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## Different mutations in the *ZmCAD2* gene underlie the maize brown-midrib1 (*bm1*) phenotype with similar effects on lignin characteristics and have potential interest for bioenergy production

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### Abstract

The maize *ZmCAD2* gene has been fully sequenced in several normal and *bm1* maize lines, highlighting a large diversity of mutations underlying the *bm1* phenotype. Mutations in three *bm1* lines (F2*bm1*, A619*bm1*, and 511J*bm1*) were found corresponding to short InDels inducing premature stop codons and truncated proteins. In two lines (511K*bm1* and 5803C*bm1*), mutations were limited to an only SNP or to a few SNP, modifying the catalytic sites, and likely inactivating the proteins. Results also established that the 5803*bm7* mutant was in fact a *bm1* mutant, with a sequence fully identical to the 5803C*bm1* sequence. The two new F7803*bm1* (natural mutant) and Ev2210*bm1* (transposon tagging *Mutator* investigations) both had a transposon insertion in the *ZmCAD2* DNA, resulting in a truncated protein, even if the mRNA was produced. The biochemical characteristics of the Ev2210*bm1* lignins corroborated the signature of *CAD2* deficiency in plants, with the presence of aldehydes and atypical compounds and linkages. Considering lignin structure and content, *CAD2* is likely a good target for the improvement of energy production based on maize and grass lignocellulosic biomass, including a greater susceptibility to environmentally friendly pretreatments, as it was shown in *bmr* sorghum. The interest in maize *bm1* hybrids for cattle feeding also should be considered because there seem to be little or limited negative effects of *CAD2* mutations on other agronomical traits.

**Keywords:** maize, brown midrib, *bm1*, lignin, ferulic acid, cell wall, degradability, biofuel, biogas

### Introduction

Maize brown-midrib (*bm*) mutants exhibit a reddish-brown pigmentation of the leaf midrib and stalk pith, associated with lignified tissues, after the plants have about five expanded leaves. The first *bm* mutants were described and investigated successively by Kiesselbach (1922), Eyster (1926), and Jorgenson (1931). The latter established that mutations of these first mutants were “due to identical factors” and that the “*bm* character” segregated as “a simple Mendelian recessive” trait. This gene was later named *bm1* when the *bm2* mutant was described by Burnham and Brinks (1932). A little later, two other genes inducing the *bm* phenotype were described as *bm3* (Emerson, 1935) and *bm4* (Burnham, 1947). The *bm2*, *bm3*, and *bm4* genes also originated from natural mutations and segregated as simple Mendelian recessive traits. The *bm1* mutation was assigned to maize chromosome 5 (bin 5.04) in position 77.5 cM, close to the centromer (Jorgenson, 1931; MaizeGDB, [www.maizegdb.org](http://www.maizegdb.org)). The three other *bm2*, *bm3*, and

*bm4* mutants were later mapped in bins 1.11, 4.05, and 9.07, respectively (MaizeGDB database). No new maize *bm* mutations were characterized for nearly 60 years after the Burnham's paper (1947), despite the mention of additional natural *bm* mutants in the MaizeGDB database. Recent allelic tests of these latter mutants highlighted the three novel *bm5*, *bm6*, and *bm7* loci (Haney et al, 2008; Ali et al, 2010). While *bm5* and *bm7* mutations are still uncharacterized, the *bm6* mutations have been mapped to a 180 kb region of bin 2.01. Ten underlying candidates were proposed (Chen et al, 2012a), including a gene encoding a zinc finger CCCH-type protein, of which different family members also colocalized with cell wall degradability QTLs in RIL progenies of Arabidopsis (Chavigneau et al, 2012) and maize (Courtial et al, unpublished data).

In the final step of maize monolignol biosynthesis, the cinnamyl alcohol dehydrogenase (*CAD*) protein(s) catalyze(s) the reduction of *p*-coumaryl, coniferyl, and sinapyl aldehydes to their corresponding alcohols, using NADPH as a cofactor, prior to their trans-

port to the wall and their polymerization into the lignin polymer. Two types of CAD genes are currently considered in plants, based on original investigations in eucalyptus. EgCAD1-type enzymes are short-chain alcohol dehydrogenases (Jornvall et al, 1995; Goffner et al, 1998), which are active as monomers on coniferaldehyde, but not sinapaldehyde (Hawkins and Boudet 1994; Damiani et al, 2005). A ZmCAD1 activity has been described in maize (Kanazawa et al, 1999), corroborating the existence of the two types of CAD enzymes also in (this) grass species. EgCAD2-type enzymes are conversely zinc-containing long-chain alcohol dehydrogenases, which are active as dimers, and are considered the predominant CAD proteins involved in lignification (Jornvall et al, 1987; Goffner et al, 1992; Hawkins and Boudet, 1994). In maize *bm1* plants, the activity of a CAD enzyme was shown to be reduced by 60 - 70% in the above-ground organs and by 90 - 97% in roots (Halpin et al, 1998). These authors therefore considered that “*bm1* is not a null mutation of the *ZmCAD*, but affects its expression, possibly through alterations in upstream or downstream non-coding regions”. Linkage analyses also showed that the *ZmCAD* locus was closely linked to RFLP markers corresponding to the position of the *bm1* mutation. Halpin et al (1998) thus concluded “that *ZmCAD* is very likely to be allelic to *bm1*”. This maize *ZmCAD* gene associated with the *bm1* mutation was latter shown to be orthologous to *EgCAD2*, and it has thus been named *ZmCAD2* (Guillaumie et al, 2007b). However, a lack of complete specificity of their polyclonal antibody left the possibility of reactions with other CAD or CAD-like proteins, and/or only partial reactions against the targeted *ZmCAD2* protein. In addition, gene expression, which was investigated based on northern analysis with degenerate primers based on the tobacco CAD2 protein sequence, of which specificity against the *ZmCAD2* sequence is likely partial, indicated a residual CAD expression in *bm1* leaves and stems. In addition, based on gene expression investigations with the MaizeWall macroarray, the *ZmCAD2* gene was shown to be under-expressed in *bm1* plantlets with a residual expression of 0.36 as compared to the control F2 line (Guillaumie et al, 2007a). This agreed closely with the residual activity described by Halpin et al (1998). In addition, two *ZmCAD2*-like genes and the *ZmCAD1* gene were simultaneously under-expressed in F2*bm1* plantlets. Similarly, based on Suppression Subtractive Hybridization (SSH) and micro-array data, Shi et al (2006) also established that several *ZmCAD* and *CAD*-like genes were down-regulated in young *bm1* plants (5-7 week-old plants).

The maize *ZmCAD2* gene (GRMZM5G844562) associated with the *bm1* mutation is located at position 98.993 Mbp on the chromosome 5 (bin 5.04) of the maize B73 genome (Schnable et al, 2009), upstream the estimated position of the chromosome 5 centromere. The maize *ZmCAD1* gene

(GRMZM2G179981) is also located in bin 5.04, in a downstream position (129.940 Mbp), upstream of the estimated position of the centromere. The latter gene only has distant paralogs, and at least one of them encodes a dihydroflavonol-4-reductase. The *ZmCAD2* also has no very close paralogs, and several of these *ZmCAD2*-like genes are now annotated as encoding mannitol dehydrogenase enzymes (maizesequence database, [www.maizesequence.org](http://www.maizesequence.org)). As a tentative conclusion, *ZmCAD1* and *ZmCAD2* are likely the two *ZmCAD* genes primarily involved in normal constitutive lignification of maize.

