Involvement of a cinnamyl alcohol dehydrogenase of *Quercus suber* in the defence response to infection by *Phytophthora cinnamomi*

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Abstract

A gene encoding a potential NADPH-dependent cinnamyl alcohol dehydrogenase (QsCAD1) (GenBank accession no: AY362455) was identified in *Quercus suber* (cork oak). Its complete cDNA sequence was obtained by RACE-PCR, starting from total RNA extracted from roots of seedlings of *Q. suber*, infected with *Phytophthora cinnamomi*, the causal agent of the decline and sudden death of *Q. suber* and *Quercus ilex* subsp. *rotundifolia* in the Iberian Peninsula. Sequence information to perform the RACE-PCR was acquired from a polymorphic fragment (C9), specifically identified by cDNA-AFLP, in leaves of epicormic shoots of a cork oak tree that suffered sudden death. RT-PCR and hybridization analysis showed that the QsCAD1 gene is up-regulated in root seedlings of *Q. suber* infected with *P. cinnamomi*. QsCAD1 has a high structural homology with VR-ERE (*Vigna radiata*), an enzyme that detoxifies eutypine (produced by *Eutypa lata*, the causal agent of eutypa dieback of grapevines), to eutypinol, and with QrCAD1 (*Q. ilex* subsp. *rotundifolia*), EgCAD1 (*Eucalyptus gunnii*), MdCAD1 (*Malus x domestica*). Taken together, these results suggest that these enzymes, and namely QsCAD1 belong to a new group of CAD potentially involved in deactivation of toxins produced by phytopathogens.

Keywords: Oak tree; *Phytophthora cinnamomi*; Cinnamyl alcohol dehydrogenase; Defence response

1. Introduction

The involvement of *Phytophthora cinnamomi* in the decline disease of cork oak (*Quercus suber*) and holm oak (*Quercus ilex* subsp. *rotundifolia*) in the Iberian Peninsula was hypothesized 10 years ago [1,2]. Recent extensive field surveys and greenhouse experiments provided evidence that *P. cinnamomi* infects the roots of these evergreen oaks. The reduction of the amount of roots available for water and nutrient uptake, caused by *Phytophthora* root rot, leads to widespread deaths of cork oak and holm oak, either directly, or by predisposing the trees to other biotic and abiotic stresses, factors usually nondeleterious to healthy plants [1,3,4]. In affected areas, some cork oaks exhibit a range of symptoms and a variable rate of disease expression, while others die suddenly. Those that show evidence of chronic disease show a gradual deterioration of the crown that starts with leaf chlorosis and dieback of leaf bearing branches. As disease progresses, anomalous leaf fall occurs. New leaves, formed on apparently dead shoots, have reduced size, resulting in thinning of the crown and reduction of leaf area. Some affected trees show stem fluxes indicative of root malfunction. Trees showing such symptoms may survive for several years, depending on their resistance to disease progression and adaptation to environmental conditions. In contrast to chronic disease decline, the sudden death process is characterized by a quick drying of the leaves and death of the tree, within few months. The causes of these contrasting behaviours, slow decline and sudden death, are not known. Until now, no efficient methods have been found that will limit the threat of the disease to the oak ecosystem.
Toxins produced by some fungi and bacteria play an important role in the expression of disease in plants. Toxin tolerance or resistance has been often associated with the capacity of the plant to metabolically detoxify the toxin [5–7]. As association between toxin tolerance and resistance to the disease has been frequently described [8,9], it is important to explain the mechanism underlying detoxification, and characterize the enzymes involved in this process.

