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Switchgrass Contains Two Cinnamyl Alcohol Dehydrogenases Involved in Lignin Formation

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Switchgrass Contains Two Cinnamyl Alcohol Dehydrogenases Involved in Lignin Formation

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Abstract Lignin content of switchgrass (*Panicum virgatum* L.), a bioenergy species, is a critical determinant of biomass quality since it can negatively impact conversion of biomass into liquid fuels via biochemical platforms. Cinnamyl alcohol dehydrogenase (CAD) is a key enzyme in lignin biosynthesis. Here, we have shown that cv. Kanlow switchgrass contains at least two closely related CAD genes (*PviCAD1* and *PviCAD2*) that code for proteins containing highly conserved domains and residues that identify them as bona fide CADs. Both recombinant proteins displayed substrate kinetics consistent with their presumed role in cell wall lignification. Proteomic and immunoblotting detected CAD containing spots in inter-

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Department of Biology, University of Nebraska-Kearney, Kearney, NE 68849, USA node protein extracts, and proteomic analyses demonstrated that both CADs were expressed. In planta CAD activity, CAD protein levels were observed at all stages of tiller development. A real-time qPCR analysis of the two CADs and one CAD-like sequence indicated that transcripts coding for PviCAD1 were present in greater abundance than those coding for PviCAD2. Transcripts for a third CAD-like sequence (*PviAroADH*) were present at intermediate levels as compared to *PviCAD1* and *CAD2*. The predicted protein sequence of PviAroADH indicated that it was an enzyme unrelated to lignification based on phylogenetic and protein modeling data.

Keywords Cinnamyl alcohol dehydrogenase · Internodes · *Panicum virgatum* L. · Proteomic identification · Recombinant enzyme characterization · Tillers · Switchgrass

Introduction

Second generation biofuels, if properly developed, are likely to play a part in alleviating carbon emissions while at the same time improving energy security [3]. In the US, switchgrass (*Panicum virgatum* L.) is being developed as a potential feedstock for these purposes, and intensive efforts are underway towards developing cultivars to serve as dedicated feedstocks. Perennial plants, such as switchgrass, have the advantage of requiring fewer chemical inputs than row crops, enhance soil organic carbon, and can be grown on marginal land. On-farm field research has also reported a highly positive overall energy balance for switchgrass grown for biomass applications [44].

Understanding the mechanisms of cell wall synthesis is important since plants bred or engineered with easier to digest cell walls may enhance fuel yields or ease downstream processing steps resulting in reduced energy requirements for bioconversion [12, 40]. Although broadly similar in composition to dicots, grass secondary cell walls have significant amounts of lignin, ferulic, and *p*-coumaric acids, which can exist freely or form ester- or ether-linkages between lignin and cell wall polysaccharides [35, 51]. As a major component of grass secondary cell walls, lignin has been implicated as a key factor in the recalcitrance of biomass towards hydrolytic enzymes used in cell wall deconstruction [10, 17, 28].

Lignin is a complex heteropolymer whose monomers are derived from the deamination of phenylalanine followed by numerous side chain modifications, hydroxylations, and *O*methylations of cinnamic acid [7]. The heteropolymer has generally been characterized as being mainly comprised of three different phenylpropanoid subunits: *p*-hydroxyphenyl (H-lignin), guaiacyl (G-lignin), and syringyl (S-lignin) [28], although other derivatives such as acylated lignin units and hydroxycinnamyl aldehydes were also found capable of being incorporated [7]. Monocot and dicot lignin generally have contained similar amounts of G and S units, but grass lignin was also found to contain a small, but significant amount of H units [51].

Lignin biosynthesis has been suggested to require a suite of at least ten enzymes [28]. A key enzyme involved in the synthesis of lignin subunits is cinnamyl alcohol dehydrogenase (CAD). Essentially, CAD catalyzes the conversion of cinnamyl aldehydes to their corresponding alcohols (H, G, and S lignin monomers) which are subsequently incorporated into the lignin polymer. CAD is a member of the alcohol dehydrogenase superfamily and has been characterized from a number of different species [19, 22, 29-32, 42, 56]. The crystal structure of an Arabidopsis CAD, AtCAD5, has been solved [55]. Disruption of CAD through natural or engineered mutations can lead to plants with altered lignin levels and differential incorporation of monolignals into lignin [2, 21, 34]. For instance, a sorghum brown midrib phenotype, bmr6, was recently shown to be caused by a nonsense mutation in CAD [38, 43] and plants with this mutation had an altered lignin composition [34, 37].

Reduction in lignin amounts or changes in lignin composition can have beneficial effects. In forage grasses, decreasing lignin resulted in enhanced dry matter digestability for grazing animals [6, 11]. These same types of changes have also been shown to enhance sugar release or actual ethanol yields from cellulosic biomass of both dicots and monocots [14, 15]. CAD occupies a central role in lignin biogenesis and alteration in its activity often results in changing lignin content, leading to plants with potentially improved quality. However, plants contain a number of CAD and CAD-like proteins which do not all participate in lignin biosynthesis [23], and it is important to biochemically characterize those CAD or CADs that can participate in lignin biosynthesis. Such studies will clearly be an important part of selective breeding of switchgrass biomass cultivars with desirable traits for biofuel production.

Materials and Methods

Identification of PviCAD1, PviCAD2, and PviAroADH

PviCADs were identified during an initial analysis of a switchgrass EST resource [49]. Several switchgrass cDNA clones were identified during single pass library sequencing as being closely related to the ZmCAD2 gene of Zea mays encoding cinnamyl alcohol dehydrogenase. However, none contained the entire coding region. Therefore, RT-PCR was performed to isolate full length coding sequences using 1 ug total RNA isolated from root tissue of the cv. 'Alamo' or stem tissue from the cv. 'Kanlow'. These were reverse transcribed with Thermoscript II reverse transcriptase (Invitrogen Corp., Carlsbad, CA, USA) according to manufacturer's instructions and amplified with the primers: 5'-CATATGGGCAGCCTGGCGTCG-3' (PviCAD1 start) and 5'-GAATTCAGTTGGCCGGCGCCC-3' (PviCAD1 stop). Products from both reactions were then cloned into pCR2.1 and sequenced. One isolate from cv. Alamo root RNA (PviCAD1, GenBank accession no. GU045611) and one from cv. Kanlow stem RNA (PviCAD2, GenBank accession no. GU045612), which fell into two separate clusters of sequences after alignment with existing switchgrass EST sequences, were chosen for further analysis. These isolates were subcloned by digestion with EcoRI and NdeI into the corresponding sites of pET28a (EMD Chemicals, Inc., Madison, WI, USA). Recombinant proteins were produced in Escherichia coli, and purified as described earlier [43]. A survey of the available EST sequences and the number of clones identified for each CAD-like sequence are shown in Table 1. We have not yet found other unique CAD-like sequences in the available EST collections.

