The Impact of Linked Selection in Chimpanzees: A Comparative Study

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Abstract

Levels of nucleotide diversity vary greatly across the genomes of most species owing to multiple factors. These include variation in the underlying mutation rates, as well as the effects of both direct and linked selection. Fundamental to interpreting the relative importance of these forces is the common observation of a strong positive correlation between nucleotide diversity and recombination rate. While indeed observed in humans, the interpretation of this pattern has been difficult in the absence of high-quality polymorphism data and recombination maps in closely related species. Here, we characterize genetic features driving nucleotide diversity in Western chimpanzees using a recently generated whole genome polymorphism data set. Our results suggest that recombination rate is the primary predictor of nucleotide variation with a strongly positive correlation. In addition, telomeric distance, regional GC-content, and regional CpG-island content are strongly negatively correlated with variation. These results are compared with humans, with both similarities and differences interpreted in the light of the estimated effective population sizes of the two species as well as their strongly differing recent demographic histories.

Key words: nucleotide diversity, selection, chimpanzee.

The Pervasive Relationship between Recombination and Variation

The correlation between nucleotide diversity and recombination rate is one of the most prevalent patterns in population genetics, and has been broadly interpreted as evidence for the strong effects of linked selection (see review of Cutter and Payseur 2013). The impact on linked neutral variation with negatively selected sites (termed Background Selection (BGS)), and with positively selected sites (termed Recurrent Hitchhiking (RHH)), is expected to be stronger in regions of low recombination (Begun and Aquadro 1992; Charlesworth et al. 1993). As such, both recurrent positive and recurrent purifying selection will serve to produce this observed relationship. Although both processes are likely at play (see Hudson 1994), and strong arguments have been made for the relative importance of one over the other in particular organisms, the pattern itself has been demonstrated to be remarkably pervasive – having been observed across mammals (e.g., Nachman 1997; Lohmueller et al. 2011), birds (e.g., Rao et al. 2011), insects (e.g., Begun and Aquadro 1992; Stump et al. 2005), fungi (e.g., Cutter and Moses 2011), plants (e.g., Dvorák et al. 1998), and viruses (e.g., Renzette et al. 2016). Although the observation is open to interpretation, an undeniable strength of BGS-based arguments is the fact that there is a far greater proportion of newly arising deleterious mutations compared to newly arising beneficial mutations across the genome, a notion already well appreciated in the early literature of the field (Timofeeff-Ressovsky 1940; Muller 1949, 1950; see review of Bank et al. 2014). Thus, the selective removal of such mutations is likely a very common process.

Given the wealth of genomic data and an inherent interest in the potential effects of segregating deleterious mutations, humans have been a centrally important organism of study in this area. Perhaps most relatedly, Hellmann et al. (2005) found recombination rate to be the best predictor of human diversity levels, as well as a strong predictor of human-chimpanzee divergence. Exploring scenarios of RHH, BGS, as well as the possibility of a mutagenic effect of recombination, the authors were unable to well-discern between these models, largely
owing to insufficient polymorphism data and recombination rate estimates in their outgroup (chimpanzees). Further, though simulations suggested a slightly better fit under models of background selection, no predictive correlation was observed in humans between diversity and gene content, an observation somewhat at odds with a BGS-based explanation.

The subsequent decade has witnessed important advances in our understanding of chimpanzee genomics, making it possible to revisit these important results in greater depth. Most previous sub-genomic comparative work has suggested that humans harbor two- to four-fold lower levels of intra-species diversity (Kaessmann et al. 1999; Deinard and Kidd 1999; Jensen-Seaman et al. 2001; Satta 2001; Stone et al. 2002; Chimpanzee Sequencing and Analysis Consortium 2005)—with the observation generally being interpreted as a somewhat larger long-term effective population size in chimpanzees (Deinard and Kidd 1999; Kaessmann et al. 2001). However, these analyses have been based on a specific set of genes or genomic locations, resulting in ascertainment concerns.

With the availability of whole genome polymorphism data from the PanMap project, and the development of a fine-scale recombination map (Auton et al. 2012), we here visit the question of the pervasiveness of linked selection in the chimpanzee genome on a large scale while avoiding such ascertainment issues. Through a whole-genome analysis of 10 Western chimpanzees, a number of notable observations emerge. First and foremost, recombination rate is a strong predictor of variation at putatively neutral sites across the genome as is exon content.

