equalizes contributions of linked SNPs by downweighting SNPs which make redundant contribution to the GRM as evidenced by off-diagonal terms in the (squared) SNP correlation matrix. We derive an analogous weighting by optimizing the weighted GRM estimator variance under tractable assumptions.

This article focuses on pedigree-free estimation of realized kinship in a homogenous population. In *Methods*, we first introduce the framework of a general class of GRM estimators, of which the classic GRM estimator ( $\hat{\Phi}_G$ ), the robust GRM estimator ( $\hat{\Phi}_R$ ), and the global Day-Williams estimator ( $\hat{\Phi}_D$ ) are all special cases. Under the assumptions of linkage equilibrium and absence of inbreeding, we propose a two-step GRM estimator ( $\hat{\Phi}_T$ ) that approximates

the minimum variance estimator within this class. In the general case of LD, we derive the variance for a weighted GRM estimator, and construct an estimator ( $\hat{\Phi}_W$ ) with certain optimality properties given the LD structure of the population. We next describe the implementation of our simulation study which compares performance of the different kinship estimators detailed above. Additionally, the optimal weights derived for  $\hat{\Phi}_W$  may also be applied to other estimators; specifically, we consider reweighted versions of  $\hat{\Phi}_L$  and  $\hat{\Phi}_H$ . Results of the simulation study are presented in *Results*. We show that the proposed estimators are very competitive against existing estimators. We conclude with a *Discussion*.

## Methods

### A general class of GRM estimators

At any autosomal locus, there are 15 possible IBD states among the four genes of two individuals. These 15 IBD states fall into nine genotypically distinct classes (see, for example, Thompson 2000). When interest lies only in the amount of sharing between (as opposed to within) the individuals, the nine classes of states condense further into four groups of states characterized by local kinship,  $\phi$ , which takes values in  $\{0, 1/4, 1/2, 1\}$ . Global realized kinship,  $\Phi$ , measures the IBD proportion shared between the individuals across the genome, and so takes values in the range  $[0, 1]$ . Since IBD results from the meiotic process, we measure this proportion in terms of genetic distance. However, since recombination is a coarse process relative to dense SNP markers, global kinship is well approximated by the average of local kinship over a large number of marker loci approximately evenly spaced in genetic distance throughout the genome.

We consider a general class of GRM estimators of the following form:

$$\hat{\Phi}(\mathbf{a}, \mathbf{w}) = \sum_{l=1}^L w_l \times \frac{x_l y_l - a_l(x_l + y_l) + 4a_l p_l - 4p_l^2}{4p_l(1 - p_l)}, \quad (1)$$

where  $L$  is the total number of marker loci,  $\mathbf{x} = (x_1, \dots, x_L)^T$  and  $\mathbf{y} = (y_1, \dots, y_L)^T$  are counts of reference alleles at each marker for the pair of individuals,  $\mathbf{p} = (p_1, \dots, p_L)^T$  are the population frequencies of the reference alleles,  $\mathbf{a} = (a_1, \dots, a_L)^T$  are multiplicative factors, and  $\mathbf{w} = (w_1, \dots, w_L)^T$  are non-negative weights satisfying  $\sum_{l=1}^L w_l = 1$ . The vectors  $\mathbf{a}$  and  $\mathbf{w}$  are parameters that distinguish different GRM estimators, while  $\mathbf{x}$  and  $\mathbf{y}$  are the data random variables and  $\mathbf{p}$  is assumed known. Note that the denominator in (1) is  $4p_l(1 - p_l)$  as opposed to  $2p_l(1 - p_l)$ , reflecting the difference between kinship coefficients and the relatedness coefficients of the numerator relationship matrix (Henderson 1976).

The classic GRM estimator is a special case of (1) with  $a_l = 2p_l$  and  $w_l = 1/L$  for all  $l$ :

$$\begin{aligned}\hat{\Phi}_G &= \frac{1}{L} \sum_{l=1}^L \frac{(x_l - 2p_l)(y_l - 2p_l)}{4p_l(1-p_l)} \\ &= \sum_{l=1}^L \frac{1}{L} \times \frac{x_l y_l - 2p_l(x_l + y_l) + 4p_l^2}{4p_l(1-p_l)}.\end{aligned}\quad (2)$$

The robust GRM estimator is a special case with  $a_l = 2p_l$  and  $w_l = 4p_l(1-p_l)/[\sum_{m=1}^L 4p_m(1-p_m)]$  for all  $l$ :

$$\begin{aligned}\hat{\Phi}_R &= \frac{\sum_{l=1}^L (x_l - 2p_l)(y_l - 2p_l)}{\sum_{l=1}^L 4p_l(1-p_l)} \\ &= \sum_{l=1}^L \frac{4p_l(1-p_l)}{\sum_{m=1}^L 4p_m(1-p_m)} \times \frac{x_l y_l - 2p_l(x_l + y_l) + 4p_l^2}{4p_l(1-p_l)}.\end{aligned}\quad (3)$$

The global Day-Williams estimator can be reparametrized (see Appendix A) into the form of (1) with  $a_l = 1$  and  $w_l = 4p_l(1-p_l)/[\sum_{m=1}^L 4p_m(1-p_m)]$  for all  $l$ :

$$\begin{aligned}\hat{\Phi}_D &= \frac{\sum_{l=1}^L x_l y_l - (x_l + y_l) + 4p_l - 4p_l^2}{\sum_{l=1}^L 4p_l(1-p_l)} \\ &= \sum_{l=1}^L \frac{4p_l(1-p_l)}{\sum_{m=1}^L 4p_m(1-p_m)} \times \frac{x_l y_l - (x_l + y_l) + 4p_l - 4p_l^2}{4p_l(1-p_l)}.\end{aligned}\quad (4)$$

In the general form of (1), write

$$Z_l(a_l) = \frac{x_l y_l - a_l(x_l + y_l) + 4a_l p_l - 4p_l^2}{4p_l(1-p_l)}$$

so that  $\hat{\Phi}(\mathbf{a}, \mathbf{w}) = \sum_{l=1}^L w_l \times Z_l(a_l)$ .

Note that  $\mathbf{Z}(\mathbf{a}) = [Z_1(a_1), \dots, Z_L(a_L)]^T$  depends on the parameters  $\mathbf{a}$  but not on  $\mathbf{w}$ . We make two basic assumptions throughout the article. First, we assume IBD genes have the same allelic types and non-IBD genes have independent allelic types. In addition, we assume exchangeability of parental lineage, so that either of the two genes from the first individual is equally likely to be IBD to either of the two genes of the second individual at the same locus. Under these assumptions, it can be shown that  $\mathbb{E}[Z_l(a_l)] = \Phi$  regardless the choice of  $a_l$  (see Appendix B, *General Case*). Thus,  $\mathbb{E}[\hat{\Phi}(\mathbf{a}, \mathbf{w})] = \Phi$  for any  $\mathbf{a}$  and  $\mathbf{w}$ .

Performance of unbiased estimators depend on their variances. For a GRM estimator in the general form of (1),

$$\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})] = \sum_{l=1}^L w_l^2 V_l(a_l) + \sum_{l \neq m} w_l w_m \text{Cov}[Z_l(a_l), Z_m(a_m)],\quad (5)$$

where  $V_l(a_l) = \text{Var}[Z_l(a_l)]$ . Computation of  $\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})]$  is intractable without simplifying assumptions. We next derive the  $\mathbf{a}$  and  $\mathbf{w}$  that minimize  $\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})]$  under different assumptions.

### Linkage equilibrium

We first assume linkage equilibrium. Although in reality there is LD, empirical results show that the relative values of variances of estimators are well approximated by those derived under this assumption (see *Discussion*). When markers are in linkage equilibrium, the allelic types at different markers on the same haplotype are independent. Equation 5 then reduces to

$$\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})] = \sum_{l=1}^L w_l^2 V_l(a_l).\quad (6)$$

To find the GRM estimator with minimal variance, Equation 6 suggests that one should first (for each  $l$ ) choose  $a_l$  that minimizes  $V_l(a_l)$ , and then choose  $\mathbf{w}$  to minimize  $\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})]$ .

