

A Genomewide Analysis of the Cinnamyl Alcohol Dehydrogenase Family in Sorghum [*Sorghum bicolor* (L.) Moench] Identifies *SbCAD2* as the *Brown midrib6* Gene

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ABSTRACT

The content and composition of the plant cell wall polymer lignin affect plant fitness, carbon sequestration potential, and agro-industrial processing. These characteristics, are heavily influenced by the supply of hydroxycinnamyl alcohol precursors synthesized by the enzyme cinnamyl alcohol dehydrogenase (CAD). In angiosperms, CAD is encoded by a multigene family consisting of members thought to have distinct roles in different stages of plant development. Due to the high sequence similarity among CAD genes, it has been challenging to identify and study the role of the individual genes without a genome sequence. Analysis of the recently released sorghum genome revealed the existence of 14 CAD-like genes at seven genomic locations. Comparisons with maize and rice revealed subtle differences in gene number, arrangement, and expression patterns. Sorghum CAD2 is the predominant CAD involved in lignification based on the phylogenetic relationship with CADs from other species and genetic evidence showing that a set of three allelic *brown midrib* (*bmr*) lignin mutants contained mutations in this gene. The impact of the mutations on the structure of the protein was assessed using molecular modeling based on X-ray crystallography data of the closely related Arabidopsis CAD5. The modeling revealed unique changes in structure consistent with the observed phenotypes of the mutants.

THE biogenesis of the lignified plant cell wall is a complex metabolic process in which the formation and modification of several cell wall polymers need to be coordinated. Lignin is synthesized via the oxidative coupling of several phenolic monomers. The traditional view holds that lignin is derived from the monolignols *p*-coumaryl, coniferyl, and sinapyl alcohol, which give rise to *p*-hydroxyphenyl (H), guaiacyl (G), and syringyl (S) residues, respectively. Small amounts of other phenolic molecules can be incorporated as well, as a function of the species, genetic make-up, and tissue (reviewed by RALPH *et al.* 2004). The last step in monolignol biosynthesis is the reduction of cinnamyl aldehyde precursors (reviewed by HUMPHREYS and CHAPPLE 2002), catalyzed by the enzyme cinnamyl alcohol dehydrogenase (CAD; EC 1.1.1.95; MANSELL *et al.* 1974). Two enzymes with cinnamyl alcohol dehydrogenase activity were identified in *Eucalyptus gunnii*, and named EgCAD1 and EgCAD2 (GOFFNER *et al.* 1992). The enzymes are structurally unrelated and share no sequence similarity. EgCAD1 is

more similar to cinnamoyl-CoA reductase (EC 1.2.1.44), its active form is a monomer, and it is active toward a range of substrates, including coniferaldehyde and benzaldehyde derivatives, but not sinapaldehyde (GOFFNER *et al.* 1992, 1998; HAWKINS and BOUDET 1994). EgCAD2 is active as a homo- or heterodimer (HAWKINS and BOUDET 1994; BOUDET *et al.* 2004) and can synthesize all three monolignols. EgCAD2 was considered the predominant CAD involved in lignification (GOFFNER *et al.* 1992), and was the basis for studies on the role of CAD in lignification in other species. Recently, however, downregulation of the tobacco ortholog of *EgCAD1* was shown to have a subtle effect on the synthesis of coniferyl alcohol (DAMIANI *et al.* 2005). Unless stated differently, further mention of CAD refers to EgCAD2-type enzymes.

In gymnosperms, which contain lignin composed primarily of G residues, the CAD activity is strongly coniferaldehyde specific. Several CAD isoforms have been identified in gymnosperm species, although it is still unclear whether those isoforms represent alleles of a single locus or paralogs (GALLIANO *et al.* 1993; MACKAY *et al.* 1997; SCHUBERT *et al.* 1998). In angiosperm species CAD is encoded by a multigene family (DIXON *et al.* 2001; LI *et al.* 2001; LYNCH *et al.* 2002; SIBOUT *et al.* 2003; TOBIAS and CHOW 2005). The Arabidopsis and rice CAD gene families consist of 9 and 12 members, respectively

Sequence data from this article have been deposited with the EMBL/GenBank Data Libraries under accession nos. FJ554573 and FJ554574.

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(TAVARES *et al.* 2000; SIBOUT *et al.* 2003; TOBIAS and CHOW 2005). Angiosperm CAD proteins are multifunctional enzymes capable of catalyzing the reduction of *p*-coumaryl, coniferyl, and sinapyl aldehydes to their corresponding alcohols (HAWKINS and BOUDET 1994; KIM *et al.* 2004). The greater variation in lignin composition in angiosperms related to maturity and tissue type, together with the presence of multiple CAD genes, has led to the hypothesis that specialized CAD proteins with differential affinities for their aldehyde substrates play a role in regulating lignin composition during development (LI *et al.* 2001). The identification of a sinapyl alcohol dehydrogenase (SAD) in aspen (*Populus tremuloides*), which has greater affinity for sinapaldehyde than coniferaldehyde (LI *et al.* 2001), provides support for this hypothesis. However, no specific requirement for SAD activity for the synthesis of S units has been identified in Arabidopsis (KIM *et al.* 2004; SIBOUT *et al.* 2005).

Phylogenetic analyses of the Arabidopsis and rice CAD gene families reveal several subgroups (TAVARES *et al.* 2000; RAES *et al.* 2003; SIBOUT *et al.* 2003; TOBIAS and CHOW 2005), one of which displays strong similarity to the gymnosperm CAD genes. This group contains the genes that have been implicated in lignification of the vascular tissue by biochemical, expression, and mutant studies, including *AtCAD4* and *AtCAD5* (SIBOUT *et al.* 2003; KIM *et al.* 2004; SIBOUT *et al.* 2005), *ZmCAD2* (HALPIN *et al.* 1998; GUILLAUMIE *et al.* 2007a), and *OsCAD2* (TOBIAS and CHOW 2005; ZHANG *et al.* 2006). The enzymes encoded by the genes belonging to other groups have lower sequence similarity to these *bona fide* CADs and present wider substrate preferences and varying degrees of enzymatic activity and expression (for review see RAES *et al.* 2003; KIM *et al.* 2004; GUILLAUMIE *et al.* 2007a). It is likely that some of the genes are involved in defense response (KIEDROWSKI *et al.* 1992; BRILL *et al.* 1999) or metabolic processes not related to the lignification of the vascular tissue.

Although modifications in lignin content and composition have been observed in the vascular tissue of mutants and transgenics with reduced CAD activity (HALPIN *et al.* 1994, 1998; BAUCHER *et al.* 1996, 1999; CHABANNES *et al.* 2001; SIBOUT *et al.* 2005), it has been difficult to assign unique functions to individual CAD genes, underscoring the complementation capacity of the CAD multienzyme network. Without clear differences in substrate specificity, the spatiotemporal control of lignin biosynthesis in Arabidopsis and maize may therefore be the result of the regulated expression of tissue-specific CAD genes (RAES *et al.* 2003; GUILLAUMIE *et al.* 2007a).

The maize *brown midrib1* (*bm1*) mutant, which has reduced CAD activity (HALPIN *et al.* 1998), reportedly has lower expression of several CAD genes (GUILLAUMIE *et al.* 2007b) and several other monolignol biosynthetic genes. This led to the hypothesis that the *Bm1* gene may encode a transcription factor. Regulatory genes impli-

cated in the regulation of lignification were shown to be reduced in expression (GUILLAUMIE *et al.* 2007b). In contrast, a phenotypically similar mutant in rice, the *golden hull and internode2* (*gh2*) mutant, is caused by a mutation in the *OsCAD2* gene, the ortholog of the maize *ZmCAD2* gene identified by HALPIN *et al.* (1998). Genetic complementation experiments demonstrated that expression of the wild-type *OsCAD2* gene was sufficient to restore normal cell wall composition of *gh2* plants (ZHANG *et al.* 2006). The *brown midrib* (*bmr*) mutants of sorghum (PORTER *et al.* 1978) are similar to the *bm* mutants of maize. The abbreviation *bmr* was adopted to distinguish it from *bm*, already in use for the sorghum *bloomless* mutants (AYYANGAR and PONNAIYA 1941). There are at least four independent *bmr* loci (BITTINGER *et al.* 1981; SABALLOS *et al.* 2008). One of the mutants, *bmr6*, displays decreased CAD activity (BUCHOLTZ *et al.* 1980; PALMER *et al.* 2008) and contains cell walls with higher levels of cinnamaldehydes (PILLONEL *et al.* 1991; SABALLOS *et al.* 2008). Aided by the release of the sorghum genome sequence (www.phytozome.net/sorghum), we report here the systematic analysis of the sorghum CAD gene family, its relationship with the gene families from other species, and the identification of the *Bmr6* gene.