The consequences of the *bm1* mutation on maize lignin content and structure were first described forty years after the description of the mutation (Kuc and Nelson, 1964; Gee et al, 1968). Mature maize *bm1* plants have a lignin content that is reduced by 10 to 20%, a slight decrease in ferulic acid (FA) esters and substantially reduced contents (about 40%) in *p*-coumaric (pCA) esters and FA ethers (Provan et al, 1997; Barrière et al, 2004a). The frequency of *p*-hydroxyphenyl (H), guaiacyl (G) and syringyl (S) thioacidolysis monomers was similar in *bm1* and normal plants, showing that the *bm1* mutation does not specifically affect one of the lignin units. However, the reduced recovery of thioacidolysis monomers reveals that the frequency of lignin units involved only in  $\beta$ -O-4 bonds was about 50% lower in *bm1* plants than in lignins of normal plants, indicating that lignins of *bm1* plants were substantially enriched in carbon-carbon inter-unit linkages (Halpin et al, 1998; Barrière et al, 2004a). Lignins of *bm1* plants are also typified by a substantial incorporation of coniferaldehyde and, to a lower extent, of sinapaldehyde and *p*-hydroxybenzaldehyde into the polymer (Jacquet, 1997; Kim et al, 2002; Kim et al, 2003; Barrière et al, 2004a). This notable incorporation of *p*-hydroxy-cinnamaldehyde-derived compounds in *bm1* lignins was in agreement with the under-expression of the *ZmCAD2* gene.

Concerns over global climate changes, together with a growing worldwide demand for energy, have highlighted the crucial need for alternative resources to replace fossil fuels. Second-generation biofuels, based on lignocellulose materials, have opened up new avenues, including a large valorization of agricultural and woody residues. The latter do not compete with food supplies, unlike biofuels based on cereal or oleaginous grains. The European Union commissioner for “Climate Action” Connie Hedegaard indeed recently said that “we cannot morally afford to build a very big industry on something that is not good for the environment or for food prices”. However, the biological conversion of cell wall carbohydrates, mainly located in the secondary lignified plant cell walls, into fermentable sugars is hindered by their association with lignins, as well as with *p*-hydroxycinnamic acids in grasses. A similar situation has existed for decades in breeding programs towards higher digestibility and energy value of forage plants. Improvement of maize

energy value, for both cattle nutrition and industrial purposes, can be based on crosses and breeding with lines for which high values of cell wall degradability have been shown (Barrière et al, 2009; Barrière et al, 2010). However, the use of mutants with similarly higher cell wall degradability is also a relevant strategy in plant breeding. A mutant gene with major effect can indeed be more easily and rapidly backcrossed in elite lines based on marker assisted selection than several quantitative traits originating from one or more genetic resources.

Based on the B73 *ZmCAD2* sequence available from the Maize Genomics Sequencing Project (MGSP, [www.maizesequence.org](http://www.maizesequence.org)), the objective of this work was to sequence the *ZmCAD2* gene in *bm1* lines in comparison with their normal counterpart, including the reference line F2bm1, several new *bm1* mutants (Ali et al, 2010), and a new natural *bm1* mutant identified in an INRA nursery. In order to strengthen the involvement of the *ZmCAD2* gene in the *bm1* phenotype, a novel maize mutant obtained from transposon tagging in the *ZmCAD2* gene was simultaneously considered and characterized during these investigations. Transcriptomic approaches were also performed in order to estimate the respective effects of different *bm1* mutations on *ZmCAD* gene expression. In addition, because conflicting results were observed in new allelism tests, the *bm7* mutant was added to the sequencing investigations. Finally, the potential of the *bm1* mutation was discussed for further improvement of plant energy value for both cattle feeding and biofuel production.

## Materials and Methods

### Origins of *bm1* and *ZmCAD2* mutant lines

The F2bm1 line was obtained after seven backcrosses by the early flint INRA F2 line of the original *bm1* source (95033-47) supplied to INRA Lusignan by LF Bauman of Purdue University (West Lafayette, Indiana, USA) in 1972. This INRA F2bm1 genetic resource was also used in investigations by Halpin et al. (1998). Seeds of A619 and A619bm1 were supplied by N de Leon of University of Wisconsin (Madison) to INRA Lusignan in 2009. Seeds of the recently identified new *bm1* mutants 5803Cbm1, 511Kbm1 (or 5803Lbm1) and 511K, 511Jbm1 (or 5803Jbm1) were obtained in 2010 and 2011 from the Maize Genetics Cooperation Stock Center (MGCSC, [www.maizegdb.org](http://www.maizegdb.org)). The *ZmCAD2* mutant was isolated by the Biogemma company from transposon tagging investigations with the *Mutator* element, in the framework of the French national Génoplante program. Seeds of this mutant (*ZmCAD2-m2210::Mu* named thereafter Ev2210bm1), and the corresponding normal control seeds (Ev2210), resulted from three generations of self-pollinations after five generations of backcrossing (BC5S3) the original mutant plant with an elite line of the Limagrain company. The new INRA F7803bm was found in 2009 in an S7 generation of a flint prog-

eny (Laborde J and Bauland C, pers comm). During the allelism tests of the F7803bm mutation, brown-midrib plants were unexpectedly obtained in both F7803bm x F2bm1 and F7803 x 5803l7bm7 crosses. Consequently, the 5803l7bm7 line was added to the current investigations.

### Biochemical characterization of the *ZmCAD2* Ev2210bm1 mutant

Plants of Ev2210bm1 *ZmCAD2* mutant and their normal Ev2210 isogenic were grown in 2007 summer nursery in Mons (Puy de Dôme, France) according to current maize cropping conditions. Cell wall and lignin analyses were carried out on whole plants without ears, harvested at silage stage maturity. Data were obtained from samples of nine normal and nine mutant plants (without ears). Two replicates were analyzed. Neutral detergent fiber (NDF, or extract-free samples) and acid detergent lignin (ADL) were measured according to Goering and Van Soest (1970). Klason lignin (KL) was estimated according to Dence and Lin (1992). KL values are usually two to four times greater in grasses than ADL estimates, with the loss of an acid-soluble lignin part in the first step of the ADL procedure (Hatfield et al, 1994; Jung et al, 1997; Hatfield and Fukushima, 2005). ADL and KL, which are constituents of the cell wall, were expressed as percentage of NDF (ADL/NDF and KL/NDF). The *in vitro* dry matter digestibility (IVDMD) was estimated according to Aufrère and Michalet-Doreau (1983). The cell wall digestibility was then estimated according to Struik (1983) and Dolstra and Medema (1990) as the *in vitro* NDF digestibility (IVNDFD) assuming that the non-NDF part of plant material was completely digestible [IVNDFD = 100 x (IVDMD - (100 - NDF)) / NDF]. Thioacidolysis was performed from extract-free samples and according to Mir Derikvand et al (2008). In addition, some thioacidolysis experiments were conducted on exhaustively permethylated samples in order to evaluate the proportion of terminal units with free phenolic groups relative to internal units, according to a previously described procedure (Lapierre et al, 1988).