We now make an additional assumption of no inbreeding. For general choices of  $a_l$ , it can be shown (see Appendix B, *Linkage Equilibrium*) that

$$\begin{aligned}V_l(a_l) &= \frac{1}{4p_l(1-p_l)} \times \left[ (a_l - 2p_l)^2 + 2\Phi(a_l - 1)^2 - \Phi \right. \\ &\quad \left. + 4\Phi(1 - \Phi)p_l(1-p_l) + (k_2 + 1)p_l(1-p_l) \right],\end{aligned}\quad (7)$$

where  $\Phi$  is the realized kinship and  $k_2$  is the realized proportion of the genome that the pair of individuals share both genes IBD. Note that the value of  $V_l(a_l)$  is asymmetric about  $p_l = 0.5$  for general choices of  $a_l$ , and thus is sensitive to the choice of reference allele. However, when  $a_l$  is any weighted average of  $2p_l$  and 1 [as in (8)],  $V_l(a_l)$  becomes invariant to the choice of reference allele. With such an  $a_l$ ,  $V_l(a_l)$  attains its minimum at  $p_l = 0.5$ , conditional on  $\Phi$  and  $k_2$  (see Appendix B, *Linkage Equilibrium*).

It follows from (7) that the optimal multiplicative factors that minimize the unweighted single-marker variances have the form

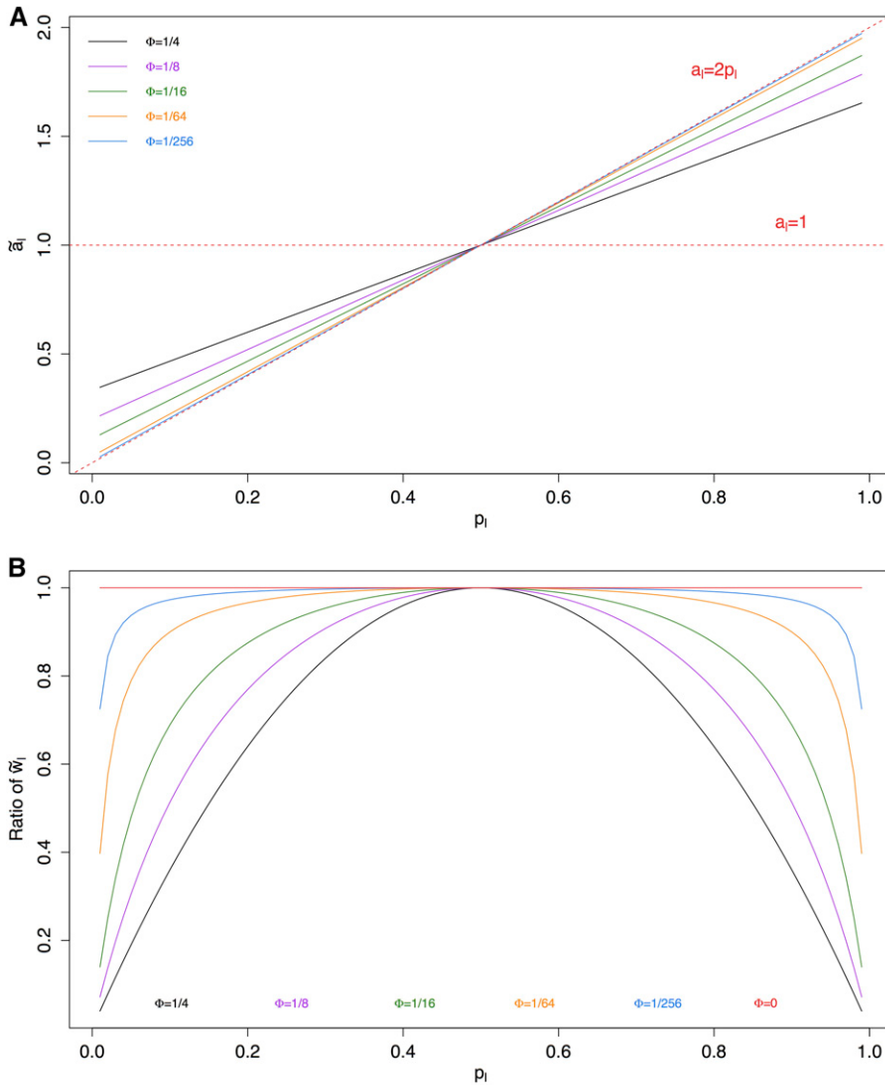
$$\tilde{a}_l = \arg \min_{a_l} V_l(a_l) = \frac{1}{1 + 2\Phi} \times 2p_l + \frac{2\Phi}{1 + 2\Phi}.\quad (8)$$

Interestingly,  $\tilde{a}_l$  is a weighted average of  $2p_l$  and 1, which are the choices used by  $\hat{\Phi}_G$ ,  $\hat{\Phi}_R$ , and  $\hat{\Phi}_D$ , respectively (compare Equations 2–4). Figure 1A shows how  $\tilde{a}_l$  varies with  $p_l$  for different  $\Phi$ 's. We see that  $a_l = 2p_l$  is optimal when  $\Phi = 0$ , whereas  $a_l = 1$  is far from optimal even when  $\Phi = (1/4)$ . Since  $\hat{\Phi}_R$  and  $\hat{\Phi}_D$  use the same weights,  $\hat{\Phi}_R$  is more efficient than  $\hat{\Phi}_D$  for  $\Phi < 1/2$  (from Equation (7)) under the assumptions of this section.

Conditional on  $\mathbf{a}$ , the optimal weights that minimize  $\text{Var}[\hat{\Phi}(\mathbf{a}, \mathbf{w})]$  have the form

$$\tilde{w}_l(\mathbf{a}) = \frac{V_l(a_l)^{-1}}{\sum_{m=1}^L V_m(a_m)^{-1}}, \quad l = 1, \dots, L.\quad (9)$$

The weights in (9) can be equivalently specified by the ratios



**Figure 1** (A)  $\tilde{a}_i$  as a function of  $p_i$  for different  $\Phi$ 's, with  $a_i = 2p_i$  and  $a_i = 1$  as reference lines. (B) Ratio of weights between a marker with allele frequency  $p_i$  and a reference marker with allele frequency 0.5, under the optimal weighting scheme  $\tilde{\mathbf{w}}(2\mathbf{p})$  for different combinations of  $(\Phi, k_2)$ . In calculation of  $\tilde{\mathbf{w}}(2\mathbf{p})$ , we used  $k_2 = (1/4)$  when  $\Phi = (1/4)$  and 0 otherwise. Note that these combinations of  $(\Phi, k_2)$  correspond to pedigree expectations of a pair of individuals that are full siblings, half siblings, first cousins, second cousins, or third cousins.

$$\frac{\tilde{w}_l(\mathbf{a})}{\tilde{w}_m(\mathbf{a})} = \frac{V_m(a_m)}{V_l(a_l)}, \quad l, m = 1, \dots, L,$$

which are functions of allele frequencies at the corresponding pairs of markers, conditional on  $\Phi, k_2$  and the choice of  $\mathbf{a}$ . For  $\mathbf{a} = 2\mathbf{p}$ , Figure 1B shows how a marker with frequency  $p_l$  is weighted relative to a marker with frequency 0.5 under the optimal weighting scheme  $\tilde{\mathbf{w}}(2\mathbf{p})$  for different combinations of  $(\Phi, k_2)$ . The optimal solution weights markers very differently, especially for large  $\Phi$ . In this case ( $\mathbf{a} = 2\mathbf{p}$ ), the uniform weighting of  $\hat{\Phi}_G$  is optimal when  $\Phi = 0$ , whereas the weighting of  $\hat{\Phi}_R$  is optimal when  $\Phi = (1/4)$  and  $k_2 = (1/4)$ .

The estimator  $\hat{\Phi}[\tilde{\mathbf{a}}, \tilde{\mathbf{w}}(\tilde{\mathbf{a}})]$  would be an obvious choice if we knew  $\tilde{\mathbf{a}}$  and  $\tilde{\mathbf{w}}(\tilde{\mathbf{a}})$ . However, elements of  $\tilde{\mathbf{a}} = (\tilde{a}_1, \dots, \tilde{a}_L)^T$  are functions of the unknown  $\Phi$ , and elements of  $\tilde{\mathbf{w}}(\tilde{\mathbf{a}}) = [\tilde{w}_1(\tilde{\mathbf{a}}), \dots, \tilde{w}_L(\tilde{\mathbf{a}})]^T$  are functions of the unknown  $\Phi, k_2$ , and  $\tilde{\mathbf{a}}$ . The closed form expressions given in Equations 7–9 motivate a two-step estimator,  $\hat{\Phi}_T$ , which approximates  $\hat{\Phi}[\tilde{\mathbf{a}}, \tilde{\mathbf{w}}(\tilde{\mathbf{a}})]$  following these two steps:

1. Obtain initial estimates of  $\Phi$  and  $k_2$  using existing methods.
2. Compute  $\tilde{\mathbf{a}}^*$  (Equation 8) and then  $\tilde{\mathbf{w}}^*(\tilde{\mathbf{a}}^*)$  (Equations 7 and 9) using these estimates of  $\Phi$  and  $k_2$ .