In order to characterize this pattern and describe underlying predictors of variation in chimpanzees, each chromosome was divided into continuous windows of 1Mb (fig. 1), and the nucleotide diversity \( \pi \) within a window was compared against multiple genomic features, many of which are hypothesized to be either directly or indirectly linked with DNA damage or repair mechanisms (i.e., recombination rate, regional GC-content, gene density, CpG-island density, simple repeats content, distance to the centromere/telomere (Fryxell and Moon 2005; Elango et al. 2008; Tyekucheva et al. 2008; Chen et al. 2010)). Further, to avoid variance generated by differing levels of selective constraint, analyses were focused on putatively neutral regions.

### Nucleotide Diversity between Autosomes in Chimpanzees

The autosomes in chimpanzees exhibit similar levels of nucleotide diversity (fig. 2)—with all autosomes being in the range of one standard deviation of the genome-wide average \( (\pi = 6.9 \times 10^{-3}) \). The contribution of different genomic factors influencing levels of variation was assessed using multiple linear regression as well as multiple logarithmic regression (see Materials and Methods section). All parameters were scaled to have variance 1 and mean 0 in order to enable an easy comparison, and the autosomes were analyzed in 1 Mb segments with at least 80% accessibility (i.e., at least 80% of sites in the 1 MB segment passed the filter criteria), utilizing the genetic map from Auton et al. (2012). Across all sites, recombination rate was the primary predictor of nucleotide variation (fig. 3), with a strongly positive correlation (table 1), indicative of an important genome-wide role for linked selection. In addition, regional CpG-island and GC-content are strongly negatively correlated with variation (table 1), consistent with their associated effects on mutation rate. CpG dinucleotides, generally exhibiting high mutation rates (i.e., transition mutation rates...
are elevated by ~30-fold in great apes (Hwang and Green 2004; Siepel and Haussler 2004; Keightley et al. 2011) owing to spontaneous methylation-dependent deamination (e.g., Cooper and Youssoufian 1988; Duret 2009), are more stable within CpG-islands than in the rest of the genome—an observation that has often been attributed to the fact that CpG dinucleotides are usually unmethylated within CpG-islands (Polak and Arndt 2008; Cohen et al. 2011). In addition, CpG-islands can play a role in gene regulation, leading to more stable CpG-sites when they are under selection (Hodgkinson and Eyre-Walker 2011). Mutation rates of methylated CpG dinucleotides are also decreased in regions of high GC-content compared to other regions in the genome, possibly due to a lower melting of the DNA duplex as methylcytosine deamination preferentially occurs on single-stranded DNA (Fryxell and Moon 2005; Elango et al. 2008). Furthermore, exon content remains a fourth but significant predictor of variation in chimpanzees, presenting a stronger correlation than has been observed in humans.

**Interpreting Similarities and Differences between Humans and Chimpanzees**

Mutation-associated recombination has been suggested as a potential explanation for the observed correlation between recombination and diversity (see Kimura and Crow 1964). Indeed, there is some evidence of this in yeast (e.g., Strathern et al. 1995). However, sequence divergence in other organisms, ranging from Drosophila to humans, does not provide support for this explanation—an observation additionally supported by the data presented here. This result is particularly strong given the recent divergence of human and chimpanzee, and the broad-scale similarity of their

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Table 1

| Recombination rate | GC content | Distance to centromere | Distance to telomere | Size | Exon content | CpG-Island content | Simple repeat content |
|-------------------|------------|------------------------|---------------------|------|-------------|-------------------|----------------------|
| \( \pi \)          | \(-0.207\) | \(-0.032\)          | \(-0.239\)          | \(-0.085\) | \(-0.256\) | \(-0.222\) | \(-0.049\)         |
| Recombination rate | 0.059      | 0.055                 | \(-0.509\)          | \(-0.172\) | \(-0.142\) | \(-0.092\) | 0.096               |
| GC content        | 0.046      | 0.046                | \(-0.389\)          | \(-0.244\) | 0.376       | 0.455             | 0.230               |
| Distance to centromere | \(-0.312\) | 0.251                | \(-0.021\)          | 0.002   | \(-0.056\) |                   |                     |
| Distance to telomere | \(0.332\) | \(-0.009\)          | \(-0.087\)          | \(-0.147\) | \(-0.271\) |                   |                     |
| Size              | \(-0.066\) | \(-0.027\)          | \(-0.271\)          | \(-0.271\) | \(-0.271\) |                   |                     |
| Exon content      | \(0.290\)  | \(0.084\)            | \(-0.092\)          | \(-0.092\) | \(-0.092\) |                   |                     |
| CpG-Island content | \(0.037\)  | \(0.037\)            | \(-0.092\)          | \(-0.092\) | \(-0.092\) |                   |                     |

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**Fig. 3** — Relative importance of each significant regressor in the model for the nucleotide diversity estimate \( \pi \) in (a) Western chimpanzees \( R^2 = 45.85\% \) and (b) humans \( R^2 = 55.69\% \). The LMG metric (Lindemann et al. 1980) was used to calculate the relative importance of each predictor by partitioning \( R^2 \) by averaging over orders (computed using Gnu R’s “rela.impo” package). Metrics were normalized to sum to 100%. Significance levels: *** 0; ** 0.001; * 0.01.
recombinational landscapes (McVicker et al. 2009; Lohmueller et al. 2011; Auton et al. 2012).

