

# Variation in Genetic Relatedness Is Determined by the Aggregate Recombination Process

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**ABSTRACT** The genomic proportion that two relatives share identically by descent—their genetic relatedness—can vary depending on the history of recombination and segregation in their pedigree. Previous calculations of the variance of genetic relatedness have defined genetic relatedness as the proportion of total genetic map length (cM) shared by relatives, and have neglected crossover interference and sex differences in recombination. Here, we consider genetic relatedness as the proportion of the total physical genome (bp) shared by relatives, and calculate its variance for general pedigree relationships, making no assumptions about the recombination process. For the relationships of grandparent-grandoffspring and siblings, the variance of genetic relatedness is a simple decreasing function of  $\bar{r}$ , the average proportion of locus pairs that recombine in meiosis. For general pedigree relationships, the variance of genetic relatedness is a function of metrics analogous to  $\bar{r}$ . Therefore, features of the aggregate recombination process that affect  $\bar{r}$  and analogs also affect variance in genetic relatedness. Such features include the number of chromosomes and heterogeneity in their size, the number of crossovers and their spatial organization along chromosomes, and sex differences in recombination. Our calculations help to explain several recent observations about variance in genetic relatedness, including that it is reduced by crossover interference (which is known to increase  $\bar{r}$ ). Our methods further allow us to calculate the neutral variance of ancestry among  $F_2$ s in a hybrid cross, enabling precise statistical inference in  $F_2$ -based tests for various kinds of selection.

**KEYWORDS** Meiosis; recombination; identity by descent; relatedness; heritability; genetic mapping; crossover interference

**V**ARIANCE in the amount of DNA shared by relatives identically by descent (IBD)—variance in genetic relatedness—is an important quantity in genetics (Thompson 2013). It translates to variance in the phenotypic similarity of relatives, and is a vital component of pedigree-based estimates of heritability and the genetic variance of traits (Visscher *et al.* 2006, 2007; Young *et al.* 2018). It is also an important consideration when estimating pedigree relationships and the degree of inbreeding from genotype data (Kardos *et al.* 2015; Wang 2016). Variance in genetic relatedness has also been hypothesized to have important consequences for the evolution of behavior (Barash *et al.* 1978) and of karyotypes and recombination rates (Sherman 1979;

Wilfert *et al.* 2007). Moreover, as we show elsewhere, variance in genetic relatedness plays a key role in selection against deleterious introgressed DNA following hybridization (Veller *et al.* 2019a).

For most pedigree relationships, genetic relatedness can vary because of variable patterns of recombination and segregation within the pedigree. For example, it is possible that a mother segregates only crossoverless paternal chromatids to an egg, in which case the resulting offspring inherits one half of its genome from its maternal grandfather and none from its maternal grandmother. On the other hand, if the mother shuffles her maternal and paternal DNA thoroughly into the egg, the offspring will be approximately equally genetically related to its maternal grandparents. Thus, intuitively, a higher degree of genetic shuffling within a pedigree leads to lower variance in genetic relatedness between relatives.

Previous theoretical calculations of the variance of genetic relatedness have largely been restricted to measuring genetic relatedness as the proportion of total genetic map length (in cM) shared IBD by relatives [e.g., Franklin (1977); Hill

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(1993b); Guo (1996); Visscher *et al.* (2006); a general treatment is given by Hill and Weir (2011)]. However, measuring genetic relatedness as the proportion of map length shared causes several problems, most notably when the genetic maps of the two sexes differ, as will typically be the case (Lenormand and Dutheil 2005; Sardell and Kirkpatrick 2020). This is easiest to appreciate for the genetic relatedness of an individual to its paternal and maternal grandparents, the values of which are determined in a paternal and a maternal meiosis, respectively. Theoretical calculations of the variance of cM genetic relatedness require the use of genetic map lengths from the relevant meioses, and thus, in these two cases, require different definitions of genetic relatedness: proportion of total male map length for relatedness to paternal grandparents, and proportion of total female map length for relatedness to maternal grandparents. Indeed, in the extreme case where crossing over is absent in one sex—say males, as in *Drosophila*—cM genetic relatedness to paternal grandparents is undefined in these calculations, because the male map length is 0 cM. Practically speaking, these problems can be sidestepped by defining cM genetic relatedness in terms of a sex-averaged genetic map, but this leads to substantial biases in theoretical calculations of its variance (Caballero *et al.* 2019).

A natural alternative that avoids such problems is to measure genetic relatedness as the proportion of the physical length of the genome (in bp) shared IBD by relatives. For many purposes, bp genetic relatedness will be the more appropriate measure (White and Hill 2020) and, unlike cM genetic relatedness, bp genetic relatedness is unambiguous when there are sex differences in recombination. Moreover, in the modern genomic era, it will often be the case that a species' genome has been sequenced before its genetic map has been elucidated, so that only bp genetic relatedness can be assayed.

Translating previous calculations of the variance of cM genetic relatedness to the variance of bp genetic relatedness would be valid only under the assumption of uniform recombination rates along chromosomes. This assumption is unrealistic for most species. For example, crossovers tend to be terminally localized along human chromosomes, especially in males (Holm and Rasmussen 1983; Bojko 1985). White and Hill (2020) have recently developed a procedure to estimate the variance of bp genetic relatedness without the assumption of uniform recombination rates. However, their method still assumes uniform recombination rates in the regions between adjacent markers, making it best applicable to high-density linkage maps (rather than low-density linkage maps or cytological data, which will be more readily available for some species).