### Allele sequencing and PCR analyses

#### DNA extraction and sequencing

DNA of each genotype was extracted from 1 g of fresh leaf using a modified CTAB protocol (CIM-MYT, 2005). Sequencing was performed for each PCR fragment in both directions by Millegen ([www.millegen.com](http://www.millegen.com), France). Contigs were constructed using Staden software [<http://staden.sourceforge.net/>] (version v4.8b1)]. Sequences were aligned using MultAlin software [<http://multalin.toulouse.inra.fr/multalin/multalin.html>] (Corpet, 1988)]. The deduced amino acid sequences were obtained using Transeq software (<http://mobyli.pasteur.fr/cgi-bin/portal.py?#forms::transeq>). Amino acid sequences were aligned similarly using MultAlin software.

**Table 1** - Primer pairs designed for amplification of the whole DNA *ZmCAD2* gene, and for expression studies, based on the B73 DNA sequence.

	Forward primer (5'→3')		Reverse primer (5'→3')	Length (bp)
<b>Gene sequencing</b>				
ZmCAD2_1F1	GACCTCCTCGAAAGACGAAA	ZmCAD2_1R1	TGTCCCATCGAATGTTTGAA	1,256
ZmCAD2_2F1	TAATTTGGGGGAGCACTCTG	ZmCAD2_2R1	ATAGCTCCGGACTGGGATG	1,281
ZmCAD2_3F1	CTCCGTGTACGGTCCAGAAT	ZmCAD2_3R1	TGGTGGTTCAACCTCACAAA	1,279
ZmCAD2_3F2	CATGACGACAGGACAACCAC	ZmCAD2_3R2	GCATGCAAGATAACGCTGAA	446
ZmCAD2_4F1	CTTATGGTGACCAGGCAAGAG	ZmCAD2_4R1	GACGCAGAACTGAAGCACCT	999
ZmCAD2_5F1	CGACTCGCTGGACTACATCA	ZmCAD2_5R1	GGTCGCGGTATTCTGCTAAG	967
<b>Expression studies</b>				
ZmCAD1aF	CTTCCTCTCCGGACAAGTGTG	ZmCAD1aR	AGCTTTCGATGGTCTCCTTGACG	-
ZmCAD2cF	GGTCATACAACGACGTCTACTCTG	ZmCAD2cR	CCGGGATCTTACCACAAACTTC	-
ZmCAD4bF	TCAAGCCCAACGGCAAGATG	ZmCAD4bR	CAGGGTCTTGTCCCAATGATGAG	-
ZmUbiquitinF	CATTGTGCCCTGTTGAACTC	ZmUbiquitinR	AACAGCAACACCACAAACCA	-

*Primer design and PCR amplifications*

Pairs of primers used for the *ZmCAD2* gene sequencing were designed based on the available maize B73 sequence (GRMZM5G844562, [www.maizegenome.org](http://www.maizegenome.org), Table 1). Fragments of nearly 0.4 to 1.5 kb were amplified, encompassing the 5'UTR, the complete coding region, 1,000 bp before start codon and 1,000 bp after stop codon, giving a *ZmCAD2* investigated region close to 5 kb long. PCR reactions were performed in a final volume of 50 µl containing 1X PCR buffer and 0.5 U Taq DNA Pol (both MP Biomedicals, ref EPTQX925), 200 µM of dNTPs, 0.32 µM of 5' oligo, and 0.32 µM of 3' oligo. Fifty ng of genomic DNA was used as template. The program comprised 4 min at 94°C, followed by 35 cycles of 30 s at 95°C, 1 min at 60°C, and 1 min at 72°C, followed by 6 min at 72°C.

*Expression analyses*

For all eight normal (F2, 511K, Ev2210) and bm1 (F2bm1, 511Kbm1, 5803Cbm1, 511Jbm1, Ev2210bm1) genotypes, plants were grown in a glasshouse at INRA Lusignan in spring 2011. Lignin pathway genes had previously been shown to be highly expressed in the below-ear internode from tassel emergence to a few days after silking (Riboulet et al, 2009), a period corresponding to the end of the elongation phase and the deposition of the secondary cell wall in this internode. Consequently, below-ear internodes (without nodes) of three representative plants were harvested when tassels were clearly emerging out of the whorl. Samples were collected five hours after sunrise. To limit experiment costs, all plants of each line were pooled after cutting internodes into fragments nearly 1 cm long. All samples were immediately frozen in liquid nitrogen and stored at -80°C.

Total RNA was extracted and treated using the plant RNeasy mini kit and RNase-Free DNase Set (Qiagen, ref 70903 and 79254). RNA (1 µg) was reverse transcribed with 200 U of SuperScript III using 5µM oligo(dT) (Invitrogen, SuperScript III First-Strand

Synthesis System for RT-PCR, ref. 18080-051) according to the manufacturer's instructions. The resulting cDNAs was diluted 1:10. In addition to the *ZmUbiquitin* (GRMZM2G102471) gene used as standard, expression was investigated for the two *ZmCAD1* and *ZmCAD2* genes, and also for the *ZmCAD4* gene (GRMZM2G118610), which was originally annotated as a *ZmCAD2*-like gene (Barrière et al, 2009), but which is now considered to encode a mannitol dehydrogenase. Gene primer pairs were designed using the QuantPrime software to anneal near the 3' end of each transcript, often in 3'UTR area, ensuring primer specificity (Table 1). Real-time reverse transcription RT-PCR was performed on a LightCycler® (Roche Diagnostics) using the SYBR Green I Master (Roche Diagnostics, ref 04707516001). The PCR reaction mixture (20 µl) consisted of 3 µl water, 10 µl SYBR Green I Master Mix, 250 nM of each primer and 5 µl diluted cDNA template. The following LightCycler experimental run protocol was used including a denaturation program (95°C for 10 min), an amplification and quantification program repeated 45 times (95°C for 10 s, 60°C for 15 s, 72°C for 25 s with a single fluorescence measurement), a melting curve program (65-95°C with a heating rate of 2.2°C per second and a continuous fluorescence measurement), and finally a cooling step to 40°C. The PCR threshold cycle number of each gene was normalized with that of the *ZmUbiquitin* reference gene to calculate the relative mRNA levels. The Fisher test ( $P < 0.1$ ) was used for statistical analyses of quantitative RT-PCR data.

**Results***Allele sequencing of the ZmCAD2 gene in normal and bm1 near-isogenic lines**F2 and F2bm1 lines*

The F2 *ZmCAD2* allele differed from the B73 *ZmCAD2* allele in the 3' region, with 12 SNPs and two 54- and 30-bp long deletions starting from positions 4,031 and 4,116 (relative to the transcriptional start site in B73), respectively. In addition, amino

acid 349 is an asparagine in F2, while it is an aspartic acid in B73. The *ZmCAD2* alleles in B73 and F2bm1 were overall similar, likely related to the fact that the original *bm1* mutation used in F2 backcrossing occurred in a dent Corn Belt line and not in a European flint one. However, a two basepair (AC) insertion was identified in exon 3 of the F2bm1 line after the C base located in B73 CDS position 392 (Supplementary Figure 1). The ORF was consequently disrupted in F2bm1 by a premature stop codon, resulting in a predicted truncated protein of 147 amino acids, in comparison to the 367 amino acid long proteins in B73 and F2 (Supplementary Figure 2). Such a truncated *ZmCAD2* protein, in which the site involved in enzyme binding to its substrate, a catalytic site, and several binding sites to cofactors, were missing is very likely to be non functional, even if produced. The AC base insertion may therefore explain the *bm1* phenotype of the F2bm1 line. In addition, in F2, F2bm1, as in all investigated lines, in B73 CDS position 951, there was a synonymous substitution of an A base (B73) by a G base (all lines), the GGA and GGG codons both encoding for a glycine.

