1	Genetic differentiation and intrinsic genomic features
2	explain variation in recombination hotspots among
3	cocoa tree populations
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10 Abstract

Our study investigates the possible drivers of recombination hotspots in *Theobroma cacao* 11 using ten recently diverged populations. This constitutes the first time that recombination 12 rates from more than two populations of the same species have been compared, providing a 13 novel view of recombination at the population-divergence time-scale. For each population, 14 a fine-scale recombination map was generated using a method based on linkage disequi-15 librium (LD). They revealed higher recombination rates in a domesticated population and 16 a population that has undergone a recent bottleneck. We address whether the pattern of 17 recombination rate variation along the chromosome is sensitive to the uncertainty in the per-18 site estimates. We find that uncertainty, as assessed from the Markov chain Monte Carlo 19 iterations is orders of magnitude smaller than the scale of variation of the recombination 20 rates genome-wide. We inferred hotspots of recombination for each population and find that 21 the genomic locations of these hotspots correlate with genetic divergence between popula-22 tions (F_{ST}) . The large majority of inferred hotspots are not shared between populations 23 (55.5%). We developed novel randomization approaches for the generation of appropriate 24 null models to understand the association between hotspots of recombination and both DNA 25 sequence motifs and genomic features. Hotspot regions contained fewer known retroelement 26 sequences than expected, and were overrepresented near transcription start and termination 27 sites. Indicating that recombination hotspots are evolving in a way that is consistent with 28 genetic divergence, but are also preferentially driven to regions of the genome that contain 29 specific features. 30

31 Introduction

Genetic variation is fundamental for evolutionary forces like selection and genetic drift to 32 act. Selection and drift also contribute to a loss of variation, which means that they must act 33 in tandem with forces that maintain variation along the genome in order for populations to 34 continue evolving over prolonged periods of time. Selection and drift also contribute to a loss 35 of variation, which means that they must act in tandem with forces that maintain variation 36 along the genome in order for populations to continue evolving over prolonged periods of 37 time. Recombination's rearranging of genetic material onto different backgrounds generates 38 a larger set of haplotype combinations on which selection can act, reducing the magnitude 39 of Hill-Robertson interference (Felsenstein, 1974). Different regimes of recombination can 40 strongly influence how efficient selection is at purging deleterious mutations and increasing 41 the frequency of beneficial mutations in the population (Felsenstein, 1974). 42

Genome-wide fine-scale recombination maps can help elucidate the distribution of recom-43 bination events along the genome (Mvers et al., 2005; Auton et al., 2012; Brunschwig et al., 44 2012; Paape et al., 2012; Choi et al., 2013; Hellsten et al., 2013; Singhal et al., 2015; Stevison 45 et al., 2016). These maps are constructed using methods that use the coalescent to leverage 46 current patterns of LD in order to estimate historical rates of recombination between sites 47 along the genome (Auton and McVean, 2007). Studies in a wide range of species have shown 48 that recombination rates are not uniform along the genome and general patterns of variation 49 have been described (Begun and Aquadro, 1992; Akhunov et al., 2003; Wu et al., 2003; An-50 derson et al., 2004; McVean et al., 2004; Mézard, 2006; Kim et al., 2007; Gore et al., 2009; 51 Schnable et al., 2009; Branca et al., 2011; Paape et al., 2012). One of these patterns is the 52 reduced recombination rate in centromeric regions of the chromosomes and the progressive 53 increase of recombination rates as the physical distance to telomeres decreases (Begun and 54 Aquadro, 1992; Akhunov et al., 2003; Wu et al., 2003; Anderson et al., 2004; Gore et al., 2009; 55

Schnable et al., 2009). Another, perhaps more interesting pattern that has been observed is
regions with unusually high rates of recombination spread throughout chromosomes: recombination hotspots (McVean et al., 2004; Brunschwig et al., 2012; Paape et al., 2012; Hellsten
et al., 2013; Stevison et al., 2016; Shanfelter et al., 2018). The importance of recombination
hotspots lies in their ability to shuffle genetic variation at higher rates than the rest of the
genome, profoundly impacting the dynamics of selection for or against specific mutations
(Felsenstein, 1974).

A variety of genomic features have been identified as being associated with regions of 63 high recombination. In Arabidopsis thaliana, Taeniopygia guttata, Poephila acuticauda, and 64 humans, hotspots have been linked to transcriptional start sites (TSSs) and transcriptional 65 termination sites (TTSs) (Myers et al., 2005; Choi et al., 2013; Singhal et al., 2015). In 66 Mimulus auttatus hotspots were found to be associated with CpG islands (short segments of 67 cytosine and guanine rich DNA, associated with promoter regions) (Hellsten et al., 2013). 68 These patterns point to recombination occurring frequently near, but not within coding 69 regions. The formation of chiasmata is important for the proper disjunction of chromosomes 70 during meiosis (Martinez-perez et al., 2008), but repeated double-strand breaks can lead to 71 an increased mutation rate (Rodgers and Mcvey, 2015). In coding regions in particular this 72 excess mutation rate can have a high evolutionary cost, due to the likelihood of deleterious 73 mutations arising being higher than that of beneficial ones (Haldane, 1937; Crow, 1970; 74 Wloch et al., 2001; Sanjuán et al., 2004; Eyre-Walker and Keightley, 2007). Recombination 75 hotspots have also been found to be correlated with particular DNA sequence motifs. In 76 some mammals, including *Mus musculus* (Brunschwig et al., 2012) and apes (Auton et al., 77 2012; Stevison et al., 2016) binding sites for PRDM9, a histone trimethylase with a DNA 78 zinc-finger binding domain, have been found to correlate with recombination hotspots. In 79 Arabidopsis, proteins have been identified that limit overall recombination rate, leading to 80 an increased recombination rate genome-wide in mutants (Fernandes et al., 2018). However, 81

these proteins have not been shown to direct recombination to particular regions, and are therefore not expected to affect the location of recombination hotspots.

