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Young inversion with multiple linked QTLs under selection in a hybrid zone

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AUTHOR CONTRIBUTIONS

CRL and TMO conceived these efforts. CRL, TMO, YY, DK, JZ, and JLG designed the study. TM and ML performed chromosome painting. CRL, KVSKP, NP, HNH, DK, YY, JT, WG, JZ, and JJ worked on molecular biology and sequencing. SF, UH, JWJ, JS, DSR, RAW, and KB; and JLG, DK, YY, JZ, JT, WG, and RAW planned and analyzed genomic and physical mapping experiments, respectively. CRL, BW, JM, UH, JLG, and TMO performed bioinformatic and evolutionary analyses. KG, CRL, and NP performed experiments with plants. CRL, BW, JM, MES, and TMO drafted the manuscript. All authors read, revised, and approved the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

DATA AVAILABILITY

The assembly and annotation of the *Boechera stricta* genome are available at https://phytozome.jgi.doe.gov/pz/portal.html#!info?alias=Org_Bstricta and have been deposited under GenBank accession number MLHT00000000. The BAC end sequences have been deposited under GenBank dbGSS accession numbers KS412618 – KS448200. The short reads of *B. stricta* (LTM, SAD12 and recombinant inbred lines) have been deposited under GenBank accession numbers SRP078672, SRP048481 and SRP079414, respectively. The short reads from the RNA-seq transcriptome have been deposited under GenBank accession numbers SRX1971488. The short reads of outgroup *Boechera holboellii* (= *B. retrofracta*) have been deposited under GenBank accession number SRP078889, and the short reads of genotyping-by-sequencing data of *B. stricta* have been deposited under GenBank accession numbers SRP075905 and SRP075997. All SNPs used in population genetic analyses have been deposited into dbSNP under accession numbers ss2136554982–ss2136641742. All other data are available in the Dryad Data Archive XXX.

CODE AVAILABILITY

All code is available in the Dryad Data Archive.

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Abstract

Fixed chromosomal inversions can reduce gene flow and promote speciation in two ways: by suppressing recombination and by carrying locally favored alleles at multiple loci. However, it is unknown whether favored mutations slowly accumulate on older inversions or if young inversions spread because they capture preexisting adaptive Quantitative Trait Loci (QTLs). By genetic mapping, chromosome painting and genome sequencing we have identified a major inversion controlling ecologically important traits in *Boechnera stricta*. The inversion arose since the last glaciation and subsequently reached local high frequency in a hybrid speciation zone. Furthermore, the inversion shows signs of positive directional selection. To test whether the inversion could have captured existing, linked QTLs, we crossed standard, collinear haplotypes from the hybrid zone and found multiple linked phenology QTLs within the inversion region. These findings provide the first direct evidence that linked, locally adapted QTLs may be captured by young inversions during incipient speciation.

Chromosome inversions play an important role in local adaptation and speciation^{1,2}, and selectively important inversions have been identified in many species^{3,4}. Selection due to different environmental factors or stages in the life cycle¹ may favor inversions carrying locally adapted alleles at several loci. In addition, established inversions are predicted to accumulate selectively important genetic differences, which may contribute to reproductive isolation during speciation¹.

Although few studies have identified the actual loci that influence selection on inversions^{2,4,5}, rearrangements may be favored due to gene alterations near breakpoints⁶, chromatin changes⁷, or combinations of advantageous, coadapted alleles⁸. Inversions suppress recombination, so locally advantageous alleles may segregate together, causing higher fitness than recombinant haplotypes⁹. Most evolutionary studies have focused on widespread, older inversions, so we have little knowledge of the evolutionary processes that guide their initial increase in frequency. For example, do inversions drift to higher frequency, and then acquire new advantageous mutations after they are common? Or are multiple linked, advantageous alleles captured in a new inversion, allowing them to spread together? Analysis of younger inversions may elucidate the evolutionary forces controlling the initial spread of chromosome inversions, which therefore influence their role in adaptation and speciation^{4,8}.

Related species often differ for chromosome inversions that carry locally-favored alleles at multiple loci^{10,11}. A key distinction among models for the evolution of inversions is whether early frequency increase is due to genetic drift or natural selection. Genetic drift might predominate initially, with subsequent accumulation of advantageous variants¹².

Alternatively, the Kirkpatrick-Barton (“KB”) model⁹ argues that linked, locally adapted alleles exist first, and subsequently are captured within a new, selectively-favored inversion¹³. In this “inversion-late” evolutionary sequence^{1,5}, linked QTLs, similar to the ancestral haplotype that gave rise to the inversion, may still exist in non-inverted genotypes⁹. Here, we test these predictions of the KB model. First, we introduce ecologically-diverged subspecies of *Boechera stricta*. Next, we examine a young inversion to infer the selective forces controlling its early increase in frequency. Finally, we cross collinear, standard genotypes from the hybrid zone to ask whether old, linked QTLs can be found within the inversion region.

Young Inversion in a hybrid zone

Previously, we showed that *Boechera stricta* (a close relative of *Arabidopsis*¹⁴) has two ecologically differentiated subspecies (“EAST” and “WEST”)^{15,16}, which form a contact zone across >200 km in the Northern Rocky Mountains. Throughout our study area, local water availability is the best predictor of habitat divergence between these subspecies¹⁵, with WEST genotypes growing in sites with more constant and abundant water supply. In and near the hybrid zone, the EAST and WEST subspecies show significant ecological differentiation across local environmental gradients, and the geographic distribution of the inversion falls within the typical range of hybrid zone habitats. The WEST subspecies has significantly faster growth rate, larger leaf area, less succulent leaves, delayed reproductive time, and longer flowering duration¹⁶, and WEST × EAST crosses found QTLs for flowering traits, leaf number, defensive chemistry, herbivore resistance, cold tolerance, overwinter survival, fecundity, and lifetime fitness^{17–20}. In addition, Q_{ST} - F_{ST} analysis showed that phenology and some morphological traits have experienced divergent selection between subspecies¹⁶.

In our initial cross¹⁷, genetic mapping in WEST × EAST recombinant inbred lines (RILs) identified a region of suppressed recombination (Supplementary Fig. 1) on linkage group 1 (LG1). Here, we combine short-read sequencing, end sequencing of Bacterial Artificial Chromosomes (BACs), linkage mapping, and physical mapping by Whole Genome Profiling to assemble a high quality genomic sequence (Supplementary Fig. 2), which enabled evolutionary analysis of the inversion. Chromosome painting (Fig. 1; Supplementary Fig. 3) cytogenetically verified the presence of an 8.4 Mb paracentric inversion in the WEST genotype (*Bs1*; Fig. 2a; Supplementary Fig. 4), spanning ~31 cM (~4% of the genome, the “inversion region”, Supplementary Table 1). We developed primers to score the inversion using PCR (Fig. 2b), showing that the common, non-inverted allele (*Bs1-std*) has the ancestral orientation found in closely-related *Capsella* and *Arabidopsis lyrata*²¹, and is found in >200 EAST and WEST populations across the species range (Fig. 3a; Dryad Data Archive). The derived, inverted allele (*Bs1-inv*) was identified in a cluster of WEST and hybrid populations in the contact zone (Fig. 3b), where it has risen to high frequency. Genotypes carrying the inversion span a narrow geographic range (~14 km, the “inversion

zone”), where pollen and geological analyses^{22,23} suggest that conditions suitable for *B. stricta* existed during the Last Glacial Maximum ~21 kya. SNPs in the inversion region show that the most similar *Bsil-inv* and *Bsil-std* haplotypes (Supplementary Fig. 5) are located only 1.7 km apart (Fig. 3b, left). Using measured mutation rates²⁴ and assuming two years per generation, the inversion dates to ~2,100 – 8,800 years (Methods). Because these widely-distributed subspecies are ecologically diverged¹⁶, while the *Bsil* inversion originated very recently, this polymorphism is compatible with the inversion-late KB model.

Previous analysis of a WEST-*inv* × EAST-*std* mapping population grown in the inversion zone found many QTLs controlling components of fitness²⁵ and strong selection on flowering time¹⁷, influenced by recent climate warming²⁶. Native WEST *Bsil-inv* alleles had higher fitness than EAST *Bsil-std* alleles²⁵. The inversion (spanning ~4% of the genome) explains 7.5% of heritable variation for lifetime fecundity, and inversion homozygotes differ by 0.56 genetic standard deviations for this trait ($t = 3.64$, $df = 165$, $P = 0.0004$). Here, we focused on effects of the inversion by examining flowering time in two F7 Near-Isogenic Line (NIL) families from this cross. We found no segregation distortion or deviation from Mendelian ratios (Supplementary Table 2) in either family, and the inversion had significant effects on flowering time in these NIL families in the greenhouse ($F_{2, 282} = 21.81$, $P < 0.001$), where the *Bsil-std* haplotype flowered ~2.0 days faster than the *Bsil-inv* haplotype (Supplementary Fig. 6). Thus, the inversion controls 40% of the average 5.0 day difference in flowering time between subspecies (Supplementary Table 2d). While this pedigree cannot resolve QTLs within the inversion, this cross shows that the inversion has significant effects on an ecologically important trait that experiences natural selection¹⁷. Gene(s) within the *Bsil* inversion control flowering in the field, and thus may contribute to reproductive isolation during speciation²⁷.

To test for phenotypic effects of the inversion or transcription changes near the breakpoints, we compared closely-related inverted and standard haplotypes from the inversion zone, using a sympatric WEST-*inv* × WEST-*std* F2 cross (Fig. 3b, left). We found high seed germination (>99%), and no *Bsil* segregation distortion or deviation from Mendelian ratios (Supplementary Table 3). MANOVA showed a significant effect of *Bsil* on a suite of phenological and morphological traits ($P = 0.005$ by permutation test, $R^2 \approx 4\%$; Supplementary Table 4), so we analyzed individual traits by ANOVA. In this cross, flowering time is delayed by 0.9 days ($P = 0.030$ by permutation test) in *Bsil-inv* homozygotes (accounting for 18% of the difference between subspecies), which may increase reproductive isolation from the early-flowering EAST genotypes¹⁶. Finally, we tested for functional changes attributable to the inversion breakpoints; these breakpoints do not disrupt existing or create new open reading frames (Supplementary Fig. 7). Using replicated F3 homozygotes from this cross, we found no significant expression differences between inversion and standard haplotypes for any of the seven genes flanking the inversion breakpoints (Supplementary Table 5), although power to detect subtle, quantitative differences is limited. Thus, this comparison of WEST-*inv* vs. WEST-*std* *Bsil* haplotypes found genetic differences for ecologically-important traits, but little evidence for functional changes at the inversion breakpoints.

