Exploring evolutionary relationships across the genome using topology weighting

Simon H. Martin* & Steven M. Van Belleghem1,2,3

1. Department of Zoology, University of Cambridge, Cambridge CB2 3EJ, United Kingdom.

2. Department of Biology, Center for Applied Tropical Ecology and Conservation, University of Puerto Rico, Rio Piedras, Puerto Rico.

3. Smithsonian Tropical Research Institute, Apartado 0843-03092, Panamá, Panama.

* Corresponding Author: shm45@cam.ac.uk
Abstract

We introduce the concept of topology weighting, a method for quantifying relationships between taxa that are not necessarily monophyletic, and visualising how these relationships change across the genome. A given set of taxa can be related in a limited number of ways, but if each taxon is represented by multiple sequences, the number of possible topologies becomes very large. Topology weighting reduces this complexity by quantifying the contribution of each 'taxon topology' to the full tree. We describe our method for topology weighting by iterative sampling of sub-trees (Twisst), and test it on both simulated and real genomic data. Overall, we show that this is an informative and versatile approach, suitable for exploring relationships in almost any genomic dataset.

Scripts to implement the method described are available at github.com/simonhmartin/twisst.
Introduction

The evolutionary history of a set of sequences is described by their genealogical tree. In recombining organisms, the genealogy varies across the genome, and this heterogeneity can provide information about historical processes like selection and gene flow. Many of the commonly-used approaches in population genomics are aimed at describing this genealogical heterogeneity. While it may be ideal to know the complete genealogy and how it varies across the genome, it is often preferable to use statistics that summarise particular features of the underlying tree, such as $F_{ST}$. However, such statistics often fail to capture the tree-like process of sequence evolution, and qualitative information about the branching order among taxa (i.e. the topology), which is relevant for many evolutionary questions.

A pattern of particular interest in evolutionary biology is when distinct taxa show discordant relationships across the genome. This discordance can reflect both incomplete lineage sorting (ILS) and introgression. Some recent genomic studies have attempted to characterise the genomic landscape of species relationships by inferring phylogenies in windows, revealing extensive topological discordance across the genome in groups such as butterflies (Martin et al. 2013), mosquitos (Fontaine et al. 2015), cats (Li et al. 2016) and cichlids (Gante et al. 2016). A more sophisticated approach that simultaneously infers genealogies and the likely recombination points at which the genealogy switches has detected similar discordance in relationships among great apes (Dutheil et al. 2009; Scally et al. 2012). These tree-based approaches are most useful when the number of sequences - and, hence, possible tree topologies - is small, making it feasible to evaluate the support for opposing evolutionary hypotheses such as gene flow and ILS. Both the inference and interpretation of trees becomes challenging when the number of sequences is large, due to the rapid escalation of topological complexity. For example, for five haplotypes, there are 15 possible unrooted, bifurcating topologies, whereas for ten haplotypes there are over two million. This poses a problem for population genomics, where studies increasingly compare multiple taxa each represented by multiple individuals.

One way to deal with the problem of increasing tree complexity is to focus only on the relationships among broader taxa (hereafter “taxon topologies”), and not among all of the sequences. This is straightforward if each taxon is completely resolved into a monophyletic clade, so that the branching patterns within each taxon can simply be ignored, but it becomes challenging when the
taxa are not reciprocally monophyletic (i.e. due to ILS or gene flow). We note that tree inference using large genomic windows, entire chromosomes or whole genomes often yields resolved monophyletic taxa, but this may be artificial as such methods are usually forced to infer a single best-supported tree, even though the region may represent multiple distinct ancestries. Statistics based on allele frequencies, such as $F_{ST}$, have the advantage of quantifying relatedness rather than simply qualitatively describing the topology. However, by ignoring topology entirely, they become difficult to interpret when the number of taxa is greater than two. There is therefore a need for descriptive methods that incorporate both the qualitative tree-like structure of relationships and the quantitative variation in taxon relationships across the genome. Examples of such methods include the so-called ABBA-BABA test (Green et al. 2010; Durand et al. 2011; Martin et al. 2015), and f statistics (Reich et al. 2009, 2012; Patterson et al. 2012), both of which evaluate the support for alternative taxon topologies using allele frequencies at single nucleotide polymorphisms (SNPs). However, because individual SNPs are only informative about two separate groupings, these methods do not scale to more than four taxa.

