Selection Signatures Underlying Dramatic Male Inflorescence Transformation During Modern Hybrid Maize Breeding

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ABSTRACT Inflorescence capacity plays a crucial role in reproductive fitness in plants, and in production of hybrid crops. Maize is a monoecious species bearing separate male and female flowers (tassel and ear, respectively). The switch from open-pollinated populations of maize to hybrid-based breeding schemes in the early 20th century was accompanied by a dramatic reduction in tassel size, and the trend has continued with modern breeding over the recent decades. The goal of this study was to identify selection signatures in genes that may underlie this dramatic transformation. Using a population of 942 diverse inbred maize accessions and a nested association mapping population comprising three 200-line biparental populations, we measured 15 tassel morphological characteristics by manual and image-based methods. Genome-wide association studies identified 242 single nucleotide polymorphisms significantly associated with measured traits. We compared 41 unselected lines from the Iowa Stiff Stalk Synthetic (BSSS) population to 21 highly selected lines developed by modern commercial breeding programs, and found that tassel size and weight were reduced significantly. We assayed genetic differences between the two groups using three selection statistics: cross population extended haplotype homozygosity, cross-population composite likelihood ratio, and fixation index. All three statistics show evidence of selection at genomic regions associated with tassel morphology relative to genome-wide null distributions. These results support the tremendous effect, both phenotypic and genotypic, that selection has had on maize male inflorescence morphology.

KEYWORDS maize; inflorescence; tassel; selection; GWAS

The male inflorescence, or tassel, of maize (Zea mays L.) is a branched structure that displays remarkable morphological diversity. Tassels vary in length, number of branches, compactness, and curvature, all of which are capable of affecting the quantity and dispersal of pollen. Inflorescence architecture is crucial to reproduction in wind pollinated plants (Friedman and Barrett 2009) such as maize. As such, tassel size and shape are likely to have been subject to both natural and artificial selection. Tassels on teosinte plants, the wild progenitor of maize, are characterized by a large number (>50) of branches (Xu et al. 2017), implying a reproductive advantage of numerous branches in natural populations. In cultivated maize, however, tassel size and branch number have decreased over the course of modern breeding (Meghji et al. 1984; Duvick 1997, 2005). Tassel size is negatively correlated with grain yield, putatively due to shading caused by large tassels (Duncan et al. 1967; Hunter et al. 1969). At high planting densities typical of modern agronomic practices, tassels can intercept enough sunlight to reduce photosynthesis lower in the canopy, which can negatively affect assimilation of photosynthate in the ear (Duncan et al. 1967; Wardlaw 1990; Hammer et al. 2009). A study in tropical maize indicated that direct selection on branch number can result in an indirect increase in grain yield (Fischer et al. 1987), showing that the relationship...
between tassel and ear may be causative and persists beyond temperate germplasm.

Previous studies have identified quantitative trait loci (QTL) and quantitative trait nucleotides associated with different aspects of tassel morphology and development that colocalize with teosinte branched1 (tb1) and teosinte glume architecture1 (tga1), which were targets of domestication that changed plant and inflorescence architecture between teosinte and modern maize (Doebley et al. 1995; Wang et al. 2005; Brown et al. 2011; Wu et al. 2016). A study mapping tassel architecture in a population of recombinant inbred lines derived from a cross between maize and teosinte identified a QTL and evidence of selection at the barren inflorescence2 (bi2) locus (Xu et al. 2017), which has been characterized previously as decreasing the number of tassel branches and spikelets (McSteen and Hake 2001). The same study identified five additional QTL that colocalized with inflorescence development genes and were in the list of genes identified as putatively selected during maize domestication by Hufford et al. (2012). These findings provide evidence for selection on tassel morphology during domestication; however, tassel morphology has continued to change postdomestication, including significant differences observed within breeding germplasm developed between the early 20th century and modern day (Meghji et al. 1984; Duvick 2005).

Until 1935, most of the maize grown in the North American Corn Belt consisted of open-pollinated populations, in which plants are allowed to reproduce freely with each other. Within 4 years, however, >90% of maize grown in Iowa had transitioned to higher-yielding hybrids created by controlled crosses, and the rest of the Corn Belt soon followed (Reif et al. 2005). The transition to single-cross hybrids was largely driven by key founder lines (B14, B37, and B73) derived from a pivotal maize population called Iowa Stiff Stalk Synthetic (BSSS), which was created in the 1930s by intermixing 16 inbred lines (Lamkey et al. 1991; Reif et al. 2005). These founder lines and their combinations, referred to as Stiff Stalk lines, became preferentially used as females in hybrid breeding schemes, while other inbred lines that combined well with them were used as males (Reif et al. 2005). These early choices laid the framework for what would eventually become the heterotic groups used in maize breeding today. Descendants of the BSSS population have been used universally across proprietary breeding programs, resulting in a number of commercial Stiff Stalk lines protected by Plant Variety Protection certificates (PVP lines) (Mikel and Dudley 2006).

The shift in breeding methods to single-cross hybrids placed an emphasis on developing inbred lines with characteristics that enhance their potential to serve as parents. These parents are subsequently crossed to each other to produce vigorous hybrid offspring. In open-pollinated populations, a competitive advantage was conferred by greater pollen production and dispersal (Friedman and Barrett 2009). Inbred line breeding greatly reduced that competitive advantage as breeders began making more controlled crosses; there is also evidence that inbreeding tends to shift resource allocation to the female inflorescence (Burd and Allen 1988). Thus, we hypothesized that selection in the Stiff Stalks by breeders between the original BSSS population and modern PVP lines acted to reduce overall tassel size.

