Maturity2, a novel regulator of flowering time in *Sorghum bicolor*, increases expression of *SbPRR37* and *SbCO* in long days delaying flowering

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Abstract

*Sorghum bicolor* is a drought-resilient facultative short-day C4 grass that is grown for grain, forage, and biomass. Adaptation of sorghum for grain production in temperate regions resulted in the selection of mutations in *Maturity* loci (*Ma*1–*Ma*6) that reduced photoperiod sensitivity and resulted in earlier flowering in long days. Prior studies identified the genes associated with *Ma*1 (*PRR37*), *Ma*3 (*PHYB*), *Ma*5 (*PHYC*) and *Ma*6 (*GHD7*) and characterized their role in the flowering time regulatory pathway. The current study focused on understanding the function and identity of *Ma*2. *Ma*2 delayed flowering in long days by selectively enhancing the expression of *SbPRR37* (*Ma*1) and *SbCO*, genes that co-repress the expression of *SbCN12*, a source of florigen. Genetic analysis identified epistatic interactions between *Ma*2 and *Ma*4 and located QTL corresponding to *Ma*2 on SBI02 and *Ma*4 on SBI10. Positional cloning and whole genome sequencing identified a candidate gene for *Ma*2, Sobic.002G302700, which encodes a SET and MYND (SYMD) domain lysine methyltransferase. Eight sorghum genotypes previously identified as recessive for *Ma*2 contained the mutated version of Sobic.002G302700 present in 80M (*ma*2) and one additional putative recessive *ma*2 allele was identified in diverse sorghum accessions.

Introduction

*Sorghum bicolor* is a drought resilient, short-day C4 grass that is grown globally for grain, forage, and biomass [1–4]. Precise control of flowering time is critical to achieve optimal yields of sorghum crops in specific target production locations/environments. Sorghum genotypes that have delayed flowering in long days due to high photoperiod sensitivity are high-yielding sources of biomass for production of biofuels and specialty bio-products [3,5]. In contrast, grain sorghum was adapted for production in temperate regions by selecting genotypes that have reduced photoperiod sensitivity resulting in earlier flowering and reduced risk of...
exposure to drought, heat, or cold temperatures during the reproductive phase. A range of flowering times are found among forage and sweet sorghums [6]. Sweet sorghum genotypes with longer vegetative growth duration have larger stems that have greater potential for sucrose accumulation [6–8].

Flowering time is regulated by development, day length, phytohormones, shading, temperature, and the circadian clock [9–11]. In the long-day plant Arabidopsis thaliana, circadian and light signals are integrated to increase the expression of FLOWERING LOCUS T (FT) and flowering in long days. FT encodes a signaling protein synthesized in leaves that moves through the phloem to the shoot apical meristem (SAM) where it interacts with FLOWERING LOCUS D (FD) and reprograms the vegetative shoot apical meristem for reproductive development [12,13]. Expression of circadian clock genes such as LATE ELONGATED HYPOCOTYL (LHY) and TIMING OF CAB1 (TOC1) regulate the expression of the clock output gene GIGANTEA (GI) and genes in the flowering time pathway [14–16]. Photoperiod and circadian clock signals are integrated to control the expression and stability of CONSTANS (CO) an activator of FT expression [17]. Under inductive long day (LD) photoperiods, CO promotes the expression of FT which induces flowering in Arabidopsis [18].

Many of the genes in the Arabidopsis flowering time pathway are found in grass species such as Oryza sativa (rice) [10], maize [19], and sorghum [3] however, the regulation of flowering time in these grasses has diverged from Arabidopsis in several important ways. Most genotypes of rice and sorghum are facultative short-day (SD) plants. In rice, the expression of the FT-like gene Heading date 3a (Hd3a) is promoted in SD [20]. In sorghum, expression of two different FT-like genes, SbCN8 and SbCN12, is induced when plants are shifted from LD to SD [21,22]. In contrast to Arabidopsis, the rice and sorghum homologs of CO (rice Heading date1, OsHd1; SbCO) repress flowering in LD [10,23]. Rice and sorghum encode two additional grass-specific regulators of flowering Ehd1 and Ghd7. Early heading date1 (Ehd1) activates the expression of FT-like genes, and Grain number, plant height and heading date7 (Ghd7) represses the expression of EHD1 and flowering [24,25]. When sorghum is grown in short days, SbEhd1 and SbCO induce the expression of SbCN8 and SbCN12, leading to floral induction [21,22,26,27].

