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Identifying the diamond in the rough: a study of allelic diversity underlying flowering time adaptation in maize landraces

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1 **Landraces (traditional varieties) of crop species are a reservoir of useful genetic**
2 **diversity, yet remain untapped due to the genetic linkage between the few useful alleles**
3 **with hundreds of undesirable alleles¹. We integrated two approaches to characterize the**
4 **genetic diversity of over 3000 maize landraces from across the Americas. First, we**
5 **mapped the genomic regions controlling latitudinal and altitudinal adaptation, identifying**
6 **1498 genes. Second, we developed and used F-One Association Mapping (FOAM) to**
7 **directly map genes controlling flowering time across 22 environments, identifying 1,005**
8 **genes. In total 65% of the SNPs associated with altitude were also associated with**
9 **flowering time. In particular, we observed many of the significant SNPs were contained in**
10 **large structural variants (inversions, centromeres, and pericentromeric regions): 29.4%**
11 **for flowering time, 58.4% for altitude and 13.1% for latitude. The combined mapping**
12 **results indicate that while floral regulatory network genes contribute substantially to field**
13 **variation, over 90% of contributing genes likely have indirect effects. Our strategy can be**
14 **used to harness the diversity of maize and other plant and animal species.**

15 Maize (*Zea mays* subsp. *mays*) is a model organism with a legacy of a hundred years of
16 cytological, genetic, and biomolecular characterization². Maize displays high levels of genetic
17 diversity with low linkage disequilibrium (LD)^{3,4}, low population differentiation⁵, prevalent
18 migration⁶ and occasional introgression from wild relatives^{7–9}. More recently, experimental
19 populations like the Nested Association Mapping (NAM) populations^{10,11}, and large association
20 panels^{4,12} have allowed mapping and deployment of useful alleles for several quantitative
21 traits^{13–16}. However, most of the founder lines from these panels correspond to highly inbred
22 improved lines, many from temperate regions, capturing only a modest fraction of the total
23 diversity present in the species. In contrast, maize landraces span numerous ecogeographic
24 areas and harbor most of the diversity of the species. Nevertheless, maize landraces like many
25 other crops traditional varieties remain largely uncharacterized by genomics.

26 This study maps genes controlling flowering with two distinct methods: (1) Each of these
27 landraces come from environments to which they are well adapted. We used this adaptation as
28 the trait to identify genes driving large scale adaptation. (2) We mapped flowering time variation
29 in controlled field experiments through a novel, rapid, experimental design called F-One
30 Association Mapping (FOAM) (Figure 1). Briefly, FOAM consists of sampling single individuals
31 across numerous populations, which are genotyped and crossed to one or a small number of
32 common parents to derive F1 families. Subsequently GWAS is performed from multi-trial F1
33 progeny evaluation. Major advantages for this design are (a) capturing thousands of alleles
34 across populations, (b) maintaining the tractability of two alleles per loci per individual, (c) ample
35 replication of alleles increasing the power and accuracy for genetic effect estimation. The main
36 limitation of FOAM is that the nested evaluation of different subsets of F1 progeny by ecological
37 zone limit the ability to accurately estimate genotype by environment interaction effects.

38 Our maize landrace FOAM population used individuals from 4,471 accessions from 35 countries
39 in the Americas (Figure 2) grouped into three adaptation classes to account for altitude
40 adaptation (low, middle and high elevation). Similarly, the common parents and evaluation sites
41 were nested within adaptation class (methods, supplemental figure 1)^{17,18}. Landrace parents
42 were genotyped for close to one million SNPs using Genotyping by Sequencing¹⁹, and missing
43 data was imputed using BEAGLE4²⁰. Of the 4,471 accessions, 3,552 yielded F1 families
44 containing both genotypic profiles and sufficient progeny, 3,633 contained detailed passport
45 information which was used for mapping large scale adaptation, and 2,603 were present in both
46 mapping studies.

47 We first explored the effects of recombination frequency and geography-driven limited dispersal
48 on the distribution of genetic diversity in the landrace parents. Using Multidimensional Scaling
49 (MDS, Methods), we observed the first axis and second axes explained only 6.1% and 1.7% of
50 of the variance respectively, consistent with the low F_{ST} in maize landraces⁵. The first axis
51 separates among Mexican landraces, consistent with Mexican landraces having a deeper
52 coalescent and greater representation in the panel. The second axis was associated to a
53 latitudinal North to South gradient across Latin America representing isolation by distance
54 (Supplemental Figure 2). In addition, a Mantel test²¹ revealed a significant correlation between
55 geographic and genetic distances (Pearson's $r= 0.46$, $P\text{-value}<0.001$), with most of the
56 association driven by altitude. Despite this, phylogenetic analysis (Methods, Supplemental
57 Figure 3) revealed that adaptation class does not drive clade membership, which indicates that

58 alleles segregate across adaptation classes, with highland adaption being polyphyletic,
59 consistent with recent reports²². To study recombination, we estimated an approximate LD
60 statistic (Methods) which shows a distribution consistent with previous recombination
61 estimates^{23,24}, with higher recombination in gene-rich regions, and lower around centromeres.
62 Each chromosome displayed a unique recombination landscape, with the presence of half a
63 dozen high LD regions (Supplemental figure 4), which together encompassed 6.1% of the base
64 pairs of genome, accounting for 2.8% of the annotated coding genes. Together, these results
65 suggest that although at large scale geography and adaptation contribute to the distribution of
66 diversity, even with the large effective population size of landraces at the genomic scale a
67 complex recombination landscape limits the free segregation of alleles through increased LD.

68 Flowering time generally plays a crucial role in local adaptation of plants, and in maize flowering
69 time is a complex trait controlled by hundreds of small effect loci, many with rich allelic series
70 ^{4,14,25–30}. We used altitude and latitude from sampling location as traits for mapping local
71 adaptation, and the significance thresholds were chosen to maximize genic overlap rate
72 between flowering, altitude, and latitude (Methods, supplemental figure 5). For altitude, we
73 observed 58.4% of the significant SNPs corresponded to regions with higher LD. In particular,
74 INV4m, the 13Mb adaptive introgression from highland teosinte into maize^{8,31} was highly
75 significant. We also observed significance for the centromeres of chromosomes 2,5,6,8 and a
76 large region upstream of the centromere on chromosome 3. Outside this low recombination
77 regions, 366 genes showed significant association with altitude. For Latitude, we observed
78 13.1% of the significant markers were contained within low recombination regions, particularly
79 the centromere of chromosomes 5. In total across all Latin America, 1498 genes showed
80 significant association with latitude, of which 395 of were shared with altitude. The minor allele
81 frequency distribution of the significant alleles indicated that many are shared across clades and
82 landraces, which was very distinct from the neutral distribution (Figure 3). These 1498 genes
83 appear to be the main contributor to large scale environmental adaptation to altitude and latitude
84 – the key drivers of flowering time.

85 To study the genetic basis of flowering time, we conducted field evaluation on F1 progeny
86 across 22 trials and 2 years in 13 locations across Mexico, with each trial containing a different
87 subset of the collection to maximize number of accessions evaluated (Methods, supplemental
88 table 1). Phenotypic data was analyzed independently for each trial using a mixed linear
89 (Methods), yielding 18,797 accession parent-environment estimates for each male and female

90 flowering time. We performed genome wide association for days to male and female flowering
91 using a mixed linear model (Methods). In total 72% of the associated SNPs were significant for
92 both male and female flowering, as expected from the overlapping genetic control¹⁴. There was
93 a significant contribution of low recombination regions in flowering time variation, parallel to that
94 of latitude and longitude, with a 20-fold enrichment for significant SNPs at high LD regions
95 (Pearson's chi-squared, p-value < 2.2e-16). In particular, significant variants included the
96 centromeres of chromosomes 3, 5, and 6, INV4m, and a 6Mb region on chromosome 3
97 beginning at 79Mb. The 6Mb region on chromosome 3 has a segregation similar to INV4, and
98 together with its increased LD suggests it might be an inversion. In NAM this putative inversion
99 and the centromere comprise a single QTL for flowering time¹⁴. For the centromere of
100 chromosome 5 there were 3 distinctive alleles segregating in the landraces, all present in the
101 NAM population (supplemental figure 6). The inverted allele of INV4m, although absent in
102 temperate material, segregates at high frequency in highland landraces (supplemental figure 7),
103 where it has very large additive effect advancing flowering by three days, the largest effect for
104 flowering time in maize to date. Both homozygous alleles from the putative inversion on
105 chromosome 3 segregate across our maize landrace panel and the NAM population. Compared
106 to INV4m, this locus displays a more modest effect on flowering time. The heterotic effect of the
107 centromere of chromosome 5 on yield³², potentially product the complementation of deleterious
108 mutations²³, suggests that the significant inversions and centromeres may similarly affect
109 flowering time through heterotic effects leading to more vigorous plants, which in maize
110 generally results in earlier flowering.

