Ancestry Informative Markers: Getting a Lot Out of a Little

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A B S T R A C T

The SNP markers have emerged as essential resources for performing genetic linkage, association, and admixture studies. Both academic and commercial groups are developing large numbers of genome-wide SNP datasets. These databases are utilised for several studies involving both structural and functional genomics. Some studies viz. admixture mapping, population structure identification and quantifying genetic diversity requires a genome-wide small informative panel of relatively evenly spaced markers. These informative markers can distinguish the ancestral origins of chromosomal segments in admixed individuals and can delineate the ancestral origin of a particular population. Thus present study reviews existing methods to select ancestry informative markers and their comparative efficiency in inferring individual ancestry level.

Key words

Ancestry informative markers, Admixture, SNP

Introduction

Inference of individual ancestry from genetic markers is helpful in diverse situations, including admixture and association mapping, forensics, prediction of medical risks, wildlife management, food safety and studies of dispersal, gene flow, and evolutionary history (Shriver et al., 1997; Davies et al., 1999; Primmer et al., 2000; Manel et al., 2002; Bamshad et al., 2003; Campbell et al., 2003; Ziv and Burchard 2003). Most of the statistical methods developed for tracing ancestry use multi-locus genotypes and allele frequencies of the population, which is either specified in advance or estimated subsequently (Smouse et al., 1982; Paetkau et al., 1995; Rannala and Mountain 1997; Cornuet et al., 1999; Pritchard et al., 2000; Guinand et al., 2002). The use of highly informative markers sometimes can reduce the genotyping rate, so, it is appropriate to measure the extent to which specific markers contribute to the ancestry inference.

The term ancestry refers to the proportion of genetic material that transcends each founder population. Ancestry-informative markers (AIMs) are those markers that can specify the likelihood of origin of a population from the...
DNA sample where the source individual is not known or is unable to proclaim their ancestry (Phillips et al., 2007). Autosomal AIMs are mainly used for studying admixture and inferring individual biogeographical ancestry (I-BGA) (Halder et al., 2008). Availability of Different densities of SNP BeadChips has led us to generate a tremendous amount of genotypic datasets. Geographic ancestry can be easily extrapolated from this genotypic data produced through SNP genotyping (Rosenberg et al., 2002; Tang et al., 2005; Paschou et al., 2007). While inferring genome-wide scans, most of the markers are found to be common and vary only to some extent among different populations. Markers which are present homogeneously within the population and possess highly differentiated frequencies are very informative for ancestry estimation (Shriver et al., 2003). Basically, there are two approaches for studying genetic admixture and ancestry, i.e., model-based (Bayesian) (Rosenberg et al., 2002; Pritchard et al., 2000) and likelihood approach (based on the allele frequencies of the admixed population) (Liu et al., 2013; Skotte et al., 2013). However, the Bayesian approach and Principal component analysis (PCA) have been the main tools of choice for studying population structure, ancestry, and diversity (Paschou et al., 2007).

**Number of AIMs required to study population structure**

The informativeness of the markers decides the number of markers needed for studies on admixture levels, population structure or genetic diversity (Ding et al., 2011). According to one study, more number of biallelic markers (SNPs) (4-10X) are required to achieve the same results in terms of efficiency and resolution as that of multi-allelic markers (microsatellites) in population structure and diversity studies (Morin et al., 2004). This problem can be addressed by dimension reduction of variables using PCA. Lewis et al., (2011) have elucidated that in most of the cases the number of genetic markers required for ancestry inference can be reduced to 1.5% of the original number of SNPs with 92% accuracy. In the study of PCA correlated SNPs for structure identification in nine human populations (Paschou et al., 2007) distributed worldwide, 50 PCA-correlated SNPs can assign the individuals to their population of origin with 100% accuracy. Recently, Getachew et al., (2017) have identified top ranked 74 AIMs on the basis of FST which could predict the admixture level in crossbred populations of Menz x Awassi and Wollo x Awassi Sheep. In an admixed population of Swiss Fleckvieh cattle, Frkonja et al., (2012) selected 48 and 96 top ranked SNPs with FST > 0.651 and 0.623, respectively and found a correlation of 0.907 and 0.924 with admixture level estimated through pedigree method. Ding et al., (2011) obtained 100 to 200 top informative SNPs using Fisher Information Content (FIC), Shannon Information Content (SIC), F statistics (FST), Informativeness for Assignment Measure (I_n), and Absolute Allele Frequency Differences (delta, δ). They observed that each method generated similar estimates of ancestry contribution in CEU and YRI admixed population. Kavakiotis et al., (2017) recently developed a method i.e., Frequent Item Feature Selection (FIFS), which was tested on a dataset of pigs. This dataset consisted of 446 individuals belonging to 14 sub-populations, genotyped at 59,436 SNPs. The SNPs selected by different methods were FIFS: 28 SNPs, Delta: 70 SNPs, Pairwise FST: 70 SNPs, I_n: 100 SNPs. Their approach successfully dealt with the problem of informative marker selection in high dimensional genomic datasets.