Recent work on recombination in apes (Stevison et al., 2016) found little correlation 84 of recombination rates in orthologous hotspot regions when looking between species, but a 85 strong correlation when comparing between two populations of the same species. This sug-86 gests that recombination hotspots are potentially changing in ways that match demographic 87 patterns, differentiating at a similar rate as genomic sequences. The identification of ten 88 recently diverged populations of the cocoa tree, *Theobroma cacao* (Motamayor et al., 2008; 89 Cornejo et al., 2018) can be leveraged to study short-term drivers of recombination hotspots. 90 These ten populations originate from different regions of South and Central America, and 91 include one fully domesticated species (Criollo), used in the production of fine chocolate, and 92 nine wilder, more resilient species which generate higher cocoa yield than the Criollo variety 93 (fig. 1) (Motamayor et al., 2008; Henderson et al., 2007; Cornejo et al., 2018). These ten 94 populations have been shown to be highly differentiated and the history of diversification has 95 been recently confirmed using genomic data (Cornejo et al., 2018). Comparing the locations 96 of hotspots between these ten populations of T. cacao can contribute to the understanding 97 of hotstpot turnover in short periods of time among highly differentiated populations. These 98 comparisons also contribute to our understanding of how demographics impact the turnover 99 of recombination hotspot locations. 100

Fine-scale, LD-based recombination maps have been constructed for a number of plant models (Paape et al., 2012; Choi et al., 2013; Hellsten et al., 2013), and in all of them a variety of correlates of recombination rate have been identified. Unlike these model plants with short generation times, *T. cacao* is a perennial woody plant with a five-year generation time (Henderson et al., 2007). Studying recombination in *T. cacao* presents particular difficulties due to its long generation time, making it difficult to directly measure rates of recombination. However, linkage-based methods allow for the estimation of historic recombination rates (Auton and McVean, 2007), facilitating the study of historical recombination in *T. cacao*. Theoretical studies have shown that population structure can generate artificially inflated measures of LD (Ohta, 1982; Li and Nei, 1974), which would be detrimental to our estimates of recombination. For this reason recombination maps were constructed independently for each population. In contrast to previous studies, which have focused primarily on recombination rates, this study attempts to describe the relationship between recombination hotspots and a variety of factors.

A series of questions were addressed in this work: (i) How are recombination rates dis-115 tributed within 10 highly differentiated populations of T. cacao, and how do they compare 116 to each other? (ii) How are hotspots distributed along the genome of each of the ten popula-117 tions of T. cacao, and can these distributions be explained by patterns of population genetic 118 differentiation? (iii) Are there identifiable DNA sequence motifs that are associated with the 119 location of recombination hotspots along the T. cacao genome? (iv) Are there genomic fea-120 tures (e.g. TSSs, TTSs, exons, introns) consistently associated with recombination hotspot 121 locations across T. cacao populations? 122

In order to address these questions, we used an LD-based method to estimate recombina-123 tion rates, since T. cacao's generation time and sheer size make performing a large number of 124 backcrosses unfeasible. We used the estimated recombination rates in a maximum likelihood 125 statistical framework to infer the location of recombination hotspots. Then we compared 126 the location of hotspots across populations and found evidence that, while hotspots gener-127 ally follow patterns of genetic differentiation, their turnover rate is faster than the rate of 128 divergence. We used resampling schemes to generate null assumptions for the content of 129 known DNA sequence motifs in ubiquitous recombination hotspots, as well as the overlap 130 of population recombination hotspots with genomic traits. The resampling schemes used to 131 identify these associations are novel in the context of this work and were designed to take 132 into account the size and distribution of elements in the genome. Our findings suggest that 133

recombination hotspot locations generally follow patterns of diversification between populations, while also having a strong tendency to occur close to TSSs and TTSs. Moreover, we
find a strong negative association between the occurrence of recombination hotspots and the
presence of retroelements.

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139 Results

140 Comparing recombination rates between populations

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Populations show a mean recombination rate r/kb between 2.1×10^{-5} and 5.25×10^{-3} 142 (table 1), with a long-right-tailed distribution (fig. 3). The extreme recombination rate 143 values affect the mean, driving it to values consistently higher than the median. The pattern 144 of recombination rates along the genome varied between populations, as can be seen in the 145 comparison of the Nanay and Purus third chromosome (fig. 4). The median 95% probability 146 interval for recombination rate across the genome for each population was found to be several 147 orders of magnitude larger than the uncertainty per site, estimated as the median 95%148 Credibility Interval of the trace for each position in the genome for that population (table 149 6).150

Overall, the recombination rate for most of the populations was higher than estimated mutation rates for multicellular eukaryotes of 10^{-6} changes per kb per generation (Lynch, 2010; Exposito-Alonso et al., 2018) (table 1). Two populations, Guianna and Criollo, were notable exceptions, having higher average recombination rates than the other populations by one and two orders of magnitude respectively. Guianna and Criollo also had a lower effective population size (N_e) (Cornejo et al., 2018) by one and two orders of magnitude respectively. However, there was no significant linear trend between mean N_e and r/kb

(p=0.09045), indicating that, for a high enough N_e , the ability to detect recombination events is not dictated by the effective population size. When Criollo and Guianna were excluded, the relationship was also not present (p = 0.6544). When all populations were included, the inbreeding coefficient (F) showed no significant linear association with mean r/kb (p = 0.3361). The average recombination rate per population was transformed from r/kb to cM/Mb (table 1). The average cM/Mb was 4.6×10^{-04} .

In order to compare the average recombination rates of the different populations, a Kruskal-Wallis test was performed for every pair of populations. The only pair of populations that did not show a significant difference in mean recombination rate was the pair of Nacional and Nanay (p = 0.3). All other pairwise comparisons were highly significant ($p < 2 \times 10^{16}$).

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170 Comparing recombination hotspot locations between populations

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The majority (55.5%) of hotspots identified were not shared between populations. The 25 most numerous sets of hotspots are represented in fig. 6. The nine largest of these are sets of hotspots unique to a single population. The hotspots unique to the remaining population (Criollo) formed the eleventh largest set.

The recombination rate in hotspot regions for nine of the populations was on average between 22 and 237% higher than the average recombination rate of the genome. The exception was Guianna, which only showed an approximately 1% increase in average recombination rate in hotspots regions when compared to that of the non-hotspot regions.

Despite the majority of hotspots not being shared between populations, pairwise Fisher's exact tests between populations indicated significantly more overlap than expected (if hotspots were randomly distributed along the genome) between hotspots for most pairs of populations (table 2). There were three comparisons that did not show significantly more overlap

than expected: Amelonado-Nacional, Amelonado-Purus, and Criollo-Nacional. A Mantel 184 test comparing distances between populations based on shared hotspots and F_{ST} values be-185 tween populations resulted in a significant correlation between them (r = 0.66, p = 0.002). 186 The correlation between eigenvectors from a correlation matrix and those of the genetic 187 covariance matrix were also explored. When all populations were included, we found that 188 the first eigenvector from the genetic covariance matrix was not significantly correlated with 189 the first eigenvector from the hotspot correlation matrix (p = 0.7055), but the second ge-190 netic eigenvector was (p = 0.009007, r = 0.7711638). However, the first eigenvector of the 191 genetic covariance matrix captured the difference between the Criollo population (the only 192 domesticated variety) and the rest of the populations. The second eigenvector explains most 193 of the natural differentiation across populations (Cornejo et al., 2018). For that reason, we 194 decided to exclude Criollo and repeat the analysis. We found that the first eigenvector from 195 the correlation matrix constructed from shared hotspot information was not significantly 196 correlated with either of the first two eigenvectors of the genetic covariance matrix when 197 Criollo was excluded (eigenvector 1: p = 0.1314, eigenvector 2: p = 0.3376). 198

