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The rate and effect of *de novo* mutations in a colonizing lineage of *Arabidopsis thaliana*

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32 Because colonizations and invasions are often associated with genetic bottlenecks, they offer an 33 opportunity to directly observe de novo mutations and their subsequent fate. North America has 34 recently been colonized by Arabidopsis thaliana, and many of the individuals found today belong to a 35 single lineage, HPG1. To determine substitution rates under natural conditions in this lineage, we have 36 sequenced 100 HPG1 genomes from plants collected between 1863 and 2006. We infer that the last 37 common HPG1 ancestor lived in the early 17th century, most likely the time when HPG1 began to 38 colonize N. America. Demographic reconstructions infer substantial population size fluctuations during 39 the past four centuries. Even though changing demographics can undermine the effect of natural 40 selection, we observed that mutations at coding sites were at lower frequency than mutations at 41 other sites, consistent with the effect of purifying selection. Exceptionally, some mutations rose to 42 high frequency and some had measurable effects in root development, consistent with positive 43 selection acting over mutations with an adaptive value. Our work showcases how by applying 44 genomics methods to a combination of modern and historic samples we can learn about plant 45 colonisations and invasions and observe "evolution in action".

46 Knowledge of mutation rates and efficacy of selection, which together determine the substitutions that can be observed in a population, is essential for understanding evolution¹. Mutation rates are required 47 to estimate the effective diversity of populations², to date historic population splits³, and to predict what 48 opportunities there are for rapid novel adaptations, for instance, to drugs or pesticides^{4,5}. Two extreme 49 approaches to discover these parameters are either short-term mutation accumulation experiments in 50 the laboratory⁶ and pedigree-based estimates⁷, or interspecific phylogenomic comparisons over millions 51 of years⁸. Since mutation rates are generally studied independently of natural selection and population 52 53 dynamics contexts, the two approaches often yield estimates that do not coincide, which has generated a heated controversy^{9,10}. An alternative not exploited so far is the analysis of naturally occurring 54 55 "evolutionary experiments" such as colonizing or invasive populations associated with a recent and strong genetic bottleneck^{11,12}. Both colonizations and invasions are increasingly common due to human 56 movement^{13,14}. The study of these natural experiments is especially powerful when time-stamped 57 samples from historic specimens can be used as internal calibration points^{15,16}. Colonizing populations 58 59 often start with very few individuals and therefore have low genetic diversity. The N. American 60 population of the self-fertilizing plant Arabidopsis thaliana is no exception, with many individuals 61 belonging to a genetically very similar lineage, haplogroup-1 (HPG1), that spread over large geographic 62 areas¹⁷. The study of the origin along with demographic and selective dynamics of mutations in HPG1 63 can further help us solve the current paradox generated by two evolutionary conjectures: "Baker's law", 64 inspired by the observation that selfing species are more successfully colonizing new environments than 65 outcrossing ones, and "Muller's ratchet", which posits that selfing populations are evolutionary dead ends, since low diversity and accumulation of more deleterious mutations should hinder their ability to 66 adapt to new environments 18-21. 67

To better understand the evolution of HPG1, we sequenced 27 herbarium specimens, from 1863–1993, and 76 live isolates of this lineage collected 1993–2006 (Fig. 1; Table S1). DNA retrieved from herbarium specimens showed biochemical features typical of ancient DNA²², which indicates that the DNA recovered from historic samples is authentic (Fig. S1, see Supplementary Online Material [SOM] for details). We mapped reads against an HPG1 pseudo-reference genome²³, focusing on single nucleotide

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polymorphisms (SNPs) because the short sequence reads of herbarium samples preclude accurate calling
 of structural variants. Genome sequences were of high quality, with herbarium samples covering
 96.8–107.2 Mb of the 119 Mb reference, and modern samples covering 108.0–108.3 Mb (Table S1). Pairs
 of herbarium genomes differed on average by 109-222 SNPs, and pairs of modern genomes by 186-299
 SNPs, that is, they ranged from 99.9997 to 99.9999 % identity .

78 A neighbor joining tree (Fig. 2A), multi-dimensional scaling (MDS) (Fig. 2B), and a parsimony 79 network (Fig. 2C) confirmed very close relatedness of the HPG1 genomes, with only three apparent 80 intra-HPG1 recombinants (Fig. 2C). Removing these resolved the reticulations in the parsimony network 81 (Fig. 2D). The remaining 100 samples (Table S1) constitute a quasi-clonal lineage mostly devoid of 82 effective recombination and population structure, and without SNPs in organellar genomes. The very low 83 genome-wide nuclear diversity (π = 0.000002, θ_W = 0.00001, 5,013 segregating sites) is two orders of magnitude lower than in the native range of the species ($\theta_W = 0.007$) (ref. ²⁴) (Table S1). The enrichment 84 of low frequency variants in the site frequency spectrum (Tajima's D = -2.84; global= -2.04) (ref. ²⁴) and 85 86 low levels of polymorphism are consistent with a recent bottleneck followed by population expansion 87 (Fig. 3). The obvious explanation is that the bottleneck corresponds to a colonization founder event, 88 likely by very few closely related individuals, or perhaps only a single plant. To describe intra-HPG1 89 relationships in a more sophisticated manner, we used Bayesian phylogenetic inference, exploiting collection dates for tip calibration of phylogenetic trees. The 76 modern individuals formed a largely 90 91 monophyletic clade, with only four interspersed herbarium samples from the second half of the 20th 92 century (Fig. 3A, B). Long branches reflected an abundance of singletons, typical of expanding 93 populations after bottlenecks.

94 To estimate the substitution rate in the HPG1 lineage, we used distance- and phylogeny-based 95 methods that take advantage of the known collection dates. One has to distinguish between the 96 mutation rate, which is the rate at which genomes change due to DNA damage, faulty repair, gene 97 conversion and replication errors, and substitution rate, which is the rate at which mutations survive and accumulate after demographic and natural selective processes²⁵. Under neutral evolution, mutation and 98 substitution rates should be equal²⁶. The simple evolutionary history of the HPG1 natural population 99 100 enables direct estimates of substitution rates, and by comparing these with mutation rates calculated in 101 controlled conditions, we can learn about demographic and selective forces. In the distance method, the 102 substitution rate is calculated from correlation between differences in collection time in historic-modern 103 sample pairs, and the number of changes between those pairs relative to a reference (Fig. 3C), scaled to 104 the size of the genome accessible to Illumina sequencing. This method resulted in an estimated rate of 2.11*10⁻⁹ substitutions site⁻¹ year⁻¹ (95% bootstrap Confidence Interval [CI]: 1.88–2.33*10⁻⁹) using 105 106 rigorous SNP calling quality thresholds. Relaxing the quality thresholds for base calling and minimum 107 genotyped rate affects both the number of called SNPs and the length of the interrogated reference sequence²⁷. These largely cancelled each other out, and our estimates were relatively stable, between 108 $2.1-3.2*10^{-9}$ substitutions site⁻¹ year⁻¹ (Table S3). The Bayesian phylogenetic approach, which uses the 109 110 collection years for tip calibration and assumes a relaxed molecular clock, yielded a similar estimate, 4.0*10⁻⁹, with confidence ranges overlapping the above estimates (95% Highest Posterior Probability 111 Density [HPPD]: $3.2-4.7*10^{-9}$). Based on the results obtained with different methods, we can confidently 112 say that the substitution rate in the wild should be between 2 to 5 $*10^{-9}$ site⁻¹ year⁻¹. We recommend 113

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114 that these rates be used to date temporal splits between populations. To be able to compare our 115 substitution rate with the mutation rate, both need to be expressed per generation. While A. thaliana is 116 an annual plant, seed bank dynamics generate a delay of average generation time at the population scale. A comprehensive study of multiple A. thaliana populations reports an average generation time of 117 1.3 vears²⁸, with a notable variance across populations. Re-scaling with the mean generation time led to 118 an adjusted substitution rate of $2.7*10^{-9}$ substitutions site⁻¹ generation⁻¹ (95% Cl 2.4-3.0*10⁻⁹) (Fig. 3E). 119 This is much lower than the rate of $7.1*10^{-9}$ mutations site⁻¹ generation⁻¹ (95% Cl 6.3–7.9*10⁻⁹) (Tables 120 121 S2, S3) that one can calculate from resequencing data for mutation accumulation (MA) lines in the Col-0 reference background grown in the greenhouse²⁹. 122

123 Differences in "per generation" rates could be caused by several factors, such as an imperfect 124 knowledge of the generation time in the wild (for rates using different generation times, see Fig. 3E). In 125 addition, mutagenic environmental factors, genome background, mutation spectrum, or methodological 126 idiosyncrasies can affect the estimates. For example, transposons, which comprise ~8% of the genome 127 and ~19% of the SNPs in greenhouse MA lines, had fewer SNPs called than expected in HPG1 (~13%). 128 This is likely due to difficulties when mapping reads to genomic areas with extensive structural variation^{30,30}a³⁰d³⁰ could have contributed to the lower substitution rate estimates for HPG1 (Fig. 3E. 129 Fig S3C. Table S3). In addition, the substitution spectrum in HPG1 is shifted to a lower 130 131 transition/transversion ratio compared to the MA lines (Fig. 3C and Fig. S3), which could be caused by 132 methylated cytosines (see Fig. S4 and SOM). Finally, an alternative evolutionary explanation for the rate 133 differences is that purifying selection slows the accumulation of mutations in the wild by removing 134 deleterious mutations (Fig. 3E). To find evidence of negative selection independently of dataset 135 comparisons, we looked at the site frequency spectrum of different annotations within the HPG1 dataset. Medium-frequency variants, which are more exposed to purifying selection³¹, were more 136 137 sharply depleted in genomic regions expected to be under greater selection constraint (genic and 138 nonsynonymous sites) than in putatively more neutral ones (intergenic or synonymous sites) (Fisher's 139 Exact test, p-value <0.05 for both comparisons, see Fig. S5). Therefore, even if we cannot say with 140 certainty that purifying selection drives the differences between HPG1 and MA rates, it must be 141 responsible for the differences between different types of sites within HPG1.

142 The substitution rate allows dating of HPG1's origin. The mean estimate from Bayesian methods 143 was the year 1597 (HPPD 95%: 1519-1660) (Fig. 3A, B). We also used a non-phylogenetic method that 144 utilizes the relationship between the average genetic distance between any two individuals, with the 145 substitution rate multiplied by twice the divergence time and the genome size; solving by the divergence 146 time in the equation we obtained 353 years. When subtracted from the average collection date of our 147 samples, the corresponding point estimate is 1625, within the confidence interval of the Bayesian 148 estimate. This corresponds to the date of the last common ancestor of HPG1, and should thus be close to 149 the time of introduction of HPG1 to N. America. (The date is older than our previous estimate, for which we had naively applied the higher greenhouse mutation rate²³). Inference of N_{e} through time suggested 150 exponential population growth until the early 19th century (Fig 3B, Fig S6C). During the 20th century the 151 N_{ρ} trajectory showed oscillating patterns between growth and bottlenecks, which are typical of selfing 152 organisms ³², and which likely led to a replacement of most HPG1 sublineages, as the modern samples 153 154 are all very closely related (Fig. 3 A, B).

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155 Since we knew both the collection years and locations of origin of the HPG1 samples, we could 156 also analyze the migration dynamics of HPG1. Although unknown sources of sampling bias could affect our analyses¹⁶, the phylogeographic models suggested that HPG1 came to cover much of its modern 157 158 range soon after its introduction to N. America (Fig. S6 A,B). We found a significant correlation between 159 collection date and both latitude and longitude (Fig. 1C), which we interpret as a net, highly dispersed, 160 movement in a northwestern direction over time. Additional support for this hypothesis comes from an 161 isolation-by-distance signal, which is most consistent with a historic westward migration and a more 162 recent reverse eastward migration (Fig. S6 E,F). The apparent source of those new migrants now 163 persisting along the East coast was the Lake Michigan area.

164 Finally, while we did not expect to easily find mutations that have helped the HPG1 lineage to 165 adapt to its new N. American environment, we wanted to determine whether any of the mutations have 166 measurable phenotypic effects. Focusing on flowering-related, reproductive and root traits of likely 167 ecological relevance, we detected significant quantitative heritable variation (Table S4). We used an 168 approach borrowed from GWAS to find SNPs that had increased in frequency (>5%) and were associated 169 with these phenotypes. Because conventional GWAS relies on recombination, which is almost absent 170 from our population, our approach could not identify individual SNPs, but only SNP cohorts distributed 171 across the genome³³. We found 79 SNPs associated with root traits, of which nine resulted in 172 nonsynonymous changes. We did not find any SNPs associated with flowering time, even though it is thought to be a key player of rapid adaptation in many annual species^{34,35}. Nineteen other SNPs, of 173 174 which four were nonsynonymous, were associated with climate variables (www.worldclim.org) even 175 after correction for latitude and longitude, and some of the hits overlapped between root traits and 176 climate variables (Table 1, Table S5, Fig. S7). Although a good number of SNPs was associated with 177 phenotypes and/or climate variables, it is not possible to confidently pinpoint individual candidate SNPs, 178 since the extent of whole-genome linkage disequilibrium (LD) in HPG1 is high (Fig. S9 B,C). However, 179 there is a gradient in the extent of LD between SNPs associated with root architecture, which could help 180 to determine particularly promising candidates for molecular characterization and quantification of 181 fitness in natural conditions (Table 1, Fig. S9 D-F). For example, the gene AT5G19330, overexpression of 182 which confers salt tolerance³⁶, contains a SNP that was unlinked to other hits and that has risen in the 183 last four centuries to a frequency of 40%. This SNP is likely to change protein function, as it leads to a 184 substitution of cysteine for tryptophan. Another nonsynonymous SNP is located in AT2G38910, encoding 185 a calcium dependent kinase that belongs to a family of factors involved in root hydraulic conductivity and phytohormone response^{37,38}. Remarkably, most derived root-associated alleles, when compared with 186 187 equally frequent neutral alleles, were first seen in older herbarium samples, most of which were 188 collected near Lake Michigan, the apparent source of modern populations (Fig. S8). Altogether, the rise 189 in frequency, older age of some de novo mutations, and their quantifiable phenotypic effects and 190 climatic correlations strengthen the hypothesis that they might have an adaptive value and were under 191 positive selection. These results favor Baker's law, which alleges that selfing species can often adapt to 192 new environments, over the evolutionary dead-end hypothesis of Muller's ratchet. Furthermore, these 193 suggestive signals of rapid adaptation via de novo mutations could change the current paradigm that 194 invasive species adapt most often from sources of standing variation, either because an incomplete

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bottleneck left residual variation or because there has been subsequent admixture with native species or
 secondary colonizers^{39,40}.