The putative PviAroADH coding region was assembled from available switchgrass EST sequences and corresponds to dbEST accession number GD038270 and NCBI Unigene assembly Pvr.3084. These sequences were identified as potentially coding for CAD-like proteins through a TBLASTN search, followed by Clustal and manual assembly of overlapping sequences. Primers based on the assembled sequence were used to amplify a product of the expected size by RT-PCR using internode RNA as a template. Identity of the product was confirmed by DNA sequencing. Despite numerous attempts, we were unable to obtain soluble, active

	Tissue									
	ST	AP	CR	FB	ET	LF/SD	RT	SL	CA	Total
PviCAD1 PviCAD2	$0 (0.000)^{a}$ 0 (0.000)	2 (0.007) 5 (0.017)	6 (0.015) 6 (0.015)	6 (0.022) 7 (0.026)	3 (0.011) 5 (0.018)	4 (0.014) 3 (0.011)	10 (0.036) 12 (0.043)	0 (0.000) 2 (0.016)	0 (0.000) 0 (0.000)	36 (0.016) 46 (0.020)
PviAroADH	6	1	3 (0.007)	2 (0.007)	0 (0.000)	2 (0.007)	5 (0.018)	1 (0.008)	0 (0.000)	22 (0.010)

 Table 1 Representation of individual cDNA clones by library

ST abiotic and biotic stress (normalized cDNA library), AP apex and stem library, CR crown library, FB early floral bud library, ET etiolated seedling library, LF/SD flowering/seed development library, RT root library, SL seedling library, CA callus (normalized cDNA library)

^a Numbers in parenthesis are % of total number of clones with high-quality sequence data

recombinant protein for PviAroADH and therefore could not document its enzymatic characteristics.

Enzyme Assays

Enzyme activity on each substrate was measured using reaction conditions similar to those previously published [19, 53] with adaptations for use with a microplate reader (BioTek Synergy HT, BioTek Instruments, Winooski, VT, USA). However, substrate levels were varied while cofactor concentration was kept constant at 200 µM in order to kinetically characterize the enzymes. Enzyme dilutions were prepared immediately before their use from 100 μ L aliquots that were stored at -80°C, and fresh dilutions were prepared approximately every 30 min in order to minimize the effects of enzyme activity loss. The amount of protein used for each assay was 3.51 and 4.73 ng for PviCAD1 and PviCAD2, respectively. Reaction rates were calculated from the first 90 s of absorbance data using PROC GLM in SAS 9.1 (SAS Institute Inc., Cary, NC, USA) and Michaelis-Menten kinetic parameters were estimated using nonlinear least squares whereby multiple initial starting values were chosen for each curve in order to ensure reliable convergence.

Plant Materials

P. virgatum cv. Kanlow N1 [52] was grown in fields at the University of Nebraska Agricultural Research Experiment Station near Mead, NE. Tillers were harvested at anthesis about 4 cm above soil surface, stripped of leaves and sheaths, separated into individual internodes, and immediately placed on dry ice for transport to the laboratory where it was stored at -80° C until used. The internode subtending the peduncle was labeled as internode 1 and the lowest (most mature internode) was labeled as internode 6 [39]. Most tillers generally possessed six well-distinguishable internodes. Tillers containing fewer than six internodes were not used.

Preparation of Internode Extracts for CAD Activity Assays

Internodes were ground using dry ice and a coffee grinder and the powdered material was used for further analyses. Centrifuge tubes (2.0 mL) were filled with approximately 300 mg of plant material, 1 mL of buffer (100 mM Tris-HCl, pH 7.5; 5 mM DTT; 5% (v/v) ethylene glycol) and 10 µL of a protease inhibitor mixture (Sigma-Aldrich #P9599). Samples were placed on ice and sonicated using a Branson Digital Sonifier 450 (Branson Ultrasonic Corp., Danbury, CT, USA) at 20 W three times with a 15 s pulse, with samples placed for 30 s into an iceethanol bath between pulses to allow for sufficient cooling. Sonicated extracts were then centrifuged for 15 min at 14,000 RPM and 4°C. Supernatant was decanted into new tubes and kept on ice until assayed for CAD activity. Protein content was determined using a colorimetric assay (Pierce 660 nm Protein Assay, Pierce Biotechnology, Rockford, IL, USA) and lysozyme was used for generating standard curves.

One- and Two-Dimensional Gel Electrophoresis

For one-dimensional SDS-PAGE, approximately 300 mg of ground internode materials were extracted as described earlier [43] and proteins were separated on 12% polyacryl-amide gels [26]. Approximately 25 μ g of protein was loaded into each well. Separated proteins were transferred to nitrocellulose membranes for immunoblotting as described below.

For two-dimensional gel electrophoresis, proteins were first extracted from 1 g samples of switchgrass internodes essentially according to a previously published method [54]. Final protein pellets were taken up in 180 μ L of sample buffer (8 M urea, 50 mM DTT, 4% CHAPS, and 0.2% ampholytes) and used for isoelectric focusing (IEF). IEF was performed using the ZOOM IPGRunner System (Invitrogen Corp., Carlsbad, CA, USA) and 7.0 cm immobilized pH gradient (IPG) strips with a pH range of

5.0-8.0. IPG strips were rehydrated overnight with 155 µL of protein extract dissolved in the sample buffer described previously. After rehydration, IPG strips were prepared for IEF using the manufacturer's protocol. IEF was conducted using a power supply (Thermo Electron Corp. PRO-6000) with the following steps: 175 V for 30 min, 175-2,000 V linear increase over 45 min, and 2,000 V for 3 h. After IEF, IPG strips were equilibrated in a buffer containing 6 M urea, 375 mM Tris-Cl, pH 8.0, 2.5% SDS, and 20% glycerol. Strips were then reduced in equilibration buffer with 65 mM DTT followed by alkylation in equilibration buffer containing 135 mM iodoacetamide. The second dimension was run on 12% polyacrylamide gels (C.B.S. Lite Slab Gel Kit, C.B.S. Scientific Company, Del Mar, CA, USA) and stained using Coomassie brilliant blue. We had conducted a larger proteomic analysis (data not shown) of several spots obtained from the two-dimensional gel shown in Fig. 5a; however, for this study we focused on the spots labeled 1-7 since this was the region expected to contain CAD proteins based on the predicted pI and molecular weights. We used the 2-D immunoblots as a guide to narrow these analyses. Protein spots numbered 1-7 in Fig. 5b were analyzed by mass spectrometry as described earlier [20].