Then  $\hat{\Phi}_T = \hat{\Phi}[\tilde{\mathbf{a}}^*, \tilde{\mathbf{w}}^*(\tilde{\mathbf{a}}^*)]$ .

In practice, existing kinship estimators such as  $\hat{\Phi}_G, \hat{\Phi}_R$ , and  $\hat{\Phi}_D$  may produce negative estimates of  $\Phi$  in Step 1. Our implementation uses  $\hat{\Phi}_G$  to obtain an initial estimate of  $\Phi$ . When this initial estimate is negative, we simply retain it as the  $\hat{\Phi}_T$  estimate. For simplicity, we set  $k_2 = 0$  when computing  $V_l(\tilde{a}_l^*)$  for all  $l$ . In principle, this affects the calculation of  $\tilde{\mathbf{w}}^*(\tilde{\mathbf{a}}^*)$  for bilateral relatives, but it makes no practical difference (see Supplemental Material, File S1, section A).

### Linkage disequilibrium

We now drop the assumptions of linkage equilibrium and absence of inbreeding. To calculate variances, we follow the approach of Sverdlov (2014), and consider the case where the pair of individuals are unrelated. Under these

assumptions, it can be shown (see Appendix B, *Linkage Disequilibrium*) that

$$\tilde{a}_l = \arg \min_{a_l} V_l(a_l) = 2p_l.$$

However, there generally does not exist an  $\mathbf{a}$  that jointly minimizes each of the unweighted covariance terms in Equation 5. Thus, we conveniently set  $\mathbf{a} = 2\mathbf{p}$  in the remainder of this section.

Let  $F_x$  and  $F_y$  denote the inbreeding coefficients of the two individuals respectively. We have

$$V_l(2p_l) = \text{Var}[Z_l(2p_l)] = \frac{1}{4}(F_x + 1)(F_y + 1) \quad (10)$$

and

$$\text{Cov}[Z_l(2p_l), Z_m(2p_m)] = \frac{1}{4}(F_x + 1)(F_y + 1)\rho_{lm}^2, \quad (11)$$

where  $\rho_{lm}$  is the genotype dosage correlation between locus  $l$  and locus  $m$ . Conveniently,  $\rho_{lm}$  only enters the expression in a squared form, so the covariance is invariant to the choice of reference allele. Combining (10) and (11), Equation 5 becomes

$$\text{Var}[\hat{\Phi}(2\mathbf{p}, \mathbf{w})] = \sum_l \sum_m w_l w_m \times \frac{1}{4}(F_x + 1)(F_y + 1)\rho_{lm}^2, \quad (12)$$

where  $\rho_{lm} = 1$  when  $l = m$ . We assume that the matrix of squared LD correlations,  $\mathbf{R} = [\rho_{lm}^2]$ , is known. The goal is to find the  $\mathbf{w}$  that minimizes  $\text{Var}[\hat{\Phi}(2\mathbf{p}, \mathbf{w})]$ . Note that the inbreeding coefficients,  $F_x$  and  $F_y$ , are part of a fixed scaling factor.

The optimization problem reduces to

$$\min_{\mathbf{w}} [\mathbf{w}^T \mathbf{R} \mathbf{w} - \mathbf{w}^T \mathbf{1}] \quad : \quad w_l \geq 0 \quad \forall l. \quad (13)$$

Presence of the second term in the objective function forces a solution that satisfies  $\mathbf{w}^T \mathbf{1} = c$  for some  $c > 0$ , which can be rescaled by  $1/c$  to obtain the final solution  $\tilde{\mathbf{w}}$ . The LD weighted GRM estimator is then  $\hat{\Phi}_W = \hat{\Phi}(2\mathbf{p}, \tilde{\mathbf{w}})$ .

The matrix  $\mathbf{R}$  is positive semidefinite (see Appendix B, *Linkage Disequilibrium*). The above minimization problem can be solved for general  $\mathbf{R}$  using standard quadratic programming procedures, and closed-form solutions exist for special cases of  $\mathbf{R}$  (see File S1, section B). However, in practice, it will be necessary to divide the large set of genome-wide SNPs into blocks. In the case where  $\mathbf{R}$  has a block-diagonal structure, the optimization problem can be solved for each block. The final solutions will be a concatenation of the rescaled block solutions. Additional details are given in Appendix B, *Linkage Disequilibrium*.

### Methods of analysis

In the simulation study, we considered six relationship types: full siblings (FS), half siblings (HS), first cousins (C1), second

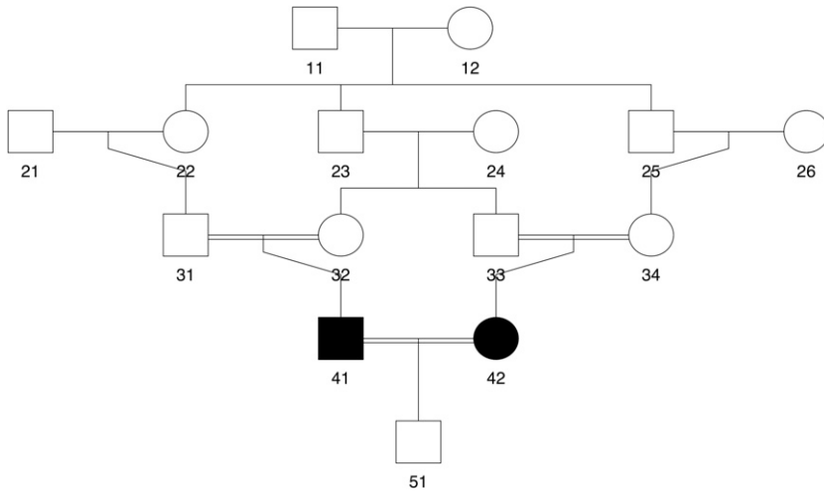
cousins (C2), third cousins (C3), and inbred cousins (IN) from the complex JV pedigree (Goddard *et al.* 1996) shown in Figure 2. Dense SNP genotypes were generated for 1000 independent pairs of each relationship type as follows:

1. Simulate recombination breakpoints and Mendelian sampling for all meioses in the smallest complete pedigree that contains the pair of relatives.
2. Assign founder genome labels (FGLs) (Sobel and Lange 1996) to founder haplotypes, and determine the inherited FGLs at all marker positions for all nonfounders with respect to the inheritance pattern simulated in step 1.
3. Sample founder haplotypes from a reference pool, and assign alleles to nonfounders with respect to both the sampled founder haplotypes and the inherited FGLs determined in step 2.
4. At each locus, combine the two alleles inherited by the related pair of individuals to create genotype data.

Data generation was implemented using the *ibd\_create* program of MORGAN version 3.3.1 (Thompson and Lewis 2016). Marker and haplotype information used in data generation were extracted from the 1000 Genomes Project Phase 3 data (1000 Genomes Project Consortium 2015). All 5008 phased haplotypes from the combined population were made available for sampling of founder haplotypes. This use of real haplotypes preserves the natural patterns of LD in the combined population. The locations of markers in Haldane cM were obtained from the Rutgers Map version 3a (Matise *et al.* 2007). A total of 169,751 markers were selected from the 22 autosomes based on spacing ( $\sim 50$  markers per cM), minor allele frequency ( $\geq 0.05$ ), and complete genotype information (no missing genotypes). The distribution of allele frequencies in the marker panel thus reflects real studies using common SNPs. The marker density was chosen to be dense enough to show patterns of LD, yet sparse enough to be attainable by older SNP arrays in human genetics and by modern SNP arrays in animal genetics.

All the kinship estimators were evaluated on the simulated data. We used PLINK version 1.07 to implement the PLINK estimator, *ibd\_haplo* program (Brown *et al.* 2012) of MORGAN version 3.3.1 to implement the HMM estimator, and our own code to implement all other estimators. Marker allele frequencies used in analysis were estimated by PLINK version 1.07 for each relationship type separately. For the MLE estimator, we adopted a very stringent convergence criterion (order of  $10^{-8}$  of the final log-likelihood) and used pedigree  $k$  coefficients as the starting configurations of the EM algorithm. The GRM estimators were computed both unconstrained to the range  $[0, 1]$  (consistent with the theory in *Methods*), and also constrained to this range.