To study the effects of demographic history more closely, shared hotspots were converted 199 to dimensions of a multiple correspondence analysis and modeled along a previously con-200 structed drift tree (Cornejo et al., 2018). The model assuming an OU process (stabilizing 201 selection) is consistent with a higher trait maintenance than the model assuming Brownian 202 motion (drift). Modeling the dimension as a Brownian motion was a better fit (AIC=79.4) 203 than modeling it as an Ornstein-Uhlenbeck (OU) process (AIC=81.4), which is consistent 204 with the small number of hotspots shared between populations. 205

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Identifying DNA sequence motifs associated with the locations of recombination hotspots 207 208

RepeatMasker was used to analyze the set of recombination hotspots that were present 209

in at least eight T. cacao populations (17 total hotspots), as well as the consensus set 210 of recombination hotstpots, and the reference genome. In order to determine whether a 211 particular set of DNA sequence repeats was overrepresented in the regions of ubiquitous 212 recombination hotspots, the percentage of DNA sequence that was identified as potentially 213 being from retroelements or DNA transposon was compared to an empirical distribution. 214 The percentage of observations from the distribution which were greater than the observed 215 are reported in table 3. While retroelements were found to be underrepresented in the ubiq-216 uitous hotspots, DNA transposons were marginally overrepresented. 217

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Identifying genomic features associated with the location of recombination hotspots

An overrepresentation of recombination hotspots was found in all ten of the populations 221 at transcriptional start sites (TSSs) and transcriptional termination sites (TTSs)(table 4). 222 The level of overrepresentation of hotstpots in particular regions was compared to a null ex-223 pectation based on simulations of hotspots of the same size as the ones detected distributed 224 randomly along chromosomes. For all populations, all 1000 simulations showed a lower pro-225 portion of overlap with TSSs and TTSs than the observed overlap. In the case of exons and 226 introns, seven populations (Contamana, Criollo, Iquitos, Maranon, Nacional, Nanay, Purus) 227 had an observed value that was lower than all, or almost all (Purus for exons), simulations. 228 Three of the remaining four populations (Amelonado, Curaray, and Nanay) had no clear 229 trend in either direction (table 2). The final population (Guianna) showed an overrepresen-230 tation of hotspots in both exons and introns. 231

233 Discussion

Understanding how recombination rates vary between recently diverged populations is an 234 important step toward disentangling the role of recombination in genetic differentiation. 235 This set of T. cacao populations presents a unique opportunity to observe recombination 236 in long-established, non-domesticated populations, as well as a recently established non-237 domesticated population (Guianna) and a domesticated population (Criollo) (Cornejo et al., 238 2018; Bartley, 2005). This model has allowed us to explore divergence patterns of recombina-239 tion hotspot at a scale below that of species divergence. Our results point to a conservation 240 of hotspots between populations that generally mirrors the patterns of genetic differentiation 241 between populations. However, we find that the correlation between genetic differentiation 242 and shared hotspots does not fully explain the observed hotspots, in part due to the ma-243 jority of hotspots being found in a single population. We find that TSSs and TTSs are 244 strongly associated with recombination hotspots in all populations, which is consistent with 245 previous findings in plants (Paape et al., 2012; Choi et al., 2013; Hellsten et al., 2013). This 246 factor seems to play an important role in determining the location of novel hotspots. Fi-247 nally, hotspots that are shared by at least eight populations appear to be associated with 248 DNA transposons, pointing to a potential mechanism for the maintenance of recombination 249 hotspots at the population-divergence time-scale. 250

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252 Comparing recombination rates between populations

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We found that the eight long-established, non-domesticated *T. cacao* populations show a recombination rate (r/kb) lower than multicellular eukaryotic mutation rates (table 1), while the other two populations (Criollo and Guianna) show unusually high average recombination rates in comparison. For all populations, the mean recombination rate was found to be lower

than the median. This is consistent with high rate outlier values; an expected result in the 258 presence of recombination hotspots. Using the effective population size for Medicago trun-259 catula from Siol et al. (2007) and the estimate of rho from Paape et al. (2012), we calculated 260 r/kb (= 4 × 10⁻³) and found that it was comparable with the rate found for the Criollo 261 population (table 1). We also calculated the median recombination rate in cM/Mb for each 262 chromosome using the Kosambi mapping function (Kosambi, 1943) over non-overlapping, 263 100 SNP windows. The average cM/Mb for all populations was 4.6×10^{-04} , which is lower 264 than has been measured for any Malvale (Kundu et al., 2015), but not as low as the lowest 265 measured for conifers (Chen et al., 2010; Stapley et al., 2017). Average recombination rates 266 in cM/Mb varied between populations from Amelonado (4.04×10^{-06}) to Criollo ($3.91 \times ^{-03}$). 267 Previous work (Cornejo et al., 2018) has shown that Criollo is the only population showing 268 a strong signature of domestication, as revealed by much higher drift than that observed 269 for other populations. Domestication has been observed to increase recombination rates, 270 particularly in plants (Ross-Ibarra, 2004), and is a possible explanation for the higher re-271 combination rate observed for the Criollo population. The high recombination rate observed 272 in Guianna can be explained in a similar way; while Guianna does not show a strong signa-273 ture of domestication, it is the most recently established population (Bartley, 2005), and it 274 has also undergone a recent bottleneck (Cornejo et al., 2018). It's possible that the Guianna 275 population is undergoing the initial stages of domestication and its increased recombination 276 is an early indicator of this. It is possible that the high recombination rates estimated for 277 Criollo and Guianna can be explained by biases in estimation caused by errors associated 278 to small samples or low genetic variation; yet, the recombination rates for Amelonado (an-279 other population with low variation) or Purus (a population with small sample size) did not 280 present this problem. Analyses exploring mutations of putative recombination suppression 281 genes (Fernandes et al., 2018) could help disentangle the nature of this extreme variation in 282 recombination rate in the Criollo and Guianna populations. 283

Despite recombination rates for eight of the ten populations being of the same order 284 of magnitude, pairwise comparisons of average rates indicated that most populations have 285 a significantly different rate of recombination from the others. The only exception were 286 Nacional and Nanay whose average rates were not significantly different from each other. 287 These two populations, however, are not more closely related to each other than they are 288 to other populations, based on sequence divergence (Cornejo et al., 2018). We interpret 289 this result as suggestive that their similarity is not due to genetic similarity, but some other 290 factors, e.g. epigenetics. 291