MATERIALS AND METHODS

Genome analysis: CAD-like genes were identified by performing a BLAST homology search (ALTSCHUL *et al.* 1990) of the sorghum genome sequence database (<http://www.phytozome.net/sorghum>) using as queries the DNA or deduced amino acid sequences of the rice *OsCAD2* (AK105011; TOBIAS and CHOW 2005), Arabidopsis *AtCAD4* and *AtCAD5* (At3g19450 and At4g34230, respectively; SIBOUT *et al.* 2003), and maize *ZmCAD2* (AJ005702.1; HALPIN *et al.* 1998; GUILLAUMIE *et al.* 2007b). The deduced amino acid sequences were used to query the translation of the sorghum genome in all six frames. The deduced amino acid sequence of the best DNA sequence match was used as a new query in both the sorghum and Arabidopsis (www.arabidopsis.org) genomes. This process was repeated until no additional CAD-like sequences were identified. Gene structure was predicted using Fgenesh+ (SALAMOV and SOLOVYEV 2000), and predicted intron/exon structures were compared with available EST sequences of rice and other grass species. Phylogenetic analysis included all rice and Arabidopsis CAD protein family members (KIM *et al.* 2004; TOBIAS and CHOW 2005). Two sorghum CAD genes (genomic accession nos. Sb02g024200.1 and Sb10g006310.1) were excluded from the analysis, because their sequences did not appear to encode functional proteins. Multiple sequence alignments were generated using CLUSTAL W version 3.2 (THOMPSON *et al.* 1994) with default parameters (<http://workbench.sdsc.edu>). A Bayesian phylogenetic analysis was performed using MrBayes software (<http://mrbayes.csit.fsu.edu/index.php>; HUELSENBECK and RONQUIST 2001) with the amino acid model set to pr = mixed, parameters lset rates = gamma, number of generations = 256,000 with sampling every 100 generations (RONQUIST *et al.* 2005). The output file was visualized for analysis using TreeView (PAGE 1996).

Plant materials and nucleic acid extraction: The *brown midrib* mutants *bmr6-ref*, *bmr6-3*, and *bmr6-27*, belonging to the

TABLE 1
Primer sequences used for *SbCAD* gene expression experiments

Gene	Genomic position		Sequence	
	Upper	Lower	Upper	Lower
<i>SbCAD2</i>	2931	3923	AAGATCTGGTCTACAACGA	GGACGACGAGCTGATCAC
<i>SbCAD4-2</i>	1257	2104	CACCTCTTCTGTGTGCTGGGATAAC	GAAATGGATCAAACGGATGGTC
<i>SbCAD4-3</i>	984	3039	GTTGTCTGGACCGGAATATGCACC	GCCGCACAGAAGTTAACCAT
<i>SbCAD5</i>	51	2574	CCCAGCGTCTCGGATCGCTT	GTCACCACTCCGGCGATCTC
<i>SbCAD6</i>	11	517	CACCAAACCACACGCAGAC	TCACCTTGTTCGCAGTAGTTCT
<i>SbCAD7</i>	148	2693	CGGGCAACGATTATGG	GTGGCATATCCCGCAGTACAG
<i>SbCAD8-1</i>	255	1019	CGGGCTTCGGSTGGG	ACGCCGGTCCACCACG
<i>Ubiquitin</i>	221	351	GGTTCGGGAGGTGGCATAGGT	AGCATGTACATTCCCAGCGGTAG

All primers are listed in 5'–3' direction.

bmr6 group (SABALLOS *et al.* 2008) and their normal isolines, were planted in 2004 and 2008 at the Purdue Agronomy Center for Research and Education (ACRE, West Lafayette, IN). DNA was extracted from leaf tissue by a modified CTAB protocol (BERNATZKY and TANKSLEY 1986). Three *bmr6-ref* near-isogenic lines (NILs) in the inbred lines BTx623, RTx430, and Kansas Collier (PEDERSEN *et al.* 2006a,b) were planted in the greenhouse at the University of Florida (Gainesville, FL). Young leaf tissue was harvested and extracted using MoBio Power Plant DNA isolation kit (Carlsbad, CA).

A population of recombinant inbred lines (RILs) were created by crossing Brown County, a non-*bmr* sweet sorghum cultivar, with the *bmr6-ref* line. The F₁ generation was selfed to produce an F₂ population of 218 plants. Individual selfed F₂ plants were advanced to the F₆ generation using the single seed descent method. The phenotype (*bmr* vs. wild type) was recorded, and DNA of the individual F₆ RILs was extracted from leaf tissue using a modified CTAB method (BERNATZKY and TANKSLEY 1986).

For expression assays, BTx623 and BTx623-*bmr6-ref* lines were planted in the greenhouse at the University of Florida. Tissue was collected at developmental stage 2 (VANDERLIP 1993). At this stage, the stalk is small and covered by the leaf sheaths; therefore, leaves and stacked internodes were harvested together to represent the shoot. In addition, the complete root ball was harvested. Tissue was also collected at stage 4 (growing point differentiation) from the apex of developing roots, the first internode (counting from the top of the stem), and the whorl. For both stages, each line was harvested in triplicate and assayed separately. The tissue was frozen in liquid nitrogen and stored at –80° until RNA extraction with the RNeasy Plant RNA extraction kit with in-column DNase digestion (QIAGEN, Carlsbad, CA). First-strand cDNA was synthesized with the iScript cDNA synthesis kit (Bio-Rad, Hercules, CA). Tissue from the *bmr6-3* mutant and the normal isolate N3, planted in the greenhouse (West Lafayette, IN), was harvested at stage 2 and processed as described above.

PCR and cloning: A homolog to the maize *ZmCAD2* gene was identified in the sorghum genome (accession no. Sb04-g005950.1) and used to design primers for cloning and gene expression experiments. The identified genomic sequence corresponded to the cDNA sequence identified by TSURUTA *et al.* (2007) (GenBank accession no. AB288109). The *SbCAD2* gene was cloned from *bmr6* lines and their respective wild-type isolines. Primer pair U168 (5' CGG CTC CGT CGT GTT C 3') and L3923 (Table 1) was used to amplify a 710-bp fragment from the 5' end of the gene using cDNA as template, with an annealing temperature of 62°. cDNA was used as template due to the technical difficulties of amplifying and

sequencing through the first intron of the gene. A 2516-bp 3' fragment (with partial overlap to the 5' fragment) was amplified from genomic DNA using primers U2237 (5' TTT CAG CAC CTT TGG GAT CGA GT 3') and L4753 (5' GCA ACA TTT GGG CAT ACG TCT 3') as above, except that the annealing temperature used was 61°.

PCR was performed with Jump Start RedAccuTaq (Sigma, St. Louis) according to the manufacturer's instructions, using a three-step program. The products were analyzed by gel electrophoresis, purified from the gel using the MinElute gel purification kit (QIAGEN), cloned in vector pT7-Blue3 (EMD-Biosciences, Gibbstown, NJ), and introduced in *Escherichia coli* NovaBlue cells (EMD-Biosciences). Plasmids with inserts were sequenced in both directions at the University of Florida's Interdisciplinary Center for Biotechnology Research on an Applied Biosystems 3700 capillary electrophoresis sequencer. The sequence from a minimum of three plasmids for each insert was compared to eliminate discrepancies in the sequences due to PCR and sequencing errors. Sequence analysis was performed using Biology Workbench 3.2 software.