A novel NADPH-dependent aldehyde reductase gene (VR-ERE) from Vigna radiata, that confers resistance to the grapevine fungal toxin eutypine [4-hydroxy-3-(3-methyl-3-buten-1-ynyl) benzaldehyde], was recently described [10]. Eutypine is produced by the mycelium of Eutypa lata, the causal agent of eutyp dieback of grapevines [11–14]. It induces a number of symptoms, including dwarfed and withered shoots, marginal necrosis of the leaves, desiccation of the inflorescences and death of branches. It was suggested [15] that eutypine is transported from the trunk through the sap to the herbaceous parts of the vine, where it exerts its toxic action. The recombinant form of VR-ERE was shown to have a high affinity for 3-substituted benzaldehydes, reducing them to alcohols. Its reducing activity was namely confirmed with eutypine, that it converts into the nontoxic corresponding benzyl alcohol, eutypinol [16]. The tolerance of some cultivars to this disease was correlated with their capacity to achieve this conversion.

The aim of the present work is to disclose cork oak genes involved in the defense response to infection by P. cinnamommi, namely in connection with the sudden death disease expression. As a first step in attempting this objective, we looked for polymorphic fragments present in mRNA profiles of leaves from adult asymptomatic trees and from trees that either exhibited symptoms of slow decline or underwent sudden death. In a second step, we have looked for the corresponding genes in roots of cork oak seedlings. The outcome of this approach was the identification of four genes, namely one that encodes a cinnamyl alcohol dehydrogenase (CAD), highly homologous to VR-ERE. The expression of CAD was analysed in infected and noninfected roots of Q. suber seedlings. The potential involvement of the uncovered CAD in the detoxification of a putative toxin produced by P. cinnamommi is discussed.

2. Material and methods

2.1. cDNA-AFLP analysis

2.1.1. Plant material

Fully expanded young leaves were collected in May, from five adult cork oak trees (BS222A-4, BS250HL-0, BS110HL-3, AS17BV-0 and AS21BV-3) located in infested sites from decline areas of Alentejo and Algarve regions of Portugal, and immediately stored at −80°C. P. cinnamommi had previously been isolated from roots and associated soil of oak trees located in those sites [3]. Trees were selected on the basis of the defoliation degree (DG), ranging from 0 (healthy tree) to 4 (dead tree), according to the scale proposed by Cadahia et al. [17]. The last figure, included in the designation of the trees, corresponds to the DG attributed at the moment of the collection of the leaves. Leaves from a cork oak tree affected by sudden death (BS222A-4) were collected from epicormic shoots present in the trunk.

2.1.2. Extraction and purification of total RNA

Total RNA was extracted from 50 mg of leaves with the RNeasy Kit from Qiagen, according to the instructions supplied by the manufacturer [18]. Total RNA was then treated with DNase I (1 U/µL; Gibco), in the presence of 2 µL of RNaseout (40 U/µL; Gibco) [19]. The reaction occurred in a final volume of 100 µL, for 30 min at 37°C. After DNA digestion, the total RNA was purified with the RNeasy Kit. The quality and the quantity of total RNA present in the samples were evaluated by UV spectrophotometry and by electrophoresis in denaturing agarose gel.

