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Abstract

We here describe a method to generate phylognies from few yet hyperpolymorphic genes. This is useful for which large collections of allel probabilities for all encountered allele types according to observed allele frees

Bootstrapping phylogenies with allele frequencies for few hyperpolymorphic loci

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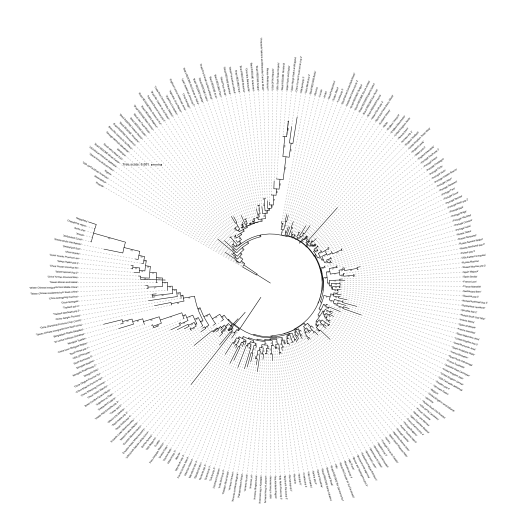
1 Introduction

Polymorphic genes of the MH complex have properties not unlike other genetic markers suitable for population studies, such as There is a historic basis for using HLA gene frequencies in population studies. Mainly, there are six highly polymorphic loci to choose between that have different frequencies between individuals and different populations. Class II genes appear to be older than the class I genes[citation needed]. Although there is some trans-species HLA gene flow between human and apes, trans speciation is relatively minor and loci dependent. Thus, most human HLA alleles, although old, have probably evolved by genetic drift, selection and mutation with gene flow and admixture contributing to the polymorphisms in the MHC region since the split between apes and humans.

It is also claimed that HLA is under selection, which raise concerns

Nonetheless, hyperpolymorphic genes such as HLA class I and II are frequently used in anthropological and population genetic studies. HLA typing has its roots in serology and thus experimentally, large body of data is available. Because of their enormous polymorphism and differences between individuals and populations, the phylogenies of HLA genes are in theory highly informative for population genetics and anthropology. From an information theoretic perspective they are a promising research target, as the amount of all possible outcomes However, current methods are based on allele frequencies alone and do not leverage the information theoretic richness during bootstrapping: Bootstrapping is based on random sampling with replacement from the original input. To the best of our knowledge, all phylogenetic toolkits that are capable of dealing with allele frequencies sample gene-wise. I.e. either a gene is sampled and all its allele frequencies (in the case of HLA genes, this can be as high as ...) are considered or the gene is discarded in a bootstrap and then none of its allele frequences are taken into account. This all-or-nothing approach works well when many loci are considered (e.g. microsatellites) but for few genes, it leads to a very coarse grained information entropy resolution.

Rich databases describing Therein, populations are described in terms of allele frequencies (see allelefrequenciens.net paper) Unfortunately, the low num-



ber of genes as biomarkers in combination with phylogenetic tools that expect high numbers of loci for bootstrapping have led to numerous works with incorrect bootstrapping. They providing false confidence into produced phylogenetic trees. The ramifications of such practices have caused severe controversies: In 2002, Piazza, Risch and Cavalli-Sforza [1] have already criticized the works of Hajjej/Arnaiz-Villena and noted that phylogenetic analysis "Using results from the analysis of a single marker, particularly one likely to have undergone selection, for the purpose of reconstructing genealogies is unreliable and unacceptable practice in population genetics"

Nonetheless, a large number of HLA-based phylogeny reconstruction publications have since been published based on one or very few marker genes [2, 3, 4, 5]. It is worth noting that HLA genes are hyperpolymorphic, and as such many columns in respective MSAs would carry information entropy for population genetics studies. Most tree reconstruction methods assume locus independence ([6]), which yet requires a rigorous examination. The problem is not necessarily the low number of gene markers as long as they are hyperpolymorphic (i.e., respectively contain multiple polymorphic nucleotide loci) and representative enough for a species' genome or population's genepool.