Evidence for Natural Selection

To test whether selection had favored this inversion, we analyzed patterns of population genetic variation (Fig. 4; Supplementary Fig. 8). *Bsil-inv* genotypes show lower LG1 polymorphism, lower Tajima's *D*, and more linkage disequilibrium (LD) than *Bsil-std* individuals. These patterns are compatible both with neutral drift in this partially inbreeding species, or with a selective sweep on the inversion²⁸. In contrast, young, derived mutations at high frequency suggest positive directional selection²⁹, so we asked whether the frequency of *Bsil-inv* is typical of frequencies of all private derived SNP alleles in the same population (the 54 similar genotypes in the left portion of Supplementary Fig. 9c and the left portion of Fig. 3b). The inversion is at relatively high frequency (0.63) in this group, but it is not fixed. We found 2,416 SNPs that, like the inversion, are confined to this population. Among these, only 63 SNPs (2.6%) have derived allele frequencies greater than the frequency of the *Bsil-inv* inversion. Hence, the frequency of the *Bsil-inv* inversion allele is higher than 97.4% of comparable derived alleles in this population – the inversion is a high frequency outlier, supporting the hypothesis of positive directional selection.

QTLs within the Inversion

During speciation, coadapted gene complexes within inversions might reduce gene flow if they preserve favorable combinations of alleles at multiple loci, reducing the frequency of disadvantageous allelic combinations. Although inversions can be engineered for proof of function in some organisms³⁰, this is infeasible in *Boechera*. However, the KB model predicts that relatives of the ancestral haplotype that gave rise to the inversion might still exist as non-inverted genotypes nearby. To test for such linked QTLs, we crossed collinear WEST-*std* × EAST-*std* genotypes from the contact zone and tested for QTLs within the inversion region, using freely-recombining F3:F4 families. We found several multivariate QTLs altering ecologically-important phenology and development traits (Fig. 5a). To clarify differences among these QTLs, we examined differences in their pleiotropic effects (Fig. 5b), using Discriminant Function Analysis (DFA) to find the axis of greatest divergence between the multivariate trait means at each QTL peak. Each DFA axis quantifies the direction of pleiotropy that controls effects of a locus, and QTLs influencing these composite traits were mapped across the inversion region. We found several distinct QTL peaks, which show divergent patterns of pleiotropy among these linked loci. Finally, we estimated the time of divergence between these parental genotypes (Fig. 5c). These results show that the QTL haplotypes diverged at least 50 – 100 Kya, and are therefore much older than the inversion, which arose less than 10 Kya.

In summary, population genetic evidence for a selective sweep corresponds with our ecological findings: the inversion affects multiple ecologically-important traits, including flowering differences that are expected to increase reproductive isolation between subspecies. The inversion occurs in a hybrid zone, and this genomic region contains multiple ecologically important QTLs, as predicted by the KB model. However, beyond the potential advantage conferred by recombination suppression⁹, closely related WEST-*inv* and WEST-*std* haplotypes show divergent phenotypic effects, which might contribute to selection in the hybrid zone. Our analysis of a young inversion is compatible with evolutionary predictions

that linked, locally adapted QTLs may be captured by new inversions and contribute to local adaptation or incipient speciation⁹.

METHODS

Study Area

The inversion was originally detected²¹ at the Lost Trail Meadow site (~2,500m elevation) about four km NW from Lost Trail Pass on the Montana-Idaho border. Pollen and geological analyses^{22,23} show that climate and vegetation in this area was strongly influenced by Pleistocene climatic changes, although the site apparently was unglaciated during the Last Glacial Maximum (LGM) about 21,000 years ago. Inferred plant communities^{22,23} suggest conditions suitable for *Boechnera stricta* may have existed at or near our study site during the LGM. We searched extensively in nearby *B. stricta* habitat; The Dryad Data Archive contains information on 122 genotypes from the inversion zone (Fig. 3), and 83 comparison genotypes from the across the species range.

Experimental Pedigrees

Details on experimental crosses and genotype locations are in the Dryad Data Archive.

DNA extraction for genomic sequencing

Seed sterilization—Seeds of *Boechnera stricta* ecotypes (LTM and SAD12) were initially surface sterilized with ethanol (200 proof) for 2 min followed by treatment with 15% sodium hypochlorite solution (with a few drops of Tween-20) for 25 min on a rotatory shaker at room temperature. Surface sterilized seeds were thoroughly washed multiple times with sterile water and were suspended in 0.1% agar. Seeds were kept for stratification in a cold room at 4°C for 4 d under dark conditions.

Raising of aseptic seedling cultures—To raise aseptic seedlings of LTM and SAD12 ecotypes, surface sterilized and stratified seedlings were inoculated in 250 ml flasks containing 50 ml of sterile ½ MS liquid medium (pH 5.7) with 0.5 g/L MES and 2% sucrose. About 70–80 seeds were inoculated per flask. To obtain etiolated seedlings, the flasks were covered with aluminum foil and kept on a bench top rotatory shaker at 110 rpm for 20 d at room temperature.

Extraction of genomic DNA from etiolated seedlings—Freshly grown 20 d old etiolated seedlings of both ecotypes were used for extraction of nuclear DNA. Nuclei isolation was performed essentially according to Prasad *et al*¹⁸ with slight modifications, as it allowed isolation of clean high molecular size nuclear DNA with minimal contamination from organelle DNA. Briefly, about 10 flasks of etiolated seedling cultures were removed from the growing medium and thoroughly washed with ice cold water and placed in ice cold ethyl ether for 3 min. Subsequently the seedlings were washed five times with ice cold TE buffer (pH 7.0). The plant material was quickly blotted on sterile filter paper and homogenized in MEB buffer using a commercial blender. The homogenate was filtered through four layers of cheesecloth followed by two layers of mira cloth. Triton-X-100 was added to the filtrate at 0.5% concentration and centrifuged to collect the pellet. The pellet

was suspended in MPDB solution and gently layered on 37.5% percoll made with MPDB. Nuclei were collected as a pellet after centrifugation, and the pellet was washed by resuspending it in 20 ml of MPBD followed by centrifugation. The nuclei pellet was again suspended in a MPDB solution and high molecular weight nuclear DNA was extracted using QIAGEN genomic tip 100/G protocol as per the manufacturer's instructions, with some modifications. The pellet consisting of nuclei was resuspended in the lysis buffer and incubated at 40°C for 15 min. RNase A and T1 were added to the suspension and further incubated at 37°C for additional 30 min. Subsequently, Proteinase K was added to the suspension at final concentration of 150 µg per ml and incubated at 3h at 45°C with gentle shaking. The suspension was cleared by centrifugation at 8,000 rpm for 15 min. The cleared suspension was added to the pre-equilibrated G-100 column and further purification was performed as per the protocol provided by the manufacturer. The eluate consisting of genomic DNA was further precipitated using isopropanol, washed with 70% ethanol and subsequently suspended in the TE (pH 8.0) buffer.

Total RNA isolation

To collect RNA from roots, the LTM-genotype seeds were surface sterilized with 12% bleach and grown under aseptic conditions on a Whatman filter paper bridges in glass tissue culture tubes containing 1/2 strength MS media with 1% sucrose. Each tube was inoculated with 2 to 3 seeds and allowed to grow. The roots obtained from the multiple seedlings were pooled and RNA was isolated with plant RNAeasy kit from Qiagen. For other tissues, we extracted RNA from soil-grown plants at several stages: 15 day old seedlings, rosette leaves from juvenile plants, cauline leaves from plants aged 2 – 3 months. Finally, from 7 month-old plants we extracted RNA from inflorescences, stems, flowers, and siliques.

For isolation of total RNA from various tissues at different developmental stages, the tissues were snap frozen in liquid nitrogen and stored at –80°C. The frozen tissue was homogenized with mortar and pestle using liquid nitrogen. About 150 mg of homogenized tissue was used for isolation. Total RNA was isolated using a Plant RNA-easy kit (Qiagen, USA). Each of the independent total RNA preparations was subjected to DNase treatment to remove traces of contaminating genomic DNA using RNase-free DNase (Promega, USA). For each tissue type, three independent isolations were pooled together and used for further analysis.

Sequencing, Assembly, and Annotation

Eighteen genomic DNA libraries and one transcriptome cDNA library were prepared (Dryad Data Archive) for sequencing using Illumina and Roche 454. The genome assembly of *B. stricta* employed Meraculous build May 2013³¹, as described on Phytozome: (https://phytozome.jgi.doe.gov/pz/portal.html#!info?alias=Org_Bstricta),

Contigs were assembled with a k-mer size of 51 and a minimum depth of 8 from 2 × 150 Illumina HiSeq reads from an unamplified 250 bp whole genome shotgun fragment library. Several discrete rounds of scaffolding were performed with libraries containing inserts ranging in size from 250bp to 40kb sequenced with various Illumina technologies. At each round of scaffolding the minimum pairing threshold was chosen by exhaustive optimization of N50 scaffold length. Assembly

gaps were closed with 2×150 Illumina HiSeq reads. The resulting assembly is comprised of 171.9 MB of contigs in 196.5 MB of scaffold.

The genome assembly was soft-masked to highlight consensus repeat family sequences predicted *de novo* by RepeatModeler. 37,384 RNAseq transcript assemblies were constructed from 118.7 million 2×150 paired-end Illumina RNAseq reads using PERTRAN (Shengqiang Shu, Joint Genome Institute in-house pipeline, <http://jgi.doe.gov/wp-content/uploads/2013/11/CSHL-PERTRAN-Shengqiang-Shu-FINAL.pdf>). Loci were determined by BLAT transcript assembly alignments, BLASTX alignments of proteins from arabi (*Arabidopsis thaliana*), rice, soybean, grape, maize, or *Chlamydomonas reinhardtii* genomes, and/or BLASTX alignments of UniProtKB/Swiss-Prot to the *B. stricta* genome. Gene models were predicated by homology-based predictors, mainly FGENESH+ and GenomeScan.