2.1.3. cDNA synthesis

The synthesis of cDNA was accomplished with the cDNA synthesis system (Roche), with modifications to the original protocol. Synthesis of the first cDNA strand was initiated with the degenerate primer [5′AGTGAAATT CT12V (V = A; C; G), comprised of an equimixturm of the three oligonucleotides [20]. Synthesis of the second cDNA strand and digestion of residual RNA were performed according to the kit protocol [21,22]. The double strand cDNA was then purified with the Qiaquick PCR purification kit (Qiagen). This methodology is designed to purify double-stranded DNA fragments from enzymatic reaction mixtures. Fragments ranging from 100 bp to 10 kb were separated from polymerases, RNases, ligases, nucleotides and salts. In the process DNA binds to the QIAquick column (silica-gel membrane), and proteins or compounds resulting from the enzymatic reactions are eliminated by washing the column with buffers. In the last step of the protocol, the DNA is eluted from the column with 50 µL of the elution buffer (10 mM Tris–Cl, pH 8.5). The volume of cDNA samples, eluted from the column, was reduced to a final volume of 20 µL by centrifugation under vacuum. cDNA samples (20 µL) were then subjected to the standard AFLP template production [23]. The cDNA was first digested with EcoRI and MseI (2.5 U/µL each) in a restriction buffer (50 mM Tris–HCl pH—7.5, 50 mM magnesium acetate, 250 mM potassium acetate), in a total volume of 25 µL for 2 h at 37°C (AFLP Core Reagent Kit, GIBCO BRL). After enzyme inactivation at 70°C for 15 min, EcoRI (5 pmol) and MseI (50 pmol) adaptors were ligated to cDNA digested fragments. The reaction was performed for 2 h at 37°C in ligase buffer, 1 U of T4 DNA ligase and ATP (0.4 mM) (AFLP Core Reagent Kit, GIBCO BRL). The cDNA ligated to the adaptor was pre-amplified using the following cycling parameters: 28 cycles of 30 s at 94°C; 60 s at 56°C and 60 s
at 72°C. The pre-amplification reaction was carried out with 15 pmol of primer EcoRI + 0, 15 pmol of primer Msel + 0.5 µL of cDNA template, 0.2 µL of Taq DNA polymerase in an appropriate Taq buffer. Aliquots of the pre-amplification reaction were run on a 1.5% agarose gel, to verify the presence of a smear of cDNA, before proceeding. The pre-amplified cDNA was diluted in water, in a ratio of 1:10 and was used as a template for the selective proceeding. The pre-amplified cDNA was diluted in water, to verify the presence of a smear of cDNA, before pre-amplification reaction were run on a 1.5% agarose gel, polymerase in an appropriate Taq buffer. Aliquots of the 15 pmol of primer Eco at 72°C. of 30 s at 94°C, cycling parameters were applied: 13 cycles with denaturation of 30s at 94°C, primer annealing of 30 s at 65°C. The annealing temperature was lowered by 0.7°C/cycle during the first 12 cycles, until reaching 56°C. Polymerization of 60s at 72°C, and then 18 cycles of 30 s at 94°C, 30 s at 56°C and 60 s at 72°C. Labelled selective amplification products were separated on standard 6% polyacrylamide sequencing gels. After electrophoresis, the gel was dried on filter paper (3MM paper; Whatmann) and exposed to X-ray film for 30h. The cDNA fragments were visualised by autoradiography, after positional marking the gel and the film.

2.2. Isolation and sequencing of fragments

Polymorphic gene fragments, present only in the mRNA profile of cork oak tree affected by sudden death, were excised from the gel. A piece of the dried gel, containing the band of interest, was cut out and soaked in 40 µL of H2O for 10 min on ice. The sample was then heated for 15 min at 95°C and cooled again on ice. After a brief centrifugation (30 s, 13,000 rpm), 5 µL of the supernatant were transferred to another tube. Re-amplification of the recovered fragment was performed under the same conditions, and with the same primer combination used in the reaction which generated the product of interest. The re-amplified PCR product was run on a 2% agarose gel, cut out and purified with the Qiaquick PCR Purification Kit (Qiagen) and finally cloned into the pCRII Topo vector, with the TA Cloning Kit (Invitrogen). Manufacturer’s instructions for these kits were followed throughout. The fragments were sequenced with an Applied Biosystems PRISM Ready Reaction DyeDeoxy Terminator cycle sequencing kit and an automated sequencer.

2.3. Rapid amplification of 5’ and 3’ complementary cDNA ends (RACE-PCR)

2.3.1. Plant material

Seeds of cork oak and holm oak were germinated at room temperature (25°C) in a mixture of vermiculite and sand. When the main root reached 8–10 cm, the seedlings were separated from the sand, washed with water and transferred to Petri dishes to be infected with P. cinnamomi. Roots still attached to the seeds, were inoculated with slices of agar containing P. cinnamomi (isolate PA20) mycelium grown for 1 week in V8 solid medium [26]. Roots in contact with slices of mycelium were covered with aluminium foil, and placed at 25°C for 24 h. The infected roots were then cut and stored at −80°C.