**Methods to find AIMs**

There are basically two different approaches. One is to estimate the assignment power of
each individual loci using available software (WHICHLOCI, Banks et al., 2003) or combinations of the different software (i.e. GAFS, Topchy et al., 2004 and BELS, Bromaghin, 2008). All these software possess few advantages and disadvantages (Helyar et al., 2011), but their primary requirement is the computational efficiency. As Helyar et al., (2011) mentioned, although, no limitations exists for the number of loci or individuals to be included in analysis, but it may be prohibitive on a desktop.

A second approach is to rank loci solely according to their informativeness, that is, the marker information content, which is the amount of information that a locus holds regarding the ancestry of an individual. The use of markers with high informativeness reduces the number of markers needed for correct assignment (Rosenberg et al., 2003). Several measures/criteria of marker informativeness have been proposed (Rosenberg et al., 2003; Ding et al., 2011), such as Delta (Shriver et al., 1997), pairwise Wright’s $F_{ST}$ given by Wright (1951), global Wright’s $F_{ST}$ by Wright (1951), pair wise Weir and Cockerham $F_{ST}$ by Weir and Cockerham (1984), global pair wise Weir and Cockerham $F_{ST}$ by Weir and Cockerham (1984), and informativeness for assignment ($I_{a}$) (Rosenberg et al., 2003). The application of different methods for marker prioritization and decision making in the construction of SNP panels is becoming more important as large high-throughput assays become readily available. A novel data mining approach, called FIFS, based on the use of frequent items for selection of the most informative markers from population genomic data has also been developed by Kavakiotis et al., (2017). It has two main components. For each sampled population the first component detects the most unique and frequently occurring genotypes while the second one picks the most appropriate among them and return the informative SNP subsets. TRES: Toolbox for Ranking and Evaluation of SNPs is another tool which is used for ranking markers and selecting set of SNPs on the basis of Wright’s $F_{ST}$, informativeness for assignment ($I_{a}$), and Absolute allele frequency difference ($\delta$) (Kavakiotis et al., 2015). Existing approaches to select a subset of informative markers entails prior knowledge of the ancestry of individuals included in the study. PCA, a powerful dimensionality reduction technique is an emerging approach to identify a small panel of informative markers. Paschou et al., (2007) developed a novel algorithm based on PCA that does not depend on any prior assumptions, which can be applied to datasets of hundreds of individuals and millions of markers.

**Methods to measure marker information content**

**Absolute allele frequency difference ($\delta$):** For a biallelic locus, suppose allele one is the reference allele, then, $\delta=|p11−p21|$. A marker with $\delta=1$ gives complete information about ancestry while a marker with $\delta=0$ carries no information. In an admixture model, this is related to the amount of linkage disequilibrium (Chakraborty and Weiss 1988); whereas in a multilocus no-admixture model this is related to the probability of correct assignment (Risch et al., 2002); Fisher information curvature criterion and ORCA for K=2 (Rosenberg et al., 2003). But it is optimum when only two populations are possible sources; and also does not take into account all available information about allele frequencies (Stephens et al., 1999; Campbell et al., 2003). Its statistical features also do not apply to the multi-allelic extension of $\delta$ (Shriver et al., 1997).

$F_{ST}$: This is a measure of population substructure and genetic divergence among subpopulations. It is excess in the probability of identity of alleles from the same population compared with randomly chosen alleles.
\[ F_{ST} = \frac{(p_{ij} - p_{2j})^2}{(p_{ij} + p_{2j})(2 - (p_{ij} + p_{2j}))} \]

Here, \( j = 1 \) or \( 2 \) is the reference allele. Values of \( F_{ST} \) can range from 0 to 1. A high \( F_{ST} \) value implies a considerable degree of differentiation between populations. This is related, for biallelic markers, to the quotient of expected posterior and prior variance of ancestry in a population equally admixed from two sources (McKeigue 1998; Molokhia et al., 2003). But it performs only slightly better than random markers (Rosenberg et al., 2001).