The likelihood of detecting Hotspots of recombination in the genome will likely be af-292 fected by the amount uncertainty in the estimates of recombination site-wise and region-wise. 293 Yet, we have been unable to identify any study where the magnitude of the uncertainty in 294 the estimates of recombination are assessed to address this issue. We have performed careful 295 comparisons and assessed the magnitude of the uncertainty in the estimation of recombi-296 nation rates to show that this uncertainty is several orders of magnitude smaller than the 297 variation in recombination rates across the genome (table 6). If the length of the chains had 298 been maintained at 40MM generations, this would have increased the uncertainty in local 299 estimates making our assessment of rate variation along the genome more challenging. 300

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302 Comparing recombination hotspot locations between populations

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Similarly to recombination rates, the location of recombination hotspots can be very informative to questions of divergence between populations. Understanding the pattern and rate of change of recombination hotspots at the population level can elucidate their role in shaping genome architecture, impacting how effectively selection operates (Felsenstein, 1974). We found that a large proportion (55.5%) of hotspots are unique to a single population, which can be seen as an indicator that the turnover rate for hotspots is faster than the time it took

the 10 populations to diverge. This variability of hotspot location between populations 310 points to demographic history not being the main driver of recombination hotspot location. 311 However, the hotspots tend to appear in similar regions, as demonstrated by the Fisher's 312 exact tests (table 2). This dichotomy can be explained by considering that the proportion 313 of the genome occupied by recombination hotspots is very low, so even a small proportion of 314 hotspots from two different populations being in the same region is enough for the Fisher's 315 exact test to recognize them as significantly similar. This small but significant similarity 316 can occur by recombination being limited in its possible positioning along the genome, but 317 not to the point of forcing hotspots to occur consistently in the same locations, and thus 318 maintaining some level of stochasticity. 319

Given the significant proportion of overlapping hotspots between populations, it was still 320 important to explore whether the similarities can be explained by shared genetic history. If 321 demographic history explained the evolution of hotspot location, it would be expected that 322 more closely related populations would have a higher percent of overlapped hotspots. A 323 significant relationship was found between population differentiation (F_{ST}) and the differ-324 entiation between populations based on shared hotspots (Mantel test, r = 0.66, p = 0.002). 325 The comparison between the hotspot correlation matrix and the genetic covariance matrix 326 supports what was found when comparing the hotspot correlation matrix to the F_{ST} matrix. 327 One caveat is that the first genetic eigenvector, which separated Criollo from the other pop-328 ulations, was not correlated with the first hotspot correlation eigenvector, indicating that 329 Criollo's domestication generated a genetic pattern that deviates from the pattern of shared 330 hotspots. This indicates that, to some extent, the genetic differentiation and the location 331 of hotspots are mirroring each other, which could be due to recombination hotspots being a 332 product of the shared history between the populations. However, since recombination rates 333 were estimated using a coalescent-based method, we expect historical relationships to be 334 represented in our findings. We also transformed the information of hotspot overlap to allow 335

for the modeling of hotspots as traits along a population tree. Our results, showing that 336 a Brownian motion model (AIC=79.4) better fits the data than a model with stabilizing 337 selection Ornstein-Uhlenbeck model (AIC=81.4), suggests in first principle that drift alone 338 could explain the evolution of the location of recombination hotspots. However, the absolute 339 number of hotspots that are shared among populations indicates that demographic history 340 alone is insufficient to explain the evolution of recombination hotspots in this species. Pre-341 vious studies looking at apes and finches have explored recombination hotspots in multiple 342 species and as many as two populations of the same species (Singhal et al., 2015; Stevison 343 et al., 2016; Shanfelter et al., 2018), but this study is the first to compare more than two 344 populations of the same species at once. The increased number of populations allows us 345 to analyze the relationship between population genetic processes and recombination. Our 346 results suggest that the pattern of gains and losses of recombination hotspots can be very 347 dynamic and the landscape of recombination changes rapidly during the process of diver-348 sification within a species. This dynamism can have a tremendous impact on the adaptive 349 dynamics of a species, and it should be taken into account, considering that theoretical stud-350 ies tend to assume that recombination rates are constant during the evolution of populations 351 (Hudson and Kaplan, 1988; Donnelly and Kurtz, 1999). 352

One conclusion that follows from these results is that, while shared recombination hotspots 353 can to some extent be explained by patterns of genetic differentiation, some of the sharing 354 can simply be due to a tendency for hotspots to arise in similar locations. It has been ob-355 served in other organisms that hotspots of recombination are frequently associated to specific 356 genomic features (including TSSs and TTSs) (Auton et al., 2013; Choi et al., 2013; Hellsten 357 et al., 2013; Myers et al., 2005; Singhal et al., 2015) or DNA sequence motifs (Auton et al., 358 2012; Brunschwig et al., 2012; Stevison et al., 2016). These factors can affect the landscape 359 of recombination, generating the patterns of shared hotspot locations between populations 360 that we are observing in T. cacao. 361

Identifying DNA sequence motifs associated with the locations of recombination hotspots

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The analysis of 17 hotspots shared between at least eight populations of T. cacao found 365 an underrepresentation of retroelements and a marginal overrepresentation of DNA trans-366 posons when compared to the entire genome. These results are not entirely surprising as 367 it has been already suggested that transposable elements (TEs) tend to be enriched in ar-368 eas of low recombination in *Drosophila* as a consequence of selection against TEs (Rizzon 369 et al., 2002). The marginal over-representation of DNA transposons in the most conserved 370 recombination hostspot is unexpected, given that all previous observations have shown a 371 reduced representation of mobile elements in areas with high recombination rate (Rizzon 372 et al., 2002). It's possible that DNA transposons are, at least in part, responsible for the 373 maintenance of recombination hotspots as populations diverge, from which we expect that 374 site-directed recombination is more frequent in these locations of the genome. However, the 375 low percentage of these sequences observed in the set of all hotspots (table 3) indicates that 376 these sequences only have a small effect on the maintenance of hotspots. It has been observed 377 in humans that short DNA motifs enriched for repeat sequences determine the location of 40 378 per cent of hotspots enriched for recurrent non-allelic homologous recombination (Mcvean, 379 2010). One potential explanation that for why natural selection does not eliminate hotspots 380 in these regions is the possibility that these regions don't produce a large enough mutational 381 load for natural selection to remove them from the population (Mcvean, 2010). 382

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³⁸⁴ Identifying genomic features associated with the location of recombination hotspots

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For all ten populations, an overrepresentation of hotspots was found in the areas immediately preceding and following transcribed regions of the chromosome. This matches the

findings of previous studies in Arabidopsis thaliana (Choi et al., 2013), Taenipygia guttata 388 and Poephila acuticauda (Singhal et al., 2015), and humans (Myers et al., 2005). The most 389 likely explanation is that recombination events within genes are selected against. The ra-390 tionale is that a recombinant chromosome that is split in the middle of a coding region will 391 have a higher risk of being inviable, and therefore not represented in the current set of chro-392 mosomes for its population. Recombination occurring in transcription start and stop sites. 393 on the other hand, does a much better job at breaking up haplotypes, while preserving the 394 functionality of coding regions. This rational is supported by previous findings of increased 395 recombination rates in these regions (Choi et al., 2013). It's also supported by results from 396 PRDM9 knock-out *Mus musculus*, which has shown a reversion to hotspots located near 397 TSSs (Brick Kevin et al., 2012). The enrichment of T. cacao hotspots in TSSs and TTSs 398 is thus a reasonable result given that zinc-finger binding motifs and potential modifiers like 399 PRDM9 have not been identified in the species. 400