#### *A619 and A619bm1 lines*

Allele sequencing was only investigated in the area overlapping the AC insertion shown in F2bm1. The same mutation was highlighted in A619bm1 as the one identified in F2bm1. This suggests that the backcrosses of the *bm1* mutation in the A619 line were made with the same *bm1* mutant allele as the one used for the F2bm1 line, which is, given the origin of these lines, likely the original *bm1*-reference (*bm1*-ref) allele.

#### *511Jbm1 line*

A CGCG four base insertion was identified in the *ZmCAD2* exon 3 of the 511Jbm1 line, which occurred after the CG bases located in positions 366-367 of the B73 CDS (Supplementary Figure 1, *bm1*-J allele). As observed in F2bm1, this insertion disrupted the ORF and induced a 127 amino acid long truncated protein (Supplementary Figure 2), which is very likely non functional, even if produced.

#### *511Kbm1 line*

Allelic variation in the *ZmCAD2* CDS of the 511Kbm1 line, in comparison to B73, was reduced to only one SNP in exon 3 substituting the GGA codon into a GAA codon (CDS bp 401), thus substituting a glycine (G) by a glutamic acid (E) as the 134<sup>th</sup> amino acid of the *ZmCAD2* protein (Supplementary Figures 1 and 2, *bm1*-K allele). This unique mutation occurred in an area encoding the between-species conserved <sup>130</sup>PTQGGFA<sup>136</sup> amino acid motif (<sup>130</sup>PTQGEFA<sup>136</sup> in the mutant), located just before the  $\beta 9$  strand. However, a 21 bp deletion was also shown in the 5'UTR, in position 143-123 bp before the B73 ATG start codon, in which was inserted a 5 bp substitution establishing an extra TATATA box 128 bp before the *ZmCAD2* ATG start codon of the 511K line, in addition to the TATATA box located 40 bp upstream. These two

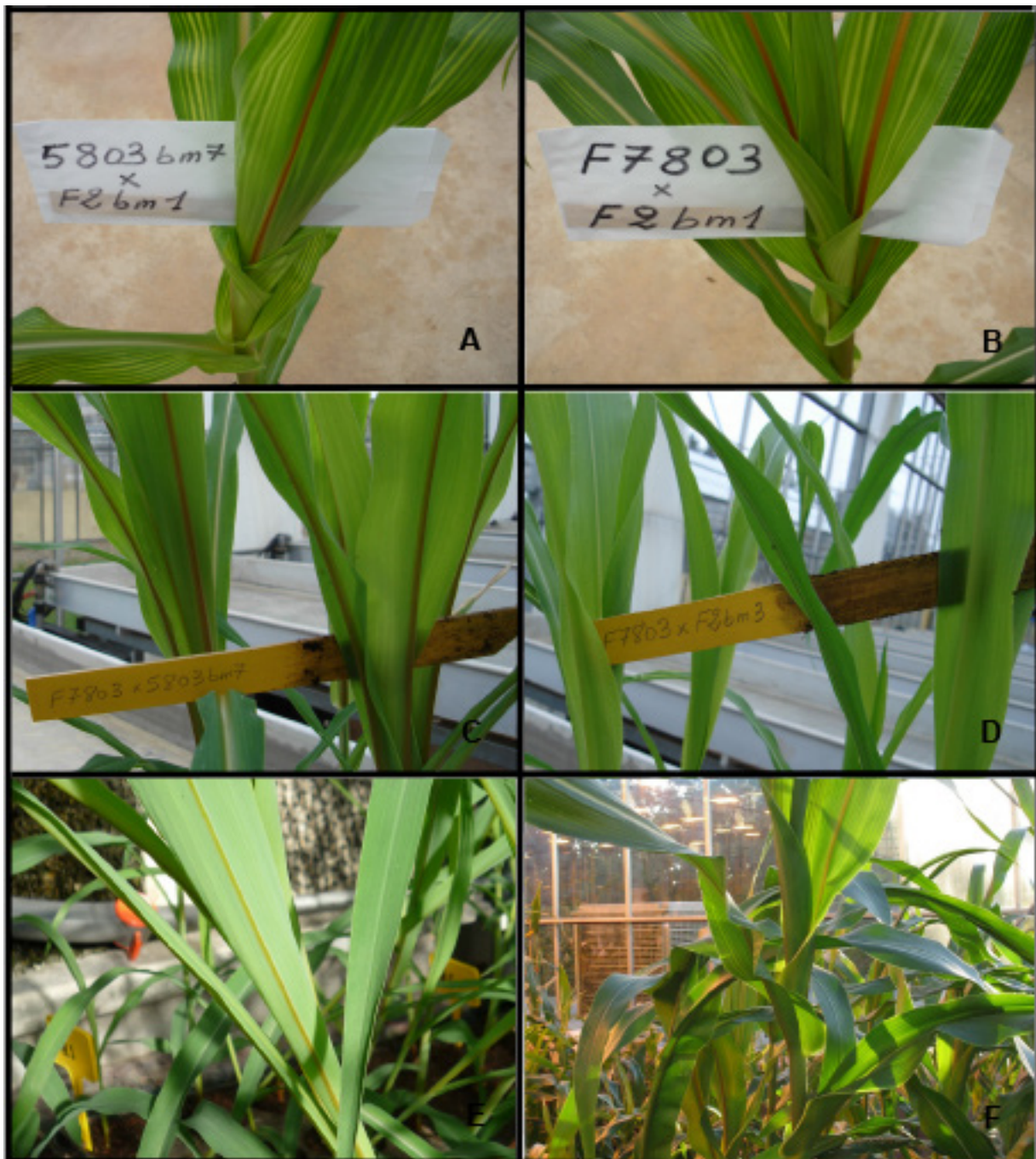
events were shown to be specific to the 511Kbm1 line and were not present in the 511K near-isogenic normal line. It was thus considered that either changes in protein structures inactivated its catalytic activity, or less likely that the transcription was reduced to a quasi-null level due to the extra TATA box.