**Expected heterozygosity:** It performs better than random markers (Rosenberg et al., 2001). It is good at measuring the amount of variation but not the differences across populations.

**The number of alleles:** It measures the amount of variation but not the differences across populations, hence is useful only for multi-allelic markers that have variation in the number of alleles such as microsatellites.

**Fisher information curvature criterion:** It enables prediction about approximate variances of ancestry estimates and the information matrix is additive across loci that are independent within populations.

It depends on unknown ancestry coefficients and requires computation for many possible parameter values; largest eigenvalue gives an upper bound that might not be generally applicable across the parameter space (Rosenberg et al., 2003).

**Shannon information criterion:** The concept of entropy can be used to develop a measure of marker informativeness (Rosenberg et al., 2003). Entropy is a measure of the uncertainty associated with a random variable and quantifies the expected information content contained in the data.

**Pairwise Kullback-Leibler divergence:** For \( K=2 \), it possesses an average potential and multi-locus extension, for assignment of an allele to one population. Contribution of specific alleles can also be measured. It requires that only two populations are possible sources and in small samples the estimate is upwardly biased (Rosenberg et al., 2003).

**Informativeness for assignment (**\( I_n **)): This measure was produced by Rosenberg et al., (2003). It potentially allocates an allele to one population and possess a natural multi-locus extension. It also facilitates measurement of specific allele contribution. Its ancestry assignment is better than random or highly heterozygous markers. But in small samples the estimate is upwardly biased.

**Informativeness for ancestry coefficients (**\( I_a **)): this was also formulated by Rosenberg et al., (2003). It is similar to \( I_n \) in statistical features, but in samples with populations of equal sample size computational efficiency decreases.

**Optimal rate of correct assignment (**ORCA **):** Gives the probability of correct assignment of an allele using the decision rule with lowest risk; has a natural multi-locus extension; enables measurement of contributions of specific alleles. But in small samples estimates are upwardly biased (Rosenberg et al., 2003).

**Comparison of different methods**

The simulation study conducted by Ding et al., (2011) revealed that \( I_n \) was the best as compared to \( \delta \), \( F_{ST} \), FIC, and SIC in selecting the set of AIMS giving the least bias and mean square error in ancestry estimation. FIC had the least overlap with other measures in selecting markers. \( F_{ST} \), FIC, and \( I_n \) gave relatively smaller and similar marker panels,
whereas SIC gave a very small panel of AIMs. FIC and SIC were more alike in picking a set of SNPs, while $\delta$, $F_{ST}$, and $I_n$ were more likely to pick the same set of SNPs, and FIC was more likely to choose SNPs that were not chosen by the other measures. FIC and SIC measures necessitate a pre-defined ancestral proportions in an admixed population, whereas $F_{ST}$, $\delta$, and $I_n$ do not. Combined information from more than one method may provide a reliable means, in selecting markers for ancestry inference (Ding et al., 2011).

Use of AIMs in detecting meat adulteration and mislabelling

Wilkinson et al., (2012) developed a panel of 96 SNPs that has the ability to authenticate pork products labelled with traditional breed names and thus expose mislabelled products. Assignment of individuals to particular breeds was extremely accurate for traditional breeds. When they studied 40 market samples with the panel of 96 SNPs, observed that 2 samples not assigned to claimed breed origin but assigned to another breed, indicating possibly mislabelled meat. The assignment to traditional breeds was also accurate for processed pork product though genotyping rate fell to 88%. Similarly, Orrù et al., (2009) found 18 SNPs, those are useful in tracing beef to the most consumed breeds in the markets of Italy. Suekawa et al., (2010) developed five breed specific DNA markers to correctly discriminate between Japanese and imported cattle for food safety based on bovine 50K SNP array.

Although millions of SNPs have been identified through which population structure, admixture, traceability and diversity is studied, the number of markers can be reduced to minimal for the same purpose based on their information content. The small panel of ancestry informative markers may greatly reduce the genotyping cost while retaining the power for assigning individuals to accurate ancestral populations. A choice of method to select AIMs, with least biasness and root mean square error plays an important role in admixture studies which in turn avoids false positive genotype-phenotype associations. The set of informative markers are also useful in meat traceability for food safety, and checking mislabelling of the meat products. The choice of method mostly depends upon the number of populations in the dataset, sample size, type of marker and number of loci involved, but we suggest a combination of all these methods to measure the marker information content and to select the desired small AIM panel accordingly.

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