Ectopic Expression of Maize Polyamine Oxidase and Pea Copper Amine Oxidase in the Cell Wall of Tobacco Plants

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To test the feasibility of altering polyamine levels by influencing their catabolic pathway, we obtained transgenic tobacco (Nicotiana tabacum) plants constitutively expressing either maize (Zea mays) polyamine oxidase (MPAO) or pea (Pisum sativum) copper amine oxidase (PCuAO), two extracellular and H₂O₂-producing enzymes. Despite the high expression levels of the transgenes in the extracellular space, the amount of free polyamines in the homozygous transgenic plants was similar to that in the wild-type ones, suggesting either a tight regulation of polyamine levels or a different compartmentalization of the two recombinant proteins and the bulk amount of endogenous polyamines. Furthermore, no change in lignification levels and plant morphology was observed in the transgenic plants compared to untransformed plants, while a small but significant change in reactive oxygen species-scavenging capacity was verified. Both the MPAO and the PCuAO tobacco transgenic plants produced high amounts of H₂O₂ only in the presence of exogenously added enzyme substrates. These observations provided evidence for the limiting amount of freely available polyamines in the extracellular space in tobacco plants under physiological conditions, which was further confirmed for untransformed maize and pea plants. The amount of H₂O₂ produced by exogenously added polyamines in cell suspensions from the MPAO transgenic plants was sufficient to induce programmed cell death, which was sensitive to catalase treatment and required gene expression and caspase-like activity. The MPAO and PCuAO transgenic plants represent excellent tools to study polyamine secretion and conjugation in the extracellular space, as well as to determine when and how polyamine catabolism actually intervenes both in cell wall development and in response to stress.

The polyamines putrescine (Put), spermidine (Spd), and spermine (Spm) are low Mᵦ metabolites naturally found in eukaryotic and prokaryotic cells (Cohen, 1998). Because of their polycationic nature at physiological pH, polyamines can bind strongly to negative charges in cellular components such as nucleic acids, various types of proteins, and acidic phospholipids (Cohen, 1998). In plant cells, polyamines can also be conjugated via an amide bond to hydroxycinnamic acids (Martin-Tanguy, 1997) and serve as precursors for secondary metabolites like nicotine. The function of the conjugated forms of polyamines is not known. However, they have long been associated with flowering (Martin-Tanguy, 1997) and plant-microbe interactions (Torrigiani et al., 1997; Mackintosh et al., 2001; Cowley and Walters, 2002).

Polyamines play important roles in DNA stabilization, RNA and protein synthesis, membrane stabilization, modulation of ion channels, and protection against oxygen radicals, and they are essential for cell homeostasis, cell growth, and tumorigenesis (Wallace et al., 2003). In particular, plant polyamines have been implicated in a variety of plant growth and developmental processes involving cell proliferation and differentiation, morphogenesis, seed dormancy and germination, tuberization, flower induction and development, fruit growth and ripening, embryogenesis, and senescence (for review, see Kumar et al., 1997; Tiburcio et al., 1997; Walden et al., 1997; Malmberg et al., 1998; Bouchereau et al., 1999). Polyamine involvement in defense mechanisms during biotic and abiotic stress (such as mineral nutrient deficiency, osmotic, salt, heat, chilling, wounding, oxidative stress, and pathogen infection) has also been demonstrated (Torrigiani et al., 1997; Yamakawa et al., 1998;
Polyamines are catabolized by the action of amine oxidases. Amine oxidases include the copper-containing amine oxidases (CuAO; EC 1.4.3.6), oxidizing the diamines Put and cadaverine at the primary amino groups, and the flavin-containing polyamine oxidases (PAO), which oxidize Spd and Spm at their secondary amino groups (Federico and Angelini, 1991). In plants, these enzymes are preferentially associated with the primary and secondary wall cells of tissues undergoing lignification, suberization, and wall stiffening (such as xylem, xylem parenchyma, endodermis, and epidermis), although their association to cortical parenchyma cell walls during specific developmental stages has also been reported (Federico and Angelini, 1991; Slocum and Furey, 1991; Liu et al., 1995; Laurenzi et al., 2002). CuAO reaction products from Put are ammonia, γ-aminobutyric acid (GABA) in a reaction catalyzed by pyrroline dehydrogenase (Flores and Filner, 1985). GABA is subsequently transaminated and oxidized to succinic acid, which is incorporated into the Kreb's cycle, thus ensuring the recycling of carbon and nitrogen from Put.

Far from being only a means of eliminating cellular polyamines, the enzymes involved in polyamine catabolism and the products deriving from their action contribute to important physiological processes (Martin-Tanguy, 1997; Šebela et al., 2001). The production of H$_2$O$_2$ through polyamine oxidation has been correlated with the oxidative burst, cell death, lignification, and suberization processes occurring during development and defense responses (Allan and Fluur, 1997; Moller and McPherson, 1998; Rea et al., 1998, 2002; Cona et al., 2003; Walters, 2003). PAO are also involved in the production of uncommon polyamines, such as norspermidine, norspermine, car-dopamine, cadohexamine, homocadopamine, and homocadohexacaine (Phillips and Kuehn, 1991). Uncommon polyamines are of interest because they may be involved in mediating growth response of various organisms under extreme environmental conditions (Phillips and Kuehn, 1991).

In the last few years, mutant plants with aberrant polyamine biosynthesis and transgenic plants exhibiting overexpression or down-regulation of enzymes involved in polyamine metabolic pathways have been used for the study of polyamine function (DeScenzo and Minocha, 1993; Kumar et al., 1996; Burtin and Michael, 1997; Watson et al., 1998; Rafart-Pedros et al., 1999; Capell et al., 2000; Noury et al., 2000; Wisniewski and Brewin, 2000; Mehta et al., 2002; Thu-Hang et al., 2002). Most of these studies have focused on the polyamine biosynthetic pathways and only a few on the catalytic pathways (Wisniewski and Brewin, 2000). Furthermore, in several of these studies, only relatively weak variations in polyamine levels have been observed, indicating the occurrence of compensatory mechanisms to maintain polyamine homeostasis.