111 Outside the structural variants, we observed 881 and 883 genes (around 2.2% of genes) with
112 significant association for days to female and days to male flowering respectively (Supplemental
113 Tables, Figure 4). To further characterize the regions associated with flowering time, we looked
114 for gene ontology enrichment and gene expression using the maize transcription
115 atlas³³(Methods), and compared the significant genes to a candidate gene list containing genes
116 characterized in other populations, known to interact in the maize flowering time regulatory
117 network³⁴ as well as the 25 members of the *Zea mays* CENTRORADIALIS (ZCN) gene family³⁵.
118 Overall the associating genes tended to be expressed in anthers, and enriched in general
119 metabolic processes, with the genes known to be part of the regulatory network being more
120 expressed in immature cob and the tip of the leaf at V5 stage and enriched for regulatory
121 processes (Supplemental figure 8). We observed a significant enrichment in flowering time

122 candidate genes compared to the rest of the genome (Fisher's Exact Test p-value = 4.3×10^{-7}).
123 In total 10 and 12 candidate genes representing the circadian clock, photoperiod, gibberellin
124 acid, and circadian clock pathways displayed significant associations with male and female
125 flowering respectively. Out of these, nine were common for both types of flowering. The most
126 significant hits corresponded to VGT1^{36,37}, one of the largest known GxE QTL, and ZCN8^{35,38},
127 the maize florigen and homolog to *FT* in *Arabidopsis* (Figure 5). ZmCCT, the largest
128 photoperiod QTL²⁹ was only modestly significant for latitude, and significant only for days to
129 female flowering, probably a result of non-inducing sampling and trial locations. In particular for
130 the gene *d8*, a locus with cryptic association with flowering time³⁴, we observed significance for
131 this gene around 50 and 100kb up and downstream the coding region for latitude, altitude, and
132 both male and female flowering, overlapping with the region previously observed to display
133 divergent selection associated with climate adaptation³⁹. In addition, the distribution of the
134 flowering time associating genes displayed a significant geography effect, with 56 and 52 genes
135 in common with altitude and latitude respectively. In general, the minor alleles for flowering time
136 tended to be associated with high elevation, and northwest coordinates, however the minor
137 allele frequency distribution of the significant SNPs was different to that of the alleles significant
138 for altitude and latitude, having a significant shift towards low frequency polymorphisms (Figure
139 3). Together, these results support the model of infrequent variants in recurrent regulatory
140 genes underlying the genetic control of flowering time variation in maize, with adaptive alleles
141 segregating across populations, and their distribution matching the fitness optimum according to
142 geographic variation. In particular, the high overlap between significant SNPs for altitude and
143 flowering time suggests that for tropical maize flowering time adaptation is very relevant for
144 changes in elevation, which affects among others spectral composition and intensity of incident
145 light, as well as the incidence of heat and cold stress. In contrast, the lower overlap between
146 latitudinal and flowering time associating SNPs could be to the sampling from non-photoperiod-
147 inducing latitudes, potentially leading to latitudinal flowering time adaptation being relevant for
148 other biotic (disease pressure) and abiotic (soil pH, precipitation) stresses.

149 We assayed the potential for predicting flowering time in the landraces using either all our high
150 density genetic markers or just the markers significantly associated with the trait. We performed
151 genome wide prediction using gBLUP independently for each trial (Methods). The average 5-
152 fold cross-validated prediction accuracy was 0.45 across trials for both male and female
153 flowering time and, and as high as 0.7 for some trials (Supplemental figure 9). Genomic

154 prediction accuracy between the top genes from GWAS was equivalent to that of 30,000
155 random evenly distributed SNPs, highlighting their potential use for breeding of the significant
156 markers. Intriguingly prediction accuracy was not correlated with our other heritability estimate
157 (Pearson cor=0.22), which could be an effect of the differences in the genetic variances and
158 sample sizes across all trials. Together the good predictive ability of the significant regions for
159 genomic selection shows the potential to greatly speed the breeding of new adapted varieties
160 with exotic beneficial alleles.

161 Crop landraces are an incredible source of diversity that will be necessary to adapt our crops to
162 next century of climate change. However, their tremendous diversity and genetic load prevent
163 them from being efficiently tapped without a genomic index. This research lays out two
164 complementary strategies for tapping this diversity. The geographic associations are powerfully
165 identifying the adaptive loci, which appear to be common and shared, and are unlikely to be
166 deleterious given their high frequency. This extensive sharing is probably the result of
167 outcrossing and extensive migration throughout Latin America in last several millennia. The
168 limitation of this approach is that correlated traits and adaptations are being co-mapped. The
169 novel FOAM field trial associations, while substantially overlapping, are showing the impacts of
170 deleterious and private mutations and their complementation in these hybrid trials. These
171 deleterious alleles have been the bane of breeders wanting to tap landrace diversity. The
172 strategy for tapping this diversity should be use the overlapping genes and alleles of the two
173 separate approaches— as these have proven to be adaptive *and* target the trait of interest. The
174 breeding could use standard genomic selection or genome editing. This provides an efficient
175 strategy to tap landraces diversity and allow our crops to adapt to faster changes than ever had
176 in the past.

Methods

Mating design and phenotypic evaluation

The mating design for the maize landrace FOAM population consisted of crossing each accession male to single cross hybrid females of matching adaptation. Leaf tissue of the landrace individual was collected for genotyping. The progeny evaluation trials were performed across 2 years in 13 locations across Mexico using an augmented row-column design, which includes systematic checks in field rows and columns⁴⁰. There were between 288 and 1928 accessions per trial, with an average of 834 (Supplemental table 1). Over half of the accessions were replicated in 5 or more trials, with a maximum value of 13 trials per accession and a minimum of 1. For each trial, each experimental row contained between 10 and 25 progeny plants. The replication across trials together with the use of systematic checks across experimental fields provides sufficient allelic replication for accurate estimation of genetic effects. Flowering time was measured in each trial following the maize standard, i.e. the number of days from planting until half of the individuals within a plot displayed silks for female flowering or anthers in half of the central spike for male flowering.

Genotyping

Accessions used as male parents were genotyped using GBS¹⁹, with ApeKI as the restriction enzyme to a replication level of ~96 individuals per sequencing plate. Approximately 8x10⁹ sequencing reads were generated using an Illumina HiSeq for the landrace accessions and sequence reads were analyzed jointly with another 40,000 maize lines as part of the GBS Build 2.7 using TASSEL⁴¹. For association analyses, missing data was imputed using BEAGLE²⁰, which has been shown to yield the best current accuracies in maize heterozygous material ($R^2=0.68$)⁴². After imputation, SNPs were filtered for minor allele frequency greater than 1%

resulting in approximately 500,000 biallelic markers across the genome. GBS non-imputed markers can be accessed at <http://hdl.handle.net/11529/10034> and imputed GBS markers at <http://hdl.handle.net/11529/10035>

Diversity Assessment

For the Mantel test²¹, we calculated the pairwise Euclidean distance matrix based on the geographical data from the accessions (latitude, longitude, and altitude, <http://mgb.cimmyt.org/gringlobal/search.aspx>). The genetic distance matrix was estimated from a genome wide random sample 30,000 non-imputed markers using TASSEL. The distance matrix was used for estimating the Neighbor-Joining tree using TASSEL. Multidimensional Scaling (MDS) was performed on the genetic distance matrix using the cmds function in R.

Recombination

Our LD statistic consisted in estimating the correlation between markers across the genome at 100 site windows using all homozygote and heterozygote non-imputed markers with the LD function on the software TASSEL. For comparing the LD and recombination values, we estimated the correlation at 1Mb sliding windows between (1) the log10 median LD estimate (2) the log median crossover probabilities estimated using the American and Chinese Nested Association Mapping populations²³, and (3) the log median population recombination rates (rho) estimated both for improved lines and landraces Hapmap2 project²⁴. Our LD estimates displayed a negative correlation with gene density ($r=-0.57$) and NAM crossover probability²³ ($r=-0.45$). We observed a modest negative correlation ($r=-0.33$) with a population genetic estimate of historical recombination (rho)^{23,24}. High-LD regions were defined based on the change in slope of global median LD (Figure 5- Figure supplement 1) as those segments that

had a median LD greater than 0.01. In total, there were 256 high LD regions encompassing 7.8% of the genome.

Genome wide association with altitude and latitude

We performed Genome Wide Association using a generalized linear model with altitude and latitude as response variables and markers filtered at 1% frequency as explanatory variables. Altitude and Latitude were recorded during field sampling of the original accessions. In order to establish a significance threshold to avoid excess of false positives, we estimated the overlap rate using the most significant flowering time GWAS SNPs. Overlap Rate was defined as the set of overlapping SNPs between the top flowering time SNPs and either altitude or latitude, divided by the union of the sets across significance thresholds from 0.001 to 0.01. Significance thresholds chosen were 0.005 for altitude and 0.01 for latitude (Supplemental figure 4). Heritability estimates were 0.88 for altitude and 0.85 for latitude, estimated LDAK⁴³ with a single Kinship matrix, estimated with all the Beagle4 imputed markers, and the matrix was estimated from the algorithm implemented in GCTA⁴⁴.

Analyses of structural variants

In order to infer the underlying haplotypes for the centromeres of chromosomes 3,5,6, as well as INV4 and the high-LD region on chromosome 3, we first estimated a genetic distance matrix for each locus using the non-imputed markers. The distance matrices were then analysed using multidimensional scaling. The centromere of chromosome 5 segregates in the landraces with three distinct homozygous haplotypes and their corresponding heterozygote pairs. The region around the centromere of chromosome 6 was 12 Mb in size, includes the centromere and a large pericentromeric region that expands out in both directions; it displayed a similar pattern to the centromere of chromosome 5 however distinct alleles were not called due to the excess of

heterozygous individuals between the homozygous classes, probably reflecting recombinant haplotypes. The centromere of chromosome 3 displayed a more complex pattern of distance than the other two associating centromeres, likely due to the presence of more than three segregating haplotypes. For INV4, we observe two distinct alleles and the heterozygote. We observed the allele is fixed in many of CIMMYT improved lines (Table 3), including those used as parents for the highland test crosses in the present experiment.

Analysis of phenotypic data

To estimate the breeding values of the landrace accession parent, for each trial a mixed linear model was fitted using restricted maximum likelihood method, in ASREML (V 3.0), using the progeny's calendar days to male or female flowering as a response variable. Of the 23 trials planted, one was excluded because flowering time data was not collected according to protocol. The models included fixed effects for checks, tester, and hybrid and a random effect of accession in a complete nested model. Including in the model the random effect of row and column and using an autoregressive model of order 1 in row and columns controlled experimental noise product of field variation. All the random effects were considered independent one from each other. The model used can be expressed as follows:

$$y_{ijklm} = \mu + \gamma_i + \lambda_j + \alpha_k + \beta_{l(k)} + \delta_{m(kl)} + \epsilon_{ij}$$

where

y_{ijklm} : is the response variable,

μ : is the overall mean,

γ_i : is the effect of the i-th row, $\gamma_i \sim N(0, \sigma_1^2)$

λ_j : is the effect of the j-th column, $\lambda_j \sim N(0, \sigma_2^2)$

α_k : is the effect of the k-th group, $k=1,..,K, K+1$, if $k \leq K$ the group is a check, the group $K+1$ is the average of testers.