Under field conditions, time to flowering in sorghum varies from ~50 to >150 days after planting (DAP) depending on genotype, planting location and date (latitude/day-length), and the environment. A tall and “ultra-late” flowering sorghum variety called Milo Maize was introduced to the United States in the late 1800s [28]. Shorter and earlier flowering Milo genotypes such as Early White Milo and Dwarf Yellow Milo were selected from the introduced Milo genotype to promote improved grain yield in temperate regions of the US [1,28,29]. Both of these Milo genotypes were later found to encode the same Ghd7 allele (ghd7-1) containing a stop codon [26]. Genetic analysis determined that mutations in three independently segregating Maturity (Ma) loci (Ma1, Ma2, Ma3) were responsible for variation in flowering times in the Milo genotypes. A cross between Early White Milo (ma1ma2ma3) and Dwarf Yellow Milo (Ma,ma2ma3) was used to construct a set of Milo maturity standards (i.e., 100M, SM100, 80M, SM80), a series of nearly isogenic lines that differ at one or more of the Maturity loci (Quinby and Karper 1945, Quinby 1966, Quinby, 1967). A fourth Maturity locus (Ma4) was discovered in crosses of Milo (Ma1) and Hegari (ma4) [30]. More recent studies identified Ma5 and Ma6 segregating in other sorghum populations [31]. Subsequent research showed that all of the Milos are dominant for Ma5 and recessive for ma6 (ghd7-1) [23,26]. In addition to these six Ma loci, many other flowering time quantitative trait loci (QTL) have been identified in sorghum [2,32–35]. Additional research has linked several of these QTL to genes such as SbEHD1 and SbCO that are potential activators of SbCN8 and SbCN12 expression, sources of florigen in sorghum.
The genes corresponding to four of the six Maturity loci have been identified. Ma₁, the locus with the greatest influence on flowering time photoperiod sensitivity, encodes SbPRR37, a pseudo-response regulator that inhibits flowering in LD [21]. Ma₃ encodes phytochrome B (phyB) [36], Ma₅ encodes phytochrome C (phyC) [23], and Ma₆ encodes Gh7 a repressor of flowering in long days [26]. The genes corresponding to Ma₂ and Ma₄ have not been identified but recessive alleles at either locus results in early flowering in long days in sorghum lines that are photoperiod sensitive and have Ma₁ genotypes [28]. Prior studies also noted that genotypes recessive for Ma₂ flower later in genotypes that are photoperiod insensitive and recessive for Ma₁ and Ma₆ [28].

In this study, the impact of Ma₂ alleles on the expression of genes in the sorghum flowering time pathway was characterized. A QTL corresponding to Ma₂ was mapped and a candidate gene for Ma₂ identified by fine mapping and genome sequencing. The results show that Ma₂ enhances SbPRR37 (Ma₁) and SbCO expression consistent with the impact of Ma₂ alleles on flowering time in genotypes that vary in Ma₁ alleles.

Methods

Plant growing conditions and populations

Seeds for all genotypes used in this study were obtained from the Sorghum Breeding Lab at Texas A&M University in College Station, TX. 100M (Ma₁Ma₂Ma₃Ma₄Ma₅ma₆) and 80M (Ma₁ma₂Ma₃Ma₄Ma₅ma₆) are sorghum maturity standards with defined maturity/flowering genotypes [1]. The maturity genotypes were selected from a cross between Early White Milo (ma₁Ma₃Ma₄Ma₅ma₆) and Dwarf Yellow Milo (Ma₁ma₂ma₃Ma₅ma₆). 100M and 80M are nearly isogenic and differ at Ma₂.

The cross of 100M and 80M was carried out by the Sorghum Breeding Lab at Texas A&M University in College Station, TX. F₁ plants were grown in the field in Puerto Rico and self-pollinated to generate the F₂ population used in this study. The 100M/80M F₂ population was planted in the spring of 2008 at the Texas A&M Agrilife Research Farm in Burleson County, Texas (near College Station, TX).

The cross of Hegari and 80M was made in the greenhouse at Texas A&M University in College Station, TX. F₁ plants were confirmed and self-pollinated to generate the F₂ population used in this study. The Hegari/80M F₂ population (n = 432) was planted in the spring of 2011 in the greenhouse in 18 L nursery pots in a 2:1 mixture of Coarse Vermiculite (SunGro Horticulture, Bellevue, WA) to brown pasture soil (American Stone and Turf, College Station, TX). All subsequent generations of Hegari/80M for fine mapping were grown in similar conditions. Greenhouse-grown plants were watered as needed and fertilized every two weeks using Peters general purpose 20-20-20 (Scotts Professional).