#### *5803Cbm1 line*

Neither insertions nor deletions were shown in the *ZmCAD2* CDS of the 5803Cbm1 line, in comparison to B73, but the two sequences differed by two synonymous and three non synonymous SNPs. Synonymous SNPs occurred in CDS positions 192 bp (CCT/CCC, proline) and 237 bp (GGG/GGA, glycine). The first non-synonymous SNP was shown in exon 1 where a TCC codon was substituted by an ACC codon (CDS bp 76), encoding a threonine (T) instead of a serine (S) as the *ZmCAD2* 26<sup>th</sup> amino acid (Supplementary Figures 1 and 2). The second non-synonymous SNP occurred in exon 3, where the TCC codon was changed into TGC (CDS bp 410), encoding a cysteine (C) in place of a serine (S) as the 137<sup>th</sup> amino acid. The latter mutation, and the one shown in exon 1, likely did not greatly change activities of the encoded enzyme, as they both occurred in between-species genomic unconserved areas (Youn et al, 2006). The third mutation also occurred in exon 3, with a codon CAC changed into CGC (CDS bp 205, which substituted a histidine (H) by an arginine (R) as 69<sup>th</sup> amino acid in the conserved amino acid motif <sup>68</sup>GHEVVGXXXXXGXXV<sup>82</sup> leading to a <sup>68</sup>GREVVG<sup>73</sup> motif in the mutant (Supplementary Figures 1 and 2, *bm1*-C allele). The latter motif is involved in the binding of the zinc ion at the catalytic site of the enzyme, and the GHE conserved amino acid sequence also corresponded to the area separating the  $\beta 4$  and  $\beta 5$  strands (Youn et al, 2006). This H to R mutation probably alters the catalytic activity of the encoded enzyme, and is consequently likely the cause of the *bm1* phenotype. In addition, a very important polymorphism was shown in the *ZmCAD2* line first intron of the 5803Cbm1 line, which differed from B73 first intron by 54 SNP and 18 Indels. These changes could also modify the *ZmCAD2* gene expression in 5803Cbm1 mutant plants. Finally, the 5'UTR of the *ZmCAD2* gene in 5803Cbm1 mutant line exhibited the same 15 bp deletion and extra TATATA box as observed in the 511Kbm1 mutant line.

#### *5803lbm7 line*

The sequence of the *ZmCAD2* gene in the 5803lbm7 line was found to be fully identical to the sequence of the 5803Cbm1 line. This result was in agreement with new allelism tests done at INRA Lusignan showing that the 5803lbm7 line gave hybrids with brown midribs in crosses with F2bm1 (Figure 1). Allelism between *bm1* and *bm7* was corroborated by another allelism test done by Hongjun Liu (Iowa State University, pers com) and it was also in agreement with unpublished previous observations of Sarah Hake (2007, pers com). The mutation in the 5803lbm7



**Figure 1** - Allelism tests between 5807bm7 and F2bm1 (A), F7803bm and F2bm1 (B), F7803bm and 5803bm7 (C), Ev2210bm and F2bm1 (E, F), all four crosses giving brown-midribs, and F7803bm and F2bm3 (D), giving normal midribs.

line is definitely allelic to *bm1* (*bm1-7* allele) and this fact is now specified in the maizegdb database ([www.maizegdb.org](http://www.maizegdb.org), locus *bm7*).

#### *F7803bm1* line

Brown-midrib plants were obtained in crosses of F7803bm with F2bm1 (Figure 1), but not in crosses with F2bm2, F2bm3, or F2bm4, and similarly not with the newly described *bm5* and *bm6* lines. However, the F7803bm line also gave plants with brown

midribs in crosses with 5803bm7 (Figure 1). This fact further validated the allelism of the two *bm1* and *bm7* mutations. In contrast to investigations with the five previous *bm1* mutants, the sequence of the *ZmCAD2* gene in the F7803bm1 line was not fully obtained, with a failure in amplifying the whole second exon. PCR products corresponding to the second exon were small, with *ZmCAD2* homologies reduced to a few base pairs, which most often cor-

responded to primer sequence. The forward primer used was located 459 bp upstream of exon 2, and the reverse primer was located 629 bp downstream of exon 2 (these two primers enabled amplification of exon 2 in normal lines). The most probable situation in F7803bm1 would therefore be an insertion of a (retro-) transposon element in the exon 2 area, preventing PCR. In addition, there was an amino acid substitution in exon 4, with a glutamic acid replaced by an aspartic acid, corresponding in B73 CDS position 924 to a substitution of a G nucleotide by a T nucleotide (Supplementary Figures 1 and 2, allele *bm1-F*). Glutamic and aspartic acids are both diacid amino acids, and such a change is not expected to be responsible for the *bm1* phenotype. Nevertheless, the probable insertion of a (retro-) transposon close to exon 2 would very likely result in a protein disrupted before the region encoded by exon 4.

*Ev2210bm1* line

In contrast to previous spontaneously occurring *bm1* mutations, the *Ev2210 ZmCAD2 bm* mutant line was obtained from transposon tagging investigations with the Mutator element. Allelism tests showed that brown-midrib plants were obtained in crosses with F2bm1 (Figure 1), but not in crosses with F2bm2, F2bm3, or F2bm4, corroborating the involvement of *ZmCAD2* mutations in *bm1* phenotype. As it was the case for F7803bm1, the full sequence of the *ZmCAD2* gene in the *Ev2210bm1* line was also not obtained. However, 24 bp of the upstream part and 26 bp of the downstream part of the *Mutator* element sequences

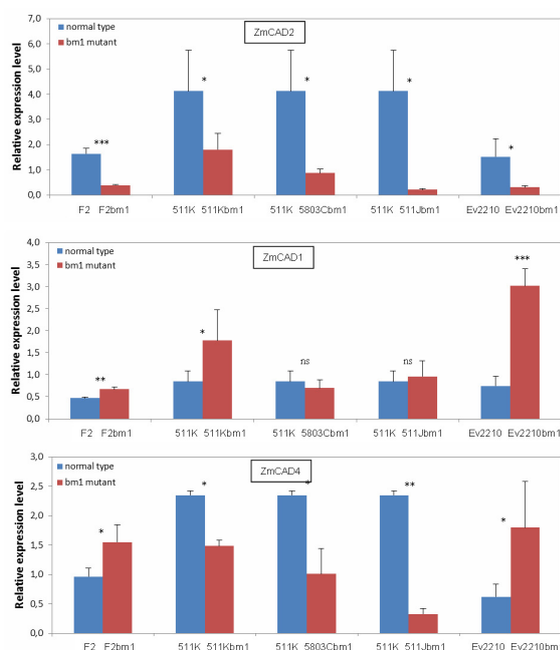
were obtained, showing that the *Mutator* element was located in this allele in exon 4, between position 678 and 679 bp of the B73 CDS. Lengths of a single copy of *Mutator* transposons are nearly 1,500 bp long, and they have a 220 bp long conserved terminally inverted repeats (TIRs). Such a long insertion, and chiefly the presence of the TIRs which cause very stable hairpin structures, would prevent an efficient PCR running. In any case, the *Mutator* element insertion induced a stop codon and disrupted the ORF, resulting in a 237 amino acid long truncated protein (Supplementary Figures 1 and 2, allele *bm1-M*), which is likely to be inactive if it is produced.