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⁴⁰² Implications for the evolutionary history of T. cacao

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Overall, our results show a large consistent pattern where recombination rates in T. ca-404 *cao* are of the same magnitude as mutation rates, but show a high diversity in location 405 and number of hotspots of recombination that cannot be explained solely by the process of 406 diversification of the populations. In fact, the results are indicative that the turnover rate of 407 hotspots is faster than the process of divergence among populations. A potential hypothesis 408 that could explain the rapid turnover of hotspots of recombination and the relative differ-409 ences in recombination among populations is that epigenetic changes control the turnover of 410 recombination in plants. This hypothesis is not unreasonable given the recent observation 411 of epigenetic control of recombination in plants (Yelina et al., 2015). Further theoretical 412 and simulation work should be done in order to better understand the implications of the 413

rapidly changing recombination hotspots in adaptive dynamics. We also show that there is 414 an overall underrepresentation of hotspots in exons and introns for most populations, which 415 is consistent with purifying selection acting against changes that could result in disruptions 416 of gene function. On the other hand, we observed an overrepresentation of hotspots in TTSs 417 and TSSs for all ten populations. This could impact the maintenance and spread of benefi-418 cial traits in the population by shuffling allelic variants of genes without causing disruption 419 of their function. We hypothesize that the enrichment of hotspots of recombination in TTSs 420 and TSSs can have an important impact in the spread of beneficial mutations across different 421 genomic backgrounds; increasing the rate of adaptation to selective pressures (e.g. selection 422 for improved pathogen response). 423

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425 Materials and Methods

426 Comparing recombination rates between populations

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Sequence data were downloaded from the Cacao Genome Database and NCBI (Accession 428 PRJNA488484), including the reference sequence for each chromosome and the full genome 429 annotation (*Theobroma cacao* cv. Matina 1-6 v1.1) (Motamayor et al., 2013). Processing 430 was done using the pipeline from (Cornejo et al., 2018) available at the github repository 431 oeco28/Cacao_Genomics. Full genome data was used from a total of 73 individuals across 432 10 populations (Cornejo et al., 2018): Criollo (N = 4, #SNPs = 309,818), Curaray (N = 433 5, #SNPs = 1,106,871), Contamana (N = 9, #SNPs = 2,097,618), Amelonado (N = 11, 434 #SNPs = 373,789), Maranon (N = 14, #SNPs = 1,783,226), Guianna (N = 9, #SNPs = 435 770,729), Iquitos (N = 7, #SNPs = 1,575,711), Purus (N = 6, #SNPs = 1,184,181), Nanay 436 (N = 10, #SNPs = 830, 885), and Nacional (N = 4, #SNPs = 718, 099). VCFTools (Danecek 437

Petr et al., 2011) was used to remove all singletons and doubletons. Only bi-allelic single
nucleotide polymorphisms (SNPs) were retained and were exported in LDhat format.

In order to estimate recombination rates we used the *interval* routine of LDhat (Auton 440 and McVean, 2007), a program that implements coalescent resampling methods to estimate 441 historical recombination rates from SNP data. To reduce computation time, each chromo-442 some was split into windows, each containing 2000 SNPs. To counteract the overestimation 443 of recombination rate produced at the ends of the windows, an overlap of 500 SNPs was left 444 between consecutive windows. The final window for each chromosome did not always match 445 the general scheme, so the final 2000 SNPs were taken (making the overlap with the second 446 to last window variable, but never less than 500 SNPs) (fig. 2). Once these windows were 447 generated, LDhat was run over each window with 100 million iterations, sampling every 10 448 thousand iterations (10,000 total points sampled), with a block penalty of 5. Lookup tables 449 with a grid of 100 points, a population mutation rate parameter (θ) of 0.1 and a number 450 of sequences (n) of 50 were used for all populations. The first 50 million iterations were 451 discarded as burn-in. Once recombination rates were calculated, 250 positions were cut off 452 from both windows involved in each overlap, so that the estimates for the first half of the 453 overlap was taken from the end of the preceding window and the estimates for the second 454 half of the overlap were taken from the beginning of the following window. The final overlap 455 in each chromosome was split in order to take only 250 SNPs from the second to last window, 456 regardless of the remaining size of the last window. The remaining rate estimates were then 457 merged in order to obtain the recombination rates for the entire chromosome. This was done 458 for each chromosome of each population. 459

The estimation of recombination rates with LDhat is approximated using a sampling scheme with a Markov Chain Monte Carlo (MCMC) algorithm as implemented in the *interval* routine. The inference of recombination rates is the result of the integration of estimated parameter values across iterations with the routine *stats*. In the majority of recent studies

where LDhat or LDhelmet are used (Myers et al., 2005; Auton et al., 2012; Brunschwig et al., 464 2012; Paape et al., 2012; Auton et al., 2013; Choi et al., 2013; Singhal et al., 2015; Stevison 465 et al., 2016), whether there is convergence of the Markov chains has not been explicitly 466 investigated. One study that we're aware of has used simulations to asses whether their 467 small sample size affected their ability to obtain reliable estimates of recombination using 468 LDhelmet (Booker et al., 2017), but did not assess the uncertainty of the estimates from 469 the MCMC process itself. We argue that evaluation of convergence is important to assess 470 the confidence in the estimated reported values, especially if there is interest in analyzing 471 the differences in recombination rate along the genome. Visual inspection of pilot runs of 472 the analysis demonstrated that convergence was not achieved after running 40M iterations, 473 which is why the length of the chains was increased to 100M iterations. Additionally we 474 explored the uncertainty in the estimates of recombination site-wise by integrating over the 475 trace of the estimates for recombination rate to infer the 95% Credibility Interval. We then 476 estimated the 95% of recombination estimates range across all sites in the genome to have 477 an overall measure of uncertainty that we compared to the median 95% Credibility Interval 478 for the trace of each position. 479

In order to compare recombination rates, the effective population size (N_e) calculated for 480 each population (Cornejo et al., 2018) was used to convert rates in $N_e r/kb$ to r/kb. Differ-481 ences in the mean genome-wide recombination rate between populations were then tested 482 using the Kruskal-Wallis test, using the kruskal.test function from the stats package in R 483 (R Core Team, 2018). There were 45 comparisons, making the Bonferroni correction cutoff 484 value: $\alpha = 0.0011$. To transform per population recombination rates from r/kb to cM/Mb, 485 we divided each chromosome into windows of 100 SNPs and used the Kosambi mapping func-486 tion (Kosambi, 1943). The median for the windows of a chromosome was then calculated, 487 and the average of each population's chromosomes was taken as that population's average 488 recombination rate in cM/Mb. 489

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Comparing recombination hotspot locations between populations

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Recombination hotspots were estimated with LDhot (Auton and McVean, 2007), a likelihoodbased program that tests whether a single distribution model or a two distribution model better explains the observed recombination rates in 1 kb sliding windows (default), for each chromosome. Each chromosome was run in its entirety, with the number of simulations (nsims) set to 1000. The resulting potential hotspots were refined by an alpha of 0.001, and overlapping hotspots were merged.