In addition, previous theoretical calculations of the variance of genetic relatedness (including those for bp genetic relatedness) have assumed that crossover interference is absent. However, it has recently been shown, by computer simulation of various forms of crossover patterning along chromosomes, that crossover interference tends to decrease

variances of genetic relatedness (Caballero *et al.* 2019). Since crossover interference is a nearly ubiquitous feature of meiosis (Hillers 2004; Otto and Payseur 2019), its neglect in previous calculations of the variance of genetic relatedness further limits their generality.

In this paper, we derive a general, assumption-free formulation for the variance of bp genetic relatedness. We show that the variance of genetic relatedness is a simple, decreasing function of certain newly developed metrics of genome-wide genetic shuffling:  $\bar{r}$  and analogs (Veller *et al.* 2019b). These metrics, in a natural and intuitive way, take into account features of the aggregate recombination process, such as the number of chromosomes and heterogeneity in their size, the number of crossovers and their location along the chromosomes, the spatial relations of crossovers with respect to each other (*e.g.*, crossover interference), and sex differences in recombination.

Our formulation of the variance of genetic relatedness in terms of  $\bar{r}$  and analogs allows the effects that the above meiotic features have on the variance of genetic relatedness to be reinterpreted—often with greater intuition—in terms of their effects on aggregate genetic shuffling. For example, the fact that crossover interference decreases the variance of genetic relatedness (Caballero *et al.* 2019) can be explained by the intuitive fact that crossover interference, by spreading crossovers out evenly along chromosomes, increases the amount of genetic shuffling that they cause (Gorlov and Gorlova 2001; Veller *et al.* 2019b).

In the calculations below, the number of loci in the genome is assumed to be very large. Loci  $i$  and  $j$  are recombinant in a random gamete with probability  $r_{ij}$  (*e.g.*,  $r_{ij} = 1/2$  if  $i$  and  $j$  are on different chromosomes). Sex-specific recombination rates,  $r_{ij}^{\circ}$  and  $r_{ij}^{\sigma}$ , are distinguished where necessary. We assume that there is no inbreeding; for a treatment of the variance of cM genetic relatedness in finite populations, in which a degree of inbreeding is inevitable, see Carmi *et al.* (2013). “Genetic relatedness” refers to bp genetic relatedness, unless specified otherwise.

## Relationships of Direct Descent

Pedigree relationships of direct descent (or “lineal” relationships) involve a single lineage, from an ancestor to one of its descendants. We will focus here on the specific example of grandparent-grandoffspring; calculations of the variance of genetic relatedness for general relationships of direct descent are given in Supplemental Material, File S1, Section S1.

### Grandparent-grandoffspring

Let the random variable  $IBD_{\text{grand}}$  be the proportion of a grandoffspring's genome inherited from a specified grandparent. Consider the gamete produced by the grandoffspring's parent, and let  $\hat{P}$  be the fraction of this gamete's genome that derives from the focal grandparent (so that, by Mendelian segregation,  $\mathbb{E}[\hat{P}] = 1/2$ ). We first wish to calculate  $\text{Var}(\hat{P})$ . To do so, we use an approach very similar to that of Hill

(1993a) and Visscher *et al.* (2006), but we define genetic relatedness in terms of bp shared rather than cM shared, and make no assumptions about the recombination process (in File S1, Section S3, we discuss technical differences between our calculations of the variance of bp genetic relatedness and previous calculations of the variance of cM genetic relatedness). We calculate (details in File S1, Section S1) that

$$\text{Var}(\hat{P}) = \frac{1}{2} \left( \frac{1}{2} - \bar{r} \right), \quad (1)$$

where  $\bar{r}$  is the probability that a randomly chosen locus pair recombines in meiosis (Veller *et al.* 2019b). Because half of the grandoffspring's genome comes from this gamete,  $IBD_{\text{grand}} = \hat{P}/2$ , so that  $\mathbb{E}[IBD_{\text{grand}}] = \mathbb{E}[\hat{P}]/2 = 1/4$  is the coefficient of relationship, and

$$\text{Var}(IBD_{\text{grand}}) = \frac{1}{4} \text{Var}(\hat{P}) = \frac{1}{8} \left( \frac{1}{2} - \bar{r} \right). \quad (2)$$

A graphical demonstration of Equation 2, based on the possible segregation patterns of a given parental meiosis, is shown in Figure 1.

Note that the formulation in Equation 2 and other such formulations in this paper apply to the whole genome, or a single chromosome, or any specific genomic region. In the latter cases,  $\bar{r}$  is the probability that a randomly chosen pair of loci within the region of interest recombine in meiosis. In addition, because the recombination process often differs between the sexes, the value of  $\bar{r}$  can differ between spermatogenesis and oogenesis. In calculating the variance of genetic relatedness between a grandoffspring and one of its maternal grandparents, the value for oogenesis,  $\bar{r}^{\text{f}}$ , would be used; the value for spermatogenesis,  $\bar{r}^{\text{m}}$ , would be used for paternal grandparents.

$\bar{r}$  can be estimated from various kinds of data, including cytological data of crossover positions at meiosis I, sequence data from gametes, and linkage maps (Veller *et al.* 2019b). We used cytological data from Lian *et al.* (2008) to calculate chromosome-specific and genome-wide values of  $\bar{r}$  in human male, and the linkage map of Kong *et al.* (2010) to calculate analogous values in human female (translating map distances to recombination rates using Kosambi's map function, which incorporates a model of crossover interference). Substituting these values of  $\bar{r}$  into Equation 2 yields the variance of genetic relatedness to paternal and maternal grandparents in humans, for each chromosome and genome-wide. Table 1 displays the standard deviations, together with the corresponding standard deviations of cM genetic relatedness, calculated by substituting the sex-specific chromosome map lengths reported by Kong *et al.* (2010) into the relevant formula of Hill and Weir (2011).