The best scored predictions for each locus were selected using multiple positive factors including EST and protein support, and one negative factor: overlap with repeats. The selected gene predictions were improved by PASA. Improvement includes adding UTRs, splicing correction, and adding alternative transcripts. PASA-improved gene model proteins were subject to protein homology analysis to above mentioned proteomes to obtain Cscore and protein coverage. Cscore is a protein BLASTP score ratio to MBH (mutual best hit) BLASTP score and protein coverage is highest percentage of protein aligned to the best of homologs. PASA-improved transcripts were selected based on Cscore, protein coverage, EST coverage, and its CDS overlapping with repeats. The transcripts were selected if its Cscore is larger than or equal to 0.5 and protein coverage larger than or equal to 0.5, or it has EST coverage, but its CDS overlapping with repeats is less than 20%. For gene models whose CDS overlaps with repeats for more than 20%, its Cscore must be at least 0.9 and homology coverage at least 70% to be selected. The selected gene models were subject to Pfam analysis and gene models whose protein is more than 30% in Pfam TE domains were removed.

There are 1,591 annotated genes in the inversion region (below), of which 408 are found in the QTL regions within the inversion.

Whole Genome Profiling (WGP)—A Hind III BAC library of *B. stricta* (Bs_LBa) was built from fresh leaf tissues from the LTM reference genotype¹⁸. The library contains 18,432 clones, arrayed into 48 – 384 well plates, with an average insert size of 150 Kb and an estimated genome coverage of 11X. The library or clones of interest from it are publicly available at the Arizona Genomics Institute (AGI) web page (<http://www.genome.arizona.edu/orders/>).

A Whole Genome Profiling (WGP) physical map of *B. stricta* was built following the protocol described by van Oeveren *et al.*³². Briefly, BAC library clone plates were arrayed into two and three dimensional pools, BAC DNA was extracted from pooled plates, BAC DNA was digested with EcoRI/MseI and ligated to their respective adapters (P5-EcoRI-tag

barcode and P7-MseI), followed by PCR amplification and sequencing with an Illumina HiSeq 2500.

Resulting sequences were deconvoluted with KeyGene® proprietary scripts (licensed to AGI), to generate the tag file (Dryad Data Archive), which were assembled with FPC v9.4³³. Four test assemblies were performed with a fixed tolerance value = 0 and cutoff values $1e^{-25}$, $1e^{-20}$, $1e^{-15}$ and $1e^{-10}$, to choose the one producing best results, based on the number of contigs and number of clones included in those contigs.

Questionable clones were eliminated from the best project ($1e^{-15}$) and the contigs were merged at a cutoff value of $1e^{-09}$. Singletons (clones that do not assemble with these parameters), were added at reduced stringency ($1e^{-09}$). This new project was subject to manually editing, after linking the tags, BAC end and genome scaffolds (v 1.0) sequences to the physical map³⁴. Using the SyMAP package³⁵, the final edited map was aligned to pseudomolecules of *B. stricta* and *Capsella rubella*³⁶. Synteny with *A. lyrata* was visualized with MUMmer³⁷.

The integration of the WGP physical map with the genetic map and sequence scaffolds (via the WGP tags and BAC end sequencing) showed high concordance among these three genomic resources. WGP enabled regions with low recombination in the linkage map to be ordered and oriented based on their correspondence with the physical map contigs. In addition, we were able to integrate some scaffolds that were not linked in the genetic map, providing more robust pseudomolecules.

Variant discovery

We aligned each genotype read to the *Boechea stricta* LTM hardmasked reference (B.strict_278_hardmasked) with BWA³⁸. We used GATK³⁹ for base quality recalibration, indel realignment, simultaneous SNP and INDEL discovery via HaplotypeCaller and joint genotyping using the default hard filtering parameters as prescribed by GATK Best Practices recommendations³⁹.

Ordering and orienting scaffolds into linkage groups

We created a genetic map based on 159 sixth-generation recombinant inbred lines (RILs) bred from two parent individuals: LTM (the reference individual) and SAD12. We (re-)sequenced LTM to a genomic coverage of ~400X using paired-end Illumina sequencing. These reads were aligned to the reference sequence using *bwa mem*⁴⁰ and *samtools mpileup*⁴¹ to visualize the alignment. We conservatively verified 105,074,494 positions as homozygous by requiring unambiguous alignment of a single base type with a depth ranging from 160–600. Next, we performed a similar analysis on a set of paired-end Illumina reads from SAD12 (depth ~170X) and compiled a catalog of 442,637 discriminatory positions throughout the genome. These markers were homozygous for different bases in LTM and SAD12, allowing identification of ancestry within the RILs. These results imply nucleotide divergence equals 0.000425 between these two accessions.

We initially aligned sequence from 199 barcoded RILs sequenced to modest (~a few X) coverage to the reference sequence and genotyped these at all sites possible within the

discriminatory catalog. During this process we eliminated a number of contaminated RILs, RILs with too little sequence for reliable genotyping, and RILs that appeared to be heterozygous over most of the genome, suggesting that the source DNA had not originated from a single RIL. The scaffolds in the assembly with sizes ≥ 20 kb were binned into 20kb blocks and each block with at least 3 genotypeable sites were genotyped as either LTM, SAD12, or heterozygous, based on the consensus of genotypeable sites. A catalog of breakpoints was constructed, as a list of scaffolds and bins where one or more RILs changed genotype. From this data we identified 9 redundant RILs, after the removal of which we were ultimately left with 159 RILs used in the analyses. Also, we found two misjoins in the original assembly, which identified themselves as being apparently unlinked neighboring bins. These were Scaffold25219, between bin 134 and 137 (removed bins 135,136 and renamed bins 137–139 to Scaffold100000 bins 0–2) and Scaffold19424 between bins 344 and 345 (renamed Scaffold19424 bins 345–391 to Scaffold200000 bins 0–46). On the final data set, 2,664 crossovers were located (16.7 per RIL), and 2.4% of all blocks were genotyped as heterozygous (an average of 3.12% is expected for F6 RILs, with a large variance).

Next, we calculated a matrix of recombination fractions, r , between each pair of bins, defined as the fraction of RILs for which the bins differ in genotype. Bins within each pair with $r < 0.15$ were clustered by a single-linkage approach. (Briefly, all pairs of bins were compared, and pairs with $r < 0.15$ were assigned to the same cluster. Two clusters were merged if any pair of members had $r < 0.15$.) This algorithm resulted in seven distinct linkage groups. The bins in each linkage group were used as markers for input into MSTmap⁴². Finally, individual bins within assembled scaffolds were re-arranged into the order on which they occur on the scaffolds, and in the orientation, if known, consistent with the genetic map. This step is necessary as the size of the 20 kb bins are often smaller than the map resolution, resulting in the bins appearing in random order on the map. During this step, the cumulative physical length of the map (in bases) was also inserted into the map. In addition, occasional unmapped bins, the position of which could be inferred from the context of adjacent mapped bins, were inserted at the appropriate locations (Dryad Data Archive). Finally, from the matrix of markers and genotypes, we inferred the locations of recombination events near and within the inversion (Supplementary Fig. 1).

Identification of centromere positions in the *Boechera stricta* reference genome

To identify the centromere position for each *B. stricta* linkage group, we downloaded sequences of 16 *A. thaliana* centromeric BAC clones and looked for their homologues in the *B. stricta* reference genome using BLAST. These 16 clones were mapped on pericentromeric regions of *B. stricta* by chromosome painting⁴³, thus positions of their homologues on *B. stricta* linkage group indicate margins of centromeres. The lengths of predicted centromere regions are 2 – 3 Mb for linkage groups 1, 2, 5, 6 and 7, and 6 – 7 Mb for linkage groups 3 and 4. As expected, all of these predicted centromere regions show low recombination rates, except parts of these regions on linkage groups 3 and 4 (Supplementary Fig. 4 and Supplementary Table 6).

Identification of a paracentric inversion by comparative chromosome painting

Painting probes—Previous F₂ genetic mapping in the LTM × SAD12 F₂ population found an extensive block of recombination suppression on LG1 (Bs1). Based on the published karyotype structure of *B. stricta* SAD12⁴³ we designed BAC-based painting probes to verify the position and orientation of ancestral genomic blocks on the seven chromosomes of *B. stricta* LTM by comparative chromosome painting (CCP). *A. thaliana* BAC contigs used as painting probes are listed in Mandáková and Lysak⁴⁴. Chromosome-specific BAC contigs were arranged and differentially labeled following the organization of genomic blocks within the reconstructed karyotype of *B. stricta*⁴³.

Whereas chromosomes Bs2-Bs7 have the identical structure, chromosome Bs1 differs between the two *B. stricta* accessions. In LTM, the upper arm of Bs1 was restructured by a paracentric inversion of approximately 8.4 Mb (~4% of the genome). In order to more precisely map the two inversion breakpoints to approximately 100 kb (approximate size of Arabidopsis BAC clones) we carried out BAC FISH experiments on pachytene chromosomes of the accessions SAD12 and LTM. The following Arabidopsis BACs were used for mapping of A1 breakpoint: F21M11 (AC003027), F20D22 (AC002411), F19P19 (AC000104), T1G11 (AC002376), F13M7 (AC004809), T7A14 (AC005322), T25N20 (AC005106), F3F20 (AC007153), T20M3 (AC009999), and T21E18 (AC024174). The A1 breakpoint was mapped between BACs F19P19 (1,125,000 – 1,230,000 bp) and T1G11 (1,233,000 – 1,333,000 bp). The following BACs were used to narrow down the C1 breakpoint: F27J15 (AC016041), F27K7 (AC084414), T27P22 (GenBank Accession missing), F11I4 (AC073555), F9P7 (AC074308), T1N15 (AC020889), F11A17 (AC007932), F21D18 (AC023673), T2J15 (AC051631), and T6B12 (AC079679); the C1 breakpoint was mapped between BACs: F9P7 (17,975,000 – 17,987,000 bp) and F11I4 (17,982,000 – 18,062,000 bp). The *A. thaliana* BAC clone T15P10 (AF167571) bearing 35S rRNA genes was used for chromosomal localization of nucleolar organizer regions (NORs). Clone pCT 4.2 [a 500-bp 5S rRNA repeat (M65137)] was used to identify 5S rDNA loci.