Here we introduce the concept of topology weighting, which offers a simple and general solution to the problem of quantifying relationships among taxa that are not necessarily monophyletic. Given a tree of relationships for a set of taxa, each represented by an arbitrary number of sequences, topology weighting quantifies the contribution of each individual 'taxon topology' to the full tree. We describe our approach to compute topology weightings, which we call Twisst (topology weighting by iterative sampling of sub-trees), and explore the utility of this approach using simulated data as well as two different genomic datasets from butterflies and fungi. Overall, we show that this concept provides a useful means to explore relationships using genomic data, both to test hypotheses and generate new ones.

Materials and Methods

**Topology weighting by iterative sampling of sub-trees**

A given set of taxa can be related in a limited number of ways. For example, for four taxa labelled A, B, C and D, there are three possible unrooted bifurcating topologies: ((A,B),C,D), ((A,C),B,D) and ((A,D),B,C) (Fig. 1). Given a tree with any number of tips (or leaves), each assigned to a particular taxon, we define the weighting of a particular 'taxon topology', $\tau$, as the fraction of all
unique sub-trees – in which each taxon is represented by a single tip – that match the particular
taxon topology:

\[ w_t = \frac{K}{N} \]  

Where \( N \) is the number of possible unique sample sets in which each taxon is represented by a
single sample. This corresponds to the product of the number of samples in each taxon. \( K \) is the
number of these unique sample sets for which the corresponding sub-tree topology matches \( \tau \).
Thus:

\[ N = \prod_{j=1}^{n} s_j, \quad K = \sum_{i=1}^{N} [x_i = \tau] \]  

In which \( n \) is the number of defined taxa (groups), \( s_j \) is the number of samples in taxon \( j \), and \( x_i \) is
the sub-tree topology corresponding to subset \( i \).

Topology weighting therefore reduces the complexity of the full tree to a number of values, each
giving the proportionate contribution of a particular taxon topology. This method has conceptual
similarities to the quartet sampling approach for comparing topologies (Estabrook et al. 1985),
except that here we do not consider all sub-trees, but only those in which each tip represents a
different taxon. Moreover, topology weighting can be applied to any number of taxa. The taxa can
be defined arbitrarily, for example by phenotype or geography, as with the 'operational taxonomic
units' used in biogeography.

We computed topology weightings using our Twisst approach, implemented by Python scripts
available for download at (https://github.com/simonhmartin/twisst). The Twisst algorithm first
computes all possible unrooted, bifurcating taxon topologies, and then determines the number of
unique sub-trees that match each topology by iteratively sampling a single individual from each
taxon and 'pruning' away all other branches and nodes. Our implementation makes use of the
Environment for Tree Exploration (ETE 3) (Huerta-Cepas et al. 2016).

The number of unique sample combinations (and corresponding sub-trees) can be very large.
However, the iterative process is sped up considerably by first collapsing monophyletic groups of
samples from the same taxon and weighting these nodes proportionately (Fig. S1). Nevertheless, if
the taxa are highly unsorted and the tree is large, it may not be possible to consider all possible
Figure 1. Topology weighting

(A) An example genealogy for four taxa (A, B, C and D), plotted as an unrooted tree on the top right. Taxa B and C are not monophyletic, due to incomplete lineage sorting and gene flow. The three possible taxon topologies are shown, along with a single example sub-tree that matches each topology. The percentage of all sub-trees matching each taxon topology (i.e. the weightings) are shown by vertical bars. (B) Topology weightings plotted across a 50 kb region of a simulated recombining chromosome. Weightings for the three topologies are stacked (they always sum to 1, as they are proportions). Changes in the weightings along the chromosome indicate regions of distinct genealogical history separated by recombinations. Below, the same data are plotted with loess smoothing (span = 2.5 kb).
sample combinations in a reasonable amount of time. In such cases, approximate weightings can be computed by randomly sampling a subset of sample combinations (Fig. S2). Random sampling for approximate weighting is performed with replacement, so that the errors fit a binomial distribution, allowing for the computation of a confidence interval (Fig. S2). Our implementation of Twisst also allows a threshold-based sampling procedure, in which sampling is repeated until a particular level of confidence is achieved. This allows further speed-ups since highly sorted trees, in which weightings are biased toward one or a few topologies, require less sampling for high confidence than unsorted or 'star-like' trees where most weightings are intermediate. For all analyses in this study, complete weightings were computed because the datasets were small enough.