Several observations emerge from Table 1. First, the variance of genetic relatedness for each individual chromosome is substantially larger than the genome-wide variance. This is because the majority of genetic shuffling in humans is due to

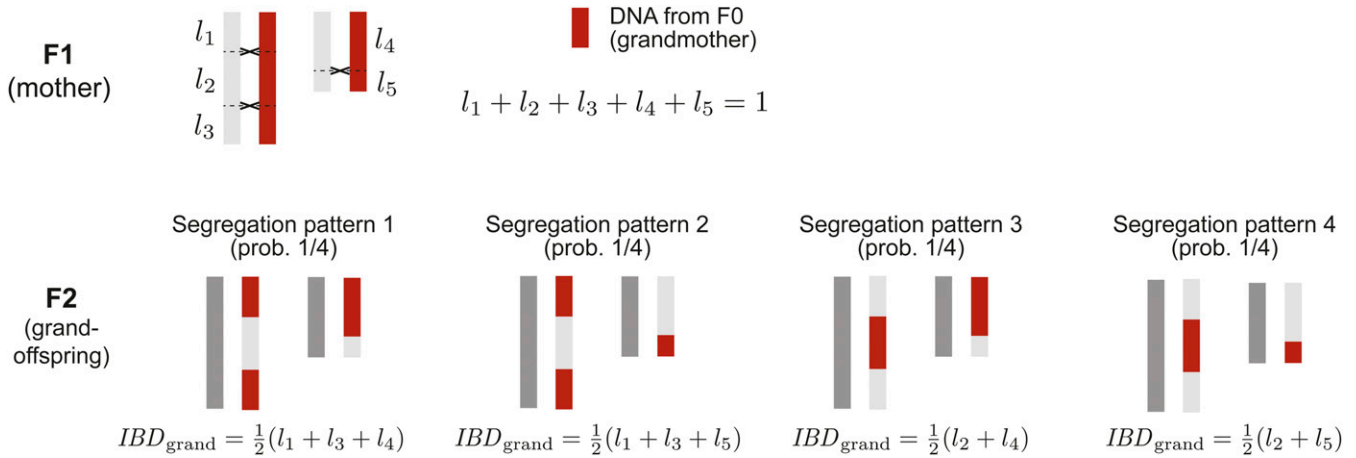
independent assortment of chromosomes, rather than crossing over (Crow 1988; Veller *et al.* 2019b). Second, the variance of genetic relatedness to a paternal grandparent is greater than to a maternal grandparent, for each chromosome and genome-wide. This is because male meiosis involves less genetic shuffling than female meiosis (lower  $\bar{r}$ ), owing to fewer crossovers and their more terminal localization along the chromosomes in males (Veller *et al.* 2019b).

In comparing the variances of bp and cM genetic relatedness, three meiotic features are relevant. Per-chromosome comparisons are affected by the location of crossovers along chromosomes (crossover distribution) and with respect to each other (crossover interference). The genome-wide comparisons are additionally influenced by independent assortment of chromosomes. We discuss the effects of these features in turn.

First, proterminal localization of crossovers in humans (especially males) reduces  $\bar{r}$  relative to a uniform distribution of crossovers (Veller *et al.* 2019b), increasing the variance of bp vs. cM genetic relatedness (since crossovers are uniformly distributed along the genetic map, by definition). To isolate this effect of nonuniform recombination rates, we artificially eliminate crossover interference in the calculation of  $\bar{r}$  by using linkage maps and Haldane's map function (which, unlike Kosambi's map function, assumes no crossover interference). Calculating  $\bar{r}$  in this way, we find that the chromosome-specific variances of bp genetic relatedness are typically larger than their corresponding cM values (File S1, Section S4), more so in males because of their more terminal distribution of crossovers.

Second, crossover interference increases  $\bar{r}$  by spreading crossovers out more evenly along chromosomes (Veller *et al.* 2019b), thus decreasing the variances of bp genetic relatedness relative to the corresponding variances of cM genetic relatedness (the calculations of which do not take into account crossover interference). Thus, in spite of the tendency of nonuniform recombination rates to increase the per-chromosome variances of bp genetic relatedness, these variances are nevertheless smaller than the corresponding variances of cM genetic relatedness when crossover interference is taken into account (Table 1). The negative effect of crossover interference on the variance of genetic relatedness was previously identified by Caballero *et al.* (2019). Interestingly, in human male, the per-chromosome variances of genetic relatedness calculated from raw (cytological) crossover data are smaller than those calculated from linkage maps using Kosambi's map function (File S1, Section S4), suggesting that Kosambi's map function does not capture the full influence of crossover interference on genetic shuffling in human male.