Chromosome preparations—Chromosome spreads were prepared according to the protocol of Lysak and Mandáková⁴⁵ with minor modifications. Entire inflorescences were fixed in ethanol: acetic acid fixative (3:1) overnight and stored in 70% ethanol at 20°C until further use. Prior to chromosome spreading, closed flower buds with white (young) anthers were rinsed in distilled water and in 1 × citrate buffer (10 × citrate buffer: 40 ml of 100 mM citric acid and 60 ml of 100 mM trisodium citrate, pH 4.8) and digested by a pectolytic enzyme mixture (0.3% cellulase, cytohellicase, and pectolyase; all Sigma Aldrich) in 1 × citrate buffer at 37°C for 3 to 6 h, and then kept in citrate buffer until used. An individual flower bud was put on a microscope slide and dissected by needles in a drop of 1 × citrate buffer to form a fine suspension. Then 15 to 30 µl of 60% acetic acid was pipetted to the cell suspension, which was spread over the slide placed on a hot plate at 50°C for ca. 30 sec to 2 min. The spread chromosomes and nuclei were then fixed by pipetting 100 µl of ethanol:acetic acid fixative (3:1) around the suspension drop. The slide was tilted to remove the fixative, dried using a hair dryer and postfixed with 4% formaldehyde in distilled water for 10 min and left to air dry.

Nick translation—DNA probes were labeled by nick translation with biotin-dUTP, digoxigenin-dUTP, or Cy3-dUTP as follows: 1 µg of DNA diluted in distilled water to 29 µl, 5 µl of nucleotide mixture (2 mM dATP, dCTP, and dGTP, 400 µM dTTP; all Roche), 5 µl of 10 × NT buffer (0.5 M Tris-HCl, pH 7.5, 50 mM MgCl₂, and 0.05% bovine serum albumin), 4 µl of 1 mM custom-made ×-dUTP (in which × is biotin, digoxigenin, or Cy3), 5 µl of 0.1 M β-mercaptoethanol, 1 µl of DNase I (2000 U mg⁻¹ diluted to 4 µg ml⁻¹; Roche), and 1 µl of DNA polymerase I (10 U µl⁻¹, Fermentas). The nick translation mixture was incubated at 15°C for 90 min (or longer) to obtain fragments of ~200–500 bp. The reaction was stopped by 1 µl of 0.5 M EDTA (pH 8.0) and by incubation at 65°C for 10 min. The labeled probes were stored at -20°C until use.

FISH and CCP

Prior to pipetting the probe to selected slides, the slides were treated with pepsin (0.1 mg ml⁻¹; Sigma Aldrich) in 0.01 M HCl for 3 – 6 min, postfixed in 4% formaldehyde in 2 × SSC (20 × saline sodium citrate: 3 M sodium chloride, 300 mM trisodium citrate, pH 7.0) for 10 min, dehydrated in an ethanol series (70, 80, and 96%; 3 min each), and air dried. The labeled BAC clones (and rDNA probes) were pipetted together and the DNA precipitated by adding 0.1 volume of 3 M sodium acetate (pH 5.2) and 2.5 volume of ice-cold 96% ethanol, kept at -20°C for at least 30 min, and centrifuged at 13,000 g at 4°C for 30 min. The pellet was dried using a desiccator and resuspended in 20 µl of hybridization buffer (50% formamide and 10% dextran sulfate in 2 × SSC) per slide. The probe and chromosomes were denatured together on a hot plate at 80°C for 2 min and hybridization was carried out by placing the slides into a moist chamber at 37°C for 40 to 63 h. Post-hybridization washing was performed in 20% formamide in 2 × SSC at 42°C. Signal detection and amplification were as follows: biotin-dUTP was detected by avidin-Texas Red (1:1000; Vector Laboratories) and amplified by goat anti-avidin-biotin (1:200, Vector Laboratories) and avidin-Texas Red; digoxigenin-dUTP was detected by mouse anti-digoxigenin (1:250; Jackson Immuno-Research Laboratories) and goat anti-mouse Alexa Fluor 488 (1:200; Molecular Probes); Cy3-dUTP was observed directly. DNA was counterstained with 4',6-diamidino-2-phenylindole (2 µg ml⁻¹) in Vectashield (Vector Laboratories). The hybridization signals were analyzed with an Olympus BX-61 epifluorescence microscope equipped with fluorochrome-specific excitation and emission filters (AHF Analysentechnik), and Zeiss Axio-Cam charged-coupled device (CCD) camera. The monochromatic images were pseudo-colored and processed using the Adobe Photoshop CS5 software (Adobe Systems).

Mapping SNPs to pseudomolecules

In the genetic map, 208 scaffolds were ordered and oriented into 7 linkage groups (LGs). Misjoins were found in original assembly of Scaffold25219 and 19424, which were split into four scaffolds, two with the original names (Scaffold25219 and 19424) and two with new names Scaffold100000 and 200000. In Scaffold25219, the first 135 bins (bin: 0 – 134; position: 12,700,000) were found to be unlinked with the last three bins (bin: 137 – 139; position: 2,740,0012,797,113 bp), while position of bins 135–136 (position: 2,700,0001 – 2,740,000) between them could not be determined. Thus, the first 135 bins were retained in Scaffold25219 (a shorter one, position: 1 – 2,720,000), the last three bins were moved to a

new Scaffold100000 (position 2,740,001 – 2,797,113 of Scaffold25219 was changed to position 1 – 57,113 of Scaffold100000), and bins 135 – 136 were excluded and not used in map. In Scaffold19424, the first 345 bins (bin: 0344; position: 1 – 6,900,000) were found to be unlinked to the remaining 47 bins (bin: 345 – 391; position: 6,900,001 – 7,828,947), thus bins 0 – 134 were kept in Scaffold19424 (position: 16,900,000) and the rest were moved into a new scaffold200000 (position 6,900,001 – 7,828,947 of Scaffold19424 was changed to position 1 – 928,947 on Scaffold200000).

Accordingly, SNP coordinates were extracted from the vcf file (scaffold name and position on scaffold) and examined whether they fell in a bin of the linkage map based on its scaffold position. If so, its position on the pseudomolecules (g') was calculated by $g' = A - |g - c|$ where A is the end position of the bin that the SNP fell in, g is the SNP's position on the scaffold, and c is the end coordinate of the bin on scaffold. For SNPs that did not fall in any bins but were located on regions 2,740,001 – 2,797,113 of Scaffold25219 or 6,900,001 – 7,828,947 of Scaffold19424, their positions on Scaffold100000 and Scaffold200000 were calculated by $g = gI - 2,740,000$ or $g = gI - 6,900,000$, respectively (gI is the position on original scaffold and g is the position on new scaffold). After that, their positions on pseudomolecules (g') were calculated as described above ($g' = A - |g - c|$).

Delimiting and genotyping the inversion

After chromosome painting narrowed down possible locations of the inversion breakpoints in *A. thaliana*, we blasted these Arabidopsis genes onto the *B. stricta* LTM genome and identified putative scaffolds containing the two breakpoints. To identify the exact breakpoints, we mapped paired-end Illumina reads from SAD12 onto the LTM scaffolds with BWA³⁸, followed by *de novo* assembly with Velvet⁴⁶ for read-pairs having one or both ends near the putative breakpoint region in the LTM genome.

Primers Inv-A, Inv-B and Inv-E were designed to amplify the inversion region using Primer 3⁴⁷. Primer Inv-D was used for Sanger sequencing the breakpoints in the two karyotypes for the confirmation of breakpoints. Locations of the breakpoints and related primers are shown in the Dryad Data Archive.

Thermocycling consisted of 94° C for 3 minutes, then 33 cycles of 94° C for 20 seconds, 55° C for 1 minute, 72° C for 30 seconds, followed by 72° C for 6 minutes. Amplified fragments were separated by electrophoresis on a 1% agarose gel stained with SYBR Safe (ThermoFisher Scientific, Waltham, MA). Fragments were ~300bp in accessions containing the inversion, and ~600bp accessions without the inversion.

Differences between subspecies

From our earlier study of ecological divergence between EAST and WEST subspecies¹⁶, we used least-square means (LSMEANS) from 24 genotypes (Table S1) to quantify differences in flowering time. In the greenhouse, EAST genotypes flower an average of 5.0 days before WEST genotypes.

Here we report on three EAST × WEST crosses, which all gave healthy advanced generation progeny in F2, F4, or F6 generations. More generally, intrinsic reproductive isolation might

be present in some instances, especially if complicated by occasional polyploidy or apomixis⁴⁸.

Effect of the inversion on NIL flowering time

In the LTM \times SAD12 (MT_{WEST-inv} \times CO_{East-std})cross¹⁷, we generated two F7 NIL families from the F6 population to investigate the inversion effect. Two F6 heterogeneous inbred families (HIFs: 116A and 120A)⁴⁹ heterozygous for the inversion and mostly homozygous elsewhere in the genome were self-fertilized, and 144 F7 plants from each parent were grown in a completely randomized design in the greenhouse. After three months, plants were vernalized in 4°C for six weeks, and flowering time was recorded. We also genotyped the inversion status in all individuals using inversion primers Inv-A, Inv-B, and Inv-C. We used fixed effects ANOVA to test Flowering Time = Family + Inversion + Family*Inversion, and verified compatibility with parametric statistical assumptions.

For families from this MT_{WEST-inv} \times CO_{East-std} cross, we also analyzed lifetime fecundity in the field, using Dryad-archived data (<http://dx.doi.org/10.5061/dryad.rp3pc>) from the Lost Trail field site in the inversion zone²⁰. We analyzed lifetime fecundity in an EAST \times WEST recombinant inbred F6 population grown in nature for three years.

Effects of the inversion in a sympatric cross

The inversion might be favored if this rearrangement alters genes around the break points and gives it higher fitness over its ancestral standard haplotypes from the same population. To test this, we generated crosses between the reference accession LTM (inversion) and accession SDM, which has the standard haplotype. These genotypes are the same subspecies (WEST) and were collected from similar habitats ~1.7 km apart. Two crosses were generated, one with LTM as mother (CL9.1) and the other with SDM as mother (CL10.1). The F1 hybrids were self-fertilized, and 1,000 F2 plants were grown in the greenhouse. The inversion was genotyped in all individuals, and we measured 11 life history traits (Survival after vernalization [Binary], Flowering [Binary], Width at 4 weeks, Width at 10 weeks, Flowering Time [days after vernalization], Flowering Width, Flowering Height, Flowering Rosette Number, Flowering Leaf Number, Fruit Number, Lifetime Fitness, and Inversion Genotype) on 994 individuals.