In order to test the feasibility of altering polyamine levels by influencing their catabolic pathway and to gain some insight into the physiological processes in which both polyamines and polyamine catabolism are involved, transgenic tobacco (Nicotiana tabacum) plants constitutively expressing either maize (Zea mays) polyamine oxidase (MPAO) or pea (Pisum sativum) copper amine oxidase (PCuAO) were obtained. The transgenic plants appeared morphologically normal, and their polyamine and lignin content was comparable to that of wild-type plants. Though exhibiting a small but significant increase in reactive oxygen species (ROS)-scavenging ability compared with wild-type plants, MPAO and PCuAO transgenic plants and cell suspensions derived from them produced high levels of H$_2$O$_2$ only in the presence of exogenously supplied enzyme substrate, indicating that polyamines in the extracellular space are limiting. Such an oxidative stress was sufficient to induce a programmed cell death (PCD)-like cell death in cell suspensions obtained from MPAO transgenic plants.

RESULTS

Molecular Characterization of MPAO and PCuAO Transgenic Tobacco Plants

Tobacco plants were transformed with MPAO-Ω-pBI or PCuAO-pBI constructs (Fig. 1A) containing the cDNA encoding for MPAO (Tavladoraki et al., 1998) or PCuAO (Tipping and McPherson, 1995), respectively, under the control of the 35S cauliflower mosaic virus promoter. In the MPAO-Ω-pBI construct, the Ω element from tobacco mosaic virus (TMV) was also inserted to obtain elevated expression levels.

Transgenic plants were selected in the presence of kanamycin, and several independent transgenic lines were obtained from two distinct transformation experiments. Primary MPAO or PCuAO transformants (T0 generation) were screened for the presence of the transgenes by PCR after alkali treatment to amplify a 0.87-kb or a 0.75-kb fragment of the MPAO or the PCuAO cDNA, respectively (Fig. 1B). No amplification product was obtained from the untransformed plants either with the MPAO- or the PCuAO-specific oligonucleotides (Fig. 1B).

MPAO and PCuAO expression was determined in the leaves of primary transformants by western-blot analysis and enzyme activity assays (Fig. 2). The two methods gave comparable results and revealed different amounts of recombinant protein among the various transgenic lines. The maximum expression level observed for recombinant MPAO (transgenic line...
MPAO-13) was 0.01 mg of MPAO per milligram of total soluble proteins, which is about 10-fold higher than that of the native enzyme in maize leaves. For recombinant PCuAO, the maximum expression level observed (transgenic line PCuAO-20) was 0.088 µg of PCuAO per milligram of total soluble proteins, which is about 10-fold less than that of the native enzyme in the pea seedlings. No protein similar to MPAO and PCuAO was detectable in untransformed plants by both enzyme activity assays and western-blot analysis utilizing anti-MPAO and anti-PCuAO polyclonal antibodies (Fig. 2).

The inheritance of the MPAO or PCuAO transgenes was studied in three transgenic lines showing different expression levels of the transgenes. Primary transformants were selfed and the seeds harvested after maturation. Upon germination on medium containing kanamycin (T1 generation), we observed for all six

![Figure 1](image.png)

**Figure 1.** Generation and molecular characterization of transgenic tobacco plants expressing MPAO or PCuAO. A, Map of the MPAO-Ω-pBI and PCuAO-pBI constructs used for Agrobacterium-mediated transformation of tobacco plants. Primers (Pr/Pf and Dr/Df) used for PCR analysis and lengths of amplified fragments are indicated. B, PCR analysis of alkali-treated leaf pieces from tobacco plants transformed with the MPAO-Ω-pBI and PCuAO-pBI constructs. Numbers represent independent MPAO and PCuAO transgenic lines. C+, positive control using purified MPAO or PCuAO cDNA; C1 and C2, negative controls using leaf pieces from untransformed plants and MPAO- or PCuAO-specific primers, respectively; M, M, marker (1-kb DNA ladder; Life Technologies/Gibco-BRL, Cleveland).

![Figure 2](image.png)

**Figure 2.** Expression levels of MPAO and PCuAO in different transgenic tobacco lines. Expression levels were determined by enzyme activity assays (top) and western-blot analysis (bottom) in crude leaf extracts. Values are mean ± SE from three replicates. Numbers represent independent MPAO and PCuAO transgenic lines. WT, untransformed tobacco plants; C+, purified MPAO or PCuAO protein. For western-blot analysis, extracts were normalized for the amount of total soluble proteins.
overexpression of maize polyamine and pea copper amine oxidase

Localization of Transgene Expression

Native MPAO and PCuAO enzymes are prevalently apoplastic proteins associated with plant cell walls. Apoplastic localization is also expected for the recombinant proteins in the transgenic plants since the whole cDNAs, including the sequences encoding for the signal peptides that guide protein entry into the endoplasmic reticulum, were inserted in the transformation constructs.

To confirm the extracellular localization of the recombinant proteins in the transgenic plants, intercellular fluids from the MPAO-13H and PCuAO-20H transgenic plants were tested for transgene accumulation both by western-blot analysis (data not shown) and enzyme activity assays (Table I). As a control, intercellular fluids were also tested for levels of the cytosolic marker Glc-6-phosphate dehydrogenase (data not shown) and extracellular peroxidases (Table I). Data demonstrated that the MPAO- and PCuAO-specific activities are at least 10-fold higher in the intercellular fluids (fractions F1 and F2; Table I) than in the crude extracts (fraction T; Table I), confirming a prevalently extracellular localization for the two recombinant enzymes. On the contrary, PAO and CuAO activity could not be detected in intercellular fluids obtained from wild-type plants under our experimental conditions. Extracellular localization of the two recombinant enzymes in the transgenic plants was also confirmed by the lack of MPAO and PCuAO enzyme activity in protoplasts obtained from these plants.