197 In summary, we have exploited whole-genome information from historic and contemporary 198 collections of a herbaceous plant to empirically characterize the effect of evolutionary forces during a 199 recent colonization. With this natural time series experiment, we could directly estimate the nuclear 200 substitution rate in wild A. thaliana populations. This parameter, which provides immediate ability to date key events of populations, is only known for one other species: Homo sapiens ⁴¹. We have 201 202 presented evidence that purifying selection is perceptible already over time scales spanning only a few 203 centuries. Although the colonizing population we have investigated has limited diversity and suffered 204 rapid fluctuations in population size, there appear to be *de novo* mutations with phenotypic effects that 205 contributed to rapid adaptation. While A. thaliana HPG1 is not an invasive species, it can teach us about 206 fundamental evolutionary processes behind successful colonizations and adaptation to new 207 environments. Our work should encourage others to search for similar natural experiments and to unlock 208 the potential of herbarium specimens to study "evolution in action".

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Online Content Methods, along with any additional Extended Data display items, are available in the online version
 of the paper; references unique to these sections appear only in the online paper.

- 212
- 213 **Supplementary Information** is available in the online version of the paper.
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222 Author Contributions H.A.B. and D.W. conceived and supervised the project, and coordinated the collaborative 223 effort. J.B. coordinated the collection of modern seed samples. C.J., B.B. and J.B. performed and analyzed flowering 224 time and seed set greenhouse experiments. C.S. and R.S. performed and analyzed root assays and seed size 225 measurements under the supervision of W.B.; C.B. and J.H. sequenced and curated modern samples, coordinated 226 by D.W.; H.A.B. coordinated the collection and analysis of herbarium samples. J.K. coordinated the extraction of 227 DNA and library preparation of herbarium samples. V.J.S. and E.R. prepared sequencing libraries from herbarium 228 specimens. C.B. called variants in HPG1. J.H. called variants in mutation accumulation lines. M.E.A. performed the 229 population and quantitative genomic analyses with supervision of R.N., C.B. and H.A.B. The paper was written by 230 M.E.A., C.B., H.A.B. and D.W. with comments from all coauthors.

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METHODS

315 Sample collection and DNA sequencing. Modern A. thaliana accessions were from the collection described by Platt and colleagues¹⁷ which identified HPG1 candidates based on 149 genome-wide SNPs 316 317 (Table S1). Herbarium specimens were directly sampled by Max Planck colleagues Jane Devos and 318 Gautam Shirsekar, or sent to us by collection curators from various herbaria (Table S1). DNA from herbarium specimens was extracted as described⁴² in a clean room facility at the University of Tübingen. 319 Two sequencing libraries with sample-specific barcodes were prepared following established protocols, 320 321 with and without repair of deaminated sites using uracil-DNA glycosylase and endonuclease VIII (refs. ^{43–45}) (see Supplementary Online Material [SOM]). DNA from modern individuals was extracted from 322 323 pools of eight siblings using the DNeasy plant mini kit (Qiagen, Hilgendorf, Germany). Genomic DNA 324 libraries were prepared using the TruSeq DNA Sample or TruSeq Nano DNA sample prep kits (Illumina, 325 San Diego, CA), and sequenced on Illumina HiSeq 2000, HiSeq 2500 or MiSeq instruments. Paired-end reads from modern samples were trimmed and quality filtered before mapping using the SHORE pipeline 326 v0.9.0 (ref. ^{46,47}). Because ancient DNA fragments are short (Fig. S1B), we merged forward and reverse 327 reads for herbarium samples after trimming, requiring a minimum of 11 bp overlap⁴⁸, and treated the 328 resulting as single-end reads. Reads were mapped with GenomeMapper v0.4.5s (ref. 49) against an HPG1 329 pseudo-reference genome²³, and against the Col-0 reference genome, and SNPs were called with 330 SHORE^{23,50} using different thresholds. Samples JK2509 to JK2531 were only mapped to the HPG1 331 332 pseudo-reference genome. Average coverage depth, number of covered genome positions, and number 333 of SNPs identified per accession relative to HPG1 are reported in Table S1. We also re-sequenced the genomes of twelve Col-0 MA lines^{50,51} (Table S2). 334

Phylogenetic methods and genome-wide statistics. We used four methods to estimate the relationships among modern accessions, and between modern and herbarium HPG1 samples: (i) multidimensional scaling (MDS); (ii) construction of a neighbor joining tree with the adegenet package in R (ref. ⁵²), with branch support assessed with 1,000 bootstrap iterations and consensus reported; (iii) construction of a parsimony network using SplitsTree v.4.12.3 (ref. ⁵³), with confidence values calculated with 1,000 bootstrap iterations; (iv) performing a Bayesian phylogenetic analysis using BEAST v.1.8 (ref. ⁵⁴) (see below).

We estimated genetic diversity as Watterson's θ (ref. ⁵⁵) and nucleotide diversity π , and the 342 difference between these two statistics as Tajimas's D (ref. 56) using DnaSP v5 (ref. 57). We calculated the 343 344 folded site frequency spectrum (SFS) as well as the unfolded SFS, for which we assigned the ancestral state using the A. lyrata genome⁵⁸. We estimated pairwise linkage disequilibrium (LD) between all 345 possible combinations of informative sites, ignoring singletons, by computing r^2 , D and D' statistics. For 346 347 the modern individuals, we calculated the recombination parameter rho $(4N_{o}r)$ and performed the four-gamete-test⁵⁹ to identify the minimum number of recombination events. All LD and recombination 348 related statistics were determined using DnaSP v5 (ref. ⁵⁷) (see SOM). 349

Substitution and mutation rate analyses. We used genome-wide nuclear SNPs to calculate pairwise "net" genetic distances using the equation $D'_{ij} = D_{ic}-D_{jc}$, where D'_{ij} is the net distance between a modern sample *i* and a herbarium sample *j*; D_{ic} the distance between the modern sample *i* and the reference

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genome *c*; and D_{jc} is the distance between a modern sample (j) and the reference genome (c). We calculated a pairwise time distance in years between the collection times, *T*'ij, and calculated the linear regression: D' = a+bT'. The slope coefficient *b* describes the number of substitution changes per year. We used either all SNPs or subsets of SNPs at different annotations (genic, intergenic etc.) appropriately scaled by accessible genome length and with confidence intervals determined by bootstrap (see SOM and Fig. S3).

359 The second approach used Bayesian phylogenetics with the tip-calibration method implemented 360 in BEAST v1.8 (ref. ⁵⁴). Our analysis optimized simultaneously and in an iterative fashion using a Monte 361 Carlo Markov Chain (MCMC) a tree topology, branch length, substitution rate, and a demographic Skygrid 362 model (Fig. 3 A,B; see SOM). The demographic model is a Bayesian nonparametric one that is optimized 363 for multiple loci and that allows for complex demographic trajectories by estimating population sizes in 364 time bins across the tree based on the number of coalescent - branching - events per bin (ref. ⁶⁰). We also performed a second analysis run using a fixed prior for substitution rate of 3*10⁻⁹ substitutions 365 site⁻¹ year⁻¹ based on our previous net distance estimate to confirm that the MCMC had the same 366 367 parameter convergence, e.g. tree topology, as in the first "estimate-all-parameters" run.

368 Inference of genome-wide selection. We separately analyzed sequences at different annotations, since 369 certain regions regions should be under a different selection regime (less evolutionary constraint) than 370 others. We compared the means and confidence intervals of substitution rates in the entire genome and 371 in intergenic regions for both datasets, HPG1 population and laboratory Col-0 MA lines. Only within the 372 HPG1 population, we also tested for an interaction between low and common allele frequency 373 polymorphisms and putatively selected and putatively neutral annotations (comparisons: entire genome 374 - intergenic, genic - intergenic, nonsynonymous - synonymous). The formal test was Fisher exact test and 375 low and common frequency SNPs were defined in all possible cutoffs from 1 to 45% allele frequency (Fig. 376 S5). The signal captured is based on the assumption that purifying selection is more efficient at 377 intermediate frequencies, pushing deleterious variants towards lower frequency in the spectrum.

378 Association analyses and dating of new mutations. We collected flowering, seed and root morphology 379 phenotypes for 63 accessions. For associations with climate parameters, we followed a similar rationale described⁶¹. 380 as We extracted information from the bioclim database 381 (http://www.worldclim.org/bioclim) at a 2.5 degrees resolution raster and intersected it with geographic 382 locations of HPG1 samples (n = 100). We performed association analyses under several models and *p*-value corrections using the R package GeneABEL (ref. ⁶²), with phenotypes and climatic variables as 383 384 response variables and SNPs as explanatory variables; appropriately correcting for covariates. Resulting 385 p-values were adjusted with an empirical p-value distribution generated from 1,000 permuted datasets, 386 or with a double Bonferroni correction: 5% / (number of SNPs + number of phenotypes tested).

- Accession numbers. Short reads have been deposited in the European Nucleotide Archive under the
 accession number XXXXX.
- 389

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434 **FIGURE LEGENDS**

Figure 1. Geographic location and temporal distribution of HPG1 samples.

(A) Sampling location of herbarium (blue) and modern individuals (green). (B) Temporal distribution of
 samples (random vertical jitter for visualization purposes). Stars indicate four herbarium accessions that
 nest within the clade of modern accessions (see Fig. 3). (C) Linear regression of latitude and longitude as
 a function of collection year (p-value of the slope and Pearson correlation coefficient are indicated)

440 Figure 2. Relationship among herbarium and modern HPG1 samples.

(A) Neighbor joining tree, consensus of 1,000 bootstrap replicates. Branch lengths indicate number of
 base substitutions (colors represent herbarium (blue) and modern individuals (green)). Scale line shows
 the equivalent branch length of 80 nucleotide changes. Note that no outgroup was included. (B) First
 two dimensions of a multidimensional scaling plot based on pairwise identity-by-state distances. Fraction
 of variance explained given in parentheses. (C, D) Network of all samples using the parsimony splits
 algorithm, before (C) and after (D) removing intra-HPG1 recombinants (in red).

Figure 3. Substitution rates and demographic history.

448 (A) Bayesian phylogenetic analyses employing the tip calibration methodology. A total of 10,000 trees 449 were superimposed as transparent lines, and the most common topology was plotted solidly. Tree 450 branches were calibrated with their corresponding collection dates. (B) Maximum Clade Credibility 451 (MCC) tree summarizing the trees in (A). The demographic model underlying the phylogenetic analysis, 452 Bayesian Skygrid reconstruction, is superimposed; the mean N_e over time is shown as a dotted line and 453 the 95% HPD is shaded grey. Note the scale line shows the equivalent branch length of 50 nucleotide 454 changes. (C) Regression between pairwise net genetic and time distances. The slope of the linear 455 regression line corresponds to the genome substitution rate per year. (D) Substitution spectra in HPG1 456 samples, compared to greenhouse-grown mutation accumulation (MA) lines. (E) Comparison of 457 genome-wide, intergenic, intronic, and genic substitution rates in HPG1 and mutation rates in 458 greenhouse-grown MA lines. Substitution rates for HPG1 were re-scaled to a per generation basis 459 assuming different generation times. Confidence intervals in HPG1 substitution rates were obtained from 460 95% confidence intervals of the slope from 1,000 bootstrap (see Table S4 for actual values).

Table 1. Genic SNPs associated with different traits.

Most SNPs first appeared in sample JK2530 collected 1922 in Indiana. For non-synonymous SNPs, the amino acid change and the Grantham score (ranging from 0 to 215), which measures the physico-chemical properties of the amino acids, are reported. All SNPs in the table were significant (p < 0.05) after raw p-values were corrected by an empirical p-value distribution from a permutation procedure. * highlights those that also passed a double Bonferroni threshold, correcting by number of SNPs and number of phenotypes (p < 0.0001). LD corresponds to how many other SNP hits are in high

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468 linkage (r^2 >0.5). See Table S5 for information on all significant SNPs and Table S4 for details on 469 phenotypes and climatic variables.