CAPS analysis: CAPS markers (KONIECZNY and AUSUBEL 1993) were based on amplification of the *SbCAD2* gene with primers U2931 and L3923 (Table 1), and JumpStart AccuTaq ReadyMix PCR reaction mixes (Sigma), using a three-step PCR program with annealing temperature of 59°. The PCR products were purified using AccuPrep PCR purification kit (Bioneer, Seoul, South Korea) and digested with restriction endonucleases according to manufacturer's directions. The enzymes *TauI* and *TauI* (Fermentas, Glen Burnie, MD) were used for the identification of the *bmr6-ref* and *bmr6-3* alleles, respectively. The digested DNA was separated on a 2% agarose gel.

Structural modeling of *SbCAD2* and its mutant versions: The crystal structure of the AtCAD5 (PDB ID: 2CF6, YOUNG *et al.* 2006) was used to build a model for *SbCAD2* and the *bmr6-3* and *bmr6-27* mutant versions. The substitutions, insertions, and deletions were superimposed on the coordinates of AtCAD5, followed by a quick energy minimization by CNS v1.1 (BRÜNGER *et al.* 1998) using potential function parameters of CHARMM19 as described previously (KIM *et al.* 2004). Substrate position was generated through the solid docking module on QUANTA (BioSYM/Micron Separations), which is based on conformational space, followed by a quick energy minimization by CNS v1.1.

Expression studies of the *CAD*-like family of genes in normal, *bmr6-ref* and *bmr6-3* mutants of sorghum: Gene-specific primer pairs flanking introns were designed to differentiate amplification resulting from cDNA and residual genomic DNA (Table 1). cDNA from shoots and roots of three plants each of BTx623/BTx623 *bmr6-ref* NILs at stages 2 and 4, and from roots and shoots of N3/*bmr6-3* NILs at stage 2 was tested for amplification of seven different sorghum *CAD* genes.

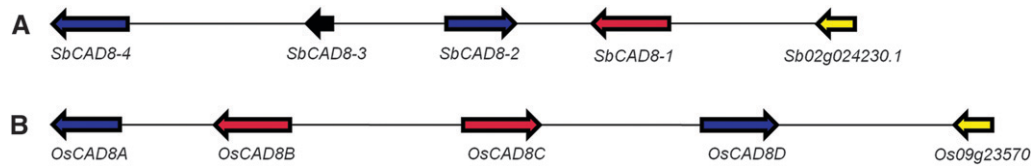


FIGURE 1.—Organization of the sorghum (A) and rice (B) *CAD8* gene clusters. Block arrows represent the genes and their orientation. Arrows with the same color represent genes more similar to each other than

to the other members of the cluster. The yellow arrows represent a gene downstream of the *CAD8* cluster present in both the rice and sorghum genomes. The scale is approximate and based on the sorghum and rice physical map. The represented sorghum and rice fragments span ~21.5 and 26 kb, respectively.

Amplification of the *Ubiquitin* cDNA was used to control for cDNA loading. The *Ubiquitin* primers were based on the maize cDNA sequence (GenBank accession no. BT016881). The reactions were carried out using RedTaq Readymix PCR reaction mix (Sigma) in a three-step PCR program consisting of 28 cycles. This was empirically determined to be within the exponential phase of amplification. PCR products were visualized in a 2% agarose gel.

RESULTS

The *CAD* family in sorghum: The availability of the sorghum genome sequence has made it relatively straightforward to identify the different members of the *CAD* gene family. An iterative search of the published sorghum genome sequence using as query the maize, rice, and *Arabidopsis bona fide* *CAD* sequences resulted in the identification of 14 *CAD-like* genes at seven different loci in the sorghum genome. Two of the loci (*SbCAD4* and *SbCAD8*) contain clusters of genes (*SbCAD4-1* to *-5*, and *SbCAD8-1* to *-4*). The DNA sequence of two members of these groups, *SbCAD4-1* and *SbCAD8-3*, are interspersed with retroelements, appear to not encode full-length proteins, and are likely pseudogenes. Sorghum genes in cluster *SbCAD8* are very similar to the genes in the rice cluster *OsCAD8*, but they differ in the gene arrangement.

The comparison of the sorghum and rice *CAD8* gene clusters is presented in Figure 1. The chromosomal location and EST information of the entire *CAD* family is presented in Table 2.

Phylogenetic analysis: To understand the structure of the *CAD* gene family and the possible functional divergence of the individual members, a phylogenetic analysis was performed using deduced amino acid sequences. The nomenclature of sorghum *CAD* genes was based on similarity with the corresponding rice orthologs (TOBIAS and CHOW 2005). In general, good correspondence between the rice and sorghum family structure was observed, with the exception of *OsCAD1*, *OsCAD3*, and *OsCAD9*, for which no sorghum orthologs were found, and *SbCAD10*, for which no unambiguous rice ortholog was identified. The predicted length of the proteins varied between 342 and 383 residues and had a maximum and a minimum identity to maize *CAD2* of 95% (*SbCAD2*) and 40% (*SbCAD4-2*), respectively.

All of the sorghum *CADs* contain the conserved GHE(X)₂G(X)₅G(X)₂V motif involved in the binding of the catalytic Zn²⁺, the GD(X)₁₀C(X)₂C(X)₂(X)₇C structural Zn²⁺ binding domain, and the conserved glycine-rich repeat G(X)GGV(L)G involved in NADP(H) cofactor binding. On the basis of these motifs, these

TABLE 2

Genomic location of the sorghum *CAD* genes and expression data based on ESTs reported in the sorghum genome database

Gene	Accession no. in genome	No. of reported ESTs	Position in genome
<i>SbCAD2</i>	Sb04g005950.1	21	4:5781208..5776540
<i>SbCAD4-1</i>	Sb10g006310.1	0	10:5750788..5749847
<i>SbCAD4-2</i>	Sb10g006300.1	0	10:5745022..5747508
<i>SbCAD4-3</i>	Sb10g006290.1	5	10:5737327..5740568
<i>SbCAD4-4</i>	Sb10g006280.1	3	10:5730469..5733611
<i>SbCAD4-5</i>	Sb10g006270.1	1	10:5711950..5714251
<i>SbCAD5</i>	Sb07g006090.1	12	7:8643832..8640087
<i>SbCAD6</i>	Sb06g001430.1	2	6:2167025..2168295
<i>SbCAD7</i>	Sb06g028240.1	5	6:57074661..57079263
<i>SbCAD8-1</i>	Sb02g024220.1	19	2:58410576..58408367
<i>SbCAD8-2</i>	Sb02g024210.1	13	2:58404166..58406013
<i>SbCAD8-3</i>	Sb02g024200.1	0	2:58401666..58401105
<i>SbCAD8-4</i>	Sb02g024190.1	9	2:58396113..58394105
<i>SbCAD10</i>	Sb08g016410.1	0	8:43662072..43660817

proteins appear to be zinc-dependent alcohol dehydrogenases and members of the medium-chain dehydrogenase/reductase (MDR) superfamily (PERSSON *et al.* 1994; RIVEROS-ROSAS *et al.* 2003). YOUN *et al.* (2006) identified 12 amino acids that constitute the proposed substrate binding site of the *bona fide* CADs in a number of angiosperms and gymnosperms. Of the 14 identified CAD-like proteins, SbCAD2 contains 10 of these 12 conserved amino acids (Figure 2). All other family members contained a varying number of amino acid substitutions in these residues, as is the case for other CAD family members in Arabidopsis, poplar, and rice (LI *et al.* 2001; RAES *et al.* 2003; BOMATI and NOEL 2005; TOBIAS and CHOW 2005).

An unrooted phylogenetic tree (Figure 3) constructed from the deduced amino acid sequences shows the relationship between the sorghum CAD family members and previously reported CADs from other species. Sorghum sequence Sb07g025220.1, the closest sequence outside the CAD family, was included in the analysis for comparison. The CAD proteins included in this analysis cluster in five groups.