Table I. Analysis of intercellular fluids from MPAO or PCuAO tobacco plants for transgene expression levels

<table>
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<th>MPAO Specific Activity</th>
<th>PCuAO Specific Activity</th>
<th>POD Specific Activity</th>
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<tbody>
<tr>
<td></td>
<td>units/mg tot. prot.</td>
<td>units/mg tot. prot.</td>
<td>units/mg tot. prot.</td>
</tr>
<tr>
<td>F1</td>
<td>3.0</td>
<td>7.6 × 10⁻²</td>
<td>2.5 × 10⁻²</td>
</tr>
<tr>
<td>F2</td>
<td>3.3</td>
<td>7.0 × 10⁻²</td>
<td>0.4 × 10⁻²</td>
</tr>
<tr>
<td>T</td>
<td>0.5</td>
<td>7.0 × 10⁻³</td>
<td>1.0 × 10⁻³</td>
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Transgene enzyme activity could be detected only upon cell wall formation (data not shown).

H₂O₂ Levels in Transgenic Plants Overexpressing MPAO or PCuAO

The homozygous transgenic lines MPAO-13H and PCuAO-20H were analyzed for H₂O₂ production in different tissues (leaf discs, stem sections, and roots) by placing them onto agar plates containing KI and starch (Olson and Varner, 1993). H₂O₂ diffusing from plant tissues is indicated by the purple-blue color in surrounding areas of the medium. Using such an assay, H₂O₂ production could not be detected in any of the tissues analyzed, either from transgenic or wild-type plants up to 24 h of incubation. Elevated levels of H₂O₂ production were observed in all tissues tested of both MPAO and PCuAO transgenic plants only upon supply of enzyme substrate (1 mM Spd and 1 mM Put, respectively; Fig. 3). On the contrary, addition of Spd or Put to the various tissues of wild-type plants did not result in detectable H₂O₂ production (Fig. 3). The level of H₂O₂ production was higher in all tissues of MPAO transgenic plants than in those of PCuAO ones. This difference reflects the expression levels of recombinant MPAO and PCuAO in the transgenic plants. Furthermore, the above results also indicate that the limiting factor for H₂O₂ production in the transgenic tobacco plants is the polyamine content of the apoplast, where the recombinant proteins are localized.

Using the same method, leaf discs from maize and pea seedlings expressing MPAO and PCuAO, respectively, were also tested for H₂O₂ production (Fig. 3). Similar to the transgenic plants, H₂O₂ was produced from maize and pea leaf discs only upon addition of Spd or Put, respectively, suggesting that polyamine transport to the apoplast is limited also in these plants.

In Situ Detection of H₂O₂ in Transgenic Plants Overexpressing MPAO or PCuAO

To exclude the possibility that the H₂O₂ detected in the transgenic segments by the KI/starch assay is a product of the recombinant enzymes liberated from the damaged cells at the tissue cutting site, H₂O₂ production in the MPAO and PCuAO transgenic plants also was evaluated in situ by allowing leaves to take up 3,3-diaminobenzidine (DAB), which in the presence of peroxidases polymerizes as soon as it comes into contact with H₂O₂, forming a brown precipitate. DAB polymerization was only observed in the transgenic plants, which had absorbed DAB in the presence of 1 mM exogenous polyamines (Fig. 4, A and C). As observed using the KI/starch assay, staining intensity was proportional to enzyme expression, and no precipitate was detected in polyamine-treated (Fig. 4, B and D) or untreated (Fig. 4G) wild-type plants. Furthermore, H₂O₂ production was not detected in transgenic leaves, which had not been treated with lines a segregation ratio of 3:1, indicating the presence of a single integration locus.

None of the transgenic plants of the T1 and T2 generations exhibited an altered morphology during normal growth. Furthermore, they exhibited similar accumulation levels of the recombinant proteins to those of the corresponding primary transformants in all tissues tested (leaves, roots, and stems). Homozygous transgenic lines of the T2 generation from primary transformants having the highest transgene expression levels (transgenic lines MPAO-13H and PCuAO-20H) were selected for further analysis.
enzyme substrate (Fig. 4, E and F), confirming that the limiting factor for H$_2$O$_2$ production in the transgenic plants is polyamine levels in the extracellular space. These results also suggest that MPAO and PCuAO proteins are functionally expressed in situ in tobacco transgenic plants.

### Polyamine and Lignin Content in MPAO and PCuAO Transgenic Plants

The leaves of the MPAO-13 and PCuAO-20 transgenic lines were analyzed for polyamine levels. Despite the high expression levels of transgenes, only a slight, statistically not significant reduction in the levels of total free polyamines was observed in the MPAO and PCuAO transgenic plants compared with untransformed plants (Table II). To determine whether the two extracellularly localized recombinant proteins interfere with polyamine levels in the apoplast, intercellular fluids were examined for polyamine content. However, polyamines could not be detected in the apoplast either of transgenic or wild-type plants, as reported previously by Yamakawa et al. (1998) and Yoda et al. (2003).

It has been suggested that H$_2$O$_2$ produced by polyamine oxidation in the cell wall may be involved in peroxidase-mediated lignification, suberization, and cell wall polymer cross-linking occurring during ontogenesis and defense responses (Møller and McPherson, 1998; Rea et al., 1998, 2002; Wisniewski et al., 2000). To investigate whether MPAO or PCuAO expression in the tobacco transgenic plants interferes with cell wall development, the transgenic plants were also histochemically analyzed for lignin content. No significant changes in lignification levels were observed in both transgenic lines compared to untransformed plants (data not shown).

### The Cellular Redox State in MPAO and PCuAO Transgenic Plants

In order to verify whether overexpression of MPAO or PCuAO in the transgenic tobacco plants affected the ROS-scavenging activity, levels of ascorbate peroxidase (APX), catalase, dehydroascorbate reductase (DHAR), ascorbate (ASC), and dehydroascorbate (DHA) were also determined in transgenic and wild-type plants.