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Table 1

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$Trait^{\dagger}$	Location (chr-bp)	Gene	Anno– tation	Protein	aa change	LD	Bonf.
G	1-958,948	AT1G03810	nonsyn	Oligonucleotide/ oligosaccharide binding	A>P, 27	53	
D	1-13,994,958	AT1G36933	transposon	Copia		49	
S	1-20,324,050	AT1G54440	intronic	RRP6-LIKE 1		11	*
D	1-23,648,407	AT1G63740	nonsyn	TIR-NLR family	Y>S, 144	46	
G	2-358,395	AT2G01820	syn	RLK family		43	*
G	2-585,918	AT2G02220	syn	PSKR1		42	*
G	2-6,034,545	AT2G14247	syn	Expressed protein		38	*
G	2-7,047,529	AT2G16270	nonsyn	Unknown protein	P>A, 27	37	*
G	2-7,186,220	AT2G16580	intronic	SAUR8		36	*
G	2-10,495,275	AT2G24680	intronic	B3 family		34	*
G	2-12,415,084	AT2G28900	intronic	OEP16		32	
S	2-16,039,488	AT2G38290	3' UTR	AMT2		8	*
S	2-16,247,290	AT2G38910	nonsyn	СРК20	A>G, 60	7	*
G	2-16,333,662	AT2G39160	nonsyn	Unknown protein	A>G, 60	29	
G	3-2,500,258	AT3G07830	syn	PGA3		28	*
G	3-3,629,794	AT3G11530	intronic	VPS55		26	*
G	3-4,269,626	AT3G13229	5' UTR	DUF868 domain		25	*
D	3-11,873,293	AT3G30219	transposon	Gypsy		0	
G & D	4-4,228,138	AT4G07440	transposon	Oligonucleotide/ oligosaccharide binding		19	
G & D	4-9,046,942	AT4G15960	nonsyn	Alpha/beta-hydrolase superfamily	A>Q, 24	18	
G & D	4-15,646,341	AT4G32410	syn	ANY1		15	
G	4-15,845,001	AT4G32840	3' UTR	PFK6		14	
D	5-4,245,213	AT5G13260	syn	Unknown protein		12	
D	5-4,500,202	AT5G13950	nonsyn	Unknown protein	A>G, 60	11	
G	5-4,797,923	AT5G14830	transposon	Retrotransposon		10	
G	5-6,508,329	AT5G19330	nonsyn	ARIA	C>W, 215	0	
G	5-11,090,365	AT5G29037	transposon	Gypsy		4	
G	5-12,312,975	AT5G32630	pseudogene	_		3	
G	5-12,358,159	AT5G32825	transposon	САСТА		2	
S	5-16,024,197	AT5G40020	intronic	Thaumatin superfamily		2	*

[†]Traits with significant associations were root gravitropism (G), root size (S), or summer precipitation, related to drought conditions.

1	bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this prepri- peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license Supplementary Information Guide for	nt (which was not
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50 **1.** Sample collection and preparation

51 Seeds from modern accessions (Table S1) were bulked at the University of Chicago. Progeny for DNA 52 extraction was grown at the Max Planck Institute for Developmental Biology. We used 2 to 8 mm² of 53 dried tissue for destructive sampling from the herbarium specimens (Table S1).

54

49

55 **2.** Authenticity of aDNA

56 First, unrepaired sequencing herbarium libraries were screened for authenticity by sequencing at low 57 coverage on Illumina HiSeq 2500 or MiSeq instruments. To verify the DNA retrieved from historical 58 samples of A. thaliana was authentic, we checked the percentage of endogenous DNA of the sample 59 (Fig. S1A) as well as typical postmortem DNA damages: high fragmentation of DNA (Fig. S1B), 60 enrichment of substitution from C to T at the first base pair (Fig. S1C) as well as purine enrichment at 61 breakpoints of DNA fragments (Fig. S1D) (for details see ¹). Sequencing to produce the final genomes 62 (101 bp paired end) was carried out on an Illumina HiSeq 2000 instrument after DNA repair by uracil-DNA glycosylase $^{2-4}$. 63

64

65 **3. SNP calling thresholds**

66 To asses the effect of SNP calling thresholds on the mutation rate, we employed three different SHORE v0.9.0 quality thresholds following previous work (see Table S4 from ref. ⁵): allowing at most 67 68 one intermediate penalty in all strains (most stringent threshold; "32-32"); requesting that at least 69 one strain had at most one intermediate penalty, while all others were allowed up to two high and 70 one intermediate penalties (intermediate stringency, "32-15"); and finally allowing one high and one 71 intermediate penalty for all strains (most lenient stringency, "24-24"). On top of that, we would 72 either allow missing information per SNP in up to 50% of accessions, or request complete 73 information (0% missing rate). Thus, the most rigorous case would be 32-32 quality and 0% missing 74 rate, and the most relaxed 24-24 quality and 50% maximum missing rate. Substitution rate 75 calculations (section 7.2) were done for datasets from all combinations of these quality parameters 76 (Fig. S3), and we chose the regular 32_15 quality threshold and complete information for the final 77 estimate (Fig 3 C, E).

78

79 4. Resequencing of Col-0 Mutation Accumulation lines

We also sequenced the genomes of twelve greenhouse-grown mutation accumulation (MA) lines, including ten that had been sequenced at lower coverage before^{5,6} (Table S2). We called SNPs, indels and structural variants (SVs), following the workflow and parameters described⁷, but without iterations. This procedure resulted in 2,203 polymorphisms shared by all lines, indicating errors in the

bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license. 84 reference sequence (12% of variants replaced N's in the TAIR9 genome) or genetic differences in the 85 founder plant of the MA population compared to the Col-0 reference genome. In addition, we 86 identified 388 segregating variants across the twelve lines (Table S2), of which 350 were singletons. 87 This analysis revealed on average 25.5 SNPs, 4.9 deletions and 3.2 insertions per MA line at the 31st 88 generation (Table S2), compared to 19.6 SNPs, 2.4 deletions and 1.0 insertions previously detected in the 30th generation with shorter read length and lower read depth⁸. The genome length accessed in 89 90 this sequencing effort, 115,954,227 bp, was used to scale the number of point mutations to a rate of 7.1×10^{-9} mutations site⁻¹ generation⁻¹ (Table S3, Fig. 3E). 91

92

93 5. Identification of *bona fide* HPG1 accessions and mutations

Before we could work with the colonizer group HPG1, we needed to carefully identify individuals that belong to other haplogroups or that have introgressions from them. We established the relationships among all samples at three levels of resolution: (i) the 149 nuclear SNPs used originally to define the HPG1 haplogroup in a global screening⁹ (Fig. S2A), (ii) SNPs in the chloroplast genome (where we did not find any variants within HPG1), (iii) and all nuclear genome SNPs (Fig. S2B-C). At these three levels we performed a multidimensional scaling (MDS) analysis and built a neighbor-joining tree using the adegenet package in R (ref. ¹⁰).

Having identified these *bona fide* HPG1 individuals, we wanted to confirm that the diversity has a legitimate origin from *de novo* mutations. For that we used the 1001 Genomes resource (1001genomes.org) to verify that the majority of HPG1-specific variants did not originate in the native Eurasian range. Subsetting the genomes from this resource to only European accessions, and limiting the SNP set to those with \geq 1% frequency of alternative alleles, there were 338 variants out of all 5,181 HPG1 variants that were also found in Europe or Asia (6%). Only one of the reported SNPs associated with phenotypes (see section 10) was among these shared variants.

108 There are several scenarios that can explain these shared SNPs. One is that some HPG1 109 individuals were moved back to Europe by humans. Another one is that parallel mutations occurred 110 in North America and Eurasia or that a reversion-mutation happened in some HPG1 individuals. 111 Given that 10% of all sites in the genome are variable in the 1001 Genomes collection, this is not an 112 implausible scenario for at least a fraction of shared SNPs. Two additional scenarios involve an origin 113 from standing European variation: (1) the shared variants come from small introgression events that 114 passed our filters above, or (2) the bottleneck was not complete, and while it left no diversity in the 115 chloroplast genome, a few hundred SNPs were passed on in the nuclear genome (given the low 116 number of variants, the colonizers could have been as many as two dozen seeds)

117 118

6. Extent of linkage disequilibrium and recombination

119 We estimated pairwise linkage disequilibrium (LD) between all possible combinations of informative 120 sites, ignoring singletons, by computing r^2 , D and D' statistics. LD decay was estimated using a linear 121 regression approach. Linkage disequilibrium parameter D' did not decay with physical distance 122 (intercept = 0.99, slope = 0.00, p < 0.0001) among all SNP pairs. Furthermore, only 0.02% of the 123 nonsingleton SNP pairs were not in complete linkage disequilibrium (D'<1), indicating extensive 124 linkage between chromosomes. We also formally estimated recombination within HPG1. The estimate was much lower $(4N_{\rho}r = \rho = 3.0 \times 10^{-6} \text{ cM bp}^{-1})$ than for a similar-sized collection of diverse 125 A. thaliana individuals from the native range¹¹ ($\rho = 7.5 \times 10^{-2}$ cM bp⁻¹). The four-gamete test¹², 126 127 which determines whether all four possible gametes (ab, aB, Ab, AB) are observed for two 128 segregating loci, revealed that all configurations of SNPs could be explained with as few as 38 129 recombination events for the 100 genomes. We argue that this number of potential recombination 130 events is sufficiently small to use phylogenetic methods with the 100 HPG1 genomes, even though 131 such methods are normally not appropriate for genome-wide analyses. Indeed, other sources of 132 failure of the four-gamete test and the violation of phylogenetic assumptions could be sequencing 133 errors, or lineage sorting of segregating sites from the ancestral population. LD and recombination 134 related statistics were determined using DnaSP v5 (ref. 13) or plink v1.90b2n (ref. 14).

135

136 **7. Substitution and mutation rate analyses**

137 <u>7.1 Greenhouse grown MA lines</u>

Mutation rates were estimated for each 31st generation greenhouse-grown MA line⁵ as the number of mutations divided by the total bp length of the genome (or a given annotation) and by 31 generations (the two MA lines with only three generations were excluded from this analysis). Mean and confidence intervals across lines are reported (Table S3). The genome length was determined as all base pairs with coverage higher or equal to 3, and a SHORE mapping quality score of at least 32 in one sample (Table S2).

144

145 <u>7.2 Natural populations of HPG1</u>

146 7.2.1 Net distances

For the "net genetic distances" method, we computed confidence intervals of the *b* regression slope coefficient (D' = a+bT') using a bootstrap with replacement of 1,000 samples to avoid over-confident confidence intervals due to lack of independence of points¹⁵. We used either all SNPs or SNPs at specific annotations to calculate different substitution rates and scaled the slope into a per-base rate using all positions (of the given annotation) that passed alternative or reference call quality

bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license. 152 thresholds rather than using a single value of genome length (Table S3). For all annotations we 153 calculated substitution rates with three quality thresholds and either full information per SNP or 154 allowing a maximum of 50% missing accessions per SNP (see section 3 and Fig. S3C). For some 155 annotations substitution rates were not reliable. For instance, in 3' and 5' UTR regions, we did not 156 have enough mutations (on average ~1 SNP difference between any pair), and thus do not report 157 these regions' rates. Transposons showed unstable substitution rate estimates that we attribute to 158 structural variation relative to the reference genome. This likely decreased our ability to map transposon reads correctly and subsequently call SNPs¹⁶. In contrast, in the MA dataset, transposon 159 160 structural variation was probably fairly low since only 31 generation separate the Col-0 reference 161 genome with each of the ten derived MA lines. This can be the reason that the number of 162 transposon SNPs identified in the MA dataset is proportionally larger than in HPG1 (Table S2 and S3). 163 Therefore, transposon substitution rates in HPG1 cannot be trusted.

164

165 7.2.2 Bayesian tip-calibration

- 166 For the second approach to estimate a substitution rate, the Bayesian phylogenetics tip-calibration 167 approach, we performed systematic runs and chain convergence assessments of different demographic and molecular clock models. We found the Skygrid demographic model¹⁷ and the 168 lognormal relaxed molecular clock¹⁸ ¹⁸ the most appropriate models. Under a relaxed molecular 169 170 clock, the substitution rate is allowed to vary across branches with a lognormal distribution. The prior used for molecular clock was a Continuous-Time Markov Chain (CTMC)^{17,19}. The analysis was carried 171 172 out remotely at CIPRES PORTAL (v3.1 www.phylo.org) using uninformative priors. The run took about 173 1,344 CPU hours and performed 1,000 million steps in a Monte Carlo Markov Chain (MCMC), 174 sampling every 100,000 steps. Burn-in was adjusted to 10% of the steps. To visualize the tree output 175 we produced a Maximum Clade Credibility (MCC) tree with a minimum posterior probability 176 threshold of 0.8 and a 10% burn-in using TreeAnnotator (part of BEAST package), and visualized the 177 MCC tree using FigTree (tree.bio.ed.ac.uk/software/figtree/) (Fig. 3B). Additionally, we used DensiTree²⁰ to simultaneously draw the 10,000 BEAST trees with the highest posterior probability 178 179 (Fig. 3A). Since all trees were drawn transparently, agreements in both topology and branch lengths 180 appear as densely colored regions, while areas with little agreement appear lighter.
- 181

182 7.2.3 Methylation status of mutated sites

As in many other species, the spectrum of *de novo* mutations in the greenhouse-grown *A. thaliana* MA lines is biased towards G:C \rightarrow A:T transitions⁸, leading to an inflated transition-to-transversion ratio (Ts/Tv). This bias is less pronounced in recent mutations in a Eurasian collection of natural accessions²¹ and in HPG1 accessions (Fig. 3D). A recent multigenerational salt stress experiment in bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license.
the greenhouse also showed a more balanced Ts/Tv (ref. ²²). These findings indicate that less benign conditions might promote a lower Ts/Tv, and one possible cause are methylation patterns, known to change under different environments²³.

190 We interrogated the potential evolutionary role of cytosine methylation in the mutability of 191 cytosine bases in the HPG1 accessions. For reference DNA methylation data, we used previously generated bisulfite-sequencing data of HPG1 strains⁷ and of Col-0 MA lines⁵, respectively. For both 192 193 datasets, methylation status was calculated as the fraction of reads with methylated cytosines by the 194 total number of reads at a certain cytosine position in the genome. Our rationale was that if 195 methylation affected mutability, the degree of methylation at positions were we find a new mutation 196 should be higher. To be sure that a given site in HPG1 was a new mutation, we only considered positions for which we could determine that state by alignment to the A. lyrata genome²⁴. The 197 198 "tested sites" were positions in HPG1 that had a mutation both from A. lyrata and A. thaliana Col-0. 199 These positions can be of two kinds, "fixed" if all HPG1 individuals carry the alternative, or 200 "segregating" if both reference and alternative alleles exist in HPG1. As control, "control set", we 201 used cytosine positions that did not vary across HPG1, A. lyrata and A. thaliana. To produce the 202 methylation distribution of the control set we randomly chose 1,000 invariant cytosine positions. For 203 the test sets, we averaged the methylation degree and compared it with the control distribution.