Group I corresponds to class I of RAES *et al.* (2003), and contains the sequences identified as the *bona fide* CAD enzymes involved in the reduction of coniferyl and sinapyl aldehydes to their corresponding alcohols for the formation of lignin in lycophytes, gymnosperms, and monocot and dicot angiosperms. SbCAD2 belongs to this group and it is closely related to the maize ZmCAD2 and rice OsCAD2 proteins. Group II contains AtCAD2, AtCAD3, AtCAD9 [following the nomenclature of KIM *et al.* (2004)], designated by RAES *et al.* (2003) as class III. Group III encompasses enzymes of diverse substrate specificity, including the proposed poplar SAD (LI *et al.* 2001) and Arabidopsis enzymes belonging to class II of RAES *et al.* (2003), involved in pathogen defense (SOMSSICH *et al.* 1996). Group IV contains the SbCAD6 and OsCAD6 sequences (TOBIAS and CHOW 2005). Both the rice and the sorghum sequences contain the SKL peroxisome targeting sequence at their C terminus. Group V contains monocot and dicot CADs. Even though there appear to be groups specific to monocots (group IV) and dicots (group II), sequence resources are probably still too limited for definitive conclusions.

The *bmr6* mutation affects the *SbCAD2* gene: The increased levels of cinnamaldehyde endgroups in the lignin of the *bmr6-ref* mutant are consistent with reduced CAD activity (BUCHOLTZ *et al.* 1980; PILLONEL *et al.* 1991; PALMER *et al.* 2008; SABALLOS *et al.* 2008). On the basis of the clustering of SbCAD2 in group I with the *bona fide* CADs, we hypothesized that the *SbCAD2* gene is the main CAD gene in sorghum and that a mutation in this gene would affect lignification.

The *SbCAD2* alleles of three different inbred lines in which the *bmr6-ref* allele had been introgressed revealed a C-to-T transition at position 2800 of the *SbCAD2*

genomic sequence. This particular nucleotide change creates a premature stop codon. The resulting predicted protein contains only 131 amino acids and lacks the nucleotide binding site and most of the residues that form the substrate binding site. Such a truncated protein is likely to be nonfunctional even if produced. The mutation creates a novel restriction site for endonuclease *TauI* (ACGC → ACGT). Restriction digest of a fragment of the gene (nucleotides 2931–3923 of the genomic DNA) resulted in three fragments (585, 407, and 24 bp) from the wild-type alleles, whereas the *bmr6-ref* allele produced four fragments (508, 363, 44, and 24 bp). This CAPS marker is therefore diagnostic for the presence of the *bmr6-ref* allele. The mutation was confirmed with the CAPS marker using independently isolated DNA from N6 and *bmr6-ref* (Figure 4). Furthermore, this CAPS marker was used in a cosegregation analysis of an F₆ RIL population derived from a cross between Brown County × *bmr6*, and indicated complete linkage between the mutated *SbCAD2* gene (*bmr6-ref* allele) and the *bmr* phenotype.

In the absence of a facile transformation protocol for sorghum, proving beyond reasonable doubt that the *SbCAD2* gene is the *Bmr6* gene requires the identification of additional independent mutations in this gene in mutants that are allelic to *bmr6-ref*. Indeed, sequencing of the *bmr6-3* allele revealed a G-to-A transition at position 3875 of the genomic sequence. This mutation results in an amino acid substitution from Gly (G) to Ser (S) at position 191 of the protein sequence. The glycine-rich motif ¹⁸⁸G(X)GGV(L)G¹⁹³ is highly conserved among studied MDR proteins. The motif is located at the first β-α-β-unit of the cofactor binding site and is presumed to participate in the binding of the pyrophosphate group of the NADP⁺-cofactor (YOUN *et al.* 2006). Although the mutated allele would encode a full-length protein, its catalytic activity may be compromised. A rice mutant with a single mutation in this same region, 185 G → D, results in complete loss of enzymatic activity, and is the causal mutation of the *gh2* phenotype (ZHANG *et al.* 2006).

The base substitution in the *bmr6-3* allele eliminates a restriction site for the enzyme *TauI*. A restriction digest of the amplified gene fragment 2931–3923 with *TauI* resulted in four fragments in the *bmr6-3* allele (796, 138, 54, and 5 bp) with a diagnostic fragment of 138 bp. This product was cleaved into two fragments (92 and 46 bp) in the wild-type allele. The mutation was confirmed with this marker using independently isolated DNA from N3 and *bmr6-3* (Figure 4).

The *bmr6-27* mutation is a nucleotide insertion after position 4288 (+G), which causes a frameshift that results in the formation of a protein with nonconservative substitutions in the 31 C-terminal residues. The mutation also created a premature stop codon that renders the protein four residues shorter than the wild-type protein. In addition to the mutation responsible for the

		10	20	30	40	50	60	70	80						
SbCAD8-2	1			MEVSKAAL	GWAARDAS	GHLSPYSFS	.RVL	RDC	DVT	TKV	LCG	VCH	TDL	LH	LKNE
SbCAD8-4	1			MEQGGKTAH	GWAARDAS	GHLSPYSFS	.R	QGD	ADV	TKV	LCG	VCH	TDL	LH	LKNE
SbCAD8-1	1			MEEKAGQAAF	GWAARDAT	GVLSPYNSFR	.R	VPK	DDV	TKV	LCG	VCH	TDL	LH	LKNE
SbCAD4-4	1			MAGE.....	KPQNGCNAA	WAARDPSG	VLS	YFN	FR	.RAV	QSS	DVA	KI	TY	CG
SbCAD4-3	1			MAGE.....	KPQHGNCNA	WAARDPSG	VLS	YFN	FR	.RP	VQSS	DVA	KI	TY	CG
SbCAD4-5	1														
SbCAD4-2	1			MAAE.....	SENSNCNA	WAARDPSG	VLS	YFN	FR	.NKA	VQSS	DVA	KI	TY	CG
SbCAD10	1														
SbCAD6	1			MEVT.....	PNHTQTVA	GWAAMNES	GKLE	PF	IF	KR	.R	ENG	VDD	VTT	KV
SbCAD7	1			MAP.VEA.....	QHPRTALAL	AHADAS	CRV	SP	IR	IS	.R	TG	DD	VA	IQ
SbCAD5	1			CSSSQISDIP	MAPVAAE	AAAKGQL	QHTG	KAV	G	L	A	H	A	D	A
SbCAD2	1														
ZmCAD2	1														
AICAD5	1														

		90	100	110	120	130	140	150	160
SbCAD8-2	55	WGNALY	VVP	GHEV	VVGV	VDV	GAG	VT	FK
SbCAD8-4	56	WGNALY	VVP	GHEV	VVGV	VDV	GAG	VT	FK
SbCAD8-1	57	WGNALY	VVP	GHEV	VVGV	VDV	GAG	VT	FK
SbCAD4-4	59	QHDSKY	PLV	PGHE	TAG	VVT	VG	AD	WK
SbCAD4-3	59	HHDSKY	PLV	PGHE	TAG	VVT	VG	AD	WK
SbCAD4-5	29	LNDSKY	PLV	PGHE	TAG	VVT	VG	AD	WK
SbCAD4-2	59	LNDSKY	PLV	PGHE	TAG	VVT	VG	AD	WK
SbCAD10	15	YTL	V	P	G	H	E	V
SbCAD6	59	WGITMY	PLV	PGHE	IT	GV	VT	V	G
SbCAD7	61	WGNALY	VVP	GHE	EAG	L	T	T	E
SbCAD5	80	WGNALY	VVP	GHE	TAG	V	V	T	E
SbCAD2	58	LGASKY	PMV	PGHE	V	V	G	V	V
ZmCAD2	58	LGASKY	PMV	PGHE	V	V	G	V	V
AICAD5	58	LGMSNY	PMV	PGHE	V	V	G	V	V

		170	180	190	200	210	220	230	240
SbCAD8-2	134	DGGFSD	V	V	VD	Q	H	V	V
SbCAD8-4	132	KGGFSD	V	V	V	S	Q	D	F
SbCAD8-1	133	KGGFSD	V	V	V	N	H	E	V
SbCAD4-4	133	KGGYSTH	I	V	V	H	E	R	Y
SbCAD4-3	133	KGGYSTH	I	V	V	H	E	R	Y
SbCAD4-5	103	KGGYSTH	I	V	V	H	E	R	Y
SbCAD4-2	133	MGGYSTH	I	V	V	H	E	R	Y
SbCAD10	84	MGGYSTH	I	V	V	H	E	R	Y
SbCAD6	133	YGGSY	M	L	V	A	H	K	R
SbCAD7	135	YGGSY	M	L	V	A	H	K	R
SbCAD5	155	YGGSY	M	L	V	A	H	K	R
SbCAD2	132	QGGFAST	M	V	D	Q	K	F	V
ZmCAD2	132	QGGFAST	M	V	D	Q	K	F	V
AICAD5	132	QGGFAST	M	V	D	Q	K	F	V