Finally, in humans, chromosome lengths are more variable when measured in bp than in cM (File S1, Section S3). This causes the contribution of independent assortment of chromosomes to  $\bar{r}$  to be smaller than if the bp lengths of the chromosomes were only as variable as the cM lengths (Veller *et al.* 2019b), which, in turn, increases the genome-wide



**Figure 1** The variance of genetic relatedness between grandoffspring and grandparent, calculated from the possible segregation patterns of a single parental meiosis. In the figure, the positions of crossovers in a maternal meiosis (and the chromatids involved) are specified, but the segregation pattern in the resulting egg (and therefore offspring) is not. Averaging across the four segregation patterns, we find  $\mathbb{E}[IBD_{\text{grand}}] = (l_1 + l_2 + l_3 + l_4 + l_5)/4 = 1/4$ , and, from Equation 1 in Veller et al. (2019b),  $\bar{r}^\sigma = (l_1 + l_3 + l_4)(l_2 + l_5) + (l_2 + l_4)(l_1 + l_3 + l_5)$ . Across the four possible segregation patterns,  $\mathbb{E}[IBD_{\text{grand}}] = 1/4$  and

$$\begin{aligned} \text{Var}(IBD_{\text{grand}}) &= \mathbb{E}[IBD_{\text{grand}}^2] - (\mathbb{E}[IBD_{\text{grand}}])^2 \\ &= \frac{1}{4} \left[ \frac{1}{4}(l_1 + l_3 + l_5)^2 + \frac{1}{4}(l_1 + l_3 + l_4)^2 + \frac{1}{4}(l_2 + l_4)^2 + \frac{1}{4}(l_2 + l_5)^2 \right] - \frac{1}{16} \\ &= \frac{1}{16} - \frac{1}{8} [(l_1 + l_3 + l_4)(l_2 + l_5) + (l_2 + l_4)(l_1 + l_3 + l_5)] = \frac{1}{8} \left( \frac{1}{2} - \bar{r}^\sigma \right), \end{aligned}$$

which is Equation 2.

variance of bp vs. cM genetic relatedness to grandparents (the mathematical details of this effect are explained in File S1, Section S3). Because of this effect, although the chromosome-specific variances of bp genetic relatedness to grandparents are substantially smaller than their cM counterparts, the genome-wide variances of bp and cM genetic relatedness are more similar (Table 1).

### Indirect Relationships

Indirect relationships involve two descendants of at least one individual in the pedigree. In the case of multiancestor pedigrees, we restrict our attention to two-ancestor pedigrees where the two ancestors were a mating pair (so that the focal descendants are, for example, full siblings, or aunt-nephew, etc.). We focus here on half-siblings and full-siblings; the calculations for general indirect relationships of this kind are given in File S1, Section S2.

#### Half-siblings

Let the random variable  $IBD_{\text{h-sib}}$  be the proportion of two half-siblings' genomes that they share IBD, if they have the same father but unrelated mothers. Then  $\mathbb{E}[IBD_{\text{h-sib}}] = 1/4$  is the coefficient of relationship, and

$$\text{Var}(IBD_{\text{h-sib}}) = \frac{1}{8} \left( \frac{1}{2} - \bar{r}_{(2)}^\sigma \right), \quad (3)$$

where  $\bar{r}_{(2)}^\sigma$  is the probability that a randomly chosen locus pair recombines when the crossovers of two of the father's meioses are pooled into one hypothetical meiosis (see Figure 2 for an example of a pooled meiosis). If the common parent were instead the mother,  $\bar{r}_{(2)}^\sigma$  would replace  $\bar{r}_{(2)}^\sigma$ . A graphical demonstration of Equation 3, based on the possible segregation patterns of two meioses in the parent, is given in Figure 2.

**Siblings:** Let the random variable  $IBD_{\text{sib}}$  be the proportion of two full-siblings' genomes that they share IBD, assuming their mother and father to be unrelated. Then  $\mathbb{E}[IBD_{\text{sib}}] = 1/2$  is the coefficient of relationship, and

$$\text{Var}(IBD_{\text{sib}}) = \frac{1}{8} \left( 1 - \bar{r}_{(2)}^\sigma - \bar{r}_{(2)}^\sigma \right). \quad (4)$$

Like  $\bar{r}$ ,  $\bar{r}_{(2)}$  can be estimated from various kinds of data, including cytological data of crossover positions at meiosis I, sequence data from gametes, and linkage maps. Table 1 lists the chromosome-specific and genome-wide standard deviations of bp genetic relatedness of human siblings, calculated using cytological data from Lian et al. (2008) for male meiosis and the linkage map of Kong et al. (2010) for female meiosis (with map distances converted to recombination rates using Kosambi's map function). Also shown are the corresponding standard deviations of cM genetic relatedness

**Table 1** Standard deviations of genetic relatedness to a paternal and maternal grandparent, and to a sibling, in humans, for both bp and cM measures of genetic relatedness

Chromosome	Grandparent				Sibling	
	Paternal		Maternal			
	bp <sup>a</sup>	cM <sup>b</sup>	bp <sup>c</sup>	cM <sup>b</sup>	bp <sup>d</sup>	cM <sup>b</sup>
1	0.145	0.156	0.115	0.126	0.146	0.147
2	0.152	0.161	0.117	0.130	0.152	0.152
3	0.164	0.168	0.126	0.138	0.165	0.162
4	0.171	0.174	0.127	0.139	0.170	0.166
5	0.173	0.176	0.128	0.142	0.173	0.169
6	0.183	0.181	0.133	0.145	0.184	0.175
7	0.179	0.178	0.135	0.148	0.181	0.176
8	0.187	0.184	0.139	0.152	0.192	0.183
9	0.186	0.186	0.149	0.157	0.199	0.188
10	0.184	0.182	0.139	0.152	0.188	0.181
11	0.187	0.189	0.142	0.157	0.193	0.190
12	0.179	0.182	0.140	0.154	0.183	0.183
13	0.177	0.192	0.152	0.170	0.192	0.204
14	0.175	0.195	0.160	0.178	0.195	0.214
15	0.180	0.197	0.154	0.172	0.196	0.209
16	0.190	0.194	0.155	0.169	0.207	0.204
17	0.193	0.195	0.150	0.168	0.204	0.204
18	0.198	0.201	0.158	0.173	0.213	0.213
19	0.199	0.203	0.178	0.182	0.226	0.223
20	0.195	0.211	0.167	0.184	0.213	0.231
21	0.198	0.219	0.189	0.205	0.235	0.260
22	0.201	0.218	0.188	0.205	0.236	0.259
Genome	0.040	0.040	0.031	0.034	0.041	0.040

<sup>a</sup> Calculated from cytological data of Lian *et al.* (2008).