Immunoblots

Proteins separated by SDS-PAGE were blotted to nitrocellulose membranes and probed with polyclonal antibodies raised to CAD as described earlier [43]. Polyclonal antibodies raised against soybean root nodule ascorbate peroxidase (AscPx) were used as loading controls. Preliminary data indicated that the signal for AscPx was relatively consistent across many different switchgrass tissues at similar protein loads (not shown) and could therefore serve as a control.

Real-Time qPCR

RNA was isolated from *P. virgatum* internodes using a previously published method [48]. After treating total RNA with RQ1-RNase-free DNase (Promega, Madison WI USA), 1 µg from each preparation was reverse transcribed using an anchored oligo dT primer mixture along with random hexamers and the Transcriptor First Strand cDNA kit (Roche Diagnostics, Indianapolis, IN, USA). Real-time PCR used an ABI Prism 7000 Sequence Detection System (Applied Biosystems, Foster City, CA, USA) and the SYBR Green PCR Master Mix. Primers used for these reactions were designed using the included Primer Express software and are given in Table 2. Reactions were performed in quadruplicate. Delta Ct values for the target

Table 2 Primer sequences

PviCAD1	FWD	5'-AGGCCAACGTTGAGCAGTA
	RVS	5'-CGATCCCTGCTGGTCTGG
PviCAD2	FWD	5'-GCGAGGTGGTGGAGGTC
	RVS	5'-CGATCCCGGCAGGCCT
PviAroADH	FWD	5'-CAGCCCCATGAAGTACCA
	RVS	5'-TTCATTGTCAGCAAGAGCG

genes were generated relative to PviCAD1 transcript level in internode 1. Three control genes were used for normalization: UCE1, eIF4 α , and GAPDH. Normalization factors were generated by geometric averaging of the control genes using geNorm version 3.4 [50], and a suitable pairwise variation (*V*) value indicated that the selected combination of control genes were stable. Normalization factors were then applied to the delta Ct values to generate relative expression levels.

Protein Modeling

Models of PviCAD1 and PviAroADH were created with SWISS-MODEL [4]. Both models used the published *Arabidopsis* CAD5 structure [55] as a template for model building (PDB ID 2CF6).

Results

Phylogenetic Analyses of Three Switchgrass CAD-Like Protein Sequences

ClustalW2 was used to align PviCAD1, PviCAD2, and PviAroADH protein sequences (Fig. 1). The results indicated that PviCAD1 and PviCAD2 share a very high degree of identity (95%) to each other, but only 45% identity to PviAroADH. In order to compare the protein sequences with other plant CADs, sequence data was obtained from publicly available databases and used to construct a phylogenetic tree (Fig. 2). The tree included a number of CAD and CAD-like proteins identified in different species including CADs from Arabidopsis (AtCAD1-9) as well as CADs from sorghum (Sorghum bicolor, SbCAD1-7). Other CADs utilized in this analysis have either been shown or suggested to directly impact lignification in sorghum (S. bicolor, bmr6 [43]), rice (Oryza sativa, OsCAD2 [56]), perennial ryegrass (Lolium perenne L., LpeCad1 [32]), corn (Z. mays, ZmCAD2 [18]), and sugarcane (Saccharum officinarum, SoCAD [45]). All of the CADs that have been shown to be involved in lignin biosynthesis clustered into a single clade. In turn, this clade further subdivided into monocot

PviCAD1	MGSLASERTVVGWAARDAAGHLSPYTYTVRNTGPEDVVVKVLYCGICHTDIHQA	54
PviCAD2	MGSLASERTVVGWAARDATGHLSPYTYTVRKTGPEDVVVKVLYCGICHTDIHQA	54
PviAroDH	MAPVAAAAEQQHTGKAAALAARDASGHLAPLTITRRSTGDDDVAIKILYCGMCHSDLHFI	60
PviCAD1	KNHLGASKYPMVPGHEVVGEVVEVGPEVSKHRVGDVVGVGVIVGCCRECRPCKANVEOYC	114
PviCAD2	KNHLGASKYPMVPRHEVVGEVVEVGPEVSKHRVGDVVGVGVIVGCCRECRPCKANVEOYC	114
PviAroDH	KNEWNNAMY PMVPGHE I AGVVTEVGKNVTKFKAGDRVGVGCMVNSCQSCDRCDEGFENHC	120
	** : ***** **:.* *.*** :*:*.:.** **** :**.	
PviCAD1	NKRIWSYNDVYTDGRPTQGGFASTMVVDQKFVVPIPAGLAPEQAAPLLCAGVTVYSPLKH	174
PviCAD2	${\tt NKRIWSYNDVYTDGRPTQGGFASTMVVDQKFVVPIPAGLAPEQAAPLLCAGVTVYSPLKH$	174
PviAroDH	${\tt RGIIFTYNSVDRDGTVTYGGYSSNVVVHERFVVRFPDAMPLDQGAPLLCAGITVYSPMKY$	180
	. *::**.* ** * **::*.:**.:*** :* .:. :*.********	
PviCAD1	FGLTAPGLRGGILGLGGVGHMGVKVAKALGHHVTVISSSSRKRAEAMDDLGADAYLVSSD	234
PviCAD2	${\tt FGLTAPGLRGGILGLGGVGH} MGVKVAKAMGH HVTVISSSSRKRAEAMDELGADAYLVSSD$	234
PviAroDH	HGLNVPGKHVGVLGLGGLGHVAVKFAKAFGMKVTVISTSPAKKQEALERLGADAFIVSKS .**** : *:*****:**:.**.***:* :*****:* :******:*	240
PviCAD1	AGAMAAAADSLDYIIDTVPVHHPLEPYLALLRLDGKHVLLGVVGEPLSFVSPMVMLGRKS	294
PviCAD2	AEAMAAAADSLDYIIDTVPVHHPLEPYLALLRLDGKHVLLGVVGEPLSFVAPMVMLGRKA	294
PviAroDH	ADEMKAAVATMDGIINTVSANMSIAPYMGLLKPNGKMIMVGLPVKPLEIPPFDLIMGNKT	300
	* * **. ::* **:**: .: **:.**: :** :::*: :**.: . :::*.*:	
PviCAD1	VTGSFIGSVDETAELLRFCVDKGLTSQIEVVKMGYVNEALERLERNDVRYRFVVDVAGSN	354
PviCAD2	VTGSFIGSIDETAELLRFCVDKGLTSQIEVVKMGYVNEALERLERNDVRYRFVVDVAGSN	354
PviAroDH	LAGSCIGGMRDTQEMIDVAAKHGVTADIEVVGAEYVNTAMERLAKADVRYRFVIDIGNTL ::** **.: :* *:::*:*:**** *** *:*** : ******:*::	360
PviCAD1	IEEQAAAAGAPAN 367	
PviCAD2	IEEATGAPAN 364	
PviAroDH	KSSE 364	