$\beta_{l(k)}$: is the effect of the l-th tester in group $K+1$

$\delta_{m(kl)}$: is the effect of the m-th accession in the tester k in group $K+1$, $\delta_{m(kl)} \sim N(0, \sigma_{kl}^2)$

ε_{ij} : is the experimental error

for the experimental error we assume the following distribution:

$\varepsilon \sim N(0, \Sigma)$, with $\Sigma = \Sigma_r \otimes \Sigma_c$ and

$$\Sigma_r = \begin{bmatrix} 1 & \rho_r^1 & \rho_r^2 & \cdots & \rho_r^{d-2} & \rho_r^{d-1} \\ \rho_r^1 & 1 & \rho_r^1 & \cdots & \rho_r^{d-3} & \rho_r^{d-2} \\ \rho_r^2 & \rho_r^1 & 1 & \cdots & \vdots & \vdots \\ \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ \rho_r^{d-2} & \rho_r^{d-3} & \rho_r^{d-4} & \cdots & 1 & \rho_r^1 \\ \rho_r^{d-1} & \rho_r^{d-2} & \rho_r^{d-3} & \cdots & \rho_r^1 & 1 \end{bmatrix} \quad \Sigma_c = \begin{bmatrix} 1 & \rho_c^1 & \rho_c^2 & \cdots & \rho_c^{d-2} & \rho_c^{d-1} \\ \rho_c^1 & 1 & \rho_c^1 & \cdots & \rho_c^{d-3} & \rho_c^{d-2} \\ \rho_c^2 & \rho_c^1 & 1 & \cdots & \vdots & \vdots \\ \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ \rho_c^{d-2} & \rho_c^{d-3} & \rho_c^{d-4} & \cdots & 1 & \rho_c^1 \\ \rho_c^{d-1} & \rho_c^{d-2} & \rho_c^{d-3} & \cdots & \rho_c^1 & 1 \end{bmatrix}$$

Genome wide association with flowering time

Association analysis was performed in two steps for all trials using a linear mixed model. For each trait (days to male and female flowering) two models were fitted, one with the trait BLUPs as response variable and another one with the standardized values of the same BLUPs. This was done in the absence of growing degree units, to verify the consistency of the results given the uneven variances for the trait across the various trials. The first step models included the fixed effects for trial (categorical); population structure in the form of 10 MDS weights

(numerical) that together explained around 13% of the genetic and 10.6% of the phenotypic variances; and the effect of the hybrid used as parent for each accession's cross. The random effect of relatedness was added to both models in the form of a kinship matrix. The kinship matrix was estimated using the same subset of SNPs as the MDS weights. The mixed model was fit using the R package EMMREML (<http://cran.r-project.org/web/packages/EMMREML/index.html>). Residuals were obtained from those models and fitted in the second step models as a response variable for the single marker analysis using R, with marker nested within trial.

The model equation used was

$$Y_{ijk} = \mu + T_i + H_{ij} + Q_{ijk} + K + \epsilon_{ijk}$$

where

y_{ijk} : is the response variable,

μ : is the overall mean,

T_i : is the effect of the i-th trial

H_j : is the effect of the hybrid parent

Q_k : is the population structure effect containing 10 weights from MDS

K : is the random effect of relatedness through kinship matrix K estimated from 30,000 random SNPs

ϵ_{ijk} : is the residual error

In the second step of the association model, the residuals from the first model were fitted as a response variable in the following model

$$Y_i = S[t] + \epsilon_i$$

Where Y is the residual, S is the SNP effect and is nested within trial t . The model tests the null hypothesis that the effect of each SNP is 0 in all trials.

$$H_0 : S = 0$$

The alternative hypothesis is that the SNP has an effect on any trial. The reason for testing this hypothesis is that the effect of each SNP can, and often does, change on value and direction depending on its segregation on the population and its phase with the causal polymorphism. We consider as significant the top one percent of the SNPs based on p-value, which all had $-\log_{10}$ p-values greater than 18.

Code availability

R implementation of the ASREML code used for estimation of breeding values can be found at
<http://data.cimmyt.org/dvn/dv/cimmytswdvn;jsessionid=c1de29cab7c37b41098fd8ad6684>

The mixed model was fit using the R package EMMREML
(<http://cran.rproject.org/web/packages/EMMREML/index.html>)

All other additional scripts are available through github user jar547@cornell.edu

Significance at genic regions

We reasoned that significance at candidate genes would depend on local LD and genotype coverage, therefore a higher proportion of significant SNPs around candidate genes would be indicative of association at the gene itself rather than at the entire LD block or because of higher genotype coverage. On that account, we looked at significant associating SNPs within 50 kb up and downstream of candidate genes. Of all the candidate genes, only PhyB1, GL15 and ZCN13 are in the high-LD set and therefore were excluded from this analysis.

Genome wide prediction was performed with using the software GAPIT⁴³. The models were run for each trial, and accuracy was measured from performing 5 fold cross validation in 10 replicates for each trial. Two models were run for each trait/trial. One model used a kinship matrix estimated 1 SNP for each of the associated genomic regions, while the other used 30,000 random SNPs for the estimation of the kinship matrix. All models included 10 MDS weights to account for population structure.

Expression across tissues

We used the transcription data from the maize atlas³³ for the following 11 tissues: 16 days after pollination embryo, 16 days after pollination endosperm, 6 days after silking primary root, tip of stage 2 leaf at V5 plant stage, base of stage 2 leaf at V5 plant stage, 13th leaf at V9 stage, 13th leaf at R2 stage , silk, anthers, Immature cob at V18 stage, 4th internode at V9 stage, and stem and shoot apical meristem at V4 stage. We used the standardized expression values, and estimated for each gene what tissue it was most expressed at. We then performed a chi-squared test for each tissue to test if there were more genes expressed at the candidate or associating genes than expected under the null model of equal levels of the global expression pattern.

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Figures

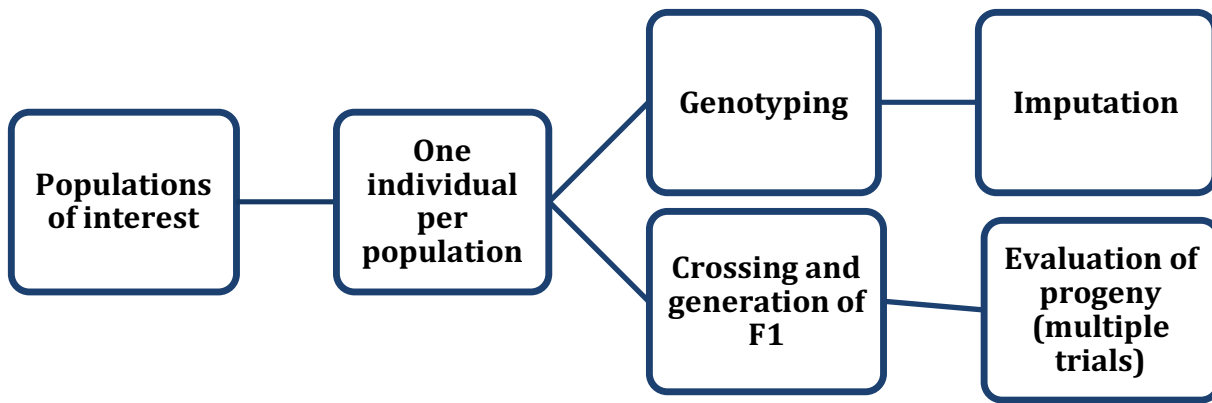


Figure 1. Experimental design. One individual from each of up to thousands of individuals is genotyped and used as parent. Progeny are then evaluated for multiple years/locations to estimate the genetic contribution of the original individual and phenotypic and genotypic data are used for Genome Wide Association

