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by

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I would not have made it through this journey without the support of many people. Title: Begin the section with a heading such as "Acknowledgements" or "Acknowledgments" centered at the top of the page.

Introduction: Write a brief introductory paragraph expressing your gratitude and acknowledging the support and contributions of individuals and organizations throughout your research journey.

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To so-and-so. . .

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CHAPTER I

GREATAPES

1.1 Introduction

Genetic variation is determined by the combined action of mutation, demographic processes, recombination and natural selection. However, there is still no consensus on the relative contributions of these processes and their interactions in shaping patterns of genetic variation. Two major open questions are: how does the influence of selection compare to other processes? And, to what degree is genetic variation influenced by beneficial versus deleterious mutations?

Genetic variation can be measured within a species or between species with two related metrics: within-species genetic diversity and between-species genetic divergence. Both can be estimated with genetic data by computing the per site average number of differences between pairs of samples within a species or between two species. and these are estimates of the mean time to coalescence. (Note that we do not discuss relative divergence, which is often measured using F_{ST} .) Evolutionary processes impact diversity and divergence in different ways, so the relationship between these carries information regarding these processes.

Natural selection directly impacts genetic diversity because it can reduce the frequencies of alleles that are deleterious (negative selection) or increase those of beneficial alleles (positive selection). Selection can also directly affect between-species genetic divergence. Deleterious alleles are more likely to be lost from the population, thus reducing divergence at the affected sites. On the other hand, beneficial alleles have a higher probability of fixation. This leads to an increase in the rate of substitution at sites under positive selection that in turn increases divergence between species. Thus, contrasting patterns of diversity and divergence

at the same time can help disentangle between modes of selection (Hudson et al., 1987). Indeed, perhaps the most widely used test for detecting adaptive evolution, the McDonald-Kreitman test, compares diversity and divergence contrasted between neutral (e.g., synonymous) and functional (e.g., non-synonymous) site classes (McDonald & Kreitman, 1991). This test and its extensions have been applied to a myriad of taxa, and it has become clear that a substantial proportion of amino acid substitutions are driven by positive selection in a number of taxa (Galtier, 2016; Ingvarsson, 2010; Slotte, 2014; N. Smith & Eyre-Walker, 2002).

Selection also disturbs genetic variation at nearby locations on the genome, and this indirect effect of selection on diversity is called "linked selection". Linked selection can be caused by at least two familiar mechanisms: genetic hitchhiking and background selection. Under genetic hitchhiking, as a beneficial mutation quickly increases in frequency in a population, its nearby genetic background is carried along, causing local reductions in levels of genetic diversity. The size of the region affected by the sweep depends on the strength of selection, which determines how fast fixation happens, and the crossover rate, because recombination allows linked sites to escape from the haplotype carrying the beneficial mutation (Kaplan et al., 1989; Maynard Smith & Haigh, 1974). Under background selection, neutral variation linked to deleterious mutations is removed from the population unless, as before, focal lineages escape via recombination (Charlesworth et al., 1993). Both of these processes leave similar footprints on patterns of within-species genetic diversity, and so attempts to determine the contributions of positive and negative selection in shaping levels of genetic variation genome-wide have proven to be difficult (Andolfatto, 2001; Y. Kim & Stephan, 2000), although the processes seem separable more locally (Schrider, 2020; Schrider & Kern, 2017). Importantly,

linked selection has more limited effects on between-species genetic divergence, as a beneficial or deleterious mutation does not affect the substitution rate of linked, neutral mutations (Birky & Walsh, 1988).

The effects of linked selection in shaping genetic variation are pervasive across genomes (Begun & Aquadro, 1992; Cai et al., 2009; Corbett-Detig et al., 2015; Lohmueller et al., 2011; Murphy et al., 2022). For example, dips in nucleotide diversity surrounding functional substitutions have been uncovered in many taxa, such as fruit flies (Kern et al., 2002; Sattath et al., 2011), rodents (Halligan et al., 2013), Capsella (Williamson et al., 2014) and maize (Beissinger et al., 2016). In Drosophila melanogaster, levels of synonymous diversity (which is putatively neutral) and amino acid divergence are negatively correlated (Andolfatto, 2007; Macpherson et al., 2007); positive selection can cause such a pattern if beneficial amino acid mutations are fixing and as they do reducing levels of linked neutral variation via selective sweeps. In contrast in humans, levels of synonymous diversity are roughly the same near amino acid substitutions and synonymous substitutions, suggesting recent, recent fixations at amino acids sites may not be the result of strongly beneficial alleles (Hernandez et al., 2011; Lohmueller et al., 2011). However, in the human genome, amino acid substitutions tend to be located in regions of lower constraint than synonymous substitutions, implying that the signal of positive selection may be confounded by the effects of background selection (Enard et al., 2014).

Two major challenges remain in the way of a fuller characterization of the effects of selection on genetic variation: (i) it is hard to model interactions between evolutionary processes (e.g., sweeps within highly constrained regions), and (ii) model identifiability is challenging for some summaries of the data (e.g., sweeps and

background selection may impact diversity in similar ways). Recent computational advances have made it possible for us to move from simpler backwards-in-time coalescent models (Hudson, 1983) to more complex and computationally demanding forward-in-time simulations, and these have provided a route to studying these hard to model interactions between evolutionary processes across multiple sites (Haller & Messer, 2019; Haller et al., 2019; Kelleher et al., 2016). Simulation-based inference can then allow us to better describe the roles of different modes of selection and other processes in shaping genomic variation. However, the problem of identifying features of the data that are informative of the strength and mode of selection still remains.

One promising approach might be to compare patterns of genetic variation in multiple species jointly as each species can be thought of as semi-independent realizations of the same evolutionary processes (c.f. Won & Hey, 2005). In speciation genomics studies, it is common to visualize large scale patterns of genetic variation along chromosomes (so-called landscapes of diversity and divergence), which may contain substantial information to help us disentangle evolutionary processes. Earlier empirical surveys have focused on the identification of regions of accentuated relative divergence between populations (Cruickshank & Hahn, 2014; Harr, 2006; Turner et al., 2005), although patches of increased divergence can be the result of myriad forces besides reproductive isolation and adaptation. Recent comparative studies have found that landscapes of diversity are highly correlated between related groups of species, such as *Ficedula* flycatchers (Burri et al., 2015; Ellegren et al., 2012), warblers (Irwin et al., 2016), stonechats (Doren et al., 2017), hummingbirds (Battey, 2020), monkeyflowers (Stankowski et al., 2019) and *Populus* (Wang et al., 2020). Burri (2017) proposed that we could capitalize on correlated

genomic landscapes to study the interplay between different forms of selection and other evolutionary processes. Neutral processes, such as incomplete lineage sorting or migration, could potentially produce significant correlations in levels of diversity across species, however strong correlations have been observed among taxa with long divergence times and without evidence of gene flow. For example, Stankowski et al. (2019) found that landscapes of diversity and divergence are highly correlated across a radiation of monkeyflowers which spans one million year (or about $10N_e$ generations, where N_e is the effective population size), far longer than what is relevant for incomplete lineage sorting (i.e., the coalescent timescale spans just a few multiples of N_e). However, a shared process that independently occurs in the branches of a group of species could maintain correlations over long timescales. For example, if two species' physical arrangement of functional elements and local recombination rates are similar, the direct and indirect effects of selection could make it so that peaks and valleys on the landscape of diversity are similar, maintaining correlation between their landscapes over evolutionary time (Burri, 2017; Delmore et al., 2018). Further, if mutational processes are heterogeneous across the genome in a manner that is shared among species, then correlated landscapes of diversity could be created through mutational variation as well.

Here, we aim to (i) describe whether and in what ways landscapes of withinspecies diversity and between-species divergence are correlated, and (ii) to tease apart the relative roles of positive and negative selection and other processes (e.g., ancestral variation, mutation rate variation) in shaping patterns of genetic variation. To understand processes driving these correlations, we employ highly realistic, chromosome-scale, forward-in-time simulations, since analytical predictions are not available. We use the great apes as a system to investigate correlated patterns of genetic variation because there is high quality population genomic data for all species (Prado-Martinez et al., 2013), the clade is about 12 million years old or $60N_e$ generations (but there have not been many chromosomal arrangements (Jauch et al., 1992)), and lastly the landscapes of gene density, recombination rate and mutation rate are roughly conserved (Kronenberg et al., 2018; Stevison et al., 2016). Our study demonstrates that correlated landscapes can be useful in distinguishing between modes of selection and the balance of direct and linked selection shaping genomic variation.

1.2 Methods

1.2.1 Genomic data. We retrieved SNP calls for ten great ape populations made on high coverage ($\sim 25 \times$) short-read sequencing data from the Great Ape Genome Project (Prado-Martinez et al., 2013), mapped onto the human reference genome (NCBI36/hg18). We analyzed 86 individuals divided into the following populations: human (n = 9 samples), bonobo (n = 13), Nigeria-Cameroon chimpanzee (n = 10), eastern chimpanzee (n = 6), central chimpanzee (n = 4), western chimpanzee (n = 4), eastern lowland gorilla (n = 3), western gorilla (n = 27), Sumatran orangutan (n = 5), Bornean orangutan (n = 5) (we excluded two samples from the original dataset: the Cross River gorilla and the chimpanzee hybrid). Prado-Martinez et al. (2013) applied several quality filters to the SNP calls (see Section 2.1 of their Supplementary Information) and, for each species, identified the genomic regions in which it would be unreliable to call SNPs (uncallable regions). For our downstream analyses, we only considered sites which were callable in all populations.

We calculated nucleotide diversity and divergence (d_{XY}) in non-overlapping 1Mb windows using scikit-allel (Miles et al., 2020). Windows in which there

were less than 40% callable sites were not used in any of the analyses. For example, this yielded 129 (out of 132) 1Mb windows in chromosome 12 in which 75% of the sites were callable on average.

To tease apart the effects of GC-biased gene conversion (gBGC), we decomposed diversity and divergence by allelic states. gBGC is expected to affect weak bases (A or T) which are disfavored when in heterozygotes which also carry a strong base (G or C). Thus, one way understand the effects of gBGC is by comparing sites which were weak to those that were strong in the ancestor (ancestrally strong alleles are not affected by gBGC, but ancestrally weak alleles can be). We assumed that the state in the ancestor of the great apes to be the state seen in rhesus macaques (genome version RheMac2) — sites without enough information in RheMac2 were excluded. Then, we computed divergence only considering sites which were ancestrally weak or ancestrally strong (Figure A.4). This approach has two major drawbacks: (i) many of the sites cannot be used because they are missing in RheMac2 and (ii) sites can be mispolarized. Thus, we came up with a second approach to tease apart the effects of gBGC on correlations between genomic landscapes. When comparing two landscapes of divergence (which encompass four species), we can classify each site by the change in state that happened without needing to polarize mutations by looking at the ancestor. For example, if we have allelic states for four species and we see A-A-T-T as the configuration of alleles at a particular site, we know that there must have been one mutation which changed the state from a weak base to another weak base (W-W). On the other hand, if we see A-G-A-A there must have been one mutation from weak to strong (W-S) (or vice-versa). Sites with multiple mutations (e.g., A-G-G-C) were removed from the analyses. Sites that did not change from W to S

(or vice-versa) are not expected to be affected by gBGC, and we refer to these as W-W or S-S mutations (Figure 1.7A). Sites where there may have been a weak to strong change (W-S mutations) may be affected by gBGC (Figure 1.7B). We only considered windows with at least 5% of callable sites in these analyses.

1.2.2 Simulations. We implemented forward-in-time Wright-Fisher simulations of the entire evolutionary history of the great apes using SLiM (Haller & Messer, 2019; Haller et al., 2019). Each branch in the great apes' tree was simulated as a single population with constant size (Figure 1.1). Population splits occurred in a single generation, and there was no contact between populations post-split. Population sizes and split times were taken from the estimates in Prado-Martinez et al. (2013). Across all our simulations, we simulated crossover events occurred with the sex-averaged rates from the deCODE genetic map (in assembly NCBI36/hg18 coordinates) (Kong et al., 2002). We then computed diversity and divergence in the same windows used for the real data using tskit (Kelleher et al., 2018; Ralph et al., 2020).

To improve run time, we simulated sister branches in parallel and recorded the final genealogies as tree sequences (Kelleher et al., 2016). Further, neutral mutations were not simulated with SLiM and were added after the fact with msprime. The resulting tree sequences were later joined and recapitated (i.e., we simulated genetic variation in the ancestor of all great apes using the coalescent) using msprime, tskit and pyslim (Kelleher et al., 2016, 2018; Rodrigues & Ralph, 2021). Despite our efforts to improve run time, our simulations of the entire history of the great apes were still incredibly costly (taking over a month to complete in many instances).

In our neutral simulations, we assumed that neutral mutations occurred at a rate of 2×10^{-8} new mutations per generation per site, uniformly across the chromosome. To understand the effects of natural selection on landscapes, we simulated beneficial and deleterious mutations only within exons, assuming that the locations of exons were shared across all great apes (Kronenberg et al., 2018) and using exon annotations from the human reference genome NCBI36/hg18. We varied the proportions of neutral, beneficial and deleterious mutations within exons, but the distribution of fitness effects for both deleterious and beneficial mutations were shared across all apes. In total, we explored 26 different parameter combinations with different simulations (see Table 1.1 and Appendix A.0.1 for the parameter space).