**Analysis of simulated chromosomes**

In order to test our method on realistic data for which the complete genealogical history is known, we simulated the evolution of recombining chromosomes using the coalescent simulator *msms* (Ewing and Hermisson 2010). All simulations involved forty haploid individuals from four taxa (10 individuals each). The four taxa split in the order (((A,B),C),D) with split times of 0.5, 1 and 1.5 (in units of 4N generations), respectively (Fig. 2A). Population size was constant throughout. In all three scenarios, unidirectional migration from C to B was simulated. The simulation was performed for a 1 Mb chromosome, with a population recombination rate (4Nr) of 0.01 or 0.001. Genealogies for each unique ancestry block (separated by recombinations) were recorded.

Three distinct evolutionary scenarios were simulated (Fig. 2A). *msms* command options are provided in Supplementary Text 1. The first was a 'Neutral' scenario, with no selection and a low migration rate from B to C of 0.1 (in units of 4Nm, where m in the fraction of B made up of migrants from C each generation). The second was an 'Adaptive introgression' scenario, which is the same as above except that a beneficial allele at a locus in the centre of the 1 Mb chromosome is allowed to move from population C into B at time 0.1. This was achieved by initiating selection at this time point on a dominant allele that was fixed in population C and absent from the other populations. A selection coefficient of 0.005 was used for both the homozygote and heterozygote, with a diploid population size of 100,000, giving a selection strength (2Ns) of 1000 for both genotypes. The third was a 'barrier locus' scenario, where the rate of migration from C to B was 5 (in units of 4Nm, as above), and a dominant allele at the central locus that is fixed in C is selected against in population B. The same selection coefficient and population size as above were used.
We simulated sequences from the simulated genealogies using *seq-gen* (Rambaut and Grass 1997). Command options are provided in Supplementary Text 1. The branch scaling factor for mutation was 0.01. Since branches in the simulated genealogies are in units of $4N$ generations, taking $N$ to be 100,000 gives $\mu$ (the per generation mutation rate) of $2.5 \times 10^{-8}$.

**Inferring trees in sliding windows**

We tested different methods for inferring trees in windows across the chromosome. The simplest approach used non-overlapping windows of a fixed number of SNPs. A range of window sizes were tested. Trees were then inferred for each window using PhyML version 3.0 (Guindon *et al.* 2010), implementing either the BIONJ neighbour-joining algorithm (Gascuel 1997) or maximum likelihood optimization of the topology and branch lengths. We also tested an approach to infer likely window breakpoints from the data. Taking the topology weightings computed from 10 SNP windows, we used the R package *GenWin* (Beissinger *et al.* 2015) to fit a beta-spline to the data and find likely inflection points, which we then used as window breakpoints and inferred a new set of trees for these. In addition, we tested the program *Saguaro*, which infers both the breakpoints and the distance matrix describing each region. Distance matrices were converted to trees using BIONJ, as above.

**Power Analyses**

An important aspect of our approach is its dependence upon reliable trees inferred from relatively short sequence windows. To investigate the power we have to infer topology weightings from short sequences, we simulated datasets under a range of sampling strategies and demographic scenarios, and then compared the true weightings to those computed using trees inferred from the simulated sequences.

Eight sampling strategies were compared, including four, five, six or ten sequences from either four or five populations (Fig. S3). For each sampling strategy, two different demographic scenarios were simulated. In the four-population scenarios, the populations split in the order $((1,2),(3,4))$, with the basal split time ($t_1$) at either 0.5 or 1 x 4N generations in the past, and the splits between populations 1 and 2 and 3 and 4 both occurring at 0.1 x 4N generations in the past ($t_2$) (Fig. S3). In the five-population cases, the populations split in the order $(((1,2),(3,4)),5)$. As above, the basal split time ($t_1$) occurs at either 0.5 or 1 x 4N generations in the past. The next split, between populations 1 and 2 and populations 3 and 4, occurs at 0.2 x 4N generations in the past ($t_{1b}$), and the final two
splits between populations 1 and 2, and 3 and 4 both occur at 0.1 x 4N generations in the past (t2) (Fig. S3).