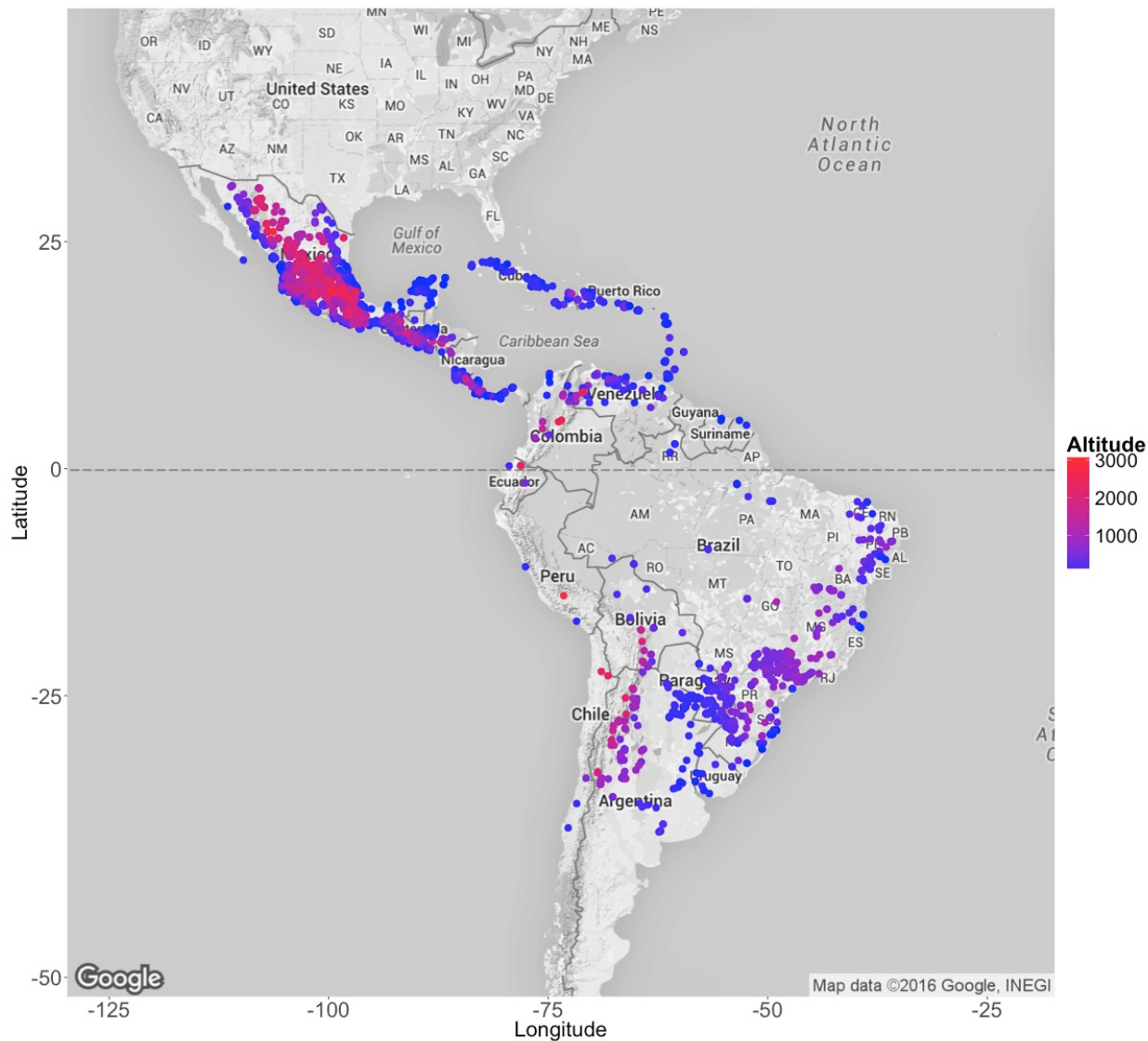


Figure 2. Geographic coordinates of original sampling sites of landrace accessions. Color gradient corresponds to altitude

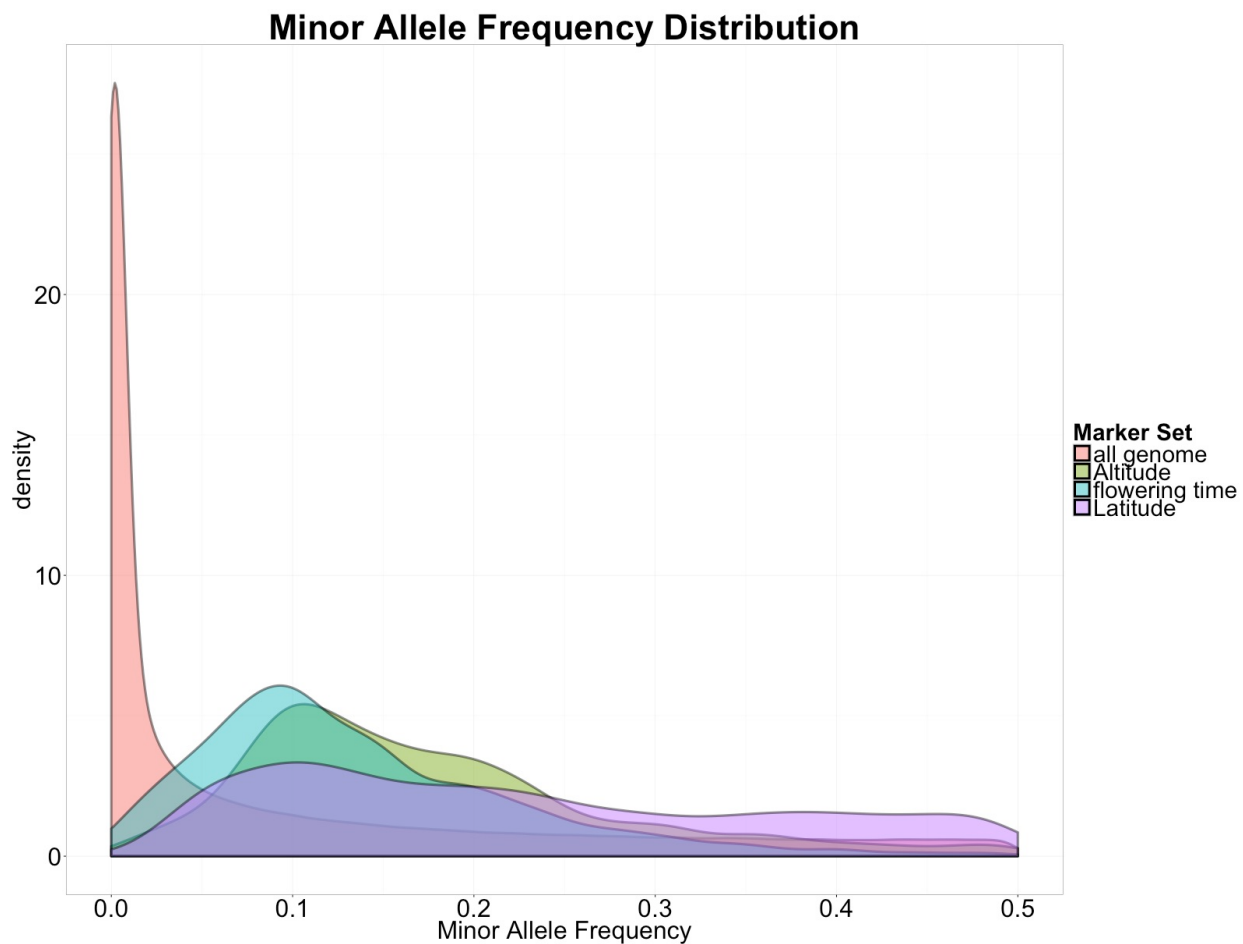


Figure 3. Minor Allele frequency distribution for all segregating SNPs, as well as the most significant SNPs for each trait.

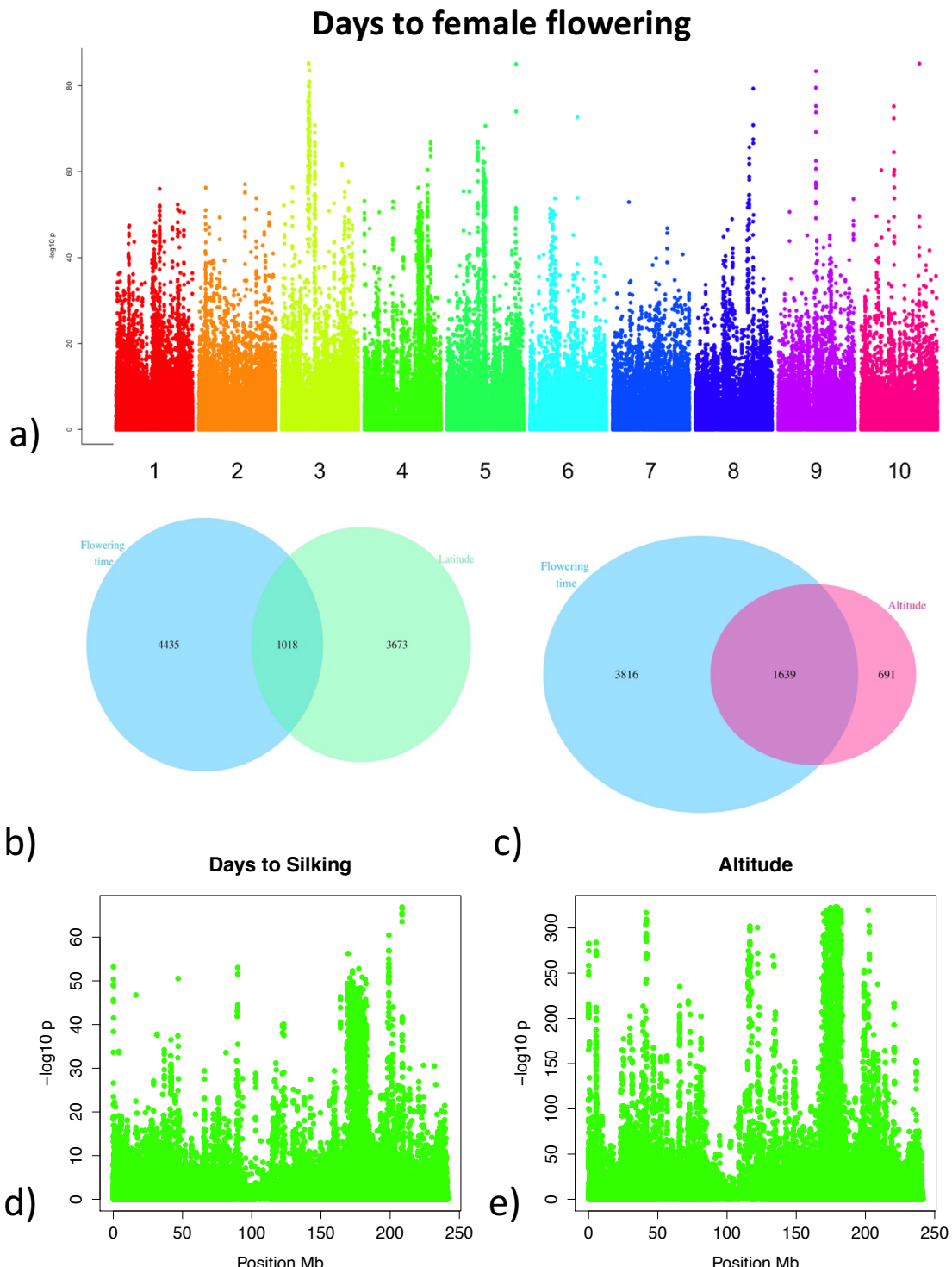


Figure 4. a) Manhattan plot for Days to silking. b) Local Manhattan plot for chromosome 4 for days to silking and c) Altitude. The large region with significance corresponds to INV4m, the

adaptive introgression from highland teosinte to highland maize d) Overlap between significant SNPs for flowering time and latitude and e) altitude