204 Ancestral cytosines with higher methylation in both A. thaliana Col-0 reference and HPG1 205 pseudo-reference methylome datasets were more likely to mutate to thymines in HPG1 (Fig. S4 A-D). 206 Additionally, the methylation degree at substitutions inside genes was higher in the HPG1 207 methylome (Fig. S4 B,D). While some $C \rightarrow T$ changes could be explained by higher spontaneous 208 deaminations known to happen more often at methylated cytosines, also $C \rightarrow A/G$ substitutions were 209 more likely to have been methylated. If this process is common enough, the Ts/Tv ratio should 210 decrease. We are far from understanding differences in Ts/Tv in natural and controlled conditions, 211 but definitely methylation status seems to have a strong statistical connection with mutability.

212

213 8. Inference of genome-wide selection

214 Since we observed differences between the two mutation accumulation (MA) datasets (the 215 laboratory Col-0 MA lines and the wild HPG1 lines), we tried to infer selection based on differences in 216 polymorphisms and substitution rates. We compared the different substitution rates and the 95% 217 bootstrap confidence intervals to assess how identical they were (Table S3). Genome-wide 218 substitution rate in the HPG1 dataset was significantly lower than that in controlled greenhouse conditions, even after correcting by the mean generation time of 1.3 years²⁵ (Table S3). However, 219 220 these differences could be due to differences in individual genomic annotations. For instance, coding 221 regions and introns were virtually identical between the two datasets, but transposons and

bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license. 222 intergenic were much lower in HPG1. Furthermore, if the true generation time is about 2-3 years (Fig. 223 3), differences in the mentioned annotations would also disappear. Therefore, we tried to investigate 224 the existence of purifying selection based on frequency equilibriums only within the HPG1 225 population. We did comparisons between pairs of annotation categories and between pairs of 226 frequency classes (i.e., low and common). In this way, genome-wide, genic and non-synonymous 227 polymorphisms were compared to matched putatively neutral annotations: intergenic and 228 synonymous sites (Fig. S5). We tested for an interaction using a Fisher's Exact test on 2x2 table 229 counts as:

230

Neutral annotation & low frequency	Neutral annotation & common frequency
Selected annotation & low frequency	Selected annotation & common frequency

235

Because any frequency cut-off is arbitrary, we computed the test with all cutoffs from 0 to 50% frequencies in 1% steps (Fig. S5). This test, which resembles in concept the MK or HKA tests, evidences that there is a depletion of common frequency polymorphisms, which are more exposed to selection, at the three putatively selected genomic levels compared to control (quasi-neutral) regions: genome-wide, genes and nonsynonymous sites.

241

9. Demography and migration of HPG1

243 <u>9.1 Skygrid coalescent</u>

244 From the Bayesian phylogenetic analyses described previously (section 7.2.2), we studied the 245 demographic model estimated via Skygrid. We reconstructed a skyline plot that depicts changes in effective population size, a measure of relative diversity, through time²⁶ (Fig. 3B). Sampling biases 246 247 could produce artefactual effects in the Skygrid plot. Nevertheless we expect these to be minor since 248 our dataset has a continuous sampling over a century (>2 samples per decade) instead of a 249 two-timepoints sampling, as it is common in ancient DNA studies where the rarity of the samples are 250 a limiting factor²⁷. An additional BEAST run was performed only with modern samples to verify that 251 the corresponding part of the tree and population sizes matched (data not shown). Implementation 252 of non-phylogenetic methodologies for demographic inference exist, e.g. Multiple Sequentially Markovian Coalescent (MSMC)²⁸, but after exploring them we concluded their resolution was 253 254 insufficient for analyses of the last several centuries. In order to compare with another method, we 255 got rough estimates of the diversity per decade as the average genetic differences between any two 256 samples per decade (Fig S6C).

258 <u>9.2. Phylogeography</u>

We performed another Bayesian phylogenetic analysis incorporating a geographic location trait^{29,30}. 259 260 For this, Brownian diffusion parameters are estimated by fitting a continuous gradient of geographic 261 locations along tree branches, starting from the leaves of the tree for which geographic locations are 262 known, i.e., the collection sites of our samples. We excluded three samples from the West coast of 263 the United States, separated over three thousand kilometers from the rest, since these do not fit a 264 gradual propagation by Brownian diffusion. We ran this analysis with the parameters described 265 previously (section 7.2.2) and sliced the resulting 3D (temporal and geographical) phylogeny at the early 16th and late 18th century using SPREAD software³¹ (Fig. S6B). Similar to before, we roughly 266 267 estimated "local diversity" for each sampling location by computing the average genetic differences 268 to the 10 closest neighbours (Fig. S6D).

269

270 <u>9.3. Isolation by distance</u>

We employed a heuristic search³² using an isolation-by-distance pattern to find the origin of 271 272 diffusion of HPG1 in North America, and compared it to the phylogeography analyses. We performed 273 a regression between genetic distances on geographic distances for all pairs of samples (genetic 274 distance ~ Euclidean geographic distances). This pattern, known as isolation by distance pattern, 275 reflects that as individuals are more geographically apart, they differ more genetically. We evaluated 276 whether this relationship was still significant for each of our samples separately (i.e., from a focal 277 sample, does genetic distance increase as geographic distance increases?). Only the significant 278 samples were retained and plotted since those points are the expected origins of migrations (Fig. S6 279 E,F). Arrows can be plotted in the direction of the maximum slope to illustrate migration trajectories. 280 This was done separately for historic and modern samples.

281

282 **10.** Phenotypic association analyses and dating of newly arisen mutations

- 283 <u>10.1. Phenotyping</u>
- 284 10.1.1 Root

Fifteen root phenotypes were scored for ≥ 10 replicates per genotype over a time-series experiment at the Gregor Mendel Institute in Vienna, using image analysis as described in detail elsewhere³³. We used the means per genotypes and per time series for association analyses.

- 288
- 289 10.1.2 Seed size

We spread the seeds of given genotypes on separate plastic square 12 x 12 cm Petri dishes. For faster image acquisition we used a cluster of eight Epson V600 scanners. The scanner cluster was operated by the BRAT Multiscan image acquisition tool bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license.
(www.gmi.oeaw.ac.at/research-groups/wolfgang-busch/resources/brat/). The resulting 1600 dpi
images were analyzed in Fiji software. Scans were converted to 8-bit binary images, thresholded
(parameters: setAutoThreshold("Default dark"); setThreshold(20, 255)) and particles analyzed
(inclusion parameters: size=0.04-0.25 circularity=0.70-1.00). The 2D seed size was measured in
square millimeters (parameters: distance=1600 known=25.4 pixel=1 unit=mm) for 2 plants per
genotype, > 500 seeds per plant.

299

300 10.1.3 Flowering in the growth chamber

301 We estimated the flowering time in growth chambers under four vernalization treatments (0, 14, 28 302 and 63 days of vernalization). We grew 6 replicates per accession divided between two complete 303 randomized blocks for each treatment. Seeds were sown on a 1:1 mixture of Premier Pro-Mix and 304 MetroMix and cold stratified for 6 days (6°C, no light). We then let plants germinate and grow at 305 18°C, 14 hours of light, 65% humidity. After 3 weeks, we transferred the plants to vernalization 306 conditions (6°C, 8 hours of light, 65% humidity). After vernalization, plants were transferred back to 307 long day conditions. Trays were rotated around the growth chambers every other day throughout the 308 experiment, under both vernalization and ambient conditions. Germination, bolting and flowering 309 dates were recorded every other day until all plants had flowered. Days till flowering or bolting times 310 were calculated from the germination date until the first flower opened and until the first flower bud 311 was developed, respectively. The average flowering time and bolting time per genotype were used 312 for association analyses.

313

314 10.1.4 Fecundity in the field

315 To investigate variation in fecundity in natural conditions, we grew three replicates of each accession 316 in a field experiment following a completely randomized block design. Seeds were sown from 317 09/20/2012 to 09/22/2012 in 66-well trays (well diameter = 4 cm) on soil from the field site where 318 plants were to be transplanted. The trays were cold stratified for seven days before being placed in a 319 cold frame at the University of Chicago (outdoors, no additional light or heat, but watered as needed 320 and protected from precipitation). Seedlings were transplanted directly into tilled ground at the 321 Warren Wood field station (41.84° N., 86.63° W.), Michigan, USA on 10/13/2012 and 10/14/2012. 322 Seedlings were watered-in and left to overwinter without further intervention. Upon maturation of 323 all fruits, stems were harvested and stored between sheets of newsprint paper. To estimate the 324 fecundity, stems were photographed on a black background and the size of each plant was estimated 325 as the number of pixels occupied by the plant on the image. This measure correlates well with the 326 total length of siliques produced, a classical estimator of fecundity in A. thaliana (Spearman's 327 rho=0.84, p-value<0.001, data not shown).

328

329 <u>10.2 Quantitative genetic analyses</u>

For 63 modern accessions, we measured time to bolting and flowering, seeds per plant, seed size, and 15 root phenotypes in common chamber or common garden settings. For all 100 accessions, climatic information from the bioclim database (<u>http://www.worldclim.org/bioclim</u>) was extracted using their geographic coordinates. For historic samples, some locations were only known by county name. In this case we assigned the geographic coordinate location of the centroid of the county.

335

336 10.2.1 Heritability

We performed association analyses using the R package GenABEL³⁴, with measured phenotypes (p =337 338 25) and climatic variables (c = 18) as response variables and SNPs as explanatory variables. A 339 Minimum Allele Frequency (MAF) cutoff of 5% was used. The number of assessed SNPs was 391 in a 340 dataset of only modern samples but with imputed genotypes for missing data using Beagle v4.0 (ref. ³⁵), and 456 SNPs with a dataset of modern and historic samples, without imputation. For all 341 342 associations, at least 63 individuals were genotyped for a specific SNP. We first investigated broad 343 sense heritability (H^2) of each trait using ANOVA partition of variance between and within lines using 344 replicates (Table S4). Significance was obtained by common F test in ANOVA. Secondly we used the 345 polygenic hglm function to fit a genome wide kinship matrix to calculate a narrow sense heritability 346 estimate (h^2) . Significance was calculated employing a likelihood ratio test comparing with a null model. In principle, h^2 is a component of H^2 , then its values should theoretically be $h^2 < H^2$; that is 347 348 not our case. Our result cannot be interpreted in this framework, since the calculation of both was not done with the same samples: for the h^2 calculation we employed genotype means whereas for 349 350 the H^2 we used multiple replicated measurements per genotype. The averaging of replicates per genotype in h^2 reduced environmental and developmental noise and thus we would expect $h^2 > H^2$. 351 We did this so the climatic estimates of h^2 , for which we only have one value per genotype, would be 352 353 comparable with the phenotypic h2 ones (Table S4).

354

355 10.2.2 Linear Models

For association analyses we first employed a linear mixed model that fitted the kinship matrix using the *mmscore* function, and only three significant SNP hits were discovered using a 5% significance threshold after False Discovery Rate correction (FDR). This was expected since we have few variants and these would have originated in an approximated phylogeny structure. We concluded that fitting the kinship matrix in our model was not appropriate since there would be no residual variation for association with specific SNPs. With this rationale we employed a fixed effects linear model using the *qtscore* function³⁶. To reduce the false-positive rate we took a conservative permutation strategy by bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license.
carrying out association with over 1,000 randomized datasets (permuting phenotypes across individuals) and used the resulting p-value distribution to correct p-values estimated with the original dataset. SNPs with p-values below 5% in the empirical p-value distribution were considered significant (Table S5). In climatic models, we included longitude and latitude as covariates to correct for any spurious association between SNPs and climate gradients created by the migratory pattern of isolation by distance.

369

370 10.2.3 Evaluation of significance

371 Significant SNPs were interspersed throughout the genome (Fig. S7) and their p-values and 372 phenotypic effects did not correlate with the minimum age of the SNPs nor with their allele 373 frequency (Fig. S10), something that could have indicated that the significance was merely driven by 374 the higher statistical power of intermediate frequency variants. Using QQ plots to assess inflation or 375 deflation of p-values, we observed generally that permutation corrected p-values were deflated. 376 Straight series of points in QQ plots indicate identical p-values for multiple SNPs, a pattern that we 377 attributed to long range LD, i.e. lack of independence (see Graphic Table S7 for trait distributions and 378 QQ plots from each association analysis). We also used a False Discovery Rate correction for the raw 379 p-values using p.adjust in R and, as a sanity check, we used a Bonferroni-corrected threshold, a 380 procedure considered over-stringent in association analyses (Table S5). This was calculated as: 5% / 381 (number of SNPs + number of traits) ~ 0.01%.

382

383 10.2.4 Context of de novo mutations associated with phenotypes

For each SNP in our dataset, we determined the ancestral and derived states, by identifying which allele was found in the oldest herbarium samples. We compared the time of emergence and the centroid of geographic distribution of the alternative alleles of SNP hits to random draws of SNPs with the same MAF filtering (5%) (Fig. S8).

388

389 10.2.5 Functional information

390 On top of phenotypic and climatic associations of SNP hits, we also provide a likely functional effect employing a commonly used amino acid matrix of biochemical effects³⁷. Functional information of 391 392 name categorization of SNP gene and ontology hits was obtained from 393 www.arabidopsis.org/portals/genAnnotation/gene structural annotation/annotation data.jsp and 394 www.arabidopsis.org/tools/bulk/go/ (Table 1 and Table S5).