The local Day-Williams method imputes local kinship directly, whereas the HMM method estimates probabilities of local IBD states. For ease of comparison, we used the most probable state from the HMM output to impute local kinship. For these two local methods, estimates of realized (global) kinships were calculated as the average of imputed local



**Figure 2** The JV pedigree. Individual 41 and 42 are the inbred quadruple cousins (IN) considered in the simulation study. Double lines indicate consanguineous marriages.

kinship across all marker loci. We simulated genotype data on an additional 300 independent pairs of cousins (100 first cousins, 100 second cousins, and 100 third cousins) for tuning purposes. For each of the two local IBD methods, a sparse grid search was implemented to find the set of tuning parameters that maximizes local kinship imputation rate over the tuning data set. The selected sets of tuning parameters were subsequently used in the actual analysis.

All LD-adjusted methods used the reference genotypes of all 2504 individuals from the 1000 Genomes Project Phase 3 data to obtain LD information. A naive LD-pruned GRM estimator ( $\hat{\Phi}_N$ ) is included as a baseline for comparison. For this estimator, LD pruning was done using PLINK version 1.07, where we sequentially threw out one of the pairs of markers that had a genotypic dosage correlation  $\rho^2 > 0.2$  (option `-r2` in PLINK). This pruning step reduced the marker set to about half of the original size, and the classic GRM was applied to the reduced marker set to obtain  $\hat{\Phi}_N$ . For the LD-weighted GRM estimator ( $\hat{\Phi}_W$ ), weights were computed in blocks of 2000 SNPs using the optimization package JuMP version 0.14.1 (Lubin and Dunning 2015) in the Julia language. Weights of the LDAK estimator ( $\hat{\Phi}_K$ ) were computed using LDAK version 4.9 with the default options.

Using a single processor on a standard desktop, the PLINK estimates or any of the GRM estimates can be computed for 1000 pairs of individuals with the selected genome-wide marker panel in a couple of minutes. The MLE estimates can take many hours to compute depending on run conditions (see *Results*). Either of the two local IBD estimates takes several hours to compute, but this is still computationally feasible. With reference genotypes from 2504 individuals as the basis for LD adjustment, LD pruning takes several minutes to implement. Weights of the LD-weighted GRM take  $<15$  min to compute, whereas weights of LDAK take  $>15$  hr to compute.

The performances of estimators were compared on the simulated data. To compare performance both across estimators and across relationship types, we summarize estimation

accuracy by the ratio of the square root of mean squared error (MSE) and the average realized kinship,

$$\frac{\sqrt{\frac{1}{1000} \sum_{m=1}^{1000} (\hat{\Phi}_m - \Phi_m)^2}}{\frac{1}{1000} \sum_{m=1}^{1000} \Phi_m}, \quad (14)$$

where  $\Phi_m$ 's are realized kinship and  $\hat{\Phi}_m$ 's are their estimates.

#### Data availability

The 1000 Genomes Project Phase 3 data are available at <http://www.1000genomes.org/data>. The Rutgers Map version 3a is available at [http://compngen.rutgers.edu/download\\_maps.shtml](http://compngen.rutgers.edu/download_maps.shtml). Information on the set of 169,751 SNP markers used in the simulation study is provided as [File S2](#).

#### Results

Table 1 summarizes the performance of different kinship estimators. The estimators are divided into two groups: Group-1 uses only marker-allele frequencies, whereas those in Group-2 use additional information. First, every estimator works better on closer relatives. This is expected given we are measuring estimation accuracies relative to the average amount of sharing (Equation 14), and the coefficients of variation are higher for remote relatives. For each estimator, the raw MSEs are in fact smaller on remote relatives (Table S4). Second, estimators that make use of additional sources of information (Group-2) generally do better than estimators that do not (Group-1). This is also expected as chromosomes are inherited as segments. Information such as marker order, genetic positions, and LD pattern are informative of the joint inheritance across markers.

Relative performances of the GRM estimators in Group-1 match our expectations under the assumption of linkage equilibrium (Figure 1, A and B). The two-step GRM estimator ( $\hat{\Phi}_T$ ) compares favorably to others on all relationship types. The classic GRM estimator ( $\hat{\Phi}_G$ ) works well when  $\Phi \approx 0$  (e.g., third cousins). The robust GRM estimator ( $\hat{\Phi}_R$ ) is preferred to

**Table 1 Estimation accuracies from simulation study as measured by the ratio ( $\times 10^2$ ) of the square root of MSE and the average realized kinship (Equation 14)**

| Additional information        | Estimator                                        | Relationship |       |       |       |        |       |
|-------------------------------|--------------------------------------------------|--------------|-------|-------|-------|--------|-------|
|                               |                                                  | FS           | HS    | C1    | C2    | C3     | IN    |
| None                          | $\hat{\Phi}_P$ PLINK                             | 1.64         | 4.66  | 10.81 | 50.76 | 178.78 | 30.30 |
|                               | $\hat{\Phi}_M$ MLE                               | 1.51         | 3.55  | 7.26  | 27.00 | 78.66  | 12.79 |
|                               | $\hat{\Phi}_D$ Global Day-Williams               | 3.12         | 6.38  | 12.34 | 49.55 | 187.68 | 7.33  |
|                               | $\hat{\Phi}_G$ Classic GRM                       | 3.00         | 4.85  | 8.32  | 30.83 | 116.22 | 5.92  |
|                               | $\hat{\Phi}_R$ Robust GRM                        | 2.42         | 4.33  | 7.99  | 31.64 | 120.56 | 5.56  |
| LD pattern                    | $\hat{\Phi}_T$ Two-step GRM                      | 2.19         | 4.26  | 7.96  | 30.84 | 116.24 | 5.50  |
|                               | $\hat{\Phi}_N$ Naive LD-pruned GRM               | 3.01         | 4.70  | 7.87  | 28.20 | 105.46 | 5.82  |
| LD pattern                    | $\hat{\Phi}_K$ LDAK                              | 2.64         | 4.10  | 7.21  | 23.74 | 85.57  | 6.31  |
| LD pattern                    | $\hat{\Phi}_W$ LD-weighted GRM                   | 1.59         | 2.60  | 4.44  | 17.33 | 65.60  | 4.82  |
| Marker order                  | $\hat{\Phi}_L$ Local Day-Williams                | 2.04         | 2.76  | 6.52  | 10.79 | 24.26  | 8.31  |
| LD pattern + marker order     | $\hat{\Phi}_{LW}$ LD-weighted local Day-Williams | 2.10         | 2.85  | 6.70  | 11.27 | 23.56  | 8.39  |
| Genetic position              | $\hat{\Phi}_H$ HMM                               | 1.77         | 3.36  | 6.04  | 17.90 | 64.97  | 4.84  |
| LD pattern + genetic position | $\hat{\Phi}_{HW}$ LD-weighted HMM                | 1.57         | 2.83  | 5.16  | 14.55 | 52.19  | 4.52  |
| Pedigree                      | $\Psi$ Pedigree kinship                          | 7.80         | 10.49 | 17.63 | 37.30 | 70.16  | 14.73 |

All estimators compared require dense SNP genotypes and reliable sources of marker allele frequencies as input. Pedigree kinship for the six relationship types are 0.25 (FS), 0.125 (HS), 0.0625 (C1), 0.0156 (C2), 0.0039 (C3), and 0.1094 (IN), respectively.

$\hat{\Phi}_G$  on close relatives, and it dominates the global Day-Williams estimator ( $\hat{\Phi}_D$ ) on all relationship types.

The MLE estimator ( $\hat{\Phi}_M$ ) stands out among estimators in Group-1. Likelihood estimators are often known to be more accurate than method-of-moment estimators (Milligan 2003; Anderson and Weir 2007). However, accuracy and computational efficiency of  $\hat{\Phi}_M$  is extremely sensitive to the starting configurations and the convergence criterion of the EM algorithm. Our implementation of  $\hat{\Phi}_M$  adopted very favorable conditions (see *Methods of analysis*). Otherwise, the results were much less accurate and computation time much longer (results not shown). For full siblings, the PLINK estimator ( $\hat{\Phi}_P$ ) is more accurate than any Group-1 GRM estimator. Since  $\hat{\Phi}_P$  estimates the  $k$  coefficients directly, this provides higher resolution on full siblings who share two IBD genes over, on average, 25% of the genome. When inbreeding is present,  $\hat{\Phi}_P$  and  $\hat{\Phi}_M$  perform poorly relative to the other Group-1 estimators. These two estimators estimate the zero-inbreeding  $k$  coefficients directly and are thus more sensitive to violation of the no-inbreeding assumption.