<sup>b</sup> Calculated from formulas in Hill and Weir (2011), using chromosome map lengths of Kong *et al.* (2010). Does not take into account crossover interference. cM relatedness to paternal and maternal grandparents defined, respectively, in terms of the male and female map; cM relatedness of siblings defined in terms of the sex-averaged map.

<sup>c</sup> Calculated from linkage maps of Kong *et al.* (2010) using Kosambi's map function.

<sup>d</sup> Calculated from cytological data of Lian *et al.* (2008) (male meiosis) and linkage maps of Kong *et al.* (2010) using Kosambi's map function (female meiosis).

of siblings, defined as the proportion of the sex-averaged genetic map that they share IBD.

As for the case of genetic relatedness to grandparents, several meiotic features affect the comparison of the variances of bp and cM genetic relatedness of siblings. First, the bp variances are increased by the proterminal distribution of crossovers along chromosomes in humans, which tends to decrease  $\bar{r}_{(2)}$ . Thus, when the variance of bp genetic relatedness of siblings is calculated using linkage maps and Haldane's map function (to eliminate the effect of crossover interference), the chromosome-specific and genome-wide estimates are substantially larger than the corresponding cM variances (File S1, Section S4). However, crossover interference, by increasing genetic shuffling, increases  $\bar{r}_{(2)}$ , thus decreasing the bp variance. These opposing effects of proterminal localization of crossovers and crossover interference roughly cancel in this case, so that our estimates of the variance of bp and cM genetic relatedness of siblings are coincidentally similar, at both the chromosome-specific and genome-wide levels (Table 1).