		250	260	270	280	290	300	310	320
SbCAD8-2	211	V	I	S	S	P	A	K	R
SbCAD8-4	209	V	I	S	S	P	A	K	R
SbCAD8-1	210	V	I	S	S	P	A	K	R
SbCAD4-4	210	V	I	S	S	P	A	K	R
SbCAD4-3	210	V	I	S	S	P	A	K	R
SbCAD4-5	180	V	I	S	S	P	A	K	R
SbCAD4-2	210	V	I	S	S	P	A	K	R
SbCAD10	162	V	I	S	S	P	A	K	R
SbCAD6	211	V	I	S	S	P	A	K	R
SbCAD7	212	V	I	S	S	P	A	K	R
SbCAD5	233	V	I	S	S	P	A	K	R
SbCAD2	209	V	I	S	S	P	A	K	R
ZmCAD2	209	V	I	S	S	P	A	K	R
AICAD5	209	V	I	S	S	P	A	K	R

		330	340	350	360	370	380	390	400
SbCAD8-2	291	I	T	G	G	R	V	A	G
SbCAD8-4	289	I	T	G	G	R	V	A	G
SbCAD8-1	290	V	P	G	K	G	V	A	G
SbCAD4-4	280	..	A	R	L	S	G	S	L
SbCAD4-3	280	..	A	R	L	S	G	S	L
SbCAD4-5	250	..	A	R	L	S	G	S	L
SbCAD4-2	280	..	A	R	L	S	G	S	L
SbCAD10	241	I	F	C	A	R	L	S	G
SbCAD6	291	I	F	C	A	R	L	S	G
SbCAD7	292	V	T	G	K	T	L	A	G
SbCAD5	313	I	M	G	N	K	L	A	G
SbCAD2	289	M	L	G	R	K	A	I	T
ZmCAD2	289	M	L	G	R	K	A	I	T
AICAD5	289	M	L	G	R	K	A	I	T

FIGURE 2.—Sequence comparison of the translated sequences of *Sorghum bicolor* CAD family members, maize ZmCAD2 (*Zea mays* sequence AJ005702.1), and Arabidopsis AtCAD5 (*Arabidopsis thaliana* sequence At4g34230). The conserved amino acids identified by YOUN *et al.* (2006) as part of the substrate binding pocket are indicated by asterisks above the sequences. The conserved glycine-rich motif G(X)GGV(L)G is indicated by a dotted line below the sequences. Italicized letters indicate completely conserved residues across all 14 sequences.

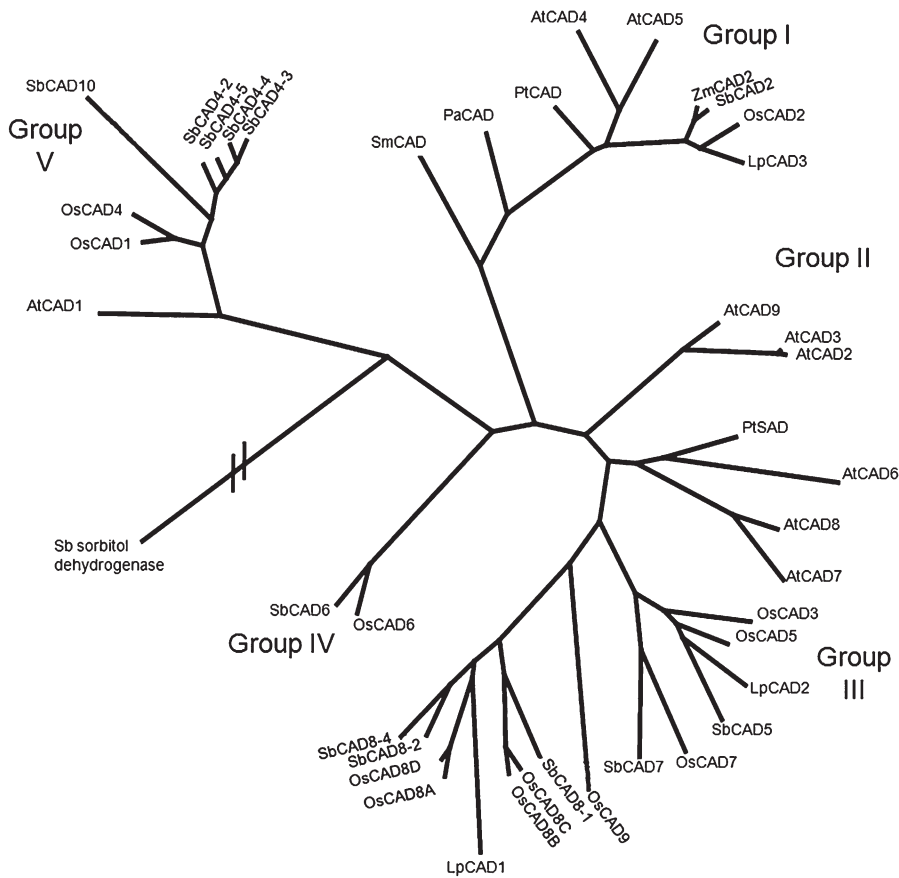


FIGURE 3.—Phylogenetic relationship between the sorghum CAD family and CAD proteins from other species. The branch lengths indicate the relative distance between sequences, with the exception of the putative sorghum sorbitol dehydrogenase, which is distantly related to the CAD family (accession no. Sb07g025220.1). The sequences used to build the tree were: *Selaginella moellendorffii* SmCAD1-1 (protein ID 450958 of the JGI Selaginella genome database); *Picea abies* PtCAD (O82035); *Zea mays* ZmCAD2 (CAA06687); *Arabidopsis thaliana* AtCAD1 (At1g72680), AtCAD2 (At2g21730), AtCAD3 (At2g21890), AtCAD4 (At3g19450), AtCAD5 (At4g34230), AtCAD6 (At4g37970), AtCAD7 (At4g37980), AtCAD8 (At4g37990), and AtCAD9 (At4g39330); *Populus tremuloides* PtSAD (AAK58693); *PtCAD* (AAF43140); *Oryza sativa* OsCAD1 (NP_001064283.1), OsCAD2 (NP_001046132.1), OsCAD3 (AAP53892), and OsCAD4 (NP_001068303.1). OsCAD5 (BK003971), OsCAD6 (CAD39907), OsCAD8 A-C (BK003972), and OsCAD9 (AAN05338). Sorghum sequences are those described in Table 2. The rice CAD gene nomenclature follows TOBIAS and CHOW (2005).

phenotype, the *bmr6-27* allele also contains a deletion of base 3077 (–T), but this is a silent mutation because of its location in the third intron of the gene. This mutation does not cause splicing abnormalities, as

sequencing of the *CAD2* cDNA from wild type and *bmr6-27* showed no sequence variation in this region.

Structural comparison of SbCAD2 and its mutant versions with AtCAD5: The amino acid sequence comparison between SbCAD2 and AtCAD5 (Figure 2) revealed 93 residue changes in total. Among those, 48 are conservative changes and 45 are nonconservative changes. A close inspection of the 3D structure of AtCAD5 (PDB ID: 2CF6) revealed that 52 of these changes are surface residues, whereas 39 occur in the secondary structure. In general, the 3D model of SbCAD2 remains very similar to that of AtCAD5, with many of the changes not having a significant impact on the overall structure (Figure 5). All the secondary structural elements observed in AtCAD5 are maintained in SbCAD2. The functional unit of AtCAD5 is a dimer tightly associated through two two-fold-related β -strands. SbCAD2 very likely maintains the dimer character on the basis of the intact nature of β D, β E, and β F. There is a minor perturbation caused by L271H in β E, which is due to a delocalization of the imidazole ring of residue 271 by the neighboring D250 and H256 residues, but the magnitude of the impact is not enough to disrupt the dimer formation at the β F contact points.