2.3.2. Extraction and purification of total RNA

Total RNA was extracted from 50 mg of roots of cork oak seedlings with RNeasy Kit from Qiagen, according to the instructions supplied by the manufacturer [18]. DNase treatment of total RNA was performed as described above (cDNA-AFLP analysis).

2.3.3. RACE-PCR

RACE-PCR (3’/5’) was performed to obtain full length cDNA of the gene comprising the C9 cDNA-AFLP fragment. Three and five prime ends were obtained with a GeneRacer Kit (Invitrogen) following the manufacturer’s instructions [27,28].

DNase treated total RNA (1 µg) extracted from roots of cork oak seedlings, infected with P. cinnamomi, was used as template in the following steps:

(1) 5’ phosphates were removed from truncated mRNA and from non mRNA with calf intestinal phosphatase (CIP). The reaction was performed for 1 h at 50°C, in a total volume of 15 µL, comprised of 1 µL CIP (10 U/µL), 1 µg of total RNA, 1 µL of RNase inhibitor (10 U/µL), 1 µL of CIP buffer (0.5 M Tris–HCl, pH 8.5; 1 mM EDTA). RNA was precipitated and washed with ethanol and resuspended in 7 µL DEPC water as described in the instruction manual of GeneRacer™ Kit (Invitrogen).

(2) Dephosphorylated RNA (6 µL) was treated with tobacco acid pyrophosphatase (TAP) to remove 5’ cap structure from intact, full-length RNA. The reaction was carried out for 1 h at 37°C in a total volume of 10 µL, comprised of 1 µL of TAP (0.5 U/µL), 1 µL of TAP buffer (0.5 M sodium acetate, pH 6; 10 mM EDTA; 1% β-mercaptoethanol; 0.1% TritonX-100), 1 µL of RNase inhibitor (10 U/µL). RNA was precipitated and washed with ethanol and resuspended in 6 µL DEPC water, as described in the instruction manual of GeneRacer™ Kit (Invitrogen).

(3) For selective ligation of an RNA oligonucleotide (RNA Oligo) to the 5’ ends of decapped mRNAs 6 µL of dephosphorylated decapped RNA were added to 0.25 µg of lyophilized RNA Oligo (5’ CGACUGGA GCACGAGGACACUGACACUGAAGGAG UAGAAA3’), the mixture was incubated for 5 min at 65°C and chilled on ice for 2 min. After a brief centrifugation (5 s, 13,000 rpm), 1 µL of RNase Inhibitor (10 U/µL), 1 µL of ligase buffer (330 mM Tris-acetate, pH 7.8; 660 mM potassium acetate, 100 mM magnesium acetate; 5 mM DTT), 1 µL of ATP (10 mM), 1 µL of T4 RNA ligase (5 U/µL) were added. The reaction mixture was incubated for 1 h at 37°C. RNA was precipitated and washed with ethanol and
resuspended in 13 μL DEPC water, as described in the instruction manual of GeneRacer™ Kit (Invitrogen).

(4) The synthesis of cDNA was primed by an oligonucleotide composed of 18 thymidine and 36 additional nucleotides (5’GCTGTCAACGATAACGTACGTAA-CGGCATGACAGTG183') (oligo-dT). Superscript™ II RNase H⁻ (GIBCO) was used as reverse transcriptase. One microlitre of oligo-dT (37.5 μM), 2 μL dNTPs (10 mM) and water were added to total RNA, to a final volume of 31 μL. The reaction mixture was incubated at 65°C for 5 min and then chilled on ice for 2 min. Then, 4 μL of First Strand Buffer (250 mM Tris–HCl, pH 8.3; 375 mM KCl; 15 mM MgCl₂, 2 μL 0.1 M of DTT, 1 μL RNaseOUT (40 U/μL), 2 μL of Superscript (200 U/μL) were added to the reaction mixture that was further incubated at 42°C for 1 h. The enzyme was inactivated at 70°C for 15 min, then 1 μL (2U) of RNase H was added and the reaction was performed at 37°C for 20 min.