To simulate local variation in mutation rates along the chromosome, we used the neutral genealogy we simulated with SLiM (and recapitated with msprime) and stripped all existing mutations from it. Using this genealogy, we added neutral mutations back with varying levels of (neutral) mutation rate variation along the chromosome (using msprime). We built mutation rate maps by sampling mutation rates for each 1Mb window independently from a normal distribution with mean 2×10^{-8} and standard deviation chosen from $\frac{\sigma}{2 \times 10^{-8}} = \{0.010, 0.017, 0.028, 0.046, 0.077, 0.129, 0.215, 0.359, 0.599, 1.000\}.$

Regime	Neutral	Deleterious only	Beneficial only	Both
Proportion of deleterious mutations	0%	10% - 70%	0%	10% - 70%
Proportion of beneficial mutations	0%	0%	0.005% - 0.5%	0.005% - 0.5%
Deleterious DFE	_	Gamma distributed with $\bar{s} = \{-0.015, -0.03\}$ and $\alpha = 0.16$	_	Gamma distributed with $\bar{s} = \{-0.015, -0.03\}$ and $\alpha = 0.16$
Beneficial DFE	_		Exponentially distributed with $\bar{s} = \{0.01, 0.005\}$	Exponentially distributed with $\bar{s} = \{0.01, 0.005\}$

Table 1.1. Range of parameters explored in the simulations. Non-neutral mutations were only allowed within exons. "DFE" refers to the distribution of fitness effects. Gamma distribution was parameterized with shape α and mean $\bar{s}=\alpha/\beta$, where β is the rate parameter.

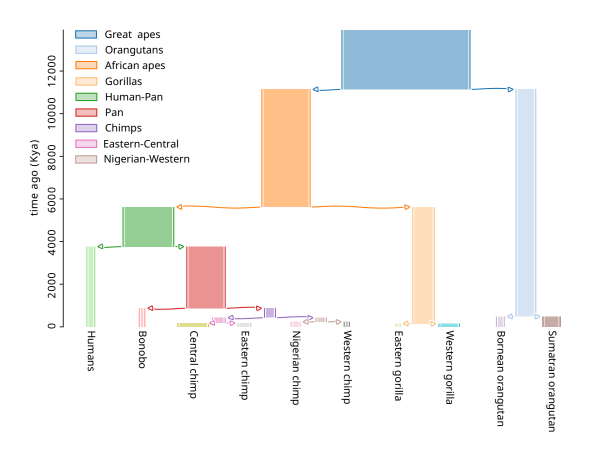


Figure 1.1. Range of parameters explored in the simulations. Arrows indicate population splits. Branch widths are proportional to population size. For example, the population size was 125,089 for the great apes branch and 7,672 for the humans branch. Figure was produced using *demesdraw* (Gower et al., 2022).

1.2.3 Visualizing correlated landscapes of diversity and

divergence. To compare landscapes of diversity and divergence along chromosomes, we computed the Spearman correlation between the landscapes across windows within a chromosome. Because of computational constraints, we focus on chromosome 12. Chromosome 12 is one of the smallest chromosomes in the great apes, there are no major inversions, and it has good variation in exon density and recombination rate. The choice was made blindly before looking at the data, but we found it behaves similarly to other chromosomes (see Figure A.6 through Figure A.27).

We expected landscapes of two closely related species to be more correlated than the landscapes of two distantly related species. Thus, the correlation between any two landscapes of diversity and divergence is expected to depend on distances between them in the phylogenetic tree. We decided to plot our correlations against distance (in generations) between the most common recent ancestor (MRCA) of each landscape. In comparing two landscapes of diversity, this amounts to the total distance between the two tips in the species tree. For instance, the phylogenetic distance dT between diversity in humans and diversity in bonobos is the sum of the lengths of the human, pan and bonobo branches in the species tree (Figure 1.1). In comparing a landscape of diversity to a landscape of divergence, this amounts to the distance between the species of the landscape of diversity and the MRCA of the two species involved in the divergence. For example, dT for the landscapes of diversity in humans and divergence between Sumatran orangutans and eastern gorillas would be the distance between the humans tip and the great apes internal node. dT for the landscapes of divergence between the orangutans and divergence between the gorillas would be the distance between the orangutan and gorilla

internal nodes. Some divergences may share branches in the tree, but these are excluded from our main figures; see subsection A.0.1 and Figure A.2.

1.3 Results

First, we will provide a qualitative view of the landscapes of diversity and divergence in the great apes. Then, we explore the correlations between landscapes in the real data and how they vary depending on phylogenetic distance. To understand the processes that can drive these correlations, we use forward-in-time simulations of the great apes history under different models (e.g., with and without natural selection). Lastly, we describe how genomic features are related to patterns of diversity and divergence in the real great apes data, and we speculate which processes can explain what we see in the data and simulations.

1.3.1 Landscapes of within-species diversity and between-species divergence. There is considerable variation in levels of genetic diversity across the great apes (Figure 1.2). Species may differ in overall levels of diversity due to population size history: species with greater historical population sizes (e.g., central chimps and western gorillas) harbor the most amount of genetic variation (Prado-Martinez et al., 2013). Levels of diversity vary along the chromosome, but do not appear to be strongly structured. Instead, diversity seems to haphazardly fluctuate up and down along the chromosome, and this variation might be attributed to neutral genealogical and mutational processes alone. A notable feature is the large dip in diversity around the 50Mb mark, which is so extensive that it almost erases the differences between-species. This dip coincides with three of the windows with the highest exon density, possibly pointing to the role of selection in shaping genetic variation in those windows.

Levels of between-species genetic divergence also vary along the genome, by an even greater amount in absolute terms. Interestingly, diversity (π) varies (along the chromosome) by about 0.2%, whereas divergence (d_{XY}) varies by more than 0.5%. Because $d_{XY} = \pi^{\rm anc} + rT$ (where $\pi^{\rm anc}$ is diversity in the ancestor, r is the substitution rate and T is the split time between the two species), this excess in variance may be due to the substitution process. Landscapes of divergence which share their most common recent ancestor (e.g., human-Bornean orangutan and bonobo-Bornean orangutan divergences — both colored in red in Figure 1.2A) overlap almost perfectly with each other. Curiously, divergence seems to accumulate faster in the ends of the chromosome, leading to a "smiley" pattern in the landscape of divergence — which is not apparent in the landscape of diversity. That is, with deeper split times, divergence in the ends of the chromosome seem to increase faster than in other regions of the genome (see how the divergences whose MRCA is the great apes look more like a convex parabola than a horizontal line in Figure 1.2A; see also Figure A.1).

In comparing landscapes across species side by side, a remarkable pattern emerges: levels of genetic diversity and divergence along chromosomes have similar peaks and troughs. To get a sense of how strong this observation is, we can compare it to one of the most well studied properties of genomic variation: the correlation between exon density and genetic diversity. We found that the correlation between human diversity and exon density is -0.2 (at the 1Mb scale), but the correlation between levels of diversity in humans and western gorillas is 0.48. Below, we dissect this observation of strong correlation between landscapes across the great apes and discuss the processes that may cause it.



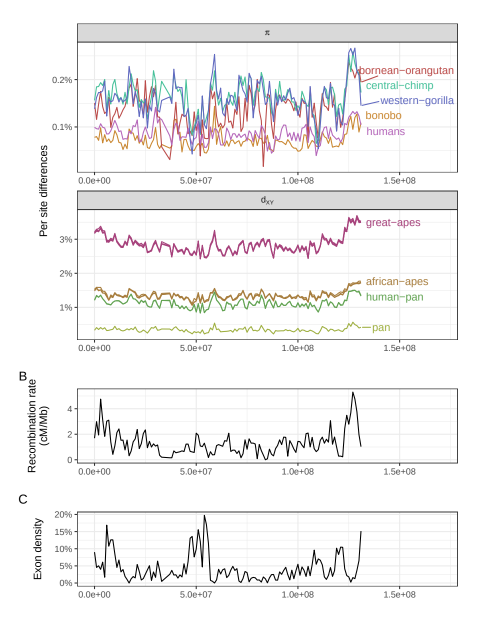


Figure 1.2. A) Landscapes of nucleotide diversity (π) and divergence (d_{XY}) in 1Mb windows along chromosome 12. Nucleotide diversity and divergence (d_{XY}) across 1Mb windows (non-overlapping) of chromosome 12 are displayed above. Lines are colored by species on the left plot and by the most common recent ancestor (MRCA) on the right. Genomic windows with less than 40% of callable sites were masked. Only a subset of the species are displayed for clarity. B) Recombination rate estimates from humans (deCODE map; Kong et al., 2010). C) Exon density along chromosome 12, computed as the percentage of callable nucleotides in a window that fall within an exon.

1.3.2Remarkable correlations between landscapes of diversity and divergence. The landscapes of diversity and divergence are highly correlated across the great apes. To interpret this signal, we first need to understand what processes can cause such correlations, and so first we describe the toy example depicted in Figure 1.3. Both genetic diversity (π) and divergence (d_{XY}) are estimates of the mean time to the most recent common ancestor (multiplied by twice the effective mutation rate). Populations V and W split recently, and so samples from one population may coalesce first with a sample from another population (e.g., samples v_2 and w_1), a pattern called incomplete lineage sorting (ILS). This causes π_V and π_W to be correlated with each other, as they share some ancestral variation (see the branch marked with * in the gene tree). The probability two samples from V coalesce before the split with W is $1 - e^{\frac{-T}{2N_e}}$, where T is the split time and N_e is the effective population size. With longer split times (T), we would expect less incomplete lineage sorting; therefore, split time (T) should be a good predictor of the correlation between two landscapes of diversity (and/or divergence). Thus, we decided to visualize correlations between landscapes of diversity and divergence by computing the phylogenetic distance dT, which is simply the distance in generation time between two statistics. For example, we define $dT(\pi_W, d_{XY}) = 2T_{VWXY} - T_{XY}$. Divergences may share branches by definition (irrespective of split times), as you can see with d_{VX} and d_{XY} (see subsection 1.2.3 for more details). In such cases, our chosen metric dT would not be a good proxy for expected correlations, so we omit such cases from our main figures. See subsection 1.2.3 and Figure A.2 for more on the correlations between landscapes that share branches.

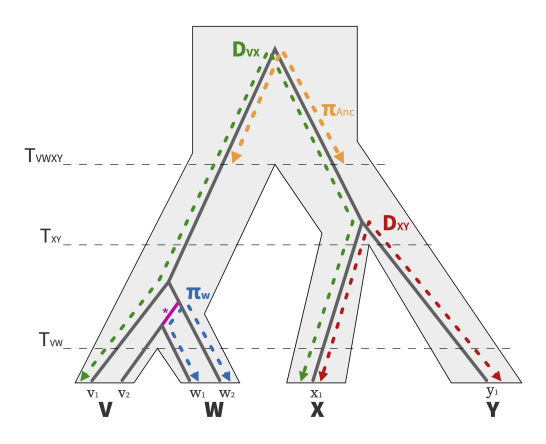


Figure 1.3. Visualizing the relationships between nucleotide diversity and divergence statistics between closely related taxa. A population and gene tree for four populations $(V,\,W,\,X,\,Y)$ are depicted with the light gray polygon and gray solid line, respectively.

Figure 1.4 shows the pairwise correlations between great apes landscapes of diversity and divergence against phylogenetic distance (dT). We see ancestral variation seems to play a role in structuring correlations between landscapes: pairs of species that recently split have their landscapes of diversity highly correlated. Surprisingly, correlations still plateau at around 0.5. We expect ancestral variation to play a minor role when comparing orangutans and chimps, which separated around $60 \times N_e$ generations ago, but their landscapes are still highly correlated. Population size history seems to affect the correlation between landscapes since the weakest correlations involve the landscape of diversity of one of the species with small historical population sizes (i.e., bonobos, eastern gorillas and western chimps).

Correlations between landscapes of divergence and diversity and between landscapes of divergence are also quite high, often surpassing 0.5, and they also decay with phylogenetic distance (dT) (see middle and right most plots in Figure 1.4). In theory, these landscapes can also be correlated due to ancestral variation. To see how ancestral variation can create correlations even between landscapes with no overlap in the tree, consider Figure 1.3: divergence between X and Y and divergence between V and W can each contain contributions from ancestral diversity if lineages have not coalesced in both branches leading from the ancestor. If a particular portion of the genome happens to have higher diversity in the ancestor, it will also have higher divergence. Since this correlation is produced by incomplete lineage sorting, it is expected to have a very small effect except when branches are short. As discussed in subsection 1.2.3, two divergences can also be correlated by definition (because they share branches in the tree). For example, when comparing human-Bornean orangutan and gorilla-Bornean orangutan

divergence we expect some correlation because these divergences share the large African apes and orangutan branches in the tree (Figure 1.1). In Figure 1.4 we excluded these comparisons where branches are shared. Such comparisons can be seen in Figure A.2. We found that even these comparisons that share branches have an excess of correlation compared to a theoretical expectation (derived from a simplified neutral model), that is the correlations are above the y = x line in Figure A.2 even for distantly related species.