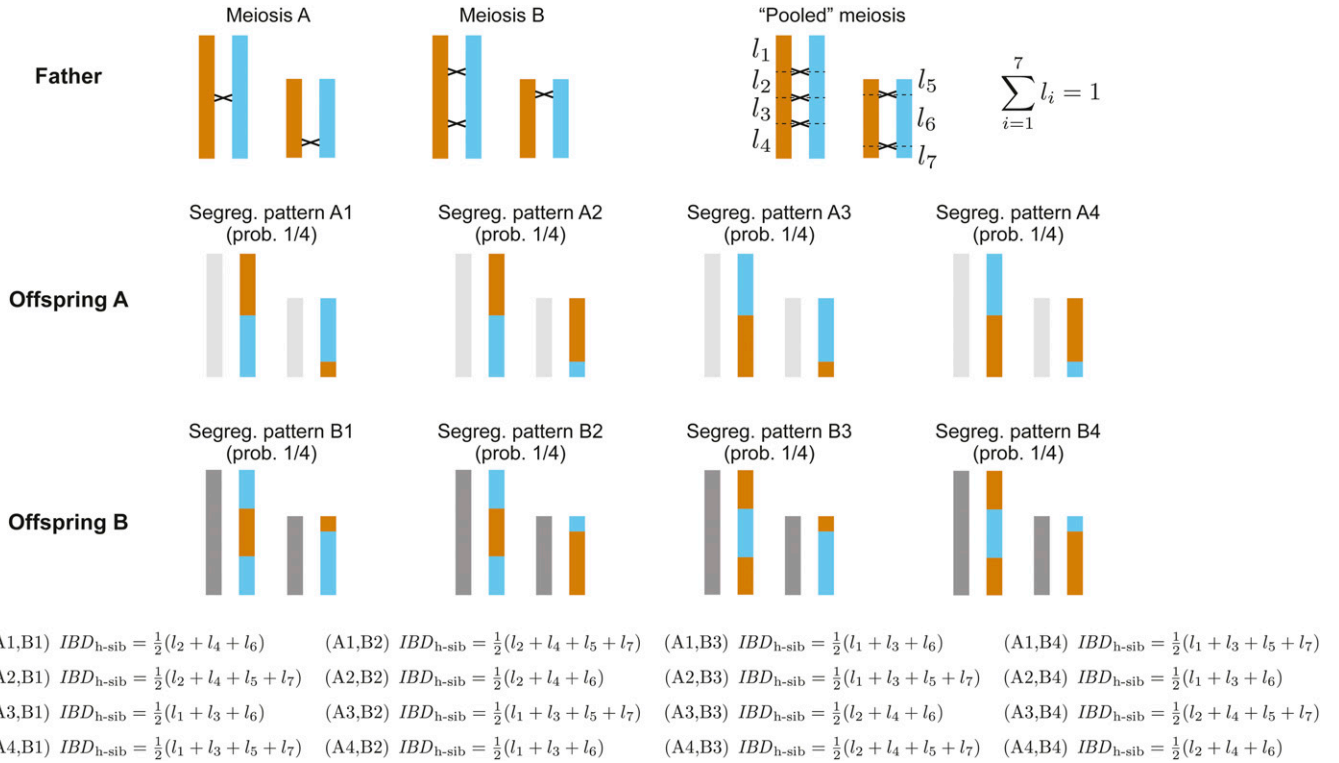
### Within- vs. cross-pedigree variance

The calculations above and in Appendices S1 and S2 are for the variance of genetic relatedness in a given instance of a specified pedigree relationship. This variance derives from the randomness of recombination and segregation in the meiotic processes of the individuals involved in that particular pedigree. For some applications, however, we are interested in the variance of genetic relatedness across instances of a specified pedigree relationship [*e.g.*, using variation in the genetic relatedness of different sibling pairs to estimate the heritability of some trait (Visscher *et al.* 2006)]. To calculate this 'population variance' of genetic relatedness, variation across individuals in their recombination processes must be taken into account. Applying the law of total variance (details in File S1, Section S5), we find that the variance of genetic relatedness across instances of a specified pedigree relationship is equal to the average within-pedigree variance. We have shown that within-pedigree variances are functions of metrics of aggregate recombination such as  $\bar{r}$  and  $\bar{r}_{(2)}$ ; to calculate the cross-pedigree variance, these metrics must simply be averaged across pedigrees.

A complication arises when using pooled recombination data (such as linkage maps) to estimate the cross-pedigree variance of genetic relatedness, because for all such metrics of aggregate recombination except  $\bar{r}$ , calculation of the metric from averaged recombination data does not return the average of the metric across pedigrees (File S1, Section S5). It is therefore technically invalid, in such cases, to use pooled recombination data to calculate the cross-pedigree variance of genetic relatedness (although it is valid in the case of grand-offspring-grandparent).

To get a sense for how large an error the use of pooled recombination data can cause, we focus on the case of paternal half-siblings. Using crossover data generated by Bell *et al.* (2020) by single-cell sequencing of large numbers of sperm from 20 human male donors, we calculated values of  $\bar{r}_{(2)}$  for each individual donor, from which we calculated a value of  $\bar{r}_{(2)}$  averaged across individuals. We also calculated a value of  $\bar{r}_{(2)}$  from recombination rates that were averaged across individuals. The values of  $\bar{r}_{(2)}$  from both individual and pooled recombination rates were calculated genome-wide and per chromosome. Using the two estimates of  $\bar{r}_{(2)}$ , we calculated the variance of genetic relatedness of paternal half-siblings according to Equation 3. We found that the values based on pooled recombination fractions differed only slightly from the correctly calculated values: the genome-wide variances differed by about 0.25%, and the chromosome-specific variances differed by comparable amounts. Details of these calculations are given in File S1, Section S5.

Therefore, the bias introduced by using linkage maps to calculate the population variance of genetic relatedness is likely to be small. Nevertheless, when available, disaggregated data in which crossover positions are inferred for individual nuclei (*e.g.*, cytological data for individual meiocytes



**Figure 2** The variance of genetic relatedness between half-siblings, calculated from the possible segregation patterns of two meioses of their common father. The positions of crossovers in the two paternal meioses (and the chromatids involved) are specified, but the segregation patterns in the resulting sperm cells (and therefore the two offspring) are not. Applying Equation 1 in Veller *et al.* (2019b) to the "pooled meiosis" in which the crossovers from the two actual meioses have been combined, we find  $\bar{r}_{(2)}^\sigma = (l_1 + l_3 + l_5 + l_7)(l_2 + l_4 + l_6) + (l_1 + l_3 + l_6)(l_2 + l_4 + l_5 + l_7)$ . Across the 16 possible segregation patterns  $(A_i, B_j)$ ,  $\mathbb{E}[IBD_{h-sib}] = 1/4$  and

$$\begin{aligned} \text{Var}(IBD_{h-sib}) &= \mathbb{E}[IBD_{h-sib}^2] - (\mathbb{E}[IBD_{h-sib}])^2 \\ &= \frac{1}{16} \left[ (l_1 + l_3 + l_5 + l_7)^2 + (l_2 + l_4 + l_6)^2 + (l_1 + l_3 + l_6)^2 + (l_2 + l_4 + l_5 + l_7)^2 \right] - \frac{1}{16} \\ &= \frac{1}{16} - \frac{1}{8} [(l_1 + l_3 + l_5 + l_7)(l_2 + l_4 + l_6) + (l_1 + l_3 + l_6)(l_2 + l_4 + l_5 + l_7)] = \frac{1}{8} \left( \frac{1}{2} - \bar{r}_{(2)}^\sigma \right), \end{aligned}$$

which is Equation 3.

or sequencing data for individual gametes) are preferable for calculating cross-pedigree variances.

### Application: Ancestry Variance and Selection Among $F_2$ s

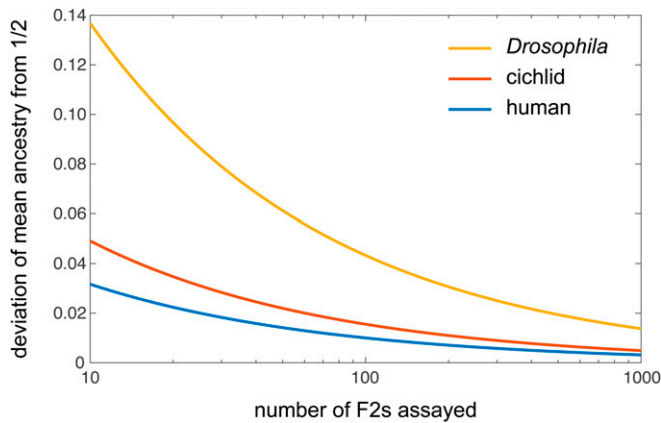
A common experimental design involves mating individuals from two lines, populations, or species (A and B) to form a hybrid "F<sub>1</sub>" generation, and then mating F<sub>1</sub>s to produce an F<sub>2</sub> generation. Every F<sub>1</sub> carries exactly half of its DNA from each species, but there is ancestry variance among F<sub>2</sub>s because of recombination and segregation in the F<sub>1</sub>s' meioses (Hill 1993a).