The obtained single strand cDNA was used as template for the amplification of 3' and 5' ends.

Amplification of the 3' end was carried out with GeneRacer 3' primer (5'GCTGTCAACGATAACGTACGTAA-CGGCATGACAGTG183') that anneals with oligo-dT and with a primer, specific to the gene comprising the C9 fragment (C9SP2; 5' CAAAGCAAATCTACTGGAAG3'). Amplification was performed in a total volume of 50 μL, comprised of 5 μL of PCR buffer (200 mM Tris–HCl, pH 7.5, 1 M KCl; 15 mM MgCl₂, 10 mM DTT, 1 mM EDTA, 1% v/v Tween 20, 1% v/v Nonidet P40), 5 μL of dNTPs (2 mM each), 2 μL of primers (10 mM), 1 μL of cDNA solution, 0.75 μL of Expand High Fidelity Polymerase (3.5 U/μL). The following cycling parameters were applied: 2 min at 95°C; 30 cycles with denaturation at 95°C for 30 s, primer annealing at 50°C for 30 s, polymerisation at 72°C for 60 s; one cycle at 72°C for 7 min.

Amplification of the 5' end was carried out with GeneRacer 5' primer (5'CGACTGGGACGCAGGAGCACTG183') that anneals with the DNA complementary to RNA Oligo and with a primer, specific to the gene comprising the C9 fragment (C9SP2; 5'TCACAGCCCTC AACAGCAGA3'). Amplifications conditions were identical to those above described for the 3' end amplification.

The products of the amplification reactions were cloned into the pCR II Topo vector, with the TA Cloning Kit (Invitrogen), and sequenced.

2.4. Qualitative analysis by RT-PCR and hybridization

2.4.1. Plant material

Seeds of cork oak and holm oak were germinated at room temperature (25°C) in a mixture of vermiculite and sand. When the main root reached 8–10 cm, the seedlings were separated from the sand and washed with water. Some roots were cut and stored at −80°C and others were transferred to Petri dishes, to be infected with *P. cinnamomi*. The infection of the root seedlings was performed as described above.

2.4.2. RT-PCR and hybridization

Total RNA (2 μg) was extracted from noninfected roots of cork oak seedlings, and from roots that were put in contact with *P. cinnamomi* mycelium for 24 h. cDNA was synthesized with Superscript II RNase H-reverse transcriptase, in a total volume of 40 μL. Two microlitres of the cDNA synthesis solution were diluted 50, 100 and 200 times, and 10 μL of these dilutions were used as cDNA template for amplification reactions. Fragments greater than 1000 bp from *QsCAD1* transcripts were amplified with a specific primer, C9SP4 for the known sequence and an oligo-dT primer. The following primers were used for the amplification of the fragment: 5'GCCGTTGACGAGCCCCCAGCA3' and 5'CAACGCTCGGTAGGTACTCG3'. PCR products were separated on 1% agarose gels and blotted to membrane (Hybond-N +) on a Trans-Blot SD semi-dry electrophoretic transfer cell. The specificity of the amplified products was confirmed by hybridization. A DNA fragment (75 bp) from *QsCAD1* gene, obtained by amplification with C9SP2 and C9SP4 primers, labelled with digoxigenin, was used as probe. Hybridization of digoxigenin-labelled probes, with the target DNA, was achieved at 42°C in the presence of DIG Easy Hybridization buffer (Roche). Digoxigenin-labelled nucleic acid hybrids were detected with a Dig Luminescent Detection Kit (Roche).