**Gene expression in *bm1* mutants and their normal counterparts**

The *ZmCAD2* gene appeared to be significantly under-expressed in all investigated *bm1* lines (F2bm1, 511Kbm1, 5803Cbm1, 511Jbm1, *Ev2210bm1*), in comparison to their normal counterparts (F2, 511K, *Ev2210*), corroborating the effect of *bm1* mutations on the expression of the *ZmCAD2* gene. In addition, the expression of the *ZmCAD1* gene was significantly up-regulated in three *bm1* mutants out of five, and especially in the *Ev2210bm1* mutant line (Figure 2). These results did not corroborate the lower *ZmCAD1* expression observed in *bm1* plants (Shi et al, 2006; Guillaumie et al, 2007a), that was, however, observed in much younger plants. The *ZmCAD4* gene, encoding a CAD-like protein with a mannitol dehydrogenase activity, had variable expression levels according to genotypes. *ZmCAD4* was over-expressed in F2bm1 and *Ev2210bm1* lines, while it was under-expressed in the three other *bm1* lines considered.

**Lignin content and structure in the *ZmCAD2 Ev2210bm1* mutant line**

The consequences of the *bm1* mutation on maize lignin content and structure have been shown to date in F2bm1 (*bm1-ref*) mutant plants (with the AC insertion), before it was firmly established that the *ZmCAD2* gene was affected in these plants. Relative to the wild-type plants, investigations in the *Ev2210bm1* mutant plants, altered in the *ZmCAD2*



**Figure 2** - Comparative expression of *ZmCAD2*, *ZmCAD1*, and *ZmCAD4* genes in normal and *bm1* maize plants (\* = P < 0.1; \*\* = P < 0.01; \*\*\* = P < 0.001, and ns = non significant differences). The *ZmUbiquitin* gene was used as standard.

**Table 2** - Cell wall traits in whole plants without ear at silage maturity stage for the normal and *ZmCAD2 Ev2210bm1* mutant plants.

	NDF	KI/NDF	ADL/NDF	IVNDFD
Normal	61.15	15.64	6.14	34.59
<i>Ev2210bm1</i>	62.01	14.55	5.24	39.27
F test	3.0 ns	15.6 *	53.4 **	26.0 **

NDF (neutral detergent fiber) as % dry matter (DM), ADL (acid detergent lignin) and Klason lignin (KI) % NDF, *in vitro* NDF digestibility (IVNDFD) as weight % of NDF. Data as mean values from 9 independent samples, differences between normal and mutant plants were non significant (ns) or significant at P < 0.01 % (\*) and 0.001 % (\*\*).

gene after transposon insertion, established that this mutant also displayed substantial variations in lignins and cell wall phenolics (Table 2). The lignin contents estimated by contents the Klason procedure and Van Soest (ADL) method were reduced by 6 and 17%, respectively, in Ev2210bm1 plants, while the cell wall content (referred to as NDF) was not changed. Substantial changes in lignin structure were also observed, with 30 - 40% fewer units only involved in  $\beta$ -O-4 bonds, as shown by the lower thioacidolysis yield (Table 3). Lignins in Ev2210bm1 plants are enriched in resistant inter-unit bonds, referred to as condensed linkages. Based on thioacidolysis investigations, the monomeric composition of the polymer was only slightly modified with a small increase in S unit frequency, a small decrease in G unit frequency, and no changes in H unit frequency. In addition to lignin-derived monomers, thioacidolysis of extractive-free samples released pCA and FA acids (Table 3). Previous studies have revealed that pCA is essentially ester-linked to lignins in maize cell walls (Ralph et al, 1994) and that most of these ester bonds do not survive the thioacidolysis procedure (Jacquet, 1997). Accordingly, the yield of thioacidolysis-released pCA accurately reflects the amount of *p*-coumarate esters. The origin of thioacidolysis-released FA is more complex because ferulate esters, which are linked to arabinoxylans, may be simultaneously linked to lignins by various bonding patterns. Anyway, obtained results suggested that the level of *p*-coumarate esters was substantially reduced in the cell walls of the Ev2210bm1 plants (nearly 30%), whereas ferulate units released by thioacidolysis were affected to a lower extent. Correlatively to lignin content and structure changes in Ev2210bm1 plants, their cell wall digestibility was improved by nearly 12%.

Following thioacidolysis and gas chromatography-mass spectrometry analysis of the lignin-derived monomers, signatures of CAD deficiency were identified in the Ev2210bm1 plants as previously observed in various CAD-deficient transgenic or mutant plants (Kim et al, 2002; Sibout et al, 2005; Thévenin et al, 2011). The mutant samples released

2 to 3 times more thioacidolysis monomers derived from coniferaldehyde end-groups than the control (Table 3). Moreover, the two G and S indene derivatives derived from coniferaldehyde and sinapaldehyde, which have been incorporated into lignins by 8-O-4 cross-coupling, were recovered in substantial amounts from Ev2210bm1 lignins and only as trace components from the control. These results corroborated those obtained in F2bm1 plants (Barrière et al, 2004a), even if aldehyde incorporations were shown to be lower in Ev2210bm1 plants than previously observed in F2bm1 plants. The difference between F2bm1 and Ev2210bm1 plants was probably related either to different penetrances of the two different types of *ZmCAD2* mutations in F2bm1 and Ev2210bm1, and/or to the different genetic backgrounds of the two lines. Among the differences, the much higher *ZmCAD1* over-expression was observed in Ev2210bm1, but not in F2bm1, plants (Figure 2).

Finally, when subjected to thioacidolysis and relatively to normal plants, Ev2210bm1 plants released 2 to 3 times higher amounts of the AG compound, which is a marker for the incorporation of free ferulic acid into lignins (Ralph et al, 2008). Despite this fact was not considered in previous years (Barrière et al, 2004a), a greater incorporation also occurred in F2bm1 lignins (data not shown), as well as in lignins of Arabidopsis CAD-C x CAD-D double mutant plants (Thévenin et al, 2011). The incorporation of ferulic acid into lignins allows new branching points with bi- and di-phenyl ether structures and consequently modifies the macro-molecular organization of the polymer (Ralph et al, 2008). The putative higher degree of branching of lignins in the Ev2210bm1 mutant was confirmed by the results obtained by thioacidolysis of permethylated samples which provides the proportion of terminal units with free phenolic groups relative to internal  $\beta$ -O-4 linked lignin units. In agreement with previous results obtained with maize lignins (Lapierre, 1993), thioacidolysis of the permethylated normal maize sample revealed that about 50% of G units are terminal units with free phenolic groups (Table 3), whereas this figure is higher for H

**Table 3** - Impact of the *ZmCAD2* Ev2210bm1 mutation on cell wall phenolics in whole plant without ear at the silage maturity stage based on thioacidolysis investigations.