Fig. 1 Alignment of PviCAD1, PviCAD2, and PviAroDH protein sequences using ClustalW2

and dicot CADs, with PviCAD1 and PviCAD2 being most closely related to the other grass CADs, namely, ZmCAD2, SoCAD, and bmr6. In contrast, PviAroADH showed significant homology to a putative corn aromatic alcohol dehydrogenase and clustered with CAD-like sequences including AtCAD8 (ELI3) that have been associated with the reduction of aromatic aldehydes and not with cell wall lignification [47].

Biochemical Characterization of Recombinant PviCAD1 and PviCAD2

Recombinant enzyme activity of PviCAD1 and PviCAD2 were assayed on a variety of substrates in order to examine substrate preferences and enzyme kinetics. Both forward (aldehyde-to-alcohol) and reverse (alcohol-to-aldehyde) reaction rates were characterized using coniferaldehyde, sinapaldehyde, coniferyl alcohol, sinapyl alcohol, and coumaryl alcohol as substrates. Results from the enzyme assays followed Michaelis–Menten kinetics for all of the tested substrates. Estimates for kinetic constants $K_{\rm m}$, $V_{\rm max}$, and $k_{\rm cat}$, and $k_{\rm cat}/K_{\rm m}$ are provided in Table 3. For the alcohol substrates, PviCAD1 had the lowest $K_{\rm m}$ at 0.98 μ M

for coniferyl alcohol followed by coumaryl (4.2 µM) and sinapyl alcohols (15.6 μ M), respectively; V_{max} values were 170 nkat mg⁻¹ for coniferyl alcohol, 320 nkat mg⁻¹ for coumaryl alcohol, and 600 nkat mg⁻¹ for sinapyl alcohol, respectively. On aldehyde substrates, PviCAD1 displayed the lowest $K_{\rm m}$ for coniferaldehyde at 10.9 μ M while the $K_{\rm m}$ for sinapaldehyde was 14.5 $\mu M.$ Also, PviCAD1 had significantly higher V_{max} values on both aldehyde substrates when compared to alcohol substrates. The V_{max} of 2,530 nkat mg⁻¹ for sinapaldehyde was significantly higher than a $V_{\rm max}$ of 1,280 nkat mg⁻¹ when coniferaldehyde was used as a substrate. Because PviCAD2 shared a highly similar sequence to PviCAD1, it was only tested on the aldehyde substrates. The results were similar to PviCAD1, although the calculated K_m values for PviCAD2 were slightly lower. Both enzymes consistently exhibited higher activity on sinapaldehyde than on coniferaldehyde. A table of CAD kinetic parameters was collected from the published literature and is presented in Table 5. The results for PviCAD1 and PviCAD2 kinetics fall within the range of CAD kinetic data that were reported. PviAroADH was not analyzed because we could not obtain active soluble protein.

Fig. 2 Phylogenetic analysis of CAD sequences. This tree contains both putative and biochemically characterized CADs. The tree was constructed as indicated previously [43]. GenBank accession numbers are provided except for Arabidopsis, Chlamydomonas, Phsycomitrella, Selaginella, and Sorghum sequences; in these cases, numbers from their corresponding genome projects were used. Arabidopsis AtCAD1 (At1g72680), AtCAD2 (At2g21730), AtCAD3 (At2g21890), AtCAD4 (At3g19450), AtCAD5 (At4g34230), AtCAD6 (At4g37970), AtCAD7 (At4g37980), AtCAD8 (At4g37990), and AtCAD9 (At4g39330); C. reinhardtii CrCAD (CHLREDRAFT 190510); E. gunnii EgCAD1 (CAA61275) and EgCAD2 (CAA46585); L. perenne LpeCAD1 (AF010290.1) and LpCAD2 (AF472592.1); N. tabacum NtCAD19 (CAA44217); O. sativa FC1 (Os04g52280) and OsCAD2 (Os02g0187800); P. patens PpCAD (87951 scaffold 163:497997..49922); P. taeda PtCAD (CAA86073); P. tremuloides PotCAD (AAF43140) and PotSAD (AAK58693); P. trichocarpa PoptrCAD4 (estExt Genewise1 v1. C LG IX2359) and PoptrCAD10 (grail3.0004034803); S. officinarum (CAA13177); S. moellendorffii SmCAD (estExt fgenesh2 pg. C 390191); S. bicolor SbCAD1 (Sb06g001430.1), SbCAD3 (Sb02g024210.1), SbCAD4 (Sb02g024190.1), SbCAD5 (Sb07g006090.1), SbCAD6 (Sb06g028240.1), SbCAD7 (Sb02g024220.1), and Bmr6 (Sb04g005950.1); and Z. mays ZmCAD2 (BM1 locus; NM 001112184). The dashed line indicates CADs that are likely to be involved in monolignol biosynthesis and corresponds to the Class I CADs identified by Barakat et al. [5]

CAD Activity in Switchgrass Internodes

Native CAD activity in switchgrass internodes and leaves was analyzed in clarified tissue homogenates. Individual internodes were analyzed because they give insight into CAD activity across different phases of stem development. In all internodes there was significantly greater activity when sinapaldehyde was used as a substrate (Fig. 3), suggesting that the CAD or CADs present in these extracts shared similar substrate preferences. Leaf and sheath extracts showed little or no activity (data not shown). CAD activity was highest in extracts from internode 1 (3.43 nmol sinapaldehyde reduced per milligram of protein per minute) and lower in all other internodes.