For circadian gene expression experiments, 100M and 80M genotypes were planted in Metro-Mix 900 (Sungro Agriculture) in 6 L pots, and thinned to 3 plants/pot after 2 weeks. Plants were grown in the greenhouse under 14 h days until 30 days after planting (DAP). After 30 days, the plants were moved into growth chambers and allowed to acclimate for 3 days. The growth chamber was set to 30°C and 14/10h Light/Dark (L/D) for the 3 days of entrainment and the first 24 h of tissue collection. The lights were changed to constant light for the second 24 h of tissue collection.

QTL mapping and multiple-QTL analysis

DNA was extracted from leaf tissue for all individuals described above as described in the FastDNA Spin Kit manual (MP Biomedicals). All individuals in each mapping or heterogeneous inbred family (HIF) population were genotyped by Digital Genotyping using FseI digestion enzyme as described in Morishige et al [37]. DNA fragments were sequenced using the Illumina GAII platform and the reads were mapped back to the sorghum reference genome (v1.0,
Phytozome v6). Genetic maps were created using MapMaker 3.0B with the Kosambi function [38]. QTL were mapped using WinQTLCartographer (v2.5.010) using composite interval mapping with a 1.0 cM walk speed and forward and backward model selection [39]. The threshold was set using 1000 permutations and \( \alpha = 0.05 \). Upon release of v3.1 of the sorghum reference genome, the QTL coordinates were updated [40].

To look for possible gene interactions multiple-QTL analysis was used in the Hegari/80M F\(_2\) population. A single QTL analysis using the EM algorithm initially identified two primary additive QTL which were used to seed model selection. The method of Manichaikul et al. [41] was employed for model selection as implemented in R/qtl for multiple-QTL analysis [42]. Computational resources on the WSGI cluster at Texas A&M were used to calculate the penalties for main effects, heavy interactions, and light interactions. These penalties were calculated from 24,000 permutations for flowering time to find a significance level of 5% in the context of a two-dimensional, two-genome scan.

**Fine mapping of the \( \text{Ma}_2 \) QTL**

All fine mapping populations for the \( \text{Ma}_2 \) QTL were derived from F\(_2\) individuals from the Hegari/80M population. The genetic distance spanning the \( \text{Ma}_2 \) locus is 2 cM corresponding to a physical distance of \(~1.8\) Mbp, so 1000 progeny would be required to obtain 20 recombinants within the \( \text{Ma}_2 \) QTL region. Six individuals that were heterozygous across the \( \text{Ma}_2 \) QTL were self-pollinated to generate six HIFs totaling 1000 F\(_3\) individuals. These individuals were grown out in the greenhouse, and flowering time was recorded. They were genotyped by Digital Genotyping as described above [37]. Two F\(_3\) individuals that had useful breakpoints with a heterozygous genotype on one side of the breakpoint were grown and self-pollinated to generate an additional round of HIFs (F\(_4\), n = 150) that were planted in the spring of 2013 and analyzed as described above. No new breakpoints were identified in the F\(_4\) generation, so this process was repeated again to generate F\(_5\) plants in the spring of 2014.

**Circadian gene expression analysis**

For the circadian gene expression analysis, 30-day-old plants were placed in a growth chamber set to 14h/10h L/D for the first 24 h and constant light for the second 24 h at 30˚C. Plants were entrained for 3 d under these growth chamber conditions before beginning tissue collection. Leaf tissue was collected and pooled from 3 plants every 3 h for 48 h. The first sample was taken at lights-on on the first day of sample collection. The experiment was repeated three times for a total of three biological replicates. RNA was extracted from each sample using the Direct-Zol RNA Miniprep Kit (Zymo Research) according to the kit instructions. cDNA was synthesized using SuperScript III kit for qRT-PCR (Invitrogen) according to the kit instructions. Primers for sorghum flowering pathway genes were developed previously, and primer sequences are available in Murphy et al [21]. Primer sequences for \( \text{Ma}_2 \) are available in S1 Table. Relative expression was determined using the comparative cycle threshold (C\(_t\)) method. Raw C\(_t\) values for each sample were normalized to C\(_t\) values for the reference gene \( \text{SbUBC} \) (Sobic.001G526600). Reference gene stability was determined previously [43]. \( \Delta\Delta\text{C}_t \) values were calculated relative to the sample with the highest expression (lowest C\(_t\) value). Relative expression values were calculated with the \( 2^{-\Delta\Delta\text{C}_t} \) method [44]. Primer specificity was tested by dissociation curve analysis and gel electrophoresis of qRT-PCR products.