To determine the set of consensus hotspots, the hotspots from all populations were merged. Two hotspots from different populations were considered to be shared if they both overlapped with the same hotspot in the consensus set. To summarize all shared hotspots, a Boolean matrix was constructed, in which a population having a hotspot that overlaps with a hotspot in the consensus list leads to an indication of presence of the consensus hotspot in that population. This matrix was used to determine hotspots shared by two or more populations.

⁵⁰⁶ A Fisher's exact test was run for each pair of populations in order to determine whether ⁵⁰⁷ hotspots for the pair of populations overlap significantly more than expected. The BED ⁵⁰⁸ files containing the location of the recombination hotspots for each pair of populations were ⁵⁰⁹ compared using Bedtools:fisher (Quinlan and Hall, 2010). The number of comparisons was ⁵¹⁰ 45, making the the Bonferroni correction cutoff value: $\alpha = 0.0011$.

In order to compare the relationships between populations based on shared hotspots we calculated Jaccard distances (distance function, philentropy package, R) (Drost, 2018) and compared them to a published F_{ST} matrix (Cornejo et al., 2018) using a Mantel test (mantel.rtest function, ade4 package, R) (Chessel et al., 2004; Dray and Dufour, 2007; Dray et al., 2007; Bougeard and Dray, 2018).

The Boolean matrix for shared hotspots was also used to explore the relationship be-516 tween hotspot similarities and genetic covariances from a previous study (Cornejo et al., 517 2018). Singletons were removed from the hotspot matrix, which was converted to a corre-518 lation matrix using the mixed.cor function from the psych package in R (Revelle, 2018). 519 The mixed.cor function was used due to its ability to calculate Pearson correlations from 520 dichotomous data. We then used the eigen function in R (R Core Team, 2018) to generate 521 eigenvectors for the hotspot correlation matrix and the genetic covariance matrix. Pearson 522 correlations between the first and second eigenvector of the genetic covariance matrix and the 523 hotspot correlation matrix were then calculated (cor.test function, stats package, R)(R 524 Core Team, 2018). This analysis was done once with all populations included, and once with 525 the Criollo population excluded before correlations were calculated. 526

In order to model the presence or absence of hotspots along a drift tree, a multiple correspondance analysis was used on the Boolean matrix of shared hotspots using the MCA function from the FactoMineR package in R Lê et al. (2008). Nine dimensions were retained and used as traits along a previously generated drift tree (Cornejo et al., 2018). Using the Rphylopars package in R (Goolsby et al., 2016), the dimensions were modeled as Brownian motion and as an Ornstein–Uhlenbeck process. The fit of the two models were compared using the AIC values for the best fitting models of each type.

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Identifying DNA sequence motifs associated with the locations of recombination hotspots 536

Motifs associated with hotspots were found using RepeatMasker (Smith et al., 2016). The entire genome, the set of consensus hotspots, and a set of ubiquitous hotspots (hotspots shared by at least eight of the populations) were examined with RepeatMasker, using normal speed and "theobroma cacao" in the species option. In order to determine whether ubiquitous hotspots were enriched for particular DNA sequences, a set of the same number and size

of sequences was randomly selected from the genome using Bedtools:shuffle (Quinlan and
Hall, 2010) and examined with RepeatMasker. This simulation was repeated one thousand
times and a null distribution against which observed values were compared was constructed
from the results.

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Identifying genomic features associated with the location of recombination hotspots

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Testing whether recombination hotspots were overrepresented near particular genomic 549 features was done by using a resampling scheme to establish null expectations and then 550 comparing the observed value to the empirical distribution. For each feature, locations were 551 retrieved and the number of observed hotspots that overlap with this feature were counted. 552 To determine whether this amount of overlapping hotspots was unusually high or low, a set of 553 hotspots that matched the number of hotspots and the size of each hotspot was simulated. 554 These simulated hotspots were placed randomly along the chromosome, using a uniform 555 distribution. The simulation was run 1000 times and the number of simulated hotspots that 556 overlap with the true genomic features was measured for each simulation. The simulations 557 generate an expected distribution of overlap with the genomic feature, and the true value 558 was then compared to the distribution. When simulated hotspots overlapped, the location 559 of one of them was sampled again. Features tested were: Transcriptional start sites (TSSs), 560 transcriptional termination sites (TTSs), exons, and introns. TSSs and TTSs are considered 561 to be the 500bp upstream and downstream of coding regions respectively. 562

The reason for the proposed novel resampling scheme is that, if the size and distribution of genomic features and hotspots were not taken into account, it would set unrealistic expectations for the overlap between features under a null model of no association. In this sense, the null model would be inappropriate and potentially inflate the false positive rate.

568 Data and code availability

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Rate and summary files from LDhat runs as well as hotspots for each population will be placed in a Dryad repository (url). Scripts for LDhat and LDhot runs as well as additional analysis is compiled in the following github repository *ejschwarzkopf/recombination-map*.

574 Tables

Population	Mean	Mean	Mean	Median	Lower Bound	Upper Bound	Mean
	$4 \mathrm{Ner/kb}$	Ne	r/kb	r/kb	(Mean r/kb)	(Mean r/kb)	$\mathrm{cM/Mb}$
Amelonado	1.58	15744	2.51e-05	2.40e-09	2.48e-05	2.54e-05	4.04e-06
Contamana	8.53	61102	3.49e-05	4.92e-06	3.48e-05	3.50e-05	7.74e-05
Criollo	14.60	695	5.25e-03	4.27 e- 03	5.23e-03	5.27e-03	3.91e-03
Curaray	10.36	58213	4.45e-05	1.78e-05	4.44e-05	4.46e-05	1.18e-04
Guianna	8.66	4651	4.65e-04	7.74e-06	4.63e-04	4.67e-04	2.74e-04
Iquitos	4.23	49984	2.11e-05	5.88e-09	2.10e-05	2.12e-05	1.84e-05
Maranon	4.09	34037	3.01e-05	1.64e-08	2.99e-05	3.02e-05	1.68e-05
Nacional	4.66	26060	4.47e-05	9.76e-08	4.44e-05	4.49e-05	4.10e-05
Nanay	6.82	42429	4.02e-05	1.51e-07	4.00e-05	4.04 e- 05	1.33e-05
Purus	5.95	17357	8.57e-05	7.74e-06	8.54e-05	8.60e-05	1.23e-04

Table 1: Recombination rates in $4N_er/kb$, r/kb, and cM/Mb for all ten *T. cacao* populations. The N_e that was used for the transformation is also reported for each population, as are the lower and upper bounds of a 95% confidence interval for r/kb.