CAD Protein Levels in Internodes

Immunoblot results of protein extracts from Kanlow N1 internodes are shown in Fig. 4 and revealed that immuno-



reactive bands at the expected size of CAD were present in all internode extracts at approximately similar levels.

Proteomic Identification of PviCAD1 and PviCAD2 and Other Proteins

Two-dimensional gel electrophoresis followed by immunoblotting and mass spectrometry of selected spots was performed using protein extracts from internode 3. The

Enzyme	Substrate	$K_{\rm m}~(\mu{\rm M})$	$V_{\rm max}~({\rm nkat}~{\rm mg}^{-1})$	$k_{\text{cat}} (\text{s}^{-1})$	$k_{\rm cat}/K_{\rm m} \; (\mu {\rm M}^{-1} \; {\rm s}^{-1})$
PviCAD1	Sinapyl alcohol	15.6±2.5	600±30	46.4	2.98
	Coumaryl alcohol	4.2±1.1	320±18	24.7	5.88
	Coniferaldehyde	10.9 ± 1.8	$1,280\pm60$	99.6	10.2
	Sinapaldehyde	14.5 ± 1.8	2,530±100	196	13.6
	Coniferyl alcohol	$0.98 {\pm} 0.1$	170±2.6	13.2	13.4
PviCAD2	Coniferaldehyde	$3.83{\pm}0.8$	$1,760\pm70$	138	35.8
	Sinapaldehyde	9.5±2.0	$2,500\pm140$	195	20.5

 Table 3
 Estimated kinetic constants



Fig. 3 CAD activity in switchgrass internodes. Internodes were numbered from least mature (top of plant; l) to most mature (bottom of plant; δ). Results are from duplicate experiments. *Error bars* represent the standard deviation (n=5 or 6)

immunoreactive region of the 2-D blots were used a guide to identify putative protein spots that could contain CAD proteins. A representative gel stained with Coomassie brilliant blue, a magnified region showing the numbered spots cut out for proteomic analysis, and an immunoblots are shown (Fig. 5a-c). Spots labeled 1 through 7 shown in Fig. 5b were excised and subjected to proteomic analysis. Proteomic analyses (Table 4) showed that peptides common to both PviCAD1 and PviCAD2 proteins were present in spots 1-4 and 7, although the number of peptides attributable to PviCAD1 and 2 were found in greatest abundance in spots 1 and 2, with a lesser number of peptides in spots 3, 4, and 7. Peptides unique to PviCAD2 were only found in spots 1 and 2. The predicted C-terminal residue for PviCAD1 (FVVDVAGSNIEEQAAAAGAPAN) was identified in spots 1-3. Taken together these data suggest that the abundance of PviCAD1-related peptides was probably greater than those attributable solely to



Fig. 4 Immunoblots of switchgrass internode extracts probed with CAD-specific polycolonal antibodies. Antibodies generated to ascorbate peroxidase (AscPx) were used a loading control. Pixel intensities for the signal arising from the two probed proteins are shown in the bottom bar graph



Fig. 5 a Two-dimensional gel of internode protein extract stained with Coomassie brilliant blue. The region of the gel selected for proteomic analyses is highlighted; **b** zoomed image of the 2-D gel with numbered protein spots that were analyzed by mass spectrometry; **c** An immunostained image from a comparable 2-D gel blotted onto nitrocellulose and developed with polyclonal antibodies to CAD is shown in the *bottom panel*. The signal arising from recombinant PviCAD1 is indicated as rCAD and was loaded in the marker lane to permit identification of potential spots that could contain native switchgrass CAD (see image above)

PviCAD2. Spots 5 and 6 were dominated by peptides arising from the large subunit of Rubisco (Table 4). Immunoblots of 2-D gels showed the presence of at least three or four immunoreactive spots at the expected size of CAD (\sim 42–45 kDa, Fig. 5c).

Real-Time qPCR Analysis of PviCAD1, PviCAD2, and PviAroADH Transcripts

Analysis of candidate reference genes using geNorm [50] indicated that UCE1, eIF4 α , and GAPDH would comprise a suitable set of reference genes. Using this approach, delta Ct values were normalized and relative expression levels of PviCAD1, PviCAD2, and PviAroADH in switchgrass internodes indicated that PviCAD1 transcripts were considerably more abundant in internodes as compared to the transcript levels for PviCAD2 (Fig. 6). Average PviCAD2 transcript levels were 44-fold lower than the PviCAD1 transcript levels in internode 1. Additionally, PviAroADH transcript was also consistently more abundant than PviCAD2. Compared to PviCAD1, relative

Table 4 Proteomic results

Protein	Sequence	Molecular weight	Ion score	Spot number
PviCAD1 and PviCAD2	TVVGWAAR	858.48	41	1,2
	HFGLTAPGLR	1,067.58	61	1,2,3,4
	GLTSQIEVVK	1,072.59	66	1,2,3
	ANVEQYCNK	1,124.5	42	1,2,3
	MGVYNEALER	1,180.55	83	1,2,3
	ANVEQYCNKR	1,280.59	38	1,2
	GGILGLGGVGHMGVK	1,350.75	66	1,2,3
	VGDVVGVGVIVGCCR	1,544.8	68	1,2,3,4
	FCVDKGLTSQIEVVK	1,721.9	1	1,2,3
	VLYCGICHTDIHQAK	1,813.86	52	1,2,3
	IWSYNDVYTDGRPTQGGFASTMVVDQK	3,050.46	76	1,2
	FVVPIPAGLAPEQAAPLLCAGVTVYSPLK	2,977.6	91	2,3
PviCAD1	NTGPEDVVVK	1,056.53	50	1,2,3,4
	ALGHHVTVISSSSR	1,449.77	71	1,2
	DAAGHLSPYTYTVR	1,549.75	74	1,2
	KSVTGSFIGSVDETAELLR	2,008.07	47	1,2,3,4,7
	FVVDVAGSNIEEQAAAAGAPAN	2,100.01	157	1,2,3
	HVLLGVVGEPLSFVSPMVMLGR	2,368.28	72	1,2,3
	YPMVPGHEVVGEVVEVGPEVSK	2,336.17	77	2,3,4
PviCAD2	KTGPEDVVVK	1,067.58	61	1,2
	AVTGSFIGSIDETAELLR	1,878.0	143	1,2
	HVLLGVVGEPLSFVAPMVMLGR	2,352.27	58	1,2
Ribulose 1,5,-bisphosphate	EITLGFVDLLR	1,274.72	78	5,6
	DDENVNSQPFMR	1,466.61	78	5,6
	DDFIEKDR	1,036.48	67	5,6
	VTPQPGVPPEEAGAAVAAESSTGTWTTVWTDGLTSLDR	3,583.86	101	5,6