Line	Thioacidolysis yield (H+G+S) ( $\mu\text{mole g}^{-1}$ KL)	Relative frequency of thioacidolysis monomers (% molar)			Minor thioacidolysis monomers ( $\mu\text{mole g}^{-1}$ KL)				Free phenolic groups (%) in $\beta$ -O-4 linkage			Thioacidolysis release <i>p</i> -OH cinnamic acids ( $\mu\text{mole g}^{-1}$ KL)	
		H units	G units	S units	coniferald end-groups	G Indene	S Indene	A <sub>6</sub>	H	G	S	pCA	FA
Normal	526.22	1.62	46.14	52.21	0.36	traces	traces	0.42	86.6	51.9	6.6	581.67	226.44
Ev2210bm1	294.89	1.60	43.57	54.82	1.18	9.54	7.27	1.27	91.5	61.3	7.8	420.22	218.89
F test	103.8**	0.2ns	12.0*	12.0*	172.4**	-	-	195.1**	6.4ns	161.3**	28.2**	40.4**	0.5ns

Thioacidolysis released H, G and S lignin monomers of coniferaldehyde (coniferald) end-groups, coniferaldehyde ether-linked at C $\beta$  (releasing G indene), sinapaldehyde ether-linked at C $\beta$  (releasing S indene), and of the marker compound (A<sub>6</sub>) for free ferulic acid incorporation in lignins. Thioacidolysis-released *p*-coumaric acid (pCA) and ferulic acid (FA). Percentage (%) of free phenolic groups in  $\beta$ -O-4 linked H, G and S lignin units, as determined by thioacidolysis of permethylated samples. Significant differences between normal and mutant plants indicated as in Table 2.

units (about 87%) and much lower for S units (less than 7%). However, in the Ev2210bm1 mutant, more than 60% of G lignin units are terminal units with free phenolic groups. The substantially higher frequency of free phenolic groups in lignins is a structural specificity which is common to all the CAD-deficient plants analyzed so far, including the F2bm1 maize mutant (Lapierre, 2010), and the signature of a lignin network disorganized in smaller domains.

## Discussion

While the *ZmCOMT* mutations were identified about 15 years ago in the maize *bm3* mutants (Vignols et al, 1995; Morrow et al, 1997), no investigations had firmly established up until now what gene was affected in *bm1* plants, even if it was greatly suspected that the *CAD2* gene was involved in the mutation(s). The sequencing of the *ZmCAD2* allele in eight *bm1* lines highlighted the diversity of genomic events underlying the *bm1* phenotype. Two- and four-base insertions, missense mutations, and transposon insertions are expected to lead to truncated or inactive *CAD2* proteins, thereby causing the *bm1* phenotype. The mutation of the *ZmCAD2* gene in *bm1* maize plants was also established in concomitant investigations in other genetic backgrounds (Chen et al, 2012b). The first intron of the *ZmCAD2 bm1-das1* mutant (Dow AgroSciences *bm1*) was shown to contain a 3,444 bp insertion, resulting in a chimeric mRNA containing a premature stop codon, and a truncated protein of 48 amino acids. The mutation identified in the *bm1* allele of the 515Dbm1 simultaneously investigated line was the same as the one observed in the F2bm1 and A619bm1 lines, consisting in an AC insertion in exon 3 that resulted in a 147 amino acid reduced protein. The AC insertion in the third *ZmCAD2* exon could thus be considered to be the first described *bm1 (bm1-ref)* mutation.

In addition, based on observations in nurseries and allelism tests, the *CAD2-bm1* mutation seems to occur or appear more frequently than other maize *bm* mutations. This fact considered from a very small number of events could actually correspond to a random effect, without any underlying biological mechanism. However, this fact could also be related i) to the complexity of the catalytic sites of the *CAD2* protein rapidly altered by a slight modification of the amino acid sequence, ii) possibly to a DNA sequence or a chromosomal position favoring transposon insertion, and iii) to a low or null effect of the *CAD2* mutation on agronomic value. Chen et al (2012b) have indeed observed that the CAG to TAG nonsense mutation occurred in *SbCAD2* of *bmr6* sorghum just one nucleotide apart from the AC insertion shown in maize *bm1* mutants. Moreover, although plants contain numerous genes coding for *CAD*-like genes, only *EgCAD2*-type, and to a lesser extent *EgCAD1*-type *CAD* enzymes, were shown with a primary physiological role in lignin biosynthesis. This situation results

from the presence in lignin-related *CAD* proteins of a few key residues permitting significant catalytic rates on monolignol precursors. The role of several of these key residues for efficient catalysis and monolignol biosynthesis had been especially established based on investigations in lead tree (*Leucaena leucocephala*) and switchgrass (Pandey et al, 2011; Saathoff et al, 2012). Correlatively, while a highly truncated protein was shown to be the determinant of the *SbCAD2 bmr6-ref* mutation in sorghum, the *bmr6-3* mutation indeed resulted from a unique mutation in the <sup>188</sup>GXGG(V/L)G<sup>193</sup> motif changed into a <sup>188</sup>GXGS(V/L)G<sup>193</sup> motif, affecting the binding affinity for the NADP+ cofactor (Saballos et al, 2009). In addition, the rice gold-hull-and-internode-2 (*gh2*) mutant, which exhibits a reddish-brown pigmentation in the hull and internodes which become golden yellow at maturation, has an altered *OsCAD2* gene with a G to A substitution in the fourth exon (Zhang et al, 2006). This substitution induced the replacement of a zero-charge glycine by an electro-negative aspartic acid, leading to an inactive protein, as confirmed by recombinant protein experiments. As observed in maize and sorghum, the *CAD* activity was drastically reduced in rice *gh2* mutant, indicating that other *CAD*-like enzymes only partly substitute the *OsCAD2* deficiency, at least in rice for coniferyl alcohol biosynthesis (Zhang et al, 2006).

All cell wall traits shown in plants of the Ev2210bm1 mutant were indeed very similar to the ones observed in the F2bm1 plants (Barrière et al, 2004a), corroborating the involvement of *ZmCAD2* mutations in the *bm1* phenotype. Both F2bm1 and the Ev2210bm1 plants had reduced contents in lignins and in *p*-coumarate esters, which are correlated with a higher cell wall digestibility. Both exhibited important structural alterations of their lignins, which correspond to the specific signature of *CAD* deficiency, namely a higher frequency of condensed bonds and of free phenolic groups and the increased incorporation of coniferaldehyde, sinapaldehyde and free ferulic acid in lignins (Lapierre et al, 2004; Ralph et al, 2008). However, in addition to the incorporation of aldehydes and atypical compounds into lignins, thioacidolysis yields indicated the synthesis of regular monolignols in *CAD2* mutant plants, reaching nearly 55 % of the normal  $\beta$ -O-4-linked monomeric units in the Ev2210bm1 maize mutant. *EgCAD1*-type and *CAD*-like genes acting on *p*-hydroxy-cinnamaldehydes, in addition to a possible residual *CAD2* activity in mutant with modified but not truncated protein, should therefore be considered. Expression data showed a probable substitution role of the *ZmCAD1* gene in Ev2210bm1 and, to a lesser extent, in F2bm1 and 511Kbm1. Corroborating the substitution by other *CAD* or *CAD*-like enzymes, the truncated *SbCAD2* protein was not detected in extracts from *bmr6* sorghum plants, but *CAD* activity was still detectable in mutant plant tissues (Pillonel et al, 1991; Saballos et al, 2009). Nev-

ertheless, if other CAD proteins (including *EgCAD1*-type) can be active on cinnamyl substrates, brown midrib or golden phenotypes and deep alterations in lignin composition establish that *EgCAD2*-type genes encode the main CAD proteins involved in the regular monolignol biosynthetic pathway of grasses.