To test the phenotypic effects of the inversion, we permuted the inversion genotypes within crosses 1,000 times, and performed MANOVA on the vector of traits. Because R only computes Type I SS for MANOVA, we obtained Type III SS by performing several MANOVAs, and reported the effect of each factor when added to the model last. We modeled MULTIVARIATE-TRAITS = CROSS + INVERSION + CROSS*INVERSION, with cross type (either CL9.1 or CL10.1 as maternal parent), three inversion genotypes (inversion, standard, or heterozygote), and their interaction as fixed-effect predictor variables. *P*-values were obtained by comparing the true Wilk's Lambda statistic to those from 1,000 permutations (Supplementary Table 4). Because MANOVA showed that multivariate means differed between *Bsil* genotypes, the univariate traits were analyzed by ANOVA with a *P* = 0.05 significance threshold⁵⁰.

Gene prediction—We obtained the 10kb region on each side of each breakpoint in the LTM reference genome (inversion) and created pseudomolecules corresponding to the corresponding regions around each breakpoint in the standard haplotype. All four sequences were submitted to the Augustus gene prediction algorithm (<http://bioinf.uni-greifswald.de/augustus/>), and we found no open reading frames spanning the breakpoints in either the inversion or standard haplotypes. Thus, the chromosomal inversion does not disrupt existing or create new open reading frames (Supplementary Fig. 7). Genes predicted by Augustus largely correspond to the transcriptome-assisted annotation in *B. stricta* genome v1.2.

Gene expression—To investigate whether the inversion event changed the expression of flanking genes, we compared two nearby populations using F2 progeny from the LTM × SDM cross (representing inversion and standard haplotypes, respectively). Based on the inversion genotypes, we chose 11 F2 plants homozygous for the inversion and 12 homozygous for standard haplotypes, providing sample sizes well above recommendations based on power analysis⁵¹. All 23 plants were from cross CL9.1 where LTM is the mother. These F2 plants were self-fertilized for one generation, and two-month-old rosette leaves from *Bsi1* homozygote F3 plants were used to compare the expression of seven genes flanking the two breakpoints and syntenic to the region in *A. thaliana*. We used Sigma Spectrum Plant Total RNA Kit to extract RNA and Thermo Scientific DyNAmo cDNA Synthesis Kit to synthesize cDNA. We used Thermo Scientific DyNAmo SYBR Green qPCR Kits for quantitative PCR (qPCR; primers in Dryad Data Archive). As in previous gene expression studies in *B. stricta*¹⁸, we used *ACTIN2* (*ACT2*) as reference and calculated the expression as $Ct = Ct_{ACT2} - Ct_{gene}$ where Ct_{ACT2} is the Ct value of *ACT2* and Ct_{gene} is the Ct value of each gene. Fold gene expression relative to *ACT2* is calculated as 2^{-Ct} . Since 2^{-Ct} has a skewed distribution, we analyzed results from both 2^{-Ct} and Ct . For each gene separately, ANOVA was performed with inversion status as fixed effect (Supplementary Table 5). In addition, we examined statistical power with JMP and with simulations in R.

Dissecting the inversion region in a collinear cross

The EAST × WEST “Parker × Ruby” cross used for QTL mapping was developed from two parents in the East-West contact zone: one in Parker Meadow (RP105, Parker, EAST subspecies, 44°37′ N, 114°31′ W) and one at Ruby Creek (RP109, Ruby, WEST subspecies, 45°33′ N, 113°46′ W). Based on nucleotide similarity (Supplementary Figure 8a, left portion), the WEST-std parent (RP109) used in this cross is very similar to the most recent common ancestor that gave rise to the inversion and its closest WEST-std relatives (IN086 and IN087). The F1 hybrid was self-fertilized to produce F2 plants, and subsequent generations were propagated by self-fertilization and single-seed descent to create 153 independent genetic lines (families). In each line, multiple F4 progeny from the same F3 plant were used in a randomized complete block design totaling 1,714 individuals. The phenotypic LSMEANS were calculated in JMP 8 (SAS, Cary, NC, USA) to represent the genotypic value for their F3 parent.

Genotyping by Sequencing—We used genotyping by sequencing (GBS)⁵² to identify SNPs in this cross. In each family, DNA from each accession was extracted from 0.1g of

young leaf tissue following a dark treatment of about three days. Tissue was stored at -80°C , flash-frozen in liquid nitrogen, and homogenized prior to genomic DNA extraction using Qiagen DNeasy Plant Mini Kits (Qiagen, Valencia, CA). DNA concentration was measured using a Qubit Fluorometer (Turner Biosystems, Invitrogen, Carlsbad, CA). (Qiagen DNeasy Plant Mini Kit) from at least ten pooled F4 individuals to represent the genotype of their F3 parent. This protocol uses an adaptor design which is compatible with TruSeq adaptors and indexes while allowing paired-end sequencing. The combination of 48 unique barcodes with four different TruSeq indexes allowed multiplexing of 192 samples (153 F3:F4 families, 19 inbred individuals of the Parker parent, and 20 inbred individuals of the Ruby parent). Sequencing used Illumina HiSeq-2000 or HiSeq-2500 at the Duke Genome Sequencing & Analysis Core Resource. From this population we obtained ~ 249 million reads with unambiguous barcodes (SRA accession SRP075905). Read pairs were assigned to genotypes and parents by custom Perl code, and low-quality bases in the end of reads were trimmed. SNPs were called using GATK Best Practices (above). In addition, these protocols were used to assay GBS SNPs (SRA accession SRP075997) in 122 genotypes across the inversion area.

Because the LTM \times SAD12 cross does not recombine in the inversion on chromosome 1, we used the Parker \times Ruby cross to infer the linkage map for bins within the inversion. The mean sequencing depth of GBS SNPs in the Parker and Ruby parents was $\sim 20\times$, with standard deviation $\sim 40\times$. We focused on 52,827 bi-allelic SNPs where the two parents have sequencing depth $\geq 6\times$ and $\geq 100\times$ (the mean depth plus two standard deviations), and are homozygous for different alleles. We binned the genome into 100kb windows, and for each individual we counted the number of reads from the two parents, and calculated the proportion of PAR-derived reads. For an individual, if the cumulative depth of all SNPs in a window is less than $20\times$, it was classified as missing, after which we removed windows with $\geq 30\%$ missing data. We used a hard-genotype cutoff based on the portion of PAR reads: PAR proportion ≤ 0.2 was defined as homozygous Ruby, PAR proportion ≥ 0.7 as homozygous PAR, and the remaining windows were classified as heterozygous. These cutoffs give the correct hard-genotype proportions for an F3 population (about 37.5% of either homozygote and 25% heterozygous genotypes). After determining the 'hard genotype calls', we calculated the proportion of the three genotypes in each window. To remove windows with excessive proportions of heterozygotes or either homozygote, we excluded windows where the proportion of the three genotypes are beyond the upper or lower 5% of their respective whole-genome distribution. Given this is a cross in the F3 generation, which has only experienced two generations of recombinations, we regarded it as extremely unlikely to have two recombination events within 1 Mb. Therefore, within each individual, if two recombination events were identified within 1 Mb, we changed the genotype calls of markers between two breakpoints to missing. The linkage map was constructed with MSTmap⁴². With only three windows (markers) dropped out, MSTmap constructed seven linkage groups, with scaffold-chromosome assignment consistent with the LTM \times SAD12 cross. The final linkage map was refined in multiple steps: we first removed markers that are $> 5\text{cM}$ away from both flanking markers and re-constructed the map. In the new map, we imputed missing marker genotype if the up-stream and down-stream markers with data have the same genotype call and are not both $> 10\text{cM}$ away from the missing position. Another

linkage map was constructed, and the imputation process was repeated since the new map had slightly changed the order of some markers, making some missing genotypes now imputable. The final linkage map was then constructed with 1010 markers. Thus, we used the Parker \times Ruby cross to infer the linkage map for bins within the inversion and we inferred the position of polymorphisms within the inversion from their positions in each ordered and oriented contig.

This collinear EAST \times WEST “Parker \times Ruby” greenhouse experiment consists of 12 blocks, each with one F4 plant from each of the 153 lines and multiple Parker and Ruby individuals ($N=1,714$). Seeds were stratified in 4°C for four weeks and planted in “Containers” (Ray Leach SC10, Stuewe & Sons Inc., Tangent, OR, USA), with soil composition and greenhouse conditions as previously described¹⁶. When rosettes were 11-weeks old, all leaves from three blocks of plants were harvested for rosette- and leaf-morphology measurements as described¹⁶. At 12 weeks of age, the remaining nine blocks were vernalized in 4°C for 6 weeks, then returned to the same greenhouse conditions for phenology measurements. All traits were measured as previous described¹⁶, except: 1) no physiological traits were measured; 2) leaf width/length ratio was used instead of leaf shape morphometrics because the leaf-shape landscape points were highly correlated¹⁶ (Supplementary Table 7).

For QTL mapping, all individual-level measurements were transformed to family-level LSMEANS in JMP ($N=153$ families). Due to the skewed distribution of most phenotypes, all traits (except binomial traits or plant stages) were log-transformed at the individual level. For greenhouse measurements, BLOCK and GENOTYPE were considered as random effects. We used R⁵³ libraries qvalue and MASS for MANOVA, False Discovery Rate, and Discriminant Function Analysis. For each marker across the inversion interval and adjacent contigs we fit MANOVA model: MULTIVARIATE-PHENOLOGY-TRAITS = BLOCK + SINGLE-MARKER-GENOTYPE. At each marker we computed Wilkes Lambda, permuted the vector of phenotypes with respect to marker genotypes 1,000 times, and finally corrected for false discovery rates across the inversion region⁵⁴.