**\( \text{Ma}_2 \) phylogenetic analysis**

Protein sequences of the closest homologs of \( \text{Ma}_2 \) were identified using BLAST analysis. Protein sequences were aligned using MUSCLE [45] and visualized using Jalview [46].
Evolutionary trees were inferred using the Neighbor-Joining method [47] in MEGA7 [48]. All positions containing gaps and missing data were eliminated.

Ma$_2$ DNA sequencing and whole genome sequence analysis

Whole genome sequence reads of 52 sorghum genotypes including 100M and 80M were obtained from Phytozome v12. Base quality score recalibration, INDEL realignment, duplicate removal, joint variant calling, and variant quality score recalibration were performed using GATK v3.3 with the RIG workflow [49]. Sobic.002G302700 was sequenced via Sanger sequencing in the genotypes in Table 1 according to the BigDye Terminator Kit (Applied Biosystems). Primers for template amplification and sequencing are provided in S1 Table.

Results

Effects of Ma$_2$ alleles on flowering pathway gene expression

The recessive ma$_2$-allele in 80M (Ma$_1$ma$_2$Ma$_3$Ma$_4$Ma$_5$ma$_6$) was previously reported to cause 80M to flower earlier than 100M (Ma$_1$Ma$_2$Ma$_3$Ma$_4$Ma$_5$ma$_6$) in long days [28]. To help elucidate how Ma$_2$ modifies flowering time, we investigated the impact of Ma$_2$ alleles on the expression of genes in sorghum’s flowering time pathway. Gene expression was analyzed by qRT-PCR using RNA isolated from 100M (Ma$_2$) and 80M (ma$_2$) leaves collected every 3 hours for one 14h light/10h dark cycle and a second 24-hour period of constant light.

SbPRR37 is a central regulator of photoperiod sensitive flowering in sorghum that acts by repressing the expression of SbCN (FT-like) genes in LD [21]. SbPRR37 expression in 100M and 80M grown in long days peaked in the morning and again in the evening as previously observed [21] (Fig 1). The amplitude of both peaks of SbPRR37 expression was reduced in 80M (ma$_2$) compared to 100M (Ma$_2$) (Fig 1A). SbCO also shows peaks of expression in the morning (dawn) and in the evening (~14h) in plants grown in LD [21] (Fig 1B). Analysis of SbCO expression in 100M and 80M showed that both peaks of SbCO expression were reduced in 80M compared to 100M (Fig 1B).

SbCN8, SbCN12, and SbCN15 are homologs of AtFT that encode florigens in sorghum [22]. Expression of SbCN8 and SbCN12 increases when sorghum plants are shifted from LD to SD, whereas SbCN15 is expressed at lower levels and shows minimal response to day length [21,26]. SbPRR37 and SbCO are co-repressors of the expression of SbCN8 and SbCN12 in long

| Genotype | Historical Ma$_2$ allele | Sequence variant | Effect on protein function |
|----------|--------------------------|------------------|---------------------------|
| 100M     | Ma$_2$                   | -                | -                         |
| SM100    | Ma$_2$                   | -                | -                         |
| SM90     | Ma$_2$                   | -                | -                         |
| Hegari   | Ma$_2$                   | -                | -                         |
| 80M      | ma$_2$                   | L141’            | Deleterious               |
| SM80     | ma$_2$                   | L141’            | Deleterious               |
| 60M      | ma$_2$                   | L141’            | Deleterious               |
| 44M      | ma$_2$                   | L141’            | Deleterious               |
| 38M      | ma$_2$                   | L141’            | Deleterious               |
| Kalo     | ma$_2$                   | L141’            | Deleterious               |
| IS3614-2 | -                        | M83T             | Deleterious               |

*Sequenced by Sanger sequencing

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days, therefore, the influence of $Ma_2$ alleles on $SbCN8/12/15$ expression was investigated [21,27]. When plants were grown in long days, expression of $SbCN12$ was ~5 fold higher in 80M compared to 100M consistent with earlier flowering in 80M (Fig 2).

Previous studies showed that $SbGhd7$ represses $SbEhd1$ expression and that alleles of $SbGHD7$ differentially affect $SbCN8$ expression (> $SbCN12$) [26]. Analysis of $SbEhd1$ and $SbGHD7$ expression in 100M and 80M showed that $Ma_2$ alleles have a limited influence on the expression of these genes (S1 Fig).