Among the germplasm used in this study are inbred lines derived from the original BSSS population, as well as a number of commercially developed inbred lines for which the Plant Variety Protection certificates have recently expired (ex-PVPs). These ex-PVP lines represent the most modern germplasm from private seed companies that is available to the public, and they can be traced back to founder lines from the BSSS population. These two sets of materials afford us an opportunity to evaluate changes in maize tassel architecture caused by the accelerated evolutionary process of modern breeding. We used single nucleotide polymorphisms (SNPs) identified as significant in association mapping as a priori selection candidates, and compared selection statistics at those SNPs to their genome-wide distribution. If selection by breeders between the BSSS and modern ex-PVPs acted to reduce tassel size, we expect that loci associated with tassel morphology phenotypes would show enrichment for selection signatures relative to the rest of the genome. Indeed, we observe an enrichment of selection statistic scores in regions containing SNPs associated with tassel morphological traits, a finding that is consistent across three different selection statistics.

Materials and Methods

Populations

This study used two populations of maize inbred lines: a diversity panel and a nested association mapping (NAM) population. The diversity panel (WiDiv-942) consists of 942 diverse lines that typically reach grain physiological maturity in the Midwest regions of the United States, and represents an expansion of the 503 line Wisconsin Diversity panel (WiDiv-503; Hirsch et al. 2014; M. Mazaheri, M. Heckwolf, B. Vaillancourt, J. Gage, B. Burdo, S. Heckwolf, K. Barry, A. Lipzen, C. Bastos Riviero, T. Kono, H. Kaepppler, E. Spalding, C. Hirsch, C. R. Buell, N. de Leon, and S. Kaepppler, unpublished data) to increase breadth and depth of genetic diversity as well as to include a greater representation of ex-PVP inbreds. The NAM population (PHW65 NAM) was styled as a smaller version of the maize NAM (McMullen et al. 2009) and consists of three biparental populations that share a common parent, the inbred line PHW65. PHW65 is an ex-PVP Lancaster-type line. The three founder lines are PHN11, Mo44, and MoG. PHN11 is an ex-PVP Iodent line closely related to PH207; Mo44 is a Missouri line derived from Mo22 and Pioneer Mexican Synthetic 17 (Flint-Garcia et al. 2005); and MoG is a Missouri line characterized by wide leaves, a heavy stalk, and high pollen production. The founder lines were chosen for their differing tassel morphologies: MoG and Mo44 have longer tassels than the recurrent parent PHW65; PHN11 has more branches than PHW65; and the MoG tassels are heavy and curved while the PHN11 tassels...
are light and more rigid. Together, the four parental lines comprise a diversity of tassel size and shape. Each biparental population consisted of 200 individuals derived by double-haploid generation from the F2 of the original parents.

**Phenotypic measurements**

The WiDiv-942 was grown in the summers of 2013 and 2014 at the University of Wisconsin’s Arlington Agricultural Research Station and in the summer of 2015 at the West Madison Agricultural Research Station. The PHW65 NAM was grown in 2014 at the Arlington Agricultural Research Station and in 2014 and 2015 at the West Madison Agricultural Research Station. All experiments were planted as randomized complete block designs with two replications at each location in 4.57 m long single row plots with 0.76 m between rows at a density of 72,000 plants per hectare.

Both populations were measured for branch number (BN), tassel length (TL), and spike length (SL) on three representative plants per plot during or after flowering. BN was quantified as the number of primary tassel branches, TL was measured as the distance in cm from the lowest tassel branch to the tip of the spike, and SL was measured as the distance in centimeter from the uppermost tassel branch to the tip of the spike. All traits were measured in all environments with the exception of SL, which was not measured in the PHW65 NAM at West Madison in 2014. Three derived tassel traits can be calculated from these measurements: branch zone length (BZ) as TL minus SL, branch density (BD) as BN divided by BZ, and spike proportion (SP) as SL divided by TL.

In addition to manual measurements, tassels measured in 2015 were also imaged and measured automatically using TIPS (Gage et al. 2017), which estimates TL and BN, as well as tortuosity (TR), compactness (CP), fractal dimension (FD), skeleton length (SK), perimeter length (PR), and tassel area (TA) from two-dimensional profile images of tassels. TR is a measure of tassel curvature expressed as a proportion, the length of a straight line from base to tip divided by the length of a spline fit along the tassel itself from base to tip; TA is the number of pixels in the tassel; CP is measured as TA divided by the area of the tassel’s convex hull; FD is a measure of overall tassel complexity; SK is a measure of overall linear length of the tassel, including the main spike and all branches; and PR measures the length of the tassel’s outline. Tassel weight (TW) was also measured as the mean dry weight of three tassels per plot.

Rather than directly use TA and the estimates of TL and BN produced by TIPS, we used partial least squares regression models to generate more accurate, image-based predictions of TL, BN, SL, and TW. Models were fit using the pls package (Mevik and Wehrens 2007) in R (R Core Team 2016). Individual tassel measurements were used for model fitting and prediction was done using all eight model components (TIPS produces eight tassel measurements). The TIPS output for the PHW65 NAM population were used as explanatory variables and the model was trained on manually measured values of TL, SL, BN, and TW in the PHW65 NAM. The resulting partial least squares regression models were used to predict image-based values of TL, SL, BN, and TW from the TIPS output for images of the WiDiv-942 population. The opposite was done as well: TIPS outputs for the WiDiv-942 population were used as explanatory variables in conjunction with manually measured values of TL, SL, BN, and TW to train the models, which were then used to predict image-based values of TL, SL, BN, and TW for the PHW65 NAM individuals. Image-based predictions of TL, SL, BN, and TW are referred to as TLp, SLp, BNp, and TWp.