As shown in Table III, APX level was significantly increased in both transgenic lines as compared to the wild-type plants, while catalase activity did not vary. Furthermore, ASC content and redox state, i.e. the ratio between the reduced form of ASC and the total ASC pool (ASC + DHA), were increased only in the PCuAO transgenic plants (Table III). Interestingly, DHAR was significantly enhanced only in these plants. This could explain the shift toward the reduced form of ASC pool in the PCuAO transgenic plants.
Moreover, since DHA is an unstable molecule, which is quickly degraded when it is not promptly reduced to ASC, the rise in DHAR activity could also explain the increased amount of the ASC pool present in the PCuAO transgenic plants.

Characterization of Cell Suspensions Obtained from MPAO Transgenic Plants

Cell suspensions were obtained from the leaves of wild-type and MPAO-13H transgenic plants and tested for $\text{H}_2\text{O}_2$ accumulation. The transgenic cell suspension maintained high expression levels of the recombinant MPAO, i.e. 20 µg of MPAO per milliliter packed cell volume. $\text{H}_2\text{O}_2$ could not be detected in the medium of either the MPAO or the wild-type cell suspension in the absence of exogenous Spd. Addition of Spd at a final concentration of 2 mM or 6 mM resulted in a steady-state accumulation of $\text{H}_2\text{O}_2$ in the culture medium of only the MPAO cell suspension (reaching a maximum concentration of about 0.18 mM with 2 mM Spd and 0.3 mM with 6 mM Spd after 0.5 h) and not in that of the wild-type one (Fig. 5). The $\text{H}_2\text{O}_2$ produced by the MPAO cell suspensions was quickly eliminated upon complete oxidation of the exogenously added Spd (Fig. 5), probably due to the presence of an efficient ROS-scavenging system (Fig. 5). Indeed, exogenously supplied $\text{H}_2\text{O}_2$ (0.35 mM) was immediately eliminated (Fig. 6). The wild-type and transgenic cell suspensions demonstrated a similar rate of $\text{H}_2\text{O}_2$ degradation (Fig. 6), which was not influenced by the presence of Spd (Fig. 6).

Addition of 2 mM or 6 mM Spd significantly enhanced cell death in the MPAO cell suspension, whereas very little cell death was observed in the wild-type cell suspension treated with the same amount of Spd or in the untreated MPAO cells (Fig. 7). To test whether the increased cell death observed in Spd-treated MPAO cells was due to an increase in $\text{H}_2\text{O}_2$ levels, cultures were preincubated with catalase before addition of Spd. Catalase treatment reduced both the steady-state levels of $\text{H}_2\text{O}_2$ generated by the oxidation of 6 mM Spd (Fig. 8A) and the cell-death response (Fig. 8B) in the MPAO cell suspension, indicating that the increased cell death observed was mainly $\text{H}_2\text{O}_2$ dependent.

Recently, it has been demonstrated that $\text{H}_2\text{O}_2$-induced cell death is a programmed event requiring de novo transcription and translation (Desikan et al., 1998; Solomon et al., 1999; Clarke et al., 2000). To determine whether the Spd-induced oxidative stress in the MPAO-expressing tobacco cell suspensions requires gene expression, cells were preincubated with cycloheximide, an inhibitor of translation, for 30 min before the induction of oxidative stress. Cycloheximide reduced cell death induced by 2 mM or 6 mM Spd by 29% and 45%, respectively (Fig. 9A), suggesting that the cell death resulting from the oxidative stress in

Table II. Free polyamine content in young leaves of wild-type and MPAO- or PCuAO-expressing tobacco plants

Polyamine levels were determined in leaves of 30-d-old wild-type and transgenic plants. The values represent the mean ± s.d. between independent plants ($n = 10$).

<table>
<thead>
<tr>
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<th>Put (nmol/g fresh weight)</th>
<th>Spd (nmol/g fresh weight)</th>
<th>Spm (nmol/g fresh weight)</th>
</tr>
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<tbody>
<tr>
<td>WT</td>
<td>275.5 ± 79.0</td>
<td>130.3 ± 50.0</td>
<td>12.8 ± 5.8</td>
</tr>
<tr>
<td>MPAO</td>
<td>217.5 ± 56.4</td>
<td>102.5 ± 35.9</td>
<td>8.3 ± 3.5</td>
</tr>
<tr>
<td>PCuAO</td>
<td>242.4 ± 59.9</td>
<td>126.2 ± 17.5</td>
<td>11.2 ± 2.6</td>
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the MPAO-expressing tobacco cell suspensions is an active process that requires the expression of new proteins. Furthermore, to verify whether the oxidative stress generated by Spd in the MPAO cell suspension triggered a signaling mechanism similar to that described for H$_2$O$_2$- and NO-dependent cell death in soybean (Glycine max) and Arabidopsis cultures (Solomon et al., 1999; Clarke et al., 2000; Tiwari et al., 2002), such suspensions were preincubated with Z-YVAD, a competitive and irreversible inhibitor of caspase-1 and -4, for 30 min before addition of Spd. Addition of the caspase inhibitor reduced the cell death induced by 2 mM Spd by about 40% and that induced by 6 mM Spd by about 28% (Fig. 9B), suggesting that the oxidative stress induced an active PCD-like pathway. In these experiments, the levels of cell death induced by Spd appeared higher than in the previous ones, probably because of the presence of dimethylsulfoxide at a final concentration of 0.2% (v/v), used as a solvent for the caspase inhibitor.

**DISCUSSION**

There is increasing evidence that PAO and CuAO may be implicated in plant development and defense responses (Rea et al., 1998; Cowley and Walters, 2002; Cona et al., 2003; Yoda et al., 2003). Considering their extracellular localization, these H$_2$O$_2$-generating enzymes may act together with peroxidases to mediate protein cross-linking and lignin deposition in plant cell walls under mechanical or physiological stress. It also has been demonstrated that H$_2$O$_2$ produced by polyamine oxidation is correlated with oxidative stress and cell death during ontogenesis and in response to biotic and abiotic stress (Allan and Fluhr, 1997; Torrigiani et al., 1997; Möller and McPherson, 1998; Cowley and Walters, 2002; Rea et al., 2002; Yoda et al., 2003). In addition, CuAO and PAO may serve to modify polyamine levels and hence influence polyamine metabolism.