The timing of the two daily peaks of $SbPrr37$ and $Sbco$ expression in sorghum is regulated by the circadian clock [21,26]. Therefore, it was possible that $Ma_2$ modifies $SbPrr37/Sbco$ expression by altering clock gene expression. However, expression of the clock genes $TOC1$ and $LHY$ was similar in 100M and 80M (S1 Fig). Taken together, these results show that $Ma_2$ is an activator of $SbPrr37$ and $SbCO$ expression in long days. Prior studies showed that co-expression of $SbPrr37$ and $SbCO$ in long days inhibits expression of $SbCN12$ and floral initiation [27]. Later flowering in sorghum genotypes that are $Ma_1Ma_2$ vs. $Ma_1ma_2$ in long days is consistent with lower $SbCN12$ expression in $Ma_1Ma_2$ genotypes.

Fig 1. Circadian expression of genes regulating flowering in $S. bicolor$ in 100M and 80M under long days. (A) Expression of $SbPrr37$ in 100M (solid black lines) and 80M (dashed red lines). The expression peaks of $SbPrr37$ are reduced in 80M. This is consistent with earlier flowering in 80M because $SbPrr37$ represses the expression of the sorghum FT-like genes. (B) Expression of $SbCO$ in 100M and 80M. Expression peaks of $SbCO$ are also reduced in 80M. This is consistent with earlier flowering in 80M because under long days $SbCO$ is a repressor of flowering. All expression values are normalized to $SbUBC$ and are the mean of 3 biological replicates.

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Genetic analysis of \( Ma_2 \) and \( Ma_4 \)

An \( F_2 \) population derived from a cross of 100M (\( Ma_2 \)) and 80M (\( ma_2 \)) was generated to map the \( Ma_2 \) locus. Because 100M and 80M are nearly isogenic lines that differ at \( Ma_2 \), only \( Ma_2 \) alleles were expected to affect flowering time in this population [28]. The \( F_2 \) population (\( n = \sim 1100 \)) segregated for flowering time in a 3:1 ratio as expected. The parental lines and \( F_2 \) individuals were genotyped by Digital Genotyping (DG) which identifies single nucleotide polymorphism (SNP) markers in thousands of sequenced sites that distinguish the parents of a population [37]. The near isogenic nature of the parental lines resulted in a very sparse genetic map that lacked coverage of large regions of the sorghum genome including all of the long arm of SBI02. In retrospect, no \( Ma_2 \) QTL for flowering time was identified using this genetic map because the gene is located on the long arm of SBI02 (see below).

To overcome the lack of DNA markers associated with the 80M/100M population, a second mapping population was created to identify the genetic locus associated with \( Ma_2 \) and \( Ma_4 \). An \( F_2 \) population (\( n = 215 \)) that would segregate for \( Ma_2 \) and \( Ma_4 \) was constructed by crossing Hegari (\( Ma_1 Ma_2 Ma_3 ma_4 Ma_5 ma_6 \)) and 80M (\( Ma_1 ma_2 Ma_3 Ma_4 Ma_5 ma_6 \)) [30,50]. The population was grown in a greenhouse under long day conditions and phenotyped for days to flowering. QTL for flowering time were identified on SBI02 and SBI10 (Fig 3). Recessive alleles of \( Ma_2 \) and \( Ma_4 \) result in earlier flowering when plants are grown in long days. The Hegari haplotype across the QTL on SBI10 was associated with early flowering therefore this QTL corresponds to \( Ma_4 \) (S2 Fig). The 80M haplotype across the QTL on SBI02 was associated with early flowering therefore the QTL on SBI02 was assigned to \( Ma_2 \).

Epistatic interactions between \( Ma_2 \) and \( Ma_4 \)

Previous studies indicated an epistatic interaction exists between \( Ma_2 \) and \( Ma_4 \) [28]. Therefore, Multiple QTL Mapping (MQM) analysis [51] was employed, using data from the Hegari/80M \( F_2 \) population, to identify additional flowering time QTL and interactions amongst the QTL as previously described [52]. MQM analysis identified the QTL for flowering time on SBI02 and
SBI10 and an additional QTL on SBI09. Additionally, an epistatic interaction was identified between \( \text{Ma}_2 \) and \( \text{Ma}_4 \) (pLOD = 42). Interaction plots showed that in a dominant \( \text{Ma}_4 \) background, a dominant allele at \( \text{Ma}_2 \) delays flowering, while in a recessive \( \text{Ma}_4 \) background, \( \text{Ma}_2 \) has a minimal impact on flowering time (Fig 4). The interaction between \( \text{Ma}_2 \) and \( \text{Ma}_4 \) identified by MQM analysis is consistent previous observations that in a recessive \( \text{ma}_4 \) background flowering is early regardless of allelic variation in \( \text{Ma}_2 \) [28].