The interest of CAD2-deficient plants in industrial processes was first shown with the demonstration of the significantly improved pulping capacity of poplar and pine CAD mutants (Baucher et al, 1996; Lapierre et al, 1999; Lapierre et al, 2000; O'Connell et al, 2002; Gill et al, 2003). In grasses, for bioenergy production, the efficiency of the *EgCAD2*-type mutation or deregulation towards significant increases of enzymatic hydrolysis and/or conversion rate into bioethanol (with or without pretreatment) has been shown in sorghum and switchgrass plants, without significant negative effects on agronomic value (Saballos et al, 2008; Corredor et al, 2009; Sattler et al, 2009; Sattler et al, 2010a; Sattler et al, 2010b; Scott et al, 2010; Saathoff et al, 2011). In maize, most investigations with *bm* genes have been done with the *bm3 COMT* mutant, because this latter induced the highest improvement in cell wall digestibility and energy value for cattle feeding (Barrière et al, 2004b). Similarly, high increases in the release of fermentable sugars were shown in *bm3* mutant plants (Vermeris et al, 2007). However, an important decrease in agronomic value has been simultaneously shown in nearly all early *bm3* investigated hybrids, a fact that could possibly be closely related to the germplasm of these ancient investigated hybrids. None of the considered hybrids were related to the lodent genetic resource, of high agronomic value and standability. Several *bm3* hybrids are now indeed available in different markets, including the US market, with proven greater efficiency in dairy cow feeding (at least 28 published investigations between 1976 and 2012, most of them in Journal of Dairy Science). The feeding value of maize *bm1* hybrids has not been extensively investigated, likely as the first digestibility measurements have shown the lower improvement of maize *bm1* genotypes in comparison to *bm3* near-isogenics (Barnes et al, 1971; Lechtenberg et al, 1972; Barrière and Argillier, 1993). Nevertheless, the agronomic value of maize *bm1* hybrids (and *CAD2* mutants) is seemingly less modified than the one of *bm3* hybrids (and *COMT* mutants). Based on INRA Lusignan unpublished data from the two old INRA260 and LG11 hybrids, average *in vivo* NDF digestibility values were equal to 50.6, 55.7, and 59.4% in normal, *bm1*, and *bm3* isogenics, while DM yields were equal to 12.7, 12.5, and 11.8 t ha<sup>-1</sup>, respectively. On the contrary, recent investigations strengthened the interest of maize *bm1*-type mutants, based on CAD down-regulation, for industrial purposes. Transgenic CAD-RNAi maize plants had stem cell walls with a slight reduction of lignin content, but the stems of deregulated plants were nevertheless more degradable than their

normal counterparts. Fermentation assays also revealed that CAD deregulated plants produced higher levels of ethanol compared to normal ones (Fornalé et al, 2012). Moreover, the degree to which lignin polymers incorporate various phenolic compounds in place of the three regular constitutive monolignols is surely underappreciated (Ralph, 2010). In addition to ferulic acid and hydroxy-cinnamaldehydes, unusual monomers including acylated hydroxycinnamyl alcohols, dihydro-hydroxycinnamyl alcohols, hydroxy-benzaldehydes and other hydroxycinnamic acids, can be incorporated into lignins of wild-type plants (Vanholme et al, 2012). Because plants could tolerate shifts in lignin composition with no or lower impact on growth than observed for reduced lignin contents, the substitution of some fraction of the three regular monolignols by unusual or alternative monomers through breeding, directed mutagenesis, or genetic engineering is thus a relevant strategy to tailor lignins in bioenergy crops so that cell walls would be more susceptible to biomass pretreatments (Eudes et al, 2012; Vanholme et al, 2012). The interest of unusual monomers incorporation into lignins for bioenergy production was also highlighted with *bmr6* sorghum. Glucose yields were indeed improved by at least 25% from sorghum biomass of *bmr6* plants, compared to normal isogenics (Saballos et al, 2008; Dien et al, 2011). Several *bmr6* (and "*bmr*") sorghum hybrids have thus been registered, or are in registration process, for both European and American markets. The higher efficiency of *bmr6* sorghum for animal feeding has also been shown from dairy cows experiments. Cows fed *bmr6* silage in their diets had 16% higher milk yields in comparison to similar diet with normal sorghum (Oliver et al, 2003). However, milk yields were equal for *bmr6* sorghum and normal maize silages, highlighting again the higher cell wall degradability in maize than in sorghum. In addition, the modification of the lignin structure could also be considered to increase cell wall degradability (Zhang et al, 2011). In a set of maize inbred lines, more condensed lignins were thus shown to be more favorable to an increased cell wall degradability than  $\beta$ -O-4 rich lignins. Finally, for both bioenergy production and cattle feeding, the successful breeding of improved maize (and C4 grasses) genotypes with more degradable cell walls, based on different lignin polymer organization, incorporation of alternative monomers, and reduced ferulate cross-linkages, requires more knowledge about phenolic compound biosynthesis in plants and about their coupling in the secondary walls.

Different tentative conclusions can be considered from the current investigations in *bm1* maize. Several very different events are responsible for the *ZmCAD2-bm1* mutation in maize. The *ZmCAD2* gene encodes the predominant CAD for monolignol production, even though other CAD gene(s) could be simultaneously involved, or may have substitutive

activity. Comparing the biochemical traits in transposon tagged and disrupted mutant highlighted the relevance of the marker characteristics that were shown in CAD2 deficient plants, including aldehyde and atypical compounds and linkages. Even if further experiments need to be done, CAD2 is likely a good target for the improvement of energy production based on maize and grass lignocellulose biomass, including also a greater susceptibility to (environmentally friendly) pretreatments (Maehara et al, 2011; Wu et al, 2011). These specific properties are related to the presence of unusual compounds incorporated into lignins, inducing a modified structure of the polymer spread over carbohydrates in smaller domains. Despite the interest of *bm1* maize hybrids for cattle feeding has not yet been established, contrarily to the proven efficiency of *bm1* plants for bioenergy production, the breeding of maize *bm1* hybrids is likely promising because the *ZmCAD2* mutations seem to induce limited unfavorable consequences on other agronomical traits, even if opposite results could be observed with a few genetic backgrounds (Lorenz et al, 2009).

For the improvement of cell wall traits for both feeding and industrial uses, genetic targets should be considered in the whole set of genes involved in secondary wall biosynthesis. Genes involved in lignin production are upstream regulated by MYB and NAC transcription factors (Sonbol et al, 2009; Fornalé et al, 2010; Zhong et al, 2011; Gray et al, 2012). Ferulate driven cross-linkages are both dependent on ferulic acid (more probably of feruloyl-CoA) and arabinoxylan biosynthesis (Barrière et al, 2009; Hatfield and Marita, 2010; Jung and Phillips, 2010; Piston et al, 2010; Jung et al, 2011). Lignified tissue patterning is regulated during plant growth by members of several gene families, including members of the zinc finger and HD-ZIP families (Barrière et al, 2009). Search for transposon-tagging mutants with improved cell wall degradability should indeed be a relevant strategy for the discovery of genes with still unknown important role in secondary wall assembly (Vermerris et al, 2007; Jung and Phillips, 2010). Mutants and favorable alleles of all these genes have then to be considered as targets during maize (and grass) breeding for enhanced energy value. However, while plant breeding for animal feeding and bioenergy production both requested the knowledge of genetic mechanisms involved in secondary wall assembly, the breeding of bioenergy plants has also to take into account the lignin network susceptibility to mild-alkali pretreatments.

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