expression values for *PviAroADH* ranged from 1.8- to 5fold lower. Expression patterns differed among the three transcripts. *PviCAD1* was most abundant in internode 1



Fig. 6 Relative expression levels of *PviCAD1*, *PviCAD2*, and *PviAroDH* in switchgrass internodes. Internode numbering as described for Fig. 3. *Error bars* represent the standard deviation

and displayed lower transcript abundance in all other internodes, with the lowest abundance in internode 4. *PviCAD2* transcript levels were low throughout the length of the tiller, with the highest abundance in internode 6. *PviAroADH* transcript levels were relatively constant across internodes 1–5 with the highest abundance in internode 6.

Detection of Motifs Present in CAD Enzymes with Good Catalytic Activity Against Hydroxycinnamylaldehydes

Based on the phylogenetic analyses, the AtCAD5 crystal structure [55], models of sorghum Bmr6 and a related sorghum CAD-like protein [43], and the enzyme activity data presented in Fig. 3 and Table 3, we looked for specific sequence motifs that could be used to distinguish CADs that are most likely to function in lignification versus CAD-like sequences that are unlikely to have significant role in secondary cell wall formation. Alignment of several grass CADs along with AtCAD8, SbCAD5, PviAroADH, and the putative *Z. mays* AroADH (enzymes not associated with

lignification, nor displaying significant in vitro activity against monolignals) indicated that there are several key conserved residues present in CADs with documented catalytic activity against coniferaldehyde or sinapaldehyde as compared to CAD-like proteins that display poor activity against these substrates (Fig. 7). The proposed mechanism for aldehyde reduction in AtCAD5 appears to involve Thr49, His52, and Asp57 [55]. Sattler et al. [43] had shown that the equivalent residue for Asp57 in monocot CADs was a histidine. As shown in Fig. 7, monocot and dicot CADs known to be associated with cell wall lignification and displaying significant catalytic activity against monolignals have the 57HL58 or 57DL58 motif. This motif is present in PviCAD1 and PviCAD2 but not in PviAroADH. Also, key residues identified in the substrate binding domain in AtCAD5 [55] and sorghum bmr6 [43] are apparently absolutely conserved in all other CADs with a demonstrated role in cell wall lignification; these include the 118IWSY121 and 274LGV276 or 274MGV276 motifs (Fig. 7). The IWSY and LGV motifs are present in PviCAD1 and 2 but absent in PviAroADH and related proteins. In these enzymes, two critical amino acid substitutions are evident, first, the leucine next to the catalytically important His57 or Asp57 has been replaced with tryptophan, and second, the tryptophan in the IWSY motif has been replaced with phenylalanine.

PviCAD1 and PviAroADH Protein Structures

PviCAD1 and PviAroADH structures were modeled using the AtCAD5 [55] structure in order to minimize template bias in the modeling. Amino acids 6 to 356 of PviCAD1 show 71.5% sequence identity with the AtCAD5 sequence, whereas amino acids 15 to 359 of PviAroADH align with only 48.1% sequence identity to AtCAD5. The residues thought to be catalytically important appeared to have the same general conformation between PviCAD1 and AtCAD5 (Fig. 8a and b). The overlay (Fig. 8c) further shows the basic similarities and differences between these two CADs; most residues have a similar orientation. However, the orientation of the catalytically important His57 is apparently different than the Asp57 of AtCAD5. Similarly, Trp119 and Tvr121 in PviCAD1 are oriented in a manner that could improve substrate binding. Results for PviAroADH (Fig. 8d) showed that, when AtCAD5 coordinates were used as a template, the alignment produced a structure that was markedly different from the active site of either AtCAD5 or PviCAD1. Significantly, the Trp64 next to the predicted catalytically important Glu63 residue appears to protrude into the putative substrate binding pocket and the replacement of Trp119 in the CADs with high catalysis against cinnamylaldehydes with Phe125 in PviAroADH appears to also change the putative substrate binding pocket relative to PviCAD1, suggesting that PviAroADH is likely to prefer substrates other than cinnamylaldehydes.

Discussion

PviCAD1 and PviCAD2 encode for the dominant CADs involved in switchgrass lignification