**Ma2 candidate gene identification**

The Hegari/80M F2 population located \( \text{Ma}_2 \) on SBI02 between 67.3 Mbp to 69.1 Mbp (Fig 5). To further delimit the \( \text{Ma}_2 \) locus, six lines from the Hegari/80M population that were heterozygous across the \( \text{Ma}_4 \) QTL but fixed across the \( \text{Ma}_4 \) locus (\( \text{Ma}_4 \text{Ma}_4 \)) were selfed to create heterogeneous inbred families (HIFs) (n = 1000 F3 plants) [53]. Analysis of these HIFs narrowed the region encoding \( \text{Ma}_2 \) to \(~600\) kb (67.72 Mb-68.33 Mb) (Fig 5). Genotypes that were still heterozygous across the delimited locus were selfed and 100 F4 plants were evaluated for
differences in flowering time. This process narrowed the Ma$_2$ locus to a region spanning ~500 kb containing 76 genes (67.72Mb-68.22Mb) (Fig 5, S2 Table).

The low rate of recombination across the Ma$_2$ locus led us to utilize whole genome sequencing in conjunction with fine mapping to identify a candidate gene for Ma$_2$. Since 100M and 80M are near isogenic lines that have very few sequence differences along the long arm of SBI02 where the Ma$_2$ QTL is located, whole genome sequences (WGS) of 100M and 80M were generated in collaboration with JGI (sequences available at https://phytozome.jgi.doe.gov).

The genome sequences were scanned for polymorphisms within the 500 kb locus spanning Ma$_2$. Only one T → A single nucleotide polymorphism (SNP) located in Sobic.002G302700 was identified that distinguished 100M and 80M within the region spanning the Ma$_2$ locus. The T → A mutation causes a Lys141 change in the third exon, resulting a truncated protein. A 500 bp DNA sequence spanning the T to A polymorphism in Sobic.002G302700 was sequenced from 80M and 100M to confirm the SNP identified by comparison of the whole genome sequences (Table 1). The T → A point mutation was present in 80M (ma$_2$) whereas 100M (Ma$_2$) encoded a functional version of Sobic.002G302700 that encodes a full length protein. Since this mutation was the only sequence variant between 100M and 80M in the fine-mapped locus, Sobic.002G302700 was identified as the best candidate gene for Ma$_2$.

Sobic.002G302700 is annotated as a SET (Suppressor of variegation, Enhancer of Zeste, Trithorax) and MYND (Myeloid-Nervy-DEAF1) (SMYD) domain-containing protein. SMYD domain family proteins in humans have been found to methylate histone lysines and non-histone targets and have roles in regulating chromatin state, transcription, signal transduction, and cell cycling [54,55]. The SET domain in SMYD-containing proteins is composed of two sub-domains that are divided by the MYND zinc-finger domain. The SET domain includes conserved sequences involved in methyltransferase activity including nine cysteine residues that are present in the protein encoded by Sobic.002G303700 (Fig 6) [56]. The MYND domain

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**Fig 5. Fine-mapping of the Ma2 QTL.** The Ma$_2$ QTL spans from 67.3 Mb to 69.1 Mb (light blue bar). Five F$_2$ individuals that were heterozygous across the Ma$_2$ QTL were self-pollinated to generate heterogeneous inbred families (HIFs) totaling 1000 F$_3$ individuals. Genotype and phenotype analysis of these HIFs narrowed the QTL region to ~600 kb (darker blue bar). Two additional rounds of fine-mapping narrowed the QTL region to ~500 kb (vertical dashed lines). This region contained 76 genes. The genotypes of relevant HIFs and the parents are shown to the left and their corresponding days to flowering are shown to the right. Blue regions correspond to the 80M genotype and red regions correspond to the Hegari genotype. Purple regions are heterozygous.

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is involved in binding DNA and is enriched in cysteine and histidine residues [57]. Protein sequence alignment of Sobic.002G302700 homologs revealed that the SYMD protein candidate for Ma2 is highly conserved across flowering plants (Fig 6).

To learn more about Ma2 regulation, the expression of Sobic.002G302700 in 100M and 80M was characterized during a 48h L:D/L:L cycle. Ma2 showed a small increase in expression from morning to evening and somewhat higher expression in 100M compared to 80M during the evening (S3 Fig).