Among the estimators of Group-2, the local Day-Williams estimator ( $\hat{\Phi}_L$ ) performs better than the others on remote relatives. However, the smoothing algorithm of  $\hat{\Phi}_L$  tends to produce downward bias (see *File S1*, section C, and *Table S5*) so that  $\hat{\Phi}_L$  must perform well on remote relatives where there is not much room for underestimation. In contrast, the HMM estimator ( $\hat{\Phi}_H$ ) generally overestimates IBD in the presence of LD (Brown *et al.* 2012). The naive LD-pruned GRM estimator ( $\hat{\Phi}_N$ ) shows slight improvement over the classic GRM estimator ( $\hat{\Phi}_G$ ), but loses to the LDAK ( $\hat{\Phi}_K$ ) on all occasions. The LD-weighted GRM estimator ( $\hat{\Phi}_W$ ) dominates both  $\hat{\Phi}_H$  and  $\hat{\Phi}_K$  in performance and loses to  $\hat{\Phi}_L$  only on remote relatives. This reflects the amount of LD present in the selected marker panel in the combined population, and shows that appropriately adjusting for patterns of LD can significantly improve the accuracy of kinship estimation.

When inbreeding is present, performance of  $\hat{\Phi}_L$  is the most affected among the Group-2 estimators. Glazner and Thompson (2015) noted that, in their example, this local IBD Day-Williams method failed to pick up short segments of complex (autozygous) IBD; the varying kinship levels across short distances seem to challenge this method (see *File S1*, section C). The performance of  $\hat{\Phi}_W$  is also affected by inbreeding. Perhaps the higher IBD levels in inbred individuals conflict with the assumption of unrelatedness in the estimator derivation.

When the LD weights are combined with the local IBD methods, there is clear improvement in performance for the HMM method (compare  $\hat{\Phi}_H$  to  $\hat{\Phi}_{HW}$ ), but less so for the Day-Williams local method (compare  $\hat{\Phi}_L$  and  $\hat{\Phi}_{LW}$ ). As noted above, the HMM overestimates IBD in high-LD regions, so that the LD weights are beneficial.

Pedigree kinship ( $\Psi$ ) is included in Table 1; it may be considered an estimator based only on the pedigree and not on genetic data. The MSE here represents the variation in realized kinship among pairs of individuals in the same pedigree relationship. The advantage of using genetic-data-based estimates of realized kinship instead of pedigree kinship is clear. Only on remote relatives (C2 and C3) do some of the marker-based estimates differ from the realized kinship by more than the pedigree values do. In fact, the results of Table 1 underplay the performance of the GRM estimators on remote relatives, since, for all the other estimators, estimates are constrained to be within the  $[0, 1]$  range. A comparison of constrained and unconstrained performance of the GRM estimators is shown in *Table S6*.

## Discussion

We have shown that improved estimators of realized kinship can be obtained (1) by optimal weighting of markers, (2) by taking physical contiguity of genome into account, and (3) by weighting on the basis of LD. In practice, the choice of

estimator largely depends on the availability of information. When one only has SNP genotypes and marker allele frequencies to work with, the two-step GRM estimator ( $\hat{\Phi}_T$ ) is both accurate and computationally efficient. If large genotyped samples from the relevant population are available, the LD-weighted GRM estimator ( $\hat{\Phi}_W$ ) is an attractive alternative. The LD weights can be computed very efficiently using existing optimization software, and the computation needs to be done only once for a population. If information on marker order or genetic positions is available, either the local Day-Williams estimator ( $\hat{\Phi}_L$ ) or the HMM estimator ( $\hat{\Phi}_H$ ) can offer a substantial increase in accuracy at the cost of longer computation time (see *Methods of analysis*). Once computed, these local IBD estimates across the genome can also be used in other analyses that use location-specific IBD: for example, in gene mapping.

In the derivation of optimal estimators, assumptions such as absence of inbreeding, linkage equilibrium, or unrelatedness were necessary to keep computations tractable. However, the proposed estimators applied well outside the initially assumed context. The two-step GRM estimator ( $\hat{\Phi}_T$ ) does not seem to be affected by the presence of inbreeding. It compares favorably to other estimators that use the same amount of information even when the assumption of linkage equilibrium is violated. The LD-weighted GRM estimator ( $\hat{\Phi}_W$ ) performed well on all relationship types considered, and is only slightly affected when related individuals happen to be inbred.

In our simulation study, relative pairs are generated as random draws from each relationship type. In practice, non-random sampling may create biases causing realized kinship to differ from the pedigree values. For instance, ascertainment by traits in human genetics and artificial selection in animal and plant genetics can result in sampled relatives being more (or less) genetically related than expected under the pedigree structure (Liu *et al.* 2003; Purcell *et al.* 2007). The comparison of pedigree vs. realized kinships in this article (Table 1) is thus an idealized best-case scenario.

Our simulation study used real haplotypes and a dense SNP marker panel. Thus LD is present in the simulated data. However, in assuming the variance form of Equation (6), the covariances due to LD are ignored (see Equation 5). To investigate the impact of LD on the relative performance of estimators, we compared the empirical and theoretical (no-LD) SDs for several estimators and relationship types. For any GRM estimator described in *Linkage equilibrium* and any relationship type (except the inbred cousins) listed in Table 1, the true variance is well approximated by empirical MSE. The theoretical (no-LD) variance (6) is computed using the simulation values of  $\Phi$  and  $k_2$ , and averaged across all 1000 pairs of that relationship type. Table S3 summarizes the results. We see that the factor by which the SD is underestimated by ignoring LD is generally smaller on remote relatives. More importantly, for a given relationship type, it is fairly consistent across estimators. This suggests that between GRM estimators that do not adjust for LD, relative efficiency computed

under the assumption of linkage equilibrium can be a good approximation to the true relative values.

Our study assumed population homogeneity so that allele frequencies and LD weights can be estimated once and applied to all pairs of relatives. This choice has the advantage of having a bigger pool of founder haplotypes available for sampling, and thus lower correlation among estimates from different simulation replicates. It also induces a more complex population structure in the simulated data, which is likely to influence performance of some estimators more than others. In limited additional experiments, we investigated performance of a subset of estimators on data simulated under either European ancestry or African ancestry, and compared results to those of Table 1 (see Table S7 and Table S8). Unsurprisingly, the GRM estimators generally benefit from the lesser structure in these more homogeneous pools of haplotypes. Relative performance among the GRM estimators that do not adjust for LD remains unchanged. The LD weighted GRM estimator ( $\hat{\Phi}_W$ ) loses to the two-step GRM estimator ( $\hat{\Phi}_T$ ) on close relatives, suggesting that the amount of LD adjustment (given the choice of marker panel and ancestry) is not enough to offset the effect of deviation from the assumption of unrelatedness. However, these results also suggest  $\hat{\Phi}_W$  is robust to population substructure and admixture; compared to other GRM estimators, its performance is much less affected by the complex structure of the combined population (compare Table 1 with Table S7 and Table S8). The HMM estimator ( $\hat{\Phi}_H$ ) does better under African ancestry (than under combined ancestry), and worse under European ancestry. This is likely due to the higher LD in the European population;  $\hat{\Phi}_H$  is sensitive to LD (Brown *et al.* 2012).

For convenience, we selected SNPs based on even genetic distance spacing and minor allele frequency (MAF). The relationship between genetic distance and LD is far from uniform, and our results on the impact of adjusting for LD show that it is a significant factor in our marker panel. The distribution of allele frequencies in our selected marker panel is quite similar to that in the commonly used OmniExpress24 genome-wide-association-study chip (see Figure S3). Compared to the OmniExpress24 autosomal markers, our selected marker panel has a slight underrepresentation for markers with  $MAF \geq 0.1$ . However, our panel was selected with a threshold of  $MAF \geq 0.05$ , whereas  $\sim 9\%$  of the OmniExpress24 autosomal markers fall below this threshold. We found that naive pruning of markers by MAF generally reduces accuracy of the two-step GRM estimator ( $\hat{\Phi}_T$ ), but can sometimes improve accuracy of other Group-1 GRM estimators depending on relationship types. Even on MAF-pruned marker sets,  $\hat{\Phi}_T$  always stands out among the Group-1 GRM estimators (results not shown). Overall, our selection of SNPs by genetic spacing and MAF create no strong biases in our estimator comparisons.