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Population	Ame	Con	Cri	Cur	Gui	Iqu	Mar	Nac	Nan
Amelonado	-	-	-	-	-	-	-	-	-
Contamana	$<\!\!2e-07$	-	-	-	-	-	-	-	-
Criollo	$<\!9e-05$	$<\!5e-13$	-	-	-	-	-	-	-
Curaray	<3e-05	< 3e-37	${<}5e-08$	-	-	_	-	-	-
Guianna	<3e-06	< 1e-37	< 7e-07	< 4e-20	-	-	-	-	-
Iquitos	<4e-08	$<\!\!6e-\!87$	<2e-11	<3e-16	<2e-29	-	-	-	-
Maranon	<6e-13	$<\!\!7e-\!77$	<2e-11	<2e-20	$<\!5e-33$	< 4e-64	-	-	-
Nacional	0.0015	<2e-43	0.0212	< 7e-14	<3e-06	$<\!\!6e-14$	< 3e-13	-	-
Nanay	0.0004	<2e-44	$<\!9e-11$	$<\!\!4e-16$	$<\!2e-21$	< 3e-39	<2e-38	< 9e-06	-
Purus	0.1782	$<\!\!4e-\!117$	$<\!\!2e-05$	<2e-29	< 1e-33	<2e-39	$<\!\!8e-\!43$	< 6e-27	< 2e-21

Table 2: Fisher's exact test p-values for pairwise comparisons of recombination hotspot locations between populations of $T.\ cacao$

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Measures	Observed %	Observed $\%$	Observed %	Mean $\%$	% Sim
	ubiquitous HS	all HS	whole genome	Sim	>ubiquitous HS
Retroelements	2.34	9.45	11.12	11.11	99.9
DNA transposons	1.94	1.64	1.10	1.10	5.4
Total	4.28	11.09	12.21	12.22	99.7

Table 3: Percentage of DNA sequences identified as either retroelements or DNA transposons, and total interspersed repeats. Observed values for the entire T. cacao genome, for all recombination hotspots (HS), and ubiquitous hotspots (hotspots in the same location in at least eight different populations). Also presented are mean percentage of these sequences for 1000 simulations of hotspots equivalent in size and count as the ubiquitous set and the percentile at which the observed value for the ubiquitous set is found in the distribution of the simulated set (Sim).

	TSSs	TTSs	Exon	Intron
	$(500 \mathrm{bp})$	$(500 \mathrm{bp})$		
Amelonado	1	1	0.602	0.527
Contamana	1	1	0.000	0.000
Criollo	1	1	0.000	0.000
Curaray	1	1	0.346	0.058
Guianna	1	1	1.000	1.000
Iquitos	1	1	0.000	0.000
Maranon	1	1	0.000	0.000
Nacional	1	1	0.000	0.000
Nanay	1	1	0.027	0.237
Purus	1	1	0.004	0.000

Table 4: Proportion of simulated chromosomes that presented a lower amount of hotspots intersecting with TSSs, TTSs, exons, and introns. TSSs and TTSs are considered to be the 500bp upstream and downstream of coding regions, respectively.

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Population	Mean Hotspot		
	Size (kb)		
Amelonado	6.9		
Contamana	6.1		
Criollo	6.1		
Curaray	5.8		
Guianna	8.6		
Iquitos	7.0		
Maranon	6.8		
Nacional	6.9		
Nanay	7.6		
Purus	6.3		
All hotspots	6.9		

Table 5: Average hotspot size (in kb) for hotspots detected in each population and average for all hotspots.

Pop	Position	Position	Genome	Genome	Position	Genome
	L95	U95	L95	U95	Range Quotient	Range Quotient
Amelonado	6.35e-10	7.67e-08	2.33e-10	3.13e-04	120.75	$1.34e{+}06$
Contamana	1.63e-06	1.34e-05	1.40e-09	2.64e-04	8.22	$1.88\mathrm{e}{+05}$
Criollo	8.75e-04	4.31e-03	5.35e-07	1.66e-02	4.92	$3.11\mathrm{e}{+04}$
Curaray	5.15e-06	2.63e-05	2.72e-09	2.02e-04	5.11	$7.40\mathrm{e}{+}04$
Guianna	9.76e-06	1.26e-04	1.81e-09	2.96e-03	12.90	$1.63\mathrm{e}{+06}$
Iquitos	3.50e-10	6.52e-08	1.58e-10	2.45e-04	186.29	$1.55\mathrm{e}{+06}$
Maranon	1.98e-09	7.40e-07	2.31e-10	3.52e-04	373.35	$1.52\mathrm{e}{+06}$
Nacional	4.80e-10	6.50e-08	2.76e-10	3.66e-04	135.60	$1.33\mathrm{e}{+06}$
Nanay	1.65e-09	3.06e-07	2.06e-10	3.52e-04	185.32	$1.71\mathrm{e}{+06}$
Purus	3.98e-08	5.24e-06	2.00e-09	6.35e-04	131.87	$3.18\mathrm{e}{+05}$

Table 6: The median of the upper and lower bounds of the 95% Credibility Interval for the trace of estimates of r from all positions in the genome are presented for each population. The upper and lower bounds of the 95% probability interval for the median estimate of r for each population is also presented. The quotients of the upper and lower bounds for each interval point to a much larger genome-wide variation in r than per-position variation in the trace for the estimate of r.

581 Figures

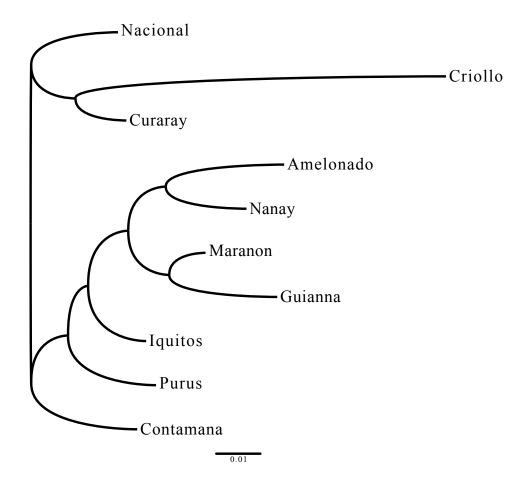


Figure 1: Drift tree for the 10 *T. cacao* populations. Modified from Cornejo et al. (2018)

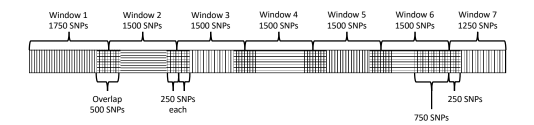


Figure 2: Example of the window layout for a 10,750 SNP chromosome. The 2,000 SNP long windows are represented by alternating horizontal and vertical lines and the overlaps between them are represented by square crosshatches. Braces above the chromosome indicate the regions from which recombination rates are extracted to generate the chromosome-wide recombination rates.