Most phylogenetic tools used for tree reconstruction from allele frequencies are not suitable for low numbers of marker genes (Phylip, DISPAN, check Gen-Pop), since they perform bootstrapping with loci (i.e., here: genes) as units. If only allele frequencies from a single gene are used, conventional bootstrapping performs resampling with replacement, draws repeatedly from only a single locus. Trivially, this single locus is in agreement with itself, and the tools report 100% branch support for all clades in the tree. False confidence Respected journals have failed to critically review this misuse of bootsrapping [4, 2]. For two genes A and B, resampling leads the combinations AA (25%), AB (50%) and BB (25%). If both A and B agree on the branch split 100% branch split support will be reported by teh software. branch split is supported by both A and B, 100 as for example is the case in [2].

We here propose a method that constructs population phylogenies for HLA allele frequencies in combination with well structured databases with deep sequencing data for those genes.

Rich databases such as AlleleFrequencies.net help to obtain population specific descriptions in terms of allele frequencies. Subsequently, mapping of alleles to probabilistic sequence representations enable the expression of equivalent information as position specific weight matrices or classic multiple sequence alignments. This mapping is facilitated by comprehensive and wellstructured databases such as IMGT. With the algorithmic transition of gene markers to nucleotide markers the number of informative loci increases by orders of magnitude. This in turn consitutes an important prerequisite for the application of classic consensus tree construction and bootstrapping, as resampling from this much larger pool of loci is statistically more meaningful.

2 Methods

We download complete multiple sequence alignments from IPD-IMGT/HLA release 3.31.0. For all genes for which multiple sequence alignments are available, we parse the respective alignment file as obtained from IMGT (ref). The Python script is available at the github repository https://github.com/HenschelLab/FreqRT.

As we are aiming to capture the entirety of genetic variability associated with HLA genes/alleles, we parse for each HLA gene g its respective multiple sequence alignment into Position Weight Matrices (PWM, [7]) by associating each sequence to the allele identified by the first two digits of the sequence identifier. This yields one PWM for each allele A, which we denote W_A .

In turn, given m selected genes, we describe a population P as a set of m PWMs W_q^P , which are the weighted sum of the respective allele PWMs

$$W_g^P = \sum_{A \in \mathcal{A}(g)} f_A^P W_A \tag{1}$$

where f_A^P is the frequency of allele A in population P and $\mathcal{A}(g)$ denotes the set of all alleles for g.

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input: Selected genes: G_1 \dots G_m, Allele frequencies f_{A_i}^{P_i} for n
              populations P_1 \dots P_n
   output: Phylogenetic tree T for P_1 \dots P_n with bootstrap values
 1 Preprocessing: Download MSAs from IMGT, \forall j generate W_{A_i}
 2 for P \in P_1 \dots P_n do
       for g \in G_1 \dots G_m do
 | W_g^P \leftarrow \sum_{A \in \mathcal{A}(g)} f_A^P W_A |
       end
 5
 6 end
 7 for bootstrap \leftarrow 1 to 1000 do
       for g \in G_1 \dots G_m do
        \overline{W_g^P} = \text{RandomResamplingColumns}(W_g^P) \text{ such that } |\overline{W_g^P}| = |W_g^P|
 9
10
       Calculate distance Matrix DM, performing pairwise Nei distance
11
       calculations between populations
       for P_i \in P_1 \dots P_n do
12
           13
14
           end
15
16
       \quad \text{end} \quad
       T_{bootstrap} = \text{TreeConstruction}(DM)
17
18 end
19 Combine bootstrap trees T_{bootstrap} to majority tree T with bootstrap
   values
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3 Conclusions

To the best of our knowledge, we produced here the most comprehensive phylogenetic trees with respect to number of populations. We have done so by carefully evaluating the confidence of each branch deploying a novel method, FreqRT. FreqRT can

The wrong practice of bootstrapping for allele frequency datasets with few genes seems to be relatively common place in the HLA community. We would hereby like to raise awareness during three time points of the publication cycle. First, software tools like Phylip, DISPAN etc ought to generate warnings, when the number of provided loci is too low to subject allele frequencies to bootstrapping. Secondly, naturally scientists creating phylogenies should know what they are doing. Finally, journals like PLoS ONE need to improve review procedures for population phylogenys built with allele frequencies, in particular those with just one or few loci.

References

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