Discriminant Function Analysis (DFA) was performed in R for a marker at the peak of each multivariate QTL (bins Scaffold-25219_50000, Scaffold-13671_1150000, and Scaffold26675_450000), defining the direction of trait variation along the axis of greatest differentiation among the multivariate means of these marker genotypes. Subsequently, at these three markers we used the eigenvector for the first DFA axis to compute the phenotypic projection of each genotype on the trait axis identified by DFA. These new trait values were used for univariate QTL mapping across the inversion region, with correction for false discovery rates across the inversion region.

Finally, we annotated SNPs by using snpEff v.4.2⁵⁵, and found 17 genes that are orthologues of flowering-time genes of *Arabidopsis thaliana*⁵⁶ within the inversion interval. See candidate genes in Supplementary Table 8 and annotated SNPs in Dryad Data Archive.

Population genetics

SNP filtering—SNPs called by GATK were filtered in each of three groups: sets consisting of 35 inversion genotypes (G_{INV}) and 87 standard genotypes (G_{STD}) from the inversion zone, and a “Reference Population” set of 83 genotypes from across the species range, outside the inversion zone (G_{RP}). Next, data from these single groups were pooled together into two datasets, the complete data set ($G_{INV}+G_{STD}+G_{RP}$) and the inversion zone genotypes ($G_{INV}+G_{STD}$). In each dataset, genotypes supported by less than 2 reads were assigned as missing, and SNPs were discarded if they met the following criteria in any of the groups: 1) detected in fewer than 50% of individuals; 2) mean depth more than 20; 3) more than one variant allele was observed; 4) sites with a proportion of heterozygous genotypes more than 15% (*B. stricta* is predominantly inbred¹⁷, hence high heterozygosity may indicate paralogous loci); 5) reference or variant alleles are indels. After filtering, 75,737 and 43,722 SNPs were retained for the complete set of genotypes $G_{INV}+G_{STD}+G_{RP}$ and for the inversion zone set of genotypes $G_{INV}+G_{STD}$, respectively. The following analyses used these SNPs.

Ancestral allelic state—Several population genetic statistics require information on the ancestral state of segregating variants. Therefore, we sequenced a *Boechea holboellii* (= *B. retrofracta*; reference genotype “Panther”, location N45 18.198, W114 22.599) individual with mean depth of ca. 400× (JGI Project ID: 1051698). Short reads were mapped to the *B. stricta* reference genome using BWA, genotypes were called using GATK with default settings, and sites were filtered out if they had depth less than 20, higher than 800, or if they were heterozygous. A Python script called ancestral alleles as the shared allele between *B. holboellii* and *B. stricta*. In total, we obtained information for ancestral states for 63,182 (83.4%) of 75,737 and 35,100 (80.3%) of 43,722 SNPs for $G_{INV}+G_{STD}+G_{RP}$ and $G_{INV}+G_{STD}$, respectively.

Windows—We partitioned the genome into 300kb windows. The sequence coverage of each window was calculated by counting the number of available sites (both variant and non-variant sites obtained from GenotypeGVCF by using `-allSite`) per window passing the quality filters, i.e., detected in at least 50% individuals, mean depth no more than 20, and proportion of heterozygous genotypes no more than 15% in each of the three groups (G_{INV} , G_{STD} , and G_{RP}). Windows were distributed on a per-scaffold basis beginning at position 1 of a scaffold and were then oriented along pseudo-molecules according to the linkage map. After excluding windows with sequence coverage less than 5 kb, 2,801 windows were retained for downstream analyses.

Phylogenetic relationship and population structure analyses—Python scripts were used to generate alignments from genotypes in the vcf file, with missing and heterozygous loci coded as “N”. With genome-wide SNPs, we verified EAST-WEST population structure using principal component analysis (PCA) with EIGENSOFT v6.0⁵⁷.

To investigate the origin of the *Bsil-inv* haplotype we used data from 901 SNPs in the inversion region of the genome to examine relationships among 35 *Bsil-inv* and 83 *Bsil-std* haplotypes. (Four admixed *Bsil-std* genotypes were excluded.) First, we constructed

neighbor-joining (NJ) trees using MEGA v6.06 with 1,000 bootstrap steps. Second, maximum-likelihood trees were constructed using RAxML version 8.0.0⁵⁸. One thousand ML trees were generated to find the best-scoring ML tree, and topological robustness was investigated using 1,000 nonparametric bootstrap replicates. NJ and ML trees were displayed by Figtree (<http://tree.bio.ed.ac.uk/software/figtree/>).

Site frequency spectrum and summary statistics—To control for error rates and variable coverage of short-read sequencing, we used a probabilistic method implemented in the software package ANGSD v0.911⁵⁹ to estimate the site frequency spectrum (SFS) and related population genetic statistics. A number of filtering steps were performed: 1) removed reads with a minimal mapping quality of 30 and bases with a minimal quality score of 30 (-minMapQ and -minQ); 2) removed sites with information from less than 50% individuals (-minInd); 3) removed sites with *P*-value higher than 1e-6 (-snp_pval); 4) assigned genotypes as missing if the depth is less than two for an individual; 5) only used genotypes with a posterior probability higher than 0.95; 6) removed sites that did not pass previous filtering criteria (above). Because *B. stricta* is predominantly inbreeding¹⁷, we estimated inbreeding coefficients for each individual in ngsTools⁶⁰ and incorporated them into the calculation of SFS in ANGSD. Using genotype likelihoods based on the GATK genotyping model⁶¹, we estimated folded and unfolded SFS and derived a set of population genetic summary statistics in 300kb windows. For the G_{INV} and G_{STD} groups, we estimated Tajima's D ⁶² based on both folded and unfolded SFS, and calculated Fay & Wu's H^2 ⁹, Fu and Li's D and F ⁶³ based on unfolded SFS. The inferred ancestral allelic state based on short reads from *B. retrofracta* genotype “Panther”, aligned to the *B. stricta* reference genome was used to estimate the unfolded SFS. We compared population genetic statistics in the inversion region, on other chromosomes (LG2 – LG7), and in Block D⁶⁴, a chromosome region with unusually high polymorphism in related species, perhaps due to clusters of NB-LRR genes⁶⁵.

F_{ST} , d_{XY} , d_A and π —Genetic differentiation (F_{ST})⁶⁶ between G_{INV} and G_{STD} groups was estimated using VCFtools v0.1.12⁶⁷, and we used Python scripts to estimate nucleotide diversity (π)⁶⁸ within each group, pairwise nucleotide divergence (d_{XY})⁶⁹ and net pairwise nucleotide divergence (d_A)⁶⁹ between groups. All parameters were calculated on a per site basis, and then averaged to obtain window-based estimates (300 kb windows). The window-based F_{ST} was calculated by averaging per site F_{ST} across all variable loci, and the window-based π , d_{XY} and d_A were estimated by averaging all sites (both variable and monomorphic) passing the initial quality filters for each window (above).

Linkage disequilibrium—The level of linkage disequilibrium (LD) between the inversion and all SNPs on chromosome 1 (including those in the inverted region) was estimated as the mean squared correlation (r^2) for 118 samples from inversion zone accessions (35 *Bsil-inv* and 83 *Bsil-std*) using plink v1.90⁷⁰. (Four admixed *Bsil-std* samples were excluded, see above.) In addition, to compare LD between the inversion and flanking SNPs, we identified ten randomly chosen comparator SNPs from LG2 – LG7, having similar derived allele frequencies (25 – 35%, close to the 29.7% frequency of *Bsil-inv* in the inversion zone), and then calculated LD between them and flanking SNPs along the chromosomes. For LG1 (Fig

5), strong LD ($r^2 > 0.4$) between the inversion and SNPs within and outside the inverted region extended to the end of the chromosome (more than 10 Mb). In contrast, LD with the ten comparator SNPs from LG2 – LG7 declined quickly to background levels (Supplementary Fig. 8b).

Age of the inversion—To estimate age of the inversion, we phased the 35 *Bsil-inv* genotypes using Beagle v4.1⁷¹ with default settings, and estimated pairwise genetic distances among haplotypes on the inverted genome region. After that, the divergence time between haplotypes was calculated by $T = d/2\mu$, where d is pairwise genetic distance and μ is the mutation rate. We used a mutation rate of 7×10^{-9} per site per generation in *Arabidopsis thaliana*²⁴) with a generation time of two years in *B. stricta*. To avoid biases due to missing data, we only considered 1,675 haplotype pairs with less than 50% missing sites in the inversion region. We approximated the lower bound for age of the inversion based on the mean pairwise distances⁷², and the upper bound from the maximum of all pairwise distances among INV genotypes.

Divergence time between QTL alleles in the collinear cross—Using the same mutation rate and generation time, we estimated $D_A = D_{xy} = 2\mu T$, and solved for T between these two alleles in 200kb non-overlapping windows⁶⁹. This window size was chosen to ensure sufficient polymorphic sites within each interval.