Distribution of Ma2 alleles in the sorghum germplasm

Recessive ma2 was originally found in the Milo background and used to construct Double Dwarf Yellow Milo (Ma1ma2ma3Ma4Ma5ma6) [28]. Double Dwarf Yellow Milo was crossed to Early White Milo (ma1Ma2ma3Ma4Ma5ma6) and the progeny selected to create 100M, 80M and the other Milo maturity standards [1,28,58]. Several of the Milo maturity standards were

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recorded as recessive Ma₂ (80M, 60M, SM80, SM60, 44M, 38M) and others as Ma₂ dominant (100M, 90M, SM100, SM90, 52M). In order to confirm the Ma₂ genotype of the maturity standards, the 500 bp sequence spanning the Lys141 mutation in Sobic.002G302700 was obtained from most of these genotypes (Table 1). Kalo was also identified as carrying a recessive allele of Ma₂. Kalo was derived from a cross of Dwarf Yellow Milo (ma₂), Pink Kafir (Ma₂), and CI432 (Ma₂), therefore it was concluded that DYM is the likely source of recessive ma₂ [28]. Sequence analysis showed that the genotypes previously identified as ma₂ including Kalo, 80M, SM80, 60M, 44M, and 38M carry the recessive mutation in Sobic.002G302700 identified in 80M. 100M, SM100, and Hegari that were identified as Ma₂, did not contain the mutated version of Sobic.002G302700 (Table 1). Additionally, sequences of Ma₂ from 52 sorghum genotypes with publicly available genome sequences were compared [40]. Sobic.002G302700 was predicted to encode functional proteins in all except one of these sorghum genotypes. A possible second recessive Ma₂ allele was found in IS3614-2 corresponding to an M83T missense mutation that was predicted to be deleterious by PROVEAN [59].

Discussion

In photoperiod sensitive sorghum genotypes, following the vegetative juvenile phase, day length has the greatest impact on flowering time under normal growing conditions. Molecular identification of the genes corresponding to Ma₁, Ma₃, Ma₅, and Ma₆ and other genes in the sorghum flowering time pathway (i.e., SbCO, SbEHD1, SbCN8/12) and an understanding of their regulation by photoperiod and the circadian clock led to the model of the flowering time pathway shown in Fig 7 [60]. The current study showed that Ma₂ represses flowering in long days by increasing the expression of SbPRR37 (Ma₁) and SbCO. The study also located QTL for Ma₂ and Ma₆, confirmed an epistatic interaction between Ma₂ and Ma₆, and identified a candidate gene for Ma₂.

In the current study, two near isogenic Milo maturity genotypes, 100M (Ma₂) and 80M (ma₂), were used to characterize how allelic variation in Ma₂ affects the expression of genes in the sorghum photoperiod regulated flowering time pathway. This analysis showed that mutation of Ma₂ significantly reduced the amplitude of the morning and evening peaks of SbPRR37 and SbCO expression without altering the timing of their expression. In parallel, the expression of SbCN12 (FT-like) increased 8-fold in leaves of 80M compared to 100M, consistent with
prior studies showing that 80M (ma₂) flowers earlier than 100M (Ma₂) in long days [28]. In contrast, expression of clock genes (TOC1, LHY) and other genes (i.e., GHD7, EHD1) in the photoperiod regulated flowering time pathway were modified to only a small extent by allelic variation in Ma₂. Based on these results, we tentatively place Ma₂ in the flowering time pathway downstream of the light sensing phytochromes and circadian clock and identify Ma₂ as a factor that enhances SbPBR37 and SbCO expression (Fig 7).

The differential increase in SbCN12 expression in 80M (vs. 100M) is consistent with inhibition of SbCN12 expression in long days by the concerted action of SbPRR37 and SbCO [27]. Genetic studies showed that floral repression mediated by SbPRR37 requires SbCO as a co-repressor [27]. Therefore, enhanced expression of both SbPBR37 (Ma₁) and SbCO by Ma₂ in Ma₁Ma₂ genotypes in long days is consistent with delayed flowering under these conditions relative to genotypes such as 80M that are Ma₁ma₂. Molecular genetic studies also showed that SbCO is an activator of SbCN12 expression and flowering in ma₁ genetic backgrounds [27]. This is consistent with the observation that ma₁Ma₂ genotypes flower earlier than ma₁ma₂ genotypes when grown in long days [28].

Interactions between Ma₂ and Ma₄

Multiple QTL (MQM) analysis of results from the population derived from Hegari/80M identified an interaction between Ma₂ and Ma₄ as well as one additional flowering QTL on SBI09. Flowering time QTL on SBI09 have been identified in other mapping populations, but the gene(s) involved have not been identified [33,34]. The interaction between Ma₂ and Ma₄ confirmed previous observations that recessive ma₄ causes accelerated flowering in long days in Ma₁Ma₂ genotypes [28]. Interestingly, the influence of Ma₂ and Ma₄ alleles on flowering time is affected by temperature [28,61]. The influence of temperature on flowering time pathway gene expression in 80M and 100M in the current study was minimized by growing plants at constant 30C. Further analysis of the temperature dependence of Ma₂ and Ma₄ on flowering time may help elucidate interactions between photoperiod and flowering time that have been previously documented [28,62]. Positional cloning of Ma₄ is underway to better understand the molecular basis of Ma₂ and Ma₄ interaction and their impact on flowering time.