2.5. Phylogenetic analysis

The amino acid sequences of CAD 1 and 2, identified in *Q. suber*, were aligned with those of their homologues in plants by the CLUSTAL W algorithm [29]. Phylogenetic analysis was conducted with the neighbour-joining method [30].

2.6. Sequence analysis

The cDNA sequences from RACE-PCR and cDNA-AFLP analysis were aligned and compared through the National Centre for Biotechnology Information (NCBI) GenBank (http://www.ncbi.nlm.nih.gov/BLAST/) with the basic local alignment search tool (BLAST) algorithm [31], the fastA programme from the GCG software (Genetics Computer Group, University of Wisconsin, Madison, 1981) and the Vector NTI 6 software™ (InforMax, Inc).

3. Results

3.1. Identification of a gene from *Q. suber* encoding a potential cinnamyl alcohol dehydrogenase

cDNA-AFLP methodology was used to identify polymorphisms in genes that are expressed in leaves of cork oak trees presenting symptoms of the decline disease, manifested as slow decline or sudden death [36]. The severity of
the disease was evaluated according to different DGs based on the scale proposed by Cadahia et al. [17]. The pattern of mRNAs extracted from young leaves of cork oak trees exhibiting various DGs, and growing in P. cinnamomi infested soil, showed a high polymorphism. Twenty-seven out of 122 up-regulated cDNA-AFLP fragments were cloned and sequenced (not shown). One was shown to have a high homology with CAD1 gene sequence and used to characterize this gene in Q. suber as described below. The mRNA was extracted from young leaves of five cork oak trees: BS222A-4, BS250HL-0, BS110HL-3, AS17BV-0 and AS21BV-3, with DGs of 4, 0, 3, 0 and 3, respectively, located in decline areas where P. cinnamomi had already been isolated with success from roots and associated soils.

In the mRNA profile obtained with primer combination I2 (EcoRI-ACC; Msel-ACC), a polymorphic fragment with 200 bp (named C9) was identified. This fragment was exclusively present in the cork oak tree BS222A-4 that was affected by sudden death (Fig. 1). The mRNA, used for the cDNA-AFLP analysis, was extracted from leaves of epicormic shoots present in the trunk. The comparison of C9 fragment nucleotide sequence with GenBank data, showed similarities of 85% and 77% with sequences of genes that code for CADs identified in Malus x domestica1 and in Eucalyptus gunnii,2 respectively. The complete cDNA sequence of the gene corresponding to the C9 fragment was obtained by RACE-PCR, starting from total RNA extracted from roots of seedlings of Q. suber infected with P. cinnamomi. The amplification was performed with primers defined in regions of the C9 fragment that are conserved in respect to the above referred to CADs. The resulting cDNA was found to be 1399 bp long. It is designated henceforth as QsCAD1. The open reading frame (978 bp) encoded a protein of 326 amino acids with a calculated molecular mass of 35 922 Da, and a calculated isoelectric point of 7.72. The 5′-UTR contained 54 nucleotides and the 3′-UTR contained 361 nucleotides (Fig. 2). Three putative polyadenylation signal sequences appeared at 33, 181 and 313 nt downstream of the TAA termination codon. QsCAD1 open reading frame codes for a protein that is 80.7% and 76.5% homologous to the CADs identified in M. x domestica and E. gunnii, respectively. QsCAD1 and C9 sequences have almost complete homology until nucleotide 111 of C9. Downstream of this nucleotide, C9 and QsCAD1 sequences diverge significantly from each other (Fig. 3).