The WiDiv-942 inbred line best linear unbiased predictions (BLUPs) for all manually measured traits, except TW, were calculated as the random genotypic effect predictions from the linear model $y = \mu + g + e + g \times e + b(e) + \epsilon$, where $y$ is the phenotypic response, $g$ is the genotypic effect of an inbred line, $e$ is the environmental effect of a year–location combination, $g \times e$ is the genotype-by-environment interaction, $b(e)$ is the effect of replicate within environment, and $\epsilon$ is the independently and identically distributed error term. The PHW65 NAM BLUPs for manually measured traits, except TW, were calculated from the model $y = \mu + g + e + p + g \times e + p \times e + b(e) + \epsilon$, where $p$ is an effect attributed to each biparental population, $p \times e$ is a population-by-environment effect, and all other terms are the same as in the WiDiv-942 model. BLUPs for image-based traits and TW for both populations were calculated from models identical to those above, but with all terms containing an environmental effect removed and a standalone replication term included. All effects were considered random, and models were fit with the R package lme4 (Bates et al. 2016). BN, BD, PR, SK, and TW were square root transformed to better meet the model assumption of normally distributed residuals, and TR was removed from further analysis due to violation of model assumptions. All BLUPs can be found in Supplemental Material, File S3.

Heritability was calculated on an entry-mean basis for all manually measured traits except for TW as $h^2 = \sigma^2_g / \sigma^2_e$, where $\sigma^2_g$ and $\sigma^2_e$ were estimated from the linear models described above and $g$ and $e$ are the number of replications and the number of environments, respectively. Repeatability was calculated similarly for all image-based traits and TW as $r = \sigma^2_g / \left( \sigma^2_g + \sigma^2_e \right)$, because image-based traits and TW were measured only in a single environment.

Genotypic and phenotypic correlations between all traits were calculated for each population. Because of differing numbers of phenotyped lines and differing numbers of environments in which the traits were evaluated, the correlations were only calculated using data from the West Madison 2015 location, and only with lines that were phenotyped for all 15 traits in that environment (693 individuals from the WiDiv-942 and 569 from the PHW65 NAM).

**Genotypic data**

The WiDiv-942 lines were genotyped by RNA sequencing as described in Hirsch et al. (2014). RNA sequencing data were
combined with RNA sequencing data from the original Hirsch et al. (2014) publication and aligned to the B73 version 4 reference genome sequence (Jiao et al. 2017) for SNP calling at 899,784 SNPs. The WiDiv-942 SNP matrix contained 30% missing data, which was imputed using fastPHASE v1.4.0 (Scheet and Stephens 2006) with the −H flag set to −3 to prevent haplotype phasing and default parameters otherwise.

Inbred lines in the PHW65 NAM population were genotyped using genotyping-by-sequencing (GBS) as described in Elshire et al. (2011), using the GBSv2 pipeline implemented in TASSEL5 (Bradbury et al. 2007), resulting in 9291 SNPs, which were imputed with FSFHap (Swarts et al. 2014) using the “cluster” algorithm with a minimum MAF of 0.3 within each biparental population, a window size/step size of 30/15 for chromosomes 1, 4, and 6, and 50/25 otherwise.

The four parents of the PHW65 NAM population were resequenced and genotyped at 23.3 million SNPs (Brohammer et al. 2018). The 31% missing SNP data in the parents was imputed using fastPHASE v1.4.0 (Scheet and Stephens 2006) with the −H flag set to −3 to prevent haplotype phasing, the −K flag set to 7, and the −T flag set to 10. The −K flag sets the number of haplotype clusters at 7 and was chosen by evaluating imputation accuracy for K values between 1 and 31 and choosing the value above which minimal gains in accuracy were observed, similar to Tian et al. (2011). The −T flag sets the number of random starts of the algorithm. As opposed to the imputation of the WiDiv-942, setting the −K and −T flags helped reduce the computational time of imputation for the larger number of resequencing SNPs. SNP calls from the four resequenced parents at 10.6 million polymorphic SNPs were projected onto their biparental progeny in the PHW65 NAM using GBS markers as a scaffold, as described in Tian et al. (2011).

Classification of public and ex-PVP Stiff Stalk lines

We identified lines that represent early Stiff Stalk germplasm, recent ex-PVP Stiff Stalk germplasm, and publicly developed Stiff Stalk germplasm that represents a temporally intermediate group between the two.

The WiDiv-942 contains 47 inbreds derived through self-pollination from the original BSSS population (referred to hereafter as BSSSC0), which represent maize Stiff Stalk germplasm from the early 20th century.

From all ex-PVP lines in the WiDiv-942, inbreds were chosen that showed genotypic evidence of being directly descended from the BSSS population (File S1). Although pedigree information was available for some ex-PVPs, the number of individuals that could be traced by pedigree back to BSSS-derived inbreds was low. Additionally, using genotypic information to identify BSSS-derived ex-PVP lines allows exclusion of individuals showing evidence of contamination from other breeding material. Population substructure within all the ex-PVP lines was estimated using ADMIXTURE (Alexander et al. 2009). The genotypic matrix consisting of 899,784 SNPs for 328 ex-PVP inbreds was first pruned for linkage disequilibrium (LD) using PLINK (Chang et al. 2015). Using a sliding window of 50 SNPs and a step size of 10 SNPs, SNPs with a $R^2$ value $>0.1$ within the window were discarded, leaving 109,605 SNPs. The pruned marker set was used in ADMIXTURE without seeding founding individuals. Ten-fold cross-validation of the errors was minimized at K groups equal to eight as an appropriate model for this marker set. These eight groups represent the resulting subpopulations; B14, B37, B73, Oh43, Lancaster, Iodent, Iodent-Lancaster, and Flint. To identify lines that both display evidence of descent from the BSSS as well as exclude lines with evidence of introgression from other germplasm pools, individuals were required to have a sum of $>95\%$ identity to the B14, B37, and B73 groups, as well as $<2\%$ identity to each of the other groups. Of the lines that met this requirement, 21 had been phenotyped and were kept for subsequent analysis. Those BSSS-descended ex-PVP lines are referred to hereafter simply as ex-PVPs. Hierarchical clustering of the ex-PVPs and BSSSC0 lines revealed six of the BSSSC0 lines did not cluster with the others. Those six were removed from further analysis, leaving 41 BSSSC0 lines.

The third group, referred to as “Public,” is comprised of 16 inbred lines from the WiDiv-942 that are known by pedigree to have been developed from the original Stiff Stalk synthetic population.