Two of the amino acid changes between AtCAD5 and SbCAD2 are in positions that may alter the enzymatic activity (Figure 6). The first of those is residue 60 M \rightarrow A. This residue resides near the surface of SbCAD2 and

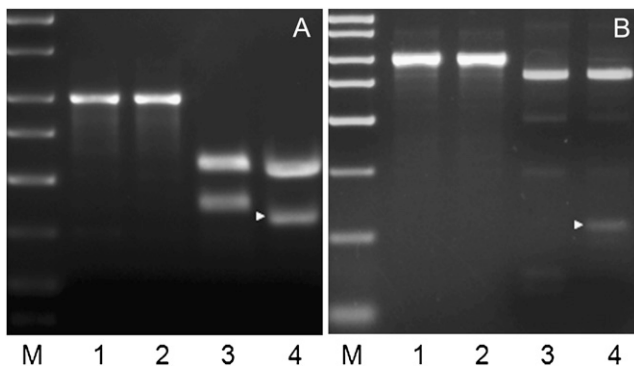


FIGURE 4.—Molecular markers can distinguish between the wild-type and mutant *SbCAD2* alleles. (A) CAPS marker specific for the *bmr6-ref* allele. Lanes 1 and 2 show the undigested 992-bp PCR product from N6 and *bmr6-ref* DNA, respectively. Lanes 3 and 4 show the respective PCR products after digestion with the restriction enzyme *TauI*. The 363-bp diagnostic fragment is indicated with a solid triangle. (B) CAPS marker specific for the *bmr6-3* allele. Lanes 1 and 2 show the undigested PCR product from N3 and *bmr6-3* DNA, respectively. Lanes 3 and 4 show the respective PCR products after digestion with the restriction enzyme *TauI*. The 138-bp diagnostic fragment is indicated with a solid triangle. The size marker is Sigma PCR Marker [catalog (cat.) no. P9577].

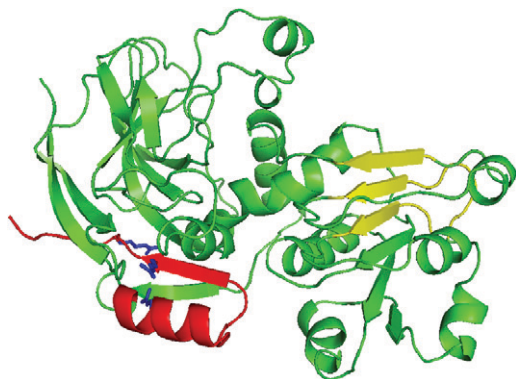


FIGURE 5.—Ribbon diagram representation for the structure of SbCAD2. The significant arginine mutations found in SbCAD2_27b are indicated in blue. The dimer contact positions for β D–F are highlighted in yellow. The three secondary structural elements, β 11, β 12, and α 6 are not maintained in the CAD model of *bmr6-27* mutant due to the degenerative hydrophobic interactions caused by the arginine residues highlighted in blue.

may possibly affect the entry of substrate. The methionine residue can act as a hydrophobic gating mechanism in concert with surrounding side chains. Thus, by replacing it with alanine, the specificity of this potential gate becomes less stringent. The second change, residue 95 C \rightarrow V, occurs in close proximity of the substrate-binding pocket. The amino acid change generates a more hydrophobic character to this pocket, which is

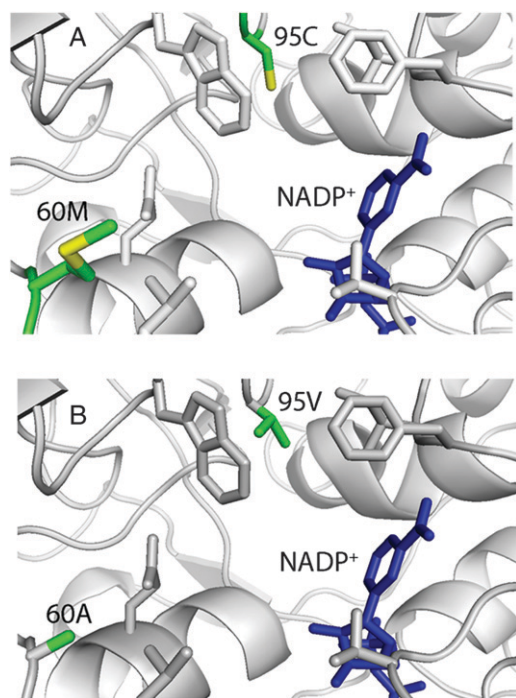


FIGURE 6.—Graphical comparison of active site residues interacting with the NADP⁺ cofactor of (A) AtCAD5 and (B) SbCAD2. Residue M60 and C95 change to A and V, respectively, whereas the other residues are conserved. This will potentially affect substrate binding and specificity.

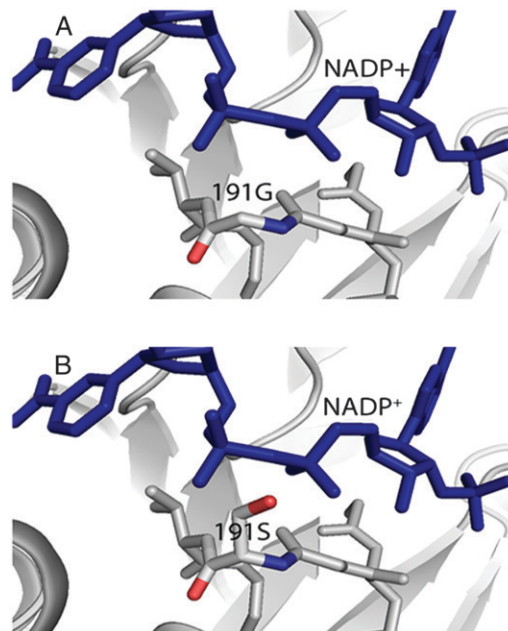


FIGURE 7.—Effect of the *bmr6-3* mutation on the structure of SbCAD2. (A) G(X)GGV(L)G motif and its interaction with the phosphate backbone of NADP⁺. (B) SbCAD2 *bmr6-3* variant at residue 191. The G191S mutation will affect the interaction between this flexible motif and phosphate groups.

filled with water molecules in the apo-form AtCAD5. Consequently, there might be fewer water molecules in the substrate-binding pocket of SbCAD2, resulting in a significant reduction of the entropic contribution to the free energy associated with substrate binding. Therefore, it is very likely that SbCAD2 has not only reduced efficiency of substrate binding, but also altered substrate specificity compared to AtCAD5.

SbCAD2 in the *bmr6-3* and *bmr6-27* mutants: The only amino acid change observed in the *bmr6-3* mutant is 191 G \rightarrow S, which is part of the highly conserved ¹⁸⁸GXGGV(L)G¹⁹³ motif. A serine residue at this site is a very rare alteration. Among the Zn-dependent medium-chain dehydrogenases in the database, this site is highly conserved with only hydrophobic residues. The unique serine substitution at position 191 in the *bmr6-3* mutant will change the characteristics of this motif and consequently affect the binding affinity for the NADP⁺ cofactor (Figure 7).

In the *bmr6-27* mutant, there is a total of 27 amino acid changes, all within the C terminus, spanning residues 331–361. Since the mutation affects a relatively small portion of the total sequence, and does not directly affect the active site of the enzyme or its dimerization domain, it is important to determine whether the changes produced structural alterations that would affect its activity. Among those 27 mutations, only 24 can be modeled in because the remaining C-terminal three residues are disordered in the AtCAD5 structure. Those remaining mutations are distributed through the secondary struc-

tures $\beta 12$ (346–349) and $\alpha 6$ (330–338). In AtCAD5, those secondary structural elements, together with $\beta 11$ (323–326) in the C terminus, maintain their interface through hydrophobic interactions. Those interfaces among $\beta 11$ and $\beta 12$ and $\alpha 6$ appear to be significantly affected by several arginine substitutions, which include 333 A \rightarrow R, 348 V \rightarrow R, and 350 V \rightarrow R. Specifically, 333 A \rightarrow R disrupts the hydrophobic area that maintains $\beta 11$, 348 V \rightarrow R disrupts the hydrophobic integrity of $\alpha 6$, while 350 V \rightarrow R perturbs a hydrophobic pocket generated by V40, P67, G134, and F135. In addition, the 346 F \rightarrow P substitution disrupts the pocket, which in turn affects the integrity of $\beta 12$.