Identification of a candidate gene for Ma₂

A mapping population derived from Hegari/80M that segregated for Ma₂ and Ma₄ enabled localization of the corresponding flowering time QTL in the sorghum genome (SBI02, Ma₂; SBI10, Ma₄). The Ma₂ QTL on SBI02 was fine-mapped using heterozygous inbred families (HIFs) from Hegari/80M. Identification of a candidate gene for Ma₂ was subsequently aided by comparison of genome sequences from the closely related 80M and 100M genotypes [28] A scan of the whole genome sequences of 100M and 80M identified only a single T to A mutation in the 500 kb region spanning the fine-mapped Ma₂ locus that caused a Lys141→ change in the third exon of Sobic.002G302700 resulting in protein truncation. Based on this information Sobic.002G302700 was tentatively identified as the best candidate gene for Ma₂.

Sobic.002G302700 encodes a SET (Suppressor of variegation, Enhancer of Zeste, Trithorax) and MYND (Myeloid-Nervy-DEAF1) (SMYD) domain containing protein. In humans, SMYD proteins act as lysine methyltransferases, and the SET domain is critical to this activity. Therefore, Ma₂ could be altering the expression of SbPBR37 and SbCO by modifying histones associated with these genes. The identification of this SMYD family protein’s involvement in flowering in sorghum as well as the identification of highly conserved homologs in other plant species suggests that Ma2 may correspond to a novel regulator of sorghum flowering. While a role for SYMD-proteins (lysine methyltransferases) as regulators of flowering time has not
been previously reported, genes encoding histone lysine demethylases (i.e., JMJ30/32) have been found to regulate temperature modulated flowering time in Arabidopsis [63].

J.R. Quinby [50] identified only one recessive allele of Ma2 among the sorghum genotypes used in the Texas sorghum breeding program. The maturity standard lines including 80M that are recessive for ma2 and the genotype Kalo were reported to be derived from the same recessive ma2 Milo genotype [28]. To confirm this, Ma2 alleles in the relevant maturity standards and Kalo were sequenced confirming that all of these ma2 genotypes carried the same mutation identified in 80M (Table 1). Among the 52 sorghum genotypes with available whole genome sequences, only 80M carried the mutation in Ma2 [40]. One possible additional allele of ma2 was identified in IS36214-2, which contained a M83T missense mutation that was predicted to be deleterious to protein function by PROVEAN [59].

In conclusion, we have shown that Ma2 represses flowering in long days by promoting the expression of the long day floral co-repressors SbPRR37 and SbCO (Fig 7). Sobic.002G302700 was identified as the best candidate for the sorghum Maturity locus Ma2. Further validation such as targeted mutation of Sobic.002G302700 in a Ma1Ma2 sorghum genotype or complementation of Ma1ma2 genotypes will be required to confirm this gene assignment. The identification of this gene and its interaction with Ma4 help elucidate an additional module of the photoperiod flowering regulation pathway in sorghum.

Supporting information

S1 Fig. Circadian expression of SbTOC1, SbLHY, SbGhd7, and SbEhd1. There were no consistent differences in expression of (A) SbTOC1, (B) SbLHY, (C) SbGhd7, and (D) SbEhd1 between 100M (solid black line) and 80M (dashed red line). (TIF)

S2 Fig. Genotype x phenotype plots for the QTL on SBI02 and SBI10. Recessive alleles of Maturity genes contribute to earlier flowering. 80M (AA) is recessive for ma2, while Hegari (BB) is dominant. Individuals genotyped AA for the QTL on SBI02 (represented by marker c2_68327634) flowered ~100 d earlier than those genotyped BB. 80M is dominant for Ma4, and individuals genotyped AA at the QTL on SBI10 (represented by marker c10_3607821) flowered ~100 d earlier than those genotyped BB. (TIF)

S3 Fig. Circadian expression of Sobic.002G302700 in 100M and 80M. The expression of Sobic.002G302700 does not cycle diurnally in 100M (solid black line) or 80M (dashed red line). There was no difference in expression between 100M and 80M in the first day. Expression was slightly elevated in 100M compared to 80M during the night and through the following morning. (TIF)

S1 Table. Ma2 (Sobic.002G302700) sequencing and qPCR primers. (DOCX)

S2 Table. Genes in the fine-mapped Ma2 QTL region. (XLSX)

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