Even as SNP panels become denser, or sequence data become available, the issue of LD remains. Additional typed loci do not provide additional relatedness information without limit. Although methods that adjust for LD will gain from

the use of additional markers, other methods may not benefit and can be adversely affected. In particular, non-LD-adjusted local methods such as the HMM estimator ( $\hat{\Phi}_H$ ) should be applied on an LD-pruned marker set to avoid overestimation of IBD. In human applications with nominally unrelated individuals, haplotypic similarities due to LD must be distinguished from cryptic relatedness. In animal and plant breeding applications, high levels of LD are a regular feature of artificially selected populations with a small effective founder population size. Therefore, methods for efficient kinship estimation in the presence of LD remain relevant even in the age of full genome sequencing.

Estimators developed in this article do not specifically deal with population substructure or admixture, but they can be generalized to do so. Thornton *et al.* (2012) and Conomos *et al.* (2016) proposed two different methods that estimate individual-specific allele frequencies which take population substructure and admixture into account. The estimated individual-specific allele frequencies were subsequently applied to the classic GRM estimator and the robust GRM estimator, respectively, for kinship estimation. The same logic may be applied to calculate LD weights that adjust for population substructure or admixture. These frequencies and LD weights can then be used with our proposed estimators. The merits of such an approach to address population substructure or admixture is a topic for future studies.

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## Appendix A: Reparameterization of the Global Day-Williams Estimator

The original form of the global Day-Williams estimator introduced in Day-Williams *et al.* (2011) is

$$\hat{\Phi}_{uv} = \frac{e_{uv} - \sum_{i=1}^m (p_i^2 + q_i^2)}{m - \sum_{i=1}^m (p_i^2 + q_i^2)},$$

where  $\Phi_{uv}$  is the kinship between individual  $u$  and  $v$ ,  $e_{uv}$  is the observed number of IBS matches between  $u$  and  $v$ ,  $m$  is the total number of markers,  $p_i$  is the reference allele frequency at marker  $i$  and  $q_i = 1 - p_i$ .  $e_{uv}$  is defined as

$$e_{uv} = \sum_{i=1}^m o_{uv}^i = \sum_{i=1}^m \frac{1}{4} (\mathbf{1}_{\{I_i=K_i\}} + \mathbf{1}_{\{I_i=L_i\}} + \mathbf{1}_{\{J_i=K_i\}} + \mathbf{1}_{\{J_i=L_i\}}),$$

where  $I_i$  and  $J_i$  are the allelic types of the individual  $u$  at marker  $i$ , and  $K_i$  and  $L_i$  are the allelic types of the individual  $v$  at marker  $i$ .

Since we are working with genotypic data from pairs of individuals, indices of individuals within a pair are exchangeable, and so are indices of genes from the same individual at a given marker. Table A1 shows the correspondence between the Day-Williams and the GRM notations at each marker position. Note that  $o_{uv}^i$  and  $[1 + (x_i - 1)(y_i - 1)]/2$  are equivalent for all possible genotype combinations. Therefore, we can rewrite  $\hat{\Phi}_{uv}$  as

**Table A1 Correspondence between notations for all possible marker genotypes**

| Notation | Day-Williams           |            | GRM          |                              |
|----------|------------------------|------------|--------------|------------------------------|
| Genotype | $(I_i, J_i, K_i, L_i)$ | $o_{uv}^i$ | $(x_i, y_i)$ | $[1 + (x_i - 1)(y_i - 1)]/2$ |
| AA,AA    | (1,1,1,1)              | 1          | (2,2)        | 1                            |
| AA,AB    | (1,1,1,2)              | 0.5        | (2,1)        | 0.5                          |
| AA,BB    | (1,1,2,2)              | 0          | (2,0)        | 0                            |
| AB,AB    | (1,2,1,2)              | 0.5        | (1,1)        | 0.5                          |
| AB,BB    | (1,2,2,2)              | 0.5        | (1,0)        | 0.5                          |
| BB,BB    | (2,2,2,2)              | 1          | (0,0)        | 1                            |

$$\begin{aligned} \hat{\Phi}_{uv} &= \frac{e_{uv} - \sum_{i=1}^m (p_i^2 + q_i^2)}{m - \sum_{i=1}^m (p_i^2 + q_i^2)} \\ &= \frac{\sum_{i=1}^m o_{uv}^i - 1 + 2p_i(1 - p_i)}{\sum_{i=1}^m 2p_i(1 - p_i)} \\ &= \frac{\sum_{i=1}^m 1 + (x_i - 1)(y_i - 1) - 2 + 4p_i(1 - p_i)}{\sum_{i=1}^m 4p_i(1 - p_i)} \\ \hat{\Phi}_D &= \frac{\sum_{i=1}^m x_i y_i - (x_i + y_i) + 4p_i - 4p_i^2}{\sum_{i=1}^m 4p_i(1 - p_i)}, \end{aligned}$$

which recovers the expression in Equation 4.

## Appendix B: Computations in Methods

### General Case

Let  $\mathbf{1}_{x_1}$  and  $\mathbf{1}_{x_2}$  be the indicator functions that the two alleles of individual 1 at marker  $l$  are the reference allele, respectively, with  $l$  omitted from the notation. Define  $\mathbf{1}_{y_1}$  and  $\mathbf{1}_{y_2}$  similarly for individual 2. It is easy to see that  $x_l = \mathbf{1}_{x_1} + \mathbf{1}_{x_2}$  and  $y_l = \mathbf{1}_{y_1} + \mathbf{1}_{y_2}$ . Recall that the basic assumptions are:

1. IBD genes are of the same allelic types, whereas non-IBD genes are of independent allelic types.
2. Either of the two genes from one individual is equal likely to be IBD to either of the two genes of the other individual at the same locus.

We have

$$\begin{aligned}\mathbb{E}[x_l] &= 2\mathbb{E}[\mathbf{1}_{x_1}] = 2p_l, \\ \mathbb{E}[x_ly_l] &= \mathbb{E}[(\mathbf{1}_{x_1} + \mathbf{1}_{x_2})(\mathbf{1}_{y_1} + \mathbf{1}_{y_2})] \\ &= 4\mathbb{E}[\mathbf{1}_{x_1}\mathbf{1}_{y_1}] \\ &= 4\Phi p_l + 4(1 - \Phi)p_l^2,\end{aligned}$$

and

$$\begin{aligned}\mathbb{E}[Z_l(a_l)] &= \mathbb{E}\left[\frac{x_ly_l - a_l(x_l + y_l) + 4a_lp_l - 4p_l^2}{4p_l(1 - p_l)}\right] \\ &= \frac{4\Phi p_l + 4(1 - \Phi)p_l^2 - a_l \times 4p_l + 4a_lp_l - 4p_l^2}{4p_l(1 - p_l)} \\ &= \Phi.\end{aligned}$$

Since

$$\mathbb{E}[\hat{\Phi}(\mathbf{a}, \mathbf{w})] = \mathbb{E}\left[\sum_{l=1}^L w_l Z_l(a_l)\right] = \sum_{l=1}^L w_l \times \Phi = \Phi,$$

$\hat{\Phi}(\mathbf{a}, \mathbf{w})$  is unbiased for any  $\mathbf{a}$  and  $\mathbf{w}$ .

## Linkage Equilibrium

We now assume linkage equilibrium and absence of inbreeding. To derive an expression for  $V_l(a_l)$  defined in (5), note that

$$\begin{aligned}\mathbb{E}[x_l^2] &= \mathbb{E}[\mathbf{1}_{x_1}^2 + 2\mathbf{1}_{x_1}\mathbf{1}_{x_2} + \mathbf{1}_{x_2}^2] \\ &= 2p_l + 2p_l^2, \\ \text{Var}[x_l] &= \mathbb{E}[x_l^2] - (\mathbb{E}[x_l])^2 \\ &= 2p_l + 2p_l^2 - 4p_l^2 \\ &= 2p_l(1 - p_l), \\ \text{Cov}[x_l, y_l] &= \mathbb{E}[(x_l - 2p_l)(y_l - 2p_l)] \\ &= \mathbb{E}[x_ly_l - 2p_l x_l - 2p_l y_l + 4p_l^2] \\ &= 4\Phi p_l(1 - p_l), \\ \mathbb{E}[x_l^2 y_l] &= \mathbb{E}[(\mathbf{1}_{x_1} + \mathbf{1}_{x_2})^2(\mathbf{1}_{y_1} + \mathbf{1}_{y_2})] \\ &= 4\mathbb{E}[\mathbf{1}_{x_1}^2 \mathbf{1}_{y_1}] + 4\mathbb{E}[\mathbf{1}_{x_1}\mathbf{1}_{x_2}\mathbf{1}_{y_1}] \\ &= 4\Phi p_l + 4(1 - \Phi)p_l^2 + 4[2\Phi p_l^2 + (1 - 2\Phi)p_l^3] \\ &= 4\Phi p_l + 4p_l^2 + 4\Phi p_l^2 + 4p_l^3 - 8\Phi p_l^3, \\ \text{Cov}[x_ly_l, x_l] &= \mathbb{E}\{[x_ly_l - 4\Phi p_l - 4(1 - \Phi)p_l^2](x_l - 2p_l)\} \\ &= \mathbb{E}[x_l^2 y_l] - 2p_l \mathbb{E}[x_ly_l] \\ &= 4\Phi p_l + 4p_l^2 + 4\Phi p_l^2 + 4p_l^3 - 8\Phi p_l^3 \\ &\quad - 8\Phi p_l^2 - 8p_l^3 + 8\Phi p_l^3 \\ &= 4\Phi p_l + 4p_l^2 - 4\Phi p_l^2 - 4p_l^3 \\ &= 4(\Phi + p_l)p_l(1 - p_l).\end{aligned}$$