Best linear unbiased estimators (BLUEs) for all lines in the WiDiv-942 were calculated by the same models used to calculate BLUPs, but with genotype fit as a fixed rather than random effect. BLUEs for the BSSSC0, Public, and ex-PVP lines were compared for each of the 15 tassel morphological traits, and significant differences were evaluated by Tukey’s honest significant difference. (Tukey 1949).

Mapping

Genome-wide association studies (GWAS) for the WiDiv-942 were performed in R (R Core Team 2016) using the multiple locus linear mixed model implemented in farmCPU (Liu et al. 2016). The first five principal components were included to account for spurious associations due to population structure, and the threshold for initial inclusion of SNPs into the model was set to allow an empirical false entry rate of 1% based on 1000 permutations. SNPs with a minor allele frequency $<0.02$ were excluded from analysis, leaving 529,018 SNPs for GWAS. SNPs with a Bonferroni-adjusted $P$-value $<0.05$ were considered significant. Because of missing phenotypic data, not all 942 individuals were included in mapping. For manual traits other than TW, 791 unique individuals were included in GWAS. For image-based traits and TW, 660 individuals were included in the GWAS.

The PHW65 NAM was mapped using stepwise model selection with the GBS SNPs followed by resampling GWAS with the 10.6 million projected parental SNPs, as described in Tian et al. (2011). The stepwise model selection step, performed using the StepwiseAdditiveModelFitterPlugin in TASSEL5 (Bradbury et al. 2007), added SNPs below an entry threshold (set by permutation for each trait, ranging from
9.6 × 10⁻⁵ to 1.5 × 10⁻⁴) and allowed terms to exit the model based on a limit set at twice the entry threshold. Residuals from the final stepwise model were calculated separately for each chromosome, with any SNPs on that chromosome excluded from the model. The residuals for each chromosome were used as the dependent variable in a resampling-based mapping method similar to that described in Valdar et al. (2006) and Tian et al. (2011), implemented with the ResamplingGWASPlugin in TASSEL5 (Bradbury et al. 2007). Briefly, 80% of the PHW65 NAM individuals were randomly selected and forward regression was used to fit SNPs that were associated with the residuals at a significance level <1 × 10⁻⁴. This process was repeated 100 times for each combination of trait and chromosome, and each SNP fitted by the resampling GWAS method was assigned a resampling model inclusion probability (RMIP) between 0 and 1, reflecting the proportion of models in which that SNP was included. A permutation-based threshold of 0.05 kept the number of false positives to ≤5 per trait, or on average, ≤1 false positive genome-wide per permutation.

**Selection statistics**

We used the 41 BSSSC0 and 21 ex-PVP lines to perform genome-wide scans for selection. The BSSSC0 lines represent maize Stiff Stalk germplasm from the early 20th century, while the ex-PVP lines represent the product of ~60 years of commercial maize breeding. We calculated cross population extended haplotype homozygosity (XP-EHH; Sabeti et al. 2007) between the two groups using the WiDiv-942 SNPs with hapbin (Maclean et al. 2015), with the −minmal argument set to 0 and the −scale argument set to 20,000 bp. Only the 877,627 SNPs with no missing data in either subpopulation were used. Scores were binned by taking the maximum XP-EHH score from each nonoverlapping 10 kb window.

Cross-population composite likelihood ratio (XP-CLR; Chen et al. 2010) was calculated with a modified version of XCLR v1.0 (Hufford et al. 2012; https://figshare.com/articles/new_fileset/757736) on the same set of SNPs as XP-EHH, with a grid size of 100 bp, a genetic window size of 0.1 cM, a maximum of 50 SNPs per window, and correlation level of 0.7. The genetic positions of the WiDiv SNPs were estimated by linear interpolation of the NAM genetic map (McMullen et al. 2009), downloaded from https://maizegdb.org/data_center/map. As with XP-EHH, the resulting XP-CLR values were also binned by using the maximum score in each nonoverlapping 10 kb window to represent that window.

Fixation index (FST) between the two groups was calculated on the same set of SNPs as XP-EHH and XP-CLR, using http://beissingerlab.github.io/docs/vectorFst.R (Beissinger et al. 2014), with a correction for small number of populations and uneven sample sizes according to Weir and Cockerham (1984). As with XP-EHH and XP-CLR scores, FST values were binned into 10 kb windows represented by the maximum score in that window.

XP-EHH, XP-CLR, and FST scores were assigned to each SNP significantly associated with each of the 15 traits by using the score for the 10 kb window to which each SNP belonged. The distribution of those scores were tested against null distributions generated by two different methods: circular permutation and window-matching permutation.

Because the distributions of selection statistics for tassel-associated SNPs could be affected by having certain SNPs physically near each other or represented more than once (associated with more than one trait), we used 10,000 iterations of a circular permutation method (Wallace et al. 2014) to create a null distribution for each selection statistic. Circular permutation is done by keeping the order and relative position of all the hits intact, while randomizing their starting position along the chromosome. The relationships between the tassel-associated SNPs are maintained, but their relationship to selection statistics along the chromosome is randomized.

Because the distribution of selection statistics in tassel-associated windows could also be an artifact of other window characteristics, we created a second null distribution for each selection statistic by randomly selecting windows from the rest of the genome that had similar characteristics to the tassel-associated windows. For each tassel-associated window, 50 windows with a similar number of genes (±1 gene), recombination rate (±20%), frequency of alleles in the BSSSC0 lines (±15%), and frequency of alleles in the ex-PVP lines (±15%) were randomly sampled without replacement. For tassel-associated windows with <50 matching windows genome-wide, the allele frequency criteria were omitted.

The distributions of selection statistics for the tassel-associated SNPs were compared to both null distributions both visually and using the Kolmogorov–Smirnov test with a two-sided P-value (Кольмогоров 1933; Smirnov 1948).