The secondary structural prediction algorithms show that both $\alpha 6$ (330–338) and $\beta 12$ (346–349) are no longer maintained. However, the amino acid changes are unlikely to affect the ability to dimerize due to their distal locations from βD , βE , and βF (Figure 5). Another significant substitution is 342 V \rightarrow R, which approaches up to 5 Å toward the S γ -atom of 47C, and could thus alter the coordination of the catalytic Zn²⁺. The various amino acid substitutions present in *bmr6-27* could result in either serious disruption of the catalytic domain or expansion of the intersecondary distances among $\beta 11$, $\beta 12$, and $\alpha 6$ (Figure 8). Even though we performed modeling for the *bmr6-27* variant of SbCAD2, the number of amino acid substitutions and the consequent potential disruption due to the change in chemical properties is beyond the capabilities of the energy minimization-modeling algorithm. Additional experiments, such as circular dichroism spectroscopy or X-ray crystallography, are required to validate the correct model for this mutant.

Semiquantitative RT-PCR analysis of *SbCAD* family members in normal and *bmr* plants: Comparison of the transcript levels of *SbCAD2* in the near-isogenic pair BTx623/BTx623-*bmr6-ref* showed that in the latter the transcript levels of *SbCAD2* were downregulated (Figure 9B). Such downregulation of transcripts with premature stop codons is known to occur as a result of nonsense-mediated RNA decay (BOUT and VERMERRIS 2003; CONTI and IZAURRALTE 2005). It is likely that the *bmr6-ref* allele is a null mutation. In contrast, missense mutations are unlikely to trigger mRNA degradation. Indeed, expression analysis showed that the transcript levels of *SbCAD2* were not severely reduced in the *bmr6-3* mutant compared to its wild-type isolate N3 (Figure 9A).

The expression of the *CAD* family genes varied according to tissue type and stage of development of the plant. Table 3 summarizes the expression differences among the growth stages and tissues tested. The presence of the *bmr6-ref* or *bmr6-3* allele of the *SbCAD2* gene in homozygous form did not alter the pattern of expression of the other *CAD* family members, with the exception of *SbCAD5*. This gene was upregulated in the shoot of all the *bmr6-ref* plants, and two of the *bmr6-3* plants at stage 2. However, one of the wild-type plants also showed

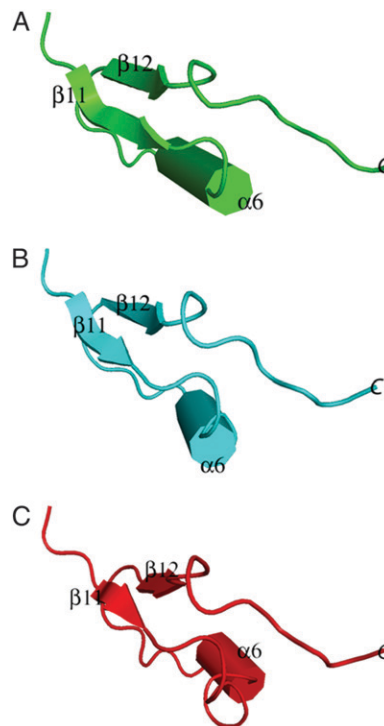


FIGURE 8.—Representation of the configurations of the secondary structural elements $\beta 11$, $\alpha 6$, and $\beta 12$ in (A) wild-type SbCAD2 and (B and C) the *bmr6-27* variant of SbCAD2. B and C represent two possible configurations of the corresponding secondary structure. In B, significant repositioning of the $\alpha 6$ -helix occurred. In C, a significant reduction of the secondary structure in all three structural elements occurred.

upregulation of this gene. Additional expression experiments with field-grown N6 and *bmr6-ref* plants confirmed strong plant-to-plant variation in the level of expression of the *SbCAD5* gene (data not shown), but in general, this gene was more frequently upregulated in *bmr6-ref* plants than in wild-type plants.

DISCUSSION

As previously shown in other plants species, *CAD* is encoded by a small gene family in sorghum. Several genes with high similarity to the *bona fide CAD* genes are present in the sorghum genome. Of the 14 *CAD* genes identified with homology searches, there is evidence that at least 10 genes are expressed, on the basis of the presence of corresponding ESTs and expression analysis. Two loci in the genome contain closely related groups of *CAD* genes. The *SbCAD8 1-4* cluster is orthologous with the rice *OsCAD8* cluster described by TOBIAS and CHOW (2005). Although the rice and sorghum genes within this cluster are highly similar, the gene arrangement differs between the species, raising the question whether the duplication event occurred prior to the divergence of the species or whether the clusters are the result of independent duplication events. The cluster is divided in two main branches, each of which contains sorghum

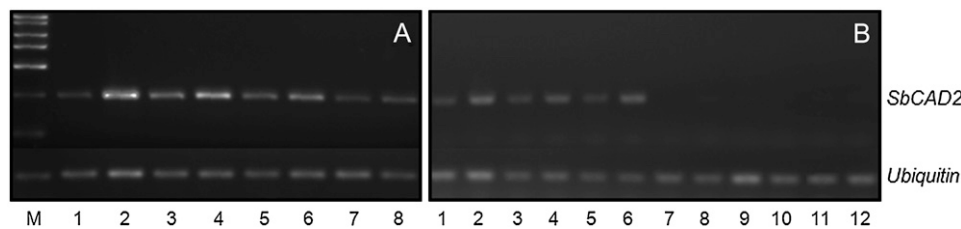


FIGURE 9.—The *bmr6-ref*, but not the *bmr6-3* mutation, causes downregulation of *SbCAD2* expression in 2-week-old plants. (Top lane) Products from RT-PCR with primers specific for a fragment of the cDNA encoding *SbCAD2*. (A) 1–4, N3 cDNA; 5–8, *bmr6-3* cDNA. (B) 1–6, BTx623 cDNA; 7–12, BTx623

bmr6-ref cDNA. Odd numbers correspond to cDNA obtained from shoots; even numbers cDNA obtained from roots. (Bottom lane) Products from RT-PCR with primers for *Ubiquitin* gene as a cDNA loading control. The size marker is Bioline Hyperladder II (cat. no. 33054).

and rice sequences. This suggests two duplication events, one before and one after speciation. This may explain the different orientation of the rice and sorghum genes. A homology search of the recently published maize draft genome sequence (<http://maizesequence.org>) with the sorghum *SbCAD8* genomic sequences revealed a single ortholog (AC197840.3_FG008). This maize gene is located on chromosome 7, which is syntenic with sorghum chromosome 2 (WILSON *et al.* 1999) on which the *SbCAD8* cluster resides. The presence of a single gene in maize, a species more closely related to sorghum than rice, is not consistent with the hypothesis that a duplication event occurred prior to the divergence of these species unless additional paralogs of AC197840.3_FG008 or remnants thereof are identified as the maize genome sequence reaches completion. The second gene cluster in sorghum (*SbCAD4 1-5*) has no reported orthologs in other species for which the genome sequence is available at this time. The closest rice sequence to the members of this group is a single gene, *OsCAD4*.

On the basis of the phylogenetic analysis, the sorghum *CAD* genes cluster into four groups. *SbCAD2* groups together with the *bona fide CADs* from other species. The high sequence similarity and conservation of important substrate binding pocket residues within this group point to a conservation of the gene function. Indeed, mutations or downregulation of the expression of genes belonging to this group have significant impact in lignin content and composition in tobacco, loblolly pine, poplar, and Arabidopsis (HALPIN *et al.* 1994; MACKAY

et al. 1997; RALPH *et al.* 1997; BAUCHER *et al.* 1999; LAPIERRE *et al.* 1999; PILATE *et al.* 2002; SIBOUT *et al.* 2003). Modeling *SbCAD2* over the crystal structure of *AtCAD5* revealed that even though *SbCAD2* contains a large number of residue changes, it remains similar in overall structure, with secondary structure preserved throughout. The close relationship of *SbCAD2* with the other members of this group supports the idea that it is the main sorghum *CAD* involved in lignification. Consistent with this hypothesis, we observed that *SbCAD2* is the predominantly expressed *CAD* gene in sorghum. In addition, a mutation in this gene is linked to the *bmr* phenotype in three independently introgressed isolines and it cosegregates with the *bmr* phenotype in a RIL population. Furthermore, sequencing of three independent *bmr6* alleles all revealed nonconservative mutations in the *SbCAD2* gene. Therefore, the evidence strongly suggests that mutations in this gene are responsible for the phenotype of the *bmr6* mutants.