Calculation of the term  $\mathbb{E}[\mathbf{1}_{x_1}\mathbf{1}_{x_2}\mathbf{1}_{y_1}]$  involves probabilities of the underlying IBD states between the two genes of individual 1 and one gene of individual 2. Given the assumption of no inbreeding, there are only three possible IBD states with probabilities given in Table B1.

**Table B1 IBD states and probabilities for the two genes of individual 1 and one random gene from individual 2**

| IBD state   | (1,2,1) | (1,2,2) | (1,2,3)     |
|-------------|---------|---------|-------------|
| Probability | $\Phi$  | $\Phi$  | $1 - 2\Phi$ |

Following the above results, the closed-form expression of the variance  $V_l(a_l)$  of Equation 5 is derived in Box B1, assuming  $0 < p_l < 1$ . For general choices of  $a_l$ ,  $V_l(a_l)$  is not symmetric about  $p_l = 0.5$ , and is thus sensitive to the choice of reference allele. The part of  $V_l(a_l)$  that is responsible for this asymmetry is

$$\begin{aligned}
\mathbb{E}[x_l^2 y_l^2] &= \mathbb{E}\left[\left(\mathbf{1}_{x_1} + \mathbf{1}_{x_2}\right)^2 \left(\mathbf{1}_{y_1} + \mathbf{1}_{y_2}\right)^2\right] \\
&= 4\mathbb{E}\left[\mathbf{1}_{x_1}^2 \mathbf{1}_{y_1}^2\right] + 8\mathbb{E}\left[\mathbf{1}_{x_1} \mathbf{1}_{x_2} \mathbf{1}_{y_1}^2\right] + 4\mathbb{E}\left[\mathbf{1}_{x_1} \mathbf{1}_{x_2} \mathbf{1}_{y_1} \mathbf{1}_{y_2}\right] \\
&= 4\Phi p_l + 4(1-\Phi)p_l^2 + 8\left[2\Phi p_l^2 + (1-2\Phi)p_l^3\right] + 4\left[k_2 p_l^2 + k_1 p_l^3 + k_0 p_l^4\right] \\
&= 4\Phi p_l + 4p_l^2 + 12\Phi p_l^2 + 8p_l^3 - 16\Phi p_l^3 + 4\left[k_2 p_l^2 + k_1 p_l^3 + k_0 p_l^4\right], \\
\text{Var}[x_l y_l - a_l(x_l + y_l)] &= \text{Var}[x_l y_l] + a_l^2 \text{Var}[x_l + y_l] - 2a_l \text{Cov}[x_l y_l, x_l + y_l] \\
&= \mathbb{E}[x_l^2 y_l^2] - \left(\mathbb{E}[x_l y_l]\right)^2 + 2a_l^2 \text{Var}[x_l] + 2a_l^2 \text{Cov}[x_l, y_l] - 4a_l \text{Cov}[x_l y_l, x_l] \\
&= 4\Phi p_l + 4p_l^2 + 12\Phi p_l^2 + 8p_l^3 - 16\Phi^2 p_l^2 - 48\Phi p_l^3 - 16p_l^4 + 32\Phi^2 p_l^3 + 32\Phi p_l^4 - 16\Phi^2 p_l^4 \\
&\quad + 4\left[k_2 p_l^2 + k_1 p_l^3 + k_0 p_l^4\right] + \left[4a_l^2 + 8a_l^2 \Phi - 16a_l(\Phi + p_l)\right] p_l(1-p_l) \\
&= 4\Phi p_l + 4p_l^2 + 12\Phi p_l^2 + 8p_l^3 - 16\Phi^2 p_l^2 - 48\Phi p_l^3 - 16p_l^4 + 32\Phi^2 p_l^3 + 32\Phi p_l^4 - 16\Phi^2 p_l^4 \\
&\quad + 4\left[k_2 p_l^2 + (4\Phi - 2k_2)p_l^3 + (1 - k_2 - 4\Phi + 2k_2)p_l^4\right] + \left[4a_l^2 + 8a_l^2 \Phi - 16a_l(\Phi + p_l)\right] p_l(1-p_l) \\
&= 4\Phi p_l + 4p_l^2 + 12\Phi p_l^2 + 8p_l^3 - 16\Phi^2 p_l^2 - 48\Phi p_l^3 - 16p_l^4 + 32\Phi^2 p_l^3 + 32\Phi p_l^4 - 16\Phi^2 p_l^4 \\
&\quad + 4\left[k_2 p_l^2 (1-p_l)^2 + 4\Phi p_l^3 + p_l^4 - 4\Phi p_l^4\right] + \left[4a_l^2 + 8a_l^2 \Phi - 16a_l(\Phi + p_l)\right] p_l(1-p_l) \\
&= 16\Phi(1-\Phi)p_l^2(1-p_l)^2 + 4p_l^2(1+3p_l)(1-p_l) + 4\Phi p_l(1-p_l) + 4k_2 p_l^2(1-p_l)^2 \\
&\quad + \left[4a_l^2 + 8a_l^2 \Phi - 16a_l(\Phi + p_l)\right] p_l(1-p_l), \\
V_l(a_l) &= \text{Var}\left[\frac{x_l y_l - a_l(x_l + y_l) + 4a_l p_l - 4p_l^2}{4p_l(1-p_l)}\right] \\
&= \frac{\text{Var}[x_l y_l - a_l(x_l + y_l)]}{16p_l^2(1-p_l)^2} \\
&= \frac{4\Phi(1-\Phi)p_l(1-p_l) + (k_2 + 1)p_l(1-p_l) + (a_l - 2p_l)^2 + 2\Phi(a_l - 1)^2 - \Phi}{4p_l(1-p_l)}.
\end{aligned}$$

**Box 1** Derivation of  $V_l(a_l)$  under the assumption of no inbreeding.

$$(a_l - 2p_l)^2 + 2\Phi(a_l - 1)^2,$$

where  $a_l$  can be a function of  $p_l$ .

Suppose, however, that  $a_l$  is a weighted average of  $2p_l$  and 1:  $a_l = b \times 2p_l + (1 - b)$  for  $b \in [0, 1]$ . Then

$$\begin{aligned}
&(a_l - 2p_l)^2 + 2\Phi(a_l - 1)^2 \\
&= (b \times 2p_l + 1 - b - 2p_l)^2 + 2\Phi(b \times 2p_l + 1 - b - 1)^2 \\
&= (2p_l - 1)^2 \left[ (b - 1)^2 + 2\Phi b^2 \right].
\end{aligned}$$

Then  $V_l(a_l)$  takes the value

$$4\Phi(1 - \Phi) + k_2 + 1 + \frac{(2p_l - 1)^2 \left[ (b-1)^2 + 2\Phi b^2 \right] - \Phi}{4p_l(1 - p_l)},$$

which is symmetric about  $p_l = 0.5$ , and it attains its minimum at  $p_l = 0.5$ , conditional on  $\Phi$  and  $k_2$ .

Returning to the general form of  $V_l(a_l)$  and setting its derivative with respect to  $a_l$  equal to 0, leads to

$$\tilde{a}_l = \frac{1}{1 + 2\Phi} \times 2p_l + \frac{2\Phi}{1 + 2\Phi}.$$

Thus the optimal  $\tilde{a}_l$  is a weighted average of  $2p_l$  and 1, and  $V_l(\tilde{a}_l)$  is invariant to the choice of reference allele.