CAD is involved in the phenylpropanoid pathway in plants and catalyzes the final step in monolignol biosynthesis by converting cinnamyl aldehydes to the corresponding cinnamyl alcohols [28]. Here, two CADs and one CAD-like gene were cloned from switchgrass EST resources. Although *PviCAD1* and *PviCAD2* were cloned from different cultivars, analysis of existing Kanlow CAD clones and

```
н
                           GASKYPMVPGHE---IWSY---LG
PviCAD1
        49 T D I H Q A K N
                          L
                                                              276
                                                             v
                        нL
          TDIHQAKN
                           GASKYPMVPGHE----IWSY----LG
PviCAD2
        49
                                                             v
                                                               276
          TDIHQAKN
                        нL
                           GASKYPMVPGHE----IWSY----LG
                                                               276
BMR6
        49
                                                             V
          TDIHQAKN
                        нL
                           GASKYPMVPGHE----IWSY----LG
ZmCAD2
        49
                                                             v
                                                              276
          тогно
                   TKN
                        ΗL
                           GASKYPMVPGHE----IWSY----MG
LpeCad1
        49
                                                             v
                                                              277
          тогно
                   т
                    ĸΝ
                        D
                         L
                                  YPMVPGHE---IWSY---MG
AtCAD5
        49
                           GMSN
                                                             v
                                                              276
PoptrCAD4 49 T D I H Q I K N
                        DL
                           G M S H Y P M V P G H E---I W S Y---M G
                                                             v
                                                              276
ZmMDH
        52 S D L H S I K N
                        ΕW
                           HNAT
                                  ҮРМVРGНЕ---I FТY---VG
                                                             L 279
                           NNAMYPMVPGHE---IFTY---VGL282
PviAroDH
        55 S D L H F I K N
                        ΕW
        61 S D L H S I K N
                        ΕW
                           G
                             NAMYPMVPGHE---IFTY---VGL290
SbCAD5
AtCAD8
        48 S D L H M V K N E W
                           G
                             мзт
                                  Y P L V P G H E---I Q T Y---V G A 275
                          W
PtSAD
        52 S D L H S I K N D
                           G F S M Y P L V P G H E---I L T Y---V G A 279
        52 S D L H S I K N D W
                           GFSMYPLVPGHE---ILTY---VGA279
Poptr10
```



and *Arabidopsis* (AtCAD5 and AtCAD8). Alignment was done using ClustalW2. The amino acids shown include those in the active site as proposed by Youn et al. [55]

Fig. 8 Comparison of key residues in the predicted active site of AtCAD5 (a), PviCAD1 (b), and PviAroDH (d). Overlay of AtCAD5 (*pink/yellow residues*) and PviCAD1 (*green/ red residues*) is shown in c. PviCAD1 and PviAroDH structures were based off of the AtCAD5 crystal structure. Only residues that are likely to be involved in substrate docking or the catalytic reaction are shown



proteomic data, especially identification of the PviCAD1 Cterminal peptide indicated that PviCAD1 sequence was identical in both cultivars. Despite the fact that many genes coding for CADs or CAD-like proteins are found in all currently known plant genomes (for example: [5, 22]), genetic and enzyme activity data from several studies support the notion that one or at most a few CADs actually function in cell wall lignification [14, 38, 43, 46, 56]. The data presented here indicate that a similar situation exists in switchgrass, where *PviCAD1* and *PviCAD2* encode functional CAD proteins that appear to be involved in switchgrass lignin biosynthesis.

CAD activity and protein was detected in all internodes, which indicated that CAD is active in lignifying tissues at different stages of maturity. These data were consistent with earlier studies that had shown lignification is greater in more mature internodes suggesting that cell wall lignification continues to take place across all stages of internode maturity. Our studies focused on the stems since these tissues comprise a significant proportion of biomass at harvest, and represent tissues with the most lignin content. Ultimately, improving quality parameters in switchgrass biomass for liquid fuels will probably result in significant reduction in stem lignin content [39].

Immunoblot results from 2-D gel electrophoresis indicated that different forms of CAD may exist in switchgrass internodes, although these differences were not fully identified. Multiple spots can be artifacts due to processes such as deamidation which are frequently encountered during sample preparation for 2-D gel analyses [41] and it is unclear if differences included CAD isozymes, posttranslational modifications, or were simply artifacts. However, our polyclonal antibodies will detect other related CAD-like proteins in tissue extracts [43]. Proteomic analyses revealed a greater abundance of peptides matching PviCAD1 suggesting that PviCAD1 was present at higher levels or peptides derived with PviCAD1 were preferentially enriched during proteomic analyses. Enrichment of PviCAD1 relative to PviCAD2 was supported by qPCR data that showed greater transcript abundance for mRNA coding for PviCAD1 relative to mRNA coding for PviCAD2. However, these results do not preclude the possibility that PviCAD2 may be present in specific tissues or have other temporally or spatially defined developmental roles as was suggested for CADs in aspen [27] and other species [5, 24]. Kanlow switchgrass is an allotetraploid cultivar [1, 33] and it is possible that PviCAD1 and PviCAD2 are encoded by the two different genomes. In many polyploid species, translation of alleles is not equal, and dominance by one or the other genome is observed [1, 13]. Our data suggests that PviCAD1 and PviCAD2 transcripts are present in all internodes. Furthermore, these tissues appear to contain active CAD proteins which could participate in the continued lignification of tissues such as the stem sclerenchyma and parenchyma. Previously published results show that these tissues are progressively more lignified in the basal and more mature internodes as compared to the topmost internode [39]. Future functional genomic studies in switchgrass should allow greater insights into these phenomena.

Relationships of PviCAD1, PviCAD2, and PviAroADH to Other CAD and CAD-Like Proteins

PviCAD1 and PviCAD2 displayed a preference for aldehyde substrates, which was unsurprising because these enzymes are expected to catalyze the final step in monolignol biosynthesis where cinnamaldehydes are converted to their corresponding alcohols. Although PviCAD1 velocity and turnover (k_{cat}) values were higher for the aldehyde substrates, the two lowest K_m values were for coniferyl and coumaryl alcohol, respectively. The low PviCAD1 K_m observed with coniferyl alcohol results in a higher k_{cat}/K_m value when compared to the other alcohols. Otherwise, PviCAD1 efficiency with the aldehydes was higher than with alcohols. PviCAD2 was

even more efficient with aldehyde substrates than PviCAD1, although the exact localization and temporal control of these two enzymes are not currently known. A similar preference for aldehyde substrates, particularly sinapaldehyde has been reported for sorghum and rice CADs [43, 56]. However, reported kinetic data for other CADs did not necessarily show this type of strong preference (Table 5). In *Arabidopsis*, AtCAD5 had only a 12.4% higher V_{max} for sinapaldehyde [22]; in *Eucalyptus gunnii* the velocity increase was 31–33%, depending upon the CAD subunit composition [19]. In contrast, CAD velocity on sinapaldehyde was markedly lower in *Populus euramericana* [42], *Picea abies* [16, 29], *Populus tremoloides* [27], and *Pinus thunbergii* [25]. Outside of monocots, the only other kinetically