395

396 10.2.6 Proof of concept examples

397 We argue that the power of an association approach relies on the fact that HPG1 lines resemble Near Isogenic Lines (NILs) produced by experimental crosses³⁸ (Fig. S9A). Similar to genome-wide 398 399 association studies (GWA), power depends on many factors, namely the noise of phenotype under 400 study, architecture of phenotypic trait, quality of genotyping, population structure, sample diversity, 401 sample size, allele frequency, and recombination. On one hand, association analyses in NILs suffer 402 from large linkage blocks, but confident results can be achieved due to accurate measurement of 403 phenotypes, limited genetic differences between any two lines, and high quality genotypes. In 404 common GWA studies such as in humans, there are multiple confounding effects. Among the 405 confounders are (1) that any two samples differ in hundreds of thousands of SNPs, and (2) that 406 historical and geographic stratification produce non-random correlations among those SNP 407 differences. This considerably complicates the identification of phenotypic effects at specific genes, 408 and power relies greatly on large sample sizes to achieve the sufficient number of recombination 409 between markers.

410 To provide support for the non-synonymous SNP on chromosome 5, at position 6,508,329 in 411 AT5G19330, we looked for pairs of lines that carry the ancestral and the derived allele, but that differ 412 in few (or no other) SNPs in the genome. When considering all genic substitutions with a minimum 413 allele frequency of 5% (Fig. S9A), we identified 20 pairs of lines differing only in the AT5G19330 SNP 414 and another linked SNP (located on a different chromosome, association p-value > 0.4). The 415 phenotypic differences in mean gravitropic score of these almost-identical pairs were significantly 416 higher than phenotypic differences among all pairs of HPG1 lines, and genetically identical pairs 417 attending to substitutions inside genes (Fig. S9A). Furthermore, this SNP was not in complete linkage with any other SNP hit ($r^2 < 0.5$) (Fig. S7D). The same approach was used to examine the SNPs in 418 419 AT1G54440 (Fig. S7E) and AT2G16580 (Fig. S7F), which represent an intermediate and a high LD 420 example.

421

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501 SUPPLEMENTAL FIGURES





503 Figure S1. Ancient-DNA-like characteristics of unrepaired herbarium libraries.

(A) Fraction of *A. thaliana* DNA in sample. (B) Median length of merged reads. (C) Fraction of
cytosine to thymine (C-to-T) substitutions at first base (5' end). (D) Relative enrichment of purines
(adenine and guanine) at 5' end breaking points. Position -1 is compared with position -5 (negative
numbers indicate genomic context before upstream reads' 5' end).





511 (A) Neighbor-joining tree built using Illumina-based SNP calls at the 149 genotyping markers 512 originally used to identify HPG1 candidates (consensus of 1,000 replicates). HPG1 accessions are 513 shown in black, whereas other North American lineages are depicted in red (see explanation below 514 for four HPG1-like accessions). (B) Neighbor-joining tree based on genome-wide SNPs (consensus of 515 1,000 replicates). Accessions colored as in (A). Note that three accessions originally classified as 516 HPG1 based on 149 SNPs (A) are placed outside this clade. A further accession (BRR7) within the 517 HPG1 main branch was a recombinant removed from the analysis. (C) First two dimensions of a 518 multidimensional scaling plot based on identity-by-state pairwise distances. Notice that black dots 519 represent multiple transparent dots overlaid, a result of multiple almost-identical HPG1 genomes. 520 Percentage of the variance explained by each dimension given in parentheses.

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523

522

524 Figure S3. Substitution spectrum and rates.

(A) "Unfolded" site frequency spectrum using *Arabidopsis lyrata* as outgroup for all transitions and
transversions. Bar plot shows proportions of different types of substitutions divided by genomic
annotation. (B) Distributions of "net" pairwise genetic distances between historic and modern
samples used to calculate mutation rates (from quality 32_15 and complete information per site).
UTRs were excluded because of the small number of SNPs. (C) Mutation rates calculated for different
genomic annotations and quality thresholds (32_32, 32_15, 24_24) and missing values (NA50:

- 531 maximum 50% missing data per SNP; COMPL: missing data 0%). Mean and 95% confidence intervals
- are shown.
534

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(A, B) Fraction of methylation of cytosines in HPG1 pseudo-reference⁷ at intergenic (A) or coding
regions (B). (C, D) Fraction of methylation of cytosines in Col-0 reference genome⁵ at intergenic (C) or
coding regions (D). In each of the four comparisons, a grey histogram represents distribution of
methylation of 1,000 random sets of invariant cytosines. Lines represent average methylation degree
at those sites in HPG1 that changed from cytosine to thymine (red). We differentiate those
substitutions that are shared - fixed - across all individuals (light red) or whose allele are present at

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- an intermediate segregating frequency (dark red). Likewise, average methylation is shown for sites
- 544 that changed from cytosine to adenine (blue) that that are fixed (light blue) or segregating (dark
- 545 blue). The fact that the average methylation is higher in new substitutions than in invariant positions
- 546 supports a connection between methylation and mutability of sites.

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We tested for an interaction in a 2x2 table of counts of SNPs using Fisher's Exact Test. The tables were built with the number of SNPs falling into each of the two annotations: genome - intergenic (A), genic - intergenic (B), and synonymous - non-synonymous (C); and two discrete allele groups assuming a minimum allele frequency (MAF) cutoff. We repeated the test by sliding the cut-off from 0 to 50% allele frequency (x axis) and we show the corresponding p-value (y axis). The dashed line indicates the 5% significance threshold.

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560 Figure S6. Phylogeographic inference in HPG1.

561 (A, B) The model infers the most probable geographic location of each of the nodes of the phylogeny 562 in Figure 3. (A) Ancestral distribution map (dark red/brown) summarizing the first ~100 years of the 563 phylogenetic tree. Clouds represent the 95% interval of the Highest Posterior Probability Density of 564 locations. (B) Current distribution map (dark blue) summarizing the last ~100 years. Clouds as in (A). 565 (C) Diversity in time. Each point represents the average number of genetic changes between a 566 sample and the other samples within a decade. The blue line shows the fit using a generalized 567 additive model and the grey shaded area the 95% confidence interval. (D) Diversity in space. Each 568 point represents the average number of genetic changes among the 10 geographically closest 569 neighbors. Genetic distances are shown qualitatively from a red (low) to blue (high) gradient. (E) 570 Origin of herbarium and modern geographic spread, determined using separate heuristic searches of bioRxiv preprint first posted online Apr. 25, 2016; doi: http://dx.doi.org/10.1101/050203. The copyright holder for this preprint (which was not peer-reviewed) is the author/funder. It is made available under a CC-BY-NC-ND 4.0 International license.
isolation-by-distance patterns. Three locations of modern samples and four locations of herbarium samples showed significant slopes (p < 0.05) in the isolation-by-distance pattern. That is, genetic distance increased when moving away from these geographic locations. For one sample of each subset, herbarium (F) and modern (G), a likely migration trajectory is depicted by an arrow and its isolation-by-distance pattern is shown.

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580 Figure S7. Density of SNPs along all chromosomes and location of SNP hits.

581 Black line shows number of SNPs per 100 kb window. Centromere locations are indicated by grey 582 shading. Vertical lines indicate SNPs associated with root phenotypes (red) and climatic variables 583 (blue) (see Table S5).

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587 Figure S8. Spatial and temporal emergence of root-associated mutations.

(A) Age distribution of derived SNPs with a significant trait association (the herbarium sample in which they were first recorded) (red), compared with genome-wide SNPs with at least 5% minor allele frequency (grey), or without frequency cutoff (black). (B) Spatial centroid of all samples carrying a derived allele. Since it is an average location, centroids can be in a body of water. Ten random draws of 50 SNPs for each category were used to produce the density lines in (A) and points in (B).

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Figure S9. Linkage disequilibrium between SNPs with significant trait associations.

600 (A-F) Linkage disequilibrium between SNPs with significant trait associations. Histogram of genetic 601 distances (A) between samples when evaluating only coding regions at 5% minimum allele frequency. Linkage disequilibrium between SNP hits measured as r^2 (B) and D' (C). Three significant 602 603 SNPs were further studied to exemplify the power of association analyses with HPG1. For each, 604 phenotypic differences between accessions that differ in the focal SNP and that are otherwise 605 virtually genetically identical are compared both with all pairs of accessions and with pairs of 606 accessions completely identical for coding regions. Below each violin plot is the histogram of linkage 607 disequilibrium of the focal SNP with all other SNP hits. The three focal SNPs evaluated are located in 608 AT5G19330 (D), AT1G54440 (E) and AT2G16580 (F).

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- ⁶¹³ Figure S10. Correlations of SNP effects and p-values with frequency and age.
- 614 Correlation between SNP frequency and p-value (A), frequency and effect (B), age and p-value (C),
- 615 age and effect **(D)**. All cases were non-significant.
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618 SUPPLEMENTAL TABLES

- 619 See appended .pdf file for Tables S1-5.
- 620 See appended .pdf file for Graphic Table S7: For each trait employed in association analyses, we
- ⁶²¹ report the histogram distribution and the QQ plot of p-values to ensure that no trait departs
- 622 exaggeratedly from the normal distribution, and that no inflation of p-values is observed (when
- 623 lambda \leq 1, there is no inflation of false positives).

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Table S1. Sample information.

(Abbreviation H* indicates herbarium samples that cluster with the mo

dern HPG1 clade rather than the historic HPG1 clade in Fig. 3., highlighted as a star in the map from Fig. 1. Abbreviations of herbarium collections or seed sources: UCONN = University of Connecticut Herbarium; CFM = Chicago Field Museum; NY = New York Botanical Garden; ABRC = Arabidopsis Biological Resources Center; OSU = Ohio State University.)

	Accession	Latitude (ºN)	Longitude (º E)	State	Date collected	Alternative name	Collector/ Herbarium	Average coverage (x)	Number of covered positions (≥3x) (mapped against HPG1 reference)	Number of covered positions (≥3x) (mapped against Col_0 reference)	SNPs vs HPG1 reference	Belongs to HPG1	Modern/ Herbarium	Column number in the available genome matrix
JK399		38.7155	-75.635591	DE	1863	888124	NY	9	105,053,631	99,889,683	142	yes	Н	101
JK366		43.1921	-77.0102	NY	1866	888144	NY	6.8	100,379,839	95,118,236	123	yes	Н	94
JK395		38.9068	-77.036667	DC	1877	888134	NY	10.3	103,620,791	98,888,406	167	yes	Н	100
JK888141		40.732007	-74.068455	NJ	1879	888141	NY	42	107,211,409	102,634,255	161	yes	Н	103
JK389		38.9068	-77.036667	DC	1888	1365363	NY	9.9	106,042,465	100,826,958	151	yes	н	98
JK362		38.9068	-77.036667	DC	1889	1365364	NY	8.8	103,997,716	98,876,320	153	yes	Н	93
JK367		40.9249	-74.0755	NJ	1890	1365344	NY	16.7	107,236,732	102,176,782	181	yes	Н	95
JK372		41.1222	-74.3569	NJ	1890	1365332	NY	14.8	106,285,178	101,480,369	163	yes	Н	96
JK1365354		38.8782	-77.09048	VA	1891	1365354	NY	36.4	106,718,326	102,458,166	169	yes	Н	88
JK376		39.97	-83.01	NY	1891	1365337	NY	12.3	105,962,154	100,840,125	145	yes	Н	97
JK351		41.15	-73.766667	NY	1894	1365333	NY	16.1	106,531,302	101,841,156	153	yes	Н	90
JK355		35.99	-83.94	TE	1896	1365374	NY	14.3	106,391,637	101,455,311	192	yes	Н	91
JK356		n/a	n/a	GA	1897	1365375	NY	5.3	90,426,010	89,296,191	n/a	no	Н	92
JK393		n/a	n/a	NC	1897	1365370	NY	30.4	102,894,430	101,298,068	n/a	no	Н	99
JK346		40.643136	-111.95177	UT	1903	102365	NY	29.1	107,223,283	102,450,446	222	yes	Н	89
JK2525		41.224343	-73.06021	СТ	1904	79391	UNCONN	12.5	105,025,845	n/a	138	yes	Н	118
JK2529		n/a	n/a	ОН	1904	176849	CFM	11.4	100,620,441	n/a	n/a	no	Н	121