For fixed  $\mathbf{a}$  [and therefore fixed  $\mathbf{V}(\mathbf{a})$ ], we can find the set of optimal weights  $\tilde{\mathbf{w}}(\mathbf{a})$  by solving the following minimization problem:

$$\min_{\mathbf{w}} \sum_{l=1}^L w_l^2 V_l(a_l) \quad : \quad \mathbf{w}^T \mathbf{1} = 1, \quad w_l \geq 0 \quad \forall l.$$

With Lagrange multipliers  $\lambda$ ,

$$\mathcal{L}(\mathbf{w}, \lambda) = \sum_{l=1}^L w_l^2 V_l(a_l) - \lambda \left( \sum_{l=1}^L w_l - 1 \right)$$

$$\frac{\partial \mathcal{L}}{\partial w_l} = 2w_l V_l(a_l) - \lambda = 0$$

$$\tilde{w}_l(\mathbf{a}) = \frac{\lambda}{2} \times V_l(a_l)^{-1}.$$

The above expression holds true for all  $l$ , implying

$$\tilde{w}_l(\mathbf{a}) = \frac{V_l(a_l)^{-1}}{\sum_{m=1}^L V_m(a_m)^{-1}}.$$

The nonnegativity constraint is automatically satisfied as  $V_l(a_l) > 0$  for all  $l$ .

### Linkage Disequilibrium

We drop the assumptions of linkage equilibrium and absence of inbreeding, but assume instead that the two individuals are unrelated, so that  $x_l \perp y_m$  are independent, for all  $l$  and  $m$ . Under the new assumptions, all of the results from *General Case* still hold, but most of the results from *Linkage Equilibrium* do not. Let the inbreeding coefficients of the two individuals be  $F_x$  and  $F_y$ , respectively,

$$\begin{aligned} \mathbb{E}[x_l^2] &= \mathbb{E}[\mathbf{1}_{x_1}^2 + 2\mathbf{1}_{x_1}\mathbf{1}_{x_2} + \mathbf{1}_{x_2}^2] \\ &= 2p_l[1 + F_x + (1 - F_x)p_l], \\ \text{Var}[x_l] &= \mathbb{E}[x_l^2] - (\mathbb{E}[x_l])^2 \\ &= 2p_l[1 + F_x + (1 - F_x)p_l] - 4p_l^2 \\ &= 2p_l(1 - p_l)(F_x + 1). \end{aligned}$$

Analogous results hold for  $y_l$ . It can be verified (line 5 of Box B1) that

$$\tilde{a}_l = \arg \min_{a_l} V_l(a_l) = \frac{\text{Cov}[x_l y_l, x_l + y_l]}{\text{Var}[x_l + y_l]}.$$

This time

$$\begin{aligned}
\tilde{a}_l &= \frac{\text{Cov}[x_l y_l, x_l] + \text{Cov}[x_l y_l, y_l]}{\text{Var}[x_l] + \text{Var}[y_l]} \\
&= \frac{(\mathbb{E}[x_l^2] - 2\mathbb{E}[x_l]\mathbb{E}[y_l] + \mathbb{E}[y_l^2]) \times 2p_l}{\mathbb{E}[x_l^2] - (\mathbb{E}[x_l])^2 + \mathbb{E}[y_l^2] - (\mathbb{E}[y_l])^2}, \\
&= 2p_l,
\end{aligned}$$

since

$$2\mathbb{E}[x_l]\mathbb{E}[y_l] = (\mathbb{E}[x_l])^2 + (\mathbb{E}[y_l])^2 = 8p_l^2.$$

Now

$$\begin{aligned}
&\text{Cov}[x_l y_l - a_l(x_l + y_l), x_m y_m - a_m(x_m + y_m)] \\
&= \text{Cov}[x_l y_l, x_m y_m] - a_m \text{Cov}[x_l y_l, x_m + y_m] \\
&\quad - a_l \text{Cov}[x_m y_m, x_l + y_l] + a_l a_m \text{Cov}[x_l + y_l, x_m + y_m],
\end{aligned}$$

where

$$\begin{aligned}
\text{Cov}[x_l y_l, x_m + y_m] &= \text{Cov}[x_l y_l, x_m] + \text{Cov}[x_l y_l, y_m] \\
&= \mathbb{E}[x_l x_m] \mathbb{E}[y_l] - \mathbb{E}[x_l] \mathbb{E}[x_m] \mathbb{E}[y_l] \\
&\quad + \mathbb{E}[y_l y_m] \mathbb{E}[x_l] - \mathbb{E}[y_l] \mathbb{E}[y_m] \mathbb{E}[x_l] \\
&= 2p_l (\mathbb{E}[x_l x_m] + \mathbb{E}[y_l y_m] - 8p_l p_m), \\
\text{Cov}[x_m y_m, x_l + y_l] &= 2p_m (\mathbb{E}[x_l x_m] + \mathbb{E}[y_l y_m] - 8p_l p_m), \\
\text{Cov}[x_l + y_l, x_m + y_m] &= \text{Cov}[x_l, x_m] + \text{Cov}[y_l, y_m] \\
&= \mathbb{E}[x_l x_m] + \mathbb{E}[y_l y_m] - 8p_l p_m.
\end{aligned}$$

Setting  $\mathbf{a} = 2\mathbf{p}$ ,

$$\begin{aligned}
&\text{Cov}[x_l y_l - 2p_l(x_l + y_l), x_m y_m - 2p_m(x_m + y_m)] \\
&= \text{Cov}[x_l y_l, x_m y_m] - 4p_l p_m (\mathbb{E}[x_l x_m] + \mathbb{E}[y_l y_m] - 8p_l p_m) \\
&= \mathbb{E}[x_l x_m] \mathbb{E}[y_l y_m] - 4p_l p_m (\mathbb{E}[x_l x_m] + \mathbb{E}[y_l y_m]) + 16p_l^2 p_m^2 \\
&= (\mathbb{E}[x_l x_m] - 4p_l p_m)(\mathbb{E}[y_l y_m] - 4p_l p_m) \\
&= \rho_{lm} \sqrt{\text{Var}[x_l] \cdot \text{Var}[x_m]} \times \rho_{lm} \sqrt{\text{Var}[y_l] \cdot \text{Var}[y_m]} \\
&= 4p_l(1-p_l)p_m(1-p_m)(F_x + 1)(F_y + 1)\rho_{lm}^2.
\end{aligned}$$

It then follows that

$$\begin{aligned}
&\text{Cov}[Z_l(2p_l), Z_m(2p_m)] \\
&= \frac{\text{Cov}[x_l y_l - 2p_l(x_l + y_l), x_m y_m - 2p_m(x_m + y_m)]}{4p_l(1-p_l) \times 4p_m(1-p_m)} \\
&= \frac{1}{4} (F_x + 1)(F_y + 1)\rho_{lm}^2,
\end{aligned}$$

$$V_l(2p_l) = \text{Var}[Z_l(2p_l)] = \frac{1}{4} (F_x + 1)(F_y + 1).$$

We assume that the matrix of squared LD correlations,  $\mathbf{R} = [\rho_{lm}^2]$ , is known. Correlation matrices are positive semidefinite. By the Schur product theorem,  $\mathbf{R}$ , the Hadamard product of a correlation matrix and itself must also be positive semidefinite.

As noted in the text (Equation 13), we solve the following minimization problem for the LD weights,

$$\min_{\mathbf{w}} [\mathbf{w}^T \mathbf{R} \mathbf{w} - \mathbf{w}^T \mathbf{1}] \quad : \quad w_l \geq 0 \quad \forall l.$$

When  $\mathbf{R}$  has a block-diagonal structure (e.g., each chromosome forms a block), we can rewrite the minimization problem in terms of subvectors and submatrices,

$$\min_{\mathbf{w}} \sum_{i=1}^n [\mathbf{w}_{(i)}^T \mathbf{R}_{(i)} \mathbf{w}_{(i)} - \mathbf{w}_{(i)}^T \mathbf{1}] \quad : \quad w_l \geq 0 \quad \forall l.$$

Since the  $\mathbf{w}_{(i)}$ 's form a partition of  $\mathbf{w}$  and the  $\mathbf{R}_{(i)}$ 's are independent submatrices of  $\mathbf{R}$ , minimization can be implemented for each block independently. We can solve for  $\mathbf{w}_{(i)}$  in block  $i$  using the corresponding submatrix  $\mathbf{R}_{(i)}$ . If  $\mathbf{w}_{(i)}^T \mathbf{1} = c_i$  for the block solutions, the final solution  $\tilde{\mathbf{w}}$  will be a concatenation of the block solutions  $\mathbf{w}_{(i)}$ 's rescaled by  $1/\sum_i c_i$ .