In each run, we used msms (Ewing and Hermisson 2010) to simulate 500 genealogies for the given sampling design and demographic scenario. For each genealogy, we computed the topology weightings using Twisst, and then generated a simulated set of sequences using seq-gen (Rambaut and Grass 1997). The sequences was then truncated at different lengths to compare tree inference using 10, 25, 50, 100, 200 or 400 SNPs. Trees were inferred in Phyml (Guindon et al. 2010), using either BIONJ, or with maximum likelihood optimisation of the topology and branch lengths. Weightings were then computed from the inferred tree, and compared to the set of true weightings using a scaled euclidean distance:

\[ d = \sqrt{\frac{\sum_{i=1}^{n} \left( \frac{w_i - \hat{x}_i}{m_i} \right)^2}{m_i^2 \sqrt{n-1}}} \]

Where \( n \) is the number of weighting values (i.e. the number of taxon topologies), \( w_i \) is the true weighting for topology \( i \) and \( \hat{x}_i \) is the inferred weighting for topology \( i \). \( m_i \) is the absolute value of the maximum possible distance from \( w_i \). This value therefore gives a distance between the true and inferred sets of weightings on a scale of 0 to 1, with 0 indicating identical values (i.e. perfect inference) and 1 indicating a maximum possible discrepancy between the true and inferred values.

Not all SNPs are phylogenetically informative, and even those that are (those that are not singletons), are not necessarily informative about the relationships among the broader taxa, which is of primary interest for topology weighting. We therefore also tested the power of inference using a subclass of “Taxon Informative Sites” (TitS), which we define as having at least two alleles present in at least two taxa. As above, simulated sequences were truncated to contain the number of TitS.

**Analysis of real genomic data**

We tested Twisst on two published genomic datasets from Neurospora spp. (ascomycete fungi) and Heliconius spp. (butterflies), selected to represent different sampling strategies (4 and 5 taxa, respectively), as well as different levels of evolutionary complexity. The Neurospora dataset (Corcoran et al. 2016) consisted of 22 aligned haploid genome sequences from Neurospora tetrasperma samples (10 of mating type A, and 12 of mating type a), along with single genomes representing two related species: Neurospora crassa, and Neurospora hispaniola. Whole genome
alignments were obtained from datadryad.org/resource/doi:10.5061/dryad.162mh. We used Lineage-10 (UK) samples of *N. tetrasperma*, as these had been shown to carry a strong signal of introgression from *N. hispaniola* (Corcoran et al. 2016). Trees were constructed for sliding windows of 50 SNPs using BIONJ as described above, with the requirement that each sample had to be genotyped at 40 or more of the 50 SNPs per window. Topology weightings were computed using Twisst, with four defined taxa: *N. tetrasperma mat a* (12 sequences), *N. tetrasperma mat A* (10), *N. crassa* (1) and *N. hispaniola* (1).

The *Heliconius* dataset consisted of 18 resequenced genomes (or 36 haploid genomes) from Martin et al. (Martin et al. 2013). These samples comprised five populations: two geographically isolated races of *Heliconius melpomene*, from Panama (*H. m. rosina*) and Peru (*H. m. amaryllis*), and their respective sympatric relatives *Heliconius cydno chioneus* (Panama) and *Heliconius timareta thelxinoe* (Peru), with which they are known to hybridize; along with two additional samples of the more distant 'silvaniform' clade to serve as outgroups. We limited our analysis to two chromosomes: 18, which carries the gene *optix*, known to be associated with red wing pattern variation; and 21, the Z sex chromosome, which has been shown to experience reduced gene flow between these species, probably due to genetic incompatibilities (Martin et al. 2013). Fastq reads were downloaded from the European Nucleotide Archive, study accession ERP002440. Reads were mapped to the *Heliconius melpomene* reference genome v2 (Davey et al. 2016) using BWA-mem (Li and Durbin 2009; Li 2013) with default parameters. Genotyping was performed using the Genome Analysis Toolkit (DePristo et al. 2011) v3 HaplotypeCaller and GenotypeGVCFs, with default parameters except that heterozygosity was set to 0.02. Phasing and imputation was performed using Beagle v4 (Browning and Browning 2007). Topology weighting was performed using Twisst, with five taxa: *H. melpomene rosina* from Panama (4 diploid samples), *H. melpomene amaryllis* from Peru (4), *H. cydno chioneus* from Panama (4) and *H. timareta thelxinoe* from Peru (4), and two outgroup samples from the 'silvaniform' clade, *H. pardalinus sergestus* and *H. pardalinus ssp. Nov.* from Peru.