3.2. Increased expression of QsCAD1 gene in roots of Q. suber infected with P. cinnamomi analysed by RT-PCR and hybridization

To observe the expression pattern of QsCAD1, total RNA extracted from roots of Q. suber seedlings, either noninfected or infected with P. cinnamomi was analysed by RT-PCR and hybridization. The synthesis of complete cDNAs was primed with a 54 bp oligonucleotide (GeneRacer™ Oligo dT Primer) containing a dT tail of 18 nucleotides and a priming site of known sequence for specific primers. In order to select specific QsCAD1 sequences, an amplification reaction was performed with serial dilutions of cDNA solution (1:50; 1:100; 1:200), and specific primers to the QsCAD1 and to the oligonucleotide used to prime the cDNA synthesis (vide Section 2). Simultaneously, control primers for the constitutively expressed gene actin were also used. These primers were designed in the more conserved regions of plant actin genes and amplified a 500 bp fragment. The amplification products were separated on an agarose gel electrophoresis,

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1GenBank accession no: AF053084.
2GenBank accession no: X88797.
transferred to a Hybond-N+ membrane and detected by hybridization with a digoxigenin labelled probe, specific and internal to the amplified fragment (Fig. 4). The intensity of the emitted hybridization signals was measured by densitometric scanning; the absorbance values were found to be directly correlated with the amount of QsCAD1 mRNA, accumulated either in noninfected roots of Q. suber seedlings or in roots infected by P. cinnamomi.

Fig. 4 shows the expression of QsCAD1 gene in non-infected roots of Q. suber seedlings and in roots exposed 24 h to infection by P. cinnamomi. The intensity of the absorbance signal is higher in the infected roots for all the cDNA dilutions decreasing, as expected, with dilution increment, indicating that the expression of QsCAD1 increased in the first 24 h of the interaction between Q. suber and P. cinnamomi.

In fact, two cDNA fragments, slightly longer than 1000 pb, were detected by hybridization with the probe specific to QsCAD1. The two amplified cDNA fragments correspond probably to two different copies of the QsCAD1 gene or to two alleles of the same gene. However, the presence of three putative polyadenylation sites located in the 3' UTR of the gene upstream from the poly(A) tail, and separated by 127 and 142 nucleotides, suggests that transcripts of different sizes are produced (Fig. 4). Similar results were obtained by Goffner et al. [32] in E. gunnii and in tobacco. The authors identified two genes in E. gunnii genome coding for CAD 1 that differed slightly in their coding regions, and isolated two distinct CAD1 cDNAs in tobacco.

The extremely high sensitivity of RT-PCR allows the detection of rare mRNAs, mRNAs in a small number of...
Figure 4. Expression of *QsCAD* gene from *Q. suber* non-infected (R) and infected (RI) roots. The infection was performed by putting the roots in contact with *P. cinnamomi* mycelia for 24 h. RT-PCR products from total RNA were separated by agarose gel electrophoresis, transferred to a Hybond N+ membrane, denatured and hybridized with a digoxigenin labelled probe. The intensity of the hybridization signals was measured by densitometric scanning. Integration values of absorbance curve’s areas are shown at the bottom. Lanes 1 and 2: molecular markers; lanes 3–8: cDNA synthesis solutions were diluted 50, 100 and 200 times as indicated and 10 μL of these dilutions were used as cDNA template for amplification reactions; lane 9: actin cDNA.
homologous (>96%) to QrCAD1 deduced from the cDNA of *Q. ilex* subsp. *rotundifolia* (our unpublished results) and to MdCAD1 (>80%), the protein deduced from the *MdCAD1* gene of *Malus x domestica*.

EgCAD1 (*E. gunnii*), MdCAD1, QsCAD1, QrCAD1 have completely divergent primary sequences from CADs present in clusters A and B, lacking all the conserved domains typical of alcohol dehydrogenases. Instead, the N-terminal part of proteins from cluster C, is the most conserved among all these proteins and VR-ERE, and includes a block of 21 amino acids corresponding to the putative NAD/NADP(H) binding site [43]. This motif is also found in proteins including cinnamoyl-CoA reductases (CCR) from *Eucalyptus gunnii*3 and dihydroflavonol

3GenBank accession nos.: CAA56103; CAA66063.
reductases (DFR) from several plant species. It is also present in the protein encoded by the drought-inducible gene CPRD14 isolated from Vigna unguiculata [44] and in the NADPH-HC toxin reductase gene HMI of maize [45].