Each F<sub>2</sub> derives from an F<sub>1</sub> mother's egg and an F<sub>1</sub> father's sperm. Let the random variables  $\hat{P}^\varnothing$  and  $\hat{P}^\sigma$  be the respective proportions of species-A DNA in the egg and sperm (measured

in bp), and let  $P$  be the proportion of species-A DNA in an F<sub>2</sub>'s genome. Then  $P = (\hat{P}^\varnothing + \hat{P}^\sigma)/2$ , and, from Equation 1,  $\text{Var}(\hat{P}^\varnothing) = \frac{1}{2}(\frac{1}{2} - \bar{r}^\sigma)$  and  $\text{Var}(\hat{P}^\sigma) = \frac{1}{2}(\frac{1}{2} - \bar{r}^\sigma)$ . Finally, because  $\hat{P}^\varnothing$  and  $\hat{P}^\sigma$  are independent, the ancestry variance among F<sub>2</sub>s is

$$\text{Var}(P) = \frac{1}{4} \left( \text{Var}(\hat{P}^\varnothing) + \text{Var}(\hat{P}^\sigma) \right) = \frac{1}{8} (1 - \bar{r}^\varnothing - \bar{r}^\sigma). \quad (5)$$

[If the F<sub>2</sub>s instead derived from a backcross of F<sub>1</sub>s to one of the parental species, the ancestry variance among F<sub>2</sub>s would be  $\frac{1}{8}(\frac{1}{2} - \bar{r})$ , with  $\bar{r}$  calculated for the sex of the F<sub>1</sub>s involved. The corresponding variance for cM-based ancestry of backcross F<sub>2</sub>s, and those in later backcross generations, have been calculated by Hill (1993a), assuming no crossover interference.]



**Figure 3** The minimum deviation of mean ancestry among  $F_2$ s that can be statistically distinguished from the null expectation of  $1/2$  at the 5% significance level.

The calculation above assumes that there is no systematic selection among  $F_2$ s in favor of alleles from one of the two species, and no meiotic drive in  $F_1$ s, both of which would shift the distribution of ancestry among  $F_2$ s toward one of the two species. For example, Matute *et al.* (2020) generated two crosses, each between one widely distributed species of *Drosophila* and one closely related island endemic. In the resulting admixed populations, island ancestry was replicably selected against over time. If viability selection plays a role in this effect, then an ancestry skew toward the widespread species would be expected among adult  $F_2$ s in these crosses.

An  $F_2$ -based test for selection of this kind would involve comparing the observed average ancestry among  $F_2$ s against the neutral null expectation of  $1/2$ . In this case, Equation 5 gives the appropriate null variance for the purpose of statistical inference; the standard error of the test is

$$SE = \frac{1}{\sqrt{8n}} \sqrt{1 - \bar{r}^{\varphi} - \bar{r}^{\sigma}}, \quad (6)$$

where  $n$  is the sample size of  $F_2$ s for which ancestry proportions have been assayed.

Substituting known values of  $\bar{r}$  into Equation 6 then shows, for a given sample of  $F_2$ s, how much their average ancestry proportion must deviate from  $1/2$  for us to reject the null hypothesis of neutrality. For example, using a linkage map generated from a cross of two closely related cichlid fish species (Feulner *et al.* 2018), together with Kosambi's map function, we calculate a sex-averaged value of  $\bar{r} = 0.4749$ . If ancestry fractions were measured for 10  $F_2$ s from this cross, then Equation 6 tells us that a 4.9% or greater deviation of the average ancestry from the null expectation of 50% would be statistically distinguishable at the 5% significance level; if 100  $F_2$ s were assayed, the threshold detectable deviation would be 1.6%. Threshold deviations for a range of sample sizes are shown in Figure 3 for the recombination processes of cichlids, humans, and *Drosophila melanogaster*. It is clearly

seen that, because of *D. melanogaster*'s low sex-averaged value of  $\bar{r} = 0.305$  (see Discussion), much greater ancestry deviations among  $F_2$ s are required for the null hypothesis of neutrality to be rejected, compared to cichlids and humans.

Note that the test described above can also be carried out for a specific genomic region of interest, by using region-specific values of  $\bar{r}$ . Alternatively, in a genome-wide scan, regions of the genome where ancestry deviations are particularly large can be statistically identified using region-specific values of  $\bar{r}$ , correcting for multiple hypothesis testing.

#### Data availability

The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article. Supplemental material available at figshare: <https://doi.org/10.25386/genetics.13138577>.

#### Discussion

Relatives of a given pedigree relationship vary in how much of their DNA they share IBD, because of variable patterns of recombination and chromosome segregation in their pedigrees. Previous theoretical calculations of the variance of genetic relatedness have been limited to measuring the genetic relatedness of two individuals as the proportion of the total genetic map (in cM) that they share IBD. Such calculations have failed to accommodate crossover interference and sex differences in recombination, both of which are near-universal features of meiosis that are known to substantially affect variance in genetic relatedness (Caballero *et al.* 2019).

Here, we have shown that when genetic relatedness is instead measured as the proportion of the total physical genome (in bp) shared IBD by relatives, the variance of genetic relatedness is determined by aggregate recombination, as quantified by  $\bar{r}$  and analogous metrics. These metrics take into account all features of the aggregate recombination process, including the number of chromosomes and heterogeneity in their size, the number of crossovers and their spatial organization along the chromosomes, and sex differences in recombination (Veller *et al.* 2019b). In addition to incorporating the above meiotic features into theoretical calculations of the variance of genetic relatedness, our treatment further allows these features' effects on genetic relatedness to be understood intuitively in terms of their effects on aggregate genetic shuffling. Several examples are discussed below.