**Results**

**Analysis of simulated chromosomes**

Topology weighting provides an informative summary of the genealogical data, and highlights differences between the three simulated scenarios (Fig. 2). As described above, there are three
possible unrooted topologies for the four taxa. Note that we illustrate these as rooted with D as the outgroup for simplicity, but the rooting is not considered when computing the weightings. In the 'Neutral' scenario, the most prevalent topology (((A,B),C),D), which reflects the population split times, has an average weighting of 71% across the chromosome. The other two topologies are both fairly rare, but one (((B,C),A),D) is more common on average (17%) than the other (((A,C),B),D) (12%). This is because the former can result from both gene flow and incomplete lineage sorting (ILS), whereas the latter can only result from ILS, as there was no simulated migration between A and C or between B and D. In the 'Adaptive introgression' scenario, the weightings are very similar to the Neutral scenario on average, but in the centre of the chromosome there is a strong excess of

Figure 2. Tests on simulated chromosomes

(A) In all three demographic scenarios, populations split in the order (((A,B),C),D), at the split times indicated (in units of 4N generations), with migration from C to B. In the 'Neutral' scenario, there is no selection and moderate migration. The 'Adaptive introgression' scenario is similar, except a beneficial allele at a locus in the centre of the chromosome is allowed to move from population C into B at time 0.1. In the 'Barrier locus' scenario, the rate of migration is high, but an allele at the central locus that is fixed in C is selected against in population B. (B) Mean weightings for the three possible topologies across the 1 Mb simulated chromosome (C) Weightings for all three topologies plotted (stacked) across the chromosome, with loess smoothing (span = 20 kb). (D) Weightings for topology (((A,B),C),D) inferred from simulated sequence data using non-overlapping 50 SNP windows and neighbour joining. Solid blue lines indicate the true values, and dashed black lines indicate the inferred values. Values are smoothed as above.
the topology (((B,C),A),D), created by the spread of a beneficial allele from population C into B. Finally, in the 'Barrier locus' scenario, high migration from C to B causes a swamping by the topology (((B,C),A),D), which has an average weighting of 65%. However, there is a broad peak at the centre of the chromosome where the split-time topology (((A,B),C),D) had not been eroded, due to selection limiting introgression.

**Inferring weightings from simulated sequence data**

Above, we computed the weightings directly from the simulated genealogies, but we are also able to show that topology weightings can be reliably estimated when the genealogies are inferred from simulated sequence data (Fig. 2D). Because neither the genealogies, nor the recombination breakpoints at which genealogies switch are known, we tested several approaches for inferring genealogies for narrow intervals across the chromosome. First, we performed extensive power analyses, covering a range of demographic scenarios and sampling designs, to explore the relationship between the number of SNPs used for tree inference and the accuracy of topology weighting. Across the range of scenarios investigated, we find a consistent lower bound of 50 SNPs to achieve >90% accuracy (Fig. S4, S5, S6). Focussing specifically on 'taxon informative sites' makes no discernible difference, probably because most SNPs in our simulations are taxon informative. These tests also indicate that neighbour-joining trees provide more accurate weightings than maximum likelihood trees, in addition to much faster computation (Fig. S4, S5, S6).