There are many processes that could maintain landscapes correlated. Above, we discussed how we expect ancestral variation to explain these correlations. The alternative would be to have a process that structures variation along chromosomes which is shared across species. Using forward-in-time simulations, we set out to (i) confirm that ancestral variation alone is not causing landscapes to remain correlated, and (ii) test which process or processes that when shared among a group of species could maintain correlations in similar ways to what we observed in the great apes' data.

1.3.3 Neutral demographic processes. To assess the extent to which ancestral variation alone could explain our observations, we performed a forward-in-time simulation of the great apes' evolutionary history. As expected, the resulting landscapes of diversity and divergence are not well correlated (Figure 1.5). Ancestral variation seems to maintain correlations between some landscapes; for instance, the landscapes of diversity in central and eastern chimps have a 0.61 correlation, the highest across all pairs of comparisons (Figure 1.5A, point a). Nevertheless, correlations between landscapes of diversity and divergence decay quickly with phylogenetic distance to 0. Some distant comparisons are moderately correlated (e.g., the landscape of diversity in Bornean orangutans and divergence

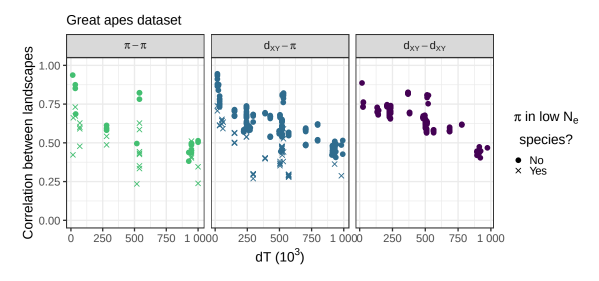


Figure 1.4. Correlations between landscapes of diversity and divergence across the great apes. Each point on the plots correspond to the (Spearman) correlation between two landscapes of diversity/divergence, computed on 1Mb windows across the entire chromosome 12. Correlations were split by type of landscapes compared $(\pi - \pi, \pi - d_{XY}, d_{XY} - d_{XY})$. dT is the phylogenetic distance (in number of generations) between the most common recent ancestor of the two landscapes compared (e.g., the dT for correlation between landscapes of diversity in humans and divergence between eastern gorillas and orangutans is distance between the humans and the great apes nodes in the phylogenetic tree, Figure 1.1). Note that species with low N_e — for which the estimated species N_e was less than 8 \times 10³: bonobos, eastern gorillas and western chimps — have a different point shape. Only comparisons for which the definition of the statistics do not overlap are shown, as explained in subsection 1.2.3.

between central and western chimps have a correlation coefficient of 0.23, see Figure 1.5A, point b), but that seems to be driven by the outlier window around 80Mb. This outlier window has a recombination rate close to 0 (Figure 1.2C), so the average nucleotide diversity over the window has a higher variance because of coalescent noise (see the extreme peaks and valleys in Figure 1.5). Recombination rate variation can create some moderate correlations, but when we look at multiple species at once it becomes clear that the mean correlation goes to 0.

- chromosomes, if this mutation rate wariation. Since mutation rate can vary along chromosomes, if this mutation rate map were shared across species, it would maintain correlations between landscapes over longer periods of time. To assess this, we used our existing simulated neutral history of the great apes and replaced all mutations assuming a common mutation rate map across all great apes: for each window, we drew a mutation rate from a normal distribution with mean 2×10^{-8} (the same as all other simulations) and standard deviation $\mu_{\rm SD}$. We found that a mutation rate map with $\mu_{\rm SD}$ close to $8\% \times 2 \times 10^{-8}$ would be needed to get correlations similar to the data (Figure 1.6C). Although mean correlations look similar to the data, we see that correlations tend to increase slightly with time in the simulations with mutation rate variation. This is expected because windows with higher mutation rate accumulate divergence faster, creating a correlation with mutation rate that gets stronger with time. In the great apes' data, however, we see a slow but steady decrease in correlations with time.
- 1.3.5 GC-biased gene conversion. A prominent feature of the landscapes of divergence in the great apes is the faster accumulation of divergence in the ends of the chromosomes (Figure 1.2). This feature was not present in any of our simulations, so we sought to understand its possible causes. Double

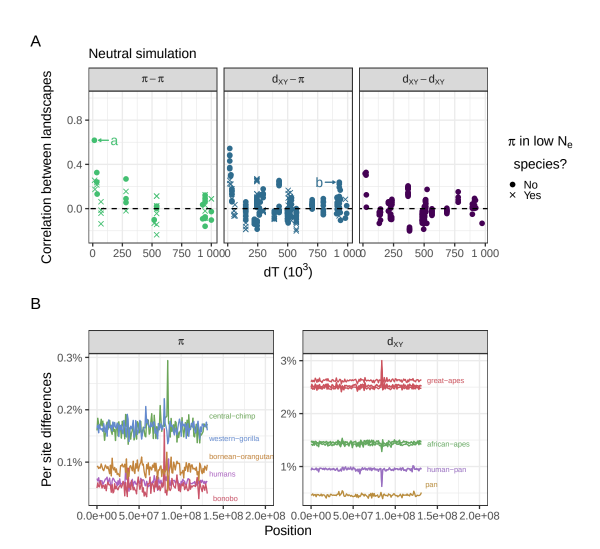


Figure 1.5. Landscapes are not well correlated in a neutral simulation. (A) Correlations between landscapes of diversity and divergence in a neutral simulation. See Figure 1.4 for more details. (B) Nucleotide diversity and divergence along the simulated neutral chromosome. See Figure 1.2A for details.

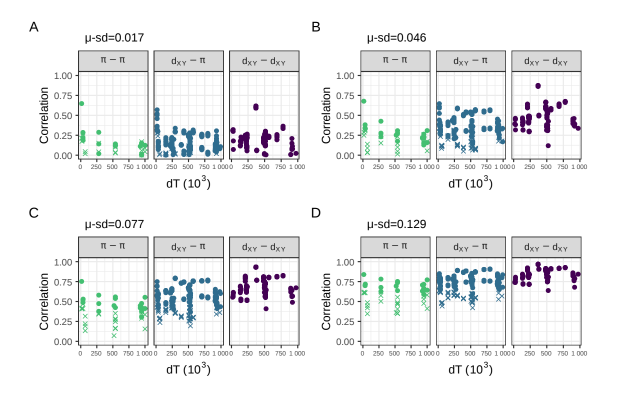


Figure 1.6. Correlations between landscapes of diversity and divergence across the great apes for simulations with variation in mutation rate along the chromosome. Panels A, B, C, and D show different simulations in which we varied the standard deviation in mutation rate between 1Mb windows, in each setting the standard deviation to the mean mutation rate (2×10^{-8}) multiplied by $\mu_{\rm SD}$. Other details are as in Figure 1.4.

strand breaks are more common at the ends of chromosomes, and these can be repaired either by crossover or gene conversion events. GC-biased gene conversion (gBGC), the process whereby weak alleles (A and T) are replaced by strong alleles (G and C) in the repair of double-stranded breaks in heterozygotes, mimics positive selection – in that it increases the probability of fixation of G and C alleles (e.g., Galtier et al., 2009). We suspected gBGC could have caused the increased rate of accumulation divergence in the ends of chromosomes, as has been observed previously (Katzman et al., 2010), and contributes to the maintenance of correlations between landscapes over long time scales.

To tease apart the effects of gBGC on correlated landscapes, we partitioned divergence by mutation type (weak to weak, strong to strong and weak to strong). If correlations are being driven by gBGC, then we would expect the correlation between landscapes of divergence to be stronger for weak to strong mutations. We found that the overall correlations are very similar across mutation types, suggesting gBGC does not play a strong role in structuring the correlations between landscapes (Figure 1.7).

1.3.6 Positive and negative natural selection. Another process whose intensity is likely correlated across all branches in the great apes tree is natural selection. If targets of selection and recombination maps are shared across species, then we would expect both the direct and indirect effects of selection to be shared across branches. It can be difficult to model natural selection in a realistic manner because we do not know precisely which locations of the genome are subject to stronger selection. Nevertheless, exons are expected to have higher density of functional mutations than other places in the genome. Thus, we ran simulations in which beneficial and deleterious mutations can happen only within

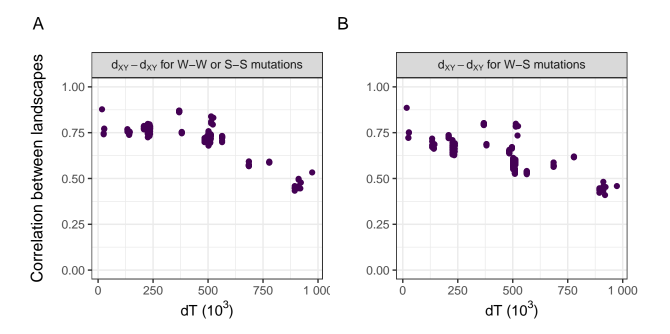


Figure 1.7. Correlations between landscapes of divergence partitioned by site type (W-W/S-S and W-S). W-W sites are sites in which the state did not change between-species (and remained weak which corresponds to A or T). Similar logic applies to S-S sites (S or strong states are G or C). W-S sites are sites in which a new mutation appeared either going from weak to strong or from strong to weak. Note these definitions do not rely on identifying the exact ancestral state, we simply compare the current states in the four species involved (two species per d_{XY} landscape). For example, if by looking at the four species we see the following states A,T,A,T the site would be classified as W-W. If we saw G,A,A,A the site would be classified as W-S. Other details are the same as in the rightmost panel in Figure 1.4.

exons. Using human annotations, we simulated the great apes' history assuming a common recombination map and exon locations. See the landscapes from the simulations in Figure 1.8.

We found that negative selection can slightly increase correlations between landscapes (Figure 1.8A-C). If 30% of all mutations within exons were strongly deleterious (mean selection coefficient $\bar{s}=-0.03$), landscapes would be weakly correlated (Figure 1.8B). The correlations between landscapes rarely surpass 0.5, even with 70% of all mutations within exons being strongly deleterious (Figure 1.8C).

Positive selection, on the other hand, can quickly increase correlations between landscapes. A beneficial mutation rate within exons of $\bar{\mu}_p = 1 \times 10^{-12}$ produced moderate correlations between landscapes (Figure 1.8D). With too much positive selection, correlations can break down because of the contrasting effects of positive selection on diversity and divergence. That is, while positive selection increases fixation rates and hence divergence between-species, its linked effects decrease diversity within the species. This can create negative correlations between landscapes, as can be seen in Figure 1.8F. Note that some correlations between landscapes of diversity and divergence remain high when the divergence is computed between closely related species (e.g., central and eastern chimps). Divergence is $d_{XY} = \pi^{\rm anc} + 2rT$, where $\pi_{\rm anc}$ is diversity in the ancestor, r is the substitution rate and T is the time since species split. Thus, for the divergences in which the two species split recently are dominated by genetic diversity in the ancestor, correlations between $\pi - d_{XY}$ remain high because $d_{XY} \simeq \pi^{\rm anc}$.

Positive and negative selection can work synergistically to produce correlated landscapes that look like the real data. For example, comparing figures Figure 1.8D,G,H which differ in rate of negatively selected mutations μ_n , it is possible to see that the correlations between landscapes start to resemble the real data with more deleterious mutations. Figure 1.8H seems to resemble the data fairly well, with $\pi - d_{XY}$ and $d_{XY} - d_{XY}$ correlations plateauing around 0.5. The $\pi - \pi$ correlations are a bit lower than the real data, however. Recent demographic events can affect genetic diversity and although our simulations are heavily parameterized with respect to the effects of selection, we are not capturing all the variation caused by more realistic demographic models. Figure 1.8D and H look very similar to each other. These have the same amount of positive selection, but the first did not have any negative selection. The major difference between them is that with negative selection there is a more clear separation between the correlations involving low N_e species, similar to what is seen in the data.