We also investigated whether the distribution of selection statistics for tassel-associated SNPs was being driven by traits characteristic of the ex-PVPs or the BSSSC0 inbreds. TL, TLP, SL, SLP, BZ, and SP are all related to the length of the tassel and, with the exception of BZ, have higher values in the ex-PVPs. BN, BNP, BD, CP, FD, SK, PR, TW, and TWP are more related to the quantity and density of branches, and have higher values in the BSSSC0 lines. The distributions of XP-EHH scores for these “length” and “branchiness” traits were examined separately to see if one or the other type of trait had undue influence on the overall distribution.

**Data availability**

Tassel images of both populations and genotypic data for the PHW65 NAM are available on the CyVerse Data Store (https://doi.org/10.7946/P2K04H) and genotypic data for the WiDiv-942 can be found at https://doi.org/10.5061/dryad.6tg356 (M. Mazaheri, M. Heckwolf, B. Vaillancourt, J. Gage, B. Burdo, S. Heckwolf, K. Barry, A. Lipzen, C. Bastos Riviero, T. Kono, H. Kaeppler, E. Spalding, C. Hirsch, C. R. Buell, J. Gage, B. Burdo, S. Heckwolf, K. Barry, A. Lipzen, C. Bastos Riviero, T. Kono, H. Kaeppler, E. Spalding, C. Hirsch, C. R. Buell, ...
N. de Leon, and S. Kaeppler, unpublished data). Code is available on GitHub at github.com/joegage/tassel_selection. Supplemental material available at Figshare: https://doi.org/10.25386/genetics.7097813.

Results

Tassels exhibit morphological diversity for traits measured manually and by image-based methods

Manual measurements of TL, SL, BN, BD, BZ, SP, and TW for >22,000 individual tassels were recorded for two populations, an association panel (WiDiv-942; M. Mazaheri, M. Heckwolf, B. Vaillancourt, J. Gage, B. Burdo, S. Heckwolf, K. Barry, A. Lipzen, C. Bastos Riviero, T. Kono, H. Kaeppler, E. Spalding, C. Hirsch, C. R. Buell, N. de Leon, and S. Kaeppler; unpublished data) and an NAM population (PHW65 NAM) with PHW65 as the common parent and PHN11, MoG, and Mo44 as founder parents. These materials were grown in three environments between 2013 and 2015 in south central Wisconsin. The WiDiv-942 association panel used in this study is comprised of 942 diverse inbred lines, while the PHW65 NAM consists of three nested biparental populations of 200 individuals each. The WiDiv-942 was chosen to capture broad phenotypic and genetic diversity, whereas the PHW65 NAM leverages controlled biparental crosses between individuals with contrasting tassel morphologies. All tassels manually measured in 2015 were also photographed and measured using the image-based tassel phenotyping system TIPS (Gage et al. 2017), resulting in image-based measurements of TL, SL, BN, and TW (referred to as TLp, SLp, BNp, and TWp, respectively), along with four other traits: CP, FD (a measure of overall complexity), SK (a measure of total linear length), and PR (the length of a tassel’s two-dimensional outline). Accuracy (Pearson’s correlation) of image-based measures for the four traits measured by both manual and image-based methods ranged from 0.71 (BN/BNp) to 0.92 (TW/TWp) in the PHW65 NAM population and from 0.81 (BN/BNp) to 0.89 (TL/TLp and TW/TWp) in the WiDiv-942 population.

All 15 tassel morphological traits measured manually or by image-based phenotyping showed variability for raw measurements within both the WiDiv-942 and PHW65 NAM populations (Figure S1 and Table 1). Excluding FD, which ranged from 1.1 to 1.6, the other 14 traits exhibited between twofold (SLp) and 36-fold (BNp) variability within the PHW65 NAM population and between twofold (SP) and 22-fold (SK) variability within the WiDiv-942 population.

The 15 traits can be broadly classified into two groups: those that contribute to tassel size by contributing to length (TL, TLp, SL, SLp, SP, and BZ) and those that contribute to tassel size by contributing to branchiness (BN, BNp, BD, CP, FD, SK, PR, TW, and TWp). Although TW and TWp are measures of mass rather than branching, both traits are affected more by changes in branchiness than changes in TL and therefore are included in the former group.

For most traits, the PHW65 NAM population has a higher or equivalent mean to the WiDiv-942 population, with the exception of TWp (0.79 g lighter), BN (1.83 fewer branches), BNp (1.07 fewer branches), and BD (0.24 fewer branches/cm). As expected, given the allelic representation contained in each of the populations, phenotypic variability within the PHW65 NAM is lower than in the WiDiv-942, with the exception of TW (SD 0.3 g greater) and BNp (SD 0.07 branches greater).

Heritabilities for manually measured traits ranged from 0.93 (BZ, BD, and SP) to 0.97 (BN) in the WiDiv-942 population and from 0.72 (BZ and SP) to 0.94 (BN and TW) in the PHW65 NAM population. Because images and TW were taken only at one location, repeatability was calculated for image-based traits and TW in place of heritability. Repeatabilities ranged from 0.72 (PR) to 0.86 (TWp) in the WiDiv-942 population and 0.63 (SLp and TLp) to 0.72 (SK) in the PHW65 NAM population (Table 1). The high heritabilities for measurements taken in multiple environments indicate large genotypic variance relative to genotype-by-environment variance and error. Similarly, high repeatabilities for traits that were measured in a single environment indicate large genotypic variance relative to error. It is unsurprising that the image-based repeatabilities were lower than the manually measured heritabilities, as manual measurements were taken in three separate environments while image-based traits were measured in one environment.

Genotypic and phenotypic correlations were calculated within the single environment in which all traits were measured (Figures S2 and S3). With the exception of CP, branchiness traits were generally positively correlated with each other in both populations. Similarly, length-related traits were all positively correlated with each other in both populations. Phenotypic and genotypic correlations were similar to each other in both populations, partly due to relatively high heritability of all traits and partly due to having been calculated within a single location.