JK401	40.643136	-111.95177	UT 19	04 102364	NY	10.4	99,572,736	94,661,828	216	yes	Н	102
JK2513	41.102121	-81.560547	OH 19	11 25	OSU	18.2	106,309,854 n	/a	176	yes	Н	108
JK2509	n/a	n/a	CT 19	17 11	OSU	15.1	102,169,546 n	/a	n/a	no	Н	104
JK2530	41.482862	-86.822602	IN 19	22 531679	CFM	22.2	107,043,540 n	/a	161	yes	Н	122
JK2526	41.666667	-73.508455	CT 19	29 79409	UNCONN	16.3	107,026,827 n	/a	161	yes	Н	119
JK2515	41.137296	-81.863779	OH 19	30 30	OSU	21.3	106,893,416 n	/a	193	yes	Н	110
JK2511	41.721618	-81.243317	OH 19	34 14	OSU	5.6	95,822,372 n	/a	109	yes	Н	106
JK2523	n/a	n/a	OH 19	40 25707	UNC	13.1	101,421,749 n	/a	n/a	no	Н	116
JK2520	n/a	n/a	OH 19	45 54051	UNC	20.3	102,831,697 n	/a	n/a	no	Н	114
JK2524	39.856783	-74.686954	NJ 19	52 63978	B UNC	13.8	100,778,282 n	/a	n/a	no	Н	117
JK2512	39.95607	-81.953309	OH 19	56 21	OSU	16.7	106,801,844 n	/a	189	yes	Н	107
JK2514	39.95607	-81.953309	OH 19	69 27	OSU	28.4	107,044,415 n	/a	219	yes	Н	109
JK2517	n/a	n/a	OH 19	81 34	OSU	21.7	102,643,436 n	/a	n/a	no	Н	112
JK2521	n/a	n/a	OH 19	92 565960	UNC	2.9	62,673,938 n	/a	n/a	no	Н	115
JK2518	41.867643	-80.789021	OH 19	93 40	OSU	14.8	106,578,197 n	/a	177	yes	Н	113
JK2531	39.856783	-74.686954	NJ 19	52 1507461	CFM	15.1	106,158,181 n	/a	177	yes	H*	123
JK2510	39.688861	-82.993218	OH 19	30 13	OSU	21	106,305,970 n	/a	178	yes	H*	105
JK2527	41.509059	-72.543694	CT 19	75 79389	UNCONN	8.3	104,089,205 n	/a	200	yes	H*	120
JK2516	39.500862	-82.472413	OH 19	80 32	OSU	18.1	106,464,569 n	/a	198	yes	H*	111
CSHL_15	40.8585	-73.4675	NY 19	93 CSHL-15	ABRC	39.3	108,189,771	105,955,885	243	yes	Μ	16
CSHL_17	40.8585	-73.4675	NY 19	93 CSHL-17	ABRC	41.5	108,194,960	105,982,511	240	yes	Μ	17
FM_10	42.4489	-76.5072	NY 19	93 FM-10	ABRC	44.6	108,203,215	106,052,866	269	yes	Μ	20
FM_11	42.4489	-76.5072	NY 19	93 FM-11	ABRC	44.4	108,214,008	106,040,276	288	yes	Μ	21
HS_12	42.373	-71.0627	MA 19	93 HS-12	ABRC	48.8	108,230,030	106,124,249	251	yes	Μ	25
HS_17	42.373	-71.0627	MA 19	93 HS-17	ABRC	55.3	108,242,062	106,155,362	254	yes	Μ	26
Kno_10	41.2816	-86.621	IN 19	93 Kno-10	ABRC	39.4	108,198,601	105,985,288	226	yes	Μ	32
KNO_15	41.2816	-86.621	IN 19	93 KNO-15	ABRC	43.6	108,219,683	106,069,077	231	yes	Μ	33
Gre_0	43.178	-85.2532	MI 19	95 Gre-0	ABRC	44.6	108,209,345	106,032,827	207	yes	Μ	22
Tul_0	43.2708	-85.2563	MI 19	95 CS6877	ABRC	31.2	108,140,393	105,806,418	221	yes	Μ	85
CS8067	41.3599	-122.755	CA 19	96 Buckhorn Pa	s: ABRC	66.4	108,260,489	106,243,277	294	yes	Μ	15
Tol_2	41.6639	-83.5553	OH 19	96 CS8022	ABRC	61	108,241,333	106,194,209	238	yes	Μ	83
Tol_3	41.6639	-83.5553	OH 19	96 CS8023	ABRC	40.2	108,184,749	105,953,559	232	yes	Μ	84
MIA_1	41.7976	-86.6691	MI 19	99 MIA-1	ABRC	73.1	108,279,881	106,291,612	234	yes	Μ	56

MIA_5	41.7976	-86.6691 MI	1999 MIA-5	ABRC	62.9	108,263,557	106,250,560	235 yes	М	57
MIC_20	41.8266	-86.4366 MI	1999 MIC-20	ABRC	39.9	108,200,416	106,010,135	237 yes	М	58
MIC_24	41.8266	-86.4366 MI	1999 MIC-24	ABRC	33.8	108,176,527	105,728,326	237 yes	М	59
Brn_10	41.9	-86.583 MI	2002 Brn-10	ABRC	33.3	108,177,381	105,905,097	243 yes	М	7
Brn_24	41.9	-86.583 MI	2002 Brn-24	ABRC	38.4	108,208,482	105,951,803	228 yes	М	8
Haz_10	41.879	-86.607 MI	2002 Haz-10	ABRC	33.8	108,154,100	105,903,700	230 yes	М	23
Haz_2	41.879	-86.607 MI	2002 Haz-2	ABRC	39.7	108,201,103	106,004,251	288 yes	М	24
Ker_4	42.184	-86.358 MI	2002 Ker-4	ABRC	32.1	108,132,127	105,806,486	261 yes	М	30
Ker_5	42.184	-86.358 MI	2002 Ker-5	ABRC	62.9	108,259,905	106,246,278	259 yes	М	31
L_R_10	41.847	-86.67 MI	2002 L-R-10	ABRC	22.4	108,062,944	105,496,224	186 yes	М	49
L_R_5	41.847	-86.67 MI	2002 L-R-5	ABRC	60.6	108,255,795	106,209,826	299 yes	М	50
Lak_12	41.8	-86.67 MI	2002 Lak-12	ABRC	37.8	108,176,901	105,775,999	237 yes	М	36
Lak_13	41.8	-86.67 MI	2002 Lak-13	ABRC	28.5	107,955,559	105,553,559	226 yes	М	37
Map_35	42.166	-86.412 MI	2002 Map-35	ABRC	64.7	108,265,863	106,224,216	290 yes	М	51
Map_42	42.166	-86.412 MI	2002 Map-42	ABRC	46	107,303,032	106,093,945 r	n/a no	М	52
Map_8	42.166	-86.412 MI	2002 Map-8	ABRC	33.4	108,155,999	105,921,907	287 yes	М	53
Mdn_10	42.051	-86.509 MI	2002 Mdn-10	ABRC	34.9	108,106,772	105,906,924 r	n/a no	М	54
Mdn_8	42.051	-86.509 MI	2002 Mdn-8	ABRC	37.4	108,199,679	105,940,666	266 yes	М	55
Paw_13	42.148	-86.431 MI	2002 Paw-13	ABRC	43	108,159,739	105,980,721	267 yes	М	70
Paw_20	42.148	-86.431 MI	2002 Paw-20	ABRC	41.3	108,218,762	106,059,867	241 yes	М	71
Riv_25	42.184	-86.382 MI	2002 Riv-25	ABRC	36.8	108,186,632	105,779,717	273 yes	М	76
Riv_26	42.184	-86.382 MI	2002 Riv-26	ABRC	35.7	108,194,281	105,958,738	260 yes	М	77
Yng_4	41.865	-86.646 MI	2002 Yng-4	ABRC	41.3	108,182,789	106,000,003	289 yes	М	86
Yng_53	41.865	-86.646 MI	2002 Yng-53	ABRC	46	108,230,553	106,125,861	191 yes	М	87
RRS_10	41.5609	-86.4251 IN	2003 RRS-10	ABRC	41.8	108,208,144	106,033,465	274 yes	М	80
DuckLkSP38	43.3431	-86.4045 MI	2004 DuckLkSP38	ABRC	37.1	108,171,751	105,932,415	253 yes	М	18
DuckLkSP40	43.3431	-86.4045 MI	2004 DuckLkSP40	ABRC	39.6	108,204,654	105,969,244	257 yes	М	19
KBS_Mac_68	42.405	-85.398 MI	2004 KBS-Mac-68	ABRC	41.3	108,181,390	105,870,424	259 yes	М	27
KBS_Mac_74	42.405	-85.398 MI	2004 KBS-Mac-74	ABRC	37.7	108,160,645	105,801,702	265 yes	М	28
MNF_Che_47	43.5251	-86.1843 MI	2004 MNF-Che-47	ABRC	27.6	108,093,393	105,596,885	281 yes	М	60
MNF_Che_49	43.5251	-86.1843 MI	2004 MNF-Che-49	ABRC	28.5	108,082,202	105,661,610	274 yes	М	61
MNF_Pin_40	43.5356	-86.1788 MI	2004 MNF-Pin-40	ABRC	47.9	108,238,775	106,099,919	287 yes	М	62
MNF_Pot_10	43.595	-86.2657 MI	2004 MNF-Pot-10	ABRC	61.4	108,189,553	106,228,588 r	n/a no	М	63

MNF_Pot_15	43.595	-86.2657 MI	2004 MNF-Pot-15	ABRC	25.2	108,543,185	107,022,924 n	/a no	Μ	64
MSGA_10	43.2749	-86.0891 MI	2004 MSGA-10	ABRC	41.9	108,191,659	106,019,404	233 yes	Μ	65
MSGA_12	43.2749	-86.0891 MI	2004 MSGA-12	ABRC	42.8	108,227,214	106,032,928	240 yes	Μ	66
MSGA_61	43.2749	-86.0891 MI	2004 MSGA-61	ABRC	45.5	108,210,152	106,077,183	247 yes	Μ	67
MuskSP_68	43.2483	-86.3368 MI	2004 MuskSP-68	ABRC	25.8	108,063,297	105,588,467	215 yes	Μ	68
MuskSP_83	43.2483	-86.3368 MI	2004 MuskSP-83	ABRC	29.9	108,099,368	105,721,042	222 yes	Μ	69
Pent_46	43.7623	-86.3929 MI	2004 Pent-46	ABRC	48.3	108,227,763	106,099,890	238 yes	Μ	72
Pent_7	43.7623	-86.3929 MI	2004 Pent-7	ABRC	55.7	108,220,625	106,144,167	240 yes	Μ	73
SLSP_67	43.665	-86.496 MI	2004 SLSP-67	ABRC	53.5	108,238,880	106,143,530	245 yes	Μ	81
SLSP_69	43.665	-86.496 MI	2004 SLSP-69	ABRC	35.5	108,160,835	105,899,252	249 yes	Μ	82
KNO2_41	41.273	-86.625 IN	2005 KNO2.41	ABRC	44.7	108,209,694	106,063,235	219 yes	Μ	34
KNO2_54	41.273	-86.625 IN	2005 KNO2.54	ABRC	44	108,212,430	105,903,373	218 yes	Μ	35
LI_EF_011	40.9064	-73.1493 NY	2005 LI-EF-011	ABRC	68.6	108,267,109	106,250,331	259 yes	Μ	38
LI_EF_018	40.9064	-73.1493 NY	2005 LI-EF-018	ABRC	39	108,244,306	105,898,497	230 yes	Μ	39
LI_OF_061	40.7777	-72.9069 NY	2005 LI-OF-061	ABRC	58	104,897,841	105,729,196 n	/a no	Μ	40
LI_RR_096	40.9447	-72.8615 NY	2005 LI-RR-096	ABRC	63.5	108,264,679	106,251,487	261 yes	Μ	41
LI_RR_097	40.9447	-72.8615 NY	2005 LI-RR-097	ABRC	40.8	108,211,310	105,992,095	249 yes	Μ	42
LI_SET_019	40.9352	-73.114 NY	2005 LI-SET-019	ABRC	29.9	108,085,297	105,737,781	259 yes	Μ	43
LI_SET_036	40.9352	-73.114 NY	2005 LI-SET-036	ABRC	41.5	108,216,592	106,006,605	238 yes	Μ	44
LI_WP_039	40.9076	-73.2089 NY	2005 LI-WP-039	ABRC	104.8	108,301,282	106,273,259	239 yes	Μ	45
LI_WP_041	40.9076	-73.2089 NY	2005 LI-WP-041	ABRC	76.5	108,287,248	106,322,146	235 yes	Μ	46
PT1_52	41.3423	-86.7368 IN	2005 PT1.52	ABRC	50.6	108,240,431	106,154,252	219 yes	Μ	74
PT1_85	41.3423	-86.7368 IN	2005 PT1.85	ABRC	46.1	108,220,150	106,097,633	233 yes	Μ	75
RMX4_118	42.036	-86.511 MI	2005 RMX4.118	ABRC	41.8	106,178,554	105,685,651 n,	/a no	Μ	78
11PNA1_14	42.0945	-86.3253 MI	2006 11PNA1.14	ABRC	47.5	108,227,783	106,133,372	276 yes	Μ	1
328PNA062	42.0945	-86.3253 MI	2006 328PNA062	ABRC	47.3	108,221,709	106,127,272	223 yes	Μ	2
627ME_13Y1	42.093	-86.359 MI	2006 n/a	ABRC	53.4	107,908,679	106,148,671 n	/a no	Μ	3
627ME_1MI1	42.093	-86.359 MI	2006 627ME-1MI1	ABRC	57.8	108,252,617	106,173,403	281 yes	Μ	4
627RMX_1MN4	42.0333	-86.5128 MI	2006 n/a	ABRC	43.6	106,799,549	105,789,469 n	/a no	Μ	5
627RMX_1MN5	42.0333	-86.5128 MI	2006 n/a	ABRC	50.6	106,885,430	105,897,441 n	/a no	Μ	6
BRR107	40.8313	-87.735 IL	2006 BRR107	ABRC	28.5	108,896,513	107,320,745 n,	/a no	Μ	9
BRR12	40.8313	-87.735 IL	2006 BRR12	ABRC	43.9	108,190,572	106,031,493	232 yes	Μ	10
BRR23	40.8313	-87.735 IL	2006 BRR23	ABRC	30.7	108,095,072	105,726,913	236 yes	М	11

BRR4	40.8313	-87.735 IL	2006	5 BRR4	ABRC	4	4.7	108,180,840	106,033,507	219 yes	М	12
BRR57	40.8313	-87.735 IL	2006	5 BRR57	ABRC	2	8.4	108,093,033	105,630,963	225 yes	М	13
BRR60	40.8313	-87.735 IL	2006	5 BRR60	ABRC	4	2.9	108,281,285	106,199,572	229 yes	М	14
KEN	41.767	-72.677 CT	n/a	KEN	ABRC	5	5.2	108,233,232	106,158,223	249 yes	М	29
LP3413_31	41.6862	-86.8513 IN	n/a	LP3413.31	ABRC	5	5.9	108,244,332	106,190,596	227 yes	М	47
LP3413_53	41.6862	-86.8513 IN	n/a	LP3413.53	ABRC	5	1.2	108,157,453	105,994,665	245 yes	М	48
RMX413_85	42.036	-86.511 MI	n/a	RMX413.85	ABRC		38	106,816,221	105,483,632 r	n/a no	М	79

Table S2. Sample information for Col-0 mutation accumulation lines.