To gain a further insight into the physiological role of the two amine oxidases in plants and to verify the possibility of interfering with polyamine homeostasis, we obtained transgenic tobacco plants overexpressing MPAO or PCuAO. Although CuAO and PAO activity has recently been detected in wild-type plants of *N. tabacum* cv Samsun at very low amounts and after enzyme enrichment or using sensitive enzymatic assays (Biondi et al., 2001; Yoda et al., 2003), enzymes with similar molecular and catalytic properties to those of MPAO or PCuAO were not detectable under our experimental conditions in the wild-type tobacco plants used in this study (*N. tabacum* cv SR1). Thus, *N. tabacum* cv SR1 plants provide a suitable system for transgenic studies on the MPAO and PCuAO genes.

In this study, we show that the MPAO and PCuAO transgenic plants and MPAO cell suspensions produce high amounts of H$_2$O$_2$, compared to the untransformed ones, only in the presence of exogenously added substrate, leading to the conclusion that the two recombinant proteins are functionally expressed in the tobacco plants and that the limiting factor for their activity in the tobacco plants is the amount of...
polyamines available in the extracellular space. Polyamine transport to the extracellular space seems to be the limiting factor for H₂O₂ production by MPAO and PCuAO not only in tobacco plants but also in maize and pea plants (Fig. 3), indicating similar regulatory mechanisms of polyamine metabolism among the various plant species.

Despite the high expression level of the transgenes in the MPAO and PCuAO transgenic plants (1% of total soluble proteins for MPAO and 0.01% for PCuAO; Fig. 2), levels of free polyamines were only slightly reduced (Table II). This is probably due to a fine regulation of polyamine levels, which is achieved via a balance of their biosynthesis, degradation, uptake, transport, and conjugation. It may also be due to inaccessibility of MPAO and PCuAO in the transgenic plants by the bulk amount of polyamines owing to a different compartmentalization. Indeed, our results demonstrated a prevalently extracellular localization of the recombinant MPAO and PCuAO proteins in the transgenic tobacco plants, similar to the native enzymes in maize and pea plants, respectively (Federico and Angelini, 1991; Laurenzi et al., 2001). On the other hand, under our experimental conditions free polyamines could not be detected in the intercellular fluids of either wild-type or transgenic tobacco plants. Furthermore, vacuoles have been proposed as temporary storage sites of polyamines (Pistocchi et al., 1988). The question thus arises as to why maize and pea plants produce high amounts of amine oxidases in the apoplast in the absence of comparably high levels of enzyme substrate. It is possible that controlled polyamine transport to the apoplast may occur during certain physiological or stress conditions. Little is known about transport of diamines and polyamines around plants or about their distribution and their roles in the extracellular environment. In TMV-infected tobacco plants, it has been demonstrated that Put and Spm accumulate in the intercellular space during hypersensitive response (HR; Yamakawa et al., 1998; Yoda et al., 2003). Furthermore, enzyme activity of Orn decarboxylase and Arg decarboxylase increases during HR in tobacco plants infected with TMV, resulting in elevated concentrations of their products and their conjugates, mainly in necrotic regions (Torrigiani et al., 1997). In several cases, conjugated polyamines are also produced in response to abiotic injury and mineral nutrient deficiency. In their conjugated forms, polyamines are often associated with a variety of phenolic cell wall components, such as hydroxycinnamic acids. It is not known whether the conjugated polyamines in the plant cell wall have a structural role or serve as polyamine storage forms or whether they are biologically active molecules. Small amounts of pectin-associated polyamines are also present in the cell wall, which have been proposed to be part of an intrinsic signaling network of the extracellular matrix by modulating the transduction of the pectic signal (Messiaen and Van Cutsem, 1999).

Lignification levels in young MPAO- and PCuAO-expressing transgenic plants appeared quite similar to those of the untransformed plants, despite the high expression levels of the transgenes. This may reflect the lack of polyamine accumulation in the apoplasm and suggests that polyamine catabolism is not involved in early stages of tobacco plant development.
Figure 8. Induction of cell death by oxidative stress in tobacco cell suspensions. A, Kinetics of H$_2$O$_2$ accumulation induced by 6 mM Spd in MPAO-expressing cell suspensions. Cells were pretreated (Spd + Cat) or not (Spd) with catalase for 30 min before addition of Spd. Data are from a single representative experiment, which was repeated in triplicate. B, Cell death was determined by Evans blue staining in wild-type (W) and MPAO-expressing (T) cell suspensions 24 h after addition (+Spd) or not (−Spd) of 6 mM Spd. Cell death was also determined in cell suspensions pretreated with catalase for 30 min before addition of 6 mM Spd (+Spd+Cat). Each point represents mean value of three independent experiments.

Figure 9. Induction of PCD by Spd-induced production of H$_2$O$_2$ in MPAO-expressing cell suspensions. Cell death was estimated by Evans blue staining in MPAO-expressing cell suspensions (T) 15 h after addition of 0, 2 mM, or 6 mM Spd. Where indicated, 40 μM cycloheximide (A) or 40 μM caspase inhibitor (B) was added to the culture medium 30 min prior to Spd administration. Values are mean ± se (n = 3). Asterisks indicate values significantly different from those of cells treated with the same amount of Spd and not with cycloheximide or caspase inhibitor P < 0.05 (by one-way ANOVA test).
under physiological conditions. However, this may not be true for all plants, as in the case of maize and pea. It is also possible that polyamine catabolism is specifically involved in plant development either during maturation and senescence or under stress conditions.