Thus, the predictive relationship between recombination and diversity in chimpanzees may be interpreted as evidence for the effects of linked selection. Importantly, RHI will have greater effects when positive selection is strong, with simulations suggesting an expected logarithmic relationship between recombination and diversity (Hellmann et al. 2005). Conversely, BGS will have the greatest effect when purifying selection is relatively weak (see Charlesworth 2012), with simulations suggesting an expected linear relationship between recombination and diversity (Hellmann et al. 2005). Interestingly, a logarithmic model is a better fit to both human and chimpanzee data, suggesting a potentially important role for positive selection in shaping genomic variation in both species (AIClog(chimp) = 5,130; AIClinear(chimp) = 5,369; AIClog(human) = 4,683; AIClinear(human) = 4,958).

Naturally, evidence is also observed for the important role of purifying selection. Note that in the weak-selection regime, a simple re-scaling of effective population size is no longer sufficient to account for BGS effects, and the site frequency spectrum may become strongly left-skewed (Ewing and Jensen 2016). Differences in this regard between these two species may be expected given their differing recent demographic histories, with human populations undergoing rapid growth (Coventry et al. 2010; Tennesen et al. 2012). More specifically, under this human demographic model, the increasing effective population size is expected to better prevent the fixation of strongly deleterious mutations, while the extreme population growth is expected to result in a larger proportion of segregating weakly deleterious mutations (Lohmueller et al. 2008; Keinan and Clark 2012; Gazave et al. 2013; Ewing and Jensen 2014; Lohmueller et al. 2014).

In order to investigate this expected difference with the genome-wide data used here, we compared the ratio of non-synonymous to synonymous polymorphic and divergent sites between chimpanzees and humans, utilizing a re-constructed ancestor as an outgroup. The mean genome-wide pWVps in humans is 1.27 and 1.26 in chimpanzees (Wilcoxon rank sum test: P value < 0.03). Conversely, mean genome-wide dN/dS in humans is 1.04 and 1.23 in chimpanzees (Wilcoxon rank sum test: P value < 2.5 × 10^{-4}). Thus, consistent with the above expectation owing to differences in recent demographic histories, a greater ratio of non-synonymous to synonymous variants is observed to be segregating in the human population, with a lesser ratio of non-synonymous to synonymous fixations on the human branch. Relatedly, there is a stronger relationship between exon content and diversity in chimpanzees than in humans. Consistent with this observation, Bataillon et al. (2015) recently argued for high rates of purifying selection in the coding regions of chimpanzee utilizing the inference framework of Eyre-Walker and Keightley (2009).

Thus, these different predictions developed in the theory literature are increasingly valuable for interpreting the influx of genomic polymorphism data from closely related species, and clearly suggest that the powerful and widely invoked correlation between recombination rate and diversity, combined with other genomic information, will be of great value in inferring differing strengths and rates of selection between species.

Materials and Methods

Diversity

A set of 5,323,301 autosomal chimpanzee single nucleotide polymorphisms (SNPs) from medium-coverage (average 9.1X) Illumina GAII sequencing of 10 Western chimpanzees (nine females and one male) was obtained from the PanMap project (Auton et al. 2012). PanMap SNP calls were subject to several filter criteria in order to minimize genotype errors (see Auton et al. 2012, SOM pp. 4–6). As the applied filter metrics can lead to the exclusion of a substantial fraction of sites in the genome, mask files, defining which nucleotides were accessible to the variant discovery in the study, were necessary to enable population genetic analysis. Mask files were generated using GATK’s “UnifiedGenotyper” and “VariantFiltration” (Version 1.0.4705) (McKenna et al. 2010; DePristo et al. 2011) using the same filter criteria.

Each chromosome was divided into continuous windows of 1Mb size and the number of nucleotide differences per site between two randomly chosen sequences (π) (Nei and Li 1979) as well as Watterson’s estimate of θ (Watterson 1975) were estimated for all windows with at least 80% accessibility (i.e., at least 80% of sites in the 1Mb segment passed the filter criteria).