Information about each Mutation Accumulation (MA) line and their number of SNPs at different annotations. Also the total number of SNPs, average number of mutations and total bp covered in the genome per annotation are reported.

MA line	Read depth	Generation	Total	SNPs	Deletions	insertions	CDS	Nonsyn	Syn	Intron	5' UTR	3' UTR	TE	Intergenic
0-4-26	57	3	7	6	1	0	0	0	0	0	0	0	1	5
0-8-87	49	3	7	5	0	2	1	1	0	1	0	0	0	3
30-109	45	31	31	23	7	1	3	3	0	3	0	0	2	15
30-119	45	31	33	26	2	5	1	1	0	1	2	0	4	18
30-29	51	31	39	26	10	3	2	1	1	3	0	1	5	15
30-39	48	31	28	18	7	3	1	1	0	1	0	1	4	11
30-49	50	31	30	23	3	4	4	4	0	0	0	0	6	13
30-59	40	31	46	31	8	7	5	2	3	2	0	0	6	18
30-69	50	31	26	21	3	2	4	3	1	1	1	1	6	8
30-79	50	31	31	25	3	3	6	4	2	2	0	0	8	9
30-89	39	31	35	27	5	3	4	3	1	1	1	0	2	19
30-99	44	31	37	35	1	1	6	5	1	2	0	2	8	17
Total SNPs				274			38	28	10	17	4	5	52	158
average (31st)			33.6	25.5	4.9	3.2	3.6	2.7	0.9	1.6	0.4	0.5	5.1	14.3
stdev (31st)			5.9	4.9	3.0	1.8	1.8	1.4	1.0	1.0	0.7	0.7	2.1	3.9
Total bp			115,954,227			30,	753,96	66		17,446,837	4,289,789	2,508,199	9,267,413	48,090,487

Table S3. Mutation rate estimates for different annotations in HPG1 and mutation accumulation lines.

Mutation rates from MA lines are compared to HPG1 substitution rates from the dataset of 32_15 quality filter and complete information (see SOM) (Abbreviations: stat, descriptive statistic; bp, base pairs; lower and upper, lower and upper 95% CI; Nonsyn. and Syn., nonsynonymous and synonymous sites; UTR, untranslated region sites; HPG1 adj., substitution rate of HPG1 adjusted by a mean generation time of 1.3 years)

Dataset	stat	CDS	Syn.	Nonsyn.	Intronic	5' UTR	3' UTR	Transposon	Intergenic	Genome
MA	mean	3.776	n/a	n/a	2.958	3.008	6.431	17.752	9.592	7.094
MA	sem	1.928	n/a	n/a	1.786	5.258	9.094	7.420	2.628	1.352
MA	lower	2.581	n/a	n/a	1.851	-0.251	0.794	13.153	7.964	6.256
MA	upper	4.971	n/a	n/a	4.065	6.267	12.067	22.351	11.221	7.932
HPG1	mean	2.149	n/a	n/a	1.540	n/a	n/a	2.290	3.029	2.114
HPG1	sem	0.108	n/a	n/a	0.165	n/a	n/a	0.536	0.173	0.119
HPG1	lower	1.943	n/a	n/a	1.231	n/a	n/a	1.314	2.698	1.871
HPG1	upper	2.364	n/a	n/a	1.874	n/a	n/a	3.309	3.368	2.344
HPG1 adj.	mean	2.794	n/a	n/a	2.002	n/a	n/a	2.977	3.938	2.748
HPG1 adj.	sem	0.140	n/a	n/a	0.214	n/a	n/a	0.697	0.225	0.154
HPG1 adj.	lower	2.526	n/a	n/a	1.600	n/a	n/a	1.708	3.508	2.432
HPG1 adj.	upper	3.073	n/a	n/a	2.436	n/a	n/a	4.302	4.378	3.047
Distributio	min	0	0	0	0	0	0	0	0	0
n of	1st qu.	2	1	1	1	0	1	2	5	9
noirwiso	median	5	3	3	3	1	2	4	10	18
	mean	5.6	3	3.1	3.8	1.2	1.9	4.3	11.3	21.1
difforences	3rd qu.	8	5	4	5	2	3	6	16	31
unterences	max.	27	17	11	15	5	7	22	43	87
Total numb	er of SNPs	971	531	448	629	74	158	656	2498	5013
Tota	l bp	32119233	n/a	n/a	18132262	2632130	4480510	6209512	43601507	108434034

Table S4. Description of phenotypic and climatic variables for association mapping analyses.

Mean and standard deviation (s.d.) across accessions for each phenotypic and climatic variables. Broad sense heritabilities (H2) were calculated from between line and within line (between replicate) variance in ANOVA. P-value corresponds to F test. Narrow sense heritabilities (h2) were calculated employing linear mixed models and kinship matrix from mean accession values. P-values correspond to Likelihood Ratio test.

Variable	Description	mean	s.d.	H2	p-value	h2	p-value
FT_V0	Time from germination until the first flower opens (days) under 0 days of vernalization	101	4.53	0.009	7.28E-03	0.017	1.97E-25
FT_V1	Time from germination until the first flower opens (days) under 14 days of vernalization	107	4.12	0.013	6.87E-04	0.395	1.83E-25
FT_V2	Time from germination until the first flower opens (days) under 28 days of vernalization	102	3.22	0.012	1.04E-03	0.429	3.37E-27
FT_V3	Time from germination until the first flower opens (days) under 63 days of vernalization	110	1.32	0.010	5.11E-03	0.226	9.52E-25
B_V0	Time from germination until the first developed bud (days) under 0 days of vernalization	88.8	4	0.013	8.99E-04	0.018	2.26E-25
B_V1	Time from germination until the first developed bud (days) under 14 days of vernalization	93.9	3.84	0.009	7.45E-03	0.340	3.98E-25
B_V2	Time from germination until the first developed bud (days) under 28 days of vernalization	89.2	2.13	0.005	6.92E-02	0.252	2.22E-25
B_V3	Time from germination until the first developed bud (days) under 63 days of vernalization	101	0.45	0.006	5.79E-02	0.177	1.99E-24
Fecundity	Pixel area of inflorescence (correlation with number of fruits, rho=0.84)	0.02	0.0042	0.001	3.56E-01	0.240	1.02E-22
seed_size	Average seed size (mm2)	0.134	0.0053	0.016	4.73E-03	0.149	3.58E-24
GR_rootLength	Average root growth rate	181	14.9	0.131	4.76E-77	0.640	3.13E-29
GR_shootArea	Average of shoot area growth rate	2279	253	0.053	2.33E-24	0.812	1.77E-31
rootLength	Average root length	467	35.8	0.048	2.01E-21	0.409	2.57E-28

dirEquivalent	Average root direction index. Score for average pixel-by- pixel deviations from growth relative to vector of gravity	0.393	0.0277	0.059	2.62E-28	0.544	1.14E-26
stdDevXY	Average root linearity coefficient of linear determination; R2 of linear regression line fitted to pixels of primary root skeleton	0.725	0.0429	0.018	4.54E-06	0.303	1.41E-25
meanRootWidth	Average root width	5.27	0.177	0.038	5.30E-16	0.359	1.52E-25
rootWidth20	Average width over first interval of the primary root length (0 to 20%) at hypocotyl/root junction	5.75	0.124	0.018	5.11E-06	0.166	3.37E-25
rootWidth40	Average width over first interval of the primary root length (20 to 40%) at hypocotyl/root junction	5.35	0.19	0.033	3.87E-13	0.291	1.76E-25
rootWidth60	Average width over first interval of the primary root length (40 to 60%) at hypocotyl/root junction	5.2	0.212	0.039	1.49E-16	0.405	6.51E-26
rootWidth80	Average width over first interval of the primary root length (60 to 80%) at hypocotyl/root junction	5.11	0.241	0.045	4.67E-20	0.381	5.47E-26
rootWidth100	Average width over first interval of the primary root length (80 to 100%) at hypocotyl/root junction	4.9	0.222	0.038	4.06E-16	0.351	8.81E-26
gravitropicDir	Average root angle between root vector and the vertical axis of the picture (assumed vector of gravity) (°)	-7.22	2.56	0.024	7.69E-09	0.210	4.68E-27
gravitropicScore	Average score for root angle intervals	0.1	0.0457	0.044	2.83E-19	0.642	7.56E-27
TotLen.EucLen	Average root tortuosity: Total root length divided by Euclidian length	1.1	0.0097	0.009	6.83E-03	0.422	2.53E-25
GR.TL	Average relative root growth rate: Root growth rate divided by total length at the earlier time point	0.673	0.0796	0.011	1.20E-03	0.393	2.69E-24
BIO1	Annual Mean Temperature (ºC x 10)	98.1	12.8	n/a	n/a	0.066	3.22E-40
BIO2	Mean Diurnal Range (Mean of monthly (max temp - min temp))	107	7.65	n/a	n/a	0.073	1.02E-40
BIO3	Isothermality (BIO2/BIO7) (x 100)	28.9	1.8	n/a	n/a	0.361	4.91E-39
BIO4	Temperature Seasonality (standard deviation x 100)	9169	483	n/a	n/a	0.383	4.68E-47
BIO5	Max Temperature of Warmest Month (°C x 10)	283	10.1	n/a	n/a	0.152	3.78E-40
BIO6	Min Temperature of Coldest Month (°C x 10)	-80.9	18	n/a	n/a	0.275	4.79E-42
BIO7	Temperature Annual Range (BIO5-BIO6) (ºC x 10)	364	17.5	n/a	n/a	0.239	6.31E-42
BIO8	Mean Temperature of Wettest Quarter (ºC x 10)	176	55.1	n/a	n/a	0.016	3.58E-43

BIO9	Mean Temperature of Driest Quarter (ºC x 10)	-7.11	48.7	n/a	n/a	0.000	3.58E-43
BIO10	Mean Temperature of Warmest Quarter (ºC x 10)	213	10.8	n/a	n/a	0.205	3.33E-40
BIO11	Mean Temperature of Coldest Quarter (ºC x 10)	-24.1	18.2	n/a	n/a	0.270	1.71E-41
BIO12	Annual Precipitation (mm)	990	109	n/a	n/a	0.219	3.94E-44
BIO13	Precipitation of Wettest Month (mm)	103	6.72	n/a	n/a	0.206	1.53E-40
BIO14	Precipitation of Driest Month (mm)	54.1	16.7	n/a	n/a	0.104	1.51E-40
BIO15	Precipitation Seasonality (Coefficient of Variation)	17.8	5.51	n/a	n/a	0.157	8.93E-40
BIO16	Precipitation of Wettest Quarter (mm)	291	19.7	n/a	n/a	0.269	1.55E-42
BIO17	Precipitation of Driest Quarter (mm)	191	44.8	n/a	n/a	0.084	3.67E-42
BIO18	Precipitation of Warmest Quarter (mm)	277	25.2	n/a	n/a	0.342	7.42E-44
BIO19	Precipitation of Coldest Quarter (mm)	197	47	n/a	n/a	0.022	2.68E-42

Table S5. SNP hits from association analyses and several descriptors.

SNP hits significant at the 5% level after permutation correction are shown. Additionally, if raw p-values pass a double Bonferroni threshold of 0.01% are marked with a "tick". (Abbreviations: nonsyn. and syn., nonsynonymous and synonymous changes; regular one-letter abbreviation was used for amino acid changes)