Members of the other groups present much more diverse substitutions in the substrate binding pocket. These substitutions would allow for a wider variety of substrate specificity by changing the size, conformation, and hydrophobicity of the binding pocket. The role of these enzymes in lignification has not yet been elucidated. Only the *Populus SAD* has been implicated directly in lignification, catalyzing the reduction of sinapyl aldehyde to sinapyl alcohol (LI *et al.* 2001; BOMATI and NOEL 2005). Wound-inducible genes in parsley (*Petroselinum crispum*) and Arabidopsis cluster

TABLE 3

Gene expression of the sorghum *CAD-like* family based on semi-quantitative RT-PCR experiments

Gene	Stage 2		Stage 4		
	Shoot	Root	Leaves	Stem	Root
<i>SbCAD2</i>	Medium	Strong	Very strong	Very strong	Strong
<i>SbCAD4-2</i>	Very weak	Weak	Very weak	Very weak	Very weak
<i>SbCAD4-3</i>	Weak	Medium	Medium	Medium	Medium
<i>SbCAD5</i>	Variable	Very weak	Weak	Weak	ND
<i>SbCAD6</i>	Weak	Weak	ND	ND	Medium
<i>SbCAD7</i>	Medium	ND	Weak	Weak	ND
<i>SbCAD8-1</i>	Strong	Weak	Medium	Medium	ND

ND, not detectable.

together in group III (SCHMELZER *et al.* 1989; TREZZINI *et al.* 1992; SOMSSICH *et al.* 1996), suggesting that the genes in this group may be involved in the production of defense-related compounds (HAWKINS and BOUDET 2003). Kinetic analysis of the Arabidopsis genes in groups II, III, and V have shown that the activity against phenylpropanoid aldehydes is much lower than the Arabidopsis CADs belonging to group I. AtCAD7 has been shown to behave as an aromatic alcohol oxidoreductase of wide substrate affinity (SOMSSICH *et al.* 1996). Consequently, these *CAD-like* genes are currently annotated as putative mannitol or benzyl alcohol dehydrogenases. However, it is evident that there is some redundancy in the CAD metabolic network. Compositional studies of Arabidopsis, rice, and sorghum mutants in which the *bona fide* CAD genes are downregulated have shown that their lignin still contains a certain amount of the normal G and S residues (SIBOUT *et al.* 2003, 2005; ZHANG *et al.* 2006; SABALLOS *et al.* 2008). Reporter gene studies of the Arabidopsis CAD family showed that AtCAD7 and AtCAD8 are expressed in lignifying tissues (KIM *et al.* 2007). In the Arabidopsis double mutant *Atcad5/Atcad4*, which lacks activity of the two *bona fide* CAD proteins, the *AtCAD1* gene is highly upregulated (SIBOUT *et al.* 2005), suggesting it may act to compensate for the lack of the CAD activity from AtCAD5 and AtCAD4. Transgenic expression of the *AtCAD9* gene under control of the *AtCAD5* promoter was able to partially rescue the phenotype of the double mutant (EUDES *et al.* 2006). In our study, we did not find consistent upregulation of any of the *SbCAD* genes of groups III, IV, or V in the *bmr6* lines, suggesting that the steady-state level of the genes in the CAD family are sufficient to produce the observed levels of G and S units in the mutant. The *SbCAD8-1* gene is highly expressed in the aerial part of the plant. In the seedling (stage 2) it is the most highly expressed CAD gene. GUILLAUMIE *et al.* (2007a) analyzed the expression of the CAD genes in maize and its relation with tissue-specific lignification patterns. The EST sequence 3071507.2.III is the likely ortholog of *SbCAD8*. The spatiotemporal expression of this gene agrees with our results, but it differs in that it is not the main CAD expressed in the aerial portion of the plant at the seedling stage. Of the identified *CAD-like* genes, the genes in the *SbCAD8* cluster are the most similar to *SbCAD2* (52–48% identity, except *SbCAD8-3*). To our knowledge, the role of the *SbCAD8* genes in sorghum has not been investigated. On the basis of the high level of expression of these genes, we hypothesize that the *SbCAD8* enzymes may be responsible for the production of S and G residues observed in the *bmr6* mutants. Studies of a *Sbcad2/Sbcad8* double mutant could help determine the role of *SbCAD8* in lignification.

At this moment, it is difficult to attribute specific functions to the other sorghum *CAD-like* genes from groups III and V on the basis of phylogenetic relationships. We observed that the expression of the *CAD-like*

genes in sorghum varies with the stage of development and tissue type, in agreement with what was observed in maize (GUILLAUMIE *et al.* 2007a). Given the evolutionary relationship and the general agreement of gene expression patterns between maize and sorghum, it is likely that their functions are conserved. One exception may be *SbCAD5*. The maize ortholog, EST 4424417.2.III, is the predominantly expressed CAD in leaves and stems of maize seedlings. In contrast, *SbCAD5* showed variable expression levels in the leaves and stems of individual plants, both in seedlings and preflowering (stage 4) plants. More *bmr6-ref* and *bmr6-3* plants showed upregulation of *SbCAD5* than their wild-type isolines. It would be tempting to speculate that the gene is upregulated to compensate for the loss of *SbCAD2* activity, but the lack of consistent upregulation in all *bmr* plants does not support this conclusion. Another possible explanation is that *SbCAD5* gene is upregulated as part of a pathogen- or other stress-response mechanism, as it clusters in the same group as the Arabidopsis pathogen-inducible defense genes *AtCAD7* and *AtCAD8*. The *bmr* phenotype is associated with decreased lignin content and results in a fitness penalty in the plant (OLIVER *et al.* 2005a,b; PEDERSEN *et al.* 2005), and it is possible that the *bmr* plants are under increased stress compared to wild-type plants. It is interesting that while in the aerial parts of both maize and sorghum seedlings the *bona fide* CAD is not the most highly expressed CAD gene, the predominant CAD genes in those tissues represent nonorthologous paralogs in the two species.

To our knowledge, no orthologs of *SbCAD6* and *OsCAD6* (group IV) have been described in the literature. *SbCAD6* is mainly expressed in the roots of stage 4 plants, although at low levels. Both *SbCAD6* and *OsCAD6* (TOBIAS and CHOW 2005) contain a predicted peroxisomal targeting signal at their C terminus. Dehydrogenase activities in the peroxisome are involved in the metabolism of alcohol, branched amino acids, fatty acids, and plant hormones (REUMANN *et al.* 2004). Detailed biochemical and expression studies are needed to assign a physiological function to this gene.

We have described three different mutant alleles of *SbCAD2*: *bmr6-ref*, a likely null allele, *bmr6-3*, which produces a full-length protein with a single disruption in the cofactor binding site, and *bmr6-27*, encoding an enzyme with a disrupted secondary structure outside the active site. Although all three mutants produce the visual *brown midrib* phenotype, it is likely that the mutations have different effects on enzyme activity. There is evidence that the *bmr6-3* allele has a stronger effect on enzymatic saccharification of stover than the *bmr6-ref* allele, whereas the latter allele results in higher coniferaldehyde levels in the midrib tissue than either the *bmr6-3* or *bmr6-27* allele (SABALLOS *et al.* 2008). Now that the molecular basis of the mutations is known, variation in penetrance can be investigated. Tailoring of the effect of the activity of the enzyme may be achieved on the basis of

the choice of allele introgressed in a particular sorghum line. The development of molecular markers specific for each of those variants will greatly facilitate the use of the alleles in genetic studies and breeding programs. There are at least six additional known *bmr6* alleles available (SABALLOS *et al.* 2008), which opens the possibility of finding more useful variants.

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