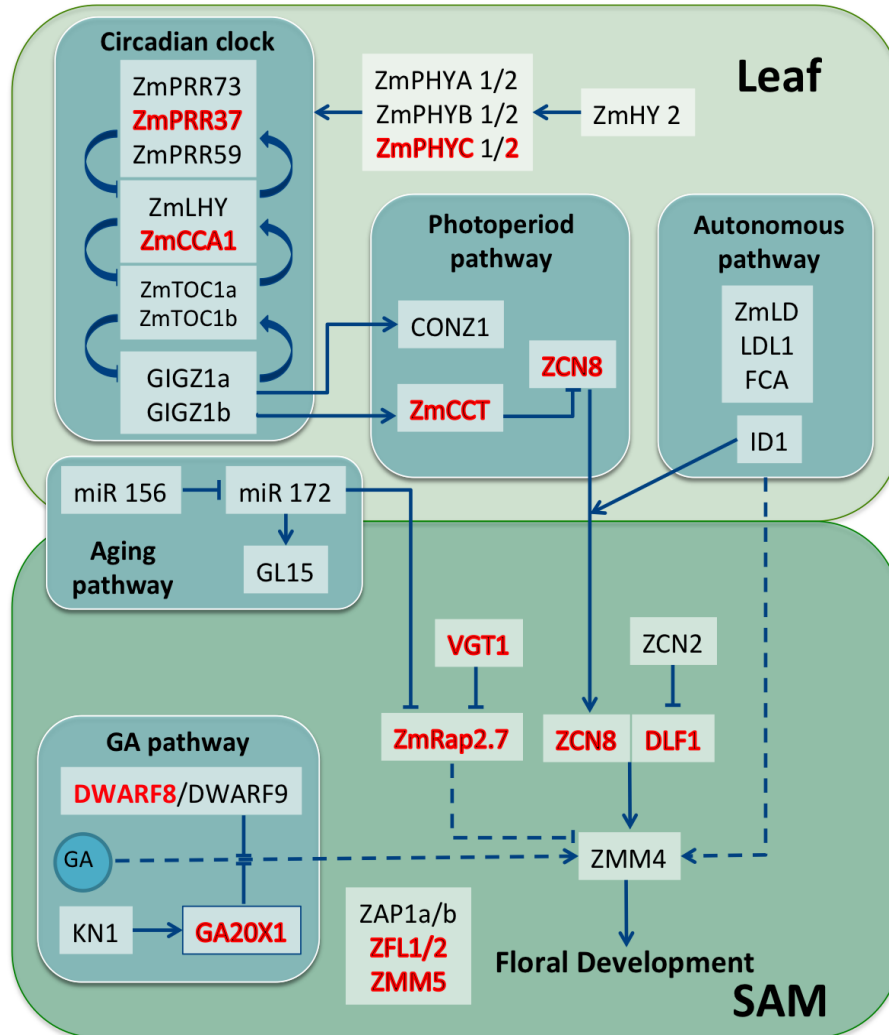
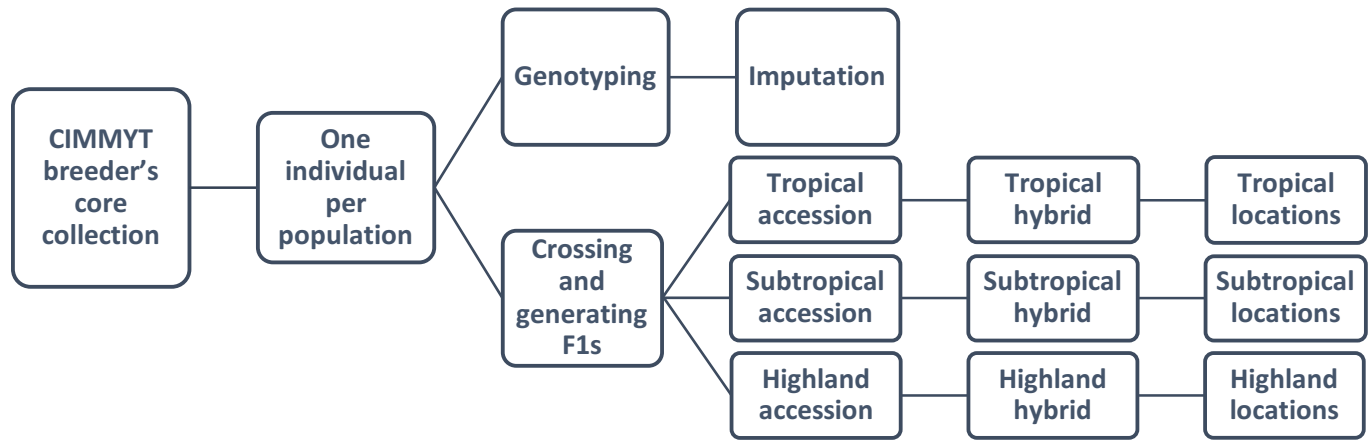
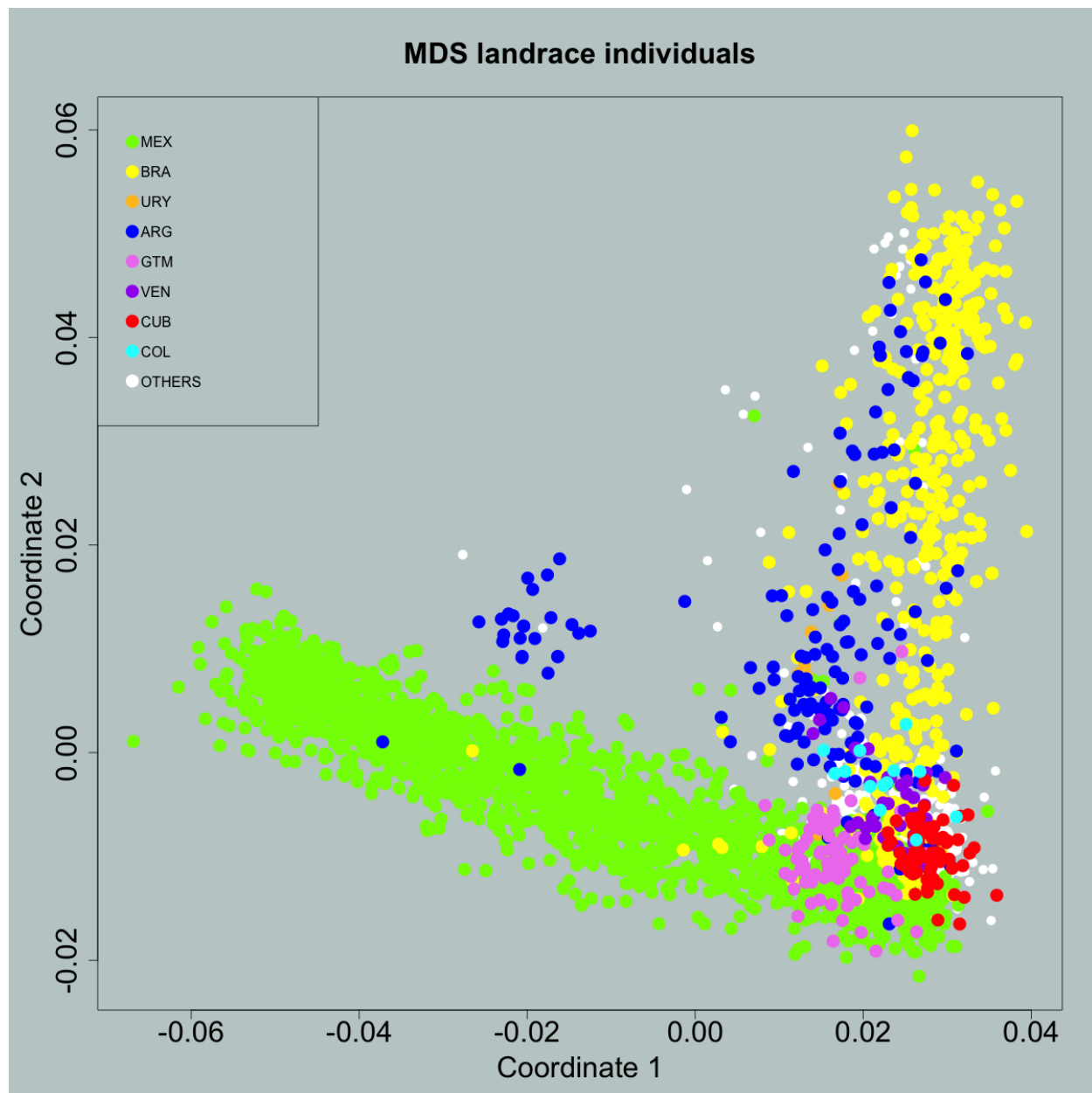


Figure 5. Flowering time pathway from Dong, et al³⁴, showing the genes involved in flowering time at the leaf and Shoot Apical Meristem (SAM). The genes highlighted in red displayed significant association with flowering time in our study.