4. Discussion

We propose that EgCAD1, MdCAD1, QsCAD1, QrCAD1 and AtCAD3 belong to a group of proteins that diverged from the CADs involved in lignification and suberization. Like VR-ERE, they are probably enzymes whose up-regulation is induced in plant–pathogen compatible interactions with or without expression of disease symptoms. Initially, it was suggested, on the basis of the enzymatic activity of EgCAD1, that CAD1 could function as an alternative enzyme in the lignin biosynthetic pathway [32]. Thus, the possibility that the substrate for QsCAD1 is host origin can not be entirely rejected. The elevated expression of the QsCAD1 could eventually be induced by compounds derived from aromatic polymers released through the action of degradative enzymes produced by the pathogen during infection. However, more recently, it was described [46] that plants down-regulated for CAD1 do not exhibit any changes in their lignin profiles but present significant modifications of their morphology and postulate that CAD1 could thus control a metabolic pathway involved in the production of aromatic substances playing a role in morphogenesis.

EgCAD1 [32] and VR-ERE [10] share a strong affinity towards a larger range of substrates than proteins grouped in cluster A and B, converting benzaldehydes more efficiently than cinnamyl aldehyde derivatives. Nevertheless, these enzymes may also have developed specifically for processing certain substances like toxins, for which they should present a higher catalytic efficiency.

The high structural homology between QsCAD1, QrCAD1, EgCAD1 and VR-ERE, suggests that they have a common enzymatic activity and a similar function. The recombinant form of VR-ERE was shown to have the capacity to reduce aldehydic compounds, namely eutypine in eutypinol [10]. Eutypine, 4-hydroxy-3-(3-methyl-3-buten-1-ynyl) benzylaldehyde is a toxin produced by E. lata, the causal agent of dieback of grapevines. The tolerance of some cultivars to this disease has been shown to be correlated with their capacity to convert eutypine into the corresponding alcohol, eutypinol, which lacks phytotoxicity. Previous studies have shown that eutypine plays an important role in the expression of the disease. It has been suggested [15] that during the growth period of grapevine, eutypine is transported from the trunk through the sap to the herbaceous parts of the vine where it induces a number of symptoms leading to the dead of the whole branch.

Dieback symptoms of grapevines resemble those exhibited by cork oak and holm oak suffering from P. cinnamomi infection. We propose that a mechanism similar to that involving eutypine in the dieback of grapevines caused by E. lata may apply to the sudden death of Quercus spp. involving P. cinnamomi. The occurrence of leaf chlorosis, wilting and death of branches, root necrosis, and sometimes tarry exudations on trunks, is followed by sudden death of crowns. Trees often die suddenly following one or two seasons of decline. These symptoms observed in the field suggest that, following infection, P. cinnamomi produces toxins that migrate through the sap from the roots to the leaves. Trees exhibiting sudden death symptoms are unable to detoxify these toxins. The homology between VR-ERE and QsCAD1, and our evidence for up-regulation of the gene in infected roots of cork oak seedlings, suggests a detoxifying role for the latter that may be inoperative in susceptible hosts, but partially or fully operative in hosts exhibiting chronic decline symptoms. The fragment C9 partially homologous to QsCAD1, isolated and identified exclusively from a cork oak tree with symptoms of sudden death, could derive from a gene coding for an inactive form of QsCAD1. The involved toxins should harbour a benzylaldehyde moiety, a common motive to substrates of VR-ERE. This is not the case of elicitors, polypeptides having toxic properties secreted by almost all Phytophthora species [47]. As a matter of fact no other toxins have been until now identified in Phytophthora species. Should, however these putative toxins be secreted by P. cinnamomi they could account for the remarkably wide host range exhibited by this species. We are presently identifying and investigating the phytotoxic nature of the substances detected in the culture medium of P. cinnamomi.

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