1.3.7 Visualizing similarity between simulations and data.

To see how a particular simulation resembles the real data, we can use figures Figure 1.4 and Figure 1.8 to compare how the patterns of all 1260 pairwise correlations between landscapes match the real data. However, it is difficult to assess the fit of the simulated scenarios to real data from such a comparison. Instead, we use principal component analysis (PCA) and create a low dimensional visualization, shown in Figure 1.9, in which each point is a simulation or the real data (shown in yellow). We created this PCA from the matrix 37×1260 in which rows are the simulations and the data, and columns are the pairwise Spearman correlations between landscapes. Unlike in the plots above, here we include the correlations between overlapping landscapes (as detailed in subsection 1.2.3) (Figure 1.9). In PC space, the data most closely resembles a subset of our

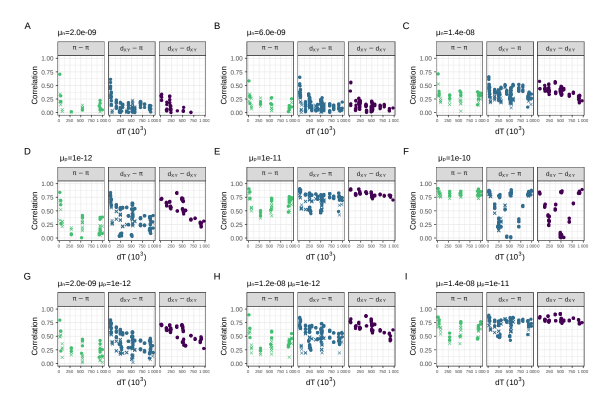


Figure 1.8. Correlations between landscapes of diversity and divergence in simulations with natural selection. (A-C) Simulations with negative selection. (D-F) Simulations with positive selection. (G-I) Simulations with both negative and positive selection. The selection parameters μ_n and μ_p are the rate of mutations in exons with negative and positive fitness effects, respectively. The mean fitness effect was $\bar{s} = -0.03$ for deleterious mutations and $\bar{s} = 0.01$ for beneficial mutations (see subsection 1.2.2 for more details). Compare to Figure 1.4.

simulations with both positive and negative selection ($\bar{\mu}_p = 1 \times 10^{-12}$ and $\bar{\mu}_n = 1.2 \times 10^{-8}$)

Correlations between genomic features and diversity and divergence. Next, we describe how two important genomic features (i.e., exon density and recombination rate) are related to diversity and divergence in the real great apes data set. The correlations between recombination rate and genetic diversity are positive in all great apes (Figure 1.10A). The strongest correlation between genetic diversity and recombination rate is seen in humans, which is unsurprising given our recombination map was estimated for humans. Recent demographic events also seem to impact the strength of the correlation; for example, the correlation between recombination rate and diversity is higher in Nigerian chimps than in western chimps, which have a much lower recent effective population size. We found that diversity is negatively correlated with exon density across all species (Figure 1.10D). Contrary to what we observed with recombination rate, the correlation between exon density and diversity was even stronger in most other apes than in humans. Species with smaller N_e tend to show weaker correlation between diversity and exon density (see Nam et al. (2017) for related findings). A striking feature of the correlations of between-species divergence and genomic features, shown in (Figure 1.10), is that the correlations get stronger with the amount of phylogenetic time that goes into the comparison (i.e., the T_{MRCA}), in a way that is roughly linear with time.

To describe why this increase in correlation with time might occur, we turn to an analytic approach. Genetic divergence (D) in the i^{th} window between two species that split t generations ago can be decomposed as:

$$D_i(t) = \pi_i(t) + R_i t + \varepsilon_i,$$

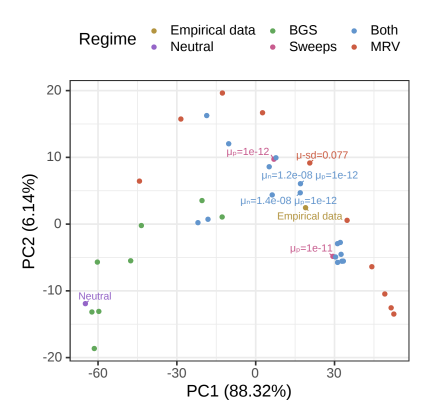


Figure 1.9. PCA visualization of data and simulations. The colors differentiate the empirical data from simulations with different parameters: Neutral refers to the simulation without any selection, BGS refers to simulations with deleterious mutations, Sweeps refers to simulations with beneficial mutations, Both refers to simulations with both beneficial and deleterious, and MRV refers to neutral simulations with variable mutation rates along the chromosome. Principal component analysis (PCA) applied to a matrix with all pairwise correlations between landscapes across the great apes (including $\pi - \pi$, $\pi - d_{XY}$ and $d_{XY} - d_{XY}$ comparisons) for the great apes dataset and simulations (with selection and with mutation rate variation). We excluded simulations with $\mu_p \geq 1 \times 10^{-10}$ from the PCA analysis because PC2 was capturing negative correlations caused by strong positive selection — as seen in Figure 1.8F.

where $\pi_i(t)$ is the genetic diversity in the ancestor at time t, R_i is the substitution rate in the window and ε_i is a contribution from genealogical and mutational noise (which has mean zero). This decomposition follows from the definition of genetic divergence as the number of mutations since the common ancestor, as depicted in Figure 1.3 (see how $D_{VX} = \pi^{\rm anc} + 2RT_{VWXY}$).

The covariance between D(t), the vector of divergences along windows, and a genomic feature X is, using bilinearity of covariance,

$$Cov(D(t), X) = Cov(\pi(t), X) + t Cov(R, X) + Cov(\varepsilon, X).$$
(1.1)

Happily, this equation predicts the linear change of the covariance with time that is seen in Figure 1.10C and perhaps Figure 1.10D. However, caution is needed because the correlation between diversity and the genomic feature $(\text{Cov}(\pi(t), X))$ may be different in different ancestors, and indeed the inferred effective population size is greater in older ancestors in the great apes (Figure 1.1).

Next consider covariances of diversity with recombination rate,

Figure 1.10C. Consulting the equation above, the fact that the covariance between
divergence and recombination rate increases with time can be caused by two
factors (taking X to be the vector of mean recombination rates along the genome):
(i) a positive covariance between substitution rates and recombination rates (Cov(R, X) > 0), and/or (ii) greater genetic diversity in longer ago ancestors $(N_e(t) \text{ larger for larger } t). \text{ It is unlikely that the increase in } N_e \text{ in more ancient}$ ancestors was sufficient to produce the dramatic increase in covariance seen in
Figure 1.10C, since it would require $Cov(\pi(t), X)$ to be far larger in the ancestral
species than is seen in any modern species. On the other hand, there are various
plausible mechanisms that would affect Cov(R, X). One factor that certainly
contributes is the "smile": we found that divergence increases faster near the ends

of the chromosomes where recombination rate is greater, probably in part because of GC-biased gene conversion. Interestingly, positive and negative selection are predicted to have opposite effects here: greater recombination rate increases the efficacy of both through reduced interference among selected alleles, so positive selection would increase substitution rate and hence increase Cov(R, X), while negative selection would decrease Cov(R, X). When considering only the middle half of the chromosome (i.e., excluding the effect of gBGC) (Figure A.5), the covariances between divergence and recombination rate flip to negative, and they continue to decrease over time. Thus, it seems that negative selection is the most important driver of divergence in the middle, whereas gBGC strongly affects the tails of the chromosome.

The covariance of diversity and exon density has a less clear pattern (Figure 1.10C), although it generally gets more strongly negative with time. This decrease could be a result of a negative covariance between substitution rates and exon density and/or an increase in the population sizes of the ancestors (if $\text{Cov}(\nu, X) < 0$, as expected since ν is relative diversity and X is now exon density). As before, positive selection in exons would be expected to produce a positive covariance between exon density and substitution rate, while negative selection would produce a negative covariance. It is hard to determine a priori which is likely to be stronger, because although negative selection is thought to be much more ubiquitous, a small amount of positive selection can have a strong effect on substitution rates. The fact that covariance generally goes down with time suggests that negative selection (i.e., constraint) is more strongly affecting substitution rates.

It is at first surprising that the correlations between exon density and divergence go up with time, but the covariances go down with time (Figure 1.10E,F). However, correlation is defined as $Cor(D_t, X) = Cov(D_t, X)/SD(D_t)SD(X)$. Thus, if the variance in divergences increases over time the correlations will decrease over time. Indeed, we see this happening as gBGC increases divergences on the ends of the chromosome faster than in the middle, leading to an increase in variance of divergence along the genome. This also explains why correlations of landscapes of very recent times are very noisy, but covariances are not. Indeed, the patterns are clearer when we exclude the tails of the chromosome (Figure A.5): there is only a modest increase in the correlation between exon density and divergence over time and the covariances go down with time more linearly.

1.4 Discussion

A central goal of population genetics is to understand the balance of evolutionary forces at work in shaping the origin and maintenance of variation within and between-species (Lewontin, 1974). While the field has been historically data-limited, with the current flood of genome sequencing data, we are poised to make progress on such old questions. Over the past decades, an important lever in understanding the relative impact of genetic drift versus selection in shaping genomic patterns of variation has been to examine the relationship between levels of diversity and genomic features, such as recombination rate and exon density. The overarching observation has been that regions of reduced crossing over generally harbor less variation than regions of increased crossing over in many but not all species (e.g., Begun & Aquadro, 1992; Corbett-Detig et al., 2015). This observation is consistent with a role for linked selection shaping patterns of variation in recombining genomes, but the relative contributions of deleterious and beneficial mutations is still largely unknown. Indeed, it seems likely that some

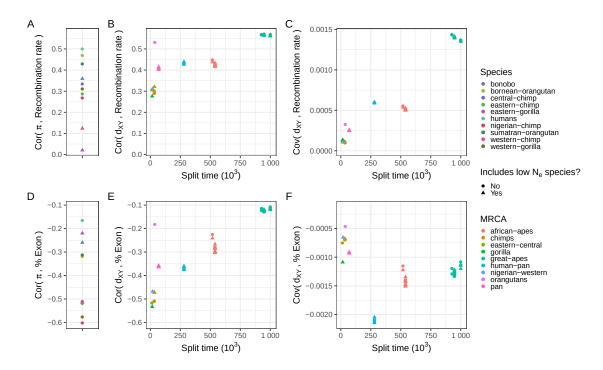


Figure 1.10. Correlations and covariances between landscapes of diversity and divergence and annotation features in the real great apes data. Exon density and recombination rates were obtained as detailed in Figure 1.2. Split time is the time distance between the two species involved in the divergence. Points are colored by the species of within-species diversity (π) in plots A and D. In plots B,C,E,F, the points are colored by the most common recent ancestor of the species for which between-species divergence was computed. Species with low N_e — for which the estimated species N_e was less than 8 × 10³: bonobos, eastern gorillas and western chimps — have a different point shape.

complex mixture of both processes shapes variation in natural populations (Kern & Hahn, 2018).

In this paper, we moved beyond genetic diversity within a single species to look at how divergence between closely related species changes with time and how this correlates with genomic features. Previous studies (e.g., Stankowski et al., 2019) looked at similar patterns (in monkeyflowers) and found strong correlations between landscapes of diversity and divergence between related species, despite deep split times. Landscapes of closely related species can remain correlated for two main reasons (i) shared ancestral variation or (ii) shared heterogeneous process. If two species recently split, their landscapes of diversity are expected to be correlated due to shared ancestral variation. If the process that structures genetic diversity along chromosomes is heterogeneous and somewhat shared between-species, then their landscapes are expected to remain correlated over longer periods of time. For example, if the effects of selection are concentrated in the same genomic regions in two species, then their landscapes of diversity will be correlated. By incorporating information from multiple species at once, we are able to pool information across species and thus increase our power to disentangle the role of different evolutionary forces. Patterns across multiple species are more likely to be robust to the idiosyncrasies of any one species, such as demographic history. For instance within-species metrics can be confounded by demography: demographic events can create spurious troughs of diversity (Simonsen et al., 1995) or exacerbate the effects of background selection on diversity (Torres et al., 2018). However, correlations between landscapes can only be produced by shared ancestral variation or shared heterogeneous process.

In the great apes, we found that landscapes of within-species diversity and between-species divergence are highly correlated across the phylogeny.

Those correlations are often stronger than those that have been historically used as evidence for the effects of selection on genetic variation. For example, the correlation between genetic diversity in humans and exon density is -0.2, yet the correlation between diversity in humans and diversity in western gorillas is 0.48. This stronger correlation may not be entirely due to shared landscape of selection — it may also be a result of shared ancestral variation (i.e., incomplete lineage sorting), mutation rate variation, and/or GC-biased gene conversion. To understand how much of the correlation between landscapes can be attributed to ancestral variation, we performed extensive simulations of the great apes' evolutionary history, and found that ancestral variation explains very little of the correlations we observed. Thus, a shared heterogeneous process seems to be needed to explain the data.

Two neutral processes can be heterogeneous along the genome and shared across species: GC-biased gene conversion and mutation. GC-biased gene conversion (gBGC) is thought to be an important factor in shaping levels of variation in humans (Chen et al., 2007; Glémin et al., 2015; Pouyet et al., 2018), and it has similar effects to those of natural selection. However, if gBGC were a major driver of correlations we would expect to see a difference in overall levels of correlation between different classes of substitution, and we do not (Figures 1.7 and A.4). As such gBGC seems to be a minor contributor to the correlations we observe, although it does seem to be leading to increased substitution rates near the telomeres (where divergences are increasing roughly 5% faster; see

Figure 1.2 and Figure A.1). In birds, an excess of divergence near telomeres has been attributed to meiotic drives (Ellegren et al., 2012).

When the history of the great apes is simulated with a shared heterogeneous mutation map, correlations between landscapes do emerge. These were as strong as seen in the data when the rates were drawn from a normal distribution with a standard deviation of the mutation rate of at least a 7.7% of the mean mutation rate. However, our mutation map was perfectly shared among was species in our simulations, so it is possible that a mutation map which changes over time might move closely to match the data. Mutation rate varies along the human genome and T. C. A. Smith et al. (2018) estimated the standard deviation of de novo mutation rate in humans at the 1Mb scale to be above 25% (with respect to the mean). However, this prior estimates of variation in de novo mutation rate did not take into account differences in callability along genomes – due to the fact that genomic regions vary in how well they can be genotyped with short-read data – which can bias inference. Our simulations showed a facet of shared mutational heterogeneity along the genome that we do not observe in real data: with variable mutation rate correlations increase over time, whereas in the real data they decrease. It is unknown how conserved mutation rate heterogeneity is across the great apes, so it remains to be seen how an evolving heterogeneous mutation rate map affects landscapes of diversity and divergence. A major driver of mutation rate variation stems from CpG dinucleotides, which have much higher mutation rates than other sites (Agarwal & Przeworski, 2021; Hodgkinson & Eyre-Walker, 2011; Nachman & Crowell, 2000). Nevertheless, when we partitioned the landscapes of divergence by mutation types, we did not see an excess of correlation between landscapes

with mutations that can be affected by CpG-induced mutation rate variation (Figures 1.7 and A.4).