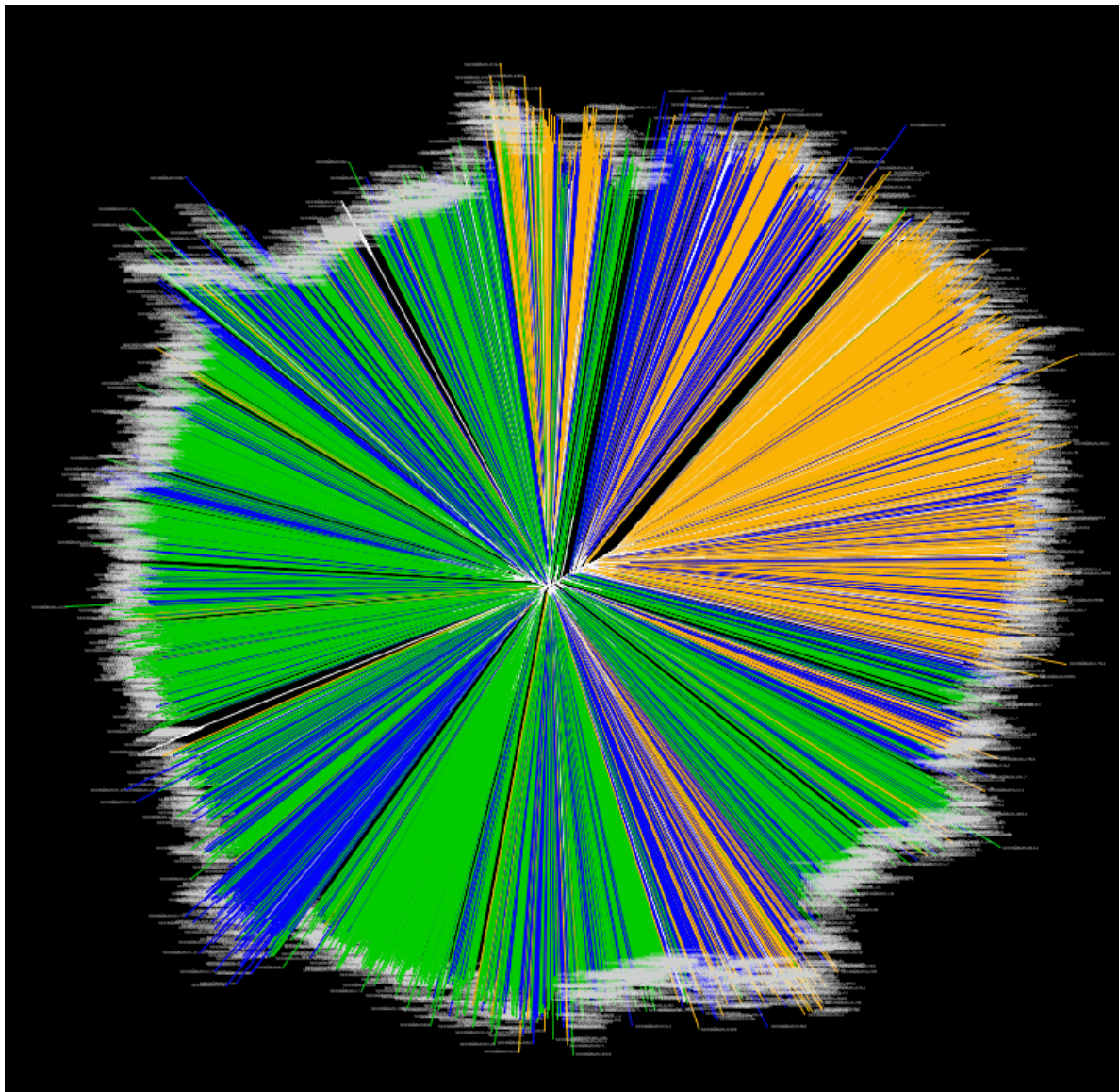
Supplemental Figures



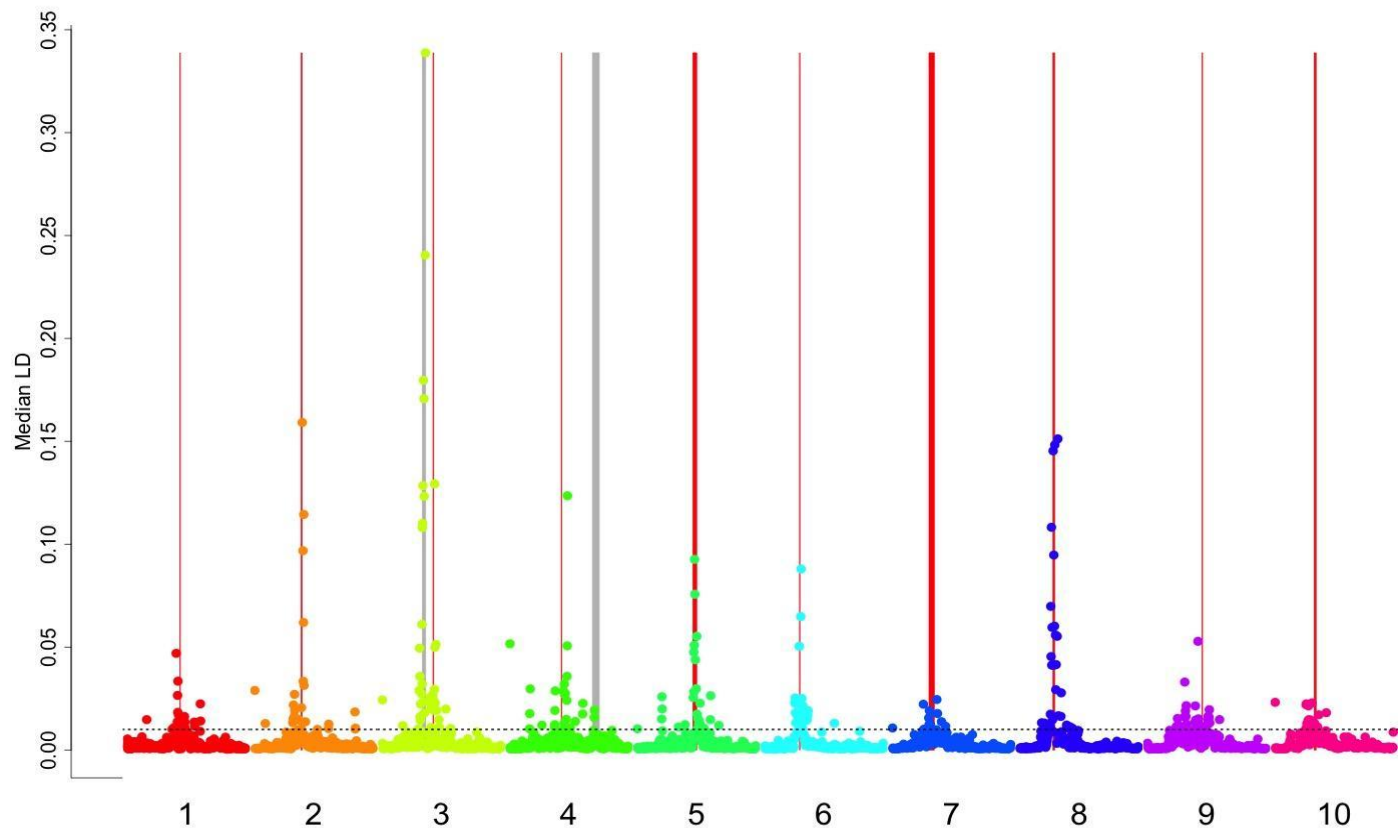
Supplemental figure 1. FOAM design with crossing and evaluation nested within adaptation.



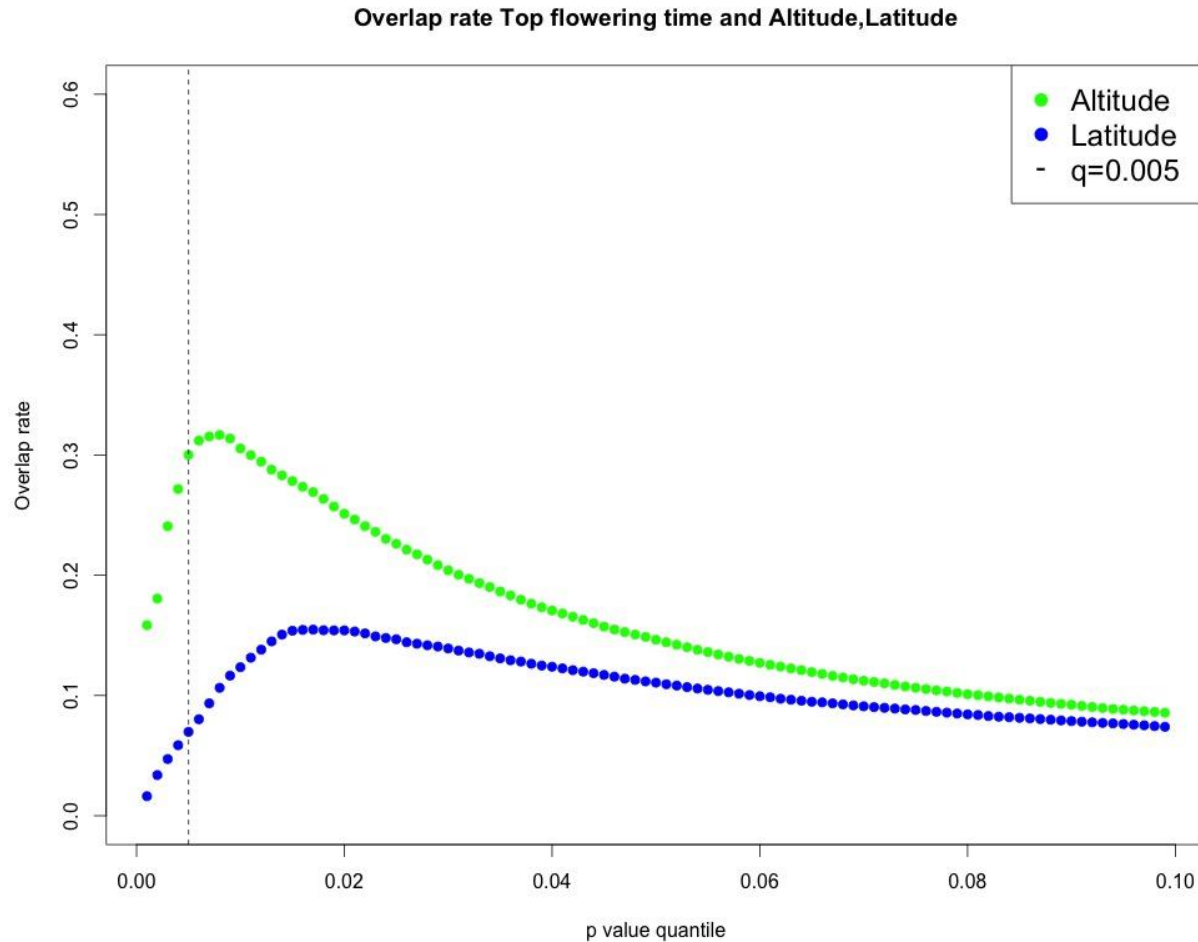
Supplemental figure 2. First 2 Principal Coordinates from Multidimensional scaling of the genetic distance among accessions