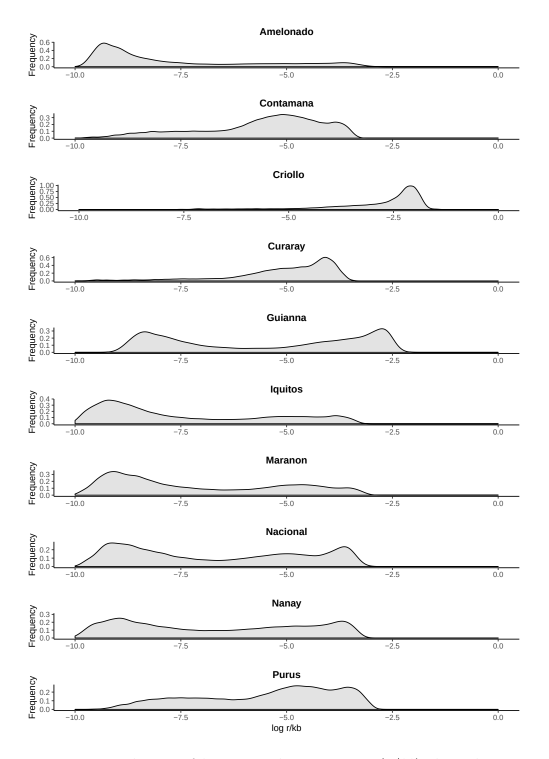


Figure 3: Distribution of log_{10} recombination rates (r/kb) along the genomes of the ten *T. cacao* populations.

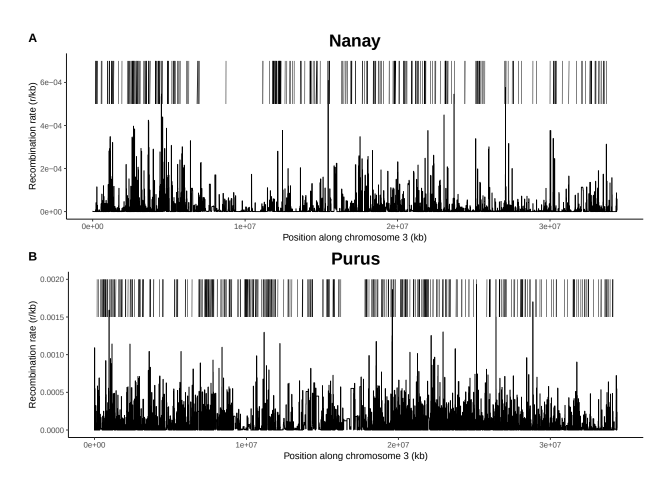


Figure 4: The third chromosomes of the Nanay (A) and Purus (B) populations were selected to exemplify the differences between populations in recombination rates (r/kb) and recombination hotspot locations (red vertical line segments).

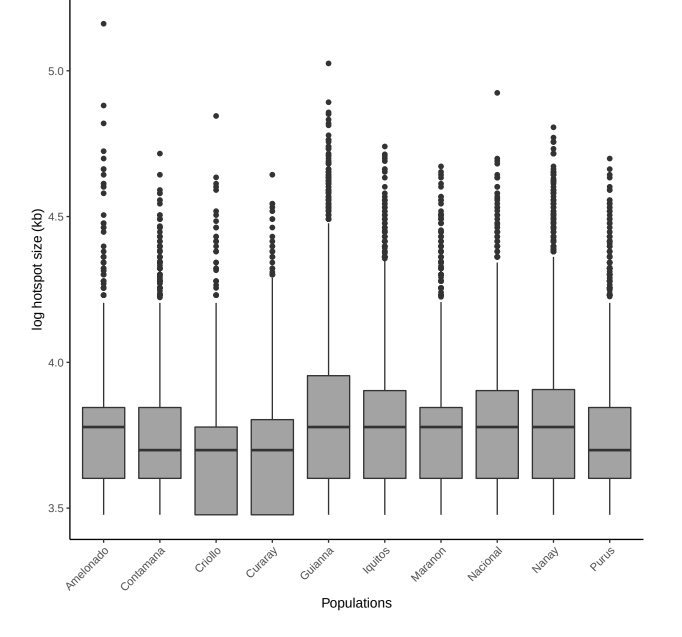


Figure 5: Boxplots of recombination hotspot sizes $(log_{10}(kb))$ by population. The horizontal line in the box represents the median value, while the points represent potential outliers.

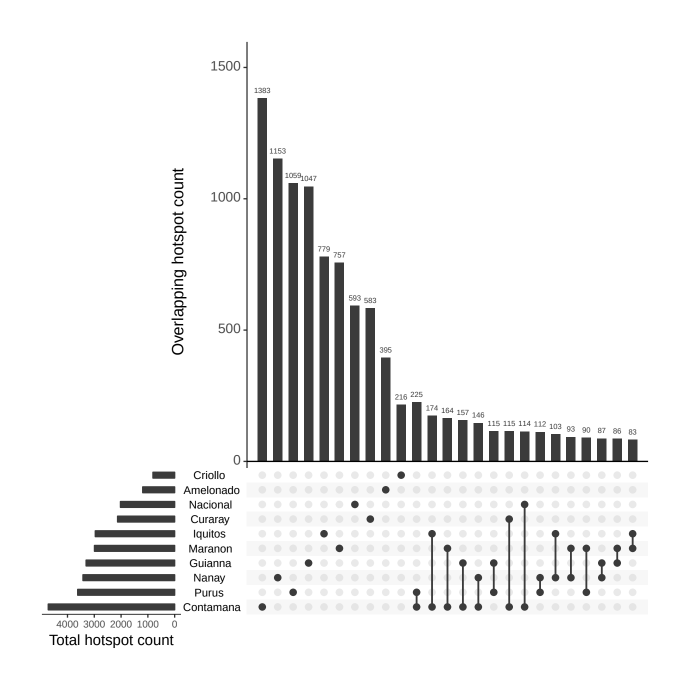


Figure 6: Upset plot showing amounts of shared hotspots. Horizontal bars represent total hotspots detected in a population, dots on the matrix represent the populations that share the hotspots represented in the vertical bar above them. The 25 largest subsets are shown.

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