#### Sex differences in recombination

In many species, male and female meiosis differ both in the number and location of crossovers [reviewed by Lenormand and Dutheil (2005); Sardell and Kirkpatrick (2020)]. In human male, there are fewer crossovers and they are more terminally localized along the chromosomes than in human female. Both factors decrease the total amount of genetic

shuffling in male meiosis (Veller *et al.* 2019b), providing an intuitive explanation of the observation of Caballero *et al.* (2019) that, in humans, relatives who are related predominantly via males have a higher variance of genetic relatedness than relatives related predominantly via females. In our calculations, for example, the standard deviation of genetic relatedness to a paternal grandparent is about 30% greater than that to a maternal grandparent (Table 1).

Such effects will be especially pronounced in species with no crossing over in one sex (*e.g.*, male *Drosophila* and female Lepidoptera). Using chromosome lengths from Release 6 of the *D. melanogaster* reference genome (Hoskins *et al.* 2015) and the female linkage map produced by Comeron *et al.* (2012) (together with Kosambi's map function), we calculate autosomal values of  $\bar{r}^{\sigma} = 0.253$  and  $\bar{r}^{\varphi} = 0.358$ . Substituting these values into Equation 2, we find that the standard deviations of (autosomal) genetic relatedness to paternal and maternal grandparents are 0.175 and 0.125, respectively—a difference of 40%.

### Chromosome number and size

Because most genetic shuffling in meiosis is due to independent assortment of chromosomes rather than crossing over (Crow 1988; Veller *et al.* 2019b), the most important contributor to cross-species differences in the variance of genetic relatedness is karyotypic differences. Two karyotypic features affect genetic shuffling, and therefore the variance of genetic relatedness.

First, the greater the number of chromosomes, the greater the genetic shuffling associated with their independent assortment. For example, humans have 22 autosomes while *D. melanogaster* has only two major autosomes. Therefore, the contribution of independent assortment to  $\bar{r}$  [and, equivalently, to  $\bar{r}_{(2)}$ ] is large in humans (0.473) but small in *D. melanogaster* (0.253). Thus, were there no crossing over in either species, the SD of the genetic relatedness of full siblings would be about 8% in humans and 25% in *D. melanogaster* (Equation 4).

Second, the more homogeneously sized the chromosomes are, the greater the genetic shuffling associated with their independent assortment. We have used this fact to explain why independent assortment of chromosomes in humans is more effective at decreasing the variance of cM vs. bp genetic relatedness, because the chromosomes are more homogeneously sized when measured in cM.

### Crossover positions

White and Hill (2020) have shown that terminal placement of crossovers tends to increase the variance of genetic relatedness, relative to more central placement of crossovers. This can be explained by the intuitive fact that a crossover near the tip of a chromosome causes less genetic shuffling than a crossover in the middle (Veller *et al.* 2019b). We have used this fact to explain why, after controlling for the effects of crossover interference, per-chromosome variances of genetic

relatedness are smaller for bp than for cM genetic relatedness in humans, because crossovers are proterminally distributed along the physical chromosome maps but, by definition, are uniformly distributed along the genetic maps.

### Crossover interference

It has recently been shown, by computer simulation of various forms of crossover patterning along chromosomes, that crossover interference tends to decrease the variance of genetic relatedness between relatives (Caballero *et al.* 2019). Veller *et al.* (2019b) demonstrated that interference among crossovers increases the amount of genetic shuffling that they cause (increasing  $\bar{r}$  and analogs). The intuition is that, when two crossovers occur very close to each other along a bivalent chromosome at meiosis I (the stage at which crossover interference operates), they cancel each other's effect on genetic shuffling, together behaving more like a single crossover. Such "stepping on toes" is prevented by crossover interference, thus increasing genetic shuffling. This provides an intuitive explanation of the result of Caballero *et al.* (2019).

Using a simulation method employed by Mancera *et al.* (2008) and Wang *et al.* (2012) to resample empirically observed crossovers in an interference-less way, Veller *et al.* (2019b) calculated that, in human male, interference among crossovers increases their contribution to  $\bar{r}$  by about 15%. By this measure, crossover interference in human male meiosis decreases the genome-wide standard deviation of genetic relatedness to a paternal grandparent from 0.043 to 0.040 (Equation 2), a decrease of about 7%.

### Crossover covariation

It has recently been shown for diverse eukaryotes that the number of crossovers per chromosome covaries positively across chromosomes within individual meiotic nuclei (Wang *et al.* 2019). This "crossover covariation" substantially increases the variance of crossover number per gamete, which will clearly affect the distribution of genetic relatedness among relatives. However, because crossover covariation does not change the (unconditional) probability that a given pair of loci are recombinant in a gamete, it does not alter  $\bar{r}$  or analogs (since these are averages of functions of individual pairwise recombination rates—see Appendices S1, S2, and S5). Therefore, crossover covariation does not affect the variance of genetic relatedness among relatives (but it will affect higher-order moments).

### Conclusion

We have shown that the variance of genetic relatedness is a function of  $\bar{r}$  and analogous metrics. Since these metrics can readily be estimated from modern cytological and sequencing data [see Veller *et al.* (2019b) and above], our results make it possible to calculate the variance of genetic relatedness in a precise, general, and unambiguous way.



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