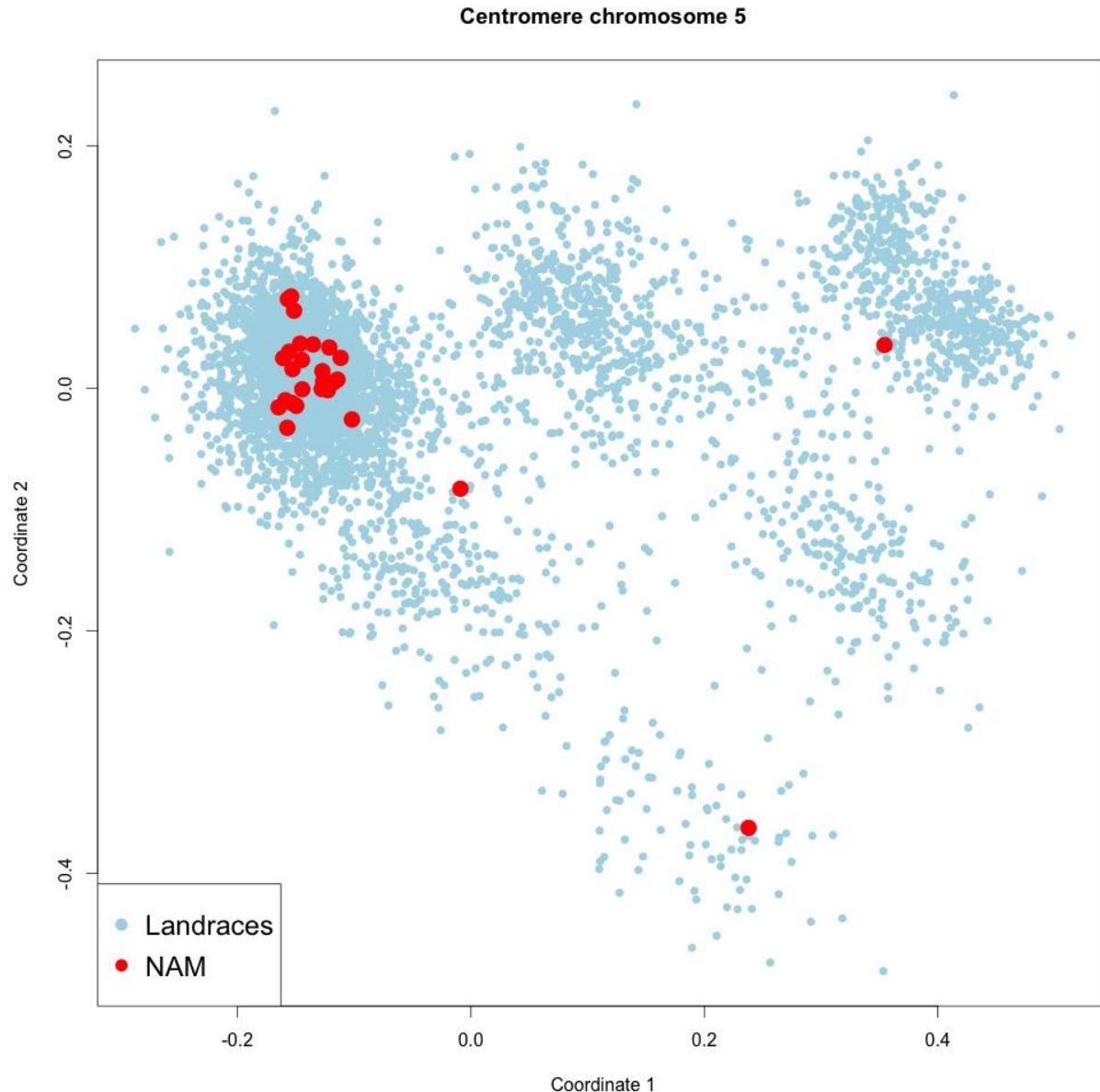
Supplemental figure 3. Neighbor-joining tree. Adaptation classes are colored green for low elevation, blue for mid elevation and orange for high elevation



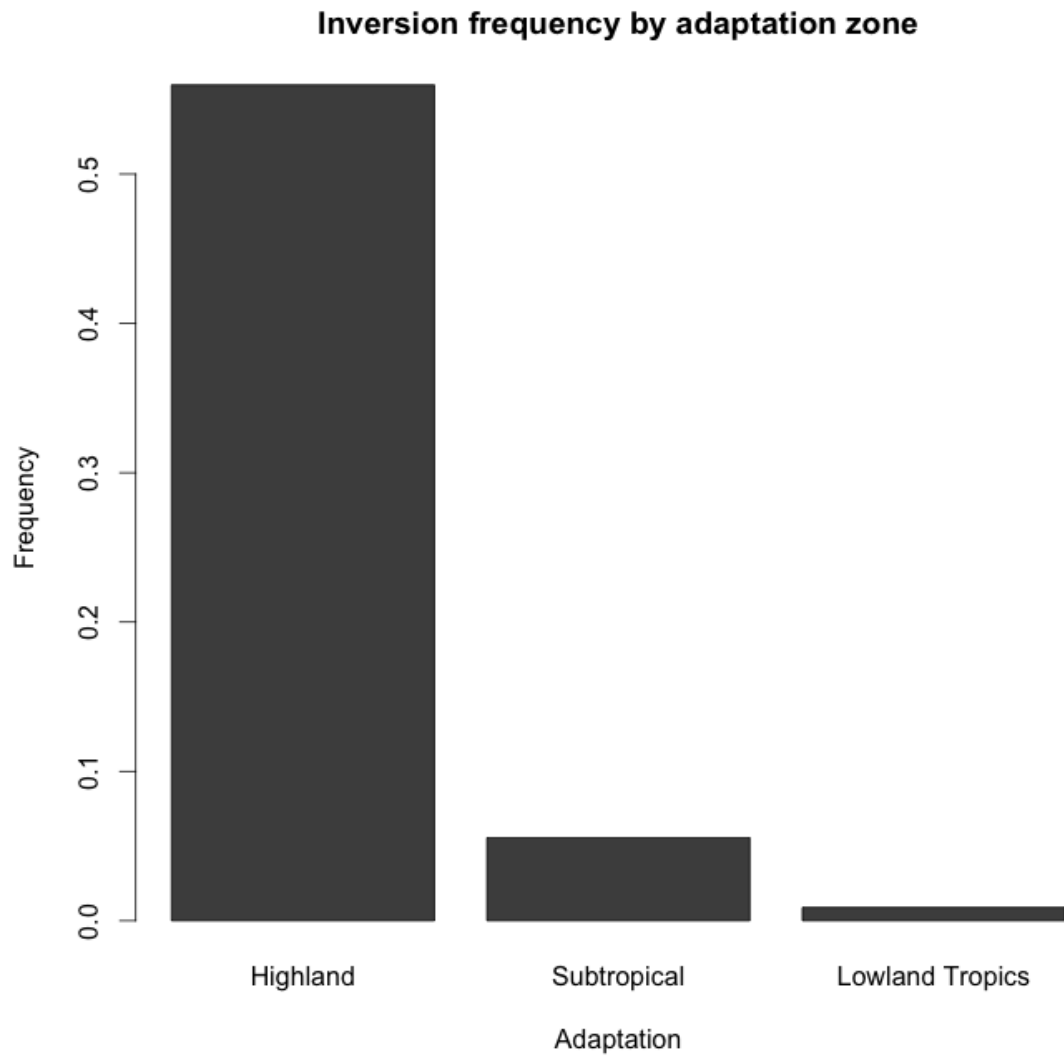
Supplemental figure 4 Genome wide view of the LD empirical threshold. Red shaded areas represent the centromeres, gray shaded areas represent inversions on chromosomes 3 and 4, and the dashed horizontal line represents the empirical LD threshold



Supplemental figure 5 Overlap rate between the top associating SNPs with flowering time and altitude, latitude at various p-value thresholds.