In order to enable a comparison with humans, autosomal genotype data, consisting of 7,906,281 SNPs from a sample of 10 Yoruban (YRI) individuals (NA18522, NA19116, NA18912, NA19093, NA18516, NA18501, NA18870, NA18498, NA18510, and NA18499) was obtained from the 1000 Genomes Low Coverage Pilot Project SNP release (1000 Genomes Project Consortium 2010). These individuals exhibit a similar sequence coverage (~7.08 X on an average) as well as an equivalent concentration of recombination rate in the fine-scale genetic map than the chimpanzee sample (Auton et al. 2012).

All chimpanzee and human SNPs were annotated using ANNOVAR (Wang et al. 2010) with the information of the chimpanzee genome build panTro2 and the human genome build hg18, respectively (extracted from the “refGene” data set of the UCSC Genome table browser (Karolchik et al. 2004)), enabling the identification of synonymous (S) and non-synonymous (N) coding variants. Thereby, the panTro2 annotation contained 2,699 transcripts for 2,574 unique genes (including 653 without coding sequence annotation),
whereas the hg18 annotation contained 52,204 transcripts for 26,452 genes (including 11,833 without coding sequence annotation). Annotations were used to calculate the ratio of non-synonymous to synonymous polymorphisms (pKNS) in both species. Mean genome-wide pKNS were calculated in continuous windows of 100 kb size, excluding information from chimpanzee chromosome 2a and 2b and the orthologous regions in human chromosome 2 due to the different histories of the chromosomes (i.e., human chromosome 2 originated from a telomeric fusion event in the human ancestral lineage (Judo et al. 1991)) as well as from chromosome 13 for which there were an insufficient number of annotated genes with polymorphisms available in the chimpanzee data set. The Wilcoxon rank sum test statistic was calculated using Gnu R's inbuilt functions.

Annotation of Genome Features

The relationship of regional nucleotide diversity levels in Western chimpanzees as well as in humans with specific genome features (namely recombination rate, GC-content, exon content, simple repeat content, CpG-island content, distance to the centromere and telomeres, as well as chromosome size) was studied to reveal whether there were any correlations between nucleotide diversity at large scales and these sequence features in either of the two species.

The length of each chromosome was determined and GC-content was measured using information obtained from the chimpanzee genome build panTro2.1 and the human genome build hg18, respectively, as downloaded from the UCSC genome browser. The distance from the middle of a given window to the centromere (typically consisting of large arrays of repetitive DNA) as well as to the closest telomere (often containing high GC content as well as high rates of recombination) was calculated to obtain information about the influence of large-scale chromosomal structure on nucleotide diversity rates. The distance was set to 0 if the entire window fell within a centromere/telomere region. Locations of centromeres and telomeres were obtained from the UCSC “Gap” database table (whereby chromosomal start and end coordinates were used to fill in missing telomere data). Genes were annotated using ANNOVAR (Wang et al. 2010) with the information of the chimpanzee genome build panTro2 and the human genome build hg18, respectively.

Regression Analyses

The contribution of the different genomic factors influencing the observed 1 Mb-scale variation in diversity levels across the genome of Western chimpanzees and humans was assessed using multiple linear regression as well as multiple logarithmic regression. Thereby, analyses were limited to intergenic 1 Mb windows with at least 80% accessibility. Estimates for diversity levels were log-transformed to be roughly normally distributed. All parameters were scaled to have variance 1 and mean 0 in order to enable an easier comparison of the different parameters (after the transformation, the slopes directly measure the strength of the relationship between the explanatory and the response variable). All possible models were analyzed and standard regression diagnostics were used to evaluate the validity of the model as well as to identify and remove outliers. The best model was chosen according to Akaike’s Information Criterion (AIC) (as implemented in Gnu R). The LMG metric (Lindemann et al. 1980) was used to calculate the relative importance of each predictor by partitioning $R^2$ by averaging over orders (computed using Gnu R’s “rela.impo” package).

Divergence

In order to identify fixed differences between the 10 Western chimpanzee individuals and humans, PanMap reads were additionally aligned against the human reference genome build hg18, using the same quality criteria than for the alignment against the chimpanzee reference genome (Auton et al. 2012). The ancestral allele for each site was determined from chimpanzee chromosome 2 due to the different histories of the chromosomes (i.e., human chromosome 2 originated from a telomeric fusion event in the human ancestral lineage (Judo et al. 1991)).

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