Natural selection can also structure genetic variation heterogeneously along the genome. In simulations, both positive and negative selection are needed for the correlations between landscapes to resemble the data. We chose exons to be the targets of selection in our simulations. Exons cover about 1% of the human genome, but in reality selection is known to affect non-coding regions as well. For example, highly conserved noncoding sequences have long been identified and characterized as functional (Bejerano et al., 2004; Katzman et al., 2007; Siepel et al., 2005). Therefore, we might expect a more realistic model to have the same amount of selection (in terms of total influx of selected mutations), but spread out over a somewhat wider region of the genome since we have omitted such sites. While that is so, conserved noncoding sequences generally occur close to coding regions of the genome. By examining the correlations between landscapes (summarized in Figure 1.9), we found that the best fitting simulation is the one with a beneficial mutation rate within exons of 1×10^{-12} and deleterious rate within exons of 1.4×10^{-8} . Our results largely agree with previous studies which found that positive selection is necessary to explain reduction in genetic diversity surrounding genes in the great apes (Nam et al., 2015, 2017).

Another way we might characterize our simulations is through examination of substitution processes. In our best fitting simulation, we get a fixation rate of beneficial mutations of around 1×10^{-9} per generation per exon base pair, what amounts to around 10% of the fixations within exons (along the human lineage) and about one new fixation of a beneficial mutation every 250 generations. Total fixation rate is decreased by around 55% relative to the rate in our neutral

simulation due to the constant removal of deleterious mutations within exons. Indeed, previous studies (Boyko et al., 2008) have estimated that around 10% of amino acid differences between humans and chimpanzees were caused by positive selection, strikingly similar to our best fitting simulation. Furthermore, we would expect to see the fixation of around 16 beneficial mutations in the past 4000 generations, which is close to the number of hard sweeps genome scans for selections have found in humans over this same time period (Schrider & Kern, 2016, 2017). Our best fitting simulation with selection assumes that 70% of mutations within exons are deleterious, similar to estimates from the site frequency spectrum (Boyko et al., 2008; Huber et al., 2017; B. Y. Kim et al., 2017). Thus while we have not done exhaustive model fitting due to computational constraints, our simulations recapitulate major patterns of variation observed in the genome.

Heterogeneous processes that correlate with a genomic feature will create differences in rates of substitution along the genome that correlate with the genomic feature. As shown in Equation (1.1), this implies that the covariance along the genome between a genomic feature and divergence is expected to increase with time, and the rate of increase is equal to the covariance between that feature and the substitution rate. (It is important to note that varying covariances with ancestral diversity can be a confounding factor, and that the observation applies to covariance, not correlation.) Indeed, the covariance between divergence and recombination rate increases roughly linearly with time (see Figure 1.10C), as expected because the rate of gBGC-induced fixations are correlated with recombination rate. Once this effect is removed (see Figure A.5F), the covariance between exon density and divergence decreases linearly with time, as we would expect due to the effects of negative selection directly removing

deleterious mutations in or near exons. The magnitude of this slope might produce a quantitative estimate of the strength of this effect, although more work is needed to disentangle confounders. It is important to contrast this observation, which applies mostly to the direct effects of selection, to other observations which also include linked effects (as discussed in Phung et al. (2016)).

While it has long been recognized that genetic variation among species might be structured similarly due to shared targets of selection, our results demonstrate that this signal contains important information about the processes at work that has yet to be utilized fully. Here we have used large-scale simulations to demonstrate the combination of forces required to patterns shared divergence and diversity as we observe it in nature, however there is clearly a need for future analytical work that might describe expected correlations across the genome given heterogeneous mutation, recombination, and selection. Further, statistical model fitting, based on theory or simulation is clearly desirable, although our experience suggests that the latter approach would prove computationally expensive.

APPENDIX

SUPPLEMENTAL MATERIAL FOR SHARED GREATAPES

Supplementary material

A.0.1 Correlation between divergences that share branches.

Landscapes of divergence can be correlated by their definition, as they can share part of their histories. In most of our analyses (except for Figure A.2), we do not show the correlations for such cases but below we describe how this sharing would affect correlations (using a simplified theory). For example, in Figure 1.3 d_{VX} and d_{XY} share the branch X; depending on how the length of the branch X compares to the total tree length, these two landscapes are bound to be correlated. Assuming that mutations follow a Poisson process and that coalescences happen instantaneously, we derive the following. There are three non-overlapping parts in the tree between these, the branch from the XY ancestor to X with length $E[\tau_X] = T_{XY}$, the branch from the XY ancestor to Y with length $E[\tau_Y] = T_{XY}$ and the branch from Y to the Y ancestor with length Y and Y are some definition of divergence and assume Y and Y and Y are Y and Y and Y are stored as Y and definition of divergence and assume Y and divergence), then

$$Cov[d_{VX}, d_{XY}] = Cov[\tau_X + \tau_V, \tau_X + \tau_Y]$$

$$= Cov[\tau_X, \tau_X] + \underbrace{Cov[\tau_X, \tau_Y]}_{0} + \underbrace{Cov[\tau_V, \tau_X]}_{0} + \underbrace{Cov[\tau_V, \tau_Y]}_{0}$$

$$= Var(\tau_X) = E[\tau_X] = T_X$$

Therefore,

$$\operatorname{Cor}[d_{VX}, d_{XY}] = \frac{\operatorname{Cov}[\tau_X + \tau_V, \tau_X + \tau_Y]}{\sqrt{\operatorname{Var}[\tau_X + \tau_V] \operatorname{Var}[\tau_X + \tau_Y]}}$$

$$= \sqrt{\frac{\operatorname{Var}[\tau_X]^2}{(\operatorname{Var}[\tau_X] + \operatorname{Var}[\tau_V])(\operatorname{Var}[\tau_X] + \operatorname{Var}[\tau_Y])}}$$

$$= \sqrt{\frac{\operatorname{Var}[\tau_X]}{\operatorname{Var}[\tau_X]} \frac{\operatorname{Var}[\tau_X]}{\operatorname{Var}[\tau_X] + \operatorname{Var}[\tau_Y]}}$$

$$= \sqrt{\frac{T_X}{T_X + T_V} \frac{T_X}{T_X + T_Y}}$$

$$= \sqrt{\frac{p_{d_{VX}} p_{d_{XY}}}{p_{d_{XY}}}}$$

where $p_{d_{VX}} = \frac{T_X}{T_X + T_V}$ is the proportion of d_{VX} that is shared with d_{XY} , and $p_{d_{XY}} = \frac{T_X}{T_X + T_Y}$ is the proportion of d_{XY} that is shared with d_{VX} .

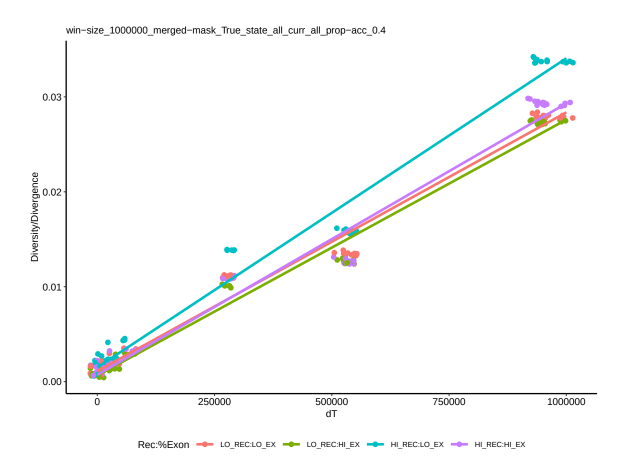


Figure A.1. Effect of exon density and recombination rate on the accumulation of genetic divergence in chromosome 12 with phylogenetic distance. Within-species genetic diversities are shown at dT=0. Mean diversity and divergences were computed for four groups depending on whether they fell or not on the top 90% percentile of recombination rate and exon density.

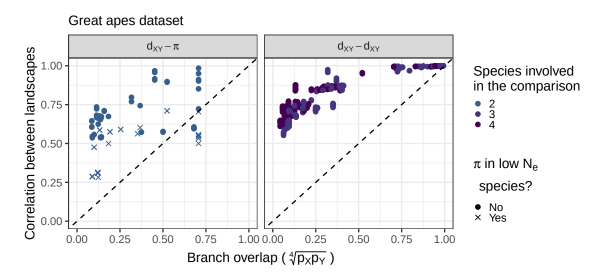


Figure A.2. Correlations between landscapes of diversity and divergence for comparisons with branch overlap. For example, diversity in humans and divergence between humans and bonobos share part of their history. Each point on the plots correspond to the (Spearman) correlation between two landscapes of diversity/divergence, computed on 1Mb windows across the entire genome. Correlations were split by type of landscapes compared ($\pi - d_{XY}, d_{XY} - d_{XY}$). The x-axis is a metric of expected branch overlap between the landscapes. See subsection A.0.1 for more information. Note that species with low N_e (bonobos, eastern gorillas and western chimps) have a different point shape. The colors reflect the number of species involved in the comparison. For example, the comparison between human-western gorilla and eastern chimp-Sumatran orangutan divergences includes four different species. On the other hand, the comparison between human-western gorilla and human-Sumatran orangutan divergences includes just three species.

μ_N	μ_P	$ar{s}_N$	$ar{ar{s}}_P$	Regime	μ_{SD}
0	0	0	0	Neutral	0
0	0	0	0	Variable μ	0.010
0	0	0	0	Variable μ	0.017
0	0	0	0	Variable μ	0.028
0	0	0	0	Variable μ	0.046
0	0	0	0	Variable μ	0.077
0	0	0	0	Variable μ	0.129
0	0	0	0	Variable μ	0.215
0	0	0	0	Variable μ	0.359
0	0	0	0	Variable μ	0.599
0	0	0	0	Variable μ	1
0	1×10^{-12}	0	1×10^{-2}	Beneficial	0
0	1×10^{-11}	0	1×10^{-2}	Beneficial	0
2×10^{-9}	0	-3×10^{-2}	0	Deleterious	0
2×10^{-9}	1×10^{-11}	-3×10^{-2}	1×10^{-2}	Both	0
2×10^{-9}	0	-1.5×10^{-2}	0	Deleterious	0
2×10^{-9}	1×10^{-11}	-1.5×10^{-2}	1×10^{-2}	Both	0
2×10^{-9}	0	-1×10^{-2}	0	Deleterious	0
2×10^{-9}	1×10^{-12}	-1×10^{-2}	5×10^{-3}	Both	0
2×10^{-9}	1×10^{-12}	-1×10^{-2}	1×10^{-2}	Both	0
2×10^{-9}	0	-3×10^{-3}	0	Deleterious	0
2×10^{-9}	1×10^{-12}		5×10^{-3}	Both	0
2×10^{-9}	1×10^{-12}	-3×10^{-3}	1×10^{-2}	Both	0
6×10^{-9}	0	-3×10^{-2}	0	Deleterious	0
6×10^{-9}	1×10^{-11}	-3×10^{-2}	1×10^{-2}	Both	0
6×10^{-9}	0	-1.5×10^{-2}	0	Deleterious	0
6×10^{-9}	1×10^{-11}	-1.5×10^{-2}	1×10^{-2}	Both	0
1.2×10^{-8}	0	-3×10^{-2}	0	Deleterious	0
1.2×10^{-8}		-3×10^{-2}		Both	0
1.2×10^{-8}	1×10^{-11}	-3×10^{-2}	1×10^{-2}	Both	0
1.2×10^{-8}	1×10^{-11}	-3×10^{-2}	1×10^{-2}	Both	0
1.4×10^{-8}	0	-3×10^{-2}	0	Deleterious	0
1.4×10^{-8}		-3×10^{-2}		Both	0
1.4×10^{-8}				Both	0
1.4×10^{-8}	1×10^{-11}	-3×10^{-2}	1×10^{-2}	Both	0

Table A.1. Parameter space explored with simulations. μ_N and μ_P are the rates of mutations under negative and positive selection, respectively. \bar{s}_N and \bar{s}_P and the mean fitness effects of negatively and positively selected mutations. μ_{SD} is the scaled standard deviation of the mutation rate map. See Table 1.1 and subsection 1.2.2 for more details.