Frequency distribution of derived alleles—We tested whether *Bsil-inv* has unusually high frequency among derived SNP alleles in the inversion zone population. For this analysis, we excluded one admixed, genetically divergent inversion individual (IN019, right half of Supplementary Fig. 9c), and analyzed *Bsil-inv* and closely related *Bsil-std* genotypes from the inversion zone (left half of Supplementary Fig. 9c, PC1 < 0.0). These genotypes comprise 54 samples (34 *Bsil-inv* and 20 *Bsil-std*), with 2,416 SNPs that, like the inversion, segregate among these genotypes and are monomorphic in the rest of our collection.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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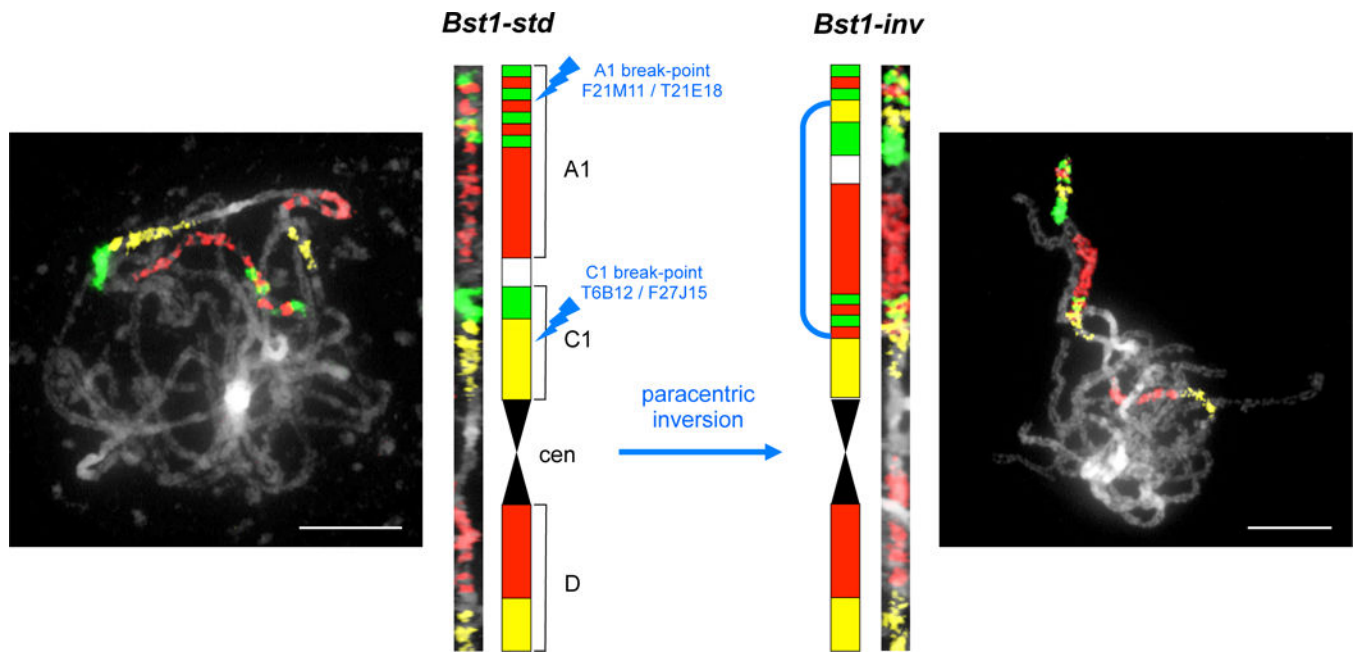


Fig. 1. Comparative Chromosome Painting of chromosome 1 in *Bst1-std* and *Bst1-inv* haplotypes
 Shown are *in situ* chromosomal localization of painting probes on *B. stricta* pachytene chromosomes, their straightened images and diagrammatic representations of the *Bst1-std* and *Bst1-inv* haplotypes. Differential labeling of BAC contigs (F21M11, T21E18, T6B12, F27J15) identifies the breakpoints and extent of the paracentric inversion in the *Bst1-inv* genotype. Scale bars, 10 μm . See Supplementary Fig. 3 for a detailed cytogenetic analysis.

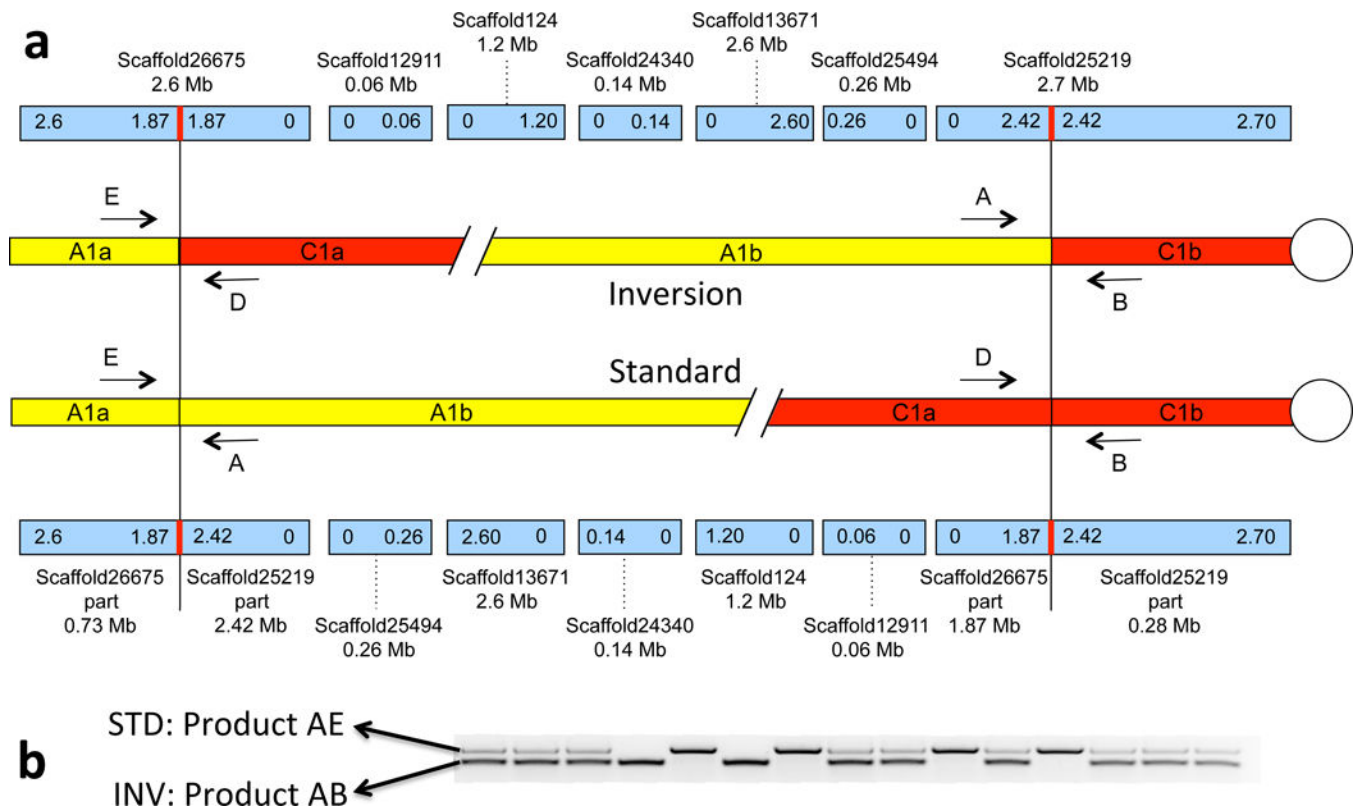


Fig. 2. Inversion region and PCR genotyping

(a) Chromosome blocks (yellow and red) are shown for the derived (*BstI-inv*, upper) and ancestral (*BstI-std*, lower) haplotypes. Centromeres are indicated by white circles. Scaffolds (blue, above and below) are labeled with their name and size. Breakpoints are shown as vertical red lines within the scaffolds. Arrows indicate PCR primers. There are 1,591 annotated genes in the inversion region. (b) Gel image showing PCR products, allowing codominant identification of inversion genotypes.

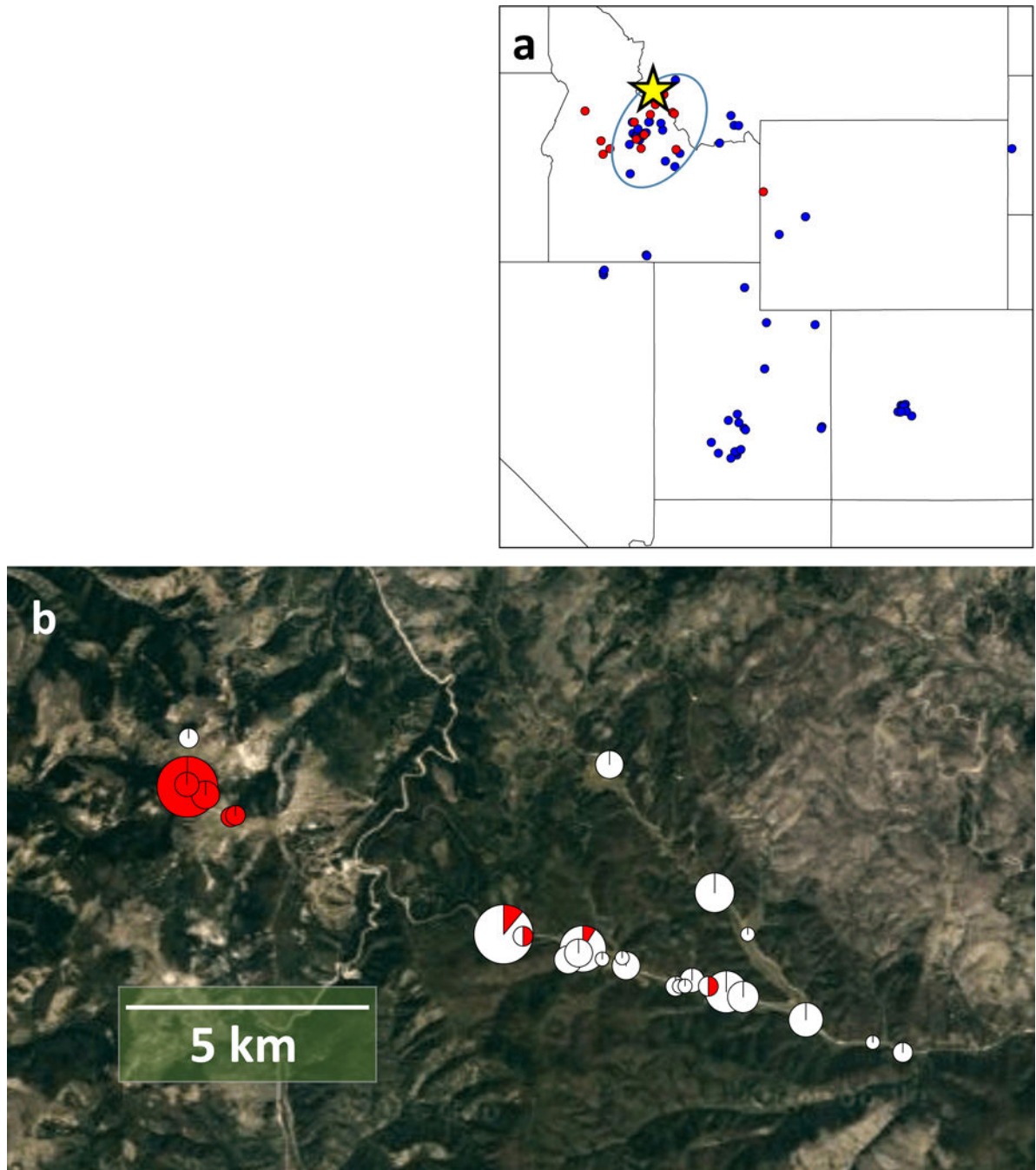


Figure 3. Geographic locations of subspecies collections and inversion zone genotypes. Geographic locations of genotyped accessions used in this study

(a) Species-wide samples, with each dot indicating one genotype from a population. Subspecies assignments show EAST (blue) and WEST (red). Ellipse indicates the contact zone where subspecies overlap, and the star shows the inversion zone. $N=83$. The collinear cross comes from within the hybrid zone (oval), outside the inversion zone (star). (b) Close-up of populations near the inversion zone (the star in Fig. 3a), with pie-diagrams indicating the inversion frequency (red) in each population, sized in proportion to sample size.