Tassel morphology has changed over the course of 20th century maize breeding

The WiDiv-942 association panel contains 41 inbred lines developed by selfing without selection from cycle 0 of the BSSS population (BSSSC0), 16 public inbred lines with pedigrees that trace entirely back to lines derived from the BSSS, and 21 ex-PVPs with genotypic evidence for descent from the BSSS (File S1). These three groups of lines allow comparison of tassel morphology over the course of modern maize breeding. The BSSSC0 lines represent the earliest material, the ex-PVPs represent the most recent and heavily selected material, while the public lines form an intermediate group.

Comparison of BLUEs for each group revealed that the mean values of public inbred BLUEs for each trait showed between a 2% (FD) and a 16% (TWp and BNp) change relative to the BSSSC0 line, while ex-PVP lines displayed between a 4% (TLp) and a 45% (TWp) change in mean relative to the BSSSC0 lines (Figure 1). For all traits except CP, the public
from RNA sequencing with minor allele frequency (SNP: one SNP for TL and TLp, four for BN and BNp, one for SL). Assures of the same phenotype were associated with the same repeatability. In 10 instances, manual and image-based measurements were expected based on the relative measures of heritability and significance associated across all traits (File S2).

Traits were mapped in the PHW65 NAM population using a resampling-based GWAS approach (Tian et al. 2011), with 529,018 SNPs discovered from RNA sequencing with minor allele frequency >0.02. Traits were mapped in the PHW65 NAM population using a resampling-based GWAS approach (Tian et al. 2011), with 10.6 million resequencing SNPs projected from parental lines onto the progeny, which were genotyped at 9291 GBS SNPs. In the PHW65 NAM population, 87 SNPs were identified as significantly associated with 12 of the 15 traits. In the PHW65 NAM population, 155 SNPs were identified as significantly associated across all traits (File S2).

The manually measured traits returned a greater number of significant associations in total than the image-based traits, as expected based on the relative measures of heritability and repeatability. In 10 instances, manual and image-based measures of the same phenotype were associated with the same SNP: one SNP for TL and TLp, four for BN and BNp, one for SL and SLp, and five for TW and TWp. One pair of SNPs each for BN and weight were identified in the WiDiv-942 population, and the rest of the colocalized SNPs were discovered in the PHW65 NAM population. The single SNP associated with TL was the same as the single SNP associated with SL; a SNP associated with weight and a SNP associated with BN were within 2.5 kb of each other. In two other instances, the manually measured and image-based traits were associated with SNPs within 5 kb of each other, and in an additional two instances they were associated with SNPs that were located within 150 kb of each other (Figure S4). Generally, there is more concordance between SNPs associated with manual and image-based measurements for traits with higher heritabilities/repeatabilities (BN and TW) than lower heritabilities/repeatabilities (SL and TL). This is to be expected, as genetic signal is stronger when heritability is high, but the pattern seen in this study may be due only to chance as BN and TW are only marginally more heritable than SL and TL.

GWAS identifies SNPs associated with tassel morphological traits

All 15 traits were mapped in the WiDiv-942 association panel using the multiple locus linear mixed model implemented in FarmCPU (Liu et al. 2016), with 529,018 SNPs discovered from RNA sequencing with minor allele frequency >0.02. Traits were mapped in the PHW65 NAM population using a resampling-based GWAS approach (Tian et al. 2011), with 10.6 million resequencing SNPs projected from parental lines onto the progeny, which were genotyped at 9291 GBS SNPs. In the WiDiv-942 population, 87 SNPs were identified as significantly associated with 12 of the 15 traits. In the PHW65 NAM population, 155 SNPs were identified as significantly associated across all traits (File S2).

The manually measured traits returned a greater number of significant associations in total than the image-based traits, as expected based on the relative measures of heritability and repeatability. In 10 instances, manual and image-based measures of the same phenotype were associated with the same SNP: one SNP for TL and TLp, four for BN and BNp, one for SL and SLp, and five for TW and TWp. One pair of SNPs each for BN and weight were identified in the WiDiv-942 population, and the rest of the colocalized SNPs were discovered in the PHW65 NAM population. The single SNP associated with TL was the same as the single SNP associated with SL; a SNP associated with weight and a SNP associated with BN were within 2.5 kb of each other. In two other instances, the manually measured and image-based traits were associated with SNPs within 5 kb of each other, and in an additional two instances they were associated with SNPs that were located within 150 kb of each other (Figure S4). Generally, there is more concordance between SNPs associated with manual and image-based measurements for traits with higher heritabilities/repeatabilities (BN and TW) than lower heritabilities/repeatabilities (SL and TL). This is to be expected, as genetic signal is stronger when heritability is high, but the pattern seen in this study may be due only to chance as BN and TW are only marginally more heritable than SL and TL.

Selection statistics show evidence for selection at SNPs associated with tassel morphology

To test for genomic signatures of selection on tassel morphology, SNPs associated with tassel characteristics in the WiDiv-942 and PHW65 NAM lines were used as a priori candidates for regions having undergone selection between the BSSSC0 lines and the ex-PVP lines. Scans for selection were conducted using LD-focused methods XP-EHH (Sabeti et al. 2007) and XP-CLR (Chen et al. 2010), and the allelic frequency change based method FST (Weir and Cockerham 1984). Genome-wide results from selection scans were binned into 10 kb windows, and windows containing SNPs associated with tassel morphology were compared to the genome-wide distribution of each statistic. Of the 242 total SNP-trait associations identified, 170 were in windows that had XP-EHH scores, 231 had XP-CLR scores, and 169 had FST scores.
scores. Not all SNP-trait associations had scores for each selection statistic because the associations were discovered in the full PHW65 NAM and WiDiv-942 populations, but may lie in regions that are monomorphic between the BSSSC0 lines and ex-PVP individuals or contain missing SNP calls. The distributions of tassel-associated XP-EHH, XP-CLR, and FST scores were significantly different from genome-wide null distributions generated from circular permutations (Kolmogorov–Smirnov test, \( P = 1.8 \times 10^{-2}, 7.8 \times 10^{-3}, \) and \( 1.9 \times 10^{-3}, \) respectively), as well as genome-wide null distributions generated by randomly sampling windows with similar characteristics (\( P = 3.9 \times 10^{-2}, 1.2 \times 10^{-3}, \) and \( 2.92 \times 10^{-3}, \) respectively), suggesting enrichment for selection signals in regions associated with tassel morphology.