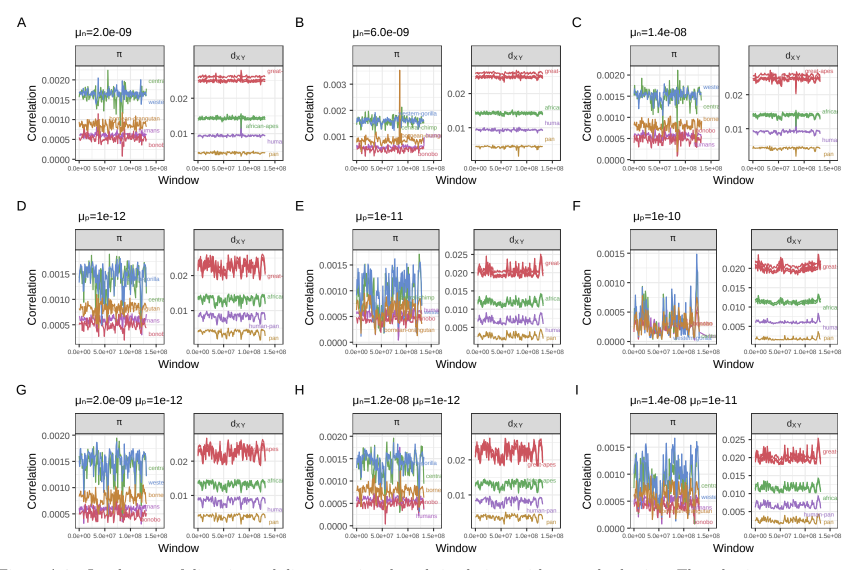
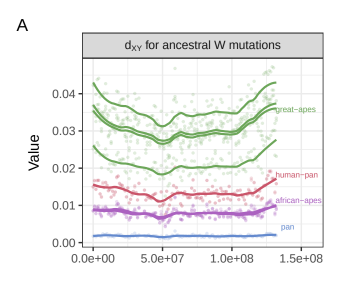


Figure A.3. Landscapes of diversity and divergence in selected simulations with natural selection. The selection parameters μ_n and μ_p are the rate of mutations in exons with negative and positive fitness effects, respectively. The mean fitness effect was $\bar{s} = -0.03$ for deleterious mutations and $\bar{s} = 0.01$ for beneficial mutations (see subsection 1.2.2 for more details). Other details are as in Figure 1.2.



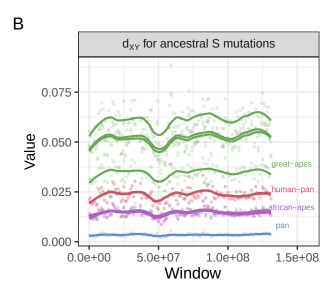


Figure A.4. Landscapes of divergence partitioned by allele state in the ancestor. Ancestral states were assumed to be the same as seen in rhesus macaques (RheMac2), and sites not called in macaques were not used. d_{XY} for W sites is simply the mean pairwise differences between samples in species X and Y per ancestral W sites (A/T). Similar reasoning applies for d_{XY} for S ancestral sites, but only considering (G/C) sites. Points were colored by the most common recent ancestor of the two species compared in each divergence. Lines were fitted using local linear regression. Note that for ancestrally weak mutations (A) there is an increase in divergence at the ends of the chromosomes, but that is not seen for ancestrally strong mutations (B).

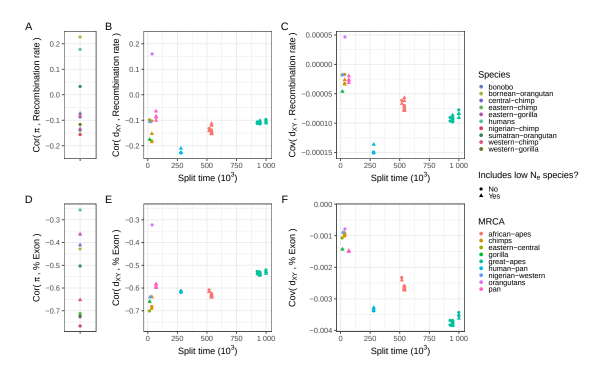


Figure A.5. Correlations and covariances between landscapes of diversity and divergence and annotation features in the real great apes data. Only windows in the middle half of chromosome 12 were included. Compare to Figure 1.10.

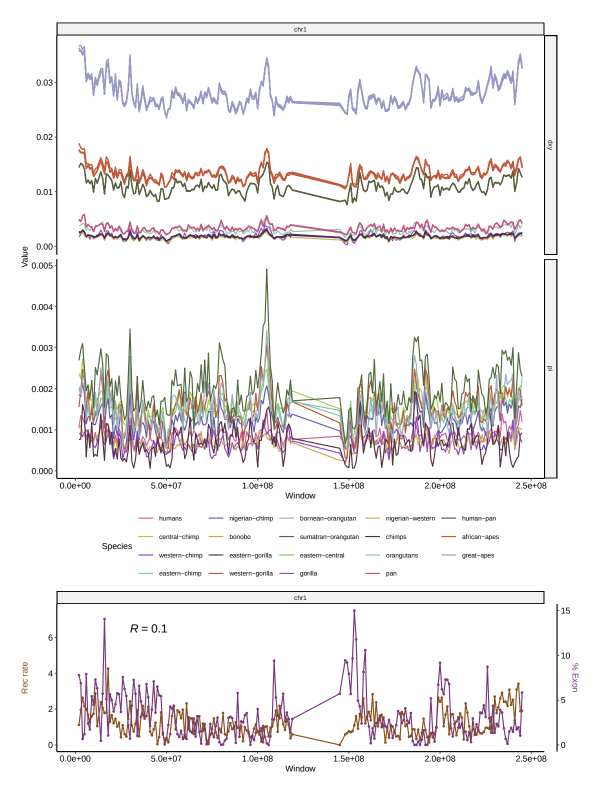


Figure A.6. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 1. See Figure 1.2 for more details.

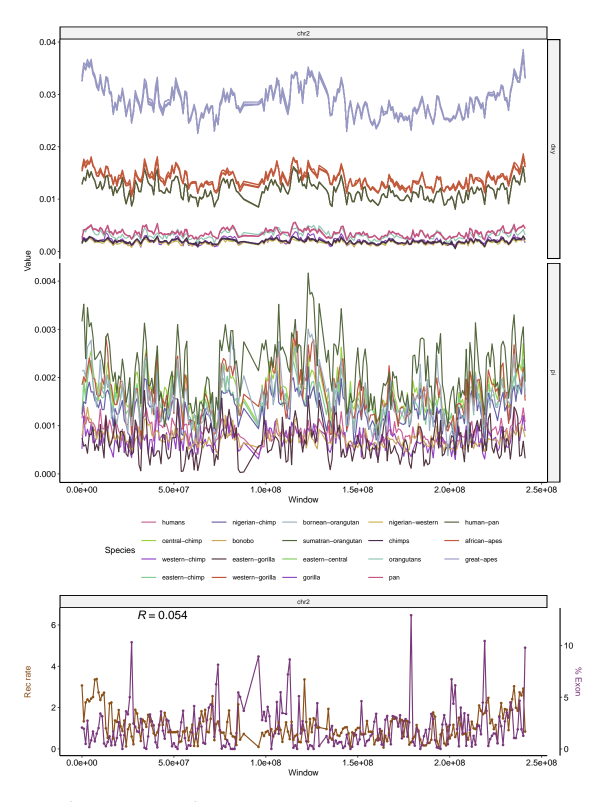


Figure A.7. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 2. See Figure 1.2 for more details.

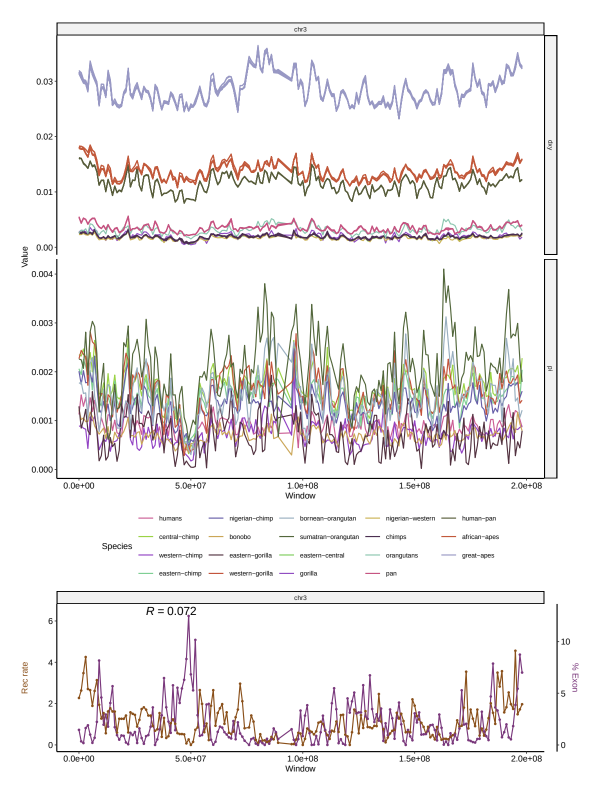


Figure A.8. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 3. See Figure 1.2 for more details.

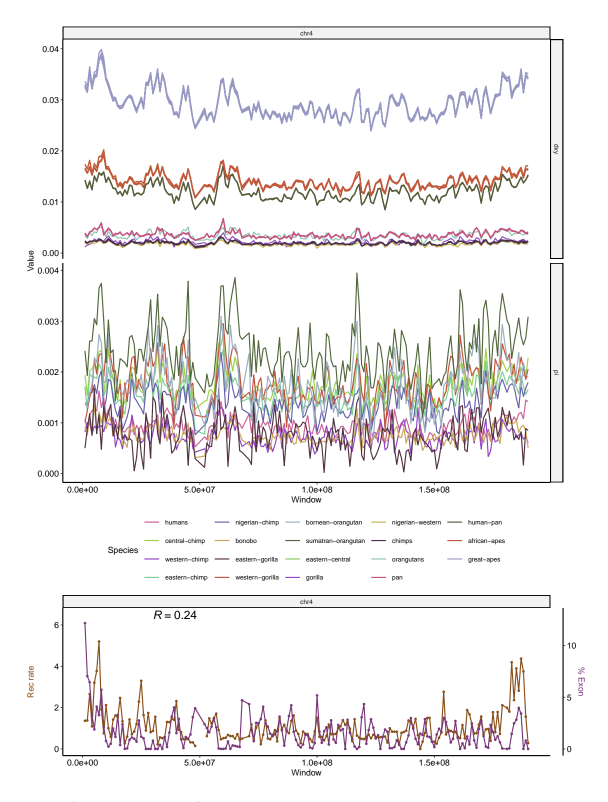


Figure A.9. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 4. See Figure 1.2 for more details.

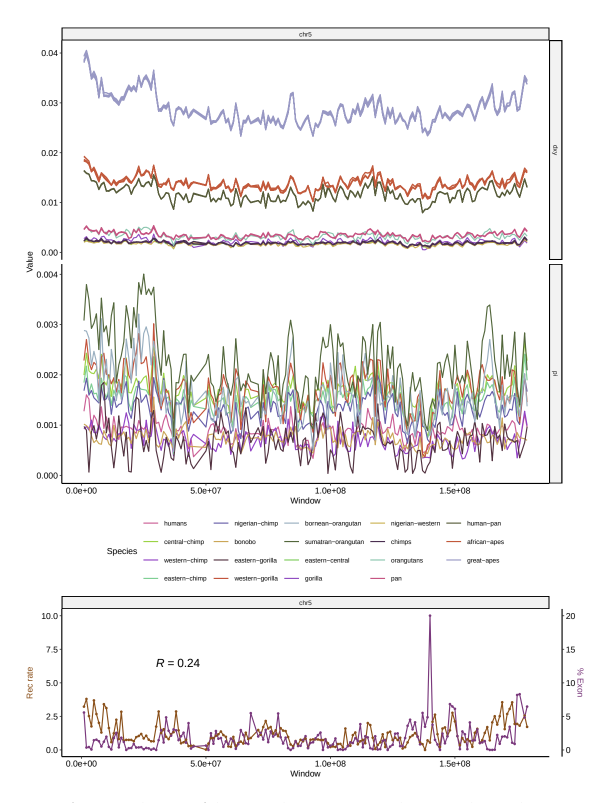


Figure A.10. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 5. See Figure 1.2 for more details.

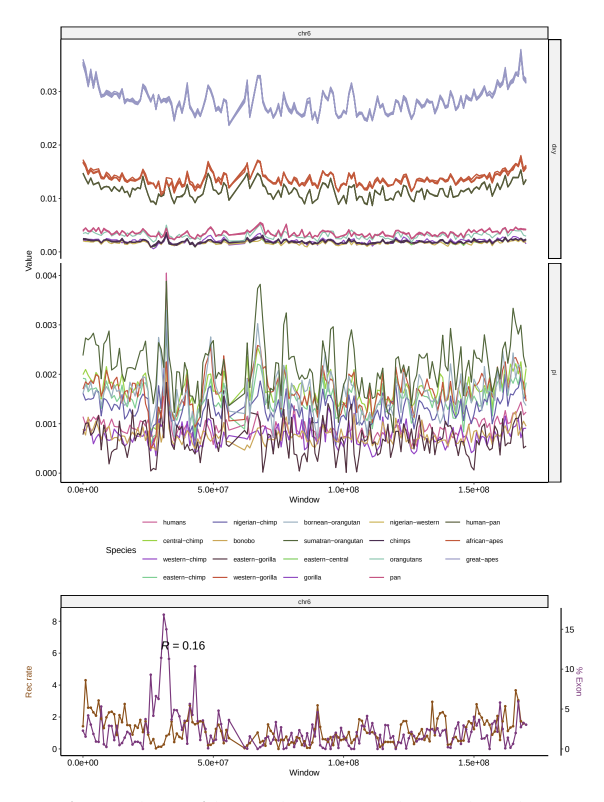


Figure A.11. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 6. See Figure 1.2 for more details.