Forested, unsuitable habitat likely limits gene flow. $N = 122$ genotypes. Individual populations range in size from 1 to 19.

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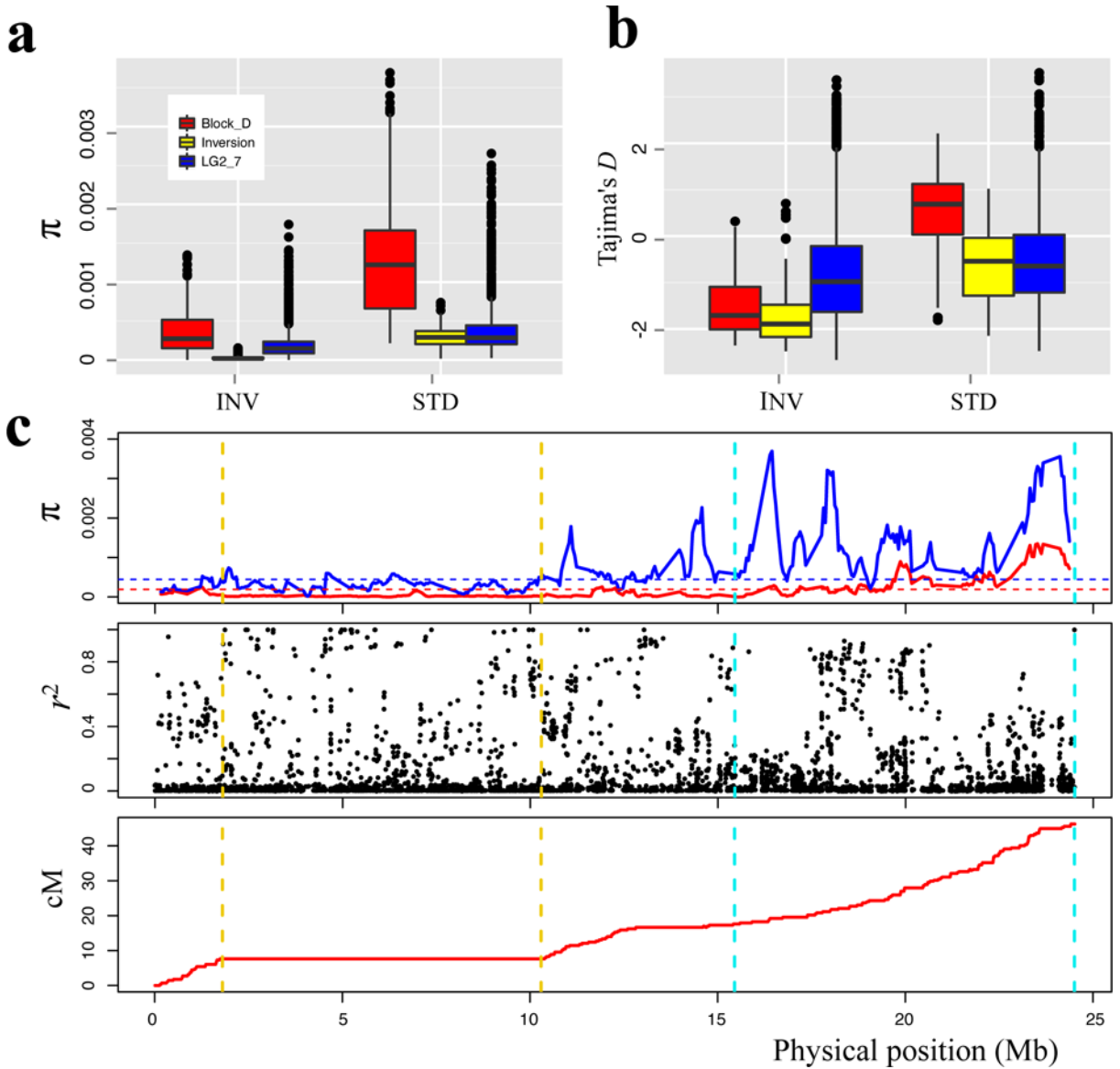


Figure 4. Population genetic variation for inversion and standard groups

(a, b) Nucleotide diversity (π) and Tajima's D of inverted genomic region (yellow), block D of chromosome 1 (red) and chromosome 2–7 (blue) in inversion (INV) and standard (STD) genotypes. Block D (south end of LG1) is treated separately because it has unusually high polymorphism in related species. In these box plots, the median is shown by a horizontal line, while the bottom and top of each box represents the first and third quartiles. The whiskers extend to 1.5 times the interquartile range. Outliers are represented by black dots. (c) Distribution of population genetic statistics along chromosome 1. Nucleotide diversity (π) in INV (red) and STD (blue) genotypes, with genome-wide averages as dashed horizontal lines. Linkage disequilibrium (R^2) between the inversion and SNPs is shown in all INV and STD genotypes from the inversion zone, and the relationship between physical and linkage maps. The dashed vertical lines mark the inverted (golden) and block D (light blue) regions. $N = 122$, except four admixed individuals were excluded from LD analysis.

LD for 10 comparator SNPs with derived allele frequencies similar to *Bsil-inv* is shown in Supplementary Fig. 8b.

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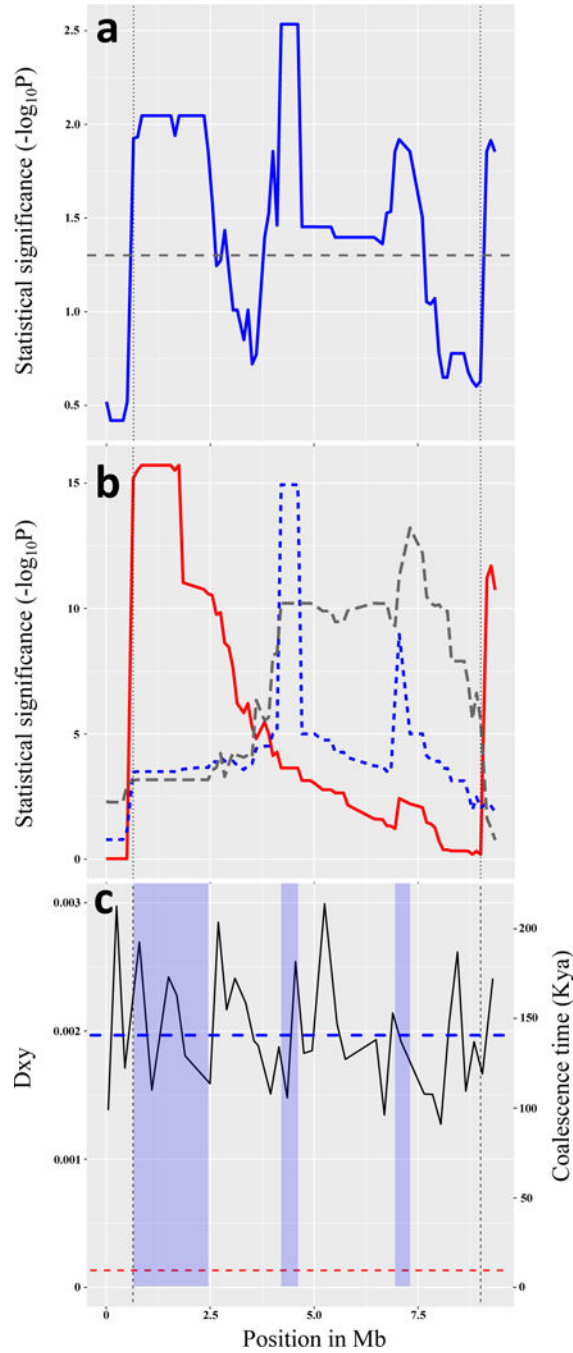


Figure 5. QTLs within the inversion

A collinear cross shows that several QTLs in the inversion region influence phenology and development traits. Inversion breakpoints are indicated by vertical lines. (a) Multivariate QTL mapping finds several QTL peaks (blue line) exceeding the significance threshold (horizontal dashed line); $N = 1,714$ F4 individuals in 153 families. (b) Testing the hypothesis that three linked QTLs have different pleiotropic effects. Plots for three QTLs (left, center, and right, shown in solid red, dotted blue, and dashed gray, respectively) quantify evidence that a locus with different patterns of pleiotropy occurs at each peak. Discriminant Function

Analysis was used to identify new composite trait axes defined at each peak, and evidence for these composite traits was mapped across the inversion region. The left and center QTLs show little overlap, suggesting different patterns of pleiotropy. (c) Comparison of molecular divergence and time to coalescence of the East and West genotypes in the collinear QTL mapping cross. The horizontal red dashed line at 8.8 Kya is the upper confidence interval for age of the inversion. The vertical axes show nucleotide divergence (D_{xy} , left axis), and coalescence time (Kya, right axis). The blue dashed line indicates the mean genome-wide values of D_{xy} and coalescence time between these East and West alleles. The horizontal axis shows position across the inversion region in Mb, beginning at marker Scaf26675_2450000. Vertical blue shading indicates the QTL regions, $\pm \log P > 1.6$ confidence intervals, with 408 annotated genes in these QTL regions. Divergence (D_{xy}) between these genotypes was calculated in 200kb non-overlapping windows, and the coalescence time was estimated using $T = D_{xy}/2\mu$, where μ is $7E-9$ per site per generation. Mean D_{xy} in the three QTLs (left to right) are 0.00206, 0.00169 and 0.00186, corresponding to coalescent times of 147 kya, 121 kya and 132 kya, respectively. The mean D_{xy} in the whole inversion region is 0.00191, and the average coalescence time is 136 kya.