XP-EHH scores for tassel-associated windows (Figure 2) were enriched for values approximately >1 and < −1, with the majority of the enrichment coming from negative scores (Figure 3 and Figure S5). Positive XP-EHH scores indicate longer, more prevalent haplotypes in the collection of lines from the BSSSC0 population, while negative XP-EHH scores indicate longer, more prevalent haplotypes in the ex-PVP set. The greater surplus of negative scores among the tassel-associated SNPs indicates extended haplotypes at those loci in the newer ex-PVP population, relative to the ancestral BSSS. The extended haplotypes in the ex-PVP population are consistent with what would be expected if beneficial alleles were recently selected, resulting in a concurrent increase in linked haplotypes due to genetic hitchhiking (Voight et al. 2006). Further, the excess of negative XP-EHH scores was driven largely by associations with phenotypes related to branchiness rather than length (Figure 4; one-sided Wilcoxon rank sum test, \( P = 2.9 \times 10^{-3} \)). Branchiness traits have more
influence on tassel size and biomass than tassel and spike length. With the exception of BZ, all the length traits increased from the BSSSC0 to the ex-PVPs, whereas all the branchiness traits decreased. As such, most of the SNPs showing evidence of selection in the ex-PVPs are associated with traits that control overall tassel size and biomass and have decreased significantly since the 1930s.

Similar to XP-EHH, XP-CLR is a haplotype-based method for identifying selective sweeps. Rather than identify alleles that have recently arisen to substantial frequency, as XP-EHH does, XP-CLR identifies alleles that have changed in frequency more quickly than expected under neutrality (Chen et al. 2010). Both methods use LD around the tested allele as a measure of the allele's age. Tassel-associated XP-CLR scores (Figure 2) show enrichment for values in the tail of the distribution (Figure 3 and Figure S5), indicating an excess of regions that may have been subject to selective sweeps.

FST, a measure of population differentiation based on allele frequencies, differs from the previous two selection statistics in that it is not haplotype based. FST values for tassel-associated windows (Figure 2) are generally higher than expected under the null distribution, with a depletion of values less than ~0.3 and enrichment for values greater than ~0.3 (Figure 3 and Figure S5).

None of these three selection statistics show strong, systematic relationships with each other on a genome-wide basis (Figure S6) or among the tassel-associated windows (Figure S7). For the 101 significant associations that were genotyped in the WiDiv-942 (and hence, in the ex-PVP and BSSSC0 lines), we hypothesized that alleles with higher frequency in the ex-PVPs would generally have allelic effects conferring more ex-PVP-like phenotypes, that is, positive values for TL, SL, TLp, SLp, and SP, and negative values for all other traits. However, we found no difference between the effect sizes of alleles with higher frequency in the ex-PVPs and alleles with higher frequency in the BSSSC0s (Welch’s two-sampled t-test, \( P = 0.87 \)). The lack of any difference could be due to the low number of SNPs we were able to directly test, or perhaps due to noise introduced by identifying SNPs that are not causative themselves but in LD with variants that directly affect tassel morphology.

Discussion

Tassel morphology of modern commercial inbred lines, which are used as parents of single cross hybrids grown in the United States Corn Belt, is dramatically different from tassel morphology seen in the open-pollinated varieties of the pre-hybrid era. We demonstrate that tassel morphology has been continually modified, with length-related traits increasing and branchiness traits decreasing. This trend is consistent between BSSSC0 inbreds and public BSSS-derived inbreds, as well as between public inbreds and BSSS-derived ex-PVPs. Through mapping of both image-based and manually measured phenotypes, we identify SNPs associated with tassel morphological traits, which collectively show enrichment for three selection statistics.

By studying traditionally measured traits, such as TL and BN, as well as traits that cannot be accurately measured manually, such as CP and FD, we generate a more comprehensive and nuanced representation of tassel morphology than in previous tassel studies. Previous QTL mapping and GWAS of maize tassel morphology have primarily emphasized tassel branching and length (e.g., Berke and Rocheford 1999;
Mickelson et al. 2002; Upadyayula et al. 2006; Brown et al. 2011; Wu et al. 2016). The novel, image-based traits used in this study display considerable variability, have high enough repeatabilities to be useful in mapping studies, and resulted in significant phenotype-genotype associations that contributed to our analysis of selection signals. The ability to add information to manual phenotyping is especially influential when manual phenotyping is laborious or inefficient. For phenotyping tassels, manual measurements are relatively easy and fast to obtain. However, image-based phenotyping measures the same traits but also contributes, with no additional time investment, supplementary measurements such as CP and FD, which would be difficult, subjective, or impossible to measure otherwise. Although tassels for this study were removed from the plants before imaging (a laborious task), such methods are a step toward higher-throughput methodologies, such as image acquisition by unmanned aerial vehicles.

By comparing tassel morphology in the BSSSC0 and ex-PVP sets considered in this study, we are able to compare two groups that bookend modern breeding of one of the most important heterotic groups in North American maize germplasm. The BSSSC0 and ex-PVP groups are characterized by unique and differing tassel morphology: relative to the BSSSC0, ex-PVP tassels are longer, less branched, and weigh less. These changes in morphology reflect an overall trend toward a tassel ideotype that has few lateral branches and produces pollen primarily by means of an extended central spike. The Stiff Stalks, traditionally used as females in breeding programs, need only produce enough pollen for self-fertilization during inbred line development and seed increases; this is reflected by the reduction of their size and weight since the inception of inbred-driven breeding schemes in the 1930s. Reduction in tassel size may have caused a decrease in the duration of pollen shed, contributing to the shortening of anthesis-silking interval between the 1930s and 1970s (Meghji et al. 1984). Previous experiments that recurrently selected the BSSS population for grain yield also observed a decrease in BN (Edwards 2011), further reinforcing the relationship between agronomic performance and reduced tassel size. Although the changes in tassel morphology caused by recent selection are well characterized, the genetic effects of selection by breeders are less studied.