In this study, a small but significant increase in the ROS-scavenging capability of the MPAO and PCuAO transgenic plants has been demonstrated relative to the wild-type ones. In particular, higher APX levels have been demonstrated in both transgenic lines compared to the wild-type tobacco plants. These data suggest that a small amount of H\textsubscript{2}O\textsubscript{2}, not detectable under our experimental conditions, may be generated by transgene expression, inducing an increase in the ROS-scavenging capacity of the transgenic plants. However, this increase was not high enough to interfere with the high rate of H\textsubscript{2}O\textsubscript{2} degradation in the MPAO cell suspensions when this ROS was exogenously added to the culture medium. In contrast to APX levels, significant variations in catalase activity between transgenic and wild-type plants were not observed. The different behavior of catalase and APX is in accordance with only a slight increase of H\textsubscript{2}O\textsubscript{2} production in transformed plants. Indeed, APX, having a much higher affinity for H\textsubscript{2}O\textsubscript{2} than catalase (Mittler, 2002), is much more sensitive even to low increase in H\textsubscript{2}O\textsubscript{2} level. The different behavior of APX and catalase could also be a consequence of their different localization within cells. Indeed, catalase, which is localized within microbodies and mitochondria, may be less sensitive to H\textsubscript{2}O\textsubscript{2} production in the plant cell wall by MPAO and PCuAO than APX, which is present in almost all cellular compartments, apoplast and cytosol included. It is intriguing that although transgene expression levels in the PCuAO transgenic plants are lower than those in the MPAO plants, ASC content and ASC redox state were enhanced relative to the untransformed plants, only in the former and not in the latter. This rise in ASC content and ASC redox state is probably due to an increased capacity of the PCuAO transgenic plants to reduce DHA, the oxidized form of ASC. Indeed, DHAR activity is also increased in these transgenic plants but not in the MPAO ones. It is possible that Put levels in tobacco plant cell walls are higher than those of Spd or Spm, and this may produce stronger oxidative stress, triggering a cellular homeostatic rebalance of the ASC recycling capability and redox state.

There is evidence that ROS play key roles in the initiation of PCD in both animals and plants, and recent work suggested that H\textsubscript{2}O\textsubscript{2} might determine the HR response in plants (Levine et al., 1994). It has been also demonstrated that H\textsubscript{2}O\textsubscript{2} can induce an apoptotic-like cell-death program in suspension cultures of soybean and Arabidopsis (Desikan et al., 1998; Solomon et al., 1999; Tiwari et al., 2002). Here, we report that despite the presence of an efficient system of ROS-scavenging in the tobacco cell suspensions, the level of oxidative stress generated by exogenously supplied substrate in MPAO-expressing cell suspensions is sufficient to induce cell death, the latter being higher in the presence of higher concentrations of enzyme substrate. The levels of H\textsubscript{2}O\textsubscript{2} accumulated in the MPAO-expressing cell suspensions (a maximum of about 0.3 mM in 1 h after addition of 6 mM Spd) was similar to that produced in Arabidopsis cell suspensions using Glc oxidase/Glc as an H\textsubscript{2}O\textsubscript{2}-generating system (Tiwari et al., 2002). Furthermore, the oxidative stress generated in the MPAO tobacco cell suspensions triggered a cell-death program requiring gene expression and caspase-like activity, similar to that observed in Arabidopsis and soybean cell suspensions (Desikan et al., 1998; Solomon et al., 1999; Tiwari et al., 2002). However, using the caspase inhibitor Z-YVAD, the reduction of cell death induced by 6 mM Spd appeared lower than that induced by 2 mM Spd. It is possible that the high oxidative stress induced by 6 mM Spd triggered cell death also via an uncontrolled necrotic pathway or via a caspase-independent pathway.

In conclusion, present data indicate that under physiological conditions the amount of freely available polyamines in the apoplast and cell wall is the limiting factor that regulates the catabolic pathway linked to polyamines. The fact that MPAO or PCuAO transgenic plants are normal both in their developmental pattern and metabolism and that an increase in cell wall polyamine content can be evidenced via H\textsubscript{2}O\textsubscript{2} production renders them an excellent tool for the study of polyamine secretion and conjugation. In addition, they may be used to determine how and when these amine oxidases actually intervene both in cell wall development and in response to stress.

MATERIAL AND METHODS

Plant Material and Growth Conditions

Plants of tobacco (Nicotiana tabacum cv Petit Havana SRI), maize (Zea mays L. cv DK 300; Monsanto, Lodi, Italy), and pea (Pisum sativum) were used throughout. Plants and cell cultures were kept in a growth chamber with an irradiance of approximately 150 µmol m\textsuperscript{-2} s\textsuperscript{-1}, a mean temperature of 24°C, and a 16-h daylength.

Vector Construction and Plant Transformation

The cDNAs encoding for MPAO (Tavladoraki et al., 1998) and for PCuAO (Tipping and McPherson, 1995) were inserted in the pBI121 vector (Jefferson et al., 1987) between restriction sites Smal and SacI, after excision of the gene encoding for β-glucuronidase, to form the MPAO-B-pBI and PCuAO-PBI constructs, respectively. In the MPAO-B-pBI construct, the translational enhancer from TMV (Gallei, 1993) was also inserted downstream from 35S cauliflower mosaic virus promoter. Subsequently, the two constructs were inserted into Agrobacterium tumefaciens strain LBA4404 through electroporation and mobilized into tobacco plants via A. tumefaciens-mediated leaf-disc transformation (Horsch et al., 1985). Shoots were regenerated from leaf discs on Murashige and Skoog medium containing 3% (v/v) Suc, 1 mg L\textsuperscript{-1} indole-3-butyric acid, 1 mg L\textsuperscript{-1} 6-benzilaminopurine, 100 mg L\textsuperscript{-1} kanamycin sulfate, 200 mg L\textsuperscript{-1} vancomycin, and 200 mg L\textsuperscript{-1} cefotaxim. Regenerated plants were transferred to rooting medium containing 1% (w/v) Suc, 100 mg L\textsuperscript{-1} kanamycin sulfate, 200 mg L\textsuperscript{-1} vancomycin, 200 mg L\textsuperscript{-1} cefotaxim, and 0.1 mg L\textsuperscript{-1} indole-3-butyric acid. Rooted plants (T0 generation) were either propagated in hormone-free medium containing 100 mg L\textsuperscript{-1} kanamycin sulfate or transferred to soil for seed preparation (T1 generation). Homozygous
transgenic lines were selected by germination of a large amount of seeds obtained from individual plants of T1 generation on medium containing kanamycin.