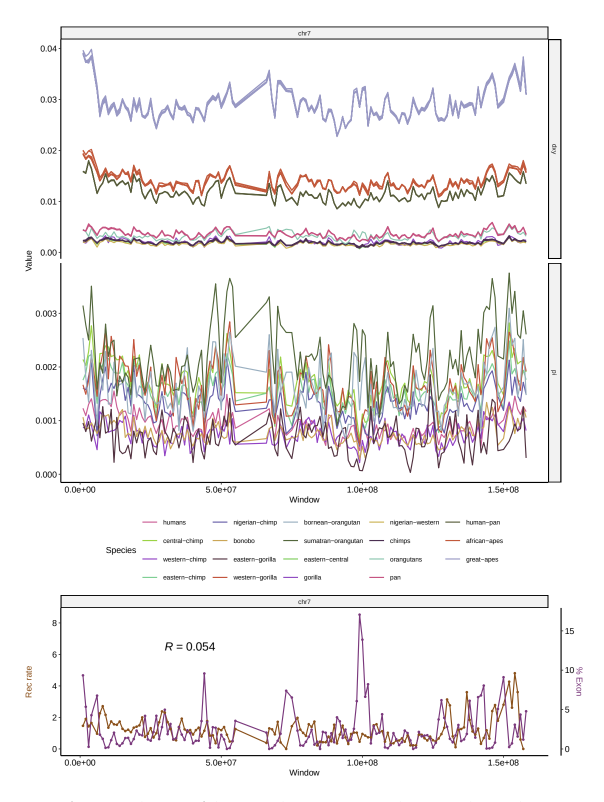


Figure A.12. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 7. See Figure 1.2 for more details.

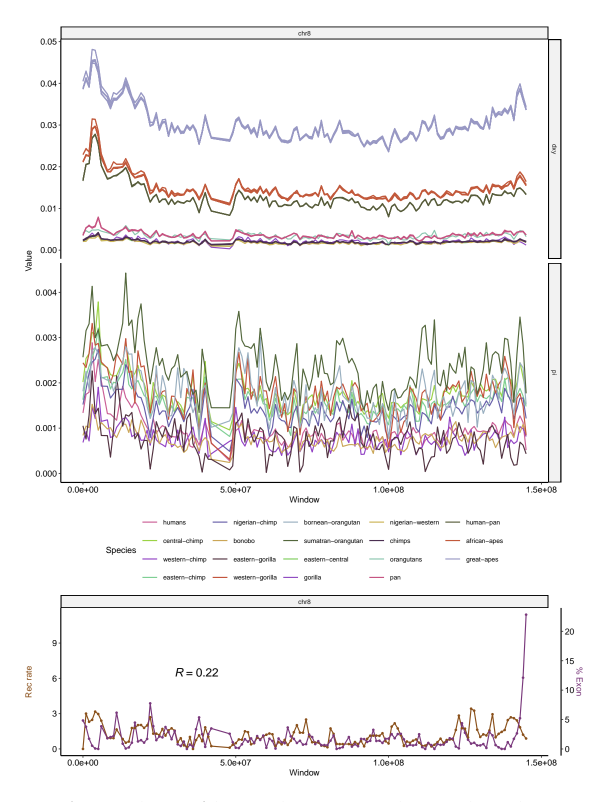


Figure A.13. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 8. See Figure 1.2 for more details.

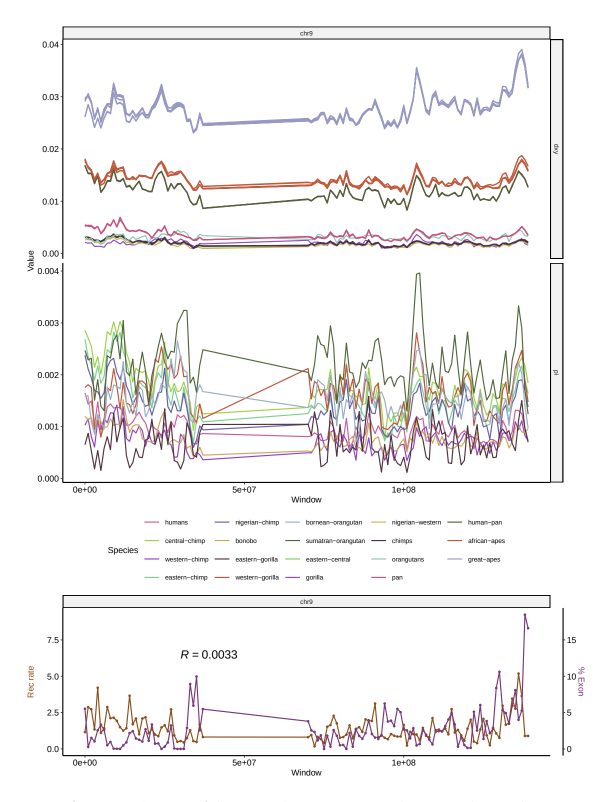


Figure A.14. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 9. See Figure 1.2 for more details.

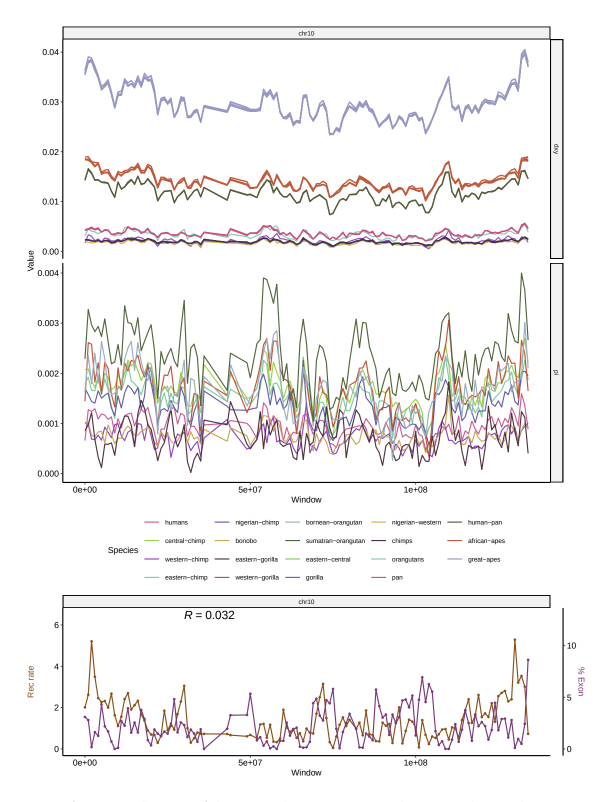


Figure A.15. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 10. See Figure 1.2 for more details.

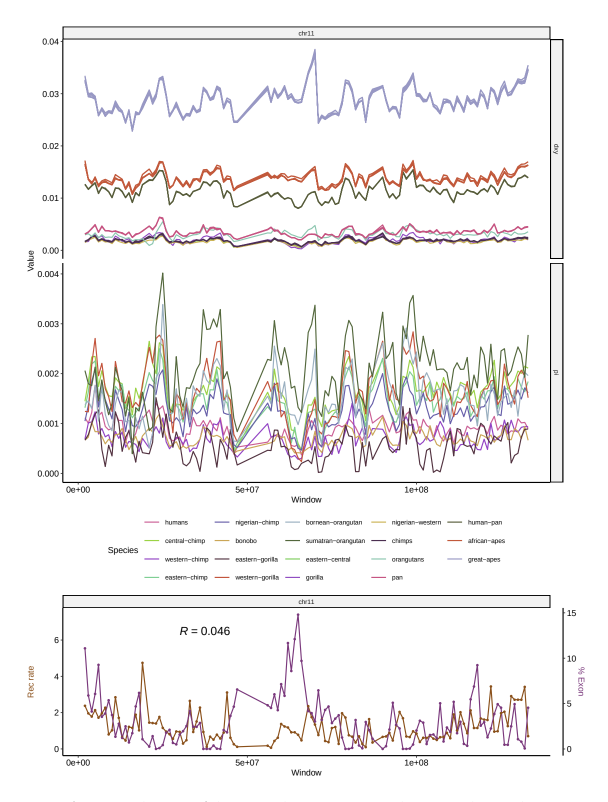


Figure A.16. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 11. See Figure 1.2 for more details.

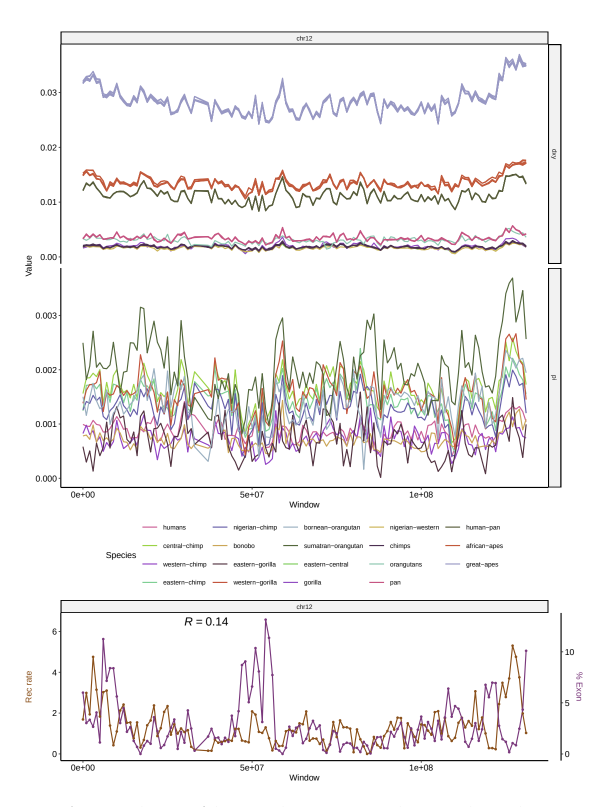


Figure A.17. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 12. See Figure 1.2 for more details.

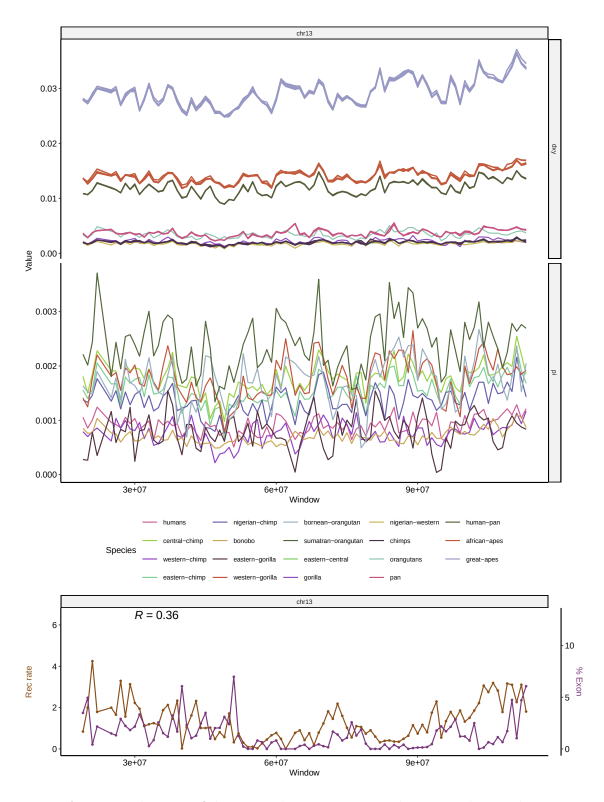


Figure A.18. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 13. See Figure 1.2 for more details.

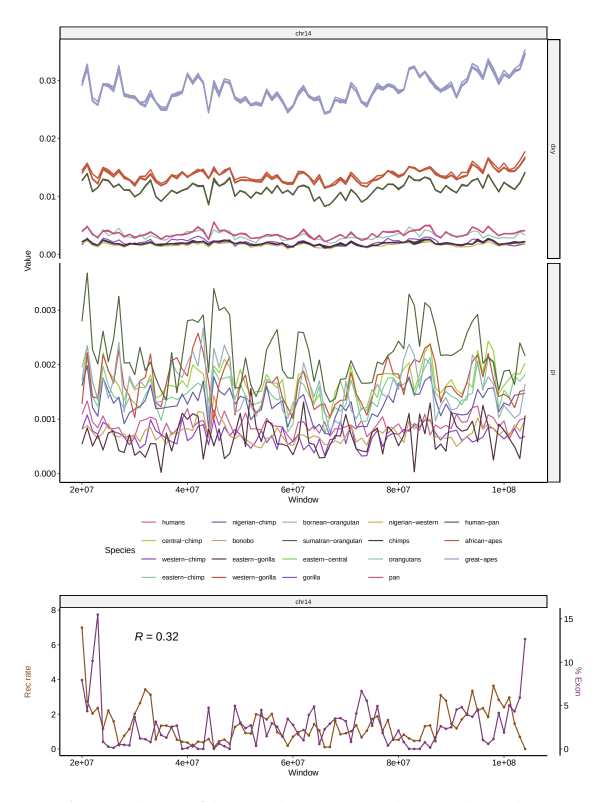


Figure A.19. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 14. See Figure 1.2 for more details.