We then analysed trees inferred for non-overlapping windows across our simulated recombining chromosomes. A fixed window size of 50 SNPs gives results that most closely approximate the true weightings (Fig. 2D). In agreement with our power analyses, the estimates are less accurate when windows with fewer than 50 SNPs are used (Fig. S7, S8). Interestingly, windows of 100 SNPs or above also result in reduced precision, with a tendency to overestimate support for the most common topology and underestimate support for others (Fig. S7, S8). This can be explained by the fact that large windows are forced to average over regions of distinct ancestry, therefore losing resolution. To confirm this hypothesis, we repeated our neutral simulation using a ten-fold lower population recombination rate. In this new dataset, 100 SNP windows are the most accurate, and even 200 SNP windows have high accuracy, while 50 SNP windows perform only marginally less well (Fig. S9, S10).
Because real recombination breakpoints are not evenly spaced, we also tested two approaches in which the window boundaries are inferred from the data itself. In our first approach, we used the R package GenWin (Beissinger et al. 2015) to fit a smooth spline to the weightings from 10-SNP windows and identify likely window boundaries as inflection points, and then inferred trees for the newly-defined window regions. The resulting topology weightings match the true weightings fairly well, but not as well as for the fixed 50 SNP windows (Fig. S11, S12). As above, this appears to be due to poor tree inference in the smallest windows. The second approach used the method Saguaro (Zamani et al. 2013), which combines a Hidden Markov Model and a Self Organising Map to infer both the trees and window boundaries. This approach poorly recapitulates the true weightings, greatly overestimating support for the most abundant topology (Fig. S11, S12). We therefore used fixed windows of 50 SNPs for all further analyses.

**Analysis of real genomic data**

The Neurospora dataset consists of four taxa (three possible topologies), and is the simpler of the two real datasets analysed (Fig. 3A,B). It was selected to test how well Twisst is able to detect the signal of a previously described adaptive introgression event from \( N. \) hispaniola into \( N. \) tetrasperma individuals of the A mating type (Corcoran et al. 2016). This introgression covers the entire (~7 Mb) non-recombining region of linkage group I (LGI). Indeed, we find a dramatic shift in the pattern of topology weightings in the central part of LGI (Fig 3C). The “species-tree” topology (topo1), which groups the two \( N. \) tetrasperma mating types as closest relatives, is strongly supported across most of the genome, but has almost no support in the central part of LGI. Instead, it is replaced by topo3, which groups mating type A individuals of \( N. \) tetrasperma with \( N. \) hispaniola with. Elsewhere, topo3 has weak support, nearly identical to that of topo2, consistent with a low level of incomplete lineage sorting throughout the genome. However, a region of LGIV also shows a weak shift in support towards topo3, potentially reflecting a separate and more subtle introgression signal.

The Heliconius dataset represents a more complex, five-taxon test case. The five taxa include an outgroup and two pairs of sympatric, non-sister taxa, between which gene flow is known to occur (Fig. 4A). Of the 15 possible topologies (Fig. 4B), the two most common across these chromosomes are ‘topo3’ and ‘topo6’. topo3 is consistent with the accepted species branching order, in which the allopatric \( H. \) cydno chioneus and \( H. \) timareta thelxinoe are sister taxa; whereas topo6 groups
populations by geography, consistent with inter-specific gene flow in both Panama and Peru. The former is by far the most prevalent throughout the Z chromosome (Fig. 4C). By contrast, support for the species topology is variable across chromosome 18, and is in places outweighed by support for topologies consistent with gene flow (topo4, topo5, topo6, topo11, topo14). In particular, there is a strong peak in the region of optix for topo11, which groups the taxa by wing pattern, and is consistent with the previously described adaptive introgression of the red-band allele between H. m. amaryllis and H. t. thelxinoe in Peru (Pardo-Diaz et al. 2012; The Heliconius Genome Consortium 2012). Another four topologies that shared at least one split with the species tree (topo1, topo2, topo10, topo15), had moderate support throughout, whereas topologies consistent with neither the species tree nor gene flow (topo7, topo8, topo9, topo12, topo13) were weakly supported throughout, and particularly weakly across the Z chromosome, implying less incomplete lineage sorting than on chromosome 18.
Discussion

Most statistics used in population genetics describe aspects of the underlying genealogy. For example, $F_{ST}$ can be interpreted as a measure of the relative rate of coalescence within sub-populations compared to the total population (Slatkin and Voeim 1991). Similarly, the $D$ statistic of the ABBA BABA test compares the relative rate of coalescence between two pairs of non-sister populations (Green et al. 2010; Durand et al. 2011). Topology weighting can be seen as a generalization of this principle, as it determines the relative frequency of all possible patterns of coalescence among samples from a set of defined taxa. Unlike the ABBA BABA test, which is based on binary trees (samples either share the same allele or not) and therefore only four taxa, topology weighting can in principle be used to summarise relationships between any number of taxa, each represented by any number of sequences. In practice, however, beyond six taxa, the
number of possible taxon topologies becomes very large, making topology weighting less practical. Nevertheless, even when the number of taxon topologies is very large, there may be value in comparing the frequencies of particular topologies of interest. There is no real limitation on the number of samples that can be analysed per taxon. Although computation of exact weightings may become infeasible for large trees, it remains possible to compute approximate weightings with small margins or error fairly rapidly. Our computational approach, Twisst, is based on a simple counting procedure, but we are confident that more efficient analytical solutions – or at least approximations – will be found.