AD kinetic	Species	Substrate	$K_{\rm m}~(\mu{\rm M})$	$V_{\rm max}~({\rm nkat}~{\rm mg}^{-1})$	Reference
	Pinus thunbergii	Coniferaldehyde Sinapaldehyde	9.1	3.3 0.073	[25]
		<i>p</i> -coumaraldehyde	30	5.7	
		Cinnamaldehyde	14	7.6	
	Picea abies	Coniferaldehyde Sinapaldehyde	3.6 83	1,724 167	[29]
		4-coumaraldehyde	12.5	2,857	
	Forsythia suspensa	Cinnamyl alcohol <i>p</i> -coumaryl alcohol	156 132	39% ^e 112%	[31]
		Coniferyl alcohol	32	100%	
	Eucalyptus gunnii	Coniferaldehyde Sinapaldehyde	4.5 ^a /5.2 ^b 6.8/2.5	2,104 ^a /1,580 ^b 2,758/2,107	[19]
		<i>p</i> -coumaraldehyde	5.1/16	1,244/2,107	
		Coniferyl alcohol	2.3/23	539/1,817	
		Sinapyl alcohol	6.6/4.5	717/562	
		p-coumaryl alcohol	35/64	812/1,150	
	Populus euramericana	Coniferaldehyde <i>p</i> -coumaraldehyde	0.77 1.2	52.2 17	[42]
		Sinapaldehyde	4.8	19.5	
	Populus tremoloides	Coniferaldehyde Sinapaldehyde	2.3° 9.1	3.5 ^c 1.7	[27]
		<i>p</i> -coumaraldehyde	6.2	2.8	
		Caffealdehyde	37	2.5	
		5-OH coniferaldehyde	17.5	2.8	
	Arabidopsis thaliana (AtCAD5) ^d	Coniferaldehyde Sinapaldehyde	35 20	157.4 177	[22]
		<i>p</i> -coumaraldehyde	13	187.3	
		Caffeylaldehyde	68	94.1	
		5-OH coniferaldehyde	22	106.9	
	Medicago sativa ^d	Coniferaldehyde Sinapaldehyde	1.3 6.9	47.8% ^e 67.2%	[9]
		Cinnamaldehyde	9.2	100%	
	Oryza sativa ^d	Coniferaldehyde Sinapaldehyde	4.4 20.8	32.9 87.0	[56]
	Lolium perenne L.	Coniferaldehyde Sinapaldehyde	1.9 6.3	57.8% ^e 75.7%	[32]
		Cinnamaldehyde	5.8	100%	

Table 5Reported CAD kinetparameters

^a HeteroCAD ^b HMW HomoCAD

^d Recombinant protein ^e Only relative values were reported

^c PtCAD

characterized enzyme for which a relatively similar increase in activity was observed is PtSAD [27], although questions have been raised about its proposed specific physiological role in angiosperms [2].

A recent analysis of Populus trichocarpa CAD genes suggested that the CAD protein sequences present in the Populus genome could be separated into three classes [5]. Based on these classes. PviCAD1 and PviCAD2 fall into Class I, which includes PoptrCAD4 and AtCAD5 and further indicates the broad conservation of the ligninpathway related CADs in plants. The phylogenetic analysis reported here places PviCAD1 and PviCAD2 into a distinct clade that contains all of the other known wellcharacterized CADs that have a role in catalyzing the last step in monolignol biosynthesis and are thus bona fide CAD genes. Also, this group further divides into two major sub-groups encompassing the monocots and dicots; Pvi-CAD1 and PviCAD2 fall into a group that contains all of the other known monocot CADs, including the recently reported sorghum BMR6 [43]. However, PviAroADH, PtSAD, and its apparent Populus ortholog Poptr10 are not included within this clade which is in agreement with Barakat et al. [5], and suggests these genes may not be involved in lignin monomer synthesis or may have other, more primary, roles such as plant defense or catalytic activity under specific conditions [8, 46]. We have been unable to produce soluble, active recombinant PviAroADH in E. coli and could not directly characterize this protein. A sorghum ortholog (Sb 02g024190; 61% identity, 5e-116) of PviAroADH displayed poor activity against monolignol and monolignal substrates [43], suggesting that Pvi-AroADH could exhibit similar properties. Future biochemical evaluation of recombinant PviAroADH should clarify these points and indicate if it could participate in the lignification process.

Sequence Motifs of Lignifying CADs and Predicted Model of PviCAD1

Sequence alignment of binding pocket and active site residues based on AtCAD5 [55], showed that among lignifying CADs (i.e., CADs shown by activity and/or genetics to function in the biosynthesis of monolignols), most amino acids were highly conserved, with the exception that in monocots His57 is present instead of the dicot Asp57 [43].

When the CAD-like sequences including PviAroADH, PtSAD, and AtCAD8 were compared to bona fide CADs, some differences were evident. First, PviCAD1 and other Class I (lignin) CADs have an invariant Gln53, which is more variable at the equivalent position in other CAD-like sequences. Second, the lignin CADs have either a His57/ Leu58 or Asp57/Leu58 motif, while the CAD-like sequences contain either Glu/Trp or Asp/Trp at equivalent positions. Finally, lignin CADs contain an invariant Trp119 which is not conserved in the other CADs. In examining the binding pocket models for PviCAD1, AtCAD5, and PviAroADH, the differences are more apparent (Fig. 8a-d). Trp119, which resides towards the back of the binding pocket, likely stabilizes the aromatic ring of cinnamaldehydes through pibonding: this residue has a similar conformation in both PviCAD1 and AtCAD5 (Fig. 8a and b, respectively). Additionally, Youn et al. [55] noted that Trp119 and Phe299 shrunk the size of the substrate binding pocket compared to PtSAD (Leu122/Gly302), which likely resulted in AtCAD5 having increased substrate specificity. These residues (Trp119/Phe299) are replaced by Phe125 and Cys305 in PviAroADH. Also, His57 and Leu58 in Pvi-CAD1 are replaced by Glu63 and Trp64 in PviAroADH. Taken together, these amino acid changes could dramatically alter the binding pocket conformation of PviAroADH (Fig. 8d) and, although a member of the alcohol dehydrogenase superfamily, this protein is likely to have a biological role that is different than PviCAD1 in switchgrass.

Given that switchgrass is a good candidate feedstock for biorefineries [36, 40] and that CAD mutants have been shown to impact lignin composition and conversion to ethanol in sorghum [15, 34, 43], the data presented here will be useful for understanding and manipulating lignin synthesis for future development of switchgrass bioenergy cultivars.

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