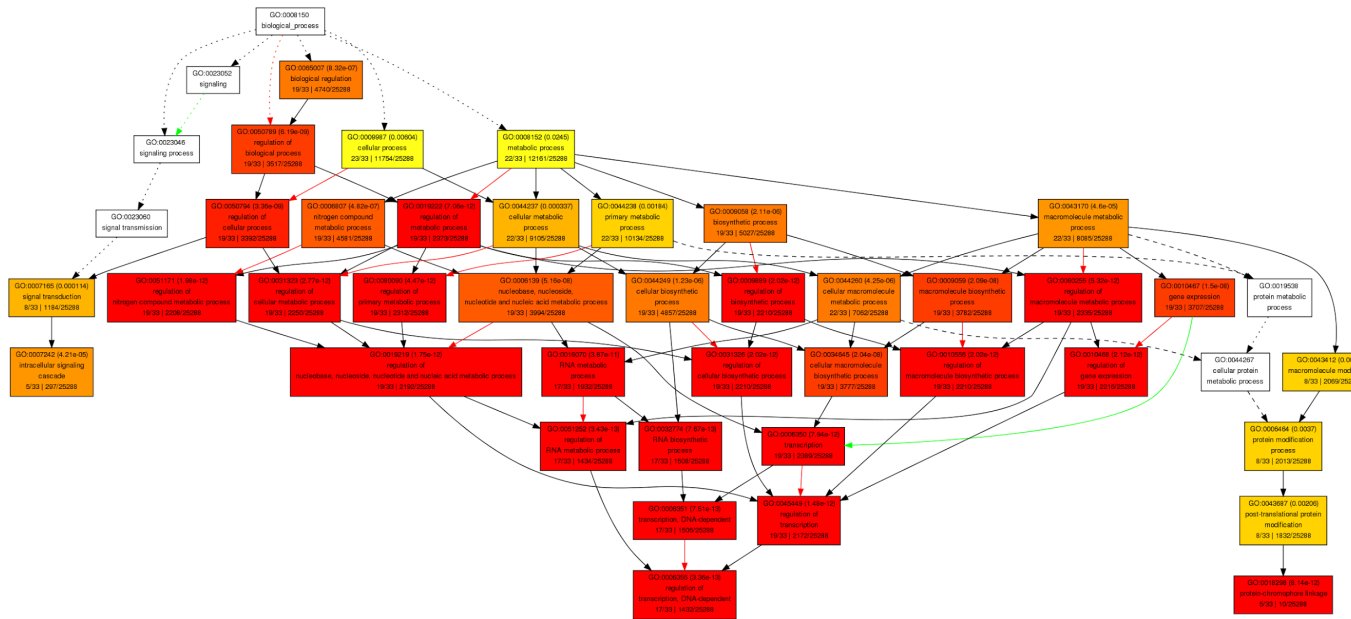


Supplemental figure 6. MDS of centromere of chromosome 5 for the FOAM landrace accessions and the NAM founders: Topright: II14H. Bottom: P39. Middle: CML333

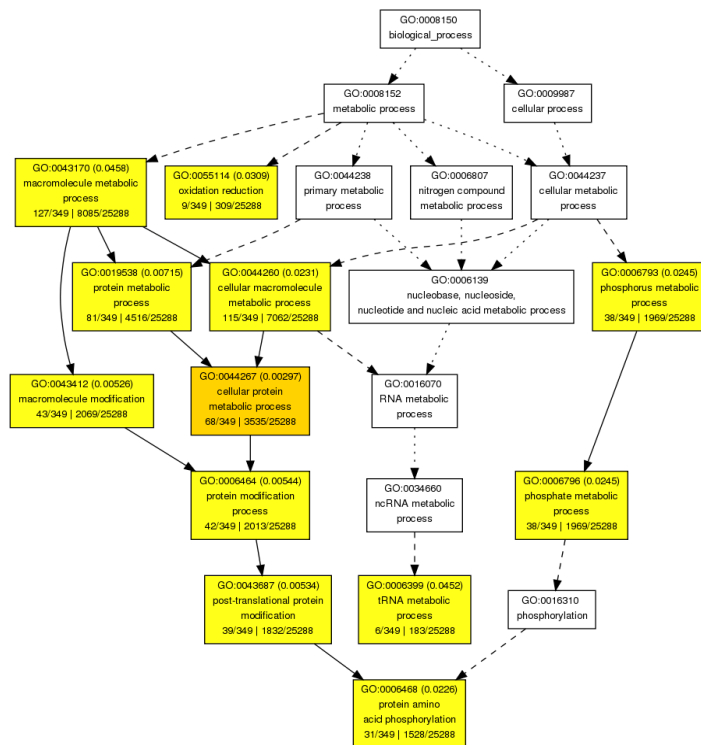


Supplemental figure 7. Frequency of INV4m according to accessions' adaptation class

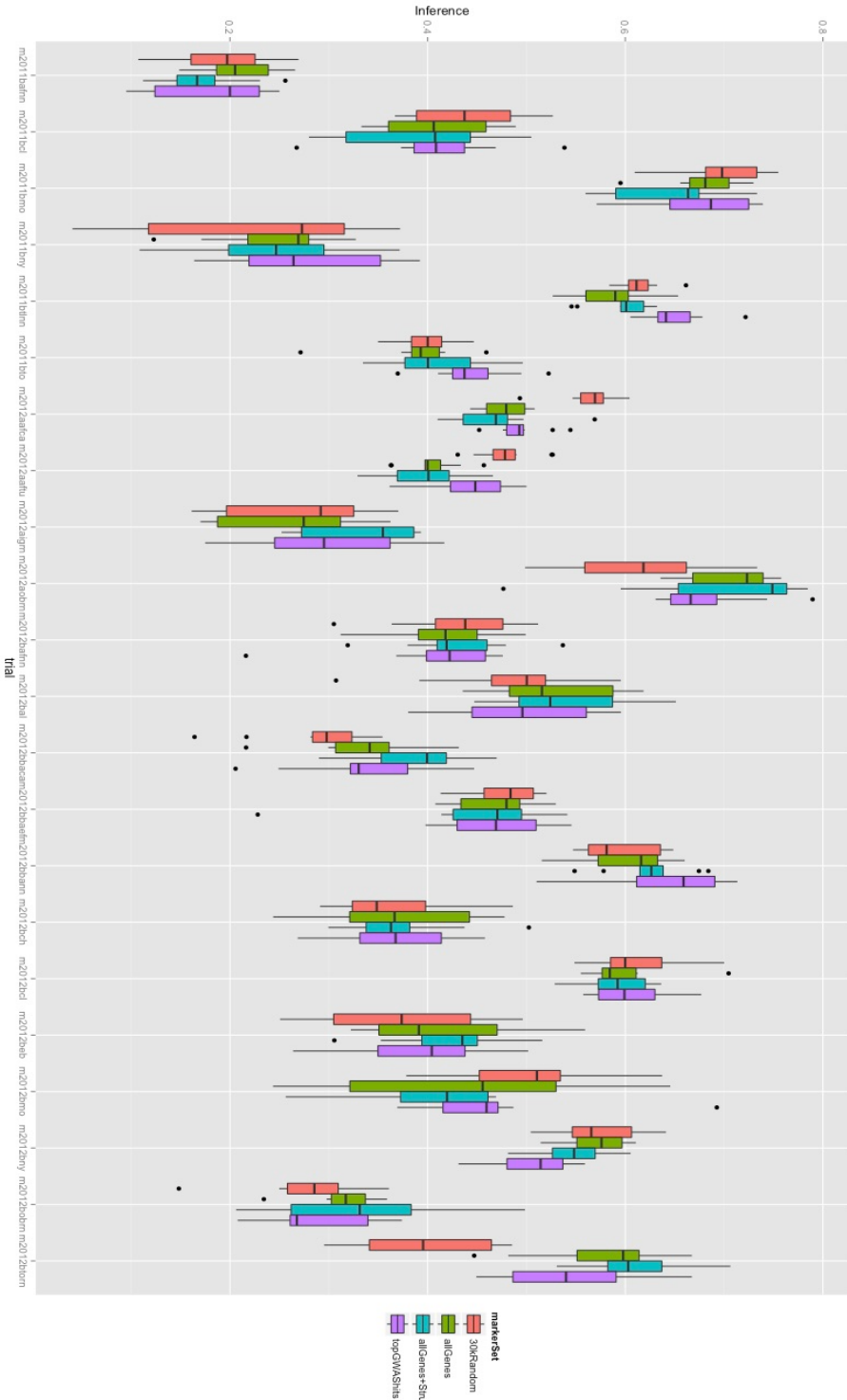
a)



b)



Supplemental figure 8. Ontology of genes for a) regulatory network genes and b) all associating gene



Supplemental figure 9. Genomic Prediction accuracy by trial

Year/Cycle	TrialCodeName	Location	Adaptation	Accessions
2011B	m2011BAFn	Agua Fria, Puebla	LOWLAND	1707
2011B	m2011BCL	Celaya, Guanajuato	SUBTROPICAL	783
2011B	m2011BMO	Tarimbaro, Michoacan	SUBTROPICAL	481
2011B	m2011BNY	San Pedro Lagunillas, Nayarit	SUBTROPICAL	551
2011B	m2011BTLn	Tlaltizapan, Morelos	SUBTROPICAL	1140
2011B	m2011BT0	Torreón, Coahuila	SUBTROPICAL	1403
2012A	m2012AAFca	Agua Fria, Puebla	LOWLAND	1921
2012A	m2012AAFTu	Agua Fria, Puebla	LOWLAND	1923
2012A	m2012AIGrn	Iguala, Guerrero	LOWLAND	749
2012A	m2012AOBrn	Obregon, Sonora	LOWLAND	452
2012B	m2012BAFn	Agua Fria, Puebla	LOWLAND	717
2012B	m2012BAL	Amoloya de Juarez, Mexico	HIGHLAND	428
2012B	m2012BBAca	El Batán, Mexico	HIGHLAND	817
2012B	m2012BBAef	El Batán, Mexico	HIGHLAND	759
2012B	m2012BBAnn	El Batán, Mexico	HIGHLAND	817
2012B	m2012BCH	Guadalupe-Victoria, Chiapas	LOWLAND	671
2012B	m2012BCL	Celaya, Guanajuato	SUBTROPICAL	805
2012B	m2012BEB	m2012BEB	SUBTROPICAL	658
2012B	m2012BMO	Numaran, Michoacan	SUBTROPICAL	282
2012B	m2012BNY	San Pedro Lagunillas, Nayarit	SUBTROPICAL	805
2012B	m2012BOBrn	Obregon, Sonora	LOWLAND	523
2012B	m2012BTOrn	Torreón, Coahuila	SUBTROPICAL	338

Supplemental Table 1. Trials location, adaptation class, and number of accessions planted