**Screening of Transgenic Plants by PCR Analysis**

Genomic PCR amplification to detect MPAO and PCuAO cDNAs in transgenic plants was performed from alkali-treated leaf pieces obtained as described by Klimyuk et al. (1993). The PCR reaction was performed in a volume of 50 µL in the presence of 67 mM Tris-HCl, pH 8.8, 16 mM (NH4)2SO4, 0.01% (v/v) Tween 20, 0.5 µM gene-specific primers, 250 µM deoxynucleotide triphosphates, and 0.5 units of Taq DNA polymerase (PolyTaq; Polymed, Firenze, Italy). After an initial denaturation step for 5 min at 94°C, 35 amplification steps were carried out (each comprising denaturation at 94°C for 1 min, annealing at 60°C for 1 min, and extension at 72°C for 1 min), followed by a 10 min extension at 72°C. The primers used for MPAO amplification were Pr (5'- GCCCATCGTCAACTCCACCCTCAAG-3') and Pr (5'-GTACGTCTGTCGTCCTCGTC-3'), and for PCuAO amplification were D1 (5'-GTACCTTATCAGACCCCTACAAG-3') and D3 (5'- CGATTGTGATCATACCTCTCTG-3'). Amplification fragments were analyzed on 1% (w/v) agarose gel.

**Preparation of Intercellular Fluids and Crude Leaf Extracts**

Leaves of tobacco plants were immersed in 10 mM sodium phosphate, pH 6.5 (for MPAO) or pH 7.0 (for PCuAO), and subjected to three consecutive rounds of vacuum for 2 min, followed by release of vacuum. Infiltrated leaves were gently dried and placed in a centrifuge tube on a grid separated from the tube bottom. Intercellular fluid was collected in the bottom of the tube after centrifugation for 15 min at 1,800 g (fraction I). Vacuum infiltration was repeated as below to obtain residual intercellular fluid (fraction II). The rest of the leaves were homogenized in 0.2 m sodium phosphate, pH 6.5 (for MPAO) or pH 7.5 (for PCuAO), and the extract clarified by centrifugation for 10 min at 10,000 g. The resulting supernatant is referred to as the extracellular extract (fraction I).

To prepare a crude protein extract, leaves were homogenized first with liquid nitrogen and then with 0.2 m sodium phosphate, pH 6.5 (for MPAO) or pH 7.0 (for PCuAO). The homogenate was centrifuged for 10 min at 10,000 g, and the supernatant is referred to as the crude leaf extract (fraction I).

All fractions obtained (intercellular fluids, intracellular extracts, and crude extracts) were tested for extracellular peroxidase activities according to Smith and Barker (1988) and for the cytosolic marker Gic-6-phosphate dehydrogenase according to Simcox et al. (1977).

**Determination of MPAO and PCuAO Enzyme Activity**

MPAO and PCuAO enzymatic activities were determined from protein extracts by following the formation of a pink adduct (εmax = 2.6 × 10^5 M⁻¹ cm⁻¹), resulting from the oxidation of 0.1 m 4-aminooantipyrine and of 1.0 m 3,5-dichloro-2-hydroxybenzenesulfonic acid (DCHBS) catalyzed by 0.08 mg (mL of homeradish peroxidase) in 0.2 m sodium phosphate buffer, pH 6.5 (mL of MPAO) or pH 7.0 (mL of PCuAO), at 25°C (Smith and Barker, 1988). One unit of enzyme represents the amount of enzyme that catalyzes the oxidation of 1 µmol of substrate per min. Expression levels were calculated from enzyme activity in the plant protein extracts considering the specific activity of the purified enzymes (80 units mg⁻¹ of protein). Quantification of total soluble proteins was performed using a protein assay kit (Bio-Rad, Hercules, CA) and bovine serum albumin as a standard.

**Western-Blot Analysis**

Western-blot analysis was performed according to Cona et al. (2003), utilizing rabbit anti-MPAO or anti-lentil CuAO polyclonal antibodies purified by affinity chromatography through a Sepharose 4B column (Amersham-Pharmacia Biotech, Uppsala) coupled to bromelain (Sigma, St. Louis).

**Polyamine Analysis**

Total free polyamine levels were determined in both whole tobacco leaves and intercellular spaces. For polyamine extraction from whole leaves, fresh leaf tissues were homogenized initially with liquid nitrogen and then with cold 5% (v/v) perchloric acid (200 mg fresh weight mL⁻¹). Crude extracts were incubated at 4°C for 18 h and were clarified by centrifugation. The supernatant was used to analyze total free polyanimes. For polyamine extraction from the intercellular fluids, 15 leaf discs (15 mm in diameter) were cut out, weighed, and washed with distilled water. Subsequently, the leaf discs were submerged in water in vacuo, and the intercellular fluid, recovered by centrifugation, was immediately mixed with perchloric acid to a final concentration of 5% (v/v).

Free polyanines were quantified after derivatization with dansyl chloride according to Smith and Davies (1985) with minor modifications.Dansylated polyanines were separated by HPLC (PU-980; Jasco, Tokyo) on a reverse-phase C18 column (Spherisorb SS OD25, 5 µm particle diameter, 4.6 × 250 mm) using an acetonitrile to water gradient (60% to 70% acetonitrile in 5.5 min, 70% to 80% in 1.5 min, 80% to 100% in 2 min, 100% for 2 min, 100% to 70% in 2 min, and 70% to 60% in 2 min at a flow rate of 1.5 mL min⁻¹). Eluted peaks were detected by a spectrofluorometer (Jasco 821-FP, excitation 365 nm, emission 510 nm), recorded, and integrated by an attached computer using the JCL6000 software (Jones Chromatography, Hengoed, UK).