One important consideration when applying this method is its dependence on the inferred trees. In most cases, the true genealogy for each distinct ancestry block is not known, and must be inferred from the sequences. Accurate inference requires multiple informative SNPs. Our tests on simulated data highlight a central difficulty when analysing recombining chromosomes: a trade-off between signal and resolution. Using larger numbers of SNPs increases our ability to infer the correct tree, but may average over genomic regions with different histories. This leads to a systematic overestimation of the weightings for more abundant topologies, whose signal tends to swamp out that of others. Using few SNPs per window allows for better resolution, but can lead to inaccuracies in tree reconstruction from insufficient signal (i.e. phylogenetic error), producing star-like trees and intermediate weightings for all topologies. Our simulations, based on realistic recombination and mutation rates, indicate that a window size of 50 SNPs provides a good compromise between signal and resolution across a range of demographic scenarios, although larger windows may be preferable if the population recombination rate is known to be low. While in some cases with high recombination rates, there may be too few mutations per recombination to accurately infer variation in genealogies across the genome, we expect that many cases will fall within a feasible range. Importantly, not all recombinations are relevant for topology weighting: only recombination events between lineages from distinct taxa (i.e. 'effective recombinations' or 'inter-taxon recombinations') can alter taxon relationships, and hence alter the topology weightings. The ability to infer the patterns of topology weighting across the genome therefore depends on the relationship between the mutation rate and the rate of inter-taxon recombination. Where possible, simulations tailored to the taxa being studied can be used to guide the choice of window size. In the future, improved methods to infer breakpoints from the data may further resolve this difficulty.
Another challenge is that diploid resequencing data should ideally be phased, so that each tip in the tree represents a distinct haplotype. Phasing can be performed using probabilistic approaches informed by patterns of linkage disequilibrium (Browning and Browning 2007; Delaneau et al. 2013). Although such methods may be error-prone across large genomic distances, they have fairly high accuracy at short ranges (Bukowicki et al. 2016), making them suited to the narrow windows used for topology weighting. Moreover, the genomic regions involving inter-taxon recombinations (i.e. those relevant for topology weighting) are more likely to be phased correctly, because they tend to involve more divergent sequences.

Topology weighting is principally a descriptive method, and can be applied with no prior knowledge of the studied samples, apart from some basis on which to define distinct groups, such as geography or phenotype. By capturing the tree-like nature of sequence evolution, it provides information that is not provided by descriptive statistics like $F_{ST}$, or clustering methods such as Structure (Pritchard et al. 2000). In addition to describing the taxon branching order, tree-based methods allow incorporation of additional parameters like a nucleotide substitution model and different rates of evolution in different parts of the tree. Unlike conventional phylogenomic methods, topology weighting captures information about fine-scale and quantitative variation across the genome. This power and resolution is highlighted in the Heliconius example studied here, where topologies supporting admixture are common across chromosome 18, but there is one narrow peak consistent with the adaptive introgression of a wing patterning allele near the gene optix, as described previously (Pardo-Diaz et al. 2012; The Heliconius Genome Consortium 2012). We note that topology weighting simply describes the signal in the data, and does not explicitly test for introgression over other causes of discordant phylogenetic signal. For example, the elevated frequency of topologies consistent with introgression across Heliconius chromosome 18 compared to the Z chromosome is only partially due to an elevated rate of gene flow on autosomes (Martin et al. 2013), but also reflects increased incomplete lineage sorting due to their larger effective population size relative to the sex chromosome. Finally, these findings also indicate that topology weighting provides a means to identify candidate loci underlying trait variation, based on clustering of taxa by phenotype. In summary, topology weighting is a simple but versatile exploratory tool that is applicable to a diverse range of questions and datasets.
Acknowledgements

We thank Pasi Rastas, Markus Möst and Joe Hanly for useful comments on the method. SHM was funded by a research fellowship from St John's College, University of Cambridge. SMVB was funded by NSF grant DEB-1257839.