Previous works have shown evidence for regions selected during domestication in animals such as pigs and chickens (Rubin et al. 2010; Yang et al. 2014); and in plants such as maize and soybean (Hufford et al. 2012; Zhou et al. 2015). Evidence has been found for differential artificial selection between breeds of cattle (Rothammer et al. 2013), as well as for postdomestication improvement in maize and soybean (Hufford et al. 2012; Zhou et al. 2015), but the latter two characterize selection between landraces and improved varieties, which cover a much larger period of selection than just modern breeding. Work in rice (Xie et al. 2015) has identified
overlap between inflorescence morphology QTL and regions showing evidence of selection in the last 50 years of breeding, but the genetic relationship between recent selection and inflorescence morphology in maize is relatively unexplored.

XP-EHH, XP-CLR, and FST scores for SNPs associated with tassel morphology traits showed deviations from permuted genome-wide null distributions, with enrichment for values that are indicative of selection. In de novo scans for selected sites, often the threshold for determining selection candidates by XP-CLR or FST is set as a quantile of the empirical distribution that limits candidates to loci in the top 0.1% or 0.01% of results (e.g., Beissinger et al. 2014; Jeong et al. 2015). XP-EHH has the agreeable property of being approximately normally distributed under a model of no selection, which allows identification of selection candidates based on P-values from the normal distribution (Sabeti et al. 2007). While not all tassel-associated SNPs have high enough XP-EHH, XP-CLR, and FST scores to be identified as selection candidates in a de novo search for selected sites, the strong enrichment for elevated scores suggests identification of regions that might be experiencing weak or ongoing selection. Genome-wide scans for selection are complicated by population dynamics and demographics that differ from expectations, and often result in identification of large genomic regions containing many features that require further investigation. Instead of approaching such an analysis with no expectations, it has been suggested (Haasl and Payseur 2016) that creating an a priori list of candidate genes can help narrow down significant results. In this study, we used associations from two comprehensive genetic mapping resources as a priori candidates to identify signatures of selection associated with a specific suite of phenotypic traits.

XP-EHH identifies putatively selected sites by identifying regions where one population contains longer haplotypes around a target SNP, which have arisen due to hitchhiking with a selected locus. We observe enrichment for both positive and negative XP-EHH scores; positive scores indicate greater haplotype homozygosity in the BSSSC0 lines, whereas negative scores indicate greater haplotype homozygosity in the ex-PVP lines. There is a substantially larger enrichment for negative XP-EHH scores than positive, consistent with the hypothesis that favorable alleles from the BSSSC0 lines were selected in the ex-PVPs, resulting in increased LD in the surrounding areas. The majority of enrichment for negative XP-EHH scores appears to be driven by branchiness traits, which may have been easier to select on (intentionally or not) by breeders aiming to reduce overall tassel size. The enrichment for positive XP-EHH scores, although not as extreme as for negative scores, indicates a number of tassel-associated loci with slower LD decay in the BSSSC0 lines than in the ex-PVP set. One possible explanation for this observation is that prior to the creation of the BSSS, during a period characterized by open-pollinated populations, heavier tassels with more branches had a reproductive advantage due to higher pollen production. During the last century of maize breeding the selective pressure on high pollen production has been removed by a hybrid breeding scheme in which inbred combining ability and hybrid grain yield are prioritized. It is possible that the BSSSC0 lines show extended haplotypes due to selection favoring alleles that confer large tassel size, whereas modern breeding schemes have removed that selective pressure and allowed or encouraged recombination in previously selected regions.

The results presented here provide evidence for selection by 20th century maize breeders on regions associated with maize tassel morphology. Whether these changes are the result of direct or indirect selection on tassel size and shape is unclear. Modern, commercially developed maize inbreds display a suite of characteristics that have systematically changed since the open-pollinated varieties of the 1930s, and it is difficult to disentangle selection for changes in tassel morphology from selection for traits that are regulated by loci pleiotropically involved in tassel morphology. Because the tassel and the ear follow a shared developmental path early in development, genes affecting morphology of one may very well have effects on the other. QTL have been discovered that exhibit pleiotropic behavior, conferring a positive relationship between tassel and ear length (Brown et al. 2011). Mutants of the inflorescence genes unbranched2 and unbranched3 have been shown to simultaneously decrease branch number and increase kernel row number, a trait that directly affects grain yield (Chuck et al. 2014), while mutants of sea floricaula/leafy2 decrease both branch number and kernel row number (Bomblies and Doebley 2006). There is also evidence for shared genetic control of tassel traits and other plant architecture traits such as leaf length (Brown et al. 2011) and leaf angle (Mickelson et al. 2002), the latter of which has also changed significantly over the course of modern maize
breeding (Duvick 2005). Although more work is needed to clarify the genetic relationships between selection and maize tassel morphology, particularly with respect to disentangling pleiotropic effects, this study is a starting point for discovering the effects artificial selection has on genetic control of inflorescences in agricultural crops.

Change in tassel size and shape between the early 20th century and modern day is a well-characterized phenomenon but the causes of that phenomenon and its potential causal effects on other agronomic traits and productivity are not fully known. Our results present evidence for enriched signals of selection in genomic regions associated with tassel morphological traits. Further, this selection took place in ~60 years between the creation of the BSSS in the 1930s and the release of ex-PVP lines in the 1980s and 1990s, a tremendous example of the efficiency of modern crop breeding in imposing drastic genetic and phenotypic changes.

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