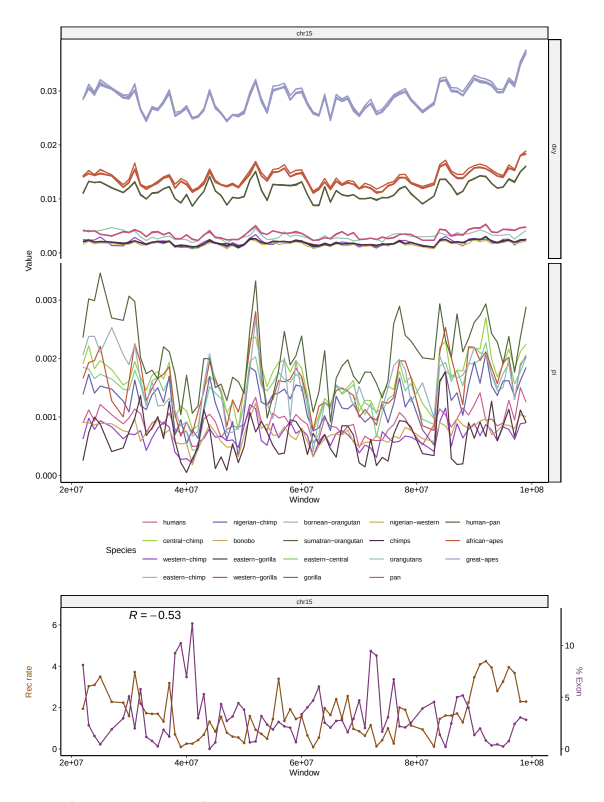


Figure A.20. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 15. See Figure 1.2 for more details.

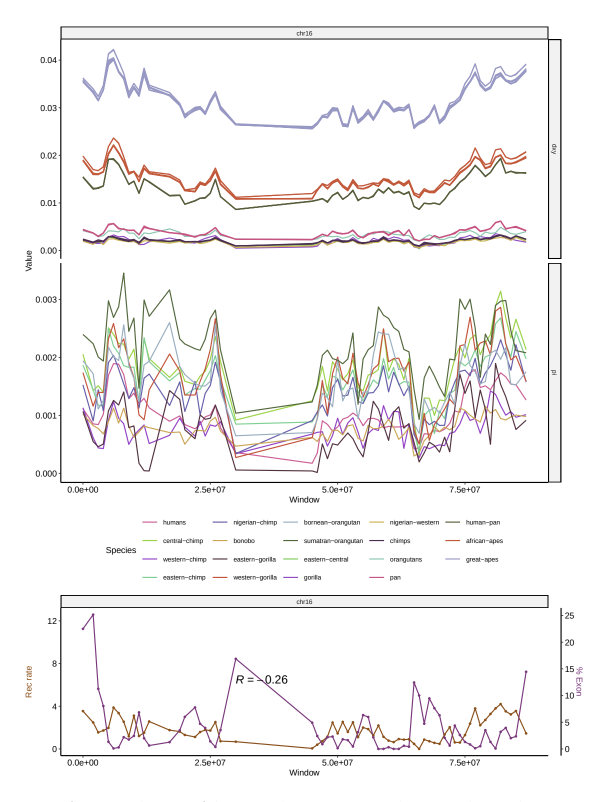


Figure A.21. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 16. See Figure 1.2 for more details.

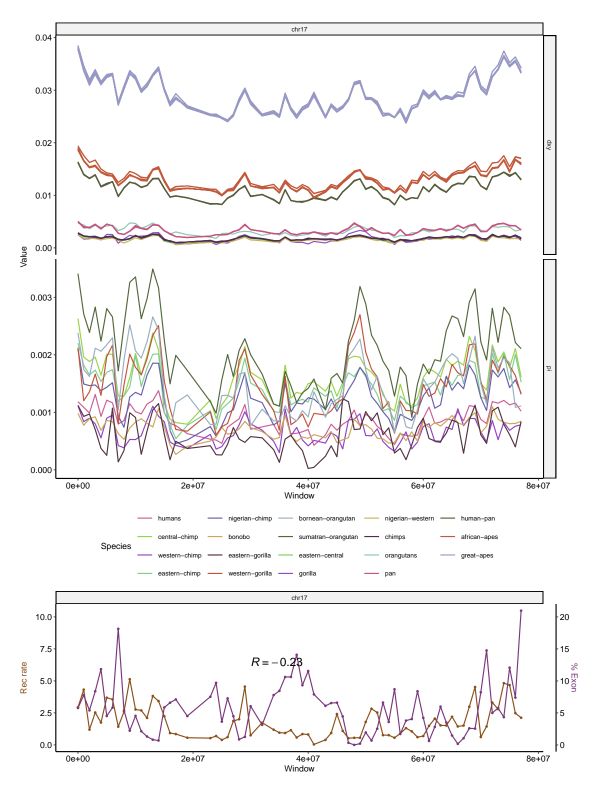


Figure A.22. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 17. See Figure 1.2 for more details.

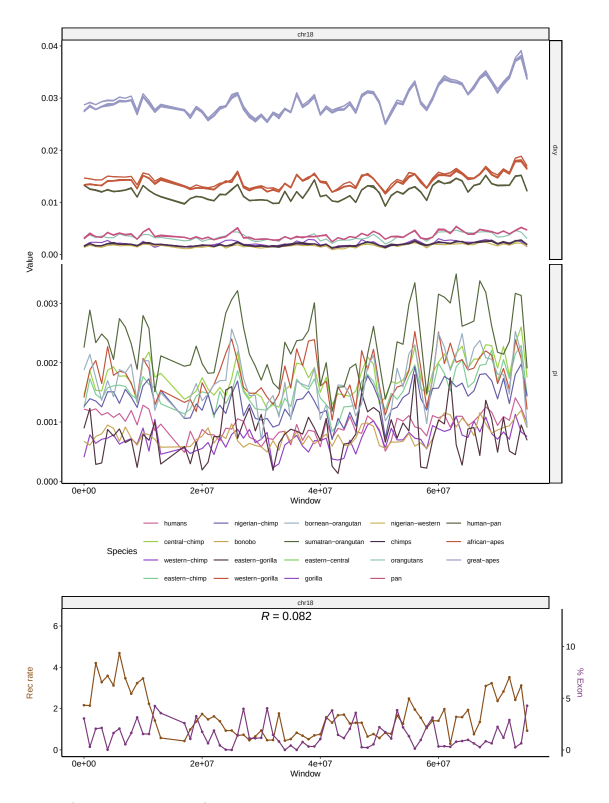


Figure A.23. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 18. See Figure 1.2 for more details.

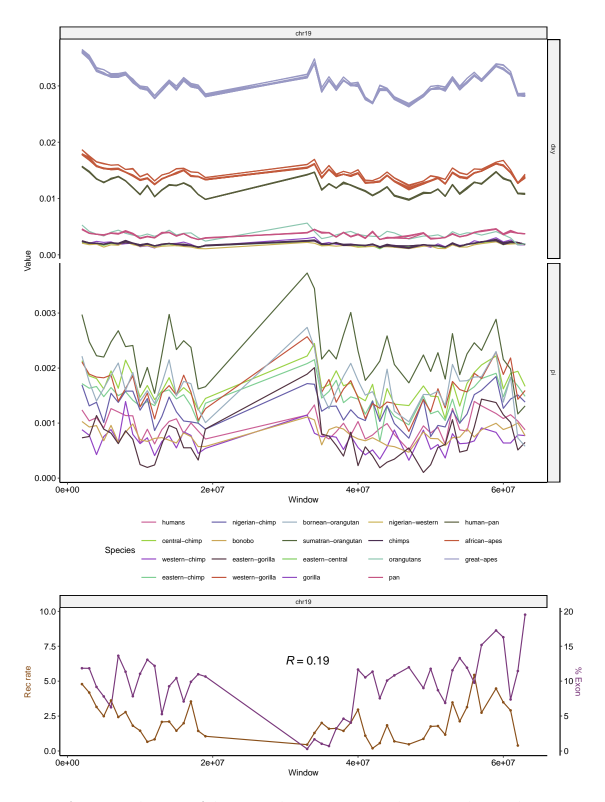


Figure A.24. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 19. See Figure 1.2 for more details.

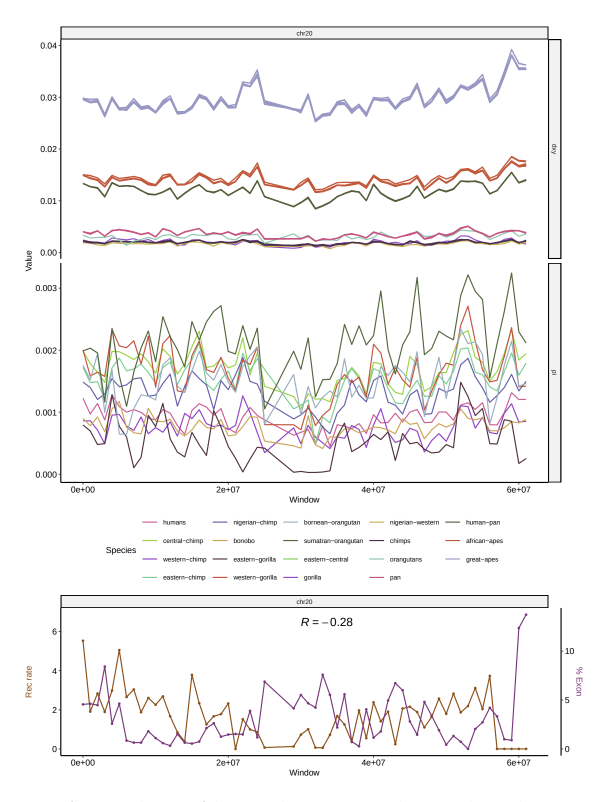


Figure A.25. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 20. See Figure 1.2 for more details.

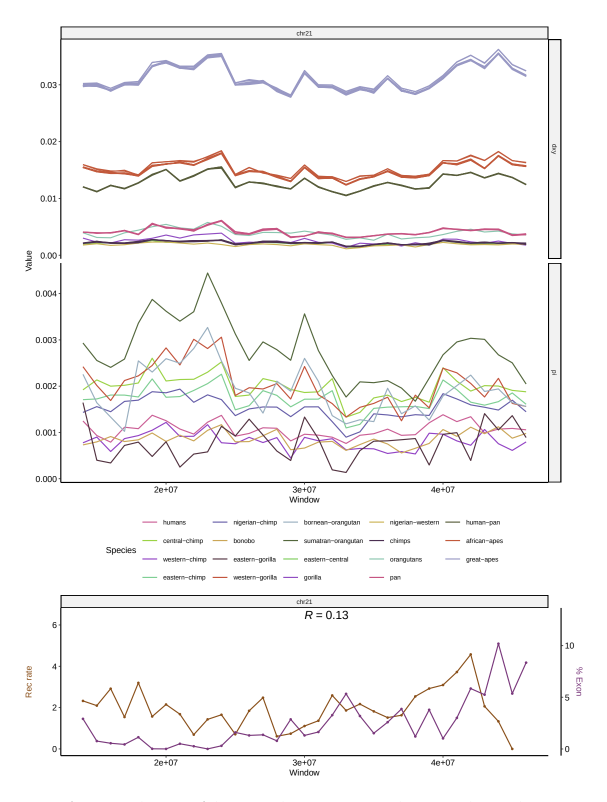


Figure A.26. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 21. See Figure 1.2 for more details.

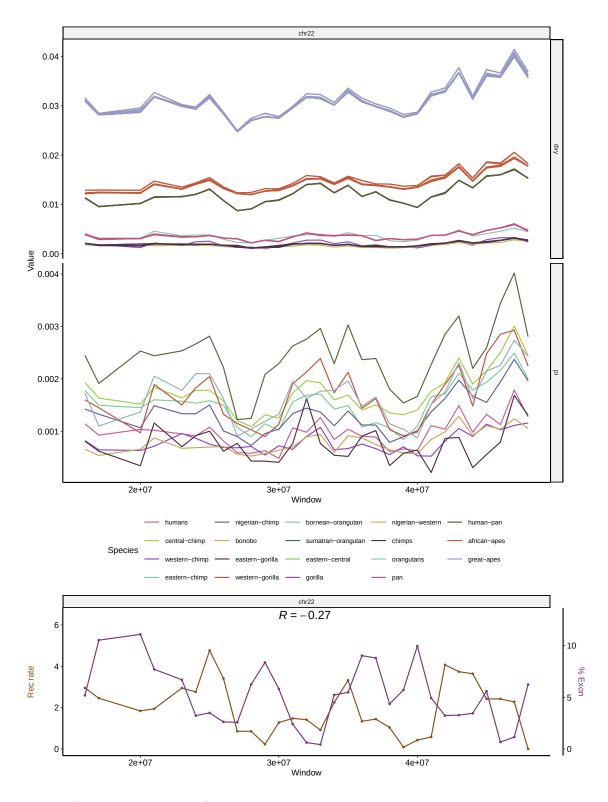


Figure A.27. Landscapes of diversity, divergence, exon density and recombination rate across chromosome 22. See Figure 1.2 for more details.

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