References


Li G., Davis B. W., Eizirik E., Murphy W. J., 2016 Phylogenomic evidence for ancient hybridization in the genomes of living cats (Felidae). Genome Res. 26: 1–11.


Reich D., Patterson N., Campbell D., Tandon A., Mazieres S., Ray N., Parra M. V, Rojas W., Duque C., Mesa N., García L. F., Triana O., Blair S., Maestre A., Dib J. C., Bravi C. M., Bailliet G., Corach D,


Supplementary Figures

Figure S1. Tree Simplification
The full tree (left) has four defined taxa of ten samples each, indicated by different colours. This equates to 10,000 ($10^4$) unique sub-trees that include a single individual from each taxon. *Twisst* reduces the number of unique sub-trees to count by collapsing monophyletic single-taxon clades and weighting them by the number of individuals present. In this example, after this process, there remain only 12 unique sub-trees to count (4x3x1x1).
Figure S2. Approximate weighting by random sampling

This example tree has four defined taxa indicated by different symbols. There are three possible taxon topologies (top right). The graph shows the estimated weighting for each topology (Y-axis) after randomly sampling a number of sub-trees from the full tree (X-axis). The 95% binomial confidence interval (calculated using the Wilson method), is shaded. Dashed lines show the true exact weightings computed by sampling all 10,000 sub-trees.
Figure S3. Demographic scenarios for power simulations

Four (left) and five (right) population simulations were performed. Split times are shown, note that two different values were tested for split time $t_1$ in both scenarios.
Figure S4. Power analyses with four equal-sized groups

Each plot in the first four columns shows the error rate, calculated as a scaled euclidean distance between the true and inferred weightings (Y-axis), plotted against the number of sites used for tree inference (X-axis). The simulated sequences were truncated to create a sequence of the correct length for tree inference. Truncation was performed either after X SNPs were observed (columns 1 and 3) or after X 'taxon-informative sites' (TitS) had been observed (columns 1 and 4). The final column gives the number of SNPs required (Y-axis) to observe a certain number of TitS (X-axis). The first two columns show results after tree inference using neighbour joining (NJ) and the next two columns show results after tree inference using maximum likelihood (ML). Each row represents a distinct sampling strategy (four groups of four samples, four groups of five samples etc.). Red and blue lines indicate the two different demographic scenarios tested, with a different split time $t_1$ (see Fig. S3).
Figure S5. Power analyses with four groups of different sizes

As in Fig. S4, except for groups of different sizes.
Figure S6. Power analyses with five equal-sized groups

As in Fig. S4, except for simulations with five groups.
Figure S7. Inferred vs true weightings using different window sizes ($\rho = 0.01$)

The true (solid coloured line) and inferred (dashed black line) weightings, plotted across the simulated 1 Mb chromosome, with loess smoothing (span 0.04). The three columns with different colours represent the three taxon topologies (see Fig. 2 in the main paper). Rows represent different window sizes (fixed number of SNPs) for tree inference (neighbour joining).
Figure S8. Weighting error rates using different window sizes ($\rho = 0.01$)

Error rate, calculated as a scaled euclidean distance between the true and observed weightings, averaged over the 1 Mb simulated shromosome. Error rates were computed after first smoothing both the observed and true weightings using loess (span = 0.04).
Figure S9. Inferred vs true weightings across simulated chromosomes ($\rho = 0.001$)

As in figure S7, except here simulations used a population recombination rate ($\rho$) of 0.001.
Figure S10. Weighting error rates using different window sizes ($\rho = 0.001$)

As in figure S8, except here simulations used a population recombination rate ($\rho$) of 0.001.
Figure S11. Inferred vs true weightings across simulated chromosomes

As in figure S7 and S9, except here comparing different methods for window-based tree inference: 50 SNP windows (first row), WinGen inference of breakpoints (second row) or Saguarro (row three).
Figure S12. Weighting error rates using different methods for window-based tree inference

As in figure S8 and S10, except here comparing different methods for window-based tree inference.