**H₂O₂ Detection by KI/Starch Assay in Plant Tissue**

H₂O₂ produced by tissues of transgenic plants was detected essentially as described by Olson and Varner (1993). Leaf discs, roots, and sections of stem were soaked in water and placed onto agar plates containing 50 mM KI and 4% (w/v) potato (Solanum tuberosum) starch. Samples were left at room temperature to allow color development. Color development was also examined after addition of 1 mM Spd or Put solutions on the plant tissue.

**H₂O₂ Detection by 3,3-Diaminobenzidine Uptake Method in Plant Leaves**

In situ H₂O₂ production was detected by an endogenous peroxidase-dependent staining procedure using DAB (Thordal-Christensen et al., 1997). Leaves were detached and placed in a solution of 1 mg mL⁻¹ DAB, pH 3.8, for 18 h under light. Subsequently, the leaves were immersed in boiling 96% (v/v) ethanol for 10 min and then stored in 96% (v/v) ethanol. H₂O₂ production was visualized as a reddish-brown coloration. H₂O₂ production was also examined in the presence of 1 mM Spd or Put in the DAB solution.

**Lignin Histochemical Analysis**

Fresh transversal sections (200 µm) were obtained from the two youngest internodes of 4- to 5-week-old transgenic and wild-type tobacco plants. Lignin deposition was detected by the phloroglucinol/HCl method according to Rea et al. (1998) using a Zeiss Axiophot optical microscope (Jena, Germany).

**Ascorbate System and H₂O₂ Scavenging Enzymes**

Leaves were collected from different three 30-d-old wild-type or transgenic forms. Attention was paid in order to select leaves in the same stage of development (about 3–4 cm in length, at the same internode level). For determination of ASC content and redox state, leaves were homogenized with 4 volumes of cold 5% (w/v) metaphosphoric acid at 4°C in a mortar. The homogenate was centrifuged at 20,000g for 15 min at 4°C, and the supernatant was collected for analysis of ASC according to Zhang and Kirkham (1996). For the assays of ASC redox enzymes and catalase, leaves were ground in liquid nitrogen and homogenized at 4°C in 2 volumes (w/v) of extraction buffer containing 50 mM Tris-HCl, pH 7.8, 0.05% (v/v) Cys, and 0.1% (w/v) bovine serum albumin. The homogenate was centrifuged at 20,000g for 15 min. The supernatant was used for enzymatic assays. Activity of L-ASC-hydrogen peroxide oxidoreductase (APX; EC 1.11.1.11) and of glutathione:dehydroascorbate oxidoreductase (DHAR: EC 1.8.5.1) was determined according to de Pinto et al. (2000). Catalase activity was evaluated according to De Gara et al. (2003).

**Cell Culture Conditions and Treatments**

Young leaves, collected from 3- to 4-week-old in vitro grown wild-type and homozygous MPAO transgenic tobacco plants, were induced to form callus.
on an agaro-solidified Murashige and Skoog medium supplemented with 3% (w/v) Suc, 1 mg L⁻¹ naphthaleneacetic acid, and 0.2 mg L⁻¹ kinetin, pH 5.7. Callus was grown in darkness at 26°C and subcultured on fresh medium every 4 weeks. After three subcultures, fast-growing callus were placed into liquid medium of the same composition. Suspensions were grown for 15 d, and large cell aggregates were eliminated by filtering through a 300-μm mesh filter. Cells settling on a 125-μm mesh filter were resuspended in fresh medium at a 1:10 dilution and grown in the same conditions giving rise to a fast-growing suspension culture. Subculturing was performed every 2 weeks, and cell suspensions were used 4 d after subculture. Before each experiment, the cells were washed with fresh culture medium and resuspended at a 1:4 dilution. For inhibitor experiments, aliquots of cells were pretreated with catalase (2,000 units mL⁻¹), with Z-VAD (40 μM), or with cyclobeximide (40 μM) for 30 min prior to Spd addition. Controls were mock treated with sterile distilled water or dimethylsulfoxide, as appropriate.

To determine H₂O₂ steady-state levels in cell cultures, 40 μL of culture medium were added to a cuvette containing 760 μL of 0.08 mg mL⁻¹ horseradish peroxidase in 0.2 M sodium phosphate buffer, pH 9.0, 100 μL of 1 mM 4-aminoantipyrine, and 100 μL of 10 mM DCHBS. Formation of the colored adduct was measured at 515 nm, after incubation for 5 min at room temperature.

Cell death was quantified by Evans blue staining, as described by Levine et al. (1994). Briefly, samples were incubated for 15 min with 0.05% (w/v) Evans blue and then washed extensively to remove unbound dye. Cells treated with Spd were washed with culture medium before staining to avoid dye precipitation by polyamines. Dye bound to dead cells was solubilized in 50% (v/v) methanol and 1% (w/v) SDS at 60°C for 30 min, and quantified by absorbance at 600 nm. For 100% cell death, the culture was heated at 100°C for 5 min.

Sequence data from this article have been deposited with the EMBL/GenBank data libraries under accession numbers AJ002204 (MaAO) and L39931 (CuAO).

ACKNOWLEDGMENTS

We thank Prof. M.J. McPherson (Centre for Plant Sciences, University of Leeds, Leeds, UK) for the gift of PCuAO cDNA clone, Dr. C. Faso (Biology Department, Università degli Studi Roma Tre) for critical reading of the manuscript, and Dr. D. Pashkoulov (Floramita SpA) for useful discussions and growth of the plants.

Received December 4, 2003; returned for revision January 20, 2004; accepted January 22, 2004.

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