Trait	Chromosome	Position Ancestral Derived	Effect	Effect standard error Sample size	p - value raw	p- value false discovery rate	p- value permutation corrected	Allele frequency	Allele frequency in modern set	Oldest herbarium individual	Longitude	Latitude Substitution type AA change	Gene	Biochemical effect (Grantham score)	Significant permutation	Significant double Bonferroni	D
dirEquivalent	1	958948 G T	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 nonsyn A->P	AT1G03810	27	✓		53
gravitropicScore	1	9925177 C T	0.033	0.010 63	7.10E-04	0.0651	0.016	0.078	0.092	1952	40.9	-82.3 interg.			✓		1
bio18	1	10187610 T C	6.830	1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 interg.			✓		52
GR_rootLength	1	12638692 C T	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.087	0.105	1952	40.9	-81.3 interg.			✓	✓	13
GR_shootArea	1	12638692 C T	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.087	0.105	1952	40.9	-81.3 interg.			✓	✓	13
GR_rootLength	1	13652509 C A	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.093	0.107	1952	40.9	-82.9 interg.			✓	1	12
GR_shootArea	1	13652509 C A	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.093	0.107	1952	40.9	-82.9 interg.			✓	✓	12
bio18	1	13904611 C T	6.570	1.756 90	1.83E-04	0.0124	0.016	0.217	0.237	1922	41.7	-85.3 interg.			✓		49
bio18	1	13994958 G A	6.830	1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 tranposon	AT1G36933		✓		49
bio18	1	17408807 C T	6.830	1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 interg.			✓		48
dirEquivalent	1	19024876 C T	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.19	0.23	1922	41.7	-85.3 interg.			✓		47
GR_shootArea	1	20324050 G A	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.087	0.105	1952	40.9	-82.9 interg.	AT1G54440		✓	✓	11
GR_rootLength	1	20324050 G A	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.087	0.105	1952	40.9	-82.9 interg.	AT1G54440		✓	✓	11
bio18	1	23648407 A C	6.830	1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 nonsynY->S	AT1G63740	144	✓		46
dirEquivalent	1	26052913 A T	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.185	0.224	1922	41.7	-85.3 interg.			✓		45
GR_shootArea	1	29696198 G A	-121.000	33.911 63	3.68E-04	0.0096	0.016	0.278	0.329	1922	41.5	-84.9 interg.			✓		42

bio16	1	29696198 G A	5.250	1.377 94	1.39E-04	0.0632	0.016	0.278	0.329	1922	41.5	-84.9 interg.		~	1	42
bio18	1	29696198 G A	6.340	1.569 94	5.36E-05	0.0124	0.004	0.278	0.329	1922	41.5	-84.9 interg.		~	1	42
GR_rootLength	1	30015381 T A	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.087	0.105	1952	40.9	-82.9 interg.		~	′ 🗸	[′] 10
GR_shootArea	1	30015381 T A	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.087	0.105	1952	40.9	-82.9 interg.		~	′ ✓	[′] 10
GR_rootLength	1	30143319 G A	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.088	0.105	1952	40.9	-82.9 interg.		~	′ √	[′] 9
GR_shootArea	1	30143319 G A	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.088	0.105	1952	40.9	-82.9 interg.		~	′ √	[′] 9
dirEquivalent	2	358395 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 syn. V->V	AT2G01820	~	′ ✓	[′] 43
dirEquivalent	2	585918 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 syn. G->G	AT2G02220	~	′ ✓	[′] 42
dirEquivalent	2	1093203 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 interg.		~	′ ✓	[′] 41
dirEquivalent	2	2176891 T C	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 interg.		~	′ ✓	<i>4</i> 0
GR_rootLength	2	3174832 T A	6.340	1.869 63	6.97E-04	0.017	0.017	0.529	0.566	1879	41.3	-84.3 interg.		~	1	0
TotLen.EucLen	2	5285907 C A	-0.006	0.002 63	3.05E-04	0.0241	0.037	0.162	0.194	1922	41.5	-85 interg.		~	′ ✓	[′] 39
dirEquivalent	2	5285907 C A	-0.019	0.005 63	2.64E-05	0.0032	0.001	0.162	0.194	1922	41.5	-85 interg.		~	′ 🗸	[′] 39
dirEquivalent	2	6034545 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 syn. S->S	AT2G14247	~	′ ✓	[′] 38
dirEquivalent	2	7047529 G T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 nonsynP->A	AT2G16270	27 🗸	′ ✓	´ 37
dirEquivalent	2	7186220 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 intron	AT2G16580	~	′ ✓	[′] 36
dirEquivalent	2	10369545 T C	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 interg.		~	′ ✓	´ 35
dirEquivalent	2	10495275 A C	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.196	0.237	1922	41.7	-85.3 intron	AT2G24680	~	′ ✓	[′] 34
dirEquivalent	2	11346211 C A	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 interg.		~	1	33
dirEquivalent	2	12415084 T A	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 intron	AT2G28900	~	•	32
dirEquivalent	2	12876361 A C	-0.015	0.004 63	1.56E-04	0.0041	0.006	0.262	0.29	1922	41.7	-84.6 interg.		~	1	31
gravitropicScore	2	12876361 A C	-0.021	0.006 63	1.08E-03	0.0651	0.027	0.262	0.29	1922	41.7	-84.6 interg.		~	1	31
bio13	2	14417366 A G	3.990	0.959 64	3.22E-05	0.0147	0.004	0.077	0	1890	39.5	-77.9 interg.		~	•	1
dirEquivalent	2	15278350 A G	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 interg.		~	•	30
GR_shootArea	2	16039488 T G	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.087	0.105	1952	40.9	-82.9 3' UTR	AT2G38290	~	′ ✓	[′] 8
GR_rootLength	2	16039488 T G	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.087	0.105	1952	40.9	-82.9 3' UTR	AT2G38290	~	′ √	[′] 8
GR_rootLength	2	16247290 G T	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.088	0.105	1952	40.9	-82.9 nonsynA->G	AT2G38910	60 🗸	′ √	[′] 7
GR_shootArea	2	16247290 G T	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.088	0.105	1952	40.9	-82.9 nonsynA->G	AT2G38910	60 🗸	′ √	[′] 7
dirEquivalent	2	16333662 G A	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 nonsyn A->G	AT2G39160	60 🗸	•	29
dirEquivalent	3	2500258 C A	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 syn. K->K	AT3G07830	~	′ ✓	[′] 28
dirEquivalent	3	3154804 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 interg.		~	′ ✓	[′] 27
dirEquivalent	3	3629794 C T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 intron	AT3G11530	~	′ ✓	[′] 26
dirEquivalent	3	4269626 G T	-0.016	0.004 63	1.15E-04	0.0032	0.006	0.194	0.237	1922	41.7	-85.3 5' UTR	AT3G13229	~	′ ✓	[′] 25
GR_shootArea	3	8873116 C T	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.097	0.118	1952	40.9	-81.9 interg.		~	′ 🗸	<i>6</i>
GR_rootLength	3	8873116 C T	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.097	0.118	1952	40.9	-81.9 interg.		~	′ √	[′] 6

GR_rootLength	3	11259214 A	T -12.1	0 3.164 63	1.33E-04	0.0037	0.003	0.088	0.105	1952	40.9	-82.9 interg.			✓	1	5
GR_shootArea	3	11259214 A	T -231.0	00 53.774 63	1.75E-05	0.0005	0.001	0.088	0.105	1952	40.9	-82.9 interg.		,	✓	✓	5
bio8	3	11873293 A	G 37.8	0 8.736 65	1.52E-05	0.0069	0.006	0.939	1	1890	41.8	-83.7 tranposon	AT3G30219	,	✓		0
GR_rootLength	3	15050751 G	A -12.1	00 3.164 63	1.33E-04	0.0037	0.003	0.108	0.105	1888	40.2	-82.5 interg.		,	✓	✓	4
GR_shootArea	3	15050751 G	A -231.0	00 53.774 63	1.75E-05	0.0005	0.001	0.108	0.105	1888	40.2	-82.5 interg.		,	✓	✓	4
dirEquivalent	3	17164638 C	A -0.0	L4 0.004 63	5.30E-04	0.0052	0.018	0.19	0.227	1922	41.7	-85.3 interg.		,	✓		24
bio18	4	279210 T	G 6.8	30 1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 interg.		,	✓		22
bio11	4	1732480 T	A -5.5	50 1.564 79	3.89E-04	0.0195	0.045	0.063	0.068	2002	41	-87.5 interg.		,	✓		2
bio4	4	1732480 T	A 224.0	00 63.967 79	4.67E-04	0.0128	0.044	0.063	0.068	2002	41	-87.5 interg.		,	✓		2
dirEquivalent	4	3355152 C	G -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		21
bio18	4	3355152 C	G 6.8	50 1.944 ##	4.25E-04	0.0124	0.035	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		21
dirEquivalent	4	3355946 G	C -0.0	.4 0.004 63	4.45E-04	0.0052	0.016	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		20
bio18	4	3355946 G	C 6.8	0 1.944 ##	4.25E-04	0.0124	0.035	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		20
dirEquivalent	4	4228138 A	G -0.0	.4 0.004 63	4.45E-04	0.0052	0.016	0.196	0.24	1922	41.7	-85.3 tranposon	AT4G07440	,	✓		19
bio18	4	4228138 A	G 6.8	30 1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 tranposon	AT4G07440	,	✓		19
dirEquivalent	4	9046942 G	C -0.0	.4 0.004 63	4.45E-04	0.0052	0.016	0.204	0.25	1922	41.7	-85.4 nonsyn H->Q	AT4G15960	24	✓		18
bio18	4	9046942 G	C 6.8	50 1.944 ##	4.25E-04	0.0124	0.035	0.204	0.25	1922	41.7	-85.4 nonsyn H->Q	AT4G15960	24	✓		18
dirEquivalent	4	11948961 T	A -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.198	0.25	1952	41.7	-85.3 interg.		,	✓		17
dirEquivalent	4	12365323 C	т -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		16
bio18	4	12365323 C	T 6.8	50 1.944 ##	4.25E-04	0.0124	0.035	0.204	0.25	1922	41.7	-85.4 interg.		,	✓		16
dirEquivalent	4	15646341 C	A -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.206	0.25	1922	41.7	-85.4 syn. E->E	AT4G32410	,	✓		15
bio18	4	15646341 C	A 6.7	1.936 99	5.14E-04	0.0124	0.042	0.206	0.25	1922	41.7	-85.4 syn. E->E	AT4G32410	,	✓		15
dirEquivalent	4	15845001 A	т -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.194	0.25	1922	41.8	-85.9 3' UTR	AT4G32840	,	✓		14
dirEquivalent	4	18249171 T	A -0.0	L4 0.004 63	4.45E-04	0.0052	0.016	0.274	0.328	1922	41.8	-85.9 interg.		,	✓		13
bio18	4	18249171 T	A 6.9	LO 2.005 71	5.62E-04	0.0124	0.047	0.274	0.328	1922	41.8	-85.9 interg.		,	✓		13
bio18	5	4245213 A	T 6.8	30 1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 syn. I->I	AT5G13260	,	✓		12
bio18	5	4500202 G	A 6.8	30 1.987 99	5.83E-04	0.0124	0.047	0.196	0.24	1922	41.7	-85.3 nonsyn A->G	AT5G13950	60	✓		11
dirEquivalent	5	4797923 A	T -0.0	.4 0.004 63	5.30E-04	0.0052	0.018	0.188	0.227	1922	41.7	-85.3 tranposon	AT5G14830	,	✓		10
dirEquivalent	5	4797976 G	A -0.0	.4 0.004 63	5.30E-04	0.0052	0.018	0.257	0.293	1922	41.7	-85.3 tranposon	AT5G14830	,	✓		10
dirEquivalent	5	4798526 A	G -0.0	.4 0.004 63	5.30E-04	0.0052	0.018	0.339	0.362	1922	41.7	-85.3 interg.		,	✓		9
gravitropicScore	5	6508329 A	G -0.0	0.006 63	5.20E-04	0.0651	0.008	0.35	0.447	1922	42	-85 nonsyn C->W	AT5G19330	215	✓		0
dirEquivalent	5	11090365 T	A -0.0	L4 0.004 63	5.30E-04	0.0052	0.018	0.186	0.224	1922	41.7	-85.3 TE	AT5G29037	,	✓		4
dirEquivalent	5	12312975 C	G -0.0	.4 0.004 63	5.30E-04	0.0052	0.018	0.185	0.224	1922	41.7	-85.3 TE	AT5G32630	,	✓		3
dirEquivalent	5	12358159 C	т -0.0	L4 0.004 63	5.30E-04	0.0052	0.018	0.186	0.224	1922	41.7	-85.3 tranposon	AT5G32825	,	✓		2
dirEquivalent	5	12409027 G	A -0.0	L4 0.004 63	5.30E-04	0.0052	0.018	0.185	0.224	1922	41.7	-85.3 interg.		,	✓		1

GR_rootLength	5	16024197 A T	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.098	0.118	1952	40.9	-81.9 intron	AT5G40020	✓	✓	2
GR_shootArea	5	16024197 A T	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.098	0.118	1952	40.9	-81.9 intron	AT5G40020	✓	✓	2
GR_shootArea	5	16109431 G A	-231.000	53.774 63	1.75E-05	0.0005	0.001	0.865	0.877	1993	42.2	-84.4 interg.		✓	✓	1
GR_rootLength	5	16109431 G A	-12.100	3.164 63	1.33E-04	0.0037	0.003	0.865	0.877	1993	42.2	-84.4 interg.		✓	✓	1
dirEquivalent	5	19099082 G C	-0.014	0.004 63	5.30E-04	0.0052	0.018	0.186	0.227	1922	41.7	-85.3 interg.		✓		0
GR_rootLength	5	20388107 A T	-10.700	3.164 63	6.94E-04	0.017	0.017	0.099	0.12	2002	41	-86.6 interg.		✓		0

Graphic Table S7

GR_rootLength







dirEquivalent

lambda 1.355



Expected (-logP)





lambda 0.054



stdDevXY



meanRootWidth



rootWidth20

lambda 0.002







lambda 0.059



lambda 0.089



rootWidth60



rootWidth80

lambda 0.068







lambda 0.04



lambda 0.059



gravitropicDir



gravitropicScore

lambda 0.478



TotLen.EucLen



lambda 0.685



GR.TL







0.0

0.0

0.5

1.5

1.0

Expected (-logP)

0

۔ 100

110

115

FT_V2

lambda 1.003











B_V0





B_V1

lambda 0.192



Expected (-logP)





B_V3



lambda 0.532





fecundity





lambda 0.074



0.0

0.0

0.2

0.4

Expected (-logP)

0.8

1.0

1.2

0.6

lambda 0.138



bio3

110

115

120

125

0

ſ 95

100





10

ß

0

Г

-100

-60

-40

-80

lambda 0.606



1.0

0.5

0.0

0.0

0.5

Expected (-logP)

1.5

1.0

2.0



lambda 0.283



0.5

0.0

0.0

0.5

1.0





Expected (-logP)

1.5

2.0

2.5

3.0






lambda 0.114



0.0

1300

0.0

0.5

900 1000 1100 1200

0

800



1.5

2.0

1.0

bio13

10

15

20

25

0.0

lambda 0.188



Expected (-logP)

1.0

1.5

0.5

bio16

240

260

280

300

320

lambda 0.221



0.0

0.5

1.0

Expected (-logP)

1.5

2.0

2.5

3.0



Expected (-logP)

bio19

lambda 0.032